# **Chainer Documentation**

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Preferred Networks, inc. and Preferred Infrastructure, inc.

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This is the Chainer documentation.

## CHAPTER 1

## Install Guide

## **Before installing Chainer**

We recommend these platforms.

- Ubuntu 14.04 LTS 64bit
- CentOS 7 64bit

Chainer is supported on Python 2.7.6+, 3.4.3+, 3.5.1+, 3.6.0+. Chainer uses C++ compiler such as g++. You need to install it before installing Chainer. This is typical installation method for each platform:

```
# Ubuntu 14.04
$ apt-get install g++
# CentOS 7
$ yum install gcc-c++
```

If you use old setuptools, upgrade it:

\$ pip install -U setuptools

## **Install Chainer**

Chainer depends on these Python packages:

- NumPy 1.9, 1.10, 1.11, 1.12
- Six 1.9

CUDA support

- CUDA 6.5, 7.0, 7.5, 8.0
- filelock

cuDNN support

• cuDNN v2, v3, v4, v5, v5.1, v6

Caffe model support

- Protocol Buffers
- protobuf>=3.0.0 is required for Py3

All these libraries are automatically installed with pip or setup.py.

Image dataset is optional

• Pillow

HDF5 serialization is optional

• h5py 2.5.0

#### Install Chainer via pip

We recommend to install Chainer via pip:

```
$ pip install chainer
```

#### **Install Chainer from source**

You can use setup.py to install Chainer from source:

```
$ tar zxf chainer-x.x.x.tar.gz
$ cd chainer-x.x.x
$ python setup.py install
```

#### When an error occurs...

Use -vvvv option with pip command. That shows all logs of installation. It may helps you:

```
$ pip install chainer -vvvv
```

#### Install Chainer with CUDA

You need to install CUDA Toolkit before installing Chainer. If you have CUDA in a default directory or set CUDA\_PATH correctly, Chainer installer finds CUDA automatically:

```
$ pip install chainer
```

**Note:** Chainer installer looks up CUDA\_PATH environment variable first. If it is empty, the installer looks for nvcc command from PATH environment variable and use its parent directory as the root directory of CUDA installation. If nvcc command is also not found, the installer tries to use the default directory for Ubuntu /usr/local/cuda.

If you installed CUDA into a non-default directory, you need to specify the directory with CUDA\_PATH environment variable:

\$ CUDA\_PATH=/opt/nvidia/cuda pip install chainer

**Warning:** If you want to use sudo to install Chainer, note that sudo command initializes all environment variables. Please specify CUDA\_PATH environment variable inside sudo like this:

```
$ sudo CUDA_PATH=/opt/nvidia/cuda pip install chainer
```

#### Install Chainer with CUDA and cuDNN

cuDNN is a library for Deep Neural Networks that NVIDIA provides. Chainer can use cuDNN. If you want to enable cuDNN, install cuDNN and CUDA before installing Chainer. We recommend you to install developer library of deb package of cuDNN.

If you want to install tar-gz version, we recommend you to install it to CUDA directory. For example if you uses Ubuntu Linux, copy .h files to include directory and .so files to lib64 directory:

```
$ cp /path/to/cudnn.h $CUDA_PATH/include
$ cp /path/to/libcudnn.so* $CUDA_PATH/lib64
```

The destination directories depend on your environment.

If you want to use cuDNN installed in other directory, please use CFLAGS, LDFLAGS and LD\_LIBRARY\_PATH environment variables before installing Chainer:

```
export CFLAGS=-I/path/to/cudnn/include
export LDFLAGS=-L/path/to/cudnn/lib
export LD_LIBRARY_PATH=/path/to/cudnn/lib:$LD_LIBRARY_PATH
```

#### Install Chainer for developers

Chainer uses Cython (>=0.24). Developers need to use Cython to regenerate C++ sources from pyx files. We recommend to use pip with -e option for editable mode:

```
$ pip install -U cython
$ cd /path/to/chainer/source
$ pip install -e .
```

Users need not to install Cython as a distribution package of Chainer only contains generated sources.

#### Support image dataset

Install Pillow manually to activate image dataset. This feature is optional:

```
$ pip install pillow
```

#### Support HDF5 serialization

Install h5py manually to activate HDF5 serialization. This feature is optional:

\$ pip install h5py

Before installing h5py, you need to install libhdf5. It depends on your environment:

```
# Ubuntu 14.04
$ apt-get install libhdf5-dev
# CentOS 7
$ yum -y install epel-release
$ yum install hdf5-devel
```

## **Uninstall Chainer**

Use pip to uninstall Chainer:

\$ pip uninstall chainer

Note: When you upgrade Chainer, pip sometimes installed various version of Chainer in site-packages. Please uninstall it repeatedly until pip returns an error.

## **Upgrade Chainer**

Just use pip with –U option:

\$ pip install -U chainer

## **Reinstall Chainer**

If you want to reinstall Chainer, please uninstall Chainer and then install it. We recommend to use --no-cache-dir option as pip sometimes uses cache:

```
$ pip uninstall chainer
$ pip install chainer --no-cache-dir
```

When you install Chainer without CUDA, and after that you want to use CUDA, please reinstall Chainer. You need to reinstall Chainer when you want to upgrade CUDA.

### **Run Chainer with Docker**

We provide the official Docker image. Use nvidia-docker command to run Chainer image with GPU. You can login to the environment with bash, and run the Python interpreter:

\$ nvidia-docker run -it chainer/chainer /bin/bash

Or, run the interpreter directly:

\$ nvidia-docker run -it chainer/chainer /usr/bin/python

## What "recommend" means?

We tests Chainer automatically with Jenkins. All supported environments are tested in this environment. We cannot guarantee that Chainer works on other environments.

## FAQ

#### The installer says "hdf5.h is not found"

You don't have libhdf5. Please install hdf5. See Before installing Chainer.

#### **MemoryError happens**

You maybe failed to install Cython. Please install it manually. See When an error occurs....

#### Examples says "cuDNN is not enabled"

You failed to build Chainer with cuDNN. If you don't need cuDNN, ignore this message. Otherwise, retry to install Chainer with cuDNN. -vvvv option helps you. See *Install Chainer with CUDA and cuDNN*.

## CHAPTER 2

## **Chainer Tutorial**

## Introduction to Chainer

This is the first section of the Chainer Tutorial. In this section, you will learn about the following things:

- Pros and cons of existing frameworks and why we are developing Chainer
- Simple example of forward and backward computation
- Usage of links and their gradient computation
- Construction of chains (a.k.a. "model" in most frameworks)
- Parameter optimization
- Serialization of links and optimizers

After reading this section, you will be able to:

- Compute gradients of some arithmetics
- · Write a multi-layer perceptron with Chainer

#### **Core Concept**

As mentioned on the front page, Chainer is a flexible framework for neural networks. One major goal is flexibility, so it must enable us to write complex architectures simply and intuitively.

Most existing deep learning frameworks are based on the "**Define-and-Run**" scheme. That is, first a network is defined and fixed, and then the user periodically feeds it with mini-batches. Since the network is statically defined before any forward/backward computation, all the logic must be embedded into the network architecture as *data*. Consequently, defining a network architecture in such systems (e.g. Caffe) follows a declarative approach. Note that one can still produce such a static network definition using imperative languages (e.g. torch.nn, Theano-based frameworks, and TensorFlow).

In contrast, Chainer adopts a **"Define-by-Run"** scheme, i.e., the network is defined on-the-fly via the actual forward computation. More precisely, Chainer stores the history of computation instead of programming logic. This strategy

enables us to fully leverage the power of programming logic in Python. For example, Chainer does not need any magic to introduce conditionals and loops into the network definitions. The Define-by-Run scheme is the core concept of Chainer. We will show in this tutorial how to define networks dynamically.

This strategy also makes it easy to write multi-GPU parallelization, since logic comes closer to network manipulation. We will review such amenities in later sections of this tutorial.

Note: In the example code of this tutorial, we assume for simplicity that the following symbols are already imported:

```
import numpy as np
import chainer
from chainer import cuda, Function, gradient_check, report, training, utils, Variable
from chainer import datasets, iterators, optimizers, serializers
from chainer import Link, Chain, ChainList
import chainer.functions as F
import chainer.links as L
from chainer.training import extensions
```

These imports appear widely in Chainer code and examples. For simplicity, we omit these imports in this tutorial.

#### Forward/Backward Computation

As described above, Chainer uses the "Define-by-Run" scheme, so forward computation itself *defines* the network. In order to start forward computation, we have to set the input array to a *Variable* object. Here we start with a simple ndarray with only one element:

```
>>> x_data = np.array([5], dtype=np.float32)
>>> x = Variable(x_data)
```

A Variable object has basic arithmetic operators. In order to compute  $y = x^2 - 2x + 1$ , just write:

>>> y = x \* \* 2 - 2 \* x + 1

The resulting y is also a Variable object, whose value can be extracted by accessing the data attribute:

```
>>> y.data
array([ 16.], dtype=float32)
```

What y holds is not only the result value. It also holds the history of computation (or computational graph), which enables us to compute its differentiation. This is done by calling its *backward()* method:

>>> y.backward()

This runs *error backpropagation* (a.k.a. *backprop* or *reverse-mode automatic differentiation*). Then, the gradient is computed and stored in the grad attribute of the input variable x:

```
>>> x.grad
array([ 8.], dtype=float32)
```

Also we can compute gradients of intermediate variables. Note that Chainer, by default, releases the gradient arrays of intermediate variables for memory efficiency. In order to preserve gradient information, pass the retain\_grad argument to the backward method:

```
>>> z = 2*x
>>> y = x**2 - z + 1
>>> y.backward(retain_grad=True)
>>> z.grad
array([-1.], dtype=float32)
```

All these computations are easily generalized to a multi-element array input. Note that if we want to start backward computation from a variable holding a multi-element array, we must set the *initial error* manually. This is done simply by setting the grad attribute of the output variable:

```
>>> x = Variable(np.array([[1, 2, 3], [4, 5, 6]], dtype=np.float32))
>>> y = x**2 - 2*x + 1
>>> y.grad = np.ones((2, 3), dtype=np.float32)
>>> y.backward()
>>> x.grad
array([[ 0., 2., 4.],
        [ 6., 8., 10.]], dtype=float32)
```

**Note:** Many functions taking *Variable* object(s) are defined in the *functions* module. You can combine them to realize complicated functions with automatic backward computation.

#### Links

In order to write neural networks, we have to combine functions with *parameters* and optimize the parameters. You can use **links** to do this. A link is an object that holds parameters (i.e. optimization targets).

The most fundamental ones are links that behave like regular functions while replacing some arguments by their parameters. We will introduce higher level links, but here think of links as simply functions with parameters.

One of the most frequently used links is the Linear link (a.k.a. *fully-connected layer* or *affine transformation*). It represents a mathematical function f(x) = Wx + b, where the matrix W and the vector b are parameters. This link corresponds to its pure counterpart *linear()*, which accepts x, W, b as arguments. A linear link from three-dimensional space to two-dimensional space is defined by the following line:

>>> f = L.Linear(3, 2)

Note: Most functions and links only accept mini-batch input, where the first dimension of the input array is considered as the *batch dimension*. In the above Linear link case, input must have shape of (N, 3), where N is the mini-batch size.

The parameters of a link are stored as attributes. Each parameter is an instance of *Variable*. In the case of the Linear link, two parameters, W and b, are stored. By default, the matrix W is initialized randomly, while the vector b is initialized with zeros.

```
>>> f.W.data
array([[ 1.01847613, 0.23103087, 0.56507462],
        [ 1.29378033, 1.07823515, -0.56423163]], dtype=float32)
>>> f.b.data
array([ 0., 0.], dtype=float32)
```

An instance of the Linear link acts like a usual function:

```
>>> x = Variable(np.array([[1, 2, 3], [4, 5, 6]], dtype=np.float32))
>>> y = f(x)
>>> y.data
array([[ 3.1757617 , 1.75755572],
            [ 8.61950684, 7.18090773]], dtype=float32)
```

Gradients of parameters are computed by the *backward()* method. Note that gradients are **accumulated** by the method rather than overwritten. So first you must clear gradients to renew the computation. It can be done by calling the *cleargrads()* method.

```
>>> f.cleargrads()
```

**Note:** *cleargrads ()* is introduced in v1.15 to replace *zerograds ()* for efficiency. *zerograds ()* is left only for backward compatibility.

Now we can compute the gradients of parameters by simply calling the backward method.

```
>>> y.grad = np.ones((2, 2), dtype=np.float32)
>>> y.backward()
>>> f.W.grad
array([[ 5., 7., 9.],
       [ 5., 7., 9.]], dtype=float32)
>>> f.b.grad
array([ 2., 2.], dtype=float32)
```

#### Write a model as a chain

Most neural network architectures contain multiple links. For example, a multi-layer perceptron consists of multiple linear layers. We can write complex procedures with parameters by combining multiple links like this:

```
>>> 11 = L.Linear(4, 3)
>>> 12 = L.Linear(3, 2)
>>> def my_forward(x):
... h = 11(x)
... return 12(h)
```

Here the L indicates the *links* module. A procedure with parameters defined in this way is hard to reuse. More Pythonic way is combining the links and procedures into a class:

```
>>> class MyProc(object):
... def __init__(self):
... self.l1 = L.Linear(4, 3)
... self.l2 = L.Linear(3, 2)
...
... def forward(self, x):
... h = self.l1(x)
... return self.l2(h)
```

In order to make it more reusable, we want to support parameter management, CPU/GPU migration, robust and flexible save/load features, etc. These features are all supported by the *Chain* class in Chainer. Then, what we have to do here is just define the above class as a subclass of Chain:

```
>>> class MyChain(Chain):
... def __init__(self):
```

**Note:** We often define a single forward method of a link by \_\_\_\_\_\_ operator. Such links and chains are callable and behave like regular functions of Variables.

It shows how a complex chain is constructed by simpler links. Links like 11 and 12 are called *child links* of MyChain. Note that Chain itself inherits Link. It means we can define more complex chains that hold MyChain objects as their child links.

Another way to define a chain is using the ChainList class, which behaves like a list of links:

```
>>> class MyChain2 (ChainList):
         def __init__(self):
. . .
              super(MyChain2, self).__init__(
. . .
                  L.Linear(4, 3),
. . .
                  L.Linear(3, 2),
. . .
              )
. . .
. . .
         def __call__(self, x):
. . .
             h = self[0](x)
. . .
              return self[1](h)
. . .
```

ChainList can conveniently use an arbitrary number of links, however if the number of links is fixed like in the above case, the Chain class is recommended as a base class.

#### Optimizer

In order to get good values for parameters, we have to optimize them by the *Optimizer* class. It runs a numerical optimization algorithm on a given link. Many algorithms are implemented in the optimizers module. Here we use the simplest one, called Stochastic Gradient Descent (SGD):

```
>>> model = MyChain()
>>> optimizer = optimizers.SGD()
>>> optimizer.use_cleargrads()
>>> optimizer.setup(model)
```

The method use\_cleargrads () is for efficiency. See use\_cleargrads () for detail.

The method setup () prepares for the optimization given a link.

Some parameter/gradient manipulations, e.g. weight decay and gradient clipping, can be done by setting *hook functions* to the optimizer. Hook functions are called after the gradient computation and right before the actual update of parameters. For example, we can set weight decay regularization by running the next line beforehand:

>>> optimizer.add\_hook(chainer.optimizer.WeightDecay(0.0005))

Of course, you can write your own hook functions. It should be a function or a callable object, taking the optimizer as the argument.

There are two ways to use the optimizer. One is using it via *Trainer*, which we will see in the following sections. The other way is using it directly. We here review the latter case. *If you are interested in getting able to use the optimizer in a simple way, skip this section and go to the next one.* 

There are two further ways to use the optimizer directly. One is manually computing gradients and then calling the update() method with no arguments. Do not forget to clear the gradients beforehand!

```
>>> x = np.random.uniform(-1, 1, (2, 4)).astype('f')
>>> model.cleargrads()
>>> # compute gradient here...
>>> loss = F.sum(model(chainer.Variable(x)))
>>> loss.backward()
>>> optimizer.update()
```

The other way is just passing a loss function to the *update()* method. In this case, *cleargrads()* is automatically called by the update method, so the user does not have to call it manually.

```
See Optimizer.update() for the full specification.
```

#### Trainer

When we want to train neural networks, we have to run *training loops* that update the parameters many times. A typical training loop consists of the following procedures:

- 1. Iterations over training datasets
- 2. Preprocessing of extracted mini-batches
- 3. Forward/backward computations of the neural networks
- 4. Parameter updates
- 5. Evaluations of the current parameters on validation datasets
- 6. Logging and printing of the intermediate results

Chainer provides a simple yet powerful way to make it easy to write such training processes. The training loop abstraction mainly consists of two components:

- Dataset abstraction. It implements 1 and 2 in the above list. The core components are defined in the dataset module. There are also many implementations of datasets and iterators in *datasets* and *iterators* modules, respectively.
- **Trainer**. It implements 3, 4, 5, and 6 in the above list. The whole procedure is implemented by *Trainer*. The way to update parameters (3 and 4) is defined by *Updater*, which can be freely customized. 5 and 6 are implemented by instances of *Extension*, which appends an extra procedure to the training loop. Users can freely customize the training procedure by adding extensions. Users can also implement their own extensions.

We will see how to use Trainer in the example section below.

#### Serializer

Before proceeding to the first example, we introduce Serializer, which is the last core feature described in this page. Serializer is a simple interface to serialize or deserialize an object. *Link*, *Optimizer*, and *Trainer* supports serialization.

Concrete serializers are defined in the serializers module. It supports NumPy NPZ and HDF5 formats.

For example, we can serialize a link object into NPZ file by the serializers.save\_npz() function:

>>> serializers.save\_npz('my.model', model)

It saves the parameters of model into the file 'my.model' in NPZ format. The saved model can be read by the *serializers.load\_npz()* function:

>>> serializers.load\_npz('my.model', model)

**Note:** Note that only the parameters and the *persistent values* are serialized by this serialization code. Other attributes are not saved automatically. You can register arrays, scalars, or any serializable objects as persistent values by the *Link.add\_persistent()* method. The registered values can be accessed by attributes of the name passed to the add\_persistent method.

The state of an optimizer can also be saved by the same functions:

```
>>> serializers.save_npz('my.state', optimizer)
>>> serializers.load_npz('my.state', optimizer)
```

**Note:** Note that serialization of optimizer only saves its internal states including number of iterations, momentum vectors of MomentumSGD, etc. It does not save the parameters and persistent values of the target link. We have to explicitly save the target link with the optimizer to resume the optimization from saved states.

Support of the HDF5 format is enabled if the h5py package is installed. Serialization and deserialization with the HDF5 format are almost identical to those with the NPZ format; just replace  $save_npz()$  and  $load_npz()$  by  $save_hdf5()$  and  $load_hdf5()$ , respectively.

#### Example: Multi-layer Perceptron on MNIST

Now you can solve a multiclass classification task using a multi-layer perceptron (MLP). We use a hand-written digits dataset called MNIST, which is one of the long-standing de facto "hello world" examples used in machine learning. This MNIST example is also found in the examples/mnist directory of the official repository. We show how to use *Trainer* to construct and run the training loop in this section.

We first have to prepare the MNIST dataset. The MNIST dataset consists of 70,000 greyscale images of size 28x28 (i.e. 784 pixels) and corresponding digit labels. The dataset is divided into 60,000 training images and 10,000 test images by default. We can obtain the vectorized version (i.e., a set of 784 dimensional vectors) by *datasets*. *get\_mnist()*.

>>> train, test = datasets.get\_mnist()
...

This code automatically downloads the MNIST dataset and saves the NumPy arrays to the \$(HOME)/.chainer directory. The returned train and test can be seen as lists of image-label pairs (strictly speaking, they are instances of *TupleDataset*).

We also have to define how to iterate over these datasets. We want to shuffle the training dataset for every *epoch*, i.e. at the beginning of every sweep over the dataset. In this case, we can use *iterators*.SerialIterator.

>>> train\_iter = iterators.SerialIterator(train, batch\_size=100, shuffle=**True**)

On the other hand, we do not have to shuffle the test dataset. In this case, we can pass shuffle=False argument to disable the shuffling. It makes the iteration faster when the underlying dataset supports fast slicing.

We also pass repeat=False, which means we stop iteration when all examples are visited. This option is usually required for the test/validation datasets; without this option, the iteration enters an infinite loop.

Next, we define the architecture. We use a simple three-layer rectifier network with 100 units per layer as an example.

```
>>> class MLP (Chain):
       def __init__(self, n_units, n_out):
. . .
            super(MLP, self).__init__(
. . .
                 # the size of the inputs to each layer will be inferred
. . .
                 l1=L.Linear(None, n_units), # n_in -> n_units
. . .
                 l2=L.Linear(None, n_units), # n_units -> n_units
. . .
                 13=L.Linear(None, n_out),
                                               # n_units -> n_out
. . .
             )
. . .
. . .
        def __call__(self, x):
. . .
            h1 = F.relu(self.l1(x))
. . .
            h2 = F.relu(self.12(h1))
. . .
            y = self.13(h2)
. . .
            return y
. . .
```

This link uses relu() as an activation function. Note that the '13' link is the final linear layer whose output corresponds to scores for the ten digits.

In order to compute loss values or evaluate the accuracy of the predictions, we define a classifier chain on top of the above MLP chain:

```
>>> class Classifier (Chain):
        def __init__(self, predictor):
. . .
             super(Classifier, self).__init__(predictor=predictor)
. . .
. . .
        def __call__(self, x, t):
. . .
            y = self.predictor(x)
. . .
            loss = F.softmax_cross_entropy(y, t)
. . .
            accuracy = F.accuracy(y, t)
. . .
             report({'loss': loss, 'accuracy': accuracy}, self)
. . .
             return loss
. . .
```

This Classifier class computes accuracy and loss, and returns the loss value. The pair of arguments x and t corresponds to each example in the datasets (a tuple of an image and a label). *softmax\_cross\_entropy()* computes the loss value given prediction and ground truth labels. *accuracy()* computes the prediction accuracy. We can set an arbitrary predictor link to an instance of the classifier.

The *report ()* function reports the loss and accuracy values to the trainer. For the detailed mechanism of collecting training statistics, see *Reporter*. You can also collect other types of observations like activation statistics in a similar ways.

Note that a class similar to the Classifier above is defined as *chainer.links.Classifier*. So instead of using the above example, we will use this predefined Classifier chain.

```
>>> model = L.Classifier(MLP(100, 10)) # the input size, 784, is inferred
>>> optimizer = optimizers.SGD()
>>> optimizer.setup(model)
```

Now we can build a trainer object.

```
>>> updater = training.StandardUpdater(train_iter, optimizer)
>>> trainer = training.Trainer(updater, (20, 'epoch'), out='result')
```

The second argument (20, 'epoch') represents the duration of training. We can use either epoch or iteration as the unit. In this case, we train the multi-layer perceptron by iterating over the training set 20 times.

In order to invoke the training loop, we just call the *run()* method.

>>> trainer.run()

This method executes the whole training sequence.

The above code just optimizes the parameters. In most cases, we want to see how the training proceeds, where we can use extensions inserted before calling the run method.

These extensions perform the following tasks:

*Evaluator* Evaluates the current model on the test dataset at the end of every epoch.

LogReport Accumulates the reported values and emits them to the log file in the output directory.

**PrintReport** Prints the selected items in the LogReport.

**ProgressBar** Shows the progress bar.

There are many extensions implemented in the *chainer.training.extensions* module. The most important one that is not included above is *snapshot()*, which saves the snapshot of the training procedure (i.e., the Trainer object) to a file in the output directory.

The example code in the *examples/mnist* directory additionally contains GPU support, though the essential part is the same as the code in this tutorial. We will review in later sections how to use GPU(s).

#### How to Write a New Network

#### **Convolutional Network for Visual Recognition Tasks**

In this section, you will learn how to write

- A small convolutional network with a model class that is inherited from Chain,
- A large convolutional network that has several building block networks with ChainList.

After reading this section, you will be able to:

· Write your own original convolutional network in Chainer

A convolutional network (ConvNet) is mainly comprised of convolutional layers. This type of network is commonly used for various visual recognition tasks, e.g., classifying hand-written digits or natural images into given object classes, detectiong objects from an image, and labeling all pixels of an image with the object classes (semantic segmenation), and so on.

In such tasks, a typical ConvNet takes a set of images whose shape is (N, C, H, W), where

- N denotes the number of images in a mini-batch,
- C denotes the number of channels of those images,
- *H* and *W* denote the height and width of those images,

respectively. Then, it typically outputs a fixed-sized vector as membership probabilities over the target object classes. It also can output a set of feature maps that have the corresponding size to the input image for a pixel labeling task, etc.

**Note:** The below example code assumes that some packages are already imported. Please see the details here: tutorial/basic.

#### LeNet5

Here, let's start by defining LeNet5 [*LeCun98*] in Chainer. This is a ConvNet model that has 5 layers comprised of 3 convolutional layers and 2 fully-connected layers. This was proposed to classify hand-written digit images in 1998. In Chainer, the model can be written as follows:

```
class LeNet5 (Chain):
   def __init__(self):
        super(LeNet5, self).___init___(
            conv1=L.Convolution2D(
                in_channels=1, out_channels=6, ksize=5, stride=1),
            conv2=L.Convolution2D(
                in_channels=6, out_channels=16, ksize=5, stride=1),
            conv3=L.Convolution2D(
                in_channels=16, out_channels=120, ksize=4, stride=1),
            fc4=L.Linear(None, 84),
            fc5=L.Linear(84, 10),
        )
        self.train = True
    def __call__(self, x):
       h = F.sigmoid(self.conv1(x))
       h = F.max_pooling_2d(h, 2, 2)
        h = F.sigmoid(self.conv2(h))
        h = F.max_pooling_2d(h, 2, 2)
        h = F.sigmoid(self.conv3(h))
        h = F.sigmoid(self.fc4(h))
        if self.train:
            return self.fc5(h)
        return F.softmax(self.fc5(h))
```

A typical way to write your network is creating a new class inherited from *Chain* class. When defining your model in this way, typically, all the layers which have trainable parameters are registered to the model by giving the objects of *Link* to the superclass's constructer as keyword arguments (see the above \_\_\_init\_\_()).

There is also another way to do the same thing. For example, *add\_link()* of *Chain* class enables to register the trainable layers (i.e., *Link* s) to the model, so that the above \_\_init\_\_() can also be written as follows:

```
def __init__(self
   super(LeNet5, self).__init__()
   self.add_link('conv1', L.Convolution2D(1, 6, 5, 1))
   self.add_link('conv2', L.Convolution2D(6, 16, 5, 1))
   self.add_link('conv3', L.Convolution2D(16, 120, 4, 1))
   self.add_link('fc4', L.Linear(None, 84))
   self.add_link('fc5', L.Linear(84, 10))
   self.train = True
```

(Argments to Convolution2D are given without keywords here for simplicity.)

The model class is instantiated before the forward and backward computations. To give input images and label vectors simply by calling the model object like a function, \_\_call\_\_() is usually defined in the model class. This method performs the forward computation of the model. Chainer uses the powerful autograd system for any computational graphs written with Links calls a corresponding *Function* inside of it), so that you don't need to explicitly write the code for backward computations in the model. Just prepare the data, then give it to the model. The way this works is the resulting output *Variable* from the forward computation has a *backward()* method to perform autograd. In the above model, \_\_call\_() has a if statement at the end to switch its behavior by the model's running mode, i.e., training mode or not. When it's in training mode, this method returns the output value of the last layer as is to compute the loss later on, otherwise it returns a prediction result by calculating *softmax()*.

If you don't want to write conv1 and the other layers more than once, you can also write the model like in this way:

```
class LeNet5 (Chain):
   def __init__(self):
        super(LeNet5, self).__init__()
        net = [('conv1', L.Convolution2D(1, 6, 5, 1))]
        net += [('_sigm1', F.Sigmoid())]
        net += [('_mpool1', F.MaxPooling2D(2, 2))]
        net += [('conv2', L.Convolution2D(6, 16, 5, 1))]
        net += [('_sigm2', F.Sigmoid())]
        net += [('_mpool2', F.MaxPooling2D(2, 2))]
       net += [('conv3', L.Convolution2D(16, 120, 4, 1))]
       net += [('_sigm3', F.Sigmoid())]
       net += [('_mpool3', F.MaxPooling2D(2, 2))]
        net += [('fc4', L.Linear(None, 84))]
        net += [('_sigm4', F.Sigmoid())]
        net += [('fc5', L.Linear(84, 10))]
        net += [('_sigm5', F.Sigmoid())]
        for n in net:
            if not n[0].startswith('_'):
                self.add_link(*n)
        self.forward = net
        self.train = True
   def __call__(self, x):
        for n, f in self.forward:
            if not n.startswith('_'):
                x = getattr(self, n)(x)
            else:
                x = f(x)
        if self.train:
            return x
        return F.softmax(x)
```

This code creates a list of all *Link* s and *Function* s after calling its superclass's constructor. Then the elements of the list are registered to this model as trainable layers when the name of an element doesn't start with \_ character. This operation can be freely replaced with many other ways because those names are just designed to select *Link* s

only from the list net easily. *Function* doesn't have any trainable parameters, so that we can't register it to the model with *add\_link()*, but we want to use *Function*'s for constructing a forward path. The list net is stored as an attribute attr:*forward* to refer it in \_\_call\_\_(). In \_\_call\_\_(), it retrieves all layers in the network from self.forward sequentially regardless of what types of object (*Link* or *Function*) it is, and gives the input variable or the intermediate output from the previous layer to the current layer. The last part of the \_\_call\_\_() to switch its behavior by the training/inference mode is the same as the former way.

#### Ways to calculate loss

When you train the model with label vector t, the loss should be calculated using the output from the model. There also are several ways to calculate the loss:

```
model = LeNet5()
# Input data and label
x = np.random.rand(32, 1, 28, 28).astype(np.float32)
t = np.random.randint(0, 10, size=(32,)).astype(np.int32)
# Forward computation
y = model(x)
# Loss calculation
loss = F.softmax_cross_entropy(y, t)
```

This is a primitive way to calculate a loss value from the output of the model. On the other hand, the loss computation can be included in the model itself by wrapping the model object (*Chain* or *ChainList* object) with a class inherited from *Chain*. The outer *Chain* should take the model defined above and register it through the constructor of its superclass or *add\_link()*. *Chain* is actually inherited from *Link*, so that *Chain* itself can also be registedred as a trainable *Link* to another *Chain*. Actually, *Classifier* class to wrap the model and add the loss computation to the model already exists. Actually, there is already a *Classifier* class that can be used to wrap the model and include the loss computation as well. It can be used like this:

```
model = L.Classifier(LeNet5())
# Foward & Loss calculation
loss = model(x, t)
```

This class takes a model object as an iput argument and registers it to a predictor property as a trained parameter. As shown above, the returned object can then be called like a function in which we pass x and t as the input arguments and the resulting loss value (which we recall is a *Variable*) is returned.

See the detailed implementation of *Classifier* from here: *chainer.links.Classifier* and check the implementation by looking at the source.

From the above examples, we can see that Chainer provides the flexibility to write our original network in many different ways. Such flexibility intends to make it intuitive for users to design new and complex models.

#### VGG16

Next, let's write some larger models in Chainer. When you write a large network consisting of several building block networks, *ChainList* is useful. First, let's see how to write a VGG16 [Simonyan14] model.

```
class VGG16(chainer.ChainList):
    def __init__(self):
        w = chainer.initializers.HeNormal()
```

```
VGGBlock(64),
            VGGBlock(128),
            VGGBlock(256, 3),
            VGGBlock(512, 3),
            VGGBlock (512, 3, True))
        self.train = True
    def __call__(self, x):
        for f in self.children():
            x = f(x, self.train)
        if self.train:
            return x
        return F.softmax(x)
class VGGBlock(chainer.Chain):
    def __init__(self, n_channels, n_convs=2, fc=False):
        w = chainer.initializers.HeNormal()
        super(VGGBlock, self).__init__(
            conv1=L.Convolution2D(None, n_channels, 3, 1, 1, initialW=w),
            conv2=L.Convolution2D(
                n_channels, n_channels, 3, 1, 1, initialW=w))
        if n_convs == 3:
            self.add_link('conv3', L.Convolution2D(
                n_channels, n_channels, 3, 1, 1, initialW=w))
        if fc:
            self.add_link('fc4', L.Linear(None, 4096, initialW=w))
            self.add_link('fc5', L.Linear(4096, 4096, initialW=w))
            self.add_link('fc6', L.Linear(4096, 1000, initialW=w))
        self.n_convs = n_convs
        self.fc = fc
    def __call__(self, x, train):
        h = F.relu(self.conv1(x))
        h = F.relu(self.conv2(h))
        if self.n_convs == 3:
            h = F.relu(self.conv3(h))
        h = F.max_pooling_2d(h, 2, 2)
        if self.fc:
            h = F.dropout(F.relu(self.fc4(h)), train=train)
            h = F.dropout(F.relu(self.fc5(h)), train=train)
            h = self.fc6(h)
        return h
```

super(VGG16, self).\_\_\_init\_\_\_(

That's it. VGG16 is a model which won the 1st place in classification + localization task at ILSVRC 2014, and since then, has become one of the standard models for many different tasks as a pre-trained model. This has 16-layers, so it's called "VGG-16", but we can write this model without writing all layers independently. Since this model consists of several building blocks that have the same architecture, we can build the whole network by re-using the building block definition. Each part of the network is consisted of 2 or 3 convolutional layers and activation function (relu()) following them, and  $max_pooling_2d()$  operations. This block is written as VGGBlock in the above example code. And the whole network just calls this block one by one in sequential manner.

#### ResNet152

How about ResNet? ResNet [*He16*] came in the following year's ILSVRC. It is a much deeper model than VGG16, having up to 152 layers. This sounds super laborious to build, but it can be implemented in almost same manner as VGG16. In the other words, it's easy. One possible way to write ResNet-152 is:

```
class ResNet152 (chainer.Chain):
   def __init__(self, n_blocks=[3, 8, 36, 3]):
        w = chainer.initializers.HeNormal()
        super(ResNet152, self).__init__(
            conv1=L.Convolution2D(
                None, 64, 7, 2, 3, initialW=w, nobias=True),
            bn1=L.BatchNormalization(64),
            res2=ResBlock(n_blocks[0], 64, 64, 256, 1),
            res3=ResBlock(n_blocks[1], 256, 128, 512),
            res4=ResBlock(n_blocks[2], 512, 256, 1024),
            res5=ResBlock(n_blocks[3], 1024, 512, 2048),
            fc6=L.Linear(2048, 1000))
        self.train = True
   def __call__(self, x):
        h = self.bn1(self.conv1(x), test=not self.train)
        h = F.max_pooling_2d(F.relu(h), 2, 2)
       h = self.res2(h, self.train)
       h = self.res3(h, self.train)
       h = self.res4(h, self.train)
       h = self.res5(h, self.train)
       h = F.average_pooling_2d(h, h.shape[2:], stride=1)
       h = self.fc6(h)
        if self.train:
            return h
        return F.softmax(h)
class ResBlock(chainer.ChainList):
   def __init__(self, n_layers, n_in, n_mid, n_out, stride=2):
       w = chainer.initializers.HeNormal()
        super(ResBlock, self).__init__()
        self.add_link(BottleNeck(n_in, n_mid, n_out, stride, True))
        for _ in range(n_layers - 1):
            self.add_link(BottleNeck(n_out, n_mid, n_out))
   def __call__(self, x, train):
        for f in self.children():
            x = f(x, train)
        return x
class BottleNeck(chainer.Chain):
   def __init__(self, n_in, n_mid, n_out, stride=1, proj=False):
        w = chainer.initializers.HeNormal()
        super(BottleNeck, self).__init__(
            conv1x1a=L.Convolution2D(
                n_in, n_mid, 1, stride, 0, initialW=w, nobias=True),
            conv3x3b=L.Convolution2D(
                n_mid, n_mid, 3, 1, 1, initialW=w, nobias=True),
            conv1x1c=L.Convolution2D(
                n_mid, n_out, 1, 1, 0, initialW=w, nobias=True),
```

```
bn_a=L.BatchNormalization (n_mid),
bn_b=L.BatchNormalization (n_mid),
bn_c=L.BatchNormalization (n_out))
if proj:
    self.add_link('convlxlr', L.Convolution2D(
        n_in, n_out, 1, stride, 0, initialW=w, nobias=True))
    self.add_link('bn_r', L.BatchNormalization(n_out))
self.proj = proj
def __call__(self, x, train):
    h = F.relu(self.bn_a(self.convlxla(x), test=not train))
    h = F.relu(self.bn_b(self.conv3x3b(h), test=not train))
    h = self.bn_c(self.convlxlc(h), test=not train)
    if self.proj:
        x = self.bn_r(self.convlxlr(x), test=not train)
    return F.relu(h + x)
```

In the BottleNeck class, depending on the value of the proj argument supplied to the initializer, it will conditionally compute a convolutional layer convlx1r which will extend the number of channels of the input x to be equal to the number of channels of the output of convlx1c, and followed by a batch normalization layer before the final ReLU layer. Writing the building block in this way improves the re-usability of a class. It switches not only the behavior in \_\_class\_\_() by flags but also the parameter registration. In this case, when proj is False, the BottleNeck doesn't have convlx1r and bn\_r layers, so the memory usage would be efficient compared to the case when it registers both anyway and just ignore them if proj is False.

Using nested Chains and ChainList for sequential part enables us to write complex and very deep models easily.

#### **Use Pre-trained Models**

Various ways to write your models were described above. It turns out that VGG16 and ResNet are very useful as general feature extractors for many kinds of tasks, including but not limited to image classification. So, Chainer provides you with the pre-trained VGG16 and ResNet-50/101/152 models with a simple API. You can use these models as follows:

```
from chainer.links import VGG16Layers
model = VGG16Layers()
```

When *VGG16Layers* is instantiated, the pre-trained parameters are automatically downloaded from the author's server. So you can immediately start to use VGG16 with pre-trained weight as a good image feature extractor. See the details of this model here: *chainer.links.VGG16Layers*.

In the case of ResNet models, there are three variations differing in the number of layers. We have chainer.links. ResNet50, chainer.links.ResNet101, and chainer.links.ResNet152 models with easy parameter loading feature. ResNet's pre-trained parameters are not available for direct downloading, so you need to download the weight from the author's web page first, and then place it into the dir \$CHAINER\_DATSET\_ROOT/pfnet/ chainer/models or your favorite place. Once the preparation is finished, the usage is the same as VGG16:

```
from chainer.links import ResNet152Layers
model = ResNet152layers()
```

Please see the details of usage and how to prepare the pre-trained weights for ResNet here: chainer.links. ResNet50

#### References

#### **Recurrent Nets and their Computational Graph**

In this section, you will learn how to write

- recurrent nets with full backprop,
- · recurrent nets with truncated backprop,
- evaluation of networks with few memory.

After reading this section, you will be able to:

- Handle input sequences of variable length
- Truncate upper stream of the network during forward computation
- · Use volatile variables to prevent network construction

#### **Recurrent Nets**

Recurrent nets are neural networks with loops. They are often used to learn from sequential input/output. Given an input stream  $x_1, x_2, \ldots, x_t, \ldots$  and the initial state  $h_0$ , a recurrent net iteratively updates its state by  $h_t = f(x_t, h_{t-1})$ , and at some or every point in time t, it outputs  $y_t = g(h_t)$ . If we expand the procedure along the time axis, it looks like a regular feed-forward network except that same parameters are repeatedly used within the network.

Here we learn how to write a simple one-layer recurrent net. The task is language modeling: given a finite sequence of words, we want to predict the next word at each position without peeking the successive words. Suppose there are 1,000 different word types, and that we use 100 dimensional real vectors to represent each word (a.k.a. word embedding).

Let's start from defining the recurrent neural net language model (RNNLM) as a chain. We can use the *chainer*. *links.LSTM* link that implements a fully-connected stateful LSTM layer. This link looks like an ordinary fully-connected layer. On construction, you pass the input and output size to the constructor:

>>> 1 = L.LSTM(100, 50)

Then, call on this instance l(x) executes one step of LSTM layer:

```
>>> l.reset_state()
>>> x = Variable(np.random.randn(10, 100).astype(np.float32))
>>> y = l(x)
```

Do not forget to reset the internal state of the LSTM layer before the forward computation! Every recurrent layer holds its internal state (i.e. the output of the previous call). At the first application of the recurrent layer, you must reset the internal state. Then, the next input can be directly fed to the LSTM instance:

```
>>> x2 = Variable(np.random.randn(10, 100).astype(np.float32))
>>> y2 = l(x2)
```

Based on this LSTM link, let's write our recurrent network as a new chain:

```
class RNN (Chain):
    def __init__ (self):
        super(RNN, self).__init__ (
        embed=L.EmbedID(1000, 100), # word embedding
        mid=L.LSTM(100, 50), # the first LSTM layer
        out=L.Linear(50, 1000), # the feed-forward output layer
```

```
)
def reset_state(self):
    self.mid.reset_state()

def __call__(self, cur_word):
    # Given the current word ID, predict the next word.
    x = self.embed(cur_word)
    h = self.mid(x)
    y = self.out(h)
    return y

rnn = RNN()
model = L.Classifier(rnn)
optimizer = optimizers.SGD()
optimizer.setup(model)
```

Here *EmbedID* is a link for word embedding. It converts input integers into corresponding fixed-dimensional embedding vectors. The last linear link out represents the feed-forward output layer.

The RNN chain implements a *one-step-forward computation*. It does not handle sequences by itself, but we can use it to process sequences by just feeding items in a sequence straight to the chain.

Suppose we have a list of word variables x\_list. Then, we can compute loss values for the word sequence by simple for loop.

```
def compute_loss(x_list):
    loss = 0
    for cur_word, next_word in zip(x_list, x_list[1:]):
        loss += model(cur_word, next_word)
    return loss
```

Of course, the accumulated loss is a Variable object with the full history of computation. So we can just call its *backward()* method to compute gradients of the total loss according to the model parameters:

```
# Suppose we have a list of word variables x_list.
rnn.reset_state()
model.cleargrads()
loss = compute_loss(x_list)
loss.backward()
optimizer.update()
```

Or equivalently we can use the compute\_loss as a loss function:

```
rnn.reset_state()
optimizer.update(compute_loss, x_list)
```

#### Truncate the Graph by Unchaining

Learning from very long sequences is also a typical use case of recurrent nets. Suppose the input and state sequence is too long to fit into memory. In such cases, we often truncate the backpropagation into a short time range. This technique is called *truncated backprop*. It is heuristic, and it makes the gradients biased. However, this technique works well in practice if the time range is long enough.

How to implement truncated backprop in Chainer? Chainer has a smart mechanism to achieve truncation, called **backward unchaining**. It is implemented in the *Variable.unchain\_backward()* method. Backward unchaining

starts from the Variable object, and it chops the computation history backwards from the variable. The chopped variables are disposed automatically (if they are not referenced explicitly from any other user object). As a result, they are no longer a part of computation history, and are not involved in backprop anymore.

Let's write an example of truncated backprop. Here we use the same network as the one used in the previous subsection. Suppose we are given a very long sequence, and we want to run backprop truncated at every 30 time steps. We can write truncated backprop using the model defined above:

```
loss = 0
count = 0
seqlen = len(x_list[1:])
rnn.reset_state()
for cur_word, next_word in zip(x_list, x_list[1:]):
    loss += model(cur_word, next_word)
    count += 1
    if count % 30 == 0 or count == seqlen:
        model.cleargrads()
        loss.backward()
        loss.unchain_backward()
        optimizer.update()
```

State is updated at model(), and the losses are accumulated to loss variable. At each 30 steps, backprop takes place at the accumulated loss. Then, the *unchain\_backward()* method is called, which deletes the computation history backward from the accumulated loss. Note that the last state of model is not lost, since the RNN instance holds a reference to it.

The implementation of truncated backprop is simple, and since there is no complicated trick on it, we can generalize this method to different situations. For example, we can easily extend the above code to use different schedules between backprop timing and truncation length.

#### Network Evaluation without Storing the Computation History

On evaluation of recurrent nets, there is typically no need to store the computation history. While unchaining enables us to walk through unlimited length of sequences with limited memory, it is a bit of a work-around.

As an alternative, Chainer provides an evaluation mode of forward computation which does not store the computation history. This is enabled by just passing volatile flag to all input variables. Such variables are called *volatile variables*.

Volatile variable is created by passing volatile='on' at the construction:

```
x_list = [Variable(..., volatile='on') for _ in range(100)] # list of 100 words
loss = compute_loss(x_list)
```

Note that we cannot call loss.backward() to compute the gradient here, since the volatile variable does not remember the computation history.

Volatile variables are also useful to evaluate feed-forward networks to reduce the memory footprint.

Variable's volatility can be changed directly by setting the Variable.volatile attribute. This enables us to combine a fixed feature extractor network and a trainable predictor network. For example, suppose we want to train a feedforward network predictor\_func, which is located on top of another fixed pre-trained network fixed\_func. We want to train predictor\_func without storing the computation history for fixed\_func. This is simply done by following code snippets (suppose x\_data and y\_data indicate input data and label, respectively):

```
x = Variable(x_data, volatile='on')
feat = fixed_func(x)
```

```
feat.volatile = 'off'
y = predictor_func(feat)
y.backward()
```

At first, the input variable x is volatile, so fixed\_func is executed in volatile mode, i.e. without memorizing the computation history. Then the intermediate variable feat is manually set to non-volatile, so predictor\_func is executed in non-volatile mode, i.e., with memorizing the history of computation. Since the history of computation is only memorized between variables feat and y, the backward computation stops at the feat variable.

**Warning:** It is not allowed to mix volatile and non-volatile variables as arguments to same function. If you want to create a variable that behaves like a non-volatile variable while can be mixed with volatile ones, use 'auto' flag instead of 'off' flag.

#### Making it with Trainer

The above codes are written with plain Function/Variable APIs. When we write a training loop, it is better to use Trainer, since we can then easily add functionalities by extensions.

Before implementing it on Trainer, let's clarify the training settings. We here use Penn Tree Bank dataset as a set of sentences. Each sentence is represented as a word sequence. We concatenate all sentences into one long word sequence, in which each sentence is separated by a special word <eos>, which stands for "End of Sequence". This dataset is easily obtained by *chainer.datasets.get\_ptb\_words()*. This function returns train, validation, and test dataset, each of which is represented as a long array of integers. Each integer represents a word ID.

Our task is to learn a recurrent neural net language model from the long word sequence. We use words in different locations to form mini-batches. It means we maintain B indices pointing to different locations in the sequence, read from these indices at each iteration, and increment all indices after the read. Of course, when one index reaches the end of the whole sequence, we turn the index back to 0.

In order to implement this training procedure, we have to customize the following components of Trainer:

- Iterator. Built-in iterators do not support reading from different locations and aggregating them into a minibatch.
- Update function. The default update function does not support truncated BPTT.

When we write a dataset iterator dedicated to the dataset, the dataset implementation can be arbitrary; even the interface is not fixed. On the other hand, the iterator must support the *Iterator* interface. The important methods and attributes to implement are batch\_size, epoch, epoch\_detail, is\_new\_epoch, iteration, \_\_next\_\_, and serialize. Following is a code from the official example in the examples/ptb directory.

```
from __future__ import division

class ParallelSequentialIterator(chainer.dataset.Iterator):
    def __init__(self, dataset, batch_size, repeat=True):
        self.dataset = dataset
        self.batch_size = batch_size
        self.epoch = 0
        self.is_new_epoch = False
        self.repeat = repeat
        self.offsets = [i * len(dataset) // batch_size for i in range(batch_size)]
        self.iteration = 0

    def __next__(self):
        length = len(self.dataset)
        if not self.repeat and self.iteration * self.batch_size >= length:
```

```
raise StopIteration
        cur_words = self.get_words()
        self.iteration += 1
        next_words = self.get_words()
        epoch = self.iteration * self.batch_size // length
        self.is_new_epoch = self.epoch < epoch</pre>
        if self.is_new_epoch:
            self.epoch = epoch
        return list(zip(cur_words, next_words))
    @property
    def epoch_detail(self):
        return self.iteration * self.batch_size / len(self.dataset)
    def get_words(self):
        return [self.dataset[(offset + self.iteration) % len(self.dataset)]
                for offset in self.offsets]
    def serialize(self, serializer):
        self.iteration = serializer('iteration', self.iteration)
        self.epoch = serializer('epoch', self.epoch)
train_iter = ParallelSequentialIterator(train, 20)
val_iter = ParallelSequentialIterator(val, 1, repeat=False)
```

Although the code is slightly long, the idea is simple. First, this iterator creates offsets pointing to positions equally spaced within the whole sequence. The i-th examples of mini-batches refer the sequence with the i-th offset. The iterator returns a list of tuples of the current words and the next words. Each mini-batch is converted to a tuple of integer arrays by the concat\_examples function in the standard updater (see the previous tutorial).

Backprop Through Time is implemented as follows.

```
def update_bptt(updater):
    loss = 0
    for i in range(35):
        batch = train_iter.__next__()
        x, t = chainer.dataset.concat_examples(batch)
        loss += model(chainer.Variable(x), chainer.Variable(t))
    model.cleargrads()
    loss.backward()
    loss.unchain_backward()    # truncate
    optimizer.update()
updater = training.StandardUpdater(train_iter, optimizer, update_bptt)
```

In this case, we update the parameters on every 35 consecutive words. The call of unchain\_backward cuts the history of computation accumulated to the LSTM links. The rest of the code for setting up Trainer is almost same as one given in the previous tutorial.

In this section we have demonstrated how to write recurrent nets in Chainer and some fundamental techniques to manage the history of computation (a.k.a. computational graph). The example in the examples/ptb directory implements truncated backprop learning of a LSTM language model from the Penn Treebank corpus. In the next section, we will review how to use GPU(s) in Chainer.

## Using GPU(s) in Chainer

In this section, you will learn about the following things:

- Relationship between Chainer and CuPy
- Basics of CuPy
- Single-GPU usage of Chainer
- Multi-GPU usage of model-parallel computing
- Multi-GPU usage of data-parallel computing

After reading this section, you will be able to:

- Use Chainer on a CUDA-enabled GPU
- Write model-parallel computing in Chainer
- Write data-parallel computing in Chainer

### **Relationship between Chainer and CuPy**

**Note:** As of the release of v1.3.0, Chainer changes its GPU backend from PyCUDA to CuPy. CuPy covers all features of PyCUDA used by Chainer, though their interfaces are not compatible.

Chainer uses *CuPy* as its backend for GPU computation. In particular, the *cupy.ndarray* class is the GPU array implementation for Chainer. CuPy supports a subset of features of NumPy with a compatible interface. It enables us to write a common code for CPU and GPU. It also supports PyCUDA-like user-defined kernel generation, which enables us to write fast implementations dedicated to GPU.

Note: The *chainer.cuda* module imports many important symbols from CuPy. For example, the cupy namespace is referred as cuda.cupy in the Chainer code. Note that the *chainer.cuda* module can be imported even if CUDA is not installed.

Chainer uses a memory pool for GPU memory allocation. As shown in the previous sections, Chainer constructs and destructs many arrays during learning and evaluating iterations. It is not well suited for CUDA architecture, since memory allocation and release in CUDA (i.e. cudaMalloc and cudaFree functions) synchronize CPU and GPU computations, which hurts performance. In order to avoid memory allocation and deallocation during the computation, Chainer uses CuPy's memory pool as the standard memory allocator. Chainer changes the default allocator of CuPy to the memory pool, so user can use functions of CuPy directly without dealing with the memory allocator.

#### Basics of cupy.ndarray

Note: CuPy does not require explicit initialization, so cuda.init() function is deprecated.

CuPy is a GPU array backend that implements a subset of NumPy interface. The *cupy.ndarray* class is in its core, which is a compatible GPU alternative of numpy.ndarray. CuPy implements many functions on *cupy.ndarray* objects. See the reference for the supported subset of NumPy API. Understanding NumPy might help utilizing most features of CuPy. See the NumPy documentation for learning it.

The main difference of *cupy.ndarray* from numpy.ndarray is that the content is allocated on the device memory. The allocation takes place on the current device by default. The current device can be changed by *cupy.cuda*. *Device* object as follows:

```
with cupy.cuda.Device(1):
    x_on_gpu1 = cupy.array([1, 2, 3, 4, 5])
```

Most operations of CuPy is done on the current device. Be careful that it causes an error to process an array on a non-current device.

Chainer provides some convenient functions to automatically switch and choose the device. For example, the *chainer.cuda.to\_gpu()* function copies a numpy.ndarray object to a specified device:

```
x_cpu = np.ones((5, 4, 3), dtype=np.float32)
x_gpu = cuda.to_gpu(x_cpu, device=1)
```

It is equivalent to the following code using CuPy:

```
x_cpu = np.ones((5, 4, 3), dtype=np.float32)
with cupy.cuda.Device(1):
    x_gpu = cupy.array(x_cpu)
```

Moving a device array to the host can be done by chainer.cuda.to\_cpu() as follows:

```
x_cpu = cuda.to_cpu(x_gpu)
```

It is equivalent to the following code using CuPy:

with x\_gpu.device: x\_cpu = x\_gpu.get()

**Note:** The *with* statements in these codes are required to select the appropriate CUDA device. If user uses only one device, these device switching is not needed. *chainer.cuda.to\_cpu()* and *chainer.cuda.to\_gpu()* functions automatically switch the current device correctly.

Chainer also provides a convenient function *chainer.cuda.get\_device\_from\_id()* and *chainer.cuda.get\_device\_from\_array()* to select a device. The former function accepts an integer or None. When None is given, it returns *a dummy device object*. Otherwise, it returns a corresponding device object. The latter function accepts CuPy array or NumPy array. When a NumPy array is given, it returns *a dummy device object*. Otherwise, it returns a corresponding device object. Otherwise, it returns a corresponding device object to the give CuPy array. The dummy device object also supports *with* statements like the above example but does nothing. Here are some other examples:

```
cuda.get_device_from_id(1).use()
x_gpul = cupy.empty((4, 3), dtype='f') # 'f' indicates float32
with cuda.get_device_from_id(1):
    x_gpul = cuda.empty((4, 3), dtype='f')
with cuda.get_device_from_array(x_gpul):
    y_gpul = x_gpu + 1
```

Since it accepts NumPy arrays, we can write a function that accepts both NumPy and CuPy arrays with correct device switching:

```
def add1(x):
    with cuda.get_device_from_array(x):
        return x + 1
```

The compatibility of CuPy with NumPy enables us to write CPU/GPU generic code. It can be made easy by the *chainer.cuda.get\_array\_module()* function. This function returns the numpy or *cupy* module based on arguments. A CPU/GPU generic function is defined using it like follows:

```
# Stable implementation of log(1 + exp(x))
def softplus(x):
    xp = cuda.get_array_module(x)
    return xp.maximum(0, x) + xp.log1p(xp.exp(-abs(x)))
```

#### **Run Neural Networks on a Single GPU**

Single-GPU usage is very simple. What you have to do is transferring *Link* and input arrays to the GPU beforehand. In this subsection, the code is based on *our first MNIST example in this tutorial*.

A Link object can be transferred to the specified GPU using the to\_gpu() method.

This time, we make the number of input, hidden, and output units configurable. The  $to_gpu()$  method also accepts a device ID like model.to\_gpu(0). In this case, the link object is transferred to the appropriate GPU device. The current device is used by default.

If we use *chainer.training.Trainer*, what we have to do is just let the updater know the device ID to send each mini-batch.

updater = training.StandardUpdater(train\_iter, optimizer, device=0)
trainer = training.Trainer(updater, (20, 'epoch'), out='result')

We also have to specify the device ID for an evaluator extension as well.

trainer.extend(extensions.Evaluator(test\_iter, model, device=0))

When we write down the training loop by hand, we have to transfer each mini-batch to the GPU manually:

```
model.to_gpu()
batchsize = 100
datasize = len(x_train)
for epoch in range(20):
    print('epoch %d' % epoch)
    indexes = np.random.permutation(datasize)
    for i in range(0, datasize, batchsize):
        x = Variable(cuda.to_gpu(x_train[indexes[i : i + batchsize]]))
        t = Variable(cuda.to_gpu(y_train[indexes[i : i + batchsize]]))
        optimizer.update(model, x, t)
```

#### Model-parallel Computation on Multiple GPUs

Parallelization of machine learning is roughly classified into two types called "model-parallel" and "data-parallel". Model-parallel means parallelizations of the computations inside the model. In contrast, data-parallel means parallelizations using data sharding. In this subsection, we show how to use the model-parallel approach on multiple GPUs in Chainer.

*Recall the MNIST example*. Now suppose that we want to modify this example by expanding the network to 6 layers with 2000 units each using two GPUs. In order to make multi-GPU computation efficient, we only make the two GPUs communicate at the third and sixth layer. The overall architecture looks like the following diagram:

```
(GPU0) input --+--> 11 --> 12 --> 13 --+--> 14 --> 15 --> 16 --+--> output

(GPU1) +--> 11 --> 12 --> 13 --+--> 14 --> 15 --> 16 --+
```

We can use the above MLP chain as following diagram:

```
(GPU0) input --+--> mlp1 --+--> mlp2 --+--> output
| | |
(GPU1) +--> mlp1 --+--> mlp2 --+
```

Let's write a link for the whole network.

```
class ParallelMLP(Chain):
   def __init__(self):
        super(ParallelMLP, self).__init__(
            # the input size, 784, is inferred
            mlp1_gpu0=MLP(1000, 2000).to_gpu(0),
            mlp1_gpu1=MLP(1000, 2000).to_gpu(1),
            # the input size, 2000, is inferred
           mlp2_gpu0=MLP(1000, 10).to_gpu(0),
           mlp2_gpu1=MLP(1000, 10).to_gpu(1),
        )
   def __call__(self, x):
       # assume x is on GPU 0
       z0 = self.mlp1_qpu0(x)
        z1 = self.mlp1_gpu1(F.copy(x, 1))
        # sync
        h0 = F.relu(z0 + F.copy(z1, 0))
        h1 = F.relu(z1 + F.copy(z0, 1))
        y0 = self.mlp2_qpu0(h0)
       y1 = self.mlp2_gpu1(h1)
        # sync
        y = y0 + F.copy(y1, 0)
        return y # output is on GPU0
```

Recall that the Link.to\_gpu() method returns the link itself. The copy() function copies an input variable to specified GPU device and returns a new variable on the device. The copy supports backprop, which just reversely transfers an output gradient to the input device.

**Note:** Above code is not parallelized on CPU, but is parallelized on GPU. This is because all the functions in the above code run asynchronously to the host CPU.

An almost identical example code can be found at examples/mnist/train\_mnist\_model\_parallel.py.

### Data-parallel Computation on Multiple GPUs with Trainer

Data-parallel computation is another strategy to parallelize online processing. In the context of neural networks, it means that a different device does computation on a different subset of the input data. In this subsection, we review the way to achieve data-parallel learning on two GPUs.

Suppose again our task is *the MNIST example*. This time we want to directly parallelize the three-layer network. The most simple form of data-parallelization is parallelizing the gradient computation for a distinct set of data. First, define a model and optimizer instances:

```
model = L.Classifier(MLP(1000, 10)) # the input size, 784, is inferred
optimizer = optimizers.SGD()
optimizer.setup(model)
```

Recall that the MLP link implements the multi-layer perceptron, and the *Classifier* link wraps it to provide a classifier interface. We used *StandardUpdater* in the previous example. In order to enable data-parallel computation with multiple GPUs, we only have to replace it with *ParallelUpdater*.

The devices option specifies which devices to use in data-parallel learning. The device with name 'main' is used as the main device. The original model is sent to this device, so the optimization runs on the main device. In the above example, the model is also cloned and sent to GPU 1. Half of each mini-batch is fed to this cloned model. After every backward computation, the gradient is accumulated into the main device, the parameter update runs on it, and then the updated parameters are sent to GPU 1 again.

See also the example code in examples/mnist/train\_mnist\_data\_parallel.py.

### **Data-parallel Computation on Multiple GPUs without Trainer**

We here introduce a way to write data-parallel computation without the help of *Trainer*. Most users can skip this section. If you are interested in how to write a data-parallel computation by yourself, this section should be informative. It is also helpful to, e.g., customize the *ParallelUpdater* class.

We again start from the MNIST example. At this time, we use a suffix like \_0 and \_1 to distinguish objects on each device. First, we define a model.

model\_0 = L.Classifier(MLP(1000, 10)) # the input size, 784, is inferred

We want to make two copies of this instance on different GPUs. The Link.to\_gpu() method runs in place, so we cannot use it to make a copy. In order to make a copy, we can use Link.copy() method.

```
import copy
model_1 = copy.deepcopy(model_0)
model_0.to_gpu(0)
model_1.to_gpu(1)
```

The *Link.copy()* method copies the link into another instance. *It just copies the link hierarchy*, and does not copy the arrays it holds.

Then, set up an optimizer:

```
optimizer = optimizers.SGD()
optimizer.setup(model_0)
```

Here we use the first copy of the model as *the master model*. Before its update, gradients of model\_1 must be aggregated to those of model\_0.

Then, we can write a data-parallel learning loop as follows:

```
batchsize = 100
datasize = len(x train)
for epoch in range(20):
    print('epoch %d' % epoch)
    indexes = np.random.permutation(datasize)
    for i in range(0, datasize, batchsize):
        x_batch = x_train[indexes[i : i + batchsize]]
        y_batch = y_train[indexes[i : i + batchsize]]
        x0 = Variable(cuda.to_gpu(x_batch[:batchsize//2], 0))
        t0 = Variable(cuda.to_gpu(y_batch[:batchsize//2], 0))
        x1 = Variable(cuda.to_gpu(x_batch[batchsize//2:], 1))
        t1 = Variable(cuda.to_gpu(y_batch[batchsize//2:], 1))
        loss 0 = model 0(x0, t0)
        loss_1 = model_1(x1, t1)
        model_0.cleargrads()
        model_1.cleargrads()
        loss_0.backward()
        loss 1.backward()
        model_0.addgrads(model_1)
        optimizer.update()
        model_1.copyparams(model_0)
```

Do not forget to clear the gradients of both model copies! One half of the mini-batch is forwarded to GPU 0, the other half to GPU 1. Then the gradients are accumulated by the *Link.addgrads()* method. This method adds the gradients of a given link to those of the self. After the gradients are prepared, we can update the optimizer in usual way. Note that the update only modifies the parameters of model\_0. So we must manually copy them to model\_1 using *Link.copyparams()* method.

**Note:** If the batch size used in one model remain the same, the scale of the gradient is roughly proportional to the number of models, when we aggregate gradients from all models by *chainer.Link.addgrads()*. So you need to adjust the batch size and/or learning rate of the optimizer accordingly.

Now you can use Chainer with GPUs. All examples in the examples directory support GPU computation, so please refer to them if you want to know more practices on using GPUs. In the next section, we will show how to define a differentiable (i.e. *backpropable*) function on Variable objects. We will also show there how to write a simple (elementwise) CUDA kernel using Chainer's CUDA utilities.

# Define your own function

In this section, you will learn about the following things:

• How to define a function on variables

- Useful tools to write a function using a GPU
- How to test the function definition

After reading this section, you will be able to:

- Write your own functions
- Define simple kernels in the function definition

### **Differentiable Functions**

Chainer provides a collection of functions in the *functions* module. It covers typical use cases in deep learning, so many existing works can be implemented with them. On the other hand, deep learning is evolving rapidly and we cannot cover all possible functions to define unseen architectures. So it is important to learn how to define your own functions.

First, suppose we want to define an elementwise function f(x, y, z) = x \* y + z. While it is possible to implement this equation using a combination of the \* and + functions, defining it as a single function may reduce memory consumption, so it is not *only* a toy example. Here we call this function *MulAdd*.

Let's start with defining MulAdd working on the CPU. Any function must inherit the *Function* class. The skeleton of a function looks like:

```
class MulAdd(Function):
    def forward_cpu(self, inputs):
        # do forward computation on CPU
        return some_tuple
    def backward_cpu(self, inputs, grad_outputs):
        # do backward computation on CPU
        return some_tuple
```

We must implement *forward\_cpu()* and *backward\_cpu()* methods. The non-self arguments of these functions are tuples of array(s), and these functions must return a tuple of array(s).

Warning: Be careful to return a tuple of arrays even if you have just one array to return.

MulAdd is simple and implemented as follows

```
class MulAdd(Function):
    def forward_cpu(self, inputs):
        x, y, z = inputs
        w = x * y + z
        return w,
    def backward_cpu(self, inputs, grad_outputs):
        x, y, z = inputs
        gw, = grad_outputs
        gx = y * gw
        gy = x * gw
        gz = gw
        return gx, gy, gz
```

As per the warning above, the forward\_cpu method returns a tuple of single element. Note that all arrays appearing in CPU functions are numpy.ndarray. The forward function is straightforward: It unpacks the input tuple,

computes the output, and packs it into a tuple. The backward function is a bit more complicated. Recall the rule of differentiation of multiplication. This example just implements the rule. Look at the return values, the function just packs the gradient of each input in same order and returns them.

By just defining the core computation of forward and backward, Function class provides a chaining logic on it (i.e. storing the history of computation, etc.).

Note: Assuming we implement a (forward) function y = f(x) which takes as input the vector  $x \in \mathbb{R}^n$  and produces as output a vector  $y \in \mathbb{R}^m$ . Then the backward method has to compute

$$\lambda_i = \sum_{j=1}^m \frac{\partial y_j}{\partial x_i} \gamma_j \text{ for } i = 1 \dots n$$

where  $\gamma$  is the grad\_outputs. Note, that the resulting vector  $\lambda$  must have the same shape as the arguments of the forward method.

Now let's define the corresponding GPU methods. You can easily predict that the methods we have to write are named forward\_gpu() and backward\_gpu():

```
class MulAdd(Function):
   def forward_cpu(self, inputs):
        . . .
    def backward_cpu(self, inputs, grad_outputs):
        . . .
   def forward_gpu(self, inputs):
        x, y, z = inputs
        w = x \star y + z
        return w,
    def backward_gpu(self, inputs, grad_outputs):
        x, y, z = inputs
        gw, = grad_outputs
        gx = y * gw
        gy = x * gw
        qz = qw
        return gx, gy, gz
```

In GPU methods, arrays are of type *cupy.ndarray*. We use arithmetic operators defined for this class. These operators implement the basic elementwise arithmetics.

You may find that the definitions of GPU methods are exactly same as those of CPU methods. In that case, we can reduce them to *forward()* and *backward()* methods

```
class MulAdd(Function):
    def forward(self, inputs):
        x, y, z = inputs
        w = x * y + z
        return w,
    def backward(self, inputs, grad_outputs):
        x, y, z = inputs
        gw, = grad_outputs
        gx = y * gw
```

gy = x \* gw gz = gw **return** gx, gy, gz

Since the *cupy.ndarray* class implements many methods of numpy.ndarray, we can write these unified methods in most cases.

The MulAdd function is used as follows:

```
x = Variable(np.random.uniform(-1, 1, (3, 2)).astype(np.float32))
y = Variable(np.random.uniform(-1, 1, (3, 2)).astype(np.float32))
z = Variable(np.random.uniform(-1, 1, (3, 2)).astype(np.float32))
w = MulAdd()(x, y, z)
```

It looks a bit ugly: we have to explicitly instantiate MulAdd before applying it to variables. We also have to be careful that one instance of MulAdd must not be used multiple times, since it acts as a node in the computational graph. In Chainer, we often define a thin wrapper Python function that hide the instantiation:

```
def muladd(x, y, z):
    return MulAdd()(x, y, z)
w = muladd(x, y, z)
```

### Unified forward/backward methods with NumPy/CuPy functions

CuPy also implements many functions that are compatible to those of NumPy. We can write unified forward/backward methods with them. Consider that we want to write a backprop-able function  $f(x, y) = \exp(x) + \exp(y)$ . We name it *ExpAdd* here. It can be written straight-forward as follows

```
class ExpAdd (Function):
   def forward_cpu(self, inputs):
       x, y = inputs
        z = np.exp(x) + np.exp(y)
        return z,
   def backward_cpu(self, inputs, grad_outputs):
        x, y = inputs
        gz, = grad_outputs
        gx = gz * np.exp(x)
       gy = gz * np.exp(y)
        return gx, gy
    def forward_gpu(self, inputs):
       cupy = cuda.cupy
        x, y = inputs
        z = cupy.exp(x) + cupy.exp(y)
       return z,
   def backward_gpu(self, inputs, grad_outputs):
        cupy = cuda.cupy
        x, y = inputs
        gz, = grad_outputs
        gx = gz * cupy.exp(x)
        gy = gz * cupy.exp(y)
```

```
return gx, gy
def expadd(x, y):
    return ExpAdd()(x, y)
```

**Note:** Here we used cuda.cupy instead of directly accessing *cupy*. This is because the cupy module cannot be imported if the CUDA is not installed. In order to keep the implementation valid in non-CUDA environment, we have to defer the access to the cupy module. Note that the *chainer.cuda* module can be imported even if the CUDA is not installed. Of course, the module in such environment is almost useless, but if the interpreter does not run through the code accessing CUDA-dedicated functions, the code is still valid.

The CPU and GPU implementations are almost same, except that numpy is replaced by *cupy* in GPU methods. We can unify these functions using the *cuda.get\_array\_module()* function. This function accepts arbitrary number of arrays, and returns an appropriate module for them. See the following code

```
class ExpAdd(Function):
    def forward(self, inputs):
        xp = cuda.get_array_module(*inputs)
        x, y = inputs
        z = xp.exp(x) + xp.exp(y)
        return z,
    def backward(self, inputs, grad_outputs):
        xp = cuda.get_array_module(*inputs)
        x, y = inputs
        gz, = grad_outputs
        gz, = grad_outputs
        gx = gz * xp.exp(x)
        gy = gz * xp.exp(y)
        return gx, gy
def expadd(x, y):
        return ExpAdd()(x, y)
```

Note that this code works correctly even if CUDA is not installed in the environment. If CUDA is not found, get\_array\_module function always returns numpy. We often use the name xp for the variadic module name, which is analogous to the abbreviation np for NumPy and cp for CuPy.

### Write an Elementwise Kernel Function

Let's turn back to the MulAdd example.

The GPU implementation of MulAdd as shown above is already fast and parallelized on GPU cores. However, it invokes two kernels during each of forward and backward computations. It might hurt performance, since the intermediate temporary arrays are read and written by possibly different GPU cores, which consumes much bandwidth. We can reduce the number of invocations by defining our own kernel. It also reduce the memory consumption.

Most functions only require elementwise operations like MulAdd. CuPy provides a useful tool to define elementwise kernels, the cupy.elementwise.ElementwiseKernel class, and Chainer wraps it by *cuda*. *elementwise()* function. Our MulAdd implementation can be improved as follows:

```
class MulAdd(Function):
    def forward_cpu(self, inputs):
        ...
```

```
def backward_cpu(self, inputs, grad_outputs):
    . . .
def forward_gpu(self, inputs):
    cupy = cuda.cupy
    x, y, z = inputs
    w = cuda.elementwise(
        'float32 x, float32 y, float32 z',
        'float32 w',
        'w = x * y + z',
        'muladd_fwd')(x, y, z)
    return w,
def backward_gpu(self, inputs, grad_outputs):
    x, y, z = inputs
    gw, = grad_outputs
    gx, gy = cuda.elementwise(
        'float32 x, float32 y, float32 gw',
        'float32 gx, float32 gy',
        ...
           gx = y \star gw;
           gy = x \star gw;
        · · · ,
        'muladd_bwd')(x, y, gw)
    qz = qw
    return gx, gy, gz
```

*cuda.elementwise()* function accepts the essential implementation of the kernel function, and returns a kernel invocation function (actually, it returns ElementwiseKernel object, which is callable). In typical usage, we pass four arguments to this function as follows:

- 1. Input argument list. This is a comma-separated string each entry of which consists of a type specification and an argument name.
- 2. Output argument list in the same format as the input argument list.
- 3. Body of *parallel loop*. We can use the input/output argument names as an element of these arrays.
- 4. Name of the kernel function, which is shown in debuggers and profilers.

Above code is not compiled on every forward/backward computation thanks to two caching mechanisms provided by *cuda.elementwise()*.

The first one is *binary caching: cuda.elementwise()* function caches the compiled binary in the \$(HOME)/. cupy/kernel\_cache directory with a hash value of the CUDA code, and reuses it if the given code matches the hash value. This caching mechanism is actually implemented in CuPy.

The second one is *upload caching*: Given a compiled binary code, we have to upload it to the current GPU in order to execute it. *cuda.elementwise()* function memoizes the arguments and the current device, and if it is called with the same arguments for the same device, it reuses the previously uploaded kernel code.

The above MulAdd code only works for float32 arrays. The ElementwiseKernel also supports the type-variadic kernel definition. In order to define variadic kernel functions, you can use *type placeholder* by placing a single character as type specifier:

```
class MulAdd(Function):
    def forward_cpu(self, inputs):
```

```
. . .
def backward_cpu(self, inputs, grad_outputs):
    . . .
def forward_gpu(self, inputs):
    cupy = cuda.cupy
    x, y, z = inputs
    w = cuda.elementwise(
        'T x, T y, T z',
        'T w',
        'w = x * y + z',
        'muladd_fwd')(x, y, z)
    return w,
def backward_gpu(self, inputs, grad_outputs):
    x, y, z = inputs
    gw, = grad_outputs
    gx, gy = cuda.elementwise(
        'T x, T y, T gw',
        'T gx, T gy',
        ...
           gx = y \star gw;
           gy = x \star gw;
        · · ·,
        'muladd_bwd')(x, y, gw)
    gz = gw
    return gx, gy, gz
```

The type placeholder T indicates an arbitrary data type that CuPy supports.

There are more functionalities on user-defined kernels in CuPy. See the CuPy documentation on user-defined kernels for more details.

### Links that wrap functions

Some functions are meant to be combined with parameters. In such case, it is useful to write a small **link** that wraps the function. We have already seen how to define a chain that wraps other links (by inheriting *Chain* class). Here we study how to define a link that does not hold any other links.

As the first example, suppose that we want to implement elementwise product function between the input array and the parameter array. It can be defined as follows:

```
class EltwiseParamProduct(Link):
    def __init__(self, shape):
        # By passing a shape of the parameter, the initializer allocates a
        # parameter variable of the shape.
        super(EltwiseParamProduct, self).__init__(W=shape)
        self.W.data[...] = np.random.randn(*shape)
    def __call__(self, x):
        return self.W * x
```

We can also initialize the parameter after the initialization by the Link.add\_param() method.

```
class EltwiseParamProduct(Link):
    def __init__(self, shape):
        super(EltwiseParamProduct, self).__init__()
        self.add_param('W', shape)
        self.W.data[...] = np.random.randn(*shape)
    def __call__(self, x):
        return self.W * x
```

Note that the initializer and the *add\_param()* method does not initialize elements of the parameter array. We have to manually initialize the elements by random values, zeros, etc.

For another example, assume we want to define a simple linear layer. It is already defined as *Linear*, so this is an educational example. The linear layer is divided into two parts: a function and its wrapper link. First, we have to define a function on variables:

```
class LinearFunction(Function):
    def forward(self, inputs):
        x, W, b = inputs
        return x.dot(W.T) + b,
    def backward(self, inputs, grad_outputs):
        x, W, b = inputs
        gy, = grad_outputs
        gx = gy.dot(W)
        gW = gy.T.dot(x)
        gb = gy.sum(axis=0)
        return gx, gW, gb
def linear(x, W, b):
    return LinearFunction()(x, W, b)
```

This function takes three arguments: input, weight, and bias. It can be used as a part of model definition, though is inconvenient since the user have to manage the weight and bias parameters directly. In order to make a convenient module, let's wrap it into a link:

```
class Linear(Link):
    def __init__(self, in_size, out_size):
        super(Linear, self).__init__(W=(out_size, in_size), b=out_size)
        self.W.data[...] = np.random.randn(out_size, in_size) / math.sqrt(in_size)
        self.b.data.fill(0)
    def __call__(self, x):
        return linear(x, self.W, self.b)
```

This link hides the parameters of the linear layer.

**Note:** An advanced tip to implement functions: if you want to preserve some information between forward and backward computations (e.g. to cache some arrays), you can store it as attributes. Be careful that it might increase the memory consumption during the whole forward-backward computation. If you want to train very large networks on a GPU with limited memory, it is not recommended to cache arrays between forward and backward. There is one exception for this: caching the output arrays does not change the memory consumption, because they are also held by the output Variable objects.

**Warning:** You should not assume a one-to-one match of calls of forward and backward. Some users may call backward more than once after one forward call.

### **Testing Function**

In order to isolate the cause of learning failure from implementation bugs, it is important to test function implementations. Chainer provides simple utilities to help writing unit tests. They are defined in the *gradient\_check* module.

The most important test utility is the *numerical\_grad()* function. This function computes the numerical gradient of given function using finite differences. It can be used as follows

```
x = np.random.randn(4, 3).astype(np.float32)
gy = np.ones((4, 3), dtype=np.float32)
f = lambda: (x * x,)
gx = gradient_check.numerical_grad(f, (x,), (gy,))
```

f is a closure that returns a tuple of array(s) computed from input arrays. The second and third arguments of  $numerical\_grad()$  are tuples of input arrays and output gradient arrays, respectively. The code above computes the numerical gradients of sum(f(x)), where sum indicates the summation over all elements. The summation can be weighted by changing gy.  $numerical\_grad()$  function also accepts additional eps argument, which indicates the quantization width of finite differences.

**Note:** *numerical\_grad()* function accepts both CPU and GPU arrays. Note that we cannot mix CPU and GPU arrays.

Another utility is *chainer.testing.assert\_allclose()* function. This is similar to numpy.testing. assert\_allclose() function. The difference is that Chainer's version accepts CPU and GPU arrays as inputs. We can mix them in one invocation of *chainer.testing.assert\_allclose()*. The default values of optional arguments are also different.

Here is a typical usage of gradient checking utilities. This is a test example of functions.relu() function

```
import unittest
from chainer import testing
class TestReLU(unittest.TestCase):
    def test_backward_cpu(self):
        x = Variable(np.random.randn(3, 2).astype(np.float32))
        y = F.relu(x)
        y.grad = np.random.randn(3, 2).astype(np.float32)
        y.backward()
        f = lambda: (F.relu(x).data,)
        gx, = gradient_check.numerical_grad(f, (x.data,), (y.grad,))
        testing.assert_allclose(gx, x.grad)
```

The first four lines of the test code are simple forward and backward computation of ReLU function. The next two lines compute numerical gradient using the same forward function without backward routine. And at last, we compare these

two results elementwise. Note that the above test code can be easily modified to test GPU version just by replacing CPU arrays to GPU arrays.

You can find many examples of function tests under tests/chainer\_tests/function\_tests directory.

# **Type check**

In this section, you will learn about the following things:

- Basic usage of type check
- Detail of type information
- · Internal mechanism of type check
- More complicated cases
- Call functions
- Typical type check example

After reading this section, you will be able to:

• Write a code to check types of input arguments of your own functions

### Basic usage of type check

When you call a function with an invalid type of array, you sometimes receive no error, but get an unexpected result by broadcasting. When you use CUDA with an illegal type of array, it causes memory corruption, and you get a serious error. These bugs are hard to fix. Chainer can check preconditions of each function, and helps to prevent such problems. These conditions may help a user to understand specification of functions.

Each implementation of *Function* has a method for type check, check\_type\_forward(). This function is called just before the forward() method of the *Function* class. You can override this method to check the condition on types and shapes of arguments.

```
check_type_forward() gets an argument in_types:
```

```
def check_type_forward(self, in_types):
    ...
```

in\_types is an instance of *TypeInfoTuple*, which is a sub-class of tuple. To get type information about the first argument, use in\_types[0]. If the function gets multiple arguments, we recommend to use new variables for readability:

```
x_type, y_type = in_types
```

In this case,  $x_type$  represents the type of the first argument, and  $y_type$  represents the second one.

We describe usage of in\_types with an example. When you want to check if the number of dimension of  $x_type$  equals to 2, write this code:

utils.type\_check.expect(x\_type.ndim == 2)

When this condition is true, nothing happens. Otherwise this code throws an exception, and the user gets a message like this:

```
Traceback (most recent call last):
...
InvalidType: Expect: in_types[0].ndim == 2
Actual: 3 != 2
```

This error message means that "ndim of the first argument expected to be 2, but actually it is 3".

## **Detail of type information**

You can access three information of x\_type.

- . shape is a tuple of ints. Each value is size of each dimension.
- .ndim is int value representing the number of dimensions. Note that ndim == len(shape)
- .dtype is numpy.dtype representing data type of the value.

You can check all members. For example, the size of the first dimension must be positive, you can write like this:

utils.type\_check.expect(x\_type.shape[0] > 0)

You can also check data types with .dtype:

utils.type\_check.expect(x\_type.dtype == np.float64)

And an error is like this:

```
Traceback (most recent call last):
...
InvalidType: Expect: in_types[0].dtype == <type 'numpy.float64'>
Actual: float32 != <type 'numpy.float64'>
```

You can also check kind of dtype. This code checks if the type is floating point

utils.type\_check.expect(x\_type.dtype.kind == 'f')

You can compare between variables. For example, the following code checks if the first argument and the second argument have the same length:

utils.type\_check.expect(x\_type.shape[1] == y\_type.shape[1])

### Internal mechanism of type check

How does it show an error message like "in\_types[0].ndim == 2"? If x\_type is an object containing ndim member variable, we cannot show such an error message because this equation is evaluated as a boolean value by Python interpreter.

Actually x\_type is a *Expr* objects, and doesn't have a ndim member variable itself. *Expr* represents a syntax tree. x\_type.ndim makes a *Expr* object representing (getattr, x\_type, 'ndim').x\_type.ndim == 2 makes an object like (eq, (getattr, x\_type, 'ndim'), 2).type\_check.expect() gets a *Expr* object and evaluates it. When it is True, it causes no error and shows nothing. Otherwise, this method shows a readable error message.

If you want to evaluate a *Expr* object, call eval() method:

actual\_type = x\_type.eval()

actual\_type is an instance of TypeInfo, while x\_type is an instance of *Expr*. In the same way, x\_type. shape[0].eval() returns an int value.

### More powerful methods

*Expr* class is more powerful. It supports all mathematical operators such as + and \*. You can write a condition that the first dimension of x\_type is the first dimension of y\_type times four:

```
utils.type_check.expect(x_type.shape[0] == y_type.shape[0] * 4)
```

When  $x_type.shape[0] == 3$  and  $y_type.shape[0] == 1$ , users can get the error message below:

```
Traceback (most recent call last):
...
InvalidType: Expect: in_types[0].shape[0] == in_types[1].shape[0] * 4
Actual: 3 != 4
```

To compare a member variable of your function, wrap a value with Variable to show readable error message:

x\_type.shape[0] == utils.type\_check.Variable(self.in\_size, "in\_size")

This code can check the equivalent condition below:

```
x_type.shape[0] == self.in_size
```

However, the latter condition doesn't know the meaning of this value. When this condition is not satisfied, the latter code shows unreadable error message:

```
InvalidType: Expect: in_types[0].shape[0] == 4 # what does '4' mean?
Actual: 3 != 4
```

Note that the second argument of utils.type\_check.Variable is only for readability.

The former shows this message:

### **Call functions**

How to check summation of all values of shape? Expr also supports function call:

```
sum = utils.type_check.Variable(np.sum, 'sum')
utils.type_check.expect(sum(x_type.shape) == 10)
```

Why do we need to wrap the function numpy.sum with utils.type\_check.Variable? x\_type.shape is not a tuple but an object of *Expr* as we have seen before. Therefore, numpy.sum(x\_type.shape) fails. We need to evaluate this function lazily.

The above example produces an error message like this:

```
Traceback (most recent call last):
...
InvalidType: Expect: sum(in_types[0].shape) == 10
Actual: 7 != 10
```

### More complicated cases

How to write a more complicated condition that can't be written with these operators? You can evaluate *Expr* and get its result value with eval() method. Then check the condition and show warning message by hand:

```
x_shape = x_type.shape.eval() # get actual shape (int tuple)
if not more_complicated_condition(x_shape):
    expect_msg = 'Shape is expected to be ...'
    actual_msg = 'Shape is ...'
    raise utils.type_check.InvalidType(expect_msg, actual_msg)
```

Please write a readable error message. This code generates the following error message:

```
Traceback (most recent call last):
...
InvalidType: Expect: Shape is expected to be ...
Actual: Shape is ...
```

### Typical type check example

We show a typical type check for a function.

First check the number of arguments:

```
utils.type_check.expect(in_types.size() == 2)
```

in\_types.size() returns a *Expr* object representing the number of arguments. You can check it in the same way.

And then, get each type:

```
x_type, y_type = in_types
```

Don't get each value before checking in\_types.size(). When the number of argument is illegal, type\_check. expect might output unuseful error messages. For example, this code doesn't work when the size of in\_types is 0:

```
utils.type_check.expect(
    in_types.size() == 2,
    in_types[0].ndim == 3,
)
```

After that, check each type:

```
utils.type_check.expect(
  x_type.dtype == np.float32,
  x_type.ndim == 3,
  x_type.shape[1] == 2,
)
```

The above example works correctly even when  $x_type.ndim == 0$  as all conditions are evaluated lazily.

# CHAPTER 3

# **Chainer Reference Manual**

# **Core functionalities**

### Variable

**class** chainer.**Variable** (*data*, *volatile=OFF*, *name=None*, *grad=None*) Array with a structure to keep track of computation.

Every variable holds a data array of type either numpy.ndarray or cupy.ndarray.

A Variable object may be constructed in two ways: by the user or by some function. When a variable is created by some function as one of its outputs, the variable holds a reference to that function. This reference is used in error backpropagation (a.k.a. backprop). It is also used in *backward unchaining*. A variable that does not hold a reference to its creator is called a *root* variable. A variable is root if it is created by the user, or if the reference is deleted by *unchain\_backward()*.

Users can disable this chaining behavior by setting the volatile flag for the initial variables. When a function gets volatile variables as its inputs, the output variables do not hold references to the function. This acts like unchaining on every function application.

### Parameters

- data (array) Initial data array.
- **volatile** (Flag) Volatility flag. String ('on', 'off', or 'auto') or boolean values can be used, too.
- **name** (*str*) Name of the variable.
- grad (array) Initial gradient array.

### Variables

- data Data array of type either numpy.ndarray or cupy.ndarray.
- grad Gradient array.

- **creator** The function who creates this variable. It is None if the variable is not created by any function.
- **volatile** Ternary *Flag* object. If 'ON', the variable does not keep track of any function applications. See *Flag* for the detail of ternary flags.

### \_\_\_abs\_\_()

Element-wise absolute.

**Returns** Output variable.

**Return type** Variable

\_\_\_add\_\_\_(*rhs*)

Element-wise addition.

Returns Output variable.

Return type Variable

\_\_\_div\_\_\_(*rhs*)

Element-wise division

Returns Output variable.

Return type Variable

\_\_\_getitem\_\_\_(x, slices)

Extract elements from array with specified shape, axes and offsets.

### **Parameters**

- **x** (Variable) A variable to be sliced.
- **slices** (*int*, *slice*, *Ellipsis*, *None*, *integer array-like*, *boolean array-like or tuple of them*) It is an integer, a slice, an ellipsis, a numpy.newaxis, an integer array-like, a boolean array-like or tuple of them.

#### Returns

Variable object which contains sliced array of x.

**Return type** *Variable* 

Note: It only supports types that are supported by CUDA's atomicAdd when an integer array is included in slices. The supported types are numpy.float32, numpy.int32, numpy.uint32, numpy. uint64 and numpy.ulonglong.

**Note:** It does not support slices that contains multiple boolean arrays.

Note: See NumPy document for details of indexing.

### \_\_len\_\_()

Returns the number of elements of the data array.

**Returns** Number of elements of the data array.

Return type int

\_matmul\_\_\_(*rhs*) Matrix multiplication.

Returns Output variable.

Return type Variable

\_mul\_\_(*rhs*)

Element-wise multiplication.

Returns Output variable.

Return type Variable

\_\_neg\_\_()

Element-wise negation.

Returns Output variable.

Return type Variable

\_\_pow\_\_\_(*rhs*)

Element-wise power function.

Returns Output variable.

Return type Variable

 $\__radd\__(rhs)$ 

Element-wise addition.

Returns Output variable.

Return type Variable

**\_\_\_rdiv\_\_**(*rhs*)

Element-wise division.

Returns Output variable.

Return type Variable

**\_\_rmatmul**\_\_ (*rhs*) Matrix multiplication.

Returns Output variable.

**Return type** Variable

**\_\_\_rmul**\_\_(*rhs*)

Element-wise multiplication.

Returns Output variable.

Return type Variable

**\_\_\_rpow\_\_**(*rhs*)

Element-wise power function.

Returns Output variable.

Return type Variable

```
\_rsub\_(rhs)
```

Element-wise subtraction.

Returns Output variable.

### **Return type** Variable

\_\_**rtruediv**\_\_(*rhs*) Element-wise division.

Returns Output variable.

### Return type Variable

 $\_sub\_(rhs)$ 

Element-wise subtraction.

Returns Output variable.

### Return type Variable

\_truediv\_(*rhs*)

Element-wise division

### Returns Output variable.

#### **Return type** Variable

#### addgrad(var)

Accumulates the gradient array from given source variable.

This method just runs self.grad += var.grad, except that the accumulation is even done across the host and different devices.

Parameters var (Variable) - Source variable.

#### **backward**(*retain\_grad=False*)

Runs error backpropagation (a.k.a. backprop) from this variable.

On backprop, *Function.backward()* is called on each *Function* object appearing in the backward graph starting from this variable. The backward graph is represented by backward references from variables to their creators, and from functions to their inputs. The backprop stops at all root variables. Some functions set None as gradients of some inputs, where further backprop does not take place at such input variables.

This method uses grad as the initial error array. User can manually set a gradient array before calling this method. If data contains only one element (i.e., it is scalar) and grad is None, then this method automatically complements 1.0 as the initial error. This is useful on starting backprop from some scalar loss value.

**Parameters retain\_grad**(*bool*) – If True, the gradient arrays of all intermediate variables are kept. Otherwise, grad of the intermediate variables are set to None on appropriate timing, which may reduce the maximum memory consumption.

In most cases of training some models, the purpose of backprop is to compute gradients of parameters, not of variables, so it is recommended to set this flag False.

### cleargrad()

Clears the gradient array.

### copydata (var)

Copies the data array from given source variable.

This method just copies the data attribute from given variable to this variable, except that the copy is even done across the host and different devices.

Parameters var (Variable) - Source variable.

### debug\_print()

Display a summary of the stored data and location of the Variable

### label

Short text that represents the variable.

### reshape(\*shape)

Returns a variable of a different shape and the same content.

### See also:

chainer.functions.reshape() for full documentation,

### set\_creator(gen\_func)

Notifies the variable that the given function is its creator.

**Parameters gen\_func** (Function) – Function object that creates this variable as one of its outputs.

### to\_cpu()

Copies the data and gradient arrays to CPU.

### to\_gpu (device=None)

Copies the data and gradient arrays to specified GPU.

Parameters device – Target device specifier. If omitted, the current device is used.

### transpose (\*axes)

Permute the dimensions of an input variable without copy.

See also:

chainer.functions.transpose() for full documentation.

### unchain\_backward()

Deletes references between variables and functions backward.

After this method completes, intermediate variables and functions that are not referenced from anywhere are deallocated by reference count GC. Also this variable itself deletes the reference to its creator function, i.e. this variable becomes root in the computation graph. It indicates that backprop after unchaining stops at this variable. This behavior is useful to implement truncated BPTT.

### zerograd()

Initializes the gradient array by zeros.

Deprecated since version v1.15: Use *cleargrad()* instead.

### Flag

```
class chainer.Flag
```

Ternary flag object for variables.

It takes three values: ON, OFF, and AUTO.

ON and OFF flag can be evaluated as a boolean value. These are converted to True and False, respectively. AUTO flag cannot be converted to boolean. In this case, ValueError is raised.

Parameters name (str, bool, or None) - Name of the flag. Following values are allowed:

- 'on', 'ON', or True for ON value
- 'off', 'OFF', or False for OFF value
- 'auto', 'AUTO', or None for AUTO value

chainer.ON = ON

Equivalent to Flag('on').

```
chainer.OFF = OFF
Equivalent to Flag('off').
```

chainer.**AUTO = AUTO** Equivalent to Flag('auto').

chainer.flag.aggregate\_flags(flags)

Returns an aggregated flag given a sequence of flags.

If both ON and OFF are found, this function raises an error. Otherwise, either of ON and OFF that appeared is returned. If all flags are AUTO, then it returns AUTO.

Parameters flags (sequence of Flag) - Input flags.

**Returns** The result of aggregation.

Return type Flag

### **Function**

### class chainer.Function

Function on variables with backpropagation ability.

All function implementations defined in *chainer.functions* inherit this class.

The main feature of this class is keeping track of function applications as a backward graph. When a function is applied to *Variable* objects, its *forward()* method is called on data fields of input variables, and at the same time it chains references from output variables to the function and from the function to its inputs.

Note: As of v1.5, a function instance cannot be used twice in any computational graphs. In order to reuse a function object multiple times, use copy.copy() before the function applications to make a copy of the instance.

This restriction also means that we cannot make a *stateful function* anymore. For example, it is now not allowed to let a function hold parameters. Define a function as a pure (stateless) procedure, and use *Link* to combine it with parameter variables.

### Example

Let x an instance of Variable and f an instance of Function taking only one argument. Then a line

```
>>> import numpy, chainer, chainer.functions as F
>>> x = chainer.Variable(numpy.zeros(10))
>>> f = F.Identity()
>>> y = f(x)
```

computes a new variable y and creates backward references. Actually, backward references are set as per the following diagram:

x <--- f <--- y

If an application of another function g occurs as

>>> g = F.Identity() >>> z = g(x)

then the graph grows with a branch:

|--- f <--- y x <-+ |--- g <--- z

Note that the branching is correctly managed on backward computation, i.e. the gradients from f and g are accumulated to the gradient of x.

Every function implementation should provide *forward\_cpu()*, *forward\_gpu()*, *backward\_cpu()* and *backward\_gpu()*. Alternatively, one can provide *forward()* and *backward()* instead of separate methods. Backward methods have default implementations that just return None, which indicates that the function is non- differentiable.

### Variables

- inputs A tuple or list of input variables.
- **outputs** A tuple or list of output variables.
- type\_check\_enable When it is True, the function checks types of input arguments. Set CHAINER\_TYPE\_CHECK environment variable 0 to disable type check, or set the variable directly in your own program.

### \_\_call\_\_(\*inputs)

Applies forward propagation with chaining backward references.

Basic behavior is expressed in documentation of Function class.

**Note:** If the data attribute of input variables exist on GPU device, then, before it calls *forward()* method, the appropriate device is selected, so in most cases implementers do not need to take care of device selection.

**Parameters inputs** – Tuple of input *Variable*, numpy.ndarray or *cupy.ndarray* objects. The volatile flags of all input variables must agree. If the input is an numpy. ndarray or a *cupy.ndarray*, it is automatically wrapped with *Variable*.

Returns One Variable object or a tuple of multiple Variable objects.

### add\_hook (hook, name=None)

Registers the function hook.

### **Parameters**

- hook (FunctionHook) Function hook to be registered.
- **name** (*str*) Name of the function hook. name must be unique among function hooks registered to the function. If None, default name of the function hook is used.

### backward(inputs, grad\_outputs)

Applies backprop to output gradient arrays.

It delegates the procedure to *backward\_cpu()* or *backward\_gpu()* by default. Which it selects is determined by the type of input arrays and output gradient arrays. Implementations of *Function* must implement either CPU/GPU methods or this method, if the function is intended to be backprop-ed.

#### Parameters

- **inputs** Tuple of input arrays.
- grad\_outputs Tuple of output gradient arrays.

**Returns** Tuple of input gradient arrays. Some or all of them can be None, if the function is not differentiable on inputs.

### Return type tuple

**Warning:** Implementations of *Function* must take care that the return value must be a tuple even if it returns only one array.

### backward\_cpu (inputs, grad\_outputs)

Applies backprop to output gradient arrays on CPU.

### Parameters

- **inputs** Tuple of input numpy.ndarray object(s).
- grad\_outputs Tuple of output gradient numpy.ndarray object(s).

**Returns** Tuple of input gradient numpy.ndarray object(s). Some or all of them can be None, if the function is not differentiable on corresponding inputs.

### Return type tuple

**Warning:** Implementations of *Function* must take care that the return value must be a tuple even if it returns only one array.

### backward\_gpu (inputs, grad\_outputs)

Applies backprop to output gradient arrays on GPU.

### Parameters

- **inputs** Tuple of input *cupy*.*ndarray* object(s).
- grad\_outputs Tuple of output gradient cupy.ndarray object(s).

**Returns** Tuple of input gradient *cupy.ndarray* object(s). Some or all of them can be None, if the function is not differentiable on corresponding inputs.

### Return type tuple

**Warning:** Implementations of *Function* must take care that the return value must be a tuple even if it returns only one array.

#### check\_type\_forward(in\_types)

Checks types of input data before forward propagation.

Before *forward()* is called, this function is called. You need to validate types of input data in this function using *the type checking utilities*.

**Parameters in\_types** (TypeInfoTuple) - The type information of input data for forward().

### delete\_hook (name)

Unregisters the function hook.

**Parameters name** (*str*) – the name of the function hook to be unregistered.

### forward(inputs)

Applies forward propagation to input arrays.

It delegates the procedure to *forward\_cpu()* or *forward\_gpu()* by default. Which it selects is determined by the type of input arrays. Implementations of *Function* must implement either CPU/GPU methods or this method.

**Parameters** inputs – Tuple of input array(s).

**Returns** Tuple of output array(s).

**Warning:** Implementations of *Function* must take care that the return value must be a tuple even if it returns only one array.

### forward\_cpu (inputs)

Applies forward propagation to input arrays on CPU.

**Parameters** inputs – Tuple of numpy.ndarray object(s).

**Returns** Tuple of numpy.ndarray object(s).

Return type tuple

**Warning:** Implementations of *Function* must take care that the return value must be a tuple even if it returns only one array.

### forward\_gpu (inputs)

Applies forward propagation to input arrays on GPU.

**Parameters** inputs – Tuple of *cupy.ndarray* object(s).

**Returns** Tuple of *cupy.ndarray* object(s).

Return type tuple

**Warning:** Implementations of *Function* must take care that the return value must be a tuple even if it returns only one array.

### label

Short text that represents the function.

The default implementation returns its type name. Each function should override it to give more information.

### local\_function\_hooks

Ordered Dictionary of registered function hooks.

Contrary to chainer.thread\_local.function\_hooks, which registers its elements to all functions, Function hooks in this property is specific to this function.

### unchain()

Purges in/out variables and this function itself from the graph.

This method is called from Variable.unchain\_backward() method.

### chainer.force\_backprop\_mode(\*args, \*\*kwds)

Enable back-propagation for Variable whose volatile is auto.

When you want to enable back-propagation in *no\_backprop\_mode()*, call this method. In this context, *Variable* object whose volatile attribute is 'auto' behaves like a **volatile** variable. That means you can disable *no\_backprop\_mode()* in this context.

If you call this method outside of *no\_backprop\_mode()* context, it changes nothing. *Variable* object with volatile='auto' behaves like a volatile variable by default.

In this example, the volatility of x and y is 'auto'. In *no\_backprop\_mode()* context, y does not have a computational graph but in *force\_backprop\_mode()* it has a graph.

```
>>> with chainer.no_backprop_mode():
    # Variable with volatile='auto' behaves like volatile='on'
    with chainer.force_backprop_mode():
        # Variable with volatile='auto' behaves like volatile='off'
        y = x + 1
```

See also:

See *no\_backprop\_mode()* for details of back-prop mode.

chainer.no\_backprop\_mode(\*args, \*\*kwds)

Disable back-propagation for Variable whose volatile is auto.

In the default setting a *Variable* object whose volatile attribute is 'auto' behaves like a **non-volatile** variable. That means such a *Variable* object builds a computational graph, consumes memory to store the graph, and you can execute back-propagation for it. With this context such a *Variable* object behaves like a **volatile** variable. So, you can easily switch training and evaluation.

In this example, the volatility of x and y is 'auto'. So, y does not have a computational graph.

```
>>> x = chainer.Variable(numpy.array([1,], 'f'), volatile='auto')
>>> with chainer.no_backprop_mode():
... y = x + 1
```

### Link and Chain

class chainer.Link (\*\*params)

Building block of model definitions.

Link is a building block of neural network models that support various features like handling parameters, defining network fragments, serialization, etc.

Link is the primitive structure for the model definitions. It supports management of parameter variables and *persistent values* that should be incorporated to serialization. Parameters are variables registered via the *add\_param()* method, or given to the initializer method. Persistent values are arrays, scalars, or any other serializable values registered via the *add\_persistent()* method.

**Note:** Whereas arbitrary serializable objects can be registered as persistent values, it is strongly recommended to just register values that should be treated as results of learning. A typical example of persistent values is ones computed during training and required for testing, e.g. running statistics for batch normalization.

Parameters and persistent values are referred by their names. They can be accessed as attributes of the links. Link class itself manages the lists of names of parameters and persistent values to distinguish parameters and persistent values from other attributes.

Link can be composed into more complex models. This composition feature is supported by child classes like *Chain* and *ChainList*. One can create a chain by combining one or more links. See the documents for these classes for details.

As noted above, Link supports the serialization protocol of the *Serializer* class. Note that only parameters and persistent values are saved and loaded. Other attributes are considered as a part of user program (i.e. a part of network definition). In order to construct a link from saved file, other attributes must be identically reconstructed by user codes.

### Example

This is a simple example of custom link definition. Chainer itself also provides many links defined under the *links* module. They might serve as examples, too.

Consider we want to define a simple primitive link that implements a fully-connected layer based on the *linear()* function. Note that this function takes input units, a weight variable, and a bias variable as arguments. Then, the fully-connected layer can be defined as follows:

```
import chainer
import chainer.functions as F
import numpy as np
class LinearLayer(chainer.Link):
    def __init__(self, n_in, n_out):
        # Parameters are initialized as a numpy array of given shape.
        super(LinearLayer, self).__init__(
            W=(n_out, n_in),
            b=(n_out,),
        )
        self.W.data[...] = np.random.randn(n_out, n_in)
        self.b.data.fill(0)
    def __call__(self, x):
        return F.linear(x, self.W, self.b)
```

This example shows that a user can define arbitrary parameters and use them in any methods. Links typically implement the \_\_call\_\_ operator.

**Parameters params** – Names, shapes, and optional dtypes of initial parameters. The keywords are used as the parameter names and the corresponding values consist either of the shape or a tuple of shape and a dtype (*shape, dtype*). If only the shape is supplied, the default dtype will be used.

**Variables name** (*str*) – Name of this link, given by the parent chain (if exists).

### add\_param (name, shape, dtype=<type 'numpy.float32'>, initializer=None)

Registers a parameter to the link.

The registered parameter is saved and loaded on serialization and deserialization, and involved in the optimization. The data and gradient of the variable are initialized by NaN arrays. If initializer is not None, the data is initialized by initializer.

If the supplied name argument corresponds to an uninitialized parameter (that is, one that was added with the *add\_uninitialized\_param()* method), name will be removed from the set of uninitialized parameters.

The parameter is set to an attribute of the link with the given name.

### **Parameters**

- **name** (*str*) Name of the parameter. This name is also used as the attribute name. Any uninitialized parameters with the same name will be removed.
- **shape** (*int* or *tuple* of *ints*) Shape of the parameter array.
- **dtype** Data type of the parameter array.
- **initializer** (chainer.initializer.Initializer) If it is not None, the data is initialized with the given initializer. Note that in this case dtype argument is ignored.

### add\_persistent (name, value)

Registers a persistent value to the link.

The registered value is saved and loaded on serialization and deserialization. The value is set to an attribute of the link.

### **Parameters**

- **name** (*str*) Name of the persistent value. This name is also used for the attribute name.
- **value** Value to be registered.

### add\_uninitialized\_param(name)

Registers an uninitialized parameter to the link.

An uninitialized parameter is defined as a parameter that has a name but that does not yet have a shape. If the shape of a parameter depends on the shape of the inputs to the <u>\_\_call\_\_</u> operator, it can be useful to defer initialization (that is, setting the shape) until the first forward call of the link. Such parameters are intended to be defined as uninitialized parameters in the initializer and then initialized during the first forward call.

An uninitialized parameter is intended to be registered to a link by calling this method in the initializer method. Then, during the first forward call, the shape of the parameter will be determined from the size of the inputs and the parameter must be initialized by calling the *add\_param()* method.

**Parameters name** – (str): Name of the uninitialized parameter.

### addgrads (link)

Accumulates gradient values from given link.

This method adds each gradient array of the given link to corresponding gradient array of this link. The accumulation is even done across host and different devices.

Parameters link (Link) – Source link object.

### children()

Returns a generator of all child links.

Returns A generator object that generates all child links.

### cleargrads()

Clears all gradient arrays.

This method should be called before the backward computation at every iteration of the optimization.

copy()

Copies the link hierarchy to new one.

The whole hierarchy rooted by this link is copied. The copy is basically shallow, except that the parameter variables are also shallowly copied. It means that the parameter variables of copied one are different from ones of original link, while they share the data and gradient arrays.

The name of the link is reset on the copy, since the copied instance does not belong to the original parent chain (even if exists).

Returns Copied link object.

### Return type Link

### copyparams (link)

Copies all parameters from given link.

This method copies data arrays of all parameters in the hierarchy. The copy is even done across the host and devices. Note that this method does not copy the gradient arrays.

Parameters link (Link) – Source link object.

#### has\_uninitialized\_params

Check if the link has uninitialized parameters.

Returns True if the link has any uninitialized parameters. Otherwise returns False.

#### Return type bool

### **links** (*skipself=False*)

Returns a generator of all links under the hierarchy.

**Parameters** skipself (bool) – If True, then the generator skips this link and starts with the first child link.

Returns A generator object that generates all links.

#### namedlinks (skipself=False)

Returns a generator of all (path, link) pairs under the hierarchy.

**Parameters** skipself (bool) – If True, then the generator skips this link and starts with the first child link.

Returns A generator object that generates all (path, link) pairs.

#### namedparams()

Returns a generator of all (path, param) pairs under the hierarchy.

**Returns** A generator object that generates all (path, parameter) pairs. The paths are relative from this link.

#### params()

Returns a generator of all parameters under the link hierarchy.

Returns A generator object that generates all parameters.

### serialize (serializer)

Serializes the link object.

Parameters serializer (AbstractSerializer) - Serializer object.

### to\_cpu()

Copies parameter variables and persistent values to CPU.

This method does not handle non-registered attributes. If some of such attributes must be copied to CPU, the link implementation must override this method to do so.

Returns: self

### to\_gpu (device=None)

Copies parameter variables and persistent values to GPU.

This method does not handle non-registered attributes. If some of such attributes must be copied to GPU, the link implementation must override this method to do so.

Parameters device - Target device specifier. If omitted, the current device is used.

Returns: self

xp

Array module for this link.

Depending on which of CPU/GPU this link is on, this property returns numpy or cupy.

### zerograds()

Initializes all gradient arrays by zero.

This method can be used for the same purpose of cleargrads, but less efficient. This method is left for backward compatibility.

Deprecated since version v1.15: Use *cleargrads()* instead.

class chainer.Chain(\*\*links)

Composable link with object-like interface.

Composability is one of the most important features of neural nets. Neural net models consist of many reusable fragments, and each model itself might be embedded into a larger learnable system. Chain enables us to write a neural net based on composition, without bothering about routine works like collecting parameters, serialization, copying the structure with parameters shared, etc.

This class actually provides a way to compose one or more links into one structure. A chain can contain one or more *child links*. Child link is a link registered to the chain with its own name. The child link is stored to an attribute of the chain with the name. User can write a whole model or a fragment of neural nets as a child class of Chain.

Each chain itself is also a link. Therefore, one can combine chains into higher-level chains. In this way, links and chains construct a *link hierarchy*. Link hierarchy forms a tree structure, where each node is identified by the path from the root. The path is represented by a string like a file path in UNIX, consisting of names of nodes on the path, joined by slashes /.

### Example

This is a simple example of custom chain definition. Chainer itself also provides some chains defined under the *links* module. They might serve as examples, too.

Consider we want to define a multi-layer perceptron consisting of two hidden layers with rectifiers as activation functions. We can use the *Linear* link as a building block:

```
import chainer
import chainer.functions as F
import chainer.links as L
class MultiLayerPerceptron(chainer.Chain):
    def __init__(self, n_in, n_hidden, n_out):
        # Create and register three layers for this MLP
        super(MultiLayerPerceptron, self).__init__(
            layer1=L.Linear(n_in, n_hidden),
            layer2=L.Linear(n_hidden, n_hidden),
            layer3=L.Linear(n_hidden, n_out),
        )
    def __call__(self, x):
```

```
# Forward propagation
h1 = F.relu(self.layer1(x))
h2 = F.relu(self.layer2(h1))
return self.layer3(h2)
```

Child links are registered via the initializer method. They also can be registered by the *add\_link()* method. The forward propagation is often implemented as The \_\_call\_\_ operator as the above example, though it is not mandatory.

**Parameters links** – Child links. The keywords are used as their names. The names are also set to the links.

\_\_\_getitem\_\_\_(*name*) Equivalent to getattr.

add link (name, link)

Registers a child link to this chain.

The registered link is saved and loaded on serialization and deserialization, and involved in the optimization. The registered link is called a child. The child link is set to an attribute of the chain with the given name.

This method also sets the name attribute of the registered link. If the given link already has the name attribute set, then it raises an error.

### Parameters

• name (str) - Name of the child link. This name is also used as the attribute name.

• link (Link) – The link object to be registered.

```
class chainer.ChainList (*links)
```

Composable link with list-like interface.

This is another example of compositional link. Unlike *Chain*, this class can be used like a list of child links. Each child link is indexed by a non-negative integer, and it maintains the current number of registered child links. The *add\_link()* method inserts a new link at the end of the list. It is useful to write a chain with arbitrary number of child links, e.g. an arbitrarily deep multi-layer perceptron.

Note that this class does not implement all methods of list.

Parameters links – Initial child links.

\_\_\_getitem\_\_\_(index)

Returns the child at given index.

**Parameters** index (*int*) – Index of the child in the list.

Returns The index-th child link.

Return type Link

```
_len_()
```

Returns a number of children.

### $add\_link(link)$

Registers a child link to this chain.

The registered link is saved and loaded on serialization and deserialization, and involved in the optimization. The registered link is called a child. The child link is accessible via children() generator, which returns a generator running through the children in registered order. This method also sets the name attribute of the registered link. If the given link already has the name attribute set, then it raises an error.

**Parameters** link (Link) – The link object to be registered.

### Optimizer

### class chainer.Optimizer

Base class of all numerical optimizers.

This class provides basic features for all optimization methods. It optimizes parameters of a *target link*. The target link is registered via the setup() method, and then the update() method updates its parameters based on a given loss function.

Each optimizer implementation must be defined as a child class of Optimizer. It must override update() method. An optimizer can use *internal states* each of which is tied to one of the parameters. State is a dictionary of serializable values (typically arrays of size same as the corresponding parameters). In order to use state dictionaries, the optimizer must override *init\_state()* method (or its CPU/GPU versions, *init\_state\_cpu()* and *init\_state\_gpu()*).

If the optimizer is based on single gradient computation (like most first-order methods), then it should inherit *GradientMethod*, which adds some features dedicated for the first order methods.

Optimizer instance also supports *hook functions*. Hook function is registered by the *add\_hook()* method. Each hook function is called in registration order in advance of the actual parameter update.

### Variables

- target Target link object. It is set by the setup () method.
- t Number of update steps. It must be incremented by the update () method.
- epoch Current epoch. It is incremented by the *new\_epoch()* method.

### accumulate\_grads (grads)

Accumulates gradients from other source.

This method just adds given gradient arrays to gradients that this optimizer holds. It is typically used in data-parallel optimization, where gradients for different shards are computed in parallel and aggregated by this method. This method correctly treats multiple GPU devices.

**Parameters** grads (*Iterable*) – Iterable of gradient arrays to be accumulated.

Deprecated since version v1.5: Use the *chainer.Link.addgrads()* method of the target link instead.

### add\_hook (hook, name=None)

Registers a hook function.

Hook function is typically called right after the gradient computation, though the timing depends on the optimization method.

### Parameters

- hook (function) Hook function. It accepts the optimizer object.
- name (str) Name of the registration. If omitted, hook.name is used by default.

### call\_hooks()

Invokes hook functions in registration order.

#### clip\_grads (maxnorm)

Clips the norm of whole gradients up to the threshold.

Parameters maxnorm (float) – Threshold of gradient L2 norm.

Deprecated since version v1.5: Use the GradientClipping hook function instead.

### compute\_grads\_norm()

Computes the norm of whole gradients.

Returns L2 norm of whole gradients, i.e. square root of sum of square of all gradient elements.

Return type float

**Warning:** This method returns a CPU-computed value, which means that this method synchronizes between CPU and GPU if at least one of the gradients reside on the GPU.

Deprecated since version v1.5.

#### init\_state (param, state)

Initializes the optimizer state corresponding to the parameter.

This method should add needed items to the state dictionary. Each optimizer implementation that uses its own states should override this method or CPU/GPU dedicated versions ( $init\_state\_cpu()$ ) and  $init\_state\_gpu()$ ).

### Parameters

- param (Variable) Parameter variable.
- **state** (*dict*) State dictionary.

### See also:

init\_state\_cpu(), init\_state\_gpu()

#### init\_state\_cpu (param, state)

Initializes the optimizer state on CPU.

This method is called from *init\_state()* by default.

### Parameters

- param (Variable) Parameter variable. Its data array is of type numpy.ndarray.
- **state** (*dict*) State dictionary.

### See also:

init\_state()

### init\_state\_gpu (param, state)

Initializes the optimizer state on GPU.

This method is called from *init\_state()* by default.

#### **Parameters**

- param (Variable) Parameter variable. Its data array is of type cupy.ndarray.
- **state** (*dict*) State dictionary.

### See also:

init\_state()

#### new\_epoch()

Starts a new epoch.

This method increments the epoch count. Note that if the optimizer depends on the epoch count, then user should call this method appropriately at the beginning of each epoch.

### prepare()

Prepares for an update.

This method initializes missing optimizer states (e.g. for newly added parameters after the set up), and copies arrays in each state dictionary to CPU or GPU according to the corresponding parameter array.

### remove\_hook (name)

Removes a hook function.

**Parameters name** (*str*) – Registered name of the hook function to remove.

### serialize (serializer)

Serializes or deserializes the optimizer.

It only saves or loads the following things:

•Optimizer states

•Global states (t and epoch)

It does not saves nor loads the parameters of the target link. They should be separately saved or loaded.

Parameters serializer (AbstractSerializer) – Serializer or deserializer object.

#### setup(link)

Sets a target link and initializes the optimizer states.

Given link is set to the target attribute. It also prepares the optimizer state dictionaries corresponding to all parameters in the link hierarchy. The existing states are discarded.

**Parameters link** (Link) – Target link object.

### update(lossfun=None, \*args, \*\*kwds)

Updates the parameters and optimizer states.

This method updates the parameters of the target link and corresponding optimizer states. The behavior of this method is different for the cases either lossfun is given or not.

If lossfun is given, then this method initializes the gradients by zeros, calls it with given extra arguments, and calls the *backward()* method of its output to compute the gradients. The implementation might call lossfun more than once.

If lossfun is not given, then this method assumes that the gradients of all parameters are already computed. An implementation that requires multiple gradient computations might raise an error on this case.

In both cases, this method invokes the update procedure for all parameters.

### **Parameters**

- **lossfun** (function) Loss function. It accepts arbitrary arguments and returns one *Variable* object that represents the loss (or objective) value. This argument can be omitted for single gradient-based methods. In this case, this method assumes gradient arrays computed.
- kwds (args,) Arguments for the loss function.

### weight\_decay (decay)

Applies weight decay to the parameter/gradient pairs.

**Parameters decay** (*float*) – Coefficient of weight decay.

Deprecated since version v1.5: Use the WeightDecay hook function instead.

#### zero\_grads()

Fills all gradient arrays by zeros.

Deprecated since version v1.5: Use the *chainer.Link.cleargrads()* method for the target link instead.

### class chainer.GradientMethod

Base class of all single gradient-based optimizers.

This is an extension of the *Optimizer* class. Typical gradient methods that just require the gradient at the current parameter vector on an update can be implemented as its child class.

An implementation of a gradient method must override the following methods:

```
•init_state() or both init_state_cpu() and init_state_gpu()
•update_one() or both update_one_cpu() and update_one_gpu()
```

Note: It is recommended to call use\_cleargrads () after creating a GradientMethod object for efficiency.

### call\_hooks()

Invokes hook functions in registration order.

### reallocate\_cleared\_grads()

Reallocate gradients cleared by cleargrad().

This method allocates arrays for all gradients which have None. This method is called before and after every optimizer hook. If an inheriting optimizer does not require this allocation, the optimizer can override this method with a blank function.

### update (lossfun=None, \*args, \*\*kwds)

Updates parameters based on a loss function or computed gradients.

This method runs in two ways.

•If lossfun is given, then use it as a loss function to compute gradients.

•Otherwise, this method assumes that the gradients are already computed.

In both cases, the computed gradients are used to update parameters. The actual update routines are defined by the update\_one() method (or its CPU/GPU versions, update\_one\_cpu() and update\_one\_gpu()).

#### update\_one (param, state)

Updates a parameter based on the corresponding gradient and state.

This method calls appropriate one from update\_param\_cpu() or update\_param\_gpu().

### **Parameters**

- param (Variable) Parameter variable.
- **state** (*dict*) State dictionary.

```
update_one_cpu (param, state)
```

Updates a parameter on CPU.

### Parameters

• param (Variable) - Parameter variable.

• **state** (*dict*) – State dictionary.

update\_one\_gpu (param, state)

Updates a parameter on GPU.

### Parameters

• param (Variable) - Parameter variable.

• **state** (*dict*) – State dictionary.

use\_cleargrads (use=True)

Enables or disables use of *cleargrads* () in update.

**Parameters use** (bool) – If True, this function enables use of *cleargrads*. If False, disables use of *cleargrads* (*zerograds* is used).

**Note:** Note that *update()* calls *zerograds()* by default for backward compatibility. It is recommended to call this method before first call of *update* because *cleargrads* is more efficient than *zerograds*.

### **Hook functions**

```
class chainer.optimizer.WeightDecay(rate)
```

Optimizer hook function for weight decay regularization.

This hook function adds a scaled parameter to the corresponding gradient. It can be used as a regularization.

**Parameters rate** (*float*) – Coefficient for the weight decay.

**Variables** rate (*float*) – Coefficient for the weight decay.

### class chainer.optimizer.Lasso(rate)

Optimizer hook function for Lasso regularization.

This hook function adds a scaled parameter to the sign of each weight. It can be used as a regularization.

**Parameters rate** (*float*) – Coefficient for the weight decay.

**Variables rate** (*float*) – Coefficient for the weight decay.

### class chainer.optimizer.GradientClipping(threshold)

Optimizer hook function for gradient clipping.

This hook function scales all gradient arrays to fit to the defined L2 norm threshold.

**Parameters threshold** (*float*) – L2 norm threshold.

Variables threshold (float) - L2 norm threshold of gradient norm.

This hook function simply adds noise generated by the noise\_func to the gradient. By default it adds time-dependent annealed Gaussian noise to the gradient at every training step:

$$g_t \leftarrow g_t + N(0, \sigma_t^2)$$

where

$$\sigma_t^2 = \frac{\eta}{(1+t)^{\gamma}}$$

with  $\eta$  selected from {0.01, 0.3, 1.0} and  $\gamma = 0.55$ .

## Parameters

- **eta** (*float*) Parameter that defines the scale of the noise, which for the default noise function is recommended to be either 0.01, 0.3 or 1.0.
- **noise\_func** (function) Noise generating function which by default is given by Adding Gradient Noise Improves Learning for Very Deep Networks.

# Serializer

## class chainer.AbstractSerializer

Abstract base class of all serializers and deserializers.

\_\_call\_\_(key, value)

Serializes or deserializes a value by given name.

This operator saves or loads a value by given name.

If this is a serializer, then the value is simply saved at the key. Note that some type information might be missed depending on the implementation (and the target file format).

If this is a descrializer, then the value is loaded by the key. The descrialization differently works on scalars and arrays. For scalars, the value argument is used just for determining the type of restored value to be converted, and the converted value is returned. For arrays, the restored elements are directly copied into the value argument. String values are treated like scalars. If the value argument is None, the type of the restored value will typically be a numpy array but can depend on the particular subclass implementation.

#### **Parameters**

- **key** (*str*) Name of the serialization entry.
- **value** (*scalar*, *array*, *None*, *or str*) Object to be (de)serialized. None is only supported by descrializers.

Returns Serialized or deserialized value.

## **\_\_\_getitem**\_\_(*key*)

Gets a child serializer.

This operator creates a \_child\_ serializer represented by the given key.

**Parameters key** (*str*) – Name of the child serializer.

## class chainer.Serializer

Base class of all serializers.

## save(obj)

Saves an object by this serializer.

This is equivalent to obj.serialize(self).

**Parameters obj** – Target object to be serialized.

## class chainer.Deserializer

Base class of all deserializers.

#### load(*obj*)

Loads an object from this deserializer.

This is equivalent to obj.serialize(self).

**Parameters obj** – Target object to be serialized.

# **Dataset abstraction**

Chainer has a support of common interface of training and validation datasets. The dataset support consists of three components: datasets, iterators, and batch conversion functions.

**Dataset** represents a set of examples. The interface is only determined by combination with iterators you want to use on it. The built-in iterators of Chainer requires the dataset to support \_\_getitem\_\_ and \_\_len\_\_ method. In particular, the \_\_getitem\_\_ method should support indexing by both an integer and a slice. We can easily support slice indexing by inheriting *DatasetMixin*, in which case users only have to implement *get\_example()* method for indexing. Some iterators also restrict the type of each example. Basically, datasets are considered as *stateless* objects, so that we do not need to save the dataset as a checkpoint of the training procedure.

**Iterator** iterates over the dataset, and at each iteration, it yields a mini batch of examples as a list. Iterators should support the *Iterator* interface, which includes the standard iterator protocol of Python. Iterators manage where to read next, which means they are *stateful*.

**Batch conversion function** converts the mini batch into arrays to feed to the neural nets. They are also responsible to send each array to an appropriate device. Chainer currently provides *concat\_examples()* as the only example of batch conversion functions.

These components are all customizable, and designed to have a minimum interface to restrict the types of datasets and ways to handle them. In most cases, though, implementations provided by Chainer itself are enough to cover the usages.

Chainer also has a light system to download, manage, and cache concrete examples of datasets. All datasets managed through the system are saved under *the dataset root directory*, which is determined by the CHAINER\_DATASET\_ROOT environment variable, and can also be set by the *set\_dataset\_root()* function.

## **Dataset representation**

See Dataset examples for dataset implementations.

```
class chainer.dataset.DatasetMixin
```

Default implementation of dataset indexing.

DatasetMixin provides the <u>\_\_\_\_\_\_getitem\_\_</u>() operator. The default implementation uses get\_example() to extract each example, and combines the results into a list. This mixin makes it easy to implement a new dataset that does not support efficient slicing.

Dataset implementation using DatasetMixin still has to provide the \_\_\_\_\_() operator explicitly.

```
__getitem___(index)
```

Returns an example or a sequence of examples.

It implements the standard Python indexing and one-dimensional integer array indexing. It uses the *get\_example()* method by default, but it may be overridden by the implementation to, for example, improve the slicing performance.

**Parameters index** (*int*, *slice*, *list or numpy.ndarray*) – An index of an example or indexes of examples.

**Returns** If index is int, returns an example created by *get\_example*. If index is either slice or one-dimensional list or numpy.ndarray, returns a list of examples created by *get\_example*.

#### Example

```
>>> import numpy
>>> from chainer import dataset
>>> class SimpleDataset(dataset.DatasetMixin):
```

```
def __init__(self, values):
. . .
           self.values = values
. . .
        def __len__(self):
. . .
           return len(self.values)
. . .
        def get_example(self, i):
. . .
            return self.values[i]
. . .
. . .
>>> ds = SimpleDataset([0, 1, 2, 3, 4, 5])
>>> ds[1] # Access by int
1
>>> ds[1:3] # Access by slice
[1, 2]
>>> ds[[4, 0]] # Access by one-dimensional integer list
[4, 0]
>>> index = numpy.arange(3)
>>> ds[index] # Access by one-dimensional integer numpy.ndarray
[0, 1, 2]
```

#### \_len\_()

Returns the number of data points.

## $\texttt{get\_example}\left(i\right)$

Returns the i-th example.

Implementations should override it. It should raise IndexError if the index is invalid.

**Parameters** i (*int*) – The index of the example.

Returns The i-th example.

# **Iterator interface**

See Iterator examples for dataset iterator implementations.

## class chainer.dataset.Iterator

Base class of all dataset iterators.

Iterator iterates over the dataset, yielding a minibatch at each iteration. Minibatch is a list of examples. Each implementation should implement an iterator protocol (e.g., the <u>\_\_next\_\_</u>() method).

Note that, even if the iterator supports setting the batch size, it does not guarantee that each batch always contains the same number of examples. For example, if you let the iterator to stop at the end of the sweep, the last batch may contain a fewer number of examples.

The interface between the iterator and the underlying dataset is not fixed, and up to the implementation.

Each implementation should provide the following attributes (not needed to be writable).

•batch\_size: Number of examples within each minibatch.

•epoch: Number of completed sweeps over the dataset.

•epoch\_detail: Floating point number version of the epoch. For example, if the iterator is at the middle of the dataset at the third epoch, then this value is 2.5.

•previous\_epoch\_detail: The value of epoch\_detail at the previous iteration. This value is None before the first iteration.

•is\_new\_epoch: True if the epoch count was incremented at the last update.

Each implementation should also support serialization to resume/suspend the iteration.

\_iter\_()

Returns self.

# \_\_\_next\_\_()

Returns the next batch.

This is a part of the iterator protocol of Python. It may raise the StopIteration exception when it stops the iteration.

## finalize()

Finalizes the iterator and possibly releases the resources.

This method does nothing by default. Implementation may override it to better handle the internal resources.

next()

Python2 alternative of \_\_\_\_next\_\_\_.

It calls \_\_\_\_\_() by default.

## serialize (serializer)

Serializes the internal state of the iterator.

This is a method to support serializer protocol of Chainer.

**Note:** It should only serialize the internal state that changes over the iteration. It should not serializes what is set manually by users such as the batch size.

# **Batch conversion function**

```
chainer.dataset.concat_examples(batch, device=None, padding=None)
```

Concatenates a list of examples into array(s).

Dataset iterator yields a list of examples. If each example is an array, this function concatenates them along the newly-inserted first axis (called *batch dimension*) into one array. The basic behavior is same for examples consisting of multiple arrays, i.e., corresponding arrays of all examples are concatenated.

For instance, consider each example consists of two arrays (x, y). Then, this function concatenates x 's into one array, and y 's into another array, and returns a tuple of these two arrays. Another example: consider each example is a dictionary of two entries whose keys are 'x' and 'y', respectively, and values are arrays. Then, this function concatenates x 's into one array, and y 's into another array, and returns a dictionary with two entries x and y whose values are the concatenated arrays.

When the arrays to concatenate have different shapes, the behavior depends on the padding value. If padding is None (default), it raises an error. Otherwise, it builds an array of the minimum shape that the contents of all arrays can be substituted to. The padding value is then used to the extra elements of the resulting arrays.

TODO(beam2d): Add an example.

- **batch** (*list*) A list of examples. This is typically given by a dataset iterator.
- **device** (*int*) Device ID to which each array is sent. Negative value indicates the host memory (CPU). If it is omitted, all arrays are left in the original device.
- **padding** Scalar value for extra elements. If this is None (default), an error is raised on shape mismatch. Otherwise, an array of minimum dimensionalities that can accommodate all arrays is created, and elements outside of the examples are padded by this value.

**Returns** Array, a tuple of arrays, or a dictionary of arrays. The type depends on the type of each example in the batch.

```
chainer.dataset.to_device(device, x)
```

Send an array to a given device.

This method send a given array to a given device. This method is used in *concat\_examples()*. You can also use this method in a custom converter method used in *Updater* and *Extension* such as *StandardUpdater* and *Evaluator*.

## Parameters

- **device** (*int* or *None*) Device ID to which an array is sent. If it is negative value, an array is sent to CPU. If it is positive, an array is sent to GPU with the given ID. If it is None, an array is left in the original device.
- x (numpy.ndarray or cupy.ndarray) An array to send.

Returns Converted array.

## **Dataset management**

```
chainer.dataset.get_dataset_root()
```

Gets the path to the root directory to download and cache datasets.

**Returns** The path to the dataset root directory.

## Return type str

```
chainer.dataset.set_dataset_root(path)
```

Sets the root directory to download and cache datasets.

There are two ways to set the dataset root directory. One is by setting the environment variable CHAINER\_DATASET\_ROOT. The other is by using this function. If both are specified, one specified via this function is used. The default dataset root is \$HOME/.chainer/dataset.

**Parameters path** (*str*) – Path to the new dataset root directory.

```
chainer.dataset.cached_download(url)
```

Downloads a file and caches it.

It downloads a file from the URL if there is no corresponding cache. After the download, this function stores a cache to the directory under the dataset root (see set\_dataset\_root(). If there is already a cache for the given URL, it just returns the path to the cache without downloading the same file.

**Parameters url** (*str*) – URL to download from.

**Returns** Path to the downloaded file.

Return type str

chainer.dataset.cache\_or\_load\_file(path, creator, loader)

Caches a file if it does not exist, or loads it otherwise.

This is a utility function used in dataset loading routines. The creator creates the file to given path, and returns the content. If the file already exists, the loader is called instead, and it loads the file and returns the content.

Note that the path passed to the creator is temporary one, and not same as the path given to this function. This function safely renames the file created by the creator to a given path, even if this function is called simultaneously by multiple threads or processes.

- **path** (*str*) Path to save the cached file.
- **creator** Function to create the file and returns the content. It takes a path to temporary place as the argument. Before calling the creator, there is no file at the temporary path.
- loader Function to load the cached file and returns the content.

**Returns** It returns the returned values by the creator or the loader.

# **Training loop abstraction**

Chainer provides a standard implementation of the training loops under the *chainer.training* module. It is built on top of many other core features of Chainer, including Variable and Function, Link/Chain/ChainList, Optimizer, Dataset, and Reporter/Summary. Compared to the training loop abstraction of other machine learning tool kits, Chainer's training framework aims at maximal flexibility, while keeps the simplicity for the typical usages. Most components are pluggable, and users can overwrite the definition.

The core of the training loop abstraction is *Trainer*, which implements the training loop itself. The training loop consists of two parts: one is *Updater*, which actually updates the parameters to train, and the other is *Extension* for arbitrary functionalities other than the parameter update.

Updater and some extensions use *dataset* and Iterator to scan the datasets and load mini batches. The trainer also uses *Reporter* to collect the observed values, and some extensions use *DictSummary* to accumulate them and computes the statistics.

You can find many examples for the usage of this training utilities from the official examples. You can also search the extension implementations from *Trainer extensions*.

## Trainer

class chainer.training.Trainer(updater, stop\_trigger=None, out='result')
The standard training loop in Chainer.

Trainer is an implementation of a training loop. Users can invoke the training by calling the *run()* method.

Each iteration of the training loop proceeds as follows.

- •Update of the parameters. It includes the mini-batch loading, forward and backward computations, and an execution of the update formula. These are all done by the update object held by the trainer.
- •Invocation of trainer extensions in the descending order of their priorities. A trigger object is attached to each extension, and it decides at each iteration whether the extension should be executed. Trigger objects are callable objects that take the trainer object as the argument and return a boolean value indicating whether the extension should be called or not.

Extensions are callable objects that take the trainer object as the argument. There are three ways to define custom extensions: inheriting the *Extension* class, decorating functions by *make\_extension()*, and defining any callable including lambda functions. See *Extension* for more details on custom extensions and how to configure them.

Users can register extensions to the trainer by calling the *extend()* method, where some configurations can be added.

•Trigger object, which is also explained above. In most cases, IntervalTrigger is used, in which case users can simply specify a tuple of the interval length and its unit, like (1000, 'iteration') or (1, 'epoch').

•The order of execution of extensions is determined by their priorities. Extensions of higher priorities are invoked earlier. There are three standard values for the priorities:

- -PRIORITY\_WRITER. This is the priority for extensions that write some records to the observation dictionary. It includes cases that the extension directly adds values to the observation dictionary, or the extension uses the *chainer.report()* function to report values to the observation dictionary.
- -PRIORITY\_EDITOR. This is the priority for extensions that edit the observation dictionary based on already reported values.
- -PRIORITY\_READER. This is the priority for extensions that only read records from the observation dictionary. This is also suitable for extensions that do not use the observation dictionary at all.

•Extensions with invoke\_before\_training flag on are also invoked at the beginning of the training loop. Extensions that update the training status (e.g., changing learning rates) should have this flag to be True to ensure that resume of the training loop correctly recovers the training status.

The current state of the trainer object and objects handled by the trainer can be serialized through the standard serialization protocol of Chainer. It enables us to easily suspend and resume the training loop.

**Note:** The serialization does not recover everything of the training loop. It only recovers the states which change over the training (e.g. parameters, optimizer states, the batch iterator state, extension states, etc.). You must initialize the objects correctly before deserializing the states.

On the other hand, it means that users can change the settings on deserialization. For example, the exit condition can be changed on the deserialization, so users can train the model for some iterations, suspend it, and then resume it with larger number of total iterations.

During the training, it also creates a *Reporter* object to store observed values on each update. For each iteration, it creates a fresh observation dictionary and stores it in the observation attribute.

Links of the target model of each optimizer are registered to the reporter object as observers, where the name of each observer is constructed as the format <optimizer name><link name>. The link name is given by the chainer.Link.namedlink() method, which represents the path to each link in the hierarchy. Other observers can be registered by accessing the reporter object via the reporter attribute.

The default trainer is *plain*, i.e., it does not contain any extensions.

#### Parameters

- updater (Updater) Updater object. It defines how to update the models.
- **stop\_trigger** Trigger that determines when to stop the training loop. If it is not callable, it is passed to IntervalTrigger.

#### Variables

- updater The updater object for this trainer.
- **stop\_trigger** Trigger that determines when to stop the training loop. The training loop stops at the iteration on which this trigger returns True.
- **observation** Observation of values made at the last update. See the Reporter class for details.
- **out** Output directory.
- **reporter** Reporter object to report observed values.

## elapsed\_time

Total time used for the training.

The time is in seconds. If the training is resumed from snapshot, it includes the time of all the previous training to get the current state of the trainer.

**extend** (*extension*, *name=None*, *trigger=None*, *priority=None*, *invoke\_before\_training=None*) Registers an extension to the trainer.

*Extension* is a callable object which is called after each update unless the corresponding trigger object decides to skip the iteration. The order of execution is determined by priorities: extensions with higher priorities are called earlier in each iteration. Extensions with the same priority are invoked in the order of registrations.

If two or more extensions with the same name are registered, suffixes are added to the names of the second to last extensions. The suffix is  $_N$  where N is the ordinal of the extensions.

See *Extension* for the interface of extensions.

#### **Parameters**

- **extension** Extension to register.
- **name** (*str*) Name of the extension. If it is omitted, the default\_name attribute of the extension is used instead. Note that the name would be suffixed by an ordinal in case of duplicated names as explained above.
- **trigger** (*tuple or Trigger*) Trigger object that determines when to invoke the extension. If it is None, extension.trigger is used instead. If it is None and the extension does not have the trigger attribute, the extension is triggered at every iteration by default. If the trigger is not callable, it is passed to IntervalTrigger to build an interval trigger.
- **priority** (*int*) Invocation priority of the extension. Extensions are invoked in the descending order of priorities in each iteration. If this is None, extension.priority is used instead.
- **invoke\_before\_training** (bool or None) If True, the extension is also invoked just before entering the training loop. If this is None, extension. invoke\_before\_training is used instead. This option is mainly used for extensions that alter the training configuration (e.g., learning rates); in such a case, resuming from snapshots require the call of extension to recover the configuration before any updates.

## get\_extension(name)

Returns the extension of a given name.

**Parameters name** (*str*) – Name of the extension.

Returns Extension.

run()

Executes the training loop.

This method is the core of Trainer. It executes the whole loop of training the models.

Note that this method cannot run multiple times for one trainer object.

# Updater

class chainer.training.Updater

Interface of updater objects for trainers.

TODO(beam2d): document it.

#### connect\_trainer(trainer)

Connects the updater to the trainer that will call it.

The typical usage of this method is to register additional links to the reporter of the trainer. This method is called at the end of the initialization of *Trainer*. The default implementation does nothing.

**Parameters trainer** (Trainer) – Trainer object to which the updater is registered.

#### finalize()

Finalizes the updater object.

This method is called at the end of training loops. It should finalize each dataset iterator used in this updater.

#### get\_all\_optimizers()

Gets a dictionary of all optimizers for this updater.

Returns Dictionary that maps names to optimizers.

Return type dict

#### get\_optimizer(name)

Gets the optimizer of given name.

Updater holds one or more optimizers with names. They can be retrieved by this method.

**Parameters name** (*str*) – Name of the optimizer.

**Returns** Optimizer of the name.

**Return type** *Optimizer* 

### serialize (serializer)

Serializes the current state of the updater object.

#### update()

Updates the parameters of the target model.

This method implements an update formula for the training task, including data loading, forward/backward computations, and actual updates of parameters.

This method is called once at each iteration of the training loop.

class chainer.training.StandardUpdater(iterator, optimizer, converter=<function concat\_examples>, device=None, loss\_func=None)

Standard implementation of Updater.

This is the standard implementation of *Updater*. It accepts one or more training datasets and one or more optimizers. The default update routine assumes that there is only one training dataset and one optimizer. Users can override this update routine by inheriting this class and overriding the update\_core() method. Each batch is converted to input arrays by concat\_examples() by default, which can also be manually set by converter argument.

- **iterator** Dataset iterator for the training dataset. It can also be a dictionary of iterators. If this is just an iterator, then the iterator is registered by the name 'main'.
- **optimizer** Optimizer to update parameters. It can also be a dictionary of optimizers. If this is just an optimizer, then the optimizer is registered by the name 'main'.
- **converter** Converter function to build input arrays. Each batch extracted by the main iterator and the device option are passed to this function. *concat\_examples()* is used by default.

- **device** Device to which the training data is sent. Negative value indicates the host memory (CPU).
- **loss\_func** Loss function. The target link of the main optimizer is used by default.

#### Variables

- **converter** Converter function.
- **loss\_func** Loss function. If it is None, the target link of the main optimizer is used instead.
- device Device to which the training data is sent.
- iteration Current number of completed updates.

#### get\_iterator(name)

Gets the dataset iterator of given name.

**Parameters name** (*str*) – Name of the dataset iterator.

**Returns** Corresponding dataset iterator.

#### **Return type** *Iterator*

Implementation of a parallel GPU Updater.

This is an implementation of *Updater* that uses multiple GPUs. It behaves similarly to *StandardUpdater*. The update routine is modified to support data-parallel computation on multiple GPUs in one machine. It is based on synchronous parallel SGD: it parallelizes the gradient computation over a mini-batch, and updates the parameters only in the main device.

## Parameters

- **iterator** Dataset iterator for the training dataset. It can also be a dictionary of iterators. If this is just an iterator, then the iterator is registered by the name 'main'.
- **optimizer** Optimizer to update parameters. It can also be a dictionary of optimizers. If this is just an optimizer, then the optimizer is registered by the name 'main'.
- **converter** Converter function to build input arrays. Each batch extracted by the main iterator is split equally between the devices and then passed with corresponding device option to this function. *concat\_examples()* is used by default.
- models Dictionary of models. The main model should be the same model attached to the 'main' optimizer.
- **devices** Dictionary of devices to which the training data is sent. The devices should be arranged in a dictionary with the same structure as models.
- **loss\_func** Loss function. The model is used as a loss function by default.

## Extension

## class chainer.training.Extension

Base class of trainer extensions.

Extension of *Trainer* is a callable object that takes the trainer object as the argument. It also provides some default configurations as its attributes, e.g. the default trigger and the default priority. This class provides a set of typical default values for these attributes.

There are three ways to define users' own extensions: inheriting this class, decorating closures by *make\_extension()*, or using any callable including lambda functions as extensions. Decorator can slightly reduce the overhead and is much easier to use, while this class provides more flexibility (for example, it can have methods to configure the behavior). Using a lambda function allows one-line coding for simple purposes, but users have to specify the configurations as arguments to *Trainer.extend()*. For a callable not inheriting this class, the default configurations of this class are used unless the user explicitly specifies them in *Trainer.extend()* method.

#### Variables

- **trigger** Default value of trigger for this extension. It is set to (1, 'iteration') by default.
- priority Default priority of the extension. It is set to PRIORITY\_READER by default.
- **invoke\_before\_training** Default flag to decide whether this extension should be invoked before the training starts. The default value is False.

#### \_\_\_call\_\_\_(trainer)

Invokes the extension.

Implementations should override this operator. This method is called at iterations which the corresponding trigger accepts.

Parameters trainer (Trainer) – Trainer object that calls this operator.

### default\_name

Default name of the extension.

It is the name of the class by default. Implementation can override this property, or provide a class attribute to hide it.

## finalize()

Finalizes the extension.

This method is called at the end of the training loop.

## serialize (serializer)

Serializes the extension state.

It is called when a trainer that owns this extension is serialized. It serializes nothing by default.

chainer.training.make\_extension(trigger=None, default\_name=None, priority=None, invoke before training=False, finalizer=None)

Decorator to make given functions into trainer extensions.

This decorator just adds some attributes to a given function. The value of the attributes are given by the arguments of this decorator.

See *Extension* for details of trainer extensions. Most of the default values of arguments also follow those for this class.

- **trigger** Default trigger of the extension.
- **default\_name** Default name of the extension. The name of a given function is used by default.
- **priority** (*int*) Default priority of the extension.
- **invoke\_before\_training** (*bool*) Default flag to decide whether the extension should be invoked before any training.

• **finalizer** – Finalizer function of this extension. The finalizer is called at the end of the training loop.

## Trigger

Trigger is a callable object to decide when to process some specific event within the training loop. It takes a Trainer object as the argument, and returns True if some event should be fired.

It is mainly used to determine when to call an extension. It is also used to determine when to quit the training loop.

chainer.training.get\_trigger(*trigger*) Gets a trigger object.

Trigger object is a callable that accepts a *Trainer* object as an argument and returns a boolean value. When it returns True, various kinds of events can occur depending on the context in which the trigger is used. For example, if the trigger is passed to the *Trainer* as the *stop trigger*, the training loop breaks when the trigger returns True. If the trigger is passed to the *extend()* method of a trainer, then the registered extension is invoked only when the trigger returns True.

This function returns a trigger object based on the argument. If trigger is already a callable, it just returns the trigger. If trigger is None, it returns a trigger that never fires. Otherwise, it passes the value to IntervalTrigger.

- **Parameters trigger** Trigger object. It can be either an already built trigger object (i.e., a callable object that accepts a trainer object and returns a bool value), or a tuple. In latter case, the tuple is passed to *IntervalTrigger*.
- **Returns** trigger if it is a callable, otherwise a *IntervalTrigger* object made from trigger.

# Debug mode

In debug mode, Chainer checks values of variables on runtime and shows more detailed error messages. It helps you to debug your programs. Instead it requires additional overhead time.

In debug mode, Chainer checks all results of forward and backward computation, and if it founds a NaN value, it raises RuntimeError. Some functions and links also check validity of input values.

```
chainer.is_debug()
    Get the debug mode.
```

Returns Return True if Chainer is in debug mode.

Return type bool

```
chainer.set_debug(debug)
Set the debug mode.
```

**Note:** This method changes global state. When you use this method on multi-threading environment, it may affects other threads.

**Parameters debug** (bool) – New debug mode.

```
class chainer.DebugMode (debug)
    Debug mode context.
```

This class provides a context manager for debug mode. When entering the context, it sets the debug mode to the value of *debug* parameter with memorizing its original value. When exiting the context, it sets the debug mode back to the original value.

Parameters debug (bool) – Debug mode used in the context.

# FunctionSet (deprecated)

```
class chainer.FunctionSet(**links)
```

Set of links (as "parameterized functions").

FunctionSet is a subclass of Chain. Function registration is done just by adding an attribute to object.

Deprecated since version v1.5: Use Chain instead.

**Note:** FunctionSet was used for manipulation of one or more parameterized functions. The concept of parameterized function is gone, and it has been replaced by *Link* and *Chain*.

```
___getitem__(key)
```

Returns an attribute by name.

**Parameters** key (*str*) – Name of the attribute.

Returns Attribute.

Example

```
>>> model = chainer.FunctionSet(l1=L.Linear(10, 10),
... l2=L.Linear(10, 10))
>>> l1 = model['l1']  # equivalent to l1 = model.l1
```

#### collect\_parameters()

Returns a tuple of parameters and gradients.

**Returns** Tuple (pair) of two tuples. The first element is a tuple of parameter arrays, and the second is a tuple of gradient arrays.

#### copy\_parameters\_from (params)

Copies parameters from another source without reallocation.

**Parameters params** (*Iterable*) – Iterable of parameter arrays.

## gradients

Tuple of gradient arrays of all registered functions.

The order of gradients is consistent with *parameters* () property.

## parameters

Tuple of parameter arrays of all registered functions.

The order of parameters is consistent with *parameters* () property.

# Utilities

# **CUDA** utilities

Device, context and memory management on CuPy.

Chainer uses CuPy (with very thin wrapper) to exploit the speed of GPU computation. Following modules and classes are imported to cuda module for convenience (refer to this table when reading chainer's source codes).

imported name	original name
chainer.cuda.cupy	сиру
chainer.cuda.ndarray	cupy.ndarray
chainer.cuda.cupy.cuda	cupy.cuda
chainer.cuda.Device	cupy.cuda.Device
chainer.cuda.Event	cupy.cuda.Event
chainer.cuda.Stream	cupy.cuda.Stream

Chainer replaces the default allocator of CuPy by its memory pool implementation. It enables us to reuse the device memory over multiple forward/backward computations, and temporary arrays for consecutive elementwise operations.

# **Devices**

chainer.cuda.get\_device(\*args)

Gets the device from a device object, an ID integer or an array object.

Note: This API is deprecated. Please use :method:'cupy.cuda.get\_device\_from\_id' or :method:'cupy.cuda.get\_device\_from\_array' instead.

This is a convenient utility to select a correct device if the type of arg is unknown (i.e., one can use this function on arrays that may be on CPU or GPU). The returned device object supports the context management protocol of Python for the *with* statement.

**Parameters args** – Values to specify a GPU device. The first device object, integer or *cupy*. *ndarray* object is used to select a device. If it is a device object, it is returned. If it is an integer, the corresponding device is returned. If it is a CuPy array, the device on which this array reside is returned. If any arguments are neither integers nor CuPy arrays, a dummy device object representing CPU is returned.

Returns Device object specified by given args.

## See also:

See *cupy.cuda.Device* for the device selection not by arrays.

chainer.cuda.get\_device\_from\_id(device\_id)
 Gets the device from an ID integer.

**Parameters** device\_id (*int* or *None*) – The ID of the device which this function returns.

```
chainer.cuda.get_device_from_array(*arrays)
```

Gets the device from a list of CuPy array or a single CuPy array.

The device on which the given CuPy array reside is returned.

**Parameters array** (*cupy.ndarray* or list of *cupy.ndarray*) – A CuPy array which this function returns the device corresponding to. If a list of :class:'cupy.ndarray's are given, it returns the first device object of an array in the list.

# CuPy array allocation and copy

Note: As of v1.3.0, the following array construction wrappers are marked as deprecated. Use the corresponding functions of the cupy module instead. The main difference of them is that the default dtype is changed from float32 to float64.

Deprecated functions	Recommended functions
chainer.cuda.empty	cupy.empty()
chainer.cuda.empty_like	<pre>cupy.empty_like()</pre>
chainer.cuda.zeros	cupy.zeros()
chainer.cuda.zeros_like	cupy.zeros_like()
chainer.cuda.ones	cupy.ones()
chainer.cuda.ones_like	cupy.ones_like()
chainer.cuda.full	cupy.full()
chainer.cuda.full_like	cupy.full_like()

chainer.cuda.copy(array, out=None, out\_device=None, stream=None)
Copies a cupy.ndarray object using the default stream.

This function can copy the device array to the destination array on another device.

## Parameters

- array (cupy.ndarray) Array to be copied.
- **out** (cupy.ndarray) Destination array. If it is not None, then out\_device argument is ignored.
- **out\_device** Destination device specifier. Actual device object is obtained by passing this value to get\_device().
- **stream** (cupy.cuda.Stream) CUDA stream.

## Returns

## Copied array.

If out is not specified, then the array is allocated on the device specified by out\_device argument.

## Return type *cupy.ndarray*

chainer.cuda.to\_cpu (*array*, *stream=None*) Copies the given GPU array to host CPU.

## copies the given OFO array to

## Parameters

- **array** Array to be sent to CPU.
- **stream** (cupy.cuda.Stream) CUDA stream.

# Returns

Array on CPU.

If given array is already on CPU, then this function just returns array without performing any copy.

## Return type numpy.ndarray

chainer.cuda.to\_gpu (array, device=None, stream=None)
 Copies the given CPU array to specified device.

#### Parameters

- **array** Array to be sent to GPU.
- **device** Device specifier.
- **stream** (cupy.cuda.Stream) CUDA stream. If not None, the copy runs asynchronously.

#### Returns

Array on GPU.

If array is already on GPU, then this function just returns array without performing any copy. Note that this function does not copy *cupy.ndarray* into specified device.

#### Return type *cupy.ndarray*

## Kernel definition utilities

#### chainer.cuda.memoize(for\_each\_device=False)

Makes a function memoizing the result for each argument and device.

This is a similar version of *cupy.memoize()*. The difference is that this function can be used in the global scope even if CUDA is not available. In such case, this function does nothing.

**Note:** This decorator acts as a dummy if CUDA is not available. It cannot be used for general purpose memoization even if for\_each\_device is set to False.

## chainer.cuda.clear\_memo()

Clears the memoized results for all functions decorated by memoize.

This function works like *cupy.clear\_memo()* as a counterpart for *chainer.cuda.memoize()*. It can be used even if CUDA is not available. In such a case, this function does nothing.

## chainer.cuda.elementwise()

Creates an elementwise kernel function.

This function uses *memoize()* to cache the kernel object, i.e. the resulting kernel object is cached for each argument combination and CUDA device.

The arguments are the same as those for *cupy*.*ElementwiseKernel*, except that the name argument is mandatory.

## chainer.cuda.reduce()

Creates a global reduction kernel function.

This function uses *memoize()* to cache the resulting kernel object, i.e. the resulting kernel object is cached for each argument combination and CUDA device.

The arguments are the same as those for *cupy*.*ReductionKernel*, except that the name argument is mandatory.

#### CPU/GPU generic code support

#### chainer.cuda.get\_array\_module(\*args)

Gets an appropriate one from numpy or cupy.

This is almost equivalent to *cupy.get\_array\_module()*. The differences are that this function can be used even if CUDA is not available and that it will return their data arrays' array module for *Variable* arguments.

Parameters args – Values to determine whether NumPy or CuPy should be used.

**Returns** *cupy* or *numpy* is returned based on the types of the arguments.

Return type module

## cuDNN support

```
chainer.cuda.set_max_workspace_size(size)
Sets the workspace size for cuDNN.
```

Check "cuDNN Library User Guide" for detail.

**Parameters size** – The workspace size for cuDNN.

```
chainer.cuda.get_max_workspace_size()
```

Gets the workspace size for cuDNN.

Check "cuDNN Library User Guide" for detail.

**Returns** The workspace size for cuDNN.

Return type int

## **Common algorithms**

```
class chainer.utils.WalkerAlias (probs)
```

Implementation of Walker's alias method.

This method generates a random sample from given probabilities  $p_1, \ldots, p_n$  in O(1) time. It is more efficient than choice (). This class works on both CPU and GPU.

**Parameters probs** (float list) – Probabilities of entries. They are normalized with *sum(probs)*.

See: Wikipedia article

## sample (shape)

Generates a random sample based on given probabilities.

**Parameters shape** (*tuple of int*) – Shape of a return value.

**Returns** Returns a generated array with the given shape. If a sampler is in CPU mode the return value is a numpy.ndarray object, and if it is in GPU mode the return value is a *cupy*. *ndarray* object.

```
to_gpu()
```

Make a sampler GPU mode.

## Reporter

#### Reporter

#### class chainer.Reporter

Object to which observed values are reported.

Reporter is used to collect values that users want to watch. The reporter object holds a mapping from value names to the actually observed values. We call this mapping *observations*.

When a value is passed to the reporter, an object called *observer* can be optionally attached. In this case, the name of the observer is added as the prefix of the value name. The observer name should be registered beforehand.

See the following example:

```
>>> from chainer import Reporter, report, report_scope
>>>
>>> reporter = Reporter()
>>> observer = object() # it can be an arbitrary (reference) object
>>> reporter.add_observer('my_observer:', observer)
>>> observation = {}
>>> with reporter.scope(observation):
...
>>> observation
{'my_observer:x': 1}
```

There are also a global API to add values:

```
>>> observation = {}
>>> with report_scope(observation):
... report({'x': 1}, observer)
...
>>> observation
{'my_observer:x': 1}
```

The most important application of Reporter is to report observed values from each link or chain in the training and validation procedures. *Trainer* and some extensions prepare their own Reporter object with the hierarchy of the target link registered as observers. We can use *report()* function inside any links and chains to report the observed values (e.g., training loss, accuracy, activation statistics, etc.).

Variables observation - Dictionary of observed values.

\_\_\_enter\_\_()

Makes this reporter object current.

\_\_exit\_\_\_(*exc\_type*, *exc\_value*, *traceback*) Recovers the previous reporter object to the current.

#### add\_observer(name, observer)

Registers an observer of values.

Observer defines a scope of names for observed values. Values observed with the observer are registered with names prefixed by the observer name.

#### Parameters

- **name** (*str*) Name of the observer.
- **observer** The observer object. Note that the reporter distinguishes the observers by their object ids (i.e., id(owner)), rather than the object equality.

## add\_observers (prefix, observers)

Registers multiple observers at once.

This is a convenient method to register multiple objects at once.

- **prefix** (*str*) Prefix of each name of observers.
- **observers** Iterator of name and observer pairs.

#### report (values, observer=None)

Reports observed values.

The values are written with the key, prefixed by the name of the observer object if given.

#### **Parameters**

- **values** (*dict*) Dictionary of observed values.
- **observer** Observer object. Its object ID is used to retrieve the observer name, which is used as the prefix of the registration name of the observed value.

#### scope (\*args, \*\*kwds)

Creates a scope to report observed values to observation.

This is a context manager to be passed to with statements. In this scope, the observation dictionary is changed to the given one.

It also makes this reporter object current.

**Parameters observation** (*dict*) – Observation dictionary. All observations reported inside of the with statement are written to this dictionary.

## chainer.get\_current\_reporter()

Returns the current reporter object.

## chainer.report (values, observer=None)

Reports observed values with the current reporter object.

Any reporter object can be set current by the with statement. This function calls the Report.report() method of the current reporter. If no reporter object is current, this function does nothing.

## Example

The most typical example is a use within links and chains. Suppose that a link is registered to the current reporter as an observer (for example, the target link of the optimizer is automatically registered to the reporter of the *Trainer*). We can report some values from the link as follows:

```
class MyRegressor(chainer.Chain):
    def __init__(self, predictor):
        super(MyRegressor, self).__init__(predictor=predictor)
    def __call__(self, x, y):
        # This chain just computes the mean absolute and squared
        # errors between the prediction and y.
        pred = self.predictor(x)
        abs_error = F.sum(F.abs(pred - y)) / len(x.data)
        loss = F.mean_squared_error(pred, y)
        # Report the mean absolute and squared errors.
        report({'abs_error': abs_error, 'squared_error': loss}, self)
        return loss
```

If the link is named 'main' in the hierarchy (which is the default name of the target link in the *StandardUpdater*), these reported values are named 'main/abs\_error' and 'main/ squared\_error'. If these values are reported inside the Evaluator extension, 'validation/' is added at the head of the link name, thus the item names are changed to 'validation/main/abs\_error' and 'validation/main/squared\_error' ('validation' is the default name of the Evaluator extension).

#### Parameters

- **values** (*dict*) Dictionary of observed values.
- **observer** Observer object. Its object ID is used to retrieve the observer name, which is used as the prefix of the registration name of the observed value.

#### chainer.report\_scope(\*args, \*\*kwds)

Returns a report scope with the current reporter.

This is equivalent to get\_current\_reporter().scope(observation), except that it does not make the reporter current redundantly.

#### Summary and DictSummary

#### class chainer. Summary

Online summarization of a sequence of scalars.

Summary computes the statistics of given scalars online.

#### add (value)

Adds a scalar value.

**Parameters value** – Scalar value to accumulate. It is either a NumPy scalar or a zerodimensional array (on CPU or GPU).

## compute\_mean()

Computes the mean.

## make\_statistics()

Computes and returns the mean and standard deviation values.

Returns Mean and standard deviation values.

#### Return type tuple

## class chainer.DictSummary

Online summarization of a sequence of dictionaries.

DictSummary computes the statistics of a given set of scalars online. It only computes the statistics for scalar values and variables of scalar values in the dictionaries.

#### add(d)

Adds a dictionary of scalars.

**Parameters d** (*dict*) – Dictionary of scalars to accumulate. Only elements of scalars, zerodimensional arrays, and variables of zero-dimensional arrays are accumulated.

## compute\_mean()

Creates a dictionary of mean values.

It returns a single dictionary that holds a mean value for each entry added to the summary.

Returns Dictionary of mean values.

Return type dict

#### make\_statistics()

Creates a dictionary of statistics.

It returns a single dictionary that holds mean and standard deviation values for every entry added to the summary. For an entry of name 'key', these values are added to the dictionary by names 'key' and 'key.std', respectively.

**Returns** Dictionary of statistics of all entries.

Return type dict

## **Experimental feature annotation**

chainer.utils.experimental(api\_name)

Declares that user is using an experimental feature.

The developer of an API can mark it as *experimental* by calling this function. When users call experimental APIs, FutureWarning is issued. The presentation of FutureWarning is disabled by setting chainer. disable\_experimental\_warning to True, which is False by default.

The basic usage is to call it in the function or method we want to mark as experimental along with the API name.

```
from chainer import utils
def f(x):
    utils.experimental('chainer.foo.bar.f')
    # concrete implementation of f follows
f(1)
```

... FutureWarning: chainer.foo.bar.f is experimental. The interface can change in...

We can also make a whole class experimental. In that case, we should call this function in its <u>\_\_\_\_\_\_init\_\_\_</u> method.

```
class C():
    def __init__(self):
        utils.experimental('chainer.foo.C')
C()
```

If we want to mark <u>\_\_\_\_\_\_</u> method only, rather than class itself, it is recommended that we explicitly feed its API name.

```
class D():
    def __init__(self):
        utils.experimental('D.__init__')
D()
```

```
... FutureWarning: D.__init__ is experimental. The interface can change in the_

→future. ...
```

Currently, we do not have any sophisticated way to mark some usage of non-experimental function as experimental. But we can support such usage by explicitly branching it.

```
def g(x, experimental_arg=None):
    if experimental_arg is not None:
        utils.experimental('experimental_arg of chainer.foo.g')
```

**Parameters** api\_name (*str*) – The name of an API marked as experimental.

# **Assertion and Testing**

Chainer provides some facilities to make debugging easy.

# Type checking utilities

Function uses a systematic type checking of the *chainer.utils.type\_check* module. It enables users to easily find bugs of forward and backward implementations. You can find examples of type checking in some function implementations.

class chainer.utils.type\_check.Expr (priority)

Abstract syntax tree of an expression.

It represents an abstract syntax tree, and isn't a value. You can get its actual value with eval() function, and get syntax representation with the \_\_str\_\_() method. Each comparison operator (e.g. ==) generates a new *Expr* object which represents the result of comparison between two expressions.

## Example

Let x and y be instances of *Expr*, then

```
>>> x = Variable(1, 'x')
>>> y = Variable(1, 'y')
>>> c = (x == y)
```

is also an instance of *Expr*. To evaluate and get its value, call *eval()* method:

```
>>> c.eval()
True
```

Call str function to get a representation of the original equation:

```
>>> str(c)
'x == y'
```

You can actually compare an expression with a value:

```
>>> (x == 1).eval()
True
```

Note that you can't use boolean operators such as and, as they try to cast expressions to boolean values:

## eval()

Evaluates the tree to get actual value.

Behavior of this function depends on an implementation class. For example, a binary operator + calls the  $\__add\__$  function with the two results of eval() function.

```
chainer.utils.type_check.expect(*bool_exprs)
```

Evaluates and tests all given expressions.

This function evaluates given boolean expressions in order. When at least one expression is evaluated as False, that means the given condition is not satisfied. You can check conditions with this function.

**Parameters bool\_exprs** (tuple of Bool expressions) – Bool expressions you want to evaluate.

class chainer.utils.type\_check.TypeInfo(shape, dtype)
 Type information of an input/gradient array.

It contains type information of an array, such as the shape of array and the number of dimensions. This information is independent of CPU or GPU array.

class chainer.utils.type\_check.TypeInfoTuple

Type information of input/gradient tuples.

It is a sub-class of tuple containing *TypeInfo*. The i-th element of this object contains type information of the i-th input/gradient data. As each element is *Expr*, you can easily check its validity.

size()

Returns an expression representing its length.

**Returns** An expression object representing length of the tuple.

Return type Expr

# Gradient checking utilities

Most function implementations are numerically tested by *gradient checking*. This method computes numerical gradients of forward routines and compares their results with the corresponding backward routines. It enables us to make the source of issues clear when we hit an error of gradient computations. The *chainer.gradient\_check* module makes it easy to implement the gradient checking.

chainer.gradient\_check.check\_backward(func, x\_data, y\_grad, params=(), eps=0.001, atol=1e-05, rtol=0.0001, no\_grads=None, dtype=None)

Test backward procedure of a given function.

This function automatically check backward-process of given function. For example, when you have a *Function* class MyFunc, that gets two arguments and returns one value, you can make its test like this:

```
>> def test_my_func(self):
>> func = MyFunc()
>> x1_data = xp.array(...)
>> x2_data = xp.array(...)
>> gy_data = xp.array(...)
>> check_backward(func, (x1_data, x2_data), gy_data)
```

This method creates *Variable* objects with x\_data and calls func with the *Variable* s to get its result as *Variable*. Then, it sets y\_grad array to grad attribute of the result and calls backward method to get gradients of the inputs. To check correctness of the gradients, the function calls  $numerical_grad()$  to calculate numerically the gradients and compares the types of gradients with *chainer.testing*. *assert\_allclose()*. If input objects (x1\_data or/and x2\_data in this example) represent integer variables, their gradients are ignored.

You can simplify a test when MyFunc gets only one argument:

>> check\_backward(func, x1\_data, gy\_data)

If MyFunc is a loss function which returns a zero-dimensional array, pass None to gy\_data. In this case, it sets 1 to grad attribute of the result:

>> check\_backward(my\_loss\_func, (x1\_data, x2\_data), None)

If MyFunc returns multiple outputs, pass all gradients for outputs as a tuple:

```
>> gy1_data = xp.array(...)
>> gy2_data = xp.array(...)
>> check_backward(func, x1_data, (gy1_data, gy2_data))
```

You can also test a *Link*. To check gradients of parameters of the link, set a tuple of the parameters to params arguments:

```
>> check_backward(my_link, (x1_data, x2_data), gy_data,
>> (my_link.W, my_link.b))
```

Note that params are not ndarray s, but Variables s.

Function objects are acceptable as func argument:

```
>> check_backward(lambda x1, x2: f(x1, x2),
>> (x1_data, x2_data), gy_data)
```

**Note:** func is called many times to get numerical gradients for all inputs. This function doesn't work correctly when func behaves randomly as it gets different gradients.

- **func** (*callable*) A function which gets *Variable* s and returns *Variable* s. func must returns a tuple of *Variable* s or one *Variable*. You can use *Function* object, *Link* object or a function satisfying the condition.
- **x\_data** (ndarray or tuple of ndarrays) A set of ndarrays to be passed to func. If x\_data is one ndarray object, it is treated as (x\_data,).
- **y\_grad** (ndarray or tuple of ndarrays or None) A set of ndarray s representing gradients of return-values of func. If y\_grad is one ndarray object, it is treated as (y\_grad,). If func is a loss-function, y\_grad should be set to None.
- **params** (Variable or tuple of ~chainder.Variable) A set of Variable s whose gradients are checked. When func is a Link object, set its parameters as params. If params is one Variable object, it is treated as (params,).
- **eps** (float) Epsilon value to be passed to numerical\_grad().
- **atol** (float) Absolute tolerance to be passed to chainer.testing. assert\_allclose().
- **rtol** (float) Relative tolerance to be passed to chainer.testing. assert\_allclose().
- no\_grads (list of bool) Flag to skip variable for gradient assertion. It should be same length as x\_data.

• dtype (*dtype*) - x\_data and y\_grad are casted to this dtype when calculating numerical gradients. Only float types and None are allowed.

See: numerical\_grad()

chainer.gradient\_check.numerical\_grad (*f*, *inputs*, *grad\_outputs*, *eps=0.001*) Computes numerical gradient by finite differences.

This function is used to implement gradient check. For usage example, see unit tests of *chainer*. *functions*.

## Parameters

- **f** (function) Python function with no arguments that runs forward computation and returns the result.
- **inputs** (*tuple of arrays*) Tuple of arrays that should be treated as inputs. Each element of them is slightly modified to realize numerical gradient by finite differences.
- **grad\_outputs** (*tuple of arrays*) Tuple of arrays that are treated as output gradients.
- **eps** (*float*) Epsilon value of finite differences.

Returns Numerical gradient arrays corresponding to inputs.

Return type tuple

# **Standard Assertions**

The assertions have same names as NumPy's ones. The difference from NumPy is that they can accept both numpy. ndarray and *cupy.ndarray*.

chainer.testing.**assert\_allclose**(*x*, *y*, *atol=1e-05*, *rtol=0.0001*, *verbose=True*) Asserts if some corresponding element of x and y differs too much.

This function can handle both CPU and GPU arrays simultaneously.

#### Parameters

- **x** Left-hand-side array.
- **y** Right-hand-side array.
- **atol** (*float*) Absolute tolerance.
- **rtol** (*float*) Relative tolerance.
- **verbose** (*bool*) If True, it outputs verbose messages on error.

# Function testing utilities

Chainer provides some utilities for testing its functions.

chainer.testing.unary\_math\_function\_unittest(func, func\_expected=None, label\_expected=None, make\_data=None)

Decorator for testing unary mathematical Chainer functions.

This decorator makes test classes test unary mathematical Chainer functions. Tested are forward and backward computations on CPU and GPU across parameterized shape and dtype.

- func (Function) Chainer function to be tested by the decorated test class.
- **func\_expected** Function used to provide expected values for testing forward computation. If not given, a corresponsing numpy function for func is implicitly picked up by its class name.
- **label\_expected** (*string*) String used to test labels of Chainer functions. If not given, the class name of func lowered is implicitly used.
- make\_data Function to customize input and gradient data used in the tests. It takes shape and dtype as its arguments, and returns a tuple of input and gradient data. By default, uniform destribution ranged [-1, 1] is used for both.

The decorated test class tests forward and backward computations on CPU and GPU across the following parameterize() ed parameters:

•shape: rank of zero, and rank of more than zero

•dtype: numpy.float16, numpy.float32 and numpy.float64

Additionally, it tests the label of the Chainer function.

Chainer functions tested by the test class decorated with the decorator should have the following properties:

•Unary, taking one parameter and returning one value

•dtype of input and output are the same

•Elementwise operation for the supplied ndarray

### Example

The following code defines a test class that tests *sin()* Chainer function, which takes a parameter with dtype of float and returns a value with the same dtype.

```
>>> import unittest
>>> from chainer import testing
>>> from chainer import functions as F
>>>
>>> @testing.unary_math_function_unittest(F.Sin())
... class TestSin(unittest.TestCase):
... pass
```

Because the test methods are implicitly injected to TestSin class by the decorator, it is enough to place pass in the class definition.

Now the test is run with nose module.

```
>>> import nose
>>> nose.run(
... defaultTest=__name__, argv=['', '-a', '!gpu'], exit=False)
True
```

To customize test data, make\_data optional parameter can be used. The following is an example of testing sqrt Chainer function, which is tested in positive value domain here instead of the default input.

```
...
>>> @testing.unary_math_function_unittest(F.Sqrt(),
... make_data=make_data)
... class TestSqrt(unittest.TestCase):
... pass
...
>>> nose.run(
... defaultTest=__name__, argv=['', '-a', '!gpu'], exit=False)
True
```

make\_data function which returns input and gradient data generated in proper value domains with given shape and dtype parameters is defined, then passed to the decorator's make\_data parameter.

# **Standard Function implementations**

Chainer provides basic *Function* implementations in the *chainer.functions* package. Most of them are wrapped by plain Python functions, which users should use.

**Note:** As of v1.5, the concept of parameterized functions are gone, and they are replaced by corresponding *Link* implementations. They are still put in the *functions* namespace for backward compatibility, though it is strongly recommended to use them via the *chainer.links* package.

# **Activation functions**

## clipped\_relu

```
chainer.functions.clipped_relu(x, z=20.0)
Clipped Rectifier Unit function.
```

For a clipping value z(>0), it computes

#### **Parameters**

- **x** (Variable or numpy.ndarray or cupy.ndarray) Input variable. A  $(s_1, s_2, ..., s_n)$ -shaped float array.
- z (float) Clipping value. (default = 20.0)

**Returns** Output variable. A  $(s_1, s_2, ..., s_n)$ -shaped float array.

Return type Variable

Example

```
>>> x = np.random.uniform(-100, 100, (10, 20)).astype('f')
>>> z = 10.0
>>> np.any(x < 0)
True
>>> np.any(x > z)
```

```
True
>>> y = F.clipped_relu(x, z=z)
>>> np.any(y.data < 0)
False
>>> np.any(y.data > z)
False
```

## crelu

chainer.functions.**crelu**(*x*, *axis=1*) Concatenated Rectified Linear Unit function.

This function is expressed as follows

$$f(x) = (\max(0, x), \max(0, -x)).$$

Here, two output values are concatenated along an axis.

See: https://arxiv.org/abs/1603.05201

## Parameters

- **x** (Variable or numpy.ndarray or cupy.ndarray) Input variable. A  $(s_1, s_2, ..., s_N)$ -shaped float array.
- **axis** (*int*) Axis that the output values are concatenated along. Default is 1.

**Returns** Output variable of concatenated array. If the axis is 1, A  $(s_1, s_2 \times 2, ..., s_N)$ -shaped float array.

**Return type** *Variable* 

#### Example

```
>>> x = np.array([[-1, 0], [2, -3]], 'f')
>>> x
array([[-1., 0.],
       [ 2., -3.]], dtype=float32)
>>> y = F.crelu(x, axis=1)
>>> y.data
array([[ 0., 0., 1., 0.],
       [ 2., 0., 0., 3.]], dtype=float32)
```

## elu

```
chainer.functions.elu(x, alpha=1.0)
Exponential Linear Unit function.
```

For a parameter  $\alpha$ , it is expressed as

$$f(x) = \begin{cases} x & \text{if } x \ge 0\\ \alpha(\exp(x) - 1) & \text{if } x < 0, \end{cases}$$

See: https://arxiv.org/abs/1511.07289

## Parameters

- **x** (Variable or numpy.ndarray or cupy.ndarray) Input variable. A  $(s_1, s_2, ..., s_N)$ -shaped float array.
- alpha (float) Parameter  $\alpha$ . Default is 1.0.

**Returns** Output variable. A  $(s_1, s_2, ..., s_N)$ -shaped float array.

Return type Variable

#### Example

```
>>> x = np.array([[-1, 0], [2, -3]], 'f')
>>> x
array([[-1., 0.],
       [ 2., -3.]], dtype=float32)
>>> y = F.elu(x, alpha=1.)
>>> y.data
array([[-0.63212055, 0. ],
       [ 2. , -0.95021296]], dtype=float32)
```

## hard\_sigmoid

chainer.functions.hard\_sigmoid(x) Element-wise hard-sigmoid function.

This function is defined as

$$f(x) = \begin{cases} 0 & \text{if } x < -2.5\\ 0.2x + 0.5 & \text{if } -2.5 < x < 2.5\\ 1 & \text{if } 2.5 < x. \end{cases}$$

**Parameters x** (*Variable* or numpy.ndarray or *cupy.ndarray*) – Input variable. A  $(s_1, s_2, ..., s_N)$ -shaped float array.

**Returns** Output variable. A  $(s_1, s_2, ..., s_N)$ -shaped float array.

**Return type** Variable

## Example

It maps the input values into the range of [0, 1].

```
>>> x = np.array([-2.6, -1, 0, 1, 2.6])
>>> x
array([-2.6, -1. , 0. , 1. , 2.6])
>>> F.hard_sigmoid(x).data
array([ 0. , 0.3, 0.5, 0.7, 1. ])
```

## leaky\_relu

chainer.functions.leaky\_relu(x, slope=0.2)
 Leaky Rectified Linear Unit function.

This function is expressed as

$$f(x) = \max(x, ax),$$

where a is a configurable slope value.

### **Parameters**

- **x** (Variable or numpy.ndarray or cupy.ndarray) Input variable. A  $(s_1, s_2, ..., s_N)$ -shaped float array.
- **slope** (*float*) Slope value *a*.

**Returns** Output variable. A  $(s_1, s_2, ..., s_N)$ -shaped float array.

**Return type** Variable

#### Example

```
>>> x = np.array([[-1, 0], [2, -3], [-2, 1]], 'f')
>>> x
array([[-1., 0.],
       [ 2., -3.],
       [-2., 1.]], dtype=float32)
>>> F.leaky_relu(x, slope=0.2).data
array([[-0.2 , 0. ],
       [ 2. , -0.60000002],
       [-0.40000001, 1. ]], dtype=float32)
```

## log\_softmax

chainer.functions.log\_softmax(x, use\_cudnn=True)

Channel-wise log-softmax function.

This function computes its logarithm of softmax along the second axis. Let  $c = (c_1, c_2, ..., c_D)$  be the slice of x along with the second axis. For each slice c, it computes the logarithm of the function f(c) defined as

$$f(c) = \frac{\exp(c)}{\sum_{d} \exp(c_d)}$$

This method is theoretically equivalent to log(softmax(x)) but is more stable.

Note: log(softmax(x)) may cause underflow when x is too small, because softmax(x) may returns 0.  $log_softmax$  method is more stable.

#### Parameters

- **x** (Variable or numpy.ndarray or cupy.ndarray) Input variable. A *n*-dimensional  $(n \ge 2)$  float array.
- **use\_cudnn** (*bool*) If True and cuDNN is enabled, then this function uses cuDNN as the core implementation.

**Returns** Output variable. A *n*-dimensional  $(n \ge 2)$  float array, which is the same shape with x.

**Return type** Variable

## See also:

softmax()

#### Example

```
>>> x = np.array([[0, 1, 2], [0, 2, 4]], 'f')
>>> x
array([[ 0., 1., 2.],
       [ 0., 2., 4.]], dtype=float32)
>>> F.log_softmax(x).data
array([[-2.40760589, -1.40760589, -0.40760589],
       [-4.14293146, -2.14293146, -0.14293146]], dtype=float32)
>>> np.allclose(F.log_softmax(x).data, F.log(F.softmax(x)).data)
True
```

## lstm

chainer.functions.lstm(c\_prev, x)

Long Short-Term Memory units as an activation function.

This function implements LSTM units with forget gates. Let the previous cell state  $c_prev$  and the input array x.

First, the input array x is split into four arrays a, i, f, o of the same shapes along the second axis. It means that x 's second axis must have 4 times the c\_prev 's second axis.

The split input arrays are corresponding to:

- $\bullet a$ : sources of cell input
- •i : sources of input gate
- •f : sources of forget gate
- •*o* : sources of output gate

Second, it computes the updated cell state c and the outgoing signal h as:

$$c = \tanh(a)\sigma(i) + c_{\text{prev}}\sigma(f),$$
  
$$h = \tanh(c)\sigma(o),$$

where  $\sigma$  is the elementwise sigmoid function. These are returned as a tuple of two variables.

This function supports variable length inputs. The mini-batch size of the current input must be equal to or smaller than that of the previous one. When mini-batch size of x is smaller than that of c, this function only updates c[0:len(x)] and doesn't change the rest of c, c[len(x):]. So, please sort input sequences in descending order of lengths before applying the function.

- **c\_prev** (*Variable* or numpy.ndarray or *cupy.ndarray*) Variable that holds the previous cell state. The cell state should be a zero array or the output of the previous call of LSTM.
- **x** (*Variable* or numpy.ndarray or *cupy.ndarray*) Variable that holds the sources of cell input, input gate, forget gate and output gate. It must have the second dimension whose size is four times of that of the cell state.

**Returns** Two *Variable* objects c and h. c is the updated cell state. h indicates the outgoing signal.

#### Return type tuple

See the original paper proposing LSTM with forget gates: Long Short-Term Memory in Recurrent Neural Networks.

#### See also:

LSTM

#### Example

Assuming y is the current incoming signal, c is the previous cell state, and h is the previous outgoing signal from an lstm function. Each of y, c and h has n\_units channels. Most typical preparation of x is:

```
>>> n_units = 100
>>> y = chainer.Variable(np.zeros((1, n_units), 'f'))
>>> h = chainer.Variable(np.zeros((1, n_units), 'f'))
>>> c = chainer.Variable(np.zeros((1, n_units), 'f'))
>>> model = chainer.Chain(w=L.Linear(n_units, 4 * n_units),
... v=L.Linear(n_units, 4 * n_units),
>>> x = model.w(y) + model.v(h)
>>> c, h = F.lstm(c, x)
```

It corresponds to calculate the input array x, or the input sources a, i, f, o, from the current incoming signal y and the previous outgoing signal h. Different parameters are used for different kind of input sources.

Note: We use the naming rule below.

- •incoming signal The formal input of the formulation of LSTM (e.g. in NLP, word vector or output of lower RNN layer). The input of *chainer.links.LSTM* is the *incoming signal*.
- •input array The array which is linear transformed from *incoming signal* and the previous outgoing signal. The *input array* contains four sources, the sources of cell input, input gate, forget gate and output gate. The input of chainer.functions.LSTM is the *input array*.

#### maxout

```
chainer.functions.maxout (x, pool_size, axis=1)
    Maxout activation function.
```

It accepts an input tensor x, reshapes the axis dimension (say the size being M \* pool\_size) into two dimensions (M, pool\_size), and takes maximum along the axis dimension.

- ★ (Variable or numpy.ndarray or cupy.ndarray) Input variable. A n-dimensional (n ≥ axis) float array. In general, its first dimension is assumed to be the minibatch dimension. The other dimensions are treated as one concatenated dimension.
- **pool\_size** (*int*) The size used for downsampling of pooling layer.
- **axis** (*int*) The axis dimension to be reshaped. The size of axis dimension should be M \* pool\_size.

**Returns** Output variable. The shape of the output is same as x except that axis dimension is transformed from M \* pool\_size to M.

Return type Variable

See also:

Maxout

#### Example

Typically, x is the output of a linear layer or a convolution layer. The following is the example where we use *maxout* () in combination with a Linear link.

```
>>> in_size, out_size, pool_size = 10, 10, 10
>>> bias = np.arange(out_size * pool_size).astype('f')
>>> l = L.Linear(in_size, out_size * pool_size, initial_bias=bias)
>>> x = np.zeros((1, in_size), 'f') # prepare data
>>> x = l(x)
>>> y = F.maxout(x, pool_size)
>>> x.shape
(1, 100)
>>> y.shape
(1, 10)
>>> x.reshape((out_size, pool_size)).data
               1., 2., 3., 4., 5.,
                                                      7.,
                                                            8.,
array([[ 0.,
                                                6.,
                                                                  9.1,
       [ 10.,
               11.,
                     12.,
                           13.,
                                 14.,
                                       15.,
                                              16.,
                                                    17.,
                                                           18.,
                                                                 19.1,
       [ 20.,
               21., 22., 23., 24., 25., 26.,
                                                     27.,
                                                           28.,
                                                                 29.1,
       ſ 30.,
               31.,
                     32.,
                           33.,
                                  34.,
                                        35.,
                                              36.,
                                                     37.,
                                                           38.,
                                                                 39.1.
       [ 40.,
               41.,
                     42.,
                           43.,
                                 44.,
                                        45.,
                                              46.,
                                                     47.,
                                                           48.,
                                                                 49.1,
                     52.,
                                  54.,
                                        55.,
                                                     57.,
               51.,
                                              56.,
                                                           58.,
                                                                 59.],
       [ 50.,
                           53.,
               61.,
                     62.,
                           63.,
                                  64.,
                                        65.,
                                              66.,
                                                     67.,
                                                           68.,
       [ 60.,
                                                                 69.1,
                     72.,
                            73.,
                                  74.,
                                        75.,
                                              76.,
                                                     77.,
       [ 70.,
               71.,
                                                           78.,
                                                                 79.1,
                                                           88.,
       [ 80.,
               81.,
                     82.,
                           83.,
                                  84.,
                                        85.,
                                              86.,
                                                     87.,
                                                                 89.1.
       [ 90.,
               91.,
                     92.,
                           93.,
                                  94.,
                                        95.,
                                              96.,
                                                     97.,
                                                           98.,
                                                                 99.]],
\rightarrow dtype=float32)
>>> y.data
array([[ 9., 19., 29., 39., 49., 59., 69., 79., 89., 99.]],
\rightarrow dtype=float32)
```

## prelu

chainer.functions.**prelu**(x, W)

Parametric ReLU function.

It accepts two arguments: an input x and a weight array W and computes the output as  $PReLU(x) = \max(x, W * x)$ , where \* is an elementwise multiplication for each sample in the batch.

When the PReLU function is combined with two-dimensional convolution, the elements of parameter *a* are typically shared across the same filter of different pixels. In order to support such usage, this function supports the shape of parameter array that indicates leading dimensions of input arrays except the batch dimension.

For example W has the shape of (2, 3, 4), x must have the shape of (B, 2, 3, 4, S1, ..., SN) where B is batch size and the number of trailing S's is arbitrary non-negative integer.

- **x** (Variable) Input variable. Its first argument is assumed to be the minibatch dimension.
- W (Variable) Weight variable.

Returns Output variable

Return type Variable

#### See also:

PReLU

## relu

chainer.functions.relu(x, use\_cudnn=True)
 Rectified Linear Unit function.

$$f(x) = \max(0, x).$$

#### **Parameters**

- **x** (Variable or numpy.ndarray or cupy.ndarray) Input variable. A  $(s_1, s_2, ..., s_N)$ -shaped float array.
- **use\_cudnn** (*bool*) If True and cuDNN is enabled, then this function uses cuDNN as the core implementation.

**Returns** Output variable. A  $(s_1, s_2, ..., s_N)$ -shaped float array.

Return type Variable

#### Example

```
>>> x = np.array([[-1, 0], [2, -3], [-2, 1]], 'f')
>>> np.any(x < 0)
True
>>> y = F.relu(x)
>>> np.any(y.data < 0)
False
>>> y.shape
(3, 2)
```

## sigmoid

```
chainer.functions.sigmoid(x, use_cudnn=True)
    Element-wise sigmoid logistic function.
```

$$f(x) = (1 + \exp(-x))^{-1}.$$

## Parameters

• **x** (Variable or numpy.ndarray or cupy.ndarray) – Input variable. A  $(s_1, s_2, ..., s_N)$ -shaped float array.

• **use\_cudnn** (*bool*) – If True and cuDNN is enabled, then this function uses cuDNN as the core implementation.

**Returns** Output variable. A  $(s_1, s_2, ..., s_N)$ -shaped float array.

Return type Variable

## Example

It maps the input values into the range of [0, 1].

```
>>> x = np.arange(-2, 3, 2).astype('f')
>>> x
array([-2., 0., 2.], dtype=float32)
>>> F.sigmoid(x).data
array([ 0.11920291, 0.5 , 0.88079709], dtype=float32)
```

#### slstm

```
chainer.functions.slstm(c_prev1, c_prev2, x1, x2)
S-LSTM units as an activation function.
```

This function implements S-LSTM unit. It is an extension of LSTM unit applied to tree structures. The function is applied to binary trees. Each node has two child nodes. It gets four arguments, previous cell states c\_prev1 and c\_prev2, and input arrays x1 and x2.

First both input arrays x1 and x2 are split into eight arrays  $a_1, i_1, f_1, o_1$ , and  $a_2, i_2, f_2, o_2$ . They have the same shape along the second axis. It means that x1 and x2 's second axis must have 4 times the length of c\_prev1 and c\_prev2.

The split input arrays are corresponding to:

- • $a_i$  : sources of cell input
- • $i_i$ : sources of input gate
- • $f_i$  : sources of forget gate
- • $o_i$  : sources of output gate

It computes the updated cell state c and the outgoing signal h as:

$$c = \tanh(a_1 + a_2)\sigma(i_1 + i_2) + c_{\text{prev}1}\sigma(f_1) + c_{\text{prev}2}\sigma(f_2),$$
  
$$h = \tanh(c)\sigma(o_1 + o_2),$$

where  $\sigma$  is the elementwise sigmoid function. The function returns c and h as a tuple.

- **c\_prev1** (*Variable* or numpy.ndarray or *cupy.ndarray*) Variable that holds the previous cell state of the first child node. The cell state should be a zero array or the output of the previous call of LSTM.
- **c\_prev2** (*Variable* or numpy.ndarray or *cupy.ndarray*) Variable that holds the previous cell state of the second child node.
- **x1** (*Variable* or numpy.ndarray or *cupy.ndarray*) Variable that holds the sources of cell input, input gate, forget gate and output gate from the first child node. It must have the second dimension whose size is four times of that of the cell state.

• **x2** (*Variable* or numpy.ndarray or *cupy.ndarray*) – Variable that holds the input sources from the second child node.

Returns Two Variable objects c and h. c is the cell state. h indicates the outgoing signal.

Return type tuple

See detail in paper: Long Short-Term Memory Over Tree Structures.

#### Example

Assuming c1, c2 is the previous cell state of children, and h1, h2 is the previous outgoing signal from children. Each of c1, c2, h1 and h2 has n\_units channels. Most typical preparation of x1, x2 is:

```
>>> n_units = 100
>>> h1 = chainer.Variable(np.zeros((1, n_units), 'f'))
>>> h2 = chainer.Variable(np.zeros((1, n_units), 'f'))
>>> c1 = chainer.Variable(np.zeros((1, n_units), 'f'))
>>> c2 = chainer.Variable(np.zeros((1, n_units), 'f'))
>>> model1 = chainer.Chain(w=L.Linear(n_units, 4 * n_units),
... v=L.Linear(n_units, 4 * n_units))
>>> model2 = chainer.Chain(w=L.Linear(n_units, 4 * n_units),
... v=L.Linear(n_units, 4 * n_units),
... v=L.Linear(n_units, 4 * n_units))
>>> x1 = model1.w(c1) + model1.v(h1)
>>> x2 = model2.w(c2) + model2.v(h2)
>>> c, h = F.slstm(c1, c2, x1, x2)
```

It corresponds to calculate the input array  $\times 1$ , or the input sources  $a_1, i_1, f_1, o_1$  from the previous cell state of first child node c1, and the previous outgoing signal from first child node h1. Different parameters are used for different kind of input sources.

## softmax

chainer.functions.**softmax**(*x*, *use\_cudnn=True*, *axis=1*)

Softmax function.

This function computes its softmax along an axis. Let  $x = (x_1, x_2, \dots, x_d)^{\top}$  be the d dimensional index array and f(x) be the d dimensional input array. For each index x of the input array f(x), it computes the probability p(x) defined as  $p(x) = \frac{\exp(f(x))}{\sum_{x_2} \exp(f(x))}$ .

#### Parameters

- **x** (Variable) Input variable.
- **use\_cudnn** (*bool*) If True and cuDNN is enabled, then this function uses cuDNN as the core implementation.
- **axis** The axis along which the softmax is to be computed.

Returns Output variable.

Return type Variable

## softplus

```
chainer.functions.softplus (x, beta=1.0)
Element-wise softplus function.
```

The softplus function is the smooth approximation of ReLU.

$$f(x) = \frac{1}{\beta} \log(1 + \exp(\beta x)),$$

where  $\beta$  is a parameter. The function becomes curved and akin to ReLU as the  $\beta$  is increasing.

#### **Parameters**

- **x** (Variable or numpy.ndarray or cupy.ndarray) Input variable. A  $(s_1, s_2, ..., s_N)$ -shaped float array.
- **beta** (float) Parameter  $\beta$ .

**Returns** Output variable. A  $(s_1, s_2, ..., s_N)$ -shaped float array.

Return type Variable

# Example

```
>>> x = np.arange(-2, 3, 2).astype('f')
>>> x
array([-2., 0., 2.], dtype=float32)
>>> F.softplus(x, beta=1.0).data
array([ 0.126928 ,  0.69314718,  2.12692809], dtype=float32)
```

## tanh

chainer.functions.tanh(x, use\_cudnn=True)
 Elementwise hyperbolic tangent function.

 $f(x) = \tanh(x).$ 

### **Parameters**

- **x** (Variable or numpy.ndarray or cupy.ndarray) Input variable. A  $(s_1, s_2, ..., s_N)$ -shaped float array.
- **use\_cudnn** (*bool*) If True and cuDNN is enabled, then this function uses cuDNN as the core implementation.

**Returns** Output variable. A  $(s_1, s_2, ..., s_N)$ -shaped float array.

Return type Variable

```
>>> x = np.arange(-1, 4, 2).astype('f')
>>> x
array([-1., 1., 3.], dtype=float32)
>>> F.tanh(x).data
array([-0.76159418, 0.76159418, 0.99505478], dtype=float32)
```

# Array manipulations

# broadcast

chainer.functions.broadcast(\*args)

Broadcast given variables.

**Parameters args** (*Variable* or numpy.ndarray or *cupy.ndarray*) – Input variables to be broadcasted. Each dimension of the shapes of the input variables must have the same size.

Returns Variable or tuple of Variable objects which are broadcasted from given arguments.

**Return type** Variable

Example

```
>>> x = np.random.uniform(0, 1, (3, 2)).astype('f')
>>> y = F.broadcast(x)
>>> np.all(x == y.data)
True
>>> z = np.random.uniform(0, 1, (3, 2)).astype('f')
>>> y, w = F.broadcast(x, z)
>>> np.all(x == y.data) & np.all(z == w.data)
True
```

## broadcast\_to

chainer.functions.broadcast\_to(x, shape)

Broadcast a given variable to a given shape.

# Parameters

- **x** (*Variable* or numpy.ndarray or *cupy.ndarray*) Input variable be broadcasted. A  $(s_1, s_2, ..., s_N)$ -shaped float array.
- **shape** (*tuple*) Tuple of int of the shape of the output variable.

Returns Output variable broadcasted to the given shape.

**Return type** *Variable* 

# cast

chainer.functions.**cast** (*x*, *typ*) Cast an input variable to a given type.

# Parameters

- **x** (*Variable* or numpy.ndarray or *cupy.ndarray*) Input variable to be casted. A  $(s_1, s_2, ..., s_N)$ -shaped float array.
- typ (str of dtype or numpy.dtype) Typecode or data type to cast.

Returns Variable holding a casted array.

**Return type** *Variable* 

# Example

```
>>> x = np.arange(0, 3, dtype=np.float64)
>>> x.dtype
dtype('float64')
>>> y = F.cast(x, np.float32)
>>> y.dtype
dtype('float32')
>>> y = F.cast(x, 'float16')
>>> y.dtype
dtype('float16')
```

# concat

chainer.functions.concat(xs, axis=1)
Concatenates given variables along an axis.

#### **Parameters**

- **xs** (tuple of *Variable* or numpy.ndarray or *cupy.ndarray*) Input variables to be concatenated. The variables must have the same shape, except in the dimension corresponding to axis.
- **axis** (*int*) The axis along which the arrays will be joined. Default is 1.

Returns The concatenated variable.

```
Return type Variable
```

### copy

chainer.functions.copy(x, dst)

Copies the input variable onto the specified device.

This function copies the array of input variable onto the device specified by dst. When dst = -1, it copies the array onto the host memory. This function supports copies from host to host, from host to device, from device to device and from device to host.

#### **Parameters**

- **x** (Variable or numpy.ndarray or cupy.ndarray) Variable to be copied.
- **dst** (*int*) Target device specifier.

Returns Output variable.

Return type Variable

### Example

```
>>> import chainer.cuda as cuda
>>> x = np.random.uniform(-1, 1, (5, 10))
>>> cuda.get_device_from_array(x).id
-1
>>> y = F.copy(x, 0) # from host to device0
>>> cuda.get_device_from_array(y.data).id
0
>>> z = F.copy(y, -1) # from device0 to host
>>> cuda.get_device_from_array(z.data).id
-1
```

## depth2space

```
chainer.functions.depth2space(X, r)
```

Computes the depth2space transformation for subpixel calculations.

# Parameters

- **X** (*Variable* or numpy.ndarray or *cupy.ndarray*) Variable holding a 4d array of shape (batch, channel \* r \* r, dim1, dim2).
- **r** (*int*) the upscaling factor.

**Returns** A variable holding the upscaled array from interspersed depth layers. The shape is (batch, channel, dim1 \* r, dim2 \* r).

### **Return type** Variable

**Note:** This can be used to compute super-resolution transformations. See https://arxiv.org/abs/1609.05158 for details.

### See also:

space2depth()

## Example

```
>>> X = np.arange(24).reshape(1, 4, 2, 3).astype('f')
>>> X.shape
(1, 4, 2, 3)
>>> X
                       2.1,
array([[[ 0., 1.,
        [ 3.,
                 4.,
                       5.]],
                7., 8.],
        [[ 6.,
        [ 9., 10., 11.]],
       [[ 12., 13., 14.],
        [ 15., 16., 17.]],
        [[ 18., 19., 20.],
        [ 21., 22., 23.]]], dtype=float32)
>>> y = F.depth2space(X, 2)
>>> y.shape
(1, 1, 4, 6)
>>> y.data
          0.,
                 6.,
                       1.,
                            7.,
                                  2.,
                                        8.1,
array([[[
               18., 13.,
                           19., 14.,
        [ 12.,
                                       20.1,
                9.,
                      4., 10.,
                                 5.,
        [ 3.,
                                      11.],
        [ 15.,
                      16.,
                            22.,
                                 17.,
                                       23.]]]], dtype=float32)
                21.,
```

# dstack

chainer.functions.**dstack** (*xs*) Concatenate variables along third axis (depth wise).

- **Parameters xs** (list of *Variable* or numpy.ndarray or *cupy.ndarray*) Input variables to be concatenated. The variables must have the same ndim. When the variables have the third axis (i.e.  $ndim \ge 3$ ), the variables must have the same shape along all but the third axis. When the variables do not have the third axis(i.e. ndim < 3), the variables must have the same shape.
- **Returns** Output variable. When the input variables have the third axis (i.e.  $ndim \ge 3$ ), the shapes of inputs and output are the same along all but the third axis. The length of third axis is the sum of the lengths of inputs' third axis. When the shape of variables are (N1, N2) (i.e. ndim = 2), the shape of output is (N1, N2, 2). When the shape of variables are (N1, ) (i.e. ndim = 1), the shape of output is (1, N1, 2). When the shape of variables are () (i.e. ndim = 0), the shape of output is (1, 1, 2).

```
Return type Variable
```

```
>>> x1 = np.array((1, 2, 3))
>>> x1.shape
(3,)
>>> x^2 = np.array((2, 3, 4))
>>> x2.shape
(3,)
>>> y = F.dstack((x1, x2))
>>> y.shape
(1, 3, 2)
>>> y.data
array([[[1, 2],
        [2, 3],
        [3, 4]])
>>> x1 = np.arange(0, 6).reshape(3, 2)
>>> x1.shape
(3, 2)
>>> x1
array([[0, 1],
      [2, 3],
      [4, 5]])
>>> x2 = np.arange(6, 12).reshape(3, 2)
>>> x2.shape
(3, 2)
>>> x2
array([[ 6, 7],
      [8, 9],
       [10, 11]])
>>> y = F.dstack([x1, x2])
>>> y.shape
(3, 2, 2)
>>> y.data
array([[[ 0, 6],
       [ 1, 7]],
       [[2, 8],
       [3, 9]],
       [[ 4, 10],
       [ 5, 11]])
>>> x1 = np.arange(0, 12).reshape(3, 2, 2)
>>> x2 = np.arange(12, 18).reshape(3, 2, 1)
>>> y = F.dstack([x1, x2])
>>> y.shape
(3, 2, 3)
>>> y.data
array([[[ 0, 1, 12],
       [2, 3, 13]],
       [[ 4, 5, 14],
       [ 6, 7, 15]],
       [[ 8, 9, 16],
```

[10, 11, 17]])

# expand\_dims

chainer.functions.expand\_dims (x, axis)

Expands dimensions of an input variable without copy.

## Parameters

- **x** (*Variable* or numpy.ndarray or *cupy.ndarray*) Input variable.
- **axis** (int) Position where new axis is to be inserted. The axis parameter is acceptable when  $-ndim 1 \le axis \le ndim$ . (ndim is the dimension of input variables). When axis < 0, the result is the same with ndim + 1 |axis|.

**Returns** Variable that holds a expanded input. The ndim of output is one grater than that of x.

**Return type** Variable

### Example

```
>>> x = np.array([1, 2, 3])
>>> x.shape
(3,)
>>> y = F.expand_dims(x, axis=0)
>>> y.shape
(1, 3)
>>> y.data
array([[1, 2, 3]])
>>> y = F.expand_dims(x, axis=1)
>>> y.shape
(3, 1)
>>> y.data
array([[1],
       [2],
       [3]])
>>> y = F.expand_dims(x, axis=-2)
>>> y.shape
(1, 3)
>>> y.data
array([[1, 2, 3]])
```

# flatten

chainer.functions.**flatten** (*x*) Flatten a given array.

Parameters x (Varaiable) – Input variable.

Returns Output variable.

**Return type** Variable

# fliplr

chainer.functions.fliplr(*a*) Flip array in the left/right direction.

**Parameters xs** (Variable) – Input variable.

Returns Output variable.

Return type Variable

# flipud

chainer.functions.flipud(a) Flip array in the up/down direction.

Parameters **xs** (Variable) – Input variable.

Returns Output variable.

Return type Variable

# get\_item

chainer.functions.get\_item(x, slices)

Extract elements from array with specified shape, axes and offsets.

# Parameters

- **x** (Variable) A variable to be sliced.
- **slices** (*int*, *slice*, *Ellipsis*, *None*, *integer array-like*, *boolean array-like or tuple of them*) It is an integer, a slice, an ellipsis, a numpy.newaxis, an integer array-like, a boolean array-like or tuple of them.

### Returns

**Variable object** which contains sliced array of x.

# Return type Variable

Note: It only supports types that are supported by CUDA's atomicAdd when an integer array is included in slices. The supported types are numpy.float32, numpy.int32, numpy.uint32, numpy.uint64 and numpy.ulonglong.

Note: It does not support slices that contains multiple boolean arrays.

Note: See NumPy document for details of indexing.

# hstack

chainer.functions.hstack(xs)

Concatenate variables horizontally (column wise).

**Parameters xs** (list of *Variable* or numpy.ndarray or *cupy.ndarray*) – Input variables to be concatenated. The variables must have the same ndim. When the variables have the second axis (i.e.  $ndim \ge 2$ ), the variables must have the same shape along all but the second axis. When the variables do not have the second axis(i.e. ndim < 2), the variables need not to have the same shape.

**Returns** Output variable. When the input variables have the second axis (i.e.  $ndim \ge 2$ ), the shapes of inputs and output are the same along all but the second axis. The length of second axis is the sum of the lengths of inputs' second axis. When the variables do not have the second axis (i.e. ndim < 2), the shape of output is (N, ) (N is the sum of the input variables' size).

Return type Variable

#### Example

```
>>> x1 = np.array((1, 2, 3))
>>> x1.shape
(3,)
>>> x^2 = np.array((2, 3, 4))
>>> x2.shape
(3,)
>>> y = F.hstack((x1, x2))
>>> y.shape
(6,)
>>> y.data
array([1, 2, 3, 2, 3, 4])
>>> x1 = np.arange(0, 12).reshape(3, 4)
>>> x1.shape
(3, 4)
>>> x1
array([[ 0, 1, 2, 3],
       [4, 5, 6, 7],
       [8, 9, 10, 11]])
>>> x2 = np.arange(12, 18).reshape(3, 2)
>>> x2.shape
(3, 2)
>>> x2
array([[12, 13],
       [14, 15],
       [16, 17]])
>>> y = F.hstack([x1, x2])
>>> y.shape
(3, 6)
>>> y.data
            1, 2, 3, 12, 13],
array([[ 0,
            5, 6, 7, 14, 15],
       [ 4,
            9, 10, 11, 16, 17]])
       [ 8,
```

# im2col

chainer.functions.**im2col** (*x*, *ksize*, *stride=1*, *pad=0*, *cover\_all=False*, *dilate=1*) Extract patches from an image based on the filter.

This function rearranges patches of an image and put them in the channel dimension of the output.

Patches are extracted at positions shifted by multiples of stride from the first position -pad for each spatial axis. The right-most (or bottom-most) patches do not run over the padded spatial size.

Notation: here is a notation.

 $\bullet n$  is the batch size.

 $\bullet c$  is the number of the input channels.

 $\bullet h$  and w are the height and width of the input image, respectively.

• $k_H$  and  $k_W$  are the height and width of the filters, respectively.

• $s_Y$  and  $s_X$  are the strides of the filter.

• $p_H$  and  $p_W$  are the spatial padding sizes.

• $d_Y$  and  $d_X$  are the dilation factors of filter application.

The output size  $(h_O, w_O)$  is determined by the following equations when cover\_all = False:

$$h_O = (h + 2p_H - k_H - (k_H - 1) * (d_Y - 1))/s_Y + 1,$$
  
$$w_O = (w + 2p_W - k_W - (k_W - 1) * (d_X - 1))/s_X + 1.$$

When cover\_all = True, the output size is determined by the following equations:

$$h_O = (h + 2p_H - k_H - (k_H - 1) * (d_Y - 1) + s_Y - 1)/s_Y + 1,$$
  
$$w_O = (w + 2p_W - k_W - (k_W - 1) * (d_X - 1) + s_X - 1)/s_X + 1.$$

### **Parameters**

- **x** (Variable) Input variable of shape (n, c, h, w).
- **ksize** (*int or pair of ints*) **Size of filters** (a.k.a. kernels). ksize=k and ksize=(k, k) are equivalent.
- **stride** (*int or pair of ints*) Stride of filter applications. stride=s and stride=(s, s) are equivalent.
- **pad** (*int or pair of ints*) Spatial padding width for input arrays. pad=p and pad=(p, p) are equivalent.
- **cover\_all** (*bool*) If True, all spatial locations are rearranged into some output pixels. It may make the output size larger.
- dilate (*int* or *pair* of *ints*) Dilation factor of filter applications. dilate=d and dilate=(d, d) are equivalent.

**Returns** Output variable whose shape is  $(n, c \cdot k_H \cdot k_W, h_O, w_O)$ 

**Return type** Variable

### pad

chainer.functions.pad(x, pad\_width, mode, \*\*keywords)
Pad an input variable.

#### **Parameters**

- x (chainer. Variable or :class:numpy.ndarray or cupy.ndarray) Input data.
- **pad\_width** (*int or array-like*) Number of values padded to the edges of each axis.
- mode (str) Specifies how the function fills the periphery of the array. constant

Pads with a constant values.

• constant\_values (int or array-like) - The values are padded for each axis.

Returns Output variable.

**Return type** *Variable* 

# permutate

chainer.functions.**permutate**(*x*, *indices*, *axis*=0, *inv*=*False*) Permutates a given variable along an axis.

This function permutate x with given indices. That means y[i] = x[indices[i]] for all i. Note that this result is same as y = x.take(indices). indices must be a permutation of [0, 1, ..., len(x) - 1].

When inv is True, indices is treated as its inverse. That means y[indices[i]] = x[i].

### Parameters

- **x** (Variable) Variable to permutate.
- indices (Variable) Indices to extract from the variable.
- **axis** (*int*) Axis that the input array is permutate along.
- **inv** (*bool*) If True, indices is treated as its inverse.

Returns Output variable.

Return type Variable

#### reshape

chainer.functions.**reshape**(*x*, *shape*)

Reshapes an input variable without copy.

# Parameters

- **x** (Variable) Input variable.
- **shape** (tuple of ints) Target shape.

# Returns

Variable that holds a reshaped version of the input variable.

**Return type** Variable

# resize\_images

```
chainer.functions.resize_images(x, output_shape)
```

Resize images to the given shape.

This function resizes 2D data to output\_shape. Currently, only bilinear interpolation is supported as the sampling method.

Notatition: here is a notation for dimensionalities.

 $\bullet n$  is the batch size.

• $c_I$  is the number of the input channels.

• *h* and *w* are the height and width of the input image, respectively.

• $h_O$  and  $w_O$  are the height and width of the output image.

### Parameters

• **x** (Variable) – Input variable of shape  $(n, c_I, h, w)$ .

• **output\_shape** (*tuple*) – This is a tuple of length 2 whose values are (h\_0, w\_0). Note that the order of height and width is opposite of the one in OpenCV.

**Returns** Resized image whose shape is  $(n, c_I, h_O, w_O)$ .

**Return type** Variable

# rollaxis

chainer.functions.**rollaxis** (*x*, *axis*, *start=0*) Roll the axis backwards to the given position.

#### **Parameters**

- **x** (Variable) Input variable.
- **axis** (*int*) The axis to roll backwards.
- **start** (*int*) The place to which the axis is moved.

**Returns** Variable whose axis is rolled.

Return type Variable

### select\_item

```
chainer.functions.select_item(x, t)
```

Select elements stored in given indices.

This function returns t.choose (x.T), that means y[i] == x[i, t[i]] for all i.

### Parameters

- **x** (Variable) Variable storing arrays.
- t (Variable) Variable storing index numbers.

**Returns** Variable that holds t-th element of x.

Return type Variable

# separate

```
chainer.functions.separate(x, axis=0)
```

Separates an array along a given axis.

This function separates an array along a given axis. For example, shape of an array is (2, 3, 4). When it separates the array with axis=1, it returns three (2, 4) arrays.

This function is an inverse of chainer.functions.stack().

Parameters

- **x** (chainer.Variable) Variable to be separated.
- **axis** (*int*) Axis along which variables are separated.

Returns Output variables.

Return type tuple of chainer. Variable

# See also:

```
chainer.functions.stack()
```

# space2depth

```
chainer.functions.space2depth(X, r)
```

Computes the space2depth transformation for subpixel calculations.

### Parameters

- X (Variable or numpy.ndarray or cupy.ndarray) Variable holding a 4d array of shape (batch, channel, dim1 \* r, dim2 \* r).
- **r** (*int*) the downscaling factor.
- **Returns** A variable holding the downscaled layer array from subpixel array sampling. The shape is (batch, channel \* r \* r, dim1, dim2).

#### **Return type** Variable

**Note:** This can be used to compute inverse super-resolution transformations. See https://arxiv.org/abs/1609. 05158 for details.

### See also:

depth2space()

```
>>> X = np.arange(24).reshape(1, 1, 4, 6).astype('f')
>>> X.shape
(1, 1, 4, 6)
>>> X
array([[[ 0., 1., 2.,
                           3.,
                               4.,
                                     5.1,
               7.,
                    8.,
                          9., 10., 11.],
        [ 6.,
        [ 12., 13., 14., 15., 16., 17.],
        [ 18., 19., 20.,
                         21., 22., 23.]]]], dtype=float32)
>>> y = F.space2depth(X, 2)
>>> y.shape
(1, 4, 2, 3)
>>> y.data
array([[[ 0., 2., 4.],
        [ 12., 14., 16.]],
       [[ 1., 3., 5.],
       [ 13., 15., 17.]],
       [[ 6., 8., 10.],
       [ 18., 20., 22.]],
       [[ 7.,
               9., 11.],
        [ 19.,
               21., 23.]]]], dtype=float32)
```

# spatial\_transformer\_grid

chainer.functions.**spatial\_transformer\_grid**(*theta*, *output\_shape*, *use\_cudnn=True*) 2D Spatial Transformer grid.

This function generates coordinates of the points sampled from an image to perform warping described in Spatial Transformer Networks.

Given a coordinate in the warped image  $(x_i^t, y_i^t)$ , the point sampled from the source image  $(x_i^s, y_i^s)$  are calculated by the following equation.

$$\begin{pmatrix} x_i^s \\ y_i^s \end{pmatrix} = \begin{pmatrix} \theta_{11} & \theta_{12} & \theta_{13} \\ \theta_{21} & \theta_{22} & \theta_{23} \end{pmatrix} \begin{pmatrix} x_i^t \\ y_i^t \\ 1 \end{pmatrix}$$

Notatition: here is a notation for dimensionalities.

 $\bullet n$  is the batch size.

• $h_O$  and  $w_O$  are the height and the width of the output image.

### **Parameters**

- theta (Variable) An array of shape (n, 2, 3). This is a batch of  $2 \times 3$  matrix used for the warping described above.
- **output\_shape** (*tuple*) A tuple of 2 elements:  $h_O, w_O$ .
- **use\_cudnn** (*bool*) If True, then this function uses cuDNN if available. Note that, cuDNN supports SpatialTransformerGrid from version 5.0.0.
- **Returns** A variable of shape  $(n, 2, h_O, w_O)$ . In the 2nd dimension, the first element is the coordinate along the x axis, and the second element is the coordinate along the y axis. All the coordinates in the image are scaled to fit range [-1, 1]. This means that the coordinate (-1, -1) corresponds to the upper-left corner of the input image.

# Return type Variable

# spatial\_transformer\_sampler

```
chainer.functions.spatial_transformer_sampler(x, grid, use_cudnn=True) 2D Spatial Transformer sampler.
```

This is a differentiable image sampler. With a set of sampling points grid and an input feature map x, this produces a sampled output feature map.

This function currently only supports bilinear interpolation as a sampling kernel.

When coordinates in grid is outside range [-1, 1], values are sampled from a zero padded input image.

Notatition: here is a notation for dimensionalities.

 $\bullet n$  is the batch size.

• $c_I$  is the number of the input channels.

• *h* and *w* are the height and width of the input image, respectively.

• $h_O$  and  $w_O$  are the height and width of the output image.

See detail in the following paper: Spatial Transformer Networks.

### **Parameters**

- **x** (Variable) Input variable of shape  $(n, c_I, h, w)$ .
- **grid** (Variable) Coordinate variable of shape  $(n, 2, h_O, w_O)$ . Each coordinate defines the spatial location in the input where a sampling kernel is applied to get the value at a particular pixel in the output. grid[idx, :, i, j] corresponds to the coordinate that is used to sample the values for an output pixel at location (i, j).

In the second dimension, the first coordinate corresponds to the location along the horizontal axis, and the second coordinate corresponds to the location along the vertical axis.

The coordinate (-1, -1) corresponds to the upper-left corner of the input image.

• **use\_cudnn** (*bool*) – If True, then this function uses cuDNN if available. Note that, cuDNN supports SpatialTransformerSampler from version 5.0.0.

**Returns** Output feature map of shape  $(n, c_I, h_O, w_O)$ .

**Return type** Variable

### split\_axis

chainer.functions.**split\_axis**(*x*, *indices\_or\_sections*, *axis*, *force\_tuple=False*) Splits given variables along an axis.

#### Parameters

- **x** (tuple of Variables) Variables to be split.
- **indices\_or\_sections** (*int or 1-D array*) If this argument is an integer, N, the array will be divided into N equal arrays along axis. If it is a 1-D array of sorted integers, it indicates the positions where the array is split.
- **axis** (*int*) Axis that the input array is split along.
- **force\_tuple** (*bool*) If True, this method returns a tuple even when the number of outputs is one.

## Returns

Tuple of Variable objects if the number of outputs is more than 1 or Variable otherwise. When force\_tuple is True, returned value is always a tuple regardless of the number of outputs.

### Return type tuple or Variable

Note: This function raises ValueError if at least one of the outputs is split to zero-size (i.e. axis-th value of its shape is zero).

#### squeeze

chainer.functions.squeeze(x, axis=None)

Remove demensions of size one from the shape of a ndarray.

#### Parameters

- x (chainer. Variable or :class:numpy.ndarray or cupy.ndarray) Input data.
- **axis** (*None or int or tuple of ints*) A subset of the single-dimensional entries in the shape to remove. If None is supplied, all of them are removed. The dimension index starts at zero. If an axis with dimension greater than one is selected, an error is raised.

Returns Variable whose dimensions of size 1 are removed.

Return type Variable

# stack

chainer.functions.**stack**(*xs*, *axis=0*) Concatenate variables along a new axis.

# Parameters

- **xs** (list of *Variable* or numpy.ndarray or *cupy.ndarray*) Input variables to be concatenated. The variables must have the same shape.
- **axis** (int) The axis along which the arrays will be stacked. The axis parameter is acceptable when  $-ndim 1 \le axis \le ndim$ . (ndim is the dimension of input variables). When axis < 0, the result is the same with ndim + 1 |axis|.

**Returns** Output variable. Let x\_1, x\_2, ..., x\_n and y be the input variables and the output variable, y[:, ..., 0, ..., :] is x\_1, y[:, ..., 1, ..., :] is x\_2 and y[:, ..., n-1, ..., :] is x\_n (The indexed axis indicates the axis).

Return type Variable

```
>>> x1 = np.arange(0, 12).reshape(3, 4)
>>> x1.shape
(3, 4)
>>> x1
array([[ 0, 1, 2, 3],
      [4, 5, 6, 7],
      [8, 9, 10, 11]])
>>> x2 = np.arange(12, 24).reshape(3, 4)
>>> x2.shape
(3, 4)
>>> x2
array([[12, 13, 14, 15],
      [16, 17, 18, 19],
      [20, 21, 22, 23]])
>>> y = F.stack([x1, x2], axis=0)
>>> y.shape
(2, 3, 4)
>>> y.data
array([[[ 0, 1, 2, 3],
       [4, 5, 6, 7],
        [8, 9, 10, 11]],
       [[12, 13, 14, 15],
       [16, 17, 18, 19],
       [20, 21, 22, 23]])
>>> y = F.stack([x1, x2], axis=1)
>>> y.shape
(3, 2, 4)
>>> y.data
array([[[ 0, 1, 2, 3],
       [12, 13, 14, 15]],
```

```
[[4, 5, 6, 7],
        [16, 17, 18, 19]],
       [[ 8, 9, 10, 11],
       [20, 21, 22, 23]])
>>> y = F.stack([x1, x2], axis=2)
>>> y.shape
(3, 4, 2)
>>> y.data
array([[[ 0, 12],
        [ 1, 13],
        [ 2, 14],
        [ 3, 15]],
       [[ 4, 16],
       [ 5, 17],
        [ 6, 18],
        [ 7, 19]],
       [[ 8, 20],
        [ 9, 21],
        [10, 22],
        [11, 23]])
>>> y = F.stack([x1, x2], axis=-1)
>>> y.shape
(3, 4, 2)
```

# swapaxes

chainer.functions.**swapaxes** (*x*, *axis1*, *axis2*) Swap two axes of a variable.

# Parameters

- **x** (Variable) Input variable.
- **axis1** (*int*) The first axis to swap.
- **axis2** (*int*) The second axis to swap.

Returns Variable whose axes are swapped.

Return type Variable

# tile

chainer.functions.tile(x, reps)

Construct an array by tiling a given array.

# Parameters

- **x** (chainer.Variable or numpy.ndarray or cupy.ndarray) Input data.
- **reps** (*int* or *tuple* of *ints*) The number of times for each axis with which x is replicated.

**Returns** Variable tiled the given array.

### **Return type** Variable

#### transpose

chainer.functions.transpose (x, axes=None)
 Permute the dimensions of an input variable without copy.

# Parameters

- **x** (Variable) Input variable.
- **axes** (*tuple of ints*) By default, reverse the dimensions, otherwise permute the axes according to the values given.

Returns Variable whose axes are permuted.

**Return type** Variable

## transpose\_sequence

#### chainer.functions.transpose\_sequence(xs)

Transpose a list of Variables.

This function transposes a list of *Variable* s and returns a list of *Variable* s. For example a user gives [(0, 1, 2, 3), (4, 5), (6)], the function returns [(0, 4, 6), (1, 5), (2), (3)]. Note that a given list needs to be sorted by each length of *Variable*.

**Parameters xs** (list of ~chainer.Variable) – Variables to transpose.

Returns Transposed list.

Return type tuple or Variable

### vstack

```
chainer.functions.vstack(xs)
```

Concatenate variables vertically (row wise).

- **Parameters xs** (list of *Variable* or numpy.ndarray or *cupy.ndarray*) Input variables to be concatenated. The variables must have the same ndim. When the variables have the second axis (i.e.  $ndim \ge 2$ ), the variables must have the same shape along all but the first axis. When the variables do not have the second axis(i.e. ndim < 2), the variables must have the same shape along all but the first axis. When the variables do not have the second axis(i.e. ndim < 2), the variables must have the same shape.
- **Returns** Output variable. When the input variables have the second axis (i.e.  $ndim \ge 2$ ), the shapes of inputs and output are the same along all but the first axis. The length of first axis is the sum of the lengths of inputs' first axis. When the variables do not have the second axis (i.e. ndim < 2), the shape of output is (2, N) (N is the size of the input variable).

**Return type** Variable

```
>>> x1 = np.array((1, 2, 3))
>>> x1.shape
(3,)
>>> x2 = np.array((2, 3, 4))
>>> x2.shape
```

```
(3,)
>>> y = F.vstack((x1, x2))
>>> y.shape
(2, 3)
>>> y.data
array([[1, 2, 3],
      [2, 3, 4]])
>>> x1 = np.arange(0, 12).reshape(3, 4)
>>> x1.shape
(3, 4)
>>> x1
array([[ 0, 1, 2, 3],
       [4, 5, 6, 7],
       [8, 9, 10, 11]])
>>> x2 = np.arange(12, 20).reshape(2, 4)
>>> x2.shape
(2, 4)
>>> x2
array([[12, 13, 14, 15],
      [16, 17, 18, 19]])
>>> y = F.vstack([x1, x2])
>>> y.shape
(5, 4)
>>> y.data
array([[ 0, 1, 2, 3],
       [4, 5, 6, 7],
       [8, 9, 10, 11],
       [12, 13, 14, 15],
       [16, 17, 18, 19]])
```

# where

chainer.functions.where(condition, x, y)

Choose elements depending on condition.

This function choose values depending on a given condition. All condition, x, and y must have the same shape.

#### Parameters

- condition (Variable) Variable containing the condition. Only boolean array is permitted.
- **x** (Variable) Variable chosen when condition is True.
- **y** (Variable) Variable chosen when condition is False.

**Returns** Variable containing chosen values.

Return type Variable

# **Neural network connections**

### bilinear

chainer.functions.**bilinear** (*e1*, *e2*, *W*, *V1=None*, *V2=None*, *b=None*) Applies a bilinear function based on given parameters.

This is a building block of Neural Tensor Network (see the reference paper below). It takes two input variables and one or four parameters, and outputs one variable.

To be precise, denote six input arrays mathematically by  $e^1 \in \mathbb{R}^{I \cdot J}$ ,  $e^2 \in \mathbb{R}^{I \cdot K}$ ,  $W \in \mathbb{R}^{J \cdot K \cdot L}$ ,  $V^1 \in \mathbb{R}^{J \cdot L}$ ,  $V^2 \in \mathbb{R}^{K \cdot L}$ , and  $b \in \mathbb{R}^L$ , where I is mini-batch size. In this document, we call  $V^1$ ,  $V^2$ , and b linear parameters.

The output of forward propagation is calculated as

$$y_{il} = \sum_{jk} e_{ij}^1 e_{ik}^2 W_{jkl} + \sum_j e_{ij}^1 V_{jl}^1 + \sum_k e_{ik}^2 V_{kl}^2 + b_l V_{kl}^2 + b_{kl} V_{k$$

Note that V1, V2, b are optional. If these are not given, then this function omits the last three terms in the above equation.

Note: This function accepts an input variable e1 or e2 of a non-matrix array. In this case, the leading dimension is treated as the batch dimension, and the other dimensions are reduced to one dimension.

Note: In the original paper, J and K must be equal and the author denotes  $[V^1V^2]$  (concatenation of matrices) by V.

#### **Parameters**

- **e1** (Variable) Left input variable.
- e2 (Variable) Right input variable.
- W (Variable) Quadratic weight variable.
- V1 (Variable) Left coefficient variable.
- V2 (Variable) Right coefficient variable.
- **b** (Variable) Bias variable.

# Returns Output variable.

#### **Return type** Variable

See: Reasoning With Neural Tensor Networks for Knowledge Base Completion [Socher+, NIPS2013].

#### convolution\_2d

This is an implementation of two-dimensional convolution in ConvNets. It takes three variables: the input image x, the filter weight W, and the bias vector b.

Notation: here is a notation for dimensionalities.

 $\bullet n$  is the batch size.

 $\bullet c_I$  and  $c_O$  are the number of the input and output channels, respectively.

• $h_I$  and  $w_I$  are the height and width of the input image, respectively.

• $h_K$  and  $w_K$  are the height and width of the filters, respectively.

• $h_P$  and  $w_P$  are the height and width of the spatial padding size, respectively.

Then the Convolution2D function computes correlations between filters and patches of size  $(h_K, w_K)$  in x. Note that correlation here is equivalent to the inner product between expanded vectors. Patches are extracted at positions shifted by multiples of stride from the first position  $(-h_P, -w_P)$  for each spatial axis. The right-most (or bottom-most) patches do not run over the padded spatial size.

Let  $(s_Y, s_X)$  be the stride of filter application. Then, the output size  $(h_O, w_O)$  is determined by the following equations:

$$h_O = (h_I + 2h_P - h_K)/s_Y + 1,$$
  
 $w_O = (w_I + 2w_P - w_K)/s_X + 1.$ 

If cover\_all option is True, the filter will cover the all spatial locations. So, if the last stride of filter does not cover the end of spatial locations, an additional stride will be applied to the end part of spatial locations. In this case, the output size  $(h_O, w_O)$  is determined by the following equations:

$$h_O = (h_I + 2h_P - h_K + s_Y - 1)/s_Y + 1,$$
  
$$w_O = (w_I + 2w_P - w_K + s_X - 1)/s_X + 1$$

If the bias vector is given, then it is added to all spatial locations of the output of convolution.

The two-dimensional convolution function is defined as follows.

#### Parameters

- **x** (Variable or numpy.ndarray or cupy.ndarray) Input variable of shape  $(n, c_I, h_I, w_I)$ .
- W (Variable or numpy.ndarray or cupy.ndarray) Weight variable of shape  $(c_O, c_I, h_K, w_K)$ .
- **b** (*Variable* or numpy.ndarray or *cupy.ndarray*) Bias variable of length *c*<sub>O</sub> (optional).
- **stride** (int or pair of int s) Stride of filter applications. stride=s and stride=(s, s) are equivalent.
- **pad** (int or pair of int s) Spatial padding width for input arrays. pad=p and pad= (p, p) are equivalent.
- use\_cudnn (bool) If True, then this function uses cuDNN if available.
- cover\_all (bool) If True, all spatial locations are convoluted into some output pixels.
- **deterministic** (*bool*) The output of this function can be non-deterministic when it uses cuDNN. If this option is True, then it forces cuDNN to use a deterministic algorithm. This option is only available for cuDNN version >= v3.

**Returns** Output variable of shape  $(n, c_O, h_O, w_O)$ .

**Return type** Variable

### See also:

Convolution2D

```
>>> n = 10
>>> c_i, c_o = 3, 1
>>> h_i, w_i = 30, 40
>>> h_k, w_k = 10, 10
>>> h_p, w_p = 5, 5
>>> x = np.random.uniform(0, 1, (n, c_i, h_i, w_i)).astype('f')
>>> x.shape
(10, 3, 30, 40)
>>> W = np.random.uniform(0, 1, (c_o, c_i, h_k, w_k)).astype('f')
>>> W.shape
(1, 3, 10, 10)
>>> b = np.random.uniform(0, 1, (c_o,)).astype('f')
>>> b.shape
(1,)
>>> s_y, s_x = 5, 7
>>> y = F.convolution_2d(x, W, b, stride=(s_y, s_x), pad=(h_p, w_p))
>>> y.shape
(10, 1, 7, 6)
>>> h_0 = int((h_i + 2 * h_p - h_k) / s_y + 1)
>>> w_o = int((w_i + 2 * w_p - w_k) / s_x + 1)
>>> y.shape == (n, c_o, h_o, w_o)
True
>>> y = F.convolution_2d(x, W, b, stride=(s_y, s_x), pad=(h_p, w_p), cover_
\rightarrowall=True)
>>> y.shape == (n, c_o, h_o, w_o + 1)
True
```

# convolution\_nd

This is an implementation of N-dimensional convolution which is generalized two-dimensional convolution in ConvNets. It takes three variables: the input x, the filter weight W and the bias vector b.

Notation: here is a notation for dimensionalities.

- $\bullet N$  is the number of spatial dimensions.
- $\bullet n$  is the batch size.
- • $c_I$  and  $c_O$  are the number of the input and output channels, respectively.
- $\bullet d_1, d_2, ..., d_N$  are the size of each axis of the input's spatial dimensions, respectively.
- • $k_1, k_2, ..., k_N$  are the size of each axis of the filters, respectively.
- $\bullet l_1, l_2, ..., l_N$  are the size of each axis of the output's spatial dimensions, respectively.
- • $p_1, p_2, ..., p_N$  are the size of each axis of the spatial padding size, respectively.

Then the convolution\_nd function computes correlations between filters and patches of size  $(k_1, k_2, ..., k_N)$  in x. Note that correlation here is equivalent to the inner product between expanded tensors. Patches are extracted at positions shifted by multiples of stride from the first position  $(-p_1, -p_2, ..., -p_N)$  for each spatial axis.

Let  $(s_1, s_2, ..., s_N)$  be the stride of filter application. Then, the output size  $(l_1, l_2, ..., l_N)$  is determined by the following equations:

$$l_n = (d_n + 2p_n - k_n)/s_n + 1 \ (n = 1, ..., N)$$

If cover\_all option is True, the filter will cover the all spatial locations. So, if the last stride of filter does not cover the end of spatial locations, an additional stride will be applied to the end part of spatial locations. In this case, the output size is determined by the following equations:

$$l_n = (d_n + 2p_n - k_n + s_n - 1)/s_n + 1 \ (n = 1, ..., N)$$

The N-dimensional convolution function is defined as follows.

#### **Parameters**

- **x** (Variable or numpy.ndarray or cupy.ndarray) Input variable of shape  $(n, c_I, d_1, d_2, ..., d_N)$ .
- W (Variable or numpy.ndarray or cupy.ndarray) Weight variable of shape  $(c_O, c_I, k_1, k_2, ..., k_N)$ .
- **b** (*Variable* or numpy.ndarray or *cupy.ndarray*) One-dimensional bias variable with length  $c_O$  (optional).
- **stride** (int or tuple of int s) Stride of filter applications  $(s_1, s_2, ..., s_N)$ . stride=s is equivalent to (s, s, ..., s).
- **pad** (int or tuple of int s) Spatial padding width for input arrays  $(p_1, p_2, ..., p_N)$ . pad=p is equivalent to (p, p, ..., p).
- **use\_cudnn** (*bool*) If True, then this function uses cuDNN if available. See below for the excact conditions.
- **cover\_all** (*bool*) If True, all spatial locations are convoluted into some output pixels. It may make the output size larger. *cover\_all* needs to be False if you want to use cuDNN.

**Returns** Output variable of shape  $(n, c_O, l_1, l_2, ..., l_N)$ .

### **Return type** Variable

**Note:** This function uses cuDNN implementation for its forward and backward computation if ALL of the following conditions are satisfied:

•cuda.cudnn\_enabled is True

•use\_cudnn is True

•The number of spatial dimensions is more than one.

•cover\_all is False

•The input's dtype is equal to the filter weight's.

•The dtype is FP16, FP32 or FP64. (FP16 is only available when cuDNN version  $\geq$  v3.)

# See also:

ConvolutionND, convolution\_2d()

```
>>> n = 10
>>> c_i, c_o = 3, 1
>>> d1, d2, d3 = 30, 40, 50
>>> k1, k2, k3 = 10, 10, 10
>>> p1, p2, p3 = 5, 5, 5
>>> x = np.random.uniform(0, 1, (n, c_i, d1, d2, d3)).astype('f')
>>> x.shape
(10, 3, 30, 40, 50)
>>> W = np.random.uniform(0, 1, (c_o, c_i, k1, k2, k3)).astype('f')
>>> W.shape
(1, 3, 10, 10, 10)
>>> b = np.random.uniform(0, 1, (c_o)).astype('f')
>>> b.shape
(1,)
>>> s1, s2, s3 = 2, 4, 6
>>> y = F.convolution_nd(x, W, b, stride=(s1, s2, s3), pad=(p1, p2, p3))
>>> y.shape
(10, 1, 16, 11, 9)
>>> l1 = int((d1 + 2 * p1 - k1) / s1 + 1)
>>> 12 = int((d2 + 2 * p2 - k2) / s2 + 1)
>>> 13 = int((d3 + 2 * p3 - k3) / s3 + 1)
>>> y.shape == (n, c_o, 11, 12, 13)
True
>>> y = F.convolution_nd(x, W, b, stride=(s1, s2, s3), pad=(p1, p2, p3), cover_
\rightarrowall=True)
>>> y.shape == (n, c_o, 11, 12, 13 + 1)
True
```

# deconvolution\_2d

```
chainer.functions.deconvolution_2d(x, W, b=None, stride=1, pad=0, outsize=None,
use_cudnn=True, deterministic=False)
```

Two dimensional deconvolution function.

This is an implementation of two-dimensional deconvolution. In most of deep learning frameworks and papers, this function is called **transposed convolution**. But because of historical reasons (e.g. paper by Ziller Deconvolutional Networks) and backward compatibility, this function is called **deconvolution** in Chainer.

It takes three variables: input image x, the filter weight W, and the bias vector b.

Notation: here is a notation for dimensionalities.

 $\bullet n$  is the batch size.

 $\bullet c_I$  and  $c_O$  are the number of the input and output channels, respectively.

• $h_I$  and  $w_I$  are the height and width of the input image, respectively.

• $h_K$  and  $w_K$  are the height and width of the filters, respectively.

• $h_P$  and  $w_P$  are the height and width of the spatial padding size, respectively.

Let  $(s_Y, s_X)$  be the stride of filter application. Then, the output size  $(h_O, w_O)$  is estimated by the following equations:

$$h_O = s_Y(h_I - 1) + h_K - 2h_P,$$
  
 $w_O = s_X(w_I - 1) + w_K - 2w_P.$ 

#### **Parameters**

- **x** (Variable or numpy.ndarray or cupy.ndarray) Input variable of shape  $(n, c_I, h_I, w_I)$ .
- W (Variable or numpy.ndarray or cupy.ndarray) Weight variable of shape  $(c_I, c_O, h_K, w_K)$ .
- **b** (*Variable* or numpy.ndarray or *cupy.ndarray*) Bias variable of length *c*<sub>O</sub> (optional).
- **stride** (int or pair of int s) Stride of filter applications. stride=s and stride=(s, s) are equivalent.
- **pad** (int or pair of int s) Spatial padding width for input arrays. pad=p and pad= (p, p) are equivalent.
- **outsize** (tuple of int) Expected output size of deconvolutional operation. It should be pair of height and width  $(h_O, w_O)$ . Default value is None and the outsize is estimated by input size, stride and pad.
- **use\_cudnn** (*bool*) If True, then this function uses cuDNN if available.
- **deterministic** (*bool*) The output of this function can be non-deterministic when it uses cuDNN. If this option is True, then it forces cuDNN to use a deterministic algorithm. This option is only available for cuDNN version >= v3.

**Returns** Output variable of shape  $(n, c_O, h_O, w_O)$ .

Return type Variable

```
>>> n = 10
>>> c_i, c_o = 1, 3
>>> h i, w i = 5, 10
>>> h_k, w_k = 10, 10
>>> h_p, w_p = 5, 5
>>> x = np.random.uniform(0, 1, (n, c_i, h_i, w_i)).astype('f')
>>> x.shape
(10, 1, 5, 10)
>>> W = np.random.uniform(0, 1, (c_i, c_o, h_k, w_k)).astype('f')
>>> W.shape
(1, 3, 10, 10)
>>> b = np.random.uniform(0, 1, c_o).astype('f')
>>> b.shape
(3,)
>>> s_y, s_x = 5, 5
>>> y = F.deconvolution_2d(x, W, b, stride=(s_y, s_x), pad=(h_p, w_p))
>>> y.shape
(10, 3, 20, 45)
>>> h_0 = s_y * (h_i - 1) + h_k - 2 * h_p
>>> w_0 = s_x * (w_i - 1) + w_k - 2 * w_p
>>> y.shape == (n, c_o, h_o, w_o)
True
```

# deconvolution\_nd

N-dimensional deconvolution function.

This is an implementation of N-dimensional deconvolution which generalizes two-dimensional one. In most of deep learning frameworks and papers, this function is called **transposed convolution**. But because of historical reasons (e.g. paper by Ziller Deconvolutional Networks) and backward compatibility, this function is called **deconvolution** in Chainer.

It takes three variables: the input x, the filter weight W, and the bias vector b.

Notation: here is a notation for dimensionalities.

 $\bullet N$  is the number of spatial dimensions.

 $\bullet n$  is the batch size.

 $\bullet c_I$  and  $c_O$  are the number of the input and output channels, respectively.

 $\bullet d_1, d_2, ..., d_N$  are the size of each axis of the input's spatial dimensions, respectively.

• $k_1, k_2, ..., k_N$  are the size of each axis of the filters, respectively.

• $p_1, p_2, ..., p_N$  are the size of each axis of the spatial padding size, respectively.

 $\bullet s_1, s_2, \dots, s_N$  are the stride of each axis of filter application, respectively.

If outsize option is None, the output size  $(l_1, l_2, ..., l_N)$  is determined by the following equations with the items in the above list:

$$l_n = s_n(d_n - 1) + k_n - 2p_n \ (n = 1, ..., N)$$

If outsize option is given, the output size is determined by outsize. In this case, the outsize  $(l_1, l_2, ..., l_N)$  must satisfy the following equations:

$$d_n = |(l_n + 2p_n - k_n)/s_n| + 1 \ (n = 1, ..., N)$$

#### **Parameters**

- **x** (Variable or numpy.ndarray or cupy.ndarray) Input variable of shape  $(n, c_I, d_1, d_2, ..., d_N)$ .
- W (Variable or numpy.ndarray or cupy.ndarray) Weight variable of shape  $(c_I, c_O, k_1, k_2, ..., k_N)$ .
- **b** (*Variable* or numpy.ndarray or *cupy.ndarray*) One-dimensional bias variable with length *c*<sub>O</sub> (optional).
- stride (int or tuple of int s) Stride of filter applications  $(s_1, s_2, ..., s_N)$ . stride=s is equivalent to (s, s, ..., s).
- pad (int or tuple of int s) Spatial padding width for input arrays  $(p_1, p_2, ..., p_N)$ . pad=p is equivalent to (p, p, ..., p).
- **outsize** (tuple of int s) Expected output size of deconvolutional operation. It should be a tuple of ints  $(l_1, l_2, ..., l_N)$ . Default value is None and the outsize is estimated by input size, stride and pad.
- **use\_cudnn** (*bool*) If True, then this function uses cuDNN if available. Note that cuDNN supports more than one-dimensional deconvolution operations only.

**Returns** Output variable of shape  $(n, c_O, l_1, l_2, ..., l_N)$ .

### Return type Variable

# See also:

links.DeconvolutionND, deconvolution\_2d()

### Example

**Example1**: the case when outsize is not given.

```
>>> n = 10
>>> c_i, c_o = 3, 1
>>> d1, d2, d3 = 5, 10, 15
>>> k1, k2, k3 = 10, 10, 10
>>> p1, p2, p3 = 5, 5, 5
>>> x = np.random.uniform(0, 1, (n, c_i, d1, d2, d3)).astype('f')
>>> x.shape
(10, 3, 5, 10, 15)
>>> W = np.random.uniform(0, 1, (c_i, c_o, k1, k2, k3)).astype('f')
>>> W.shape
(3, 1, 10, 10, 10)
>>> b = np.random.uniform(0, 1, (c_o)).astype('f')
>>> b.shape
(1,)
>>> s1, s2, s3 = 2, 4, 6
>>> y = F.deconvolution_nd(x, W, b, stride=(s1, s2, s3), pad=(p1, p2, p3))
>>> y.shape
(10, 1, 8, 36, 84)
>>> 11 = s1 * (d1 - 1) + k1 - 2 * p1
>>> 12 = s2 * (d2 - 1) + k2 - 2 * p2
>>> 13 = s3 * (d3 - 1) + k3 - 2 * p3
>>> y.shape == (n, c_o, 11, 12, 13)
True
```

**Example2**: the case when outsize is given.

```
>>> n = 10
>>> c_i, c_o = 3, 1
>>> d1, d2, d3 = 5, 10, 15
>>> k1, k2, k3 = 10, 10, 10
>>> p1, p2, p3 = 5, 5, 5
>>> x = np.random.uniform(0, 1, (n, c_i, d1, d2, d3)).astype('f')
>>> x.shape
(10, 3, 5, 10, 15)
>>> W = np.random.uniform(0, 1, (c_i, c_o, k1, k2, k3)).astype('f')
>>> W.shape
(3, 1, 10, 10, 10)
>>> b = np.random.uniform(0, 1, (c_o)).astype('f')
>>> b.shape
(1,)
>>> s1, s2, s3 = 2, 4, 6
>>> 11, 12, 13 = 9, 38, 87
>>> d1 == int((l1 + 2 * p1 - k1) / s1) + 1
True
>>> d2 == int((12 + 2 * p2 - k2) / s2) + 1
True
>>> d3 == int((13 + 2 * p3 - k3) / s3) + 1
True
>>> y = F.deconvolution_nd(x, W, b, stride=(s1, s2, s3), pad=(p1, p2, p3),_
→outsize=(11, 12, 13))
```

```
>>> y.shape
(10, 1, 9, 38, 87)
>>> y.shape == (n, c_o, 11, 12, 13)
True
```

# depthwise\_convolution\_2d

```
chainer.functions.depthwise_convolution_2d(x, W, b=None, stride=1, pad=0)
Two-dimensional depthwise convolution function.
```

This is an implementation of two-dimensional depthwise convolution. It takes two or three variables: the input image x, the filter weight W, and optionally, the bias vector b.

Notation: here is a notation for dimensionalities.

- $\bullet n$  is the batch size.
- • $c_I$  is the number of the input.
- • $c_M$  is the channel multiplier.

•h and w are the height and width of the input image, respectively.

• $h_O$  and  $w_O$  are the height and width of the output image, respectively.

• $k_H$  and  $k_W$  are the height and width of the filters, respectively.

### Parameters

- **x** (chainer.Variable or numpy.ndarray or cupy.ndarray) Input variable of shape  $(n, c_I, h, w)$ .
- W (Variable) Weight variable of shape  $(c_M, c_I, k_H, k_W)$ .
- **b** (Variable) Bias variable of length  $c_M * c_I$  (optional).
- **stride** (*int or pair of ints*) Stride of filter applications. stride=s and stride=(s, s) are equivalent.
- **pad** (*int* or *pair* of *ints*) Spatial padding width for input arrays. pad=p and pad=(p, p) are equivalent.

**Returns** Output variable. Its shape is  $(n, c_I * c_M, h_O, w_O)$ .

# **Return type** Variable

Like Convolution2D, DepthwiseConvolution2D function computes correlations between filters and patches of size  $(k_H, k_W)$  in x. But unlike Convolution2D, DepthwiseConvolution2D does not add up input channels of filters but concatenates them. For that reason, the shape of outputs of depthwise convolution are  $(n, c_I * c_M, h_O, w_O)$ ,  $c_M$  is called channel\_multiplier.

 $(h_O, w_O)$  is determined by the equivalent equation of Convolution2D.

If the bias vector is given, then it is added to all spatial locations of the output of convolution.

See: L. Sifre. Rigid-motion scattering for image classification

# See also:

DepthwiseConvolution2D

### Example

```
>>> x = np.random.uniform(0, 1, (2, 3, 4, 7))
>>> W = np.random.uniform(0, 1, (2, 3, 3, 3))
>>> b = np.random.uniform(0, 1, (6,))
>>> y = F.depthwise_convolution_2d(x, W, b)
>>> y.shape
(2, 6, 2, 5)
```

# dilated\_convolution\_2d

Two-dimensional dilated convolution function.

This is an implementation of two-dimensional dilated convolution in ConvNets. It takes three variables: the input image x, the filter weight W, and the bias vector b.

Notation: here is a notation for dimensionalities.

 $\bullet n$  is the batch size.

• $c_I$  and  $c_O$  are the number of the input and output, respectively.

 $\bullet h$  and w are the height and width of the input image, respectively.

• $k_H$  and  $k_W$  are the height and width of the filters, respectively.

### **Parameters**

- **x** (Variable) Input variable of shape  $(n, c_I, h, w)$ .
- W (Variable) Weight variable of shape  $(c_O, c_I, k_H, k_W)$ .
- **b** (Variable) Bias variable of length  $c_O$  (optional).
- **stride** (*int or pair of ints*) Stride of filter applications. stride=s and stride=(s, s) are equivalent.
- **pad** (*int or pair of ints*) Spatial padding width for input arrays. pad=p and pad=(p, p) are equivalent.
- **dilate** (*int* or *pair* of *ints*) Dilation factor of filter applications. dilate=d and dilate=(d, d) are equivalent.
- **use\_cudnn** (*bool*) If True, then this function uses cuDNN if available.
- **cover\_all** (*bool*) If True, all spatial locations are convoluted into some output pixels. It may make the output size larger.

**Returns** Output variable.

## Return type Variable

The two-dimensional dilated convolution function is defined as follows. Then the DilatedConvolution2D function computes correlations between filters and patches of size  $(k_H, k_W)$  in x. Patches here are extracted at intervals of the dilation factor. Note that correlation here is equivalent to the inner product between expanded vectors. Patches are extracted at intervals of the dilation factor and at positions shifted by multiples of stride from the first position -pad for each spatial axis. The right-most (or bottom-most) patches do not run over the padded spatial size.

Let  $(s_Y, s_X)$  be the stride of filter application,  $(p_H, p_W)$  the spatial padding size, and  $(d_Y, d_X)$  the dilation factor of filter application. Then, the output size  $(h_O, w_O)$  is determined by the following equations:

$$h_O = (h + 2p_H - k_H - (k_H - 1) * (d_Y - 1))/s_Y + 1,$$
  
$$w_O = (w + 2p_W - k_W - (k_W - 1) * (d_X - 1))/s_X + 1.$$

If the bias vector is given, then it is added to all spatial locations of the output of convolution.

### See also:

DilatedConvolution2D

### embed\_id

```
chainer.functions.embed_id(x, W, ignore_label=None)
```

Efficient linear function for one-hot input.

This function implements so called *word embedding*. It takes two arguments: a set of IDs (words)  $\times$  in B dimensional integer vector, and a set of all ID (word) embeddings W in  $V \times d$  float32 matrix. It outputs  $B \times d$  matrix whose i-th column is the  $\times$  [i]-th column of W.

This function is only differentiable on the input W.

### Parameters

- **x** (Variable) Batch vectors of IDs.
- W (Variable) Representation of each ID (a.k.a. word embeddings).
- **ignore\_label** (*int or None*) If ignore\_label is an int value, i-th column of return value is filled with 0.

Returns Output variable.

Return type Variable

# See also:

EmbedID

# linear

chainer.functions.linear(x, W, b=None)

Linear function, or affine transformation.

It accepts two or three arguments: an input minibatch x, a weight matrix W, and optionally a bias vector b. It computes

$$Y = xW^{\top} + b.$$

### **Parameters**

- **x** (*Variable* or numpy.ndarray or *cupy.ndarray*) Input variable, which is a  $(s_B, s_1, s_2, ..., s_n)$ -shaped float array. Its first dimension  $(s_B)$  is assumed to be the *minibatch dimension*. The other dimensions are treated as concatenated one dimension whose size must be  $(s_1 * ... * s_n = N)$ .
- W (Variable or numpy.ndarray or cupy.ndarray) Weight variable of shape (M, N), where  $(N = s_1 * ... * s_n)$ .

• **b** (*Variable* or numpy.ndarray or *cupy.ndarray*) – Bias variable (optional) of shape (*M*,).

**Returns** Output variable. A float array with shape of  $(s_B, M)$ .

**Return type** Variable

### See also:

Linear

# Example

```
>>> x = np.random.uniform(0, 1, (3, 4)).astype('f')
>>> W = np.random.uniform(0, 1, (5, 4)).astype('f')
>>> b = np.random.uniform(0, 1, (5,)).astype('f')
>>> y = F.linear(x, W, b)
>>> y.shape
(3, 5)
```

# n\_step\_bigru

chainer.functions.**n\_step\_bigru**(*n\_layers*, *dropout\_ratio*, *hx*, *ws*, *bs*, *xs*, *train=True*, *use\_cudnn=True*)

Stacked Bi-directional Gated Recurrent Unit function.

This function calculates stacked Bi-directional GRU with sequences. This function gets an initial hidden state  $h_0$ , an input sequence x, weight matrices W, and bias vectors b. This function calculates hidden states  $h_t$  for each time t from input  $x_t$ .

$$\begin{split} r_t^f &= \sigma(W_0^f x_t + W_3^f h_{t-1} + b_0^f + b_3^f) \\ z_t^f &= \sigma(W_1^f x_t + W_4^f h_{t-1} + b_1^f + b_4^f) \\ h_t^{f'} &= \tanh(W_2^f x_t + b_2^f + r_t^f \cdot (W_5^f h_{t-1} + b_5^f)) \\ h_t^f &= (1 - z_t^f) \cdot h_t^{f'} + z_t^f \cdot h_{t-1} \\ r_t^b &= \sigma(W_0^b x_t + W_3^b h_{t-1} + b_0^b + b_3^b) \\ z_t^b &= \sigma(W_1^b x_t + W_4^b h_{t-1} + b_1^b + b_4^b) \\ h_t^{b'} &= \tanh(W_2^b x_t + b_2^b + r_t^b \cdot (W_5^b h_{t-1} + b_5^b)) \\ h_t^b &= (1 - z_t^b) \cdot h_t^{b'} + z_t^b \cdot h_{t-1} \\ h_t &= [h_t^f; h_t^f] \end{split}$$

where  $W^{f}$  is weight matrices for forward-GRU,  $W^{b}$  is weight matrices for backward-GRU.

As the function accepts a sequence, it calculates  $h_t$  for all t with one call. Six weight matrices and six bias vectors are required for each layers. So, when S layers exists, you need to prepare 6S weight matrices and 6S bias vectors.

If the number of layers  $n_{layers}$  is greather than 1, input of k-th layer is hidden state  $h_t$  of k-1-th layer. Note that all input variables except first layer may have different shape from the first layer.

#### **Parameters**

- n\_layers (int) Number of layers.
- **dropout\_ratio** (*float*) Dropout ratio.

- hx (chainer.Variable) Variable holding stacked hidden states. Its shape is (S, B, N) where S is number of layers and is equal to n\_layers, B is mini-batch size, and N is dimention of hidden units.
- **ws** (list of list of chainer.Variable) Weight matrices. ws[i] represents weights for i-th layer. Each ws[i] is a list containing six matrices. ws[i][j] is corresponding with W\_j in the equation. Only ws[0][j] where 0 <= j < 3 is (I, N) shape as they are multiplied with input variables. All other matrices has (N, N) shape.
- **bs** (list of list of chainer.Variable) Bias vectors. bs[i] represents biases for i-th layer. Each bs[i] is a list containing six vectors. bs[i][j] is corresponding with b\_j in the equation. Shape of each matrix is (N, ) where N is dimension of hidden units.
- **xs** (list of chainer.Variable) A list of Variable holding input values. Each element xs[t] holds input value for time t. Its shape is (B\_t, I), where B\_t is minibatch size for time t, and I is size of input units. Note that this functions supports variable length sequences. When sequences has different lengths, sort sequences in descending order by length, and transpose the sorted sequence. transpose\_sequence() transpose a list of Variable() holding sequence. So xs needs to satisfy xs[t].shape[0] >= xs[t + 1].shape[0].
- **train** (*bool*) If True, this function executes dropout.
- use\_cudnn (bool) If True, this function uses cuDNN if available.
- use\_bi\_direction (bool) If True, this function uses Bi-direction GRU.

# Returns

**This functions returns a tuple concaining three elements,** hy and ys. - hy is an updated hidden states whose shape is same as hx. - ys is a list of *Variable*. Each element

ys[t] holds hidden states of the last layer corresponding to an input xs[t]. Its shape is (B\_t, N) where B\_t is mini-batch size for time t, and N is size of hidden units. Note that B\_t is the same value as xs[t].

# Return type tuple

# n\_step\_bilstm

chainer.functions.n\_step\_bilstm(n\_layers, dropout\_ratio, hx, cx, ws, bs, xs, train=True, use\_cudnn=True) Stacked Bi-directional Long Short-Term Memory function.

This function calculates stacked Bi-directional LSTM with sequences. This function gets an initial hidden state  $h_0$ , an initial cell state  $c_0$ , an input sequence x, weight matrices W, and bias vectors b. This function calculates

hidden states  $h_t$  and  $c_t$  for each time t from input  $x_t$ .

$$\begin{split} i_t^f &= \\ \sigma(W_0^f x_t + W_4^f h_{t-1} + b_0^f + b_4^f), \\ f_t^f &= \\ \sigma(W_1^f x_t + W_5^f h_{t-1} + b_1^f + b_5^f), \\ o_t^f &= \\ \sigma(W_2^f x_t + W_6^f h_{t-1} + b_2^f + b_6^f), \\ a_t^f &= \\ \\ tanh(W_3^f x_t + W_7^f h_{t-1} + b_3^f + b_7^f), \\ c_t^f &= \\ f_t^f \cdot c_{t-1}^f + i_t^f \cdot a_t^f, \\ h_t^f &= \\ o_t^f \cdot tanh(c_t^f), \\ i_t^b &= \\ \sigma(W_0^b x_t + W_4^b h_{t-1} + b_0^b + b_4^b), \\ f_t^b &= \\ \sigma(W_1^b x_t + W_5^b h_{t-1} + b_1^b + b_5^b), \\ o_t^b &= \\ \sigma(W_2^b x_t + W_6^b h_{t-1} + b_2^b + b_6^b), \\ a_t^b &= \\ tanh(W_3^b x_t + W_7^b h_{t-1} + b_3^b + b_7^b), \\ c_t^b &= \\ f_t^b \cdot c_{t-1}^b + i_t^b \cdot a_t^b, \\ h_t^b &= \\ o_t^b \cdot tanh(c_t^b), \\ h_t &= \\ [h^f; h^b] \end{split}$$

where  $W^{f}$  is weight matrices for forward-LSTM,  $W^{b}$  is weight matrices for backward-LSTM.

As the function accepts a sequence, it calculates  $h_t$  for all t with one call. Eight weight matrices and eight bias vectors are required for each layers. So, when S layers exists, you need to prepare 8S weight matrices and 8S bias vectors.

If the number of layers  $n_{layers}$  is greather than 1, input of k-th layer is hidden state  $h_t$  of k-1-th layer. Note that all input variables except first layer may have different shape from the first layer.

# **Parameters**

- n\_layers (int) Number of layers.
- dropout\_ratio (float) Dropout ratio.
- hx (chainer.Variable) Variable holding stacked hidden states. Its shape is (S, B, N) where S is number of layers and is equal to n\_layers, B is mini-batch size, and N is dimention of hidden units.

- **cx** (chainer.Variable) Variable holding stacked cell states. It has the same shape as hx.
- ws (list of list of chainer.Variable) Weight matrices. ws [i] represents weights for i-th layer. Each ws [i] is a list containing eight matrices. ws [i] [j] is corresponding with W\_j in the equation. Only ws [0] [j] where 0 <= j < 4 is (I, N) shape as they are multiplied with input variables. All other matrices has (N, N) shape.
- **bs** (list of list of chainer.Variable) Bias vectors. bs[i] represents biases for i-th layer. Each bs[i] is a list containing eight vectors. bs[i][j] is corresponding with b\_j in the equation. Shape of each matrix is (N, ) where N is dimension of hidden units.
- **xs** (list of chainer.Variable) A list of Variable holding input values. Each element xs[t] holds input value for time t. Its shape is (B\_t, I), where B\_t is minibatch size for time t, and I is size of input units. Note that this functions supports variable length sequences. When sequences has different lengths, sort sequences in descending order by length, and transpose the sorted sequence. transpose\_sequence() transpose a list of Variable() holding sequence. So xs needs to satisfy xs[t].shape[0] >= xs[t + 1].shape[0].
- train (bool) If True, this function executes dropout.
- use\_cudnn (bool) If True, this function uses cuDNN if available.

### Returns

This functions returns a tuple concaining three elements, hy, cy and ys. - hy is an updated hidden states whose shape is same as hx. - cy is an updated cell states whose shape is same as cx. - ys is a list of *Variable*. Each element

ys[t] holds hidden states of the last layer corresponding to an input xs[t]. Its shape is (B\_t, N) where B\_t is mini-batch size for time t, and N is size of hidden units. Note that B\_t is the same value as xs[t].

# Return type tuple

# n\_step\_birnn

chainer.functions.n\_step\_birnn(n\_layers, dropout\_ratio, hx, ws, bs, xs, train=True, use\_cudnn=True, activation='tanh')

Stacked Bi-directional RNN function for sequence inputs.

This function calculates stacked Bi-directional RNN with sequences. This function gets an initial hidden state  $h_0$ , an initial cell state  $c_0$ , an input sequence x, weight matrices W, and bias vectors b. This function calculates hidden states  $h_t$  and  $c_t$  for each time t from input  $x_t$ .

$$\begin{aligned} h_{t}^{f} &= \\ f(W_{0}^{f}x_{t} + W_{1}^{f}h_{t-1} + b_{0}^{f} + b_{1}^{f}), \\ h_{t}^{b} &= \\ f(W_{0}^{b}x_{t} + W_{1}^{b}h_{t-1} + b_{0}^{b} + b_{1}^{b}), \\ h_{t} &= \\ & [h_{t}^{f}; h_{t}^{f}], \end{aligned}$$

where f is an activation function.

Weight matrices W contains two matrices  $W^f$  and  $W^b$ .  $W^f$  is weight matrices for forward directional RNN.  $W^b$  is weight matrices for backward directional RNN.

 $W_1^f$  contains  $W_0^f$  for an input sequence and  $W_1^f$  for a hidden state.  $W^b$  contains  $W_0^b$  for an input sequence and  $W_1^b$  for a hidden state.

Bias matrices b contains two matrices  $b^f$  and  $b^f$ .  $b^f$  contains  $b_0^f$  for an input sequence and  $b_1^f$  for a hidden state.  $b^b$  contains  $b_0^b$  for an input sequence and  $b_1^b$  for a hidden state.

As the function accepts a sequence, it calculates  $h_t$  for all t with one call. Two weight matrices and two bias vectors are required for each layer. So, when S layers exist, you need to prepare 2S weight matrices and 2S bias vectors.

If the number of layers  $n_{layers}$  is greather than 1, input of k-th layer is hidden state  $h_t$  of k-1-th layer. Note that all input variables except first layer may have different shape from the first layer.

# Parameters

- n\_layers (*int*) Number of layers.
- **dropout\_ratio** (*float*) Dropout ratio.
- hx (chainer.Variable) Variable holding stacked hidden states. Its shape is (S, B, N) where S is number of layers and is equal to n\_layers, B is mini-batch size, and N is dimention of hidden units.
- ws (list of list of chainer.Variable) Weight matrices. ws[i + di] represents weights for i-th layer. Note that di = 0 for forward-RNN and di = 1 for backward-RNN. Each ws[i + di] is a list containing two matrices. ws[i + di][j] is corresponding with W^{ff\_j if di = 0 and corresponding with W^{b\_j if di = 1 in the equation. Only ws[0][j] and ws[1][j] where 0 <= j < 1 are (I, N) shape as they are multiplied with input variables. All other matrices has (N, N) shape.
- **bs** (list of list of chainer.Variable)-Bias vectors. bs[i + di] represnents biases for i-th layer. Note that di = 0 for forward-RNN and di = 1 for backward-RNN. Each bs[i + di] is a list containing two vectors. bs[i + di][j] is corresponding with b^{f}\_j if di = 0 and corresponding with b^{b}\_j if di = 1 in the equation. Shape of each matrix is (N, ) where N is dimention of hidden units.
- **xs** (list of chainer.Variable) A list of Variable holding input values. Each element xs[t] holds input value for time t. Its shape is (B\_t, I), where B\_t is minibatch size for time t, and I is size of input units. Note that this functions supports variable length sequences. When sequences has different lengths, sort sequences in descending order by length, and transpose the sorted sequence. transpose\_sequence() transpose a list of Variable() holding sequence. So xs needs to satisfy xs[t].shape[0] >= xs[t + 1].shape[0].
- **train** (*bool*) If True, this function executes dropout.
- use\_cudnn (bool) If True, this function uses cuDNN if available.
- **activation** (*str*) Activation function name. Please select tanh or relu.

# Returns

This functions returns a tuple concaining three elements, hy and ys.

- hy is an updated hidden states whose shape is same as hx.
- ys is a list of *Variable*. Each element ys[t] holds hidden states of the last layer corresponding to an input xs[t]. Its shape is (B\_t, N) where B\_t is mini-batch size for time t, and N is size of hidden units. Note that B\_t is the same value as xs[t].

### Return type tuple

# n\_step\_gru

chainer.functions.**n\_step\_gru**(*n\_layers*, *dropout\_ratio*, *hx*, *ws*, *bs*, *xs*, *train=True*, *use\_cudnn=True*)

Stacked Uni-directional Gated Recurrent Unit function.

This function calculates stacked Uni-directional GRU with sequences. This function gets an initial hidden state  $h_0$ , an input sequence x, weight matrices W, and bias vectors b. This function calculates hidden states  $h_t$  for each time t from input  $x_t$ .

$$r_{t} = \sigma(W_{0}x_{t} + W_{3}h_{t-1} + b_{0} + b_{3})$$

$$z_{t} = \sigma(W_{1}x_{t} + W_{4}h_{t-1} + b_{1} + b_{4})$$

$$h'_{t} = \tanh(W_{2}x_{t} + b_{2} + r_{t} \cdot (W_{5}h_{t-1} + b_{5}))$$

$$h_{t} = (1 - z_{t}) \cdot h'_{t} + z_{t} \cdot h_{t-1}$$

As the function accepts a sequence, it calculates  $h_t$  for all t with one call. Six weight matrices and six bias vectors are required for each layers. So, when S layers exists, you need to prepare 6S weight matrices and 6S bias vectors.

If the number of layers  $n_{layers}$  is greather than 1, input of k-th layer is hidden state  $h_t$  of k-1-th layer. Note that all input variables except first layer may have different shape from the first layer.

#### Parameters

- **n\_layers** (*int*) Number of layers.
- dropout\_ratio (float) Dropout ratio.
- **hx** (chainer.Variable) Variable holding stacked hidden states. Its shape is (S, B, N) where S is number of layers and is equal to n\_layers, B is mini-batch size, and N is dimention of hidden units.
- **ws** (list of list of chainer.Variable) Weight matrices. ws[i] represents weights for i-th layer. Each ws[i] is a list containing six matrices. ws[i][j] is corresponding with W\_j in the equation. Only ws[0][j] where 0 <= j < 3 is (I, N) shape as they are multiplied with input variables. All other matrices has (N, N) shape.
- **bs** (list of list of chainer.Variable) Bias vectors. bs[i] represents biases for i-th layer. Each bs[i] is a list containing six vectors. bs[i][j] is corresponding with b\_j in the equation. Shape of each matrix is (N, ) where N is dimension of hidden units.
- **xs** (list of chainer.Variable) A list of Variable holding input values. Each element xs[t] holds input value for time t. Its shape is (B\_t, I), where B\_t is minibatch size for time t, and I is size of input units. Note that this functions supports variable length sequences. When sequences has different lengths, sort sequences in descending order by length, and transpose the sorted sequence. transpose\_sequence() transpose a list of Variable() holding sequence. So xs needs to satisfy xs[t].shape[0] >= xs[t + 1].shape[0].
- **train** (*bool*) If True, this function executes dropout.
- use\_cudnn (bool) If True, this function uses cuDNN if available.

# Returns

**This functions returns a tuple concaining three elements,** hy and ys. - hy is an updated hidden states whose shape is same as hx. - ys is a list of *Variable*. Each element

ys[t] holds hidden states of the last layer corresponding to an input xs[t]. Its shape is (B\_t, N) where B\_t is mini-batch size for time t, and N is size of hidden units. Note that B\_t is the same value as xs[t].

# Return type tuple

# n\_step\_lstm

Stacked Uni-directional Long Short-Term Memory function.

This function calculates stacked Uni-directional LSTM with sequences. This function gets an initial hidden state  $h_0$ , an initial cell state  $c_0$ , an input sequence x, weight matrices W, and bias vectors b. This function calculates hidden states  $h_t$  and  $c_t$  for each time t from input  $x_t$ .

$$i_{t} = \sigma(W_{0}x_{t} + W_{4}h_{t-1} + b_{0} + b_{4})$$

$$f_{t} = \sigma(W_{1}x_{t} + W_{5}h_{t-1} + b_{1} + b_{5})$$

$$o_{t} = \sigma(W_{2}x_{t} + W_{6}h_{t-1} + b_{2} + b_{6})$$

$$a_{t} = \tanh(W_{3}x_{t} + W_{7}h_{t-1} + b_{3} + b_{7})$$

$$c_{t} = f_{t} \cdot c_{t-1} + i_{t} \cdot a_{t}$$

$$h_{t} = o_{t} \cdot \tanh(c_{t})$$

As the function accepts a sequence, it calculates  $h_t$  for all t with one call. Eight weight matrices and eight bias vectors are required for each layers. So, when S layers exists, you need to prepare 8S weight matrices and 8S bias vectors.

If the number of layers  $n_{layers}$  is greather than 1, input of k-th layer is hidden state  $h_t$  of k-1-th layer. Note that all input variables except first layer may have different shape from the first layer.

- n\_layers (int) Number of layers.
- **dropout\_ratio** (*float*) Dropout ratio.
- hx (chainer.Variable) Variable holding stacked hidden states. Its shape is (S, B, N) where S is number of layers and is equal to n\_layers, B is mini-batch size, and N is dimention of hidden units.
- **cx** (chainer.Variable) Variable holding stacked cell states. It has the same shape as hx.
- **ws** (list of list of chainer.Variable) Weight matrices. ws[i] represents weights for i-th layer. Each ws[i] is a list containing eight matrices. ws[i][j] is corresponding with W\_j in the equation. Only ws[0][j] where 0 <= j < 4 is (I, N) shape as they are multiplied with input variables. All other matrices has (N, N) shape.
- **bs** (list of list of chainer.Variable) Bias vectors. bs[i] represents biases for i-th layer. Each bs[i] is a list containing eight vectors. bs[i][j] is corresponding with b\_j in the equation. Shape of each matrix is (N, ) where N is dimension of hidden units.
- **xs** (list of chainer.Variable) A list of Variable holding input values. Each element xs[t] holds input value for time t. Its shape is (B\_t, I), where B\_t is minibatch size for time t, and I is size of input units. Note that this functions supports variable length sequences. When sequences has different lengths, sort sequences in descending order by length, and transpose the sorted sequence. transpose\_sequence() transpose a list of Variable() holding sequence. So xs needs to satisfy xs[t].shape[0] >= xs[t + 1].shape[0].
- **train** (*bool*) If True, this function executes dropout.

• **use\_cudnn** (*bool*) – If True, this function uses cuDNN if available.

### Returns

**This functions returns a tuple concaining three elements,** hy, cy and ys. - hy is an updated hidden states whose shape is same as hx. - cy is an updated cell states whose shape is same as cx. - ys is a list of *Variable*. Each element

ys[t] holds hidden states of the last layer corresponding to an input xs[t]. Its shape is (B\_t, N) where B\_t is mini-batch size for time t, and N is size of hidden units. Note that B\_t is the same value as xs[t].

Return type tuple

### See also:

chainer.functions.lstm()

# n\_step\_rnn

chainer.functions.n\_step\_rnn(n\_layers, dropout\_ratio, hx, ws, bs, xs, train=True, use\_cudnn=True, activation='tanh') Stacked Uni-directional RNN function for sequence inputs.

This function calculates stacked Uni-directional RNN with sequences. This function gets an initial hidden state  $h_0$ , an initial cell state  $c_0$ , an input sequence x, weight matrices W, and bias vectors b. This function calculates hidden states  $h_t$  and  $c_t$  for each time t from input  $x_t$ .

$$h_t = f(W_0 x_t + W_1 h_{t-1} + b_0 + b_1)$$

where f is an activation function.

Weight matrices W contains two matrices  $W_0$  and  $W_1$ .  $W_0$  is a parameter for an input sequence.  $W_1$  is a parameter for a hidden state. Bias matrices b contains two matrices  $b_0$  and  $b_1$ .  $b_0$  is a parameter for an input sequence.  $b_1$  is a parameter for a hidden state.

As the function accepts a sequence, it calculates  $h_t$  for all t with one call. Two weight matrices and two bias vectors are required for each layer. So, when S layers exist, you need to prepare 2S weight matrices and 2S bias vectors.

If the number of layers  $n_{layers}$  is greather than 1, input of k-th layer is hidden state  $h_t$  of k-1-th layer. Note that all input variables except first layer may have different shape from the first layer.

- n\_layers (int) Number of layers.
- dropout\_ratio (float) Dropout ratio.
- hx (chainer.Variable) Variable holding stacked hidden states. Its shape is (S, B, N) where S is number of layers and is equal to n\_layers, B is mini-batch size, and N is dimention of hidden units.
- ws (list of list of chainer.Variable) Weight matrices. ws [i] represents weights for i-th layer. Each ws [i] is a list containing two matrices. ws [i] [j] is corresponding with W\_j in the equation. Only ws [0] [j] where 0 <= j < 1 is (I, N) shape as they are multiplied with input variables. All other matrices has (N, N) shape.
- **bs** (list of list of chainer.Variable) Bias vectors. bs[i] represents biases for i-th layer. Each bs[i] is a list containing two vectors. bs[i][j] is corresponding with b\_j in the equation. Shape of each matrix is (N, ) where N is dimension of hidden units.

- **xs** (list of chainer.Variable) A list of Variable holding input values. Each element xs[t] holds input value for time t. Its shape is (B\_t, I), where B\_t is minibatch size for time t, and I is size of input units. Note that this functions supports variable length sequences. When sequences has different lengths, sort sequences in descending order by length, and transpose the sorted sequence. transpose\_sequence() transpose a list of Variable() holding sequence. So xs needs to satisfy xs[t].shape[0] >= xs[t + 1].shape[0].
- train (bool) If True, this function executes dropout.
- **use\_cudnn** (*bool*) If True, this function uses cuDNN if available.
- **activation** (*str*) Activation function name. Please select tanh or relu.

### Returns

### This functions returns a tuple concaining three elements, hy and ys.

- hy is an updated hidden states whose shape is same as hx.
- ys is a list of *Variable*. Each element ys[t] holds hidden states of the last layer corresponding to an input xs[t]. Its shape is (B\_t, N) where B\_t is mini-batch size for time t, and N is size of hidden units. Note that B\_t is the same value as xs[t].

Return type tuple

# **Evaluation functions**

### accuracy

chainer.functions.**accuracy**(*y*, *t*, *ignore\_label=None*) Computes multiclass classification accuracy of the minibatch.

### Parameters

- $\mathbf{y}$  (Variable or numpy.ndarray or cupy.ndarray) Array whose (i, j, k, ...)-th element indicates the score of the class j at the (i, k, ...)-th sample. The prediction label  $\hat{t}$  is calculated by the formula  $\hat{t}(i, k, ...) = \operatorname{argmax}_{i} y(i, j, k, ...)$ .
- t (Variable or numpy.ndarray or cupy.ndarray of numpy.int32) Array of ground truth labels.
- **ignore\_label** (*int or None*) Skip calculating accuracy if the true label is ignore\_label.

**Returns** A variable holding a scalar array of the accuracy.

## Return type Variable

Note: This function is non-differentiable.

### Example

We show the most common case, when y is the two dimensional array.

# binary\_accuracy

chainer.functions.**binary\_accuracy**(y, t)

Computes binary classification accuracy of the minibatch.

## Parameters

- **y** (Variable) Variable holding a matrix whose i-th element indicates the score of positive at the i-th example.
- t (Variable) Variable holding an int32 vector of ground truth labels. If t [i] == -1, corresponding x [i] is ignored. Accuracy is zero if all ground truth labels are -1.

Returns A variable holding a scalar array of the accuracy.

**Return type** Variable

Note: This function is non-differentiable.

## classification\_summary

```
chainer.functions.classification_summary(y, t, label_num=None, beta=1.0, ignore_label=-
```

*Calculates Precision, Recall, F beta Score, and support.* 

This function calculates the following quantities for each class.

•Precision:  $\frac{\text{tp}}{\text{tp+fp}}$ 

•Recall:  $\frac{\text{tp}}{\text{tp+tn}}$ 

•F beta Score: The weighted harmonic average of Precision and Recall.

•Support: The number of instances of each ground truth label.

Here, tp, fp, and tn stand for the number of true positives, false positives, and true negative, respectively.

label\_num specifies the number of classes, that is, each value in t must be an integer in the range of [0, label\_num). If label\_num is None, this function regards label\_num as a maximum of in t plus one.

ignore\_label determines which instances should be ignored. Specifically, instances with the given label are not taken into account for calculating the above quantities. By default, it is set to -1 so that all instances are taken into consideration, as labels are supposed to be non-negative integers. Setting ignore\_label to a non-negative integer less than label\_num is illegal and yields undefined behavior. In the current implementation, it arises RuntimeWarning and ignore\_label-th entries in output arrays do not contain correct quantities.

- y (Variable) Variable holding a vector of scores.
- t (Variable) Variable holding a vector of ground truth labels.
- label\_num (int) The number of classes.
- **beta** (*float*) The parameter which determines the weight of precision in the F-beta score.
- **ignore\_label** (*int*) Instances with this label are ignored.
- **Returns** 4-tuple of ~chainer.Variable of size (label\_num,). Each element represents precision, recall, F beta score, and support of this minibatch.

## r2\_score

chainer.functions.**r2\_score** (*pred*, *true*, *sample\_weight=None*, *multioutput='uniform\_average'*) Computes R^2(coefficient of determination) regression score function.

### Parameters

- pred (Variable) Variable holding a vector, matrix or tensor of estimated target values.
- **true** (Variable) Variable holding a vector, matrix or tensor of correct target values.
- **sample\_weight** This argument is for compatibility with scikit-learn's implementation of r2\_score. Current implementation admits None only.
- multioutput (*string*)-['uniform\_average', 'raw\_values']. if 'uniform\_average', this function returns an average of R^2 score of multiple output. If 'raw\_average', this function return a set of R^2 score of multiple output.
- **Returns** A Variable holding a scalar array of the R^2 score if 'multioutput' is 'uniform\_average' or a vector of R^2 scores if 'multioutput' is 'raw\_values'.

### **Return type** Variable

Note: This function is non-differentiable.

# Loss functions

## bernoulli\_nll

chainer.functions.bernoulli\_nll(x, y, reduce='sum')

Computes the negative log-likelihood of a Bernoulli distribution.

This function calculates the negative log-likelihood of a Bernoulli distribution.

$$-\log B(x;p) = -\sum_{i} \{x_i \log(p_i) + (1-x_i) \log(1-p_i)\},\$$

where  $p = \sigma(y), \sigma(\cdot)$  is a sigmoid function, and B(x; p) is a Bernoulli distribution.

The output is a variable whose value depends on the value of the option reduce. If it is 'no', it holds the elementwise loss values. If it is 'sum', loss values are summed up.

**Note:** As this function uses a sigmoid function, you can pass a result of fully-connected layer (that means Linear) to this function directly.

### Parameters

- **x** (*Variable* or numpy.ndarray or *cupy.ndarray*) Input variable.
- **y** (*Variable* or numpy.ndarray or *cupy.ndarray*) A variable representing the parameter of Bernoulli distribution.
- **reduce** (*str*) Reduction option. Its value must be either 'sum' or 'no'. Otherwise, ValueError is raised.
- **Returns** A variable representing the negative log-likelihood. If reduce is 'no', the output variable holds array whose shape is same as one of (hence both of) input variables. If it is 'sum', the output variable holds a scalar value.

**Return type** Variable

### black\_out

chainer.functions.black\_out (x, t, W, samples, reduce='mean')
BlackOut loss function.

BlackOut loss function is defined as

$$-\log(p(t)) - \sum_{s \in S} \log(1 - p(s)),$$

where t is the correct label, S is a set of negative examples and  $p(\cdot)$  is likelihood of a given label. And, p is defined as

$$p(y) = \frac{\exp(W_y^{\top} x)}{\sum_{s \in samples} \exp(W_s^{\top} x)}$$

The output is a variable whose value depends on the value of the option reduce. If it is 'no', it holds the no loss values. If it is 'mean', this function takes a mean of loss values.

### Parameters

- **x** (Variable) Batch of input vectors. Its shape should be (N, D).
- t (Variable) Vector of ground truth labels. Its shape should be (N, ). Each elements v should satisfy  $0 \ge v \ge V$  or -1 where V is the number of label types.
- W (Variable) Weight matrix. Its shape should be (V, D)
- **samples** (Variable) Negative samples. Its shape should be (N, S) where S is the number of negative samples.
- reduce (str) Reduction option. Its value must be either
- or 'mean'. Otherwise, ('no')-

:param ValueError is raised.:

**Returns** A variable object holding loss value(s). If reduce is 'no', the output variable holds an array whose shape is (N,). If it is 'mean', it holds a scalar.

#### **Return type** Variable

See: BlackOut: Speeding up Recurrent Neural Network Language Models With Very Large Vocabularies

### See also:

BlackOut.

## connectionist\_temporal\_classification

```
chainer.functions.connectionist_temporal_classification(x, t, blank_symbol,
input_length=None,
label_length=None, re-
duce='mean')
```

Connectionist Temporal Classification loss function.

Connectionist Temporal Classification(CTC) [Graves2006] is a loss function of sequence labeling where the alignment between the inputs and target is unknown. See also [Graves2012]

The output is a variable whose value depends on the value of the option reduce. If it is 'no', it holds the samplewise loss values. If it is 'mean', it takes the mean of loss values.

### Parameters

- **x** (sequence of Variable) RNN output at each time. x must be a list of Variable s. Each element of x, x[i] is a Variable representing output of RNN at time i.
- t (Variable) Expected label sequence.
- **blank\_symbol** (*int*) Index of blank\_symbol. This value must be non-negative.
- **input\_length** (Variable) Length of valid sequence for each of mini batch x (optional). If input\_length is skipped, It regards that all of x is valid input.
- **label\_length** (Variable) Length of valid sequence for each of mini batch t (optional). If label\_length is skipped, It regards that all of t is valid input.
- **reduce** (*str*) Reduction option. Its value must be either 'mean' or 'no'. Otherwise, ValueError is raised.
- **Returns** A variable holding a scalar value of the CTC loss. If reduce is 'no', the output variable holds array whose shape is (*B*,) where *B* is the number of samples. If it is 'mean', it holds a scalar.

Return type Variable

**Note:** You need to input x without applying to activation functions(e.g. softmax function), because this function applies softmax functions to x before calculating CTC loss to avoid numerical limitations. You also need to apply softmax function to forwarded values before you decode it.

**Note:** This function is differentiable only by x.

Note: This function supports (batch, sequence, 1-dimensional input)-data.

### contrastive

chainer.functions.contrastive(x0, x1, y, margin=1, reduce='mean')

Computes contrastive loss.

It takes a pair of samples and a label as inputs. The label is 1 when those samples are similar, or 0 when they are dissimilar.

Let N and K denote mini-batch size and the dimension of input variables, respectively. The shape of both input variables x0 and x1 should be (N, K). The loss value of the *n*-th sample pair  $L_n$  is

$$L_n = \frac{1}{2} \left( y_n d_n^2 + (1 - y_n) \max(\text{margin} - d_n, 0)^2 \right)$$

where  $d_n = \|\mathbf{x}_{0n} - \mathbf{x}_{1n}\|_2$ ,  $\mathbf{x}_{0n}$  and  $\mathbf{x}_{1n}$  are *n*-th K-dimensional vectors of x 0 and x 1.

The output is a variable whose value depends on the value of the option reduce. If it is 'no', it holds the elementwise loss values. If it is 'mean', this function takes a mean of loss values.

#### **Parameters**

- **x0** (Variable) The first input variable. The shape should be (N, K), where N denotes the mini-batch size, and K denotes the dimension of x0.
- **x1** (Variable) The second input variable. The shape should be the same as x0.
- **y** (Variable) Labels. All values should be 0 or 1. The shape should be (N, ), where N denotes the mini-batch size.
- margin (float) A parameter for contrastive loss. It should be positive value.
- **reduce** (*str*) Reduction option. Its value must be either 'mean' or 'no'. Otherwise, ValueError is raised.
- **Returns** A variable holding the loss value(s) calculated by the above equation. If reduce is 'no', the output variable holds array whose shape is same as one of (hence both of) input variables. If it is 'mean', the output variable holds a scalar value.

### Return type Variable

**Note:** This cost can be used to train siamese networks. See Learning a Similarity Metric Discriminatively, with Application to Face Verification for details.

# crf1d

chainer.functions.crfld(cost, xs, ys, reduce='mean')

Calculates negative log-likelihood of linear-chain CRF.

It takes a transition cost matrix, a sequence of costs, and a sequence of labels. Let  $c_{st}$  be a transition cost from a label s to a label t,  $x_{it}$  be a cost of a label t at position i, and  $y_i$  be an expected label at position i. The negative log-likelihood of linear-chain CRF is defined as

$$L = -\left(\sum_{i=1}^{l} x_{iy_i} + \sum_{i=1}^{l-1} c_{y_i y_{i+1}} - \log(Z)\right),$$

where l is the length of the input sequence and Z is the normalizing constant called partition function.

**Note:** When you want to calculate the negative log-likelihood of sequences which have different lengths, sort the sequences in descending order of lengths and transpose the sequences. For example, you have three input sequences:

```
>>> a1 = a2 = a3 = a4 = np.random.uniform(-1, 1, 3).astype('f')
>>> b1 = b2 = b3 = np.random.uniform(-1, 1, 3).astype('f')
>>> c1 = c2 = np.random.uniform(-1, 1, 3).astype('f')
```

>>> a = [a1, a2, a3, a4]
>>> b = [b1, b2, b3]
>>> c = [c1, c2]

where al and all other variables are arrays with (K, ) shape. Make a transpose of the sequences:

```
>>> x1 = np.stack([a1, b1, c1])
>>> x2 = np.stack([a2, b2, c2])
>>> x3 = np.stack([a3, b3])
>>> x4 = np.stack([a4])
```

and make a list of the arrays:

>>> xs = [x1, x2, x3, x4]

You need to make label sequences in the same fashion. And then, call the function:

```
>>> cost = chainer.Variable(
... np.random.uniform(-1, 1, (3, 3)).astype('f'))
>>> ys = [np.zeros(x.shape[0:1], dtype='i') for x in xs]
>>> loss = F.crfld(cost, xs, ys)
```

It calculates mean of the negative log-likelihood of the three sequences.

The output is a variable whose value depends on the value of the option reduce. If it is 'no', it holds the elementwise loss values. If it is 'mean', it holds mean of the loss values.

### **Parameters**

- **cost** (Variable) A  $K \times K$  matrix which holds transition cost between two labels, where K is the number of labels.
- **xs** (*list of Variable*) Input vector for each label. len (xs) denotes the length of the sequence, and each *Variable* holds a  $B \times K$  matrix, where B is mini-batch size, K is the number of labels. Note that B s in all the variables are not necessary the same, i.e., it accepts the input sequences with different lengths.
- **ys** (*list of Variable*) Expected output labels. It needs to have the same length as xs. Each *Variable* holds a *B* integer vector. When x in xs has the different *B*, correspoding y has the same *B*. In other words, ys must satisfy ys[i].shape == xs[i]. shape[0:1] for all i.
- **reduce** (*str*) Reduction option. Its value must be either 'mean' or 'no'. Otherwise, ValueError is raised.

### Returns

A variable holding the average negative log-likelihood of the input sequences.

Return type Variable

**Note:** See detail in the original paper: Conditional Random Fields: Probabilistic Models for Segmenting and Labeling Sequence Data.

```
chainer.functions.argmax_crfld(cost, xs)
```

Computes a state that maximizes a joint probability of the given CRF.

### Parameters

- **cost** (Variable) A  $K \times K$  matrix which holds transition cost between two labels, where K is the number of labels.
- **xs** (*list of Variable*) Input vector for each label. len (xs) denotes the length of the sequence, and each *Variable* holds a  $B \times K$  matrix, where B is mini-batch size, K is the number of labels. Note that B s in all the variables are not necessary the same, i.e., it accepts the input sequences with different lengths.

# Returns

A tuple of Variable object s and a list ps. The shape of s is (B,), where B is the mini-batch size. i-th element of s, s[i], represents log-likelihood of i-th data. ps is a list of numpy.ndarray or *cupy.ndarray*, and denotes the state that maximizes the point probability. len(ps) is equal to len(xs), and shape of each ps[i] is the mini-batch size of the corresponding xs[i]. That means, ps[i].shape == xs[i]. shape[0:1].

### Return type tuple

### cross\_covariance

```
chainer.functions.cross_covariance (y, z, reduce='half_squared_sum')
Computes the sum-squared cross-covariance penalty between y and z
```

Computes the sum-squared cross-covariance penalty between  $\ensuremath{\mathtt{y}}$  and  $\ensuremath{\mathtt{z}}$ 

The output is a variable whose value depends on the value of the option reduce. If it is 'no', it holds the covariant matrix that has as many rows (resp. columns) as the dimension of y (resp.z). If it is 'half\_squared\_sum', it holds the half of the Frobenius norm (i.e. L2 norm of a matrix flattened to a vector) of the covarianct matrix.

#### Parameters

- **y** (Variable) Variable holding a matrix where the first dimension corresponds to the batches.
- **z** (Variable) Variable holding a matrix where the first dimension corresponds to the batches.
- **reduce** (*str*) Reduction option. Its value must be either 'half\_squared\_sum' or 'no'. Otherwise, ValueError is raised.
- Returns A variable holding the cross covariance loss. If reduce is 'no', the output variable holds 2-dimensional array matrix of shape (M, N) where M (resp. N) is the number of columns of y (resp. z). If it is 'half\_squared\_sum', the output variable holds a scalar value.

Return type Variable

Note: This cost can be used to disentangle variables. See https://arxiv.org/abs/1412.6583v3 for details.

# gaussian\_kl\_divergence

```
chainer.functions.gaussian_kl_divergence (mean, ln_var, reduce='sum')
Computes the KL-divergence of Gaussian variables from the standard one.
```

Given two variable mean representing  $\mu$  and  $\ln_var$  representing  $\log(\sigma^2)$ , this function calculates the KLdivergence in elementwise manner between the given multi-dimensional Gaussian  $N(\mu, S)$  and the standard Gaussian N(0, I)

$$D_{\mathbf{KL}}(N(\mu, S) \parallel N(0, I)),$$

where S is a diagonal matrix such that  $S_{ii} = \sigma_i^2$  and I is an identity matrix.

The output is a variable whose value depends on the value of the option reduce. If it is 'no', it holds the elementwise loss values. If it is 'sum', loss values are summed up.

#### **Parameters**

- mean (Variable or numpy.ndarray or cupy.ndarray) A variable representing mean of given gaussian distribution, μ.
- **ln\_var** (*Variable* or numpy.ndarray or *cupy.ndarray*) A variable representing logarithm of variance of given gaussian distribution,  $\log(\sigma^2)$ .
- **reduce** (*str*) Reduction option. Its value must be either 'sum' or 'no'. Otherwise, ValueError is raised.
- **Returns** A variable representing KL-divergence between given gaussian distribution and the standard gaussian. If reduce is 'no', the output variable holds array whose shape is same as one of (hence both of) input variables. If it is 'sum', the output variable holds a scalar value.

**Return type** Variable

### gaussian\_nll

chainer.functions.gaussian\_nll(x, mean, ln\_var, reduce='sum')

Computes the negative log-likelihood of a Gaussian distribution.

Given two variable mean representing  $\mu$  and  $ln_var$  representing  $log(\sigma^2)$ , this function computes in elementwise manner the negative log-likelihood of x on a Gaussianx distribution  $N(\mu, S)$ ,

$$-\log N(x;\mu,\sigma^2) = \log\left(\sqrt{(2\pi)^D |S|}\right) + \frac{1}{2}(x-\mu)^\top S^{-1}(x-\mu),$$

where D is a dimension of x and S is a diagonal matrix where  $S_{ii} = \sigma_i^2$ .

The output is a variable whose value depends on the value of the option reduce. If it is 'no', it holds the elementwise loss values. If it is 'sum', loss values are summed up.

#### **Parameters**

- **x** (*Variable* or numpy.ndarray or *cupy.ndarray*) Input variable.
- mean (Variable or numpy.ndarray or cupy.ndarray) A variable representing mean of a Gaussian distribution, μ.
- $\ln_{var}(Variable \text{ or numpy.ndarray or } cupy.ndarray) A variable representing logarithm of variance of a Gaussian distribution, <math>\log(\sigma^2)$ .
- **reduce** (*str*) Reduction option. Its value must be either 'sum' or 'no'. Otherwise, ValueError is raised.
- **Returns** A variable representing the negative log-likelihood. If reduce is 'no', the output variable holds array whose shape is same as one of (hence both of) input variables. If it is 'sum', the output variable holds a scalar value.

### Return type Variable

### hinge

chainer.functions.hinge (x, t, norm='L1', reduce='mean')
Computes the hinge loss for a one-of-many classification task.

$$L = \frac{1}{N} \sum_{n=1}^{N} \sum_{k=1}^{K} \left[ \max(0, 1 - \delta \{ t_n = k \} x_{nk}) \right]^p$$

where N denotes the batch size and K is the number of classes of interest,

$$\delta\{\text{condition}\} = \begin{cases} 1 & \text{if condition is true} \\ -1 & \text{otherwise,} \end{cases}$$

and

$$p = \begin{cases} 1 & \text{if norm} = L1\\ 2 & \text{if norm} = L2 \end{cases}$$

The output is a variable whose value depends on the value of the option reduce. If it is 'no', it holds the elementwise loss values. If it is 'mean', it takes the mean of loss values.

### Parameters

- **x** (Variable) Input variable. The shape of x should be (N, K).
- t (Variable) The N-dimensional label vector with values  $t_n \in \{0, 1, 2, ..., K-1\}$ . The shape of t should be (N,).
- **norm** (*string*) Specifies norm type. Either 'L1' or 'L2' is acceptable.
- **reduce** (*str*) Reduction option. Its value must be either 'mean' or 'no'. Otherwise, ValueError is raised.
- **Returns** A variable object holding a scalar array of the hinge loss L. If reduce is 'no', the output variable holds array whose shape is same as one of (hence both of) input variables. If it is 'mean', the output variable holds a scalar value.

**Return type** Variable

# huber\_loss

chainer.functions.huber\_loss (*x*, *t*, *delta*, *reduce='sum\_along\_second\_axis'*) Loss function which is less sensitive to outliers in data than MSE.

$$a = x - t$$

and

$$L_{\delta}(a) = \begin{cases} \frac{1}{2}a^2 & \text{if } |\mathbf{a}| \le \delta\\ \delta(|a| - \frac{1}{2}\delta) & \text{otherwise,} \end{cases}$$

The output is a variable whose value depends on the value of the option reduce. If it is 'no', it holds the elementwise loss values. If it is 'sum\_along\_second\_axis', loss values are summed up along the second axis (i.e. axis=1).

- **x** (Variable) Input variable. The shape of x should be (N, K).
- t (Variable) Target variable for regression. The shape of t should be (N, K).
- delta (float) Constant variable for huber loss function as used in definition.
- **reduce** (*str*) Reduction option. Its value must be either 'sum\_along\_second\_axis' or 'no'. Otherwise, ValueError is raised.
- **Returns** A variable object holding a scalar array of the huber loss  $L_{\delta}$ . If reduce is 'no', the output variable holds array whose shape is same as one of (hence both of) input variables. If it is 'sum\_along\_second\_axis', the shape of the array is same as the input variables, except the second axis is removed.

### **Return type** Variable

See: Huber loss - Wikipedia.

### mean\_absolute\_error

chainer.functions.mean\_absolute\_error(x0, x1)

Mean absolute error function.

This function computes mean absolute error between two variables. The mean is taken over the minibatch.

### mean\_squared\_error

```
chainer.functions.mean_squared_error(x0, x1)
```

Mean squared error function.

This function computes mean squared error between two variables. The mean is taken over the minibatch. Note that the error is not scaled by 1/2.

### negative\_sampling

chainer.functions.negative\_sampling (x, t, W, sampler, sample\_size, reduce='sum') Negative sampling loss function.

In natural language processing, especially language modeling, the number of words in a vocabulary can be very large. Therefore, you need to spend a lot of time calculating the gradient of the embedding matrix.

By using the negative sampling trick you only need to calculate the gradient for a few sampled negative examples.

The objective function is below:

$$f(x,p) = \log \sigma(x^{\top} w_p) + k E_{i \sim P(i)} [\log \sigma(-x^{\top} w_i)],$$

where  $\sigma(\cdot)$  is a sigmoid function,  $w_i$  is the weight vector for the word *i*, and *p* is a positive example. It is approximated with *k* examples *N* sampled from probability P(i), like this:

$$f(x,p) \approx \log \sigma(x^{\top} w_p) + \sum_{n \in N} \log \sigma(-x^{\top} w_n).$$

Each sample of N is drawn from the word distribution P(w). This is calculated as  $P(w) = \frac{1}{Z}c(w)^{\alpha}$ , where c(w) is the unigram count of the word w,  $\alpha$  is a hyper-parameter, and Z is the normalization constant.

- **x** (Variable) Batch of input vectors.
- t (Variable) Vector of ground truth labels.
- W (Variable) Weight matrix.
- **sampler** (*FunctionType*) Sampling function. It takes a shape and returns an integer array of the shape. Each element of this array is a sample from the word distribution. A *WalkerAlias* object built with the power distribution of word frequency is recommended.
- **sample\_size** (*int*) Number of samples.
- **reduce** (*str*) Reduction option. Its value must be either 'sum' or 'no'. Otherwise, ValueError is raised.
- **Returns** A variable holding the loss value(s) calculated by the above equation. If reduce is 'no', the output variable holds array whose shape is same as one of (hence both of) input variables. If it is 'sum', the output variable holds a scalar value.

## Return type Variable

See: Distributed Representations of Words and Phrases and their Compositionality

### See also:

NegativeSampling.

### sigmoid\_cross\_entropy

chainer.functions.**sigmoid\_cross\_entropy**(*x*, *t*, *use\_cudnn=True*, *normalize=True*, *reduce='mean'*)

Computes cross entropy loss for pre-sigmoid activations.

## Parameters

- **x** (Variable) A variable object holding a matrix whose (i, j)-th element indicates the unnormalized log probability of the j-th unit at the i-th example.
- t (Variable) Variable holding an int32 vector of ground truth labels. If t [i] == -1, corresponding x [i] is ignored. Loss is zero if all ground truth labels are -1.
- **normalize** (*bool*) Variable holding a boolean value which determines the normalization constant. If true, this function normalizes the cross entropy loss across all instances. If else, it only normalizes along a batch size.
- **reduce** (*str*) Variable holding a *str* which determines whether to reduce the shape of the input. If it is 'mean', it computes the sum of cross entropy and normalize it according to normalize option. If is is 'no', this function computes cross entropy for each instance and does not normalize it (normalize option is ignored). In this case, the loss value of the ignored instance, which has -1 as its target value, is set to 0.
- **Returns** A variable object holding an array of the cross entropy. If reduce is 'mean', it is a scalar array. If reduce is 'no', the shape is same as x.

**Return type** Variable

**Note:** This function is differentiable only by x.

### softmax\_cross\_entropy

Computes cross entropy loss for pre-softmax activations.

#### Parameters

- **x** (Variable) Variable holding a multidimensional array whose element indicates unnormalized log probability: the first axis of the variable represents the number of samples, and the second axis represents the number of classes. While this function computes a usual softmax cross entropy if the number of dimensions is equal to 2, it computes a cross entropy of the replicated softmax if the number of dimensions is greater than 2.
- t (Variable) Variable holding an int32 vector of ground truth labels. If t[i] == ignore\_label, corresponding x[i] is ignored.
- **normalize** (*bool*) If True, this function normalizes the cross entropy loss across all instances. If False, it only normalizes along a batch size.
- **cache\_score** (bool) When it is True, the function stores result of forward computation to use it on backward computation. It reduces computational cost though consumes more memory.
- **class\_weight** (*ndarray* or *ndarray*) An array that contains constant weights that will be multiplied with the loss values along with the second dimension. The shape of this array should be (x.shape[1],). If this is not None, each class weight class\_weight[i] is actually multiplied to y[:, i] that is the corresponding logsoftmax output of x and has the same shape as x before calculating the actual loss value.
- **ignore\_label** (*int*) Label value you want to ignore. Its default value is -1. See description of the argument *t*.
- **reduce** (*str*) A string that determines whether to reduce the loss values. If it is 'mean', it computes the sum of the individual cross entropy and normalize it according to normalize option. If it is 'no', this function computes cross entropy for each instance and does not normalize it (normalize option is ignored). In this case, the loss value of the ignored instance, which has ignore\_label as its target value, is set to 0.
- **Returns** A variable holding a scalar array of the cross entropy loss. If reduce is 'mean', it is a scalar array. If reduce is 'no', the shape is same as that of x.

### Return type Variable

**Note:** This function is differentiable only by x.

# triplet

chainer.functions.triplet (anchor, positive, negative, margin=0.2, reduce='mean')
Computes triplet loss.

It takes a triplet of variables as inputs, a, p and n: anchor, positive example and negative example respectively. The triplet defines a relative similarity between samples. Let N and K denote mini-batch size and the dimension of input variables, respectively. The shape of all input variables should be (N, K).

$$L(a, p, n) = \frac{1}{N} \left( \sum_{i=1}^{N} \max\{d(a_i, p_i) - d(a_i, n_i) + \text{margin}, 0\} \right)$$

where  $d(x_i, y_i) = \|\mathbf{x}_i - \mathbf{y}_i\|_2^2$ .

The output is a variable whose value depends on the value of the option reduce. If it is 'no', it holds the elementwise loss values. If it is 'mean', this function takes a mean of loss values.

### **Parameters**

- **anchor** (Variable) The anchor example variable. The shape should be (N, K), where N denotes the minibatch size, and K denotes the dimension of the anchor.
- **positive** (Variable) The positive example variable. The shape should be the same as anchor.
- **negative** (Variable) The negative example variable. The shape should be the same as anchor.
- margin (float) A parameter for triplet loss. It should be a positive value.
- **reduce** (*str*) Reduction option. Its value must be either 'mean' or 'no'. Otherwise, ValueError is raised.
- **Returns** A variable holding a scalar that is the loss value calculated by the above equation. If reduce is 'no', the output variable holds array whose shape is same as one of (hence both of) input variables. If it is 'mean', the output variable holds a scalar value.

## Return type Variable

**Note:** This cost can be used to train triplet networks. See Learning Fine-grained Image Similarity with Deep Ranking for details.

# **Mathematical functions**

## arccos

```
chainer.functions.arccos(x)
Elementwise arccosine function.
```

 $y_i = \arccos x_i.$ 

Parameters x (Variable) – Input variable.

Returns Output variable.

Return type Variable

# arcsin

chainer.functions.**arcsin**(*x*) Elementwise arcsine function.

 $y_i = \arcsin x_i.$ 

**Parameters x** (Variable) – Input variable.

**Returns** Output variable.

Return type Variable

# arctan

chainer.functions.**arctan**(x) Elementwise arctangent function.

 $y_i = \arctan x_i.$ 

Parameters x (Variable) - Input variable.

**Returns** Output variable.

**Return type** *Variable* 

# argmax

chainer.functions.**argmax** (*x*, *axis=None*) Returns index which holds maximum of array elements over a given axis.

### **Parameters**

- **x** (Variable) Array to find maximum elements.
- **axis** (*None or int*) Axis over which a max is performed. The default (axis = None) is perform a max over all the dimensions of the input array.

Returns Output variable.

Return type Variable

# argmin

chainer.functions.argmin(x, axis=None)

Returns index which holds minimum of array elements over a given axis.

### Parameters

- **x** (Variable) Array to find minimum elements.
- **axis** (*None or int*) Axis over which a min is performed. The default (axis = None) is perform a min over all the dimensions of the input array.

Returns Output variable.

Return type Variable

# average

chainer.functions.average (x, axis=None, weights=None, keepdims=False) Calculate weighted average of array elements over a given axis.

### **Parameters**

• **x** (Variable) – Elements to sum.

- **axis** (*None or int*) Axis which the method is performed. With the default (axis = None) it performs a mean over all the dimensions of the input array.
- weights (*None or* chainer.Variable) An array holding weights to calculate weighted average. If it is None, all weights are assumed to be one. When axis is None, weights must have the same shape of x. And when axis is int, it must be 1-D array satisfing weights.shape == (x.shape[axis],).
- keepdims (bool) If True, the specified axes are remained as axes of length one.

Returns Output variable.

Return type Variable

## batch\_inv

chainer.functions.batch\_inv(a)

Computes the inverse of a batch of square matrices.

**Parameters a** (Variable) – Input array to compute the inverse for. Shape of the array should be (m, n, n) where m is the number of matrices in the batch, and n is the dimensionality of a square matrix.

Returns Inverse of every matrix in the batch of matrices.

**Return type** Variable

## batch\_l2\_norm\_squared

chainer.functions.batch\_12\_norm\_squared(x)

L2 norm (a.k.a. Euclidean norm) squared.

This function implements the square of L2 norm on a vector. No reduction along batch axis is done.

**Parameters x** (Variable) – Input variable. The first dimension is assumed to be the *minibatch dimension*. If x has more than two dimensions all but the first dimension are flattened to one dimension.

Returns Two dimensional output variable.

Return type Variable

## batch\_matmul

chainer.functions.**batch\_matmul** (*a*, *b*, *transa=False*, *transb=False*) Computes the batch matrix multiplications of two sets of arrays.

- a (Variable) The left operand of the batch matrix multiplications. A 2-D array of shape (B, N) is considered as B  $N \times 1$  matrices. A 3-D array of shape (B, M, N) is considered as B  $M \times N$  matrices.
- **b** (Variable) The right operand of the batch matrix multiplications. Its array is treated as matrices in the same way as a's array.
- transa (bool) If True, transpose each matrix in a.
- transb (bool) If True, transpose each matrix in b.

### Returns

The result of the batch matrix multiplications as a 3-D array.

Return type Variable

# bias

```
chainer.functions.bias(x, y, axis=1)
```

Elementwise summation with broadcasting.

Computes a elementwise summation of two input variables, with the shape of the latter variable broadcasted to match the shape of the former. axis is the first axis of the first variable along which the second variable is applied.

The term "broadcasting" here comes from Caffe's bias layer so the "broadcasting" with the following arguments:

x : 100 x 3 x 40 x 60 y : 3 x 40 axis : 1

is equivalent to the following numpy broadcasting:

x : 100 x 3 x 40 x 60 y : 1 x 3 x 40 x 1

Note that how the axis indicates to which axis of x we apply y.

### Parameters

- **x** (Variable) Input variable to be summed.
- **y** (Variable) Input variable to sum, broadcasted.
- **axis** (*int*) The first axis of x along which y is applied.

Returns Output variable.

Return type Variable

## ceil

chainer.functions.**ceil**(*x*) Elementwise ceil function.

 $y_i = \lceil x_i \rceil$ 

Parameters x (Variable) - Input variable.

Returns Output variable.

Return type Variable

# clip

chainer.functions.**clip** (*x*, *x\_min*, *x\_max*) Clips (limits) elements of input variable.

Given an interval [x\_min, xmax], elements outside the interval are clipped to the interval edges.

## Parameters

- **x** (Variable) Input variable to be clipped.
- **x\_min** (*float*) Minimum value.
- **x\_max** (float) Maximum value.

Returns Output variable.

Return type Variable

### cos

```
chainer.functions.cos(x)
Elementwise cos function.
```

# cosh

chainer.functions.cosh(x)
 Elementwise hyperbolic cosine function.

 $y_i = \cosh x_i.$ 

Parameters x (Variable) – Input variable.

Returns Output variable.

Return type Variable

# ехр

chainer.functions.**exp**(*x*) Elementwise exponential function.

# fmod

```
chainer.functions.fmod(x, divisor)
Elementwise mod function.
```

 $y_i = x_i \mod \text{divisor.}$ 

### Parameters

- **x** (Variable) Input variable.
- divisor (Variable) Input divisor.

Returns Output variable.

Return type Variable

# floor

```
chainer.functions.floor(x)
Elementwise floor function.
```

 $y_i = \lfloor x_i \rfloor$ 

Parameters x (Variable) - Input variable.

Returns Output variable.

Return type Variable

# identity

chainer.functions.identity(\*inputs)
 Just returns input variables.

# inv

chainer.functions.**inv**(*a*) Computes the inverse of square matrix.

**Parameters a** (Variable) – Input array to compute the inverse for. Shape of the array should be (n, n) where n is the dimensionality of a square matrix.

Returns Matrix inverse of a.

Return type Variable

# linear\_interpolate

chainer.functions.linear\_interpolate(*p*, *x*, *y*) Elementwise linear-interpolation function.

This function is defined as

$$f(p, x, y) = px + (1 - p)y.$$

# Parameters

- **p** (Variable) Input variable.
- **x** (Variable) Input variable.
- **y** (Variable) Input variable.

Returns Output variable.

Return type Variable

# log

chainer.functions.log(x) Elementwise natural logarithm function.

# log10

```
chainer.functions.log10(x)
Elementwise logarithm function to the base 10.
```

 $y_i = \log_{10} x_i.$ 

Parameters x (Variable) - Input variable.

Returns Output variable.

Return type Variable

# log1p

chainer.functions.log1p(x) Elementwise natural logarithm plus one function.

# log2

chainer.functions.**log2** (*x*) Elementwise logarithm function to the base 2.

 $y_i = \log_2 x_i.$ 

**Parameters x** (Variable) – Input variable.

Returns Output variable.

**Return type** *Variable* 

# logsumexp

```
chainer.functions.logsumexp(x, axis=None)
```

Log-sum-exp of array elements over a given axis.

This function calculates logarithm of sum of exponential of array elements.

$$y_i = \log\left(\sum_j \exp(x_{ij})\right)$$

## Parameters

- **x** (Variable) Elements to log-sum-exp.
- **axis** (*None*, *int*, *or tuple of int*) Axis which a sum is performed. The default (axis = None) is perform a sum over all the dimensions of the input array.

Returns Output variable.

## Return type Variable

# matmul

chainer.functions.**matmul** (*a*, *b*, *transa=False*, *transb=False*) Computes the matrix multiplication of two arrays.

## Parameters

- **a** (Variable) The left operand of the matrix multiplication. A 1-D array of shape (N, ) is considered as an  $N \times 1$  matrix. A 2-D array of shape (M, N) is considered as an  $M \times N$  matrix.
- **b** (Variable) The right operand of the matrix multiplication. Its array is treated as a matrix in the same way as a's array.
- transa (bool) If True, transpose a.
- transb (bool) If True, transpose b.

### Returns

The result of the matrix multiplication as a 2-D array.

Return type Variable

### max

chainer.functions.**max** (*x*, *axis=None*, *keepdims=False*) Maximum of array elements over a given axis.

## Parameters

- **x** (Variable) Array to be maximized.
- **axis** (*None*, *int*, *or tuple of int*) Axis over which a max is performed. The default (axis = None) is perform a max over all the dimensions of the input array.

Returns Output variable.

Return type Variable

### maximum

```
chainer.functions.maximum(x1,x2)
```

Element-wise maximum of input variables.

### **Parameters**

- **x1** (Variable) Input variables to be compared.
- **x2** (Variable) Input variables to be compared.

Returns Output variable.

Return type Variable

### mean

chainer.functions.mean (x, axis=None, weights=None, keepdims=False) Calculate weighted average of array elements over a given axis.

- **x** (Variable) Elements to sum.
- **axis** (*None or int*) Axis which the method is performed. With the default (axis = None) it performs a mean over all the dimensions of the input array.
- weights (*None or* chainer.Variable) An array holding weights to calculate weighted average. If it is None, all weights are assumed to be one. When axis is None, weights must have the same shape of x. And when axis is int, it must be 1-D array satisfing weights.shape == (x.shape[axis],).
- **keepdims** (*bool*) If True, the specified axes are remained as axes of length one.

Returns Output variable.

**Return type** *Variable* 

## min

chainer.functions.min (x, axis=None, keepdims=False) Minimum of array elements over a given axis.

### **Parameters**

- **x** (Variable) Array to be minimized.
- **axis** (*None*, *int*, *or tuple of int*) Axis over which a min is performed. The default (axis = None) is perform a min over all the dimensions of the input array.

Returns Output variable.

**Return type** Variable

## minimum

chainer.functions.minimum (x1, x2) Element-wise minimum of input variables.

### **Parameters**

- **x1** (Variable) Input variables to be compared.
- **x2** (Variable) Input variables to be compared.

Returns Output variable.

**Return type** Variable

### rsqrt

```
chainer.functions.rsqrt(x)
```

Computes elementwise reciprocal of square root of input  $x_i$ .

$$y_i = \frac{1}{\sqrt{x_i}}.$$

Parameters x (Variable) – Input variable.

Returns Output variable.

Return type Variable

See also:

sqrt()

### scale

```
chainer.functions.scale (x, y, axis=1)
```

Elementwise product with broadcasting.

Computes a elementwise product of two input variables, with the shape of the latter variable broadcasted to match the shape of the former. axis is the first axis of the first variable along which the second variable is applied.

The term "broadcasting" here comes from Caffe's scale layer so the "broadcasting" with the following arguments:

x : 100 x 3 x 40 x 60 y : 3 x 40 axis : 1

is equivalent to the following numpy broadcasting:

x : 100 x 3 x 40 x 60 y : 1 x 3 x 40 x 1

Note that how the axis indicates to which axis of x we apply y.

### Parameters

- **x** (Variable) Input variable to be scaled.
- y (Variable) Input variable to scale, broadcasted.
- **axis** (*int*) The first axis of x along which y is applied.

Returns Output variable.

Return type Variable

# sin

```
chainer.functions.sin(x)
    Elementwise sin function.
```

# sinh

chainer.functions.sinh(x)
 Elementwise hyperbolic sine function.

 $y_i = \sinh x_i.$ 

Parameters x (Variable) - Input variable.
Returns Output variable.
Return type Variable

### sqrt

```
chainer.functions.sqrt(x)
    Elementwise square root function.
```

 $y_i = \sqrt{x_i}.$ 

If the value of  $x_i$  is negative, it returns Nan for  $y_i$  respect to underlying numpy and cupy specification.

**Parameters x** (Variable) – Input variable.

Returns Output variable.

Return type Variable

# square

chainer.functions.**square**(x) Elementwise square function.

 $y_i = x_i^2$ .

**Parameters** x (chainer.Variable or numpy.ndarray or cupy.ndarray) – Input variable.

Returns Output variable.

Return type Variable

### squared\_difference

```
chainer.functions.squared_difference (x1, x2)
Squared difference of input variables.
```

**Parameters** 

- **x1** (Variable) Input variables to be compared.
- **x2** (Variable) Input variables to be compared.

**Returns** (x1 - x2) + 2 element-wise.

**Return type** *Variable* 

### sum

chainer.functions.**sum**(*x*, *axis=None*, *keepdims=False*) Sum of array elements over a given axis.

## **Parameters**

- **x** (Variable) Elements to sum.
- **axis** (*None*, *int*, *or tuple of int*) Axis which a sum is performed. The default (axis = None) is perform a sum over all the dimensions of the input array.
- **keepdims** (*bool*) If True, the specified axes are remained as axes of length one.

Returns Output variable.

**Return type** *Variable* 

# tanh

Hyperbolic tangent function is described in "Activation functions" section.

See also:

tanh()

### tan

chainer.functions.**tan**(*x*) Elementwise tan function.

# Noise injections

### dropout

```
chainer.functions.dropout(x, ratio=0.5, train=True)
```

Drops elements of input variable randomly.

This function drops input elements randomly with probability ratio and scales the remaining elements by factor 1 / (1 - ratio). In testing mode, it does nothing and just returns x.

## **Parameters**

- **x** (Variable) Input variable.
- **ratio** (*float*) Dropout ratio.
- train (bool) If True, executes dropout. Otherwise, does nothing.

Returns Output variable.

## Return type Variable

See the paper by G. Hinton: Improving neural networks by preventing co-adaptation of feature detectors.

# gaussian

```
chainer.functions.gaussian(mean, ln_var)
```

Gaussian sampling function.

It takes mean  $\mu$  and logarithm of variance  $\log(\sigma^2)$  as input and output a sample drawn from gaussian  $N(\mu, \sigma)$ .

## **Parameters**

- **mean** (Variable) Input variable representing mean μ.
- ln\_var (Variable) Input variable representing logarithm of variance  $log(\sigma^2)$ .

Returns Output variable.

Return type Variable

## simplified\_dropconnect

chainer.functions.simplified\_dropconnect(x, W, b=None, ratio=0.5, train=True, mask=None)

Linear unit regularized by simplified dropconnect.

Simplified dropconnect drops weight matrix elements randomly with probability ratio and scales the remaining elements by factor 1 / (1 - ratio). Which element is dropped depends on each sample. It accepts two or three arguments: an input minibatch x, a weight matrix W, and optionally a bias vector b. It computes  $Y = xW^{\top} + b$ .

In testing mode, zero will be used as simplified dropconnect ratio instead of ratio.

Notice: This implementation cannot be used for reproduction of the paper. There is a difference between the current implementation and the original one. The original version uses sampling with gaussian distribution before passing activation function, whereas the current implementation averages before activation.

### **Parameters**

- **x** (chainer.Variable or numpy.ndarray or cupy.ndarray) Input variable. Its first dimension n is assumed to be the *minibatch dimension*. The other dimensions are treated as concatenated one dimension whose size must be N.
- W (Variable) Weight variable of shape (M, N).
- **b** (Variable) Bias variable (optional) of shape (M,).
- **ratio** (*float*) Dropconnect ratio.
- **train** (*bool*) If True, executes simplified dropconnect. Otherwise, simplified dropconnect function works as a linear function.
- **:param mask (None or chainer.Variable or numpy.ndarray or: cupy.ndarray):** If None, randomized dropconnect mask is generated. Otherwise, The mask must be (n, M, N) shaped array. Main purpose of this option is debugging. *mask* array will be used as a dropconnect mask.

Returns Output variable.

## Return type Variable

### See also:

### Dropconnect

### See also:

Li, W., Matthew Z., Sixin Z., Yann L., Rob F. (2013). Regularization of Neural Network using DropConnect. International Conference on Machine Learning. URL

# Normalization functions

## batch\_normalization

chainer.functions.batch\_normalization(x, gamma, beta, eps=2e-05, running\_mean=None,

running\_var=None, decay=0.9, use\_cudnn=True)

Batch normalization function.

It takes the input variable x and two parameter variables gamma and beta. The parameter variables must both have the same dimensionality, which is referred to as the channel shape. This channel shape corresponds to the dimensions in the input which are not averaged over. Since the first dimension of the input corresponds

to the batch size, the second dimension of x will correspond to the first dimension of the channel shape, the third dimension of x will correspond to the second channel dimension (if it exists) and so on. Therefore, the dimensionality of the input must be at least one plus the number of channel dimensions. The total effective "batch size" will then be considered to be the product of all dimensions in x except for the channel dimensions.

As an example, if the input is four dimensional and the parameter variables are one dimensional, then it is assumed that the first dimension of the input is the batch size, the second dimension is the channel size, and the remaining two dimensions are considered to be spatial dimensions that will be averaged over along with the batch size in the batch normalization computations. That is, the total batch size will be considered to be the product of all input dimensions except the second dimension.

Note: If this function is called, it will not be possible to access the updated running mean and variance statistics, because they are members of the function object, which cannot be accessed by the caller. If it is desired to access the updated running statistics, it is necessary to get a new instance of the function object, call the object, and then access the running\_mean and/or running\_var attributes. See the corresponding Link class for an example of how to do this.

## Parameters

- **x** (Variable) Input variable.
- gamma (Variable) Scaling parameter of normalized data.
- beta (Variable) Shifting parameter of scaled normalized data.
- **eps** (*float*) Epsilon value for numerical stability.
- running\_mean (array) Running average of the mean. This is a running average of the mean over several mini-batches using the decay parameter. If None, the running average is not computed. If this is None, then running\_var must also be None.
- **running\_var** (array) Running average of the variance. This is a running average of the variance over several mini-batches using the decay parameter. If None, the running average is not computed. If this is None, then running\_mean must also be None.
- decay (float) Decay rate of moving average. It is used during training.
- **use\_cudnn** (*bool*) If True and cuDNN is enabled, then this function uses cuDNN as the core implementation.

See: Batch Normalization: Accelerating Deep Network Training by Reducing Internal Covariate Shift

### See also:

links.BatchNormalization

## fixed\_batch\_normalization

Batch normalization function with fixed statistics.

This is a variant of batch normalization, where the mean and variance statistics are given by the caller as fixed variables. This is used on testing mode of the batch normalization layer, where batch statistics cannot be used for prediction consistency.

- **x** (Variable) Input variable.
- gamma (Variable) Scaling parameter of normalized data.
- **beta** (Variable) Shifting parameter of scaled normalized data.

- mean (Variable) Shifting parameter of input.
- **var** (Variable) Square of scaling parameter of input.
- **eps** (*float*) Epsilon value for numerical stability.
- **use\_cudnn** (*bool*) If True and cuDNN is enabled, then this function uses cuDNN as the core implementation.

## See also:

functions.batch\_normalization(), links.BatchNormalization

### local\_response\_normalization

chainer.functions.local\_response\_normalization (x, n=5, k=2, alpha=0.0001, beta=0.75) Local response normalization across neighboring channels.

This function implements normalization across channels. Let x an input image with N channels. Then, this function computes an output image y by following formula:

$$y_{i} = \frac{x_{i}}{\left(k + \alpha \sum_{j=\max 1, i-n/2}^{\min N, i+n/2} x_{j}^{2}\right)^{\beta}}$$

### **Parameters**

- **x** (Variable) Input variable.
- **n** (*int*) Normalization window width.
- **k** (float) Smoothing parameter.
- **alpha** (*float*) Normalizer scaling parameter.
- **beta** (*float*) Normalizer power parameter.

Returns Output variable.

Return type Variable

See: Section 3.3 of ImageNet Classification with Deep Convolutional Neural Networks

# normalize

```
chainer.functions.normalize(x, eps=1e-05, axis=1)
```

L2 norm squared (a.k.a. Euclidean norm).

This function implements L2 normalization on a vector along the given axis. No reduction is done along the normalization axis.

In the case when axis=1 and x is a vector of dimension (N, K), where N and K denote mini-batch size and the dimension of the input variable, this function computes an output vector y by the following equation:

$$y_i = \frac{x_i}{\|x_i\|_2 + \epsilon}$$

eps is used to avoid division by zero when norm of x along the given axis is zero.

The default value of axis is determined for backward compatibility.

- **x** (Variable) Two dimensional output variable. The first dimension is assumed to be the mini-batch dimension.
- **eps** (*float*) Epsilon value for numerical stability.
- **axis** (*int*) Axis along which to normalize.

**Returns** The output variable which has the same shape as x.

Return type Variable

# **Spatial pooling**

# average\_pooling\_2d

chainer.functions.average\_pooling\_2d(x, ksize, stride=None, pad=0, use\_cudnn=True) Spatial average pooling function.

This function acts similarly to Convolution2D, but it computes the average of input spatial patch for each channel without any parameter instead of computing the inner products.

### **Parameters**

- **x** (Variable) Input variable.
- **ksize** (*int or pair of ints*) Size of pooling window. ksize=k and ksize=(k, k) are equivalent.
- **stride** (*int or pair of ints or None*) Stride of pooling applications. stride=s and stride=(s, s) are equivalent. If None is specified, then it uses same stride as the pooling window size.
- **pad** (*int or pair of ints*) Spatial padding width for the input array. pad=p and pad=(p, p) are equivalent.
- **use\_cudnn** (*bool*) If True and cuDNN is enabled, then this function uses cuDNN as the core implementation.

Returns Output variable.

Return type Variable

**Note:** This function currently does not support cover\_all mode as max\_pooling\_2d(). Average pooling runs in non-cover-all mode.

# average\_pooling\_nd

chainer.functions.average\_pooling\_nd(x, ksize, stride=None, pad=0, use\_cudnn=True) N-dimensionally spatial average pooling function.

This function provides a N-dimensionally generalized version of average\_pooling\_2d(). This acts similarly to ConvolutionND, but it computes the average of input spatial patch for each channel without any parameter instead of computing the inner products.

### **Parameters**

• x (Variable) - Input variable.

- ksize (int or tuple of ints) Size of pooling window. ksize=k and ksize=(k, k, ..., k) are equivalent.
- **stride** (*int or tuple of ints or None*) Stride of pooling applications. stride=s and stride=(s, s, ..., s) are equivalent. If None is specified, then it uses same stride as the pooling window size.
- **pad** (*int* or *tuple* of *ints*) Spatial padding width for the input array. pad=p and pad=(p, p, ..., p) are equivalent.
- **use\_cudnn** (*bool*) If True and cuDNN is enabled, then this function uses cuDNN as the core implementation. cuDNN supports more than one-dimensional pooling.

Returns Output variable.

Return type Variable

**Note:** This function currently does not support cover\_all mode as *max\_pooling\_nd()*. Average pooling runs in non-cover-all mode.

## max\_pooling\_2d

chainer.functions.max\_pooling\_2d(x, ksize, stride=None, pad=0, cover\_all=True, use\_cudnn=True)

Spatial max pooling function.

This function acts similarly to Convolution2D, but it computes the maximum of input spatial patch for each channel without any parameter instead of computing the inner products.

## **Parameters**

- **x** (Variable) Input variable.
- **ksize** (*int or pair of ints*) Size of pooling window. ksize=k and ksize=(k, k) are equivalent.
- **stride** (*int or pair of ints or None*) Stride of pooling applications. stride=s and stride=(s, s) are equivalent. If None is specified, then it uses same stride as the pooling window size.
- **pad** (*int or pair of ints*) Spatial padding width for the input array. pad=p and pad=(p, p) are equivalent.
- **cover\_all** (*bool*) If True, all spatial locations are pooled into some output pixels. It may make the output size larger.
- **use\_cudnn** (*bool*) If True and cuDNN is enabled, then this function uses cuDNN as the core implementation.

Returns Output variable.

**Return type** Variable

### max\_pooling\_nd

chainer.functions.**max\_pooling\_nd**(*x*, *ksize*, *stride=None*, *pad=0*, *cover\_all=True*, *use\_cudnn=True*) N-dimensionally spatial max pooling function. This function provides a N-dimensionally generalized version of  $max_pooling_2d()$ . This acts similarly to ConvolutionND, but it computes the maximum of input spatial patch for each channel without any parameter instead of computing the inner products.

## Parameters

- **x** (Variable) Input variable.
- **ksize** (*int or tuple of ints*) **Size of pooling window**. ksize=k and ksize=(k, k, ..., k) are equivalent.
- **stride** (*int or tuple of ints or None*) Stride of pooling applications. stride=s and stride=(s,s, ..., s) are equivalent. If None is specified, then it uses same stride as the pooling window size.
- **pad** (*int or tuple of ints*) Spatial padding width for the input array. pad=p and pad=(p, p, ..., p) are equivalent.
- **cover\_all** (*bool*) If True, all spatial locations are pooled into some output pixels. It may make the output size larger.
- **use\_cudnn** (*bool*) If True and cuDNN is enabled, then this function uses cuDNN as the core implementation. cuDNN supports more than one-dimensional pooling.

Returns Output variable.

## Return type Variable

# roi\_pooling\_2d

chainer.functions.**roi\_pooling\_2d**(*x*, *rois*, *outh*, *outw*, *spatial\_scale*) Spatial Region of Interest (ROI) pooling function.

This function acts similarly to MaxPooling2D, but it computes the maximum of input spatial patch for each channel with the region of interest.

## Parameters

- **x** (Variable) Input variable. The shape is expected to be 4 dimentional: (n: batch, c: channel, h, height, w: width).
- **rois** (Variable) Input roi variable. The shape is expected to be (n: data size, 5), and each datum is set as below: (batch\_index, x\_min, y\_min, x\_max, y\_max).
- **outh** (*int*) Height of output image after pooled.
- **outw** (*int*) Width of output image after pooled.
- **spatial\_scale** (*float*) **Scale** of the roi is resized.

Returns Output variable.

## Return type Variable

See the original paper proposing ROIPooling: Fast R-CNN.

# spatial\_pyramid\_pooling\_2d

Spatial pyramid pooling function.

It outputs a fixed-length vector regardless of input feature map size.

It performs pooling operation to the input 4D-array  $\times$  with different kernel sizes and padding sizes, and then flattens all dimensions except first dimension of all pooling results, and finally concatenates them along second dimension.

At *i*-th pyramid level, the kernel size  $(k_h^{(i)}, k_w^{(i)})$  and padding size  $(p_h^{(i)}, p_w^{(i)})$  of pooling operation are calculated as below:

$$\begin{aligned} k_h^{(i)} &= \lceil b_h/2^i \rceil, \\ k_w^{(i)} &= \lceil b_w/2^i \rceil, \\ p_h^{(i)} &= (2^i k_h^{(i)} - b_h)/2, \\ p_w^{(i)} &= (2^i k_w^{(i)} - b_w)/2. \end{aligned}$$

where  $\lceil \cdot \rceil$  denotes the ceiling function, and  $b_h, b_w$  are height and width of input variable x, respectively. Note that index of pyramid level *i* is zero-based.

See detail in paper: Spatial Pyramid Pooling in Deep Convolutional Networks for Visual Recognition.

### **Parameters**

- **x** (Variable) Input variable. The shape of x should be (batchsize, # of channels, height, width).
- **pyramid\_height** (*int*) Number of pyramid levels
- **pooling\_class** (*MaxPooling2D* or *AveragePooling2D*) Only MaxPooling2D class can be available for now.
- **use\_cudnn** (*bool*) If True and cuDNN is enabled, then this function uses cuDNN as the core implementation.

### Returns

**Output variable. The shape of the output variable** will be  $(batchsize, c \sum_{h=0}^{H-1} 2^{2h}, 1, 1)$ , where c is the number of channels of input variable x and H is the number of pyramid levels.

Return type Variable

**Note:** This function uses some pooling classes as components to perform spatial pyramid pooling. Now it supports only MaxPooling2D as elemental pooling operator so far.

## unpooling\_2d

chainer.functions.unpooling\_2d(x, ksize, stride=None, pad=0, outsize=None, cover\_all=True) Inverse operation of pooling for 2d array.

This function acts similarly to Deconvolution2D, but it spreads input 2d array's value without any parameter instead of computing the inner products.

- **x** (Variable) Input variable.
- **ksize** (*int or pair of ints*) Size of pooling window. ksize=k and ksize=(k, k) are equivalent.
- **stride** (*int*, *pair* of *ints* or *None*) Stride of pooling applications. stride=s and stride=(s, s) are equivalent. If None is specified, then it uses same stride as the pooling window size.

- **pad** (*int or pair of ints*) Spatial padding width for the input array. pad=p and pad=(p, p) are equivalent.
- **outsize** (*None or pair of ints*) Expected output size (height, width) of array after the operation. If None, the size (height or width) is estimated from the size of input array in first batch with get\_deconv\_outsize(). If outsize is not None, the result of outsize applied to get\_conv\_outsize() must be equal to the shape of the 2d array in the input batch x.
- cover\_all (bool) If True, the output size may be smaller than the size if cover\_all is False. This flag serves to align behavior to the pooling functions which can cover all input locations, see max\_pooling\_2d() and convolution\_2d().

Returns Output variable.

Return type Variable

### upsampling\_2d

```
chainer.functions.upsampling_2d(x, indexes, ksize, stride=None, pad=0, outsize=None, cover_all=True)
```

Upsampling using pooling indices.

This function produces an upsampled image using pooling indices.

### Example

It should be noted that you need to specify use\_cudnn=False when you create MaxPooling2D object because if cuDNN used for operating max pooling, indexes is never created and stored in the MaxPooling2D object.

```
>>> p = F.MaxPooling2D(2, 2, use_cudnn=False)
>>> x = np.arange(1, 37).reshape(1, 1, 6, 6).astype('f')
>>> x = chainer.Variable(x)
>>> x.data
array([[[ 1., 2.,
                     3.,
                          4.,
                               5.,
                                      6.1.
                    9., 10., 11., 12.],
        [ 7.,
              8.,
        [ 13., 14., 15., 16., 17., 18.],
        [ 19., 20., 21., 22., 23., 24.],
        [ 25., 26., 27., 28., 29., 30.],
        [ 31., 32., 33., 34., 35., 36.]]]], dtype=float32)
```

This is the original x before max pooling.

```
>>> pooled_x = p(x)
>>> pooled_x.data
array([[[ 8., 10., 12.],
        [ 20., 22., 24.],
        [ 32., 34., 36.]]]], dtype=float32)
```

This is the output of the max pooling operation. upsampling\_2d needs indexes array stored in the max pooling object p.

```
>>> upsampled_x = F.upsampling_2d(
... pooled_x, p.indexes, p.kh, p.sy, p.ph, x.shape[2:])
>>> upsampled_x.shape
(1, 1, 6, 6)
>>> upsampled_x.data
```

array([[[	0.,	0.,	0.,	0.,	0.,	0.],		
]	Ο.,	8.,	0.,	10.,	0.,	12.],		
[	0.,	0.,	0.,	0.,	0.,	0.],		
]	0.,	20.,	0.,	22.,	0.,	24.],		
]	0.,	0.,	0.,	0.,	0.,	0.],		
] [	0.,	32.,	0.,	34.,	0.,	36.]]]],	dtype=float32)	

# **Parameters**

- **x** (Variable) Input variable.
- indexes (*ndarray* or ndarray) Index array that was used to calculate x with MaxPooling2D.
- **ksize** (*int* or (*int*, *int*)) ksize attribute of MaxPooling2D object that is used to calculate x
- **stride** (*int* or (*int*, *int*)) stride attribute of MaxPooling2D object that is used to calculate x
- **pad** (*int* or (*int*, *int*)) pad attribute of MaxPooling2D object that is used to calculate x
- **outsize**((*int*, *int*)) Expected output size (height, width).
- cover\_all (bool) Whether cover\_all is used in the MaxPooling2D object or not.

Returns Output variable.

Return type Variable

# **Utility functions**

# forget

```
chainer.functions.forget (func, *xs)
```

Call a function without storing internal results.

On a forward propagation Chainer stores all internal results of Function on a computational graph as they are required on backward-propagation. These results consume too much memory when the internal results are too large. This method **forgets** such internal results on forward propagation, and still supports back-propagation with recalculation.

In a forward propagation, this method calls a given function with given variables without creating a computational graph. That means, no internal results are stored. In a backward propagation this method calls the given function again to create a computational graph to execute back-propagation.

This method reduces internal memory usage. Instead it requires more calculation time as it calls the function twice.

## Example

Let f be a function defined as:

```
>>> def f(a, b):
... return a + b + a
```

and, x and y be *Variable*:

```
>>> x = chainer.Variable(np.random.uniform(-1, 1, 5).astype('f'))
>>> y = chainer.Variable(np.random.uniform(-1, 1, 5).astype('f'))
```

When z is calculated as z = f(x, y), its internal result x + y is stored in memory. Instead if you call f with forget ():

>>> z = F.forget(f, x, y)

internal x + y is forgotten.

**Note:** The method does not support functions behaving randomly, such as *dropout()* and *negative\_sampling()*. It is because first results of these function differ from the second one.

### **Parameters**

- **func** (*callable*) A function to call. It needs to be called with *Variable* object(s) and to return a *Variable* object or a tuple of *Variable* objects.
- **xs** (Variable) Argument variables of the function.

**Returns** A variable func returns. If it returns a tuple, the method returns a tuple too.

Return type Variable

# **Standard Link implementations**

Chainer provides many *Link* implementations in the *chainer.links* package.

Note: Some of the links are originally defined in the *chainer.functions* namespace. They are still left in the namespace for backward compatibility, though it is strongly recommended to use them via the *chainer.links* package.

## Learnable connections

### **Bias**

class chainer.links.Bias (axis=1, shape=None)

Broadcasted elementwise summation with learnable parameters.

Computes a elementwise summation as bias() function does except that its second input is a learnable bias parameter b the link has.

- **axis** (*int*) The first axis of the first input of *bias* () function along which its second input is applied.
- **shape** (*tuple of ints*) Shape of the learnable bias parameter. If None, this link does not have learnable parameters so an explicit bias needs to be given to its \_\_call\_\_ method's second input.

See *bias()* for details.

Variables b (Variable) - Bias parameter if shape is given. Otherwise, no attributes.

**\_call\_\_\_**(\*xs) Applies broadcasted elementwise summation.

**Parameters xs** (*list of Variables*) – Input variables whose length should be one if the link has a learnable bias parameter, otherwise should be two.

### **Bilinear**

class chainer.links.Bilinear(left\_size, right\_size, out\_size, nobias=False, initialW=None, initial\_bias=None)

Bilinear layer that performs tensor multiplication.

Bilinear is a primitive link that wraps the *bilinear()* functions. It holds parameters W, V1, V2, and b corresponding to the arguments of *bilinear()*.

#### Parameters

- left\_size (*int*) Dimension of input vector  $e^1(J)$
- right\_size (*int*) Dimension of input vector  $e^2(K)$
- **out\_size** (*int*) Dimension of output vector *y* (*L*)
- nobias (bool) If True, parameters V1, V2, and b are omitted.
- **initialW** (3-D numpy array) Initial value of W. Shape of this argument must be (left\_size, right\_size, out\_size). If None, W is initialized by centered Gaussian distribution properly scaled according to the dimension of inputs and outputs. May also be a callable that takes numpy.ndarray or cupy.ndarray and edits its value.
- initial\_bias (tuple) Initial values of  $V^1$ ,  $V^2$  and b. The length this argument must be 3. Each element of this tuple must have the shapes of (left\_size, output\_size), (right\_size, output\_size), and (output\_size,), respectively. If None,  $V^1$  and  $V^2$  is initialized by scaled centered Gaussian distributions and b is set to 0. May also be a tuple of callables that take numpy.ndarray or cupy.ndarray and edit its value.

### See also:

See chainer.functions.bilinear() for details.

### Variables

- W (Variable) Bilinear weight parameter.
- V1 (Variable) Linear weight parameter for the first argument.
- V2 (Variable) Linear weight parameter for the second argument.
- **b** (Variable) Bias parameter.

### \_\_\_call\_\_\_(*e1*, *e2*)

Applies the bilinear function to inputs and the internal parameters.

- **e1** (Variable) Left input.
- e2 (Variable) Right input.

Returns Output variable.

Return type Variable

## Convolution2D

Two-dimensional convolutional layer.

This link wraps the *convolution\_2d()* function and holds the filter weight and bias vector as parameters.

#### **Parameters**

- **in\_channels** (*int or None*) Number of channels of input arrays. If None, parameter initialization will be deferred until the first forward data pass at which time the size will be determined.
- **out\_channels** (*int*) Number of channels of output arrays.
- **ksize** (*int or pair of ints*) Size of filters (a.k.a. kernels). ksize=k and ksize=(k, k) are equivalent.
- **stride** (*int or pair of ints*) Stride of filter applications. stride=s and stride=(s, s) are equivalent.
- **pad** (*int or pair of ints*) Spatial padding width for input arrays. pad=p and pad=(p, p) are equivalent.
- wscale (float) Scaling factor of the initial weight.
- **bias** (*float*) Initial bias value.
- nobias (bool) If True, then this link does not use the bias term.
- **use\_cudnn** (*bool*) If True, then this link uses cuDNN if available.
- initialW (4-D array) Initial weight value. If None, then this function uses Gaussian distribution scaled by w\_scale to initialize weight. May also be a callable that takes numpy.ndarray or cupy.ndarray and edits its value.
- initial\_bias (1-D array) Initial bias value. If None, then this function uses bias to initialize bias. May also be a callable that takes numpy.ndarray or cupy. ndarray and edits its value.
- **deterministic** (*bool*) The output of this link can be non-deterministic when it uses cuDNN. If this option is True, then it forces cuDNN to use a deterministic algorithm. This option is only available for cuDNN version >= v4.

#### See also:

See chainer.functions.convolution\_2d() for the definition of two-dimensional convolution.

### Variables

- W (Variable) Weight parameter.
- **b** (Variable) Bias parameter.

 $\_call\_(x)$ 

Applies the convolution layer.

**Parameters x** (Variable) – Input image.

Returns Output of the convolution.

Return type Variable

### **ConvolutionND**

N-dimensional convolution layer.

This link wraps the *convolution\_nd()* function and holds the filter weight and bias vector as parameters.

#### **Parameters**

- ndim (*int*) Number of spatial dimensions.
- in\_channels (*int*) Number of channels of input arrays.
- out\_channels (*int*) Number of channels of output arrays.
- **ksize** (*int or tuple of ints*) **Size of filters** (a.k.a. kernels). ksize=k and ksize=(k, k, ..., k) are equivalent.
- **stride** (*int or tuple of ints*) Stride of filter application. stride=s and stride=(s, s, ..., s) are equivalent.
- **pad** (*int* or tuple of *ints*) Spatial padding width for input arrays. pad=p and pad=(p, p, ..., p) are equivalent.
- **initialW** Value used to initialize the filter weight. May be an initializer instance or another value that *init\_weight()* helper function can take.
- **initial\_bias** Value used to initialize the bias vector. May be an initializer instance or another value except None that *init\_weight()* helper function can take. If None is given, this link does not use the bias vector.
- **use\_cudnn** (*bool*) If True, then this link uses cuDNN if available. See *convolution\_nd()* for exact conditions of cuDNN availability.
- **cover\_all** (*bool*)-If True, all spatial locations are convoluted into some output pixels. It may make the output size larger. cover\_all needs to be False if you want to use cuDNN.

### See also:

See  $convolution_nd()$  for the definition of N-dimensional convolution. See  $convolution_2d()$  for the definition of two-dimensional convolution.

### Variables

- W (Variable) Weight parameter.
- **b** (Variable) Bias parameter. If initial\_bias is None, set to None.

 $\__call\__(x)$ 

Applies N-dimensional convolution layer.

Parameters x (Variable) - Input image.
Returns Output of convolution.
Return type Variable

## Deconvolution2D

Two dimensional deconvolution function.

This link wraps the *deconvolution\_2d()* function and holds the filter weight and bias vector as parameters.

### Parameters

- **in\_channels** (*int or None*) Number of channels of input arrays. If None, parameter initialization will be deferred until the first forward data pass at which time the size will be determined.
- **out\_channels** (*int*) Number of channels of output arrays.
- **ksize** (*int or pair of ints*) Size of filters (a.k.a. kernels). ksize=k and ksize=(k, k) are equivalent.
- **stride** (*int or pair of ints*) Stride of filter applications. stride=s and stride=(s, s) are equivalent.
- **pad** (*int or pair of ints*) Spatial padding width for input arrays. pad=p and pad=(p, p) are equivalent.
- wscale (float) Scaling factor of the initial weight.
- **bias** (*float*) Initial bias value.
- nobias (bool) If True, then this function does not use the bias term.
- **outsize** (tuple) Expected output size of deconvolutional operation. It should be pair of height and width  $(out_H, out_W)$ . Default value is None and the outsize is estimated by input size, stride and pad.
- **use\_cudnn** (*bool*) If True, then this function uses cuDNN if available.
- initialW (4-D array) Initial weight value. If None, then this function uses Gaussian distribution scaled by w\_scale to initialize weight. May also be a callable that takes numpy.ndarray or cupy.ndarray and edits its value.
- initial\_bias (1-D array) Initial bias value. If None, then this function uses bias to initialize bias. May also be a callable that takes numpy.ndarray or cupy. ndarray and edits its value.
- **deterministic** (bool) The output of this link can be non-deterministic when it uses cuDNN. If this option is True, then it forces cuDNN to use a deterministic algorithm. This option is only available for cuDNN version >= v4.

The filter weight has four dimensions  $(c_I, c_O, k_H, k_W)$  which indicate the number of input channels, output channels, height and width of the kernels, respectively. The filter weight is initialized with i.i.d. Gaussian random samples, each of which has zero mean and deviation  $\sqrt{1/(c_I k_H k_W)}$  by default. The deviation is scaled by wscale if specified.

The bias vector is of size  $c_0$ . Its elements are initialized by bias argument. If nobias argument is set to True, then this function does not hold the bias parameter.

#### See also:

See chainer.functions.deconvolution\_2d() for the definition of two-dimensional convolution.

### **DeconvolutionND**

N-dimensional deconvolution function.

This link wraps *deconvolution\_nd()* function and holds the filter weight and bias vector as its parameters.

#### **Parameters**

- ndim (*int*) Number of spatial dimensions.
- in\_channels (*int*) Number of channels of input arrays.
- **out\_channels** (*int*) Number of channels of output arrays.
- **ksize** (*int or tuple of ints*) **Size of filters** (a.k.a. kernels). ksize=k and ksize=(k, k, ..., k) are equivalent.
- **stride** (*int or tuple of ints*) Stride of filter application. stride=s and stride=(s, s, ..., s) are equivalent.
- **pad** (*int* or tuple of *ints*) Spatial padding width for input arrays. pad=p and pad=(p, p, ..., p) are equivalent.
- **outsize** (*tuple of ints*) Expected output size of deconvolutional operation. It should be a tuple of ints that represents the output size of each dimension. Default value is None and the outsize is estimated with input size, stride and pad.
- **initialW** Value used to initialize the filter weight. May be an initializer instance of another value the same with that *init\_weight()* function can take.
- **initial\_bias** Value used to initialize the bias vector. May be an initializer instance or another value except None the same with that *init\_weight()* function can take. If None is supplied, this link does not use the bias vector.
- **use\_cudnn** (*bool*) If True, then this link uses cuDNN if available.

### See also:

deconvolution\_nd()

### Variables

- W (Variable) Weight parameter.
- b (Variable) Bias parameter. If initial\_bias is None, set to None.

### DepthwiseConvolution2D

Two-dimensional depthwise convolutional layer.

This link wraps the *depthwise\_convolution\_2d()* function and holds the filter weight and bias vector as parameters.

#### **Parameters**

- **in\_channels** (*int*) Number of channels of input arrays. If None, parameter initialization will be deferred until the first forward data pass at which time the size will be determined.
- **channel\_multiplier** (*int*) **Channel** multiplier number. Number of output arrays equal in\_channels \* channel\_multiplier.
- **ksize** (*int or pair of ints*) Size of filters (a.k.a. kernels). ksize=k and ksize=(k, k) are equivalent.
- **stride** (*int or pair of ints*) Stride of filter applications. stride=s and stride=(s, s) are equivalent.
- **pad** (*int or pair of ints*) Spatial padding width for input arrays. pad=p and pad=(p, p) are equivalent.
- nobias (bool) If True, then this link does not use the bias term.
- initialW (4-D array) Initial weight value. If None, the default initializer is used. May also be a callable that takes numpy.ndarray or cupy.ndarray and edits its value.
- initial\_bias (1-D array) Initial bias value. If None, the bias is set to 0. May also be a callable that takes numpy.ndarray or cupy.ndarray and edits its value.

### See also:

See chainer.functions.depthwise\_convolution\_2d().

### Variables

- W (Variable) Weight parameter.
- **b** (Variable) Bias parameter.

### DilatedConvolution2D

Two-dimensional dilated convolutional layer.

This link wraps the dilated\_convolution\_2d() function and holds the filter weight and bias vector as parameters.

- **in\_channels** (*int or None*) Number of channels of input arrays. If None, parameter initialization will be deferred until the first forward data pass at which time the size will be determined.
- **out\_channels** (*int*) Number of channels of output arrays.
- ksize (int or pair of ints) Size of filters (a.k.a. kernels). ksize=k and ksize=(k, k) are equivalent.
- **stride** (*int or pair of ints*) Stride of filter applications. stride=s and stride=(s, s) are equivalent.

- **pad** (*int* or *pair* of *ints*) Spatial padding width for input arrays. pad=p and pad=(p, p) are equivalent.
- dilate (*int* or pair of *ints*) Dilation factor of filter applications. dilate=d and dilate=(d, d) are equivalent.
- wscale (float) Scaling factor of the initial weight.
- **bias** (float) Initial bias value.
- nobias (bool) If True, then this link does not use the bias term.
- use\_cudnn (bool) If True, then this link uses cuDNN if available.
- initialW (4-D array) Initial weight value. If None, then this function uses scaled Gaussian distribution to initialize weight. May also be a callable that takes numpy. ndarray or cupy.ndarray and edits its value.
- initial\_bias (1-D array) Initial bias value. If None, then this function uses bias to initialize bias. May also be a callable that takes numpy.ndarray or cupy. ndarray and edits its value.

See *chainer.functions.dilated\_convolution\_2d()* for the definition of two-dimensional dilated convolution.

### Variables

- W (Variable) Weight parameter.
- **b** (Variable) Bias parameter.

### \_\_\_call\_\_\_(*x*)

Applies the convolution layer.

**Parameters x** (Variable) – Input image.

Returns Output of the convolution.

Return type Variable

### EmbedID

**class** chainer.links.**EmbedID** (*in\_size*, *out\_size*, *initialW=None*, *ignore\_label=None*)

Efficient linear layer for one-hot input.

This is a link that wraps the  $embed_id()$  function. This link holds the ID (word) embedding matrix W as a parameter.

- in\_size (*int*) Number of different identifiers (a.k.a. vocabulary size).
- **out\_size** (*int*) Size of embedding vector.
- **initialW** (2-D array) Initial weight value. If None, then the matrix is initialized from the standard normal distribution. May also be a callable that takes numpy.ndarray or cupy.ndarray and edits its value.
- **ignore\_label** (*int or None*) If ignore\_label is an int value, i-th column of return value is filled with 0.

```
chainer.functions.embed_id()
```

Variables W (Variable) – Embedding parameter matrix.

 $\_call_(x)$ 

Extracts the word embedding of given IDs.

Parameters x (Variable) – Batch vectors of IDs.

Returns Batch of corresponding embeddings.

Return type Variable

## GRU

class chainer.links.GRU (n\_units, n\_inputs=None, init=None, inner\_init=None, bias\_init=0)
Stateless Gated Recurrent Unit function (GRU).

GRU function has six parameters  $W_r$ ,  $W_z$ , W,  $U_r$ ,  $U_z$ , and U. All these parameters are  $n \times n$  matrices, where n is the dimension of hidden vectors.

Given two inputs a previous hidden vector h and an input vector x, GRU returns the next hidden vector h' defined as

$$r = \sigma(W_r x + U_r h),$$

$$z = \sigma(W_z x + U_z h),$$

$$\bar{h} = tanh(W x + U(r \odot h)),$$

$$h' = (1 - z) \odot h + z \odot \bar{h},$$

where  $\sigma$  is the sigmoid function, and  $\odot$  is the element-wise product.

GRU does not hold the value of hidden vector h. So this is stateless. Use StatefulGRU as a stateful GRU.

#### Parameters

- **n\_units** (*int*) Dimension of hidden vector *h*.
- **n\_inputs** (*int*) Dimension of input vector x. If None, it is set to the same value as n\_units.

See:

- On the Properties of Neural Machine Translation: Encoder-Decoder Approaches [Cho+, SSST2014].
- Empirical Evaluation of Gated Recurrent Neural Networks on Sequence Modeling [Chung+NIPS2014 DLWorkshop].

### See also:

StatefulGRU

#### Highway

Highway module.

In highway network, two gates are added to the ordinal non-linear transformation  $(H(x) = activate(W_h x + b_h))$ . One gate is the transform gate  $T(x) = \sigma(W_t x + b_t)$ , and the other is the carry gate C(x). For simplicity, the author defined C = 1 - T. Highway module returns y defined as

$$y = activate(W_h x + b_h) \odot \sigma(W_t x + b_t) + x \odot (1 - \sigma(W_t x + b_t))$$

The output array has the same spatial size as the input. In order to satisfy this,  $W_h$  and  $W_t$  must be square matrices.

#### **Parameters**

- in\_out\_size (*int*) Dimension of input and output vectors.
- **nobias** (*bool*) If True, then this function does not use the bias.
- **activate** Activation function of plain array. *tanh* is also available.
- **init\_Wh** (2-D array) Initial weight value of plain array. If None, then this function uses Gaussian distribution scaled by w\_scale to initialize  $W_h$ . May also be a callable that takes numpy.ndarray or "cupy.ndarray" and edits its value.
- init\_bh  $(1-D \ array)$  Initial bias value of plain array. If None, then this function uses zero vector to initialize  $b_h$ . May also be a callable that takes numpy.ndarray or cupy.ndarray and edits its value.
- init\_Wt  $(2-D \ array)$  Initial weight value of transform array. If None, then this function uses Gaussian distribution scaled by w\_scale to initialize  $W_t$ . May also be a callable that takes numpy.ndarray or cupy.ndarray and edits its value.
- init\_bt (1-D array) Initial bias value of transform array. Default value is -1 vector. May also be a callable that takes numpy.ndarray or cupy.ndarray and edits its value. Negative value is recommended by the author of the paper. (e.g. -1, -3, ...).

### See: Highway Networks.

#### $\_call_(x)$

Computes the output of the Highway module.

**Parameters x** (Variable) – Input variable.

**Returns** Output variable. Its array has the same spatial size and the same minibatch size as the input array.

**Return type** Variable

### Inception

Inception module of GoogLeNet.

It applies four different functions to the input array and concatenates their outputs along the channel dimension. Three of them are 2D convolutions of sizes 1x1, 3x3 and 5x5. Convolution paths of 3x3 and 5x5 sizes have 1x1 convolutions (called projections) ahead of them. The other path consists of 1x1 convolution (projection) and 3x3 max pooling.

The output array has the same spatial size as the input. In order to satisfy this, Inception module uses appropriate padding for each convolution and pooling.

See: Going Deeper with Convolutions.

### Parameters

- in\_channels (*int*) Number of channels of input arrays.
- **out1** (*int*) Output size of 1x1 convolution path.
- **proj3** (*int*) Projection size of 3x3 convolution path.
- **out3** (*int*) Output size of 3x3 convolution path.
- **proj5** (*int*) Projection size of 5x5 convolution path.
- **out5** (*int*) Output size of 5x5 convolution path.
- **proj\_pool** (*int*) Projection size of max pooling path.
- **conv\_init** A callable that takes numpy.ndarray or cupy.ndarray and edits its value. It is used for initialization of the convolution matrix weights. Maybe be None to use default initialization.
- **bias\_init** A callable that takes numpy.ndarray or cupy.ndarray and edits its value. It is used for initialization of the convolution bias weights. Maybe be None to use default initialization.

#### \_\_\_call\_\_(x)

Computes the output of the Inception module.

Parameters x (Variable) – Input variable.

**Returns** Output variable. Its array has the same spatial size and the same minibatch size as the input array. The channel dimension has size out1 + out3 + out5 + proj\_pool.

Return type Variable

#### **InceptionBN**

inception module of the new GoogLeinet with Batchnormalization.

This chain acts like *Inception*, while InceptionBN uses the *BatchNormalization* on top of each convolution, the 5x5 convolution path is replaced by two consecutive 3x3 convolution applications, and the pooling method is configurable.

See: Batch Normalization: Accelerating Deep Network Training by Reducing Internal Covariate Shift.

- **in\_channels** (*int*) Number of channels of input arrays.
- **out1** (*int*) Output size of the 1x1 convolution path.
- **proj3** (*int*) Projection size of the single 3x3 convolution path.
- **out3** (*int*) Output size of the single 3x3 convolution path.
- proj33 (*int*) Projection size of the double 3x3 convolutions path.
- **out33** (*int*) Output size of the double 3x3 convolutions path.

- pooltype (*str*) Pooling type. It must be either 'max' or 'avg'.
- proj\_pool (bool) If True, do projection in the pooling path.
- **stride** (*int*) Stride parameter of the last convolution of each path.
- **conv\_init** A callable that takes numpy.ndarray or cupy.ndarray and edits its value. It is used for initialization of the convolution matrix weights. Maybe be None to use default initialization.
- **dtype** (*numpy.dtype*) **Type** to use in ~batch\_normalization. BatchNormalization.

### Inception

- **Variables train** (*bool*) If True, then batch normalization layers are used in training mode. If False, they are used in testing mode.
- $\_call\_(x, test=None)$

Computes the output of the InceptionBN module.

### Parameters

- **x** (Variable) An input variable.
- **test** (*bool*) If True, batch normalization layers run in testing mode; if test is omitted, not self.train is used as test.

### Linear

class chainer.links.Linear(in\_size, out\_size, wscale=1, bias=0, nobias=False, initialW=None, initial bias=None)

Linear layer (a.k.a. fully-connected layer).

This is a link that wraps the *linear()* function, and holds a weight matrix W and optionally a bias vector b as parameters.

The weight matrix W is initialized with i.i.d. Gaussian samples, each of which has zero mean and deviation  $\sqrt{1/2}$ 

#### Parameters

•in\_size (*int or None*) – Dimension of input vectors. If None, parameter initialization will be deferred until the first forward data pass at which time the size will be determined.

•out\_size (*int*) – Dimension of output vectors.

•wscale (float) – Scaling factor of the weight matrix.

•bias (float) – Initial bias value.

•nobias (bool) – If True, then this function does not use the bias.

•initialW (2-D array) - Initial weight value. If None, then this function uses Gaussian distribution scaled by w\_scale to initialize weight. May also be a callable that takes numpy.ndarray or cupy. ndarray and edits its value.

•initial\_bias (1-D array) - Initial bias value. If None, then this function uses bias to initialize bias. May also be a callable that takes numpy.ndarray or cupy.ndarray and edits its value.

### See also:

linear()

### Variables

• W (Variable) - Weight parameter.

• **b** (Variable) – Bias parameter.

 $\_call_(x)$ 

Applies the linear layer.

**Parameters x** (Variable) – Batch of input vectors.

Returns Output of the linear layer.

Return type Variable

## LSTM

class chainer.links.LSTM(in\_size, out\_size, \*\*kwargs)
Fully-connected LSTM layer.

This is a fully-connected LSTM layer as a chain. Unlike the *lstm()* function, which is defined as a stateless activation function, this chain holds upward and lateral connections as child links.

It also maintains *states*, including the cell state and the output at the previous time step. Therefore, it can be used as a *stateful LSTM*.

This link supports variable length inputs. The mini-batch size of the current input must be equal to or smaller than that of the previous one. The mini-batch size of c and h is determined as that of the first input x. When mini-batch size of i-th input is smaller than that of the previous input, this link only updates c[0:len(x)] and h[0:len(x)] and doesn't change the rest of c and h. So, please sort input sequences in descending order of lengths before applying the function.

#### **Parameters**

- **in\_size** (*int*) Dimension of input vectors. If None, parameter initialization will be deferred until the first forward data pass at which time the size will be determined.
- **out\_size** (*int*) Dimensionality of output vectors.
- **lateral\_init** A callable that takes numpy.ndarray or cupy.ndarray and edits its value. It is used for initialization of the lateral connections. Maybe be None to use default initialization.
- upward\_init A callable that takes numpy.ndarray or cupy.ndarray and edits its value. It is used for initialization of the upward connections. Maybe be None to use default initialization.
- **bias\_init** A callable that takes numpy.ndarray or cupy.ndarray and edits its value It is used for initialization of the biases of cell input, input gate and output gate.and gates of the upward connection. Maybe a scalar, in that case, the bias is initialized by this value. Maybe be None to use default initialization.
- **forget\_bias\_init** A callable that takes numpy.ndarray or cupy.ndarray and edits its value It is used for initialization of the biases of the forget gate of the upward connection. Maybe a scalar, in that case, the bias is initialized by this value. Maybe be None to use default initialization.

### Variables

- upward (Linear) Linear layer of upward connections.
- **lateral** (Linear) Linear layer of lateral connections.
- c (Variable) Cell states of LSTM units.
- h (Variable) Output at the previous time step.

### $\_call\_(x)$

Updates the internal state and returns the LSTM outputs.

**Parameters x** (Variable) – A new batch from the input sequence.

Returns Outputs of updated LSTM units.

### **Return type** Variable

### reset\_state()

Resets the internal state.

It sets None to the  $\ensuremath{\mathsf{c}}$  and  $\ensuremath{\mathsf{h}}$  attributes.

## $set_state(c, h)$

Sets the internal state.

It sets the c and h attributes.

### **Parameters**

- c (Variable) A new cell states of LSTM units.
- h (Variable) A new output at the previous time step.

### MLPConvolution2D

### **NStepBiLSTM**

class chainer.links.NStepBiLSTM(n\_layers, in\_size, out\_size, dropout, use\_cudnn=True)
 Stacked Bi-directional LSTM for sequnces.

This link is stacked version of Bi-directional LSTM for sequences. It calculates hidden and cell states of all layer at end-of-string, and all hidden states of the last layer for each time.

Unlike chainer.functions.n\_step\_bilstm(), this function automatically sort inputs in descending order by length, and transpose the seuqnece. Users just need to call the link with a list of chainer.Variable holding sequences.

### Parameters

- **n\_layers** (*int*) Number of layers.
- **in\_size** (*int*) Dimensionality of input vectors.
- **out\_size** (*int*) Dimensionality of hidden states and output vectors.
- **dropout** (*float*) Dropout ratio.
- use\_cudnn (bool) Use cuDNN.

### See also:

chainer.functions.n\_step\_bilstm()

### **NStepBiRNNReLU**

class chainer.links.NStepBiRNNReLU(n\_layers, in\_size, out\_size, dropout, use\_cudnn=True)

Stacked Bi-directional RNN for sequnces.

This link is stacked version of Bi-directional RNN for sequences. Note that the activation function is relu. It calculates hidden and cell states of all layer at end-of-string, and all hidden states of the last layer for each time.

Unlike *chainer.functions.n\_step\_birnn()*, this function automatically sort inputs in descending order by length, and transpose the seuqnece. Users just need to call the link with a list of *chainer.Variable* holding sequences.

### Parameters

- n\_layers (int) Number of layers.
- **in\_size** (*int*) Dimensionality of input vectors.
- **out\_size** (*int*) Dimensionality of hidden states and output vectors.
- **dropout** (*float*) Dropout ratio.
- use\_cudnn (bool) Use cuDNN.

### See also:

chainer.functions.n\_step\_birnn()

### **NStepBiRNNTanh**

class chainer.links.NStepBiRNNTanh (n\_layers, in\_size, out\_size, dropout, use\_cudnn=True)
Stacked Bi-directional RNN for sequnces.

This link is stacked version of Bi-directional RNN for sequences. Note that the activation function is tanh. It calculates hidden and cell states of all layer at end-of-string, and all hidden states of the last layer for each time.

Unlike *chainer.functions.n\_step\_birnn()*, this function automatically sort inputs in descending order by length, and transpose the seuqnece. Users just need to call the link with a list of *chainer.Variable* holding sequences.

#### Parameters

- n\_layers (int) Number of layers.
- **in\_size** (*int*) Dimensionality of input vectors.
- **out\_size** (*int*) Dimensionality of hidden states and output vectors.
- **dropout** (*float*) Dropout ratio.
- use\_cudnn (bool) Use cuDNN.

## See also:

chainer.functions.n\_step\_birnn()

### **NStepGRU**

class chainer.links.NStepGRU (n\_layers, in\_size, out\_size, dropout, use\_cudnn=True)
 Stacked Uni-directional GRU for sequnces.

This link is stacked version of Uni-directional GRU for sequences. It calculates hidden and cell states of all layer at end-of-string, and all hidden states of the last layer for each time.

Unlike *chainer.functions.n\_step\_gru()*, this function automatically sort inputs in descending order by length, and transpose the seuqnece. Users just need to call the link with a list of *chainer.Variable* holding sequences.

#### **Parameters**

• n\_layers (*int*) – Number of layers.

- **in\_size** (*int*) Dimensionality of input vectors.
- **out\_size** (*int*) Dimensionality of hidden states and output vectors.
- **dropout** (*float*) Dropout ratio.
- use\_cudnn (bool) Use cuDNN.

chainer.functions.n\_step\_gru()

### **NStepLSTM**

class chainer.links.NStepLSTM(n\_layers, in\_size, out\_size, dropout, use\_cudnn=True)
 Stacked Uni-directional LSTM for sequnces.

This link is stacked version of Uni-directional LSTM for sequences. It calculates hidden and cell states of all layer at end-of-string, and all hidden states of the last layer for each time.

Unlike *chainer.functions.n\_step\_lstm()*, this function automatically sort inputs in descending order by length, and transpose the seuqnece. Users just need to call the link with a list of *chainer.Variable* holding sequences.

### Parameters

- n\_layers (*int*) Number of layers.
- **in\_size** (*int*) Dimensionality of input vectors.
- **out\_size** (*int*) Dimensionality of hidden states and output vectors.
- **dropout** (*float*) Dropout ratio.
- use\_cudnn (bool) Use cuDNN.

### See also:

chainer.functions.n\_step\_lstm()

### **NStepRNNReLU**

class chainer.links.NStepRNNReLU(n\_layers, in\_size, out\_size, dropout, use\_cudnn=True)
 Stacked Uni-directional RNN for sequnces.

This link is stacked version of Uni-directional RNN for sequences. Note that the activation function is relu. It calculates hidden and cell states of all layer at end-of-string, and all hidden states of the last layer for each time.

Unlike *chainer.functions.n\_step\_rnn()*, this function automatically sort inputs in descending order by length, and transpose the seuqnece. Users just need to call the link with a list of *chainer.Variable* holding sequences.

- **n\_layers** (*int*) Number of layers.
- **in\_size** (*int*) Dimensionality of input vectors.
- **out\_size** (*int*) Dimensionality of hidden states and output vectors.
- **dropout** (*float*) Dropout ratio.
- use\_cudnn (bool) Use cuDNN.

chainer.functions.n\_step\_rnn()

### **NStepRNNTanh**

class chainer.links.NStepRNNTanh (n\_layers, in\_size, out\_size, dropout, use\_cudnn=True)
 Stacked Uni-directional RNN for sequnces.

This link is stacked version of Uni-directional RNN for sequences. Note that the activation function is tanh. It calculates hidden and cell states of all layer at end-of-string, and all hidden states of the last layer for each time.

Unlike *chainer.functions.n\_step\_rnn()*, this function automatically sort inputs in descending order by length, and transpose the seuqnece. Users just need to call the link with a list of *chainer.Variable* holding sequences.

#### **Parameters**

- n\_layers (*int*) Number of layers.
- **in\_size** (*int*) Dimensionality of input vectors.
- **out\_size** (*int*) Dimensionality of hidden states and output vectors.
- **dropout** (*float*) Dropout ratio.
- use\_cudnn (bool) Use cuDNN.

### See also:

chainer.functions.n\_step\_rnn()

### Scale

**class** chainer.links.**Scale** (*axis=1*, *W\_shape=None*, *bias\_term=False*, *bias\_shape=None*) Broadcasted elementwise product with learnable parameters.

Computes a elementwise product as scale() function does except that its second input is a learnable weight parameter W the link has.

#### Parameters

- **axis** (*int*) The first axis of the first input of *scale()* function along which its second input is applied.
- W\_shape (tuple of ints) Shape of learnable weight parameter. If None, this link does not have learnable weight parameter so an explicit weight needs to be given to its \_\_call\_\_ method's second input.
- **bias\_term** (bool) Whether to also learn a bias (equivalent to Scale link + Bias link).
- **bias\_shape** (*tuple of ints*) Shape of learnable bias. If W\_shape is None, this should be given to determine the shape. Otherwise, the bias has the same shape W\_shape with the weight parameter and bias\_shape is ignored.

### See also:

See *scale()* for details.

#### Variables

• W (Variable) - Weight parameter if W\_shape is given. Otherwise, no W attribute.

• bias (Bias) - Bias term if bias\_term is True. Otherwise, no bias attribute.

#### \_\_\_call\_\_(\*xs)

Applies broadcasted elementwise product.

**Parameters xs** (*list of Variables*) – Input variables whose length should be one if the link has a learnable weight parameter, otherwise should be two.

### **StatefulGRU**

class chainer.links.StatefulGRU(in\_size, out\_size, init=None, inner\_init=None, bias\_init=0)
Stateful Gated Recurrent Unit function (GRU).

Stateful GRU function has six parameters  $W_r$ ,  $W_z$ , W,  $U_r$ ,  $U_z$ , and U. All these parameters are  $n \times n$  matrices, where n is the dimension of hidden vectors.

Given input vector x, Stateful GRU returns the next hidden vector h' defined as

$$r = \sigma(W_r x + U_r h),$$

$$z = \sigma(W_z x + U_z h),$$

$$\bar{h} = tanh(W x + U(r \odot h)),$$

$$h' = (1 - z) \odot h + z \odot \bar{h},$$

where h is current hidden vector.

As the name indicates, StatefulGRU is *stateful*, meaning that it also holds the next hidden vector h' as a state. Use GRU as a stateless version of GRU.

#### Parameters

- **in\_size** (*int*) Dimension of input vector *x*.
- **out\_size** (*int*) Dimension of hidden vector *h*.
- **init** A callable that takes numpy.ndarray or cupy.ndarray and edits its value. It is used for initialization of the GRU's input units (W). Maybe be *None* to use default initialization.
- **inner\_init** A callable that takes numpy.ndarray or cupy.ndarray and edits its value. It is used for initialization of the GRU's inner recurrent units (U). Maybe be None to use default initialization.
- **bias\_init** A callable or scalar used to initialize the bias values for both the GRU's inner and input units. Maybe be None to use default initialization.

Variables h (Variable) - Hidden vector that indicates the state of *StatefulGRU*.

### See also:

GRU

### StatefulPeepholeLSTM

```
class chainer.links.StatefulPeepholeLSTM(in_size, out_size)
Fully-connected LSTM layer with peephole connections.
```

This is a fully-connected LSTM layer with peephole connections as a chain. Unlike the *LSTM* link, this chain holds peep\_i, peep\_f and peep\_o as child links besides upward and lateral.

Given a input vector x, Peephole returns the next hidden vector h' defined as

$$a = \\ \tanh(upwardx + lateralh), \\ i = \\ \sigma(upwardx + lateralh + peep_ic), \\ f = \\ \sigma(upwardx + lateralh + peep_fc), \\ c' = \\ a \odot i + f \odot c, \\ o = \\ \sigma(upwardx + lateralh + peep_oc'), \\ h' = \\ o \tanh(c'), \end{cases}$$

where  $\sigma$  is the sigmoid function,  $\odot$  is the element-wise product, c is the current cell state, c' is the next cell state and h is the current hidden vector.

#### **Parameters**

- **in\_size** (*int*) Dimension of the input vector *x*.
- **out\_size** (*int*) Dimension of the hidden vector *h*.

### Variables

- upward (Linear) Linear layer of upward connections.
- lateral (Linear) Linear layer of lateral connections.
- **peep\_i** (Linear) Linear layer of peephole connections to the input gate.
- peep\_f (Linear) Linear layer of peephole connections to the forget gate.
- peep\_o (Linear) Linear layer of peephole connections to the output gate.
- c (Variable) Cell states of LSTM units.
- h (Variable) Output at the current time step.

#### $\_call_(x)$

Updates the internal state and returns the LSTM outputs.

**Parameters x** (Variable) – A new batch from the input sequence.

Returns Outputs of updated LSTM units.

### Return type Variable

### reset\_state()

Resets the internal states.

It sets None to the c and h attributes.

### **StatelessLSTM**

Stateless LSTM layer.

This is a fully-connected LSTM layer as a chain. Unlike the *lstm()* function, this chain holds upward and lateral connections as child links. This link doesn't keep cell and hidden states.

### **Parameters**

- **in\_size** (*int or None*) Dimension of input vectors. If None, parameter initialization will be deferred until the first forward data pass at which time the size will be determined.
- **out\_size** (*int*) Dimensionality of output vectors.

#### Variables

- upward (chainer.links.Linear) Linear layer of upward connections.
- lateral (chainer.links.Linear) Linear layer of lateral connections.

#### $\_call\_(c, h, x)$

Returns new cell state and updated output of LSTM.

#### Parameters

- c (Variable) Cell states of LSTM units.
- **h** (Variable) Output at the previous time step.
- **x** (Variable) A new batch from the input sequence.

### Returns

Returns (c\_new, h\_new), where c\_new represents new cell state, and h\_new is updated output of LSTM units.

Return type tuple of ~chainer.Variable

## Activation/loss/normalization functions with parameters

### **BatchNormalization**

This link wraps the batch\_normalization () and fixed\_batch\_normalization () functions.

It runs in three modes: training mode, fine-tuning mode, and testing mode.

In training mode, it normalizes the input by *batch statistics*. It also maintains approximated population statistics by moving averages, which can be used for instant evaluation in testing mode.

In fine-tuning mode, it accumulates the input to compute *population statistics*. In order to correctly compute the population statistics, a user must use this mode to feed mini-batches running through whole training dataset.

In testing mode, it uses pre-computed population statistics to normalize the input variable. The population statistics is approximated if it is computed by training mode, or accurate if it is correctly computed by fine-tuning mode.

### **Parameters**

- **size** (*int* or tuple of *ints*) **Size** (or shape) of channel dimensions.
- decay (float) Decay rate of moving average. It is used on training.
- **eps** (*float*) Epsilon value for numerical stability.
- **dtype** (*numpy*. *dtype*) Type to use in computing.
- **use\_gamma** (*bool*) If True, use scaling parameter. Otherwise, use unit(1) which makes no effect.
- **use\_beta** (*bool*) If True, use shifting parameter. Otherwise, use unit(0) which makes no effect.
- use\_cudnn (bool) If True, then this link uses cuDNN if available.

See: Batch Normalization: Accelerating Deep Network Training by Reducing Internal Covariate Shift

### See also:

batch\_normalization(), fixed\_batch\_normalization()

### Variables

- gamma (Variable) Scaling parameter.
- **beta** (Variable) Shifting parameter.
- avg\_mean (Variable) Population mean.
- avg\_var (Variable) Population variance.
- **N** (*int*) Count of batches given for fine-tuning.
- decay (float) Decay rate of moving average. It is used on training.
- **eps** (*float*) Epsilon value for numerical stability. This value is added to the batch variances.
- use\_cudnn (bool) If True, then this link uses cuDNN if available.

#### **\_\_\_\_call\_\_**(*x*, *test=False*, *finetune=False*)

Invokes the forward propagation of BatchNormalization.

BatchNormalization accepts additional arguments, which controls three different running mode.

#### Parameters

- **x** (Variable) Input variable.
- **test** (*bool*) If True, BatchNormalization runs in testing mode; it normalizes the input using pre-computed statistics.
- **finetune** (*bool*) If finetune is True and test is False, BatchNormalization runs in fine-tuning mode; it accumulates the input array to compute population statistics for normalization, and normalizes the input using batch statistics.

If test is False, then BatchNormalization runs in training mode; it computes moving averages of mean and variance for evaluation during training, and normalizes the input using batch statistics.

#### start finetuning()

Resets the population count for collecting population statistics.

This method can be skipped if it is the first time to use the fine-tuning mode. Otherwise, this method should be called before starting the fine-tuning mode again.

### LayerNormalization

class chainer.links.LayerNormalization (size=None, eps=1e-06, initial\_gamma=None, ini*tial beta=None*) Layer normalization layer on outputs of linear functions.

This link implements a "layer normalization" layer which normalizes the input units by statistics that are computed along the second axis, scales and shifts them. Parameter initialization will be deferred until the first forward data pass at which time the size will be determined.

#### **Parameters**

- size (int) Size of input units. If None, parameter initialization will be deferred until the first forward data pass at which time the size will be determined.
- **eps** (*float*) Epsilon value for numerical stability of normalization.
- initial\_gamma (Initializer) Initializer for scaling vector. If None, then the vector is filled by 1. If a scalar, the vector is filled by it. If numpy.ndarray, the vector is set by it.
- initial beta (Initializer) Initializer for shifting vector. If None, then the vector is filled by 0. If a scalar, the vector is filled by it. If numpy.ndarray, the vector is set by it.

### Variables

- gamma (Variable) Scaling parameter.
- **beta** (Variable) Shifting parameter.
- **eps** (*float*) Epsilon value for numerical stability.

See: Layer Normalization

### $\_call_(x)$

Apply layer normalization to given input.

**Parameters x** (Variable) – Batch vectors. Shape of this value must be (*batch\_size, unit\_size*), e.g., the output of linear().

**Returns** Output of the layer normalization.

**Return type** Variable

### **BinaryHierarchicalSoftmax**

```
class chainer.links.BinaryHierarchicalSoftmax(in size, tree)
```

Hierarchical softmax layer over binary tree.

In natural language applications, vocabulary size is too large to use softmax loss. Instead, the hierarchical softmax uses product of sigmoid functions. It costs only  $O(\log(n))$  time where n is the vocabulary size in average.

At first a user need to prepare a binary tree whose each leaf is corresponding to a word in a vocabulary. When a word x is given, exactly one path from the root of the tree to the leaf of the word exists. Let path(x) =

 $((e_1, b_1), \ldots, (e_m, b_m))$  be the path of x, where  $e_i$  is an index of *i*-th internal node, and  $b_i \in \{-1, 1\}$  indicates direction to move at *i*-th internal node (-1 is left, and 1 is right). Then, the probability of x is given as below:

$$P(x) = \prod_{\substack{(e_i, b_i) \in \text{path}(x) \\ (e_i, b_i) \in \text{path}(x)}} P(b_i | e_i)$$
$$= \prod_{\substack{(e_i, b_i) \in \text{path}(x)}} \sigma(b_i x^\top w_{e_i}),$$

where  $\sigma(\cdot)$  is a sigmoid function, and w is a weight matrix.

This function costs  $O(\log(n))$  time as an average length of paths is  $O(\log(n))$ , and O(n) memory as the number of internal nodes equals n - 1.

#### Parameters

- in\_size (int) Dimension of input vectors.
- **tree** A binary tree made with tuples like ((1, 2), 3).

Variables W (Variable) – Weight parameter matrix.

See: Hierarchical Probabilistic Neural Network Language Model [Morin+, AISTAT2005].

 $\_call\_(x, t)$ 

Computes the loss value for given input and ground truth labels.

#### **Parameters**

- **x** (Variable) Input to the classifier at each node.
- t (Variable) Batch of ground truth labels.

Returns Loss value.

**Return type** Variable

```
static create_huffman_tree (word_counts)
```

Makes a Huffman tree from a dictionary containing word counts.

This method creates a binary Huffman tree, that is required for *BinaryHierarchicalSoftmax*. For example, {0: 8, 1: 5, 2: 6, 3: 4} is converted to ((3, 1), (2, 0)).

**Parameters word\_counts** (dict of int key and int or float values) – Dictionary representing counts of words.

Returns Binary Huffman tree with tuples and keys of word\_coutns.

### BlackOut

```
class chainer.links.BlackOut (in_size, counts, sample_size)
BlackOut loss layer.
```

#### See also:

black\_out () for more detail.

- **in\_size** (*int*) Dimension of input vectors.
- counts (*int list*) Number of each identifiers.
- **sample\_size** (*int*) Number of negative samples.

Variables W (Variable) – Weight parameter matrix.

## CRF1d

```
class chainer.links.CRF1d(n_label)
```

Linear-chain conditional random field loss layer.

This link wraps the crf1d() function. It holds a transition cost matrix as a parameter.

**Parameters n\_label** (*int*) – Number of labels.

#### See also:

crf1d() for more detail.

Variables cost (Variable) – Transition cost parameter.

#### argmax (xs)

Computes a state that maximizes a joint probability.

Parameters xs (list of Variable) – Input vector for each label.

Returns

A tuple of *Variable* representing each log-likelihood and a list representing the argmax path.

Return type tuple

#### See also:

See crfld\_argmax() for more detail.

### SimplifiedDropconnect

class chainer.links.SimplifiedDropconnect(in\_size, out\_size, ratio=0.5, nobias=False, initialW=None, initial\_bias=None)

Fully-connected layer with simplified dropconnect regularization.

Notice: This implementation cannot be used for reproduction of the paper. There is a difference between the current implementation and the original one. The original version uses sampling with gaussian distribution before passing activation function, whereas the current implementation averages before activation.

#### **Parameters**

- **in\_size** (*int*) Dimension of input vectors. If None, parameter initialization will be deferred until the first forward data pass at which time the size will be determined.
- **out\_size** (*int*) Dimension of output vectors.
- nobias (bool) If True, then this link does not use the bias term.
- initialW (3-D array or None) Initial weight value. If None, the default initializer is used. May also be a callable that takes numpy.ndarray or cupy.ndarray and edits its value.
- initial\_bias (2-D array, float or None) Initial bias value. If None, the bias is set to 0. May also be a callable that takes numpy.ndarray or cupy.ndarray and edits its value.

#### Variables

- W (Variable) Weight parameter.
- **b** (Variable) Bias parameter.

simplified\_dropconnect()

#### See also:

Li, W., Matthew Z., Sixin Z., Yann L., Rob F. (2013). Regularization of Neural Network using DropConnect. International Conference on Machine Learning. URL

**\_\_\_\_call\_\_**(x, train=True, mask=None)

Applies the simplified dropconnect layer.

### Parameters

- **x** (chainer.Variable or numpy.ndarray or cupy.ndarray) Batch of input vectors. Its first dimension n is assumed to be the *minibatch dimension*.
- train (bool) If True, executes simplified dropconnect. Otherwise, simplified dropconnect link works as a linear unit.

:param mask (None or chainer.Variable or numpy.ndarray or: cupy.ndarray): If None, randomized simplified dropconnect mask is generated. Otherwise, The mask must be (n, M, N) shaped array. Main purpose of this option is debugging. *mask* array will be used as a dropconnect mask.

Returns Output of the simplified dropconnect layer.

Return type Variable

#### PReLU

class chainer.links.PReLU(shape=(), init=0.25)
Parametric ReLU function as a link.

### Parameters

- **shape** (*tuple of ints*) Shape of the parameter array.
- **init** (*float*) Initial parameter value.

See the paper for details: Delving Deep into Rectifiers: Surpassing Human-Level Performance on ImageNet Classification.

### See also:

chainer.functions.prelu()

Variables ₩ (Variable) – Coefficient of parametric ReLU.

#### \_\_\_call\_\_\_(x)

Applies the parametric ReLU activation function.

Parameters x (Variable) – Input variable.

Returns Output of the parametric ReLU function.

**Return type** *Variable* 

#### Maxout

class chainer.links.Maxout (in\_size, out\_size, pool\_size, wscale=1, initialW=None, initial\_bias=0)
Fully-connected maxout layer.

Let M, P and N be an input dimension, a pool size, and an output dimension, respectively. For an input vector x of size M, it computes

$$Y_i = \max_i (W_{ij} \cdot x + b_{ij}).$$

Here W is a weight tensor of shape (M, P, N), b an optional bias vector of shape (M, P) and  $W_{ij}$ . is a subvector extracted from W by fixing first and second dimensions to i and j, respectively. Minibatch dimension is omitted in the above equation.

As for the actual implementation, this chain has a Linear link with a (M \* P, N) weight matrix and an optional M \* P dimensional bias vector.

#### Parameters

- **in\_size** (*int*) Dimension of input vectors.
- **out\_size** (*int*) Dimension of output vectors.
- **pool\_size** (*int*) Number of channels.
- wscale (float) Scaling factor of the weight matrix.
- initialW (3-D array or None) Initial weight value. If None, then this function uses Gaussian distribution scaled by w\_scale to initialize weight.
- initial\_bias (2-D array, float or None) Initial bias value. If it is float, initial bias is filled with this value. If None, bias is omitted.

Variables *linear* (Link) – The Linear link that performs affine transformation.

#### See also:

#### maxout()

### See also:

Goodfellow, I., Warde-farley, D., Mirza, M., Courville, A., & Bengio, Y. (2013). Maxout Networks. In Proceedings of the 30th International Conference on Machine Learning (ICML-13) (pp. 1319-1327). URL

### $\_call_(x)$

Applies the maxout layer.

**Parameters x** (Variable) – Batch of input vectors.

Returns Output of the maxout layer.

**Return type** *Variable* 

#### **NegativeSampling**

class chainer.links.NegativeSampling(in\_size, counts, sample\_size, power=0.75)

Negative sampling loss layer.

This link wraps the *negative\_sampling()* function. It holds the weight matrix as a parameter. It also builds a sampler internally given a list of word counts.

#### Parameters

• **in\_size** (*int*) – Dimension of input vectors.

- counts (int list) Number of each identifiers.
- **sample\_size** (*int*) Number of negative samples.
- **power** (float) Power factor  $\alpha$ .

negative\_sampling() for more detail.

Variables W (Variable) – Weight parameter matrix.

**\_\_\_\_call\_\_** (*x*, *t*, *reduce='sum'*)

Computes the loss value for given input and ground truth labels.

#### **Parameters**

- **x** (Variable) Input of the weight matrix multiplication.
- t (Variable) Batch of ground truth labels.
- **reduce** (*str*) Reduction option. Its value must be either 'sum' or 'no'. Otherwise, ValueError is raised.

Returns Loss value.

Return type Variable

## Machine learning models

### Classifier

```
class chainer.links.Classifier(predictor, lossfun=<function softmax_cross_entropy>, acc-
fun=<function accuracy>)
```

A simple classifier model.

This is an example of chain that wraps another chain. It computes the loss and accuracy based on a given input/label pair.

### **Parameters**

- **predictor** (Link) Predictor network.
- **lossfun** (function) Loss function.
- **accfun** (function) Function that computes accuracy.

### Variables

- predictor (Link) Predictor network.
- **lossfun** (function) Loss function.
- accfun (function) Function that computes accuracy.
- **y** (Variable) Prediction for the last minibatch.
- **loss** (Variable) Loss value for the last minibatch.
- *accuracy* (Variable) Accuracy for the last minibatch.
- **compute\_accuracy** (*bool*) If True, compute accuracy on the forward computation. The default value is True.

\_call\_(\*args)

Computes the loss value for an input and label pair.

It also computes accuracy and stores it to the attribute.

Parameters args (list of ~chainer.Variable) – Input minibatch.

The all elements of args but last one are features and the last element corresponds to ground truth labels. It feeds features to the predictor and compare the result with ground truth labels.

Returns Loss value.

**Return type** Variable

## **Pre-trained models**

Pre-trained models are mainly used to achieve a good performance with a small dataset, or extract a semantic feature vector. Although CaffeFunction automatically loads a pre-trained model released as a caffemodel, the following link models provide an interface for automatically converting caffemodels, and easily extracting semantic feature vectors.

For example, to extract the feature vectors with VGG16Layers, which is a common pre-trained model in the field of image recognition, users need to write the following few lines:

```
from chainer.links import VGG16Layers
from PIL import Image
model = VGG16Layers()
img = Image.open("path/to/image.jpg")
feature = model.extract([img], layers=["fc7"])["fc7"]
```

where fc7 denotes a layer before the last fully-connected layer. Unlike the usual links, these classes automatically load all the parameters from the pre-trained models during initialization.

### VGG16Layers

During initialization, this chain model automatically downloads the pre-trained caffemodel, convert to another chainer model, stores it on your local directory, and initializes all the parameters with it. This model would be useful when you want to extract a semantic feature vector from a given image, or fine-tune the model on a different dataset. Note that this pre-trained model is released under Creative Commons Attribution License.

If you want to manually convert the pre-trained caffemodel to a chainer model that can be specified in the constructor, please use convert\_caffemodel\_to\_npz classmethod instead.

**Parameters pretrained\_model** (*str*) – the destination of the pre-trained chainer model serialized as a .npz file. If this argument is specified as auto, it automatically downloads the caffemodel from the internet. Note that in this case the converted chainer model is stored on \$CHAINER\_DATASET\_ROOT/pfnet/chainer/models directory, where \$CHAINER\_DATASET\_ROOT is set as \$HOME/.chainer/dataset unless you specify another value as a environment variable. The converted chainer model is automatically used from the second time. If the argument is specified as None, all the parameters are not initialized by the pre-trained model, but the default initializer used in the original paper, i.e., chainer. initializers.Normal(scale=0.01). Variables available\_layers (list of str) - The list of available layer names used by \_\_\_\_\_call\_\_\_ and extract methods.

**\_\_\_\_call\_\_**(*x*, *layers=['prob']*, *test=True*)

Computes all the feature maps specified by layers.

#### **Parameters**

- **x** (Variable) Input variable.
- layers (list of str) The list of layer names you want to extract.
- **test** (*bool*) If True, dropout runs in test mode.
- **Returns** A directory in which the key contains the layer name and the value contains the corresponding feature map variable.

Return type Dictionary of ~chainer.Variable

classmethod convert\_caffemodel\_to\_npz (path\_caffemodel, path\_npz)

Converts a pre-trained caffemodel to a chainer model.

#### **Parameters**

- **path\_caffemodel** (*str*) Path of the pre-trained caffemodel.
- path\_npz (str) Path of the converted chainer model.

**extract** (*images, layers=['fc7'], size=(224, 224), test=True, volatile=OFF*) Extracts all the feature maps of given images.

The difference of directly executing \_\_\_\_\_\_\_ is that it directly accepts images as an input and automatically transforms them to a proper variable. That is, it is also interpreted as a shortcut method that implicitly calls prepare and \_\_\_\_\_\_functions.

### **Parameters**

- images (iterable of PIL. Image or numpy.ndarray) Input images.
- layers (list of str) The list of layer names you want to extract.
- **size** (*pair of ints*) The resolution of resized images used as an input of CNN. All the given images are not resized if this argument is None, but the resolutions of all the images should be the same.
- test (bool) If True, dropout runs in test mode.
- volatile (Flag) Volatility flag used for input variables.
- **Returns** A directory in which the key contains the layer name and the value contains the corresponding feature map variable.

**Return type** Dictionary of ~chainer.Variable

predict (images, oversample=True)

Computes all the probabilities of given images.

#### **Parameters**

- images (iterable of PIL. Image or numpy.ndarray) Input images.
- **oversample** (*bool*) If True, it averages results across center, corners, and mirrors. Otherwise, it uses only the center.

**Returns** Output that contains the class probabilities of given images.

**Return type** Variable

chainer.links.model.vision.vgg.**prepare** (*image*, *size*=(224, 224))

Converts the given image to the numpy array for VGG models.

Note that you have to call this method before <u>call</u> because the pre-trained vgg model requires to resize the given image, covert the RGB to the BGR, subtract the mean, and permute the dimensions before calling.

### Parameters

- **image** (*PIL.Image or numpy.ndarray*) Input image. If an input is numpy. ndarray, its shape must be (height, width), (height, width, channels), or (channels, height, width), and the order of the channels must be RGB.
- **size** (*pair of ints*) Size of converted images. If None, the given image is not resized.

Returns The converted output array.

Return type numpy.ndarray

## GoogLeNet

class chainer.links.GoogLeNet (pretrained\_model='auto')

A pre-trained GoogLeNet model provided by BVLC [1].

When you specify the path of the pre-trained chainer model serialized as a .npz file in the constructor, this chain model automatically initializes all the parameters with it. This model would be useful when you want to extract a semantic feature vector per image, or fine-tune the model on a different dataset.

If you want to manually convert the pre-trained caffemodel to a chainer model that can be specified in the constructor, please use convert\_caffemodel\_to\_npz classmethod instead.

GoogLeNet, which is also called Inception-v1, is an architecture of convolutional neural network proposed in 2014. This model is relatively lightweight and requires small memory footprint during training compared with modern architectures such as ResNet. Therefore, if you fine-tune your network based on a model pre-trained by Imagenet and need to train it with large batch size, GoogLeNet may be useful. On the other hand, if you just want an off-the-shelf classifier, we recommend you to use ResNet50 or other models since they are more accurate than GoogLeNet.

- **Parameters pretrained\_model** (str) the destination of the pre-trained chainer model serialized as a .npz file. If this argument is specified as auto, it automatically downloads the caffemodel from the internet. Note that in this case the converted chainer model is stored on \$CHAINER\_DATASET\_ROOT/pfnet/chainer/models directory, where \$CHAINER\_DATASET\_ROOT is set as \$HOME/.chainer/dataset unless you specify another value as a environment variable. The converted chainer model is automatically used from the second time. If the argument is specified as None, all the parameters are not initialized by the pre-trained model, but the default initializer used in BVLC, i.e., chainer. initializers.LeCunUniform(scale=1.0). Note that, in Caffe, when weight\_filler is specified as "xavier" type without variance\_norm parameter, the weights are initialized by Uniform(-s, s), where  $s = \sqrt{\frac{3}{fan_{in}}}$  and  $fan_{in}$  is the number of input units. This corresponds to LeCunUniform in Chainer but not GlorotUniform.
- Variables available\_layers (list of str) The list of available layer names used by \_\_\_\_\_call\_\_\_ and extract methods.

**\_\_\_\_call\_\_**(x, layers=['prob'], train=False)

Computes all the feature maps specified by layers.

### Parameters

• **x** (Variable) – Input variable. It should be prepared by

- function. (prepare) -
- layers (list of str) The list of layer names you want to extract.
- train (bool) If True, Dropout runs in training mode.

**Returns** A directory in which the key contains the layer name and the value contains the corresponding feature map variable.

**Return type** Dictionary of ~chainer.Variable

classmethod convert\_caffemodel\_to\_npz (path\_caffemodel, path\_npz)

Converts a pre-trained caffemodel to a chainer model.

#### **Parameters**

- **path\_caffemodel** (*str*) Path of the pre-trained caffemodel.
- path\_npz (str) Path of the converted chainer model.
- **extract** (*images, layers=['pool5'], size=(224, 224), train=False, volatile=OFF*) Extracts all the feature maps of given images.

The difference of directly executing \_\_call\_\_ is that it directly accepts images as an input and automatically transforms them to a proper variable. That is, it is also interpreted as a shortcut method that implicitly calls prepare and \_\_call\_\_ functions.

#### Parameters

- images (iterable of PIL.Image or numpy.ndarray) Input images.
- layers (list of str) The list of layer names you want to extract.
- **size** (*pair of ints*) The resolution of resized images used as an input of CNN. All the given images are not resized if this argument is None, but the resolutions of all the images should be the same.
- train (bool) If True, Dropout runs in training mode.
- volatile (Flag) Volatility flag used for input variables.
- **Returns** A directory in which the key contains the layer name and the value contains the corresponding feature map variable.

Return type Dictionary of ~chainer.Variable

#### predict (images, oversample=True)

Computes all the probabilities of given images.

#### **Parameters**

- images (iterable of PIL. Image or numpy.ndarray) Input images.
- **oversample** (*bool*) If True, it averages results across center, corners, and mirrors. Otherwise, it uses only the center.

Returns Output that contains the class probabilities of given images.

### Return type Variable

chainer.links.model.vision.googlenet.**prepare** (*image*, *size*=(224, 224)) Converts the given image to the numpy array for GoogLeNet.

Note that you have to call this method before <u>call</u> because the pre-trained GoogLeNet model requires to resize the given image, covert the RGB to the BGR, subtract the mean, and permute the dimensions before calling.

### Parameters

- **image** (*PIL.Image or numpy.ndarray*) Input image. If an input is numpy. ndarray, its shape must be (height, width), (height, width, channels), or (channels, height, width), and the order of the channels must be RGB.
- **size** (*pair of ints*) Size of converted images. If None, the given image is not resized.

Returns The converted output array.

Return type numpy.ndarray

### **Residual Networks**

When you specify the path of the pre-trained chainer model serialized as a .npz file in the constructor, this chain model automatically initializes all the parameters with it. This model would be useful when you want to extract a semantic feature vector per image, or fine-tune the model on a different dataset. Note that unlike VGG16Layers, it does not automatically download a pre-trained caffemodel. This caffemodel can be downloaded at GitHub.

If you want to manually convert the pre-trained caffemodel to a chainer model that can be specified in the constructor, please use convert\_caffemodel\_to\_npz classmethod instead.

### Parameters

- **pretrained\_model** (*str*) the destination of the pre-trained chainer model serialized as a .npz file. If this argument is specified as auto, it automatically loads and converts the caffemodel from \$CHAINER\_DATASET\_ROOT/pfnet/chainer/models/ ResNet-{n-layers}-model.caffemodel, where \$CHAINER\_DATASET\_ROOT is set as \$HOME/.chainer/dataset unless you specify another value by modifying the environment variable and {n\_layers} is replaced with the specified number of layers given as the first argment to this costructor. Note that in this case the converted chainer model is stored on the same directory and automatically used from the next time. If this argument is specified as None, all the parameters are not initialized by the pre-trained model, but the default initializer used in the original paper, i.e., chainer.initializers. HeNormal(scale=1.0).
- **n\_layers** (*int*) The number of layers of this model. It should be either 50, 101, or 152.

Variables available\_layers (list of str) - The list of available layer names used by \_\_\_\_\_call\_\_\_ and extract methods.

**\_\_\_call\_\_**(x, layers=['prob'], test=True)

Computes all the feature maps specified by layers.

#### Parameters

- **x** (Variable) Input variable.
- layers (list of str) The list of layer names you want to extract.
- **test** (*bool*) If True, BarchNormalization runs in test mode.
- **Returns** A directory in which the key contains the layer name and the value contains the corresponding feature map variable.

Return type Dictionary of ~chainer.Variable

**classmethod convert\_caffemodel\_to\_npz** (*path\_caffemodel*, *path\_npz*, *n\_layers=50*) Converts a pre-trained caffemodel to a chainer model.

#### Parameters

- **path\_caffemodel** (*str*) Path of the pre-trained caffemodel.
- **path\_npz** (*str*) Path of the converted chainer model.

**extract** (*images, layers=['pool5'], size=(224, 224), test=True, volatile=OFF*) Extracts all the feature maps of given images.

The difference of directly executing \_\_call\_\_ is that it directly accepts images as an input and automatically transforms them to a proper variable. That is, it is also interpreted as a shortcut method that implicitly calls prepare and \_\_call\_\_ functions.

#### **Parameters**

- images (iterable of PIL. Image or numpy.ndarray) Input images.
- layers (list of str) The list of layer names you want to extract.
- **size** (*pair of ints*) The resolution of resized images used as an input of CNN. All the given images are not resized if this argument is None, but the resolutions of all the images should be the same.
- **test** (bool) If True, BatchNormalization runs in test mode.
- volatile (Flag) Volatility flag used for input variables.

**Returns** A directory in which the key contains the layer name and the value contains the corresponding feature map variable.

Return type Dictionary of ~chainer.Variable

#### predict (images, oversample=True)

Computes all the probabilities of given images.

### Parameters

- images (iterable of PIL. Image or numpy.ndarray) Input images.
- **oversample** (*bool*) If True, it averages results across center, corners, and mirrors. Otherwise, it uses only the center.

**Returns** Output that contains the class probabilities of given images.

#### **Return type** Variable

### class chainer.links.ResNet50Layers (pretrained\_model='auto')

A pre-trained CNN model with 50 layers provided by MSRA [1].

When you specify the path of the pre-trained chainer model serialized as a .npz file in the constructor, this chain model automatically initializes all the parameters with it. This model would be useful when you want to extract a semantic feature vector per image, or fine-tune the model on a different dataset. Note that unlike VGG16Layers, it does not automatically download a pre-trained caffemodel. This caffemodel can be downloaded at GitHub.

If you want to manually convert the pre-trained caffemodel to a chainer model that can be specified in the constructor, please use convert\_caffemodel\_to\_npz classmethod instead.

ResNet50 has 25,557,096 trainable parameters, and it's 58% and 43% fewer than ResNet101 and ResNet152, respectively. On the other hand, the top-5 classification accuracy on ImageNet dataset drops only 0.7% and 1.1% from ResNet101 and ResNet152, respectively. Therefore, ResNet50 may have the best balance between the accuracy and the model size. It would be basically just enough for many cases, but some advanced models

for object detection or semantic segmentation use deeper ones as their building blocks, so these deeper ResNets are here for making reproduction work easier.

- Parameters pretrained\_model (str) the destination of the pre-trained chainer model serialized as a .npz file. If this argument is specified as auto, it automatically loads and converts the caffemodel from \$CHAINER\_DATASET\_ROOT/pfnet/chainer/ models/ResNet-50-model.caffemodel, where \$CHAINER\_DATASET\_ROOT is set as \$HOME/.chainer/dataset unless you specify another value by modifying the environment variable. Note that in this case the converted chainer model is stored on the same directory and automatically used from the next time. If this argument is specified as None, all the parameters are not initialized by the pre-trained model, but the default initializer used in the original paper, i.e., chainer.initializers.HeNormal(scale=1.0).
- **Variables available\_layers** (*list of str*) The list of available layer names used by \_\_\_\_\_call\_\_\_ and extract methods.

class chainer.links.ResNet101Layers (pretrained\_model='auto')

A pre-trained CNN model with 101 layers provided by MSRA [1].

When you specify the path of the pre-trained chainer model serialized as a .npz file in the constructor, this chain model automatically initializes all the parameters with it. This model would be useful when you want to extract a semantic feature vector per image, or fine-tune the model on a different dataset. Note that unlike VGG16Layers, it does not automatically download a pre-trained caffemodel. This caffemodel can be downloaded at GitHub.

If you want to manually convert the pre-trained caffemodel to a chainer model that can be specified in the constructor, please use convert\_caffemodel\_to\_npz classmethod instead.

ResNet101 has 44,549,224 trainable parameters, and it's 43% fewer than ResNet152 model, while the top-5 classification accuracy on ImageNet dataset drops 1.1% from ResNet152. For many cases, ResNet50 may have the best balance between the accuracy and the model size.

- Parameters pretrained\_model (str) the destination of the pre-trained chainer model serialized as a .npz file. If this argument is specified as auto, it automatically loads and converts the caffemodel from \$CHAINER\_DATASET\_ROOT/pfnet/chainer/ models/ResNet-101-model.caffemodel, where \$CHAINER\_DATASET\_ROOT is set as \$HOME/.chainer/dataset unless you specify another value by modifying the environment variable. Note that in this case the converted chainer model is stored on the same directory and automatically used from the next time. If this argument is specified as None, all the parameters are not initialized by the pre-trained model, but the default initializer used in the original paper, i.e., chainer.initializers.HeNormal(scale=1.0).
- Variables available\_layers (list of str) The list of available layer names used by \_\_\_\_\_\_call\_\_\_ and extract methods.

class chainer.links.ResNet152Layers (pretrained\_model='auto')

A pre-trained CNN model with 152 layers provided by MSRA [1].

When you specify the path of the pre-trained chainer model serialized as a .npz file in the constructor, this chain model automatically initializes all the parameters with it. This model would be useful when you want to extract a semantic feature vector per image, or fine-tune the model on a different dataset. Note that unlike VGG16Layers, it does not automatically download a pre-trained caffemodel. This caffemodel can be downloaded at GitHub.

If you want to manually convert the pre-trained caffemodel to a chainer model that can be specified in the constructor, please use convert\_caffemodel\_to\_npz classmethod instead.

ResNet152 has 60,192,872 trainable parameters, and it's the deepest ResNet model and it achieves the best result on ImageNet classification task in ILSVRC 2015.

- **Parameters pretrained\_model** (*str*) the destination of the pre-trained chainer model serialized as a .npz file. If this argument is specified as auto, it automatically loads and converts the caffemodel from \$CHAINER\_DATASET\_ROOT/pfnet/chainer/models/ResNet-152-model.caffemodel, where \$CHAINER\_DATASET\_ROOT is set as \$HOME/.chainer/dataset unless you specify another value by modifying the environment variable. Note that in this case the converted chainer model is stored on the same directory and automatically used from the next time. If this argument is specified as None, all the parameters are not initialized by the pre-trained model, but the default initializer used in the original paper, i.e., chainer.initializers.HeNormal(scale=1.0).
- **Variables available\_layers** (*list of str*) The list of available layer names used by \_\_\_\_\_call\_\_\_ and extract methods.

chainer.links.model.vision.resnet.**prepare** (*image*, *size*=(224, 224)) Converts the given image to the numpy array for ResNets.

Note that you have to call this method before <u>call</u> because the pre-trained resnet model requires to resize the given image, covert the RGB to the BGR, subtract the mean, and permute the dimensions before calling.

#### Parameters

- **image** (*PIL.Image or numpy.ndarray*) Input image. If an input is numpy. ndarray, its shape must be (height, width), (height, width, channels), or (channels, height, width), and the order of the channels must be RGB.
- **size** (*pair of ints*) Size of converted images. If None, the given image is not resized.

Returns The converted output array.

Return type numpy.ndarray

## **Deprecated links**

### **Parameter**

```
class chainer.links.Parameter (array)
```

Link that just holds a parameter and returns it.

Deprecated since version v1.5: The parameters are stored as variables as of v1.5. Use them directly instead.

**Parameters array** – Initial parameter array.

**Variables W** (Variable) – Parameter variable.

\_call\_(volatile='off')

Returns the parameter variable.

**Parameters volatile** (Flag) – The volatility of the returned variable.

Returns A copy of the parameter variable with given volatility.

Return type Variable

# **Optimizers**

```
class chainer.optimizers.AdaDelta (rho=0.95, eps=1e-06)
Zeiler's ADADELTA.
```

See: http://www.matthewzeiler.com/pubs/googleTR2012/googleTR2012.pdf

**class** chainer.optimizers.**AdaGrad** (*lr=0.001*, *eps=1e-08*) AdaGrad implementation.

See: http://jmlr.org/papers/v12/duchi11a.html

class chainer.optimizers.Adam(alpha=0.001, beta1=0.9, beta2=0.999, eps=1e-08)
Adam optimization algorithm.

See: https://arxiv.org/abs/1412.6980v8

- class chainer.optimizers.MomentumSGD (lr=0.01, momentum=0.9) Classical momentum SGD.
- class chainer.optimizers.NesterovAG (lr=0.01, momentum=0.9)
   Nesterov's Accelerated Gradient.

Formulated as the linear combination coefficients of the velocity and gradient contributions at each iteration.

See: https://arxiv.org/abs/1212.0901

- class chainer.optimizers.RMSprop (lr=0.01, alpha=0.99, eps=1e-08) Hinton's RMSprop.
- class chainer.optimizers.RMSpropGraves (lr=0.0001, alpha=0.95, momentum=0.9, eps=0.0001)
  Alex Graves's RMSprop.

See https://arxiv.org/abs/1308.0850

- class chainer.optimizers.SGD (*lr=0.01*) Vanilla Stochastic Gradient Descent.
- class chainer.optimizers.SMORMS3 (lr=0.001, eps=1e-16) Simon Funk's SMORMS3.

See http://sifter.org/~simon/journal/20150420.html.

# Serializers

## Serialization in NumPy NPZ format

NumPy serializers can be used in arbitrary environments that Chainer runs with. It consists of asymmetric serializer/deserializer due to the fact that numpy.savez() does not support online serialization. Therefore, serialization requires two-step manipulation: first packing the objects into a flat dictionary, and then serializing it into npz format.

Serializer for dictionary.

This is the standard serializer in Chainer. The hierarchy of objects are simply mapped to a flat dictionary with keys representing the paths to objects in the hierarchy.

**Note:** Despite of its name, this serializer DOES NOT serialize the object into external files. It just build a flat dictionary of arrays that can be fed into numpy.savez() and numpy.savez\_compressed(). If you want to use this serializer directly, you have to manually send a resulting dictionary to one of these functions.

- target (dict) The dictionary that this serializer saves the objects to. If target is None, then a new dictionary is created.
- **path** (*str*) The base path in the hierarchy that this serializer indicates.

Variables target (dict) - The target dictionary. Once the serialization completes, this dictionary
 can be fed into numpy.savez() or numpy.savez\_compressed() to serialize it in the
 NPZ format.

class chainer.serializers.NpzDeserializer (npz, path='', strict=True)
 Deserializer for NPZ format.

This is the standard deserializer in Chainer. This deserializer can be used to read an object serialized by  $save_npz()$ .

### Parameters

- **npz** *npz* file object.
- **path** The base path that the deserialization starts from.
- **strict** (*bool*) If True, the deserializer raises an error when an expected value is not found in the given NPZ file. Otherwise, it ignores the value and skip deserialization.

chainer.serializers.save\_npz(filename, obj, compression=True)

Saves an object to the file in NPZ format.

This is a short-cut function to save only one object into an NPZ file.

### Parameters

- **filename** (*str*) Target file name.
- obj Object to be serialized. It must support serialization protocol.
- **compression** (*bool*) If True, compression in the resulting zip file is enabled.

### chainer.serializers.load\_npz(filename, obj)

Loads an object from the file in NPZ format.

This is a short-cut function to load from an .npz file that contains only one object.

### **Parameters**

- filename (*str*) Name of the file to be loaded.
- **obj** Object to be deserialized. It must support serialization protocol.

### Serialization in HDF5 format

```
class chainer.serializers.HDF5Serializer(group, compression=4)
    Serializer for HDF5 format.
```

This is the standard serializer in Chainer. The chain hierarchy is simply mapped to HDF5 hierarchical groups.

Parameters

- group (*h5py.Group*) The group that this serializer represents.
- **compression** (*int*) Gzip compression level.

```
class chainer.serializers.HDF5Deserializer(group, strict=True)
```

Deserializer for HDF5 format.

This is the standard deserializer in Chainer. This deserializer can be used to read an object serialized by *HDF5Serializer*.

### **Parameters**

- group (h5py. Group) The group that the deserialization starts from.
- **strict** (*bool*) If True, the deserializer raises an error when an expected value is not found in the given HDF5 file. Otherwise, it ignores the value and skip deserialization.

```
chainer.serializers.save_hdf5(filename, obj, compression=4)
```

Saves an object to the file in HDF5 format.

This is a short-cut function to save only one object into an HDF5 file. If you want to save multiple objects to one HDF5 file, use *HDF5Serializer* directly by passing appropriate h5py.Group objects.

### Parameters

- **filename** (*str*) Target file name.
- obj Object to be serialized. It must support serialization protocol.
- **compression** (*int*) Gzip compression level.

### chainer.serializers.load\_hdf5(filename, obj)

Loads an object from the file in HDF5 format.

This is a short-cut function to load from an HDF5 file that contains only one object. If you want to load multiple objects from one HDF5 file, use *HDF5Deserializer* directly by passing appropriate h5py. Group objects.

### Parameters

- filename (*str*) Name of the file to be loaded.
- **obj** Object to be deserialized. It must support serialization protocol.

# **Function hooks**

Chainer provides a function-hook mechanism that enriches the behavior of forward and backward propagation of *Function*.

### **Base class**

### $class \verb+ chainer.function.FunctionHook+$

Base class of hooks for Functions.

*FunctionHook* is an callback object that is registered to *Function*. Registered function hooks are invoked before and after forward and backward operations of each function.

Function hooks that derive *FunctionHook* are required to implement four methods: forward\_preprocess(), forward\_postprocess(), backward\_preprocess(), and backward\_postprocess(). By default, these methods do nothing.

Specifically, when \_\_\_\_\_\_() method of some function is invoked, forward\_preprocess() (resp. forward\_postprocess()) of all function hooks registered to this function are called before (resp. after) forward propagation.

Likewise, when *backward()* of some *Variable* is invoked, *backward\_preprocess()* (resp. *backward\_postprocess()*) of all function hooks registered to the function which holds this variable as a gradient are called before (resp. after) backward propagation.

There are two ways to register FunctionHook objects to Function objects.

First one is to use with statement. Function hooks hooked in this way are registered to all functions within with statement and are unregistered at the end of with statement.

### Example

The following code is a simple example in which we measure the elapsed time of a part of forward propagation procedure with *TimerHook*, which is a subclass of *FunctionHook*.

```
>>> from chainer import function_hooks
>>> class Model (chainer.Chain):
        def __call__(self, x1):
. . .
            return F.exp(self.l(x1))
. . .
>>> model1 = Model(l=L.Linear(10, 10))
>>> model2 = Model(l=L.Linear(10, 10))
>>> x = chainer.Variable(np.zeros((1, 10), 'f'))
>>> with chainer.function_hooks.TimerHook() as m:
        = modell(x) 
. . .
        y = model2(x)
. . .
       print("Total time : " + str(m.total_time()))
. . .
        model3 = Model(l=L.Linear(10, 10))
. . .
. . .
        z = model3(y)
Total time : ...
```

In this example, we measure the elapsed times for each forward propagation of all functions in model1 and model2 (specifically, LinearFunction and Exp of model1 and model2). Note that model3 is not a target of measurement as *TimerHook* is unregistered before forward propagation of model3.

**Note:** Chainer stores the dictionary of registered function hooks as a thread local object. So, function hooks registered are different depending on threads.

The other one is to register directly to Function object with  $add_hook$  () method. Function hooks registered in this way can be removed by  $delete_hook$  () method. Contrary to former registration method, function hooks are registered only to the function which  $add_hook$  () is called.

**Parameters name** (*str*) – Name of this function hook.

**backward\_postprocess** (*function*, *in\_data*, *out\_grad*) Callback function invoked after backward propagation.

canoack runction invoked after backward pro-

### Parameters

- function (Function) Function object to which the function hook is registered.
- **in\_data** (tuple of numpy.ndarray or tuple of cupy.ndarray) Input of forward propagation.
- **out\_grad** (tuple of numpy.ndarray or tuple of cupy.ndarray) Gradient data of backward propagation.

backward\_preprocess (function, in\_data, out\_grad)

Callback function invoked before backward propagation.

#### Parameters

• function (Function) – Function object to which the function hook is registered.

- **in\_data** (tuple of numpy.ndarray or tuple of cupy.ndarray) Input data of forward propagation.
- **out\_grad** (tuple of numpy.ndarray or tuple of cupy.ndarray) Gradient data of backward propagation.

### forward\_postprocess (function, in\_data)

Callback function invoked after forward propagation.

#### Parameters

- function (Function) Function object to which the function hook is registered.
- **in\_data** (tuple of numpy.ndarray or tuple of cupy.ndarray) Input data of forward propagation.

### forward\_preprocess (function, in\_data)

Callback function invoked before forward propagation.

### Parameters

- function (Function) Function object to which the function hook is registered.
- **in\_data** (tuple of numpy.ndarray or tuple of cupy.ndarray) Input data of forward propagation.

### **Concrete function hooks**

Function hook that prints debug information.

This function hook outputs the debug information of input arguments of forward and backward methods involved in the hooked functions at preprocessing time (that is, just before each method is called).

Unlike simple "debug print" technique, where users insert print functions at every function to be inspected, we can show the information of all functions involved with single with statement.

Further, this hook enables us to show the information of backward methods without inserting print functions into Chainer's library code.

### Variables

- **sep** Separator of print function.
- end Character to be added at the end of print function.
- **file** Output file\_like object that that redirect to.
- **flush** If True, this hook forcibly flushes the text stream at the end of preprocessing.

### Example

The basic usage is to use it with with statement.

```
>>> from chainer import function_hooks
>>> l = L.Linear(10, 10)
>>> x = chainer.Variable(np.zeros((1, 10), 'f'))
>>> with chainer.function_hooks.PrintHook():
... y = l(x)
... z = F.sum(y)
... z.backward()
```

In this example, PrintHook shows the debug information of forward propagation of LinearFunction (which is implicitly called by 1) and Sum (called by F.sum) and backward propagation of z and y.

class chainer.function\_hooks.TimerHook

Function hook for measuring elapsed time of functions.

**Variables call\_history** – List of measurement results. It consists of pairs of the function that calls this hook and the elapsed time the function consumes.

total\_time()

Returns total elapsed time in seconds.

# Weight Initializers

Weight initializer is an instance of Initializer that destructively edits the contents of numpy.ndarray or *cupy.ndarray*. Typically, weight initializers are passed to \_\_\_\_init\_\_\_ of *Link* and initializes its the weights and biases.

### **Base class**

```
class chainer.initializer.Initializer(dtype=None)
    Initializes array.
```

It initializes the given array.

Variables dtype – Data type specifier. It is for type check in \_\_call\_\_ function.

\_\_call\_\_(array)

Initializes given array.

This method destructively changes the value of array. The derived class is required to implement this method. The algorithms used to make the new values depend on the concrete derived classes.

**Parameters array** (*numpy.ndarray or* cupy.ndarray) – An array to be initialized by this initializer.

### **Concrete initializers**

```
class chainer.initializers.Identity(scale=1.0, dtype=None)
```

Initializes array with the identity matrix.

It initializes the given array with the constant multiple of the identity matrix. Note that arrays to be passed must be 2D squared matrices.

**Variables** *scale* (*scalar*) – A constant to be multiplied to identity matrices.

#### Variables

- **fill\_value** (*scalar or numpy.ndarray or* cupy.ndarray) A constant to be assigned to the initialized array. Broadcast is allowed on this assignment.
- **dtype** Data type specifier.

chainer.initializers.Zero(dtype=None)

Returns initializer that initializes array with the all-zero array.

Parameters dtype – Data type specifier.

Returns An initialized array.

Return type numpy.ndarray or cupy.ndarray

chainer.initializers.One(dtype=None)

Returns initializer that initializes array with the all-one array.

Parameters dtype – Data type specifier.

Returns An initialized array.

**Return type** numpy.ndarray or *cupy.ndarray* 

**class** chainer.initializers.**Normal** (*scale=0.05*, *dtype=None*) Initializes array with a normal distribution.

Each element of the array is initialized by the value drawn independently from Gaussian distribution whose mean is 0, and standard deviation is scale.

### **Parameters**

- **scale** (*float*) Standard deviation of Gaussian distribution.
- **dtype** Data type specifier.

class chainer.initializers.GlorotNormal(scale=1.0, dtype=None)

Initializes array with scaled Gaussian distribution.

Each element of the array is initialized by the value drawn independently from Gaussian distribution whose mean is 0, and standard deviation is  $scale \times \sqrt{\frac{2}{fan_{in}+fan_{out}}}$ , where  $fan_{in}$  and  $fan_{out}$  are the number of input and output units, respectively.

Reference: Glorot & Bengio, AISTATS 2010

### Parameters

- **scale** (*float*) A constant that determines the scale of the standard deviation.
- **dtype** Data type specifier.

class chainer.initializers.HeNormal(scale=1.0, dtype=None)

Initializes array with scaled Gaussian distribution.

Each element of the array is initialized by the value drawn independently from Gaussian distribution whose mean is 0, and standard deviation is  $scale \times \sqrt{\frac{2}{fan_{in}}}$ , where  $fan_{in}$  is the number of input units.

Reference: He et al., https://arxiv.org/abs/1502.01852

**Parameters** 

- scale (float) A constant that determines the scale of the standard deviation.
- **dtype** Data type specifier.

class chainer.initializers.Orthogonal (scale=1.1, dtype=None)

Initializes array with an orthogonal system.

This initializer first makes a matrix of the same shape as the array to be initialized whose elements are drawn independently from standard Gaussian distribution. Next, it applies Singular Value Decomposition (SVD) to the matrix. Then, it initializes the array with either side of resultant orthogonal matrices, depending on the shape of the input array. Finally, the array is multiplied by the constant scale.

If the ndim of the input array is more than 2, we consider the array to be a matrix by concatenating all axes except the first one.

The number of vectors consisting of the orthogonal system (i.e. first element of the shape of the array) must be equal to or smaller than the dimension of each vector (i.e. second element of the shape of the array).

### Variables

- *scale* (*float*) A constant to be multiplied by.
- dtype Data type specifier.

Reference: Saxe et al., https://arxiv.org/abs/1312.6120

```
class chainer.initializers.Uniform(scale=0.05, dtype=None)
```

Initializes array with a scaled uniform distribution.

Each element of the array is initialized by the value drawn independently from uniform distribution [-scale, scale].

#### Variables

- scale (float) A constant that determines the scale of the uniform distribution.
- dtype Data type specifier.

class chainer.initializers.LeCunUniform(scale=1.0, dtype=None)

Initializes array with a scaled uniform distribution.

Each element of the array is initialized by the value drawn independently from uniform distribution [-s,s] where  $s = scale \times \sqrt{\frac{3}{fan_{in}}}$ . Here  $fan_{in}$  is the number of input units.

Reference: LeCun 98, Efficient Backprop http://yann.lecun.com/exdb/publis/pdf/lecun-98b.pdf

#### Variables

- scale (float) A constant that determines the scale of the uniform distribution.
- **dtype** Data type specifier.

class chainer.initializers.GlorotUniform(scale=1.0, dtype=None)

Initializes array with a scaled uniform distribution.

Each element of the array is initialized by the value drawn independently from uniform distribution [-s,s] where  $s = scale \times \sqrt{\frac{6}{fan_{in} + fan_{out}}}$ . Here,  $fan_{in}$  and  $fan_{out}$  are the number of input and output units, respectively.

#### Variables

- scale (float) A constant that determines the scale of the uniform distribution.
- dtype Data type specifier.

**class** chainer.initializers.**HeUniform** (*scale=1.0*, *dtype=None*) Initializes array with scaled uniform distribution.

Each element of the array is initialized by the value drawn independently from uniform distribution [-s,s]where  $s = scale \times \sqrt{\frac{6}{fan_{in}}}$ . Here,  $fan_{in}$  is the number of input units.

Variables

- *scale* (*float*) A constant that determines the scale of the uniform distribution.
- dtype Data type specifier.

### **Helper function**

### chainer.init\_weight (weights, initializer, scale=1.0)

Helper function for initialization of the weight tensor.

This function accepts several types of initializer, prepares the appropriate ~chainer.Initializer if necessary, and does the initialization.

### Parameters

- weights (numpy.ndarray or cupy.ndarray) Weight tensor to be initialized.
- **initializer** The value used to initialize the data. May be None (in which case *HeNormal* is used as an initializer), a scalar to set all values to, an numpy.ndarray to be assigned, or a callable that takes numpy.ndarray or *cupy.ndarray* and edits its value.
- **scale** (*scalar*) A constant to multiply initializer by.

# **Dataset examples**

The most basic *dataset* implementation is an array. Both NumPy and CuPy arrays can be used directly as datasets.

In many cases, though, the simple arrays are not enough to write the training procedure. In order to cover most of such cases, Chainer provides many built-in implementations of datasets.

These built-in datasets are divided into two groups. One is a group of general datasets. Most of them are wrapper of other datasets to introduce some structures (e.g., tuple or dict) to each data point. The other one is a group of concrete, popular datasets. These concrete examples use the downloading utilities in the *chainer.dataset* module to cache downloaded and converted datasets.

### **General datasets**

General datasets are further divided into three types.

The first one is *DictDataset* and *TupleDataset*, both of which combine other datasets and introduce some structures on them.

The second one is *SubDataset*, which represents a subset of an existing dataset. It can be used to separate a dataset for hold-out validation or cross validation. Convenient functions to make random splits are also provided.

The third one is *TransformDataset*, which wraps around a dataset by applying a function to data indexed from the underlying dataset. It can be used to modify behavior of a dataset that is already prepared.

The last one is a group of domain-specific datasets. Currently, ImageDataset and LabeledImageDataset are provided for datasets of images.

### DictDataset

class chainer.datasets.DictDataset(\*\*datasets)

Dataset of a dictionary of datasets.

It combines multiple datasets into one dataset. Each example is represented by a dictionary mapping a key to an example of the corresponding dataset.

**Parameters datasets** – Underlying datasets. The keys are used as the keys of each example. All datasets must have the same length.

### **TupleDataset**

```
class chainer.datasets.TupleDataset(*datasets)
```

Dataset of a tuple of datasets.

It combines multiple datasets into one dataset. Each example is represented by a tuple whose *i*-th item corresponds to the *i*-th dataset.

**Parameters datasets** – Underlying datasets. The i-th one is used for the i-th item of each example. All datasets must have the same length.

### SubDataset

class chainer.datasets.SubDataset(dataset, start, finish, order=None)
Subset of a base dataset.

SubDataset defines a subset of a given base dataset. The subset is defined as an interval of indexes, optionally with a given permutation.

If order is given, then the i-th example of this dataset is the order[start + i]-th example of the base dataset, where i is a non-negative integer. If order is not given, then the i-th example of this dataset is the start + i-th example of the base dataset. Negative indexing is also allowed: in this case, the term start + i is replaced by finish + i.

SubDataset is often used to split a dataset into training and validation subsets. The training set is used for training, while the validation set is used to track the generalization performance, i.e. how the learned model works well on unseen data. We can tune hyperparameters (e.g. number of hidden units, weight initializers, learning rate, etc.) by comparing the validation performance. Note that we often use another set called test set to measure the quality of the tuned hyperparameter, which can be made by nesting multiple SubDatasets.

There are two ways to make training-validation splits. One is a single split, where the dataset is split just into two subsets. It can be done by  $split_dataset()$  or  $split_dataset_random()$ . The other one is a k-fold cross validation, in which the dataset is divided into k subsets, and k different splits are generated using each of the k subsets as a validation set and the rest as a training set. It can be done by  $get_cross_validation_datasets()$ .

### Parameters

- **dataset** Base dataset.
- **start** (*int*) The first index in the interval.
- **finish** (*int*) The next-to-the-last index in the interval.
- **order** (sequence of ints) Permutation of indexes in the base dataset. If this is None, then the ascending order of indexes is used.

chainer.datasets.split\_dataset(dataset, split\_at, order=None)

Splits a dataset into two subsets.

This function creates two instances of *SubDataset*. These instances do not share any examples, and they together cover all examples of the original dataset.

- **dataset** Dataset to split.
- **split\_at** (*int*) Position at which the base dataset is split.
- **order** (*sequence of ints*) Permutation of indexes in the base dataset. See the document of *SubDataset* for details.

### Returns

**Two** *SubDataset* objects. The first subset represents the examples of indexes order[:split\_at] while the second subset represents the examples of indexes order[split\_at:].

### Return type tuple

chainer.datasets.split\_dataset\_random(dataset, first\_size, seed=None)

Splits a dataset into two subsets randomly.

This function creates two instances of *SubDataset*. These instances do not share any examples, and they together cover all examples of the original dataset. The split is automatically done randomly.

### **Parameters**

- **dataset** Dataset to split.
- **first\_size** (*int*) Size of the first subset.
- **seed** (*int*) Seed the generator used for the permutation of indexes. If an integer being convertible to 32 bit unsigned integers is specified, it is guaranteed that each sample in the given dataset always belongs to a specific subset. If None, the permutation is changed randomly.

### Returns

Two *SubDataset* objects. The first subset contains first\_size examples randomly chosen from the dataset without replacement, and the second subset contains the rest of the dataset.

#### Return type tuple

```
chainer.datasets.get_cross_validation_datasets(dataset,n_fold, order=None)
```

Creates a set of training/test splits for cross validation.

This function generates  $n_fold$  splits of the given dataset. The first part of each split corresponds to the training dataset, while the second part to the test dataset. No pairs of test datasets share any examples, and all test datasets together cover the whole base dataset. Each test dataset contains almost same number of examples (the numbers may differ up to 1).

#### **Parameters**

- dataset Dataset to split.
- **n\_fold** (*int*) Number of splits for cross validation.
- **order** (sequence of ints) Order of indexes with which each split is determined. If it is None, then no permutation is used.

Returns List of dataset splits.

Return type list of tuples

chainer.datasets.get\_cross\_validation\_datasets\_random(*dataset*, *n\_fold*, *seed=None*) Creates a set of training/test splits for cross validation randomly.

This function acts almost same as get\_cross\_validation\_dataset(), except automatically generating random permutation.

- dataset Dataset to split.
- **n\_fold** (*int*) Number of splits for cross validation.

• **seed** (*int*) – Seed the generator used for the permutation of indexes. If an integer beging convertible to 32 bit unsigned integers is specified, it is guaranteed that each sample in the given dataset always belongs to a specific subset. If None, the permutation is changed randomly.

Returns List of dataset splits.

**Return type** list of tuples

### **TransformDataset**

### class chainer.datasets.TransformDataset(dataset, transform)

Dataset that indexes the base dataset and transforms the data.

This dataset wraps the base dataset by modifying the behavior of the base dataset's \_\_getitem\_\_(). Arrays returned by \_\_getitem\_\_() of the base dataset with integer as an argument are transformed by the given function transform. Also, \_\_len\_\_() returns the integer returned by the base dataset's \_\_len\_\_().

The function transform takes, as an argument, in\_data, which is the output of the base dataset's \_\_\_\_\_getitem\_\_(), and returns the transformed arrays as output. Please see the following example.

```
>>> from chainer.datasets import get_mnist
>>> from chainer.datasets import TransformDataset
>>> dataset, _ = get_mnist()
>>> def transform(in_data):
... img, label = in_data
... img -= 0.5 # scale to [-0.5, -0.5]
... return img, label
>>> dataset = TransformDataset(dataset, transform)
```

### **Parameters**

- dataset The underlying dataset. The index of this dataset corresponds to the index of the base dataset. This object needs to support functions \_\_getitem\_() and \_\_len\_() as described above.
- **transform** (*callable*) A function that is called to transform values returned by the underlying dataset's \_\_\_\_\_().

### **ImageDataset**

**class** chainer.datasets.**ImageDataset** (*paths*, *root='.'*, *dtype=<type 'numpy.float32'>*) Dataset of images built from a list of paths to image files.

This dataset reads an external image file on every call of the <u>\_\_getitem\_()</u> operator. The paths to the image to retrieve is given as either a list of strings or a text file that contains paths in distinct lines.

Each image is automatically converted to arrays of shape channels, height, width, where channels represents the number of channels in each pixel (e.g., 1 for grey-scale images, and 3 for RGB-color images).

**Note:** This dataset requires the Pillow package being installed. In order to use this dataset, install Pillow (e.g. by using the command pip install Pillow). Be careful to prepare appropriate libraries for image formats you want to use (e.g. libpng for PNG images, and libjpeg for JPG images).

- **paths** (*str* or *list* of *strs*) If it is a string, it is a path to a text file that contains paths to images in distinct lines. If it is a list of paths, the i-th element represents the path to the i-th image. In both cases, each path is a relative one from the root path given by another argument.
- **root** (*str*) Root directory to retrieve images from.
- **dtype** Data type of resulting image arrays.

### LabeledImageDataset

class chainer.datasets.LabeledImageDataset (pairs, root='.', dtype=<type 'numpy.float32'>, label dtype=<type 'numpy.int32'>)

Dataset of image and label pairs built from a list of paths and labels.

This dataset reads an external image file like *ImageDataset*. The difference from *ImageDataset* is that this dataset also returns a label integer. The paths and labels are given as either a list of pairs or a text file contains paths/labels pairs in distinct lines. In the latter case, each path and corresponding label are separated by white spaces. This format is same as one used in Caffe.

**Note:** This dataset requires the Pillow package being installed. In order to use this dataset, install Pillow (e.g. by using the command pip install Pillow). Be careful to prepare appropriate libraries for image formats you want to use (e.g. libpng for PNG images, and libjpeg for JPG images).

### **Parameters**

- **pairs** (*str* or *list* of *tuples*) If it is a string, it is a path to a text file that contains paths to images in distinct lines. If it is a list of pairs, the *i*-th element represents a pair of the path to the *i*-th image and the corresponding label. In both cases, each path is a relative one from the root path given by another argument.
- **root** (*str*) Root directory to retrieve images from.
- **dtype** Data type of resulting image arrays.
- **label\_dtype** Data type of the labels.

### **Concrete datasets**

### **MNIST**

Gets the MNIST dataset.

MNIST is a set of hand-written digits represented by grey-scale 28x28 images. In the original images, each pixel is represented by one-byte unsigned integer. This function scales the pixels to floating point values in the interval [0, scale].

This function returns the training set and the test set of the official MNIST dataset. If withlabel is True, each dataset consists of tuples of images and labels, otherwise it only consists of images.

#### Parameters

• withlabel (bool) – If True, it returns datasets with labels. In this case, each example is a tuple of an image and a label. Otherwise, the datasets only contain images.

- ndim (*int*) Number of dimensions of each image. The shape of each image is determined depending on ndim as follows:
  - ndim == 1: the shape is (784,)
  - ndim == 2: the shape is (28, 28)
  - ndim == 3: the shape is (1, 28, 28)
- **scale** (*float*) Pixel value scale. If it is 1 (default), pixels are scaled to the interval [0, 1].
- **dtype** Data type of resulting image arrays.
- **label\_dtype** Data type of the labels.
- **Returns** A tuple of two datasets. If withlabel is True, both datasets are *TupleDataset* instances. Otherwise, both datasets are arrays of images.

### **CIFAR10/100**

```
chainer.datasets.get_cifar10 (withlabel=True, ndim=3, scale=1.0)
Gets the CIFAR-10 dataset.
```

CIFAR-10 is a set of small natural images. Each example is an RGB color image of size 32x32, classified into 10 groups. In the original images, each component of pixels is represented by one-byte unsigned integer. This function scales the components to floating point values in the interval [0, scale].

This function returns the training set and the test set of the official CIFAR-10 dataset. If withlabel is True, each dataset consists of tuples of images and labels, otherwise it only consists of images.

#### Parameters

- withlabel (bool) If True, it returns datasets with labels. In this case, each example is a tuple of an image and a label. Otherwise, the datasets only contain images.
- ndim (int) Number of dimensions of each image. The shape of each image is determined depending on ndim as follows:
  - ndim == 1: the shape is (3072,)
  - ndim == 3: the shape is (3, 32, 32)
- **scale** (*float*) Pixel value scale. If it is 1 (default), pixels are scaled to the interval [0, 1].
- **Returns** A tuple of two datasets. If withlabel is True, both datasets are *TupleDataset* instances. Otherwise, both datasets are arrays of images.

chainer.datasets.get\_cifar100(withlabel=True, ndim=3, scale=1.0)

Gets the CIFAR-100 dataset.

CIFAR-100 is a set of small natural images. Each example is an RGB color image of size 32x32, classified into 100 groups. In the original images, each component pixels is represented by one-byte unsigned integer. This function scales the components to floating point values in the interval [0, scale].

This function returns the training set and the test set of the official CIFAR-100 dataset. If withlabel is True, each dataset consists of tuples of images and labels, otherwise it only consists of images.

#### Parameters

• withlabel (bool) – If True, it returns datasets with labels. In this case, each example is a tuple of an image and a label. Otherwise, the datasets only contain images.

- ndim (*int*) Number of dimensions of each image. The shape of each image is determined depending on ndim as follows:
- ndim == 1: the shape is (3072,)
- ndim == 3: the shape is (3, 32, 32)
- **scale** (*float*) Pixel value scale. If it is 1 (default), pixels are scaled to the interval [0, 1].
- **Returns** A tuple of two datasets. If withlabel is True, both are *TupleDataset* instances. Otherwise, both datasets are arrays of images.

### **Penn Tree Bank**

```
chainer.datasets.get_ptb_words()
```

Gets the Penn Tree Bank dataset as long word sequences.

Penn Tree Bank is originally a corpus of English sentences with linguistic structure annotations. This function uses a variant distributed at https://github.com/tomsercu/lstm, which omits the annotation and splits the dataset into three parts: training, validation, and test.

This function returns the training, validation, and test sets, each of which is represented as a long array of word IDs. All sentences in the dataset are concatenated by End-of-Sentence mark '<eos>', which is treated as one of the vocabulary.

Returns Int32 vectors of word IDs.

Return type tuple of numpy.ndarray

### See also:

Use get\_ptb\_words\_vocabulary() to get the mapping between the words and word IDs.

chainer.datasets.get\_ptb\_words\_vocabulary()

Gets the Penn Tree Bank word vocabulary.

### Returns

**Dictionary that maps words to corresponding word IDs. The IDs are** used in the Penn Tree Bank long sequence datasets.

### Return type dict

### See also:

See  $get_ptb_words$  () for the actual datasets.

# **Iterator examples**

Chainer provides some iterators that implement typical strategies to create mini-batches by iterating over datasets. *SerialIterator* is the simplest one, which extract mini batches in the main thread. *MultiprocessIterator* is a parallelized version of *SerialIterator*. It maintains worker subprocesses to load the next mini-batch in parallel.

### SerialIterator

**class** chainer.iterators.**SerialIterator** (*dataset*, *batch\_size*, *repeat=True*, *shuffle=True*) Dataset iterator that serially reads the examples.

This is a simple implementation of *Iterator* that just visits each example in either the order of indexes or a shuffled order.

To avoid unintentional performance degradation, the shuffle option is set to True by default. For validation, it is better to set it to False when the underlying dataset supports fast slicing. If the order of examples has an important meaning and the updater depends on the original order, this option should be set to False.

This iterator saves -1 instead of None in snapshots since some serializers do not support None.

#### **Parameters**

- dataset Dataset to iterate.
- **batch\_size** (*int*) Number of examples within each batch.
- **repeat** (*bool*) If True, it infinitely loops over the dataset. Otherwise, it stops iteration at the end of the first epoch.
- **shuffle** (*bool*) If True, the order of examples is shuffled at the beginning of each epoch. Otherwise, examples are extracted in the order of indexes.

### **MultiprocessIterator**

Dataset iterator that loads examples in parallel.

This is an implementation of *Iterator* that loads examples with worker processes. It uses the standard multiprocessing module to parallelize the loading. The dataset is sent to the worker processes in the standard way using pickle.

Note that this iterator effectively prefetches the examples for the next batch asynchronously after the current batch is returned.

This iterator saves -1 instead of None in snapshots since some serializers do not support None.

- **dataset** (*Dataset*) **Dataset** to iterate.
- **batch\_size** (*int*) Number of examples within each batch.
- **repeat** (*bool*) If True, it infinitely loops over the dataset. Otherwise, it stops iteration at the end of the first epoch.
- **shuffle** (*bool*) If True, the order of examples is shuffled at the beginning of each epoch. Otherwise, examples are extracted in the order of indexes.
- **n\_processes** (*int*) Number of worker processes. The number of CPUs is used by default.
- **n\_prefetch** (*int*) Number of prefetch batches.
- **shared\_mem** (*int*) The size of using shared memory per data. If None, size is adjusted automatically.

# **Trainer extensions**

### dump\_graph

chainer.training.extensions.dump\_graph(root\_name, out\_name='cg.dot', variable\_style=None, function\_style=None)

Returns a trainer extension to dump a computational graph.

This extension dumps a computational graph. The graph is output in DOT language.

It only dumps a graph at the first iteration by default.

### Parameters

- **root\_name** (*str*) Name of the root of the computational graph. The root variable is retrieved by this name from the observation dictionary of the trainer.
- **out\_name** (*str*) Output file name.
- **variable\_style** (*dict*) Dot node style for variables. Each variable is rendered by an octagon by default.
- **function\_style** (*dict*) Dot node style for functions. Each function is rendered by a rectangular by default.

### See also:

See build\_computational\_graph() for the variable\_style and function\_style arguments.

### **Evaluator**

Trainer extension to evaluate models on a validation set.

This extension evaluates the current models by a given evaluation function. It creates a *Reporter* object to store values observed in the evaluation function on each iteration. The report for all iterations are aggregated to *DictSummary*. The collected mean values are further reported to the reporter object of the trainer, where the name of each observation is prefixed by the evaluator name. See *Reporter* for details in naming rules of the reports.

Evaluator has a structure to customize similar to that of *StandardUpdater*. The main differences are:

•There are no optimizers in an evaluator. Instead, it holds links to evaluate.

•An evaluation loop function is used instead of an update function.

•Preparation routine can be customized, which is called before each evaluation. It can be used, e.g., to initialize the state of stateful recurrent networks.

There are two ways to modify the evaluation behavior besides setting a custom evaluation function. One is by setting a custom evaluation loop via the eval\_func argument. The other is by inheriting this class and overriding the evaluate() method. In latter case, users have to create and handle a reporter object manually. Users also have to copy the iterators before using them, in order to reuse them at the next time of evaluation.

This extension is called at the end of each epoch by default.

- **iterator** Dataset iterator for the validation dataset. It can also be a dictionary of iterators. If this is just an iterator, the iterator is registered by the name 'main'.
- **target** Link object or a dictionary of links to evaluate. If this is just a link object, the link is registered by the name 'main'.
- **converter** Converter function to build input arrays. *concat\_examples()* is used by default.
- device Device to which the training data is sent. Negative value indicates the host memory (CPU).
- **eval\_hook** Function to prepare for each evaluation process. It is called at the beginning of the evaluation. The evaluator extension object is passed at each call.
- **eval\_func** Evaluation function called at each iteration. The target link to evaluate as a callable is used by default.

#### Variables

- **converter** Converter function.
- device Device to which the training data is sent.
- eval\_hook Function to prepare for each evaluation process.
- **eval\_func** Evaluation function called at each iteration.

#### **\_\_\_call\_\_**(trainer=None)

Executes the evaluator extension.

Unlike usual extensions, this extension can be executed without passing a trainer object. This extension reports the performance on validation dataset using the *report()* function. Thus, users can use this extension independently from any trainer by manually configuring a *Reporter* object.

**Parameters trainer** (Trainer) – Trainer object that invokes this extension. It can be omitted in case of calling this extension manually.

#### Returns

**Result dictionary that contains mean statistics of values** reported by the evaluation function.

#### Return type dict

#### evaluate()

Evaluates the model and returns a result dictionary.

This method runs the evaluation loop over the validation dataset. It accumulates the reported values to *DictSummary* and returns a dictionary whose values are means computed by the summary.

Users can override this method to customize the evaluation routine.

### Returns

**Result dictionary. This dictionary is further reported via** *report()* without specifying any observer.

#### **Return type** dict

get\_all\_iterators()

Returns a dictionary of all iterators.

#### get\_all\_targets()

Returns a dictionary of all target links.

get\_iterator (*name*) Returns the iterator of the given name.

get\_target (name)
 Returns the target link of the given name.

### **ExponentialShift**

Trainer extension to exponentially shift an optimizer attribute.

This extension exponentially increases or decreases the specified attribute of the optimizer. The typical use case is an exponential decay of the learning rate.

This extension is also called before the training loop starts by default.

### Parameters

- **attr** (*str*) Name of the attribute to shift.
- **rate** (*float*) Rate of the exponential shift. This value is multiplied to the attribute at each call.
- **init** (*float*) Initial value of the attribute. If it is None, the extension extracts the attribute at the first call and uses it as the initial value.
- target (float) Target value of the attribute. If the attribute reaches this value, the shift stops.
- **optimizer** (Optimizer) Target optimizer to adjust the attribute. If it is None, the main optimizer of the updater is used.

### LinearShift

class chainer.training.extensions.LinearShift (attr, value\_range, time\_range, optimizer=None)

Trainer extension to change an optimizer attribute linearly.

This extension changes an optimizer attribute from the first value to the last value linearly within a specified duration. The typical use case is warming up of the momentum coefficient.

For example, suppose that this extension is called at every iteration, and value\_range == (x, y) and time\_range == (i, j). Then, this extension keeps the attribute to be x up to the i-th iteration, linearly shifts the value to y by the j-th iteration, and then keeps the value to be y after the j-th iteration.

This extension is also called before the training loop starts by default.

- **attr** (*str*) Name of the optimizer attribute to adjust.
- value\_range (tuple of float) The first and the last values of the attribute.
- time\_range (tuple of ints) The first and last counts of calls in which the attribute is adjusted.
- **optimizer** (Optimizer) Target optimizer object. If it is None, the main optimizer of the trainer is used.

### LogReport

class chainer.training.extensions.LogReport (keys=None, trigger=(1, 'epoch'), postprocess=None, log\_name='log')

Trainer extension to output the accumulated results to a log file.

This extension accumulates the observations of the trainer to *DictSummary* at a regular interval specified by a supplied trigger, and writes them into a log file in JSON format.

There are two triggers to handle this extension. One is the trigger to invoke this extension, which is used to handle the timing of accumulating the results. It is set to 1, 'iteration' by default. The other is the trigger to determine when to emit the result. When this trigger returns True, this extension appends the summary of accumulated values to the list of past summaries, and writes the list to the log file. Then, this extension makes a new fresh summary object which is used until the next time that the trigger fires.

It also adds some entries to each result dictionary.

- •'epoch' and 'iteration' are the epoch and iteration counts at the output, respectively.
- •'elapsed\_time' is the elapsed time in seconds since the training begins. The value is taken from Trainer.elapsed\_time.

### Parameters

- **keys** (*iterable of strs*) Keys of values to accumulate. If this is None, all the values are accumulated and output to the log file.
- trigger Trigger that decides when to aggregate the result and output the values. This is distinct from the trigger of this extension itself. If it is a tuple in the form <int>, 'epoch' or <int>, 'iteration', it is passed to IntervalTrigger.
- **postprocess** Callback to postprocess the result dictionaries. Each result dictionary is passed to this callback on the output. This callback can modify the result dictionaries, which are used to output to the log file.
- **log\_name** (*str*) Name of the log file under the output directory. It can be a format string: the last result dictionary is passed for the formatting. For example, users can use '{iteration}' to separate the log files for different iterations. If the log name is None, it does not output the log to any file.

### log

The current list of observation dictionaries.

### observe\_lr

chainer.training.extensions.**observe\_lr**(*optimizer\_name='main'*, *observation\_key='lr'*) Returns a trainer extension to record the learning rate.

#### Parameters

- **optimizer\_name** (*str*) Name of optimizer whose learning rate is recorded.
- **observation\_key** (*str*) Key of observation to record.

Returns The extension function.

### observe value

chainer.training.extensions.observe\_value (observation\_key, target\_func) Returns a trainer extension to continuously record a value.

#### **Parameters**

- **observation\_key** (*str*) Key of observation to record.
- target\_func (function) Function that returns the value to record. It must take one argument: :class:~chainer.training.Trainer object.

**Returns** The extension function.

### snapshot

chainer.training.extensions.**snapshot**(*savefun=<function*) filesave\_npz>, *name='snapshot\_iter\_{.updater.iteration}', trigger=(1,* 'epoch'))

Returns a trainer extension to take snapshots of the trainer.

This extension serializes the trainer object and saves it to the output directory. It is used to support resuming the training loop from the saved state.

This extension is called once for each epoch by default. The default priority is -100, which is lower than that of most built-in extensions.

Note: This extension first writes the serialized object to a temporary file and then rename it to the target file name. Thus, if the program stops right before the renaming, the temporary file might be left in the output directory.

### **Parameters**

- savefun Function to save the trainer. It takes two arguments: the output file path and the trainer object.
- **filename** (*str*) Name of the file into which the trainer is serialized. It can be a format string, where the trainer object is passed to the str.format() method.
- trigger Trigger that decides when to take snapshot. It can be either an already built trigger object (i.e., a callable object that accepts a trainer object and returns a bool value), or a tuple in the form <int>, 'epoch' or <int>, 'iteration'. In latter case, the tuple is passed to IntervalTrigger.

### snapshot object

chainer.training.extensions.snapshot\_object (target, savefun=<function filename, save\_npz>, trigger=(1, 'epoch'))

Returns a trainer extension to take snapshots of a given object.

This extension serializes the given object and saves it to the output directory.

This extension is called once for each epoch by default. The default priority is -100, which is lower than that of most built-in extensions.

#### **Parameters**

• target – Object to serialize.

- filename (*str*) Name of the file into which the object is serialized. It can be a format string, where the trainer object is passed to the *str.format()* method. For example, 'snapshot\_{.updater.iteration}' is converted to 'snapshot\_10000' at the 10,000th iteration.
- **savefun** Function to save the object. It takes two arguments: the output file path and the object to serialize.
- **trigger** Trigger that decides when to take snapshot. It can be either an already built trigger object (i.e., a callable object that accepts a trainer object and returns a bool value), or a tuple in the form <int>, 'epoch' or <int>, 'iteration'. In latter case, the tuple is passed to IntervalTrigger.

Returns An extension function.

### PlotReport

class chainer.training.extensions.PlotReport	<b>=</b> ( <i>y_keys</i> ,	$x_key = 'it$	eration', tr	ig-
	ger=(1,	'epoch'),	postprocess=No.	ne,
	file_name='plot.png',		marker="	'x',
Trainer extension to output plots	grid=True	)		

Trainer extension to output plots.

This extension accumulates the observations of the trainer to *DictSummary* at a regular interval specified by a supplied trigger, and plot a graph with using them.

There are two triggers to handle this extension. One is the trigger to invoke this extension, which is used to handle the timing of accumulating the results. It is set to 1, 'iteration' by default. The other is the trigger to determine when to emit the result. When this trigger returns True, this extension appends the summary of accumulated values to the list of past summaries, and writes the list to the log file. Then, this extension makes a new fresh summary object which is used until the next time that the trigger fires.

It also adds 'epoch' and 'iteration' entries to each result dictionary, which are the epoch and iteration counts at the output.

**Warning:** If your environment needs to specify a backend of matplotlib explicitly, please call matplotlib.use before importing Chainer. For example:

```
import matplotlib
matplotlib.use('Agg')
```

import chainer

Then, once chainer.training.extensions is imported, matplotlib.use will have no effect.

For the details, please see here: http://matplotlib.org/faq/usage\_faq.html#what-is-a-backend

- **y\_keys** (*iterable of strs*) Keys of values regarded as y. If this is None, nothing is output to the graph.
- **x\_key** (*str*) Keys of values regarded as x. The default value is 'iteration'.
- **trigger** Trigger that decides when to aggregate the result and output the values. This is distinct from the trigger of this extension itself. If it is a tuple in the form <int>, 'epoch' or <int>, 'iteration', it is passed to IntervalTrigger.

- **postprocess** Callback to postprocess the result dictionaries. Figure object, Axes object, and all plot data are passed to this callback in this order. This callback can modify the figure.
- **file\_name** (*str*) Name of the figure file under the output directory. It can be a format string.
- marker (*str*) The marker used to plot the graph. Default is 'x'. If None is given, it draws with no markers.
- grid (bool) Set the axis grid on if True. Default is True.

### **PrintReport**

Trainer extension to print the accumulated results.

This extension uses the log accumulated by a *LogReport* extension to print specified entries of the log in a human-readable format.

### **Parameters**

- entries (list of str) List of keys of observations to print.
- **log\_report** (*str or* LogReport) Log report to accumulate the observations. This is either the name of a LogReport extensions registered to the trainer, or a LogReport instance to use internally.
- **out** Stream to print the bar. Standard output is used by default.

### **ProgressBar**

Trainer extension to print a progress bar and recent training status.

This extension prints a progress bar at every call. It watches the current iteration and epoch to print the bar.

- training\_length (tuple) Length of whole training. It consists of an integer and either 'epoch' or 'iteration'. If this value is omitted and the stop trigger of the trainer is IntervalTrigger, this extension uses its attributes to determine the length of the training.
- update\_interval (int) Number of iterations to skip printing the progress bar.
- **bar\_length** (*int*) Length of the progress bar in characters.
- **out** Stream to print the bar. Standard output is used by default.

# **Trainer triggers**

### Interval

class chainer.training.triggers.IntervalTrigger(period, unit)

Trigger based on a fixed interval.

This trigger accepts iterations divided by a given interval. There are two ways to specify the interval: per iterations and epochs. *Iteration* means the number of updates, while *epoch* means the number of sweeps over the training dataset. Fractional values are allowed if the interval is a number of epochs; the trigger uses the *iteration* and *epoch\_detail* attributes defined by the updater.

For the description of triggers, see get\_trigger().

### Parameters

- **period** (*int or float*) Length of the interval. Must be an integer if unit is 'iteration'.
- unit (*str*) Unit of the length specified by period. It must be either 'iteration' or 'epoch'.

### \_\_\_call\_\_(trainer)

Decides whether the extension should be called on this iteration.

**Parameters trainer** (Trainer) – Trainer object that this trigger is associated with. The updater associated with this trainer is used to determine if the trigger should fire.

### Returns

True if the corresponding extension should be invoked in this iteration.

Return type bool

### ManualSchedule

class chainer.training.triggers.ManualScheduleTrigger (points, unit)

Trigger invoked at specified point(s) of iterations or epochs.

This trigger accepts iterations or epochs indicated by given point(s). There are two ways to specify the point(s): iteration and epoch. iteration means the number of updates, while epoch means the number of sweeps over the training dataset. Fractional values are allowed if the point is a number of epochs; the trigger uses the iteration and epoch\_detail attributes defined by the updater.

### Parameters

- **points** (*int*, *float*, *or list of int or float*) time of the trigger. Must be an integer or list of integer if unit is 'iteration'.
- **unit** (*str*) Unit of the time specified by points. It must be either 'iteration' or 'epoch'.

```
___call__(trainer)
```

Decides whether the extension should be called on this iteration.

**Parameters trainer** (Trainer) – Trainer object that this trigger is associated with. The updater associated with this trainer is used to determine if the trigger should fire.

### Returns

True if the corresponding extension should be invoked in this iteration.

### Return type bool

### Minimum and maximum values

class chainer.training.triggers.MaxValueTrigger(key, trigger=(1, 'epoch'))
Trigger invoked when specific value becomes maximum.

For example you can use this trigger to take snapshot on the epoch the validation accuracy is maximum.

#### Parameters

- **key** (*str*) Key of value. The trigger fires when the value associated with this key becomes maximum.
- **trigger** Trigger that decides the comparison interval between current best value and new value. This must be a tuple in the form of <int>, 'epoch' or <int>, 'iteration' which is passed to *IntervalTrigger*.

class chainer.training.triggers.MinValueTrigger(key, trigger=(1, 'epoch'))
Trigger invoked when specific value becomes minimum.

For example you can use this trigger to take snapshot on the epoch the validation loss is minimum.

#### **Parameters**

- **key** (*str*) Key of value. The trigger fires when the value associated with this key becomes minimum.
- **trigger** Trigger that decides the comparison interval between current best value and new value. This must be a tuple in the form of <int>, 'epoch' or <int>, 'iteration' which is passed to *IntervalTrigger*.

# **Caffe Reference Model Support**

Caffe is a popular framework maintained by BVLC at UC Berkeley. It is widely used by computer vision communities, and aims at fast computation and easy usage without any programming. The BVLC team provides trained reference models in their Model Zoo, one of the reason why this framework gets popular.

Chainer can import the reference models and emulate the network by *Link* implementations. This functionality is provided by the *chainer.links.caffe.CaffeFunction* class.

class chainer.links.caffe.CaffeFunction(model\_path)

Caffe emulator based on the model file of Caffe.

Given a protocol buffers file of a Caffe model, this class loads and emulates it on *Variable* objects. It supports the official reference models provided by BVLC.

Note: protobuf>=3.0.0 is required if you use Python 3 because protobuf 2 is not supported on Python 3.

Note: CaffeFunction ignores the following layers:

•Layers that CaffeFunction does not support (including data layers)

•Layers that have no top blobs

•Layers whose bottom blobs are incomplete (i.e., some or all of them are not given nor computed)

**Warning:** It does not support full compatibility against Caffe. Some layers and configurations are not implemented in Chainer yet, though the reference models provided by the BVLC team are supported except data layers.

### Example

Consider we want to extract the (unnormalized) log class probability of given images using BVLC reference CaffeNet. The model can be downloaded from:

http://dl.caffe.berkeleyvision.org/bvlc\_reference\_caffenet.caffemodel

We want to compute the fc8 blob from the data blob. It is simply written as follows:

```
# Load the model
func = CaffeFunction('path/to/bvlc_reference_caffenet.caffemodel')
# Minibatch of size 10
x_data = numpy.ndarray((10, 3, 227, 227), dtype=numpy.float32)
... # (Fill the minibatch here)
# Forward the pre-trained net
x = Variable(x_data)
y, = func(inputs={'data': x}, outputs=['fc8'])
```

The result y contains the Variable corresponding to the fc8 blob. The computational graph is memorized as a usual forward computation in Chainer, so we can run backprop through this pre-trained net.

**Parameters model\_path** (*str*) – Path to the binary-proto model file of Caffe.

**Variables forwards** (*dict*) – A mapping from layer names to corresponding functions.

\_\_\_\_call\_\_\_(inputs, outputs, disable=(), train=True) Executes a sub-network of the network.

This function acts as an interpreter of the network definition for Caffe. On execution, it interprets each layer one by one, and if the bottom blobs are already computed, then emulates the layer and stores output blobs as *Variable* objects.

### Parameters

- **inputs** (*dict*) A dictionary whose key-value pairs indicate initial correspondences between blob names and *Variable* objects.
- **outputs** (*Iterable*) A list of blob names whose corresponding *Variable* objects are returned.
- **disable** (*Iterable*) A list of layer names that will be ignored during the forward computation.
- **train** (*bool*) If True, this function emulates the TRAIN phase of the Caffe layers. Otherwise, it emulates the TEST phase.

### Returns

A tuple of output Variable objects corresponding to elements of the outputs argument.

Return type tuple

# **Visualization of Computational Graph**

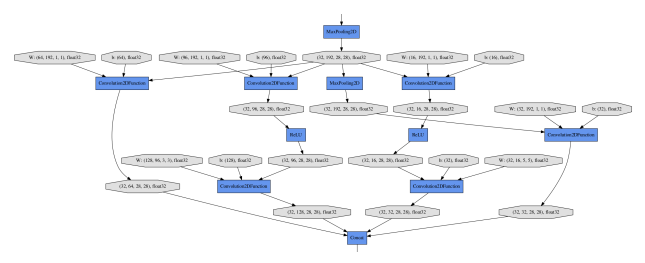
As neural networks get larger and complicated, it gets much harder to confirm if their architectures are constructed properly. Chainer supports visualization of computational graphs. Users can generate computational graphs by invoking *build\_computational\_graph()*. Generated computational graphs are dumped to specified format (Currently Dot Language is supported).

Basic usage is as follows:

```
import chainer.computational_graph as c
...
g = c.build_computational_graph(vs)
with open('path/to/output/file', 'w') as o:
        o.write(g.dump())
```

where vs is list of Variable instances and g is an instance of *ComputationalGraph*. This code generates the computational graph that are backward-reachable (i.e. reachable by repetition of steps backward) from at least one of vs.

Here is an example of (a part of) the generated graph (inception(3a) in GoogLeNet). This example is from example/ imagenet.



Builds a graph of functions and variables backward-reachable from outputs.

- **outputs** (*list*) nodes from which the graph is constructed. Each element of outputs must be either Variable object or Function object.
- **remove\_split** (*bool*) It must be True. This argument is left for backward compatibility.
- **variable\_style** (*dict*) Dot node style for variable. Possible keys are 'shape', 'color', 'fillcolor', 'style', and etc.
- **function\_style** (*dict*) Dot node style for function.
- **rankdir** (*str*) Direction of the graph that must be TB (top to bottom), BT (bottom to top), LR (left to right) or RL (right to left).
- **remove\_variable** (*bool*) If *True*, :class: <sup>4</sup>~chainer.Variable's are removed from the resulting computational graph. Only :class: <sup>4</sup>~chainer.Function's are shown in the output.
- **show\_name** (*bool*) If *True*, the *name* attribute of each node is added to the label of the node. Default is *True*.

#### Returns

A graph consisting of nodes and edges that are backward-reachable from at least one of outputs.

If unchain\_backward was called in some variable in the computational graph before this function, backward step is stopped at this variable.

For example, suppose that computational graph is as follows:

```
|--> f ---> y
x --+
|--> g ---> z
```

Let outputs = [y, z]. Then the full graph is emitted.

Next, let outputs = [y]. Note that z and g are not backward-reachable from y. The resulting graph would be following:

x ---> f ---> y

See TestGraphBuilder for details.

Return type ComputationalGraph

**Note:** The default behavior of ComputationalGraph has been changed from v1.23.0, so that it ouputs the richest representation of a graph as default, namely, styles are set and names of functions and variables are shown. To reproduce the same result as previous versions (<= v1.22.0), please specify *variable\_style=None*, *function\_style=None*, and *show\_name=False* explicitly.

```
class chainer.computational_graph.ComputationalGraph (nodes,
                                                                                   edges,
                                                                                                vari-
                                                                     able_style={'shape':
                                                                                                 'oc-
                                                                     tagon', 'style': 'filled',
                                                                                                 fill-
                                                                                 '#E0E0E0'},
                                                                     color':
                                                                                                func-
                                                                     tion style={'shape':
                                                                                             'record',
                                                                                 'filled',
                                                                     'style':
                                                                                            'fillcolor':
                                                                                       rankdir='TB',
                                                                     '#6495ED'},
                                                                     remove_variable=False,
                                                                     show_name=True)
```

Class that represents computational graph.

Note: We assume that the computational graph is directed and acyclic.

#### **Parameters**

- nodes (list) List of nodes. Each node is either Variable object or Function object.
- edges (list) List of edges. Each edge consists of pair of nodes.
- **variable\_style** (*dict*) Dot node style for variable.
- **function\_style** (*dict*) Dot node style for function.
- **rankdir** (*str*) Direction of the graph that must be TB (top to bottom), BT (bottom to top), LR (left to right) or RL (right to left).
- **remove\_variable** (*bool*) If *True*, :class: <sup>4</sup>~chainer.Variable's are removed from the resulting computational graph. Only :class: <sup>4</sup>~chainer.Function's are shown in the output.
- **show\_name** (*bool*) If *True*, the *name* attribute of each node is added to the label of the node. Default is *True*.

**Note:** The default behavior of ComputationalGraph has been changed from v1.23.0, so that it ouputs the richest representation of a graph as default, namely, styles are set and names of functions and variables are shown. To reproduce the same result as previous versions (<= v1.22.0), please specify *variable\_style=None*, *function\_style=None*, and *show\_name=False* explicitly.

### dump (format='dot')

Dumps graph as a text.

Parameters

- **format** (*str*) The graph language name of the output.
- it must be 'dot'. (Currently,)-

**Returns** The graph in specified format.

Return type str

## **Environment variables**

Here are the environment variables Chainer uses.

CHAINER_CUDNSet 0 to disable cuDNN in Chainer. Otherwise cuDNN is enabled automatically.		
CHAINER_SEED Default seed value of random number generators for CUDA. If it is not set, the seed value is		
	generated from Python random module. Set an integer value in decimal format.	
CHAINER_TYPE_Sett@ tox disable type checking. Otherwise type checking is enabled automatically. See		
	Function for details.	

# CHAPTER 4

# CuPy Reference Manual

This is the official documentation of CuPy, a multi-dimensional array on CUDA with a subset of NumPy interface.

# **CuPy Overview**

CuPy is an implementation of NumPy-compatible multi-dimensional array on CUDA. CuPy consists of the core multidimensional array class, *cupy.ndarray*, and many functions on it. It supports a subset of numpy.ndarray interface that is enough for Chainer.

The following is a brief overview of supported subset of NumPy interface:

- Basic indexing (indexing by ints, slices, newaxes, and Ellipsis)
- Element types (dtypes): bool\_, (u)int{8, 16, 32, 64}, float{16, 32, 64}
- Most of the array creation routines
- · Reshaping and transposition
- · All operators with broadcasting
- All Universal functions (a.k.a. ufuncs) for elementwise operations except those for complex numbers
- Dot product functions (except einsum) using cuBLAS
- Reduction along axes (sum, max, argmax, etc.)

CuPy also includes following features for performance:

- · Customizable memory allocator, and a simple memory pool as an example
- User-defined elementwise kernels
- User-defined reduction kernels
- · cuDNN utilities

CuPy uses on-the-fly kernel synthesis: when a kernel call is required, it compiles a kernel code optimized for the shapes and dtypes of given arguments, sends it to the GPU device, and executes the kernel. The compiled code is cached to \$(HOME)/.cupy/kernel\_cache directory (this cache path can be overwritten by setting the CUPY\_CACHE\_DIR environment variable). It may make things slower at the first kernel call, though this slow down will be resolved at the second execution. CuPy also caches the kernel code sent to GPU device within the process, which reduces the kernel transfer time on further calls.

### A list of supported attributes, properties, and methods of ndarray

### **Memory layout**

base ctypes itemsize flags nbytes shape size strides

Data type

dtype

### Other attributes

Т

### **Array conversion**

tolist() tofile() dump() dumps() astype() copy() view() fill()

### Shape manipulation

reshape() transpose() swapaxes() ravel() squeeze()

### Item selection and manipulation

```
take() diagonal()
```

### Calculation

max() argmax() min() argmin() clip() trace() sum() mean() var() std() prod() dot()

### Arithmetic and comparison operations

\_\_lt\_\_() \_\_le\_\_() \_\_gt\_\_() \_\_ge\_\_() \_\_eq\_\_() \_\_ne\_\_() \_\_nonzero\_\_() \_\_neg\_\_() \_\_pos\_\_() \_\_abs\_\_() \_\_invert\_\_() \_\_add\_\_() \_\_sub\_\_() \_\_mul\_\_() \_\_div\_\_() \_\_truediv\_\_() \_\_floordiv\_\_() \_\_mod\_\_() \_\_divmod\_\_() \_\_pow\_\_() \_\_lshift\_\_() \_\_rshift\_\_() \_\_and\_\_() \_\_or\_\_() \_\_xor\_\_() \_\_iadd\_\_() \_\_isub\_\_() \_\_imul\_\_() \_\_idiv\_\_() \_\_itruediv\_\_() \_\_ifloordiv\_\_() \_\_imod\_\_() \_\_ipow\_\_() \_\_ilshift\_\_() \_\_irshift\_\_() \_\_iand\_\_() \_\_ior\_\_() \_\_ixor\_\_()

### **Special methods**

```
__copy__() __deepcopy__() __reduce__() __array__() __len__() __getitem__()
__setitem__() __int__() __long__() __float__() __oct__() __hex__() __repr__()
__str__()
```

### **Memory transfer**

get() set()

### A list of supported routines of cupy module

### Array creation routines

```
empty() empty_like() eye() identity() ones() ones_like() zeros() zeros_like()
full() full_like()
array() asarray() ascontiguousarray() copy()
arange() linspace()
diag() diagflat()
```

### Array manipulation routines

```
copyto()
reshape() ravel()
rollaxis() swapaxes() transpose()
atleast_1d() atleast_2d() atleast_3d() broadcast broadcast_arrays()
broadcast_to() expand_dims() squeeze()
column_stack() concatenate() dstack() hstack() vstack()
array_split() dsplit() hsplit() split() vsplit()
roll()
```

### **Binary operations**

bitwise\_and bitwise\_or bitwise\_xor invert left\_shift right\_shift

### **Indexing routines**

take() diagonal()

### Input and output

```
load() save() savez() savez_compressed()
array_repr() array_str()
```

### Linear algebra

```
dot() vdot() inner() outer() tensordot()
trace()
```

### **Logic functions**

isfinite isinf isnan
logical\_and logical\_or logical\_not logical\_xor
greater greater\_equal less less\_equal equal not\_equal

### **Mathematical functions**

sin cos tan arcsin arccos arctan hypot arctan2 deg2rad rad2deg degrees radians
sinh cosh tanh arcsinh arccosh arctanh
rint floor ceil trunc
sum() prod()
exp expm1 exp2 log log10 log2 log1p logaddexp logaddexp2
signbit copysign ldexp frexp nextafter
add reciprocal negative multiply divide power subtract true\_divide floor\_divide fmod
mod modf remainder
clip() sqrt square absolute sign maximum minimum fmax fmin

### Sorting, searching, and counting

argmax() argmin() count\_nonzero() nonzero() flatnonzero() where()

### **Statistics**

```
amin() amax()
mean() var() std()
bincount()
```

### Padding

pad()

### **External Functions**

scatter\_add()

### Other

asnumpy()

# Multi-Dimensional Array (ndarray)

#### class cupy.ndarray

Multi-dimensional array on a CUDA device.

This class implements a subset of methods of numpy.ndarray. The difference is that this class allocates the array content on the current GPU device.

### Parameters

- **shape** (tuple of ints) Length of axes.
- dtype Data type. It must be an argument of numpy.dtype.
- memptr (cupy.cuda.MemoryPointer) Pointer to the array content head.
- **strides** (tuple of ints) The strides for axes.
- order ({ 'C', 'F'}) Row-major (C-style) or column-major (Fortran-style) order.

### Variables

- **base** (*None or* cupy.ndarray) Base array from which this array is created as a view.
- data (cupy.cuda.MemoryPointer) Pointer to the array content head.
- **dtype** (*numpy.dtype*) **Dtype** object of element type.

See also:

Data type objects (dtype)

• **size** (*int*) – Number of elements this array holds.

This is equivalent to product over the shape tuple.

See also:

numpy.ndarray.size

т

Shape-reversed view of the array.

If ndim < 2, then this is just a reference to the array itself.

\_\_abs\_\_\_
\_\_add\_\_\_
x.\_\_add\_\_(y) <==> x+y
\_\_\_and\_\_\_
x.\_\_and\_\_(y) <==> x&y
\_\_\_delitem\_\_\_
x.\_\_delitem\_\_(y) <==> del x[y]

\_\_\_div\_\_\_ x.\_\_div\_\_(y) <==> x/y

\_\_divmod\_\_\_

```
____eq___
x.__eq__(y) <==> x==y
___float___
___floordiv___
x.__floordiv__(y) <==> x//y
___ge___
x.__ge__(y) <==> x>=y
___getitem__
```

x.\_\_getitem\_\_(y) <==> x[y]

Supports both basic and advanced indexing.

Note: Currently, it does not support slices that consists of more than one boolean arrays

**Note:** CuPy handles out-of-bounds indices differently from NumPy. NumPy handles them by raising an error, but CuPy wraps around them.

>>> a = cupy.arange(3)
>>> a[[1, 3]]
array([1, 0])

### \_\_\_gt\_\_\_

 $x.\_gt\_(y) <==> x>y$ 

### \_\_hex\_\_

\_\_\_iadd\_\_\_(y) <==> x+=y

\_\_\_iand\_\_\_

 $x.\_iand\_(y) <==> x \&=y$ 

\_\_idiv\_\_

 $x.\_idiv\_(y) \iff x/=y$ 

\_\_ifloordiv\_\_\_

x.\_\_ifloordiv\_\_(y) <==> x//=y

\_\_ilshift\_

x.\_\_ilshift\_\_(y) <==> x<<=y

\_\_\_imod\_\_\_

x.\_\_imod\_\_(y) <==> x%=y

\_\_imul\_\_\_ x.\_\_imul\_\_(y) <==> x\*=y

\_\_int\_\_

\_\_invert\_\_ x.\_\_invert\_\_() <==> ~x

\_\_ior\_\_\_

 $x.\_ior\_(y) <==> x|=y$ 

\_\_ipow\_\_ x.\_\_ipow\_\_(y) <==> x\*\*=y \_\_irshift x.\_\_irshift\_\_(y) <==> x>>=y \_\_isub\_\_\_  $x.\_isub\_(y) \iff x-=y$ \_\_\_itruediv\_\_\_ x.\_\_itruediv\_\_(y)  $\leq x/=y$ \_\_ixor\_\_  $x.\_ixor\_(y) \iff x^=y$ \_\_le\_\_ x.\_\_le\_\_(y) <==> x<=y \_\_len\_\_ long \_\_lshift\_\_  $x.\_lshift\_(y) <==> x << y$ \_\_1t\_\_  $x.\_lt\_(y) \leq x < y$ \_\_mod\_\_\_  $x.\_mod\_(y) \leq x\%y$ \_\_mul\_\_  $x.\_mul\_(y) \iff x*y$ \_\_ne\_\_  $x.\_ne\_(y) \le x!=y$ \_\_neg\_\_ x.\_\_neg\_\_() <==> -x \_\_nonzero\_\_ x.\_\_nonzero\_\_() <=> x != 0\_\_oct\_\_ \_\_or\_\_  $x.\_or\_(y) \iff x|y$ \_\_pos\_\_ x.\_\_pos\_\_() <==> +x \_\_pow\_\_\_ \_\_\_radd\_\_\_  $x.\_radd\_(y) \le y+x$ \_\_\_rand\_\_\_  $x.\_rand\_(y) \iff y\&x$ \_\_\_rdiv\_\_\_  $x.\_rdiv\_(y) \iff y/x$ rdivmod \_\_\_repr\_\_\_

\_\_\_rfloordiv x.\_\_rfloordiv\_\_(y)  $\leq = y//x$ \_\_\_rlshift\_ x.\_\_rlshift\_\_(y) <==> y<<x rmod  $x.\_rmod\_(y) \iff y\%x$ rmul  $x.\_rmul\_(y) \iff y^*x$ \_ror\_\_  $x.\_ror\_(y) \iff y|x$ \_\_\_rpow\_\_\_ \_\_\_\_rrshift\_\_\_ x.\_\_rrshift\_\_(y) <==> y>>x \_\_\_rshift\_\_ x.\_\_rshift\_\_(y) <==> x>>y rsub  $x\_rsub\_(y) \iff y-x$ \_\_\_rtruediv\_\_\_ x.\_\_rtruediv\_\_(y)  $\leq = y/x$ \_\_rxor\_\_ x.\_\_rxor\_\_(y) <==> y^x \_\_\_setitem\_\_\_ x.\_\_setitem\_\_(slices, y) <==> x[slices] = y Supports both basic and advanced indexing.

Note: Currently, it does not support slices that consists of more than one boolean arrays

**Note:** CuPy handles out-of-bounds indices differently from NumPy when using integer array indexing. NumPy handles them by raising an error, but CuPy wraps around them.

```
>>> import cupy
>>> x = cupy.arange(3)
>>> x[[1, 3]] = 10
>>> x
array([10, 10, 2])
```

**Note:** The behavior differs from NumPy when integer arrays in slices reference the same location multiple times. In that case, the value that is actually stored is undefined.

```
>>> import cupy; import numpy
>>> a = cupy.zeros((2,))
>>> i = cupy.arange(10000) % 2
>>> v = cupy.arange(10000).astype(numpy.float)
>>> a[i] = v
```

>>> a
array([ 9150., 9151.])

On the other hand, NumPy stores the value corresponding to the last index among the indices referencing duplicate locations.

```
>>> import numpy
>>> a_cpu = numpy.zeros((2,))
>>> i_cpu = numpy.arange(10000) % 2
>>> v_cpu = numpy.arange(10000).astype(numpy.float)
>>> a_cpu[i_cpu] = v_cpu
>>> a_cpu
array([ 9998., 9999.])
```

\_\_str\_\_

```
__sub___
```

x.\_\_sub\_\_(y) <==> x-y

# <u>\_\_\_\_\_\_truediv\_\_\_</u>

x.\_\_truediv\_\_(y)  $\leq x/y$ 

\_\_xor\_\_

 $x.\_xor\_(y) \leq x^y$ 

# argmax()

Returns the indices of the maximum along a given axis.

#### See also:

cupy.argmax() for full documentation, numpy.ndarray.argmax()

#### argmin()

Returns the indices of the minimum along a given axis.

#### See also:

cupy.argmin() for full documentation, numpy.ndarray.argmin()

#### astype()

Casts the array to given data type.

# Parameters

- dtype Type specifier.
- **copy** (bool) If it is False and no cast happens, then this method returns the array itself. Otherwise, a copy is returned.
- **Returns** If copy is False and no cast is required, then the array itself is returned. Otherwise, it returns a (possibly casted) copy of the array.

Note: This method currently does not support order, casting, and subok arguments.

#### See also:

numpy.ndarray.astype()

#### clip()

Returns an array with values limited to [a\_min, a\_max].

#### See also:

cupy.clip() for full documentation, numpy.ndarray.clip()

# copy()

Returns a copy of the array.

**Parameters order** ({ 'C', 'F'}) – Row-major (C-style) or column-major (Fortran-style) order. This function currently does not support order 'A' and 'K'.

#### See also:

cupy.copy() for full documentation, numpy.ndarray.copy()

# cstruct

C representation of the array.

This property is used for sending an array to CUDA kernels. The type of returned C structure is different for different dtypes and ndims. The definition of C type is written in cupy/carray.cuh.

# device

CUDA device on which this array resides.

#### diagonal()

Returns a view of the specified diagonals.

# See also:

cupy.diagonal() for full documentation, numpy.ndarray.diagonal()

#### dot ()

Returns the dot product with given array.

#### See also:

cupy.dot() for full documentation, numpy.ndarray.dot()

#### dump()

Dumps a pickle of the array to a file.

Dumped file can be read back to cupy.ndarray by cupy.load().

#### dumps()

Dumps a pickle of the array to a string.

#### fill()

Fills the array with a scalar value.

Parameters value - A scalar value to fill the array content.

See also:

numpy.ndarray.fill()

# flags

Object containing memory-layout information.

It only contains c\_contiguous, f\_contiguous, and owndata attributes. All of these are read-only. Accessing by indexes is also supported.

See also:

numpy.ndarray.flags

#### flatten()

Returns a copy of the array flatten into one dimension.

It currently supports C-order only.

**Returns** A copy of the array with one dimension.

Return type cupy.ndarray

# See also:

numpy.ndarray.flatten()

#### get()

Returns a copy of the array on host memory.

**Parameters stream** (cupy.cuda.Stream) – CUDA stream object. If it is given, the copy runs asynchronously. Otherwise, the copy is synchronous.

Returns Copy of the array on host memory.

Return type numpy.ndarray

# itemsize

Size of each element in bytes.

# See also:

numpy.ndarray.itemsize

## max()

Returns the maximum along a given axis.

#### See also:

cupy.amax() for full documentation, numpy.ndarray.max()

#### mean()

Returns the mean along a given axis.

# See also:

cupy.mean() for full documentation, numpy.ndarray.mean()

#### min()

Returns the minimum along a given axis.

#### See also:

cupy.amin() for full documentation, numpy.ndarray.min()

#### nbytes

Size of whole elements in bytes.

It does not count skips between elements.

#### See also:

numpy.ndarray.nbytes

# ndim

Number of dimensions.

a.ndim is equivalent to len (a.shape).

#### See also:

numpy.ndarray.ndim

#### nonzero()

Return the indices of the elements that are non-zero.

Returned Array is containing the indices of the non-zero elements in that dimension.

Returns Indices of elements that are non-zero.

#### **Return type** tuple of arrays

#### See also:

numpy.nonzero()

# prod()

Returns the product along a given axis.

#### See also:

cupy.prod() for full documentation, numpy.ndarray.prod()

#### ravel()

Returns an array flattened into one dimension.

# See also:

cupy.ravel() for full documentation, numpy.ndarray.ravel()

# reduced\_view()

Returns a view of the array with minimum number of dimensions.

**Parameters dtype** – Data type specifier. If it is given, then the memory sequence is reinterpreted as the new type.

Returns A view of the array with reduced dimensions.

# Return type cupy.ndarray

#### repeat()

Returns an array with repeated arrays along an axis.

#### See also:

cupy.repeat() for full documentation, numpy.ndarray.repeat()

# reshape()

Returns an array of a different shape and the same content.

# See also:

cupy.reshape() for full documentation, numpy.ndarray.reshape()

# scatter\_add()

Adds given values to specified elements of an array.

#### See also:

cupy.scatter\_add() for full documentation.

#### set()

Copies an array on the host memory to *cupy.ndarray*.

- **arr** (*numpy.ndarray*) The source array on the host memory.
- **stream** (cupy.cuda.Stream) CUDA stream object. If it is given, the copy runs asynchronously. Otherwise, the copy is synchronous.

#### shape

Lengths of axes.

Setter of this property involves reshaping without copy. If the array cannot be reshaped without copy, it raises an exception.

# squeeze()

Returns a view with size-one axes removed.

#### See also:

cupy.squeeze() for full documentation, numpy.ndarray.squeeze()

#### ${\tt std}\,(\,)$

Returns the standard deviation along a given axis.

#### See also:

cupy.std() for full documentation, numpy.ndarray.std()

## strides

Strides of axes in bytes.

# See also:

numpy.ndarray.strides

# ${\tt sum}\,(\,)$

Returns the sum along a given axis.

# See also:

cupy.sum() for full documentation, numpy.ndarray.sum()

#### swapaxes()

Returns a view of the array with two axes swapped.

## See also:

cupy.swapaxes() for full documentation, numpy.ndarray.swapaxes()

# take()

Returns an array of elements at given indices along the axis.

#### See also:

cupy.take() for full documentation, numpy.ndarray.take()

# tofile()

Writes the array to a file.

# See also:

numpy.ndarray.tolist()

#### tolist()

Converts the array to a (possibly nested) Python list.

Returns The possibly nested Python list of array elements.

Return type list

## See also:

numpy.ndarray.tolist()

#### trace()

Returns the sum along diagonals of the array.

# See also:

cupy.trace() for full documentation, numpy.ndarray.trace()

# transpose()

Returns a view of the array with axes permuted.

#### See also:

cupy.transpose() for full documentation, numpy.ndarray.reshape()

#### var()

Returns the variance along a given axis.

#### See also:

cupy.var() for full documentation, numpy.ndarray.var()

#### view()

Returns a view of the array.

**Parameters** dtype – If this is different from the data type of the array, the returned view reinterpret the memory sequence as an array of this type.

Returns A view of the array. A reference to the original array is stored at the base attribute.

#### Return type *cupy.ndarray*

#### See also:

numpy.ndarray.view()

#### cupy.asnumpy(a, stream=None)

Returns an array on the host memory from an arbitrary source array.

#### **Parameters**

- **a** Arbitrary object that can be converted to numpy.ndarray.
- **stream** (cupy.cuda.Stream) CUDA stream object. If it is specified, then the deviceto-host copy runs asynchronously. Otherwise, the copy is synchronous. Note that if a is not a *cupy.ndarray* object, then this argument has no effect.

Returns Converted array on the host memory.

Return type numpy.ndarray

# **Universal Functions (ufunc)**

CuPy provides universal functions (a.k.a. ufuncs) to support various elementwise operations. CuPy's ufunc supports following features of NumPy's one:

- Broadcasting
- Output type determination
- Casting rules

CuPy's ufunc currently does not provide methods such as reduce, accumulate, reduceat, outer, and at.

# **Ufunc class**

class cupy.ufunc

Universal function.

# Variables

- **name** (*str*) The name of the universal function.
- **nin** (*int*) Number of input arguments.
- **nout** (*int*) Number of output arguments.
- **nargs** (*int*) Number of all arguments.
- \_\_\_call\_\_()

Applies the universal function to arguments elementwise.

#### **Parameters**

- **args** Input arguments. Each of them can be a *cupy.ndarray* object or a scalar. The output arguments can be omitted or be specified by the out argument.
- **out** (cupy.ndarray) Output array. It outputs to new arrays default.
- **dtype** Data type specifier.

Returns Output array or a tuple of output arrays.

## types

A list of type signatures.

Each type signature is represented by type character codes of inputs and outputs separated by '->'.

# Available ufuncs

# Math operations

add subtract multiply divide logaddexp logaddexp2 true\_divide floor\_divide negative power remainder mod fmod absolute rint sign exp exp2 log log2 log10 expm1 log1p sqrt square reciprocal

# **Trigonometric functions**

sin cos tan arcsin arccos arctan arctan2 hypot sinh cosh tanh arcsinh arccosh arctanh deg2rad rad2deg

# **Bit-twiddling 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 fmax fmin

# **Floating point values**

isfinite isinf isnan signbit copysign nextafter modf ldexp frexp fmod floor ceil trunc

# ufunc.at

Currently, CuPy does not support at for ufuncs in general. However, *cupy.scatter\_add()* can substitute add. at as both behave identically.

# **Routines**

The following pages describe NumPy-compatible routines. These functions cover a subset of NumPy routines.

# **Array Creation Routines**

# **Basic creation routines**

```
cupy.empty(shape, dtype=<type 'float'>, order='C')
```

Returns an array without initializing the elements.

#### Parameters

- **shape** (*tuple of ints*) Dimensionalities of the array.
- **dtype** Data type specifier.
- order ({ 'C', 'F'}) Row-major (C-style) or column-major (Fortran-style) order.

**Returns** A new array with elements not initialized.

Return type *cupy.ndarray* 

# See also:

```
numpy.empty()
```

```
cupy.empty_like(a, dtype=None)
```

Returns a new array with same shape and dtype of a given array.

This function currently does not support order and subok options.

#### **Parameters**

- **a** (cupy.ndarray) Base array.
- dtype Data type specifier. The data type of a is used by default.

Returns A new array with same shape and dtype of a with elements not initialized.

Return type *cupy.ndarray* 

#### See also:

```
numpy.empty_like()
```

cupy.eye (N, M=None, k=0, dtype=<type 'float'>)

Returns a 2-D array with ones on the diagonals and zeros elsewhere.

- N (*int*) Number of rows.
- M(int) Number of columns. M == N by default.
- **k** (*int*) Index of the diagonal. Zero indicates the main diagonal, a positive index an upper diagonal, and a negative index a lower diagonal.
- **dtype** Data type specifier.

Returns A 2-D array with given diagonals filled with ones and zeros elsewhere.

Return type *cupy.ndarray* 

# See also:

numpy.eye()

```
cupy.identity (n, dtype=<type 'float'>)
Returns a 2-D identity array.
```

It is equivalent to eye (n, n, dtype).

# Parameters

- **n** (*int*) Number of rows and columns.
- **dtype** Data type specifier.

Returns A 2-D identity array.

Return type *cupy.ndarray* 

#### See also:

```
numpy.identity()
```

```
cupy.ones(shape, dtype=<type 'float'>)
```

Returns a new array of given shape and dtype, filled with ones.

This function currently does not support order option.

## Parameters

- **shape** (*tuple of ints*) Dimensionalities of the array.
- **dtype** Data type specifier.

**Returns** An array filled with ones.

Return type *cupy.ndarray* 

# See also:

numpy.ones()

#### cupy.ones\_like(a, dtype=None)

Returns an array of ones with same shape and dtype as a given array.

This function currently does not support order and subok options.

# Parameters

- **a** (cupy.ndarray) Base array.
- dtype Data type specifier. The dtype of a is used by default.

**Returns** An array filled with ones.

Return type *cupy.ndarray* 

## See also:

numpy.ones\_like()

cupy.zeros (shape, dtype=<type 'float'>, order='C')

Returns a new array of given shape and dtype, filled with zeros.

# Parameters

- **shape** (tuple of ints) Dimensionalities of the array.
- **dtype** Data type specifier.
- order ({ 'C', 'F'}) Row-major (C-style) or column-major (Fortran-style) order.

Returns An array filled with ones.

**Return type** *cupy.ndarray* 

# See also:

numpy.zeros()

#### cupy.zeros\_like(*a*, *dtype=None*)

Returns an array of zeros with same shape and dtype as a given array.

This function currently does not support order and subok options.

# Parameters

- **a** (cupy.ndarray) Base array.
- dtype Data type specifier. The dtype of a is used by default.

Returns An array filled with ones.

# **Return type** *cupy.ndarray*

#### See also:

```
numpy.zeros_like()
```

# cupy.full(shape, fill\_value, dtype=None)

Returns a new array of given shape and dtype, filled with a given value.

This function currently does not support order option.

# Parameters

- **shape** (tuple of ints) Dimensionalities of the array.
- **fill\_value** A scalar value to fill a new array.
- **dtype** Data type specifier.

**Returns** An array filled with fill\_value.

Return type *cupy.ndarray* 

# See also:

numpy.full()

# cupy.full\_like(a, fill\_value, dtype=None)

Returns a full array with same shape and dtype as a given array.

This function currently does not support order and subok options.

- a (cupy.ndarray) Base array.
- fill\_value A scalar value to fill a new array.
- dtype Data type specifier. The dtype of a is used by default.

**Returns** An array filled with fill\_value.

Return type *cupy.ndarray* 

#### See also:

numpy.full\_like()

# Creation from other data

cupy.array(obj, dtype=None, copy=True, ndmin=0)

Creates an array on the current device.

This function currently does not support the order and subok options.

# Parameters

- **obj** *cupy.ndarray* object or any other object that can be passed to numpy. array().
- **dtype** Data type specifier.
- **copy** (*bool*) If False, this function returns obj if possible. Otherwise this function always returns a new array.
- **ndmin** (*int*) Minimum number of dimensions. Ones are inserted to the head of the shape if needed.

Returns An array on the current device.

Return type *cupy.ndarray* 

# See also:

numpy.array()

```
cupy.asarray(a, dtype=None)
```

Converts an object to array.

This is equivalent to array(a, dtype, copy=False). This function currently does not support the order option.

#### **Parameters**

- **a** The source object.
- **dtype** Data type specifier. It is inferred from the input by default.

Returns An array on the current device. If a is already on the device, no copy is performed.

Return type *cupy.ndarray* 

# See also:

numpy.asarray()

# cupy.asanyarray(a, dtype=None)

Converts an object to array.

This is currently equivalent to *asarray()*, since there is no subclass of ndarray in CuPy. Note that the original numpy.asanyarray() returns the input array as is if it is an instance of a subtype of numpy.ndarray.

# See also:

cupy.asarray(), numpy.asanyarray()

cupy.ascontiguousarray(a, dtype=None)

Returns a C-contiguous array.

# Parameters

- a (cupy.ndarray) Source array.
- **dtype** Data type specifier.

Returns If no copy is required, it returns a. Otherwise, it returns a copy of a.

Return type *cupy.ndarray* 

#### See also:

```
numpy.ascontiguousarray()
```

cupy.copy(\*args, \*\*kwargs)

# **Numerical ranges**

```
cupy.arange (start, stop=None, step=1, dtype=None)
```

Returns an array with evenly spaced values within a given interval.

Values are generated within the half-open interval [start, stop). The first three arguments are mapped like the range built-in function, i.e. start and step are optional.

# Parameters

- **start** Start of the interval.
- **stop** End of the interval.
- **step** Step width between each pair of consecutive values.
- dtype Data type specifier. It is inferred from other arguments by default.

Returns The 1-D array of range values.

**Return type** *cupy.ndarray* 

#### See also:

numpy.arange()

cupy.linspace (start, stop, num=50, endpoint=True, retstep=False, dtype=None)

Returns an array with evenly-spaced values within a given interval.

Instead of specifying the step width like *cupy.arange()*, this function requires the total number of elements specified.

- **start** Start of the interval.
- **stop** End of the interval.
- **num** Number of elements.
- **endpoint** (*bool*) If True, the stop value is included as the last element. Otherwise, the stop value is omitted.

- **retstep** (*bool*) If True, this function returns (array, step). Otherwise, it returns only the array.
- dtype Data type specifier. It is inferred from the start and stop arguments by default.

Returns The 1-D array of ranged values.

## Return type *cupy.ndarray*

cupy.logspace (*start*, *stop*, *num=50*, *endpoint=True*, *base=10.0*, *dtype=None*) Returns an array with evenly-spaced values on a log-scale.

Instead of specifying the step width like *cupy.arange()*, this function requires the total number of elements specified.

#### **Parameters**

- **start** Start of the interval.
- **stop** End of the interval.
- **num** Number of elements.
- **endpoint** (*bool*) If True, the stop value is included as the last element. Otherwise, the stop value is omitted.
- **base** (*float*) Base of the log space. The step sizes between the elements on a log-scale are the same as base.
- dtype Data type specifier. It is inferred from the start and stop arguments by default.

Returns The 1-D array of ranged values.

# Return type *cupy.ndarray*

# cupy.meshgrid(\*xi, \*\*kwargs)

Return coordinate matrices from coordinate vectors.

Given one-dimensional coordinate arrays x1, x2, ..., xn, this function makes N-D grids.

For one-dimensional arrays x1, x2, ..., xn with lengths Ni = len(xi), this function returns (N1, N2, N3, ..., Nn) shaped arrays if indexing='ij' or (N2, N1, N3, ..., Nn) shaped arrays if indexing='xy'.

Unlike NumPy, CuPy currently only supports 1-D arrays as inputs. Also, CuPy does not support sparse option yet.

#### **Parameters**

- **xi** (tuple of ndarrays) 1-D arrays representing the coordinates of a grid.
- **indexing** ({'xy', 'ij'}, optional) Cartesian ('xy', default) or matrix ('ij') indexing of output.
- **copy** (*bool*, *optional*) If False, a view into the original arrays are returned. Default is True.

Returns list of cupy.ndarray

# See also:

numpy.meshgrid()

# Matrix creation

cupy.**diag**(v, k=0)

Returns a diagonal or a diagonal array.

#### Parameters

- **v** (*array-like*) Array or array-like object.
- **k** (*int*) Index of diagonals. Zero indicates the main diagonal, a positive value an upper diagonal, and a negative value a lower diagonal.
- **Returns** If v indicates a 1-D array, then it returns a 2-D array with the specified diagonal filled by v. If v indicates a 2-D array, then it returns the specified diagonal of v. In latter case, if v is a *cupy.ndarray* object, then its view is returned.

Return type *cupy.ndarray* 

# See also:

numpy.diag()

#### cupy.diagflat(v, k=0)

Creates a diagonal array from the flattened input.

#### **Parameters**

- **v** (*array-like*) Array or array-like object.
- **k** (*int*) Index of diagonals. See *cupy.diag()* for detail.

Returns A 2-D diagonal array with the diagonal copied from v.

**Return type** *cupy.ndarray* 

# **Array Manipulation Routines**

#### **Basic manipulations**

```
cupy.copyto(dst, src, casting='same_kind', where=None)
```

Copies values from one array to another with broadcasting.

This function can be called for arrays on different devices. In this case, casting, where, and broadcasting is not supported, and an exception is raised if these are used.

#### **Parameters**

- dst (cupy.ndarray) Target array.
- **src** (cupy.ndarray) Source array.
- **casting** (*str*) **Casting rule**. See numpy.can\_cast() for detail.
- where (*cupy.ndarray of bool*) If specified, this array acts as a mask, and an element is copied only if the corresponding element of where is True.

# See also:

```
numpy.copyto()
```

# Shape manipulation

#### cupy.**reshape**(*a*, *newshape*)

Returns an array with new shape and same elements.

It tries to return a view if possible, otherwise returns a copy.

This function currently does not support order option.

# Parameters

- a (cupy.ndarray) Array to be reshaped.
- **newshape** (*int* or tuple of *ints*) The new shape of the array to return. If it is an integer, then it is treated as a tuple of length one. It should be compatible with a.size. One of the elements can be -1, which is automatically replaced with the appropriate value to make the shape compatible with a.size.

**Returns** A reshaped view of a if possible, otherwise a copy.

Return type *cupy.ndarray* 

#### See also:

numpy.reshape()

# cupy.ravel(a)

Returns a flattened array.

It tries to return a view if possible, otherwise returns a copy.

This function currently does not support order option.

Parameters a (cupy.ndarray) – Array to be flattened.

**Returns** A flattened view of a if possible, otherwise a copy.

Return type *cupy.ndarray* 

#### See also:

numpy.ravel()

# Transposition

cupy.rollaxis(a, axis, start=0)

Moves the specified axis backwards to the given place.

#### Parameters

- **a** (cupy.ndarray) Array to move the axis.
- **axis** (*int*) The axis to move.
- **start** (*int*) The place to which the axis is moved.

Returns A view of a that the axis is moved to start.

#### **Return type** *cupy.ndarray*

# See also:

# numpy.rollaxis()

cupy.swapaxes (a, axis1, axis2)

Swaps the two axes.

# Parameters

- a (cupy.ndarray) Array to swap the axes.
- **axis1** (*int*) The first axis to swap.
- **axis2** (*int*) The second axis to swap.

Returns A view of a that the two axes are swapped.

#### Return type cupy.ndarray

#### See also:

numpy.swapaxes()

```
cupy.transpose (a, axes=None)
Permutes the dimensions of an array.
```

#### **Parameters**

- a (cupy.ndarray) Array to permute the dimensions.
- **axes** (*tuple of ints*) Permutation of the dimensions. This function reverses the shape by default.

**Returns** A view of a that the dimensions are permuted.

Return type *cupy.ndarray* 

# See also:

numpy.transpose()

# **Edit dimensionalities**

#### cupy.atleast\_1d(\*arys)

Converts arrays to arrays with dimensions >= 1.

- **Parameters arys** (*tuple of arrays*) Arrays to be converted. All arguments must be *cupy.ndarray* objects. Only zero-dimensional array is affected.
- **Returns** If there are only one input, then it returns its converted version. Otherwise, it returns a list of converted arrays.

#### See also:

numpy.atleast\_1d()

#### cupy.atleast\_2d(\*arys)

Converts arrays to arrays with dimensions  $\geq 2$ .

If an input array has dimensions less than two, then this function inserts new axes at the head of dimensions to make it have two dimensions.

**Parameters arys** (*tuple of arrays*) – Arrays to be converted. All arguments must be *cupy.ndarray* objects.

**Returns** If there are only one input, then it returns its converted version. Otherwise, it returns a list of converted arrays.

#### See also:

numpy.atleast\_2d()

#### cupy.atleast\_3d(\*arys)

Converts arrays to arrays with dimensions  $\geq 3$ .

If an input array has dimensions less than three, then this function inserts new axes to make it have three dimensions. The place of the new axes are following:

•If its shape is (), then the shape of output is (1, 1, 1).

•If its shape is (N, ), then the shape of output is (1, N, 1).

•If its shape is (M, N), then the shape of output is (M, N, 1).

•Otherwise, the output is the input array itself.

- **Parameters arys** (*tuple of arrays*) Arrays to be converted. All arguments must be *cupy.ndarray* objects.
- **Returns** If there are only one input, then it returns its converted version. Otherwise, it returns a list of converted arrays.

#### See also:

numpy.atleast\_3d()

# class cupy.broadcast

Object that performs broadcasting.

CuPy actually uses this class to support broadcasting in various operations. Note that this class does not provide an iterator.

**Parameters arrays** (tuple of arrays) – Arrays to be broadcasted.

Variables

- **shape** (tuple of ints) The broadcasted shape.
- **nd** (*int*) Number of dimensions of the broadcasted shape.
- **size** (*int*) Total size of the broadcasted shape.
- **values** (*list of arrays*) The broadcasted arrays.

#### See also:

numpy.broadcast

#### cupy.broadcast\_arrays(\*args)

Broadcasts given arrays.

**Parameters args** (*tuple of arrays*) – Arrays to broadcast for each other.

**Returns** A list of broadcasted arrays.

**Return type** list

See also:

#### numpy.broadcast\_arrays()

cupy.broadcast\_to(array, shape)

Broadcast an array to a given shape.

# Parameters

- array (cupy.ndarray) Array to broadcast.
- **shape** (*tuple of int*) The shape of the desired array.

Returns Broadcasted view.

Return type *cupy.ndarray* 

#### See also:

numpy.broadcast\_to()

cupy.expand\_dims (a, axis) Expands given arrays.

- a (cupy.ndarray) Array to be expanded.
- **axis** (*int*) Position where new axis is to be inserted.

# Returns

# The number of dimensions is one greater than that of the input array.

**Return type** *cupy.ndarray* 

#### See also:

numpy.expand\_dims()

#### cupy.squeeze(a, axis=None)

Removes size-one axes from the shape of an array.

#### Parameters

- a (cupy.ndarray) Array to be reshaped.
- **axis** (*int* or *tuple* of *ints*) Axes to be removed. This function removes all size-one axes by default. If one of the specified axes is not of size one, an exception is raised.

Returns An array without (specified) size-one axes.

Return type *cupy.ndarray* 

#### See also:

numpy.squeeze()

#### Changing kind of array

#### cupy.asfortranarray(a, dtype=None)

Return an array laid out in Fortran order in memory.

#### **Parameters**

- **a** (ndarray) The input array.
- **dtype** (*str* or *dtype* object, *optional*) By default, the data-type is inferred from the input data.

**Returns** The input *a* in Fortran, or column-major, order.

#### **Return type** *ndarray*

## See also:

```
numpy.asfortranarray()
```

# Joining arrays along axis

```
cupy.column_stack(tup)
```

Stacks 1-D and 2-D arrays as columns into a 2-D array.

A 1-D array is first converted to a 2-D column array. Then, the 2-D arrays are concatenated along the second axis.

```
Parameters tup (sequence of arrays) - 1-D or 2-D arrays to be stacked.
```

Returns A new 2-D array of stacked columns.

#### Return type *cupy.ndarray*

#### See also:

numpy.column\_stack()

cupy.concatenate(tup, axis=0)

Joins arrays along an axis.

# Parameters

- **tup** (*sequence of arrays*) Arrays to be joined. All of these should have same dimensionalities except the specified axis.
- **axis** (*int*) The axis to join arrays along.

Returns Joined array.

Return type cupy.ndarray

## See also:

numpy.concatenate()

#### cupy.vstack(tup)

Stacks arrays vertically.

If an input array has one dimension, then the array is treated as a horizontal vector and stacked along the additional axis at the head. Otherwise, the array is stacked along the first axis.

**Parameters tup** (*sequence of arrays*) – Arrays to be stacked. Each array is converted by *cupy.atleast\_2d()* before stacking.

Returns Stacked array.

Return type cupy.ndarray

#### See also:

numpy.dstack()

#### cupy.hstack(tup)

Stacks arrays horizontally.

If an input array has one dimension, then the array is treated as a horizontal vector and stacked along the first axis. Otherwise, the array is stacked along the second axis.

**Parameters** tup (sequence of arrays) – Arrays to be stacked.

Returns Stacked array.

Return type cupy.ndarray

#### See also:

numpy.hstack()

cupy.dstack(*tup*)

Stacks arrays along the third axis.

**Parameters tup** (sequence of arrays) – Arrays to be stacked. Each array is converted by cupy.atleast\_3d() before stacking.

Returns Stacked array.

Return type *cupy.ndarray* 

#### See also:

numpy.dstack()

cupy.**stack**(*tup*, *axis=0*)

Stacks arrays along a new axis.

# Parameters

- tup (sequence of arrays) Arrays to be stacked.
- **axis** (*int*) Axis along which the arrays are stacked.

Returns Stacked array.

Return type *cupy.ndarray* 

#### See also:

numpy.stack()

# Splitting arrays along axis

#### cupy.array\_split (ary, indices\_or\_sections, axis=0)

Splits an array into multiple sub arrays along a given axis.

This function is almost equivalent to *cupy.split()*. The only difference is that this function allows an integer sections that does not evenly divide the axis.

# See also:

cupy.split() for more detail, numpy.array\_split()

## cupy.split (ary, indices\_or\_sections, axis=0)

Splits an array into multiple sub arrays along a given axis.

# Parameters

- ary (cupy.ndarray) Array to split.
- **indices\_or\_sections** (*int or sequence of ints*) A value indicating how to divide the axis. If it is an integer, then is treated as the number of sections, and the axis is evenly divided. Otherwise, the integers indicate indices to split at. Note that the sequence on the device memory is not allowed.
- **axis** (*int*) Axis along which the array is split.

**Returns** A list of sub arrays. Each array is a view of the corresponding input array.

#### See also:

```
numpy.split()
```

```
cupy.vsplit(ary, indices_or_sections)
```

Splits an array into multiple sub arrays along the first axis.

This is equivalent to split with axis=0.

# See also:

cupy.split() for more detail, numpy.dsplit()

# cupy.hsplit(ary, indices\_or\_sections)

Splits an array into multiple sub arrays horizontally.

This is equivalent to split with axis=0 if ary has one dimension, and otherwise that with axis=1.

# See also:

cupy.split() for more detail, numpy.hsplit()

cupy.dsplit (ary, indices\_or\_sections)

Splits an array into multiple sub arrays along the third axis.

This is equivalent to split with axis=2.

# See also:

```
cupy.split() for more detail, numpy.dsplit()
```

# Repeating part of arrays along axis

```
cupy.tile(A, reps)
```

Construct an array by repeating A the number of times given by reps.

#### **Parameters**

- A (cupy.ndarray) Array to transform.
- **reps** (*int* or *tuple*) The number of repeats.

Returns Transformed array with repeats.

Return type *cupy.ndarray* 

#### See also:

```
numpy.tile()
```

cupy.repeat (a, repeats, axis=None)

Repeat arrays along an axis.

# Parameters

- **a** (cupy.ndarray) Array to transform.
- repeats (int, list or tuple) The number of repeats.
- **axis** (*int*) The axis to repeat.

Returns Transformed array with repeats.

**Return type** *cupy.ndarray* 

#### See also:

numpy.repeat()

# **Rearranging elements**

cupy.flip(a, axis)

Reverse the order of elements in an array along the given axis.

Note that flip function has been introduced since NumPy v1.12. The contents of this document is the same as the original one.

- **a** (ndarray) Input array.
- **axis** (*int*) Axis in array, which entries are reversed.

Returns Output array.

Return type *ndarray* 

# See also:

numpy.flip()

cupy.fliplr(a)

Flip array in the left/right direction.

Flip the entries in each row in the left/right direction. Columns are preserved, but appear in a different order than before.

Parameters a (ndarray) – Input array.

Returns Output array.

Return type *ndarray* 

See also:

numpy.fliplr()

# cupy.flipud(a)

Flip array in the up/down direction.

Flip the entries in each column in the up/down direction. Rows are preserved, but appear in a different order than before.

Parameters a (ndarray) – Input array.

Returns Output array.

Return type *ndarray* 

#### See also:

numpy.flipud()

cupy.roll(a, shift, axis=None)

Roll array elements along a given axis.

#### Parameters

- a (ndarray) Array to be rolled.
- **shift** (*int*) The number of places by which elements are shifted.
- **axis** (*int* or *None*) The axis along which elements are shifted. If axis is None, the array is flattened before shifting, and after that it is reshaped to the original shape.

Returns Output array.

Return type *ndarray* 

# See also:

```
numpy.roll()
```

cupy.**rot90** (*a*, *k*=1, *axes*=(0, 1))

Rotate an array by 90 degrees in the plane specified by axes.

Note that axes argument has been introduced since NumPy v1.12. The contents of this document is the same as the original one.

# Parameters

• **a** (ndarray) – Array of two or more dimensions.

- **k** (*int*) Number of times the array is rotated by 90 degrees.
- **axes** (tuple of ints): The array is rotated in the plane defined by the axes. Axes must be different.

Returns Output array.

Return type *ndarray* 

#### See also:

numpy.rot90()

# **Binary Operations**

# **Elementwise bit operations**

# cupy.bitwise\_and = <ufunc 'cupy\_bitwise\_and'>

Computes the bitwise AND of two arrays elementwise.

Only integer and boolean arrays are handled.

See also:

numpy.bitwise\_and

# cupy.bitwise\_or = <ufunc 'cupy\_bitwise\_or'>

Computes the bitwise OR of two arrays elementwise.

Only integer and boolean arrays are handled.

# See also:

numpy.bitwise\_or

# cupy.bitwise\_xor = <ufunc 'cupy\_bitwise\_xor'> Computes the bitwise XOR of two arrays elementwise.

Only integer and boolean arrays are handled.

See also:

numpy.bitwise\_xor

#### cupy.invert = <ufunc 'cupy\_invert'>

Computes the bitwise NOT of an array elementwise.

Only integer and boolean arrays are handled.

#### See also:

numpy.invert

# cupy.left\_shift = <ufunc 'cupy\_left\_shift'>

Shifts the bits of each integer element to the left.

Only integer arrays are handled.

See also:

#### numpy.left\_shift

cupy.right\_shift = <ufunc 'cupy\_right\_shift'>
 Shifts the bits of each integer element to the right.

Only integer arrays are handled

## See also:

numpy.right\_shift

# **Indexing Routines**

```
cupy.take(a, indices, axis=None, out=None)
```

Takes elements of an array at specified indices along an axis.

This is an implementation of "fancy indexing" at single axis.

This function does not support mode option.

# Parameters

- a (cupy.ndarray) Array to extract elements.
- indices (int or array-like) Indices of elements that this function takes.
- **axis** (*int*) The axis along which to select indices. The flattened input is used by default.
- **out** (cupy.ndarray) Output array. If provided, it should be of appropriate shape and dtype.

Returns The result of fancy indexing.

Return type cupy.ndarray

#### See also:

numpy.take()

```
cupy.diagonal(a, offset=0, axis1=0, axis2=1)
```

Returns specified diagonals.

This function extracts the diagonals along two specified axes. The other axes are not changed. This function returns a writable view of this array as NumPy 1.10 will do.

# Parameters

- a (cupy.ndarray) Array from which the diagonals are taken.
- **offset** (*int*) Index of the diagonals. Zero indicates the main diagonals, a positive value upper diagonals, and a negative value lower diagonals.
- **axis1** (*int*) The first axis to take diagonals from.
- **axis2** (*int*) The second axis to take diagonals from.

Returns A view of the diagonals of a.

# Return type cupy.ndarray

#### See also:

```
numpy.diagonal()
```

```
cupy.ix_(*args)
```

Construct an open mesh from multiple sequences.

This function takes N 1-D sequences and returns N outputs with N dimensions each, such that the shape is 1 in all but one dimension and the dimension with the non-unit shape value cycles through all N dimensions.

Using *ix*\_one can quickly construct index arrays that will index the cross product. a[cupy.ix\_([1,3], [2,5])] returns the array [[a[1,2] a[1,5]], [a[3,2] a[3,5]]].

**Parameters \*args** – 1-D sequences

**Returns** N arrays with N dimensions each, with N the number of input sequences. Together these arrays form an open mesh.

Return type tuple of ndarrays

# **Examples**

```
>>> a = cupy.arange(10).reshape(2, 5)
>>> a
array([[0, 1, 2, 3, 4],
        [5, 6, 7, 8, 9]])
>>> ixgrid = cupy.ix_([0,1], [2,4])
>>> ixgrid
(array([[0],
        [1]]), array([[2, 4]]))
```

See also:

numpy.ix\_()

#### cupy.fill\_diagonal (a, val, wrap=False)

Fill the main diagonal of the given array of any dimensionality.

For an array a with a.ndim > 2, the diagonal is the list of locations with indices a[i, i, ..., i] all identical. This function modifies the input array in-place, it does not return a value.

# Parameters

- a (cupy.ndarray) The array, at least 2-D.
- **val** (*scalar*) The value to be written on the diagonal. Its type must be compatible with that of the array a.
- wrap (bool) If specified, the diagonal is "wrapped" after N columns. This affects only tall matrices.

#### **Examples**

# See also:

numpy.fill\_diagonal()

# cupy.c\_ = <cupy.indexing.generate.CClass object>

Translates slice objects to concatenation along the second axis.

This is a CuPy object that corresponds to  $cupy.r_()$ , which is useful because of its common occurrence. In particular, arrays will be stacked along their last axis after being upgraded to at least 2-D with 1's post-pended to the shape (column vectors made out of 1-D arrays).

For detailed documentation, see  $r_()$ .

This implementation is partially borrowed from NumPy's one.

```
Parameters a function, so takes no parameters (Not) -
```

Returns Joined array.

Return type *cupy.ndarray* 

#### See also:

numpy.c\_()

# **Examples**

```
>>> a = cupy.array([[1, 2, 3]], dtype=np.int32)
>>> b = cupy.array([[4, 5, 6]], dtype=np.int32)
>>> cupy.c_[a, 0, 0, b]
array([[1, 2, 3, 0, 0, 4, 5, 6]], dtype=int32)
```

## cupy.r\_ = <cupy.indexing.generate.RClass object>

Translates slice objects to concatenation along the first axis.

This is a simple way to build up arrays quickly. If the index expression contains comma separated arrays, then stack them along their first axis.

This object can build up from normal CuPy arrays. Therefore, the other objects (e.g. writing strings like '2,3,4', or using imaginary numbers like [1,2,3j], or using string integers like '-1') are not implemented yet compared with NumPy.

This implementation is partially borrowed from NumPy's one.

```
Parameters a function, so takes no parameters (Not) -
```

Returns Joined array.

Return type cupy.ndarray

# See also:

numpy.r\_()

# **Examples**

```
>>> a = cupy.array([1, 2, 3], dtype=np.int32)
>>> b = cupy.array([4, 5, 6], dtype=np.int32)
>>> cupy.r_[a, 0, 0, b]
array([1, 2, 3, 0, 0, 4, 5, 6], dtype=int32)
```

# Input and Output

# **NPZ** files

```
cupy.load(file, mmap_mode=None)
```

Loads arrays or pickled objects from .npy, .npz or pickled file.

This function just calls numpy.load and then sends the arrays to the current device. NPZ file is converted to NpzFile object, which defers the transfer to the time of accessing the items.

# Parameters

- file (file-like object or string) The file to read.
- mmap\_mode (*None*, 'r+', 'r', 'w+', 'c') If not None, memory-map the file to construct an intermediate numpy.ndarray object and transfer it to the current device.
- **Returns** CuPy array or NpzFile object depending on the type of the file. NpzFile object is a dictionary-like object with the context manager protocol (which enables us to use *with* statement on it).

# See also:

numpy.load()

```
cupy.save (file, arr)
```

Saves an array to a binary file in .npy format.

#### **Parameters**

- file (file or str) File or filename to save.
- **arr** (*array\_like*) Array to save. It should be able to feed to *cupy.asnumpy()*.

#### See also:

#### numpy.save()

#### cupy.savez (file, \*args, \*\*kwds)

Saves one or more arrays into a file in uncompressed .npz format.

Arguments without keys are treated as arguments with automatic keys named arr\_0, arr\_1, etc. corresponding to the positions in the argument list. The keys of arguments are used as keys in the .npz file, which are used for accessing NpzFile object when the file is read by *cupy.load()* function.

#### **Parameters**

- file (file or str) File or filename to save.
- **\*args** Arrays with implicit keys.
- **\*\*kwds** Arrays with explicit keys.

#### See also:

#### numpy.savez()

#### cupy.savez\_compressed (file, \*args, \*\*kwds)

Saves one or more arrays into a file in compressed .npz format.

It is equivalent to *cupy.savez()* function except the output file is compressed.

#### See also:

cupy.savez() for more detail, numpy.savez\_compressed()

# String formatting

cupy.array\_repr (arr, max\_line\_width=None, precision=None, suppress\_small=None) Returns the string representation of an array.

- **arr** (*array\_like*) Input array. It should be able to feed to *cupy.asnumpy()*.
- **max\_line\_width** (*int*) The maximum number of line lengths.

- **precision** (*int*) Floating point precision. It uses the current printing precision of NumPy.
- suppress\_small (bool) If True, very small numbers are printed as zeros

Returns The string representation of arr.

Return type str

#### See also:

```
numpy.array_repr()
```

cupy.array\_str(*arr*, *max\_line\_width=None*, *precision=None*, *suppress\_small=None*) Returns the string representation of the content of an array.

# Parameters

- **arr** (*array\_like*) Input array. It should be able to feed to *cupy.asnumpy()*.
- **max\_line\_width** (*int*) The maximum number of line lengths.
- **precision** (*int*) Floating point precision. It uses the current printing precision of NumPy.
- **suppress\_small** (*bool*) If True, very small number are printed as zeros.

# See also:

numpy.array\_str()

# **Linear Algebra**

# Matrix and vector products

```
cupy.dot (a, b, out=None)
```

Returns a dot product of two arrays.

For arrays with more than one axis, it computes the dot product along the last axis of a and the second-to-last axis of b. This is just a matrix product if the both arrays are 2-D. For 1-D arrays, it uses their unique axis as an axis to take dot product over.

#### Parameters

- **a** (cupy.ndarray) The left argument.
- **b** (cupy.ndarray) The right argument.
- **out** (cupy.ndarray) **Output** array.

**Returns** The dot product of a and b.

Return type *cupy.ndarray* 

# See also:

```
numpy.dot()
```

```
cupy.vdot (a, b)
```

Returns the dot product of two vectors.

The input arrays are flattened into 1-D vectors and then it performs inner product of these vectors.

#### Parameters

• **a** (cupy.ndarray) – The first argument.

• **b** (cupy.ndarray) – The second argument.

Returns Zero-dimensional array of the dot product result.

Return type *cupy.ndarray* 

# See also:

numpy.vdot()

#### cupy.**inner**(*a*, *b*)

Returns the inner product of two arrays.

It uses the last axis of each argument to take sum product.

#### Parameters

- **a** (cupy.ndarray) The first argument.
- **b** (cupy.ndarray) The second argument.

**Returns** The inner product of a and b.

Return type cupy.ndarray

#### See also:

numpy.inner()

cupy.outer(*a*, *b*, *out=None*)

Returns the outer product of two vectors.

The input arrays are flattened into 1-D vectors and then it performs outer product of these vectors.

#### **Parameters**

- a (cupy.ndarray) The first argument.
- **b** (cupy.ndarray) The second argument.
- **out** (cupy.ndarray) **Output** array.

**Returns** 2-D array of the outer product of a and b.

Return type *cupy.ndarray* 

# See also:

numpy.outer()

# cupy.tensordot (a, b, axes=2)

Returns the tensor dot product of two arrays along specified axes.

This is equivalent to compute dot product along the specified axes which are treated as one axis by reshaping.

- a (cupy.ndarray) The first argument.
- **b** (cupy.ndarray) The second argument.
- axes -
  - If it is an integer, then axes axes at the last of a and the first of b are used.
  - If it is a pair of sequences of integers, then these two sequences specify the list of axes for a and b. The corresponding axes are paired for sum-product.
- out (cupy.ndarray) Output array.

**Returns** The tensor dot product of a and b along the axes specified by axes.

**Return type** *cupy.ndarray* 

# See also:

numpy.tensordot()

# **Decompositions**

cupy.linalg.cholesky(a)

Cholesky decomposition.

Decompose a given two-dimensional square matrix into L \* L.T, where L is a lower-triangular matrix and . T is a conjugate transpose operator. Note that in the current implementation a must be a real matrix, and only float32 and float64 are supported.

Parameters a (cupy.ndarray) - The input matrix with dimension (N, N)

# See also:

```
numpy.linalg.cholesky()
```

cupy.linalg.gr(a, mode='reduced')

QR decomposition.

Decompose a given two-dimensional matrix into Q \* R, where Q is an orthonormal and R is an upper-triangular matrix.

# **Parameters**

- **a** (cupy.ndarray) The input matrix.
- mode (str) The mode of decomposition. Currently 'reduced', 'complete', 'r', and 'raw' modes are supported. The default mode is 'reduced', and decompose a matrix A = (M, N) into Q, R with dimensions (M, K), (K, N), where K = min (M, N).

# See also:

```
numpy.linalg.qr()
```

```
cupy.linalg.svd(a, full_matrices=True, compute_uv=True)
```

Singular Value Decomposition.

Factorizes the matrix a as u \* np.diag(s) \* v, where u and v are unitary and s is an one-dimensional array of a's singular values.

#### Parameters

- a (cupy.ndarray) The input matrix with dimension (M, N).
- full\_matrices (bool) If True, it returns U and V with dimensions (M, M) and (N, N). Otherwise, the dimensions of U and V are respectively (M, K) and (K, N), where K = min (M, N).
- **compute\_uv** (*bool*) If True, it only returns singular values.

# See also:

```
numpy.linalg.svd()
```

# Norms etc.

cupy.trace(a, offset=0, axis1=0, axis2=1, dtype=None, out=None)
Returns the sum along the diagonals of an array.

It computes the sum along the diagonals at axis1 and axis2.

# Parameters

- **a** (cupy.ndarray) Array to take trace.
- **offset** (*int*) Index of diagonals. Zero indicates the main diagonal, a positive value an upper diagonal, and a negative value a lower diagonal.
- **axis1** (*int*) The first axis along which the trace is taken.
- **axis2** (*int*) The second axis along which the trace is taken.
- **dtype** Data type specifier of the output.
- **out** (cupy.ndarray) **Output** array.

Returns The trace of a along axes (axis1, axis2).

Return type *cupy.ndarray* 

# See also:

numpy.trace()

# **Logic Functions**

# **Infinities and NaNs**

# cupy.isfinite = <ufunc 'cupy\_isfinite'>

Tests finiteness elementwise.

Each element of returned array is True only if the corresponding element of the input is finite (i.e. not an infinity nor NaN).

# See also:

numpy.isfinite

#### cupy.isinf = <ufunc 'cupy\_isinf'>

Tests if each element is the positive or negative infinity.

#### See also:

numpy.isinf

```
cupy.isnan = <ufunc 'cupy_isnan'>
Tests if each element is a NaN.
```

#### See also:

numpy.isnan

# Logic operations

cupy.logical\_and = <ufunc 'cupy\_logical\_and'> Computes the logical AND of two arrays.

# See also:

numpy.logical\_and

cupy.logical\_or = <ufunc 'cupy\_logical\_or'> Computes the logical OR of two arrays.

See also:

numpy.logical\_or

cupy.logical\_not = <ufunc 'cupy\_logical\_not'> Computes the logical NOT of an array.

# See also:

numpy.logical\_not

```
cupy.logical_xor = <ufunc 'cupy_logical_xor'>
Computes the logical XOR of two arrays.
```

# See also:

numpy.logical\_xor

# **Comparison operations**

```
cupy.greater = <ufunc 'cupy_greater'>
```

Tests elementwise if x1 > x2.

# See also:

numpy.greater

```
cupy.greater_equal = <ufunc 'cupy_greater_equal'>
Tests elementwise if x1 >= x2.
```

See also:

numpy.greater\_equal

cupy.less = <ufunc 'cupy\_less'>

Tests elementwise if x1 < x2.

# See also:

# numpy.less

```
cupy.less_equal = <ufunc 'cupy_less_equal'>
Tests elementwise if x1 <= x2.
```

See also:

numpy.less\_equal

```
cupy.equal = <ufunc 'cupy_equal'>
```

Tests elementwise if x1 = x2.

See also:

numpy.equal

#### cupy.not\_equal = <ufunc 'cupy\_not\_equal'> Tests elementwise if x1 != x2.

See also:

numpy.equal

# **Mathematical Functions**

# **Trigonometric functions**

```
cupy.sin = <ufunc 'cupy_sin'>
Elementwise sine function.
```

#### See also:

numpy.sin

cupy.cos = <ufunc 'cupy\_cos'> Elementwise cosine function.

#### See also:

# numpy.cos

cupy.tan = <ufunc 'cupy\_tan'> Elementwise tangent function.

#### See also:

# numpy.tan

```
cupy.arcsin = <ufue (cupy_arcsin'>
Elementwise inverse-sine function (a.k.a. arcsine function).
```

#### See also:

## numpy.arcsin

# cupy.arccos = <ufunc 'cupy\_arccos'>

Elementwise inverse-cosine function (a.k.a. arccosine function).

#### See also:

# numpy.arccos

#### cupy.arctan = <ufunc 'cupy\_arctan'>

Elementwise inverse-tangent function (a.k.a. arctangent function).

#### See also:

numpy.arctan

# cupy.hypot = <ufunc 'cupy\_hypot'>

Computes the hypoteneous of orthogonal vectors of given length.

This is equivalent to sqrt(x1 + x2 + x2 + x2), while this function is more efficient.

#### See also:

numpy.hypot

cupy.arctan2 = <ufunc 'cupy\_arctan2'>
Elementwise inverse-tangent of the ratio of two arrays.

#### See also:

numpy.arctan2

cupy.deg2rad = <ufunc 'cupy\_deg2rad'>
 Converts angles from degrees to radians elementwise.

See also:

numpy.deg2rad,numpy.radians

cupy.rad2deg = <ufunc 'cupy\_rad2deg'>
 Converts angles from radians to degrees elementwise.

See also:

numpy.rad2deg,numpy.degrees

cupy.degrees = <ufunc 'cupy\_rad2deg'>
 Converts angles from radians to degrees elementwise.

See also:

numpy.rad2deg,numpy.degrees

```
cupy.radians = <ufunc 'cupy_deg2rad'>
    Converts angles from degrees to radians elementwise.
```

# See also:

numpy.deg2rad,numpy.radians

# **Hyperbolic functions**

```
cupy.sinh = <ufunc 'cupy_sinh'>
Elementwise hyperbolic sine function.
```

# See also:

numpy.sinh

cupy.cosh = <ufunc 'cupy\_cosh'>
 Elementwise hyperbolic cosine function.

# See also:

numpy.cosh

cupy.tanh = <ufunc 'cupy\_tanh'>
Elementwise hyperbolic tangent function.

# See also:

numpy.tanh

cupy.arcsinh = <ufunc 'cupy\_arcsinh'>
Elementwise inverse of hyperbolic sine function.

# See also:

numpy.arcsinh

cupy.arccosh = <ufunc 'cupy\_arccosh'>
Elementwise inverse of hyperbolic cosine function.

See also:

numpy.arccosh

cupy.arctanh = <ufunc 'cupy\_arctanh'>
Elementwise inverse of hyperbolic tangent function.

See also:

numpy.arctanh

# Rounding

cupy.rint = <ufunc 'cupy\_rint'>

Rounds each element of an array to the nearest integer.

# See also:

numpy.rint

cupy.floor = <ufunc 'cupy\_floor'>

Rounds each element of an array to its floor integer.

See also:

numpy.floor

cupy.ceil = <ufunc 'cupy\_ceil'>

Rounds each element of an array to its ceiling integer.

# See also:

numpy.ceil

cupy.trunc = <ufunc 'cupy\_trunc'>

Rounds each element of an array towards zero.

# See also:

numpy.trunc

# Sums and products

cupy.sum(\*args, \*\*kwargs)

cupy.prod(\*args, \*\*kwargs)

# **Exponential and logarithm functions**

# cupy.exp = <ufunc 'cupy\_exp'>

Elementwise exponential function.

See also:

numpy.exp

cupy.expm1 = <ufunc 'cupy\_expm1'> Computes exp(x) - 1 elementwise.

#### See also:

numpy.expm1

cupy.exp2 = <ufunc 'cupy\_exp2'> Elementwise exponentiation with base 2.

See also:

numpy.exp2

cupy.log = <ufunc 'cupy\_log'>
 Elementwise natural logarithm function.

See also:

numpy.log

cupy.log10 = <ufunc 'cupy\_log10'> Elementwise common logarithm function.

See also:

numpy.log10

```
cupy.log2 = <ufunc 'cupy_log2'>
Elementwise binary logarithm function.
```

#### See also:

numpy.log2

```
cupy.log1p = <ufunc 'cupy_log1p'>
Computes log(1 + x) elementwise.
```

See also:

numpy.log1p

```
cupy.logaddexp = <ufunc 'cupy_logaddexp'>
```

Computes log(exp(x1) + exp(x2)) elementwise.

See also:

numpy.logaddexp

```
cupy.logaddexp2 = <ufunc 'cupy_logaddexp2'>
```

Computes log2 (exp2(x1) + exp2(x2)) elementwise.

See also:

numpy.logaddexp2

# **Floating point manipulations**

# cupy.signbit = <ufunc 'cupy\_signbit'>

Tests elementwise if the sign bit is set (i.e. less than zero).

# See also:

numpy.signbit

cupy.copysign = <ufunc 'cupy\_copysign'>

Returns the first argument with the sign bit of the second elementwise.

## See also:

numpy.copysign

cupy.ldexp = <ufunc 'cupy\_ldexp'>

Computes x1 \* 2 \*\* x2 elementwise.

See also:

numpy.ldexp

## cupy.frexp = <ufunc 'cupy\_frexp'>

Decomposes each element to mantissa and two's exponent.

This ufunc outputs two arrays of the input dtype and the int dtype.

See also:

numpy.frexp

## cupy.nextafter = <ufunc 'cupy\_nextafter'>

Computes the nearest neighbor float values towards the second argument.

See also:

numpy.nextafter

## **Arithmetic operations**

# cupy.negative = <ufunc 'cupy\_negative'>

Takes numerical negative elementwise.

## See also:

## numpy.negative

cupy.add = <ufunc 'cupy\_add'> Adds two arrays elementwise.

See also:

#### numpy.add

cupy.subtract = <ufunc 'cupy\_subtract'>
 Subtracts arguments elementwise.

#### See also:

## numpy.subtract

cupy.multiply = <ufunc 'cupy\_multiply'> Multiplies two arrays elementwise.

#### See also:

numpy.multiply

## cupy.divide = <ufunc 'cupy\_divide'> Divides arguments elementwise.

See also:

numpy.divide

## cupy.true\_divide = <ufunc 'cupy\_true\_divide'> Elementwise true division (i.e. division as floating values).

See also:

numpy.true\_divide

cupy.floor\_divide = <ufunc 'cupy\_floor\_divide'> Elementwise floor division (i.e. integer quotient).

See also:

numpy.floor\_divide

cupy.power = <ufunc 'cupy\_power'>
 Computes x1 \*\* x2 elementwise.

See also:

numpy.power

```
cupy . fmod = <ufunc 'cupy_fmod'>
    Computes the remainder of C division elementwise.
```

See also:

numpy.fmod

```
cupy.mod = <ufunc 'cupy_remainder'>
Computes the remainder of Python division elementwise.
```

See also:

numpy.remainder

```
cupy.remainder = <ufunc 'cupy_remainder'>
Computes the remainder of Python division elementwise.
```

See also:

numpy.remainder

```
cupy.modf = <ufunc 'cupy_modf'>
```

Extracts the fractional and integral parts of an array elementwise.

This ufunc returns two arrays.

See also:

numpy.modf

```
cupy.reciprocal = <ufunc 'cupy_reciprocal'>
Computes 1 / x elementwise.
```

See also:

numpy.reciprocal

## **Miscellaneous**

cupy.clip(\*args, \*\*kwargs)

cupy.sqrt = <ufunc 'cupy\_sqrt'>

## cupy . square = <ufunc 'cupy\_square'> Elementwise square function.

## See also:

## numpy.square

```
cupy.absolute = <ufunc 'cupy_absolute'>
Elementwise absolute value function.
```

#### See also:

#### numpy.absolute

```
cupy.sign = <ufunc 'cupy_sign'>
Elementwise sign function.
```

It returns -1, 0, or 1 depending on the sign of the input.

## See also:

numpy.sign

## cupy.maximum = <ufunc 'cupy\_maximum'>

Takes the maximum of two arrays elementwise.

If NaN appears, it returns the NaN.

## See also:

numpy.maximum

## cupy.minimum = <ufunc 'cupy\_minimum'>

Takes the minimum of two arrays elementwise.

If NaN appears, it returns the NaN.

## See also:

numpy.minimum

## cupy.fmax = <ufunc 'cupy\_fmax'>

Takes the maximum of two arrays elementwise.

If NaN appears, it returns the other operand.

## See also:

numpy.fmax

## cupy.fmin = <ufunc 'cupy\_fmin'>

Takes the minimum of two arrays elementwise.

If NaN appears, it returns the other operand.

See also:

numpy.fmin

# Random Sampling (cupy.random)

CuPy's random number generation routines are based on cuRAND. They cover a small fraction of numpy.random.

The big difference of *cupy.random* from numpy.random is that *cupy.random* supports dtype option for most functions. This option enables us to generate float32 values directly without any space overhead.

## Sample random data

cupy.random.choice (a, size=None, replace=True, p=None)

Returns an array of random values from a given 1-D array.

Each element of the returned array is independently sampled from a according to p or uniformly.

## Parameters

- **a** (1-D array-like or int) If an array-like, a random sample is generated from its elements. If an int, the random sample is generated as if a was cupy.arange (n)
- **size** (*int* or tuple of *ints*) The shape of the array.
- replace (boolean) Whether the sample is with or without replacement
- **p** (1-D array-like) The probabilities associated with each entry in a. If not given the sample assumes a uniform distribution over all entries in a.

## Returns

An array of a values distributed according to p or uniformly.

Return type cupy.ndarray

## See also:

numpy.random.choice()

cupy.random.rand(\*size, \*\*kwarg)

Returns an array of uniform random values over the interval [0, 1).

Each element of the array is uniformly distributed on the half-open interval [0, 1). All elements are identically and independently distributed (i.i.d.).

## Parameters

- **size** (*tuple of ints*) The shape of the array.
- **dtype** Data type specifier. Only numpy.float32 and numpy.float64 types are allowed. The default is numpy.float64.

Returns A random array.

Return type cupy.ndarray

## See also:

numpy.random.rand()

```
cupy.random.randn(*size, **kwarg)
```

Returns an array of standard normal random values.

Each element of the array is normally distributed with zero mean and unit variance. All elements are identically and independently distributed (i.i.d.).

## Parameters

- **size** (tuple of ints) The shape of the array.
- **dtype** Data type specifier. Only numpy.float32 and numpy.float64 types are allowed. The default is numpy.float64.

Returns An array of standard normal random values.

Return type *cupy.ndarray* 

## See also:

numpy.random.randn()

cupy.random.randint(low, high=None, size=None)

Returns a scalar or an array of integer values over [low, high).

Each element of returned values are independently sampled from uniform distribution over left-close and rightopen interval [low, high).

## Parameters

- low (*int*) If high is not None, it is the lower bound of the interval. Otherwise, it is the **upper** bound of the interval and lower bound of the interval is set to 0.
- high (*int*) Upper bound of the interval.
- **size** (None or int or tuple of ints) The shape of returned value.

**Returns** If size is None, it is single integer sampled. If size is integer, it is the 1D-array of length size element. Otherwise, it is the array whose shape specified by size.

Return type int or cupy.ndarray of ints

```
cupy.random.random_integers (low, high=None, size=None)
```

Return a scalar or an array of integer values over [low, high]

Each element of returned values are independently sampled from uniform distribution over closed interval [low, high].

#### Parameters

- low (*int*) If high is not None, it is the lower bound of the interval. Otherwise, it is the **upper** bound of the interval and the lower bound is set to 1.
- high (*int*) Upper bound of the interval.
- size (None or int or tuple of ints) The shape of returned value.

**Returns** If size is None, it is single integer sampled. If size is integer, it is the 1D-array of length size element. Otherwise, it is the array whose shape specified by size.

**Return type** int or cupy.ndarray of ints

cupy.random.random\_sample(size=None, dtype=<type 'float'>)

Returns an array of random values over the interval [0, 1).

This is a variant of cupy.random.rand().

## Parameters

- **size** (*int* or tuple of *ints*) The shape of the array.
- **dtype** Data type specifier. Only numpy.float32 and numpy.float64 types are allowed.

Returns An array of uniformly distributed random values.

Return type *cupy.ndarray* 

#### See also:

numpy.random.random\_sample()

cupy.random.random(size=None, dtype=<type 'float'>)

Returns an array of random values over the interval [0, 1).

This is a variant of cupy.random.rand().

#### Parameters

- **size** (*int* or *tuple* of *ints*) The shape of the array.
- **dtype** Data type specifier. Only numpy.float32 and numpy.float64 types are allowed.

Returns An array of uniformly distributed random values.

Return type *cupy.ndarray* 

## See also:

numpy.random.random\_sample()

cupy.random.ranf (size=None, dtype=<type 'float'>)
Returns an array of random values over the interval [0, 1).

This is a variant of cupy.random.rand().

#### **Parameters**

- **size** (*int* or tuple of *ints*) The shape of the array.
- **dtype** Data type specifier. Only numpy.float32 and numpy.float64 types are allowed.

Returns An array of uniformly distributed random values.

Return type *cupy.ndarray* 

## See also:

```
numpy.random.random_sample()
```

```
cupy.random.sample(size=None, dtype=<type 'float'>)
    Returns an array of random values over the interval [0, 1).
```

This is a variant of cupy.random.rand().

## Parameters

- size (int or tuple of ints) The shape of the array.
- **dtype** Data type specifier. Only numpy.float32 and numpy.float64 types are allowed.

Returns An array of uniformly distributed random values.

Return type cupy.ndarray

## See also:

```
numpy.random.random_sample()
```

## **Distributions**

```
cupy.random.gumbel (loc=0.0, scale=1.0, size=None, dtype=<type 'float'>)
Returns an array of samples drawn from a Gumbel distribution.
```

The samples are drawn from a Gumbel distribution with location loc and scale scale. Its probability density function is defined as

$$f(x) = \frac{1}{\eta} \exp\left\{-\frac{x-\mu}{\eta}\right\} \exp\left[-\exp\left\{-\frac{x-\mu}{\eta}\right\}\right]$$

where  $\mu$  is loc and  $\eta$  is scale.

## **Parameters**

- loc (float) The location of the mode  $\mu$ .
- **scale** (*float*) The scale parameter  $\eta$ .
- **size** (*int or tuple of ints*) The shape of the array. If None, a zerodimensional array is generated.
- **dtype** Data type specifier. Only numpy.float32 and numpy.float64 types are allowed.

Returns Samples drawn from the Gumbel destribution.

#### **Return type** *cupy.ndarray*

#### See also:

```
numpy.random.gumbel()
```

cupy.random.lognormal (*mean=0.0*, *sigma=1.0*, *size=None*, *dtype=<type 'float'>*) Returns an array of samples drawn from a log normal distribution.

The samples are natural log of samples drawn from a normal distribution with mean mean and deviation sigma.

## Parameters

- **mean** (*float*) Mean of the normal distribution.
- **sigma** (*float*) Standard deviation of the normal distribution.
- **size** (*int or tuple of ints*) The shape of the array. If None, a zerodimensional array is generated.
- **dtype** Data type specifier. Only numpy.float32 and numpy.float64 types are allowed.

Returns Samples drawn from the log normal distribution.

#### Return type *cupy.ndarray*

## See also:

```
numpy.random.lognormal()
```

cupy.random.**normal** (*loc=0.0*, *scale=1.0*, *size=None*, *dtype=<type* '*float*'>) Returns an array of normally distributed samples.

#### **Parameters**

- **loc** (float or array\_like of floats) Mean of the normal distribution.
- **scale** (*float* or *array\_like* of *floats*) Standard deviation of the normal distribution.
- **size** (*int or tuple of ints*) The shape of the array. If None, a zerodimensional array is generated.
- **dtype** Data type specifier. Only numpy.float32 and numpy.float64 types are allowed.

**Returns** Normally distributed samples.

## Return type cupy.ndarray

## See also:

```
numpy.random.normal()
```

cupy.random.standard\_normal(size=None, dtype=<type 'float'>)

Returns an array of samples drawn from the standard normal distribution.

This is a variant of cupy.random.randn().

## Parameters

- **size** (*int or tuple of ints*) The shape of the array. If None, a zerodimensional array is generated.
- **dtype** Data type specifier.

Returns Samples drawn from the standard normal distribution.

## Return type *cupy.ndarray*

## See also:

```
numpy.random.standard_normal()
```

cupy.random.uniform (*low=0.0*, *high=1.0*, *size=None*, *dtype=<type 'float'>*) Returns an array of uniformly-distributed samples over an interval.

Samples are drawn from a uniform distribution over the half-open interval [low, high).

## Parameters

- **low** (*float*) Lower end of the interval.
- high (float) Upper end of the interval.
- **size** (*int or tuple of ints*) The shape of the array. If None, a zerodimensional array is generated.
- **dtype** Data type specifier.

Returns Samples drawn from the uniform distribution.

Return type *cupy.ndarray* 

## See also:

numpy.random.uniform()

## Random number generator

## cupy.random.seed(seed=None)

Resets the state of the random number generator with a seed.

This function resets the state of the global random number generator for the current device. Be careful that generators for other devices are not affected.

**Parameters seed** (*None or int*) - Seed for the random number generator. If None, it uses os.urandom() if available or time.clock() otherwise. Note that this function does not support seeding by an integer array.

## cupy.random.get\_random\_state()

Gets the state of the random number generator for the current device.

If the state for the current device is not created yet, this function creates a new one, initializes it, and stores it as the state for the current device.

Returns The state of the random number generator for the device.

Return type RandomState

class cupy.random.RandomState (seed=None, method=100)

Portable container of a pseudo-random number generator.

An instance of this class holds the state of a random number generator. The state is available only on the device which has been current at the initialization of the instance.

Functions of *cupy.random* use global instances of this class. Different instances are used for different devices. The global state for the current device can be obtained by the *cupy.random.get\_random\_state()* function.

## Parameters

- **seed** (*None or int*) Seed of the random number generator. See the *seed()* method for detail.
- **method** (*int*) Method of the random number generator. Following values are available:

```
cupy.cuda.curand.CURAND_RNG_PSEUDO_DEFAULT
cupy.cuda.curand.CURAND_RNG_XORWOW
cupy.cuda.curand.CURAND_RNG_MRG32K3A
cupy.cuda.curand.CURAND_RNG_MTGP32
cupy.cuda.curand.CURAND_RNG_MT19937
cupy.cuda.curand.CURAND_RNG_PHILOX4_32_10
```

## **choice** (*a*, *size=None*, *replace=True*, *p=None*)

Returns an array of random values from a given 1-D array.

#### See also:

cupy.random.choice() for full document, numpy.random.choice()

#### interval (mx, size)

Generate multiple integers independently sampled uniformly from [0, mx].

#### **Parameters**

- **mx** (*int*) Upper bound of the interval
- **size** (*None or int or tuple*) Shape of the array or the scalar returned.
- **Returns** If None, an *cupy.ndarray* with shape () is returned. If int, 1-D array of length size is returned. If tuple, multi-dimensional array with shape size is returned. Currently, each element of the array is numpy.int32.

## Return type int or cupy.ndarray

**lognormal** (*mean=0.0*, *sigma=1.0*, *size=None*, *dtype=<type* '*float*'>) Returns an array of samples drawn from a log normal distribution.

### See also:

cupy.random.lognormal() for full documentation, numpy.random.RandomState. lognormal()

**normal** (*loc=0.0*, *scale=1.0*, *size=None*, *dtype=<type 'float'>*) Returns an array of normally distributed samples.

#### See also:

cupy.random.normal() for full documentation, numpy.random.RandomState.normal()

## rand(\*size, \*\*kwarg)

Returns uniform random values over the interval [0, 1).

See also:

cupy.random.rand() for full documentation, numpy.random.RandomState.rand()

**randn** (\**size*, \*\**kwarg*)

Returns an array of standard normal random values.

## See also:

cupy.random.randn() for full documentation, numpy.random.RandomState.randn()

#### random\_sample (size=None, dtype=<type 'float'>)

Returns an array of random values over the interval [0, 1).

See also:

cupy.random.random\_sample() for full documentation, numpy.random.RandomState. random\_sample()

#### seed(seed=None)

Resets the state of the random number generator with a seed.

#### See also:

cupy.random.seed() for full documentation, numpy.random.RandomState.seed()

## standard\_normal (size=None, dtype=<type 'float'>)

Returns samples drawn from the standard normal distribution.

### See also:

cupy.random.standard\_normal() for full documentation, numpy.random.RandomState. standard\_normal()

**uniform** (*low=0.0*, *high=1.0*, *size=None*, *dtype=<type 'float'>*) Returns an array of uniformly-distributed samples over an interval.

#### See also:

```
cupy.random.uniform() for full documentation, numpy.random.RandomState.
uniform()
```

## Sorting, Searching, and Counting

cupy.**argmax** (*a*, *axis=None*, *dtype=None*, *out=None*, *keepdims=False*) Returns the indices of the maximum along an axis.

#### **Parameters**

- a (cupy.ndarray) Array to take argmax.
- **axis** (*int*) Along which axis to find the maximum. a is flattened by default.
- **dtype** Data type specifier.
- **out** (cupy.ndarray) **Output** array.
- keepdims (bool) If True, the axis axis is preserved as an axis of length one.

Returns The indices of the maximum of a along an axis.

## Return type cupy.ndarray

#### See also:

```
numpy.argmax()
```

```
cupy.argmin(a, axis=None, dtype=None, out=None, keepdims=False)
```

Returns the indices of the minimum along an axis.

## Parameters

- a (cupy.ndarray) Array to take argmin.
- **axis** (*int*) Along which axis to find the minimum. a is flattened by default.
- **dtype** Data type specifier.
- **out** (cupy.ndarray) **Output** array.
- keepdims (bool) If True, the axis axis is preserved as an axis of length one.

Returns The indices of the minimum of a along an axis.

Return type *cupy.ndarray* 

## See also:

## numpy.argmin()

## cupy.count\_nonzero(*a*, *axis=None*)

Counts the number of non-zero values in the array.

## Parameters

- a (cupy.ndarray) The array for which to count non-zeros.
- **axis** (*int or tuple*, *optional*) Axis or tuple of axes along which to count nonzeros. Default is None, meaning that non-zeros will be counted along a flattened version of a

## Returns

**Number of non-zero values in the array** along a given axis. Otherwise, the total number of non-zero values in the array is returned.

Return type int or cupy.ndarray of int

## cupy.nonzero(a)

Return the indices of the elements that are non-zero.

Returns a tuple of arrays, one for each dimension of a, containing the indices of the non-zero elements in that dimension.

## Parameters a (cupy.ndarray) - array

Returns Indices of elements that are non-zero.

## Return type tuple of arrays

## See also:

numpy.nonzero()

## cupy.flatnonzero(a)

Return indices that are non-zero in the flattened version of a.

This is equivalent to a.ravel().nonzero()[0].

Parameters a (cupy.ndarray) – input array

Returns Output array, containing the indices of the elements of a.ravel() that are non-zero.

Return type cupy.ndarray

### See also:

```
numpy.flatnonzero()
```

```
cupy.where(*args, **kwargs)
```

## **Statistics**

## **Order statistics**

cupy.amin(\*args, \*\*kwargs)

cupy.amax(\*args, \*\*kwargs)

cupy.nanmin(a, axis=None, out=None, keepdims=False)

Returns the minimum of an array along an axis ignoring NaN.

When there is a slice whose elements are all NaN, a RuntimeWarning is raised and NaN is returned.

## Parameters

- **a** (cupy.ndarray) Array to take the minimum.
- **axis** (*int*) Along which axis to take the minimum. The flattened array is used by default.
- **out** (cupy.ndarray) **Output** array.
- **keepdims** (bool) If True, the axis is remained as an axis of size one.

Returns The minimum of a, along the axis if specified.

Return type *cupy.ndarray* 

## See also:

## numpy.nanmin()

cupy.nanmax(a, axis=None, out=None, keepdims=False)

Returns the maximum of an array along an axis ignoring NaN.

When there is a slice whose elements are all NaN, a RuntimeWarning is raised and NaN is returned.

## Parameters

- a (cupy.ndarray) Array to take the maximum.
- **axis** (*int*) Along which axis to take the maximum. The flattened array is used by default.
- **out** (cupy.ndarray) **Output** array.
- **keepdims** (*bool*) If True, the axis is remained as an axis of size one.

Returns The maximum of a, along the axis if specified.

Return type *cupy.ndarray* 

## See also:

numpy.nanmax()

## Means and variances

cupy.mean (*a*, *axis=None*, *dtype=None*, *out=None*, *keepdims=False*) Returns the arithmetic mean along an axis.

## Parameters

- a (cupy.ndarray) Array to compute mean.
- **axis** (*int*) Along which axis to compute mean. The flattened array is used by default.
- **dtype** Data type specifier.
- **out** (cupy.ndarray) **Output** array.
- **keepdims** (*bool*) If True, the axis is remained as an axis of size one.

Returns The mean of the input array along the axis.

#### Return type cupy.ndarray

#### See also:

## numpy.mean()

cupy.var(*a*, *axis=None*, *dtype=None*, *out=None*, *ddof=0*, *keepdims=False*) Returns the variance along an axis.

### **Parameters**

- **a** (cupy.ndarray) Array to compute variance.
- **axis** (*int*) Along which axis to compute variance. The flattened array is used by default.
- dtype Data type specifier.
- **out** (cupy.ndarray) **Output** array.
- **keepdims** (bool) If True, the axis is remained as an axis of size one.

**Returns** The variance of the input array along the axis.

### Return type cupy.ndarray

## See also:

#### numpy.var()

cupy.std(a, axis=None, dtype=None, out=None, ddof=0, keepdims=False)

Returns the standard deviation along an axis.

#### Parameters

- **a** (cupy.ndarray) Array to compute standard deviation.
- **axis** (*int*) Along which axis to compute standard deviation. The flattened array is used by default.
- dtype Data type specifier.
- **out** (cupy.ndarray) **Output** array.
- **keepdims** (*bool*) If True, the axis is remained as an axis of size one.

Returns The standard deviation of the input array along the axis.

### Return type cupy.ndarray

#### See also:

numpy.std()

## **Histograms**

cupy.bincount (x, weights=None, minlength=None)

Count number of occurrences of each value in array of non-negative ints.

## Parameters

- **x** (cupy.ndarray) Input array.
- weights (cupy.ndarray) Weights array which has the same shape as x.
- minlength (*int*) A minimum number of bins for the output array.

#### Returns

The result of binning the input array. The length of output is equal to max(cupy. max(x) + 1, minlength).

Return type *cupy.ndarray* 

## See also:

```
numpy.bincount()
```

## **External Functions**

## cupy.scatter\_add(a, slices, value)

Adds given values to specified elements of an array.

It adds value to the specified elements of a. If all of the indices target different locations, the operation of scatter\_add() is equivalent to a[slices] = a[slices] + value. If there are multiple elements targeting the same location, scatter\_add() uses all of these values for addition. On the other hand, a[slices] = a[slices] + value only adds the contribution from one of the indices targeting the same location.

Note that just like an array indexing, negative indices are interpreted as counting from the end of an array.

Also note that scatter\_add() behaves identically to numpy.add.at().

## Example

```
>>> import numpy
>>> import cupy
>>> a = cupy.zeros((6,), dtype=numpy.float32)
>>> i = cupy.array([1, 0, 1])
>>> v = cupy.array([1, 1., 1.])
>>> cupy.scatter_add(a, i, v);
>>> a
array([ 1., 2., 0., 0., 0., 0.], dtype=float32)
```

#### **Parameters**

• a (ndarray) - An array that gets added.

- **slices** It is integer, slices, ellipsis, numpy.newaxis, integer array-like, boolean array-like or tuple of them. It works for slices used for *cupy.ndarray.\_\_getitem\_\_()* and *cupy.ndarray.\_\_setitem\_\_()*.
- **v** (*array-like*) Values to increment a at referenced locations.

Note: It only supports types that are supported by CUDA's atomicAdd when an integer array is included in slices. The supported types are numpy.float32, numpy.int32, numpy.uint32, numpy.uint64 and numpy.ulonglong.

Note: scatter\_add() does not raise an error when indices exceed size of axes. Instead, it wraps indices.

#### See also:

numpy.add.at().

# NumPy-CuPy Generic Code Support

#### cupy.get\_array\_module(\*args)

Returns the array module for arguments.

This function is used to implement CPU/GPU generic code. If at least one of the arguments is a *cupy*. *ndarray* object, the *cupy* module is returned.

Parameters args – Values to determine whether NumPy or CuPy should be used.

**Returns** cupy or numpy is returned based on the types of the arguments.

Return type module

#### Example

A NumPy/CuPy generic function can be written as follows

```
>>> def softplus(x):
... xp = cupy.get_array_module(x)
... return xp.maximum(0, x) + xp.log1p(xp.exp(-abs(x)))
```

# Low-Level CUDA Support

## **Device management**

```
class cupy.cuda.Device
```

Object that represents a CUDA device.

This class provides some basic manipulations on CUDA devices.

It supports the context protocol. For example, the following code is an example of temporarily switching the current device:

with Device(0): do\_something\_on\_device\_0()

After the with statement gets done, the current device is reset to the original one.

**Parameters device** (*int or* cupy.cuda.Device) – Index of the device to manipulate. Be careful that the device ID (a.k.a. GPU ID) is zero origin. If it is a Device object, then its ID is used. The current device is selected by default.

**Variables** id(int) - ID of this device.

## compute\_capability

Compute capability of this device.

The capability is represented by a string containing the major index and the minor index. For example, compute capability 3.5 is represented by the string '35'.

## cublas\_handle

The cuBLAS handle for this device.

The same handle is used for the same device even if the Device instance itself is different.

## cusolver\_handle

The cuSOLVER handle for this device.

The same handle is used for the same device even if the Device instance itself is different.

## synchronize()

Synchronizes the current thread to the device.

### use()

Makes this device current.

If you want to switch a device temporarily, use the *with* statement.

## Memory management

```
class cupy.cuda.Memory
```

Memory allocation on a CUDA device.

This class provides an RAII interface of the CUDA memory allocation.

#### **Parameters**

- device (cupy.cuda.Device) Device whose memory the pointer refers to.
- **size** (*int*) Size of the memory allocation in bytes.

## \_\_int\_

Returns the pointer value to the head of the allocation.

## \_\_long\_\_

## class cupy.cuda.MemoryPointer

Pointer to a point on a device memory.

An instance of this class holds a reference to the original memory buffer and a pointer to a place within this buffer.

#### **Parameters**

- mem (Memory) The device memory buffer.
- offset (*int*) An offset from the head of the buffer to the place this pointer refers.

### Variables

- device (cupy.cuda.Device) Device whose memory the pointer refers to.
- mem (Memory) The device memory buffer.
- **ptr** (*int*) Pointer to the place within the buffer.

## \_\_add\_\_

Adds an offset to the pointer.

#### \_\_iadd\_\_

Adds an offset to the pointer in place.

## \_\_int\_

Returns the pointer value.

## \_isub\_\_

Subtracts an offset from the pointer in place.

#### \_\_long\_\_

\_\_\_**radd\_\_** x.\_\_radd\_\_(y) <==> y+x

#### rsub

 $x\_rsub\_(y) \iff y-x$ 

## \_\_sub\_\_

Subtracts an offset from the pointer.

#### copy\_from()

Copies a memory sequence from a (possibly different) device or host.

This function is a useful interface that selects appropriate one from *copy\_from\_device()* and *copy\_from\_host()*.

## **Parameters**

• **mem** (*ctypes.c\_void\_p* or cupy.cuda.MemoryPointer) - Source memory pointer.

• **size** (*int*) – Size of the sequence in bytes.

## copy\_from\_async()

Copies a memory sequence from an arbitrary place asynchronously.

This function is a useful interface that selects appropriate one from *copy\_from\_device\_async()* and *copy\_from\_host\_async()*.

#### Parameters

- **mem** (*ctypes.c\_void\_p or* cupy.cuda.MemoryPointer) Source memory pointer.
- **size** (*int*) Size of the sequence in bytes.
- **stream** (cupy.cuda.Stream) CUDA stream.

#### copy\_from\_device()

Copies a memory sequence from a (possibly different) device.

#### **Parameters**

- **src** (cupy.cuda.MemoryPointer) Source memory pointer.
- **size** (*int*) Size of the sequence in bytes.

### copy\_from\_device\_async()

Copies a memory from a (possibly different) device asynchronously.

#### Parameters

- **src** (cupy.cuda.MemoryPointer) Source memory pointer.
- **size** (*int*) Size of the sequence in bytes.
- **stream** (cupy.cuda.Stream) CUDA stream.

## copy\_from\_host()

Copies a memory sequence from the host memory.

#### **Parameters**

- **mem** (*ctypes.c\_void\_p*) Source memory pointer.
- **size** (*int*) Size of the sequence in bytes.

#### copy\_from\_host\_async()

Copies a memory sequence from the host memory asynchronously.

#### **Parameters**

- **mem** (*ctypes.c\_void\_p*) Source memory pointer. It must be a pinned memory.
- **size** (*int*) Size of the sequence in bytes.
- **stream** (cupy.cuda.Stream) CUDA stream.

#### copy\_to\_host()

Copies a memory sequence to the host memory.

#### **Parameters**

- **mem** (*ctypes.c\_void\_p*) Target memory pointer.
- **size** (*int*) Size of the sequence in bytes.

## copy\_to\_host\_async()

Copies a memory sequence to the host memory asynchronously.

#### **Parameters**

- **mem** (*ctypes.c\_void\_p*) Target memory pointer. It must be a pinned memory.
- **size** (*int*) Size of the sequence in bytes.
- **stream** (cupy.cuda.Stream) CUDA stream.

#### memset()

Fills a memory sequence by constant byte value.

## **Parameters**

- value (int) Value to fill.
- **size** (*int*) Size of the sequence in bytes.

#### memset\_async()

Fills a memory sequence by constant byte value asynchronously.

### Parameters

- **value** (*int*) Value to fill.
- **size** (*int*) Size of the sequence in bytes.
- **stream** (cupy.cuda.Stream) CUDA stream.

#### cupy.cuda.alloc()

Calls the current allocator.

Use *set\_allocator()* to change the current allocator.

**Parameters** size (*int*) – Size of the memory allocation.

Returns Pointer to the allocated buffer.

#### **Return type** *MemoryPointer*

#### cupy.cuda.set\_allocator()

Sets the current allocator.

**Parameters allocator** (function) – CuPy memory allocator. It must have the same interface as the *cupy.cuda.alloc()* function, which takes the buffer size as an argument and returns the device buffer of that size.

#### class cupy.cuda.MemoryPool

Memory pool for all devices on the machine.

A memory pool preserves any allocations even if they are freed by the user. Freed memory buffers are held by the memory pool as *free blocks*, and they are reused for further memory allocations of the same sizes. The allocated blocks are managed for each device, so one instance of this class can be used for multiple devices.

**Note:** When the allocation is skipped by reusing the pre-allocated block, it does not call cudaMalloc and therefore CPU-GPU synchronization does not occur. It makes interleaves of memory allocations and kernel invocations very fast.

**Note:** The memory pool holds allocated blocks without freeing as much as possible. It makes the program hold most of the device memory, which may make other CUDA programs running in parallel out-of-memory situation.

**Parameters allocator** (function) – The base CuPy memory allocator. It is used for allocating new blocks when the blocks of the required size are all in use.

```
free_all_blocks()
Release free blocks.
```

**free\_all\_free**() Release free blocks.

## malloc()

Allocates the memory, from the pool if possible.

This method can be used as a CuPy memory allocator. The simplest way to use a memory pool as the default allocator is the following code:

```
set_allocator(MemoryPool().malloc)
```

**Parameters** size (*int*) – Size of the memory buffer to allocate in bytes.

Returns Pointer to the allocated buffer.

**Return type** *MemoryPointer* 

## n\_free\_blocks()

Count the total number of free blocks.

Returns The total number of free blocks.

**Return type** int

## **Streams and events**

class cupy.cuda.Stream(null=False, non\_blocking=False)
 CUDA stream.

This class handles the CUDA stream handle in RAII way, i.e., when an Stream instance is destroyed by the GC, its handle is also destroyed.

## Parameters

- **null** (*bool*) If True, the stream is a null stream (i.e. the default stream that synchronizes with all streams). Otherwise, a plain new stream is created.
- **non\_blocking** (*bool*) If True, the stream does not synchronize with the NULL stream.

Variables ptr (cupy.cuda.runtime.Stream) - Raw stream handle. It can be passed to the CUDA Runtime API via ctypes.

## add\_callback (callback, arg)

Adds a callback that is called when all queued work is done.

### Parameters

- **callback** (function) Callback function. It must take three arguments (Stream object, int error status, and user data object), and returns nothing.
- **arg** (*object*) Argument to the callback.

done

True if all work on this stream has been done.

record (event=None)

Records an event on the stream.

**Parameters event** (*None or* cupy.cuda.Event) – CUDA event. If None, then a new plain event is created and used.

Returns The recorded event.

Return type cupy.cuda.Event

See also:

cupy.cuda.Event.record()

#### synchronize()

Waits for the stream completing all queued work.

#### wait\_event (event)

Makes the stream wait for an event.

The future work on this stream will be done after the event.

**Parameters event** (cupy.cuda.Event) - CUDA event.

**class** cupy.cuda.**Event** (*block=False*, *disable\_timing=False*, *interprocess=False*) CUDA event, a synchronization point of CUDA streams.

This class handles the CUDA event handle in RAII way, i.e., when an Event instance is destroyed by the GC, its handle is also destroyed.

## Parameters

- **block** (*bool*) If True, the event blocks on the *synchronize()* method.
- **disable\_timing** (*bool*) If True, the event does not prepare the timing data.
- interprocess (bool) If True, the event can be passed to other processes.

Variables ptr (cupy.cuda.runtime.Stream) – Raw stream handle. It can be passed to the CUDA Runtime API via ctypes.

## done

True if the event is done.

record (stream=None)

Records the event to a stream.

**Parameters stream** (cupy.cuda.Stream) – CUDA stream to record event. The null stream is used by default.

See also:

cupy.cuda.Stream.record()

## synchronize()

Synchronizes all device work to the event.

If the event is created as a blocking event, it also blocks the CPU thread until the event is done.

cupy.cuda.get\_elapsed\_time(start\_event, end\_event)

Gets the elapsed time between two events.

## Parameters

- **start\_event** (Event) Earlier event.
- end\_event (Event) Later event.

Returns Elapsed time in milliseconds.

Return type float

## Profiler

cupy.cuda.profile(\*args, \*\*kwds)

Enable CUDA profiling during with statement.

This function enables profiling on entering a with statement, and disables profiling on leaving the statement.

```
>>> with cupy.cuda.profile():
... # do something you want to measure
... pass
```

```
cupy.cuda.profiler.initialize()
    Initialize the CUDA profiler.
```

This function initialize the CUDA profiler. See the CUDA document for detail.

#### Parameters

- **config\_file** (*str*) Name of the configuration file.
- **output\_file** (*str*) Name of the coutput file.
- **output\_mode** (*int*) cupy.cuda.profiler.cudaKeyValuePair or cupy. cuda.profiler.cudaCSV.

#### cupy.cuda.profiler.start()

Enable profiling.

A user can enable CUDA profiling. When an error occurs, it raises an exception.

See the CUDA document for detail.

## cupy.cuda.profiler.stop()

Disable profiling.

A user can disable CUDA profiling. When an error occurs, it raises an exception.

See the CUDA document for detail.

```
cupy.cuda.nvtx.Mark()
```

Marks an instantaneous event (marker) in the application.

Markes are used to describe events at a specific time during execution of the application.

**Parameters** 

- **message** (*str*) Name of a marker.
- id\_color (*int*) ID of color for a marker.

```
cupy.cuda.nvtx.MarkC()
```

Marks an instantaneous event (marker) in the application.

Markes are used to describe events at a specific time during execution of the application.

#### Parameters

- **message** (*str*) Name of a marker.
- **color** (*uint32*) Color code for a marker.

## cupy.cuda.nvtx.RangePush()

Starts a nestead range.

Ranges are used to describe events over a time span during execution of the application. The duration of a range is defined by the corresponding pair of RangePush\*() to RangePop() calls.

## **Parameters**

- **message** (*str*) Name of a range.
- id\_color (*int*) ID of color for a range.

```
cupy.cuda.nvtx.RangePushC()
```

Starts a nestead range.

Ranges are used to describe events over a time span during execution of the application. The duration of a range is defined by the corresponding pair of RangePush\*() to RangePop() calls.

#### **Parameters**

- **message** (*str*) Name of a range.
- color (uint 32) ARGB color for a range.

cupy.cuda.nvtx.RangePop()

Ends a nestead range.

Ranges are used to describe events over a time span during execution of the application. The duration of a range is defined by the corresponding pair of RangePush\*() to RangePop() calls.

# Kernel binary memoization

#### cupy.memoize()

Makes a function memoizing the result for each argument and device.

This decorator provides automatic memoization of the function result.

```
Parameters for_each_device (bool) – If True, it memoizes the results for each device. Otherwise, it memoizes the results only based on the arguments.
```

#### cupy.clear\_memo()

Clears the memoized results for all functions decorated by memoize.

# **User-Defined Kernels**

CuPy provides easy ways to define two types of CUDA kernels: elementwise kernels and reduction kernels. We first describe how to define and call elementwise kernels, and then describe how to define and call reduction kernels.

## **Basics of elementwise kernels**

An elementwise kernel can be defined by the *ElementwiseKernel* class. The instance of this class defines a CUDA kernel which can be invoked by the \_\_call\_\_ method of this instance.

A definition of an elementwise kernel consists of four parts: an input argument list, an output argument list, a loop body code, and the kernel name. For example, a kernel that computes a squared difference  $f(x, y) = (x - y)^2$  is defined as follows:

The argument lists consist of comma-separated argument definitions. Each argument definition consists of a *type specifier* and an *argument name*. Names of NumPy data types can be used as type specifiers.

Note: n, i, and names starting with an underscore \_ are reserved for the internal use.

The above kernel can be called on either scalars or arrays with broadcasting:

```
>>> x = cupy.arange(10, dtype=np.float32).reshape(2, 5)
>>> y = cupy.arange(5, dtype=np.float32)
>>> squared_diff(x, y)
array([[ 0.,  0.,  0.,  0.,  0.],
      [ 25.,  25.,  25.,  25.]], dtype=float32)
>>> squared_diff(x, 5)
array([[ 25.,  16.,  9.,  4.,  1.],
      [ 0.,  1.,  4.,  9.,  16.]], dtype=float32)
```

Output arguments can be explicitly specified (next to the input arguments):

```
>>> z = cupy.empty((2, 5), dtype=np.float32)
>>> squared_diff(x, y, z)
array([[ 0., 0., 0., 0., 0.],
      [ 25., 25., 25., 25., 25.]], dtype=float32)
```

## **Type-generic kernels**

If a type specifier is one character, then it is treated as a **type placeholder**. It can be used to define a type-generic kernels. For example, the above squared\_diff kernel can be made type-generic as follows:

```
>>> squared_diff_generic = cupy.ElementwiseKernel(
... 'T x, T y',
... 'T z',
... 'z = (x - y) * (x - y)',
... 'squared_diff_generic')
```

Type placeholders of a same character in the kernel definition indicate the same type. The actual type of these placeholders is determined by the actual argument type. The ElementwiseKernel class first checks the output arguments and then the input arguments to determine the actual type. If no output arguments are given on the kernel invocation, then only the input arguments are used to determine the type.

The type placeholder can be used in the loop body code:

```
>>> squared_diff_generic = cupy.ElementwiseKernel(
... 'T x, T y',
... 'T z',
... 'T diff = x - y;
... z = diff * diff;
... ''',
... 'squared_diff_generic')
```

More than one type placeholder can be used in a kernel definition. For example, the above kernel can be further made generic over multiple arguments:

```
>>> squared_diff_super_generic = cupy.ElementwiseKernel(
... 'X x, Y y',
... 'Z z',
... 'Z = (x - y) * (x - y)',
... 'squared_diff_super_generic')
```

Note that this kernel requires the output argument explicitly specified, because the type Z cannot be automatically determined from the input arguments.

## **Raw argument specifiers**

The ElementwiseKernel class does the indexing with broadcasting automatically, which is useful to define most elementwise computations. On the other hand, we sometimes want to write a kernel with manual indexing for some arguments. We can tell the ElementwiseKernel class to use manual indexing by adding the raw keyword preceding the type specifier.

We can use the special variable i and method \_ind.size() for the manual indexing. i indicates the index within the loop. \_ind.size() indicates total number of elements to apply the elementwise operation. Note that it represents the size **after** broadcast operation.

For example, a kernel that adds two vectors with reversing one of them can be written as follows:

```
>>> add_reverse = cupy.ElementwiseKernel(
... 'T x, raw T y', 'T z',
... 'z = x + y[_ind.size() - i - 1]',
... 'add_reverse')
```

(Note that this is an artificial example and you can write such operation just by z = x + y[::-1] without defining a new kernel). A raw argument can be used like an array. The indexing operator  $y[\_ind.size() - i - 1]$  involves an indexing computation on y, so y can be arbitrarily shaped and strode.

Note that raw arguments are not involved in the broadcasting. If you want to mark all arguments as raw, you must specify the size argument on invocation, which defines the value of \_ind.size().

## **Reduction kernels**

Reduction kernels can be defined by the *ReductionKernel* class. We can use it by defining four parts of the kernel code:

- 1. Identity value: This value is used for the initial value of reduction.
- 2. Mapping expression: It is used for the pre-processing of each element to be reduced.
- 3. Reduction expression: It is an operator to reduce the multiple mapped values. The special variables a and b are used for its operands.
- 4. Post mapping expression: It is used to transform the resulting reduced values. The special variable a is used as its input. Output should be written to the output parameter.

ReductionKernel class automatically inserts other code fragments that are required for an efficient and flexible reduction implementation.

For example, L2 norm along specified axes can be written as follows:

```
>>> 12norm_kernel = cupy.ReductionKernel(
        'T x', # input params
. . .
        'T y', # output params
. . .
        'x * x', # map
. . .
        'a + b', # reduce
. . .
        'y = sqrt(a)', # post-reduction map
. . .
        '0', # identity value
. . .
        'l2norm' # kernel name
. . .
...)
>>> x = cupy.arange(10, dtype='f').reshape(2, 5)
>>> l2norm_kernel(x, axis=1)
array([ 5.47722578, 15.96871948], dtype=float32)
```

Note: raw specifier is restricted for usages that the axes to be reduced are put at the head of the shape. It means, if you want to use raw specifier for at least one argument, the axis argument must be 0 or a contiguous increasing sequence of integers starting from 0, like (0, 1), (0, 1, 2), etc.

## Reference

## class cupy.ElementwiseKernel

User-defined elementwise kernel.

This class can be used to define an elementwise kernel with or without broadcasting.

The kernel is compiled at an invocation of the <u>\_\_\_\_\_\_</u>() method, which is cached for each device. The compiled binary is also cached into a file under the \$HOME/.cupy/kernel\_cache/directory with a hashed file name. The cached binary is reused by other processes.

## **Parameters**

- in\_params (*str*) Input argument list.
- out\_params (*str*) Output argument list.
- operation (*str*) The body in the loop written in CUDA-C/C++.
- **name** (*str*) Name of the kernel function. It should be set for readability of the performance profiling.
- **reduce\_dims** (bool) If False, the shapes of array arguments are kept within the kernel invocation. The shapes are reduced (i.e., the arrays are reshaped without copy to the minimum dimension) by default. It may make the kernel fast by reducing the index calculations.
- **options** (*list*) Options passed to the nvcc command.
- **preamble** (*str*) Fragment of the CUDA-C/C++ code that is inserted at the top of the cu file.
- **loop\_prep** (*str*) Fragment of the CUDA-C/C++ code that is inserted at the top of the kernel function definition and above the for loop.
- **after\_loop** (*str*) Fragment of the CUDA-C/C++ code that is inserted at the bottom of the kernel function definition.

\_\_call\_

Compiles and invokes the elementwise kernel.

The compilation runs only if the kernel is not cached. Note that the kernels with different argument dtypes or dimensions are not compatible. It means that single ElementwiseKernel object may be compiled into multiple kernel binaries.

## Parameters

- **args** Arguments of the kernel.
- **size** (*int*) Range size of the indices. If specified, the variable n is set to this value. Otherwise, the result of broadcasting is used to determine the value of n.

**Returns** Arrays are returned according to the out\_params argument of the \_\_init\_\_\_method.

## class cupy.ReductionKernel

User-defined reduction kernel.

This class can be used to define a reduction kernel with or without broadcasting.

The kernel is compiled at an invocation of the <u>\_\_\_\_\_\_</u>() method, which is cached for each device. The compiled binary is also cached into a file under the \$HOME/.cupy/kernel\_cache/directory with a hashed file name. The cached binary is reused by other processes.

#### **Parameters**

- in\_params (*str*) Input argument list.
- out\_params (*str*) Output argument list.
- **map\_expr** (*str*) Mapping expression for input values.
- **reduce\_expr** (*str*) Reduction expression.
- **post\_map\_expr** (*str*) Mapping expression for reduced values.
- identity (*str*) Identity value for starting the reduction.
- **name** (*str*) Name of the kernel function. It should be set for readability of the performance profiling.
- **reduce\_type** (*str*) Type of values to be used for reduction. This type is used to store the special variables a.
- **reduce\_dims** (*bool*) If True, input arrays are reshaped without copy to smaller dimensions for efficiency.
- **preamble** (*str*) Fragment of the CUDA-C/C++ code that is inserted at the top of the cu file.
- **options** (*tuple of str*) Additional compilation options.

## \_\_\_call\_\_()

Compiles and invokes the reduction kernel.

The compilation runs only if the kernel is not cached. Note that the kernels with different argument dtypes, ndims, or axis are not compatible. It means that single ReductionKernel object may be compiled into multiple kernel binaries.

**Parameters args** – Arguments of the kernel.

**Returns** Arrays are returned according to the out\_params argument of the \_\_init\_\_\_method.

# **Testing Modules**

CuPy offers testing utilities to support unit testing. They are under namespace *cupy.testing*.

## **Standard Assertions**

The assertions have same names as NumPy's ones. The difference from NumPy is that they can accept both numpy. ndarray and cupy.ndarray.

cupy.testing.**assert\_allclose** (*actual*, *desired*, *rtol=1e-07*, *atol=0*, *err\_msg=''*, *verbose=True*) Raises an AssertionError if objects are not equal up to desired tolerance.

## Parameters

- actual (numpy.ndarray or cupy.ndarray) The actual object to check.
- desired (numpy.ndarray or cupy.ndarray) The desired, expected object.
- **rtol** (*float*) Relative tolerance.
- **atol** (*float*) Absolute tolerance.
- **err\_msg** (*str*) The error message to be printed in case of failure.
- **verbose** (bool) If True, the conflicting values are appended to the error message.

#### See also:

```
numpy.testing.assert_allclose()
```

cupy.testing.**assert\_array\_almost\_equal** (*x*, *y*, *decimal=6*, *err\_msg=''*, *verbose=True*) Raises an AssertionError if objects are not equal up to desired precision.

#### **Parameters**

- **x** (*numpy.ndarray* or cupy.ndarray) The actual object to check.
- **y** (*numpy.ndarray* or cupy.ndarray) The desired, expected object.
- **decimal** (*int*) Desired precision.
- **err\_msg** (*str*) The error message to be printed in case of failure.
- **verbose** (*bool*) If True, the conflicting values are appended to the error message.

## See also:

numpy.testing.assert\_array\_almost\_equal()

## cupy.testing.assert\_array\_almost\_equal\_nulp(x, y, nulp=1)

Compare two arrays relatively to their spacing.

## Parameters

- **x** (*numpy.ndarray or* cupy.ndarray) The actual object to check.
- y (numpy.ndarray or cupy.ndarray) The desired, expected object.
- **nulp** (*int*) The maximum number of unit in the last place for tolerance.

## See also:

## numpy.testing.assert\_array\_almost\_equal\_nulp()

cupy.testing.assert\_array\_max\_ulp(a, b, maxulp=1, dtype=None)
Check that all items of arrays differ in at most N Units in the Last Place.

## **Parameters**

- a (numpy.ndarray or cupy.ndarray) The actual object to check.
- **b** (numpy.ndarray or cupy.ndarray) The desired, expected object.
- **maxulp** (*int*) The maximum number of units in the last place that elements of a and b can differ.
- **dtype** (*numpy*.*dtype*) Data-type to convert a and b to if given.

## See also:

## numpy.testing.assert\_array\_max\_ulp()

cupy.testing.**assert\_array\_equal** (*x*, *y*, *err\_msg=``*, *verbose=True*) Raises an AssertionError if two array\_like objects are not equal.

#### **Parameters**

- **x** (numpy.ndarray or cupy.ndarray) The actual object to check.
- y (numpy.ndarray or cupy.ndarray) The desired, expected object.
- **err\_msg** (*str*) The error message to be printed in case of failure.
- **verbose** (*bool*) If True, the conflicting values are appended to the error message.

## See also:

```
numpy.testing.assert_array_equal()
```

cupy.testing.assert\_array\_list\_equal (xlist, ylist, err\_msg='', verbose=True)
Compares lists of arrays pairwise with assert\_array\_equal.

#### **Parameters**

- **x** (*array\_like*) Array of the actual objects.
- **y** (*array\_like*) Array of the desired, expected objects.
- **err\_msg** (*str*) The error message to be printed in case of failure.
- **verbose** (*bool*) If True, the conflicting values are appended to the error message.

Each element of x and y must be either numpy.ndarray or *cupy.ndarray*. x and y must have same length. Otherwise, this function raises AssertionError. It compares elements of x and y pairwise with *assert\_array\_equal()* and raises error if at least one pair is not equal.

#### See also:

```
numpy.testing.assert_array_equal()
```

cupy.testing.**assert\_array\_less** (*x*, *y*, *err\_msg=*'', *verbose=True*) Raises an AssertionError if array\_like objects are not ordered by less than.

#### **Parameters**

- **x** (*numpy.ndarray or* cupy.ndarray) The smaller object to check.
- y (numpy.ndarray or cupy.ndarray) The larger object to compare.
- **err\_msg** (*str*) The error message to be printed in case of failure.
- **verbose** (*bool*) If True, the conflicting values are appended to the error message.

#### See also:

```
numpy.testing.assert_array_less()
```

## NumPy-CuPy Consistency Check

The following decorators are for testing consistency between CuPy's functions and corresponding NumPy's ones.

Decorator that checks NumPy results and CuPy ones are close.

## Parameters

- **rtol** (*float*) Relative tolerance.
- **atol** (float) Absolute tolerance
- **err\_msg** (*str*) The error message to be printed in case of failure.
- **verbose** (*bool*) If True, the conflicting values are appended to the error message.
- **name** (*str*) Argument name whose value is either numpy or cupy module.
- type\_check (bool) If True, consistency of dtype is also checked.
- accept\_error (bool, Exception or tuple of Exception) Sepcify acceptable errors. When both NumPy test and CuPy test raises the same type of errors, and the type of the errors is specified with this argument, the errors are ignored and not raised. If it is True all error types are acceptable. If it is False no error is acceptable.

Decorated test fixture is required to return the arrays whose values are close between numpy case and cupy case. For example, this test case checks numpy.zeros and cupy.zeros should return same value.

```
>>> import unittest
>>> from cupy import testing
>>> @testing.gpu
... class TestFoo (unittest.TestCase):
. . .
         @testing.numpy_cupy_allclose()
. . .
         def test_foo(self, xp):
. . .
             # ...
. . .
             # Prepare data with xp
. . .
             # ...
. . .
. . .
             xp_result = xp.zeros(10)
. . .
             return xp_result
. . .
```

See also:

```
cupy.testing.assert_allclose()
```

```
cupy.testing.numpy_cupy_array_almost_equal(decimal=6, err_msg='', verbose=True,
name='xp', type_check=True, ac-
cept_error=False)
```

Decorator that checks NumPy results and CuPy ones are almost equal.

### Parameters

- **decimal** (*int*) Desired precision.
- **err\_msg** (*str*) The error message to be printed in case of failure.
- **verbose** (*bool*) If True, the conflicting values are appended to the error message.
- **name** (*str*) Argument name whose value is either numpy or cupy module.
- type\_check (bool) If True, consistency of dtype is also checked.

• accept\_error (bool, Exception or tuple of Exception) – Sepcify acceptable errors. When both NumPy test and CuPy test raises the same type of errors, and the type of the errors is specified with this argument, the errors are ignored and not raised. If it is True all error types are acceptable. If it is False no error is acceptable.

Decorated test fixture is required to return the same arrays in the sense of *cupy.testing*. assert array almost equal() (except the type of array module) even if xp is numpy or cupy.

#### See also:

cupy.testing.assert\_array\_almost\_equal()

Decorator that checks results of NumPy and CuPy are equal w.r.t. spacing.

#### **Parameters**

- **nulp** (*int*) The maximum number of unit in the last place for tolerance.
- **name** (*str*) Argument name whose value is either numpy or cupy module.
- type\_check (bool) If True, consistency of dtype is also checked.
- accept\_error (bool, Exception or tuple of Exception) Sepcify acceptable errors. When both NumPy test and CuPy test raises the same type of errors, and the type of the errors is specified with this argument, the errors are ignored and not raised. If it is True all error types are acceptable. If it is False no error is acceptable.

Decorated test fixture is required to return the same arrays in the sense of *cupy.testing. assert\_array\_almost\_equal\_nulp()* (except the type of array module) even if xp is numpy or cupy.

#### See also:

cupy.testing.assert\_array\_almost\_equal\_nulp()

cupy.testing.numpy\_cupy\_array\_max\_ulp (maxulp=1, dtype=None, name='xp', type\_check=True, accept\_error=False) Decorator that checks results of NumPy and CuPy ones are equal w.r.t. ulp.

#### **Parameters**

- **maxulp** (*int*) The maximum number of units in the last place that elements of resulting two arrays can differ.
- **dtype** (*numpy.dtype*) Data-type to convert the resulting two array to if given.
- **name** (*str*) Argument name whose value is either numpy or cupy module.
- type\_check (bool) If True, consistency of dtype is also checked.
- accept\_error (bool, Exception or tuple of Exception) Sepcify acceptable errors. When both NumPy test and CuPy test raises the same type of errors, and the type of the errors is specified with this argument, the errors are ignored and not raised. If it is True all error types are acceptable. If it is False no error is acceptable.

Decorated test fixture is required to return the same arrays in the sense of *assert\_array\_max\_ulp()* (except the type of array module) even if xp is numpy or cupy.

## See also:

cupy.testing.assert\_array\_max\_ulp()

```
cupy.testing.numpy_cupy_array_equal(err_msg=``, verbose=True, name=`xp`, type_check=True, accept_error=False)
```

Decorator that checks NumPy results and CuPy ones are equal.

## Parameters

- **err\_msg** (*str*) The error message to be printed in case of failure.
- **verbose** (*bool*) If True, the conflicting values are appended to the error message.
- **name** (*str*) Argument name whose value is either numpy or cupy module.
- **type\_check** (*bool*) If True, consistency of dtype is also checked.
- accept\_error (bool, Exception or tuple of Exception) Sepcify acceptable errors. When both NumPy test and CuPy test raises the same type of errors, and the type of the errors is specified with this argument, the errors are ignored and not raised. If it is True all error types are acceptable. If it is False no error is acceptable.

Decorated test fixture is required to return the same arrays in the sense of *numpy\_cupy\_array\_equal()* (except the type of array module) even if xp is numpy or cupy.

## See also:

```
cupy.testing.assert_array_equal()
```

cupy.testing.numpy\_cupy\_array\_list\_equal (*err\_msg=``*, *verbose=True*, *name='xp'*) Decorator that checks the resulting lists of NumPy and CuPy's one are equal.

## Parameters

- **err\_msg** (*str*) The error message to be printed in case of failure.
- **verbose** (*bool*) If True, the conflicting values are appended to the error message.
- name (str) Argument name whose value is either numpy or cupy module.

Decorated test fixture is required to return the same list of arrays (except the type of array module) even if xp is numpy or cupy.

## See also:

```
cupy.testing.assert_array_list_equal()
```

```
cupy.testing.numpy_cupy_array_less(err_msg='', verbose=True, name='xp', type_check=True,
```

*accept\_error=False*) Decorator that checks the CuPy result is less than NumPy result.

## **Parameters**

- **err\_msg** (*str*) The error message to be printed in case of failure.
- **verbose** (*bool*) If True, the conflicting values are appended to the error message.
- **name** (*str*) Argument name whose value is either numpy or cupy module.
- **type\_check** (*bool*) If True, consistency of dtype is also checked.
- accept\_error (bool, Exception or tuple of Exception) Sepcify acceptable errors. When both NumPy test and CuPy test raises the same type of errors, and the type of the errors is specified with this argument, the errors are ignored and not raised. If it is True all error types are acceptable. If it is False no error is acceptable.

Decorated test fixture is required to return the smaller array when xp is cupy than the one when xp is numpy.

## See also:

```
cupy.testing.assert_array_less()
```

cupy.testing.numpy\_cupy\_raises(name='xp')

Decorator that checks the NumPy and CuPy throw same errors.

### **Parameters**

- **name** (*str*) Argument name whose value is either
- or cupy module. (numpy) -

Decorated test fixture is required throw same errors even if xp is numpy or cupy.

See also:

```
cupy.testing.assert_array_less()
```

## **Parameterized dtype Test**

The following decorators offer the standard way for parameterized test with respect to single or the combination of dtype(s).

```
cupy.testing.for_dtypes(dtypes, name='dtype')
Decorator for parameterized dtype test.
```

#### **Parameters**

- dtypes (list of dtypes) dtypes to be tested.
- **name** (*str*) Argument name to which specified dtypes are passed.

This decorator adds a keyword argument specified by name to the test fixture. Then, it runs the fixtures in parallel by passing the each element of dtypes to the named argument.

cupy.testing.for\_all\_dtypes (name='dtype', no\_float16=False, no\_bool=False)
Decorator that checks the fixture with all dtypes.

#### **Parameters**

- **name** (*str*) Argument name to which specified dtypes are passed.
- no\_float16 (bool) If, True, numpy. float16 is omitted from candidate dtypes.
- no\_bool (bool) If, True, numpy.bool\_ is omitted from candidate dtypes.

```
dtypes to be tested: numpy.float16 (optional), numpy.float32, numpy.float64, numpy.
dtype('b'), numpy.dtype('h'), numpy.dtype('i'), numpy.dtype('l'), numpy.
dtype('q'), numpy.dtype('B'), numpy.dtype('H'), numpy.dtype('I'), numpy.
dtype('L'), numpy.dtype('Q'), and numpy.bool_(optional).
```

The usage is as follows. This test fixture checks if cPickle successfully reconstructs *cupy.ndarray* for various dtypes. dtype is an argument inserted by the decorator.

```
>>> import unittest
>>> from cupy import testing
>>> @testing.gpu
... class TestNpz (unittest.TestCase):
. . .
        @testing.for_all_dtypes()
. . .
        def test_pickle(self, dtype):
. . .
            a = testing.shaped_arange((2, 3, 4), dtype=dtype)
. . .
            s = six.moves.cPickle.dumps(a)
. . .
            b = six.moves.cPickle.loads(s)
. . .
            testing.assert_array_equal(a, b)
. . .
```

Typically, we use this decorator in combination with decorators that check consistency between NumPy and CuPy like *cupy\_testing.numpy\_cupy\_allclose()*. The following is such an example.

```
>>> import unittest
>>> from cupy import testing
>>> @testing.gpu
... class TestMean(unittest.TestCase):
...
...
...
@testing.for_all_dtypes()
...
@testing.numpy_cupy_allclose()
...
def test_mean_all(self, xp, dtype):
...
a = testing.shaped_arange((2, 3), xp, dtype)
...
return a.mean()
```

#### See also:

cupy.testing.for\_dtypes()

cupy.testing.**for\_float\_dtypes** (*name='dtype'*, *no\_float16=False*) Decorator that checks the fixture with all float dtypes.

#### **Parameters**

- **name** (*str*) Argument name to which specified dtypes are passed.
- no\_float16 (bool) If, True, numpy. float16 is omitted from candidate dtypes.

dtypes to be tested are numpy.float16 (optional), numpy.float32, and numpy.float64.

#### See also:

cupy.testing.for\_dtypes(), cupy.testing.for\_all\_dtypes()

## cupy.testing.for\_signed\_dtypes(name='dtype')

Decorator that checks the fixture with signed dtypes.

**Parameters name** (*str*) – Argument name to which specified dtypes are passed.

dtypes to be tested are numpy.dtype('b'), numpy.dtype('h'), numpy.dtype('i'), numpy. dtype('l'), and numpy.dtype('q').

#### See also:

cupy.testing.for\_dtypes(), cupy.testing.for\_all\_dtypes()

#### cupy.testing.for\_unsigned\_dtypes(name='dtype')

Decorator that checks the fixture with all dtypes.

**Parameters name** (*str*) – Argument name to which specified dtypes are passed.

dtypes to be tested are numpy.dtype('B'), numpy.dtype('H'),

numpy.dtype('I'), numpy.dtype('L'), and numpy.dtype('Q').

#### See also:

cupy.testing.for\_dtypes(), cupy.testing.for\_all\_dtypes()

cupy.testing.for\_int\_dtypes (name='dtype', no\_bool=False)

Decorator that checks the fixture with integer and optionally bool dtypes.

#### Parameters

- **name** (*str*) Argument name to which specified dtypes are passed.
- no\_bool (bool) If True, numpy.bool\_ is omitted from candidate dtypes.

```
dtypes to be tested are numpy.dtype('b'), numpy.dtype('h'), numpy.dtype('i'), numpy.
dtype('l'), numpy.dtype('q'), numpy.dtype('B'), numpy.dtype('H'), numpy.
dtype('I'), numpy.dtype('L'), numpy.dtype('Q'), and numpy.bool_(optional).
```

#### See also:

cupy.testing.for\_dtypes(), cupy.testing.for\_all\_dtypes()

cupy.testing.for\_dtypes\_combination(types, names=('dtype', ), full=None)
Decorator that checks the fixture with a product set of dtypes.

#### **Parameters**

- types (list of dtypes) dtypes to be tested.
- **names** (list of str) Argument names to which dtypes are passed.
- **full** (*bool*) If True, then all combinations of dtypes will be tested. Otherwise, the subset of combinations will be tested (see the description below).

Decorator adds the keyword arguments specified by names to the test fixture. Then, it runs the fixtures in parallel with passing (possibly a subset of) the product set of dtypes. The range of dtypes is specified by types.

The combination of dtypes to be tested changes depending on the option full. If full is True, all combinations of types are tested. Sometimes, such an exhaustive test can be costly. So, if full is False, only the subset of possible combinations is tested. Specifically, at first, the shuffled lists of types are made for each argument name in names. Let the lists be D1, D2, ..., Dn where n is the number of arguments. Then, the combinations to be tested will be zip (D1, ..., Dn). If full is None, the behavior is switched by setting the environment variable CUPY\_TEST\_FULL\_COMBINATION=1.

For example, let types be [float16, float32, float64] and names be ['a\_type', 'b\_type']. If full is True, then the decorated test fixture is executed with all 2<sup>3</sup> patterns. On the other hand, if full is False, shuffled lists are made for a\_type and b\_type. Suppose the lists are (16, 64, 32) for a\_type and (32, 64, 16) for b\_type (prefixes are removed for short). Then the combinations of (a\_type, b\_type) to be tested are (16, 32), (64, 64) and (32, 16).

Decorator that checks the fixture with a product set of all dtypes.

## **Parameters**

- **names** (list of str) Argument names to which dtypes are passed.
- **no\_float16** (*bool*) If True, numpy.float16 is omitted from candidate dtypes.
- no\_bool (bool) If True, numpy.bool\_ is omitted from candidate dtypes.
- **full** (*bool*) If True, then all combinations of dtypes will be tested. Otherwise, the subset of combinations will be tested (see description in *cupy.testing. for\_dtypes\_combination()*).

#### See also:

cupy.testing.for\_dtypes\_combination()

cupy.testing.for\_signed\_dtypes\_combination (names=('dtype', ), full=None)
Decorator for parameterized test w.r.t. the product set of signed dtypes.

#### **Parameters**

• **names** (list of str) – Argument names to which dtypes are passed.

• **full** (bool) - If True, then all combinations of dtypes will be tested. Otherwise, the subset of combinations will be tested (see description in *cupy.testing.* for\_dtypes\_combination()).

#### See also:

#### cupy.testing.for\_dtypes\_combination()

cupy.testing.for\_unsigned\_dtypes\_combination (names=('dtype', ), full=None)
Decorator for parameterized test w.r.t. the product set of unsigned dtypes.

### **Parameters**

- **names** (list of str) Argument names to which dtypes are passed.
- **full** (*bool*) If True, then all combinations of dtypes will be tested. Otherwise, the subset of combinations will be tested (see description in *cupy.testing. for\_dtypes\_combination()*).

## See also:

#### cupy.testing.for\_dtypes\_combination()

cupy.testing.for\_int\_dtypes\_combination (names=('dtype', ), no\_bool=False, full=None)
Decorator for parameterized test w.r.t. the product set of int and boolean.

### Parameters

- **names** (*list of str*) Argument names to which dtypes are passed.
- no\_bool (bool) If True, numpy.bool\_ is omitted from candidate dtypes.
- **full** (*bool*) If True, then all combinations of dtypes will be tested. Otherwise, the subset of combinations will be tested (see description in *cupy.testing. for\_dtypes\_combination()*).

## See also:

cupy.testing.for\_dtypes\_combination()

## **Parameterized order Test**

The following decorators offer the standard way to parameterize tests with orders.

```
cupy.testing.for_orders(orders, name='order')
```

Decorator to parameterize tests with order.

## **Parameters**

- orders (list of orders) orders to be tested.
- **name** (*str*) Argument name to which the specified order is passed.

This decorator adds a keyword argument specified by name to the test fixtures. Then, the fixtures run by passing each element of orders to the named argument.

```
cupy.testing.for_CF_orders (name='order')
```

Decorator that checks the fixture with orders 'C' and 'F'.

**Parameters name** (*str*) – Argument name to which the specified order is passed.

## See also:

```
cupy.testing.for_all_dtypes()
```

#### **Environment variables**

Here are the environment variables Chainer uses.

```
CUPY_CACHE_DPath to the directory to store kernel cache. $ (HOME) /.cupy.kernel_cache is used by default. See CuPy Overview for detail.
```

#### For install

These environment variables are only used during installation.

CUDA\_PATPath to the directory containing CUDA. The parent of the directory containing nvcc is used as default. When nvcc is not found, /usr/local/cuda is used. See *Install Chainer with CUDA* for details.

#### **Difference between CuPy and NumPy**

The interface of CuPy is designed to obey that of NumPy. However, there are some differences.

#### Cast behavior from float to integer

Some casting behaviors from float to integer are not defined in C++ specification. The casting from a negative float to unsigned integer and infinity to integer is one of such eamples. The behavior of NumPy depends on your CPU architecture. This is Intel CPU result.

```
>>> np.array([-1], dtype='f').astype('I')
array([4294967295], dtype=uint32)
>>> cupy.array([-1], dtype='f').astype('I')
array([0], dtype=uint32)
```

```
>>> np.array([float('inf')], dtype='f').astype('i')
array([-2147483648], dtype=int32)
>>> cupy.array([float('inf')], dtype='f').astype('i')
array([2147483647], dtype=int32)
```

#### **Boolean values squared**

In NumPy implementation,  $x \star 2$  is calculated using multiplication operator as  $x \star x$ . Because the result of the multiplication of boolean values is boolean, True  $\star 2$  return boolean value. However, when you use power operator with other arguments, it returns int values. If we aligned the behavior of the squared boolean values of CuPy to that of NumPy, we would have to check their values in advance of the calculation. But it would be slow because it would force CPUs to wait until the calculation on GPUs end. So we decided not to check its value.

```
>>> np.array([True]) ** 2
array([ True], dtype=bool)
>>> cupy.array([True]) ** 2
array([1])
```

#### Random methods support dtype argument

NumPy's random value generator does not support dtype option and it always resturns a float 32 value. We support the option in CuPy because cuRAND, which is used in CuPy, supports any types of float values.

```
>>> np.random.randn(dtype='f')
Traceback (most recent call last):
   File "<stdin>", line 1, in <module>
TypeError: randn() got an unexpected keyword argument 'dtype'
>>> cupy.random.randn(dtype='f')
array(0.10689262300729752, dtype=float32)
```

#### **Out-of-bounds indices**

CuPy handles out-of-bounds indices differently by default from NumPy when using integer array indexing. NumPy handles them by raising an error, but CuPy wraps around them.

```
>>> x = np.array([0, 1, 2])
>>> x[[1, 3]] = 10
Traceback (most recent call last):
    File "<stdin>", line 1, in <module>
IndexError: index 3 is out of bounds for axis 1 with size 3
>>> x = cupy.array([0, 1, 2])
>>> x[[1, 3]] = 10
>>> x
array([10, 10, 2])
```

#### Duplicate values in indices

CuPy's \_\_\_setitem\_\_ behaves differently from NumPy when integer arrays reference the same location multiple times. In that case, the value that is actually stored is undefined. Here is an example of CuPy.

```
>>> a = cupy.zeros((2,))
>>> i = cupy.arange(10000) % 2
>>> v = cupy.arange(10000).astype(np.float)
>>> a[i] = v
>>> a
array([ 9150., 9151.])
```

NumPy stores the value corresponding to the last element among elements referencing duplicate locations.

```
>>> a_cpu = np.zeros((2,))
>>> i_cpu = np.arange(10000) % 2
>>> v_cpu = np.arange(10000).astype(np.float)
>>> a_cpu[i_cpu] = v_cpu
>>> a_cpu
array([ 9998., 9999.])
```

## CHAPTER 5

#### Chainer Contribution Guide

This is a guide for all contributions to Chainer. The development of Chainer is running on the official repository at GitHub. Anyone that wants to register an issue or to send a pull request should read through this document.

#### **Classification of Contributions**

There are several ways to contribute to Chainer community:

- 1. Registering an issue
- 2. Sending a pull request (PR)
- 3. Sending a question to Chainer User Group
- 4. Open-sourcing an external example
- 5. Writing a post about Chainer

This document mainly focuses on 1 and 2, though other contributions are also appreciated.

#### **Release and Milestone**

We are using GitHub Flow as our basic working process. In particular, we are using the master branch for our development, and releases are made as tags.

Releases are classified into three groups: major, minor, and revision. This classification is based on following criteria:

- Major update contains disruptive changes that break the backward compatibility.
- Minor update contains additions and extensions to the APIs keeping the supported backward compatibility.
- Revision update contains improvements on the API implementations without changing any API specification.

The release classification is reflected into the version number x.y.z, where x, y, and z corresponds to major, minor, and revision updates, respectively.

We set a milestone for an upcoming release. The milestone is of name 'vX.Y.Z', where the version number represents a revision release at the outset. If at least one *feature* PR is merged in the period, we rename the milestone to represent a minor release (see the next section for the PR types).

See also API Compatibility Policy.

#### **Issues and PRs**

Issues and PRs are classified into following categories:

- Bug: bug reports (issues) and bug fixes (PRs)
- Enhancement: implementation improvements without breaking the interface
- Feature: feature requests (issues) and their implementations (PRs)
- NoCompat: disrupts backward compatibility
- **Test**: test fixes and updates
- Document: document fixes and improvements
- Example: fixes and improvements on the examples
- Install: fixes installation script
- Contribution-Welcome: issues that we request for contribution (only issues are categorized to this)
- Other: other issues and PRs

Issues and PRs are labeled by these categories. This classification is often reflected into its corresponding release category: Feature issues/PRs are contained into minor/major releases and NoCompat issues/PRs are contained into major releases, while other issues/PRs can be contained into any releases including revision ones.

On registering an issue, write precise explanations on what you want Chainer to be. Bug reports must include necessary and sufficient conditions to reproduce the bugs. Feature requests must include **what** you want to do (and **why** you want to do, if needed). You can contain your thoughts on **how** to realize it into the feature requests, though **what** part is most important for discussions.

**Warning:** If you have a question on usages of Chainer, it is highly recommended to send a post to Chainer User Group instead of the issue tracker. The issue tracker is not a place to share knowledge on practices. We may redirect question issues to Chainer User Group.

If you can write code to fix an issue, send a PR to the master branch. Before writing your code for PRs, read through the *Coding Guidelines*. The description of any PR must contain a precise explanation of **what** and **how** you want to do; it is the first documentation of your code for developers, a very important part of your PR.

Once you send a PR, it is automatically tested on Travis CI for Linux and Mac OS X, and on AppVeyor for Windows. Your PR need to pass at least the test for Linux on Travis CI. After the automatic test passes, some of the core developers will start reviewing your code. Note that this automatic PR test only includes CPU tests.

**Note:** We are also running continuous integration with GPU tests for the master branch. Since this service is running on our internal server, we do not use it for automatic PR tests to keep the server secure.

Even if your code is not complete, you can send a pull request as a *work-in-progress PR* by putting the [WIP] prefix to the PR title. If you write a precise explanation about the PR, core developers and other contributors can join the discussion about how to proceed the PR.

#### **Coding Guidelines**

We use PEP8 and a part of OpenStack Style Guidelines related to general coding style as our basic style guidelines.

To check your code, use autopep8 and flake8 command installed by hacking package:

```
$ pip install autopep8 hacking
$ autopep8 --global-config .pep8 path/to/your/code.py
$ flake8 path/to/your/code.py
```

To check Cython code, use .flake8.cython configuration file:

\$ flake8 --config=.flake8.cython path/to/your/cython/code.pyx

The autopep8 supports automatically correct Python code to conform to the PEP 8 style guide:

\$ autopep8 --in-place --global-config .pep8 path/to/your/code.py

The flake8 command lets you know the part of your code not obeying our style guidelines. Before sending a pull request, be sure to check that your code passes the flake8 checking.

Note that flake8 command is not perfect. It does not check some of the style guidelines. Here is a (not-complete) list of the rules that flake8 cannot check.

- Relative imports are prohibited. [H304]
- Importing non-module symbols is prohibited.
- Import statements must be organized into three parts: standard libraries, third-party libraries, and internal imports. [H306]

In addition, we restrict the usage of *shortcut symbols* in our code base. They are symbols imported by packages and sub-packages of chainer. For example, chainer.Variable is a shortcut of chainer.variable. Variable. It is not allowed to use such shortcuts in the "chainer" library implementation. Note that you can still use them in tests and examples directories. Also note that you should use shortcut names of CuPy APIs in Chainer implementation.

Once you send a pull request, your coding style is automatically checked by Travis-CI. The reviewing process starts after the check passes.

The CuPy is designed based on NumPy's API design. CuPy's source code and documents contain the original NumPy ones. Please note the followings when writing the document.

- In order to identify overlapping parts, it is preferable to add some remarks that this document is just copied or altered from the original one. It is also preferable to briefly explain the specification of the function in a short paragraph, and refer to the corresponding function in NumPy so that users can read the detailed document. However, it is possible to include a complete copy of the document with such a remark if users cannot summarize in such a way.
- If a function in CuPy only implements a limited amount of features in the original one, users should explicitly describe only what is implemented in the document.

### **Testing Guidelines**

Testing is one of the most important part of your code. You must test your code by unit tests following our testing guidelines. Note that we are using the nose package and the mock package for testing, so install nose and mock before writing your code:

\$ pip install nose mock

In order to run unit tests at the repository root, you first have to build Cython files in place by running the following command:

\$ python setup.py develop

Once the Cython modules are built, you can run unit tests simply by running nosetests command at the repository root:

\$ nosetests

It requires CUDA by default. In order to run unit tests that do not require CUDA, pass --attr='!gpu' option to the nosetests command:

\$ nosetests path/to/your/test.py --attr='!gpu'

Some GPU tests involve multiple GPUs. If you want to run GPU tests with insufficient number of GPUs, specify the number of available GPUs by --eval-attr='gpu<N' where N is a concrete integer. For example, if you have only one GPU, launch nosetests by the following command to skip multi-GPU tests:

\$ nosetests path/to/gpu/test.py --eval-attr='gpu<2'</pre>

Some tests spend too much time. If you want to skip such tests, pass --attr='!slow' option to the nosetests command:

\$ nosetests path/to/your/test.py --attr='!slow'

Tests are put into the tests/chainer\_tests, tests/cupy\_tests and tests/install\_tests directories. These have the same structure as that of chainer, cupy and install directories, respectively. In order to enable test runner to find test scripts correctly, we are using special naming convention for the test subdirectories and the test scripts.

- The name of each subdirectory of tests must end with the \_tests suffix.
- The name of each test script must start with the test\_prefix.

Following this naming convention, you can run all the tests by just typing nosetests at the repository root:

```
$ nosetests
```

Or you can also specify a root directory to search test scripts from:

```
$ nosetests tests/chainer_tests # to just run tests of Chainer
$ nosetests tests/cupy_tests # to just run tests of CuPy
$ nosetests tests/install_tests # to just run tests of installation modules
```

If you modify the code related to existing unit tests, you must run appropriate commands.

**Note:** CuPy tests include type-exhaustive test functions which take long time to execute. If you are running tests on a multi-core machine, you can parallelize the tests by following options:

\$ nosetests --processes=12 --process-timeout=1000 tests/cupy\_tests

The magic numbers can be modified for your usage. Note that some tests require many CUDA compilations, which require a bit long time. Without the process-timeout option, the timeout is set shorter, causing timeout failures

for many test cases.

There are many examples of unit tests under the tests directory. They simply use the unittest package of the standard library.

Even if your patch includes GPU-related code, your tests should not fail without GPU capability. Test functions that require CUDA must be tagged by the chainer.testing.attr.gpu decorator (or cupy.testing.attr.gpu for testing CuPy APIs):

The functions tagged by the gpu decorator are skipped if --attr='!gpu' is given. We also have the chainer. testing.attr.cudnn decorator to let nosetests know that the test depends on cuDNN.

The test functions decorated by gpu must not depend on multiple GPUs. In order to write tests for multiple GPUs, use chainer.testing.attr.multi\_gpu() or cupy.testing.attr.multi\_gpu() decorators instead:

```
import unittest
from chainer.testing import attr

class TestMyFunc(unittest.TestCase):
    ...
    @attr.multi_gpu(2)  # specify the number of required GPUs here
    def test_my_two_gpu_func(self):
    ...
```

If your test requires too much time, add chainer.testing.attr.slow decorator. The test functions decorated by slow are skipped if --attr='!slow' is given:

**Note:** If you want to specify more than two attributes, separate them with a comma such as --attr='!gpu, ! slow'. See detail in the document of nose.

Once you send a pull request, your code is automatically tested by Travis-CI with -attr='!gpu,!slow' option. Since Travis-CI does not support CUDA, we cannot check your CUDA-related code automatically. The reviewing process starts after the test passes. Note that reviewers will test your code without the option to check CUDA-related code.

**Note:** Some of numerically unstable tests might cause errors irrelevant to your changes. In such a case, we ignore the failures and go on to the review process, so do not worry about it.

## CHAPTER 6

### **API** Compatibility Policy

This document expresses the design policy on compatibilities of Chainer APIs. Development team should obey this policy on deciding to add, extend, and change APIs and their behaviors.

This document is written for both users and developers. Users can decide the level of dependencies on Chainer's implementations in their codes based on this document. Developers should read through this document before creating pull requests that contain changes on the interface. Note that this document may contain ambiguities on the level of supported compatibilities.

## **Targeted Versions**

This policy is applied to Chainer of versions v1.5.1 and higher. Note that this policy is not applied to Chainer of lower versions.

### **Versioning and Backward Compatibilities**

The updates of Chainer are classified into three levels: major, minor, and revision. These types have distinct levels of backward compatibilities.

- Major update contains disruptive changes that break the backward compatibility.
- Minor update contains addition and extension to the APIs keeping the supported backward compatibility.
- Revision update contains improvements on the API implementations without changing any API specifications.

Note that we do not support full backward compatibility, which is almost infeasible for Python-based APIs, since there is no way to completely hide the implementation details.

## **Processes to Break Backward Compatibilities**

#### Deprecation, Dropping, and Its Preparation

Any APIs may be *deprecated* at some minor updates. In such a case, the deprecation note is added to the API documentation, and the API implementation is changed to fire deprecation warning (if possible). There should be another way to reimplement the same things previously written with the deprecated APIs.

Any APIs may be marked as *to be dropped in the future*. In such a case, the dropping is stated in the documentation with the major version number on which the API is planned to be dropped, and the API implementation is changed to fire the future warning (if possible).

The actual dropping should be done through the following steps:

- Make the API deprecated. At this point, users should not need the deprecated API in their new application codes.
- After that, mark the API as *to be dropped in the future*. It must be done in the minor update different from that of the deprecation.
- At the major version announced in the above update, drop the API.

Consequently, it takes at least two minor versions to drop any APIs after the first deprecation.

#### **API Changes and Its Preparation**

Any APIs may be marked as *to be changed in the future* for changes without backward compatibility. In such a case, the change is stated in the documentation with the version number on which the API is planned to be changed, and the API implementation is changed to fire the future warning on the certain usages.

The actual change should be done in the following steps:

- Announce that the API will be changed in the future. At this point, the actual version of change need not be accurate.
- After the announcement, mark the API as *to be changed in the future* with version number of planned changes. At this point, users should not use the marked API in their new application codes.
- At the major update announced in the above update, change the API.

## **Experimental APIs**

Thanks to many contributors, we have introduced many new features to Chainer.

However, we have sometimes released new features only to later notice that their APIs are not appropriate. The objective of experimental APIs is to avoid such issues by allowing the developer to mark any newly added API as experimental.

Any newly added API can be marked as *experimental*. Any API that is not experimental is called *stable* in this document.

**Note:** Undocumented behaviors are not considered as APIs. So they are not experimental nor stable. The treatment of undocumented behaviors are described in *Undocumented behaviors* section.

Chainer can change the interfaces and documents of experimental APIs at **any** version up. This change is not considered as a break of backward compatibility. Chainer can promote an experimental API to become stable at any **minor** or **major** version up. Once experimental APIs become stable, they cannot revert to experimental again.

When users use experimental APIs for the first time, warnings are raised once for each experimental API, unless users explicitly disable the emission of the warnings in advance.

See the document of *chainer.utils.experimental()* how developers mark APIs as experimental and how users enable or disable the warnings practically.

**Note:** It is up to developers if APIs should be annotated as experimental or not. We recommend to make the APIs experimental if they implement large modules or make a decision from several design choices.

#### **Supported Backward Compatibility**

This section defines backward compatibilities that minor updates must maintain.

#### **Documented Interface**

Chainer has the official API documentation. Many applications can be written based on the documented features. We support backward compatibilities of documented features. In other words, codes only based on the documented features run correctly with minor/revision-updated versions.

Developers are encouraged to use apparent names for objects of implementation details. For example, attributes outside of the documented APIs should have one or more underscores at the prefix of their names.

#### **Undocumented behaviors**

Behaviors of Chainer implementation not stated in the documentation are undefined. Undocumented behaviors are not guaranteed to be stable between different minor/revision versions.

Minor update may contain changes to undocumented behaviors. For example, suppose an API X is added at the minor update. In the previous version, attempts to use X cause AttributeError. This behavior is not stated in the documentation, so this is undefined. Thus, adding the API X in minor version is permissible.

Revision update may also contain changes to undefined behaviors. Typical example is a bug fix. Another example is an improvement on implementation, which may change the internal object structures not shown in the documentation. As a consequence, even revision updates do not support compatibility of pickling, unless the full layout of pickled objects is clearly documented.

#### **Documentation Error**

Compatibility is basically determined based on the documentation, though it sometimes contains errors. It may make the APIs confusing to assume the documentation always stronger than the implementations. We therefore may fix the documentation errors in any updates that may break the compatibility in regard to the documentation.

**Note:** Developers MUST NOT fix the documentation and implementation of the same functionality at the same time in revision updates as "bug fix". Such a change completely breaks the backward compatibility. If you want to fix the

bugs in both sides, first fix the documentation to fit it into the implementation, and start the API changing procedure described above.

#### **Object Attributes and Properties**

Object attributes and properties are sometimes replaced by each other at minor updates. It does not break the user codes, except the codes depend on how the attributes and properties are implemented.

#### **Functions and Methods**

Methods may be replaced by callable attributes keeping the compatibility of parameters and return values in minor updates. It does not break the user codes, except the codes depend on how the methods and callable attributes are implemented.

#### **Exceptions and Warnings**

The specifications of raising exceptions are considered as a part of standard backward compatibilities. No exception is raised in the future versions with correct usages that the documentation allows, unless the API changing process is completed.

On the other hand, warnings may be added at any minor updates for any APIs. It means minor updates do not keep backward compatibility of warnings.

#### Model Format Compatibility

Objects serialized by official serializers that Chainer provides are correctly loaded with the higher (future) versions. They might not be correctly loaded with Chainer of the lower versions.

**Note:** Current serialization APIs do not support versioning (at least in v1.6.1). It prevents us from introducing changes in the layout of objects that support serialization. We are discussing about introducing versioning in serialization APIs.

#### Installation Compatibility

The installation process is another concern of compatibilities. We support environmental compatibilities in the following ways.

- Any changes of dependent libraries that force modifications on the existing environments must be done in major updates. Such changes include following cases:
  - dropping supported versions of dependent libraries (e.g. dropping cuDNN v2)
  - adding new mandatory dependencies (e.g. adding h5py to setup\_requires)
- Supporting optional packages/libraries may be done in minor updates (e.g. supporting h5py in optional features).

**Note:** The installation compatibility does not guarantee that all the features of Chainer correctly run on supported environments. It may contain bugs that only occurs in certain environments. Such bugs should be fixed in some updates.

## CHAPTER 7

Tips and FAQs

# It takes too long time to compile a computational graph. Can I skip it?

Chainer does not compile computational graphs, so you cannot skip it, or, I mean, you have already skipped it :).

It seems you have actually seen on-the-fly compilations of CUDA kernels. CuPy compiles kernels on demand to make kernels optimized to the number of dimensions and element types of input arguments. Pre-compilation is not available, because we have to compile an exponential number of kernels to support all CuPy functionalities. This restriction is unavoidable because Python cannot call CUDA/C++ template functions in generic way. Note that every framework using CUDA require compilation at some point; the difference between other statically-compiled frameworks (such as cutorch) and Chainer is whether a kernel is compiled at installation or at the first use.

These compilations should run only at the first use of the kernels. The compiled binaries are cached to the \$ (HOME) / .cupy/kernel\_cache directory by default. If you see that compilations run every time you run the same script, then the caching is failed. Please check that the directory is kept as is between multiple executions of the script. If your home directory is not suited to caching the kernels (e.g. in case that it uses NFS), change the kernel caching directory by setting the CUPY\_CACHE\_DIR environment variable to an appropriate path. See *CuPy Overview* for more details.

### mnist example does not converge in CPU mode on Mac OS X

Many users reported that mnist example does not work correctly on Mac OS X. We are suspecting it is caused by vecLib, that is a default BLAS library installed on Mac OS X.

Note: Mac OS X is not officially supported. I mean it is not tested continuously on our test server.

We recommend to use other BLAS libraries such as OpenBLAS. We empirically found that it fixes this problem. It is necessary to reinstall NumPy to use replaced BLAS library. Here is an instruction to install NumPy with OpenBLAS using Homebrew.

```
$ brew tap homebrew/science
$ brew install openblas
$ brew install numpy --with-openblas
```

If you want to install NumPy with pip, use site.cfg file.

You can check if NumPy uses OpenBLAS with numpy.show\_config method. Check if *blas\_opt\_info* refers to *openblas*.

```
>>> import numpy
>>> numpy.show_config()
lapack_opt_info:
   libraries = ['openblas', 'openblas']
   library_dirs = ['/usr/local/opt/openblas/lib']
   define_macros = [('HAVE_CBLAS', None)]
   language = c
blas_opt_info:
   libraries = ['openblas', 'openblas']
    library_dirs = ['/usr/local/opt/openblas/lib']
   define_macros = [('HAVE_CBLAS', None)]
   language = c
openblas_info:
   libraries = ['openblas', 'openblas']
   library_dirs = ['/usr/local/opt/openblas/lib']
   define_macros = [('HAVE_CBLAS', None)]
   language = c
openblas_lapack_info:
   libraries = ['openblas', 'openblas']
   library_dirs = ['/usr/local/opt/openblas/lib']
   define_macros = [('HAVE_CBLAS', None)]
   language = c
blas_mkl_info:
   NOT AVAILABLE
```

See detail about this problem in issue #704.

## CHAPTER 8

## Comparison with Other Frameworks

## A table for quick comparison

This table compares Chainer with other popular deep learning frameworks. We hope it helps you to choose an appropriate framework for the demand.

**Note:** This chart may be out-dated, since the developers of Chainer do not perfectly follow the latest development status of each framework. Please report us if you find an out-dated cell. Requests for new comparison axes are also welcome.

		Chainer	Theano-based	Torch7	Caffe
Specs	Scripting	Python	Python	LuaJIT	Python
	Net definition language	Python	Python	LuaJIT	Protocol Buffers
	Define-by-Run scheme	Y			
	CPU Array backend	NumPy	NumPy	Tensor	
	GPU Array backend	CuPy	CudaNdarray <sup>1</sup>	CudaTensor	
NNs	Reverse-mode AD	Y	Y	Y	Y
	Basic RNN support	Y	Y	Y(nnx)	#2033
	Variable-length loops	Y	Y(scan)		
	Stateful RNNs <sup>2</sup>	Y	Y	Y <sup>6</sup>	
	Per-batch architectures	Y			
Perf	CUDA support	Y	Y	Y	Y
	cuDNN support	Y	Y	Y(cudnn.torch)	Y
	FFT-based convolution		Y	Y(fbcunn)	#544
	CPU/GPU generic coding <sup>3</sup>	Y	4	Y	
	Multi GPU (data parallel)	Y	Y <sup>7</sup>	Y(fbcunn)	Y
	Multi GPU (model parallel)	Y	Y <sup>8</sup>	Y(fbcunn)	
Misc	Type checking	Y	Y	Y	N/A
	Model serialization	Y	Y(pickle)	Y	Y
	Caffe reference model	Y	5	Y(loadcaffe)	Y

### **Benchmarks**

We are preparing for the benchmarks.

<sup>&</sup>lt;sup>1</sup> They are also developing libgpuarray

<sup>&</sup>lt;sup>2</sup> Stateful RNN is a type of RNN implementation that maintains states in the loops. It should enable us to use the states arbitrarily to update them. <sup>6</sup> Also available in the Torch RNN package

<sup>&</sup>lt;sup>3</sup> This row shows whether each array API supports unified codes for CPU and GPU.

<sup>&</sup>lt;sup>4</sup> The array backend of Theano does not have compatible interface with NumPy, though most users write code on Theano variables, which is generic for CPU and GPU.

<sup>&</sup>lt;sup>7</sup> Via Platoon

<sup>&</sup>lt;sup>8</sup> Experimental as May 2016

<sup>&</sup>lt;sup>5</sup> Depending on the frameworks.

## CHAPTER 9

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### CuPy

The CuPy is designed based on NumPy's API. CuPy's source code and documents contain the original NumPy ones.

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