SdePy Package Documentation

Release 1.0.0

MC

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Part I Getting Started

CHAPTER 1

SdePy

The SdePy package provides tools to state and numerically integrate Ito Stochastic Differential Equations (SDEs), including equations with time-dependent parameters, time-dependent correlations, and stochastic jumps, and to compute with, and extract statistics from, their realized paths.

Several preset processes are provided, including lognormal, Ornstein-Uhlenbeck, Hull-White n-factor, Heston, and jump-diffusion processes.

Computations are fully vectorized across paths, via NumPy and SciPy, making live sessions with 100000 paths reasonably fluent on single cpu hardware.

This package came out of practical need, so expect a flexible tool that gets real-life things done. On the other hand, not every part of it is clean and polished, so expect rough edges, and the occasional bug (please report!).

Developers are committed to the stability of the public API, here again out of practical need to safeguard dependencies.

1.1 Start here

- Installation: pip install sdepy
- Quick Guide (as code)
- Documentation (as pdf)
- Source
- License
- Bug Reports

1.2 License

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File: sdepy/doc/_templates/autosummary/class.rst License: 3-clause BSD

For details, see sdepy/doc/_templates/autosummary/LICENSE.txt

4 Chapter 1. SdePy

CHAPTER 2

Quick Guide

2.1 Install and import

Install using pip install sdepy, or copy the package source code in a directory in your Python path. Import as

```
>>> import sdepy
>>> from sdepy import * # safe and handy for interactive sessions
>>> import numpy as np
>>> import scipy
>>> import matplotlib.pyplot as plt # optional, if plots are needed
```

2.2 How to state an SDE

Here follows a bare-bone definition of a Stochastic Differential Equation (SDE), in this case a Ornstein-Uhlenbeck process:

```
>>> @integrate
... def my_process(t, x, theta=1., k=1., sigma=1.):
... return {'dt': k*(theta - x), 'dw': sigma}
```

This represents the SDE dX = k* (theta -X) *dt + sigma*dW(t), where theta, k and sigma are parameters and dW(t) are Wiener process increments. A further 'dn' or 'dj' entry in the returned dictionary would allow for Poisson or compound Poisson jumps.

A number of preset processes are provided, including lognormal processes, Hull-White n-factor processes, Heston processes, and jump-diffusion processes.

2.3 How to integrate an SDE

Now my_process is a class, a subclass of the cooperating SDE and integrator classes:

```
>>> issubclass(my_process, integrator), issubclass(my_process, SDE)
(True, True)
```

It is to be instantiated with a number of parameters, including the SDE parameters theta, k and sigma; its instances are callable, given a timeline they will integrate and return the process along it. Decorating my_process with kunfc allows for more concise handling of parameters:

```
>>> myp = kfunc(my_process)
>>> iskfunc(myp)
True
```

It is best explained by examples:

1. **Scalar process** in 100000 paths, with default parameters, computed at 5 time points, using 100 steps in between:

2. **Vector process** with three components and **correlated Wiener increments** (same parameters, paths, timeline and steps as above):

```
>>> corr = ((1, .2, -.3), (.2, 1, .1), (-.3, .1, 1))
>>> x = my_process(x0=1, vshape=3, corr=corr,
... paths=100*1000, steps=100)(coarse_timeline)
>>> x.shape
(5, 3, 100000)
```

3. Vector process with **time-dependent parameters and correlations**, computed on a fine-grained timeline and 10000 paths, using one integration step for each point in the timeline (no steps parameter):

```
>>> timeline = np.linspace(0., 1., 101)
>>> corr = lambda t: ((1, .2, -.1*t), (.2, 1, .1), (-.1*t, .1, 1))
>>> theta, k, sigma = (lambda t: 2-t, lambda t: 2/(t+1), lambda t: np.sin(t/2))
>>> x = my_process(x0=1, vshape=3, corr=corr,
... theta=theta, k=k, sigma=sigma, paths=10*1000)(timeline)
>>> x.shape
(101, 3, 10000)
>>> gr = plt.plot(timeline, x[:, 0, :4]) # inspect a few paths
>>> plt.show(gr)
```

4. A scalar process with **path-dependent initial conditions and parameters**, integrated **backwards** (i0=-1):

```
>>> x0 = np.random.random(10*1000)
>>> sigma = 1 + np.random.random(10*1000)
>>> x = my_process(x0=x0, sigma=sigma, paths=10*1000,
... i0=-1)(timeline)
>>> x.shape
(101, 10000)
>>> (x[-1, :] == x0).all()
True
```

5. A scalar process computed on a **10 x 15 grid of parameters** sigma and k (note that the shape of the initial conditions and of each parameter should be broadcastable to the values of the process across paths, i.e. to shape vshape + (paths,)):

```
>>> sigma = np.linspace(0., 1., 10).reshape(10, 1, 1)
>>> k = np.linspace(1., 2., 15).reshape(1, 15, 1)
>>> x = my_process(x0=1, theta=2, k=k, sigma=sigma, vshape=(10, 15),
... paths=10*1000)(coarse_timeline)
```

(continues on next page)

```
>>> x.shape
(5, 10, 15, 10000)
>>> gr = plt.plot(coarse_timeline, x[:, 5, ::2, :].mean(axis=-1))
>>> plt.show()
```

In the example above, set steps=100 to go from inaccurate and fast, to meaningful and slow (the plot illustrates the k-dependence of average process values).

6. Processes generated using **integration results as stochasticity sources** (mind using consistent vshape and paths, and synchronizing timelines):

```
>>> my_dw = integrate(lambda t, x: {'dw': 1}) (vshape=1, paths=10000) (timeline)
>>> p = myp(dw=my_dw, vshape=3, paths=10000,
... x0=1, sigma=((1,), (2,), (3,))) # using myp = kfunc(my_process)
>>> x = p(timeline)
>>> x.shape
(101, 3, 10000)
```

Now, x1, x2, x3 = x[:, 0], x[:, 1], x[:, 2] have different sigma, but share the same dw increments, as can be seen plotting a path:

```
>>> k = 0 # path to be plotted
>>> gr = plt.plot(timeline, x[:, :, k])
>>> plt.show()
```

If more integrations steps are needed between points in the output timeline, use steps to keep the integration timeline consistent with the one of my_dw:

```
>>> x = p(coarse_timeline, steps=timeline)
>>> x.shape
(5, 3, 10000)
```

7. Using stochasticity sources with memory (mind using consistent vshape and paths):

```
>>> my_dw = true_wiener_source(paths=10000)
>>> p = myp(x0=1, k=1, sigma=1, dw=my_dw, paths=10000)

>>> t1 = np.linspace(0., 1., 30)
>>> t2 = np.linspace(0., 1., 100)
>>> t3 = t = np.linspace(0., 1., 300)
>>> x1, x2, x3 = p(t1), p(t2), p(t3)
>>> y1, y2, y3 = p(t, theta=1.5), p(t, theta=1.75), p(t, theta=2)
```

These processes share the same underlying Wiener increments: x1, x2, x3 illustrate SDE integration convergence as steps become smaller, and y1, y2, y3 illustrate how k affects paths, all else being equal:

```
>>> i = 0 # path to be plotted

>>> gr = plt.plot(t, x1(t)[:, i], t, x2(t)[:, i], t, x3(t)[:, i])

>>> gr = plt.plot(t, y1[:, i], t3, y2[:, i], t3, y3[:, i])

>>> plt.show()
```

2.4 How to handle the integration output

SDE integrators return process instances, a subclass of np.ndarray with a timeline stored in the t attribute (note the shape of x, repeatedly used in the examples below):

```
>>> coarse_timeline = (0., 0.25, 0.5, 0.75, 1.0)
>>> timeline = np.linspace(0., 1., 101)

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```

continues on next page)

```
>>> x = my_process(x0=1, vshape=3, paths=1000)(timeline)
>>> x.shape
(101, 3, 1000)
```

x is a process instance, based on the given timeline:

```
>>> type(x)
<class 'sdepy.infrastructure.process'>
>>> np.isclose(timeline, x.t).all()
True
```

Whenever possible, a process will store references, not copies, of timeline and values. In fact,

```
>>> timeline is x.t
True
```

The first axis is reserved for the timeline, the last for paths, and axes in the middle match the shape of process values:

```
>>> x.shape == x.t.shape + x.vshape + (x.paths,)
True
```

Calling processes interpolates in time (the result is an array, not a process):

```
>>> y = x(coarse_timeline)
>>> y.shape
(5, 3, 1000)
>>> type(y)
<class 'numpy.ndarray'>
```

All array methods, including indexing, work as usual (no overriding), and return NumPy arrays:

```
>>> type(x[0])
<class 'numpy.ndarray'>
>>> type(x.mean(axis=0))
<class 'numpy.ndarray'>
```

You can slice processes along time, values and paths with special indexing:

```
>>> y = x['t', ::2] # time indexing
>>> y.shape
(51, 3, 1000)
>>> y = x['v', 0] # values indexing
>>> y.shape
(101, 1000)
>>> y = x['p', :10] # paths indexing
>>> y.shape
(101, 3, 10)
```

The output of a special indexing operation is a process:

```
>>> isinstance(y, process)
True
```

Smart indexing is allowed. To select paths that cross x=0 at some point and for some component, use:

```
>>> i_negative = x.min(axis=(0, 1)) < 0
>>> y = x['p', i_negative]
```

(continues on next page)

```
>>> y.shape == (101, 3, i_negative.sum())
True
```

You can do algebra with processes that either share the same timeline, or are constant (a process with a one-point timeline is assumed to be constant), and either have the same number of paths, or are deterministic (with one path):

```
>>> x_const = x['t', 0] # a constant process
>>> x_one_path = x['p', 0] # a process with one path
>>> y = np.exp(x) - x_const
>>> z = np.maximum(x, x_one_path)
>>> isinstance(y, process), isinstance(z, process)
(True, True)
```

When integrating SDEs, the SDE parameters and/or stochasticity sources accept processes as valid values (mind using deterministic processes, or synchronizing the number of paths, and make sure that the shape of values do broadcast together). To use a realization of my_process as the volatility of a 3-component lognormal process, do as follows:

```
>>> stochastic_vol = my_process(x0=1, paths=10*1000)(timeline)
>>> stochastic_vol_x = lognorm_process(x0=1, vshape=3, paths=10*1000,
... mu=0, sigma=stochastic_vol)(timeline)
```

Processes have specialized methods, and may be analyzed, and their statistics cumulated across multiple runs, using the montecarlo class. Some examples follow:

1. Cumulative probability distribution function at t=0.5 of the process values of x across paths:

```
>>> cdf = x.cdf(0.5, x=np.linspace(-2, 2, 100)) # an array
```

2. Characteristic function at t=0.5 of the same distribution:

```
>>> chf = x.chf(0.5, u=np.linspace(-2, 2, 100)) # an array
```

3. Standard deviation across paths:

```
>>> std = x.pstd() # a one-path process
>>> std.shape
(101, 3, 1)
```

4. Maximum value reached along the timeline:

```
>>> xmax = x.tmax() # a constant process
>>> xmax.shape
(1, 3, 1000)
```

5. A linearly interpolated, or Gaussian kernel estimate (default) of the probability distribution function (pdf) and its cumulated values (cdf) across paths, at a given time point, may be obtained using the montecarlo class:

```
>>> y = x(1)[0] # 0-th component of x at time t=1
>>> a = montecarlo(y, bins=30)
>>> ygrid = np.linspace(y.min(), y.max(), 200)
>>> gr = plt.plot(ygrid, a.pdf(ygrid), ygrid, a.cdf(ygrid))
>>> gr = plt.plot(ygrid, a.pdf(ygrid, method='interp', kind='nearest'))
>>> plt.show()
```

6. A montecarlo instance can be used to cumulate the results of multiple simulations, across multiple components of process values:

2.5 Example - Stochastic Runge-Kutta

Minimal implementation of a basic stochastic Runge-Kutta integration, scheme, as a subclass of integrator (the A and dZ methods below are the standardized way in which equations are exposed to integrators):

SDE of a lognormal process, as a subclass of SDE, and classes that integrate it with the default integration method (p1) and via my_integrator (p2):

```
>>> class my_SDE(SDE):
... def sde(self, t, x): return {'dt': 0, 'dw': x}
>>> class p1(my_SDE, integrator): pass
>>> class p2(my_SDE, my_integrator): pass
```

Comparison of integration errors, as the integration from t=0 to t=1 is carried out with an increasing number of steps:

```
>>> np.random.seed(1)
>>> args = dict(dw=true_wiener_source(paths=100), paths=100, x0=10)
>>> timeline = (0, 1)
>>> steps = np.array((2, 3, 5, 10, 20, 30, 50, 100,
                       200, 300, 500, 1000, 2000, 3000))
>>> exact = lognorm_process(mu=0, sigma=1, **args)(timeline)[-1].mean()
>>> errors = np.abs(np.array([
        [p1(**args, steps=s)(timeline)[-1].mean()/exact - 1,
. . .
         p2(**args, steps=s)(timeline)[-1].mean()/exact - 1]
       for s in steps]))
>>> ax = plt.axes(label=0); ax.set_xscale('log'); ax.set_yscale('log')
>>> gr = ax.plot(steps, errors)
>>> plt.show()
>>> print('euler error: \{:.2e\}\n rk error: \{:.2e\}'.format(errors[-1,0], errors[-
\hookrightarrow 1, 1]))
euler error: 1.70e-03
  rk error: 8.80e-06
```

2.6 Example - Fokker-Planck Equation

Monte Carlo integration of partial differential equations, illustrated in the simplest example of the heat equation diff(u, t) - k*diff(u, x, 2) == 0, for the function u(x, t), i.e. the Fokker-Planck equation for the SDE dX(t) = sqrt(2*k)*dW(t). Initial conditions at t=t0, two examples:

```
1. u(x, t0) = 1 for lb < x < hb and 0 otherwise,
2. u(x, t0) = sin(x).
```

Setup:

```
>>> from numpy import exp, sin
>>> from scipy.special import erf
>>> from scipy.integrate import quad
>>> np.random.seed(1)
>>> k = .5
>>> x0, x1 = 0, 10;
>>> t0, t1 = 0, 1
>>> 1b, hb = 4, 6
```

Exact green function and solutions, to be checked against results:

```
>>> def green_exact(y, s, x, t):
       return \exp(-(x - y)**2/(4*k*(t - s)))/sqrt(4*np.pi*k*(t - s))
>>> def u1_exact(x, t):
        return (erf((x - 1b)/2/sqrt(k*(t - t0))) - erf((x - hb)/2/sqrt(k*(t - u))))
\hookrightarrowt0))))/2
>>> def u2_exact(x, t):
        return \exp(-k*(t-t0))*\sin(x)
```

Realization of the needed stochastic process, by backward integration from a grid of final values of x at t=t1, using the preset wiener_process class (the steps keyword is added as a reminder of the setup needed for less-than-trivial equations, it does not actually make a difference here):

```
>>> xgrid = np.linspace(x0, x1, 51)
>>> tgrid = np.linspace(t0, t1, 5)
>>> xp = wiener_process(paths=10000,
                sigma=sqrt(2*k), steps=100,
. . .
                vshape=xgrid.shape, x0=xgrid[..., np.newaxis],
. . .
                i0=-1) (timeline=tgrid)
```

Computation of the green function and of the solution u(x, t1) (note the liberal use of scipy.integrate. quad below, enabled by the smoothness of the Gaussian kernel estimate a [i, j].pdf):

```
>>> a = montecarlo(xp, bins=100)
>>> def green(y, i, j):
       """green function from (y=y, s=tgrid[i]) to (x=xgrid[j], t=t1)"""
       return a[i, j].pdf(y)
>>> u1, u2 = np.empty(51), np.empty(51)
>>> for j in range(51):
       u1[j] = quad(lambda y: green(y, 0, j), lb, hb)[0]
       u2[j] = quad(lambda y: sin(y)*green(y, 0, j), -np.inf, np.inf)[0]
```

Comparison against exact values:

```
\rightarrow \rightarrow y = np.linspace(x0, x1, 500)
>>> for i, j in ((1, 20), (2, 30), (3, 40)):
        gr = plt.plot(y, green(y, i, j),
. . .
                        y, green_exact(y, tgrid[i], xgrid[j], t1), ':')
. . .
>>> plt.show()
>>> gr = plt.plot(xgrid, u1, y, u1_exact(y, t1), ':')
```

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2.7 Example - Basket Lookback Option

Take a basket of 4 financial securities, with risk-neutral probabilities following lognormal processes in the Black-Sholes framework. Correlations, dividend yields and term structure of volatility (will be linearly interpolated) are given below:

```
>>> corr = [
    [1,
               0.50, 0.37, 0.35],
. . .
        [0.50, 1, 0.47, 0.46],
. . .
        [0.37, 0.47, 1, 0.19],
. . .
                     0.19, 1]]
        [0.35, 0.46,
. . .
>>> dividend_yield = process(c=(0.20, 4.40, 0., 4.80))/100
>>> riskfree = 0 # to keep it simple
>>> vol_timepoints = (0.1, 0.2, 0.5, 1, 2, 3)
>>> vol = np.array([
       [0.40, 0.38, 0.30, 0.28, 0.27, 0.27],
. . .
       [0.31, 0.29, 0.22, 0.16, 0.18, 0.21],
. . .
       [0.24, 0.22, 0.19, 0.19, 0.21, 0.22],
      [0.35, 0.31, 0.21, 0.18, 0.19, 0.19]])
>>> sigma = process(t=vol_timepoints, v=vol.T)
>>> sigma.shape
(6, 4, 1)
```

The prices of the securities at the end of each quarter for the next 2 years, simulated across 50000 independent paths and their antithetics (odd_wiener_source is used), are:

A call option knocks in if any of the securities reaches a price below 80 at any quarter (starting from 100), and pays the lookback maximum attained by the basket (equally weighted), minus 105, if positive. Its price is:

```
>>> x_worst = x.min(axis=1)
>>> x_mean = x.mean(axis=1)
>>> down_and_in_paths = (x_worst.min(axis=0) < 80)
>>> lookback_x_mean = x_mean.max(axis=0)
>>> payoff = np.maximum(0, lookback_x_mean - 105)
>>> payoff[np.logical_not(down_and_in_paths)] = 0
>>> a = montecarlo(payoff, use='even')
>>> print(a)
4.997 +/- 0.027
```

Part II **API Documentation**

CHAPTER 3

Overview

This package provides tools to state and numerically integrate Ito Stochastic Differential Equations (SDEs), including equations with time-dependent parameters, time-dependent correlations, and stochastic jumps, and to compute with, and extract statistics from, their realized paths.

Package contents:

- 1. A set of tools to ease computations with stochastic processes, as obtained from numerical integration of the corresponding SDE, is provided via the process and montecarlo classes (see *Infrastructure*):
 - The process class, a subclass of numpy.ndarray representing a sequence of values in time, realized in one or several paths. Algebraic manipulations and ufunc computations are supported for instances that share the same timeline, or are constant, and comply with numpy broadcasting rules. Interpolation along the timeline is supported via callability of process instances. Process-specific functionalities, such as averaging and indexing along time or across paths, are delegated to process-specific methods, attributes and properties (no overriding of numpy.ndarray operations).
 - The montecarlo class, as an aid to cumulate the results of several Monte Carlo simulations of a given stochastic variable, and to extract summary estimates for its probability distribution function and statistics.
- 2. Numerical realizations of the differentials commonly found as stochasticity sources in SDEs, are provided via the source class and its subclasses, with or without memory of formerly invoked realizations (see *Stochasticity Sources*).
- 3. A general framework for stochastic step by step simulations, and for numerical SDE integration, is provided via the paths_generator class, and its cooperating subclasses integrator, SDE and SDEs (see SDE Integration Framework). The full API allows for extensive customization of preprocessing, post-processing, stochasticity sources instantiation and handling, integration algorithms etc. The integrate decorator provides a simple and concise interface to handle standard use cases, via Euler-Maruyama integration.
- 4. Several preset stochastic processes are provided, including lognormal, Ornstein-Uhlenbeck, Hull-White n-factor, Heston, and jump-diffusion processes (see *Stochastic Processes*). Each process consists of a process generator class, a subclass of integrator and SDE, named with a _process suffix, and a definition of the underlying SDE, a subclass of SDE or SDEs, named with a _SDE suffix.
- 5. Several analytical results relating to the preset stochastic processes are made available, as a general reference and for testing purposes (see *Analytical Results*). They are limited to the case of constant process parameters, and with some further limitations on the parameters' domains. Function arguments are consistent with those of the corresponding processes. Suffixes _pdf, _cdf and _chf stand respectively for

- probability distribution function, cumulative probability distribution function, and characteristic function. Black-Scholes formulae for the valuation of call and put options have been included (with prefix bs).
- 6. As an aid to interactive and notebook sessions, shortcuts are provided for stochasticity sources and preset processes (see *Shortcuts*). Shortcuts have been wrapped as "kfuncs", objects with managed keyword arguments that simplify interactive workflow when frequent parameters tuning operations are needed (see kfunc decorator documentation). Analytical results are wrapped as kfuncs as well.

For all sources and processes, values can take any shape, scalar or multidimensional. Correlated multivariate stochasticity sources are supported. Poisson jumps are supported, and may be compounded with any random variable supported by scipy.stats. Time-varying process parameters (correlations, intensity of Poisson processes, volatilities etc.) are allowed whenever applicable. process instances act as valid stochasticity source realizations (as does any callable object complying with a source protocol), and may be passed as a source specification when computing the realization of a given process.

Computations are fully vectorized across paths, providing an efficient infrastructure for simulating a large number of process realizations. Less so, for large number of time steps: integrating 100 time steps across one million paths takes seconds, one million time steps across 100 paths takes minutes.

Infrastructure

process([t, x, v, c, dtype])	Array representation of a process (a subclass of
	numpy.ndarray).
<pre>montecarlo([sample, axis, bins, range, use,])</pre>	Summary statistics of Monte Carlo simulations.

4.1 sdepy.process

class sdepy.**process** (t=0., *, x=None, v=None, c=None, dtype=None) Array representation of a process (a subclass of numpy.ndarray).

If p is a process instance, p[i, ..., k] is the value that the k-th path of the represented process takes at time p.t[i]. The first and last indexes of p are reserved for the timeline and paths respectively. A process should contain no less than 1 time point and 1 path. Zero or more middle indexes refer to the values that the process takes at each given time and path.

If p has N time points, paths is its number of paths and vshape is the shape of its values at any given time point and path, then p. shape is (N_*) + vshape + (paths,). N, vshape, paths are inferred at instantiation from the shape of t and x, v or c parameters.

Parameters

- **t** [array-like] Timeline of the process, as a one dimensional array with shape (N,), in increasing order. Defaults to 0.
- x [array-like, optional] Values of the process along the timeline and across paths. Should broadcast to (N,) + vshape + (paths,). The shapes of t and of the firs index of x must match. One and only one of x, v, c must be provided upon process creation, as a keyword argument.
- v [array-like, optional] Values of a deterministic process along the timeline. Should broadcast to (N,) + vshape. The shapes of t and of the firs index of v must match.
- **c** [array-like, optional] Value of a constant, single-path process, with shape vshape. Each time point of the resulting process contains a copy of c.

dtype [data-type, optional] Data-type of the values of the process. x, v or c will be converted to dtype if need be.

Notes

A reference and not a copy of t, x, v, c is stored if possible.

A process is a subclass of numpy.ndarray, where its values as an array are the process values along the timeline and across paths. All numpy.ndarray methods, attributes and properties are guaranteed to act upon such values, as would those of the parent class. Such no-overriding commitment is intended to safeguard predictability of array operations on process instances; process-specific functionalities are delegated to process-specific methods, attributes and properties.

A process with a single time point is assumed to be constant.

Processes have the __array_priority__ attribute set to 1.0 by default. Ufuncs acting on a process, or on a process and an array, or on different processes sharing the same timeline, or on different processes one of which is constant, return a process with the timeline of the original process(es) passed as a reference. Ufuncs calls on different processes fail if non constant processes do not share the same timeline (interpolation should be handled explicitly), or in case broadcasting rules would result in mixing time, values and/or paths axes.

Let p be a process instance. Standard numpy indexing acts on the process values and returns numpy.ndarray instances: in fact, p[i] is equivalent to p.x[i], i.e. the same as p.view(numpy.ndarray)[i]. Process-specific indexing is addressed via the following syntax, where i can be an integer, a multi-index or smart indexing reference consistent with the process shape:

- p['t', i]: timeline indexing, roughly equivalent to process (t=p.t[i], x=p.x[i, ..., :])
- p['v', i]: values indexing, roughly equivalent to process (t=p.t, x=p.x[:, i, :])
- p['p', i]: paths indexing, roughly equivalent to process(t=p.t, x=p.x[:, ..., i])

Attributes

- * Process values, viewed as a numpy.ndarray.
- **paths** Number of paths of the process (coincides with the size of the last dimension of the process).
- **vshape** Shape of the values of the process.
- **tx** Timeline of the process, reshaped to be broadcastable to the process values and paths across time.
- dt Process timeline increments, as returned by numpy.diff.
- **dtx** Process timeline increments, as returned by numpy.diff, reshaped to be broadcastable to the process values.
- t [array] Stores the timeline of the process.
- interp_kind [str] Stores the default interpolation kind, passed upon interpolation (interp
 and __call__ methods) to scipy.interpolate.interpld unless a specific kind is provided. Defaults to 'linear'.

Methods

<pre>interp(*[, kind])</pre>	Interpolation in time of the process values.
call(s[, ds, kind])	Interpolation in time of process values or incre-
	ments.
getitem(key)	See documentation of the process class.

Continued on next page

Table 2 – continued from previous page

rebase(t, *[, kind])	Change the process timeline to t, using interpola-
repase(t, [, kmd])	tion.
shapeas(vshape_or_process)	Reshape process values according to the given tar-
Shapeas(vshape_or_process)	get shape.
pcopy(**args)	Copy timeline and values of the process (args are
peopy (mgs)	passed to numpy.ndarray.copy).
xcopy(**args)	Copy values of the process, share timeline (args
xcopy(digs)	are passed to numpy.ndarray.copy).
(**omgo)	
tcopy(**args)	Copy timeline of the process, share values.
<pre>pmin([out])</pre>	One path process exposing for each time point the minimum process value attained across paths.
pmax([out])	One path process exposing for each time point the
	maximum process value attained across paths.
psum([dtype, out])	One path process exposing for each time point the
	sum of process values across paths.
<pre>pmean([dtype, out])</pre>	One path process exposing for each time point the
	mean of process values across paths.
<pre>pvar([dtype, out, ddof])</pre>	One path process exposing for each time point the
[variance of process values across paths.
pstd([dtype, out, ddof])	One path process exposing for each time point the
ps ea([atype, out, ador])	standard deviation of process values across paths.
tmin([out])	Constant process exposing for each path the mini-
emin([out])	mum process value attained along time.
tmax([out])	Constant process exposing for each path the maxi-
chax([out])	mum process value attained along time.
tsum([dtype, out])	Constant process exposing for each path the sum
esum([atype, out])	of process values along time.
tmean([dtype, out])	Constant process exposing for each path the mean
tmean([diype, out])	
(dt dd. fl)	of process values along time.
tvar([dtype, out, ddof])	Constant process exposing for each path the vari-
11.60	ance of process values along time.
tstd([dtype, out, ddof])	Constant process exposing for each path the stan-
45.4	dard deviation of process values along time.
tdiff([dt_exp, fwd])	Process increments along the timeline, optionally
	weighted by time increments.
tder()	Forward looking derivative of the given process,
	linearly interpolated between time points.
tint()	Integral of the given process, linearly interpolated
	between time points.
chf([t, u])	Characteristic function of the probability distribu-
	tion of process values.
cdf([t, x])	Cumulative probability distribution function of
	process values.
	<u> </u>

4.1.1 sdepy.process.interp

process.interp(*, kind=None)

Interpolation in time of the process values.

Returns a callable f, as returned by scipy.interpolate.interpld, such that f(s) approximates the value of the process at time point s. f refers to the process timeline and values, without storing copies. s may be of any shape.

Parameters

kind [string] An interpolation kind as accepted by scipy.interpolate.
interpld. If None, defaults to the interp_kind class attribute.

4.1. sdepy.process

Returns

f [callable] f, as returned by scipy.interpolate.interp1d, such that f(s) approximates the value of the process at time point s. f refers to the process timeline and values, without storing copies.

```
s may be of any shape: if p is a process instance, p.interp()(s).shape ==
s.shape + p.vshape + (p.paths,).
```

In case p has a single time point, interpolation is not handled via scipy. interpolate.interpld; the process is assumed to be constant in time, and f is a function object behaving accordingly.

See also:

```
process.__call__
```

Notes

The process is extrapolated as constant outside the timeline boundaries.

If p is a process instance, p.interp(s) is an array, not a process. If an interpolated process is needed, it should be explicitly created using q = process(s, x=p(s)), or its shorthand q = p.rebase(s).

4.1.2 sdepy.process. call

```
process.__call__(s, ds=None, *, kind=None)
```

Interpolation in time of process values or increments.

If p is a process instance and f = p.interp(kind):

- p(s) returns f(s),
- p(s, ds) returns f(s + ds) f(s).

See also:

```
process.interp
```

4.1.3 sdepy.process.rebase

```
process.rebase(t, *, kind=None)
```

Change the process timeline to t, using interpolation.

A new process is returned with timeline t and values set to the calling process values, interpolated at t using process.interp with the given interpolation kind.

If t is a scalar, a constant process is returned.

4.1.4 sdepy.process.shapeas

```
process.shapeas (vshape_or_process)
```

Reshape process values according to the given target shape.

Returns a process pointing to the same data as the calling process, adding new 1-dimensional axes, or removing existing 1-dimensional axes to the left of the first dimension of process values, as needed to make the returned process broadcastable to a process with values of the given shape.

To achieve broadcastability the unaffected dimensions, including the shape of the timeline and the number of paths, have to be compatible.

Raises

ValueError [if requested to remove a non 1-dimensional axis]

4.1.5 sdepy.process.pcopy

```
process.pcopy(**args)
```

Copy timeline and values of the process (args are passed to numpy.ndarray.copy).

4.1.6 sdepy.process.xcopy

```
process.xcopy(**args)
```

Copy values of the process, share timeline (args are passed to numpy.ndarray.copy).

4.1.7 sdepy.process.tcopy

```
process.tcopy(**args)
```

Copy timeline of the process, share values. (args are passed to numpy.ndarray.copy).

4.1.8 sdepy.process.pmin

```
process.pmin(out=None)
```

One path process exposing for each time point the minimum process value attained across paths.

4.1.9 sdepy.process.pmax

```
process.pmax(out=None)
```

One path process exposing for each time point the maximum process value attained across paths.

4.1.10 sdepy.process.psum

```
process.psum(dtype=None, out=None)
```

One path process exposing for each time point the sum of process values across paths.

4.1.11 sdepy.process.pmean

```
process.pmean (dtype=None, out=None)
```

One path process exposing for each time point the mean of process values across paths.

4.1.12 sdepy.process.pvar

```
process.pvar(dtype=None, out=None, ddof=0)
```

One path process exposing for each time point the variance of process values across paths.

4.1.13 sdepy.process.pstd

```
process.pstd(dtype=None, out=None, ddof=0)
```

One path process exposing for each time point the standard deviation of process values across paths.

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4.1.14 sdepy.process.tmin

```
process.tmin(out=None)
```

Constant process exposing for each path the minimum process value attained along time.

4.1.15 sdepy.process.tmax

```
process.tmax(out=None)
```

Constant process exposing for each path the maximum process value attained along time.

4.1.16 sdepy.process.tsum

```
process.tsum(dtype=None, out=None)
```

Constant process exposing for each path the sum of process values along time.

4.1.17 sdepy.process.tmean

```
process.tmean (dtype=None, out=None)
```

Constant process exposing for each path the mean of process values along time.

4.1.18 sdepy.process.tvar

```
process.tvar(dtype=None, out=None, ddof=0)
```

Constant process exposing for each path the variance of process values along time.

4.1.19 sdepy.process.tstd

```
process.tstd(dtype=None, out=None, ddof=0)
```

Constant process exposing for each path the standard deviation of process values along time.

4.1.20 sdepy.process.tdiff

```
process.tdiff(dt_exp=0, fwd=True)
```

Process increments along the timeline, optionally weighted by time increments.

Parameters

dt_exp [int or float, optional] Exponent applied to time increment weights. If 0, returns process increments. If 1, approximates a time derivative. If 0.5, approximates realized volatility.

fwd [bool, optional] If True, the differences are forward-looking

Returns

```
q [process] If p is a process shaped (N,) + p.vshape + (p.paths,), with
timelinet,p.tdiff(dt_exp, fwd) returns a process q, shaped (N-1,) + p.
vshape + (p.paths,) with values
```

```
q[i] = (p[i+1] - p[i])/(t[i+1] - t[i])**dt_exp
```

If fwd evaluates to True, q[i] is assigned to time point t[i] (q stores at t[i] the increments of p looking forwards) or to t[i+1] otherwise (increments looking backwards).

See also:

```
tder, tint
```

Notes

if p is a process instance realizing a solution of the SDE dp(t) = sigma(t) *dw(t) across several paths, then

```
p.tdiff(dt_exp=0.5).pstd()
```

is a 1-path process that estimates sigma(t).

4.1.21 sdepy.process.tder

```
process.tder()
```

Forward looking derivative of the given process, linearly interpolated between time points.

```
Shorthand for p.tdiff(dt_exp=1).
```

See also:

```
tdiff, tint
```

Notes

```
p.tder().tint() equals, within rounding errors, p['t', :-1] - p['t', 0]
```

4.1.22 sdepy.process.tint

```
process.tint()
```

Integral of the given process, linearly interpolated between time points.

See also:

```
tdiff, tder
```

Notes

```
p.tin().tder() equals, within rounding errors, p['t', :-1]
```

4.1.23 sdepy.process.chf

```
process.chf (t=None, u=None)
```

Characteristic function of the probability distribution of process values.

p.chf(t, u) estimates the characteristic function of interpolated process values p(t) at time(s) t. p.chf(u) is a shorthand for p.chf(p.t, u) (no interpolation).

Parameters

- ${f t}$ [array-like, optional] Time points at which to compute the characteristic function. If omitted or None, the entire process timeline is used.
- **u** [array-like, mandatory] Values at which to evaluate the characteristic function.

Returns

```
array Returns an array, with shape t.shape + u.shape + vshape, where vshape is the shape of values of the calling process p, containing the average across paths of exp(1j*u*p(t)).
```

4.1.24 sdepy.process.cdf

```
process.cdf (t=None, x=None)
```

Cumulative probability distribution function of process values.

p.cdf(t, x) estimates the cumulative probability distribution function of interpolated process values p(t) at time(s) t. p.cdf(x) is a shorthand for p.cdf(p.t, x) (no interpolation).

Parameters

- **t** [array-like, optional] Time points along the process timeline. If omitted or None, the entire process timeline is used.
- x [array-like, mandatory] Values at which to evaluate the cumulative probability distribution function.

Returns

array Returns an array, with shape t.shape + x.shape + vshape, where vshape is the shape of the values of the calling process p, containing the average across paths of 1 if p(t) <= x else 0.

4.2 sdepy.montecarlo

Summary statistics of Monte Carlo simulations.

Compute, store and cumulate results of Monte Carlo simulations across multiple runs. Cumulated results include mean, standard deviation, standard error, skewness, kurtosis, and 1d-histograms of the distribution of outcomes. Probability distribution function estimates are provided, based on the cumulated histograms.

Parameters

- **sample** [array-like, optional] Initial data set to be summarized. If None, an empty instance is provided, initialized with the given parameters.
- **axis** [integer, optional] Axis of the given sample enumerating single data points (paths, or different realizations of a simulated process or event). Defaults to the last axis of the sample.
- use [{'all', 'even', 'odd'}, optional] If 'all' (default), the data set is processed as is. If 'even' or 'odd', the sample x is assumed to consist of antithetic values along the specified axis, assumed of even size 2*N, where x[0], x[1], ... is antithetic respectively to x[N], x[N+1], ... Summary operations are then applied to a sample of size N consisting of the half-sum ('even') or half-difference ('odd') of antithetic values.
- **bins** [array-like, or int, or str, optional] Bins used to evaluate the counts' cumulated distribution are computed, against the first data set encountered, according to the bins parameter:
 - If int or str, it dictates the number of bins or their determination method, as passed to numpy.histogram when processing the first sample.
 - If array-like, overrides range, setting explicit bins' boundaries, so that bins[i][j] is the lower bound of the j-th bin used for the distribution of the i-th component of data points.
 - If None, no distribution data will be computed.

Defaults to 100.

range [(float, float) or None, optional] Bins range specification, as passed to numpy.
histogram.

dtype [data-type, optional] Data type used for cumulating moments. If None, the data-type of the first sample is used, if of float kind, or float otherwise.

ctype [data-type, optional] Data type used for cumulating histogram counts. Defaults to numpy.int64.

Notes

The shape of cumulated statistics is set as the shape of the data points of the first data set processed (shape of the first sample after summarizing along the paths axis). When cumulating subsequent samples, broadcasting rules apply.

Indexing can be used to access single values or slices of the stored data. Given a montecarlo instance a, a [i] is a new instance referencing statistics of the i-th component of data summarized in a (no copying).

The first data set encountered fixes the histogram bins. Points of subsequent data sets that fall outside the bins, while properly taken into account in summary statistics (mean, standard error etc.), are ignored when building cumulated histograms and probability distribution functions. Their number is accounted for in the outpaths property and outerr method.

Histograms and distributions, and the related outpaths and outerr, must be invoked on single-valued montecarlo instances. For multiple valued simulations, use indexing to select the value to be addressed (e.g. a[i].histogram()).

Attributes

paths Number of cumulated sample data points (0 for an empty instance).

vshape Shape of cumulated sample data points.

shape Shape of cumulated sample data set, rearranged with averaging axis as last axis.

outpaths Data points fallen outside of the bins' boundaries.

- m Shortcut for the mean method.
- **s** Shortcut for the std method.
- e Shortcut for the stderr method.

stats Dictionary of cumulated statistics.

- h Shortcut for the histogram method.
- $\begin{tabular}{ll} \textbf{dh} & Shortcut for the {\tt density_histogram} & method. \end{tabular}$

Methods

update(sample[, axis])	Add the given sample to the montecarlo simula-
	tion.
mean()	Mean of cumulated sample data points.
var()	Variance of cumulated sample data points.
std()	Standard deviation of cumulated sample data
	points.
skew()	Skewness of cumulated sample data points.
kurtosis()	Kurtosis of cumulated sample data points.
stderr()	Standard error of the mean of cumulated sample
	data points.
histogram()	Distribution of the cumulated sample data, as a
	counts histogram.
density_histogram()	Distribution of the cumulated sample data, as a
	normalized counts histogram.
•	Continued on post page

Continued on next page

Table 3 – continued from previous page

pdf(x[, method, bandwidth, kind])	Normalized sample probability density function,
	evaluated at x.
cdf(x[, method, bandwidth, kind])	Cumulative sample probability density function,
	evaluated at x.
outerr()	Fraction of cumulated data points fallen outside of
	the bins' boundaries.

4.2.1 sdepy.montecarlo.update

```
montecarlo.update (sample, axis=-1)
```

Add the given sample to the montecarlo simulation.

Combines the given sample data with summary statistics obtained (if any) from former samples to which the montecarlo instance was exposed at instantiation and at previous calls to this method. Updates cumulated statistics and histograms accordingly.

Parameters

sample [array-like] Data set to be summarized.

axis [integer, optional] Axis of the given sample enumerating single data points (paths, or different realizations of a simulated process or event). Defaults to the last axis of the sample.

4.2.2 sdepy.montecarlo.mean

```
montecarlo.mean()
```

Mean of cumulated sample data points.

4.2.3 sdepy.montecarlo.var

```
montecarlo.var()
```

Variance of cumulated sample data points.

4.2.4 sdepy.montecarlo.std

```
montecarlo.std()
```

Standard deviation of cumulated sample data points.

4.2.5 sdepy.montecarlo.skew

```
montecarlo.skew()
```

Skewness of cumulated sample data points.

4.2.6 sdepy.montecarlo.kurtosis

```
montecarlo.kurtosis()
```

Kurtosis of cumulated sample data points.

4.2.7 sdepy.montecarlo.stderr

```
montecarlo.stderr()
    Standard error of the mean of cumulated sample data points.
    a.stderr() equals a.std()/sqrt(a.paths - 1).
```

4.2.8 sdepy.montecarlo.histogram

```
montecarlo.histogram()
```

Distribution of the cumulated sample data, as a counts histogram.

Returns a (counts, bins) tuple of arrays representing the one-dimensional histogram data of the distribution of cumulated samples (as returned by numpy.histogram).

4.2.9 sdepy.montecarlo.density_histogram

```
montecarlo.density histogram()
```

Distribution of the cumulated sample data, as a normalized counts histogram.

Returns a (counts, bins) tuple of arrays representing the one-dimensional density histogram data of the distribution of cumulated samples (as returned by numpy.histogram, the sum of the counts times the bins' widths is 1).

May systematically over-estimate the probability distribution within the bins' boundaries if part of the cumulated samples data (accounted for in the outpaths property and outerr method) fall outside.

4.2.10 sdepy.montecarlo.pdf

```
montecarlo.pdf (x, method='gaussian\_kde', bandwidth=1.0, kind='linear')
Normalized sample probability density function, evaluated at x.
```

Parameters

x [array-like] Values at which to evaluate the pdf.

method [{'gaussian_kde', 'interp'}] Specifies the method used to estimate the pdf value. One of: 'gaussian_kde' (default), smooth Gaussian kernel density estimate of the probability density function; 'interp', interpolation of density histogram values, of the given kind.

bandwidth [float] The bandwidth of Gaussian kernels is set to bandwidth times each bin width.

kind [str] Interpolation kind for the 'interp' method, passed to scipy.
interpolate.intepld.

Returns

array An estimate of the sample probability density function of the cumulated sample data, at the given 'x' values, according to the stated method.

Notes

For the 'gaussian_kde' method, kernels are computed at bins midpoints, weighted according to the density histogram counts, using in each bin a bandwidth set to bandwidth times the bin width. The resulting pdf:

- Has support on the real line.
- Integrates exactly to 1.

• May not closely track the density histogram counts.

For the 'interp' method, the pdf evaluates to the density histogram counts at each bin midpoint, and to 0 at the bins boundaries and outside. The resulting pdf:

- Has support within the bins boundaries.
- Is intended to track the density histogram counts.
- Integrates close to, but not exactly equal to, 1.

May systematically overestimate the probability distribution within the bins' boundaries if part of the cumulated samples data (accounted for in the outpaths property and outerr method) fall above or below the bins boundaries.

4.2.11 sdepy.montecarlo.cdf

```
montecarlo.cdf (x, method='gaussian\_kde', bandwidth=1.0, kind='linear') Cumulative sample probability density function, evaluated at x.
```

See pdf method documentation.

Notes

For the 'gaussian_kde' method, the integral of the Gaussian kernels is expressed in terms of scipy. special.erf, and coincides with the integral of the pdf computed with the same method.

For the 'interp' method, the cdf evaluates as follows:

- At bin endpoints, to the cumulated density histogram values weighed by the bins width.
- Below the bins boundaries, to 0.
- Above the bins boundaries, to 1.

It is close to, but not exactly equal to, the integral of the pdf computed with the same method.

4.2.12 sdepy.montecarlo.outerr

```
montecarlo.outerr()
```

Fraction of cumulated data points fallen outside of the bins' boundaries.

CHAPTER 5

Stochasticity Sources

source(*[, paths, vshape, dtype])	Base class for stochasticity sources.
wiener_source(*[, paths, vshape, dtype,])	dw, a source of standard Wiener process (Brownian
wienei_source('[, paths, vshape, dtype,])	motion) increments.
(*[· · · · · · · · · · · · · · · · · · ·
poisson_source(*[, paths, vshape, dtype, lam])	dn, a source of Poisson process increments.
cpoisson_source(*[, paths, vshape, dtype,])	dj, a source of compound Poisson process increments (jumps).
odd_wiener_source(*[, paths, vshape])	dw, a source of standard Wiener process (Brownian
	motion) increments with antithetic paths exposing op-
	posite increments (averages exactly to 0 across paths).
even_poisson_source(*[, paths, vshape])	dn, a source of Poisson process increments with anti-
	thetic paths exposing identical increments.
even_cpoisson_source(*[, paths, vshape])	dj, a source of compound Poisson process increments
	(jumps) with antithetic paths exposing identical incre-
	ments.
true_source(*[, paths, vshape, dtype, rtol,])	Base class for stochasticity sources with memory.
true_wiener_source(*[, paths, vshape,])	dw, source of standard Wiener process (brownian mo-
	tion) increments with memory.
true_poisson_source(*[, paths, vshape,])	dn, a source of Poisson process increments with mem-
	ory.
<pre>true_cpoisson_source(*[, paths, vshape,])</pre>	dj, a source of compound Poisson process increments
	(jumps) with memory.
norm_rv([a, b])	Normal distribution with mean a and standard devia-
	tion b, possibly time-dependent.
uniform_rv([a, b])	Uniform distribution between a and b, possibly time-
	dependent.
exp_rv([a])	Exponential distribution with scale a, possibly time-
*	dependent.
double_exp_rv([a, b, pa])	Double exponential distribution, with scale a with
<u> </u>	probability pa, and -b with probability (1 - pa), possi-
	bly time-dependent.
rvmap(f, y)	Map f to random variates of distribution y, possibly
± \ / \ /	time-dependent.
	mine dependents

5.1 sdepy.source

class sdepy.**source**(*, paths=1, vshape=(), dtype=None)
Base class for stochasticity sources.

Parameters

paths [int] Number of paths (last dimension) of the source realizations.

vshape [tuple of int] Shape of source values.

dtype [data-type] Data type of source values. Defaults to None.

Returns

array Once instantiated as dz, dz (t, dt) returns a random realization of the stochasticity source increments from time t to time t + dt, with shape (t + dt).shape + vshape + (paths,). For sources with memory (true_source class and subclasses), dz (t) returns the realized value at time t of the source process, according to initial conditions set at instantiation. The definition of source specific parameters, and computation of actual source realizations, are delegated to subclasses. Defaults to an array of numpy.nan.

Notes

Any callable object dz (t, dt), with attributes paths and vshape, returning arrays broadcastable to shape t_shape + vshape + (paths,), where t_shape is the shape of t and/or dt, complies with the source protocol. Such object may be passed to any of the process realization classes, to be used as a stochasticity source in integrating or computing the relevant SDE solution. process instances, in particular, may be used as stochasticity sources.

When calling dz (t, dt), t and/or dt can take any shape.

Attributes

- **size** Returns the number of stored scalar values from previous evaluations, or 0 for sources without memory.
- t Returns a copy of the time points at which source values have been stored from previous evaluations, as an array, or an empty array for sources without memory.

Methods

call(t[, dt])	Realization of stochasticity source values or incre-
	ments.

5.1.1 sdepy.source.__call__

```
source.___call___(t, dt=None)
```

Realization of stochasticity source values or increments.

5.2 sdepy.wiener_source

class sdepy.wiener_source(*, paths=1, vshape=(), dtype=None, corr=None, rho=None) dw, a source of standard Wiener process (Brownian motion) increments.

Parameters

paths [int] Number of paths (last dimension) of the source realizations.

vshape [tuple of int] Shape of source values.

dtype [data-type] Data type of source values. Defaults to None.

corr [array-like, or callable, or None] Correlation matrix of the standard Wiener process increments, possibly time-dependent, or None for no correlations, or for correlations specified by the rho parameter. If not None, overrides rho. If corr is a square matrix of shape (M, M), or callable with corr (t) evaluating to such matrix, the last dimension of the source values must be of size M (vshape[-1] == M), and increments along the last axis of the source values will be correlated accordingly.

rho [array-like, or callable, or None] Correlations of the standard Wiener process increments, possibly time-dependent, or None for no correlations. If rho is scalar,
or callable with rho(t) evaluating to a scalar, M=2 is assumed, and corr=((1,
rho), (rho, 1)). If rho is a vector of shape (K,), or a callable with rho(t)
evaluating to such vector, M=2*K is assumed, and the M source values along the last
vshape dimension are correlated so that rho[i] correlates the i-th and K+i-th values, other correlations being zero (corr = array((I, R), (R, I)) where I
= numpy.eye(K) and R = numpy.diag(rho)).

Returns

array Once instantiated as dw, dw(t, dt) returns a random realization of standard
Wiener process increments from time t to time t + dt, with shape (t + dt).
shape + vshape + (paths,). The increments are normal variates with mean 0,
either independent with standard deviation sqrt(dt), or correlated with covariance
matrix corr*dt, or corr(t + dt/2)*dt (the latter approximates the integral of
corr(t) from t to t + dt).

See also:

source

Notes

Realizations across different t and/or dt array elements, and/or across different paths, and/or along axes of the source values other than the last axis of vshape, are independent. corr should be a correlation matrix with unit diagonal elements and off-diagonal correlation coefficients, not a covariance matrix.

corr and rho values with a trailing one-dimensional paths axis are accepted, of shape (M, M, 1) or (M/2, 1) respectively. This last axis is ignored: this allows for deterministic process instances (single path processes) to be passed as valid corr or rho values. Path dependent corr and rho are not supported.

For time-dependent correlations, dw(t, dt) approximates the increments of a process w(t) obeying the SDE dw(t) = corr(t)*dz(t), where z(t) are standard uncorrelated Wiener processes.

Attributes

corr [array, or callable] Stores the correlation matrix used computing increments. May expose either a reference to corr, if provided explicitly, or an appropriate object, in case rho was specified.

Methods

___call___(t, dt)

See wiener_source class documentation.

5.2.1 sdepy.wiener source. call

```
wiener_source.__call__(t, dt)
```

See wiener_source class documentation.

5.3 sdepy.poisson_source

class sdepy.**poisson_source** (*, paths=1, vshape=(), dtype=<class 'int'>, lam=1.0) dn, a source of Poisson process increments.

Parameters

paths [int] Number of paths (last dimension) of the source realizations.

vshape [tuple of int] Shape of source values.

dtype [data-type] Data type of source values. Defaults to int.

lam [array-like, or callable] Intensity of the Poisson process, possibly time-dependent. Should be an array of non-negative values, broadcastable to shape vshape + (paths,), or a callable with lam(t) evaluating to such array.

Returns

array Once instantiated as dn, dn(t, dt) returns a random realization of Poisson process increments from time t to time t + dt, with shape (t + dt).shape +
vshape + (paths,). The increments are independent Poisson variates with mean
lam*dt, or lam(t + dt/2)*dt (the latter approximates the integral of lam(t)
from t to t + dt).

See also:

source

Attributes

- **size** Returns the number of stored scalar values from previous evaluations, or 0 for sources without memory.
- **t** Returns a copy of the time points at which source values have been stored from previous evaluations, as an array, or an empty array for sources without memory.

Methods

__call___(t, dt)

See poisson_source class documentation.

5.3.1 sdepy.poisson_source.__call__

```
poisson_source.__call__(t, dt)

See poisson_source class documentation.
```

5.4 sdepy.cpoisson_source

class sdepy.**cpoisson_source**(*, paths=1, vshape=(), dtype=None, dn=None, ptype=<class 'int'>, lam=1.0, y=None)
dj, a source of compound Poisson process increments (jumps).

Parameters

paths [int] Number of paths (last dimension) of the source realizations.

vshape [tuple of int] Shape of source values.

dtype [data-type] Data type of source values. Defaults to None.

dn [source or source class, optional] If given, dn is used as the underlying source of Poisson process increments, overriding the ptype and lam parameters.

ptype [data-type] Data type of Poisson process increments. Defaults to int.

lam [array-like, or callable] Intensity of the underlying Poisson process, possibly time-dependent. See poisson_source class documentation.

- y [distribution, or callable, or None] Distribution of random variates to be compounded with the Poisson process increments, possibly time-dependent. May be any scipy.stats distribution instance, or any object exposing an rvs (shape) method to generate independent random variates of the given shape, or a callable with y(t) evaluating to such object. The following preset distributions may be specified, possibly with time-varying parameters:
 - y=norm_rv(a, b) normal distribution with mean a and standard deviation b.
 - y=uniform_rv(a, b) uniform distribution between a and b.
 - y=exp_rv(a) exponential distribution with scale a.
 - y=double_exp_rv(a, b, pa) double exponential distribution, with scale a with probability pa, and -b with probability 1 pa.

where a, b, pa are array-like with values in the appropriate domains, broadcastable to a shape vshape + (paths,), or callables with a(t), b(t), pa(t) evaluating to such arrays. If None, defaults to uniform_rv(a=0, b=1).

Returns

array Once instantiated as dj, dj(t, dt) returns a random realization of compound Poisson process increments from time t to time t + dt, with shape (t + dt). shape + vshape + (paths,). The increments are independent compound Poisson variates, consisting of the sum of N independent y or y(t + dt/2) variates, where N is a Poisson variate with mean lam*dt, or lam(t + dt/2)*dt (approximates each variate being taken from y at the time of the corresponding Poisson process event).

See also:

 $poisson_source, source, norm_rv, uniform_rv, exp_rv, double_exp_rv, rvmap$

Notes

Preset distributions norm_rv, uniform_rv, exp_rv, double_exp_rv behave as follows:

- If all parameters are array-like, return an object with an rvs method as described above, and with methods mean, std, var, exp_mean with signature (), returning the mean, standard deviation, variance and mean of the exponential of the random variate.
- If any parameter is callable, returns a callable y such that y (t) evaluates to the corresponding distribution with parameter values at time t.

To compound the Poisson process increments with a function f(z), or time-dependent function f(t, z), of a given random variate z, one can pass y=rvmap(f, z) to compound_poisson.

[ToDo: make a note on martingale correction using exp_mean]

Attributes

- y [distribution, or callable] Stores the distribution used computing the Poisson process increments
- **dn_value** [array of int] After each realization, this attribute stores the underlying Poisson process increments.
- **y_value** [list of array] After each realization, this attribute stores the underlying y random variates.

Methods

___call__(t, dt)

See cpoisson_source class documentation.

5.4.1 sdepy.cpoisson_source.__call__

```
cpoisson_source.__call__(t, dt)
```

See cpoisson_source class documentation.

5.5 sdepy.odd_wiener_source

class sdepy.odd_wiener_source(*, paths=2, vshape=(), **args)

dw, a source of standard Wiener process (Brownian motion) increments with antithetic paths exposing opposite increments (averages exactly to 0 across paths).

Once instantiated as dw with paths= $2 \times K$ paths, x = dw(t, dt) consists of leading K paths with independent increments, and trailing K paths consisting of a copy of the leading paths with sign reversed (x[..., i] = -x[..., K + i]).

See also:

wiener_source

Attributes

- **size** Returns the number of stored scalar values from previous evaluations, or 0 for sources without memory.
- t Returns a copy of the time points at which source values have been stored from previous evaluations, as an array, or an empty array for sources without memory.

Methods

__call__

5.6 sdepy.even poisson source

class sdepy.even_poisson_source(*, paths=2, vshape=(), **args)

dn, a source of Poisson process increments with antithetic paths exposing identical increments.

Once instantiated as dn with paths=2*K paths, x = dn(t, dt) consists of leading K paths with independent increments, and trailing K paths consisting of a copy of the leading paths: (x[..., i] = x[..., K + i]). Intended to be used together with odd_wiener_source to generate antithetic paths in jump-diffusion processes.

See also:

source, poisson_source

Attributes

- **size** Returns the number of stored scalar values from previous evaluations, or 0 for sources without memory.
- **t** Returns a copy of the time points at which source values have been stored from previous evaluations, as an array, or an empty array for sources without memory.

Methods

call	

5.7 sdepy.even_cpoisson_source

class sdepy.even_cpoisson_source(*, paths=2, vshape=(), **args)

dj, a source of compound Poisson process increments (jumps) with antithetic paths exposing identical increments.

Once instantiated as dj with paths=2*K paths, x = dj(t, dt) consists of leading K paths with independent increments, and trailing K paths consisting of a copy of the leading paths: x[..., i] equals x[..., K + i]. Intended to be used together with odd_wiener_source to generate antithetic paths in jump-diffusion processes.

See also:

source, cpoisson_source

Attributes

- **size** Returns the number of stored scalar values from previous evaluations, or 0 for sources without memory.
- **t** Returns a copy of the time points at which source values have been stored from previous evaluations, as an array, or an empty array for sources without memory.

Methods

__call__

5.8 sdepy.true_source

class sdepy.true_source (*, paths=1, vshape=(), dtype=None, rtol='max', t0=0.0, z0=0.0)

Base class for stochasticity sources with memory.

Parameters

paths, vshape, dtype See source class documentation.

- **rtol** [float, or 'max'] relative tolerance used in assessing the coincidence of t with the time of a previously stored realization of the source. If set to max, the resolution of the float type is used.
- **t0, z0** [array-like] z0 is the initial value of the source at time t0.

Returns

array Once instantiated as dz, dz (t) returns the realized value at time t of the source process, such that dz (t0) = z0, with shape (t + dt).shape + vshape + (paths,), as specified by subclasses. dz (t, dt) returns dz (t + dt) - dz (t). New values of dz (t) should follow a probability distribution conditional on values realized in previous calls. Defaults to an array of numpy.nan.

See also:

source

Attributes

- **size** Returns the number of stored scalar values from previous evaluations, or 0 for sources without memory.
- **t** Returns a copy of the time points at which source values have been stored from previous evaluations, as an array, or an empty array for sources without memory.

Methods

getitem(index)	Reference to a sub-array or element of the source
	values sharing the same memory of past realiza-
	tions.
new_inside(z1, z2, t1, t2, s)	Generate a new process increment, at a time s be-
	tween those of formerly realized values.
$new_outside(\mathbf{z},\mathbf{t},\mathbf{s})$	Generate a new process increment, at a time s
	above or below those of formerly realized values.

5.8.1 sdepy.true source.new inside

true_source.new_inside (z1, z2, t1, t2, s)

Generate a new process increment, at a time s between those of formerly realized values.

Parameters

- **z1**, **z2** [array] Formerly realized values of the source at times t1, t2 respectively.
- **t1, t2** [float] t1, t2 are the times of former realizations closest to s, with t1 < s < t2.

Returns

array Value of the source at s, conditional on formerly realized value z1 at t1 and z2 at t2. Should be defined by subclasses. Defaults to an array of numpy.nan.

5.8.2 sdepy.true_source.new_outside

```
true_source.new_outside (z, t, s)
```

Generate a new process increment, at a time s above or below those of formerly realized values.

Parameters

- **z** [array] Formerly realized value of the source at time t.
- $t,s \hspace{0.1cm} \mbox{[float]} \hspace{0.1cm} \mbox{t} \hspace{0.1cm} \mbox{is the highest (lowest) time of former realizations, and s is above (below) \pm.}$

Returns

array Value of the source at s, conditional on formerly realized value z at t. Should be defined by subclasses. Defaults to an array of numpy.nan.

5.9 sdepy.true_wiener_source

```
class sdepy.true_wiener_source (*, paths=1, vshape=(), dtype=None, corr=None, rho=None, rtol='max', t0=0.0, z0=0.0)
dw, source of standard Wiener process (brownian motion) increments with memory.
```

Parameters

paths, vshape, dtype, corr, rho

See wiener_source class documentation.

rtol, t0, z0 See true_source class documentation.

Returns

array Once instantiated as dw, dw (t) returns z0 plus a realization of the standard Wiener process increment from time t0 to t, and dw (t, dt) returns dw (t + dt) - dw (t). The returned values follow a probability distribution conditional on values realized in previous calls.

Notes

For time-dependent correlations the result is approximate, mind running a first evaluation on a sequence of consecutive closely spaced time points in the region of interest.

Attributes

- **size** Returns the number of stored scalar values from previous evaluations, or 0 for sources without memory.
- **t** Returns a copy of the time points at which source values have been stored from previous evaluations, as an array, or an empty array for sources without memory.

Methods

See source and true_source methods.

5.10 sdepy.true_poisson_source

```
class sdepy.true_poisson_source(*, paths=1, vshape=(), dtype=<class 'int'>, lam=1.0, rtol='max', t0=0.0, z0=0)
dn, a source of Poisson process increments with memory.
```

Parameters

```
paths, vshape, dtype, lam See poisson_source class documentation.
rtol, t0, z0 See true source class documentation.
```

Returns

array Once instantiated as dn, dn(t) returns z0 plus a realization of Poisson process increments from time t0 to t, and dn(t, dt) returns dn(t + dt) - dn(t). The returned values follow a probability distribution conditional on the realized values in previous calls.

See also:

```
source, poisson_source, true_source
```

Notes

For time-dependent intensity lam(t) the result is approximate, mind running a first evaluation on a sequence of consecutive closely spaced time points in the region of interest.

Attributes

- **size** Returns the number of stored scalar values from previous evaluations, or 0 for sources without memory.
- **t** Returns a copy of the time points at which source values have been stored from previous evaluations, as an array, or an empty array for sources without memory.

Methods

See "source" and "true_source" methods.

5.11 sdepy.true_cpoisson_source

```
class sdepy.true_cpoisson_source (*, paths=1, vshape=(), dtype=None, rtol='max', t0=0.0, z0=0.0, dn=None, ptype=<class 'int'>, lam=1.0, v=None)
```

dj, a source of compound Poisson process increments (jumps) with memory.

Parameters

```
paths, vshape, dtype, dn, ptype, lam, y See cpoisson_source class documentation. rtol, t0, z0 See true_source class documentation.
```

Returns

array Once instantiated as dj, dj(t) returns z0 plus a realization of compound Poisson process increments from time t0 to t, and dj(t, dt) returns dj(t + dt) - dj(t). The returned values follow a probability distribution conditional on the realized values in previous calls.

See also:

source, cpoisson_source, true_source

Notes

For time-dependent intensity lam(t) and compounding random variable y(t) the result is approximate, mind running a first evaluation on a sequence of consecutive closely spaced time points in the region of interest.

Attributes

- **size** Returns the number of stored scalar values from previous evaluations, or 0 for sources without memory.
- t Returns a copy of the time points at which source values have been stored from previous evaluations, as an array, or an empty array for sources without memory.

Methods

See "source" and "true_source" methods.

5.12 sdepy.norm_rv

```
sdepy.norm_rv (a=0, b=1)
```

Normal distribution with mean a and standard deviation b, possibly time-dependent.

```
Wraps scipy.stats.norm(loc=a, scale=b).
```

See also:

cpoisson_source

5.13 sdepy.uniform rv

```
sdepy.uniform_rv(a=0, b=1)
```

Uniform distribution between a and b, possibly time-dependent.

```
Wraps scipy.stats.uniform(loc=a, scale=b-a).
```

See also:

cpoisson_source

5.14 sdepy.exp_rv

```
sdepy.exp_rv(a=1)
```

Exponential distribution with scale a, possibly time-dependent.

Wraps scipy.stats.expon(scale=a). The probability distribution function is:

```
• if a > 0, pdf(x) = a \times exp(-a \times x), with support in [0, inf)
```

• if
$$a < 0$$
, pdf(x) = $-a \times exp(a \times x)$, with support in $(-inf, 0]$

See also:

cpoisson_source

5.15 sdepy.double_exp_rv

```
sdepy.double_exp_rv (a=1, b=1, pa=0.5)
```

Double exponential distribution, with scale a with probability pa, and -b with probability (1 - pa), possibly time-dependent.

Double exponential distribution, with probability distribution

```
• for x in [0, inf), pdf(x) = pa*exp(-a*x)*a
```

• for x in
$$(-\inf, 0)$$
, pdf $(x) = (1-pa) *exp(b*x) *b$

where a and b are positive and pa is in [0, 1].

See also:

cpoisson_source

5.16 sdepy.rvmap

```
sdepy.rvmap(f, y)
```

Map f to random variates of distribution y, possibly time-dependent.

Parameters

- **f** [callable] Callable with signature f(y), or f(t, y) or f(s, y), to be mapped to the random variates of y or y(t)
- ${\bf y}$ [distribution, or callable] Distribution, possibly time-dependent, as accepted by cpoisson_source.

Returns

new_y [Distribution, or callable] An object with and rvs(shape) method,
 or a callable with new_y(t) evaluating to such object, as accepted by
 cpoisson_source. new_y.rvs(shape), or new_y(t).rvs(shape),
 returns f(y.rvs(shape)), or f([t,] y(t).rvs(shape).

See also:

cpoisson_source, norm_rv, uniform_rv, exp_rv, double_exp_rv

Notes

new_y does not provide any mean, std, var, exp_mean method.

To be recognized as time-dependent, f should have its first parameter named t or s.

SDE Integration Framework

<pre>paths_generator(*[, paths, xshape, wshape,</pre>	Step by step generation of stochastic simulations
])	across multiple paths, intended for subclassing.
<pre>integrator(*[, paths, xshape, wshape,])</pre>	Step by step numerical integration of Ito Stochastic
	Differential Equations (SDEs), intended for subclass-
	ing.
SDE(*[, paths, vshape, dtype, steps, i0,])	Class representation of a user defined Stochastic Dif-
	ferential Equation (SDE), intended for subclassing.
SDEs(*[, paths, vshape, dtype, steps, i0,])	Class representation of a user defined system of
	Stochastic Differential Equations (SDEs), intended
	for subclassing.
<pre>integrate([sde, q, sources, log, addaxis])</pre>	Decorator for Ito Stochastic Differential Equation
	(SDE) integration.

6.1 sdepy.paths_generator

Step by step generation of stochastic simulations across multiple paths, intended for subclassing.

Given a number of requested paths, shapes and output timeline, encapsulates the low level tasks of memory allocation and step by step iteration along the timeline.

The definition of the target iteration steps (pace method), initialization (begin method), computation of next step (next method), storing results at points on the requested timeline (store method), cleaning up (end method) and evaluation of a final result to be returned (exit method), are delegated to subclasses.

Instances are callables with signature (timeline) that iterate subclass methods along the given timeline, using the configuration set out at instantiation.

Parameters

paths [int] Size of last axis of the allocated arrays (number of paths of the simulation).

xshape [int or tuple of int] Shape of values that will be stored at each point of the output timeline.

wshape [int or tuple of int] Shape of working space used for step by step iteration.

dtype [data-type] Data-type of the output and working space.

steps [iterable, or int, or None] Specification of the time points to be touched during the simulation (as defined by the pace method). Default behaviour is:

- if None, the simulated steps coincide with the timeline;
- if int, the simulated steps touch all timeline points, as well as steps equally spaced points between the minimum and maximum point in the timeline;
- if iterable, the simulated steps touch all timeline points, as well as all values in steps between the minimum and maximum points in the timeline.
- i0 [int] Index along the timeline at which the simulation starts. The timeline is assumed to be in ascending order. The simulation is performed backwards from timeline[i0] to timeline[0], and forwards from timeline[i0] to timeline[-1].
- **info** [dict, or None] Diagnostic information about the simulation is stored in this dictionary and is accessible as the info attribute. If None, a new empty dict is used.
- **getinfo** [bool] If True (default), records basic information in the info attribute about the last performed simulation (if the simulation is both backwards and forwards, the information pertains to the forwards part only). Used by subclasses to enable/disable diagnostic info generation.

Returns

simulation results Once instantiated as p, p (timeline) runs the simulation along the given timeline, based on parameters of instantiation, returning results as determined by subclass methods. Defaults to (tt, xx) where tt is a reference to timeline and xx is an array of numpy.nan of the requested shape.

See also:

integrator, SDE, SDEs

Notes

All initialization parameters are stored as attributes of the same name, and may be accessed by subclasses.

During the simulation, a itervars attribute is present, pointing to a dictionary that contains the following items, to be used by subclass methods (double letters refer to values along the entire timeline, single letters refer to single time points):

- steps_tt: an array of all time points to be touched by the simulation. It consolidates the output timeline and the time points to be touched, as specified by steps.
- tt: the output timeline.
- xx: simulation output, an array of shape tt.shape + xshape + (paths,). xx[i] is the simulated value at time tt[i].
- sw: working space for time points, an array of shape (depth,).
- xw: working space for paths generation, an array of objects of shape (depth,), where each of xw[k] is an array of shape wshape + (paths,).
- reverse: True if the simulation runs backwards, False otherwise. If True, steps_tt and tt are in descending order.
- i : such that tt[i] is the next point along the timeline that will be reached (when invoking next), or the point that was just reached (when invoking store).

Note that:

• sw and xw are rolled at each iteration: subclass methods should not rely on storing references to their elements across iterations.

- xw[k][...] = value broadcasts value into the allocated memory of xw[k] (this is usually what you want), xw[k] = value stores value, as an object, in xw[k] (avoid).
- xx and xw[k] are initialized to arrays filled with numpy.nan.

Attributes

depth [int] Number of time points to be stored in the working space. Defaults to 2.

Methods

call(timeline)	Run the simulation along the given timeline.
pace(timeline)	Target integration steps for the current integration.
begin()	Set initial conditions.
next()	Numerical simulation step.
store(i, k)	Store the current integration step into the integra-
	tion output.
end()	End of iteration optional tasks.
exit(tt, xx)	Final tasks and construction of the output value(s).

6.1.1 sdepy.paths generator. call

 $\verb|paths_generator.__call__(\textit{timeline})|$

Run the simulation along the given timeline.

Parameters

timeline [array-like] A one dimensional array of strictly increasing numbers, defining the timeline of the simulation.

Returns

Simulation results, as specified by subclass methods.

6.1.2 sdepy.paths generator.pace

paths_generator.pace(timeline)

Target integration steps for the current integration.

Parameters

timeline [array] Requested simulation timeline, cast as an array of float data-type.

Returns

array Target time points to be touched during the simulation (typically, more thinly spaced than the output time points in timeline, based on the steps parameter), to be merged with timeline.

May be overridden by subclasses. For default behaviour,

see paths_generator class documentation.

6.1.3 sdepy.paths_generator.begin

paths_generator.begin()

Set initial conditions.

Given the time points sw[0], ..., sw[depth - 2], should define and store in the working space the corresponding initial values xw[0], ..., xw[depth - 2]. Note that when begin

gets called, sw[depth - 1] and sw[depth - 1] are undefined and will be respectively set, and computed, at the first iteration.

Notes

It is called once for each backwards and forwards simulation, after memory allocation and before starting the iteration along the time points in steps_tt.

Outline of expected code for depth=2:

```
# access itervars
iv = self.itervars
sw, xw = iv['sw'], iv['xw']

# this is the initial time, taken from the
# simulation timeline
t0 = sw[0]
assert t0 == iv['steps_tt'][0] == iv['tt'][0]

# store the initial condition
xw[0][...] = 1.
```

Must be provided by subclasses.

6.1.4 sdepy.paths_generator.next

```
paths_generator.next()
Numerical simulation step.
```

Given the points sw[0], ..., sw[depth - 2] and the corresponding values xw[0], ..., xw[depth - 2], should:

- Optionally modify the target next time point sw[depth 1], to a value between sw[depth 2] and sw[depth 1] (this allows for adaptive time steps, with the constraint of touching all point specified by the output timeline and the steps parameter).
- 2. Compute the corresponding value xw[depth 1]

Notes

It is called once per iteration step.

Outline of expected code for depth=2:

```
# access itervars
iv = self.itervars
sw, xw = iv['sw'], iv['xw']

# get starting values, and time step to be taken
s0, x0 = sw[0], xw[0]
ds = sw[1] - sw[0]

# compute and store the next step
xw[1][...] = x0 + ds
```

Must be provided by subclasses.

6.1.5 sdepy.paths_generator.store

```
paths_generator.store (i, k)
```

Store the current integration step into the integration output.

Should take the k-th value in the working space xw, transform it if need be, and store it as the output xx[i] at the output time point tt[i].

Parameters

- i [int] Index of the output timeline point to set as output.
- **k** [int] Index of the working space point to use as input.

Notes

It is called initially to store the initial values that belong to the output timeline, among those put into the working space by begin, and later during the iteration, each time the simulation touches a point on the output timeline.

Outline of expected code for xshape == wshape and an exponentiation transformation:

```
# access itervars
iv = self.itervars
sw, xw = iv['sw'], iv['xw']
xx = iv['xx']

# this is the current time, also found
# along the output timeline
s = sw[k]
assert s == iv['tt'][i]

# transform and store
np.exp(xw[k], out=xx[i])
```

Must be provided by subclasses.

6.1.6 sdepy.paths generator.end

```
paths_generator.end()
```

End of iteration optional tasks.

It is called once for each backwards and forwards simulation, once the final point in the output timeline has been reached and the simulation ends.

After it is called, itervars are deleted.

May be provided by subclasses.

6.1.7 sdepy.paths_generator.exit

```
paths_generator.exit(tt, xx)
```

Final tasks and construction of the output value(s).

Parameters

- **tt** [array] Output timeline. It is the timeline passed to the __call__ method, cast as an array, with its original data-type (if the data-type is of integer kind, the simulation is carried out using floats).
- **xx** [array] Output values along the timeline, as computed by next and stored by store methods.

Notes

It is called once, after backwards and/or forwards simulations have been completed, and its return value is returned.

Default implementation:

```
return tt, xx
```

May be provided by subclasses.

6.2 sdepy.integrator

Step by step numerical integration of Ito Stochastic Differential Equations (SDEs), intended for subclassing.

For usage, see the SDE class documentation.

This class encapsulates SDE integration methods, and cooperates with the SDE class, that should always have precedence in method resolution order. As long as the respective APIs are complied with, a new integrator stated as an integrator subclass will interoperate with existing SDEs (as described by SDE subclasses), and a new SDE will interoperate with existing integrators.

Parameters

paths, xshape, wshape, dtype, steps, i0, info, getinfo See paths_generator class
documentation.

method [string] Integration method. Defaults to 'euler', for the Euler-Maruyama method (at present, this single method is supported). It is stored as an attribute of the same name.

Returns

process Once instantiated as p, p (timeline) performs the integration along the given timeline, based on parameters of instantiation, returning the resulting process as determined by the cooperating SDE subclass and the chosen integration method. Defaults to a process of numpy.nan along the given timeline.

See also:

```
paths_generator, SDE, SDEs
```

Notes

The equation to be integrated is exposed to the integration algorithm in a standardized form, via methods A and dZ delegated to a cooperating SDE class. The latter should take care of equation parameters, initial conditions, expected paths and shapes, and should instantiate all necessary stochasticity sources.

The integration method is exposed as the next method to the paths generator parent class.

If the getinfo attribute is set to True, at each integration step the following items are added to the itervars dictionary, made available to subclasses to track the integration progress:

- last_t: starting time point of the last integration step.
- last_dt: time increment of the last integration step.
- last_x: starting value of the process, at time last_t.
- last_A: dictionary of the last computed values of the SDE terms, at time last_t.

- last_dZ: dictionary of the last realized SDE stochasticity source values, cumulated in the interval from last_t to last_t + last_dt.
- new_x: computed value of the process, at time last_t + last_dt.

This becomes relevant in case the output timeline is coarse (e.g. just the initial and final time) but diagnostic information is needed about all integration steps performed (e.g., to track how often the process has changed sign, or to count the number of realized jumps).

Methods

A(t, x)	Value of the SDE terms at time t and process value
	х.
dZ(t, dt)	Value of the SDE differentials at time t, for time
	increment dt.
next()	Perform an integration step with the requested
	method.
euler_next()	Euler-Maruyama integration step.

6.2.1 sdepy.integrator.A

```
integrator. \mathbf{A}(t, x)
```

Value of the SDE terms at time t and process value x.

Example of expected code for the SDE dx = (1 - x) * dt + 2 * dw (t):

```
return {
   'dt': (1 - x),
   'dw': 2
   }
```

The SDE class takes care of casting user-specified equations into this format.

6.2.2 sdepy.integrator.dZ

```
integrator.dZ(t, dt)
```

Value of the SDE differentials at time t, for time increment dt.

Example of expected code for the SDE dx = (1 - x) * dt + 2 * dw (t), where x has two components:

```
shape = (2, self.paths)
return {
    'dt': dt,
    'dw': wiener_source(vshape=2, paths=self.paths)(0, dt)
    }
```

The SDE class takes care of instantiating user-specified stochasticity sources and casting them into this format.

6.2.3 sdepy.integrator.next

```
integrator.next()
```

Perform an integration step with the requested method.

6.2.4 sdepy.integrator.euler next

```
integrator.euler_next()
    Euler-Maruyama integration step.
```

6.3 sdepy.SDE

Class representation of a user defined Stochastic Differential Equation (SDE), intended for subclassing.

This class aims to provide an easy to use and flexible interface, allowing to specify user-defined SDEs and expose them in a standardized form to the cooperating integrator class (the latter should always follow in method resolution order). A minimal definition of an Ornstein-Uhlenbeck process is as follows:

```
>>> from sdepy import SDE, integrator
>>> class my_process(SDE, integrator):
...     def sde(self, t, x, theta=1., k=1., sigma=1.):
...     return {'dt': k*(theta - x), 'dw': sigma}
```

An SDE is stated as a dictionary, containing for each differential the value of the corresponding coefficient:

```
dx = f(t, x)*dt + g(t, x)*dw + h(t, x)*dj
```

translates to:

```
{'dt': f(t, x), 'dw': g(t, x), 'dj': h(t, x)}
```

Instances are callables with signature (timeline) that integrate the SDE along the given timeline, using the configuration set out in the instantiation parameters:

```
>>> P = my_process(x0=1, sigma=0.5, paths=100*1000, steps=100)
>>> x = P(timeline=(0., 0.5, 1.))
>>> x.shape
(3, 100000)
```

Subclasses can specify or customize: the equation and its parameters (sde method), initial conditions and preprocessing (init method and log attribute), shape of the values to be computed and stored (shapes method), stochastic differentials appearing in the equation (sources attribute) and their parameters and initialization (methods source_dt, source_dw, source_dn, source_dj, or any custom source_{id} method for a corresponding differential '{id}' declared in sources and used as a key in sde return values), optional non array-like parameters (more method), how to store results at points on the requested timeline (let method), and postprocessing (result method and log attribute).

Parameters

paths [int] Number of paths of the process.

vshape [int or tuple of int] Shape of the values of the process.

dtype [data-type, optional] Data-type of the process. Defaults to the numpy default.

steps [iterable, or int, or None] Specification of the time points to be touched during integration (as accepted by a cooperating integrator class). Default behaviour is:

- if None, the simulated steps coincide with the timeline;
- if int, the simulated steps touch all timeline points, as well as steps equally spaced points between the minimum and maximum point in the timeline;
- if iterable, the simulated steps touch all timeline points, as well as all values in steps between the minimum and maximum points in the timeline.

- i0 [int] Index along the timeline at which the integration starts. The timeline is assumed to be in ascending order. Initial conditions are set at timeline[i0], the integration is performed backwards from timeline[i0] to timeline[0], and forwards from timeline[i0] to timeline[-1].
- **info** [dict, optional] Diagnostic information about the integration is stored in this dictionary and is accessible as the info attribute. Defaults to a new empty dict.
- **getinfo** [bool] If True, subclass methods info_begin, info_next, info_store, info_end are invoked during integration. Defaults to True.
- method [str] Integration method, as accepted by the integrator cooperating class.
- **args [SDE-specific parameters] SDE parameters and initial conditions, as implied by the signature of sde, init and more methods, and stochasticity sources parameters, as implied by the signature of source_{id} methods. Each keyword should be used once (e.g. corr, a source_dw parameter, cannot be used as the name of a SDE parameter).

Returns

process Once instantiated as p, p (timeline) performs the integration along the given timeline, based on parameters of instantiation, and returns the resulting process as defined by subclass methods. Defaults to a process of numpy.nan along the given timeline.

See also:

paths_generator, integrator, SDEs

Notes

Custom stochastic differentials used in the SDE should be recognized, and treated appropriately, by the chosen integration method. This may require customization of the next method of the integrator class.

All named initialization parameters (paths, steps etc.) are stored as attributes.

Notes on SDE-specific parameters:

- init parameters are converted to arrays via np.asarray.
- sde and source quantitative parameters may be array-like, or time dependent with signature (t).
- both are converted to arrays via np.asarray, and for both, their constant value, or values at each time point, should be broadcastable to a shape wshape + (paths,).
- more parameters undergo no further initialization, before being made available to the shapes and more methods.

If getinfo is True, the invoked info subclass methods may initialize and cumulate diagnostic information in items of the info dictionary, based on read-only access of the internal variables set during integration by paths_generator and integrator cooperating classes, as exposed in the itervars attribute.

Attributes

sources [set or dict] As a class attribute, holds the names of the differentials 'dz' expected to appear in the equation. As an instance attribute, sources['dz'] is an object, complying with the source protocol, that instantiates the differential 'dz' used during integration. sources['dz'](t, dt) is computed at every step for each 'dz' in sources, as required by the chosen integration method.

args [dict] Stores parameters passed as **args upon initialization of the SDE.

log [bool] If True, the natural logarithm of the initial values set by the init method is taken as the initial value of the integration, and the result of the integration is exponentiated

6.3. sdepy.SDE 49

back before serving it to the result method. The sde should expose the appropriate equation for integrating the logarithm of the intended process.

Methods

sde(t, x) Stochastic Differential Equation (SDE) to be in grated. shapes(vshape) Shape of the values to be computed and storage upon integration of the SDE. source_dt() Setup a source of deterministic increments, to used as 'dt' during integration. source_dw([dw, corr, rho]) Setup a source of standard Wiener process (Brain motion) increments, to be used as 'dw' durintegration.	ored o be
Shape of the values to be computed and sto upon integration of the SDE. Source_dt() Setup a source of deterministic increments, to used as 'dt' during integration. Source_dw([dw, corr, rho]) Setup a source of standard Wiener process (Brinian motion) increments, to be used as 'dw' du	o be
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Source_dw([dw, corr, rho]) Setup a source of standard Wiener process (Br nian motion) increments, to be used as 'dw' du	
nian motion) increments, to be used as 'dw' du	
	ring
integration	
source_dn([dn, ptype, lam]) Setup a source of Poisson process increments	, to
be used as 'dn' during integration.	
source_dj([dj, dn, ptype, lam, y]) Set up a source of compound Poisson process	
crements (jumps), to be used as 'dj' during i	nte-
gration.	
more() Further optional non array parameters, and init	tial-
izations.	
init(t, out_x[, x0]) Set initial conditions for SDE integration.	
let(t, out_x, x) Store the value of the integrated process at t	
point t belonging to the requested output time.	ine.
result(tt, xx) Compute the integration output.	
info_begin() Optional diagnostic information logging function	ion,
called before the integration begins.	
info_next() Optional diagnostic information logging function	ion,
called after each integration step.	
info_store() Optional diagnostic information logging function	ion,
called after each invocation of the let method.	
info_end() Optional diagnostic information logging function	
called after the integration has been completed	•

6.3.1 sdepy.SDE.sde

 \mathtt{SDE} . \mathbf{sde} (t, x)

Stochastic Differential Equation (SDE) to be integrated.

Parameters

- t [float] Time point at which the SDE should be evaluated.
- \boldsymbol{x} [array] Values that the stochastic process takes at time t.

sde_args [zero or more arrays, as keyword arguments] SDE parameters, as implied by the sde method signature. Passed upon instantiation of the SDE class as possibly time-dependent array-like, these parameters are served to the sde method once evaluated at t and converted to arrays via np.asarray.

Returns

sde_terms [dict of arrays] Contains, for each differential stated in the source attribute, the value of the corresponding coefficient in the represented SDE.

Notes

x should be treated as read-only.

6.3.2 sdepy.SDE.shapes

SDE.shapes (vshape)

Shape of the values to be computed and stored upon integration of the SDE.

Parameters

vshape [int or tuple of int] Shape of the values of the integration result, as requested upon instantiation of SDE.

Returns

vshape [int or tuple of int] Confirms or overrides the given vshape.

xshape [int or tuple of int] Shape of the values stored during integration at the output time points. out_x array passed to the let method has shape xshape +
 (paths,). Defaults to vshape.

wshape [int or tuple of int] Shape of the working space used during integration. x values
 passed to the sde and let methods have shape wshape + (paths,). Defaults
 to vshape.

Notes

xshape and wshape are passed to the parent paths generator class.

hull_white_SDE and heston_SDE classes illustrate use cases for different values of vshape, xshape and/or wshape.

6.3.3 sdepy.SDE.source_dt

```
SDE.source_dt()
```

Setup a source of deterministic increments, to be used as 'dt' during integration.

Returns

```
An object "z" complying with the "source" protocol, such that z(t, dt) = dt.
```

6.3.4 sdepy.SDE.source_dw

```
\verb"SDE.source_dw" (\textit{dw=None}, \textit{corr=None}, \textit{rho=None})
```

Setup a source of standard Wiener process (Brownian motion) increments, to be used as 'dw' during integration.

Parameters

dw [source, or source subclass, or None] If an object complying with the source protocol, it is returned (corr and rho are ignored). If a source subclass, it is instantiated with the given parameters, and returned. If None, a new instance of wiener_source is returned, with the given parameters.

corr, rho [see wiener_source documentation]

Returns

An object complying with the "source" protocol,

6.3. sdepy.SDE 51

instantiating the requested stochasticity source.

The shape of source values is set to "wshape".

See also:

wiener_source

6.3.5 sdepy.SDE.source_dn

SDE.source_dn (dn=None, ptype=<class 'int'>, lam=1.0)

Setup a source of Poisson process increments, to be used as 'dn' during integration.

Parameters

dn [source, or source subclass, or None] If an object complying with the source protocol, it is returned (ptype and lam are ignored). If a source subclass, it is instantiated with the given parameters, and returned. If None, a new instance of poisson_source is returned, with the given parameters.

ptype, lam [see poisson_source documentation]

Returns

An object complying with the "source" protocol,

instantiating the requested stochasticity source.

The shape of source values is set to "wshape".

See also:

poisson_source

6.3.6 sdepy.SDE.source_dj

SDE.source_dj (dj=None, dn=None, ptype=<class 'int'>, lam=1.0, y=None)

Set up a source of compound Poisson process increments (jumps), to be used as 'dj' during integration.

Parameters

dj [source, or source subclass, or None] If an object complying with the source protocol, it is returned (ptype, lam and y are ignored). If a source subclass, it is instantiated with the given parameters, and returned. If None, a new instance of cpoisson_source is returned, with the given parameters.

ptype, lam, y [see cpoisson_source documentation]

Returns

An object complying with the "source" protocol,

instantiating the requested stochasticity source.

The shape of source values is set to "wshape".

See also:

cpoisson_source

6.3.7 sdepy.SDE.more

SDE.more()

Further optional non array parameters, and initializations.

Parameters

more_args [zero or more keyword arguments] Further, possibly non array-like, SDE parameters, as implied by the more method signature. Passed upon instantiation of the SDE class, are served to the more method and made available to other methods as items in the args attribute.

Notes

The factors parameter of hull_white_SDE illustrates a use case for the more method.

6.3.8 sdepy.SDE.init

```
SDE.init (t, out_x, x0=1.0)
```

Set initial conditions for SDE integration.

Parameters

t [float] Time point at which initial conditions should be imposed.

out_x [array] Array, shaped wshape + (paths,), where initial conditions are to be stored.

init_args [zero or more arrays, as keyword arguments] Initialization parameters, as implied by the init method signature. Passed upon instantiation of the SDE class as array-like, these parameters are served to the init method converted to arrays via np.asarray.

Notes

The default implementation has a single x0 parameter, and sets out_x[...] = x0.

6.3.9 sdepy.SDE.let

```
SDE.let (t, out\_x, x)
```

Store the value of the integrated process at time point t belonging to the requested output timeline.

Parameters

t [float] Time point to which the integration result x refers.

 out_x [array] Array, shaped xshape + (paths,), where the result x is to be stored.

x [array] Integration result at time t, shaped wshape + (paths,)

Notes

The default implementation sets $out_x[...] = x$.

In case xshape != wshape, this method should operate as needed in order to store in $out_x a$ value broadcastable to its shape (e.g. it might store in $out_x only some of the components of x).$

x should be treated as read-only.

6.3.10 sdepy.SDE.result

```
SDE . result (tt, xx)
```

Compute the integration output.

Parameters

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```
tt [array] Output integration timeline.
```

```
xx [array] Integration result, shaped tt.shape + xshape + (paths,).
```

Returns

result Final result, returned to the user.

Notes

The default implementation returns sdepy.process(t=tt, x=xx). In case vshape != xshape, this method should operate as needed in order to return a process with values shaped as yshape (e.g. it might return a function of the components of xx).

6.3.11 sdepy.SDE.info begin

```
SDE.info_begin()
```

Optional diagnostic information logging function, called before the integration begins.

6.3.12 sdepy.SDE.info next

```
SDE.info_next()
```

Optional diagnostic information logging function, called after each integration step.

6.3.13 sdepy.SDE.info_store

```
SDE.info_store()
```

Optional diagnostic information logging function, called after each invocation of the let method.

6.3.14 sdepy.SDE.info_end

```
SDE.info_end()
```

Optional diagnostic information logging function, called after the integration has been completed.

6.4 sdepy.SDEs

```
class sdepy.SDEs(*, paths=1, vshape=(), dtype=None, steps=None, i0=0, info=None, get-
info=True, method='euler', **args)
```

Class representation of a user defined system of Stochastic Differential Equations (SDEs), intended for subclassing.

The parent SDE class represents a single SDE, scalar or multidimensional: by an appropriate choice of the vshape parameter, and composition of equation values, it suffices to describe any system of SDEs.

Its SDEs subclass is added for convenience of representation: it allows to state each equation separately and to retrieve separate processes as a result. The number of equations must be stated as the q attribute. The vshape parameter is taken as the common shape of values in each equation in the system.

A minimal definition of a lognormal process x with stochastic volatility y is as follows:

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```
... {'dt': 0, 'dw': xi*y})

>>> P = my_process(x0=(1., 2.), xi=0.5, vshape=5,
... paths=100*1000, steps=100, )

>>> x, y = P(timeline=(0., 0.5, 1.))

>>> x.shape, y.shape
((3, 5, 100000), (3, 5, 100000))
```

See also:

SDE, integrator, paths_generator

Notes

By default, the stochasticity sources of each component equation are realized independently, even if represented in the sde output by the same key ('dw' in the example above).

The way stochasticity sources are instantiated and dispatched to each equation, and how correlations of the Wiener source are set via the corr parameter, depend on the value of the addaxis attribute:

- If True, source values have shape vshape + (q,), and the [kk, i] component of source values is dispatched to the kk component of equation i (kk is a multiindex spanning shape vshape). If given, corr must be of shape (q, q) and correlates corresponding components across equations.
- If addaxis is False (default) and N is the size of the last axis of vshape, the values of the sources have shape vshape[:-1] + (N*q,), and the [kk, i*N + h] component of the source values is dispatched to the [kk, h] component of equation i (kk is a multiindex spanning shape vshape[:-1], and h is in range (N)). If given, corr must be of shape (N*q, N*q), and correlates all last components of all equations to each other.

After instantiation, stochasticity sources and correlation matrices may be inspected as follows:

```
>>> P = my_process(vshape=(), rho=0.5)
>>> P.sources['dw'].vshape
(2,)
>>> P.sources['dw'].corr.shape
(2, 2)
>>> P.sources['dw'].corr[0, 1]
0.5
```

Attributes

q [int] Number of equations.

addaxis [bool] Affects the internal representation of the equations: if True, a last axis of size q is added to vshape, if False, components are stacked onto the last axis of vshape. Defaults to False. It is forced to True if the process components have scalar values.

Methods

pack(xs)	Packs the given arrays (one per equation) into a single array.
unpack(X)	Unpacks the given array into multiple arrays (one per equation).

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6.4.1 sdepy.SDEs.pack

```
SDEs.pack (xs)
```

Packs the given arrays (one per equation) into a single array.

Parameters

xs [list of arrays] List of self.q arrays to be packed according to the addaxis attribute setting.

Returns

X [array] Array packing the given xs along its second-last dimension (the last dimension enumerates paths).

6.4.2 sdepy.SDEs.unpack

```
SDEs.unpack (X)
```

Unpacks the given array into multiple arrays (one per equation).

Parameters

X [array] Array with a last dimension enumerating paths, and a second last dimension to be unpacked according to the addaxis attribute setting.

Returns

x, y, ... [list of arrays] List of self.q arrays, unpacking the given X.

6.5 sdepy.integrate

sdepy.integrate(sde=None, *, q=None, sources=None, log=False, addaxis=False)
Decorator for Ito Stochastic Differential Equation (SDE) integration.

Decorates a function representing the SDE or SDEs into the corresponding integrator (a subclass of SDE or SDEs and of integrator).

Parameters

- **sde** [function] Function to be wrapped. Its signature and values should be as expected for the sde method of the SDE or SDEs classes.
- q [int] Number of equations. If None, attempts a test evaluation of sde to find out.
- **sources** [set] Stochasticity sources used in the equation. If None, attempts a test evaluation of sde to find out.

log [bool] Sets the log attribute for the wrapping class.

addaxis [bool] Sets the addaxis attribute for the wrapping class.

Examples

```
>>> from sdepy import integrate
>>> @integrate
... def my_process(t, x, theta=1., k=1., sigma=1.):
... return {'dt': k*(theta - x), 'dw': sigma}
```

```
>>> P = my_process(x0=1, sigma=0.5, paths=100*1000, steps=100)
>>> x = P(timeline=(0., 0.5, 1.))
>>> x.shape
(3, 100000)
```

CHAPTER 7

Stochastic Processes

Lognorm_process([paths, vshape, dtype,]) Lognormal process	wiener_process([paths, vshape, dtype,])	Wiener process (Brownian motion) with drift.
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		with compound Poisson logarithmic jumps).
Continued on next page	<pre>merton_jumpdiff_SDE(*[, paths, vshape,])</pre>	SDE for a Merton jump-diffusion process.
		Continued on next page

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Table 1 – continued from previous page

kou_jumpdiff_SDE(*[, paths, vshape, dtype, SDE for a double exponential (Kou) jump-diffusion process.

7.1 sdepy.wiener_process

class sdepy.wiener_process (paths=1, vshape=(), dtype=None, steps=None, i0=0, info=None, getinfo=True, method='euler', x0=0., mu=0., sigma=1., dw=None, corr=None, rho=None)

Wiener process (Brownian motion) with drift.

Generates a process x(t) that solves the following SDE:

```
dx(t) = mu(t)*dt + sigma(t)*dw(t, dt)
```

where dw(t, dt) are standard Wiener process increments with correlation matrix specified by corr(t) or rho(t).x0, SDE parameters and dw(t, dt) should broadcast to vshape + (paths,).

Parameters

paths, vshape, dtype, steps, i0, info, getinfo, method See SDE class documentation.

x0 [array-like] Initial condition.

mu, sigma [array-like, or callable] SDE parameters.

dw, corr, rho Specification of stochasticity source of Wiener process increments. See SDE.source_dw documentation.

Returns

x [process] Once instantiated as p, p(timeline) performs the integration along the given timeline, based on parameters of instantiation, and returns the resulting process.

See also:

```
SDE, SDE.source_dw, wiener_source, wiener_SDE
```

Attributes

See SDE class documentation.

Methods

See SDE class documentation.

7.2 sdepy.lognorm_process

class $sdepy.lognorm_process(paths=1, vshape=(), dtype=None, steps=None, i0=0, info=None, getinfo=True, method='euler', x0=1., mu=0., sigma=1., dw=None, corr=None, rho=None)$

Lognormal process.

Generates a process x(t) that solves the following SDE:

```
dx(t) = mu(t)*x(t)*dt + sigma(t)*x(t)*dw(t, dt)
```

where dw(t, dt) are standard Wiener process increments with correlation matrix specified by corr(t) or rho(t). x0, SDE parameters and dw(t, dt) should broadcast to vshape + (paths,). x0 should be positive.

Parameters

paths, vshape, dtype, steps, i0, info, getinfo, method See SDE class documentation.

x0 [array-like] Initial condition.

mu, sigma [array-like, or callable] SDE parameters.

dw, corr, rho Specification of stochasticity source of Wiener process increments. See SDE.source dw documentation.

Returns

x [process] Once instantiated as p, p(timeline) performs the integration along the given timeline, based on parameters of instantiation, and returns the resulting process.

See also:

SDE, SDE.source_dw, wiener_source, wiener_SDE

Notes

x(t) is obtained via Euler-Maruyama numerical integration of the following equivalent SDE for a (t) = log(x(t)):

```
da(t) = (mu(t) - sigma(t) **2/2) *dt + sigma(t) *dw(t, dt)
```

Attributes

See SDE class documentation.

Methods

See SDE class documentation.

7.3 sdepy.ornstein_uhlenbeck_process

```
 \begin{array}{lll} \textbf{class} & \texttt{sdepy.ornstein\_uhlenbeck\_process} \ (paths=1, & vshape=(), & dtype=None, \\ & steps=None, & i0=0, & info=None, & getinfo=True, \\ & method='euler', & x0=0., & theta=0., & k=1., \\ & sigma=1., & dw=None, & corr=None, & rho=None) \end{array}
```

Ornstein-Uhlenbeck process (mean-reverting Brownian motion).

Generates a process x(t) that solves the following SDE:

```
dx(t) = k(t) * (theta(t) - x(t)) * dt + sigma(t) * dw(t, dt)
```

where dw(t, dt) are standard Wiener process increments with correlation matrix specified by corr(t) or rho(t). x0, SDE parameters and dw(t, dt) should broadcast to vshape + (paths,).

Parameters

paths, vshape, dtype, steps, i0, info, getinfo, method See SDE class documentation.

x0 [array-like] Initial condition.

theta, k, sigma [array-like, or callable] SDE parameters.

dw, corr, rho Specification of stochasticity source of Wiener process increments. See SDE.source_dw documentation.

Returns

x [process] Once instantiated as p, p(timeline) performs the integration along the given timeline, based on parameters of instantiation, and returns the resulting process.

See also:

```
SDE, SDE.source_dw, wiener_source, ornstein_uhlenbeck_SDE
```

Attributes

See SDE class documentation.

Methods

See SDE class documentation.

7.4 sdepy.hull_white_process

```
class sdepy.hull_white_process (paths=1, vshape=(), dtype=None, steps=None, i0=0, info=None, getinfo=True, method='euler', factors=1, x0=0., theta=0., k=1., sigma=1., dw=None, corr=None, rho=None)
```

F-factors Hull-White process (sum of F correlated mean-reverting Brownian motions).

Generates a process x(t) that solves the following SDE:

where $dw_i(t, dt)$ are standard Wiener process increments with correlations $dw_i(t, dt) * dw_j(t, dt) = corr(t)[i, j]$. x0, SDE parameters and dw(t, dt) should broadcast to vshape + (factors, paths).

Parameters

paths, vshape, dtype, steps, i0, info, getinfo, method See SDE class documentation.

x0 [array-like] Initial condition.

theta, k, sigma [array-like, or callable] SDE parameters.

dw, corr, rho Specification of stochasticity source of Wiener process increments. See SDE.source_dw documentation.

See also:

```
SDE, SDE.source_dw, wiener_source, hull_white_SDE, ornstein_uhlenbeck_process
```

Attributes

See SDE class documentation.

Methods

See SDE class documentation.

7.5 sdepy.hull_white_1factor_process

class sdepy.hull_white_lfactor_process (paths=1, vshape=(), dtype=None, steps=None, i0=0, info=None, getinfo=True, method='euler', x0=0., theta=0., k=1., sigma=1., dw=None, corr=None, rho=None)

1-factor Hull-White process (F=1 Hull-White process with F-index collapsed to a scalar). See hull_white_process class documentation.

See also:

hull_white_process, ornstein_uhlenbeck_process

Notes

Class added for naming convenience. Differs from a hull_white_process with factors=1 in that the last index of the process parameters has not been reserved to enumerate factors, and no factors parameter is present. Synonymous with ornstein_uhlenbeck_process.

Attributes

See SDE class documentation.

Methods

See SDE class documentation.

7.6 sdepy.cox_ingersoll_ross_process

```
class sdepy.cox_ingersoll_ross_process(paths=1, vshape=(), dtype=None, steps=None, i0=0, info=None, getinfo=True, method='euler', x0=1., theta=1., k=1., xi=1., dw=None, corr=None, rho=None)
```

Cox-Ingersoll-Ross mean reverting process.

Generates a process x(t) that solves the following SDE:

```
dx(t) = k(t) * (theta(t) - x(t)) * dt + xi(t) * sqrt(x(t)) * dw(t, dt)
```

where dw(t, dt) are standard Wiener process increments with correlation matrix specified by corr(t) or rho(t). x0, SDE parameters and dw(t, dt) should broadcast to vshape + (paths,). x0, theta, k should be positive.

Parameters

paths, vshape, dtype, steps, i0, info, getinfo, method See SDE class documentation.

x0 [array-like] Initial condition.

theta, k, xi [array-like, or callable] SDE parameters.

dw, corr, rho Specification of stochasticity source of Wiener process increments. See SDE.source_dw documentation.

Returns

x [process] Once instantiated as p, p(timeline) performs the integration along the given timeline, based on parameters of instantiation, and returns the resulting process.

See also:

```
SDE, SDE.source_dw, wiener_source, cox_ingersoll_ross_SDE
```

Attributes

See SDE class documentation.

Methods

See SDE class documentation.

7.7 sdepy.full_heston_process

```
class sdepy.full_heston_process (paths=1, vshape=(), dtype=None, steps=None, i0=0, info=None, getinfo=True, method='euler', x0=1., mu=0., sigma=1., y0=1., theta=1., k=1., xi=1., dw=None, corr=None, rho=None)
```

Heston stochastic volatility process (returns both process and volatility).

Generates processes x(t) and an y(t) that solve the following SDEs:

where, if N = vshape[-1] is the size of the last dimension of x(t), y(t), and dw(t, dt) are standard Wiener process increments with shape vshape + (2*N, paths):

```
dw(t)[..., i, :]*dw(t)[..., j, :] = corr(t)[..., i, j]*dt
dw_x(t) = dw(t)[..., :N, :],
dw_y(t) = dw(t)[..., N:, :],
```

x0 and SDE parameters should broadcast to vshape + (paths,). dw(t, dt) should broadcast to vshape[:-1] + (2*vshape[-1], paths). x0, y0, theta, k should be positive.

Parameters

paths, vshape, dtype, steps, i0, info, getinfo, method See SDE class documentation.

x0, y0 [array-like] Initial conditions for x (t) and y (t) processes respectively.

mu, sigma, theta, k, xi [array-like, or callable] SDE parameters.

dw, corr, rho Specification of stochasticity source of Wiener process increments. See SDE.source dw documentation.

Returns

x, y [processes] Once instantiated as p, p (timeline) performs the integration along the given timeline, based on parameters of instantiation, and returns the resulting processes.

See also:

```
SDE, SDE.source_dw, wiener_source, full_heston_SDE
```

Notes

x(t), y(t) are obtained via Euler-Maruyama numerical integration of the above SDE for y(t) and of the following equivalent SDE for a(t) = log(x(t)), handling negative values of y(t) via the full truncation algorithm [1]:

```
da(t) = (mu(t) - y(t) * sigma(t) * *2/2) * dt + sqrt(y(t)) * dw_x(t)
```

References

[1]

Attributes

See SDE class documentation.

Methods

See SDE class documentation.

7.8 sdepy.heston_process

```
class sdepy.heston_process (paths=1, vshape=(), dtype=None, steps=None, i0=0, info=None, getinfo=True, method='euler', x0=1., mu=0., sigma=1., y0=1., theta=1., k=1., xi=1., dw=None, corr=None, rho=None) Heston stochastic volatility process (stores and returns process only).
```

Generates a process as in full_heston_process (see its documentation), storing and returning the x(t) component only.

Parameters

paths, vshape, dtype, steps, i0, info, getinfo, method See SDE class documentation.

x0, mu, sigma, y0, theta, k, xi, dw, corr, rho See full_heston_process class documentation.

Returns

x [process] Once instantiated as p, p(timeline) performs the integration along the given timeline, based on parameters of instantiation, and returns the resulting process.

See also:

```
full_heston_process
```

Attributes

See SDE class documentation.

Methods

See SDE class documentation.

7.9 sdepy.jumpdiff_process

```
class sdepy.jumpdiff_process (paths=1, vshape=(), dtype=None, steps=None, i0=0, info=None, getinfo=True, method='euler', x0=1., mu=0., sigma=1., dw=None, corr=None, rho=None, dj=None, dn=None, ptype=int, lam=1., y=None)

Jump-diffusion process (lognormal process with compound Poisson logarithmic jumps).
```

Generates a process x(t) that solves the following SDE (see [1]):

```
dx(t) = mu(t)*x(t)*dt + sigma(t)*x(t)*dw(t, dt) + x(t)*dj(t, dt)
```

where dw(t, dt) are standard Wiener process increments with correlation matrix specified by corr(t) or rho(t), and dj(t, dt) are increments of a Poisson process with intensity lam(t), compounded with random variates distributed as exp(y(t)) - 1.

Parameters

paths, vshape, dtype, steps, i0, info, getinfo, method See SDE class documentation.

x0 [array-like] Initial condition.

mu, sigma [array-like, or callable] SDE parameters.

dw, corr, rho Specification of stochasticity source of Wiener process increments. See SDE.source_dw documentation.

dj, **dn**, **ptype**, **lam**, **y** Specification of stochasticity source of compound Poisson process increments. See SDE.source_dj documentation.

See also:

```
SDE, SDE.source_dw, SDE.source_dj, wiener_source, cpoisson_source, jumpdiff_SDE
```

Notes

The drift of the mean value $x_{mean}(t)$ of x(t) is mu(t) + nu(t), i.e. $dx_{mean}(t)/dt = x_{mean}(t)*(mu(t) + nu(t))$, where:

```
nu(t) = lam(t) * (y_exp_mean(t) - 1)
y_exp_mean(t) = average of exp(y(t))
```

x(t) is obtained via Euler-Maruyama numerical integration of the following equivalent SDE for a (t) = log(x(t)):

```
da(t) = (mu(t) - sigma(t) **2/2) *dt + x(t) *sigma(t) *dw(t, dt) + x(t) *dh(t, dt)
```

where dh(t, dt) are increments of a Poisson process with intensity lam(t) compounded with random variates distributed as y(t).

References

[1]

Attributes

See SDE class documentation.

Methods

See SDE class documentation.

7.10 sdepy.merton_jumpdiff_process

class sdepy.merton_jumpdiff_process (paths=1, vshape=(), dtype=None, steps=None, i0=0, info=None, getinfo=True, method='euler', x0=1., mu=0., sigma=1., dw=None, corr=None, rho=None, dj=None, dn=None, ptype=int, lam=1., a=0., b=1.)

Merton jump-diffusion process (jump-diffusion process with normal jump size distribution).

Same as $jumpdiff_process$, where the y parameter is specialized to $norm_rv(a, b)$, a normal variate with mean a(t) and standard deviation b(t).

See also:

jumpdiff_process, norm_rv

Attributes

See SDE class documentation.

Methods

See SDE class documentation.

7.11 sdepy.kou_jumpdiff_process

class sdepy.kou_jumpdiff_process (paths=1, vshape=(), dtype=None, steps=None, i0=0, info=None, getinfo=True, method='euler', x0=1., mu=0., sigma=1., dw=None, corr=None, rho=None, dj=None, dn=None, ptype=int, lam=1., a=0.5, b=0.5, pa=0.5)

Double exponential (Kou) jump-diffusion process (jump-diffusion process with double exponential jump size distribution).

Same as jumpdiff_process, where the y parameter is specialized to double_exp_rv(a, b, pa), a double exponential variate with scale a(t) with probability pa(t), and -b(t) with probability (1 - pa(t)).

See also:

```
jumpdiff_process, double_exp_rv
```

Attributes

See SDE class documentation.

Methods

See SDE class documentation.

7.12 sdepy.wiener_SDE

```
class sdepy.wiener_SDE(*, paths=1, vshape=(), dtype=None, steps=None, i0=0, info=None, get-info=True, method='euler', **args)
SDE for a Wiener process (Brownian motion) with drift.
```

See also:

wiener_process

Attributes

See SDE class documentation.

Methods

See SDE class documentation.

7.13 sdepy.lognorm_SDE

class sdepy.lognorm_SDE (*, paths=1, vshape=(), dtype=None, steps=None, i0=0, info=None, getinfo=True, method='euler', **args)

SDE for a lognormal process with drift.

See also:

lognorm_process

Attributes

See SDE class documentation.

Methods

See SDE class documentation.

7.14 sdepy.ornstein_uhlenbeck_SDE

```
class sdepy.ornstein\_uhlenbeck\_SDE(*, paths=1, vshape=(), dtype=None, steps=None, i0=0, info=None, getinfo=True, method='euler', **args)
```

SDE for an Ornstein-Uhlenbeck process.

See also:

ornstein_uhlenbeck_process

Attributes

See SDE class documentation.

Methods

See SDE class documentation.

7.15 sdepy.hull_white_SDE

class $sdepy.hull_white_SDE(*, paths=1, vshape=(), dtype=None, steps=None, i0=0, info=None, getinfo=True, method='euler', **args)$ SDE for an F-factors Hull White process.

See also:

hull_white_process

Attributes

See SDE class documentation.

Methods

See SDE class documentation.

7.16 sdepy.cox_ingersoll_ross_SDE

SDE for a Cox-Ingersoll-Ross mean reverting process.

See also:

cox_ingersoll_ross_process

Attributes

See SDE class documentation.

Methods

See SDE class documentation.

7.17 sdepy.full_heston_SDE

class $sdepy.full_heston_SDE(*, paths=1, vshape=(), dtype=None, steps=None, i0=0, info=None, getinfo=True, method='euler', **args)$ SDE for a Heston stochastic volatility process.

See also:

full_heston_process, heston_process

Attributes

See SDE class documentation.

Methods

See SDE class documentation.

7.18 sdepy.heston_SDE

See also:

heston_process, full_heston_process, full_heston_SDE

Attributes

See SDE class documentation.

Methods

See SDE class documentation.

7.19 sdepy.jumpdiff_SDE

class sdepy.jumpdiff_SDE (*, paths=1, vshape=(), dtype=None, steps=None, i0=0, info=None, getinfo=True, method='euler', **args)

SDE for a jump-diffusion process (lognormal process with compound Poisson logarithmic jumps).

See also:

jumpdiff_process

Attributes

See SDE class documentation.

Methods

See SDE class documentation.

7.20 sdepy.merton_jumpdiff_SDE

See also:

```
merton_jumpdiff_process, jumpdiff_SDE
```

Attributes

See SDE class documentation.

Methods

See SDE class documentation.

7.21 sdepy.kou_jumpdiff_SDE

 $\begin{array}{ll} \textbf{class} & \texttt{sdepy.kou_jumpdiff_SDE} \ (*, & paths=1, & vshape=(), & dtype=None, & steps=None, & i0=0, \\ & & info=None, & getinfo=True, & method='euler', & **args') \\ & & \texttt{SDE} \ for \ a \ double \ exponential} \ (\texttt{Kou}) \ jump-diffusion \ process. \\ \end{array}$

See also:

kou_jumpdiff_process, jumpdiff_SDE

Attributes

See SDE class documentation.

Methods

See SDE class documentation.

CHAPTER 8

Analytical Results

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independent parameters. Standard deviation of values at time t of a lognormal process (as per the lognorm_process class) with time-independent parameters. lognorm_pdf(t, x, *[, x0, mu, sigma]) Probability distribution function of values at time t of a lognormal process (as per the lognorm_process class) with time-independent parameters, evaluated at x.	10g110111 <u>1</u> 041(1, 10, 1111, 01g1111)	
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process (as per the lognorm_process class) with time-independent parameters. Probability distribution function of values at time t of a lognormal process (as per the lognorm_process class) with time-independent parameters, evaluated at x.	lognorm std(t *[v() mu sigma])	
independent parameters. Probability distribution function of values at time t of a lognormal process (as per the lognorm_process class) with time-independent parameters, evaluated at x.	i ognorn <u>e</u> sea(t, [, xo, mu, sigmu])	•
Probability distribution function of values at time t of a lognormal process (as per the lognorm_process class) with time-independent parameters, evaluated at x.		
of a lognormal process (as per the lognorm_process class) with time-independent parameters, evaluated at x.	lognorm ndf(t v *[v0 my sigma])	
class) with time-independent parameters, evaluated at x.	rognorm_par(i, x, ·[, xo, mu, sigma])	
X.		
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Table 1 – continue	ed from previous page
lognorm_cdf(t, x, *[, x0, mu, sigma])	Cumulative probability distribution function of val-
	ues at time t of a lognormal process (as per the log-
	norm_process class) with time-independent parame-
	ters, evaluated at x.
lognorm_log_chf(t, u, *[, x0, mu, sigma])	Characteristic function of the probability distribution
iognorm_rog_enr(t, u, [, xo, mu, sigmu])	of values at time t of the logarithm of a lognormal
	process (as per the lognorm_process class) with time-
	independent parameters, evaluated at u.
oruh_mean(t, *[, x0, theta, k, sigma])	Mean of values at time t of an Ornstein-Uhlenbeck
orum_mean(t, [, xo, theta, k, sigma])	process (as per the ornstein_uhlenbeck_process class)
	with time-independent parameters.
oruh_var(t, *[, x0, theta, k, sigma])	Variance of values at time t of an Ornstein-Uhlenbeck
orun_var(t, '[, xo, theta, k, signia])	
	process (as per the ornstein_uhlenbeck_process class)
7 (1/4) (0 (1 (4 (1 (1 (1 (1 (1 (1 (1 (1 (1 (1 (1 (1 (1	with time-independent parameters.
oruh_std(t, *[, x0, theta, k, sigma])	Standard deviation of values at time t of an
	Ornstein-Uhlenbeck process (as per the orn-
	stein_uhlenbeck_process class) with time-
1 15(4) 1 (1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	independent parameters.
oruh_pdf(t, x, *[, x0, theta, k, sigma])	Probability distribution function of values at time
	t of an Ornstein-Uhlenbeck process (as per the
	ornstein_uhlenbeck_process class) with time-
	independent parameters, evaluated at x.
oruh_cdf(t, x, *[, x0, theta, k, sigma])	Cumulative probability distribution function of val-
	ues at time t of an Ornstein-Uhlenbeck process (as
	per the ornstein_uhlenbeck_process class) with time-
	independent parameters, evaluated at x.
hw2f_mean(t, *[, x0, theta, k, sigma, rho])	Mean of values at time t of a Hull-White 2-factors pro-
	cess (as per the hull_white_process class) with time-
	independent parameters.
hw2f_var(t, *[, x0, theta, k, sigma, rho])	Variance of values at time t of a Hull-White 2-factors
	process (as per the hull_white_process class) with
	time-independent parameters.
$hw2f_std(t, *[, x0, theta, k, sigma, rho])$	Standard deviation of values at time t of a Hull-White
	2-factors process (as per the hull_white_process
	class) with time-independent parameters.
$hw2f_pdf(t, x, *[, x0, theta, k, sigma, rho])$	Probability distribution function of values at time
	t of a Hull-White 2-factors process (as per the
	hull_white_process class) with time-independent pa-
	rameters, evaluated at x.
$hw2f_cdf(**args)$	Cumulative probability distribution function of values
	at time t of a Hull-White 2-factors process (as per the
	hull_white_process class) with time-independent pa-
	rameters, evaluated at x.
$cir_{mean}(t, *[, x0, theta, k, xi])$	Mean of values at time t of a Cox-Ingersoll-Ross
	process (as per the cox_ingersoll_ross_process class)
	with time-independent parameters.
cir_var(t, *[, x0, theta, k, xi])	Variance of values at time t of a Cox-Ingersoll-Ross
	process (as per the cox_ingersoll_ross_process class)
	with time-independent parameters.
$cir_std(t, *[, x0, theta, k, xi])$	Standard deviation of values at time t of
	a Cox-Ingersoll-Ross process (as per the
	cox_ingersoll_ross_process class) with time-
	· ·
	independent parameters.

Continued on next page

Table 1 – continued	from previous page
$cir_pdf(t, x, *[, x0, theta, k, xi])$	Probability distribution function of values at
	time t of a Cox-Ingersoll-Ross process (as per
	the cox_ingersoll_ross_process class) with time-
	independent parameters, evaluated at x.
$heston_log_mean(t, *[, x0, mu, sigma, y0,])$	Mean of the logarithm of values at time t of a Hes-
	ton process (as per the full_heston_process class) with
	time-independent parameters.
heston_log_var(**args)	Variance of the logarithm of values at time t of a Hes-
	ton process (as per the full_heston_process class) with
	time-independent parameters.
$\textit{heston_log_std}(t, *[, x0, mu, sigma, y0,])$	Standard deviation of the logarithm of values at time
	t of a Heston process (as per the full_heston_process
	class) with time-independent parameters.
$heston_log_pdf(t, logx, *[, x0, mu, sigma,])$	Probability distribution function of values at time t
	of the logarithm of a Heston process, (as per the
	full_heston_process class) with time-independent pa-
	rameters, evaluated at logx.
$\textcolor{red}{heston_log_chf(t, u, *[, x0, mu, sigma, y0,])}$	Characteristic function of the probability distribution
	of values at time t of the logarithm of a Heston pro-
	cess (as per the full_heston_process class), with time-
	independent parameters, evaluated at u.
$mjd_log_pdf(t, logx, *[, x0, mu, sigma,])$	Probability distribution function of values at time t of
	the logarithm of a Merton jump-diffusion process (as
	per the merton_jumpdiff_process class), with time-
	independent parameters, evaluated at logx.
mjd_log_chf(t, u, *[, x0, mu, sigma, lam, a, b])	Characteristic function of the probability distribu-
	tion of values at time t of the logarithm of a
	Merton jump-diffusion process (as per the mer-
	ton_jumpdiff_process class), with time-independent
7 (4 × F 0 · · · · · · · · · · · · · · · · · ·	parameters, evaluated at u.
kou_mean(t, *[, x0, mu, sigma, lam, a, b, pa])	Mean of values at time t of a double expo-
	nential (Kou) jump-diffusion process (as per the
	kou_jumpdiff_process class) with time-independent
1	parameters.
$kou_log_pdf(t, logx, *[, x0, mu, sigma,])$	Probability distribution function of values at time t of
	the logarithm of a double-exponential (Kou) jump-
	diffusion process (as per the kou_jumpdiff_process class), with time-independent parameters, evaluated
$kou_log_chf(t, u, *[, x0, mu, sigma, lam,])$	at logx. Characteristic function of the probability distribution
KOU_109_CH1(t, u, 1[, x0, mu, sigma, iam,])	of values at time t of the logarithm of a Kou jump-
	diffusion process, (as per the kou_jumpdiff_process
	class) with time-independent parameters, evaluated at
	u.
bsd1d2(k, t, *[, x0, r, q, sigma])	Black-Scholes d1 and d2 coefficients.
$\frac{bscall(k, t, *[, x0, t, q, sigma])}{bscall(k, t, *[, x0, t, q, sigma])}$	Black-Scholes call option value.
bscall_delta(k, t, *[, x0, r, q, sigma])	Black-Scholes call option delta.
bsput(k, t, *[, x0, r, q, sigma])	Black-Scholes put option value.
$\frac{bsput(k, t, [t, x_0, t, q, sigma])}{bsput_delta(k, t, *[t, x_0, t, q, sigma])}$	Black-Scholes put option delta.
zopac_actea(k, t, [, No, 1, q, sigina])	Diack Scholes put option dela.

8.1 sdepy.wiener_mean

class $sdepy.wiener_mean(t, *, x0=0., mu=0., sigma=1.)$

Mean of values at time t of a Wiener process (as per the wiener_process class) with time-independent parameters.

See also: wiener_process **Attributes** params **Methods** call 8.2 sdepy.wiener_var class $sdepy.wiener_var(t, *, x0=0., mu=0., sigma=1.)$ Variance of values at time t of a Wiener process (as per the wiener_process class) with time-independent parameters. See also: wiener_process **Attributes** params **Methods** call 8.3 sdepy.wiener_std class sdepy.wiener_std(t, *, x0=0., mu=0., sigma=1.) Standard deviation of values at time t of a Wiener process (as per the wiener_process class) with timeindependent parameters. See also: wiener_process Attributes params

call

Methods

8.4 sdepy.wiener_pdf

```
class sdepy.wiener_pdf(t, x, *, x0=0., mu=0., sigma=1.)
```

Probability distribution function of values at time t of a Wiener process (as per the wiener_process class) with time-independent parameters, evaluated at x.

See also:

wiener_process

Attributes

params

Methods

__call__

8.5 sdepy.wiener cdf

```
class sdepy.wiener_cdf(t, x, *, x0=0., mu=0., sigma=1.)
```

Cumulative probability distribution function of values at time t of a Wiener process (as per the wiener_process class) with time-independent parameters, evaluated at x.

See also:

wiener_process

Attributes

params

Methods

__call__

8.6 sdepy.wiener_chf

```
class sdepy.wiener_chf(t, u, *, x0=0., mu=0., sigma=1.)
```

Characteristic function of the probability distribution of values at time t of a Wiener process (as per the wiener_process class) with time-independent parameters, evaluated at u.

See also:

wiener_process

Attributes

params

Methods

8.7 sdepy.lognorm_mean

```
class sdepy.lognorm_mean(t, *, x0=1., mu=0., sigma=1.)
```

Mean of values at time t of a lognormal process (as per the lognorm_process class) with time-independent parameters.

See also:

lognorm_process

Attributes

params

Methods

__call__

8.8 sdepy.lognorm var

class sdepy.lognorm_var(t, *, x0=1., mu=0., sigma=1.)

Variance of values at time t of a lognormal process (as per the lognorm_process class) with time-independent parameters.

See also:

lognorm_process

Attributes

params

Methods

__call__

8.9 sdepy.lognorm_std

class $sdepy.lognorm_std(t, *, x0=1., mu=0., sigma=1.)$

Standard deviation of values at time t of a lognormal process (as per the lognorm_process class) with time-independent parameters.

See also:

lognorm_process

Attributes

params

Methods

8.10 sdepy.lognorm_pdf

class sdepy.lognorm_pdf(t, x, *, x0=1., mu=0., sigma=1.)

Probability distribution function of values at time t of a lognormal process (as per the lognorm_process class) with time-independent parameters, evaluated at x.

See also:

lognorm_process

Attributes

params

Methods

__call__

8.11 sdepy.lognorm cdf

class $sdepy.lognorm_cdf(t, x, *, x0=1., mu=0., sigma=1.)$

Cumulative probability distribution function of values at time t of a lognormal process (as per the lognorm_process class) with time-independent parameters, evaluated at x.

See also:

lognorm_process

Attributes

params

Methods

__call__

8.12 sdepy.lognorm_log_chf

 $\verb"class" sdepy.lognorm_log_chf" (t, u, *, x0=1., mu=0., sigma=1.)$

Characteristic function of the probability distribution of values at time t of the logarithm of a lognormal process (as per the lognorm_process class) with time-independent parameters, evaluated at u.

See also:

lognorm_process

Attributes

params

Methods

8.13 sdepy.oruh_mean

```
class sdepy.oruh_mean(t, *, x0=0., theta=0., k=1., sigma=1.)
```

Mean of values at time t of an Ornstein-Uhlenbeck process (as per the ornstein_uhlenbeck_process class) with time-independent parameters.

See also:

ornstein_uhlenbeck_process

Attributes

params

Methods

__call__

8.14 sdepy.oruh var

```
class sdepy.oruh_var(t, *, x0=0., theta=0., k=1., sigma=1.)
```

Variance of values at time t of an Ornstein-Uhlenbeck process (as per the ornstein_uhlenbeck_process class) with time-independent parameters.

See also:

ornstein_uhlenbeck_process

Attributes

params

Methods

__call__

8.15 sdepy.oruh_std

```
class sdepy.oruh_std(t, *, x0=0., theta=0., k=1., sigma=1.)
```

Standard deviation of values at time t of an Ornstein-Uhlenbeck process (as per the ornstein_uhlenbeck_process class) with time-independent parameters.

See also:

ornstein_uhlenbeck_process

Attributes

params

Methods

8.16 sdepy.oruh_pdf

class sdepy.oruh_pdf(t, x, *, x0=0., theta=0., k=1., sigma=1.)

Probability distribution function of values at time t of an Ornstein-Uhlenbeck process (as per the ornstein_uhlenbeck_process class) with time-independent parameters, evaluated at x.

See also

ornstein_uhlenbeck_process

Attributes

params

Methods

__call__

8.17 sdepy.oruh cdf

class sdepy.oruh_cdf(t, x, *, x0=0., theta=0., k=1., sigma=1.)

Cumulative probability distribution function of values at time t of an Ornstein-Uhlenbeck process (as per the ornstein_uhlenbeck_process class) with time-independent parameters, evaluated at x.

See also:

ornstein_uhlenbeck_process

Attributes

params

Methods

_call__

8.18 sdepy.hw2f_mean

class $sdepy.hw2f_mean(t, *, x0=(0., 0.), theta=(0., 0.), k=(1., 1.), sigma=(1., 1.), rho=0.)$

Mean of values at time t of a Hull-White 2-factors process (as per the hull_white_process class) with time-independent parameters.

See also:

hull_white_process

Attributes

params

Methods

8.19 sdepy.hw2f_var

class sdepy.**hw2f_var**(t, *, x0=(0, 0.), theta=(0, 0.), k=(1, 1.), sigma=(1, 1.), rho=0.)

Variance of values at time t of a Hull-White 2-factors process (as per the hull_white_process class) with time-independent parameters.

See also:

```
hull_white_process
```

Attributes

params

Methods

__call__

8.20 sdepy.hw2f std

 $\textbf{class} \ \, \texttt{sdepy.hw2f_std}\,(t,\, *, x0 = (0.,\, 0.), \, theta = (0.,\, 0.), \, k = (1.,\, 1.), \, sigma = (1.,\, 1.), \, rho = 0.)$

Standard deviation of values at time t of a Hull-White 2-factors process (as per the hull_white_process class) with time-independent parameters.

See also:

```
hull_white_process
```

Attributes

params

Methods

_call__

8.21 sdepy.hw2f_pdf

class $sdepy.hw2f_pdf(t, x, *, x0=(0., 0.), theta=(0., 0.), k=(1., 1.), sigma=(1., 1.), rho=0.)$

Probability distribution function of values at time t of a Hull-White 2-factors process (as per the hull_white_process class) with time-independent parameters, evaluated at x.

See also:

hull_white_process

Attributes

params

Methods

8.22 sdepy.hw2f_cdf

```
class sdepy.hw2f_cdf(**args)
```

Cumulative probability distribution function of values at time t of a Hull-White 2-factors process (as per the hull_white_process class) with time-independent parameters, evaluated at x.

See also:

```
hull_white_process
```

Attributes

params

Methods

__call__

8.23 sdepy.cir mean

```
class sdepy.cir_mean(t, *, x0=1., theta=1., k=1., xi=1.)
```

Mean of values at time t of a Cox-Ingersoll-Ross process (as per the cox_ingersoll_ross_process class) with time-independent parameters.

See also:

```
cox_ingersoll_ross_process
```

Attributes

params

Methods

__call__

8.24 sdepy.cir_var

```
class sdepy.cir_var(t, *, x0=1., theta=1., k=1., xi=1.)
```

Variance of values at time t of a Cox-Ingersoll-Ross process (as per the cox_ingersoll_ross_process class) with time-independent parameters.

See also:

```
cox_ingersoll_ross_process
```

Attributes

params

Methods

8.25 sdepy.cir_std

```
class sdepy.cir_std(t, *, x0=1., theta=1., k=1., xi=1.)
```

Standard deviation of values at time t of a Cox-Ingersoll-Ross process (as per the cox_ingersoll_ross_process class) with time-independent parameters.

See also

```
cox_ingersoll_ross_process
```

Attributes

params

Methods

__call__

8.26 sdepy.cir pdf

```
class sdepy.cir_pdf (t, x, *, x0=1., theta=1., k=1., xi=1.)
```

Probability distribution function of values at time t of a Cox-Ingersoll-Ross process (as per the cox_ingersoll_ross_process class) with time-independent parameters, evaluated at x.

See also:

```
cox_ingersoll_ross_process
```

Attributes

params

Methods

__call__

8.27 sdepy.heston_log_mean

```
class sdepy.heston_log_mean(t, *, x0=1., mu=0., sigma=1., y0=1., theta=1., k=1., xi=1., rho=0)
```

Mean of the logarithm of values at time t of a Heston process (as per the full_heston_process class) with time-independent parameters.

See also:

```
full_heston_process
```

Attributes

params

Methods

__call__

8.28 sdepy.heston_log_var

class sdepy.heston_log_var(**args)

Variance of the logarithm of values at time t of a Heston process (as per the full_heston_process class) with time-independent parameters.

See also:

full_heston_process

Attributes

params

Methods

_call__

8.29 sdepy.heston_log_std

class sdepy.heston_log_std(t, *, x0=1., mu=0., sigma=1., y0=1., theta=1., k=1., xi=1., rho=0.)

Standard deviation of the logarithm of values at time t of a Heston process (as per the full_heston_process class) with time-independent parameters.

See also:

full_heston_process

Attributes

params

Methods

__call__

8.30 sdepy.heston_log_pdf

class sdepy.heston_log_pdf(t, logx, *, x0=1., mu=0., sigma=1., y0=1., theta=1., k=1., xi=1., rho=0.)

Probability distribution function of values at time t of the logarithm of a Heston process, (as per the full_heston_process class) with time-independent parameters, evaluated at logx.

See also:

full_heston_process

Notes

Estimate by numerical integration, using scipy.integrate.quad, of the closed-form characteristic function heston_log_chf. Integration errors are not reported/checked. Either t or logx must be a scalar.

Attributes

params

Methods

_call__

8.31 sdepy.heston_log_chf

class sdepy.heston_log_chf(t, u, *, x0=1., mu=0., sigma=1., y0=1., theta=1., k=1., xi=1., rho=0.)

Characteristic function of the probability distribution of values at time t of the logarithm of a Heston process (as per the full_heston_process class), with time-independent parameters, evaluated at u.

See also:

full_heston_process

Attributes

params

Methods

_call__

8.32 sdepy.mjd_log_pdf

class $sdepy.mjd_log_pdf(t, logx, *, x0=1., mu=0., sigma=1., lam=1., a=0.0, b=1.)$

Probability distribution function of values at time t of the logarithm of a Merton jump-diffusion process (as per the merton_jumpdiff_process class), with time-independent parameters, evaluated at logx.

See also:

 $jumpdiff_process, merton_jumpdiff_process, mjd_log_chf$

Notes

Estimate by numerical integration, using scipy.integrate.quad, of the closed-form characteristic function mjd_log_chf. Integration errors are not reported/checked. Either t or logx must be a scalar.

Attributes

params

Methods

__call__

8.33 sdepy.mjd_log_chf

class sdepy.**mjd_log_chf** (t, u, *, x0=1., mu=0., sigma=1., lam=1., a=0.0, b=1.)

Characteristic function of the probability distribution of values at time t of the logarithm of a Merton jumpdiffusion process (as per the merton_jumpdiff_process class), with time-independent parameters, evaluated at u.

See also:

jumpdiff_process, merton_jumpdiff_process

Attributes

params

Methods

__call__

8.34 sdepy.kou mean

class sdepy.**kou_mean** (t, *, x0=1., mu=0., sigma=1., lam=1., a=0.5, b=0.5, pa=0.5)

Mean of values at time t of a double exponential (Kou) jump-diffusion process (as per the kou_jumpdiff_process class) with time-independent parameters.

See also:

jumpdiff_process, kou_jumpdiff_process

Attributes

params

Methods

__call__

8.35 sdepy.kou_log_pdf

class sdepy.kou_log_pdf(t, logx, *, x0=1., mu=0., sigma=1., lam=1., pa=0.5, a=0.5, b=0.5)

Probability distribution function of values at time t of the logarithm of a double-exponential (Kou) jump-diffusion process (as per the kou_jumpdiff_process class), with time-independent parameters, evaluated at logx.

See also:

jumpdiff_process, kou_jumpdiff_process, kou_log_chf

Notes

Estimate by numerical integration, using scipy.integrate.quad, of the closed-form characteristic function kou_log_chf. Integration errors are not reported/checked. Either t or logx must be a scalar.

Attributes

params

Methods

_call__

8.36 sdepy.kou_log_chf

class sdepy.**kou_log_chf** (t, u, *, x0=1., mu=0., sigma=1., lam=1., a=0.5, b=0.5, pa=0.5)

Characteristic function of the probability distribution of values at time t of the logarithm of a Kou jump-diffusion process, (as per the kou_jumpdiff_process class) with time-independent parameters, evaluated at u.

See also:

```
jumpdiff_process, kou_jumpdiff_process
```

Attributes

params

Methods

__call__

8.37 sdepy.bsd1d2

```
class sdepy.bsd1d2 (k, t, *, x0=1., r=0., q=0., sigma=1.) Black-Scholes d1 and d2 coefficients.
```

See also:

bscall

Attributes

params

Methods

call

8.38 sdepy.bscall

```
class sdepy.bscall (k, t, *, x0=1., r=0., q=0., sigma=1.) Black-Scholes call option value.
```

Parameters

- k [array-like] Strike.
- t [array-like] Time to maturity.
- **x0** [array-like] Initial value of underlying security.
- r [array-like] Risk-free rate.
- q [array-like] Dividend yield of underlying security.
- sigma [array-like] Volatility of underlying security.

Returns

array Risk neutral valuation at time s=0 of an European call option paying max(x(t) - k, 0) at maturity, where the price x(s) of the underlying security follows a lognormal process with x(0) = x0 and volatility sigma.

See also:

```
bsd1d2, bscall_delta, bsput, bsput_delta
```

Notes

bscall(k, t, x0, r, q, sigma) returns:

```
bscall_value = x0*exp(-q*t)*norm.cdf(d1) + k*exp(-r*t)*norm.cdf(d2)
```

where cdf is scipy.stats.norm.cdf and d1, d2 = bsd1d2(k, t, x0, r, q, sigma) are given as:

Attributes

params

Methods

_call__

8.39 sdepy.bscall_delta

```
sdepy.bscall_delta (k, t, *, x0=1., r=0., q=0., sigma=1.) Black-Scholes call option delta.
```

See also:

bscall

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8.40 sdepy.bsput

```
class sdepy.bsput (k, t, *, x0=1., r=0., q=0., sigma=1.)
Black-Scholes put option value.

See also:
bscall

Attributes
params

Methods
```

8.41 sdepy.bsput_delta

Methods

```
class sdepy.bsput_delta (k, t, *, x0=1., r=0., q=0., sigma=1.)
Black-Scholes put option delta.
See also:

bscall

Attributes

params
```

CHAPTER 9

Shortcuts

Stochasticity sources and preset processes may be addressed using the following shortcuts:

Full name	Shortcut
wiener_source	dw
poisson_source	dn
cpoisson_source	dj
odd_wiener_source	odd_dw
even_poisson_source	even_dn
even_cpoisson_source	even_dj
true_wiener_source	true_dw
true_poisson_source	true_dn
true_cpoisson_source	true_dj
wiener_process	wiener
lognorm_process	lognorm
ornstein_uhlenbeck_process	oruh
hull_white_process	hwff
hull_white_1factor_process	hw1f
cox_ingersoll_ross_process	cir
full_heston_process	heston_xy
heston_process	heston
jumpdiff_process	jumpdiff
merton_jumpdiff_process	mjd
kou_jumpdiff_process	kou

Shortcuts have been wrapped as "kfuncs", objects with managed keyword arguments (see kfunc decorator documentation below).

Analytical results are named according to the shortcut of the corresponding process (e.g. lognorm_mean, lognorm_cdf etc. from the lognorm shortcut) and are wrapped as kfuncs as well.

kfunc([f, nvar])	Decorator to wrap classes or functions as objects with
	managed keyword arguments.
iskfunc(cls_or_object)	Tests if the given class or instance has been wrapped
	as a kfunc.

9.1 sdepy.kfunc

```
sdepy.kfunc(f=None, *, nvar=None)
```

Decorator to wrap classes or functions as objects with managed keyword arguments.

This decorator, intended as an aid to interactive and notebook sessions, wraps a callable, class or function, as a "kfunc" object that handles separately its parameters (keyword-only), whose values are stored in the object, and its variables (positional or keyword), always provided upon evaluation.

Syntax:

```
@kfunc
class my_class:
    def __init___(self, **kwparams):
        ...
    def __call__(self, *var, **kwvar):
        ...
@kfunc(nvar=k)
def my_function(*var, **kwargs):
        ...
```

After decoration, my_class is a kfunc with kwparams as parameters, and with var and kwvar as variables, and my_function is a kfunc with the first k of var, kwargs as variables, and the remaining kwargs as parameters. For usage, see examples below.

Examples

Wrap wiener_source into a kfunc, named dw:

```
>>> from sdepy import wiener_source, kfunc
>>> dw = kfunc(wiener_source)
```

Instantiate dw and evaluate it (this is business as usual):

```
>>> my_instance = dw(paths=100, dtype=np.float32)
>>> x = my_instance(t=0, dt=1)
>>> x.shape, x.dtype
((100,), dtype('float32'))
```

Inspect kfunc parameters stored in my_instance:

```
>>> my_instance.params
{'paths': 100, 'vshape': (), 'dtype': <class 'numpy.float32'>, 'corr':_

None, 'rho': None}
```

Evaluate my_instance changing some parameters (call the instance with one or more):

```
>>> x = my_instance(t=0, dt=1, paths=999)
>>> x.shape, x.dtype
((999,), dtype('float32'))
```

Parameters stored in my_instance are not affected:

```
>>> my_instance.paths == my_instance.params['paths'] == 100
True
```

Create a new instance, changing some parameters and keeping those already set (call the instance without passing any variables):

```
>>> new_instance = my_instance(vshape=2, rho=0.5)
>>> new_instance.params
{'paths': 100, 'vshape': 2, 'dtype': <class 'numpy.float32'>, 'corr':_

None, 'rho': 0.5}
```

Instantiate and evaluate at once (pass one or more variables to the class constructor):

```
>>> x = dw(0, 1, paths=100, dtype=np.float32)
>>> x.shape, x.dtype
((100,), dtype('float32'))
```

As long as variables are passed by name, order doesn't matter (omitted variables take default values, if any):

```
>>> x = dw(paths=100, dtype=np.float32, dt=1, t=0)
>>> x.shape, x.dtype
((100,), dtype('float32'))
```

Attributes

params [dictionary] Parameter values stored in the instance (read-only). For wrapped SDE subclasses, also includes default values of all SDE-specific parameters, as stored in the args attribute.

9.2 sdepy.iskfunc

```
sdepy.iskfunc(cls_or_object)
```

Tests if the given class or instance has been wrapped as a kfunc.

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