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Brian is a simulator for spiking neural networks. It is written in the Python programming language and is available on almost all platforms. We believe that a simulator should not only save the time of processors, but also the time of scientists. Brian is therefore designed to be easy to learn and use, highly flexible and easily extensible.

To get an idea of what writing a simulation in Brian looks like, take a look at a simple example, or run our interactive demo.

Once you have a feel for what is involved in using Brian, we recommend you start by following the installation instructions, and in case you are new to the Python programming language, having a look at Running Brian scripts. Then, go through the tutorials, and finally read the User Guide.

While reading the documentation, you will see the names of certain functions and classes are highlighted links (e.g. PoissonGroup). Clicking on these will take you to the “reference documentation”. This section is automatically generated from the code, and includes complete and very detailed information, so for new users we recommend sticking to the User’s guide. However, there is one feature that may be useful for all users. If you click on, for example, PoissonGroup, and scroll down to the bottom, you’ll get a list of all the example code that uses PoissonGroup. This is available for each class or method, and can be helpful in understanding how a feature works.

Finally, if you’re having problems, please do let us know at our support page.

Please note that all interactions (e.g. via the mailing list or on github) should adhere to our Code of Conduct.

Contents:
1.1 Installation

We recommend users to use the Anaconda distribution by Continuum Analytics. Its use will make the installation of Brian 2 and its dependencies simpler, since packages are provided in binary form, meaning that they don’t have to be build from the source code at your machine. Furthermore, our automatic testing on the continuous integration services travis and appveyor are based on Anaconda, we are therefore confident that it works under this configuration.

However, Brian 2 can also be installed independent of Anaconda, either with other Python distributions (Enthought Canopy, Python(x,y) for Windows, ...) or simply based on Python and pip (see Installation with pip below).

1.1.1 Installation with Anaconda

Installing Anaconda

Download the Anaconda distribution for your Operating System. Note that the choice between Python 2.7 and Python 3.x is not very important at this stage. Anaconda allows you to create a Python 3 environment from Python 2 Anaconda and vice versa.

After the installation, make sure that your environment is configured to use the Anaconda distribution. You should have access to the conda command in a terminal and running python (e.g. from your IDE) should show a header like this, indicating that you are using Anaconda’s Python interpreter:
Installing Brian 2

Note: The provided Brian 2 packages are only for 64bit systems. If you want to install Brian 2 in a 32bit environment, please use the Installation with pip instead.

You can either install Brian 2 in the Anaconda root environment, or create a new environment for Brian 2 (https://conda.io/projects/conda/en/latest/user-guide/tasks/manage-environments.html). The latter has the advantage that you can update (or not update) the dependencies of Brian 2 independently from the rest of your system.

Brian 2 is not part of the main Anaconda distribution, but built using the community-maintained conda-forge project. You will therefore have to to install it from the conda-forge channel. To do so, use:

```bash
cflush install -c conda-forge brian2
```

You can also permanently add the channel to your list of channels:

```bash
cconda config --add channels conda-forge
```

This has only to be done once. After that, you can install and update the brian2 packages as any other Anaconda package:

```bash
cconda install brian2
```

Installing other useful packages

There are various packages that are useful but not necessary for working with Brian. These include: matplotlib (for plotting), nose (for running the test suite), ipython and jupyter-notebook (for an interactive console). To install them from anaconda, simply do:

```bash
cconda install matplotlib nose ipython notebook
```

You should also have a look at the brian2tools package, which contains several useful functions to visualize Brian 2 simulations and recordings. You can install it with pip or anaconda, similar to Brian 2 itself (but as of now, it is not included in the conda-forge channel, you therefore have to install it from our own brian-team channel), e.g. with:

```bash
cconda install -c brian-team brian2tools
```

1.1.2 Installation with pip

If you decide not to use Anaconda, you can install Brian 2 from the Python package index: https://pypi.python.org/pypi/Brian2

To do so, use the pip utility:
You might want to add the `--user` flag, to install Brian 2 for the local user only, which means that you don’t need administrator privileges for the installation.

Note that when installing `brian2` from source with pip, support for using numerical integration with the GSL requires a working installation of the GSL development libraries (e.g. the package `libgsl-dev` on Debian/Ubuntu Linux).

### 1.1.3 Requirements for C++ code generation

C++ code generation is highly recommended since it can drastically increase the speed of simulations (see Computational methods and efficiency for details). To use it, you need a C++ compiler and either Cython or weave (only available for Python 2.x). Cython/weave will be automatically installed if you perform the installation via Anaconda, as recommended. Otherwise you can install them in the usual way, e.g. using `pip install cython` or `pip install weave`.

**Linux and OS X**

On Linux and Mac OS X, the conda package will automatically install a C++ compiler. But even if you install Brian from source, you will most likely already have a working C++ compiler installed on your system (try calling `g++ --version` in a terminal). If not, use your distribution’s package manager to install a g++ package.

**Windows**

On Windows, the necessary steps to get Runtime code generation (i.e. Cython/weave) to work depend on the Python version you are using (also see the notes in the Python wiki):

- **Python >= 3.5**
  - Make sure that your `setuptools` package has at least version 34.4.0 (use `conda update setuptools` when using Anaconda, or `pip install --upgrade setuptools` when using pip).

- **Python 2.7**
  - Download and install the Microsoft Visual C++ Compiler for Python 2.7

For Standalone code generation, you can either use the compiler installed above or any other version of Visual Studio – in this case, the Python version does not matter.

Try running the test suite (see Testing Brian below) after the installation to make sure everything is working as expected.

### 1.1.4 Development version

To run the latest development code, you can directly clone the git repository at github (https://github.com/brian-team/brian2) and then run `pip install -e .` to install Brian in “development mode”. With this installation, updating the git repository is in general enough to keep up with changes in the code, i.e. it is not necessary to install it again.

Another option is to use pip to directly install from github:

```
pip install https://github.com/brian-team/brian2/archive/master.zip
```
1.1.5 Testing Brian

If you have the nose testing utility installed, you can run Brian’s test suite:

```python
import brian2
brian2.test()
```

It should end with “OK”, showing a number of skipped tests but no errors or failures. For more control about the tests that are run see the developer documentation on testing.

1.2 Running Brian scripts

Brian scripts are standard Python scripts, and can therefore be run in the same way. For interactive, explorative work, you might want to run code in a jupyter notebook or in an ipython shell; for running finished code, you might want to execute scripts through the standard Python interpreter; finally, for working on big projects spanning multiple files, a dedicated integrated development environment for Python could be a good choice. We will briefly describe all these approaches and how they relate to Brian’s examples and tutorial that are part of this documentation. Note that none of these approaches are specific to Brian, so you can also search for more information in any of the resources listed on the Python website.

- Jupyter notebook
- IPython shell
- Python interpreter
- Integrated development environment (IDE)

1.2.1 Jupyter notebook

The Jupyter Notebook is an open-source web application that allows you to create and share documents that contain live code, equations, visualizations and narrative text.

(from jupyter.org)

Jupyter notebooks are a great tool to run Brian code interactively, and include the results of the simulations, as well as additional explanatory text in a common document. Such documents have the file ending .ipynb, and in Brian we use this format to store the Tutorials. These files can be displayed by github (see e.g. the first Brian tutorial), but in this case you can only see them as a static website, not edit or execute any of the code.

To make the full use of such notebooks, you have to run them using the jupyter infrastructure. The easiest option is to use the free mybinder.org web service, which allows you to try out Brian without installing it on your own machine. Links to run the tutorials on this infrastructure are provided as “launch binder” buttons on the Tutorials page, and also for each of the Examples at the top of the respective page (e.g. Example: COBAHH). To run notebooks on your own machine, you need an installation of the jupyter notebook software on your own machine, as well as Brian itself (see the Installation instructions for details). To open an existing notebook, you have to download it to your machine. For the Brian tutorials, you find the necessary links on the Tutorials page. When you have downloaded/installed everything necessary, you can start the jupyter notebook from the command line (using Terminal on OS X/Linux, Command Prompt on Windows):

```
jupyter notebook
```
this will open the “Notebook Dashboard” in your default browser, from which you can either open an existing notebook or create a new one. In the notebook, you can then execute individual “code cells” by pressing \texttt{SHIFT+ENTER} on your keyboard, or by pressing the play button in the toolbar.

For more information, see the \texttt{jupyter notebook documentation}.

\subsection*{1.2.2 IPython shell}

An alternative to using the jupyter notebook is to use the interactive Python shell \texttt{IPython}, which runs in the Terminal/Command Prompt. You can use it to directly type Python code interactively (each line will be executed as soon as you press \texttt{ENTER}), or to run Python code stored in a file. Such files typically have the file ending \texttt{.py}. You can either create it yourself in a text editor of your choice (e.g. by copying&pasting code from one of the \texttt{Examples}), or by downloading such files from places such as github (e.g. the Brian examples), or ModelDB. You can then run them from within IPython via:

\begin{verbatim}
%run filename.py
\end{verbatim}

\subsection*{1.2.3 Python interpreter}

The most basic way to run Python code is to run it through the standard Python interpreter. While you can also use this interpreter interactively, it is much less convenient to use than the IPython shell or the jupyter notebook described above. However, if all you want to do is to run an existing Python script (e.g. one of the Brian \texttt{Examples}), then you can do this by calling:

\begin{verbatim}
python filename.py
\end{verbatim}

in a Terminal/Command Prompt.

\subsection*{1.2.4 Integrated development environment (IDE)}

Python is a widely used programming language, and is therefore support by a wide range of integrated development environments (IDE). Such IDEs provide features that are very convenient for developing complex projects, e.g. they integrate text editor and interactive Python console, graphical debugging tools, etc. Popular environments include Spyder, PyCharm, and Visual Studio Code, for an extensive list see the Python wiki.

\section*{1.3 Release notes}

\subsection*{1.3.1 Brian 2.3}

This release contains the usual mix of bug fixes and new features (see below), but also makes some important changes to the Brian 2 code base to pave the way for the full Python 2 -> 3 transition (the source code is now directly compatible with Python 2 and Python 3, without the need for any translation at install time). Please note that this release will be the last release that supports Python 2, given that Python 2 reaches end-of-life in January 2020. Brian now also uses \texttt{pytest} as its testing framework, since the previously used \texttt{nose} package is not maintained anymore. Since brian2hears has been released as an independent package, using \texttt{brian2.hears} as a “bridge” to Brian 1’s \texttt{brian.hears} package is now deprecated.

Finally, the Brian project has adopted the “Contributor Covenant” \texttt{Contributor Covenant Code of Conduct}, pledging “to make participation in our community a harassment-free experience for everyone”.

\section*{1.3. Release notes}
New features

• The `restore()` function can now also restore the state of the random number generator, allowing for exact reproducibility of stochastic simulations (#1134)
• The functions `expm1()`, `log1p()`, and `exprel()` can now be used (#1133)
• The system for calling random number generating functions has been generalized (see Functions with context-dependent return values), and a new `poisson` function for Poisson-distributed random numbers has been added (#1111)
• New versions of Visual Studio are now supported for standalone mode on Windows (#1135)

Selected improvements and bug fixes

• `run_regularly` operations are now included in the network, even if they are created after the parent object was added to the network (#1009). Contributed by Vigneswaran Chandrasekaran.
• No longer incorrectly classify some equations as having “multiplicative noise” (#968). Contributed by Vigneswaran Chandrasekaran.
• Brian is now compatible with Python 3.8 (#1130), and doctests are compatible with numpy 1.17 (#1120)
• Progress reports for repeated runs have been fixed (#1116), thanks to Ronaldo Nunes for reporting the issue.
• `SpikeGeneratorGroup` now correctly works with `restore()` (#1084), thanks to Tom Achache for reporting the issue.
• An indexing problem in `PopulationRateMonitor` has been fixed (#1119).
• Handling of equations referring to $-\infty$ has been fixed (#1061).
• Long simulations recording more than ~2 billion data points no longer crash with a segmentation fault (#1136), thanks to Rike-Benjamin Schuppner for reporting the issue.

Backward-incompatible changes

• The fix for `run_regularly` operations (#1009, see above) entails a change in how objects are stored within `Network` objects. Previously, `Network.objects` stored a complete list of all objects, including objects such as `StateUpdater` that – often invisible to the user – are a part of major objects such as `NeuronGroup`. Now, `Network.objects` only stores the objects directly provided by the user (`NeuronGroup`, `Synapses`, `StateMonitor`, …), the dependent objects (`StateUpdater`, `Threshold`, …) are taken into account at the time of the run. This might break code in some corner cases, e.g. when removing a `StateUpdater` from `Network.objects` via `Network.remove`.
• The `brian2.hears` interface to Brian 1’s `brian.hears` package has been deprecated.

Infrastructure and documentation improvements

• The same code base is used on Python 2 and Python 3 (#1073).
• The test framework uses `pytest` (#1127).
• We have adapted a Code of Conduct (#1113), thanks to Tapasweni Pathak for the suggestion.
Contributions

Github code, documentation, and issue contributions (ordered by the number of contributions):

- Marcel Stimberg (@mstimberg)
- Dan Goodman (@thesamovar)
- Vigneswaran Chandrasekaran (@Vigneswaran-Chandrasekaran)
- Moritz Orth (@morth)
- Tristan Stöber (@tristanstoeber)
- @ulyssek
- Wilhelm Braun (@wilhelmbraun)
- @flomlo
- Rike-Benjamin Schuppner (@Debilski)
- @sdeiss
- Ben Evans (@bdevans)
- Tapasweni Pathak (@tapaswenipathak)
- @jonathanoesterle
- Richard C Gerkin (@rgerkin)
- Christian Behrens (@chbehrens)
- Romain Brette (@romainbrette)
- XiaoquinNUDT (@XiaoquinNUDT)
- Dylan Muir (@DylanMuir)
- Aleksandra Teska (@alTeska)
- Felix Z. Hoffmann (@felix11h)
- @baixiaotian63648995
- Carlos de la Torre (@c-torre)
- Sam Mathias (@sammosummo)
- @Marghepano
- Simon Brodeur (@sbrodeur)
- Alex Dimitrov (@adimitr)

Other contributions outside of github (ordered alphabetically, apologies to anyone we forgot...):

- Ronaldo Nunes
- Tom Achache

1.3.2 Brian 2.2.2.1

This is a bug-fix release that fixes several bugs and adds a few minor new features. We recommend all users of Brian 2 to upgrade.

1.3. Release notes
As always, please report bugs or suggestions to the github bug tracker (https://github.com/brian-team/brian2/issues) or to the brian-development mailing list (brian-development@googlegroups.com).

[Note that the original upload of this release was version 2.2.2, but due to a mistake in the released archive, it has been uploaded again as version 2.2.2.1]

**Selected improvements and bug fixes**

- Fix an issue with the synapses generator syntax (#1037).
- Fix an incorrect error when using a SpikeGeneratorGroup with a long period (#1041). Thanks to Kévin Cuallado-Keltsch for reporting this issue.
- Improve the performance of SpikeGeneratorGroup by avoiding a conversion from time to integer time step (#1043). This time step is now also available to user code as t_in_timesteps.
- Function definitions for weave/Cython/C++ standalone can now declare additional header files and libraries. They also support a new sources argument to use a function definition from an external file. See the Functions documentation for details.
- For convenience, single-neuron subgroups can now be created with a single index instead of with a slice (e.g. neurongroup[3] instead of neurongroup[3:4]).
- Fix an issue when -inf is used in an equation (#1061).

**Contributions**

Github code, documentation, and issue contributions (ordered by the number of contributions):

- Marcel Stimberg (@mstimberg)
- Dan Goodman (@thesamovar)
- Felix Z. Hoffmann (@Felix11H)
- @wjsx0914
- Kévin Cuallado-Keltsch (@kevincuallado)
- Romain Cazé (@rcaze)
- Daphne (@daphn3cor)
- Erik (@parenthetical-e)
- @RahulMaram
- Eghbal Hosseini (@eghbalhosseini)
- Martino Sorbaro (@martinosoph)
- Mihir Vaidya (@MihirVaidya94)
- @helloolingling
- Volodimir Slobodyanyuk (@vslbody)
- Peter Duggins (@psipeter)
1.3.3 Brian 2.2.1

This is a bug-fix release that fixes a few minor bugs and incompatibilities with recent versions of the dependencies. We recommend all users of Brian 2 to upgrade.

As always, please report bugs or suggestions to the github bug tracker (https://github.com/brian-team/brian2/issues) or to the brian-development mailing list (brian-development@googlegroups.com).

**Selected improvements and bug fixes**

- Work around problems with the latest version of `py-cpuinfo` on Windows (#990, #1020) and no longer require it for Linux and OS X.
- Avoid warnings with newer versions of Cython (#1030) and correctly build the Cython spike queue for Python 3.7 (#1026), thanks to Fleur Zeldenrust and Ankur Sinha for reporting these issues.
- Fix error messages for `SyntaxError` exceptions in jupyter notebooks (#964).

**Dependency and packaging changes**

- Conda packages in `conda-forge` are now available for Python 3.7 (but no longer for Python 3.5).
- Linux and OS X no longer depend on the `py-cpuinfo` package.
- Source packages on `pypi` now require a recent Cython version for installation.

**Contributions**

Github code, documentation, and issue contributions (ordered by the number of contributions):

- Marcel Stimberg (@mstimberg)
- Dan Goodman (@thesamovar)
- Christopher (@Chris-Currin)
- Peter Duggins (@psipeter)
- Paola Suárez (@psrmx)
- Ankur Sinha (@sanjayankur31)
- @JingjinW
- Denis Alevi (@denisalevi)
- @lemonade117
- @wjx0914
- Sven Leach (@SvennoNito)
- svadams (@svadams)
- @ghaessig
- Varshith Sreeramdas (@varshiths)
1.3.4 Brian 2.2

This release fixes a number of important bugs and comes with a number of performance improvements. It also makes sure that simulation no longer give platform-dependent results for certain corner cases that involve the division of integers. These changes can break backwards-compatibility in certain cases, see below. We recommend all users of Brian 2 to upgrade.

As always, please report bugs or suggestions to the github bug tracker (https://github.com/brian-team/brian2/issues) or to the brian-development mailing list (brian-development@googlegroups.com).

Selected improvements and bug fixes

- Divisions involving integers now use floating point division, independent of Python version and code generation target. The `//` operator can now be used in equations and expressions to denote flooring division (#984).
- Simulations can now use single precision instead of double precision floats in simulations (#981, #1004). This is mostly intended for use with GPU code generation targets.
- The `timestep`, introduced in version 2.1.3, was further optimized for performance, making the refractoriness calculation faster (#996).
- The `lastupdate` variable is only automatically added to synaptic models when event-driven equations are used, reducing the memory and performance footprint of simple synaptic models (#1003). Thanks to Denis Alevi for bringing this up.
- A `from brian2 import *` imported names unrelated to Brian, and overwrote some Python builtins such as `dir` (#969). Now, fewer names are imported (but note that this still includes numpy and plotting tools: Importing Brian).
- The `exponential_euler` state updater is no longer failing for systems of equations with differential equations that have trivial, constant right-hand-sides (#1010). Thanks to Peter Duggins for making us aware of this issue.

Backward-incompatible changes

- Code that divided integers (e.g. `N/10`) with a C-based code generation target, or with the `numpy` target on Python 2, will now use floating point division instead of flooring division (i.e., Python 3 semantics). A warning will notify the user of this change, use either the flooring division operator (`N//10`), or the `int` function (`int(N/10)`) to make the expression unambiguous.
- Code that directly referred to the `lastupdate` variable in synaptic statements, without using any event-driven variables, now has to manually add `lastupdate : second` to the equations and update the variable at the end of `on_pre` and/or `on_post` with `lastupdate = t`.
- Code that relied on `from brian2 import *` also importing unrelated names such as `sympy`, now has to import such names explicitly.

Documentation improvements

- Various small fixes and additions (e.g. installation instructions, available functions, fixes in examples)
- A new example, Izhikevich 2007, provided by Guillaume Dumas.
Contributions

Github code, documentation, and issue contributions (ordered by the number of contributions):

- Marcel Stimberg (@mstimberg)
- Dan Goodman (@thesamovar)
- Denis Alevi (@denisalevi)
- Thomas Nowotny (@tnowotny)
- @neworderofjamie
- Paul Brodersen (@paulbrodersen)
- @matrec4
- svadams (@svadams)
- XiaoquinNUDT (@XiaoquinNUDT)
- Peter Duggins (@psipeter)
- @nh17937
- Patrick Nave (@pnaive95)
- @AI-pha
- Guillaume Dumas (@deep-introspection)
- @godelicbach
- @galharth

1.3.5 Brian 2.1.3.1

This is a bug-fix release that fixes two bugs in the recent 2.1.3 release:

- Fix an inefficiency in the newly introduced `timestep` function when using the numpy target (#965)
- Fix inefficiencies in the unit system that could lead to slow operations and high memory use (#967). Thanks to Kaustab Pal for making us aware of the issue.

1.3.6 Brian 2.1.3

This is a bug-fix release that fixes a number of important bugs (see below), but does not introduce any new features. We recommend all users of Brian 2 to upgrade.

As always, please report bugs or suggestions to the github bug tracker (https://github.com/brian-team/brian2/issues) or to the brian-development mailing list (brian-development@googlegroups.com).

Selected improvements and bug fixes

- The Cython cache on disk now uses significantly less space by deleting unnecessary source files (set the `codegen.runtime.cython.delete_source_files` preference to `False` if you want to keep these files for debugging). In addition, a warning will be given when the Cython or weave cache exceeds a configurable size (`codegen.max_cache_dir_size`). The `clear_cache` function is provided to delete files from the cache (#914).
• The C++ standalone mode now respects the `profile` option and therefore no longer collects profiling information by default. This can speed up simulations in certain cases (#935).

• The exact number of time steps that a neuron stays in the state of refractoriness after a spike could vary by up to one time step when the requested refractory time was a multiple of the simulation time step. With this fix, the number of time steps is ensured to be as expected by making use of a new `timestep` function that avoids floating point rounding issues (#949, first reported by @zhouyanasd in issue #943).

• When `restore()` was called twice for a network, spikes that were not yet delivered to their target were not restored correctly (#938, reported by @zhouyanasd).

• `SpikeGeneratorGroup` now uses a more efficient method for sorting spike indices and times, leading to a much faster preparation time for groups that store many spikes (#948).

• Fix a memory leak in `TimedArray` (#923, reported by Wilhelm Braun).

• Fix an issue with summed variables targeting subgroups (#925, reported by @AI-pha).

• Fix the use of `run_regularly` on subgroups (#922, reported by @AI-pha).

• Improve performance for `SpatialNeuron` by removing redundant computations (#910, thanks to Moritz Augustin for making us aware of the issue).

• Fix linked variables that link to scalar variables (#916)

• Fix warnings for numpy 1.14 and avoid compilation issues when switching between versions of numpy (#913)

• Fix problems when using logical operators in code generated for the numpy target which could lead to issues such as wrongly connected synapses (#901, #900).

**Backward-incompatible changes**

• No longer allow `delay` as a variable name in a synaptic model to avoid ambiguity with respect to the synaptic delay. Also no longer allow access to the `delay` variable in synaptic code since there is no way to distinguish between pre- and post-synaptic delay (#927, reported by Denis Alevi).

• Due to the changed handling of refractoriness (see bug fixes above), simulations that make use of refractoriness will possibly no longer give exactly the same results. The preference `legacy.refractory_timing` can be set to `True` to reinstate the previous behaviour.

**Infrastructure and documentation improvements**

• From this version on, conda packages will be available on conda-forge. For a limited time, we will copy over packages to the `brian-team` channel as well.

• Conda packages are no longer tied to a specific numpy version (PR #954)

• New example ([Brunel & Wang, 2001](#)) contributed by Teo Stocco and Alex Seeholzer.

**Contributions**

Github code, documentation, and issue contributions (ordered by the number of contributions):

• Marcel Stimberg (@mstimberg)

• Dan Goodman (@thesamovar)

• Teo Stocco (@zifeo)

• Dylan Muir (@DylanMuir)
1.3.7 Brian 2.1.2

This is another bug fix release that fixes a major bug in *Equations*' substitution mechanism (#896). Thanks to Teo Stocco for reporting this issue.

1.3.8 Brian 2.1.1

This is a bug fix release that re-activates parts of the caching mechanism for code generation that had been erroneously deactivated in the previous release.
1.3.9 Brian 2.1

This release introduces two main new features: a new “GSL integration” mode for differential equation that offers to integrate equations with variable-timestep methods provided by the GNU Scientific Library, and caching for the run preparation phase that can significantly speed up simulations. It also comes with a newly written tutorial, as well as additional documentation and examples.

As always, please report bugs or suggestions to the github bug tracker (https://github.com/brian-team/brian2/issues) or to the brian-development mailing list (brian-development@googlegroups.com).

New features

- New numerical integration methods with variable time-step integration, based on the GNU Scientific Library (see Numerical integration). Contributed by Charlee Fletterman, supported by 2017’s Google Summer of Code program.
- New caching mechanism for the code generation stage (application of numerical integration algorithms, analysis of equations and statements, etc.), reducing the preparation time before the actual run, in particular for simulations with multiple \texttt{run()} statements.

Selected improvements and bug fixes

- Fix a rare problem in Cython code generation caused by missing type information (#893)
- Fix warnings about improperly closed files on Python 3.6 (#892; reported and fixed by Teo Stocco)
- Fix an error when using numpify integer types for synaptic indexing (#888)
- Fix an error in numpify codegen target, triggered when assigning to a variable with an unfilled condition (#887)
- Fix an error when repeatedly referring to subexpressions in multiline statements (#880)
- Shorten long arrays in warning messages (#874)
- Enable the use of \texttt{if} in the shorthand generator syntax for \texttt{Synapses.connect} (#873)
- Fix the meaning of \texttt{i} and \texttt{j} in synapses connecting to/from other synapses (#854)

Backward-incompatible changes and deprecations

- In C++ standalone mode, information about the number of synapses and spikes will now only be displayed when built with \texttt{debug=True} (#882).
- The \texttt{linear} state updater has been renamed to \texttt{exact} to avoid confusion (#877). Users are encouraged to use \texttt{exact}, but the name \texttt{linear} is still available and does not raise any warning or error for now.
- The \texttt{independent} state updater has been marked as deprecated and might be removed in future versions.

Infrastructure and documentation improvements

- A new, more advanced, \texttt{tutorial} “about managing the slightly more complicated tasks that crop up in research problems, rather than the toy examples we’ve been looking at so far.”
- Additional documentation on \texttt{Custom events} and \texttt{Converting from integrated form to ODEs} (including example code for typical synapse models).
- New example code reproducing published findings (Platkiewicz and Brette, 2011; Stimberg et al., 2018)
• Fixes to the sphinx documentation creation process, the documentation can be downloaded as a PDF once again (705 pages!)
• Conda packages now have support for numpy 1.13 (but support for numpy 1.10 and 1.11 has been removed)

Contributions

Github code, documentation, and issue contributions (ordered by the number of contributions):
  • Marcel Stimberg (@mstimberg)
  • Charlee Fletterman (@CharleeSF)
  • Dan Goodman (@thesamovar)
  • Teo Stocco (@zifeo)
  • @k47h4

Other contributions outside of github (ordered alphabetically, apologies to anyone we forgot...):
  • Chaofei Hong
  • Lucas (“lucasedst”)

1.3.10 Brian 2.0.2.1

Fixes a bug in the tutorials’ HTML rendering on readthedocs.org (code blocks were not displayed). Thanks to Flora Bouchacourt for making us aware of this problem.

1.3.11 Brian 2.0.2

New features

• molar and liter (as well as litre, scaled versions of the former, and a few useful abbreviations such as mM) have been added as new units (#574).
• A new module brian2.units.constants provides physical constants such as the Faraday constants or the gas constant (see Constants for details).
• SpatialNeuron now supports non-linear membrane currents (e.g. Goldman–Hodgkin–Katz equations) by linearizing them with respect to v.
• Multi-compartmental models can access the capacitive current via I_c in their equations (#677)
• A new function scheduling_summary() that displays information about the scheduling of all objects (see Scheduling for details).
• Introduce a new preference to pass arguments to the make/nmake command in C++ standalone mode (devices.cpp_standalone.extra_make_args_unix for Linux/OS X and devices.cpp_standalone.extra_make_args_windows for Windows). For Linux/OS X, this enables parallel compilation by default.
• Anaconda packages for Brian 2 are now available for Python 3.6 (but Python 3.4 support has been removed).
Selected improvements and bug fixes

- Work around low performance for certain C++ standalone simulations on Linux, due to a bug in glibc (see #803). Thanks to Oleg Strikov (@xj8z) for debugging this issue and providing the workaround that is now in use.
- Make exact integration of event-driven synaptic variables use the linear numerical integration algorithm (instead of independent), fixing rare occasions where integration failed despite the equations being linear (#801).
- Better error messages for incorrect unit definitions in equations.
- Various fixes for the internal representation of physical units and the unit registration system.
- Fix a bug in the assignment of state variables in subtrees of SpatialNeuron (#822)
- Numpy target: fix an indexing error for a SpikeMonitor that records from a subgroup (#824)
- Summed variables targeting the same post-synaptic variable now raise an error (previously, only the one executed last was taken into account, see #766).
- Fix bugs in synapse generation affecting Cython (#781) respectively numpy (#835)
- C++ standalone simulations with many objects no longer fail on Windows (#787)

Backwards-incompatible changes

- celsius has been removed as a unit, because it was ambiguous in its relation to kelvin and gave wrong results when used as an absolute temperature (and not a temperature difference). For temperature differences, you can directly replace celsius by kelvin. To convert an absolute temperature in degree Celsius to Kelvin, add the zero_celsius constant from brian2.units.constants (#817).
- State variables are no longer allowed to have names ending in _pre or _post to avoid confusion with references to pre- and post-synaptic variables in Synapses (#818)

Changes to default settings

- In C++ standalone mode, the clean argument now defaults to False, meaning that make clean will not be executed by default before building the simulation. This avoids recompiling all files for unchanged simulations that are executed repeatedly. To return to the previous behaviour, specify clean=True in the device.build call (or in set_device if your script does not have an explicit device.build).

Contributions

Github code, documentation, and issue contributions (ordered by the number of contributions):
- Marcel Stimberg (@mstimberg)
- Dan Goodman (@thesamovar)
- Thomas McColgan (@phreeza)
- Daan Sprenkels (@dsprenkels)
- Romain Brette (@romainbrette)
- Oleg Strikov (@xj8z)
- Charlee Fletterman (@CharleeSF)
• Meng Dong (@whenov)
• Denis Alevi (@denisalevi)
• Mihir Vaidya (@MihirVaidya94)
• Adam (@ffa)
• Sourav Singh (@souravsingh)
• Nick Hale (@nik849)
• Cody Greer (@Cody-G)
• Jean-Sébastien Dessureault (@jsdessureault)
• Michele Giugliano (@mgiugliano)
• Teo Stocco (@zifeo)
• Edward Betts (@EdwardBetts)

Other contributions outside of github (ordered alphabetically, apologies to anyone we forgot . . .):

• Christopher Nolan
• Regimantas Jurkus
• Shailesh Appukuttan

1.3.12 Brian 2.0.1

This is a bug-fix release that fixes a number of important bugs (see below), but does not introduce any new features. We recommend all users of Brian 2 to upgrade.

As always, please report bugs or suggestions to the github bug tracker (https://github.com/brian-team/brian2/issues) or to the brian-development mailing list (brian-development@googlegroups.com).

Improvements and bug fixes

• Fix PopulationRateMonitor for recordings from subgroups (#772)
• Fix SpikeMonitor for recordings from subgroups (#777)
• Check that string expressions provided as the rates argument for PoissonGroup have correct units.
• Fix compilation errors when multiple run statements with different report arguments are used in C++ standalone mode.
• Several documentation updates and fixes

Contributions

Code and documentation contributions (ordered by the number of commits):

• Marcel Stimberg (@mstimberg)
• Dan Goodman (@thesamovar)
• Alex Seeholzer (@flinz)
• Meng Dong (@whenov)

Testing, suggestions and bug reports (ordered alphabetically, apologies to anyone we forgot . . .):
1.3.13 Brian 2.0 (changes since 1.4)

Major new features

- Much more flexible model definitions. The behaviour of all model elements can now be defined by arbitrary equations specified in standard mathematical notation.
- Code generation as standard. Behind the scenes, Brian automatically generates and compiles C++ code to simulate your model, making it much faster.
- “Standalone mode”. In this mode, Brian generates a complete C++ project tree that implements your model. This can be then be compiled and run entirely independently of Brian. This leads to both highly efficient code, as well as making it much easier to run simulations on non-standard computational hardware, for example on robotics platforms.
- Multicompartmental modelling.
- Python 2 and 3 support.

New features

- Installation should now be much easier, especially if using the Anaconda Python distribution. See Installation.
- Many improvements to Synapses which replaces the old Connection object in Brian 1. This includes: synapses that are triggered by non-spike events; synapses that target other synapses; huge speed improvements thanks to using code generation; new “generator syntax” when creating synapses is much more flexible and efficient. See Synapses.
- New model definitions allow for much more flexible refractoriness. See Refractoriness.
- SpikeMonitor and StateMonitor are now much more flexible, and cover a lot of what used to be covered by things like MultiStateMonitor, etc. See Recording during a simulation.
- Multiple event types. In addition to the default spike event, you can create arbitrary events, and have these trigger code blocks (like reset) or synaptic events. See Custom events.
- New units system allows arrays to have units. This eliminates the need for a lot of the special casing that was required in Brian 1. See Physical units.
- Indexing variable by condition, e.g. you might write G.v['x>0'] to return all values of variable v in NeuronGroup G where the group’s variable x>0. See State variables.
- “Magic” run() system has been greatly simplified and is now much more transparent. In addition, if there is any ambiguity about what the user wants to run, an error will be raised rather than making a guess. This makes it much safer. In addition, there is now a store()/restore() mechanism that simplifies restarting simulations and managing separate training/testing runs. See Running a simulation.
- Changing an external variable between runs now works as expected, i.e. something like tau=1*ms; run(100*ms); tau=5*ms; run(100*ms). In Brian 1 this would have used tau=1*ms for both runs. More generally, in Brian 2 there is now better control over namespaces. See Namespaces.
- New “shared” variables with a single value shared between all neurons. See Shared variables.
• New `Group.run_regularly` method for a codegen-compatible way of doing things that used to be done with `network_operation()` (which can still be used). See `Regular operations`.

• New system for handling externally defined functions. They have to specify which units they accept in their arguments, and what they return. In addition, you can easily specify the implementation of user-defined functions in different languages for code generation. See `Functions`.

• State variables can now be defined as integer or boolean values. See `Equations`.

• State variables can now be exported directly to Pandas data frame. See `Storing state variables`.

• New generalised “flags” system for giving additional information when defining models. See `Flags`.

• `TimedArray` now allows for 2D arrays with arbitrary indexing. See `Timed arrays`.

• Better support for using Brian in IPython/Jupyter. See, for example, `start_scope()`.

• New preferences system. See `Preferences`.

• Random number generation can now be made reliably reproducible. See `Random numbers`.

• New profiling option to see which parts of your simulation are taking the longest to run. See `Profiling`.

• New logging system allows for more precise control. See `Logging`.

• New ways of importing Brian for advanced Python users. See `Importing Brian`.

• Improved control over the order in which objects are updated during a run. See `Custom progress reporting`.

• Users can now easily define their own numerical integration methods. See `State update`.

• Support for parallel processing using the OpenMP version of standalone mode. Note that all Brian tests pass with this, but it is still considered to be experimental. See `Multi-threading with OpenMP`.

### Backwards incompatible changes

See `Detailed Brian 1 to Brian 2 conversion notes`.

### Behind the scenes changes

• All user models are now passed through the code generation system. This allows us to be much more flexible about introducing new target languages for generated code to make use of non-standard computational hardware. See `Code generation`.

• New standalone/device mode allows generation of a complete project tree that can be compiled and built independently of Brian and Python. This allows for even more flexible use of Brian on non-standard hardware. See `Devices`.

• All objects now have a unique name, used in code generation. This can also be used to access the object through the `Network` object.

### Contributions

Full list of all Brian 2 contributors, ordered by the time of their first contribution:

• Dan Goodman (@thesamovar)

• Marcel Stimberg (@mstimberg)

• Romain Brette (@romainbrette)

• Cyrille Rossant (@rossant)
1.3.14 Brian 2.0 (changes since 2.0rc3)

New features

- A new flag `constant over dt` can be applied to subexpressions to have them only evaluated once per timestep (see *Models and neuron groups*). This flag is mandatory for stateful subexpressions, e.g. expressions using `rand()` or `randn` (#720, #721)

Improvements and bug fixes

- Fix `EventMonitor.values` and `SpikeMonitor.spike_trains` to always return sorted spike/event times (#725).
- Respect the `active` attribute in C++ standalone mode (#718).
- More consistent check of compatible time and dt values (#730).
- Attempting to set a synaptic variable or to start a simulation with synapses without any preceding connect call now raises an error (#737).
- Improve the performance of coordinate calculation for *Morphology* objects, which previously made plotting very slow for complex morphologies (#741).
- Fix a bug in *SpatialNeuron* where it did not detect non-linear dependencies on v, introduced via point currents (#743).

Infrastructure and documentation improvements

- An interactive demo, tutorials, and examples can now be run in an interactive jupyter notebook on the mybinder platform, without any need for a local Brian installation (#736). Thanks to Ben Evans for the idea and help with the implementation.
- A new extensive guide for converting Brian 1 simulations to Brian 2 user coming from Brian 1: Changes for Brian 1 users
- A re-organized *User’s guide*, with clearer indications which information is important for new Brian users.
Contributions

Code and documentation contributions (ordered by the number of commits):

- Marcel Stimberg (@mstimberg)
- Dan Goodman (@thesamovar)
- Benjamin Evans (@bdevans)

Testing, suggestions and bug reports (ordered alphabetically, apologies to anyone we forgot...):

- Chaofei Hong
- Daniel Bliss
- Jacopo Bono
- Ruben Tikidji-Hamburyan

1.3.15 Brian 2.0rc3

This is another “release candidate” for Brian 2.0 that fixes a range of bugs and introduces better support for random numbers (see below). We are getting close to the final Brian 2.0 release, the remaining work will focus on bug fixes, and better error messages and documentation.

As always, please report bugs or suggestions to the github bug tracker (https://github.com/brian-team/brian2/issues) or to the brian-development mailing list (brian-development@googlegroups.com).

New features

- Brian now comes with its own `seed()` function, allowing to seed the random number generator and thereby to make simulations reproducible. This function works for all code generation targets and in runtime and standalone mode. See Random numbers for details.

- Brian can now export/import state variables of a group or a full network to/from a pandas DataFrame and comes with a mechanism to extend this to other formats. Thanks to Dominik Krzeminski for this contribution (see #306).

Improvements and bug fixes

- Use a Mersenne-Twister pseudorandom number generator in C++ standalone mode, replacing the previously used low-quality random number generator from the C standard library (see #222, #671 and #706).

- Fix a memory leak in code running with the weave code generation target, and a smaller memory leak related to units stored repetitively in the UnitRegistry.

- Fix a difference of one timestep in the number of simulated timesteps between runtime and standalone that could arise for very specific values of dt and t (see #695).

- Fix standalone compilation failures with the most recent gcc version which defaults to C++14 mode (see #701)

- Fix incorrect summation in synapses when using the `summed` flag and writing to pre-synaptic variables (see #704).

- Make synaptic pathways work when connecting groups that define nested subexpressions, instead of failing with a cryptic error message (see #707).
Contributions

Code and documentation contributions (ordered by the number of commits):

- Marcel Stimberg (@mstimberg)
- Dominik Krzemiński (@dokato)
- Dan Goodman (@thesamovar)
- Martino Sorbaro (@martinosorb)

Testing, suggestions and bug reports (ordered alphabetically, apologies to anyone we forgot...):

- Craig Henriquez
- Daniel Bliss
- David Higgins
- Gordon Erlebacher
- Max Gillett
- Moritz Augustin
- Sami Abdul-Wahid

1.3.16 Brian 2.0rc1

This is a bug fix release that we release only about two weeks after the previous release because that release introduced a bug that could lead to wrong integration of stochastic differential equations. Note that standard neuronal noise models were not affected by this bug, it only concerned differential equations implementing a “random walk”. The release also fixes a few other issues reported by users, see below for more information.

Improvements and bug fixes

- Fix a regression from 2.0b4: stochastic differential equations without any non-stochastic part (e.g. $dx/dt = xi/sqrt(ms)$) were not integrated correctly (see #686).
- Repeatedly calling `restore()` (or `Network.restore`) no longer raises an error (see #681).
- Fix an issue that made `PoissonInput` refuse to run after a change of dt (see #684).
- If the `rates` argument of `PoissonGroup` is a string, it will now be evaluated at every time step instead of once at construction time. This makes time-dependent rate expressions work as expected (see #660).

Contributions

Code and documentation contributions (ordered by the number of commits):

- Marcel Stimberg (@mstimberg)

Testing, suggestions and bug reports (ordered alphabetically, apologies to anyone we forgot...):

- Cian O’Donnell
- Daniel Bliss
- Ibrahim Ozturk
- Olivia Gozel
1.3.17 Brian 2.0rc

This is a release candidate for the final Brian 2.0 release, meaning that from now on we will focus on bug fixes and documentation, without introducing new major features or changing the syntax for the user. This release candidate itself does however change a few important syntax elements, see “Backwards-incompatible changes” below.

As always, please report bugs or suggestions to the github bug tracker (https://github.com/brian-team/brian2/issues) or to the brian-development mailing list (brian-development@googlegroups.com).

Major new features

- New “generator syntax” to efficiently generate synapses (e.g. one-to-one connections), see Creating synapses for more details.
- For synaptic connections with multiple synapses between a pair of neurons, the number of the synapse can now be stored in a variable, allowing its use in expressions and statements (see Creating synapses).
- Synapses can now target other Synapses objects, useful for some models of synaptic modulation.
- The Morphology object has been completely re-worked and several issues have been fixed. The new Section object allows to model a section as a series of truncated cones (see Creating a neuron morphology).
- Scripts with a single run() call, no longer need an explicit device.build() call to run with the C++ standalone device. A set_device() in the beginning is enough and will trigger the build call after the run (see Standalone code generation).
- All state variables within a Network can now be accessed by Network.get_states and Network.set_states and the store()/restore() mechanism can now store the full state of a simulation to disk.
- Stochastic differential equations with multiplicative noise can now be integrated using the Euler-Heun method (heun). Thanks to Jan-Hendrik Schleimer for this contribution.
- Error messages have been significantly improved: errors for unit mismatches are now much clearer and error messages triggered during the initialization phase point back to the line of code where the relevant object (e.g. a NeuronGroup) was created.
- PopulationRateMonitor now provides a smooth_rate method for a filtered version of the stored rates.

Improvements and bug fixes

- In addition to the new synapse creation syntax, sparse probabilistic connections are now created much faster.
- The time for the initialization phase at the beginning of a run() has been significantly reduced.
- Multicompartmental simulations with a large number of compartments are now simulated more efficiently and are making better use of several processor cores when OpenMP is activated in C++ standalone mode. Thanks to Moritz Augustin for this contribution.
- Simulations will use compiler settings that optimize performance by default.
- Objects that have user-specified names are better supported for complex simulation scenarios (names no longer have to be unique at all times, but only across a network or across a standalone device).
- Various fixes for compatibility with recent versions of numpy and sympy.
Important backwards-incompatible changes

- The argument names in `Synapses.connect` have changed and the first argument can no longer be an array of indices. To connect based on indices, use `Synapses.connect(i=source_indices, j=target_indices)`. See *Creating synapses* and the documentation of `Synapses.connect` for more details.

- The actions triggered by pre-synaptic and post-synaptic spikes are now described by the `on_pre` and `on_post` keyword arguments (instead of `pre` and `post`).

- The `Morphology` object no longer allows to change attributes such as length and diameter after its creation. Complex morphologies should instead be created using the `Section` class, allowing for the specification of all details.

- `Morphology` objects that are defined with coordinates need to provide the start point (relative to the end point of the parent compartment) as the first coordinate. See *Creating a neuron morphology* for more details.

- For simulations using the C++ standalone mode, no longer call `Device.build` (if using a single `run()` call), or use `set_device()` with `build_on_run=False` (see *Standalone code generation*).

Infrastructure improvements

- Our test suite is now also run on Mac OS-X (on the Travis CI platform).

Contributions

Code and documentation contributions (ordered by the number of commits):

- Marcel Stimberg (@mstimberg)
- Dan Goodman (@thesamovar)
- Moritz Augustin (@moritzaugustin)
- Jan-Hendrik Schleimer (@ttxtea)
- Romain Cazé (@rcaze)
- Konrad Wartke (@Kwartke)
- Romain Brette (@romainbrette)

Testing, suggestions and bug reports (ordered alphabetically, apologies to anyone we forgot...):

- Chaofei Hong
- Kees de Leeuw
- Luke Y Prince
- Myung Seok Shim
- Owen Mackwood
- Github users: @epaxon, @flinz, @mariomulansky, @martinosorb, @neuralyzer, @oleskiw, @prcastro, @sudoankit
1.3.18 Brian 2.0b4

This is the fourth (and probably last) beta release for Brian 2.0. This release adds a few important new features and fixes a number of bugs so we recommend all users of Brian 2 to upgrade. If you are a user new to Brian, we also recommend to directly start with Brian 2 instead of using the stable release of Brian 1. Note that the new recommended way to install Brian 2 is to use the Anaconda distribution and to install the Brian 2 conda package (see Installation). This is however still a Beta release, please report bugs or suggestions to the github bug tracker (https://github.com/brian-team/brian2/issues) or to the brian-development mailing list (brian-development@googlegroups.com).

Major new features

• In addition to the standard threshold/reset, groups can now define “custom events”. These can be recorded with the new EventMonitor (a generalization of SpikeMonitor) and Synapses can connect to these events instead of the standard spike event. See Custom events for more details.

• SpikeMonitor and EventMonitor can now also record state variable values at the time of spikes (or custom events), thereby offering the functionality of StateSpikeMonitor from Brian 1. See Recording variables at spike time for more details.

• The code generation modes that interact with C++ code (weave, Cython, and C++ standalone) can now be more easily configured to work with external libraries (compiler and linker options, header files, etc.). See the documentation of the cpp_prefs module for more details.

Improvements and bug fixes

• Cython simulations no longer interfere with each other when run in parallel (thanks to Daniel Bliss for reporting and fixing this).

• The C++ standalone now works with scalar delays and the spike queue implementation deals more efficiently with them in general.

• Dynamic arrays are now resized more efficiently, leading to faster monitors in runtime mode.

• The spikes generated by a SpikeGeneratorGroup can now be changed between runs using the set_spikes method.

• Multi-step state updaters now work correctly for non-autonomous differential equations.

• PoissonInput now correctly works with multiple clocks (thanks to Daniel Bliss for reporting and fixing this).

• The get_states method now works for StateMonitor. This method provides a convenient way to access all the data stored in the monitor, e.g. in order to store it on disk.

• C++ compilation is now easier to get to work under Windows, see Installation for details.

Important backwards-incompatible changes

• The custom_operation method has been renamed to run_regularly and can now be called without the need for storing its return value.

• StateMonitor will now by default record at the beginning of a time step instead of at the end. See Recording variables continuously for details.

• Scalar quantities now behave as python scalars with respect to in-place modifications (augmented assignments). This means that \( x = 3\text{ mV}; \ y = x; \ y += 1\text{ mV} \) will no longer increase the value of the variable \( x \) as well.
Infrastructure improvements

- We now provide conda packages for Brian 2, making it very easy to install when using the Anaconda distribution (see Installation).

Contributions

Code and documentation contributions (ordered by the number of commits):

- Marcel Stimberg (@mstimberg)
- Dan Goodman (@thesamovar)
- Daniel Bliss (@dabliss)
- Romain Brette (@romainbrette)

Testing, suggestions and bug reports (ordered alphabetically, apologies to everyone we forgot...):

- Daniel Bliss
- Damien Drix
- Rainer Engelken
- Beatriz Herrera Figueredo
- Owen Mackwood
- Augustine Tan
- Ot de Wiljes

1.3.19 Brian 2.0b3

This is the third beta release for Brian 2.0. This release does not add many new features but it fixes a number of important bugs so we recommend all users of Brian 2 to upgrade. If you are a user new to Brian, we also recommend to directly start with Brian 2 instead of using the stable release of Brian 1.

This is however still a Beta release, please report bugs or suggestions to the github bug tracker (https://github.com/brian-team/brian2/issues) or to the brian-development mailing list (brian-development@googlegroups.com).

Major new features

- A new PoissonInput class for efficient simulation of Poisson-distributed input events.

Improvements

- The order of execution for pre and post statements happening in the same time step was not well defined (it fell back to the default alphabetical ordering, executing post before pre). It now explicitly specifies the order attribute so that pre gets executed before post (as in Brian 1). See the Synapses documentation for details.
- The default schedule that is used can now be set via a preference (core.network.default_schedule). New automatically generated scheduling slots relative to the explicitly defined ones can be used, e.g. before_resets or after_synapses. See Scheduling for details.
• The `scipy` package is no longer a dependency (note that `weave` for compiled C code under Python 2 is now available in a separate package). Note that multicompartmental models will still benefit from the `scipy` package if they are simulated in pure Python (i.e. with the `numpy` code generation target) – otherwise Brian 2 will fall back to a numpy-only solution which is significantly slower.

**Important bug fixes**

• Fix `SpikeGeneratorGroup` which did not emit all the spikes under certain conditions for some code generation targets (#429)

• Fix an incorrect update of pre-synaptic variables in synaptic statements for the `numpy` code generation target (#435).

• Fix the possibility of an incorrect memory access when recording a subgroup with `SpikeMonitor` (#454).

• Fix the storing of results on disk for C++ standalone on Windows – variables that had the same name when ignoring case (e.g. `i` and `I`) where overwriting each other (#455).

**Infrastructure improvements**

• Brian 2 now has a chat room on gitter: https://gitter.im/brian-team/brian2

• The sphinx documentation can now be built from the release archive file

• After a big cleanup, all files in the repository have now simple LF line endings (see https://help.github.com/articles/dealing-with-line-endings/ on how to configure your own machine properly if you want to contribute to Brian).

**Contributions**

Code and documentation contributions (ordered by the number of commits):

• Marcel Stimberg (@mstimberg)

• Dan Goodman (@thesamovar)

• Konrad Wartke (@kwartke)

Testing, suggestions and bug reports (ordered alphabetically, apologies to everyone we forgot . . . ):

• Daniel Bliss

• Owen Mackwood

• Ankur Sinha

• Richard Tomsett

**1.3.20 Brian 2.0b2**

This is the second beta release for Brian 2.0, we recommend all users of Brian 2 to upgrade. If you are a user new to Brian, we also recommend to directly start with Brian 2 instead of using the stable release of Brian 1.

This is however still a Beta release, please report bugs or suggestions to the github bug tracker (https://github.com/brian-team/brian2/issues) or to the brian-development mailing list (brian-development@googlegroups.com).
Major new features

- Multi-compartmental simulations can now be run using the *Standalone code generation* mode (this is not yet well-tested, though).

- The implementation of *TimedArray* now supports two-dimensional arrays, i.e. different input per neuron (or synapse, etc.), see *Timed arrays* for details.

- Previously, not setting a code generation target (using the *codegen.target* preference) would mean that the *numpy* target was used. Now, the default target is *auto*, which means that a compiled language (*weave* or *cython*) will be used if possible. See *Computational methods and efficiency* for details.

- The implementation of *SpikeGeneratorGroup* has been improved and it now supports a *period* argument to repeatedly generate a spike pattern.

Improvements

- The selection of a numerical algorithm (if none has been specified by the user) has been simplified. See *Numerical integration* for details.

- Expressions that are shared among neurons/synapses are now updated only once instead of for every neuron/synapse which can lead to performance improvements.

- On Windows, The Microsoft Visual C compiler is now supported in the *cpp_standalone* mode, see the respective notes in the *Installation* and *Computational methods and efficiency* documents.

- Simulation runs (using the standard “runtime” device) now collect profiling information. See *Profiling* for details.

Infrastructure and documentation improvements

- *Tutorials for beginners* in the form of ipython notebooks (currently only covering the basics of neurons and synapses) are now available.

- The *Examples* in the documentation now include the images they generated. Several examples have been adapted from Brian 1.

- The code is now automatically tested on Windows machines, using the *appveyor* service. This complements the Linux testing on *travis*.

- Using a version of a dependency (e.g. *sympy*) that we don’t support will now raise an error when you import *brian2* – see *Dependency checks* for more details.

- Test coverage for the *cpp_standalone* mode has been significantly increased.

Important bug fixes

- The preparation time for complicated equations has been significantly reduced.

- The string representation of small physical quantities has been corrected (#361)

- Linking variables from a group of size 1 now works correctly (#383)
Contributions

Code and documentation contributions (ordered by the number of commits):

- Marcel Stimberg (@mstimberg)
- Dan Goodman (@thesamovar)
- Romain Brette (@romainbrette)
- Pierre Yger (@yger)

Testing, suggestions and bug reports (ordered alphabetically, apologies to everyone we forgot…):

- Conor Cox
- Gordon Erlebacher
- Konstantin Mergenthaler

1.3.21 Brian 2.0beta

This is the first beta release for Brian 2.0 and the first version of Brian 2.0 we recommend for general use. From now on, we will try to keep changes that break existing code to a minimum. If you are a user new to Brian, we’d recommend to start with the Brian 2 beta instead of using the stable release of Brian 1.

This is however still a Beta release, please report bugs or suggestions to the github bug tracker (https://github.com/brian-team/brian2/issues) or to the brian-development mailing list (brian-development@googlegroups.com).

Major new features

- New classes Morphology and SpatialNeuron for the simulation of Multicompartment models
- A temporary “bridge” for brian.hears that allows to use its Brian 1 version from Brian 2 (Brian Hears)
- Cython is now a new code generation target, therefore the performance benefits of compiled code are now also available to users running simulations under Python 3.x (where scipy.weave is not available)
- Networks can now store their current state and return to it at a later time, e.g. for simulating multiple trials starting from a fixed network state (Continuing/repeating simulations)
- C++ standalone mode: multiple processors are now supported via OpenMP (Multi-threading with OpenMP), although this code has not yet been well tested so may be inaccurate.
- C++ standalone mode: after a run, state variables and monitored values can be loaded from disk transparently. Most scripts therefore only need two additional lines to use standalone mode instead of Brian’s default runtime mode (Standalone code generation).

Syntax changes

- The syntax and semantics of everything around simulation time steps, clocks, and multiple runs have been cleaned up, making reinit obsolete and also making it unnecessary for most users to explicitly generate Clock objects – instead, a dt keyword can be specified for objects such as NeuronGroup (Running a simulation)
- The scalar flag for parameters/subexpressions has been renamed to shared
- The “unit” for boolean variables has been renamed from bool to boolean
- C++ standalone: several keywords of CPPStandaloneDevice.build have been renamed

1.3. Release notes
• The preferences are now accessible via `prefs` instead of `brian_prefs`
• The runner method has been renamed to `custom_operation`

**Improvements**

• Variables can now be linked across `NeuronGroups` ([Linked variables](#))
• More flexible progress reporting system, progress reporting also works in the C++ standalone mode ([Progress reporting](#))
• State variables can be declared as `integer` ([Equation strings](#))

**Bug fixes**

57 github issues have been closed since the alpha release, of which 26 had been labeled as bugs. We recommend all users of Brian 2 to upgrade.

**Contributions**

Code and documentation contributions (ordered by the number of commits):

• Marcel Stimberg (@mstimberg)
• Dan Goodman (@thesamovar)
• Romain Brette (@romainbrette)
• Pierre Yger (@yger)
• Werner Beroux (@wernight)

Testing, suggestions and bug reports (ordered alphabetically, apologies to everyone we forgot…):

• Guillaume Bellec
• Victor Benichoux
• Laureline Logiaco
• Konstantin Mergenthaler
• Maurizio De Pitta
• Jan-Hendrick Schleimer
• Douglas Sterling
• Katharina Wilmes

## 1.4 Changes for Brian 1 users

- *Physical units*
- *Unported packages*
- *Replacement packages*
In most cases, Brian 2 works in a very similar way to Brian 1 but there are some important differences to be aware of. The major distinction is that in Brian 2 you need to be more explicit about the definition of your simulation in order to avoid inadvertent errors. In some cases, you will now get a warning in other even an error – often the error/warning message describes a way to resolve the issue.

Specific examples how to convert code from Brian 1 can be found in the document Detailed Brian 1 to Brian 2 conversion notes.

### 1.4.1 Physical units

The unit system now extends to arrays, e.g. `np.arange(5) * mV` will retain the units of volts and not discard them as Brian 1 did. Brian 2 is therefore also more strict in checking the units. For example, if the state variable `v` uses the unit of volt, the statement `G.v = np.random(len(G)) / 1000.` will now raise an error. For consistency, units are returned everywhere, e.g. in monitors. If `mon` records a state variable `v`, `mon.t` will return a time in seconds and `mon.v` the stored values of `v` in units of volts.

If you need a pure numpy array without units for further processing, there are several options: if it is a state variable or a recorded variable in a monitor, appending an underscore will refer to the variable values without units, e.g. `mon.t_` returns pure floating point values. Alternatively, you can remove units by diving by the unit (e.g. `mon.t / second`) or by explicitly converting it (`np.asarray(mon.t)`).

Here’s an overview showing a few expressions and their respective values in Brian 1 and Brian 2:

<table>
<thead>
<tr>
<th>Expression</th>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>1 * mV</code></td>
<td><code>1.0 * mvolt</code></td>
<td><code>1.0 * mvolt</code></td>
</tr>
<tr>
<td><code>np.array(1) * mV</code></td>
<td><code>0.001</code></td>
<td><code>1.0 * mvolt</code></td>
</tr>
<tr>
<td><code>np.array([1]) * mV</code></td>
<td><code>array([0.001])</code></td>
<td><code>array([1.0]) * mvolt</code></td>
</tr>
<tr>
<td><code>np.mean(np.arange(5) * mV)</code></td>
<td><code>0.002</code></td>
<td><code>2.0 * mvolt</code></td>
</tr>
<tr>
<td><code>(np.arange(2) * mV) &gt;= 1 * mV</code></td>
<td><code>array([False, True], dtype=bool)</code></td>
<td><code>array([False, True], dtype=bool)</code></td>
</tr>
<tr>
<td><code>(np.arange(2) * mV)[0] &gt;= 1 * mV</code></td>
<td><code>False</code></td>
<td><code>False</code></td>
</tr>
<tr>
<td><code>(np.arange(2) * mV)[1] &gt;= 1 * mV</code></td>
<td><code>DimensionMismatchError</code></td>
<td><code>True</code></td>
</tr>
</tbody>
</table>

### 1.4.2 Unported packages

The following packages have not (yet) been ported to Brian 2. If your simulation critically depends on them, you should consider staying with Brian 1 for now.

- `brian.tools`
- `brian.library.modelfitting`
- `brian.library.electrophysiology`

### 1.4.3 Replacement packages

The following packages that were included in Brian 1 have now been split into separate packages.

- `brian.hears` has been updated to `brian2hears`. Note that there is a legacy package `brian2.hears` included in `brian2`, but this is now deprecated and will be removed in a future release. For now, see `Brian Hears` for details.
1.4.4 Removed classes/functions and their replacements

In Brian 2, we have tried to keep the number of classes/functions to a minimum, but make each of them flexible enough to encompass a large number of use cases. A lot of the classes and functions that existed in Brian 1 have therefore been removed. The following table lists (most of) the classes that existed in Brian 1 but do no longer exist in Brian 2. You can consult it when you get a NameError while converting an existing script from Brian 1. The third column links to a document with further explanation and the second column gives either:

1. the equivalent class in Brian 2 (e.g. StateMonitor can record multiple variables now and therefore replaces MultiStateMonitor);

2. the name of a Brian 2 class in square brackets (e.g. [Synapses] for STDP), this means that the class can be used as a replacement but needs some additional code (e.g. explicitly specified STDP equations). The “More details” document should help you in making the necessary changes;

3. “string expression”, if the functionality of a previously existing class can be expressed using the general string expression framework (e.g. threshold=VariableThreshold('Vt', 'V') can be replaced by threshold='V > Vt');

4. a link to the relevant github issue if no equivalent class/function does exist so far in Brian 2;

5. a remark such as “obsolete” if the particular class/function is no longer needed.

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
<th>More details</th>
</tr>
</thead>
<tbody>
<tr>
<td>AdEx</td>
<td>[Equations]</td>
<td>Library models (Brian 1 -&gt; 2 conversion)</td>
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<tr>
<td>aEIF</td>
<td>[Equations]</td>
<td>Library models (Brian 1 -&gt; 2 conversion)</td>
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<td>AERSpikeMonitor</td>
<td>#298</td>
<td>Monitors (Brian 1 -&gt; 2 conversion)</td>
</tr>
<tr>
<td>alpha_conductance</td>
<td>[Equations]</td>
<td>Library models (Brian 1 -&gt; 2 conversion)</td>
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<tr>
<td>alpha_current</td>
<td>[Equations]</td>
<td>Library models (Brian 1 -&gt; 2 conversion)</td>
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<tr>
<td>alpha_synapse</td>
<td>[Equations]</td>
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<td>[SpikeMonitor]</td>
<td>Monitors (Brian 1 -&gt; 2 conversion)</td>
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<td>#443</td>
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### Table 1 – continued from previous page

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<tr>
<td>VariableThreshold</td>
<td>string expression</td>
<td>Neural models (Brian 1 \rightarrow 2 conversion)</td>
</tr>
</tbody>
</table>

### List of detailed instructions

#### Detailed Brian 1 to Brian 2 conversion notes

These documents are only relevant for former users of Brian 1. If you do not have any Brian 1 code to convert, go directly to the main *User’s guide.*
Neural models (Brian 1 –> 2 conversion)

The syntax for specifying neuron models in a NeuronGroup changed in several details. In general, a string-based syntax (that was already optional in Brian 1) consistently replaces the use of classes (e.g. VariableThreshold) or guessing (e.g. which variable does threshold=50*mV check).

### Threshold and Reset

String-based thresholds are now the only possible option and replace all the methods of defining threshold/reset in Brian 1:

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>threshold=-50<em>mV, reset=-70</em>mV)</td>
<td>threshold='v &gt; -50<em>mV →', reset='v = -70</em>mV')</td>
</tr>
<tr>
<td>threshold=Threshold(-50<em>mV, state='v'), reset=Reset(-70</em>mV, state='w')</td>
<td>threshold='v &gt; -50<em>mV →', reset='v = -70</em>mV')</td>
</tr>
<tr>
<td>threshold=VariableThreshold(state='v', threshold_state='vt'),</td>
<td>threshold='v &gt; vt', reset='v = vr')</td>
</tr>
<tr>
<td>reset=VariableThreshold(state='v', resetvaluestate='vr'))</td>
<td></td>
</tr>
</tbody>
</table>
### Table 2 – continued from previous page

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>`group = NeuronGroup(N, 'rate : Hz',</td>
<td>`group = NeuronGroup(N, 'rate : Hz',</td>
</tr>
<tr>
<td>threshold=PoissonThreshold(state='rate ')</td>
<td>threshold='rand()&lt;rate*dt')</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>There’s no direct equivalent for the “functional threshold/reset”</td>
<td></td>
</tr>
<tr>
<td>mechanism from Brian 1. In simple cases, they can be</td>
<td></td>
</tr>
<tr>
<td>implemented using the general string expression/statement mechanism</td>
<td></td>
</tr>
<tr>
<td>(note that in Brian 1, <code>reset=myreset</code> is equivalent to <code>reset=FunReset (myreset)</code>):</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>P.v_[spikes] = -</td>
<td>threshold='v &gt; -50*mV</td>
</tr>
<tr>
<td>70*mV+rand(len(spikes))<em>5</em>mV</td>
<td>reset='-70*mV +rand()<em>5</em>mV')</td>
</tr>
<tr>
<td>`group = NeuronGroup(N, 'dv/dt = -v / tau : volt',</td>
<td></td>
</tr>
<tr>
<td>threshold=-50*mV,</td>
<td></td>
</tr>
<tr>
<td>reset=myreset)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>return (v &gt; -50*mV) &amp; (rand(N) &gt; 0.5)</td>
<td>threshold='v &gt; -50*mV</td>
</tr>
<tr>
<td></td>
<td>reset='v = -70*mV')</td>
</tr>
<tr>
<td>`group = NeuronGroup(N, 'dv/dt = -v / tau : volt',</td>
<td></td>
</tr>
<tr>
<td>threshold=SimpleFunThreshold(mythreshold,</td>
<td></td>
</tr>
<tr>
<td>state='v'),</td>
<td></td>
</tr>
<tr>
<td>reset=-70*mV)</td>
<td></td>
</tr>
</tbody>
</table>

For more complicated cases, you can use the general mechanism for *User-provided functions* that Brian 2 provides. The only caveat is that you’d have to provide an implementation of the function in the code generation target language which is by default C++ or Cython. However, in the default *Runtime code generation* mode, you can choose different code generation targets for different parts of your simulation. You can thus switch the code generation target for the threshold/reset mechanism to *numpy* while leaving the default target for the rest of the simulation in place. The details of this process and the correct definition of the functions (e.g. `global_reset` needs a “dummy” return value) are somewhat cumbersome at the moment and we plan to make them more straightforward in the future. Also note that if you use this kind of mechanism extensively, you’ll lose all the performance advantage that Brian 2’s code generation mechanism provides (in addition to not being able to use *Standalone code generation* mode at all).
**Brian 1**

```python
def single_threshold(v):
    # Only let a single neuron spike
    crossed_threshold = np.nonzero(v > -50*mV)[0]
    should_spike = np.zeros(len(P), dtype=np.bool)
    if len(crossed_threshold):
        choose = np.random.randint(len(crossed_threshold))
        should_spike[crossed_threshold[choose]] = True
    return should_spike
def global_reset(P, spikes):
    # Reset everything
    if len(spikes):
        P.v_[:] = -70*mV
neurons = NeuronGroup(N, 'dv/dt = -(v - threshold)/tau : volt',
                      threshold=SimpleFunThreshold(single_threshold, state='v'),
                      reset=global_reset)
```

**Brian 2**

```python
@check_units(v=volt, result=bool)
def single_threshold(v):
    pass  # ... (identical to Brian 1)

@check_units(spikes=1, result=1)
def global_reset(spikes):
    # Reset everything
    if len(spikes):
        neurons.v_[:] = -0.07
    neurons = NeuronGroup(N, 'dv/dt = -(v - threshold)/tau : volt',
                       threshold='single_threshold(v)',
                       reset='dummy = global_reset(i)')
```

For an example how to translate `EmpiricalThreshold`, see the section on “Refractoriness” below.

### Refractoriness

For a detailed description of Brian 2’s refractoriness mechanism see Refractoriness.

In Brian 1, refractoriness was tightly linked with the reset mechanism and some combinations of refractoriness and reset were not allowed. The standard refractory mechanism had two effects during the refractoriness: it prevented the refractory cell from spiking and it clamped a state variable (normally the membrane potential of the cell). In Brian 2, refractoriness is independent of reset and the two effects are specified separately: the `refractory` keyword specifies the time (or an expression evaluating to a time) during which the cell does not spike, and the *(unless refractory)* flag marks one or more variables to be clamped during the refractory period. To correctly translate the standard refractory mechanism from Brian 1, you’ll therefore need to specify both:

**Brian 1**

```python
group = NeuronGroup(N, 'dv/dt = (I - v)/tau : volt',
                    threshold=-50*mV, reset=-70*mV, refractory=3*ms)
```

**Brian 2**

```python
group = NeuronGroup(N, 'dv/dt = (I - v)/tau : volt (unless refractory)',
                    threshold='v > -50*mV', reset='v = -70*mV', refractory=3*ms)
```
More complex refractoriness mechanisms based on SimpleCustomRefractoriness and CustomRefractoriness can be translated using string expressions or user-defined functions, see the remarks in the preceding section on “Threshold and Reset”.

Brian 2 no longer has an equivalent to the EmpiricalThreshold class (which detects at the first threshold crossing but ignores all following threshold crossings for a certain time after that). However, the standard refractoriness mechanism can be used to implement the same behaviour, since it does not reset/clamp any value if not explicitly asked for it (which would be fatal for Hodgkin-Huxley type models):

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>group = NeuronGroup(N,'''</td>
<td>group = NeuronGroup(N,'''</td>
</tr>
<tr>
<td>dv/dt = (I_L - I_Na -</td>
<td>dv/dt = (I_L - I_Na -</td>
</tr>
<tr>
<td>\rightarrow I_K + I)/Cm : volt</td>
<td>\rightarrow I_K + I)/Cm : volt</td>
</tr>
<tr>
<td>...'''</td>
<td>...'''</td>
</tr>
<tr>
<td>threshold=EmpiricalThreshold(threshold=20*mV,</td>
<td>threshold='v &gt; -20*mV</td>
</tr>
<tr>
<td>\rightarrow</td>
<td>\rightarrow</td>
</tr>
<tr>
<td>\rightarrow refractory=1*ms,</td>
<td>refractory=1*ms)</td>
</tr>
<tr>
<td>\rightarrow state='v')})</td>
<td></td>
</tr>
</tbody>
</table>

**Subgroups**

The class NeuronGroup in Brian 2 does no longer provide a subgroup method, the only way to construct subgroups is therefore the slicing syntax (that works in the same way as in Brian 1):

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>group = NeuronGroup(4000, ...)</td>
<td>group = NeuronGroup(4000, ...)</td>
</tr>
<tr>
<td>group_exc = group.subgroup(3200)</td>
<td>group_exc = group[:3200]</td>
</tr>
<tr>
<td>group_inh = group.subgroup(800)</td>
<td>group_inh = group[3200:]</td>
</tr>
</tbody>
</table>

**Linked Variables**

For a description of Brian 2’s mechanism to link variables between groups, see *Linked variables*.

Linked variables need to be explicitly annotated with the (linked) flag in Brian 2:
**Synapses (Brian 1 → 2 conversion)**

In Brian 2, the `Synapses` class is the only class to model synaptic connections, you will therefore have to convert all uses of Brian 1’s `Connection` class. The `Connection` class increases a post-synaptic variable by a certain amount (the “synaptic weight”) each time a pre-synaptic spike arrives. This has to be explicitly specified when using the `Synapses` class, the equivalent to the basic `Connection` usage is:

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>conn = Connection(source, target, 'ge')</td>
<td>conn = Synapses(source, target, 'w :</td>
</tr>
<tr>
<td></td>
<td>siemens', on_pre='ge += w')</td>
</tr>
</tbody>
</table>

Note that the variable `w`, which stores the synaptic weight, has to have the same units as the post-synaptic variable (in this case: `ge`) that it increases.

**Creating synapses and setting weights**

With the `Connection` class, creating a synapse and setting its weight is a single process whereas with the `Synapses` class those two steps are separate. There is no direct equivalent to the convenience functions `connect_full`, `connect_random` and `connect_one_to_one`, but you can easily implement the same functionality with the general mechanism of `Synapses.connect`:
### Brian 1

<table>
<thead>
<tr>
<th>conn1 = Connection(source, target, 'ge')</th>
<th>conn1[3, 5] = 3*nS</th>
</tr>
</thead>
<tbody>
<tr>
<td>conn1[3, 5] = 3*nS</td>
<td>conn1 = Synapses(source, target, 'w : siemens',</td>
</tr>
<tr>
<td></td>
<td>on_pre='ge += w')</td>
</tr>
<tr>
<td>conn1.connect(i=3, j=5)</td>
<td>conn1.w[3, 5] = 3<em>nS  # (or conn1.w = 3</em>nS)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>conn2 = Connection(source, target, 'ge')</th>
<th>conn2 = ... # see above</th>
</tr>
</thead>
<tbody>
<tr>
<td>conn2.connect_full(source, target, 5*nS)</td>
<td>conn2.connect()</td>
</tr>
<tr>
<td>conn2.w = 5*nS</td>
<td>conn2.w = 5*nS</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>conn3 = Connection(source, target, 'ge')</th>
<th>conn3 = ... # see above</th>
</tr>
</thead>
<tbody>
<tr>
<td>conn3.connect_random(source, target,</td>
<td>conn3.connect(p=0.02)</td>
</tr>
<tr>
<td>sparseness=0.02,</td>
<td>conn3.w = 2*nS</td>
</tr>
<tr>
<td>weight=2*ns)</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>conn4 = Connection(source, target, 'ge')</th>
<th>conn4 = ... # see above</th>
</tr>
</thead>
<tbody>
<tr>
<td>conn4.connect_one_to_one(source, target,</td>
<td>conn4.connect(j='i')</td>
</tr>
<tr>
<td>weight=4*nS)</td>
<td>conn4.w = 4*nS</td>
</tr>
</tbody>
</table>

| conn5 = IdentityConnection(source,  | conn5 = Synapses(source, target,  |
| target,  | 'w : siemens (shared)') |
| weight=3*nS) | conn5.w = 3*nS |

### Weight matrices

Brian 2's `Synapses` class does not support setting the weights of a neuron with a weight matrix. However, `Synapses.connect` creates the synapses in a predictable order (first all synapses for the first pre-synaptic cell, then all synapses for the second pre-synaptic cell, etc.), so a reshaped “flat” weight matrix can be used:
However note that if your weight matrix can be described mathematically (e.g. random as in the example above), then you should not create a weight matrix in the first place but use Brian 2’s mechanism to set variables based on mathematical expressions (in the above case: conn5.w = 'rand()'). Especially for big connection matrices this will have better performance, since it will be executed in generated code. You should only resort to explicit weight matrices when there is no alternative (e.g. to load weights from previous simulations).

In Brian 1, you can restrict the functions connect, connect_random, etc. to subgroups. Again, there is no direct equivalent to this in Brian 2, but the general string syntax allows you to make connections conditional on logical statements that refer to pre-/post-synaptic indices and can therefore also used to restrict the connection to a subgroup of cells. When you set the synaptic weights, you can however use subgroups to restrict the subset of weights you want to set.

### Connections defined by functions

Brian 1 allowed you to pass in a function as the value for the weight argument in a `connect` call (and also for the sparseness argument in `connect_random`). You should be able to replace such use cases by the the general, string-expression based method:
Brian 1 Documentation, Release 2.3

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
</table>
| conn8 = Connection(source, target, 'ge')
conn8.connect_full(source, target,
weight=lambda i,
→j:(1+cos(i-j))*2*nS) | conn8 = Synapses(source, target, 'w:
→siemens',
on_pre='ge += w')
conn8.connect()
conn8.w = '(1 + cos(i - j))*2*nS'

| conn9 = Connection(source, target, 'ge')
conn9.connect_random(source, target,
sparseness=0.02,
→weight=lambda:rand()*nS) | conn9 = ...  # see above
conn9.connect(p=0.02)
conn9.w = 'rand()*nS'

| conn10 = Connection(source, target, 'ge')
conn10.connect_random(source, target,
sparseness=lambda:
→i,j:exp(-abs(i-j)*.1),
weight=2*ns) | conn10 = ...  # see above
conn10.connect(p='exp(-abs(i - j)*.1)')
conn10.w = 2*nS

### Delays

The specification of delays changed in several aspects from Brian 1 to Brian 2: In Brian 1, delays where homogeneous by default, and heterogeneous delays had to be marked by delay=True, together with the specification of the maximum delay. In Brian 2, homogeneous delays are the default and you do not have to state the maximum delay. Brian 1’s syntax of specifying a pair of values to get randomly distributed delays in that range is no longer supported, instead use Brian 2’s standard string syntax:

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
</table>
| conn11 = Connection(source, target, 'ge',
→ delay=True,
max_delay=5*ms)
conn11.connect_full(source, target,
→weight=3*nS,
delay=(0*ms, 5*ms)) | conn11 = Synapses(source, target, 'w :
→siemens',
on_pre='ge += w')
conn11.connect()
conn11.w = 3*nS
conn11.delay = 'rand()*5*ms'

### Modulation

In Brian 2, there’s no need for the modulation keyword that Brian 1 offered, you can describe the modulation as part of the on_pre action:

1.4. Changes for Brian 1 users
### Structure

There’s no equivalent for Brian 1’s `structure` keyword in Brian 2, synapses are always stored in a sparse data structure. There is currently no support for changing synapses at run time (i.e. the “dynamic” structure of Brian 1).

#### Converting Brian 1’s Synapses class

Brian 2’s `Synapses` class works for the most part like the class of the same name in Brian 1. There are however some differences in details, listed below:

#### Synaptic models

The basic syntax to define a synaptic model is unchanged, but the keywords `pre` and `post` have been renamed to `on_pre` and `on_post`, respectively.

#### Lumped variables (summed variables)

The syntax to define lumped variables (we use the term “summed variables” in Brian 2) has been changed: instead of assigning the synaptic variable to the neuronal variable you’ll have to include the summed variable in the synaptic equations with the flag `(summed)`:  

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>stdp_syn = Synapses(inputs, neurons, ...</code></td>
<td></td>
</tr>
<tr>
<td><code>model='' w:1</code></td>
<td></td>
</tr>
<tr>
<td><code>dApre/dt = -Apre/ taupre : 1 (event-driven)</code></td>
<td></td>
</tr>
<tr>
<td><code>dApost/dt = -Apost/ taupost : 1 (event-driven)'''</code></td>
<td></td>
</tr>
<tr>
<td><code>pre='''ge += w</code></td>
<td></td>
</tr>
<tr>
<td><code>Apre += delta_Apre</code></td>
<td></td>
</tr>
<tr>
<td><code>Apost += delta_Apost</code></td>
<td></td>
</tr>
<tr>
<td><code> Apostle, 0, gmax)'''</code></td>
<td></td>
</tr>
<tr>
<td><code>post='''Apost +=</code></td>
<td></td>
</tr>
<tr>
<td><code>delta_Apost</code></td>
<td></td>
</tr>
<tr>
<td><code>Apre, 0, gmax)'''</code></td>
<td></td>
</tr>
<tr>
<td><code>w = clip(w +</code></td>
<td></td>
</tr>
<tr>
<td><code>w = clip(w +</code></td>
<td></td>
</tr>
<tr>
<td><code>w = clip(w +</code></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>stdp_syn = Synapses(inputs, neurons, ...</code></td>
<td></td>
</tr>
<tr>
<td><code>model='' w:1</code></td>
<td></td>
</tr>
<tr>
<td><code>dApre/dt = -Apre/ taupre : 1 (event-driven)</code></td>
<td></td>
</tr>
<tr>
<td><code>dApost/dt = -Apost/ taupost : 1 (event-driven)'''</code></td>
<td></td>
</tr>
<tr>
<td><code>on_pre='''ge += w</code></td>
<td></td>
</tr>
<tr>
<td><code>Apre += delta_Apre</code></td>
<td></td>
</tr>
<tr>
<td><code>Apost += delta_Apost</code></td>
<td></td>
</tr>
<tr>
<td><code>Apost, 0, gmax)'''</code></td>
<td></td>
</tr>
<tr>
<td><code>on_post='''Apost +=</code></td>
<td></td>
</tr>
<tr>
<td><code>delta_Apost</code></td>
<td></td>
</tr>
<tr>
<td><code>Apre, 0, gmax)'''</code></td>
<td></td>
</tr>
<tr>
<td><code>w = clip(w +</code></td>
<td></td>
</tr>
<tr>
<td><code>w = clip(w +</code></td>
<td></td>
</tr>
<tr>
<td><code>w = clip(w +</code></td>
<td></td>
</tr>
</tbody>
</table>
Brian 1

# a non-linear synapse (e.g. NMDA)
neurons = NeuronGroup(1, model='''
    dv/dt = (gtot - v)/(10*ms) : 1
    gtot : 1''')
syn = Synapses(inputs, neurons,
    model='''
    dg/dt = -a*g+b*x*(1-g) : 1
dx/dt = -c*x : 1
    w : 1 # synaptic weight''',
    on_pre='x += w')
neurons.gtot=S.g

Creating synapses

In Brian 1, synapses were created by assigning True or an integer (the number of synapses) to an indexed Synapses object. In Brian 2, all synapse creation goes through the Synapses.connect function. For examples how to create more complex connection patterns, see the section on translating Connections objects above.

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>syn = Synapses(...) # single synapse syn[3, 5] = True</td>
<td>syn = Synapses(...) # single synapse syn.connect(i=3, j=5)</td>
</tr>
<tr>
<td># all-to-all connections syn[:, :] = True</td>
<td># all-to-all connections syn.connect()</td>
</tr>
<tr>
<td># all to neuron number 1 syn[:, 1] = True</td>
<td># all to neuron number 1 syn.connect(j='1')</td>
</tr>
<tr>
<td># multiple synapses syn[4, 7] = 3</td>
<td># multiple synapses syn.connect(i=4, j=7, n=3)</td>
</tr>
<tr>
<td># connection probability 2% syn[:, :] = 0.02</td>
<td># connection probability 2% syn.connect(p=0.02)</td>
</tr>
</tbody>
</table>

Multiple pathways

As Brian 1, Brian 2 supports multiple pre- or post-synaptic pathways, with separate pre-/post-codes and delays. In Brian 1, you have to specify the pathways as tuples and can then later access them individually by using their index. In Brian 2, you specify the pathways as a dictionary, i.e. by giving them individual names which you can then later use to access them (the default pathways are called pre and post):
Monitoring synaptic variables

Both in Brian 1 and Brian 2, you can record the values of synaptic variables with a StateMonitor. You no longer have to call an explicit indexing function, but you can directly provide an appropriately indexed Synapses object. You can now also use the same technique to index the StateMonitor object to get the recorded values, see the respective section in the Synapses documentation for details.

Inputs (Brian 1 –> 2 conversion)

Brian 2 documentation

For the main documentation about adding external stimulation to a network, see the document Input stimuli.

- Poisson Input
- Spike generation
- Arbitrary time-dependent input (TimedArray)
**Poisson Input**

Brian 2 provides the same two groups that Brian 1 provided: `PoissonGroup` and `PoissonInput`. The mechanism for inhomogeneous Poisson processes has changed: instead of providing a Python function of time, you’ll now have to provide a string expression that is evaluated at every time step. For most use cases, this should allow a direct translation:

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>rates = lambda t:(1+cos(2*pi*t*1*Hz))*10*Hz</code></td>
<td><code>rates = '(1 + cos(2*pi*t*1*Hz)*10*Hz)'</code></td>
</tr>
<tr>
<td><code>group = PoissonGroup(100, rates=rates)</code></td>
<td><code>group = PoissonGroup(100, rates=rates)</code></td>
</tr>
</tbody>
</table>

For more complex rate modulations, the expression can refer to *User-provided functions* and/or you can replace the `PoissonGroup` by a general `NeuronGroup` with a threshold condition `rand()<rates*dt` (which allows you to store per-neuron attributes).

There is currently no direct replacement for the more advanced features of `PoissonInput` (`record`, `freeze`, `copies`, `jitter`, and `reliability` keywords), but various workarounds are possible, e.g. by directly using a `BinomialFunction` in the equations. For example, you can get the functionality of the `freeze` keyword (identical Poisson events for all neurons) by storing the input in a shared variable and then distribute the input to all neurons:

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>group = NeuronGroup(10, 'dv/dt = -v/(10*ms) : 1'</code></td>
<td><code>group = NeuronGroup(10, ''dv/dt = -v / (10*ms) : 1</code></td>
</tr>
<tr>
<td><code>input = PoissonInput(group, N=1000, 'rate=1*Hz,</code></td>
<td><code>shared_input = BinomialFunction(n=1000, p=1*Hz*group.dt)</code></td>
</tr>
<tr>
<td><code>weight=0.1, state='v</code></td>
<td><code>group.run_regularly('''shared_input = poisson_input()*0.1</code></td>
</tr>
<tr>
<td><code>freeze=True)</code></td>
<td><code>v += shared_input''</code>)</td>
</tr>
</tbody>
</table>

**Spike generation**

`SpikeGeneratorGroup` provides mostly the same functionality as in Brian 1. In contrast to Brian 1, there is only one way to specify which neurons spike and when – you have to provide the index array and the times array as separate arguments:
Brian 1

```python
gen1 = SpikeGeneratorGroup(2, [(0, 0*ms),
    → (1, 1*ms)])
gen2 = SpikeGeneratorGroup(2, [(array([0,
    → 1]), 0*ms),
    → (array([0,
    → 1]), 1*ms)])
gen3 = SpikeGeneratorGroup(2, (array([0,
    → 1]),
    → array([0,
    → 1]))*ms))
gen4 = SpikeGeneratorGroup(2, array([[0,
    → 0.0],
    → [0.001]]))
```

Brian 2

```python
gen1 = SpikeGeneratorGroup(2, [0, 1], [0,
    → 1]*ms)
gen2 = SpikeGeneratorGroup(2, [0, 1, 0,
    → 1], [0, 0, 1,
    → 1]*ms)
gen3 = SpikeGeneratorGroup(2, [0, 1], [0,
    → 1]*ms)
gen4 = SpikeGeneratorGroup(2, [0, 1], [0,
    → 1]*ms)
```

**Note:** For large arrays, make sure to provide a `Quantity` array (e.g. `[0, 1, 2]*ms`) and not a list of `Quantity` values (e.g. `[0*ms, 1*ms, 2*ms]`). A list has first to be translated into an array which can take a considerable amount of time for a list with many elements.

There is no direct equivalent of the Brian 1 option to use a generator that updates spike times online. The easiest alternative in Brian 2 is to pre-calculate the spikes and then use a standard `SpikeGeneratorGroup`. If this is not possible (e.g. there are too many spikes to fit in memory), then you can workaround the restriction by using custom code (see `User-provided functions` and `Arbitrary Python code (network operations)`).

### Arbitrary time-dependent input (TimedArray)

For a detailed description of the `TimedArray` mechanism in Brian 2, see `Timed arrays`.

In Brian 1, timed arrays where special objects that could be assigned to a state variable and would then be used to update this state variable at every time step. In Brian 2, a timed array is implemented using the standard `Functions` mechanism which has the advantage that more complex access patterns can be implemented (e.g. by not using `t` as an argument, but something like `t - delay`). This syntax was possible in Brian 1 as well, but was disadvantageous for performance and had other limits (e.g. no unit support, no linear integration). In Brian 2, these disadvantages no longer apply and the function syntax is therefore the only available syntax. You can convert the old-style Brian 1 syntax to Brian 2 as follows:

**Warning:** The example below does not correctly translate the changed semantics of `TimedArray` related to the time. In Brian 1, `TimedArray([0, 1, 2], dt=10*ms)` will return 0 for `t<5*ms`, 1 for `5*ms<=t<15*ms`, and 2 for `t>=15*ms`. Brian 2 will return 0 for `t<10*ms`, 1 for `10*ms<=t<20*ms`, and 2 for `t>=20*ms`. 

---

**Chapter 1. Introduction**
Monitors (Brian 1 -> 2 conversion)

### Brian 1

```
# same input for all neurons
eqs = '''
    dv/dt = (I - v)/tau : volt
    I : volt
    
    group = NeuronGroup(1, model=eqs,
        reset=0*mV,
        threshold=15*mV)

    group.I = TimedArray(linspace(0*mV, 20*mV, 100),
        dt=10*ms)

    # same input for all neurons
    eqs = '''
    dv/dt = (I(t) - v)/tau : volt
    
    group = NeuronGroup(1, model=eqs,
        reset='v = 0*mV',
        threshold='v > 15*mV')
```

```
# neuron-specific input
eqs = '''
    dv/dt = (I - v)/tau : volt
    I : volt
    
    group = NeuronGroup(5, model=eqs,
        reset=0*mV,
        threshold=15*mV)

    values = (linspace(0*mV, 20*mV, 100)[:,
        None] *
        linspace(0, 1, 5))

    group.I = TimedArray(values, dt=10*ms)

    # neuron-specific input
    values = (linspace(0*mV, 20*mV, 100)[:,
        None] *
        linspace(0, 1, 5))

    I = TimedArray(values, dt=10*ms)

    eqs = '''
    dv/dt = (I(t, i) - v)/tau : volt
    
    group = NeuronGroup(5, model=eqs,
        reset='v = 0*mV',
        threshold='v > 15*mV')
```

### Brian 2

```
# same input for all neurons
I = TimedArray(linspace(0*mV, 20*mV, 100),
    dt=10*ms)

# same input for all neurons
eqs = '''
    dv/dt = (I(t) - v)/tau : volt
    
    group = NeuronGroup(1, model=eqs,
        reset='v = 0*mV',
        threshold='v > 15*mV')
```

```
# neuron-specific input
values = (linspace(0*mV, 20*mV, 100)[:,
    None] *
    linspace(0, 1, 5))

I = TimedArray(values, dt=10*ms)

eqs = '''
    dv/dt = (I(t, i) - v)/tau : volt
    
    group = NeuronGroup(5, model=eqs,
        reset='v = 0*mV',
        threshold='v > 15*mV')
```

Monitors (Brian 1 -> 2 conversion)

Brian 2 documentation

For the main documentation about recording network activity, see the document *Recording during a simulation*.

- Monitoring spiking activity
- Monitoring variables

Monitoring spiking activity

The main class to record spiking activity is *SpikeMonitor* which is created in the same way as in Brian 1. However, the internal storage and retrieval of spikes is different. In Brian 1, spikes were stored as a list of pairs (i, t) of the index and time of each spike. In Brian 2, spikes are stored as two arrays i and t, storing the indices and times. You can access these arrays as attributes of the monitor, there’s also a convenience attribute t that returns both at the same time. The following table shows how the spike indices and times can be retrieved in various forms in Brian 1 and Brian 2:
You can also access the spike times for individual neurons. In Brian 1, you could directly index the monitor which is no longer allowed in Brian 2. Instead, ask for a dictionary of spike times and index the returned dictionary:

```python
# dictionary of spike times for each neuron:
spike_dict = mon.spiketimes
# all spikes for neuron 3:
spikes_3 = spike_dict[3]  # (no units)
spikes_3 = mon[3]  # alternative (no units)
```

In Brian 2, `SpikeMonitor` also provides the functionality of the Brian 1 classes `SpikeCounter` and `PopulationSpikeCounter`. If you are only interested in the counts and not in the individual spike events, use `record=False` to save the memory of storing them:

```python
counter = SpikeCounter(group)
pop_counter = PopulationSpikeCounter(group)
#... do the run
# Number of spikes for neuron 3:
count_3 = counter[3]
# Total number of spikes:
total_spikes = pop_counter.nspikes
counter = SpikeMonitor(group, record=False)
#... do the run
# Number of spikes for neuron 3
count_3 = counter.count[3]
# Total number of spikes:
total_spikes = counter.num_spikes
```

Currently Brian 2 provides no functionality to calculate statistics such as correlations or histograms online, there is no equivalent to the following classes that existed in Brian 1: `AutoCorrelogram`, `CoincidenceCounter`, `CoincidenceMatrixCounter`, `ISIHistogramMonitor`, `VanRossumMetric`. You will therefore have to be calculate the corresponding statistics manually after the simulation based on the information stored in the `SpikeMonitor`. If you use the default `Runtime code generation`, you can also create a new Python class that calculates the statistic online (see this example from a Brian 2 tutorial).

You can also access the spike times for individual neurons. In Brian 1, you could directly index the monitor which is no longer allowed in Brian 2. Instead, ask for a dictionary of spike times and index the returned dictionary:

```python
# dictionary of spike times for each neuron:
spike_dict = mon.spiketimes
# all spikes for neuron 3:
spikes_3 = spike_dict[3]  # (no units)
spikes_3 = mon[3]  # alternative (no units)
```

In Brian 2, `SpikeMonitor` also provides the functionality of the Brian 1 classes `SpikeCounter` and `PopulationSpikeCounter`. If you are only interested in the counts and not in the individual spike events, use `record=False` to save the memory of storing them:

```python
counter = SpikeCounter(group)
pop_counter = PopulationSpikeCounter(group)
#... do the run
# Number of spikes for neuron 3:
count_3 = counter[3]
# Total number of spikes:
total_spikes = pop_counter.nspikes
counter = SpikeMonitor(group, record=False)
#... do the run
# Number of spikes for neuron 3
count_3 = counter.count[3]
# Total number of spikes:
total_spikes = counter.num_spikes
```

Currently Brian 2 provides no functionality to calculate statistics such as correlations or histograms online, there is no equivalent to the following classes that existed in Brian 1: `AutoCorrelogram`, `CoincidenceCounter`, `CoincidenceMatrixCounter`, `ISIHistogramMonitor`, `VanRossumMetric`. You will therefore have to be calculate the corresponding statistics manually after the simulation based on the information stored in the `SpikeMonitor`. If you use the default `Runtime code generation`, you can also create a new Python class that calculates the statistic online (see this example from a Brian 2 tutorial).

**Note**: The text has been reformatted into a readable format.
Monitoring variables

Single variables are recorded with a `StateMonitor` in the same way as in Brian 1, but the times and variable values are accessed differently:

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>mon = StateMonitor(group, 'v', record=True)</code></td>
<td><code>mon = StateMonitor(group, 'v', record=True)</code></td>
</tr>
<tr>
<td># ... do the run</td>
<td># ... do the run</td>
</tr>
<tr>
<td># plot the trace of neuron 3:</td>
<td># plot the trace of neuron 3:</td>
</tr>
<tr>
<td>plot(mon.times/ms, mon[3]/mV)</td>
<td>plot(mon.t/ms, mon[3].v/mV)</td>
</tr>
<tr>
<td># plot the traces of all neurons:</td>
<td># plot the traces of all neurons:</td>
</tr>
<tr>
<td>plot(mon.times/ms, mon.values.T/mV)</td>
<td>plot(mon.t/ms, mon.v.T/mV)</td>
</tr>
</tbody>
</table>

Further differences:

- `StateMonitor` now records in the 'start' scheduling slot by default. This leads to a more intuitive correspondence between the recorded times and the values: in Brian 1 (where `StateMonitor` recorded in the 'end' slot) the recorded value at 0ms was not the initial value of the variable but the value after integrating it for a single time step. The disadvantage of this new default is that the very last value at the end of the last time step of a simulation is not recorded anymore. However, this value can be manually added to the monitor by calling `StateMonitor.record_single_timestep`.

- To not record every time step, use the `dt` argument (as for all other classes) instead of specifying a number of timesteps.

- Using `record=False` does no longer provide mean and variance of the recorded variable.

In contrast to Brian 1, `StateMonitor` can now record multiple variables and therefore replaces Brian 1’s `MultiStateMonitor`:

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>mon = MultiStateMonitor(group, ['v', 'w'], record=True)</code></td>
<td><code>mon = StateMonitor(group, ['v', 'w'], record=True)</code></td>
</tr>
<tr>
<td># ... do the run</td>
<td># ... do the run</td>
</tr>
<tr>
<td># plot the traces of v and w for neuron 3</td>
<td># plot the traces of v and w for neuron 3</td>
</tr>
<tr>
<td>plot(mon['v'].times/ms, mon['v'][3]/mV)</td>
<td>plot(mon.t/ms, mon[3].v/mV)</td>
</tr>
<tr>
<td>plot(mon['w'].times/ms, mon['w'][3]/mV)</td>
<td>plot(mon.t/ms, mon[3].w/mV)</td>
</tr>
</tbody>
</table>

To record variable values at the times of spikes, Brian 2 no longer provides a separate class as Brian 1 did (StateSpikeMonitor). Instead, you can use `SpikeMonitor` to record additional variables (in addition to the neuron index and the spike time):
Note that there is no equivalent to StateHistogramMonitor, you will have to calculate the histogram from the recorded values or write your own custom monitor class.

Networks and clocks (Brian 1 → 2 conversion)

Brian 2 documentation

For the main documentation about running simulations, controlling the simulation timestep, etc., see the document *Running a simulation*.

- Clocks and timesteps
- Networks

Clocks and timesteps

Brian’s system of handling clocks has substantially changed. For details about the new system in place see *Setting the simulation time step*. The main differences to Brian 1 are:

- There is no more “clock guessing” – objects either use the defaultclock or a dt/clock value that was explicitly specified during their construction.

- In Brian 2, the time step is allowed to change after the creation of an object and between runs – the relevant value is the value in place at the point of the run() call.

- It is rarely necessary to create an explicit Clock object, most of the time you should use the defaultclock or provide a dt argument during the construction of the object.

- There’s only one Clock class, the (deprecated) FloatClock, RegularClock, etc. classes that Brian 1 provided no longer exist.

- It is no longer possible to (re-)set the time of a clock explicitly, there is no direct equivalent of Clock.reinit and reinit_default_clock. To start a completely new simulation after you have finished a previous one, either create a new Network or use the start_scope() mechanism. To “rewind” a simulation to a previous point, use the new store()/restore() mechanism. For more details, see below and *Running a simulation*. 

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
<tbody>
<tr>
<td># We assume that &quot;group&quot; has a varying threshold</td>
<td># We assume that &quot;group&quot; has a varying threshold</td>
</tr>
<tr>
<td>mon = StateSpikeMonitor(group, 'v')</td>
<td>mon = SpikeMonitor(group, variables='v')</td>
</tr>
<tr>
<td># ... do the run</td>
<td># ... do the run</td>
</tr>
<tr>
<td># plot the mean v at spike time for each neuron</td>
<td># plot the mean v at spike time for each neuron</td>
</tr>
<tr>
<td>mean_values = [mean(mon.values('v', idx)) for idx in range(len(group))]</td>
<td>values = mon.values('v')</td>
</tr>
<tr>
<td>plot(mean_values/mV, 'o')</td>
<td>plot(mean_values/mV, 'o')</td>
</tr>
</tbody>
</table>
Networks

Both Brian 1 and Brian 2 offer two ways to run a simulation: either by explicitly creating a `Network` object, or by using a `MagicNetwork`, i.e., a simple `run()` statement.

Explicit network

The mechanism to create explicit `Network` objects has not changed significantly from Brian 1 to Brian 2. However, creating a new `Network` will now also automatically reset the clock back to 0s, and stricter checks no longer allow the inclusion of the same object in multiple networks.

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>group = ...</td>
<td>group = ...</td>
</tr>
<tr>
<td>mon = ...</td>
<td>mon = ...</td>
</tr>
<tr>
<td>net = Network(group, mon)</td>
<td>net = Network(group, mon)</td>
</tr>
<tr>
<td>net.run(1*ms)</td>
<td>net.run(1*ms)</td>
</tr>
<tr>
<td>reinit()</td>
<td># new network starts at 0s</td>
</tr>
<tr>
<td>group = ...</td>
<td>group = ...</td>
</tr>
<tr>
<td>mon = ...</td>
<td>mon = ...</td>
</tr>
<tr>
<td>net = Network(group, mon)</td>
<td>net = Network(group, mon)</td>
</tr>
<tr>
<td>net.run(1*ms)</td>
<td>net.run(1*ms)</td>
</tr>
</tbody>
</table>

“Magic” network

For most simple, “flat”, scripts (see e.g., the Examples), the `run()` statement in Brian 2 automatically collects all the Brian objects (`NeuronGroup`, etc.) into a “magic” network in the same way as Brian 1 did. The logic behind this collection has changed, though, with important consequences for more complex simulation scripts: in Brian 1, the magic network includes all Brian objects that have been `created` in the same execution frame as the `run()` call. Objects that are created in other functions could be added using `magic_return` and `magic_register`. In Brian 2, the magic network contains all Brian objects that are `visible` in the same execution frame as the `run()` call. The advantage of the new system is that it is clearer what will be included in the network and there is no danger of including previously created, but no longer needed, objects in a simulation. E.g., in the following example, a common mistake in Brian 1 was to not include the `clear()`, which meant that each run not only simulated the current objects, but also all objects from previous loop iterations. Also, without the `reinit_default_clock()`, each run would start at the end time of the previous run. In Brian 2, this loop does not need any explicit clearing up, each `run()` will only simulate the object that it “sees” (group1, group2, syn, and mon) and start each simulation at 0s:

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>for r in range(100):</code></td>
<td><code>for r in range(100):</code></td>
</tr>
<tr>
<td>reinit_default_clock()</td>
<td></td>
</tr>
<tr>
<td>clear()</td>
<td></td>
</tr>
<tr>
<td>group1 = NeuronGroup(...)</td>
<td>group1 = NeuronGroup(...)</td>
</tr>
<tr>
<td>group2 = NeuronGroup(...)</td>
<td>group2 = NeuronGroup(...)</td>
</tr>
<tr>
<td>syn = Synapses(group1, group2, ...)</td>
<td>syn = Synapses(group1, group2, ...)</td>
</tr>
<tr>
<td>mon = SpikeMonitor(group2)</td>
<td>mon = SpikeMonitor(group2)</td>
</tr>
<tr>
<td>run(1*second)</td>
<td>run(1*second)</td>
</tr>
</tbody>
</table>
There is no replacement for the `magic_return` and `magic_register` functions. If the returned object is stored in a variable at the level of the `run()` call, then it is no longer necessary to use `magic_return`, as the returned object is “visible” at the level of the `run()` call:

```
Brian 1
@magic_return
def f():
    return PoissonGroup(100,
        rates=100*Hz)
pg = f()  # needs magic_return
mon = SpikeMonitor(pg)
run(100*ms)
Brian 2
def f():
    return PoissonGroup(100,
        rates=100*Hz)
png = f()  # is "visible" and will be included
mon = SpikeMonitor(png)
run(100*ms)
```

The general recommendation is however: if your script is complex (multiple functions/files/classes) and you are not sure whether some objects will be included in the magic network, use an explicit `Network` object.

Note that one consequence of the “is visible” approach is that objects stored in containers (lists, dictionaries, …) will not be automatically included in Brian 2. Use an explicit `Network` object to get around this restriction:

```
Brian 1
groups = {'exc': NeuronGroup(...),
            'inh': NeuronGroup(...)}
...
run(5*ms)
Brian 2
groups = {'exc': NeuronGroup(...),
            'inh': NeuronGroup(...)}
...
et = Network(groups)
et.run(5*ms)
```

### External constants

In Brian 2, external constants are taken from the surrounding namespace at the point of the `run()` call and not when the object is defined (for other ways to define the namespace, see `External variables`). This allows to easily change external constants between runs, in contrast to Brian 1 where the whether this worked or not depended on details of the model (e.g. whether linear integration was used):
Preferences (Brian 1 -> 2 conversion)

Brian 2 documentation

For the main documentation about preferences, see the document Preferences.

In Brian 1, preferences were set either with the function set_global_preferences or by creating a module somewhere on the Python path called brian_global_config.py.

Setting preferences

The function set_global_preferences no longer exists in Brian 2. Instead, importing from brian2 gives you a variable prefs that can be used to set preferences. For example, in Brian 1 you would write:

```python
set_global_preferences(weavecompiler='gcc')
```

In Brian 2 you would write:

```python
prefs.codegen.cpp.compiler = 'gcc'
```

Configuration file

The module brian_global_config.py is not used by Brian 2, instead we search for configuration files in the current directory, user directory or installation directory. In Brian you would have a configuration file that looks like this:

```python
from brian.globalprefs import *
set_global_preferences(weavecompiler='gcc')
```

In Brian 2 you would have a file like this:
Preference name changes

- **defaultclock**: removed because it led to unclear behaviour of scripts.
- **useweave_linear_diffeq**: removed because it was no longer relevant.
- **useweave**: now replaced by `codegen.target`.
- **weavecompiler**: now replaced by `codegen.cpp.compiler`.
- **gcc_options**: now replaced by `codegen.cpp.extra_compile_args_gcc`.
- **openmp**: now replaced by `devices.cpp_standalone.openmp_threads`.
- **usecodegen***: removed because it was no longer relevant.
- **usenewpropagate**: removed because it was no longer relevant.
- **usecstdp**: removed because it was no longer relevant.
- **brianhears_usegpu**: removed because Brian Hears doesn’t exist in Brian 2.
- **magic_useframes**: removed because it was no longer relevant.

Multicompartmental models (Brian 1 → 2 conversion)

Brian 2 documentation

Support for multicompartmental models is now an integral part of Brian 2 (an early version of it was included as an experimental module in Brian 1). See the document *Multicompartment models*.

Brian 1 offered support for simple multi-compartmental models in the `compartments` module. This module allowed you to combine the equations for several compartments into a single `Equations` object. This is only a suitable solution for simple morphologies (e.g. “ball-and-stick” models) but has the advantage over using `SpatialNeuron` that you can have several of such neurons in a `NeuronGroup`.

If you already have a definition of a model using Brian 1’s `compartments` module, then you can simply print out the equations and use them directly in Brian 2. For simple models, writing the equations without that help is rather straightforward anyway:
### Library models (Brian 1 → 2 conversion)

- **Neuron models**
- **Ionic currents**
- **Synapses**

#### Neuron models

The neuron models in Brian 1’s `brian.library.IF` package are nothing more than shorthands for equations. The following table shows how the models from Brian 1 can be converted to explicit equations (and reset statements in the case of the adaptive exponential integrate-and-fire model) for use in Brian 2. The examples include a “current” $I$ (depending on the model not necessarily in units of Ampère) and could e.g. be used to plot the f-I curve of the neuron.

#### Perfect integrator

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>eqs = (perfect_IF(tau=10*ms) +</td>
<td>tau = 10*ms</td>
</tr>
<tr>
<td>Current(’I : volt’))</td>
<td>eqs = ’’dvm/dt = I/tau : volt</td>
</tr>
<tr>
<td>group = NeuronGroup(N, eqs,</td>
<td>I : volt’’</td>
</tr>
<tr>
<td>’\textprime\textprime’,</td>
<td></td>
</tr>
<tr>
<td>reset=’v = -70*mV’)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Leaky integrate-and-fire neuron

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
</table>
| eqs = (leaky_IF(tau=10*ms, El=-70*mV) + Current('I : volt'))
| group = ... # see above |

```
tau = 10*ms; El = -70*mV
eqs = '''dvm/dt = ((El - vm) + I)/tau : volt
       I : volt'''
group = ... # see above
```

Exponential integrate-and-fire neuron

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
</table>
| eqs = (exp_IF(C=1*nF, gL=30*nS, EL=-70*mV, VT=-50*mV, DeltaT=2*mV) + Current('I : amp'))
| group = ... # see above |

```
C = 1*nF; gL = 30*nS; EL = -70*mV; VT = -50*mV; DeltaT = 2*mV
eqs = '''dvm/dt = (gL*EL - vm) + gL*DeltaT*exp((vm-VT)/DeltaT) + I)/C : volt
        I : amp'''
group = ... # see above
```

Quadratic integrate-and-fire neuron

<table>
<thead>
<tr>
<th>Brian 1</th>
<th>Brian 2</th>
</tr>
</thead>
</table>
| eqs = (quadratic_IF(C=1*nF, a=5*nS/mV, EL=-70*mV, VT=-50*mV) + Current('I : amp'))
| group = ... # see above |

```
C = 1*nF; a=5*nS/mV; EL=-70*mV; VT=-50*mV
eqs = '''dvm/dt = (a*(vm-EL)*(vm-VT) + I)/C : volt
        I : amp'''
group = ... # see above
```
Izhikevich neuron

### Brian 1

```python
eqs = (Izhikevich(a=0.02/ms, b=0.2/ms) + Current('I : volt/second'))
group = ... # see above
```

### Brian 2

```
a = 0.02/ms; b = 0.2/ms
eqs = '''dvm/dt = (0.04/ms/mV)*vm**2+(5/ms)*vm+140*mV/ms-w + I : volt
dw/dt = a*(b*vm-w) : volt/second
I : volt/second'''
group = ... # see above
```

Adaptive exponential integrate-and-fire neuron (“Brette-Gerstner model”)

### Brian 1

```python
# AdEx, aEIF, and Brette_Gerstner all refer to the same model
eqs = (aEIF(C=1*nF, gL=30*nS, EL=-70*mV, VT=-50*mV, DeltaT=2*mV, tauw=150*ms, a=4*nS) + Current('I:amp'))
group = NeuronGroup(N, eqs,
   threshold='v > -20*mV',
   reset=AdaptiveReset(Vr=-70*mV, b=0.08*nA))
```

### Brian 2

```
C = 1*nF; gL = 30*nS; EL = -70*mV; VT = -50*mV; DeltaT = 2*mV; tauw = 150*ms; a = 4*nS
eqs = '''dvm/dt = (gL*(EL-vm)+gL*DeltaT*exp((vm-VT)/DeltaT) -w + I)/C : volt
dw/dt=(a*(vm-EL)-w)/tauw : amp
I : amp'''
group = NeuronGroup(N, eqs,
   threshold='vm > -20*mV',
   reset='vm=-70*mV; w += 0.08*nA')
```

Ionic currents

Brian 1’s functions for ionic currents, provided in `brian.library.ionic_currents` correspond to the following equations (note that the currents follow the convention to use a shifted membrane potential, i.e. the membrane potential at rest is 0mV):
from brian.library.ionic_currents import *
defaultclock.dt = 0.01*ms
eqs_leak = leak_current(gl=60*nS, El=10.6*mV, current_name='I_leak')
eqs_K = K_current_HH(gmax=7.2*uS, EK=-12*mV, current_name='I_K')
eqs_Na = Na_current_HH(gmax=24*uS, ENa=115*mV, current_name='I_Na')
eqs = (MembraneEquation(C=200*pF) +
    eqs_leak + eqs_K + eqs_Na +
    Current('I_inj : amp'))

defaultclock.dt = 0.01*ms
gl = 60*nS; El = 10.6*mV
eqs_leak = Equations('I_leak = gl*(El - vm) : amp')
g_K = 7.2*uS; EK = -12*mV
eqs_K = Equations('''I_K = g_K*n**4*(EK-vm) : amp''')
dn/dt = alphan*(1-n)-betan*n : 1
alphan = 0.01*(10*mV-vm)/(exp(10.1*vm/mV)-1)/mV/ms : Hz
betan = 0.125*exp(-0.0125*vm/mV)/ms : Hz

C = 200*pF
eqs = Equations('''dvm/dt = (I_leak + I_K + I_Na + I_inj)/C : volt''') + eqs_leak + eqs_K + eqs_Na

Synapses

Brian 1’s synaptic models, provided in brian.library.synapses can be converted to the equivalent Brian 2 equations as follows:
Current-based synapses

Brian 1

\[ \text{syn\_eqs} = \text{exp\_current('s', \texttt{tau}=5*ms, } \texttt{current\_name='I\_syn')} \]
\[ \text{eqs} = (\text{MembraneEquation}(C=1*nF) + \text{Current('Im = gl*(El-vm) : amp') + syn\_eqs}) \]
\[ \text{group} = \text{NeuronGroup}(N, \text{eqs, threshold='vm>-50*mV', reset='vm=-70*mV'}) \]
\[ \text{syn} = \text{Synapses(source, group, pre='s += 1*nA')} \]
\[ \# ... connect synapses, etc. \]

Brian 2

\[ \text{tau} = 5*ms \]
\[ \text{syn\_eqs} = \text{Equations('dI\_syn/dt = -I\_syn/tau : amp')} \]
\[ \text{eqs} = (\text{Equations('dvm/dt = (gl*(El - vm) + I\_syn)/C : volt') + syn\_eqs}) \]
\[ \text{group} = \text{NeuronGroup}(N, \text{eqs, threshold='vm>-50*mV', reset='vm=-70*mV'}) \]
\[ \text{syn} = \text{Synapses(source, group, on\_pre='I\_syn += 1*nA')} \]
\[ \# ... connect synapses, etc. \]

Conductance-based synapses

Brian 1

\[ \text{syn\_eqs} = \text{alpha\_current('s', \texttt{tau}=2.5*ms, } \texttt{current\_name='I\_syn')} \]
\[ \text{eqs} = ... \# remaining code as above \]

Brian 2

\[ \text{tau} = 2.5*ms \]
\[ \text{syn\_eqs} = \text{Equations(''dI\_syn/dt = (s - I\_syn)/tau : amp')} \]
\[ \text{ds/dt = -s/tau : amp''}) \]
\[ \text{group} = \text{NeuronGroup}(N, \text{eqs, threshold='vm>-50*mV', reset='vm=-70*mV'}) \]
\[ \text{syn} = \text{Synapses(source, group, on\_pre='s += 1*nA')} \]
\[ \# ... connect synapses, etc. \]

Continued on next page
### Brian Hears

Deprecated since version 2.2.2.2: Use the `brian2hears` package instead.

This module is designed for users of the Brian 1 library “Brian Hears”. It allows you to use Brian Hears with Brian 2 with only a few modifications (although it’s not compatible with the “standalone” mode of Brian 2). The way it works is by acting as a “bridge” to the version in Brian 1. To make this work, you must have a copy of Brian 1 installed (preferably the latest version), and import Brian Hears using:

```python
from brian2.hears import *
```

Many scripts will run without any changes, but there are a few caveats to be aware of. Mostly, the problems are due to the fact that the units system in Brian 2 is not 100% compatible with the units system of Brian 1.

`FilterbankGroup` now follows the rules for `NeuronGroup` in Brian 2, which means some changes may be necessary to match the syntax of Brian 2, for example, the following would work in Brian 1 Hears:

```python
# Leaky integrate-and-fire model with noise and refractoriness
eqs = '''
   dv/dt = (I-v)/(1*ms)+0.2*xi*(2/(1*ms))**.5 : 1
   I : 1
'''

anf = FilterbankGroup(ihc, 'I', eqs, reset=0, threshold=1, refractory=5*ms)
```

However, in Brian 2 Hears you would need to do:

```python
# Leaky integrate-and-fire model with noise and refractoriness
eqs = '''
   dv/dt = (I-v)/(1*ms)+0.2*xi*(2/(1*ms))**.5 : 1 (unless refractory)
   I : 1
'''

anf = FilterbankGroup(ihc, 'I', eqs, reset='v=0', threshold='v>1', refractory=5*ms)
```
Slicing sounds no longer works. Previously you could do, e.g. `sound[:20*ms]` but with Brian 2 you would need to do `sound.slice(0*ms, 20*ms)`.

In addition, some functions may not work correctly with Brian 2 units. In most circumstances, Brian 2 units can be used interchangeably with Brian 1 units in the bridge, but in some cases it may be necessary to convert units from one format to another, and to do that you can use the functions `convert_unit_b1_to_b2` and `convert_unit_b2_to_b1`.

## 1.5 Known issues

In addition to the issues noted below, you can refer to our bug tracker on GitHub.

### List of known issues

- **Cannot find msvcr90d.dll**
- “AttributeError: MSVCCompiler instance has no attribute ‘compiler_cxx’”
- “Missing compiler_cxx fix for MSVCCompiler”
- Problems with numerical integration
- Jupyter notebooks and C++ standalone mode progress reporting
- Parallel Brian simulations with the `weave` code generation target
- Parallel Brian simulations with C++ standalone
- Slow C++ standalone simulations
- Cython fails with compilation error on OS X: `error: use of undeclared identifier 'isinf'`
- CMD windows open when running Brian on Windows with the Spyder 3 IDE

### 1.5.1 Cannot find msvcr90d.dll

If you see this message coming up, find the file `PythonDir\Lib\site-packages\numpy\distutils\mingw32ccompiler.py` and modify the line `msvcr_dbg_success = build_msvcr_library(debug=True)` to read `msvcr_dbg_success = False` (you can comment out the existing line and add the new line immediately after).

### 1.5.2 “AttributeError: MSVCCompiler instance has no attribute ‘compiler_cxx’”

This is caused by a bug in some versions of numpy on Windows. The easiest solution is to update to the latest version of numpy.

If that isn’t possible, a hacky solution is to modify the numpy code directly to fix the problem. The following change may work. Modify line 388 of `numpy/distutils/ccompiler.py` from `elif not self.compiler_cxx:` to `elif not hasattr(self, 'compiler_cxx') or not self.compiler_cxx:`. If the line number is different, it should be nearby. Search for `elif not self.compiler_cxx` in that file.
1.5.3 “Missing compiler_cxx fix for MSVCCompiler”

If you keep seeing this message, do not worry. It’s not possible for us to hide it, but doesn’t indicate any problems.

1.5.4 Problems with numerical integration

In some cases, the automatic choice of numerical integration method will not be appropriate, because of a choice of parameters that couldn’t be determined in advance. In this case, typically you will get nan (not a number) values in the results, or large oscillations. In this case, Brian will generate a warning to let you know, but will not raise an error.

1.5.5 Jupyter notebooks and C++ standalone mode progress reporting

When you run simulations in C++ standalone mode and enable progress reporting (e.g. by using `report='text'` as a keyword argument), the progress will not be displayed in the jupyter notebook. If you started the notebook from a terminal, you will find the output there. Unfortunately, this is a tricky problem to solve at the moment, due to the details of how the jupyter notebook handles output.

1.5.6 Parallel Brian simulations with the `weave` code generation target

When using the `weave` code generation target (the default runtime target on Python 2.x, see Runtime code generation for details), you should avoid running multiple Brian simulations in parallel. The `weave` package caches compiled files, but this cache is not prepared for multiple concurrent updates. If two Python scripts (or two processes started from the same Python script, e.g. via the `multiprocessing` package) try to store compilation results at the same time, `weave` will crash with an error message. The `numpy` and `cython` targets are not affected by this problem.

1.5.7 Parallel Brian simulations with C++ standalone

Simulations using the C++ standalone device will create code and store results in a dedicated directory (`output`, by default). If you run multiple simulations in parallel, you have to take care that these simulations do not use the same directory – otherwise, everything from compilation errors to incorrect results can happen. Either chose a different directory name for each simulation and provide it as the `directory` argument to the `set_device` or `build` call, or use `directory=None` which will use a randomly chosen unique temporary directory (in `/tmp` on Unix-based systems) for each simulation. If you need to know the directory name, you can access it after the simulation run via `device.project_dir`.

1.5.8 Slow C++ standalone simulations

Some versions of the GNU standard library (in particular those used by recent Ubuntu versions) have a bug that can dramatically slow down simulations in C++ standalone mode on modern hardware (see #803). As a workaround, Brian will set an environment variable `LD_BIND_NOW` during the execution of standalone simulations which changes the way the library is linked so that it does not suffer from this problem. If this environment variable leads to unwanted behaviour on your machine, change the `prefs.devices.cpp_standalone.run_environment_variables` preference.

1.5.9 Cython fails with compilation error on OS X: error: use of undeclared identifier 'isinf'

Try setting the environment variable `MACOSX_DEPLOYMENT_TARGET=10.9`.
1.5.10 CMD windows open when running Brian on Windows with the Spyder 3 IDE

This is due to the interaction with the integrated ipython terminal. Either change the run configuration to “Execute in an external system terminal” or patch the internal Python function used to spawn processes as described in github issue #1140.

1.6 Support

If you are stuck with a problem using Brian, please do get in touch at our email support list.

You can save time by following this procedure when reporting a problem:

1. Do try to solve the problem on your own first. Read the documentation, including using the search feature, index and reference documentation.

2. Search the mailing list archives to see if someone else already had the same problem.

3. Before writing, try to create a minimal example that reproduces the problem. You’ll get the fastest response if you can send just a handful of lines of code that show what isn’t working.

1.7 Contributor Covenant Code of Conduct

1.7.1 Our Pledge

In the interest of fostering an open and welcoming environment, we as contributors and maintainers pledge to making participation in our project and our community a harassment-free experience for everyone, regardless of age, body size, disability, ethnicity, sex characteristics, gender identity and expression, level of experience, education, socio-economic status, nationality, personal appearance, race, religion, or sexual identity and orientation.

1.7.2 Our Standards

Examples of behavior that contributes to creating a positive environment include:

- Using welcoming and inclusive language
- Being respectful of differing viewpoints and experiences
- Gracefully accepting constructive criticism
- Focusing on what is best for the community
- Showing empathy towards other community members

Examples of unacceptable behavior by participants include:

- The use of sexualized language or imagery and unwelcome sexual attention or advances
- Trolling, insulting/derogatory comments, and personal or political attacks
- Public or private harassment
- Publishing others’ private information, such as a physical or electronic address, without explicit permission
- Other conduct which could reasonably be considered inappropriate in a professional setting
1.7.3 Our Responsibilities

Project maintainers are responsible for clarifying the standards of acceptable behavior and are expected to take appropriate and fair corrective action in response to any instances of unacceptable behavior.

Project maintainers have the right and responsibility to remove, edit, or reject comments, commits, code, wiki edits, issues, and other contributions that are not aligned to this Code of Conduct, or to ban temporarily or permanently any contributor for other behaviors that they deem inappropriate, threatening, offensive, or harmful.

1.7.4 Scope

This Code of Conduct applies both within project spaces and in public spaces when an individual is representing the project or its community. Examples of representing a project or community include using an official project e-mail address, posting via an official social media account, or acting as an appointed representative at an online or offline event. Representation of a project may be further defined and clarified by project maintainers.

1.7.5 Enforcement

Instances of abusive, harassing, or otherwise unacceptable behavior may be reported by contacting the project team at team@briansimulator.org. All complaints will be reviewed and investigated and will result in a response that is deemed necessary and appropriate to the circumstances. The project team is obligated to maintain confidentiality with regard to the reporter of an incident. Further details of specific enforcement policies may be posted separately.

Project maintainers who do not follow or enforce the Code of Conduct in good faith may face temporary or permanent repercussions as determined by other members of the project’s leadership.

1.7.6 Attribution

This Code of Conduct is adapted from the Contributor Covenant, version 1.4, available at https://www.contributor-covenant.org/version/1/4/code-of-conduct.html

For answers to common questions about this code of conduct, see https://www.contributor-covenant.org/faq
The tutorial consists of a series of Jupyter Notebooks\(^1\).

For more information about how to use Jupyter Notebooks, see the Jupyter Notebook documentation.

2.1 Introduction to Brian part 1: Neurons

All Brian scripts start with the following. If you’re trying this notebook out in the Jupyter notebook, you should start by running this cell.

```python
from brian2 import *
```

Later we’ll do some plotting in the notebook, so we activate inline plotting in the notebook by doing this:

```python
%matplotlib inline
```

If you are not using the Jupyter notebook to run this example (e.g. you are using a standard Python terminal, or you copy&paste these example into an editor and run them as a script), then plots will not automatically be displayed. In this case, call the `show()` command explicitly after the plotting commands.

2.1.1 Units system

Brian has a system for using quantities with physical dimensions:

```python
20*volt
```

\[20.0 \text{ V}\]

All of the basic SI units can be used (volt, amp, etc.) along with all the standard prefixes (m=milli, p=pico, etc.), as well as a few special abbreviations like mV for millivolt, pF for picofarad, etc.

\(^1\) Formerly known as “IPython Notebooks”.
Also note that combinations of units with work as expected:

\[10 \text{nA} \times 5 \text{Mohm}\]

50.0 mV

And if you try to do something wrong like adding amps and volts, what happens?

\[5 \text{amp} + 10 \text{volt}\]

```
DimensionMismatchError
Traceback (most recent call last)
<ipython-input-8-245c0c0332d1> in <module>()
----> 1 5*amp+10*volt
/home/marcel/programming/brian2/brian2/units/fundamentalunits.pyc in __add__(self, other)
     1427     return self._binary_operation(other, operator.add,
     1428         fail_for_mismatch=True,
--> 1429         operator_str='+')
     1430     def __radd__(self, other):
     1431     def __add__(self, other):

/home/marcel/programming/brian2/brian2/units/fundamentalunits.pyc in fail_for_dimension_mismatch(obj1, obj2, error_message, **error_quantities)
     184   raise DimensionMismatchError(error_message, dim1)
     185     else:
--> 186   raise DimensionMismatchError(error_message, dim1, dim2)
     187     return dim1, dim2
```

(continues on next page)
If you haven’t see an error message in Python before that can look a bit overwhelming, but it’s actually quite simple and it’s important to know how to read these because you’ll probably see them quite often.

You should start at the bottom and work up. The last line gives the error type `DimensionMismatchError` along with a more specific message (in this case, you were trying to add together two quantities with different SI units, which is impossible).

Working upwards, each of the sections starts with a filename (e.g. `C:\Users\Dan\...`) with possibly the name of a function, and then a few lines surrounding the line where the error occurred (which is identified with an arrow).

The last of these sections shows the place in the function where the error actually happened. The section above it shows the function that called that function, and so on until the first section will be the script that you actually run. This sequence of sections is called a traceback, and is helpful in debugging.

If you see a traceback, what you want to do is start at the bottom and scan up the sections until you find your own file because that’s most likely where the problem is. (Of course, your code might be correct and Brian may have a bug in which case, please let us know on the email support list.)

### 2.1.2 A simple model

Let’s start by defining a simple neuron model. In Brian, all models are defined by systems of differential equations. Here’s a simple example of what that looks like:

```python
tau = 10*ms
eqs = '''
dv/dt = (1-v)/tau : 1
'''
```

In Python, the notation ```'''``` is used to begin and end a multi-line string. So the equations are just a string with one line per equation. The equations are formatted with standard mathematical notation, with one addition. At the end of a line you write : `unit` where `unit` is the SI unit of that variable. Note that this is not the unit of the two sides of the equation (which would be 1/second), but the unit of the variable defined by the equation, i.e. in this case `v`.

Now let’s use this definition to create a neuron.

```python
G = NeuronGroup(1, eqs)
```

In Brian, you only create groups of neurons, using the class `NeuronGroup`. The first two arguments when you create one of these objects are the number of neurons (in this case, 1) and the defining differential equations.

Let’s see what happens if we didn’t put the variable `tau` in the equation:

```python
eqs = '''
dv/dt = 1-v : 1
'''
G = NeuronGroup(1, eqs)
run(100*ms)
```

BrianObjectExceptionTraceback (most recent call last)

```
<ipython-input-11-97ed109f5888> in <module>()
```
3 '''
4 G = NeuronGroup(1, eqs)
5 run(100*ms)
6
/home/marcel/programming/brian2/brian2/units/fundamentalunits.pyc in new_f(*args, **kwds)
   2373         get_
   2374         dimensions(newkeyset[k])
   2375         result = f(*args, **kwds)
   2376         if 'result' in au:
   2377             if au['result'] == bool:

/home/marcel/programming/brian2/brian2/core/magic.pyc in run(duration, report, report_period, namespace, profile, level)
   369     return magic_network.run(duration, report=report, report_period=report_period, namespace=namespace, profile=profile, level=level+2)
   370     run.__module__ = __name__
   371
/home/marcel/programming/brian2/brian2/core/base.pyc in device_override_decorated_function(*args, **kwds)
   274         return getattr(curdev, name)(*args, **kwds)
   275     else:
   276         return func(*args, **kwds)
   277         device_override_decorated_function.__doc__ = func.__doc__

/home/marcel/programming/brian2/brian2/units/fundamentalunits.pyc in new_f(*args, **kwds)
   2373         get_
   2374         dimensions(newkeyset[k])
   2375         result = f(*args, **kwds)
   2376         if 'result' in au:
   2377             if au['result'] == bool:

/home/marcel/programming/brian2/brian2/core/network.pyc in run(self, duration, report, report_period, namespace, profile, level)
(continues on next page)
namespace = get_local_namespace(level=level+3)

self.before_run(namespace)

if len(self.objects)==0:

/home/marcel/programming/brian2/brian2/core/base.pyc in device_override_decorated_function
   274     return getattr(curdev, name)(*args, **kwds)
   275     else:
--> 276     return func(*args, **kwds)
   277   278 device_override_decorated_function.__doc__ = func.__doc__

/home/marcel/programming/brian2/brian2/core/network.pyc in before_run(self, run_namespace)
   841     obj.before_run(run_namespace)
   842     except Exception as ex:
--> 843     raise brian_object_exception("An error occurred when preparing an object.", obj, ex)
   844   845     # Check that no object has been run as part of another network before

BrianObjectException: Original error and traceback:
Traceback (most recent call last):
  File "/home/marcel/programming/brian2/brian2/core/network.py", line 841, in before_run
    obj.before_run(run_namespace)
  File "/home/marcel/programming/brian2/brian2/groups/neurongroup.py", line 811, in before_run
    self.equations.check_units(self, run_namespace=run_namespace)
  File "/home/marcel/programming/brian2/brian2/equations/equations.py", line 963, in check_units
    *ex.dims)
DimensionMismatchError: Inconsistent units in differential equation defining variable v:
Expression 1-v does not have the expected unit hertz (unit is 1).

Error encountered with object named "neurongroup_1".
Object was created here (most recent call only, full details in debug log):
  File "<ipython-input-11-97ed109f5888>", line 4, in <module>
    G = NeuronGroup(1, eqs)

An error occurred when preparing an object. DimensionMismatchError: Inconsistent units in differential equation defining variable v:
Expression 1-v does not have the expected unit hertz (unit is 1).
(See above for original error message and traceback.)

An error is raised, but why? The reason is that the differential equation is now dimensionally inconsistent. The left hand side \( \frac{dv}{dt} \) has units of 1/\( \text{second} \) but the right hand side \( 1-v \) is dimensionless. People often find this behaviour of Brian confusing because this sort of equation is very common in mathematics. However, for quantities with physical dimensions it is incorrect because the results would change depending on the unit you measured it in. For time, if you measured it in seconds the same equation would behave differently to how it would if you measured time in milliseconds. To avoid this, we insist that you always specify dimensionally consistent equations.
Now let’s go back to the good equations and actually run the simulation.

```python
start_scope()

tau = 10*ms
eqs = '''
dv/dt = (1-v)/tau : 1
'''

G = NeuronGroup(1, eqs)
run(100*ms)
```

INFO No numerical integration method specified for group 'neurongroup', using method 'exact' (took 0.05s). [brian2.stateupdaters.base.method_choice]

First off, ignore that `start_scope()` at the top of the cell. You’ll see that in each cell in this tutorial where we run a simulation. All it does is make sure that any Brian objects created before the function is called aren’t included in the next run of the simulation.

Secondly, you’ll see that there is an “INFO” message about not specifying the numerical integration method. This is harmless and just to let you know what method we chose, but we’ll fix it in the next cell by specifying the method explicitly.

So, what has happened here? Well, the command `run(100*ms)` runs the simulation for 100 ms. We can see that this has worked by printing the value of the variable `v` before and after the simulation.

```python
start_scope()

G = NeuronGroup(1, eqs, method='exact')
print('Before v = %s' % G.v[0])
run(100*ms)
print('After v = %s' % G.v[0])
```

```
Before v = 0.0
After v = 0.9999546000702376
```

By default, all variables start with the value 0. Since the differential equation is $\frac{dv}{dt} = \frac{(1-v)}{\tau}$ we would expect after a while that $v$ would tend towards the value 1, which is just what we see. Specifically, we’d expect $v$ to have the value $1 - \exp(-t/\tau)$. Let’s see if that’s right.

```python
print('Expected value of v = %s' % (1-exp(-100*ms/tau)))
```

```
Expected value of v = 0.9999546000702375
```

Good news, the simulation gives the value we’d expect!

Now let’s take a look at a graph of how the variable $v$ evolves over time.

```python
start_scope()

G = NeuronGroup(1, eqs, method='exact')
M = StateMonitor(G, 'v', record=True)
run(30*ms)
plot(M.t/ms, M.v[0])
xlabel('Time (ms)')
ylabel('v');
```
This time we only ran the simulation for 30 ms so that we can see the behaviour better. It looks like it’s behaving as expected, but let’s just check that analytically by plotting the expected behaviour on top.

```python
start_scope()
G = NeuronGroup(1, eqs, method='exact')
M = StateMonitor(G, 'v', record=0)
run(30*ms)
plot(M.t/ms, M.v[0], 'C0', label='Brian')
plot(M.t/ms, 1-exp(-M.t/tau), 'C1--', label='Analytic')
xlabel('Time (ms)')
ylabel('v')
legend();
```
As you can see, the blue (Brian) and dashed orange (analytic solution) lines coincide.

In this example, we used the object StateMonitor object. This is used to record the values of a neuron variable while the simulation runs. The first two arguments are the group to record from, and the variable you want to record from. We also specify record=0. This means that we record all values for neuron 0. We have to specify which neurons we want to record because in large simulations with many neurons it usually uses up too much RAM to record the values of all neurons.

Now try modifying the equations and parameters and see what happens in the cell below.

```python
start_scope()

tau = 10*ms
eqs = ''
dv/dt = (sin(2*pi*100*Hz*t)-v)/tau : 1
''

# Change to Euler method because exact integrator doesn't work here
G = NeuronGroup(1, eqs, method='euler')
M = StateMonitor(G, 'v', record=0)

G.v = 5 # initial value

run(60*ms)

plot(M.t/ms, M.v[0])
xlabel('Time (ms)')
ylabel('v');```
2.1.3 Adding spikes

So far we haven’t done anything neuronal, just played around with differential equations. Now let’s start adding spiking behaviour.

```python
start_scope()

tau = 10*ms
eqs = '''
dv/dt = (1-v)/tau : 1
'''

G = NeuronGroup(1, eqs, threshold='v>0.8', reset='v = 0', method='exact')

M = StateMonitor(G, 'v', record=0)
run(50*ms)
plot(M.t/ms, M.v[0])
xlabel('Time (ms)')
ylabel('v');```

---

2.1. Introduction to Brian part 1: Neurons
We’ve added two new keywords to the NeuronGroup declaration: threshold='v>0.8' and reset='v = 0'. What this means is that when v>0.8 we fire a spike, and immediately reset v = 0 after the spike. We can put any expression and series of statements as these strings.

As you can see, at the beginning the behaviour is the same as before until v crosses the threshold v>0.8 at which point you see it reset to 0. You can’t see it in this figure, but internally Brian has registered this event as a spike. Let’s have a look at that.

```python
start_scope()
G = NeuronGroup(1, eqs, threshold='v>0.8', reset='v = 0', method='exact')
spikemon = SpikeMonitor(G)
run(50*ms)
print('Spike times: %s' % spikemon.t[:])
```

Spike times: [16. 32.1 48.2] ms

The SpikeMonitor object takes the group whose spikes you want to record as its argument and stores the spike times in the variable t. Let’s plot those spikes on top of the other figure to see that it’s getting it right.

```python
start_scope()
G = NeuronGroup(1, eqs, threshold='v>0.8', reset='v = 0', method='exact')
statemon = StateMonitor(G, 'v', record=0)
spikemon = SpikeMonitor(G)
run(50*ms)
plot(statemon.t/ms, statemon.v[0])
```

(continues on next page)
Here we’ve used the \texttt{axvline} command from \texttt{matplotlib} to draw an orange, dashed vertical line at the time of each spike recorded by the \texttt{SpikeMonitor}.

Now try changing the strings for \texttt{threshold} and \texttt{reset} in the cell above to see what happens.

\subsection*{2.1.4 Refractoriness}

A common feature of neuron models is refractoriness. This means that after the neuron fires a spike it becomes refractory for a certain duration and cannot fire another spike until this period is over. Here’s how we do that in Brian.

\begin{verbatim}
start_scope()

tau = 10*ms
eqs = ''
dv/dt = (1-v)/tau : 1 (unless refractory)
''

G = NeuronGroup(1, eqs, threshold='v>0.8', reset='v = 0', refractory=5*ms, method='exact')

statemon = StateMonitor(G, 'v', record=0)
spikemon = SpikeMonitor(G)

run(50*ms)

plot(statemon.t/ms, statemon.v[0])
\end{verbatim}
for t in spikemon.t:
    axvline(t/ms, ls='--', c='C1', lw=3)
xlabel('Time (ms)')
ylabel('v');

As you can see in this figure, after the first spike, $v$ stays at 0 for around 5 ms before it resumes its normal behaviour. To do this, we’ve done two things. Firstly, we’ve added the keyword `refractory=5*ms` to the `NeuronGroup` declaration. On its own, this only means that the neuron cannot spike in this period (see below), but doesn’t change how $v$ behaves. In order to make $v$ stay constant during the refractory period, we have to add `(unless refractory)` to the end of the definition of $v$ in the differential equations. What this means is that the differential equation determines the behaviour of $v$ unless it’s refractory in which case it is switched off.

Here’s what would happen if we didn’t include `(unless refractory)`. Note that we’ve also decreased the value of $\tau$ and increased the length of the refractory period to make the behaviour clearer.

```
start_scope()

tau = 5*ms
eqs = ''
dv/dt = (1-v)/tau : 1
''

G = NeuronGroup(1, eqs, threshold='v>0.8', reset='v = 0', refractory=15*ms, method='exact')

statemon = StateMonitor(G, 'v', record=0)
spikemon = SpikeMonitor(G)

run(50*ms)

plot(statemon.t/ms, statemon.v[0])
```
for t in spikemon.t:
    axvline(t/ms, ls='--', c='C1', lw=3)
axhline(0.8, ls=':', c='C2', lw=3)
xlabel('Time (ms)')
ylabel('v')
print("Spike times: \\n" % spikemon.t[:])

Spike times: [ 8. 23. 38.] ms

So what’s going on here? The behaviour for the first spike is the same: v rises to 0.8 and then the neuron fires a spike at time 8 ms before immediately resetting to 0. Since the refractory period is now 15 ms this means that the neuron won’t be able to spike again until time 8 + 15 = 23 ms. Immediately after the first spike, the value of v now instantly starts to rise because we didn’t specify (unless refractory) in the definition of dv/dt. However, once it reaches the value 0.8 (the dashed green line) at time roughly 8 ms it doesn’t fire a spike even though the threshold is v>0.8. This is because the neuron is still refractory until time 23 ms, at which point it fires a spike.

Note that you can do more complicated and interesting things with refractoriness. See the full documentation for more details about how it works.

### 2.1.5 Multiple neurons

So far we’ve only been working with a single neuron. Let’s do something interesting with multiple neurons.

start_scope()
N = 100
tau = 10*ms
eqs = ''
\[ \frac{dv}{dt} = \frac{2-v}{\tau} \]

(continues on next page)
This shows a few changes. Firstly, we’ve got a new variable \( N \) determining the number of neurons. Secondly, we added the statement \( G.v = \text{’rand()} \) before the run. What this does is initialise each neuron with a different uniform random value between 0 and 1. We’ve done this just so each neuron will do something a bit different. The other big change is how we plot the data in the end.

As well as the variable \( \text{spikemon.t} \) with the times of all the spikes, we’ve also used the variable \( \text{spikemon.i} \) which gives the corresponding neuron index for each spike, and plotted a single black dot with time on the x-axis and neuron index on the y-value. This is the standard “raster plot” used in neuroscience.

### 2.1.6 Parameters

To make these multiple neurons do something more interesting, let’s introduce per-neuron parameters that don’t have a differential equation attached to them.

```python
start_scope()
N = 100
tau = 10*ms
```
v0_max = 3.
duration = 1000*ms

eqs = ''
dv/dt = (v0-v)/tau : 1 (unless refractory)
v0 : 1
''

G = NeuronGroup(N, eqs, threshold='v>1', reset='v=0', refractory=5*ms, method='exact')
M = SpikeMonitor(G)

G.v0 = 'i*v0_max/(N-1)'
run(duration)

figure(figsize=(12,4))
s subplot(121)
plot(M.t/ms, M.i, '.k')
xlabel('Time (ms)')
ylabel('Neuron index')
s subplot(122)
plot(G.v0, M.count/duration)
xlabel('v0')
ylabel('Firing rate (sp/s)');

The line v0 : 1 declares a new per-neuron parameter v0 with units 1 (i.e. dimensionless).
The line G.v0 = 'i*v0_max/(N-1)' initialises the value of v0 for each neuron varying from 0 up to v0_max.
The symbol i when it appears in strings like this refers to the neuron index.

So in this example, we’re driving the neuron towards the value v0 exponentially, but when v crosses v>1, it fires a spike and resets. The effect is that the rate at which it fires spikes will be related to the value of v0. For v0<1 it will never fire a spike, and as v0 gets larger it will fire spikes at a higher rate. The right hand plot shows the firing rate as a function of the value of v0. This is the I-f curve of this neuron model.

Note that in the plot we’ve used the count variable of the SpikeMonitor: this is an array of the number of spikes each neuron in the group fired. Dividing this by the duration of the run gives the firing rate.
2.1.7 Stochastic neurons

Often when making models of neurons, we include a random element to model the effect of various forms of neural noise. In Brian, we can do this by using the symbol $xi$ in differential equations. Strictly speaking, this symbol is a “stochastic differential” but you can sort of thinking of it as just a Gaussian random variable with mean 0 and standard deviation 1. We do have to take into account the way stochastic differentials scale with time, which is why we multiply it by $\tau^{-0.5}$ in the equations below (see a textbook on stochastic differential equations for more details). Note that we also changed the method keyword argument to use 'euler' (which stands for the Euler-Maruyama method); the 'exact' method that we used earlier is not applicable to stochastic differential equations.

```python
start_scope()

N = 100
tau = 10*ms
v0_max = 3.
duration = 1000*ms
sigma = 0.2

eqs = '''
dv/dt = (v0-v)/tau+sigma*xi*tau**-0.5 : 1 (unless refractory)
v0 : 1
'''
G = NeuronGroup(N, eqs, threshold='v>1', reset='v=0', refractory=5*ms, method='euler')
M = SpikeMonitor(G)
G.v0 = 'i*v0_max/(N-1)'
run(duration)

figure(figsize=(12,4))
subplot(121)
plot(M.t/ms, M.i, '.k')
xlabel('Time (ms)')
ylabel('Neuron index')
subplot(122)
plot(G.v0, M.count/duration)
xlabel('v0')
ylabel('Firing rate (sp/s)');
```

That’s the same figure as in the previous section but with some noise added. Note how the curve has changed shape: instead of a sharp jump from firing at rate 0 to firing at a positive rate, it now increases in a sigmoidal fashion. This is
because no matter how small the driving force the randomness may cause it to fire a spike.

2.1.8 End of tutorial

That’s the end of this part of the tutorial. The cell below has another example. See if you can work out what it is doing and why. Try adding a StateMonitor to record the values of the variables for one of the neurons to help you understand it.

You could also try out the things you’ve learned in this cell.

Once you’re done with that you can move on to the next tutorial on Synapses.

```python
start_scope()
N = 1000
tau = 10*ms
vr = -70*mV
vt0 = -50*mV
delta_vt0 = 5*mV
tau_t = 100*ms
sigma = 0.5*(vt0-vr)
v_drive = 2*(vt0-vr)
duration = 100*ms

eqs = '"
dv/dt = (v_drive+vr-v)/tau + sigma*xn*tau**-0.5 : volt
dvt/dt = (vt0-vt)/tau_t : volt
"

reset = '"
v = vr
vt += delta_vt0
"

G = NeuronGroup(N, eqs, threshold='v>vt', reset=reset, refractory=5*ms, method='euler')
spikemon = SpikeMonitor(G)
G.v = 'rand()*(vt0-vr)+vr'
G.vt = vt0
run(duration)
_ = hist(spikemon.t/ms, 100, histtype='stepfilled', facecolor='k', weights=ones(len(spikemon))/(N*defaultclock.dt))
xlabel('Time (ms)')
ylabel('Instantaneous firing rate (sp/s)');
```
2.2 Introduction to Brian part 2: Synapses

If you haven’t yet read part 1: Neurons, go read that now.

As before we start by importing the Brian package and setting up matplotlib for IPython:

```python
from brian2 import *
%matplotlib inline
```

2.2.1 The simplest Synapse

Once you have some neurons, the next step is to connect them up via synapses. We’ll start out with doing the simplest possible type of synapse that causes an instantaneous change in a variable after a spike.

```python
start_scope()

eqs = '''
dv/dt = (I-v)/tau : 1
I : 1
tau : second
'''
G = NeuronGroup(2, eqs, threshold='v>1', reset='v = 0', method='exact')
G.I = [2, 0]
G.tau = [10, 100]*ms

# Comment these two lines out to see what happens without Synapses
S = Synapses(G, G, on_pre='v_post += 0.2')
S.connect(i=0, j=1)
```

(continues on next page)
There are a few things going on here. First of all, let’s recap what is going on with the NeuronGroup. We’ve created two neurons, each of which has the same differential equation but different values for parameters I and tau. Neuron 0 has $I=2$ and $\tau=10\,\text{ms}$ which means that is driven to repeatedly spike at a fairly high rate. Neuron 1 has $I=0$ and $\tau=100\,\text{ms}$ which means that on its own - without the synapses - it won’t spike at all (the driving current I is 0). You can prove this to yourself by commenting out the two lines that define the synapse.

Next we define the synapses:

```
Synapses(source, target, ...) means that we are defining a synaptic model that goes from source to target. In this case, the source and target are both the same, the group G. The syntax `on_pre='v_post += 0.2'` means that when a spike occurs in the presynaptic neuron (hence `on_pre`) it causes an instantaneous change to happen $v_post += 0.2$. The `_post` means that the value of $v$ referred to is the post-synaptic value, and it is increased by 0.2. So in total, what this model says is that whenever two neurons in G are connected by a synapse, when the source neuron fires a spike the target neuron will have its value of $v$ increased by 0.2.
```

However, at this point we have only defined the synapse model, we haven’t actually created any synapses. The next line `S.connect(i=0, j=1)` creates a synapse from neuron 0 to neuron 1.

### 2.2.2 Adding a weight

In the previous section, we hard coded the weight of the synapse to be the value 0.2, but often we would to allow this to be different for different synapses. We do that by introducing synapse equations.
This example behaves very similarly to the previous example, but now there’s a synaptic weight variable \( w \). The string \\
'w : 1' is an equation string, precisely the same as for neurons, that defines a single dimensionless parameter \( w \). We changed the behaviour on a spike to \\
on_pre='v_post += w' now, so that each synapse can behave differently depending on the value of \( w \). To illustrate this, we’ve made a third neuron which behaves precisely the same as the second neuron, and connected neuron 0 to both neurons 1 and 2. We’ve also set the weights via \( S.w = 'j*0.2' \). When \( i \) and \( j \) occur in the context of synapses, \( i \) refers to the source neuron index, and \( j \) to the target index.
neuron index. So this will give a synaptic connection from 0 to 1 with weight $0.2 = 0.2 \times 1$ and from 0 to 2 with weight $0.4 = 0.2 \times 2$.

### 2.2.3 Introducing a delay

So far, the synapses have been instantaneous, but we can also make them act with a certain delay.

```python
start_scope()

eqs = ''
dv/dt = (I-v)/tau : 1
I : 1
tau : second
''
G = NeuronGroup(3, eqs, threshold='v>1', reset='v = 0', method='exact')
G.I = [2, 0, 0]
G.tau = [10, 100, 100]*ms

S = Synapses(G, G, 'w : 1', on_pre='v_post += w')
S.connect(i=0, j=[1, 2])
S.w = 'j*0.2'
S.delay = 'j*2*ms'

M = StateMonitor(G, 'v', record=True)

run(50*ms)

plot(M.t/ms, M.v[0], label='Neuron 0')
plot(M.t/ms, M.v[1], label='Neuron 1')
plot(M.t/ms, M.v[2], label='Neuron 2')
xlabel('Time (ms)')
ylabel('v')
legend();
```
As you can see, that’s as simple as adding a line `s.delay = 'j*2*ms'` so that the synapse from 0 to 1 has a delay of 2 ms, and from 0 to 2 has a delay of 4 ms.

### 2.2.4 More complex connectivity

So far, we specified the synaptic connectivity explicitly, but for larger networks this isn’t usually possible. For that, we usually want to specify some condition.

```python
start_scope()
N = 10
G = NeuronGroup(N, 'v:1')
S = Synapses(G, G)
S.connect(condition='i!=j', p=0.2)
```

Here we’ve created a dummy neuron group of N neurons and a dummy synapses model that doesn’t actually do anything just to demonstrate the connectivity. The line `S.connect(condition='i!=j', p=0.2)` will connect all pairs of neurons i and j with probability 0.2 as long as the condition `i!=j` holds. So, how can we see that connectivity? Here’s a little function that will let us visualise it.

```python
def visualise_connectivity(S):
    Ns = len(S.source)
    Nt = len(S.target)
    figure(figsize=(10, 4))
    subplot(121)
    plot(zeros(Ns), arange(Ns), 'ok', ms=10)
    plot(ones(Nt), arange(Nt), 'ok', ms=10)
    for i, j in zip(S.i, S.j):
        plot([0, 1], [i, j], '-k')
    xticks([0, 1], ['Source', 'Target'])
ylabel('Neuron index')
```

(continues on next page)
There are two plots here. On the left hand side, you see a vertical line of circles indicating source neurons on the left, and a vertical line indicating target neurons on the right, and a line between two neurons that have a synapse. On the right hand side is another way of visualising the same thing. Here each black dot is a synapse, with x value the source neuron index, and y value the target neuron index.

Let’s see how these figures change as we change the probability of a connection:

```python
start_scope()
N = 10
G = NeuronGroup(N, 'v:1')
for p in [0.1, 0.5, 1.0]:
    S = Synapses(G, G)
    S.connect(condition='i!=j', p=p)
    visualise_connectivity(S)
suptitle('p = ' + str(p))
```
And let’s see what another connectivity condition looks like. This one will only connect neighbouring neurons.

```python
start_scope()

N = 10
G = NeuronGroup(N, 'v:1')
S = Synapses(G, G)
S.connect(condition='abs(i-j)<4 and i!=j')
visualise_connectivity(S)
```

Try using that cell to see how other connectivity conditions look like.

You can also use the generator syntax to create connections like this more efficiently. In small examples like this, it doesn’t matter, but for large numbers of neurons it can be much more efficient to specify directly which neurons should be connected than to specify just a condition. Note that the following example uses `skip_if_invalid` to avoid errors at the boundaries (e.g. do not try to connect the neuron with index 1 to a neuron with index -2).
If each source neuron is connected to precisely one target neuron (which would be normally used with two separate groups of the same size, not with identical source and target groups as in this example), there is a special syntax that is extremely efficient. For example, 1-to-1 connectivity looks like this:

```python
s = Synapses(G, G)
s.connect(j='i')
visualise_connectivity(s)
```
You can also do things like specifying the value of weights with a string. Let’s see an example where we assign each neuron a spatial location and have a distance-dependent connectivity function. We visualise the weight of a synapse by the size of the marker.

```python
start_scope()

N = 30
euron_spacing = 50*umetre
width = N/4.0*neuron_spacing

# Neuron has one variable x, its position
G = NeuronGroup(N, 'x : metre')
G.x = 'i*neuron_spacing'

# All synapses are connected (excluding self-connections)
S = Synapses(G, G, 'w : 1')
S.connect(condition='i!=j')

# Weight varies with distance
S.w = 'exp(-(x_pre-x_post)**2/(2*width**2))'

scatter(S.x_pre/um, S.x_post/um, S.w*20)
xlabel('Source neuron position (um)')
ylabel('Target neuron position (um)');
```
Now try changing that function and seeing how the plot changes.

### 2.2.5 More complex synapse models: STDP

Brian’s synapse framework is very general and can do things like short-term plasticity (STP) or spike-timing dependent plasticity (STDP). Let’s see how that works for STDP.

STDP is normally defined by an equation something like this:

\[
\Delta w = \sum_{t_{\text{pre}}} \sum_{t_{\text{post}}} W(t_{\text{post}} - t_{\text{pre}})
\]

That is, the change in synaptic weight \( w \) is the sum over all presynaptic spike times \( t_{\text{pre}} \) and postsynaptic spike times \( t_{\text{post}} \) of some function \( W \) of the difference in these spike times. A commonly used function \( W \) is:

\[
W(\Delta t) = \begin{cases} 
A_{\text{pre}}e^{-\Delta t/\tau_{\text{pre}}} & \Delta t > 0 \\
A_{\text{post}}e^{\Delta t/\tau_{\text{post}}} & \Delta t < 0 
\end{cases}
\]

This function looks like this:

```python
tau_pre = tau_post = 20*ms
A_pre = 0.01
A_post = -A_pre*1.05
delta_t = linspace(-50, 50, 100)*ms
W = where(delta_t>0, A_pre*exp(-delta_t/tau_pre), A_post*exp(delta_t/tau_post))
plot(delta_t/ms, W)
xlabel(r'$\Delta t$ (ms)')
ylabel('W')
axhline(0, ls='-', c='k');```

![Graph showing synaptic weight changes](image.png)
Simulating it directly using this equation though would be very inefficient, because we would have to sum over all pairs of spikes. That would also be physiologically unrealistic because the neuron cannot remember all its previous spike times. It turns out there is a more efficient and physiologically more plausible way to get the same effect.

We define two new variables $a_{\text{pre}}$ and $a_{\text{post}}$ which are “traces” of pre- and post-synaptic activity, governed by the differential equations:

\[
\tau_{\text{pre}} \frac{da_{\text{pre}}}{dt} = -a_{\text{pre}} \tag{2.1}
\]

\[
\tau_{\text{post}} \frac{da_{\text{post}}}{dt} = -a_{\text{post}} \tag{2.2}
\]

When a presynaptic spike occurs, the presynaptic trace is updated and the weight is modified according to the rule:

\[
a_{\text{pre}} \rightarrow a_{\text{pre}} + A_{\text{pre}} \quad w \rightarrow w + (\tau_{\text{post}} A_{\text{pre}}) \tag{2.3}
\]

When a postsynaptic spike occurs:

\[
a_{\text{post}} \rightarrow a_{\text{post}} + A_{\text{post}} \quad w \rightarrow w + (\tau_{\text{pre}} A_{\text{post}}) \tag{2.5}
\]

To see that this formulation is equivalent, you just have to check that the equations sum linearly, and consider two cases: what happens if the presynaptic spike occurs before the postsynaptic spike, and vice versa. Try drawing a picture of it.

Now that we have a formulation that relies only on differential equations and spike events, we can turn that into Brian code.

```brian2
code
start_scope()

taupre = taupost = 20*ms
wmax = 0.01
```

(continues on next page)
There are a few things to see there. Firstly, when defining the synapses we’ve given a more complicated multi-line string defining three synaptic variables ($w$, $apre$ and $apost$). We’ve also got a new bit of syntax there, \(\text{(event-driven)}\) after the definitions of $apre$ and $apost$. What this means is that although these two variables evolve continuously over time, Brian should only update them at the time of an event (a spike). This is because we don’t need the values of $apre$ and $apost$ except at spike times, and it is more efficient to only update them when needed.

Next we have a \texttt{on\_pre=...} argument. The first line is $v\_post += w$: this is the line that actually applies the synaptic weight to the target neuron. The second line is $apre += Apre$ which encodes the rule above. In the third line, we’re also encoding the rule above but we’ve added one extra feature: we’ve clamped the synaptic weights between a minimum of 0 and a maximum of $wmax$ so that the weights can’t get too large or negative. The function $\text{clip}(x, \text{low}, \text{high})$ does this.

Finally, we have a \texttt{on\_post=...} argument. This gives the statements to calculate when a post-synaptic neuron fires. Note that we do not modify $v$ in this case, only the synaptic variables.

Now let’s see how all the variables behave when a presynaptic spike arrives some time before a postsynaptic spike.
```python
w = clip(w+apost, 0, wmax)
'''
on_post=''
apost += Apost
w = clip(w+apre, 0, wmax)
''', method='linear')
S.connect(i=0, j=1)
M = StateMonitor(S, ['w', 'apre', 'apost'], record=True)
run(30*ms)
figure(figsize=(4, 8))
subplot(211)
plot(M.t/ms, M.apre[0], label='apre')
plot(M.t/ms, M.apost[0], label='apost')
legend()
subplot(212)
plot(M.t/ms, M.w[0], label='w')
legend(loc='best')
xlabel('Time (ms)');```
A couple of things to note here. First of all, we’ve used a trick to make neuron 0 fire a spike at time 10 ms, and neuron 1 at time 20 ms. Can you see how that works?

Secondly, we’ve replaced the \textit{(event-driven)} by \textit{(clock-driven)} so you can see how $a_{pre}$ and $a_{post}$ evolve over time. Try reverting this change and see what happens.

Try changing the times of the spikes to see what happens.

Finally, let’s verify that this formulation is equivalent to the original one.

```python
start_scope()

taupre = taupost = 20*ms
Apre = 0.01
```

(continues on next page)
Apost = -Apre*taupre/taupost*1.05

tmax = 50*ms

N = 100

# Presynaptic neurons G spike at times from 0 to tmax
# Postsynaptic neurons G spike at times from tmax to 0
# So difference in spike times will vary from -tmax to +tmax
G = NeuronGroup(N, 'tspike:second', threshold='t>tspike', refractory=100*ms)
H = NeuronGroup(N, 'tspike:second', threshold='t>tspike', refractory=100*ms)

G.tspike = 'i*tmax/(N-1)'
H.tspike = '((N-1-i)*tmax/(N-1)'

S = Synapses(G, H,
            '''
            w : 1
            dapre/dt = -apre/taupre : 1 (event-driven)
            dapost/dt = -apost/taupost : 1 (event-driven)
            ''',
            on_pre='''
            apre += Apre
            w = w+apost
            ''',
            on_post='''
            apost += Apost
            w = w+apre
            ''')

S.connect(j='i')

run(tmax+1*ms)

plot((H.tspike-G.tspike)/ms, S.w)
xlabel(r'$\Delta t$ (ms)')
ylabel(r'$\Delta w$')
axhline(0, ls='-', c='k');
Can you see how this works?

### 2.6 End of tutorial

### 2.3 Introduction to Brian part 3: Simulations

If you haven’t yet read parts 1 and 2 on Neurons and Synapses, go read them first.

This tutorial is about managing the slightly more complicated tasks that crop up in research problems, rather than the toy examples we’ve been looking at so far. So we cover things like inputting sensory data, modelling experimental conditions, etc.

As before we start by importing the Brian package and setting up matplotlib for IPython:

```python
from brian2 import *
%matplotlib inline
```

#### 2.3.1 Multiple runs

Let’s start by looking at a very common task: doing multiple runs of a simulation with some parameter that changes. Let’s start off with something very simple, how does the firing rate of a leaky integrate-and-fire neuron driven by Poisson spiking neurons change depending on its membrane time constant? Let’s set that up.

```python
# remember, this is here for running separate simulations in the same notebook
start_scope()
# Parameters
num_inputs = 100
input_rate = 10*Hz
weight = 0.1
```
# Range of time constants

tau_range = linspace(1, 10, 30)*ms

# Use this list to store output rates
output_rates = []

# Iterate over range of time constants
for tau in tau_range:
    # Construct the network each time
    P = PoissonGroup(num_inputs, rates=input_rate)
    eqs = '''
dv/dt = -v/tau : 1
'''
    G = NeuronGroup(1, eqs, threshold='v>1', reset='v=0', method='exact')
    S = Synapses(P, G, on_pre='v += weight')
    S.connect()
    M = SpikeMonitor(G)
    # Run it and store the output firing rate in the list
    run(1*second)
    output_rates.append(M.num_spikes/second)

# And plot it
plot(tau_range/ms, output_rates)
xlabel(r'$\tau$ (ms)')
ylabel('Firing rate (sp/s)');

Now if you’re running the notebook, you’ll see that this was a little slow to run. The reason is that for each loop, you’re recreating the objects from scratch. We can improve that by setting up the network just once. We store a copy of the state of the network before the loop, and restore it at the beginning of each iteration.

start_scope()
num_inputs = 100
input_rate = 10*Hz
weight = 0.1

(continues on next page)
tau_range = linspace(1, 10, 30)*ms
output_rates = []
# Construct the network just once
P = PoissonGroup(num_inputs, rates=input_rate)
eqs = ""
   dv/dt = -v/tau : 1
   
G = NeuronGroup(1, eqs, threshold='v>1', reset='v=0', method='exact')
S = Synapses(P, G, on_pre='v += weight')
S.connect()
M = SpikeMonitor(G)
# Store the current state of the network
store()
for tau in tau_range:
   # Restore the original state of the network
   restore()
   # Run it with the new value of tau
   run(1*second)
   output_rates.append(M.num_spikes/second)
plot(tau_range/ms, output_rates)
xlabel(r'$\tau$ (ms)')
ylabel('Firing rate (sp/s)');

That's a very simple example of using store and restore, but you can use it in much more complicated situations. For example, you might want to run a long training run, and then run multiple test runs afterwards. Simply put a store after the long training run, and a restore before each testing run.

You can also see that the output curve is very noisy and doesn’t increase monotonically like we’d expect. The noise is coming from the fact that we run the Poisson group afresh each time. If we only wanted to see the effect of the time constant, we could make sure that the spikes were the same each time (although note that really, you ought to do multiple runs and take an average). We do this by running just the Poisson group once, recording its spikes, and then creating a new SpikeGeneratorGroup that will output those recorded spikes each time.
start_scope()
num_inputs = 100
input_rate = 10*Hz
weight = 0.1
tau_range = linspace(1, 10, 30)*ms
output_rates = []
# Construct the Poisson spikes just once
P = PoissonGroup(num_inputs, rates=input_rate)
MP = SpikeMonitor(P)
# We use a Network object because later on we don't
# want to include these objects
net = Network(P, MP)
net.run(1*second)
# And keep a copy of those spikes
spikes_i = MP.i
spikes_t = MP.t
# Now construct the network that we run each time
# SpikeGeneratorGroup gets the spikes that we created before
SGG = SpikeGeneratorGroup(num_inputs, spikes_i, spikes_t)
eqs = '
    dv/dt = -v/tau : 1
    
G = NeuronGroup(1, eqs, threshold='v>1', reset='v=0', method='exact')
S = Synapses(SGG, G, on_pre='v += weight')
S.connect()
M = SpikeMonitor(G)
# Store the current state of the network
net = Network(SGG, G, S, M)
net.store()
for tau in tau_range:
    # Restore the original state of the network
    net.restore()
    # Run it with the new value of tau
    net.run(1*second)
    output_rates.append(M.num_spikes/second)
plot(tau_range/ms, output_rates)
xlabel(r'$\tau$ (ms)')
ylabel('Firing rate (sp/s)');
You can see that now there is much less noise and it increases monotonically because the input spikes are the same each time, meaning we’re seeing the effect of the time constant, not the random spikes.

Note that in the code above, we created Network objects. The reason is that in the loop, if we just called run it would try to simulate all the objects, including the Poisson neurons \( P \), and we only want to run that once. We use Network to specify explicitly which objects we want to include.

The techniques we’ve looked at so far are the conceptually most simple way to do multiple runs, but not always the most efficient. Since there’s only a single output neuron in the model above, we can simply duplicate that output neuron and make the time constant a parameter of the group.

```python
start_scope()
num_inputs = 100
input_rate = 10*Hz
weight = 0.1
tau_range = linspace(1, 10, 30)*ms
num_tau = len(tau_range)
P = PoissonGroup(num_inputs, rates=input_rate)
# We make tau a parameter of the group
eqs = '
    dv/dt = -v/tau : 1
    tau : second
''
# And we have num_tau output neurons, each with a different tau
G = NeuronGroup(num_tau, eqs, threshold='v>1', reset='v=0', method='exact')
G.tau = tau_range
S = Synapses(P, G, on_pre='v += weight')
S.connect()
M = SpikeMonitor(G)
# Now we can just run once with no loop
run(1*second)
output_rates = M.count/second  # firing rate is count/duration
plot(tau_range/ms, output_rates)
```

(continues on next page)
You can see that this is much faster again! It’s a little bit more complicated conceptually, and it’s not always possible to do this trick, but it can be much more efficient if it’s possible.

Let’s finish with this example by having a quick look at how the mean and standard deviation of the interspike intervals depends on the time constant.

```python
trains = M.spike_trains()
isi_mu = full(num_tau, nan)*second
isi_std = full(num_tau, nan)*second
for idx in range(num_tau):
    train = diff(trains[idx])
    if len(train)>1:
        isi_mu[idx] = mean(train)
        isi_std[idx] = std(train)
errorbar(tau_range/ms, isi_mu/ms, yerr=isi_std/ms)
xlabel(r'$\tau$ (ms)')
ylabel('Interspike interval (ms)');
```
Notice that we used the `spike_trains()` method of `SpikeMonitor`. This is a dictionary with keys being the indices of the neurons and values being the array of spike times for that neuron.

### 2.3.2 Changing things during a run

Imagine an experiment where you inject current into a neuron, and change the amplitude randomly every 10 ms. Let’s see if we can model that using a Hodgkin-Huxley type neuron.

```python
start_scope()

# Parameters
area = 20000*umetre**2
Cm = 1*ufarad*cm**-2 * area
gl = 5e-5*siemens*cm**-2 * area
E_l = -65*mV
E_K = -90*mV
E_Na = 50*mV
g_Na = 100*msiemens*cm**-2 * area
g_Kd = 30*msiemens*cm**-2 * area
V_T = -63*mV

# The model
eqs_HH = '''
dv/dt = (gl*(E_l-v) - g_Na*(m*m*m)*h*(v-E_Na) - g_Kd*(n*n*n*n)*(v-E_K) + I)/Cm : volt
dm/dt = 0.32*(mV**-1)*(13.*mV-v+VT)/
      (exp((13.*mV-v+VT)/(4.*mV)) - 1.)/ms*(1-m)-0.28*(mV**-1)*(v-V_T-40.*mV)/
      (exp((v-V_T-40.*mV)/(5.*mV))) - 1)/ms*m : 1
dn/dt = 0.032*(mV**-1)*(15.*mV-v+VT)/
      (exp((15.*mV-v+VT)/(5.*mV)) - 1.)/ms*(1-n)-5*exp((10.*mV-v+VT)/(40.*mV))/ms*n : 1
dh/dt = 0.128*exp((17.*mV-v+VT)/(18.*mV))/ms*(1-h)-4./(1+exp((40.*mV-v+VT)/(5.*mV)))/ms*h : 1
I : amp'''
```

(continues on next page)
In the code above, we used a loop over multiple runs to achieve this. That’s fine, but it’s not the most efficient way to do it because each time we call `run` we have to do a lot of initialisation work that slows everything down. It also won’t work as well with the more efficient standalone mode of Brian. Here’s another way.

```python
start_scope()
group = NeuronGroup(1, eqs_HH,
    threshold='v > -40*mV',
    refractory='v > -40*mV',
    method='exponential_euler')
group.v = El
statemon = StateMonitor(group, 'v', record=True)
spikemon = SpikeMonitor(group, variables='v')
figure(figsize=(9, 4))
    for l in range(5):
        group.I = rand()*50*nA
        run(10*ms)
        axvline(l*10, ls='--', c='k')
axhline(El/mV, ls='-', c='lightgray', lw=3)
plot(statemon.t/ms, statemon.v[0]/mV, '-b')
plot(spikemon.t/ms, spikemon.v/mV, 'ob')
xlabel('Time (ms)')
ylabel('v (mV)');
```

In the code above, we used a loop over multiple runs to achieve this. That’s fine, but it’s not the most efficient way to do it because each time we call `run` we have to do a lot of initialisation work that slows everything down. It also won’t work as well with the more efficient standalone mode of Brian. Here’s another way.

```python
start_scope()
group = NeuronGroup(1, eqs_HH,
    threshold='v > -40*mV',
    refractory='v > -40*mV',
    method='exponential_euler')
group.v = El
statemon = StateMonitor(group, 'v', record=True)
spikemon = SpikeMonitor(group, variables='v')
    # we replace the loop with a `run_regularly`
group.run_regularly('I = rand()*50*nA', dt=10*ms)
run(50*ms)
figure(figsize=(9, 4))
    # we keep the loop just to draw the vertical lines
    for l in range(5):
```

(continues on next page)
We’ve replaced the loop that had multiple run calls with a run_regularly. This makes the specified block of code run every dt=10*ms. The run_regularly lets you run code specific to a single NeuronGroup, but sometimes you might need more flexibility. For this, you can use network_operation which lets you run arbitrary Python code (but won’t work with the standalone mode).

```python
start_scope()
group = NeuronGroup(1, eqs_HH,
    threshold='v > -40*mV',
    refractory='v > -40*mV',
    method='exponential_euler')
group.v = El
statemon = StateMonitor(group, 'v', record=True)
spikemon = SpikeMonitor(group, variables='v')
# we replace the loop with a network_operation
@network_operation(dt=10*ms)
def change_I():
    group.I = rand()*50*nA
run(50*ms)
figure(figsize=(9, 4))
for l in range(5):
    axvline(l*10, ls='--', c='k')
axhline(El/mV, ls='-', c='lightgray', lw=3)
plot(statemon.t/ms, statemon.v[0]/mV, '-b')
plot(spikemon.t/ms, spikemon.v/mV, 'ob')
xlabel('Time (ms)')
ylabel('v (mV)');
```
Now let’s extend this example to run on multiple neurons, each with a different capacitance to see how that affects the behaviour of the cell.

```python
def start_scope():
    N = 3
    eqs_HH_2 = '''
    dv/dt = (gl*(El-v) - g_na*(m**3)*h*(v-ENa) - g_kd*(n**4)*(v-EK) + I)/C : volt
    dm/dt = 0.32*(mV**-1)*(13.*mV-v+VT)/
        (exp((13.*mV-v+VT)/(4.*mV))-1.)/ms*(1.-m)-0.28*(mV**-1)*(v-VT-40.*mV)/
        (exp((v-VT-40.*mV)/(5.*mV))-1.)/ms*m : 1
    dn/dt = 0.032*(mV**-1)*(15.*mV-v+VT)/
        (exp((15.*mV-v+VT)/(5.*mV))-1.)/ms*(1.-n)-.5*exp((10.*mV-v+VT)/(40.*mV))/ms*n : 1
    dh/dt = 0.128*exp((17.*mV-v+VT)/(18.*mV))/ms*(1.-h)-4./(1+exp((40.*mV-v+VT)/(5.*mV)))/
        ms*h : 1
    I : amp
    C : farad
    '''
    group = NeuronGroup(N, eqs_HH_2,
        threshold='v > -40*mV',
        refractory='v > -40*mV',
        method='exponential_euler')
    group.v = El
    # initialise with some different capacitances
    group.C = array([0.8, 1, 1.2])*ufarad*cm**-2*area
    statemon = StateMonitor(group, variables=True, record=True)
    # we go back to run_regularly
    group.run_regularly('I = rand()*50*nA', dt=10*ms)
    run(50*ms)
    figure(figsize=(9, 4))
    for l in range(5):
        axvline(l*10, ls='--', c='k')
        axhline(El/mV, ls='-', c='lightgray', lw=3)
        plot(statemon.t/ms, statemon.v.T/mV, '-')
    xlabel('Time (ms)')
    ylabel('v (mV)');
```

2.3. Introduction to Brian part 3: Simulations
So that runs, but something looks wrong! The injected currents look like they’re different for all the different neurons!
Let’s check:

```python
plot(statemon.t/ms, statemon.I.T/nA, '-')
xlabel('Time (ms)')
ylabel('I (nA)');
```

Sure enough, it’s different each time. But why? We wrote `group.run_regularly('I = rand()*50*nA', dt=10*ms)` which seems like it should give the same value of I for each neuron. But, like threshold and reset statements, `run_regularly` code is interpreted as being run separately for each neuron, and because I is a parameter, it can be different for each neuron. We can fix this by making I into a shared variable, meaning it has the same value for each neuron.
start_scope()
N = 3
eqs_HH_3 = '''
dv/dt = (gl*(El-v) - g_na*(m*m*m)*h*(v-ENa) - g_kd*(n*n*n*n)*(v-EK) + I)/C : volt
dm/dt = 0.32*(mV**-1)*(13.*mV-v+VT)/
    (exp((13.*mV-v+VT)/(4.*mV))-1.)/ms*(1-m)-0.28*(mV**-1)*(v-VT-40.*mV)/
    (exp((v-VT-40.*mV)/(5.*mV))-1.)/ms*m : 1
dn/dt = 0.032*(mV**-1)*(15.*mV-v+VT)/
    (exp((15.*mV-v+VT)/(5.*mV))-1.)/ms*(1.-n)-.5*exp((10.*mV-v+VT)/(40.*mV))/ms*n : 1
dh/dt = 0.128*exp((17.*mV-v+VT)/(18.*mV))/ms*(1.-h)-4./(1+exp((40.*mV-v+VT)/(5.*mV)))/
    ms*h : 1
I : amp (shared) # everything is the same except we've added this shared
C : farad
'''

group = NeuronGroup(N, eqs_HH_3,
    threshold='v > -40*mV',
    refractory='v > -40*mV',
    method='exponential_euler')
group.v = El
group.C = array([0.8, 1, 1.2])*ufarad*cm**-2*area
statemon = StateMonitor(group, 'v', record=True)
group.run_regularly('I = rand()*50*nA', dt=10*ms)
run(50*ms)

figure(figsize=(9, 4))
for l in range(5):
    axvline(l*10, ls='--', c='k')
    axhline(El/mV, ls='-', c='lightgray', lw=3)
plot(statemon.t/ms, statemon.v.T/mV, '-')
xlabel('Time (ms)')
ylabel('v (mV)');

Ahh, that’s more like it!
2.3.3 Adding input

Now let’s think about a neuron being driven by a sinusoidal input. Let’s go back to a leaky integrate-and-fire to simplify the equations a bit.

```python
start_scope()
A = 2.5
f = 10*Hz
tau = 5*ms
eqs = ''
dv/dt = (I-v)/tau : 1
I = A*sin(2*pi*f*t) : 1
''
G = NeuronGroup(1, eqs, threshold='v>1', reset='v=0', method='euler')
M = StateMonitor(G, variables=True, record=True)
run(200*ms)
plot(M.t/ms, M.v[0], label='v')
plot(M.t/ms, M.I[0], label='I')
xlabel('Time (ms)')
ylabel('v')
legend(loc='best');
```

So far, so good and the sort of thing we saw in the first tutorial. Now, what if that input current were something we had recorded and saved in a file? In that case, we can use TimedArray. Let’s start by reproducing the picture above but using TimedArray.

```python
start_scope()
A = 2.5
f = 10*Hz
tau = 5*ms
# Create a TimedArray and set the equations to use it
t_recorded = arange(int(200*ms/defaultclock.dt))*defaultclock.dt
I_recorded = TimedArray(A*sin(2*pi*f*t_recorded), dt=defaultclock.dt)
```
eqs = '''
  dv/dt = (I-v)/tau : 1
  I = I_recorded(t) : 1
'''

G = NeuronGroup(1, eqs, threshold='v>1', reset='v=0', method='exact')
M = StateMonitor(G, variables=True, record=True)
run(200*ms)
plot(M.t/ms, M.v[0], label='v')
plot(M.t/ms, M.I[0], label='I')
xlabel('Time (ms)')
ylabel('v')
legend(loc='best');

Note that for the example where we put the sin function directly in the equations, we had to use the method='euler' argument because the exact integrator wouldn't work here (try it!). However, TimedArray is considered to be constant over its time step and so the linear integrator can be used. This means you won’t get the same behaviour from these two methods for two reasons. Firstly, the numerical integration methods exact and euler give slightly different results. Secondly, sin is not constant over a timestep whereas TimedArray is.

Now just to show that TimedArray works for arbitrary currents, let’s make a weird “recorded” current and run it on that.

```python
start_scope()
A = 2.5
f = 10*Hz
tau = 5*ms
# Let's create an array that couldn't be
# reproduced with a formula
num_samples = int(200*ms/defaultclock.dt)
I_arr = zeros(num_samples)
for _ in range(100):
    a = randint(num_samples)
```

(continues on next page)
I_arr[a:a+100] = rand()
I_recorded = TimedArray(A*I_arr, dt=defaultclock.dt)
eqs = ''
dv/dt = (I-v)/tau : 1
I = I_recorded(t) : 1
''
G = NeuronGroup(1, eqs, threshold='v>1', reset='v=0', method='exact')
M = StateMonitor(G, variables=True, record=True)
run(200*ms)
plot(M.t/ms, M.v[0], label='v')
plot(M.t/ms, M.I[0], label='I')
xlabel('Time (ms)')
ylabel('v')
legend(loc='best');

Finally, let’s finish on an example that actually reads in some data from a file. See if you can work out how this example works.

start_scope()
from matplotlib.image import imread
img = (1-imread('brian.png'))[::-1, :, 0].T
num_samples, N = img.shape
ta = TimedArray(img, dt=1*ms) # 228
A = 1.5
tau = 2*ms
eqs = ''
dv/dt = (A*ta(t, i)-v)/tau+0.8*xi*tau**-0.5 : 1
''
G = NeuronGroup(N, eqs, threshold='v>1', reset='v=0', method='euler')
M = SpikeMonitor(G)
run(num_samples*ms)
plot(M.t/ms, M.i, '.k', ms=3)
xlim(0, num_samples)
ylim(0, N)
xlabel('Time (ms)')
ylabel('Neuron index');
3.1 Importing Brian

After installation, Brian is available in the brian2 package. By doing a wildcard import from this package, i.e.:

```
from brian2 import *
```

you will not only get access to the brian2 classes and functions, but also to everything in the pylab package, which includes the plotting functions from matplotlib and everything included in numpy/scipy (e.g. functions such as arange, linspace, etc.).

The following topics are not essential for beginners.

3.1.1 Precise control over importing

If you want to use a wildcard import from Brian, but don’t want to import all the additional symbols provided by pylab, you can use:

```
from brian2.only import *
```

Note that whenever you use something different from the most general `from brian2 import *` statement, you should be aware that Brian overwrites some numpy functions with their unit-aware equivalents (see Units). If you combine multiple wildcard imports, the Brian import should therefore be the last import. Similarly, you should not import and call overwritten numpy functions directly, e.g. by using `import numpy as np` followed by `np.sin` since this will not use the unit-aware versions. To make this easier, Brian provides a brian2.numpy_package
that provides access to everything in numpy but overwrites certain functions. If you prefer to use prefixed names, the recommended way of doing the imports is therefore:

```python
import brian2.numpy_ as np
import brian2.only as br2
```

Note that it is safe to use e.g. `np.sin` and `numpy.sin` after a `from brian2 import *`.

### 3.1.2 Dependency checks

Brian will check the dependency versions during import and raise an error for an outdated dependency. An outdated dependency does not necessarily mean that Brian cannot be run with it, it only means that Brian is untested on that version. If you want to force Brian to run despite the outdated dependency, set the `core.outdated_dependency_error` preference to `False`. Note that this cannot be done in a script, since you do not have access to the preferences before importing `brian2`. See *Preferences* for instructions how to set preferences in a file.

### 3.2 Physical units

- Using units
- Removing units
- Temperatures
- Constants
- Importing units
- In-place operations on quantities

Brian includes a system for physical units. The base units are defined by their standard SI unit names: amp/ampere, kilogram/kilogramme, second, metre/meter, mole/mol, kelvin, and candela. In addition to these base units, Brian defines a set of derived units: coulomb, farad, gram/gramme, hertz, joule, liter/litre, molar, pascal, ohm, siemens, volt, watt, together with prefixed versions (e.g. msiemens = 0.001*siemens) using the prefixes p, n, µ, k, M, G, T (two exceptions to this rule: kilogram is not defined with any additional prefixes, and metre and meter are additionally defined with the “centi” prefix, i.e. cmetre/cmeter). For convenience, a couple of additional useful standard abbreviations such as cm (instead of cmetre/cmeter), nS (instead of nsiemens), ms (instead of msecond), Hz (instead of hertz), mM (instead of mmolar) are included. To avoid clashes with common variable names, no one-letter abbreviations are provided (e.g. you can use mV or nS, but not V or S).

#### 3.2.1 Using units

You can generate a physical quantity by multiplying a scalar or vector value with its physical unit:

```python
>>> tau = 20*ms
>>> print(tau)
20. ms
>>> rates = [10, 20, 30]*Hz
>>> print(rates)
[ 10.  20.  30.] Hz
```
Brian will check the consistency of operations on units and raise an error for dimensionality mismatches:

```python
>>> tau += 1  # ms? second?
Traceback (most recent call last):
  ...  
  DimensionMismatchError: Cannot calculate ... += 1, units do not match (units are second and 1).
>>> 3*kg + 3*amp
Traceback (most recent call last):
  ...  
  DimensionMismatchError: Cannot calculate 3. kg + 3. A, units do not match (units are kilogram and amp).
```

Most Brian functions will also complain about non-specified or incorrect units:

```python
>>> G = NeuronGroup(10, 'dv/dt = -v/tau: volt', dt=0.5)
Traceback (most recent call last):
  ...  
  DimensionMismatchError: Function "__init__" expected a quantitity with unit second for argument "dt" but got 0.5 (unit is 1).
```

Numpy functions have been overwritten to correctly work with units (see the developer documentation for more details):

```python
>>> print(mean(rates))
20. Hz
>>> print(rates.repeat(2))
[ 10. 10. 20. 20. 30. 30.] Hz
```

### 3.2.2 Removing units

There are various options to remove the units from a value (e.g. to use it with analysis functions that do not correctly work with units)

- Divide the value by its unit (most of the time the recommended option because it is clear about the scale)
- Transform it to a pure numpy array in the base unit by calling `asarray()` (no copy) or `array` (copy)
- Directly get the unitless value of a state variable by appending an underscore to the name

```python
>>> tau/ms
20.0
>>> asarray(rates)
array([ 10., 20., 30.])
>>> G = NeuronGroup(5, 'dv/dt = -v/tau: volt')
>>> print(G.v_[::])
[ 0. 0. 0. 0. 0.]
```

### 3.2.3 Temperatures

Brian only supports temperatures defined in °K, using the provided `kelvin` unit object. Other conventions such as °C, or °F are not compatible with Brian’s unit system, because they cannot be expressed as a multiplicative scaling of the SI base unit kelvin (their zero point is different). However, in biological experiments and modeling, temperatures are typically reported in °C. How to use such temperatures depends on whether they are used as temperature differences or as absolute temperatures:

3.2. Physical units
temperature differences  Their major use case is the correction of time constants for differences in temperatures based on the Q10 temperature coefficient. In this case, all temperatures can directly use kelvin even though the temperatures are reported in Celsius, since temperature differences in Celsius and Kelvin are identical.

absolute temperatures  Equations such as the Goldman–Hodgkin–Katz voltage equation have a factor that depends on the absolute temperature measured in Kelvin. To get this temperature from a temperature reported in °C, you can use the zero_celsius constant from the brian2.units.constants package (see below):

```
from brian2.units.constants import zero_celsius

celsius_temp = 27
abs_temp = celsius_temp*kelvin + zero_celsius
```

Note:  Earlier versions of Brian had a celsius unit which was in fact identical to kelvin. While this gave the correct results for temperature differences, it did not correctly work for absolute temperatures. To avoid confusion and possible misinterpretation, the celsius unit has therefore been removed.

### 3.2.4 Constants

The brian2.units.constants package provides a range of physical constants that can be useful for detailed biological models. Brian provides the following constants:

<table>
<thead>
<tr>
<th>Constant</th>
<th>Symbol(s)</th>
<th>Brian name</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Avogadro constant</td>
<td>( N_A, L )</td>
<td>avogadro_constant</td>
<td>6.022140857 \times 10^{23} \text{ mol}^{-1}</td>
</tr>
<tr>
<td>Boltzmann constant</td>
<td>( k )</td>
<td>boltzmann_constant</td>
<td>1.38064852 \times 10^{-23} \text{ J K}^{-1}</td>
</tr>
<tr>
<td>Electric constant</td>
<td>( \epsilon_0 )</td>
<td>electric_constant</td>
<td>8.854187817 \times 10^{-12} \text{ F m}^{-1}</td>
</tr>
<tr>
<td>Electron mass</td>
<td>( m_e )</td>
<td>electron_mass</td>
<td>9.10938356 \times 10^{-31} \text{ kg}</td>
</tr>
<tr>
<td>Elementary charge</td>
<td>( e )</td>
<td>elementary_charge</td>
<td>1.6021766208 \times 10^{-19} \text{ C}</td>
</tr>
<tr>
<td>Faraday constant</td>
<td>( F )</td>
<td>faraday_constant</td>
<td>96485.33289 \text{ C mol}^{-1}</td>
</tr>
<tr>
<td>Gas constant</td>
<td>( R )</td>
<td>gas_constant</td>
<td>8.3144598 \text{ J mol}^{-1} \text{ K}^{-1}</td>
</tr>
<tr>
<td>Magnetic constant</td>
<td>( \mu_0 )</td>
<td>magnetic_constant</td>
<td>12.566370614 \times 10^{-7} \text{ N A}^{-2}</td>
</tr>
<tr>
<td>Molar mass constant</td>
<td>( M_u )</td>
<td>molar_mass_constant</td>
<td>1 \times 10^{-3} \text{ kg mol}^{-1}</td>
</tr>
<tr>
<td>0°C</td>
<td></td>
<td>zero_celsius</td>
<td>273.15 K</td>
</tr>
</tbody>
</table>

Note that these constants are not imported by default, you will have to explicitly import them from brian2.units.constants. During the import, you can also give them shorter names using Python’s from ... import ... as ... syntax. For example, to calculate the \( \frac{RT}{F} \) factor that appears in the Goldman–Hodgkin–Katz voltage equation you can use:

```
from brian2 import *
from brian2.units.constants import zero_celsius, gas_constant as R, faraday_constant as F

celsius_temp = 27
T = celsius_temp*kelvin + zero_celsius
factor = R*T/F
```

The following topics are not essential for beginners.
3.2.5 Importing units

Brian generates standard names for units, combining the unit name (e.g. “siemens”) with a prefix (e.g. “m”), and also generates squared and cubed versions by appending a number. For example, the units “msiemens”, “siemens2”, “usiemens3” are all predefined. You can import these units from the package `brian2.units.allunits`—accordingly, an `from brian2.units.allunits import *` will result in everything from `Ylumen3` (cubed yotta lumen) to `ymol` (yocto mole) being imported.

A better choice is normally to do `from brian2.units import *` or `import everything from brian2 import *` which only imports the units mentioned in the introductory paragraph (base units, derived units, and some standard abbreviations).

3.2.6 In-place operations on quantities

In-place operations on quantity arrays change the underlying array, in the same way as for standard numpy arrays. This means, that any other variables referencing the same object will be affected as well:

```python
>>> q = [1, 2] * mV
>>> r = q
>>> q += 1*mV
>>> q
array([ 2., 3.]) * mvolt
>>> r
array([ 2., 3.]) * mvolt
```

In contrast, scalar quantities will never change the underlying value but instead return a new value (in the same way as standard Python scalars):

```python
>>> x = 1*mV
>>> y = x
>>> x *= 2
>>> x
2. * mvolt
>>> y
1. * mvolt
```

3.3 Models and neuron groups

For Brian 1 users

See the document *Neural models (Brian 1 –> 2 conversion)* for details how to convert Brian 1 code.

- Model equations
- Noise
- Threshold and reset
3.3.1 Model equations

The core of every simulation is a NeuronGroup, a group of neurons that share the same equations defining their properties. The minimum NeuronGroup specification contains the number of neurons and the model description in the form of equations:

\[
G = \text{NeuronGroup}(10, \ 'dv/dt = -v/(10*ms) : volt')
\]

This defines a group of 10 leaky integrators. The model description can be directly given as a (possibly multi-line) string as above, or as an Equations object. For more details on the form of equations, see Equations. Brian needs the model to be given in the form of differential equations, but you might see the integrated form of synapses in some textbooks and papers. See Converting from integrated form to ODEs for details on how to convert between these representations.

Note that model descriptions can make reference to physical units, but also to scalar variables declared outside of the model description itself:

\[
\tau = 10*\text{ms}
G = \text{NeuronGroup}(10, \ 'dv/dt = -v/\tau : volt')
\]

If a variable should be taken as a parameter of the neurons, i.e. if it should be possible to vary its value across neurons, it has to be declared as part of the model description:

\[
G = \text{NeuronGroup}(10, \ ''dv/dt = -v/\tau : volt}
\quad \tau : \text{second'''}
\]

To make complex model descriptions more readable, named subexpressions can be used:

\[
G = \text{NeuronGroup}(10, \ ''dv/dt = I_{\text{leak}} / C_m : volt}
\quad I_{\text{leak}} = g_L*(E_L - v) : \text{amp'''}
\]

For a list of some standard model equations, see Neural models (Brian 1 \rightarrow 2 conversion).

3.3.2 Noise

In addition to ordinary differential equations, Brian allows you to introduce random noise by specifying a stochastic differential equation. Brian uses the physicists’ notation used in the Langevin equation, representing the “noise” as a term \(\xi(t)\), rather than the mathematicians’ stochastic differential \(dW_t\). The following is an example of the Ornstein-Uhlenbeck process that is often used to model a leaky integrate-and-fire neuron with a stochastic current:

\[
G = \text{NeuronGroup}(10, \ 'dv/dt = -v/\tau + \sigma*\sqrt{2/\tau} \cdot \xi : volt')
\]
You can start by thinking of $x_i$ as just a Gaussian random variable with mean 0 and standard deviation 1. However, it scales in an unusual way with time and this gives it units of $1/\sqrt{\text{second}}$. You don’t necessarily need to understand why this is, but it is possible to get a reasonably simple intuition for it by thinking about numerical integration: see below.

### 3.3.3 Threshold and reset

To emit spikes, neurons need a threshold. Threshold and reset are given as strings in the `NeuronGroup` constructor:

```python
tau = 10*ms
G = NeuronGroup(10, 'dv/dt = -v/tau : volt', threshold='v > -50*mV', reset='v = -70*mV')
```

Whenever the threshold condition is fulfilled, the reset statements will be executed. Again, both threshold and reset can refer to physical units, external variables and parameters, in the same way as model descriptions:

```python
v_r = -70*mV  # reset potential
G = NeuronGroup(10, '''dv/dt = -v/tau : volt
    v_th : volt  # neuron-specific threshold''',
    threshold='v > v_th', reset='v = v_r')
```

You can also create non-spike events. See *Custom events* for more details.

### 3.3.4 Refractoriness

To make a neuron non-excitable for a certain time period after a spike, the refractory keyword can be used:

```python
G = NeuronGroup(10, 'dv/dt = -v/tau : volt', threshold='v > -50*mV',
    reset='v = -70*mV', refractory=5*ms)
```

This will not allow any threshold crossing for a neuron for 5ms after a spike. The refractory keyword allows for more flexible refractoriness specifications, see *Refractoriness* for details.

### 3.3.5 State variables

Differential equations and parameters in model descriptions are stored as state variables of the `NeuronGroup`. In addition to these variables, Brian also defines two variables automatically:

- $i$ The index of a neuron.
- $N$ The total number of neurons.

All state variables can be accessed and set as an attribute of the group. To get the values without physical units (e.g. for analysing data with external tools), use an underscore after the name:

```python
>>> G = NeuronGroup(10, '''dv/dt = -v/tau : volt
    tau : second''', name='neurons')
>>> G.v = -70*mV
>>> G.v
<neurons.v: array([-70., -70., -70., -70., -70., -70., -70., -70.]) * 
   → mV>
>>> G.v_  # values without units
<neurons.v_: array([-0.07, -0.07, -0.07, -0.07, -0.07, -0.07, -0.07, -0.07])>
```
The value of state variables can also be set using string expressions that can refer to units and external variables, other state variables or mathematical functions:

```python
>>> G.tau = '5*ms + (1.0*i/N)*5*ms'
>>> G.tau
<neurons.tau: array([ 5. , 5.5, 6. , 6.5, 7. , 7.5, 8. , 8.5, 9. , 9.5]) * msecond>
```

You can also set the value only if a condition holds, for example:

```python
>>> G.v['tau>7.25*ms'] = -60*mV
>>> G.v
<neurons.v: array([-70., -70., -70., -70., -70., -60., -60., -60., -60., -60.]) * mvolt>
```

### 3.3.6 Subgroups

It is often useful to refer to a subset of neurons, this can be achieved using Python’s slicing syntax:

```python
G = NeuronGroup(10, '''dv/dt = -v/tau : volt
        tau : second''',
    threshold='v > -50*mV',
    reset='v = -70*mV')

# Create subgroups
G1 = G[:5]
G2 = G[5:]

# This will set the values in the main group, subgroups are just "views"
G1.tau = 10*ms
G2.tau = 20*ms
```

Here `G1` refers to the first 5 neurons in `G`, and `G2` to the second 5 neurons. In general `G[i:j]` refers to the neurons with indices from `i` to `j-1`, as in general in Python.

For convenience, you can also use a single index, i.e. `G[i]` is equivalent to `G[i:i+1]`. In some situations, it can be easier to provide a list of indices instead of a slice, Brian therefore also allows for this syntax. Note that this is restricted to cases that are strictly equivalent with slicing syntax, e.g. you can write `G[[3, 4, 5]]` instead of `G[3:6]`, but you cannot write `G[[3, 5, 7]]` or `G[[5, 4, 3]]`.

Subgroups can be used in most places where regular groups are used, e.g. their state variables or spiking activity can be recorded using monitors, they can be connected via `Synapses`, etc. In such situations, indices (e.g. the indices of the neurons to record from in a `StateMonitor`) are relative to the subgroup, not to the main group.

The following topics are not essential for beginners.

### 3.3.7 Shared variables

Sometimes it can also be useful to introduce shared variables or subexpressions, i.e. variables that have a common value for all neurons. In contrast to external variables (such as `Cm` above), such variables can change during a run, e.g.
by using `run_regularly()`. This can be for example used for an external stimulus that changes in the course of a run:

```python
>>> G = NeuronGroup(10, '''
shared_input : volt (shared)
... 
  dv/dt = (-v + shared_input)/tau : volt
... 
  tau : second''', name='neurons')
```

Note that there are several restrictions around the use of shared variables: they cannot be written to in contexts where statements apply only to a subset of neurons (e.g. reset statements, see below). If a code block mixes statements writing to shared and vector variables, then the shared statements have to come first.

By default, subexpressions are re-evaluated whenever they are used, i.e. using a subexpression is completely equivalent to substituting it. Sometimes it is useful to instead only evaluate a subexpression once and then use this value for the rest of the time step. This can be achieved by using the \texttt{(constant over dt)} flag. This flag is mandatory for subexpressions that refer to stateful functions like \texttt{rand()} which notably allows them to be recorded with a \texttt{StateMonitor} – otherwise the monitor would record a different instance of the random number than the one that was used in the equations.

For shared variables, setting by string expressions can only refer to shared values:

```python
>>> G.shared_input = '(4.0/N)*mV'
>>> G.shared_input
<neurons.shared_input: 0.4 * mvolt>
```

### 3.3.8 Storing state variables

Sometimes it can be convenient to access multiple state variables at once, e.g. to set initial values from a dictionary of values or to store all the values of a group on disk. This can be done with the \texttt{get_states()} and \texttt{set_states()} methods:

```python
>>> group = NeuronGroup(5, '''
    dv/dt = -v/tau : 1
... 
    tau : second''', name='neurons')
>>> initial_values = {
...    'v': [0, 1, 2, 3, 4],
...    'tau': [10, 20, 10, 20, 10]*ms
...}
>>> group.set_states(initial_values)
>>> group.v[:]
array([ 0., 1., 2., 3., 4.])
>>> group.tau[:]
array([ 10., 20., 10., 20., 10.]) * msecond
>>> states = group.get_states()
>>> states['v']
array([ 0., 1., 2., 3., 4.])
```

The data (without physical units) can also be exported/imported to/from Pandas data frames (needs an installation of pandas):

```python
>>> df = group.get_states(units=False, format='pandas')
>>> df
    N   dt  i   t  tau  v
0   5 0.0001  0  0.0  0.01  0.0
1   5 0.0001  1  0.0  0.02  1.0
2   5 0.0001  2  0.0  0.01  2.0
3   5 0.0001  3  0.0  0.02  3.0
4   5 0.0001  4  0.0  0.01  4.0
```

(continues on next page)
3.3.9 Linked variables

A `NeuronGroup` can define parameters that are not stored in this group, but are instead a reference to a state variable in another group. For this, a group defines a parameter as linked and then uses `linked_var()` to specify the linking. This can for example be useful to model shared noise between cells:

```python
ingp = NeuronGroup(1, 'dnoise/dt = -noise/tau + tau***0.5*xi : 1')
neurons = NeuronGroup(100, '''noise : 1 (linked)
dv/dt = (-v + noise_strength*noise)/tau : volt''')
neurons.noise = linked_var(inp, 'noise')
```

If the two groups have the same size, the linking will be done in a 1-to-1 fashion. If the source group has the size one (as in the above example) or if the source parameter is a shared variable, then the linking will be done as 1-to-all. In all other cases, you have to specify the indices to use for the linking explicitly:

```python
# two inputs with different phases
inp = NeuronGroup(2, '''phase : 1
dx/dt = 1*mV/ms*sin(2*pi*100*Hz*t-phase) : volt''')
inp.phase = [0, pi/2]

neurons = NeuronGroup(100, '''inp : volt (linked)
dv/dt = (-v + inp) / tau : volt''')
# Half of the cells get the first input, other half gets the second
neurons.inp = linked_var(inp, 'x', index=repeat([0, 1], 50))
```

3.3.10 Time scaling of noise

Suppose we just had the differential equation

\[ \frac{dx}{dt} = \xi \]

To solve this numerically, we could compute

\[ x(t + dt) = x(t) + \xi_1 \]

where \( \xi_1 \) is a normally distributed random number with mean 0 and standard deviation 1. However, what happens if we change the time step? Suppose we used a value of \( dt/2 \) instead of \( dt \). Now, we compute

\[ x(t + dt) = x(t + dt/2) + \xi_1 = x(t) + \xi_2 + \xi_1 \]

The mean value of \( x(t + dt) \) is 0 in both cases, but the standard deviations are different. The first method \( x(t + dt) = x(t) + \xi_1 \) gives \( x(t + dt) \) a standard deviation of 1, whereas the second method \( x(t + dt) = x(t + dt/2) + \xi_1 = x(t) + \xi_2 + \xi_1 \) gives \( x(t) \) a variance of \( 1+1=2 \) and therefore a standard deviation of \( \sqrt{2} \).
In order to solve this problem, we use the rule \( x(t + dt) = x(t) + \sqrt{dt} \xi_1 \), which makes the mean and standard deviation of the value at time \( t \) independent of \( dt \). For this to make sense dimensionally, \( \xi \) must have units of \( 1/\sqrt{\text{second}} \).

For further details, refer to a textbook on stochastic differential equations.

### 3.4 Numerical integration

By default, Brian chooses an integration method automatically, trying to solve the equations exactly first (for linear equations) and then resorting to numerical algorithms. It will also take care of integrating stochastic differential equations appropriately.

Note that in some cases, the automatic choice of integration method will not be appropriate, because of a choice of parameters that couldn’t be determined in advance. In this case, typically you will get nan (not a number) values in the results, or large oscillations. In this case, Brian will generate a warning to let you know, but will not raise an error.

#### 3.4.1 Method choice

You will get an INFO message telling you which integration method Brian decided to use, together with information about how much time it took to apply the integration method to your equations. If other methods have been tried but were not applicable, you will also see the time it took to try out those other methods. In some cases, checking other methods (in particular the 'exact' method which attempts to solve the equations analytically) can take a considerable amount of time – to avoid wasting this time, you can always chose the integration method manually (see below). You can also suppress the message by raising the log level or by explicitly suppressing 'method_choice' log messages – for details, see Logging.

If you prefer to chose an integration algorithm yourself, you can do so using the method keyword for NeuronGroup, Synapses, or SpatialNeuron. The complete list of available methods is the following:

- 'exact': exact integration for linear equations (alternative name: 'linear')
- 'exponential_euler': exponential Euler integration for conditionally linear equations
- 'euler': forward Euler integration (for additive stochastic differential equations using the Euler-Maruyama method)
- 'rk2': second order Runge-Kutta method (midpoint method)
- 'rk4': classical Runge-Kutta method (RK4)
- 'milstein': derivative-free Milstein method for solving stochastic differential equations with diagonal multiplicative noise

**Note:** The 'independent' integration method (exact integration for a system of independent equations, where all the equations can be analytically solved independently) should no longer be used and might be removed in future versions of Brian.

**Note:** The following methods are still considered experimental

- 'gsl': default integrator when choosing to integrate equations with the GNU Scientific Library ODE solver: the rkf45 method. Uses an adaptable time step by default.
Brian 2 Documentation, Release 2.3

- 'gsl_rkf45': Runge-Kutta-Fehlberg method. A good general-purpose integrator according to the GSL documentation. Uses an adaptable time step by default.
- 'gsl_rk4': Fourth order Runge-Kutta method using GSL. Uses an adaptable time step by default.
- 'gsl_rk8pd': Runge-Kutta Prince-Dormand method using GSL. Uses an adaptable time step by default.

The following topics are not essential for beginners.

### 3.4.2 Technical notes

Each class defines its own list of algorithms it tries to apply. `NeuronGroup` and `Synapses` will use the first suitable method out of the methods 'exact', 'euler' and 'heun' while `SpatialNeuron` objects will use 'exact', 'exponential_euler', 'rk2' or 'heun'.

You can also define your own numerical integrators, see State update for details.

### 3.4.3 GSL stateupdaters

The stateupdaters preceded with the gsl tag use ODE solvers defined in the GNU Scientific Library. The benefit of using these integrators over the ones written by Brian internally, is that they are implemented with an adaptable timestep. Integrating with an adaptable timestep comes with two advantages:

- These methods check whether the estimated error of the solutions returned fall within a certain error bound. For the non-gsl integrators there is currently no such check.
- Systems no longer need to be simulated with just one time step. That is, a bigger timestep can be chosen and the integrator will reduce the timestep when increased accuracy is required. This is particularly useful for systems where both slow and fast time constants coexist, as is the case with for example (networks of neurons with) Hodgkin-Huxley equations. Note that Brian's timestep still determines the resolution for monitors, spike timing, spike propagation etc. Hence, in a network, the simulation error will therefore still be on the order of \( \Delta t \). The benefit is that short time constants occurring in equations no longer dictate the network time step.

In addition to a choice between different integration methods, there are a few more options that can be specified when using GSL. These options can be specified by sending a dictionary as the `method_options` key upon initialization of the object using the integrator (`NeuronGroup`, `Synapses` or `SpatialNeuron`). The available method options are:

- 'adaptable_timestep': whether or not to let GSL reduce the timestep to achieve the accuracy defined with the 'absolute_error' and 'absolute_error_per_variable' options described below. If this is set to False, the timestep is determined by Brian (i.e. the \( \Delta t \) of the respective clock is used, see Scheduling). If the resulted estimated error exceeds the set error bounds, the simulation is aborted. When using cython or weave this is reported with an IntegrationError. Defaults to True.
- 'absolute_error': each of the methods has a way of estimating the error that is the result of using numerical integration. You can specify the maximum size of this error to be allowed for any of the to-be-integrated variables in base units with this keyword. Note that giving very small values makes the simulation slow and...
might result in unsuccessful integration. In the case of using the 'absolute_error_per_variable' option, this is the error for variables that were not specified individually. Defaults to 1e-6.

- 'absolute_error_per_variable': specify the absolute error per variable in its own units. Variables for which the error is not specified use the error set with the 'absolute_error' option. Defaults to None.

- 'max_steps': The maximal number of steps that the integrator will take within a single “Brian timestep” in order to reach the given error criterion. Can be set to 0 to not set any limits. Note that without limits, it can take a very long time until the integrator figures out that it cannot reach the desired error level. This will manifest as a simulation that appears to be stuck. Defaults to 100.

- 'use_last_timestep': with the 'adaptable_timestep' option set to True, GSL tries different time steps to find a solution that satisfies the set error bounds. It is likely that for Brian’s next time step the GSL time step will be somewhat similar per neuron (e.g. active neurons will have a shorter GSL time step than inactive neurons). With this option set to True, the time step GSL found to satisfy the set error bounds is saved per neuron and given to GSL again in Brian’s next time step. This also means that the final time steps are saved in Brian’s memory and can thus be recorded with the StateMonitor: it can be accessed under '_last_timestep'. Note that some extra memory is required to keep track of the last time steps. Defaults to True.

- 'save_failed_steps': if 'adaptable_timestep' is set to True, each time GSL tries a time step and it results in an estimated error that exceeds the set bounds, one is added to the '_failed_steps' variable. For purposes of investigating what happens within GSL during an integration step, we offer the option of saving this variable. Defaults to False.

- 'save_step_count': the same goes for the total number of GSL steps taken in a single Brian time step: this is optionally saved in the '_step_count' variable. Defaults to False.

Note that at the moment recording '_last_timestep', '_failed_steps', or '_step_count' requires a call to run() (e.g. with 0 ms) to trigger the code generation process, before the call to StateMonitor.

More information on the GSL ODE solver itself can be found in its documentation.

### 3.5 Equations

- **Equation strings**
- **Arithmetic operations and functions**
- **External variables**
- **Flags**
- **List of special symbols**
- **Event-driven equations**
- **Equation objects**
- **Examples of Equation objects**

### 3.5.1 Equation strings

Equations are used both in NeuronGroup and Synapses to:

- define state variables
- define continuous-updates on these variables, through differential equations
Note: Brian models are defined by systems of first order ordinary differential equations, but you might see the integrated form of synapses in some textbooks and papers. See *Converting from integrated form to ODEs* for details on how to convert between these representations.

Equations are defined by multiline strings.

An Equation is a set of single lines in a string:

1. \( \frac{dx}{dt} = f \) : unit (differential equation)
2. \( x = f \) : unit (subexpression)
3. \( x \) : unit (parameter)

Each equation may be spread out over multiple lines to improve formatting. Comments using `#` may also be included. Subunits are not allowed, i.e., one must write `volt`, not `mV`. This is to make it clear that the values are internally always saved in the basic units, so no confusion can arise when getting the values out of a `NeuronGroup` and discarding the units. Compound units are of course allowed as well (e.g. `farad/meter**2`). There are also three special “units” that can be used: 1 denotes a dimensionless floating point variable, `boolean` and `integer` denote dimensionless variables of the respective kind.

Note: For molar concentration, the base unit that has to be used in the equations is `mmolar` (or `mM`), not `molar`. This is because 1 molar is \( 10^3 \text{ mol/m}^3 \) in SI units (i.e., it has a “scale” of \( 10^3 \)), whereas 1 millimolar corresponds to 1 \( \text{mol/m}^3 \).

Some special variables are defined: `t`, `dt` (time) and `xi` (white noise). Variable names starting with an underscore and a couple of other names that have special meanings under certain circumstances (e.g. names ending in `_pre` or `_post`) are forbidden.

For stochastic equations with several `xi` values it is necessary to make clear whether they correspond to the same or different noise instantiations. To make this distinction, an arbitrary suffix can be used, e.g. using `xi_1` several times refers to the same variable, `xi_2` (or `xi_inh`, `xi_alpha`, etc.) refers to another. An error will be raised if you use more than one plain `xi`. Note that noise is always independent across neurons, you can only work around this restriction by defining your noise variable as a shared parameter and update it using a user-defined function (e.g. with `run_regularly`), or create a group that models the noise and link to its variable (see *Linked variables*).

### 3.5.2 Arithmetic operations and functions

Equation strings can make use of standard arithmetic operations for numerical values, using the Python 3 syntax. The supported operations are `+`, `-`, `*`, `/` (floating point division), `//` (flooring division), `%` (remainder), `**` (power). For variable assignments, e.g. in reset statements, the corresponding in-place assignments such as `+=` can be used as well. For comparisons, the operations `==` (equality), `!=` (inequality), `<`, `<=`, `>`, and `>=` are available. Truth values can be combined using `and` and `or`, or negated using `not`. Note that Brian does not support any operations specific to integers, e.g. “bitwise AND” or shift operations.

**Warning:** Brian versions up to 2.1.3.1 did not support `//` as the floor division operator and potentially used different semantics for the `/` operator depending on whether Python 2 or 3 was used. To write code that correctly and unambiguously works with both newer and older Brian versions, you can use expressions such as `1.0*a/b` to enforce floating point division (if one of the operands is a floating point number, both Python 2 and 3 will use floating point division), or `floor(a/b)` to enforce flooring division Note that the `floor` function always returns a floating point value, if it is important that the result is an integer value, additionally wrap it with the `int` function.
Brian also supports standard mathematical functions with the same names as used in the numpy library (e.g. `exp`, `sqrt`, `abs`, `clip`, `sin`, `cos`, ...) – for a full list see Default functions. Note that support for such functions is provided by Brian itself and the translation to the various code generation targets is automatically taken care of. You should therefore refer to them directly by name and not as e.g. `np.sqrt` or `numpy.sqrt`, regardless of the way you imported Brian or numpy. This also means that you cannot directly refer to arbitrary functions from numpy or other libraries. For details on how to extend the support to non-default functions see User-provided functions.

### 3.5.3 External variables

Equations defining neuronal or synaptic equations can contain references to external parameters or functions. These references are looked up at the time that the simulation is run. If you don’t specify where to look them up, it will look in the Python local/global namespace (i.e. the block of code where you call `run()`). If you want to override this, you can specify an explicit “namespace”. This is a Python dictionary with keys being variable names as they appear in the equations, and values being the desired value of that variable. This namespace can be specified either in the creation of the group or when you can the `run()` function using the `namespace` keyword argument.

The following three examples show the different ways of providing external variable values, all having the same effect in this case:

```python
# Explicit argument to the NeuronGroup
G = NeuronGroup(1, 'dv/dt = -v / tau : 1', namespace={'tau': 10*ms})
net = Network(G)
net.run(10*ms)

# Explicit argument to the run function
G = NeuronGroup(1, 'dv/dt = -v / tau : 1')
net = Network(G)
net.run(10*ms, namespace={'tau': 10*ms})

# Implicit namespace from the context
G = NeuronGroup(1, 'dv/dt = -v / tau : 1')
net = Network(G)
tau = 10*ms
net.run(10*ms)
```

See Namespaces for more details.

The following topics are not essential for beginners.

### 3.5.4 Flags

A flag is a keyword in parentheses at the end of the line, which qualifies the equations. There are several keywords:

- **event-driven** this is only used in Synapses, and means that the differential equation should be updated only at the times of events. This implies that the equation is taken out of the continuous state update, and instead a event-based state update statement is generated and inserted into event codes (pre and post). This can only qualify differential equations of synapses. Currently, only one-dimensional linear equations can be handled (see below).
unless refractory  this means the variable is not updated during the refractory period. This can only qualify differential equations of neuron groups.

constant  this means the parameter will not be changed during a run. This allows optimizations in state updaters. This can only qualify parameters.

constant over dt  this means that the subexpression will be only evaluated once at the beginning of the time step. This can be useful to e.g. approximate a non-linear term as constant over a time step in order to use the linear numerical integration algorithm. It is also mandatory for subexpressions that refer to stateful functions like rand() to make sure that they are only evaluated once (otherwise e.g. recording the value with a StateMonitor would re-evaluate it and therefore not record the same values that are used in other places). This can only qualify subexpressions.

shared  this means that a parameter or subexpression is not neuron-/synapse-specific but rather a single value for the whole NeuronGroup or Synapses. A shared subexpression can only refer to other shared variables.

linked  this means that a parameter refers to a parameter in another NeuronGroup. See Linked variables for more details.

Multiple flags may be specified as follows:

\[
\frac{dx}{dt} = f : \text{unit} \ (\text{flag1,flag2})
\]

3.5.5 List of special symbols

The following lists all of the special symbols that Brian uses in equations and code blocks, and their meanings.

dt  Time step width

i  Index of a neuron (NeuronGroup) or the pre-synaptic neuron of a synapse (Synapses)

j  Index of a post-synaptic neuron of a synapse

lastspike  Last time that the neuron spiked (for refractoriness)

lastupdate  Time of the last update of synaptic variables in event-driven equations (only defined when event-driven equations are used).

N  Number of neurons (NeuronGroup) or synapses (Synapses). Use N_pre or N_post for the number of presynaptic or postsynaptic neurons in the context of Synapses.

not_refractory  Boolean variable that is normally true, and false if the neuron is currently in a refractory state

t  Current time

t_in_timesteps  Current time measured in time steps

xi, xi_  Stochastic differential in equations

3.5.6 Event-driven equations

Equations defined as event-driven are completely ignored in the state update. They are only defined as variables that can be externally accessed. There are additional constraints:

- An event-driven variable cannot be used by any other equation that is not also event-driven.
- An event-driven equation cannot depend on a differential equation that is not event-driven (directly, or indirectly through subexpressions). It can depend on a constant parameter.

Currently, automatic event-driven updates are only possible for one-dimensional linear equations, but this may be extended in the future.
3.5.7 Equation objects

The model definitions for NeuronGroup and Synapses can be simple strings or Equations objects. Such objects can be combined using the add operator:

```python
eqs = Equations('dx/dt = (y-x)/tau : volt')
eqs += Equations('dy/dt = -y/tau: volt')
```

Equations allow for the specification of values in the strings, but does this by simple string replacement, e.g. you can do:

```python
eqs = Equations('dx/dt = x/tau : volt', tau=10*ms)
```

but this is exactly equivalent to:

```python
eqs = Equations('dx/dt = x/(10*ms) : volt')
```

The Equations object does some basic syntax checking and will raise an error if two equations defining the same variable are combined. It does not however do unit checking, checking for unknown identifiers or incorrect flags – all this will be done during the instantiation of a NeuronGroup or Synapses object.

3.5.8 Examples of Equation objects

Concatenating equations

```python
>>> membrane_eqs = Equations('dv/dt = -(v + I)/ tau : volt')
>>> eqs1 = membrane_eqs + Equations('''I = sin(2*pi*freq*t) : volt
... freq : Hz''')
>>> eqs2 = membrane_eqs + Equations('''I : volt''')
>>> print(eqs1)
I = sin(2*pi*freq*t) : V
dv/dt = -(v + I)/ tau : V
freq : Hz
>>> print(eqs2)
dv/dt = -(v + I)/ tau : V
I : V
```

Substituting variable names

```python
>>> general_equation = 'dg/dt = -g / tau : siemens'
>>> eqs_exc = Equations(general_equation, g='g_e', tau='tau_e')
>>> eqs_inh = Equations(general_equation, g='g_i', tau='tau_i')
>>> print(eqs_exc)
dg_e/dt = -g_e / tau_e : S
>>> print(eqs_inh)
dg_i/dt = -g_i / tau_i : S
```

Inserting values

```python
>>> eqs = Equations('dv/dt = mu/tau + sigma/tau**.5*xi : volt',
... mu=-65*mV, sigma=3*mV, tau=10*ms)
>>> print(eqs)
```
3.6 Refractoriness

- **Defining the refractory period**
- **Defining model behaviour during refractoriness**
- **Arbitrary refractoriness**

Brian allows you to model the absolute refractory period of a neuron in a flexible way. The definition of refractoriness consists of two components: the amount of time after a spike that a neuron is considered to be refractory, and what changes in the neuron during the refractoriness.

### 3.6.1 Defining the refractory period

The refractory period is specified by the `refractory` keyword in the `NeuronGroup` initializer. In the simplest case, this is simply a fixed time, valid for all neurons:

```
G = NeuronGroup(N, model='...', threshold='...', reset='...',
refractory=2*ms)
```

Alternatively, it can be a string expression that evaluates to a time. This expression will be evaluated after every spike and allows for a changing refractory period. For example, the following will set the refractory period to a random duration between 1ms and 3ms after every spike:

```
G = NeuronGroup(N, model='...', threshold='...', reset='...',
refractory='(1 + 2*rand())*ms')
```

In general, modelling a refractory period that varies across neurons involves declaring a state variable that stores the refractory period per neuron as a model parameter. The refractory expression can then refer to this parameter:

```
G = NeuronGroup(N, model='''...
    refractory : second'', threshold='...',
    reset='...', refractory='refractory')
# Set the refractory period for each cell
G.refractory = ...
```

This state variable can also be a dynamic variable itself. For example, it can serve as an adaptation mechanism by increasing it after every spike and letting it relax back to a steady-state value between spikes:

```
refractory_0 = 2*ms
tau_refractory = 50*ms
G = NeuronGroup(N, model='''...
    drefractory/dt = (refractory_0 - refractory) / tau_
    ➔refractory : second'',
    threshold='...', refractory='refractory',
    reset='''...
    refractory += 1*ms'''}
G.refractory = refractory_0
```

In some cases, the condition for leaving the refractory period is not easily expressed as a certain time span. For example, in a Hodgkin-Huxley type model the threshold is only used for counting spikes and the refractoriness is used to prevent to count multiple spikes for a single threshold crossing (the threshold condition would evaluate to `True` for several time points). When a neuron should leave the refractory period is not easily expressed as a time span but more...
naturally as a condition that the neuron should remain refractory for as long as it stays above the threshold. This can be achieved by using a string expression for the `refractory` keyword that evaluates to a boolean condition:

```plaintext
G = NeuronGroup(N, model='...', threshold='v > -20*mV',
                 refractory='v >= -20*mV')
```

The `refractory` keyword should be read as “stay refractory as long as the condition remains true”. In fact, specifying a time span for the refractoriness will be automatically transformed into a logical expression using the current time `t` and the time of the last spike `lastspike`. Specifying `refractory=2*ms` is basically equivalent to specifying `refractory='(t - lastspike) <= 2*ms'`. However, this expression can give inconsistent results for the common case that the refractory period is a multiple of the simulation timestep. Due to floating point impreciseness, the actual value of `t - lastspike` can be slightly above or below a multiple of the simulation time step; comparing it directly to the refractory period can therefore lead to an end of the refractory one time step sooner or later. To avoid this issue, the actual code used for the above example is equivalent to `refractory='timestep(t - lastspike, dt) <= timestep(2*ms, dt)'`. The `timestep` function is provided by Brian and takes care of converting a time into a time step in a safe way.

New in version 2.1.3: The `timestep` function is now used to avoid floating point issues in the refractoriness calculation. To restore the previous behaviour, set the `legacy.refractory_timing` preference to `True`.

### 3.6.2 Defining model behaviour during refractoriness

The refractoriness definition as described above only has a single effect by itself: threshold crossings during the refractory period are ignored. In the following model, the variable `v` continues to update during the refractory period but it does not elicit a spike if it crosses the threshold:

```plaintext
G = NeuronGroup(N, 'dv/dt = -v / tau : 1',
                 threshold='v > 1', reset='v=0',
                 refractory=2*ms)
```

There is also a second implementation of refractoriness that is supported by Brian, one or several state variables can be clamped during the refractory period. To model this kind of behaviour, variables that should stop being updated during refractoriness can be marked with the `(unless refractory)` flag:

```plaintext
G = NeuronGroup(N, '''dv/dt = -(v + w)/ tau_v : 1 (unless refractory)
                     dw/dt = -w / tau_w : 1''',
                 threshold='v > 1', reset='v=0; w+=0.1', refractory=2*ms)
```

In the above model, the `v` variable is clamped at 0 for 2ms after a spike but the adaptation variable `w` continues to update during this time. In addition, a variable of a neuron that is in its refractory period is `read-only`: incoming synapses or other code will have no effect on the value of `v` until it leaves its refractory period.

The following topics are not essential for beginners.

### 3.6.3 Arbitrary refractoriness

In fact, arbitrary behaviours can be defined using Brian’s refractoriness mechanism. A `NeuronGroup` with refractoriness automatically defines two variables:
**not_refractory**  A boolean variable stating whether a neuron is allowed to spike.

**lastspike**  The time of the last spike of the neuron.

The variable **not_refractory** is updated at every time step by checking the refractoriness condition — for a refractoriness defined by a time period, this means comparing **lastspike** to the current time t. The **not_refractory** variable is then used to implement the refractoriness behaviour. Specifically, the **threshold** condition is replaced by **threshold and not_refractory** and differential equations that are marked as (unless refractory) are multiplied by int(not_refractory) (so that they have the value 0 when the neuron is refractory).

This **not_refractory** variable is also available to the user to define more sophisticated refractoriness behaviour. For example, the following code updates the w variable with a different time constant during refractoriness:

```python
G = NeuronGroup(N, '''dv/dt = -(v + w)/ tau_v : 1 (unless refractory)
   dw/dt = (-w / tau_active)*int(not_refractory) + (-w / tau_ref)*(1 - int(not_refractory)) : 1''',
   threshold='v > 1', reset='v=0; w+=0.1', refractory=2*ms)
```

### 3.7 Synapses

For Brian 1 users

*Synapses* is now the only class for defining synaptic interactions, it replaces *Connection, STDP*, etc. See the document *Synapses (Brian 1 → 2 conversion)* for details how to convert Brian 1 code.

- **Defining synaptic models**
- **Creating synapses**
- **Accessing synaptic variables**
- **Delays**
- **Monitoring synaptic variables**
- **Synaptic connection/weight matrices**
- **Creating synapses with the generator syntax**
- **Summed variables**
- **Creating multi-synapses**
- **Multiple pathways**
- **Numerical integration**
- **Technical notes**

#### 3.7.1 Defining synaptic models

The most simple synapse (adding a fixed amount to the target membrane potential on every spike) is described as follows:
w = 1 mV
S = Synapses(P, Q, on_pre='v += w')

This defines a set of synapses between NeuronGroup P and NeuronGroup Q. If the target group is not specified, it is identical to the source group by default. The on_pre keyword defines what happens when a presynaptic spike arrives at a synapse. In this case, the constant w is added to variable v. Because v is not defined as a synaptic variable, it is assumed by default that it is a postsynaptic variable, defined in the target NeuronGroup Q. Note that this does not create synapses (see Creating Synapses), only the synaptic models.

To define more complex models, models can be described as string equations, similar to the models specified in NeuronGroup:

\[ S = \text{Synapses}(P, Q, \text{model}='w : \text{volt}', \text{on}_\text{pre}='v += w') \]

The above specifies a parameter w, i.e. a synapse-specific weight. Note that to avoid confusion, synaptic variables cannot have the same name as a pre- or post-synaptic variables.

Synapses can also specify code that should be executed whenever a postsynaptic spike occurs (keyword on_post) and a fixed (pre-synaptic) delay for all synapses (keyword delay).

As shown above, variable names that are not referring to a synaptic variable are automatically understood to be post-synaptic variables. To explicitly specify that a variable should be from a pre- or post-synaptic neuron, append the suffix _pre or _post. An alternative but equivalent formulation of the on_pre statement above would therefore be v_post += w.

**Model syntax**

The model follows exactly the same syntax as for NeuronGroup. There can be parameters (e.g. synaptic variable w above), but there can also be named subexpressions and differential equations, describing the dynamics of synaptic variables. In all cases, synaptic variables are created, one value per synapse.

Brian also automatically defines a number of synaptic variables that can be used in equations, on_pre and on_post statements, as well as when assigning to other synaptic variables:

- i The index of the pre-synaptic source of a synapse.
- j The index of the post-synaptic target of a synapse.
- N The total number of synapses.
- NIncoming The total number of synapses connected to the post-synaptic target of a synapse.
- NOutgoing The total number of synapses outgoing from the pre-synaptic source of a synapse.
- lastupdate The last time this synapse has applied an on_pre or on_post statement. There is normally no need to refer to this variable explicitly, it is used to implement Event-driven updates (see below). It is only defined when event-driven equations are used.

**Event-driven updates**

By default, differential equations are integrated in a clock-driven fashion, as for a NeuronGroup. This is potentially very time consuming, because all synapses are updated at every timestep and Brian will therefore emit a warning. If you are sure about integrating the equations at every timestep (e.g. because you want to record the values continuously), then you should specify the flag (clock-driven). To ask Brian 2 to simulate differential equations in an event-driven fashion use the flag (event-driven). A typical example is pre- and postsynaptic traces in STDP:
Here, Brian updates the value of $A_{\text{pre}}$ for a given synapse only when this synapse receives a spike, whether it is presynaptic or postsynaptic. More precisely, the variables are updated every time either the `on_pre` or `on_post` code is called for the synapse, so that the values are always up to date when these codes are executed.

Automatic event-driven updates are only possible for a subset of equations, in particular for one-dimensional linear equations. These equations must also be independent of the other ones, that is, a differential equation that is not event-driven cannot depend on an event-driven equation (since the values are not continuously updated). In other cases, the user can write event-driven code explicitly in the update codes (see below).

### Pre and post codes

The `on_pre` code is executed at each synapse receiving a presynaptic spike. For example:

```python
on_pre='v+=w'
```

adds the value of synaptic variable $w$ to postsynaptic variable $v$. Any sort of code can be executed. For example, the following code defines stochastic synapses, with a synaptic weight $w$ and transmission probability $p$:

```python
S=Synapses(input,neurons,model='''w : 1
p : 1.0
on_pre="v+=w*(rand()<p)"
''')
```

The code means that $w$ is added to $v$ with probability $p$. The code may also include multiple lines.

Similarly, the `on_post` code is executed at each synapse where the postsynaptic neuron has fired a spike.

### 3.7.2 Creating synapses

Creating a `Synapses` instance does not create synapses, it only specifies their dynamics. The following command creates a synapse between neuron 5 in the source group and neuron 10 in the target group:

```python
S.connect(i=5, j=10)
```

Multiple synaptic connections can be created in a single statement:

```python
S.connect()
S.connect(i=[1, 2], j=[3, 4])
S.connect(i=numpy.arange(10), j=1)
```

The first statement connects all neuron pairs. The second statement creates synapses between neurons 1 and 3, and between neurons 2 and 4. The third statement creates synapses between the first ten neurons in the source group and neuron 1 in the target group.

### Conditional

One can also create synapses by giving (as a string) the condition for a pair of neurons $i$ and $j$ to be connected by a synapse, e.g. you could connect neurons that are not very far apart with:

```python
S.connect(condition='abs(i-j)<=5')
```
The string expressions can also refer to pre- or postsynaptic variables. This can be useful for example for spatial connectivity: assuming that the pre- and postsynaptic groups have parameters x and y, storing their location, the following statement connects all cells in a 250 um radius:

```
S.connect(condition='sqrt((x_pre-x_post)**2 + (y_pre-y_post)**2) < 250*umeter')
```

**Probabilistic**

Synapse creation can also be probabilistic by providing a `p` argument, providing the connection probability for each pair of synapses:

```
S.connect(p=0.1)
```

This connects all neuron pairs with a probability of 10%. Probabilities can also be given as expressions, for example to implement a connection probability that depends on distance:

```
S.connect(condition='i != j',
          p='p_max*exp(-(x_pre-x_post)**2+(y_pre-y_post)**2 / (2*(125*umeter)**2))')
```

If this statement is applied to a `Synapses` object that connects a group to itself, it prevents self-connections (`i != j`) and connects cells with a probability that is modulated according to a 2-dimensional Gaussian of the distance between the cells.

**One-to-one**

You can specify a mapping from i to any function f(i), e.g. the simplest way to give a 1-to-1 connection would be:

```
S.connect(j='i')
```

This mapping can also use a restricting condition with `if`, e.g. to connect neurons 0, 2, 4, 6, ... to neurons 0, 1, 2, 3, ... you could write:

```
S.connect(j='int(i/2) if i % 2 == 0')
```

### 3.7.3 Accessing synaptic variables

Synaptic variables can be accessed in a similar way as `NeuronGroup` variables. They can be indexed with two indexes, corresponding to the indexes of pre and postsynaptic neurons, or with string expressions (referring to i and j as the pre-/post-synaptic indices, or to other state variables of the synapse or the connected neurons). Note that setting a synaptic variable always refers to the synapses that currently exist, i.e. you have to set them after the relevant `Synapses.connect` call.

Here are a few examples:

```
S.w[2, 5] = 1*nS
S.w[1, : ] = 2*nS
S.w = 1*nS # all synapses assigned
S.w[2, 3] = (1*nS, 2*nS)
S.w[group1, group2] = "(1+cos(i-j))*2*nS"
S.w[:, :] = 'rand()\times nS'
S.w['abs(x_pre-x_post) < 250*umetre'] = 1*nS
```
Assignments can also refer to *pre-defined variables*, e.g., to normalize synaptic weights. For example, after the following assignment the sum of weights of all synapses that a neuron receives is identical to 1, regardless of the number of synapses it receives:

```plaintext
syn.w = '1.0/N_incoming'
```

Note that it is also possible to index synaptic variables with a single index (integer, slice, or array), but in this case synaptic indices have to be provided.

### 3.7.4 Delays

There is a special synaptic variable that is automatically created: `delay`. It is the propagation delay from the presynaptic neuron to the synapse, i.e., the presynaptic delay. This is just a convenience syntax for accessing the delay stored in the presynaptic pathway: `pre.delay`. When there is a postsynaptic code (keyword `post`), the delay of the postsynaptic pathway can be accessed as `post.delay`.

The delay variable(s) can be set and accessed in the same way as other synaptic variables. The same semantics as for other synaptic variables apply, which means in particular that the delay is only set for the synapses that have been already created with `Synapses.connect`. If you want to set a global delay for all synapses of a `Synapses` object, you can directly specify that delay as part of the `Synapses` initializer:

```plaintext
synapses = Synapses(sources, targets, '...', on_pre='...', delay=1*ms)
```

When you use this syntax, you can still change the delay afterwards by setting `synapses.delay`, but you can only set it to another scalar value. If you need different delays across synapses, do not use this syntax but instead set the delay variable as any other synaptic variable (see above).

### 3.7.5 Monitoring synaptic variables

A `StateMonitor` object can be used to monitor synaptic variables. For example, the following statement creates a monitor for variable `w` for the synapses 0 and 1:

```plaintext
M = StateMonitor(S, 'w', record=[0,1])
```

Note that these are *synapse* indices, not neuron indices. More convenient is to directly index the `Synapses` object, Brian will automatically calculate the indices for you in this case:

```plaintext
M = StateMonitor(S, 'w', record=S[0,:]) # all synapses originating from neuron 0
M = StateMonitor(S, 'w', record=S['i!=j']) # all synapses excluding autapses
M = StateMonitor(S, 'w', record=S['w>0']) # all synapses with non-zero weights (at this time)
```

You can also record a synaptic variable for all synapses by passing `record=True`.

The recorded traces can then be accessed in the usual way, again with the possibility to index the `Synapses` object:

```plaintext
plot(M.t / ms, M[S[0]].w / nS) # first synapse
plot(M.t / ms, M[S[0,:]].w / nS) # all synapses originating from neuron 0
plot(M.t / ms, M[S['w>0*nS']].w / nS) # all synapses with non-zero weights (at this time)
```

Note (for users of Brian’s advanced standalone mode only): the use of the `Synapses` object for indexing and `record=True` only work in the default runtime modes. In standalone mode (see *Standalone code generation*), the synapses have not yet been created at this point, so Brian cannot calculate the indices.
The following topics are not essential for beginners.

### 3.7.6 Synaptic connection/weight matrices

Brian does not directly support specifying synapses by using a matrix, but you always have to use a “sparse” format, where each connection is defined by its source and target indices. However, you can easily convert between the two formats. Assuming you have a connection matrix $C$ of size $N \times M$, where $N$ is the number of presynaptic cells, and $M$ the number of postsynaptic cells, with each entry being 1 for a connection, and 0 otherwise. You can convert this matrix to arrays of source and target indices, which you can then provide to Brian’s `connect` function:

```
C = ...  # The connection matrix as a numpy array of 0's and 1's
sources, targets = C.nonzero()
synapses = Synapses(...)  
synapses.connect(i=sources, j=targets)
```

Similarly, you can transform the flat array of values stored in a synapse into a matrix form. For example, to get a matrix with all the weight values $w$, with NaN values where no synapse exists:

```
synapses = Synapses(source_group, target_group,
                     '''...
                         w : 1  # synaptic weight''', ...)  
# ...
# Run e.g. a simulation with plasticity that changes the weights
run(...)  
# Create a matrix to store the weights and fill it with NaN
W = np.full((len(source_group), len(target_group)), np.nan)
# Insert the values from the Synapses object
W[synapses.i[:], synapses.j[:]] = synapses.w[:]
```

### 3.7.7 Creating synapses with the generator syntax

The most general way of specifying a connection is using the generator syntax, e.g. to connect neuron $i$ to all neurons $j$ with $0 \leq j \leq i$:

```
S.connect(j='k for k in range(0, i+1)')
```

There are several parts to this syntax. The general form is:

```
j='EXPR for VAR in RANGE if COND'
```

Here `EXPR` can be any integer-valued expression. `VAR` is the name of the iteration variable (any name you like can be specified here). The `if COND` part is optional and lets you give an additional condition that has to be true for the synapse to be created. Finally, `RANGE` can be either:

1. a Python `range`, e.g. `range(N)` is the integers from 0 to $N-1$, `range(A, B)` is the integers from $A$ to $B-1$, `range(low, high, step)` is the integers from `$low$` to `$high-1$` with steps of size `$step$`, or
2. it can be a random sample sample(N, p=0.1) gives a random sample of integers from 0 to N-1 with 10% probability of each integer appearing in the sample. This can have extra arguments like range, e.g. sample(low, high, step, p=0.1) will give each integer in range(low, high, step) with probability 10%.

If you try to create an invalid synapse (i.e. connecting neurons that are outside the correct range) then you will get an error, e.g. you might like to try to do this to connect each neuron to its neighbours:

```python
S.connect(j='i+(-1)**k for k in range(2)')
```

However this won’t work at for i=0 it gives j=-1 which is invalid. There is an option to just skip any synapses that are outside the valid range:

```python
S.connect(j='i+(-1)**k for k in range(2)', skip_if_invalid=True)
```

### 3.7.8 Summed variables

In many cases, the postsynaptic neuron has a variable that represents a sum of variables over all its synapses. This is called a “summed variable”. An example is nonlinear synapses (e.g. NMDA):

```python
neurons = NeuronGroup(1, model='''dv/dt=(gtot-v)/(10*ms) : 1
gtot : 1''')
S = Synapses(input, neurons,
model='''dg/dt=-a*g+b*x*(1-g) : 1
gtot_post = g : 1 (summed)
dx/dt=-c*x : 1
w : 1 # synaptic weight''', on_pre='x+=w')
```

Here, each synapse has a conductance $g$ with nonlinear dynamics. The neuron’s total conductance is $gtot$. The line stating $gtot_post = g : 1$ (summed) specifies the link between the two: $gtot$ in the postsynaptic group is the summer over all variables $g$ of the corresponding synapses. What happens during the simulation is that at each time step, presynaptic conductances are summed for each neuron and the result is copied to the variable $gtot$. Another example is gap junctions:

```python
neurons = NeuronGroup(N, model='''dv/dt=(v0-v+Igap)/tau : 1
Igap : 1''')
S=Synapses(neurons,model='''w:1 # gap junction conductance
Igap_post = w*(v_pre-v_post): 1 (summed)''')
```

Here, $Igap$ is the total gap junction current received by the postsynaptic neuron.

Note that you cannot target the same post-synaptic variable from more than one Synapses object. To work around this restriction, use multiple post-synaptic variables that are then summed up:

```python
neurons = NeuronGroup(1, model='''dv/dt=(gtot-v)/(10*ms) : 1
gtot = gtot1 + gtot2: 1
gtot1 : 1
gtot2 : 1''')
S1 = Synapses(input, neurons,
model='''dg/dt=-a1*g+b1*x*(1-g) : 1
gtot1_post = g : 1 (summed)
dx/dt=-c1*x : 1
w : 1 # synaptic weight''', on_pre='x+=w')
S2 = Synapses(input, neurons,
model='''dg/dt=-a2*g+b2*x*(1-g) : 1
''', on_pre='x+=w')
```

(continues on next page)
3.7.9 Creating multi-synapses

It is also possible to create several synapses for a given pair of neurons:

```python
S.connect(i=numpy.arange(10), j=1, n=3)
```

This is useful for example if one wants to have multiple synapses with different delays. To distinguish multiple variables connecting the same pair of neurons in synaptic expressions and statements, you can create a variable storing the synapse index with the `multisynaptic_index` keyword:

```python
syn = Synapses(source_group, target_group, model='w : 1', on_pre='v += w', multisynaptic_index='synapse_number')
```

```python
S.connect(i=numpy.arange(10), j=1, n=3)
```

```python
syn.delay = '1*ms + synapse_number*2*ms'
```

This index can then be used to set/get synapse-specific values:

```python
S.delay = '(synapse_number + 1)*ms)'  # Set delays between 1 and 10ms
S.w['synapse_number<5'] = 0.5
S.w['synapse_number>=5'] = 1
```

It also enables three-dimensional indexing, the following statement has the same effect as the last one above:

```python
S.w[ :, :, 5:] = 1
```

3.7.10 Multiple pathways

It is possible to have multiple pathways with different update codes from the same presynaptic neuron group. This may be interesting in cases when different operations must be applied at different times for the same presynaptic spike, e.g. for a STDP rule that shifted in time. To do this, specify a dictionary of pathway names and codes:

```python
on_pre={'pre_transmission': 'ge+=w',
       'pre_plasticity': '''w=clip(w+Apost,0,inf)
Apre+=dApre'''}
```

This creates two pathways with the given names (in fact, specifying `on_pre=code` is just a shorter syntax for `on_pre={'pre': code}`) through which the delay variables can be accessed. The following statement, for example, sets the delay of the synapse between the first neurons of the source and target groups in the `pre_plasticity` pathway:

```python
S.pre_plasticity.delay[0,0] = 3*ms
```

As mentioned above, `pre` pathways are generally executed before `post` pathways. The order of execution of several `pre` (or `post`) pathways with the same delay is however arbitrary, and simply based on the alphabetical ordering of their names (i.e. `pre_plasticity` will be executed before `pre_transmission`). To explicitly specify the order, set the `order` attribute of the pathway, e.g.:
S.pre_transmission.order = -2

will make sure that the pre_transmission code is executed before the pre_plasticity code in each time step.

Multiple pathways can also be useful for abstract models of synaptic currents, e.g. modelling them as rectangular currents:

```python
synapses = Synapses(...,
    on_pre={'up': 'I_syn_post += 1*nA',
            'down': 'I_syn_post -= 1*nA'},
    delays={'up': 0*ms, 'down': 5*ms}  # 5ms-wide rectangular current
)
```

### 3.7.11 Numerical integration

#### Differential equation flags

For the integration of differential equations, one can use the same keywords as for `NeuronGroup`.

**Note:** Declaring a subexpression as *(constant over dt)* means that it will be evaluated each timestep for all synapses, potentially a very costly operation.

#### Explicit event-driven updates

As mentioned above, it is possible to write event-driven update code for the synaptic variables. This can also be done manually, by defining the variable `lastupdate` and referring to the predefined variable `t` (current time). Here’s an example for short-term plasticity – but note that using the automatic event-driven approach from above is usually preferable:

```python
S=Synapses(input,neuron,
    model='''x : 1
    u : 1
    w : 1
    lastupdate : second'','
    on_pre='''u=U+(u-U)*exp(-(t-lastupdate)/tauf)
    x=1+(x-1)*exp(-(t-lastupdate)/taud)
    i+=w*u*x
    x*=(1-u)
    u+=U*(1-u)
    lastupdate = t'''
)
```

By default, the `pre` pathway is executed before the `post` pathway (both are executed in the 'synapses' scheduling slot, but the `pre` pathway has the `order` attribute -1, whereas the `post` pathway has `order` 1. See `Scheduling` for more details).

### 3.7.12 Technical notes
How connection arguments are interpreted

If conditions for connecting neurons are combined with both the \( n \) (number of synapses to create) and the \( p \) (probability of a synapse) keywords, they are interpreted in the following way:

For every pair \( i, j \):
  - if condition\((i, j)\) is fulfilled:
    - Evaluate \( p(i, j) \)
    - If uniform random number between 0 and 1 < \( p(i, j) \):
      - Create \( n(i, j) \) synapses for \((i, j)\)

With the generator syntax \( j='EXPR\ for\ VAR\ in\ RANGE\ if\ COND' \), the interpretation is:

For every \( i \):
  - for every \( VAR\ in\ RANGE\):
    - \( j = EXPR \)
    - if \( COND \):
      - Create \( n(i, j) \) synapses for \((i, j)\)

Note that the arguments in \( RANGE \) can only depend on \( i \) and the values of presynaptic variables. Similarly, the expression for \( j, EXPR \) can depend on \( i \), presynaptic variables, and on the iteration variable \( VAR \). The condition \( COND \) can depend on anything (presynaptic and postsynaptic variables).

With the 1-to-1 mapping syntax \( j='EXPR'\) the interpretation is:

For every \( i \):
  - \( j = EXPR \)
  - Create \( n(i, j) \) synapses for \((i, j)\)

Efficiency considerations

If you are connecting a single pair of neurons, the direct form \( \text{connect}(i=5, j=10) \) is the most efficient. However, if you are connecting a number of neurons, it will usually be more efficient to construct an array of \( i \) and \( j \) values and have a single \( \text{connect}(i=i, j=j) \) call.

For large connections, you should use one of the string based syntaxes where possible as this will generate compiled low-level code that will be typically much faster than equivalent Python code.

If you are expecting a majority of pairs of neurons to be connected, then using the condition-based syntax is optimal, e.g. \( \text{connect}(\text{condition}='i!=j') \). However, if relatively few neurons are being connected then the 1-to-1 mapping or generator syntax will be better. For 1-to-1, \( \text{connect}(j='i') \) will always be faster than \( \text{connect}(\text{condition}='i==j') \) because the latter has to evaluate all \( N\times2 \) pairs \((i, j)\) and check if the condition is true, whereas the former only has to do \( O(N) \) operations.

One tricky problem is how to efficiently generate connectivity with a probability \( p(i, j) \) that depends on both \( i \) and \( j \), since this requires \( N\times N \) computations even if the expected number of synapses is proportional to \( N \). Some tricks for getting around this are shown in Example: efficient_gaussian_connectivity.

3.8 Input stimuli

For Brian 1 users

See the document Inputs (Brian 1 -> 2 conversion) for details how to convert Brian 1 code.
There are various ways of providing “external” input to a network.

### 3.8.1 Poisson inputs

For generating spikes according to a Poisson point process, `PoissonGroup` can be used, e.g.:

```python
P = PoissonGroup(100, np.arange(100)*Hz + 10*Hz)
G = NeuronGroup(100, 'dv/dt = -v / (10*ms) : 1')
S = Synapses(P, G, on_pre='v+=0.1')
S.connect(j='i')
```

See *More on Poisson inputs* below for further information.

For simulations where the individually generated spikes are just used as a source of input to a neuron, the `PoissonInput` class provides a more efficient alternative: see *Efficient Poisson inputs via PoissonInput* below for details.

### 3.8.2 Spike generation

You can also generate an explicit list of spikes given via arrays using `SpikeGeneratorGroup`. This object behaves just like a `NeuronGroup` in that you can connect it to other groups via a `Synapses` object, but you specify three bits of information: `N` the number of neurons in the group; `indices` an array of the indices of the neurons that will fire; and `times` an array of the same length as `indices` with the times that the neurons will fire a spike. The `indices` and `times` arrays are matching, so for example `indices=[0, 2, 1]` and `times=[1*ms, 2*ms, 3*ms]` means that neuron 0 fires at time 1 ms, neuron 2 fires at 2 ms and neuron 1 fires at 3 ms. Example use:

```python
indices = array([0, 2, 1])
times = array([1, 2, 3])*ms
G = SpikeGeneratorGroup(3, indices, times)
```

The spikes that will be generated by `SpikeGeneratorGroup` can be changed between runs with the `set_spikes` method. This can be useful if the input to a system should depend on its previous output or when running multiple trials with different input:

```python
inp = SpikeGeneratorGroup(N, indices, times)
G = NeuronGroup(N, '...')
feedforward = Synapses(inp, G, '...', on_pre='...')
feedforward.connect(j='i')
recurrent = Synapses(G, G, '...', on_pre='...')
recurrent.connect('i!=j')
spike_mon = SpikeMonitor(G)
```
3.8.3 Explicit equations

If the input can be explicitly expressed as a function of time (e.g., a sinusoidal input current), then its description can be directly included in the equations of the respective group:

```brian2
G = NeuronGroup(100, '''
    dv/dt = (-v + I)/(10*ms) : 1
    rates : Hz # each neuron's input has a different rate
    size : 1  # and a different amplitude
    I = size*sin(2*pi*rates*t) : 1'''
    
    G.rates = '10*Hz + i*Hz'
    G.size = '(100-i)/100. + 0.1'
```

3.8.4 Timed arrays

If the time dependence of the input cannot be expressed in the equations in the way shown above, it is possible to create a `TimedArray`. This acts as a function of time where the values at given time points are given explicitly. This can be especially useful to describe non-continuous stimulation. For example, the following code defines a `TimedArray` where stimulus blocks consist of a constant current of random strength for 30ms, followed by no stimulus for 20ms. Note that in this particular example, numerical integration can use exact methods, since it can assume that the `TimedArray` is a constant function of time during a single integration time step.

```python
stimulus = TimedArray(np.hstack([[c, c, c, 0, 0]
                                 for c in np.random.rand(1000)]),
                       dt=10*ms)
G = NeuronGroup(100, 'dv/dt = (-v + stimulus(t))/(10*ms) : 1',
                threshold='v>1', reset='v=0')
G.v = '0.5*rand()'  # different initial values for the neurons
```

TimedArray can take a one-dimensional value array (as above) and therefore return the same value for all neurons or it can take a two-dimensional array with time as the first and (neuron/synapse/...) index as the second dimension.

In the following, this is used to implement shared noise between neurons, all the “even neurons” get the first noise instantiation, all the “odd neurons” get the second:

```python
runtime = 1*second
stimulus = TimedArray(np.random.rand(int(runtime/defaultclock.dt), 2),
                       dt=defaultclock.dt)
G = NeuronGroup(100, 'dv/dt = (-v + stimulus(t, i % 2))/(10*ms) : 1',
                threshold='v>1', reset='v=0')
```
3.8.5 Regular operations

An alternative to specifying a stimulus in advance is to run explicitly specified code at certain points during a simulation. This can be achieved with `run_regularly()`. One can think of these statements as equivalent to reset statements but executed unconditionally (i.e. for all neurons) and possibly on a different clock than the rest of the group. The following code changes the stimulus strength of half of the neurons (randomly chosen) to a new random value every 50ms. Note that the statement uses logical expressions to have the values only updated for the chosen subset of neurons (where the newly introduced auxiliary variable `change` equals 1):

```python
G = NeuronGroup(100, '''
  dv/dt = (-v + I)/(10*ms) : 1
  I : 1  # one stimulus per neuron'''
)
G.run_regularly('''
  change = int(rand() < 0.5)
  I = change*(rand()*2) + (1-change)*I'','
  dt=50*ms)
```

The following topics are not essential for beginners.

3.8.6 More on Poisson inputs

Setting rates for Poisson inputs

`PoissonGroup` takes either a constant rate, an array of rates (one rate per neuron, as in the example above), or a string expression evaluating to a rate as an argument.

If the given value for `rates` is a constant, then using `PoissonGroup(N, rates)` is equivalent to:

```python
NeuronGroup(N, 'rates : Hz', threshold='rand()<rates*dt')
```

and setting the group's `rates` attribute.

If `rates` is a string, then this is equivalent to:

```python
NeuronGroup(N, 'rates = ... : Hz', threshold='rand()<rates*dt')
```

with the respective expression for the rates. This expression will be evaluated at every time step and therefore allows the use of time-dependent rates, i.e. inhomogeneous Poisson processes. For example, the following code (see also `Timed arrays`) uses a `TimedArray` to define the rates of a `PoissonGroup` as a function of time, resulting in five 100ms blocks of 100 Hz stimulation, followed by 100ms of silence:

```python
stimulus = TimedArray(np.tile([100.0, 0.], 5)*Hz, dt=100.*ms)
P = PoissonGroup(1, rates='stimulus(t)')
```

Note that, as can be seen in its equivalent `NeuronGroup` formulation, a `PoissonGroup` does not work for high rates where more than one spike might fall into a single timestep. Use several units with lower rates in this case (e.g. use `PoissonGroup(10, 1000*Hz)` instead of `PoissonGroup(1, 10000*Hz)`).
Efficient Poisson inputs via PoissonInput

For simulations where the PoissonGroup is just used as a source of input to a neuron (i.e., the individually generated spikes are not important, just their impact on the target cell), the PoissonInput class provides a more efficient alternative: instead of generating spikes, PoissonInput directly updates a target variable based on the sum of independent Poisson processes:

```python
G = NeuronGroup(100, 'dv/dt = -v / (10*ms) : 1')
P = PoissonInput(G, 'v', 100, 100*Hz, weight=0.1)
```

The PoissonInput class is however more restrictive than PoissonGroup, it only allows for a constant rate across all neurons (but you can create several PoissonInput objects, targeting different subgroups). It internally uses BinomialFunction which will draw a random number each time step, either from a binomial distribution or from a normal distribution as an approximation to the binomial distribution if \( np > 5 \land n(1 - p) > 5 \), where \( n \) is the number of inputs and \( p = dt \cdot rate \) the spiking probability for a single input.

### 3.8.7 Arbitrary Python code (network operations)

If none of the above techniques is general enough to fulfill the requirements of a simulation, Brian allows you to write a NetworkOperation, an arbitrary Python function that is executed every time step (possible on a different clock than the rest of the simulation). This function can do arbitrary operations, use conditional statements etc. and it will be executed as it is (i.e. as pure Python code even if weave code generation is active). Note that one cannot use network operations in combination with the C++ standalone mode. Network operations are particularly useful when some condition or calculation depends on operations across neurons, which is currently not possible to express in abstract code. The following code switches input on for a randomly chosen single neuron every 50 ms:

```python
G = NeuronGroup(10, '''dv/dt = (-v + active*I)/(10*ms) : 1
I = sin(2*pi*100*Hz*t) : 1 (shared) # single input
active : 1 # will be set in the network operation''')
@network_operation(dt=50*ms)
def update_active():
    index = np.random.randint(10)
    G.active_ = 0  # the underscore switches off unit checking
    G.active_[index] = 1
```

Note that the network operation (in the above example: update_active) has to be included in the Network object if one is constructed explicitly.

Only functions with zero or one arguments can be used as a NetworkOperation. If the function has one argument then it will be passed the current time \( t \):

```python
@network_operation(dt=1*ms)
def update_input(t):
    if t>50*ms and t<100*ms:
        pass  # do something
```

Note that this is preferable to accessing defaultclock.t from within the function – if the network operation is not running on the defaultclock itself, then that value is not guaranteed to be correct.

Instance methods can be used as network operations as well, however in this case they have to be constructed explicitly, the network_operation() decorator cannot be used:

```python
class Simulation(object):
    def __init__(self, data):
        self.data = data
```

(continues on next page)
3.9 Recording during a simulation

For Brian 1 users

See the document Monitors (Brian 1 –> 2 conversion) for details how to convert Brian 1 code.

- Recording spikes
- Recording variables at spike time
- Recording variables continuously
- Recording population rates
- Getting all data
- Recording values for a subset of the run
- Freeing up memory in long recordings
- Recording random subsets of neurons

Recording variables during a simulation is done with “monitor” objects. Specifically, spikes are recorded with `SpikeMonitor`, the time evolution of variables with `StateMonitor` and the firing rate of a population of neurons with `PopulationRateMonitor`.

3.9.1 Recording spikes

To record spikes from a group `G` simply create a `SpikeMonitor` via `SpikeMonitor(G)`. After the simulation, you can access the attributes `i`, `t`, `num_spikes` and `count` of the monitor. The `i` and `t` attributes give the array of neuron indices and times of the spikes. For example, if `M.i==[0, 2, 1]` and `M.t==[1*ms, 2*ms, 3*ms]` it means that neuron 0 fired a spike at 1 ms, neuron 2 fired a spike at 2 ms, and neuron 1 fired a spike at 3 ms. Alternatively, you can also call the `spike_trains` method to get a dictionary mapping neuron indices to arrays of spike times, i.e. in the above example, `spike_trains = M.spike_trains(); spike_trains[1]` would return `array([ 3.]) * msecond`. The `num_spikes` attribute gives the total number of spikes recorded, and `count` is an array of the length of the recorded group giving the total number of spikes recorded from each neuron.

Example:

```python
G = NeuronGroup(N, model='...')
M = SpikeMonitor(G)
```
run(runtime)
plot(M.t/ms, M.i, '.')

If you are only interested in summary statistics but not the individual spikes, you can set the `record` argument to `False`. You will then not have access to `i` and `t` but you can still get the `count` and the total number of spikes (`num_spikes`).

### 3.9.2 Recording variables at spike time

By default, a `SpikeMonitor` only records the time of the spike and the index of the neuron that spiked. Sometimes it can be useful to additionally record other variables, e.g. the membrane potential for models where the threshold is not at a fixed value. This can be done by providing an extra `variables` argument, the recorded variable can then be accessed as an attribute of the `SpikeMonitor`, e.g.:

```python
G = NeuronGroup(10, 'v : 1', threshold='rand()<100*Hz*dt')
G.run_regularly('v = rand()')
M = SpikeMonitor(G, variables=['v'])
run(100*ms)
plot(M.t/ms, M.v, '.')
```

To conveniently access the values of a recorded variable for a single neuron, the `SpikeMonitor.values` method can be used that returns a dictionary with the values for each neuron:

```python
G = NeuronGroup(N, '''dv/dt = (1-v)/(10*ms) : 1
v_th : 1''',
    threshold='v > v_th',
    reset='''v=0
v_th = clip(v_th + rand()*0.2 - 0.1, 0.1, 0.9)''')
G.v_th = 0.5
spike_mon = SpikeMonitor(G, variables='v')
run(1*second)
print('Threshold crossing values for neuron 0: {}'.format(v_values[0]))
hist(spike_mon.v, np.arange(0, 1, .1))
show()
```

**Note:** Spikes are not the only events that can trigger recordings, see *Custom events*.

### 3.9.3 Recording variables continuously

To record how a variable evolves over time, use a `StateMonitor`, e.g. to record the variable `v` at every time step and plot it for neuron 0:

```python
G = NeuronGroup(...)
M = StateMonitor(G, 'v', record=True)
run(...)  
plot(M.t/ms, M.v[0]/mV)
```

In general, you specify the group, variables and indices you want to record from. You specify the variables with a string or list of strings, and the indices either as an array of indices or `True` to record all indices (but beware because this may take a lot of memory).
After the simulation, you can access these variables as attributes of the monitor. They are 2D arrays with shape (num_indices, num_times). The special attribute \( t \) is an array of length num_times with the corresponding times at which the values were recorded.

Note that you can also use `StateMonitor` to record from `Synapses` where the indices are the synapse indices rather than neuron indices.

In this example, we record two variables \( v \) and \( u \), and record from indices 0, 10 and 100. Afterwards, we plot the recorded values of \( v \) and \( u \) from neuron 0:

```python
G = NeuronGroup(...)
M = StateMonitor(G, ('v', 'u'), record=[0, 10, 100])
run(...)
plot(M.t/ms, M.v[0]/mV, label='v')
plot(M.t/ms, M.u[0]/mV, label='u')
```

There are two subtly different ways to get the values for specific neurons: you can either index the 2D array stored in the attribute with the variable name (as in the example above) or you can index the monitor itself. The former will use an index relative to the recorded neurons (e.g. \( M.v[1] \) will return the values for the second recorded neuron which is the neuron with the index 10 whereas \( M.v[10] \) would raise an error because only three neurons have been recorded), whereas the latter will use an absolute index corresponding to the recorded group (e.g. \( M[1].v \) will raise an error because the neuron with the index 1 has not been recorded and \( M[10].v \) will return the values for the neuron with the index 10). If all neurons have been recorded (e.g. with `record=True`) then both forms give the same result.

Note that for plotting all recorded values at once, you have to transpose the variable values:

```python
plot(M.t/ms, M.v.T/mV)
```

Note: In contrast to Brian 1, the values are recorded at the beginning of a time step and not at the end (you can set the `when` argument when creating a `StateMonitor`, details about scheduling can be found here: Custom progress reporting).

### 3.9.4 Recording population rates

To record the time-varying firing rate of a population of neurons use `PopulationRateMonitor`. After the simulation the monitor will have two attributes \( t \) and \( rate \), the latter giving the firing rate at each time step corresponding to the time in \( t \). For example:

```python
G = NeuronGroup(...)
M = PopulationRateMonitor(G)
run(...)
plot(M.t/ms, M.rate/Hz)
```

To get a smoother version of the rate, use `PopulationRateMonitor.smooth_rate`.

The following topics are not essential for beginners.
3.9.5 Getting all data

Note that all monitors are implement as “groups”, so you can get all the stored values in a monitor with the `get_states` method, which can be useful to dump all recorded data to disk, for example:

```python
import pickle
group = NeuronGroup(...)
state_mon = StateMonitor(group, 'v', record=...)
run(...)
data = state_mon.get_states(['t', 'v'])
with open('state_mon.pickle', 'w') as f:
    pickle.dump(data, f)
```

3.9.6 Recording values for a subset of the run

Monitors can be created and deleted between runs, e.g. to ignore the first second of your simulation in your recordings you can do:

```python
# Set up network without monitor
run(1*second)
state_mon = StateMonitor(...)  
run(...)  # Continue run and record with the StateMonitor
```

Alternatively, you can set the monitor’s `active` attribute as explained in the `Scheduling` section.

3.9.7 Freeing up memory in long recordings

Creating and deleting monitors can also be useful to free memory during a long recording. The following will do a simulation run, dump the monitor data to disk, delete the monitor and finally continue the run with a new monitor:

```python
import pickle
# Set up network
state_mon = StateMonitor(...)  
run(...)  # a long run
data = state_mon.get_states(...)  
with open('first_part.data', 'w') as f:
    pickle.dump(data, f)

del state_mon

del data
state_mon = StateMonitor(...)  
run(...)  # another long run
```

Note that this technique cannot be applied in `standalone mode`.

3.9.8 Recording random subsets of neurons

In large networks, you might only be interested in the activity of a random subset of neurons. While you can specify a record argument for a `StateMonitor` that allows you to select a subset of neurons, this is not possible for `SpikeMonitor/EventMonitor` and `PopulationRateMonitor`. However, Brian allows you to record with these monitors from a subset of neurons by using a `subgroup`:

```python

group = NeuronGroup(1000, ...)  
spike_mon = SpikeMonitor(group[::100])  # only record first 100 neurons
```
It might seem like a restriction that such a subgroup has to be contiguous, but the order of neurons in a group does not have any meaning as such: in a randomly ordered group of neurons, any contiguous group of neurons can be considered a random subset. If some aspects of your model do depend on the position of the neuron in a group (e.g. a ring model, where neurons are connected based on their distance in the ring, or a model where initial values or parameters span a range of values in a regular fashion), then this requires an extra step: instead of using the order of neurons in the group directly, or depending on the neuron index \(i\), create a new, shuffled, index variable as part of the model definition and then depend on this index instead:

```python
group = NeuronGroup(10000, '''
    index : integer (constant)'''
)
indices = group.i[:]
np.random.shuffle(indices)
group.index = indices
# Then use 'index' in string expressions or use it as an index array
# for initial values/parameters defined as numpy arrays
```

If this solution is not feasible for some reason, there is another approach that works for a `SpikeMonitor/EventMonitor`. You can add an additional flag to each neuron, stating whether it should be recorded or not. Then, you define a new custom event that is identical to the event you are interested in, but additionally requires the flag to be set. E.g. to only record the spikes of neurons with the `to_record` attribute set:

```python
group = NeuronGroup(..., '''
    to_record : boolean (constant)''',
    threshold='...', reset='...',
    events={'recorded_spike': '...', and to_record'})
group.to_record = ...
mon_events = EventMonitor(group, 'recorded_spike')
```

Note that this solution will evaluate the threshold condition for each neuron twice, and is therefore slightly less efficient. There's one additional caveat: you'll have to manually include and `not refractory` in your events definition if your neuron uses refractoriness. This is done automatically for the threshold condition, but not for any user-defined events.

### 3.10 Running a simulation

For Brian 1 users

See the document *Networks and clocks (Brian 1 -> 2 conversion)* for details how to convert Brian 1 code.
To run a simulation, one either constructs a new Network object and calls its Network.run method, or uses the “magic” system and a plain run() call, collecting all the objects in the current namespace.

Note that Brian has several different ways of running the actual computations, and choosing the right one can make orders of magnitude of difference in terms of simplicity and efficiency. See Computational methods and efficiency for more details.

### 3.10.1 Networks

In most straightforward simulations, you do not have to explicitly create a Network object but instead can simply call run() to run a simulation. This is what is called the “magic” system, because Brian figures out automatically what you want to do.

When calling run(), Brian runs the collect() function to gather all the objects in the current context. It will include all the objects that are “visible”, i.e. that you could refer to with an explicit name:

```python
G = NeuronGroup(10, 'dv/dt = -v / tau : volt')
S = Synapses(G, G, model='w:1', on_pre='v+=w')
S.connect('i!=j')
mon = SpikeMonitor(G)
run(10*ms)  # will include G, S, mon
```

Note that it will not automatically include objects that are “hidden” in containers, e.g. if you store several monitors in a list. Use an explicit Network object in this case. It might be convenient to use the collect() function when creating the Network object in that case:

```python
G = NeuronGroup(10, 'dv/dt = -v / tau : volt')
S = Synapses(G, G, model='w:1', on_pre='v+=w')
S.connect('i!=j')
monitors = [SpikeMonitor(G), StateMonitor(G, 'v', record=True)]
# a simple run would not include the monitors
net = Network(collect())  # automatically include G and S
net.add(monitors)  # manually add the monitors
net.run(10*ms)
```

### 3.10.2 Setting the simulation time step

To set the simulation time step for every simulated object, set the dt attribute of the defaultclock which is used by all objects that do not explicitly specify a clock or dt value during construction:

```python
defaultclock.dt = 0.05*ms
```

If some objects should use a different clock (e.g. to record values with a StateMonitor not at every time step in a long running simulation), you can provide a dt argument to the respective object:

```python
s_mon = StateMonitor(group, 'v', record=True, dt=1*ms)
```

To sum up:

- Set defaultclock.dt to the time step that should be used by most (or all) of your objects.
• Set dt explicitly when creating objects that should use a different time step.

Behind the scenes, a new Clock object will be created for each object that defines its own dt value.

3.10.3 Progress reporting

Especially for long simulations it is useful to get some feedback about the progress of the simulation. Brian offers a few built-in options and an extensible system to report the progress of the simulation. In the Network.run or run() call, two arguments determine the output: report and report_period. When report is set to 'text' or 'stderr', the progress will be printed to the standard output, when it is set to 'stderr', it will be printed to “standard error”. There will be output at the start and the end of the run, and during the run in report_period intervals. It is also possible to do custom progress reporting.

3.10.4 Continuing/repeating simulations

To store the current state of the simulation, call store() (use the Network.store method for a Network). You can store more than one snapshot of a system by providing a name for the snapshot; if store() is called without a specified name, 'default' is used as the name. To restore the state, use restore().

The following simple example shows how this system can be used to run several trials of an experiment:

```python
# set up the network
G = NeuronGroup(...)
...
spike_monitor = SpikeMonitor(G)

# Snapshots the state
store()

# Run the trials
spike_counts = []
for trial in range(3):
    restore()  # Restore the initial state
    run(...)
    # store the results
    spike_counts.append(spike_monitor.count)
```

The following schematic shows how multiple snapshots can be used to run a network with a separate “train” and “test” phase. After training, the test is run several times based on the trained network. The whole process of training and testing is repeated several times as well:

```python
# set up the network
G = NeuronGroup(..., '''
    test_input : amp
    ...'''
S = Synapses(..., '''
    plastic : boolean {shared}
    ...'''
G.v = ...
S.connect(...)
S.w = ...

# First snapshot at t=0
store('initialized')
```

(continues on next page)
# Run 3 complete trials
for trial in range(3):
    # Simulate training phase
    restore('initialized')
    S.plastic = True
    run(...)

    # Snapshot after learning
    store('after_learning')

# Run 5 tests after the training
for test_number in range(5):
    restore('after_learning')
    S.plastic = False  # switch plasticity off
    G.test_input = test_inputs[test_number]
    # monitor the activity now
    spike_mon = SpikeMonitor(G)
    run(...)
    # Do something with the result
    # ...

The following topics are not essential for beginners.

### 3.10.5 Multiple magic runs

When you use more than a single `run()` statement, the magic system tries to detect which of the following two situations applies:

1. You want to continue a previous simulation
2. You want to start a new simulation

For this, it uses the following heuristic: if a simulation consists only of objects that have not been run, it will start a new simulation starting at time 0 (corresponding to the creation of a new `Network` object). If a simulation only consists of objects that have been simulated in the previous `run()` call, it will continue that simulation at the previous time.

If neither of these two situations apply, i.e., the network consists of a mix of previously run objects and new objects, an error will be raised. If this is not a mistake but intended (e.g. when a new input source and synapses should be added to a network at a later stage), use an explicit `Network` object.

In these checks, “non-invalidating” objects (i.e. objects that have `BrianObject.invalidates_magic_network` set to `False`) are ignored, e.g. creating new monitors is always possible.

Note that if you do not want to run an object for the complete duration of your simulation, you can create the object in the beginning of your simulation and then set its `active` attribute. For details, see the `Scheduling` section below.
3.10.6 Changing the simulation time step

You can change the simulation time step after objects have been created or even after a simulation has been run:

```python
defaultclock.dt = 0.1*ms
# Set the network
# ...
run(initial_time)
defaultclock.dt = 0.01*ms
run(full_time - initial_time)
```

To change the time step between runs for objects that do not use the `defaultclock`, you cannot directly change their `dt` attribute (which is read-only) but instead you have to change the `dt` of the `clock` attribute. If you want to change the `dt` value of several objects at the same time (but not for all of them, i.e. when you cannot use `defaultclock.dt`) then you might consider creating a `Clock` object explicitly and then passing this clock to each object with the `clock` keyword argument (instead of `dt`). This way, you can later change the `dt` for several objects at once by assigning a new value to `Clock.dt`.

Note that a change of `dt` has to be compatible with the internal representation of clocks as an integer value (the number of elapsed time steps). For example, you can simulate an object for 100ms with a time step of 0.1ms (i.e. for 1000 steps) and then switch to a `dt` of 0.5ms, the time will then be internally represented as 200 steps. You cannot, however, switch to a `dt` of 0.3ms, because 100ms are not an integer multiple of 0.3ms.

3.10.7 Profiling

To get an idea which parts of a simulation take the most time, Brian offers a basic profiling mechanism. If a simulation is run with the `profile=True` keyword argument, it will collect information about the total simulation time for each `CodeObject`. This information can then be retrieved from `Network.profiling_info`, which contains a list of `(name, time)` tuples or a string summary can be obtained by calling `profiling_summary()`. The following example shows profiling output after running the CUBA example (where the neuronal state updates take up the most time):

```bash
>>> profiling_summary(show=5)  # show the 5 objects that took the longest
Profiling summary
============
neurongroup_stateupdater  5.54 s   61.32 %
synapses_pre   1.39 s   15.39 %
synapses_1_pre 1.03 s   11.37 %
spikemonitor  0.59 s   6.55 %
neurongroup_thresholder  0.33 s   3.66 %
```

3.10.8 Scheduling

Every simulated object in Brian has three attributes that can be specified at object creation time: `dt`, `when`, and `order`. The time step of the simulation is determined by `dt`, if it is specified, or otherwise by `defaultclock.dt`. Changing this will therefore change the `dt` of all objects that don’t specify one. Alternatively, a `clock` object can be specified directly, this can be useful if a clock should be shared between several objects – under most circumstances, however, a user should not have to deal with the creation of `Clock` objects and just define `dt`.

During a single time step, objects are updated in an order according first to their `when` argument’s position in the schedule. This schedule is determined by `Network.schedule` which is a list of strings, determining “execution slots” and their order. It defaults to: `['start', 'groups', 'thresholds', 'synapses', 'resets', 'end']`. In addition to the names provided in the schedule, names such as `before_thresholds` or `after_synapses` can be used that are understood as slots in the respective positions. The default for the `when`
attribute is a sensible value for most objects (resets will happen in the reset slot, etc.) but sometimes it make sense to change it, e.g. if one would like a StateMonitor, which by default records in the start slot, to record the membrane potential before a reset is applied (otherwise no threshold crossings will be observed in the membrane potential traces).

Finally, if during a time step two objects fall in the same execution slot, they will be updated in ascending order according to their order attribute, an integer number defaulting to 0. If two objects have the same when and order attribute then they will be updated in an arbitrary but reproducible order (based on the lexicographical order of their names).

Note that objects that don’t do any computation by themselves but only act as a container for other objects (e.g. a NeuronGroup which contains a StateUpdater, a Resetter and a Thresholder), don’t have any value for when, but pass on the given values for dt and order to their containing objects.

If you want your simulation object to run only for a particular time period of the whole simulation, you can use the active attribute. For example, this can be useful when you want a monitor to be active only for some time out of a long simulation:

```python
# Set up the network
# ...
monitor = SpikeMonitor(...)
monitor.active = False
run(long_time*seconds)  # not recording
monitor.active = True
run(required_time*seconds)  # recording
```

To see how the objects in a network are scheduled, you can use the scheduling_summary() function:

```python
>>> group = NeuronGroup(10, 'dv/dt = -v/(10*ms) : 1', threshold='v > 1',
...                      reset='v = 0')
>>> mon = StateMonitor(group, 'v', record=True, dt=1*ms)
>>> scheduling_summary()

<table>
<thead>
<tr>
<th>object</th>
<th>part of</th>
<th>Clock</th>
</tr>
</thead>
<tbody>
<tr>
<td>dt</td>
<td></td>
<td></td>
</tr>
<tr>
<td>when</td>
<td></td>
<td></td>
</tr>
<tr>
<td>order</td>
<td></td>
<td></td>
</tr>
<tr>
<td>active</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------------------+-------------------+-------</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
| statemonitor (StateMonitor) | statemonitor (StateMonitor) | 1. ms (every
-10 steps) | start | 0 | yes
| neurongroup_stateupdater (StateUpdater) | neurongroup (NeuronGroup) | 100. us |
| neurongroup_thresholder (Threshold) | neurongroup (NeuronGroup) | 100. us |
| neurongroup_resetter (Reseter) | neurongroup (NeuronGroup) | 100. us |
```

As you can see in the output above, the StateMonitor will only record the membrane potential every 10 time steps, but when it does, it will do it at the start of the time step, before the numerical integration, the thresholding, and the reset operation takes place.

Every new Network starts a simulation at time 0; Network.t is a read-only attribute, to go back to a previous moment in time (e.g. to do another trial of a simulation with a new noise instantiation) use the mechanism described below.

### 3.10.9 Store/restore

Note that Network.run, Network.store and Network.restore (or run(), store(), restore()) are the only way of affecting the time of the clocks. In contrast to Brian1, it is no longer necessary (nor possible) to directly set the time of the clocks or call a reinit function.
The state of a network can also be stored on disk with the optional filename argument of \texttt{Network.store()} and \texttt{restore()}. This way, you can run the initial part of a simulation once, store it to disk, and then continue from this state later. Note that the \texttt{store()/restore()} mechanism does not re-create the network as such, you still need to construct all the \texttt{NeuronGroup, Synapses, StateMonitor, ...} objects, restoring will only re-store all the state variable values (membrane potential, conductances, synaptic connections/weights/delays, ...). This restoration does however restore the internal state of the objects as well, e.g. spikes that have not been delivered yet because of synaptic delays will be delivered correctly.

### 3.11 Multicompartment models

**For Brian 1 users**

See the document \textit{Multicompartmental models (Brian 1 -> 2 conversion)} for details how to convert Brian 1 code.

It is possible to create neuron models with a spatially extended morphology, using the \texttt{SpatialNeuron} class. A \texttt{SpatialNeuron} is a single neuron with many compartments. Essentially, it works as a \texttt{NeuronGroup} where elements are compartments instead of neurons.

A \texttt{SpatialNeuron} is specified by a morphology (see \textit{Creating a neuron morphology}) and a set of equations for transmembrane currents (see \textit{Creating a spatially extended neuron}).

#### 3.11.1 Creating a neuron morphology

**Schematic morphologies**

Morphologies can be created combining geometrical objects:

```python
soma = Soma(diameter=30*um)
cylinder = Cylinder(diameter=1*um, length=100*um, n=10)
```

The first statement creates a single iso-potential compartment (i.e. with no axial resistance within the compartment), with its area calculated as the area of a sphere with the given diameter. The second one specifies a cylinder consisting of 10 compartments with identical diameter and the given total length.

For more precise control over the geometry, you can specify the length and diameter of each individual compartment, including the diameter at the start of the section (i.e. for \(n\) compartments: \(n\) length and \(n+1\) diameter values) in a \texttt{Section} object:

```python
section = Section(diameter=[6, 5, 4, 3, 2, 1]*um, length=[10, 10, 10, 5, 5]*um, n=5)
```

The individual compartments are modeled as truncated cones, changing the diameter linearly between the given diameters over the length of the compartment. Note that the \texttt{diameter} argument specifies the values at the nodes between the compartments, but accessing the \texttt{diameter} attribute of a \texttt{Morphology} object will return the diameter at the center of the compartment (see the note below).

The following table summarizes the different options to create schematic morphologies (the black compartment before the start of the section represents the parent compartment with diameter 15 \(\mu\)m, not specified in the code below):
<table>
<thead>
<tr>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>Soma</td>
</tr>
<tr>
<td># Soma always has a single compartment</td>
</tr>
<tr>
<td>Soma(diameter=30*um)</td>
</tr>
<tr>
<td>Cylinder</td>
</tr>
<tr>
<td># Each compartment has fixed length and diameter</td>
</tr>
<tr>
<td>Cylinder(n=5, diameter=10*um, length=50*um)</td>
</tr>
<tr>
<td>Section</td>
</tr>
<tr>
<td># Length and diameter individually defined for each compartment (at start and end)</td>
</tr>
<tr>
<td>Section(n=5, diameter=[15, 5, 10, 5, 10, 5]*um, length=[10, 20, 5, 5, 10]*um)</td>
</tr>
</tbody>
</table>

**Note:** For a **Section**, the diameter argument specifies the diameter *between* the compartments (and at the beginning/end of the first/last compartment). The corresponding values can therefore be later retrieved from the **Morphology** via the `start_diameter` and `end_diameter` attributes. The diameter attribute of a **Morphology** does correspond to the diameter at the midpoint of the compartment. For a **Cylinder**, `start_diameter`, `diameter`, and `end_diameter` are of course all identical.

The tree structure of a morphology is created by attaching **Morphology** objects together:

```python
morpho = Soma(diameter=30*um)
morpho.axon = Cylinder(length=100*um, diameter=1*um, n=10)
morpho.dendrite = Cylinder(length=50*um, diameter=2*um, n=5)
```

These statements create a morphology consisting of a cylindrical axon and a dendrite attached to a spherical soma. Note that the names `axon` and `dendrite` are arbitrary and chosen by the user. For example, the same morphology can be created as follows:

```python
morpho = Soma(diameter=30*um)
morpho.output_process = Cylinder(length=100*um, diameter=1*um, n=10)
morpho.input_process = Cylinder(length=50*um, diameter=2*um, n=5)
```

The syntax is recursive, for example two sections can be added at the end of the dendrite as follows:

```python
morpho.dendrite.branch1 = Cylinder(length=50*um, diameter=1*um, n=3)
morpho.dendrite.branch2 = Cylinder(length=50*um, diameter=1*um, n=3)
```

Equivalently, one can use an indexing syntax:

```python
morpho['dendrite']['branch1'] = Cylinder(length=50*um, diameter=1*um, n=3)
morpho['dendrite']['branch2'] = Cylinder(length=50*um, diameter=1*um, n=3)
```
The names given to sections are completely up to the user. However, names that consist of a single digit (1 to 9) or the letters L (for left) and R (for right) allow for a special short syntax: they can be joined together directly, without the needs for dots (or dictionary syntax) and therefore allow to quickly navigate through the morphology tree (e.g. morpho.LRLLR is equivalent to morpho.L.R.L.L.R). This short syntax can also be used to create trees:

```python
>>> morpho = Soma(diameter=30*um)
>>> morpho.L = Cylinder(length=10*um, diameter=1*um, n=3)
>>> morpho.L1 = Cylinder(length=5*um, diameter=1*um, n=3)
>>> morpho.L2 = Cylinder(length=5*um, diameter=1*um, n=3)
>>> morpho.L3 = Cylinder(length=5*um, diameter=1*um, n=3)
>>> morpho.R = Cylinder(length=10*um, diameter=1*um, n=3)
>>> morpho.RL = Cylinder(length=5*um, diameter=1*um, n=3)
>>> morpho.RR = Cylinder(length=5*um, diameter=1*um, n=3)
```

The above instructions create a dendritic tree with two main sections, three sections attached to the first section and two to the second. This can be verified with the `Morphology.topology` method:

```python
>>> morpho.topology()
( ) [root]
`---| .L
  `---| .L.1
  `---| .L.2
  `---| .L.3
`---| .R
  `---| .R.L
  `---| .R.R
```

Note that an expression such as `morpho.L` will always refer to the entire subtree. However, accessing the attributes (e.g. `diameter`) will only return the values for the given section.

**Note:** To avoid ambiguities, do not use names for sections that can be interpreted in the abbreviated way detailed above. For example, do not name a child section `L1` (which will be interpreted as the first child of the child `L`).

The number of compartments in a section can be accessed with `morpho.n` (or `morpho.L.n`, etc.), the number of total sections and compartments in a subtree can be accessed with `morpho.total_sections` and `morpho.total_compartments` respectively.

### Adding coordinates

For plotting purposes, it can be useful to add coordinates to a `Morphology` that was created using the “schematic” approach described above. This can be done by calling the `generate_coordinates` method on a morphology, which will return an identical morphology but with additional 2D or 3D coordinates. By default, this method creates a morphology according to a deterministic algorithm in 2D:

```python
new_morpho = morpho.generate_coordinates()
```
To get more “realistic” morphologies, this function can also be used to create morphologies in 3D where the orientation of each section differs from the orientation of the parent section by a random amount:

```python
new_morpho = morpho.generate_coordinates(section_randomness=25)
```

This algorithm will base the orientation of each section on the orientation of the parent section and then randomly perturb this orientation. More precisely, the algorithm first chooses a random vector orthogonal to the orientation of the parent section. Then, the section will be rotated around this orthogonal vector by a random angle, drawn from an exponential distribution with the \( \beta \) parameter (in degrees) given by `section_randomness`. This \( \beta \) parameter specifies both the mean and the standard deviation of the rotation angle. Note that no maximum rotation angle is enforced, values for `section_randomness` should therefore be reasonably small (e.g. using a `section_randomness` of 45 would already lead to a probability of \(~14\%\) that the section will be rotated by more than 90 degrees, therefore making the section go “backwards”).
In addition, also the orientation of each compartment within a section can be randomly varied:

```python
new_morpho = morpho.generate_coordinates(section_randomness=25,
                                        compartment_randomness=15)
```

The algorithm is the same as the one presented above, but applied individually to each compartment within a section (still based on the orientation on the parent section, not on the orientation of the previous compartment).

### Complex morphologies

Morphologies can also be created from information about the compartment coordinates in 3D space. Such morphologies can be loaded from a .swc file (a standard format for neuronal morphologies; for a large database of morphologies in this format see http://neuromorpho.org):

```python
morpho = Morphology.from_file('corticalcell.swc')
```

To manually create a morphology from a list of points in a similar format to SWC files, see `Morphology.from_points`.

Morphologies that are created in such a way will use standard names for the sections that allow for the short syntax shown in the previous sections: if a section has one or two child sections, then they will be called L and R, otherwise they will be numbered starting at 1.

Morphologies with coordinates can also be created section by section, following the same syntax as for “schematic” morphologies:

```python
soma = Soma(diameter=30*um, x=50*um, y=20*um)
cylinder = Cylinder(n=10, x=[0, 100]*um, diameter=1*um)
section = Section(n=5,
                  x=[0, 10, 20, 30, 40, 50]*um,
                  y=[0, 10, 20, 30, 40, 50]*um,
                  z=[0, 10, 10, 10, 10, 10]*um,
                  diameter=[6, 5, 4, 3, 2, 1]*um)
```

Note that the x, y, z attributes of `Morphology` and `SpatialNeuron` will return the coordinates at the midpoint of each compartment (as for all other attributes that vary over the length of a compartment, e.g. diameter or distance), but during construction the coordinates refer to the start and end of the section (`Cylinder`), respectively to the coordinates of the nodes between the compartments (`Section`).

A few additional remarks:

1. In the majority of simulations, coordinates are not used in the neuronal equations, therefore the coordinates are purely for visualization purposes and do not affect the simulation results in any way.
2. Coordinate specification cannot be combined with length specification – lengths are automatically calculated from the coordinates.
3. The coordinate specification can also be 1- or 2-dimensional (as in the first two examples above), the unspecified coordinate will use 0 μm.

4. All coordinates are interpreted relative to the parent compartment, i.e. the point (0 μm, 0 μm, 0 μm) refers to the end point of the previous compartment. Most of the time, the first element of the coordinate specification is therefore 0 μm, to continue a section where the previous one ended. However, it can be convenient to use a value different from 0 μm for sections connecting to the Soma to make them (visually) connect to a point on the sphere surface instead of the center of the sphere.

### 3.11.2 Creating a spatially extended neuron

A SpatialNeuron is a spatially extended neuron. It is created by specifying the morphology as a Morphology object, the equations for transmembrane currents, and optionally the specific membrane capacitance $C_m$ and intracellular resistivity $R_i$:

```python
gL = 1e-4*siemens/cm**2
EL = -70*mV
eqs = ''
Im=gL * (EL - v) : amp/meter**2
I : amp (point current)
''
neuron = SpatialNeuron(morphology=morpho, model=eqs, Cm=1*uF/cm**2, Ri=100*ohm*cm)
neuron.v = EL + 10*mV
```

Several state variables are created automatically: the SpatialNeuron inherits all the geometrical variables of the compartments (length, diameter, area, volume), as well as the distance variable that gives the distance to the soma. For morphologies that use coordinates, the x, y and z variables are provided as well. Additionally, a state variable $C_m$ is created. It is initialized with the value given at construction, but it can be modified on a compartment per compartment basis (which is useful to model myelinated axons). The membrane potential is stored in state variable $v$.

Note that for all variable values that vary across a compartment (e.g. distance, x, y, z, v), the value that is reported is the value at the midpoint of the compartment.

The key state variable, which must be specified at construction, is $Im$. It is the total transmembrane current, expressed in units of current per area. This is a mandatory line in the definition of the model. The rest of the string description may include other state variables (differential equations or subexpressions) or parameters, exactly as in NeuronGroup. At every timestep, Brian integrates the state variables, calculates the transmembrane current at every point on the neuronal morphology, and updates $v$ using the transmembrane current and the diffusion current, which is calculated based on the morphology and the intracellular resistivity. Note that the transmembrane current is a surfacic current, not the total current in the compartment. This choice means that the model equations are independent of the number of compartments chosen for the simulation. The space and time constants can obtained for any point of the neuron with the space_constant respectively time_constant attributes:

```python
l = neuron.space_constant[0]
tau = neuron.time_constant[0]
```

The calculation is based on the local total conductance (not just the leak conductance), therefore, it can potentially vary during a simulation (e.g. decrease during an action potential). The reported value is only correct for compartments with a cylindrical geometry, though, it does not give reasonable values for compartments with strongly varying diameter.

To inject a current $I$ at a particular point (e.g. through an electrode or a synapse), this current must be divided by the area of the compartment when inserted in the transmembrane current equation. This is done automatically when the flag point current is specified, as in the example above. This flag can apply only to subexpressions or parameters with amp units. Internally, the expression of the transmembrane current $Im$ is simply augmented with $+I/area$. A current can then be injected in the first compartment of the neuron (generally the soma) as follows:
neuron.I[0] = 1*nA

State variables of the *SpatialNeuron* include all the compartments of that neuron (including subtrees). Therefore, the statement neuron.v = EL + 10*mV sets the membrane potential of the entire neuron at -60 mV.

Subtrees can be accessed by attribute (in the same way as in *Morphology* objects):

neuron.axon.gNa = 10*gL

Note that the state variables correspond to the entire subtree, not just the main section. That is, if the axon had branches, then the above statement would change gNa on the main section and all the sections in the subtree. To access the main section only, use the attribute main:

neuron.axon.main.gNa = 10*gL

A typical use case is when one wants to change parameter values at the soma only. For example, inserting an electrode current at the soma is done as follows:

neuron.main.I = 1*nA

A part of a section can be accessed as follows:

initial_segment = neuron.axon[10*um:50*um]

Finally, similar to the way that you can refer to a subset of neurons of a *NeuronGroup*, you can also index the *SpatialNeuron* object itself, e.g. to get a group representing only the first compartment of a cell (typically the soma), you can use:

soma = neuron[0]

In the same way as for sections, you can also use slices, either with the indices of compartments, or with the distance from the root:

first_compartments = neurons[:3]
first_compartments = neurons[0*um:30*um]

However, note that this is restricted to contiguous indices which most of the time means that all compartments indexed in this way have to be part of the same section. Such indices can be acquired directly from the morphology:

axon = neurons[morpho.axon.indices[:]]

or, more concisely:

axon = neurons[morpho.axon]

**Synaptic inputs**

There are two methods to have synapses on *SpatialNeuron*. The first one to insert synaptic equations directly in the neuron equations:

```python
eqs=''
Im = gL * (EL - v) : amp/meter**2
Is = gs * (Es - v) : amp (point current)
dgs/dt = -gs/taus : siemens
''
neuron = SpatialNeuron(morphology=morpho, model=eqs, Cm=1*uF/cm**2, Ri=100*ohm*cm)
```
Note that, as for electrode stimulation, the synaptic current must be defined as a point current. Then we use a `Synapses` object to connect a spike source to the neuron:

```python
S = Synapses(stimulation, neuron, on_pre='gs += w')
S.connect(i=0, j=50)
S.connect(i=1, j=100)
```

This creates two synapses, on compartments 50 and 100. One can specify the compartment number with its spatial position by indexing the morphology:

```python
S.connect(i=0, j=morpho[25*um])
S.connect(i=1, j=morpho.axon[30*um])
```

In this method for creating synapses, there is a single value for the synaptic conductance in any compartment. This means that it will fail if there are several synapses onto the same compartment and synaptic equations are nonlinear.

The second method, which works in such cases, is to have synaptic equations in the `Synapses` object:

```python
eqs='''
Im = gL * (EL - v) : amp/meter**2
Is = gs * (Es - v) : amp (point current)
gs : siemens 
'''
neuron = SpatialNeuron(morphology=morpho, model=eqs, Cm=1 * uF / cm ** 2, Ri=100 * ohm / cm)
S = Synapses(stimulation, neuron, model='''dg/dt = -g/taus : siemens
          gs_post = g : siemens (summed)''',
          on_pre='g += w')
```

Here each synapse (instead of each compartment) has an associated value $g$, and all values of $g$ for each compartment (i.e., all synapses targeting that compartment) are collected into the compartmental variable $gs$.

### Detecting spikes

To detect and record spikes, we must specify a threshold condition, essentially in the same way as for a `NeuronGroup`:

```python
neuron = SpatialNeuron(morphology=morpho, model=eqs, threshold='v > 0*mV', refractory='v > -10*mV')
```

Here spikes are detected when the membrane potential $v$ reaches 0 mV. Because there is generally no explicit reset in this type of model (although it is possible to specify one), $v$ remains above 0 mV for some time. To avoid detecting spikes during this entire time, we specify a refractory period. In this case no spike is detected as long as $v$ is greater than -10 mV. Another possibility could be:

```python
neuron = SpatialNeuron(morphology=morpho, model=eqs, threshold='m > 0.5', refractory='m > 0.4')
```

where $m$ is the state variable for sodium channel activation (assuming this has been defined in the model). Here a spike is detected when half of the sodium channels are open.

With the syntax above, spikes are detected in all compartments of the neuron. To detect them in a single compartment, use the `threshold_location` keyword:

```python
neuron = SpatialNeuron(morphology=morpho, model=eqs, threshold='m > 0.5', threshold_location=30, refractory='m > 0.4')
```
In this case, spikes are only detecting in compartment number 30. Reset then applies locally to that compartment (if a reset statement is defined). Again the location of the threshold can be specified with spatial position:

```python
neuron = SpatialNeuron(morphology=morpho, model=eqs, threshold='m > 0.5',
                      threshold_location=morpho.axon[30*um],
                      refractory='m > 0.4')
```

**Subgroups**

In the same way that you can refer to a subset of neurons in a `NeuronGroup`, you can also refer to a subset of compartments in a `SpatialNeuron`.

### 3.12 Computational methods and efficiency

- **Runtime code generation**
- **Standalone code generation**
- **Compiler settings**

Brian has several different methods for running the computations in a simulation. The default mode is *Runtime code generation*, which runs the simulation loop in Python but compiles and executes the modules doing the actual simulation work (numerical integration, synaptic propagation, etc.) in a defined target language. Brian will select the best available target language automatically. On Windows, to ensure that you get the advantages of compiled code, read the instructions on installing a suitable compiler in *Windows*. Runtime mode has the advantage that you can combine the computations performed by Brian with arbitrary Python code specified as *NetworkOperation*.

The fact that the simulation is run in Python means that there is a (potentially big) overhead for each simulated time step. An alternative is to run Brian in with *Standalone code generation* — this is in general faster (for certain types of simulations much faster) but cannot be used for all kinds of simulations. To enable this mode, add the following line after your Brian import, but before your simulation code:

```python
set_device('cpp_standalone')
```

For detailed control over the compilation process (both for runtime and standalone code generation), you can change the *Cleaning up after a run* that are used.

The following topics are not essential for beginners.

#### 3.12.1 Runtime code generation

Code generation means that Brian takes the Python code and strings in your model and generates code in one of several possible different languages and actually executes that. The target language for this code generation process is set in the `codegen.target` preference. By default, this preference is set to `auto`, meaning that it will chose a compiled language target if possible and fall back to Python otherwise (it will also raise a warning in this case, set `codegen.target`
to 'numpy' explicitly to avoid this warning). There are two compiled language targets for Python 2.x, 'weave' (needing a working installation of a C++ compiler) and 'cython' (needing the Cython package in addition); for Python 3.x, only 'cython' is available. If you want to chose a code generation target explicitly (e.g. because you want to get rid of the warning that only the Python fallback is available), set the preference to 'numpy', 'weave' or 'cython' at the beginning of your script:

```python
from brian2 import *
prefs.codegen.target = 'numpy'  # use the Python fallback
```

See Preferences for different ways of setting preferences.

Warning:
Do not use the weave code generation targets when running multiple simulations in parallel. See Known issues for more details.

Caching
When you run code with weave or cython for the first time, it will take some time to compile the code. For short simulations, this can make these targets to appear slow compared to the numpy target where such compilation is not necessary. However, the compiled code is stored on disk and will be re-used for later runs, making these simulations start faster. If you run many simulations with different code (e.g. Brian’s test suite), this code can take quite a bit of space on the disk. During the import of the brian2 package, we check whether the size of the disk cache exceeds the value set by the codegen.max_cache_dir_size preference (by default, 1GB) and display a message if this is the case. You can clear the disk cache manually, or use the clear_cache function, e.g. clear_cache('cython').

3.12.2 Standalone code generation
Brian supports generating standalone code for multiple devices. In this mode, running a Brian script generates source code in a project tree for the target device/language. This code can then be compiled and run on the device, and modified if needed. At the moment, the only “device” supported is standalone C++ code. In some cases, the speed gains can be impressive, in particular for smaller networks with complicated spike propagation rules (such as STDP).

To use the C++ standalone mode, you only have to make very small changes to your script. The exact change depends on whether your script has only a single run() (or Network.run) call, or several of them:

**Single run call**
At the beginning of the script, i.e. after the import statements, add:

```python
set_device('cpp_standalone')
```

The CPPStandaloneDevice.build function will be automatically called with default arguments right after the run() call. If you need non-standard arguments then you can specify them as part of the set_device() call:

```python
set_device('cpp_standalone', directory='my_directory', debug=True)
```

**Multiple run calls**
At the beginning of the script, i.e. after the import statements, add:
set_device('cpp_standalone', build_on_run=False)

After the last `run()` call, call `device.build()` explicitly:

```python
device.build(directory='output', compile=True, run=True, debug=False)
```

The `build` function has several arguments to specify the output directory, whether or not to compile and run the project after creating it and whether or not to compile it with debugging support or not.

**Multiple builds**

To run multiple full simulations (i.e. multiple `device.build` calls, not just multiple `run()` calls as discussed above), you have to reinitialize the device again:

```python
device.reinit()
device.activate()
```

Note that the device “forgets” about all previously set build options provided to `set_device()` (most importantly the `build_on_run` option, but also e.g. the directory), you’ll have to specify them as part of the `Device.activate` call. Also, `Device.activate` will reset the `defaultclock`, you’ll therefore have to set its `dt` after the `activate` call if you want to use a non-default value.

**Limitations**

Not all features of Brian will work with C++ standalone, in particular Python based network operations and some array based syntax such as `S.w[0, :] = ...` will not work. If possible, rewrite these using string based syntax and they should work. Also note that since the Python code actually runs as normal, code that does something like this may not behave as you would like:

```python
results = []
for val in vals:
    # set up a network
    run()
    results.append(result)
```

The current C++ standalone code generation only works for a fixed number of `run` statements, not with loops. If you need to do loops or other features not supported automatically, you can do so by inspecting the generated C++ source code and modifying it, or by inserting code directly into the main loop as follows:

```python
device.insert_code('main', '''
cout << "Testing direct insertion of code." << endl;
''')
```

**Variables**

After a simulation has been run (after the `run()` call if `set_device()` has been called with `build_on_run` set to `True` or after the `Device.build` call with `run` set to `True`), state variables and monitored variables can be accessed using standard syntax, with a few exceptions (e.g. string expressions for indexing).
Multi-threading with OpenMP

**Warning:** OpenMP code has not yet been well tested and so may be inaccurate.

When using the C++ standalone mode, you have the opportunity to turn on multi-threading, if your C++ compiler is compatible with OpenMP. By default, this option is turned off and only one thread is used. However, by changing the preferences of the codegen.cpp_standalone object, you can turn it on. To do so, just add the following line in your python script:

```
prefs.devices.cpp_standalone.openmp_threads = XX
```

XX should be a positive value representing the number of threads that will be used during the simulation. Note that the speedup will strongly depend on the network, so there is no guarantee that the speedup will be linear as a function of the number of threads. However, this is working fine for networks with not too small timestep (dt > 0.1ms), and results do not depend on the number of threads used in the simulation.

Customizing the build process

In standalone mode, a standard “make file” is used to orchestrate the compilation and linking. To provide additional arguments to the `make` command (respectively `nmake` on Windows), you can use the `devices.cpp_standalone.extra_make_args_unix` or `devices.cpp_standalone.extra_make_args_windows` preference. On Linux, this preference is by default set to `['-j']` to enable parallel compilation. Note that you can also use these arguments to overwrite variables in the make file, e.g. to use clang instead of the default gcc compiler:

```
prefs.devices.cpp_standalone.extra_make_args_unix += ['CC=clang++']
```

Cleaning up after a run

Standalone simulations store all results of a simulation (final state variable values and values stored in monitors) to disk. These results can take up quite significant amount of space, and you might therefore want to delete these results when you do not need them anymore. You can do this by using the device’s `delete` method:

```
device.delete()
```

Be aware that deleting the data will make all access to state variables fail, including the access to values in monitors. You should therefore only delete the data after doing all analysis/plotting that you are interested in.

By default, this function will delete both the generated code and the data, i.e. the full project directory. If you want to keep the code (which typically takes up little space compared to the results), exclude it from the deletion:

```
device.delete(code=False)
```

If you added any additional files to the project directory manually, these will not be deleted by default. To delete the full directory regardless of its content, use the `force` option:

```
device.delete(force=True)
```

**Note:** When you initialize state variables with concrete values (and not with a string expression), they will be stored to disk from your Python script and loaded from disk at the beginning of the standalone run. Since these values
are necessary for the compiled binary file to run, they are considered “code” from the point of view of the `delete` function.

### 3.12.3 Compiler settings

If using C++ code generation (either via weave, cython or standalone), the compiler settings can make a big difference for the speed of the simulation. By default, Brian uses a set of compiler settings that switches on various optimizations and compiles for running on the same architecture where the code is compiled. This allows the compiler to make use of as many advanced instructions as possible, but reduces portability of the generated executable (which is not usually an issue).

If there are any issues with these compiler settings, for example because you are using an older version of the C++ compiler or because you want to run the generated code on a different architecture, you can change the settings by manually specifying the `codegen.cpp.extra_compile_args` preference (or by using `codegen.cpp.extra_compile_args_gcc` or `codegen.cpp.extra_compile_args_msvc` if you want to specify the settings for either compiler only).

### 3.13 Converting from integrated form to ODEs

Brian requires models to be expressed as systems of first order ordinary differential equations, and the effect of spikes to be expressed as (possibly delayed) one-off changes. However, many neuron models are given in integrated form. For example, one form of the Spike Response Model (SRM; Gerstner and Kistler 2002) is defined as

\[ V(t) = \sum_i w_i \sum_{t_i} \text{PSP}(t - t_i) + V_{\text{rest}} \]

where \( V(t) \) is the membrane potential, \( V_{\text{rest}} \) is the rest potential, \( w_i \) is the synaptic weight of synapse \( i \), and \( t_i \) are the timings of the spikes coming from synapse \( i \), and PSP is a postsynaptic potential function.

An example PSP is the \( \alpha \)-function \( \text{PSP}(t) = (t/\tau) e^{-t/\tau} \). For this function, we could rewrite the equation above in the following ODE form:

\[
\tau \frac{dV}{dt} = V_{\text{rest}} - V + g \\
\tau \frac{dg}{dt} = -g \\
g \leftarrow g + w_i \quad \text{upon spike from synapse} \ i
\]

This could then be written in Brian as:

```python
eqs = '''
dV/dt = (V_rest-V+g)/tau : 1
dg/dt = -g/tau : 1
'''
G = NeuronGroup(N, eqs, ...)
... S = Synapses(G, G, 'w : 1', on_pre='g += w')
```

To see that these two formulations are the same, you first solve the problem for the case of a single synapse and a single spike at time 0. The initial conditions at \( t = 0 \) will be \( V(0) = V_{\text{rest}}, g(0) = w \).

To solve these equations, let’s substitute \( s = t/\tau \) and take derivatives with respect to \( s \) instead of \( t \), set \( u = V - V_{\text{rest}} \), and assume \( w = 1 \). This gives us the equations \( u' = g - u, g' = -g \) with initial conditions \( u(0) = 0, g(0) = 1 \). At this point, you can either consult a textbook on solving linear systems of differential equations, or just plug this into Wolfram Alpha to get the solution \( g(s) = e^{-s}, u(s) = se^{-s} \) which is equal to the PSP given above.
Now we use the linearity of these differential equations to see that it also works when \( w \neq 0 \) and for summing over multiple spikes at different times.

In general, to convert from integrated form to ODE form, see Köhn and Wörgötter (1998), Sánchez-Montañás (2001), and Jahnke et al. (1999). However, for some simple and widely used types of synapses, use the list below. In this list, we assume synapses are postsynaptic potentials, but you can replace \( V(t) \) with a current or conductance for postsynaptic currents or conductances. In each case, we give the Brian code with unitless variables, where eqs is the differential equations for the target NeuronGroup, and on_pre is the argument to Synapses.

### Exponential synapse \( V(t) = e^{-t/\tau} \):

```plaintext
eqs = '''
dV/dt = -V/tau : 1
'''
on_pre = 'V += w'
```

### Alpha synapse \( V(t) = (t/\tau)e^{-t/\tau} \):

```plaintext
eqs = '''
dV/dt = (x-V)/tau : 1
dx/dt = -x/tau : 1
'''
on_pre = 'x += w'
```

\( V(t) \) reaches a maximum value of \( w/e \) at time \( t = \tau \).

### Biexponential synapse \( V(t) = \frac{\tau_2}{\tau_2 - \tau_1} \left( e^{-t/\tau_1} - e^{-t/\tau_2} \right) \):

```plaintext
eqs = '''
dV/dt = ((tau_2 / tau_1)**(tau_1 / (tau_2 - tau_1))**x-V)/tau_1 : 1
dx/dt = -x/tau_2 : 1
'''
on_pre = 'x += w'
```

\( V(t) \) reaches a maximum value of \( w \) at time \( t = \frac{\tau_2}{\tau_2 - \tau_1} \log \left( \frac{\tau_2}{\tau_1} \right) \).

### STDP

The weight update equation of the standard STDP is also often stated in an integrated form and can be converted to an ODE form. This is covered in Tutorial 2.
This section has additional information on details not covered in the User’s guide.

4.1 Functions

- Default functions
- User-provided functions
  - Python code generation
  - Other code generation targets
  - Dependencies between functions
  - Additional compiler arguments
  - Arrays vs. scalar values in user-provided functions
  - Functions with context-dependent return values
  - Additional namespace
  - Data types
  - External source files

All equations, expressions and statements in Brian can make use of mathematical functions. However, functions have to be prepared for use with Brian for two reasons: 1) Brian is strict about checking the consistency of units, therefore every function has to specify how it deals with units; 2) functions need to be implemented differently for different code generation targets.

Brian provides a number of default functions that are already prepared for use with numpy and C++ and also provides a mechanism for preparing new functions for use (see below).
4.1.1 Default functions

The following functions (stored in the DEFAULT_FUNCTIONS dictionary) are ready for use:

- Random numbers: `rand` (random numbers drawn from a uniform distribution between 0 and 1), `randn` (random numbers drawn from the standard normal distribution, i.e. with mean 0 and standard deviation 1), and `poisson` (discrete random numbers from a Poisson distribution with rate parameter $\lambda$)
- Elementary functions: `sqrt`, `exp`, `log`, `log10`, `abs`, `sign`
- Trigonometric functions: `sin`, `cos`, `tan`, `sinh`, `cosh`, `tanh`, `arcsin`, `arccos`, `arctan`
- Functions for improved numerical accuracy: `expm1` (calculates $\exp(x) - 1$, more accurate for $x$ close to 0), `log1p` (calculates $\log(1 + x)$, more accurate for $x$ close to 0), and `exprel` (calculates $(\exp(x) - 1)/x$, more accurate for $x$ close to 0, and returning 1.0 instead of NaN for $x == 0$)
- General utility functions: `clip`, `floor`, `ceil`

Brian also provides a special purpose function `int`, which can be used to convert an expression or variable into an integer value. This is especially useful for boolean values (which will be converted into 0 or 1), for example to have a conditional evaluation as part of an equation or statement which sometimes allows to circumvent the lack of an if statement. For example, the following reset statement resets the variable $v$ to either $v_{r1}$ or $v_{r2}$, depending on the value of $w$: `'v = v_{r1} * \text{int}(w <= 0.5) + v_{r2} * \text{int}(w > 0.5)'`

Finally, the function `timestep` is a function that takes a time and the length of a time step as an input and returns an integer corresponding to the respective time step. The advantage of using this function over a simple division is that it slightly shifts the time before dividing to avoid floating point issues. This function is used as part of the Refractoriness mechanism.

4.1.2 User-provided functions

Python code generation

If a function is only used in contexts that use Python code generation, preparing a function for use with Brian only means specifying its units. The simplest way to do this is to use the `check_units()` decorator:

```python
@check_units(x1=meter, y1=meter, x2=meter, y2=meter, result=meter)
def distance(x1, y1, x2, y2):
    return sqrt((x1 - x2)**2 + (y1 - y2)**2)
```

Another option is to wrap the function in a `Function` object:

```python
def distance(x1, y1, x2, y2):
    return sqrt((x1 - x2)**2 + (y1 - y2)**2)
# wrap the distance function
distance = Function(distance, arg_units=[meter, meter, meter, meter],
                    return_unit=meter)
```

The use of Brian’s unit system has the benefit of checking the consistency of units for every operation but at the expense of performance. Consider the following function, for example:

```python
@check_units(I=amp, result=Hz)
def piecewise_linear(I):
    return clip((I-1*nA) * 50*Hz/nA, 0*Hz, 100*Hz)
```

When Brian runs a simulation, the state variables are stored and passed around without units for performance reasons. If the above function is used, however, Brian adds units to its input argument so that the operations inside the function do not fail with dimension mismatches. Accordingly, units are removed from the return value so that the function
output can be used with the rest of the code. For better performance, Brian can alter the namespace of the function when it is executed as part of the simulation and remove all the units, then pass values without units to the function. In the above example, this means making the symbol nA refer to 1e-9 and Hz to 1. To use this mechanism, add the decorator `implementation()` with the `discard_units` keyword:

```python
@implementation('numpy', discard_units=True)
@check_units(I=amp, result=Hz)
def piecewise_linear(I):
    return clip((I-1*nA) * 50*Hz/nA, 0*Hz, 100*Hz)
```

Note that the use of the function outside of simulation runs is not affected, i.e. using `piecewise_linear` still requires a current in Ampere and returns a rate in Hertz. The `discard_units` mechanism does not work in all cases, e.g. it does not work if the function refers to units as `brian2.nA` instead of `nA`, if it uses imports inside the function (e.g. `from brian2 import nA`), etc. The `discard_units` can also be switched on for all functions without having to use the `implementation()` decorator by setting the `codegen.runtime.numpy.discard_units` preference.

### Other code generation targets

To make a function available for other code generation targets (e.g. C++), implementations for these targets have to be added. This can be achieved using the `implementation()` decorator. The form of the code (e.g. a simple string or a dictionary of strings) necessary is target-dependent, for C++ both options are allowed, a simple string will be interpreted as filling the `support_code` block. Note that both `cpp` and `weave` can be used to provide C++ implementations, the first should be used for generic C++ implementations, and the latter if weave-specific code is necessary. An implementation for the C++ target could look like this:

```python
@implementation('cpp', '')
    double piecewise_linear(double I) {
        if (I < 1e-9)
            return 0;
        if (I > 3e-9)
            return 100;
        return (I/1e-9 - 1) * 50;
    }

@check_units(I=amp, result=Hz)
def piecewise_linear(I):
    return clip((I-1*nA) * 50*Hz/nA, 0*Hz, 100*Hz)
```

Alternatively, `FunctionImplementation` objects can be added to the `Function` object. The same sort of approach as for C++ works for Cython using the `cython` target. The example above would look like this:

```python
@implementation('cython', '')
    cdef double piecewise_linear(double I):
        if I<1e-9:
            return 0.0
        elif I>3e-9:
            return 100.0
        return (I/1e-9-1)*50

@check_units(I=amp, result=Hz)
def piecewise_linear(I):
    return clip((I-1*nA) * 50*Hz/nA, 0*Hz, 100*Hz)
```
Dependencies between functions

The code generation mechanism for user-defined functions only adds the source code for a function when it is necessary. If a user-defined function refers to another function in its source code, it therefore has to explicitly state this dependency so that the code of the dependency is added as well:

```python
@implementation ('cpp', '''
    double rectified_linear(double x)
    {
        return clip(x, 0, INFINITY);
    }''',
    dependencies={'clip': DEFAULT_FUNCTIONS['clip']}
)
@check_units(x=1, result=1)
def rectified_linear(x):
    return np.clip(x, 0, np.inf)
```

Note: The dependency mechanism is unnecessary for the numpy code generation target, since functions are defined as actual Python functions and not as code given in a string.

Additional compiler arguments

If the code for a function needs additional compiler options to work, e.g. to link to an external library, these options can be provided as keyword arguments to the `@implementation` decorator. E.g. to link C++ code to the `foo` library which is stored in the directory `/usr/local/foo`, use:

```python
@implementation ('cpp', '...',
    libraries=['foo'], library_dirs=['/usr/local/foo'])
```

These arguments can also be used to refer to external source files, see below. Equivalent arguments can also be set as global `Preferences` in which case they apply to all code and not only to code referring to the respective function. Note that in C++ standalone mode, all files are compiled together, and therefore the additional compiler arguments provided to functions are always combined with the preferences into a common set of settings that is applied to all code.

The list of currently supported additional arguments (for further explications, see the respective `Preferences` and the Python documentation of the `distutils.core.Extension` class):

<table>
<thead>
<tr>
<th>keyword</th>
<th>weave</th>
<th>C++ standalone</th>
<th>Cython</th>
</tr>
</thead>
<tbody>
<tr>
<td>headers</td>
<td>☑</td>
<td>☑</td>
<td></td>
</tr>
<tr>
<td>sources</td>
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<td>runtime_library_dirs</td>
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</tr>
</tbody>
</table>

Arrays vs. scalar values in user-provided functions

Equations, expressions and abstract code statements are always implicitly referring to all the neurons in a `NeuronGroup`, all the synapses in a `Synapses` object, etc. Therefore, function calls also apply to more than a single value. The way in which this is handled differs between code generation targets that support vectorized expressions (e.g. the numpy target) and targets that don’t (e.g. the weave target or the cpp_standalone mode).
If the code generation target supports vectorized expressions, it will receive an array of values. For example, in the piecewise_linear example above, the argument I will be an array of values and the function returns an array of values. For code generation without support for vectorized expressions, all code will be executed in a loop (over neurons, over synapses, ...), the function will therefore be called several times with a single value each time.

In both cases, the function will only receive the “relevant” values, meaning that if for example a function is evaluated as part of a reset statement, it will only receive values for the neurons that just spiked.

Functions with context-dependent return values

When using the numpy target, functions have to return an array of values (e.g. one value for each neuron). In some cases, the number of values to return cannot be deduced from the function’s arguments. Most importantly, this is the case for random numbers: a call to rand() has to return one value for each neuron if it is part of a neuron’s equations, but only one value for each neuron that spiked during the time step if it is part of the reset statement. Such function are said to “auto vectorise”, which means that their implementation receives an additional array argument _vectorisation_idx; the length of this array determines the number of values the function should return. This argument is also provided to functions for other code generation targets, but in these cases it is a single value (e.g. the index of the neuron), and is currently ignored. To enable this property on a user-defined function, you’ll currently have to manually create a Function object:

```python
def exponential_rand(l, _vectorisation_idx):
    '''Generate a number from an exponential distribution using inverse transform sampling'''
    uniform = np.random.rand(len(_vectorisation_idx))
    return -(1/l)*np.log(1 - uniform)

exponential_rand = Function(exponential_rand, arg_units=[1], return_unit=1,
stateless=False, auto_vectorise=True)
```

Implementations for other code generation targets can then be added using the add_implementation mechanism:

```python
cpp_code = '''
double exponential_rand(double l, int _vectorisation_idx)
{
    double uniform = rand(_vectorisation_idx);
    return -(1/l)*log(1 - uniform);
}
'''
exponential_rand.implementations.add_implementation('cpp', cpp_code,
dependencies={'rand': DEFAULT_˓
→FUNCTIONS['rand'],
'log': DEFAULT_˓
→FUNCTIONS['log']})
```

Note that by referring to the rand function, the new random number generator will automatically generate reproducible random numbers if the seed() function is use to set its seed. Restoring the random number state with restore() will have the expected effect as well.

Additional namespace

Some functions need additional data to compute a result, e.g. a TimedArray needs access to the underlying array. For the numpy target, a function can simply use a reference to an object defined outside the function, there is no need to explicitly pass values in a namespace. For the other code language targets, values can be passed in the namespace argument of the implementation() decorator or the add_implementation method. The namespace values are then accessible in the function code under the given name, prefixed with _namespace. Note that this mechanism
should only be used for numpy arrays or general objects (e.g. function references to call Python functions from weave or Cython code). Scalar values should be directly included in the function code, by using a “dynamic implementation” (see add_dynamic_implementation).

See TimedArray and BinomialFunction for examples that use this mechanism.

Data types

By default, functions are assumed to take any type of argument, and return a floating point value. If you want to put a restriction on the type of an argument, or specify that the return type should be something other than float, either declare it as a Function (and see its documentation on specifying types) or use the declare_types() decorator, e.g.:

```python
@check_units(a=1, b=1, result=1)
@declare_types(a='integer', result='highest')
def f(a, b):
    return a*b
```

This is potentially important if you have functions that return integer or boolean values, because Brian’s code generation optimisation step will make some potentially incorrect simplifications if it assumes that the return type is floating point.

External source files

Code for functions can also be provided via external files in the target language. This can be especially useful for linking to existing code without having to include it a second time in the Python script. For C++-based code generation targets (i.e. weave and the C++ standalone mode), the external code should be in a file that is provided as an argument to the sources keyword, together with a header file whose name is provided to headers (see the note for the codegen.cpp.headers preference about the necessary format). Since the main simulation code is compiled and executed in a different directory, you should also point the compiler towards the directory of the header file via the include_dirs keyword. For the same reason, use an absolute path for the source file. For example, the piecewise_linear function from above can be implemented with external files as follows:

```python
//file: piecewise_linear.h
double piecewise_linear(double);

//file: piecewise_linear.cpp
double piecewise_linear(double I) {
    if (I < 1e-9)
        return 0;
    if (I > 3e-9)
        return 100;
    return (I/1e-9 - 1) * 50;
}
```

# Python script

```python
# Get the absolute directory of this Python script, the C++ files are expected to be stored alongside of it
import os
current_dir = os.path.abspath(os.path.dirname(__file__))

@implementation('cpp', '// all code in piecewise_linear.cpp',
              sources=[os.path.join(current_dir,
```
For Cython, the process is very similar (see the Cython documentation for general information). The name of the header file does not need to be specified, it is expected to have the same name as the source file (except for the .pxd extension). The source and header files will be automatically copied to the cache directory where Cython files are compiled, they therefore have to be imported as top-level modules, regardless of whether the executed Python code is itself in a package or module.

A Cython equivalent of above’s C++ example can be written as:

```python
# file: piecewise_linear.pxd
cdef double piecewise_linear(double)

# file: piecewise_linear.pyx
cdef double piecewise_linear(double I):
    if I<1e-9:
        return 0.0
    elif I>3e-9:
        return 100.0
    return (I/1e-9-1)*50
```

# Python script

```python
# Get the absolute directory of this Python script, the Cython files # are expected to be stored alongside of it
import os
current_dir = os.path.abspath(os.path.dirname(__file__))

@implementation('cython',
               'from piecewise_linear cimport piecewise_linear',
               sources=[os.path.join(current_dir, 'piecewise_linear.pyx')])
@check_units(I=amp, result=Hz)
def piecewise_linear(I):
    return clip((I-1*nA) * 50*Hz/nA, 0*Hz, 100*Hz)
```

### 4.2 Preferences

Brian has a system of global preferences that affect how certain objects behave. These can be set either in scripts by using the `prefs` object or in a file. Each preference looks like `codegen.c.compiler`, i.e. dotted names.

#### 4.2.1 Accessing and setting preferences

Preferences can be accessed and set either keyword-based or attribute-based. The following are equivalent:

```python
prefs['codegen.c.compiler'] = 'gcc'
prefs.codegen.c.compiler = 'gcc'
```
Using the attribute-based form can be particularly useful for interactive work, e.g. in ipython, as it offers autocompletion and documentation. In ipython, `prefs.codegen.c?` would display a docstring with all the preferences available in the `codegen.c` category.

### 4.2.2 Preference files

Preferences are stored in a hierarchy of files, with the following order (each step overrides the values in the previous step but no error is raised if one is missing):

- The global defaults are stored in the installation directory.
- The user default are stored in `~/.brian/user_preferences` (which works on Windows as well as Linux). The `~` symbol refers to the user directory.
- The file `brian_preferences` in the current directory.

The preference files are of the following form:

```
[a.b.c = 1]
# Comment line
[a]
b.d = 2
[a.b]
b.e = 3
```

This would set preferences `a.b.c=1, a.b.d=2 and a.b.e=3`.

### 4.2.3 List of preferences

Brian itself defines the following preferences (including their default values):

**GSL**

Directory containing GSL code

**GSL.directory = None** Set path to directory containing GSL header files (gsl_odeiv2.h etc.) If this directory is already in Python’s include (e.g. because of conda installation), this path can be set to None.

**codegen**

Code generation preferences

`codegen.loop_invariant_optimisations = True`

Whether to pull out scalar expressions out of the statements, so that they are only evaluated once instead of once for every neuron/synapse/... Can be switched off, e.g. because it complicates the code (and the same optimisation is already performed by the compiler) or because the code generation target does not deal well with it. Defaults to True.

`codegen.max_cache_dir_size = 1000`

The size of a directory (in MB) with cached code for weave or Cython that triggers a warning. Set to 0 to never get a warning.

`codegen.string_expression_target = 'numpy'`
Default target for the evaluation of string expressions (e.g. when indexing state variables). Should normally not be changed from the default numpy target, because the overhead of compiling code is not worth the speed gain for simple expressions.

Accepts the same arguments as `codegen.target`, except for 'auto'

```
codegen.target = 'auto'
```

Default target for code generation.

Can be a string, in which case it should be one of:

- 'auto' the default, automatically chose the best code generation target available.
- 'weave' uses `scipy.weave` to generate and compile C++ code, should work anywhere where `gcc` is installed and available at the command line.
- 'cython', uses the Cython package to generate C++ code. Needs a working installation of Cython and a C++ compiler.
- 'numpy' works on all platforms and doesn’t need a C compiler but is often less efficient.

Or it can be a `CodeObject` class.

### codegen.cpp

C++ compilation preferences

```
codegen.cpp.compiler = ''
```

Compiler to use (uses default if empty)

Should be gcc or msvc.

```
codegen.cpp.define_macros = []
```

List of macros to define; each macro is defined using a 2-tuple, where ‘value’ is either the string to define it to or None to define it without a particular value (equivalent of “#define FOO” in source or -DFOO on Unix C compiler command line).

```
codegen.cpp.extra_compile_args = None
```

Extra arguments to pass to compiler (if None, use either `extra_compile_args_gcc` or `extra_compile_args_msvc`).

```
codegen.cpp.extra_compile_args_gcc = ['-w', '-O3', '-ffast-math', '-fno-finite-math-only', '-march=native']
```

Extra compile arguments to pass to GCC compiler

```
codegen.cpp.extra_compile_args_msvc = ['/Ox', '/w', '', '/MP']
```

Extra compile arguments to pass to MSVC compiler (the default /arch: flag is determined based on the processor architecture)

```
codegen.cpp.extra_link_args = []
```

Any extra platform- and compiler-specific information to use when linking object files together.

```
codegen.cpp.headers = []
```

A list of strings specifying header files to use when compiling the code. The list might look like ["<vectors>","my_header"]''. Note that the header strings need to be in a form than can be pasted at the end of a #include statement in the C++ code.

```
codegen.cpp.include_dirs = []
```
Include directories to use. Note that $prefix/include will be appended to the end automatically, where $prefix is Python's site-specific directory prefix as returned by sys.prefix.

codegen.cpp.libraries = []

List of library names (not filenames or paths) to link against.

codegen.cpp.library_dirs = []

List of directories to search for C/C++ libraries at link time. Note that $prefix/lib will be appended to the end automatically, where $prefix is Python's site-specific directory prefix as returned by sys.prefix.

codegen.cpp.msvc_architecture = ''

MSVC architecture name (or use system architecture by default).

Could take values such as x86, amd64, etc.

codegen.cpp.msvc_vars_location = ''

Location of the MSVC command line tool (or search for best by default).

codegen.cpp.runtime_library_dirs = []

List of directories to search for C/C++ libraries at run time.

codegen.generators

Codegen generator preferences (see subcategories for individual languages)

codegen.generators.cpp

C++ codegen preferences

codegen.generators.cpp.flush_denormals = False

Adds code to flush denormals to zero.

The code is gcc and architecture specific, so may not compile on all platforms. The code, for reference is:

```c
#define CSR_FLUSH_TO_ZERO (1 << 15)
unsigned csr = __builtin_ia32_stmxcsr();
csr |= CSR_FLUSH_TO_ZERO;
__builtin_ia32_ldmxcsr(csr);
```


codegen.generators.cpp.restrict_keyword = '__restrict'

The keyword used for the given compiler to declare pointers as restricted.

This keyword is different on different compilers, the default works for gcc and MSVS.

codegen.runtime

Runtime codegen preferences (see subcategories for individual targets)

codegen.runtime.cython

Cython runtime codegen preferences

codegen.runtime.cython.cache_dir = None

Location of the cache directory for Cython files. By default, will be stored in a brian_extensions subdirectory where Cython inline stores its temporary files (the result of get_cython_cache_dir()).
codegen.runtime.cython.delete_source_files = True

Whether to delete source files after compiling. The Cython source files can take a significant amount of
disk space, and are not used anymore when the compiled library file exists. They are therefore deleted by
default, but keeping them around can be useful for debugging.

codegen.runtime.cython.multiprocess_safe = True

Whether to use a lock file to prevent simultaneous write access to cython .pyx and .so files.

codegen.runtime.npumpy

Numpy runtime codegen preferences

codegen.runtime.npumpy.discard_units = False

Whether to change the namespace of user-specified functions to remove units.

core

Core Brian preferences

core.default_float_dtype = float64

Default dtype for all arrays of scalars (state variables, weights, etc.).
core.default_integer_dtype = int32

Default dtype for all arrays of integer scalars.
core.outdated_dependency_error = True

Whether to raise an error for outdated dependencies (True) or just a warning (False).

core.network

Network preferences

core.network.default_schedule = ['start', 'groups', 'thresholds', 'synapses', 'resets', 'end']

Default schedule used for networks that don’t specify a schedule.

devices

Device preferences

devices.cpp_standalone

C++ standalone preferences

devices.cpp_standalone.extra_make_args_unix = ['-j']

Additional flags to pass to the GNU make command on Linux/OS-X. Defaults to “-j” for parallel compi-
lation.
devices.cpp_standalone.extra_make_args_windows = []

Additional flags to pass to the nmake command on Windows. By default, no additional flags are passed.
devices.cpp_standalone.openmp_spatialneuron_strategy = None

DEPRECATED. Previously used to chose the strategy to parallelize the solution of the three tridiagonal
systems for multicompartamental neurons. Now, its value is ignored.
devices.cpp_standalone.openmp_threads = 0
The number of threads to use if OpenMP is turned on. By default, this value is set to 0 and the C++ code is generated without any reference to OpenMP. If greater than 0, then the corresponding number of threads are used to launch the simulation.

```python
devices.cpp_standalone.run_environment_variables = {'LD_BIND_NOW': '1'}
```

Dictionary of environment variables and their values that will be set during the execution of the standalone code.

### legacy

Preferences to enable legacy behaviour

```python
legacy.refractory_timing = False
```

Whether to use the semantics for checking the refractoriness condition that were in place up until (including) version 2.1.2. In that implementation, refractory periods that were multiples of dt could lead to a varying number of refractory timesteps due to the nature of floating point comparisons). This preference is only provided for exact reproducibility of previously obtained results, new simulations should use the improved mechanism which uses a more robust mechanism to convert refractoriness into timesteps. Defaults to `False`.

### logging

Logging system preferences

```python
logging.console_log_level = 'INFO'
```

What log level to use for the log written to the console.

Has to be one of CRITICAL, ERROR, WARNING, INFO, DEBUG or DIAGNOSTIC.

```python
logging.delete_log_on_exit = True
```

Whether to delete the log and script file on exit.

If set to `True` (the default), log files (and the copy of the main script) will be deleted after the brian process has exited, unless an uncaught exception occurred. If set to `False`, all log files will be kept.

```python
logging.file_log = True
```

Whether to log to a file or not.

If set to `True` (the default), logging information will be written to a file. The log level can be set via the `logging.file_log_level` preference.

```python
logging.file_log_level = 'DIAGNOSTIC'
```

What log level to use for the log written to the log file.

In case file logging is activated (see `logging.file_log`), which log level should be used for logging. Has to be one of CRITICAL, ERROR, WARNING, INFO, DEBUG or DIAGNOSTIC.

```python
logging.save_script = True
```

Whether to save a copy of the script that is run.

If set to `True` (the default), a copy of the currently run script is saved to a temporary location. It is deleted after a successful run (unless `logging.delete_log_on_exit` is `False`) but is kept after an uncaught exception occurred. This can be helpful for debugging, in particular when several simulations are running in parallel.

```python
logging.std_redirection = True
```
Whether or not to redirect stdout/stderr to null at certain places.

This silences a lot of annoying compiler output, but will also hide error messages making it harder to debug problems. You can always temporarily switch it off when debugging. If `logging.std_redirection_to_file` is set to `True` as well, then the output is saved to a file and if an error occurs the name of this file will be printed.

```plaintext
logging.std_redirection_to_file = True
```

Whether to redirect stdout/stderr to a file.

If both `logging.std_redirection` and this preference are set to `True`, all standard output/error (most importantly output from the compiler) will be stored in files and if an error occurs the name of this file will be printed. If `logging.std_redirection` is `True` and this preference is `False`, then all standard output/error will be completely suppressed, i.e. neither be displayed nor stored in a file.

The value of this preference is ignore if `logging.std_redirection` is set to `False`.

### 4.3 Logging

Brian uses a logging system to display warnings and general information messages to the user, as well as writing them to a file with more detailed information, useful for debugging. Each log message has one of the following “log levels”:

- **ERROR** Only used when an exception is raised, i.e. an error occurs and the current operation is interrupted. *Example:* You use a variable name in an equation that Brian does not recognize.

- **WARNING** Brian thinks that something is most likely a bug, but it cannot be sure. *Example:* You use a `Synapses` object without any synapses in your simulation.

- **INFO** Brian wants to make the user aware of some automatic choice that it did for the user. *Example:* You did not specify an integration method for a `NeuronGroup` and therefore Brian chose an appropriate method for you.

- **DEBUG** Additional information that might be useful when a simulation is not working as expected. *Example:* The integration timestep used during the simulation.

- **DIAGNOSTIC** Additional information useful when tracking down bugs in Brian itself. *Example:* The generated code for a `CodeObject`.

By default, all messages are written to the log file and all messages of level `INFO` and above are displayed on the console. To change what messages are displayed, see below.

---

**Note:** By default, the log file is deleted after a successful simulation run, i.e. when the simulation exited without an error. To keep the log around, set the `logging.delete_log_on_exit` preference to `False`.

### 4.3.1 Showing/hiding log messages

If you want to change what messages are displayed on the console, you can call a method of the method of `BrianLogger`:

```plaintext
BrianLogger.log_level_debug()  # now also display debug messages
```

It is also possible to suppress messages for certain sub-hierarchies by using `BrianLogger.suppress_hierarchy`:

---
# Suppress code generation messages on the console
BrianLogger.suppress_hierarchy('brian2.codegen')

# Suppress preference messages even in the log file
BrianLogger.suppress_hierarchy('brian2.core.preferences', filter_log_file=True)

Similarly, messages ending in a certain name can be suppressed with `BrianLogger.suppress_name`:

```python
# Suppress resolution conflict warnings
BrianLogger.suppress_name('resolution_conflict')
```

These functions should be used with care, as they suppresses messages independent of the level, i.e. even warning and error messages.

## 4.3.2 Preferences

You can also change details of the logging system via Brian’s `Preferences` system. With this mechanism, you can switch the logging to a file off completely (by setting `logging.file_log` to `False`) or have it log less messages (by setting `logging.file_log_level` to a level higher than `DIAGNOSTIC`) – this can be important for long-running simulations where the log might otherwise take up a lot of disk space. For a list of all preferences related to logging, see the documentation of the `brian2.utils.logger` module.

**Warning:** Most of the logging preferences are only taken into account during the initialization of the logging system which takes place as soon as `brian2` is imported. Therefore, if you use e.g. `prefs.logging.file_log = False` in your script, this will not have the intended effect! Instead, set these preferences in a file (see `Preferences`).

## 4.4 Namespaces

Equations can contain references to external parameters or functions. During the initialisation of a `NeuronGroup` or a `Synapses` object, this namespace can be provided as an argument. This is a group-specific namespace that will only be used for names in the context of the respective group. Note that units and a set of standard functions are always provided and should not be given explicitly. This namespace does not necessarily need to be exhaustive at the time of the creation of the `NeuronGroup/Synapses`, entries can be added (or modified) at a later stage via the `namespace` attribute (e.g. `G.namespace['tau'] = 10*ms`).

At the point of the call to the `Network.run` namespace, any group-specific namespace will be augmented by the “run namespace”. This namespace can be either given explicitly as an argument to the `run` method or it will be taken from the locals and globals surrounding the call. A warning will be emitted if a name is defined in more than one namespace.

To summarize: an external identifier will be looked up in the context of an object such as `NeuronGroup` or `Synapses`. It will follow the following resolution hierarchy:

1. Default unit and function names.
2. Names defined in the explicit group-specific namespace.
3. Names in the run namespace which is either explicitly given or the implicit namespace surrounding the run call.

Note that if you completely specify your namespaces at the `Group` level, you should probably pass an empty dictionary as the namespace argument to the `run` call – this will completely switch off the “implicit namespace” mechanism.
The following three examples show the different ways of providing external variable values, all having the same effect in this case:

```python
# Explicit argument to the NeuronGroup
G = NeuronGroup(1, 'dv/dt = -v / tau : 1', namespace={'tau': 10*ms})
net = Network(G)
net.run(10*ms)

# Explicit argument to the run function
G = NeuronGroup(1, 'dv/dt = -v / tau : 1')
net = Network(G)
net.run(10*ms, namespace={'tau': 10*ms})

# Implicit namespace from the context
G = NeuronGroup(1, 'dv/dt = -v / tau : 1')
net = Network(G)
tau = 10*ms
net.run(10*ms)
```

External variables are free to change between runs (but not during one run), the value at the time of the `run()` call is used in the simulation.

### 4.5 Custom progress reporting

#### 4.5.1 Progress reporting

For custom progress reporting (e.g. graphical output, writing to a file, etc.), the `report` keyword accepts a callable (i.e. a function or an object with a `__call__` method) that will be called with four parameters:

- `elapsed`: the total (real) time since the start of the run
- `completed`: the fraction of the total simulation that is completed, i.e. a value between 0 and 1
- `start`: The start of the simulation (in biological time)
- `duration`: the total duration (in biological time) of the simulation

The function will be called every `report_period` during the simulation, but also at the beginning and end with `completed` equal to 0.0 and 1.0, respectively.

For the C++ standalone mode, the same standard options are available. It is also possible to implement custom progress reporting by directly passing the code (as a multi-line string) to the `report` argument. This code will be filled into a progress report function template, it should therefore only contain a function body. The simplest use of this might look like:

```python
net.run(duration, report='std::cout << (int)(completed*100.) << "$ completed" << std::endl;')
```

### Examples of custom reporting

#### Progress printed to a file

```python
from brian2.core.network import TextReport
report_file = open('report.txt', 'w')
file_reporter = TextReport(report_file)
```

(continues on next page)
**4.6 Random numbers**

Brian provides two basic functions to generate random numbers that can be used in model code and equations: `rand()`, to generate uniformly generated random numbers between 0 and 1, and `randn()`, to generate random numbers from a standard normal distribution (i.e. normally distributed numbers with a mean of 0 and a standard deviation of 1). All other stochastic elements of a simulation (probabilistic connections, Poisson-distributed input generated by `PoissonGroup` or `PoissonInput`, differential equations using the noise term $\xi$, ...) will internally make use of these two basic functions.

For **Runtime code generation**, random numbers are generated by `numpy.random.rand` and `numpy.random.randn` respectively, which uses a Mersenne-Twister pseudorandom number generator. When the `numpy` code generation target is used, these functions are called directly, but for `weave` and `cython`, Brian uses a internal buffers for uniformly and normally distributed random numbers and calls the `numpy` functions whenever all numbers from this buffer have been used. This avoids the overhead of switching between C code and Python code for each random number. For **Standalone code generation**, the random number generation is based on “randomkit”, the same Mersenne-Twister implementation that is used by `numpy`. The source code of this implementation will be included in every generated standalone project.

**4.6.1 Seeding and reproducibility**
Runtime mode

As explained above, Runtime code generation makes use of numpy’s random number generator. In principle, using numpy.random.seed therefore permits reproducing a stream of random numbers. However, for weave and cython, Brian’s buffer complicates the matter a bit: if a simulation sets numpy’s seed, uses 10000 random numbers, and then resets the seed, the following 10000 random numbers (assuming the current size of the buffer) will come out of the pre-generated buffer before numpy’s random number generation functions are called again and take into account the seed set by the user. Instead, users should use the seed() function provided by Brian 2 itself, this will take care of setting numpy’s random seed and empty Brian’s internal buffers. This function also has the advantage that it will continue to work when the simulation is switched to standalone code generation (see below). Note that random numbers are not guaranteed to be reproducible across different code generation targets or different versions of Brian, especially if several sources of randomness are used in the same CodeObject (e.g. two noise variables in the equations of a NeuronGroup). This is because Brian does not guarantee the order of certain operations (e.g. should it first generate all random numbers for the first noise variable for all neurons, followed by the random numbers for the second noise variable for all neurons or rather first the random numbers for all noise variables of the first neuron, then for the second neuron, etc.) Since all random numbers are coming from the same stream of random numbers, the order of getting the numbers out of this stream matter.

Standalone mode

For Standalone code generation, Brian’s seed() function will insert code to set the random number generator seed into the generated code. The code will be generated at the position where the seed() call was made, allowing detailed control over the seeding. For example the following code would generate identical initial conditions every time it is run, but the noise generated by the xi variable would differ:

```python
G = NeuronGroup(10, 'dv/dt = -v/(10*ms) + 0.1*xi/sqrt(ms) : 1')
seed(4321)
G.v = 'rand()' 
seed()
run(100*ms)
```

**Note:** In standalone mode, seed() will not set numpy’s random number generator. If you use random numbers in the Python script itself (e.g. to generate a list of synaptic connections that will be passed to the standalone code as a pre-calculated array), then you have to explicitly call numpy.random.seed yourself to make these random numbers reproducible.

**Note:** Seeding should lead to reproducible random numbers even when using OpenMP with multiple threads (for repeated simulations with the same number of threads), but this has not been rigorously tested. Use at your own risk.

4.7 Custom events

4.7.1 Overview

In most simulations, a NeuronGroup defines a threshold on its membrane potential that triggers a spike event. This event can be monitored by a SpikeMonitor, it is used in synaptic interactions, and in integrate-and-fire models it also leads to the execution of one or more reset statements.

Sometimes, it can be useful to define additional events, e.g. when an ion concentration in the cell crosses a certain threshold. This can be done with the custom events system in Brian, which is illustrated in this diagram.
You can see in this diagram that the source NeuronGroup has four types of events, called spike, evt_other, evt_mon and evt_run. The event spike is the default event. It is triggered when you include threshold='...' in a NeuronGroup, and has two potential effects. Firstly, when the event is triggered it causes the reset code to run, specified by reset='...'. Secondly, if there are Synapses connected, it causes the on_pre on on_post code to run (depending if the NeuronGroup is presynaptic or postsynaptic for those Synapses).

In the diagram though, we have three additional event types. We’ve included several event types here to make it clearer, but you could use the same event for different purposes. Let’s start with the first one, evt_other. To understand this, we need to look at the Synapses object in a bit more detail. A Synapses object has multiple pathways associated to it. By default, there are just two, called pre and post. The pre pathway is activated by presynaptic spikes, and the post pathway by postsynaptic spikes. Specifically, the spike event on the presynaptic NeuronGroup triggers the pre pathway, and the spike event on the postsynaptic NeuronGroup triggers the post pathway. In the example in the diagram, we have created a new pathway called other, and the evt_other event in the presynaptic NeuronGroup triggers this pathway. Note that we can arrange this however we want. We could have spike trigger the other pathway if we wanted to, or allow it to trigger both the pre and other pathways. We could also allow evt_other to trigger the pre pathway. See below for details on the syntax for this.

The third type of event in the example is named evt_mon and this is connected to an EventMonitor which works exactly the same way as SpikeMonitor (which is just an EventMonitor attached by default to the event spike).

Finally, the fourth type of event in the example is named evt_run, and this causes some code to be run in the NeuronGroup triggered by the event. To add this code, we call NeuronGroup.run_on_event. So, when you set reset='...', this is equivalent to calling NeuronGroup.run_on_event with the spike event.

### 4.7.2 Details
Defining an event

This can be done with the **events** keyword in the **NeuronGroup** initializer:

```python
group = NeuronGroup(N, '...', threshold='...', reset='...',
    events={'custom_event': 'x > x_th'})
```

In this example, we define an event with the name `custom_event` that is triggered when the `x` variable crosses the threshold `x_th`. Note that you can define any number of custom events. Each event is defined by its name as the key, and its condition as the value of the dictionary.

Recording events

Custom events can be recorded with an **EventMonitor**:

```python
event_mon = EventMonitor(group, 'custom_event')
```

Such an **EventMonitor** can be used in the same way as a **SpikeMonitor** – in fact, creating the **SpikeMonitor** is basically identical to recording the spike event with an **EventMonitor**. An **EventMonitor** is not limited to record the event time/neuron index, it can also record other variables of the model at the time of the event:

```python
event_mon = EventMonitor(group, 'custom_event', variables=['var1', 'var2'])
```

Triggering NeuronGroup code

If the event should trigger a series of statements (i.e. the equivalent of **reset** statements), this can be added by calling **run_on_event**:

```python
group.run_on_event('custom_event', 'x=0')
```

Triggering synaptic pathways

When neurons are connected by **Synapses**, the **pre** and **post** pathways are triggered by spike events on the presynaptic and postsynaptic **NeuronGroup** by default. It is possible to change which pathway is triggered by which event by providing an **on_event** keyword that either specifies which event to use for all pathways, or a specific event for each pathway (where non-specified pathways use the default spike event):

```python
synapse_1 = Synapses(group, another_group, '...', on_pre='...', on_event='custom_event')
```

The code above causes all pathways to be triggered by an event named `custom_event` instead of the default spike.

```python
synapse_2 = Synapses(group, another_group, '...', on_pre='...', on_post='...',
    on_event={'pre': 'custom_event'})
```

In the code above, only the **pre** pathway is triggered by the `custom_event` event.

We can also create new pathways and have them be triggered by custom events. For example:

```python
synapse_3 = Synapses(group, another_group, '...',
    on_pre={'pre': '....',
            'custom_pathway': '....'},
    on_event={'pre': 'spike',
              'custom_pathway': 'custom_event'})
```
In this code, the default pre pathway is still triggered by the spike event, but there is a new pathway called custom_pathway that is triggered by the custom_event event.

**Scheduling**

By default, custom events are checked after the spiking threshold (in the after_thresholds slots) and statements are executed after the reset (in the after_resets slots). The slot for the execution of custom event-triggered statements can be changed when it is added with the usual when and order keyword arguments (see Scheduling for details). To change the time when the condition is checked, use NeuronGroup.set_event_schedule.

4.8 State update

In Brian, a state updater transforms a set of equations into an abstract state update code (and therefore is automatically target-independent). In general, any function (or callable object) that takes an Equations object and returns abstract code (as a string) can be used as a state updater and passed to the NeuronGroup constructor as a method argument.

The more common use case is to specify no state updater at all or chose one by name, see Choice of state updaters below.

4.8.1 Explicit state update

Explicit state update schemes can be specified in mathematical notation, using the ExplicitStateUpdater class. A state updater scheme contains a series of statements, defining temporary variables and a final line (starting with x_new =), giving the updated value for the state variable. The description can make reference to t (the current time), dt (the size of the time step), x (value of the state variable), and f(x, t) (the definition of the state variable x, assuming dx/dt = f(x, t)). In addition, state updaters supporting stochastic equations additionally make use of dW (a normal distributed random variable with variance dt) and g(x, t), the factor multiplied with the noise variable, assuming dx/dt = f(x, t) + g(x, t) * \xi.

Using this notation, simple forward Euler integration is specified as:

```plaintext
x_new = x + dt * f(x, t)
```

A Runge-Kutta 2 (midpoint) method is specified as:

```plaintext
k = dt * f(x,t)
\text{\textbf{x}}_{\text{new}} = x + dt \times f(x + k/2, t + dt/2)
```

When creating a new state updater using ExplicitStateUpdater, you can specify the stochastic keyword argument, determining whether this state updater does not support any stochastic equations (None, the default), stochastic equations with additive noise only ('additive'), or arbitrary stochastic equations ('multiplicative'). The provided state updaters use the Stratonovich interpretation for stochastic equations (which is the correct interpretation if the white noise source is seen as the limit of a coloured noise source with a short time constant). As a result of this, the simple Euler-Maruyama scheme (x_{new} = x + dt*f(x, t) + dW*g(x, t)) will only be used for additive noise.

An example for a general state updater that handles arbitrary multiplicative noise (under Stratonovich interpretation) is the derivative-free Milstein method:

```plaintext
x_{support} = x + dt*f(x, t) + dt^{.5} * g(x, t)  
g_{support} = g(x_{support}, t)  
k = (1/2*dt^{.5})*(g_{support} - g(x, t)) * (dW^{*2})    
x_{new} = x + dt*f(x,t) + g(x, t) * dW + k
```
Note that a single line in these descriptions is only allowed to mention \( g(x, t) \), respectively \( f(x, t) \) only once (and you are not allowed to write, for example, \( g(f(x, t), t) \)). You can work around these restrictions by using intermediate steps, defining temporary variables, as in the above examples for \texttt{milstein} and \texttt{rk2}.

### 4.8.2 Choice of state updaters

As mentioned in the beginning, you can pass arbitrary callables to the method argument of a \texttt{NeuronGroup}, as long as this callable converts an \texttt{Equations} object into abstract code. The best way to add a new state updater, however, is to register it with brian and provide a method to determine whether it is appropriate for a given set of equations. This way, it can be automatically chosen when no method is specified and it can be referred to with a name (i.e. you can pass a string like \texttt{'euler'} to the method argument instead of importing \texttt{euler} and passing a reference to the object itself).

If you create a new state updater using the \texttt{ExplicitStateUpdater} class, you have to specify what kind of stochastic equations it supports. The keyword argument \texttt{stochastic} takes the values \texttt{None} (no stochastic equation support, the default), \texttt{'additive'} (support for stochastic equations with additive noise), \texttt{'multiplicative'} (support for arbitrary stochastic equations).

After creating the state updater, it has to be registered with \texttt{StateUpdateMethod}:

```python
new_state_updater = ExplicitStateUpdater('...', stochastic='additive')
StateUpdateMethod.register('mymethod', new_state_updater)
```

The preferred way to do write new general state updaters (i.e. state updaters that cannot be described using the explicit syntax described above) is to extend the \texttt{StateUpdateMethod} class (but this is not strictly necessary, all that is needed is an object that implements a \texttt{__call__} method that operates on an \texttt{Equations} object and a dictionary of variables). Optionally, the state updater can be registered with \texttt{StateUpdateMethod} as shown above.

### 4.8.3 Implicit state updates

**Note:** All of the following is just here for future reference, it’s not implemented yet.

Implicit schemes often use Newton-Raphson or fixed point iterations. These can also be defined by mathematical statements, but the number of iterations is dynamic and therefore not easily vectorised. However, this might not be a big issue in C, GPU or even with Numba.

**Backward Euler**

Backward Euler is defined as follows:

```plaintext
x(t+dt) = x(t) + dt*f(x(t+dt), t+dt)
```

This is not a executable statement because the RHS depends on the future. A simple way is to perform fixed point iterations:

```plaintext
x(t+dt) = x(t)
x(t+dt) = x(t) + dt*dx = f(x(t+dt), t+dt)  until increment<tolerance
```

This includes a loop with a different number of iterations depending on the neuron.
4.9 How Brian works

In this section we will briefly cover some of the internals of how Brian works. This is included here to understand the general process that Brian goes through in running a simulation, but it will not be sufficient to understand the source code of Brian itself or to extend it to do new things. For a more detailed view of this, see the documentation in the Developer's guide.

4.9.1 Clock-driven versus event-driven

Brian is a clock-driven simulator. This means that the simulation time is broken into an equally spaced time grid, 0, dt, 2*dt, 3*dt, .... At each time step t, the differential equations specifying the models are first integrated giving the values at time t+dt. Spikes are generated when a condition such as \( v > v_t \) is satisfied, and spikes can only occur on the time grid.

The advantage of clock driven simulation is that it is very flexible (arbitrary differential equations can be used) and computationally efficient. However, the time grid approximation can lead to an overestimate of the amount of synchrony that is present in a network. This is usually not a problem, and can be managed by reducing the time step dt, but it can be an issue for some models.

Note that the inaccuracy introduced by the spike time approximation is of order O(dt), so the total accuracy of the simulation is of order O(dt) per time step. This means that in many cases, there is no need to use a higher order numerical integration method than forward Euler, as it will not improve the order of the error beyond O(dt). See State update for more details of numerical integration methods.

Some simulators use an event-driven method. With this method, spikes can occur at arbitrary times instead of just on the grid. This method can be more accurate than a clock-driven simulation, but it is usually substantially more computationally expensive (especially for larger networks). In addition, they are usually more restrictive in terms of the class of differential equations that can be solved.

For a review of some of the simulation strategies that have been used, see Brette et al. 2007.

4.9.2 Code overview

The user-visible part of Brian consists of a number of objects such as NeuronGroup, Synapses, Network, etc. These are all written in pure Python and essentially work to translate the user specified model into the computational engine. The end state of this translation is a collection of short blocks of code operating on a namespace, which are called in a sequence by the Network. Examples of these short blocks of code are the “state updaters” which perform numerical integration, or the synaptic propagation step. The namespaces consist of a mapping from names to values, where the possible values can be scalar values, fixed-length or dynamically sized arrays, and functions.

4.9.3 Syntax layer

The syntax layer consists of everything that is independent of the way the final simulation is computed (i.e. the language and device it is running on). This includes things like NeuronGroup, Synapses, Network, Equations, etc.

The user-visible part of this is documented fully in the User's guide and the Advanced guide. In particular, things such as the analysis of equations and assignment of numerical integrators. The end result of this process, which is passed to the computational engine, is a specification of the simulation consisting of the following data:

- A collection of variables which are scalar values, fixed-length arrays, dynamically sized arrays, and functions. These are handled by Variable objects detailed in Variables and indices. Examples: each state variable of a NeuronGroup is assigned an ArrayVariable; the list of spike indices stored by a SpikeMonitor is assigned a DynamicArrayVariable; etc.
• A collection of code blocks specified via an “abstract code block” and a template name. The “abstract code block” is a sequence of statements such as $v = vr$ which are to be executed. In the case that say, $v$ and $vr$ are arrays, then the statement is to be executed for each element of the array. These abstract code blocks are either given directly by the user (in the case of neuron threshold and reset, and synaptic pre and post codes), or generated from differential equations combined with a numerical integrator. The template name is one of a small set (around 20 total) which give additional context. For example, the code block $a = b$ when considered as part of a “state update” means execute that for each neuron index. In the context of a reset statement, it means execute it for each neuron index of a neuron that has spiked. Internally, these templates need to be implemented for each target language/device, but there are relatively few of them.

• The order of execution of these code blocks, as defined by the Network.

4.9.4 Computational engine

The computational engine covers everything from generating to running code in a particular language or on a particular device. It starts with the abstract definition of the simulation resulting from the syntax layer described above.

The computational engine is described by a Device object. This is used for allocating memory, generating and running code. There are two types of device, “runtime” and “standalone”. In runtime mode, everything is managed by Python, even if individual code blocks are in a different language. Memory is managed using numpy arrays (which can be passed as pointers to use in other languages). In standalone mode, the output of the process (after calling Device.build) is a complete source code project that handles everything, including memory management, and is independent of Python.

For both types of device, one of the key steps that works in the same way is code generation, the creation of a compilable and runnable block of code from an abstract code block and a collection of variables. This happens in two stages: first of all, the abstract code block is converted into a code snippet, which is a syntactically correct block of code in the target language, but not one that can run on its own (it doesn’t handle accessing the variables from memory, etc.). This code snippet typically represents the inner loop code. This step is handled by a CodeGenerator object. In some cases it will involve a syntax translation (e.g. the Python syntax $x**y$ in C++ should be $\text{pow}(x, y)$).

The next step is to insert this code snippet into a template to form a compilable code block. This code block is then passed to a runtime CodeObject. In the case of standalone mode, this doesn’t do anything, but for runtime devices it handles compiling the code and then running the compiled code block in the given namespace.

4.10 Interfacing with external code

Some neural simulations benefit from a direct connections to external libraries, e.g. to support real-time input from a sensor (but note that Brian currently does not offer facilities to assure real-time processing) or to perform complex calculations during a simulation run.

If the external library is written in Python (or is a library with Python bindings), then the connection can be made either using the mechanism for User-provided functions, or using a network operation.

In case of C/C++ libraries, only the User-provided functions mechanism can be used. On the other hand, such simulations can use the same user-provided C++ code to run both with the runtime weave target and with the Standalone code generation mode. In addition to that code, one generally needs to include additional header files and use compiler/linker options to interface with the external code. For this, several preferences can be used that will be taken into account for weave, cython and the cpp_standalone device. These preferences are mostly equivalent to the respective keyword arguments for Python’s distutils.core.Extension class, see the documentation of the cpp_prefs module for more details.
Examples

5.1 Example: COBAHH

This is an implementation of a benchmark described in the following review paper:


Benchmark 3: random network of HH neurons with exponential synaptic conductances

Clock-driven implementation (no spike time interpolation)

R. Brette - Dec 2007

```python
from brian2 import *

# Parameters
area = 20000*umetre**2
Cm = (1*ufarad*cm**-2) * area
gl = (5e-5*siemens*cm**-2) * area

E1 = -60*mV
EK = -90*mV
ENa = 50*mV
g_na = (100*msiemens*cm**-2) * area
g_kd = (30*msiemens*cm**-2) * area
VT = -63*mV

# Time constants
taue = 5*ms
taui = 10*ms

# Reversal potentials
Ee = 0*mV
Ei = -80*mV
```

(continues on next page)
we = 6*nS  # excitatory synaptic weight
wi = 67*nS  # inhibitory synaptic weight

# The model
eqs = Equations('''
dv/dt = (gl*(El-v)+ge*(Ee-v)+gi*(Ei-v)-
g_na*(m*m*m)*h*(v-ENa)-
g_kd*(n*n*n*n)*(v-EK))/Cm : volt
dm/dt = alpha_m*(1-m)-beta_m*m : 1
dn/dt = alpha_n*(1-n)-beta_n*n : 1
dh/dt = alpha_h*(1-h)-beta_h*h : 1
dge/dt = -ge*(1./taue) : siemens
dgi/dt = -gi*(1./taui) : siemens
alpha_m = 0.32*(mV**-1)*4*mV/exprel((13*mV-v+VT)/(4*mV))/ms : Hz
beta_m = 0.28*(mV**-1)*5*mV/exprel((v-VT-40*mV)/(5*mV))/ms : Hz
alpha_h = 0.128*exp((17*mV-v+VT)/(18*mV))/ms : Hz
beta_h = 4./(1+exp((40*mV-v+VT)/(5*mV)))/ms : Hz
alpha_n = 0.032*(mV**-1)*5*mV/exprel((15*mV-v+VT)/(5*mV))/ms : Hz
beta_n = .5*exp((10*mV-v+VT)/(40*mV))/ms : Hz
''')

P = NeuronGroup(4000, model=eqs, threshold='v>-20*mV', refractory=3*ms,
method='exponential_euler')

Pe = P[:3200]
Pi = P[3200:]
Ce = Synapses(Pe, P, on_pre='ge+=we')
Ci = Synapses(Pi, P, on_pre='gi+=wi')
Ce.connect(p=0.02)
Ci.connect(p=0.02)

# Initialization
P.v = 'El + (randn() * 5 - 5)*mV'
P.ge = '((randn() * 1.5 + 4) * 10.*nS'
P.gi = 'randn() * 12 + 20) * 10.*nS'

# Record a few traces
trace = StateMonitor(P, 'v', record=[1, 10, 100])
run(1 * second, report='text')
plot(trace.t/ms, trace[1].v/mV)
plot(trace.t/ms, trace[10].v/mV)
plot(trace.t/ms, trace[100].v/mV)
xlabel('t (ms)')
ylabel('v (mV)')
show()
5.2 Example: CUBA

This is a Brian script implementing a benchmark described in the following review paper:


Benchmark 2: random network of integrate-and-fire neurons with exponential synaptic currents.

Clock-driven implementation with exact subthreshold integration (but spike times are aligned to the grid).

```python
from brian2 import *

taum = 20*ms
taue = 5*ms
taui = 10*ms
Vt = -50*mV
Vr = -60*mV
El = -49*mV

eqs = '''
   dv/dt = (ge+gi-(v-El))/taum : volt (unless refractory)
   (continues on next page)'''
```

5.2. Example: CUBA 201
\[
\frac{d e}{d t} = -\frac{e}{\tau_e} \text{ volt} \\
\frac{d i}{d t} = -\frac{i}{\tau_i} \text{ volt}
\]

\[
P = \text{NeuronGroup}(4000, \text{eqs}, \text{threshold='v>Vt'}, \text{reset='v = Vr'}, \text{refractory=5*ms, method='exact'}))
\]

\[
P.v = 'Vr + \text{rand()} \times (Vt - Vr)'
\]

\[
P.ge = 0\text{mV} \\
P.gi = 0\text{mV}
\]

\[
we = (60\times0.27/10)\text{mV} \# \text{excitatory synaptic weight (voltage)}
\]

\[
wi = (-20\times4.5/10)\text{mV} \# \text{inhibitory synaptic weight}
\]

\[
Ce = \text{Synapses}(P, P, \text{on_pre='ge += we'})
\]

\[
Ci = \text{Synapses}(P, P, \text{on_pre='gi += wi'})
\]

\[
Ce\text{.connect('i<3200', p=0.02)}
\]

\[
Ci\text{.connect('i>=3200', p=0.02)}
\]

\[
s\_\text{mon} = \text{SpikeMonitor}(P)
\]

\[
\text{run}(1 \times \text{second})
\]

\[
\text{plot}(s\_\text{mon}.t/\text{ms}, s\_\text{mon}.i, ',k')
\]

\[
\text{xlabel('Time (ms)')}
\]

\[
\text{ylabel('Neuron index')}
\]

\[
\text{show()}
\]
5.3 Example: IF_curve_Hodgkin_Huxley

Input-Frequency curve of a HH model.

Network: 100 unconnected Hodgkin-Huxley neurons with an input current I. The input is set differently for each neuron. This simulation should use exponential Euler integration.

```python
from brian2 import *
	num_neurons = 100

duration = 2*second

# Parameters
area = 20000*umetre**2
Cm = 1*ufarad*cm**-2 * area
gl = 5e-5*siemens*cm**-2 * area
El = -65*mV
EK = -90*mV
ENa = 50*mV
g_na = 100*msiemens*cm**-2 * area
g_kd = 30*msiemens*cm**-2 * area
VT = -63*mV

# The model
eqs = Equations(''
    dv/dt = (gl*(El-v) - g_na*(m*m*m)*h*(v-ENa) - g_kd*(n*n*n*n)*(v-EK) + I)/Cm : volt
    dm/dt = 0.32*(mV**-1)*4*mV/exprel((13.*mV-v+VT)/(4*mV))/ms*(1-m)-0.28*(mV**-1)*5*mV/
            ->exprel((v-VT-40.*mV)/(5*mV))/ms*m : 1
    dn/dt = 0.032*(mV**-1)*5*mV/exprel((15.*mV-v+VT)/(5*mV))/ms*(1-n)-.5*exp((10.*mV-
            ->v+VT)/(40.*mV))/ms*n : 1
    dh/dt = 0.128*exp((17.*mV-v+VT)/(18.*mV))/ms*(1-h)-4./(1+exp((40.*mV-v+VT)/(5.*mV)))/
            ->ms*h : 1
    I : amp
''')

# Threshold and refractoriness are only used for spike counting
group = NeuronGroup(num_neurons, eqs,
    threshold='v > -40*mV',
    refractory='v > -40*mV',
    method='exponential_euler')

group.v = El
group.I = '0.7*nA * i / num_neurons'

monitor = SpikeMonitor(group)

run(duration)

plot(group.I/nA, monitor.count / duration)
xlabel('I (nA)')
ylabel('Firing rate (sp/s)')
show()
```

5.3. Example: IF_curve_Hodgkin_Huxley
5.4 Example: IF_curve_LIF

Input-Frequency curve of a IF model.

Network: 1000 unconnected integrate-and-fire neurons (leaky IF) with an input parameter v0. The input is set differently for each neuron.

```python
from brian2 import *

n = 1000
duration = 1*second
tau = 10*ms
eqs = '``
   dv/dt = (v0 - v) / tau : volt (unless refractory)
v0 : volt
   '``

group = NeuronGroup(n, eqs, threshold='v > 10*mV', reset='v = 0*mV',
   refractory=5*ms, method='exact')

monitor = SpikeMonitor(group)
```
(continues on next page)
run(duration)
plot(group.v0/mV, monitor.count / duration)
xlabel('v0 (mV)')
ylabel('Firing rate (sp/s)')
show()

5.5 Example: adaptive_threshold

A model with adaptive threshold (increases with each spike)

```python
from brian2 import *

eqs = '''
dv/dt = -v/(10*ms) : volt
dvt/dt = (10*mV-vt)/(15*ms) : volt
'''

reset = ''
v = 0*mV
```
vt += 3*mV
```

IF = NeuronGroup(1, model=eqs, reset=reset, threshold='v>vt',
    method='exact')
IF.vt = 10*mV
PG = PoissonGroup(1, 500 * Hz)

C = Synapses(PG, IF, on_pre='v += 3*mV')
C.connect()

Mv = StateMonitor(IF, 'v', record=True)
Mvt = StateMonitor(IF, 'vt', record=True)
# Record the value of v when the threshold is crossed
M_crossings = SpikeMonitor(IF, variables='v')
run(2*second, report='text')

subplot(1, 2, 1)
plot(Mv.t / ms, Mv[0].v / mV)
plot(Mvt.t / ms, Mvt[0].vt / mV)
ylabel('v (mV)')
xlabel('t (ms)')
# zoom in on the first 100ms
xlim(0, 100)
subplot(1, 2, 2)
hist(M_crossings.v / mV, bins=np.arange(10, 20, 0.5))
xlabel('v at threshold crossing (mV)')
show()
5.6 Example: non_reliability

Reliability of spike timing.

See e.g. Mainen & Sejnowski (1995) for experimental results in vitro.

Here: a constant current is injected in all trials.

```python
from brian2 import *

N = 25
tau = 20*ms
sigma = .015
eqs_neurons = '''
dx/dt = (1.1 - x) / tau + sigma * (2 / tau)**.5 * xi : 1 (unless refractory)
'''
neurons = NeuronGroup(N, model=eqs_neurons, threshold='x > 1', reset='x = 0',
                        refractory=5*ms, method='euler')
spikes = SpikeMonitor(neurons)
run(500*ms)
plot(spikes.t/ms, spikes.i, '.k')
```

(continues on next page)
5.7 Example: phase_locking

Phase locking of IF neurons to a periodic input.

```python
from brian2 import *

tau = 20*ms
n = 100
b = 1.2  # constant current mean, the modulation varies
freq = 10*Hz

eqs = '"
    dv/dt = (-v + a * sin(2 * pi * freq * t) + b) / tau : 1
    a : 1
"

neurons = NeuronGroup(n, model=eqs, threshold='v > 1', reset='v = 0',
```

method='euler')
neurons.v = 'rand()' 
neurons.a = '0.05 + 0.7*i/n'
S = SpikeMonitor(neurons)
trace = StateMonitor(neurons, 'v', record=50)

run(1000*ms)
subplot(211)
plot(S.t/ms, S.i, '.k')
xlabel('Time (ms)')
ylabel('Neuron index')
subplot(212)
plot(trace.t/ms, trace.v.T)
xlabel('Time (ms)')
ylabel('v')
tight_layout()
show()

5.8 Example: reliability

Reliability of spike timing.
See e.g. Mainen & Sejnowski (1995) for experimental results in vitro.

```python
from brian2 import *

# The common noisy input
N = 25
tau_input = 5*ms
input = NeuronGroup(1, 'dx/dt = -x / tau_input + (2 /tau_input)**.5 * xi : 1')

# The noisy neurons receiving the same input
tau = 10*ms
sigma = .015
eqs_neurons = '''
dx/dt = (0.9 + .5 * I - x) / tau + sigma * (2 / tau)**.5 * xi : 1
I : 1 (linked)
'''
neurons = NeuronGroup(N, model=eqs_neurons, threshold='x > 1',
                        reset='x = 0', refractory=5*ms, method='euler')
neurons.x = 'rand()'
neurons.I = linked_var(input, 'x') # input.x is continuously fed into neurons.I
spikes = SpikeMonitor(neurons)

run(500*ms)
plt.plot(spikes.t/ms, spikes.i, '.k')
xlabel('Time (ms)')
ylabel('Neuron index')
show()
```
5.9 advanced

5.9.1 Example: compare_GSL_to_conventional

Example using GSL ODE solvers with a variable time step and comparing it to the Brian solver.

For highly accurate simulations, i.e. simulations with a very low desired error, the GSL simulation with a variable time step can be faster because it uses a low time step only when it is necessary. In biologically detailed models (e.g. of the Hodgkin-Huxley type), the relevant time constants are very short around an action potential, but much longer when the neuron is near its resting potential. The following example uses a very simple neuron model (leaky integrate-and-fire), but simulates a change in relevant time constants by changing the actual time constant every 10ms, independently for each of 100 neurons. To accurately simulate this model with a fixed time step, the time step has to be very small, wasting many unnecessary steps for all the neurons where the time constant is long.

Note that using the GSL ODE solver is much slower, if both methods use a comparable number of steps, i.e. if the desired accuracy is low enough so that a single step per “Brian time step” is enough.

```python
from brian2 import *
import time

# Run settings
```
```python
start_dt = .1 * ms
method = 'rk2'
error = 1.e-6  # requested accuracy

def runner(method, dt, options=None):
    seed(0)
    I = 5
    group = NeuronGroup(100, '''dv/dt = (-v + I)/tau : 1
tau : second''',
                        method=method,
                        method_options=options,
                        dt=dt)
    group.run_regularly('''v = rand()
tau = 0.1*ms + rand()*9.9*ms''', dt=10*ms)
    rec_vars = ['v', 'tau']
    if 'gsl' in method:
        rec_vars += ['_step_count']
    net = Network(group)
    net.run(0 * ms)
    mon = StateMonitor(group, rec_vars, record=True, dt=start_dt)
    net.add(mon)
    start = time.time()
    net.run(1 * second)
    mon.add_attribute('run_time')
    mon.run_time = time.time() - start
    return mon

lin = runner('linear', start_dt)
method_options = {'save_step_count': True,
                  'absolute_error': error,
                  'max_steps': 10000}
gsl = runner('gsl_%s' % method, start_dt, options=method_options)

print("Running with GSL integrator and variable time step:")
print('Run time: %.3f s' % gsl.run_time)

# check gsl error
assert np.max(np.abs(lin.v - gsl.v)) < error, "Maximum error gsl integration too large: %f" % np.max(np.abs(lin.v - gsl.v))
print("average step count: %.1f" % np.mean(gsl._step_count))
print("average absolute error: %g" % np.mean(np.abs(gsl.v - lin.v)))

print("Running with exact integration and fixed time step:")
dt = start_dt
count = 0
dts = []
avg_errors = []
max_errors = []
runtimes = []
while True:
    print('Using dt: %s' % str(dt))
    brian = runner(method, dt)
    print(\tRun time: %.3f s' % brian.run_time)
```

(continues on next page)
avg_errors.append(np.mean(np.abs(brian.v - lin.v)))
max_errors.append(np.max(np.abs(brian.v - lin.v)))
dts.append(dt)
runtimes.append(brian.run_time)
if np.max(np.abs(brian.v - lin.v)) > error:
    print('Error too high (%g), decreasing dt' % np.max(np.abs(brian.v - lin.v)))
    dt *= .5
count += 1
else:
    break
print("Desired error level achieved:")
print("average step count: %.2fs" % (start_dt / dt))
print("average absolute error: %.1g" % np.mean(np.abs(brian.v - lin.v)))
print('Run time: %.3fs' % brian.run_time)
if brian.run_time > gsl.run_time:
    print("This is %.1f times slower than the simulation with GSL's variable "
        "time step method." % (brian.run_time / gsl.run_time))
else:
    print("This is %.1f times faster than the simulation with GSL's variable "
        "time step method." % (gsl.run_time / brian.run_time))

fig, (ax1, ax2) = plt.subplots(1, 2)
ax2.axvline(1e-6, color='gray')
for label, gsl_error, std_errors, ax in [('average absolute error', np.mean(np.abs(gsl.v - lin.v))), ('maximum absolute error', np.max(np.abs(gsl.v - lin.v))), avg_errors, ax1),
    ax.set(xscale='log', yscale='log')
    ax.plot([], [], 'o', color='C0', label='fixed time step')  # for the legend entry
    for (error, runtime, dt) in zip(std_errors, runtimes, dts):
        ax.plot(error, runtime, 'o', color='C0')
        ax.annotate('%s' % str(dt), xy=(error, runtime), xytext=(2.5, 5),
            textcoords='offset points', color='C0')
    ax.plot(gsl_error, gsl.run_time, 'o', color='C1', label='variable time step (GSL)')
ax1.set_xlabel(label, xlim=(10**-10, 10**1))
ax1.set_ylabel('runtime (s)')
ax2.legend(loc='lower left')
plt.show()
5.9.2 Example: custom_events

Example demonstrating the use of custom events.

Here we have three neurons, the first is Poisson spiking and connects to neuron G, which in turn connects to neuron H. Neuron G has two variables v and g, and the incoming Poisson spikes cause an instantaneous increase in variable g. g decays rapidly, and in turn causes a slow increase in v. If v crosses a threshold, it causes a standard spike and reset. If g crosses a threshold, it causes a custom event \texttt{gspike}, and if it returns below that threshold it causes a custom event \texttt{end\_gspike}. The standard spike event when v crosses a threshold causes an instantaneous increase in variable x in neuron H (which happens through the standard \texttt{pre} pathway in the synapses), and the gspike event causes an increase in variable y (which happens through the custom pathway \texttt{gpath}).

```python
from brian2 import *

# Input Poisson spikes
inp = PoissonGroup(1, rates=250*Hz)

# First group G

# eqs_G = '...
dv/dt = (g-v)/(50*ms) : 1
dg/dt = -g/(10*ms) : 1
allow_gspike : boolean

G = NeuronGroup(1, eqs_G, threshold='v>1',
```

(continues on next page)
reset='v = 0; g = 0; allow_gspike = True';
events={'gspike': 'g>1 and allow_gspike',
       'end_gspike': 'g<1 and not allow_gspike'}
G.run_on_event('gspike', 'allow_gspike = False')
G.run_on_event('end_gspike', 'allow_gspike = True')

# Second group H
eqs_H = '''
dx/dt = -x/(10*ms) : 1
dy/dt = -y/(10*ms) : 1
'''
H = NeuronGroup(1, eqs_H)

# Synapses from input Poisson group to G
Sin = Synapses(inp, G, on_pre='g += 0.5')
Sin.connect()

# Synapses from G to H
S = Synapses(G, H,
on_pre={'pre': 'x += 1',
       'gpath': 'y += 1'},
on_event={'pre': 'spike',
         'gpath': 'gspike'})
S.connect()

# Monitors
Mstate = StateMonitor(G, ('v', 'g'), record=True)
Mgspike = EventMonitor(G, 'gspike', 'g')
Mspike = SpikeMonitor(G, 'v')
MHstate = StateMonitor(H, ('x', 'y'), record=True)

# Initialise and run
G.allow_gspike = True
run(500*ms)

# Plot
figure(figsize=(10, 4))
subplot(121)
plot(Mstate.t/ms, Mstate.g[0], '-g', label='g')
plot(Mstate.t/ms, Mstate.v[0], '-b', lw=2, label='V')
plot(Mspike.t/ms, Mspike.v, 'ob', label='_nolegend_')
plot(Mgspike.t/ms, Mgspike.g, 'og', label='_nolegend_')
xlabel('Time (ms)')
title('Presynaptic group G')
legend(loc='best')
subplot(122)
plot(MHstate.t/ms, MHstate.y[0], '-r', label='y')
plot(MHstate.t/ms, MHstate.x[0], '-k', lw=2, label='x')
xlabel('Time (ms)')
title('Postsynaptic group H')
legend(loc='best')
tight_layout()
show()
5.9.3 Example: exprel_function

Show the improved numerical accuracy when using the `exprel()` function in rate equations.

Rate equations for channel opening/closing rates often include a term of the form \( \frac{x}{\exp(x)-1} \). This term is problematic for two reasons:

- It is not defined for \( x = 0 \) (where it should equal to 1 for continuity);
- For values \( x \approx 0 \), there is a loss of accuracy.

For better accuracy, and to avoid issues at \( x = 0 \), Brian provides the function `exprel()`, which is equivalent to \( \frac{\exp(x)-1}{x} \), but with better accuracy and the expected result at \( x = 0 \). In this example, we demonstrate the advantage of expressing a typical rate equation from the HH model with `exprel()`.

```python
from brian2 import *

# Dummy group to evaluate the rate equation at various points
eqs = '''v : volt
    # opening rate from the HH model
    alpha_simple = 0.32*(mV**-1)*(-50*mV-v)/(exp((-50*mV-v)/(4*mV))-1.)/ms : Hz
    alpha_improved = 0.32*(mV**-1)*4*mV/exprel((-50*mV-v)/(4*mV))/ms : Hz'''
neuron = NeuronGroup(1000, eqs)

# Use voltage values around the problematic point
neuron.v = np.linspace(-50 - .5e-6, -50 + .5e-6, len(neuron))*mV

fig, ax = plt.subplots()
ax.plot((neuron.v + 50*mV)/nvolt, neuron.alpha_simple, '.', label=r'$\alpha_{\text{simple}}$')
ax.plot((neuron.v + 50*mV)/nvolt, neuron.alpha_improved, 'k', label=r'$\alpha_{\text{improved}}$')
ax.legend()
ax.set(xlabel='$v$ relative to -50mV (nV)', ylabel=r'$\alpha$ (Hz)')
ax.ticklabel_format(useOffset=False)
plt.tight_layout()
plt.show()
```
5.9.4 Example: float_32_64_benchmark

Benchmark showing the performance of float32 versus float64.

```python
from brian2 import *
from brian2.devices.device import reset_device, reinit_devices

# CUBA benchmark
def run_benchmark(name):
    if name=='CUBA':
        taum = 20*ms
        taue = 5*ms
        taui = 10*ms
        Vt = -50*mV
        Vr = -60*mV
        El = -49*mV

        eqs = ''
        dv/dt  = (ge+gi-(v-El))/taum : volt (unless refractory)
        dge/dt = -ge/taue : volt
        dgi/dt = -gi/taui : volt
```

(continues on next page)
P = NeuronGroup(4000, model=eqs, threshold='v>-20*mV', refractory=3*ms, method='exponential_euler')

(continues on next page)
Pe = P[0:3200]
Pi = P[3200:]
Ce = Synapses(Pe, P, on_pre='ge+=we')
Ci = Synapses(Pi, P, on_pre='gi+=wi')
Ce.connect(p=0.02)
Ci.connect(p=0.02)

# Initialization
P.v = 'El + (randn() * 5 - 5)*mV'
P.ge = '(randn() * 1.5 + 4) * 10.*nS'
P.gi = '(randn() * 12 + 20) * 10.*nS'

run(1 * second, profile=True)

return sum(t for name, t in magic_network.profiling_info)

def generate_results(num_repeats):
    results = {}

    for name in ['CUBA', 'COBA']:
        for target in ['numpy', 'cython', 'weave']:
            for dtype in [float32, float64]:
                prefs.codegen.target = target
                prefs.core.default_float_dtype = dtype
                times = [run_benchmark(name) for repeat in range(num_repeats)]
                results[name, target, dtype.__name__] = amin(times)

    for name in ['CUBA', 'COBA']:
        for dtype in [float32, float64]:
            times = []
            for _ in range(num_repeats):
                reset_device()
                reinit_devices()
                set_device('cpp_standalone', directory=None, with_output=False)
                prefs.core.default_float_dtype = dtype
                times.append(run_benchmark(name))
                results[name, 'cpp_standalone', dtype.__name__] = amin(times)

    return results

results = generate_results(3)

bar_width = 0.9
names = ['CUBA', 'COBA']
targets = ['numpy', 'cython', 'weave', 'cpp_standalone']
precisions = ['float32', 'float64']

figure(figsize=(8, 8))
for j, name in enumerate(names):
    subplot(2, 2, 1+2*j)
    title(name)
    index = arange(len(targets))
    for i, precision in enumerate(precisions):
        bar(index+i*bar_width/len(precisions),
            [results[name, target, precision] for target in targets],
            bar_width/len(precisions), label=precision, align='edge')
ylabel('Time (s)')
if j:
    xticks(index+0.5*bar_width, targets, rotation=45)
else:
    xticks(index+0.5*bar_width, ('',)*len(targets))
    legend(loc='best')

subplot(2, 2, 2+2*j)
index = arange(len(precisions))
for i, target in enumerate(targets):
    bar(index+i*bar_width/len(targets),
        [results[name, target, precision] for precision in precisions],
        bar_width/len(targets), label=target, align='edge')
ylabel('Time (s)')
if j:
    xticks(index+0.5*bar_width, precisions, rotation=45)
else:
    xticks(index+0.5*bar_width, ('',)*len(precisions))
    legend(loc='best')
tight_layout()
show()

5.9.5 Example: opencv_movie

An example that uses a function from external C library (OpenCV in this case). Works for all C-based code generation targets (i.e. for weave and cpp_standalone device) and for numpy (using the Python bindings).

This example needs a working installation of OpenCV 3.x and its Python bindings. It has been tested on 64 bit Linux in a conda environment with packages from the conda-forge channels (opencv 3.4.4, x264 1!152.20180717, ffmpeg 4.1).

import os
import urllib2
import cv2  # Import OpenCV2

from brian2 import *

defaultclock.dt = 1*ms
prefs.codegen.target = 'weave'
prefs.logging.std_redirection = False
set_device('cpp_standalone', clean=True)
filename = os.path.abspath('Megamind.avi')

if not os.path.exists(filename):
    print('Downloading the example video file')
    response = urllib2.urlopen('http://docs.opencv.org/2.4/_downloads/Megamind.avi')
    data = response.read()
    with open(filename, 'wb') as f:
        f.write(data)

video = cv2.VideoCapture(filename)
width, height, frame_count = (int(video.get(cv2.CAP_PROP_FRAME_WIDTH)),
    int(video.get(cv2.CAP_PROP_FRAME_HEIGHT)),
(continues on next page)
fps = 24

time_between_frames = 1/second/fps

@implementation

('cpp', ''

double* get_frame(bool new_frame)
{
    // The following initializations will only be executed once
    static cv::VideoCapture source("VIDEO_FILENAME");
    static cv::Mat frame;
    static double* grayscale_frame = (double*)malloc(VIDEO_WIDTH*VIDEO_HEIGHT*sizeof(double));
    if (new_frame)
    {
        source >> frame;
        double mean_value = 0;
        for (int row=0; row<VIDEO_HEIGHT; row++)
            for (int col=0; col<VIDEO_WIDTH; col++)
            {
                const double grayscale_value = (frame.at<cv::Vec3b>(row, col)[0] +
                                                  frame.at<cv::Vec3b>(row, col)[1] +
                                                  frame.at<cv::Vec3b>(row, col)[2])/(3.0*128);
                mean_value += grayscale_value / (VIDEO_WIDTH * VIDEO_HEIGHT);
                grayscale_frame[row*VIDEO_WIDTH + col] = grayscale_value;
            }
        // subtract the mean
        for (int i=0; i<VIDEO_HEIGHT*VIDEO_WIDTH; i++)
            grayscale_frame[i] -= mean_value;
    }
    return grayscale_frame;
}

double video_input(const int x, const int y)
{
    // Get the current frame (or a new frame in case we are asked for the first
    // element
    double* frame = get_frame(x==0 && y==0);
    return frame[y*VIDEO_WIDTH + x];
}
'''.replace('VIDEO_FILENAME', filename),

libraries=['opencv_core',
           'opencv_highgui',
           'opencv_videoio'],

headers=['<opencv2/core/core.hpp>',
          '<opencv2/highgui/highgui.hpp>'],

declare_macros=[('VIDEO_WIDTH', width),
                 ('VIDEO_HEIGHT', height)]

@check_units(x=1, y=1, result=1)
def video_input(x, y):
    # we assume this will only be called in the custom operation (and not for
    # example in a reset or synaptic statement), so we don't need to do indexing
    # but we can directly return the full result
    _, frame = video.read()
    grayscale = frame.mean(axis=2)
    grayscale /= 128.  # scale everything between 0 and 2
    return grayscale.ravel() - grayscale.ravel().mean()}
N = width * height
tau, tau_th = 10*ms, time_between_frames
G = NeuronGroup(N, '''
    dv/dt = (-v + I)/tau : 1
    dv_th/dt = -v_th/tau_th : 1
    row : integer (constant)
    column : integer (constant)
    I : 1 # input current''',
    threshold='v>v_th', reset='v=0; v_th = 3*v_th + 1.0',
    method='exact')
G.v_th = 1
G.row = 'i//width'
G.column = 'i%width'

G.run_regularly('I = video_input(column, row)',
    dt=time_between_frames)
mon = SpikeMonitor(G)
runtime = frame_count*time_between_frames
run(runtime, report='text')

# Avoid going through the whole Brian2 indexing machinery too much
i, t, row, column = mon.i[:], mon.t[:], G.row[:], G.column[:]
import matplotlib.animation as animation
# TODO: Use overlapping windows
stepsize = 100*ms

def next_spikes():
    step = next_spikes.step
    if step*stepsize > runtime:
        next_spikes.step=0
        raise StopIteration()
    spikes = i[(t>=step*stepsize) & (t<(step+1)*stepsize)]
    next_spikes.step += 1
    yield column[spikes], row[spikes]
next_spikes.step = 0

fig, ax = plt.subplots()
dots, = ax.plot([], [], 'k.', markersize=2, alpha=.25)
ax.set_xlim(0, width)
ax.set_ylim(0, height)
ax.invert_yaxis()

def run(data):
    x, y = data
    dots.set_data(x, y)

ani = animation.FuncAnimation(fig, run, next_spikes, blit=False, repeat=True,
    repeat_delay=1000)
plt.show()

5.9.6 Example: stochastic_odes

Demonstrate the correctness of the “derivative-free Milstein method” for multiplicative noise.
from brian2 import *
# We only get exactly the same random numbers for the exact solution and the
# simulation if we use the numpy code generation target
prefs.codegen.target = 'numpy'

# setting a random seed makes all variants use exactly the same Wiener process
seed = 12347

X0 = 1
mu = 0.5/second # drift
sigma = 0.1/second # diffusion
runtime = 1*second

def simulate(method, dt):
    '''simulate geometrical Brownian with the given method'''
    np.random.seed(seed)
    G = NeuronGroup(1, 'dX/dt = (mu - 0.5*second*sigma**2)*X + X*sigma*xi*second**0.5: 
˓→1',
                   dt=dt, method=method)
    G.X = X0
    mon = StateMonitor(G, 'X', record=True)
    net = Network(G, mon)
    net.run(runtime)
    return mon.t_[:], mon.X.flatten()

def exact_solution(t, dt):
    '''Return the exact solution for geometrical Brownian motion at the given
time points'''
    # Remove units for simplicity
    my_mu = float(mu)
    my_sigma = float(sigma)
    dt = float(dt)
    t = asarray(t)
    np.random.seed(seed)
    # We are calculating the values at the *start* of a time step, as when using
    # a StateMonitor. Therefore the Brownian motion starts with zero
    brownian = np.hstack([0, cumsum(sqrt(dt) * np.random.randn(len(t)-1))])
    return (X0 * exp((my_mu - 0.5*my_sigma**2)*t + dt) + my_sigma*brownian)

def main():
    figure(1, figsize=(16, 7))
    figure(2, figsize=(16, 7))
    methods = ['milstein', 'heun']
    dts = [1*ms, 0.5*ms, 0.2*ms, 0.1*ms, 0.05*ms, 0.025*ms, 0.01*ms, 0.005*ms]
    rows = floor(sqrt(len(dts)))
    cols = ceil(1.0 * len(dts) / rows)
    errors = dict([(method, zeros(len(dts))) for method in methods])
    (continues on next page)
for dt_idx, dt in enumerate(dts):
    print('dt: $s' % dt)
trajectories = []
# Test the numerical methods
for method in methods:
    t, trajectories[method] = simulate(method, dt)
# Calculate the exact solution
exact = exact_solution(t, dt)

for method in methods:
    # plot the trajectories
    figure(1)
    subplot(rows, cols, dt_idx+1)
    plot(t, trajectories[method], label=method, alpha=0.75)

    # determine the mean absolute error
    errors[method][dt_idx] = mean(abs(trajectories[method] - exact))

    # plot the difference to the real trajectory
    figure(2)
    subplot(rows, cols, dt_idx+1)
    plot(t, trajectories[method] - exact, label=method, alpha=0.75)

    figure(1)
    plot(t, exact, color='gray', lw=2, label='exact', alpha=0.75)
    title('dt = $s' % str(dt))
xticks([])

figure(1)
legend(frameon=False, loc='best')
tight_layout()

figure(2)
legend(frameon=False, loc='best')
tight_layout()

figure(3)
for method in methods:
    plot(array(dts) / ms, errors[method], 'o', label=method)
legend(frameon=False, loc='best')
yscale('log')
xscale('log')
xlabel('dt (ms)')
ylabel('Mean absolute error')
tight_layout()
show()
5.10 compartmental

5.10.1 Example: bipolar_cell

A pseudo MSO neuron, with two dendrites and one axon (fake geometry).

```python
from brian2 import *

# Morphology
morpho = Soma(30*um)
morpho.axon = Cylinder(diameter=1*um, length=300*um, n=100)
morpho.L = Cylinder(diameter=1*um, length=100*um, n=50)
morpho.R = Cylinder(diameter=1*um, length=150*um, n=50)

# Passive channels
gL = 1e-4*siemens/cm**2
EL = -70*mV
eqs='''
Im = gL * (EL - v) : amp/meter**2
I : amp (point current)

(continues on next page)
```
neuron = SpatialNeuron(morphology=morpho, model=eqs,
                      Cm=1*uF/cm**2, Ri=100*ohm*cm, method='exponential_euler')
neuron.v = EL
neuron.I = 0*amp

# Monitors
mon_soma = StateMonitor(neuron, 'v', record=[0])
mon_L = StateMonitor(neuron.L, 'v', record=True)
mon_R = StateMonitor(neuron, 'v', record=morpho.R[75*um])

run(1*ms)
neuron.I[morpho.L[50*um]] = 0.2*nA  # injecting in the left dendrite
run(5*ms)
neuron.I = 0*amp
run(50*ms, report='text')

subplot(211)
plot(mon_L.t/ms, mon_soma[0].v/mV, 'k')
plot(mon_L.t/ms, mon_L[morpho.L[50*um]].v/mV, 'r')
plot(mon_L.t/ms, mon_R[morpho.R[75*um]].v/mV, 'b')
ylabel('v (mV)')
subplot(212)
for x in linspace(0*um, 100*um, 10, endpoint=False):
    plot(mon_L.t/ms, mon_L[morpho.L[x]].v/mV)
xlabel('Time (ms)')
ylabel('v (mV)')
show()
5.10.2 Example: bipolar_with_inputs

A pseudo MSO neuron, with two dendrites (fake geometry). There are synaptic inputs.

```python
from brian2 import *

# Morphology
morpho = Soma(30*um)
morpho.L = Cylinder(diameter=1*um, length=100*um, n=50)
morpho.R = Cylinder(diameter=1*um, length=100*um, n=50)

# Passive channels
gL = 1e-4*siemens/cm**2
EL = -70*mV
Es = 0*mV
eqs=''
Im = gL*(EL-v) : amp/meter**2
Is = gs*(Es-v) : amp (point current)
gs : siemens
''

neuron = SpatialNeuron(morphology=morpho, model=eqs,
```
Cm=1*uF/cm**2, Ri=100*ohm*cm, method='exponential_euler')

neuron.v = EL

# Regular inputs
stimulation = NeuronGroup(2, 'dx/dt = 300*Hz : 1', threshold='x>1', reset='x=0',
method='euler')
stimulation.x = [0, 0.5]    # Asynchronous

# Synapses
taus = 1*ms
w = 20*nS
S = Synapses(stimulation, neuron, model='''dg/dt = -g/taus : siemens (clock-driven)
gs_post = g : siemens (summed)'',
on_pre='g += w', method='exact')
S.connect(i=0, j=morpho.L[-1])
S.connect(i=1, j=morpho.R[-1])

# Monitors
mon_soma = StateMonitor(neuron, 'v', record=[0])
mon_L = StateMonitor(neuron.L, 'v', record=True)
mon_R = StateMonitor(neuron.R, 'v',
record=morpho.R[-1])
run(50*ms, report='text')

subplot(211)
plot(mon_L.t/ms, mon_soma[0].v/mV, 'k')
plot(mon_L.t/ms, mon_L[morpho.L[-1]].v/mV, 'r')
plot(mon_L.t/ms, mon_R[morpho.R[-1]].v/mV, 'b')
ylabel('v (mV)')
subplot(212)
for x in linspace(0*um, 100*um, 10, endpoint=False):
    plot(mon_L.t/ms, mon_L[morpho.L[x]].v/mV)
xlabel('Time (ms)')
ylabel('v (mV)')
show()
5.10.3 Example: bipolar_with_inputs2

A pseudo MSO neuron, with two dendrites (fake geometry). There are synaptic inputs.

Second method.

```python
from brian2 import *

# Morphology
morpho = Soma(30*um)
morpho.L = Cylinder(diameter=1*um, length=100*um, n=50)
morpho.R = Cylinder(diameter=1*um, length=100*um, n=50)

# Passive channels
gL = 1e-4*siemens/cm**2
EL = -70*mV
Es = 0*mV
taus = 1*ms
eqs=''
Im = gL*(EL-v) : amp/meter**2
Is = gs*(Es-v) : amp (point current)
dgs/dt = -gs/taus : siemens
```
neuron = SpatialNeuron(morphology=morpho, model=eqs,
    Cm=1*uF/cm**2, Ri=100*ohm*cm, method='exponential_euler')
neuron.v = EL

# Regular inputs
stimulation = NeuronGroup(2, 'dx/dt = 300*Hz : 1', threshold='x>1', reset='x=0',
    method='euler')
stimulation.x = [0, 0.5] # Asynchronous

# Synapses
w = 20*nS
S = Synapses(stimulation, neuron, on_pre='gs += w')
S.connect(i=0, j=morpho.L[99.9*um])
S.connect(i=1, j=morpho.R[99.9*um])

# Monitors
mon_soma = StateMonitor(neuron, 'v', record=[0])
mon_L = StateMonitor(neuron.L, 'v', record=True)
mon_R = StateMonitor(neuron, 'v', record=morpho.R[99.9*um])
run(50*ms, report='text')

subplot(211)
plot(mon_L.t/ms, mon_soma[0].v/mV, 'k')
plot(mon_L.t/ms, mon_L[morpho.L[99.9*um]].v/mV, 'r')
plot(mon_L.t/ms, mon_R[morpho.R[99.9*um]].v/mV, 'b')
ylabel('v (mV)')

subplot(212)
for i in [0, 5, 10, 15, 20, 25, 30, 35, 40, 45]:
    plot(mon_L.t/ms, mon_L.v[i, :]/mV)
xlabel('Time (ms)')
ylabel('v (mV)')
show()
5.10.4 Example: cylinder

A short cylinder with constant injection at one end.

```python
from brian2 import *
defaultclock.dt = 0.01*ms

# Morphology
diameter = 1*um
length = 300*um
Cm = 1*uF/cm**2
Ri = 150*ohm*cm
N = 200
morpho = Cylinder(diameter=diameter, length=length, n=N)

# Passive channels
gL = 1e-4*siemens/cm**2
EL = -70*mV
eqs = '''
I_m = gL * (EL - v) : amp/meter**2
I : amp (point current)
```

(continues on next page)
'\[
\text{neuron = SpatialNeuron(morphology=morpho, model=eqs, Cm=Cm, Ri=Ri,}
\text{ method='exponential_euler')}\]
\[
\text{neuron.v = EL}\]
\]
\[
1a = neuron.space_constant[0]
\text{print("Electrotonic length: \\
\$s\" % 1a)}\]
\[
\text{neuron.I[0] = 0.02*nA \# injecting at the left end}
\text{run(100*ms, report='text')}\]
\]
\[
\text{plot(neuron.distance/um, neuron.v/mV, 'kx')}\]
\text{\# Theory}
\[
x = neuron.distance
\text{ra = la * 4 * Ri / (pi * diameter**2)}
\text{theory = EL + ra * neuron.I[0] * cosh((length - x) / la) / sinh(length / la)}
\text{plot(x/um, theory/mV, 'r')}\]
\text{xlabel('x (um)')}\]
\text{ylabel('v (mV)')}\]
\text{show()}
\]
5.10.5 Example: hh_with_spikes


Spikes are recorded along the axon, and then velocity is calculated.

```python
from brian2 import *
from scipy import stats

defaultclock.dt = 0.01*ms

morpho = Cylinder(length=10*cm, diameter=2*238*um, n=1000, type='axon')

El = 10.613*mV
ENa = 115*mV
EK = -12*mV
gl = 0.3*msiemens/cm**2
gNa0 = 120*msiemens/cm**2
gK = 36*msiemens/cm**2

# Typical equations
eqs = '''
# The same equations for the whole neuron, but possibly different parameter values
# distributed transmembrane current
Im = gl * (El-v) + gNa * m**3 * h * (ENa-v) + gK * n**4 * (EK-v) : amp/meter**2
I : amp (point current) # applied current
dm/dt = alpham * (1-m) - betam * m : 1
dn/dt = alphan * (1-n) - betan * n : 1
dh/dt = alphah * (1-h) - betah * h : 1
alpham = (0.1/mV) * 10*mV/exprel((-v+25*mV)/(10*mV))/ms : Hz
betam = 4 * exp(-v/(18*mV))/ms : Hz
alphan = (0.01/mV) * 10*mV/exprel((-v+10*mV)/(10*mV))/ms : Hz
betan = 0.125*exp(-v/(80*mV))/ms : Hz

gNa : siemens/meter**2
'''

neuron = SpatialNeuron(morphology=morpho, model=eqs, method="exponential_euler",
refractory="m > 0.4", threshold="m > 0.5",
Cm=1*uF/cm**2, Ri=35.4*ohm*cm)

neuron.v = 0*mV
neuron.h = 1
neuron.m = 0
neuron.n = .5
neuron.I = 0*amp
neuron.gNa = gNa0

M = StateMonitor(neuron, 'v', record=True)
spikes = SpikeMonitor(neuron)

run(50*ms, report='text')
neuron.I[0] = 1*uA # current injection at one end

run(3*ms)
neuron.I = 0*amp
run(50*ms, report='text')

# Calculation of velocity
```

(continues on next page)
slopes, intercept, r_value, p_value, std_err = stats.linregress(spikes.t/second, neuron.distance[spikes.i]/meter)

print("Velocity = %.2f m/s" % slope)

subplot(211)
for i in range(10):
    plot(M.t/ms, M.v.T[:, i*100]/mV)
ylabel('v')

subplot(212)
plot(spikes.t/ms, spikes.i*neuron.length[0]/cm, '.k')
plot(spikes.t/ms, (intercept+slope*(spikes.t/second))/cm, 'r')
xlabel('Time (ms)')
ylabel('Position (cm)')
show()
morpho = Cylinder(length=10*cm, diameter=2*238*um, n=1000, type='axon')

E1 = 10.613*mV
ENa = 115*mV
EK = -12*mV
gl = 0.3*siemens/cm**2
gNa0 = 120*siemens/cm**2
gK = 36*siemens/cm**2

# Typical equations
eqs = '''
# The same equations for the whole neuron, but possibly different parameter values
# distributed transmembrane current
Im = gl * (E1-v) + gNa * m**3 * h * (ENa-v) + gK * n**4 * (EK-v) : amp/meter**2
I : amp (point current) # applied current
dm/dt = alpham * (1-m) - betam * m : 1
dn/dt = alphan * (1-n) - betan * n : 1
dh/dt = alphah * (1-h) - betah * h : 1
alpham = (0.1/mV) * 10*mV/exprel((-v+25*mV)/(10*mV))/ms : Hz
betam = 4 * exp(-v/(18*mV))/ms : Hz
alphah = 0.07 * exp(-v/(20*mV))/ms : Hz
betah = 1/(exp((-v+30*mV) / (10*mV)) + 1)/ms : Hz
alphan = (0.01/mV) * 10*mV/exprel((-v+10*mV)/(10*mV))/ms : Hz
betan = 0.125*exp(-v/(80*mV))/ms : Hz
gNa : siemens/meter**2
'''

neuron = SpatialNeuron(morphology=morpho, model=eqs, Cm=1*uF/cm**2,
Ri=35.4*ohm*cm, method="exponential_euler")

neuron.v = 0*mV
neuron.h = 1
neuron.m = 0
neuron.n = .5
neuron.I = 0
neuron.gNa = gNa0
neuron[5*cm:10*cm].gNa = 0*siemens/cm**2
M = StateMonitor(neuron, 'v', record=True)

run(50*ms, report='text')
neuron.I[0] = 1*uA  # current injection at one end
run(3*ms)
neuron.I = 0*amp
run(100*ms, report='text')
for i in range(75, 125, 1):
    plot(cumsum(neuron.length)/cm, i+(1./60)*M.v[:, i*5]/mV, 'k')
yticks([])
ylabel('Time [major] v (mV) [minor]')
xlabel('Position (cm)')
axis('tight')
show()
5.10.7 Example: infinite_cable

An (almost) infinite cable with pulse injection in the middle.

```python
from brian2 import *

defaultclock.dt = 0.001*ms

# Morphology
diameter = 1*um
Cm = 1*uF/cm**2
Ri = 100*ohm*cm
N = 500
morpho = Cylinder(diameter=diameter, length=3*mm, n=N)

# Passive channels
gL = 1e-4*siemens/cm**2
EL = -70*mV
eqs = ''
I_m = gL * (EL-v) : amp/meter**2
I : amp (point current)

...''
```

(continues on next page)
neuron = SpatialNeuron(morphology=morpho, model=eqs, Cm=Cm, Ri=Ri,  
   method = 'exponential_euler')
neuron.v = EL

# membrane time constant
print("Time constant: \$s\) % taum
la = neuron.space_constant[0]
print("Characteristic length: \$s\) % la)

# Monitors
mon = StateMonitor(neuron, 'v', record=range(0, N//2, 20))

neuron.I[len(neuron) // 2] = 1*nA  # injecting in the middle
run(0.02*ms)
neuron.I = 0*amp
run(10*ms, report='text')

t = mon.t
plot(t/ms, mon.v/mV, 'k')

# Theory (incorrect near cable ends)
for i in range(0, len(neuron)//2, 20):
   x = (len(neuron)/2 - i) * morpho.length[0]
   theory = (1/(la*Cm*pi*diameter) * sqrt(taum / (4*pi*(t + defaultclock.dt)))) *  
   exp(-{t+defaultclock.dt}/taum -  
   taum / (4*(t+defaultclock.dt))*(x/la)**2))
   theory = EL + theory * 1*nA * 0.02*ms
   plot(t/ms, theory/mV, 'r')

xlabel('Time (ms)')
ylabel('v (mV)')
show()
5.10.8 Example: lfp

Hodgkin-Huxley equations (1952)

We calculate the extracellular field potential at various places.

```python
from brian2 import *
defaultclock.dt = 0.01*ms
morpho = Cylinder(x=[0, 10]*cm, diameter=2*238*um, n=1000, type='axon')

El = 10.613* mV
ENa = 115*mV
EK = -12*mV
gl = 0.3*msiemens/cm**2
gNa0 = 120*msiemens/cm**2
gK = 36*msiemens/cm**2

# Typical equations
eqs = '''
# The same equations for the whole neuron, but possibly different parameter values
# distributed transmembrane current
Im = gl * (El-v) + gNa * m**3 * h * (ENa-v) + gK * n**4 * (EK-v) : amp/meter**2
I : amp (point current) # applied current
'''
```

(continues on next page)
$$\frac{dm}{dt} = \alpha_m \cdot (1-m) - \beta_m \cdot m : 1$$

$$\frac{dn}{dt} = \alpha_n \cdot (1-n) - \beta_n \cdot n : 1$$

$$\frac{dh}{dt} = \alpha_h \cdot (1-h) - \beta_h \cdot h : 1$$

$$\alpha_m = (0.1/mV) \cdot 10*mV/\text{exprel}((-v+25*mV)/(10*mV))/ms : Hz$$

$$\beta_m = 4 \cdot \text{exp}(-v/(18*mV))/ms : Hz$$

$$\alpha_n = (0.01/mV) \cdot 10*mV/\text{exprel}((-v+10*mV)/(10*mV))/ms : Hz$$

$$\beta_n = 0.125\cdot\text{exp}(-v/(80*mV))/ms : Hz$$

$$g_{Na} : \text{siemens/meter}^2$$

```python
neuron = SpatialNeuron(morphology=morpho, model=eqs, Cm=1*uF/cm**2, 
Ri=35.4*ohm*cm, method="exponential_euler")
neuron.v = 0*mV
neuron.h = 1
neuron.m = 0
neuron.n = 0.5
neuron.I = 0
neuron.gNa = gNa0
neuron[5*cm:10*cm].gNa = 0*siemens/cm**2
M = StateMonitor(neuron, 'v', record=True)
```

```python
# LFP recorder
Ne = 5 # Number of electrodes
sigma = 0.3*siemens/meter # Resistivity of extracellular field (0.3-0.4 S/m)
lfp = NeuronGroup(Ne,model='''v : volt
x : meter
y : meter
z : meter''')
lfp.x = 7*cm # Off center (to be far from stimulating electrode)
lfp.y = [1*mm, 2*mm, 4*mm, 8*mm, 16*mm]
S = Synapses(neuron,lfp,model='''w : ohm*meter**2 (constant) # Weight in the LFP
\text{calculation}
\text{v_post} = w*(Ic_pre-Im_pre) : volt (summed)''')
S.summed_updaters['v_post'].when = 'after_groups' # otherwise Ic has not yet been updated for the current time step.
S.connect()
S.w = 'area_pre/(4*pi*sigma)/((x_pre-x_post)**2+(y_pre-y_post)**2+(z_pre-z_post)**2)*.5'
Mlfp = StateMonitor(lfp,'v',record=True)
```

```python
run(50*ms, report='text')
nearon.I[0] = 1*uA # current injection at one end
run(3*ms)
nearon.I = 0*amp
run(100*ms, report='text')
```

```python
subplot(211)
for i in range(10):
    plot(M.t/ms,M.v[i*100]/mV)
ylabel('\text{SV}_m$ (mV)')
subplot(212)
for i in range(5):
    plot(M.t/ms,Mlfp.v[i]/mV)
ylabel('LFP (mV)')
```

(continues on next page)
5.10.9 Example: morphotest

Demonstrate the usage of the Morphology object.

```python
from brian2 import *

# Morphology
morpho = Soma(30*um)
morpho.L = Cylinder(diameter=1*um, length=100*um, n=5)
morpho.LL = Cylinder(diameter=1*um, length=20*um, n=2)
morpho.R = Cylinder(diameter=1*um, length=100*um, n=5)

# Passive channels
gL = 1e-4*siemens/cm**2
EL = -70*mV
eqs = ''
Im = gL * (EL-v) : amp/meter**2
```

(continues on next page)
from brian2 import *

defaultclock.dt = 0.01*ms

# Passive channels
gL = 1e-4*siemens/cm**2
EL = -70*mV

# Morphology
diameter = 1*um
length = 300*um
Cm = 1*uF/cm**2
Ri = 150*ohm*cm
N = 500

rm = 1 / (gL * pi * diameter)  # membrane resistance per unit length
ra = (4 * Ri) / (pi * diameter**2)  # axial resistance per unit length
la = sqrt(rm / ra)  # space length
morpho = Cylinder(diameter=diameter, length=length, n=N)
d1 = 0.5*um
L1 = 200*um

rm = 1 / (gL * pi * d1)  # membrane resistance per unit length
ra = (4 * Ri) / (pi * d1**2)  # axial resistance per unit length
l1 = sqrt(rm / ra)  # space length
morpho.L = Cylinder(diameter=d1, length=L1, n=N)
d2 = (diameter**1.5 - d1**1.5)**(1. / 1.5)
rm = 1/(gL * pi * d2)  # membrane resistance per unit length
ra = (4 * Ri) / (pi * d2**2)  # axial resistance per unit length
l2 = sqrt(rm / ra)  # space length
L2 = (L1 / l1) * l2
morpho.R = Cylinder(diameter=d2, length=L2, n=N)

eqs=''
Im = gL * (EL-v) : amp/meter**2
I : amp (point current)
'''

neuron = SpatialNeuron(morphology=morpho, model=eqs, Cm=Cm, Ri=Ri,
                       method='exponential_euler')
neuron.v = EL
neuron.I[0] = 0.02*nA # injecting at the left end
run(100*ms, report='text')

plot(neuron.main.distance/um, neuron.main.v/mV, 'k')
plot(neuron.L.distance/um, neuron.L.v/mV, 'k')
plot(neuron.R.distance/um, neuron.R.v/mV, 'k')
# Theory
x = neuron.main.distance
ra = la * 4 * Ri/(pi * diameter**2)
l = length/la + L1/l1
theory = EL + ra*neuron.I[0]*cosh(l - neuron.main.distance[-1]/la -
       (x - neuron.main.distance[-1])/l1)/sinh(l)
plot(x/um, theory/mV, 'r')
x = neuron.L.distance
theory = (EL+ra*neuron.I[0]*cosh(l - neuron.main.distance[-1]/la -
       (x - neuron.main.distance[-1])/l1)/sinh(l))
plot(x/um, theory/mV, 'r')
x = neuron.R.distance
theory = (EL+ra*neuron.I[0]*cosh(l - neuron.main.distance[-1]/la -
       (x - neuron.main.distance[-1])/l2)/sinh(l))
plot(x/um, theory/mV, 'r')
xlabel('x (um)')
ylabel('v (mV)')
show()
5.10.11 Example: spike_initiation

Ball and stick with Na and K channels

```python
from brian2 import *

defaultclock.dt = 0.025*ms

# Morphology
morpho = Soma(30*um)
morpho.axon = Cylinder(diameter=1*um, length=300*um, n=100)

# Channels
gL = 1e-4*siemens/cm**2
EL = -70*mV
ENa = 50*mV
ka = 6*mV
ki = 6*mV
va = -30*mV
vi = -50*mV
EK = -90*mV
vk = -20*mV
kk = 8*mV
va = -30*mV

eqs = '''
Im = gL*(EL-v)+gNa*m*h*(ENa-v)+gK*n*(EK-v) : amp/meter**2
dm/dt = (minf-m)/(0.3*ms) : 1 # simplified Na channel
dh/dt = (hinf-h)/(3*ms) : 1 # inactivation
dn/dt = (ninf-n)/(5*ms) : 1 # K+
minf = 1/(1+exp((va-v)/ka)) : 1
hinf = 1/(1+exp((v-vi)/ki)) : 1
ninf = 1/(1+exp((vk-v)/kk)) : 1
I : amp (point current)
gNa : siemens/meter**2
gK : siemens/meter**2
'''

neuron = SpatialNeuron(morphology=morpho, model=eqs,
                        Cm=1*uF/cm**2, Ri=100*ohm*cm, method='exponential_euler')

neuron.v = -65*mV
neuron.I = 0*amp
neuron.axon[30*um:60*um].gNa = 700*gL
neuron.axon[30*um:60*um].gK = 700*gL

# Monitors
mon=StateMonitor(neuron, 'v', record=True)

run(1*ms)
neuron.main.I = 0.15*nA
run(50*ms)
neuron.I = 0*amp
run(95*ms, report='text')

plot(mon.t/ms, mon.v[0]/mV, 'r')
plot(mon.t/ms, mon.v[20]/mV, 'g')
plot(mon.t/ms, mon.v[40]/mV, 'b')
plot(mon.t/ms, mon.v[60]/mV, 'k')
```

(continues on next page)
5.11 frompapers

5.11.1 Example: Brette_2004

Phase locking in leaky integrate-and-fire model

Fig. 2A from:


This shows the phase-locking structure of a LIF driven by a sinusoidal current. When the current crosses the threshold ($a<3$), the model almost always phase locks (in a measure-theoretical sense).
from brian2 import *

# defaultclock.dt = 0.01*ms  # for a more precise picture
N = 2000
tau = 100*ms
freq = 1/tau

eqs = '''
dv/dt = (-v + a + 2*sin(2*pi*t/tau))/tau : 1
a : 1
'''

neurons = NeuronGroup(N, eqs, threshold='v>1', reset='v=0', method='euler')
neurons.a = linspace(2, 4, N)

run(5*second, report='text')  # discard the first spikes (wait for convergence)
S = SpikeMonitor(neurons)
run(5*second, report='text')

i, t = S.it
plot((t % tau)/tau, neurons.a[i], ',')
xlabel('Spike phase')
ylabel('Parameter a')
show()
5.11.2 Example: Brette_Gerstner_2005

Adaptive exponential integrate-and-fire model.

http://www.scholarpedia.org/article/Adaptive_exponential_integrate-and-fire_model


```python
from brian2 import *

# Parameters
C = 281 * pF
gL = 30 * nS
tau_m = C / gL
EL = -70.6 * mV
VT = -50.4 * mV
DeltaT = 2 * mV
Vcut = VT + 5 * DeltaT

# Pick an electrophysiological behaviour
tauw, a, b, Vr = 144*ms, 4*nS, 0.0805*nA, -70.6*mV # Regular spiking (as in the paper)
tauw, a, b, Vr = 20*ms, 4*nS, 0.5*nA, VT+5*mV # Bursting
tauw, a, b, Vr = 144*ms, 2*C/(144*ms), 0*nA, -70.6*mV # Fast spiking

eqs = ""
    dvm/dt = (gL*(EL - vm) + gL*DeltaT*exp((vm - VT)/DeltaT) + I - w)/C : volt
    dw/dt = (a*(vm - EL) - w)/tauw : amp
    I : amp
""

neuron = NeuronGroup(1, model=eqs, threshold='vm>Vcut',
                     reset="vm=Vr; w+=b", method='euler')

neuron.vm = EL
trace = StateMonitor(neuron, 'vm', record=0)
spikes = SpikeMonitor(neuron)

run(20 * ms)
neuron.I = 1*nA
run(100 * ms)
neuron.I = 0*nA
run(20 * ms)

# We draw nicer spikes
vm = trace[0].vm[:]
for t in spikes.t:
    i = int(t / defaultclock.dt)
    vm[i] = 20*mV

plot(trace.t / ms, vm / mV)
xlabel('time (ms)')
ylabel('membrane potential (mV)')
show()
```
5.11.3 Example: Brette_Guigon_2003

Reliability of spike timing

Adapted from Fig. 10D.E of

This shows that reliability of spike timing is a generic property of spiking neurons, even those that are not leaky. This is a non-physiological model which can be leaky or anti-leaky depending on the sign of the input $I$.

All neurons receive the same fluctuating input, scaled by a parameter $p$ that varies across neurons. This shows:

1. reproducibility of spike timing
2. robustness with respect to deterministic changes (parameter)
3. increased reproducibility in the fluctuation-driven regime (input crosses the threshold)

```python
from brian2 import *

N = 500

(continues on next page)```
tau = 33*ms
taux = 20*ms
sigma = 0.02

eqs_input = '''
\[\frac{dx}{dt} = \frac{-x}{taux} + \left(\frac{2}{taux}\right)^{0.5}x_i \right] : 1
'''
eqs = '''
\[\frac{dv}{dt} = \frac{(v*I + 1)/tau + sigma\left(\frac{2}{tau}\right)^{0.5}x_i}{1}
I = 0.5 + 3*p*B \right] : 1
B = 2/(1 + \exp(-2*x)) - 1 : 1 \text{(shared)}
p : 1
x : 1 \text{(linked)}
'''

input = NeuronGroup(1, eqs_input, method='euler')
neurons = NeuronGroup(N, eqs, threshold='v>1', reset='v=0', method='euler')
neurons.p = '1.0*i/N'
neurons.v = 'rand()'
neurons.x = linked_var(input, 'x')

M = StateMonitor(neurons, 'B', record=0)
S = SpikeMonitor(neurons)
run(1000*ms, report='text')

subplot(211)
# The input
plot(M.t/ms, M[0].B)
xticks([])
title('shared input')
subplot(212)
plot(S.t/ms, neurons.p[S.i], ',')
plot([0, 1000], [0.5, 0.5], color='C1')
xlabel('time (ms)')
ylabel('p')
title('spiking activity')
show()
5.11.4 Example: Brunel_Hakim_1999

Dynamics of a network of sparsely connected inhibitory current-based integrate-and-fire neurons. Individual neurons fire irregularly at low rate but the network is in an oscillatory global activity regime where neurons are weakly synchronized.


```
from brian2 import *

N = 5000
Vr = 10*mV
theta = 20*mV
tau = 20*ms
delta = 2*ms
tauerefr = 2*ms
duration = .1*second
C = 1000
sparseness = float(C)/N
J = .1*mV
muext = 25*mV
sigmaext = 1*mV
```

(continues on next page)
eqs = ""
   dV/dt = (-V+muext + sigmaext * sqrt(tau) * xi)/tau : volt
   """

group = NeuronGroup(N, eqs, threshold='V>theta',
   reset='V=Vr', refractory=taurefr, method='euler')

group.V = Vr
conn = Synapses(group, group, on_pre='V += -J', delay=delta)
conn.connect(p=sparseness)
M = SpikeMonitor(group)
LFP = PopulationRateMonitor(group)

run(duration)

subplot(211)
plot(M.t/ms, M.i, '.')
xlim(0, duration/ms)

subplot(212)
plot(LFP.t/ms, LFP.smooth_rate(window='flat', width=0.5*ms)/Hz)
xlim(0, duration/ms)

show()
5.11.5 Example: Brunel_Wang_2001

Sample-specific persistent activity

Five subpopulations with three selective and one reset stimuli example. Analog to figure 6b in the paper.


```python
from brian2 import *

# populations
N = 1000
N_E = int(N * 0.8)  # pyramidal neurons
N_I = int(N * 0.2)  # interneurons

# voltage
V_L = -70. * mV
V_thr = -50. * mV
V_reset = -55. * mV
V_E = 0. * mV
V_I = -70. * mV

# membrane capacitance
C_m_E = 0.5 * nF
C_m_I = 0.2 * nF

# membrane leak
g_m_E = 25. * nS
g_m_I = 20. * nS

# refractory period
tau_rp_E = 2. * ms
tau_rp_I = 1. * ms

# external stimuli
rate = 3 * Hz
C_ext = 800

# synapses
C_E = N_E
C_I = N_I

# AMPA (excitatory)
g_AMPA_ext_E = 2.08 * nS
g_AMPA_rec_E = 0.104 * nS * 800. / N_E

# NMDA (excitatory)
g_NMDA_E = 0.327 * nS * 800. / N_E
g_NMDA_I = 0.258 * nS * 800. / N_E
```

(continues on next page)
alpha = 0.5 / ms
Mg2 = 1.

# GABAergic (inhibitory)
g_GABA_E = 1.25 * nS * 200. / N_I
g_GABA_I = 0.973 * nS * 200. / N_I
tau_GABA = 10. * ms

# subpopulations
f = 0.1
p = 5
N_sub = int(N_E * f)
N_non = int(N_E * (1. - f * p))
w_plus = 2.1
w_minus = 1. - f * (w_plus - 1.) / (1. - f)

# modeling
eqs_E = '''
dv / dt = (- g_m_E * (v - V_L) - I_syn) / C_m_E : volt (unless refractory)
I_syn = I_AMPA_ext + I_AMPA_rec + I_NMDA_rec + I_GABA_rec : amp
I_AMPA_ext = g_AMPA_ext_E * (v - V_E) * s_AMPA_ext : amp
I_AMPA_rec = g_AMPA_rec_E * (v - V_E) * 1 * s_AMPA : amp
ds_AMPA_ext / dt = - s_AMPA_ext / tau_AMPA : 1
ds_AMPA / dt = - s_AMPA / tau_AMPA : 1
I_NMDA_rec = g_NMDA_E * (v - V_E) / (1 + Mg2 * exp(-0.062 * v / mV) / 3.57) * s_NMDA_tot : amp
s_NMDA_tot : 1
I_GABA_rec = g_GABA_E * (v - V_I) * s_GABA : amp
ds_GABA / dt = - s_GABA / tau_GABA : 1
'''
eqs_I = '''
dv / dt = (- g_m_I * (v - V_L) - I_syn) / C_m_I : volt (unless refractory)
I_syn = I_AMPA_ext + I_AMPA_rec + I_NMDA_rec + I_GABA_rec : amp
I_AMPA_ext = g_AMPA_ext_I * (v - V_E) * s_AMPA_ext : amp
I_AMPA_rec = g_AMPA_rec_I * (v - V_E) * 1 * s_AMPA : amp
ds_AMPA_ext / dt = - s_AMPA_ext / tau_AMPA : 1
ds_AMPA / dt = - s_AMPA / tau_AMPA : 1
I_NMDA_rec = g_NMDA_I * (v - V_E) / (1 + Mg2 * exp(-0.062 * v / mV) / 3.57) * s_NMDA_tot : amp
s_NMDA_tot : 1
I_GABA_rec = g_GABA_I * (v - V_I) * s_GABA : amp
ds_GABA / dt = - s_GABA / tau_GABA : 1
'''

P_E = NeuronGroup(N_E, eqs_E, threshold='v > V_thr', reset='v = V_reset', refractory=tau_rp_E, method='euler')
P_E.v = V_L
P_I = NeuronGroup(N_I, eqs_I, threshold='v > V_thr', reset='v = V_reset', refractory=tau_rp_I, method='euler')
(continues on next page)
P.I.v = V_L

eqs_glut = '''
s_NMDA_tot_post = w * s_NMDA : 1 (summed)
ds_NMDA / dt = - s_NMDA / tau_NMDA_decay + alpha * x * (1 - s_NMDA) : 1 (clock-driven)
dx / dt = - x / tau_NMDA_rise : 1 (clock-driven)
w : 1
'''
eqs_pre_glut = '''
s_AMPA += w
x += 1
'''
eqs_pre_gaba = '''
s_GABA += 1
'''
eqs_pre_ext = '''
s_AMPA_ext += 1
'''

# E to E
C_E_E = Synapses(P_E, P_E, model=eqs_glut, on_pre=eqs_pre_glut, method='euler')
C_E_E.connect('i != j')
C_E_E.w[:] = 1

for pi in range(N_non, N_non + p * N_sub, N_sub):
    # internal other subpopulation to current nonselective
    C_E_E.w[C_E_E.indices[:, pi:pi + N_sub]] = w_minus

    # internal current subpopulation to current subpopulation
    C_E_E.w[C_E_E.indices[pi:pi + N_sub, pi:pi + N_sub]] = w_plus

# E to I
C_E_I = Synapses(P_E, P_I, model=eqs_glut, on_pre=eqs_pre_glut, method='euler')
C_E_I.connect()
C_E_I.w[:] = 1

# I to I
C_I_I = Synapses(P_I, P_I, on_pre=eqs_pre_gaba, method='euler')
C_I_I.connect('i != j')

# I to E
C_I_E = Synapses(P_I, P_E, on_pre=eqs_pre_gaba, method='euler')
C_I_E.connect()

# external noise
C_P_E = PoissonInput(P_E, 's_AMPA_ext', C_ext, rate, '1')
C_P_I = PoissonInput(P_I, 's_AMPA_ext', C_ext, rate, '1')

# at 1s, select population 1
C_selection = int(f * C_ext)
rate_selection = 25 * Hz
stimuli1 = TimedArray(np.r_[np.zeros(40), np.ones(2), np.zeros(100)], dt=25 * ms)
input1 = PoissonInput(P_E[N_non:N_non + N_sub], 's_AMPA_ext', C_selection, rate_selection, 'stimuli1(t)')
# at 2s, select population 2
stimuli2 = TimedArray(np.r_[np.zeros(80), np.ones(2), np.zeros(100)], dt=25 * ms)
input2 = PoissonInput(P_E[N_non + N_sub:N_non + 2 * N_sub], 's_AMPA_ext', C_selection, 
                   rate_selection, 'stimuli2(t)')

# at 4s, reset selection
stimuli_reset = TimedArray(np.r_[np.zeros(120), np.ones(2), np.zeros(100)], dt=25 * ms)
input_reset_I = PoissonInput(P_E, 's_AMPA_ext', C_ext, rate_selection, 'stimuli_reset(t)')
input_reset_E = PoissonInput(P_I, 's_AMPA_ext', C_ext, rate_selection, 'stimuli_reset(t)')

# monitors
N_activity_plot = 15
sp_E_sels = [SpikeMonitor(P_E[pi:pi + N_activity_plot]) for pi in range(N_non, N_non + 
                  p * N_sub, N_sub)]
sp_E  = SpikeMonitor(P_E[:N_activity_plot])
sp_I  = SpikeMonitor(P_I[:N_activity_plot])

r_E_sels = [PopulationRateMonitor(P_E[pi:pi + N_sub]) for pi in range(N_non, N_non + 
                  p * N_sub, N_sub)]
r_E = PopulationRateMonitor(P_E[:N_non])
r_I = PopulationRateMonitor(P_I)

# simulate, can be long >120s
net = Network(collect())
net.add(sp_E_sels)
net.add(r_E_sels)
net.run(4 * second, report='stdout')

# plotting
title('Population rates')
xlabel('ms')
ylabel('Hz')

plot(r_E.t / ms, r_E.smooth_rate(width=25 * ms) / Hz, label='nonselective')
plot(r_I.t / ms, r_I.smooth_rate(width=25 * ms) / Hz, label='inhibitory')

for i, r_E_sel in enumerate(r_E_sels[::-1]):
    plot(r_E_sel.t / ms, r_E_sel.smooth_rate(width=25 * ms) / Hz, label='selective {}'.format(p - i))

legend()
figure()

title('Population activities ({} neurons/pop)'.format(N_activity_plot))
xlabel('ms')
yticks([])

plot(sp_E.t / ms, sp_E.i + (p + 1) * N_activity_plot, '.', markersize=2, label='nonselective')
plot(sp_I.t / ms, sp_I.i + p * N_activity_plot, '.', markersize=2, label='inhibitory')

for i, sp_E_sel in enumerate(sp_E_sels[::-1]):
    plot(sp_E_sel.t / ms, sp_E_sel.i + (p - i - 1) * N_activity_plot, '.', 
          markersize=2, label='selective {}'.format(p - i))
legend()

show()
5.11.6 Example: Clopath_et_al_2010_homeostasis

This code contains an adapted version of the voltage-dependent triplet STDP rule from: Clopath et al., Connectivity reflects coding: a model of voltage-based STDP with homeostasis, Nature Neuroscience, 2010 (http://dx.doi.org/10.1038/nn.2479)

The plasticity rule is adapted for a leaky integrate & fire model in Brian2. More specifically, the filters v_lowpass1 and v_lowpass2 are incremented by a constant at every post-synaptic spike time, to compensate for the lack of an actual spike in the integrate & fire model.

As an illustration of the rule, we simulate the competition between inputs projecting on a downstream neuron. We would like to note that the parameters have been chosen arbitrarily to qualitatively reproduce the behavior of the original work, but need additional fitting.

We kindly ask to cite the article when using the model presented below.

This code was written by Jacopo Bono, 12/2015

```python
from brian2 import *

###############################################
# PLASTICITY MODEL
###############################################
```
### Plasticity Parameters

- $V_{\text{rest}} = -70.\text{mV}$  
  
  # resting potential

- $V_{\text{thresh}} = -55.\text{mV}$  
  
  # spiking threshold

- $\text{Theta}_\text{low} = V_{\text{rest}}$  
  
  # depolarization threshold for plasticity

- $x_{\text{reset}} = 1.$  
  
  # spike trace reset value

- $\tau_{x} = 15.\text{ms}$  
  
  # spike trace time constant

- $A_{\text{LTD}} = 1.5e^{-4}$  
  
  # depression amplitude

- $A_{\text{LTP}} = 1.5e^{-2}$  
  
  # potentiation amplitude

- $\tau_{\text{lowpass1}} = 40.\text{ms}$  
  
  # timeconstant for low-pass filtered voltage

- $\tau_{\text{lowpass2}} = 30.\text{ms}$  
  
  # timeconstant for low-pass filtered voltage

- $v_{\text{target}} = 12.\text{mV}^2$  
  
  # target depolarisation

### Plasticity Equations

- Equations executed at every timestep

  
  ```
  \text{Syn\_model} = {'''
  \text{w}\_\text{ampa}:1  
  \hspace{1cm} \# synaptic weight (ampa synapse)
  
  '''
  ```

- Equations executed only when a presynaptic spike occurs

  ```
  \text{Pre\_eq} = {'''
  \hspace{1cm} g_{\text{ampa\_post}} += w_{\text{ampa}}*\text{ampa\_max\_cond}  
  \hspace{1cm} \# increment synaptic conductance
  
  \hspace{1cm} A_{\text{LTD\_u}} = A_{\text{LTD}}*(v_{\text{homeo}}**2/v_{\text{target}})  
  \hspace{1cm} \# metaplasticity
  
  \hspace{1cm} w_{\text{minus}} = A_{\text{LTD\_u}}*(v_{\text{lowpass1\_post}}/\text{mV} - \text{Theta}_\text{low}/\text{mV})*\text{int}(v_{\text{lowpass1\_post}}/\text{mV} - \text{Theta}_\text{low}/\text{mV} > 0)  
  \hspace{1cm} \# synaptic depression
  
  \hspace{1cm} w_{\text{ampa}} = \text{clip}(w_{\text{ampa}}-w_{\text{minus}}, 0, w_{\text{max}})  
  \hspace{1cm} \# hard bounds
  
  '''
  ```

- Equations executed only when a postsynaptic spike occurs

  ```
  \text{Post\_eq} = {'''
  \hspace{1cm} v_{\text{lowpass1}} += 10.\text{mV}  
  \hspace{1cm} \# mimics the depolarisation effect due to a spike
  
  \hspace{1cm} v_{\text{lowpass2}} += 10.\text{mV}  
  \hspace{1cm} \# mimics the depolarisation effect due to a spike
  
  \hspace{1cm} v_{\text{homeo}} += 0.1.\text{mV}  
  \hspace{1cm} \# mimics the depolarisation effect due to a spike
  
  \hspace{1cm} w_{\text{plus}} = A_{\text{LTP}}*x_{\text{trace\_pre}}*(v_{\text{lowpass2\_post}}/\text{mV} - \text{Theta}_\text{low}/\text{mV})*\text{int}(v_{\text{lowpass2\_post}}/\text{mV} - \text{Theta}_\text{low}/\text{mV} > 0)  
  \hspace{1cm} \# synaptic potentiation
  
  \hspace{1cm} w_{\text{ampa}} = \text{clip}(w_{\text{ampa}}+w_{\text{plus}}, 0, w_{\text{max}})  
  \hspace{1cm} \# hard bounds
  
  '''
  ```

### Neuron parameters

- $g_{\text{leak}} = 30.\text{nS}$  
  
  # leak conductance

- $C = 300.\text{pF}$  
  
  # membrane capacitance

- $\tau_{\text{AMPA}} = 2.\text{ms}$  
  
  # AMPA synaptic timeconstant
E_AMPA = 0.*mV  # reversal potential AMPA
ampa_max_cond = 5.e-8*siemens  # Ampa maximal conductance
w_max = 1.  # maximal ampa weight

### Neuron Equations

# We connect 10 presynaptic neurons to 1 downstream neuron

dv/dt = (gleak*(V_rest-v) + I_ext + I_syn)/C: volt  # voltage

# downstream neuron
eqs_neurons = '''

# input neurons
eqs_inputs = '''

dv/dt = gleak*(V_rest-v)/C: volt  # voltage

# Simulation

#### Parameters

defaultclock.dt = 500.*us  # timestep
Nr_neurons = 1  # Number of downstream neurons
Nr_inputs = 5  # Number of input neurons
input_rate = 35*Hz  # Rates
init_weight = 0.5  # initial synaptic weight
final_t = 20.*second  # end of simulation
input_time = 100.*ms  # duration of an input

### Create neuron objects

Nrn_downstream = NeuronGroup(Nr_neurons, eqs_neurons, threshold='v>V_thresh',
                           reset='v=V_rest;x_trace+=x_reset/(taux/ms)',
                           method='euler')

Nrn_input = NeuronGroup(Nr_inputs, eqs_inputs, threshold='rand()<rates*dt',
                        reset='v=V_rest;x_trace+=x_reset/(taux/ms)',
                        method='exact')

### create Synapses
Syn = Synapses(Nrns_input, Nrn_downstream,
    model=Syn_model,
    on_pre=Pre_eq,
    on_post=Post_eq
)

Syn.connect(i=numpy.arange(Nr_inputs), j=0)

### Monitors and storage
W_evolution = StateMonitor(Syn, 'w_ampa', record=True)

### Run
# Initial values
Nrn_downstream.v = V_rest
Nrn_downstream.v_lowpass1 = V_rest
Nrn_downstream.v_lowpass2 = V_rest
Nrn_downstream.v_homeo = 0
Nrn_downstream.I_ext = 0.*amp
Nrn_downstream.x_trace = 0.
Nrns_input.v = V_rest
Nrns_input.x_trace = 0.
Syn.w_ampa = init_weight

# Switch on a different input every 100ms
Nrns_input.run_regularly('''
    selected_index = int(floor(rand()*Nr_inputs))
    rates = input_rate * int(selected_index == i) # All rates are zero except for the selected neuron
    ''', dt=input_time)
run(final_t, report='text')

# Plots
stitle = 'Synaptic Competition'
fig = figure(figsize=(8, 5))
for kk in range(Nr_inputs):
    plt.plot(W_evolution.t, W_evolution.w_ampa[kk], '-', linewidth=2)
xlabel('Time [ms]', fontsize=22)
ylabel('Weight [a.u.]', fontsize=22)
plt.subplots_adjust(bottom=0.2, left=0.15, right=0.95, top=0.85)
title(stitle, fontsize=22)
plt.show()
5.11.7 Example: Clopath_et_al_2010_no_homeostasis

This code contains an adapted version of the voltage-dependent triplet STDP rule from: Clopath et al., Connectivity reflects coding: a model of voltage-based STDP with homeostasis, Nature Neuroscience, 2010 (http://dx.doi.org/10.1038/nn.2479)

The plasticity rule is adapted for a leaky integrate & fire model in Brian2. In particular, the filters $v_{\text{lowpass1}}$ and $v_{\text{lowpass2}}$ are incremented by a constant at every post-synaptic spike time, to compensate for the lack of an actual spike in the integrate & fire model. Moreover, this script does not include the homeostatic metaplasticity.

As an illustration of the Rule, we simulate a plot analogous to figure 2b in the above article, showing the frequency dependence of plasticity as measured in: Sjöström et al., Rate, timing and cooperativity jointly determine cortical synaptic plasticity. Neuron, 2001. We would like to note that the parameters have been chosen arbitrarily to qualitatively reproduce the behavior of the original works, but need additional fitting.

We kindly ask to cite both articles when using the model presented below.

This code was written by Jacopo Bono, 12/2015

```python
from brian2 import *

# PLASTICITY MODEL

#### Plasticity Parameters

V_rest = -70.*mV # resting potential
V_thresh = -50.*mV # spiking threshold
```

(continues on next page)
Theta_low = V_rest # depolarization threshold for plasticity
x_reset = 1. # spike trace reset value
tau = 15.*ms # spike trace time constant
A_LTD = 1.5e-4 # depression amplitude
A_LTP = 1.5e-2 # potentiation amplitude
tau_lowpass1 = 40.*ms # timeconstant for low-pass filtered voltage
tau_lowpass2 = 30.*ms # timeconstant for low-pass filtered voltage

#### Plasticity Equations

# equations executed at every timestep
Syn_model =

w_ampa:1 # synaptic weight (ampa synapse)

# equations executed only when a presynaptic spike occurs
Pre_eq =

g_ampa_post += w_ampa*ampa_max_cond # increment synaptic conductance
w_minus = A_LTD*(v_lowpass1_post/mV - Theta_low/mV)*int(v_lowpass1_post/mV -
Theta_low/mV > 0) # synaptic depression
w_ampa = clip(w_ampa-w_minus,0,w_max) # hard bounds

# equations executed only when a postsynaptic spike occurs
Post_eq =

v_lowpass1 += 10*mV # mimics the depolarisation by a spike
v_lowpass2 += 10*mV # mimics the depolarisation by a spike
w_plus = A_LTP*x_trace_pre*(v_lowpass2_post/mV - Theta_low/mV)*int(v_
lowpass2_post/mV - Theta_low/mV > 0) # synaptic potentiation
w_ampa = clip(w_ampa+w_plus,0,w_max) # hard bounds

# I&F Parameters and equations

### Neuron parameters

gleak = 30.*nS # leak conductance
C = 300.*pF # membrane capacitance
tau_AMPA = 2.*ms # AMPA synaptic timeconstant
E_AMPA = 0.*mV # reversal potential AMPA
ampa_max_cond = 5.e-10*siemens # Ampa maximal conductance
w_max = 1. # maximal ampa weight

### Neuron Equations
eqs_neurons = '''
\begin{align*}
\frac{dv}{dt} &= (g_{\text{leak}}(V_{\text{rest}}-v) + I_{\text{ext}} + I_{\text{syn}})/C: \text{volt} \quad \# \text{voltage} \\
\frac{dv_{\text{lowpass1}}}{dt} &= (v-v_{\text{lowpass1}})/\tau_{\text{lowpass1}}: \text{volt} \quad \# \text{low-pass filter of the voltage} \\
\frac{dv_{\text{lowpass2}}}{dt} &= (v-v_{\text{lowpass2}})/\tau_{\text{lowpass2}}: \text{volt} \quad \# \text{low-pass filter of the voltage} \\
I_{\text{ext}} : \text{amp} \quad \# \text{external current} \\
I_{\text{syn}} &= g_{\text{ampa}}(E_{\text{AMPA}}-v): \text{amp} \quad \# \text{synaptic current} \\
g_{\text{ampa}}/dt &= -g_{\text{ampa}}/\tau_{\text{AMPA}}: \text{siemens} \quad \# \text{synaptic conductance} \\
\frac{dx_{\text{trace}}}{dt} &= -x_{\text{trace}}/\tau_{\text{aux}}:1 \quad \# \text{spike trace}
\end{align*}
'''

'''

# Simulation

### Parameters

defaultclock.dt = 100.*\mu\text{sec} \quad \# \text{timestep}

Nr_neurons = 2 \quad \# \text{Number of neurons}

rate_array = [1., 5., 10., 15., 20., 30., 50.]*Hz \quad \# \text{Rates}

init_weight = 0.5 \quad \# \text{initial synaptic weight}

reps = 15 \quad \# \text{Number of pairings}

### Create neuron objects

Nrns = NeuronGroup(Nr_neurons, eqs_neurons, threshold='v>V_{\text{thresh}}', reset='v=V_{\text{rest}}; x_{\text{trace}}+=x_{\text{reset}}/(\tau_{\text{aux}}/\text{ms})', method='euler')

#### create Synapses

Syn = Synapses(Nrns, Nrns, model=Syn_model, on_pre=Pre_eq, on_post=Post_eq)

Syn.connect('i!=j')

#### Monitors and storage

weight_result = np.zeros((2,len(rate_array))) \quad \# \text{to save the final weights}

#### Run

for jj, rate in enumerate(rate_array):

    # Calculate interval between pairs
    pair_interval = 1./rate - 10*\text{ms}
    print('Starting simulations for \$$.\text{rate}$$ \% rate)

    # Initial values
    Nrns.v = V_{\text{rest}}
    Nrns.v_{\text{lowpass1}} = V_{\text{rest}}

(continues on next page)
Nrns.v_lowpass2 = V_rest
Nrns.I_ext = 0*amp
Nrns.x_trace = 0.
Syn.w_ampa = init_weight

# loop over pairings
for ii in range(reps):
    # 1st SPIKE
    Nrns.v[0] = V_thresh + 1*mV
    # 2nd SPIKE
    run(10*ms)
    Nrns.v[1] = V_thresh + 1*mV
    # run
    run(pair_interval)
    print('Pair %d out of %d' % (ii+1, reps))

#store weight changes
weight_result[0, jj] = 100.*Syn.w_ampa[0]/init_weight
weight_result[1, jj] = 100.*Syn.w_ampa[1]/init_weight

# Plots
figure(figsize=(8, 5))
plot(rate_array, weight_result[0, :], '-', linewidth=2, color=scolor)
plot(rate_array, weight_result[1, :], ':', linewidth=2, color=scolor)
xlabel('Pairing frequency [Hz]', fontsize=22)
ylabel('Normalised Weight [%]', fontsize=22)
legend(["Pre-Post", 'Post-Pre'], loc='best')
subplots_adjust(bottom=0.2, left=0.15, right=0.95, top=0.85)
title(stitle)
show()
5.11.8 Example: Destexhe_et_al_1998

Reproduces Figure 12 (simplified three-compartment model) from the following paper:
Dendritic Low-Threshold Calcium Currents in Thalamic Relay Cells Alain Destexhe, Mike Neubig, Daniel Ulrich, John Huguenard Journal of Neuroscience 15 May 1998, 18 (10) 3574-3588

The original NEURON code is available on ModelDB: https://senselab.med.yale.edu/modeldb/ShowModel.cshtml?model=279

Reference for the original morphology:

Available at NeuroMorpho.org:

Notes

- Completely removed the “Fast mechanism for submembranal Ca++ concentration (cai)” – it did not affect the results presented here
- Time constants for the I_T current are slightly different from the equations given in the paper – the paper calculation seems to be based on 36 degree Celsius but the temperature that is used is 34 degrees.
• To reproduce Figure 12C, the “presence of dendritic shunt conductances” meant setting $g_L$ to 0.15 mS/cm$^2$ for the whole neuron.

• Other small discrepancies with the paper – values from the NEURON code were used whenever different from the values stated in the paper.

```python
from __future__ import print_function
from brian2 import *
from brian2.units.constants import (zero_celsius, faraday_constant as F,
                                     gas_constant as R)

defaultclock.dt = 0.01*ms

VT = -52*mV
El = -76.5*mV  # from code, text says: -69.85*mV
E_Na = 50*mV
E_K = -100*mV
C_d = 7.954  # dendritic correction factor

T = 34*kelvin + zero_celsius  # 34 degC (current-clamp experiments)
tadj_HH = 3.0**((34-36)/10.0)  # temperature adjustment for Na & K (original recordings at 36 degC)
tadj_m_T = 2.5**((34-24)/10.0)
tadj_h_T = 2.5**((34-24)/10.0)
shift_I_T = -1*mV

gamma = F/(R*T)  # R=gas constant, F=Faraday constant
Z_CA = 2      # Valence of Calcium ions
Ca_i = 240*nM  # intracellular Calcium concentration
Ca_o = 2*mM   # extracellular Calcium concentration

eqs = Equations(''
Im = gl*(El-v) - I_Na - I_K - I_T: amp/meter**2
I_inj : amp (point current)
gl : siemens/meter**2

# HH-type currents for spike initiation
g_Na : siemens/meter**2
g_K : siemens/meter**2
I_Na = g_Na * m**3 * h * (v-E_Na) : amp/meter**2
I_K = g_K * n**4 * (v-E_K) : amp/meter**2
v2 = v - VT : volt  # shifted membrane potential (Traub convention)

# Low-threshold Calcium current (I_T) -- nonlinear function of voltage
I_T = P_Ca * m_I*T**2*h_T * G_Ca : amp/meter**2
P_Ca = meter/second  # maximum Permeability to Calcium
G_Ca = Z_Ca**2*F*v*gamma*(Ca_i - Ca_o*exp(-Z_Ca*gamma*v))/(1 - exp(-Z_Ca*gamma*v)) : coulomb/meter**3
```

(continues on next page)
\[
\begin{align*}
\frac{dm_T}{dt} &= -(m_T - m_T_{\infty})/\tau_m_T : 1 \\
\frac{dh_T}{dt} &= -(h_T - h_T_{\infty})/\tau_h_T : 1 \\
m_T_{\infty} &= \frac{1}{1 + \exp(-{(v/mV + 56)/6.2})} : 1 \\
h_T_{\infty} &= \frac{1}{1 + \exp((v/mV + 15.8)/18.2))} : 1 \\
\tau_m_T &= (0.612 + 1.0/(\exp(-(v/mV + 131)/16.7) + \exp((v/mV + 15.8)/18.2))) \times ms / tadj_m_T: second \\
\tau_h_T &= (int(v<-81*mV) \times \exp((v/mV + 466)/66.6) + int(v>=-81*mV) \times (28 + \exp(-(v/mV + 21)/10.5))) \times ms / tadj_h_T: second \\
\end{align*}
\]

```python
# Simplified three-compartment morphology
morpho = Cylinder(x=[0, 38.42]*um, diameter=26*um)
morpho.dend = Cylinder(x=[0, 12.49]*um, diameter=10.28*um)
morpho.dend.distal = Cylinder(x=[0, 84.67]*um, diameter=8.5*um)
neuron = SpatialNeuron(morpho, eqs, Cm=0.88*uF/cm**2, Ri=173*ohm*cm,
                      method='exponential_euler')
neuron.v = -74*mV
# Only the soma has Na/K channels
neuron.main.g_Na = 100*msiemens/cm**2
neuron.main.g_K = 100*msiemens/cm**2
# Apply the correction factor to the dendrites
neuron.dend.Cm *= C_d
neuron.m_T = 'm_T_{\infty}'
neuron.h_T = 'h_T_{\infty}'
mon = StateMonitor(neuron, ['v'], record=True)
store('initial state')

def do_experiment(currents, somatic_density, dendritic_density,
                   dendritic_conductance=0.0379*msiemens/cm**2,
                   HH_currents=True):
    restore('initial state')
    voltages = []
    neuron.P_Ca = somatic_density
    neuron.dend.distal.P_Ca = dendritic_density * C_d
    # dendritic conductance (shunting conductance used for Fig 12C)
    neuron.gl = dendritic_conductance
    neuron.dend.gl = dendritic_conductance * C_d
    if not HH_currents:
        # Shut off spiking (for Figures 12B and 12C)
        neuron.g_Na = 0*msiemens/cm**2
        neuron.g_K = 0*msiemens/cm**2
    run(180*ms)
    store('before current')
    for current in currents:
        restore('before current')
        neuron.main.I_inj = current
        print('...', end='')
        run(320*ms)
        voltages.append(mon[morpho].v[:])  # somatic voltage
    return voltages
```

(continues on next page)
## Run the various variants of the model to reproduce Figure 12

```python
mpl.rcParams['lines.markersize'] = 3.0
fig, axes = plt.subplots(2, 2)

print('Running experiments for Figure A1 ', end='')
voltages = do_experiment([50, 75]*pA, somatic_density=1.7e-5*cm/second, dendritic_density=1.7e-5*cm/second)
print(' done. ')
cut_off = 100*ms  # Do not display first part of simulation
axes[0, 0].plot((mon.t - cut_off) / ms, voltages[0] / mV, color='gray')
axes[0, 0].plot((mon.t - cut_off) / ms, voltages[1] / mV, color='black')
axes[0, 0].set(xlim=(0, 400), ylim=(-80, 40), xticks=[], title='A1: Uniform T-current density', ylabel='Voltage (mV)')
axes[0, 0].spines['right'].set_visible(False)
axes[0, 0].spines['top'].set_visible(False)

print('Running experiments for Figure A2 ', end='')
voltages = do_experiment([50, 75]*pA, somatic_density=1.7e-5*cm/second, dendritic_density=9.5e-5*cm/second)
print(' done. ')
cut_off = 100*ms  # Do not display first part of simulation
axes[1, 0].plot((mon.t - cut_off) / ms, voltages[0] / mV, color='gray')
axes[1, 0].plot((mon.t - cut_off) / ms, voltages[1] / mV, color='black')
axes[1, 0].set(xlim=(0, 400), ylim=(-80, 40), xticks=[], title='A2: High T-current density in dendrites', xlabel='Time (ms)', ylabel='Voltage (mV)')
axes[1, 0].spines['right'].set_visible(False)
axes[1, 0].spines['top'].set_visible(False)

print('Running experiments for Figure B ', end='')
currents = np.linspace(0, 200, 41)*pA
voltages_somatic = do_experiment(currents, somatic_density=56.36e-5*cm/second, dendritic_density=0*cm/second, HH_currents=False)
voltages_somatic_dendritic = do_experiment(currents, somatic_density=1.7e-5*cm/second, dendritic_density=9.5e-5*cm/second, HH_currents=False)
print(' done. ')
maxima_somatic = Quantity(voltages_somatic).max(axis=1)
maxima_somatic_dendritic = Quantity(voltages_somatic_dendritic).max(axis=1)
axes[0, 1].yaxis.tick_right()
axes[0, 1].plot(currents/pA, maxima_somatic/mV, 'o-', color='black', label='Somatic only')
axes[0, 1].plot(currents/pA, maxima_somatic_dendritic/mV, 's-', color='black', label='Somatic & dendritic')
axes[0, 1].set(xlabel='Injected current (pA)', ylabel='Peak LTS (mV)', ylim=(-80, 0))
axes[0, 1].legend(loc='best', frameon=False)

print('Running experiments for Figure C ', end='')
currents = np.linspace(200, 400, 41)*pA
voltages_somatic = do_experiment(currents, somatic_density=56.36e-5*cm/second, dendritic_density=0*cm/second, dendritic_conductance=0.15*msiemens/cm**2, HH_currents=False)
voltages_somatic_dendritic = do_experiment(currents, somatic_density=1.7e-5*cm/second, dendritic_density=9.5e-5*cm/second, HH_currents=False)
```

(continues on next page)
dendritic_conductance=0.15*msiemens/cm**2,
HH_currents=False)
print(' done.')
maxima_somatic = Quantity(voltages_somatic).max(axis=1)
maxima_somatic_dendritic = Quantity(voltages_somatic_dendritic).max(axis=1)
axes[1, 1].yaxis.tick_right()
axes[1, 1].plot(currents/pA, maxima_somatic/mV,
    'o-', color='black', label='Somatic only')
axes[1, 1].plot(currents/pA, maxima_somatic_dendritic/mV,
    's-', color='black', label='Somatic & dendritic')
axes[1, 1].set(xlabel='Injected current (pA)', ylabel='Peak LTS (mV)',
    ylim=(-80, 0))
axes[1, 1].legend(loc='best', frameon=False)
plt.tight_layout()
plt.show()
Synfire chains


```python
from brian2 import *

duration = 100*ms

# Neuron model parameters
Vr = -70*mV
Vt = -55*mV
taum = 10*ms
tauvsp = 0.325*ms
weight = 4.86*mV

# Neuron model
eqs = Equations(''
    dV/dt = -(V-Vr)+x*(1./taum) : volt
    dx/dt = (-x+y)*(1./taupsp) : volt
    dy/dt = -y*(1./taupsp)+25.27*mV/ms+(39.24*mV/ms**0.5)*xi : volt''
)

# Neuron groups
n_groups = 10
group_size = 100
P = NeuronGroup(N=n_groups*group_size, model=eqs,
    threshold='V>Vt', reset='V=Vr', refractory=1*ms,
    method='euler')

Pinput = SpikeGeneratorGroup(85, np.arange(85),
    np.random.randn(85)*1*ms + 50*ms)

# The network structure
S = Synapses(P, P, on_pre='y+=weight')
S.connect(j='k for k in range((int(i/group_size)+1)*group_size, (int(i/group_size)+2)*group_size) 'if i<N_pre-group_size')
Sinput = Synapses(Pinput, P[:group_size], on_pre='y+=weight')
Sinput.connect()

# Record the spikes
Mgp = SpikeMonitor(P)
Minput = SpikeMonitor(Pinput)

# Setup the network, and run it
P.V = 'Vr + rand() *(Vt - Vr)'
run(duration)

plot(Mgp.t/ms, 1.0*Mgp.i/group_size, '.')
plot([0, duration/ms], np.arange(n_groups).repeat(2).reshape(-1, 2).T, 'k-')
ylabel('group number')
yticks(np.arange(n_groups))
xlabel('time (ms)')
show()
```
5.11.10 Example: Hindmarsh_Rose_1984

Burst generation in the Hindmarsh-Rose model. Reproduces Figure 6 of:


```python
from brian2 import *

defualtclock.dt = time_unit/10

x_1 = -1.6  # leftmost equilibrium point of the model without adaptation
a = 1; b = 3; c = 1; d = 5
r = 0.001; s = 4

eqs = ''

dx/dt = (y - a*x**3 + b*x**2 + I - z)/time_unit : 1
dy/dt = (c - d*x**2 - y)/time_unit : 1
dz/dt = r*(s*(x - x_1) - z)/time_unit : 1
I : 1 (constant)
```

(continues on next page)
# We run the model with three different currents
neuron = NeuronGroup(3, eqs, method='rk4')

# Set all variables to their equilibrium point
neuron.x = x_1
neuron.y = 'c - d*x**2'
neuron.z = 'r*(s*(x - x_1))'

# Set the constant current input
neuron.I = [0.4, 2, 4]

# Record the "membrane potential"
mon = StateMonitor(neuron, 'x', record=True)

run(2100*time_unit)

ax_top = plt.subplot2grid((2, 3), (0, 0), colspan=3)
ax_bottom_l = plt.subplot2grid((2, 3), (1, 0), colspan=2)
ax_bottom_r = plt.subplot2grid((2, 3), (1, 2))

for ax in [ax_top, ax_bottom_l, ax_bottom_r]:
    ax.spines['top'].set_visible(False)
    ax.spines['right'].set_visible(False)
    ax.set(ylim=(-2, 2), yticks=[-2, 0, 2])

ax_top.plot(mon.t/time_unit, mon.x[0])
ax_bottom_l.plot(mon.t/time_unit, mon.x[1])
ax_bottom_l.set_xlim(700, 2100)

ax_bottom_r.plot(mon.t/time_unit, mon.x[2])
ax_bottom_r.set_xlim(1400, 2100)
ax_bottom_r.set_yticks([])

plt.show()
5.11.11 Example: Izhikevich_2007

STDP modulated with reward
Adapted from Fig. 1c of:


Note: The variable “mode” can switch the behavior of the synapse from “Classical STDP” to “Dopamine modulated STDP”.

Author: Guillaume Dumas (Institut Pasteur) Date: 2018-08-24

```python
from brian2 import *

# Parameters
simulation_duration = 6 * second

## Neurons
taum = 10*ms
Ee = 0*mV
vt = -54*mV
vr = -60*mV
```

(continues on next page)
\[ E_l = -74\, \text{mV} \]
\[ \tau_{ae} = 5\, \text{ms} \]

### STDP

\[ \tau_{a\text{pre}} = 20\, \text{ms} \]
\[ \tau_{a\text{post}} = \tau_{a\text{pre}} \]
\[ g_{\text{max}} = 0.01 \]
\[ dA_{\text{pre}} = 0.01 \]
\[ dA_{\text{post}} = -dA_{\text{pre}} \cdot \tau_{a\text{pre}} / \tau_{a\text{post}} \cdot 1.05 \]
\[ dA_{\text{post}} *= g_{\text{max}} \]
\[ dA_{\text{pre}} *= g_{\text{max}} \]

### Dopamine signaling

\[ \tau_{uc} = 1000\, \text{ms} \]
\[ \tau_{ud} = 200\, \text{ms} \]
\[ \tau_{us} = 1\, \text{ms} \]
\[ \epsilon_{\text{dopa}} = 5\times10^{-3} \]

# Setting the stage

### Stimuli section

\[ \text{input\_indices} = \text{array([0, 1, 0, 1, 1, 0,} \]
\[ 0, 1, 0, 1, 1, 0]) \]
\[ \text{input\_times} = \text{array([500, 550, 1000, 1010, 1500, 1510,} \]
\[ 3500, 3550, 4000, 4010, 4500, 4510])\, \text{ms} \]
\[ \text{input} = \text{SpikeGeneratorGroup}(2, \text{input\_indices}, \text{input\_times}) \]

\[ \text{neurons} = \text{NeuronGroup}(2, \text{'''dv/dt = (ge * (Ee-v) + El - v) / taum : volt} \]
\[ \text{dge/dt = -ge / taue : 1''}',} \]
\[ \text{threshold='v>vt', reset='v = vr',} \]
\[ \text{method='exact')} \]

\[ \text{neurons.v} = \text{vr} \]
\[ \text{neurons\_monitor} = \text{SpikeMonitor(neurons)} \]

\[ \text{synapse} = \text{Synapses(input, neurons,} \]
\[ \text{model='''s: volt''',} \]
\[ \text{on\_pre='v += s')} \]
\[ \text{synapse.connect(i=[0, 1], j=[0, 1])} \]
\[ \text{synapse.s} = 100. \times \text{mV} \]

### STDP section

\[ \text{synapse\_stdp} = \text{Synapses(neurons, neurons,} \]
\[ \text{model='''mode: 1} \]
\[ \text{dc/dt = -c / tauc : 1 (clock-driven)} \]
\[ \text{dd/dt = -d / taud : 1 (clock-driven)} \]
\[ \text{ds/dt = mode * c * d / taus : 1 (clock-driven)} \]
\[ \text{dApre/dt = -Apre / taupre : 1 (event-driven)} \]
\[ \text{dApost/dt = -Apost / taupost : 1 (event-driven)}''',} \]
\[ \text{on\_pre='''ge += s} \]
\[ \text{Apre += dApre} \]
\[ \text{c = clip(c + mode * Apost, -gmax, gmax)} \]
\[ \text{s = clip(s + (1-mode) * Apost, -gmax, gmax)} \]
\[ '''\text{,''} \]
\[ \text{on\_post='''Apost += dApost} \]
\[ \text{c = clip(c + mode * Apre, -gmax, gmax)} \]
\[ \text{s = clip(s + (1-mode) * Apre, -gmax, gmax)} \]
\[ '''\text{,''} \]
method='euler'
)
synapse_stdp.connect(i=0, j=1)
synapse_stdp.mode = 0
synapse_stdp.s = 1e-10
synapse_stdp.c = 1e-10
synapse_stdp.d = 0
synapse_stdp_monitor = StateMonitor(synapse_stdp, ['s', 'c', 'd'], record=[0])

## Dopamine signaling section

dopamine_indices = array([0, 0, 0])
dopamine_times = array([3520, 4020, 4520])*ms
dopamine = SpikeGeneratorGroup(1, dopamine_indices, dopamine_times)
dopamine_monitor = SpikeMonitor(dopamine)
reward = Synapses(dopamine, synapse_stdp, model='''
on_pre='''d_post += epsilon_dopa''',
method='exact')
reward.connect()

# Simulation

## Classical STDP

run(simulation_duration/2)

## Dopamine modulated STDP

run(simulation_duration/2)

# Visualisation

dopamine_indices, dopamine_times = dopamine_monitor.it
neurons_indices, neurons_times = neurons_monitor.it
figure(figsize=(12,6))
subplot(411)
plot([0.05, 2.95], [2.7, 2.7], linewidth=5, color='k')
text(1.5, 3, 'Classical STDP', horizontalalignment='center', fontsize=20)
plot([3.05, 5.95], [2.7, 2.7], linewidth=5, color='k')
text(4.5, 3, 'Dopamine modulated STDP', horizontalalignment='center', fontsize=20)
plot(neurons_times, neurons_indices, 'ob')
plot(dopamine_times, dopamine_indices + 2, 'or')
xlim([0, simulation_duration/second])
ylim([-0.5, 4])
yticks([0, 1, 2], ['Pre-neuron', 'Post-neuron', 'Reward'])
xticks([])

subplot(412)
plot(synapse_stdp_monitor.t/second, synapse_stdp_monitor.d.T/gmax, 'r-')
xlim([0, simulation_duration/second])
ylabel('Extracellular dopamine d(t)')
xticks([])

subplot(413)
plot(synapse_stdp_monitor.t/second, synapse_stdp_monitor.c.T/gmax, 'b-')
xlim([0, simulation_duration/second])
ylabel('Eligibility trace c(t)')
xticks([])

subplot(414)
plot(synapse_stdp_monitor.t/second, synapse_stdp_monitor.s.T/gmax, 'g-')
xlim([0, simulation_duration/second])
ylabel('Synaptic strength s(t)')
xlabel('Time (s)')

(continues on next page)
5.11.12 Example: Kremer_et_al_2011_barrel_cortex


Development of direction maps with pinwheels in the barrel cortex. Whiskers are deflected with random moving bars. N.B.: network construction can be long.

```python
from brian2 import *
import time
t1 = time.time()

# PARAMETERS
# Neuron numbers
M4, M23exc, M23inh = 22, 25, 12  # size of each barrel (in neurons)
N4, N23exc, N23inh = M4**2, M23exc**2, M23inh**2  # neurons per barrel
barrelarraysize = 5  # Choose 3 or 4 if memory error
Nbarrels = barrelarraysize**2

# Stimulation
stim_change_time = 5*ms
Fmax = .5/stim_change_time  # maximum firing rate in layer 4 (.5 spike / stimulation)

# Neuron parameters
tau_m, tau_e, tau_i = 10*ms, 2*ms, 25*ms
E_l = -70*mV
Vt, vt_inc, tau_vt = -55*mV, 2*mV, 50*ms  # adaptive threshold

# STDP
taup, taud = 5*ms, 25*ms
```

(continues on next page)
Ap, Ad = .05, -.04

# EPSPs/IPSPs
EPSP, IPSP = 1*mV, -1*mV
EPSC = EPSP * (taue/taum)**(taum/(taue-taum))
IPSC = IPSP * (taui/taum)**(taum/(taui-taum))
Ap, Ad = Ap*EPSC, Ad*EPSC

# Layer 4, models the input stimulus
eqs_layer4 =
'''
rate = int(is_active)*clip(cos(direction - selectivity), 0, inf)*Fmax: Hz
is_active = abs((barrel_x + 0.5 - bar_x) * cos(direction) + (barrel_y + 0.5 - bar_y) -
    sin(direction)) < 0.5: boolean
barrel_x : integer # The x index of the barrel
barrel_y : integer # The y index of the barrel
selectivity : 1
# Stimulus parameters (same for all neurons)
bar_x = cos(direction)*(t - stim_start_time)/(5*ms) + stim_start_x : 1 (shared)
bar_y = sin(direction)*(t - stim_start_time)/(5*ms) + stim_start_y : 1 (shared)
direction : 1 (shared) # direction of the current stimulus
stim_start_time : second (shared) # start time of the current stimulus
stim_start_x : 1 (shared) # start position of the stimulus
stim_start_y : 1 (shared) # start position of the stimulus
'''
layer4 = NeuronGroup(N4*Nbarrels, eqs_layer4, threshold='rand() < rate*dt',
    method='euler', name='layer4')
layer4.barrel_x = '(i // N4) % barrelarraysize + 0.5'
layer4.barrel_y = 'i // (barrelarraysize*N4) + 0.5'
layer4.selectivity = '(i%N4)/(1.0*N4)*2*pi' # for each barrel, selectivity between 0-
    and 2*pi
stimradius = (11+1)*.5
# Chose a new randomly oriented bar every 60ms
runner_code =
'''
direction = rand()*2*pi
stim_start_x = barrelarraysize / 2.0 - cos(direction)*stimradius
stim_start_y = barrelarraysize / 2.0 - sin(direction)*stimradius
stim_start_time = t
'''
layer4.run_regularly(runner_code, dt=60*ms, when='start')

# Layer 2/3
# Model: IF with adaptive threshold
eqs_layer23 =
'''
dv/dt=(ge+gi+El-v)/taum : volt
dge/dt=-ge/taue : volt
dgi/dt=-gi/taui : volt
dvt/dt=(Vt-vt)/tauvt : volt # adaptation
barrel_idx : integer
x : 1 # in "barrel width" units
y : 1 # in "barrel width" units
'''
layer23 = NeuronGroup(Nbarrels*(N23exc+N23inh), eqs_layer23,
    threshold='v>vT', reset='v = El; vt += vt_inc',
    refractory=2*ms, method='euler', name='layer23')
layer23.v = El
layer23.vt = Vt

(continues on next page)
# Subgroups for excitatory and inhibitory neurons in layer 2/3
layer23exc = layer23[:Nbarrels*N23exc]
layer23inh = layer23[Nbarrels*N23exc:]

# Layer 2/3 excitatory
# The units for x and y are the width/height of a single barrel
layer23exc.x = '(i % (barrelarraysize*M23exc)) * (1.0/M23exc)'
layer23exc.y = '(i // (barrelarraysize*M23exc)) * (1.0/M23exc)'
layer23exc.barrel_idx = 'floor(x) + floor(y)*barrelarraysize'

# Layer 2/3 inhibitory
layer23inh.x = '(i % (barrelarraysize*M23inh)) * (1.0/M23inh)'
layer23inh.y = '(i // (barrelarraysize*M23inh)) * (1.0/M23inh)'
layer23inh.barrel_idx = 'floor(x) + floor(y)*barrelarraysize'

print("Building synapses, please wait...")
# Feedforward connections (plastic)
feedforward = Synapses(layer4, layer23exc,
model='''w:volt
\[dA_source/dt = -A_source/taup : volt (event-driven)\]
\[dA_target/dt = -A_target/taud : volt (event-driven)\]
''',
on_pre='''ge+=w
A_source += Ap
w = clip(w+A_target, 0, EPSC)''',
on_post='''
A_target += Ad
w = clip(w+A_source, 0, EPSC)''',
name='feedforward')

# Connect neurons in the same barrel with 50% probability
feedforward.connect('(barrel_x_pre + barrelarraysize*barrel_y_pre) == barrel_idx_post', p=0.5)
feedforward.w = EPSC*.5
print('excitatory lateral')
# Excitatory lateral connections
recurrent_exc = Synapses(layer23exc, layer23, model='w:volt', on_pre='ge+=w',
name='recurrent_exc')
recurrent_exc.connect(p='.15*exp(-.5*(((x_pre-x_post)/.4)**2+((y_pre-y_post)/.4)**2))
',
recurrent_exc.w['j<Nbarrels*N23exc'] = EPSC*.3 # excitatory->excitatory
recurrent_exc.w['j>=Nbarrels*N23exc'] = EPSC # excitatory->inhibitory

# Inhibitory lateral connections
print('inhibitory lateral')
recurrent_inh = Synapses(layer23inh, layer23exc, on_pre='gi+=IPSC',
name='recurrent_inh')
recurrent_inh.connect(p='exp(-.5*(((x_pre-x_post)/.2)**2+((y_pre-y_post)/.2)**2))
')

if get_device().__class__.__name__=='RuntimeDevice':
    print('Total number of connections')
    print('feedforward: %d' % len(feedforward))
    print('recurrent exc: %d' % len(recurrent_exc))
    print('recurrent inh: %d' % len(recurrent_inh))
t2 = time.time()
print("Construction time: \%.1fs" % (t2 - t1))

run(5*second, report='text')

# Calculate the preferred direction of each cell in layer23 by doing a
# vector average of the selectivity of the projecting layer4 cells, weighted
# by the synaptic weight.
_r = bincount(feedforward.j,
weights=feedforward.w * cos(feedforward.selectivity_pre)/feedforward.N_incoming,
minlength=len(layer23exc))
_i = bincount(feedforward.j,
weights=feedforward.w * sin(feedforward.selectivity_pre)/feedforward.N_incoming,
minlength=len(layer23exc))
selectivity_exc = (arctan2(_r, _i) % (2*pi))*180./pi

scatter(layer23.x[:Nbarrels*N23exc], layer23.y[:Nbarrels*N23exc],
c=selectivity_exc[:Nbarrels*N23exc],
edgecolors='none', marker='s', cmap='hsv')
vlines(np.arange(barrelarraysize), 0, barrelarraysize, 'k')
hlines(np.arange(barrelarraysize), 0, barrelarraysize, 'k')
clim(0, 360)
colorbar()
show()
5.11.13 Example: Morris_Lecar_1981

Morris-Lecar model

Reproduces Fig. 9 of:


```python
from brian2 import *
set_device('cpp_standalone')
defaulclock.dt = 0.01*ms

g_L = 2*mS
g_Ca = 4*mS
g_K = 8*mS
V_L = -50*mV
V_Ca = 100*mV
V_K = -70*mV
lambda_n__max = 1.0/(15*ms)
V_1 = 10*mV
V_2 = 15*mV  # Note that Figure caption says ~15 which seems to be a typo
V_3 = -1*mV

(continues on next page)```
V_4 = 14.5*mV
C = 20*uF

# V,N-reduced system (Eq. 9 in article), note that the variables M and N (and lambda_ →N, etc.)
# have been renamed to m and n to better match the Hodgkin-Huxley convention, and
# because N has
# a reserved meaning in Brian (number of neurons)
eqs = '''
dV/dt = (-g_L*(V - V_L) - g_Ca*m_inf*(V - V_Ca) - g_K*n*(V - V_K) + I)/C : volt
dn/dt = lambda_n*(n_inf - n) : 1
m_inf = 0.5*(1 + tanh((V - V_1)/V_2)) : 1
n_inf = 0.5*(1 + tanh((V - V_3)/V_4)) : 1
lambda_n = lambda_n__max*cosh((V - V_3)/(2*V_4)) : Hz
I : amp
'''
neuron = NeuronGroup(17, eqs, method='exponential_euler')
neuron.I = (np.arange(17)*25+100)*uA
neuron.V = V_L
neuron.n = 'n_inf'
mon = StateMonitor(neuron, ['V', 'n'], record=True)
run_time = 220*ms
run(run_time)
fig, (ax1, ax2) = plt.subplots(1, 2, gridspec_kw={'right': 0.95, 'bottom': 0.15}, figsize=(6.4, 3.2))
fig.subplots_adjust(wspace=0.4)
for line_no, idx in enumerate([0, 4, 12, 15]):
    color = 'C%d' % line_no
    ax1.plot(mon.t/ms, mon.V[idx]/mV, color=color)
    ax1.text(225, mon.V[idx][-1]/mV, '%.0f' % (neuron.I[idx]/uA), color=color)
ax1.set(xlim=(0, 220), ylim=(-50, 50), xlabel='time (ms)')
ax1.set_ylabel('V (mV)', rotation=0)
ax1.spines['right'].set_visible(False)
ax1.spines['top'].set_visible(False)
# dV/dt nullclines
V = linspace(-50, 50, 100)*mV
for line_no, (idx, color) in enumerate([(0, 'C0'), (4, 'C1'), (8, 'C4'), (12, 'C2'), (16, 'C5')]):
    n_null = (g_L*(V - V_L) + g_Ca*0.5*(1 + tanh((V - V_1)/V_2))*(V - V_Ca) - neuron.I[idx])/(-g_K*(V - V_K))
    ax2.plot(V/mV, n_null, color=color)
    ax2.text(V[20+5*line_no]/mV, n_null[20+5*line_no]+0.01, '%.0f' % (neuron.I[idx]/uA), color=color)
# dn/dt nullcline
n_null = 0.5*(1 + tanh((V - V_3)/V_4))
ax2.plot(V/mV, n_null, color='k')
ax2.set(xlim=(-50, 50), ylim=(0, 1), xlabel='V (mV)'
ax2.set_ylabel('n', rotation=0)
ax2.spines['right'].set_visible(False)
ax2.spines['top'].set_visible(False)
plt.show()
5.11.14 Example: Platkiewicz_Brette_2011

Slope-threshold relationship with noisy inputs, in the adaptive threshold model

Fig. 5E,F from:


```python
from scipy import optimize
from scipy.stats import linregress
from brian2 import *

N = 200  # 200 neurons to get more statistics, only one is shown
duration = 1*second
# --Biophysical parameters
ENa = 60*mV
EL = -70*mV
vT = -55*mV
Vi = -63*mV
tauh = 5*ms
tau = 5*ms
ka = 5*mV
ki = 6*mV
a = ka / ki
tauI = 5*ms
mu = 15*mV
sigma = 6*mV / sqrt(tauI / (tauI + tau))

# --Theoretical prediction for the slope-threshold relationship (approximation: \( a = 1 + \epsilon \))
```

(continues on next page)
thresh = \lambda \text{ slope, a: } \text{ Vi - slope } \times \text{ tauh } \times \log(1 + (\text{ Vi - vT} / a) / (\text{ slope } \times \text{ tauh}))
\# \text{ ------Exact calculation of the slope-threshold relationship}
\# \text{ (note that optimize.fsolve does not work with units, we therefore let th be a}
\# \text{ unitless quantity, i.e. the value in volt).}
thresh_ex = \lambda \text{ s: } \text{ optimize.fsolve (lambda th: (a} \times \text{ tauh}\times \exp((\text{ Vi-}\text{ th}\times \text{ volt})/(\text{ s} \times \text{ tauh})) - \\
\text{ - th}\times \text{ volt}\times (1-a) - a} \times (\text{ s} \times \text{ tauh} + \text{ Vi}) + \text{ vT}) / \text{ volt},
\text{ thresh(s, a)} \times \text{ volt}

eqs = ""
\text{ dv/dt= (EL-v+mu+sigma*I)/tau : volt}
\text{ dtheta/dt=(vT+a*clip(v-Vi, 0*mV, inf*mV)-theta)/tauh : volt}
\text{ dI/dt=-I/tauI+(2/tauI)**.5*xi : 1 # Ornstein-Uhlenbeck}
""
neurons = NeuronGroup(N, eqs, threshold="v>theta", reset='v=EL',
\text{ refractory=5*ms})
neurons.v = EL
neurons.theta = vT
neurons.I = 0
S = SpikeMonitor(neurons)
M = StateMonitor(neurons, 'v', record=True)
Mt = StateMonitor(neurons, 'theta', record=0)
run(duration, report='text')
\# Linear regression gives depolarization slope before spikes
\text{ tx = M.t[ M.t > 0*second] \& (M.t < 1.5 * tauh)]}
\text{ slope, threshold = [], []}
\text{ for (i, t) in zip(S.i, S.t):
\text{ ind = (M.t < t) \& (M.t > t - tauh)
\text{ mx = M.v[i, ind]
\text{ s, __, __, __, __ = linregress(tx[:len(mx)])/ms, mx/mV)
\text{ slope.append(s)
\text{ threshold.append(mx[-1])}
\# Figure
\text{ fig, (ax1, ax2) = plt.subplots(1, 2, figsize=(10, 4))}
\text{ ax1.plot(M.t/ms, M.v[0]/mV, 'k')
\text{ ax1.plot(M.t/ms, M.theta[0]/mV, 'r')
\# Display spikes on the trace}
\text{ spike_timesteps = np.round(S.t[S.i == 0]/defaultclock.dt).astype(int)
\text{ ax1.vlines(S.t[S.i == 0]/ms,
\text{ M.v[0, spike_timesteps]/mV,
\text{ 0, color='r'})
\text{ ax1.plot(S.t[S.i == 0]/ms, M.v[0, spike_timesteps]/mV, 'ro', ms=3)
\text{ ax1.set(xlabel='Time (ms)', ylabel='Voltage (mV)', xlim=(0, 500),
\text{ ylim=(-75, -35))}
\text{ ax2.plot(slope, Quantity(threshold)/mV, 'r.')}
\text{ sx = linspace(0.5*mV/ms, 4*mV/ms, 100)
\text{ t = Quantity([thresh_ex(s)) for s in sx])
\text{ ax2.plot(sx/(mV/ms), t/mV, 'k')
\text{ ax2.set(xlim=(0.5, 4), xlabel='Depolarization slope (mV/ms)',
\text{ ylabel='Threshold (mV)')}
\text{ fig.tight_layout()}
"""
5.11.15 Example: Rossant_et_al_2011bis

5.11.16 Distributed synchrony example

Fig. 14 from:

5000 independent E/I Poisson inputs are injected into a leaky integrate-and-fire neuron. Synchronous events, following an independent Poisson process at 40 Hz, are considered, where 15 E Poisson spikes are randomly shifted to be synchronous at those events. The output firing rate is then significantly higher, showing that the spike timing of less than 1% of the excitatory synapses have an important impact on the postsynaptic firing.

```brian2
from brian2 import *

# neuron parameters
theta = -55*mV
El = -65*mV
vmean = -65*mV
taum = 5*ms
taue = 3*ms
taui = 10*ms
eqs = Equations(""
    dv/dt = (ge+gi-(v-El))/taum : volt
    dge/dt = -ge/taue : volt
    dgi/dt = -gi/taui : volt
""

# input parameters
p = 15
ne = 4000
```

(continues on next page)
ni = 1000
lambda_c = 40*Hz
lambda_e = lambda_i = 1*Hz

# synapse parameters
we = .5*mV/(tau_m/tau_e)**(tau_m/(tau_e-tau_m))
wi = (v_mean-El-lambda_e*ne*we*tau_e)/(lambda_e*ni*tau_i)

# NeuronGroup definition

group = NeuronGroup(N=2, model=eqs, reset='v = El',
                 threshold='v>theta',
                 refractory=5*ms, method='exact')
group.v = El
group.ge = group.gi = 0

# independent E/I Poisson inputs
p1 = PoissonInput(group[0:1], 'ge', N=ne, rate=lambda_e, weight=we)
p2 = PoissonInput(group[0:1], 'gi', N=ni, rate=lambda_i, weight=wi)

# independent E/I Poisson inputs + synchronous E events
p3 = PoissonInput(group[1:], 'ge', N=ne, rate=lambda_e-(p*1.0/ne)*lambda_c, weight=we)
p4 = PoissonInput(group[1:], 'gi', N=ni, rate=lambda_i, weight=wi)
p5 = PoissonInput(group[1:], 'ge', N=1, rate=lambda_c, weight=p*we)

# run the simulation
M = SpikeMonitor(group)
SM = StateMonitor(group, 'v', record=True)
BrianLogger.log_level_info()
run(1*second)

# plot trace and spikes
for i in [0, 1]:
    spikes = (M.t[M.i == i] - defaultclock.dt)/ms
    val = SM[i].v
    subplot(2,1,i+1)
    plot(SM.t/ms, val)
    plot(tile(spikes, (2,1)),
         vstack((val[array(spikes, dtype=int)],
                zeros(len(spikes)))), 'C0')
    title("$s: %d spikes/second" % {
                "uncorrelated inputs": M.count[0],
                "correlated inputs": M.count[1]})

tight_layout()
show()
5.11.17 Example: Rothman_Manis_2003

Cochlear neuron model of Rothman & Manis


All model types differ only by the maximal conductances.

Adapted from their Neuron implementation by Romain Brette

```python
from brian2 import *

#defaultclock.dt=0.025*ms # for better precision

Simulation parameters: choose current amplitude and neuron type
(from type1c, typelt, type12, type21, type2, type2o)

neuron_type = 'type1c'
Ipulse = 250*pA
```

(continues on next page)
C = 12*pF
Eh = -43*mV
EK = -70*mV  # -77*mV in mod file
EL = -65*mV
ENa = 50*mV
nf = 0.85  # proportion of n vs p kinetics
zss = 0.5  # steady state inactivation of glt
temp = 22.  # temperature in degree celcius
q10 = 3. ** ((temp - 22) / 10.)
# hcno current (octopus cell)
frac = 0.0
qt = 4.5 ** ((temp - 33.) / 10.)

# Maximal conductances of different cell types in nS
maximal_conductances = dict(
    type1c=(1000, 150, 0, 0, 0.5, 0, 2),
    type1t=(1000, 80, 0, 65, 0.5, 0, 2),
    type12=(1000, 150, 20, 0, 2, 0, 2),
    type21=(1000, 150, 35, 0, 3.5, 0, 2),
    type2=(1000, 150, 200, 0, 20, 0, 2),
    type2o=(1000, 150, 600, 0, 0, 40, 2)  # octopus cell
)

gnabar, gkhtbar, gktbar, gkabar, ghbar, gbarno, gl = [x * nS for x in maximal_conductances[neuron_type]]

# Classical Na channel
eqs_na = ```
inga = gnabar*m**3*h*(ENa-v) : amp
dm/dt=q10*(minf-m)/mtau : 1
dh/dt=q10*(hinf-h)/htau : 1
minf = 1./(1+exp(-(vu + 38.) / 7.)) : 1
hinf = 1./(1+exp((vu + 65.) / 6.)) : 1
mtau = ((10. / (5*exp((vu+60.) / 18.) + 36.*exp(-(vu+60.) / 25.))) + 0.04)*ms : second
htau = ((100. / (7*exp((vu+60) / 11.) + 21*exp(-(vu+60) / 23.))) + 0.6)*ms : second
```  

# KHT channel (delayed-rectifier K+)
eqs_kht = ```
inght = gkhtbar*(nf*n**2 + (1-nf)*p)*(EK-v) : amp
dn/dt=q10*(ninf-n)/ntau : 1
dp/dt=q10*(pinf-p)/ptau : 1
ninf = (1 + exp(-(vu + 15) / 5.))**-0.5 : 1
pinf = 1. / (1 + exp(-(vu + 23) / 6.)) : 1
ntau = ((100. / (11*exp((vu+60) / 24.) + 21*exp(-(vu+60) / 23.))) + 0.7)*ms : second
ptau = ((100. / (4*exp((vu+60) / 32.) + 5*exp(-(vu+60) / 22.))) + 5)*ms : second
```  

# Ih channel (subthreshold adaptive, non-inactivating)
eqs_ih = ```
ingh = ghbar*r*(Eh-v) : amp
dr/dt=q10*(rinf-r)/rtau : 1
rinf = 1. / (1+exp((vu + 76.) / 7.)) : 1
rtau = ((100000. / (237.*exp((vu+60.) / 12.) + 17.*exp(-(vu+60.) / 14.))) + 25.)*ms : second
```  

(continues on next page)
# KLT channel (low threshold K+)
eqs_klt = ""
  iklt = gkltbar*w**4*z*(Ek-v) : amp
  dw/dt=q10*(winf-w)/wtau : 1
  dz/dt=q10*(zinf-z)/ztau : 1
  winf = (1. / (1 + exp(-(vu + 48.) / 6.)))**0.25 : 1
  zinf = zss + ((1.-zss) / (1 + exp((vu + 71.) / 10.))) : 1
  wtau = (((100. / (6.*exp((vu+60.) / 6.) + 16.*exp(-(vu+60.) / 45.))) + 1.5)*ms : second
  ztau = (((1000. / (exp((vu+60.) / 20.) + exp(-(vu+60.) / 8.))) + 50)*ms : second
"

# Ka channel (transient K+)
eqs_ka = ""
  ik = gkabar*a**4*b*c*(Ek-v) : amp
  da/dt=q10*(ainf-a)/atau : 1
  db/dt=q10*(binf-b)/btau : 1
  dc/dt=q10*(cinf-c)/ctau : 1
  ainf = (1. / (1 + exp(-(vu + 31) / 6.)))**0.25 : 1
  binf = 1. / (1 + exp((vu + 66) / 7.))**0.5 : 1
  cinf = 1. / (1 + exp((vu + 66) / 7.))**0.5 : 1
  atau = (((100. / (7*exp((vu+60) / 14.) + 29*exp(-(vu+60) / 24.))) + 0.1)*ms : second
  btau = (((1000. / (14*exp((vu+60) / 27.) + 29*exp(-(vu+60) / 24.))) + 1)*ms : second
  ctau = (((90. / (1 + exp((-66-vu) / 17.))) + 10)*ms : second
"

# Leak

eqs_leak = ""
  ileak = gl*(El-v) : amp
"

# h current for octopus cells

eqs_hcno = ""
  ihcno = gbarno*(h1*frac + h2*(1-frac))*(Eh-v) : amp
  dh1/dt=(hinfno-h1)/tau1 : 1
  dh2/dt=(hinfno-h2)/tau2 : 1
  hinfno = 1./(1+exp((vu+66.)/7.)) : 1
  tau1 = bet1/(qt*0.008*(1+alp1))*ms : second
  tau2 = bet2/(qt*0.0029*(1+alp2))*ms : second
  alp1 = exp(1e-3*3*(vu+50)*9.648e4/(8.315*(273.16+temp))) : 1
  bet1 = exp(1e-3*3*0.3*(vu+50)*9.648e4/(8.315*(273.16+temp))) : 1
  alp2 = exp(1e-3*3*(vu+84)*9.648e4/(8.315*(273.16+temp))) : 1
  bet2 = exp(1e-3*3*0.6*(vu+84)*9.648e4/(8.315*(273.16+temp))) : 1
"

eqs = ""
  dv/dt = (ileak + ina + ikht + iklt + ika + ih + ihcno + I)/C : volt
  vu = v/mV : 1 # unitless v
  I : amp
"

eqs += eqs_leak + eqs_ka + eqs_na + eqs_ih + eqs_klt + eqs_kht + eqs_hcno

neuron = NeuronGroup(1, eqs, method='exponential_euler')
neuron.v = El

run(50*ms, report='text') # Go to rest

(continues on next page)
M = StateMonitor(neuron, 'v', record=0)
neuron.I = Ipulse

run(100*ms, report='text')

plot(M.t / ms, M[0].v / mV)
xlabel('t (ms)')
ylabel('v (mV)')
show()

5.11.18 Example: Sturzl_et_al_2000

Adapted from Theory of Arachnid Prey Localization W. Sturzl, R. Kempter, and J. L. van Hemmen PRL 2000

Poisson inputs are replaced by integrate-and-fire neurons

Romain Brette

```python
from brian2 import *

# Parameters
degree = 2 * pi / 360.
```

5.11. frompapers
duration = 500*ms
R = 2.5*cm  # radius of scorpion
vr = 50*meter/second  # Rayleigh wave speed
phi = 144*degree  # angle of prey
A = 250*Hz
deltaI = .7*ms  # inhibitory delay
gamma = (22.5 + 45 * arange(8)) * degree  # leg angle
delay = R / vr * (1 - cos(phi - gamma))  # wave delay

# Wave (vector w)
time = arange(int(duration / defaultclock.dt) + 1) * defaultclock.dt
Dtot = 0.
w = 0.
for f in arange(150, 451)*Hz:
    D = exp(-(f/Hz - 300) ** 2 / (2 * (50 ** 2)))
    rand_angle = 2 * pi * rand()
    w += 100 * D * cos(2 * pi * f * time + rand_angle)
    Dtot += D
w = .01 * w / Dtot

# Rates from the wave
rates = TimedArray(w, dt=defaultclock.dt)

# Leg mechanical receptors
tau_legs = 1 * ms
sigma = .01
eqs_legs = ""
    dv/dt = (1 + rates(t - d) - v)/tau_legs + sigma*(2./tau_legs)**.5*xi:1
d : second
""
legs = NeuronGroup(8, model=eqs_legs, threshold='v > 1', reset='v = 0',
    refractory=1*ms, method='euler')
legs.d = delay
spikes_legs = SpikeMonitor(legs)

# Command neurons
tau = 1 * ms
taus = 1.001 * ms
wex = 7
winh = -2
eqs_neuron = ""
    dv/dt = (x - v)/tau : 1
dx/dt = (y - x)/taus : 1 # alpha currents
dy/dt = -y/taus : 1
""
neurons = NeuronGroup(8, model=eqs_neuron, threshold='v>1', reset='v=0',
    method='exact')
synapses_ex = Synapses(legs, neurons, on_pre='y+=wex')
synapses_ex.connect(j='i')
synapses_inh = Synapses(legs, neurons, on_pre='y+=winh', delay=deltaI)
synapses_inh.connect(abs(((j - i) % N_post) - N_post/2) <= 1')
spikes = SpikeMonitor(neurons)
run(duration, report='text')
nspikes = spikes.count
phi_est = imag(log(sum(nspikes * exp(gamma * 1j))))
print("True angle (deg): %.2f" % (phi/degree))
print("Estimated angle (deg): %.2f" % (phi_est/degree))
rmmax = amax(nspikes)/duration/Hz
polar(concatenate((gamma, [gamma[0] + 2 * pi])),
      concatenate((nspikes, [nspikes[0]])) / duration / Hz,
      c='k')
axvline(phi, ls='-', c='g')
axvline(phi_est, ls='-', c='b')
show()

5.11.19 Example: Touboul_Brette_2008

Chaos in the AdEx model

Fig. 8B from: Touboul, J. and Brette, R. (2008). Dynamics and bifurcations of the adaptive exponential integrate-and-fire model. Biological Cybernetics 99(4-5):319-34.

This shows the bifurcation structure when the reset value is varied (vertical axis shows the values of w at spike times for a given a reset value Vr).
from brian2 import *

defaultclock.dt = 0.01*ms

C = 281*pF
gL = 30*nS
EL = -70.6*mV
VT = -50.4*mV
DeltaT = 2*mV
tauw = 40*ms
a = 4*nS
b = 0.08*nA
I = .8*nA
Vcut = VT + 5 * DeltaT  # practical threshold condition
N = 200

eqs = 
"dvm/dt=(gL*(EL-vm)+gL*DeltaT*exp((vm-VT)/DeltaT)+I-w)/C : volt
dw/dt=(a*(vm-EL)-w)/tauw : amp
Vr:volt"

neuron = NeuronGroup(N, model=eqs, threshold='vm > Vcut',
                      reset="vm = Vr; w += b", method='euler')
neuron.vm = EL
neuron.w = a * (neuron.vm - EL)
neuron.Vr = linspace(-48.3 * mV, -47.7 * mV, N)  # bifurcation parameter

init_time = 3*second
run(init_time, report='text')  # we discard the first spikes

states = StateMonitor(neuron, "w", record=True, when='start')
spikes = SpikeMonitor(neuron)
run(1 * second, report='text')

# Get the values of Vr and w for each spike
Vr = neuron.Vr[spikes.i]
w = states.w[spikes.i, int_((spikes.t-init_time)/defaultclock.dt)]

figure()
plot(Vr / mV, w / nA, '.k')
xlabel('Vr (mV)')
ylabel('w (nA)')
show()
5.11.20 Example: Vogels_et_al_2011

Inhibitory synaptic plasticity in a recurrent network model

(F. Zenke, 2011) (from the 2012 Brian twister)

Adapted from:


```python
from brian2 import *

# Defining network model parameters
# ###########################################
NE = 8000  # Number of excitatory cells
NI = NE/4   # Number of inhibitory cells

tau_ampa = 5.0*ms  # Glutamatergic synaptic time constant
tau_gaba = 10.0*ms  # GABAergic synaptic time constant
```

(continues on next page)
epsilon = 0.02  # Sparseness of synaptic connections

tau_stdp = 20*ms  # STDP time constant

simtime = 10*second  # Simulation time

# # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # #
# Neuron model
# # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # #

gl = 10.0*siemens  # Leak conductance
el = -60*mV  # Resting potential
er = -80*mV  # Inhibitory reversal potential
vt = -50.*mV  # Spiking threshold
memc = 200.0*pifarad  # Membrane capacitance
bgcurrent = 200*pA  # External current

eqs_neurons=''
dv/dt=(-gl*(v-el)-(g_ampa*v+g_gaba*(v-er))+bgcurrent)/memc : volt (unless refractory)
dg_ampa/dt = -g_ampa/tau_ampa : siemens
dg_gaba/dt = -g_gaba/tau_gaba : siemens''

# # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # #
# Initialize neuron group
# # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # #

neurons = NeuronGroup(NE+NI, model=eqs_neurons, threshold='v > vt',
                       reset='v=el', refractory=5*ms, method='euler')
Pe = neurons[:NE]
Pi = neurons[NE:]

# # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # #
# Connecting the network
# # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # #

con_e = Synapses(Pe, neurons, on_pre='g_ampa += 0.3*nS')
con_e.connect(p=epsilon)
con_ii = Synapses(Pi, Pi, on_pre='g_gaba += 3*nS')
con_ii.connect(p=epsilon)

# # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # #
# Inhibitory Plasticity
# # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # # #
eqs_stdp_inhib = '''
    w : 1
    dApre/dt=-Apre/tau_stdp : 1 (event-driven)
    dApost/dt=-Apost/tau_stdp : 1 (event-driven)
    '''
    alpha = 3*Hz*tau_stdp*2  # Target rate parameter
    gmax = 100  # Maximum inhibitory weight

con_ie = Synapses(Pi, Pe, model=eqs_stdp_inhib,
                  on_pre='''Apre += 1.
                      w = clip(w+(Apost-alpha)*eta, 0, gmax)
                      g_gaba += w*nS''','
                  p=epsilon)
on_post='''Apost += 1.
    w = clip(w+Apre*eta, 0, gmax)
''

con_ie.connect(p=epsilon)
con_ie.w = 1e-10

# Setting up monitors
sm = SpikeMonitor(Pe)

# Run without plasticity
eta = 0  # Learning rate
run(1*second)

# Run with plasticity
eta = 1e-2  # Learning rate
run(simtime-1*second, report='text')

# Make plots
i, t = sm.it
subplot(211)
plot(t/ms, i, 'k.', ms=0.25)
title("Before")
xlabel(""")
yticks([])
xlim(0.8*1e3, 1*1e3)
subplot(212)
plot(t/ms, i, 'k.', ms=0.25)
xlabel("time (ms)")
yticks([])
title("After")
xlim((simtime-0.2*second)/ms, simtime/ms)
show()
5.11.21 Example: Wang_Buszaki_1996

Wang-Buszaki model


Note that implicit integration (exponential Euler) cannot be used, and therefore simulation is rather slow.

```python
from brian2 import *

defaultclock.dt = 0.01*ms
Cm = 1*uF # /cm**2
Iapp = 2*uA
gL = 0.1*msiemens
EL = -65*mV
ENa = 55*mV
EK = -90*mV
gNa = 35*msiemens
gK = 9*msiemens
```

(continues on next page)
\texttt{eqs = \'''}
\texttt{dv/dt = (-gNa*m**3*h*(v-ENa)-gK*n**4*(v-EK)-gL*(v-EL)+Iapp)/Cm : volt}
\texttt{m = alpha_m/(alpha_m+beta_m) : 1}
\texttt{alpha_m = 0.1/mV*10*mV/exprel(-(v+35*mV)/(10*mV))/ms : Hz}
\texttt{beta_m = 4*exp(-(v+60*mV)/(18*mV))/ms : Hz}
\texttt{dh/dt = 5*(alpha_h*(1-h)-beta_h*h) : 1}
\texttt{alpha_h = 0.07*exp(-(v+58*mV)/(20*mV))/ms : Hz}
\texttt{beta_h = 1./(exp(-0.1/mV*(v+28*mV))+1)/ms : Hz}
\texttt{dn/dt = 5*(alpha_n*(1-n)-beta_n*n) : 1}
\texttt{alpha_n = 0.01/mV*10*mV/exprel(-(v+34*mV)/(10*mV))/ms : Hz}
\texttt{beta_n = 0.125*exp(-(v+44*mV)/(80*mV))/ms : Hz}
\texttt{'''}

\texttt{neuron = NeuronGroup(1, eqs, method='exponential_euler')}
\texttt{neuron.v = -70*mV}
\texttt{neuron.h = 1}
\texttt{M = StateMonitor(neuron, 'v', record=0)}

\texttt{run(100*ms, report='text')}

\texttt{plot(M.t/ms, M[0].v/mV)}
\texttt{show()}
5.12 frompapers/Brette_2012

5.12.1 Example: Fig1


Fig 1C-E. Somatic voltage-clamp in a ball-and-stick model with Na channels at a particular location.

```python
from brian2 import *
from params import *

defaultclock.dt = 0.025*ms

# Morphology
morpho = Soma(50*um)  # chosen for a target Rm
morpho.axon = Cylinder(diameter=1*um, length=300*um, n=300)

location = 40*um  # where Na channels are placed
duration = 500*ms

# Channels
eqs='''
Im = gL*(EL - v) + gclamp*(vc - v) + gNa*m*(ENa - v) : amp/meter**2
dm/dt = (minf - m) / taum: 1  # simplified Na channel
minf = 1 / (1 + exp((va - v) / ka)) : 1
gclamp : siemens/meter**2
GNa : siemens/meter**2
vc = EL + 50*mV * t/duration : volt (shared)  # Voltage clamp with a ramping voltage
'''

neuron = SpatialNeuron(morphology=morpho, model=eqs, Cm=Cm, Ri=Ri)
compartment = morpho.axon[location]
neuron.v = EL
neuron.gclamp[0] = gL*500
neuron.gNa[compartment] = gNa_0/neuron.area[compartment]

# Monitors
mon = StateMonitor(neuron, ['v', 'vc', 'm'], record=True)

run(duration, report='text')

subplot(221)
plot(mon[0].vc/mV,
     -(mon[0].vc - mon[0].v)*(neuron.gclamp[0])*neuron.area[0]/nA, 'k')
xlabel('V (mV)')
ylabel('I (nA)')
xlim(-75, -45)
title('I-V curve')

subplot(222)
plot(mon[0].vc/mV, mon[compartment].m, 'k')
xlabel('V (mV)')
ylabel('m')
```

(continues on next page)
title('Activation curve (m(V))')

subplot(223)
# Number of simulation time steps for each volt increment in the voltage-clamp
dt_per_volt = len(mon.t)/(50*mV)
for v in [-64*mV, -61*mV, -58*mV, -55*mV]:
    plot(mon.v[:,1:100,int(dt_per_volt*(v - EL))]/mV, 'k')
xlabel('Distance from soma (um)')
ylabel('V (mV)')

subplot(224)
plot(mon[compartment].v/mV, mon[compartment].v/mV, 'k--') # Diagonal
plot(mon[0].v/mV, mon[compartment].v/mV, 'k')
xlabel('Vs (mV)')
ylabel('Va (mV)')
tight_layout()
show()

5.12.2 Example: Fig3AB

Fig. 3. A, B. Kink with only Nav1.6 channels

```python
from brian2 import *
from params import *

codegen.target='numpy'

defaultclock.dt = 0.025*ms

# Morphology
morpho = Soma(50*um)  # chosen for a target Rm
morpho.axon = Cylinder(diameter=1*um, length=300*um, n=300)

location = 40*um  # where Na channels are placed

# Channels
eqs='''
Im = gL*(EL - v) + gNa*m*(ENa - v) : amp/meter**2
dm/dt = (minf - m) / taum : 1  # simplified Na channel
minf = 1 / (1 + exp((va - v) / ka)) : 1
gNa : siemens/meter**2
Iin : amp (point current)
'''

neuron = SpatialNeuron(morphology=morpho, model=eqs, Cm=Cm, Ri=Ri,
                       method="exponential_euler")

compartment = morpho.axon[location]
neuron.v = EL
neuron.gNa[compartment] = gNa_0/neuron.area[compartment]
M = StateMonitor(neuron, ['v', 'm'], record=True)

run(20*ms, report='text')
neuron.Iin[0] = gL * 20*mV * neuron.area[0]
run(80*ms, report='text')

subplot(121)
plot(M.t/ms, M[0].v/mV, 'r')
plot(M.t/ms, M[compartment].v/mV, 'k')
plot(M.t/ms, M[compartment].m*(80+60)-80, 'k--')  # open channels
ylim(-80, 60)
xlabel('Time (ms)')
ylabel('V (mV)')
title('Voltage traces')

subplot(122)
dm = diff(M[0].v) / defaultclock.dt
dm40 = diff(M[compartment].v) / defaultclock.dt
plot((M[0].v/mV)[1:], dm/(volt/second), 'r')
plot((M[compartment].v/mV)[1:], dm40/(volt/second), 'k')
xlim(-80, 40)
xlabel('V (mV)')
ylabel('dV/dt (V/s)')
title('Phase plot')
tight_layout()
show()
```
5.12.3 Example: Fig3CF


Fig. 3C-F. Kink with Nav1.6 and Nav1.2

```python
from brian2 import *
from params import *

defaultclock.dt = 0.01*ms

# Morphology
morpho = Soma(50*um) # chosen for a target Rm
morpho.axon = Cylinder(diameter=1*um, length=300*um, n=300)

location16 = 40*um # where Nav1.6 channels are placed
location12 = 15*um # where Nav1.2 channels are placed

va2 = va + 15*mV # depolarized Nav1.2

# Channels
duration = 100*ms
```

(continues on next page)
eqs='''
Im = gL * (EL - v) + gNa*m*(ENa - v) + gNa2*m2*(ENa - v) : amp/meter**2
dm/dt = (minf - m) / taum : 1 # simplified Na channel
minf = 1 / (1 + exp((va - v) / ka)) : 1
dm2/dt = (minf2 - m2) / taum : 1 # simplified Na channel, Nav1.2
minf2 = 1 / (1 + exp((va2 - v) / ka)) : 1
gNa : siemens/meter**2
gNa2 : siemens/meter**2 # Nav1.2
Iin : amp (point current)
'''

neuron = SpatialNeuron(morphology=morpho, model=eqs, Cm=Cm, Ri=Ri,
                       method="exponential_euler")
compartment16 = morpho.axon[location16]
compartment12 = morpho.axon[location12]
neuron.v = EL
neuron.gNa[compartment16] = gNa_0/neuron.area[compartment16]
neuron.gNa2[compartment12] = 20*gNa_0/neuron.area[compartment12]

# Monitors
M = StateMonitor(neuron, ['v', 'm', 'm2'], record=True)
run(20*ms, report='text')
neuron.Iin[0] = gL * 20*mV * neuron.area[0]
run(80*ms, report='text')

subplot(221)
plot(M.t/ms, M[0].v/mV, 'r')
plot(M.t/ms, M[compartment16].v/mV, 'k--') # open channels
ylim(-80, 60)
xlabel('Time (ms)')
ylabel('V (mV)')
title('Voltage traces')

subplot(222)
plot(M[0].v/mV, 1 / (1 + exp((va - M[0].v) / ka)), 'k--')
plot(M[0].v/mV, M[compartment12].m2, 'r--')
xlim(-70, 0)
xlabel('V (mV)')
ylabel('m')
title('Activation curves')

subplot(223)
dm = diff(M[0].v) / defaultclock.dt
dm40 = diff(M[compartment16].v) / defaultclock.dt
plot((M[0].v/mV)[1:], dm/(volt/second), 'r')
plot((M[compartment16].v/mV)[1:], dm40/(volt/second), 'k')
xlim(-80, 40)
xlabel('V (mV)')
ylabel('dV/dt (V/s)')
title('Phase plot')

subplot(224)
plot((M[0].v/mV)[1:], dm/(volt/second), 'r')
plot((M[compartment16].v/mV)[1:], dm40/(volt/second), 'k')

(continues on next page)
plot((M[0].v/mV)[1:], 10 + 0*xm/(volt/second), 'k--')
xlim(-70, -40)
ylim(0, 20)
xlabel('V (mV)')
ylabel('dV/dt (V/s)')
title('Phase plot(zoom)')
tight_layout()
show()

5.12.4 Example: Fig4


Fig. 4E-F. Spatial distribution of Na channels. Tapering axon near soma.

from brian2 import *
from params import *

defaultclock.dt = 0.025*ms

(continues on next page)
# Morphology
morpho = Soma(50*um)  # chosen for a target Rm
# Tapering (change this for the other figure panels)
diameters = hstack([linspace(4, 1, 11), ones(290)])*um
morpho.axon = Section(diameter=diameters, length=ones(300)*um, n=300)

# Na channels
Na_start = (25 + 10)*um
Na_end = (40 + 10)*um
linear_distribution = True  # True is F, False is E

duration = 500*ms

# Channels
eqs='''
Im = gL*(EL - v) + gclamp*(vc - v) + gNa*m*(ENa - v) : amp/meter**2
dm/dt = (minf - m) / taum: 1  # simplified Na channel
minf = 1 / (1 + exp((va - v) / ka)) : 1
gclamp : siemens/meter**2
gNa : siemens/meter**2
vc = EL + 50*mV * t / duration : volt (shared)  # Voltage clamp with a ramping voltage command
'''

neuron = SpatialNeuron(morphology=morpho, model=eqs, Cm=Cm, Ri=Ri,
method="exponential_euler")
compartments = morpho.axon[Na_start:Na_end]
neuron.v = EL
neuron.gclamp[0] = gL*500
if linear_distribution:
    profile = linspace(1, 0, len(compartments))
else:
    profile = ones(len(compartments))
profile = profile / sum(profile)  # normalization

neuron.gNa[compartments] = gNa_0 * profile / neuron.area[compartments]

# Monitors
mon = StateMonitor(neuron, 'v', record=True)
run(duration, report='text')

dt_per_volt = len(mon.t) / (50*mV)
for v in [-64*mV, -61*mV, -58*mV, -55*mV, -52*mV]:
    plot(mon.v[:100, int(dt_per_volt * (v - EL))]/mV, 'k')
xlim(0, 50+10)
ylim(-65, -25)
ylabel('V (mV)')
xlabel('Location (um)')
title('Voltage across axon')
tight_layout()
show()
5.12.5 Example: Fig5A


Fig. 5A. Voltage trace for current injection, with an additional reset when a spike is produced.

Trick: to reset the entire neuron, we use a set of synapses from the spike initiation compartment where the threshold condition applies to all compartments, and the reset operation \( v = EL \) is applied there every time a spike is produced.

```python
from brian2 import *
from params import *

defaultclock.dt = 0.025*ms
duration = 500*ms

# Morphology
morpho = Soma(50*um)  # chosen for a target Rm
morpho.axon = Cylinder(diameter=1*um, length=300*um, n=300)

# Input
taux = 5*ms
sigmax = 12*mV
```

(continues on next page)
xx0 = 7*mV

compartment = 40

# Channels
eqs = '''
Im = gL * (EL - v) + gNa * m * (ENa - v) + gLx * (xx0 + xx) : amp/meter**2  
dm/dt = (minf - m) / taum : 1 # simplified Na channel  
minf = 1 / (1 + exp((va - v) / ka)) : 1  
gNa : siemens/meter**2  
gLx : siemens/meter**2  
dxx/dt = -xx / taux + sigmax * (2 / taux)**.5 *xi : volt  
'''

neuron = SpatialNeuron(morphology=morpho, model=eqs, Cm=Cm, Ri=Ri,  
threshold='m>0.5', threshold_location=compartment,  
refractory=5*ms)

neuron.v = EL  
neuron.gLx[0] = gL  
neuron.gNa[compartment] = gNa_0 / neuron.area[compartment]

# Reset the entire neuron when there is a spike
reset = Synapses(neuron, neuron, on_pre='v = EL')  
reset.connect('i == compartment') # Connects the spike initiation compartment to all compartments

# Monitors
S = SpikeMonitor(neuron)  
M = StateMonitor(neuron, 'v', record=0)  
run(duration, report='text')

# Add spikes for display
v = M[0].v
for t in S.t:  
    v[int(t / defaultclock.dt)] = 50*mV

plot(M.t/ms, v/mV, 'k')
tight_layout()  
show()
5.12.6 Example: params

Parameters for spike initiation simulations.

```python
from brian2.units import *

# Passive parameters
EL = -75*mV
S = 7.85e-9*meter**2  # area (sphere of 50 um diameter)
Cm = 0.75*uF/cm**2
gL = 1. / (30000*ohm*cm**2)
Ri = 150*ohm*cm

# Na channels
ENa = 60*mV
ka = 6*mV
va = -40*mV
gNa_0 = gL * 2*S
taum = 0.1*ms
```
5.13 frompapers/Stimberg_et_al_2018

5.13.1 Example: example_1_COBA

Modeling neuron-glia interactions with the Brian 2 simulator Marcel Stimberg, Dan F. M. Goodman, Romain Brette, Maurizio De Pittà bioRxiv 198366; doi: https://doi.org/10.1101/198366

Figure 1: Modeling of neurons and synapses.

Randomly connected networks with conductance-based synapses (COBA; see Brunel, 2000). Synapses exhibit short-time plasticity (Tsodyks, 2005; Tsodyks et al., 1998).

```python
from brian2 import *
import sympy
import plot_utils as pu

seed(11922)  # to get identical figures for repeated runs

# Model parameters
# General parameters
duration = 1.0*second  # Total simulation time
sim_dt = 0.1*ms  # Integrator/sampling step
N_e = 3200  # Number of excitatory neurons
N_i = 800  # Number of inhibitory neurons

### Neuron parameters
E_l = -60*mV  # Leak reversal potential
g_l = 9.99*nS  # Leak conductance
E_e = 0*mV  # Excitatory synaptic reversal potential
E_i = -80*mV  # Inhibitory synaptic reversal potential
C_m = 198*pF  # Membrane capacitance
tau_e = 5*ms  # Excitatory synaptic time constant
tau_i = 10*ms  # Inhibitory synaptic time constant
tau_r = 5*ms  # Refractory period
I_ex = 150*pA  # External current
V_th = -50*mV  # Firing threshold
V_r = E_l  # Reset potential

### Synapse parameters
w_e = 0.05*nS  # Excitatory synaptic conductance
w_i = 1.0*nS  # Inhibitory synaptic conductance
U_0 = 0.6  # Synaptic release probability at rest
Omega_d = 2.0/second  # Synaptic depression rate
Omega_f = 3.33/second  # Synaptic facilitation rate

# Model definition
# Set the integration time (in this case not strictly necessary, since we are
# using the default value)
defaultclock.dt = sim_dt

### Neurons
(continues on next page)
```
neuron_eqs = '''
v/dt = (g_l*(E_l-v) + g_e*(E_e-v) + g_i*(E_i-v) + 
       I_ex)/C_m : volt (unless refractory)
dq_e/dt = -g_e/tau_e : siemens # post-synaptic exc. conductance
dq_i/dt = -g_i/tau_i : siemens # post-synaptic inh. conductance
'''
neurons = NeuronGroup(N_e + N_i, model=neuron_eqs,
                      threshold='v>V_th', reset='v=V_r',
                      refractory='tau_r', method='euler')
# Random initial membrane potential values and conductances
neurons.v = 'E_l + rand()*(V_th-E_l)'
neurons.g_e = 'rand()w_e'
neurons.g_i = 'rand()*w_i'
exc_neurons = neurons[:N_e]
inh_neurons = neurons[N_e:]

### Synapses
synapses_eqs = '''
# Usage of releasable neurotransmitter per single action potential:
du_S/dt = -Omega_f * u_S : 1 (event-driven)
# Fraction of synaptic neurotransmitter resources available:
dx_S/dt = Omega_d *(1 - x_S) : 1 (event-driven)
'''
synapses_action = '''
u_S += U_0 * (1 - u_S)
r_S = u_S * x_S
x_S -= r_S
'''
exc_syn = Synapses(exc_neurons, neurons, model=synapses_eqs,
on_pre=synapses_action+'g_e_post += w_e*r_S')
inh_syn = Synapses(inh_neurons, neurons, model=synapses_eqs,
on_pre=synapses_action+'g_i_post += w_i*r_S')
exc_syn.connect(p=0.05)
inh_syn.connect(p=0.2)
# Start from "resting" condition: all synapses have fully-replenished
# neurotransmitter resources
exc_syn.x_S = 1
inh_syn.x_S = 1

# Monitors
# Note that we could use a single monitor for all neurons instead, but in this
# way plotting is a bit easier in the end
exc_mon = SpikeMonitor(exc_neurons)
inh_mon = SpikeMonitor(inh_neurons)

### We record some additional data from a single excitatory neuron
ni = 50
# Record conductances and membrane potential of neuron ni
state_mon = StateMonitor(exc_neurons, ['v', 'g_e', 'g_i'], record=ni)
# We make sure to monitor synaptic variables after synapse are updated in order
# to use simple recurrence relations to reconstruct them. Record all synapses
# originating from neuron ni
synapse_mon = StateMonitor(exc_syn, ['u_S', 'x_S'],
                           record=exc_syn[ni, :], when='after_synapses')
# Simulation run
run(duration, report='text')

# Analysis and plotting
plt.style.use('figures.mplstyle')

## Spiking activity (w/ rate)
fig1, ax = plt.subplots(nrows=2, ncols=1, sharex=False,
                         gridspec_kw={'height_ratios': [3, 1],
                                     'left': 0.18, 'bottom': 0.18, 'top': 0.95,
                                     'hspace': 0.1},
                         figsize=(3.07, 3.07))
ax[0].plot(exc_mon.t[exc_mon.i <= N_e//4]/ms,
            exc_mon.i[exc_mon.i <= N_e//4], '|', color='C0')
ax[0].plot(inh_mon.t[inh_mon.i <= N_i//4]/ms,
            inh_mon.i[inh_mon.i <= N_i//4]+N_e//4, '|', color='C1')
pu.adjust_spines(ax[0], ['left'])
ax[0].set(xlim=(0.,duration/ms), ylim=(0,(N_e+N_i)//4), ylabel='neuron index')

# Generate frequencies
bin_size = 1*ms
spk_count, bin_edges = np.histogram(np.r_[exc_mon.t/ms, inh_mon.t/ms],
                                     int(duration/ms))
rate = double(spk_count)/(N_e + N_i)/bin_size/Hz
ax[1].plot(bin_edges[:-1], rate, '-', color='k')
pu.adjust_spines(ax[1], ['left', 'bottom'])
ax[1].set(xlim=(0.,duration/ms), ylim=(0, 10.),
          xlabel='time (ms)', ylabel='rate (Hz)')
pu.adjust_ylabels(ax, x_offset=-0.18)

## Dynamics of a single neuron
fig2, ax = plt.subplots(4, sharex=False,
                        gridspec_kw={'left': 0.27, 'bottom': 0.18, 'top': 0.95,
                                     'hspace': 0.2},
                        figsize=(3.07, 3.07))
### Postsynaptic conductances
ax[0].plot(state_mon.t/ms, state_mon.g_e[0]/nS, color='C0')
ax[0].plot(state_mon.t/ms, -state_mon.g_i[0]/nS, color='C1')
ax[0].plot([state_mon.t[0]/ms, state_mon.t[-1]/ms], [0, 0], color='grey',
            linestyle=':')
pu.adjust_spines(ax[0], ['left'])
ax[0].set(xlim=(0.,duration/ms), ylim=(-5.0,0.25),
          ylabel='postsyn. conduct. ('+sympy.latex(nS)+')')

### Membrane potential
ax[1].axhline(V_th/mV, color='C2', linestyle=':')
# Threshold
# Artificially insert spikes
ax[1].plot(state_mon.t/ms, state_mon.v[0]/mV, color='black')
ax[1].vlines(exc_mon.t[exc_mon.i == ni]/ms, V_th/mV, 0, color='black')
pu.adjust_spines(ax[1], ['left'])
ax[1].set(xlim=(0., duration/ms), ylim=(-1+V_r/mV,0.),
          ylabel='postsyn. conduct. \n$'+sympy.latex(nS)+'$')
(continues on next page)
ylabel='membrane potential ($0$') format (sympy.latex(mV))

### Synaptic variables

# Retrieves indexes of spikes in the synaptic monitor using the fact that we are sampling spikes and synaptic variables by the same dt
spk_index = np.in1d(synapse_mon.t, exc_mon.t[exc_mon.i == ni])
ax[2].plot(synapse_mon.t[spk_index]/ms, synapse_mon.x_S[0][spk_index], '.', ms=4, color='C3')
ax[2].plot(synapse_mon.t[spk_index]/ms, synapse_mon.u_S[0][spk_index], '.', ms=4, color='C4')
# Super-impose reconstructed solutions

time = synapse_mon.t  # time vector
tspk = Quantity(synapse_mon.t, copy=True)  # Spike times
for ts in exc_mon.t[exc_mon.i == ni]:
    tspk[time >= ts] = ts
ax[2].plot(synapse_mon.t/ms, 1 + (synapse_mon.x_S[0]-1)*exp(-(time-tspk)*Omega_d), '-', color='C3')
ax[2].plot(synapse_mon.t/ms, synapse_mon.u_S[0]*exp(-(time-tspk)*Omega_f), '-', color='C4')
# Adjust axis
pu.adjust_spines(ax[2], ['left'])
ax[2].set(xlim=(0., duration/ms), ylim=(-0.05, 1.05),
ylabel='synaptic variables $u_S, x_S$')

nspikes = np.sum(spk_index)
x_S_spike = synapse_mon.x_S[0][spk_index]
u_S_spike = synapse_mon.u_S[0][spk_index]
aax[3].vlines(synapse_mon.t[spk_index]/ms, np.zeros(nspikes),
x_S_spike*u_S_spike/(1-u_S_spike))
pu.adjust_spines(ax[3], ['left', 'bottom'])
aax[3].set(xlim=(0., duration/ms), ylim=(-0.01, 0.62),
yticks=np.arange(0, 0.62, 0.2), xlabel='time (ms)', ylabel='$r_S$')

pu.adjust_ylabels(ax, x_offset=-0.20)

plt.show()
5.13.2 Example: example_2_gchi_astrocyte

Modeling neuron-glia interactions with the Brian 2 simulator Marcel Stimberg, Dan F. M. Goodman, Romain Brette, Maurizio De Pitta bioRxiv 198366; doi: https://doi.org/10.1101/198366

Figure 2: Modeling of synaptically-activated astrocytes

Two astrocytes (one stochastic and the other deterministic) activated by synapses (connecting “dummy” groups of neurons) (see De Pitta et al., 2009)

```python
from brian2 import *
import plot_utils as pu
```

(continues on next page)
set_device('cpp_standalone', directory=None)  # Use fast "C++ standalone mode"
seed(790824)  # to get identical figures for repeated runs

#---------------------------------------------------------------
# Model parameters
#---------------------------------------------------------------
## General parameters
duration = 30*second  # Total simulation time
sim_dt = 1*ms  # Integrator/sampling step

## Neuron parameters
f_0 = 0.5*Hz  # Spike rate of the "source" neurons

## Synapse parameters
rho_c = 0.001  # Synaptic vesicle-to-extracellular space volume ratio
Y_T = 500*mmolar  # Total vesicular neurotransmitter concentration
Omega_c = 40/second  # Neurotransmitter clearance rate

## Astrocyte parameters
# --- Calcium fluxes
G_P = 0.9*umolar/second  # Maximal Ca\(^{2+}\) uptake rate by SERCAs
K_P = 0.1 * umolar  # Ca\(^{2+}\) affinity of SERCAs
C_T = 2*umolar  # Total cell free Ca\(^{2+}\) content
rhostress_A = 0.18  # ER-to-cytoplasm volume ratio
Omega_C = 6/second  # Maximal rate of Ca\(^{2+}\) release by IP\(_3\)Rs
Omega_L = 0.1/second  # Maximal rate of Ca\(^{2+}\) leak from the ER
# --- IP\(_3\)R kinetics
d_1 = 0.13*umolar  # IP\(_3\) binding affinity
d_2 = 1.05*umolar  # Ca\(^{2+}\) inactivation dissociation constant
G_2 = 0.2/second  # IP\(_3\)R binding rate for Ca\(^{2+}\) inhibition
d_3 = 0.9434*umolar  # IP\(_3\) dissociation constant
d_5 = 0.08*umolar  # Ca\(^{2+}\) activation dissociation constant
# --- Agonist-dependent IP\(_3\) production
O_beta = 5*umolar/second  # Maximal rate of IP\(_3\) production by PLC\(_\beta\)
O_N = 0.3/umolar/second  # Agonist binding rate
Omega_N = 0.5/second  # Maximal inactivation rate
K_KC = 0.5*umolar  # Ca\(^{2+}\) affinity of PKC
zeta = 10  # Maximal reduction of receptor affinity by PKC
# --- IP\(_3\) production
O_delta = 0.2*umolar/second  # Maximal rate of IP\(_3\) production by PLC\(_\delta\)
kappa_delta = 1.5 * umolar  # Inhibition constant of PLC\(_\delta\) by IP\(_3\)
K_delta = 0.3*umolar  # Ca\(^{2+}\) affinity of PLC\(_\delta\)
# --- IP\(_3\) degradation
Omega_5P = 0.1/second  # Maximal rate of IP\(_3\) degradation by IP-5P
K_D = 0.5*umolar  # Ca\(^{2+}\) affinity of IP\(_3\)-3K
K_3K = 1*umolar  # IP\(_3\) affinity of IP\(_3\)-3K
O_3K = 4.5*umolar/second  # Maximal rate of IP\(_3\) degradation by IP\(_3\)-3K
# --- IP\(_3\) external production
F_ex = 0.09*umolar/second  # Maximal exogenous IP3 flow
I_Theta = 0.3*umolar  # Threshold gradient for IP\(_3\) diffusion
omega_I = 0.05*umolar  # Scaling factor of diffusion

# Model definition
#---------------------------------------------------------------
defaultclock.dt = sim_dt  # Set the integration time

(continues on next page)
### "Neurons"

(We are only interested in the activity of the synapse, so we replace the neurons by trivial "dummy" groups)

#### Regular spiking neuron

source_neurons = NeuronGroup(1, 'dx/dt = f_0 : 1', threshold='x>1', reset='x=0', method='euler')

#### Dummy neuron

target_neurons = NeuronGroup(1, '')

### Synapses

(Our synapse model is trivial, we are only interested in its neurotransmitter release)

synapses_eqs = 'dY_S/dt = -Omega_c * Y_S : mmolar (clock-driven)'
synapses_action = 'Y_S += rho_c * Y_T'
synapses = Synapses(source_neurons, target_neurons, model=synapses_eqs, on_pre=synapses_action, method='exact')
synapses.connect()

### Astrocytes

(We are modelling two astrocytes, the first is deterministic while the second displays stochastic dynamics)

astro_eqs = ''

# Fraction of activated astrocyte receptors:

dGamma_A/dt = O_N * Y_S * (1 - Gamma_A) - Omega_N*(1 + zeta * C/(C + K_KC)) * Gamma_A : 1

# IP_3 dynamics:

dI/dt = J_beta + J_delta - J_3K - J_5P + J_ex : mmolar/second

J_beta = O_beta * Gamma_A : mmolar/second

J_delta = O_delta/(1 + I/kappa_delta) * C**4/(C**4 + K_delta**4) : mmolar/second

J_3K = O_3K * C**4/(C**4 + K_D**4) * I/(I + K_3K) : mmolar/second

J_5P = Omega_5P*I : mmolar/second

delta_I_bias = I - I_bias : mmolar

J_ex = -F_ex/2*(1 + tanh((abs(delta_I_bias) - I_Theta)/omega_I)) * sign(delta_I_bias) : mmolar/second

I_bias : mmolar (constant)

# Ca^2+-induced Ca^2+ release:

dC/dt = J_r + J_l - J_p : mmolar/second

# IP3R de-inactivation probability

dh/dt = (h_inf - h_clipped)/tau_h * (1 + noise*xi*tau_h**0.5) : 1

h_clipped = clip(h,0,1) : 1

J_r = (Omega_C * m_inf**3 * h_clipped**3) * (C_T - (1 + rho_A)*C) : mmolar/second

J_l = Omega_L * (C_T - (1 + rho_A)*C) : mmolar/second

J_p = O_P * C**2/(C**2 + K_P**2) : mmolar/second

m_inf = I/(I + d_1) * C/(C + d_5) : 1

h_inf = Q_2/(Q_2 + C) : 1

tau_h = 1/(O_2 * (Q_2 + C)) : second

Q_2 = d_2 * (I + d_1)/(I + d_3) : mmolar

# Neurotransmitter concentration in the extracellular space

Y_S : mmolar
# Noise flag
noise : 1 (constant)

# Milstein integration method for the multiplicative noise
astrocytes = NeuronGroup(2, astro_eqs, method='milstein')
astrocytes.h = 0.9  # IP3Rs are initially mostly available for CICR

# The first astrocyte is deterministic ("zero noise"), the second stochastic
astrocytes.noise = [0, 1]

# Connection between synapses and astrocytes (both astrocytes receive the
# same input from the synapse). Note that in this special case, where each
# astrocyte is only influenced by the neurotransmitter from a single synapse,
# the '(linked)' variable mechanism could be used instead. The mechanism used
# below is more general and can add the contribution of several synapses.
ecs_syn_to_astro = Synapses(synapses, astrocytes,

    'Y_S_post = Y_S_pre * mmolar (summed)')
ecs_syn_to_astro.connect()

# Monitors
astro_mon = StateMonitor(astrocytes, variables=['Gamma_A', 'C', 'h', 'I'],
record=True)

# Simulation run
run(duration, report='text')

# Analysis and plotting
from matplotlib.ticker import FormatStrFormatter
plt.style.use('figures.mplstyle')

# Plot Gamma_A
fig, ax = plt.subplots(4, 1, figsize=(6.26894, 6.26894*0.66))
ax[0].plot(astro_mon.t/second, astro_mon.Gamma_A.T)
ax[0].set(xlim=(0., duration/second), ylim=[-0.05, 1.02], yticks=[0.0, 0.5, 1.0],
ylabel=r'$\Gamma_{A}$')
pu.adjust_spines(ax[0], ['left'])

# Plot I
ax[1].plot(astro_mon.t/second, astro_mon.I.T/umolar)
ax[1].set(xlim=(0., duration/second), ylim=[-0.1, 5.0],
yticks=range(0.0, 5.1, 1.),
ylabel=r'$I$ ($\mu M$)')
ax[1].yaxis.set_major_formatter(FormatStrFormatter('%.1f'))
ax[1].legend(["deterministic", "stochastic"], loc='upper left')
pu.adjust_spines(ax[1], ['left'])

# Plot C
ax[2].plot(astro_mon.t/second, astro_mon.C.T/umolar)
ax[2].set(xlim=(0., duration/second), ylim=[-0.1, 1.3],
ylabel=r'$C$ ($\mu M$)')
pu.adjust_spines(ax[2], ['left'])

5.13. frompapers/Stimberg_et_al_2018
# Plot h
ax[3].plot(astro_mon.t/second, astro_mon.h.T)
ax[3].set(xlim=(0., duration/second),
    ylim=[0.4, 1.02],
    ylabel='h', xlabel='time ($s$)')
uu.adjust_spines(ax[3], ['left', 'bottom'])

uu.adjust_ylabels(ax, x_offset=-0.1)
plt.show()
### Model parameters

```
# General parameters
duration = transient + 600*ms  # Total simulation time
sim_dt = 1*ms  # Integrator/sampling step

# Synapse parameters
rho_c = 0.005  # Synaptic vesicle-to-extracellular space volume ratio
Y_T = 500*mmolar  # Total vesicular neurotransmitter concentration
Omega_c = 40/second  # Neurotransmitter clearance rate
U_0__star = 0.6  # Resting synaptic release probability
Omega_f = 3.33/second  # Synaptic facilitation rate
Omega_d = 2.0/second  # Synaptic depression rate

# --- Presynaptic receptors
O_G = 1.5/umolar/second  # Agonist binding (activating) rate
Omega_G = 0.5/(60*second)  # Agonist release (deactivating) rate

# Astrocyte parameters
# --- Calcium fluxes
O_P = 0.9*umolar/second  # Maximal Ca^2+ uptake rate by SERCAs
K_P = 0.05 * umolar  # Ca^2+ affinity of SERCAs
C_T = 2*umolar  # Total cell free Ca^2+ content
rho_A = 0.18  # ER-to-cytoplasm volume ratio
Omega_C = 6/second  # Maximal rate of Ca^2+ release by IP_3Rs
Omega_L = 0.1/second  # Maximal rate of Ca^2+ leak from the ER

# --- IP_3 kinetics
d_1 = 0.13*umolar  # IP_3 binding affinity
d_2 = 1.05*umolar  # Ca^2+ inactivation dissociation constant
O_2 = 0.2/umolar/second  # IP_3R binding rate for Ca^2+ inhibition
d_3 = 0.9434*umolar  # IP_3 dissociation constant
d_5 = 0.08*umolar  # Ca^2+ activation dissociation constant

# --- IP_3 production
O_delta = 0.6*umolar/second  # Maximal rate of IP_3 production by PLCdelta
kappa_delta = 1.5* umolar  # Inhibition constant of PLC_delta by IP_3
K_delta = 0.1*umolar  # Ca^2+ affinity of PLCdelta

# --- IP_3 degradation
Omega_5P = 0.05/second  # Maximal rate of IP_3 degradation by IP-5P
K_D = 0.7*umolar  # Ca^2+ affinity of IP3-3K
K_3K = 1.0*umolar  # IP_3 affinity of IP_3-3K
O_3K = 4.5*umolar/second  # Maximal rate of IP_3 degradation by IP_3-3K

# --- IP_3 diffusion
F_ex = 2.0*umolar/second  # Maximal exogenous IP3 flow
I_Theta = 0.3*umolar  # Threshold gradient for IP_3 diffusion
omega_I = 0.05*umolar  # Scaling factor of diffusion

# --- Gliotransmitter release and time course
C_Theta = 0.5*umolar  # Ca^2+ threshold for exocytosis
Omega_A = 0.6/second  # Gliotransmitter recycling rate
U_A = 0.6  # Gliotransmitter release probability
G_T = 200*mmolar  # Total vesicular gliotransmitter concentration
rho_e = 6.5e-4  # Astrocytic vesicle-to-extracellular volume ratio
Omega_e = 60/second  # Gliotransmitter clearance rate
alpha = 0.0  # Gliotransmission nature
```

(continues on next page)
# Model definition

```python
# defaultclock.dt = sim_dt   # Set the integration time

### "Neurons"
# We are only interested in the activity of the synapse, so we replace the
# neurons by trivial "dummy" groups
spikes = [0, 50, 100, 150, 200,
          300, 310, 320, 330, 340, 350, 360, 370, 380, 390, 400]*ms
spikes += transient  # allow for some initial transient
source_neurons = SpikeGeneratorGroup(1, np.zeros(len(spikes)), spikes)
target_neurons = NeuronGroup(1, '')

### Synapses
# Note that the synapse does not actually have any effect on the post-synaptic
# target
# Also note that for easier plotting we do not use the "event-driven" flag here,
# even though the value of u_S and x_S only needs to be updated on the arrival
# of a spike
synapses_eqs = ''

# Neurotransmitter
dY_S/dt = -Omega_c * Y_S : mmolar (clock-driven)
# Fraction of activated presynaptic receptors
dGamma_S/dt = O_G * G_A * (1 - Gamma_S) -
              Omega_G * Gamma_S : 1 (clock-driven)
# Usage of releasable neurotransmitter per single action potential:
du_S/dt = -Omega_f * u_S : 1 (clock-driven)
# Fraction of synaptic neurotransmitter resources available:
dx_S/dt = Omega_d *(1 - x_S) : 1 (clock-driven)
# released synaptic neurotransmitter resources:
r_S : 1
# gliotransmitter concentration in the extracellular space:
G_A : mmolar
'''
synapses_action = '''
U_0 = (1 - Gamma_S) * U_0__star + alpha * Gamma_S
u_S += U_0 * (1 - u_S)
r_S = u_S * x_S
x_S -= r_S
Y_S += rho_c * Y_T * r_S
'''
synapses = Synapses(source_neurons, target_neurons,
                     model=synapses_eqs, on_pre=synapses_action,
                     method='exact')

# We create three synapses, only the second and third ones are modulated by astrocytes
synapses.connect(True, n=3)

### Astrocytes
# The astrocyte emits gliotransmitter when its Ca^2+ concentration crosses
# a threshold
astro_eqs = '''

# IP_3 dynamics:
dI/dt = J_delta - J_3K - J_5P - J_ex : mmolar
J_delta = D_delta/(1 + I/kappa_delta) * C**2/(C**2 + K_delta**2) : mmolar/second
J_3K = D_3K * C**4/(C**4 + K_D**4) * I/(I + K_3K) : mmolar/second
J_5P = D_5P * I : mmolar/second
# Exogenous stimulation
```
\[ \text{delta}_I_{\text{bias}} = I - I_{\text{bias}} : \text{mmolar} \]
\[ J_{\text{ex}} = -F_{\text{ex}}/2 \times (1 + \tanh((\text{abs}(\text{delta}_I_{\text{bias}}) - I_{\text{Theta}})/\omega_I)) \times \text{sign}(\text{delta}_I_{\text{bias}}) : \text{mmolar/second} \]
\[ I_{\text{bias}} : \text{mmolar} \quad \text{constant} \]

# Ca^{2+}-induced Ca^{2+} release:
\[ \frac{dC}{dt} = \left( \Omega_C \times m_{\text{inf}}^3 \times h^3 + \Omega_L \right) \times (C_T - (1 + \rho_A) \times C) - \Omega_P \times C/(C^2 + K_P^2) : \text{mmolar} \]
\[ \frac{dh}{dt} = \frac{h_{\text{inf}} - h}{\tau_h} : 1 \quad \text{IP3R de-inactivation probability} \]
\[ m_{\text{inf}} = \frac{I/(I + d_1) \times C/(C + d_5)}{1} \]
\[ h_{\text{inf}} = \frac{Q_2}{Q_2 + C} : 1 \]
\[ \tau_h = \frac{1}{O_2 \times (Q_2 + C)} : \text{second} \]
\[ Q_2 = \frac{d_2 \times (I + d_3 \times I)}{I + d_1} : \text{mmolar} \]

# Fraction of gliotransmitter resources available:
\[ \frac{dx_A}{dt} = \Omega_A \times (1 - x_A) : 1 \]

# Gliotransmitter concentration in the extracellular space:
\[ \frac{dG_A}{dt} = -\Omega_{e} \times G_A : \text{mmolar} \]
\[ \text{glio_release} = '' \\
G_A += \rho_e \times G_T \times U_A \times x_A \\
x_A -= U_A \times x_A \\
'' \]

# The following formulation makes sure that a "spike" is only triggered at the first threshold crossing -- the astrocyte is considered "refractory" (i.e., not allowed to trigger another event) as long as the Ca^{2+} concentration remains above threshold
# The gliotransmitter release happens when the threshold is crossed, in Brian terms it can therefore be considered a "reset"

\[ \text{astrocyte} = \text{NeuronGroup}(2, \text{astro_eqs}, \quad \theta_r='C>C_Theta', \quad \text{refractory}='C>C_Theta', \quad \text{reset}=$\text{glio_release}$, \quad \text{method}='\text{rk4}') \]

# Different length of stimulation
\[ \text{astrocyte.x_A} = 1.0 \]
\[ \text{astrocyte.h} = 0.9 \]
\[ \text{astrocyte.I} = 0.4 \times \text{umolar} \]
\[ \text{astrocyte.I}_{\text{bias}} = \text{np.asarray}([0.8, 1.25]) \times \text{umolar} \]

# Connection between astrocytes and the second synapse. Note that in this special case, where the synapse is only influenced by the gliotransmitter from a single astrocyte, the "linked" variable mechanism could be used instead.
# The mechanism used below is more general and can add the contribution of several astrocytes
\[ \text{ecs astro to syn} = \text{Synapses(astrocyte, synapses, \quad \text{G}_A_{\text{post}} = \text{G}_A_{\text{pre}} : \text{mmolar (summed)})} \]

# Connect second and third synapse to a different astrocyte
\[ \text{ecs astro to syn.connect(j='i+1')} \]

# Monitors
# Note that we cannot use "record=True" for synapses in C++ standalone mode -- the StateMonitor needs to know the number of elements to record from during its initialization, but in C++ standalone mode, no synapses have been created yet. We therefore explicitly state to record from the three synapses.
\[ \text{syn_mon} = \text{StateMonitor(synapses, variables=['u_S', 'x_S', 'r_S', 'Y_S'])} \]
record=[0, 1, 2])
ast_mon = StateMonitor(astrocyte, variables=['C', 'G_A'], record=True)

# Simulation run
run(duration, report='text')

# Analysis and plotting
from matplotlib import cycler
plt.style.use('figures.mplstyle')
fig, ax = plt.subplots(nrows=7, ncols=1, figsize=(6.26894, 6.26894 * 1.2),
                       gridspec_kw={'height_ratios': [3, 2, 1, 1, 3, 3, 3],
                                    'top': 0.98, 'bottom': 0.08,
                                    'left': 0.15, 'right': 0.95})

## Ca^2+ traces of the two astrocytes
ax[0].plot((ast_mon.t-transient)/second, ast_mon.C[0]/umolar, '-', color='C2')
ax[0].plot((ast_mon.t-transient)/second, ast_mon.C[1]/umolar, '-', color='C3')

## Add threshold for gliotransmitter release
ax[0].plot(np.asarray([-transient/second,0.0]),
            np.asarray([C_Theta,C_Theta])/umolar, ':', color='gray')
ax[0].set(xlim=[-transient/second, 0.0], yticks=[0., 0.4, 0.8, 1.2],
          ylabel=r'$C$ ($\mu$M)')
pu.adjust_spines(ax[0], ['left'])

## Gliotransmitter concentration in the extracellular space
ax[1].plot((ast_mon.t-transient)/second, ast_mon.G_A[0]/umolar, '-', color='C2')
ax[1].plot((ast_mon.t-transient)/second, ast_mon.G_A[1]/umolar, '-', color='C3')
ax[1].set(yticks=[0., 50., 100.], xlim=[-transient/second, 0.0],
         xticklabels=[], yticklabels=['time (s)'], ylabel=r'$G_A$ ($\mu$M)')
pu.adjust_spines(ax[1], ['left', 'bottom'])

## Turn off one axis to display x-labeling of ax[1] correctly
ax[2].axis('off')

## Synaptic stimulation
ax[3].vlines((spikes-transient)/ms, 0, 1, clip_on=False)
ax[3].set(xlim=(0, (duration-transient)/ms))
ax[3].axis('off')

## Synaptic variables
# Use a custom cycle that uses black as the first color
prop_cycle = cycler(color='k').concat(matplotlib.rcParams['axes.prop_cycle'][2:])
ax[4].set(xlim=(0, (duration-transient)/ms), ylim=[0., 1.],
         yticks=matplotlib.arange(0, 1.1, .25), ylabel='$u_S$',
         prop_cycle=prop_cycle)
ax[4].plot((syn_mon.t-transient)/ms, syn_mon.u_S.T)
pu.adjust_spines(ax[4], ['left'])

ax[5].set(xlim=(0, (duration-transient)/ms), ylim=[-0.05, 1.],
         yticks=matplotlib.arange(0, 1.1, .25), ylabel='$x_S$',
         prop_cycle=prop_cycle)
ax[5].plot((syn_mon.t-transient)/ms, syn_mon.x_S.T)
pu.adjust_spines(ax[5], ['left'])

ax[6].set(xlim=(0, (duration-transient)/ms), ylim=(-5., 1500),
    xticks=np.arange(0, (duration-transient)/ms, 100), xlabel='time (ms)',
    yticks=[0, 500, 1000, 1500], ylabel=r'$Y_S$ ($\mu$M)',
    prop_cycle=prop_cycle)
ax[6].plot((syn_mon.t-transient)/ms, syn_mon.Y_S/umolar)
ax[6].legend(['no gliotransmission',
               'weak gliotransmission',
               'stronger gliotransmission'], loc='upper right')
pu.adjust_spines(ax[6], ['left', 'bottom'])
pu.adjust_ylabels(ax, x_offset=-0.11)
plt.show()
5.13.4 Example: example_4_rsmean

Modeling neuron-glia interactions with the Brian 2 simulator Marcel Stimberg, Dan F. M. Goodman, Romain Brette, Maurizio De Pittà bioRxiv 198366; doi: https://doi.org/10.1101/198366

Figure 4C: Closed-loop gliotransmission.
I/O curves in terms of average per-spike release vs. rate of stimulation for three synapses: one without gliotransmission, and the other two with open- and close-loop gliotransmission.

```python
from brian2 import *
import plot_utils as pu

set_device('cpp_standalone', directory=None)  # Use fast "C++ standalone mode"
seed(1929)  # to get identical figures for repeated runs

# Model parameters
### General parameters
N_synapses = 100
N_astro = 2
transient = 15*second
duration = transient + 180*second  # Total simulation time
sim_dt = 1*ms  # Integrator/sampling step

### Neuron parameters

### Synapse parameters
rho_c = 0.005  # Synaptic vesicle-to-extracellular space volume ratio
Y_T = 500*mmolar  # Total vesicular neurotransmitter concentration
Omega_c = 40/second  # Neurotransmitter clearance rate
U_0__star = 0.6  # Resting synaptic release probability
Omega_f = 3.33/second  # Synaptic facilitation rate
Omega_d = 2.0/second  # Synaptic depression rate

# --- Presynaptic receptors
O_G = 1.5/umolar/second  # Agonist binding (activating) rate
Omega_G = 0.5/(60*second)  # Agonist release (deactivating) rate

### Astrocyte parameters
# --- Calcium fluxes
G_P = 0.9*umolar/second  # Maximal Ca^2+ uptake rate by SERCAs
K_P = 0.05*umolar  # Ca^2+ affinity of SERCAs
C_T = 2*umolar  # Total cell free Ca^2+ content
rho_A = 0.18  # ER-to-cytoplasm volume ratio
Omega_C = 6/second  # Maximal rate of Ca^2+ release by IP_3Rs
Omega_L = 0.1/second  # Maximal rate of Ca^2+ leak from the ER

# --- IP_3R kinectics
d_1 = 0.13*umolar  # IP_3 binding affinity
d_2 = 1.05*umolar  # Ca^2+ inactivation dissociation constant
G_2 = 0.2/umolar/second  # IP_3R binding rate for Ca^2+ inhibition
d_3 = 0.9434*umolar  # IP_3 dissociation constant
d_5 = 0.08*umolar  # Ca^2+ activation dissociation constant

# --- IP_3 production

# --- Agonist-dependent IP_3 production
O_beta = 3.2*umolar/second  # Maximal rate of IP_3 production by PLCbeta
O_N = 0.3/umolar/second  # Agonist binding rate
Omega_N = 0.5/second  # Maximal inactivation rate
K_KC = 0.5*umolar  # Ca^2+ affinity of PKC
zeta = 10  # Maximal reduction of receptor affinity by PKC

# --- Endogenous IP3 production
O_delta = 0.6*umolar/second  # Maximal rate of IP_3 production by PLCdelta
kappa_delta = 1.5*umolar  # Inhibition constant of PLC_delta by IP_3
```

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K_delta = 0.1*umolar  # Ca^2+ affinity of PLCdelta
--- IP_3 degradation
Omega_5P = 0.05/second  # Maximal rate of IP_3 degradation by IP-5P
K_D = 0.7*umolar  # Ca^2+ affinity of IP3-3K
K_3K = 1.0*umolar  # IP_3 affinity of IP_3-3K
Q_3K = 4.5*umolar/second  # Maximal rate of IP_3 degradation by IP_3-3K
--- IP_3 diffusion
F_ex = 2.0*umolar/second  # Maximal exogenous IP3 flow
I_Theta = 0.3*umolar  # Threshold gradient for IP_3 diffusion
omega_I = 0.05*umolar  # Scaling factor of diffusion
--- Gliotransmitter release and time course
C_Theta = 0.5*umolar  # Ca^2+ threshold for exocytosis
Omega_A = 0.6/second  # Gliotransmitter recycling rate
U_A = 0.6  # Gliotransmitter release probability
G_T = 200*mmolar  # Total vesicular gliotransmitter concentration
rho_e = 6.5e-4  # Astrocytic vesicle-to-extracellular volume ratio
Omega_e = 60/second  # Gliotransmitter clearance rate
alpha = 0.0  # Gliotransmission nature

defaultclock.dt = sim_dt  # Set the integration time

f_vals = np.logspace(-1, 2, N_synapses)*Hz
source_neurons = PoissonGroup(N_synapses, rates=f_vals)
target_neurons = NeuronGroup(N_synapses, '')

### Synapses
# Note that the synapse does not actually have any effect on the post-synaptic
# target
# Also note that for easier plotting we do not use the "event-driven" flag here,
# even though the value of u_S and x_S only needs to be updated on the arrival
# of a spike
synapses_eqs = ''
# Neurotransmitter
dY_S/dt = -Omega_c * Y_S : mmolar (clock-driven)
# Fraction of activated presynaptic receptors
dGamma_S/dt = O_G * G_A * (1 - Gamma_S) - Omega_G * Gamma_S : 1 (clock-driven)
# Usage of releasable neurotransmitter per single action potential:
du_S/dt = -Omega_f * u_S : 1 (event-driven)
# Fraction of synaptic neurotransmitter resources available for release:
dx_S/dt = Omega_d *(1 - x_S) : 1 (event-driven)
r_S : 1 # released synaptic neurotransmitter resources
G_A : mmolar # gliotransmitter concentration in the extracellular space

synapses_action = ''
U_0 = (1 - Gamma_S) * U_0__star + alpha * Gamma_S
u_S += U_0 * (1 - u_S)
x_S -= r_S
Y_S += rho_c * Y_T * r_S

synapses = Synapses(source_neurons, target_neurons,
model=synapses_eqs, on_pre=synapses_action,
method='exact')

# We create three synapses per connection; only the first two are modulated by
the astrocyte however. Note that we could also create three synapses per connection with a single connect call by using connect(j='i', n=3), but this would create synapses arranged differently (synapses connection pairs (0, 0), (0, 0), (0, 0), (1, 1), (1, 1), (1, 1), ...), instead of connections (0, 0), (1, 1), ..., (0, 0), (1, 1), ..., (0, 0), (1, 1), ...). Making the later connection descriptions more complicated.

```python
synapses.connect(j='i') # closed-loop modulation
synapses.connect(j='i') # open modulation
synapses.connect(j='i') # no modulation
synapses.x_S = 1.0
```

### Astrocytes

The astrocyte emits gliotransmitter when its Ca^2+ concentration crosses a threshold:

```python
astro_eqs = '''
# Fraction of activated astrocyte receptors:
dGamma_A/dt = O_N * Y_S * (1 - Gamma_A) -
                Omega_N*(1 + zeta * C/(C + K_KC)) * Gamma_A : 1

# IP_3 dynamics:
dI/dt = J_beta + J_delta - J_3K - J_5P + J_ex : mmolar
J_beta = O_beta * Gamma_A : mmolar/second
J_delta = O_delta/(1 + I/kappa_delta) * C**2/(C**2 + K_delta**2) : mmolar/second
J_3K = O_3K * C**4/(C**4 + K_D**4) * I/(I + K_3K) : mmolar/second
J_5P = Omega_5P*I : mmolar/second
delta_I_bias = I - I_bias : mmolar
J_ex = -F_ex/2*(1 + tanh((abs(delta_I_bias) - I_Theta)/omega_I)) * sign(delta_I_bias) : mmolar/second
I_bias : mmolar (constant)

# Ca^2+-induced Ca^2+ release:
dC/dt = (Omega_C * m_inf**3 * h**3 + Omega_L) * (C_T - (1 + rho_A)*C) -
      O_P * C**2/(C**2 + K_P**2) : mmolar
dh/dt = (h_inf - h)/tau_h : 1  # IP3R de-inactivation probability
m_inf = I/(I + d_1) * C/(C + d_5) : 1
h_inf = Q_2/(Q_2 + C) : 1
tau_h = 1/(O_2 * (Q_2 + C)) : second
Q_2 = d_2 * (I + d_1)/(I + d_3) : mmolar

# Fraction of gliotransmitter resources available for release
dx_A/dt = Omega_A * (1 - x_A) : 1
# gliotransmitter concentration in the extracellular space
dG_A/dt = -Omega_e*G_A : mmolar
# neurotransmitter concentration in the extracellular space
Y_S : mmolar

'''
glio_release = '''
G_A += rho_e * G_T * U_A * x_A
x_A -= U_A * x_A
'''

astrocyte = NeuronGroup(N_astro*N_synapses, astro_eqs,
                        # The following formulation makes sure that a "spike" is only triggered at the first threshold crossing
                        threshold="C>C_Theta",
                        refractory="C>C_Theta",
                        # The gliotransmitter release happens when the threshold
                        glio_release = glio_release
                        )
```

(continues on next page)
# is crossed, in Brian terms it can therefore be
# considered a "reset"
reset=glio_release,
    method='rk4')
astrocyte.h = 0.9
astrocyte.x_A = 1.0
# Only the second group of N_synapses astrocytes are activated by external stimulation
astrocyte.I_bias = (np.r_[np.zeros(N_synapses), np.ones(N_synapses)])*1.0*umolar

## Connections
ecs_syn_to_astro = Synapses(synapses, astrocyte,
    'Y_S_post = Y_S_pre : mmolar (summed)')
# Connect the first N_synapses synapses--astrocyte pairs
ecs_syn_to_astro.connect(j='i if i < N_synapses')

ecs_astro_to_syn = Synapses(astrocyte, synapses,
    'G_A_post = G_A_pre : mmolar (summed)')
# Connect the first N_synapses astrocytes--pairs
# (closed-loop configuration)
ecs_astro_to_syn.connect(j='i if i < N_synapses')
# Connect the second N_synapses astrocyte--synapses pairs
# (open-loop configuration)
ecs_astro_to_syn.connect(j='i if i >= N_synapses and i < 2*N_synapses')

# Monitors
syn_mon = StateMonitor(synapses, 'r_S',
    record=np.arange(N_synapses*(N_astro+1)))

# Simulation run
run(duration, report='text')

# Analysis and plotting
plt.style.use('figures.mplstyle')
fig, ax = plt.subplots(nrows=4, ncols=1, figsize=(3.07, 3.07*1.33), sharex=False,
    gridspec_kw={'height_ratios': [1, 3, 3, 3],
                 'top': 0.98, 'bottom': 0.12,
                 'left': 0.22, 'right': 0.93})

## Turn off one axis to display accordingly to the other figure in example_4_synrel.py
ax[0].axis('off')

ax[1].errorbar(f_vals/Hz, np.mean(syn_mon.r_S[2*N_synapses:], axis=1),
    np.std(syn_mon.r_S[2*N_synapses:], axis=1),
    fnt='o', color='black', lw=0.5)
ax[1].set(xlim=(0.08, 100), xscale='log',
    ylim=(0., 0.7),
    ylabel=r'$\langle r_S \rangle$')
pu.adjust_spines(ax[1], ['left'])

ax[2].errorbar(f_vals/Hz, np.mean(syn_mon.r_S[N_synapses:2*N_synapses], axis=1),
5.13.5 Example: example_4_synrel

Modeling neuron-glia interactions with the Brian 2 simulator Marcel Stimberg, Dan F. M. Goodman, Romain Brette, Maurizio De Pittà bioRxiv 198366; doi: https://doi.org/10.1101/198366

Figure 4B: Closed-loop gliotransmission.

Extracellular neurotransmitter concentration (averaged across 500 synapses) for three step increases of the presynaptic
rate, for three synapses: one without gliotransmission, and the other two with open- and close-loop gliotransmission.

```python
from brian2 import *

import plot_utils as pu

set_device('cpp_standalone', directory=None)  # Use fast "C++ standalone mode"
seed(16283)  # to get identical figures for repeated runs

# Model parameters
N_synapses = 500
N_astro = 2
duration = 20*second  # Total simulation time
sim_dt = 1*ms  # Integrator/sampling step

### General parameters
rho_c = 0.005  # Synaptic vesicle-to-extracellular space volume ratio
Y_T = 500*mmolar  # Total vesicular neurotransmitter concentration
Omega_c = 40/second  # Neurotransmitter clearance rate
U_0__star = 0.6  # Resting synaptic release probability
Omega_f = 3.33/second  # Synaptic facilitation rate
Omega_d = 2.0/second  # Synaptic depression rate

### Synapse parameters

### Astrocyte parameters
# --- Calcium fluxes
O_P = 0.9*umolar/second  # Maximal Ca^2+ uptake rate by SERCAs
K_P = 0.05*umolar  # Ca^2+ affinity of SERCAs
C_T = 2*umolar  # Total cell free Ca^2+ content
rho_A = 0.18  # ER-to-cytoplasm volume ratio
Omega_A = 6/second  # Maximal rate of Ca^2+ release by IP_3Rs
Omega_L = 0.1/second  # Maximal rate of Ca^2+ leak from the ER

# --- IP_3R kinetics
d_1 = 0.13*umolar  # IP_3 binding affinity
d_2 = 1.05*umolar  # Ca^2+ inactivation dissociation constant
O_2 = 0.2/umolar/second  # IP_3 binding rate for Ca^2+ inhibition
d_3 = 0.9434*umolar  # IP_3 dissociation constant
d_5 = 0.08*umolar  # Ca^2+ activation dissociation constant

# --- IP_3 production
O_beta = 3.2*umolar/second  # Maximal rate of IP_3 production by PLCbeta
O_N = 0.3/umolar/second  # Agonist binding rate
Omega_N = 0.5/second  # Maximal inactivation rate
K_KC = 0.5*umolar  # Ca^2+ affinity of PKC
zeta = 10  # Maximal reduction of receptor affinity by PKC

# --- Endogenous IP3 production
O_delta = 0.6*umolar/second  # Maximal rate of IP_3 production by PLCdelta
kappa_delta = 1.5*umolar  # Inhibition constant of PLC_delta by IP_3
K_delta = 0.1*umolar  # Ca^2+ affinity of PLCdelta

# --- IP_3 diffusion
```

(continues on next page)
\[ F = 2 \text{umolar/second} \]  # GJC IP_3 permeability
\[ I_{\Theta} = 0.3 \text{umolar} \]  # Threshold gradient for IP_3 diffusion
\[ \omega_I = 0.05 \text{umolar} \]  # Scaling factor of diffusion

--- IP_3 degradation
\[ \Omega_{5P} = 0.05/\text{second} \]  # Maximal rate of IP_3 degradation by IP-5P
\[ K_D = 0.7 \text{umolar} \]  # Ca^{2+} affinity of IP3-3K
\[ K_{3K} = 1.0 \text{umolar} \]  # IP_3 affinity of IP_3-3K
\[ \Omega_{3K} = 4.5 \text{umolar/second} \]  # Maximal rate of IP_3 degradation by IP_3-3K

--- IP_3 diffusion
\[ F_{\text{ex}} = 2.0 \text{umolar/second} \]  # Maximal exogenous IP3 flow
\[ I_{\Theta} = 0.3 \text{umolar} \]  # Threshold gradient for IP_3 diffusion
\[ \omega_I = 0.05 \text{umolar} \]  # Scaling factor of diffusion

--- Gliotransmitter release and time course
\[ C_{\Theta} = 0.5 \text{umolar} \]  # Ca^{2+} threshold for exocytosis
\[ \Omega_A = 0.6/\text{second} \]  # Gliotransmitter recycling rate
\[ U_A = 0.6 \]  # Gliotransmitter release probability
\[ G_T = 200 \text{mmolar} \]  # Total vesicular gliotransmitter concentration
\[ \rho_e = 6.5 \times 10^{-4} \]  # Astrocytic vesicle-to-extracellular volume ratio
\[ \Omega_e = 60/\text{second} \]  # Gliotransmitter clearance rate
\[ \alpha = 0.0 \]  # Gliotransmission nature

defaultclock.dt = sim_dt  # Set the integration time
rate_in = TimedArray([0.011, 0.11, 1.1, 11] * Hz, dt=5*second)
synapses = Synapses(source_neurons, target_neurons, 5.13. frompapers/Stimberg_et_al_2018
(continues on next page)
We create three synapses per connection: only the first two are modulated by the astrocyte however. Note that we could also create three synapses per connection with a single connect call by using connect(j='i', n=3), but this would create synapses arranged differently (synapses connection pairs (0, 0), (0, 0), (1, 1), (1, 1), (1, 1), ..., instead of connections (0, 0), (1, 1), ..., (0, 0), (1, 1), ..., (0, 0), (1, 1), ...). Making the later connection descriptions more complicated.

```python
synapses.connect(j='i')  # closed-loop modulation
synapses.connect(j='i')  # open modulation
synapses.connect(j='i')  # no modulation
```

```python
synapses.x_S = 1.0
```

### Astrocytes

The astrocyte emits gliotransmitter when its Ca^2+ concentration crosses a threshold.

```
astro_eqs = ```

# Fraction of activated astrocyte receptors:
\[
\frac{d\Gamma_A}{dt} = O_N \times Y_S \times (1 - \Gamma_A) - \Omega_N \times (1 + zeta \times C/(C + K_KC)) \times \Gamma_A : 1
\]

# IP_3 dynamics:
\[
\frac{dI}{dt} = J_beta + J_delta - J_3K - J_5P + J_ex : \text{mmolar/second}
\]

\[
J_beta = O_beta \times \Gamma_A : \text{mmolar/second}
\]

\[
J_delta = O_delta/(1 + I/kappa_delta) \times C^{2}/(C^{2} + K_delta^{2}) : \text{mmolar/second}
\]

\[
J_3K = O_3K \times C^{4}/(C^{4} + K_D^{4}) \times I/(I + K_3K) : \text{mmolar/second}
\]

\[
J_5P = Omega_5P*I : \text{mmolar/second}
\]

\[
delta_I_bias = I - I_bias : \text{mmolar}
\]

\[
J_ex = -F_ex/2 \times (1 + \tanh((abs(delta_I_bias) - I_Theta)/omega_I)) \times \text{sign(delta_I_bias)} : \text{mmolar/second}
\]

\[
I_bias : \text{mmolar (constant)}
\]

# Ca^2+-induced Ca^2+ release:
\[
\frac{dC}{dt} = (Omega_C \times m_inf \times \gamma_{2} \times h^{3} + Omega_L) \times (C_T - (1 + rho_A) \times C) - O_P \times C^{2}/(C^{2} + K_P^{2}) : \text{mmolar/second}
\]

\[
\frac{dh}{dt} = (h_{inf} - h)/tau_h : 1 \quad \# \text{IP3R de-inactivation probability}
\]

\[
m_{inf} = I/(I + d_1) \times C/(C + d_5) : 1
\]

\[
h_{inf} = Q_2/(Q_2 + \gamma_{2} + C) : 1
\]

\[
tau_h = 1/(O_2 \times (Q_2 + \gamma_{2} + C)) : \text{second}
\]

\[
Q_2 = d_2 \times (I + d_1)/(I + d_3) : \text{mmolar}
\]

# Fraction of gliotransmitter resources available for release
\[
\frac{dx_A}{dt} = Omega_A \times (1 - x_A) : 1
\]

# gliotransmitter concentration in the extracellular space
\[
\frac{dG_A}{dt} = -Omega_e \times G_A : \text{mmolar}
\]

# Neurotransmitter concentration in the extracellular space
\[
Y_S : \text{mmolar}
\]

```
glio_release = ```

\[
G_A += rho_e \times G_T \times U_A \times x_A
\]

\[
x_A -= U_A \times x_A
\]

astrocyte = NeuronGroup(N_astro*N_synapses, astro_eqs,
    # The following formulation makes sure that a "spike" is only triggered at the first threshold crossing
    ...)
threshold='C>C_Theta',
refractory='C>C_Theta',
# The gliotransmitter release happens when the threshold 
is crossed, in Brian terms it can therefore be 
# considered a "reset"
reset=glio_release,
method='rk4')

astrocyte.h = 0.9
astrocyte.x_A = 1.0
# Only the second group of N_synapses astrocytes are activated by external stimulation
astrocyte.I_bias = (np.r_[np.zeros(N_synapses), np.ones(N_synapses)])*1.0*umolar

## Connections
ecs_syn_to_astro = Synapses(synapses, astrocyte,
    'Y_S_post = Y_S_pre : mmolar (summed)')
ecs_syn_to_astro.connect(j='i if i < N_synapses')
ecs_astro_to_syn = Synapses(astrocyte, synapses,
    'G_A_post = G_A_pre : mmolar (summed)')
ecs_astro_to_syn.connect(j='i if i < N_synapses')
ecs_astro_to_syn.connect(j='i if i >= N_synapses and i < 2*N_synapses')

# Monitors
syn_mon = StateMonitor(synapses, 'Y_S',
    record=np.arange(N_synapses*(N_astro+1)), dt=10*ms)

# Simulation run
run(duration, report='text')

# Analysis and plotting
plt.style.use('figures.mplstyle')
fig, ax = plt.subplots(nrows=4, ncols=1, figsize=(3.07, 3.07*1.33),
    sharex=True,
    gridspec_kw={'height_ratios': [1, 3, 3, 3],
    'top': 0.98, 'bottom': 0.12,
    'left': 0.24, 'right': 0.95})
ax[0].semilogy(syn_mon.t/second, rate_in(syn_mon.t), '-', color='black')
ax[0].set(xlim=(0, duration/second), ylim=(0.01, 12),
yticks=[0.01, 0.1, 1, 10], ylabel=r'$\nu_{in}$ (Hz)')
pu.adjust_spines(ax[0], ['left'])

ax[1].plot(syn_mon.t/second,
    np.mean(syn_mon.Y_S[2*N_synapses:]/umolar, axis=0),
    '-', color='black')
ax[1].set(xlim=(0, duration/second), ylim=(-5, 260),
yticks=np.arange(0, 260, 50),
ylabel=r'$\langle Y_S \rangle$ ($\mu$M)')
pu.adjust_spines(ax[1], ['left'])

5.13. frompapers/Stimberg_et_al_2018
ax[1].legend(['no gliotransmission'], loc='upper left')
pu.adjust_spines(ax[1], ['left'])

ax[2].plot(syn_mon.t/second,
    np.mean(syn_mon.Y_S[N_synapses:2*N_synapses]/umolar, axis=0),
    '-', color='C2')
ax[2].set(xlim=(0, duration/second), ylim=(-3, 150),
    yticks=np.arange(0, 151, 25),
    ylabel=r'$\langle Y_S \rangle$ ($\mu$M)')
ax[2].legend(['open-loop gliotransmission'], loc='upper left')
pu.adjust_spines(ax[2], ['left'])

ax[3].plot(syn_mon.t/second,
    np.mean(syn_mon.Y_S[:N_synapses]/umolar, axis=0),
    '-', color='C3')
ax[3].set(xlim=(0, duration/second), ylim=(-2, 150),
    xticks=np.arange(0., duration/second+1, 5.0),
    yticks=np.arange(0, 151, 25),
    xlabel='time (s)', ylabel=r'$\langle Y_S \rangle$ ($\mu$M)')
ax[3].legend(['closed-loop gliotransmission'], loc='upper left')
pu.adjust_spines(ax[3], ['left', 'bottom'])
pu.adjust_ylabels(ax, x_offset=-0.22)

plt.show()
5.13.6 Example: example_5_astro_ring

Modeling neuron-glia interactions with the Brian 2 simulator Marcel Stimberg, Dan F. M. Goodman, Romain Brette, Maurizio De Pittà bioRxiv 198366; doi: https://doi.org/10.1101/198366

Figure 5: Astrocytes connected in a network.

Intercellular calcium wave propagation in a ring of 50 astrocytes connected by bidirectional gap junctions (see Goldberg et al., 2010)

```python
from brian2 import *

import plot_utils as pu

set_device('cpp_standalone', directory=None)  # Use fast "C++ standalone mode"

# Model parameters
### General parameters
duration = 4000*second  # Total simulation time
sim_dt = 50*ms  # Integrator/sampling step

### Astrocyte parameters
# --- Calcium fluxes
O_P = 0.9*umolar/second  # Maximal Ca^2+ uptake rate by SERCAs
K_P = 0.05 * umolar  # Ca^2+ affinity of SERCAs
C_T = 2*umolar  # Total cell free Ca^2+ content
rho_A = 0.18  # ER-to-cytoplasm volume ratio
Omega_C = 6/second  # Maximal rate of Ca^2+ release by IP_3Rs
Omega_L = 0.1/second  # Maximal rate of Ca^2+ leak from the ER

# --- IP_3R kinetics
d_1 = 0.13*umolar  # IP_3 binding affinity
d_2 = 1.05*umolar  # Ca^2+ inactivation dissociation constant
O_2 = 0.2/umolar/second  # IP_3R binding rate for Ca^2+ inhibition
d_3 = 0.9434*umolar  # IP_3 dissociation constant
d_5 = 0.08*umolar  # Ca^2+ activation dissociation constant

# --- IP_3 production
O_delta = 0.6*umolar/second  # Maximal rate of IP_3 production by PLCdelta
kappa_delta = 1.5* umolar  # Inhibition constant of PLC_delta by IP_3
K_delta = 0.1*umolar  # Ca^2+ affinity of PLCdelta

# --- IP_3 degradation
Omega_5P = 0.05/second  # Maximal rate of IP_3 degradation by IP-5P
K_D = 0.7*umolar  # Ca^2+ affinity of IP3-3K
K_3K = 1.0*umolar  # IP_3 affinity of IP_3-3K
O_3K = 4.5*umolar/second  # Maximal rate of IP_3 degradation by IP_3-3K

# --- IP_3 diffusion
F_ex = 0.09*umolar/second  # Maximal exogenous IP3 flow
F = 0.09*umolar/second  # GJC IP_3 permeability
L_Theta = 0.3*umolar  # Threshold gradient for IP_3 diffusion
omega_I = 0.05*umolar  # Scaling factor of diffusion

# Model definition
defaultclock.dt = sim_dt  # Set the integration time
```

(continues on next page)
### Astrocytes

astro_eqs = '

dI/dt = J_delta - J_3K - J_5P + J_ex + J_coupling : mmolar

\[ J_{\text{delta}} = \frac{O_{\text{delta}}}{1 + \frac{I}{\kappa_{\text{delta}}}} \times \frac{C^2}{C^2 + K_{\text{delta}}^2} : \text{mmolar/second} \]

\[ J_{3K} = O_{3K} \times \frac{C^4}{C^4 + K_D^4} \times \frac{I}{I + K_{3K}} : \text{mmolar/second} \]

\[ J_{5P} = \Omega_{5P} \times I : \text{mmolar/second} \]

# Exogenous stimulation (rectangular wave with period of 50s and duty factor 0.4)

\[ \text{stimulus} = \text{int}(t \% (50*\text{second})) < 20*\text{second} : 1 \]

\[ \text{delta}_I_{\text{bias}} = I - I_{\text{bias}} \times \text{stimulus} : \text{mmolar} \]

\[ J_{\text{ex}} = -\frac{F_{\text{ex}}}{2} \times (1 + \tanh((\text{abs(delta}_I_{\text{bias}}) - I_{\Theta})/\omega_{I})) \times \text{sign(delta}_I_{\text{bias}}) : \text{mmolar/second} \]

# Diffusion between astrocytes

\[ J_{\text{coupling}} : \text{mmolar/second} \]

# Ca^2+--induced Ca^2+ release:

\[ dC/dt = J_r + J_l - J_p : \text{mmolar} \]

\[ dh/dt = \frac{h_{\inf} - h}{\tau_h} : 1 \]

\[ J_r = (\Omega_C \times m_{\inf}^3 \times h^3) \times (C_T - (1 + \rho_A) \times C) : \text{mmolar/second} \]

\[ J_l = \Omega_L \times (C_T - (1 + \rho_A) \times C) : \text{mmolar/second} \]

\[ m_{\inf} = \frac{I}{I + d_1} \times \frac{C}{C + d_5} : 1 \]

\[ h_{\inf} = \frac{Q_2}{Q_2 + C} : 1 \]

\[ \tau_h = \frac{1}{\Omega_2 \times (Q_2 + C)} : \text{second} \]

\[ Q_2 = d_2 \times \frac{(I + d_1)/(I + d_3)}{d_2} : \text{mmolar} \]

# External IP_3 drive

\[ I_{\text{bias}} : \text{mmolar (constant)} \]

---

N_astro = 50 # Total number of astrocytes in the network

astrocytes = NeuronGroup(N_astro, astro_eqs, method='rk4')

# Asymmetric stimulation on the 50th cell to get some nice chaotic patterns

astrocytes.I_bias[N_astro//2] = 1.0*umolar

astrocytes.h = 0.9

# Diffusion between astrocytes

astro_to_astro_eqs = '

dC/dt = J_r + J_l - J_p : \text{mmolar}

dh/dt = \frac{h_{\inf} - h}{\tau_h} : 1

J_r = (\Omega_C \times m_{\inf}^3 \times h^3) \times (C_T - (1 + \rho_A) \times C) : \text{mmolar/second}

J_l = \Omega_L \times (C_T - (1 + \rho_A) \times C) : \text{mmolar/second}

m_{\inf} = \frac{I}{I + d_1} \times \frac{C}{C + d_5} : 1

h_{\inf} = \frac{Q_2}{Q_2 + C} : 1

\tau_h = \frac{1}{\Omega_2 \times (Q_2 + C)} : \text{second}

Q_2 = d_2 \times \frac{(I + d_1)/(I + d_3)}{d_2} : \text{mmolar}

# External IP_3 drive

I_{\text{bias}} : \text{mmolar (constant)}

---

-----------------------------

# Monitors

astro_mon = StateMonitor(astrocytes, variables=['C'], record=True)

# Simulation run

run(duration, report='text')
## Analysis and plotting

```python
# Use a consistent style and figure layout
plt.style.use('figures.mplstyle')

fig, ax = plt.subplots(nrows=1, ncols=1, figsize=(6.26894, 6.26894 * 0.66),
                      gridspec_kw={'left': 0.1, 'bottom': 0.12})

scaling = 1.2
step = 10

ax.plot(astro_mon.t/second,
        (astro_mon.C[0:N_astro//2-1].T/astro_mon.C.max() +
         np.arange(N_astro//2-1)*scaling), color='black')
ax.plot(astro_mon.t/second,
        (astro_mon.C[N_astro//2:].T/astro_mon.C.max() +
         np.arange(N_astro//2, N_astro)*scaling),
        color='black')
ax.plot(astro_mon.t/second,
         np.arange(N_astro//2-1, N_astro//2)*scaling),
        color='C0')

ax.set(xlim=(0., duration/second), ylim=(0, (N_astro+1.5)*scaling),
       xticks=np.arange(0., duration/second, 500), xlabel='time (s)',
       yticks=np.arange(0.5*scaling, (N_astro + 1.5)*scaling, step*scaling),
       yticklabels=[str(yt) for yt in np.arange(0, N_astro + 1, step)],
       ylabel='$C/C_{max}$ (cell index)')

plt.show()
```

5.13. frompapers/Stimberg_et_al_2018 335
5.13.7 Example: example_6_COBA_with_astro

Modeling neuron-glia interactions with the Brian 2 simulator Marcel Stimberg, Dan F. M. Goodman, Romain Brette, Maurizio De Pittà bioRxiv 198366; doi: https://doi.org/10.1101/198366

Figure 6: Recurrent neuron-glial network.

Randomly connected COBA network (see Brunel, 2000) with excitatory synapses modulated by release-increasing gliotransmission from a randomly connected network of astrocytes.

```python
from brian2 import *
import plot_utils as pu
set_device('cpp_standalone', directory=None)  # Use fast "C++ standalone mode"
seed(28371)  # to get identical figures for repeated runs

# Model parameters
# General parameters
N_e = 3200  # Number of excitatory neurons
N_i = 800  # Number of inhibitory neurons
N_a = 3200  # Number of astrocytes
size = 3.75*mmeter  # Length and width of the square lattice
distance = 50*umeter  # Distance between neurons

# Neuron parameters
E_l = -60*mV  # Leak reversal potential
g_l = 9.99*nS  # Leak conductance
E_e = 0*mV  # Excitatory synaptic reversal potential
E_i = -80*mV  # Inhibitory synaptic reversal potential
C_m = 198*pF  # Membrane capacitance
tau_e = 5*ms  # Excitatory synaptic time constant
tau_i = 10*ms  # Inhibitory synaptic time constant
tau_r = 5*ms  # Refractory period
I_ex = 100*pA  # External current
V_th = -50*mV  # Firing threshold
V_r = E_l  # Reset potential

# Synapse parameters
rho_c = 0.005  # Synaptic vesicle-to-extracellular space volume ratio
Y_T = 500.*mmolar  # Total vesicular neurotransmitter concentration
Omega_c = 40/second  # Neurotransmitter clearance rate
U_0_star = 0.6  # Resting synaptic release probability
Omega_f = 3.33/second  # Synaptic facilitation rate
Omega_d = 2.0/second  # Synaptic depression rate
w_e = 0.05*nS  # Excitatory synaptic conductance
w_i = 1.0*nS  # Inhibitory synaptic conductance
# Presynaptic receptors
O_G = 1.5/umolar/second  # Agonist binding (activating) rate
Omega_G = 0.5/(60*second)  # Agonist release (deactivating) rate

# Astrocyte parameters
# Calcium fluxes
```

(continues on next page)
O_P = 0.9*umolar/second # Maximal Ca^2+ uptake rate by SERCAs
K_P = 0.05*umolar # Ca^2+ affinity of SERCAs
C_T = 2*umolar # Total cell free Ca^2+ content
rho_A = 0.18 # ER-to-cytoplasm volume ratio
Omega_C = 6/second # Maximal rate of Ca^2+ release by IP_3Rs
Omega_L = 0.1/second # Maximal rate of Ca^2+ leak from the ER

# --- IP_3R kinetics
d_1 = 0.13*umolar # IP_3 binding affinity
d_2 = 1.05*umolar # Ca^2+ inactivation dissociation constant
G_2 = 0.2/umolar/second # IP_3R binding rate for Ca^2+ inhibition
d_3 = 0.9434*umolar # IP_3 dissociation constant
d_5 = 0.08*umolar # Ca^2+ activation dissociation constant

# --- IP_3 production
# --- Agonist-dependent IP_3 production
O_beta = 0.5*umolar/second # Maximal rate of IP_3 production by PLCbeta
O_N = 0.3/umolar/second # Agonist binding rate
Omega_N = 0.5/second # Maximal inactivation rate
K_KC = 0.5*umolar # Ca^2+ affinity of PKC
zeta = 10 # Maximal reduction of receptor affinity by PKC

# --- Endogenous IP3 production
G_delta = 1.2*umolar/second # Maximal rate of IP_3 production by PLCdelta
kappa_delta = 1.5*umolar # Inhibition constant of PLC_delta by IP_3
K_delta = 0.1*umolar # Ca^2+ affinity of PLCdelta

# --- IP_3 degradation
Omega_5P = 0.05/second # Maximal rate of IP_3 degradation by IP-5P
K_D = 0.7*umolar # Ca^2+ affinity of IP3-3K
K_3K = 1.0*umolar # IP_3 affinity of IP3-3K
G_3K = 4.5*umolar/second # Maximal rate of IP_3 degradation by IP_3-3K

# --- IP_3 diffusion
F = 0.09*umolar/second # GJC IP_3 permeability
I_Theta = 0.3*umolar # Threshold gradient for IP_3 diffusion
omega_I = 0.05*umolar # Scaling factor of diffusion

# --- Gliotransmitter release and time course
C_Theta = 0.5*umolar # Ca^2+ threshold for exocytosis
Omega_A = 0.6/second # Gliotransmitter recycling rate
U_A = 0.6 # Gliotransmitter release probability
G_T = 200*mmolar # Total vesicular gliotransmitter concentration
rho_e = 6.5e-4 # Astrocytic vesicle-to-extracellular volume ratio
Omega_e = 60/second # Gliotransmitter clearance rate
alpha = 0.0 # Gliotransmission nature

# Define HF stimulus
stimulus = TimedArray([[1.0, 1.2, 1.0, 1.0], dt=2*second])

dv/dt = (g_l*(E_l-v) + g_e*(E_e-v) + g_i*(E_i-v) + I_ex*stimulus(t))/C_m : volt

(continues on next page)
\[ \frac{dg_e}{dt} = -\frac{g_e}{\tau_e} : \text{siemens} \quad \# \text{post-synaptic excitatory conductance} \]

\[ \frac{dg_i}{dt} = -\frac{g_i}{\tau_i} : \text{siemens} \quad \# \text{post-synaptic inhibitory conductance} \]

Neuron position in space:
\[ x : \text{meter (constant)} \]
\[ y : \text{meter (constant)} \]

```
neurons = NeuronGroup(N_e + N_i, model=neuron_eqs,
    threshold='v>V_th', reset='v=V_r',
    refractory='tau_r', method='euler')
```

```
exc_neurons = neurons[:N_e]
inh_neurons = neurons[N_e:]
```

```
# Arrange excitatory neurons in a grid
N_rows = int(sqrt(N_e))
N_cols = N_e//N_rows
grid_dist = (size / N_cols)
exc_neurons.x = '(i // N_rows)*grid_dist - N_rows/2.0*grid_dist'
exc_neurons.y = '(i % N_rows)*grid_dist - N_cols/2.0*grid_dist'

# Random initial membrane potential values and conductances
neurons.v = 'E_l + rand()*(V_th-E_l)'
negons.g_e = 'rand()*(w_e)
negons.g_i = 'rand()*(w_i)
```

```
### Synapses

synapses_eqs = '''
# Neurotransmitter
\[ \frac{dY_S}{dt} = -\Omega_c \times Y_S : \text{mmolar (clock-driven)} \]

# Fraction of activated presynaptic receptors
\[ \frac{dGamma_S}{dt} = O_G \times G_A \times (1 - Gamma_S) - \Omega_G \times Gamma_S : 1 \quad \text{(clock-driven)} \]

# Usage of releasable neurotransmitter per single action potential:
\[ \frac{du_S}{dt} = -\Omega_f \times u_S : 1 \quad \text{(event-driven)} \]

# Fraction of synaptic neurotransmitter resources available for release:
\[ \frac{dx_S}{dt} = \Omega_d \times (1 - x_S) : 1 \quad \text{(event-driven)} \]

U_0 = 1

# released synaptic neurotransmitter resources:
\[ r_S : 1 \]

# gliotransmitter concentration in the extracellular space:
\[ G_A : \text{mmolar} \]

# which astrocyte covers this synapse?
astrocyte_index : integer (constant)
```

```
astrocyte_index = 0

synapses_action = '''
U_0 = (1 - Gamma_S) \times U_0__star + alpha \times Gamma_S
\[ u_S += U_0 \times (1 - u_S) \]
\[ r_S = u_S \times x_S \]
\[ x_S -= r_S \]
\[ Y_S += rho_c \times Y_T \times r_S \]
```

```
exc_syn = Synapses(exc_neurons, neurons, model=synapses_eqs,
    on_pre=synapses_action+'g_e_post += w_e*r_S',
    method='exact')
```

```
exc_syn.connect(True, p=0.05)
```

```
inhib_syn = Synapses(inh_neurons, neurons, model=synapses_eqs,
    on_pre=synapses_action+'g_i_post += w_i*r_S',
    method='exact')
```

```
inhib_syn.connect(True, p=0.2)
```

(continues on next page)
# Connect excitatory synapses to an astrocyte depending on the position of the
# post-synaptic neuron
N_rows_a = int(sqrt(N_a))
N_cols_a = N_a/N_rows_a
grid_dist = size / N_rows_a
exc_syn.astrocyte_index = ('int(x_post/grid_dist) + '
                           'N_cols_a*int(y_post/grid_dist)')

### Astrocytes

# The astrocyte emits gliotransmitter when its Ca^2+ concentration crosses
# a threshold
astro_eqs = '''
frac_active_A = O_N * Y_S * (1 - clip(Gamma_A,0,1)) -
               Omega_N*(1 + zeta * C/(C + K_KC)) * clip(Gamma_A,0,1)

# Intracellular IP_3
dI/dt = J_beta + J_delta - J_3K - J_5P + J_coupling : mmolar/second
J_beta = O_beta * Gamma_A : mmolar/second
J_delta = O_delta/(1 + I/kappa_delta) * C**2/(C**2 + K_delta**2) : mmolar/second
J_3K = O_3K * C**4/(C**4 + K_D**4) * I/(I + K_3K) : mmolar/second
J_5P = Omega_5P*I : mmolar/second

# Ca^2+-induced Ca^2+ release:
dC/dt = J_r + J_l - J_p : mmolar
J_r = (Omega_C * m_inf**3 * h**3) * (C_T - (1 + rho_A)*C) : mmolar/second
J_l = Omega_L * (C_T - (1 + rho_A)*C) : mmolar/second
J_p = O_P * C**2/(C**2 + K_P**2) : mmolar/second
m_inf = I/(I + d_1) * C/(C + d_5) : 1
h_inf = Q_2/(Q_2 + C) : 1
tau_h = 1/(O_2 * (Q_2 + C)) : second
Q_2 = d_2 * (I + d_1)/(I + d_3) : mmolar

# Fraction of gliotransmitter resources available for release:
dx_A/dt = Omega_A * (1 - x_A) : 1

# gliotransmitter concentration in the extracellular space:
dG_A/dt = -Omega_e*G_A : mmolar
# Neurotransmitter concentration in the extracellular space:
Y_S : mmolar

# The astrocyte position in space
x : meter (constant)
y : meter (constant)
```
glio_release = '''
G_A += rho_e * G_T * U_A * x_A
x_A -= U_A * x_A
```

astrocytes = NeuronGroup(N_a, astro_eqs,
    # The following formulation makes sure that a "spike" is
    # only triggered at the first threshold crossing
    threshold='C>C_Theta',
    refractory='C>C_Theta',
    # The gliotransmitter release happens when the threshold
    # is crossed, in Brian terms it can therefore be
    # considered a "reset"
    reset=glio_release,
method='rk4',
dt=1e-2*second)

# Arrange astrocytes in a grid
astrocytes.x = '(i // N_rows_a)*grid_dist - N_rows_a/2.0*grid_dist'
astrocytes.y = '(i % N_rows_a)*grid_dist - N_cols_a/2.0*grid_dist'

# Add random initialization
astrocytes.C = 0.01*umolar
astrocytes.h = 0.9
astrocytes.I = 0.01*umolar
astrocytes.x_A = 1.0

ecs_astro_to_syn = Synapses(astrocytes, exc_syn,
   'G_A_post = G_A_pre : mmolar (summed)')
ecs_astro_to_syn.connect('i == astrocyte_index_post')
ecs_syn_to_astro = Synapses(exc_syn, astrocytes,
   'Y_S_post = Y_S_pre/N_incoming : mmolar (summed)')
ecs_syn_to_astro.connect('astrocyte_index_pre == j')

# Diffusion between astrocytes
astro_to_astro_eqs = ''
  delta_I = I_post - I_pre : mmolar
  J_coupling_post = -(1 + tanh((abs(delta_I) - I_Theta)/omega_I))*
    sign(delta_I)*F/2 : mmolar/second (summed)
''
astro_to_astro = Synapses(astrocytes, astrocytes,
   model=astro_to_astro_eqs)
astro_to_astro.connect('i != j and '
  'sqrt((x_pre-x_post)**2 +
    (y_pre-y_post)**2) < 75*um')

# Monitors

# Note that we could use a single monitor for all neurons instead, but this
# way plotting is a bit easier in the end
exc_mon = SpikeMonitor(exc_neurons)
inh_mon = SpikeMonitor(inh_neurons)
ast_mon = SpikeMonitor(astrocytes)

# Simulation run
run(duration, report='text')

# Plot of Spiking activity
fig, ax = plt.subplots(nrows=3, ncols=1, sharex=True, figsize=(6.26894, 6.26894*0.8),
gridspec_kw={'height_ratios': [1, 6, 2],
    'left': 0.12, 'top': 0.97})
time_range = np.linspace(0, duration/second, duration/second*100)*second
ax[0].plot(time_range, I_ex*stimulus(time_range)/pA, 'k')
ax[0].set(xlim=(0, duration/second), ylim=(98, 122),
yticks=[100, 120], ylabel='$I_{ex}$ (pA)')
```
pu.adjust_spines(ax[0], ['left'])

## We only plot a fraction of the spikes
fraction = 4
ax[1].plot(exc_mon.t[exc_mon.i <= N_e//fraction]/second, 
           exc_mon.i[exc_mon.i <= N_e//fraction], '|', color='C0')
ax[1].plot(inh_mon.t[inh_mon.i <= N_i//fraction]/second, 
           inh_mon.i[inh_mon.i <= N_i//fraction]+N_e//fraction, '|', color='C1')
ax[1].plot(ast_mon.t[ast_mon.i <= N_a//fraction]/second, 
           ast_mon.i[ast_mon.i <= N_a//fraction]+(N_e+N_i)//fraction, 
           '|', color='C2')
ax[1].set(xlim=(0, duration/second), ylim=[0,(N_e+N_i+N_a)//fraction], 
          yticks=np.arange(0, (N_e+N_i+N_a)//fraction+1, 250),
          ylabel='cell index')
pu.adjust_spines(ax[1], ['left'])

# Generate frequencies
bin_size = 1*ms
spk_count, bin_edges = np.histogram(np.r_[exc_mon.t/second, inh_mon.t/second],
                                     int(duration/bin_size))
rate = 1.0*spk_count/(N_e + N_i)/bin_size/Hz
rate[rate<0.001] = 0.001 # Fix 0 lower bound for log scale
ax[2].semilogy(bin_edges[:-1], rate, '-', color='k')
pu.adjust_spines(ax[2], ['left', 'bottom'])
ax[2].set(xlim=(0, duration/second), ylim=(0.1, 150),
          xticks=np.arange(0,9), yticks=[0.1, 1, 10, 100],
          xlabel='time (s)', ylabel='rate (Hz)')
ax[2].get_yaxis().set_major_formatter(ScalarFormatter())
pu.adjust_ylabels(ax, x_offset=-0.11)
plt.show()
```
Module with useful functions for making publication-ready plots.

```python
def adjust_spines(ax, spines, position=5, smart_bounds=False):
    r""
    Set custom visibility and position of axes
    
    ax : Axes
    Axes handle
    spines : List
    String list of 'left', 'bottom', 'right', 'top' spines to show
    position : Integer
    Number of points for position of axis
    r""
    for loc, spine in ax.spines.items():
        if loc in spines:
            spine.set_position(('outward', position))
            spine.set_smart_bounds(smart_bounds)
        else:
            spine.set_color('none') # don't draw spine
```

(continues on next page)
# turn off ticks where there is no spine
if 'left' in spines:
    ax.yaxis.set_ticks_position('left')
elif 'right' in spines:
    ax.yaxis.set_ticks_position('right')
else:
    # no yaxis ticks
    ax.yaxis.set_ticks([])
    ax.tick_params(axis='y', which='both', left='off', right='off')

if 'bottom' in spines:
    ax.xaxis.set_ticks_position('bottom')
elif 'top' in spines:
    ax.xaxis.set_ticks_position('top')
else:
    # no xaxis ticks
    ax.xaxis.set_ticks([])
    ax.tick_params(axis='x', which='both', bottom='off', top='off')


def adjust_ylabels(ax, x_offset=0):
    
    Scan all ax list and identify the outmost y-axis position. Setting all the labels to that position + x_offset.
    
    xc = 0.0
    for a in ax:
        xc = min(xc, (a.yaxis.get_label()).get_position()[0])

    for a in ax:
        a.yaxis.set_label_coords(xc + x_offset,
                                 (a.yaxis.get_label()).get_position()[1])


5.14 standalone

5.14.1 Example: STDP_standalone

Spike-timing dependent plasticity. Adapted from Song, Miller and Abbott (2000) and Song and Abbott (2001).
This example is modified from synapses_STDP.py and writes a standalone C++ project in the directory STDP_standalone.

```python
from brian2 import *
set_device('cpp_standalone', directory='STDP_standalone')

N = 1000
taum = 10*ms
taupre = 20*ms
taupost = taupre
```

(continues on next page)
\[E_e = 0 \text{mV}\]
\[v_t = -54 \text{mV}\]
\[v_r = -60 \text{mV}\]
\[E_l = -74 \text{mV}\]
\[\tau_e = 5 \text{ms}\]
\[F = 15 \text{Hz}\]
\[g_{\text{max}} = .01\]
\[dA_{\text{pre}} = .01\]
\[dA_{\text{post}} = \frac{-dA_{\text{pre}}}{\tau_{\text{pre}}/\tau_{\text{post}} \times 1.05}\]
\[dA_{\text{post}} = g_{\text{max}}\]
\[dA_{\text{pre}} = g_{\text{max}}\]

eqs\_neurons = '':'
\[\frac{dv}{dt} = \frac{ge \times (E_e-v_r) + E_l - v}{\tau_{\text{m}}} : \text{volt}\]
\[\frac{dge}{dt} = -ge / \tau_{\text{e}} : 1\]
''

input = PoissonGroup(N, rates=F)
neurons = NeuronGroup(1, eqs\_neurons, threshold='v>v_t', reset='v=v_r',
method='\text{exact}')
S = Synapses(input, neurons,
  
  \[\frac{dA_{\text{pre}}}{dt} = \frac{-A_{\text{pre}}}{\tau_{\text{pre}}} : 1 \ (\text{event-driven})\]
  \[\frac{dA_{\text{post}}}{dt} = \frac{-A_{\text{post}}}{\tau_{\text{post}}} : 1 \ (\text{event-driven})\]
  ,
  on\_pre='''ge += w
        A_{\text{pre}} += dA_{\text{pre}}
        w = \text{clip}(w + A_{\text{post}}, 0, g_{\text{max}})''',
  on\_post='''A_{\text{post}} += dA_{\text{post}}
        w = \text{clip}(w + A_{\text{pre}}, 0, g_{\text{max}})'''
  )
S.connect()
S.w = 'rand() \times g_{\text{max}}'
mon = StateMonitor(S, 'w', record=[0, 1])
s_mon = SpikeMonitor(input)
run(100*\text{second}, \text{report}='\text{text}')

subplot(311)
plot(S.w / g_{\text{max}}, '.k')
ylabel('Weight / g_{\text{max}}')
xlabel('Synapse index')
subplot(312)
hist(S.w / g_{\text{max}}, 20)
xlabel('Weight / g_{\text{max}}')
subplot(313)
plot(mon.t/\text{second}, mon.w.T/g_{\text{max}})
xlabel('Time (s)')
ylabel('Weight / g_{\text{max}}')
tight\_layout()
show()
5.14.2 Example: cuba_openmp

Run the cuba.py example with OpenMP threads.

```
from brian2 import *

set_device('cpp_standalone', directory='CUBA')
prefs.devices.cpp_standalone.openmp_threads = 4

taum = 20*ms
taue = 5*ms
taui = 10*ms
Vt = -50*mV
Vr = -60*mV
El = -49*mV

eqs = ''
dv/dt = (ge+gi-(v-El))/taum : volt (unless refractory)
dge/dt = -ge/taue : volt (unless refractory)
dgi/dt = -gi/taui : volt (unless refractory)
''
```

(continues on next page)
P = NeuronGroup(4000, eqs, threshold='v>Vt', reset='v = Vr', refractory=5*ms, method='exact')
P.v = 'Vr + rand() * (Vt - Vr)'
P.ge = 0*mV
P.gi = 0*mV
we = (60*0.27/10)*mV  # excitatory synaptic weight (voltage)
wi = (-20*4.5/10)*mV  # inhibitory synaptic weight
Ce = Synapses(P, P, on_pre='ge += we')
Ci = Synapses(P, P, on_pre='gi += wi')
Ce.connect('i<3200', p=0.02)
Ci.connect('i>=3200', p=0.02)
s_mon = SpikeMonitor(P)
run(1 * second)
plot(s_mon.t/ms, s_mon.i, ',k')
xlabel('Time (ms)')
ylabel('Neuron index')
show()
5.15 synapses

5.15.1 Example: STDP

Spike-timing dependent plasticity

Adapted from Song, Miller and Abbott (2000) and Song and Abbott (2001)

```python
from brian2 import *

N = 1000
taum = 10*ms
taupre = 20*ms
taupost = taupre
Ee = 0*mV
vt = -54*mV
vr = -60*mV
E1 = -74*mV
taue = 5*ms
F = 15*Hz
gmax = .01
dApre = .01
dApost = -dApre * taupre / taupost * 1.05
dApost *= gmax
dApre *= gmax

eqs_neurons = '''
    dv/dt = (ge * (Ee-vr) + El - v) / taum : volt
    dge/dt = -ge / taue : 1
'''

input = PoissonGroup(N, rates=F)
neurons = NeuronGroup(1, eqs_neurons, threshold='v>vt', reset='v = vr',
    method='exact')
S = Synapses(input, neurons,
    '''
    w : 1
    dApre/dt = -Apre / taupre : 1 (event-driven)
    dApost/dt = -Apost / taupost : 1 (event-driven)''',
    on_pre='''
        ge += w
        Apre += dApre
        w = clip(w + Apost, 0, gmax)''',
    on_post='''
        Apost += dApost
        w = clip(w + Apre, 0, gmax)''',
)
S.connect()
S.w = 'rand() * gmax'
mon = StateMonitor(S, 'w', record=[0, 1])
s_mon = SpikeMonitor(input)
run(100*second, report='text')

subplot(311)
plot(S.w / gmax, '.k')
ylabel('Weight / gmax')
xlabel('Synapse index')
```

(continues on next page)
5.15.2 Example: efficient_gaussian_connectivity

An example of turning an expensive `Synapses.connect` operation into three cheap ones using a mathematical trick.

Consider the connection probability between neurons i and j given by the Gaussian function $p = e^{-\alpha(i-j)^2}$ (for some constant $\alpha$). If we want to connect neurons with this probability, we can very simply do:

```python
S.connect(p='exp(-alpha*(i-j)*2)')
```

However, this has a problem. Although we know that this will create $O(N)$ synapses if $N$ is the number of neurons, because we have specified $p$ as a function of $i$ and $j$, we have to evaluate $p(i, j)$ for every pair $(i, j)$, and
therefore it takes $O(N^2)$ operations.

Our first option is to take a cutoff, and say that if $p < q$ for some small $q$, then we assume that $p \approx 0$. We can work out which $j$ values are compatible with a given value of $i$ by solving $e^{-\alpha(i-j)^2} < q$ which gives $|i-j| < \sqrt{-\log(q)/\alpha} = w$. Now we implement the rule using the generator syntax to only search for values between $i-w$ and $i+w$, except that some of these values will be outside the valid range of values for $j$ so we set skip_if_invalid=True. The connection code is then:

```python
S.connect(j='k for k in range(i-w, i+w) if rand()<exp(-alpha*(i-j)**2)',
           skip_if_invalid=True)
```

This is a lot faster (see graph labelled “Limited” for this algorithm).

However, it may be a problem that we have to specify a cutoff and so we will lose some synapses doing this: it won’t be mathematically exact. This isn’t a problem for the Gaussian because $w$ grows very slowly with the cutoff probability $q$, but for other probability distributions with more weight in the tails, it could be an issue.

If we want to be exact, we can still do a big improvement. For the case $i-w \leq j \leq i+w$ we use the same connection code, but we also handle the case $|i-j| > w$. This time, we note that we want to create a synapse with probability $p(i-j)$ and we can rewrite this as $p(i-j)/p(w) \cdot p(w)$. If $|i-j| > w$ then this is a product of two probabilities $p(i-j)/p(w)$ and $p(w)$. So in the region $|i-j| > w$ a synapse will be created if two random events both occur, with these two probabilities. This might seem a little strange until you notice that one of the two probabilities $p(w)$ doesn’t depend on $i$ or $j$. This lets us use the much more efficient sample algorithm to generate a set of candidate $j$ values, and then add the additional test $\text{rand()} < p(i-j)/p(w)$. Here’s the code for that:

```python
w = int(ceil(sqrt(log(q)/\alpha)))
S.connect(j='k for k in range(i-w, i+w) if rand()<exp(-alpha*(i-j)**2)',
           skip_if_invalid=True)
pmax = exp(-0.1*w**2)
S.connect(j='k for k in sample(0, i-w, p=pmax) if rand()<exp(-alpha*(i-j)**2)/pmax',
           skip_if_invalid=True)
S.connect(j='k for k in sample(i+w, N_post, p=pmax) if rand()<exp(-alpha*(i-j)**2)/pmax',
           skip_if_invalid=True)
```

This “Divided” method is also much faster than the naive method, and is mathematically correct. Note though that this method is still $O(N^2)$ but the constants are much, much smaller and this will usually be sufficient. It is possible to take the ideas developed here even further and get even better scaling, but in most cases it’s unlikely to be worth the effort.

The code below shows these examples written out, along with some timing code and plots for different values of $N$.

```python
from brian2 import *
import time

def naive(N):
    G = NeuronGroup(N, 'v:1', threshold='v>1', name='G')
    S = Synapses(G, G, on_pre='v += 1', name='S')
    S.connect(p='exp(-0.1*(i-j)**2)')

def limited(N, q=0.001):
    G = NeuronGroup(N, 'v:1', threshold='v>1', name='G')
    S = Synapses(G, G, on_pre='v += 1', name='S')
    w = int(ceil(sqrt(log(q)/\alpha)))
    S.connect(j='k for k in range(i-w, i+w) if rand()<exp(-alpha*(i-j)**2)',
           skip_if_invalid=True)

def divided(N, q=0.001):
```

(continues on next page)
G = NeuronGroup(N, 'v:1', threshold='v>1', name='G')
S = Synapses(G, G, on_pre='v += 1', name='S')

w = int(ceil(sqrt(log(q)/-0.1)))
S.connect (j='k for k in range(i-w, i+w) if rand()<exp(-0.1*(i-j)**2)', skip_if_invalid=True)
pmax = exp(-0.1*w**2)
S.connect (j='k for k in sample(0, i-w, p=pmax) if rand()<exp(-0.1*(i-j)**2)/pmax', skip_if_invalid=True)
S.connect (j='k for k in sample(i+w, N_post, p=pmax) if rand()<exp(-0.1*(i-j)**2)/pmax', skip_if_invalid=True)

def repeated_run(f, N, repeats):
    start_time = time.time()
    for _ in range(repeats):
        f(N)
    end_time = time.time()
    return (end_time-start_time)/repeats

N = array([100, 500, 1000, 5000, 10000, 20000])
repeats = array([100, 10, 10, 1, 1, 1])*3
naive(10)
limited(10)
divided(10)

print 'Starting naive'
loglog(N, [repeated_run(naive, n, r) for n, r in zip(N, repeats)],
       label='Naive', lw=2)
print 'Starting limit'
loglog(N, [repeated_run(limited, n, r) for n, r in zip(N, repeats)],
       label='Limited', lw=2)
print 'Starting divided'
loglog(N, [repeated_run(divided, n, r) for n, r in zip(N, repeats)],
       label='Divided', lw=2)
xlabel('N')
ylabel('Time (s)')
legend(loc='best', frameon=False)
show()
5.15.3 Example: gapjunctions

Neurons with gap junctions.

```python
from brian2 import *

n = 10
v0 = 1.05
tau = 10*ms

eqs = '''
dv/dt = (v0 - v + Igap) / tau : 1
Igap : 1 # gap junction current '''

neurons = NeuronGroup(n, eqs, threshold='v > 1', reset='v = 0',
                       method='exact')
neurons.v = 'i * 1.0 / (n-1)'
trace = StateMonitor(neurons, 'v', record=[0, 5])
S = Synapses(neurons, neurons, '''
             w : 1 # gap junction conductance
             (continues on next page)
```
from brian2 import *

5.15.4 Example: jeffress

Jeffress model, adapted with spiking neuron models. A sound source (white noise) is moving around the head. Delay differences between the two ears are used to determine the azimuth of the source. Delays are mapped to a neural place code using delay lines (each neuron receives input from both ears, with different delays).
defaultclock.dt = .02*ms

# Sound
sound = TimedArray(10 * randn(50000), dt=defaultclock.dt) # white noise

# Ears and sound motion around the head (constant angular speed)
sound_speed = 300*metre/second
interaural_distance = 20*cm # big head!
max_delay = interaural_distance / sound_speed
print("Maximum interaural delay: %s" % max_delay)
angular_speed = 2 * pi / second # 1 turn/second
tau_ear = 1*ms
sigma_ear = .1
eqs_ears = '''
dx/dt = (sound(t-delay)-x)/tau_ear+sigma_ear*(2./tau_ear)**.5*xi : 1 #unless
\rightarrow refractory
delay = distance*sin(theta) : second
distance : second # distance to the centre of the head in time units
dtheta/dt = angular_speed : radian
'''
ears = NeuronGroup(2, eqs_ears, threshold='x>1', reset='x = 0',
refractory=2.5*ms, name='ears', method='euler')
ears.distance = [-.5 * max_delay, .5 * max_delay]
traces = StateMonitor(ears, 'delay', record=True)

# Coincidence detectors
num_neurons = 30
tau = 1*ms
sigma = .1
eqs_neurons = '''
dv/dt = -v / tau + sigma * (2 / tau)**.5 * xi : 1
'''
neurons = NeuronGroup(num_neurons, eqs_neurons, threshold='v>1',
reset='v = 0', name='neurons', method='euler')
synapses = Synapses(ears, neurons, on_pre='v += .5')
synapses.connect()
synapses.delay['i==0'] = '(1.0*j)/(num_neurons-1)*1.1*max_delay'
synapses.delay['i==1'] = '(1.0*(num_neurons-j-1))/(num_neurons-1)*1.1*max_delay'
spikes = SpikeMonitor(neurons)
run(1000*ms)

# Plot the results
i, t = spikes.it
subplot(2, 1, 1)
plot(t/ms, i, '.
xlabel('Time (ms)')
ylabel('Neuron index')
xlim(0, 1000)
subplot(2, 1, 2)
plot(traces.t/ms, traces.delay.T/ms)
xlabel('Time (ms)')
ylabel('Input delay (ms)')
xlim(0, 1000)
tight_layout()
5.15.5 Example: licklider

Spike-based adaptation of Licklider’s model of pitch processing (autocorrelation with delay lines) with phase locking.

```python
from brian2 import *

defaulClock.dt = .02 * ms

# Ear and sound
max_delay = 20*ms # 50 Hz
tau_ear = 1*ms
sigma_ear = 0.0
eqs_ear = ''
  dx/dt = (sound-x)/tau_ear+0.1*(2./tau_ear)**.5*xi : 1 (unless refractory)
sound = 5*sin(2*pi*frequency*t)**3 : 1 # nonlinear distortion
#sound = 5*(sin(4*pi*frequency*t)+.5*sin(6*pi*frequency*t)) : 1 # missing fundamental
frequency = (200+200*t*Hz)*Hz # Hz # increasing pitch
  ...
receptors = NeuronGroup(2, eqs_ear, threshold='x>1', reset='x=0',
```

(continues on next page)
refractory=2*ms, method='euler')

# Coincidence detectors
min_freq = 50*Hz
max_freq = 1000*Hz
num_neurons = 300
tau = 1*ms
sigma = .1
eqs_neurons = '''
dv/dt = -v/tau+sigma*(2./tau)**.5*xi : 1
'''

neurons = NeuronGroup(num_neurons, eqs_neurons, threshold='v>1', reset='v=0', method='euler')
synapses = Synapses(receptors, neurons, on_pre='v += 0.5')
synapses.connect()
synapses.delay = 'i*1.0/exp(log(min_freq/Hz)+(j*1.0/(num_neurons-1))*log(max_freq/min_freq))*second'

spikes = SpikeMonitor(neurons)
run(500*ms)
plot(spikes.t/ms, spikes.i, '.k')
xlabel('Time (ms)')
ylabel('Frequency')
yticks([0, 99, 199, 299], array(1. / synapses.delay[i, [0, 99, 199, 299]], dtype=int))
show()
5.15.6 Example: nonlinear

NMDA synapses.

```python
from brian2 import *

a = 1 / (10*ms)
b = 1 / (10*ms)
c = 1 / (10*ms)

input = NeuronGroup(2, 'dv/dt = 1/(10*ms) : 1', threshold='v>1', reset='v = 0', method='euler')
neurons = NeuronGroup(1, '''dv/dt = (g-v)/(10*ms) : 1
g : 1''', method='exact')

S = Synapses(input, neurons,'''
dg_syn/dt = -a*g_syn+b*x*(1-g_syn) : 1 (clock-driven)
g_post = g_syn : 1 (summed)
dx/dt=-c*x : 1 (clock-driven)
w : 1 # synaptic weight
''', on_pre='x += w') # NMDA synapses
S.connect()
```

(continues on next page)
S.w = [1., 10.]
input.v = [0., 0.5]

M = StateMonitor(S, 'g',
    # If not using standalone mode, this could also simply be
    # record=True
    record=np.arange(len(input)*len(neurons)))
Mn = StateMonitor(neurons, 'g', record=0)

run(1000*ms)

subplot(2, 1, 1)
plot(M.t/ms, M.g.T)
xlabel('Time (ms)')
ylabel('g_syn')

subplot(2, 1, 2)
plot(Mn.t/ms, Mn[0].g)
ylabel('Time (ms)')
ylabel('g')
tight_layout()
show()
5.15.7 Example: spatial_connections

A simple example showing how string expressions can be used to implement spatial (deterministic or stochastic) connection patterns.

```python
from brian2 import *

rows, cols = 20, 20
G = NeuronGroup(rows * cols, '''x : meter
    y : meter''')

# initialize the grid positions
grid_dist = 25*umeter
G.x = '(i // rows) * grid_dist - rows/2.0 * grid_dist'
G.y = '(i % rows) * grid_dist - cols/2.0 * grid_dist'

# Deterministic connections
distance = 120*umeter
S_deterministic = Synapses(G, G)
S_deterministic.connect('sqrt((x_pre - x_post)**2 + (y_pre - y_post)**2) < distance')

# Random connections (no self-connections)
S_stochastic = Synapses(G, G)
S_stochastic.connect('i != j',
    p='1.5 * exp(-((x_pre-x_post)**2 + (y_pre-y_post)**2)/
    --> (2*(60*umeter)**2))')

figure(figsize=(12, 6))
# Show the connections for some neurons in different colors
for color in ['g', 'b', 'm']:
    subplot(1, 2, 1)
    neuron_idx = np.random.randint(0, rows*cols)
    plot(G.x[S_deterministic.j[neuron_idx, :]] / umeter,
     G.y[S_deterministic.j[neuron_idx, :]] / umeter, color + '.')
    subplot(1, 2, 2)
    plot(G.x[S_stochastic.j[neuron_idx, :]] / umeter,
     G.y[S_stochastic.j[neuron_idx, :]] / umeter, color + '.')

for idx, t in enumerate(['deterministic connections',
    'random connections']):
    subplot(1, 2, idx + 1)
    xlim((-rows/2.0 * grid_dist) / umeter, (rows/2.0 * grid_dist) / umeter)
    ylim((-cols/2.0 * grid_dist) / umeter, (cols/2.0 * grid_dist) / umeter)
title(t)
xlabel('x')
ylabel('y', rotation='horizontal')
axis('equal')

tight_layout()
show()
```
5.15.8 Example: state_variables

Set state variable values with a string (using code generation).

```python
from brian2 import *

G = NeuronGroup(100, 'v:volt', threshold='v>-50*mV')
G.v = '(sin(2*pi*i/N) - 70 + 0.25*randn()) * mV'
S = Synapses(G, G, 'w : volt', on_pre='v += w')
S.connect()

space_constant = 200.0
S.w['i > j'] = 'exp(-(i - j)**2/space_constant) * mV'

# Generate a matrix for display
w_matrix = np.zeros((len(G), len(G)))
w_matrix[S.i[:,], S.j[:,]] = S.w[:,]

subplot(1, 2, 1)
plot(G.v[:,] / mV)
xlabel('Neuron index')
ylabel('v')
subplot(1, 2, 2)
imshow(w_matrix)
xlabel('i')
ylabel('j')
title('Synaptic weight')
tight_layout()
show()
```
5.15.9 Example: synapses

A simple example of using *Synapses*.

```python
from brian2 import *

G1 = NeuronGroup(10, 'dv/dt = -v / (10*ms) : 1,
                  threshold='v > 1', reset='v=0.', method='exact')
G1.v = 1.2
G2 = NeuronGroup(10, 'dv/dt = -v / (10*ms) : 1,
                  threshold='v > 1', reset='v=0', method='exact')
syn = Synapses(G1, G2, 'dw/dt = -w / (50*ms): 1 (event-driven)', on_pre='v += w')
syn.connect('i == j', p=0.75)
# Set the delays
syn.delay = '1*ms + i*ms + 0.25*ms * randn()'
# Set the initial values of the synaptic variable
syn.w = 1
mon = StateMonitor(G2, 'v', record=True)
```

(continues on next page)
```python
run(20*ms)
plot(mon.t/ms, mon.v.T)
xlabel('Time (ms)')
ylabel('v')
show()
```
CHAPTER 6

brian2 package

Brian 2
Functions

\texttt{check\_cache}(\texttt{target})

6.1 \texttt{check\_cache} function

\texttt{(Shortest import: from brian2.\_\_init\_\_ import check\_cache)}

\texttt{brian2.\_\_init\_\_.check\_cache(\texttt{target})}

\texttt{clear\_cache}(\texttt{target})

Cleans the on-disk cache with the compiled files for a given code generation target.

6.2 \texttt{clear\_cache} function

\texttt{(Shortest import: from brian2.\_\_init\_\_ import clear\_cache)}

\texttt{brian2.\_\_init\_\_.clear\_cache(\texttt{target})}

Cleans the on-disk cache with the compiled files for a given code generation target.

\textbf{Parameters}

\texttt{target} : \texttt{str}

The code generation target (e.g. \texttt{'weave'} or \texttt{'cython'})

\textbf{Raises}

\texttt{ValueError} If the given code generation target does not have an on-disk cache
**IOError** If the cache directory contains unexpected files, suggesting that deleting it would also delete files unrelated to the cache.

### 6.3 hears module

This is only a bridge for using Brian 1 hears with Brian 2.

Deprecated since version 2.2.2.2: Use the `brian2hears` package instead.

**NOTES:**

- Slicing sounds with Brian 2 units doesn’t work, you need to either use Brian 1 units or replace calls to `sound[:20*ms]` with `sound.slice(None, 20*ms).etc.`

**TODO:** handle properties (e.g. `sound.duration`)

Not working examples:

- `time_varying_filter1` (care with units)

**Exported members:** `convert_unit_b1_to_b2`, `convert_unit_b2_to_b1`

**Classes**

<table>
<thead>
<tr>
<th>BridgeSound</th>
<th>We add a new method <code>slice</code> because slicing with units can’t work with Brian 2 units.</th>
</tr>
</thead>
</table>

#### 6.3.1 BridgeSound class

*(Shortest import: from brian2.hears import BridgeSound)*

**class** `brian2.hears.BridgeSound`

**Bases:** `brian2.hears.new_class`

We add a new method `slice` because slicing with units can’t work with Brian 2 units.

**Methods**

- `slice(*args)`

**Details**

- `slice(*args)`

`FilterbankGroup`(*filterbank, targetvar, ...*)

**Methods**

#### 6.3.2 FilterbankGroup class

*(Shortest import: from brian2.hears import FilterbankGroup)*
class brian2.hears.FilterbankGroup(filterbank, targetvar, *args, **kwds)
Bases: brian2.groups.neurongroup.NeuronGroup

Methods

reinit()

Details

reinit()

Sound alias of brian2.hears.BridgeSound

6.3.3 Sound class

(Shortest import: from brian2.hears import Sound)
brian2.hears.Sound
   alias of brian2.hears.BridgeSound

WrappedSound alias of brian2.hears.wrap_units_class.<locals>.new_class

6.3.4 WrappedSound class

(Shortest import: from brian2.hears import WrappedSound)
brian2.hears.WrappedSound
   alias of brian2.hears.wrap_units_class.<locals>.new_class

Functions

convert_unit_b1_to_b2(val)

6.3.5 convert_unit_b1_to_b2 function

(Shortest import: from brian2.hears import convert_unit_b1_to_b2)
brian2.hears.convert_unit_b1_to_b2(val)

convert_unit_b2_to_b1(val)

6.3.6 convert_unit_b2_to_b1 function

(Shortest import: from brian2.hears import convert_unit_b2_to_b1)
brian2.hears.convert_unit_b2_to_b1(val)
**6.3.7 modify_arg function**

(Shortest import: from brian2.hears import modify_arg)

brian2.hears.modify_arg(arg)

Modify arguments to make them compatible with Brian 1.
- Arrays of units are replaced with straight arrays
- Single values are replaced with Brian 1 equivalents
- Slices are handled so we can use e.g. sound[:20*ms]

The second part was necessary because some functions/classes test if an object is an array or not to see if it is a sequence, but because brian2.Quantity derives from ndarray this was causing problems.

**6.3.8 wrap_units function**

(Shortest import: from brian2.hears import wrap_units)

brian2.hears.wrap_units(f)

Wrap a function to convert units into a form that Brian 1 can handle. Also, check the output argument, if it is a b1h.Sound wrap it.

**6.3.9 wrap_units_class function**

(Shortest import: from brian2.hears import wrap_units_class)

brian2.hears.wrap_units_class(_C)

Wrap a class to convert units into a form that Brian 1 can handle in all methods

**6.3.10 wrap_units_property function**

(Shortest import: from brian2.hears import wrap_units_property)

brian2.hears.wrap_units_property(p)
6.4 numpy_ module

A dummy package to allow importing numpy and the unit-aware replacements of numpy functions without having to know which functions are overwritten.

This can be used for example as `import brian2.numpy_ as np`

Exported members: `ModuleDeprecationWarning, VisibleDeprecationWarning, __version__, show_config(), char, rec, memmap, newaxis, ndarray, flatiter, nditer, nested_iters, ufunc, arange(), array, zeros, count_nonzero(), empty, broadcast, dtype, fromstring, fromfile, frombuffer, int_asbuffer, where() ... (628 more members)`

6.5 only module

A dummy package to allow wildcard import from brian2 without also importing the pylab (numpy + matplotlib) namespace.

Usage: `from brian2.only import *`

Exported members: `get_logger(), BrianLogger, std_silent, Trackable, Nameable, SpikeSource, linked_var(), DEFAULT_FUNCTIONS, Function, implementation(), declare_types(), PreferenceError, BrianPreference, prefs, brian_prefs, Clock, defaultclock, Equations, Expression, Statements, BrianObject, BrianObjectException, Network, profiling_summary(), scheduling_summary() ... (304 more members)`

Functions

| `restore_initial_state()` | Restores internal Brian variables to the state they are in when Brian is imported |

6.5.1 restore_initial_state function

(Shortest import: `from brian2 import restore_initial_state`)

`brian2.only.restore_initial_state()`

Restores internal Brian variables to the state they are in when Brian is imported

- Resets `defaultclock.dt = 0.1*ms`, `BrianGlobalPreferences._restore` preferences, and set `BrianObject._scope_current_key` back to 0.

6.6 Subpackages

6.6.1 codegen package

Package providing the code generation framework.

Exported members: `NumpyCodeObject, WeaveCodeObject, CythonCodeObject`
**codeobject module**

Module providing the base `CodeObject` and related functions.

**Exported members:** `CodeObject, constant_or_scalar`

**Classes**

`CodeObject(owner, code, variables, ...[, name])` Executable code object.

### CodeObject class

*(Shortest import: from brian2.codegen.codeobject import CodeObject)*

```python
class brian2.codegen.codeobject.CodeObject(  
    owner, code, variables,  
    variable_indices, template_name, template_source, compiler_kwds,  
    name='codeobject*'
)
```

**Bases:** `brian2.core.names.Nameable`

Executable code object.

The code can either be a string or a `brian2.codegen.templates.MultiTemplate`.

After initialisation, the code is compiled with the given namespace using `code.compile(namespace)`.

Calling `code(key1=val1, key2=val2)` executes the code with the given variables inserted into the namespace.

**Attributes**

| class_name | A short name for this type of `CodeObject` |
| generator_class | The `CodeGenerator` class used by this `CodeObject` |

**Methods**

- `__call__(**kwds)` Call self as a function.
- `compile()`
- `is_available()` Whether this target for code generation is available.
- `run()` Runs the code in the namespace.
- `update_namespace()` Update the namespace for this timestep.

**Details**

- `class_name` A short name for this type of `CodeObject`
- `generator_class` The `CodeGenerator` class used by this `CodeObject`
- `__call__(**kwds)` Call self as a function.
- `compile()`
**classmethod is_available()**  
Whether this target for code generation is available. Should use a minimal example to check whether code generation works in general.

**run()**  
Runs the code in the namespace.

**Returns return_value : dict**  
A dictionary with the keys corresponding to the output_variables defined during the call of CodeGenerator.code_object.

**update_namespace()**  
Update the namespace for this timestep. Should only deal with variables where the reference changes every timestep, i.e. where the current reference in namespace is not correct.

**Functions**

<table>
<thead>
<tr>
<th>function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>check_compiler_kwds</td>
<td>Internal function to check the provided compiler keywords against the list of understood keywords.</td>
</tr>
</tbody>
</table>

**check_compiler_kwds function**

(Shortest import: from brian2.codegen.codeobject import check_compiler_kwds)

brian2.codegen.codeobject.check_compiler_kwds(compiler_kwds, accepted_kwds, target)

Internal function to check the provided compiler keywords against the list of understood keywords.

**Parameters**

- **compiler_kwds**: dict  
  Dictionary of compiler keywords and respective list of values.

- **accepted_kwds**: list of str  
  The compiler keywords understood by the code generation target

- **target**: str  
  The name of the code generation target (used for the error message).

**Raises**

- **ValueError** If a compiler keyword is not understood

<table>
<thead>
<tr>
<th>function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>constant_or_scalar</td>
<td>Convenience function to generate code to access the value of a variable.</td>
</tr>
</tbody>
</table>

**constant_or_scalar function**

(Shortest import: from brian2.codegen.codeobject import constant_or_scalar)

brian2.codegen.codeobject.constant_or_scalar(varname, variable)

Convenience function to generate code to access the value of a variable. Will return 'varname' if the variable is a constant, and array_name[0] if it is a scalar array.
create_runner_codeobj(group, code, ...[, ...])  Create a CodeObject for the execution of code in the context of a Group.

create_runner_codeobj function

(Shortest import: from brian2.codegen.codeobject import create_runner_codeobj)

brian2.codegen.codeobject.create_runner_codeobj(group, code, template_name, run_namespace, user_code=None, variable_indices=None, name=None, check_units=True, needed_variables=None, additional_variables=None, template_kwds=None, override_conditional_write=None, codeobj_class=None)

Create a CodeObject for the execution of code in the context of a Group.

Parameters group : Group
    The group where the code is to be run.

code : str or dict of str
    The code to be executed.

template_name : str
    The name of the template to use for the code.

run_namespace : dict-like
    An additional namespace that is used for variable lookup (either an explicitly defined namespace or one taken from the local context).

user_code : str, optional
    The code that had been specified by the user before other code was added automatically. If not specified, will be assumed to be identical to code.

variable_indices : dict-like, optional
    A mapping from Variable objects to index names (strings). If none is given, uses the corresponding attribute of group.

name : str, optional
    A name for this code object, will use group + '_codeobject*' if none is given.

check_units : bool, optional
    Whether to check units in the statement. Defaults to True.

needed_variables : list of str, optional
    A list of variables that are neither present in the abstract code, nor in the USES_VARIABLES statement in the template. This is only rarely necessary, an example being a StateMonitor where the names of the variables are neither known to the template nor included in the abstract code statements.

additional_variables : dict-like, optional
A mapping of names to `Variable` objects, used in addition to the variables saved in group.

`template_kwds`: dict, optional

A dictionary of additional information that is passed to the template.

`override_conditional_write`: list of str, optional:

A list of variable names which are used as conditions (e.g. for refractoriness) which should be ignored.

`codeobj_class`: class, optional

The `CodeObject` class to run code with. If not specified, defaults to the group's `codeobj_class` attribute.

**cppPrefs module**

Preferences related to C++ compilation

**Preferences**

C++ compilation preferences

```python
codegen.cpp.compiler = ''
```

Compiler to use (uses default if empty)

Should be gcc or msvc.

```python
codegen.cpp.define_macros = []
```

List of macros to define; each macro is defined using a 2-tuple, where 'value' is either the string to define it to or None to define it without a particular value (equivalent of "#define FOO" in source or -DFOO on Unix C compiler command line).

```python
codegen.cpp.extra_compile_args = None
```

Extra arguments to pass to compiler (if None, use either extra_compile_args_gcc or extra_compile_args_msvc).

```python
codegen.cpp.extra_compile_args_gcc
  = ['-w', '-O3', '-ffast-math', '-fno-finite-math-only', '-march=native']
```

Extra compile arguments to pass to GCC compiler

```python
codegen.cpp.extra_compile_args_msvc = ['/Ox', '/w', '', '/MP']
```

Extra compile arguments to pass to MSVC compiler (the default /arch: flag is determined based on the processor architecture)

```python
codegen.cpp.extra_link_args = []
```

Any extra platform- and compiler-specific information to use when linking object files together.

```python
codegen.cpp.headers = []
```

A list of strings specifying header files to use when compiling the code. The list might look like ["<vector>"","my_header"长长的]. Note that the header strings need to be in a form than can be pasted at the end of a #include statement in the C++ code.

```python
codegen.cpp.include_dirs = []
```

6.6. Subpackages
Include directories to use. Note that $prefix/include will be appended to the end automatically, where $prefix is Python’s site-specific directory prefix as returned by sys.prefix.

codegen.cpp.libraries = []
List of library names (not filenames or paths) to link against.

codegen.cpp.library_dirs = []
List of directories to search for C/C++ libraries at link time. Note that $prefix/lib will be appended to the end automatically, where $prefix is Python’s site-specific directory prefix as returned by sys.prefix.

codegen.cpp.msvc_architecture = ''
MSVC architecture name (or use system architecture by default).
Could take values such as x86, amd64, etc.

codegen.cpp.msvc_vars_location = ''
Location of the MSVC command line tool (or search for best by default).

codegen.cpp.runtime_library_dirs = []
List of directories to search for C/C++ libraries at run time.

Exported members: get_compiler_and_args, get_msvc_env, compiler_supports_c99, C99Check

Classes

| C99Check(name) | Helper class to create objects that can be passed as an availability_check to a FunctionImplementation. |

**C99Check class**

(Shortest import: from brian2.codegen.cpp_prefs import C99Check)

class brian2.codegen.cpp_prefs.C99Check(name)
Bases: object

Helper class to create objects that can be passed as an availability_check to a FunctionImplementation.

Methods

```python
__call__(*args, **kwargs)  # Call self as a function.
```

Details

```python
__call__(*args, **kwargs)  # Call self as a function.
```

Functions

```python
compiler_supports_c99()
```
**compiler_supports_c99 function**

(Shortest import: from brian2.codegen.cpp_prefs import compiler_supports_c99)

brian2.codegen.cpp_prefs.compiler_supports_c99()

---

**get_compiler_and_args function**

(Shortest import: from brian2.codegen.cpp_prefs import get_compiler_and_args)

brian2.codegen.cpp_prefs.get_compiler_and_args()

Returns the computed compiler and compilation flags

---

**get_msvc_env function**

(Shortest import: from brian2.codegen.cpp_prefs import get_msvc_env)

brian2.codegen.cpp_prefs.get_msvc_env()

---

**update_for_cross_compilation function**

(Shortest import: from brian2.codegen.cpp_prefs import update_for_cross_compilation)

brian2.codegen.cpp_prefs.update_for_cross_compilation(library_dirs, extra_compile_args, extra_link_args, logger=None)

Update the compiler arguments to allow cross-compilation for 32bit on a 64bit Linux system.

**Parameters**

- **library_dirs**: list
  List of library directories (will be modified in-place).
- **extra_compile_args**: list
  List of extra compile args (will be modified in-place).
- **extra_link_args**: list
  List of extra link args (will be modified in-place).
- **logger**: BrianLogger, optional
  The logger to use for the INFO message. Defaults to None (no message).
get_cpu_flags module

This script is used to ask for the CPU flags on Windows. We use this instead of importing the cpuinfo package, because recent versions of py-cpuinfo use the multiprocessing module, and any import of cpuinfo that is not within a 

```python
if __name__ == '__main__':
```

block will lead to the script being executed twice.

The CPU flags are printed to stdout encoded as JSON.

optimisation module

Simplify and optimise sequences of statements by rewriting and pulling out loop invariants.

Exported members: `optimise_statements, ArithmeticSimplifier, Simplifier`

Classes

```python
ArithmeticSimplifier(variables)
```

Carries out the following arithmetic simplifications:

ArithmeticSimplifier class

(Shortest import: from brian2.codegen.optimisation import ArithmeticSimplifier)

```python
class brian2.codegen.optimisation.ArithmeticSimplifier(variables):
    Bases: brian2.parsing.bast.BrianASTRenderer
```

Carries out the following arithmetic simplifications:

1. Constant evaluation (e.g. \(\exp(0) = 1\)) by attempting to evaluate the expression in an “assumptions namespace”
2. Binary operators, e.g. \(0 \times x = 0, 1 \times x = x\), etc. You have to take care that the dtypes match here, e.g. if \(x\) is an integer, then \(1.0 \times x\) shouldn’t be replaced with \(x\) but left as \(1.0 \times x\).

Parameters `variables`: dict of (str, Variable)

Usual definition of variables.

assumptions : sequence of str

Additional assumptions that can be used in simplification, each assumption is a string statement. These might be the scalar statements for example.

Methods

```python
render_BinOp(node)
render_node(node)
```

Assumes that the node has already been fully processed by BrianASTRenderer

Details

```python
render_BinOp (node)
render_node (node)
```

Assumes that the node has already been fully processed by BrianASTRenderer
Simplifier class

(Shortest import: from brian2.codegen.optimisation import Simplifier)

class brian2.codegen.optimisation.Simplifier(variables, scalar_statements, extra_lio_prefix="")

Bases: brian2.parsing.bast.BrianASTRenderer

Carry out arithmetic simplifications (see ArithmeticSimplifier) and loop invariants

Parameters

variables : dict of (str, Variable)

Usual definition of variables.

scalar_statements : sequence of Statement

Predefined scalar statements that can be used as part of simplification

Notes

After calling render_expr on a sequence of expressions (coming from vector statements typically), this object will have some new attributes:

loop_invariants [OrderedDict of (expression, varname)] varname will be of the form _lio_N where N is some integer, and the expressions will be strings that correspond to scalar-only expressions that can be evaluated outside of the vector block.

loop_invariant_dtypes [dict of (varname, dtypename)] dtypename will be one of 'boolean', 'integer', 'float'.

Methods

render_expr(expr)

render_node(node) Assumes that the node has already been fully processed by BrianASTRenderer

Details

render_expr(expr)

render_node(node) Assumes that the node has already been fully processed by BrianASTRenderer

Functions

cancel_identical_terms(primary, inverted) Cancel terms in a collection, e.g.

cancel_identical_terms function

(Shortest import: from brian2.codegen.optimisation import cancel_identical_terms)
brian2.codegen.optimisation.cancel_identical_terms(primary, inverted)

Cancel terms in a collection, e.g. a+b-a should be cancelled to b

Simply renders the nodes into expressions and removes whenever there is a common expression in primary and inverted.

**Parameters**

- **primary**: list of AST nodes
  
  These are the nodes that are positive with respect to the operator, e.g. in x*y/z it would be [x, y].

- **inverted**: list of AST nodes
  
  These are the nodes that are inverted with respect to the operator, e.g. in x*y/z it would be [z].

**Returns**

- **primary**: list of AST nodes
  
  Primary nodes after cancellation

- **inverted**: list of AST nodes
  
  Inverted nodes after cancellation

---

$brian2$.codegen.optimisation.collect$(node)$

Attempts to collect commutative operations into one and simplifies them.

For example, if x and y are scalars, and z is a vector, then (x*z)*y should be rewritten as (x*y)*z to minimise the number of vector operations. Similarly, ((x*2)*3)*4 should be rewritten as x*24.

Works for either multiplication/division or addition/subtraction nodes.

The final output is a subexpression of the following maximal form:

$$ (((\text{numerical_value}*(\text{product of scalars})/\text{(product of scalars)})*(\text{product of vectors})/(\text{product of vectors})/\text{product of vectors})/\text{product of vectors})/\text{product of vectors})/\text{product of vectors}) \ldots $$

Any possible cancellations will have been done.

**Parameters**

- **node**: Brian AST node
  
  The node to be collected/simplified.

**Returns**

- **node**: Brian AST node
  
  Simplified node.

---

$brian2$.codegen.optimisation.collect_commutative$(node, primary, inverted, ...)$

**collect_commutative function**

(*Shortest import: from brian2.codegen.optimisation import collect_commutative*)

---
evaluate_expr function

(Shortest import: from brian2.codegen.optimisation import evaluate_expr)

brian2.codegen.optimisation.evaluate_expr(expr, ns)

Try to evaluate the expression in the given namespace

Returns either (value, True) if successful, or (expr, False) otherwise.

expression_complexity function

(Shortest import: from brian2.codegen.optimisation import expression_complexity)

brian2.codegen.optimisation.expression_complexity(expr, variables)

optimise_statements function

(Shortest import: from brian2.codegen.optimisation import optimise_statements)

brian2.codegen.optimisation.optimise_statements(scalar_statements, ...

Optimise a sequence of scalar and vector statements

Performs the following optimisations:

1. Constant evaluations (e.g. exp(0) to 1). See evaluate_expr.
2. Arithmetic simplifications (e.g. 0*x to 0). See ArithmeticSimplifier, collect().
3. Pulling out loop invariants (e.g. v*exp(-dt/tau) to a=exp(-dt/tau) outside the loop and v*a inside). See Simplifier.
4. Boolean simplifications (allowing the replacement of expressions with booleans with a sequence of if/thens). See Simplifier.

Parameters

scalar_statements : sequence of Statement

Statements that only involve scalar values and should be evaluated in the scalar block.

vector_statements : sequence of Statement

Statements that involve vector values and should be evaluated in the vector block.

variables : dict of (str, Variable)

Definition of the types of the variables.

blockname : str, optional
Name of the block (used for LIO constant prefixes to avoid name clashes)

**Returns** `new_scalar_statements` : sequence of `Statement`

As above but with loop invariants pulled out from vector statements

`new_vector_statements` : sequence of `Statement`

Simplified/optimised versions of statements

---

### reduced_node function

(Shortest import: `from brian2.codegen.optimisation import reduced_node`)

brian2.codegen.optimisation.reduced_node(terms, op)

Reduce a sequence of terms with the given operator

For example, if terms were [a, b, c] and op was multiplication then the reduction would be (a*b)*c.

**Parameters**

- **terms** : list
  - AST nodes.
- **op** : AST node
  - Could be `ast.Mult` or `ast.Add`.

**Examples**

```python
>>> import ast
>>> nodes = [ast.Name(id='x'), ast.Name(id='y'), ast.Name(id='z')]
>>> ast.dump(reduced_node(nodes, ast.Mult), annotate_fields=False)
"BinOp(BinOp(Name('x'), Mult(), Name('y')), Mult(), Name('z'))"
>>> nodes = [ast.Name(id='x')]
>>> ast.dump(reduced_node(nodes, ast.Add), annotate_fields=False)
"Name('x')"
```

---

### permutation_analysis module

Module for analysing synaptic pre and post code for synapse order independence.

**Exported members:** `OrderDependenceError, check_for_order_independence`

**Classes**

---

**OrderDependenceError class**

(Shortest import: `from brian2.codegen.permutation_analysis import OrderDependenceError`)

class brian2.codegen.permutation_analysis.OrderDependenceError
Bases: `Exception`

Functions

```python
_check_for_order_independence(statements, ...)
```

Check that the sequence of statements doesn’t depend on the order in which the indices are iterated through.

**check_for_order_independence function**

```python
from brian2.codegen.permutation_analysis import check_for_order_independence
brian2.codegen.permutation_analysis.check_for_order_independence(statements, variables, indices)
```

Check that the sequence of statements doesn’t depend on the order in which the indices are iterated through.

**statements module**

Module providing the `Statement` class.

**Classes**

```python
Statement(var, op, expr, comment, dtype[, ...])
```

A single line mathematical statement.

**Statement class**

```python
from brian2.codegen.statements import Statement

class brian2.codegen.statements.Statement(var, op, expr, comment, dtype, constant=False, subexpression=False, scalar=False)
```

A single line mathematical statement.

The structure is `var op expr`.

**Parameters**

- **var**: `str`
  The left hand side of the statement, the value being written to.

- **op**: `str`
  The operation, can be any of the standard Python operators (including `+=` etc.) or a special operator `:=` which means you are defining a new symbol (whereas `=` means you are setting the value of an existing symbol).

- **expr**: `str, Expression`
  The right hand side of the statement.

- **dtype**: `dtype`
  The numpy dtype of the value or array `var()`.

- **constant**: `bool`, optional
  Set this flag to `True` if the value will not change (only applies for `op=':='`).

---

**6.6. Subpackages**

---
**subexpression**: bool, optional

Set this flag to True if the variable is a subexpression. In some languages (e.g. Python) you can use this to save a memory copy, because you don’t need to do \( \text{lhs}[:] = \text{rhs} \) but a redefinition \( \text{lhs} = \text{rhs} \).

**scalar**: bool, optional

Set this flag to True if \( \text{var()} \) and \( \text{expr} \) are scalar.

**Notes**

Will compute the following attribute:

**inplace** True or False depending if the operation is in-place or not.

Boolean simplification notes:

Will initially set the attribute `used_boolean_variables` to `None`. This is set by `optimise_statements` when it is called on a sequence of statements to the list of boolean variables that are used in this expression. In addition, the attribute `boolean_simplified_expressions` is set to a dictionary with keys consisting of a tuple of pairs (var, value) where var is the name of the boolean variable (will be in `used_boolean_variables`) and var is True or False. The values of the dictionary are strings representing the simplified version of the expression if each var=value substitution is made for that key. The keys will range over all possible values of the set of boolean variables. The complexity of the original statement is set as the attribute `complexity_std`, and the complexity of the simplified versions are in the dictionary `complexities` (with the same keys).

This information can be used to generate code that replaces a complex expression that varies depending on the value of one or more boolean variables with an if/then sequence where each subexpression is simplified. It is optional to use this (e.g. the numpy codegen does not, but the weave and cython ones do).

**targets module**

Module that stores all known code generation targets as `codegen_targets`.

Exported members: `codegen_targets`

**templates module**

Exported members: `Templater`

**Classes**

| `CodeObjectTemplate(template, template_source)` | Single template object returned by `Templater` and used for final code generation |

**CodeObjectTemplate class**

(Shortest import: from brian2.codegen.templates import CodeObjectTemplate)

class brian2.codegen.templates.CodeObjectTemplate(template, template_source)

Bases: `object`

Single template object returned by `Templater` and used for final code generation
Should not be instantiated by the user, but only directly by `Templater`.

**Notes**

The final code is obtained from this by calling the template (see `__call__`).

**Attributes**

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Description</th>
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</thead>
<tbody>
<tr>
<td><code>allows_scalar_write</code></td>
<td>Does this template allow writing to scalar variables?</td>
</tr>
<tr>
<td><code>iterate_all</code></td>
<td>The indices over which the template iterates completely</td>
</tr>
<tr>
<td><code>variables</code></td>
<td>The set of variables in this template</td>
</tr>
<tr>
<td><code>writes_read_only</code></td>
<td>Read-only variables that are changed by this template</td>
</tr>
</tbody>
</table>

**Methods**

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>__call__</code></td>
<td>Return a usable code block or blocks from this template.</td>
</tr>
</tbody>
</table>

**Details**

- **`allows_scalar_write`**
  - Does this template allow writing to scalar variables?

- **`iterate_all`**
  - The indices over which the template iterates completely

- **`variables`**
  - The set of variables in this template

- **`writes_read_only`**
  - Read-only variables that are changed by this template

**`__call__`**

```python
__call__(scalar_code, vector_code, **kwds)
```

Return a usable code block or blocks from this template.

**Parameters**

- `scalar_code` : dict
  - Dictionary of scalar code blocks.

- `vector_code` : dict
  - Dictionary of vector code blocks

- `**kwds` : dict
  - Additional parameters to pass to the template

**Notes**

Returns either a string (if macros were not used in the template), or a `MultiTemplate` (if macros were used).
LazyTemplateLoader class

(Shortest import: from brian2.codegen.templates import LazyTemplateLoader)

class brian2.codegen.templates.LazyTemplateLoader(
    environment, extension
)

Bases: object

Helper object to load templates only when they are needed.

Methods

get_template(name)

Details

gt_template(name)

MultiTemplate class

(Shortest import: from brian2.codegen.templates import MultiTemplate)

class brian2.codegen.templates.MultiTemplate(module)

Bases: collections.abc.Mapping

Code generated by a CodeObjectTemplate with multiple blocks

Each block is a string stored as an attribute with the block name. The object can also be accessed as a dictionary.

Templater class

(Shortest import: from brian2.codegen.templates import Templater)

class brian2.codegen.templates.Templater(
    package_name, extension[, env Globals]
)

Bases: object

Class to load and return all the templates a CodeObject defines.

Parameters package_name: str, tuple of str

The package where the templates are saved. If this is a tuple then each template will be searched in order starting from the first package in the tuple until the template is found. This allows for derived templates to be used. See also derive.
env_globals : dict (optional)

A dictionary of global values accessible by the templates. Can be used for providing utility functions. In all cases, the filter ‘autoindent’ is available (see existing templates for example usage).

Notes

Templates are accessed using templater.template_base_name (the base name is without the file extension). This returns a CodeObjectTemplate.

Methods

```
def derive(package_name[, extension, env_globals])
```

Return a new Templater derived from this one, where the new package name and globals overwrite the old.

Details

```
def derive (package_name, extension=None, env_globals=None)
```

Return a new Templater derived from this one, where the new package name and globals overwrite the old.

Functions

```
autoindent(code)
```

autoindent function

(Shortest import: from brian2.codegen.templates import autoindent)

brian2.codegen.templates.autoindent(code)

```
autoindent_postfilter(code)
```

autoindent_postfilter function

(Shortest import: from brian2.codegen.templates import autoindent_postfilter)

brian2.codegen.templates.autoindent_postfilter(code)

```
variables_to_array_names(variables[...])
```

variables_to_array_names function

(Shortest import: from brian2.codegen.templates import variables_to_array_names)

brian2.codegen.templates.variables_to_array_names(variables, access_data=True)
translation module

This module translates a series of statements into a language-specific syntactically correct code block that can be inserted into a template.

It infers whether or not a variable can be declared as constant, etc. It should handle common subexpressions, and so forth.

The input information needed:

- The sequence of statements (a multiline string) in standard mathematical form
- The list of known variables, common subexpressions and functions, and for each variable whether or not it is a value or an array, and if an array what the dtype is.
- The dtype to use for newly created variables
- The language to translate to

Exported members: `analyse_identifiers, get_identifiers_recursively`

Classes

`LineInfo(**kwds)` A helper class, just used to store attributes.

LineInfo class

(Shortest import: from brian2.codegen.translation import LineInfo)

class brian2.codegen.translation.LineInfo(**kwds)
    Bases: object

    A helper class, just used to store attributes.

Functions

`analyse_identifiers(code, variables[, recursive])` Analyses a code string (sequence of statements) to find all identifiers by type.

analyse_identifiers function

(Shortest import: from brian2.codegen.translation import analyse_identifiers)

brian2.codegen.translation.analyse_identifiers(code, variables, recursive=False)

Analyses a code string (sequence of statements) to find all identifiers by type.

In a given code block, some variable names (identifiers) must be given as inputs to the code block, and some are created by the code block. For example, the line:

```
a = b+c
```

This could mean to create a new variable a from b and c, or it could mean modify the existing value of a from b or c, depending on whether a was previously known.

Parameters  

- **code**: str
  The code string, a sequence of statements one per line.

- **variables**: dict of Variable, set of names
Specifiers for the model variables or a set of known names

**recursive**: bool, optional

Whether to recurse down into subexpressions (defaults to False).

**Returns**

`newly_defined` : set

A set of variables that are created by the code block.

`used_known` : set

A set of variables that are used and already known, a subset of the `known` parameter.

`unknown` : set

A set of variables which are used by the code block but not defined by it and not previously known. Should correspond to variables in the external namespace.

```python
get_identifiers_recursively(expressions,...)  # Gets all the identifiers in a list of expressions, recursing
down into subexpressions.
```

**get_identifiers_recursively function**

(Shortest import: from brian2.codegen.translation import get_identifiers_recursively)

```python
brian2.codegen.translation.get_identifiers_recursively(expressions, variables, include_numbers=False)
```

 Gets all the identifiers in a list of expressions, recursing down into subexpressions.

 **Parameters**

`expressions` : list of str

List of expressions to check.

`variables` : dict-like

Dictionary of Variable objects

`include_numbers` : bool, optional

Whether to include number literals in the output. Defaults to False.

```python
is_scalar_expression(expr, variables)  # Whether the given expression is scalar.
```

**is_scalar_expression function**

(Shortest import: from brian2.codegen.translation import is_scalar_expression)

```python
brian2.codegen.translation.is_scalar_expression(expr, variables)
```

Whether the given expression is scalar.

 **Parameters**

`expr` : str

The expression to check

`variables` : dict-like

Variable and Function object for all the identifiers used in expr

 **Returns**

`scalar` : bool
Whether \texttt{expr} is a scalar expression

\texttt{make_statements} (\texttt{code}, \texttt{variables}, \texttt{dtype}[, ...])

Turn a series of abstract code statements into Statement objects, inferring whether each line is a set/declare operation, whether the variables are constant or not, and handling the caching of subexpressions.

\textbf{make\textunderscore statements function}

(\textit{Shortest import}: from \texttt{brian2.codegen.translation} import \texttt{make\textunderscore statements})

\texttt{brian2.codegen.translation.make\textunderscore statements} (\texttt{code}, \texttt{variables}, \texttt{dtype}, \texttt{optimise=True}, \texttt{block\textunderscore name=”)}

Turn a series of abstract code statements into Statement objects, inferring whether each line is a set/declare operation, whether the variables are constant or not, and handling the caching of subexpressions.

**Parameters**

\textbf{code : str}

A (multi-line) string of statements.

\textbf{variables : dict-like}

A dictionary of with \texttt{Variable} and \texttt{Function} objects for every identifier used in the \texttt{code}.

\textbf{dtype : dtype}

The data type to use for temporary variables

\textbf{optimise : bool, optional}

Whether to optimise expressions, including pulling out loop invariant expressions and putting them in new scalar constants. Defaults to \texttt{False}, since this function is also used just to in contexts where we are not interested by this kind of optimisation. For the main code generation stage, its value is set by the \texttt{codegen.loop\_invariant\_optimisations} preference.

\textbf{blockname : str, optional}

A name for the block (used to name intermediate variables to avoid name clashes when multiple blocks are used together)

**Returns**

---

\textbf{scalar\textunderscore statements, vector\textunderscore statements} : (list of Statement, list of Statement)

Lists with statements that are to be executed once and statements that are to be executed once for every neuron/synapse/… (or in a vectorised way)

**Notes**

If \texttt{optimise} is \texttt{True}, then \texttt{scalar\textunderscore statements} may include newly introduced scalar constants that have been identified as loop-invariant and have therefore been pulled out of the vector statements. The resulting statements will also use augmented assignments where possible, i.e. a statement such as \texttt{w = w + 1} will be replaced by \texttt{w += 1}. Also, statements involving booleans will have additional information added to them (see \texttt{Statement} for details) describing how the statement can be reformulated as a sequence of if/then statements. Calls \texttt{optimise\_statements}.  

---

386 Chapter 6. \texttt{brian2} package
Subpackages

generators package

GSL_generator module

GSLCodeGenerators for code that uses the ODE solver provided by the GNU Scientific Library (GSL)

Exported members: GSLCodeGenerator, GSLWeaveCodeGenerator, GSLCythonCodeGenerator

Classes

| GSLCodeGenerator(variables, ...[, ...]) | GSL code generator. |

GSLCodeGenerator class

(Shortest import: from brian2.codegen.generators import GSLCodeGenerator)

class brian2.codegen.generators.GSL_generator.GSLCodeGenerator(variables, variable_indices, owner, iterate_all, codeobj_class, name, template_name, override_conditional_write=None, allows_scalar_write=False)

Bases: object

GSL code generator.

Notes

Approach is to first let the already existing code generator for a target language do the bulk of the translating from abstract_code to actual code. This generated code is slightly adapted to render it GSL compatible. The most critical part here is that the vector_code that is normally contained in a loop in the `main()` is moved to the function `_GSL_func` that is sent to the GSL integrator. The variables used in the vector_code are added to a struct named `dataholder` and their values are set from the Brian namespace just before the scalar code block.

Methods

| add_gsl_variables_as_non_scalar(diff_vars) | Add gsl variables as non-scalar. |
| add_meta_variables(options)                |                                    |
| c_data_type(dtype)                         | Get string version of object dtype that is attached to Brian variables. |
| diff_var_to_replace(diff_vars)             | Add differential variable-related strings that need to be replaced to go from normal brian to GSL code |

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### Table 59 – continued from previous page

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<th>Function Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>find_differential_variables(code)</code></td>
<td>Find the variables that were tagged <code>_gsl_{var}_f{ind}</code> and return var, ind pairs.</td>
</tr>
<tr>
<td><code>find_function_names()</code></td>
<td>Return a list of used function names in the self.variables dictionary.</td>
</tr>
<tr>
<td><code>find_undefined_variables(statements)</code></td>
<td>Find identifiers that are not in self.variables dictionary.</td>
</tr>
<tr>
<td><code>find_used_variables(statements, other_variables)</code></td>
<td>Find all the variables used in the right hand side of the given expressions.</td>
</tr>
<tr>
<td><code>get_dimension_code(diff_num)</code></td>
<td>Generate code for function that sets the dimension of the ODE system.</td>
</tr>
<tr>
<td><code>initialize_array(varname, values)</code></td>
<td>Initialize a static array with given floating point values.</td>
</tr>
<tr>
<td><code>is_constant_and_cpp_standalone(var_obj)</code></td>
<td>Check whether self.cpp_standalone and variable is Constant.</td>
</tr>
<tr>
<td><code>is_cpp_standalone()</code></td>
<td>Check whether we're running with cpp_standalone.</td>
</tr>
<tr>
<td><code>make_function_code(lines)</code></td>
<td>Add lines of GSL translated vector code to <code>non-changing</code> _GSL_func code.</td>
</tr>
<tr>
<td><code>scale_array_code(diff_vars, method_options)</code></td>
<td>Return code for definition of _GSL_scale_array in generated code.</td>
</tr>
<tr>
<td><code>to_replace_vector_vars(variables_in_vector)</code></td>
<td>Create dictionary containing key, value pairs with to be replaced text to translate from conventional Brian to GSL.</td>
</tr>
<tr>
<td><code>translate(code, dtype)</code></td>
<td>Translates an abstract code block into the target language.</td>
</tr>
<tr>
<td><code>translate_scalar_code(code_lines,...)</code></td>
<td>Translate scalar code: if calculated variables are used in the vector_code their value is added to the variable in the _dataholder.</td>
</tr>
<tr>
<td><code>translate_vector_code(code_lines, to_replace)</code></td>
<td>Translate vector code to GSL compatible code by substituting fragments of code.</td>
</tr>
<tr>
<td><code>unpack_namespace(variables_in_vector, ...[, ...])</code></td>
<td>Write code that unpacks Brian namespace to cython/cpp namespace.</td>
</tr>
<tr>
<td><code>unpack_namespace_single(var_obj, in_vector,...)</code></td>
<td>Writes the code necessary to pull single variable out of the Brian namespace into the generated code.</td>
</tr>
<tr>
<td><code>var_init_lhs(var, type)</code></td>
<td>Get string version of the left hand side of an initializing expression.</td>
</tr>
<tr>
<td><code>write_dataholder(variables_in_vector)</code></td>
<td>Return string with full code for _dataholder struct.</td>
</tr>
<tr>
<td><code>write_dataholder_single(var_obj)</code></td>
<td>Return string declaring a single variable in the _dataholder struct.</td>
</tr>
<tr>
<td><code>yvector_code(diff_vars)</code></td>
<td>Generate code for function dealing with GSLs y vector.</td>
</tr>
</tbody>
</table>

### Details

#### add_gsl_variables_as_non_scalar(diff_vars)

Add _gsl variables as non-scalar.

In GSLStateUpdater the differential equation variables are substituted with GSL tags that describe the information needed to translate the conventional Brian code to GSL compatible code. This function tells Brian that the variables that contain these tags should always be vector variables. If we don’t do this, Brian renders the tag-variables as scalar if no vector variables are used in the right hand side of the expression.

**Parameters**

- **diff_vars**: dict
dictionary with variables as keys and differential equation index as value
add_meta_variables *(options)*

c_data_type *(dtype)*
Get string version of object dtype that is attached to Brian variables. c pp_generator already has this
function, but the Cython generator does not, but we need it for GSL code generation.

diff_var_to_replace *(diff_vars)*
Add differential variable-related strings that need to be replaced to go from normal brian to GSL code
From the code generated by Brian’s ‘normal’ generators (cpp_generator or cython_generator a few bits
of text need to be replaced to get GSL compatible code. The bits of text related to differential equation
variables are put in the replacer dictionary in this function.

Parameters **diff_vars** : dict
dictionary with variables as keys and differential equation index as value

Returns **to_replace** : dict
dictionary with strings that need to be replaced as keys and the strings that will replace them as values

find_differential_variables *(code)*
Find the variables that were tagged _gsl_{var}_f{ind} and return var, ind pairs.
GSLStateUpdater tagged differential variables and here we extract the information given in these tags.

Parameters **code** : list of strings
A list of strings containing gsl tagged variables

Returns **diff_vars** : dict
A dictionary with variable names as keys and differential equation index as value

find_function_names()
Return a list of used function names in the self.variables dictionary
Functions need to be ignored in the GSL translation process, because the brian generator already suffi-
ciently dealt with them. However, the brian generator also removes them from the variables dict, so there
is no way to check whether an identifier is a function after the brian translation process. This function
is called before this translation process and the list of function names is stored to be used in the GSL
translation.

Returns **function_names** : list
list of strings that are function names used in the code

find_undefined_variables *(statements)*
Find identifiers that are not in self.variables dictionary.
Brian does not save the _lio_ variables it uses anywhere. This is problematic for our GSL implementa-
tion because we save the lio variables in the _dataholder struct (for which we need the datatype of the
variables). This function adds the left hand side variables that are used in the vector code to the variable
dictionary as AuxiliaryVariables (all we need later is the datatype).

Parameters **statements** : list
list of statement objects (need to have the dtype attribute)

Notes
I keep self.variables and other_variables separate so I can distinguish what variables are in
the Brian namespace and which ones are defined in the code itself.
**find_used_variables** *(statements, other_variables)*  
Find all the variables used in the right hand side of the given expressions.

- **Parameters**  
  *statements*: list  
  list of statement objects

- **Returns**  
  *used_variables*: dict  
  dictionary of variables that are used as variable name (str), Variable pairs.

**get_dimension_code** *(diff_num)*  
Generate code for function that sets the dimension of the ODE system.

GSL needs to know how many differential variables there are in the ODE system. Since the current approach is to have the code in the vector loop the same for all simulations, this dimension is set by an external function. The code for this set_dimension function is written here. It is assumed the code will be the same for each target language with the exception of some syntactical differences.

- **Parameters**  
  *diff_num*: int  
  Number of differential variables that describe the ODE system

- **Returns**  
  *set_dimension_code*: str  
  The code describing the target language function in a single string

**initialize_array** *(varname, values)*  
Initialize a static array with given floating point values. E.g. in C++, when called with arguments array and [1.0, 3.0, 2.0], this method should return double array[] = {1.0, 3.0, 2.0}.

- **Parameters**  
  *varname*: str  
  The name of the array variable that should be initialized

- **Parameters**  
  *values*: list of float  
  The values that should be assigned to the array

- **Returns**  
  *code*: str  
  One or more lines of array initialization code.

**is_constant_and_cpp_standalone** *(var_obj)*  
Check whether self.cpp_standalone and variable is Constant.

This check is needed because in the case of using the cpp_standalone device we do not want to apply our GSL variable conversion (var -> _GSL_dataholder.var), because the cpp_standalone code generation process involves replacing constants with their actual value (‘freezing’). This results in code that looks like (if for example var = 1.2): _GSL_dataholder.1.2 = 1.2 and _GSL_dataholder->1.2. To prevent repetitive calls to get_device() etc. the outcome of is_cpp_standalone is saved.

- **Parameters**  
  *var_obj*: Variable  
  instance of brian Variable class describing the variable

- **Returns**  
  *is_cpp_standalone*: bool  
  whether the used device is cpp_standalone and the given variable is an instance of Constant

**is_cpp_standalone** ()  
Check whether we’re running with cpp_standalone.

Test if get_device() is instance CPPStandaloneDevice.

- **Returns**  
  *is_cpp_standalone*: bool
whether currently using cpp_standalone device

See also:

**is_constant_and_cpp_standalone** uses the returned value

**make_function_code** *(lines)*

Add lines of GSL translated vector code to ‘non-changing’ _GSL_func code.

Adds nonchanging aspects of GSL _GSL_func code to lines of code written somewhere else *(translate_vector_code)*. Here these lines are put between the non-changing parts of the code and the target-language specific syntax is added.

**Parameters**

- lines : str
  Code containing GSL version of equations

**Returns**

- function_code : str
  code describing _GSL_func that is sent to GSL integrator.

**scale_array_code** *(diff_vars, method_options)*

Return code for definition of _GSL_scale_array in generated code.

**Parameters**

- diff_vars : dict
dictionary with variable name (str) as key and differential variable index (int) as value
- method_options : dict
dictionary containing integrator settings

**Returns**

- code : str
  full code describing a function returning a array containing doubles with the absolute errors for each differential variable (according to their assigned index in the GSL State-Updater)

**to_replace_vector_vars** *(variables_in_vector, ignore=frozenset())*

Create dictionary containing key, value pairs with to be replaced text to translate from conventional Brian to GSL.

**Parameters**

- variables_in_vector : dict
dictionary with variable name (str), Variable pairs of variables occurring in vector code
- ignore : set, optional
  set of strings with variable names that should be ignored

**Returns**

- to_replace : dict
dictionary with strings that need to be replaced i.e. _lio_1 will be _GSL_dataholder._lio_1 (in cython) or _GSL_dataholder->_lio_1 (cpp)

**Notes**

t will always be added because GSL defines its own t. i.e. for cpp: `{’const t = _ptr_array_defaultclock_t[0];’ : ‘’}`

**translate** *(code, dttype)*

Translates an abstract code block into the target language.
**translate_scalar_code** *(code_lines, variables_in_scalar, variables_in_vector)*

Translate scalar code: if calculated variables are used in the vector_code their value is added to the variable in the _dataholder.

**Parameters**
- *code_lines*: list
  - list of strings containing scalar code
- *variables_in_vector*: dict
  - dictionary with variable name (str), *Variable* pairs of variables occurring in vector code
- *variables_in_scalar*: dict
  - dictionary with variable name (str), *Variable* pairs of variables occurring in scalar code

**Returns**
- *scalar_code*: str
  - code fragment that should be injected in the main before the loop

**translate_vector_code** *(code_lines, to_replace)*

Translate vector code to GSL compatible code by substituting fragments of code.

**Parameters**
- *code_lines*: list
  - list of strings describing the vector_code
- *to_replace*: dict
  - dictionary with to be replaced strings (see to_replace_vector_vars and to_replace_diff_vars)

**Returns**
- *vector_code*: str
  - New code that is now to be added to the function that is sent to the GSL integrator

**unpack_namespace** *(variables_in_vector, variables_in_scalar, ignore=frozenset())*

Write code that unpacks Brian namespace to cython/cpp namespace.

For vector code this means putting variables in _dataholder (i.e. _GSL_dataholder->var or _GSL_dataholder.var = ...) Note that code is written so a variable could occur both in scalar and vector code

**Parameters**
- *variables_in_vector*: dict
  - dictionary with variable name (str), *Variable* pairs of variables occurring in vector code
- *variables_in_scalar*: dict
  - dictionary with variable name (str), *Variable* pairs of variables occurring in scalar code
- *ignore*: set, optional
  - set of string names of variables that should be ignored

**Returns**
- *unpack_namespace_code*: str
  - code fragment unpacking the Brian namespace (setting variables in the _dataholder struct in case of vector)

**unpack_namespace_single** *(var_obj, in_vector, in_scalar)*

Writes the code necessary to pull single variable out of the Brian namespace into the generated code.
The code created is significantly different between cpp and cython, so I decided to not make this function general over all target languages (i.e. in contrast to most other functions that only have syntactical differences)

```python
var_init_lhs(var, type)
Get string version of the left hand side of an initializing expression

Parameters
    var : str
    type : str

Returns
    code : str
    For cpp returns type + var, while for cython just var
```

```python
write_dataholder(variables_in_vector)
Return string with full code for _dataholder struct.

Parameters
    variables_in_vector : dict
dictionary containing variable name as key and Variable as value

Returns
    code : str
code for _dataholder struct
```

```python
write_dataholder_single(var_obj)
Return string declaring a single variable in the _dataholder struct.

Parameters
    var_obj : Variable

Returns
    code : str
    string describing this variable object as required for the _dataholder struct (e.g. double* _array_neurongroup_v)
```

```python
yvector_code(diff_vars)
Generate code for function dealing with GSLs y vector.

The values of differential variables have to be transferred from Brian’s namespace to a vector that is given to GSL. The transferring from Brian –> y and back from y –> Brian after integration happens in separate functions. The code for this is written here.

Parameters
    diff_vars : dictionary
    Dictionary containing variable names as keys (str) and differential variable index as value

Returns
    yvector_code : str
    The code for the two functions (_fill_y_vector and _empty_y_vector) as single string.
```

```python
GSLCythonCodeGenerator(variables, ...[, ...])
```

### Methods

#### GSLCythonCodeGenerator class

(Shortest import: from brian2.codegen.generators import GSLCythonCodeGenerator)
class brian2.codegen.generators.GSL_generator.GSLCythonCodeGenerator(variables, variable_indices, owner, iterate_all, codeobj_class, name, template_name, override_conditional_write=None, allows_scalar_write=False)

Bases: brian2.codegen.generators.GSL_generator.GSLCodeGenerator

Methods

<table>
<thead>
<tr>
<th>Method Name</th>
<th>Description</th>
</tr>
</thead>
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<tr>
<td><code>c_data_type(dtype)</code></td>
<td>Get string version of object dtype that is attached to Brian variables.</td>
</tr>
<tr>
<td><code>get_array_name(var[, access_data])</code></td>
<td>Initialize a static array with given floating point values.</td>
</tr>
<tr>
<td><code>initialize_array(varname, values)</code></td>
<td>Writes the code necessary to pull single variable out of the Brian namespace into the generated code.</td>
</tr>
<tr>
<td><code>unpack_namespace_single(var_obj, in_vector, ...)</code></td>
<td>Get string version of the left hand side of an initializing expression</td>
</tr>
<tr>
<td><code>var_init_lhs(var, type)</code></td>
<td></td>
</tr>
<tr>
<td><code>var_replace_diff_var_lhs(var, ind)</code></td>
<td></td>
</tr>
</tbody>
</table>

Details

**c_data_type (dtype)**
Get string version of object dtype that is attached to Brian variables. c pp_generator already has this function, but the Cython generator does not, but we need it for GSL code generation.

**static get_array_name (var, access_data=True)**

**initialize_array (varname, values)**
Initialize a static array with given floating point values. E.g. in C++, when called with arguments array and [1.0, 3.0, 2.0], this method should return double array[] = {1.0, 3.0, 2.0}.

Parameters

- **varname**: str
  - The name of the array variable that should be initialized

- **values**: list of float
  - The values that should be assigned to the array

Returns

- **code**: str
  - One or more lines of array initialization code.

**unpack_namespace_single (var_obj, in_vector, in_scalar)**
Writes the code necessary to pull single variable out of the Brian namespace into the generated code.
The code created is significantly different between cpp and cython, so I decided to not make this function general over all target languages (i.e. in contrast to most other functions that only have syntactical differences)

**var_init_lhs**(var, type)
Get string version of the left hand side of an initializing expression

**Parameters**
- **var**: str
- **type**: str

**Returns**
- **code**: str
  For cpp returns type + var, while for cython just var

**var_replace_diff_var_lhs**(var, ind)

---

**GSLWeaveCodeGenerator**(variables, ...[, ...])

### Methods

**GSLWeaveCodeGenerator class**

*(Shortest import: from brian2.codegen.generators import GSLWeaveCodeGenerator)*

**class** brian2.codegen.generators.GSL_generator.GSLWeaveCodeGenerator**(variables, variable_indices, owner, iterate_all, codeobj_class, name, template_name, override_conditional_write=None, allows_scalar_write=False)**

**Bases:** brian2.codegen.generators.GSL_generator.GSLCodeGenerator

### Methods

<table>
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<td><strong>c_data_type</strong>(dtype)</td>
<td>Get string version of object dtype that is attached to Brian variables.</td>
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<tr>
<td><strong>get_array_name</strong>(var[, access_data])</td>
<td>Initialize a static array with given floating point values.</td>
</tr>
<tr>
<td><strong>initialize_array</strong>(varname, values)</td>
<td>Writes the code necessary to pull single variable out of the Brian namespace into the generated code.</td>
</tr>
<tr>
<td><strong>unpack_namespace_single</strong>(var_obj, in_vector, ...)</td>
<td>Get string version of the left hand side of an initializing expression</td>
</tr>
</tbody>
</table>

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<table>
<thead>
<tr>
<th>Function</th>
<th>Parameters</th>
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</tr>
</thead>
<tbody>
<tr>
<td>var_replace_diff_var_lhs</td>
<td>(var, ind)</td>
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<td>static get_array_name (var, access_data=True)</td>
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<td></td>
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</tr>
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<td></td>
<td>Parameters varname : str</td>
<td>The name of the array variable that should be initialized</td>
</tr>
<tr>
<td></td>
<td>values : list of float</td>
<td>The values that should be assigned to the array</td>
</tr>
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<td></td>
<td></td>
<td>Returns code: str</td>
</tr>
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<td></td>
<td>One or more lines of array initialization code.</td>
</tr>
<tr>
<td>unpack_namespace_single</td>
<td>(var_obj, in_vector, in_scalar)</td>
<td>Writes the code necessary to pull single variable out of the Brian namespace</td>
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<td>into the generated code.</td>
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<td></td>
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<td></td>
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<tr>
<td></td>
<td>Parameters var : str</td>
<td></td>
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<tr>
<td></td>
<td>type : str</td>
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<td></td>
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<tr>
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<td>just var</td>
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<td>var_replace_diff_var_lhs</td>
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Functions

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<tr>
<th>Function</th>
<th>Parameters</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>valid_gsl_dir (val)</td>
<td>Validate given string to be path containing</td>
<td>Validate given string to be path containing required GSL files.</td>
</tr>
<tr>
<td></td>
<td>required GSL files.</td>
<td></td>
</tr>
</tbody>
</table>

valid_gsl_dir function

(Shortest import: from brian2.codegen.generators.GSL_generator import valid_gsl_dir)

brian2.codegen.generators.GSL_generator.valid_gsl_dir(val)

Validate given string to be path containing required GSL files.
base module

Base class for generating code in different programming languages, gives the methods which should be overridden to implement a new language.

Exported members: CodeGenerator

Classes

CodeGenerator(variables, variable_indices, ...) Base class for all languages.

CodeGen class

(Shortest import: from brian2.codegen.generators import CodeGenerator)

class brian2.codegen.generators.base.CodeGenerator(variables, variable_indices, owner, iterate_all, codeobj_class, name, template_name, override_conditional_write=None, allows_scalar_write=False)

Bases: object

Base class for all languages.

See definition of methods below.

TODO: more details here

Methods

array_read_write(statements) Helper function, gives the set of ArrayVariables that are read from and written to in the series of statements.

arrays_helper(statements) Combines the two helper functions array_read_write and get_conditional_write_vars, and updates the read set.

determine_keywords() A dictionary of values that is made available to the templated.

get_array_name(var[, access_data]) Get a globally unique name for a ArrayVariable.

get_conditional_write_vars() Helper function, returns a dict of mappings (varname, condition_var_name) indicating that when varname is written to, it should only be when condition_var_name is True.

has_repeated_indices(statements) Whether any of the statements potentially uses repeated indices (e.g.

translate(code, dtype) Translates an abstract code block into the target language.

translate_expression(expr) Translate the given expression string into a string in the target language, returns a string.

translate_one_statement_sequence(statements) Continued on next page
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<table>
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<tr>
<th>Method</th>
<th>Description</th>
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</thead>
<tbody>
<tr>
<td>translate_statement(statement)</td>
<td>Translate a single line Statement into the target language, returns a string.</td>
</tr>
<tr>
<td>translate_statement_sequence(...)</td>
<td>Translate a sequence of Statement into the target language, taking care to declare variables, etc.</td>
</tr>
</tbody>
</table>

Details

array_read_write(statements)
Helper function, gives the set of ArrayVariables that are read from and written to in the series of statements. Returns the pair read, write of sets of variable names.

arrays_helper(statements)
Combines the two helper functions array_read_write and get_conditional_write_vars, and updates the read set.

determine_keywords()
A dictionary of values that is made available to the templated. This is used for example by the CPPCodeGenerator to set up all the supporting code.

static get_array_name(var, access_data=True)
Get a globally unique name for a ArrayVariable.

   Parameters  var: ArrayVariable
   
   The variable for which a name should be found.

   access_data: bool, optional
   
   For DynamicArrayVariable objects, specifying True here means the name for the underlying data is returned. If specifying False, the name of object itself is returned (e.g. to allow resizing).

   Returns:
   
   name : str
   
   A unique name for var().

get_conditional_write_vars()
Helper function, returns a dict of mappings (varname, condition_var_name) indicating that when varname is written to, it should only be when condition_var_name is True.

has_repeated_indices(statements)
Whether any of the statements potentially uses repeated indices (e.g. pre- or postsynaptic indices).

translate(code, dtype)
Translates an abstract code block into the target language.

translate_expression(expr)
Translate the given expression string into a string in the target language, returns a string.

translate_one_statement_sequence(statements, scalar=False)

translate_statement(statement)
Translate a single line Statement into the target language, returns a string.

translate_statement_sequence(scalar_statements, vector_statements)
Translate a sequence of Statement into the target language, taking care to declare variables, etc. if necessary.
Returns a tuple \((\text{scalar\_code}, \text{vector\_code}, \text{kwds})\) where \(\text{scalar\_code}\) is a list of the lines of code executed before the inner loop, \(\text{vector\_code}\) is a list of the lines of code in the inner loop, and \(\text{kwds}\) is a dictionary of values that is made available to the template.

**cpp_generator module**

**Exported members:** CPPCodeGenerator, c_data_type

**Classes**

<table>
<thead>
<tr>
<th>CPPCodeGenerator(*args, **kwds)</th>
<th>C++ language</th>
</tr>
</thead>
</table>

**CPPCodeGenerator class**

*(Shortest import: from brian2.codegen.generators import CPPCodeGenerator)*

**class** brian2.codegen.generators.cpp_generator.CPPCodeGenerator(*args, **kwds)

**Bases:** brian2.codegen.generators.base.CodeGenerator

C++ language

C++ code templates should provide Jinja2 macros with the following names:

- **main** The main loop.
- **support_code** The support code (function definitions, etc.), compiled in a separate file.

For user-defined functions, there are two keys to provide:

- **support_code** The function definition which will be added to the support code.
- **hashdefine_code** The \#define code added to the main loop.

See TimedArray for an example of these keys.

**Attributes**

| flush_denormals |
| restrict |

**Methods**

<table>
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<th>denormals_to_zero_code()</th>
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</thead>
<tbody>
<tr>
<td>determine_keywords()</td>
</tr>
<tr>
<td>A dictionary of values that is made available to the templated.</td>
</tr>
<tr>
<td>get_array_name(var[, access_data])</td>
</tr>
<tr>
<td>Get a globally unique name for a ArrayVariable.</td>
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<tr>
<td>translate_expression(expr)</td>
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<tr>
<td>Translate the given expression string into a string in the target language, returns a string.</td>
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<tr>
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</tr>
</tbody>
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- **translate_statement_sequence**(*sc_statements,...*)
  Translate a sequence of Statement into the target language, taking care to declare variables, etc.

- **translate_to_declarations**(*read, write, indices*)

- **translate_to_read_arrays**(*read, write, indices*)

- **translate_to_statements**(*statements,...*)

- **translate_to_write_arrays**(*write*)

### Details

- **flush_denormals**

- **restrict**

- **denormals_to_zero_code**()

- **determine_keywords**()
  A dictionary of values that is made available to the templated. This is used for example by the CPPCodeGenerator to set up all the supporting code.

- **static get_array_name**(*var, access_data=True*)
  Get a globally unique name for a ArrayVariable.
  
  **Parameters**
  
  var : ArrayVariable
  The variable for which a name should be found.

  access_data : bool, optional
  For DynamicArrayVariable objects, specifying True here means the name for the underlying data is returned. If specifying False, the name of object itself is returned (e.g. to allow resizing).

  **Returns**

  name : str
  A uniqe name for var().

- **translate_expression**(*expr*)
  Translate the given expression string into a string in the target language, returns a string.

- **translate_one_statement_sequence**(*statements, scalar=False*)

- **translate_statement**(*statement*)
  Translate a single line Statement into the target language, returns a string.

- **translate_statement_sequence**(*sc_statements, ve_statements*)
  Translate a sequence of Statement into the target language, taking care to declare variables, etc. if necessary.

  Returns a tuple (scalar_code, vector_code, kwds) where scalar_code is a list of the lines of code executed before the inner loop, vector_code is a list of the lines of code in the inner loop, and kwds is a dictionary of values that is made available to the template.

- **translate_to_declarations**(*read, write, indices*)

- **translate_to_read_arrays**(*read, write, indices*)

- **translate_to_statements**(*statements, conditional_write_vars*)
Functions

\textbf{\texttt{\texttt{c_data_type}}} (\texttt{\texttt{dtype}}) \hspace{1cm} \text{Gives the C language specifier for numpy data types.}

\begin{itemize}
  \item \textbf{\texttt{\texttt{c_data_type}} function}
  \end{itemize}

\texttt{(Shortest import: from brian2.codegen.generators import \texttt{\texttt{c_data_type}})}

\begin{verbatim}
\texttt{\texttt{brian2.codegen.generators.cpp_generator.c_data_type(dtype)}}
\end{verbatim}

Gives the C language specifier for numpy data types. For example, \texttt{numpy.int32} maps to \texttt{int32\_t} in C.

\textbf{\texttt{\texttt{cython_generator}} module}

\textit{Exported members: \texttt{\texttt{CythonCodeGenerator}}}

\textbf{Classes}

\begin{verbatim}
\texttt{\texttt{CythonCodeGenerator}(*args, **kwds)} \hspace{1cm} \text{Cython code generator}
\end{verbatim}

\textbf{\texttt{\texttt{CythonCodeGenerator}} class}

\texttt{(Shortest import: from brian2.codegen.generators.cython_generator import \texttt{\texttt{CythonCodeGenerator}})}

\begin{verbatim}
\texttt{\texttt{class brian2.codegen.generators.cython_generator.CythonCodeGenerator(*args, **kwds)}}
\end{verbatim}

\texttt{\texttt{Bases: brian2.codegen.generators.base.CodeGenerator}}

\begin{verbatim}
\texttt{\texttt{Cython code generator}}
\end{verbatim}

\textbf{Methods}

\begin{verbatim}
\texttt{\texttt{determine_keywords()}} \hspace{1cm} \text{A dictionary of values that is made available to the templated.}
\texttt{\texttt{translate_expression(expr)}} \hspace{1cm} \text{Translate the given expression string into a string in \texttt{\texttt{the target language}}, returns a string.}
\texttt{\texttt{translate_one_statement_sequence(statements)}} \hspace{1cm} \text{Translate a single line \texttt{\texttt{Statement}} into the target language, returns a string.}
\texttt{\texttt{translate_statement(statement)}} \hspace{1cm} \text{Translate a single line \texttt{\texttt{Statement}} into the target language, returns a string.}
\texttt{\texttt{translate_statement_sequence(sc_statements, ...)}} \hspace{1cm} \text{Translate a sequence of \texttt{\texttt{Statement}} into the target language, taking care to declare variables, etc.}
\texttt{\texttt{translate_to_read_arrays(read, indices)}} \hspace{1cm} \texttt{translate_to_statements(statements, ...)}
\texttt{\texttt{translate_to_write_arrays(write)}}
\end{verbatim}
Details

determine_keywords()
A dictionary of values that is made available to the templated. This is used for example by the
CPPCodeGenerator to set up all the supporting code

translate_expression(expr)
Translate the given expression string into a string in the target language, returns a string.

translate_one_statement_sequence(statements, scalar=False)

translate_statement(statement)
Translate a single line Statement into the target language, returns a string.

translate_statement_sequence(sc_statements, ve_statements)
Translate a sequence of Statement into the target language, taking care to declare variables, etc. if
necessary.

    Returns a tuple (scalar_code, vector_code, kwds) where scalar_code is a list of the lines
    of code executed before the inner loop, vector_code is a list of the lines of code in the inner loop, and
    kwds is a dictionary of values that is made available to the template.

translate_to_read_arrays(read, indices)

translate_to_statements(statements, conditional_write_vars)

translate_to_write_arrays(write)

CythonNodeRenderer([auto_vectorise])

Methods

CythonNodeRenderer class

(Shortest import: from brian2.codegen.generators.cython_generator import
CythonNodeRenderer)

class brian2.codegen.generators.cython_generator.CythonNodeRenderer(auto_vectorise=None)
Bases: brian2.parsing.rendering.NodeRenderer

    Methods

    render_BinOp(node)
    render_Name(node)
    render_NameConstant(node)

Details

    render_BinOp(node)
    render_Name(node)
    render_NameConstant(node)

Functions
get_cpp_dtype(obj)

get_cpp_dtype function

(Shortest import: from brian2.codegen.generators.cython_generator import get_cpp_dtype)

brian2.codegen.generators.cython_generator.get_cpp_dtype(obj)

get_numpy_dtype(obj)

get_numpy_dtype function

(Shortest import: from brian2.codegen.generators.cython_generator import get_numpy_dtype)

brian2.codegen.generators.cython_generator.get_numpy_dtype(obj)

numpy_generator module

Exported members: NumpyCodeGenerator

Classes

class NumpyCodeGenerator(variables, [...][, [...]])

NumpyCodeGenerator class

(Shortest import: from brian2.codegen.generators import NumpyCodeGenerator)

class brian2.codegen.generators.numpy_generator.NumpyCodeGenerator(variables,
variable_indices, owner, iterate_all, codeobj_class, name,
template_name, override_conditional_write=None,
alows_scalar_write=False)

Bases: brian2.codegen.generators.base.CodeGenerator

Numpy language

Essentially Python but vectorised.
Methods

conditional_write(line, stmt, variables, ...)
determine_keywords()  A dictionary of values that is made available to the templated.
read_arrays(read, write, indices, variables, ...)
translate_expression(expr)  Translate the given expression string into a string in the target language, returns a string.
translate_one_statement_sequence(statements)
translate_statement(statement)  Translate a single line Statement into the target language, returns a string.
ufunc_at_vectorisation(statement, variables, ...)
vectorise_code(statements, variables, ...[, ...])
write_arrays(statements, read, write, ...)

Details

conditional_write(line, stmt, variables, conditional_write_vars, created_vars)
determine_keywords()  A dictionary of values that is made available to the templated. This is used for example by the CPPCodeGenerator to set up all the supporting code.
read_arrays(read, write, indices, variables, variable_indices)
translate_expression(expr)  Translate the given expression string into a string in the target language, returns a string.
translate_one_statement_sequence(statements, scalar=False)
translate_statement(statement)  Translate a single line Statement into the target language, returns a string.
ufunc_at_vectorisation(statement, variables, indices, conditional_write_vars, created_vars, used_variables)
vectorise_code(statements, variables, variable_indices, index='idx')
write_arrays(statements, read, write, variables, variable_indices)

VectorisationError

VectorisationError class

(Shortest import: from brian2.codegen.generators.numpy_generator import VectorisationError)  class brian2.codegen.generators.numpy_generator.VectorisationError  Bases: Exception

Functions

ceil_func(value)
**ceil_func function**

(Shortest import: from brian2.codegen.generators.numpy_generator import ceil_func)

brian2.codegen.generators.numpy_generator.ceil_func(value)

**clip_func function**

(Shortest import: from brian2.codegen.generators.numpy_generator import clip_func)

brian2.codegen.generators.numpy_generator.clip_func(array, a_min, a_max)

**floor_func function**

(Shortest import: from brian2.codegen.generators.numpy_generator import floor_func)

brian2.codegen.generators.numpy_generator.floor_func(value)

**int_func function**

(Shortest import: from brian2.codegen.generators.numpy_generator import int_func)

brian2.codegen.generators.numpy_generator.int_func(value)

**poisson_func function**

(Shortest import: from brian2.codegen.generators.numpy_generator import poisson_func)

brian2.codegen.generators.numpy_generator.poisson_func(lam, vectorisation_idx)

**rand_func function**

(Shortest import: from brian2.codegen.generators.numpy_generator import rand_func)

brian2.codegen.generators.numpy_generator.rand_func(vectorisation_idx)
**randn_func**

**randn_func function**

(Shortest import: from brian2.codegen.generators.numpy_generator import randn_func)

brian2.codegen.generators.numpy_generator.randn_func(vectorisation_idx)

**runtime package**

Runtime targets for code generation.

**Subpackages**

**GSLcython_rt package**

**GSLcython_rt module**

Module containing the Cython CodeObject for code generation for integration using the ODE solver provided in the GNU Scientific Library (GSL)

Exported members: GSLCythonCodeObject, IntegrationError

**Classes**

**GSLCompileError**

(Shortest import: from brian2.codegen.runtime.GSLcython_rt.GSLcython_rt import GSLCompileError)

class brian2.codegen.runtime.GSLcython_rt.GSLcython_rt.GSLCompileError
    Bases: Exception

GSLCythonCodeObject(owne, code, variables, ...)

**Methods**

**GSLCythonCodeObject class**

(Shortest import: from brian2.codegen import GSLCythonCodeObject)
class brian2.codegen.runtime.GSLcython_rt.GSLcython_rt.GSLCythonCodeObject

    (owner, code, variables, variable_indices, template_name, template_source, compiler_kwds, name='cython_code_object*)


Methods

    compile()

Details

    compile()

IntegrationError

Error used to signify that GSL was unable to complete integration (only works for cython)

IntegrationError class

(Shortest import: from brian2.codegen import IntegrationError)

class brian2.codegen.runtime.GSLcython_rt.GSLcython_rt.IntegrationError

    Bases: Exception

    Error used to signify that GSL was unable to complete integration (only works for cython)

GSLweave_rt package

GSLweave_rt module

Module containing the Weave CodeObject for code generation for integration using the ODE solver provided in the GNU Scientific Library (GSL)

Exported members: GSLWeaveCodeObject

Classes

GSLCompileError
GSLCompileError class

(Shortest import: from brian2.codegen.runtime.GSLweave_rt.GSLweave_rt import GSLCompileError)

class brian2.codegen.runtime.GSLweave_rt.GSLweave_rt.GSLCompileError
    Bases: Exception

GSLWeaveCodeObject class

(Shortest import: from brian2.codegen import GSLWeaveCodeObject)

class brian2.codegen.runtime.GSLweave_rt.GSLweave_rt.GSLWeaveCodeObject(own
    er, code, variables, ...)

    Bases: brian2.codegen.runtime.weave_rt.weave_rt.WeaveCodeObject

    Methods

    run() Runs the code in the namespace.

    Details

        run ()
        Runs the code in the namespace.

            Returns return_value : dict
            A dictionary with the keys corresponding to the output_variables defined during
            the call of CodeGenerator.code_object.

cython_rt package

cython_rt module

Exported members: CythonCodeObject
Classes

*CythonCodeObject*(owner, code, variables, ...)  Execute code using Cython.

**CythonCodeObject class**

*(Shortest import: from brian2 import CythonCodeObject)*

class brian2.codegen.runtime.cython_rt.cython_rt.CythonCodeObject (owner, code, variables, variable_indices, template_name, template_source, compiler_kwds, name='cython_code_object*')

Bases: brian2.codegen.runtime.numpy_rt.numpy_rt.NumpyCodeObject

Execute code using Cython.

**Methods**

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>compile()</td>
<td></td>
</tr>
<tr>
<td><em>is_available()</em></td>
<td>Whether this target for code generation is available.</td>
</tr>
<tr>
<td>run()</td>
<td>Runs the code in the namespace.</td>
</tr>
<tr>
<td><em>update_namespace()</em></td>
<td>Update the namespace for this timestep.</td>
</tr>
<tr>
<td><em>variables_to_namespace()</em></td>
<td></td>
</tr>
</tbody>
</table>

**Details**

**compile()**

*classmethod is_available()*

Whether this target for code generation is available. Should use a minimal example to check whether code generation works in general.

**run()**

Runs the code in the namespace.

**Returns**

return_value : dict

A dictionary with the keys corresponding to the output_variables defined during the call of CodeGenerator.code_object.

**update_namespace()**

Update the namespace for this timestep. Should only deal with variables where the reference changes every timestep, i.e. where the current reference in namespace is not correct.

**variables_to_namespace()**
**extension_manager module**

Cython automatic extension builder/manager

Inspired by IPython’s Cython cell magics, see: https://github.com/ipython/ipython/blob/master/IPython/extensions/cythonmagic.py

Exported members: cython_extension_manager

Classes

```python
CythonExtensionManager()
```

**Attributes**

**CythonExtensionManager class**

*(Shortest import: from brian2.codegen.runtime.cython_rt.extension_manager import CythonExtensionManager)*

```python
class brian2.codegen.runtime.cython_rt.extension_manager.CythonExtensionManager
Bases: object
```

**Attributes**

```python
so_ext
The extension suffix for compiled modules.
```

**Methods**

```python
create_extension(code[, force, name,...])
```

**Details**

```python
so_ext
The extension suffix for compiled modules.
```

```python
create_extension(code, force=False, name=None, define_macros=None, include_dirs=None, library_dirs=None, runtime_library_dirs=None, extra_compile_args=None, extra_link_args=None, libraries=None, compiler=None, sources=None, owner_name="")
```

**Functions**

```python
get_cython_cache_dir()
```

**get_cython_cache_dir function**

*(Shortest import: from brian2.codegen.runtime.cython_rt.extension_manager import get_cython_cache_dir)*
brian2.codegen.runtime.cython_rt.extension_manager.get_cython_cache_dir

**get_cython_extensions** function

(Shortest import: from brian2.codegen.runtime.cython_rt.extension_manager import get_cython_extensions)

brian2.codegen.runtime.cython_rt.extension_manager.get_cython_extensions()

**simplify_path_env_var** function

(Shortest import: from brian2.codegen.runtime.cython_rt.extension_manager import simplify_path_env_var)

brian2.codegen.runtime.cython_rt.extension_manager.simplify_path_env_var(path)

**Objects**

cython_extension_manager

(Numpy runtime implementation.)

**Preferences**

Numpy runtime codegen preferences

codegen.runtime.numpy.discard_units = False

Whether to change the namespace of user-specified functions to remove units.

**numpy_rt**

Numpy runtime implementation.

**Modules**

Numpy runtime codegen preferences

codegen.runtime.numpy.discard_units = False

Whether to change the namespace of user-specified functions to remove units.
LazyArange

LazyArange(stop[, start, indices])
A class that can be used as a `arange` replacement (with an implied step size of 1) but does not actually create an array of values until necessary.

LazyArange class

(Shortest import: from brian2.codegen.runtime.numpy_rt.numpy_rt import LazyArange)

class brian2.codegen.runtime.numpy_rt.numpy_rt.LazyArange(stop, start=0, indices=None)

Bases: collections.abc.Iterable

A class that can be used as a `arange` replacement (with an implied step size of 1) but does not actually create an array of values until necessary. It is somewhat similar to the `range()` function in Python 3, but does not use a generator. It is tailored to a special use case, the `_vectorisation_idx` variable in numpy templates, and not meant for general use. The `_vectorisation_idx` is used for stateless function calls such as `rand()` and for the numpy codegen target determines the number of values produced by such a call. This will often be the number of neurons or synapses, and this class avoids creating a new array of that size at every code object call when all that is needed is the length of the array.

Examples

```python
>>> from brian2.codegen.runtime.numpy_rt.numpy_rt import LazyArange
>>> ar = LazyArange(10)
>>> len(ar)
10
>>> len(ar[:5])
5
>>> type(ar[:5])
<class 'brian2.codegen.runtime.numpy_rt.numpy_rt.LazyArange'>
>>> ar[5]
5
>>> for value in ar[3:7]:
...    print(value)
...
3
4
5
6
>>> len(ar[np.array([1, 2, 3])])
3
```

NumpyCodeObject

NumpyCodeObject(owner, code, variables,...)
Execute code using Numpy

NumpyCodeObject class

(Shortest import: from brian2 import NumpyCodeObject)
```python
class brian2.codegen.runtime.numpy_rt.numpy_rt.NumpyCodeObject(  
    owner, code, variables, variable_indices,  
    template_name, template_source, compiler_kwds,  
    name='numpy_code_object*')
```

Bases: `brian2.codegen.codeobject.CodeObject`

Execute code using Numpy

Default code for Brian because it works on all platforms.

### Methods

- **compile()**
- **is_available()** Whether this target for code generation is available.
- **run()** Runs the code in the namespace.
- **update_namespace()** Update the namespace for this timestep.
- **variables_to_namespace()**

### Details

**compile()**

```python
classmethod is_available()  
    Whether this target for code generation is available. Should use a minimal example to check whether code  
    generation works in general.
```

**run()**

Runs the code in the namespace.

**Returns** `return_value : dict`

A dictionary with the keys corresponding to the `output_variables` defined during the call of `CodeGenerator.code_object`.

**update_namespace()**

Update the namespace for this timestep. Should only deal with variables where the reference changes every timestep, i.e. where the current reference in `namespace` is not correct.

**variables_to_namespace()**

### weave_rt package

Runtime C++ code generation via weave.

### weave_rt module

Module providing `WeaveCodeObject`.

**Exported members:** `WeaveCodeObject, WeaveCodeGenerator`

**Classes**

6.6. Subpackages
**WeaveCodeGenerator class**

*(Shortest import: from brian2.codegen import WeaveCodeGenerator)*

```python
class brian2.codegen.runtime.weave_rt.WeaveCodeGenerator(*args, **kwds):
    Bases: brian2.codegen.generators.cpp_generator.CPPCodeGenerator
```

**WeaveCodeObject class**

*(Shortest import: from brian2 import WeaveCodeObject)*

```python
class brian2.codegen.runtime.weave_rt.WeaveCodeObject(owner, code, variables, ...):
    Bases: brian2.codegen.codeobject.CodeObject
```

Weave code object

The code should be a `MultiTemplate` object with two macros defined, `main` (for the main loop code) and `support_code` for any support code (e.g. function definitions).

**Methods**

- `compile()`
- `is_available()` Whether this target for code generation is available.
- `run()` Runs the code in the namespace.
- `update_namespace()` Update the namespace for this timestep.
- `variables_to_namespace()`

**Details**

`compile()`

```python
classmethod is_available()
    Whether this target for code generation is available. Should use a minimal example to check whether code generation works in general.
```

`run()` Runs the code in the namespace.

**Returns** `return_value` : `dict`

A dictionary with the keys corresponding to the `output_variables` defined during
the call of CodeGenerator.code_object.

**update_namespace()**
Update the namespace for this timestep. Should only deal with variables where the reference changes every timestep, i.e. where the current reference in namespace is not correct.

**variables_to_namespace()**

Functions

---

**get_weave_cache_dir()**

(Shortest import: from brian2.codegen.runtime.weave_rt.weave_rt import get_weave_cache_dir)

brian2.codegen.runtime.weave_rt.weave_rt.get_weave_cache_dir()

---

**get_weave_extensions()**

(Shortest import: from brian2.codegen.runtime.weave_rt.weave_rt import get_weave_extensions)

brian2.codegen.runtime.weave_rt.weave_rt.get_weave_extensions()

---

**weave_data_type(dtype)**
Gives the C language specifier for numpy data types using weave.

---

**weave_data_type()**

(Shortest import: from brian2.codegen.runtime.weave_rt.weave_rt import weave_data_type)

brian2.codegen.runtime.weave_rt.weave_rt.weave_data_type(dtype)
Gives the C language specifier for numpy data types using weave. For example, numpy.int32 maps to long in C.

---

### 6.6.2 core package

Essential Brian modules, in particular base classes for all kinds of brian objects.

#### Built-in preferences

Core Brian preferences

core.default_float_dtype = float64
Default dtype for all arrays of scalars (state variables, weights, etc.).
core.default_integer_dtype = int32
    Default dtype for all arrays of integer scalars.
core.outdated_dependency_error = True
    Whether to raise an error for outdated dependencies (True) or just a warning (False).

**base module**

*Exported members: BrianObject, BrianObjectException*

**Classes**

| BrianObject(**kwds) | All Brian objects derive from this class, defines magic tracking and update. |

**BrianObject class**

*(Shortest import: from brian2 import BrianObject)*

class brian2.core.base.BrianObject(**kwds)
    Bases: brian2.core.names.Nameable
        All Brian objects derive from this class, defines magic tracking and update.
        See the documentation for Network for an explanation of which objects get updated in which order.

        **Parameters dt:** Quantity, optional
            The time step to be used for the simulation. Cannot be combined with the clock argument.
        
        **clock:** Clock, optional
            The update clock to be used. If neither a clock, nor the dt argument is specified, the defaultclock will be used.
        
        **when:** str, optional
            In which scheduling slot to simulate the object during a time step. Defaults to 'start'.
        
        **order:** int, optional
            The priority of this object for operations occurring at the same time step and in the same scheduling slot. Defaults to 0.
        
        **name:** str, optional
            A unique name for the object - one will be assigned automatically if not provided (of the form brianobject_1, etc.).

    **Notes**

        The set of all BrianObject objects is stored in BrianObject.__instances__().
Attributes

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>_clock</td>
<td>The clock used for simulating this object</td>
</tr>
<tr>
<td>_creation_stack</td>
<td>A string indicating where this object was created (traceback with any parts of Brian code removed)</td>
</tr>
<tr>
<td>_network</td>
<td>Used to remember the Network in which this object has been included before, to raise an error if it is included in a new Network</td>
</tr>
<tr>
<td>_scope_current_key</td>
<td>Global key value for ipython cell restrict magic</td>
</tr>
<tr>
<td>_scope_key</td>
<td>The scope key is used to determine which objects are collected by magic</td>
</tr>
<tr>
<td>active</td>
<td>Whether or not the object should be run</td>
</tr>
<tr>
<td>add_to_magic_network</td>
<td>Whether or not the object should be added to a MagicNetwork.</td>
</tr>
<tr>
<td>clock</td>
<td>The Clock determining when the object should be updated</td>
</tr>
<tr>
<td>code_objects</td>
<td>The list of CodeObject contained within the BrianObject.</td>
</tr>
<tr>
<td>contained_objects</td>
<td>The list of objects contained within the BrianObject.</td>
</tr>
<tr>
<td>invalidates_magic_network</td>
<td>Whether or not MagicNetwork is invalidated when a new BrianObject of this type is added</td>
</tr>
<tr>
<td>name</td>
<td>The unique name for this object.</td>
</tr>
<tr>
<td>order</td>
<td>The order in which objects with the same clock and when should be updated</td>
</tr>
<tr>
<td>updaters</td>
<td>The list of Updater that define the runtime behaviour of this object.</td>
</tr>
<tr>
<td>when</td>
<td>The ID string determining when the object should be updated in Network.run.</td>
</tr>
</tbody>
</table>

Methods

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>add_dependency(obj)</td>
<td>Add an object to the list of dependencies.</td>
</tr>
<tr>
<td>after_run()</td>
<td>Optional method to do work after a run is finished.</td>
</tr>
<tr>
<td>before_run(run_namespace)</td>
<td>Optional method to prepare the object before a run.</td>
</tr>
<tr>
<td>run()</td>
<td></td>
</tr>
</tbody>
</table>

Details

<table>
<thead>
<tr>
<th>Name</th>
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<tr>
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<td>The clock used for simulating this object</td>
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<tr>
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<td>A string indicating where this object was created (traceback with any parts of Brian code removed)</td>
</tr>
<tr>
<td>_network</td>
<td>Used to remember the Network in which this object has been included before, to raise an error if it is included in a new Network</td>
</tr>
<tr>
<td>_scope_current_key</td>
<td>Global key value for ipython cell restrict magic</td>
</tr>
<tr>
<td>_scope_key</td>
<td></td>
</tr>
</tbody>
</table>
The scope key is used to determine which objects are collected by magic

**active**
Whether or not the object should be run.
Inactive objects will not have their `update` method called in `Network.run`. Note that setting or unsetting the `active` attribute will set or unset it for all `contained_objects`.

**add_to_magic_network**
Whether or not the object should be added to a `MagicNetwork`. Note that all objects in `BrianObject.contained_objects` are automatically added when the parent object is added, therefore e.g. `NeuronGroup` should set `add_to_magic_network` to `True`, but it should not be set for all the dependent objects such as `StateUpdater`.

**clock**
The `Clock` determining when the object should be updated.
Note that this cannot be changed after the object is created.

**code_objects**
The list of `CodeObject` contained within the `BrianObject`.
TODO: more details.
Note that this attribute cannot be set directly, you need to modify the underlying list, e.g. `obj.code_objects.extend([A, B])`.

**contained_objects**
The list of objects contained within the `BrianObject`.
When a `BrianObject` is added to a `Network`, its contained objects will be added as well. This allows for compound objects which contain a mini-network structure.
Note that this attribute cannot be set directly, you need to modify the underlying list, e.g. `obj.contained_objects.extend([A, B])`.

**invalidates_magic_network**
Whether or not `MagicNetwork` is invalidated when a new `BrianObject` of this type is added.

**name**
The unique name for this object.
Used when generating code. Should be an acceptable variable name, i.e. starting with a letter character and followed by alphanumeric characters and `_`.

**order**
The order in which objects with the same clock and `when` should be updated.

**updaters**
The list of `Updater` that define the runtime behaviour of this object.
TODO: more details.
Note that this attribute cannot be set directly, you need to modify the underlying list, e.g. `obj.updaters.extend([A, B])`.

**when**
The ID string determining when the object should be updated in `Network.run`.

**add_dependency**(obj)
Add an object to the list of dependencies. Takes care of handling subgroups correctly (i.e., adds its parent object).

**Parameters**

- `obj`: `BrianObject`
The object that this object depends on.

```python
after_run()
```
Optional method to do work after a run is finished.
Called by `Network.after_run` after the main simulation loop terminated.

```python
before_run(run_namespace)
```
Optional method to prepare the object before a run.

```
run()
```

```python
BrianObjectException(message, brianobj, ...)  # High level exception that adds extra Brian-specific information to exceptions
```

**BrianObjectException class**

*(Shortest import: from brian2 import BrianObjectException)*

```python
class brian2.core.base.BrianObjectException(message, brianobj, original_exception)
```
Bases: `Exception`

High level exception that adds extra Brian-specific information to exceptions

This exception should only be raised at a fairly high level in Brian code to pass information back to the user. It adds extra information about where a `BrianObject` was defined to better enable users to locate the source of problems.

You should use the `brian_object_exception` function to raise this, and it should only be raised in an `except` block handling a prior exception.

**Parameters**

- `message` : str
  Additional error information to add to the original exception.

- `brianobj` : BrianObject
  The object that caused the error to happen.

- `original_exception` : Exception
  The original exception that was raised.

**Functions**

```python
brian_object_exception(message, brianobj, ...)
```
Returns a `BrianObjectException` derived from the original exception.

**brian_object_exception function**

*(Shortest import: from brian2.core.base import brian_object_exception)*

```python
brian2.core.base.brian_object_exception(message, brianobj, original_exception)
Returns a `BrianObjectException` derived from the original exception.
```

Creates a new class derived from the class of the original exception and `BrianObjectException`. This allows exception handling code to respond both to the original exception class and `BrianObjectException`.  

6.6. Subpackages
See :class:`BrianObjectException` for arguments and notes.

.. function:: device_override(name)

   Decorates a function/method to allow it to be overridden by the current :class:`Device`.

   .. code:: python

      import brian2
      @brian2.core.base.device_override('example')
      def my_function(arg):
         pass

   The name is the function name in the :class:`Device` to use as an override if it exists.
   The returned function has an additional attribute `original_function` which is a reference to the original, undecorated function.

   Parameters:
   name : str
      An explicit name, if not specified gives an automatically generated name.

.. function:: weakproxy_with_fallback(obj)

   Attempts to create a weakproxy to the object, but falls back to the object if not possible.

   .. code:: python

      from brian2 import weakproxy_with_fallback
      proxy = weakproxy_with_fallback(my_object)

   Parameters:
   obj : any
      The object to wrap.

---

**clocks module**

Clocks for the simulator.

*Exported members: Clock, defaultclock*

**Classes**

.. code:: python

   class Clock(dt[, name])

   An object that holds the simulation time and the time step.

   Parameters:
   dt : float
      The time step of the simulation as a float
   name : str, optional
      An explicit name, if not specified gives an automatically generated name.
Notes

Clocks are run in the same `Network.run` iteration if \( t \) is the same. The condition for two clocks to be considered as having the same time is \( \text{abs}(t1-t2) < \text{epsilon} \times \text{abs}(t1) \), a standard test for equality of floating point values. The value of \text{epsilon} is \( 1e-14 \).

Attributes

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>( dt )</td>
<td>The time step of the simulation in seconds.</td>
</tr>
<tr>
<td>( dt_)</td>
<td>The time step of the simulation as a float (in seconds)</td>
</tr>
<tr>
<td>( \text{epsilon}__\text{dt} )</td>
<td>The relative difference for times (in terms of ( dt )) so that they are considered identical.</td>
</tr>
</tbody>
</table>

Methods

`set_interval(self, start, end)`
Set the start and end time of the simulation.

Details

- \( dt \)
  - The time step of the simulation in seconds.
- \( dt_\)
  - The time step of the simulation as a float (in seconds)
- \( \text{epsilon}_\_\text{dt} \)
  - The relative difference for times (in terms of \( dt \)) so that they are considered identical.

`set_interval(self, start, end)`
Set the start and end time of the simulation.

Sets the start and end value of the clock precisely if possible (using \text{epsilon}) or rounding up if not. This assures that multiple calls to `Network.run` will not re-run the same time step.

Tutorials and examples using this

- Example `COBAHH`
- Example `CUBA`

**DefaultClockProxy class**

(Shortest import: from brian2.core.clocks import DefaultClockProxy)

```python
class brian2.core.clocks.DefaultClockProxy
    Bases: object

    Method proxy to access the defaultclock of the currently active device
```

6.6. Subpackages
Function: **check_dt**

| **check_dt**(new_dt, old_dt, target_t) | Check that the target time can be represented equally well with the new dt. |

**check_dt function**

(SHORTEST IMPORT: from brian2.core.clocks import check_dt)

brian2.core.clocks.check_dt(new_dt, old_dt, target_t)

Check that the target time can be represented equally well with the new dt.

**Parameters**

- **new_dt**: float
  
  The new dt value

- **old_dt**: float
  
  The old dt value

- **target_t**: float
  
  The target time

**Raises**

- **ValueError** If using the new dt value would lead to a difference in the target time of more than Clock.
  
  epsilon_dt times new_dt (by default, 0.01% of the new dt).

**Examples**

```
>>> from brian2 import *
>>> check_dt(float(17*ms), float(0.1*ms), float(0*ms))  # For t=0s, every dt is fine
>>> check_dt(float(0.05*ms), float(0.1*ms), float(10*ms)) # t=10*ms can be represented with the new dt
>>> check_dt(float(0.2*ms), float(0.1*ms), float(10.1*ms)) # t=10.1ms cannot be represented with dt=0.2ms # doctest: +ELLIPSIS
Traceback (most recent call last):
...
ValueError: Cannot set dt from 100. us to 200. us, the time 10.1 ms is not a multiple of 200. us
```

**Objects**

- **defaultclock**: The standard clock, used for objects that do not specify any clock or dt

**defaultclock object**

(SHORTEST IMPORT: from brian2 import defaultclock)

brian2.core.clocks.defaultclock = <brian2.core.clocks.DefaultClockProxy object>

The standard clock, used for objects that do not specify any clock or dt
core_preferences module

Definitions, documentation, default values and validation functions for core Brian preferences.

Functions

   default_float_dtype_validator(dtype)

default_float_dtype_validator function

(Shortest import: from brian2.core.core_preferences import default_float_dtype_validator)

brian2.core.core_preferences.default_float_dtype_validator(dtype)

   dtype_repr(dtype)

dtype_repr function

(Shortest import: from brian2.core.core_preferences import dtype_repr)

brian2.core.core_preferences.dtype_repr(dtype)

functions module

Exported members: DEFAULT_FUNCTIONS, Function, implementation(), declare_types()

Classes

   Function(pyfunc[, sympy_func, arg_units, ...])  An abstract specification of a function that can be used as part of model equations, etc.

Function class

(Shortest import: from brian2 import Function)

class brian2.core.functions.Function(pyfunc, sympy_func=None, arg_units=None, return_unit=None, arg_types=None, return_type=None, stateless=True, auto_vectorise=False)

Bases: object

An abstract specification of a function that can be used as part of model equations, etc.

   Parameters pyfunc : function
       A Python function that is represented by this Function object.

       sympy_func: sympy.Function, optional
       A corresponding sympy function (if any). Allows functions to be interpreted by sympy and potentially make simplifications. For example, sqrt(x**2) could be replaced by abs(x).

       arg_units : list of Unit, optional
If `pyfunc` does not provide unit information (which typically means that it was not annotated with a `check_units()` decorator), the units of the arguments have to specified explicitly using this parameter.

**return_unit**: `Unit` or callable, optional

Same as for `arg_units`: if `pyfunc` does not provide unit information, this information has to be provided explicitly here. `return_unit` can either be a specific `Unit`, if the function always returns the same unit, or a function of the input units, e.g. a “square” function would return the square of its input units, i.e. `return_unit` could be specified as `lambda u: u**2`.

**arg_types**: list of str, optional

Similar to `arg_units`, but gives the type of the argument rather than its unit. In the current version of Brian arguments are specified by one of the following strings: ‘boolean’, ‘integer’, ‘float’, ‘any’. If `arg_types` is not specified, ‘any’ will be assumed. In future versions, a more refined specification may be possible. Note that any argument with a type other than float should have no units. If

**return_type**: str, optional

Similar to `return_unit` and `arg_types`. In addition to ‘boolean’, ‘integer’ and ‘float’ you can also use ‘highest’ which will return the highest type of its arguments. You can also give a function, as for `return_unit`. If the return type is not specified, it is assumed to be ‘float’.

**stateless**: bool, optional

Whether this function does not have an internal state, i.e. if it always returns the same output when called with the same arguments. This is true for mathematical functions but not true for `rand()`, for example. Defaults to True.

**auto_vectorise**: bool, optional

Whether the implementations of this function should get an additional argument (not specified in abstract code) that can be used to determine the number of values that should be returned (for the numpy target), or an index potentially useful for generating deterministic values independent of the order of vectorisation (for all other targets). The main use case are random number functions, e.g. equations refer to `rand()`, but the generate code will actually call `rand(_vectorisation_idx)`. Defaults to False.

**Notes**

If a function should be usable for code generation targets other than Python/numpy, implementations for these target languages have to be added using the `implementation` decorator or using the `add_implementations` function.

**Attributes**

- `implementations`: Stores implementations for this function in a `FunctionImplementationContainer`

**Methods**
__call__(*args) Call self as a function.

is_locally_constant(dt) Return whether this function (if interpreted as a function of time) should be considered constant over a timestep.

Details

implementations
Stores implementations for this function in a FunctionImplementationContainer

__call__(*args)
Call self as a function.

is_locally_constant(dt)
Return whether this function (if interpreted as a function of time) should be considered constant over a timestep. This is most importantly used by TimedArray so that linear integration can be used. In its standard implementation, always returns False.

Parameters dt : float
The length of a timestep (without units).

Returns constant : bool
Whether the results of this function can be considered constant over one timestep of length dt.

FunctionImplementation([name, code, ...]) A simple container object for function implementations.

FunctionImplementation class

(Shortest import: from brian2.core.functions import FunctionImplementation)

class brian2.core.functions.FunctionImplementation (name=None, code=None, namespace=None, dependencies=None, availability_check=None, dynamic=False, compiler_kwds=None)

Bases: object

A simple container object for function implementations.

Parameters name : str, optional
The name of the function in the target language. Should only be specified if the function has to be renamed for the target language.

code : language-dependent, optional
A language dependent argument specifying the implementation in the target language, e.g. a code string or a dictionary of code strings.

namespace : dict-like, optional
A dictionary of mappings from names to values that should be added to the namespace of a CodeObject using the function.

dependencies : dict-like, optional
A mapping of names to Function objects, for additional functions needed by this function.

**availability_check**: callable, optional

A function that will be called to check whether the function should be made available (e.g. depending on whether it is supported by the compiler). The function should do nothing if the function is available, or raise a NotImplementedError with a message explaining why it isn’t.

**dynamic**: bool, optional

Whether this code/namespace is dynamic, i.e. generated for each new context it is used in. If set to True, code and namespace have to be callable with a Group as an argument and are expected to return the final code and namespace. Defaults to False.

### Methods

- `get_code(owner)`
- `get_namespace(owner)`

### Details

- `get_code(owner)`
- `get_namespace(owner)`

**FunctionImplementationContainer**

Helper object to store implementations and give access in a dictionary-like fashion, using CodeGenerator implementations as a fallback for CodeObject implementations.

### FunctionImplementationContainer class

(SHORTEST IMPORT: from brian2.core.functions import FunctionImplementationContainer)

class brian2.core.functions.FunctionImplementationContainer (function)

Bases: collections.abc.Mapping

Helper object to store implementations and give access in a dictionary-like fashion, using CodeGenerator implementations as a fallback for CodeObject implementations.

### Methods

- `add_dynamic_implementation(target, code[...])` Adds an “dynamic implementation” for this function.
- `add_implementation(target, code[...])`
- `add_numpy_implementation(wrapped_func[, ...])` Add a numpy implementation to a Function.
Details

**add_dynamic_implementation** (target, code, namespace=None, dependencies=None, availabil-
ity_check=None, name=None, compiler_kwds=None)

Adds an “dynamic implementation” for this function. code and namespace arguments are expected to
be callables that will be called in `Network.before_run` with the owner of the CodeObject as an
argument. This allows to generate code that depends on details of the context it is run in, e.g. the dt of a
clock.

**add_implementation** (target, code, namespace=None, dependencies=None, availabil-
ity_check=None, name=None, compiler_kwds=None)

**add_numpy_implementation** (wrapped_func, dependencies=None, discard_units=None, com-
piler_kwds=None)

Add a numpy implementation to a `Function`.

**Parameters**

function : `Function`

The function description for which an implementation should be added.

wrapped_func : callable

The original function (that will be used for the numpy implementation)

dependencies : list of `Function`, optional

A list of functions this function needs.

discard_units : bool, optional

See `implementation()`.

---

**SymbolicConstant** (name, sympy_obj, value)  
Class for representing constants (e.g.

---

**SymbolicConstant class**

(Shortest import: from brian2.core.functions import SymbolicConstant)

class brian2.core.functions.SymbolicConstant (name, sympy_obj, value)

    Bases: brian2.core.variables.Constant

    Class for representing constants (e.g. pi) that are understood by sympy.

---

**exprel**  
Represents \((\exp(x) - 1)/x\).

---

**exprel class**

(Shortest import: from brian2.core.functions import exprel)

class brian2.core.functions.exprel

    Bases: sympy.core.function.Function

    Represents \((\exp(x) - 1)/x\).

    The benefit of using `exprel(x)` over \((\exp(x) - 1)/x\) is that the latter is prone to cancellation under finite
    precision arithmetic when \(x\) is close to zero, and cannot be evaluated when \(x\) is equal to zero.
Methods

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>eval(arg)</code></td>
<td>Returns a canonical form of cls applied to arguments args.</td>
</tr>
<tr>
<td><code>fdiff(argindex)</code></td>
<td>Returns the first derivative of this function.</td>
</tr>
</tbody>
</table>

Details

```python
classmethod eval(arg)
```

Returns a canonical form of cls applied to arguments args.

The `eval()` method is called when the class cls is about to be instantiated and it should return either some simplified instance (possible of some other class), or if the class cls should be unmodified, return None.

```python
fdiff(argindex=1)
```

Returns the first derivative of this function.

Tutorials and examples using this

- Example `COBAHH`
- Example `IF_curve_Hodgkin_Huxley`
- Example `compartmental hh_with_spikes`
- Example `compartmental hodgkin_huxley_1952`
- Example `compartmental lfp`
- Example `advanced float_32_64_benchmark`
- Example `advanced exprel_function`
- Example `frompapers Wang_Buszaki_1996`

Functions

```python
declare_types(**types)
```

Decorator to declare argument and result types for a function

Implementation

```python
(Shortest import: from brian2 import declare_types)

brian2.core.functions.declare_types(**types)
```

Decorator to declare argument and result types for a function

Usage is similar to `check_units()` except that types must be one of `{VALID_ARG_TYPES}` and the result type must be one of `{VALID_RETURN_TYPES}`. Unspecified argument types are assumed to be 'all' (i.e. anything is permitted), and an unspecified result type is assumed to be 'float'. Note that the 'highest' option for result type will give the highest type of its argument, e.g. if the arguments were boolean and integer then the result would be integer, if the arguments were integer and float it would be float.

```python
implementation(target[, code, namespace, ...])
```

A simple decorator to extend user-written Python functions to work with code generation in other languages.
implementation function

(Shortest import: from brian2 import implementation)

brian2.core.functions.implementation(target, code=None, namespace=None, dependencies=None, discard_units=None, name=None, **compiler_kwds)

A simple decorator to extend user-written Python functions to work with code generation in other languages.

Parameters:

- **target**: str
  - Name of the code generation target (e.g. 'weave') for which to add an implementation.

- **code**: str or dict-like, optional
  - What kind of code the target language expects is language-specific, e.g. C++ code allows for a dictionary of code blocks instead of a single string.

- **namespaces**: dict-like, optional
  - A namespace dictionary (i.e. a mapping of names to values) that should be added to a CodeObject namespace when using this function.

- **dependencies**: dict-like, optional
  - A mapping of names to Function objects, for additional functions needed by this function.

- **discard_units**: bool, optional
  - Numpy functions can internally make use of the unit system. However, during a simulation run, state variables are passed around as unitless values for efficiency. If discard_units is set to False, input arguments will have units added to them so that the function can still use units internally (the units will be stripped away from the return value as well). Alternatively, if discard_units is set to True, the function will receive unitless values as its input. The namespace of the function will be altered to make references to units (e.g. ms) refer to the corresponding floating point values so that no unit mismatch errors are raised. Note that this system cannot work in all cases, e.g. it does not work with functions that internally imports values (e.g. does from brian2 import ms) or access values with units indirectly (e.g. uses brian2.ms instead of ms). If no value is given, defaults to the preference setting codegen.runtime.numpy.discard_units.

- **name**: str, optional
  - The name of the function in the target language. Should only be specified if the function has to be renamed for the target language.

- **compiler_kwds**: dict, optional
  - Additional keyword arguments will be transferred to the code generation stage, e.g. for C++-based targets, the code can make use of additional header files by providing a list of strings as the headers argument.

Notes

While it is in principle possible to provide a numpy implementation as an argument for this decorator, this is normally not necessary – the numpy implementation should be provided in the decorated function.
If this decorator is used with other decorators such as `check_units()` or `declare_types()`, it should be the uppermost decorator (that is, the last one to be applied).

**Examples**

Sample usage:

```python
@implementation('cpp',"
    #include<math.h>
    inline double usersin(double x)
    {
        return sin(x);
    }
"
)
def usersin(x):
    return sin(x)
```

**timestep function**

*(Shortest import: from brian2.core.functions import timestep)*

`brian2.core.functions.timestep(t, dt)`

Converts a given time to an integer time step. This function slightly shifts the time before dividing it by `dt` to make sure that multiples of `dt` do not end up in the preceding time step due to floating point issues. This function is used in the refractoriness calculation.

New in version 2.1.3.

**Parameters**

- `t` : np.ndarray, float, Quantity
  - The time to convert.
- `dt` : float or Quantity
  - The length of a simulation time step.

**Returns**

- `ts` : np.ndarray, np.int64
  - The time step corresponding to the given time.

**Notes**

This function cannot handle infinity values, use big values instead (e.g. a `NeuronGroup` will use `-1e4*second` as the value of the `lastspike` variable for neurons that never spiked).

**magic module**

*Exported members: MagicNetwork, magic_network, MagicError, run(), stop(), collect(), store(), restore(), start_scope()*

**Classes**
MagicError

Error that is raised when something goes wrong in MagicNetwork

MagicNetwork class

(Shortest import: from brian2 import MagicError)

class brian2.core.magic.MagicError
    Bases: Exception
    Error that is raised when something goes wrong in MagicNetwork
    See notes to MagicNetwork for more details.

MagicNetwork() Network that automatically adds all Brian objects

MagicNetwork class

(Shortest import: from brian2 import MagicNetwork)

class brian2.core.magic.MagicNetwork
    Bases: brian2.core.network.Network
    Network that automatically adds all Brian objects
    In order to avoid bugs, this class will occasionally raise MagicError when the intent of the user is not clear. See the notes below for more details on this point. If you persistently see this error, then Brian is not able to safely guess what you intend to do, and you should use a Network object and call Network.run explicitly.
    Note that this class cannot be instantiated by the user, there can be only one instance magic_network of MagicNetwork.
    See also:
    Network, collect(), run(), stop(), store(), restore()

Notes

All Brian objects that are visible at the point of the run() call will be included in the network. This class is designed to work in the following two major use cases:

1. You create a collection of Brian objects, and call run() to run the simulation. Subsequently, you may call run() again to run it again for a further duration. In this case, the Network.t time will start at 0 and for the second call to run() will continue from the end of the previous run.

2. You have a loop in which at each iteration, you create some Brian objects and run a simulation using them. In this case, time is reset to 0 for each call to run().

In any other case, you will have to explicitly create a Network object yourself and call Network.run on this object. Brian has a built in system to guess which of the cases above applies and behave correctly. When it is not possible to safely guess which case you are in, it raises MagicError. The rules for this guessing system are explained below.

If a simulation consists only of objects that have not been run, it will assume that you want to start a new simulation. If a simulation only consists of objects that have been simulated in the previous run() call, it will continue that simulation at the previous time.

6.6. Subpackages 431
If neither of these two situations apply, i.e., the network consists of a mix of previously run objects and new objects, an error will be raised.

In these checks, “non-invalidating” objects (i.e. objects that have `BrianObject.invalidates_magic_network` set to `False`) are ignored, e.g. creating new monitors is always possible.

Methods

```plaintext
add(*objs)  You cannot add objects directly to MagicNetwork
after_run()  
check_dependencies()  
get_states([units, format, subexpressions, ...])  See Network.get_states.
remove(*objs)  You cannot remove objects directly from MagicNetwork
restore([name, filename, ...])  See Network.restore.
run(duration[, report, report_period, ...])  Runs the simulation for the given duration.
set_states(values[, units, format, level])  See Network.set_states.
store([name, filename, level])  See Network.store.
```

Details

```plaintext
add(*objs)
   You cannot add objects directly to MagicNetwork

after_run()

check_dependencies()

get_states(units=True, format='dict', subexpressions=False, level=0)
   See Network.get_states.

remove(*objs)
   You cannot remove objects directly from MagicNetwork

restore(name='default', filename=None, restore_random_state=False, level=0)
   See Network.restore.

run(duration, report=None, report_period=60*second, namespace=None, level=0)
   Runs the simulation for the given duration.

Parameters

duration: Quantity
   The amount of simulation time to run for.

report: {None, 'text', 'stdout', 'stderr', function}, optional
   How to report the progress of the simulation. If None, do not report progress. If 'text' or 'stdout' is specified, print the progress to stdout. If 'stderr' is specified, print the progress to stderr. Alternatively, you can specify a callback `callable(elapsed, complete, duration)` which will be passed the amount of time elapsed as a `Quantity`, the fraction complete from 0.0 to 1.0 and the total duration of the simulation (in biological time). The function will always be called at the beginning and the end (i.e. for fractions 0.0 and 1.0), regardless of the `report_period`.

report_period: Quantity
```
How frequently (in real time) to report progress.

**namespace**: dict-like, optional

A namespace that will be used in addition to the group-specific namespaces (if defined). If not specified, the locals and globals around the run function will be used.

**profile**: bool, optional

Whether to record profiling information (see `Network.profiling_info`). Defaults to `False`.

**level**: int, optional

How deep to go up the stack frame to look for the locals/global (see `namespace` argument). Only used by run functions that call this run function, e.g. `MagicNetwork.run` to adjust for the additional nesting.

**Notes**

The simulation can be stopped by calling `Network.stop` or the global `stop()` function.

**set_states** *(values, units=True, format='dict', level=0)*

See `Network.set_states`.

**store** *(name='default', filename=None, level=0)*

See `Network.store`.

**Functions**

<table>
<thead>
<tr>
<th><code>collect([level])</code></th>
<th>Return the list of <code>BrianObject</code> s that will be simulated if <code>run()</code> is called.</th>
</tr>
</thead>
</table>

**collect function**

*(Shortest import: from brian2 import collect)*

```python
brian2.core.magic.collect (level=0)
```

Return the list of `BrianObject` s that will be simulated if `run()` is called.

**Parameters**

- **level**: int, optional
  
  How much further up to go in the stack to find the objects. Needs only to be specified if `collect()` is called as part of a function and should be increased by 1 for every level of nesting. Defaults to 0.

**Returns**

- **objects**: set of `BrianObject`
  
  The objects that will be simulated.

**get_objects_in_namespace**(level)

Get all the objects in the current namespace that derive from `BrianObject`.

**get_objects_in_namespace function**

*(Shortest import: from brian2.core.magic import get_objects_in_namespace)*

6.6. Subpackages
brian2.core.magic.get_objects_in_namespace(level)
Get all the objects in the current namespace that derive from BrianObject. Used to determine the objects for the MagicNetwork.

Parameters level : int, optional
How far to go back to get the locals/globals. Each function/method call should add 1 to this argument, functions/method with a decorator have to add 2.

Returns objects : set
A set with weak references to the BrianObjects in the namespace.

restore([name, filename, restore_random_state]) Restore the state of the network and all included objects.

(Shortest import: from brian2 import restore)

brian2.core.magic.restore(name='default', filename=None, restore_random_state=False)
Restore the state of the network and all included objects.

Parameters name : str, optional
The name of the snapshot to restore, if not specified uses 'default'.

filename : str, optional
The name of the file from where the state should be restored. If not specified, it is expected that the state exist in memory (i.e. Network.store was previously called without the filename argument).

restore_random_state : bool, optional
Whether to restore the state of the random number generator. If set to True, going back to an earlier state of the simulation will continue exactly where it left off, even if the simulation is stochastic. If set to False (the default), random numbers are independent between runs (except for explicitly set random seeds), regardless of whether store() / restore() has been used or not. Note that this also restores numpy’s random number generator (since it is used internally by Brian), but it does not restore Python’s builtin random number generator in the random module.

See Also:
——–
Network.restore :

run(duration[, report, report_period, ...]) Runs a simulation with all “visible” Brian objects for the given duration.

(Shortest import: from brian2 import run)

brian2.core.magic.run(duration, report=None, report_period=10*second, namespace=None, level=0)
Runs a simulation with all “visible” Brian objects for the given duration. Calls collect() to gather all the
objects, the simulation can be stopped by calling the global `stop()` function.

In order to avoid bugs, this function will occasionally raise `MagicError` when the intent of the user is not clear. See the notes to `MagicNetwork` for more details on this point. If you persistently see this error, then Brian is not able to safely guess what you intend to do, and you should use a `Network` object and call `Network.run` explicitly.

**Parameters**

- **duration**: `Quantity`
  
  The amount of simulation time to run for. If the network consists of new objects since the last time `run()` was called, the start time will be reset to 0. If `run()` is called twice or more without changing the set of objects, the second and subsequent runs will start from the end time of the previous run. To explicitly reset the time to 0, do `magic_network.t = 0*second`.

- **report**: `{None, ‘stdout’, ‘stderr’, ‘graphical’, function}`, optional
  
  How to report the progress of the simulation. If None, do not report progress. If stdout or stderr is specified, print the progress to stdout or stderr. If graphical, Tkinter is used to show a graphical progress bar. Alternatively, you can specify a callback function `(elapsed, complete)` which will be passed the amount of time elapsed (in seconds) and the fraction complete from 0 to 1.

- **report_period**: `Quantity`
  
  How frequently (in real time) to report progress.

- **profile**: `bool`, optional
  
  Whether to record profiling information (see `Network.profiling_info`). Defaults to False.

- **namespace**: `dict-like`, optional
  
  A namespace in which objects which do not define their own namespace will be run. If not namespace is given, the locals and globals around the run function will be used.

- **level**: `int`, optional
  
  How deep to go down the stack frame to look for the locals/global (see `namespace` argument). Only necessary under particular circumstances, e.g. when calling the run function as part of a function call or lambda expression. This is used in tests, e.g.: `assert_raises(MagicError, lambda: run(1*ms, level=3))`.

**Raises**

- **MagicError** Error raised when it was not possible for Brian to safely guess the intended use. See `MagicNetwork` for more details.

**See also:**

- `Network.run`, `MagicNetwork`, `collect()`, `start_scope()`, `stop()`

---

**start_scope()**

Starts a new scope for magic functions

---

**start_scope function**

*(Shortest import: from brian2 import start_scope)*
brian2.core.magic.start_scope()
    Starts a new scope for magic functions
    All objects created before this call will no longer be automatically included by the magic functions such as run().

stop()
    Stops all running simulations.

stop function

(Shortest import: from brian2 import stop)
brian2.core.magic.stop()
    Stops all running simulations.
    See also:
    Network.stop, run(), reinit

store([name, filename])
    Store the state of the network and all included objects.
store function

(Shortest import: from brian2 import store)
brian2.core.magic.store(name='default', filename=None)
    Store the state of the network and all included objects.
    Parameters name : str, optional
        A name for the snapshot, if not specified uses 'default'.
    filename : str, optional
        A filename where the state should be stored. If not specified, the state will be stored in memory.
    See also:
    Network.store

Objects

magic_network
    Automatically constructed MagicNetwork of all Brian objects

magic_network object

(Shortest import: from brian2 import magic_network)
brian2.core.magic.magic_network = MagicNetwork()
    Automatically constructed MagicNetwork of all Brian objects
names module

Exported members: Nameable

Classes

Nameable(name) Base class to find a unique name for an object

Nameable class

(Shortest import: from brian2 import Nameable)

class brian2.core.names.Nameable(name)
    Bases: brian2.core.tracking.Trackable

    Base class to find a unique name for an object
    
    If you specify a name explicitly, and it has already been taken, a \texttt{ValueError} is raised. You can also specify
    a name with a wildcard asterisk in the end, i.e. in the form '\texttt{name*}'. It will then try \texttt{name} first but if this is
    already specified, it will try \texttt{name_1}, \texttt{name_2}, etc. This is the default mechanism used by most core objects
    in Brian, e.g. \texttt{NeuronGroup} uses a default name of '\texttt{neurongroup*}'.

    \begin{description}
    \item[Parameters name : str]
        An name for the object, possibly ending in \texttt{*} to specify that variants of this name should
        be tried if the name (without the asterisk) is already taken. If (and only if) the name for
        this object has already been set, it is also possible to call the initialiser with \texttt{None} for
        the \texttt{name} argument. This situation can arise when a class derives from multiple classes
        that derive themselves from \texttt{Nameable} (e.g. \texttt{Group} and \texttt{CodeRunner}) and their
        initialisers are called explicitly.
    \end{description}

    \begin{description}
    \item[Raises \texttt{ValueError}]
        If the name is already taken.
    \end{description}

    Attributes

    \begin{description}
    \item[id]
        A unique id for this object.
    \item[name]
        The unique name for this object.
    \end{description}

    Methods

    \begin{description}
    \item[assign_id()]
        Assign a new id to this object.
    \end{description}

    Details

    \begin{description}
    \item[id]
        A unique id for this object.
    \item[name]
        In contrast to names, which may be reused, the id stays unique. This is used in the dependency checking
        to not have to deal with the chore of comparing weak references, weak proxies and strong references.
    \end{description}
The unique name for this object.

Used when generating code. Should be an acceptable variable name, i.e. starting with a letter character and followed by alphanumeric characters and _.

assign_id()

Assign a new id to this object. Under most circumstances, this method should only be called once at the creation of the object to generate a unique id. In the case of the MagicNetwork, however, the id should change when a new, independent set of objects is simulated.

Functions

find_name(name)

find_name function

(Shortest import: from brian2.core.names import find_name)

brian2.core.names.find_name(name)

namespace module

Implementation of the namespace system, used to resolve the identifiers in model equations of NeuronGroup and Synapses

Exported members: get_local_namespace(), DEFAULT_FUNCTIONS, DEFAULT_UNITS, DEFAULT_CONSTANTS

Functions

get_local_namespace(level)

get_local_namespace function

(Shortest import: from brian2 import get_local_namespace)

brian2.core.namespace.get_local_namespace(level)

Get the surrounding namespace.

Parameters level : int, optional

How far to go back to get the locals/globals. Each function/method call should add 1 to this argument, functions/method with a decorator have to add 2.

Returns namespace : dict

The locals and globals at the given depth of the stack frame.

network module

Module defining the Network object, the basis of all simulation runs.
Preferences

Network preferences

core.network.default_schedule = ['start', 'groups', 'thresholds', 'synapses', 'resets', 'end']

Default schedule used for networks that don’t specify a schedule.

Exported members: Network, profiling_summary(), scheduling_summary()

Classes

Network(*objs[, name]) The main simulation controller in Brian

Network class

(Shortest import: from brian2 import Network)

class brian2.core.network.Network(*objs, name='network*')

Bases: brian2.core.names.Nameable

The main simulation controller in Brian

Network handles the running of a simulation. It contains a set of Brian objects that are added with add. The run method actually runs the simulation. The main run loop, determining which objects get called in what order is described in detail in the notes below. The objects in the Network are accesible via their names, e.g. net['neurongroup'] would return the NeuronGroup with this name.

Parameters objs : (BrianObject, container), optional

A list of objects to be added to the Network immediately, see add.

name : str, optional

An explicit name, if not specified gives an automatically generated name

See also:

MagicNetwork, run(), stop()

Notes

The main run loop performs the following steps:

1. Prepare the objects if necessary, see prepare.
2. Determine the end time of the simulation as t + \"duration\". 
3. Determine which set of clocks to update. This will be the clock with the smallest value of t. If there are several with the same value, then all objects with these clocks will be updated simultaneously. Set t to the clock time.
4. If the t value of these clocks is past the end time of the simulation, stop running. If the Network.stop method or the stop() function have been called, stop running. Set t to the end time of the simulation.
5. For each object whose clock is set to one of the clocks from the previous steps, call the update method. This method will not be called if the active flag is set to False. The order in which the objects are called is described below.
6. Increase Clock.t by Clock.dt for each of the clocks and return to step 2.
The order in which the objects are updated in step 4 is determined by the `Network.schedule` and the objects `when` and `order` attributes. The `schedule` is a list of string names. Each `when` attribute should be one of these strings, and the objects will be updated in the order determined by the schedule. The default schedule is `['start', 'groups', 'thresholds', 'synapses', 'resets', 'end']`. In addition to the names provided in the schedule, automatic names starting with `before_` and `after_` can be used. That means that all objects with `when=='before_start'' will be updated first, then those with `when=='start',when=='after_start',when=='before_groups',when=='groups' and so forth. If several objects have the same `when` attribute, then the order is determined by the `order` attribute (lower first).

### Attributes

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<th>Description</th>
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<td>Stored state of objects (store/restore)</td>
</tr>
<tr>
<td><code>objects</code></td>
<td>The set of objects in the Network, should not normally be modified directly.</td>
</tr>
<tr>
<td><code>profiling_info</code></td>
<td>The time spent in executing the various <code>CodeObject</code>s.</td>
</tr>
<tr>
<td><code>schedule</code></td>
<td>List of <code>when</code> slots in the order they will be updated, can be modified.</td>
</tr>
<tr>
<td><code>sorted_objects</code></td>
<td>The sorted objects of this network in the order defined by the schedule.</td>
</tr>
<tr>
<td><code>t</code></td>
<td>Current simulation time in seconds (<code>Quantity</code>)</td>
</tr>
<tr>
<td><code>t_</code></td>
<td>Current time as a float</td>
</tr>
</tbody>
</table>

### Methods

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
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<td><code>add(*objs)</code></td>
<td>Add objects to the Network</td>
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<td><code>after_run()</code></td>
<td>Prepares the Network for a run.</td>
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<td><code>before_run(namespace)</code></td>
<td>Prepares the Network for a run.</td>
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<tr>
<td><code>get_profiling_info(**kwds)</code></td>
<td>The only reason this is not directly implemented in <code>profiling_info</code> is to allow devices (e.g.</td>
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<tr>
<td><code>get_states([units, format, subexpressions, . . .])</code></td>
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<td><code>remove(*objs)</code></td>
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<tr>
<td><code>run(duration[, report, report_period, . . .])</code></td>
<td>Runs the simulation for the given duration.</td>
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<tr>
<td><code>scheduling_summary()</code></td>
<td>Return a <code>SchedulingSummary</code> object, representing the scheduling information for all objects included in the network.</td>
</tr>
<tr>
<td><code>set_states(values[, units, format, level])</code></td>
<td>Set the state variables of objects in the network.</td>
</tr>
<tr>
<td><code>stop()</code></td>
<td>Stops the network from running, this is reset the next time <code>Network.run</code> is called.</td>
</tr>
<tr>
<td><code>store([name, filename])</code></td>
<td>Store the state of the network and all included objects.</td>
</tr>
</tbody>
</table>
Details

_stored_state
Stored state of objects (store/restore)

objects
The set of objects in the Network, should not normally be modified directly. Note that in a MagicNetwork, this attribute only contains the objects during a run: it is filled in before_run and emptied in after_run.

profiling_info
The time spent in executing the various CodeObject s.

A list of (name, time) tuples, containing the name of the CodeObject and the total execution time for simulations of this object (as a Quantity with unit second). The list is sorted descending with execution time.

Profiling has to be activated using the profile keyword in run() or Network.run.

schedule
List of when slots in the order they will be updated, can be modified.

See notes on scheduling in Network. Note that additional when slots can be added, but the schedule should contain at least all of the names in the default schedule: ['start', 'groups', 'thresholds', 'synapses', 'resets', 'end'].

The schedule can also be set to None, resetting it to the default schedule set by the core.network.default_schedule preference.

sorted_objects
The sorted objects of this network in the order defined by the schedule.

Objects are sorted first by their when attribute, and secondly by the order attribute. The order of the when attribute is defined by the schedule. In addition to the slot names defined in the schedule, automatic slot names starting with before_ and after_ can be used (e.g. the slots ['groups', 'thresholds'] allow to use ['before_groups', 'groups', 'after_groups', 'before_thresholds', 'thresholds', 'after_thresholds'])

Final ties are resolved using the objects’ names, leading to an arbitrary but deterministic sorting.

Current simulation time in seconds (Quantity)

Current time as a float

Add objects to the Network

Parameters objs: (BrianObject, container)

The BrianObject or container of Brian objects to be added. Specify multiple objects, or lists (or other containers) of objects. Containers will be added recursively. If the container is a dict then it will add the values from the dictionary but not the keys. If you want to add the keys, do add(objs.keys()).

after_run()

Prepares the Network for a run.

Objects in the Network are sorted into the correct running order, and their BrianObject. before_run methods are called.
Parameters run_namespace: dict-like, optional

A namespace in which objects which do not define their own namespace will be run.

check_dependencies()

get_profiling_info(**kwds)

The only reason this is not directly implemented in profiling_info is to allow devices (e.g. CPPStandaloneDevice) to overwrite this.

get_states(units=True, format='dict', subexpressions=False, read_only_variables=True, level=0)

Return a copy of the current state variable values of objects in the network. The returned arrays are copies of the actual arrays that store the state variable values, therefore changing the values in the returned dictionary will not affect the state variables.

Parameters vars: list of str, optional

The names of the variables to extract. If not specified, extract all state variables (except for internal variables, i.e. names that start with '_'). If the subexpressions argument is True, the current values of all subexpressions are returned as well.

units: bool, optional

Whether to include the physical units in the return value. Defaults to True.

format: str, optional

The output format. Defaults to 'dict'.

subexpressions: bool, optional

Whether to return subexpressions when no list of variable names is given. Defaults to False. This argument is ignored if an explicit list of variable names is given in vars.

read_only_variables: bool, optional

Whether to return read-only variables (e.g. the number of neurons, the time, etc.). Setting it to False will assure that the returned state can later be used with set_states. Defaults to True.

level: int, optional

How much higher to go up the stack to resolve external variables. Only relevant if extracting subexpressions that refer to external variables.

Returns values: dict

A dictionary mapping object names to the state variables of that object, in the specified format.

See also:

VariableOwner.get_states

remove(*objs)

Remove an object or sequence of objects from a Network.

Parameters objs: (BrianObject, container)

The BrianObject or container of Brian objects to be removed. Specify multiple objects, or lists (or other containers) of objects. Containers will be removed recursively.

restore(name='default', filename=None, restore_random_state=False)

Retore the state of the network and all included objects.

Parameters name: str, optional
The name of the snapshot to restore, if not specified uses 'default'.

filename : str, optional

The name of the file from where the state should be restored. If not specified, it is expected that the state exist in memory (i.e. Network.store was previously called without the filename argument).

restore_random_state : bool, optional

Whether to restore the state of the random number generator. If set to True, going back to an earlier state of the simulation will continue exactly where it left off, even if the simulation is stochastic. If set to False (the default), random numbers are independent between runs (except for explicitly set random seeds), regardless of whether store()/restore() has been used or not. Note that this also restores numpy’s random number generator (since it is used internally by Brian), but it does not restore Python’s built-in random number generator in the random module.

run (duration, report=None, report_period=60*second, namespace=None, level=0)

Runs the simulation for the given duration.

Parameters duration : Quantity

The amount of simulation time to run for.


How to report the progress of the simulation. If None, do not report progress. If 'text' or 'stdout' is specified, print the progress to stdout. If 'stderr' is specified, print the progress to stderr. Alternatively, you can specify a callback callable(elapsed, complete, duration) which will be passed the amount of time elapsed as a Quantity, the fraction complete from 0.0 to 1.0 and the total duration of the simulation (in biological time). The function will always be called at the beginning and the end (i.e. for fractions 0.0 and 1.0), regardless of the report_period.

report_period : Quantity

How frequently (in real time) to report progress.

namespace : dict-like, optional

A namespace that will be used in addition to the group-specific namespaces (if defined). If not specified, the locals and globals around the run function will be used.

profile : bool, optional

Whether to record profiling information (see Network.profiling_info). Defaults to False.

level : int, optional

How deep to go up the stack frame to look for the locals/global (see namespace argument). Only used by run functions that call this run function, e.g. MagicNetwork.run to adjust for the additional nesting.

Notes

The simulation can be stopped by calling Network.stop or the global stop() function.
Return a `SchedulingSummary` object, representing the scheduling information for all objects included in the network.

**Returns summary:** `SchedulingSummary`

Object representing the scheduling information.

**set_states** (*values*, *units=True, format='dict', level=0*)

Set the state variables of objects in the network.

**Parameters**

- **values**: dict
  
  A dictionary mapping object names to objects of `format`, setting the states of this object.
  
  - **units**: bool, optional
    
    Whether the `values` include physical units. Defaults to `True`.
  
  - **format**: str, optional
    
    The format of `values`. Defaults to `'dict'`.
  
  - **level**: int, optional
    
    How much higher to go up the stack to resolve external variables. Only relevant when using string expressions to set values.

**See also:**

- `Group.set_states`

**stop** ()

Stops the network from running, this is reset the next time `Network.run` is called.

**store** (*name='default', filename=None*)

Store the state of the network and all included objects.

**Parameters**

- **name**: str, optional
  
  A name for the snapshot, if not specified uses `'default'`.
  
- **filename**: str, optional
  
  A filename where the state should be stored. If not specified, the state will be stored in memory.

**Notes**

The state stored to disk can be restored with the `Network.restore` function. Note that it will only restore the internal state of all the objects (including undelivered spikes) – the objects have to exist already and they need to have the same name as when they were stored. Equations, thresholds, etc. are not stored – this is therefore not a general mechanism for object serialization. Also, the format of the file is not guaranteed to work across platforms or versions. If you are interested in storing the state of a network for documentation or analysis purposes use `Network.get_states` instead.

**Tutorials and examples using this**

- Tutorial `3-intro-to-brian-simulations`
- Example `IF_curve_LIF`
- Example `IF_curve_Hodgkin_Huxley`
- Example `advanced/stochastic_odes`
- Example `advanced/compare_GSL_to_conventional`
- Example `frompapers/Brunel_Wang_2001`

`ProfilingSummary(net[, show])`  
Class to nicely display the results of profiling.

### ProfilingSummary class

`(Shortest import: from brian2.core.network import ProfilingSummary)`

```python
class brian2.core.network.ProfilingSummary(net, show=None)
Bases: object
```

Class to nicely display the results of profiling. Objects of this class are returned by `profiling_summary()`.

**Parameters**

- **net**: `Network`
  The `Network` object to profile.
- **show**: `int`, optional
  The number of results to show (the longest results will be shown). If not specified, all results will be shown.

**See also:**
- `Network.profiling_info`

### SchedulingSummary class

`(Shortest import: from brian2.core.network import SchedulingSummary)`

```python
class brian2.core.network.SchedulingSummary(objects)
Bases: object
```

Object representing the schedule that is used to simulate the objects in a network. Objects of this type are returned by `scheduling_summary()`, they should not be created manually by the user.

**Parameters**

- **objects**: list of `BrianObject`
  The sorted list of objects that are simulated by the network.

### TextReport class

`(Shortest import: from brian2.core.network import TextReport)`

```python
class brian2.core.network.TextReport(stream)
```

Helper object to report simulation progress in `Network.run`.

### 6.6. Subpackages
Helper object to report simulation progress in `Network.run`.

**Parameters**

- **stream**: file
  The stream to write to, commonly `sys.stdout` or `sys.stderr`.

**Methods**

- **__call__(elapsed, completed, start, duration)**: Call self as a function.

**Functions**

- **profiling_summary([net, show])**:
  Returns a `ProfilingSummary` of the profiling info for a run.

  - **Parameters**
    - **net**: `{Network, None}` optional
      The `Network` object to profile, or `magic_network` if not specified.
    - **show**: int
      The number of results to show (the longest results will be shown). If not specified, all results will be shown.

- **schedule_propagation_offset([net])**:
  Returns the minimal time difference for a post-synaptic effect after a spike.

  - **Parameters**
    - **net**: `Network`
      The network to check (uses the magic network if not specified).
Returns offset: Quantity

The minimum spike propagation delay: 0*ms for the standard schedule but \( dt \) for schedules where synapses precedes thresholds.

Notes

This function always returns 0*ms or defaultclock.dt – no attempt is made to deal with other clocks.

\[
scheduling\_summary([\text{net}])
\]

Returns a SchedulingSummary object, representing the scheduling information for all objects included in the given Network (or the “magic” network, if none is specified).

scheduling_summary function

(Shortest import: from brian2 import scheduling_summary)

brian2.core.network.scheduling_summary(net=None)

Returns a SchedulingSummary object, representing the scheduling information for all objects included in the given Network (or the “magic” network, if none is specified). The returned objects can be printed or converted to a string to give an ASCII table representation of the schedule. In a Jupyter notebook, the output can be displayed as a HTML table.

Parameters net: Network, optional

The network for which the scheduling information should be displayed. Defaults to the “magic” network.

Returns summary: SchedulingSummary

An object that represents the scheduling information.

operations module

Exported members: NetworkOperation, network_operation()

Classes

NetworkOperation(function[, dt, clock, ...]) Object with function that is called every time step.

NetworkOperation class

(Shortest import: from brian2 import NetworkOperation)

class brian2.core.operations.NetworkOperation (function, \( dt=None \), clock=None, when='start', order=0, name='networkoperation*')

Bases: brian2.core.base.BrianObject

Object with function that is called every time step.

Parameters function: function

The function to call every time step, should take either no arguments in which case it
is called as `function()` or one argument, in which case it is called with the current
`Clock time (Quantity)`.

**dt**: `Quantity`, optional

The time step to be used for the simulation. Cannot be combined with the `clock`
argument.

**clock**: `Clock`, optional

The update clock to be used. If neither a clock, nor the `dt` argument is specified, the
`defaultclock` will be used.

**when**: `str`, optional

In which scheduling slot to execute the operation during a time step. Defaults to
'start'.

**order**: `int`, optional

The priority of this operation for operations occurring at the same time step and in the
same scheduling slot. Defaults to 0.

See also:

`network_operation()`, `Network`, `BrianObject`

Attributes

**function**

The function to be called each time step

Methods

**run()**

Details

**function**

The function to be called each time step

**run()**

Functions

**network_operation([when])**

Decorator to make a function get called every time step of a simulation.

**network_operation function**

(*Shortest import: from brian2 import network_operation*)

*brian2.core.operations.network_operation(when=\textit{None})*

Decorator to make a function get called every time step of a simulation.

The function being decorated should either have no arguments, or a single argument which will be called with
the current time \( t \).
Parameters **dt**: *Quantity*, optional

The time step to be used for the simulation. Cannot be combined with the **clock** argument.

**clock**: *Clock*, optional

The update clock to be used. If neither a clock, nor the **dt** argument is specified, the **defaultclock** will be used.

**when**: *str*, optional

In which scheduling slot to execute the operation during a time step. Defaults to 'start'.

**order**: *int*, optional

The priority of this operation for operations occurring at the same time step and in the same scheduling slot. Defaults to 0.

See also:

*NetworkOperation*, *Network*, *BrianObject*

Notes

Converts the function into a **NetworkOperation**.

If using the form:

```python
@network_operations(when='end')
def f():
    ...
```

Then the arguments to network_operation must be keyword arguments.

Examples

Print something each time step: ```python
>>> from brian2 import * >>> @network_operation ... def f(): ... print('something') ... >>> net = Network(f)
```

Print the time each time step:

```python
>>> @network_operation ... def f(t): ... print('The time is', t) ... >>> net = Network(f)
```

Specify a dt, etc.:

```python
>>> @network_operation(dt=0.5*ms, when='end') ... def f(): ... print('This will happen at the end of each timestep.') ... >>> net = Network(f)
```
preferences module

Brian global preferences are stored as attributes of a $\text{BrianGlobalPreferences}$ object $\text{prefs}$.  

Exported members: $\text{PreferenceError, BrianPreference, prefs, brianPrefs}$

Classes

$\text{BrianGlobalPreferences}()$  Class of the $\text{prefs}$ object.

BrianGlobalPreferences class

($\text{Shortest import: from brian2.core.preferences import BrianGlobalPreferences}$)

class $\text{brian2.core.preferences.BrianGlobalPreferences}$

Bases: collections.abc.MutableMapping

Class of the $\text{prefs}$ object.

Used for getting/setting/validating/registering preference values. All preferences must be registered via $\text{register_preferences}$. To get or set a preference, you can either use a dictionary-based or an attribute-based interface:

$\text{prefs['core.default_float_dtype']} = \text{float32}$
$\text{prefs.core.default_float_dtype} = \text{float32}$

Preferences can be read from files, see $\text{load_preferences}$ and $\text{read_preference_file}$. Note that $\text{load_preferences}$ is called automatically when Brian has finished importing.

Attributes

<table>
<thead>
<tr>
<th>as_file</th>
<th>Get a Brian preference doc file format string for the current preferences</th>
</tr>
</thead>
<tbody>
<tr>
<td>defaults_as_file</td>
<td>Get a Brian preference doc file format string for the default preferences</td>
</tr>
<tr>
<td>toplevel_categories</td>
<td>The toplevel preference categories</td>
</tr>
</tbody>
</table>

Methods

<table>
<thead>
<tr>
<th>check_all_validated()</th>
<th>Checks that all preferences that have been set have been validated.</th>
</tr>
</thead>
<tbody>
<tr>
<td>do_validation()</td>
<td>Validates preferences that have not yet been validated.</td>
</tr>
<tr>
<td>eval_pref(value)</td>
<td>Evaluate a string preference in the units namespace</td>
</tr>
<tr>
<td>get_documentation([basename, link_targets])</td>
<td>Generates a string documenting all preferences with the given basename.</td>
</tr>
<tr>
<td>load_preferences()</td>
<td>Load all the preference files, but do not validate them.</td>
</tr>
<tr>
<td>read_preference_file(file)</td>
<td>Reads a Brian preferences file</td>
</tr>
<tr>
<td>register_preferences(prefbasename, ...)</td>
<td>Registers a set of preference names, docs and validation functions.</td>
</tr>
</tbody>
</table>

Continued on next page
### Details

**as_file**
Get a Brian preference doc file format string for the current preferences

**defaults_as_file**
Get a Brian preference doc file format string for the default preferences

**toplevel_categories**
The toplevel preference categories

**check_all_validated()**
Checks that all preferences that have been set have been validated.

Logs a warning if not. Should be called by `Network.run` or other key Brian functions.

**do_validation()**
Validates preferences that have not yet been validated.

**eval_pref(value)**
Evaluate a string preference in the units namespace

**get_documentation(basename=None, link_targets=True)**
Generates a string documenting all preferences with the given `basename`. If no `basename` is given, all preferences are documented.

**load_preferences()**
Load all the preference files, but do not validate them.

Preference files are read in the following order:

1. `brian2/default_preferences` from the Brian installation directory.
2. `~/.brian/user_preferences` from the user's home directory
3. `.brian_preferences` from the current directory

Files that are missing are ignored. Preferences read at each step override preferences from previous steps.

#### See also:

**read_preference_file**

**read_preference_file(file)**
Reads a Brian preferences file

The file format for Brian preferences is a plain text file of the form:

```
a.b.c = 1
# Comment line
[a]
b.d = 2
[a.b]
e = 3
```

Blank and comment lines are ignored, all others should be of one of the following two forms:

```
key = value
[section]
```
**eval** is called on the values, so strings should be written as, e.g. `'3'` rather than `3`. The eval is called with all unit names available. Within a section, the section name is prepended to the key. So in the above example, it would give the following unvalidated dictionary:

```
{'a.b.c': 1,
 'a.b.d': 2,
 'a.b.e': 3,
}
```

**Parameters**

- **file**: file, str
  - The file object or filename of the preference file.

**register_preferences** *(prefbasename, prefbasedoc, **prefs)*

Registers a set of preference names, docs and validation functions.

- **Parameters**
  - **prefbasename**: str
    - The base name of the preference.
  - **prefbasedoc**: str
    - Documentation for this base name
  - ****prefs**: dict of (name, BrianPreference) pairs
    - The preference names to be defined. The full preference name will be `prefbasename.name`, and the `BrianPreference` value is used to define the default value, docs, and validation function.

**Raises**

- **PreferenceError** If the base name is already registered.

**See also:**

- BrianPreference

**reset_to_defaults** ()

Resets the parameters to their default values.

**BrianGlobalPreferencesView** *(basename, all_prefs)*

A class allowing for accessing preferences in a subcategory.

**BrianGlobalPreferencesView class**

(Shortest import: from brian2.core.preferences import BrianGlobalPreferencesView)

**class** brian2.core.preferences.BrianGlobalPreferencesView *(basename, all_prefs)*

A class allowing for accessing preferences in a subcategory. It forwards requests to `BrianGlobalPreferences` and provides documentation and autocompletion support for all preferences in the given category. This object is used to allow accessing preferences via attributes of the **prefs** object.

- **Parameters**
  - **basename**: str
The name of the preference category. Has to correspond to a key in BrianGlobalPreferences.pref_register.

**all_prefs**: BrianGlobalPreferences

A reference to the main object storing the preferences.

---

**BrianPreference**(default, docs[, validator, ...]) Used for defining a Brian preference.

### BrianPreference class

(Shortest import: from brian2 import BrianPreference)

```python
class brian2.core.preferences.BrianPreference(default, docs, validator=None, representor=<built-in function repr>)
```

Bases: object

Used for defining a Brian preference.

**Parameters default**: object

The default value.

**docs**: str

Documentation for the preference value.

**validator**: func

A function that True or False depending on whether the preference value is valid or not. If not specified, uses the DefaultValidator for the default value provided (check if the class is the same, and for Quantity objects, whether the units are consistent).

**representor**: func

A function that returns a string representation of a valid preference value that can be passed to `eval`. By default, uses `repr` which works in almost all cases.

---

**DefaultValidator**(value) Default preference validator

### DefaultValidator class

(Shortest import: from brian2.core.preferences import DefaultValidator)

```python
class brian2.core.preferences.DefaultValidator(value)
```

Bases: object

Default preference validator

Used by BrianPreference as the default validator if none is given. First checks if the provided value is of the same class as the default value, and then if the default is a Quantity, checks that the units match.

### Methods

**__call__**(value) Call self as a function.
Details

__call__(value)
Call self as a function.

ErrorRaiser class

(Shortest import: from brian2.core.preferences import ErrorRaiser)

class brian2.core.preferences.ErrorRaiser
    Bases: object

PreferenceError class

(Shortest import: from brian2 import PreferenceError)

class brian2.core.preferences.PreferenceError
    Bases: Exception
    Exception relating to the Brian preferences system.

Functions

check_preference_name(name) Make sure that a preference name is valid.

check_preference_name function

(Shortest import: from brian2.core.preferences import check_preference_name)

brian2.core.preferences.check_preference_name(name)
    Make sure that a preference name is valid. This currently checks that the name does not contain illegal characters and does not clash with method names such as “keys” or “items”.

    Parameters name : str
        The name to check.

    Raises

        PreferenceError In case the name is invalid.

parse_preference_name(name) Split a preference name into a base and end name.

parse_preference_name function

(Shortest import: from brian2.core.preferences import parse_preference_name)
brian2.core.preferences.parse_preference_name(name)
Split a preference name into a base and end name.

**Parameters**

name : str
The full name of the preference.

**Returns**

basename : str
The first part of the name up to the final ..

dename : str
The last part of the name from the final . onwards.

**Examples**

```python
>>> parse_preference_name('core.weave_compiler')
('core', 'weave_compiler')
>>> parse_preference_name('codegen.cpp.compiler')
('codegen.cpp', 'compiler')
```

**Objects**

brian_prefs

**brian_prefs object**

(Shortest import: from brian2 import brian_prefs)

brian2.core.preferences.brian_prefs = <brian2.core.preferences.ErrorRaiser object>

```python
prefs
```

**prefs object**

(Shortest import: from brian2 import prefs)

brian2.core.preferences.prefs = <BrianGlobalPreferences with top-level categories:  "logging", "GSL", "devices", "codegen", "core", "legacy">

Preference categories:

**logging** Logging system preferences
**GSL** Directory containing GSL code
**devices** Device preferences
**codegen** Code generation preferences
**core** Core Brian preferences
**legacy** Preferences to enable legacy behaviour

**spikesource module**

Exported members: SpikeSource

---

6.6. Subpackages
Classes

<table>
<thead>
<tr>
<th><strong>SpikeSource</strong></th>
<th>A source of spikes.</th>
</tr>
</thead>
</table>

**SpikeSource class**

*(Shortest import: from brian2 import SpikeSource)*

class brian2.core.spikesource.SpikeSource

Bases: object

A source of spikes.

An object that can be used as a source of spikes for objects such as SpikeMonitor, Synapses, etc.

The defining properties of SpikeSource are that it should have:

- A length that can be extracted with `len(obj)`, where the maximum spike index possible is `len(obj)-1`.
- An attribute `spikes`, an array of ints each from 0 to `len(obj)-1` with no repeats (but possibly not in sorted order). This should be updated each time step.
- A `clock` attribute, this will be used as the default clock for objects with this as a source.

**spikes**

An array of ints, each from 0 to `len(obj)-1` with no repeats (but possibly not in sorted order). Updated each time step.

**clock**

The clock on which the spikes will be updated.

**tracking module**

Expoted members: Trackable

Classes

<table>
<thead>
<tr>
<th><strong>InstanceFollower</strong></th>
<th>Keep track of all instances of classes derived from Trackable</th>
</tr>
</thead>
</table>

**InstanceFollower class**

*(Shortest import: from brian2.core.tracking import InstanceFollower)*

class brian2.core.tracking.InstanceFollower

Bases: object

Keep track of all instances of classes derived from Trackable

The variable `__instancesets__` is a dictionary with keys which are class objects, and values which are InstanceTrackerSet, so `__instanceset__[cls]` is a set tracking all of the instances of class `cls` (or a subclass).
Methods

```
add(value)
get(cls)
```

Details

```
add(value)
get(cls)
```

### InstanceTrackerSet class

*(Shortest import: from brian2.core.tracking import InstanceTrackerSet)*

```python
class brian2.core.tracking.InstanceTrackerSet
    Bases: set
    A set of weakref.ref to all existing objects of a certain class.
    Should not normally be directly used.
```

Methods

```
add(value)    Adds a weakref.ref to the value
remove(value) Removes the value (which should be a weakref) if it is in the set
```

Details

```
add(value)    Adds a weakref.ref to the value
remove(value) Removes the value (which should be a weakref) if it is in the set
```

Sometimes the value will have been removed from the set by clear, so we ignore KeyError in this case.

### Trackable class

*(Shortest import: from brian2 import Trackable)*

```python
class brian2.core.tracking.Trackable
    Bases: object
```

**Classes derived from this will have their instances tracked.**
Classes derived from this will have their instances tracked.

The \texttt{classmethod \_\_\_instances\_\_()} will return a \texttt{InstanceTrackerSet} of the instances of that class, and its subclasses.

\textbf{variables module}

\textbf{Exported members:} \texttt{Variable, Constant, ArrayVariable, DynamicArrayVariable, Subexpression, AuxiliaryVariable, VariableView, Variables, LinkedVariable, linked\_var()}

\textbf{Classes}

\begin{description}
\item[\texttt{ArrayVariable}(name, owner, size, device[, ...])] An object providing information about a model variable stored in an array (for example, all state variables).
\end{description}

\textbf{ArrayVariable class}

\texttt{(Shortest import: from brian2.core.variables import ArrayVariable)}

\begin{verbatim}
class brian2.core.variables.ArrayVariable(name, owner, size, device, dimensions=Dimension(), dtype=None, constant=False, scalar=False, read_only=False, dynamic=False, unique=False)

Bases: brian2.core.variables.Variable
\end{verbatim}

An object providing information about a model variable stored in an array (for example, all state variables). Most of the time \texttt{Variables.add_array} should be used instead of instantiating this class directly.

\begin{description}
\item[\texttt{name} : \texttt{str}]
The name of the variable. Note that this refers to the \textit{original} name in the owning group. The same variable may be known under other names in other groups (e.g. the variable \texttt{v} of a NeuronGroup is known as \texttt{v\_post} in a Synapse connecting to the group).
\item[\texttt{dimensions} : Dimension, optional]
The physical dimensions of the variable
\item[\texttt{owner} : Nameable]
The object that “owns” this variable, e.g. the NeuronGroup or Synapses object that declares the variable in its model equations.
\item[\texttt{size} : int]
The size of the array
\item[\texttt{device} : Device]
The device responsible for the memory access.
\item[\texttt{dtype} : dtype, optional]
The dtype used for storing the variable. If none is given, defaults to \texttt{core.default_float_dtype}.
\item[\texttt{constant} : bool, optional]
Whether the variable’s value is constant during a run. Defaults to \texttt{False}.
\end{description}
scalar : bool, optional

Whether this array is a 1-element array that should be treated like a scalar (e.g. for a single delay value across synapses). Defaults to False.

read_only : bool, optional

Whether this is a read-only variable, i.e. a variable that is set internally and cannot be changed by the user. Defaults to False.

unique : bool, optional

Whether the values in this array are all unique. This information is only important for variables used as indices and does not have to reflect the actual contents of the array but only the possibility of non-uniqueness (e.g. synaptic indices are always unique but the corresponding pre- and post-synaptic indices are not). Defaults to False.

Attributes

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>conditional_write</td>
<td>Another variable, on which the write is conditioned (e.g. device)</td>
</tr>
<tr>
<td>device</td>
<td>The Device responsible for memory access.</td>
</tr>
<tr>
<td>size</td>
<td>The size of this variable.</td>
</tr>
<tr>
<td>unique</td>
<td>Whether all values in this arrays are necessarily unique (only relevant for index variables).</td>
</tr>
</tbody>
</table>

Methods

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>get_addressable_value(name, group)</td>
<td>Get the value (without units) of this variable in a form that can be indexed in the context of a group.</td>
</tr>
<tr>
<td>get_addressable_value_with_unit(name, group)</td>
<td>Get the value (with units) of this variable in a form that can be indexed in the context of a group.</td>
</tr>
<tr>
<td>get_len()</td>
<td>Get the length of the value associated with the variable or 0 for a scalar variable.</td>
</tr>
<tr>
<td>get_value()</td>
<td>Return the value associated with the variable (without units).</td>
</tr>
<tr>
<td>set_conditional_write(var)</td>
<td>Set the value associated with the variable.</td>
</tr>
</tbody>
</table>

Details

conditional_write

Another variable, on which the write is conditioned (e.g. a variable denoting the absence of refractoriness)

device

The Device responsible for memory access.

size

The size of this variable.

unique

Whether all values in this arrays are necessarily unique (only relevant for index variables).

get_addressable_value(name, group)

Get the value (without units) of this variable in a form that can be indexed in the context of a group. For
example, if a postsynaptic variable $x$ is accessed in a synapse $S$ as $S.x_{\text{post}}$, the synaptic indexing scheme can be used.

**Parameters**

- **name**: str
  
  The name of the variable

- **group**: Group
  
  The group providing the context for the indexing. Note that this group is not necessarily the same as $\text{Variable.owner}$: a variable owned by a NeuronGroup can be indexed in a different way if accessed via a Synapses object.

**Returns**

- **variable**: object
  
  The variable in an indexable form (without units).

**get_addressable_value_with_unit** *(name, group)*

Get the value (with units) of this variable in a form that can be indexed in the context of a group. For example, if a postsynaptic variable $x$ is accessed in a synapse $S$ as $S.x_{\text{post}}$, the synaptic indexing scheme can be used.

**Parameters**

- **name**: str
  
  The name of the variable

- **group**: Group
  
  The group providing the context for the indexing. Note that this group is not necessarily the same as $\text{Variable.owner}$: a variable owned by a NeuronGroup can be indexed in a different way if accessed via a Synapses object.

**Returns**

- **variable**: object
  
  The variable in an indexable form (with units).

**get_len** *

Get the length of the value associated with the variable or 0 for a scalar variable.

**get_value** *

Return the value associated with the variable (without units). This is the way variables are accessed in generated code.

**set_conditional_write**(var)

**set_value**(value)

Set the value associated with the variable.

---

**AuxiliaryVariable** *(name[, dimensions, dtype, ...]*)

Variable description for an auxiliary variable (most likely one that is added automatically to abstract code, e.g.)

---

**AuxiliaryVariable class**

*(Shortest import: from brian2.core.variables import AuxiliaryVariable)*

**class** brian2.core.variables.AuxiliaryVariable *(name, dimensions=Dimension(), dtype=None, scalar=False)*

Bases: brian2.core.variables.Variable

Variable description for an auxiliary variable (most likely one that is added automatically to abstract code, e.g. _cond for a threshold condition), specifying its type and unit for code generation. Most of the time
Variables.add_auxiliary_variable should be used instead of instantiating this class directly.

**Parameters**

- **name**: str
  
  The name of the variable

- **dimensions**: Dimension, optional
  
  The physical dimensions of the variable.

- **dtype**: dtype, optional
  
  The dtype used for storing the variable. If none is given, defaults to `core.default_float_dtype`.

- **scalar**: bool, optional
  
  Whether the variable is a scalar value (`True`) or vector-valued, e.g. defined for every neuron (`False`). Defaults to `False`.

**Methods**

- **get_value()**: Return the value associated with the variable (without units).

**Details**

**get_value()**

Return the value associated with the variable (without units). This is the way variables are accessed in generated code.

**Constant**(name, value[, dimensions, owner])

A scalar constant (e.g.

**Constant class**

(Shortest import: from brian2.core.variables import Constant)

```python
class brian2.core.variables.Const...]
owner=...)
```

A scalar constant (e.g. the number of neurons `N`). Information such as the dtype or whether this variable is a boolean are directly derived from the `value`. Most of the time `Variables.add_constant` should be used instead of instantiating this class directly.

**Parameters**

- **name**: str
  
  The name of the variable

- **dimensions**: Dimension, optional
  
  The physical dimensions of the variable. Note that the variable itself (as referenced by `value`) should never have units attached.

- **value**: reference to the variable value:
  
  The value of the constant.

- **owner**: Nameable, optional
The object that “owns” this variable, for constants that belong to a specific group, e.g. the N constant for a NeuronGroup. External constants will have None (the default value).

Attributes

| value | The constant’s value |

Methods

| get_value() | Return the value associated with the variable (without units). |

Details

| value | The constant’s value |
| get_value() | Return the value associated with the variable (without units). This is the way variables are accessed in generated code. |

DynamicArrayVariable (name, owner, size, device)

An object providing information about a model variable stored in a dynamic array (used in Synapses).

DynamicArrayVariable class

(Shortest import: from brian2.core.variables import DynamicArrayVariable)

class brian2.core.variables.DynamicArrayVariable (name, owner, size, device, dimensions=Dimension(), dtype=None, constant=False, needs_reference_update=False, resize_along_first=False, scalar=False, read_only=False, unique=False)

Bases: brian2.core.variables.ArrayVariable

An object providing information about a model variable stored in a dynamic array (used in Synapses). Most of the time Variables.add_dynamic_array should be used instead of instantiating this class directly.

Parameters

name : ‘str’

The name of the variable. Note that this refers to the original name in the owning group. The same variable may be known under other names in other groups (e.g. the variable v of a NeuronGroup is known as v_post in a Synapse connecting to the group).

dimensions: Dimension, optional

The physical dimensions of the variable.

owner: Nameable
The object that “owns” this variable, e.g. the NeuronGroup or Synapses object that declares the variable in its model equations.

**size**: int or tuple of int

The (initial) size of the variable.

**device**: Device

The device responsible for the memory access.

**dtype**: dtype, optional

The dtype used for storing the variable. If none is given, defaults to `core.default_float_dtype`.

**constant**: bool, optional

Whether the variable’s value is constant during a run. Defaults to False.

**needs_reference_update**: bool, optional

Whether the code objects need a new reference to the underlying data at every time step. This should be set if the size of the array can be changed by other code objects. Defaults to False.

**scalar**: bool, optional

Whether this array is a 1-element array that should be treated like a scalar (e.g. for a single delay value across synapses). Defaults to False.

**read_only**: bool, optional

Whether this is a read-only variable, i.e. a variable that is set internally and cannot be changed by the user. Defaults to False.

**unique**: bool, optional

Whether the values in this array are all unique. This information is only important for variables used as indices and does not have to reflect the actual contents of the array but only the possibility of non-uniqueness (e.g. synaptic indices are always unique but the corresponding pre- and post-synaptic indices are not). Defaults to False.

### Attributes

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>dimensions</strong></td>
<td>The number of dimensions</td>
</tr>
<tr>
<td><strong>ndim</strong></td>
<td></td>
</tr>
<tr>
<td><strong>needs_reference_update</strong></td>
<td>Whether this variable needs an update of the reference to the underlying data whenever it is passed to a code object</td>
</tr>
<tr>
<td><strong>resize_along_first</strong></td>
<td>Whether this array will be only resized along the first dimension</td>
</tr>
</tbody>
</table>

### Methods

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>resize(new_size)</strong></td>
<td>Resize the dynamic array.</td>
</tr>
</tbody>
</table>
Details

dimensions

ndim

The number of dimensions

needs_reference_update

Whether this variable needs an update of the reference to the underlying data whenever it is passed to a code object

resize_along_first

Whether this array will be only resized along the first dimension

resize(new_size)

Resize the dynamic array. Calls self.device.resize to do the actual resizing.

Parameters new_size : int or tuple of int

The new size.

LinkedVariable(group, name, variable[, index]) A simple helper class to make linking variables explicit.

LinkedVariable class

(Shortest import: from brian2.core.variables import LinkedVariable)

class brian2.core.variables.LinkedVariable(group, name, variable, index=None)

Bases: object

A simple helper class to make linking variables explicit. Users should use linked_var() instead.

Parameters group : Group

The group through which the variable is accessed (not necessarily the same as variable.owner.

name : str

The name of variable in group (not necessarily the same as variable.name).

variable : Variable

The variable that should be linked.

index : str or ndarray, optional

An indexing array (or the name of a state variable), providing a mapping from the entries in the link source to the link target.

Subexpression(name, owner, expr, device[...]) An object providing information about a named subexpression in a model.

Subexpression class

(Shortest import: from brian2.core.variables import Subexpression)
**class** brian2.core.variables.Subexpression(name, owner, expr, device, dimensions=Dimension(), dtype=None, scalar=False)

**Bases:** brian2.core.variables.Variable

An object providing information about a named subexpression in a model. Most of the time Variables.add_subexpression should be used instead of instantiating this class directly.

**Parameters**

- **name** : str
  The name of the subexpression.

- **dimensions** : Dimension, optional
  The physical dimensions of the subexpression.

- **owner** : Group
  The group to which the expression refers.

- **expr** : str
  The subexpression itself.

- **device** : Device
  The device responsible for the memory access.

- **dtype** : dtype, optional
  The dtype used for the expression. Defaults to core.default_float_dtype.

- **scalar** : bool, optional
  Whether this is an expression only referring to scalar variables. Defaults to False

**Attributes**

- **device**
  The Device responsible for memory access

- **expr**
  The expression defining the subexpression

- **identifiers**
  The identifiers used in the expression

**Methods**

- **get_addressable_value**(name, group)
  Get the value (without units) of this variable in a form that can be indexed in the context of a group.

- **get_addressable_value_with_unit**(name, group)
  Get the value (with units) of this variable in a form that can be indexed in the context of a group.

**Details**

- **device**
  The Device responsible for memory access

- **expr**
  The expression defining the subexpression

- **identifiers**
  The identifiers used in the expression
**get_addressable_value**(name, group)

Get the value (without units) of this variable in a form that can be indexed in the context of a group. For example, if a postsynaptic variable $x$ is accessed in a synapse $S$ as $S.x_{post}$, the synaptic indexing scheme can be used.

**Parameters**

- **name**: str
  - The name of the variable

- **group**: Group
  - The group providing the context for the indexing. Note that this group is not necessarily the same as Variable.owner: a variable owned by a NeuronGroup can be indexed in a different way if accessed via a Synapses object.

**Returns**

- **variable**: object
  - The variable in an indexable form (without units).

**get_addressable_value_with_unit**(name, group)

Get the value (with units) of this variable in a form that can be indexed in the context of a group. For example, if a postsynaptic variable $x$ is accessed in a synapse $S$ as $S.x_{post}$, the synaptic indexing scheme can be used.

**Parameters**

- **name**: str
  - The name of the variable

- **group**: Group
  - The group providing the context for the indexing. Note that this group is not necessarily the same as Variable.owner: a variable owned by a NeuronGroup can be indexed in a different way if accessed via a Synapses object.

**Returns**

- **variable**: object
  - The variable in an indexable form (with units).

---

**Variable**(name[, dimensions, owner, dtype, ...])

An object providing information about model variables (including implicit variables such as $t$ or $xi$).

---

**Variable class**

(Shortest import: from brian2.core.variables import Variable)

```python
class brian2.core.variables.Variable(name, dimensions=Dimension(), owner=None, dtype=None, scalar=False, constant=False, read_only=False, dynamic=False, array=False)
```

Bases: brian2.utils.caching.CacheKey

An object providing information about model variables (including implicit variables such as $t$ or $xi$). This class should never be instantiated outside of testing code, use one of its subclasses instead.

**Parameters**

- **name**: 'str'
  - The name of the variable. Note that this refers to the original name in the owning group.
  - The same variable may be known under other names in other groups (e.g. the variable $v$ of a NeuronGroup is known as $v_{post}$ in a Synapse connecting to the group).

- **dimensions**: Dimension, optional
  - The physical dimensions of the variable.
owner : Nameable, optional

The object that “owns” this variable, e.g. the NeuronGroup or Synapses object that declares the variable in its model equations. Defaults to None (the value used for Variable objects without an owner, e.g. external Constants).

dtype : dtype, optional

The dtype used for storing the variable. Defaults to the preference core. default_scalar.dtype.

scalar : bool, optional

Whether the variable is a scalar value (True) or vector-valued, e.g. defined for every neuron (False). Defaults to False.

constant : bool, optional:

Whether the value of this variable can change during a run. Defaults to False.

read_only : bool, optional

Whether this is a read-only variable, i.e. a variable that is set internally and cannot be changed by the user (this is used for example for the variable N, the number of neurons in a group). Defaults to False.

array : bool, optional

Whether this variable is an array. Allows for simpler check than testing isinstance(var, ArrayVariable). Defaults to False.

Attributes

<table>
<thead>
<tr>
<th>attribute</th>
<th>description</th>
</tr>
</thead>
<tbody>
<tr>
<td>array</td>
<td>Whether the variable is an array</td>
</tr>
<tr>
<td>constant</td>
<td>Whether the variable is constant during a run</td>
</tr>
<tr>
<td>dim</td>
<td>The variable’s dimensions.</td>
</tr>
<tr>
<td>dtype</td>
<td>The dtype used for storing the variable.</td>
</tr>
<tr>
<td>dtype_str</td>
<td>String representation of the numpy dtype</td>
</tr>
<tr>
<td>dynamic</td>
<td>Whether the variable is dynamically sized (only for non-scalars)</td>
</tr>
<tr>
<td>is_boolean</td>
<td></td>
</tr>
<tr>
<td>is_integer</td>
<td></td>
</tr>
<tr>
<td>name</td>
<td>The variable’s name.</td>
</tr>
<tr>
<td>owner</td>
<td>The Group to which this variable belongs.</td>
</tr>
<tr>
<td>read_only</td>
<td>Whether the variable is read-only</td>
</tr>
<tr>
<td>scalar</td>
<td>Whether the variable is a scalar</td>
</tr>
<tr>
<td>unit</td>
<td>The Unit of this variable</td>
</tr>
</tbody>
</table>

Methods

<table>
<thead>
<tr>
<th>method</th>
<th>description</th>
</tr>
</thead>
<tbody>
<tr>
<td>get_addressable_value(name, group)</td>
<td>Get the value (without units) of this variable in a form that can be indexed in the context of a group.</td>
</tr>
<tr>
<td>get_addressable_value_with_unit(name, group)</td>
<td>Get the value (with units) of this variable in a form that can be indexed in the context of a group.</td>
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6.6. Subpackages
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<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
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<tbody>
<tr>
<td><code>get_len()</code></td>
<td>Get the length of the value associated with the variable or 0 for a scalar variable.</td>
</tr>
<tr>
<td><code>get_value()</code></td>
<td>Return the value associated with the variable (without units).</td>
</tr>
<tr>
<td><code>get_value_with_unit()</code></td>
<td>Return the value associated with the variable (with units).</td>
</tr>
<tr>
<td><code>set_value(value)</code></td>
<td>Set the value associated with the variable.</td>
</tr>
</tbody>
</table>

**Details**

- **array**
  Whether the variable is an array

- **constant**
  Whether the variable is constant during a run

- **dim**
  The variable’s dimensions.

- **dtype**
  The dtype used for storing the variable.

- **dtype_str**
  String representation of the numpy dtype

- **dynamic**
  Whether the variable is dynamically sized (only for non-scalars)

- **is_boolean**

- **is_integer**

- **name**
  The variable’s name.

- **owner**
  The Group to which this variable belongs.

- **read_only**
  Whether the variable is read-only

- **scalar**
  Whether the variable is a scalar

- **unit**
  The Unit of this variable

**get_addressable_value**(name, group)
Get the value (without units) of this variable in a form that can be indexed in the context of a group. For example, if a postsynaptic variable \( x \) is accessed in a synapse \( S \) as \( S.x\_post \), the synaptic indexing scheme can be used.

**Parameters**

- **name**: str
  The name of the variable

- **group**: Group
  The group providing the context for the indexing. Note that this group is not necessarily the same as `Variable.owner`: a variable owned by a `NeuronGroup` can be indexed in a different way if accessed via a `Synapses` object.
Returns variable : object

The variable in an indexable form (without units).

get_addressable_value_with_unit (name, group)
Get the value (with units) of this variable in a form that can be indexed in the context of a group. For example, if a postsynaptic variable $x$ is accessed in a synapse $S$ as $S.x_{\text{post}}$, the synaptic indexing scheme can be used.

Parameters name : str
The name of the variable

group : Group
The group providing the context for the indexing. Note that this group is not necessarily the same as `Variable.owner`: a variable owned by a `NeuronGroup` can be indexed in a different way if accessed via a `Synapses` object.

Returns variable : object
The variable in an indexable form (with units).

get_len ()
Get the length of the value associated with the variable or 0 for a scalar variable.

get_value ()
Return the value associated with the variable (without units). This is the way variables are accessed in generated code.

get_value_with_unit ()
Return the value associated with the variable (with units).

set_value (value)
Set the value associated with the variable.

VariableView (name, variable, group[, dimensions])
A view on a variable that allows to treat it as an numpy array while allowing special indexing (e.g.

VariableView class

(Shortest import: from brian2.core.variables import VariableView)

class brian2.core.variables.VariableView (name, variable, group, dimensions=None)

Bases: object
A view on a variable that allows to treat it as an numpy array while allowing special indexing (e.g. with strings) in the context of a Group.

Parameters name : str
The name of the variable (not necessarily the same as `variable.name`).

variable : Variable
The variable description.

group : Group
The group through which the variable is accessed (not necessarily the same as `variable.owner`).

dimensions : Dimension, optional
The physical dimensions to be used for the variable, should be \texttt{None} when a variable is accessed without units (e.g. when accessing \texttt{G.var_}).

### Attributes

- **\texttt{dtype}**
- **\texttt{shape}**
- **\texttt{unit}** The \texttt{Unit} of this variable

### Methods

- **\texttt{get\_item(item[, level, namespace])}** Get the value of this variable.
- **\texttt{get\_subexpression\_with\_index\_array(**kwds)}**
- **\texttt{get\_with\_expression(**kwds)}** Gets a variable using a string expression.
- **\texttt{get\_with\_index\_array(**kwds)}**
- **\texttt{set\_item(item, value[, level, namespace])}** Set this variable.
- **\texttt{set\_subexpression\_with\_index\_array(**kwds)}**
- **\texttt{set\_with\_expression(**kwds)}** Sets a variable using a string expression.
- **\texttt{set\_with\_expression\_conditional(**kwds)}** Sets a variable using a string expression and string condition.
- **\texttt{set\_with\_index\_array(**kwds)}**

### Details

- **\texttt{dtype}**
- **\texttt{shape}**
- **\texttt{unit}** The \texttt{Unit} of this variable

**\texttt{get\_item(item, level=0, namespace=None)}** Get the value of this variable. Called by \texttt{__getitem__}.

**Parameters**

- **item** : slice, \texttt{ndarray} or string
  - The index for the setting operation
- **level** : int, optional
  - How much farther to go up in the stack to find the implicit namespace (if used, see \texttt{run\_namespace}).
- **namespace** : dict-like, optional
  - An additional namespace that is used for variable lookup (if not defined, the implicit namespace of local variables is used).

**\texttt{get\_subexpression\_with\_index\_array(**kwds)}**

**\texttt{get\_with\_expression(**kwds)}** Gets a variable using a string expression. Is called by \texttt{VariableView.get\_item} for statements such as \texttt{print(G.v['g_syn > 0'])}.

**Parameters**

- **code** : str
An expression that states a condition for elements that should be selected. Can contain references to indices, such as \(i\) or \(j\) and to state variables. For example: '\(i>3\) and \(v>0\text{ mV}\).

**run_namespace**: dict-like

An additional namespace that is used for variable lookup (either an explicitly defined namespace or one taken from the local context).

**get_with_index_array** (**kwds**)

**set_item** (*item, value, level=0, namespace=None*)

Set this variable. This function is called by \_\_setitem\_ but there is also a situation where it should be called directly: if the context for string-based expressions is higher up in the stack, this function allows to set the level argument accordingly.

**Parameters**

- **item**: slice, ndarray or string
  - The index for the setting operation
- **value**: Quantity, ndarray or number
  - The value for the setting operation
- **level**: int, optional
  - How much farther to go up in the stack to find the implicit namespace (if used, see run_namespace).
- **namespace**: dict-like, optional
  - An additional namespace that is used for variable lookup (if not defined, the implicit namespace of local variables is used).

**set_with_expression** (**kwds**)

Sets a variable using a string expression. Is called by VariableView.set_item for statements such as \(S\cdot\text{var}[:, :, :] = \exp(-\text{abs}(i-j)/\text{space_constant})\cdot\text{nS}\).

**Parameters**

- **item**: ndarray
  - The indices for the variable (in the context of this group).
- **code**: str
  - The code that should be executed to set the variable values. Can contain references to indices, such as \(i\) or \(j\)
- **run_namespace**: dict-like, optional
  - An additional namespace that is used for variable lookup (if not defined, the implicit namespace of local variables is used).
- **check_units**: bool, optional
  - Whether to check the units of the expression.
- **run_namespace**: dict-like, optional
  - An additional namespace that is used for variable lookup (if not defined, the implicit namespace of local variables is used).

**set_with_expression_conditional** (**kwds**)

Sets a variable using a string expression and string condition. Is called by VariableView.set_item for statements such as \(S\cdot\text{var}'i!=j' = \exp(-\text{abs}(i-j)/\text{space_constant})\cdot\text{nS}'

**Parameters**

- **cond**: str
The string condition for which the variables should be set.

code : str

The code that should be executed to set the variable values.

run_namespace : dict-like, optional

An additional namespace that is used for variable lookup (if not defined, the implicit
namespace of local variables is used).

check_units : bool, optional

Whether to check the units of the expression.

set_with_index_array(**kwds)

Variables(owner[, default_index]) A container class for storing Variable objects.

Variables class

(Shortest import: from brian2.core.variables import Variables)

class brian2.core.variables.Variables(owner, default_index='_idx')

Bases: collections.abc.Mapping

A container class for storing Variable objects. Instances of this class are used as the Group.variables
attribute and can be accessed as (read-only) dictionaries.

Parameters

owner : Nameable

The object (typically a Group) “owning” the variables.

default_index : str, optional

The index to use for the variables (only relevant for ArrayVariable and
DynamicArrayVariable). Defaults to '_idx'.

Attributes

indices A dictionary given the index name for every array
name

owner A reference to the Group owning these variables

Methods

add_arange(name, size[, start, dtype, ...]) Add an array, initialized with a range of integers.

add_array(name, size[, dimensions, values, ...]) Add an array (initialized with zeros).

add_arrays(names, size[, dimensions, dtype, ...
...]) Adds several arrays (initialized with zeros) with the
same attributes (size, units, etc.).

add_auxiliary_variable(name[, dimensions, ...]) Add an auxiliary variable (most likely one that is
added automatically to abstract code, e.g.

add_constant(name, value[, dimensions]) Add a scalar constant (e.g.

add_dynamic_array(name, size[, dimensions, ...]) Add a dynamic array.

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### Table 212 – continued from previous page

<table>
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<th>Function</th>
<th>Description</th>
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<td><code>add_object(name, obj)</code></td>
<td>Add an arbitrary Python object.</td>
</tr>
<tr>
<td><code>add_reference(name, group[, varname, index])</code></td>
<td>Add a reference to a variable defined somewhere else (possibly under a different name).</td>
</tr>
<tr>
<td><code>add_references(group, varnames[, index])</code></td>
<td>Add all <code>Variable</code> objects from a name to <code>Variable</code> mapping with the same name as in the original mapping.</td>
</tr>
<tr>
<td><code>add_referred_subexpression(name, group, ...)</code></td>
<td></td>
</tr>
<tr>
<td><code>add_subexpression(name, expr[, dimensions, ...])</code></td>
<td>Add a named subexpression.</td>
</tr>
<tr>
<td><code>create_clock_variables(clock[, prefix])</code></td>
<td>Convenience function to add the ( t ) and ( dt ) attributes of a clock.</td>
</tr>
</tbody>
</table>

#### Details

**indices**

A dictionary given the index name for every array name

**owner**

A reference to the `Group` owning these variables

**add_range**

```python
add_range(name, size, start=0, dtype=<class 'numpy.int32'>, constant=True, read_only=True, unique=True, index=None)
```

Add an array, initialized with a range of integers.

**Parameters**

- **name** : str
  The name of the variable.

- **size** : int
  The size of the array.

- **start** : int
  The start value of the range.

- **dtype** : dtype, optional
  The dtype used for storing the variable. If none is given, defaults to `np.int32`.

- **constant** : bool, optional
  Whether the variable’s value is constant during a run. Defaults to `True`.

- **read_only** : bool, optional
  Whether this is a read-only variable, i.e. a variable that is set internally and cannot be changed by the user. Defaults to `True`.

- **index** : str, optional
  The index to use for this variable. Defaults to `Variables.default_index`.

- **unique** : bool, optional
  See `ArrayVariable`. Defaults to `True` here.

**add_array**

```python
add_array(name, size, dimensions=Dimension(), values=None, dtype=None, constant=False, read_only=False, scalar=False, unique=False, index=None)
```

Add an array (initialized with zeros).

**Parameters**

- **name** : str
  The name of the variable.
The name of the variable.

dimensions: Dimension, optional

The physical dimensions of the variable.

size: int

The size of the array.

values: ndarray, optional

The values to initialize the array with. If not specified, the array is initialized to zero.

dtype: dtype, optional

The dtype used for storing the variable. If none is given, defaults to \texttt{core.default_float_dtype}.

constant: bool, optional

Whether the variable’s value is constant during a run. Defaults to False.

scalar: bool, optional

Whether this is a scalar variable. Defaults to False, if set to True, also implies that size() equals 1.

read_only: bool, optional

Whether this is a read-only variable, i.e. a variable that is set internally and cannot be changed by the user. Defaults to False.

index: str, optional

The index to use for this variable. Defaults to \texttt{Variables.default_index}.

unique: bool, optional

See \texttt{ArrayVariable}. Defaults to False.

\texttt{add_arrays (names, size, dimensions=Dimension(), dtype=None, constant=False, read_only=False, scalar=False, unique=False, index=None)}

Adds several arrays (initialized with zeros) with the same attributes (size, units, etc.).

**Parameters**

names: list of str

The names of the variable.

dimensions: Dimension, optional

The physical dimensions of the variable.

size: int

The sizes of the arrays.

dtype: dtype, optional

The dtype used for storing the variables. If none is given, defaults to \texttt{core.default_float_dtype}.

constant: bool, optional

Whether the variables’ values are constant during a run. Defaults to False.

scalar: bool, optional

Whether these are scalar variables. Defaults to False, if set to True, also implies that size() equals 1.
read_only : bool, optional
Whether these are read-only variables, i.e. variables that are set internally and cannot
be changed by the user. Defaults to False.

index : str, optional
The index to use for these variables. Defaults to Variables.default_index.

unique : bool, optional
See ArrayVariable. Defaults to False.

add_auxiliary_variable (name, dimensions=Dimension(), dtype=None, scalar=False)
Add an auxiliary variable (most likely one that is added automatically to abstract code, e.g. _cond for a
threshold condition), specifying its type and unit for code generation.

Parameters name : str
The name of the variable

dimensions : Dimension
The physical dimensions of the variable.

dtype : dtype, optional
The dtype used for storing the variable. If none is given, defaults to
core.default_float_dtype.

scalar : bool, optional
Whether the variable is a scalar value (True) or vector-valued, e.g. defined for every
neuron (False). Defaults to False.

add_constant (name, value, dimensions=Dimension())
Add a scalar constant (e.g. the number of neurons N).

Parameters name : str
The name of the variable

value: reference to the variable value :
The value of the constant.

dimensions : Dimension, optional
The physical dimensions of the variable. Note that the variable itself (as referenced by
value) should never have units attached.

add_dynamic_array (name, size, dimensions=Dimension(), values=None, dtype=None, constant=False, needs_reference_update=False, resize_along_first=False, read_only=False, unique=False, scalar=False, index=None)
Add a dynamic array.

Parameters name : str
The name of the variable.

dimensions : Dimension, optional
The physical dimensions of the variable.

size : int or tuple of int
The (initial) size of the array.

values : ndarray, optional
The values to initialize the array with. If not specified, the array is initialized to zero.

**dtype**: `dtype`, optional

The dtype used for storing the variable. If none is given, defaults to `core.default_float_dtype`.

**constant**: bool, optional

Whether the variable’s value is constant during a run. Defaults to `False`.

**needs_reference_update**: bool, optional

Whether the code objects need a new reference to the underlying data at every time step. This should be set if the size of the array can be changed by other code objects. Defaults to `False`.

**scalar**: bool, optional

Whether this is a scalar variable. Defaults to `False`, if set to `True`, also implies that `size()` equals 1.

**read_only**: bool, optional

Whether this is a read-only variable, i.e. a variable that is set internally and cannot be changed by the user. Defaults to `False`.

**index**: str, optional

The index to use for this variable. Defaults to `Variables.default_index`.

**unique**: bool, optional

See `DynamicArrayVariable`. Defaults to `False`.

**add_object**(name, obj)

Add an arbitrary Python object. This is only meant for internal use and therefore only names starting with an underscore are allowed.

**Parameters**

- **name**: str

  The name used for this object (has to start with an underscore).

- **obj**: object

  An arbitrary Python object that needs to be accessed directly from a `CodeObject`.

**add_reference**(name, group, varname=None, index=None)

Add a reference to a variable defined somewhere else (possibly under a different name). This is for example used in `Subgroup` and `Synapses` to refer to variables in the respective `NeuronGroup`.

**Parameters**

- **name**: str

  The name of the variable (in this group, possibly a different name from `var.name`).

- **group**: `Group`

  The group from which `var()` is referenced.

- **varname**: str, optional

  The variable to refer to. If not given, defaults to `name`.

- **index**: str, optional

  The index that should be used for this variable (defaults to `Variables.default_index`).
add_references \( (\text{group, varnames, index=None}) \)
Add all \texttt{Variable} objects from a name to \texttt{Variable} mapping with the same name as in the original mapping.

**Parameters**
- **group**: \texttt{Group}
The group from which the variables are referenced
- **varnames**: iterable of str
  The variables that should be referred to in the current group
- **index**: str, optional
  The index to use for all the variables (defaults to \texttt{Variables.default_index})

add_referred_subexpression \( (\text{name, group, subexpr, index}) \)

add_subexpression \( (\text{name, expr, dimensions=Dimension(), dtype=None, scalar=False, index=None}) \)
Add a named subexpression.

**Parameters**
- **name**: str
  The name of the subexpression.
- **dimensions**: \texttt{Dimension}
  The physical dimensions of the subexpression.
- **expr**: str
  The subexpression itself.
- **dtype**: \texttt{dtype}, optional
  The dttype used for the expression. Defaults to \texttt{core.default_float_dtype}.
- **scalar**: bool, optional
  Whether this is an expression only referring to scalar variables. Defaults to \texttt{False}
- **index**: str, optional
  The index to use for this variable. Defaults to \texttt{Variables.default_index}.

create_clock_variables \( (\text{clock, prefix=""}) \)
Convenience function to add the \texttt{t} and \texttt{dt} attributes of a clock.

**Parameters**
- **clock**: \texttt{Clock}
The clock that should be used for \texttt{t} and \texttt{dt}.
- **prefix**: str, optional
  A prefix for the variable names. Used for example in monitors to not confuse the dynamic array of recorded times with the current time in the recorded group.

Functions

\begin{center}
\begin{tabular}{ll}
\texttt{get_dtype(obj)} & Helper function to return the \texttt{numpy.dtype} of an arbitrary object.
\end{tabular}
\end{center}
get_dtype function

(Shortest import: from brian2.core.variables import get_dtype)

brian2.core.variables.get_dtype(obj)
   Helper function to return the numpy.dtype of an arbitrary object.
   Parameters  obj: object
               Any object (but typically some kind of number or array).
   Returns  dtype: numpy.dtype
               The type of the given object.

get_dtype_str(val)  Returns canonical string representation of the dtype of a value or dtype

get_dtype_str function

(Shortest import: from brian2.core.variables import get_dtype_str)

brian2.core.variables.get_dtype_str(val)
   Returns canonical string representation of the dtype of a value or dtype
   Returns  dtype_str: str
               The numpy dtype name

linked_var(group_or_variable[, name, index])  Represents a link target for setting a linked variable.

linked_var function

(Shortest import: from brian2 import linked_var)

brian2.core.variables.linked_var(group_or_variable, name=None, index=None)
   Represents a link target for setting a linked variable.
   Parameters  group_or_variable: NeuronGroup or VariableView
               Either a reference to the target NeuronGroup (e.g. G) or a direct reference to a
               VariableView object (e.g. G.v). In case only the group is specified, name has
               to be specified as well.
   name: str, optional
         The name of the target variable, necessary if group_or_variable is a
         NeuronGroup.
   index: str or ndarray, optional
          An indexing array (or the name of a state variable), providing a mapping from the entries
          in the link source to the link target.
Examples

```python
>>> from brian2 import *

>>> G1 = NeuronGroup(10, 'dv/dt = -v / (10*ms) : volt')
>>> G2 = NeuronGroup(10, 'v : volt (linked)')
>>> G2.v = linked_var(G1, 'v')
>>> G2.v = linked_var(G1.v)  # equivalent
```

`variables_by_owner(variables, owner)`

`variables_by_owner` function

`(Shortest import: from brian2.core.variables import variables_by_owner)`

brian2.core.variables.**variables_by_owner**(variables, owner)

### 6.6.3 devices package

Package providing the “devices” infrastructure.

**device module**

Module containing the `Device` base class as well as the `RuntimeDevice` implementation and some helper functions to access/set devices.

Exported members: `Device`, `RuntimeDevice`, `get_device()`, `set_device()`, `all_devices`, `reinit_devices`, `reinit_and_delete`, `reset_device`, `device`, `seed()`

Classes

- `CurrentDeviceProxy` Method proxy for access to the currently active device
  
  `(Shortest import: from brian2.devices.device import CurrentDeviceProxy)`

  **class** brian2.devices.device.**CurrentDeviceProxy**

  **Bases:** object

  Method proxy for access to the currently active device

- `Device()` Base Device object.
  
  `(Shortest import: from brian2.devices.device import Device)`

  **class** brian2.devices.device.**Device**

  **Bases:** object

  Base Device object.
Attributes

**network_schedule**
The network schedule that this device supports.

Methods

**activate**([build_on_run=True,**kwargs])**
Called when this device is set as the current device.

**add_array**(var)
Add an array to this device.

**build**(**kwargs**)
For standalone projects, called when the project is ready to be built.

**code_object**(owner, name, abstract_code,**...)**
Return CodeObject class according to input/default settings.

**code_object_class**([codeobj_class,**fallback_pref**])
Return CodeObject class according to input/default settings.

**delete**(data, code, directory,**force**)**
Delete code and/or data generated/stored by the device.

**fill_with_array**(var, arr)
Fill an array with the values given in another array.

**get_array_name**(var,**access_data**)**
Return a globally unique name for var().

**get_len**(array)
Return the length of the array.

**get_random_state**( )
Return a (pickable) representation of the current random number generator state.

**init_with_arange**(var, start, dtype)
Initialize an array with an integer range.

**init_with_zeros**(var, dtype)
Initialize an array with zeros.

**insert_code**(slot, code)
Insert code directly into a given slot in the device.

**insert_device_code**(slot, code)
Insert code directly into a given slot in the device.

**reinit**( )
Reinitialize the device.

**resize**(var, new_size)
Resize a DynamicArrayVariable.

**resize_along_first**(var, new_size)
Resize a DynamicArrayVariable.

**seed**(seed)
Set the seed for the random number generator.

**set_random_state**(state)
Reset the random number generator state to a previously stored state (see Device.get_random_state).

**spike_queue**(source_start, source_end)
Create and return a new SpikeQueue for this Device.

Details

**network_schedule**
The network schedule that this device supports. If the device only supports a specific, fixed schedule, it has to set this attribute to the respective schedule (see Network.schedule for details). If it supports arbitrary schedules, it should be set to None (the default).

**activate**(build_on_run=True,**kwargs**)
Called when this device is set as the current device.

**add_array**(var)
Add an array to this device.

Parameters

**var**: ArrayVariable
The array to add.

**build**(**kwargs**)
For standalone projects, called when the project is ready to be built. Does nothing for runtime mode.
code_object (owner, name, abstract_code, variables, template_name, variable_indices, codeobj_class=None, template_kwds=None, override_conditional_write=None, compiler_kwds=None)
code_object_class (codeobj_class=None, fallback_pref='codegen.target')
   Return CodeObject class according to input/default settings

   Parameters codeobj_class : a CodeObject class, optional
      If this is keyword is set to None or no arguments are given, this method will return the
default.

codeobj_class : str, optional
   String describing which attribute of prefs to access to retrieve the ‘default’ target. Usually
   this is codegen.target, but in some cases we want to use object-specific targets such
   as codegen.string_expression_target.

   Returns codeobj_class : class
      The CodeObject class that should be used

delete (data=True, code=True, directory=True, force=False)
   Delete code and/or data generated/stored by the device.

   Parameters data : bool, optional
      Whether to delete the data generated by the simulation (final values of state variables,
data stored in monitors, etc.). Defaults to True.

code : bool, optional
   Whether to delete the code generated by the simulation. Includes the numerical values
   used for initialization of state variables in assignments not using strings. Defaults to
   True.

directory : bool, optional
   Whether to delete the project directory generated by the simulation. Will not delete
directories that contain files not created by Brian unless the force option is specified.
   Defaults to True.

   force : bool, optional
      Whether to delete the project directory with all its content, even if it contains files that
      were not created by Brian. Useful only when the directory option is set to True as
      well. Defaults to False.

fill_with_array (var, arr)
   Fill an array with the values given in another array.

   Parameters var : ArrayVariable
      The array to fill.

   arr : ndarray
      The array values that should be copied to var().

get_array_name (var, access_data=True)
   Return a globally unique name for var().

   Parameters access_data : bool, optional
For `DynamicArrayVariable` objects, specifying `True` here means the name for the underlying data is returned. If specifying `False`, the name of object itself is returned (e.g. to allow resizing).

**Returns name**: str

The name for `var()`.

**get_len**(array)

Return the length of the array.

**Parameters**

- **array**: `ArrayVariable`

  The array for which the length is requested.

**Returns l**: int

The length of the array.

**get_random_state()**

Return a (pickable) representation of the current random number generator state. Providing the returned object (e.g. a dict) to `Device.set_random_state` should restore the random number generator state.

**Returns state**:

The state of the random number generator in a representation that can be passed as an argument to `Device.set_random_state`.

**init_with_arange**(var, start, dtype)

Initialize an array with an integer range.

**Parameters**

- **var**: `ArrayVariable`

  The array to fill with the integer range.

- **start**: int

  The start value for the integer range

- **dtype**: `dtype`

  The data type to use for the array.

**init_with_zeros**(var, dtype)

Initialize an array with zeros.

**Parameters**

- **var**: `ArrayVariable`

  The array to initialize with zeros.

- **dtype**: `dtype`

  The data type to use for the array.

**insert_code**(slot, code)

Insert code directly into a given slot in the device. By default does nothing.

**insert_device_code**(slot, code)

**reinit**()

Reinitialize the device. For standalone devices, clears all the internal state of the device.

**resize**(var, new_size)

Resize a `DynamicArrayVariable`.

**Parameters**

- **var**: `DynamicArrayVariable`

  The variable that should be resized.
new_size : int
    The new size of the variable

resize_along_first (var, new_size)

seed (seed=None)
    Set the seed for the random number generator.

    Parameters seed : int, optional
        The seed value for the random number generator, or None (the default) to set a random seed.

set_random_state (state)
    Reset the random number generator state to a previously stored state (see Device.get_random_state).

    Parameters state :
        A random number generator state as provided by Device.get_random_state.

spike_queue (source_start, source_end)
    Create and return a new SpikeQueue for this Device.

    Parameters source_start : int
        The start index of the source group (necessary for subgroups)

    source_end : int
        The end index of the source group (necessary for subgroups)

Dummy

Dummy class

(Shortest import: from brian2.devices.device import Dummy)

class brian2.devices.device.Dummy
    Bases: object
    Dummy object

    Methods

___call___(*args, **kwds) Call self as a function.

Details

___call___(*args, **kwds)
    Call self as a function.

Tutorials and examples using this

- Example advanced/exprel_function
- Example frompapers/Stimberg_et_al_2018/example_2_gchiAstrocyte
The default device used in Brian, state variables are stored as numpy arrays in memory.

### RuntimeDevice class

*(Shortest import: from brian2.devices import RuntimeDevice)*

```python
class brian2.devices.device.RuntimeDevice
    Bases: brian2.devices.device.Device
```

The default device used in Brian, state variables are stored as numpy arrays in memory.

#### Attributes

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<td>Mapping from Variable objects to numpy arrays (or DynamicArray objects).</td>
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#### Methods

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<th>Method</th>
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<td><code>add_array(var)</code></td>
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<td><code>get_array_name(var[, access_data])</code></td>
<td>Return a globally unique name for var().</td>
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<td><code>get_random_state()</code></td>
<td>Return a (pickable) representation of the current random number generator state.</td>
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<td><code>get_value(var[, access_data])</code></td>
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<td><code>resize(var, new_size)</code></td>
<td>Resize a DynamicArrayVariable.</td>
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<td><code>resize_along_first(var, new_size)</code></td>
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<td><code>set_random_state(state)</code></td>
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<td><code>set_value(var, value)</code></td>
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<tr>
<td><code>spike_queue(source_start, source_end)</code></td>
<td>Create and return a new SpikeQueue for this Device.</td>
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#### Details

**arrays**

Mapping from Variable objects to numpy arrays (or DynamicArray objects). Arrays in this dictionary will disappear as soon as the last reference to the Variable object used as a key is gone.

**add_array** *(var)*

Add an array to this device.

**Parameters** *var*: ArrayVariable

The array to add.

**fill_with_array** *(var, arr)*
Fill an array with the values given in another array.

**Parameters**

- **var : ArrayVariable**
  - The array to fill.

- **arr : ndarray**
  - The array values that should be copied to `var()`.

**get_array_name** *(var, access_data=True)*

Return a globally unique name for `var()`.

**Parameters**

- **access_data : bool, optional**
  - For `DynamicArrayVariable` objects, specifying `True` here means the name for the underlying data is returned. If specifying `False`, the name of object itself is returned (e.g. to allow resizing).

**Returns**

- **name : str**
  - The name for `var()`.

**get_random_state** *

Return a (pickable) representation of the current random number generator state. Providing the returned object (e.g. a dict) to `Device.set_random_state` should restore the random number generator state.

**Returns**

- **state**
  - The state of the random number generator in a representation that can be passed as an argument to `Device.set_random_state`.

**get_value** *(var, access_data=True)*

**init_with_arange** *(var, start, dtype)*

Initialize an array with an integer range.

**Parameters**

- **var : ArrayVariable**
  - The array to fill with the integer range.

- **start : int**
  - The start value for the integer range

- **dtype : dtype**
  - The data type to use for the array.

**init_with_zeros** *(var, dtype)*

Initialize an array with zeros.

**Parameters**

- **var : ArrayVariable**
  - The array to initialize with zeros.

- **dtype : dtype**
  - The data type to use for the array.

**resize** *(var, new_size)*

Resize a `DynamicArrayVariable`.

**Parameters**

- **var : DynamicArrayVariable**
  - The variable that should be resized.

- **new_size : int**
The new size of the variable

\texttt{resize\_along\_first}(\textit{var, new\_size})

\texttt{seed}(\textit{seed=None})

Set the seed for the random number generator.

**Parameters** \texttt{seed} : int, optional

The seed value for the random number generator, or \texttt{None} (the default) to set a random seed.

\texttt{set\_random\_state}(\textit{state})

Reset the random number generator state to a previously stored state (see \texttt{Device.get\_random\_state}).

**Parameters** \texttt{state} :

A random number generator state as provided by \texttt{Device.get\_random\_state}.

\texttt{set\_value}(\textit{var, value})

\texttt{spike\_queue}(\textit{source\_start, source\_end})

Create and return a new \texttt{SpikeQueue} for this \texttt{Device}.

**Parameters** \texttt{source\_start} : int

The start index of the source group (necessary for subgroups)

\texttt{source\_end} : int

The end index of the source group (necessary for subgroups)

**Tutorials and examples using this**

- Example \texttt{from\_papers/Kremer\_et\_al\_2011\_barrel\_cortex}

**Functions**

\texttt{auto\_target()}

Automatically chose a code generation target (invoked when the \texttt{codegen.target} preference is set to 'auto').

**auto\_target function**

*(Shortest import: from brian2.devices.device import auto\_target)*

\texttt{brian2.devices.device.auto\_target()}

Automatically chose a code generation target (invoked when the \texttt{codegen.target} preference is set to 'auto'). Caches its result so it only does the check once. Prefers weave > cython > numpy.

**Returns** \texttt{target} : class derived from \texttt{CodeObject}

The target to use

\texttt{get\_device()}

Gets the active \texttt{Device} object
**get_device function**

*(Shortest import: from brian2 import get_device)*

```python
brian2.devices.device.get_device()
```

Gets the active `Device` object.

**reinit_and_delete function**

*(Shortest import: from brian2.devices import reinit_and_delete)*

```python
brian2.devices.device.reinit_and_delete()
```

Calls `reinit_devices` and additionally deletes the files left behind by the standalone mode in the temporary directory. Silently suppresses errors that occur while deleting the directory.

**reinit_devices function**

*(Shortest import: from brian2.devices import reinit_devices)*

```python
brian2.devices.device.reinit_devices()
```

Reinitialize all devices, call `Device.activate` again on the current device and reset the preferences. Used as a “teardown” function in testing, if users want to reset their device (e.g. for multiple standalone runs in a single script), calling `device.reinit()` followed by `device.activate()` should normally be sufficient.

**Notes**

This also resets the `defaultclock`, i.e. a non-standard `dt` has to be set again.

```python
reset_device([device])
```

Reset to a previously used device.

**reset_device function**

*(Shortest import: from brian2.devices import reset_device)*

```python
brian2.devices.device.reset_device(device=None)
```

Reset to a previously used device. Restores also the previously specified build options (see `set_device()`) for the device. Mostly useful for internal Brian code and testing on various devices.

**Parameters**

- `device`: `Device` or str, optional

  The device to go back to. If none is specified, go back to the device chosen with `set_device()` before the current one.
**seed**([seed])  
Set the seed for the random number generator.

**seed function**

*(Shortest import: from brian2 import seed)*

brian2.devices.device.seed(seed=None)

Set the seed for the random number generator.

**Parameters**  
*seed* : int, optional

The seed value for the random number generator, or *None* (the default) to set a random seed.

**Notes**

This function delegates the call to *Device.seed* of the current device.

**set_device**(device[, build_on_run])  
Set the device used for simulations.

**set_device function**

*(Shortest import: from brian2 import set_device)*

brian2.devices.device.set_device(device, build_on_run=True, **kwargs)

Set the device used for simulations.

**Parameters**  
*device* : *Device* or str

The *Device* object or the name of the device.

*build_on_run* : bool, optional

Whether a call to *run()* (or *Network.run*) should directly trigger a *Device.build*. This is only relevant for standalone devices and means that a run call directly triggers the start of a simulation. If the simulation consists of multiple run calls, set *build_on_run* to False and call *Device.build* explicitly. Defaults to True.

*kwargs* : dict, optional

Only relevant when *build_on_run* is True: additional arguments that will be given to the *Device.build* call.

**Objects**

*active_device*  
The currently active device (set with *set_device()*)

**active_device object**

*(Shortest import: from brian2.devices.device import active_device)*

brian2.devices.device.active_device = <brian2.devices.device.RuntimeDevice object>

The currently active device (set with *set_device()*).
**device object**

(Shortest import: from brian2 import device)

```python
def device: device = <brian2.devices.device.CurrentDeviceProxy object>
Proxy object to access methods of the current device
```

**runtime_device object**

(Shortest import: from brian2.devices.device import runtime_device)

```python
def runtime_device: runtime_device = <brian2.devices.device.RuntimeDevice object>
The default device used in Brian, state variables are stored as numpy arrays in memory.
```

**Subpackages**

**cpp_standalone package**

Package implementing the C++ “standalone” Device and CodeObject.

**GSLcodeobject module**

Module containing CPPStandalone CodeObject for code generation for integration using the ODE solver provided in the GNU Scientific Library

**Classes**

```python
def GSLCPPStandaloneCodeObject(owner, code, ...
```

**GSLCPPStandaloneCodeObject class**

(Shortest import: from brian2.devices.cpp_standalone import GSLCPPStandaloneCodeObject)
class brian2.devices.cpp_standalone.GSLcodeobject.GSLCPPStandaloneCodeObject (owner, code, variables, variable_indices, template_name, template_source, compiler_kwds, name='codeobject*'))

Bases: brian2.codegen.codeobject.CodeObject

codeobject module

Module implementing the C++ “standalone” CodeObject

Exported members: CPPStandaloneCodeObject

Classes

CPPStandaloneCodeObject( )

CPPStandaloneCodeObject class

(Shortest import: from brian2.devices.cpp_standalone import CPPStandaloneCodeObject)

class brian2.devices.cpp_standalone.codeobject.CPPStandaloneCodeObject (owner, code, variables, variable_indices, template_name, template_source, compiler_kwds, name='codeobject*'))

Bases: brian2.codegen.codeobject.CodeObject

C++ standalone code object

The code should be a MultiTemplate object with two macros defined, main (for the main loop code) and support_code for any support code (e.g. function definitions).

Methods
__call__(**kwds)  Call self as a function.

run()  Runs the code in the namespace.

Details

__call__(**kwds)
Call self as a function.

run()
Runs the code in the namespace.

Returns return_value : dict
A dictionary with the keys corresponding to the output_variables defined during the call of CodeGenerator.code_object.

Functions

generate_rand_code(rand_func, owner)

generate_rand_code function

(Shortest import: from brian2.devices.cpp_standalone.codeobject import generate_rand_code)
brian2.devices.cpp_standalone.codeobject.generate_rand_code(rand_func, owner)

openmp_pragma(pragma_type)

openmp_pragma function

(Shortest import: from brian2.devices.cpp_standalone.codeobject import openmp_pragma)
brian2.devices.cpp_standalone.codeobject.openmp_pragma(pragma_type)

device module

Classes

CPPStandaloneDevice()  The Device used for C++ standalone simulations.

CPPStandaloneDevice class

(Shortest import: from brian2.devices.cpp_standalone.device import CPPStandaloneDevice)
class brian2.devices.cpp_standalone.device.CPPStandaloneDevice
    Bases: brian2.devices.device.Device

    The Device used for C++ standalone simulations.
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<td>List of all arrays to be filled with numbers (list of (var, varname, start) tuples)</td>
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<tr>
<td><code>array_cache</code></td>
<td>Dictionary mapping <code>ArrayVariable</code> objects to their value or to <code>None</code> if the value (potentially) depends on executed code.</td>
</tr>
<tr>
<td><code>arrays</code></td>
<td>Dictionary mapping <code>ArrayVariable</code> objects to their globally unique name</td>
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<tr>
<td><code>build_on_run</code></td>
<td>Whether a run should trigger a build</td>
</tr>
<tr>
<td><code>build_options</code></td>
<td>Build options</td>
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<td>Code lines that have been manually added with <code>device.insert_code()</code> Dictionary mapping slot names to lists of lines.</td>
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<td>List of all dynamic arrays Dictionary mapping <code>DynamicArrayVariable</code> objects with 1 dimension to their globally unique name</td>
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<td>Dictionary mapping <code>DynamicArrayVariable</code> objects with 2 dimensions to their globally unique name</td>
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<td>Whether to generate profiling information (stored in an instance variable to be accessible during CodeObject generation)</td>
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<td><code>has Been run</code></td>
<td>Whether the simulation has been run</td>
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<td>Cached environment variables for MSVC</td>
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<tr>
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<td><code>project_dir</code></td>
<td>The directory which contains the generated code and results</td>
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<td>Dict of all static saved arrays</td>
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<td>List of all arrays to be filled with zeros (list of (var, varname) )</td>
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### Methods

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<td><code>add_array(var)</code></td>
<td>Add an array to this device.</td>
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<td><code>build([directory, compile, run, debug, ...])</code></td>
<td>Build the project</td>
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<tr>
<td><code>code_object(owner, name, abstract_code, ...)</code></td>
<td>Return CodeObject class (either CPPStandaloneCodeObject class or input)</td>
</tr>
<tr>
<td><code>code_object_class([codeobj_class, fallback_pref])</code></td>
<td></td>
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<td><code>compile_source(directory, compiler, debug, clean)</code></td>
<td></td>
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<td><code>copy_source_files(writer, directory)</code></td>
<td>Delete code and/or data generated/stored by the device.</td>
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<td><code>fill_with_array(var, arr)</code></td>
<td>Fill an array with the values given in another array.</td>
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</tr>
<tr>
<td><code>resize</code></td>
<td><code>var, new_size</code> Resize a DynamicArrayVariable.</td>
</tr>
<tr>
<td><code>run</code></td>
<td><code>directory, with_output, run_args</code></td>
</tr>
<tr>
<td><code>run_function</code></td>
<td><code>name[, include_in_parent]</code> Context manager to divert code into a function</td>
</tr>
<tr>
<td><code>seed</code></td>
<td><code>[seed]</code> Set the seed for the random number generator.</td>
</tr>
<tr>
<td><code>static_array</code></td>
<td><code>name, arr</code></td>
</tr>
<tr>
<td><code>variableview_get_subexpression_with_index_array</code></td>
<td></td>
</tr>
<tr>
<td><code>variableview_get_with_expression</code></td>
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</tr>
<tr>
<td><code>variableview_set_with_index_array</code></td>
<td></td>
</tr>
<tr>
<td><code>write_static_arrays</code></td>
<td><code>directory</code></td>
</tr>
</tbody>
</table>

### Details

**arange_arrays**

List of all arrays to be filled with numbers (list of (var, varname, start) tuples)

**array_cache**

Dictionary mapping `ArrayVariable` objects to their value or to `None` if the value (potentially) depends on executed code. This mechanism allows to access state variables in standalone mode if their value is known at run time.

**arrays**

Dictionary mapping `ArrayVariable` objects to their globally unique name.

**build_on_run**

Whether a run should trigger a build.

**build_options**

Build options.

**code_lines**

Code lines that have been manually added with `device.insert_code()` Dictionary mapping slot names to lists of lines. Note that the main slot is handled separately as part of `main_queue`.

**dynamic_arrays**

List of all dynamic arrays Dictionary mapping DynamicArrayVariable objects with 1 dimension to their globally unique name.
**dynamic_arrays_2d**
Dictionary mapping DynamicArrayVariable objects with 2 dimensions to their globally unique name

**enable_profiling**
Whether to generate profiling information (stored in an instance variable to be accessible during CodeObject generation)

**has_been_run**
Whether the simulation has been run

**msvc_env**
cached environment variables for MSVC

**profiled_codeobjects**
CodeObjects that use profiling (users can potentially enable profiling only for a subset of runs)

**project_dir**
The directory which contains the generated code and results

**static_arrays**
Dict of all static saved arrays

**zero_arrays**
List of all arrays to be filled with zeros (list of (var, varname) )

**add_array** *(var)*
Add an array to this device.

**Parameters**
*var*: ArrayVariable
The array to add.

**build** *(directory='output', compile=True, run=True, debug=False, clean=False, with_output=True, additional_source_files=None, run_args=None, direct_call=True, **kwds)*
Build the project

TODO: more details

**Parameters**
*directory*: str, optional
The output directory to write the project to, any existing files will be overwritten. If the given directory name is None, then a temporary directory will be used (used in the test suite to avoid problems when running several tests in parallel). Defaults to 'output'.

*compile*: bool, optional
Whether or not to attempt to compile the project. Defaults to True.

*run*: bool, optional
Whether or not to attempt to run the built project if it successfully builds. Defaults to True.

*debug*: bool, optional
Whether to compile in debug mode. Defaults to False.

*with_output*: bool, optional
Whether or not to show the stdout of the built program when run. Output will be shown in case of compilation or runtime error. Defaults to True.

*clean*: bool, optional
Whether or not to clean the project before building. Defaults to False.
additional_source_files : list of str, optional
A list of additional .cpp files to include in the build.
direct_call : bool, optional
Whether this function was called directly. Is used internally to distinguish an automatic build due to the build_on_run option from a manual device.build call.

check_openmp_compatible(nb_threads)
code_object(owner, name, abstract_code, variables, template_name, variable_indices, codeobj_class=None, template_kwds=None, override_conditional_write=None, compiler_kwds=None)
code_object_class(codeobj_class=None, fallback_pref=None)
Return CodeObject class (either CPPStandaloneCodeObject class or input)

Parameters
codeobj_class : a CodeObject class, optional
If this is keyword is set to None or no arguments are given, this method will return the default (CPPStandaloneCodeObject class).
fallback_pref : str, optional
For the cpp_standalone device this option is ignored.

Returns
codeobj_class : class
The CodeObject class that should be used

compile_source(directory, compiler, debug, clean)
copy_source_files(writer, directory)
delete(code=True, data=True, directory=True, force=False)
Delete code and/or data generated/stored by the device.

Parameters
data : bool, optional
Whether to delete the data generated by the simulation (final values of state variables, data stored in monitors, etc.). Defaults to True.
code : bool, optional
Whether to delete the code generated by the simulation. Includes the numerical values used for initialization of state variables in assignments not using strings. Defaults to True.
directory : bool, optional
Whether to delete the project directory generated by the simulation. Will not delete directories that contain files not created by Brian unless the force option is specified. Defaults to True.
force : bool, optional
Whether to delete the project directory with all its content, even if it contains files that were not created by Brian. Useful only when the directory option is set to True as well. Defaults to False.

fill_with_array(var, arr)
Fill an array with the values given in another array.

Parameters
var : ArrayVariable
The array to fill.
arr: ndarray

The array values that should be copied to var().

find_synapses()
freeze(code, ns)
generate_codeobj_source(writer)
generate_main_source(writer)
generate_makefile(writer, compiler, compiler_flags, linker_flags, nb_threads, debug)
generate_network_source(writer, compiler)
generate_objects_source(writer, arange_arrays, synapses, static_array_specs, networks)
generate_run_source(writer)
generate_synapses_classes_source(writer)

get_array_filename(var, basedir='results')
Return a file name for a variable.

    Parameters var: ArrayVariable

        The variable to get a filename for.

    basedir : str

        The base directory for the filename, defaults to 'results'.

Returns:
———:

    filename : str

        A filename of the form 'results/'+varname+'_'+str(hash(varname)),
where varname is the name returned by get_array_name.

Notes

The reason that the filename is not simply 'results/' + varname is that this could lead to file names
that are not unique in file systems that are not case sensitive (e.g. on Windows).

get_array_name(var, access_data=True)
Return a globally unique name for var().

    Parameters access_data : bool, optional

        For DynamicArrayVariable objects, specifying True here means the name for
the underlying data is returned. If specifying False, the name of object itself is re-
turned (e.g. to allow resizing).

get_value(var, access_data=True)

generate_objects_source(var, start, dtype)
Initialize an array with an integer range.

    Parameters var: ArrayVariable

        The array to fill with the integer range.

    start : int
The start value for the integer range

```python
dtype: dtype
```

The data type to use for the array.

```python
init_with_zeros(var, dtype)
```

Initialize an array with zeros.

**Parameters**

- **var**: `ArrayVariable`
  - The array to initialize with zeros.

- **dtype**: `dtype`
  - The data type to use for the array.

```python
insert_code(slot, code)
```

Insert code directly into main.cpp

```python
network_get_profiling_info(net)
```

```python
network_restore(net, *args, **kwds)
```

```python
network_run(net, duration, report=None, report_period=10. * second, namespace=None, profile=False, level=0, **kwds)
```

```python
network_store(net, *args, **kwds)
```

```python
reinit()
```

Reinitialize the device. For standalone devices, clears all the internal state of the device.

```python
resize(var, new_size)
```

Resize a `DynamicArrayVariable`.

**Parameters**

- **var**: `DynamicArrayVariable`
  - The variable that should be resized.

- **new_size**: `int`
  - The new size of the variable

```python
run(directory, with_output, run_args)
```

```python
run_function(name, include_in_parent=True)
```

Context manager to divert code into a function

Code that happens within the scope of this context manager will go into the named function.

**Parameters**

- **name**: `str`
  - The name of the function to divert code into.

- **include_in_parent**: `bool`
  - Whether or not to include a call to the newly defined function in the parent context.

```python
seed(seed=None)
```

Set the seed for the random number generator.

**Parameters**

- **seed**: `int`, optional
  - The seed value for the random number generator, or `None` (the default) to set a random seed.

```python
static_array(name, arr)
```
variableview_get_subexpression_with_index_array (variableview, item, run_namespace=None)

variableview_get_with_expression (variableview, code, run_namespace=None)

variableview_set_with_index_array (variableview, item, value, check_units)

write_static_arrays (directory)

CPPWriter (project_dir)

Methods

CPPWriter class

(Shortest import: from brian2.devices.cpp_standalone.device import CPPWriter)

class brian2.devices.cpp_standalone.device.CPPWriter (project_dir)

Bases: object

Methods

write (filename, contents)

Details

write (filename, contents)

RunFunctionContext (name, include_in_parent)

RunFunctionContext class

(Shortest import: from brian2.devices.cpp_standalone.device import RunFunctionContext)

class brian2.devices.cpp_standalone.device.RunFunctionContext (name, include_in_parent)

Bases: object

Functions

invert_dict (x)

invert_dict function

(Shortest import: from brian2.devices.cpp_standalone.device import invert_dict)

brian2.devices.cpp_standalone.device.invert_dict (x)

Objects
cpp_standalone_device object

(Shortest import: from brian2.devices.cpp_standalone import cpp_standalone_device)

brian2.devices.cpp_standalone.device.cpp_standalone_device = <brian2.devices.cpp_standalone.device.CPPStandaloneDevice object>

The Device used for C++ standalone simulations.

6.6.4 equations package

Module handling equations and “code strings”, expressions or statements, used for example for the reset and threshold definition of a neuron.

Exported members: Equations, Expression, Statements

codestrings module

Module defining CodeString, a class for a string of code together with information about its namespace. Only serves as a parent class, its subclasses Expression and Statements are the ones that are actually used.

Exported members: Expression, Statements

Classes

CodeString(code) A class for representing “code strings”, i.e.

CodeString class

(Shortest import: from brian2.equations.codestrings import CodeString)

class brian2.equations.codestrings.CodeString(code)

Bases: collections.abc.Hashable

A class for representing "code strings", i.e. a single Python expression or a sequence of Python statements.

Parameters code : str

The code string, may be an expression or a statement(s) (possibly multi-line).

Attributes

code The code string

Details

code The code string
Expression class

(Shortest import: from brian2 import Expression)

class brian2.equations.codestrings.Expression (code=None, sympy_expression=None)

   Bases: brian2.equations.codestrings.CodeString

   Class for representing an expression.

   Parameters code : str, optional
       The expression. Note that the expression has to be written in a form that is parseable by sympy. Alternatively, a sympy expression can be provided (in the sympy_expression argument).

   sympy_expression : sympy expression, optional
       A sympy expression. Alternatively, a plain string expression can be provided (in the code argument).

Attributes

stochastic_variables

   Stochastic variables in this expression

Methods

split_stochastic()

   Split the expression into a stochastic and non-stochastic part.

Details

stochastic_variables

   Stochastic variables in this expression

split_stochastic()

   Split the expression into a stochastic and non-stochastic part.

   Splits the expression into a tuple of one Expression objects f (the non-stochastic part) and a dictionary mapping stochastic variables to Expression objects. For example, an expression of the form f + g * xi_1 + h * xi_2 would be returned as: (f, {'xi_1': g, 'xi_2': h}) Note that the Expression objects for the stochastic parts do not include the stochastic variable itself.

   Returns (f, d): (Expression, dict)

       A tuple of an Expression object and a dictionary, the first expression being the non-stochastic part of the equation and the dictionary mapping stochastic variables (xi or starting with xi_) to Expression objects. If no stochastic variable is present in the code string, a tuple (self, None) will be returned with the unchanged Expression object.
### Statements class

*(Shortest import: from brian2 import Statements)*

**class**  

```python
class brian2.equations.codestrings.Statements(code)
    Bases: brian2.equations.codestrings.CodeString
```

Class for representing statements.

**Parameters**  

- `code` : str  
    The statement or statements. Several statements can be given as a multi-line string or separated by semicolons.

**Notes**

Currently, the implementation of this class does not add anything to `CodeString`, but it should be used instead of that class for clarity and to allow for future functionality that is only relevant to statements and not to expressions.

**Functions**

- `is_constant_over_dt(expression, variables, ...)`: Check whether an expression can be considered as constant over a time step.

### is_constant_over_dt function

*(Shortest import: from brian2.equations.codestrings import is_constant_over_dt)*

**brian2.equations.codestrings.is_constant_over_dt(expression, variables, dt_value)**

Check whether an expression can be considered as constant over a time step. This is not the case when the expression either:

1. contains the variable $t$ (except as the argument of a function that can be considered as constant over a time step, e.g. a `TimedArray` with a dt equal to or greater than the dt used to evaluate this expression)
2. refers to a stateful function such as `rand()`.

**Parameters**

- `expression` : sympy.Expr  
    The (sympy) expression to analyze
- `variables` : dict  
    The variables dictionary.
- `dt_value` : float or None  
    The length of a timestep (without units), can be `None` if the time step is not yet known.

**Returns**

- `is_constant` : bool  
    Whether the expression can be considered to be constant over a time step.
equations module

Exported members: Equations

Classes

<table>
<thead>
<tr>
<th>EquationError</th>
<th>Exception type related to errors in an equation definition.</th>
</tr>
</thead>
</table>

EquationError class

(Shortest import: from brian2.equations.equations import EquationError)

class brian2.equations.equations.EquationError
   Bases: Exception
       Exception type related to errors in an equation definition.

<table>
<thead>
<tr>
<th>Equations(eqns, **kwds)</th>
<th>Container that stores equations from which models can be created.</th>
</tr>
</thead>
</table>

Equations class

(Shortest import: from brian2 import Equations)

class brian2.equations.equations.Equations(eqns, **kwds)
   Bases: collections.abc.Hashable, collections.abc.Mapping
       Container that stores equations from which models can be created.

String equations can be of any of the following forms:

1. \( \frac{dx}{dt} = f \) : unit (flags) (differential equation)
2. \( x = f \) : unit (flags) (equation)
3. \( x \) : unit (flags) (parameter)

String equations can span several lines and contain Python-style comments starting with 

Parameters eqs: str or list of SingleEquation objects

A multiline string of equations (see above) – for internal purposes also a list of SingleEquation objects can be given. This is done for example when adding new equations to implement the refractory mechanism. Note that in this case the variable names are not checked to allow for “internal names”, starting with an underscore.

kwds: keyword arguments:

Keyword arguments can be used to replace variables in the equation string. Arguments have to be of the form varname=replacement, where varname has to correspond to a variable name in the given equation. The replacement can be either a string (replacing a name with a new name, e.g. \( \tau = \tau_e \)) or a value (replacing the variable name with the value, e.g. \( \tau = \tau_e \) or \( \tau = 10 \text{ms} \)).
### Attributes

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>_substituted_expressions</code></td>
<td>Cache for equations with the subexpressions substituted</td>
</tr>
<tr>
<td><code>diff_eq_expressions</code></td>
<td>A list of (variable name, expression) tuples of all differential equations.</td>
</tr>
<tr>
<td><code>diff_eq_names</code></td>
<td>All differential equation names.</td>
</tr>
<tr>
<td><code>dimensions</code></td>
<td>Dictionary of all internal variables and their corresponding physical dimensions.</td>
</tr>
<tr>
<td><code>eq_expressions</code></td>
<td>A list of (variable name, expression) tuples of all equations.</td>
</tr>
<tr>
<td><code>eq_names</code></td>
<td>All equation names (including subexpressions).</td>
</tr>
<tr>
<td><code>identifier_checks</code></td>
<td>A set of functions that are used to check identifiers (class attribute).</td>
</tr>
<tr>
<td><code>identifiers</code></td>
<td>Set of all identifiers used in the equations, excluding the variables defined in the equations</td>
</tr>
<tr>
<td><code>is_stochastic</code></td>
<td>Whether the equations are stochastic.</td>
</tr>
<tr>
<td><code>names</code></td>
<td>All variable names defined in the equations.</td>
</tr>
<tr>
<td><code>ordered</code></td>
<td>A list of all equations, sorted according to the order in which they should be updated</td>
</tr>
<tr>
<td><code>parameter_names</code></td>
<td>All parameter names.</td>
</tr>
<tr>
<td><code>stochastic_type</code></td>
<td>Returns the type of stochastic differential equations (additive or multiplicative).</td>
</tr>
<tr>
<td><code>stochastic_variables</code></td>
<td>All subexpression names.</td>
</tr>
</tbody>
</table>

### Methods

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>check_flags(allowed_flags[, incompatible_flags])</code></td>
<td>Check the list of flags.</td>
</tr>
<tr>
<td><code>check_identifier(identifier)</code></td>
<td>Perform all the registered checks.</td>
</tr>
<tr>
<td><code>check_identifiers()</code></td>
<td>Check all identifiers for conformity with the rules.</td>
</tr>
<tr>
<td><code>check_units(group, run_namespace)</code></td>
<td>Check all the units for consistency.</td>
</tr>
<tr>
<td><code>get_substituted_expressions([variables, ...])</code></td>
<td>Return a list of (varname, expr) tuples, containing all differential equations and optionally subexpressions with all the subexpression variables substituted with the respective expressions.</td>
</tr>
<tr>
<td><code>register_identifier_check(func)</code></td>
<td>Register a function for checking identifiers.</td>
</tr>
<tr>
<td><code>substitute(**kwds)</code></td>
<td></td>
</tr>
</tbody>
</table>

### Details

- ** `_substituted_expressions`**
  Cache for equations with the subexpressions substituted

- ** `diff_eq_expressions`**
  A list of (variable name, expression) tuples of all differential equations.

- ** `diff_eq_names`**
  All differential equation names.

- ** `dimensions`**
Dictionary of all internal variables and their corresponding physical dimensions.

**eq_expressions**
A list of (variable name, expression) tuples of all equations.

**eq_names**
All equation names (including subexpressions).

**identifier_checks**
A set of functions that are used to check identifiers (class attribute). Functions can be registered with the static method `Equations.register_identifier_check` and will be automatically used when checking identifiers.

**identifiers**
Set of all identifiers used in the equations, excluding the variables defined in the equations.

**is_stochastic**
Whether the equations are stochastic.

**names**
All variable names defined in the equations.

**ordered**
A list of all equations, sorted according to the order in which they should be updated.

**parameter_names**
All parameter names.

**stochastic_type**
Returns the type of stochastic differential equations (additive or multiplicative). The system is only classified as additive if all equations have only additive noise (or no noise).

Returns type: str
Either `None` (no noise variables), 'additive' (factors for all noise variables are independent of other state variables or time), 'multiplicative' (at least one of the noise factors depends on other state variables and/or time).

**stochastic_variables**

**subexpr_names**
All subexpression names.

**check_flags** `(allowed_flags, incompatible_flags=None)`
Check the list of flags.

Parameters allowed_flags: dict
A dictionary mapping equation types (PARAMETER, DIFFERENTIAL_EQUATION, SUBEXPRESSION) to a list of strings (the allowed flags for that equation type).

incompatible_flags: list of tuple
A list of flag combinations that are not allowed for the same equation.

Notes:
---

Not specifying allowed flags for an equation type is the same as specifying an empty list for it.
Raise

**ValueError** If any flags are used that are not allowed.

**static check_identifier** *(identifier)*

Perform all the registered checks. Checks can be registered via `Equations`. `register_identifier_check`.

**Parameters**

`identifier` : str

The identifier that should be checked

Raise

**ValueError** If any of the registered checks fails.

**check_identifiers** ()

Check all identifiers for conformity with the rules.

Raise

**ValueError** If an identifier does not conform to the rules.

See also:

`Equations.check_identifier` The function that is called for each identifier.

**check_units** *(group, run_namespace)*

Check all the units for consistency.

**Parameters**

`group` : `Group`

The group providing the context

`run_namespace` : dict-like, optional

An additional namespace that is used for variable lookup (if not defined, the implicit namespace of local variables is used).

`level` : int, optional

How much further to go up in the stack to find the calling frame

Raise

**DimensionMismatchError** In case of any inconsistencies.

**get_substituted_expressions** *(variables=None, include_subexpressions=False)*

Return a list of `(varname, expr)` tuples, containing all differential equations (and optionally subexpressions) with all the subexpression variables substituted with the respective expressions.

**Parameters**

`variables` : dict, optional

A mapping of variable names to `Variable/Function` objects.

`include_subexpressions` : bool

Whether also to return substituted subexpressions. Defaults to `False`.

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Returns `expr_tuples` : list of (str, CodeString)

A list of (varname, expr) tuples, where expr is a CodeString object with all subexpression variables substituted with the respective expression.

```
static register_identifier_check (func)
```

Register a function for checking identifiers.

```
Parameters func : callable
```

The function has to receive a single argument, the name of the identifier to check, and raise a ValueError if the identifier violates any rule.

```
substitute (**kwds)
```

Tutorials and examples using this

- Example `COBAHH`
- Example `IF_curve_Hodgkin_Huxley`
- Example `advanced/float_32_64_benchmark`
- Example `frompapers/Rossant_et_al_2011bis`
- Example `frompapers/Diesmann_et_al_1999`
- Example `frompapers/Hindmarsh_Rose_1984`
- Example `frompapers/Clopath_et_al_2010_homeostasis`
- Example `frompapers/Clopath_et_al_2010_no_homeostasis`
- Example `frompapers/Destexhe_et_al_1998`

```
SingleEquation (type, varname, dimensions[, ...])
```

Class for internal use, encapsulates a single equation or parameter.

**SingleEquation class**

```
(Shortest import: from brian2.equations.equations import SingleEquation)

class brian2.equations.equations.SingleEquation (type, varname, dimensions, var_type='float', expr=None, flags=None)
```

```
Bases: collections.abc.Hashable, brian2.utils.caching.CacheKey
```

Class for internal use, encapsulates a single equation or parameter.

```
Note: This class should never be used directly, it is only useful as part of the Equations class.
```

```
Parameters type : {PARAMETER, DIFFERENTIAL_EQUATION, SUBEXPRESSION}
```

The type of the equation.

```
varname : str
```

The variable that is defined by this equation.

```
dimensions : Dimension
```

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The physical dimensions of the variable

```python
var_type: {FLOAT, INTEGER, BOOLEAN}
```

The type of the variable (floating point value or boolean).

```python
expr: Expression, optional
```

The expression defining the variable (or `None` for parameters).

```python
flags: list of str, optional:
```

A list of flags that give additional information about this equation. What flags are possible depends on the type of the equation and the context.

### Attributes

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>identifiers</code></td>
<td>All identifiers in the RHS of this equation.</td>
</tr>
<tr>
<td><code>stochastic_variables</code></td>
<td>Stochastic variables in the RHS of this equation.</td>
</tr>
<tr>
<td><code>unit</code></td>
<td>The <code>Unit</code> of this equation.</td>
</tr>
</tbody>
</table>

### Details

- **identifiers**: All identifiers in the RHS of this equation.
- **stochastic_variables**: Stochastic variables in the RHS of this equation
- **unit**: The `Unit` of this equation.

### Functions

**check_identifier_basic function**

```python
check_identifier_basic(identifier)
```

Check an identifier (usually resulting from an equation string provided by the user) for conformity with the rules.

**Parameters**

- **identifier**: str

The identifier that should be checked

### check_identifier_basic function

(Shortest import: from brian2.equations.equations import check_identifier_basic)
Raises

*SyntaxError*  If the identifier does not conform to the above rules.

<table>
<thead>
<tr>
<th>Function Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>check_identifier_constants</code></td>
<td>Make sure that identifier names do not clash with function names.</td>
</tr>
<tr>
<td><code>check_identifier_functions</code></td>
<td>Make sure that identifier names do not clash with function names.</td>
</tr>
<tr>
<td><code>check_identifier_reserved</code></td>
<td>Check that an identifier is not using a reserved special variable name.</td>
</tr>
<tr>
<td><code>check_identifier_units</code></td>
<td>Make sure that identifier names do not clash with unit names.</td>
</tr>
</tbody>
</table>

**check_identifier_constants function**

*(Shortest import: from brian2.equations.equations import check_identifier_constants)*

```python
brian2.equations.equations.check_identifier_constants(identifier)
```

Make sure that identifier names do not clash with function names.

**check_identifier_functions function**

*(Shortest import: from brian2.equations.equations import check_identifier_functions)*

```python
brian2.equations.equations.check_identifier_functions(identifier)
```

Make sure that identifier names do not clash with function names.

**check_identifier_reserved function**

*(Shortest import: from brian2.equations.equations import check_identifier_reserved)*

```python
brian2.equations.equations.check_identifier_reserved(identifier)
```

Check that an identifier is not using a reserved special variable name. The special variables are: `t`, `dt`, and `xi`, as well as everything starting with `xi_`.

**Parameters**

- `identifier` : `str`
  
  The identifier that should be checked

**Raises**

*SyntaxError*  If the identifier is a special variable name.
**check_identifier_units function**

(Shortest import: from brian2.equations.equations import check_identifier_units)

brian2.equations.equations.check_identifier_units(identifier)

Make sure that identifier names do not clash with unit names.

**check_subexpressions function**

(Shortest import: from brian2.equations.equations import check_subexpressions)

brian2.equations.equations.check_subexpressions(group, equations, run_namespace)

Checks the subexpressions in the equations and raises an error if a subexpression refers to stateful functions without being marked as “constant over dt”.

Parameters

- **group**: Group
  The group providing the context.
- **equations**: Equations
  The equations to check.
- **run_namespace**: dict
  The run namespace for resolving variables.

Raises

**SyntaxError** For subexpressions not marked as “constant over dt” that refer to stateful functions.

**dimensions_and_type_from_string function**

(Shortest import: from brian2.equations.equations import dimensions_and_type_from_string)

brian2.equations.equations.dimensions_and_type_from_string(unit_string)

Returns the physical dimensions that results from evaluating a string like “siemens / metre ** 2”, allowing for the special string “1” to signify dimensionless units, the string “boolean” for a boolean and “integer” for an integer variable.

Parameters

- **unit_string**: str
  The string that should evaluate to a unit
Returns `d, type:(Dimension, {FLOAT, INTEGER or BOOL})`

The resulting physical dimensions and the type of the variable.

**Raises**

`ValueError` If the string cannot be evaluated to a unit.

---

**extract_constant_subexpressions** *(eqs)*

**extract_constant_subexpressions function**

*(Shortest import: from brian2.equations.equations import extract_constant_subexpressions)*

brian2.equations.equations.extract_constant_subexpressions(eqs)

**is_stateful**(expression, variables) Whether the given expression refers to stateful functions (and is therefore not guaranteed to give the same result if called repetively).

**is_stateful function**

*(Shortest import: from brian2.equations.equations import is_stateful)*

brian2.equations.equations.is_stateful(expression, variables) Whether the given expression refers to stateful functions (and is therefore not guaranteed to give the same result if called repetively).

**Parameters**

- `expression`: sympy.Expression
  The sympy expression to check.
- `variables`: dict
  The dictionary mapping variable names to Variable or Function objects.

**Returns**

- `stateful`: bool
  True, if the given expression refers to a stateful function like `rand()` and False otherwise.

---

**parse_string_equations** *(eqns)* Parse a string defining equations.

**parse_string_equations function**

*(Shortest import: from brian2.equations.equations import parse_string_equations)*

brian2.equations.equations.parse_string_equations(eqs) Parse a string defining equations.

**Parameters**

- `eqns`: str
  The (possibly multi-line) string defining the equations. See the documentation of the
Equations class for details.

Returns equations : dict

A dictionary mapping variable names to Equations objects

refractory module

Module implementing Brian’s refractory mechanism.

Exported members: add_refractoriness

Functions

add_refractoriness(eqs)  
Extends a given set of equations with the refractory mechanism.

Parameters eqs : Equations

The equations without refractory mechanism.

Returns new_eqs : Equations

New equations, with added parameters and changed differential equations having the “unless refractory” flag.

check_identifier_refractory(identifier)  
Check that the identifier is not using a name reserved for the refractory mechanism.

Parameters identifier : str

The identifier to check.

Raises

ValueError If the identifier is a variable name used for the refractory mechanism.

6.6. Subpackages
**unitcheck module**

Utility functions for handling the units in *Equations*.

**Exported members:** `unit_from_expression`, `check_dimensions`, `check_units_statements`

**Functions**

- **check_dimensions** function

  *(Shortest import: from brian2.equations.unitcheck import check_dimensions)*

  ```python
  brian2.equations.unitcheck.check_dimensions(expression, dimensions, ...)
  ```

  Compares the physical dimensions of an expression to expected dimensions in a given namespace.

  **Parameters**

  - **expression**: str
    The expression to evaluate.
  - **dimensions**: Dimension
    The expected physical dimensions for the expression.
  - **variables**: dict
    Dictionary of all variables (including external constants) used in the expression.

  **Raises**

  - **KeyError** In case one or more identifiers cannot be resolved.
  - **DimensionMismatchError** If an unit mismatch occurs during the evaluation.

- **check_units_statements** function

  *(Shortest import: from brian2.equations.unitcheck import check_units_statements)*

  ```python
  brian2.equations.unitcheck.check_units_statements(code, variables)
  ```

  Check the units for a series of statements. Setting a model variable has to use the correct unit. For newly introduced temporary variables, the unit is determined and used to check the following statements to ensure consistency.

  **Parameters**

  - **code**: str
    The statements as a (multi-line) string
  - **variables**: dict of Variable objects
    The information about all variables used in `code` (including Constant objects for external variables)
 Raises

**KeyError**  In case one of the identifiers cannot be resolved.

**DimensionMismatchError**  If an unit mismatch occurs during the evaluation.

### 6.6.5 groups package

Package providing groups such as *NeuronGroup* or *PoissonGroup*.

*Exported members:*  *CodeRunner, Group, VariableOwner, NeuronGroup*

### group module

*Exported members:*  *Group, VariableOwner, CodeRunner*

**Classes**

| CodeRunner(group, template[, code, ...]) | A “code runner” that runs a CodeObject every timestep and keeps a reference to the Group. |

### CodeRunner class

*(Shortest import: from brian2 import CodeRunner)*

```
class brian2.groups.group.CodeRunner(group, template, code=", user_code=None, dt=None, clock=None, when='start', order=0, name='coderunner*', check_units=True, template_kwds=None, needed_variables=None, override_conditional_write=None, codeobj_class=None, generate_empty_code=True)
```

*Bases: brian2.core.base.BrianObject*

A “code runner” that runs a CodeObject every timestep and keeps a reference to the *Group*. Used in *NeuronGroup* for Thresholder, Resetter and StateUpdater.

On creation, we try to run the before_run method with an empty additional namespace (see *Network. before_run*). If the namespace is already complete this might catch unit mismatches.

**Parameters**

- **group**: *Group*
  
  The group to which this object belongs.

- **template**: *Template*
  
  The template that should be used for code generation.

- **code**: *str, optional*
  
  The abstract code that should be executed every time step. The *update_abstract_code* method might generate this code dynamically before every run instead.

- **dt**: *Quantity, optional*
  
  The time step to be used for the simulation. Cannot be combined with the *clock* argument.
**user_code** : str, optional

The abstract code as specified by the user, i.e. without any additions of internal code that the user not necessarily knows about. This will be used for warnings and error messages.

**clock** : *Clock*, optional

The update clock to be used. If neither a clock, nor the dt argument is specified, the *defaultclock* will be used.

**when** : str, optional

In which scheduling slot to execute the operation during a time step. Defaults to 'start'.

**order** : int, optional

The priority of this operation for operations occurring at the same time step and in the same scheduling slot. Defaults to 0.

**name** : str, optional

The name for this object.

**check_units** : bool, optional

Whether the units should be checked for consistency before a run. Is activated (True) by default but should be switched off for state updaters (units are already checked for the equations and the generated abstract code might have already replaced variables with their unit-less values).

**template_kwds** : dict, optional

A dictionary of additional information that is passed to the template.

**needed_variables**: list of str, optional:

A list of variables that are neither present in the abstract code, nor in the USES_VARIABLES statement in the template. This is only rarely necessary, an example being a StateMonitor where the names of the variables are neither known to the template nor included in the abstract code statements.

**override_conditional_write**: list of str, optional:

A list of variable names which are used as conditions (e.g. for refractoriness) which should be ignored.

**codeobj_class** : class, optional

The CodeObject class to run code with. If not specified, defaults to the group’s codeobj_class attribute.

**generate_empty_code** : bool, optional

Whether to generate a CodeObject if there is no abstract code to execute. Defaults to True but should be switched off e.g. for a StateUpdater when there is nothing to do.

### Methods

**before_run**(run_namespace) Optional method to prepare the object before a run.

Continued on next page
update_abstract_code \(\text{(run_namespace)}\)

Update the abstract code for the code object. Will be called in before_run and should update the CodeRunner.abstract_code attribute.

Todo

Does nothing by default.

Group(**kwds)

Methods

Group class

(Shortest import: from brian2 import Group)

class brian2.groups.group.Group(**kwds)
    Bases: brian2.groups.group.VariableOwner, brian2.core.base.BrianObject

Methods

custom_operation(*args, **kwds)
resolve_all(identifiers, run_namespace[, ...])  Resolve a list of identifiers.
run_regularly(code[, dt, clock, when, ...])  Run abstract code in the group’s namespace.
runner(*args, **kwds)

Details

custom_operation(*args, **kwds)
resolve_all (identifiers, run_namespace, user_identifiers=None, additional_variables=None)  Resolve a list of identifiers. Calls Group._resolve for each identifier.

Parameters identifiers : iterable of str
    The names to look up.

    run_namespace : dict-like, optional
        An additional namespace that is used for variable lookup (if not defined, the implicit namespace of local variables is used).

    user_identifiers : iterable of str, optional
        The names in identifiers that were provided by the user (i.e. are part of user-specified equations, abstract code, etc.). Will be used to determine when to issue
namespace conflict warnings. If not specified, will be assumed to be identical to identifiers.

**additional_variables** : dict-like, optional

An additional mapping of names to Variable objects that will be checked before Group.variables.

**Returns variables** : dict of Variable or Function

A mapping from name to Variable/Function object for each of the names given in identifiers

**Raises**

**KeyError** If one of the names in identifier cannot be resolved

```python
def run_regularly(code, dt=None, clock=None, when='start', order=0, name=None, codeobj_class=None)
```

Run abstract code in the group’s namespace. The created CodeRunner object will be automatically added to the group, it therefore does not need to be added to the network manually. However, a reference to the object will be returned, which can be used to later remove it from the group or to set it to inactive.

**Parameters code** : str

The abstract code to run.

dt : Quantity, optional

The time step to use for this custom operation. Cannot be combined with the clock argument.

clock : Clock, optional

The update clock to use for this operation. If neither a clock nor the dt argument is specified, defaults to the clock of the group.

when : str, optional

When to run within a time step, defaults to the 'start' slot.

name : str, optional

A unique name, if non is given the name of the group appended with ‘run_regularly’, ‘run_regularly_1’, etc. will be used. If a name is given explicitly, it will be used as given (i.e. the group name will not be prepended automatically).

codeobj_class : class, optional

The CodeObject class to run code with. If not specified, defaults to the group’s codeobj_class attribute.

**Returns obj** : CodeRunner

A reference to the object that will be run.
```

```
IndexWrapper class

(Shortest import: from brian2.groups.group import IndexWrapper)

class brian2.groups.group.IndexWrapper(group)
    Bases: object

    Convenience class to allow access to the indices via indexing syntax. This allows for example to get all indices for synapses originating from neuron 10 by writing synapses.indices[10, :] instead of synapses._indices.((10, slice(None)).

Indexing class

(Shortest import: from brian2.groups.group import Indexing)

class brian2.groups.group.Indexing(group, default_index='_idx')
    Bases: object

    Object responsible for calculating flat index arrays from arbitrary group- specific indices. Stores strong references to the necessary variables so that basic indexing (i.e. slicing, integer arrays/values, ...) works even when the respective VariableOwner no longer exists. Note that this object does not handle string indexing.

Methods

__call__(item, index_var)
    Return flat indices to index into state variables from arbitrary group specific indices.

Details

__call__(item=slice(None, None, None), index_var=None)
    Return flat indices to index into state variables from arbitrary group specific indices. In the default implementation, raises an error for multidimensional indices and transforms slices into arrays.

    Parameters item : slice, array, int
        The indices to translate.

    Returns indices: numpy.ndarray
        The flat indices corresponding to the indices given in item.

See also:

SynapticIndexing

VariableOwner class

(Shortest import: from brian2 import VariableOwner)
class brian2.groups.group.VariableOwner(name)

Bases: brian2.core.names.Nameable

Mix-in class for accessing arrays by attribute.

# TODO: Overwrite the __dir__ method to return the state variables # (should make autocompletion work)

Methods

add_attribute(name) Add a new attribute to this group.

check_variable_write(variable) Function that can be overwritten to raise an error if writing to a variable should not be allowed.

get_states([vars, units, format, ...]) Return a copy of the current state variable values.

set_states(values[, units, format, level]) Set the state variables.

state(name[, use_units, level]) Return the state variable in a way that properly supports indexing in the context of this group.

Details

add_attribute(name)  
Add a new attribute to this group. Using this method instead of simply assigning to the new attribute name is necessary because Brian will raise an error in that case, to avoid bugs passing unnoticed (misspelled state variable name, un-declared state variable, ...).

Parameters name : str  
The name of the new attribute

Raises

AttributeError If the name already exists as an attribute or a state variable.

check_variable_write(variable)  
Function that can be overwritten to raise an error if writing to a variable should not be allowed. Note that this does not deal with incorrect writes that are general to all kind of variables (incorrect units, writing to a read-only variable, etc.). This function is only used for type-specific rules, e.g. for raising an error in Synapses when writing to a synaptic variable before any connect call.

By default this function does nothing.

Parameters variable : Variable  
The variable that the user attempts to set.

get_states(var=\text{None}, \quad \text{units}=True, \quad \text{format}='\text{dict}', \quad \text{subexpressions}=\text{False}, \quad \text{read_only_variables}=\text{True}, \quad \text{level}=0)  
Return a copy of the current state variable values. The returned arrays are copies of the actual arrays that store the state variable values, therefore changing the values in the returned dictionary will not affect the state variables.

Parameters vars : list of str, optional  
The names of the variables to extract. If not specified, extract all state variables (except for internal variables, i.e. names that start with \text{'\_'}). If the subexpressions argument is \text{True}, the current values of all subexpressions are returned as well.

units : bool, optional
Whether to include the physical units in the return value. Defaults to True.

**format**: str, optional

The output format. Defaults to 'dict'.

**subexpressions**: bool, optional

Whether to return subexpressions when no list of variable names is given. Defaults to False. This argument is ignored if an explicit list of variable names is given in vars.

**read_only_variables**: bool, optional

Whether to return read-only variables (e.g. the number of neurons, the time, etc.). Setting it to False will assure that the returned state can later be used with `set_states`. Defaults to True.

**level**: int, optional

How much higher to go up the stack to resolve external variables. Only relevant if extracting subexpressions that refer to external variables.

**Returns values**: dict or specified format

The variables specified in vars, in the specified format.

**set_states**(values, units=True, format='dict', level=0)

Set the state variables.

**Parameters values**: depends on format

The values according to format.

**units**: bool, optional

Whether the values include physical units. Defaults to True.

**format**: str, optional

The format of values. Defaults to 'dict'

**level**: int, optional

How much higher to go up the stack to resolve external variables. Only relevant when using string expressions to set values.

**state**(name, use_units=True, level=0)

Return the state variable in a way that properly supports indexing in the context of this group

**Parameters name**: str

The name of the state variable

**use_units**: bool, optional

Whether to use the state variable’s unit.

**level**: int, optional

How much farther to go down in the stack to find the namespace.

**Returns** :

**———** :

**var**: VariableView or scalar value

The state variable’s value that can be indexed (for non-scalar values).
Functions

`get_dtype(equation[, dtype])` Helper function to interpret the `dtype` keyword argument in `NeuronGroup` etc.

---

**get_dtype function**

*(Shortest import: from brian2.groups.group import get_dtype)*

*brian2.groups.group.get_dtype(equation, dtype=None)*

Helper function to interpret the `dtype` keyword argument in `NeuronGroup` etc.

**Parameters**

- `equation`: `SingleEquation`  
  The equation for which a dtype should be returned

- `dtype`: `dtype` or dict, optional  
  Either the `dtype` to be used as a default dtype for all float variables (instead of the `core.default_float_dtype` preference) or a dictionary stating the `dtype` for some variables; all other variables will use the preference default

**Returns**

- `d`: `dtype`  
  The dtype for the variable defined in `equation`

---

**neurongroup module**

*Exported members: NeuronGroup*

**Classes**

- `NeuronGroup(N, model[, method, ...])` A group of neurons.

---

**NeuronGroup class**

*(Shortest import: from brian2 import NeuronGroup)*

*class brian2.groups.neurongroup.NeuronGroup(N, model[, method, ...])*

A group of neurons.

**Parameters**

- `N`: int  
  Number of neurons in the group.

- `model`: (str, `Equations`)  
  The differential equations defining the group

- `method`: (str, function), optional
The numerical integration method. Either a string with the name of a registered method (e.g. “euler”) or a function that receives an `Equations` object and returns the corresponding abstract code. If no method is specified, a suitable method will be chosen automatically.

**threshold** : str, optional

The condition which produces spikes. Should be a single line boolean expression.

**reset** : str, optional

The (possibly multi-line) string with the code to execute on reset.

**refractory** : {str, `Quantity`}, optional

Either the length of the refractory period (e.g. `2*ms`), a string expression that evaluates to the length of the refractory period after each spike (e.g. `'(1 + rand())*ms'`), or a string expression evaluating to a boolean value, given the condition under which the neuron stays refractory after a spike (e.g. `'v > -20*mV'`)

**events** : dict, optional

User-defined events in addition to the “spike” event defined by the `threshold`. Has to be a mapping of strings (the event name) to strings (the condition) that will be checked.

**namespace** : dict, optional

A dictionary mapping identifier names to objects. If not given, the namespace will be filled in at the time of the call of `Network.run`, with either the values from the `namespace` argument of the `Network.run` method or from the local context, if no such argument is given.

**dtype** : (dtype, dict), optional

The `numpy.dtype` that will be used to store the values, or a dictionary specifying the type for variable names. If a value is not provided for a variable (or no value is provided at all), the preference setting `core.default_float_dtype` is used.

**codeobj_class** : class, optional

The `CodeObject` class to run code with.

**dt** : `Quantity`, optional

The time step to be used for the simulation. Cannot be combined with the `clock` argument.

**clock** : `Clock`, optional

The update clock to be used. If neither a clock, nor the `dt` argument is specified, the `defaultclock` will be used.

**order** : int, optional

The priority of of this group for operations occurring at the same time step and in the same scheduling slot. Defaults to 0.

**name** : str, optional

A unique name for the group, otherwise use `neurongroup_0`, etc.
Notes

NeuronGroup contains a StateUpdater, Thresholder and Resetter, and these are run at the ‘groups’, ‘thresholds’ and ‘resets’ slots (i.e. the values of their when attribute take these values). The order attribute will be passed down to the contained objects but can be set individually by setting the order attribute of the state_updater, thresholder and resetter attributes, respectively.

Attributes

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>_refractory</td>
<td>The refractory condition or timespan</td>
</tr>
<tr>
<td>event_codes</td>
<td>Code that is triggered on events (e.g. reset)</td>
</tr>
<tr>
<td>events</td>
<td>Events supported by this group</td>
</tr>
<tr>
<td>method_choice</td>
<td>The state update method selected by the user</td>
</tr>
<tr>
<td>namespace</td>
<td>The group-specific namespace</td>
</tr>
<tr>
<td>resetter</td>
<td>Reset neurons which have spiked (or perform arbitrary actions for user-defined events)</td>
</tr>
<tr>
<td>spikes</td>
<td>The spikes returned by the most recent thresholding operation.</td>
</tr>
<tr>
<td>state_updater</td>
<td>Performs numerical integration step</td>
</tr>
<tr>
<td>subexpression_updater</td>
<td>Update the “constant over a time step” subexpressions</td>
</tr>
<tr>
<td>thresholder</td>
<td>Checks the spike threshold (or arbitrary user-defined events)</td>
</tr>
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<td>user_equations</td>
<td>The original equations as specified by the user (i.e.</td>
</tr>
</tbody>
</table>

Methods

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
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<tbody>
<tr>
<td>before_run([run_namespace])</td>
<td>Optional method to prepare the object before a run.</td>
</tr>
<tr>
<td>run_on_event(event, code[, when, order])</td>
<td>Run code triggered by a custom-defined event (see NeuronGroup documentation for the specification of events). The created Resetter object will be automatically added to the group, it therefore does not need to be added to the network manually.</td>
</tr>
<tr>
<td>set_event_schedule(event[, when, order])</td>
<td>Change the scheduling slot for checking the condition of an event.</td>
</tr>
<tr>
<td>state(name[, use_units, level])</td>
<td>Return the state variable in a way that properly supports indexing in the context of this group</td>
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namespace
    The group-specific namespace

resetter
    Reset neurons which have spiked (or perform arbitrary actions for user-defined events)

spikes
    The spikes returned by the most recent thresholding operation.

state_updater
    Performs numerical integration step

subexpression_updater
    Update the “constant over a time step” subexpressions

thresholder
    Checks the spike threshold (or arbitrary user-defined events)

user_equations
    The original equations as specified by the user (i.e. without the multiplied \( \text{int}(\text{not_refractory}) \) term for equations marked as \( \text{unless refractory} \))

before_run \((\text{run_namespace} = \text{None})\)
    Optional method to prepare the object before a run.
    TODO

run_on_event \((\text{event, code, when} = \text{'after_resets'}, \text{order} = \text{None})\)
    Run code triggered by a custom-defined event (see NeuronGroup documentation for the specification of events). The created Resetter object will be automatically added to the group, it therefore does not need to be added to the network manually. However, a reference to the object will be returned, which can be used to later remove it from the group or to set it to inactive.

    Parameters event : str
        The name of the event that should trigger the code
    code : str
        The code that should be executed
    when : str, optional
        The scheduling slot that should be used to execute the code. Defaults to 'after_resets'.
    order : int, optional
        The order for operations in the same scheduling slot. Defaults to the order of the NeuronGroup.

    Returns obj : Resetter
        A reference to the object that will be run.

set_event_schedule \((\text{event, when} = \text{'after_thresholds'}, \text{order} = \text{None})\)
    Change the scheduling slot for checking the condition of an event.

    Parameters event : str
        The name of the event for which the scheduling should be changed
    when : str, optional
        The scheduling slot that should be used to check the condition. Defaults to 'after_thresholds'.
order : int, optional
The order for operations in the same scheduling slot. Defaults to the order of the
NeuronGroup.

state(name, use_units=True, level=0)
Return the state variable in a way that properly supports indexing in the context of this group

Parameters name : str
The name of the state variable

use_units : bool, optional
Whether to use the state variable’s unit.

level : int, optional
How much farther to go down in the stack to find the namespace.

Returns :

var : VariableView or scalar value
The state variable’s value that can be indexed (for non-scalar values).

Tutorials and examples using this

• Tutorial 2-intro-to-brian-synapses
• Tutorial 3-intro-to-brian-simulations
• Tutorial 1-intro-to-brian-neurons
• Example reliability
• Example IF_curve_LIF
• Example phase_locking
• Example COBAHH
• Example non_reliability
• Example IF_curve_Hodgkin_Huxley
• Example CUBA
• Example adaptive_threshold
• Example compartmental/bipolar_with_inputs
• Example compartmental/bipolar_with_inputs2
• Example compartmental/lfp
• Example advanced/stochastic_odes
• Example advanced/compare_GSL_to_conventional
• Example advanced/float_32_64_benchmark
• Example advanced/exprel_function
• Example advanced/custom_events
• Example advanced/opencv_movie
• Example synapses/licklider
• Example synapses/efficient_gaussian_connectivity
• Example synapses/gapjunctions
• Example synapses/nonlinear
• Example synapses/synapses
• Example synapses/state_variables
• Example synapses/jeffress
• Example synapses/spatial_connections
• Example synapses/STDP
• Example standalone/cuba_openmp
• Example standalone/STDP_standalone
• Example frompapers/Rossant_et_al_2011bis
• Example frompapers/Diesmann_et_al_1999
• Example frompapers/Hindmarsh_Rose_1984
• Example frompapers/Clopath_et_al_2010_homeostasis
• Example frompapers/Wang_Buszaki_1996
• Example frompapers/Morris_Lecar_1981
• Example frompapers/Izhikevich_2007
• Example frompapers/Vogels_et_al_2011
• Example frompapers/Brette_Gerstner_2005
• Example frompapers/Brette_Guigon_2003
• Example frompapers/Kremer_et_al_2011_barrel_cortex
• Example frompapers/Rothman_Manis_2003
• Example frompapers/Brunel_Hakim_1999
• Example frompapers/Clopath_et_al_2010_no_homeostasis
• Example frompapers/Touboul_Brette_2008
• Example frompapers/Sturzl_et_al_2000
• Example frompapers/Platkiewicz_Brette_2011
• Example frompapers/Brette_2004
• Example frompapers/Brunel_Wang_2001
• Example frompapers/Stimberg_et_al_2018/example_6_COBA_with_astro
• Example frompapers/Stimberg_et_al_2018/example_1_COBA
• Example frompapers/Stimberg_et_al_2018/example_5_astro_ring
• Example frompapers/Stimberg_et_al_2018/example_2_gchi_astrocyte
• Example frompapers/Stimberg_et_al_2018/example_4_synrel
Reseter class

(Shortest import: from brian2.groups.neurongroup import Reseter)

class brian2.groups.neurongroup.Reseter(group[, when=’resets’, order=None, event=’spike’]):

Bases: brian2.groups.group.CodeRunner

The CodeRunner that applies the reset statement(s) to the state variables of neurons that have spiked in this timestep.

Methods

update_abstract_code(run_namespace)

Update the abstract code for the code object.

Details

update_abstract_code (run_namespace)

Update the abstract code for the code object. Will be called in before_run and should update the CodeRunner.abstract_code attribute.

Does nothing by default.

StateUpdater class

(Shortest import: from brian2.groups.neurongroup import StateUpdater)

class brian2.groups.neurongroup.StateUpdater(group, method[, method_options=None]):

Bases: brian2.groups.group.CodeRunner

The CodeRunner that updates the state variables of a NeuronGroup at every timestep.

Methods

update_abstract_code(run_namespace)

Update the abstract code for the code object.
Details

**update_abstract_code** *(run_namespace)*

Update the abstract code for the code object. Will be called in `before_run` and should update the `CodeRunner.abstract_code` attribute.

Does nothing by default.

<table>
<thead>
<tr>
<th>SubexpressionUpdater <strong>group</strong>, <strong>subexpressions</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>The <code>CodeRunner</code> that updates the state variables storing the values of subexpressions that have been marked as &quot;constant over dt&quot;.</td>
</tr>
</tbody>
</table>

SubexpressionUpdater class

(Shortest import: `from brian2.groups.neurongroup import SubexpressionUpdater`)

class brian2.groups.neurongroup.SubexpressionUpdater *(group, **subexpressions**, when='before_start')*

Bases: brian2.groups.group.CodeRunner

The `CodeRunner` that updates the state variables storing the values of subexpressions that have been marked as "constant over dt".

<table>
<thead>
<tr>
<th>Thresholder <strong>group</strong>, <strong>when</strong>, <strong>event</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>The <code>CodeRunner</code> that applies the threshold condition to the state variables of a <code>NeuronGroup</code> at every timestep and sets its <code>spikes</code> and <code>refractory_until</code> attributes.</td>
</tr>
</tbody>
</table>

Threshold class

(Shortest import: `from brian2.groups.neurongroup import Thresholder`)

class brian2.groups.neurongroup.Thresholder *(group, when='thresholds', event='spike')*

Bases: brian2.groups.group.CodeRunner

The `CodeRunner` that applies the threshold condition to the state variables of a `NeuronGroup` at every timestep and sets its `spikes` and `refractory_until` attributes.

Methods

**update_abstract_code** *(run_namespace)*

Update the abstract code for the code object.

Details

**update_abstract_code** *(run_namespace)*

Update the abstract code for the code object. Will be called in `before_run` and should update the `CodeRunner.abstract_code` attribute.

Does nothing by default.

Functions

6.6. Subpackages
**check_identifier_pre_post** function

(Shortest import: from brian2.groups.neurongroup import check_identifier_pre_post)

brian2.groups.neurongroup.\texttt{check_identifier_pre_post}\((\texttt{identifier})\)

Do not allow names ending in \_pre or \_post to avoid confusion.

---

**to_start_stop** function

(Shortest import: from brian2.groups.neurongroup import to_start_stop)

brian2.groups.neurongroup.\texttt{to_start_stop}\((\texttt{item, N})\)

Helper function to transform a single number, a slice or an array of contiguous indices to a start and stop value. This is used to allow for some flexibility in the syntax of specifying subgroups in \texttt{NeuronGroup} and \texttt{SpatialNeuron}.

**Parameters**

- **item**: slice, int or sequence
  
  The slice, index, or sequence of indices to use. Note that a sequence of indices has to be a sorted ascending sequence of subsequent integers.

- **N**: int
  
  The total number of elements in the group.

**Returns**

- **start**: int
  
  The start value of the slice.

- **stop**: int
  
  The stop value of the slice.

**Examples**

```python
>>> from brian2.groups.neurongroup import to_start_stop
>>> to_start_stop(slice(3, 6), 10)
(3, 6)
>>> to_start_stop(slice(3, None), 10)
(3, 10)
>>> to_start_stop(5, 10)
(5, 6)
>>> to_start_stop([3, 4, 5], 10)
Traceback (most recent call last):
  File "stdin", line 1, in <module>
IndexError: Subgroups can only be constructed using contiguous indices.
```
**subgroup module**

Exported members: `Subgroup`

Classes

```python
Subgroup(source, start, stop[, name]) Subgroup of any Group
```

**Subgroup class**

*(Shortest import: from brian2 import Subgroup)*

class brian2.groups.subgroup.Subgroup (source, start, stop, name=None)

Bases: brian2.groups.group.Group, brian2.core.spikesource.SpikeSource

Subgroup of any Group

Parameters

- `source : SpikeSource`
  The source object to subgroup.
- `start, stop : int`
  Select only spikes with indices from `start` to `stop-1`.
- `name : str, optional`
  A unique name for the group, or use `source.name+'_subgroup_0'`, etc.

Attributes

- `spikes`

Details

- `spikes`

**6.6.6 importexport package**

Package providing import/export support.

Exported members: `ImportExport`

**dictlike module**

Module providing `DictImportExport` and `PandasImportExport` (requiring a working installation of pandas).

Classes

```python
DictImportExport An importer/exporter for variables in format of dict of numpy arrays.
```
DictImportExport class

(Shortest import: from brian2.importexport import DictImportExport)

class brian2.importexport.dictlike.DictImportExport
    Bases: brian2.importexport.importexport.ImportExport

    An importer/exporter for variables in format of dict of numpy arrays.

Attributes

<table>
<thead>
<tr>
<th>name</th>
<th>Abstract property giving a method name.</th>
</tr>
</thead>
</table>

Methods

<table>
<thead>
<tr>
<th>export_data</th>
<th>Abstract static export data method with two obligatory parameters.</th>
</tr>
</thead>
<tbody>
<tr>
<td>import_data</td>
<td>Import and set state variables.</td>
</tr>
</tbody>
</table>

Details

<table>
<thead>
<tr>
<th>name</th>
<th>Abstract property giving a method name.</th>
</tr>
</thead>
</table>

static export_data(group, variables[, units, level])

Abstract static export data method with two obligatory parameters. It should return a copy of the current state variable values. The returned arrays are copies of the actual arrays that store the state variable values, therefore changing the values in the returned dictionary will not affect the state variables.

Parameters:
- **group**: Group
  - Group object.
- **variables**: list of str
  - The names of the variables to extract.

static import_data(group, data[, units, level])

Import and set state variables.

Parameters:
- **group**: Group
  - Group object.
- **data**: dict_like
  - Data to import with variable names.

PandasImportExport

An importer/exporter for variables in pandas DataFrame format.

PandasImportExport class

(Shortest import: from brian2.importexport import PandasImportExport)
class brian2.importexport.dictlike.PandasImportExport
Bases: brian2.importexport.importexport.ImportExport

An importer/exporter for variables in pandas DataFrame format.

Attributes

- **name**: Abstract property giving a method name.

Methods

- **export_data**(group, variables[, units, level]) Abstract static export data method with two obligatory parameters.
- **import_data**(group, data[, units, level]) Import and set state variables.

Details

- **name**: Abstract property giving a method name.

**static export_data**(group, variables, units=True, level=0)
Abstract static export data method with two obligatory parameters. It should return a copy of the current state variable values. The returned arrays are copies of the actual arrays that store the state variable values, therefore changing the values in the returned dictionary will not affect the state variables.

- **Parameters**
  - **group**: Group
  - **variables**: list of str

**static import_data**(group, data, units=True, level=0)
Import and set state variables.

- **Parameters**
  - **group**: Group
  - **data**: dict_like

importexport module

Module defining the `ImportExport` class that enables getting state variable data in and out of groups in various formats (see `Group.get_states` and `Group.set_states`).

Classes

- **ImportExport**: Class for registering new import/export methods (via static methods).
ImportExport class

(Shortest import: from brian2 import ImportExport)

class brian2.importexport.importexport.ImportExport
   Bases: object

   Class for registering new import/export methods (via static methods). Also the base class that should
   be extended for such methods (ImportExport.export_data, ImportExport.import_data, and
   ImportExport.name have to be overwritten).

   See also:
   VariableOwner.get_states, VariableOwner.set_states

Attributes

| methods     | A dictionary mapping import/export methods names to ImportExport objects |
| name        | Abstract property giving a method name.                               |

Methods

| export_data(group, variables) | Asbtract static export data method with two obligatory parameters. |
| import_data(group, data)     | Import and set state variables.                                    |
| register(importerexporter)   | Register a import/export method.                                   |

Details

methods
   A dictionary mapping import/export methods names to ImportExport objects

name
   Abstract property giving a method name.

static export_data(group, variables)
   Asbtract static export data method with two obligatory parameters. It should return a copy of the current
   state variable values. The returned arrays are copies of the actual arrays that store the state variable values,
   therefore changing the values in the returned dictionary will not affect the state variables.

   Parameters group : Group
      Group object.

   variables : list of str
      The names of the variables to extract.

static import_data(group, data)
   Import and set state variables.

   Parameters group : Group
      Group object.

   data : dict_like
Data to import with variable names.

**static register**(importerexporter)

Register a import/export method. Registered methods can be referred to via their name.

**Parameters** importerexporter: ImportExport

The importerexporter object, e.g. an DictImportExport.

### 6.6.7 input package

Classes for providing external input to a network.

**Exported members:** BinomialFunction, PoissonGroup, PoissonInput, SpikeGeneratorGroup, TimedArray

#### binomial module

Implementation of BinomialFunction

**Exported members:** BinomialFunction

**Classes**

| BinomialFunction(n, p[, approximate, name]) | A function that generates samples from a binomial distribution. |

#### BinomialFunction class

*(Shortest import: from brian2 import BinomialFunction)*

**class** brian2.input.binomial.BinomialFunction(n, p, approximate=True, name='binomial*')

**Bases:** brian2.core.functions.Function, brian2.core.names.Nameable

A function that generates samples from a binomial distribution.

**Parameters**

- **n** : int
  
  Number of samples

- **p** : float
  
  Probablility

  **approximate** : bool, optional

  Whether to approximate the binomial with a normal distribution if \( np > 5 \land n(1-p) > 5 \). Defaults to True.

**Attributes**
**implementations**

Container for implementing functions for different targets. This container can be extended by other code-generation targets/devices. The key has to be the name of the target, the value a function that takes three parameters (n, p, use_normal) and returns a tuple of (code, dependencies).

---

**Details**

**implementations**

Container for implementing functions for different targets. This container can be extended by other code-generation targets/devices. The key has to be the name of the target, the value a function that takes three parameters (n, p, use_normal) and returns a tuple of (code, dependencies).

---

**poissongroup module**

*Exported members:* PoissonGroup

**Classes**

| PoissonGroup(**kwds**) | Poisson spike source |

---

**PoissonGroup class**

(*Shortest import:* from brian2 import PoissonGroup)

```python
class brian2.input.poissongroup.PoissonGroup(**kwds)
    Bases: brian2.groups.group.Group, brian2.core.spikesource.SpikeSource
```

Poisson spike source

**Parameters**

- **N**: int
  
  Number of neurons

- **rates**: Quantity, str
  
  Single rate, array of rates of length N, or a string expression evaluating to a rate. This string expression will be evaluated at every time step, it can therefore be time-dependent (e.g. refer to a TimedArray).

- **dt**: Quantity, optional
  
  The time step to be used for the simulation. Cannot be combined with the clock argument.

- **clock**: Clock, optional
  
  The update clock to be used. If neither a clock, nor the dt argument is specified, the defaultclock will be used.

- **when**: str, optional
  
  When to run within a time step, defaults to the 'thresholds' slot.

- **order**: int, optional
The priority of this group for operations occurring at the same time step and in the same scheduling slot. Defaults to 0.

**name**: str, optional

Unique name, or use poissongroup, poissongroup_1, etc.

### Attributes

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>namespace</td>
<td>The group-specific namespace</td>
</tr>
<tr>
<td>spikes</td>
<td>The spikes returned by the most recent thresholding operation.</td>
</tr>
</tbody>
</table>

### Methods

**before_run([run_namespace])**

Optional method to prepare the object before a run.

### Details

**namespace**

The group-specific namespace

**spikes**

The spikes returned by the most recent thresholding operation.

**before_run**(run_namespace=None)

Optional method to prepare the object before a run.

TODO

### Tutorials and examples using this

- Tutorial 3-intro-to-brian-simulations
- Example adaptive_threshold
- Example advanced/custom_events
- Example synapses/STDP
- Example standalone/STDP_standalone
- Example frompapers/Stimberg_et_al_2018/example_4_synrel
- Example frompapers/Stimberg_et_al_2018/example_4_rsmean

### poissoninput module

*Exported members: PoissonInput*

*Classes*

| PoissonInput(target, target_var, N, rate, weight) | Adds independent Poisson input to a target variable of a Group. |

---

6.6. Subpackages
PoissonInput class

(Shortest import: from brian2 import PoissonInput)

class brian2.input.poissoninput.PoissonInput(target, target_var, N, rate, weight, when='synapses', order=0)
Bases: brian2.groups.group.CodeRunner

Adds independent Poisson input to a target variable of a Group. For large numbers of inputs, this is much more efficient than creating a PoissonGroup. The synaptic events are generated randomly during the simulation and are not preloaded and stored in memory. All the inputs must target the same variable, have the same frequency and same synaptic weight. All neurons in the target Group receive independent realizations of Poisson spike trains.

Parameters target : Group
The group that is targeted by this input.

target_var : str
The variable of target that is targeted by this input.

N : int
The number of inputs

rate : Quantity
The rate of each of the inputs

weight : str or Quantity
Either a string expression (that can be interpreted in the context of target) or a Quantity that will be added for every event to the target_var of target. The unit has to match the unit of target_var

when : str, optional
When to update the target variable during a time step. Defaults to the synapses scheduling slot.

order : int, optional
The priority of of the update compared to other operations occurring at the same time step and in the same scheduling slot. Defaults to 0.

Attributes

<table>
<thead>
<tr>
<th>N</th>
<th>The number of inputs</th>
</tr>
</thead>
<tbody>
<tr>
<td>rate</td>
<td>The rate of each input</td>
</tr>
</tbody>
</table>

Methods

before_run(run_namespace) Optional method to prepare the object before a run.

Details

N
The number of inputs
rate
   The rate of each input

before_run (run_namespace)
   Optional method to prepare the object before a run.
   TODO

Tutorials and examples using this

• Example frompapers/Rossant_et_al_2011bis
• Example frompapers/Brunel_Wang_2001

spikegeneratorgroup module

Module defining SpikeGeneratorGroup.

Exported members: SpikeGeneratorGroup

Classes

SpikeGeneratorGroup(N, indices, times[, dt, . . . ])
A group emitting spikes at given times.

SpikeGeneratorGroup class

(Shortest import: from brian2 import SpikeGeneratorGroup)

class brian2.input.spikegeneratorgroup.SpikeGeneratorGroup(N, indices, times, dt=None, clock=None, period=0*second, when='thresholds', order=0, sorted=False, name='spikegeneratorgroup*', codeobj_class=None)

Bases: brian2.groups.group.Group, brian2.groups.group.CodeRunner, brian2.core.spikesource.SpikeSource

A group emitting spikes at given times.

Parameters
    N : int
       The number of “neurons” in this group
    indices : array of integers
       The indices of the spiking cells
    times : Quantity
       The spike times for the cells given in indices. Has to have the same length as indices.
    period : Quantity, optional
If this is specified, it will repeat spikes with this period. A period of 0s means not repeating spikes.

**dt**: Quantity, optional

The time step to be used for the simulation. Cannot be combined with the clock argument.

**clock**: Clock, optional

The update clock to be used. If neither a clock, nor the dt argument is specified, the defaultclock will be used.

**when**: str, optional

When to run within a time step, defaults to the 'thresholds' slot.

**order**: int, optional

The priority of of this group for operations occurring at the same time step and in the same scheduling slot. Defaults to 0.

**sorted**: bool, optional

Whether the given indices and times are already sorted. Set to True if your events are already sorted (first by spike time, then by index), this can save significant time at construction if your arrays contain large numbers of spikes. Defaults to False.

**Notes**

- If sorted is set to True, the given arrays will not be copied (only affects runtime mode).

**Attributes**

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>_neuron_index</td>
<td>Array of spiking neuron indices.</td>
</tr>
<tr>
<td>_previous_dt</td>
<td>Remember the dt we used the last time when we checked the spike bins to not repeat the work for multiple runs with the same dt</td>
</tr>
<tr>
<td>_spike_time</td>
<td>Array of spiking neuron times.</td>
</tr>
<tr>
<td>_spikes_changed</td>
<td>“Dirty flag” that will be set when spikes are changed after the before_run check</td>
</tr>
<tr>
<td>spikes</td>
<td>The spikes returned by the most recent thresholding operation.</td>
</tr>
</tbody>
</table>

**Methods**

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>before_run</td>
<td>Optional method to prepare the object before a run.</td>
</tr>
<tr>
<td>set_spikes</td>
<td>Change the spikes that this group will generate.</td>
</tr>
</tbody>
</table>

**Details**

- _neuron_index
  - Array of spiking neuron indices.

- _previous_dt
  - Remember the dt we used the last time when we checked the spike bins to not repeat the work for multiple runs with the same dt.
runs with the same dt

_spikes_time
Array of spiking neuron times.

_spikes_changed
“Dirty flag” that will be set when spikes are changed after the before_run check

spikes
The spikes returned by the most recent thresholding operation.

before_run (run_namespace)
Optional method to prepare the object before a run.

TODO

set_spikes (indices, times, period=0*second, sorted=False)
Change the spikes that this group will generate.

This can be used to set the input for a second run of a model based on the output of a first run (if the input for the second run is already known before the first run, then all the information should simply be included in the initial SpikeGeneratorGroup initializer call, instead).

Parameters

indices : array of integers
The indices of the spiking cells

times : Quantity
The spike times for the cells given in indices. Has to have the same length as indices.

period : Quantity, optional
If this is specified, it will repeat spikes with this period. A period of 0s means not repeating spikes.

sorted : bool, optional
Whether the given indices and times are already sorted. Set to True if your events are already sorted (first by spike time, then by index), this can save significant time at construction if your arrays contain large numbers of spikes. Defaults to False.

Tutorials and examples using this

- Tutorial 3-intro-to-brian-simulations
- Example frompapers/Diesmann_et_al_1999
- Example frompapers/Izhikevich_2007
- Example frompapers/Stimberg_et_al_2018/example_3_io_synapse

timedarray module

Implementation of TimedArray.

Exported members: TimedArray

Classes
TimedArray class

*(Shortest import: from brian2 import TimedArray)*

class brian2.input.timedarray.TimedArray(values, dt[, name=None])


A function of time built from an array of values. The returned object can be used as a function, including in model equations etc. The resulting function has to be called as function_name(t) if the provided value array is one-dimensional and as function_name(t, i) if it is two-dimensional.

Parameters

values : ndarray or Quantity

An array of values providing the values at various points in time. This array can either be one- or two-dimensional. If it is two-dimensional it's first dimension should be the time.

dt : Quantity

The time distance between values in the values array.

name : str, optional

A unique name for this object, see Nameable for details. Defaults to '_timedarray*'.

Notes

For time values corresponding to elements outside of the range of values provided, the first respectively last element is returned.

Examples

```python
>>> from brian2 import *
>>> ta = TimedArray([1, 2, 3, 4] * mV, dt=0.1*ms)
>>> print(ta(0.3*ms))
4. mV
>>> G = NeuronGroup(1, 'v = ta(t) : volt')
>>> mon = StateMonitor(G, 'v', record=True)
>>> net = Network(G, mon)
>>> net.run(1*ms)  # doctest: +ELLIPSIS...
... >>> print(mon[0].v)
[ 1.  2.  3.  4.  4.  4.  4.  4.  4.  4.  4.  4.  4.  4.  4.  4.  4.  4.  4.  4. ] mV
>>> ta2d = TimedArray([[1, 2], [3, 4], [5, 6]]*mV, dt=0.1*ms)
>>> G = NeuronGroup(4, 'v = ta2d(t, i%2) : volt')
>>> mon = StateMonitor(G, 'v', record=True)
>>> net = Network(G, mon)
>>> net.run(0.2*ms)  # doctest: +ELLIPSIS...
... >>> print(mon.v[:])
[[ 1.  3.]]
```
Methods

**is_locally_constant**(dt)  
Return whether this function (if interpreted as a function of time) should be considered constant over a timestep.

**Details**

**is_locally_constant**(dt)  
Return whether this function (if interpreted as a function of time) should be considered constant over a timestep. This is most importantly used by *TimedArray* so that linear integration can be used. In its standard implementation, always returns False.

**Parameters**  
dt : float  
The length of a timestep (without units).

**Returns**  
customant : bool  
Whether the results of this function can be considered constant over one timestep of length dt.

**Tutorials and examples using this**

- Tutorial 3-intro-to-brian-simulations
- Example synapses/jeffress
- Example frompapers/Sturzl_et_al_2000
- Example frompapers/Brunel_Wang_2001
- Example frompapers/Stimberg_et_al_2018/example_6_COBA_with_astro
- Example frompapers/Stimberg_et_al_2018/example_4_synrel

**6.6.8 memory package**

dynamicarray module

TODO: rewrite this (verbatim from Brian 1.x), more efficiency

**Exported members:** DynamicArray, DynamicArray1D

**Classes**

DynamicArray(shape[, dtype, factor, ...])  
An N-dimensional dynamic array class
DynamicArray class

(Shortest import: from brian2.memory.dynamicarray import DynamicArray)

class brian2.memory.dynamicarray.DynamicArray(shape, dtype=<class 'float'>, factor=2, use_numpy_resize=False, refcheck=True)

Bases: object

An N-dimensional dynamic array class

The array can be resized in any dimension, and the class will handle allocating a new block of data and copying when necessary.

Warning: The data will NOT be contiguous for >1D arrays. To ensure this, you will either need to use 1D arrays, or to copy the data, or use the shrink method with the current size (although note that in both cases you negate the memory and efficiency benefits of the dynamic array).

Initialisation arguments:

shape, dtype The shape and dtype of the array to initialise, as in Numpy. For 1D arrays, shape can be a single int, for ND arrays it should be a tuple.

factor The resizing factor (see notes below). Larger values tend to lead to more wasted memory, but more computationally efficient code.

use_numpy_resize, refcheck Normally, when you resize the array it creates a new array and copies the data. Sometimes, it is possible to resize an array without a copy, and if this option is set it will attempt to do this. However, this can cause memory problems if you are not careful so the option is off by default. You need to ensure that you do not create slices of the array so that no references to the memory exist other than the main array object. If you are sure you know what you’re doing, you can switch this reference check off. Note that resizing in this way is only done if you resize in the first dimension.

The array is initialised with zeros. The data is stored in the attribute data which is a Numpy array.

Some numpy methods are implemented and can work directly on the array object, including len(arr), arr[...] and arr[...] = .... In other cases, use the data attribute.

Notes

The dynamic array returns a data attribute which is a view on the larger _data attribute. When a resize operation is performed, and a specific dimension is enlarged beyond the size in the _data attribute, the size is increased to the larger of cursize*factor and newsize. This ensures that the amortized cost of increasing the size of the array is O(1).

Examples

>>> x = DynamicArray((2, 3), dtype=int)
>>> x[:1] = 1
>>> x.resize((3, 3))
>>> x[:1] += 1
>>> x.resize((3, 4))
>>> x[:1] += 1
>>> x.resize((4, 4))
>>> x[:1] += 1

(continues on next page)
### Methods

**resize**(newshape)  
Resizes the data to the new shape, which can be a different size to the current data, but should have the same rank, i.e.

**resize_along_first**(newshape)

**shrink**(newshape)  
Reduces the data to the given shape, which should be smaller than the current shape.

### Details

**resize**(newshape)  
Resizes the data to the new shape, which can be a different size to the current data, but should have the same rank, i.e. same number of dimensions.

**resize_along_first**(newshape)  
Reduces the data to the given shape, which should be smaller than the current shape. **resize()** can also be used with smaller values, but it will not shrink the allocated memory, whereas **shrink** will reallocate the memory. This method should only be used infrequently, as if it is used frequently it will negate the computational efficiency benefits of the DynamicArray.

**DynamicArray1D**(shape[, dtype, factor, ...])  
Version of **DynamicArray** with specialised **resize** method designed to be more efficient.

### DynamicArray1D class

(Shortest import: from brian2.memory.dynamicarray import DynamicArray1D)

**class** brian2.memory.dynamicarray.DynamicArray1D**(shape[, dtype=<class 'float'>, factor=2, use_numpy_resize=False, refcheck=True])**

Bases: brian2.memory.dynamicarray.DynamicArray

Version of **DynamicArray** with specialised **resize** method designed to be more efficient.

### Methods

**resize**(newshape)  
Resizes the data to the new shape, which can be a different size to the current data, but should have the same rank, i.e.
Details

**resize**(newshape)

Resizes the data to the new shape, which can be a different size to the current data, but should have the same rank, i.e. same number of dimensions.

Functions

**getslcies**(shape[, from_start])

**getslcies function**

*(Shortest import: from brian2.memory.dynamicarray import getslices)*

brian2.memory.dynamicarray.getslcies(shape, from_start=True)

### 6.6.9 monitors package

Base package for all monitors, i.e. objects to record activity during a simulation run.

**Exported members:** `SpikeMonitor, EventMonitor, StateMonitor, PopulationRateMonitor`

**ratemonitor module**

**Exported members:** `PopulationRateMonitor`

**Classes**

<table>
<thead>
<tr>
<th>PopulationRateMonitor(source[, name, ...])</th>
<th>Record instantaneous firing rates, averaged across neurons from a <code>NeuronGroup</code> or other spike source.</th>
</tr>
</thead>
</table>

**PopulationRateMonitor class**

*(Shortest import: from brian2 import PopulationRateMonitor)*

class brian2.monitors.ratemonitor.PopulationRateMonitor(source, name='ratemonitor*', codeobj_class=None, dtype=<class 'numpy.float64'>)  

Bases: `brian2.groups.group.Group, brian2.groups.group.CodeRunner`

Record instantaneous firing rates, averaged across neurons from a `NeuronGroup` or other spike source.

**Parameters**

<table>
<thead>
<tr>
<th>source</th>
<th><em>(NeuronGroup, SpikeSource)</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>name</td>
<td>str, optional</td>
</tr>
<tr>
<td>codeobj_class</td>
<td>class, optional</td>
</tr>
</tbody>
</table>

The source of spikes to record.

A unique name for the object, otherwise will use `source.name+_ratemonitor_0`, etc.
The `CodeObject` class to run code with.

**dtype**: dtype, optional

The dtype to use to store the `rate` variable. Defaults to `float64`, i.e. double precision.

**Notes**:

Currently, this monitor can only monitor the instantaneous firing rates at each time step of the source clock. Any binning/smoothing of the firing rates has to be done manually afterwards.

**Attributes**

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>source</code></td>
<td>The group we are recording from</td>
</tr>
</tbody>
</table>

**Methods**

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>reinit()</code></td>
<td>Clears all recorded rates</td>
</tr>
<tr>
<td><code>resize(new_size)</code></td>
<td></td>
</tr>
<tr>
<td><code>smooth_rate(self, window, width)</code></td>
<td>Return a smooth version of the population rate.</td>
</tr>
</tbody>
</table>

**Details**

**source**

The group we are recording from

**reinit()**

Clears all recorded rates

**resize(new_size)**

**smooth_rate** *(self, window='gaussian', width=None)*

Return a smooth version of the population rate.

**Parameters**

- `window` : str, ndarray
  The window to use for smoothing. Can be a string to chose a predefined window (`'flat'` for a rectangular, and `'gaussian'` for a Gaussian-shaped window). In this case the width of the window is determined by the `width` argument. Note that for the Gaussian window, the `width` parameter specifies the standard deviation of the Gaussian, the width of the actual window is `4*width + dt` (rounded to the nearest `dt`). For the flat window, the width is rounded to the nearest odd multiple of `dt` to avoid shifting the rate in time. Alternatively, an arbitrary window can be given as a numpy array (with an odd number of elements). In this case, the width in units of time depends on the `dt` of the simulation, and no `width` argument can be specified. The given window will be automatically normalized to a sum of 1.

- `width` : `Quantity`, optional
  The width of the window in seconds (for a predefined window).

**Returns**

- `rate` : `Quantity`
The population rate in Hz, smoothed with the given window. Note that the rates are smoothed and not re-binned, i.e. the length of the returned array is the same as the length of the rate attribute and can be plotted against the PopulationRateMonitor's t attribute.

Tutorials and examples using this

- Example frompapers/Brunel_Hakim_1999
- Example frompapers/Brunel_Wang_2001

spikemonitor module

Exported members: EventMonitor, SpikeMonitor

Classes

| EventMonitor(source, event[, variables, ...]) | Record events from a NeuronGroup or another event source. |

EventMonitor class

(Shortest import: from brian2 import EventMonitor)

class brian2.monitors.spikemonitor.EventMonitor(source, event, variables=None, record=True, when=None, order=None, name='eventmonitor*', codeobj_class=None)

Bases: brian2.groups.group.Group, brian2.groups.group.CodeRunner

Record events from a NeuronGroup or another event source.

The recorded events can be accessed in various ways: the attributes i and t store all the indices and event times, respectively. Alternatively, you can get a dictionary mapping neuron indices to event trains, by calling the event_trains method.

Parameters

source: NeuronGroup, SpikeSource

The source of events to record.

event: str

The name of the event to record

variables: str or sequence of str, optional

Which variables to record at the time of the event (in addition to the index of the neuron).

Can be the name of a variable or a list of names.

record: bool, optional

Whether or not to record each event in i and t (the count will always be recorded). Defaults to True.

when: str, optional

When to record the events, by default records events in the same slot where the event is emitted.
order : int, optional

The priority of this group for operations occurring at the same time step and in the same scheduling slot. Defaults to the order where the event is emitted + 1, i.e. it will be recorded directly afterwards.

name : str, optional

A unique name for the object, otherwise will use source. name+'_eventmonitor_0', etc.

codeobj_class : class, optional

The CodeObject class to run code with.

See also:
SpikeMonitor

Attributes

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>count</td>
<td>The array of event counts (length = size of target group)</td>
</tr>
<tr>
<td>event</td>
<td>The event that we are listening to</td>
</tr>
<tr>
<td>it</td>
<td>Returns the pair (i, t).</td>
</tr>
<tr>
<td>it_</td>
<td>Returns the pair (i, t_).</td>
</tr>
<tr>
<td>num_events</td>
<td>Returns the total number of recorded events.</td>
</tr>
<tr>
<td>record</td>
<td>Whether to record times and indices of events</td>
</tr>
<tr>
<td>record_variables</td>
<td>The additional variables that will be recorded</td>
</tr>
<tr>
<td>source</td>
<td>The source we are recording from</td>
</tr>
</tbody>
</table>

Methods

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>all_values()</td>
<td>Return a dictionary mapping recorded variable names (including t) to a dictionary mapping neuron indices to arrays of variable values at the time of the events (sorted by time).</td>
</tr>
<tr>
<td>event_trains()</td>
<td>Return a dictionary mapping event indices to arrays of event times.</td>
</tr>
<tr>
<td>reinit()</td>
<td>Clears all recorded spikes</td>
</tr>
<tr>
<td>resize(new_size)</td>
<td></td>
</tr>
<tr>
<td>values(var)</td>
<td>Return a dictionary mapping neuron indices to arrays of variable values at the time of the events (sorted by time).</td>
</tr>
</tbody>
</table>

Details

count

The array of event counts (length = size of target group)

event

The event that we are listening to

it

Returns the pair (i, t).
it_
   Returns the pair (i, t_).

num_events
   Returns the total number of recorded events.

record
   Whether to record times and indices of events

record_variables
   The additional variables that will be recorded

source
   The source we are recording from

all_values()
   Return a dictionary mapping recorded variable names (including t) to a dictionary mapping neuron indices to arrays of variable values at the time of the events (sorted by time). This is equivalent to (but more efficient than) calling values for each variable and storing the result in a dictionary.

   Returns all_values : dict

   Dictionary mapping variable names to dictionaries which themselves are mapping neuron indicies to arrays of variable values at the time of the events.

Examples

```python
>>> from brian2 import *
>>> G = NeuronGroup(2, """"counter1 : integer
...     counter2 : integer
...     max_value : integer"""",
...     threshold='counter1 >= max_value',
...     reset='counter1 = 0')
>>> G.run_regularly('counter1 += 1; counter2 += 1')  # doctest: +ELLIPSIS
CodeRunner(...)
>>> G.max_value = [50, 100]
>>> mon = EventMonitor(G, event='spike', variables='counter2')
>>> run(10*ms)
>>> all_values = mon.all_values()
>>> print(all_values['counter2'][0])
[ 50 100]
>>> print(all_values['t'][1])
[ 9.9] ms
```

event_trains()
   Return a dictionary mapping event indices to arrays of event times. Equivalent to calling values('t').

   Returns event_trains : dict

   Dictionary that stores an array with the event times for each neuron index.

See also:

SpikeMonitor.spike_trains

reinit()
   Clears all recorded spikes

resize(new_size)
values (var)
Return a dictionary mapping neuron indices to arrays of variable values at the time of the events (sorted by time).

Parameters

var : str
The name of the variable.

Returns

values : dict
Dictionary mapping each neuron index to an array of variable values at the time of the events

Examples

```python
>>> from brian2 import *
>>> G = NeuronGroup(2, "counter1 : integer
... counter2 : integer
... max_value : integer",
... threshold='counter1 >= max_value',
... reset='counter1 = 0')
>>> G.run_regularly('counter1 += 1; counter2 += 1')  # doctest: +ELLIPSIS
CodeRunner(...)
>>> G.max_value = [50, 100]
>>> mon = EventMonitor(G, event='spike', variables='counter2')
>>> run(10*ms)
>>> counter2_values = mon.values('counter2')
>>> print(counter2_values[0])
[ 50 100]
>>> print(counter2_values[1])
[100]
```

Tutorials and examples using this

- Example advanced/custom_events

SpikeMonitor class

(Shortest import: from brian2 import SpikeMonitor)

class brian2.monitors.spikemonitor.SpikeMonitor(source[, variables, record, ...])
Record spikes from a NeuronGroup or other spike source.

Bases: brian2.monitors.spikemonitor.EventMonitor

Record spikes from a NeuronGroup or other spike source.

The recorded spikes can be accessed in various ways (see Examples below): the attributes i and t store all the indices and spike times, respectively. Alternatively, you can get a dictionary mapping neuron indices to spike trains, by calling the spike_trains method. If you record additional variables with the variables argument, these variables can be accessed by their name (see Examples).
Parameters **source**: *(NeuronGroup, SpikeSource)*

The source of spikes to record.

**variables**: str or sequence of str, optional

Which variables to record at the time of the spike (in addition to the index of the neuron). Can be the name of a variable or a list of names.

**record**: bool, optional

Whether or not to record each spike in \(i\) and \(t\) (the *count* will always be recorded). Defaults to True.

**when**: str, optional

When to record the events, by default records events in the same slot where the event is emitted.

**order**: int, optional

The priority of this group for operations occurring at the same time step and in the same scheduling slot. Defaults to the order where the event is emitted + 1, i.e. it will be recorded directly afterwards.

**name**: str, optional

A unique name for the object, otherwise will use *source*. name+'_spikemonitor_0', etc.

**codeobj_class**: class, optional

The CodeObject class to run code with.

**Examples**

```python
>>> from brian2 import *
>>> spikes = SpikeGeneratorGroup(3, [0, 1, 2], [0, 1, 2]*ms)
>>> spike_mon = SpikeMonitor(spikes)
>>> net = Network(spikes, spike_mon)
>>> net.run(3*ms)
>>> print(spike_mon.i[:])
[0 1 2]
>>> print(spike_mon.t[:])
[ 0. 1. 2.] ms
>>> print(spike_mon.t_[:])
[ 0. 0.001 0.002]
>>> from brian2 import *
>>> G = NeuronGroup(2, """counter1 : integer
counter2 : integer
max_value : integer""",
threshold='counter1 >= max_value',
reset='counter1 = 0')
>>> G.run_regularly('counter1 += 1; counter2 += 1')  # doctest: +ELLIPSIS
CodeRunner(...)
>>> G.max_value = [50, 100]
>>> mon = SpikeMonitor(G, variables='counter2')
>>> net = Network(G, mon)
>>> net.run(10*ms)
>>> print(mon.i[:])
[0 0 1]
```

(continues on next page)
Attributes

<table>
<thead>
<tr>
<th>count</th>
<th>The array of spike counts (length = size of target group)</th>
</tr>
</thead>
<tbody>
<tr>
<td>num_spikes</td>
<td>Returns the total number of recorded spikes.</td>
</tr>
</tbody>
</table>

Methods

| all_values()  | Return a dictionary mapping recorded variable names (including t) to a dictionary mapping neuron indices to arrays of variable values at the time of the spikes (sorted by time). |
| spike_trains()| Return a dictionary mapping spike indices to arrays of spike times. |
| values(var)   | Return a dictionary mapping neuron indices to arrays of variable values at the time of the spikes (sorted by time). |

Details

**count**
The array of spike counts (length = size of target group)

**num_spikes**
Returns the total number of recorded spikes.

**all_values()**
Return a dictionary mapping recorded variable names (including t) to a dictionary mapping neuron indices to arrays of variable values at the time of the spikes (sorted by time). This is equivalent to (but more efficient than) calling `values` for each variable and storing the result in a dictionary.

**Returns** all_values : dict

Dictionary mapping variable names to dictionaries which themselves are mapping neuron indices to arrays of variable values at the time of the spikes.

Examples

```python
>>> from brian2 import *
>>> G = NeuronGroup(2, """"counter1 : integer
... counter2 : integer
... max_value : integer""",
... threshold='counter1 >= max_value',
... reset='counter1 = 0')
>>> G.run_regularly('counter1 += 1; counter2 += 1')  # doctest: +ELLIPSIS
CodeRunner(...)
>>> G.max_value = [50, 100]
```

(continues on next page)
>>> mon = SpikeMonitor(G, variables='counter2')
>>> run(10*ms)
>>> all_values = mon.all_values()
>>> print(all_values['counter2'][0])
[ 50 100]
>>> print(all_values['t'][1])
[ 9.9] ms

spike_trains()  
Return a dictionary mapping spike indices to arrays of spike times.  

Returns spike_trains : dict  
Dictionary that stores an array with the spike times for each neuron index.

Examples

>>> from brian2 import *
>>> spikes = SpikeGeneratorGroup(3, [0, 1, 2], [0, 1, 2]*ms)
>>> spike_mon = SpikeMonitor(spikes)
>>> run(3*ms)
>>> spike_trains = spike_mon.spike_trains()
>>> spike_trains[1]
array([ 1.]) * msecond

values(var)  
Return a dictionary mapping neuron indices to arrays of variable values at the time of the spikes (sorted by time).  

Parameters var : str  
The name of the variable.  

Returns values : dict  
Dictionary mapping each neuron index to an array of variable values at the time of the spikes.

Examples

>>> from brian2 import *
>>> G = NeuronGroup(2, """"counter1 : integer  
... counter2 : integer  
... max_value : integer"""",  
... threshold='counter1 >= max_value',  
... reset='counter1 = 0')
>>> G.run_regularly('counter1 += 1; counter2 += 1')  
# doctest: +ELLIPSIS
CodeRunner(...)  
>>> G.max_value = [50, 100]
>>> mon = SpikeMonitor(G, variables='counter2')
>>> run(10*ms)
>>> counter2_values = mon.values('counter2')
>>> print(counter2_values[0])
[ 50 100]
>>> print(counter2_values[1])
[100]
Tutorials and examples using this

- Tutorial 3-intro-to-brian-simulations
- Tutorial 1-intro-to-brian-neurons
- Example reliability
- Example IF_curve_LIF
- Example phase_locking
- Example non_reliability
- Example IF_curve_Hodgkin_Huxley
- Example CUBA
- Example adaptive_threshold
- Example compartmental/hh_with_spikes
- Example advanced/custom_events
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- Example synapses/jeffress
- Example synapses/STDP
- Example standalone/cuba_openmp
- Example standalone/STDP_standalone
- Example frompapers/Rossant_et_al_2011bis
- Example frompapers/Diesmann_et_al_1999
- Example frompapers/Izhikevich_2007
- Example frompapers/Vogels_et_al_2011
- Example frompapers/Brette_Gerstner_2005
- Example frompapers/Brette_Guigon_2003
- Example frompapers/Brunei_Hakim_1999
- Example frompapers/Touboul_Brette_2008
- Example frompapers/Sturzl_et_al_2000
- Example frompapers/Platkiewicz_Brette_2011
- Example frompapers/Brette_2004
- Example frompapers/Brunei_Wang_2001
- Example frompapers/Brette_2012/Fig5A
- Example frompapers/Stimberg_et_al_2018/example_6_COBA_with_astro
- Example frompapers/Stimberg_et_al_2018/example_1_COBA
statemonitor module

Exported members: StateMonitor

Classes

StateMonitor(source, variables, record[, ...]) Record values of state variables during a run

StateMonitor class

(Shortest import: from brian2 import StateMonitor)

class brian2.monitors.statemonitor.StateMonitor(source, variables, record, dt=None, clock=None, when='start', order=0, name='statemonitor*', codeobj_class=None)

Bases: brian2.groups.group.Group, brian2.groups.group.CodeRunner

Record values of state variables during a run

To extract recorded values after a run, use the t attribute for the array of times at which values were recorded, and variable name attribute for the values. The values will have shape (len(indices), len(t)), where indices are the array indices which were recorded. When indexing the StateMonitor directly, the returned object can be used to get the recorded values for the specified indices, i.e. the indexing semantic refers to the indices in source, not to the relative indices of the recorded values. For example, when recording only neurons with even numbers, mon[[0, 2]].v will return the values for neurons 0 and 2, whereas mon.v[[0, 2]] will return the values for the first and third recorded neurons, i.e. for neurons 0 and 4.

Parameters

source : Group

Which object to record values from.

variables : str, sequence of str, True

Which variables to record, or True to record all variables (note that this may use a great deal of memory).

record : bool, sequence of ints

Which indices to record, nothing is recorded for False, everything is recorded for True (warning: may use a great deal of memory), or a specified subset of indices.

dt : Quantity, optional

The time step to be used for the monitor. Cannot be combined with the clock argument.

clock : Clock, optional

The update clock to be used. If neither a clock, nor the dt argument is specified, the clock of the source() will be used.

when : str, optional

At which point during a time step the values should be recorded. Defaults to 'start'.

order : int, optional

The priority of of this group for operations occurring at the same time step and in the same scheduling slot. Defaults to 0.

name : str, optional
A unique name for the object, otherwise will use source.name+'statemonitor_0', etc.

codeobj_class: CodeObject, optional

The CodeObject class to create.

Notes

Since this monitor by default records in the 'start' time slot, recordings of the membrane potential in integrate-and-fire models may look unexpected: the recorded membrane potential trace will never be above threshold in an integrate-and-fire model, because the reset statement will have been applied already. Set the when keyword to a different value if this is not what you want.

Note that record=True only works in runtime mode for synaptic variables. This is because the actual array of indices has to be calculated and this is not possible in standalone mode, where the synapses have not been created yet at this stage. Consider using an explicit array of indices instead, i.e. something like record=np.arange(n_synapses).

Examples

Record all variables, first 5 indices:

```python
eqs = ""
dV/dt = (2-V)/(10*ms) : 1
""
threshold = 'V>1'
reset = 'V = 0'
G = NeuronGroup(100, eqs, threshold=threshold, reset=reset)
G.V = rand(len(G))
M = StateMonitor(G, True, record=range(5))
run(100*ms)
plot(M.t, M.V.T)
show()
```

Attributes

- `record` The array of recorded indices
- `record_variables` The variables to record

Methods

- `record_single_timestep()` Records a single time step.
- `reinit()`
- `resize(new_size)`

Details

- `record` The array of recorded indices
- `record_variables`
The variables to record

`record_single_timestep()`

Records a single time step. Useful for recording the values at the end of the simulation – otherwise a `StateMonitor` will not record the last simulated values since its `when` attribute defaults to `'start'`, i.e. the last recording is at the beginning of the last time step.

**Notes**

This function will only work if the `StateMonitor` has been already run, but a run with a length of 0·ms does suffice.

**Examples**

```python
>>> from brian2 import *
>>> G = NeuronGroup(1, 'dv/dt = -v/(5*ms) : 1')
>>> G.v = 1
>>> mon = StateMonitor(G, 'v', record=True)
>>> run(0.5*ms)
>>> print(np.array_str(mon.v[:,], precision=3))
[[ 1.  0.98  0.961  0.942  0.923]]
>>> print(mon.t[:,])
[ 0. 100. 200. 300. 400.] us
>>> print(np.array_str(G.v[:,], precision=3))  # last value had not been
˓→recorded
[ 0.905]
>>> mon.record_single_timestep()
>>> print(mon.t[:,])
[ 0. 100. 200. 300. 400. 500.] us
>>> print(np.array_str(mon.v[:,], precision=3))
[[ 1.  0.98  0.961  0.942  0.923  0.905]]
```

`reinit()`

`resize(new_size)`

**Tutorials and examples using this**

- Tutorial 2-intro-to-brian-synapses
- Tutorial 3-intro-to-brian-simulations
- Tutorial 1-intro-to-brian-neurons
- Example phase_locking
- Example COBAHH
- Example adaptive_threshold
- Example compartmental/spike_initiation
- Example compartmental/bipolar_with_inputs
- Example compartmental/bipolar_with_inputs2
- Example compartmental/hh_with_spikes
- Example compartmental/infinite_cable
• Example compartmental/hodgkin_huxley_1952
• Example compartmental/bipolar_cell
• Example compartmental/lfp
• Example advanced/stochastic_odes
• Example advanced/compare_GSL_to_conventional
• Example advanced/custom_events
• Example synapses/gapjunctions
• Example synapses/nonlinear
• Example synapses/synapses
• Example synapses/jeffress
• Example synapses/STDP
• Example standalone/STDP_standalone
• Example frompapers/Rossant_et_al_2011bis
• Example frompapers/Hindmarsh_Rose_1984
• Example frompapers/Clopath_et_al_2010_homeostasis
• Example frompapers/Wang_Buszaki_1996
• Example frompapers/Morris_Lecar_1981
• Example frompapers/Izhikevich_2007
• Example frompapers/Brette_Gersten_2005
• Example frompapers/Brette_Guigon_2003
• Example frompapers/Rothman_Manis_2003
• Example frompapers/Desette_et_al_1998
• Example frompapers/Touboul_Brette_2008
• Example frompapers/Platkiewicz_Brette_2011
• Example frompapers/Brette_2012/Fig3CF
• Example frompapers/Brette_2012/Fig3AB
• Example frompapers/Brette_2012/Fig5A
• Example frompapers/Brette_2012/Fig1
• Example frompapers/Brette_2012/Fig4
• Example frompapers/Stimberg_et_al_2018/example_1_COBA
• Example frompapers/Stimberg_et_al_2018/example_5_astro_ring
• Example frompapers/Stimberg_et_al_2018/example_2_gchi_astrocyte
• Example frompapers/Stimberg_et_al_2018/example_4_synrel
• Example frompapers/Stimberg_et_al_2018/example_4_rsmean
• Example frompapers/Stimberg_et_al_2018/example_3_io_synapse
StateMonitorView class

(State import: from brian2.monitors.statemonitor import StateMonitorView)

class brian2.monitors.statemonitor.StateMonitorView(monitor, item)
    Bases: object

6.6.10 parsing package

bast module

Brian AST representation

This is a standard Python AST representation with additional information added.

Exported members: brian_ast, BrianASTRenderer, dtype_hierachy

Classes

BrianASTRenderer(variables[, copy_variables]) This class is modelled after NodeRenderer - see there for details.

BrianASTRenderer class

(State import: from brian2.parsing.bast import BrianASTRenderer)

class brian2.parsing.bast.BrianASTRenderer(variables, copy_variables=True)
    Bases: object
        This class is modelled after NodeRenderer - see there for details.

Methods

render_BinOp(node)
render_BoolOp(node)
render_Call(node)
render_Compare(node)
render_Constant(node)
render_Name(node)
render_NameConstant(node)
render_Num(node)
render_UnaryOp(node)
render_node(node)

Details

render_BinOp(node)
render_BoolOp(node)
render_Call(node)
render_Compare(node)
render_Constant(node)
render_Name(node)
render_NameConstant(node)
render_Num(node)
render_UnaryOp(node)
render_node(node)

Functions

\texttt{brian\_ast(expr, variables)} \quad \text{Returns an AST tree representation with additional information}

\textbf{brian\_ast function}

(\textit{Shortest import:} from brian2.parsing.bast import brian\_ast)

\begin{verbatim}

brian2.parsing.bast.brian_ast(expr, variables)

Returns an AST tree representation with additional information

Each node will be a standard Python \texttt{ast} node with the following additional attributes:

\texttt{dtype} One of 'boolean', 'integer' or 'float', referring to the data type of the value of this node.

\texttt{scalar} Either \texttt{True} or \texttt{False} if the node uses any vector-valued variables.

\texttt{complexity} An integer representation of the computational complexity of the node. This is a very rough representation used for things like $2 \times (x+y)$ is less complex than $2 \times x + 2 \times y$ and $\exp(x)$ is more complex than $2 \times x$ but shouldn’t be relied on for fine distinctions between expressions.

\textbf{Parameters} \texttt{expr} : str

The expression to convert into an AST representation

\texttt{variables} : dict

The dictionary of \texttt{Variable} objects used in the expression.

\end{verbatim}

\texttt{brian\_dtype\_from\_dtype(dtype)} \quad \text{Returns ‘boolean’, ‘integer’ or ‘float’}

\textbf{brian\_dtype\_from\_dtype function}

(\textit{Shortest import:} from brian2.parsing.bast import brian\_dtype\_from\_dtype)

\begin{verbatim}

brian2.parsing.bast.brian\_dtype\_from\_dtype(dtype)

Returns ‘boolean’, ‘integer’ or ‘float’

\end{verbatim}

\texttt{brian\_dtype\_from\_value(value)} \quad \text{Returns ‘boolean’, ‘integer’ or ‘float’}
brian_dtype_from_value function

(Shortest import: from brian2.parsing.bast import brian_dtype_from_value)

brian2.parsing.bast.brian_dtype_from_value(value)

Returns ‘boolean’, ‘integer’ or ‘float’

is_boolean(value)

is_boolean function

(Shortest import: from brian2.parsing.bast import is_boolean)

brian2.parsing.bast.is_boolean(value)

is_boolean_dtype(obj)

is_boolean_dtype function

(Shortest import: from brian2.parsing.bast import is_boolean_dtype)

brian2.parsing.bast.is_boolean_dtype(obj)

is_float(value)

is_float function

(Shortest import: from brian2.parsing.bast import is_float)

brian2.parsing.bast.is_float(value)

is_float_dtype(obj)

is_float_dtype function

(Shortest import: from brian2.parsing.bast import is_float_dtype)

brian2.parsing.bast.is_float_dtype(obj)

is_integer(value)

is_integer function

(Shortest import: from brian2.parsing.bast import is_integer)

brian2.parsing.bast.is_integer(value)
is_integer_dtype(obj)

is_integer_dtype function

(Shortest import: from brian2.parsing.bast import is_integer_dtype)

brian2.parsing.bast.is_integer_dtype(obj)

dependencies module

Exported members: abstract_code_dependencies

Functions

abstract_code_dependencies(code[, . . .]) Analyses identifiers used in abstract code blocks

abstract_code_dependencies function

(Shortest import: from brian2.parsing.dependencies import abstract_code_dependencies)

brian2.parsing.dependencies.abstract_code_dependencies(code, known_vars=None, known_funcs=None)

Analyses identifiers used in abstract code blocks

Parameters code : str

The abstract code block.

known_vars : set

The set of known variable names.

known_funcs : set

The set of known function names.

Returns results : namedtuple with the following fields

all The set of all identifiers that appear in this code block, including functions.
read The set of values that are read, excluding functions.
write The set of all values that are written to.
funcs The set of all function names.
known_all The set of all identifiers that appear in this code block and are known.
known_read The set of known values that are read, excluding functions.
known_write The set of known values that are written to.
knownfuncs The set of known functions that are used.
unknown_read The set of all unknown variables whose values are read. Equal to read-known_vars.
unknown_write The set of all unknown variables written to. Equal to write-known_vars.
unknown_funcs The set of all unknown function names, equal to \( \text{funcs-known_funcs} \).

undefined_read The set of all unknown variables whose values are read before they are written to. If this set is nonempty it usually indicates an error, since a variable that is read should either have been defined in the code block (in which case it will appear in newly_defined) or already be known.

newly_defined The set of all variable names which are newly defined in this abstract code block.

\[
\text{get_read_write_funcs}(\text{parsed\_code})
\]

get_read_write_funcs function

(Shortest import: from brian2.parsing.dependencies import get_read_write_funcs)

brian2.parsing.dependencies.get_read_write_funcs(parsed_code)

expressions module

Exported members: parse_expression_dimensions

Functions

\[
\text{is_boolean_expression}(\text{expr}, \text{variables})\quad \text{Determines if an expression is of boolean type or not}
\]

is_boolean_expression function

(Shortest import: from brian2.parsing.expressions import is_boolean_expression)

brian2.parsing.expressions.is_boolean_expression(expr, variables)

Determines if an expression is of boolean type or not

Parameters

expr : str

The expression to test

variables : dict-like of Variable

The variables used in the expression.

Returns

isbool : bool

Whether or not the expression is boolean.

Raises

SyntaxError If the expression ought to be boolean but is not, for example \( x<y \) and \( z \) where \( z \) is not a boolean variable.

Notes

We test the following cases recursively on the abstract syntax tree:
• The node is a boolean operation. If all the subnodes are boolean expressions we return `True`, otherwise we raise the `SyntaxError`.

• The node is a function call, we return `True` or `False` depending on whether the function description has the `_returns_bool` attribute set.

• The node is a variable name, we return `True` or `False` depending on whether `is_boolean` attribute is set or if the name is `True` or `False`.

• The node is a comparison, we return `True`.

• The node is a unary operation, we return `True` if the operation is `not`, otherwise `False`.

• Otherwise we return `False`.

---

**parse_expression_dimensions**

(Shortest import: `from brian2.parsing.expressions import parse_expression_dimensions`

brian2.parsing.expressions.**parse_expression_dimensions** *(expr, variables)*

Returns the unit value of an expression, and checks its validity

**Parameters**

- **expr**: str
  
The expression to check.

- **variables**: dict
  
Dictionary of all variables used in the `expr` (including `Constant` objects for external variables)

**Returns**

- **unit**: Quantity
  
The output unit of the expression

**Raises**

- **SyntaxError** If the expression cannot be parsed, or if it uses `a ** b` for `b` anything other than a constant number.

- **DimensionMismatchError** If any part of the expression is dimensionally inconsistent.

---

**functions module**

*Exported members:* `AbstractCodeFunction`, `abstract_code_from_function`, `extract_abstract_code_functions`, `substitute_abstract_code_functions`

**Classes**

- `AbstractCodeFunction` *(name, args, code,...)*
  
The information defining an abstract code function
AbstractCodeFunction class

(Shortest import: from brian2.parsing.functions import AbstractCodeFunction)

class brian2.parsing.functions.AbstractCodeFunction(name, args, code, return_expr)
    Bases: object

The information defining an abstract code function
Has attributes corresponding to initialisation parameters

Parameters

name : str
    The function name.

args : list of str
    The arguments to the function.

code : str
    The abstract code string consisting of the body of the function less the return statement.

return_expr : str or None
    The expression returned, or None if there is nothing returned.

FunctionRewriter class

(Shortest import: from brian2.parsing.functions import FunctionRewriter)

class brian2.parsing.functions.FunctionRewriter(func[, numcalls=0])
    Bases: ast.NodeTransformer

Inlines a function call using temporary variables
numcalls is the number of times the function rewriter has been called so far, this is used to make sure that when
recursively inlining there is no name aliasing. The substitute_abstract_code_functions ensures that this is kept
up to date between recursive runs.

The pre attribute is the set of lines to be inserted above the currently being processed line, i.e. the inline code.

The visit method returns the current line processed so that the function call is replaced with the output of the
inlining.

Methods

visit_Call(node)

Details

visit_Call (node)

VarRewriter(pre)
    Rewrites all variable names in names by prepending pre
VarRewriter class

(Shortest import: from brian2.parsing.functions import VarRewriter)

```python
class brian2.parsing.functions.VarRewriter(pre):
    Bases: ast.NodeTransformer
    Rewrites all variable names in names by prepending pre
```

Methods

- `visit_Call(node)`
- `visit_Name(node)`

Details

- `visit_Call(node)`
- `visit_Name(node)`

Functions

- `abstract_code_from_function(func)` Converts the body of the function to abstract code

abstract_code_from_function function

(Shortest import: from brian2.parsing.functions import abstract_code_from_function)

```python
brian2.parsing.functions.abstract_code_from_function(func)
```

- Converts the body of the function to abstract code

  Parameters  
  `func` : function, str or ast.FunctionDef

  The function object to convert. Note that the arguments to the function are ignored.

  Returns  
  `func` : AbstractCodeFunction

  The corresponding abstract code function

Raises

- `SyntaxError` If unsupported features are used such as if statements or indexing.

- `extract_abstract_code_functions(code)` Returns a set of abstract code functions from function definitions.

extract_abstract_code_functions function

(Shortest import: from brian2.parsing.functions import extract_abstract_code_functions)

```python
brian2.parsing.functions.extract_abstract_code_functions(code)
```
Returns a set of abstract code functions from function definitions. Returns all functions defined at the top level and ignores any other code in the string.

**Parameters**
- `code`: str
  
  The code string defining some functions.

**Returns**
- `funcs`: dict

A mapping \((\text{name}, \func)\) for \(\func\) an AbstractCodeFunction.

### `substitute_abstract_code_functions` function

(Shortest import: `from brian2.parsing.functions import substitute_abstract_code_functions`)

brian2.parsing.functions.\texttt{substitute\_abstract\_code\_functions}(\texttt{code}, \texttt{funcs})

Performs inline substitution of all the functions in the code

**Parameters**
- `code`: str

  The abstract code to make inline substitutions into.

- `funcs`: list, dict or set of AbstractCodeFunction

  The function substitutions to use, note in the case of a dict, the keys are ignored and the function name is used.

**Returns**
- `code`: str

  The code with inline substitutions performed.

### rendering module

**Exported members:** NodeRenderer, NumpyNodeRenderer, CPPNodeRenderer, SympyNodeRenderer, get_node_value

**Classes**

CPPNodeRenderer([auto_vectorise])

**Methods**

### CPPNodeRenderer class

(Shortest import: `from brian2.parsing.rendering import CPPNodeRenderer`)

class brian2.parsing.rendering.\texttt{CPPNodeRenderer}(\texttt{auto\_vectorise=\texttt{None}})

**Bases:** brian2.parsing.rendering.NodeRenderer
Methods

rendert_{Assign\_name}(node)
rendert_{BinOp\_name}(node)
rendert_{Name\_name}(node)
rendert_{NameConstant\_name}(node)

Details

rendert_{Assign\_name}(node)
rendert_{BinOp\_name}(node)
rendert_{Name\_name}(node)
rendert_{NameConstant\_name}(node)

NodeRenderer() (auto_vectorise)

Methods

NodeRenderer class

(Shortest import: from brian2.parsing.rendering import NodeRenderer)

class brian2.parsing.rendering.\texttt{NodeRenderer}(auto_vectorise=\texttt{None})
  Bases: \texttt{object}

Methods

rendert_{Assign\_name}(node)
rendert_{AugAssign\_name}(node)
rendert_{BinOp\_name}(node)
rendert_{BinOp\_parentheses}(left, right, op)
rendert_{BoolOp\_name}(node)
rendert_{Call\_name}(node)
rendert_{Compare\_name}(node)
rendert_{Constant\_name}(node)
rendert_{Name\_name}(node)
rendert_{NameConstant\_name}(node)
rendert_{Num\_name}(node)
rendert_{UnaryOp\_name}(node)
rendert_{code\_name}(code)
rendert_{element\_parentheses\_name}(node) Render an element with parentheses around it or leave them away for numbers, names and function calls.
rendert_{expr\_name}(expr[, strip])
rendert_{func\_name}(node)
rendert_{node\_name}(node)
Details

render_Assign (node)
render_AugAssign (node)
render_BinOp (node)
render_BinOp_parentheses (left, right, op)
render_BoolOp (node)
render_Call (node)
render_Compare (node)
render_Constant (node)
render_Name (node)
render_NameConstant (node)
render_Num (node)
render_UnaryOp (node)
render_code (code)
render_element_parentheses (node)
    Render an element with parentheses around it or leave them away for numbers, names and function calls.
render_expr (expr, strip=True)
render_func (node)
render_node (node)

NumpyNodeRenderer([auto_vectorise])

Methods

NumpyNodeRenderer class

(Shortest import: from brian2.parsing.rendering import NumpyNodeRenderer)
class brian2.parsing.rendering.NumpyNodeRenderer (auto_vectorise=None)
    Bases: brian2.parsing.rendering.NodeRenderer

Methods

render_UnaryOp (node)

Details

render_UnaryOp (node)
**SympyNodeRenderer**([auto_vectorise])

**Methods**

**SympyNodeRenderer class**

*(Shortest import: from brian2.parsing.rendering import SympyNodeRenderer)*

```python
class brian2.parsing.rendering.SympyNodeRenderer(auto_vectorise=None)

Bases: brian2.parsing.rendering.NodeRenderer
```

**Methods**

- `render_BinOp(node)`
- `render_BoolOp(node)`
- `render_Call(node)`
- `render_Compare(node)`
- `render_Name(node)`
- `render_NameConstant(node)`
- `render_Num(node)`
- `render_UnaryOp(node)`
- `render_func(node)`

**Details**

- `render_BinOp(node)`
- `render_BoolOp(node)`
- `render_Call(node)`
- `render_Compare(node)`
- `render_Name(node)`
- `render_NameConstant(node)`
- `render_Num(node)`
- `render_UnaryOp(node)`
- `render_func(node)`

**Functions**

- `get_node_value(node)`

  Helper function to mask differences between Python versions

---

**get_node_value function**

*(Shortest import: from brian2.parsing.rendering import get_node_value)*
brian2.parsing.rendering.get_node_value(node)
Helper function to mask differences between Python versions

**statements module**

Functions

<table>
<thead>
<tr>
<th><strong>parse_statement</strong></th>
<th>(code)</th>
<th>Parses a single line of code into “var op expr”.</th>
</tr>
</thead>
</table>

**parse_statement function**

*(Shortest import: from brian2.parsing.statements import parse_statement)*

brian2.parsing.statements.parse_statement(code)
Parses a single line of code into “var op expr”.

**Parameters**

code : str
A string containing a single statement of the form var op expr # comment, where the # comment part is optional.

**Returns**

var, op, expr, comment : str, str, str, str
The four parts of the statement.

**Examples**

```python
>>> parse_statement('v = -65*mV # reset the membrane potential')
('v', '=', '-65*mV', 'reset the membrane potential')
>>> parse_statement('v += dt*(-v/tau)')
('v', '+=', 'dt*(-v/tau)', '')
```

**sympytools module**

Utility functions for parsing expressions and statements.

**Classes**

<table>
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<tr>
<th><strong>CustomSympyPrinter</strong></th>
<th>([settings])</th>
<th>Printer that overrides the printing of some basic sympy objects.</th>
</tr>
</thead>
</table>

**CustomSympyPrinter class**

*(Shortest import: from brian2.parsing.sympytools import CustomSympyPrinter)*

class brian2.parsing.sympytools.CustomSympyPrinter (settings=None)

**Bases**: sympy.printing.str.StrPrinter

Printer that overrides the printing of some basic sympy objects. E.g. print “a and b” instead of “And(a, b)”.

**Functions**
check_expression_for_multiple_stateful_functions function

(Shortest import: from brian2.parsing.sympytools import check_expression_for_multiple_stateful_functions)

brian2.parsing.sympytools.check_expression_for_multiple_stateful_functions(expr, variables)

expression_complexity(expr[, complexity]) Returns the complexity of an expression (either string or sympy)

expression_complexity function

(Shortest import: from brian2.parsing.sympytools import expression_complexity)

brian2.parsing.sympytools.expression_complexity(expr, complexity=None)

Returns the complexity of an expression (either string or sympy)

The complexity is defined as 1 for each arithmetic operation except divide which is 2, and all other operations are 20. This can be overridden using the complexity argument.

Note: calling this on a statement rather than an expression is likely to lead to errors.

Parameters expr: ‘sympy.Expr‘ or str :

The expression.

complexity: None or dict (optional) :

A dictionary mapping expression names to their complexity, to overwrite default behaviour.

Returns complexity: int :

The complexity of the expression.

str_to_sympy(expr[, variables]) Parses a string into a sympy expression.

str_to_sympy function

(Shortest import: from brian2.parsing.sympytools import str_to_sympy)

brian2.parsing.sympytools.str_to_sympy(expr, variables=None)

Parses a string into a sympy expression. There are two reasons for not using sympify directly: 1) sympify does a from sympy import *, adding all functions to its namespace. This leads to issues when trying to use sympy function names as variable names. For example, both beta and factor – quite reasonable names for variables – are sympy functions, using them as variables would lead to a parsing error. 2) We want to use a common syntax across expressions and statements, e.g. we want to allow to use and (instead of &) and function names like ceil (instead of ceiling).

Parameters expr : str

The string expression to parse.
variables: dict, optional

Dictionary mapping variable/function names in the expr to their respective Variable/Function objects.

Returns s_expr:

A sympy expression

Raises

SyntaxError In case of any problems during parsing.

sympy_to_str function

(Shortest import: from brian2.parsing.sympytools import sympy_to_str)

brian2.parsing.sympytools.sympy_to_str(sympy_expr)

Converts a sympy expression into a string. This could be as easy as str(sympy_expr) but it is possible
that the sympy expression contains functions like Abs (for example, if an expression such as sqrt(x**2)
appeared somewhere). We do want to re-translate Abs into abs in this case.

Parameters sympy_expr: sympy.core.expr.Expr

The expression that should be converted to a string.

Returns:

str_expr: str

A string representing the sympy expression.

Objects

PRINTER Printer that overrides the printing of some basic sympy objects.

PRINTER object

(Shortest import: from brian2.parsing.sympytools import PRINTER)

brian2.parsing.sympytools.PRINTER = <brian2.parsing.sympytools.CustomSympyPrinter object>

Printer that overrides the printing of some basic sympy objects. E.g. print “a and b” instead of “And(a, b)”.

6.6.11 random package

6.6.12 spatialneuron package

Exported members: Morphology, Soma, Cylinder, Section, SpatialNeuron
**morphology module**

**Exported members:** *Morphology, Section, Cylinder, Soma*

**Classes**

| Children(owner) | Helper class to represent the children (sub trees) of a section. |

**Children class**

*(Shortest import: from brian2.spatialneuron.morphology import Children)*

**class** brian2.spatialneuron.morphology.**Children**(owner)

**Bases:** object

Helper class to represent the children (sub trees) of a section. Can be used like a dictionary (mapping names to *Morphology* objects), but iterates over the values (sub trees) instead of over the keys (names).

**Methods**

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>add</strong>(name, subtree[, automatic_name])</td>
<td>Add a new child to the morphology.</td>
</tr>
<tr>
<td><strong>name</strong>(child)</td>
<td>Return the given name (i.e. not the automatic name such as 1) for a child subtree.</td>
</tr>
<tr>
<td><strong>remove</strong>(name)</td>
<td>Remove a subtree from this morphology.</td>
</tr>
</tbody>
</table>

**Details**

**add**(name, subtree, automatic_name=False)

Add a new child to the morphology.

**Parameters**

- **name**: str
  
  The name (e.g. "axon", "soma") to use for this sub tree.

- **subtree**: *Morphology*
  
  The sub tree to link as a child.

- **automatic_name**: bool, optional
  
  Whether to chose a new name automatically, if a subtree of the same name already exists (uses e.g. "dend2" instead "dend"). Defaults to False and will raise an error instead.

**name**(child)

Return the given name (i.e. not the automatic name such as 1) for a child subtree.

**Parameters**

- **child**: *Morphology*
  
  The given name for the child.

**remove**(name)

Remove a subtree from this morphology.

**Parameters**

- **name**: str
The name of the sub tree to remove.

\[
\text{Cylinder(**kwds)} \quad \text{A cylindrical section.}
\]

**Cylinder class**

(*Shortest import: from brian2 import Cylinder*)

```python
class brian2.spatialneuron.morphology.Cylinder(**kwds):
    Bases: brian2.spatialneuron.morphology.Section
```

A cylindrical section. For sections with more complex geometry (varying length and/or diameter of each compartment), use the `Section` class.

**Parameters**
- **diameter**: `Quantity`  
  The diameter of the cylinder.
- **n**: `int`, optional  
  The number of compartments in this section. Defaults to 1.
- **length**: `Quantity`, optional  
  The length of the cylinder. Cannot be combined with the specification of coordinates.
- **x**: `Quantity`, optional  
  A sequence of two values, the start and the end point of the cylinder. The coordinates are interpreted as relative to the end point of the parent compartment (if any), so in most cases the start point should be \( 0 \times \text{um} \). The common exception is a cylinder connecting to a `Soma`, here the start point can be used to make the cylinder start at the surface of the sphere instead of at its center. You can specify all of \( x, y, \) or \( z \) to specify a morphology in 3D, or only one or two out of them to specify a morphology in 1D or 2D.
- **y**: `Quantity`, optional  
  See \( x \)
- **z**: `Quantity`, optional  
  See \( x \)
- **type**: `str`, optional  
  The type (e.g. "axon") of this `Cylinder`.

**Attributes**

- **area**: The membrane surface area of each compartment in this section.
- **diameter**: The diameter at the middle of each compartment in this section.
- **end_diameter**: The diameter at the end of each compartment in this section.
- **r_length_1**: The geometry-dependent term to calculate the conductance between the start and the midpoint of each compartment.

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Table 382 – continued from previous page

<table>
<thead>
<tr>
<th>Variable</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$r_{length_2}$</td>
<td>The geometry-dependent term to calculate the conductance between the midpoint and the end of each compartment.</td>
</tr>
<tr>
<td>start_diameter</td>
<td>The diameter at the start of each compartment in this section.</td>
</tr>
<tr>
<td>volume</td>
<td>The volume of each compartment in this section.</td>
</tr>
</tbody>
</table>

Methods

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>copy_section()</td>
<td>Create a copy of the current section (attributes of this section only, not re-creating the parent/children relation)</td>
</tr>
</tbody>
</table>

Details

**area**

The membrane surface area of each compartment in this section. The surface area of each compartment is calculated as $\pi dl$, where $l$ is the length of the compartment, and $d$ is its diameter. Note that this surface area does not contain the area of the two disks at the two sides of the cylinder.

**diameter**

The diameter at the middle of each compartment in this section.

**end_diameter**

The diameter at the end of each compartment in this section.

**$r_{length_1}$**

The geometry-dependent term to calculate the conductance between the start and the midpoint of each compartment. Dividing this value by the Intracellular resistivity gives the conductance.

**$r_{length_2}$**

The geometry-dependent term to calculate the conductance between the midpoint and the end of each compartment. Dividing this value by the Intracellular resistivity gives the conductance.

**start_diameter**

The diameter at the start of each compartment in this section.

**volume**

The volume of each compartment in this section. The volume of each compartment is calculated as $\pi d^2 l$, where $l$ is the length of the compartment, and $d$ is its diameter.

**copy_section()**

Create a copy of the current section (attributes of this section only, not re-creating the parent/children relation)

Returns copy: Morphology

A copy of this section (without the links to the parent/children)

Tutorials and examples using this

- Example compartmental/spike_initiation
- Example compartmental/morphotest
- Example compartmental/bipolar_with_inputs

6.6. Subpackages
Morphology (**kwds)

Neuronal morphology (tree structure).

**Morphology class**

(*Shortest import: from brian2 import Morphology*)

```python
class brian2.spatialneuron.morphology.Morphology(**kwds)
    Bases: object
```

Neuronal morphology (tree structure).

The data structure is a tree where each node is an un-branched section consisting of a number of connected compartments, each one defined by its geometrical properties (length, area, diameter, position).

**Notes**

You cannot create objects of this class, create a Soma, a Section, or a Cylinder instead.

**Attributes**

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>area</td>
<td>The membrane surface area of each compartment in this section.</td>
</tr>
<tr>
<td>children</td>
<td>The children (as a Children object) of this section.</td>
</tr>
<tr>
<td>coordinates</td>
<td>Array with all coordinates at the start- and end-points of each compartment in this section.</td>
</tr>
<tr>
<td>coordinates_</td>
<td>Array with all coordinates (as unitless floating point numbers) at the start- and end-points of each compartment in this section.</td>
</tr>
<tr>
<td>diameter</td>
<td>The diameter at the middle of each compartment in this section.</td>
</tr>
<tr>
<td>Variable</td>
<td>Description</td>
</tr>
<tr>
<td>---------------</td>
<td>------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>distance</td>
<td>The total distance between the midpoint of each compartment and the root of the morphology.</td>
</tr>
<tr>
<td>end_distance</td>
<td>The distance to the root of the morphology at the end of this section.</td>
</tr>
<tr>
<td>end_x</td>
<td>The x coordinate at the end of each compartment.</td>
</tr>
<tr>
<td>end_x_</td>
<td>The x coordinate (as a unitless floating point number) at the end of each compartment.</td>
</tr>
<tr>
<td>end_y</td>
<td>The y coordinate at the end of each compartment.</td>
</tr>
<tr>
<td>end_y_</td>
<td>The y coordinate (as a unitless floating point number) at the end of each compartment.</td>
</tr>
<tr>
<td>end_z</td>
<td>The z coordinate at the end of each compartment.</td>
</tr>
<tr>
<td>end_z_</td>
<td>The z coordinate (as a unitless floating point number) at the end of each compartment.</td>
</tr>
<tr>
<td>length</td>
<td>The length of each compartment in this section.</td>
</tr>
<tr>
<td>n</td>
<td>The number of compartments in this section.</td>
</tr>
<tr>
<td>parent</td>
<td>The parent section of this section.</td>
</tr>
<tr>
<td>r_length_1</td>
<td>The geometry-dependent term to calculate the conductance between the start and the midpoint of each compartment.</td>
</tr>
<tr>
<td>r_length_2</td>
<td>The geometry-dependent term to calculate the conductance between the midpoint and the end of each compartment.</td>
</tr>
<tr>
<td>start_x</td>
<td>The x coordinate at the beginning of each compartment.</td>
</tr>
<tr>
<td>start_x_</td>
<td>The x coordinate (as a unitless floating point number) at the beginning of each compartment.</td>
</tr>
<tr>
<td>start_y</td>
<td>The y coordinate at the beginning of each compartment.</td>
</tr>
<tr>
<td>start_y_</td>
<td>The y coordinate (as a unitless floating point number) at the beginning of each compartment.</td>
</tr>
<tr>
<td>start_z</td>
<td>The z coordinate at the beginning of each compartment.</td>
</tr>
<tr>
<td>start_z_</td>
<td>The z coordinate (as a unitless floating point number) at the beginning of each compartment.</td>
</tr>
<tr>
<td>total_compartments</td>
<td>The total number of compartments in this subtree (i.e. total sections)</td>
</tr>
<tr>
<td>total_sections</td>
<td>The total number of sections in this subtree.</td>
</tr>
<tr>
<td>volume</td>
<td>The volume of each compartment in this section.</td>
</tr>
<tr>
<td>x</td>
<td>The x coordinate at the midpoint of each compartment.</td>
</tr>
<tr>
<td>x_</td>
<td>The x coordinate (as a unitless floating point number) at the midpoint of each compartment.</td>
</tr>
<tr>
<td>y</td>
<td>The y coordinate at the midpoint of each compartment.</td>
</tr>
<tr>
<td>y_</td>
<td>The y coordinate (as a unitless floating point number) at the midpoint of each compartment.</td>
</tr>
<tr>
<td>z</td>
<td>The y coordinate at the midpoint of each compartment.</td>
</tr>
<tr>
<td>z_</td>
<td>The z coordinate (as a unitless floating point number) at the midpoint of each compartment.</td>
</tr>
</tbody>
</table>
### Methods

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>copy_section()</code></td>
<td>Create a copy of the current section (attributes of this section only, not re-creating the parent/children relation)</td>
</tr>
<tr>
<td><code>from_file(filename[, spherical_soma])</code></td>
<td>Convencience method to load a morphology from a given file.</td>
</tr>
<tr>
<td><code>from_points(points[, spherical_soma])</code></td>
<td>Create a morphology from a sequence of points (similar to the SWC format, see Morphology. from_swc_file).</td>
</tr>
<tr>
<td><code>from_swc_file(filename[, spherical_soma])</code></td>
<td>Load a morphology from a SWC file.</td>
</tr>
<tr>
<td><code>generate_coordinates([section_randomness, ...])</code></td>
<td>Create a new Morphology, with coordinates filled in place where the previous morphology did not have any.</td>
</tr>
<tr>
<td><code>topology()</code></td>
<td>Return a representation of the topology</td>
</tr>
</tbody>
</table>

### Details

**area**
The membrane surface area of each compartment in this section.

**children**
The children (as a `Children` object) of this section.

**coordinates**
Array with all coordinates at the start- and end-points of each compartment in this section. The array has size $(n + 1) \times 3$, where $n$ is the number of compartments in this section. Each row is one point (start point of first compartment, end point of first compartment, end point of second compartment, …), with the columns being the x, y, and z coordinates. Returns `None` for morphologies without coordinates.

**coordinates_**
Array with all coordinates (as unitless floating point numbers) at the start- and end-points of each compartment in this section. The array has size $(n + 1) \times 3$, where $n$ is the number of compartments in this section. Each row is one point (start point of first compartment, end point of first compartment, end point of second compartment, …), with the columns being the x, y, and z coordinates. Returns `None` for morphologies without coordinates.

**diameter**
The diameter at the middle of each compartment in this section.

**distance**
The total distance between the midpoint of each compartment and the root of the morphology.

**end_distance**
The distance to the root of the morphology at the end of this section.

**end_x**
The x coordinate at the end of each compartment. Returns `None` for morphologies without coordinates.

**end_x_**
The x coordinate (as a unitless floating point number) at the end of each compartment. Returns `None` for morphologies without coordinates.

**end_y**
The y coordinate at the end of each compartment. Returns `None` for morphologies without coordinates.
end_y
The y coordinate (as a unitless floating point number) at the end of each compartment. Returns None for morphologies without coordinates.

d_end_z
The z coordinate at the end of each compartment. Returns None for morphologies without coordinates.

d_end_z
The z coordinate (as a unitless floating point number) at the end of each compartment. Returns None for morphologies without coordinates.

length
The length of each compartment in this section.

n
The number of compartments in this section.

parent
The parent section of this section.

r_length_1
The geometry-dependent term to calculate the conductance between the start and the midpoint of each compartment. Dividing this value by the Intracellular resistivity gives the conductance.

r_length_2
The geometry-dependent term to calculate the conductance between the midpoint and the end of each compartment. Dividing this value by the Intracellular resistivity gives the conductance.

start_x
The x coordinate at the beginning of each compartment. Returns None for morphologies without coordinates.

start_x
The x coordinate (as a unitless floating point number) at the beginning of each compartment. Returns None for morphologies without coordinates.

start_y
The y coordinate at the beginning of each compartment. Returns None for morphologies without coordinates.

start_y
The y coordinate (as a unitless floating point number) at the beginning of each compartment. Returns None for morphologies without coordinates.

start_z
The z coordinate at the beginning of each compartment. Returns None for morphologies without coordinates.

start_z
The z coordinate (as a unitless floating point number) at the beginning of each compartment. Returns None for morphologies without coordinates.

total_compartments
The total number of compartments in this subtree (i.e. the number of compartments in this section plus all the compartments in the sections deeper in the tree).

total_sections
The total number of sections in this subtree.

volume
The total number of sections in this subtree.
The x coordinate at the midpoint of each compartment. Returns None for morphologies without coordinates.

The x coordinate (as a unitless floating point number) at the midpoint of each compartment. Returns None for morphologies without coordinates.

The y coordinate at the midpoint of each compartment. Returns None for morphologies without coordinates.

The y coordinate (as a unitless floating point number) at the midpoint of each compartment. Returns None for morphologies without coordinates.

The z coordinate at the midpoint of each compartment. Returns None for morphologies without coordinates.

The z coordinate (as a unitless floating point number) at the midpoint of each compartment. Returns None for morphologies without coordinates.

Create a copy of the current section (attributes of this section only, not re-creating the parent/children relation)

Returns copy: Morphology

A copy of this section (without the links to the parent/children)

Convencience method to load a morphology from a given file. At the moment, only SWC files are supported, calling this function is therefore equivalent to calling Morphology.from_swc_file directly.

Parameters filename : str

The name of a file storing a morphology.

spherical_soma : bool, optional

Whether to model the soma as a sphere.

Returns :

morphology: Morphology

The morphology stored in the given file.

Create a morphology from a sequence of points (similar to the SWC format, see Morphology.from_swc_file). Each point has to be a 7-tuple: (index, name, x, y, z, diameter, parent)

Note that the values should not use units, but are instead all taken to be in micrometers.

Parameters points : sequence of 7-tuples

The points of the morphology.

spherical_soma : bool, optional
Whether to model a soma as a sphere.

Returns:

- morphology: Morphology

Notes

This format closely follows the SWC format (see `Morphology.from_swc_file`) with two differences: the `type` should be a string (e.g. 'soma') instead of an integer and the 6-th element should be the diameter and not the radius.

```python
static from_swc_file (filename, spherical_soma=True)
```

Load a morphology from a SWC file. A large database of morphologies in this format can be found at http://neuromorpho.org

The format consists of an optional header of lines starting with # (ignored), followed by a sequence of points, each described in a line following the format:

```
index type x y z radius parent
```

- `index` is an integer label (starting at 1) that identifies the current point and increases by one each line.
- `type` is an integer representing the type of the neural segment. The only type that changes the interpretation by Brian is the type 1 which signals a soma. Types 2 (axon), 3 (dendrite), and 4 (apical dendrite) are used to give corresponding names to the respective sections. All other types are ignored.
- `x`, `y`, and `z` are the cartesian coordinates at each point and `r` is its radius. `parent` refers to the index of the parent point or is -1 for the root point.

Parameters:

- `filename`: str
  - The name of the SWC file.
- `spherical_soma`: bool, optional
  - Whether to model the soma as a sphere.

Returns:

- `morpho`: Morphology
  - The morphology stored in the given file.

```python
generate_coordinates (section_randomness=0.0, compartment_randomness=0.0, overwrite_existing=False)
```

Create a new `Morphology`, with coordinates filled in place where the previous morphology did not have any. This is mostly useful for plotting a morphology, it does not affect its electrical properties.

Parameters:

- `section_randomness`: float, optional
  - The randomness when deciding the direction vector for each new section. The given number is the $\beta$ parameter of an exponential distribution (in degrees) which will be used to determine the deviation from the direction of the parent section. If the given value equals 0 (the default), then a deterministic algorithm will be used instead.
- `compartment_randomness`: float, optional
  - The randomness when deciding the direction vector for each compartment within a section. The given number is the $\beta$ parameter of an exponential distribution (in degrees) which will be used to determine the deviation from the main direction of the current section. If the given value equals 0 (the default), then all compartments will be along a straight line.
overwrite_existing : bool, optional

Whether to overwrite existing coordinates in the morphology. This is by default set to False, meaning that only sections that do not currently have any coordinates set will get new coordinates. This allows to conveniently generate a morphology that can be plotted for a morphology that is based on points but also has artificially added sections (the most common case: an axon added to a reconstructed morphology). If set to True, all sections will get new coordinates. This can be useful to either get a schematic representation of the morphology (with section_randomness and compartment_randomness both 0) or to simply generate a new random variation of a morphology (which will still be electrically equivalent, of course).

Returns morpho_with_coordinates : Morphology

The same morphology, but with coordinates

topology()

Return a representation of the topology

Returns topology : Topology

An object representing the topology (can be converted to a string by using str(...) or simply by printing it with print.)

Tutorials and examples using this

• Example compartmental/spike_initiation
• Example compartmental/morphotest
• Example compartmental/bipolar_with_inputs
• Example compartmental/bipolar_with_inputs2
• Example compartmental/infinite_cable
• Example compartmental/cylinder
• Example compartmental/bipolar_cell
• Example compartmental/rall
• Example frompapers/Brette_2012/Fig3CF
• Example frompapers/Brette_2012/Fig3AB
• Example frompapers/Brette_2012/Fig5A
• Example frompapers/Brette_2012/Fig1
• Example frompapers/Brette_2012/Fig4

MorphologyIndexWrapper(morphology) A simpler version of IndexWrapper, not allowing for string indexing (Morphology is not a Group).

MorphologyIndexWrapper class

(Shortest import: from brian2.spatialneuron.morphology import MorphologyIndexWrapper)
class brian2.spatialneuron.morphology.MorphologyIndexWrapper(morphology)
A simpler version of `IndexWrapper`, not allowing for string indexing (`Morphology` is not a `Group`). It allows to use `morphology.indices[...]` instead of `morphology[...]._indices()`.

```python
Node(index, comp_name, x, y, z, diameter, ...)
```

### Attributes

**Node class**

*(Shortest import: from brian2.spatialneuron.morphology import Node)*

```python
class brian2.spatialneuron.morphology.Node:
    Bases: tuple

    Attributes
    -------

    child, comp_name, diameter, index, parent, x, y, z
    Aliases for field numbers 0, 1, 5, 0, 6, 2, 3, 4

    children
    Alias for field number 7
```

### Details

- **children**
  - Alias for field number 7
- **comp_name**
  - Alias for field number 1
- **diameter**
  - Alias for field number 5
- **index**
  - Alias for field number 0
- **parent**
  - Alias for field number 6
- **x**
  - Alias for field number 2
- **y**
  - Alias for field number 3
- **z**
  - Alias for field number 4
Section(**kwds)**

A section (unbranched structure), described as a sequence of truncated cones with potentially varying diameters and lengths per compartment.

---

### Section class

*(Shortest import: from brian2 import Section)*

class brian2.spatialneuron.morphology.Section(**kwds)

Bases: brian2.spatialneuron.morphology.Morphology

A section (unbranched structure), described as a sequence of truncated cones with potentially varying diameters and lengths per compartment.

**Parameters**

- **diameter**: Quantity
  
  Either a single value (the constant diameter along the whole section), or a value of length \(n+1\). When \(n+1\) values are given, they will be interpreted as the diameters at the start of the first compartment and the diameters at the end of each compartment (which is equivalent to: the diameter at the start of each compartment and the diameter at the end of the last compartment).

- **n**: int, optional
  
  The number of compartments in this section. Defaults to 1.

- **length**: Quantity, optional
  
  Either a single value (the total length of the section), or a value of length \(n\), the length of each individual compartment. Cannot be combined with the specification of coordinates.

- **x**: Quantity, optional
  
  \(n+1\) values, specifying the x coordinates of the start point of the first compartment and the end-points of all compartments (which is equivalent to: the start point of all compartments and the end point of the last compartment). The coordinates are interpreted as relative to the end point of the parent compartment (if any), so in most cases the start point should be 0*um. The common exception is a cylinder connecting to a Soma, here the start point can be used to make the cylinder start at the surface of the sphere instead of at its center. You can specify all of \(x\), \(y\), or \(z\) to specify a morphology in 3D, or only one or two out of them to specify a morphology in 1D or 2D.

- **y**: Quantity, optional
  
  See \(x\)

- **z**: Quantity, optional
  
  See \(x\)

- **type**: str, optional
  
  The type (e.g. "axon") of this Section.

---

**Attributes**
### area
The membrane surface area of each compartment in this section.

### diameter
The diameter at the middle of each compartment in this section.

### distance
The total distance between the midpoint of each compartment and the root of the morphology.

### end_diameter
The diameter at the end of each compartment in this section.

### end_distance
The distance to the root of the morphology at the end of this section.

### end_x_
The x coordinate (as a unitless floating point number) at the end of each compartment.

### end_y_
The y coordinate (as a unitless floating point number) at the end of each compartment.

### end_z_
The z coordinate (as a unitless floating point number) at the end of each compartment.

### length
The length of each compartment in this section.

### r_length_1
The geometry-dependent term to calculate the conductance between the start and the midpoint of each compartment.

### r_length_2
The geometry-dependent term to calculate the conductance between the midpoint and the end of each compartment.

### start_diameter
The diameter at the start of each compartment in this section.

### start_x_
The x coordinate (as a unitless floating point number) at the beginning of each compartment.

### start_y_
The y coordinate (as a unitless floating point number) at the beginning of each compartment.

### start_z_
The z coordinate (as a unitless floating point number) at the beginning of each compartment.

### volume
The volume of each compartment in this section.

### x_
The x coordinate (as a unitless floating point number) at the midpoint of each compartment.

### y_
The y coordinate (as a unitless floating point number) at the midpoint of each compartment.

### z_
The z coordinate (as a unitless floating point number) at the midpoint of each compartment.

### Methods

#### copy_section()
Create a copy of the current section (attributes of this section only, not re-creating the parent/children relation)

### Details

#### area
The membrane surface area of each compartment in this section. The surface area of each compartment is calculated as \( \frac{\pi}{2} (d_1 + d_2) \sqrt{\left(\frac{(d_1 - d_2)^2}{4} + l^2\right)} \), where \( l \) is the length of the compartment, and \( d_1 \) and \( d_2 \) are the diameter at the start and end of the compartment, respectively. Note that this surface area does not contain...
the area of the two disks at the two sides of the truncated cone.

diameter
- The diameter at the middle of each compartment in this section.

distance
- The total distance between the midpoint of each compartment and the root of the morphology.

end_diameter
- The diameter at the end of each compartment in this section.

end_distance
- The distance to the root of the morphology at the end of this section.

end_x_
- The x coordinate (as a unitless floating point number) at the end of each compartment. Returns None for morphologies without coordinates.

end_y_
- The y coordinate (as a unitless floating point number) at the end of each compartment. Returns None for morphologies without coordinates.

end_z_
- The z coordinate (as a unitless floating point number) at the end of each compartment. Returns None for morphologies without coordinates.

length
- The length of each compartment in this section.

r_length_1
- The geometry-dependent term to calculate the conductance between the start and the midpoint of each compartment. Dividing this value by the Intracellular resistivity gives the conductance.

r_length_2
- The geometry-dependent term to calculate the conductance between the midpoint and the end of each compartment. Dividing this value by the Intracellular resistivity gives the conductance.

start_diameter
- The diameter at the start of each compartment in this section.

start_x_
- The x coordinate (as a unitless floating point number) at the beginning of each compartment. Returns None for morphologies without coordinates.

start_y_
- The y coordinate (as a unitless floating point number) at the beginning of each compartment. Returns None for morphologies without coordinates.

start_z_
- The z coordinate (as a unitless floating point number) at the beginning of each compartment. Returns None for morphologies without coordinates.

volume
- The volume of each compartment in this section. The volume of each compartment is calculated as $\frac{\pi}{12}(d_1^3 + d_1d_2 + d_2^3)$, where $l$ is the length of the compartment, and $d_1$ and $d_2$ are the diameter at the start and end of the compartment, respectively.

x_
- The x coordinate (as a unitless floating point number) at the midpoint of each compartment. Returns None for morphologies without coordinates.
y_ The y coordinate (as a unitless floating point number) at the midpoint of each compartment. Returns None for morphologies without coordinates.

z_ The z coordinate (as a unitless floating point number) at the midpoint of each compartment. Returns None for morphologies without coordinates.

copy_section() Create a copy of the current section (attributes of this section only, not re-creating the parent/children relation)

Returns copy: Morphology

A copy of this section (without the links to the parent/children)

Tutorials and examples using this

- Example from papers/Brette_2012/Fig4

Soma(**kwds) A spherical, iso-potential soma.

Soma class

(Shortest import: from brian2 import Soma)

class brian2.spatialneuron.morphology.Soma(**kwds)
    Bases: brian2.spatialneuron.morphology.Morphology

A spherical, iso-potential soma.

Parameters

diameter : Quantity
    Diameter of the sphere.

x : Quantity, optional
    The x coordinate of the position of the soma.

y : Quantity, optional
    The y coordinate of the position of the soma.

z : Quantity, optional
    The z coordinate of the position of the soma.

type : str, optional
    The type of this section, defaults to 'soma'.

Attributes

area The membrane surface area of this section (as an array of length 1).
diameter The diameter of this section (as an array of length 1).
### Table 394 – continued from previous page

<table>
<thead>
<tr>
<th>Variable</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>distance</td>
<td>The total distance between the midpoint of this section and the root of the morphology.</td>
</tr>
<tr>
<td>end_distance</td>
<td>The distance to the root of the morphology at the end of this section.</td>
</tr>
<tr>
<td>end_x_</td>
<td>The x-coordinate of the current section (as an array of length 1).</td>
</tr>
<tr>
<td>end_y_</td>
<td>The y-coordinate of the current section (as an array of length 1).</td>
</tr>
<tr>
<td>end_z_</td>
<td>The z-coordinate of the current section (as an array of length 1).</td>
</tr>
<tr>
<td>length</td>
<td>The “length” (equal to (diameter)) of this section (as an array of length 1).</td>
</tr>
<tr>
<td>r_length_1</td>
<td>The geometry-dependent term to calculate the conductance between the start and the midpoint of each compartment.</td>
</tr>
<tr>
<td>r_length_2</td>
<td>The geometry-dependent term to calculate the conductance between the midpoint and the end of each compartment.</td>
</tr>
<tr>
<td>start_x_</td>
<td>The x-coordinate of the current section (as an array of length 1).</td>
</tr>
<tr>
<td>start_y_</td>
<td>The y-coordinate of the current section (as an array of length 1).</td>
</tr>
<tr>
<td>start_z_</td>
<td>The z-coordinate of the current section (as an array of length 1).</td>
</tr>
<tr>
<td>volume</td>
<td>The volume of this section (as an array of length 1).</td>
</tr>
<tr>
<td>x_</td>
<td>The x-coordinate of the current section (as an array of length 1).</td>
</tr>
<tr>
<td>y_</td>
<td>The y-coordinate of the current section (as an array of length 1).</td>
</tr>
<tr>
<td>z_</td>
<td>The z-coordinate of the current section (as an array of length 1).</td>
</tr>
</tbody>
</table>

### Methods

**copy_section()**

Create a copy of the current section (attributes of this section only, not re-creating the parent/children relation).

### Details

**area**
The membrane surface area of this section (as an array of length 1).

**diameter**
The diameter of this section (as an array of length 1).

**distance**
The total distance between the midpoint of this section and the root of the morphology. The *Soma* is most likely the root of the morphology, and therefore the **distance** is 0.

**end_distance**
The distance to the root of the morphology at the end of this section. Note that since a *Soma* is modeled
as a point (see docs of $x$, etc.), it does not add anything to the total distance, e.g. a section connecting to a Soma has a distance of 0 um at its start.

**end_x**
The x-coordinate of the current section (as an array of length 1). Note that a Soma is modelled as a “point” with finite surface/volume, equivalent to that of a sphere with the given diameter. It’s start-, midpoint-, and end-coordinates are therefore identical.

**end_y**
The y-coordinate of the current section (as an array of length 1). Note that a Soma is modelled as a “point” with finite surface/volume, equivalent to that of a sphere with the given diameter. It’s start-, midpoint-, and end-coordinates are therefore identical.

**end_z**
The z-coordinate of the current section (as an array of length 1). Note that a Soma is modelled as a “point” with finite surface/volume, equivalent to that of a sphere with the given diameter. It’s start-, midpoint-, and end-coordinates are therefore identical.

**length**
The “length” (equal to diameter) of this section (as an array of length 1).

**r_length_1**
The geometry-dependent term to calculate the conductance between the start and the midpoint of each compartment. Returns a fixed (high) value for a Soma, corresponding to a section with very low intracellular resistance.

**r_length_2**
The geometry-dependent term to calculate the conductance between the midpoint and the end of each compartment. Returns a fixed (high) value for a Soma, corresponding to a section with very low intracellular resistance.

**start_x**
The x-coordinate of the current section (as an array of length 1). Note that a Soma is modelled as a “point” with finite surface/volume, equivalent to that of a sphere with the given diameter. It’s start-, midpoint-, and end-coordinates are therefore identical.

**start_y**
The y-coordinate of the current section (as an array of length 1). Note that a Soma is modelled as a “point” with finite surface/volume, equivalent to that of a sphere with the given diameter. It’s start-, midpoint-, and end-coordinates are therefore identical.

**start_z**
The z-coordinate of the current section (as an array of length 1). Note that a Soma is modelled as a “point” with finite surface/volume, equivalent to that of a sphere with the given diameter. It’s start-, midpoint-, and end-coordinates are therefore identical.

**volume**
The volume of this section (as an array of length 1).

**x**
The x-coordinate of the current section (as an array of length 1). Note that a Soma is modelled as a “point” with finite surface/volume, equivalent to that of a sphere with the given diameter. It’s start-, midpoint-, and end-coordinates are therefore identical.

**y**
The y-coordinate of the current section (as an array of length 1). Note that a Soma is modelled as a “point” with finite surface/volume, equivalent to that of a sphere with the given diameter. It’s start-, midpoint-, and end-coordinates are therefore identical.
The z-coordinate of the current section (as an array of length 1). Note that a Soma is modelled as a “point” with finite surface/volume, equivalent to that of a sphere with the given diameter. It’s start-, midpoint-, and end-coordinates are therefore identical.

**copy_section()**

Create a copy of the current section (attributes of this section only, not re-creating the parent/children relation)

**Returns** copy: Morphology

A copy of this section (without the links to the parent/children)

---

**Tutorials and examples using this**

- Example `compartmental/spike_initiation`
- Example `compartmental/morphotest`
- Example `compartmental/bipolar_with_inputs`
- Example `compartmental/bipolar_with_inputs2`
- Example `compartmental/bipolar_cell`
- Example `frompapers/Brette_2012/Fig3CF`
- Example `frompapers/Brette_2012/Fig3AB`
- Example `frompapers/Brette_2012/Fig5A`
- Example `frompapers/Brette_2012/Fig1`
- Example `frompapers/Brette_2012/Fig4`

**SubMorphology** (morphology, i, j) A view on a subset of a section in a morphology.

**SubMorphology class**

(Shortest import: from brian2.spatialneuron.morphology import SubMorphology)

**class** brian2.spatialneuron.morphology.SubMorphology (morphology, i, j)

Bases: object

A view on a subset of a section in a morphology.

**Attributes**

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>area</td>
<td>The membrane surface area of each compartment in this sub-section.</td>
</tr>
<tr>
<td>diameter</td>
<td>The diameter at the middle of each compartment in this sub-section.</td>
</tr>
<tr>
<td>distance</td>
<td>The total distance between the midpoint of each compartment in this sub-section and the root of the morphology.</td>
</tr>
</tbody>
</table>

Continued on next page
### Table 397 – continued from previous page

<table>
<thead>
<tr>
<th>Variable</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>end_x</code></td>
<td>The x coordinate at the end of each compartment in this sub-section.</td>
</tr>
<tr>
<td><code>end_x_</code></td>
<td>The x coordinate (as a unitless floating point number) at the end of each compartment in this sub-section.</td>
</tr>
<tr>
<td><code>end_y</code></td>
<td>The y coordinate at the end of each compartment in this sub-section.</td>
</tr>
<tr>
<td><code>end_y_</code></td>
<td>The y coordinate (as a unitless floating point number) at the end of each compartment in this sub-section.</td>
</tr>
<tr>
<td><code>end_z</code></td>
<td>The z coordinate at the end of each compartment in this sub-section.</td>
</tr>
<tr>
<td><code>end_z_</code></td>
<td>The z coordinate (as a unitless floating point number) at the end of each compartment in this sub-section.</td>
</tr>
<tr>
<td><code>length</code></td>
<td>The length of each compartment in this sub-section.</td>
</tr>
<tr>
<td><code>n</code></td>
<td>The number of compartments in this sub-section.</td>
</tr>
<tr>
<td><code>n_sections</code></td>
<td>The number of sections in this sub-section (always 1).</td>
</tr>
<tr>
<td><code>r_length_1</code></td>
<td>The geometry-dependent term to calculate the conductance between the start and the midpoint of each compartment in this sub-section.</td>
</tr>
<tr>
<td><code>r_length_2</code></td>
<td>The geometry-dependent term to calculate the conductance between the midpoint and the end of each compartment in this sub-section.</td>
</tr>
<tr>
<td><code>start_x</code></td>
<td>The x coordinate at the beginning of each compartment in this sub-section.</td>
</tr>
<tr>
<td><code>start_x_</code></td>
<td>The x coordinate (as a unitless floating point number) at the beginning of each compartment in this sub-section.</td>
</tr>
<tr>
<td><code>start_y</code></td>
<td>The y coordinate at the beginning of each compartment in this sub-section.</td>
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<tr>
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<td>The z coordinate at the beginning of each compartment in this sub-section.</td>
</tr>
<tr>
<td><code>start_z_</code></td>
<td>The z coordinate (as a unitless floating point number) at the beginning of each compartment in this sub-section.</td>
</tr>
<tr>
<td><code>volume</code></td>
<td>The volume of each compartment in this sub-section.</td>
</tr>
<tr>
<td><code>x</code></td>
<td>The x coordinate at the midpoint of each compartment in this sub-section.</td>
</tr>
<tr>
<td><code>x_</code></td>
<td>The x coordinate (as a unitless floating point number) at the midpoint of each compartment in this sub-section.</td>
</tr>
<tr>
<td><code>y</code></td>
<td>The y coordinate at the midpoint of each compartment in this sub-section.</td>
</tr>
<tr>
<td><code>y_</code></td>
<td>The y coordinate (as a unitless floating point number) at the midpoint of each compartment in this sub-section.</td>
</tr>
<tr>
<td><code>z</code></td>
<td>The z coordinate at the midpoint of each compartment in this sub-section.</td>
</tr>
</tbody>
</table>

**Continued on next page**
### Table 397 – continued from previous page

<table>
<thead>
<tr>
<th>Field</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>z_</code></td>
<td>The z coordinate (as a unitless floating point number) at the midpoint of each compartment in this sub-section.</td>
</tr>
</tbody>
</table>

#### Details

<table>
<thead>
<tr>
<th>Field</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>area</strong></td>
<td>The membrane surface area of each compartment in this sub-section.</td>
</tr>
<tr>
<td><strong>diameter</strong></td>
<td>The diameter at the middle of each compartment in this sub-section.</td>
</tr>
<tr>
<td><strong>distance</strong></td>
<td>The total distance between the midpoint of each compartment in this sub-section and the root of the morphology.</td>
</tr>
<tr>
<td><strong>end_x</strong></td>
<td>The x coordinate at the end of each compartment in this sub-section. Returns None for morphologies without coordinates.</td>
</tr>
<tr>
<td><strong>end_x_</strong></td>
<td>The x coordinate (as a unitless floating point number) at the end of each compartment in this sub-section. Returns None for morphologies without coordinates.</td>
</tr>
<tr>
<td><strong>end_y</strong></td>
<td>The y coordinate at the end of each compartment in this sub-section. Returns None for morphologies without coordinates.</td>
</tr>
<tr>
<td><strong>end_y_</strong></td>
<td>The y coordinate (as a unitless floating point number) at the end of each compartment in this sub-section. Returns None for morphologies without coordinates.</td>
</tr>
<tr>
<td><strong>end_z</strong></td>
<td>The z coordinate at the end of each compartment in this sub-section. Returns None for morphologies without coordinates.</td>
</tr>
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<td><strong>end_z_</strong></td>
<td>The z coordinate (as a unitless floating point number) at the end of each compartment in this sub-section. Returns None for morphologies without coordinates.</td>
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<tr>
<td><strong>length</strong></td>
<td>The length of each compartment in this sub-section.</td>
</tr>
<tr>
<td><strong>n</strong></td>
<td>The number of compartments in this sub-section.</td>
</tr>
<tr>
<td><strong>n_sections</strong></td>
<td>The number of sections in this sub-section (always 1).</td>
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<tr>
<td><strong>r_length_1</strong></td>
<td>The geometry-dependent term to calculate the conductance between the start and the midpoint of each compartment in this sub-section. Dividing this value by the Intracellular resistivity gives the conductance.</td>
</tr>
<tr>
<td><strong>r_length_2</strong></td>
<td>The geometry-dependent term to calculate the conductance between the midpoint and the end of each compartment in this sub-section. Dividing this value by the Intracellular resistivity gives the conductance.</td>
</tr>
<tr>
<td><strong>start_x</strong></td>
<td>The x coordinate at the beginning of each compartment in this sub-section. Returns None for morphologies without coordinates.</td>
</tr>
</tbody>
</table>
**start_x**
The x coordinate (as a unitless floating point number) at the beginning of each compartment in this sub-section. Returns *None* for morphologies without coordinates.

**start_y**
The y coordinate at the beginning of each compartment in this sub-section. Returns *None* for morphologies without coordinates.

**start_y**
The y coordinate (as a unitless floating point number) at the beginning of each compartment in this sub-section. Returns *None* for morphologies without coordinates.

**start_z**
The x coordinate at the beginning of each compartment in this sub-section. Returns *None* for morphologies without coordinates.

**start_z**
The z coordinate (as a unitless floating point number) at the beginning of each compartment in this sub-section. Returns *None* for morphologies without coordinates.

**volume**
The volume of each compartment in this sub-section.

x
The x coordinate at the midpoint of each compartment in this sub-section. Returns *None* for morphologies without coordinates.

x_
The x coordinate (as a unitless floating point number) at the midpoint of each compartment in this sub-section. Returns *None* for morphologies without coordinates.

y
The y coordinate at the midpoint of each compartment in this sub-section. Returns *None* for morphologies without coordinates.

y_
The y coordinate (as a unitless floating point number) at the midpoint of each compartment in this sub-section. Returns *None* for morphologies without coordinates.

z
The z coordinate at the midpoint of each compartment in this sub-section. Returns *None* for morphologies without coordinates.

z_
The z coordinate (as a unitless floating point number) at the midpoint of each compartment in this sub-section. Returns *None* for morphologies without coordinates.

| Topology(morphology) | A representation of the topology of a Morphology. |

**Topology class**

*(Shortest import: from brian2.spatialneuron.morphology import Topology)*

class brian2.spatialneuron.morphology.Topology(morphology)

Bases: object

A representation of the topology of a Morphology. Has a useful string representation, inspired by NEURON’s topology function.
spatialneuron module

Exported members: SpatialNeuron

Classes

<table>
<thead>
<tr>
<th>FlatMorphology(morphology)</th>
<th>Container object to store the flattened representation of a morphology.</th>
</tr>
</thead>
</table>

FlatMorphology class

(Shortest import: from brian2.spatialneuron.spatialneuron import FlatMorphology)

class brian2.spatialneuron.spatialneuron.FlatMorphology(morphology)

Bases: object

Container object to store the flattened representation of a morphology. Note that all values are stored as numpy arrays without unit information (i.e. in base units).

<table>
<thead>
<tr>
<th>SpatialNeuron([morphology, model, ...])</th>
<th>A single neuron with a morphology and possibly many compartments.</th>
</tr>
</thead>
</table>

SpatialNeuron class

(Shortest import: from brian2 import SpatialNeuron)

class brian2.spatialneuron.spatialneuron.SpatialNeuron(morphology=None, 
model=None, threshold=None, refractory=False, reset=None, 
events=None, threshold_location=None, dt=None, clock=None, 
order=0, Cm=0.009 * metre ** -4 * kilogram ** -1 * second ** 4 * amp ** 2, 
Ri=1.5 * metre ** 3 * kilogram * second ** -3 * amp ** -2, 
name='spatialneuron*', 
dtype=None, namespace=None, 
method=('exact', 'exponential_euler', 'rk2', 'heun'), 
method_options=None)

Bases: brian2.groups.neurongroup.NeuronGroup

A single neuron with a morphology and possibly many compartments.

Parameters

<table>
<thead>
<tr>
<th>morphology</th>
<th>Morphology</th>
</tr>
</thead>
<tbody>
<tr>
<td>The morphology of the neuron.</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>model</th>
<th>(str, Equations)</th>
</tr>
</thead>
<tbody>
<tr>
<td>The equations defining the group.</td>
<td></td>
</tr>
</tbody>
</table>
method : (str, function), optional

The numerical integration method. Either a string with the name of a registered method (e.g. "euler") or a function that receives an `Equations` object and returns the corresponding abstract code. If no method is specified, a suitable method will be chosen automatically.

threshold : str, optional

The condition which produces spikes. Should be a single line boolean expression.

threshold_location : (int, `Morphology`), optional

Compartment where the threshold condition applies, specified as an integer (compartment index) or a `Morphology` object corresponding to the compartment (e.g. `morpho.axon[10*um]`). If unspecified, the threshold condition applies at all compartments.

Cm : `Quantity`, optional

Specific capacitance in uF/cm**2 (default 0.9). It can be accessed and modified later as a state variable. In particular, its value can differ in different compartments.

Ri : `Quantity`, optional

Intracellular resistivity in ohm.cm (default 150). It can be accessed as a shared state variable, but modified only before the first run. It is uniform across the neuron.

reset : str, optional

The (possibly multi-line) string with the code to execute on reset.

events : dict, optional

User-defined events in addition to the “spike” event defined by the `threshold`. Has to be a mapping of strings (the event name) to strings (the condition) that will be checked.

refractory : {str, `Quantity`}, optional

Either the length of the refractory period (e.g. 2*ms), a string expression that evaluates to the length of the refractory period after each spike (e.g. '(1 + rand())*ms'), or a string expression evaluating to a boolean value, given the condition under which the neuron stays refractory after a spike (e.g. 'v > -20*mV')

namespace : dict, optional

A dictionary mapping identifier names to objects. If not given, the namespace will be filled in at the time of the call of `Network.run`, with either the values from the namespace argument of the `Network.run` method or from the local context, if no such argument is given.

dtype : (dtype, dict), optional

The `numpy.dtype` that will be used to store the values, or a dictionary specifying the type for variable names. If a value is not provided for a variable (or no value is provided at all), the preference setting `core.default_float_dtype` is used.

dt : `Quantity`, optional

The time step to be used for the simulation. Cannot be combined with the `clock` argument.

clock : `Clock`, optional
The update clock to be used. If neither a clock, nor the `dt` argument is specified, the `defaultclock` will be used.

**order**: int, optional

The priority of of this group for operations occurring at the same time step and in the same scheduling slot. Defaults to 0.

**name**: str, optional

A unique name for the group, otherwise use `spatialneuron_0`, etc.

**Attributes**

| user_equations | The original equations as specified by the user (i.e. before inserting point-currents into the membrane equation, before adding all the internally used variables and constants, etc.). |

**Methods**

| spatialneuron_attribute(neuron, name) | Selects a subtree from `SpatialNeuron` neuron and returns a `SpatialSubgroup`. |
| spatialneuron_segment(neuron, item) | Selects a segment from `SpatialNeuron` neuron, where item is a slice of either compartment indexes or distances. |

**Details**

**user_equations**

The original equations as specified by the user (i.e. before inserting point-currents into the membrane equation, before adding all the internally used variables and constants, etc.).

**static spatialneuron_attribute (neuron, name)**

Selects a subtree from `SpatialNeuron` neuron and returns a `SpatialSubgroup`. If it does not exist, returns the `Group` attribute.

**static spatialneuron_segment (neuron, item)**

Selects a segment from `SpatialNeuron` neuron, where item is a slice of either compartment indexes or distances. Note a: segment is not a `SpatialNeuron`, only a `Group`.

**Tutorials and examples using this**

- Example `compartmental/spike_initiation`
- Example `compartmental/morphotest`
- Example `compartmental/bipolar_with_inputs`
- Example `compartmental/bipolar_with_inputs2`
- Example `compartmental/hh_with_spikes`
- Example `compartmental/infinite_cable`
- Example `compartmental/cylinder`
- Example `compartmental/hodgkin_huxley_1952`
- Example `compartmental/bipolar_cell`
SpatialStateUpdater class

```python
SpatialStateUpdater(group, method, clock[, ...])
```

The CodeRunner that updates the state variables of a SpatialNeuron at every timestep.

**Methods**

```python
before_run(run_namespace)
```

Optional method to prepare the object before a run.

**Details**

```python
before_run(run_namespace)
```

Optional method to prepare the object before a run.

TODO

SpatialSubgroup class

```python
SpatialSubgroup(source, start, stop, morphology) => A subgroup of a SpatialNeuron.
```

**Parameters**

- **source**: int
  
  First compartment.
stop : int
        Ending compartment, not included (as in slices).

morphology : Morphology
        Morphology corresponding to the subgroup (not the full morphology).

name : str, optional
        Name of the subgroup.

6.6.13 stateupdaters package

Module for transforming model equations into “abstract code” that can be then be further translated into executable
code by the codegen module.

Exported members: StateUpdateMethod, linear, exact, independent, milstein, heun, euler, rk2, rk4, ExplicitStateUpdater, exponential_euler, gsl_rk2, gsl_rk4, gsl_rkf45, gsl_rkck, gsl_rk8pd

GSL module

Module containg the StateUpdateMethod for integration using the ODE solver provided in the GNU Scientific Library
(GSL)

Exported members: gsl_rk2, gsl_rk4, gsl_rkf45, gsl_rkck, gsl_rk8pd

Classes

GSLContainer(method_options, integrator[, ...])
        Class that contains information (equation- or integrator-related) required for later code generation

GSLContainer class

(Shortest import: from brian2.stateupdaters.GSL import GSLContainer)

class brian2.stateupdaters.GSL.GSLContainer(method_options, integrator, abstract_code=None, needed_variables=[],
variable_flags=[])
        Bases: object
        Class that contains information (equation- or integrator-related) required for later code generation

Methods

_call_(obj)  Transfer the code object class saved in self to the object sent as an argument.

get_codeobj_class()  Return codeobject class based on target language and device.
Details

```python
__call__(obj)
```
Transfer the code object class saved in self to the object sent as an argument.

This method is returned when calling `GSLStateUpdater`. This class inherits from `StateUpdateMethod` which originally only returns abstract code. However, with GSL this returns a method because more is needed than just the abstract code: the state updater requires its own `CodeObject` that is different from the other `NeuronGroup` objects. This method adds this `CodeObject` to the `StateUpdater` object (and also adds the variables ‘t’, ‘dt’, and other variables that are needed in the `GSLCodeGenerator`.

**Parameters**

- `obj`: `GSLStateUpdater`
  the object that the codeobj_class and other variables need to be transferred to

**Returns**

- `abstract_code`: str
  The abstract code (translated equations), that is returned conventionally by brian and used for later code generation in the `CodeGenGenerator.translate` method.

```python
get_codeobj_class()
```
Return codeobject class based on target language and device.

Choose which version of the GSL `CodeObject` to use. If `isinstance(device, CPPStandaloneDevice)`, then we want the `GSLCPPStandaloneCodeObject`. Otherwise the return value is based on `prefs.codegen.target`.

**Returns**

- `code_object`: class
  The respective `CodeObject` class (i.e. either `GSLWeaveCodeObject`, `GSLCythonCodeObject`, or `GSLCPPStandaloneCodeObject`).

---

**GSLStateUpdater class**

*(Shortest import: from brian2.stateupdaters.GSL import GSLStateUpdater)*

```python
class brian2.stateupdaters.GSL.GSLStateUpdater(integrator)
Bases: brian2.stateupdaters.base.StateUpdateMethod
```
A statupdater that rewrites the differential equations so that the GSL generator knows how to write the code in the target language.

New in version 2.1.

**Methods**

```python
__call__(equations[, variables, method_options])
```
Translate equations to `abstract_code`. 
Details

```python
__call__(equations, variables=None, method_options=None)
```
Translate equations to abstract_code.

**Parameters**
- `equations`: `Equations` object containing the equations that describe the ODE system
- `variables`: dict
dictionary containing str, `Variable` pairs

**Returns**
- `method`: callable
  Method that needs to be called with `StateUpdater` to add `CodeObject` class and some other variables so these can be sent to the `CodeGenerator`

**Objects**

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>gsl_rk2</code></td>
<td>A statupdater that rewrites the differential equations so that the GSL generator knows how to write the code in the target language.</td>
</tr>
</tbody>
</table>

**gsl_rk2 object**

(Shortest import: from brian2 import gsl_rk2)

brian2.stateupdaters.GSL.gsl_rk2 = <brian2.stateupdaters.GSL.GSLStateUpdater object>
A statupdater that rewrites the differential equations so that the GSL generator knows how to write the code in the target language.
New in version 2.1.

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>gsl_rk4</code></td>
<td>A statupdater that rewrites the differential equations so that the GSL generator knows how to write the code in the target language.</td>
</tr>
</tbody>
</table>

**gsl_rk4 object**

(Shortest import: from brian2 import gsl_rk4)

brian2.stateupdaters.GSL.gsl_rk4 = <brian2.stateupdaters.GSL.GSLStateUpdater object>
A statupdater that rewrites the differential equations so that the GSL generator knows how to write the code in the target language.
New in version 2.1.

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>gsl_rk8pd</code></td>
<td>A statupdater that rewrites the differential equations so that the GSL generator knows how to write the code in the target language.</td>
</tr>
</tbody>
</table>

**gsl_rk8pd object**

(Shortest import: from brian2 import gsl_rk8pd)
brian2.stateupdaters.GSL.gsl_rk8pd = <brian2.stateupdaters.GSL.GSLStateUpdater object>
A statupdater that rewrites the differential equations so that the GSL generator knows how to write the code in
the target language.
New in version 2.1.

**gsl_rkck**

A statupdater that rewrites the differential equations so that the GSL generator knows how to write the code in
the target language.

**gsl_rkck object**

(Shortest import: from brian2 import gsl_rkck)

brian2.stateupdaters.GSL.gsl_rkck = <brian2.stateupdaters.GSL.GSLStateUpdater object>
A statupdater that rewrites the differential equations so that the GSL generator knows how to write the code in
the target language.
New in version 2.1.

**gsl_rkf45**

A statupdater that rewrites the differential equations so that the GSL generator knows how to write the code in
the target language.

**gsl_rkf45 object**

(Shortest import: from brian2 import gsl_rkf45)

brian2.stateupdaters.GSL.gsl_rkf45 = <brian2.stateupdaters.GSL.GSLStateUpdater object>
A statupdater that rewrites the differential equations so that the GSL generator knows how to write the code in
the target language.
New in version 2.1.

**base module**

*Exported members: StateUpdateMethod*

**Classes**

*StateUpdateMethod*

**Attributes**

**StateUpdateMethod class**

(Shortest import: from brian2 import StateUpdateMethod)

class brian2.stateupdaters.base.StateUpdateMethod
    Bases: object
Attributes

**stateupdaters**

A dictionary mapping state updater names to `StateUpdateMethod` objects

Methods

```python
__call__(equations[, variables, method_options])

Generate abstract code from equations.

Parameters:
- **equations**: `Equations`
  The model equations.
- **variables**: dict, optional
  The `Variable` objects for the model variables.
- **method_options**: dict, optional
  Additional options specific to the state updater.

Returns:
- **code**: str
  The abstract code performing a state update step.

**apply_stateupdater(equations, variables, method)**

Applies a given state updater to equations.

Parameters:
- **equations**: `Equations`
  The model equations.
- **variables**: dict
  The dictionary of `Variable` objects, describing the internal model variables.
- **method**: {callable, str, list of str}
  A method for the state updater.

**register(name, stateupdater)**

Register a state updater.

Details

**stateupdaters**

A dictionary mapping state updater names to `StateUpdateMethod` objects

```python
__call__(equations, variables=\_\_\_None\_\_, method_options=\_\_\_None\_\_)

Generate abstract code from equations. The method also gets the the variables because some state updaters have to check whether variable names reflect other state variables (which can change from timestep to timestep) or are external values (which stay constant during a run) For convenience, this arguments are optional – this allows to directly see what code a state updater generates for a set of equations by simply writing `euler(eqs)`, for example.

Parameters:
- **equations**: `Equations`
  The model equations.
- **variables**: dict, optional
  The `Variable` objects for the model variables.
- **method_options**: dict, optional
  Additional options specific to the state updater.

Returns:
- **code**: str
  The abstract code performing a state update step.

```
```
A callable usable as a state updater, the name of a registered state updater or a list of names of state updaters.

**Returns**  
`abstract_code`: str  
The code integrating the given equations.

```python
static register(name, stateupdater)
```
Register a state updater. Registered state updaters can be referred to via their name.

**Parameters**  
`name`: str  
A short name for the state updater (e.g. 'euler')  
`stateupdater`: StateUpdaterMethod  
The state updater object, e.g. an `ExplicitStateUpdater`.

---

**UnsupportedEquationsException**

---

**UnsupportedEquationsException class**

*(Shortest import: from brian2.stateupdaters.base import UnsupportedEquationsException)*

```python
class brian2.stateupdaters.base.UnsupportedEquationsException
    Bases: Exception

Functions

```extract_method_options``(method_options, ...)
Helper function to check `method_options` against options understood by this state updater, and setting default values for all unspecified options.

---

**extract_method_options function**

*(Shortest import: from brian2.stateupdaters.base import extract_method_options)*

```python
brian2.stateupdaters.base.extract_method_options(method_options, default_options)
```
Helper function to check `method_options` against options understood by this state updater, and setting default values for all unspecified options.

**Parameters**  
`method_options`: dict or None  
The options that the user specified for the state update.

`default_options`: dict  
The default option values for this state updater (each admissible option needs to be present in this dictionary). To specify that a state updater does not take any options, provide an empty dictionary as the argument.

**Returns**  
`options`: dict  
The final dictionary with all the options either at their default or at the user-specified value.
Raises

**KeyError** If the user specifies an option that is not understood by this state updater.

Examples

```python
>>> options = extract_method_options({'a': True}, default_options={'b': False, 'c': False})  # doctest: +IGNORE_EXCEPTION_DETAIL
Traceback (most recent call last):
... KeyErrors: 'method_options specifies "a", but this is not an option for this state updater. Available options are: "b", "c".'

>>> options = extract_method_options({'a': True}, default_options={})  # doctest: +IGNORE_EXCEPTION_DETAIL
Traceback (most recent call last):
... KeyErrors: 'method_options specifies "a", but this is not an option for this state updater. This state updater does not accept any options.'

>>> options = extract_method_options({'a': True}, default_options={'a': False, 'b': False})
>>> sorted(options.items())
[('a', True), ('b', False)]
```

**exact module**

Exact integration for linear equations.

*Exported members:* `linear, exact, independent`

**Classes**

- **IndependentStateUpdater**
  A state update for equations that do not depend on other state variables, i.e. 1-dimensional differential equations.
  The individual equations are solved by sympy.

  Deprecated since version 2.1: This method might be removed from future versions of Brian.

**Methods**

- **__call__**(equations[, variables, method_options]) Generate abstract code from equations.
Details

___call___(equations, variables=None, method_options=None)
Generate abstract code from equations. The method also gets the the variables because some state updaters have to check whether variable names reflect other state variables (which can change from timestep to timestep) or are external values (which stay constant during a run) For convenience, this arguments are optional – this allows to directly see what code a state updater generates for a set of equations by simply writing euler(eqs), for example.

Parameters

  equations : Equations
    The model equations.
  variables : dict, optional
    The Variable objects for the model variables.
  method_options : dict, optional
    Additional options specific to the state updater.

Returns

  code : str
    The abstract code performing a state update step.

LinearStateUpdater
A state updater for linear equations.

LinearStateUpdater class

(Shortest import: from brian2.stateupdaters.exact import LinearStateUpdater)

class brian2.stateupdaters.exact.LinearStateUpdater
  Bases: brian2.stateupdaters.base.StateUpdateMethod

  A state updater for linear equations. Derives a state updater step from the analytical solution given by sympy. Uses the matrix exponential (which is only implemented for diagonalizable matrices in sympy).

Methods

___call___(equations[, variables, method_options])  Generate abstract code from equations.

Details

___call___(equations, variables=None, method_options=None)
Generate abstract code from equations. The method also gets the the variables because some state updaters have to check whether variable names reflect other state variables (which can change from timestep to timestep) or are external values (which stay constant during a run) For convenience, this arguments are optional – this allows to directly see what code a state updater generates for a set of equations by simply writing euler(eqs), for example.

Parameters

  equations : Equations
    The model equations.
variables: dict, optional
The Variable objects for the model variables.

method_options: dict, optional
Additional options specific to the state updater.

Returns:

code: str
The abstract code performing a state update step.

Functions

get_linear_system(eqs, variables) Convert equations into a linear system using sympy.

get_linear_system function

(Shortest import: from brian2.stateupdaters.exact import get_linear_system)

brian2.stateupdaters.exact.get_linear_system(eqs, variables)
Convert equations into a linear system using sympy.

Parameters eqs: Equations
The model equations.

Returns (diff_eq_names, coefficients, constants): (list of str, sympy.Matrix, sympy.Matrix)
A tuple containing the variable names (diff_eq_names) corresponding to the rows of the matrix coefficients and the vector constants, representing the system of equations in the form M * X + B

Raises

ValueError If the equations cannot be converted into an M * X + B form.

Objects

exact A state updater for linear equations.

exact object

(Shortest import: from brian2 import exact)

brian2.stateupdaters.exact.exact = LinearStateUpdater()
A state updater for linear equations. Derives a state updater step from the analytical solution given by sympy. Uses the matrix exponential (which is only implemented for diagonalizable matrices in sympy).
**independent**

A state update for equations that do not depend on other state variables, i.e.

**independent object**

```python
brian2.stateupdaters.exact.independent = <brian2.stateupdaters.exact.IndependentStateUpdater object>
```

A state update for equations that do not depend on other state variables, i.e. 1-dimensional differential equations. The individual equations are solved by sympy.

Deprecated since version 2.1: This method might be removed from future versions of Brian.

**linear**

A state updater for linear equations.

**linear object**

```python
brian2.stateupdaters.exact.linear = LinearStateUpdater()
```

A state updater for linear equations. Derives a state updater step from the analytical solution given by sympy. Uses the matrix exponential (which is only implemented for diagonalizable matrices in sympy).

**explicit module**

*Exported members:* **milstein**, **heun**, **euler**, **rk2**, **rk4**, **ExplicitStateUpdater**

**Classes**

```python
ExplicitStateUpdater(description[, ...])
```

An object that can be used for defining state updaters via a simple description (see below).

**ExplicitStateUpdater class**

```python
class brian2.stateupdaters.explicit.ExplicitStateUpdater (description, stochastic=None, custom_check=None)
```

**Bases:** **brian2.stateupdaters.base.StateUpdateMethod**

An object that can be used for defining state updaters via a simple description (see below). Resulting instances can be passed to the **method** argument of the **NeuronGroup** constructor. As other state updater functions the **ExplicitStateUpdater** objects are callable, returning abstract code when called with an **Equations** object.

A description of an explicit state updater consists of a (multi-line) string, containing assignments to variables and a final “x_new = ...”, stating the integration result for a single timestep. The assignments can be used to define an arbitrary number of intermediate results and can refer to \( f(x, t) \) (the function being integrated, as a function of \( x \), the previous value of the state variable and \( t \), the time) and \( dt \), the size of the timestep.

For example, to define a Runge-Kutta 4 integrator (already provided as **rk4**), use:
\[
\begin{align*}
    k_1 &= dt \cdot f(x, t) \\
    k_2 &= dt \cdot f(x + k_1/2, t + dt/2) \\
    k_3 &= dt \cdot f(x + k_2/2, t + dt/2) \\
    k_4 &= dt \cdot f(x + k_3, t + dt) \\
    x_{\text{new}} &= x + \frac{(k_1 + 2k_2 + 2k_3 + k_4)}{6}
\end{align*}
\]

Note that for stochastic equations, the function \( f \) only corresponds to the non-stochastic part of the equation. The additional function \( g \) corresponds to the stochastic part that has to be multiplied with the stochastic variable \( x_i \) (a standard normal random variable – if the algorithm needs a random variable with a different variance/mean you have to multiply/add it accordingly). Equations with more than one stochastic variable do not have to be treated differently, the part referring to \( g \) is repeated for all stochastic variables automatically.

Stochastic integrators can also make reference to \( dW \) (a normal distributed random number with variance \( dt \)) and \( g(x, t) \), the stochastic part of an equation. A stochastic state updater could therefore use a description like:

\[
x_{\text{new}} = x + dt \cdot f(x, t) + g(x, t) \cdot dW
\]

For simplicity, the same syntax is used for state updaters that only support additive noise, even though \( g(x, t) \) does not depend on \( x \) or \( t \) in that case.

There are some restrictions on the complexity of the expressions (but most can be worked around by using intermediate results as in the above Runge-Kutta example): Every statement can only contain the functions \( f \) and \( g \) once; The expressions have to be linear in the functions, e.g. you can use \( dt \cdot f(x, t) \) but not \( f(x, t)^2 \).

**Parameters**

- **description**: str
  
  A state updater description (see above).

- **stochastic**: \{None, 'additive', 'multiplicative'\}
  
  What kind of stochastic equations this state updater supports: None means no support of stochastic equations, 'additive' means only equations with additive noise and 'multiplicative' means supporting arbitrary stochastic equations.

**Raises**

- **ValueError**: If the parsing of the description failed.

**See also:**

- euler, rk2, rk4, milstein

**Notes**

Since clocks are updated after the state update, the time \( t \) used in the state update step is still at its previous value. Enumerating the states and discrete times, \( x_{\text{new}} = x + dt \cdot f(x, t) \) is therefore understood as \( x_{i+1} = x_i + dt \cdot f(x_i, t_i) \), yielding the correct forward Euler integration. If the integrator has to refer to the time at the end of the timestep, simply use \( t + dt \) instead of \( t \).

**Attributes**

<table>
<thead>
<tr>
<th>DESCRIPTION</th>
<th>A complete state updater description</th>
</tr>
</thead>
<tbody>
<tr>
<td>EXPRESSION</td>
<td>A single expression</td>
</tr>
</tbody>
</table>

Continued on next page
Table 429 – continued from previous page

<table>
<thead>
<tr>
<th>OUTPUT</th>
<th>The last line of a state updater description</th>
</tr>
</thead>
<tbody>
<tr>
<td>STATEMENT</td>
<td>An assignment statement</td>
</tr>
<tr>
<td>TEMP_VAR</td>
<td>Legal names for temporary variables</td>
</tr>
</tbody>
</table>

**Methods**

<table>
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</tr>
<tr>
<td><strong>call</strong></td>
<td>Apply a state updater description to model equations.</td>
</tr>
</tbody>
</table>

**replace_func**

Replace a single occurrence of $f(x, t)$ or $g(x, t)$: `expr` is the non-stochastic (in the case of $f$) or stochastic part ($g$) of the expression defining the right-hand-side of the differential equation describing `var()`. 

**Details**

**DESCRIPTION**

A complete state updater description

**EXPRESSION**

A single expression

**OUTPUT**

The last line of a state updater description

**STATEMENT**

An assignment statement

**TEMP_VAR**

Legal names for temporary variables

**__call__**(eqs[, variables, method_options])

Apply a state updater description to model equations.

**replace_func**(x, t, expr, temp_vars, eq_symbols)

Used to replace a single occurrence of $f(x, t)$ or $g(x, t)$: `expr` is the non-stochastic (in the case of $f$) or stochastic part ($g$) of the expression defining the right-hand-side of the differential equation describing `var()`.

**DESCRIPTION**

Requires all given `ParseExpression`s to be found in the given order. Expressions may be separated by whitespace. May be constructed using the `+` operator. May also be constructed using the `-` operator, which will suppress backtracking.

Example:

```python
integer = Word(nums)
name_expr = OneOrMore(Word(alphas))

expr = And([integer("id"), name_expr("name"), integer("age")])
# more easily written as:
expr = integer("id") + name_expr("name") + integer("age")
```

**EXPRESSION**

Token for matching strings that match a given regular expression. Defined with string specifying the regular expression in a form recognized by the stdlib Python re module. If the given regex contains named groups (defined using `(?P<name>...)`), these will be preserved as named parse results.

If instead of the Python stdlib re module you wish to use a different RE module (such as the `regex`
module), you can replace it by either building your Regex object with a compiled RE that was compiled using regex:

Example:

```python
realnum = Regex(r"[+-]?\d+\.\d*")
date = Regex(r'(\P{year}\d{4})-(\P{month}\d{2})-(\P{day}\d{2})')
# ref: https://stackoverflow.com/questions/267399/how-do-you-match-only-valid-
roman = Regex(r"[M{0,4}(CM|CD|D?{0,3})(XC|XL|L?X{0,3})(IX|IV|V?I?{0,3})]")
# use regex module instead of stdlib re module to construct a Regex using
import regex
parser = pp.Regex(regex.compile(r'\[0-9\]'))
```

**OUTPUT ()**

Converter to return the matched tokens as a list - useful for returning tokens of ZeroOrMore and OneOrMore expressions.

Example:

```python
ident = Word(alphas)
um = Word(nums)
term = ident | num
func = ident + Optional(delimitedList(term))
print(func.parseString("fn a, b, 100"))  # -> ['fn', 'a', 'b', '100']

func = ident + Group(Optional(delimitedList(term)))
print(func.parseString("fn a, b, 100"))  # -> ['fn', ['a', 'b', '100']]
```

**STATEMENT ()**

Converter to return the matched tokens as a list - useful for returning tokens of ZeroOrMore and OneOrMore expressions.

Example:

```python
ident = Word(alphas)
um = Word(nums)
term = ident | num
func = ident + Optional(delimitedList(term))
print(func.parseString("fn a, b, 100"))  # -> ['fn', 'a', 'b', '100']

func = ident + Group(Optional(delimitedList(term)))
print(func.parseString("fn a, b, 100"))  # -> ['fn', ['a', 'b', '100']]
```

**TEMP_VAR()**

Requires all given ParseExpressions to be found in the given order. Expressions may be separated by whitespace. May be constructed using the ‘+’ operator. May also be constructed using the ‘-’ operator, which will suppress backtracking.

Example:

```python
integer = Word(nums)
name_expr = OneOrMore(Word(alphas))
expr = And([integer("id"), name_expr("name"), integer("age")])
# more easily written as:
expr = integer("id") + name_expr("name") + integer("age")
```
__call__ (eqs, variables=None, method_options=None)

Apply a state updater description to model equations.

Parameters eqs: Equations

The equations describing the model

variables: dict-like, optional

The Variable objects for the model. Ignored by the explicit state updater.

method_options : dict, optional

Additional options to the state updater (not used at the moment for the explicit state updaters).

Examples

```python
>>> from brian2 import *
>>> eqs = Equations('dv/dt = -v / tau : volt')
>>> print(euler(eqs))
_\dd{v}{t} = -dt*v/tau + v
v = _\dd{v}{t}
```

replace_func (x, t, expr, temp_vars, eq_symbols, stochastic_variable=None)

Used to replace a single occurrence of f(x, t) or g(x, t): expr is the non-stochastic (in the case of f) or stochastic part (g) of the expression defining the right-hand-side of the differential equation describing var(). It replaces the variable var() with the value given as x and t by the value given for t. Intermediate variables will be replaced with the appropriate replacements as well.

For example, in the rk2 integrator, the second step involves the calculation of f(k/2 + x, dt/2 + t). If var() is v and expr is -v / tau, this will result in -(k_v/2 + v) / tau.

Note that this deals with only one state variable var(), given as an argument to the surrounding _generate_RHS function.

Functions

diagonal_noise(equations, variables) Checks whether we deal with diagonal noise, i.e.

diagonal_noise function

(Shortest import: from brian2.stateupdaters.explicit import diagonal_noise)

brian2.stateupdaters.explicit.diagonal_noise (equations, variables)

Checks whether we deal with diagonal noise, i.e. one independent noise variable per variable.
 Raises

UnsupportedEquationsException If the noise is not diagonal.

\[
\text{split_expression} (\text{expr}) \quad \text{Split an expression into a part containing the function } f \text{ and another one containing the function } g.
\]

split_expression function

\texttt{split_expression} (\texttt{expr})

Split an expression into a part containing the function \( f \) and another one containing the function \( g \). Returns a tuple of the two expressions (as sympy expressions).

\textbf{Parameters} \ \texttt{expr} : \texttt{str}

An expression containing references to functions \( f \) and \( g \).

\textbf{Returns} \ (\texttt{non_stochastic}, \texttt{stochastic}) : \texttt{tuple of sympy expressions}

A pair of expressions representing the non-stochastic (containing function-independent terms and terms involving \( f \)) and the stochastic part of the expression (terms involving \( g \) and/or \( dW \)).

**Examples**

\begin{verbatim}
>>> split_expression('dt * __f(__x, __t)')
(dt*__f(__x, __t), None)

>>> split_expression('dt * __f(__x, __t) + __dW * __g(__x, __t)')
(dt*__f(__x, __t), __dW*__g(__x, __t))

>>> split_expression('1/(2*dt**.5)*(__g_support - __g(__x, __t))*(__dW**2)')
(0, __dW**2*__g_support*dt**(-0.5)/2 - __dW**2*dt**(-0.5)*__g(__x, __t)/2)
\end{verbatim}

Objects

euler

Forward Euler state updater

euler object

\texttt{euler} = \texttt{ExplicitStateUpdater} ('''x_new = __dW*__g(__x, __t) + __x + dt*__f(__x, __t)''', \texttt{stochastic}='additive')

Forward Euler state updater

heun

Stochastic Heun method (for multiplicative Stratonovic SDEs with non-diagonal diffusion matrix)

heun object

\texttt{heun} = \texttt{ExplicitStateUpdater} ('''x_new = __dW*__g(__x, __t) + __x + dt*__f(__x, __t)''', \texttt{stochastic}='additive')

Stochastic Heun method (for multiplicative Stratonovic SDEs with non-diagonal diffusion matrix)
brian2.stateupdaters.explicit.heun = ExplicitStateUpdater('''__x_support = __dW*__g(__x, __t) + __x __g_support = __g(__x_support, __t + dt) x_new = 0.5*__dW*(__g_support + __g(__x, __t)) + __x + dt*__f(__x, __t)''', stochastic='multiplicative')

Stochastic Heun method (for multiplicative Stratonovic SDEs with non-diagonal diffusion matrix)

milstein

Derivative-free Milstein method

milstein object

(Shortest import: from brian2 import milstein)

brian2.stateupdaters.explicit.milstein = ExplicitStateUpdater('''__x_support = __x + dt**0.5*__g(__x, __t) + dt*__f(__x, __t) __g_support = ... - __g(__x, __t)) x_new = __dW*__g(__x, __t) + __k + __x + dt*__f(__x, __t)''', stochastic='multiplicative')

Derivative-free Milstein method

rk2

Second order Runge-Kutta method (midpoint method)

rk2 object

(Shortest import: from brian2 import rk2)

brian2.stateupdaters.explicit.rk2 = ExplicitStateUpdater('''__k = dt*__f(__x, __t) x_new = __x + dt*__f(0.5*__k + __x, __t + 0.5*dt)''', stochastic=None)

Second order Runge-Kutta method (midpoint method)

rk4

Classical Runge-Kutta method (RK4)

rk4 object

(Shortest import: from brian2 import rk4)

brian2.stateupdaters.explicit.rk4 = ExplicitStateUpdater('''__k_1 = dt*__f(__x, __t) __k_2 = dt*__f(0.5*__k_1 + __x, __t + 0.5*dt) __k_3 = ... + 0.333333333333333*__k_2 + 0.333333333333333*__k_3 + 0.166666666666667*__k_4 + __x''', stochastic=None)

Classical Runge-Kutta method (RK4)

exponential_euler module

Exported members: exponential_euler

Classes

ExponentialEulerStateUpdater A state updater for conditionally linear equations, i.e.

ExponentialEulerStateUpdater class

(Shortest import: from brian2.stateupdaters.exponential_euler import ExponentialEulerStateUpdater)

class brian2.stateupdaters.exponential_euler.ExponentialEulerStateUpdater Bases: brian2.stateupdaters.base.StateUpdateMethod

A state updater for conditionally linear equations, i.e. equations where each variable only depends linearly on itself (but possibly non-linearly on other variables). Typical Hodgkin-Huxley equations fall into this category, it is therefore the default integration method used in the GENESIS simulator, for example.
Methods

__call__(equations[], variables, method_options)  Generate abstract code from equations.

Details

__call__(equations, variables=None, method_options=None)
Generate abstract code from equations. The method also gets the the variables because some state updaters
have to check whether variable names reflect other state variables (which can change from timestep to
timestep) or are external values (which stay constant during a run) For convenience, this arguments are
optional – this allows to directly see what code a state updater generates for a set of equations by simply
writing euler(eqs), for example.

Parameters equations: Equations
The model equations.
variables: dict, optional
The Variable objects for the model variables.
method_options: dict, optional
Additional options specific to the state updater.

Returns:

———

code: str
The abstract code performing a state update step.

Functions

get_conditionally_linear_system(eqs[], variables=None)  Convert equations into a linear system using sympy.

get_conditionally_linear_system function

(Shortest import: from brian2.stateupdaters.exponential_euler import
get_conditionally_linear_system)

brian2.stateupdaters.exponential_euler.get_conditionally_linear_system(eqs, variables=None)

Convert equations into a linear system using sympy.

Parameters eqs: Equations
The model equations.

Returns coefficients: dict of (sympy expression, sympy expression) tuples
For every variable x, a tuple (M, B) containing the coefficients M and B (as sympy
expressions) for M * x + B
Raises

**ValueError** If one of the equations cannot be converted into a \( M \times x + B \) form.

Examples

```python
>>> from brian2 import Equations
>>> eqs = Equations('...
... dv/dt = (-v + w**2) / tau : 1
... dw/dt = -w / tau : 1
... **")
>>> system = get_conditionally_linear_system(eqs)
>>> print(system['v'])
(-1/tau, w**2.0/tau)
>>> print(system['w'])
(-1/tau, 0)
```

Objects

- **exponential_euler**

  A state updater for conditionally linear equations, i.e.

  exponential_euler object

  *(Shortest import: from brian2 import exponential_euler)*

  brian2.stateupdaters.exponential_euler.exponential_euler = <brian2.stateupdaters.exponential_euler.ExponentialEulerStateUpdater object>

  A state updater for conditionally linear equations, i.e. equations where each variable only depends linearly on itself (but possibly non-linearly on other variables). Typical Hodgkin-Huxley equations fall into this category, it is therefore the default integration method used in the GENESIS simulator, for example.

6.6.14 synapses package

Package providing synapse support.

*Exported members:* Synapses

**parse_synaptic_generator_syntax module**

*Exported members:* parse_synapse_generator

**Functions**

- **handle_range(*args, **kwds)** Checks the arguments/keywords for the range iterator

**handle_range function**

*(Shortest import: from brian2.synapses.parse_synaptic_generator_syntax import handle_range)*

brian2.synapses.parse_synaptic_generator_syntax.handle_range(*args, **kwds)

Checks the arguments/keywords for the range iterator
Should have 1-3 positional arguments.
Returns a dict with keys low, high, step. Default values are low=0, step=1.

handle_sample(*args, **kwds) Checks the arguments/keywords for the sample iterator

handle_sample function

(Shortest import: from brian2.synapses.parse_synaptic_generator_syntax import handle_sample)

brian2.synapses.parse_synaptic_generator_syntax.handle_sample(*args, **kwds)
Checks the arguments/keywords for the sample iterator

Should have 1-3 positional arguments and 1 keyword argument (either p or size).
Returns a dict with keys low, high, step, sample_size, p, size. Default values are low=0, step=1`. Sample size will be either `'random` or `fixed`. In the first case, p will have a value and size will be None (and vice versa for the second case).

parse_synapse_generator function

(Shortest import: from brian2.synapses.parse_synaptic_generator_syntax import parse_synapse_generator)

brian2.synapses.parse_synaptic_generator_syntax.parse_synapse_generator(expr)
Returns a parsed form of a synapse generator expression.

The general form is:

```
    element for iteration_variable in iterator_func(...) 
```

or

```
    element for iteration_variable in iterator_func(...) if if_expression 
```

Returns a dictionary with keys:

- **original_expression** The original expression as a string.
- **element** As above, a string expression.
- **iteration_variable** A variable name, as above.
- **iterator_func** String. Either range or sample.
- **if_expression** String expression or None.
- **iterator_kwds** Dictionary of key/value pairs representing the keywords. See `handle_range` and `handle_sample`.

spikequeue module

The spike queue class stores future synaptic events produced by a given presynaptic neuron group (or postsynaptic for backward propagation in STDP).
Exported members: SpikeQueue

Classes

SpikeQueue class

(Shortest import: from brian2.synapses.spikequeue import SpikeQueue)

class brian2.synapses.spikequeue.SpikeQueue(source_start, source_end)
    Bases: object

Data structure saving the spikes and taking care of delays.

Parameters

source_start: int
    The start of the source indices (for subgroups)

source_end: int
    The end of the source indices (for subgroups)

Notes:

---

**Data structure**: A spike queue is implemented as a 2D array ‘X’ that is circular in the time direction (rows) and dynamic in the events direction (columns). The row index corresponding to the current timestep is ‘currentime’. Each element contains the target synapse index.

**Offsets**: Offsets are used to solve the problem of inserting multiple synaptic events with the same delay. This is difficult to vectorise. If there are n synaptic events with the same delay, these events are given an offset between 0 and n-1, corresponding to their relative position in the data structure.

Attributes

__dt
    The dt used for storing the spikes (will be set in prepare)

__source_end
    The end of the source indices (for subgroups)

__source_start
    The start of the source indices (for subgroups)

currenttime
    The current time (in time steps)

n
    number of events in each time step

Methods
Brian 2 Documentation, Release 2.3

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>advance()</td>
<td>Advances by one timestep</td>
</tr>
<tr>
<td>peek()</td>
<td>Returns the all the synaptic events corresponding to the current time, as an array of synapse indexes.</td>
</tr>
<tr>
<td>prepare()</td>
<td>Prepare the data structures</td>
</tr>
<tr>
<td>push(sources)</td>
<td>Push spikes to the queue.</td>
</tr>
</tbody>
</table>

Details

_\text{dt}_
The dt used for storing the spikes (will be set in \text{prepare})

_\text{source}_\text{end}_
The end of the source indices (for subgroups)

_\text{source}_\text{start}_
The start of the source indices (for subgroups)

\text{current}_\text{time}_
The current time (in time steps)

n
number of events in each time step

\text{advance}()
Advances by one timestep

\text{peek}()
Returns the all the synaptic events corresponding to the current time, as an array of synapse indexes.

\text{prepare}(\text{delays}, \text{dt}, \text{synapse}_\text{sources})
Prepare the data structures

This is called every time the network is run. The size of the of the data structure (number of rows) is adjusted to fit the maximum delay in \text{delays}, if necessary. A flag is set if delays are homogeneous, in which case insertion will use a faster method implemented in \text{insert_homogeneous}.

\text{push}(\text{sources})
Push spikes to the queue.

Parameters \text{sources} : ndarray of int
The indices of the neurons that spiked.

\textit{synapses module}

Exported members: \textit{Synapses}

Classes

\textit{StateUpdater}(\text{group}, \text{method}, \text{clock}, \text{order}[, . . . ])
The CodeRunner that updates the state variables of a \textit{Synapses} at every timestep.

\textit{StateUpdater class}

(\textit{Shortest import}: from brian2.synapses.synapses import \textit{StateUpdater})
class brian2.synapses.synapses.StateUpdater(group, method, clock, order, method_options=None)

Bases: brian2.groups.group.CodeRunner

The CodeRunner that updates the state variables of a Synapses at every timestep.

Methods

update_abstract_code([run_namespace, level])
Update the abstract code for the code object.

Details

update_abstract_code (run_namespace=None, level=0)
Update the abstract code for the code object. Will be called in before_run and should update the CodeRunner.abstract_code attribute.

Does nothing by default.

SummedVariableUpdater class

(Shortest import: from brian2.synapses.synapses import SummedVariableUpdater)

class brian2.synapses.synapses.SummedVariableUpdater(expression, target_varname, synapses, target, target_size_name, index_var)

Bases: brian2.groups.group.CodeRunner

The CodeRunner that updates a value in the target group with the sum over values in the Synapses object.

Synapses class

(Shortest import: from brian2 import Synapses)

class brian2.synapses.synapses.Synapses(source, target=None, model=None, on_pre=None, pre=None, on_post=None, post=None, connect=None, delay=None, on_event='spike', multisynaptic_index=None, namespace=None, dtype=None, codeobj_class=None, dt=None, clock=None, order=0, method=('exact', 'euler', 'heun'), method_options=None, name='synapses*')

Bases: brian2.groups.group.Group

Class representing synaptic connections.

Creating a new Synapses object does by default not create any synapses, you have to call the Synapses.
connect method for that.

**Parameters**

- **source**: SpikeSource
  The source of spikes, e.g. a NeuronGroup.
- **target**: Group, optional
  The target of the spikes, typically a NeuronGroup. If none is given, the same as source()
- **model**: str, Equations, optional
  The model equations for the synapses.
- **on_pre**: str, dict, optional
  The code that will be executed after every pre-synaptic spike. Can be either a single (possibly multi-line) string, or a dictionary mapping pathway names to code strings. In the first case, the pathway will be called pre and made available as an attribute of the same name. In the latter case, the given names will be used as the pathway/attribute names. Each pathway has its own code and its own delays.
- **pre**: str, dict, optional
  Deprecated. Use on_pre instead.
- **on_post**: str, dict, optional
  The code that will be executed after every post-synaptic spike. Same conventions as for on_pre: the default name for the pathway is post.
- **post**: str, dict, optional
  Deprecated. Use on_post instead.
- **delay**: Quantity, dict, optional
  The delay for the “pre” pathway (same for all synapses) or a dictionary mapping pathway names to delays. If a delay is specified in this way for a pathway, it is stored as a single scalar value. It can still be changed afterwards, but only to a single scalar value. If you want to have delays that vary across synapses, do not use the keyword argument, but instead set the delays via the attribute of the pathway, e.g. S.pre.delay = ... (or S.delay = ... as an abbreviation), S.post.delay = .... etc.
- **on_event**: str or dict, optional
  Define the events which trigger the pre and post pathways. By default, both pathways are triggered by the 'spike' event, i.e. the event that is triggered by the threshold condition in the connected groups.
- **multisynaptic_index**: str, optional
  The name of a variable (which will be automatically created) that stores the “synapse number”. This number enumerates all synapses between the same source and target so that they can be distinguished. For models where each source-target pair has only a single connection, this number only wastes memory (it would always default to 0), it is therefore not stored by default. Defaults to None (no variable).
- **namespace**: dict, optional
  A dictionary mapping identifier names to objects. If not given, the namespace will be filled in at the time of the call of Network.run, with either the values from the namespace argument of the Network.run method or from the local context, if no such argument is given.
**dtype**: `dtype`, dict, optional

The `numpy.dtype` that will be used to store the values, or a dictionary specifying the type for variable names. If a value is not provided for a variable (or no value is provided at all), the preference setting `core.default_float_dtype` is used.

**codeobj_class**: class, optional

The `CodeObject` class to use to run code.

**dt**: `Quantity`, optional

The time step to be used for the update of the state variables. Cannot be combined with the `clock` argument.

**clock**: `Clock`, optional

The update clock to be used. If neither a clock, nor the `dt` argument is specified, the `defaultclock` will be used.

**order**: int, optional

The priority of of this group for operations occurring at the same time step and in the same scheduling slot. Defaults to 0.

**method**: str, `StateUpdateMethod`, optional

The numerical integration method to use. If none is given, an appropriate one is automatically determined.

**name**: str, optional

The name for this object. If none is given, a unique name of the form `synapses`, `synapses_1`, etc. will be automatically chosen.

### Attributes

<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
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<tbody>
<tr>
<td><code>_connect_called</code></td>
<td>remember whether connect was called to raise an error if an assignment to a synaptic variable is attempted without a preceding connect.</td>
</tr>
<tr>
<td><code>_pathways</code></td>
<td>List of all <code>SynapticPathway</code> objects</td>
</tr>
<tr>
<td><code>_registered_variables</code></td>
<td>Set of <code>Variable</code> objects that should be resized when the number of synapses changes</td>
</tr>
<tr>
<td><code>_synaptic_updaters</code></td>
<td>List of names of all updaters, e.g.</td>
</tr>
<tr>
<td><code>delay</code></td>
<td>The presynaptic delay (if a pre-synaptic pathway exists).</td>
</tr>
<tr>
<td><code>delay_</code></td>
<td>The presynaptic delay without unit information (if apre-synaptic pathway exists).</td>
</tr>
<tr>
<td><code>events</code></td>
<td>“Events” for all the pathways</td>
</tr>
<tr>
<td><code>namespace</code></td>
<td>The group-specific namespace</td>
</tr>
<tr>
<td><code>state_updater</code></td>
<td>Performs numerical integration step</td>
</tr>
<tr>
<td><code>subexpression_updater</code></td>
<td>Update the “constant over a time step” subexpressions</td>
</tr>
<tr>
<td><code>summed_updaters</code></td>
<td>“Summed variable” mechanism – sum over all synapses of a pre-/postsynaptic target</td>
</tr>
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</table>
## Methods

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>before_run</code><em>(run_namespace)</em></td>
<td>Optional method to prepare the object before a run.</td>
</tr>
<tr>
<td><code>check_variable_write</code>(variable)</td>
<td>Checks that <code>Synapses.connect</code> has been called before setting a synaptic variable.</td>
</tr>
<tr>
<td><code>connect</code>(<strong>kwds</strong>)</td>
<td>Add synapses.</td>
</tr>
<tr>
<td><code>register_variable</code>(variable)</td>
<td>Register a <code>DynamicArray</code> to be automatically resized when the size of the indices change.</td>
</tr>
<tr>
<td><code>unregister_variable</code>(variable)</td>
<td>Unregister a <code>DynamicArray</code> from the automatic resizing mechanism.</td>
</tr>
</tbody>
</table>

## Details

- `_connect_called`
  - remember whether connect was called to raise an error if an assignment to a synaptic variable is attempted without a preceding connect.

- `_pathways`
  - List of all `SynapticPathway` objects

- `_registered_variables`
  - Set of `Variable` objects that should be resized when the number of synapses changes

- `_synaptic_updaters`
  - List of names of all updaters, e.g. ['pre', 'post']

- `delay`
  - The presynaptic delay (if a pre-synaptic pathway exists).

- `delay_`
  - The presynaptic delay without unit information (if apre-synaptic pathway exists).

- `events`
  - “Events” for all the pathways

- `namespace`
  - The group-specific namespace

- `state_updater`
  - Performs numerical integration step

- `subexpression_updater`
  - Update the “constant over a time step” subexpressions

- `summed_updaters`
  - “Summed variable” mechanism – sum over all synapses of a pre-/postsynaptic target

- `before_run`*(run_namespace)*
  - Optional method to prepare the object before a run.

  TODO

- `check_variable_write`(variable)
  - Checks that `Synapses.connect` has been called before setting a synaptic variable.

  **Parameters**

  - `variable`: `Variable`

    The variable that the user attempts to set.
_raises

**TypeError** If `Synapses.connect` has not been called yet.

```python
connect(**kwds)
```

Add synapses.

See `Synapses` for details.

Parameters

- **condition** : str, bool, optional
  A boolean or string expression that evaluates to a boolean. The expression can depend on indices `i` and `j` and on pre- and post-synaptic variables. Can be combined with arguments `n`, and `p` but not `i` or `j`.

- **i** : int, ndarray of int, optional
  The presynaptic neuron indices (in the form of an index or an array of indices). Must be combined with `j` argument.

- **j** : int, ndarray of int, str, optional
  The postsynaptic neuron indices. It can be an index or array of indices if combined with the `i` argument, or it can be a string generator expression.

- **p** : float, str, optional
  The probability to create `n` synapses wherever the `condition` evaluates to true. Cannot be used with generator syntax for `j`.

- **n** : int, str, optional
  The number of synapses to create per pre/post connection pair. Defaults to 1.

- **skip_if_invalid** : bool, optional
  If set to True, rather than raising an error if you try to create an invalid/out of range pair `(i, j)` it will just quietly skip those synapses.

- **namespace** : dict-like, optional
  A namespace that will be used in addition to the group-specific namespaces (if defined). If not specified, the locals and globals around the run function will be used.

- **level** : int, optional
  How deep to go up the stack frame to look for the locals/global (see `namespace` argument).

Examples

```python
>>> from brian2 import *
>>> import numpy as np

>>> G = NeuronGroup(10, 'dv/dt = -v / tau : 1', threshold='v>1', reset='v=0')
>>> S = Synapses(G, G, 'w:1', on_pre='v+=w')
>>> S.connect(condition='i != j')  # all-to-all but no self-connections
>>> S.connect(i=0, j=0)  # connect neuron 0 to itself
>>> S.connect(i=np.array([1, 2]), j=np.array([2, 1]))  # connect 1->2 and 2->1
>>> S.connect()  # connect all-to-all
>>> S.connect(condition='i != j', p=0.1)  # Connect neurons with 10% probability, exclude self-connections
```

(continues on next page)
register_variable (variable)
Register a DynamicArray to be automatically resized when the size of the indices change. Called automatically when a SynapticArrayVariable specifier is created.

unregister_variable (variable)
Unregister a DynamicArray from the automatic resizing mechanism.

Tutorials and examples using this

- Tutorial 2-intro-to-brian-synapses
- Tutorial 3-intro-to-brian-simulations
- Tutorial 1-intro-to-brian-neurons
- Example COBAAH
- Example CUBA
- Example adaptive_threshold
- Example compartmental/bipolar_with_inputs
- Example compartmental/bipolar_with_inputs2
- Example compartmental/lfp
- Example advanced/float_32_64_benchmark
- Example advanced/custom_events
- Example synapses/licklider
- Example synapses/efficient_gaussian_connectivity
- Example synapses/gapjunctions
- Example synapses/nonlinear
- Example synapses/synapses
- Example synapses/state_variables
- Example synapses/jeffress
- Example synapses/spatial_connections
- Example synapses/STDP
- Example standalone/cuba_openmp
- Example standalone/STDP_standalone
- Example frompapers/Diesmann_et_al_1999
- Example frompapers/Clopath_et_al_2010_homeostasis
SynapticIndexing class

(Shortest import: from brian2.synapses.synapses import SynapticIndexing)

class brian2.synapses.synapses.SynapticIndexing(synapses)
    Bases: object

    Methods

    __call__(index, index_var)
    Returns synaptic indices for index, which can be a tuple of indices (including arrays and slices), a single index or a string.

    Details

    __call__(index=None, index_var='idx')
    Returns synaptic indices for index, which can be a tuple of indices (including arrays and slices), a single index or a string.
**SynapticPathway**(synapses, code, prepost[,...])  
The CodeRunner that applies the pre/post statement(s) to the state variables of synapses where the pre-/postsynaptic group spiked in this time step.

---

**SynapticPathway class**

*(Shortest import: from brian2.synapses.synapses import SynapticPathway)*

class brian2.synapses.synapses.SynapticPathway(synapses, code, prepost, objname=None, delay=None, event='spike')

Bases: brian2.groups.group.CodeRunner, brian2.groups.group.Group

The CodeRunner that applies the pre/post statement(s) to the state variables of synapses where the pre-/postsynaptic group spiked in this time step.

**Parameters**

synapses: Synapses  
Reference to the main Synapses object

prepost: {'pre', 'post'}  
Whether this object should react to pre- or postsynaptic spikes

objname: str, optional  
The name to use for the object, will be appendend to the name of synapses to create a name in the sense of Nameable. If None is provided (the default), prepost will be used.

delay: Quantity, optional  
A scalar delay (same delay for all synapses) for this pathway. If not given, delays are expected to vary between synapses.

**Attributes**

_initialise_queue_codeobj  
The CodeObject initialising the SpikeQueue at the begin of a run

queue  
The SpikeQueue

**Methods**

before_run(**kwds)

check_variable_write(variable)  
Function that can be overwritten to raise an error if writing to a variable should not be allowed.

initialise_queue()

push_spikes()

update_abstract_code(**kwds)

**Details**

_initialise_queue_codeobj  
The CodeObject initialising the SpikeQueue at the begin of a run

queue
The SpikeQueue

```python
before_run(**kwds)
```

```python
check_variable_write(variable)
```

Function that can be overwritten to raise an error if writing to a variable should not be allowed. Note that this does not deal with incorrect writes that are general to all kind of variables (incorrect units, writing to a read-only variable, etc.). This function is only used for type-specific rules, e.g. for raising an error in Synapses when writing to a synaptic variable before any connect call.

By default this function does nothing.

**Parameters**

- `variable`: Variable
  The variable that the user attempts to set.

```python
initialise_queue()
push_spikes()
update_abstract_code(**kwds)
```

---

### SynapticSubgroup class

(Shortest import: from brian2.synapses.synapses import SynapticSubgroup)

```python
class brian2.synapses.synapses.SynapticSubgroup(synapses, indices)
```

Bases: object

A simple subgroup of Synapses that can be used for indexing.

**Parameters**

- `indices`: ndarray of int
  The synaptic indices represented by this subgroup.

- `synaptic_pre`: DynamicArrayVariable
  References to all pre-synaptic indices. Only used to throw an error when new synapses where added after creating this object.

**Functions**

```python
find_synapses(index, synaptic_neuron)
```

---

### find_synapses function

(Shortest import: from brian2.synapses.synapses import find_synapses)

```python
brian2.synapses.synapses.find_synapses(index, synaptic_neuron)
```

**slice_to_test(x)**

Returns a testing function corresponding to whether an index is in slice x.
slice_to_test function

(Shortest import: from brian2.synapses.synapses import slice_to_test)

brian2.synapses.synapses.slice_to_test(x)
Returns a testing function corresponding to whether an index is in slice x. x can also be an int.

6.6.15 units package

The unit system.

Exported members: pamp, namp, uamp, mamp, amp, kamp, Mamp, Gamp, Tamp, kelvin, kilogram, pmetre, nmetre, umetre, mmetre, metre, kmetre, Mmetre, Gmetre, Tmetre, pmeter, nmeter, umeter, mmeter, meter ... (218 more members)

allunits module

THIS FILE IS AUTOMATICALLY GENERATED BY A STATIC CODE GENERATION TOOL DO NOT EDIT BY HAND
Instead edit the template:

dev/tools/static_codegen/unitstemplate.py

Exported members: metre, meter, kilogram, second, amp, amper, kelvin, mole, mol, candle, kilogramme, gram, gramme, molar, radian, steradian, hertz, newton, pascal, joule, watt, coulomb, volt, farad, ohm ... (2045 more members)

Objects

celsius A dummy object to raise errors when celsius is used.

celsius object

(Shortest import: from brian2.units.allunits import celsius)

brian2.units.allunits.celsius = <brian2.units.allunits._Celsius object>
A dummy object to raise errors when celsius is used. The use of celsius can lead to ambiguities when mixed with temperatures in kelvin, so its use is no longer supported. See github issue #817 for details.

constants module

A module providing some physical units as Quantity objects. Note that these units are not imported by wildcard imports (e.g. from brian2 import *), they have to be imported explicitly. You can use import ... as ... to import them with shorter names, e.g.:

```
from brian2.units.constants import faraday_constant as F
```
The available constants are:
<table>
<thead>
<tr>
<th>Constant</th>
<th>Symbol(s)</th>
<th>Brian name</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Avogadro constant</td>
<td>$N_A, L$</td>
<td><code>avogadro_constant</code></td>
<td>$6.022140857 \times 10^{23} \text{ mol}^{-1}$</td>
</tr>
<tr>
<td>Boltzmann constant</td>
<td>$k$</td>
<td><code>boltzmann_constant</code></td>
<td>$1.38064852 \times 10^{-23} \text{ J K}^{-1}$</td>
</tr>
<tr>
<td>Electric constant</td>
<td>$\epsilon_0$</td>
<td><code>electric_constant</code></td>
<td>$8.854187817 \times 10^{-12} \text{ F m}^{-1}$</td>
</tr>
<tr>
<td>Electron mass</td>
<td>$m_e$</td>
<td><code>electron_mass</code></td>
<td>$9.10938356 \times 10^{-31} \text{ kg}$</td>
</tr>
<tr>
<td>Elementary charge</td>
<td>$e$</td>
<td><code>elementary_charge</code></td>
<td>$1.6021766208 \times 10^{-19} \text{ C}$</td>
</tr>
<tr>
<td>Faraday constant</td>
<td>$F$</td>
<td><code>faraday_constant</code></td>
<td>$96485.33289 \text{ C mol}^{-1}$</td>
</tr>
<tr>
<td>Gas constant</td>
<td>$R$</td>
<td><code>gas_constant</code></td>
<td>$8.3144598 \text{ J mol}^{-1} \text{ K}^{-1}$</td>
</tr>
<tr>
<td>Magnetic constant</td>
<td>$\mu_0$</td>
<td><code>magnetic_constant</code></td>
<td>$12.566370614 \times 10^{-7} \text{ N A}^{-2}$</td>
</tr>
<tr>
<td>Molar mass constant</td>
<td>$M_u$</td>
<td><code>molar_mass_constant</code></td>
<td>$1 \times 10^{-3} \text{ kg mol}^{-1}$</td>
</tr>
<tr>
<td>0°C</td>
<td></td>
<td><code>zero_celsius</code></td>
<td>$273.15 \text{ K}$</td>
</tr>
</tbody>
</table>

**fundamentalunits module**

Exported members: `DimensionMismatchError`, `get_or_create_dimension()`, `get_dimensions()`, `is_dimensionless()`, `have_same_dimensions()`, `in_unit()`, `in_best_unit()`, `Quantity`, `Unit`, `register_new_unit()`, `check_units()`, `is_scalar_type()`, `get_unit()`

**Classes**

```python
Dimension(dims) Stores the indices of the 7 basic SI unit dimension (length, mass, etc.).
```

**Dimension class**

(Shortest import: from brian2.units.fundamentalunits import Dimension)

```python
class brian2.units.fundamentalunits.Dimension(dims)
Bases: object
```

Stores the indices of the 7 basic SI unit dimension (length, mass, etc.). Provides a subset of arithmetic operations appropriate to dimensions: multiplication, division and powers, and equality testing.

**Parameters**

- **dims**: sequence of `float`  
  The dimension indices of the 7 basic SI unit dimensions.

**Notes**

Users shouldn’t use this class directly, it is used internally in Quantity and Unit. Even internally, never use `Dimension(...)` to create a new instance, use `get_or_create_dimension()` instead. This function makes sure that only one Dimension instance exists for every combination of indices, allowing for a very fast dimensionality check with `is`.

**Attributes**

```python
dim Returns the Dimension object itself.
```

Continued on next page
Table 464 – continued from previous page

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>is_dimensionless</strong></td>
<td>Whether this Dimension is dimensionless.</td>
</tr>
</tbody>
</table>

**Methods**

```
get_dimension(d)        # Return a specific dimension.
```

**Details**

- **dim**
  Returns the `Dimension` object itself. This can be useful, because it allows to check for the dimension of an object by checking its `dim` attribute – this will return a `Dimension` object for `Quantity`, `Unit` and `Dimension`.

- **is_dimensionless**
  Whether this Dimension is dimensionless.

**Notes**

Normally, instead one should check dimension for being identical to `DIMENSIONLESS`.

```
get_dimension(d)        # Return a specific dimension.
```

**Parameters**

- **d**: str
  A string identifying the SI basic unit dimension. Can be either a description like “length” or a basic unit like “m” or “metre”.

**Returns**

- **dim**: float
  The dimensionality of the dimension `d`.

```
DimensionMismatchError(description, *dims)  # Exception class for attempted operations with inconsistent dimensions.
```

**DimensionMismatchError class**

*(Shortest import: from brian2 import DimensionMismatchError)*

```
class brian2.units.fundamentalunits.DimensionMismatchError (description, *dims)
```

**Bases:** Exception

Exception class for attempted operations with inconsistent dimensions.

For example, `3*mvolt + 2*amp` raises this exception. The purpose of this class is to help catch errors based on incorrect units. The exception will print a representation of the dimensions of the two inconsistent objects that were operated on.

**Parameters**

- **description**: str
  A description of the type of operation being performed, e.g. Addition, Multiplication, etc.

- **dims**: Dimension
The physical dimensions of the objects involved in the operation, any number of them is possible

Tutorials and examples using this

- Tutorial 1-intro-to-brian-neurons

<table>
<thead>
<tr>
<th>Quantity</th>
<th>A number with an associated physical dimension.</th>
</tr>
</thead>
</table>

Quantity class

(Shortest import: from brian2 import Quantity)

class brian2.units.fundamentalunits.Quantity
Bases: numpy.ndarray, object

A number with an associated physical dimension. In most cases, it is not necessary to create a Quantity object by hand, instead use multiplication and division of numbers with the constant unit names second, kilogram, etc.

See also:
Unit

Notes

The Quantity class defines arithmetic operations which check for consistency of dimensions and raise the DimensionMismatchError exception if they are inconsistent. It also defines default and other representations for a number for printing purposes.

See the documentation on the Unit class for more details about the available unit names like mvolt, etc.

Casting rules

The rules that define the casting operations for Quantity object are:

1. Quantity op Quantity = Quantity Performs dimension checking if appropriate
2. (Scalar or Array) op Quantity = Quantity Assumes that the scalar or array is dimensionless

There is one exception to the above rule, the number 0 is interpreted as having “any dimension”.

Examples

```python
>>> from brian2 import *
>>> I = 3 * amp  # I is a Quantity object
>>> R = 2 * ohm  # same for R
>>> I * R
6. * volt
>>> (I * R).in_unit(mvolt)
'6000. mV'
>>> (I * R) / mvolt
6000.0
>>> X = I + R  # doctest: +IGNORE_EXCEPTION_DETAIL
Traceback (most recent call last):
```
... DimensionMismatchError: Addition, dimensions were (A) (m^2 kg s^-3 A^-2)
```python
>>> Is = np.array([1, 2, 3]) * amp
>>> Is * R
array([ 2., 4., 6.]) * volt
>>> np.asarray(Is * R) # gets rid of units
array([ 2., 4., 6.])
```

Attributes

<table>
<thead>
<tr>
<th>attribute</th>
<th>description</th>
</tr>
</thead>
<tbody>
<tr>
<td>dimensions</td>
<td>The physical dimensions of this quantity.</td>
</tr>
<tr>
<td>is_dimensionless</td>
<td>Whether this is a dimensionless quantity.</td>
</tr>
<tr>
<td>dim</td>
<td>The physical dimensions of this quantity.</td>
</tr>
</tbody>
</table>

Methods

<table>
<thead>
<tr>
<th>method</th>
<th>description</th>
</tr>
</thead>
<tbody>
<tr>
<td>with_dimensions(value, *args, **keywords)</td>
<td>Create a Quantity object with dim.</td>
</tr>
<tr>
<td>has_same_dimensions(other)</td>
<td>Return whether this object has the same dimensions as another.</td>
</tr>
<tr>
<td>in_unit(u[, precision, python_code])</td>
<td>Represent the quantity in a given unit.</td>
</tr>
<tr>
<td>in_best_unit([precision, python_code])</td>
<td>Represent the quantity in the “best” unit.</td>
</tr>
</tbody>
</table>

Details

- **dimensions**: The physical dimensions of this quantity.
- **is_dimensionless**: Whether this is a dimensionless quantity.
- **dim**: The physical dimensions of this quantity.

**static with_dimensions(value, *args, **keywords)**
Create a `Quantity` object with dim.

Parameters:
- **value**: {array_like, number}
The value of the dimension
- **args**: `{Dimension, sequence of float}`
Either a single argument (a `Dimension`) or a sequence of 7 values.
- **kwds**: Keywords defining the dim, see `Dimension` for details.

Returns:
- **q**: `Quantity`
A `Quantity` object with the given dim
Examples

All of these define an equivalent Quantity object:

```python
>>> from brian2 import *
>>> Quantity.with_dimensions(2, get_or_create_dimension(length=1))
2.*metre
>>> Quantity.with_dimensions(2, length=1)
2.*metre
>>> 2 * metre
2.*metre
```

`has_same_dimensions(other)`
Return whether this object has the same dimensions as another.

**Parameters**

other : \{Quantity, array-like, number\}
The object to compare the dimensions against.

**Returns**

same : bool
True if other has the same dimensions.

`in_unit(u, precision=None, python_code=False)`
Represent the quantity in a given unit. If python_code is True, this will return valid python code, i.e. a string like 5.0 * um ** 2 instead of 5.0 um^2

**Parameters**

u : \{Quantity, Unit\}
The unit in which to show the quantity.

precision : int, optional
The number of digits of precision (in the given unit, see Examples). If no value is given, numpy's get_printoptions() value is used.

python_code : bool, optional
Whether to return valid python code (True) or a human readable string (False, the default).

**Returns**

s : str
String representation of the object in unit u.

See also:

`in_unit()`

Examples

```python
>>> from brian2.units import *
>>> from brian2.units.stdunits import *
>>> x = 25.123456 * mV
>>> x.in_unit(volt)
'0.02512346 V'
>>> x.in_unit(volt, 3)
'0.025 V'
>>> x.in_unit(mV, 3)
'25.123 mV'
```
in_best_unit (precision=None, python_code=False, *regs)

Represent the quantity in the “best” unit.

Parameters

- **python_code**: bool, optional
  
  If set to False (the default), will return a string like `5.0 \text{ um}^2` which is not a valid Python expression. If set to True, it will return `5.0 \times \text{ um} \times 2` instead.

- **precision**: int, optional
  
  The number of digits of precision (in the best unit, see Examples). If no value is given, numpy’s `get_printoptions()` value is used.

- **regs**: `UnitRegistry` objects
  
  The registries where to search for units. If none are given, the standard, user-defined and additional registries are searched in that order.

Returns

- **representation**: str
  
  A string representation of this `Quantity`.

See also:

- `in_best_unit()`

Examples

```python
>>> from brian2.units import *

>>> x = 0.00123456 * volt

>>> x.in_best_unit()
'1.23456 \text{ mV}'

>>> x.in_best_unit(3)
'1.235 \text{ mV}'
```

Tutorials and examples using this

- Example frompapers/Destexhe_et_al_1998
- Example frompapers/Platkiewicz_Brette_2011
- Example frompapers/Stimberg_et_al_2018/example_1_COBA

Unit (value[, dim, scale, name, dispname, ...])

A physical unit.

Unit class

(Shortest import: from brian2 import Unit)

```python
class brian2.units.fundamentalunits.Unit (value, dim=None, scale=0, name=None, dispname=None, latexname='', iscompound=False)
```
A physical unit.

Normally, you do not need to worry about the implementation of units. They are derived from the `Quantity` object with some additional information (name and string representation).

Basically, a unit is just a number with given dimensions, e.g. mV = 0.001 with the dimensions of voltage. The units module defines a large number of standard units, and you can also define your own (see below).

The unit class also keeps track of various things that were used to define it so as to generate a nice string representation of it. See below.

When creating scaled units, you can use the following prefixes:

<table>
<thead>
<tr>
<th>Factor</th>
<th>Name</th>
<th>Prefix</th>
</tr>
</thead>
<tbody>
<tr>
<td>$10^{24}$</td>
<td>yotta</td>
<td>Y</td>
</tr>
<tr>
<td>$10^{21}$</td>
<td>zetta</td>
<td>Z</td>
</tr>
<tr>
<td>$10^{18}$</td>
<td>exa</td>
<td>E</td>
</tr>
<tr>
<td>$10^{15}$</td>
<td>peta</td>
<td>P</td>
</tr>
<tr>
<td>$10^{12}$</td>
<td>tera</td>
<td>T</td>
</tr>
<tr>
<td>$10^{9}$</td>
<td>giga</td>
<td>G</td>
</tr>
<tr>
<td>$10^{6}$</td>
<td>mega</td>
<td>M</td>
</tr>
<tr>
<td>$10^{3}$</td>
<td>kilo</td>
<td>k</td>
</tr>
<tr>
<td>$10^{2}$</td>
<td>hecto</td>
<td>h</td>
</tr>
<tr>
<td>$10^{1}$</td>
<td>deka</td>
<td>da</td>
</tr>
<tr>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$10^{-1}$</td>
<td>deci</td>
<td>d</td>
</tr>
<tr>
<td>$10^{-2}$</td>
<td>centi</td>
<td>c</td>
</tr>
<tr>
<td>$10^{-3}$</td>
<td>milli</td>
<td>m</td>
</tr>
<tr>
<td>$10^{-6}$</td>
<td>micro</td>
<td>µ (mu in SI)</td>
</tr>
<tr>
<td>$10^{-9}$</td>
<td>nano</td>
<td>n</td>
</tr>
<tr>
<td>$10^{-12}$</td>
<td>pico</td>
<td>p</td>
</tr>
<tr>
<td>$10^{-15}$</td>
<td>femto</td>
<td>f</td>
</tr>
<tr>
<td>$10^{-18}$</td>
<td>atto</td>
<td>a</td>
</tr>
<tr>
<td>$10^{-21}$</td>
<td>zepto</td>
<td>z</td>
</tr>
<tr>
<td>$10^{-24}$</td>
<td>yocto</td>
<td>y</td>
</tr>
</tbody>
</table>

**Defining your own**

It can be useful to define your own units for printing purposes. So for example, to define the newton metre, you write

```python
>>> from brian2 import *
>>> from brian2.units.allunits import newton
>>> Nm = newton * metre
```

You can then do

```python
>>> (1*Nm).in_unit(Nm)
'1. N m'
```

New “compound units”, i.e. units that are composed of other units will be automatically registered and from then on used for display. For example, imagine you define total conductance for a membrane, and the total area of that membrane:
If you now ask for the conductance density, you will get an “ugly” display in basic SI dimensions, as Brian does not know of a corresponding unit:

```python
>>> conductance/area
0.5 * metre ** -4 * kilogram ** -1 * second ** -3 * amp ** 2
```

By using an appropriate unit once, it will be registered and from then on used for display when appropriate:

```python
>>> usiemens/cm**2
usiemens / cmetre ** 2
>>> conductance/area  # same as before, but now Brian knows about uS/cm^2
50. * usiemens / cmetre ** 2
```

Note that user-defined units cannot override the standard units (volt, second, etc.) that are predefined by Brian. For example, the unit Nm has the dimensions “length²·mass/time²”, and therefore the same dimensions as the standard unit joule. The latter will be used for display purposes:

```python
>>> 3*joule
3. * joule
>>> 3*Nm
3. * joule
```

### Attributes

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>_dispname</code></td>
<td>The display name of this unit.</td>
</tr>
<tr>
<td><code>_latexname</code></td>
<td>A LaTeX expression for the name of this unit.</td>
</tr>
<tr>
<td><code>_name</code></td>
<td>The full name of this unit.</td>
</tr>
<tr>
<td><code>dim</code></td>
<td>The Dimensions of this unit</td>
</tr>
<tr>
<td><code>dispname</code></td>
<td>The display name of the unit</td>
</tr>
<tr>
<td><code>iscompound</code></td>
<td>Whether this unit is a combination of other units.</td>
</tr>
<tr>
<td><code>latexname</code></td>
<td>The LaTeX name of the unit</td>
</tr>
<tr>
<td><code>name</code></td>
<td>The name of the unit</td>
</tr>
<tr>
<td><code>scale</code></td>
<td>The scale for this unit (as the integer exponent of 10), i.e.</td>
</tr>
</tbody>
</table>

### Methods

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>create(dim, name, dispname[, latexname, scale])</code></td>
<td>Create a new named unit.</td>
</tr>
<tr>
<td><code>create_scaled_unit(baseunit, scalefactor)</code></td>
<td>Create a scaled unit from a base unit.</td>
</tr>
<tr>
<td><code>set_display_name(name)</code></td>
<td>Sets the display name for the unit.</td>
</tr>
<tr>
<td><code>set_latex_name(name)</code></td>
<td>Sets the LaTeX name for the unit.</td>
</tr>
<tr>
<td><code>set_name(name)</code></td>
<td>Sets the name for the unit.</td>
</tr>
</tbody>
</table>

### Details

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>_dispname</code></td>
<td>The display name of this unit.</td>
</tr>
</tbody>
</table>
_\text{name}
   A LaTeX expression for the name of this unit.

_\text{dim}
   The full name of this unit.

dim
   The Dimensions of this unit

dispname
   The display name of the unit

\text{iscompound}
   Whether this unit is a combination of other units.

latexname
   The LaTeX name of the unit

name
   The name of the unit

scale
   The scale for this unit (as the integer exponent of 10), i.e. a scale of 3 means $10^3$, e.g. for a “k” prefix.

\text{static create}(\text{dim}, \text{name}, \text{dispname}, \text{latexname} = None, \text{scale} = 0)
   Create a new named unit.

   \text{Parameters dim} : \text{Dimension}
      The dimensions of the unit.

   \text{name} : \text{str}
      The full name of the unit, e.g. 'volt'

   \text{dispname} : \text{str}
      The display name, e.g. 'V'

   \text{latexname} : \text{str}, optional
      The name as a LaTeX expression (math mode is assumed, do not add $ signs or similar), e.g. '\omega'. If no \text{latexname} is specified, \text{dispname} will be used.

   \text{scale} : \text{int}, optional
      The scale of this unit as an exponent of 10, e.g. -3 for a unit that is 1/1000 of the base scale. Defaults to 0 (i.e. a base unit).

   \text{Returns u} : \text{Unit}
      The new unit.

\text{static create_scaled_unit}(\text{baseunit}, \text{scalefactor})
   Create a scaled unit from a base unit.

   \text{Parameters baseunit} : \text{Unit}
      The unit of which to create a scaled version, e.g. volt, amp.

   \text{scalefactor} : \text{str}
      The scaling factor, e.g. "m" for mvolt, mamp

   \text{Returns u} : \text{Unit}
      The new unit.
**set_display_name**(name)
Sets the display name for the unit.
Depreciated since version 2.1: Create a new unit with `Unit.create` instead.

**set_latex_name**(name)
Sets the LaTeX name for the unit.
Depreciated since version 2.1: Create a new unit with `Unit.create` instead.

**set_name**(name)
Sets the name for the unit.
Depreciated since version 2.1: Create a new unit with `Unit.create` instead.

---

**UnitRegistry**()
Stores known units for printing in best units.

---

**UnitRegistry class**

(_Shortest import: from brian2.units.fundamentalunits import UnitRegistry_)

class brian2.units.fundamentalunits.UnitRegistry
    Bases: object
    Stores known units for printing in best units.
    All a user needs to do is to use the `register_new_unit()` function.
    Default registries:
    The units module defines three registries, the standard units, user units, and additional units. Finding best units is done by first checking standard, then user, then additional. New user units are added by using the `register_new_unit()` function.
    Standard units includes all the basic non-compound unit names built in to the module, including volt, amp, etc. Additional units defines some compound units like newton metre (Nm) etc.

**Methods**

<table>
<thead>
<tr>
<th>Method</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>add(u)</code></td>
<td>Add a unit to the registry</td>
</tr>
<tr>
<td><code>__getitem__(x)</code></td>
<td>Returns the best unit for quantity x</td>
</tr>
</tbody>
</table>

**Details**

**add**(u)
Add a unit to the registry

**__getitem__**(x)
Returns the best unit for quantity x

The algorithm is to consider the value:

\[
m = \frac{\text{abs}(x/u)}
\]

for all matching units u. We select the unit where this ratio is the closest to 10 (if it is an array with several values, we select the unit where the deviations from that are the smallest. More precisely, the unit that minimizes the sum of \((\log_{10}(m) - 1)^2\) over all entries).

**Functions**
check_units function

(Shortest import: from brian2 import check_units)

Decorator to check units of arguments passed to a function

Raises

DimensionMismatchError  In case the input arguments or the return value do not have the expected dimensions.

TypeError  If an input argument or return value was expected to be a boolean but is not.

Notes

This decorator will destroy the signature of the original function, and replace it with the signature (*args, **kwds). Other decorators will do the same thing, and this decorator critically needs to know the signature of the function it is acting on, so it is important that it is the first decorator to act on a function. It cannot be used in combination with another decorator that also needs to know the signature of the function.

Note that the bool type is “strict”, i.e. it expects a proper boolean value and does not accept 0 or 1. This is not the case the other way round, declaring an argument or return value as “1” does allow for a True or False value.

Examples

```python
>>> from brian2.units import *
>>> @check_units(I=amp, R=ohm, wibble=metre, result=volt)
... def getvoltage(I, R, **k):
...     return I*R
... getvoltage(1*amp, 1*ohm, wibble=1)
Traceback (most recent call last):
  ... DimensionMismatchError: Function "getvoltage" variable "wibble" has wrong dimensions, dimensions were (1) (m)
```

You don’t have to check the units of every variable in the function, and you can define what the units should be for variables that aren’t explicitly named in the definition of the function. For example, the code above checks that the variable wibble should be a length, so writing

```python
>>> getvoltage(1*amp, 1*ohm, wibble=1)  # doctest: +IGNORE_EXCEPTION_DETAIL
```

fails, but

```python
>>> getvoltage(1*amp, 1*ohm, wibble=1*metre)
1. * volt
```

passes. String arguments or None are not checked
By using the special name \texttt{result}, you can check the return value of the function.

You can also use \texttt{1} or \texttt{bool} as a special value to check for a unitless number or a boolean value, respectively:

\begin{verbatim}
>>> @check_units(value=1, absolute=bool, result=bool) ... def is_high(value, absolute=False): ... if absolute: ... return abs(value) >= 5 ... else: ... return value >= 5
\end{verbatim}

This will then again raise an error if the argument if not of the expected type: >>> is_high(7) True >>> is_high(-7. True) True >>> is_high(3, 4) # doctest: +IGNORE_EXCEPTION_DETAIL Traceback (most recent call last): … TypeError: Function “is_high” expected a boolean value for argument “absolute” but got 4.

\begin{verbatim}
fail_for_dimension_mismatch(obj1[, obj2, …])
\end{verbatim}

\textbf{fail\_for\_dimension\_mismatch function}

\textit{(Shortest import): from brian2.units.fundamentalunits import fail_for_dimension_mismatch}

\begin{verbatim}
brian2.units.fundamentalunits.fail_for_dimension_mismatch(obj1[, obj2=None, …], error_message=None, **error_quantities)
\end{verbatim}

Compare the dimensions of two objects.

\textbf{Parameters} \texttt{obj1}, \texttt{obj2} : {array-like, \texttt{Quantity}}

The object to compare. If \texttt{obj2} is \texttt{None}, assume it to be dimensionless

\texttt{error\_message} : str, optional

An error message that is used in the DimensionMismatchError

\texttt{error\_quantities} : dict mapping str to \texttt{Quantity}, optional

Quantities in this dictionary will be converted using the \texttt{\_short\_str} helper method and inserted into the \texttt{error\_message} (which should have placeholders with the corresponding names). The reason for doing this in a somewhat complicated way instead of directly including all the details in \texttt{error\_messsage} is that converting large quantity arrays to strings can be rather costly and we don’t want to do it if no error occured.

\textbf{Returns} \texttt{dim1, dim2} : \texttt{Dimension, Dimension}

The physical dimensions of the two arguments (so that later code does not need to get the dimensions again).

\textbf{Raises}

\texttt{DimensionMismatchError} If the dimensions of \texttt{obj1} and \texttt{obj2} do not match (or, if \texttt{obj2} is \texttt{None}, in case \texttt{obj1} is not dimensionless).

\textbf{Notes}

Implements special checking for \texttt{0}, treating it as having “any dimensions”.
**get_dimensions** function

* (Shortest import: from brian2 import get_dimensions)

```
get_dimensions(obj)  # Return the dimensions of any object that has them.
```

Slightly more general than `Quantity.dimensions` because it will return `DIMENSIONLESS` if the object is of number type but not a `Quantity` (e.g. a `float` or `int`).

- **Parameters**
  - `obj`: object
    - The object to check.

- **Returns**
  - `dim`: 'Dimension' :
    - The physical dimensions of the `obj`.

**get_or_create_dimension** function

* (Shortest import: from brian2 import get_or_create_dimension)

```
get_or_create_dimension(*args, **kwds)  # Create a new Dimension object or get a reference to an existing one.
```

Create a new Dimension object or get a reference to an existing one. This function takes care of only creating new objects if they were not created before and otherwise returning a reference to an existing object. This allows to compare dimensions very efficiently using `is`.

- **Parameters**
  - `args`: sequence of `float`
    - A sequence with the indices of the 7 elements of an SI dimension.
  - `kwds`: keyword arguments
    - a sequence of `keyword=value` pairs where the keywords are the names of the SI dimensions, or the standard unit.

**Notes**

The 7 units are (in order):
- Length, Mass, Time, Electric Current, Temperature, Quantity of Substance, Luminosity

and can be referred to either by these names or their SI unit names, e.g. `length`, `metre`, and `m` all refer to the same thing here.

**Examples**

The following are all definitions of the dimensions of force
>>> from brian2 import *
>>> get_or_create_dimension(length=1, mass=1, time=-2)
metre * kilogram * second ** -2
>>> get_or_create_dimension(m=1, kg=1, s=-2)
metre * kilogram * second ** -2
>>> get_or_create_dimension([1, 1, -2, 0, 0, 0, 0])
metre * kilogram * second ** -2

g**et_unit**(d)
Find an unscaled unit (e.g.

get_unit function

(*Shortest import: from brian2 import get_unit*)
brian2.units.fundamentalunits.get_unit(d)

Find an unscaled unit (e.g. volt but not mvolt) for a Dimension.

Parameters d : Dimension
The dimension to find a unit for.

Returns u : Unit
A registered unscaled Unit for the dimensions d, or a new Unit if no unit was found.

g**et_unit_for_display**(d)
Return a string representation of an appropriate unscaled unit or '1' for a dimensionless quantity.

get_unit_for_display function

(*Shortest import: from brian2.units.fundamentalunits import get_unit_for_display*)
brian2.units.fundamentalunits.get_unit_for_display(d)

Return a string representation of an appropriate unscaled unit or '1' for a dimensionless quantity.

Parameters d : Dimension
The dimension to find a unit for.

Returns s : str
A string representation of the respective unit or the string '1'.

have_same_dimensions(obj1, obj2)
Test if two values have the same dimensions.

have_same_dimensions function

(*Shortest import: from brian2 import have_same_dimensions*)
brian2.units.fundamentalunits.have_same_dimensions(obj1, obj2)

Test if two values have the same dimensions.

Parameters obj1, obj2 : {Quantity, array-like, number}
The values of which to compare the dimensions.
Returns same: bool
True if obj1 and obj2 have the same dimensions.

\texttt{in\_best\_unit(x[, \text{precision}])} Represent the value in the “best” unit.

**in\_best\_unit function**

(*Shortest import: from brian2 import in\_best\_unit*)

\texttt{brian2\_units\_fundamentalunits.\texttt{in\_best\_unit}(x, \text{precision=None})}

Represent the value in the “best” unit.

**Parameters**
- \texttt{x} : \{\texttt{Quantity}, array-like, number\}
  
  The value to display

- \texttt{precision} : int, optional
  
  The number of digits of precision (in the best unit, see Examples). If no value is given, numpy’s \texttt{get\_printoptions()} value is used.

**Returns**
- \text{representation} : str
  
  A string representation of this \texttt{Quantity}.

**See also:**

\texttt{Quantity.in\_best\_unit}

**Examples**

```python
>>> from brian2\_units import *
>>> in\_best\_unit(0.00123456 * volt)
'1.23456 mV'
>>> in\_best\_unit(0.00123456 * volt, 2)
'1.23 mV'
>>> in\_best\_unit(0.123456)
'0.123456'
>>> in\_best\_unit(0.123456, 2)
'0.12'
```

\texttt{in\_unit(x, u[, \text{precision}])} Display a value in a certain unit with a given precision.

**in\_unit function**

(*Shortest import: from brian2 import in\_unit*)

\texttt{brian2\_units\_fundamentalunits.\texttt{in\_unit}(x, u, \text{precision=None})}

Display a value in a certain unit with a given precision.

**Parameters**
- \texttt{x} : \{\texttt{Quantity}, array-like, number\}
  
  The value to display

- \texttt{u} : \{\texttt{Quantity.Unit}\}
The unit to display the value \( x \) in.

**precision : int, optional**

The number of digits of precision (in the given unit, see Examples). If no value is given, numpy’s `get_printoptions()` value is used.

**Returns s : str**

A string representation of \( x \) in units of \( u \).

See also:

`Quantity.in_unit`

**Examples**

```python
>>> from brian2 import *
>>> in_unit(3 * volt, mvolt)
'3000. mV'
>>> in_unit(123123 * msecond, second, 2)
'123.12 s'
>>> in_unit(10 * uA/cm**2, nA/um**2)
'1.00000000e-04 nA/um^2'
>>> in_unit(10 * mV, ohm * amp)
'0.01 ohm A'
>>> in_unit(10 * nS, ohm) # doctest: +NORMALIZE_WHITESPACE
... # doctest: +IGNORE_EXCEPTION_DETAIL
Traceback (most recent call last):
... DimensionMismatchError: Non-matching unit for method "in_unit",
dimensions were (m^-2 kg^-1 s^3 A^2) (m^2 kg s^-3 A^-2)
```

**is_dimensionless**

Test if a value is dimensionless or not.

**is_dimensionless function**

(Shortest import: from brian2 import is_dimensionless)

`brian2.units.fundamentalunits.is_dimensionless(obj)`

Test if a value is dimensionless or not.

**Parameters obj : object**

The object to check.

**Returns dimensionless : bool**

True if `obj` is dimensionless.

**is_scalar_type**

Tells you if the object is a 1d number type.

**is_scalar_type function**

(Shortest import: from brian2 import is_scalar_type)
is_scalar_type(obj)
Tells you if the object is a 1d number type.

Parameters  
obj: object
The object to check.

Returns  
scalar: bool
True if obj is a scalar that can be interpreted as a dimensionless Quantity.

quantity_with_dimensions(floatval, dims)
Create a new Quantity with the given dimensions.

quantity_with_dimensions function

(Shortest import: from brian2.units.fundamentalunits import quantity_with_dimensions)

register_new_unit(u)
Register a new unit for automatic displaying of quantities

register_new_unit function

(Shortest import: from brian2 import register_new_unit)
The unit that should be registered.

Examples

```python
>>> from brian2 import *

2.0 farad/metre**2
2.0 * metre**-4 * kilogram**-1 * second**4 * amp**2
>>> register_new_unit(pfarad / mmetre**2)
2.0 farad/metre**2
2000000.0 pfarad/mmetre**2
```

```
wrap_function_change_dimensions(func, ...)  Returns a new function that wraps the given function func so that it changes the dimensions of its input.
```

**wrap_function_change_dimensions function**

*(Shortest import: from brian2.units.fundamentalunits import wrap_function_change_dimensions)*

```
brian2.units.fundamentalunits.wrap_function_change_dimensions(func, change_dim_func)

Returns a new function that wraps the given function func so that it changes the dimensions of its input. Quantities are transformed to unitless numpy arrays before calling func, the output is a quantity with the original dimensions passed through the function change_dim_func. A typical use would be a sqrt function that uses lambda d: d ** 0.5 as change_dim_func.

These transformations apply only to the very first argument, all other arguments are ignored/untouched.
```

```
wrap_function_dimensionless(func)  Returns a new function that wraps the given function func so that it raises a DimensionMismatchError if the function is called on a quantity with dimensions (excluding dimensionless quantities).
```

**wrap_function_dimensionless function**

*(Shortest import: from brian2.units.fundamentalunits import wrap_function_dimensionless)*

```
brian2.units.fundamentalunits.wrap_function_dimensionless(func)

Returns a new function that wraps the given function func so that it raises a DimensionMismatchError if the function is called on a quantity with dimensions (excluding dimensionless quantities). Quantities are transformed to unitless numpy arrays before calling func.

These checks/transformations apply only to the very first argument, all other arguments are ignored/untouched.
```

```
wrap_function_keep_dimensions(func)  Returns a new function that wraps the given function func so that it keeps the dimensions of its input.
```

**wrap_function_keep_dimensions function**
wrap_function_keep_dimensions function

(Shortest import: from brian2.units.fundamentalunits import wrap_function_keep_dimensions)

brian2.units.fundamentalunits.wrap_function_keep_dimensions(func)

Returns a new function that wraps the given function \texttt{func} so that it keeps the dimensions of its input. Quantities are transformed to unitless numpy arrays before calling \texttt{func}, the output is a quantity with the original dimensions re-attached.

These transformations apply only to the very first argument, all other arguments are ignored/untouched, allowing to work functions like \texttt{sum} to work as expected with additional \texttt{axis} etc. arguments.

\begin{verbatim}
wrap_function_remove_dimensions(func) Returns a new function that wraps the given function func so that it removes any dimensions from its input.
\end{verbatim}

wrap_function_remove_dimensions function

(Shortest import: from brian2.units.fundamentalunits import wrap_function_remove_dimensions)

brian2.units.fundamentalunits.wrap_function_remove_dimensions(func)

Returns a new function that wraps the given function \texttt{func} so that it removes any dimensions from its input. Useful for functions that are returning integers (indices) or booleans, irrespective of the datatype contained in the array.

These transformations apply only to the very first argument, all other arguments are ignored/untouched.

\textbf{Objects}

\begin{verbatim}
DIMENSIONLESS The singleton object for dimensionless Dimensions.
\end{verbatim}

DIMENSIONLESS object

(Shortest import: from brian2.units.fundamentalunits import DIMENSIONLESS)

brian2.units.fundamentalunits.DIMENSIONLESS = Dimension()

The singleton object for dimensionless Dimensions.

\begin{verbatim}
additional_unit_register UnitRegistry containing additional units (newton*metre, farad / metre, ...)
\end{verbatim}

additional_unit_register object

(Shortest import: from brian2.units.fundamentalunits import additional_unit_register)

brian2.units.fundamentalunits.additional_unit_register = <brian2.units.fundamentalunits.UnitRegistry object>

UnitRegistry containing additional units (newton*metre, farad / metre, ...)
**standard_unit_register object**

(Shortest import: from brian2.units.fundamentalunits import standard_unit_register)

brian2.units.fundamentalunits.standard_unit_register = <brian2.units.fundamentalunits.UnitRegistry object>

*UnitRegistry* containing all the standard units (metre, kilogram, um...)

**user_unit_register object**

(Shortest import: from brian2.units.fundamentalunits import user_unit_register)

brian2.units.fundamentalunits.user_unit_register = <brian2.units.fundamentalunits.UnitRegistry object>

*UnitRegistry* containing all units defined by the user

**stdunits module**

Optional short unit names

This module defines the following short unit names:

mV, mA, uA (micro_amp), nA, pA, mF, nF, pF, uS, ms, Hz, kHz, MHz, cm, cm^2, cm^3, mm, mm^2, mm^3, um, um^2, um^3

*Exported members:* mV, mA, uA, nA, pA, mF, nF, pF, uS, ms, Hz, kHz, MHz, cm, cm^2, cm^3, mm, mm^2, mm^3, um, um^2, um^3... (3 more members)

**unitsafefunctions module**

Unit-aware replacements for numpy functions.

*Exported members:* log(), log10(), exp(), expm1(), log1p(), exprel(), sin(), cos(), tan(), arcsin(), arccos(), arctan(), sinh(), cosh(), tanh(), arcsinh(), arccosh(), arctanh(), diagonal(), ravel(), trace(), dot(), where(), ones_like(), zeros_like()... (2 more members)

**Functions**

**arange([start[, stop[, step[, dtype]]]])**

Return evenly spaced values within a given interval.

**arange function**

(Shortest import: from brian2 import arange)

brian2.units.unitsafefunctions.arange([start[, stop[, step[, dtype=]]])

Return evenly spaced values within a given interval.
Values are generated within the half-open interval \([\text{start}, \text{stop})\) (in other words, the interval including \(\text{start}\) but excluding \(\text{stop}\)). For integer arguments the function is equivalent to the Python built-in \texttt{range} function, but returns an ndarrray rather than a list.

When using a non-integer step, such as 0.1, the results will often not be consistent. It is better to use \texttt{numpy.linspace} for these cases.

**Parameters**

- \texttt{start} : number, optional
  
  Start of interval. The interval includes this value. The default start value is 0.

- \texttt{stop} : number
  
  End of interval. The interval does not include this value, except in some cases where \texttt{step} is not an integer and floating point round-off affects the length of \texttt{out}.

- \texttt{step} : number, optional
  
  Spacing between values. For any output \texttt{out}, this is the distance between two adjacent values, \(\text{out}[i+1] - \text{out}[i]\). The default step size is 1. If \texttt{step} is specified as a position argument, \texttt{start} must also be given.

- \texttt{dtype} : dtype
  
  The type of the output array. If \texttt{dtype} is not given, infer the data type from the other input arguments.

**Returns**

- \texttt{arange} : ndarray
  
  Array of evenly spaced values.

  For floating point arguments, the length of the result is \(\text{ceil}((\text{stop} - \text{start})/\text{step})\). Because of floating point overflow, this rule may result in the last element of \texttt{out} being greater than \texttt{stop}.

**See also:**

- \texttt{linspace()} Evenly spaced numbers with careful handling of endpoints.
- \texttt{ogrid} Arrays of evenly spaced numbers in N-dimensions.
- \texttt{mgrid} Grid-shaped arrays of evenly spaced numbers in N-dimensions.

**Examples**

```python
>>> np.arange(3)
anarray([0, 1, 2])

>>> np.arange(3.0)
anarray([ 0., 1., 2.])

>>> np.arange(3,7)
anarray([3, 4, 5, 6])

>>> np.arange(3,7,2)
anarray([3, 5])
```

\texttt{arccos(x, /[, out, where, casting, order, ...])} \hspace{1cm} \text{Trigonometric inverse cosine, element-wise.}

**arccos function**

(\textit{Shortest import: from brian2 import arccos})
brian2.units.unitsafefunctions.arccos(x, /, out=None, *, where=True, casting='same_kind', order='K', dtype=None, subok=True, outobj=None)

Trigonometric inverse cosine, element-wise.

The inverse of \( \cos() \) so that, if \( y = \cos(x) \), then \( x = \arccos(y) \).

Parameters

- **x**: array_like
  \( x \)-coordinate on the unit circle. For real arguments, the domain is \([-1, 1]\).

- **out**: ndarray, None, or tuple of ndarray and None, optional
  A location into which the result is stored. If provided, it must have a shape that the inputs broadcast to. If not provided or None, a freshly-allocated array is returned. A tuple (possible only as a keyword argument) must have length equal to the number of outputs.

- **where**: array_like, optional
  Values of True indicate to calculate the ufunc at that position, values of False indicate to leave the value in the output alone.

- ****kwargs:
  For other keyword-only arguments, see the ufunc docs.

Returns

- **angle**: ndarray
  The angle of the ray intersecting the unit circle at the given \( x \)-coordinate in radians \([0, \pi]\). This is a scalar if \( x \) is a scalar.

See also:

- \( \cos() \), \( \arctan() \), \( \arcsin() \), emath.arccos

Notes

- \( \arccos() \) is a multivalued function: for each \( x \) there are infinitely many numbers \( z \) such that \( \cos(z) = x \). The convention is to return the angle \( z \) whose real part lies in \([0, \pi]\).

- For real-valued input data types, \( \arccos() \) always returns real output. For each value that cannot be expressed as a real number or infinity, it yields nan and sets the invalid floating point error flag.

- For complex-valued input, \( \arccos() \) is a complex analytic function that has branch cuts \([-\infty, -1]\) and \([1, \infty]\) and is continuous from above on the former and from below on the latter.

- The inverse \( \cos() \) is also known as \( \acos \) or \( \cos^{-1} \).

References


Examples

We expect the arccos of 1 to be 0, and of -1 to be \( \pi \):

```python
>>> np.arccos([1, -1])
array([ 0.        ,  3.14159265])
```
## arccosh

Inverse hyperbolic cosine, element-wise.

**arccosh**<br>

*arccosh(x, out=None, *, where=True, casting='same_kind', order='K', dtype=None, subok=True, signature=None, extobj=None)\n*

Inverse hyperbolic cosine, element-wise.

**Parameters**

- **x**: array_like
  
  Input array.

- **out**: ndarray, None, or tuple of ndarray and None, optional
  
  A location into which the result is stored. If provided, it must have a shape that the inputs broadcast to. If not provided or None, a freshly-allocated array is returned. A tuple (possible only as a keyword argument) must have length equal to the number of outputs.

- **where**: array_like, optional
  
  Values of True indicate to calculate the ufunc at that position, values of False indicate to leave the value in the output alone.

- ****kwarg**: 
  
  For other keyword-only arguments, see the ufunc docs.

**Returns**

- **arccosh**: ndarray
  
  Array of the same shape as `x`. This is a scalar if `x` is a scalar.

See also:

- `cosh()`, `arcsinh()`, `sinh()`, `arctanh()`, `tanh()`

**Notes**

`arccosh()` is a multivalued function: for each `x` there are infinitely many numbers `z` such that \( \cosh(z) = x \). The convention is to return the `z` whose imaginary part lies in \([-\pi, \pi]\) and the real part in \([0, \infty]\).

For real-valued input data types, `arccosh()` always returns real output. For each value that cannot be expressed as a real number or infinity, it yields nan and sets the invalid floating point error flag.

For complex-valued input, `arccosh()` is a complex analytical function that has a branch cut \([-\infty, 1]\) and is continuous from above on it.
References

[R13], [R14]

Examples

```python
g>>> np.arccosh([np.e, 10.0])
aarray([ 1.65745445, 2.99322285])
g>>> np.arccosh(1)
0.0
```

`arcsin(x, [out, where, casting, order, ...])`  
Inverse sine, element-wise.

**arcsin function**

(Shortest import: from brian2 import arcsin)

```
brian2.units.unitsafefunctions.arcsin(x, /, out=None, *, where=True, casting='same_kind', order='K', dtype=None, subok=True, signature=None, extobj=None)
```

Inverse sine, element-wise.

**Parameters**  
`x` : array_like  
y-coordinate on the unit circle.

`out` : ndarray, None, or tuple of ndarray and None, optional  
A location into which the result is stored. If provided, it must have a shape that the inputs broadcast to. If not provided or None, a freshly-allocated array is returned. A tuple (possible only as a keyword argument) must have length equal to the number of outputs.

`where` : array_like, optional  
Values of True indicate to calculate the ufunc at that position, values of False indicate to leave the value in the output alone.

**kwargs** :
For other keyword-only arguments, see the ufunc docs.

**Returns**  
`angle` : ndarray  
The inverse sine of each element in x, in radians and in the closed interval [-pi/2, pi/2]. This is a scalar if x is a scalar.

**See also:**

`sin()`, `cos()`, `arccos()`, `tan()`, `arctan()`, `arctan2`, `emath.arcsin`

**Notes**

`arcsin()` is a multivalued function: for each x there are infinitely many numbers z such that $\sin(z) = x$. The convention is to return the angle z whose real part lies in [-pi/2, pi/2].
For real-valued input data types, \textit{arcsin} always returns real output. For each value that cannot be expressed as a real number or infinity, it yields \texttt{nan} and sets the \texttt{invalid} floating point error flag.

For complex-valued input, \textit{arcsin()} is a complex analytic function that has, by convention, the branch cuts [-inf, -1] and [1, inf] and is continuous from above on the former and from below on the latter.

The inverse sine is also known as \texttt{asin} or \texttt{sin^{-1}}.

\textbf{References}


\textbf{Examples}

\begin{verbatim}
>>> np.arcsin(1) # pi/2 1.5707963267948966
>>> np.arcsin(-1) # -pi/2 -1.5707963267948966
>>> np.arcsin(0) 0.0
\end{verbatim}

\texttt{arcsinh(x, /[, out, where, casting, order, ...])} Inverse hyperbolic sine element-wise.

\textbf{arcsinh function}

(\textit{Shortest import: from brian2 import arcsinh})

\begin{verbatim}
brian2.units.unitsafefunctions.arcsinh(x, /, out=None, *, where=True, casting='same_kind', order='K', dtype=None, subok=True[, signature, extobj])
\end{verbatim}

Inverse hyperbolic sine element-wise.

\textbf{Parameters} \texttt{x} : array_like

Input array.

\texttt{out} : ndarray, None, or tuple of ndarray and None, optional

A location into which the result is stored. If provided, it must have a shape that the inputs broadcast to. If not provided or \texttt{None}, a freshly-allocated array is returned. A tuple (possible only as a keyword argument) must have length equal to the number of outputs.

\texttt{where} : array_like, optional

Values of True indicate to calculate the ufunc at that position, values of False indicate to leave the value in the output alone.

**\texttt{kwargs}** :

For other keyword-only arguments, see the ufunc docs.

\textbf{Returns} \texttt{out} : ndarray or scalar

Array of the same shape as \texttt{x}. This is a scalar if \texttt{x} is a scalar.
Notes

arcsinh() is a multivalued function: for each \( x \) there are infinitely many numbers \( z \) such that \( \sinh(z) = x \). The convention is to return the \( z \) whose imaginary part lies in \([-\pi/2, \pi/2]\).

For real-valued input data types, arcsinh() always returns real output. For each value that cannot be expressed as a real number or infinity, it returns nan and sets the invalid floating point error flag.

For complex-valued input, arccos() is a complex analytical function that has branch cuts \([1j, \text{inf}j]\) and \([-1j, -\text{inf}j]\) and is continuous from the right on the former and from the left on the latter.

The inverse hyperbolic sine is also known as asinh or sinh^-1.

References

[R15], [R16]

Examples

```python
>>> np.arcsinh(np.array([np.e, 10.0]))
array([ 1.72538256, 2.99822295])
```

arctan function

(Shortest import: from brian2 import arctan)

brian2.units.unitsafefunctions.arctan(x, /[, out, where, casting, order, ...])  
Trigonometric inverse tangent, element-wise.

The inverse of tan, so that if \( y = \tan(x) \) then \( x = \arctan(y) \).

Parameters

- **x** : array_like
  - out : ndarray, None, or tuple of ndarray and None, optional
    - A location into which the result is stored. If provided, it must have a shape that the inputs broadcast to. If not provided or None, a freshly-allocated array is returned. A tuple (possible only as a keyword argument) must have length equal to the number of outputs.
  - where : array_like, optional
    - Values of True indicate to calculate the ufunc at that position, values of False indicate to leave the value in the output alone.
  - **kwargs** :
    - For other keyword-only arguments, see the ufunc docs.

Returns

- **out** : ndarray or scalar
  - Out has the same shape as \( x \). Its real part is in \([-\pi/2, \pi/2]\) (arctan(+/-inf) returns +/-\pi/2). This is a scalar if \( x \) is a scalar.
See also:

arctan2 The “four quadrant” arctan of the angle formed by \((x, y)\) and the positive \(x\)-axis.

angle() Argument of complex values.

Notes

\(\text{arctan}()\) is a multi-valued function: for each \(x\) there are infinitely many numbers \(z\) such that \(\tan(z) = x\). The convention is to return the angle \(z\) whose real part lies in \([-\pi/2, \pi/2]\).

For real-valued input data types, \(\text{arctan}()\) always returns real output. For each value that cannot be expressed as a real number or infinity, it yields \(\text{nan}\) and sets the \text{invalid} floating point error flag.

For complex-valued input, \(\text{arctan}()\) is a complex analytic function that has \([1j, \inf j]\) and \([-1j, -\inf j]\) as branch cuts, and is continuous from the left on the former and from the right on the latter.

The inverse tangent is also known as \(\text{atan}\) or \(\tan^{-1}\).

References


Examples

We expect the arctan of 0 to be 0, and of 1 to be \(\pi/4\):

```python
>>> np.arctan([0, 1])
array([ 0. , 0.78539816])
```

```python
>>> np.pi/4
0.78539816339744828
```

Plot arctan:

```python
>>> import matplotlib.pyplot as plt
>>> x = np.linspace(-10, 10)
>>> plt.plot(x, np.arctan(x))
>>> plt.axis('tight')
>>> plt.show()
```

\(\text{arctan}(x, /[, out, where, casting, order, ...])\) Inverse hyperbolic tangent element-wise.

\(\text{arctanh}\) function

(Shortest import: from brian2 import arctanh)

```python
brian2.units.unitsafefunctions.arctanh(x, /, out=None, *, where=True, casting='same_kind', order='K', dtype=None, subok=True[, signature, extobj])
```

Inverse hyperbolic tangent element-wise.

**Parameters**

\(x\) : array_like
Input array.

**out** : ndarray, None, or tuple of ndarray and None, optional

A location into which the result is stored. If provided, it must have a shape that the inputs broadcast to. If not provided or `None`, a freshly-allocated array is returned. A tuple (possible only as a keyword argument) must have length equal to the number of outputs.

**where** : array_like, optional

Values of True indicate to calculate the ufunc at that position, values of False indicate to leave the value in the output alone.

**kwargs** :

For other keyword-only arguments, see the ufunc docs.

**Returns**

**out** : ndarray or scalar

Array of the same shape as `x`. This is a scalar if `x` is a scalar.

See also:

emath.arctanh

Notes

`arctanh()` is a multivalued function: for each `x` there are infinitely many numbers `z` such that `tanh(z) = x`. The convention is to return the `z` whose imaginary part lies in `[-pi/2, pi/2]`.

For real-valued input data types, `arctanh()` always returns real output. For each value that cannot be expressed as a real number or infinity, it yields `nan` and sets the `invalid` floating point error flag.

For complex-valued input, `arctanh()` is a complex analytical function that has branch cuts `[-1, -inf]` and `[1, inf]` and is continuous from above on the former and from below on the latter.

The inverse hyperbolic tangent is also known as `atanh` or `tanh^-1`.

References

[R17], [R18]

Examples

```python
>>> np.arctanh([0, -0.5])
array([ 0. , -0.54930614])
```

cos(x, /[, out, where, casting, order, ...]) Cosine element-wise.

**cos function**

(Shortest import: from brian2 import cos)

brian2.units.unitsafefunctions.cos(x, /, out=None, *, where=True, casting='same_kind', order='K', dtype=None, subok=True[, signature, extobj])

Cosine element-wise.
**Parameters**

**x** : array_like

Input array in radians.

**out** : ndarray, None, or tuple of ndarray and None, optional

A location into which the result is stored. If provided, it must have a shape that the inputs broadcast to. If not provided or None, a freshly-allocated array is returned. A tuple (possible only as a keyword argument) must have length equal to the number of outputs.

**where** : array_like, optional

Values of True indicate to calculate the ufunc at that position, values of False indicate to leave the value in the output alone.

**kwargs** :

For other keyword-only arguments, see the ufunc docs.

**Returns**

**y** : ndarray

The corresponding cosine values. This is a scalar if x is a scalar.

**Notes**

If out is provided, the function writes the result into it, and returns a reference to out. (See Examples)

**References**


**Examples**

```python
>>> np.cos(np.array([0, np.pi/2, np.pi]))
array([ 1.00000000e+00, 6.12303177e-17, -1.00000000e+00])
```

# Example of providing the optional output parameter
>>> out2 = np.cos([0.1], out1)

# Example of ValueError due to provision of shape mis-matched 'out'
>>> np.cos(np.zeros((3,3)),np.zeros((2,2)))
Traceback (most recent call last):
  File "<stdin>", line 1, in <module>
ValueError: operands could not be broadcast together with shapes (3,3) (2,2)
```

$\cosh(x, /[, out, where, casting, order, ...])$ Hyperbolic cosine, element-wise.

**cosh function**

*(Shortest import: from brian2 import cosh)*

6.6. Subpackages
brian2.units.unitsafefunctions.cosh(x, out=None, *, where=True, casting='same_kind', order='K', dtype=None, subok=True, signature=None, extobj=None)

Hyperbolic cosine, element-wise.
Equivalent to 1/2 * (np.exp(x) + np.exp(-x)) and np.cos(1j*x).

Parameters

- **x**: array_like
  Input array.

- **out**: ndarray, None, or tuple of ndarray and None, optional
  A location into which the result is stored. If provided, it must have a shape that the inputs broadcast to. If not provided or None, a freshly-allocated array is returned. A tuple (possible only as a keyword argument) must have length equal to the number of outputs.

- **where**: array_like, optional
  Values of True indicate to calculate the ufunc at that position, values of False indicate to leave the value in the output alone.

- ****kwargs:*
  For other keyword-only arguments, see the ufunc docs.

Returns

- **out**: ndarray or scalar
  Output array of same shape as x. This is a scalar if x is a scalar.

Examples

```python
>>> np.cosh(0)
1.0
```

The hyperbolic cosine describes the shape of a hanging cable:

```python
>>> import matplotlib.pyplot as plt
>>> x = np.linspace(-4, 4, 1000)
>>> plt.plot(x, np.cosh(x))
>>> plt.show()
```

**diagonal**

(Shortest import: from brian2 import diagonal)

brian2.units.unitsafefunctions.diagonal(a, offset=0, axis1=0, axis2=1)

Return specified diagonals.

If a is 2-D, returns the diagonal of a with the given offset, i.e., the collection of elements of the form a[i, i+offset]. If a has more than two dimensions, then the axes specified by axis1 and axis2 are used to determine the 2-D sub-array whose diagonal is returned. The shape of the resulting array can be determined by removing axis1 and axis2 and appending an index to the right equal to the size of the resulting diagonals.

In versions of NumPy prior to 1.7, this function always returned a new, independent array containing a copy of the values in the diagonal.
In NumPy 1.7 and 1.8, it continues to return a copy of the diagonal, but depending on this fact is deprecated. Writing to the resulting array continues to work as it used to, but a FutureWarning is issued.

Starting in NumPy 1.9 it returns a read-only view on the original array. Attempting to write to the resulting array will produce an error.

In some future release, it will return a read/write view and writing to the returned array will alter your original array. The returned array will have the same type as the input array.

If you don’t write to the array returned by this function, then you can just ignore all of the above.

If you depend on the current behavior, then we suggest copying the returned array explicitly, i.e., use `np.diagonal(a).copy()` instead of just `np.diagonal(a)`. This will work with both past and future versions of NumPy.

**Parameters**

- `a` : array_like
  - Array from which the diagonals are taken.
- `offset` : int, optional
  - Offset of the diagonal from the main diagonal. Can be positive or negative. Defaults to main diagonal (0).
- `axis1` : int, optional
  - Axis to be used as the first axis of the 2-D sub-arrays from which the diagonals should be taken. Defaults to first axis (0).
- `axis2` : int, optional
  - Axis to be used as the second axis of the 2-D sub-arrays from which the diagonals should be taken. Defaults to second axis (1).

**Returns**

- `array_of_diagonals` : ndarray
  - If `a` is 2-D, then a 1-D array containing the diagonal and of the same type as `a` is returned unless `a` is a `matrix`, in which case a 1-D array rather than a (2-D) `matrix` is returned in order to maintain backward compatibility.
  - If `a.ndim > 2`, then the dimensions specified by `axis1` and `axis2` are removed, and a new axis inserted at the end corresponding to the diagonal.

**Raises**

- `ValueError` If the dimension of `a` is less than 2.

**See also:**

- `diag()` MATLAB work-a-like for 1-D and 2-D arrays.
- `diagflat()` Create diagonal arrays.
- `trace()` Sum along diagonals.

**Examples**
>>> a = np.arange(4).reshape(2,2)

```python
>>> a
array([[0, 1],
       [2, 3]])
```

```python
>>> a.diagonal()
array([0, 3])
```

```python
>>> a.diagonal(1)
array([1])
```

A 3-D example:

```python
>>> a = np.arange(8).reshape(2,2,2); a
array([[[0, 1],
        [2, 3]],
       [[4, 5],
        [6, 7]]])
```

```python
>>> a.diagonal(0, 0, 0) # Main diagonals of two arrays created by skipping ...
array([[0, 6],
       [1, 7]])
```

The sub-arrays whose main diagonals we just obtained; note that each corresponds to fixing the right-most (column) axis, and that the diagonals are “packed” in rows.

```python
>>> a[:,:,0] # main diagonal is [0 6]
array([[0, 2],
       [4, 6]])
```

```python
>>> a[:,:,1] # main diagonal is [1 7]
array([[1, 3],
       [5, 7]])
```

**dot**

**dot function**

((Shortest import: from brian2 import dot)

brian2.units.unitsafefunctions.dot (a, b, out=None)

Dot product of two arrays. Specifically,

- If both a and b are 1-D arrays, it is inner product of vectors (without complex conjugation).
- If both a and b are 2-D arrays, it is matrix multiplication, but using matmul() or a @ b is preferred.
- If either a or b is 0-D (scalar), it is equivalent to multiply() and using numpy.multiply(a, b) or a * b is preferred.
- If a is an N-D array and b is a 1-D array, it is a sum product over the last axis of a and b.
- If a is an N-D array and b is an M-D array (where M>=2), it is a sum product over the last axis of a and the second-to-last axis of b:

```python
    dot(a, b)[i,j,k,m] = sum(a[i,j,:]*b[k,:,m])
```

**Parameters**

- **a**: array_like
First argument.

\[ b : \text{array_like} \]

Second argument.

\[ \text{out} : \text{ndarray, optional} \]

Output argument. This must have the exact kind that would be returned if it was not used. In particular, it must have the right type, must be C-contiguous, and its dtype must be the dtype that would be returned for \( \text{dot}(a, b) \). This is a performance feature. Therefore, if these conditions are not met, an exception is raised, instead of attempting to be flexible.

**Returns** \( \text{output} : \text{ndarray} \)

Returns the dot product of \( a \) and \( b \). If \( a \) and \( b \) are both scalars or both 1-D arrays then a scalar is returned; otherwise an array is returned. If \( \text{out} \) is given, then it is returned.

**Raises**

**ValueError** If the last dimension of \( a \) is not the same size as the second-to-last dimension of \( b \).

**See also:**

- \text{vdot} Complex-conjugating dot product.
- \text{tensordot} Sum products over arbitrary axes.
- \text{einsum} Einstein summation convention.
- \text{matmul} ‘@’ operator as method with out parameter.

**Examples**

```python
>>> np.dot(3, 4)
12
```

Neither argument is complex-conjugated:

```python
>>> np.dot([2j, 3j], [2j, 3j])
(-13+0j)
```

For 2-D arrays it is the matrix product:

```python
>>> a = [[1, 0], [0, 1]]
>>> b = [[4, 1], [2, 2]]
>>> np.dot(a, b)
array([[4, 1],
       [2, 2]])
```

```python
>>> a = np.arange(3*4*5*6).reshape((3,4,5,6))
>>> b = np.arange(3*4*5*6)[::-1].reshape((5,4,6,3))
>>> np.dot(a, b)[2,3,2,1,2,2]
499128
>>> sum(a[2,3,2,:] * b[1,2,:,2])
499128
```
\[ \exp(x, \text{out}, \text{where}, \text{casting}, \text{order}, \ldots) \] Calculate the exponential of all elements in the input array.

\textbf{exp function}

(\textit{Shortest import: from brian2 import exp})

\begin{verbatim}
brian2.units.unitsafefunctions.exp(x, out=None, *, where=True, casting='same_kind', order='K', dtype=None, subok=True, signature, extobj)
\
Calculate the exponential of all elements in the input array.

\textbf{Parameters} x : array_like
\hspace{1cm} Input values.

out : ndarray, None, or tuple of ndarray and None, optional
\hspace{1cm} A location into which the result is stored. If provided, it must have a shape that the inputs broadcast to. If not provided or None, a freshly-allocated array is returned. A tuple (possible only as a keyword argument) must have length equal to the number of outputs.

where : array_like, optional
\hspace{1cm} Values of True indicate to calculate the ufunc at that position, values of False indicate to leave the value in the output alone.

**kwargs :
\hspace{1cm} For other keyword-only arguments, see the ufunc docs.

\textbf{Returns} out : ndarray or scalar
\hspace{1cm} Output array, element-wise exponential of x. This is a scalar if x is a scalar.

See also:

expm1 () Calculate \( \exp(x) - 1 \) for all elements in the array.

exp2 Calculate \( 2^x \) for all elements in the array.

\textbf{Notes}

The irrational number \( e \) is also known as Euler's number. It is approximately 2.718281, and is the base of the natural logarithm, \( \ln \) (this means that, if \( x = \ln y = \log_e y \), then \( e^x = y \). For real input, \( \exp(x) \) is always positive.

For complex arguments, \( x = a + ib \), we can write \( e^{x} = e^{a}e^{ib} \). The first term, \( e^{a} \), is already known (it is the real argument, described above). The second term, \( e^{ib} \), is \( \cos b + i \sin b \), a function with magnitude 1 and a periodic phase.

\textbf{References}

[R19], [R20]
Examples

Plot the magnitude and phase of $\exp(x)$ in the complex plane:

```python
>>> import matplotlib.pyplot as plt

>>> x = np.linspace(-2*np.pi, 2*np.pi, 100)
>>> xx = x + 1j * x[:, np.newaxis] # a + ib over complex plane
>>> out = np.exp(xx)

>>> plt.subplot(121)
>>> plt.imshow(np.abs(out),
... extent=[-2*np.pi, 2*np.pi, -2*np.pi, 2*np.pi], cmap='gray')
>>> plt.title('Magnitude of exp(x)')

>>> plt.subplot(122)
>>> plt.imshow(np.angle(out),
... extent=[-2*np.pi, 2*np.pi, -2*np.pi, 2*np.pi], cmap='hsv')
>>> plt.title('Phase (angle) of exp(x)')
>>> plt.show()
```

`linspace`(start, stop[, num, endpoint, ...]) Return evenly spaced numbers over a specified interval.

### linspace function

*(Shortest import: from brian2 import linspace)*

brian2.units.unit_safe_functions` . linspace(start, stop=50, endpoint=True, retstep=False, dtype=None, axis=0)

Return evenly spaced numbers over a specified interval.

Returns `num` evenly spaced samples, calculated over the interval `[start, stop()]`.

The endpoint of the interval can optionally be excluded.

Changed in version 1.16.0: Non-scalar `start` and `stop()` are now supported.

**Parameters**

- `start`: array_like
  The starting value of the sequence.

- `stop`: array_like
  The end value of the sequence, unless `endpoint` is set to False. In that case, the sequence consists of all but the last of `num + 1` evenly spaced samples, so that `stop()` is excluded. Note that the step size changes when `endpoint` is False.

- `num`: int, optional
  Number of samples to generate. Default is 50. Must be non-negative.

- `endpoint`: bool, optional
  If True, `stop()` is the last sample. Otherwise, it is not included. Default is True.

- `retstep`: bool, optional
  If True, return `(samples, step())`, where `step()` is the spacing between samples.

- `dtype`: dtype, optional
The type of the output array. If `dtype` is not given, infer the data type from the other input arguments.

New in version 1.9.0.

`axis` : int, optional

The axis in the result to store the samples. Relevant only if start or stop are array-like. By default (0), the samples will be along a new axis inserted at the beginning. Use -1 to get an axis at the end.

New in version 1.16.0.

Returns `samples` : ndarray

There are `num` equally spaced samples in the closed interval `[start, stop]` or the half-open interval `[start, stop)` (depending on whether `endpoint` is True or False).

`step` : float, optional

Only returned if `retstep` is True

Size of spacing between samples.

See also:

`arange()` Similar to `linspace()`, but uses a step size (instead of the number of samples).

`geomspace()` Similar to `linspace()`, but with numbers spaced evenly on a log scale (a geometric progression).

`logspace()` Similar to `geomspace()`, but with the end points specified as logarithms.

Examples

```python
>>> np.linspace(2.0, 3.0, num=5)
array([ 2. , 2.25, 2.5 , 2.75, 3. ])
>>> np.linspace(2.0, 3.0, num=5, endpoint=False)
array([ 2. , 2.2, 2.4, 2.6, 2.8])
>>> np.linspace(2.0, 3.0, num=5, retstep=True)
(array([ 2. , 2.25, 2.5 , 2.75, 3. ]), 0.25)
```

Graphical illustration:

```python
>>> import matplotlib.pyplot as plt
...
>>> N = 8
>>> y = np.zeros(N)
>>> x1 = np.linspace(0, 10, N, endpoint=True)
>>> x2 = np.linspace(0, 10, N, endpoint=False)
>>> plt.plot(x1, y, 'o')
...[<matplotlib.lines.Line2D object at 0x...>]
>>> plt.plot(x2, y + 0.5, 'o')
...[<matplotlib.lines.Line2D object at 0x...>]
>>> plt.ylim([-0.5, 1])
...(-0.5, 1)
>>> plt.show()
```
log function

(Shortest import: from brian2 import log)

brian2.units.unitsafefunctions.log(x, out=None, *, where=True, casting='same_kind', order='K', dtype=None, subok=True[, signature, extobj])

Natural logarithm, element-wise.

The natural logarithm \( \log() \) is the inverse of the exponential function, so that \( \log(\exp(x)) = x \). The natural logarithm is logarithm in base \( e \).

Parameters x : array_like
    Input value.

out : ndarray, None, or tuple of ndarray and None, optional
    A location into which the result is stored. If provided, it must have a shape that the inputs broadcast to. If not provided or None, a freshly-allocated array is returned. A tuple (possible only as a keyword argument) must have length equal to the number of outputs.

where : array_like, optional
    Values of True indicate to calculate the ufunc at that position, values of False indicate to leave the value in the output alone.

**kwargs :
    For other keyword-only arguments, see the ufunc docs.

Returns y : ndarray
    The natural logarithm of x, element-wise. This is a scalar if x is a scalar.

See also:
    log10(), log2, log1p(), emath.log

Notes

Logarithm is a multivalued function: for each \( x \) there is an infinite number of \( z \) such that \( \exp(z) = x \). The convention is to return the \( z \) whose imaginary part lies in \([-\pi, \pi]\).

For real-valued input data types, \( \log() \) always returns real output. For each value that cannot be expressed as a real number or infinity, it yields nan and sets the invalid floating point error flag.

For complex-valued input, \( \log() \) is a complex analytical function that has a branch cut \([-\infty, 0]\) and is continuous from above on it. \( \log() \) handles the floating-point negative zero as an infinitesimal negative number, conforming to the C99 standard.

References

[R21], [R22]
Examples

```python
>>> np.log([1, np.e, np.e**2, 0])
array([ 0., 1., 2., -Inf])
```

```
ravel(a[, order])
```

Return a contiguous flattened array.

ravel function

(Shortest import: from brian2 import ravel)

brian2.units.unitsafefunctions.ravel(a, order='C')

Return a contiguous flattened array.

A 1-D array, containing the elements of the input, is returned. A copy is made only if needed.

As of NumPy 1.10, the returned array will have the same type as the input array. (for example, a masked array will be returned for a masked array input)

**Parameters**

- **a**: array_like
  - Input array. The elements in `a` are read in the order specified by `order`, and packed as a 1-D array.

- **order**: {'C', 'F', 'A', 'K'}, optional
  - The elements of `a` are read using this index order. 'C' means to index the elements in row-major, C-style order, with the last axis index changing fastest, back to the first axis index changing slowest. 'F' means to index the elements in column-major, Fortran-style order, with the first index changing fastest, and the last index changing slowest. Note that the ‘C’ and ‘F’ options take no account of the memory layout of the underlying array, and only refer to the order of axis indexing. ‘A’ means to read the elements in Fortran-like index order if `a` is Fortran contiguous in memory, C-like order otherwise. ‘K’ means to read the elements in the order they occur in memory, except for reversing the data when strides are negative. By default, ‘C’ index order is used.

**Returns**

- **y**: array_like
  - `y` is an array of the same subtype as `a`, with shape `(a.size,)`. Note that matrices are special cased for backward compatibility, if `a` is a matrix, then `y` is a 1-D ndarray.

See also:

- `ndarray.flat` 1-D iterator over an array.
- `ndarray.flatten` 1-D array copy of the elements of an array in row-major order.
- `ndarray.reshape` Change the shape of an array without changing its data.

Notes

In row-major, C-style order, in two dimensions, the row index varies the slowest, and the column index the quickest. This can be generalized to multiple dimensions, where row-major order implies that the index along the first axis varies slowest, and the index along the last quickest. The opposite holds for column-major, Fortran-style index ordering.

When a view is desired in as many cases as possible, `arr.reshape(-1)` may be preferable.
Examples

It is equivalent to `reshape(-1, order=order)`.

```python
>>> x = np.array([[1, 2, 3], [4, 5, 6]])
>>> print(np.ravel(x))
[1 2 3 4 5 6]
```

```python
>>> print(x.reshape(-1))
[1 2 3 4 5 6]
```

```python
>>> print(np.ravel(x, order='F'))
[1 4 2 5 3 6]
```

When `order` is ‘A’, it will preserve the array’s ‘C’ or ‘F’ ordering:

```python
>>> print(np.ravel(x.T))
[1 4 2 5 3 6]
>>> print(np.ravel(x.T, order='A'))
[1 2 3 4 5 6]
```

When `order` is ‘K’, it will preserve orderings that are neither ‘C’ nor ‘F’, but won’t reverse axes:

```python
>>> a = np.arange(3)[::-1]; a
array([2, 1, 0])
>>> a.ravel(order='C')
array([2, 1, 0])
>>> a.ravel(order='K')
array([2, 1, 0])
```

```python
>>> a = np.arange(12).reshape(2,3,2).swapaxes(1,2); a
array([[[ 0, 2, 4],
        [ 1, 3, 5]],
        [[ 6, 8, 10],
        [ 7, 9, 11]]])
>>> a.ravel(order='C')
array([ 0, 2, 4, 1, 3, 5, 6, 8, 10, 7, 9, 11])
>>> a.ravel(order='K')
array([ 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11])
```

```
\[
\sin(x, /[, out, where, casting, order, ...])
\]

Trigonometric sine, element-wise.

**sin function**

(Shortest import: from brian2 import sin)

brian2.units.unitsafefunctions.sin(x, /, out=None, *, where=True, casting='same_kind', order='K', dtype=None, subok=True[, signature, extobj])

Trigonometric sine, element-wise.

Parameters

- **x**: array_like
  Angle, in radians (2\(\pi\) rad equals 360 degrees).
- **out**: ndarray, None, or tuple of ndarray and None, optional
A location into which the result is stored. If provided, it must have a shape that the inputs broadcast to. If not provided or None, a freshly-allocated array is returned. A tuple (possible only as a keyword argument) must have length equal to the number of outputs.

**where**: array_like, optional

Values of True indicate to calculate the ufunc at that position, values of False indicate to leave the value in the output alone.

**kwargs**:

For other keyword-only arguments, see the ufunc docs.

**Returns**

- **y**: array_like

  The sine of each element of x. This is a scalar if x is a scalar.

**See also**:

-arcsin(), sinh(), cos()—

**Notes**

The sine is one of the fundamental functions of trigonometry (the mathematical study of triangles). Consider a circle of radius 1 centered on the origin. A ray comes in from the +x axis, makes an angle at the origin (measured counter-clockwise from that axis), and departs from the origin. The y coordinate of the outgoing ray’s intersection with the unit circle is the sine of that angle. It ranges from -1 for x = 3π/2 to +1 for π/2. The function has zeroes where the angle is a multiple of π. Sines of angles between π and 2π are negative. The numerous properties of the sine and related functions are included in any standard trigonometry text.

**Examples**

Print sine of one angle:

```python
>>> np.sin(np.pi/2.)
1.0
```

Print sines of an array of angles given in degrees:

```python
>>> np.sin(np.array((0., 30., 45., 60., 90.)) * np.pi / 180.)
array([ 0.0, 0.5, 0.70710678, 0.8660254, 1.])
```

Plot the sine function:

```python
>>> import matplotlib.pyplot as plt
>>> x = np.linspace(-np.pi, np.pi, 201)
>>> plt.plot(x, np.sin(x))
>>> plt.xlabel('Angle [rad]')
>>> plt.ylabel('sin(x)')
>>> plt.axis('tight')
>>> plt.show()
```

**sinh(x[, out, where, casting, order, ...])** — Hyperbolic sine, element-wise.
**sinh function**

*(Shortest import: from brian2 import sinh)*

```python
brian2.units.unitsafefunctions.sinh(x, out=None, *, where=True, casting='same_kind', order='K', dtype=None, subok=True, signature=None, extobj=None)
```

Hyperbolic sine, element-wise.

Equivalent to \( \frac{1}{2} \left( \exp(x) - \exp(-x) \right) \) or \(-i \sin(1j x)\).

**Parameters**

- **x**: array_like
  - Input array.
- **out**: ndarray, None, or tuple of ndarray and None, optional
  - A location into which the result is stored. If provided, it must have a shape that the inputs broadcast to. If not provided or None, a freshly-allocated array is returned. A tuple (possible only as a keyword argument) must have length equal to the number of outputs.
- **where**: array_like, optional
  - Values of True indicate to calculate the ufunc at that position, values of False indicate to leave the value in the output alone.
- ****kwargs**:
  - For other keyword-only arguments, see the ufunc docs.

**Returns**

- **y**: ndarray
  - The corresponding hyperbolic sine values. This is a scalar if x is a scalar.

**Notes**

If out is provided, the function writes the result into it, and returns a reference to out. (See Examples)

**References**


**Examples**

```python
>>> np.sinh(0)
0.0
>>> np.sinh(np.pi*1j/2)
1j
>>> np.sinh(np.pi*1j)  # (exact value is 0)
1.2246063538223773e-016j
>>> # Discrepancy due to vagaries of floating point arithmetic.
```

```python
>>> # Example of providing the optional output parameter
>>> out2 = np.sinh([0.1], out1)
>>> out2 is out1
True
```
## Example of ValueError due to provision of shape mis-matched `out`

```python
>>> np.sinh(np.zeros((3,3)),np.zeros((2,2)))
Traceback (most recent call last):
  File "<stdin>", line 1, in <module>
ValueError: operands could not be broadcast together with shapes (3,3) (2,2)
```

---

### tan

#### tan function

*(Shortest import: from brian2 import tan)*

```python
brian2.units.unitsafefunctions.tan(x, /, out=None, *, where=True, casting='same_kind', order='K', dtype=None, subok=True[, signature, extobj])
```

Compute tangent element-wise.

Equivalent to `np.sin(x)/np.cos(x)` element-wise.

**Parameters**

- `x`: array_like
  - Input array.
- `out`: ndarray, None, or tuple of ndarray and None, optional
  - A location into which the result is stored. If provided, it must have a shape that the inputs broadcast to. If not provided or `None`, a freshly-allocated array is returned. A tuple (possible only as a keyword argument) must have length equal to the number of outputs.
- `where`: array_like, optional
  - Values of True indicate to calculate the ufunc at that position, values of False indicate to leave the value in the output alone.

**kwargs:

For other keyword-only arguments, see the ufunc docs.

**Returns**

- `y`: ndarray
  - The corresponding tangent values. This is a scalar if `x` is a scalar.

#### Notes

If `out` is provided, the function writes the result into it, and returns a reference to `out`. (See Examples)

#### References


#### Examples
tanh(x, /[, out, where, casting, order, ...])  Compute hyperbolic tangent element-wise.

**tanh function**

(Shortest import: from brian2 import tanh)

brian2.units.unitsafefunctions.tanh(x, /, out=None, *, where=True, casting='same_kind', order='K', dtype=None, subok=True[, signature, extobj])

Compute hyperbolic tangent element-wise.

Equivalent to np.sinh(x)/np.cosh(x) or -1j * np.tan(1j*x).

**Parameters**

- **x**: array_like
  
  Input array.

- **out**: ndarray, None, or tuple of ndarray and None, optional
  
  A location into which the result is stored. If provided, it must have a shape that the inputs broadcast to. If not provided or None, a freshly-allocated array is returned. A tuple (possible only as a keyword argument) must have length equal to the number of outputs.

- **where**: array_like, optional
  
  Values of True indicate to calculate the ufunc at that position, values of False indicate to leave the value in the output alone.

**kwargs:

For other keyword-only arguments, see the ufunc docs.

**Returns**

- **y**: ndarray
  
  The corresponding hyperbolic tangent values. This is a scalar if x is a scalar.

**Notes**

If out is provided, the function writes the result into it, and returns a reference to out. (See Examples)
Brian 2 Documentation, Release 2.3

References
[R23], [R24]
Examples
>>> np.tanh((0, np.pi*1j, np.pi*1j/2))
array([ 0. +0.00000000e+00j, 0. -1.22460635e-16j,

0. +1.63317787e+16j])

>>> # Example of providing the optional output parameter illustrating
>>> # that what is returned is a reference to said parameter
>>> out2 = np.tanh([0.1], out1)
>>> out2 is out1
True
>>> # Example of ValueError due to provision of shape mis-matched `out`
>>> np.tanh(np.zeros((3,3)),np.zeros((2,2)))
Traceback (most recent call last):
File "<stdin>", line 1, in <module>
ValueError: operands could not be broadcast together with shapes (3,3) (2,2)

trace(a[, offset, axis1, axis2, dtype, out])

Return the sum along diagonals of the array.

trace function
(Shortest import: from brian2 import trace)
brian2.units.unitsafefunctions.trace(a, offset=0, axis1=0, axis2=1, dtype=None, out=None)
Return the sum along diagonals of the array.
If a is 2-D, the sum along its diagonal with the given offset is returned, i.e., the sum of elements a[i,
i+offset] for all i.
If a has more than two dimensions, then the axes specified by axis1 and axis2 are used to determine the 2-D
sub-arrays whose traces are returned. The shape of the resulting array is the same as that of a with axis1 and
axis2 removed.
Parameters a : array_like
Input array, from which the diagonals are taken.
offset : int, optional
Offset of the diagonal from the main diagonal. Can be both positive and negative.
Defaults to 0.
axis1, axis2 : int, optional
Axes to be used as the first and second axis of the 2-D sub-arrays from which the
diagonals should be taken. Defaults are the first two axes of a.
dtype : dtype, optional
Determines the data-type of the returned array and of the accumulator where the elements are summed. If dtype has the value None and a is of integer type of precision less
than the default integer precision, then the default integer precision is used. Otherwise,
the precision is the same as that of a.
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Chapter 6. brian2 package


out : ndarray, optional

Array into which the output is placed. Its type is preserved and it must be of the right shape to hold the output.

Returns sum_along_diagonals : ndarray

If a is 2-D, the sum along the diagonal is returned. If a has larger dimensions, then an array of sums along diagonals is returned.

See also:
diag(), diagonal(), diagflat()

Examples

```python
>>> np.trace(np.eye(3))
3.0
>>> a = np.arange(8).reshape((2,2,2))
>>> np.trace(a)
array([6, 8])
>>> a = np.arange(24).reshape((2,2,2,3))
>>> np.trace(a).shape
(2, 3)
```

where(condition, [x, y])

Return elements chosen from x or y depending on condition.

Note: When only condition is provided, this function is a shorthand for np.asarray(condition). nonzero(). Using nonzero() directly should be preferred, as it behaves correctly for subclasses. The rest of this documentation covers only the case where all three arguments are provided.

Parameters condition : array_like, bool

Where True, yield x, otherwise yield y.

x, y : array_like

Values from which to choose. x, y and condition need to be broadcastable to some shape.

Returns out : ndarray

An array with elements from x where condition is True, and elements from y elsewhere.

See also:

6.6. Subpackages
choose()

nonzero() The function that is called when x and y are omitted

Notes

If all the arrays are 1-D, where() is equivalent to:

```python
[xv if c else yv
 for c, xv, yv in zip(condition, x, y)]
```

Examples

```python
>>> a = np.arange(10)
>>> a
array([0, 1, 2, 3, 4, 5, 6, 7, 8, 9])
>>> np.where(a < 5, a, 10*a)
array([0, 1, 2, 3, 4, 50, 60, 70, 80, 90])
```

This can be used on multidimensional arrays too:

```python
>>> np.where([[True, False], [True, True]],
... [[1, 2], [3, 4]],
... [[9, 8], [7, 6]])
array([[1, 8],
       [3, 4]])
```

The shapes of x, y, and the condition are broadcast together:

```python
>>> x, y = np.ogrid[:3, :4]
>>> np.where(x < y, x, 10 + y)  # both x and 10+y are broadcast
array([[10, 0, 0, 0],
       [10, 11, 1, 1],
       [10, 11, 12, 2]])
```

```python
>>> a = np.array([[0, 1, 2],
... [0, 2, 4],
... [0, 3, 6]])
>>> np.where(a < 4, a, -1)  # -1 is broadcast
array([[0, 1, 2],
       [0, 2, -1],
       [0, 3, -1]])
```

```
wrap_function_to_method(func) Wraps a function so that it calls the corresponding method on the Quantities object (if called with a Quantities object as the first argument).

wrap_function_to_method function

(Shortest import: from brian2.units.unitsafefunctions import wrap_function_to_method)

brian2.units.unitsafefunctions.wrap_function_to_method(func)
Wraps a function so that it calls the corresponding method on the Quantities object (if called with a Quantities object as the first argument). All other arguments are left untouched.

### 6.6.16 utils package

Utility functions for Brian.

*Exported members:* `get_logger()`, `BrianLogger`, `std_silent`

#### arrays module

Helper module containing functions that operate on numpy arrays.

**Functions**

<table>
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<tr>
<th><code>calc_repeats</code>&lt;br&gt;<code>(delay)</code></th>
<th>Calculates offsets corresponding to an array, where repeated values are subsequently numbered, i.e.</th>
</tr>
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</table>

**calc_repeats function**

*(Shortest import: from brian2.utils.arrays import calc_repeats)*

brian2.utils.arrays.`calc_repeats`<br>`(delay)`

Calculates offsets corresponding to an array, where repeated values are subsequently numbered, i.e. if there n identical values, the returned array will have values from 0 to n-1 at their positions. The code is complex because tricks are needed for vectorisation.

This function is used in the Python `SpikeQueue` to calculate the offset array for the insertion of spikes with their respective delays into the queue and in the numpy code for synapse creation to calculate how many synapses for each source-target pair exist.

**Examples**

```python
>>> import numpy as np
>>> print(calc_repeats(np.array([7, 5, 7, 3, 7, 5])))
[0 0 1 0 2 1]
```

#### caching module

Module to support caching of function results to memory (used to cache results of parsing, generation of state update code, etc.). Provides the `cached` decorator.

**Classes**

<table>
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<tr>
<th><code>CacheKey</code></th>
<th>Mixin class for objects that will be used as keys for caching (e.g.</th>
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**CacheKey class**

*(Shortest import: from brian2.utils.caching import CacheKey)*
class brian2.utils.caching.CacheKey
    Bases: object

Mixin class for objects that will be used as keys for caching (e.g. Variable objects) and have to define a certain “identity” with respect to caching. This “identity” is different from standard Python hashing and equality checking: a Variable for example would be considered “identical” for caching purposes regardless which object (e.g. NeuronGroup) it belongs to (because this does not matter for parsing, creating abstract code, etc.) but this of course matters for the values it refers to and therefore for comparison of equality to other variables.

Classes that mix in the CacheKey class should re-define the _cache_irrelevant_attributes attribute to note all the attributes that should be ignored. The property _state_tuple will refer to a tuple of all attributes that were not excluded in such a way; this tuple will be used as the key for caching purposes.

Attributes

__cache_irrelevant_attributes__
Set of attributes that should not be considered for caching of state update code, etc.

Details

__cache_irrelevant_attributes__
Set of attributes that should not be considered for caching of state update code, etc.

Functions

cached(func)
Decorator to cache a function so that it will not be re-evaluated when called with the same arguments.

cached function

(Shortest import: from brian2.utils.caching import cached)

brian2.utils.caching.cached(func)
Decorator to cache a function so that it will not be re-evaluated when called with the same arguments. Uses the _hashable function to make arguments usable as a dictionary key even though they mutable (lists, dictionaries, etc.).

Parameters

func : function
    The function to decorate.

Returns

decorated : function
    The decorated function.

Notes

This is not a general-purpose caching decorator in any way comparable to functools.lru_cache or joblib’s caching functions. It is very simplistic (no maximum cache size, no normalization of calls, e.g. foo(3) and foo(x=3) are not considered equivalent function calls) and makes very specific assumptions for our use case. Most importantly, Variable objects are considered to be identical when they refer to the same object, even though the actually stored values might have changed.
environment module

Functions

running_from_ipython()  Check whether we are currently running under ipython.

running_from_ipython function

(Shortest import: from brian2.utils.environment import running_from_ipython)

brian2.utils.environment.running_from_ipython()

   Check whether we are currently running under ipython.

   Returns  ipython : bool

       Whether running under ipython or not.

filetools module

File system tools

Exported members:  ensure_directory, ensure_directory_of_file, in_directory, copy_directory

Classes

in_directory(new_dir)  Safely temporarily work in a subdirectory

in_directory class

(Shortest import: from brian2.utils.filetools import in_directory)

class  brian2.utils.filetools.in_directory(new_dir)

   Bases: object

   Safely temporarily work in a subdirectory

   Usage:

   with in_directory(directory):
       ... do stuff here

   Guarantees that the code in the with block will be executed in directory, and that after the block is completed
   we return to the original directory.

Functions

copy_directory(source, target)  Copies directory source to target.

copy_directory function

(Shortest import: from brian2.utils.filetools import copy_directory)

brian2.utils.filetools.copy_directory(source, target)
Copies directory source to target.

### ensure_directory function

*(Shortest import: from brian2.utils.filetools import ensure_directory)*

```python
d = ensure_directory(d)
```

Ensures that a given directory exists (creates it if necessary).

### ensure_directory_of_file function

*(Shortest import: from brian2.utils.filetools import ensure_directory_of_file)*

```python
d = ensure_directory_of_file(f)
```

Ensures that a directory exists for filename to go in (creates if necessary), and returns the directory path.

### logger module

Brian’s logging module.

### Preferences

Logging system preferences

```python
logging.console_log_level = 'INFO'
```

What log level to use for the log written to the console.

Has to be one of CRITICAL, ERROR, WARNING, INFO, DEBUG or DIAGNOSTIC.

```python
logging.delete_log_on_exit = True
```

Whether to delete the log and script file on exit.

If set to `True` (the default), log files (and the copy of the main script) will be deleted after the brian process has exited, unless an uncaught exception occurred. If set to `False`, all log files will be kept.

```python
logging.file_log = True
```

Whether to log to a file or not.

If set to `True` (the default), logging information will be written to a file. The log level can be set via the `logging.file_log_level` preference.

```python
logging.file_log_level = 'DIAGNOSTIC'
```

What log level to use for the log written to the log file.

In case file logging is activated (see `logging.file_log`), which log level should be used for logging. Has to be one of CRITICAL, ERROR, WARNING, INFO, DEBUG or DIAGNOSTIC.
logging.save_script = True

Whether to save a copy of the script that is run.

If set to True (the default), a copy of the currently run script is saved to a temporary location. It is deleted after a successful run (unless logging.delete_log_on_exit is False) but is kept after an uncaught exception occurred. This can be helpful for debugging, in particular when several simulations are running in parallel.

logging.std_redirection = True

Whether or not to redirect stdout/stderr to null at certain places.

This silences a lot of annoying compiler output, but will also hide error messages making it harder to debug problems. You can always temporarily switch it off when debugging. If logging.std_redirection_to_file is set to True as well, then the output is saved to a file and if an error occurs the name of this file will be printed.

logging.std_redirection_to_file = True

Whether to redirect stdout/stderr to a file.

If both logging.std_redirection and this preference are set to True, all standard output/error (most importantly output from the compiler) will be stored in files and if an error occurs the name of this file will be printed. If logging.std_redirection is True and this preference is False, then all standard output/error will be completely suppressed, i.e. neither be displayed nor stored in a file.

The value of this preference is ignore if logging.std_redirection is set to False.

Exported members: get_logger(), BrianLogger, std_silent

Classes

BrianLogger(name)

Convenience object for logging.

BrianLogger class

(Shortest import: from brian2 import BrianLogger)

class brian2.utils.logger.BrianLogger(name)

Bases: object

Convenience object for logging. Call get_logger() to get an instance of this class.

Parameters name : str

The name used for logging, normally the name of the module.

Attributes

__log_messages

Class attribute for remembering log messages that should only be displayed once

exception_occured

Class attribute to remember whether any exception occurred

file_handler

The logging.FileHandler responsible for logging to the temporary log file

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### Methods

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<th>Description</th>
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<td><code>debug(msg[, name_suffix, once])</code></td>
<td>Log a debug message.</td>
</tr>
<tr>
<td><code>diagnostic(msg[, name_suffix, once])</code></td>
<td>Log a diagnostic message.</td>
</tr>
<tr>
<td><code>error(msg[, name_suffix, once])</code></td>
<td>Log an error message.</td>
</tr>
<tr>
<td><code>info(msg[, name_suffix, once])</code></td>
<td>Log an info message.</td>
</tr>
<tr>
<td><code>initialize()</code></td>
<td>Initialize Brian’s logging system.</td>
</tr>
<tr>
<td><code>log_level_debug()</code></td>
<td>Set the log level to “debug”.</td>
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<tr>
<td><code>log_level_diagnostic()</code></td>
<td>Set the log level to “diagnostic”.</td>
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<tr>
<td><code>log_level_error()</code></td>
<td>Set the log level to “error”.</td>
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<td><code>log_level_info()</code></td>
<td>Set the log level to “info”.</td>
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<td><code>log_level_warn()</code></td>
<td>Set the log level to “warn”.</td>
</tr>
<tr>
<td><code>suppress_hierarchy(name[, filter_log_file])</code></td>
<td>Suppress all log messages in a given hierarchy.</td>
</tr>
<tr>
<td><code>suppress_name(name[, filter_log_file])</code></td>
<td>Suppress all log messages with a given name.</td>
</tr>
<tr>
<td><code>warn(msg[, name_suffix, once])</code></td>
<td>Log a warn message.</td>
</tr>
</tbody>
</table>

### Details

- **_log_messages**
  Class attribute for remembering log messages that should only be displayed once

- **exception_occured**
  Class attribute to remember whether any exception occurred

- **file_handler**
  The `logging.FileHandler` responsible for logging to the temporary log file

- **tmp_log**
  The name of the temporary log file (by default deleted after the run if no exception occurred), if any

- **tmp_script**
  The name of the temporary copy of the main script file (by default deleted after the run if no exception occurred), if any

- **debug** *(msg, name_suffix=None, once=False)*
  Log a debug message.

  **Parameters**
  - **msg** : str
    The message to log.
  - **name_suffix** : str, optional
    A suffix to add to the name, e.g. a class or function name.
  - **once** : bool, optional
    Whether this message should be logged only once and not repeated if sent another time.

- **diagnostic** *(msg, name_suffix=None, once=False)*
  Log a diagnostic message.
**Parameters**

*msg*: str

The message to log.

*name_suffix*: str, optional

A suffix to add to the name, e.g. a class or function name.

*once*: bool, optional

Whether this message should be logged only once and not repeated if sent another time.

**error** (*msg*, *name_suffix*=*None*, *once*=*False*)

Log an error message.

*Parameters*

*msg*: str

The message to log.

*name_suffix*: str, optional

A suffix to add to the name, e.g. a class or function name.

*once*: bool, optional

Whether this message should be logged only once and not repeated if sent another time.

**info** (*msg*, *name_suffix*=*None*, *once*=*False*)

Log an info message.

*Parameters*

*msg*: str

The message to log.

*name_suffix*: str, optional

A suffix to add to the name, e.g. a class or function name.

*once*: bool, optional

Whether this message should be logged only once and not repeated if sent another time.

**static initialize()**

Initialize Brian’s logging system. This function will be called automatically when Brian is imported.

**static log_level_debug()**

Set the log level to “debug”.

**static log_level_diagnostic()**

Set the log level to “diagnostic”.

**static log_level_error()**

Set the log level to “error”.

**static log_level_info()**

Set the log level to “info”.

**static log_level_warn()**

Set the log level to “warn”.

**static suppress_hierarchy** (*name*, *filter_log_file*=*False*)

Suppress all log messages in a given hierarchy.

*Parameters*

*name*: str

Suppress all log messages in the given *name* hierarchy. For example, specifying 'brian2' suppresses all messages logged by Brian, specifying 'brian2codegen' suppresses all messages generated by the code generation modules.
filter_log_file : bool, optional
Whether to suppress the messages also in the log file. Defaults to False meaning that suppressed messages are not displayed on the console but are still saved to the log file.

static suppress_name (name, filter_log_file=False)
Suppress all log messages with a given name.

Parameters name : str
Suppress all log messages ending in the given name. For example, specifying 'resolution_conflict' would suppress messages with names such as brian2.equations.codestrings.CodeString.resolution_conflict or brian2.equations.equations.Equations.resolution_conflict.

filter_log_file : bool, optional
Whether to suppress the messages also in the log file. Defaults to False meaning that suppressed messages are not displayed on the console but are still saved to the log file.

warn (msg, name_suffix=None, once=False)
Log a warn message.

Parameters msg : str
The message to log.

name_suffix : str, optional
A suffix to add to the name, e.g. a class or function name.

once : bool, optional
Whether this message should be logged only once and not repeated if sent another time.

Tutorials and examples using this

- Example frompapers/Rossant_et_al_2011bis

HierarchyFilter (name)
A class for suppressing all log messages in a subtree of the name hierarchy.

HierarchyFilter class

(Shortest import: from brian2.utils.logger import HierarchyFilter)

class brian2.utils.logger.HierarchyFilter (name)
Bases: object
A class for suppressing all log messages in a subtree of the name hierarchy. Does exactly the opposite as the logging.Filter class, which allows messages in a certain name hierarchy to pass.

Parameters name : str
The name hierarchy to suppress. See BrianLogger.suppress_hierarchy for details.
### Methods

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<tbody>
<tr>
<td><code>filter(record)</code></td>
<td>Filter out all messages in a subtree of the name hierarchy.</td>
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</tbody>
</table>

### Details

**filter** *(record)*

Filter out all messages in a subtree of the name hierarchy.

### LogCapture class

*(Shortest import: from brian2.utils.logger import LogCapture)*

```python
class brian2.utils.logger.LogCapture(log_list[, log_level])
```

A class for capturing log warnings. This class is used by `catch_logs` to allow testing in a similar way as with `warnings.catch_warnings`.

#### Methods

<table>
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<tr>
<th>Method</th>
<th>Description</th>
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<tbody>
<tr>
<td><code>emit(record)</code></td>
<td>Do whatever it takes to actually log the specified logging record.</td>
</tr>
<tr>
<td><code>install()</code></td>
<td>Install this handler to catch all warnings.</td>
</tr>
<tr>
<td><code>uninstall()</code></td>
<td>Uninstall this handler and re-connect the previously installed handlers.</td>
</tr>
</tbody>
</table>

#### Details

**emit** *(record)*

Do whatever it takes to actually log the specified logging record.

This version is intended to be implemented by subclasses and so raises a `NotImplementedError`.

**install()**

Install this handler to catch all warnings. Temporarily disconnect all other handlers.

**uninstall()**

Uninstall this handler and re-connect the previously installed handlers.

### NameFilter class

*(Shortest import: from brian2.utils.logger import NameFilter)*

```python
class brian2.utils.logger.NameFilter(name)
```

A class for suppressing log messages ending with a certain name.
Bases: object

A class for suppressing log messages ending with a certain name.

Parameters name : str

The name to suppress. See BrianLogger.suppress_name for details.

Methods

filter(record) Filter out all messages ending with a certain name.

Details

filter(record)
Filter out all messages ending with a certain name.

catch_logs([log_level]) A context manager for catching log messages.

catch_logs class

(Shortest import: from brian2.utils.logger import catch_logs)

class brian2.utils.logger.catch_logs(log_level=30)
Bases: object

A context manager for catching log messages. Use this for testing the messages that are logged. Defaults to catching warning/error messages and this is probably the only real use case for testing. Note that while this context manager is active, all log messages are suppressed. Using this context manager returns a list of (log level, name, message) tuples.

Parameters log_level : int or str, optional

The log level above which messages are caught.

Examples

```python
>>> logger = get_logger('brian2.logtest')
>>> logger.warn('An uncaught warning') # doctest: +SKIP
WARNING brian2.logtest: An uncaught warning
>>> with catch_logs() as l:
...    logger.warn('a caught warning')
...    print('l contains: %s' % l)
...  
l contains: [('WARNING', 'brian2.logtest', 'a caught warning')]
```

std_silent([alwaysprint]) Context manager that temporarily silences stdout and stderr but keeps the output saved in a temporary file and writes it if an exception is raised.
std_silent class

(Shortest import: from brian2 import std_silent)

```python
class brian2.utils.logger.std_silent(alwaysprint=False)
    Bases: object

    Context manager that temporarily silences stdout and stderr but keeps the output saved in a temporary file and writes it if an exception is raised.
```

Attributes

- `dest_stderr`
- `dest_stdout`

Methods

- `close()`

Details

- `dest_stderr = None`
- `dest_stdout = None`
- `classmethod close()`

Functions

- `brian_excepthook(exc_type, exc_obj, exc_tb)` Display a message mentioning the debug log in case of an uncaught exception.

brian_excepthook function

(Shortest import: from brian2.utils.logger import brian_excepthook)

```python
brian2.utils.logger.brian_excepthook(exc_type, exc_obj, exc_tb)
    Display a message mentioning the debug log in case of an uncaught exception.
```

- `clean_up_logging()` Shutdown the logging system and delete the debug log file if no error occured.

clean_up_logging function

(Shortest import: from brian2.utils.logger import clean_up_logging)

```python
brian2.utils.logger.clean_up_logging()
    Shutdown the logging system and delete the debug log file if no error occured.
```
**get_logger**

```python
get_logger(module_name)
```

Get an object that can be used for logging.

**get_logger function**

*(Shortest import: from brian2 import get_logger)*

```python
brian2.utils.logger.get_logger(module_name='brian2')
```

Get an object that can be used for logging.

- **Parameters**
  - `module_name`: str
    - The name used for logging, should normally be the module name as returned by `__name__`.

- **Returns**
  - `logger`: BrianLogger

**log_level_validator**

```python
log_level_validator(log_level)
```

**log_level_validator function**

*(Shortest import: from brian2.utils.logger import log_level_validator)*

```python
brian2.utils.logger.log_level_validator(log_level)
```

**stringtools module**

- **Exported members:**
  - indent, deindent, word_substitute, replace, get_identifiers, strip_empty_lines, stripped_deindented_lines, strip_empty_leading_and_trailing_lines, code_representation, SpellChecker

- **Classes**
  - `SpellChecker`:
    ```python
    class brian2.utils.stringtools.SpellChecker(words[, alphabet])
    ```

A simple spell checker that will be used to suggest the correct name if the user made a typo (e.g. for state variable names).

- **Parameters**
  - `words`: iterable of str
    - The known words
  - `alphabet`: iterable of str, optional
    - The allowed characters. Defaults to the characters allowed for identifiers, i.e. ascii characters, digits and the underscore.
**Methods**

- `edits1(word)`
- `known(words)`
- `known_edits2(word)`
- `suggest(word)`

**Details**

- `edits1(word)`
- `known(words)`
- `known_edits2(word)`
- `suggest(word)`

**Functions**

- `code_representation(code)`
  Returns a string representation for several different formats of code

  **code_representation function**

  *(Shortest import: from brian2.utils.stringtools import code_representation)*

  ```python
  brian2.utils.stringtools.code_representation(code)
  Returns a string representation for several different formats of code
  Formats covered include: - A single string - A list of statements/strings - A dict of strings - A dict of lists of statements/strings
  ```

- `deindent(text[, numtabs, spacespertab, ...])`
  Returns a copy of the string with the common indentation removed.

  **deindent function**

  *(Shortest import: from brian2.utils.stringtools import deindent)*

  ```python
  brian2.utils.stringtools.deindent(text, numtabs=None, spacespertab=4, docstring=False)
  Returns a copy of the string with the common indentation removed.
  Note that all tab characters are replaced with spacespertab spaces.
  If the docstring flag is set, the first line is treated differently and is assumed to be already correctly tabulated.
  If the numtabs option is given, the amount of indentation to remove is given explicitly and not the common indentation.
  ```

**Examples**

Normal strings, e.g. function definitions:
>>> multiline = """" def f(x):
... return x**2"
>>> print(multiline)
def f(x):
    return x**2
>>> print(deindent(multiline))
def f(x):
    return x**2
>>> print(deindent(multiline, docstring=True))
def f(x):
    return x**2
>>> print(deindent(multiline, numtabs=1, spacespertab=2))
def f(x):
    return x**2

Docstrings:

```python
>>> docstring = """"First docstring line.
... This line determines the indentation."""
>>> print(docstring)
First docstring line.
>>> print(deindent(docstring, docstring=True))
First docstring line.
```

---

```python
get_identifiers(expr[, include_numbers]) Return all the identifiers in a given string expr, that is everything that matches a programming language variable like expression, which is here implemented as the regexp \b[A-Za-z_] [A-Za-z0-9_] \b.
```

---

### get_identifiers function

*(Shortest import: from brian2.utils.stringtools import get_identifiers)*

brian2.utils.stringtools.get_identifiers(expr, include_numbers=False)

Return all the identifiers in a given string expr, that is everything that matches a programming language variable like expression, which is here implemented as the regexp \b[A-Za-z_] [A-Za-z0-9_] \b.

**Parameters**

- `expr` : str
  - The string to analyze
- `include_numbers` : bool, optional
  - Whether to include number literals in the output. Defaults to `False`.

**Returns**

- `identifiers` : set
  - A set of all the identifiers (and, optionally, numbers) in `expr`.

**Examples**

```python
>>> expr = '3-a*_b+c5+8+f(A -.3e-10, tau_2)*17'
>>> ids = get_identifiers(expr)
```

(continues on next page)
print(sorted(list(ids)))
['A', '_b', 'a', 'c5', 'f', 'tau_2']

ids = get_identifiers(expr, include_numbers=True)
print(sorted(list(ids)))
['.3e-10', '17', '3', '8', 'A', '_b', 'a', 'c5', 'f', 'tau_2']

---

**indent**

Indents a given multiline string.

**indent function**

*(Shortest import: from brian2.utils.stringtools import indent)*

brian2.utils.stringtools.indent(text[, numtabs, spacespertab, tab])
Indents a given multiline string.

By default, indentation is done using spaces rather than tab characters. To use tab characters, specify the tab character explicitly, e.g.:

```python
indent(text, tab='    ')
```

Note that in this case spacespertab is ignored.

**Examples**

```python
>>> multiline = """def f(x):
...     return x*x"""
>>> print(multiline)
def f(x):
    return x*x
>>> print(indent(multiline))
def f(x):
    return x*x
>>> print(indent(multiline, numtabs=2))
def f(x):
    return x*x
>>> print(indent(multiline, spacespertab=2))
def f(x):
    return x*x
>>> print(indent(multiline, tab='####'))
####def f(x):
####     return x*x
```

---

**replace**

Applies a dictionary of substitutions.

**replace function**

*(Shortest import: from brian2.utils.stringtools import replace)*

brian2.utils.stringtools.replace(s, substitutions)
Applies a dictionary of substitutions. Simpler than word_substitute, it does not attempt to only replace words.

---

6.6. Subpackages
strip_empty_leading_and_trailing_lines function

(Shortest import: from brian2.utils.stringtools import strip_empty_leading_and_trailing_lines)

brian2.utils.stringtools.strip_empty_leading_and_trailing_lines(s)

Removes all empty leading and trailing lines in the multi-line string s.

strip_empty_lines function

(Shortest import: from brian2.utils.stringtools import strip_empty_lines)

brian2.utils.stringtools.strip_empty_lines(s)

Removes all empty lines from the multi-line string s.

Examples

```python
>>> multiline = "")A string with
... an empty line.""
>>> print(strip_empty_lines(multiline))
A string with
an empty line.
```

stripped_deindented_lines function

(Shortest import: from brian2.utils.stringtools import stripped_deindented_lines)

brian2.utils.stringtools.stripped_deindented_lines(code)

Returns a list of the lines in a multi-line string, deindented.

word_substitute function

(Shortest import: from brian2.utils.stringtools import word_substitute)

brian2.utils.stringtools.word_substitute(expr, substitutions)

Applies a dict of word substitutions.
The dict substitutions consists of pairs (word, rep) where each word appearing in expr is replaced by rep. Here a ‘word’ means anything matching the regexp \bword\b.

### Examples

```python
given expr = 'a*_b+c5+8+f(A)'
given print(word_substitute(expr, {'a':'banana', 'f':'func'}))
result is banana*_b+c5+8+func(A)
```

### topsort module

*Exported members: topsort*

### Functions

<table>
<thead>
<tr>
<th>Function</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>topsort</strong></td>
<td>Topologically sort a graph</td>
</tr>
</tbody>
</table>

### topsort function

*(Shortest import: from brian2.utils.topsort import topsort)*

```python
brian2.utils.topsort.topsort(graph)
```

#### Topologically sort a graph

The graph should be of the form `{node: [list of nodes], ...}`.
This section is intended as a guide to how Brian functions internally for people developing Brian itself, or extensions to Brian. It may also be of some interest to others wishing to better understand how Brian works internally.

## 7.1 Coding guidelines

The basic principles of developing Brian are:

1. For the user, the emphasis is on making the package flexible, readable and easy to use. See the paper “The Brian simulator” in Frontiers in Neuroscience for more details.

2. For the developer, the emphasis is on keeping the package maintainable by a small number of people. To this end, we use stable, well maintained, existing open source packages whenever possible, rather than writing our own code.

### 7.1.1 Development workflow

Brian development is done in a git repository on github. Continuous integration testing is provided by travis CI, code coverage is measured with coveralls.io.

### The repository structure

Brian’s repository structure is very simple, as we are normally not supporting older versions with bugfixes or other complicated things. The master branch of the repository is the basis for releases, a release is nothing more than adding a tag to the branch, creating the tarball, etc. The master branch should always be in a deployable state, i.e. one should be able to use it as the base for everyday work without worrying about random breakages due to updates. To ensure this, no commit ever goes into the master branch without passing the test suite before (see below). The only exception to this rule is if a commit not touches any code files, e.g. additions to the README file or to the documentation (but even in this case, care should be taken that the documentation is still built correctly).

For every feature that a developer works on, a new branch should be opened (normally based on the master branch), with a descriptive name (e.g. add-numba-support). For developers that are members of “brian-team”, the branch
should ideally be created in the main repository. This way, one can easily get an overview over what the “core team” is currently working on. Developers who are not members of the team should fork the repository and work in their own repository (if working on multiple issues/features, also using branches).

**Implementing a feature/fixing a bug**

Every new feature or bug fix should be done in a dedicated branch and have an issue in the issue database. For bugs, it is important to not only fix the bug but also to introduce a new test case (see *Testing*) that makes sure that the bug will not ever be reintroduced by other changes. It is often a good idea to first define the test cases (that should fail) and then work on the fix so that the tests pass. As soon as the feature/fix is complete or as soon as specific feedback on the code is needed, open a “pull request” to merge the changes from your branch into *master*. In this pull request, others can comment on the code and make suggestions for improvements. New commits to the respective branch automatically appear in the pull request which makes it a great tool for iterative code review. Even more useful, travis will automatically run the test suite on the result of the merge. As a reviewer, always wait for the result of this test (it can take up to 30 minutes or so until it appears) before doing the merge and never merge when a test fails. As soon as the reviewer (someone from the core team and not the author of the feature/fix) decides that the branch is ready to merge, he/she can merge the pull request and optionally delete the corresponding branch (but it will be hidden by default, anyway).

**Useful links**

- The Brian repository: [https://github.com/brian-team/brian2](https://github.com/brian-team/brian2)
- Travis testing for Brian: [https://travis-ci.org/brian-team/brian2](https://travis-ci.org/brian-team/brian2)
- Code Coverage for Brian: [https://coveralls.io/github/brian-team/brian2](https://coveralls.io/github/brian-team/brian2)
- github’s documentation on pull requests: [https://help.github.com/articles/using-pull-requests](https://help.github.com/articles/using-pull-requests)

**7.1.2 Coding conventions**

**General recommendations**

Syntax is chosen as much as possible from the user point of view, to reflect the concepts as directly as possible. Ideally, a Brian script should be readable by someone who doesn’t know Python or Brian, although this isn’t always possible. Function, class and keyword argument names should be explicit rather than abbreviated and consistent across Brian. See Romain’s paper *On the design of script languages for neural simulators* for a discussion.

We use the PEP-8 coding conventions for our code. This in particular includes the following conventions:

- Use 4 spaces instead of tabs per indentation level
- Use spaces after commas and around the following binary operators: assignment (=), augmented assignment (+=, -= etc.), comparisons (==, <, >, !=, <>, <=, >=, in, not in, is, is not), Booleans (and, or, not).
- Do not use spaces around the equals sign in keyword arguments or when specifying default values. Neither put spaces immediately inside parentheses, brackets or braces, immediately before the open parenthesis that starts the argument list of a function call, or immediately before the open parenthesis that starts an indexing or slicing.
- Avoid using a backslash for continuing lines whenever possible, instead use Python’s implicit line joining inside parentheses, brackets and braces.
- The core code should only contain ASCII characters, no encoding has to be declared
• imports should be on different lines (e.g. do not use import sys, os) and should be grouped in the following order, using blank lines between each group:
  1. standard library imports
  2. third-party library imports (e.g. numpy, scipy, sympy, ...)
  3. brian imports

• Use absolute imports for everything outside of “your” package, e.g. if you are working in brian2.equations, import functions from the stringtools modules via from brian2.utils.stringtools import .... Use the full path when importing, e.g. do from brian2.units.fundamentalunits import seconds instead of from brian2 import seconds.

• Use “new-style” relative imports for everything in “your” package, e.g. in brian2.codegen.functions.py import the Function class as from .specifiers import Function.

• Do not use wildcard imports (from brian2 import *), instead import only the identifiers you need, e.g. from brian2 import NeuronGroup, Synapses. For packages like numpy that are used a lot, use import numpy as np. But note that the user should still be able to do something like from brian2 import * (and this style can also be freely used in examples and tests, for example). Modules always have to use the __all__ mechanism to specify what is being made available with a wildcard import. As an exception from this rule, the main brian2/__init__.py may use wildcard imports.

Python 2 vs. Python 3

Brian is written in Python 2 but runs on Python 3 using the 2to3 conversion tool (which is automatically applied if Brian is installed using the standard python setup.py install mechanism). To make this possible without too much effort, Brian no longer supports Python 2.5 and can therefore make use of a couple of forward-compatible (but backward-incompatible) idioms introduced in Python 2.6. The Porting to Python 3 book is available online and has a lot of information on these topics. Here are some things to keep in mind when developing Brian:

• If you are working with integers and using division, consider using // for flooring division (default behaviour for / in python 2) and switch the behaviour of / to floating point division by using from __future__ import division.

• If importing modules from the standard library (which have changed quite a bit from Python 2 to Python 3), only use simple import statements like import itertools instead of from itertools import izip–2to3 is otherwise unable to make the correct conversion.

• If you are using the print statement (which should only occur in tests, in particular doctests – always use the Logging framework if you want to present messages to the user otherwise), try “cheating” and use the functional style in Python 2, i.e. write print('some text') instead of print 'some text'. More complicated print statements should be avoided, e.g instead of print >>sys.stderr, 'Error message use sys.stderr.write('Error message\n') (or, again, use logging).

• Exception stacktraces look a bit different in Python 2 and 3: For non-standard exceptions, Python 2 only prints the Exception class name (e.g. DimensionMismatchError) whereas Python 3 prints the name including the module name (e.g. brian2.units.fundamentalunits.DimensionMismatchError). This will make doctests fail that match the exception message. In this case, write the doctest in the style of Python 2 but add the doctest directive #doctest: +IGNORE_EXCEPTION_DETAIL to the statement leading to the exception. This unfortunately has the side effect of also ignoring the text of the exception, but it will still fail for an incorrect exception type.

• If you write code reading and writing strings to files, make sure you make the distinction between bytes and unicode (see “separate binary data and strings”) In general, strings within Brian are unicode strings and only converted to bytes when reading from or writing to a file (or something like a network stream, for example).

• If you are sorting lists or dictionaries, have a look at “when sorting, use key instead of cmp”
• Make sure to define a __hash__ function for objects that define an __eq__ function (and to define it consistently). Python 3 is more strict about this, an object with __eq__ but without __hash__ is unhashable.

7.1.3 Representing Brian objects

__repr__ and __str__

Every class should specify or inherit useful __repr__ and __str__ methods. The __repr__ method should give the “official” representation of the object; if possible, this should be a valid Python expression, ideally allowing for eval(repr(x)) == x. The __str__ method on the other hand, gives an “informal” representation of the object. This can be anything that is helpful but does not have to be Python code. For example:

```python
>>> import numpy as np
>>> ar = np.array([1, 2, 3]) * mV
>>> print(ar) # uses __str__
[ 1.  2.  3.] mV
>>> ar # uses __repr__
array([ 1.,  2.,  3.]) * mvolt
```

If the representation returned by __repr__ is not Python code, it should be enclosed in <...>, e.g. a Synapses representation might be <Synapses object with 64 synapses>.

If you don’t want to make the distinction between __repr__ and __str__, simply define only a __repr__ function, it will be used instead of __str__ automatically (no need to write __str__ = __repr__). Finally, if you include the class name in the representation (which you should in most cases), use self.__class__.__name__ instead of spelling out the name explicitly – this way it will automatically work correctly for subclasses. It will also prevent you from forgetting to update the class name in the representation if you decide to rename the class.

LaTeX representations with sympy

Brian objects dealing with mathematical expressions and equations often internally use sympy. Sympy’s latex function does a nice job of converting expressions into LaTeX code, using fractions, root symbols, etc. as well as converting greek variable names into corresponding symbols and handling sub- and superscripts. For the conversion of variable names to work, they should use an underscore for subscripts and two underscores for superscripts:

```python
>>> from sympy import latex, Symbol
>>> tau_1__e = Symbol('tau_1__e')
>>> print(latex(tau_1__e))
\tau^{e}_{1}
```

Sympy’s printer supports formatting arbitrary objects, all they have to do is to implement a _latex method (no trailing underscore). For most Brian objects, this is unnecessary as they will never be formatted with sympy’s LaTeX printer. For some core objects, in particular the units, is is useful, however, as it can be reused in LaTeX representations for ipython (see below). Note that the _latex method should not return $ or \begin{equation} (sympy’s method includes a mode argument that wraps the output automatically).

Representations for ipython

“Old” ipython console

In particular for representations involving arrays or lists, it can be useful to break up the representation into chunks, or indent parts of the representation. This is supported by the ipython console’s “pretty printer”. To make this work for
a class, add a `_repr_pretty_(self, p, cycle)` (note the single underscores) method. You can find more information in the ipython documentation.

“New” ipython console (qtconsole and notebook)

The new ipython consoles, the qtconsole and the ipython notebook support a much richer set of representations for objects. As Brian deals a lot with mathematical objects, in particular the LaTeX and to a lesser extent the HTML formatting capabilities of the ipython notebook are interesting. To support LaTeX representation, implement a `_repr_latex_` method returning the LaTeX code (including \$$, \begin{equation} \) or similar). If the object already has a `_latex` method (see LaTeX representations with sympy above), this can be as simple as:

```python
def _repr_latex_(self):
    return sympy.latex(self, mode='inline')  # wraps the expression in \$ .. \$
```

The LaTeX rendering only supports a single mathematical block. For complex objects, e.g. `NeuronGroup` it might be useful to have a richer representation. This can be achieved by returning HTML code from `_repr_html_` – this HTML code is processed by MathJax so it can include literal LaTeX code that will be transformed before it is rendered as HTML. An object containing two equations could therefore be represented with a method like this:

```python
def _repr_html_(self):
    return '''
<h3> Equation 1 </h3>
{eq_1}
<h3> Equation 2 </h3>
{eq_2}'''.format(eq_1=sympy.latex(self.eq_1, mode='equation'),
                     eq_2=sympy.latex(self.eq_2, mode='equation'))
```

7.1.4 Defensive programming

One idea for Brian 2 is to make it so that it’s more likely that errors are raised rather than silently causing weird bugs. Some ideas in this line:

- `Synapses.source` should be stored internally as a weakref `Synapses._source`, and `Synapses.source` should be a computed attribute that dereferences this weakref. Like this, if the source object isn’t kept by the user, `Synapses` won’t store a reference to it, and so won’t stop it from being deallocated.

- We should write an automated test that takes a piece of correct code like:

```python
NeuronGroup(N, eqs, reset='V>Vt')
```

and tries replacing all arguments by nonsense arguments, it should always raise an error in this case (forcing us to write code to validate the inputs). For example, you could create a new `NonsenseObject` class, and do this:

```python
nonsense = NonsenseObject()
NeuronGroup(nonsense, eqs, reset='V>Vt')
NeuronGroup(N, nonsense, reset='V>Vt')
NeuronGroup(N, eqs, nonsense)
```

In general, the idea should be to make it hard for something incorrect to run without raising an error, preferably at the point where the user makes the error and not in some obscure way several lines later.

The preferred way to validate inputs is one that handles types in a Pythonic way. For example, instead of doing something like:
If not isinstance(arg, (float, int)):
    raise TypeError(...)

Do something like:

arg = float(arg)

(or use try/except to raise a more specific error). In contrast to the isinstance check it does not make any assumptions about the type except for its ability to be converted to a float.

This approach is particularly useful for numpy arrays:

arr = np.asarray(arg)

(or np.asanyarray if you want to allow for array subclasses like arrays with units or masked arrays). This approach has also the nice advantage that it allows all “array-like” arguments, e.g. a list of numbers.

### 7.1.5 Documentation

It is very important to maintain documentation. We use the Sphinx documentation generator tools. The documentation is all handwritten. Sphinx source files are stored in the docs_sphinx folder. The HTML files can be generated via the script dev/tools/docs/build_html_brian2.py and end up in the docs folder.

Most of the documentation is stored directly in the Sphinx source text files, but reference documentation for important Brian classes and functions are kept in the documentation strings of those classes themselves. This is automatically pulled from these classes for the reference manual section of the documentation. The idea is to keep the definitive reference documentation near the code that it documents, serving as both a comment for the code itself, and to keep the documentation up to date with the code.

The reference documentation includes all classes, functions and other objects that are defined in the modules and only documents them in the module where they were defined. This makes it possible to document a class like Quantity only in brian2.units.fundamentalunits and not additionally in brian2.units and brian2. This mechanism relies on the __module__ attribute, in some cases, in particular when wrapping a function with a decorator (e.g. check_units), this attribute has to be set manually:

```python
foo.__module__ = __name__
```

Without this manual setting, the function might not be documented at all or in the wrong module.

In addition to the reference, all the examples in the examples folder are automatically included in the documentation.

Note that you can directly link to github issues using :issue:`issue number`, e.g. writing :issue:`33` links to a github issue about running benchmarks for Brian 2: #33. This feature should rarely be used in the main documentation, reserve its use for release notes and important known bugs.

### Docstrings

Every module, class, method or function has to start with a docstring, unless it is a private or special method (i.e. starting with _ or __) and it is obvious what it does. For example, there is normally no need to document __str__ with “Return a string representation.”.

For the docstring format, we use the our own sphinx extension (in brian2/sphinxext) based on numpydoc, allowing to write docstrings that are well readable both in sourcecode as well as in the rendered HTML. We generally follow the format used by numpy.
When the docstring uses variable, class or function names, these should be enclosed in single backticks. Class and function/method names will be automatically linked to the corresponding documentation. For classes imported in the main brian2 package, you do not have to add the package name, e.g. writing `NeuronGroup` is enough. For other classes, you have to give the full path, e.g. `brian2.units.fundamentalunits.UnitRegistry`.

If it is clear from the context where the class is (e.g. within the documentation of `UnitRegistry`), consider using the ~ abbreviation: `~brian2.units.fundamentalunits.UnitRegistry` displays only the class name: `UnitRegistry`. Note that you do not have to enclose the exception name in a “Raises” or “Warns” section, or the class/method/function name in a “See Also” section in backticks, they will be automatically linked (putting backticks there will lead to incorrect display or an error message).

Inline source fragments should be enclosed in double backticks.

Class docstrings follow the same conventions as method docstrings and should document the `__init__` method, the `__init__` method itself does not need a docstring.

### Documenting functions and methods

The docstring for a function/method should start with a one-line description of what the function does, without referring to the function name or the names of variables. Use a “command style” for this summary, e.g. “Return the result.” instead of “Returns the result.” If the signature of the function cannot be automatically extracted because of an decorator (e.g. `check_units()`), place a signature in the very first row of the docstring, before the one-line description.

For methods, do not document the `self` parameter, nor give information about the method being static or a class method (this information will be automatically added to the documentation).

### Documenting classes

Class docstrings should use the same “Parameters” and “Returns” sections as method and function docstrings for documenting the `__init__` constructor. If a class docstring does not have any “Attributes” or “Methods” section, these sections will be automatically generated with all documented (i.e. having a docstring), public (i.e. not starting with _) attributes respectively methods of the class. Alternatively, you can provide these sections manually. This is useful for example in the `Quantity` class, which would otherwise include the documentation of many `ndarray` methods, or when you want to include documentation for functions like `__getitem__` which would otherwise not be documented. When specifying these sections, you only have to state the names of documented methods/attributes but you can also provide direct documentation. For example:

```
Attributes
--------
foo
bar
baz

This is a description.
```

This can be used for example for class or instance attributes which do not have “classical” docstrings. However, you can also use a special syntax: When defining class attributes in the class body or instance attributes in `__init__` you can use the following variants (here shown for instance attributes):

```python
def __init__(a, b, c):
    #: The docstring for the instance attribute a.
    #: Can also span multiple lines
    self.a = a

    self.b = b #: The docstring for self.b (only one line).
```

(continues on next page)
Long example of a function docstring

This is a very long docstring, showing all the possible sections. Most of the time no See Also, Notes or References section is needed:

```python
def foo(var1, var2, long_var_name='hi'):
    """
    A one-line summary that does not use variable names or the function name.

    Several sentences providing an extended description. Refer to variables using back-ticks, e.g. `var1`.

    Parameters
    ----------
    var1 : array_like
        Array_like means all those objects -- lists, nested lists, etc. -- that can be converted to an array. We can also refer to variables like `var1`.
    var2 : int
        The type above can either refer to an actual Python type (e.g. `int`), or describe the type of the variable in more detail, e.g. `array` or `array_like`.
    Long_variable_name : {'hi', 'ho'}, optional
        Choices in brackets, default first when optional.

    Returns
    -------
    describe : type
        Explanation
    output : type
        Explanation
    tuple : type
        Explanation
    items : type
        even more explaining

    Raises
    -----:
    BadException
        Because you shouldn’t have done that.

    See Also
    --------
    otherfunc : relationship (optional)
    newfunc : Relationship (optional), which could be fairly long, in which case the line wraps here.
    thirdfunc, fourthfunc, fifthfunc

    Notes
    -----:
    Notes about the implementation algorithm (if needed).
    """
```

(continues on next page)
This can have multiple paragraphs.

You may include some math:

.. math:: X(e^{j\omega}) = x(n)e^{-j\omega n}

And even use a greek symbol like :math:`\omega` inline.

References
---------
Cite the relevant literature, e.g. [1]_. You may also cite these references in the notes section above.


Examples
--------
These are written in doctest format, and should illustrate how to use the function.

```python
>>> a=[1,2,3]
>>> print([x + 3 for x in a])
[4, 5, 6]
>>> print("a\nb")
a
b
```

pass

### 7.1.6 Logging

For a description of logging from the users point of view, see Logging.

Logging in Brian is based on the logging module in Python’s standard library.

Every brian module that needs logging should start with the following line, using the `get_logger()` function to get an instance of `BrianLogger`:

```python
logger = get_logger(__name__)
```

In the code, logging can then be done via:

```python
logger.diagnostic('A diagnostic message')
logger.debug('A debug message')
logger.info('An info message')
logger.warn('A warning message')
logger.error('An error message')
```

If a module logs similar messages in different places or if it might be useful to be able to suppress a subset of messages
in a module, add an additional specifier to the logging command, specifying the class or function name, or a method name including the class name (do not include the module name, it will be automatically added as a prefix):

```python
logger.debug('A debug message', 'CodeString')
logger.debug('A debug message', 'NeuronGroup.update')
logger.debug('A debug message', 'reinit')
```

If you want to log a message only once, e.g. in a function that is called repeatedly, set the optional `once` keyword to `True`:

```python
logger.debug('Will only be shown once', once=True)
logger.debug('Will only be shown once', once=True)
```

The output of debugging looks like this in the log file:

```
```

and like this on the console (if the log level is set to “debug”):

```
DEBUG A debug message [brian2.equations.equations.CodeString]
```

**Log level recommendations**

**diagnostic**  Low-level messages that are not of any interest to the normal user but useful for debugging Brian itself. A typical example is the source code generated by the code generation module.

**debug**  Messages that are possibly helpful for debugging the user’s code. For example, this shows which objects were included in the network, which clocks the network uses and when simulations start and stop.

**info**  Messages which are not strictly necessary, but are potentially helpful for the user. In particular, this will show messages about the chosen state updater and other information that might help the user to achieve better performance and/or accuracy in the simulations (e.g. using (event-driven) in synaptic equations, avoiding incompatible dt values between `TimedArray` and the `NeuronGroup` using it, ...)

**warn**  Messages that alert the user to a potential mistake in the code, e.g. two possible solutions for an identifier in an equation. It can also be used to make the user aware that he/she is using an experimental feature, an unsupported compiler or similar. In this case, normally the `once=True` option should be used to raise this warning only once. As a rule of thumb, “common” scripts like the examples provided in the examples folder should normally not lead to any warnings.

**error**  This log level is not used currently in Brian, an exception should be raised instead. It might be useful in “meta-code”, running scripts and catching any errors that occur.

The default log level shown to the user is `info`. As a general rule, all messages that the user sees in the default configuration (i.e., `info` and `warn` level) should be avoidable by simple changes in the user code, e.g. the renaming of variables, explicitly specifying a state updater instead of relying on the automatic system, adding (clock-driven)/(event-driven) to synaptic equations, etc.

**Testing log messages**

It is possible to test whether code emits an expected log message using the `catch_logs` context manager. This is normally not necessary for debug and info messages, but should be part of the unit tests for warning messages (`catch_logs` by default only catches warning and error messages):
with catch_logs() as logs:
    # code that is expected to trigger a warning
    # ...
    assert len(logs) == 1
    # logs contains tuples of (log level, name, message)
    assert logs[0][0] == 'WARNING' and logs[0][1].endswith('warning_type')

7.1.7 Testing

Brian uses the pytest package for its testing framework.

Running the test suite

The pytest tool automatically finds tests in the code. However, to deal with the different code generation targets, and correctly set up tests for standalone mode, it is recommended to use Brian’s builtin test function that calls pytest appropriately:

```python
>>> import brian2
>>> brian2.test()
```

By default, this runs the test suite for all available (runtime) code generation targets. If you only want to test a specific target, provide it as an argument:

```python
>>> brian2.test('numpy')
```

If you want to test several targets, use a list of targets:

```python
>>> brian2.test(['weave', 'cython'])
```

In addition to the tests specific to a code generation target, the test suite will also run a set of independent tests (e.g. parsing of equations, unit system, utility functions, etc.). To exclude these tests, set the `test_codegen_independent` argument to False. Not all available tests are run by default, tests that take a long time are excluded. To include these, set `long_tests` to True.

To run the C++ standalone tests, you have to set the `test_standalone` argument to the name of a standalone device. If you provide an empty argument for the runtime code generation targets, you will only run the standalone tests:

```python
>>> brian2.test([], test_standalone='cpp_standalone')
```

Writing tests

Generally speaking, we aim for a 100% code coverage by the test suite. Less coverage means that some code paths are never executed so there’s no way of knowing whether a code change broke something in that path.

Unit tests

The most basic tests are unit tests, tests that test one kind of functionality or feature. To write a new unit test, add a function called `test_...` to one of the `test_...` files in the `brian2.tests` package. Test files should roughly correspond to packages, test functions should roughly correspond to tests for one function/method/feature. In the test functions, use assertions that will raise an `AssertionError` when they are violated, e.g.
G = NeuronGroup(42, model='dv/dt = -v / (10*ms) : 1')
assert len(G) == 42

When comparing arrays, use the array_equal() function from numpy.testing.utils which takes care of comparing types, shapes and content and gives a nicer error message in case the assertion fails. Never make tests depend on external factors like random numbers – tests should always give the same result when run on the same codebase. You should not only test the expected outcome for the correct use of functions and classes but also that errors are raised when expected. For that you can use pytest's raises function with which you can define a block of code that should raise an exception of a certain type:

```python
with pytest.raises(DimensionMismatchError):
    3*volt + 5*second
```

You can also check whether expected warnings are raised, see the documentation of the logging mechanism for details.

For simple functions, doctests (see below) are a great alternative to writing classical unit tests.

By default, all tests are executed for all selected runtime code generation targets (see Running the test suite above). This is not useful for all tests, some basic tests that for example test equation syntax or the use of physical units do not depend on code generation and need therefore not to be repeated. To execute such tests only once, they can be annotated with a codegen_independent marker, using the mark decorator:

```python
import pytest
from brian2 import NeuronGroup

@ pytest.mark.codegen_independent
def test_simple():
    # Test that the length of a NeuronGroup is correct
    group = NeuronGroup(5, '')
    assert len(group) == 5
```

Tests that are not “codegen-independent” are by default only executed for the runtimes device, i.e. not for the cpp_standalone device, for example. However, many of those tests follow a common pattern that is compatible with standalone devices as well: they set up a network, run it, and check the state of the network afterwards. Such tests can be marked as standalone_compatible, using the mark decorator in the same way as for codegen_independent tests:

```python
import pytest
from numpy.testing.utils import assert_equal
from brian2 import *

@ pytest.mark.standalone_compatible
def test_simple_run():
    # Check that parameter values of a neuron don't change after a run
    group = NeuronGroup(5, 'v : volt')
    group.v = 'i*mV'
    run(1*ms)
    assert_equal(group.v[:,], np.arange(5)*mV)
```

Tests that have more than a single run function but are otherwise compatible with standalone mode (e.g. they don’t need access to the number of synapses or results of the simulation before the end of the simulation), can be marked as standalone_compatible and multiple_runs. They then have to use an explicit device.build(...) call of the form shown below:

```python
import pytest
from numpy.testing.utils import assert_equal
from brian2 import *
```

(continues on next page)
@pytest.mark.standalone_compatible
@ pytest.mark.multiple_runs

def test_multiple_runs():
    # Check that multiple runs advance the clock as expected
    group = NeuronGroup(5, 'v : volt')
    mon = StateMonitor(group, 'v', record=True)
    run(1 * ms)
    run(1 * ms)
    device.build(direct_call=False, **device.build_options)
    assert_equal(defaultclock.t, 2 * ms)
    assert_equal(mon.t[0], 0 * ms)
    assert_equal(mon.t[-1], 2 * ms - defaultclock.dt)

Tests can also be written specifically for a standalone device (they then have to include the set_device call and possibly the build call explicitly). In this case tests have to be annotated with the name of the device (e.g. 'cpp_standalone') and with 'standalone_only' to exclude this test from the runtime tests. Such code would look like this for a single run() call, i.e. using the automatic “build on run” feature:

```python
import pytest
from brian2 import *

@ pytest.mark.cpp_standalone
@ pytest.mark.standalone_only

def test_cpp_standalone():
    set_device('cpp_standalone', directory=None)
    # set up simulation
    # run simulation
    run(...)
    # check simulation results
```

If the code uses more than one run() statement, it needs an explicit build call:

```python
import pytest
from brian2 import *

@ pytest.mark.cpp_standalone
@ pytest.mark.standalone_only

def test_cpp_standalone():
    set_device('cpp_standalone', build_on_run=False)
    # set up simulation
    # run simulation
    run(...)
    # do something
    # run again
    run(...)
    device.build(directory=None)
    # check simulation results
```
Summary

<table>
<thead>
<tr>
<th>@pytest.mark marker</th>
<th>Executed for devices</th>
<th>explicit use of device</th>
</tr>
</thead>
<tbody>
<tr>
<td>codegen_independent</td>
<td>independent of devices</td>
<td>none</td>
</tr>
<tr>
<td>none</td>
<td>Runtime targets</td>
<td>none</td>
</tr>
<tr>
<td>standalone_compatible</td>
<td>Runtime and standalone</td>
<td>none</td>
</tr>
<tr>
<td>standalone_compatible, multiple_runs</td>
<td>Runtime and standalone</td>
<td>device.build(direct_call=False, **device.build_options)</td>
</tr>
<tr>
<td>cpp_standalone, standalone_only</td>
<td>C++ standalone device</td>
<td>set_device('cpp_standalone') ... device.build(directory=None)</td>
</tr>
<tr>
<td>my_device, standalone_only</td>
<td>“My device”</td>
<td>set_device('my_device') ... device.build(directory=None)</td>
</tr>
</tbody>
</table>

Doctests

Doctests are executable documentation. In the Examples block of a class or function documentation, simply write code copied from an interactive Python session (to do this from ipython, use %doctestmode), e.g.:

```python
>>> from brian2.utils.stringtools import word_substitute
>>> expr = 'a*_b+c5+8+f(A)'
>>> print(word_substitute(expr, {'a':'banana', 'f':'func'}))
banana*_b+c5+8+func(A)
```

During testing, the actual output will be compared to the expected output and an error will be raised if they don’t match. Note that this comparison is strict, e.g. trailing whitespace is not ignored. There are various ways of working around some problems that arise because of this expected exactness (e.g. the stacktrace of a raised exception will never be identical because it contains file names), see the doctest documentation for details.

Doctests can (and should) not only be used in docstrings, but also in the hand-written documentation, making sure that the examples actually work. To turn a code example into a doc test, use the .. doctest:: directive, see Equations for examples written as doctests. For all doctests, everything that is available after from brian2 import * can be used directly. For everything else, add import statements to the doctest code or – if you do not want the import statements to appear in the document – add them in a .. testsetup:: block. See the documentation for Sphinx’s doctest extension for more details.

Doctests are a great way of testing things as they not only make sure that the code does what it is supposed to do but also that the documentation is up to date!

Correctness tests

[These do not exist yet for brian2]. Unit tests test a specific function or feature in isolation. In addition, we want to have tests where a complex piece of code (e.g. a complete simulation) is tested. Even if it is sometimes impossible to really check whether the result is correct (e.g. in the case of the spiking activity of a complex network), a useful check is also whether the result is consistent. For example, the spiking activity should be the same when using code generation for Python or C++. Or, a network could be pickled before running and then the result of the run could be compared to a second run that starts from the unpickled network.

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7.2 Units

7.2.1 Casting rules

In Brian 1, a distinction is made between scalars and numpy arrays (including scalar arrays): Scalars could be multiplied with a unit, resulting in a Quantity object whereas the multiplication of an array with a unit resulted in a (unitless) array. Accordingly, scalars were considered as dimensionless quantities for the purpose of unit checking (e.g., $1 + 1 \text{ mV}$ raised an error) whereas arrays were not (e.g. $\text{array}(1) + 1 \text{ mV}$ resulted in $1.001$ without any errors). Brian 2 no longer makes this distinction and treats both scalars and arrays as dimensionless for unit checking and make all operations involving quantities return a quantity:

```python
>>> 1 + 1*second
Traceback (most recent call last):
...  
DimensionMismatchError: Cannot calculate 1. s + 1, units do not match (units are, → second and 1).
```

```python
>>> np.array([1]) + 1*second
Traceback (most recent call last):
...  
DimensionMismatchError: Cannot calculate 1. s + [1], units do not match (units are, → second and 1).
```

```python
>>> 1*second + 1*second
2. * second
```

```python
>>> np.array([1])*second + 1*second
array([ 2.]) * second
```

As one exception from this rule, a scalar or array $0$ is considered as having “any unit”, i.e. $0 + 1 * \text{second}$ will result in $1 * \text{second}$ without a dimension mismatch error and $0 == 0 * \text{mV}$ will evaluate to True. This seems reasonable from a mathematical viewpoint and makes some sources of error disappear. For example, the Python builtin sum (not numpy’s version) adds the value of the optional argument start, which defaults to $0$, to its main argument. Without this exception, `sum([1 * \text{mV}, 2 * \text{mV}])` would therefore raise an error.

The above rules also apply to all comparisons (e.g. $==$ or $<$) with one further exception: $\text{inf}$ and $-\text{inf}$ also have “any unit”, therefore an expression like $v <= \text{inf}$ will never raise an exception (and always return True).

7.2.2 Functions and units

**ndarray methods**

All methods that make sense on quantities should work, i.e. they check for the correct units of their arguments and return quantities with units were appropriate. Most of the methods are overwritten using thin function wrappers:

**wrap_function_keep_dimension**: Strips away the units before giving the array to the method of ndarray, then reattaches the unit to the result (examples: sum, mean, max)

**wrap_function_change_dimension**: Changes the dimensions in a simple way that is independent of function arguments, the shape of the array, etc. (examples: sqrt, var, power)

**wrap_function_dimensionless**: Raises an error if the method is called on a quantity with dimensions (i.e. it works on dimensionless quantities).

**List of methods**

all, any, argmax, argsort, clip, compress, conj, conjugate, copy, cumsum, diagonal, dot, dump, dumps, fill, flatten, getfield, item, itemset, max, mean, min, newbyteorder, nonzero,
prod, ptp, put, ravel, repeat, reshape, round, searchsorted, setasflat, setfield, setflags, sort, squeeze, std, sum, take, tolist, trace, transpose, var, view

Notes

- Methods directly working on the internal data buffer (setfield, getfield, newbyteorder) ignore the dimensions of the quantity.
- The type of a quantity cannot be int, therefore astype does not quite work when trying to convert the array into integers.
- choose is only defined for integer arrays and therefore does not work.
- tostring and tofile only return/save the pure array data without the unit (but you can use dump or dumps to pickle a quantity array).
- resize does not work: ValueError: cannot resize this array: it does not own its data.
- cumprod would result in different dimensions for different elements and is therefore forbidden.
- item returns a pure Python float by definition.
- itemset does not check for units.

Numpy ufuncs

All of the standard numpy ufuncs (functions that operate element-wise on numpy arrays) are supported, meaning that they check for correct units and return appropriate arrays. These functions are often called implicitly, for example when using operators like < or **.

Math operations: add, subtract, multiply, divide, logaddexp, logaddexp2, true_divide, floor_divide, negative, power, remainder, mod, fmod, absolute, rint, sign, conj, conjugate, exp, exp2, log, log2, log10, expm1, log1p, sqrt, square, reciprocal, ones_like

Trigonometric functions: sin, cos, tan, arcsin, arccos, arctan, arctan2, hypot, sinh, cosh, tanh, arcsinh, arccosh, arctanh, deg2rad, rad2deg

Bitwise functions: bitwise_and, bitwise_or, bitwise_xor, invert, left_shift, right_shift

Comparison functions: greater, greater_equal, less, less_equal, not_equal, equal, logical_and, logical_or, logical_xor, logical_not, maximum, minimum

Floating functions: isreal, iscomplex, isfinite, isinf, isnan, floor, ceil, trunc, fmod

Not taken care of yet: signbit, copysign, nextafter, modf, ldexp, frexp

Notes

- Everything involving log or exp, as well as trigonometric functions only works on dimensionless array (for arctan2 and hypot this is questionable, though).
- Unit arrays can only be raised to a scalar power, not to an array of exponents as this would lead to differing dimensions across entries. For simplicity, this is enforced even for dimensionless quantities.
- Bitwise functions never works on quantities (numpy will by itself throw a TypeError because they are floats not integers).
- All comparisons only work for matching dimensions (with the exception of always allowing comparisons to 0) and return a pure boolean array.
- All logical functions treat quantities as boolean values in the same way as floats are treated as boolean: Any non-zero value is True.
Numpy functions

Many numpy functions are functional versions of ndarray methods (e.g. \texttt{mean}, \texttt{sum}, \texttt{clip}). They therefore work automatically when called on quantities, as numpy propagates the call to the respective method.

There are some functions in numpy that do not propagate their call to the corresponding method (because they use \texttt{np.asarray} instead of \texttt{np.asanyarray}, which might actually be a bug in numpy): \texttt{trace}, \texttt{diagonal}, \texttt{ravel}, \texttt{dot}. For these, wrapped functions in \texttt{unitsafefunctions.py} are provided.

Wrapped numpy functions in \texttt{unitsafefunctions.py}

These functions are thin wrappers around the numpy functions to correctly check for units and return quantities when appropriate:

\texttt{log, exp, sin, cos, tan, arcsin, arccos, arctan, sinh, cosh, tanh, arcsinh, arccosh, arctanh, diagonal, ravel, trace, dot}

Numpy functions that work unchanged

This includes all functional counterparts of the methods mentioned above (with the exceptions mentioned above). Some other functions also work correctly, as they are only using functions/methods that work with quantities:

- \texttt{linspace, diff, digitize}\footnote{But does not care about the units of its input.}
- \texttt{trim_zeros, fliplr, flipud, roll, rot90, shuffle}
- \texttt{corrcoef}\footnote{But does not care about the units of its input.}

Numpy functions that return a pure numpy array instead of quantities

- \texttt{arange}
- \texttt{cov}
- \texttt{random.permutation}
- \texttt{histogram, histogram2d}
- \texttt{cross, inner, outer}
- \texttt{where}

Numpy functions that do something wrong

- \texttt{insert, delete} (return a quantity array but without units)
- \texttt{correlate} (returns a quantity with wrong units)
- \texttt{histogramdd} (raises a DimensionMismatchError)

User-defined functions and units

For performance and simplicity reasons, code within the Brian core does not use Quantity objects but unitless numpy arrays instead. See \textit{Adding support for new functions} for details on how to make use user-defined functions with Brian’s unit system.

\footnote{But does not care about the units of its input.}
7.3 Equations and namespaces

7.3.1 Equation parsing

Parsing is done via pyparsing, for now find the grammar at the top of the brian2.equations.equations file.

7.3.2 Variables

Each Brian object that saves state variables (e.g. NeuronGroup, Synapses, StateMonitor) has a variables attribute, a dictionary mapping variable names to Variable objects (in fact a Variables object, not a simple dictionary). Variable objects contain information about the variable (name, dtype, units) as well as access to the variable’s value via a get_value method. Some will also allow setting the values via a corresponding set_value method. These objects can therefore act as proxies to the variables’ “contents”.

Variable objects provide the “abstract namespace” corresponding to a chunk of “abstract code”, they are all that is needed to check for syntactic correctness, unit consistency, etc.

7.3.3 Namespaces

The namespace attribute of a group can contain information about the external (variable or function) names used in the equations. It specifies a group-specific namespace used for resolving names in that group. At run time, this namespace is combined with a “run namespace”. This namespace is either explicitly provided to the Network.run method, or the implicit namespace consisting of the locals and globals around the point where the run function is called is used. This namespace is then passed down to all the objects via Network.before_fun which calls all the individual BrianObject.before_run methods with this namespace.

7.4 Variables and indices

7.4.1 Introduction

To be able to generate the proper code out of abstract code statements, the code generation process has to have access to information about the variables (their type, size, etc.) as well as to the indices that should be used for indexing arrays (e.g. a state variable of a NeuronGroup will be indexed differently in the NeuronGroup state updater and in synaptic propagation code). Most of this information is stored in the variables attribute of a VariableOwner (this includes NeuronGroup, Synapses, PoissonGroup and everything else that has state variables). The variables attribute can be accessed as a (read-only) dictionary, mapping variable names to Variable objects storing the information about the respective variable. However, it is not a simple dictionary but an instance of the Variables class. Let’s have a look at its content for a simple example:

```python
>>> tau = 10*ms
>>> G = NeuronGroup(10, 'dv/dt = -v / tau : volt')
>>> for name, var in sorted(G.variables.items()):
...     print('|$s : $s' % (name, var))
...
N : <Constant(dimensions=Dimension(), dtype=int64, scalar=True, constant=True, read_only=True)>
dt : <ArrayVariable(dimensions=second, dtype=float, scalar=True, constant=True, read_only=True)>
i : <ArrayVariable(dimensions=Dimension(), dtype=int32, scalar=False, constant=True, read_only=True)>
```

(continues on next page)
The state variable $v$ we specified for the NeuronGroup is represented as an ArrayVariable, all the other variables were added automatically. There’s another array $i$, the neuronal indices (simply an array of integers from 0 to 9), that is used for string expressions involving neuronal indices. The constant $N$ represents the total number of neurons.

At the first sight it might be surprising that $t$, the current time of the clock and $dt$, its timestep, are ArrayVariable objects as well. This is because those values can change during a run (for $t$) or between runs (for $dt$), and storing them as arrays with a single value (note the scalar=True) is the easiest way to share this value – all code accessing it only needs a reference to the array and can access its only element.

The information stored in the Variable objects is used to do various checks on the level of the abstract code, i.e. before any programming language code is generated. Here are some examples of errors that are caught this way:

```python
>>> G.v = 3*ms  # G.variables['v'].unit is volt
DimensionMismatchError: v should be set with a value with units volt, but got 3. ms
  (unit is second).
>>> G.N = 5  # G.variables['N'] is read-only
TypeError: Variable N is read-only
```

### 7.4.2 Creating variables

Each variable that should be accessible as a state variable and/or should be available for use in abstract code has to be created as a Variable. For this, first a Variables container with a reference to the group has to be created, individual variables can then be added using the various add_... methods:

```python
self.variables = Variables(self)
sel.variables.add_array('an_array', unit=volt, size=100)
sel.variables.add_constant('N', unit=Unit(1), value=self._N, dtype=np.int32)
sel.variables.create_clock_variables(self.clock)
```

As an additional argument, array variables can be specified with a specific index (see Indices below).

### 7.4.3 References

For each variable, only one Variable object exists even if it is used in different contexts. Let’s consider the following example:

```python
>>> G = NeuronGroup(5, 'dv/dt = -v / tau : volt', threshold='v > 1', reset='v = 0', name='neurons')
>>> subG = G[2:]  
>>> S = Synapses(G, G, on_pre='v+=1*mV', name='synapses')
>>> S.connect()
```

All allow an access to the state variable $v$ (note the different shapes, these arise from the different indices used, see below):
In all of these cases, the Variables object stores references to the same ArrayVariable object:

```python
>>> id(G.variables['v'])
108610960
>>> id(subG.variables['v'])
108610960
>>> id(S.variables['v'])
108610960
```

Such a reference can be added using Variables.add_reference, note that the name used for the reference is not necessarily the same as in the original group, e.g. in the above example S.variables also stores references to v under the names v_pre and v_post.

### 7.4.4 Indices

In subgroups and especially in synapses, the transformation of abstract code into executable code is not straightforward because it can involve variables from different contexts. Here is a simple example:

```python
>>> G = NeuronGroup(5, 'dv/dt = -v / tau : volt', threshold='v > 1', reset='v = 0')
>>> S = Synapses(G, G, 'w : volt', on_pre='v+=w')
```

The seemingly trivial operation v+=w involves the variable v of the NeuronGroup and the variable w of the Synapses object which have to be indexed in the appropriate way. Since this statement is executed in the context of S, the variable indices stored there are relevant:

```python
>>> S.variables.indices['w']
'_idx'
>>> S.variables.indices['v']
'_postsynaptic_idx'
```

The index _idx has a special meaning and always refers to the “natural” index for a group (e.g. all neurons for a NeuronGroup, all synapses for a Synapses object, etc.). All other indices have to refer to existing arrays:

```python
>>> S.variables['_postsynaptic_idx']
<DynamicArrayVariable(dimensions=Dimension(), dtype=<class 'numpy.int32'>, scalar=False, constant=True, read_only=True)>
```

In this case, _postsynaptic_idx refers to a dynamic array that stores the postsynaptic targets for each synapse (since it is an array itself, it also has an index. It is defined for each synapse so its index is _idx – in fact there is currently no support for an additional level of indirection in Brian: a variable representing an index has to have _idx as its own index). Using this index information, the following C++ code (slightly simplified) is generated:

```c++
for(int _spiking_synapse_idx=0;
    _spiking_synapse_idx<_num_spiking_synapses;
    _spiking_synapse_idx++)
{
    const int _idx = _spiking_synapses[_spiking_synapse_idx];
}
```

(continues on next page)
In this case, the “natural” index _idx iterates over all the synapses that received a spike (this is defined in the template) and _postsynaptic_idx refers to the postsynaptic targets for these synapses. The variables w and v are then pulled out of their respective arrays with these indices so that the statement v += w; does the right thing.

### 7.4.5 Getting and setting state variables

When a state variable is accessed (e.g. using G.v), the group does not return a reference to the underlying array itself but instead to a VariableView object. This is because a state variable can be accessed in different contexts and indexing it with a number/array (e.g. obj.v[0]) or a string (e.g. obj.v['i>3']) can refer to different values in the underlying array depending on whether the object is the NeuronGroup, a Subgroup or a Synapses object.

The __setitem__ and __getitem__ methods in VariableView delegate to VariableView.set_item and VariableView.get_item respectively (which can also be called directly under special circumstances). They analyze the arguments (is the index a number, a slice or a string? Is the target value an array or a string expression?) and delegate the actual retrieval/setting of the values to a specific method:

- Getting with a numerical (or slice) index (e.g. G.v[0]): VariableView.get_with_index_array
- Getting with a string index (e.g. G.v['i>3']): VariableView.get_with_expression
- Setting with a numerical (or slice) index and a numerical target value (e.g. G.v[5:] = -70*mV): VariableView.set_with_index_array
- Setting with a numerical (or slice) index and a string expression value (e.g. G.v[5:] = (-70+i)*mV): VariableView.set_with_expression
- Setting with a string index and a string expression value (e.g. G.v['i>5'] = (-70+i)*mV): VariableView.set_with_expression_conditional

These methods are annotated with the device_override decorator and can therefore be implemented in a different way in certain devices. The standalone device, for example, overrides all the getting functions and the setting with index arrays. Note that for standalone devices, the “setter” methods do not actually set the values but only note them down for later code generation.

### 7.4.6 Additional variables and indices

The variables stored in the variables attribute of a VariableOwner can be used everywhere (e.g. in the state updater, in the threshold, the reset, etc.). Objects that depend on these variables, e.g. the Thresholder of a NeuronGroup add additional variables, in particular AuxiliaryVariables that are automatically added to the abstract code: a threshold condition v > 1 is converted into the statement _cond = v > 1; to specify the meaning of the variable _cond for the code generation stage (in particular, C++ code generation needs to know the data type) an AuxiliaryVariable object is created.

In some rare cases, a specific variable_indices dictionary is provided that overrides the indices for variables stored in the variables attribute. This is necessary for synapse creation because the meaning of the variables changes in this context: an expression v>0 does not refer to the v variable of all the connected postsynaptic variables, as it does under other circumstances in the context of a Synapses object, but to the v variable of all possible targets.
7.5 Preferences system

Each preference looks like `codegen.c.compiler`, i.e. dotted names. Each preference has to be registered and validated. The idea is that registering all preferences ensures that misspellings of a preference value by a user causes an error, e.g. if they wrote `codegen.c.compiler` it would raise an error. Validation means that the value is checked for validity, so `codegen.c.compiler = 'gcc'` would be allowed, but `codegen.c.compiler = 'hcc'` would cause an error.

An additional requirement is that the preferences system allows for extension modules to define their own preferences, including extending the existing core brian preferences. For example, an extension might want to define `extension.*` but it might also want to define a new language for codegen, e.g. `codegen.lisp.*`. However, extensions cannot add preferences to an existing category.

7.5.1 Accessing and setting preferences

Preferences can be accessed and set either keyword-based or attribute-based. To set/get the value for the preference example mentioned before, the following are equivalent:

```python
prefs['codegen.c.compiler'] = 'gcc'
prefs.codegen.c.compiler = 'gcc'
if prefs['codegen.c.compiler'] == 'gcc':
    ...
if prefs.codegen.c.compiler == 'gcc':
    ...
```

Using the attribute-based form can be particularly useful for interactive work, e.g. in ipython, as it offers autocompletion and documentation. In ipython, `prefs.codegen.c?` would display a docstring with all the preferences available in the `codegen.c` category.

7.5.2 Preference files

Preferences are stored in a hierarchy of files, with the following order (each step overrides the values in the previous step but no error is raised if one is missing):

- The global defaults are stored in the installation directory.
- The user default are stored in `~/.brian/preferences` (which works on Windows as well as Linux).
- The file `brian_preferences` in the current directory.

7.5.3 Registration

Registration of preferences is performed by a call to `BrianGlobalPreferences.register_preferences`, e.g.:

```python
register_preferences(
    'codegen.c',
    'Code generation preferences for the C language',
    'compiler'= BrianPreference(
        validator=is_compiler,
        docs='...',
        default='gcc'),
(continues on next page)```
The first argument `codegen.c` is the base name, and every preference of the form `codegen.c.*` has to be registered by this function (preferences in subcategories such as `codegen.c.somethingelse.*` have to be specified separately). In other words, by calling `register_preferences`, a module takes ownership of all the preferences with one particular base name. The second argument is a descriptive text explaining what this category is about. The preferences themselves are provided as keyword arguments, each set to a `BrianPreference` object.

### 7.5.4 Validation functions

A validation function takes a value for the preference and returns `True` (if the value is a valid value) or `False`. If no validation function is specified, a default validator is used that compares the value against the default value: Both should belong to the same class (e.g. int or str) and, in the case of a `Quantity` have the same unit.

### 7.5.5 Validation

Setting the value of a preference with a registered base name instantly triggers validation. Trying to set an unregistered preference using keyword or attribute access raises an error. The only exception from this rule is when the preferences are read from configuration files (see below). Since this happens before the user has the chance to import extensions that potentially define new preferences, this uses a special function (`_set_preference`). In this case, for base names that are not yet registered, validation occurs when the base name is registered. If, at the time that `Network.run` is called, there are unregistered preferences set, a `PreferenceError` is raised.

### 7.5.6 File format

The preference files are of the following form:

```
# Preference file

# Comment line
[a]
b.d = 2
[b.e]  
```

This would set preferences `a.b.c=1`, `a.b.d=2` and `a.b.e=3`.

### 7.5.7 Built-in preferences

Brian itself defines the following preferences:

**GSL**

Directory containing GSL code

`GSL.directory = None` Set path to directory containing GSL header files (gsl_odeiv2.h etc.) If this directory is already in Python’s include (e.g. because of conda installation), this path can be set to None.
**codegen**

Code generation preferences

```python
codegen.loop_invariant_optimisations = True
```

Whether to pull out scalar expressions out of the statements, so that they are only evaluated once instead of once for every neuron/synapse/... Can be switched off, e.g. because it complicates the code (and the same optimisation is already performed by the compiler) or because the code generation target does not deal well with it. Defaults to True.

```python
codegen.max_cache_dir_size = 1000
```

The size of a directory (in MB) with cached code for weave or Cython that triggers a warning. Set to 0 to never get a warning.

```python
codegen.string_expression_target = 'numpy'
```

Default target for the evaluation of string expressions (e.g. when indexing state variables). Should normally not be changed from the default numpy target, because the overhead of compiling code is not worth the speed gain for simple expressions.

Accepts the same arguments as `codegen.target`, except for 'auto'.

```python
codegen.target = 'auto'
```

Default target for code generation.

Can be a string, in which case it should be one of:

- 'auto' the default, automatically chose the best code generation target available.
- 'weave' uses scipy.weave to generate and compile C++ code, should work anywhere where gcc is installed and available at the command line.
- 'cython', uses the Cython package to generate C++ code. Needs a working installation of Cython and a C++ compiler.
- 'numpy' works on all platforms and doesn’t need a C compiler but is often less efficient.

Or it can be a `CodeObject` class.

**codegen.cpp**

C++ compilation preferences

```python
codegen.cpp.compiler = ''
```

Compiler to use (uses default if empty)

Should be gcc or msvc.

```python
codegen.cpp.define_macros = []
```

List of macros to define; each macro is defined using a 2-tuple, where ‘value’ is either the string to define it to or None to define it without a particular value (equivalent of “#define FOO” in source or -DFOO on Unix C compiler command line).

```python
codegen.cpp.extra_compile_args = None
```

Extra arguments to pass to compiler (if None, use either `extra_compile_args_gcc` or `extra_compile_args_msvc`).

```python
codegen.cpp.extra_compile_args_gcc = ['-w', '-O3', '-ffast-math', '-fno-finite-math-only', '-march=native']
```

Extra compile arguments to pass to GCC compiler
codegen.cpp.extra_compile_args_msvc = ['./Ox', '/w', '','/MP']

Extra compile arguments to pass to MSVC compiler (the default /arch: flag is determined based on the processor architecture)

codegen.cpp.extra_link_args = []

Any extra platform- and compiler-specific information to use when linking object files together.

codegen.cpp.headers = []

A list of strings specifying header files to use when compiling the code. The list might look like ["<vector>", "my_header"]. Note that the header strings need to be in a form than can be pasted at the end of a #include statement in the C++ code.

codegen.cpp.include_dirs = []

Include directories to use. Note that $prefix/include will be appended to the end automatically, where $prefix is Python's site-specific directory prefix as returned by sys.prefix.

codegen.cpp.libraries = []

List of library names (not filenames or paths) to link against.

codegen.cpp.library_dirs = []

List of directories to search for C/C++ libraries at link time. Note that $prefix/lib will be appended to the end automatically, where $prefix is Python’s site-specific directory prefix as returned by sys.prefix.

codegen.cpp.msvc_architecture = ''

MSVC architecture name (or use system architectue by default).
    Could take values such as x86, amd64, etc.

codegen.cpp.msvc_vars_location = ''

Location of the MSVC command line tool (or search for best by default).

codegen.cpp.runtime_library_dirs = []

List of directories to search for C/C++ libraries at run time.

codegen.generators

Codegen generator preferences (see subcategories for individual languages)

codegen.generators.cpp

C++ codegen preferences

codegen.generators.cpp.flush_denormals = False

Adds code to flush denormals to zero.

The code is gcc and architecture specific, so may not compile on all platforms. The code, for reference is:

```c
#define CSR_FLUSH_TO_ZERO (1 << 15)
unsigned csr = __builtin_ia32_stmxcsr();
csr |= CSR_FLUSH_TO_ZERO;
__builtin_ia32_ldmxcsr(csr);
```


codegen.generators.cpp.restrict_keyword = '__restrict'

7.5. Preferences system
The keyword used for the given compiler to declare pointers as restricted.
This keyword is different on different compilers, the default works for gcc and MSVS.

codegen.runtime
Runtime codegen preferences (see subcategories for individual targets)

codegen.runtime.cython
Cython runtime codegen preferences
codegen.runtime.cython.cache_dir = None
Location of the cache directory for Cython files. By default, will be stored in a
brian_extensions subdirectory where Cython inline stores its temporary files (the result of
get_cython_cache_dir()).
codegen.runtime.cython.delete_source_files = True
Whether to delete source files after compiling. The Cython source files can take a significant amount of
disk space, and are not used anymore when the compiled library file exists. They are therefore deleted by
default, but keeping them around can be useful for debugging.
codegen.runtime.cython.multiprocess_safe = True
Whether to use a lock file to prevent simultaneous write access to cython .pyx and .so files.

codegen.runtime.numpy
Numpy runtime codegen preferences
codegen.runtime.numpy.discard_units = False
Whether to change the namespace of user-specified functions to remove units.

core
Core Brian preferences
core.default_float_dtype = float64
Default dtype for all arrays of scalars (state variables, weights, etc.).
core.default_integer_dtype = int32
Default dtype for all arrays of integer scalars.
core.outdated_dependency_error = True
Whether to raise an error for outdated dependencies (True) or just a warning (False).

core.network
Network preferences
core.network.default_schedule = ['start', 'groups', 'thresholds', 'synapses',
'resets', 'end']
Default schedule used for networks that don’t specify a schedule.
devices

Device preferences

devices.cpp_standalone

C++ standalone preferences

devices.cpp_standalone.extra_make_args_unix = ['-j']

Additional flags to pass to the GNU make command on Linux/OS-X. Defaults to “-j” for parallel compilation.

devices.cpp_standalone.extra_make_args_windows = []

Additional flags to pass to the nmake command on Windows. By default, no additional flags are passed.

devices.cpp_standalone.openmp_spatialneuron_strategy = None

DEPRECATED. Previously used to chose the strategy to parallelize the solution of the three tridiagonal systems for multicompartmental neurons. Now, its value is ignored.

devices.cpp_standalone.openmp_threads = 0

The number of threads to use if OpenMP is turned on. By default, this value is set to 0 and the C++ code is generated without any reference to OpenMP. If greater than 0, then the corresponding number of threads are used to launch the simulation.

devices.cpp_standalone.run_environment_variables = {'LD_BIND_NOW': '1'}

Dictionary of environment variables and their values that will be set during the execution of the standalone code.

legacy

Preferences to enable legacy behaviour

legacy.refractory_timing = False

Whether to use the semantics for checking the refractoriness condition that were in place up until (including) version 2.1.2. In that implementation, refractory periods that were multiples of dt could lead to a varying number of refractory timesteps due to the nature of floating point comparisons). This preference is only provided for exact reproducibility of previously obtained results, new simulations should use the improved mechanism which uses a more robust mechanism to convert refractoriness into timesteps. Defaults to False.

logging

Logging system preferences

logging.console_log_level = 'INFO'

What log level to use for the log written to the console.

Has to be one of CRITICAL, ERROR, WARNING, INFO, DEBUG or DIAGNOSTIC.

logging.delete_log_on_exit = True

Whether to delete the log and script file on exit.

If set to True (the default), log files (and the copy of the main script) will be deleted after the brian process has exited, unless an uncaught exception occured. If set to False, all log files will be kept.

7.5. Preferences system
logging.file_log = True
    Whether to log to a file or not.
    If set to True (the default), logging information will be written to a file. The log level can be set via the
    logging.file_log_level preference.

logging.file_log_level = 'DIAGNOSTIC'
    What log level to use for the log written to the log file.
    In case file logging is activated (see logging.file_log), which log level should be used for logging. Has to be one of CRITICAL, ERROR, WARNING, INFO, DEBUG or DIAGNOSTIC.

logging.save_script = True
    Whether to save a copy of the script that is run.
    If set to True (the default), a copy of the currently run script is saved to a temporary location. It is deleted after a successful run (unless logging.delete_log_on_exit is False) but is kept after an uncaught exception occurred. This can be helpful for debugging, in particular when several simulations are running in parallel.

logging.std_redirection = True
    Whether or not to redirect stdout/stderr to null at certain places.
    This silences a lot of annoying compiler output, but will also hide error messages making it harder to debug problems. You can always temporarily switch it off when debugging. If logging.std_redirection_to_file is set to True as well, then the output is saved to a file and if an error occurs the name of this file will be printed.

logging.std_redirection_to_file = True
    Whether to redirect stdout/stderr to a file.
    If both logging.std_redirection and this preference are set to True, all standard output/error (most importantly output from the compiler) will be stored in files and if an error occurs the name of this file will be printed. If logging.std_redirection is True and this preference is False, then all standard output/error will be completely suppressed, i.e. neither be displayed nor stored in a file.
    The value of this preference is ignore if logging.std_redirection is set to False.

7.6 Adding support for new functions

For a description of Brian’s function system from the user point of view, see Functions.

The default functions available in Brian are stored in the DEFAULT_FUNCTIONS dictionary. New Function objects can be added to this dictionary to make them available to all Brian code, independent of its namespace.

To add a new implementation for a code generation target, a FunctionImplementation can be added to the Function.implementations dictionary. The key for this dictionary has to be either a CodeGenerator class object, or a CodeObject class object. The CodeGenerator of a CodeObject (e.g. CPPCodeGenerator for WeaveCodeObject) is used as a fallback if no implementation specific to the CodeObject class exists.

If a function is already provided for the target language (e.g. it is part of a library imported by default), using the same name, all that is needed is to add an empty FunctionImplementation object to mark the function as implemented. For example, exp is a standard function in C++:

```python
DEFAULT_FUNCTIONS['exp'].implementations[CPPCodeGenerator] = FunctionImplementation()
```
Some functions are implemented but have a different name in the target language. In this case, the `FunctionImplementation` object only has to specify the new name:

```python
DEFAULT_FUNCTIONS['arcsin'].implementations[CPPCodeGenerator] =
    FunctionImplementation('asin')
```

Finally, the function might not exist in the target language at all, in this case the code for the function has to be provided, the exact form of this code is language-specific. In the case of C++, it’s a dictionary of code blocks:

```python
clip_code = {'support_code': '''
    double _clip(const float value, const float a_min, const float a_max)
    {
        if (value < a_min)
            return a_min;
        if (value > a_max)
            return a_max;
        return value;
    }
'''
    }
DEFAULT_FUNCTIONS['clip'].implementations[CPPCodeGenerator] = FunctionImplementation(
    '_clip',
    code=clip_code)
```

### 7.7 Code generation

The generation of a code snippet is done by a `CodeGenerator` class. The templates are stored in the `CodeObject.templater` attribute, which is typically implemented as a subdirectory of templates. The compilation and running of code is done by a `CodeObject`. See the sections below for each of these.

#### 7.7.1 Code path

The following gives an outline of the key steps that happen for the code generation associated to a `NeuronGroup` `StateUpdater`. The items in grey are Brian core functions and methods and do not need to be implemented to create a new code generation target or device. The parts in yellow are used when creating a new device. The parts in green relate to generating code snippets from abstract code blocks. The parts in blue relate to creating new templates which these snippets are inserted into. The parts in red relate to creating new runtime behaviour (compiling and running generated code).
In brief, what happens can be summarised as follows. `Network.run` will call `BrianObject.before_run` on each of the objects in the network. Objects such as `StateUpdater`, which is a subclass of `CodeRunner` use this spot to generate and compile their code. The process for doing this is to first create the abstract code block, done in the `StateUpdater.update_abstract_code` method. Then, a `CodeObject` is created with this code block. In doing so, Brian will call out to the currently active `Device` to get the `CodeObject` and `CodeGenerator` classes associated to the device, and this hierarchy of calls gives several hooks which can be changed to implement new targets.

### 7.7.2 Code generation

To implement a new language, or variant of an existing language, derive a class from `CodeGenerator`. Good examples to look at are the `NumpyCodeGenerator`, `CPPCodeGenerator` and `CythonCodeGenerator` classes in the `brian2.codegen.generators` package. Each `CodeGenerator` has a `class_name` attribute which is a string used by the user to refer to this code generator (for example, when defining function implementations).

The derived `CodeGenerator` class should implement the methods marked as `NotImplemented` in the base `CodeGenerator` class. `CodeGenerator` also has several handy utility methods to make it easier to write these, see the existing examples to get an idea of how these work.

### 7.7.3 Syntax translation

One aspect of writing a new language is that sometimes you need to translate from Python syntax into the syntax of another language. You are free to do this however you like, but we recommend using a `NodeRenderer` class which allows you to iterate over the abstract syntax tree of an expression. See examples in `brian2.parsing.rendering`.

### 7.7.4 Templates

In addition to snippet generation, you need to create templates for the new language. See the `templates` directories in `brian2.codegen.runtime.*` for examples of these. They are written in the Jinja2 templating system. The
location of these templates is set as the `CodeObject.templater` attribute. Examples such as CPPCodeObject show how this is done.

### 7.7.5 Code objects

To allow the final code block to be compiled and run, derive a class from `CodeObject`. This class should implement the placeholder methods defined in the base class. The class should also have attributes `templater` (which should be a `Templater` object pointing to the directory where the templates are stored), `generator_class` (which should be the `CodeGenerator` class), and `class_name` (which should be a string the user can use to refer to this code generation target).

### 7.7.6 Default functions

You will typically want to implement the default functions such as the trigonometric, exponential and `rand` functions. We usually put these implementations either in the same module as the `CodeGenerator` class or the `CodeObject` class depending on whether they are language-specific or runtime target specific. See those modules for examples of implementing these functions.

### 7.7.7 Code guide

- `brian2.codegen`: everything related to code generation
- `brian2.codegen.generators`: snippet generation, including the `CodeGenerator` classes and default function implementations.
- `brian2.codegen.runtime`: templates, compilation and running of code, including `CodeObject` and default function implementations.
- `brian2.core.functions, brian2.core.variables`: these define the values that variable names can have.
- `brian2.parsing`: tools for parsing expressions, etc.
- `brian2.parsing.rendering`: AST tools for rendering expressions in Python into different languages.
- `brian2.utils`: various tools for string manipulation, file management, etc.

### 7.7.8 Additional information

For some additional (older, but still accurate) notes on code generation:

#### Older notes on code generation

The following is an outline of how the Brian 2 code generation system works, with indicators as to which packages to look at and which bits of code to read for a clearer understanding.

We illustrate the global process with an example, the creation and running of a single `NeuronGroup` object:

- Parse the equations, add refractoriness to them: this isn’t really part of code generation.
- Allocate memory for the state variables.
- Create `Threshold, Resetter, and StateUpdater` objects.
  - Determine all the variable and function names used in the respective abstract code blocks and templates
– Determine the abstract namespace, i.e. determine a Variable or Function object for each name.

– Create a CodeObject based on the abstract code, template and abstract namespace. This will generate code in the target language and the namespace in which the code will be executed.

- At runtime, each object calls CodeObject.__call__ to execute the code.

Stages of code generation

Equations to abstract code

In the case of Equations, the set of equations are combined with a numerical integration method to generate an abstract code block (see below) which represents the integration code for a single time step.

An example of this would be converting the following equations:

```python
eqs = '''
dv/dt = (v0-v)/tau : volt (unless refractory)
v0 : volt
'''

group = NeuronGroup(N, eqs, threshold='v>10*mV',
                        reset='v=0*mV', refractory=5*ms)
```

into the following abstract code using the exponential_euler method (which is selected automatically):

```python
not_refractory = 1*{(t - lastspike) > 0.005000}
_BA_v = -v0
_v = -_BA_v + (_BA_v + v)*exp(-dt*not_refractory/tau)
v = _v
```

The code for this stage can be seen in NeuronGroup.__init__, StateUpdater.__init__, and StateUpdater.update_abstract_code in the brian2.groups.neurongroup, and the StateUpdateMethod classes defined in the brian2.stateupdaters package.

For more details, see State update.

Abstract code

‘Abstract code’ is just a multi-line string representing a block of code which should be executed for each item (e.g. each neuron, each synapse). Each item is independent of the others in abstract code. This allows us to later generate code either for vectorised languages (like numpy in Python) or using loops (e.g. in C++).

Abstract code is parsed according to Python syntax, with certain language constructs excluded. For example, there cannot be any conditional or looping statements at the moment, although support for this is in principle possible and may be added later. Essentially, all that is allowed at the moment is a sequence of arithmetical a = b*c style statements.

Abstract code is provided directly by the user for threshold and reset statements in NeuronGroup and for pre/post spiking events in Synapses.

Abstract code to snippet

We convert abstract code into a ‘snippet’, which is a small segment of code which is syntactically correct in the target language, although it may not be runnable on its own (that’s handled by insertion into a ‘template’ later). This is handled by the CodeGenerator object in brian2.codegen.generators. In the case of converting into
python/numpy code this typically doesn’t involve any changes to the code at all because the original code is in Python syntax. For conversion to C++, we have to do some syntactic transformations (e.g. \(a^b\) is converted to `pow(a, b)`), and add declarations for certain variables (e.g. converting \(x = y \times z\) into `const double x = y * z;`).

An example of a snippet in C++ for the equations above:

```c++
const double v0 = _ptr_array_neurongroup_v0[_neuron_idx];
const double lastspike = _ptr_array_neurongroup_lastspike[_neuron_idx];
bool not_refractory = _ptr_array_neurongroup_not_refractory[_neuron_idx];
double v = _ptr_array_neurongroup_v[_neuron_idx];
not_refractory = 1 * (t - lastspike > 0.0050000000000000001);
const double _BA_v = -(v0);
const double _v = -(_BA_v) + (_BA_v + v) * exp(-(dt) * not_refractory / tau);
v = _v;
_ptr_array_neurongroup_not_refractory[_neuron_idx] = not_refractory;
_ptr_array_neurongroup_v[_neuron_idx] = v;
```

The code path that includes snippet generation will be discussed in more detail below, since it involves the concepts of namespaces and variables which we haven’t covered yet.

### Snippet to code block

The final stage in the generation of a runnable code block is the insertion of a snippet into a template. These use the Jinja2 template specification language. This is handled in `brian2.codegen.templates`.

An example of a template for Python thresholding:

```jinja
# USES_VARIABLES { not_refractory, lastspike, t }
{% for line in code_lines %}
{{line}}
{% endfor %}
_return_values, = _cond.nonzero()
# Set the neuron to refractory
not_refractory[_return_values] = False
lastspike[_return_values] = t
```

and the output code from the example equations above:

```python
# USES_VARIABLES { not_refractory, lastspike, t }
v = _array_neurongroup_v
_cond = v > 10 * mV
_return_values, = _cond.nonzero()
# Set the neuron to refractory
not_refractory[_return_values] = False
lastspike[_return_values] = t
```

### Code block to executing code

A code block represents runnable code. Brian operates in two different regimes, either in runtime or standalone mode. In runtime mode, memory allocation and overall simulation control is handled by Python and numpy, and code objects operate on this memory when called directly by Brian. This is the typical way that Brian is used, and it allows for a rapid development cycle. However, we also support a standalone mode in which an entire project workspace is generated for a target language or device by Brian, which can then be compiled and run independently of Brian. Each mode has different templates, and does different things with the outputted code blocks. For runtime mode, in Python/numpy code is executed by simply calling the `exec` statement on the code block in a given namespace. For
C++/weave code, the `scipy.weave.inline` function is used. In standalone mode, the templates will typically each be saved into different files.

**Key concepts**

**Namespaces**

In general, a namespace is simply a mapping/dict from names to values. In Brian we use the term ‘namespace’ in two ways: the high level “abstract namespace” maps names to objects based on the `Variables` or `Function` class. In the above example, `v` maps to an `ArrayVariable` object, `tau` to a `Constant` object, etc. This namespace has all the information that is needed for checking the consistency of units, to determine which variables are boolean or scalar, etc. During the `CodeObject` creation, this abstract namespace is converted into the final namespace in which the code will be executed. In this namespace, `v` maps to the numpy array storing the state variable values (without units) and `tau` maps to a concrete value (again, without units). See *Equations and namespaces* for more details.

**Variable**

`Variable` objects contain information about the variable they correspond to, including details like the data type, whether it is a single value or an array, etc.

See `brian2.core.variables` and, e.g. `Group._create_variables`, `NeuronGroup._create_variables`.

**Templates**

Templates are stored in Jinja2 format. They come in one of two forms, either they are a single template if code generation only needs to output a single block of code, or they define multiple Jinja macros, each of which is a separate code block. The `CodeObject` should define what type of template it wants, and the names of the macros to define. For examples, see the templates in the directories in `brian2/codegen/runtime`. See `brian2.codegen.templates` for more details.

**Code guide**

This section includes a guide to the various relevant packages and subpackages involved in the code generation process.

`codegen` Stores the majority of all code generation related code.

- `codegen.functions` Code related to including functions - built-in and user-defined - in generated code.
- `codegen.generators` Each `CodeGenerator` is defined in a module here.
- `codegen.runtime` Each runtime `CodeObject` and its templates are defined in a package here.

`core`

- `core.variables` The `Variable` types are defined here.

`equations` Everything related to *Equations*.

`groups` All `Group` related stuff is in here. The `Group.resolve` methods are responsible for determining the abstract namespace.

`parsing` Various tools using Python’s `ast` module to parse user-specified code. Includes syntax translation to various languages in `parsing.rendering`. 
stateupdaters Everything related to generating abstract code blocks from integration methods is here.

7.8 Devices

This document describes how to implement a new Device for Brian. This is a somewhat complicated process, and you should first be familiar with devices from the user point of view (Computational methods and efficiency) as well as the code generation system (Code generation).

We wrote Brian’s devices system to allow for two major use cases, although it can potentially be extended beyond this. The two use cases are:

1. Runtime mode. In this mode, everything is managed by Python, including memory management (using numpy by default) and running the simulation. Actual computational work can be carried out in several different ways, including numpy, weave or Cython.
2. Standalone mode. In this mode, running a Brian script leads to generating an entire source code project tree which can be compiled and run independently of Brian or Python.

Runtime mode is handled by RuntimeDevice and is already implemented, so here I will mainly discuss standalone devices. A good way to understand these devices is to look at the implementation of CPPStandaloneDevice (the only one implemented in the core of Brian). In many cases, the simplest way to implement a new standalone device would be to derive a class from CPPStandaloneDevice and overwrite just a few methods.

7.8.1 Memory management

Memory is managed primarily via the Device.add_array, Device.get_value and Device.set_value methods. When a new array is created, the add_array method is called, and when trying to access this memory the other two are called. The RuntimeDevice uses numpy to manage the memory and returns the underlying arrays in these methods. The CPPStandaloneDevice just stores a dictionary of array names but doesn’t allocate any memory. This information is later used to generate code that will allocate the memory, etc.

7.8.2 Code objects

As in the case of runtime code generation, computational work is done by a collection of CodeObject s. In CPPStandaloneDevice, each code object is converted into a pair of .cpp and .h files, and this is probably a fairly typical way to do it. For this device, it just uses the same code generation routines as for the runtime C++ device weave.

7.8.3 Building

The method Device.build is used to generate the project. This can be implemented any way you like, although looking at CPPStandaloneDevice.build is probably a good way to get an idea of how to do it.

7.8.4 Device override methods

Several functions and methods in Brian are decorated with the device_override decorator. This mechanism allows a standalone device to override the behaviour of any of these functions by implementing a method with the name provided to device_override. For example, the CPPStandaloneDevice uses this to override Network.run as CPPStandaloneDevice.network_run.
7.8.5 Other methods

There are some other methods to implement, including initialising arrays, creating spike queues for synaptic propagation. Take a look at the source code for these.

7.9 Multi-threading with OpenMP

The following is an outline of how to make C++ standalone templates compatible with OpenMP, and therefore make them work in a multi-threaded environment. This should be considered as an extension to Code generation, that has to be read first. The C++ standalone mode of Brian is compatible with OpenMP, and therefore simulations can be launched by users with one or with multiple threads. Therefore, when adding new templates, the developers need to make sure that those templates are properly handling the situation if launched with OpenMP.

7.9.1 Key concepts

All the simulations performed with the C++ standalone mode can be launched with multi-threading, and make use of multiple cores on the same machine. Basically, all the Brian operations that can easily be performed in parallel, such as computing the equations for NeuronGroup, Synapses, and so on can and should be split among several threads. The network construction, so far, is still performed only by one single thread, and all created objects are shared by all the threads.

7.9.2 Use of #pragma flags

In OpenMP, all the parallelism is handled thanks to extra comments, added in the main C++ code, under the form:

```
#pragma omp ...
```

But to avoid any dependencies in the code that is generated by Brian when OpenMP is not activated, we are using functions that will only add those comments, during code generation, when such a multi-threading mode is turned on. By default, nothing will be inserted.

Translational of the #pragma commands

All the translations from openmp_pragma() calls in the C++ templates are handled in the file devices/cpp_standalone/codeobject.py In this function, you can see that all calls with various string inputs will generate #pragma statements inserted into the C++ templates during code generation. For example:

```
{{ openmp_pragma('static') }}
```

will be transformed, during code generation, into:

```
#pragma omp for schedule(static)
```

You can find the list of all the translations in the core of the openmp_pragma() function, and if some extra translations are needed, they should be added here.

Execution of the OpenMP code

In this section, we are explaining the main ideas behind the OpenMP mode of Brian, and how the simulation is executed in such a parallel context. As can be seen in devices/cpp_standalone/templates/main.cpp,
the appropriate number of threads, defined by the user, is fixed at the beginning of the main function in the C++ code with:

```cpp
{{ openmp_pragma('set_num_threads') }}
```

equivalent to (thanks to the openmp_pragam() function defined above): nothing if OpenMP is turned off (default), and to:

```cpp
omp_set_dynamic(0);
omp_set_num_threads(nb_threads);
```

otherwise. When OpenMP creates a parallel context, this is the number of threads that will be used. As said, network creation is performed without any calls to OpenMP, on one single thread. Each template that wants to use parallelism has to add `{{ openmp_pragma{('parallel')}}` to create a general block that will be executed in parallel or `{{ openmp_pragma{('parallel-static')}}` to execute a single loop in parallel.

### 7.9.3 How to make your template use OpenMP parallelism

To design a parallel template, such as for example `devices/cpp_standalone/templates/common_group.cpp`, you can see that as soon as you have loops that can safely be split across nodes, you just need to add an openmp command in front of those loops:

```cpp
{{openmp_pragma('parallel-static')}}
for(int _idx=0; _idx<N; _idx++)
{
    ...
}
```

By doing so, OpenMP will take care of splitting the indices and each thread will loop only on a subset of indices, sharing the load. By default, the scheduling use for splitting the indices is static, meaning that each node will get the same number of indices: this is the faster scheduling in OpenMP, and it makes sense for `NeuronGroup` or `Synapses` because operations are the same for all indices. By having a look at examples of templates such as `devices/cpp_standalone/templates/statemonitor.cpp`, you can see that you can merge portions of code executed by only one node and portions executed in parallel. In this template, for example, only one node is recording the time and extending the size of the arrays to store the recorded values:

```cpp
{{_dynamic_t}}.push_back(_clock_t);
// Resize the dynamic arrays
{{_recorded}}.resize(_new_size, _num_indices);
```

But then, values are written in the arrays by all the nodes:

```cpp
{{ openmp_pragma('parallel-static') }}
for (int _i = 0; _i < _num_indices; _i++)
{
    ....
}
```

In general, operations that manipulate global data structures, e.g. that use `push_back` for a `std::vector`, should only be executed by a single thread.

### 7.9.4 Synaptic propagation in parallel

7.9. Multi-threading with OpenMP
General ideas

With OpenMP, synaptic propagation is also multi-threaded. Therefore, we have to modify the `SynapticPathway` objects, handling spike propagation. As can be seen in `devices/cpp_standalone/templates/synapses_classes.cpp`, such an object, created during run time, will be able to get the number of threads decided by the user:

```
_nb_threads = {{ openmp_pragma('get_num_threads') }};
```

By doing so, a `SynapticPathway`, instead of handling only one `SpikeQueue`, will be divided into `_nb_threads` `SpikeQueue`s, each of them handling a subset of the total number of connections. All the calls to `SynapticPathway` object are performed from within parallel blocks in the `synapses` and `synapses_push_spikes` template, we have to take this parallel context into account. This is why all the function of the `SynapticPathway` object are taking care of the node number:

```
void push(int *spikes, unsigned int nspikes)
{
    queue[{{ openmp_pragma('get_thread_num') }}]->push(spikes, nspikes);
}
```

Such a method for the `SynapticPathway` will make sure that when spikes are propagated, all the threads will propagate them to their connections. By default, again, if OpenMP is turned off, the queue vector has size 1.

Preparation of the `SynapticPathway`

Here we are explaining the implementation of the `prepare()` method for `SynapticPathway`:

```
{{ openmp_pragma('parallel') }}
{
    unsigned int length;
    if ({{ openmp_pragma('get_thread_num') }} == _nb_threads - 1)
        length = n_synapses - (unsigned int) {{ openmp_pragma('get_thread_num') }}*n_synapses/_nb_threads;
    else
        length = (unsigned int) n_synapses/_nb_threads;

    unsigned int padding = {{ openmp_pragma('get_thread_num') }}*(n_synapses/_nb_threads);

    queue[{{ openmp_pragma('get_thread_num') }}]->openmp_padding = padding;
    queue[{{ openmp_pragma('get_thread_num') }}]->prepare(&real_delays[padding], &sources[padding], length, _dt);
}
```

Basically, each threads is getting an equal number of synapses (except the last one, that will get the remaining ones, if the number is not a multiple of `n_threads`), and the queues are receiving a padding integer telling them what part of the synapses belongs to each queue. After that, the parallel context is destroyed, and network creation can continue. Note that this could have been done without a parallel context, in a sequential manner, but this is just speeding up everything.

Selection of the spikes

Here we are explaining the implementation of the `peek()` method for `SynapticPathway`. This is an example of concurrent access to data structures that are not well handled in parallel, such as `std::vector`. When `peek()` is called, we need to return a vector of all the neuron spiking at that particular time. Therefore, we need to ask every
queue of the SynapticPathway what are the id of the spiking neurons, and concatenate them. Because those ids are stored in vectors with various shapes, we need to loop over nodes to perform this concatenate, in a sequential manner:

```c
{{ openmp_pragma('static-ordered') }}
for(int _thread=0; _thread < {{ openmp_pragma('get_num_threads') }}; _thread++)
{
    {{ openmp_pragma('ordered') }}
    {
        if (_thread == 0)
            all_peek.clear();
        all_peek.insert(all_peek.end(), queue[_thread]->peek()->begin(), queue[_thread]->peek()->end());
    }
}
```

The loop, with the keyword ‘static-ordered’, is therefore performed such that node 0 enters it first, then node 1, and so on. Only one node at a time is executing the block statement. This is needed because vector manipulations can not be performed in a multi-threaded manner. At the end of the loop, all_peek is now a vector where all sub queues have written the id of spiking cells, and therefore this is the list of all spiking cells within the SynapticPathway.

### 7.9.5 Compilation of the code

One extra file needs to be modified, in order for OpenMP implementation to work. This is the makefile devices/cpp_standalone/templates/makefile. As one can simply see, the CFLAGS are dynamically modified during code generation thanks to:

```c
{{ openmp_pragma('compilation') }}
```

If OpenMP is activated, this will add the following dependencies:

```c
-fopenmp
```

such that if OpenMP is turned off, nothing, in the generated code, does depend on it.

### 7.10 Solving differential equations with the GNU Scientific Library

Conventionally, Brian generates its own code performing Numerical integration according to the chosen algorithm (see the section on Code generation). Another option is to let the differential equation solvers defined in the GNU Scientific Library (GSL) solve the given equations. In addition to offering a few extra integration methods, the GSL integrator comes with the option of having an adaptable timestep. The latter functionality can have benefits for the speed with which large simulations can be run. This is because it allows the use of larger timesteps for the overhead loops in Python, without losing the accuracy of the numerical integration at points where small timesteps are necessary. In addition, a major benefit of using the ODE solvers from GSL is that an estimation is performed on how wrong the current solution is, so that simulations can be performed with some confidence on accuracy. (Note however that the confidence of accuracy is based on estimation!)

#### 7.10.1 StateUpdateMethod
Translation of equations to abstract code

The first part of Brian’s code generation is the translation of equations to what we call ‘abstract code’. In the case of Brian’s stateupdaters so far, this abstract code describes the calculations that need to be done to update differential variables depending on their equations as is explained in the section on State update. In the case of preparing the equations for GSL integration this is a bit different. Instead of writing down the computations that have to be done to reach the new value of the variable after a time step, the equations have to be described in a way that GSL understands. The differential equations have to be defined in a function and the function is given to GSL. This is best explained with an example. If we have the following equations (taken from the adaptive threshold example):

\[
\begin{align*}
\frac{dv}{dt} &= -v/(10\times\text{ms}) : \text{volt} \\
\frac{dvt}{dt} &= (10\times\text{mV} - vt)/(15\times\text{ms}) : \text{volt}
\end{align*}
\]

We would describe the equations to GSL as follows:

\[
\begin{align*}
v &= y[0] \\
vt &= y[1] \\
f[0] &= -v/(10\times10^{-3}) \\
f[1] &= (10\times10^{-3} - vt)
\end{align*}
\]

Each differential variable gets an index. Its value at any time is saved in the \(y\)-array and the derivatives are saved in the \(f\)-array. However, doing this translation in the stateupdater would mean that Brian has to deal with variable descriptions that contain array accessing: something that for example sympy doesn’t do. Because we still want to use Brian’s existing parsing and checking mechanisms, we needed to find a way to describe the abstract code with only ‘normal’ variable names. Our solution is to replace the \(y[0], f[0]\), etc. with a ‘normal’ variable name that is later replaced just before the final code generation (in the GSLCodeGenerator). It has a tag and all the information needed to write the final code. As an example, the GSL abstract code for the above equations would be:

\[
\begin{align*}
v &= \_\text{gsl\_y0} \\
v &= \_\text{gsl\_y1} \\
\_\text{gsl\_f0} &= -v/(10\times10^{-3}) \\
\_\text{gsl\_f1} &= (10\times10^{-3} - vt)
\end{align*}
\]

In the GSLCodeGenerator these tags get replaced by the actual accessing of the arrays.

Return value of the StateUpdateMethod

So far, for each each code generation language (numpy, weave, cython) there was just one set of rules of how to translate abstract code to real code, described in its respective CodeObject and CodeGenerator. If the target language is set to weave, the stateupdater will use the WeaveCodeObject, just like other objects such as the StateMonitor. However, to achieve the above described translations of the abstract code generated by the StateUpdateMethod, we need a special WeaveCodeObject for the stateupdater alone (which at its turn can contain the special CodeGenerator), and this CodeObject should be selected based on the chosen StateUpdateMethod.

In order to achieve CodeObject selection based on the chosen stateupdater, the StateUpdateMethod returns a class that can be called with an object, and the appropriate CodeObject is added as an attribute to the given object. The return value of this callable is the abstract code describing the equations in a language that makes sense to the GSLCodeGenerator.

7.10.2 GSLCodeObject

Each target language has its own GSLCodeObject that is derived from the already existing code object of its language. There are only minimal changes to the already existing code object:
• Overwrite stateupdate template: a new version of the stateupdate template is given (stateupdate.cpp for weave/C++ standalone and stateupdate.pyx for cython).

• Have a GSL specific generator_class: GSLWeaveCodeGenerator or GSLCythonCodeGenerator

• Add the attribute original_generator_class: the conventional target-language generator is used to do the bulk of the translation to get from abstract code to language-specific code.

This defining of GSL-specific code objects also allowed us to catch compilation errors so we can give the user some information on that it might be GSL-related (overwriting the compile() method in the case of cython and the run() method for weave). In the case of the C++ CodeObject such overriding wasn’t really possible so compilation errors in this case might be quite undescrptive.

### 7.10.3 GSLCodeGenerator

This is where the magic happens. Roughly 1000 lines of code define the translation of abstract code to code that uses the GNU Scientific Library’s ODE solvers to achieve state updates.

Upon a call to run(), the code objects necessary for the simulation get made. The code for this is described in the device. Part of making the code objects is generating the code that describes the code objects. This starts with a call to translate, which in the case of GSL brings us to the GSLCodeGenerator.translate(). This method is built up as follows:

• Some GSL-specific preparatory work:
  - Check whether the equations contain variable names that are reserved for the GSL code.
  - Add the ‘gsl tags’ (see section on StateUpdateMethod) to the variables known to Brian as non-scalars. This is necessary to ensure that all equations containing ‘gsl tags’ are considered vector equations, and thus added to Brian’s vector code.
  - Add GSL integrator meta variables as official Brian variables, so these are also taken into account upon translation. The meta variables that are possible are described in the user manual (e.g. GSL’s step taken in a single overhead step ‘_step_count’).
  - Save function names. The original generators delete the function names from the variables dictionary once they are processed. However, we need to know later in the GSL part of the code generation whether a certain encountered variable name refers to a function or not.

• Brian’s general preparatory work. This piece of code is directly copied from the base CodeGenerator and is thus similar to what is done normally.

• A call to original_generator.translate() to get the abstract code translated into code that is target-language specific.

• A lot of statements to translate the target-language specific code to GSL-target-language specific code, described in more detail below.

The biggest difference between conventional Brian code and GSL code is that the stateupdate-describing lines are contained directly in the main() or in a separate function, respectively. In both cases, the equations describing the system refer to parameters that are in the Brian namespace (e.g. “dv/dt = -v/tau” needs access to “tau”). How can we access Brian’s namespace in this separate function that is needed with GSL?

To explain the solution we first need some background information on this ‘separate function’ that is given to the GSL integrators: _GSL_func. This function always gets three arguments:

• double t: the current time. This is relevant when the equations are dependent on time.

• const double _GSL_y[]: an array containing the current values of the differential variables (const because the cannot be changed by _GSL_func itself).
• **double f[]:** an array containing the derivatives of the differential variables (i.e. the equations describing the differential system).

• **void * params:** a pointer.

The pointer can be a pointer to whatever you want, and can thus point to a data structure containing the system parameters (such as tau). To achieve a structure containing all the parameters of the system, a considerable amount of code has to be added/changed to that generated by conventional Brian:

- The data structure, _GSL_dataholder, has to be defined with all variables needed in the vector code. For this reason, also the datatype of each variable is required.
  - This is done in the method GSLCodeGenerator.write_dataholder

- Instead of referring to the variables by their name only (e.g. \( \frac{dv}{dt} = -\frac{v}{\tau} \)), the variables have to be accessed as part of the data structure (e.g. \( \frac{dv}{dt} = -v/_GSL_dataholder->\tau \) in the case of weave/cpp). Also, as mentioned earlier, we want to translate the 'gsl tags' to what they should be in the final code (e.g. _gsl_f0 to f[0]).
  - This is done in the method GSLCodeGenerator.translate_vector_code. It works based on the to_replace dictionary (generated in the methods GSLCodeGenerator.diff_var_to_replace and GSLCodeGenerator.to_replace_vector_vars) that simply contains the old variables as keys and new variables as values, and is given to the word_replace function.

- The values of the variables in the data structure have to be set to the values of the variables in the Brian namespace.
  - This is done in the method GSLCodeGenerator.unpack_namespace, and for the ‘scalar’ variables that require calculation first it is done in the method GSLCodeGenerator.translate_scalar_code.

In addition, a few more ‘support’ functions are generated for the GSL script:

- **int _set_dimension(size_t * dimension):** sets the dimension of the system. Required for GSL.

- **double* _assign_memory_y():** allocates the right amount of memory for the y array (also according to the dimension of the system).

- **int _fill_y_vector(_dataholder* _GSL_dataholder, double* _GSL_y, int _idx):** pulls out the values for each differential variable out of the ‘Brian’ array into the y-vector. This happens in the vector loop (e.g. \( y[0] = _GSL_dataholder->_ptr_array_neurongroup_v[_idx]; \) for weave/C++).

- **int _empty_y_vector(_dataholder* _GSL_dataholder, double* _GSL_y, int _idx):** the opposite of _fill_y_vector. Pulls final numerical solutions from the y array and gives it back to Brian’s namespace.

- **double* _set_GSL_scale_array():** sets the array bound for each differential variable, for which the values are based on method_options['absolute_error'] and method_options['absolute_error_per_variable'].

All of this is written in support functions so that the vector code in the main() can stay almost constant for any simulation.

### 7.10.4 Stateupdate templates

There is many extra things that need to be done for each simulation when using GSL compared to conventional Brian stateupdaters. These are summarized in this section.

Things that need to be done for every type of simulation (either before, in or after main()):
- Cython-only: define the structs and functions that we will be using in cython language (for weave these definitions already sit in GSL’s own header files that are included).

- Prepare the `gsl_odeiv2_system`: give function pointer, set dimension, give pointer to `_GSL_dataholder` as params.

- Allocate the driver (name for the struct that contains the info necessary to perform GSL integration)

- Define dt.

Things that need to be done every loop iteration for every type of simulation:

- Define t and t1 (t + dt).

- Transfer the values in the Brian arrays to the y-array that will be given to GSL.

- Set `_GSL_dataholder._idx` (in case we need to access array variables in `_GSL_func`).

- Initialize the driver (reset counters, set `dt_start`).

- Apply driver (either with adaptable- or fixed time step).

- Optionally save certain meta-variables

- Transfer values from GSL’s y-vector to Brian arrays
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