# **py-pde Documentation**

Release unknown

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# CONTENTS

1	Getti	ng started 3
	1.1	When (not) to use the package
	1.2	Installation
		1.2.1         Install using pip         3
		1.2.2 Install using conda
		1.2.3 Install from source
		Required prerequisites
		Optional packages
		Downloading <i>py-pde</i>
	1.3	Package overview
2	Exan	nples 7
-	2.1	Plotting a vector field
	2.2	Solving Laplace's equation in 2d
	2.3	Plotting a scalar field in cylindrical coordinates
	2.4	Solving Poisson's equation in 1d       10
	2.5	Simple diffusion equation
	2.6	Kuramoto-Sivashinsky - Using PDE class    12
	2.7	Spherically symmetric PDE       13
	2.8	Diffusion on a Cartesian grid
	2.9	Stochastic simulation       16
	2.10	Time-dependent boundary conditions
	2.11	Setting boundary conditions
	2.12	1D problem - Using PDE class   19
	2.13	Heterogeneous boundary conditions
	2.14	Brusselator - Using the <i>PDE</i> class
	2.15	Writing and reading trajectory data
	2.16	Diffusion equation with spatial dependence
	2.17	Using simulation trackers
	2.18	Schrödinger's Equation
	2.19	Kuramoto-Sivashinsky - Using custom class
	2.20	Custom Class for coupled PDEs
	2.21	1D problem - Using custom class
	2.22	Visualizing a scalar field
	2.23	Kuramoto-Sivashinsky - Compiled methods
	2.24	Solver comparison
	2.25	Custom PDE class: SIR model
	2.26	Brusselator - Using custom class
3	User	manual 41

i

	3.1	Mathema	ical basics	41
		3.1.1 <b>(</b>	Curvilinear coordinates	41
		Po	lar coordinates	41
		Sp	herical coordinates	41
		C		42
		-		42
				43
	3.2			43
				43
				43
			6	44
				44
				 44
	3.3			45
	5.5		6	45
				+5 46
			1	40 47
			1	48 48
			1	48
				49
			1	49
			1	49 7 9
				50
				51
	3.4			52
			01	52
				53
				54
	3.5	Contribut		55
		3.5.1 \$	1 0	55
		3.5.2 H	Extending functionality	55
		3.5.3 I	Design choices	55
		3.5.4 (	Coding style	55
		3.5.5 H	Running unit tests	56
	3.6	Citing the	package	56
	3.7	-		57
				57
				57
		3.7.3 (	Dur Responsibilities	57
			1	57
				58
				58
		5.7.0 1		00
4	Refer	ence man	Jal	59
	4.1	pde.fields	package	59
		1		60
				74
		· · ·		80
				85
				89
	4.2	1		94
				95
		-		95
				95 96
		D		-0

		pde.grids.boundaries.axes module
		pde.grids.boundaries.axis module
		pde.grids.boundaries.local module
	4.2.2	pde.grids.coordinates package
		pde.grids.coordinates.base module
		pde.grids.coordinates.bipolar module
		pde.grids.coordinates.bispherical module
		pde.grids.coordinates.cartesian module
		pde.grids.coordinates.cylindrical module
		pde.grids.coordinates.polar module
		pde.grids.coordinates.spherical module
	4.2.3	pde.grids.operators package
		pde.grids.operators.cartesian module
		pde.grids.operators.common module
		pde.grids.operators.cylindrical_sym module
		pde.grids.operators.polar_sym module
		pde.grids.operators.spherical_sym module
	4.2.4	pde.grids.base module
	4.2.5	pde.grids.cartesian module
	4.2.6	pde.grids.cylindrical module
	4.2.7	pde.grids.spherical module
4.3	pde.pd	les package
	4.3.1	pde.pdes.allen_cahn module
	4.3.2	pde.pdes.base module
	4.3.3	pde.pdes.cahn_hilliard module
	4.3.4	pde.pdes.diffusion module
	4.3.5	pde.pdes.kpz_interface module
	4.3.6	pde.pdes.kuramoto_sivashinsky module
	4.3.7	pde.pdes.laplace module
	4.3.8	pde.pdes.pde module
	4.3.9	pde.pdes.swift_hohenberg module
	4.3.10	
4.4		lvers package
4.4	4.4.1	pde.solvers.adams_bashforth module
	4.4.2	pde.solvers.base module
	4.4.2	pde.solvers.controller module
	4.4.3	pde.solvers.crank_nicolson module
		pde.solvers.explicit module
	4.4.5	
	4.4.7	pde.solvers.explicit_mpi module
	4.4.7	I I I I I I I I I I I I I I I I I I I
15		1 12
4.5	*	orage package
	4.5.1	pde.storage.base module
	4.5.2	pde.storage.file module
	4.5.3	pde.storage.memory module
1.6	4.5.4	pde.storage.movie module
4.6		ols package
	4.6.1	pde.tools.cache module
	4.6.2	pde.tools.config module
	4.6.3	pde.tools.cuboid module
	4.6.4	pde.tools.docstrings module
	4.6.5	pde.tools.expressions module
	4.6.6	pde.tools.ffmpeg module
	4.6.7	pde.tools.math module

Python 1	Module I	ndex	271
	4.8.2	pde.visualization.plotting module	264
	4.8.1	pde.visualization.movies module	
4.8	pde.vist	alization package	
	4.7.4	pde.trackers.trackers module	253
	4.7.3	pde.trackers.interrupts module	249
	4.7.2	pde.trackers.interactive module	247
	4.7.1	pde.trackers.base module	244
4.7	pde.trac	kers package	243
	4.6.16	pde.tools.typing module	243
	4.6.15	pde.tools.spectral module	242
	4.6.14	pde.tools.plotting module	237
	4.6.13	pde.tools.parse_duration module	237
	4.6.12	pde.tools.parameters module	234
	4.6.11	pde.tools.output module	233
	4.6.10	pde.tools.numba module	231
	4.6.9	pde.tools.mpi module	229
	4.6.8	pde.tools.misc module	226

Index

273

The *py-pde* python package provides methods and classes useful for solving partial differential equations (PDEs) of the form

$$\partial_t u(\boldsymbol{x},t) = \mathcal{D}[u(\boldsymbol{x},t)] + \eta(u,\boldsymbol{x},t) ,$$

where  $\mathcal{D}$  is a (non-linear) operator containing spatial derivatives that defines the time evolution of a (set of) physical fields u with possibly tensorial character, which depend on spatial coordinates x and time t. The framework also supports stochastic differential equations in the Itô representation, where the noise is represented by  $\eta$  above.



The main audience for the package are researchers and students who want to investigate

the behavior of a PDE and get an intuitive understanding of the role of the different terms and the boundary conditions. To support this, *py-pde* evaluates PDEs using the methods of lines with a finite-difference approximation of the differential operators. Consequently, the mathematical operator  $\mathcal{D}$  can be naturally translated to a function evaluating the evolution rate of the PDE.

#### Contents

#### CHAPTER

### **GETTING STARTED**

### 1.1 When (not) to use the package

*py-pde* provides a straight-forward way to simulate partial differential equations (PDEs) using a finite-difference scheme. Advantages of this approach include:

- Supports non-linear PDEs with complex boundary conditions.
- Direct specification of the evolution equations using a syntax that is similar to the underlying mathematical equations.
- Supports collections of multiple fields of vectorial or tensorial character.

However, there are of course also disadvantages to the approach taken by *py-pde*:

- Only suited for simple geometries (like rectangles, disks, or cylinders) with a fixed discretization.
- Optimized for PDEs that describe the time-evolution of a physical system. Neither time-independent systems nor integro-differential equation are fully supported.
- Finite-differences can lead to numerical instabilities.
- Provided generic solvers might not be most suitable choice for particular equations.

# 1.2 Installation

The *py-pde* package is developed for python 3.9 and has been tested up to version 3.12 under Linux, Windows, and macOS. Before you can start using the package, you need to install it using one of the following methods.

#### 1.2.1 Install using pip

The package is available on pypi, so you should be able to install it by running

```
pip install py-pde
```

In order to have all features of the package available, you might also want to install the following optional packages:

```
pip install h5py pandas pyfftw tqdm
```

Moreover, **ffmpeg** needs to be installed and for creating movies.

#### 1.2.2 Install using conda

The *py-pde* package is also available on conda using the *conda-forge* channel. You can thus install it using

conda install -c conda-forge py-pde

This installation includes many dependencies to have most features of py-pde.

#### 1.2.3 Install from source

Installing from source can be necessary if the pypi installation does not work or if the latest source code should be installed from github.

#### **Required prerequisites**

The code builds on other python packages, which need to be installed for *py-pde* to function properly. The required packages are listed in the table below:

Package	Minimal version	Usage
matplotlib	3.1	Visualizing results
numba	0.59	Just-in-time compilation to accelerate numerics
numpy	1.22	Handling numerical data
scipy	1.10	Miscellaneous scientific functions
sympy	1.9	Dealing with user-defined mathematical expressions
tqdm	4.66	Display progress bars during calculations

The simplest way to install these packages is to use the requirements.txt in the base folder:

```
pip install -r requirements.txt
```

Alternatively, these package can be installed via your operating system's package manager, e.g. using **macports**, **homebrew**, or **conda**. The package versions given above are minimal requirements, although this is not tested systematically. Generally, it should help to install the latest version of the package.

#### **Optional packages**

The following packages should be installed to use some miscellaneous features:

Package	Minimal version	Usage
ffmpeg-python	0.2	Reading and writing videos
h5py	2.10	Storing data in the hierarchical file format
ipywidgets	8	Jupyter notebook support
mpi4py	3	Parallel processing using MPI
napari	0.4.8	Displaying images interactively
numba-mpi	0.22	Parallel processing using MPI+numba
pandas	2	Handling tabular data
pyfftw	0.12	Faster Fourier transforms
rocket-fft	0.2.4	Numba-compiled fast Fourier transforms

For making movies, the **ffmpeg** should be available. Additional packages might be required for running the tests in the folder tests and to build the documentation in the folder docs. These packages are listed in the files requirements.txt in the respective folders.

#### Downloading py-pde

The package can be simply checked out from github.com/zwicker-group/py-pde. To import the package from any python session, it might be convenient to include the root folder of the package into the PYTHONPATH environment variable.

This documentation can be built by calling the **make html** in the docs folder. The final documentation will be available in docs/build/html. Note that a LaTeX documentation can be build using **make latexpdf**.

#### 1.3 Package overview

The main aim of the *pde* package is to simulate partial differential equations in simple geometries. Here, the time evolution of a PDE is determined using the method of lines by explicitly discretizing space using fixed grids. The differential operators are implemented using the finite difference method. For simplicity, we consider only regular, orthogonal grids, where each axis has a uniform discretization and all axes are (locally) orthogonal. Currently, we support simulations on *CartesianGrid*, *PolarSymGrid*, *SphericalSymGrid*, and *CylindricalSymGrid*, with and without periodic boundaries where applicable.

Fields are defined by specifying values at the grid points using the classes *ScalarField*, *VectorField*, and *Ten-sor2Field*. These classes provide methods for applying differential operators to the fields, e.g., the result of applying the Laplacian to a scalar field is returned by calling the method *laplace()*, which returns another instance of *ScalarField*, whereas *gradient()* returns a *VectorField*. Combining these functions with ordinary arithmetics on fields allows to represent the right hand side of many partial differential equations that appear in physics. Importantly, the differential operators work with flexible boundary conditions.

The PDEs to solve are represented as a separate class inheriting from *PDEBase*. One example defined in this package is the diffusion equation implemented as *DiffusionPDE*, but more specific situations need to be implemented by the user. Most notably, PDEs can be specified by their expression using the convenient *PDE* class.

The PDEs are solved using solver classes, where a simple explicit solver is implemented by *ExplicitSolver*, but more advanced implementations can be done. To obtain more details during the simulation, trackers can be attached to the solver instance, which analyze intermediate states periodically. Typical trackers include *ProgressTracker* (display simulation progress), *PlotTracker* (display images of the simulation), and *SteadyStateTracker* (aborting simulation when a stationary state is reached). Others can be found in the *trackers* module. Moreover, we provide *MemoryStorage* and *FileStorage*, which can be used as trackers to store the intermediate state to memory and to a file, respectively.

#### CHAPTER

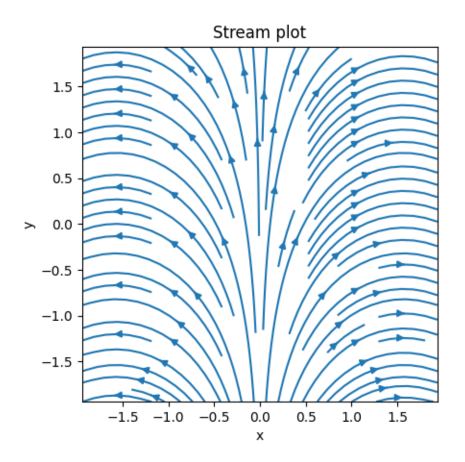
TWO

#### **EXAMPLES**

These are example scripts using the *py-pde* package, which illustrates some of the most important features of the package.

# 2.1 Plotting a vector field

This example shows how to initialize and visualize the vector field  $u = (\sin(x), \cos(x))$ .



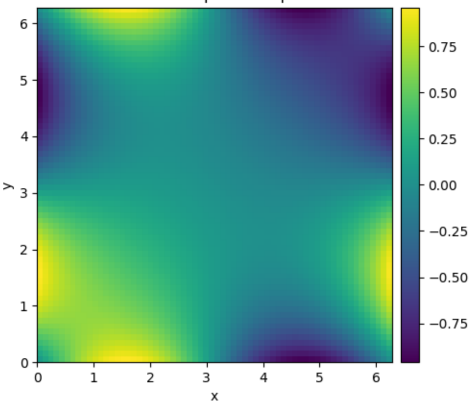


```
field = VectorField.from_expression(grid, ["sin(x)", "cos(x)"])
field.plot(method="streamplot", title="Stream plot")
```

Total running time of the script: (0 minutes 0.521 seconds)

# 2.2 Solving Laplace's equation in 2d

This example shows how to solve a 2d Laplace equation with spatially varying boundary conditions.



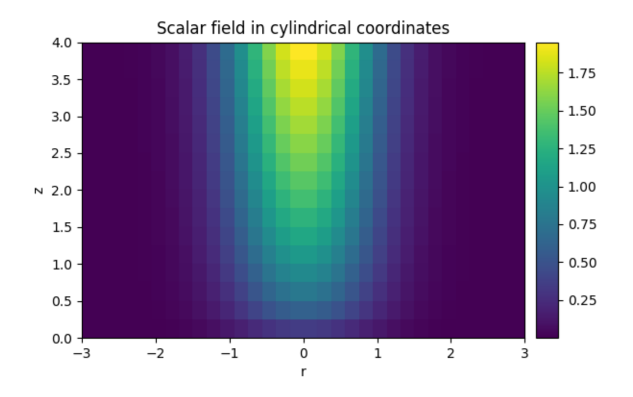
#### Solution to Laplace's equation

import numpy as np
from pde import CartesianGrid, solve\_laplace\_equation
grid = CartesianGrid([[0, 2 \* np.pi]] \* 2, 64)
bcs = [{"value": "sin(y)"}, {"value": "sin(x)"}]
res = solve\_laplace\_equation(grid, bcs)
res.plot()

Total running time of the script: (0 minutes 0.647 seconds)

#### 2.3 Plotting a scalar field in cylindrical coordinates

This example shows how to initialize and visualize the scalar field  $u = \sqrt{z} \exp(-r^2)$  in cylindrical coordinates.



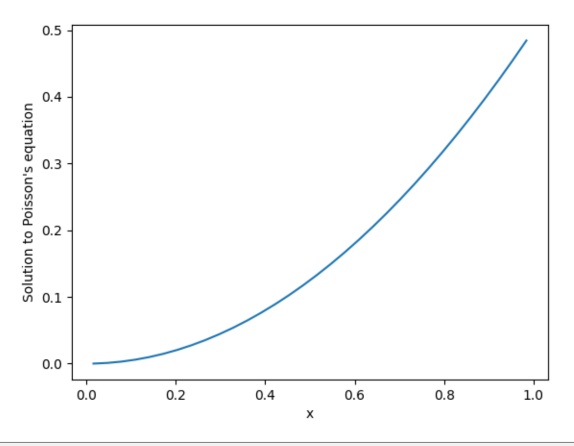
from pde import CylindricalSymGrid, ScalarField
grid = CylindricalSymGrid(radius=3, bounds\_z=[0, 4], shape=16)
field = ScalarField.from\_expression(grid, "sqrt(z) \* exp(-r\*\*2)")

field.plot(title="Scalar field in cylindrical coordinates")

#### Total running time of the script: (0 minutes 0.384 seconds)

# 2.4 Solving Poisson's equation in 1d

This example shows how to solve a 1d Poisson equation with boundary conditions.

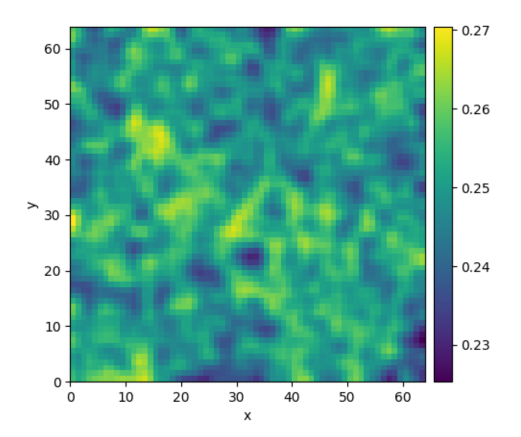


from pde import CartesianGrid, ScalarField, solve\_poisson\_equation
grid = CartesianGrid([[0, 1]], 32, periodic=False)
field = ScalarField(grid, 1)
result = solve\_poisson\_equation(field, bc=[{"value": 0}, {"derivative": 1}])
result.plot()

Total running time of the script: (0 minutes 0.101 seconds)

# 2.5 Simple diffusion equation

This example solves a simple diffusion equation in two dimensions.



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0%	0.02/10.0 [00:09<1:17:40, 467.01s/it]
7%	0.708/10.0 [00:09<02:02, 13.21s/it]
6781	6.697/10.0 [00:09<00:04, 1.41s/it]
6781	6.697/10.0 [00:09<00:04, 1.42s/it]
100%	10.0/10.0 [00:09<00:00, 1.05it/s]
100%	10.0/10.0 [00:09<00:00, 1.05it/s]

```
from pde import DiffusionPDE, ScalarField, UnitGrid
grid = UnitGrid([64, 64]) # generate grid
state = ScalarField.random_uniform(grid, 0.2, 0.3) # generate initial condition
```

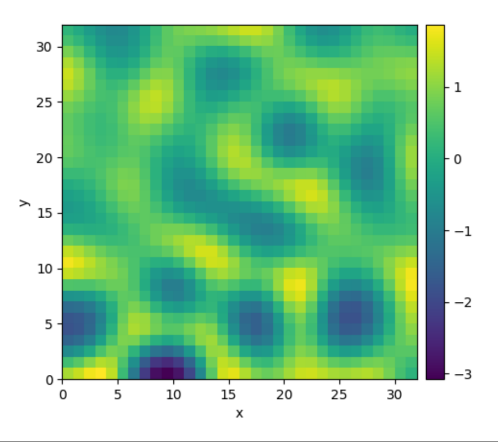
```
eq = DiffusionPDE(diffusivity=0.1) # define the pde
result = eq.solve(state, t_range=10)
result.plot()
```

Total running time of the script: (0 minutes 9.677 seconds)

# 2.6 Kuramoto-Sivashinsky - Using PDE class

This example implements a scalar PDE using the *PDE*. We here consider the Kuramoto–Sivashinsky equation, which for instance describes the dynamics of flame fronts:

$$\partial_t u = -\frac{1}{2} |\nabla u|^2 - \nabla^2 u - \nabla^4 u$$



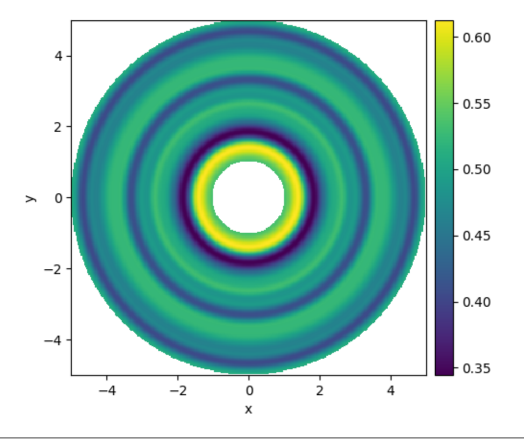
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181	0.12/10.0 [00:24<33:14, 201.85s/it]
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Total running time of the script: (0 minutes 24.413 seconds)

### 2.7 Spherically symmetric PDE

This example illustrates how to solve a PDE in a spherically symmetric geometry.

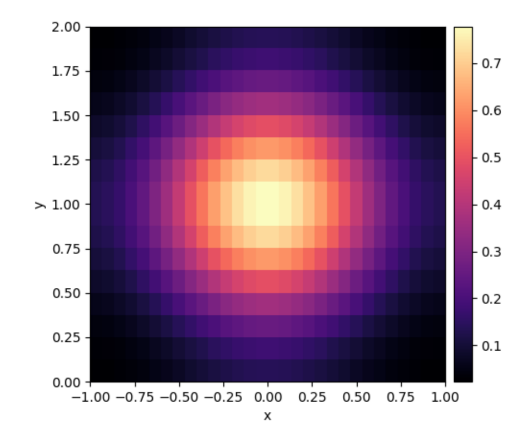


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43%	0.043/0.1 [00:02<00:02, 47.74s/it]
43%	0.043/0.1 [00:02<00:02, 47.76s/it]
100%	0.1/0.1 [00:02<00:00, 20.54s/it]
100%	0.1/0.1 [00:02<00:00, 20.54s/it]

```
from pde import DiffusionPDE, ScalarField, SphericalSymGrid
grid = SphericalSymGrid(radius=[1, 5], shape=128) # generate grid
state = ScalarField.random_uniform(grid) # generate initial condition
eq = DiffusionPDE(0.1) # define the PDE
result = eq.solve(state, t_range=0.1, dt=0.001)
result.plot(kind="image")
```

Total running time of the script: (0 minutes 2.226 seconds)

# 2.8 Diffusion on a Cartesian grid



This example shows how to solve the diffusion equation on a Cartesian grid.

	0%	0/1.0 [00:00 , ?it/s]</th
	Initializing:	0%    0/1.0 [00:00 , ?it/s]</th
	0%	0/1.0 [00:03 , ?it/s]</td
	18	0.01/1.0 [00:04<06:37, 401.55s/it]
	3%	0.03/1.0 [00:04<02:09, 133.86s/it]
	3%	0.03/1.0 [00:04<02:09, 133.91s/it]
	100%	1.0/1.0 [00:04<00:00, 4.02s/it]
	100%	1.0/1.0 [00:04<00:00, 4.02s/it]
- 1		

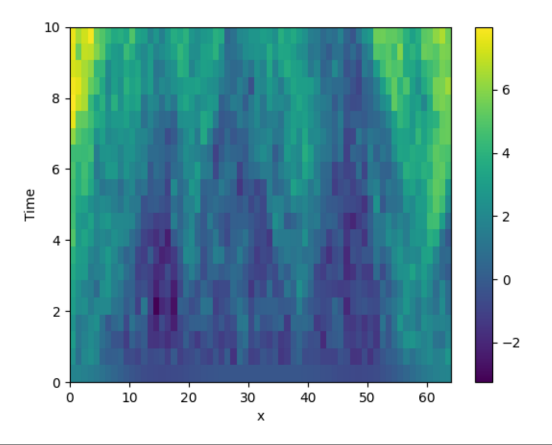
```
from pde import CartesianGrid, DiffusionPDE, ScalarField
grid = CartesianGrid([[-1, 1], [0, 2]], [30, 16]) # generate grid
state = ScalarField(grid) # generate initial condition
state.insert([0, 1], 1)
```

```
eq = DiffusionPDE(0.1) # define the pde
result = eq.solve(state, t_range=1, dt=0.01)
result.plot(cmap="magma")
```

Total running time of the script: (0 minutes 4.169 seconds)

### 2.9 Stochastic simulation

This example illustrates how a stochastic simulation can be done.

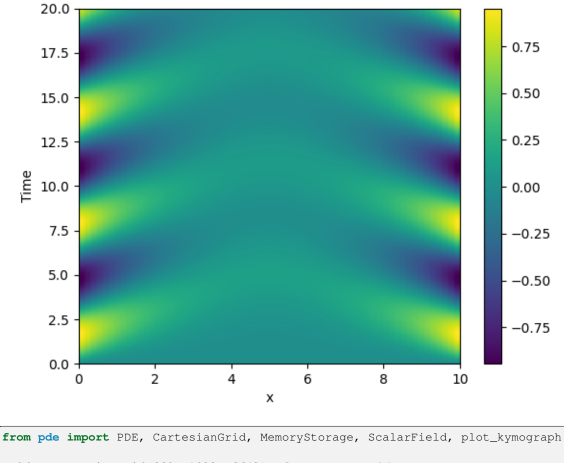


```
from pde import KPZInterfacePDE, MemoryStorage, ScalarField, UnitGrid, plot_kymograph
grid = UnitGrid([64]) # generate grid
state = ScalarField.random_harmonic(grid) # generate initial condition
eq = KPZInterfacePDE(noise=1) # define the SDE
storage = MemoryStorage()
eq.solve(state, t_range=10, dt=0.01, tracker=storage.tracker(0.5))
plot_kymograph(storage)
```

Total running time of the script: (0 minutes 4.843 seconds)

# 2.10 Time-dependent boundary conditions

This example solves a simple diffusion equation in one dimensions with time-dependent boundary conditions.

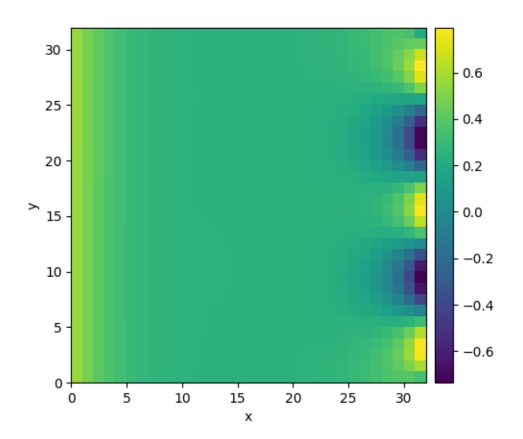


grid = CartesianGrid([[0, 10]], [64]) # generate grid state = ScalarField(grid) # generate initial condition eq = PDE({"c": "laplace(c)"}, bc={"value\_expression": "sin(t)"}) storage = MemoryStorage() eq.solve(state, t\_range=20, dt=1e-4, tracker=storage.tracker(0.1)) # plot the trajectory as a space-time plot plot\_kymograph(storage)

Total running time of the script: (0 minutes 8.670 seconds)

# 2.11 Setting boundary conditions

This example shows how different boundary conditions can be specified.



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0%	0/10.0 [00:04 , ?it/s]</td
0%	0.005/10.0 [00:04<2:28:59, 894.36s/it]
0%	0.03/10.0 [00:04<24:46, 149.07s/it]
14%	1.355/10.0 [00:04<00:28, 3.30s/it]
1481	1.355/10.0 [00:04<00:28, 3.32s/it]
100%	10.0/10.0 [00:04<00:00, 2.22it/s]
100%	10.0/10.0 [00:04<00:00, 2.22it/s]

```
from pde import DiffusionPDE, ScalarField, UnitGrid
grid = UnitGrid([32, 32], periodic=[False, True]) # generate grid
state = ScalarField.random_uniform(grid, 0.2, 0.3) # generate initial condition
```

```
# set boundary conditions `bc` for all axes
bc_x_left = {"derivative": 0.1}
bc_x_right = {"value": "sin(y / 2)"}
bc_x = [bc_x_left, bc_x_right]
bc_y = "periodic"
eq = DiffusionPDE(bc=[bc_x, bc_y])
result = eq.solve(state, t_range=10, dt=0.005)
result.plot()
```

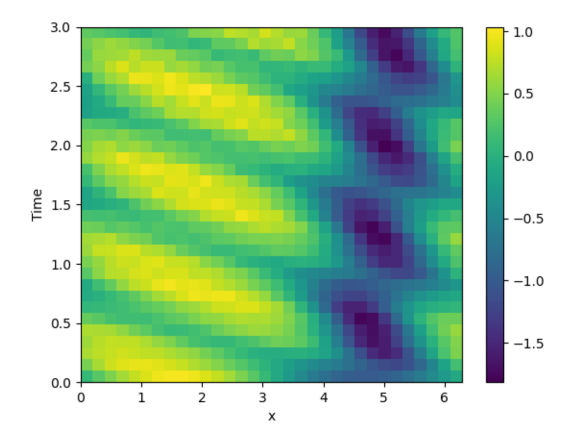
Total running time of the script: (0 minutes 4.646 seconds)

#### 2.12 1D problem - Using PDE class

This example implements a PDE that is only defined in one dimension. Here, we chose the Korteweg-de Vries equation, given by

$$\partial_t \phi = 6\phi \partial_x \phi - \partial_x^3 \phi$$

which we implement using the PDE.



```
from math import pi
from pde import PDE, CartesianGrid, MemoryStorage, ScalarField, plot_kymograph
# initialize the equation and the space
eq = PDE({"\u03c6"", "6 * \u03c6 * \u03c6 d_dx(\u03c6) - laplace(d_dx(\u03c6))"})
grid = CartesianGrid([[0, 2 * pi]], [32], periodic=True)
state = ScalarField.from_expression(grid, "sin(x)")
# solve the equation and store the trajectory
storage = MemoryStorage()
eq.solve(state, t_range=3, solver="scipy", tracker=storage.tracker(0.1))
# plot the trajectory as a space-time plot
plot_kymograph(storage)
```

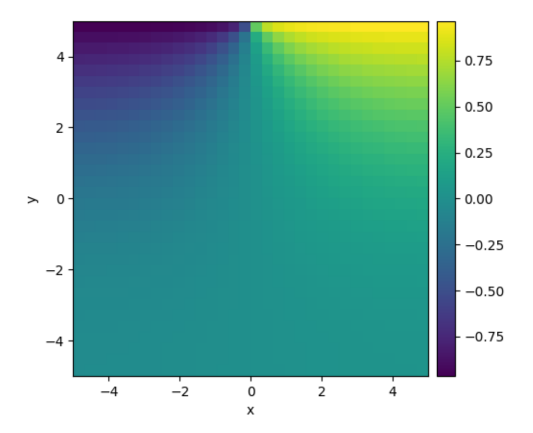
Total running time of the script: (0 minutes 8.274 seconds)

### 2.13 Heterogeneous boundary conditions

This example implements a spatially coupled SIR model with the following dynamics for the density of susceptible, infected, and recovered individuals:

$$\partial_t s = D\nabla^2 s - \beta i s$$
$$\partial_t i = D\nabla^2 i + \beta i s - \gamma i$$
$$\partial_t r = D\nabla^2 r + \gamma i$$

Here, D is the diffusivity,  $\beta$  the infection rate, and  $\gamma$  the recovery rate.



0%	0/10.0 [00:00 , ?it/s]</th
Initializing:	0%    0/10.0 [00:00 , ?it/s]</th
081	0/10.0 [00:00 , ?it/s]</td
3%	0.27/10.0 [00:00<00:07, 1.38it/s]
981	0.87/10.0 [00:00<00:02, 3.67it/s]
38%	3.83/10.0 [00:00<00:00, 8.82it/s]
38%	3.83/10.0 [00:00<00:01, 4.56it/s]
100%	10.0/10.0 [00:00<00:00, 11.91it/s]
100%	10.0/10.0 [00:00<00:00, 11.91it/s]

```
import numpy as np
from pde import CartesianGrid, DiffusionPDE, ScalarField
# define grid and an initial state
grid = CartesianGrid([[-5, 5], [-5, 5]], 32)
field = ScalarField(grid)
# define the boundary conditions, which here are calculated from a function
(continues on next page)
```

```
def bc_value(adjacent_value, dx, x, y, t):
    """return boundary value"""
    return np.sign(x)

bc_x = "derivative"
bc_y = ["derivative", {"value_expression": bc_value}]

# define and solve a simple diffusion equation
eq = DiffusionPDE(bc=[bc_x, bc_y])
res = eq.solve(field, t_range=10, dt=0.01, backend="numpy")
res.plot()
```

Total running time of the script: (0 minutes 0.983 seconds)

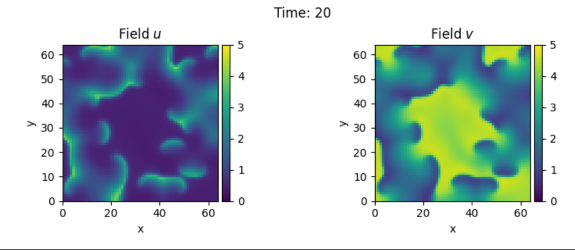
#### 2.14 Brusselator - Using the PDE class

This example uses the PDE class to implement the Brusselator with spatial coupling,

$$\partial_t u = D_0 \nabla^2 u + a - (1+b)u + vu^2$$
$$\partial_t v = D_1 \nabla^2 v + bu - vu^2$$

Here,  $D_0$  and  $D_1$  are the respective diffusivity and the parameters a and b are related to reaction rates.

Note that the same result can also be achieved with a *full implementation of a custom class*, which allows for more flexibility at the cost of code complexity.



#### from pde import PDE, FieldCollection, PlotTracker, ScalarField, UnitGrid

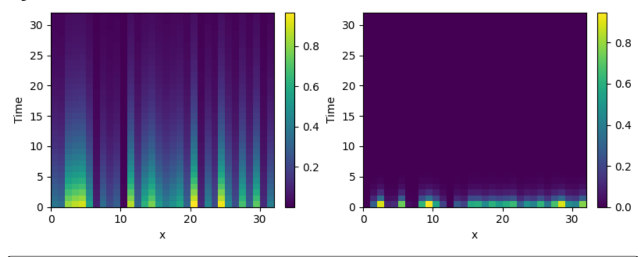
```
# define the PDE
a, b = 1, 3
d0, d1 = 1, 0.1
eq = PDE(
        {
            "u": f"{d0} * laplace(u) + {a} - ({b} + 1) * u + u**2 * v",
            "v": f"{d1} * laplace(v) + {b} * u - u**2 * v",
        }
)
```

```
# initialize state
grid = UnitGrid([64, 64])
u = ScalarField(grid, a, label="Field $u$")
v = b / a + 0.1 * ScalarField.random_normal(grid, label="Field $v$")
state = FieldCollection([u, v])
# simulate the pde
tracker = PlotTracker(interrupts=1, plot_args={"vmin": 0, "vmax": 5})
sol = eq.solve(state, t_range=20, dt=1e-3, tracker=tracker)
```

Total running time of the script: (0 minutes 27.736 seconds)

#### 2.15 Writing and reading trajectory data

This example illustrates how to store intermediate data to a file for later post-processing. The storage frequency is an argument to the tracker.



```
import pde
```

```
# define grid, state and pde
grid = pde.UnitGrid([32])
state = pde.FieldCollection(
    [pde.ScalarField.random_uniform(grid), pde.VectorField.random_uniform(grid)]
)
eq = pde.PDE({"s": "-0.1 * s", "v": "-v"})
# get a temporary file to write data to
path = NamedTemporaryFile(suffix=".hdf5")
# run a simulation and write the results
writer = pde.FileStorage(path.name, write_mode="truncate")
eq.solve(state, t_range=32, dt=0.01, tracker=writer.tracker(1))
# read the simulation back in again
```

(continues on next page)

from tempfile import NamedTemporaryFile

```
reader = pde.FileStorage(path.name, write_mode="read_only")
pde.plot_kymographs(reader)
```

Total running time of the script: (0 minutes 5.118 seconds)

### 2.16 Diffusion equation with spatial dependence

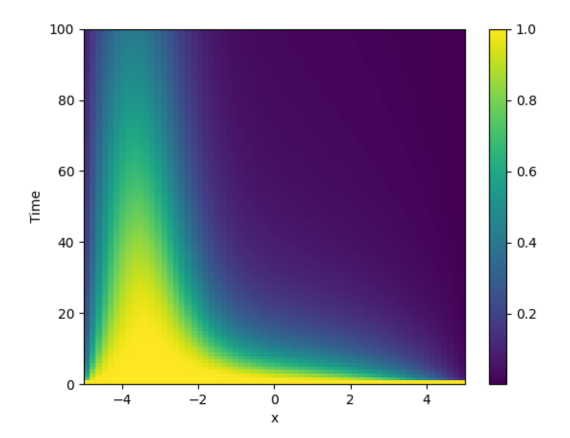
This example solve the Diffusion equation with a heterogeneous diffusivity:

$$\partial_t c = \nabla \big( D(\boldsymbol{r}) \nabla c \big)$$

using the *PDE* class. In particular, we consider D(x) = 1.01 + tanh(x), which gives a low diffusivity on the left side of the domain.

Note that the naive implementation,  $PDE({"c": "divergence((1.01 + tanh(x)) * gradi$  $ent(c))"})$ , has numerical instabilities. This is because two finite difference approximations are nested. To arrive at a more stable numerical scheme, it is advisable to expand the divergence,

$$\partial_t c = D\nabla^2 c + \nabla D.\nabla c$$

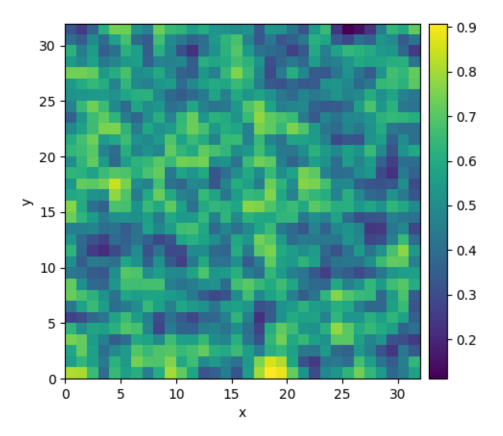


```
from pde import PDE, CartesianGrid, MemoryStorage, ScalarField, plot_kymograph
# Expanded definition of the PDE
diffusivity = "1.01 + tanh(x)"
term_1 = f"({diffusivity}) * laplace(c)"
term_2 = f"dot(gradient({diffusivity}), gradient(c))"
eq = PDE({"c": f"{term_1} + {term_2}"}, bc={"value": 0})
grid = CartesianGrid([[-5, 5]], 64) # generate grid
field = ScalarField(grid, 1) # generate initial condition
storage = MemoryStorage() # store intermediate information of the simulation
res = eq.solve(field, 100, dt=1e-3, tracker=storage.tracker(1)) # solve the PDE
plot_kymograph(storage) # visualize the result in a space-time plot
```

Total running time of the script: (0 minutes 12.294 seconds)

### 2.17 Using simulation trackers

This example illustrates how trackers can be used to analyze simulations.



Time: 3

0%	0/3.0 [00:00 , ?it/s]</th
Initializing:	0%    0/3.0 [00:00 , ?it/s]</th
0%	0/3.0 [00:04 , ?it/s]</th
3%	0.1/3.0 [00:04<02:12, 45.70s/it]
7%	0.2/3.0 [00:04<01:03, 22.85s/it]
238	0.7/3.0 [00:04<00:15, 6.53s/it]
23%	0.7/3.0 [00:04<00:15, 6.79s/it]
100%	3.0/3.0 [00:04<00:00, 1.58s/it]
100%	3.0/3.0 [00:04<00:00, 1.58s/it]
532.9734026319	691
532.9734026319	691
532.9734026319	691
532.9734026319	691

```
import pde
grid = pde.UnitGrid([32, 32]) # generate grid
state = pde.ScalarField.random_uniform(grid) # generate initial condition
storage = pde.MemoryStorage()
trackers = [
    "progress", # show progress bar during simulation
    "steady_state", # abort when steady state is reached
   storage.tracker(interrupts=1), # store data every simulation time unit
   pde.PlotTracker(show=True), # show images during simulation
   # print some output every 5 real seconds:
   pde.PrintTracker(interrupts=pde.RealtimeInterrupts(duration=5)),
]
eq = pde.DiffusionPDE(0.1) # define the PDE
eq.solve(state, 3, dt=0.1, tracker=trackers)
for field in storage:
   print(field.integral)
```

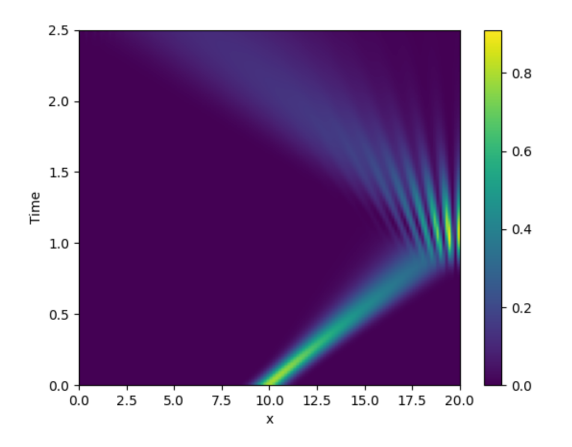
Total running time of the script: (0 minutes 4.812 seconds)

### 2.18 Schrödinger's Equation

This example implements a complex PDE using the *PDE*. We here chose the Schrödinger equation without a spatial potential in non-dimensional form:

$$i\partial_t\psi = -\nabla^2\psi$$

Note that the example imposes Neumann conditions at the wall, so the wave packet is expected to reflect off the wall.



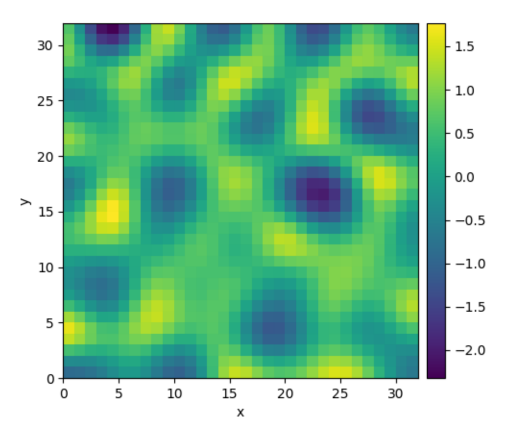
```
from math import sqrt
from pde import PDE, CartesianGrid, MemoryStorage, ScalarField, plot_kymograph
grid = CartesianGrid([[0, 20]], 128, periodic=False)  # generate grid
# create a (normalized) wave packet with a certain form as an initial condition
initial_state = ScalarField.from_expression(grid, "exp(I * 5 * x) * exp(-(x - 10)**2)
...,")
initial_state /= sqrt(initial_state.to_scalar("norm_squared").integral.real)
eq = PDE({"\u03c4": f"I * laplace(\u03c4)"})  # define the pde
# solve the pde and store intermediate data
storage = MemoryStorage()
eq.solve(initial_state, t_range=2.5, dt=1e-5, tracker=[storage.tracker(0.02)])
# visualize the results as a space-time plot
plot_kymograph(storage, scalar="norm_squared")
```

Total running time of the script: (0 minutes 6.702 seconds)

# 2.19 Kuramoto-Sivashinsky - Using custom class

This example implements a scalar PDE using a custom class. We here consider the Kuramoto–Sivashinsky equation, which for instance describes the dynamics of flame fronts:

$$\partial_t u = -\frac{1}{2} |\nabla u|^2 - \nabla^2 u - \nabla^4 u$$



0%	0/10.0 [00:00 , ?it/s]</th <th></th>	
Initializing:	0%    0/10.0 [00:00 , ?it/s]</th <th></th>	
0%	0/10.0 [00:00 , ?it/s]</th <th></th>	
3%	0.27/10.0 [00:00<00:12, 1.26s/it]	
7%	0.72/10.0 [00:00<00:04, 1.87it/s]	
28%	2.83/10.0 [00:00<00:01, 4.70it/s]	
7481	7.36/10.0 [00:01<00:00, 6.86it/s]	
7481	7.36/10.0 [00:01<00:00, 5.47it/s]	
100%	10.0/10.0 [00:01<00:00, 7.43it/s]	
100%	10.0/10.0 [00:01<00:00, 7.43it/s]	
(		

```
from pde import PDEBase, ScalarField, UnitGrid

class KuramotoSivashinskyPDE(PDEBase):
    """Implementation of the normalized Kuramoto-Sivashinsky equation"""
    def evolution_rate(self, state, t=0):
        """implement the python version of the evolution equation"""
        state_lap = state.laplace(bc="auto_periodic_neumann")
        state_grad = state.gradient(bc="auto_periodic_neumann")
        state_grad = state.gradient(bc="auto_periodic_neumann")
        return -state_grad.to_scalar("squared_sum") / 2 - state_lap - state_lap2

grid = UnitGrid([32, 32])  # generate grid
state = ScalarField.random_uniform(grid)  # generate initial condition

eq = KuramotoSivashinskyPDE()  # define the pde
result = eq.solve(state, t_range=10, dt=0.01)
result.plot()
```

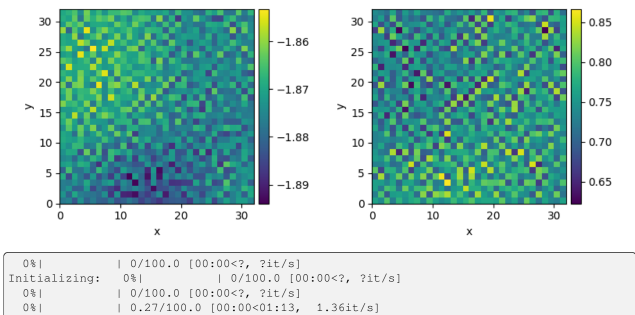
Total running time of the script: (0 minutes 1.494 seconds)

#### 2.20 Custom Class for coupled PDEs

This example shows how to solve a set of coupled PDEs, the spatially coupled FitzHugh–Nagumo model, which is a simple model for the excitable dynamics of coupled Neurons:

$$\partial_t u = \nabla^2 u + u(u - \alpha)(1 - u) + w$$
$$\partial_t w = \epsilon u$$

Here,  $\alpha$  denotes the external stimulus and  $\epsilon$  defines the recovery time scale. We implement this as a custom PDE class below.



18	0.86/100.0 [00:00<00:27, 3.56it/s]
481	3.69/100.0 [00:00<00:11, 8.20it/s]
10%	9.9/100.0 [00:00<00:08, 10.72it/s]
1981	18.92/100.0 [00:01<00:07, 10.55it/s]
298	28.58/100.0 [00:02<00:06, 10.96it/s]
3981	39.3/100.0 [00:03<00:05, 11.45it/s]
518	51.11/100.0 [00:04<00:04, 11.20it/s]
62%	62.2/100.0 [00:05<00:03, 11.50it/s]
7481	74.28/100.0 [00:06<00:02, 11.48it/s]
86%	85.99/100.0 [00:07<00:01, 11.49it/s]
98%	97.63/100.0 [00:08<00:00, 11.64it/s]
98%	97.63/100.0 [00:08<00:00, 11.33it/s]
100%	100.0/100.0 [00:08<00:00, 11.60it/s]
100%	100.0/100.0 [00:08<00:00, 11.60it/s]

```
from pde import FieldCollection, PDEBase, UnitGrid
class FitzhughNagumoPDE(PDEBase):
    """FitzHugh-Nagumo model with diffusive coupling"""
   def __init__(self, stimulus=0.5, τ=10, a=0, b=0, bc="auto_periodic_neumann"):
       super().__init__()
        self.bc = bc
        self.stimulus = stimulus
        self.\tau = \tau
        self.a = a
        self.b = b
    def evolution_rate(self, state, t=0):
        v, w = state # membrane potential and recovery variable
        v_t = v.laplace(bc=self.bc) + v - v**3 / 3 - w + self.stimulus
        w_t = (v + self.a - self.b * w) / self.\tau
        return FieldCollection([v_t, w_t])
grid = UnitGrid([32, 32])
state = FieldCollection.scalar_random_uniform(2, grid)
eq = FitzhughNagumoPDE()
result = eq.solve(state, t_range=100, dt=0.01)
result.plot()
```

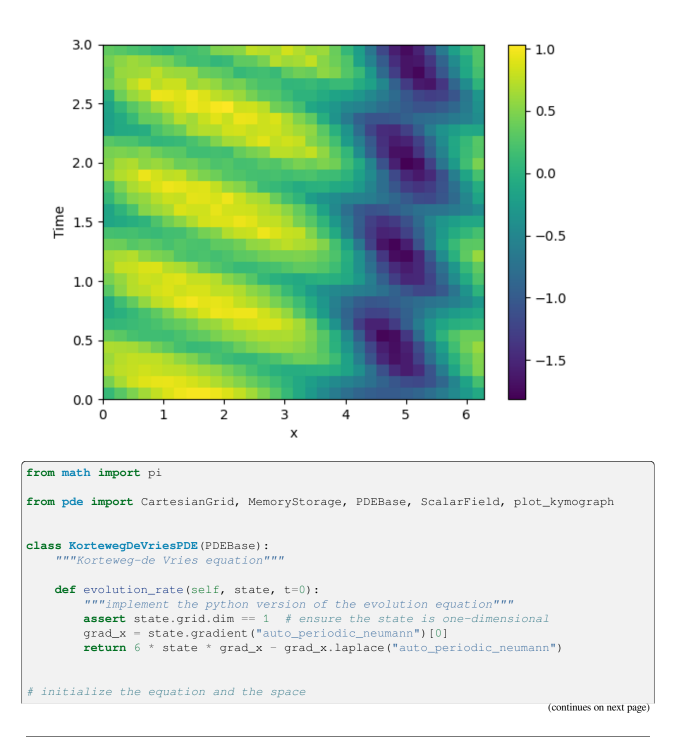
Total running time of the script: (0 minutes 8.950 seconds)

# 2.21 1D problem - Using custom class

This example implements a PDE that is only defined in one dimension. Here, we chose the Korteweg-de Vries equation, given by

$$\partial_t \phi = 6\phi \partial_x \phi - \partial_x^3 \phi$$

which we implement using a custom PDE class below.

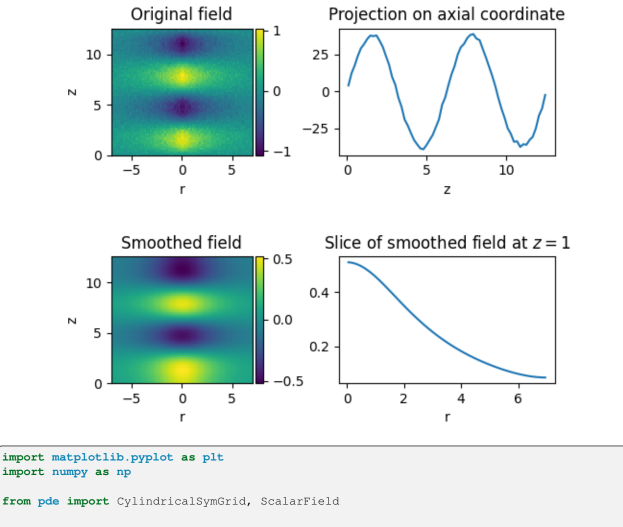


```
grid = CartesianGrid([[0, 2 * pi]], [32], periodic=True)
state = ScalarField.from_expression(grid, "sin(x)")
# solve the equation and store the trajectory
storage = MemoryStorage()
eq = KortewegDeVriesPDE()
eq.solve(state, t_range=3, solver="scipy", tracker=storage.tracker(0.1))
# plot the trajectory as a space-time plot
plot_kymograph(storage)
```

Total running time of the script: (0 minutes 2.528 seconds)

# 2.22 Visualizing a scalar field

This example displays methods for visualizing scalar fields.



(continues on next page)

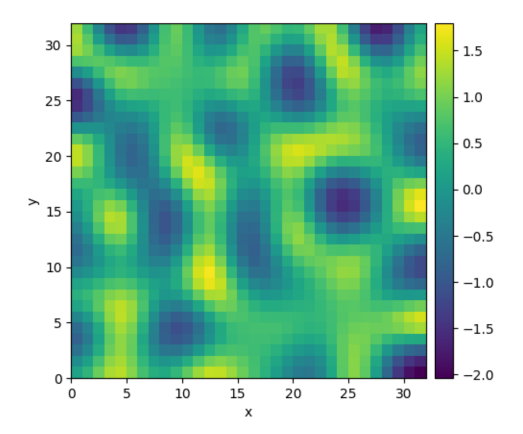
```
# create a scalar field with some noise
grid = CylindricalSymGrid(7, [0, 4 * np.pi], 64)
data = ScalarField.from_expression(grid, "sin(z) * exp(-r / 3)")
data += 0.05 * ScalarField.random_normal(grid)
# manipulate the field
smoothed = data.smooth() # Gaussian smoothing to get rid of the noise
projected = data.project("r") # integrate along the radial direction
sliced = smoothed.slice({"z": 1}) # slice the smoothed data
# create four plots of the field and the modifications
fig, axes = plt.subplots(nrows=2, ncols=2)
data.plot(ax=axes[0, 0], title="Original field")
smoothed.plot(ax=axes[1, 0], title="Smoothed field")
projected.plot(ax=axes[0, 1], title="Projection on axial coordinate")
sliced.plot(ax=axes[1, 1], title="Slice of smoothed field at $z=1$")
plt.subplots_adjust(hspace=0.8)
plt.show()
```

Total running time of the script: (0 minutes 0.316 seconds)

# 2.23 Kuramoto-Sivashinsky - Compiled methods

This example implements a scalar PDE using a custom class with a numba-compiled method for accelerated calculations. We here consider the Kuramoto–Sivashinsky equation, which for instance describes the dynamics of flame fronts:

$$\partial_t u = -\frac{1}{2} |\nabla u|^2 - \nabla^2 u - \nabla^4 u$$



0%	0/10.0 [00:00 , ?it/s]</th
Initializing:	0%    0/10.0 [00:00 , ?it/s]</td
0%	0/10.0 [00:09 , ?it/s]</td
0%	0.01/10.0 [00:11<3:06:09, 1118.08s/it]
0%	0.02/10.0 [00:11<1:32:59, 559.06s/it]
281	0.18/10.0 [00:11<10:10, 62.12s/it]
80%	8.01/10.0 [00:11<00:02, 1.40s/it]
80%	8.01/10.0 [00:11<00:02, 1.40s/it]
100%	10.0/10.0 [00:11<00:00, 1.12s/it]
100%	10.0/10.0 [00:11<00:00, 1.12s/it]

```
import numba as nb
from pde import PDEBase, ScalarField, UnitGrid
class KuramotoSivashinskyPDE (PDEBase):
    """Implementation of the normalized Kuramoto-Sivashinsky equation"""
    def __init__(self, bc="auto_periodic_neumann"):
```

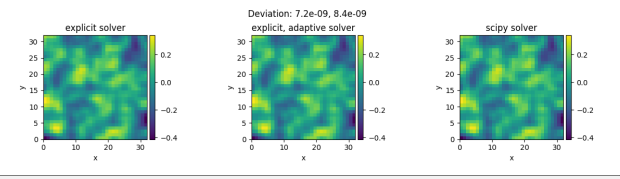
(continues on next page)

```
super().__init__()
        self.bc = bc
    def evolution_rate(self, state, t=0):
        """implement the python version of the evolution equation"""
        state_lap = state.laplace(bc=self.bc)
        state_lap2 = state_lap.laplace(bc=self.bc)
        state_grad_sq = state.gradient_squared(bc=self.bc)
        return -state_grad_sq / 2 - state_lap - state_lap2
    def _make_pde_rhs_numba(self, state):
        """nunmba-compiled implementation of the PDE"""
        gradient_squared = state.grid.make_operator("gradient_squared", bc=self.bc)
        laplace = state.grid.make_operator("laplace", bc=self.bc)
        @nb.njit
        def pde_rhs(data, t):
            return -0.5 * gradient_squared(data) - laplace(data + laplace(data))
        return pde_rhs
grid = UnitGrid([32, 32]) # generate grid
state = ScalarField.random_uniform(grid) # generate initial condition
eq = KuramotoSivashinskyPDE() # define the pde
result = eq.solve(state, t_range=10, dt=0.01)
result.plot()
```

Total running time of the script: (0 minutes 11.355 seconds)

# 2.24 Solver comparison

This example shows how to set up solvers explicitly and how to extract diagnostic information.



```
Diagnostic information from first run:
{'controller': {'t_start': 0, 't_end': 1.0, 'profiler': {'solver': 0.

$\obsigmath{totop} 05391846700001679, 'tracker': 4.970599997022873e-05, 'compilation': 4.

$\obsigmath{totop} 4027449630000035}, 'jit_count': {'make_stepper': 11, 'simulation': 0}, 'solver_start

$\obsigmath{totop} : '2024-04-04 06:59:14.746044', 'successful': True, 'stop_reason': 'Reached final_

$\operatormath{totop} : 'solver_duration': '0:00:00.053963', 't_final': 1.0, 'process_count': 1},

$\operatormath{totop} = package_version': 'unknown', 'solver': {'class': 'ExplicitSolver', 'pde_class':

$\operatormath{totop} = package_version': 'unknown', 'steps': 1000, 'state_modifications': 0.0, 'stochastic

(continues on next page)
```

```
Diagnostic information from second run:
{'controller': {'t_start': 0, 't_end': 1.0, 'profiler': {'solver': 0.
→19512004099999558, 'tracker': 4.7649999999066495e-05, 'compilation': 1.
→1542392590000077}, 'jit_count': {'make_stepper': 2, 'simulation': 0}, 'solver_start
→': '2024-04-04 06:59:15.955103', 'successful': True, 'stop_reason': 'Reached final_

white output and the set of the set o
→ 'DiffusionPDE', 'dt': 0.001, 'dt_adaptive': True, 'steps': 12, 'stochastic': False,
→16639903703254955, 'mean': 0.0833333333333334, 'std': 0.05138013183612336, 'count
Diagnostic information from third run:
{'controller': {'t_start': 0, 't_end': 1.0, 'profiler': {'solver': 0.
→0034982200000115427, 'tracker': 4.478599998947175e-05, 'compilation': 0.
→6280662980000216}, 'jit_count': {'make_stepper': 1, 'simulation': 0}, 'solver_start
→time', 'solver_duration': '0:00:00.003541', 't_final': 1.0, 'process_count': 1},
-- 'package_version': 'unknown', 'solver': {'class': 'ScipySolver', 'pde_class':
→ 'DiffusionPDE', 'dt': None, 'steps': 50, 'stochastic': False, 'backend': 'numba'}}
```

```
import pde
```

```
# initialize the grid, an initial condition, and the PDE
grid = pde.UnitGrid([32, 32])
field = pde.ScalarField.random_uniform(grid, -1, 1)
eq = pde.DiffusionPDE()
# try the explicit solver
solver1 = pde.ExplicitSolver(eq)
controller1 = pde.Controller(solver1, t_range=1, tracker=None)
sol1 = controller1.run(field, dt=1e-3)
sol1.label = "explicit solver"
print("Diagnostic information from first run:")
print(controller1.diagnostics)
print()
# try an explicit solver with adaptive time steps
solver2 = pde.ExplicitSolver(eq, scheme="runge-kutta", adaptive=True)
controller2 = pde.Controller(solver2, t_range=1, tracker=None)
sol2 = controller2.run(field, dt=1e-3)
sol2.label = "explicit, adaptive solver"
print("Diagnostic information from second run:")
print(controller2.diagnostics)
print()
# try the standard scipy solver
solver3 = pde.ScipySolver(eq)
controller3 = pde.Controller(solver3, t_range=1, tracker=None)
```

(continues on next page)

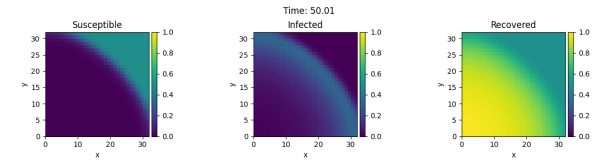
Total running time of the script: (0 minutes 8.355 seconds)

# 2.25 Custom PDE class: SIR model

This example implements a spatially coupled SIR model with the following dynamics for the density of susceptible, infected, and recovered individuals:

$$\begin{split} \partial_t s &= D\nabla^2 s - \beta is \\ \partial_t i &= D\nabla^2 i + \beta is - \gamma i \\ \partial_t r &= D\nabla^2 r + \gamma i \end{split}$$

Here, D is the diffusivity,  $\beta$  the infection rate, and  $\gamma$  the recovery rate.



081	0/50.0 [00:00 , ?it/s]</th
Initializing:	0%    0/50.0 [00:00 , ?it/s]</td
0%	0/50.0 [00:00 , ?it/s]</td
081	0.02/50.0 [00:00<34:18, 41.18s/it]
0%	0.04/50.0 [00:00<17:12, 20.68s/it]
1%	0.47/50.0 [00:00<01:33, 1.89s/it]
481	2.14/50.0 [00:01<00:25, 1.90it/s]
11%	5.59/50.0 [00:01<00:12, 3.48it/s]
21%	10.56/50.0 [00:02<00:09, 4.16it/s]
31%	15.71/50.0 [00:03<00:07, 4.80it/s]
4381	21.73/50.0 [00:04<00:05, 4.98it/s]
55%	27.51/50.0 [00:05<00:04, 5.32it/s]
68%	33.91/50.0 [00:06<00:02, 5.37it/s]
80%	39.9/50.0 [00:07<00:01, 5.57it/s]
93%	46.4/50.0 [00:08<00:00, 5.58it/s]
93%	46.4/50.0 [00:09<00:00, 5.12it/s]
100%	50.0/50.0 [00:09<00:00, 5.52it/s]
100%	50.0/50.0 [00:09<00:00, 5.52it/s]
(	

```
from pde import FieldCollection, PDEBase, PlotTracker, ScalarField, UnitGrid
class SIRPDE(PDEBase):
   """SIR-model with diffusive mobility"""
   def init (
       self, beta=0.3, gamma=0.9, diffusivity=0.1, bc="auto_periodic_neumann"
    ):
       super().__init__()
        self.beta = beta # transmission rate
       self.gamma = gamma # recovery rate
       self.diffusivity = diffusivity # spatial mobility
        self.bc = bc # boundary condition
   def get_state(self, s, i):
        """generate a suitable initial state"""
       norm = (s + i).data.max() # maximal density
        if norm > 1:
           s /= norm
           i /= norm
        s.label = "Susceptible"
        i.label = "Infected"
        # create recovered field
       r = ScalarField(s.grid, data=1 - s - i, label="Recovered")
       return FieldCollection([s, i, r])
   def evolution_rate(self, state, t=0):
       s, i, r = state
        diff = self.diffusivity
       ds_dt = diff * s.laplace(self.bc) - self.beta * i * s
       di_dt = diff * i.laplace(self.bc) + self.beta * i * s - self.gamma * i
        dr_dt = diff * r.laplace(self.bc) + self.gamma * i
        return FieldCollection([ds_dt, di_dt, dr_dt])
eq = SIRPDE(beta=2, gamma=0.1)
# initialize state
grid = UnitGrid([32, 32])
s = ScalarField(grid, 1)
i = ScalarField(grid, 0)
i.data[0, 0] = 1
state = eq.get_state(s, i)
# simulate the pde
tracker = PlotTracker(interrupts=10, plot_args={"vmin": 0, "vmax": 1})
sol = eq.solve(state, t_range=50, dt=1e-2, tracker=["progress", tracker])
```

Total running time of the script: (0 minutes 9.390 seconds)

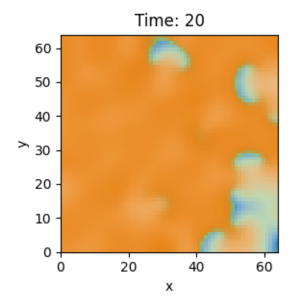
# 2.26 Brusselator - Using custom class

This example implements the Brusselator with spatial coupling,

$$\partial_t u = D_0 \nabla^2 u + a - (1+b)u + vu^2$$
$$\partial_t v = D_1 \nabla^2 v + bu - vu^2$$

Here,  $D_0$  and  $D_1$  are the respective diffusivity and the parameters a and b are related to reaction rates.

Note that the PDE can also be implemented using the *PDE* class; see *the example*. However, that implementation is less flexible and might be more difficult to extend later.



```
import numba as nb
import numpy as np
from pde import FieldCollection, PDEBase, PlotTracker, ScalarField, UnitGrid
class BrusselatorPDE(PDEBase):
    """Brusselator with diffusive mobility"""
   def __init__(self, a=1, b=3, diffusivity=[1, 0.1], bc="auto_periodic_neumann"):
       super().__init__()
        self.a = a
        self.b = b
        self.diffusivity = diffusivity # spatial mobility
        self.bc = bc # boundary condition
   def get_initial_state(self, grid):
        """prepare a useful initial state"""
        u = ScalarField(grid, self.a, label="Field $u$")
        v = self.b / self.a + 0.1 * ScalarField.random_normal(grid, label="Field $v$")
        return FieldCollection([u, v])
    def evolution_rate(self, state, t=0):
        """pure python implementation of the PDE"""
        u, v = state
```

(continues on next page)

```
rhs = state.copy()
        d0, d1 = self.diffusivity
        rhs[0] = d0 * u.laplace(self.bc) + self.a - (self.b + 1) * u + u**2 * v
        rhs[1] = d1 * v.laplace(self.bc) + self.b * u - u**2 * v
        return rhs
    def _make_pde_rhs_numba(self, state):
        """nunmba-compiled implementation of the PDE"""
        d0, d1 = self.diffusivity
        a, b = self.a, self.b
        laplace = state.grid.make_operator("laplace", bc=self.bc)
        @nb.njit
        def pde_rhs(state_data, t):
           u = state_data[0]
            v = state_data[1]
            rate = np.empty_like(state_data)
            rate[0] = d0 * laplace(u) + a - (1 + b) * u + v * u**2
            rate[1] = d1 * laplace(v) + b * u - v * u**2
            return rate
        return pde_rhs
# initialize state
grid = UnitGrid([64, 64])
eq = BrusselatorPDE(diffusivity=[1, 0.1])
state = eq.get_initial_state(grid)
# simulate the pde
tracker = PlotTracker(interrupts=1, plot_args={"kind": "merged", "vmin": 0, "vmax": 5}
\rightarrow)
sol = eq.solve(state, t_range=20, dt=1e-3, tracker=tracker)
```

Total running time of the script: (0 minutes 8.958 seconds)

## CHAPTER

# THREE

# **USER MANUAL**

# 3.1 Mathematical basics

To solve partial differential equations (PDEs), the *py-pde* package provides differential operators to express spatial derivatives. These operators are implemented using the finite difference method to support various boundary conditions. The time evolution of the PDE is then calculated using the method of lines by explicitly discretizing space using the grid classes. This reduces the PDEs to a set of ordinary differential equations, which can be solved using standard methods as described below.

## 3.1.1 Curvilinear coordinates

The package supports multiple curvilinear coordinate systems. They allow to exploit symmetries present in physical systems. Consequently, many grids implemented in *py-pde* inherently assume symmetry of the described fields. However, a drawback of curvilinear coordinates are the fact that the basis vectors now depend on position, which makes tensor fields less intuitive and complicates the expression of differential operators. To avoid confusion, we here specify the used coordinate systems explicitly:

## **Polar coordinates**

Polar coordinates describe points by a radius r and an angle  $\phi$  in a two-dimensional coordinates system. They are defined by the transformation

$$\begin{cases} x = r\cos(\phi) \\ y = r\sin(\phi) \end{cases} \text{ for } r \in [0,\infty] \text{ and } \phi \in [0,2\pi) \end{cases}$$

The associated symmetric grid *PolarSymGrid* assumes that fields only depend on the radial coordinate r. Note that vector and tensor fields can still have components in the polar direction. In particular, vector fields still have two components:  $\vec{v}(r) = v_r(r)\vec{e}_r + v_{\phi}(r)\vec{e}_{\phi}$ .

## **Spherical coordinates**

Spherical coordinates describe points by a radius r, an azimuthal angle  $\theta$ , and a polar angle  $\phi$ . The conversion to ordinary Cartesian coordinates reads

$$\begin{cases} x = r\sin(\theta)\cos(\phi) \\ y = r\sin(\theta)\sin(\phi) & \text{ for } r \in [0,\infty], \ \theta \in [0,\pi], \text{ and } \phi \in [0,2\pi) \\ z = r\cos(\theta) \end{cases}$$

The associated symmetric grid *SphericalSymGrid* assumes that fields only depend on the radial coordinate *r*. Note that vector and tensor fields can still have components in the two angular direction.

**Warning:** Not all results of differential operators on vectorial and tensorial fields can be expressed in terms of fields that only depend on the radial coordinate r. In particular, the gradient of a vector field can only be calculated if the azimuthal component of the vector field vanishes. Similarly, the divergence of a tensor field can only be taken in special situations.

### Cylindrical coordinates

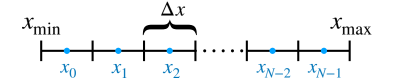
Cylindrical coordinates describe points by a radius r, an axial coordinate z, and a polar angle  $\phi$ . The conversion to ordinary Cartesian coordinates reads

$$\begin{cases} x = r \cos(\phi) \\ y = r \sin(\phi) & \text{ for } r \in [0, \infty], z \in \mathbb{R}, \text{ and } \phi \in [0, 2\pi) \\ z = z \end{cases}$$

The associated symmetric grid CylindricalSymGrid assumes that fields only depend on the coordinates r and z. Vector and tensor fields still specify all components in the three-dimensional space.

**Warning:** The order of components in the vector and tensor fields defined on cylindrical grids is different than in ordinary math. While it is common to use  $(r, \phi, z)$ , we here use the order  $(r, z, \phi)$ . It might thus be best to access components by name instead of index.

## 3.1.2 Spatial discretization



The finite differences scheme used by *py-pde* is currently restricted to orthogonal coordinate systems with uniform discretization. Because of the orthogonality, each axis of the grid can be discretized independently. For simplicity, we only consider uniform grids, where the support points are spaced equidistantly along a given axis, i.e., the discretization  $\Delta x$  is constant. If a given axis covers values in a range  $[x_{\min}, x_{\max}]$ , a discretization with N support points can then be though of as covering the axis with N equal-sized boxes; see inset. Field values are then specified for each box, i.e., the support points lie at the centers of the box:

$$x_i = x_{\min} + \left(i + \frac{1}{2}\right) \Delta x \quad \text{for} \quad i = 0, \dots, N-1$$
$$\Delta x = \frac{x_{\max} - x_{\min}}{N}$$

which is also indicated in the inset.

Differential operators are implemented using the usual second-order central differences. This requires the introducing of virtual support points at  $x_{-1}$  and  $x_N$ , which can be determined from the boundary conditions at  $x = x_{\min}$  and  $x = x_{\max}$ , respectively. The field classes automate this transparently. However, if you need more control over boundary conditions, you can access the full underlying data using the field.\_data\_full, which will have N + 2 entries along an axis

that has N support points. In this case, the first and last entries (data\_full[0] and data\_full[N + 1]) denote the lower and upper virtual point, respectively. The actual field data can be obtained using data\_full[1:-1] or the field.data attribute for convenience. Note that functions evaluating differential operators generally expect the full data as input while they return only valid data.

## 3.1.3 Temporal evolution

Once the fields have been discretized, the PDE reduces to a set of coupled ordinary differential equations (ODEs), which can be solved using standard methods. This reduction is also known as the method of lines. The *py-pde* package implements the simple Euler scheme and a more advanced Runge-Kutta scheme in the *ExplicitSolver* class. For the simple implementations of these explicit methods, the user typically specifies a fixed time step, although adaptive methods, which adjust the time step automatically, are also often used and available in the package. One problem with explicit solvers is that they require small time steps to stably evolve some PDEs; such PDEs are then often called 'stiff'. Stiff PDEs can sometimes be solved more efficiently by using implicit methods. This package provides a simple implementation of the Backward Euler method in the *ImplicitSolver* class. Finally, more advanced methods are available by wrapping the scipy.integrate.solve\_ivp() in the *ScipySolver* class.

# 3.2 Basic usage

We here describe the typical workflow to solve a PDE using *py-pde*. Throughout this section, we assume that the package has been imported using import pde.

# 3.2.1 Defining the geometry

The state of the system is described in a discretized geometry, also known as a *grid*. The package focuses on simple geometries, which work well for the employed finite difference scheme. Grids are defined by instance of various classes that capture the symmetries of the underlying space. In particular, the package offers Cartesian grids of *1* to *3* dimensions via *CartesianGrid*, as well as curvilinear coordinate for spherically symmetric systems in two dimension (*PolarSym-Grid*) and three dimensions (*SphericalSymGrid*), as well as the special class *CylindricalSymGrid* for a cylindrical geometry which is symmetric in the angle.

All grids allow to set the size of the underlying geometry and the number of support points along each axis, which determines the spatial resolution. Moreover, most grids support periodic boundary conditions. For example, a rectangular grid with one periodic boundary condition can be specified as

grid = pde.CartesianGrid([[0, 10], [0, 5]], [20, 10], periodic=[**True, False**])

This grid will have a rectangular shape of 10x5 with square unit cells of side length 0.5. Note that the grid will only be periodic in the *x*-direction.

## 3.2.2 Initializing a field

Fields specifying the values at the discrete points of the grid defined in the previous section. Most PDEs discussed in the package describe a scalar variable, which can be encoded th class *ScalarField*. However, tensors with rank 1 (vectors) and rank 2 are also supported using *VectorField* and *Tensor2Field*, respectively. In any case, a field is initialized using a pre-defined grid, e.g., field = pde.ScalarField(grid). Optional values allow to set the value of the grid, as well as a label that is later used in plotting, e.g., field1 = pde.ScalarField(grid, data=1, label="Ones"). Moreover, fields can be initialized randomly (field2 = pde.ScalarField. random\_normal(grid, mean=0.5)) or from a mathematical expression, which may depend on the coordinates of the grid (field3 = pde.ScalarField.from\_expression(grid, "x \* y")).

All field classes support basic arithmetic operations and can be used much like numpy arrays. Moreover, they have methods for applying differential operators, e.g., the result of applying the Laplacian to a scalar field is returned by calling the method *laplace()*, which returns another instance of *ScalarField*, whereas *gradient()* returns a VectorField. Combining these functions with ordinary arithmetics on fields allows to represent the right hand side of many partial differential equations that appear in physics. Importantly, the differential operators work with flexible boundary conditions.

# 3.2.3 Specifying the PDE

PDEs are also instances of special classes and a number of classical PDEs are already pre-defined in the module pde. pdes. Moreover, the special class PDE allows defining PDEs by simply specifying the expression on their right hand side. To see how this works in practice, let us consider the Kuramoto–Sivashinsky equation,  $\partial_t u = -\nabla^4 u - \nabla^2 u - \frac{1}{2}|\nabla u|^2$ , which describes the time evolution of a scalar field u. A simple implementation of this equation reads

eq = pde.PDE({"u": "-gradient\_squared(u) / 2 - laplace(u + laplace(u))"})

Here, the argument defines the evolution rate for all fields (in this case only u). The expression on the right hand side can contain typical mathematical functions and the operators defined by the package.

# 3.2.4 Running the simulation

To solve the PDE, we first need to generate an initial condition, i.e., the initial values of the fields that are evolved forward in time by the PDE. This field also defined the geometry on which the PDE is solved. In the simplest case, the solution is then obtain by running

result = eq.solve(field, t\_range=10, dt=1e-2)

Here, *t\_range* specifies the duration over which the PDE is considered and *dt* specifies the time step. The *result* field will be defined on the same grid as the initial condition *field*, but instead contain the data value at the final time. Note that all intermediate states are discarded in the simulation above and no information about the dynamical evolution is retained. To study the dynamics, one can either analyze the evolution on the fly or store its state for subsequent analysis. Both these tasks are achieved using *trackers*, which analyze the simulation periodically. For instance, to store the state for some time points in memory, one uses

Note that we also included the special identifier "progress" in the list of trackers, which shows a progress bar during the simulation. Another useful tracker is "plot" which displays the state on the fly.

# 3.2.5 Analyzing the results

Sometimes is suffices to plot the final result, which can be done using result.plot(). The final result can of course also be analyzed quantitatively, e.g., using result.average to obtain its mean value. If the intermediate states have been saved as indicated above, they can be analyzed subsequently:

```
for time, field in storage.items():
    print(f"t={time}, field={field.magnitude}")
```

Moreover, a movie of the simulation can be created using pde.movie(storage, filename=FILE), where *FILE* determines where the movie is written.

# 3.3 Advanced usage

# 3.3.1 Boundary conditions

A crucial aspect of partial differential equations are boundary conditions, which need to be specified at the domain boundaries. For the simple domains contained in *py-pde*, all boundaries are orthogonal to one of the axes in the domain, so boundary conditions need to be applied to both sides of each axis. Here, the lower side of an axis can have a differnt condition than the upper side. For instance, one can enforce the value of a field to be 4 at the lower side and its derivative (in the outward direction) to be 2 on the upper side using the following code:

```
bc_lower = {"value": 4}
bc_upper = {"derivative": 2}
bc = [bc_lower, bc_upper]
grid = pde.UnitGrid([16])
field = pde.ScalarField(grid)
field.laplace(bc)
```

Here, the Laplace operator applied to the field in the last line will respect the boundary conditions. Note that it suffices to give one condition if both sides of the axis require the same condition. For instance, to enforce a value of 3 on both side, one could simply use  $bc = \{ value': 3 \}$ . Vectorial boundary conditions, e.g., to calculate the vector gradient or tensor divergence, can have vectorial values for the boundary condition. Generally, only the normal components at a boundary need to be specified if an operator reduces the rank of a field, e.g., for divergences. Otherwise, e.g., for gradients and Laplacians, the full field needs to be specified at the boundary.

Boundary values that depend on space can be set by specifying a mathematical expression, which may depend on the coordinates of all axes:

```
# two different conditions for lower and upper end of x-axis
bc_x = [{"derivative": 0.1}, {"value": "sin(y / 2)"}]
# the same condition on the lower and upper end of the y-axis
bc_y = {"value": "sqrt(1 + cos(x))"}
grid = UnitGrid([32, 32])
field = pde.ScalarField(grid)
field.laplace(bc=[bc_x, bc_y])
```

**Warning:** To interpret arbitrary expressions, the package uses exec(). It should therefore not be used in a context where malicious input could occur.

Inhomogeneous values can also be specified by directly supplying an array, whose shape needs to be compatible with the boundary, i.e., it needs to have the same shape as the grid but with the dimension of the axis along which the boundary is specified removed.

There exist also special boundary conditions that impose a more complex value of the field  $(bc='value_expression')$  or its derivative  $(bc='derivative_expression')$ . Beyond the spatial coordinates that are already supported for the constant conditions above, the expressions of these boundary conditions can depend on the time variable t. Moreover, these boundary conditions also except python functions (with signature *adjacent\_value, dx, \*coords, t*), thus greatly enlarging the flexibility with which boundary conditions can be expressed. Note that PDEs need to supply the current time *t* when setting the boundary conditions, e.g., when applying the differential operators. The pre-defined PDEs and the general class *PDE* already support time-dependent boundary conditions.

One important aspect about boundary conditions is that they need to respect the periodicity of the underlying grid. For instance, in a 2d grid with one periodic axis, the following boundary condition can be used:

```
grid = pde.UnitGrid([16, 16], periodic=[True, False])
field = pde.ScalarField(grid)
bc = ["periodic", {"derivative": 0}]
field.laplace(bc)
```

For convenience, this typical situation can be described with the special boundary condition *auto\_periodic\_neumann*, e.g., calling the Laplace operator using field.laplace("auto\_periodic\_neumann") is identical to the example above. Similarly, the special condition *auto\_periodic\_dirichlet* enforces periodic boundary conditions or Dirichlet boundary condition (vanishing value), depending on the periodicity of the underlying grid.

In summary, we have the following options for boundary conditions on a field c

Name	Condition	Example	
Dirichlet	c = 0	"dirichlet" or "value"	
	c = const	{"value": 1.5}	
	c = f(x, t)	{"value_expression": "sin(x)"}	
	c = f(x, t)	{"value_expression": func} with function	
		func(value, dx, *coords, t)	
Neumann	$\partial_n c = 0$	"neumann" or "derivative"	
	$\partial_n c = \text{const}$	{"derivative": -2}	
	$\partial_n c = f(x, t)$	{"derivative_expression": "exp(t)"}	
Robin	$\partial_n c + \text{value} \cdot c = \text{const}$	{"type": "mixed", "value": 2,	
		"const": 7}	
	$\partial_n c + \text{value} \cdot c = \text{const}$	{"type": "mixed_expression", "value":	
		"exp(t)", "const": "3 * x"}	
Curvature	$\partial_n^2 c = \text{const}$	{"curvature": 3}	
Periodic	c(0) = c(L)	"periodic"	
Anti-periodic	c(0) = -c(L)	"anti-periodic"	
Periodic or Dirichlet	c(0) = c(L)  or  c = 0	"auto_periodic_dirichlet"	
Periodic or Neu- mann	$c(0) = c(L)$ or $\partial_n c = 0$	"auto_periodic_neumann"	

Table	1:	Supported	boundary	conditions
-------	----	-----------	----------	------------

Here,  $\partial_n$  denotes a derivative in outward normal direction, f denotes an arbitrary function given by an expression (see next section), x denotes coordinates along the boundary, t denotes time.

# 3.3.2 Expressions

Expressions are strings that describe mathematical expressions. They can be used in several places, most prominently in defining PDEs using *PDE*, in creating fields using *from\_expression()*, and in defining boundary conditions; see section above. Expressions are parsed using sympy, so the expected syntax is defined by this python package. While we describe some common use cases below, it might be best to test the abilities using the *evaluate()* function.

**Warning:** To interpret arbitrary expressions, the package uses exec(). It should therefore not be used in a context where malicious input could occur.

Simple expressions can contain many standard mathematical functions, e.g.,  $sin(a) + b^{**2}$  is a valid expression. *PDE* and *evaluate()* furthermore accept differential operators defined in this package. Note that operators need to be specified with their full name, i.e., *laplace* for a scalar Laplacian and *vector\_laplace* for a Laplacian operating on a vector field. Moreover, the dot product between two vector fields can be denoted by using dot(field1, field2) in the expression, and outer(field1, field2) calculates an outer product. In this case, boundary conditons for the operators can be specified using the *bc* argument, in which case the same boundary conditions are applied to all operators. The additional argument *bc\_ops* provides a more fine-grained control, where conditions for each individual operator can be specified.

Field expressions can also directly depend on spatial coordinates. For instance, if a field is defined on a two-dimensional Cartesian grid, the variables x and y denote the local coordinates. To initialize a step profile in the x-direction, one can use either (x > 5) or heaviside (x - 5, 0.5), where the second argument denotes the returned value in case the first argument is 0. Finally, expressions for equations in *PDE* can explicitly depend on time, which is denoted by the variable t.

Expressions also support user-defined functions via the *user\_funcs* argument, which is a dictionary that maps the name of a function to an actual implementation. Finally, constants can be defined using the *consts* argument. Constants can either be individual numbers or spatially extended data, which provide values for each grid point. Note that in the latter case only the actual grid data should be supplied, i.e., the *data* attribute of a potential field class.

# 3.3.3 Custom PDE classes

To implement a new PDE in a way that all of the machinery of *py-pde* can be used, one needs to subclass *PDEBase* and overwrite at least the *evolution\_rate()* method. A simple implementation for the Kuramoto–Sivashinsky equation could read

A slightly more advanced example would allow for attributes that for instance define the boundary conditions and the diffusivity:

We here replaced the call to to\_scalar('squared\_sum') by a dot product with itself (using the @ notation), which is equivalent. Note that the numpy implementation of the right hand side of the PDE is rather slow since it runs mostly in pure python and constructs a lot of intermediate field classes. While such an implementation is helpful for testing initial ideas, actual computations should be performed with compiled PDEs as described below.

# 3.3.4 Low-level operators

This section explains how to use the low-level version of the field operators. This is necessary for the numba-accelerated implementations described above and it might be necessary to use parts of the *py-pde* package in other packages.

## **Differential operators**

Applying a differential operator to an instance of *ScalarField* is a simple as calling field.laplace(bc), where *bc* denotes the boundary conditions. Calling this method returns another *ScalarField*, which in this case contains the discretized Laplacian of the original field. The equivalent call using the low-level interface is

```
apply_laplace = field.grid.make_operator("laplace", bc)
laplace_data = apply_laplace(field.data)
```

Here, the first line creates a function apply\_laplace for the given grid field.grid and the boundary conditions *bc*. This function can be applied to numpy.ndarray instances, e.g. field.data. Note that the result of this call is again a numpy.ndarray.

Similarly, a gradient operator can be defined

```
grid = UnitGrid([6, 8])
apply_gradient = grid.make_operator("gradient", bc="auto_periodic_neumann")
data = np.random.random((6, 8))
gradient_data = apply_gradient(data)
assert gradient_data.shape == (2, 6, 8)
```

Note that this example does not even use the field classes. Instead, it directly defines a *grid* and the respective gradient operator. This operator is then applied to a random field and the resulting numpy.ndarray represents the 2-dimensional vector field.

The make\_operator method of the grids generally supports the following differential operators: 'laplacian', 'gradient', 'gradient\_squared', 'divergence', 'vector\_gradient', 'vector\_laplace', and 'tensor\_divergence'. Moreover, generic operators that perform a derivative along a single axis are supported: Specifying 'd\_dx' for instance performs a single derivative along the *x*-direction, 'd\_dy\_forward' uses a forward derivative along the *y*-direction, and 'd\_d2r' performs a second derivative in *r*-direction. A complete list of operators supported by a certain grid class can be obtained from the class property GridClass.operators. New operators can be added using the class method GridClass.register\_operator().

## **Field integration**

The integral of an instance of *ScalarField* is usually determined by accessing the property field.integral. Since the integral of a discretized field is basically a sum weighted by the cell volumes, calculating the integral using only numpy is easy:

```
cell_volumes = field.grid.cell_volumes
integral = (field.data * cell_volumes).sum()
```

Note that cell\_volumes is a simple number for Cartesian grids, but is an array for more complicated grids, where the cell volume is not uniform.

## **Field interpolation**

The fields defined in the *py-pde* package also support linear interpolation by calling field.interpolate(point). Similarly to the differential operators discussed above, this call can also be translated to code that does not use the full package:

```
grid = UnitGrid([6, 8])
interpolate = grid.make_interpolator_compiled(bc="auto_periodic_neumann")
data = np.random.random((6, 8))
value = interpolate(data, np.array([3.5, 7.9]))
```

We first create a function interpolate, which is then used to interpolate the field data at a certain point. Note that the coordinates of the point need to be supplied as a numpy.ndarray and that only the interpolation at single points is supported. However, iteration over multiple points can be fast when the loop is compiled with numba.

## **Inner products**

For vector and tensor fields, *py-pde* defines inner products that can be accessed conveniently using the @-syntax: field1 @ field2 determines the scalar product between the two fields. The package also provides an implementation for an dot-operator:

```
grid = UnitGrid([6, 8])
field1 = VectorField.random_normal(grid)
field2 = VectorField.random_normal(grid)
dot_operator = field1.make_dot_operator()
result = dot_operator(field1.data, field2.data)
assert result.shape == (6, 8)
```

Here, result is the data of the scalar field resulting from the dot product.

# 3.3.5 Numba-accelerated PDEs

The compiled operators introduced in the previous section can be used to implement a compiled method for the evolution rate of PDEs. As an example, we now extend the class KuramotoSivashinskyPDE introduced above:

```
from pde.tools.numba import jit
class KuramotoSivashinskyPDE (PDEBase) :
   def __init__(self, diffusivity=1, bc="auto_periodic_neumann", bc_laplace="auto_

→periodic_neumann"):

        """ initialize the class with a diffusivity and boundary conditions
        for the actual field and its second derivative """
       self.diffusivity = diffusivity
       self.bc = bc
        self.bc_laplace = bc_laplace
    def evolution_rate(self, state, t=0):
        """ numpy implementation of the evolution equation """
        state_lapacian = state.laplace(bc=self.bc)
        state_gradient = state.gradient(bc="auto_periodic_neumann")
        return (- state_lapacian.laplace(bc=self.bc_laplace)
                - state_lapacian
                - 0.5 * self.diffusivity * (state_gradient @ state_gradient))
   def _make_pde_rhs_numba(self, state):
        """ the numba-accelerated evolution equation """
        # make attributes locally available
        diffusivity = self.diffusivity
        # create operators
        laplace_u = state.grid.make_operator("laplace", bc=self.bc)
        gradient_u = state.grid.make_operator("gradient", bc=self.bc)
        laplace2_u = state.grid.make_operator("laplace", bc=self.bc_laplace)
        dot = VectorField(state.grid).make_dot_operator()
        0jit
        def pde_rhs(state_data, t=0):
            """ compiled helper function evaluating right hand side """
            state_lapacian = laplace_u(state_data)
            state_grad = gradient_u(state_data)
            return (- laplace2_u(state_lapacian)
                    - state_lapacian
                    - diffusivity / 2 * dot(state_grad, state_grad))
        return pde_rhs
```

To activate the compiled implementation of the evolution rate, we simply have to overwrite the \_make\_pde\_rhs\_numba() method. This method expects an example of the state class (e.g., an instance of *ScalarField*) and returns a function that calculates the evolution rate. The *state* argument is necessary to define the grid and the dimensionality of the data that the returned function is supposed to be handling. The implementation of the compiled function is split in several parts, where we first copy the attributes that are required by the implementation. This is necessary, since numba freezes the values when compiling the function, so that in the example above the diffusivity cannot be altered without recompiling. In the next step, we create all operators that we need subsequently. Here, we use the boundary conditions defined by the attributes, which requires two different laplace operators, since their boundary

conditions might differ. In the last step, we define the actual implementation of the evolution rate as a local function that is compiled using the jit decorator. Here, we use the implementation shipped with *py-pde*, which sets some default values. However, we could have also used the usual numba implementation. It is important that the implementation of the evolution rate only uses python constructs that numba can compile.

One advantage of the numba compiled implementation is that we can now use loops, which will be much faster than their python equivalents. For instance, we could have written the dot product in the last line as an explicit loop:

```
[...]
   def _make_pde_rhs_numba(self, state):
       """ the numba-accelerated evolution equation """
       # make attributes locally available
       diffusivity = self.diffusivity
       # create operators
       laplace_u = state.grid.make_operator("laplace", bc=self.bc)
       gradient_u = state.grid.make_operator("gradient", bc=self.bc)
       laplace2_u = state.grid.make_operator("laplace", bc=self.bc_laplace)
       dot = VectorField(state.grid).make_dot_operator()
       dim = state.grid.dim
       0jit
       def pde_rhs(state_data, t=0):
           """ compiled helper function evaluating right hand side """
           state_lapacian = laplace_u(state_data)
           state_grad = gradient_u(state_data)
           result = - laplace2_u(state_lapacian) - state_lapacian
           for i in range(state_data.size):
               for j in range(dim):
                    result.flat[i] -= diffusivity / 2 * state_grad[j].flat[i]**2
           return result
       return pde_rhs
```

Here, we extract the total number of elements in the state using its size attribute and we obtain the dimensionality of the space from the grid attribute dim. Note that we access numpy arrays using their flat attribute to provide an implementation that works for all dimensions.

## 3.3.6 Configuration parameters

Configuration parameters affect how the package behaves. They can be set using a dictionary-like interface of the configuration config, which can be imported from the base package. Here is a list of all configuration options that can be adjusted in the package:

### numba.debug

Determines whether numba uses the debug mode for compilation. If enabled, this emits extra information that might be useful for debugging. (Default value: False)

### numba.fastmath

Determines whether the fastmath flag is set during compilation. If enabled, some mathematical operations might be faster, but less precise. This flag does not affect infinity detection and NaN handling. (Default value: True)

### numba.multithreading

Determines whether multiple threads are used in numba-compiled code. Enabling this option accelerates a small subset of operators applied to fields defined on large grids. (Default value: True)

#### numba.multithreading\_threshold

Minimal number of support points of grids before multithreading is enabled in numba compilations. Has no effect when `numba.multithreading` is `False`. (**Default value: 65536**)

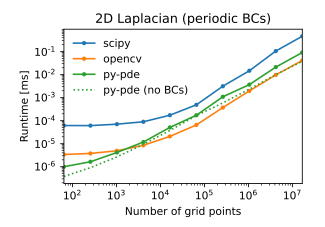
Tip: To disable parallel computing in the package, the following code could be added to the start of the script:

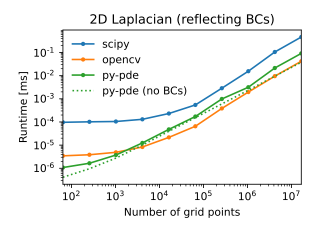
```
from pde import config
config["numba.multithreading"] = False
# actual code using py-pde
```

# 3.4 Performance

## 3.4.1 Measuring performance

The performance of the *py-pde* package depends on many details and general statements are thus difficult to make. However, since the core operators are just-in-time compiled using numba, many operations of the package proceed at performances close to most compiled languages. For instance, a simple Laplace operator applied to fields defined on a Cartesian grid has performance that is similar to the operators supplied by the popular OpenCV package. The following figures illustrate this by showing the duration of evaluating the Laplacian on grids of increasing number of support points for two different boundary conditions (lower duration is better):





Note that the call overhead is lower in the *py-pde* package, so that the performance on small grids is particularly good. However, realistic use-cases probably need more complicated operations and it is thus always necessary to profile the respective code. This can be done using the function <code>estimate\_computation\_speed()</code> or the traditional timeit, profile, or even more sophisticated profilers like pyinstrument.

# 3.4.2 Improving performance

Beside the underlying implementation of the operators, a major factor for performance is numerical problem at hand and the methods that are used to solve it. As a rule of thumb, simulations run faster when there are fewer degrees of freedom. In the case of partial differential equations, this often means using a coarser grid with fewer support points. However, there often also is an lower bound to the number of support points if structures of a certain length scales need to be resolved. Reducing the number of support points not only reduces the number of variables to be treated, but it can also allow for larger time steps. This is particularly transparent for the simple diffusion equation, where a von Neumann stability analysis reveals that the maximal time step scales as one over the discretization length squared! Choosing the right time step obviously also affects performance of a simulation. The package supports automatic choice of suitable time steps, using adaptive stepping schemes. To enable those, it's best to specify an initial time step, like so

eq.solve(t\_range=10, dt=1e-3, adaptive=True)

An additional advantage of this choice is that it selects *ExplicitSolver*, which is also compiled with numba for speed. Alternatively, if only *t\_range* is specified, the generic scipy-solver *ScipySolver*, which can be significantly slower.

Additional factors influencing the performance of the package include the compiler used for numpy, scipy, and of course numba. Moreover, the BLAS and LAPACK libraries might make a difference. The package has some basic support for multithreading, which can be accelerated using the *Threading Building Blocks* library. Finally, it can help to install the intel short vector math library (SVML). However, this is not distributed with **macports** and might thus be more difficult to enable.

Using macports, one could for instance install the following variants of typical packages

```
port install py37-numpy +gcc8+openblas
port install py37-scipy +gcc8+openblas
port install py37-numba +tbb
```

Note that you can disable the automatic multithreading via Configuration parameters.

# 3.4.3 Multiprocessing using MPI

The package also supports parallel simulations of PDEs using the Message Passing Interface (MPI), which allows combining the power of CPU cores that do not share memory. To use this advanced simulation technique, a working implementation of MPI needs to be installed on the computer. Usually, this is done automatically, when the optional package numba-mpi is installed via *pip* or *conda*.

To run simulations in parallel, the special solver *ExplicitMPISolver* needs to be used and the entire script needs to be started using mpiexec. Taken together, a minimal example reads

```
from pde import DiffusionPDE, ScalarField, UnitGrid
grid = UnitGrid([64, 64])
state = ScalarField.random_uniform(grid, 0.2, 0.3)
eq = DiffusionPDE(diffusivity=0.1)
result = eq.solve(state, t_range=10, dt=0.1, method="explicit_mpi")
if result is not None: # restrict the output to the main node
    result.plot()
```

Saving this script as *multiprocessing.py*, we can evoke a parallel simulation using

mpiexec -n 2 python3 multiprocessing.py

Here, the number 2 determines the number of cores that will be used. Note that macOS might require an additional hint on how to connect the processes even when they are run on the same machine (e.g., your workstation). It might help to run mpiexec  $-n \ 2$  -host localhost: 2 python3 multiprocessing.py in this case.

In the example above, two python processes will start in parallel and run independently at first. In particular, both processes will load all packages and create the initial *state* field as well as the PDE class *eq*. Once the *explicit\_mpi* solver is evoked, the processes will start communicating. *py-pde* will split up the full grid into two sub-grids, in this case of shape 32x64, distribute the associated sub-fields to both processes and ask each process to evolve the PDE for their sub-field. Note that boundary conditions are treated and boundary values are exchanged between neighboring sub-grids automatically. To avoid confusion, trackers will only be used on one process and also the result is only returned in one process to avoid problems where multiple process write data simultaneously. Consequently, the example above checked whether *result is None* (in which case the corresponding process is a child process) and only resumes analysis when the result is actually present.

The automatic treatment tries to use sensible default values, so typical simulations work out of the box. However, in some situations it might be advantageous to adjust these values. For instance, the decomposition of the grid can be affected by an argument *decomposition*, which can be passed to the *solve()* method or the *ExplicitMPISolver*. The argument should be a list with one integer for each axis in the grid, which specifies how often the particular axis is divided.

**Warning:** The automatic division of the grid into sub-grids can lead to unexpected behavior, particularly in custom PDEs that were not designed for this use case. As a rule of thumb, all local operations are fine (since they can be performed on each subgrid), while global operations might need synchronization between all subgrids. One example is integration, which has been implemented properly in *py-pde*. Consequently, it is safe to use *integral*.

# 3.5 Contributing code

# 3.5.1 Structure of the package

The functionality of the *pde* package is split into multiple sub-package. The domain, together with its symmetries, periodicities, and discretizations, is described by classes defined in *grids*. Discretized fields are represented by classes in *fields*, which have methods for differential operators with various boundary conditions collected in *boundaries*. The actual pdes are collected in *pdes* and the respective solvers are defined in *solvers*.

# 3.5.2 Extending functionality

All code is build on a modular basis, making it easy to introduce new classes that integrate with the rest of the package. For instance, it is simple to define a new partial differential equation by subclassing *PDEBase*. Alternatively, PDEs can be defined by specifying their evolution rates using mathematical expressions by creating instances of the class *PDE*. Moreover, new grids can be introduced by subclassing *GridBase*. It is also possible to only use parts of the package, e.g., the discretized differential operators from *operators*.

New operators can be associated with grids by registering them using *register\_operator()*. For instance, to create a new operator for the cylindrical grid one needs to define a factory function that creates the operator. This factory function takes an instance of *Boundaries* as an argument and returns a function that takes as an argument the actual data array for the grid. Note that the grid itself is an attribute of *Boundaries*. This operator would be registered with the grid by calling CylindricalSymGrid.register\_operator("operator", make\_operator), where the first argument is the name of the operator and the second argument is the factory function.

# 3.5.3 Design choices

The data layout of field classes (subclasses of *FieldBase*) was chosen to allow for a simple decomposition of different fields and tensor components. Consequently, the data is laid out in memory such that spatial indices are last. For instance, the data of a vector field field defined on a 2d Cartesian grid will have three dimensions and can be accessed as field.data[vector\_component, x, y], where vector\_component is either 0 or 1.

# 3.5.4 Coding style

The coding style is enforced using isort and black. Moreover, we use Google Style docstrings, which might be best learned by example. The documentation, including the docstrings, are written using reStructuredText, with examples in the following cheatsheet. To ensure the integrity of the code, we also try to provide many test functions, contained in the separate sub-folder tests. These tests can be ran using scripts in the scripts subfolder in the root folder. This folder also contain a script tests\_types.sh, which uses mypy to check the consistency of the python type annotations. We use these type annotations for additional documentation and they have also already been useful for finding some bugs.

We also have some conventions that should make the package more consistent and thus easier to use. For instance, we try to use properties instead of getter and setter methods as often as possible. Because we use a lot of numba justin-time compilation to speed up computations, we need to pass around (compiled) functions regularly. The names of the methods and functions that make such functions, i.e. that return callables, should start with 'make\_\*' where the wildcard should describe the purpose of the function being created.

# 3.5.5 Running unit tests

The pde package contains several unit tests, collection in the tests folder in the project root. These tests ensure that basic functions work as expected, in particular when code is changed in future versions. To run all tests, there are a few convenience scripts in the root directory scripts. The most basic script is tests\_run.sh, which uses pytest to run the tests. Clearly, the python package pytest needs to be installed. There are also additional scripts that for instance run tests in parallel (needs the python package pytest\_xdist installed), measure test coverage (needs package pytest\_cov installed), and make simple performance measurements. Moreover, there is a script test\_types.sh, which uses mypy to check the consistency of the python type annotations and there is a script format\_code.sh, which formats the code automatically to adhere to our style.

Before committing a change to the code repository, it is good practice to run the tests, check the type annotations, and the coding style with the scripts described above.

# 3.6 Citing the package

To cite or reference *py-pde* in other work, please refer to the publication in the Journal of Open Source Software. Here are the respective bibliographic records in Bibtex format:

```
@article{py-pde,
Author = {David Zwicker},
Doi = {10.21105/joss.02158},
Journal = {Journal of Open Source Software},
Number = {48},
Pages = {2158},
Publisher = {The Open Journal},
Title = {py-pde: A Python package for solving partial differential equations},
Url = {https://doi.org/10.21105/joss.02158},
Volume = {5},
Year = {2020}
}
```

and in RIS format:

```
TY - JOUR
AU - Zwicker, David
JO - Journal of Open Source Software
IS - 48
SP - 2158
PB - The Open Journal
T1 - py-pde: A Python package for solving partial differential equations
UR - https://doi.org/10.21105/joss.02158
VL - 5
PY - 2020
```

# 3.7 Code of Conduct

# 3.7.1 Our Pledge

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

# 3.7.2 Our Standards

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

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

Examples of unacceptable behavior by participants include:

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

# 3.7.3 Our Responsibilities

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

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

# 3.7.4 Scope

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

# 3.7.5 Enforcement

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

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

# 3.7.6 Attribution

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

For answers to common questions about this code of conduct, see https://www.contributor-covenant.org/faq

# CHAPTER

FOUR

# **REFERENCE MANUAL**

The py-pde package provides classes and methods for solving partial differential equations.

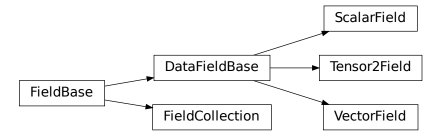
Subpackages:

# 4.1 pde.fields package

Defines fields, which contain the actual data stored on a discrete grid.

ScalarField	Scalar field discretized on a grid
VectorField	Vector field discretized on a grid
Tensor2Field	Tensor field of rank 2 discretized on a grid
FieldCollection	Collection of fields defined on the same grid

Inheritance structure of the classes:



The details of the classes are explained below:

## 4.1.1 pde.fields.base module

Defines base classes of fields, which are discretized on grids

#### **class DataFieldBase** (grid, data='zeros', \*, label=None, dtype=None, with\_ghost\_cells=False)

Bases: FieldBase

abstract base class for describing fields of single entities

#### Parameters

- grid (GridBase) Grid defining the space on which this field is defined.
- data (Number or ndarray, optional) Field values at the support points of the grid. The flag with\_ghost\_cells determines whether this data array contains values for the ghost cells, too. The resulting field will contain real data unless the *data* argument contains complex values. Special values are "zeros" or None, initializing the field with zeros, and "empty", just allocating memory with unspecified values.
- label (str, optional) Name of the field
- **dtype** (*numpy* dtype) The data type of the field. If omitted, it will be determined from *data* automatically.
- with\_ghost\_cells (bool) Indicates whether the ghost cells are included in data

**apply\_operator** (*operator*, *bc*, *out=None*, \*, *label=None*, *args=None*, \*\**kwargs*)

apply a (differential) operator and return result as a field

#### **Parameters**

- **operator** (*str*) An identifier determining the operator. Note that not all grids support the same operators.
- bc (BoundariesData / None) Boundary conditions applied to the field before applying the operator. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by *['value': NUM]*) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by *['derivative': DERIV]*) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documentation*. If the special value *None* is given, no boundary conditions are enforced. The user then needs to ensure that the ghost cells are set accordingly.
- **out** (*DataFieldBase*, optional) Optional field to which the result is written.
- **label** (*str*, *optional*) Name of the returned field
- args (dict) Additional arguments for the boundary conditions
- **\*\*kwargs** Additional arguments affecting how the operator behaves.

#### Returns

Field data after applying the operator. This field is identical to *out* if this argument was specified.

#### **Return type**

DataFieldBase

#### property average: int | float | complex | ndarray

the average of data

This is calculated by integrating each component of the field over space and dividing by the grid volume

### Туре

float or ndarray

copy (\*, label=None, dtype=None)

return a new field with the data (but not the grid) copied

#### Parameters

- label (str, optional) Name of the returned field
- **dtype** (*numpy dtype*) The data type of the field. If omitted, it will be determined from *data* automatically or the dtype of the current field is used.
- **self**(TDataField)-

#### Returns

A copy of the current field

#### **Return type**

DataFieldBase

### property data\_shape: tuple[int, ...]

the shape of the data at each grid point

### Type

tuple

### property fluctuations: int | float | complex | ndarray

quantification of the average fluctuations

The fluctuations are defined as the standard deviation of the data scaled by the cell volume. This definition makes the fluctuations independent of the discretization. It corresponds to the physical scaling available in the random\_normal().

#### Returns

A tensor with the same rank of the field, specifying the fluctuations of each component of the tensor field individually. Consequently, a simple scalar is returned for a *ScalarField*.

#### Return type

ndarray

#### Туре

float or ndarray

classmethod from\_state(attributes, data=None)

create a field from given state.

### Parameters

- attributes (dict) The attributes that describe the current instance
- data (ndarray, optional) Data values at the support points of the grid defining the field

### Returns

The instance created from the stored state

#### **Return type**

DataFieldBase

### get\_boundary\_values (axis, upper, bc=None)

get the field values directly on the specified boundary

### Parameters

- **axis** (*int*) The axis perpendicular to the boundary
- upper (bool) Whether the boundary is at the upper side of the axis
- bc (BoundariesData / None) The boundary conditions applied to the field. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by *{'value': NUM}*) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by *{'derivative': DERIV}*) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documentation*. If the special value *None* is given, no boundary conditions are enforced. The user then needs to ensure that the ghost cells are set accordingly.

### Returns

The discretized values on the boundary

### **Return type**

ndarray

## classmethod get\_class\_by\_rank(rank)

return a DataFieldBase subclass describing a field with a given rank

### Parameters

rank (int) - The rank of the tensor field

Returns

The DataField class that corresponds to the rank

### **Return type**

type[DataFieldBase]

get\_image\_data (scalar='auto', transpose=False, \*\*kwargs)

return data for plotting an image of the field

### Parameters

- **scalar** (*str* or *int*) The method for extracting scalars as described in DataFieldBase.to\_scalar().
- transpose (bool) Determines whether the transpose of the data should is plotted
- \*\*kwargs Additional parameters are forwarded to grid.get\_image\_data

### Returns

Information useful for plotting an image of the field

### **Return type**

dict

### get\_line\_data (scalar='auto', extract='auto')

return data for a line plot of the field

### Parameters

- **scalar** (*str* or *int*) The method for extracting scalars as described in DataFieldBase.to\_scalar().
- **extract** (*str*) The method used for extracting the line data. See the docstring of the grid method *get\_line\_data* to find supported values.

#### Returns

Information useful for performing a line plot of the field

**Return type** 

dict

get\_vector\_data (transpose=False, \*\*kwargs)

return data for a vector plot of the field

#### **Parameters**

• **\*\*kwargs** – Additional parameters are forwarded to grid.get\_image\_data

• transpose (bool) -

#### Returns

Information useful for plotting an vector field

Return type

dict

### insert (point, amount)

adds an (integrated) value to the field at an interpolated position

### Parameters

- **point** (ndarray) The point inside the grid where the value is added. This is given in grid coordinates.
- **amount** (Number or ndarray) The amount that will be added to the field. The value describes an integrated quantity (given by the field value times the discretization volume). This is important for consistency with different discretizations and in particular grids with non-uniform discretizations.

#### **Return type**

None

abstract property integral: int | float | complex | ndarray

integral of the scalar field over space

#### interpolate (point, \*, bc=None, fill=None)

interpolate the field to points between support points

#### Parameters

- **point** (ndarray) The points at which the values should be obtained. This is given in grid coordinates.
- **bc** (BoundariesData / None) The boundary conditions applied to the field, which affects values close to the boundary. If omitted, the argument *fill* is used to determine values outside the domain. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by *[value': NUM]*) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by

*{'derivative': DERIV}*) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documentation*. If the special value *None* is given, no boundary conditions are enforced. The user then needs to ensure that the ghost cells are set accordingly.

• **fill** (*Number*, *optional*) – Determines how values out of bounds are handled. If None, a *ValueError* is raised when out-of-bounds points are requested. Otherwise, the given value is returned.

#### Returns

the values of the field

#### **Return type**

ndarray

interpolate\_to\_grid (grid, \*, bc=None, fill=None, label=None)

interpolate the data of this field to another grid.

#### **Parameters**

- grid (GridBase) The grid of the new field onto which the current field is interpolated.
- bc (BoundariesData / None) The boundary conditions applied to the field, which affects values close to the boundary. If omitted, the argument *fill* is used to determine values outside the domain. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by *{'value': NUM}*) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by *{'derivative': DERIV}*) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documentation*. If the special value *None* is given, no boundary conditions are enforced. The user then needs to ensure that the ghost cells are set accordingly.
- **fill** (*Number*, *optional*) Determines how values out of bounds are handled. If *None*, a *ValueError* is raised when out-of-bounds points are requested. Otherwise, the given value is returned.
- label (str, optional) Name of the returned field
- **self**(*TDataField*) -

#### Returns

Field of the same rank as the current one.

## Return type

TDataField

#### property magnitude: float

determine the (scalar) magnitude of the field

This is calculated by getting a scalar field using the default arguments of the  $to\_scalar()$  method, averaging the result over the whole grid, and taking the absolute value.

#### Туре

float

#### make\_dot\_operator(backend='numba', \*, conjugate=True)

return operator calculating the dot product between two fields

This supports both products between two vectors as well as products between a vector and a tensor.

### Parameters

- backend (str) Can be numba or numpy, deciding how the function is constructed
- conjugate (bool) Whether to use the complex conjugate for the second operand

### Returns

function that takes two instance of ndarray, which contain the discretized data of the two operands. An optional third argument can specify the output array to which the result is written.

### **Return type**

Callable[[ndarray, ndarray, ndarray | None], ndarray]

### make\_interpolator(\*, fill=None, with\_ghost\_cells=False)

returns a function that can be used to interpolate values.

### Parameters

- **fill** (*Number*, *optional*) Determines how values out of bounds are handled. If *None*, a *ValueError* is raised when out-of-bounds points are requested. Otherwise, the given value is returned.
- with\_ghost\_cells (bool) Flag indicating that the interpolator should work on the full data array that includes values for the ghost points. If this is the case, the boundaries are not checked and the coordinates are used as is.

### Returns

A function which returns interpolated values when called with arbitrary positions within the space of the grid.

### **Return type**

*Callable*[[*ndarray*, *ndarray*], int | float | complex | *ndarray*]

visualize the field

## Parameters

- kind (*str*) Determines the visualizations. Supported values are *image*, *line*, *vector*, or *interactive*. Alternatively, *auto* determines the best visualization based on the field itself.
- **title** (*str*) Title of the plot. If omitted, the title might be chosen automatically.
- filename (str, optional) If given, the plot is written to the specified file.
- **action** (*str*) Decides what to do with the final figure. If the argument is set to *show*, matplotlib.pyplot.show() will be called to show the plot. If the value is *none*, the figure will be created, but not necessarily shown. The value *close* closes the figure, after saving it to a file when *filename* is given. The default value *auto* implies that the plot is shown if it is not a nested plot call.
- **ax\_style** (*dict*) Dictionary with properties that will be changed on the axis after the plot has been drawn by calling matplotlib.pyplot.setp(). A special item i this dictionary is *use\_offset*, which is flag that can be used to control whether offset are shown along the axes of the plot.
- **fig\_style** (*dict*) Dictionary with properties that will be changed on the figure after the plot has been drawn by calling matplotlib.pyplot.setp(). For instance, using fig\_style={'dpi': 200} increases the resolution of the figure.

- **ax** (matplotlib.axes.Axes) Figure axes to be used for plotting. The special value "create" creates a new figure, while "reuse" attempts to reuse an existing figure, which is the default.
- \*\*kwargs All additional keyword arguments are forwarded to the actual plotting function determined by kind.

### Returns

Instance that contains information to update the plot with new data later.

Return type *PlotReference* 

Tip: Typical additional arguments for the various plot kinds include

- kind == "line":
  - scalar: Sets method for extracting scalars as described in DataFieldBase.to\_scalar().
  - *extract*: Method used for extracting the line data.
  - ylabel: Label of the y-axis.
  - ylim: Data limits of the y-axis.
  - Additional arguments are passed to matplotlib.pyplot.plot()
- kind == "image":
  - colorbar: Determines whether a colorbar is shown
  - scalar: Sets method for extracting scalars as described in DataFieldBase.to\_scalar().
  - transpose Determines whether the transpose of the data is plotted
  - Most remaining arguments are passed to matplotlib.pyplot.imshow()
- kind == `"vector":
  - method Can be either quiver or streamplot
  - transpose Determines whether the transpose of the data is plotted
  - max\_points Sets max. number of points along each axis in quiver plots
  - Additional arguments are passed to matplotlib.pyplot.quiver() or matplotlib. pyplot.streamplot().

**classmethod random\_colored** (*grid, exponent=0, scale=1, \*, label=None, dtype=None, rng=None*) create a field of random values with colored noise

The spatially correlated values obey

$$\langle c_i(\mathbf{k})c_j(\mathbf{k}')\rangle = \Gamma^2 |\mathbf{k}|^{\nu} \delta_{ij} \delta(\mathbf{k} - \mathbf{k}')$$

in spectral space, where k is the wave vector. The special case  $\nu = 0$  corresponds to white noise. Note that the spatial correlations always assume periodic boundary conditions (even if the underlying grid does not) and that the components of tensor fields are uncorrelated.

#### **Parameters**

• grid (GridBase) - Grid defining the space on which this field is defined

- **exponent** (*float*) Exponent  $\nu$  of the power spectrum
- scale (float) Scaling factor  $\Gamma$  determining noise strength
- label (str, optional) Name of the returned field
- **dtype** (*numpy dtype*) The data type of the field. If omitted, it defaults to *double*.
- **rng** (Generator) Random number generator (default: default\_rng())

#### **Return type**

TDataField

classmethod random\_harmonic (grid, modes=3, harmonic=<ufunc 'cos'>, axis\_combination=<ufunc 'multiply'>, \*, label=None, dtype=None, rng=None)

create a random field build from harmonics

The resulting fields will be highly correlated in space and can thus serve for testing differential operators.

With the default settings, the resulting field  $c_i(\mathbf{x})$  is given by

$$c_i(\mathbf{x}) = \prod_{\alpha=1}^N \sum_{j=1}^M a_{ij\alpha} \cos\left(\frac{2\pi x_\alpha}{jL_\alpha}\right) ,$$

where N is the number of spatial dimensions, each with length  $L_{\alpha}$ , M is the number of modes given by *modes*, and  $a_{ij\alpha}$  are random amplitudes, chosen from a uniform distribution over the interval [0, 1].

Note that the product could be replaced by a sum when  $axis\_combination = numpy.add$  and the cos() could be any other function given by the parameter *harmonic*.

#### **Parameters**

- grid (GridBase) Grid defining the space on which this field is defined
- modes (*int*) Number M of harmonic modes
- harmonic (*callable*) Determines which harmonic function is used. Typical values are numpy.sin() and numpy.cos(), which basically relate to different boundary conditions applied at the grid boundaries.
- **axis\_combination** (*callable*) Determines how values from different axis are combined. Typical choices are numpy.multiply() and numpy.add() resulting in products and sums of the values along axes, respectively.
- **label** (*str*, *optional*) Name of the returned field
- dtype (numpy dtype) The data type of the field. If omitted, it defaults to double.
- **rng** (Generator) Random number generator (default: default\_rng())

#### **Return type**

TDataField

create field with normal distributed random values

These values are uncorrelated in space. A complex field is returned when either *mean* or *std* is a complex number. In this case, the real and imaginary parts of these arguments are used to determine the distribution of the real and imaginary parts of the resulting field. Consequently, ScalarField.random\_normal(grid, 0, 1 + 1j) creates a complex field where the real and imaginary parts are chosen from a standard normal distribution.

- grid (GridBase) Grid defining the space on which this field is defined
- mean (float) Mean of the Gaussian distribution
- std (float) Standard deviation of the Gaussian distribution.
- **scaling** (*str*) Determines how the values are scaled. Possible choices are 'none' (values are drawn from a normal distribution with given mean and standard deviation) or 'physical' (the variance of the random number is scaled by the inverse volume of the grid cell; this is for instance useful for concentration fields, which vary less in larger volumes).
- label (str, optional) Name of the returned field
- **dtype** (*numpy dtype*) The data type of the field. If omitted, it defaults to *double* if both *mean* and *std* are real, otherwise it is *complex*.
- rng (Generator) Random number generator (default: default\_rng())

#### **Return type**

TDataField

**classmethod random\_uniform** (grid, vmin=0, vmax=1, \*, label=None, dtype=None, rng=None) create field with uniform distributed random values

These values are uncorrelated in space.

#### Parameters

- grid (GridBase) Grid defining the space on which this field is defined
- **vmin** (*float*) Smallest possible random value
- **vmax** (*float*) Largest random value
- label (str, optional) Name of the returned field
- **dtype** (*numpy dtype*) The data type of the field. If omitted, it defaults to *double* if both *vmin* and *vmax* are real, otherwise it is *complex*.
- rng (Generator) Random number generator (default: default\_rng())

#### Return type

TDataField

## rank: int

## set\_ghost\_cells (bc, \*, set\_corners=False, args=None)

set the boundary values on virtual points for all boundaries

- **bc** (*str* or *list* or *tuple* or *dict*) The boundary conditions applied to the field. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by *{'value': NUM})* and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by *{'derivative': DERIV})* are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documentation*.
- **set\_corners** (*bool*) Determines whether the corner cells are set using interpolation
- args Additional arguments that might be supported by special boundary conditions.

#### Return type None

#### smooth (sigma=1, \*, out=None, label=None)

applies Gaussian smoothing with the given standard deviation

This function respects periodic boundary conditions of the underlying grid, using reflection when no periodicity is specified.

#### **Parameters**

- sigma (float) Gives the standard deviation of the smoothing in real length units (default:
   1)
- **out** (FieldBase, *optional*) Optional field into which the smoothed data is stored. Setting this to the input field enables in-place smoothing.
- label (str, optional) Name of the returned field
- **self**(TDataField)-

## Returns

Field with smoothed data. This is stored at out if given.

## Return type

TDataField

abstract to\_scalar(scalar='auto', \*, label=None)

return scalar variant of the field

#### **Parameters**

- scalar (str)-
- label (str / None) -

## **Return type**

ScalarField

## classmethod unserialize\_attributes (attributes)

unserializes the given attributes

## Parameters

attributes (dict) – The serialized attributes

## Returns

The unserialized attributes

## Return type

dict

## class FieldBase (grid, data, \*, label=None)

# Bases: object

abstract base class for describing (discretized) fields

- grid (GridBase) Grid defining the space on which this field is defined
- data (ndarray, optional) Field values at the support points of the grid and the ghost cells
- label (str, optional) Name of the field

**apply** (*func*, *out=None*, \*, *label=None*, *evaluate\_args=None*)

applies a function/expression to the data and returns it as a field

### **Parameters**

- **func** (*callable or str*) The (vectorized) function being applied to the data or an expression that can be parsed using sympy (evaluate() is used in this case). The local field values can be accessed using the field labels for a field collection and via the variable *c* otherwise.
- **out** (FieldBase, *optional*) Optional field into which the data is written
- label (str, optional) Name of the returned field
- evaluate\_args (dict) Additional arguments passed to evaluate(). Only used when *func* is a string.
- self(TField)-

#### Returns

Field with new data. Identical to out if given.

#### Return type

FieldBase

#### assert\_field\_compatible (other, accept\_scalar=False)

checks whether other is compatible with the current field

#### **Parameters**

- other (FieldBase) The other field this one is compared to
- accept\_scalar(bool, optional) Determines whether it is acceptable that other is an instance of ScalarField.

## **Return type**

None

## property attributes: dict[str, Any]

describes the state of the instance (without the data)

# Туре

dict

## property attributes\_serialized: dict[str, str]

serialized version of the attributes

#### Туре

dict

## conjugate()

returns complex conjugate of the field

#### Returns

the complex conjugated field

Return type

## FieldBase

Parameters

self(TField) -

abstract copy (\*, label=None, dtype=None)

return a new field with the data (but not the grid) copied

## **Parameters**

- label (str, optional) Name of the returned field
- **dtype** (*numpy dtype*) The data type of the field. If omitted, it will be determined from *data* automatically or the dtype of the current field is used.
- **self**(*TField*)-

#### Returns

A copy of the current field

#### **Return type**

DataFieldBase

## property data: ndarray

discretized data at the support points

Туре

ndarray

```
property dtype: dtype[Any] | None | type[Any] | _SupportsDType[dtype[Any]]
| str | tuple[Any, int] | tuple[Any, SupportsIndex |
Sequence[SupportsIndex]] | list[Any] | _DTypeDict | tuple[Any, Any]
```

the numpy dtype of the underlying data

Type

DTypeLike

classmethod from\_file(filename)

create field from data stored in a file

Field can be written to a file using *FieldBase.to\_file()*.

## Example

Write a field to a file and then read it back:

```
field = pde.ScalarField(...)
field.write_to("test.hdf5")
```

field\_copy = pde.FieldBase.from\_file("test.hdf5")

## Parameters

**filename** (*str*) – Path to the file being read

Returns

The field with the appropriate sub-class

Return type

FieldBase

## classmethod from\_state(attributes, data=None)

create a field from given state.

- **attributes** (*dict*) The attributes that describe the current instance
- data (ndarray, optional) Data values at the support points of the grid defining the field

The field created from the state

## **Return type**

FieldBase

## abstract get\_image\_data()

return data for plotting an image of the field

## **Parameters**

- **scalar** (*str* or *int*) The method for extracting scalars as described in DataFieldBase.to\_scalar().
- **transpose** (bool) Determines whether the transpose of the data should is plotted
- \*\*kwargs Additional parameters are forwarded to grid.get\_image\_data

#### Returns

Information useful for plotting an image of the field

## **Return type**

dict

abstract get\_line\_data (scalar='auto', extract='auto')

return data for a line plot of the field

## Parameters

- **scalar** (*str* or *int*) The method for extracting scalars as described in DataFieldBase.to\_scalar().
- **extract** (*str*) The method used for extracting the line data. See the docstring of the grid method *get\_line\_data* to find supported values.

#### Returns

Information useful for performing a line plot of the field

#### **Return type**

dict

## property grid: GridBase

The grid on which the field is defined

## Type

base,GridBase

## property imag: TField

Imaginary part of the field

#### Туре

FieldBase

## property is\_complex: bool

whether the field contains real or complex data

Туре

bool

property label: str | None

the name of the field

Туре

str

abstract plot(\*args, \*\*kwargs)

visualize the field

plot\_interactive (viewer\_args=None, \*\*kwargs)

create an interactive plot of the field using napari

For a detailed description of the launched program, see the napari webpage.

#### **Parameters**

- **viewer\_args** (*dict*) Arguments passed to napari.viewer.Viewer to affect the viewer.
- \*\*kwargs Extra arguments passed to the plotting function

Return type None

property real: TField

Real part of the field

Type

FieldBase

```
split_mpi(decomposition=-1)
```

splits the field onto subgrids in an MPI run

In a normal serial simulation, the method simply returns the field itself. In contrast, in an MPI simulation, the field provided on the main node is split onto all nodes using the given decomposition. The field data provided on all other nodes is not used.

## **Parameters**

• **decomposition** (*list of ints*) – Number of subdivision in each direction. Should be a list of length *field.grid.num\_axes* specifying the number of nodes for this axis. If one value is -1, its value will be determined from the number of available nodes. The default value decomposed the first axis using all available nodes

self(TField) -

#### Returns

The part of the field that corresponds to the subgrid associated with the current MPI node.

# Return type

FieldBase

to\_file (filename, \*\*kwargs)

store field in a file

The extension of the filename determines what format is being used. If it ends in .h5 or .hdf, the Hierarchical Data Format is used. The other supported format are images, where only the most typical formats are supported.

To load the field back from the file, you may use FieldBase.from\_file().

Example

Write a field to a file and then read it back:

```
field = pde.ScalarField(...)
field.write_to("test.hdf5")
field_copy = pde.FieldBase.from_file("test.hdf5")
```

#### **Parameters**

- filename (*str*) Path where the data is stored
- \*\*kwargs Additional parameters may be supported for some formats

## **Return type**

None

## classmethod unserialize\_attributes(attributes)

unserializes the given attributes

## Parameters

attributes (dict) – The serialized attributes

Returns The unserialized attributes

**Return type** 

dict

property writeable: bool

whether the field data can be changed or not

Type bool

#### exception RankError

#### Bases: TypeError

error indicating that the field has the wrong rank

# 4.1.2 pde.fields.collection module

Defines a collection of fields to represent multiple fields defined on a common grid.

class FieldCollection (fields, \*, copy\_fields=False, label=None, labels=None, dtype=None)

Bases: FieldBase

Collection of fields defined on the same grid

**Note:** All fields in a collection must have the same data type. This might lead to up-casting, where for instance a combination of a real-valued and a complex-valued field will be both stored as complex fields.

#### **Parameters**

• **fields** (sequence or mapping of DataFieldBase) - Sequence or mapping of the individual fields. If a mapping is used, the keys set the names of the individual fields.

- **copy\_fields** (bool) Flag determining whether the individual fields given in *fields* are copied. Note that fields are always copied if some of the supplied fields are identical. If fields are copied the original fields will be left untouched. Conversely, if *copy\_fields* == *False*, the original fields are modified so their data points to the collection. It is thus basically impossible to have fields that are linked to multiple collections at the same time.
- label (str) Label of the field collection
- **labels** (*list of str*) Labels of the individual fields. If omitted, the labels from the *fields* argument are used.
- **dtype** (*numpy dtype*) The data type of the field. All the numpy dtypes are supported. If omitted, it will be determined from *data* automatically.

#### append (\*fields, label=None)

create new collection with appended field(s)

#### **Parameters**

- **fields** (*FieldCollection* or *DataFieldBase*) A sequence of single fields or collection of fields that will be appended to the fields in the current collection. The data of all fields will be copied.
- label (str) Label of the new field collection. If omitted, the current label is used

#### Returns

A new field collection, which combines the current one with fields given by *fields*.

## **Return type**

FieldCollection

#### assert\_field\_compatible(other, accept\_scalar=False)

checks whether other is compatible with the current field

## Parameters

- **other** (FieldBase) Other field this is compared to
- accept\_scalar (bool, optional) Determines whether it is acceptable that other is an instance of ScalarField.

## property attributes: dict[str, Any]

describes the state of the instance (without the data)

## Туре

dict

## property attributes\_serialized: dict[str, str]

serialized version of the attributes

Type dict

# property averages: list

averages of all fields

## **copy** (\*, *label=None*, *dtype=None*)

return a copy of the data, but not of the grid

## Parameters

• **label** (*str*, *optional*) – Name of the returned field

• **dtype** (*numpy dtype*) – The data type of the field. If omitted, it will be determined from *data* automatically.

• **self**(FieldCollection) -

## **Return type**

FieldCollection

## property fields: list[DataFieldBase]

the fields of this collection

Type list

create a field collection from classes and data

## **Parameters**

- **field\_classes** (*list*) List of the classes that define the individual fields
- data (ndarray, optional) Data values of all fields at support points of the grid
- grid (GridBase) Grid defining the space on which this field is defined.
- with\_ghost\_cells (bool) Indicates whether the ghost cells are included in data
- **label** (*str*) Label of the field collection
- **labels** (*list of str*) Labels of the individual fields. If omitted, the labels from the *fields* argument are used.
- **dtype** (*numpy dtype*) The data type of the field. All the numpy dtypes are supported. If omitted, it will be determined from *data* automatically.

classmethod from\_scalar\_expressions (grid, expressions, \*, user\_funcs=None, consts=None, label=None, labels=None, dtype=None)

create a field collection on a grid from given expressions

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur.

- grid (GridBase) Grid defining the space on which this field is defined
- **expressions** (*list of str*) A list of mathematical expression, one for each field in the collection. The expressions determine the values as a function of the position on the grid. The expressions may contain standard mathematical functions and they may depend on the axes labels of the grid. More information can be found in the *expression documentation*.
- **user\_funcs** (*dict*, *optional*) A dictionary with user defined functions that can be used in the expression
- **consts** (*dict*, *optional*) A dictionary with user defined constants that can be used in the expression. The values of these constants should either be numbers or ndarray.
- **label** (*str*, *optional*) Name of the whole collection
- labels (list of str, optional) Names of the individual fields

• **dtype** (*numpy dtype*) – The data type of the field. All the numpy dtypes are supported. If omitted, it will be determined from *data* automatically.

## **Return type**

FieldCollection

classmethod from\_state(attributes, data=None)

create a field collection from given state.

## Parameters

- **attributes** (*dict*) The attributes that describe the current instance
- data (ndarray, optional) Data values at support points of the grid defining all fields

# Return type

FieldCollection

# get\_image\_data(index=0, \*\*kwargs)

return data for plotting an image of the field

#### Parameters

- index (int) Index of the field whose data is returned
- \*\*kwargs Arguments forwarded to the get\_image\_data method

#### Returns

Information useful for plotting an image of the field

Return type

dict

## get\_line\_data (index=0, scalar='auto', extract='auto')

return data for a line plot of the field

## Parameters

- **index** (*int*) Index of the field whose data is returned
- **scalar** (*str* or *int*) The method for extracting scalars as described in DataFieldBase.to\_scalar().
- **extract** (*str*) The method used for extracting the line data. See the docstring of the grid method *get\_line\_data* to find supported values.

## Returns

Information useful for performing a line plot of the field

## **Return type**

dict

## property integrals: list

integrals of all fields

## interpolate\_to\_grid (grid, \*, fill=None, label=None)

interpolate the data of this field collection to another grid.

## Parameters

• grid (GridBase) – The grid of the new field onto which the current field is interpolated.

- **fill** (*Number*, *optional*) Determines how values out of bounds are handled. If None, a *ValueError* is raised when out-of-bounds points are requested. Otherwise, the given value is returned.
- label (str, optional) Name of the returned field collection

Interpolated data

#### **Return type**

FieldCollection

## property labels: \_FieldLabels

the labels of all fields

**Note:** The attribute returns a special class \_FieldLabels to allow specific manipulations of the field labels. The returned object behaves much like a list, but assigning values will modify the labels of the fields in the collection.

#### Туре

\_FieldLabels

#### property magnitudes: ndarray

scalar magnitudes of all fields

#### Туре

ndarray

visualize all the fields in the collection

- **kind** (*str* or *list* of *str*) Determines the kind of the visualizations. Supported values are *image*, *line*, *vector*, *interactive*, or *merged*. Alternatively, *auto* determines the best visualization based on each field itself. Instead of a single value for all fields, a list with individual values can be given, unless *merged* is chosen.
- **figsize** (*str or tuple of numbers*) Determines the figure size. The figure size is unchanged if the string *default* is passed. Conversely, the size is adjusted automatically when *auto* is passed. Finally, a specific figure size can be specified using two values, using matplotlib.figure.Figure.set\_size\_inches().
- **arrangement** (*str*) Determines how the subpanels will be arranged. The default value *horizontal* places all subplots next to each other. The alternative value *vertical* puts them below each other.
- **title** (*str*) Title of the plot. If omitted, the title might be chosen automatically. This is shown above all panels.
- **constrained\_layout** (bool) Whether to use *constrained\_layout* in matplotlib.pyplot.figure() call to create a figure. This affects the layout of all plot elements. Generally, spacing might be better with this flag enabled, but it can also lead to problems when plotting multiple plots successively, e.g., when creating a movie.
- filename (*str*, *optional*) If given, the figure is written to the specified file.

- **action** (*str*) Decides what to do with the final figure. If the argument is set to *show*, matplotlib.pyplot.show() will be called to show the plot. If the value is *none*, the figure will be created, but not necessarily shown. The value *close* closes the figure, after saving it to a file when *filename* is given. The default value *auto* implies that the plot is shown if it is not a nested plot call.
- **fig\_style** (*dict*) Dictionary with properties that will be changed on the figure after the plot has been drawn by calling matplotlib.pyplot.setp(). For instance, using fig\_style={'dpi': 200} increases the resolution of the figure.
- **fig** (matplotlib.figures.Figure) Figure that is used for plotting. If omitted, a new figure is created.
- **subplot\_args** (*list*) Additional arguments for the specific subplots. Should be a list with a dictionary of arguments for each subplot. Supplying an empty dict allows to keep the default setting of specific subplots.
- \*\*kwargs All additional keyword arguments are forwarded to the actual plotting function of all subplots.

Instances that contain information to update all the plots with new data later.

#### **Return type**

List of PlotReference

classmethod scalar\_random\_uniform (num\_fields, grid, vmin=0, vmax=1, \*, label=None, labels=None, rng=None)

create scalar fields with random values between vmin and vmax

## Parameters

- num\_fields (int) The number of fields to create
- grid (GridBase) Grid defining the space on which the fields are defined
- **vmin** (*float*) Lower bound. Can be complex to create complex fields
- **vmax** (*float*) Upper bound. Can be complex to create complex fields
- label (str, optional) Name of the field collection
- labels (list of str, optional) Names of the individual fields
- **rng** (Generator) Random number generator (default: default\_rng())

#### **Return type**

FieldCollection

## smooth (sigma=1, \*, out=None, label=None)

applies Gaussian smoothing with the given standard deviation

This function respects periodic boundary conditions of the underlying grid, using reflection when no periodicity is specified.

#### sigma (float):

Gives the standard deviation of the smoothing in real length units (default: 1)

# out (FieldCollection, optional):

Optional field into which the smoothed data is stored

## label (str, optional):

Name of the returned field

Field collection with smoothed data, stored at out if given.

## Parameters

- sigma(float)-
- out (FieldCollection / None) -
- label (str / None) -

Return type

FieldCollection

#### classmethod unserialize\_attributes (attributes)

unserializes the given attributes

Parameters attributes (dict) – The serialized attributes Returns

The unserialized attributes

Return type

# 4.1.3 pde.fields.scalar module

Defines a scalar field over a grid

class ScalarField (grid, data='zeros', \*, label=None, dtype=None, with\_ghost\_cells=False)

Bases: DataFieldBase

Scalar field discretized on a grid

## Parameters

- grid (GridBase) Grid defining the space on which this field is defined.
- data (Number or ndarray, optional) Field values at the support points of the grid. The flag with\_ghost\_cells determines whether this data array contains values for the ghost cells, too. The resulting field will contain real data unless the *data* argument contains complex values. Special values are "zeros" or None, initializing the field with zeros, and "empty", just allocating memory with unspecified values.
- label (str, optional) Name of the field
- **dtype** (*numpy dtype*) The data type of the field. If omitted, it will be determined from *data* automatically.
- with\_ghost\_cells (bool) Indicates whether the ghost cells are included in data

create a scalar field on a grid from a given expression

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur.

## **Parameters**

- grid (GridBase) Grid defining the space on which this field is defined
- **expression** (*str*) Mathematical expression for the scalar value as a function of the position on the grid. The expression may contain standard mathematical functions and it may depend on the axes labels of the grid. More information can be found in the *expression documentation*.
- **user\_funcs** (*dict*, *optional*) A dictionary with user defined functions that can be used in the expression
- **consts** (*dict*, *optional*) A dictionary with user defined constants that can be used in the expression. The values of these constants should either be numbers or ndarray.
- **label** (*str*, *optional*) Name of the field
- **dtype** (*numpy dtype*) The data type of the field. If omitted, it will be determined from *data* automatically.

## **Return type**

ScalarField

#### classmethod from\_image (path, bounds=None, periodic=False, \*, label=None)

create a scalar field from an image

## Parameters

- path (Path or str) The path to the image file
- **bounds** (*tuple*, *optional*) Gives the coordinate range for each axis. This should be two tuples of two numbers each, which mark the lower and upper bound for each axis.
- **periodic** (*bool or list*) Specifies which axes possess periodic boundary conditions. This is either a list of booleans defining periodicity for each individual axis or a single boolean value specifying the same periodicity for all axes.
- label (str, optional) Name of the field

## **Return type**

## ScalarField

## get\_boundary\_field(index, bc=None, \*, label=None)

get the field on the specified boundary

- **index** (*str or tuple*) Index specifying the boundary. Can be either a string given in *boundary\_names*, like "left", or a tuple of the axis index perpendicular to the boundary and a boolean specifying whether the boundary is at the upper side of the axis or not, e.g., (1, True).
- **bc** (*BoundariesData | None*) The boundary conditions applied to the field. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by *['value': NUM]*) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by *['derivative': DERIV]*) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis

and a vanishing derivative otherwise. More information can be found in the *boundaries doc-umentation*. If the special value *None* is given, no boundary conditions are enforced. The user then needs to ensure that the ghost cells are set accordingly.

• label (str) - Label of the returned field

### Returns

The field on the boundary

#### **Return type**

ScalarField

## gradient (bc, out=None, \*\*kwargs)

apply gradient operator and return result as a field

#### Parameters

- bc (BoundariesData / None) The boundary conditions applied to the field. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by *['value': NUM]*) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by *['derivative': DERIV]*) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documentation*. If the special value *None* is given, no boundary conditions are enforced. The user then needs to ensure that the ghost cells are set accordingly.
- out (VectorField, optional) Optional vector field to which the result is written.
- **label** (*str*, *optional*) Name of the returned field

## Returns

result of applying the operator

## **Return type**

VectorField

#### gradient\_squared(bc, out=None, \*\*kwargs)

apply squared gradient operator and return result as a field

This evaluates  $|\nabla \phi|^2$  for the scalar field  $\phi$ 

- bc (BoundariesData / None) The boundary conditions applied to the field. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by *['value': NUM]*) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by *['derivative': DERIV]*) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documentation*. If the special value *None* is given, no boundary conditions are enforced. The user then needs to ensure that the ghost cells are set accordingly.
- out (ScalarField, optional) Optional vector field to which the result is written.
- **label** (*str*, *optional*) Name of the returned field

• **central** (*bool*) – Determines whether a central difference approximation is used for the gradient operator or not. If not, the squared gradient is calculated as the mean of the squared values of the forward and backward derivatives, which thus includes the value at a support point in the result at the same point.

## Returns

the squared gradient of the field

#### Return type

ScalarField

#### property integral: int | float | complex

integral of the scalar field over space

#### Туре

Number

interpolate\_to\_grid (grid, \*, bc=None, fill=None, label=None)

interpolate the data of this scalar field to another grid.

#### **Parameters**

- grid (GridBase) The grid of the new field onto which the current field is interpolated.
- bc (BoundariesData / None) The boundary conditions applied to the field, which affects values close to the boundary. If omitted, the argument *fill* is used to determine values outside the domain. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by *{'value': NUM}*) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by *{'derivative': DERIV}*) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documentation*. If the special value *None* is given, no boundary conditions are enforced. The user then needs to ensure that the ghost cells are set accordingly.
- **fill** (*Number*, *optional*) Determines how values out of bounds are handled. If *None*, a *ValueError* is raised when out-of-bounds points are requested. Otherwise, the given value is returned.
- **label** (*str*, *optional*) Name of the returned field

#### • **self**(ScalarField)-

#### Returns

Field of the same rank as the current one.

#### **Return type**

ScalarField

laplace (bc, out=None, \*\*kwargs)

apply Laplace operator and return result as a field

#### Parameters

bc (BoundariesData / None) – The boundary conditions applied to the field. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value

NUM (specified by *{'value': NUM}*) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by *{'derivative': DERIV}*) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries doc-umentation*. If the special value *None* is given, no boundary conditions are enforced. The user then needs to ensure that the ghost cells are set accordingly.

- **out** (ScalarField, optional) Optional scalar field to which the result is written.
- **label** (*str*, *optional*) Name of the returned field
- **backend** (*str*) The backend (e.g., 'numba' or 'scipy') used for this operator.

#### Returns

the Laplacian of the field

#### Return type

ScalarField

project (axes, method='integral', label=None)

project scalar field along given axes

## Parameters

- **axes** (*list of str*) The names of the axes that are removed by the projection operation. The valid names for a given grid are the ones in the GridBase.axes attribute.
- **method** (*str*) The projection method. This can be either 'integral' to integrate over the removed axes or 'average' to perform an average instead.
- label (str, optional) The label of the returned field

#### Returns

The projected data in a scalar field with a subgrid of the original grid.

## **Return type** *ScalarField*

#### rank: int = 0

slice (position, \*, method='nearest', label=None)

slice data at a given position

**Note:** This method should not be used to evaluate fields right at the boundary since it does not respect boundary conditions. Use *get\_boundary\_field()* to obtain the values directly on the boundary.

#### **Parameters**

- position (dict) Determines the location of the slice using a dictionary supplying coordinate values for a subset of axes. Axes not mentioned in the dictionary are retained and form the slice. For instance, in a 2d Cartesian grid, *position* = {'x': 1} slices along the ydirection at x=1. Additionally, the special positions 'low', 'mid', and 'high' are supported to reference relative positions along the axis.
- **method** (*str*) The method used for slicing. Currently, we only support *nearest*, which takes data from cells defined on the grid.
- **label** (*str*, *optional*) The label of the returned field

## Returns

The sliced data in a scalar field with a subgrid of the original grid.

## **Return type**

ScalarField

to\_scalar(scalar='auto', \*, label=None)

return a modified scalar field by applying method scalar

## Parameters

- **scalar** (*str* or *callable*) Determines the method used for obtaining the scalar. If this is a callable, it is simply applied to self.data and a new scalar field with this data is returned. Alternatively, pre-defined methods can be selected using strings. Here, *abs* and *norm* denote the norm of each entry of the field, while *norm\_squared* returns the squared norm. The default *auto* is to return a (unchanged) copy of a real field and the norm of a complex field.
- label (str, optional) Name of the returned field

## Returns

Scalar field after applying the operation

## **Return type**

ScalarField

# 4.1.4 pde.fields.tensorial module

Defines a tensorial field of rank 2 over a grid

```
class Tensor2Field (grid, data='zeros', *, label=None, dtype=None, with_ghost_cells=False)
```

Bases: DataFieldBase

Tensor field of rank 2 discretized on a grid

**Warning:** Components of the tensor field are given in the local basis. While the local basis is identical to the global basis in Cartesian coordinates, the local basis depends on position in curvilinear coordinate systems. Moreover, the field always contains all components, even if the underlying grid assumes symmetries.

- grid (GridBase) Grid defining the space on which this field is defined.
- data (Number or ndarray, optional) Field values at the support points of the grid. The flag with\_ghost\_cells determines whether this data array contains values for the ghost cells, too. The resulting field will contain real data unless the *data* argument contains complex values. Special values are "zeros" or None, initializing the field with zeros, and "empty", just allocating memory with unspecified values.
- label (str, optional) Name of the field
- **dtype** (*numpy dtype*) The data type of the field. If omitted, it will be determined from *data* automatically.
- with\_ghost\_cells (bool) Indicates whether the ghost cells are included in data

#### divergence (bc, out=None, \*\*kwargs)

apply tensor divergence and return result as a field

The tensor divergence is a vector field  $v_{\alpha}$  resulting from a contracting of the derivative of the tensor field  $t_{\alpha\beta}$ :

$$v_{\alpha} = \sum_{\beta} \frac{\partial t_{\alpha\beta}}{\partial x_{\beta}}$$

## Parameters

- bc (BoundariesData / None) The boundary conditions applied to the field. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by *{'value': NUM}*) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by *{'derivative': DERIV}*) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documentation*. If the special value *None* is given, no boundary conditions are enforced. The user then needs to ensure that the ghost cells are set accordingly.
- **out** (VectorField, *optional*) Optional scalar field to which the result is written.
- label (str, optional) Name of the returned field
- **\*\*kwargs** Additional arguments affecting how the operator behaves.

#### Returns

result of applying the operator

## **Return type**

VectorField

dot (other, out=None, \*, conjugate=True, label='dot product')

calculate the dot product involving a tensor field

This supports the dot product between two tensor fields as well as the product between a tensor and a vector. The resulting fields will be a tensor or vector, respectively.

#### Parameters

- other (VectorField or Tensor2Field) the second field
- **out** (VectorField or Tensor2Field, optional) Optional field to which the result is written.
- conjugate (bool) Whether to use the complex conjugate for the second operand
- label (str, optional) Name of the returned field

#### Returns

VectorField or Tensor2Field: result of applying the dot operator

#### **Return type**

VectorField | Tensor2Field

classmethod from\_expression (grid, expressions, \*, user\_funcs=None, consts=None, label=None, dtype=None)

create a tensor field on a grid from given expressions

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur.

## Parameters

- grid (GridBase) Grid defining the space on which this field is defined
- **expressions** (*list* of *str*) A 2d list of mathematical expression, one for each component of the tensor field. The expressions determine the values as a function of the position on the grid. The expressions may contain standard mathematical functions and they may depend on the axes labels of the grid. More information can be found in the *expression documentation*.
- **user\_funcs** (*dict*, *optional*) A dictionary with user defined functions that can be used in the expression
- **consts** (*dict*, *optional*) A dictionary with user defined constants that can be used in the expression. The values of these constants should either be numbers or ndarray.
- **label** (*str*, *optional*) Name of the field
- **dtype** (*numpy dtype*) The data type of the field. If omitted, it will be determined from *data* automatically.

#### **Return type**

Tensor2Field

## property integral: ndarray

integral of each component over space

## Туре

ndarray

visualize all the components of this tensor field

- **kind** (*str or list of str*) Determines the kind of the visualizations. Supported values are *image* or *line*. Alternatively, *auto* determines the best visualization based on the grid.
- **title** (*str*) Title of the plot. If omitted, the title might be chosen automatically. This is shown above all panels.
- **constrained\_layout** (bool) Whether to use *constrained\_layout* in matplotlib.pyplot.figure() call to create a figure. This affects the layout of all plot elements. Generally, spacing might be better with this flag enabled, but it can also lead to problems when plotting multiple plots successively, e.g., when creating a movie.
- filename (*str*, *optional*) If given, the figure is written to the specified file.
- **action** (*str*) Decides what to do with the final figure. If the argument is set to *show*, matplotlib.pyplot.show() will be called to show the plot. If the value is *none*, the figure will be created, but not necessarily shown. The value *close* closes the figure, after saving it to a file when *filename* is given. The default value *auto* implies that the plot is shown if it is not a nested plot call.

- **fig\_style** (*dict*) Dictionary with properties that will be changed on the figure after the plot has been drawn by calling matplotlib.pyplot.setp(). For instance, using fig\_style={'dpi': 200} increases the resolution of the figure.
- **fig** (matplotlib.figures.Figure) Figure that is used for plotting. If omitted, a new figure is created.
- \*\*kwargs All additional keyword arguments are forwarded to the actual plotting function of all subplots.

Instances that contain information to update all the plots with new data later.

## **Return type**

2d list of PlotReference

#### rank: int = 2

#### symmetrize (make\_traceless=False, inplace=False)

symmetrize the tensor field in place

#### **Parameters**

- make\_traceless (bool) Determines whether the result is also traceless
- **inplace** (*bool*) Flag determining whether to symmetrize the current field or return a new one

#### Returns

result of the operation

## **Return type**

Tensor2Field

#### to\_scalar(scalar='auto', \*, label='scalar`{scalar}`')

return scalar variant of the field

The invariants of the tensor field A are

$$I_{1} = \operatorname{tr}(\boldsymbol{A})$$

$$I_{2} = \frac{1}{2} \left[ (\operatorname{tr}(\boldsymbol{A})^{2} - \operatorname{tr}(\boldsymbol{A}^{2})) \right]$$

$$I_{3} = \operatorname{det}(\boldsymbol{A})$$

where tr denotes the trace and det denotes the determinant. Note that the three invariants can only be distinct and non-zero in three dimensions. In two dimensional spaces, we have the identity  $2I_2 = I_3$  and in onedimensional spaces, we have  $I_1 = I_3$  as well as  $I_2 = 0$ .

#### Parameters

- **scalar** (*str*) The method to calculate the scalar. Possible choices include *norm* (the default chosen when the value is *auto*), *min*, *max*, *squared\_sum*, *norm\_squared*, *trace* (or *invariant1*), *invariant2*, and *determinant* (or *invariant3*)
- **label** (*str*, *optional*) Name of the returned field

#### Returns

the scalar field after applying the operation

#### **Return type**

ScalarField

## trace (label='trace')

return the trace of the tensor field as a scalar field

#### Parameters

label(str, optional) - Name of the returned field

#### Returns

scalar field of traces

#### Return type

ScalarField

#### transpose (label='transpose')

return the transpose of the tensor field

#### Parameters

label (str, optional) - Name of the returned field

#### Returns

transpose of the tensor field

## Return type

Tensor2Field

# 4.1.5 pde.fields.vectorial module

Defines a vectorial field over a grid

```
class VectorField (grid, data='zeros', *, label=None, dtype=None, with_ghost_cells=False)
```

Bases: DataFieldBase

Vector field discretized on a grid

**Warning:** Components of the vector field are given in the local basis. While the local basis is identical to the global basis in Cartesian coordinates, the local basis depends on position in curvilinear coordinate systems. Moreover, the field always contains all components, even if the underlying grid assumes symmetries.

## Parameters

- grid (GridBase) Grid defining the space on which this field is defined.
- data (Number or ndarray, optional) Field values at the support points of the grid. The flag with\_ghost\_cells determines whether this data array contains values for the ghost cells, too. The resulting field will contain real data unless the *data* argument contains complex values. Special values are "zeros" or None, initializing the field with zeros, and "empty", just allocating memory with unspecified values.
- label (str, optional) Name of the field
- **dtype** (*numpy dtype*) The data type of the field. If omitted, it will be determined from *data* automatically.
- with\_ghost\_cells (bool) Indicates whether the ghost cells are included in data

## divergence(bc, out=None, \*\*kwargs)

apply divergence operator and return result as a field

- bc (BoundariesData / None) The boundary conditions applied to the field. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by *['value': NUM]*) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by *['derivative': DERIV]*) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documentation*. If the special value *None* is given, no boundary conditions are enforced. The user then needs to ensure that the ghost cells are set accordingly.
- **out** (ScalarField, *optional*) Optional scalar field to which the result is written.
- label (str, optional) Name of the returned field
- **\*\*kwargs** Additional arguments affecting how the operator behaves.

Divergence of the field

#### **Return type**

ScalarField

dot (other, out=None, \*, conjugate=True, label='dot product')

calculate the dot product involving a vector field

This supports the dot product between two vectors fields as well as the product between a vector and a tensor. The resulting fields will be a scalar or vector, respectively.

#### Parameters

- other (VectorField or Tensor2Field) the second field
- **out** (ScalarField *or* VectorField, *optional*) Optional field to which the result is written.
- conjugate (bool) Whether to use the complex conjugate for the second operand
- label (str, optional) Name of the returned field

#### Returns

ScalarField or VectorField: result of applying the operator

## Return type

ScalarField | VectorField

classmethod from\_expression (grid, expressions, \*, user\_funcs=None, consts=None, label=None, dtype=None)

create a vector field on a grid from given expressions

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur.

- grid (GridBase) Grid defining the space on which this field is defined
- **expressions** (*list of str*) A list of mathematical expression, one for each component of the vector field. The expressions determine the values as a function of the position

on the grid. The expressions may contain standard mathematical functions and they may depend on the axes labels of the grid. More information can be found in the *expression documentation*.

- **user\_funcs** (*dict*, *optional*) A dictionary with user defined functions that can be used in the expression
- **consts** (*dict*, *optional*) A dictionary with user defined constants that can be used in the expression. The values of these constants should either be numbers or ndarray.
- label (str, optional) Name of the field
- **dtype** (*numpy dtype*) The data type of the field. If omitted, it will be determined from *data* automatically.

#### Return type

VectorField

#### classmethod from\_scalars (fields, \*, label=None, dtype=None)

create a vector field from a list of ScalarFields

Note that the data of the scalar fields is copied in the process

#### Parameters

- fields (list) The list of (compatible) scalar fields
- label (str, optional) Name of the returned field
- **dtype** (*numpy dtype*) The data type of the field. If omitted, it will be determined from *data* automatically.

#### Returns

the resulting vector field

## **Return type**

VectorField

get\_vector\_data (transpose=False, max\_points=None, \*\*kwargs)

return data for a vector plot of the field

#### Parameters

- **transpose** (bool) Determines whether the transpose of the data should be plotted.
- max\_points (*int*) The maximal number of points that is used along each axis. This option can be used to sub-sample the data.
- \*\*kwargs Additional parameters forwarded to grid.get\_image\_data

#### Returns

Information useful for plotting an vector field

#### **Return type**

dict

#### gradient (bc, out=None, \*\*kwargs)

apply vector gradient operator and return result as a field

The vector gradient field is a tensor field  $t_{\alpha\beta}$  that specifies the derivatives of the vector field  $v_{\alpha}$  with respect to all coordinates  $x_{\beta}$ .

- bc (BoundariesData / None) The boundary conditions applied to the field. Boundary conditions need to determine all components of the vector field. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by {'value': NUM}) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by {'derivative': DERIV}) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the boundaries documentation. If the special value None is given, no boundary conditions are enforced. The user then needs to ensure that the ghost cells are set accordingly.
- **out** (VectorField, optional) Optional vector field to which the result is written.
- **label** (*str*, *optional*) Name of the returned field
- **\*\*kwargs** Additional arguments affecting how the operator behaves.

Gradient of the field

## **Return type**

Tensor2Field

## property integral: ndarray

integral of each component over space

## Туре

ndarray

interpolate\_to\_grid (grid, \*, bc=None, fill=None, label=None)

interpolate the data of this vector field to another grid.

## Parameters

- grid (GridBase) The grid of the new field onto which the current field is interpolated.
- **bc** (BoundariesData / None) The boundary conditions applied to the field, which affects values close to the boundary. If omitted, the argument *fill* is used to determine values outside the domain. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by {value': NUM}) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by {'derivative': DERIV}) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documentation*. If the special value *None* is given, no boundary conditions are enforced. The user then needs to ensure that the ghost cells are set accordingly.
- **fill** (*Number*, *optional*) Determines how values out of bounds are handled. If *None*, a *ValueError* is raised when out-of-bounds points are requested. Otherwise, the given value is returned.
- label (str, optional) Name of the returned field
- **self** (VectorField) -

## Returns

Field of the same rank as the current one.

#### **Return type**

VectorField

laplace (bc, out=None, \*\*kwargs)

apply vector Laplace operator and return result as a field

The vector Laplacian is a vector field  $L_{\alpha}$  containing the second derivatives of the vector field  $v_{\alpha}$  with respect to the coordinates  $x_{\beta}$ :

$$L_{\alpha} = \sum_{\beta} \frac{\partial^2 v_{\alpha}}{\partial x_{\beta} \partial x_{\beta}}$$

## **Parameters**

- bc (BoundariesData / None) The boundary conditions applied to the field. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by {value': NUM}) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by {'derivative': DERIV}) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the boundaries documentation. If the special value None is given, no boundary conditions are enforced. The user then needs to ensure that the ghost cells are set accordingly.
- out (VectorField, optional) Optional vector field to which the result is written.
- **label** (*str*, *optional*) Name of the returned field
- **\*\*kwargs** Additional arguments affecting how the operator behaves.

#### Returns

Laplacian of the field

#### Return type

VectorField

#### make\_outer\_prod\_operator(backend='numba')

return operator calculating the outer product of two vector fields

Warning: This function does not check types or dimensions.

#### Parameters

**backend** (*str*) – The backend (e.g., 'numba' or 'scipy') used for this operator.

## Returns

function that takes two instance of ndarray, which contain the discretized data of the two operands. An optional third argument can specify the output array to which the result is written. Note that the returned function is jitted with numba for speed.

#### **Return type**

Callable[[ndarray, ndarray, ndarray | None], ndarray]

## outer\_product (other, out=None, \*, label=None)

calculate the outer product of this vector field with another

- other (VectorField) The second vector field
- **out** (*Tensor2Field*, optional) Optional tensorial field to which the result is written.
- label (str, optional) Name of the returned field

result of the operation

#### Return type

Tensor2Field

```
rank: int = 1
```

to\_scalar (scalar='auto', \*, label='scalar `{scalar}`')

return scalar variant of the field

## **Parameters**

- **scalar** (*str*) Choose the method to use. Possible choices are *norm*, *max*, *min*, *squared\_sum*, *norm\_squared*, or an integer specifying which component is returned (indexing starts at 0). The default value *auto* picks the method automatically: The first (and only) component is returned for real fields on one-dimensional spaces, while the norm of the vector is returned otherwise.
- label (str, optional) Name of the returned field

#### Returns

The scalar field after applying the operation

## **Return type**

pde.fields.scalar.ScalarField

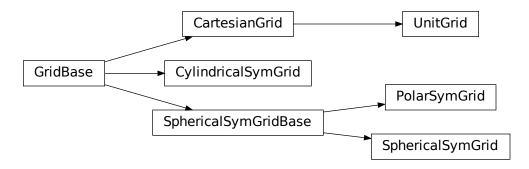
# 4.2 pde.grids package

Grids define the domains on which PDEs will be solved. In particular, symmetries, periodicities, and the discretizations are defined by the underlying grid.

We only consider regular, orthogonal grids, which are constructed from orthogonal coordinate systems with equidistant discretizations along each axis. The dimension of the space that the grid describes is given by the attribute dim. Cartesian coordinates can be mapped to grid coordinates and the corresponding discretization cells using the method transform().

UnitGrid	d-dimensional Cartesian grid with unit discretization in all directions
CartesianGrid	d-dimensional Cartesian grid with uniform discretization for each axis
PolarSymGrid	2-dimensional polar grid assuming angular symmetry
SphericalSymGrid	3-dimensional spherical grid assuming spherical symme- try
CylindricalSymGrid	3-dimensional cylindrical grid assuming polar symmetry

Inheritance structure of the classes:



Subpackages:

# 4.2.1 pde.grids.boundaries package

This package contains classes for handling the boundary conditions of fields.

# **Boundary conditions**

The mathematical details of boundary conditions for partial differential equations are treated in more detail in the documentation document. Since the *pde* package only supports orthogonal grids, boundary conditions need to be applied at the end of each axis. Consequently, methods expecting boundary conditions typically receive a list of conditions for each axes:

```
field = ScalarField(UnitGrid([16, 16], periodic=[True, False]))
field.laplace(bc=[bc_x, bc_y])
```

If an axis is periodic (like the first one in the example above), the only valid boundary conditions are 'periodic' and its cousin 'anti-periodic', which imposes opposite signs on both sides. For non-periodic axes (e.g., the second axis), different boundary conditions can be specified for the lower and upper end of the axis, which is done using a tuple of two conditions. Typical choices for individual conditions are Dirichlet conditions that enforce a value NUM (specified by *{ 'value': NUM}*) and Neumann conditions that enforce the value DERIV for the derivative in the normal direction (specified by *{ 'derivative': DERIV}*). The specific choices for the example above could be

```
bc_x = "periodic"
bc_y = ({"value": 2}, {"derivative": -1})
```

which enforces a value of 2 at the lower side of the y-axis and a derivative (in outward normal direction) of -1 on the upper side. Instead of plain numbers, which enforce the same condition along the whole boundary, expressions can be used to support inhomogeneous boundary conditions. These mathematical expressions are given as a string that can be parsed by *sympy*. They can depend on all coordinates of the grid. An alternative boundary condition to the example above could thus read

```
bc_y = ({"value": "y**2"}, {"derivative": "-sin(x)"})
```

**Warning:** To interpret arbitrary expressions, the package uses exec(). It should therefore not be used in a context where malicious input could occur.

Inhomogeneous values can also be specified by directly supplying an array, whose shape needs to be compatible with the boundary, i.e., it needs to have the same shape as the grid but with the dimension of the axis along which the boundary is specified removed.

The package also supports mixed boundary conditions (depending on both the value and the derivative of the field) and imposing a second derivative. An example is

which enforces the condition  $\partial_n c + 2c = 7$  and  $\partial_n^2 c = 2$  onto the field c on the lower and upper side of the axis, respectively.

Beside the full specification of the boundary conditions, various short-hand notations are supported. If both sides of an axis have the same boundary condition, only one needs to be specified (instead of the tuple). For instance,  $bc_y = \{ value': 2 \}$  imposes a value of 2 on both sides of the y-axis. Similarly, if all axes have the same boundary conditions, only one axis needs to be specified (instead of the list). For instance, the following example

```
field = ScalarField(UnitGrid([16, 16], periodic=False))
field.laplace(bc={"value": 2})
```

imposes a value of 2 on all sides of the grid. Finally, the special values 'auto\_periodic\_neumann' and 'auto\_periodic\_dirichlet' impose periodic boundary conditions for periodic axis and a vanishing derivative or value otherwise. For example,

```
field = ScalarField(UnitGrid([16, 16], periodic=[True, False]))
field.laplace(bc="auto_periodic_neumann")
```

enforces periodic boundary conditions on the first axis, while the second one has standard Neumann conditions.

**Note:** Derivatives are given relative to the outward normal vector, such that positive derivatives correspond to a function that increases across the boundary.

## **Boundaries overview**

The *boundaries* package defines the following classes:

#### Local boundary conditions:

- DirichletBC: Imposing a constant value of the field at the boundary
- *ExpressionValueBC*: Imposing the value of the field at the boundary given by an expression or a python function
- NeumannBC: Imposing a constant derivative of the field in the outward normal direction at the boundary
- *ExpressionDerivativeBC*: Imposing the derivative of the field in the outward normal direction at the boundary given by an expression or a python function
- *MixedBC*: Imposing the derivative of the field in the outward normal direction proportional to its value at the boundary

- *ExpressionMixedBC*: Imposing the derivative of the field in the outward normal direction proportional to its value at the boundary with coefficients given by expressions or python functions
- CurvatureBC: Imposing a constant second derivative (curvature) of the field at the boundary

There are corresponding classes that only affect the normal component of a field, which can be useful when dealing with vector and tensor fields: *NormalDirichletBC*, *NormalNeumannBC*, *NormalMixedBC*, and *NormalCurva-tureBC*.

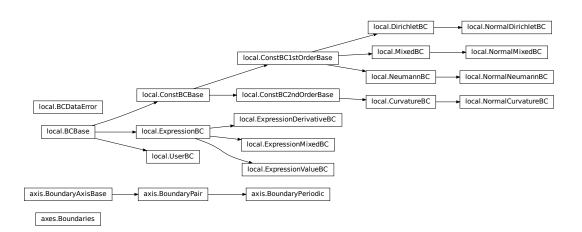
## Boundaries for an axis:

- BoundaryPair: Uses the local boundary conditions to specify the two boundaries along an axis
- BoundaryPeriodic: Indicates that an axis has periodic boundary conditions

## Boundaries for all axes of a grid:

• Boundaries: Collection of boundaries to describe conditions for all axes

## Inheritance structure of the classes:



The details of the classes are explained below:

## pde.grids.boundaries.axes module

This module handles the boundaries of all axes of a grid. It only defines *Boundaries*, which acts as a list of *BoundaryAxisBase*.

## class Boundaries(boundaries)

Bases: list

class that bundles all boundary conditions for all axes

initialize with a list of boundaries

## property boundaries: Iterator[BCBase]

iterator over all non-periodic boundaries

#### check\_value\_rank(rank)

check whether the values at the boundaries have the correct rank

#### **Parameters**

**rank** (*int*) – The tensorial rank of the field for this boundary condition

Return type None

## **Throws:**

RuntimeError: if any value does not have rank rank

## copy()

create a copy of the current boundaries

Return type

Boundaries

#### classmethod from\_data (grid, boundaries, rank=0)

Creates all boundaries from given data

## Parameters

- grid (GridBase) The grid with which the boundary condition is associated
- **boundaries** (*str* or *list* or *tuple* or *dict*) Data that describes the boundaries. This can either be a list of specifications for each dimension or a single one, which is then applied to all dimensions. The boundary for a dimensions can be specified by one of the following formats:
  - string specifying a single type for all boundaries
  - dictionary specifying the type and values for all boundaries
  - tuple pair specifying the low and high boundary individually
- rank (int) The tensorial rank of the field for this boundary condition

## **Return type**

Boundaries

## classmethod get\_help()

Return information on how boundary conditions can be set

## **Return type**

str

```
get_mathematical_representation (field_name='C')
```

return mathematical representation of the boundary condition

```
Parameters
field_name(str)-
```

**Return type** 

str

## grid: GridBase

grid for which boundaries are defined

#### Туре

GridBase

#### make\_ghost\_cell\_setter()

return function that sets the ghost cells on a full array

## **Return type**

GhostCellSetter

## property periodic: list[bool]

a boolean array indicating which dimensions are periodic according to the boundary conditions

Type

ndarray

set\_ghost\_cells (data\_full, \*, set\_corners=False, args=None)

set the ghost cells for all boundaries

#### Parameters

- data\_full (ndarray) The full field data including ghost points
- set\_corners (bool) Determines whether the corner cells are set using interpolation
- args Additional arguments that might be supported by special boundary conditions.

#### **Return type**

None

#### pde.grids.boundaries.axis module

This module handles the boundaries of a single axis of a grid. There are generally only two options, depending on whether the axis of the underlying grid is defined as periodic or not. If it is periodic, the class *BoundaryPeriodic* should be used, while non-periodic axes have more option, which are represented by *BoundaryPair*.

## class BoundaryAxisBase(low, high)

Bases: object

base class for defining boundaries of a single axis in a grid

#### Parameters

- low (BCBase) Instance describing the lower boundary
- high (BCBase) Instance describing the upper boundary

#### property axis: int

The axis along which the boundaries are defined

#### Туре

int

## copy()

return a copy of itself, but with a reference to the same grid

# Return type

BoundaryAxisBase

## classmethod get\_help()

Return information on how boundary conditions can be set

#### **Return type**

str

## get\_mathematical\_representation ( $field_name='C'$ )

return mathematical representation of the boundary condition

```
Parameters
field_name(str)-
```

**Return type** 

tuple[str, str]

## get\_sparse\_matrix\_data(idx)

sets the elements of the sparse representation of this condition

## **Parameters**

idx (tuple) - The index of the point that must lie on the boundary condition

#### Returns

A constant value and a dictionary with indices and factors that can be used to calculate this virtual point

#### **Return type**

float, dict

property grid: GridBase

Underlying grid

Туре

GridBase

high: BCBase

Boundary condition at upper end

## Туре

BCBase

#### low: BCBase

Boundary condition at lower end

#### Туре

BCBase

## make\_ghost\_cell\_setter()

return function that sets the ghost cells for this axis on a full array

Return type

GhostCellSetter

## property periodic: bool

whether the axis is periodic

Туре

bool

## property rank: int

rank of the associated boundary condition

Туре

int

#### set\_ghost\_cells (data\_full, \*, args=None)

set the ghost cell values for all boundaries

- data\_full (ndarray) The full field data including ghost points
- args Additional arguments that might be supported by special boundary conditions.

Return type

None

## class BoundaryPair (low, high)

## Bases: BoundaryAxisBase

represents the two boundaries of an axis along a single dimension

#### Parameters

- **low** (*BCBase*) Instance describing the lower boundary
- high (BCBase) Instance describing the upper boundary

#### check\_value\_rank(rank)

check whether the values at the boundaries have the correct rank

#### Parameters

**rank** (*int*) – The tensorial rank of the field for this boundary condition

Return type None

#### **Throws:**

RuntimeError: if the value does not have rank rank

#### classmethod from\_data (grid, axis, data, rank=0)

create boundary pair from some data

#### **Parameters**

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated
- data (str or dict) Data that describes the boundary pair
- rank (int) The tensorial rank of the field for this boundary condition

#### Returns

the instance created from the data

## **Return type**

BoundaryPair

#### **Throws:**

ValueError if data cannot be interpreted as a boundary pair

#### **class** BoundaryPeriodic (grid, axis, flip\_sign=False)

Bases: BoundaryPair

represent a periodic axis

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated
- **flip\_sign** (bool) Impose different signs on the two sides of the boundary

#### check\_value\_rank(rank)

check whether the values at the boundaries have the correct rank

## Parameters

**rank** (*int*) – The tensorial rank of the field for this boundary condition

Return type None

#### copy()

return a copy of itself, but with a reference to the same grid

# Return type

BoundaryPeriodic

## property flip\_sign

Whether different signs are imposed on the two sides of the boundary

#### Туре

bool

## get\_boundary\_axis (grid, axis, data, rank=0)

return object representing the boundary condition for a single axis

## Parameters

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated
- data (str or tuple or dict) Data describing the boundary conditions for this axis
- **rank** (*int*) The tensorial rank of the field for this boundary condition

#### Returns

Appropriate boundary condition for the axis

#### **Return type**

BoundaryAxisBase

## pde.grids.boundaries.local module

This module contains classes for handling a single boundary of a non-periodic axis. Since an axis has two boundary, we simply distinguish them by a boolean flag *upper*, which is *True* for the side of the axis with the larger coordinate.

The module currently supports the following standard boundary conditions:

- *DirichletBC*: Imposing the value of a field at the boundary
- NeumannBC: Imposing the derivative of a field in the outward normal direction at the boundary
- *MixedBC*: Imposing the derivative of a field in the outward normal direction proportional to its value at the boundary
- *CurvatureBC*: Imposing the second derivative (curvature) of a field at the boundary

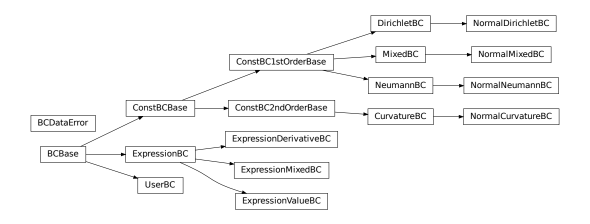
There are also variants of these boundary conditions that only affect the normal components of a vector or tensor field: *NormalDirichletBC*, *NormalNeumannBC*, *NormalMixedBC*, and *NormalCurvatureBC*.

Finally, there are more specialized classes, which offer greater flexibility, but might also require a slightly deeper understanding for proper use:

- *ExpressionValueBC*: Imposing the value of a field at the boundary based on a mathematical expression or a python function.
- *ExpressionDerivativeBC*: Imposing the derivative of a field in the outward normal direction at the boundary based on a mathematical expression or a python function.
- *ExpressionMixedBC*: Imposing a mixed (Robin) boundary condition using mathematical expressions or python functions.
- UserBC: Allows full control for setting virtual points, values, or derivatives. The boundary conditions are never enforced automatically. It is thus the user's responsibility to ensure virtual points are set correctly before operators are applied. To set boundary conditions a dictionary {TARGET: value} must be supplied as argument *args* to set\_ghost\_cells() or the numba equivalent. Here, *TARGET* determines how the *value* is interpreted and what boundary condition is actually enforced: the value of the virtual points directly (*virtual\_point*), the value of the field at the boundary (*value*) or the outward derivative of the field at the boundary (*derivative*).

Note that derivatives are generally given in the direction of the outward normal vector, such that positive derivatives correspond to a function that increases across the boundary.

# Inheritance structure of the classes:



### **class** BCBase (grid, axis, upper, \*, rank=0)

#### Bases: object

represents a single boundary in an BoundaryPair instance

### Parameters

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated
- **upper** (bool) Flag indicating whether this boundary condition is associated with the upper side of an axis or not. In essence, this determines the direction of the local normal vector of the boundary.
- **rank** (*int*) The tensorial rank of the field for this boundary condition

# property axis\_coord: float

value of the coordinate that defines this boundary condition

Type float

# check\_value\_rank(rank)

check whether the values at the boundaries have the correct rank

### Parameters

**rank** (*int*) – The tensorial rank of the field for this boundary condition

Return type None

### **Throws:**

RuntimeError: if the value does not have rank rank

**copy** (*upper=None*, *rank=None*)

# Parameters

- **self** (*TBC*) -
- upper (bool / None) -
- rank (int / None) -

**Return type** 

TBC

### classmethod from\_data (grid, axis, upper, data, \*, rank=0)

create boundary from some data

### **Parameters**

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated
- **upper** (bool) Indicates whether this boundary condition is associated with the upper or lower side of the axis.
- data (str or dict) Data that describes the boundary
- **rank** (*int*) The tensorial rank of the field for this boundary condition

#### Returns

the instance created from the data

### **Return type**

BCBase

### Throws:

ValueError if data cannot be interpreted as a boundary condition

classmethod from\_dict (grid, axis, upper, data, \*, rank=0)

create boundary from data given in dictionary

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated
- **upper** (bool) Indicates whether this boundary condition is associated with the upper or lower side of the axis.

- data (dict) The dictionary defining the boundary condition
- **rank** (*int*) The tensorial rank of the field for this boundary condition

Return type BCBase

**classmethod from str**(grid, axis, upper, condition, \*, rank=0, \*\*kwargs)

creates boundary from a given string identifier

# Parameters

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated
- **upper** (*bool*) Indicates whether this boundary condition is associated with the upper or lower side of the axis.
- condition (*str*) Identifies the boundary condition
- **rank** (*int*) The tensorial rank of the field for this boundary condition
- **\*\*kwargs** Additional arguments passed to the constructor

Return type BCBase

classmethod get\_help()

Return information on how boundary conditions can be set

Return type str

get\_mathematical\_representation (field\_name='C')

return mathematical representation of the boundary condition

```
Parameters
field_name(str)-
```

Return type str

get\_sparse\_matrix\_data(*idx*)

Parameters
idx(tuple[int, ...])-

### **Return type**

tuple[float, dict[int, float]]

get\_virtual\_point (arr, idx=None)

```
Parameters
```

idx(tuple[int, ...] | None)-

Return type

float

# homogeneous: bool

determines whether the boundary condition depends on space

Type

bool

### make\_adjacent\_evaluator()

returns a function evaluating the value adjacent to a given point

Deprecated since version Since: 2023-12-19

#### Returns

A function with signature (arr\_1d, i\_point, bc\_idx), where *arr\_1d* is the one-dimensional data array (the data points along the axis perpendicular to the boundary), *i\_point* is the index into this array for the current point and bc\_idx are the remaining indices of the current point, which indicate the location on the boundary plane. The result of the function is the data value at the adjacent point along the axis associated with this boundary condition in the upper (lower) direction when *upper* is True (False).

# **Return type**

function

### make\_ghost\_cell\_sender()

return function that might mpi\_send data to set ghost cells for this boundary

Return type

GhostCellSetter

# make\_ghost\_cell\_setter()

return function that sets the ghost cells for this boundary

Return type GhostCellSetter

# abstract make\_virtual\_point\_evaluator()

returns a function evaluating the value at the virtual support point

# Returns

A function that takes the data array and an index marking the current point, which is assumed to be a virtual point. The result is the data value at this point, which is calculated using the boundary condition.

Return type function

runction

names: list[str]

identifiers used to specify the given boundary class

# Туре

list

# normal: bool = False

determines whether the boundary condition only affects normal components.

If this flag is *False*, boundary conditions must specify values for all components of the field. If *True*, only the normal components at the boundary are specified.

Type

bool

### property periodic: bool

whether the boundary condition is periodic

Туре

bool

# abstract set\_ghost\_cells(data\_full, \*, args=None)

set the ghost cell values for this boundary

# Parameters

- data\_full (ndarray) The full field data including ghost points
- **args** (ndarray) Determines what boundary conditions are set. *args* should be set to {TARGET: value}. Here, *TARGET* determines how the *value* is interpreted and what boundary condition is actually enforced: the value of the virtual points directly (*virtual\_point*), the value of the field at the boundary (*value*) or the outward derivative of the field at the boundary (*derivative*).

# Return type

None

# to\_subgrid(subgrid)

converts this boundary condition to one valid for a given subgrid

# **Parameters**

• subgrid (GridBase) - Grid of the new boundary conditions

• **self** (*TBC*) -

# Returns

Boundary conditions valid on the subgrid

# **Return type**

BCBase

# exception BCDataError

Bases: ValueError

exception that signals that incompatible data was supplied for the BC

### **class ConstBC1stOrderBase** (grid, axis, upper, \*, rank=0, value=0)

Bases: ConstBCBase

represents a single boundary in an BoundaryPair instance

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur. However, the function is safe when *value* cannot be an arbitrary string.

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated
- **upper** (bool) Flag indicating whether this boundary condition is associated with the upper side of an axis or not. In essence, this determines the direction of the local normal vector of the boundary.
- **rank** (*int*) The tensorial rank of the field for this boundary condition
- normal (bool) Flag indicating whether the condition is only applied in the normal direction.
- **value** (float or str or ndarray) a value stored with the boundary condition. The interpretation of this value depends on the type of boundary condition. If value is a single value

(or tensor in case of tensorial boundary conditions), the same value is applied to all points. Inhomogeneous boundary conditions are possible by supplying an expression as a string, which then may depend on the axes names of the respective grid.

# get\_sparse\_matrix\_data(idx)

sets the elements of the sparse representation of this condition

# Parameters

idx (tuple) – The index of the point that must lie on the boundary condition

### Returns

A constant value and a dictionary with indices and factors that can be used to calculate this virtual point

Return type

float, dict

### get\_virtual\_point (arr, idx=None)

calculate the value of the virtual point outside the boundary

# Parameters

- **arr** (*array*) The data values associated with the grid
- idx (tuple) The index of the point to evaluate. This is a tuple of length grid.num\_axes with the either -1 or dim as the entry for the axis associated with this boundary condition. Here, dim is the dimension of the axis. The index is optional if dim == 1.

# Returns

Value at the virtual support point

# **Return type**

float

### abstract get\_virtual\_point\_data(compiled=False)

return data suitable for calculating virtual points

### Parameters

**compiled** (bool) – Flag indicating whether a compiled version is required, which automatically takes updated values into account when it is used in numba-compiled code.

# Returns

the data structure associated with this virtual point

### **Return type**

tuple

# make\_adjacent\_evaluator()

returns a function evaluating the value adjacent to a given point

Deprecated since version Since: 2023-12-19

# Returns

A function with signature (arr\_1d, i\_point, bc\_idx), where *arr\_1d* is the one-dimensional data array (the data points along the axis perpendicular to the boundary), *i\_point* is the index into this array for the current point and bc\_idx are the remaining indices of the current point, which indicate the location on the boundary plane. The result of the function is the data value at the adjacent point along the axis associated with this boundary condition in the upper (lower) direction when *upper* is True (False).

### Return type

function

# make\_virtual\_point\_evaluator()

returns a function evaluating the value at the virtual support point

# Returns

A function that takes the data array and an index marking the current point, which is assumed to be a virtual point. The result is the data value at this point, which is calculated using the boundary condition.

Return type function

set\_ghost\_cells (data\_full, \*, args=None)

set the ghost cell values for this boundary

# **Parameters**

- data\_full (ndarray) The full field data including ghost points
- **args** (ndarray) Determines what boundary conditions are set. *args* should be set to {TARGET: value}. Here, *TARGET* determines how the *value* is interpreted and what boundary condition is actually enforced: the value of the virtual points directly (*virtual\_point*), the value of the field at the boundary (*value*) or the outward derivative of the field at the boundary (*derivative*).

Return type None

**class** ConstBC2ndOrderBase (grid, axis, upper, \*, rank=0, value=0)

Bases: ConstBCBase

abstract base class for boundary conditions of 2nd order

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur. However, the function is safe when *value* cannot be an arbitrary string.

### Parameters

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated
- **upper** (bool) Flag indicating whether this boundary condition is associated with the upper side of an axis or not. In essence, this determines the direction of the local normal vector of the boundary.
- **rank** (*int*) The tensorial rank of the field for this boundary condition
- **normal** (*bool*) Flag indicating whether the condition is only applied in the normal direction.
- **value** (float or str or ndarray) a value stored with the boundary condition. The interpretation of this value depends on the type of boundary condition. If value is a single value (or tensor in case of tensorial boundary conditions), the same value is applied to all points. Inhomogeneous boundary conditions are possible by supplying an expression as a string, which then may depend on the axes names of the respective grid.

# get\_sparse\_matrix\_data(idx)

sets the elements of the sparse representation of this condition

# Parameters

idx (tuple) - The index of the point that must lie on the boundary condition

# Returns

A constant value and a dictionary with indices and factors that can be used to calculate this virtual point

**Return type** 

float, dict

# get\_virtual\_point (arr, idx=None)

calculate the value of the virtual point outside the boundary

# Parameters

- arr (array) The data values associated with the grid
- idx (tuple) The index of the point to evaluate. This is a tuple of length grid.num\_axes with the either -1 or dim as the entry for the axis associated with this boundary condition. Here, dim is the dimension of the axis. The index is optional if dim == 1.

# Returns

Value at the virtual support point

# **Return type**

float

# abstract get\_virtual\_point\_data()

return data suitable for calculating virtual points

### Returns

the data structure associated with this virtual point

**Return type** 

tuple

### make\_adjacent\_evaluator()

returns a function evaluating the value adjacent to a given point

Deprecated since version Since: 2023-12-19

### Returns

A function with signature (arr\_1d, i\_point, bc\_idx), where *arr\_1d* is the one-dimensional data array (the data points along the axis perpendicular to the boundary), *i\_point* is the index into this array for the current point and bc\_idx are the remaining indices of the current point, which indicate the location on the boundary plane. The result of the function is the data value at the adjacent point along the axis associated with this boundary condition in the upper (lower) direction when *upper* is True (False).

### Return type

function

### make\_virtual\_point\_evaluator()

returns a function evaluating the value at the virtual support point

### Returns

A function that takes the data array and an index marking the current point, which is assumed to be a virtual point. The result is the data value at this point, which is calculated using the boundary condition.

# **Return type**

function

set\_ghost\_cells(data\_full, \*, args=None)

set the ghost cell values for this boundary

# Parameters

- data\_full (ndarray) The full field data including ghost points
- **args** (ndarray) Determines what boundary conditions are set. *args* should be set to {TARGET: value}. Here, *TARGET* determines how the *value* is interpreted and what boundary condition is actually enforced: the value of the virtual points directly (*virtual\_point*), the value of the field at the boundary (*value*) or the outward derivative of the field at the boundary (*derivative*).

Return type None

**class ConstBCBase** (grid, axis, upper, \*, rank=0, value=0)

# Bases: BCBase

base class representing a boundary whose virtual point is set from constants

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur. However, the function is safe when *value* cannot be an arbitrary string.

# **Parameters**

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated
- **upper** (bool) Flag indicating whether this boundary condition is associated with the upper side of an axis or not. In essence, this determines the direction of the local normal vector of the boundary.
- **rank** (*int*) The tensorial rank of the field for this boundary condition
- **normal** (*bool*) Flag indicating whether the condition is only applied in the normal direction.
- **value** (float or str or ndarray) a value stored with the boundary condition. The interpretation of this value depends on the type of boundary condition. If value is a single value (or tensor in case of tensorial boundary conditions), the same value is applied to all points. Inhomogeneous boundary conditions are possible by supplying an expression as a string, which then may depend on the axes names of the respective grid.

**copy** (*upper=None*, *rank=None*, *value=None*)

return a copy of itself, but with a reference to the same grid

# Parameters

- **self** (ConstBCBase) -
- upper (bool / None) -
- rank (int / None) -
- value(float | ndarray | str | None)-

# **Return type**

ConstBCBase

# link\_value(value)

link value of this boundary condition to external array

# Parameters

value (ndarray) -

# to\_subgrid(subgrid)

converts this boundary condition to one valid for a given subgrid

# Parameters

- subgrid (GridBase) Grid of the new boundary conditions
- **self** (ConstBCBase) -

# Returns

Boundary conditions valid on the subgrid

# **Return type**

ConstBCBase

### property value: ndarray

# value\_is\_linked: bool

flag that indicates whether the value associated with this boundary condition is linked to ndarray managed by external code.

# Туре

bool

# **class** CurvatureBC (grid, axis, upper, \*, rank=0, value=0)

Bases: ConstBC2ndOrderBase

represents a boundary condition imposing the 2nd normal derivative at the boundary

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur. However, the function is safe when *value* cannot be an arbitrary string.

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated
- **upper** (bool) Flag indicating whether this boundary condition is associated with the upper side of an axis or not. In essence, this determines the direction of the local normal vector of the boundary.
- **rank** (*int*) The tensorial rank of the field for this boundary condition
- **normal** (*bool*) Flag indicating whether the condition is only applied in the normal direction.
- **value** (float or str or ndarray) a value stored with the boundary condition. The interpretation of this value depends on the type of boundary condition. If value is a single value (or tensor in case of tensorial boundary conditions), the same value is applied to all points. Inhomogeneous boundary conditions are possible by supplying an expression as a string, which then may depend on the axes names of the respective grid.

 $\texttt{get\_mathematical\_representation} (\textit{field\_name='C'})$ 

return mathematical representation of the boundary condition

```
Parameters
field_name(str)-
```

Return type

str

```
get_virtual_point_data()
```

return data suitable for calculating virtual points

# Returns

the data structure associated with this virtual point

# **Return type**

tuple

names: list[str] = ['curvature', 'second\_derivative', 'extrapolate']

identifiers used to specify the given boundary class

Type list

**class DirichletBC** (grid, axis, upper, \*, rank=0, value=0)

Bases: ConstBC1stOrderBase

represents a boundary condition imposing the value

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur. However, the function is safe when *value* cannot be an arbitrary string.

### **Parameters**

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated
- **upper** (bool) Flag indicating whether this boundary condition is associated with the upper side of an axis or not. In essence, this determines the direction of the local normal vector of the boundary.
- rank (int) The tensorial rank of the field for this boundary condition
- **normal** (*bool*) Flag indicating whether the condition is only applied in the normal direction.
- **value** (float or str or ndarray) a value stored with the boundary condition. The interpretation of this value depends on the type of boundary condition. If value is a single value (or tensor in case of tensorial boundary conditions), the same value is applied to all points. Inhomogeneous boundary conditions are possible by supplying an expression as a string, which then may depend on the axes names of the respective grid.

```
get_mathematical_representation (field\_name='C')
```

return mathematical representation of the boundary condition

```
Parameters
field_name(str)-
```

**Return type** 

str

```
get_virtual_point_data(compiled=False)
```

return data suitable for calculating virtual points

### Parameters

**compiled** (*bool*) – Flag indicating whether a compiled version is required, which automatically takes updated values into account when it is used in numba-compiled code.

### Returns

the data structure associated with this virtual point

Return type

tuple

# names: list[str] = ['value', 'dirichlet']

identifiers used to specify the given boundary class

Туре

list

**class ExpressionBC** (grid, axis, upper, \*, rank=0, value=0, const=0, target='virtual\_point', user\_funcs=None, value\_cell=None)

# Bases: BCBase

represents a boundary whose virtual point is calculated from an expression

The expression is given as a string and will be parsed by sympy or a function that is optionally compiled with numba. The expression can contain typical mathematical operators and may depend on the value at the last support point next to the boundary (*value*), spatial coordinates defined by the grid marking the boundary point (e.g., x or r), and time t.

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur.

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated
- **upper** (bool) Flag indicating whether this boundary condition is associated with the upper side of an axis or not. In essence, this determines the direction of the local normal vector of the boundary.
- **rank** (*int*) The tensorial rank of the field for this boundary condition
- value (float or str or callable) An expression that determines the value of the boundary condition. Alternatively, this can be a function with signature (value, dx, \*coords, t) that determines the value of target from the field value value (the value of the adjacent cell unless value\_cell is specified), the spatial discretization dx in the direction perpendicular to the wall, the spatial coordinates of the wall point, and time t. Ideally, this function should be numba-compilable since simulations might otherwise be very slow.
- **const** (*float or str or callable*) An expression similar to *value*, which is only used for mixed (Robin) boundary conditions. Note that the implementation currently does not support that one argument is given as a callable function while the other is defined via an expression, so both need to have the same type.

- target (*str*) Selects which value is actually set. Possible choices include *value*, *derivative*, *mixed*, and *virtual\_point*.
- user\_funcs (dict, optional) A dictionary with user defined functions that can be used in expressions
- **value\_cell**(*int*) Determines which cells is read to determine the field value that is used as *value* in the expression or the function call. The default (*None*) specifies the adjacent cell.

# copy (upper=None, rank=None)

return a copy of itself, but with a reference to the same grid

### Parameters

- **self**(ExpressionBC)-
- upper (bool / None) -
- rank (int / None) -

Return type

ExpressionBC

# get\_mathematical\_representation (field\_name='C')

return mathematical representation of the boundary condition

```
Parameters
field_name(str)-
```

Return type

```
get_sparse_matrix_data(idx)
```

```
Parameters
```

idx(tuple[int, ...])-

# Return type

tuple[float, dict[int, float]]

### get\_virtual\_point (arr, idx=None)

### Parameters

idx(tuple[int, ...] | None)-

# **Return type**

float

# make\_adjacent\_evaluator()

returns a function evaluating the value adjacent to a given point

Deprecated since version Since: 2023-12-19

### Returns

A function with signature (arr\_1d, i\_point, bc\_idx), where *arr\_1d* is the one-dimensional data array (the data points along the axis perpendicular to the boundary), *i\_point* is the index into this array for the current point and bc\_idx are the remaining indices of the current point, which indicate the location on the boundary plane. The result of the function is the data value at the adjacent point along the axis associated with this boundary condition in the upper (lower) direction when *upper* is True (False).

# **Return type**

function

# make\_virtual\_point\_evaluator()

returns a function evaluating the value at the virtual support point

# Returns

A function that takes the data array and an index marking the current point, which is assumed to be a virtual point. The result is the data value at this point, which is calculated using the boundary condition.

Return type function

### names: list[str] = ['virtual\_point']

identifiers used to specify the given boundary class

Туре

list

# set\_ghost\_cells (data\_full, \*, args=None)

set the ghost cell values for this boundary

# Parameters

- data\_full (ndarray) The full field data including ghost points
- **args** (ndarray) Determines what boundary conditions are set. *args* should be set to {TARGET: value}. Here, *TARGET* determines how the *value* is interpreted and what boundary condition is actually enforced: the value of the virtual points directly (*virtual\_point*), the value of the field at the boundary (*value*) or the outward derivative of the field at the boundary (*derivative*).

### **Return type**

None

# to\_subgrid(subgrid)

converts this boundary condition to one valid for a given subgrid

### Parameters

- **subgrid** (GridBase) Grid of the new boundary conditions
- **self**(ExpressionBC) -

#### Returns

Boundary conditions valid on the subgrid

### **Return type**

BCBase

**class ExpressionDerivativeBC** (grid, axis, upper, \*, rank=0, value=0, target='derivative', user funcs=None, value cell=None)

Bases: *ExpressionBC* 

represents a boundary whose outward derivative is calculated from an expression

The expression is given as a string and will be parsed by sympy. The expression can contain typical mathematical operators and may depend on the value at the last support point next to the boundary (*value*), spatial coordinates defined by the grid marking the boundary point (e.g., x or r), and time t.

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur.

# Parameters

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated
- **upper** (bool) Flag indicating whether this boundary condition is associated with the upper side of an axis or not. In essence, this determines the direction of the local normal vector of the boundary.
- rank (int) The tensorial rank of the field for this boundary condition
- **value** (*float or str or callable*) An expression that determines the value of the boundary condition. Alternatively, this can be a function with signature (*value, dx, \*coords, t*) that determines the value of *target* from the field value *value* (the value of the adjacent cell unless *value\_cell* is specified), the spatial discretization *dx* in the direction perpendicular to the wall, the spatial coordinates of the wall point, and time *t*. Ideally, this function should be numba-compilable since simulations might otherwise be very slow.
- **const** (*float or str or callable*) An expression similar to *value*, which is only used for mixed (Robin) boundary conditions. Note that the implementation currently does not support that one argument is given as a callable function while the other is defined via an expression, so both need to have the same type.
- target (*str*) Selects which value is actually set. Possible choices include *value*, *derivative*, *mixed*, and *virtual\_point*.
- user\_funcs (dict, optional) A dictionary with user defined functions that can be used in expressions
- **value\_cell**(*int*) Determines which cells is read to determine the field value that is used as *value* in the expression or the function call. The default (*None*) specifies the adjacent cell.

# names: list[str] = ['derivative\_expression', 'derivative\_expr']

identifiers used to specify the given boundary class

# Туре

list

**class ExpressionMixedBC** (grid, axis, upper, \*, rank=0, value=0, const=0, target='mixed', user\_funcs=None, value\_cell=None)

# Bases: ExpressionBC

represents a boundary whose outward derivative is calculated from an expression

The expression is given as a string and will be parsed by sympy. The expression can contain typical mathematical operators and may depend on the value at the last support point next to the boundary (*value*), spatial coordinates defined by the grid marking the boundary point (e.g., x or r), and time t.

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur.

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated

- **upper** (bool) Flag indicating whether this boundary condition is associated with the upper side of an axis or not. In essence, this determines the direction of the local normal vector of the boundary.
- **rank** (*int*) The tensorial rank of the field for this boundary condition
- value (float or str or callable) An expression that determines the value of the boundary condition. Alternatively, this can be a function with signature (value, dx, \*coords, t) that determines the value of target from the field value value (the value of the adjacent cell unless value\_cell is specified), the spatial discretization dx in the direction perpendicular to the wall, the spatial coordinates of the wall point, and time t. Ideally, this function should be numba-compilable since simulations might otherwise be very slow.
- **const** (*float or str or callable*) An expression similar to *value*, which is only used for mixed (Robin) boundary conditions. Note that the implementation currently does not support that one argument is given as a callable function while the other is defined via an expression, so both need to have the same type.
- target (*str*) Selects which value is actually set. Possible choices include *value*, *derivative*, *mixed*, and *virtual\_point*.
- user\_funcs (dict, optional) A dictionary with user defined functions that can be used in expressions
- **value\_cell**(*int*) Determines which cells is read to determine the field value that is used as *value* in the expression or the function call. The default (*None*) specifies the adjacent cell.

```
names: list[str] = ['mixed_expression', 'mixed_expr', 'robin_expression',
'robin_expr']
```

identifiers used to specify the given boundary class

Type list

**class ExpressionValueBC** (grid, axis, upper, \*, rank=0, value=0, target='value', user\_funcs=None, value\_cell=None)

Bases: ExpressionBC

represents a boundary whose value is calculated from an expression

The expression is given as a string and will be parsed by sympy. The expression can contain typical mathematical operators and may depend on the value at the last support point next to the boundary (*value*), spatial coordinates defined by the grid marking the boundary point (e.g., x or r), and time t.

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur.

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated
- **upper** (bool) Flag indicating whether this boundary condition is associated with the upper side of an axis or not. In essence, this determines the direction of the local normal vector of the boundary.
- **rank** (*int*) The tensorial rank of the field for this boundary condition

- value (float or str or callable) An expression that determines the value of the boundary condition. Alternatively, this can be a function with signature (value, dx, \*coords, t) that determines the value of target from the field value value (the value of the adjacent cell unless value\_cell is specified), the spatial discretization dx in the direction perpendicular to the wall, the spatial coordinates of the wall point, and time t. Ideally, this function should be numba-compilable since simulations might otherwise be very slow.
- **const** (*float* or str or callable) An expression similar to value, which is only used for mixed (Robin) boundary conditions. Note that the implementation currently does not support that one argument is given as a callable function while the other is defined via an expression, so both need to have the same type.
- target (str) Selects which value is actually set. Possible choices include value, derivative, mixed, and virtual\_point.
- user\_funcs (dict, optional) A dictionary with user defined functions that can be used in expressions
- **value\_cell**(*int*) Determines which cells is read to determine the field value that is used as *value* in the expression or the function call. The default (*None*) specifies the adjacent cell.

```
names: list[str] = ['value_expression', 'value_expr']
```

identifiers used to specify the given boundary class

# Туре

list

```
class MixedBC (grid, axis, upper, *, rank=0, value=0, const=0)
```

Bases: ConstBC1stOrderBase

represents a mixed (or Robin) boundary condition imposing a derivative in the outward normal direction of the boundary that is given by an affine function involving the actual value:

 $\partial_n c + \gamma c = \beta$ 

Here, c is the field to which the condition is applied,  $\gamma$  quantifies the influence of the field and  $\beta$  is the constant term. Note that  $\gamma = 0$  corresponds to Dirichlet conditions imposing  $\beta$  as the derivative. Conversely,  $\gamma \to \infty$  corresponds to imposing a zero value on c.

This condition can be enforced by using one of the following variants

```
bc = {'mixed': VALUE}
bc = {'type': 'mixed', 'value': VALUE, 'const': CONST}
```

where VALUE corresponds to  $\gamma$  and CONST to  $\beta$ .

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated
- **upper** (bool) Flag indicating whether this boundary condition is associated with the upper side of an axis or not. In essence, this determines the direction of the local normal vector of the boundary.
- rank (int) The tensorial rank of the field for this boundary condition
- value (float or str or array) The parameter  $\gamma$  quantifying the influence of the field onto its normal derivative. If value is a single value (or tensor in case of tensorial boundary conditions), the same value is applied to all points. Inhomogeneous boundary conditions are

possible by supplying an expression as a string, which then may depend on the axes names of the respective grid.

• **const** (float or ndarray or str) – The parameter  $\beta$  determining the constant term for the boundary condition. Supports the same input as *value*.

**copy** (*upper=None*, *rank=None*, *value=None*, *const=None*)

return a copy of itself, but with a reference to the same grid

### Parameters

- self (MixedBC) -
- upper (bool / None) -
- rank (int / None) -
- value(float | ndarray | str | None)-
- const (float | ndarray | str | None) -

Return type MixedBC

```
get_mathematical_representation (field_name='C')
```

return mathematical representation of the boundary condition

```
Parameters
field_name(str)-
```

Return type

str

# get\_virtual\_point\_data(compiled=False)

return data suitable for calculating virtual points

### Parameters

**compiled** (*bool*) – Flag indicating whether a compiled version is required, which automatically takes updated values into account when it is used in numba-compiled code.

# Returns

the data structure associated with this virtual point

#### **Return type**

tuple

```
names: list[str] = ['mixed', 'robin']
```

identifiers used to specify the given boundary class

### Туре

list

```
to_subgrid(subgrid)
```

converts this boundary condition to one valid for a given subgrid

# Parameters

- subgrid (GridBase) Grid of the new boundary conditions
- self (MixedBC) -

### Returns

Boundary conditions valid on the subgrid

# **Return type**

ConstBCBase

# **class** NeumannBC (grid, axis, upper, \*, rank=0, value=0)

Bases: ConstBC1stOrderBase

represents a boundary condition imposing the derivative in the outward normal direction of the boundary

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur. However, the function is safe when *value* cannot be an arbitrary string.

# **Parameters**

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated
- **upper** (bool) Flag indicating whether this boundary condition is associated with the upper side of an axis or not. In essence, this determines the direction of the local normal vector of the boundary.
- rank (int) The tensorial rank of the field for this boundary condition
- **normal** (*bool*) Flag indicating whether the condition is only applied in the normal direction.
- **value** (float or str or ndarray) a value stored with the boundary condition. The interpretation of this value depends on the type of boundary condition. If value is a single value (or tensor in case of tensorial boundary conditions), the same value is applied to all points. Inhomogeneous boundary conditions are possible by supplying an expression as a string, which then may depend on the axes names of the respective grid.

### get\_mathematical\_representation (field\_name='C')

return mathematical representation of the boundary condition

```
Parameters
field_name(str)-
```

### **Return type**

str

# get\_virtual\_point\_data(compiled=False)

return data suitable for calculating virtual points

# Parameters

**compiled** (bool) – Flag indicating whether a compiled version is required, which automatically takes updated values into account when it is used in numba-compiled code.

### Returns

the data structure associated with this virtual point

**Return type** 

tuple

# names: list[str] = ['derivative', 'neumann']

identifiers used to specify the given boundary class

# Туре

list

# **class** NormalCurvatureBC (grid, axis, upper, \*, rank=0, value=0)

# Bases: CurvatureBC

represents a boundary condition imposing the 2nd normal derivative onto the normal components at the boundary

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur. However, the function is safe when *value* cannot be an arbitrary string.

### **Parameters**

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated
- **upper** (bool) Flag indicating whether this boundary condition is associated with the upper side of an axis or not. In essence, this determines the direction of the local normal vector of the boundary.
- **rank** (*int*) The tensorial rank of the field for this boundary condition
- normal (bool) Flag indicating whether the condition is only applied in the normal direction.
- **value** (float or str or ndarray) a value stored with the boundary condition. The interpretation of this value depends on the type of boundary condition. If value is a single value (or tensor in case of tensorial boundary conditions), the same value is applied to all points. Inhomogeneous boundary conditions are possible by supplying an expression as a string, which then may depend on the axes names of the respective grid.

# names: list[str] = ['normal\_curvature']

identifiers used to specify the given boundary class

# Туре

list

```
normal: bool = True
```

determines whether the boundary condition only affects normal components.

If this flag is *False*, boundary conditions must specify values for all components of the field. If *True*, only the normal components at the boundary are specified.

# Туре

bool

class NormalDirichletBC (grid, axis, upper, \*, rank=0, value=0)

# Bases: DirichletBC

represents a boundary condition imposing the value on normal components

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur. However, the function is safe when *value* cannot be an arbitrary string.

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated

- **upper** (bool) Flag indicating whether this boundary condition is associated with the upper side of an axis or not. In essence, this determines the direction of the local normal vector of the boundary.
- **rank** (*int*) The tensorial rank of the field for this boundary condition
- normal (bool) Flag indicating whether the condition is only applied in the normal direction.
- **value** (float or str or ndarray) a value stored with the boundary condition. The interpretation of this value depends on the type of boundary condition. If value is a single value (or tensor in case of tensorial boundary conditions), the same value is applied to all points. Inhomogeneous boundary conditions are possible by supplying an expression as a string, which then may depend on the axes names of the respective grid.

```
names: list[str] = ['normal_value', 'normal_dirichlet',
'dirichlet_normal']
```

identifiers used to specify the given boundary class

Type list

#### normal: bool = True

determines whether the boundary condition only affects normal components.

If this flag is *False*, boundary conditions must specify values for all components of the field. If *True*, only the normal components at the boundary are specified.

Туре

bool

**class** NormalMixedBC (grid, axis, upper, \*, rank=0, value=0, const=0)

Bases: MixedBC

represents a mixed (or Robin) boundary condition setting the derivative of the normal components in the outward normal direction of the boundary using an affine function involving the actual value:

$$\partial_n c + \gamma c = \beta$$

Here, c is the field to which the condition is applied,  $\gamma$  quantifies the influence of the field and  $\beta$  is the constant term. Note that  $\gamma = 0$  corresponds to Dirichlet conditions imposing  $\beta$  as the derivative. Conversely,  $\gamma \to \infty$  corresponds to imposing a zero value on c.

This condition can be enforced by using one of the following variants

```
bc = {'mixed': VALUE}
bc = {'type': 'mixed', 'value': VALUE, 'const': CONST}
```

where *VALUE* corresponds to  $\gamma$  and *CONST* to  $\beta$ .

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated
- **upper** (bool) Flag indicating whether this boundary condition is associated with the upper side of an axis or not. In essence, this determines the direction of the local normal vector of the boundary.
- **rank** (*int*) The tensorial rank of the field for this boundary condition

- **value** (*float* or str or array) The parameter  $\gamma$  quantifying the influence of the field onto its normal derivative. If *value* is a single value (or tensor in case of tensorial boundary conditions), the same value is applied to all points. Inhomogeneous boundary conditions are possible by supplying an expression as a string, which then may depend on the axes names of the respective grid.
- **const** (float or ndarray or str) The parameter  $\beta$  determining the constant term for the boundary condition. Supports the same input as *value*.

```
names: list[str] = ['normal_mixed', 'normal_robin']
```

identifiers used to specify the given boundary class

Type list

### normal: bool = True

determines whether the boundary condition only affects normal components.

If this flag is *False*, boundary conditions must specify values for all components of the field. If *True*, only the normal components at the boundary are specified.

Type bool

**class** NormalNeumannBC (grid, axis, upper, \*, rank=0, value=0)

Bases: NeumannBC

represents a boundary condition imposing the derivative of normal components in the outward normal direction of the boundary

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur. However, the function is safe when *value* cannot be an arbitrary string.

# **Parameters**

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated
- **upper** (bool) Flag indicating whether this boundary condition is associated with the upper side of an axis or not. In essence, this determines the direction of the local normal vector of the boundary.
- **rank** (*int*) The tensorial rank of the field for this boundary condition
- **normal** (*bool*) Flag indicating whether the condition is only applied in the normal direction.
- **value** (float or str or ndarray) a value stored with the boundary condition. The interpretation of this value depends on the type of boundary condition. If value is a single value (or tensor in case of tensorial boundary conditions), the same value is applied to all points. Inhomogeneous boundary conditions are possible by supplying an expression as a string, which then may depend on the axes names of the respective grid.

```
names: list[str] = ['normal_derivative', 'normal_neumann',
'neumann_normal']
```

identifiers used to specify the given boundary class

#### Type list

### normal: bool = True

determines whether the boundary condition only affects normal components.

If this flag is *False*, boundary conditions must specify values for all components of the field. If *True*, only the normal components at the boundary are specified.

Type bool

**class UserBC** (grid, axis, upper, \*, rank=0)

```
Bases: BCBase
```

represents a boundary whose virtual point are set by the user.

Boundary conditions will only be set when a dictionary {TARGET: value} is supplied as argument *args* to *set\_ghost\_cells()* or the numba equivalent. Here, *TARGET* determines how the *value* is interpreted and what boundary condition is actually enforced: the value of the virtual points directly (*virtual\_point*), the value of the field at the boundary (*value*) or the outward derivative of the field at the boundary (*derivative*).

**Warning:** This implies that the boundary conditions are never enforced automatically, e.g., when evaluating an operator. It is thus the user's responsibility to ensure virtual points are set correctly before operators are applied.

#### **Parameters**

- grid (GridBase) The grid for which the boundary conditions are defined
- **axis** (*int*) The axis to which this boundary condition is associated
- **upper** (bool) Flag indicating whether this boundary condition is associated with the upper side of an axis or not. In essence, this determines the direction of the local normal vector of the boundary.
- rank (int) The tensorial rank of the field for this boundary condition

copy (upper=None, rank=None)

return a copy of itself, but with a reference to the same grid

### Parameters

- **self** (*TBC*) -
- upper (bool / None) -
- rank (int | None) -

Return type

```
get_mathematical_representation (field_name='C')
```

return mathematical representation of the boundary condition

```
Parameters
field_name(str)-
Return type
str
```

# make\_ghost\_cell\_setter()

return function that sets the ghost cells for this boundary

# **Return type**

GhostCellSetter

# make\_virtual\_point\_evaluator()

returns a function evaluating the value at the virtual support point

### Returns

A function that takes the data array and an index marking the current point, which is assumed to be a virtual point. The result is the data value at this point, which is calculated using the boundary condition.

# Return type

function

# names: list[str] = ['user']

identifiers used to specify the given boundary class

Type

list

# set\_ghost\_cells (data\_full, \*, args=None)

set the ghost cell values for this boundary

### **Parameters**

- data\_full (ndarray) The full field data including ghost points
- **args** (ndarray) Determines what boundary conditions are set. *args* should be set to {TARGET: value}. Here, *TARGET* determines how the *value* is interpreted and what boundary condition is actually enforced: the value of the virtual points directly (*virtual\_point*), the value of the field at the boundary (*value*) or the outward derivative of the field at the boundary (*derivative*).

# **Return type**

None

```
to_subgrid(subgrid)
```

converts this boundary condition to one valid for a given subgrid

# Parameters

- subgrid (GridBase) Grid of the new boundary conditions
- **self** (*TBC*) -

### Returns

Boundary conditions valid on the subgrid

### **Return type**

BCBase

# registered\_boundary\_condition\_classes()

returns all boundary condition classes that are currently defined

### Returns

a dictionary with the names of the boundary condition classes

### **Return type**

dict

# registered\_boundary\_condition\_names()

returns all named boundary conditions that are currently defined

### Returns

a dictionary with the names of the boundary conditions that can be used

**Return type** 

dict

# 4.2.2 pde.grids.coordinates package

Package collecting classes representing orthonormal coordinate systems

BipolarCoordinates	2-dimensional bipolar coordinates
BisphericalCoordinates	3-dimensional bispherical coordinates
CartesianCoordinates	n-dimensional Cartesian coordinates
CylindricalCoordinates	n-dimensional Cartesian coordinates
PolarCoordinates	2-dimensional polar coordinates
SphericalCoordinates	3-dimensional spherical coordinates

# pde.grids.coordinates.base module

# class CoordinatesBase

Bases: object

Base class for orthonormal coordinate systems

# axes: list[str]

name of each coordinate axis

Type list

basis\_rotation (points)

returns rotation matrix rotating basis vectors to Cartesian coordinates

# Parameters

points (ndarray) - Coordinates of the point(s)

# Returns

Rotation matrices for all points. The returnd array has the shape  $(dim, dim) + points\_shape$ , assuming *points* has the shape *points\\_shape* + (dim,).

# **Return type**

ndarray

**cell\_volume** (*c\_low*, *c\_high*)

calculate the volume between coordinate lines

### **Parameters**

- **c\_low** (ndarray) Lower values of the coordinate lines enclosing the volume
- c\_high (ndarray) Upper values of the coordinate lines enclosing the volume

### Returns

Enclosed volumes for all given cells

# Return type

ndarray

# coordinate\_limits: list[tuple[float, float]]

the limits of each coordinate axis

# Type

list of tuple

# dim: int

spatial dimension of the coordinate system

Type int

### distance (p1, p2)

Calculate the distance between two points

# Parameters

• p1 (ndarray) - First position

• p2 (ndarray) - Second position

# Returns

Distance between the two positions

Return type

float

# mapping\_jacobian (points)

returns the Jacobian matrix of the coordinate mapping

### Parameters

points (ndarray) - Coordinates of the point(s)

### Returns

The Jacobian

# **Return type**

ndarray

# metric (points)

calculate the metric tensor at coordinate points

# Parameters

points (ndarray) - The coordinates of the points

# Returns

Metric tensor at the points

# **Return type**

ndarray

# $\texttt{pos\_diff}\left(pl,p2\right)$

return Cartesian vector(s) pointing from p1 to p2

- p1 (ndarray) First point(s)
- p2 (ndarray) Second point(s)

### Returns

The difference vectors between the points with periodic boundary conditions applied.

### **Return type**

ndarray

# pos\_from\_cart (points)

convert Cartesian coordinates to coordinates in this system

# Parameters

points (ndarray) – Points given in Cartesian coordinates.

### Returns

Points given in the coordinates of this system

### Return type

ndarray

# pos\_to\_cart (points)

convert coordinates to Cartesian coordinates

# Parameters

points (ndarray) - The coordinates of points in the current coordinate system

### Returns

Cartesian coordinates of the points

# **Return type**

ndarray

# scale\_factors (points)

calculate the scale factors at various points

### Parameters

points (ndarray) - The coordinates of the points

# Returns

Scale factors at the points

# Return type

ndarray

# vec\_to\_cart (points, components)

convert the vectors at given points to a Cartesian basis

# **Parameters**

- points (ndarray) The coordinates of the point(s) where the vectors are specified.
- components (ndarray) The components of the vectors at the given points

# Returns

The vectors specified at the same position but with components given in Cartesian coordinates.

### Return type

ndarray

# volume\_factor(points)

calculate the volume factors at various points

### Parameters

points (ndarray) - Coordinates of the point(s)

**Returns** Volume factors at the points

Return type

ndarray

# exception DimensionError

Bases: ValueError

exception indicating that dimensions were inconsistent

# pde.grids.coordinates.bipolar module

# class BipolarCoordinates (scale\_parameter=1)

Bases: CoordinatesBase

2-dimensional bipolar coordinates

# Parameters scale\_parameter(float)-

axes: list[str] = [' $\sigma$ ', ' $\tau$ ']

name of each coordinate axis

Type list

```
coordinate_limits: list[tuple[float, float]] = [(0, 6.283185307179586),
(-inf, inf)]
```

the limits of each coordinate axis

Туре

list of tuple

dim: int = 2

spatial dimension of the coordinate system

Туре

int

# pde.grids.coordinates.bispherical module

```
class BisphericalCoordinates(scale_parameter=1)
```

Bases: CoordinatesBase

3-dimensional bispherical coordinates

```
Parameters
    scale_parameter(float) -
axes: list[str] = ['σ', 'τ', 'ψ']
    name of each coordinate axis
```

Type list

```
coordinate_limits: list[tuple[float, float]] = [(0, 3.141592653589793),
(-inf, inf), (0, 6.283185307179586)]
```

the limits of each coordinate axis

Туре

list of tuple

# dim: int = 3

spatial dimension of the coordinate system

Type int

# pde.grids.coordinates.cartesian module

### class CartesianCoordinates(dim)

```
Bases: CoordinatesBase
```

n-dimensional Cartesian coordinates

# Parameters

dim (int) – Dimension of the Cartesian coordinate system

# pde.grids.coordinates.cylindrical module

### class CylindricalCoordinates

Bases: CoordinatesBase

n-dimensional Cartesian coordinates

```
axes: list[str] = ['r', '\psi', 'z']
```

name of each coordinate axis

Type list

```
coordinate_limits: list[tuple[float, float]] = [(0, inf), (0,
6.283185307179586), (-inf, inf)]
```

the limits of each coordinate axis

# Туре

list of tuple

# dim: int = 3

spatial dimension of the coordinate system

Type int

# pde.grids.coordinates.polar module

### class PolarCoordinates

Bases: CoordinatesBase

2-dimensional polar coordinates

axes:  $list[str] = ['r', '\psi']$ 

name of each coordinate axis

Type list

```
coordinate_limits: list[tuple[float, float]] = [(0, inf), (0,
6.283185307179586)]
```

the limits of each coordinate axis

# Туре

list of tuple

dim: int = 2

spatial dimension of the coordinate system

Type int

### pde.grids.coordinates.spherical module

### class SphericalCoordinates

Bases: CoordinatesBase

3-dimensional spherical coordinates

```
axes: list[str] = ['r', '\theta', '\psi']
```

name of each coordinate axis

Type list

```
coordinate_limits: list[tuple[float, float]] = [(0, inf), (0,
3.141592653589793), (0, 6.283185307179586)]
```

the limits of each coordinate axis

Type

list of tuple

```
dim: int = 3
```

spatial dimension of the coordinate system

Type int

major\_axis = 0

# 4.2.3 pde.grids.operators package

Package collecting modules defining discretized operators for different grids.

These operators can either be used directly or they are imported by the respective methods defined on fields and grids.

cartesian	This module implements differential operators on Carte- sian grids
cylindrical_sym	This module implements differential operators on cylin- drical grids
polar_sym	This module implements differential operators on polar grids
spherical_sym	This module implements differential operators on spher- ical grids
common.make_derivative	make a derivative operator along a single axis using numba compilation
common.make_derivative2	make a second-order derivative operator along a single axis

# pde.grids.operators.cartesian module

This module implements differential operators on Cartesian grids

make_laplace	make a Laplace operator on a Cartesian grid
make_gradient	make a gradient operator on a Cartesian grid
make_divergence	make a divergence operator on a Cartesian grid
make_vector_gradient	make a vector gradient operator on a Cartesian grid
make_vector_laplace	make a vector Laplacian on a Cartesian grid
make_tensor_divergence	make a tensor divergence operator on a Cartesian grid
make_poisson_solver	make a operator that solves Poisson's equation

# make\_divergence (grid, backend='auto')

make a divergence operator on a Cartesian grid

# Parameters

- grid (CartesianGrid) The grid for which the operator is created
- **backend** (*str*) Backend used for calculating the divergence operator. If backend='auto', a suitable backend is chosen automatically.

# Returns

A function that can be applied to an array of values

# **Return type**

OperatorType

# make\_gradient (grid, backend='auto')

make a gradient operator on a Cartesian grid

- grid (CartesianGrid) The grid for which the operator is created
- **backend** (*str*) Backend used for calculating the gradient operator. If backend='auto', a suitable backend is chosen automatically.

# Returns

A function that can be applied to an array of values

# **Return type**

OperatorType

# make\_laplace (grid, backend='auto')

make a Laplace operator on a Cartesian grid

### Parameters

- grid (CartesianGrid) The grid for which the operator is created
- **backend** (*str*) Backend used for calculating the Laplace operator. If backend='auto', a suitable backend is chosen automatically.

### Returns

A function that can be applied to an array of values

# **Return type**

OperatorType

# make\_poisson\_solver(bcs, method='auto')

make a operator that solves Poisson's equation

# Parameters

- **bcs** (Boundaries) {ARG\_BOUNDARIES\_INSTANCE}
- **method** (*str*) Method used for calculating the tensor divergence operator. If method='auto', a suitable method is chosen automatically.

### Returns

A function that can be applied to an array of values

### **Return type**

OperatorType

### make\_tensor\_divergence (grid, backend='numba')

make a tensor divergence operator on a Cartesian grid

# Parameters

- grid (CartesianGrid) The grid for which the operator is created
- **backend** (*str*) Backend used for calculating the tensor divergence operator.

### Returns

A function that can be applied to an array of values

### **Return type**

OperatorType

# make\_vector\_gradient (grid, backend='numba')

make a vector gradient operator on a Cartesian grid

# Parameters

- grid (CartesianGrid) The grid for which the operator is created
- **backend** (*str*) Backend used for calculating the vector gradient operator.

### Returns

A function that can be applied to an array of values

# **Return type**

OperatorType

make\_vector\_laplace (grid, backend='numba')

make a vector Laplacian on a Cartesian grid

# Parameters

- grid (CartesianGrid) The grid for which the operator is created
- **backend** (*str*) Backend used for calculating the vector Laplace operator.

# Returns

A function that can be applied to an array of values

# **Return type**

OperatorType

# pde.grids.operators.common module

Common functions that are used by many operators

# make\_derivative (grid, axis=0, method='central')

make a derivative operator along a single axis using numba compilation

# Parameters

- grid (GridBase) The grid for which the operator is created
- **axis** (*int*) The axis along which the derivative will be taken
- **method** (*str*) The method for calculating the derivative. Possible values are 'central', 'forward', and 'backward'.

# Returns

A function that can be applied to an full array of values including those at ghost cells. The result will be an array of the same shape containing the actual derivatives at the valid (interior) grid points.

### **Return type**

# OperatorType

# make\_derivative2 (grid, axis=0)

make a second-order derivative operator along a single axis

# Parameters

- grid (GridBase) The grid for which the operator is created
- **axis** (*int*) The axis along which the derivative will be taken

### Returns

A function that can be applied to an full array of values including those at ghost cells. The result will be an array of the same shape containing the actual derivatives at the valid (interior) grid points.

### **Return type**

OperatorType

make\_general\_poisson\_solver (matrix, vector, method='auto')

make an operator that solves Poisson's problem

# Parameters

• matrix - The (sparse) matrix representing the laplace operator on the given grid.

- vector The constant part representing the boundary conditions of the Laplace operator.
- **method** (*str*) The chosen method for implementing the operator

# Returns

A function that can be applied to an array of values to obtain the solution to Poisson's equation where the array is used as the right hand side

# **Return type**

OperatorType

### make\_laplace\_from\_matrix (matrix, vector)

make a Laplace operator using matrix vector products

# Parameters

- matrix (Sparse) matrix representing the laplace operator on the given grid
- vector Constant part representing the boundary conditions of the Laplace operator

#### Returns

A function that can be applied to an array of values to obtain the result of applying the linear operator *matrix* and the offset given by *vector*.

# **Return type**

Callable[[ndarray, ndarray | None], ndarray]

### uniform\_discretization (grid)

returns the uniform discretization or raises RuntimeError

### **Parameters**

grid (GridBase) - The grid whose discretization is tested

### Raises

RuntimeError -

### Returns

the common discretization of all axes

# **Return type**

float

# pde.grids.operators.cylindrical\_sym module

This module implements differential operators on cylindrical grids

make_laplace	make a discretized laplace operator for a cylindrical grid
make_gradient	make a discretized gradient operator for a cylindrical grid
make_divergence	make a discretized divergence operator for a cylindrical grid
make_vector_gradient	make a discretized vector gradient operator for a cylindri- cal grid
make_vector_laplace	make a discretized vector laplace operator for a cylindrical grid
make_tensor_divergence	make a discretized tensor divergence operator for a cylin- drical grid
make_poisson_solver	make a operator that solves Poisson's equation

# make\_divergence (grid)

make a discretized divergence operator for a cylindrical grid

The cylindrical grid assumes polar symmetry, so that fields only depend on the radial coordinate r and the axial coordinate z. Here, the first axis is along the radius, while the second axis is along the axis of the cylinder. The radial discretization is defined as  $r_i = (i + \frac{1}{2})\Delta r$  for  $i = 0, ..., N_r - 1$ .

# Parameters

grid (CylindricalSymGrid) – The grid for which the operator is created

### Returns

A function that can be applied to an array of values

Return type

OperatorType

# make\_gradient (grid)

make a discretized gradient operator for a cylindrical grid

The cylindrical grid assumes polar symmetry, so that fields only depend on the radial coordinate r and the axial coordinate z. Here, the first axis is along the radius, while the second axis is along the axis of the cylinder. The radial discretization is defined as  $r_i = (i + \frac{1}{2})\Delta r$  for  $i = 0, ..., N_r - 1$ .

### **Parameters**

grid (CylindricalSymGrid) – The grid for which the operator is created

# Returns

A function that can be applied to an array of values

# Return type

OperatorType

# make\_gradient\_squared (grid, central=True)

make a discretized gradient squared operator for a cylindrical grid

The cylindrical grid assumes polar symmetry, so that fields only depend on the radial coordinate r and the axial coordinate z. Here, the first axis is along the radius, while the second axis is along the axis of the cylinder. The radial discretization is defined as  $r_i = (i + \frac{1}{2})\Delta r$  for  $i = 0, ..., N_r - 1$ .

# Parameters

- grid (CylindricalSymGrid) The grid for which the operator is created
- **central** (bool) Whether a central difference approximation is used for the gradient operator. If this is False, the squared gradient is calculated as the mean of the squared values of the forward and backward derivatives.

### Returns

A function that can be applied to an array of values

# **Return type**

OperatorType

### make\_laplace(grid)

make a discretized laplace operator for a cylindrical grid

The cylindrical grid assumes polar symmetry, so that fields only depend on the radial coordinate r and the axial coordinate z. Here, the first axis is along the radius, while the second axis is along the axis of the cylinder. The radial discretization is defined as  $r_i = (i + \frac{1}{2})\Delta r$  for  $i = 0, ..., N_r - 1$ .

### Parameters

grid (CylindricalSymGrid) – The grid for which the operator is created

### Returns

A function that can be applied to an array of values

# **Return type**

OperatorType

### make\_poisson\_solver(bcs, method='auto')

make a operator that solves Poisson's equation

The cylindrical grid assumes polar symmetry, so that fields only depend on the radial coordinate r and the axial coordinate z. Here, the first axis is along the radius, while the second axis is along the axis of the cylinder. The radial discretization is defined as  $r_i = (i + \frac{1}{2})\Delta r$  for  $i = 0, ..., N_r - 1$ .

### Parameters

- **bcs** (*Boundaries*) Specifies the boundary conditions applied to the field. This must be an instance of *Boundaries*, which can be created from various data formats using the class method *from\_data()*.
- method (str) The chosen method for implementing the operator

### Returns

A function that can be applied to an array of values

#### Return type

OperatorType

# make\_tensor\_divergence (grid)

make a discretized tensor divergence operator for a cylindrical grid

The cylindrical grid assumes polar symmetry, so that fields only depend on the radial coordinate r and the axial coordinate z. Here, the first axis is along the radius, while the second axis is along the axis of the cylinder. The radial discretization is defined as  $r_i = (i + \frac{1}{2})\Delta r$  for  $i = 0, ..., N_r - 1$ .

# Parameters

grid (CylindricalSymGrid) – The grid for which the operator is created

### Returns

A function that can be applied to an array of values

### **Return type**

OperatorType

#### make\_vector\_gradient (grid)

make a discretized vector gradient operator for a cylindrical grid

The cylindrical grid assumes polar symmetry, so that fields only depend on the radial coordinate r and the axial coordinate z. Here, the first axis is along the radius, while the second axis is along the axis of the cylinder. The radial discretization is defined as  $r_i = (i + \frac{1}{2})\Delta r$  for  $i = 0, ..., N_r - 1$ .

### Parameters

grid (CylindricalSymGrid) – The grid for which the operator is created

### Returns

A function that can be applied to an array of values

**Return type** 

OperatorType

# make\_vector\_laplace (grid)

make a discretized vector laplace operator for a cylindrical grid

The cylindrical grid assumes polar symmetry, so that fields only depend on the radial coordinate r and the axial coordinate z. Here, the first axis is along the radius, while the second axis is along the axis of the cylinder. The radial discretization is defined as  $r_i = (i + \frac{1}{2})\Delta r$  for  $i = 0, ..., N_r - 1$ .

#### Parameters

grid (CylindricalSymGrid) – The grid for which the operator is created

#### Returns

A function that can be applied to an array of values

Return type

OperatorType

## pde.grids.operators.polar\_sym module

This module implements differential operators on polar grids

make_laplace	make a discretized laplace operator for a polar grid
make_gradient	make a discretized gradient operator for a polar grid
make_divergence	make a discretized divergence operator for a polar grid
make_vector_gradient	make a discretized vector gradient operator for a polar grid
make_tensor_divergence	make a discretized tensor divergence operator for a polar grid
make_poisson_solver	make a operator that solves Poisson's equation

## make\_divergence (grid)

make a discretized divergence operator for a polar grid

The polar grid assumes polar symmetry, so that fields only depend on the radial coordinate r. The radial discretization is defined as  $r_i = r_{\min} + (i + \frac{1}{2})\Delta r$  for  $i = 0, ..., N_r - 1$ , where  $r_{\min}$  is the radius of the inner boundary, which is zero by default. Note that the radius of the outer boundary is given by  $r_{\max} = r_{\min} + N_r \Delta r$ .

#### Parameters

grid (PolarSymGrid) – The polar grid for which this operator will be defined

#### Returns

A function that can be applied to an array of values

## **Return type**

OperatorType

## make\_gradient (grid)

make a discretized gradient operator for a polar grid

The polar grid assumes polar symmetry, so that fields only depend on the radial coordinate r. The radial discretization is defined as  $r_i = r_{\min} + (i + \frac{1}{2})\Delta r$  for  $i = 0, ..., N_r - 1$ , where  $r_{\min}$  is the radius of the inner boundary, which is zero by default. Note that the radius of the outer boundary is given by  $r_{\max} = r_{\min} + N_r \Delta r$ .

#### Parameters

grid (PolarSymGrid) – The polar grid for which this operator will be defined

#### Returns

A function that can be applied to an array of values

#### **Return type**

OperatorType

### make\_gradient\_squared(grid, central=True)

make a discretized gradient squared operator for a polar grid

The polar grid assumes polar symmetry, so that fields only depend on the radial coordinate r. The radial discretization is defined as  $r_i = r_{\min} + (i + \frac{1}{2})\Delta r$  for  $i = 0, ..., N_r - 1$ , where  $r_{\min}$  is the radius of the inner boundary, which is zero by default. Note that the radius of the outer boundary is given by  $r_{\max} = r_{\min} + N_r \Delta r$ .

#### Parameters

- grid (PolarSymGrid) The polar grid for which this operator will be defined
- **central** (*bool*) Whether a central difference approximation is used for the gradient operator. If this is False, the squared gradient is calculated as the mean of the squared values of the forward and backward derivatives.

#### Returns

A function that can be applied to an array of values

#### **Return type**

OperatorType

## make\_laplace(grid)

make a discretized laplace operator for a polar grid

The polar grid assumes polar symmetry, so that fields only depend on the radial coordinate r. The radial discretization is defined as  $r_i = r_{\min} + (i + \frac{1}{2})\Delta r$  for  $i = 0, ..., N_r - 1$ , where  $r_{\min}$  is the radius of the inner boundary, which is zero by default. Note that the radius of the outer boundary is given by  $r_{\max} = r_{\min} + N_r \Delta r$ .

#### Parameters

grid (PolarSymGrid) – The polar grid for which this operator will be defined

#### Returns

A function that can be applied to an array of values

#### **Return type**

OperatorType

#### make\_poisson\_solver(bcs, method='auto')

make a operator that solves Poisson's equation

The polar grid assumes polar symmetry, so that fields only depend on the radial coordinate r. The radial discretization is defined as  $r_i = r_{\min} + (i + \frac{1}{2})\Delta r$  for  $i = 0, ..., N_r - 1$ , where  $r_{\min}$  is the radius of the inner boundary, which is zero by default. Note that the radius of the outer boundary is given by  $r_{\max} = r_{\min} + N_r \Delta r$ .

#### Parameters

- **bcs** (*Boundaries*) Specifies the boundary conditions applied to the field. This must be an instance of *Boundaries*, which can be created from various data formats using the class method *from\_data()*.
- **method** (*str*) The chosen method for implementing the operator

#### Returns

A function that can be applied to an array of values

#### **Return type**

OperatorType

## make\_tensor\_divergence (grid)

make a discretized tensor divergence operator for a polar grid

The polar grid assumes polar symmetry, so that fields only depend on the radial coordinate r. The radial discretization is defined as  $r_i = r_{\min} + (i + \frac{1}{2})\Delta r$  for  $i = 0, ..., N_r - 1$ , where  $r_{\min}$  is the radius of the inner boundary, which is zero by default. Note that the radius of the outer boundary is given by  $r_{\max} = r_{\min} + N_r \Delta r$ .

## Parameters

grid (PolarSymGrid) – The polar grid for which this operator will be defined

## Returns

A function that can be applied to an array of values

## Return type

OperatorType

## make\_vector\_gradient (grid)

make a discretized vector gradient operator for a polar grid

The polar grid assumes polar symmetry, so that fields only depend on the radial coordinate r. The radial discretization is defined as  $r_i = r_{\min} + (i + \frac{1}{2})\Delta r$  for  $i = 0, ..., N_r - 1$ , where  $r_{\min}$  is the radius of the inner boundary, which is zero by default. Note that the radius of the outer boundary is given by  $r_{\max} = r_{\min} + N_r \Delta r$ .

## **Parameters**

grid (PolarSymGrid) – The polar grid for which this operator will be defined

## Returns

A function that can be applied to an array of values

## Return type

OperatorType

## pde.grids.operators.spherical\_sym module

This module implements differential operators on spherical grids

make_laplace	make a discretized laplace operator for a spherical grid
make_gradient	make a discretized gradient operator for a spherical grid
make_divergence	make a discretized divergence operator for a spherical grid
make_vector_gradient	make a discretized vector gradient operator for a spherical grid
make_tensor_divergence	make a discretized tensor divergence operator for a spher- ical grid
make_poisson_solver	make a operator that solves Poisson's equation

## make\_divergence (grid, safe=True, conservative=True)

make a discretized divergence operator for a spherical grid

The spherical grid assumes spherical symmetry, so that fields only depend on the radial coordinate r. The radial discretization is defined as  $r_i = r_{\min} + (i + \frac{1}{2})\Delta r$  for  $i = 0, \ldots, N_r - 1$ , where  $r_{\min}$  is the radius of the inner boundary, which is zero by default. Note that the radius of the outer boundary is given by  $r_{\max} = r_{\min} + N_r \Delta r$ .

**Warning:** This operator ignores the  $\theta$ -component of the field when calculating the divergence. This is because the resulting scalar field could not be expressed on a SphericalSymGrid.

#### **Parameters**

• grid (SphericalSymGrid) – The polar grid for which this operator will be defined

- **safe** (bool) Add extra checks for the validity of the input
- **conservative** (bool) Flag indicating whether the operator should be conservative (which results in slightly slower computations). Conservative operators ensure mass conservation.

## Returns

A function that can be applied to an array of values

#### **Return type**

OperatorType

## make\_gradient (grid)

make a discretized gradient operator for a spherical grid

The spherical grid assumes spherical symmetry, so that fields only depend on the radial coordinate r. The radial discretization is defined as  $r_i = r_{\min} + (i + \frac{1}{2})\Delta r$  for  $i = 0, ..., N_r - 1$ , where  $r_{\min}$  is the radius of the inner boundary, which is zero by default. Note that the radius of the outer boundary is given by  $r_{\max} = r_{\min} + N_r \Delta r$ .

#### Parameters

grid (SphericalSymGrid) – The polar grid for which this operator will be defined

#### Returns

A function that can be applied to an array of values

#### **Return type**

OperatorType

#### make\_gradient\_squared(grid, central=True)

make a discretized gradient squared operator for a spherical grid

The spherical grid assumes spherical symmetry, so that fields only depend on the radial coordinate r. The radial discretization is defined as  $r_i = r_{\min} + (i + \frac{1}{2})\Delta r$  for  $i = 0, ..., N_r - 1$ , where  $r_{\min}$  is the radius of the inner boundary, which is zero by default. Note that the radius of the outer boundary is given by  $r_{\max} = r_{\min} + N_r \Delta r$ .

#### Parameters

- grid (SphericalSymGrid) The polar grid for which this operator will be defined
- **central** (*bool*) Whether a central difference approximation is used for the gradient operator. If this is False, the squared gradient is calculated as the mean of the squared values of the forward and backward derivatives.

#### Returns

A function that can be applied to an array of values

#### **Return type**

OperatorType

#### make\_laplace (grid, conservative=True)

make a discretized laplace operator for a spherical grid

The spherical grid assumes spherical symmetry, so that fields only depend on the radial coordinate r. The radial discretization is defined as  $r_i = r_{\min} + (i + \frac{1}{2})\Delta r$  for  $i = 0, ..., N_r - 1$ , where  $r_{\min}$  is the radius of the inner boundary, which is zero by default. Note that the radius of the outer boundary is given by  $r_{\max} = r_{\min} + N_r \Delta r$ .

- grid (SphericalSymGrid) The polar grid for which this operator will be defined
- **conservative** (bool) Flag indicating whether the laplace operator should be conservative (which results in slightly slower computations). Conservative operators ensure mass conservation.

## Returns

A function that can be applied to an array of values

## **Return type**

OperatorType

## make\_poisson\_solver(bcs, method='auto')

make a operator that solves Poisson's equation

The polar grid assumes polar symmetry, so that fields only depend on the radial coordinate r. The radial discretization is defined as  $r_i = r_{\min} + (i + \frac{1}{2})\Delta r$  for  $i = 0, ..., N_r - 1$ , where  $r_{\min}$  is the radius of the inner boundary, which is zero by default. Note that the radius of the outer boundary is given by  $r_{\max} = r_{\min} + N_r \Delta r$ .

#### **Parameters**

- **bcs** (*Boundaries*) Specifies the boundary conditions applied to the field. This must be an instance of *Boundaries*, which can be created from various data formats using the class method *from\_data()*.
- **method** (*str*) The chosen method for implementing the operator

#### Returns

A function that can be applied to an array of values

## **Return type**

## OperatorType

#### make\_tensor\_divergence (grid, safe=True, conservative=False)

make a discretized tensor divergence operator for a spherical grid

The spherical grid assumes spherical symmetry, so that fields only depend on the radial coordinate r. The radial discretization is defined as  $r_i = r_{\min} + (i + \frac{1}{2})\Delta r$  for  $i = 0, ..., N_r - 1$ , where  $r_{\min}$  is the radius of the inner boundary, which is zero by default. Note that the radius of the outer boundary is given by  $r_{\max} = r_{\min} + N_r \Delta r$ .

#### **Parameters**

- grid (SphericalSymGrid) The polar grid for which this operator will be defined
- **safe** (*bool*) Add extra checks for the validity of the input
- **conservative** (bool) Flag indicating whether the operator should be conservative (which results in slightly slower computations). Conservative operators ensure mass conservation.

#### Returns

A function that can be applied to an array of values

#### **Return type**

OperatorType

#### make\_tensor\_double\_divergence (grid, safe=True, conservative=True)

make a discretized tensor double divergence operator for a spherical grid

The spherical grid assumes spherical symmetry, so that fields only depend on the radial coordinate r. The radial discretization is defined as  $r_i = r_{\min} + (i + \frac{1}{2})\Delta r$  for  $i = 0, ..., N_r - 1$ , where  $r_{\min}$  is the radius of the inner boundary, which is zero by default. Note that the radius of the outer boundary is given by  $r_{\max} = r_{\min} + N_r \Delta r$ .

- grid (SphericalSymGrid) The polar grid for which this operator will be defined
- **safe** (*bool*) Add extra checks for the validity of the input

• **conservative** (bool) – Flag indicating whether the operator should be conservative (which results in slightly slower computations). Conservative operators ensure mass conservation.

## Returns

A function that can be applied to an array of values

### **Return type**

OperatorType

## make\_vector\_gradient (grid, safe=True)

make a discretized vector gradient operator for a spherical grid

**Warning:** This operator ignores the two angular components of the field when calculating the gradient. This is because the resulting field could not be expressed on a SphericalSymGrid.

The spherical grid assumes spherical symmetry, so that fields only depend on the radial coordinate r. The radial discretization is defined as  $r_i = r_{\min} + (i + \frac{1}{2})\Delta r$  for  $i = 0, ..., N_r - 1$ , where  $r_{\min}$  is the radius of the inner boundary, which is zero by default. Note that the radius of the outer boundary is given by  $r_{\max} = r_{\min} + N_r \Delta r$ .

## Parameters

- grid (SphericalSymGrid) The polar grid for which this operator will be defined
- **safe** (bool) Add extra checks for the validity of the input

#### Returns

A function that can be applied to an array of values

#### **Return type**

OperatorType

## 4.2.4 pde.grids.base module

Defines the base class for all grids

## exception DomainError

Bases: ValueError

exception indicating that point lies outside domain

## class GridBase

Bases: object

Base class for all grids defining common methods and interfaces

initialize the grid

#### assert\_grid\_compatible(other)

checks whether other is compatible with the current grid

## Parameters

other (GridBase) – The grid compared to this one

Raises

ValueError – if grids are not compatible

Return type None

## axes: list[str]

Names of all axes that are described by the grid

Туре

list

## property axes\_bounds: tuple[tuple[float, float], ...]

lower and upper bounds of each axis

Type

tuple

## property axes\_coords: tuple[ndarray, ...]

coordinates of the cells for each axis

Туре

tuple

## axes\_symmetric: list[str] = []

The names of the additional axes that the fields do not depend on, e.g. along which they are constant.

Type list

#### boundary\_names: dict[str, tuple[int, bool]] = {}

Names of boundaries to select them conveniently

Туре

dict

## c: CoordinatesBase

Coordinates of the grid

Туре

CoordinatesBase

## cell\_coords

coordinate values for all axes of each cell

Туре

#### ndarray

## cell\_volume\_data: Sequence[FloatNumerical] | None

Information about the size of discretization cells

Type list

## cell\_volumes

volume of each cell

## Туре

ndarray

#### compatible\_with (other)

tests whether this grid is compatible with other grids.

Grids are compatible when they cover the same area with the same discretization. The difference to equality is that compatible grids do not need to have the same periodicity in their boundaries.

## Parameters

other (GridBase) - The other grid to test against

## Returns

Whether the grid is compatible

Return type bool

contains\_point (points, \*, coords='cartesian', full=False)

check whether the point is contained in the grid

## Parameters

- point (ndarray) Coordinates of the point
- coords (*str*) The coordinate system in which the points are given
- full (bool) Indicates whether coordinates along symmetric axes are specified
- **points** (*ndarray*) -

## Returns

A boolean array indicating which points lie within the grid

## **Return type**

ndarray

## coordinate\_arrays

for each axes: coordinate values for all cells

## Type

tuple

## coordinate\_constraints: list[int] = []

axes that not described explicitly

## Type list

## copy()

return a copy of the grid

Return type GridBase

## difference\_vector (p1, p2, \*, coords='grid')

return Cartesian vector(s) pointing from p1 to p2

In case of periodic boundary conditions, the shortest vector is returned.

## Parameters

- **p1** (ndarray) First point(s)
- p2 (ndarray) Second point(s)
- **coords** (*str*) The coordinate system in which the points are specified. Valid values are *cartesian, cell, and grid; see transform()*.

## Returns

The difference vectors between the points with periodic boundary conditions applied.

## **Return type**

ndarray

## difference\_vector\_real (p1, p2)

## **Parameters**

• **p1** (ndarray) -

• **p2** (ndarray) -

## **Return type**

ndarray

## property dim: int

The spatial dimension in which the grid is embedded

Туре

int

## property discretization: ndarray

the linear size of a cell along each axis

Туре

numpy.array

## distance (p1, p2, \*, coords='grid')

Calculate the distance between two points given in real coordinates

This takes periodic boundary conditions into account if necessary.

## **Parameters**

- **p1** (ndarray) First position
- p2 (ndarray) Second position
- **coords** (*str*) The coordinate system in which the points are specified. Valid values are *cartesian, cell,* and *grid*; see *transform()*.

## Returns

Distance between the two positions

#### **Return type**

float

 $distance_real(p1, p2)$ 

## **Parameters**

- p1 (ndarray) -
- p2 (ndarray) -

Return type

float

## classmethod from\_bounds (bounds, shape, periodic)

## **Parameters**

- **bounds** (Sequence [tuple [float, float]]) -
- **shape**(Sequence[int])-
- periodic (Sequence [bool]) -

## **Return type**

GridBase

#### classmethod from\_state(state)

create a field from a stored state.

#### **Parameters**

**state** (*str* or *dict*) – The state from which the grid is reconstructed. If *state* is a string, it is decoded as JSON, which should yield a *dict*.

## Returns

Grid re-created from the state

## Return type

GridBase

#### get\_axis\_index (key, allow\_symmetric=True)

return the index belonging to an axis

## **Parameters**

- key (int or str) The index or name of an axis
- allow\_symmetric (bool) Whether axes with assumed symmetry are included

## Returns

The index of the axis

#### **Return type**

int

get\_boundary\_conditions (bc='auto\_periodic\_neumann', rank=0)

constructs boundary conditions from a flexible data format

#### Parameters

- **bc** (*str* or *list* or *tuple* or *dict*) The boundary conditions applied to the field. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by *{'value': NUM})* and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by *{'derivative': DERIV})* are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documentation*.
- **rank** (*int*) The tensorial rank of the value associated with the boundary conditions.

#### Returns

The boundary conditions for all axes.

## **Return type**

Boundaries

## Raises

- ValueError If the data given in *bc* cannot be read
- **PeriodicityError** If the boundaries are not compatible with the periodic axes of the grid.

get\_image\_data(data)

return a 2d-image of the data

### Parameters

data (ndarray) - The values at the grid points

## Returns

A dictionary with information about the data convenient for plotting.

Return type

## get\_line\_data (data, extract='auto')

return a line cut through the grid

## Parameters

- data (ndarray) The values at the grid points
- **extract** (*str*) Determines which cut is done through the grid. Possible choices depend on the actual grid.

#### Returns

A dictionary with information about the line cut, which is convenient for plotting.

**Return type** 

dict

```
get_random_point (*, boundary_distance=0, coords='cartesian', rng=None)
```

return a random point within the grid

#### **Parameters**

- **boundary\_distance** (*float*) The minimal distance this point needs to have from all boundaries.
- **coords** (*str*) Determines the coordinate system in which the point is specified. Valid values are *cartesian*, *cell*, and *grid*; see *transform()*.
- rng (Generator) Random number generator (default: default\_rng())

## Returns

The coordinates of the random point

## Return type

ndarray

## get\_vector\_data (data, \*\*kwargs)

return data to visualize vector field

## **Parameters**

- data (ndarray) The vectorial values at the grid points
- **\*\*kwargs** Arguments forwarded to get\_image\_data().

## Returns

A dictionary with information about the data convenient for plotting.

## **Return type**

dict

#### integrate (data, axes=None)

Integrates the discretized data over the grid

## Parameters

• data (ndarray) - The values at the support points of the grid that need to be integrated.

• **axes** (list of int, optional) – The axes along which the integral is performed. If omitted, all axes are integrated over.

## Returns

The values integrated over the entire grid

#### **Return type**

ndarray

iter\_mirror\_points (point, with\_self=False, only\_periodic=True)

generates all mirror points corresponding to point

## Parameters

- point (ndarray) The point within the grid
- with\_self (bool) Whether to include the point itself
- only\_periodic (bool) Whether to only mirror along periodic axes

#### Returns

A generator yielding the coordinates that correspond to mirrors

#### **Return type**

Generator

#### make\_cell\_volume\_compiled (flat\_index=False)

return a compiled function returning the volume of a grid cell

#### Parameters

**flat\_index** (*bool*) – When True, cell\_volumes are indexed by a single integer into the flattened array.

## Returns

returning the volume of the chosen cell

## **Return type**

function

#### make\_inserter\_compiled (\*, with\_ghost\_cells=False)

return a compiled function to insert values at interpolated positions

#### **Parameters**

with\_ghost\_cells (bool) – Flag indicating that the interpolator should work on the full data array that includes values for the grid points. If this is the case, the boundaries are not checked and the coordinates are used as is.

#### Returns

A function with signature (data, position, amount), where *data* is the numpy array containing the field data, position is denotes the position in grid coordinates, and *amount* is the that is to be added to the field.

#### Return type

callable

## make\_integrator()

return function that can be used to integrates discretized data over the grid

If this function is used in a multiprocessing run (using MPI), the integrals are performed on all subgrids and then accumulated. Each process then receives the same value representing the global integral.

## Returns

A function that takes a numpy array and returns the integral with the correct weights given by the cell volumes.

#### **Return type**

callable

## make\_normalize\_point\_compiled (reflect=True)

return a compiled function that normalizes a point

Here, the point is assumed to be specified by the physical values along the non-symmetric axes of the grid. Normalizing points is useful to make sure they lie within the domain of the grid. This function respects periodic boundary conditions and can also reflect points off the boundary.

## **Parameters**

**reflect** (*bool*) – Flag determining whether coordinates along non-periodic axes are reflected to lie in the valid range. If *False*, such coordinates are left unchanged and only periodic boundary conditions are enforced.

## Returns

A function that takes a ndarray as an argument, which describes the coordinates of the points. This array is modified in-place!

## **Return type**

callable

## make\_operator (operator, bc, \*\*kwargs)

return a compiled function applying an operator with boundary conditions

The returned function takes the discretized data on the grid as an input and returns the data to which the operator *operator* has been applied. The function only takes the valid grid points and allocates memory for the ghost points internally to apply the boundary conditions specified as *bc*. Note that the function supports an optional argument *out*, which if given should provide space for the valid output array without the ghost cells. The result of the operator is then written into this output array. The function also accepts an optional parameter *args*, which is forwarded to *set\_ghost\_cells*.

## Parameters

- **operator** (*str*) Identifier for the operator. Some examples are 'laplace', 'gradient', or 'divergence'. The registered operators for this grid can be obtained from the *operators* attribute.
- **bc** (*str* or *list* or *tuple* or *dict*) The boundary conditions applied to the field. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by *{'value': NUM})* and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by *{'derivative': DERIV})* are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documentation*.
- \*\*kwargs Specifies extra arguments influencing how the operator is created.

#### Returns

the function that applies the operator. This function has the signature (arr: np.ndarray, out: np.ndarray = None, args=None).

#### **Return type**

callable

## make\_operator\_no\_bc(operator, \*\*kwargs)

return a compiled function applying an operator without boundary conditions

A function that takes the discretized full data as an input and an array of valid data points to which the result of applying the operator is written.

Note: The resulting function does not check whether the ghost cells of the input array have been supplied with sensible values. It is the responsibility of the user to set the values of the ghost cells beforehand. Use this function only if you absolutely know what you're doing. In all other cases, make\_operator() is probably the better choice.

#### **Parameters**

- operator (*str*) Identifier for the operator. Some examples are 'laplace', 'gradient', or 'divergence'. The registered operators for this grid can be obtained from the operators attribute.
- **\*\*kwargs** Specifies extra arguments influencing how the operator is created.

## Returns

the function that applies the operator. This function has the signature (arr: np.ndarray, out: np.ndarray), so they out array need to be supplied explicitly.

#### **Return type** callable

## **normalize\_point** (*point*, \*, *reflect=False*)

normalize grid coordinates by applying periodic boundary conditions

Here, points are assumed to be specified by the physical values along the non-symmetric axes of the grid, e.g., by grid coordinates. Normalizing points is useful to make sure they lie within the domain of the grid. This function respects periodic boundary conditions and can also reflect points off the boundary if *reflect = True*.

#### **Parameters**

- point (ndarray) Coordinates of a single point or an array of points, where the last axis denotes the point coordinates (e.g., a list of points).
- reflect (bool) Flag determining whether coordinates along non-periodic axes are reflected to lie in the valid range. If *False*, such coordinates are left unchanged and only periodic boundary conditions are enforced.

## Returns

The respective coordinates with periodic boundary conditions applied.

## **Return type**

ndarrav

## num\_axes: int

Number of axes that are not assumed symmetrically

Туре int

#### property num\_cells: int

the number of cells in this grid

Туре int

#### property numba\_type: str

represents type of the grid data in numba signatures

Type str

## operators: set[str] = {}

names of all operators defined for this grid

Туре

set

## property periodic: list[bool]

Flags that describe which axes are periodic

Type list

## plot()

visualize the grid

Return type None

## point\_from\_cartesian (points, \*, full=False)

convert points given in Cartesian coordinates to grid coordinates

## Parameters

- points (ndarray) Points given in Cartesian coordinates.
- full (bool) Indicates whether coordinates along symmetric axes are specified

#### Returns

Points given in the coordinates of the grid

## **Return type**

ndarray

## point\_to\_cartesian (points, \*, full=False)

convert coordinates of a point in grid coordinates to Cartesian coordinates

#### Parameters

- points (ndarray) The grid coordinates of the points
- full (bool) Indicates whether coordinates along symmetric axes are specified

## Returns

The Cartesian coordinates of the point

## **Return type**

## ndarray

classmethod register\_operator(name, factory\_func=None, rank\_in=0, rank\_out=0)

register an operator for this grid

## Example

The method can either be used directly:

GridClass.register\_operator("operator", make\_operator)

or as a decorator for the factory function:

#### **Parameters**

- **name** (*str*) The name of the operator to register
- **factory\_func** (*callable*) A function with signature (grid: GridBase, \*\*kwargs), which takes a grid object and optional keyword arguments and returns an implementation of the given operator. This implementation is a function that takes a ndarray of discretized values as arguments and returns the resulting discretized data in a ndarray after applying the operator.
- **rank\_in** (*int*) The rank of the input field for the operator
- **rank\_out** (*int*) The rank of the field that is returned by the operator

## property shape: tuple[int, ...]

the number of support points of each axis

Type

tuple of int

slice (indices)

return a subgrid of only the specified axes

## Parameters

indices (list) – Indices indicating the axes that are retained in the subgrid

Returns

The subgrid

Return type

GridBase

## abstract property state: dict[str, Any]

all information required for reconstructing the grid

## Туре

dict

## property state\_serialized: str

JSON-serialized version of the state of this grid

Туре

str

#### **transform** (*coordinates*, *source*, *target*, \*, *full=False*)

converts coordinates from one coordinate system to another

Supported coordinate systems include the following:

## • cartesian:

Cartesian coordinates where each point carries *dim* values. These are the true physical coordinates in space.

• grid:

Coordinates values in the coordinate system defined by the grid. A point is thus characterized by *grid.num\_axes* values.

• cell:

Normalized grid coordinates based on indexing the discretization cells. A point is characterized by *grid.num\_axes* values and the range of values for a given axis is between 0 and N, where N is the number of grid points. Consequently, the integral part of the cell coordinate denotes the cell, while the fractional part denotes the relative position within the cell. In particular, the cell center is located at i + 0.5 with i = 0, ..., N-1.

**Note:** Some conversion might involve projections if the coordinate system imposes symmetries. For instance, converting 3d Cartesian coordinates to grid coordinates in a spherically symmetric grid will only return the radius from the origin. Conversely, converting these grid coordinates back to 3d Cartesian coordinates will only return coordinates along a particular ray originating at the origin.

## Parameters

- coordinates (ndarray) The coordinates to convert
- **source** (*str*) The source coordinate system
- **target** (*str*) The target coordinate system
- full (bool) Indicates whether coordinates along symmetric axes are specified

## Returns

The transformed coordinates

#### **Return type**

ndarray

#### property typical\_discretization: float

the average side length of the cells

Type float

uniform\_cell\_volumes

returns True if all cell volumes are the same

## Туре

bool

property volume: float

total volume of the grid

## Туре

float

**class OperatorInfo** (*factory*, *rank\_in*, *rank\_out*, *name=''*)

Bases: NamedTuple

stores information about an operator

Create new instance of OperatorInfo(factory, rank\_in, rank\_out, name)

## Parameters

- factory (OperatorFactory) -
- rank\_in(int)-
- rank\_out (int) -
- name (str)-

## factory: OperatorFactory

Alias for field number 0

## name: str

Alias for field number 3

#### rank\_in: int

Alias for field number 1

#### rank\_out: int

Alias for field number 2

## exception PeriodicityError

Bases: RuntimeError

exception indicating that the grid periodicity is inconsistent

## discretize\_interval (x\_min, x\_max, num)

construct a list of equidistantly placed intervals

The discretization is defined as

$$x_{i} = x_{\min} + \left(i + \frac{1}{2}\right) \Delta x \quad \text{for} \quad i = 0, \dots, N-1$$
$$\Delta x = \frac{x_{\max} - x_{\min}}{N}$$

where N is the number of intervals given by *num*.

#### Parameters

- **x\_min** (*float*) Minimal value of the axis
- **x\_max** (float) Maximal value of the axis
- **num** (*int*) Number of intervals

## Returns

(midpoints, dx): the midpoints of the intervals and the used discretization dx.

## **Return type**

tuple

## registered\_operators()

returns all operators that are currently defined

#### Returns

a dictionary with the names of the operators defined for each grid class

#### **Return type**

dict

## 4.2.5 pde.grids.cartesian module

Cartesian grids of arbitrary dimension.

```
class CartesianGrid (bounds, shape, periodic=False)
```

Bases: GridBase

d-dimensional Cartesian grid with uniform discretization for each axis

The grids can be thought of as a collection of n-dimensional boxes, called cells, of equal length in each dimension. The bounds then defined the total volume covered by these cells, while the cell coordinates give the location of the box centers. We index the boxes starting from 0 along each dimension. Consequently, the cell  $i - \frac{1}{2}$  corresponds to the left edge of the covered interval and the index  $i + \frac{1}{2}$  corresponds to the right edge, when the dimension is covered by d boxes.

In particular, the discretization along dimension k is defined as

$$x_{i}^{(k)} = x_{\min}^{(k)} + \left(i + \frac{1}{2}\right) \Delta x^{(k)} \quad \text{for} \quad i = 0, \dots, N^{(k)} - 1$$
$$\Delta x^{(k)} = \frac{x_{\max}^{(k)} - x_{\min}^{(k)}}{N^{(k)}}$$

where  $N^{(k)}$  is the number of cells along this dimension. Consequently, cells have dimension  $\Delta x^{(k)}$  and cover the interval  $[x_{\min}^{(k)}, x_{\max}^{(k)}]$ .

## Parameters

- **bounds** (*list of tuple*) Give the coordinate range for each axis. This should be a tuple of two number (lower and upper bound) for each axis. The length of *bounds* thus determines the grid dimension.
- **shape** (*list*) The number of support points for each axis. The length of *shape* needs to match the grid dimension.
- **periodic** (*bool or list*) Specifies which axes possess periodic boundary conditions. This is either a list of booleans defining periodicity for each individual axis or a single boolean value specifying the same periodicity for all axes.

```
boundary_names: dict[str, tuple[int, bool]] = {'back': (2, False),
'bottom': (1, False), 'front': (2, True), 'left': (0, False), 'right':
(0, True), 'top': (1, True)}
```

Names of boundaries to select them conveniently

Type dict

## property cell\_volume\_data

size associated with each cell

cuboid: Cuboid

**difference\_vector** (*p1*, *p2*, \*, *coords='grid'*)

return Cartesian vector(s) pointing from p1 to p2

In case of periodic boundary conditions, the shortest vector is returned.

- p1 (ndarray) First point(s)
- p2 (ndarray) Second point(s)

• **coords** (*str*) – The coordinate system in which the points are specified. Valid values are *cartesian, cell,* and *grid*; see *transform()*.

## Returns

The difference vectors between the points with periodic boundary conditions applied.

#### **Return type**

ndarray

classmethod from\_bounds (bounds, shape, periodic)

## Parameters

- **bounds** (*tuple*) Give the coordinate range for each axis. This should be a tuple of two number (lower and upper bound) for each axis. The length of *bounds* thus determines the grid dimension.
- **shape** (*tuple*) The number of support points for each axis. The length of *shape* needs to match the grid dimension.
- **periodic** (*bool or list*) Specifies which axes possess periodic boundary conditions. This is either a list of booleans defining periodicity for each individual axis or a single boolean value specifying the same periodicity for all axes.

## Returns

representing the region chosen by bounds

#### **Return type**

CartesianGrid

## classmethod from\_state(state)

create a field from a stored state.

## Parameters

**state** (*dict*) – The state from which the grid is reconstructed.

## Returns

the grid re-created from the state data

## **Return type**

CartesianGrid

## get\_image\_data(data)

return a 2d-image of the data

## **Parameters**

data (ndarray) – The values at the grid points

## Returns

A dictionary with information about the data convenient for plotting.

### **Return type**

dict

## get\_line\_data(data, extract='auto')

return a line cut through the given data

- data (ndarray) The values at the grid points
- **extract** (*str*) Determines which cut is done through the grid. Possible choices are (default is *cut\_0*):

- *cut\_*#: return values along the axis specified by # and use the mid point along all other axes.
- project\_#: average values for all axes, except axis #.

Here, # can either be a zero-based index (from 0 to dim-1) or a letter denoting the axis.

#### Returns

A dictionary with information about the line cut, which is convenient for plotting.

#### **Return type**

dict

get\_random\_point (\*, boundary\_distance=0, coords='cartesian', rng=None)

return a random point within the grid

#### Parameters

- **boundary\_distance** (*float*) The minimal distance this point needs to have from all boundaries.
- **coords** (*str*) Determines the coordinate system in which the point is specified. Valid values are *cartesian*, *cell*, and *grid*; see *transform()*.
- rng (Generator) Random number generator (default: default\_rng())

## Returns

The coordinates of the point

Return type

ndarray

## get\_vector\_data (data, \*\*kwargs)

return data to visualize vector field

## **Parameters**

- data (ndarray) The vectorial values at the grid points
- **\*\*kwargs** Arguments forwarded to get\_image\_data().

#### Returns

A dictionary with information about the data convenient for plotting.

#### Return type

dict

## iter\_mirror\_points (point, with\_self=False, only\_periodic=True)

generates all mirror points corresponding to point

## Parameters

- point (ndarray) The point within the grid
- with\_self (bool) Whether to include the point itself
- only\_periodic (bool) Whether to only mirror along periodic axes

#### Returns

A generator yielding the coordinates that correspond to mirrors

#### **Return type**

Generator

plot (\*args, title=None, filename=None, action='auto', ax\_style=None, fig\_style=None, ax=None, \*\*kwargs)
visualize the grid

## **Parameters**

- title (*str*) Title of the plot. If omitted, the title might be chosen automatically.
- filename (*str*, *optional*) If given, the plot is written to the specified file.
- **action** (*str*) Decides what to do with the final figure. If the argument is set to *show*, matplotlib.pyplot.show() will be called to show the plot. If the value is *none*, the figure will be created, but not necessarily shown. The value *close* closes the figure, after saving it to a file when *filename* is given. The default value *auto* implies that the plot is shown if it is not a nested plot call.
- **ax\_style** (*dict*) Dictionary with properties that will be changed on the axis after the plot has been drawn by calling matplotlib.pyplot.setp(). A special item i this dictionary is *use\_offset*, which is flag that can be used to control whether offset are shown along the axes of the plot.
- **fig\_style** (*dict*) Dictionary with properties that will be changed on the figure after the plot has been drawn by calling matplotlib.pyplot.setp(). For instance, using fig\_style={'dpi': 200} increases the resolution of the figure.
- **ax** (matplotlib.axes.Axes) Figure axes to be used for plotting. The special value "create" creates a new figure, while "reuse" attempts to reuse an existing figure, which is the default.
- **\*\*kwargs** Extra arguments are passed on the to the matplotlib plotting routines, e.g., to set the color of the lines

## slice (indices)

return a subgrid of only the specified axes

## Parameters

indices (list) – Indices indicating the axes that are retained in the subgrid

Returns The subgrid

8

Return type CartesianGrid

property state: dict[str, Any]

the state of the grid

## Type

dict

property volume: float

total volume of the grid

Type float

class UnitGrid (shape, periodic=False)

Bases: CartesianGrid

d-dimensional Cartesian grid with unit discretization in all directions

The grids can be thought of as a collection of d-dimensional cells of unit length. The *shape* parameter determines how many boxes there are in each direction. The cells are enumerated starting with 0, so the last cell has index

n-1 if there are *n* cells along a dimension. A given cell *i* extends from coordinates *i* to i+1, so the midpoint is at  $i+\frac{1}{2}$ , which is the cell coordinate. Taken together, the cells covers the interval [0, n] along this dimension.

## **Parameters**

- **shape** (*list*) The number of support points for each axis. The dimension of the grid is given by *len(shape)*.
- **periodic** (*bool or list*) Specifies which axes possess periodic boundary conditions. This is either a list of booleans defining periodicity for each individual axis or a single boolean value specifying the same periodicity for all axes.

## classmethod from\_state(state)

create a field from a stored state.

```
Parameters
state (dict) – The state from which the grid is reconstructed.
```

Return type UnitGrid

slice (indices)

return a subgrid of only the specified axes

## Parameters

indices (list) - Indices indicating the axes that are retained in the subgrid

## Returns

The subgrid

## Return type

UnitGrid

## property state: dict[str, Any]

the state of the grid

#### Type dict

to\_cartesian()

convert unit grid to CartesianGrid

#### Returns

The equivalent cartesian grid

## Return type

CartesianGrid

## 4.2.6 pde.grids.cylindrical module

Cylindrical grids with azimuthal symmetry

**class** CylindricalSymGrid (*radius*, *bounds\_z*, *shape*, *periodic\_z=False*)

Bases: GridBase

3-dimensional cylindrical grid assuming polar symmetry

The polar symmetry implies that states only depend on the radial and axial coordinates r and z, respectively. These are discretized uniformly as

$$r_{i} = R_{\text{inner}} + \left(i + \frac{1}{2}\right)\Delta r \qquad \text{for} \quad i = 0, \dots, N_{r} - 1 \qquad \text{with} \quad \Delta r = \frac{R_{\text{outer}} - R_{\text{inner}}}{N_{r}}$$
$$z_{j} = z_{\min} + \left(j + \frac{1}{2}\right)\Delta z \qquad \text{for} \quad j = 0, \dots, N_{z} - 1 \qquad \text{with} \quad \Delta z = \frac{z_{\max} - z_{\min}}{N_{z}}$$

where  $R_{\text{outer}}$  is the outer radius of the grid,  $R_{\text{inner}}$  corresponds to a possible inner radius (which is zero by default), and  $z_{\min}$  and  $z_{\max}$  denote the respective lower and upper bounds of the axial direction, so that  $z_{\max} - z_{\min}$  is the total height. The two axes are discretized by  $N_r$  and  $N_z$  support points, respectively.

**Warning:** The order of components in the vector and tensor fields defined on this grid are still  $(r, \phi, z)$ . To avoid any confusion it might thus be best to access components by name instead of index, e.g., use field['z'] instead of field[2].

### Parameters

- **radius** (float or tuple of floats) radius  $R_{outer}$  in case a simple float is given. If a tuple is supplied it is interpreted as the inner and outer radius,  $(R_{inner}, R_{outer})$ .
- **bounds\_z** (*tuple*) The lower and upper bound of the z-axis
- **shape** (*tuple*) The number of support points in r and z direction, respectively. The same number is used for both if a single value is given.
- **periodic\_z** (bool) Determines whether the z-axis has periodic boundary conditions.

```
boundary_names: dict[str, tuple[int, bool]] = {'bottom': (1, False),
'inner': (0, False), 'outer': (0, True), 'top': (1, True)}
```

Names of boundaries to select them conveniently

## Type

dict

```
c: CoordinatesBase = CylindricalCoordinates()
```

Coordinates of the grid

## Туре

CoordinatesBase

```
cell_volume_data: Sequence[FloatNumerical] | None
```

Information about the size of discretization cells

## Туре

list

```
coordinate_constraints: list[int] = [0, 1]
```

axes that not described explicitly

Type list

```
difference_vector (p1, p2, *, coords='grid')
```

return Cartesian vector(s) pointing from p1 to p2

In case of periodic boundary conditions, the shortest vector is returned.

- p1 (ndarray) First point(s)
- p2 (ndarray) Second point(s)
- **coords** (*str*) The coordinate system in which the points are specified. Valid values are *cartesian, cell,* and *grid*; see *transform()*.

## Returns

The difference vectors between the points with periodic boundary conditions applied.

### **Return type**

ndarray

classmethod from\_bounds (bounds, shape, periodic)

## **Parameters**

- **bounds** (*tuple*) Give the coordinate range for each axis. This should be a tuple of two number (lower and upper bound) for each axis. The length of *bounds* must be 2.
- **shape** (*tuple*) The number of support points for each axis. The length of *shape* needs to be 2.
- **periodic** (*bool* or *list*) Specifies which axes possess periodic boundary conditions. The first entry is ignored.

## Returns

grid representing the region chosen by bounds

Return type

CylindricalSymGrid

## classmethod from\_state(state)

create a field from a stored state.

#### Parameters

state (dict) – The state from which the grid is reconstructed.

## Return type

CylindricalSymGrid

## get\_cartesian\_grid (mode='valid')

return a Cartesian grid for this Cylindrical one

#### **Parameters**

**mode** (str) – Determines how the grid is determined. Setting it to 'valid' only returns points that are fully resolved in the cylindrical grid, e.g., the cylinder is circumscribed. Conversely, 'full' returns all data, so the cylinder is inscribed.

## Returns

The requested grid

## **Return type**

pde.grids.cartesian.CartesianGrid

## get\_image\_data(data)

return a 2d-image of the data

## Parameters

data (ndarray) - The values at the grid points

## Returns

A dictionary with information about the image, which is convenient for plotting.

## **Return type**

dict

## get\_line\_data(data, extract='auto')

return a line cut for the cylindrical grid

## Parameters

- data (ndarray) The values at the grid points
- **extract** (*str*) Determines which cut is done through the grid. Possible choices are (default is *cut\_axial*):
  - $cut_z$  or  $cut_axial$ : values along the axial coordinate for r = 0.
  - *project\_z* or *project\_axial*: average values for each axial position (radial average).
  - project\_r or project\_radial: average values for each radial position (axial average)

#### Returns

A dictionary with information about the line cut, which is convenient for plotting.

## Return type

dict

get\_random\_point (\*, boundary\_distance=0, avoid\_center=False, coords='cartesian', rng=None)
return a random point within the grid

#### **Parameters**

- **boundary\_distance** (*float*) The minimal distance this point needs to have from all boundaries.
- **avoid\_center** (*bool*) Determines whether the boundary distance should also be kept from the center, i.e., whether points close to the center are returned.
- **coords** (*str*) Determines the coordinate system in which the point is specified. Valid values are *cartesian*, *cell*, and *grid*; see *transform()*.
- rng (Generator) Random number generator (default: default\_rng())

## Returns

The coordinates of the point

#### **Return type**

ndarray

## property has\_hole: bool

whether the inner radius is larger than zero

## Туре

bool

iter\_mirror\_points (point, with\_self=False, only\_periodic=True)

generates all mirror points corresponding to point

#### **Parameters**

- point (ndarray) The point within the grid
- with\_self (bool) Whether to include the point itself
- only\_periodic (bool) Whether to only mirror along periodic axes

## Returns

A generator yielding the coordinates that correspond to mirrors

Return type

Generator

property length: float

length of the cylinder

Туре

float

property radius: float | tuple[float, float]

radius of the sphere

Type float

slice (indices)

return a subgrid of only the specified axes

Parameters

indices (list) – Indices indicating the axes that are retained in the subgrid

Returns

CartesianGrid or PolarSymGrid: The subgrid

Return type CartesianGrid | PolarSymGrid

property state: dict[str, Any]

all information required for reconstructing the grid

Туре

dict

```
property volume: float
```

total volume of the grid

Туре

float

## 4.2.7 pde.grids.spherical module

Spherically-symmetric grids in 2 and 3 dimensions. These are grids that only discretize the radial direction, assuming symmetry with respect to all angles. This choice implies that differential operators might not be applicable to all fields. For instance, the divergence of a vector field on a spherical grid can only be represented as a scalar field on the same grid if the  $\theta$ -component of the vector field vanishes.

class PolarSymGrid(radius, shape)

Bases: SphericalSymGridBase

2-dimensional polar grid assuming angular symmetry

The angular symmetry implies that states only depend on the radial coordinate r, which is discretized uniformly as

$$r_i = R_{\text{inner}} + \left(i + \frac{1}{2}\right)\Delta r$$
 for  $i = 0, \dots, N-1$  with  $\Delta r = \frac{R_{\text{outer}} - R_{\text{inner}}}{N}$ 

where  $R_{\text{outer}}$  is the outer radius of the grid and  $R_{\text{inner}}$  corresponds to a possible inner radius, which is zero by default. The radial direction is discretized by N support points.

- radius (float or tuple of floats) Radius  $R_{\text{outer}}$  in case a simple float is given. If a tuple is supplied it is interpreted as the inner and outer radius,  $(R_{\text{inner}}, R_{\text{outer}})$ .
- **shape** (*tuple or int*) The number N of support points along the radial coordinate.

## c: *CoordinatesBase* = PolarCoordinates()

Coordinates of the grid

Type

CoordinatesBase

```
coordinate_constraints: list[int] = [0, 1]
```

axes that not described explicitly

#### Type list

class SphericalSymGrid(radius, shape)

Bases: SphericalSymGridBase

3-dimensional spherical grid assuming spherical symmetry

The symmetry implies that states only depend on the radial coordinate r, which is discretized as follows:

$$r_i = R_{\text{inner}} + \left(i + \frac{1}{2}\right)\Delta r$$
 for  $i = 0, \dots, N-1$  with  $\Delta r = \frac{R_{\text{outer}} - R_{\text{inner}}}{N}$ 

where  $R_{\text{outer}}$  is the outer radius of the grid and  $R_{\text{inner}}$  corresponds to a possible inner radius, which is zero by default. The radial direction is discretized by N support points.

**Warning:** Not all results of differential operators on vectorial and tensorial fields can be expressed in terms of fields that only depend on the radial coordinate r. In particular, the gradient of a vector field can only be calculated if the azimuthal component of the vector field vanishes. Similarly, the divergence of a tensor field can only be taken in special situations.

#### **Parameters**

- **radius** (float or tuple of floats) Radius  $R_{\text{outer}}$  in case a simple float is given. If a tuple is supplied it is interpreted as the inner and outer radius,  $(R_{\text{inner}}, R_{\text{outer}})$ .
- **shape** (*tuple or int*) The number N of support points along the radial coordinate.

#### c: CoordinatesBase = SphericalCoordinates()

Coordinates of the grid

## Туре

```
CoordinatesBase
```

## coordinate\_constraints: list[int] = [0, 1, 2]

axes that not described explicitly

Type list

#### class SphericalSymGridBase(radius, shape)

Bases: GridBase

Base class for d-dimensional spherical grids with angular symmetry

The angular symmetry implies that states only depend on the radial coordinate r, which is discretized uniformly as

$$r_i = R_{\text{inner}} + \left(i + \frac{1}{2}\right)\Delta r$$
 for  $i = 0, \dots, N-1$  with  $\Delta r = \frac{R_{\text{outer}} - R_{\text{inner}}}{N}$ 

where  $R_{\text{outer}}$  is the outer radius of the grid and  $R_{\text{inner}}$  corresponds to a possible inner radius, which is zero by default. The radial direction is discretized by N support points.

#### **Parameters**

- radius (float or tuple of floats) Radius  $R_{\text{outer}}$  in case a simple float is given. If a tuple is supplied it is interpreted as the inner and outer radius,  $(R_{\text{inner}}, R_{\text{outer}})$ .
- **shape** (tuple or int) The number N of support points along the radial coordinate.

boundary\_names: dict[str, tuple[int, bool]] = {'inner': (0, False),
'outer': (0, True)}

Names of boundaries to select them conveniently

## Туре

dict

#### cell\_volume\_data: Sequence[FloatNumerical] | None

Information about the size of discretization cells

## Туре

list

classmethod from\_bounds (bounds, shape, periodic)

#### **Parameters**

- **bounds** (*tuple*) Give the coordinate range for the radial axis.
- **shape** (*tuple*) The number of support points for the radial axis
- **periodic**(tuple[bool]) -

#### Returns

represents the region chosen by bounds

#### **Return type**

SphericalGridBase

## classmethod from\_state(state)

create a field from a stored state.

#### Parameters

**state** (*dict*) – The state from which the grid is reconstructed.

## **Return type**

## SphericalSymGridBase

#### get\_cartesian\_grid (mode='valid', num=None)

return a Cartesian grid for this spherical one

- mode (*str*) Determines how the grid is determined. Setting it to 'valid' (or 'inscribed') only returns points that are fully resolved in the spherical grid, e.g., the Cartesian grid is inscribed in the sphere. Conversely, 'full' (or 'circumscribed') returns all data, so the Cartesian grid is circumscribed.
- **num** (*int*) Number of support points along each axis of the returned grid.

## Returns

The requested grid

## **Return type**

pde.grids.cartesian.CartesianGrid

get\_image\_data (data, \*, performance\_goal='speed', fill\_value=0, masked=True)

return a 2d-image of the data

## Parameters

- data (ndarray) The values at the grid points
- **performance\_goal** (*str*) Determines the method chosen for interpolation. Possible options are *speed* and *quality*.
- **fill\_value** (*float*) The value assigned to invalid positions (those inside the hole or outside the region).
- masked (*bool*) Whether a numpy.ma.MaskedArray is returned for the data instead of the normal ndarray.

## Returns

dict: A dictionary with information about the image, which is convenient for plotting.

## Return type

dict[str, Any]

get\_line\_data(data, extract='auto')

return a line cut along the radial axis

## **Parameters**

- data (ndarray) The values at the grid points
- **extract** (*str*) Determines which cut is done through the grid. This parameter is mainly supplied for a consistent interface and has no effect for polar grids.

## Returns

A dictionary with information about the line cut, which is convenient for plotting.

## Return type

dict[str, Any]

get\_random\_point (\*, boundary\_distance=0, avoid\_center=False, coords='cartesian', rng=None)

return a random point within the grid

Note that these points will be uniformly distributed in the volume, implying they are not uniformly distributed on the radial axis.

## Parameters

- **boundary\_distance** (*float*) The minimal distance this point needs to have from all boundaries.
- **avoid\_center** (*bool*) Determines whether the boundary distance should also be kept from the center, i.e., whether points close to the center are returned.
- **coords** (*str*) Determines the coordinate system in which the point is specified. Valid values are *cartesian*, *cell*, and *grid*; see *transform()*.
- **rng** (Generator) Random number generator (default: default\_rng())

## Returns

The coordinates of the point

#### **Return type**

ndarray

## property has\_hole: bool

returns whether the inner radius is larger than zero

plot (\*args, title=None, filename=None, action='auto', ax\_style=None, fig\_style=None, ax=None, \*\*kwargs)
visualize the spherically symmetric grid in two dimensions

#### Parameters

- **title** (*str*) Title of the plot. If omitted, the title might be chosen automatically.
- filename (str, optional) If given, the plot is written to the specified file.
- **action** (*str*) Decides what to do with the final figure. If the argument is set to *show*, matplotlib.pyplot.show() will be called to show the plot. If the value is *none*, the figure will be created, but not necessarily shown. The value *close* closes the figure, after saving it to a file when *filename* is given. The default value *auto* implies that the plot is shown if it is not a nested plot call.
- **ax\_style** (*dict*) Dictionary with properties that will be changed on the axis after the plot has been drawn by calling matplotlib.pyplot.setp(). A special item i this dictionary is *use\_offset*, which is flag that can be used to control whether offset are shown along the axes of the plot.
- **fig\_style** (*dict*) Dictionary with properties that will be changed on the figure after the plot has been drawn by calling matplotlib.pyplot.setp(). For instance, using fig\_style={'dpi': 200} increases the resolution of the figure.
- **ax** (matplotlib.axes.Axes) Figure axes to be used for plotting. The special value "create" creates a new figure, while "reuse" attempts to reuse an existing figure, which is the default.
- **\*\*kwargs** Extra arguments are passed on the to the matplotlib plotting routines, e.g., to set the color of the lines

## property radius: float | tuple[float, float]

radius of the sphere

Type float

property state: dict[str, Any]

the state of the grid

## Туре

state

property volume: float

total volume of the grid

Туре

float

volume\_from\_radius (radius, dim)

Return the volume of a sphere with a given radius

- radius (float or ndarray) Radius of the sphere
- **dim** (*int*) Dimension of the space

Returns Volume of the sphere

```
Return type
float or ndarray
```

## 4.3 pde.pdes package

Package that defines PDEs describing physical systems.

The examples in this package are often simple version of classical PDEs to demonstrate various aspects of the *py-pde* package. Clearly, not all extensions to these PDEs can be covered here, but this should serve as a starting point for custom investigations.

Publicly available methods should take fields with grid information and also only return such methods. There might be corresponding private methods that deal with raw data for faster simulations.

PDE	PDE defined by mathematical expressions
AllenCahnPDE	A simple Allen-Cahn equation
CahnHilliardPDE	A simple Cahn-Hilliard equation
DiffusionPDE	A simple diffusion equation
KPZInterfacePDE	The Kardar–Parisi–Zhang (KPZ) equation
KuramotoSivashinskyPDE	The Kuramoto-Sivashinsky equation
SwiftHohenbergPDE	The Swift-Hohenberg equation
WavePDE	A simple wave equation

Additionally, we offer two solvers for typical elliptical PDEs:

solve_laplace_equation	Solve Laplace's equation on a given grid.
solve_poisson_equation	Solve Laplace's equation on a given grid

## 4.3.1 pde.pdes.allen\_cahn module

A Allen-Cahn equation

**class AllenCahnPDE** (*interface\_width=1*, *bc='auto\_periodic\_neumann'*)

Bases: PDEBase

A simple Allen-Cahn equation

The mathematical definition is

$$\partial_t c = \gamma \nabla^2 c - c^3 + c$$

where c is a scalar field and  $\gamma$  sets the (squared) interfacial width.

- interface\_width (float) The diffusivity of the described species
- **bc** (*BoundariesData*) The boundary conditions applied to the field. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end

(using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by *['value': NUM]*) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by *['derivative': DERIV]*) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documenta-tion*.

## evolution\_rate(state, t=0)

evaluate the right hand side of the PDE

#### Parameters

- state (ScalarField) The scalar field describing the concentration distribution
- t (float) The current time point

## Returns

Scalar field describing the evolution rate of the PDE

#### **Return type**

ScalarField

## explicit\_time\_dependence: bool | None = False

Flag indicating whether the right hand side of the PDE has an explicit time dependence.

## Туре

bool

## property expression: str

the expression of the right hand side of this PDE

Туре

```
str
interface_width: float
```

## 4.3.2 pde.pdes.base module

Base class for defining partial differential equations

class PDEBase (\*, noise=0, rng=None)

Bases: object

base class for defining partial differential equations (PDEs)

Custom PDEs can be implemented by subclassing *PDEBase* to specify the evolution rate. In the simple case of deterministic PDEs, the methods *PDEBase.evolution\_rate()* and *PDEBase.\_make\_pde\_rhs\_numba()* need to be overwritten for supporting the *numpy* and *numba* backend, respectively.

- **noise** (float or ndarray) Variance of the additive Gaussian white noise that is supported for all PDEs by default. If set to zero, a deterministic partial differential equation will be solved. Different noise magnitudes can be supplied for each field in coupled PDEs.
- **rng** (Generator) Random number generator (default: default\_rng()) used for stochastic simulations. Note that this random number generator is only used for numpy function, while compiled numba code uses the random number generator of numba. Moreover, in simulations using multiprocessing, setting the same generator in all processes might yield unintended correlations in the simulation results.

**Note:** If more complicated noise structures are required, the methods *PDEBase.noise\_realization()* and *PDEBase.\_make\_noise\_realization\_numba()* need to be overwritten for the *numpy* and *numba* backend, respectively.

#### cache\_rhs: bool = False

Flag indicating whether the right hand side of the equation should be cached. If True, the same implementation is used in subsequent calls to *solve*. Note that the cache is only invalidated when the grid of the underlying state changes. Consequently, the simulation might lead to wrong results if the parameters of the PDE are changed after the first call. This option is thus disabled by default and should be used with care.

Туре

bool

#### check\_implementation: bool = True

Flag determining whether numba-compiled functions should be checked against their numpy counter-parts. This can help with implementing a correct compiled version for a PDE class.

Туре

bool

**check\_rhs\_consistency** (*state*, *t=0*, \*, *tol=1e-07*, *rhs\_numba=None*, \*\*kwargs)

check the numba compiled right hand side versus the numpy variant

## **Parameters**

- **state** (FieldBase) The state for which the evolution rates should be compared
- t (float) The associated time point
- **tol** (*float*) Acceptance tolerance. The check passes if the evolution rates differ by less then this value
- **rhs\_numba** (*callable*) The implementation of the numba variant that is to be checked. If omitted, an implementation is obtained by calling PDEBase. \_make\_pde\_rhs\_numba\_cached().

Return type None

#### complex\_valued: bool = False

Flag indicating whether the right hand side is a complex-valued PDE, which requires all involved variables to have complex data type.

#### Туре

bool

## diagnostics: dict[str, Any]

Diagnostic information (available after the PDE has been solved)

Туре

dict

## abstract evolution\_rate(state, t=0)

evaluate the right hand side of the PDE

- **state** (*FieldBase*) The field at the current time point
- t (float) The current time point

#### Returns

Field describing the evolution rate of the PDE

## **Return type**

FieldBase

#### explicit\_time\_dependence: bool | None = None

Flag indicating whether the right hand side of the PDE has an explicit time dependence.

Type bool

#### property is\_sde: bool

flag indicating whether this is a stochastic differential equation

The BasePDF class supports additive Gaussian white noise, whose magnitude is controlled by the *noise* property. In this case, *is\_sde* is *True* if *self.noise* != 0.

## Туре

bool

## make\_modify\_after\_step(state)

returns a function that can be called to modify a state

This function is applied to the state after each integration step when an explicit stepper is used. The default behavior is to not change the state.

#### **Parameters**

**state** (FieldBase) – An example for the state from which the grid and other information can be extracted

#### Returns

Function that can be applied to a state to modify it and which returns a measure for the corrections applied to the state

#### **Return type**

Callable[[ndarray], float]

## make\_pde\_rhs (state, backend='auto', \*\*kwargs)

return a function for evaluating the right hand side of the PDE

#### Parameters

- **state** (FieldBase) An example for the state from which the grid and other information can be extracted.
- **backend** (*str*) Determines how the function is created. Accepted values are 'numpy' and 'numba'. Alternatively, 'auto' lets the code pick the optimal backend.

#### Returns

Function determining the right hand side of the PDE

# Return type callable

make\_sde\_rhs (state, backend='auto', \*\*kwargs)

return a function for evaluating the right hand side of the SDE

#### Parameters

• state (FieldBase) - An example for the state from which information can be extracted

• **backend** (*str*) – Determines how the function is created. Accepted values are 'numpy' and 'numba'. Alternatively, 'auto' lets the code pick the optimal backend.

## Returns

Function determining the deterministic part of the right hand side of the PDE together with a noise realization.

## **Return type**

Callable[[ndarray, float], tuple[ndarray, ndarray]]

**noise\_realization** (*state*, *t*=0, \*, *label='Noise realization'*)

returns a realization for the noise

## Parameters

- state (ScalarField) The scalar field describing the concentration distribution
- t (float) The current time point
- **label** (*str*) The label for the returned field

## Returns

Scalar field describing the evolution rate of the PDE

#### Return type

ScalarField

**solve** (*state*, *t\_range*, *dt=None*, *tracker='auto'*, \*, *solver='explicit'*, *ret\_info=False*, \*\**kwargs*)

solves the partial differential equation

The method constructs a suitable solver (*SolverBase*) and controller (Controller) to advance the state over the temporal range specified by *t\_range*. This method only exposes the most common functions, so explicit construction of these classes might offer more flexibility.

- **state** (*FieldBase*) The initial state (which also defines the spatial grid).
- **t\_range** (*float* or *tuple*) Sets the time range for which the PDE is solved. This should typically be a tuple of two numbers, (*t\_start, t\_end*), specifying the initial and final time of the simulation. If only a single value is given, it is interpreted as *t\_end* and the time range is (0, *t\_end*).
- dt (float) Time step of the chosen stepping scheme. If *None*, a default value based on the stepper will be chosen. If an adaptive stepper is used (supported by *ScipySolver* and *ExplicitSolver*), *dt* sets the initial time step.
- **tracker** (*TrackerCollectionDataType*) Defines trackers that process the state of the simulation at specified times. A tracker is either an instance of *Tracker-Base* or a string identifying a tracker (possible identifiers can be obtained by calling *get\_named\_trackers()*). Multiple trackers can be specified as a list. The default value *auto* checks the state for consistency (tracker 'consistency') and displays a progress bar (tracker 'progress') when tqdm is installed. More general trackers are defined in *trackers*, where all options are explained in detail. In particular, the time points where the tracker analyzes data can be chosen when creating a tracker object explicitly.
- **solver** (*SolverBase* or str) Specifies the method for solving the differential equation. This can either be an instance of *SolverBase* or a descriptive name like 'explicit' or 'scipy'. The valid names are given by *pde.solvers.registered\_solvers()*. Details of the solvers and additional features (like adaptive time steps) are explained in *solvers*.

- **ret\_info** (*bool*) Flag determining whether diagnostic information about the solver process should be returned. Note that the same information is also available as the *diag-nostics* attribute.
- **\*\*kwargs** Additional keyword arguments are forwarded to the solver class chosen with the *solver* argument. In particular, *ExplicitSolver* supports several *schemes* and an adaptive stepper can be enabled using adaptive=True. Conversely, *ScipySolver* accepts the additional arguments of scipy.integrate.solve\_ivp().

### Returns

The state at the final time point. If  $ret_info == True$ , a tuple with the final state and a dictionary with additional information is returned. Note that *None* instead of a field is returned in multiprocessing simulations if the current node is not the main MPI node.

# Return type

FieldBase

# expr\_prod (factor, expression)

helper function for building an expression with an (optional) pre-factor

# Parameters

- **factor** (*float*) The value of the prefactor
- **expression** (*str*) The remaining expression

### Returns

The expression with the factor appended if necessary

### **Return type**

str

# 4.3.3 pde.pdes.cahn\_hilliard module

### A Cahn-Hilliard equation

**class** CahnHilliardPDE (*interface\_width=1*, *bc\_c='auto\_periodic\_neumann'*, *bc\_mu='auto\_periodic\_neumann'*)

Bases: PDEBase

A simple Cahn-Hilliard equation

The mathematical definition is

$$\partial_t c = \nabla^2 \left( c^3 - c - \gamma \nabla^2 c \right)$$

where c is a scalar field taking values on the interval [-1, 1] and  $\gamma$  sets the (squared) interfacial width.

- **interface\_width** (*float*) The width of the interface between the separated phases. This defines a characteristic length in the simulation. The grid needs to resolve this length of a stable simulation.
- bc\_c (BoundariesData) The boundary conditions applied to the field. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by *['value': NUM]*) and Neumann conditions enforcing the value DERIV for the

derivative in the normal direction (specified by *{'derivative': DERIV}*) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documenta-tion*.

• **bc\_mu** (*BoundariesData*) – The boundary conditions applied to the chemical potential associated with the scalar field c. Supports the same options as  $bc_c$ .

#### evolution\_rate(state, t=0)

evaluate the right hand side of the PDE

#### Parameters

- state (ScalarField) The scalar field describing the concentration distribution
- t (float) The current time point

#### Returns

Scalar field describing the evolution rate of the PDE

#### **Return type**

ScalarField

#### explicit\_time\_dependence: bool | None = False

Flag indicating whether the right hand side of the PDE has an explicit time dependence.

# Туре

bool

#### property expression: str

the expression of the right hand side of this PDE

Type str

# 4.3.4 pde.pdes.diffusion module

#### A simple diffusion equation

**class DiffusionPDE** (*diffusivity=1*, \*, *bc='auto\_periodic\_neumann'*, *noise=0*, *rng=None*)

Bases: PDEBase

A simple diffusion equation

The mathematical definition is

$$\partial_t c = D\nabla^2 c$$

where c is a scalar field and D denotes the diffusivity.

- **diffusivity** (*float*) The diffusivity of the described species
- bc (BoundariesData) The boundary conditions applied to the field. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by *{'value': NUM}*) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by *{'derivative': DERIV}*) are supported. Note

that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documenta-tion*.

- **noise** (*float*) Variance of the (additive) noise term
- **rng** (Generator) Random number generator (default: default\_rng()) used for stochastic simulations. Note that this random number generator is only used for numpy function, while compiled numba code uses the random number generator of numba. Moreover, in simulations using multiprocessing, setting the same generator in all processes might yield unintended correlations in the simulation results.

# evolution\_rate(state, t=0)

evaluate the right hand side of the PDE

#### Parameters

- state (ScalarField) The scalar field describing the concentration distribution
- t (float) The current time point

#### Returns

Scalar field describing the evolution rate of the PDE

# **Return type**

ScalarField

#### explicit\_time\_dependence: bool | None = False

Flag indicating whether the right hand side of the PDE has an explicit time dependence.

#### Туре

bool

#### property expression: str

the expression of the right hand side of this PDE

Type str

# 4.3.5 pde.pdes.kpz\_interface module

The Kardar-Parisi-Zhang (KPZ) equation describing the evolution of an interface

class KPZInterfacePDE (nu=0.5, lmbda=1, \*, bc='auto\_periodic\_neumann', noise=0, rng=None)

Bases: PDEBase

The Kardar-Parisi-Zhang (KPZ) equation

The mathematical definition is

$$\partial_t h = \nu \nabla^2 h + \frac{\lambda}{2} \left( \nabla h \right)^2 + \eta(\mathbf{r}, t)$$

where h is the height of the interface in Monge parameterization. The dynamics are governed by the two parameters  $\nu$  and  $\lambda$ , while  $\eta$  is Gaussian white noise, whose strength is controlled by the *noise* argument.

- **nu** (float) Parameter  $\nu$  for the strength of the diffusive term
- **Imbda** (float) Parameter  $\lambda$  for the strenth of the gradient term

- **bc** (BoundariesData) The boundary conditions applied to the field. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by *{'value': NUM}*) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by *{'derivative': DERIV}*) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documentation*.
- **noise** (*float*) Variance of the (additive) noise term
- **rng** (Generator) Random number generator (default: default\_rng()) used for stochastic simulations. Note that this random number generator is only used for numpy function, while compiled numba code uses the random number generator of numba. Moreover, in simulations using multiprocessing, setting the same generator in all processes might yield unintended correlations in the simulation results.

#### evolution\_rate(state, t=0)

evaluate the right hand side of the PDE

#### **Parameters**

- **state** (ScalarField) The scalar field describing the concentration distribution
- t (float) The current time point

#### Returns

Scalar field describing the evolution rate of the PDE

#### **Return type**

ScalarField

#### explicit\_time\_dependence: bool | None = False

Flag indicating whether the right hand side of the PDE has an explicit time dependence.

#### Туре

bool

# property expression: str

the expression of the right hand side of this PDE

Туре

str

# 4.3.6 pde.pdes.kuramoto\_sivashinsky module

The Kardar-Parisi-Zhang (KPZ) equation describing the evolution of an interface

class KuramotoSivashinskyPDE (nu=1, \*, bc='auto\_periodic\_neumann', bc\_lap=None, noise=0, rng=None)

Bases: PDEBase

The Kuramoto-Sivashinsky equation

The mathematical definition is

$$\partial_t u = -\nu \nabla^4 u - \nabla^2 u - \frac{1}{2} \left( \nabla h \right)^2 + \eta(\boldsymbol{r}, t)$$

where u is the height of the interface in Monge parameterization. The dynamics are governed by the parameters  $\nu$ , while  $\eta$  is Gaussian white noise, whose strength is controlled by the *noise* argument.

#### Parameters

- **nu** (float) Parameter  $\nu$  for the strength of the fourth-order term
- **bc** (BoundariesData) The boundary conditions applied to the field. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by *{'value': NUM}*) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by *{'derivative': DERIV}*) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documentation*.
- **bc\_lap** (*BoundariesData* / *None*) The boundary conditions applied to the second derivative of the scalar field c. If *None*, the same boundary condition as *bc* is chosen. Otherwise, this supports the same options as *bc*.
- **noise** (*float*) Variance of the (additive) noise term
- **rng** (Generator) Random number generator (default: default\_rng()) used for stochastic simulations. Note that this random number generator is only used for numpy function, while compiled numba code uses the random number generator of numba. Moreover, in simulations using multiprocessing, setting the same generator in all processes might yield unintended correlations in the simulation results.

# evolution\_rate(state, t=0)

evaluate the right hand side of the PDE

### Parameters

- **state** (ScalarField) The scalar field describing the concentration distribution
- t (float) The current time point

#### Returns

Scalar field describing the evolution rate of the PDE

#### **Return type**

ScalarField

## explicit\_time\_dependence: bool | None = False

Flag indicating whether the right hand side of the PDE has an explicit time dependence.

# Туре

bool

# property expression: str

the expression of the right hand side of this PDE

Туре

str

# 4.3.7 pde.pdes.laplace module

Solvers for Poisson's and Laplace's equation

#### **solve\_laplace\_equation** (grid, bc, label="Solution to Laplace's equation")

Solve Laplace's equation on a given grid.

This is implemented by calling *solve\_poisson\_equation()* with a vanishing right hand side.

#### **Parameters**

- grid (GridBase) The grid on which the equation is solved
- **bc** (BoundariesData) The boundary conditions applied to the field. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by {value': NUM}) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by {'derivative': DERIV}) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documentation*.
- **label** (*str*) The label of the returned field.

#### Returns

The field that solves the equation. This field will be defined on the given grid.

#### **Return type**

ScalarField

solve\_poisson\_equation (rhs, bc, label="Solution to Poisson's equation", \*\*kwargs)

Solve Laplace's equation on a given grid

Denoting the current field by u, we thus solve for f, defined by the equation

$$\nabla^2 u(\boldsymbol{r}) = -f(\boldsymbol{r})$$

with boundary conditions specified by bc.

Note: In case of periodic or Neumann boundary conditions, the right hand side f(r) needs to satisfy the following condition

$$\int f \,\mathrm{d}V = \oint g \,\mathrm{d}S \;,$$

where g denotes the function specifying the outwards derivative for Neumann conditions. Note that for periodic boundaries g vanishes, so that this condition implies that the integral over f must vanish for neutral Neumann or periodic conditions.

- **rhs** (*ScalarField*) The scalar field *f* describing the right hand side
- bc (BoundariesData) The boundary conditions applied to the field. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end

(using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by *['value': NUM]*) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by *['derivative': DERIV]*) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documenta-tion*.

• **label** (*str*) – The label of the returned field.

#### Returns

The field u that solves the equation. This field will be defined on the same grid as *rhs*.

#### **Return type**

ScalarField

# 4.3.8 pde.pdes.pde module

Defines a PDE class whose right hand side is given as a string

Bases: PDEBase

PDE defined by mathematical expressions

#### variables

The name of the variables (i.e., fields) in the order they are expected to appear in the state.

Type

tuple

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur.

- **rhs** (*dict*) The expressions defining the evolution rate. The dictionary keys define the name of the fields whose evolution is considered, while the values specify their evolution rate as a string that can be parsed by sympy. These expression may contain variables (i.e., the fields themselves, spatial coordinates of the grid, and *t* for the time), standard local mathematical operators defined by sympy, and the operators defined in the *pde* package. Note that operators need to be specified with their full name, i.e., *laplace* for a scalar Laplacian and *vector\_laplace* for a Laplacian operating on a vector field. Moreover, the dot product between two vector fields can be denoted by using *dot(field1, field2)* in the expression, an outer product is calculated using *outer(field1, field2)*, and *integral(field)* denotes an integral over a field. More information can be found in the *expression documentation*.
- bc (BoundariesData) Boundary conditions for the operators used in the expression. The conditions here are applied to all operators that do not have a specialized condition given in bc\_ops. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by {'value': NUM}) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by {'derivative': DERIV})

are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documentation*.

- bc\_ops (dict) Special boundary conditions for some operators. The keys in this dictionary specify where the boundary condition will be applied. The keys follow the format "VARI-ABLE:OPERATOR", where VARIABLE specifies the expression in *rhs* where the boundary condition is applied to the operator specified by OPERATOR. For both identifiers, the wildcard symbol "\*" denotes that all fields and operators are affected, respectively. For instance, the identifier "c:\*" allows specifying a condition for all operators of the field named *c*.
- **user\_funcs** (*dict*, *optional*) A dictionary with user defined functions that can be used in the expressions in *rhs*.
- **consts** (*dict*, *optional*) A dictionary with user defined constants that can be used in the expression. These can be either scalar numbers or fields defined on the same grid as the actual simulation.
- **noise** (float or ndarray) Variance of additive Gaussian white noise. The default value of zero implies deterministic partial differential equations will be solved. Different noise magnitudes can be supplied for each field in coupled PDEs by either specifying a sequence of numbers or a dictionary with values for each field.
- **rng** (Generator) Random number generator (default: default\_rng()) used for stochastic simulations. Note that this random number generator is only used for numpy function, while compiled numba code uses the random number generator of numba. Moreover, in simulations using multiprocessing, setting the same generator in all processes might yield unintended correlations in the simulation results.

**Note:** The order in which the fields are given in *rhs* defines the order in which they need to appear in the *state* variable when the evolution rate is calculated. Note that *dict* keep the insertion order since Python version 3.7, so a normal dictionary can be used to define the equations.

### evolution\_rate(state, t=0.0)

evaluate the right hand side of the PDE

## Parameters

- **state** (FieldBase) The field describing the state of the PDE
- t (float) The current time point

### Returns

Field describing the evolution rate of the PDE

# Return type

FieldBase

# property expressions: dict[str, str]

show the expressions of the PDE

# 4.3.9 pde.pdes.swift\_hohenberg module

The Swift-Hohenberg equation

**class** SwiftHohenbergPDE (*rate=0.1*, *kc2=1.0*, *delta=1.0*, \*, *bc='auto\_periodic\_neumann'*, *bc\_lap=None*) Bases: *PDEBase* 

Dases. I DEDase

The Swift-Hohenberg equation

The mathematical definition is

$$\partial_t c = \left[\epsilon - \left(k_c^2 + \nabla^2\right)^2\right]c + \delta c^2 - c^3$$

where c is a scalar field and  $\epsilon$ ,  $k_c^2$ , and  $\delta$  are parameters of the equation.

#### Parameters

- **rate** (*float*) The bifurcation parameter  $\epsilon$
- kc2 (float) Squared wave vector  $k_c^2$  of the linear instability
- delta (float) Parameter  $\delta$  of the non-linearity
- bc (BoundariesData) The boundary conditions applied to the field. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by *['value': NUM]*) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by *['derivative': DERIV]*) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documentation*.
- **bc\_lap** (*BoundariesData* / *None*) The boundary conditions applied to the second derivative of the scalar field c. If *None*, the same boundary condition as *bc* is chosen. Otherwise, this supports the same options as *bc*.

### evolution\_rate(state, t=0)

evaluate the right hand side of the PDE

#### **Parameters**

- state (ScalarField) The scalar field describing the concentration distribution
- t (float) The current time point

#### Returns

Scalar field describing the evolution rate of the PDE

Return type

ScalarField

### explicit\_time\_dependence: bool | None = False

Flag indicating whether the right hand side of the PDE has an explicit time dependence.

#### Туре

bool

# property expression: str

the expression of the right hand side of this PDE

Type str

# 4.3.10 pde.pdes.wave module

A simple wave equation

**class** WavePDE (*speed=1*, *bc='auto\_periodic\_neumann'*)

Bases: PDEBase

A simple wave equation

The mathematical definition,  $\partial_t^2 u = c^2 \nabla^2 u$ , is implemented as two first-order equations,

$$\partial_t u = v$$
$$\partial_t v = c^2 \nabla^2 u$$

where c sets the wave speed and v is an auxiallary field. Note that the class expects an initial condition specifying both fields, which can be created using the *WavePDE.get\_initial\_condition()* method. The result will also return two fields.

#### **Parameters**

- **speed** (*float*) The speed *c* of the wave
- **bc** (BoundariesData) The boundary conditions applied to the field u. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non-periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by *{value': NUM}*) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by *{'derivative': DERIV}*) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documenta-tion*.

evolution\_rate(state, t=0)

evaluate the right hand side of the PDE

# Parameters

- **state** (FieldCollection) The fields u and v
- t (float) The current time point

#### Returns

Fields describing the evolution rates of the PDE

#### **Return type**

FieldCollection

### explicit\_time\_dependence: bool | None = False

Flag indicating whether the right hand side of the PDE has an explicit time dependence.

Type bool

#### property expressions: dict[str, str]

the expressions of the right hand side of this PDE

Type dict

```
get_initial_condition (u, v=None)
```

create a suitable initial condition

#### Parameters

- u (ScalarField) The initial density on the grid
- **v** (ScalarField, optional) The initial rate of change. This is assumed to be zero if the value is omitted.

### Returns

The combined fields u and v, suitable for the simulation

#### Return type

FieldCollection

# 4.4 pde.solvers package

Solvers define how a PDE is solved, i.e., how the initial state is advanced in time.

Controller	class controlling a simulation
ExplicitSolver	various explicit PDE solvers
ExplicitMPISolver	various explicit PDE solve using MPI
ImplicitSolver	implicit (backward) Euler PDE solver
CrankNicolsonSolver	Crank-Nicolson solver
AdamsBashforthSolver	explicit Adams-Bashforth multi-step solver
ScipySolver	PDE solver using scipy.integrate.
	solve_ivp().
registered_solvers	returns all solvers that are currently registered

**class** AdamsBashforthSolver (*pde*, \*, *backend='auto'*)

Bases: SolverBase

explicit Adams-Bashforth multi-step solver

#### **Parameters**

- pde (PDEBase) The partial differential equation that should be solved
- **backend** (*str*) Determines how the function is created. Accepted values are 'numpy` and 'numba'. Alternatively, 'auto' lets the code decide for the most optimal backend.

name = 'adams-bashforth'

```
class Controller (solver, t_range, tracker='auto')
```

Bases: object

class controlling a simulation

The controller calls a solver to advance the simulation into the future and it takes care of trackers that analyze and modify the state periodically.

## Parameters

• **solver** (*SolverBase*) – Solver instance that is used to advance the simulation in time

- **t\_range** (*float* or *tuple*) Sets the time range for which the simulation is run. If only a single value *t\_end* is given, the time range is assumed to be [0, *t\_end*].
- **tracker** (*TrackerCollectionDataType*) Defines trackers that process the state of the simulation at specified times. A tracker is either an instance of *Tracker-Base* or a string identifying a tracker (possible identifiers can be obtained by calling *get\_named\_trackers()*). Multiple trackers can be specified as a list. The default value *auto* checks the state for consistency (tracker 'consistency') and displays a progress bar (tracker 'progress') when tqdm is installed. More general trackers are defined in *trackers*, where all options are explained in detail. In particular, the time points where the tracker analyzes data can be chosen when creating a tracker object explicitly.

### diagnostics: dict[str, Any]

diagnostic information (available after simulation finished)

# Туре

dict

run (initial\_state, dt=None)

run the simulation

Diagnostic information about the solver are available in the *diagnostics* property after this function has been called.

#### **Parameters**

- **initial\_state** (*FieldBase*) The initial state of the simulation. This state will be copied and thus not modified by the simulation. Instead, the final state will be returned and trackers can be used to record intermediate states.
- dt (float) Time step of the chosen stepping scheme. If *None*, a default value based on the stepper will be chosen.

#### Returns

The state at the final time point. If multiprocessing is used, only the main node will return the state. All other nodes return None.

#### **Return type**

TState | None

#### property t\_range: tuple[float, float]

start and end time of the simulation

# Туре

tuple

**class CrankNicolsonSolver** (*pde*, \*, *maxiter=100*, *maxerror=0.0001*, *explicit\_fraction=0*, *backend='auto'*)

# Bases: SolverBase

Crank-Nicolson solver

- **pde** (*PDEBase*) The instance describing the pde that needs to be solved
- maxiter (*int*) The maximal number of iterations per step
- **maxerror** (float) The maximal error that is permitted in each step
- **explicit\_fraction** (*float*) Hyperparameter determining the fraction of explicit time stepping in the implicit step. *explicit\_fraction* == 0 is the simple Crank-Nicolson scheme, while *explicit\_fraction* == 1 reduces to the explicit Euler method. Intermediate values can improve convergence.

• **backend** (*str*) – Determines how the function is created. Accepted values are 'numpy` and 'numba'. Alternatively, 'auto' lets the code decide for the most optimal backend.

#### name = 'crank-nicolson'

```
class ExplicitSolver (pde, scheme='euler', *, backend='auto', adaptive=False, tolerance=0.0001)
```

Bases: AdaptiveSolverBase

various explicit PDE solvers

### Parameters

- **pde** (*PDEBase*) The instance describing the pde that needs to be solved
- **scheme** (*str*) Defines the explicit scheme to use. Supported values are 'euler' and 'runge-kutta' (or 'rk' for short).
- **backend** (*str*) Determines how the function is created. Accepted values are 'numpy` and 'numba'. Alternatively, 'auto' lets the code decide for the most optimal backend.
- **adaptive** (bool) When enabled, the time step is adjusted during the simulation using the error tolerance set with *tolerance*.
- **tolerance** (*float*) The error tolerance used in adaptive time stepping. This is used in adaptive time stepping to choose a time step which is small enough so the truncation error of a single step is below *tolerance*.

```
name = 'explicit'
```

**class** ImplicitSolver (*pde*, *maxiter=100*, *maxerror=0.0001*, *backend='auto'*)

Bases: SolverBase

implicit (backward) Euler PDE solver

#### **Parameters**

- **pde** (*PDEBase*) The instance describing the pde that needs to be solved
- **maxiter** (*int*) The maximal number of iterations per step
- **maxerror** (float) The maximal error that is permitted in each step
- **backend** (*str*) Determines how the function is created. Accepted values are 'numpy` and 'numba'. Alternatively, 'auto' lets the code decide for the most optimal backend.

name = 'implicit'

### class ScipySolver(pde, backend='auto', \*\*kwargs)

Bases: SolverBase

PDE solver using scipy.integrate.solve\_ivp().

This class is a thin wrapper around scipy.integrate.solve\_ivp(). In particular, it supports all the methods implemented by this function and exposes its arguments, so details can be controlled.

- pde (*PDEBase*) The instance describing the pde that needs to be solved
- **backend** (*str*) Determines how the function is created. Accepted values are 'numpy` and 'numba'. Alternatively, 'auto' lets the code decide for the most optimal backend.
- \*\*kwargs All extra arguments are forwarded to scipy.integrate.solve\_ivp().

#### make\_stepper (state, dt=None)

return a stepper function

#### **Parameters**

- **state** (FieldBase) An example for the state from which the grid and other information can be extracted.
- dt (float) Initial time step for the simulation. If *None*, the solver will choose a suitable initial value.

#### Returns

Function that can be called to advance the *state* from time *t\_start* to time *t\_end*.

#### **Return type**

Callable[[FieldBase, float, float], float]

```
name = 'scipy'
```

#### registered\_solvers()

returns all solvers that are currently registered

#### Returns

List with the names of the solvers

**Return type** 

list of str

# 4.4.1 pde.solvers.adams\_bashforth module

Defines an explicit Adams-Bashforth solver

#### class AdamsBashforthSolver (pde, \*, backend='auto')

Bases: SolverBase

explicit Adams-Bashforth multi-step solver

# Parameters

- pde (PDEBase) The partial differential equation that should be solved
- **backend** (*str*) Determines how the function is created. Accepted values are 'numpy` and 'numba'. Alternatively, 'auto' lets the code decide for the most optimal backend.

info: dict[str, Any]

```
name = 'adams-bashforth'
```

# 4.4.2 pde.solvers.base module

Package that contains base classes for solvers.

Beside the abstract base class defining the interfaces, we also provide *AdaptiveSolverBase*, which contains methods for implementing adaptive solvers.

class AdaptiveSolverBase (pde, \*, backend='auto', adaptive=True, tolerance=0.0001)

Bases: SolverBase

base class for adaptive time steppers

#### Parameters

- **pde** (*PDEBase*) The instance describing the pde that needs to be solved
- **backend** (*str*) Determines how the function is created. Accepted values are 'numpy` and 'numba'. Alternatively, 'auto' lets the code decide for the most optimal backend.
- **adaptive** (bool) When enabled, the time step is adjusted during the simulation using the error tolerance set with *tolerance*.
- **tolerance** (*float*) The error tolerance used in adaptive time stepping. This is used in adaptive time stepping to choose a time step which is small enough so the truncation error of a single step is below *tolerance*.

```
dt_max: float = 1000000000.0
```

maximal time step that the adaptive solver will use

Туре

float

dt\_min: float = 1e-10

minimal time step that the adaptive solver will use

Type float

#### make\_stepper (state, dt=None)

return a stepper function using an explicit scheme

#### Parameters

- **state** (*FieldBase*) An example for the state from which the grid and other information can be extracted
- dt (float) Time step used (Uses SolverBase.dt\_default if None). This sets the initial time step for adaptive solvers.

#### Returns

Function that can be called to advance the *state* from time *t\_start* to time *t\_end*. The function call signature is (*state: numpy.ndarray, t\_start: float, t\_end: float*)

#### **Return type**

Callable[[FieldBase, float, float], float]

#### exception ConvergenceError

Bases: RuntimeError

indicates that an implicit step did not converge

#### **class SolverBase** (*pde*, \*, *backend='auto'*)

Bases: object

base class for PDE solvers

- pde (*PDEBase*) The partial differential equation that should be solved
- **backend** (*str*) Determines how the function is created. Accepted values are 'numpy` and 'numba'. Alternatively, 'auto' lets the code decide for the most optimal backend.

#### dt\_default: float = 0.001

default time step used if no time step was specified

# Туре

float

# classmethod from\_name (name, pde, \*\*kwargs)

create solver class based on its name

Solver classes are automatically registered when they inherit from *SolverBase*. Note that this also requires that the respective python module containing the solver has been loaded before it is attempted to be used.

#### **Parameters**

- **name** (*str*) The name of the solver to construct
- pde (PDEBase) The partial differential equation that should be solved
- **\*\*kwargs** Additional arguments for the constructor of the solver

#### Returns

An instance of a subclass of SolverBase

#### **Return type**

SolverBase

# make\_stepper (state, dt=None)

return a stepper function using an explicit scheme

#### Parameters

- **state** (*FieldBase*) An example for the state from which the grid and other information can be extracted
- dt (float) Time step used (Uses SolverBase.dt\_default if None)

#### Returns

Function that can be called to advance the *state* from time *t\_start* to time *t\_end*. The function call signature is (*state: numpy.ndarray, t\_start: float, t\_end: float*)

### **Return type**

Callable[[FieldBase, float, float], float]

```
registered_solvers = ['AdamsBashforthSolver', 'AdaptiveSolverBase',
'CrankNicolsonSolver', 'ExplicitMPISolver', 'ExplicitSolver',
'ImplicitSolver', 'ScipySolver', 'adams-bashforth', 'crank-nicolson',
'explicit', 'explicit_mpi', 'implicit', 'scipy']
```

# 4.4.3 pde.solvers.controller module

Defines a class controlling the simulations of PDEs

**class** Controller (*solver*, *t\_range*, *tracker='auto'*)

Bases: object

class controlling a simulation

The controller calls a solver to advance the simulation into the future and it takes care of trackers that analyze and modify the state periodically.

- **solver** (*SolverBase*) Solver instance that is used to advance the simulation in time
- **t\_range** (*float* or *tuple*) Sets the time range for which the simulation is run. If only a single value *t\_end* is given, the time range is assumed to be [0, *t\_end*].
- **tracker** (*TrackerCollectionDataType*) Defines trackers that process the state of the simulation at specified times. A tracker is either an instance of *Tracker-Base* or a string identifying a tracker (possible identifiers can be obtained by calling *get\_named\_trackers()*). Multiple trackers can be specified as a list. The default value *auto* checks the state for consistency (tracker 'consistency') and displays a progress bar (tracker 'progress') when tqdm is installed. More general trackers are defined in *trackers*, where all options are explained in detail. In particular, the time points where the tracker analyzes data can be chosen when creating a tracker object explicitly.

#### diagnostics: dict[str, Any]

diagnostic information (available after simulation finished)

Туре

dict

info: dict[str, Any]

run (initial\_state, dt=None)

run the simulation

Diagnostic information about the solver are available in the *diagnostics* property after this function has been called.

#### **Parameters**

- **initial\_state** (*FieldBase*) The initial state of the simulation. This state will be copied and thus not modified by the simulation. Instead, the final state will be returned and trackers can be used to record intermediate states.
- dt (float) Time step of the chosen stepping scheme. If *None*, a default value based on the stepper will be chosen.

#### Returns

The state at the final time point. If multiprocessing is used, only the main node will return the state. All other nodes return None.

# **Return type**

TState | None

#### property t\_range: tuple[float, float]

start and end time of the simulation

Туре

tuple

# 4.4.4 pde.solvers.crank\_nicolson module

Defines a Crank-Nicolson solver

```
class CrankNicolsonSolver (pde, *, maxiter=100, maxerror=0.0001, explicit_fraction=0, backend='auto')
```

Bases: SolverBase

Crank-Nicolson solver

Parameters

- **pde** (*PDEBase*) The instance describing the pde that needs to be solved
- maxiter (*int*) The maximal number of iterations per step
- **maxerror** (*float*) The maximal error that is permitted in each step
- **explicit\_fraction** (*float*) Hyperparameter determining the fraction of explicit time stepping in the implicit step. *explicit\_fraction* == 0 is the simple Crank-Nicolson scheme, while *explicit\_fraction* == 1 reduces to the explicit Euler method. Intermediate values can improve convergence.
- **backend** (*str*) Determines how the function is created. Accepted values are 'numpy' and 'numba'. Alternatively, 'auto' lets the code decide for the most optimal backend.

info: dict[str, Any]

name = 'crank-nicolson'

# 4.4.5 pde.solvers.explicit module

Defines an explicit solver supporting various methods

class ExplicitSolver (pde, scheme='euler', \*, backend='auto', adaptive=False, tolerance=0.0001)

Bases: AdaptiveSolverBase

various explicit PDE solvers

#### Parameters

- pde (*PDEBase*) The instance describing the pde that needs to be solved
- **scheme** (*str*) Defines the explicit scheme to use. Supported values are 'euler' and 'runge-kutta' (or 'rk' for short).
- **backend** (*str*) Determines how the function is created. Accepted values are 'numpy` and 'numba'. Alternatively, 'auto' lets the code decide for the most optimal backend.
- **adaptive** (bool) When enabled, the time step is adjusted during the simulation using the error tolerance set with *tolerance*.
- **tolerance** (*float*) The error tolerance used in adaptive time stepping. This is used in adaptive time stepping to choose a time step which is small enough so the truncation error of a single step is below *tolerance*.

info: dict[str, Any]

```
name = 'explicit'
```

# 4.4.6 pde.solvers.explicit\_mpi module

Defines an explicit solver using multiprocessing via MPI

**class ExplicitMPISolver** (pde, scheme='euler', decomposition=-1, \*, backend='auto', adaptive=False, tolerance=0.0001)

Bases: ExplicitSolver

various explicit PDE solve using MPI

**Warning:** This solver can only be used if MPI is properly installed. In particular, python scripts then need to be started using mpirun or mpiexec. Please refer to the documentation of your MPI distribution for details.

The main idea of the solver is to take the full initial state in the main node (ID 0) and split the grid into roughly equal subgrids. The main node then distributes these subfields to all other nodes and each node creates the right hand side of the PDE for itself (and independently). Each node then advances the PDE independently, ensuring proper coupling to neighboring nodes via special boundary conditions, which exchange field values between sub grids. This is implemented by the get\_boundary\_conditions() method of the sub grids, which takes the boundary conditions for the full grid and creates conditions suitable for the specific sub grid on the given node. The trackers (and thus all input and output) are only handled on the main node.

**Warning:** The function providing the right hand side of the PDE needs to support MPI. This is automatically the case for local evaluations (which only use the field value at the current position), for the differential operators provided by *pde*, and integration of fields. Similarly, *modify\_after\_step* can only be used to do local modifications since the field data supplied to the function is local to each MPI node.

#### Example

A minimal example using the MPI solver is

```
from pde import DiffusionPDE, ScalarField, UnitGrid
grid = UnitGrid([64, 64])
state = ScalarField.random_uniform(grid, 0.2, 0.3)
eq = DiffusionPDE(diffusivity=0.1)
result = eq.solve(state, t_range=10, dt=0.1, method="explicit_mpi")
if result is not None: # restrict the output to the main node
result.plot()
```

Saving this script as *multiprocessing.py*, a parallel simulation is started by

mpiexec -n 2 python3 multiprocessing.py

Here, the number 2 determines the number of cores that will be used. Note that macOS might require an additional hint on how to connect the processes even when they are run on the same machine (e.g., your workstation). It might help to run mpiexec -n 2 -host localhost python3 multiprocessing.py in this case

#### Parameters

• pde (*PDEBase*) – The instance describing the pde that needs to be solved

- **scheme** (*str*) Defines the explicit scheme to use. Supported values are 'euler' and 'runge-kutta' (or 'rk' for short).
- **decomposition** (*list of ints*) Number of subdivision in each direction. Should be a list of length *grid.num\_axes* specifying the number of nodes along this axis. If one value is -1, its value will be determined from the number of available nodes. The default value decomposed the first axis using all available nodes.
- **backend** (*str*) Determines how the function is created. Accepted values are 'numpy` and 'numba'. Alternatively, 'auto' lets the code decide for the most optimal backend.
- **adaptive** (*bool*) When enabled, the time step is adjusted during the simulation using the error tolerance set with *tolerance*.
- **tolerance** (*float*) The error tolerance used in adaptive time stepping. This is used in adaptive time stepping to choose a time step which is small enough so the truncation error of a single step is below *tolerance*.

#### make\_stepper (state, dt=None)

return a stepper function using an explicit scheme

#### Parameters

- **state** (*FieldBase*) An example for the state from which the grid and other information can be extracted
- dt (float) Time step of the explicit stepping. If *None*, this solver specifies 1e-3 as a default value.

#### Returns

Function that can be called to advance the *state* from time *t\_start* to time *t\_end*. The function call signature is (*state: numpy.ndarray, t\_start: float, t\_end: float*)

### **Return type**

Callable[[FieldBase, float, float], float]

name = 'explicit\_mpi'

# 4.4.7 pde.solvers.implicit module

Defines an implicit Euler solver

```
class ImplicitSolver (pde, maxiter=100, maxerror=0.0001, backend='auto')
```

Bases: SolverBase

implicit (backward) Euler PDE solver

- **pde** (*PDEBase*) The instance describing the pde that needs to be solved
- maxiter (int) The maximal number of iterations per step
- **maxerror** (*float*) The maximal error that is permitted in each step
- **backend** (*str*) Determines how the function is created. Accepted values are 'numpy` and 'numba'. Alternatively, 'auto' lets the code decide for the most optimal backend.

```
info: dict[str, Any]
```

```
name = 'implicit'
```

# 4.4.8 pde.solvers.scipy module

Defines a solver using scipy.integrate

#### class ScipySolver(pde, backend='auto', \*\*kwargs)

Bases: SolverBase

PDE solver using scipy.integrate.solve\_ivp().

This class is a thin wrapper around scipy.integrate.solve\_ivp(). In particular, it supports all the methods implemented by this function and exposes its arguments, so details can be controlled.

#### Parameters

- **pde** (*PDEBase*) The instance describing the pde that needs to be solved
- **backend** (*str*) Determines how the function is created. Accepted values are 'numpy` and 'numba'. Alternatively, 'auto' lets the code decide for the most optimal backend.
- \*\*kwargs All extra arguments are forwarded to scipy.integrate.solve\_ivp().

info: dict[str, Any]

#### make\_stepper (state, dt=None)

return a stepper function

#### Parameters

- **state** (FieldBase) An example for the state from which the grid and other information can be extracted.
- dt (float) Initial time step for the simulation. If *None*, the solver will choose a suitable initial value.

#### Returns

Function that can be called to advance the *state* from time *t\_start* to time *t\_end*.

#### **Return type**

Callable[[FieldBase, float, float], float]

name = 'scipy'

# 4.5 pde.storage package

Module defining classes for storing simulation data.

get_memory_storage	a context manager that can be used to create a MemoryS-
	torage
MemoryStorage	store discretized fields in memory
FileStorage	store discretized fields in a hdf5 file
MovieStorage	store discretized fields in a movie file

# 4.5.1 pde.storage.base module

Base classes for storing data

#### **class StorageBase** (*info=None*, *write\_mode='truncate\_once'*)

Bases: object

base class for storing time series of discretized fields

These classes store time series of *FieldBase*, i.e., they store the values of the fields at particular time points. Iterating of the storage will return the fields in order and individual time points can also be accessed.

#### **Parameters**

- **info** (*dict*) Supplies extra information that is stored in the storage
- write\_mode (*str*) Determines how new data is added to already existing one. Possible values are: 'append' (data is always appended), 'truncate' (data is cleared every time this storage is used for writing), or 'truncate\_once' (data is cleared for the first writing, but subsequent data using the same instances are appended). Alternatively, specifying 'readonly' will disable writing completely.

# append (field, time=None)

add field to the storage

### Parameters

- field (FieldBase) The field that is added to the storage
- **time** (*float*, *optional*) The time point

#### **Return type**

None

### apply (func, out=None, \*, progress=False)

applies function to each field in a storage

#### Parameters

- **func** (*callable*) The function to apply to each stored field. The function must either take as a single argument the field or as two arguments the field and the associated time point. In both cases, it should return a field.
- **out** (*StorageBase*) Storage to which the output is written. If omitted, a new *Memo-ryStorage* is used and returned
- progress (bool) Flag indicating whether the progress is shown during the calculation

#### Returns

The new storage that contains the data after the function func has been applied

### **Return type**

StorageBase

clear (clear\_data\_shape=False)

truncate the storage by removing all stored data.

#### Parameters

**clear\_data\_shape** (bool) – Flag determining whether the data shape is also deleted.

Return type None **copy** (*out=None*, \*, *progress=False*)

copies all fields in a storage to a new one

#### **Parameters**

- **out** (*StorageBase*) Storage to which the output is written. If omitted, a new *Memo-ryStorage* is used and returned
- progress (bool) Flag indicating whether the progress is shown during the calculation

#### Returns

The new storage that contains the copied data

#### **Return type**

StorageBase

#### data: Any

## property data\_shape: tuple[int, ...]

the current data shape.

Raises

RuntimeError – if data\_shape was not set

```
property dtype: dtype[Any] | None | type[Any] | _SupportsDType[dtype[Any]]
| str | tuple[Any, int] | tuple[Any, SupportsIndex |
Sequence[SupportsIndex]] | list[Any] | _DTypeDict | tuple[Any, Any]
```

the current data type.

Raises

**RuntimeError** – if data\_type was not set

#### end\_writing()

finalize the storage after writing

Return type None

#### extract\_field(field\_id, label=None)

extract the time course of a single field from a collection

This method makes a copy of the underlying data.

#### **Parameters**

- **field\_id** (*int or str*) The index into the field collection. This determines which field of the collection is returned. Instead of a numerical index, the field label can also be supplied. If there are multiple fields with the same label, only the first field is returned.
- **label** (*str*) The label of the returned field. If omitted, the stored label is used.

# Returns

a storage instance that contains the data for the single field

#### **Return type**

MemoryStorage

#### extract\_time\_range (t\_range=None)

extract a particular time interval

**Note:** This might return a view into the original data, so modifying the returned data can also change the underlying original data.

### Parameters

**t\_range** (*float or tuple*) – Determines the range of time points included in the result. If only a single number is given, all data up to this time point are included.

# Returns

a storage instance that contains the extracted data.

#### **Return type**

MemoryStorage

# property grid: GridBase | None

the grid associated with this storage

This returns None if grid was not stored in self.info.

Туре

GridBase

#### property has\_collection: bool

whether the storage is storing a collection

Туре

bool

#### items()

iterate over all times and stored fields, returning pairs

# Return type

Iterator[tuple[float, FieldBase]]

# property shape: tuple[int, ...] | None

the shape of the stored data

#### start\_writing(field, info=None)

initialize the storage for writing data

#### Parameters

- **field**(FieldBase) An example of the data that will be written to extract the grid and the data\_shape
- info (dict) Supplies extra information that is stored in the storage

Return type None

#### times: Sequence[float]

**tracker** (*interrupts=1*, \*, *transformation=None*, *interval=None*)

create object that can be used as a tracker to fill this storage

#### **Parameters**

• **interrupts** (InterruptsBase / float / str / Sequence[float] / ndarray) – Determines when the tracker interrupts the simulation. A single numbers determines an interval (measured in the simulation time unit) of regular interruption. A string is interpreted as a duration in real time assuming the format 'hh:mm:ss'. A list of numbers is taken as explicit simulation time points. More fine- grained contol is possible by passing an instance of classes defined in *interrupts*.

• **transformation** (*callable*, *optional*) – A function that transforms the current state into a new field or field collection, which is then stored. This allows to store derived quantities of the field during calculations. The argument needs to be a callable function taking 1 or 2 arguments. The first argument always is the current field, while the optional second argument is the associated time.

### Returns

The tracker that fills the current storage

#### Return type

StorageTracker

#### Example

The *transformation* argument allows storing additional fields:

```
def add_to_state(state):
    transformed_field = state.smooth(1)
    return field.append(transformed_field)

storage = pde.MemoryStorage()
tracker = storage.tracker(1, transformation=add_to_state)
eq.solve(..., tracker=tracker)
```

In this example, storage will contain a trajectory of the fields of the simulation as well as the smoothed fields. Other transformations are possible by defining appropriate add\_to\_state()

#### view\_field(field\_id)

returns a view into this storage focusing on a particular field

Note: Modifying data returned by the view will modify the underlying storage

#### **Parameters**

**field\_id** (*int or str*) – The index into the field collection. This determines which field of the collection is returned. Instead of a numerical index, the field label can also be supplied. If there are multiple fields with the same label, only the first field is returned.

#### Returns

A view into the storage only returning a single field

Return type StorageView

```
write_mode: Literal['append', 'readonly', 'truncate', 'truncate_once']
```

**class StorageTracker** (*storage*, *interrupts=1*, \*, *transformation=None*, *interval=None*)

Bases: TrackerBase

Tracker that stores data in special storage classes

# storage

The underlying storage class through which the data can be accessed

# Туре

StorageBase

#### **Parameters**

- **storage** (*StorageBase*) Storage instance to which the data is written
- interrupts (InterruptData) Determines when the tracker interrupts the simulation. A single numbers determines an interval (measured in the simulation time unit) of regular interruption. A string is interpreted as a duration in real time assuming the format 'hh:mm:ss'. A list of numbers is taken as explicit simulation time points. More fine- grained contol is possible by passing an instance of classes defined in *interrupts*.
- **transformation** (*callable*, *optional*) A function that transforms the current state into a new field or field collection, which is then stored. This allows to store derived quantities of the field during calculations. The argument needs to be a callable function taking 1 or 2 arguments. The first argument always is the current field, while the optional second argument is the associated time.

### finalize(info=None)

finalize the tracker, supplying additional information

#### Parameters

**info** (*dict*) – Extra information from the simulation

# **Return type**

None

### **handle** (*field*, t)

handle data supplied to this tracker

# Parameters

- field (FieldBase) The current state of the simulation
- t (float) The associated time

#### **Return type**

None

initialize (field, info=None)

#### Parameters

- field (FieldBase) An example of the data that will be analyzed by the tracker
- **info** (*dict*) Extra information from the simulation

# Returns

The first time the tracker needs to handle data

#### Return type

float

#### class StorageView (storage, \*, field)

Bases: object

represents a view into a storage that extracts a particular field

- **storage** (*StorageBase*) The storage providing the basic data
- **field** (*int* or *str*) The index into the field collection determining which field of the collection is returned. Instead of a numerical index, the field label can also be supplied. If there are multiple fields with the same label, only the first field is returned.

```
property grid: GridBase | None
```

```
has_collection: bool = False
```

items()

iterate over all times and stored fields, returning pairs

```
Return type
Iterator[tuple[float, DataFieldBase]]
property times: Sequence[float]
```

# 4.5.2 pde.storage.file module

Defines a class storing data on the file system using the hierarchical data format (hdf)

Bases: StorageBase

store discretized fields in a hdf5 file

#### Parameters

- filename (*str*) The path to the hdf5-file where the data is stored
- **info** (*dict*) Supplies extra information that is stored in the storage
- write\_mode (*str*) Determines how new data is added to already existing data. Possible values are: 'append' (data is always appended), 'truncate' (data is cleared every time this storage is used for writing), or 'truncate\_once' (data is cleared for the first writing, but appended subsequently). Alternatively, specifying 'readonly' will disable writing completely.
- max\_length (*int*, *optional*) Maximal number of entries that will be stored in the file. This can be used to preallocate data, which can lead to smaller files, but is also less flexible. Giving max\_length = None, allows for arbitrarily large data, which might lead to larger files.
- **compression** (bool) Whether to store the data in compressed form. Automatically enabled chunked storage.
- **keep\_opened** (bool) Flag indicating whether the file should be kept opened after each writing. If *False*, the file will be closed after writing a dataset. This keeps the file in a consistent state, but also requires more work before data can be written.
- **check\_mpi** (bool) If True, files will only be opened in the main node for an parallel simulation using MPI. This flag has no effect in serial code.

### clear (clear\_data\_shape=False)

truncate the storage by removing all stored data.

#### Parameters

**clear\_data\_shape** (bool) – Flag determining whether the data shape is also deleted.

#### close()

close the currently opened file

#### Return type None

# property data

The actual data for all time

# Туре

# ndarray

### end\_writing()

finalize the storage after writing.

This makes sure the data is actually written to a file when self.keep\_opened == False

### **Return type**

None

#### start\_writing(field, info=None)

initialize the storage for writing data

# Parameters

- **field**(FieldBase) An example of the data that will be written to extract the grid and the data\_shape
- **info** (*dict*) Supplies extra information that is stored in the storage

### **Return type**

# None

# property times

The times at which data is available

# Туре

ndarray

# 4.5.3 pde.storage.memory module

Defines a class storing data in memory.

```
class MemoryStorage (times=None, data=None, *, info=None, field_obj=None, write_mode='truncate_once')
Bases: StorageBase
```

store discretized fields in memory

- times (ndarray) Sequence of times for which data is known
- data (list of ndarray) The field data at the given times
- field\_obj (FieldBase) An instance of the field class store data for a single time point.
- **info** (*dict*) Supplies extra information that is stored in the storage
- write\_mode (*str*) Determines how new data is added to already existing data. Possible values are: 'append' (data is always appended), 'truncate' (data is cleared every time this storage is used for writing), or 'truncate\_once' (data is cleared for the first writing, but appended subsequently). Alternatively, specifying 'readonly' will disable writing completely.

### clear (clear\_data\_shape=False)

truncate the storage by removing all stored data.

```
Parameters 
clear_data_shape (bool) – Flag determining whether the data shape is also deleted.
```

Return type None

classmethod from\_collection (storages, label=None, \*, rtol=1e-05, atol=1e-08)

combine multiple memory storages into one

This method can be used to combine multiple time series of different fields into a single representation. This requires that all time series contain data at the same time points.

#### **Parameters**

- **storages** (*list*) A collection of instances of *StorageBase* whose data will be concatenated into a single MemoryStorage
- **label** (*str*, *optional*) The label of the instances of FieldCollection that represent the concatenated data
- rtol (float) Relative tolerance used when checking times for merging
- atol (float) Absolute tolerance used when checking times for merging

#### Returns

Storage containing all the data.

### **Return type**

MemoryStorage

classmethod from\_fields (times=None, fields=None, info=None, write\_mode='truncate\_once')
create MemoryStorage from a list of fields

### Parameters

- times (ndarray) Sequence of times for which data is known
- fields (list of FieldBase) The fields at all given time points
- info (dict) Supplies extra information that is stored in the storage
- write\_mode (*str*) Determines how new data is added to already existing data. Possible values are: 'append' (data is always appended), 'truncate' (data is cleared every time this storage is used for writing), or 'truncate\_once' (data is cleared for the first writing, but appended subsequently). Alternatively, specifying 'readonly' will disable writing completely.

### **Return type**

MemoryStorage

start\_writing(field, info=None)

initialize the storage for writing data

# Parameters

- field (FieldBase) An instance of the field class store data for a single time point.
- info (dict) Supplies extra information that is stored in the storage

**Return type** 

None

#### get\_memory\_storage (field, info=None)

a context manager that can be used to create a MemoryStorage

#### Example

This can be used to quickly store data:

```
with get_memory_storage(field_class) as storage:
    storage.append(numpy_array0, 0)
    storage.append(numpy_array1, 1)
# use storage thereafter
```

#### Parameters

- **field** (FieldBase) An instance of the field class store data for a single time point.
- **info** (*dict*) Supplies extra information that is stored in the storage

#### Yields

MemoryStorage

# 4.5.4 pde.storage.movie module

Defines a class storing data on the file system as a compressed movie

This package requires the optional ffmpeg-python package to use FFmpeg for reading and writing movies.

**class MovieStorage** (filename, \*, vmin=0, vmax=1, bits\_per\_channel=8, video\_format='auto', bitrate=-1, info=None, write\_mode='truncate\_once', write\_times=False, loglevel='warning')

#### Bases: StorageBase

store discretized fields in a movie file

This storage only works when the *ffmpeg* program and ffmpeg is installed. The default codec is FFV1, which supports lossless compression for various configurations. Not all video players support this codec, but VLC usually works quite well.

Note that important metainformation is stored as a comment in the movie, so this data must not be deleted or altered if the video should be read again.

**Warning:** This storage potentially compresses data and can thus lead to loss of some information. The data quality depends on many parameters, but most important are the bits per channel of the video format and the range that is encoded (determined by *vmin* and *vmax*).

Note also that selecting individual time points might be quite slow since the video needs to be read from the beginning each time. Instead, it is much more efficient to process entire videos (by iterating over them or using *items()*).

#### Parameters

• **filename** (*str*) – The path where the movie is stored. The file extension determines the container format of the movie. The standard codec FFV1 plays well with the ".avi", ".mkv", and ".mov" container format.

- **vmin** (*float or array*) Lowest values that are encoded (per field). Smaller values are clipped to this value.
- **vmax** (*float* or *array*) Highest values that are encoded (per field). Larger values are clipped to this value.
- **bits\_per\_channel** (*int*) The number of bits used per color channel. Typical values are 8 and 16. The relative accuracy of stored values is 0.01 and 0.0001, respectively.
- video\_format (*str*) Identifier for a video format from *formats*, which determines the number of channels, the bit depth of individual colors, and the codec. The special value *auto* tries to find a suitable format automatically, taking *bits\_per\_channel* into account.
- **bitrate** (*float*) The bitrate of the movie (in kilobits per second). The default value of -1 let's the encoder choose an appropriate bit rate.
- **info** (*dict*) Supplies extra information that is stored in the storage alongside additional information necessary to reconstruct fields and grids.
- write\_mode (*str*) Determines how new data is added to already existing data. Possible values are: 'append' (data is always appended), 'truncate' (data is cleared every time this storage is used for writing), or 'truncate\_once' (data is cleared for the first writing, but appended subsequently). Alternatively, specifying 'readonly' will disable writing completely.
- write\_times (bool) Flag determining whether timestamps are written to a file. If True, a separate file with name filename + ".times" is created where the times are written as plain text. Without these timestamps, the time information might be inaccurate.
- **loglevel** (*str*) FFmpeg log level determining the amount of data sent to stdout. The default only emits warnings and errors, but setting this to *"info"* can be useful to get additioanl information about the encoding.

#### clear()

truncate the storage by removing all stored data.

#### close()

close the currently opened file

Return type None

#### property data

The actual data for all time

#### Туре

ndarray

# end\_writing()

finalize the storage after writing

Return type

None

#### items()

iterate over all times and stored fields, returning pairs

#### Return type

Iterator[tuple[float, FieldBase]]

#### start\_writing(field, info=None)

initialize the storage for writing data

#### **Parameters**

- **field**(FieldBase) An example of the data that will be written to extract the grid and the data\_shape
- info (dict) Supplies extra information that is stored in the storage

# Return type

None

# times: Sequence[float]

The times at which data is available

### Type

#### ndarray

# tracker (interrupts=1, \*, transformation=None)

create object that can be used as a tracker to fill this storage

### Parameters

- **interrupts** (InterruptsBase / float / str / Sequence[float] / ndarray) Determines when the tracker interrupts the simulation. A single numbers determines an interval (measured in the simulation time unit) of regular interruption. A string is interpreted as a duration in real time assuming the format 'hh:mm:ss'. A list of numbers is taken as explicit simulation time points. More fine- grained contol is possible by passing an instance of classes defined in *interrupts*.
- **transformation** (*callable*, *optional*) A function that transforms the current state into a new field or field collection, which is then stored. This allows to store derived quantities of the field during calculations. The argument needs to be a callable function taking 1 or 2 arguments. The first argument always is the current field, while the optional second argument is the associated time.

### Returns

The tracker that fills the current storage

# **Return type**

StorageTracker

### Example

The transformation argument allows storing additional fields:

```
def add_to_state(state):
    transformed_field = state.smooth(1)
    return field.append(transformed_field)

storage = pde.MemoryStorage()
tracker = storage.tracker(1, transformation=add_to_state)
eq.solve(..., tracker=tracker)
```

In this example, storage will contain a trajectory of the fields of the simulation as well as the smoothed fields. Other transformations are possible by defining appropriate  $add_to_state()$ 

# 4.6 pde.tools package

Package containing several tools required in py-pde

cache	Functions, classes, and decorators for managing caches
config	Handles configuration variables of the package
cuboid	An n-dimensional, axes-aligned cuboid
docstrings	Methods for automatic transformation of docstrings
expressions	Handling mathematical expressions with sympy
ffmpeg	Functions for interacting with FFmpeg
math	Auxiliary mathematical functions
misc	Miscellaneous python functions
mpi	Auxillary functions and variables for dealing with MPI
	multiprocessing
numba	Helper functions for just-in-time compilation with numba
output	Python functions for handling output
parameters	Infrastructure for managing classes with parameters
parse_duration	Parsing time durations from strings
plotting	Tools for plotting and controlling plot output using context
	managers
spectral	Functions making use of spectral decompositions
typing	Provides support for mypy type checking of the package

# 4.6.1 pde.tools.cache module

Functions, classes, and decorators for managing caches

cached_property	Decorator to use a method as a cached property
cached_method	Decorator to enable caching of a method
hash_mutable	return hash also for (nested) mutable objects.
hash_readable	return human readable hash also for (nested) mutable ob-
	jects.
make_serializer	returns a function that serialize data with the given method.
make_unserializer	returns a function that unserialize data with the given method
DictFiniteCapacity	cache with a limited number of items
SerializedDict	a key value database which is stored on the disk

class DictFiniteCapacity(\*args, \*\*kwargs)

 $Bases: \, \texttt{OrderedDict}$ 

cache with a limited number of items

### check\_length()

ensures that the dictionary does not grow beyond its capacity

default\_capacity: int = 100

update ( $[E, ]^{**F}$ )  $\rightarrow$  None. Update D from dict/iterable E and F.

If E is present and has a .keys() method, then does: for k in E: D[k] = E[k] If E is present and lacks a .keys() method, then does: for k, v in E: D[k] = v In either case, this is followed by: for k in F: D[k] = F[k]

**class** SerializedDict (key\_serialization='pickle', value\_serialization='pickle', storage\_dict=None)

Bases: MutableMapping

a key value database which is stored on the disk

This class provides hooks for converting arbitrary keys and values to strings, which are then stored in the database.

provides a dictionary whose keys and values are serialized

### **Parameters**

- **key\_serialization** (*str*) Determines the serialization method for keys
- value\_serialization (str) Determines the serialization method for values
- **storage\_dict** (*dict*) Can be used to chose a different dictionary for the underlying storage mechanism, e.g., storage\_dict = PersistentDict()

**class cached\_method** (*factory=None*, *extra\_args=None*, *ignore\_args=None*, *hash\_function='hash\_mutable'*, *doc=None*, *name=None*)

Bases: \_class\_cache

Decorator to enable caching of a method

The function is only called the first time and each successive call returns the cached result of the first call.

#### Example

The decorator can be used like so:

#### class Foo:

```
@cached_method
def bar(self):
    return "Cached"
foo = Foo()
result = foo.bar()
```

The data is stored in a dictionary named \_cache\_methods attached to the instance of each object. The cache can thus be cleared by setting self.\_cache\_methods = {}. The cache of specific property can be cleared using self.\_cache\_methods[property\_name] = {}, where property\_name is the name of the property.

decorator that caches calls in a dictionary attached to the instances. This can be used with most classes

#### Example

An example for using the class is:

```
class Foo():
```

```
@cached_property()
def property(self):
    return "Cached property"
```

(continues on next page)

(continued from previous page)

```
@cached_method()
  def method(self):
        return "Cached method"
foo = Foo()
foo.property
foo.method()
```

The cache can be cleared by setting foo.\_cache\_methods = {} if the cache factory is a simple dict, i.e, if factory == None. Alternatively, each cached method has a clear\_cache\_of\_obj() method, which clears the cache of this particular method. In the example above we could thus call foo.bar. clear\_cache\_of\_obj(foo) to clear the cache.

Note that the object instance has to be passed as a parameter, since the method bar() is defined on the class, not the instance, i.e., we could also call Foo.bar.clear\_cache\_of\_obj(foo). To clear the cache from within a method, one can thus call self.method\_name.clear\_cache\_of\_obj(self), where *method\_name* is the name of the method whose cache is cleared

# Example

An advanced example is:

```
class Foo():
    def get_cache(self, name):
        # `name` is the name of the method to cache
        return DictFiniteCapacity()
    @cached_method(factory='get_cache')
    def foo(self):
        return "Cached"
```

- **factory** (*callable*) Function/class creating an empty cache. *dict* by default. This can be used with user-supplied storage backends by. The cache factory should return a dict-like object that handles the cache for the given method.
- **extra\_args** (*list*) List of attributes of the class that are included in the cache key. They are then treated as if they are supplied as arguments to the method. This is important to include when the result of a method depends not only on method arguments but also on instance attributes.
- **ignore\_args** (*list*) List of keyword arguments that are not included in the cache key. These should be arguments that do not influence the result of a method, e.g., because they only affect how intermediate results are displayed.
- hash\_function (str) An identifier determining what hash function is used on the arguments
- doc (str) Optional string giving the docstring of the decorated method
- **name** (*str*) Optional string giving the name of the decorated method

**class cached\_property** (*factory=None*, *extra\_args=None*, *ignore\_args=None*, *hash\_function='hash\_mutable'*, *doc=None*, *name=None*)

Bases: \_class\_cache

Decorator to use a method as a cached property

The function is only called the first time and each successive call returns the cached result of the first call.

#### Example

Here is an example for how to use the decorator:

```
class Foo():
    @cached_property
    def bar(self):
        return "Cached"
foo = Foo()
result = foo.bar
```

The data is stored in a dictionary named <u>\_cache\_methods</u> attached to the instance of each object. The cache can thus be cleared by setting self.\_cache\_methods = {}. The cache of specific property can be cleared using self.\_cache\_methods[property\_name] = {}, where property\_name is the name of the property.

Adapted from <https://wiki.python.org/moin/PythonDecoratorLibrary>.

decorator that caches calls in a dictionary attached to the instances. This can be used with most classes

#### Example

An example for using the class is:

```
class Foo():
    @cached_property()
    def property(self):
        return "Cached property"
    @cached_method()
    def method(self):
        return "Cached method"
foo = Foo()
foo.property
foo.method()
```

The cache can be cleared by setting foo.\_cache\_methods = {} if the cache factory is a simple dict, i.e, if factory == None. Alternatively, each cached method has a clear\_cache\_of\_obj() method, which clears the cache of this particular method. In the example above we could thus call foo.bar. clear\_cache\_of\_obj(foo) to clear the cache.

Note that the object instance has to be passed as a parameter, since the method bar() is defined on the class, not the instance, i.e., we could also call Foo.bar.clear\_cache\_of\_obj(foo). To clear the cache from within

a method, one can thus call self.method\_name.clear\_cache\_of\_obj(self), where *method\_name* is the name of the method whose cache is cleared

### Example

An advanced example is:

```
class Foo():
    def get_cache(self, name):
        # `name` is the name of the method to cache
        return DictFiniteCapacity()
    @cached_method(factory='get_cache')
    def foo(self):
        return "Cached"
```

#### **Parameters**

- **factory** (*callable*) Function/class creating an empty cache. *dict* by default. This can be used with user-supplied storage backends by. The cache factory should return a dict-like object that handles the cache for the given method.
- **extra\_args** (*list*) List of attributes of the class that are included in the cache key. They are then treated as if they are supplied as arguments to the method. This is important to include when the result of a method depends not only on method arguments but also on instance attributes.
- **ignore\_args** (*list*) List of keyword arguments that are not included in the cache key. These should be arguments that do not influence the result of a method, e.g., because they only affect how intermediate results are displayed.
- hash\_function (str) An identifier determining what hash function is used on the arguments
- doc (*str*) Optional string giving the docstring of the decorated method
- **name** (*str*) Optional string giving the name of the decorated method

### hash\_mutable(obj)

return hash also for (nested) mutable objects.

### Notes

This function might be a bit slow, since it iterates over all containers and hashes objects recursively. Moreover, the returned value might change with each run of the python interpreter, since the hash values of some basic objects, like *None*, change with each instance of the interpreter.

#### **Parameters**

**obj** – A general python object

#### Returns

A hash value associated with the data of *obj* 

Return type int

#### hash\_readable(obj)

return human readable hash also for (nested) mutable objects.

This function returns a JSON-like representation of the object. The function might be a bit slow, since it iterates over all containers and hashes objects recursively. Note that this hash function tries to return the same value for equivalent objects, but it does not ensure that the objects can be reconstructed from this data.

### Parameters

**obj** – A general python object

#### Returns

A hash value associated with the data of *obj* 

Return type

str

### make\_serializer(method)

returns a function that serialize data with the given method. Note that some of the methods destroy information and cannot be reverted.

### **Parameters**

method (str) - An identifier determining the serializer that will be returned

### Returns

A function that serializes objects

Return type callable

### make\_unserializer(method)

returns a function that unserialize data with the given method

This is the inverse function of *make\_serializer()*.

### Parameters

**method** (str) – An identifier determining the unserializer that will be returned

### Returns

A function that serializes objects

## Return type

callable

### objects\_equal(a, b)

compares two objects to see whether they are equal

In particular, this uses numpy.array\_equal() to check for numpy arrays

#### Parameters

- a The first object
- b The second object

#### Returns

Whether the two objects are considered equal

### **Return type**

bool

## 4.6.2 pde.tools.config module

Handles configuration variables of the package

Config	class handling the package configuration
get_package_versions	tries to load certain python packages and returns their ver-
	sion
parse_version_str	helper function converting a version string into a list of
	integers
check_package_version	checks whether a package has a sufficient version
packages_from_requirements	read package names from a requirements file
environment	obtain information about the compute environment

### **class** Config(*items=None*, *mode='update'*)

Bases: UserDict

class handling the package configuration

### Parameters

- **items** (*dict*, *optional*) Configuration values that should be added or overwritten to initialize the configuration.
- mode (str) Defines the mode in which the configuration is used. Possible values are
  - insert: any new configuration key can be inserted
  - update: only the values of pre-existing items can be updated
  - locked: no values can be changed

Note that the items specified by *items* will always be inserted, independent of the mode.

### to\_dict()

convert the configuration to a simple dictionary

### Returns

A representation of the configuration in a normal dict.

### **Return type**

dict

### check\_package\_version (package\_name, min\_version)

checks whether a package has a sufficient version

### Parameters

- package\_name (str)-
- min\_version (str) -

### environment()

obtain information about the compute environment

#### Returns

information about the python installation and packages

### **Return type**

dict

#### get\_package\_versions (packages, \*, na\_str='not available')

tries to load certain python packages and returns their version

### Parameters

- packages (list) The names of all packages
- **na\_str** (*str*) Text to return if package is not available

### Returns

Dictionary with version for each package name

#### **Return type**

dict

### packages\_from\_requirements(requirements\_file)

read package names from a requirements file

### Parameters

requirements\_file (str or Path) - The file from which everything is read

#### Returns

list of package names

Return type list[str]

#### parse\_version\_str(ver\_str)

helper function converting a version string into a list of integers

Parameters ver\_str(str)-Return type

list[int]

## 4.6.3 pde.tools.cuboid module

An n-dimensional, axes-aligned cuboid

This module defines the *Cuboid* class, which represents an n-dimensional cuboid that is aligned with the axes of a Cartesian coordinate system.

**class** Cuboid (*pos*, *size*, *mutable=True*)

Bases: object

class that represents a cuboid in n dimensions

defines a cuboid from a position and a size vector

#### Parameters

- **pos** (*list*) The position of the lower left corner. The length of this list determines the dimensionality of space
- **size** (*list*) The size of the cuboid along each dimension.
- mutable (bool) Flag determining whether the cuboid parameters can be changed

property bounds: tuple[tuple[float, float], ...]

#### buffer (amount=0, inplace=False)

dilate the cuboid by a certain amount in all directions

```
Parameters
    amount (float | ndarray) -
```

Return type Cuboid

### property centroid

#### contains\_point (points)

returns a True when points are within the Cuboid

### **Parameters**

points (ndarray) - List of point coordinates

#### Returns

list of booleans indicating which points are inside

### **Return type**

ndarray

#### copy()

Return type Cuboid

### property corners: tuple[ndarray, ndarray]

return coordinates of two extreme corners defining the cuboid

#### property diagonal: float

returns the length of the diagonal

### property dim: int

### classmethod from\_bounds (bounds, \*\*kwargs)

create cuboid from bounds

#### Parameters

**bounds** (*list*) – Two dimensional array of axes bounds

#### Returns

cuboid with positive size

## Return type

Cuboid

### classmethod from\_centerpoint (centerpoint, size, \*\*kwargs)

create cuboid from two points

### **Parameters**

- centerpoint (list) Coordinates of the center
- **size** (*list*) Size of the cuboid

### Returns

cuboid with positive size

### Return type

Cuboid

#### classmethod from\_points (p1, p2, \*\*kwargs)

create cuboid from two points

### **Parameters**

- p1 (list) Coordinates of first corner point
- p2 (list) Coordinates of second corner point

### Returns

cuboid with positive size

Return type Cuboid

### property mutable: bool

property size: ndarray

#### property surface\_area: float

surface area of a cuboid in n dimensions.

The surface area is the volume of the (n-1)-dimensional hypercubes that bound the current cuboid:

- n = 1: the number of end points (2)
- n = 2: the perimeter of the rectangle
- n = 3: the surface area of the cuboid

### property vertices: list[list[float]]

return the coordinates of all the corners

#### property volume: float

### asanyarray\_flags (data, dtype=None, writeable=True)

turns data into an array and sets the respective flags.

A copy is only made if necessary

#### Parameters

- data (ndarray) numpy array that whose flags are adjusted
- dtype (dtype [Any] | None | type [Any] | \_SupportsD-Type [dtype [Any]] | str | tuple [Any, int] | tuple [Any, SupportsIndex | Sequence [SupportsIndex]] | list [Any] | \_DTypeDict | tuple [Any, Any]) - the resulant dtype
- writeable (bool) Flag determining whether the results is writable

#### Returns

array with same data as *data* but with flags adjusted.

#### **Return type**

ndarray

## 4.6.4 pde.tools.docstrings module

Methods for automatic transformation of docstrings

get_text_block	return a single text block
replace_in_docstring	replace a text in a docstring using the correct indentation
fill_in_docstring	decorator that replaces text in the docstring of a function

### $\texttt{fill\_in\_docstring}(f)$

decorator that replaces text in the docstring of a function

Parameters f (TFunc) –

## Return type

TFunc

get\_text\_block (identifier)

return a single text block

### Parameters

identifier (str) – The name of the text block

### Returns

the text block as one long line.

### **Return type**

str

### replace\_in\_docstring(f, token, value, docstring=None)

replace a text in a docstring using the correct indentation

### Parameters

- **f** (*callable*) The function with the docstring to handle
- token (*str*) The token to search for
- **value** (*str*) The replacement string
- docstring (*str*) A docstring that should be used instead of f.\_\_doc\_\_\_

#### Returns

The function with the modified docstring

### **Return type**

callable

## 4.6.5 pde.tools.expressions module

Handling mathematical expressions with sympy

This module provides classes representing expressions that can be provided as human-readable strings and are converted to numpy and numba representations using sympy.

parse_number	return a number compiled from an expression
ScalarExpression	describes a mathematical expression of a scalar quantity
TensorExpression	describes a mathematical expression of a tensorial quan- tity
evaluate	evaluate an expression involving fields

class ExpressionBase (expression, signature=None, \*, user\_funcs=None, consts=None, repl=None)

Bases: object

abstract base class for handling expressions

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur.

#### **Parameters**

- expression (sympy.core.basic.Basic) A sympy expression or array. This could for instance be an instance of Expr or NDimArray.
- **signature** (*list of str*, *optional*) The signature defines which variables are expected in the expression. This is typically a list of strings identifying the variable names. Individual names can be specified as list, in which case any of these names can be used. The first item in such a list is the definite name and if another name of the list is used, the associated variable is renamed to the definite name. If signature is *None*, all variables in *expressions* are allowed.
- **user\_funcs** (*dict*, *optional*) A dictionary with user defined functions that used in the expression.
- **consts** (*dict*, *optional*) A dictionary with user defined constants that can be used in the expression. The values of these constants should either be numbers or ndarray.
- **repl** (*dict*, *optional*) Replacements that are applied to symbols before turning the expression into a python equivalent.

### property complex: bool

whether the expression contains the imaginary unit I

### Type

bool

#### property constant: bool

whether the expression is a constant

Туре

bool

## depends\_on(variable)

determine whether the expression depends on variable

```
Parameters
variable (str) – the name of the variable to check for
```

#### Returns

whether the variable appears in the expression

## Return type

bool

### property expression: str

the expression in string form

Туре

str

### get\_compiled (single\_arg=False)

return numba function evaluating expression

### Parameters

**single\_arg** (bool) – Determines whether the function takes all variables in a single argument as an array or whether all variables need to be supplied separately.

#### Returns

the compiled function

Return type function

property rank: int

the rank of the expression

Type int

#### abstract property shape: tuple[int, ...]

the shape of the tensor

Туре

#### tuple

**class** ScalarExpression (*expression=0*, *signature=None*, \*, *user\_funcs=None*, *consts=None*, *repl=None*, *explicit\_symbols=None*, *allow\_indexed=False*)

Bases: ExpressionBase

describes a mathematical expression of a scalar quantity

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur.

#### **Parameters**

- **expression** (*str or float*) The expression, either a number or a string that sympy can parse.
- **signature** (*list of str*) The signature defines which variables are expected in the expression. This is typically a list of strings identifying the variable names. Individual names can be specified as lists, in which case any of these names can be used. The first item in such a list is the definite name and if another name of the list is used, the associated variable is renamed to the definite name. If signature is *None*, all variables in *expressions* are allowed.
- user\_funcs (dict, optional) A dictionary with user defined functions that used in the expression.
- **consts** (*dict*, *optional*) A dictionary with user defined constants that can be used in the expression. The values of these constants should either be numbers or ndarray.

- **repl** (*dict*, *optional*) Replacements that are applied to symbols before turning the expression into a python equivalent.
- **explicit\_symbols** (*list of str*) List of symbols that need to be interpreted as general sympy symbols
- **allow\_indexed** (*bool*) Whether to allow indexing of variables. If enabled, array variables are allowed to be indexed using square bracket notation.

### copy()

return a copy of the current expression

### Return type

ScalarExpression

### derivatives

differentiate the expression with respect to all variables

### differentiate(var)

return the expression differentiated with respect to var

Parameters var (str) -

Return type

Scalar Expression

### property is\_zero: bool

returns whether the expression is zero

Type bool

```
shape: tuple[int, ...] = ()
```

property value: int | float | complex

the value for a constant expression

Type float

**class TensorExpression** (*expression*, *signature=None*, \*, *user\_funcs=None*, *consts=None*, *repl=None*, *explicit\_symbols=None*)

Bases: ExpressionBase

describes a mathematical expression of a tensorial quantity

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur.

### **Parameters**

- **expression** (*str or float*) The expression, either a number or a string that sympy can parse.
- **signature** (*list of str*) The signature defines which variables are expected in the expression. This is typically a list of strings identifying the variable names. Individual names can be specified as list, in which case any of these names can be used. The first item in such

a list is the definite name and if another name of the list is used, the associated variable is renamed to the definite name. If signature is *None*, all variables in *expressions* are allowed.

- user\_funcs (dict, optional) A dictionary with user defined functions that used in the expression.
- **consts** (*dict*, *optional*) A dictionary with user defined constants that can be used in the expression. The values of these constants should either be numbers or ndarray.
- **repl** (*dict*, *optional*) Replacements that are applied to symbols before turning the expression into a python equivalent.
- **explicit\_symbols** (*list of str*) List of symbols that need to be interpreted as general sympy symbols

#### derivatives

differentiate the expression with respect to all variables

### differentiate(var)

return the expression differentiated with respect to var

Parameters

**var**(*str*)-

Return type TensorExpression

get\_compiled\_array(single\_arg=True)

compile the tensor expression such that a numpy array is returned

#### Parameters

**single\_arg** (*bool*) – Whether the compiled function expects all arguments as a single array or whether they are supplied individually.

#### **Return type**

Callable[[ndarray, ndarray | None], ndarray]

#### property rank: int

rank of the tensor expression

Туре

int

property shape: tuple[int, ...]

the shape of the tensor

## Туре

tuple

#### property value

the value for a constant expression

evaluate an expression involving fields

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur.

### Parameters

- **expression** (*str*) The expression, which is parsed by sympy. The expression may contain variables (i.e., fields and spatial coordinates of the grid), standard local mathematical operators defined by sympy, and the operators defined in the *pde* package. Note that operators need to be specified with their full name, i.e., *laplace* for a scalar Laplacian and *vector\_laplace* for a Laplacian operating on a vector field. Moreover, the dot product between two vector fields can be denoted by using *dot(field1, field2)* in the expression, and *outer(field1, field2)* calculates an outer product. More information can be found in the *expression documentation*.
- **fields** (dict or *FieldCollection*) Dictionary of the fields involved in the expression. The dictionary keys specify the field names allowed in *expression*. Alternatively, *fields* can be a *FieldCollection* with unique labels.
- **bc** (BoundariesData) Boundary conditions for the operators used in the expression. The conditions here are applied to all operators that do not have a specialized condition given in *bc\_ops*. Boundary conditions are generally given as a list with one condition for each axis. For periodic axes, only periodic boundary conditions are allowed (indicated by 'periodic' and 'anti-periodic'). For non- periodic axes, different boundary conditions can be specified for the lower and upper end (using a tuple of two conditions). For instance, Dirichlet conditions enforcing a value NUM (specified by {'value': NUM}) and Neumann conditions enforcing the value DERIV for the derivative in the normal direction (specified by {'derivative': DERIV}) are supported. Note that the special value 'natural' imposes periodic boundary conditions for periodic axis and a vanishing derivative otherwise. More information can be found in the *boundaries documentation*.
- bc\_ops (dict) Special boundary conditions for some operators. The keys in this dictionary specify the operator to which the boundary condition will be applied.
- user\_funcs (dict, optional) A dictionary with user defined functions that can be used in the expressions in *rhs*.
- **consts** (*dict*, *optional*) A dictionary with user defined constants that can be used in the expression. These can be either scalar numbers or fields defined on the same grid as the actual simulation.
- **label** (*str*) Name of the field that is returned.

#### Returns

The resulting field. The rank of the returned field (and thus the precise class) is determined automatically.

#### Return type

pde.fields.base.DataFieldBase

#### parse\_number (expression, variables=None)

return a number compiled from an expression

**Warning:** This implementation uses exec() and should therefore not be used in a context where malicious input could occur.

#### **Parameters**

- expression (str or Number) An expression that can be interpreted as a number
- variables (dict) A dictionary of values that replace variables in the expression

Returns

the calculated value

Return type Number

## 4.6.6 pde.tools.ffmpeg module

Functions for interacting with FFmpeg

FFmpegFormat	defines a FFmpeg format used for storing field data in a video
formats	dict of pre-defined FFmpegFormat formats
find_format	find a defined FFmpegFormat that satisifies the require-
	ments

**class FFmpegFormat** (*pix\_fmt\_file*, *pix\_fmt\_data*, *channels*, *bits\_per\_channel*, *dtype*, *codec='ffv1'*)

Bases: object

defines a FFmpeg format used for storing field data in a video

**Note:** All pixel formats supported by FFmpeg can be obtained by running ffmpeg -pix\_fmts. However, not all pixel formats are supported by all codecs. Supported pixel formats are listed in the output of ffmpeg -h encoder=<ENCODER>, where <*ENCODER*> is one of the encoders listed in ffmpeg -codecs.

### Parameters

- pix\_fmt\_file (str)-
- pix\_fmt\_data(str)-
- channels (int) -
- **bits\_per\_channel** (*int*) -
- dtype (dtype[Any] | None | type[Any] | \_SupportsD-Type[dtype[Any]] | str | tuple[Any, int] | tuple[Any, SupportsIndex | Sequence[SupportsIndex]] | list[Any] | \_DTypeDict | tuple[Any, Any])-
- codec (str)-

### bits\_per\_channel: int

number of bits per color channel in this pixel format

Type int

### property bytes\_per\_channel: int

number of bytes per color channel

Type int

### channels: int

number of color channels in this pixel format

Type int

### codec: str = 'ffv1'

name of the codec that supports this pixel format

Type str

```
data_from_frame (frame_data)
```

converts data stored in a color channel to normalized data

```
Parameters
frame_data (ndarray) -
```

### data\_to\_frame (normalized\_data)

converts normalized data to data being stored in a color channel

```
Parameters
    normalized_data(ndarray) -
```

Return type ndarray

```
dtype: dtype[Any] | None | type[Any] | _SupportsDType[dtype[Any]] | str |
tuple[Any, int] | tuple[Any, SupportsIndex | Sequence[SupportsIndex]] |
list[Any] | _DTypeDict | tuple[Any, Any]
```

numpy dtype corresponding to the data of a single channel

#### property max\_value: float | int

maximal value stored in a color channel

### pix\_fmt\_data: str

name of the pixel format used in the frame data

Type str

#### pix\_fmt\_file: str

name of the pixel format used in the codec

Туре

str

```
find_format (channels, bits_per_channel=8)
```

find a defined FFmpegFormat that satisifies the requirements

### Parameters

- channels (int) Minimal number of color channels
- bits\_per\_channel (int) Minimal number of bits per channel

#### Returns

Identifier for a format that satisifies the requirements (but might have more channels or more bits per channel then requested. *None* is returned if no format can be identified.

### **Return type**

str

```
formats = {'gbrap16le': FFmpegFormat(pix_fmt_file='gbrap16le',
    pix_fmt_data='gbrap16le', channels=4, bits_per_channel=16,
    dtype=dtype('uint16'), codec='ffv1'), 'gbrp16le':
    FFmpegFormat(pix_fmt_file='gbrp16le', pix_fmt_data='gbrp16le', channels=3,
    bits_per_channel=16, dtype=dtype('uint16'), codec='ffv1'), 'gray':
    FFmpegFormat(pix_fmt_file='gray', pix_fmt_data='gray', channels=1,
    bits_per_channel=8, dtype=<class 'numpy.uint8'>, codec='ffv1'), 'gray16le':
    FFmpegFormat(pix_fmt_file='gray16le', pix_fmt_data='gray16le', channels=1,
    bits_per_channel=16, dtype=dtype('uint16'), codec='ffv1'), 'rgb24':
    FFmpegFormat(pix_fmt_file='rgb24', pix_fmt_data='rgb24', channels=3,
    bits_per_channel=8, dtype=<class 'numpy.uint8'>, codec='ffv1'), 'rgb32':
    FFmpegFormat(pix_fmt_file='rgb32', pix_fmt_data='rgb32', channels=4,
    bits_per_channel=8, dtype=<class 'numpy.uint8'>, codec='ffv1')}
```

dict of pre-defined *FFmpegFormat* formats

### 4.6.7 pde.tools.math module

Auxiliary mathematical functions

#### class OnlineStatistics (\*args, \*\*kwargs)

Bases: OnlineStatistics

class for using an online algorithm for calculating statistics

```
class_type = jitclass.OnlineStatistics#7fbb2c768b10<min:float64,
max:float64,mean:float64,_mean2:float64,count:uint64>
```

#### count: int

recorded number of items

Type

#### mean: float

recorded mean

Type float

class SmoothData1D (x, y, sigma=None)

Bases: object

allows smoothing data in 1d using a Gaussian kernel of defined width

The data is given a pairs of x and y, the assumption being that there is an underlying relation y = f(x).

initialize with data

#### Parameters

- **x** List of x values
- y List of y values
- **sigma** (*float*) The size of the smoothing window in units of *x*. If omitted, the average distance of x values multiplied by *sigma\_auto\_scale* is used.

```
property bounds: tuple[float, float]
```

return minimal and maximal x values

```
derivative (xs)
```

return the derivative of the smoothed values for the positions xs

Note that this value

```
Parameters
xs (list of ndarray) – the x-values
```

Returns

The associated values of the derivative

Return type ndarray

### sigma\_auto\_scale: float = 10

scale for setting automatic values for sigma

Туре

float

## 4.6.8 pde.tools.misc module

Miscellaneous python functions

module_available	check whether a python module is available
ensure_directory_exists	creates a folder if it not already exists
preserve_scalars	decorator that makes vectorized methods work with scalars
decorator_arguments	make a decorator usable with and without arguments:
import_class	import a class or module given an identifier
classproperty	decorator that can be used to define read-only properties
	for classes.
hybridmethod	descriptor that can be used as a decorator to allow calling a method both as a classmethod and an instance method
estimate_computation_speed	estimates the computation speed of a function
hdf_write_attributes	write (JSON-serialized) attributes to a hdf file
number	convert a value into a float or complex number
get_common_dtype	returns a dtype in which all arguments can be represented
number_array	convert an array with arbitrary dtype either to np.double or np.cdouble

### class classproperty(fget=None, doc=None)

Bases: property

decorator that can be used to define read-only properties for classes.

This is inspired by the implementation of astropy, see astropy.org.

### Example

The decorator can be used much like the *property* decorator:

```
class Test():
```

item: str = 'World'

(continues on next page)

(continued from previous page)

```
@classproperty
def message(cls):
    return 'Hello ' + cls.item
print(Test.message)
```

deleter (fdel)

Descriptor to obtain a copy of the property with a different deleter.

getter (fget)

Descriptor to obtain a copy of the property with a different getter.

setter (fset)

Descriptor to obtain a copy of the property with a different setter.

#### decorator\_arguments (decorator)

make a decorator usable with and without arguments:

The resulting decorator can be used like @decorator or @decorator(\*args, \*\*kwargs)

Inspired by https://stackoverflow.com/a/14412901/932593

#### **Parameters**

decorator (Callable) – the decorator that needs to be modified

Returns

the decorated function

Return type Callable

### ensure\_directory\_exists(folder)

creates a folder if it not already exists

### Parameters

folder (str) – path of the new folder

#### estimate\_computation\_speed (func, \*args, \*\*kwargs)

estimates the computation speed of a function

#### Parameters

func (callable) – The function to call

### Returns

the number of times the function can be calculated in one second. The inverse is thus the runtime in seconds per function call

#### **Return type**

float

### get\_common\_dtype(\*args)

returns a dtype in which all arguments can be represented

#### Parameters

\*args – All items (arrays, scalars, etc) to be checked

Returns: numpy.cdouble if any entry is complex, otherwise np.double

#### hdf\_write\_attributes (hdf\_path, attributes=None, raise\_serialization\_error=False)

write (JSON-serialized) attributes to a hdf file

### **Parameters**

- hdf\_path Path to a group or dataset in an open HDF file
- **attributes** (*dict*) Dictionary with values written as attributes
- **raise\_serialization\_error** (*bool*) Flag indicating whether serialization errors are raised or silently ignored

### Return type

None

### class hybridmethod (fclass, finstance=None, doc=None)

#### Bases: object

descriptor that can be used as a decorator to allow calling a method both as a classmethod and an instance method

Adapted from https://stackoverflow.com/a/28238047

classmethod (fclass)

instancemethod (finstance)

### import\_class(identifier)

import a class or module given an identifier

#### Parameters

**identifier** (*str*) – The identifier can be a module or a class. For instance, calling the function with the string *identifier* == *'numpy.linalg.norm'* is roughly equivalent to running *from numpy.linalg import norm* and would return a reference to *norm*.

### module\_available (module\_name)

check whether a python module is available

### Parameters

module\_name (str) - The name of the module

#### Returns

True if the module can be imported and False otherwise

**Return type** 

bool

### number (value)

convert a value into a float or complex number

### Parameters

**value** (*Number or str*) – The value which needs to be converted

#### **Return type**

int | float | complex

### **Result:**

Number: A complex number or a float if the imaginary part vanishes

#### number\_array (data, dtype=None, copy=True)

convert an array with arbitrary dtype either to np.double or np.cdouble

#### Parameters

- **data** (ndarray) The data that needs to be converted to a float array. This can also be any iterable of numbers.
- **dtype** (*numpy dtype*) The data type of the field. All the numpy dtypes are supported. If omitted, it will be determined from *data* automatically.
- **copy** (bool) Whether the data must be copied (in which case the original array is left untouched). Note that data will always be copied when changing the dtype.

#### Returns

An array with the correct dtype

### Return type

ndarray

#### preserve\_scalars(method)

decorator that makes vectorized methods work with scalars

This decorator allows to call functions that are written to work on numpy arrays to also accept python scalars, like *int* and *float*. Essentially, this wrapper turns them into an array and unboxes the result.

#### **Parameters**

method (TFunc) – The method being decorated

#### Returns

The decorated method

Return type TFunc

#### skipUnlessModule (module\_names)

decorator that skips a test when a module is not available

#### Parameters

module\_names (str) - The name of the required module(s)

#### Returns

A function, so this can be used as a decorator

### **Return type**

Callable[[TFunc], TFunc]

### 4.6.9 pde.tools.mpi module

Auxillary functions and variables for dealing with MPI multiprocessing

**Warning:** These functions are mostly no-ops unless MPI is properly installed and python code was started using mpirun or mpiexec. Please refer to the documentation of your MPI distribution for details.

mpi_send	send data to another MPI node
mpi_recv	receive data from another MPI node
mpi_allreduce	combines data from all MPI nodes

### initialized: bool = False

Flag determining whether mpi was initialized (and is available)

#### Type bool

is\_main: bool = True

Flag indicating whether the current process is the main process (with ID 0)

### Туре

bool

### mpi\_allreduce (data, operator=None)

combines data from all MPI nodes

Note that complex datatypes and user-defined functions are not properly supported.

### Parameters

- data Data being send from this node to all others
- operator (*int* / *str* / *None*) The operator used to combine all data. Possible options are summarized in the IntEnum numba\_mpi.Operator.

### Returns

The accumulated data

### mpi\_recv(data, source, tag)

receive data from another MPI node

### Parameters

- data A buffer into which the received data is written
- **dest** (*int*) The ID of the sending node
- tag (int) A numeric tag identifying the message

### **Return type**

None

### mpi\_send(data, dest, tag)

send data to another MPI node

### Parameters

- data The data being send
- **dest** (*int*) The ID of the receiving node
- **tag** (*int*) A numeric tag identifying the message

### **Return type**

None

### ol\_mpi\_allreduce(data, operator=None)

overload the mpi\_allreduce function

### Parameters

operator (int | str | None) -

ol\_mpi\_recv(data, source, tag)

overload the *mpi\_recv* function

### Parameters

- source (int) -
- **tag**(*int*)-

```
ol_mpi_send(data, dest, tag)
```

overload the *mpi\_send* function

### **Parameters**

- **dest** (*int*) -
- **tag**(*int*)-

### parallel\_run: bool = False

Flag indicating whether the current run is using multiprocessing

Type bool

rank: int = 0

ID of the current process

Туре

int

```
size: int = 1
```

Total process count

Type int

## 4.6.10 pde.tools.numba module

Helper functions for just-in-time compilation with numba

### class Counter(value=0)

Bases: object

helper class for implementing JIT\_COUNT

We cannot use a simple integer for this, since integers are immutable, so if one imports JIT\_COUNT from this module it would always stay at the fixed value it had when it was first imported. The workaround would be to import the symbol every time the counter is read, but this is error-prone. Instead, we implement a thin wrapper class around an int, which only supports reading and incrementing the value. Since this object is now mutable it can be used easily. A disadvantage is that the object needs to be converted to int before it can be used in most expressions.

```
Parameters
```

**value** (int) -

increment()

```
flat_idx(arr, i)
```

helper function allowing indexing of scalars as if they arrays

Parameters

• arr (ndarray) -

• **i**(*int*)-

**Return type** 

int | float | complex

### get\_common\_numba\_dtype(\*args)

returns a numba numerical type in which all arrays can be represented

#### Parameters

**\*args** – All items to be tested

Returns: numba.complex128 if any entry is complex, otherwise numba.double

jit (function, signature=None, parallel=False, \*\*kwargs)

apply nb.jit with predefined arguments

### Parameters

- function (TFunc) The function which is jitted
- signature Signature of the function to compile
- parallel (bool) Allow parallel compilation of the function
- \*\*kwargs Additional arguments to nb.jit

#### Returns

Function that will be compiled using numba

Return type TFunc

### make\_array\_constructor (arr)

returns an array within a jitted function using basic information

#### **Parameters**

arr (ndarray) - The array that should be accessible within jit

#### **Return type**

*Callable*[[], *ndarray*]

**Warning:** A reference to the array needs to be retained outside the numba code to prevent garbage collection from removing the array

#### numba\_dict (data=None)

converts a python dictionary to a numba typed dictionary

```
Parameters
```

```
data(dict[str, Any] | None)-
```

### Return type

Dict | None

#### numba\_environment()

return information about the numba setup used

### Returns

(dict) information about the numba setup

**Return type** 

dict[str, Any]

### **ol\_flat\_idx** (*arr*, *i*)

helper function allowing indexing of scalars as if they arrays

### random\_seed(seed=0)

sets the seed of the random number generator of numpy and numba

```
Parameters
seed (int) - Sets random seed
```

Return type None

## 4.6.11 pde.tools.output module

Python functions for handling output

get_progress_bar_class	returns a class that behaves as progress bar.
display_progress	displays a progress bar when iterating
in_jupyter_notebook	checks whether we are in a jupyter notebook
BasicOutput	class that writes text line to stdout
JupyterOutput	class that writes text lines as html in a jupyter cell

class BasicOutput (stream=<\_io.TextIOWrapper name='<stdout>' mode='w' encoding='utf-8'>)

Bases: OutputBase

class that writes text line to stdout

### **Parameters**

**stream** – The stream where the lines are written

### show()

shows the actual text

### class JupyterOutput (header=", footer=")

#### Bases: OutputBase

class that writes text lines as html in a jupyter cell

#### Parameters

- header (*str*) The html code written before all lines
- **footer** (*str*) The html code written after all lines

### show()

shows the actual text

### class OutputBase

Bases: object

base class for output management

### abstract show()

shows the actual text

### display\_progress (iterator, total=None, enabled=True, \*\*kwargs)

displays a progress bar when iterating

### Parameters

• iterator (*iter*) – The iterator

- total (*int*) Total number of steps
- enabled (bool) Flag determining whether the progress is display
- \*\*kwargs All extra arguments are forwarded to the progress bar class

### Returns

A class that behaves as the original iterator, but shows the progress alongside iteration.

### get\_progress\_bar\_class (fancy=True)

returns a class that behaves as progress bar.

This either uses classes from the optional *tqdm* package or a simple version that writes dots to stderr, if the class it not available.

#### **Parameters**

**fancy** (*bool*) – Flag determining whether a fancy progress bar should be used in jupyter notebooks (if ipywidgets is installed)

### in\_jupyter\_notebook()

checks whether we are in a jupyter notebook

Return type

bool

## 4.6.12 pde.tools.parameters module

Infrastructure for managing classes with parameters

One aim is to allow easy management of inheritance of parameters.

Parameter	class representing a single parameter
DeprecatedParameter	a parameter that can still be used normally but is depre- cated
HideParameter	a helper class that allows hiding parameters of the parent classes
Parameterized	a mixin that manages the parameters of a class
get_all_parameters	get a dictionary with all parameters of all registered classes

**class DeprecatedParameter** (*name*, *default\_value=None*, *cls=<class 'object'>*, *description=''*, *hidden=False*, *extra=None*)

### Bases: Parameter

a parameter that can still be used normally but is deprecated

initialize a parameter

Parameters

- **name** (*str*) The name of the parameter
- default\_value The default value
- **cls** The type of the parameter, which is used for conversion
- **description** (*str*) A string describing the impact of this parameter. This description appears in the parameter help
- hidden (bool) Whether the parameter is hidden in the description summary

• extra (dict) – Extra arguments that are stored with the parameter

#### class HideParameter (name)

```
Bases: object
```

a helper class that allows hiding parameters of the parent classes

Parameters

**name** (*str*) – The name of the parameter

**class Parameter** (*name*, *default\_value=None*, *cls=<class 'object'>*, *description="*, *hidden=False*, *extra=None*)

Bases: object

class representing a single parameter

initialize a parameter

Parameters

- **name** (*str*) The name of the parameter
- default\_value The default value
- **cls** The type of the parameter, which is used for conversion
- **description** (*str*) A string describing the impact of this parameter. This description appears in the parameter help
- hidden (bool) Whether the parameter is hidden in the description summary
- **extra** (*dict*) Extra arguments that are stored with the parameter

#### convert (value=None)

converts a value into the correct type for this parameter. If value is not given, the default value is converted.

Note that this does not make a copy of the values, which could lead to unexpected effects where the default value is changed by an instance.

### Parameters

**value** – The value to convert

Returns

The converted value, which is of type *self.cls* 

class Parameterized(parameters=None)

Bases: object

a mixin that manages the parameters of a class

initialize the parameters of the object

### Parameters

**parameters** (*dict*) - A dictionary of parameters to change the defaults. The allowed parameters can be obtained from *get\_parameters()* or displayed by calling *show\_parameters()*.

### get\_parameter\_default (name)

return the default value for the parameter with name

### Parameters

**name** (*str*) – The parameter name

**classmethod** get\_parameters (*include\_hidden=False*, *include\_deprecated=False*, *sort=True*) return a dictionary of parameters that the class supports

### Parameters

- include\_hidden (bool) Include hidden parameters
- include\_deprecated (bool) Include deprecated parameters
- **sort** (*bool*) Return ordered dictionary with sorted keys

#### Returns

a dictionary of instance of *Parameter* with their names as keys.

### **Return type**

dict

```
parameters_default: Sequence[Parameter | HideParameter] = []
```

**show\_parameters** (description=None, sort=False, show\_hidden=False, show\_deprecated=False)

show all parameters in human readable format

### Parameters

- **description** (*bool*) Flag determining whether the parameter description is shown. The default is to show the description only when we are in a jupyter notebook environment.
- **sort** (bool) Flag determining whether the parameters are sorted
- **show\_hidden** (bool) Flag determining whether hidden parameters are shown
- **show\_deprecated** (*bool*) Flag determining whether deprecated parameters are shown
- **default\_value** (bool) Flag determining whether the default values or the current values are shown

All flags default to False.

#### get\_all\_parameters (data='name')

get a dictionary with all parameters of all registered classes

#### Parameters

data (*str*) – Determines what data is returned. Possible values are 'name', 'value', or 'description', to return the respective information about the parameters.

#### **Return type**

dict[str, Any]

### sphinx\_display\_parameters (app, what, name, obj, options, lines)

helper function to display parameters in sphinx documentation

#### Example

This function should be connected to the 'autodoc-process-docstring' event like so:

app.connect('autodoc-process-docstring', sphinx\_display\_parameters)

## 4.6.13 pde.tools.parse\_duration module

Parsing time durations from strings

This module provides a function that parses time durations from strings. It has been copied from the django software, which comes with the following notes:

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### parse\_duration(value)

Parse a duration string and return a datetime.timedelta.

### Parameters

**value** (str) - A time duration given as text. The preferred format for durations is '%d %H:%M:%S.%f'. This function also supports ISO 8601 representation and PostgreSQL's day-time interval format.

### Returns

An instance representing the duration.

### **Return type**

datetime.timedelta

## 4.6.14 pde.tools.plotting module

Tools for plotting and controlling plot output using context managers

add_scaled_colorbar	add a vertical color bar to an image plot
disable_interactive	context manager disabling the interactive mode of mat- plotlib
plot_on_axes	decorator for a plot method or function that uses a single axes
plot_on_figure	decorator for a plot method or function that fills an entire figure
PlotReference	contains all information to update a plot element
BasicPlottingContext	basic plotting using just matplotlib
JupyterPlottingContext	plotting in a jupyter widget using the <i>inline</i> backend
get_plotting_context	returns a suitable plotting context
napari_add_layers	adds layers to a napari viewer

**class BasicPlottingContext** (*fig\_or\_ax=None*, *title=None*, *show=True*)

Bases: PlottingContextBase

basic plotting using just matplotlib

### **Parameters**

- fig\_or\_ax If axes are given, they are used. If a figure is given, it is set as active.
- title (*str*) The shown in the plot
- **show** (bool) Flag determining whether plots are actually shown

### class JupyterPlottingContext(title=None, show=True)

Bases: PlottingContextBase

plotting in a jupyter widget using the inline backend

#### **Parameters**

- title (*str*) The shown in the plot
- **show** (bool) Flag determining whether plots are actually shown

### close()

close the plot

### supports\_update: bool = False

flag indicating whether the context supports that plots can be updated with out redrawing the entire plot. The jupyter backend (*inline*) requires replotting of the entire figure, so an update is not supported.

### **class PlotReference** (*ax*, *element*, *parameters=None*)

Bases: object

contains all information to update a plot element

### Parameters

- **ax** (matplotlib.axes.Axes) The axes of the element
- element (matplotlib.artist.Artist) The actual element
- parameters (dict) Parameters to recreate the plot element

ax

element

#### parameters

#### class PlottingContextBase (*title=None*, *show=True*)

Bases: object

base class of the plotting contexts

### Example

The context wraps calls to the matplotlib.pyplot interface:

```
context = PlottingContext()
with context:
    plt.plot(...)
    plt.xlabel(...)
```

#### **Parameters**

- title (*str*) The shown in the plot
- **show** (bool) Flag determining whether plots are actually shown

#### close()

close the plot

```
fig: mpl_figure.Figure | None
```

#### supports\_update: bool = True

flag indicating whether the context supports that plots can be updated with out redrawing the entire plot

```
add_scaled_colorbar (axes_image, ax=None, aspect=20, pad_fraction=0.5, label=", **kwargs)
```

add a vertical color bar to an image plot

The height of the colorbar is now adjusted to the plot, so that the width determined by *aspect* is now given relative to the height. Moreover, the gap between the colorbar and the plot is now given in units of the fraction of the width by *pad\_fraction*.

Inspired by https://stackoverflow.com/a/33505522/932593

#### Parameters

- axes\_image(matplotlib.cm.ScalarMappable)-Mappable object, e.g., returned from matplotlib.pyplot.imshow()
- **ax** (matplotlib.axes.Axes) The current figure axes from which space is taken for the colorbar. If omitted, the axes in which the *axes\_image* is shown is taken.
- **aspect** (*float*) The target aspect ratio of the colorbar
- **pad\_fraction** (*float*) Width of the gap between colorbar and image
- **label** (*str*) Set a label for the colorbar
- \*\*kwargs Additional parameters are passed to colorbar call

#### Returns

The resulting Colorbar object

#### **Return type**

Colorbar

### disable\_interactive()

context manager disabling the interactive mode of matplotlib

This context manager restores the previous state after it is done. Details of the interactive mode are described in matplotlib.interactive().

#### get\_plotting\_context (context=None, title=None, show=True)

returns a suitable plotting context

#### Parameters

- **context** An instance of *PlottingContextBase* or an instance of matplotlib. axes.Axes or matplotlib.figure.Figure to determine where the plotting will happen. If omitted, the context is determined automatically.
- title (*str*) The title shown in the plot
- **show** (bool) Determines whether the plot is shown while the simulation is running. If *False*, the files are created in the background.

#### Returns

The plotting context

### **Return type**

PlottingContextBase

### in\_ipython()

try to detect whether we are in an ipython shell, e.g., a jupyter notebook

### **Return type**

bool

### napari\_add\_layers (viewer, layers\_data)

adds layers to a napari viewer

#### Parameters

- viewer (napar i.viewer.Viewer) The napari application
- **layers\_data** (*dict*) Data for all layers that will be added.

#### napari\_viewer (grid, run=None, close=False, \*\*kwargs)

creates an napari viewer for interactive plotting

#### Parameters

- grid (pde.grids.base.GridBase) The grid defining the space
- **run** (bool) Whether to run the event loop of napari.
- **close** (bool) Whether to close the viewer immediately (e.g. for testing)
- \*\*kwargs Extra arguments are passed to napari. Viewer

### **Return type**

Generator[napari.viewer.Viewer, None, None]

#### class nested\_plotting\_check

### Bases: object

context manager that checks whether it is the root plotting call

#### Example

The context manager can be used in plotting calls to check for nested plotting calls:

```
with nested_plotting_check() as is_outermost_plot_call:
    make_plot(...) # could potentially call other plotting methods
    if is_outermost_plot_call:
        plt.show()
```

plot\_on\_axes (wrapped=None, update\_method=None)

decorator for a plot method or function that uses a single axes

This decorator adds typical options for creating plots that fill a single axes. These options are available via keyword arguments. To avoid redundancy in describing these options in the docstring, the placeholder *{PLOT\_ARGS}* can be added to the docstring of the wrapped function or method and will be replaced by the appropriate text. Note that the decorator can be used on both functions and methods.

### Example

The following example illustrates how this decorator can be used to implement plotting for a given class. In particular, supplying the *update\_method* will allow efficient dynamical plotting:

```
class State:
    def __init__(self) -> None:
        self.data = np.arange(8)
    def _update_plot(self, reference):
        reference.element.set_ydata(self.data)
    @plot_on_axes(update_method='_update_plot')
    def plot(self, ax):
        line, = ax.plot(np.arange(8), self.data)
        return PlotReference(ax, line)
@plot_on_axes
def make_plot(ax):
        ax.plot(...)
```

When *update\_method* is absent, the method can still be used for plotting, but dynamic updating, e.g., by pde. trackers.PlotTracker, is not possible.

#### **Parameters**

- wrapped (callable) Function to be wrapped
- **update\_method** (*callable or str*) Method to call to update the plot. The argument of the new method will be the result of the initial call of the wrapped method.

#### plot\_on\_figure (wrapped=None, update\_method=None)

decorator for a plot method or function that fills an entire figure

This decorator adds typical options for creating plots that fill an entire figure. This decorator adds typical options for creating plots that fill a single axes. These options are available via keyword arguments. To avoid redundancy in describing these options in the docstring, the placeholder *[PLOT\_ARGS]* can be added to the docstring of the

wrapped function or method and will be replaced by the appropriate text. Note that the decorator can be used on both functions and methods.

### Example

The following example illustrates how this decorator can be used to implement plotting for a given class. In particular, supplying the *update\_method* will allow efficient dynamical plotting:

```
class State:
    def __init__(self) -> None:
        self.data = np.random.random((2, 8))
    def _update_plot(self, reference):
        ref1, ref2 = reference
        ref1.element.set_ydata(self.data[0])
        ref2.element.set_ydata(self.data[1])
    @plot_on_figure(update_method='_update_plot')
    def plot(self, fig):
        ax1, ax2 = fig.subplots(1, 2)
        l1, = ax1.plot(np.arange(8), self.data[0])
        12, = ax2.plot(np.arange(8), self.data[1])
        return [PlotReference(ax1, 11), PlotReference(ax2, 12)]
@plot_on_figure
def make_plot(fig):
    . . .
```

When *update\_method* is not supplied, the method can still be used for plotting, but dynamic updating, e.g., by pde.trackers.PlotTracker, is not possible.

### **Parameters**

- wrapped (callable) Function to be wrapped
- **update\_method** (*callable or str*) Method to call to update the plot. The argument of the new method will be the result of the initial call of the wrapped method.

## 4.6.15 pde.tools.spectral module

Functions making use of spectral decompositions

make_colored_noise	Return a function creating an array of random values that
	obey

**make\_colored\_noise** (*shape*, *dx*=1.0, *exponent*=0, *scale*=1, *rng*=None)

Return a function creating an array of random values that obey

$$\langle c(m{k})c(m{k'})
angle = \Gamma^2 |m{k}|^
u \delta(m{k}-m{k'})$$

in spectral space on a Cartesian grid. The special case  $\nu = 0$  corresponds to white noise. For simplicity, the correlations respect periodic boundary conditions.

**Parameters** 

- **shape** (*tuple of ints*) Number of supports points in each spatial dimension. The number of the list defines the spatial dimension.
- **dx** (*float* or *list* of *floats*) Discretization along each dimension. A uniform discretization in each direction can be indicated by a single number.
- **exponent** (*float*) Exponent  $\nu$  of the power spectrum
- scale (float) Scaling factor  $\Gamma$  determining noise strength
- **rng** (Generator) Random number generator (default: default\_rng())

### Returns

a function returning a random realization

## Return type

callable

## 4.6.16 pde.tools.typing module

Provides support for mypy type checking of the package

```
class AdjacentEvaluator(*args, **kwargs)
```

Bases: Protocol

```
class CellVolume (*args, **kwargs)
Bases: Protocol
```

```
class GhostCellSetter(*args, **kwargs)
Bases: Protocol
```

```
class OperatorFactory (*args, **kwargs)
```

Bases: Protocol

a factory function that creates an operator for a particular grid

```
class OperatorType(*args, **kwargs)
```

Bases: Protocol

an operator that acts on an array

```
class VirtualPointEvaluator(*args, **kwargs)
```

Bases: Protocol

# 4.7 pde.trackers package

Classes for tracking simulation results in controlled interrupts

Trackers are classes that periodically receive the state of the simulation to analyze, store, or output it. The trackers defined in this module are:

CallbackTracker	Tracker calling a function periodically
ProgressTracker	Tracker showing the progress of the simulation
PrintTracker	Tracker printing data to a stream (default: stdout)
PlotTracker	Tracker plotting data on screen, to files, or writes a movie
LivePlotTracker	PlotTracker with defaults for live plotting
DataTracker	Tracker storing custom data obtained by calling a function
SteadyStateTracker	Tracker aborting the simulation once steady state is reached
RuntimeTracker	Tracker interrupting the simulation once a duration has passed
ConsistencyTracker	Tracker interrupting the simulation when the state is not finite
InteractivePlotTracker	Tracker showing the state interactively in napari

Some trackers can also be referenced by name for convenience when using them in simulations. The lit of supported names is returned by get\_named\_trackers().

Multiple trackers can be collected in a *TrackerCollection*, which provides methods for handling them efficiently. Moreover, custom trackers can be implemented by deriving from *TrackerBase*. Note that trackers generally receive a view into the current state, implying that they can adjust the state by modifying it in-place. Moreover, trackers can abort the simulation by raising the special exception StopIteration.

For each tracker, the time at which the simulation is interrupted can be decided using one of the following classes:

FixedInterrupts	class representing a list of interrupt times
ConstantInterrupts	class representing equidistantly spaced time interrupts
LogarithmicInterrupts	class representing logarithmically spaced time interrupts
RealtimeInterrupts	class representing time interrupts spaced equidistantly in
	real time

In particular, interrupts can be specified conveniently using parse\_interrupt().

## 4.7.1 pde.trackers.base module

Base classes for trackers

#### exception FinishedSimulation

Bases: StopIteration

exception for signaling that simulation finished successfully

**class TrackerBase** (*interrupts*=1, \*, *interval*=None)

Bases: object

base class for implementing trackers

#### **Parameters**

**interrupts** (*InterruptData*) – Determines when the tracker interrupts the simulation. A single numbers determines an interval (measured in the simulation time unit) of regular interruption. A string is interpreted as a duration in real time assuming the format 'hh:mm:ss'. A list of numbers is taken as explicit simulation time points. More fine- grained contol is possible by passing an instance of classes defined in *interrupts*.

### finalize (info=None)

finalize the tracker, supplying additional information

### Parameters

**info** (*dict*) – Extra information from the simulation

Return type None

### classmethod from\_data(data, \*\*kwargs)

create tracker class from given data

### Parameters

data (str or TrackerBase) - Data describing the tracker

### Returns

An instance representing the tracker

Return type

TrackerBase

### **abstract** handle (*field*, *t*)

handle data supplied to this tracker

### Parameters

- field (FieldBase) The current state of the simulation
- t (float) The associated time

## Return type

None

### initialize (field, info=None)

initialize the tracker with information about the simulation

#### Parameters

- field (FieldBase) An example of the data that will be analyzed by the tracker
- **info** (*dict*) Extra information from the simulation

### Returns

The first time the tracker needs to handle data

Return type

float

### class TrackerCollection(trackers=None)

### Bases: object

List of trackers providing methods to handle them efficiently

### trackers

List of the trackers in the collection

Type list

### Parameters

trackers (list [TrackerBase] / None) - List of trackers that are to be handled.

### finalize (info=None)

finalize the tracker, supplying additional information

### Parameters

**info** (*dict*) – Extra information from the simulation

Return type None

#### classmethod from\_data(data, \*\*kwargs)

create tracker collection from given data

### **Parameters**

```
data (Sequence [TrackerBase | str] | TrackerBase | str | None) – Data describing the tracker collection
```

### Returns

An instance representing the tracker collection

### **Return type**

TrackerCollection

handle (state, t, atol=1e-08)

handle all trackers

### Parameters

- **state** (FieldBase) The current state of the simulation
- t (float) The associated time
- **atol** (*float*) An absolute tolerance that is used to determine whether a tracker should be called now or whether the simulation should be carried on more timesteps. This is basically used to predict the next time to decided which one is closer.

### Returns

The next time the simulation needs to be interrupted to handle a tracker.

### **Return type**

float

### initialize (field, info=None)

initialize the tracker with information about the simulation

### Parameters

- field (FieldBase) An example of the data that will be analyzed by the tracker
- **info** (*dict*) Extra information from the simulation

#### Returns

The first time the tracker needs to handle data

Return type

float

### time\_next\_action: float

The time of the next interrupt of the simulation

Туре

float

#### tracker\_action\_times: list[float]

Times at which the trackers need to be handled next

Type list

```
get_named_trackers()
```

returns all named trackers

Returns

a mapping of names to the actual tracker classes.

**Return type** 

dict

## 4.7.2 pde.trackers.interactive module

Special module for defining an interactive tracker that uses napari to display fields

**class** InteractivePlotTracker (*interrupts='0:01'*, \*, *close=True*, *show\_time=False*, *interval=None*)

Bases: TrackerBase

Tracker showing the state interactively in napari

**Note:** The interactive tracker uses the python multiprocessing module to run napari externally. The multiprocessing module has limitations on some platforms, which requires some care when writing your own programs. In particular, the main method needs to be safe-guarded so that the main module can be imported again after spawning a new process. An established pattern that works is to introduce a function *main* in your code, which you call using the following pattern

```
def main():
    # here goes your main code
if __name__ == "__main__":
    main()
```

The last two lines ensure that the *main* function is only called when the module is run initially and not again when it is re-imported.

#### **Parameters**

- **interrupts** (*InterruptData*) Determines when the tracker interrupts the simulation. A single numbers determines an interval (measured in the simulation time unit) of regular interruption. A string is interpreted as a duration in real time assuming the format 'hh:mm:ss'. A list of numbers is taken as explicit simulation time points. More fine- grained contol is possible by passing an instance of classes defined in *interrupts*.
- **close** (bool) Flag indicating whether the napari window is closed automatically at the end of the simulation. If *False*, the tracker blocks when *finalize* is called until the user closes napari manually.
- **show\_time** (bool) Whether to indicate the time

#### finalize(info=None)

finalize the tracker, supplying additional information

#### Parameters

**info** (*dict*) – Extra information from the simulation

Return type

None

## handle (state, t)

handle data supplied to this tracker

#### Parameters

- **state** (FieldBase) The current state of the simulation
- t (float) The associated time

## **Return type**

None

#### initialize (state, info=None)

initialize the tracker with information about the simulation

#### Parameters

- state (FieldBase) An example of the data that will be analyzed by the tracker
- **info** (*dict*) Extra information from the simulation

#### Returns

The first time the tracker needs to handle data

## **Return type**

float

name = 'interactive'

#### class NapariViewer(state, t\_initial=None)

#### Bases: object

allows viewing and updating data in a separate napari process

#### Parameters

- **state** (*pde.fields.base.FieldBase*) The initial state to be shown
- **t\_initial** (*float*) The initial time. If *None*, no time will be shown.

#### close (force=True)

closes the napari process

## Parameters

**force** (bool) – Whether to force closing of the napari program. If this is *False*, this method blocks until the user closes napari manually.

## update (state, t)

update the state in the napari viewer

- state (pde.fields.base.FieldBase) The new state
- t (float) Current time

napari\_process (data\_channel, initial\_data, t\_initial=None, viewer\_args=None)

multiprocessing.Process running napari

#### **Parameters**

- data\_channel (multiprocessing.Queue) queue instance to receive data to view
- **initial\_data** (*dict*) Initial data to be shown by napari. The layers are named according to the keys in the dictionary. The associated value needs to be a tuple, where the first item is a string indicating the type of the layer and the second carries the associated data
- **t\_initial** (*float*) Initial time
- viewer\_args (dict) Additional arguments passed to the napari viewer

## 4.7.3 pde.trackers.interrupts module

Module defining classes for time interrupts for trackers

The provided interrupt classes are:

ConstantInterrupts	class representing equidistantly spaced time interrupts
FixedInterrupts	class representing a list of interrupt times
LogarithmicInterrupts	class representing logarithmically spaced time interrupts
RealtimeInterrupts	class representing time interrupts spaced equidistantly in
	real time

#### class ConstantInterrupts (*dt*=1, *t\_start=None*)

## Bases: InterruptsBase

class representing equidistantly spaced time interrupts

## Parameters

- dt (float) The duration between subsequent interrupts. This is measured in simulation time units.
- **t\_start** (*float*, *optional*) The time after which the tracker becomes active. If omitted, the tracker starts recording right away. This argument can be used for an initial equilibration period during which no data is recorded.

#### copy()

return a copy of this instance

## initialize(t)

initialize the interrupt class

#### Parameters

**t** (*float*) – The starting time of the simulation

#### Returns

The first time the simulation needs to be interrupted

## **Return type**

float

### next(t)

computes the next time point

#### **Parameters**

t (*float*) – The current time point of the simulation. The returned next time point lies later than this time, so interrupts might be skipped.

### Returns

The next time point

## **Return type**

float

## class FixedInterrupts(interrupts)

```
Bases: \ \textit{InterruptsBase}
```

class representing a list of interrupt times

#### Parameters

interrupts(np.ndarray | Sequence[float])-

## copy()

return a copy of this instance

#### initialize(t)

initialize the interrupt class

## Parameters

t (float) – The starting time of the simulation

#### Returns

The first time the simulation needs to be interrupted

#### Return type

float

#### next(t)

computes the next time point

#### Parameters

t (*float*) – The current time point of the simulation. The returned next time point lies later than this time, so interrupts might be skipped.

#### Returns

The next time point

#### **Return type**

float

## class InterruptsBase

Bases: object

base class for implementing interrupts

## abstract copy()

return a copy of this instance

## Parameters

self(TInterrupt) -

#### **Return type**

TInterrupt

#### dt: float

current time difference between interrupts

Туре

float

## abstract initialize(t)

initialize the interrupt class

#### Parameters

t (float) – The starting time of the simulation

#### Returns

The first time the simulation needs to be interrupted

#### **Return type**

float

## abstract next(t)

computes the next time point

#### Parameters

t (*float*) – The current time point of the simulation. The returned next time point lies later than this time, so interrupts might be skipped.

#### Returns

The next time point

#### **Return type**

float

#### class LogarithmicInterrupts (*dt\_initial=1*, *factor=1*, *t\_start=None*)

Bases: ConstantInterrupts

class representing logarithmically spaced time interrupts

#### Parameters

- **dt\_initial** (*float*) The initial duration between subsequent interrupts. This is measured in simulation time units.
- **factor** (*float*) The factor by which the time between interrupts is increased every time. Values larger than one lead to time interrupts that are increasingly further apart.
- **t\_start** (*float*, *optional*) The time after which the tracker becomes active. If omitted, the tracker starts recording right away. This argument can be used for an initial equilibration period during which no data is recorded.

#### next(t)

computes the next time point

#### Parameters

t (float) – The current time point of the simulation. The returned next time point lies later than this time, so interrupts might be skipped.

#### Returns

The next time point

#### **Return type**

float

#### class RealtimeInterrupts (duration, dt\_initial=0.01)

Bases: ConstantInterrupts

class representing time interrupts spaced equidistantly in real time

This spacing is only achieved approximately and depends on the initial value set by *dt\_initial* and the actual variation in computation speed.

#### Parameters

- **duration** (*float* or *str*) The duration (in real seconds) that the interrupts should be spaced apart. The duration can also be given as a string, which is then parsed using the function parse\_duration().
- **dt\_initial** (*float*) The initial duration between subsequent interrupts. This is measured in simulation time units.

#### initialize(t)

initialize the interrupt class

#### Parameters

t (float) – The starting time of the simulation

#### Returns

The first time the simulation needs to be interrupted

Return type

float

#### next(t)

computes the next time point

#### **Parameters**

t (*float*) – The current time point of the simulation. The returned next time point lies later than this time, so interrupts might be skipped.

#### Returns

The next time point

**Return type** 

float

## parse\_interrupt(data)

create interrupt class from various data formats

### Parameters

**data** (str or number or *InterruptsBase*) – Data determining the interrupt class. If this is a *InterruptsBase*, it is simply returned, numbers imply *ConstantInterrupts*, a string is parsed as a time for *RealtimeInterrupts*, and lists are interpreted as *FixedInterrupts*.

#### Returns

An instance that represents the interrupt

#### Return type

InterruptsBase

## 4.7.4 pde.trackers.trackers module

Module defining classes for tracking results from simulations.

The trackers defined in this module are:

CallbackTracker	Tracker calling a function periodically
ProgressTracker	Tracker showing the progress of the simulation
PrintTracker	Tracker printing data to a stream (default: stdout)
PlotTracker	Tracker plotting data on screen, to files, or writes a movie
LivePlotTracker	PlotTracker with defaults for live plotting
DataTracker	Tracker storing custom data obtained by calling a function
SteadyStateTracker	Tracker aborting the simulation once steady state is reached
RuntimeTracker	Tracker interrupting the simulation once a duration has passed
ConsistencyTracker	Tracker interrupting the simulation when the state is not finite
MaterialConservationTracker	Tracking interrupting the simulation when material con- servation is broken

**class CallbackTracker** (*func*, *interrupts*=1, \*, *interval*=None)

Bases: TrackerBase

Tracker calling a function periodically

### Example

The callback tracker can be used to check for conditions during the simulation:

```
def check_simulation(state, time):
    if state.integral < 0:
        raise StopIteration
tracker = CallbackTracker(check_simulation, interval="0:10")</pre>
```

Adding tracker to the simulation will perform a check every 10 real time seconds. If the integral of the entire state falls below zero, the simulation will be aborted.

- **func** (*Callable*) The function to call periodically. The function signature should be (*state*) or (*state*, *time*), where *state* contains the current state as an instance of *FieldBase* and *time* is a float value indicating the current time. Note that only a view of the state is supplied, implying that a copy needs to be made if the data should be stored. The function can thus adjust the state by modifying it in-place and it can even interrupt the simulation by raising the special exception StopIteration.
- interrupts (InterruptData) Determines when the tracker interrupts the simulation. A single numbers determines an interval (measured in the simulation time unit) of regular interruption. A string is interpreted as a duration in real time assuming the format 'hh:mm:ss'. A list of numbers is taken as explicit simulation time points. More fine- grained contol is possible by passing an instance of classes defined in *interrupts*.

#### **handle** (*field*, t)

handle data supplied to this tracker

#### Parameters

- field (FieldBase) The current state of the simulation
- t (float) The associated time

Return type None

#### **class ConsistencyTracker** (*interrupts=None*, \*, *interval=None*)

#### Bases: TrackerBase

Tracker interrupting the simulation when the state is not finite

#### Parameters

**interrupts** (*InterruptData | None*) – Determines when the tracker interrupts the simulation. A single numbers determines an interval (measured in the simulation time unit) of regular interruption. A string is interpreted as a duration in real time assuming the format 'hh:mm:ss'. A list of numbers is taken as explicit simulation time points. More fine- grained contol is possible by passing an instance of classes defined in *interrupts*. The default value *None* checks for consistency approximately every (real) second.

#### **handle** (*field*, t)

handle data supplied to this tracker

#### **Parameters**

- field (FieldBase) The current state of the simulation
- t (float) The associated time

Return type

None

```
name = 'consistency'
```

**class** DataTracker(func, interrupts=1, \*, filename=None, interval=None)

Bases: CallbackTracker

Tracker storing custom data obtained by calling a function

#### Example

The data tracker can be used to gather statistics during the run

```
def get_statistics(state, time):
    return {"mean": state.data.mean(), "variance": state.data.var()}
data_tracker = DataTracker(get_statistics, interval=10)
```

Adding data\_tracker to the simulation will gather the statistics every 10 time units. After the simulation, the final result will be accessable via the *data* attribute or conveniently as a pandas from the *dataframe* attribute.

#### times

The time points at which the data is stored

Type list

#### data

The actually stored data, which is a list of the objects returned by the callback function.

Туре

list

#### Parameters

- **func** (*Callable*) The function to call periodically. The function signature should be (*state*) or (*state, time*), where *state* contains the current state as an instance of FieldBase and *time* is a float value indicating the current time. Note that only a view of the state is supplied, implying that a copy needs to be made if the data should be stored. Typical return values of the function are either a single number, a numpy array, a list of number, or a dictionary to return multiple numbers with assigned labels.
- interrupts (InterruptData) Determines when the tracker interrupts the simulation. A single numbers determines an interval (measured in the simulation time unit) of regular interruption. A string is interpreted as a duration in real time assuming the format 'hh:mm:ss'. A list of numbers is taken as explicit simulation time points. More fine- grained contol is possible by passing an instance of classes defined in *interrupts*.
- **filename** (*str*) A path to a file to which the data is written at the end of the tracking. The data format will be determined by the extension of the filename. '.pickle' indicates a python pickle file storing a tuple (*self.times, self.data*), whereas any other data format requires pandas.

#### property dataframe: pandas.DataFrame

the data in a dataframe

If *func* returns a dictionary, the keys are used as column names. Otherwise, the returned data is enumerated starting with '0'. In any case the time point at which the data was recorded is stored in the column 'time'.

#### Туре

pandas.DataFrame

#### finalize (info=None)

finalize the tracker, supplying additional information

## Parameters

**info** (*dict*) – Extra information from the simulation

## **Return type**

None

#### handle (field, t)

handle data supplied to this tracker

#### **Parameters**

- field (FieldBase) The current state of the simulation
- t (float) The associated time

#### Return type None

to\_file (filename, \*\*kwargs)
 store data in a file

The extension of the filename determines what format is being used. For instance, '.pickle' indicates a python pickle file storing a tuple (*self.times, self.data*), whereas any other data format requires pandas. Supported formats include 'csv', 'json'.

#### Parameters

- filename (*str*) Path where the data is stored
- **\*\*kwargs** Additional parameters may be supported for some formats

**class** LivePlotTracker (*interrupts='0:03'*, \*, *show=True*, *max\_fps=2*, *interval=None*, \*\*kwargs)

Bases: PlotTracker

PlotTracker with defaults for live plotting

The only difference to *PlotTracker* are the changed default values, where output is by default shown on screen and the *interval* is set something more suitable for interactive plotting. In particular, this tracker can be enabled by simply listing 'plot' as a tracker.

### Parameters

- **interrupts** (*InterruptData*) Determines when the tracker interrupts the simulation. A single numbers determines an interval (measured in the simulation time unit) of regular interruption. A string is interpreted as a duration in real time assuming the format 'hh:mm:ss'. A list of numbers is taken as explicit simulation time points. More fine- grained contol is possible by passing an instance of classes defined in *interrupts*.
- **title** (*str*) Text to show in the title. The current time point will be appended to this text, so include a space for optimal results.
- **output\_file** (*str*, *optional*) Specifies a single image file, which is updated periodically, so that the progress can be monitored (e.g. on a compute cluster)
- **output\_folder** (*str*, *optional*) Specifies a folder to which all images are written. The files will have names with increasing numbers.
- **movie\_file** (*str*, *optional*) Specifies a filename to which a movie of all the frames is written after the simulation.
- **show** (bool, optional) Determines whether the plot is shown while the simulation is running. If *False*, the files are created in the background. This option can slow down a simulation severely.
- max\_fps (float) Determines the maximal rate (frames per second) at which the plots are updated. Some plots are skipped if the tracker receives data at a higher rate. A larger value (e.g., *math.inf*) can be used to ensure every frame is drawn, which might penalizes the overall performance.
- **plot\_args** (*dict*) Extra arguments supplied to the plot call. For example, this can be used to specify axes ranges when a single panel is shown. For instance, the value {'ax\_style': {'ylim': (0, 1)}} enforces the y-axis to lie between 0 and 1.

```
name = 'plot'
```

**class** MaterialConservationTracker (*interrupts=1*, *atol=0.0001*, *rtol=0.0001*, \*, *interval=None*)

Bases: TrackerBase

Tracking interrupting the simulation when material conservation is broken

#### Parameters

• **interrupts** (*InterruptData*) – Determines when the tracker interrupts the simulation. A single numbers determines an interval (measured in the simulation time unit) of regular

interruption. A string is interpreted as a duration in real time assuming the format 'hh:mm:ss'. A list of numbers is taken as explicit simulation time points. More fine- grained contol is possible by passing an instance of classes defined in *interrupts*.

- **atol** (*float*) Absolute tolerance for amount deviations
- rtol (float) Relative tolerance for amount deviations

#### **handle** (*field*, t)

handle data supplied to this tracker

#### **Parameters**

- field (FieldBase) The current state of the simulation
- t (float) The associated time

#### **Return type**

None

initialize (field, info=None)

#### Parameters

- field (FieldBase) An example of the data that will be analyzed by the tracker
- **info** (*dict*) Extra information from the simulation

### Returns

The first time the tracker needs to handle data

Return type

float

## name = 'material\_conservation'

Bases: TrackerBase

Tracker plotting data on screen, to files, or writes a movie

This tracker can be used to create movies from simulations or to simply update a single image file on the fly (i.e. to monitor simulations running on a cluster). The default values of this tracker are chosen with regular output to a file in mind.

#### Example

To create a movie while running the simulation, you can use

```
movie_tracker = PlotTracker(interval=10, movie="my_movie.mp4")
eq.solve(..., tracker=movie_tracker)
```

This will create the file *my\_movie.mp4* during the simulation. Note that you can display the frames interactively by setting show=True.

#### **Parameters**

• **interrupts** (*InterruptData*) – Determines when the tracker interrupts the simulation. A single numbers determines an interval (measured in the simulation time unit) of regular interruption. A string is interpreted as a duration in real time assuming the format 'hh:mm:ss'. A list of numbers is taken as explicit simulation time points. More fine- grained contol is possible by passing an instance of classes defined in *interrupts*.

- **title** (*str or callable*) Title text of the figure. If this is a string, it is shown with a potential placeholder named *time* being replaced by the current simulation time. Conversely, if *title* is a function, it is called with the current state and the time as arguments. This function is expected to return a string.
- **output\_file** (*str*, *optional*) Specifies a single image file, which is updated periodically, so that the progress can be monitored (e.g. on a compute cluster)
- movie (str or Movie) Create a movie. If a filename is given, all frames are written to this file in the format deduced from the extension after the simulation ran. If a Movie is supplied, frames are appended to the instance.
- **show** (*bool*, *optional*) Determines whether the plot is shown while the simulation is running. If set to *None*, the images are only shown if neither *output\_file* nor *movie* is set, otherwise they are kept hidden. Note that showing the plot can slow down a simulation severely.
- tight\_layout (bool) Determines whether tight\_layout () is used.
- max\_fps(float) Determines the maximal rate (frames per second) at which the plots are updated in real time during the simulation. Some plots are skipped if the tracker receives data at a higher rate. A larger value (e.g., *math.inf*) can be used to ensure every frame is drawn, which might penalizes the overall performance.
- plot\_args (dict) Extra arguments supplied to the plot call. For example, this can be used to specify axes ranges when a single panel is shown. For instance, the value {'ax\_style': {'ylim': (0, 1)}} enforces the y-axis to lie between 0 and 1.

**Note:** If an instance of *Movie* is given as the *movie* argument, it can happen that the movie is not written to the file when the simulation ends. This is because, the movie could still be extended by appending frames. To write the movie to a file call its *save()* method. Beside adding frames before and after the simulation, an explicit movie object can also be used to adjust the output. For instance, the following example code creates a movie with a framerate of 15, a resolution of 200 dpi, and a bitrate of 6000 kilobits per second:

```
movie = Movie("movie.mp4", framerate=15, dpi=200, bitrate=6000)
eq.solve(..., tracker=PlotTracker(1, movie=movie))
movie.save()
```

finalize (info=None)

finalize the tracker, supplying additional information

Parameters info (dict) – Extra information from the simulation Return type

None

handle (state, t)

handle data supplied to this tracker

- **state** (FieldBase) The current state of the simulation
- t (float) The associated time

## Return type

None

#### initialize (state, info=None)

initialize the tracker with information about the simulation

#### Parameters

- state (FieldBase) An example of the data that will be analyzed by the tracker
- **info** (*dict*) Extra information from the simulation

#### Returns

The first time the tracker needs to handle data

## Return type

float

Bases: TrackerBase

Tracker printing data to a stream (default: stdout)

#### Parameters

- **interrupts** (*InterruptData*) Determines when the tracker interrupts the simulation. A single numbers determines an interval (measured in the simulation time unit) of regular interruption. A string is interpreted as a duration in real time assuming the format 'hh:mm:ss'. A list of numbers is taken as explicit simulation time points. More fine- grained contol is possible by passing an instance of classes defined in *interrupts*.
- **stream** (*IO*[*str*]) The stream used for printing

#### **handle** (*field*, t)

handle data supplied to this tracker

#### **Parameters**

- field (FieldBase) The current state of the simulation
- t (float) The associated time

#### **Return type**

None

#### name = 'print'

#### **class ProgressTracker** (*interrupts=None*, \*, *fancy=True*, *ndigits=5*, *leave=True*, *interval=None*)

### Bases: TrackerBase

Tracker showing the progress of the simulation

- **interrupts** (*InterruptData | None*) Determines when the tracker interrupts the simulation. A single numbers determines an interval (measured in the simulation time unit) of regular interruption. A string is interpreted as a duration in real time assuming the format 'hh:mm:ss'. A list of numbers is taken as explicit simulation time points. More fine- grained contol is possible by passing an instance of classes defined in *interrupts*. The default value *None* updates the progress bar approximately every (real) second.
- fancy (bool) Flag determining whether a fancy progress bar should be used in jupyter notebooks (if ipywidgets is installed)

- ndigits (*int*) The number of digits after the decimal point that are shown maximally.
- **leave** (bool) Whether to leave the progress bar after the simulation has finished (default: True)

#### finalize (info=None)

finalize the tracker, supplying additional information

#### Parameters

**info** (*dict*) – Extra information from the simulation

## **Return type**

None

## **handle** (*field*, t)

handle data supplied to this tracker

#### **Parameters**

- field (FieldBase) The current state of the simulation
- t (float) The associated time

#### **Return type**

None

## initialize (field, info=None)

initialize the tracker with information about the simulation

## Parameters

- field (FieldBase) An example of the data that will be analyzed by the tracker
- info (dict) Extra information from the simulation

#### Returns

The first time the tracker needs to handle data

**Return type** 

float

#### name = 'progress'

**class RuntimeTracker** (*max\_runtime*, *interrupts=1*, \*, *interval=None*)

Bases: TrackerBase

Tracker interrupting the simulation once a duration has passed

#### Parameters

- **max\_runtime** (*float* or *str*) The maximal runtime of the simulation. If the runtime is exceeded, the simulation is interrupted. Values can be either given as a number (interpreted as seconds) or as a string, which is then parsed using the function *parse\_duration()*.
- interrupts (InterruptData) Determines when the tracker interrupts the simulation. A single numbers determines an interval (measured in the simulation time unit) of regular interruption. A string is interpreted as a duration in real time assuming the format 'hh:mm:ss'. A list of numbers is taken as explicit simulation time points. More fine- grained contol is possible by passing an instance of classes defined in *interrupts*.

#### **handle** (*field*, t)

handle data supplied to this tracker

- field (FieldBase) The current state of the simulation
- t (float) The associated time

## Return type

None

initialize (field, info=None)

#### Parameters

- field (FieldBase) An example of the data that will be analyzed by the tracker
- **info** (*dict*) Extra information from the simulation

#### Returns

The first time the tracker needs to handle data

## **Return type**

float

**class SteadyStateTracker** (*interrupts=None*, *atol=1e-08*, *rtol=1e-05*, \*, *progress=False*, *evolution\_rate=None*, *interval=None*)

## Bases: TrackerBase

Tracker aborting the simulation once steady state is reached

Steady state is obtained when the state does not change anymore, i.e., when the evolution rate is close to zero. If the argument *evolution\_rate* is specified, it is used to calculate the evolution rate directly. If it is omitted, the evolution rate is estaimted by comparing the current state *cur* to the state *prev* at the previous time step. In both cases, convergence is assumed when the absolute value of the evolution rate falls below atol + rtol \* cur for all points. Here, *atol* and *rtol* denote absolute and relative tolerances, respectively.

#### Parameters

- **interrupts** (*InterruptData | None*) Determines when the tracker interrupts the simulation. A single numbers determines an interval (measured in the simulation time unit) of regular interruption. A string is interpreted as a duration in real time assuming the format 'hh:mm:ss'. A list of numbers is taken as explicit simulation time points. More fine- grained contol is possible by passing an instance of classes defined in *interrupts*. The default value *None* checks for the steady state approximately every (real) second.
- **atol** (*float*) Absolute tolerance that must be reached to abort the simulation
- rtol (float) Relative tolerance that must be reached to abort the simulation
- **progress** (bool) Flag indicating whether the progress towards convergence is shown graphically during the simulation
- **evolution\_rate** (*callable*) Function to evaluate the current evolution rate. If omitted, the evolution rate is estimate from the change in the state variable, which can be less accurate. A suitable form of the function is returned by *eq.make\_pde\_rhs(state)* when *eq* is the PDE class.

#### **handle** (*field*, t)

handle data supplied to this tracker

- **field** (FieldBase) The current state of the simulation
- t (float) The associated time

Return type None	
<pre>name = 'steady_state'</pre>	
<pre>progress_bar_format = 'Convergence: {pe [{elapsed}&lt;{remaining}]'</pre>	ercentage:3.0f}% {bar}
determines the format of the progress bar shown when	$n \ progress = True$

4.8 pde.visualization package

Functions and classes for visualizing simulations.

movies	Functions for creating movies of simulation results
plotting	Functions and classes for plotting simulation data

## 4.8.1 pde.visualization.movies module

Functions for creating movies of simulation results

Movie	Class for creating movies from matplotlib figures using ffmpeg
movie_scalar	produce a movie for a simulation of a scalar field
movie_multiple	produce a movie for a simulation with n components
movie	produce a movie by simply plotting each frame

**class** Movie (filename, framerate=30, dpi=None, \*\*kwargs)

Bases: object

Class for creating movies from matplotlib figures using ffmpeg

Note: Internally, this class uses matplotlib.animation.FFMpegWriter. Note that the *ffmpeg* program needs to be installed in a system path, so that *matplotlib* can find it.

**Warning:** The movie is only fully written after the *save()* method has been called. To aid with this, it is best practice to use a contextmanager:

```
with Movie("output.mp4") as movie:
    movie.add_figure()
```

- **filename** (*str*) The filename where the movie is stored. The suffix of this path also determines the default movie codec.
- **framerate** (*float*) The number of frames per second, which determines how fast the movie will appear to run.
- **dpi** (*float*) The resolution of the resulting movie

• **\*\*kwargs** - Additional parameters are used to initialize matplotlib.animation. FFMpegWriter. Here, we can for instance set the bit rate of the resulting video using the *bitrate* parameter.

### add\_figure (fig=None)

adds the figure *fig* as a frame to the current movie

#### Parameters

fig (Figure) - The plot figure that is added to the movie

#### classmethod is\_available()

check whether the movie infrastructure is available

#### Returns

True if movies can be created

### **Return type**

bool

#### save()

convert the recorded images to a movie using ffmpeg

**movie** (*storage*, *filename*, \*, *progress=True*, *show\_time=True*, *plot\_args=None*, *movie\_args=None*)

produce a movie by simply plotting each frame

#### Parameters

- **storage** (*StorageBase*) The storage instance that contains all the data for the movie
- **filename** (*str*) The filename to which the movie is written. The extension determines the format used.
- progress (bool) Flag determining whether the progress of making the movie is shown.
- **show\_time** (bool) Whether to show the simulation time in the movie
- plot\_args (dict) Additional arguments for the function plotting the state
- movie\_args (dict) Additional arguments for *Movie*. For example, this can be used to set the resolution (*dpi*), the framerate (*framerate*), and the bitrate (*bitrate*).

#### **Return type**

None

**movie\_multiple** (storage, filename, quantities=None, scale='automatic', progress=True)

produce a movie for a simulation with n components

- **storage** (*StorageBase*) The storage instance that contains all the data for the movie
- **filename** (*str*) The filename to which the movie is written. The extension determines the format used.
- **quantities** A 2d list of quantities that are shown in a rectangular arrangement. If *quantities* is a simple list, the panels will be rendered as a single row. Each panel is defined by a dictionary, where the mandatory item 'source' defines what is being shown. Here, an integer specifies the component that is extracted from the field while a function is evaluate with the full state as an input and the result is shown. Additional items in the dictionary can be 'title' (setting the title of the panel), 'scale' (defining the color range shown; these are typically two numbers defining the lower and upper bound, but if only one is given the range [0, scale] is assumed), and 'cmap' (defining the colormap being used).

- scale (*str*, *float*, *tuple of float*) Flag determining how the range of the color scale is determined. In the simplest case a tuple of numbers marks the lower and upper end of the scalar values that will be shown. If only a single number is supplied, the range starts at zero and ends at the given number. Additionally, the special value 'automatic' determines the range from the range of scalar values.
- progress (bool) Flag determining whether the progress of making the movie is shown.

#### **Return type**

None

**movie\_scalar** (*storage*, *filename*, *scale='automatic'*, *extras=None*, *progress=True*, *tight=False*, *show=True*) produce a movie for a simulation of a scalar field

#### **Parameters**

- **storage** (*StorageBase*) The storage instance that contains all the data for the movie
- **filename** (*str*) The filename to which the movie is written. The extension determines the format used.
- scale (*str*, *float*, *tuple of float*) Flag determining how the range of the color scale is determined. In the simplest case a tuple of numbers marks the lower and upper end of the scalar values that will be shown. If only a single number is supplied, the range starts at zero and ends at the given number. Additionally, the special value 'automatic' determines the range from the range of scalar values.
- **extras** (*dict*, *optional*) Additional functions that are evaluated and shown for each time step. The key of the dictionary is used as a panel title.
- progress (bool) Flag determining whether the progress of making the movie is shown.
- tight (*bool*) Whether to call matplotlib.pyplot.tight\_layout(). This affects the layout of all plot elements.
- **show** (bool) Flag determining whether images are shown during making the movie

## Return type

None

## 4.8.2 pde.visualization.plotting module

Functions and classes for plotting simulation data

ScalarFieldPlot	class managing compound plots of scalar fields
plot_magnitudes	plot spatially averaged quantities as a function of time
plot_kymograph	plots a single kymograph from stored data
plot_kymographs	plots kymographs for all fields stored in storage
plot_interactive	plots stored data interactively using the napari viewer

class ScalarFieldPlot (fields, quantities=None, scale='automatic', fig=None, title=None, tight=False, show=True)

Bases: object

class managing compound plots of scalar fields

#### Parameters

• **fields** (*FieldBase*) – Collection of fields

- **quantities** A 2d list of quantities that are shown in a rectangular arrangement. If *quantities* is a simple list, the panels will be rendered as a single row. Each panel is defined by a dictionary, where the mandatory item 'source' defines what is being shown. Here, an integer specifies the component that is extracted from the field while a function is evaluate with the full state as an input and the result is shown. Additional items in the dictionary can be 'title' (setting the title of the panel), 'scale' (defining the color range shown; these are typically two numbers defining the lower and upper bound, but if only one is given the range [0, scale] is assumed), and 'cmap' (defining the colormap being used).
- **scale** (*str*, *float*, *tuple of float*) Flag determining how the range of the color scale is determined. In the simplest case a tuple of numbers marks the lower and upper end of the scalar values that will be shown. If only a single number is supplied, the range starts at zero and ends at the given number. Additionally, the special value 'automatic' determines the range from the range of scalar values.
- ( (fig) class:matplotlib.figure.Figure): Figure to be used for plotting. If `None, a new figure is created.
- **title** (*str*) Title of the plot.
- tight (bool) Whether to call matplotlib.pyplot.tight\_layout(). This affects the layout of all plot elements.
- **show** (bool) Flag determining whether to show a plot. If *False*, the plot is kept in the background, which can be useful if it only needs to be written to a file.

**classmethod from\_storage** (*storage*, *quantities=None*, *scale='automatic'*, *tight=False*, *show=True*) create ScalarFieldPlot from storage

#### Parameters

- **storage** (*StorageBase*) Instance of the storage class that contains the data
- **quantities** A 2d list of quantities that are shown in a rectangular arrangement. If *quantities* is a simple list, the panels will be rendered as a single row. Each panel is defined by a dictionary, where the mandatory item 'source' defines what is being shown. Here, an integer specifies the component that is extracted from the field while a function is evaluate with the full state as an input and the result is shown. Additional items in the dictionary can be 'title' (setting the title of the panel), 'scale' (defining the color range shown; these are typically two numbers defining the lower and upper bound, but if only one is given the range [0, scale] is assumed), and 'cmap' (defining the colormap being used).
- **scale** (*str*, *float*, *tuple of float*) Flag determining how the range of the color scale is determined. In the simplest case a tuple of numbers marks the lower and upper end of the scalar values that will be shown. If only a single number is supplied, the range starts at zero and ends at the given number. Additionally, the special value 'automatic' determines the range from the range of scalar values.
- **tight** (*bool*) Whether to call matplotlib.pyplot.tight\_layout(). This affects the layout of all plot elements.
- **show** (bool) Flag determining whether to show a plot. If *False*, the plot is kept in the background.

#### Returns

ScalarFieldPlot

#### Return type

ScalarFieldPlot

#### make\_movie (storage, filename, progress=True)

make a movie from the data stored in storage

#### **Parameters**

- **storage** (*StorageBase*) The storage instance that contains all the data for the movie
- **filename** (*str*) The filename to which the movie is written. The extension determines the format used.
- progress (bool) Flag determining whether the progress of making the movie is shown.

## Return type

None

## savefig(path, \*\*kwargs)

save plot to file

#### **Parameters**

- **path** (*str*) The path to the file where the image is written. The file extension determines the image format
- **\*\*kwargs** Additional arguments are forwarded to matplotlib.figure.Figure. savefig().

#### update (fields, title=None)

update the plot with the given fields

#### Parameters

- **fields** (FieldBase) The field or field collection of which the defined quantities are shown.
- title (str, optional) The title of this view. If None, the current title is not changed.

#### **Return type**

None

#### extract\_field(fields, source=None, check\_rank=None)

Extracts a single field from a possible collection.

#### Parameters

- fields (FieldBase) The field from which data is extracted
- **source** (*int* or *callable*, *optional*) Determines how a field is extracted from *fields*. If *None*, *fields* is passed as is, assuming it is already a scalar field. This works for the simple, standard case where only a single *ScalarField* is treated. Alternatively, *source* can be an integer, indicating which field is extracted from an instance of FieldCollection. Lastly, *source* can be a function that takes *fields* as an argument and returns the desired field.
- **check\_rank** (*int*, *optional*) Can be given to check whether the extracted field has the correct rank (0 = ScalarField, 1 = VectorField, ...).

#### Returns

The extracted field

#### **Return type**

DataFieldBase

#### plot\_interactive (storage, time\_scaling='exact', viewer\_args=None, \*\*kwargs)

plots stored data interactively using the napari viewer

#### Parameters

- **storage** (StorageBase) The storage instance that contains all the data
- time\_scaling (*str*) Defines how the time axis is scaled. Possible options are "exact" (the actual time points are used), or "scaled" (the axis is scaled so that it has similar dimension to the spatial axes). Note that the spatial axes will never be scaled.
- **viewer\_args** (*dict*) Arguments passed to napari.viewer.Viewer to affect the viewer.
- **\*\*kwargs** Extra arguments passed to the plotting function

plots a single kymograph from stored data

The kymograph shows line data stacked along time. Consequently, the resulting image shows space along the horizontal axis and time along the vertical axis.

- **storage** (StorageBase) The storage instance that contains all the data
- **field\_index** (*int*) An index to choose a single field out of many in a collection stored in *storage*. This option should not be used if only a single field is stored in a collection.
- **scalar** (*str*) The method for extracting scalars as described in DataFieldBase. to\_scalar().
- **extract** (*str*) The method used for extracting the line data. See the docstring of the grid method *get\_line\_data* to find supported values.
- **colorbar** (*bool*) Whether to show a colorbar or not
- **transpose** (bool) Determines whether the transpose of the data should is plotted
- **title** (*str*) Title of the plot. If omitted, the title might be chosen automatically.
- filename (*str*, *optional*) If given, the plot is written to the specified file.
- **action** (*str*) Decides what to do with the final figure. If the argument is set to *show*, matplotlib.pyplot.show() will be called to show the plot. If the value is *none*, the figure will be created, but not necessarily shown. The value *close* closes the figure, after saving it to a file when *filename* is given. The default value *auto* implies that the plot is shown if it is not a nested plot call.
- **ax\_style** (*dict*) Dictionary with properties that will be changed on the axis after the plot has been drawn by calling matplotlib.pyplot.setp(). A special item i this dictionary is *use\_offset*, which is flag that can be used to control whether offset are shown along the axes of the plot.
- **fig\_style** (*dict*) Dictionary with properties that will be changed on the figure after the plot has been drawn by calling matplotlib.pyplot.setp(). For instance, using fig\_style={'dpi': 200} increases the resolution of the figure.
- **ax** (matplotlib.axes.Axes) Figure axes to be used for plotting. The special value "create" creates a new figure, while "reuse" attempts to reuse an existing figure, which is the default.
- \*\*kwargs Additional keyword arguments are passed to matplotlib.pyplot. imshow().

## Returns

The reference to the plot

### **Return type**

#### PlotReference

plots kymographs for all fields stored in storage

The kymograph shows line data stacked along time. Consequently, the resulting image shows space along the horizontal axis and time along the vertical axis.

#### **Parameters**

- storage (StorageBase) The storage instance that contains all the data
- **scalar** (*str*) The method for extracting scalars as described in DataFieldBase. to\_scalar().
- **extract** (*str*) The method used for extracting the line data. See the docstring of the grid method *get\_line\_data* to find supported values.
- colorbar (bool) Whether to show a colorbar or not
- **transpose** (bool) Determines whether the transpose of the data should is plotted
- **resize\_fig** (bool) Whether to resize the figure to adjust to the number of panels
- **title** (*str*) Title of the plot. If omitted, the title might be chosen automatically. This is shown above all panels.
- **constrained\_layout** (*bool*) Whether to use *constrained\_layout* in matplotlib. pyplot.figure() call to create a figure. This affects the layout of all plot elements. Generally, spacing might be better with this flag enabled, but it can also lead to problems when plotting multiple plots successively, e.g., when creating a movie.
- **filename** (*str*, *optional*) If given, the figure is written to the specified file.
- **action** (*str*) Decides what to do with the final figure. If the argument is set to *show*, matplotlib.pyplot.show() will be called to show the plot. If the value is *none*, the figure will be created, but not necessarily shown. The value *close* closes the figure, after saving it to a file when *filename* is given. The default value *auto* implies that the plot is shown if it is not a nested plot call.
- **fig\_style** (*dict*) Dictionary with properties that will be changed on the figure after the plot has been drawn by calling matplotlib.pyplot.setp(). For instance, using fig\_style={'dpi': 200} increases the resolution of the figure.
- **fig** (matplotlib.figures.Figure) Figure that is used for plotting. If omitted, a new figure is created.
- **\*\*kwargs** Additional keyword arguments are passed to the calls to matplotlib. pyplot.imshow().

#### Returns

The references to all plots

## **Return type**

list of *PlotReference* 

plot\_magnitudes (storage, quantities=None, \*args, title=None, filename=None, action='auto', ax\_style=None, fig\_style=None, ax=None, \*\*kwargs)

plot spatially averaged quantities as a function of time

For scalar fields, the default is to plot the average value while the averaged norm is plotted for vector fields.

#### Parameters

- **storage** (StorageBase) Instance of *StorageBase* that contains the simulation data that will be plotted
- **quantities** A 2d list of quantities that are shown in a rectangular arrangement. If *quan-tities* is a simple list, the panels will be rendered as a single row. Each panel is defined by a dictionary, where the mandatory item 'source' defines what is being shown. Here, an integer specifies the component that is extracted from the field while a function is evaluate with the full state as an input and the result is shown. Additional items in the dictionary can be 'title' (setting the title of the panel), 'scale' (defining the color range shown; these are typically two numbers defining the lower and upper bound, but if only one is given the range [0, scale] is assumed), and 'cmap' (defining the colormap being used).
- **title** (*str*) Title of the plot. If omitted, the title might be chosen automatically.
- filename (*str*, *optional*) If given, the plot is written to the specified file.
- **action** (*str*) Decides what to do with the final figure. If the argument is set to *show*, matplotlib.pyplot.show() will be called to show the plot. If the value is *none*, the figure will be created, but not necessarily shown. The value *close* closes the figure, after saving it to a file when *filename* is given. The default value *auto* implies that the plot is shown if it is not a nested plot call.
- **ax\_style** (*dict*) Dictionary with properties that will be changed on the axis after the plot has been drawn by calling matplotlib.pyplot.setp(). A special item i this dictionary is *use\_offset*, which is flag that can be used to control whether offset are shown along the axes of the plot.
- **fig\_style** (*dict*) Dictionary with properties that will be changed on the figure after the plot has been drawn by calling matplotlib.pyplot.setp(). For instance, using fig\_style={'dpi': 200} increases the resolution of the figure.
- **ax** (matplotlib.axes.Axes) Figure axes to be used for plotting. The special value "create" creates a new figure, while "reuse" attempts to reuse an existing figure, which is the default.
- **\*\*kwargs** All remaining parameters are forwarded to the *ax.plot* method

#### Returns

The reference to the plot

#### **Return type**

PlotReference

#### Indices and tables

- genindex
- modindex
- search

# **PYTHON MODULE INDEX**

# f

pde.fields,59
pde.fields.base,60
pde.fields.collection,74
pde.fields.scalar,80
pde.fields.tensorial,85
pde.fields.vectorial,89

# g

```
pde.grids,94
pde.grids.base,144
pde.grids.boundaries,95
pde.grids.boundaries.axes,97
pde.grids.boundaries.axis,99
pde.grids.boundaries.local, 102
pde.grids.cartesian,157
pde.grids.coordinates, 127
pde.grids.coordinates.base,127
pde.grids.coordinates.bipolar,130
pde.grids.coordinates.bispherical, 130
pde.grids.coordinates.cartesian, 131
pde.grids.coordinates.cylindrical, 131
pde.grids.coordinates.polar, 132
pde.grids.coordinates.spherical, 132
pde.grids.cylindrical, 161
pde.grids.operators, 133
pde.grids.operators.cartesian, 133
pde.grids.operators.common, 135
pde.grids.operators.cylindrical_sym, 136
pde.grids.operators.polar_sym, 139
pde.grids.operators.spherical_sym, 141
pde.grids.spherical, 165
```

# р

```
pde, 59
pde.pdes, 170
pde.pdes.allen_cahn, 170
pde.pdes.base, 171
pde.pdes.cahn_hilliard, 175
pde.pdes.diffusion, 176
pde.pdes.kpz_interface, 177
pde.pdes.kuramoto_sivashinsky, 178
```

pde.pdes.laplace, 180
pde.pdes.pde, 181
pde.pdes.swift\_hohenberg, 183
pde.pdes.wave, 184

## S

pde.solvers, 185 pde.solvers.adams\_bashforth, 188 pde.solvers.base, 188 pde.solvers.controller, 190 pde.solvers.crank\_nicolson, 192 pde.solvers.explicit, 192 pde.solvers.explicit\_mpi, 193 pde.solvers.scipy, 195 pde.storage, 195 pde.storage.base, 196 pde.storage.file, 201 pde.storage.memory, 202 pde.storage.movie, 204

## t

```
pde.tools, 207
pde.tools.cache, 207
pde.tools.config,213
pde.tools.cuboid,214
pde.tools.docstrings, 217
pde.tools.expressions, 217
pde.tools.ffmpeg, 223
pde.tools.math, 225
pde.tools.misc, 226
pde.tools.mpi, 229
pde.tools.numba,231
pde.tools.output, 233
pde.tools.parameters, 234
pde.tools.parse_duration, 237
pde.tools.plotting, 237
pde.tools.spectral, 242
pde.tools.typing, 243
pde.trackers, 243
pde.trackers.base, 244
pde.trackers.interactive, 247
```

pde.trackers.interrupts, 249
pde.trackers.trackers, 253

## ۷

pde.visualization, 262
pde.visualization.movies, 262
pde.visualization.plotting, 264

# INDEX

## A

AdamsBashforthSolver (class in pde.solvers), 185 AdamsBashforthSolver (class in pde.solvers.adams\_bashforth), 188 AdaptiveSolverBase (class in pde.solvers.base), 188 add figure() (Movie method), 263 add\_scaled\_colorbar() (in module pde.tools.plotting), 239 AdjacentEvaluator (class in pde.tools.typing), 243 AllenCahnPDE (class in pde.pdes.allen cahn), 170 append() (FieldCollection method), 75 append() (StorageBase method), 196 apply() (FieldBase method), 69 apply() (StorageBase method), 196 apply\_operator() (DataFieldBase method), 60 asanyarray\_flags() (in module pde.tools.cuboid), 216 assert\_field\_compatible() (FieldBase method), 70 assert\_field\_compatible() (FieldCollection method), 75 assert\_grid\_compatible() (GridBase method), 144 attributes (FieldBase property), 70 attributes (FieldCollection property), 75 attributes\_serialized (FieldBase property), 70 attributes\_serialized (FieldCollection property), 75 average (DataFieldBase property), 60 averages (FieldCollection property), 75 ax (PlotReference attribute), 238 axes (BipolarCoordinates attribute), 130 axes (BisphericalCoordinates attribute), 130 axes (CoordinatesBase attribute), 127 axes (CylindricalCoordinates attribute), 131 axes (GridBase attribute), 145 axes (PolarCoordinates attribute), 132 axes (SphericalCoordinates attribute), 132 axes\_bounds (GridBase property), 145 axes\_coords (GridBase property), 145 axes symmetric (GridBase attribute), 145 axis (BoundaryAxisBase property), 99

axis\_coord (BCBase property), 103

## В

BasicOutput (class in pde.tools.output), 233 BasicPlottingContext (class in pde.tools.plotting), 238 basis\_rotation() (CoordinatesBase method), 127 BCBase (class in pde.grids.boundaries.local), 103 BCDataError, 107 BipolarCoordinates (class in pde.grids.coordinates.bipolar), 130 BisphericalCoordinates (class in pde.grids.coordinates.bispherical), 130 bits\_per\_channel (FFmpegFormat attribute), 223 boundaries (Boundaries property), 97 Boundaries (class in pde.grids.boundaries.axes), 97 boundary\_names (CartesianGrid attribute), 157 boundary\_names (CylindricalSymGrid attribute), 162 boundary names (GridBase attribute), 145 boundary\_names (SphericalSymGridBase attribute), 167 BoundaryAxisBase (class in pde.grids.boundaries.axis), 99 BoundaryPair (class in pde.grids.boundaries.axis), 101 BoundaryPeriodic (class in pde.grids.boundaries.axis), 101 bounds (Cuboid property), 214 bounds (SmoothData1D property), 225 buffer() (Cuboid method), 214 bytes\_per\_channel (FFmpegFormat property), 223

# С

c (CylindricalSymGrid attribute), 162 c (GridBase attribute), 145 c (PolarSymGrid attribute), 166 c (SphericalSymGrid attribute), 166 cache\_rhs (PDEBase attribute), 172 cached\_method (class in pde.tools.cache), 208 cached\_property (class in pde.tools.cache), 209 CahnHilliardPDE (class in pde.pdes.cahn\_hilliard), 175 CallbackTracker (class in pde.trackers.trackers), 253 CartesianCoordinates (class pde.grids.coordinates.cartesian), 131 CartesianGrid (class in pde.grids.cartesian), 157 cell\_coords (GridBase attribute), 145 cell volume() (CoordinatesBase method), 127 cell volume data (CartesianGrid property), 157 cell volume data (CylindricalSymGrid attribute), 162 cell\_volume\_data (GridBase attribute), 145 cell\_volume\_data (SphericalSymGridBase attribute), 167 cell\_volumes (GridBase attribute), 145 CellVolume (class in pde.tools.typing), 243 centroid (Cuboid property), 215 channels (FFmpegFormat attribute), 223 check\_implementation (PDEBase attribute), 172 check\_length() (DictFiniteCapacity method), 207 check package version() module (in pde.tools.config), 213 check rhs consistency() (PDEBase method), 172 check value rank() (BCBase method), 104 check\_value\_rank() (Boundaries method), 97 check value rank() (BoundaryPair method), 101 check value rank() (BoundaryPeriodic method), 101 class\_type (OnlineStatistics attribute), 225 classmethod() (hybridmethod method), 228 classproperty (class in pde.tools.misc), 226 clear() (FileStorage method), 201 clear() (MemoryStorage method), 202 clear() (MovieStorage method), 205 clear() (StorageBase method), 196 close() (FileStorage method), 201 close() (JupyterPlottingContext method), 238 close() (MovieStorage method), 205 close() (NapariViewer method), 248 close() (PlottingContextBase method), 239 codec (FFmpegFormat attribute), 224 compatible\_with() (GridBase method), 145 complex (ExpressionBase property), 218 complex valued (PDEBase attribute), 172 Config (class in pde.tools.config), 213 conjugate() (FieldBase method), 70 ConsistencyTracker (class in pde.trackers.trackers), 254 constant (ExpressionBase property), 218 ConstantInterrupts (class in pde.trackers.interrupts), 249 ConstBC1stOrderBase (class in pde.grids.boundaries.local), 107 ConstBC2ndOrderBase (class in pde.grids.boundaries.local), 109 ConstBCBase (*class in pde.grids.boundaries.local*), 111

in contains\_point() (Cuboid method), 215 contains\_point() (GridBase method), 146 Controller (class in pde.solvers), 185 Controller (class in pde.solvers.controller), 190 ConvergenceError, 189 convert () (Parameter method), 235 coordinate arrays (GridBase attribute), 146 coordinate constraints (CylindricalSymGrid attribute), 162 coordinate\_constraints (GridBase attribute), 146 coordinate\_constraints (PolarSymGrid attribute), 166 coordinate\_constraints (SphericalSymGrid attribute), 166 coordinate\_limits (BipolarCoordinates attribute), 130 (BisphericalCoordinates coordinate\_limits attribute), 130 coordinate limits (CoordinatesBase attribute), 128 coordinate limits (*CylindricalCoordinates* tribute), 131 coordinate limits (PolarCoordinates attribute), 132 coordinate limits (SphericalCoordinates attribute), 132 CoordinatesBase (class in pde.grids.coordinates.base), 127 copy() (BCBase method), 104 copy() (Boundaries method), 98 copy () (BoundaryAxisBase method), 99 copy() (BoundaryPeriodic method), 102 copy() (ConstantInterrupts method), 249 copy() (ConstBCBase method), 111 copy() (Cuboid method), 215 copy() (DataFieldBase method), 61 copy() (ExpressionBC method), 115 copy () (FieldBase method), 70 copy() (FieldCollection method), 75 copy() (FixedInterrupts method), 250 copy() (GridBase method), 146 copy() (InterruptsBase method), 250 copy() (MixedBC method), 120 copy() (ScalarExpression method), 220 copy() (StorageBase method), 196 copy() (UserBC method), 125 corners (Cuboid property), 215 count (OnlineStatistics attribute), 225 Counter (class in pde.tools.numba), 231 CrankNicolsonSolver (class in pde.solvers), 186 CrankNicolsonSolver (class in pde.solvers.crank\_nicolson), 192 cuboid (CartesianGrid attribute), 157 Cuboid (class in pde.tools.cuboid), 214 CurvatureBC (class in pde.grids.boundaries.local), 112

CylindricalCoordinates (class in pde.grids.coordinates.cylindrical), 131 CylindricalSymGrid (class in pde.grids.cylindrical), 161

# D

data (DataTracker attribute), 254 data (FieldBase property), 71 data (FileStorage property), 202 data (MovieStorage property), 205 data (StorageBase attribute), 197 data\_from\_frame() (FFmpegFormat method), 224 data\_shape (DataFieldBase property), 61 data\_shape (StorageBase property), 197 data\_to\_frame() (FFmpegFormat method), 224 DataFieldBase (class in pde. fields. base), 60 dataframe (DataTracker property), 255 DataTracker (class in pde.trackers.trackers), 254 decorator\_arguments() (in module pde.tools.misc), 227 default\_capacity (DictFiniteCapacity attribute), 207 deleter() (classproperty method), 227 depends on () (ExpressionBase method), 218 DeprecatedParameter (class pde.tools.parameters), 234 derivative() (SmoothData1D method), 225 derivatives (ScalarExpression attribute), 220 derivatives (TensorExpression attribute), 221 diagnostics (Controller attribute), 186, 191 diagnostics (PDEBase attribute), 172 diagonal (Cuboid property), 215 DictFiniteCapacity (class in pde.tools.cache), 207 difference\_vector() (CartesianGrid method), 157 difference\_vector() (CylindricalSymGrid method), 162 difference\_vector() (GridBase method), 146 difference\_vector\_real() (GridBase method), 146 differentiate() (ScalarExpression method), 220 differentiate() (TensorExpression method), 221 DiffusionPDE (class in pde.pdes.diffusion), 176 dim (BipolarCoordinates attribute), 130 dim (BisphericalCoordinates attribute), 131 dim (CoordinatesBase attribute), 128 dim (Cuboid property), 215 dim (CylindricalCoordinates attribute), 131 dim (GridBase property), 147 dim (PolarCoordinates attribute), 132 dim (SphericalCoordinates attribute), 132 DimensionError, 130 DirichletBC (class in pde.grids.boundaries.local), 113 disable\_interactive() module (in pde.tools.plotting), 239 discretization (GridBase property), 147

- distance() (CoordinatesBase method), 128
- distance() (GridBase method), 147
- distance\_real() (GridBase method), 147
- divergence() (Tensor2Field method), 85
  divergence() (VectorField method), 89
- DomainError, 144
- dot () (*Tensor2Field method*), 86
- dot () (VectorField method), 90
- dt (InterruptsBase attribute), 250
- dt\_default (SolverBase attribute), 189
- dt\_max (AdaptiveSolverBase attribute), 189
- dt\_min (AdaptiveSolverBase attribute), 189
- dtype (FFmpegFormat attribute), 224
- dtype (FieldBase property), 71
- dtype (StorageBase property), 197

## Е

element (PlotReference attribute), 238 end\_writing() (FileStorage method), 202 in end\_writing() (MovieStorage method), 205 end\_writing() (StorageBase method), 197 ensure\_directory\_exists() module (in pde.tools.misc), 227 environment variable PYTHONPATH, 5 environment() (in module pde.tools.config), 213 estimate\_computation\_speed() (in module pde.tools.misc), 227 evaluate() (in module pde.tools.expressions), 221 evolution\_rate() (AllenCahnPDE method), 171 evolution\_rate() (CahnHilliardPDE method), 176 evolution\_rate() (DiffusionPDE method), 177 evolution\_rate() (KPZInterfacePDE method), 178 (KuramotoSivashinskyPDE evolution\_rate() method), 179 evolution\_rate() (PDE method), 182 evolution rate() (PDEBase method), 172 evolution\_rate() (SwiftHohenbergPDE method), 183 evolution rate() (WavePDE method), 184 explicit time dependence (AllenCahnPDE attribute), 171 explicit\_time\_dependence (CahnHilliardPDE attribute), 176 (DiffusionPDE explicit\_time\_dependence attribute), 177 explicit\_time\_dependence (KPZInterfacePDE attribute), 178 explicit\_time\_dependence (KuramotoSivashinskyPDE attribute), 179

explicit_time_dependence (PDEBase attribute),	FixedInterrupts (class in pde.trackers.interrupts),	
173	250	
explicit_time_dependence (SwiftHohenbergPDE	<pre>flat_idx() (in module pde.tools.numba), 231</pre>	
attribute), 183	flip_sign (BoundaryPeriodic property), 102	
<pre>explicit_time_dependence (WavePDE attribute),</pre>	fluctuations (DataFieldBase property), 61	
184	formats (in module pde.tools.ffmpeg), 224	
ExplicitMPISolver (class in	from_bounds() (CartesianGrid class method), 158	
pde.solvers.explicit_mpi), 193	from_bounds() (Cuboid class method), 215	
ExplicitSolver (class in pde.solvers), 187	<pre>from_bounds() (CylindricalSymGrid class method),</pre>	
ExplicitSolver (class in pde.solvers.explicit), 192	163	
<pre>expr_prod() (in module pde.pdes.base), 175</pre>	from_bounds() ( <i>GridBase class method</i> ), 147	
expression (AllenCahnPDE property), 171	<pre>from_bounds() (SphericalSymGridBase class method),</pre>	
expression (CahnHilliardPDE property), 176	167	
expression ( <i>DiffusionPDE property</i> ), 177	<pre>from_centerpoint() (Cuboid class method), 215</pre>	
expression (ExpressionBase property), 219	<pre>from_collection() (MemoryStorage class method),</pre>	
expression (KPZInterfacePDE property), 178	203	
expression (KuramotoSivashinskyPDE property), 179	from_data() ( <i>BCBase class method</i> ), 104	
expression (SwiftHohenbergPDE property), 183	from_data() (Boundaries class method), 98	
ExpressionBase (class in pde.tools.expressions), 218	from_data() ( <i>BoundaryPair class method</i> ), 101	
ExpressionBC (class in pde.grids.boundaries.local),	<pre>from_data() (FieldCollection class method), 76</pre>	
114	from_data() ( <i>TrackerBase class method</i> ), 245	
ExpressionDerivativeBC (class in	from_data() ( <i>TrackerCollection class method</i> ), 246	
pde.grids.boundaries.local), 116	from_dict()( <i>BCBase class method</i> ), 104	
ExpressionMixedBC (class in	<pre>from_expression() (ScalarField class method), 80</pre>	
pde.grids.boundaries.local), 117	<pre>from_expression() (Tensor2Field class method), 86</pre>	
expressions (PDE property), 182	<pre>from_expression() (VectorField class method), 90</pre>	
expressions (WavePDE property), 184	<pre>from_fields() (MemoryStorage class method), 203</pre>	
ExpressionValueBC (class in	<pre>from_file() (FieldBase class method), 71</pre>	
pde.grids.boundaries.local), 118	<pre>from_image() (ScalarField class method), 81</pre>	
<pre>extract_field() (in module</pre>	from_name() (SolverBase class method), 190	
pde.visualization.plotting), 266	<pre>from_points() (Cuboid class method), 215</pre>	
<pre>extract_field() (StorageBase method), 197</pre>	<pre>from_scalar_expressions() (FieldCollection</pre>	
<pre>extract_time_range() (StorageBase method), 197</pre>	class method), 76	
_	<pre>from_scalars() (VectorField class method), 91</pre>	
F	<pre>from_state() (CartesianGrid class method), 158</pre>	
factory (OperatorInfo attribute), 156	<pre>from_state() (CylindricalSymGrid class method), 163</pre>	
EEmpogEormat (class in nde tools ffmnee) 223	from state() (DataFieldBase class method). 61	

FFmpegFormat (class in pde.tools.ffmpeg), 223
FieldBase (class in pde.fields.base), 69
FieldCollection (class in pde.fields.collection), 74
fields (FieldCollection property), 76
fig (PlottingContextBase attribute), 239
FileStorage (class in pde.storage.file), 201
<pre>fill_in_docstring() (in module</pre>
pde.tools.docstrings), 217
finalize()(DataTracker method), 255
finalize() (InteractivePlotTracker method), 247
finalize() (PlotTracker method), 258
finalize() (ProgressTracker method), 260
finalize() (StorageTracker method), 200
finalize() (TrackerBase method), 244
finalize() (TrackerCollection method), 245
<pre>find_format() (in module pde.tools.ffmpeg), 224</pre>
FinishedSimulation, 244

iss method), 215 alSymGrid class method), class method), 147 SymGridBase class method), *uboid class method*), 215 noryStorage class method), method), 104 lass method), 98 r class method), 101 on class method), 76 class method), 245 ction class method), 246 method), 104 arField class method), 80 sor2Field class method), 86 torField class method), 90 orage class method), 203 ss method), 71 class method), 81 ass method), 190 iss method), 215 ns() (FieldCollection eld class method), 91 rid class method), 158 SymGrid class method), 163 from state() (DataFieldBase class method), 61 from\_state() (FieldBase class method), 71 from\_state() (FieldCollection class method), 77 from\_state() (GridBase class method), 147 from state() (SphericalSymGridBase class method), 167 from\_state() (UnitGrid class method), 161 from\_storage() (ScalarFieldPlot class method), 265 from\_str() (BCBase class method), 105 G get\_all\_parameters() module (in pde.tools.parameters), 236 get\_axis\_index() (GridBase method), 148 module get\_boundary\_axis() (in

pde.grids.boundaries.axis), 102

get\_boundary\_conditions() (GridBase method), 148

get\_boundary\_field() (ScalarField method), 81 get\_boundary\_values() (DataFieldBase method), 61 (CylindricalSymGrid get\_cartesian\_grid() method), 163 get cartesian grid() (SphericalSymGridBase method). 167 get\_class\_by\_rank() (DataFieldBase class method), 62 get\_common\_dtype() (in module pde.tools.misc), 227 get\_common\_numba\_dtype() (in module pde.tools.numba), 231 get\_compiled() (ExpressionBase method), 219 get\_compiled\_array() (TensorExpression method), 221get\_help() (BCBase class method), 105 get\_help() (Boundaries class method), 98 get help() (BoundaryAxisBase class method), 99 get\_image\_data() (CartesianGrid method), 158 get image data() (CylindricalSymGrid method), 163 get\_image\_data() (DataFieldBase method), 62 get\_image\_data() (FieldBase method), 72 get image data() (FieldCollection method), 77 get\_image\_data() (GridBase method), 148 get\_image\_data() (SphericalSymGridBase method), 168 get\_initial\_condition() (WavePDE method), 185 get\_line\_data() (CartesianGrid method), 158 get\_line\_data() (CylindricalSymGrid method), 164 get\_line\_data() (DataFieldBase method), 62 get\_line\_data() (FieldBase method), 72 get\_line\_data() (FieldCollection method), 77 get line data() (GridBase method), 149 get\_line\_data() (SphericalSymGridBase method), 168 get\_mathematical\_representation() (BCBase method), 105 get\_mathematical\_representation() (Boundaries method), 98 get mathematical representation() (BoundaryAxisBase method), 99 get\_mathematical\_representation() (CurvatureBC method), 112 get\_mathematical\_representation() (DirichletBC method), 113 get\_mathematical\_representation() (ExpressionBC method), 115 get\_mathematical\_representation() (MixedBC method), 120 get mathematical representation() (NeumannBC method), 121 get\_mathematical\_representation()

(UserBC method), 125 get\_memory\_storage() module (in pde.storage.memory), 203 get\_named\_trackers() (in module pde.trackers.base), 247 get package versions() module (in pde.tools.config), 213 get\_parameter\_default() (Parameterized method), 235 get\_parameters () (Parameterized class method), 235 get\_plotting\_context() (in module pde.tools.plotting), 240 module get\_progress\_bar\_class() (in pde.tools.output), 234 get\_random\_point() (CartesianGrid method), 159 get\_random\_point() (CylindricalSymGrid method), 164 get random point () (GridBase method), 149 get\_random\_point() (SphericalSymGridBase method), 168 get\_sparse\_matrix\_data() (BCBase method), 105 get\_sparse\_matrix\_data() (BoundaryAxisBase method). 100 get\_sparse\_matrix\_data() (ConstBC1stOrderBase method), 108 get\_sparse\_matrix\_data() (ConstBC2ndOrderBase method), 109 get\_sparse\_matrix\_data() (ExpressionBC method), 115 get\_text\_block() (in module pde.tools.docstrings), 217 get\_vector\_data() (CartesianGrid method), 159 get\_vector\_data() (DataFieldBase method), 63 get vector data() (GridBase method), 149 get\_vector\_data() (VectorField method), 91 get\_virtual\_point() (BCBase method), 105 get\_virtual\_point() (ConstBC1stOrderBase method), 108 get\_virtual\_point() (ConstBC2ndOrderBase *method*), 110 get\_virtual\_point() (ExpressionBC method), 115 get\_virtual\_point\_data() (ConstBC1stOrderBase method), 108 (Conget\_virtual\_point\_data() stBC2ndOrderBase method), 110 get\_virtual\_point\_data() (CurvatureBC *method*), 113 get\_virtual\_point\_data() (DirichletBC method), 114 get\_virtual\_point\_data() (MixedBC method), 120 get\_virtual\_point\_data() (NeumannBC method), 121

getter() (classproperty method), 227 GhostCellSetter (class in pde.tools.typing), 243 gradient() (ScalarField method), 82 gradient\_() (VectorField method), 91 gradient\_squared() (ScalarField method), 82 grid (Boundaries attribute), 98 grid (BoundaryAxisBase property), 100 grid (FieldBase property), 72 grid (StorageBase property), 198 grid (StorageView property), 201 GridBase (class in pde.grids.base), 144

# Η

handle() (CallbackTracker method), 253 handle() (ConsistencyTracker method), 254 handle() (DataTracker method), 255 handle() (InteractivePlotTracker method), 248 handle() (MaterialConservationTracker method), 257 handle() (PlotTracker method), 258 handle() (PrintTracker method), 259 handle() (ProgressTracker method), 260 handle() (RuntimeTracker method), 260 handle() (SteadyStateTracker method), 261 handle() (StorageTracker method), 200 handle() (TrackerBase method), 245 handle() (TrackerCollection method), 246 has\_collection (StorageBase property), 198 has\_collection (Storage View attribute), 201 has\_hole (CylindricalSymGrid property), 164 has\_hole (SphericalSymGridBase property), 169 hash\_mutable() (in module pde.tools.cache), 211 hash\_readable() (in module pde.tools.cache), 211 hdf\_write\_attributes() (in module pde.tools.misc), 227 HideParameter (class in pde.tools.parameters), 235 high (BoundaryAxisBase attribute), 100 homogeneous (BCBase attribute), 105 hybridmethod (class in pde.tools.misc), 228

# I

imag (FieldBase property), 72 ImplicitSolver (class in pde.solvers), 187 ImplicitSolver (class in pde.solvers.implicit), 194 import\_class() (in module pde.tools.misc), 228 in\_ipython() (in module pde.tools.plotting), 240 in\_jupyter\_notebook() (in module pde.tools.output), 234 increment() (Counter method), 231 info (AdamsBashforthSolver attribute), 188 info (Controller attribute), 191 info (CrankNicolsonSolver attribute), 192 info (ExplicitSolver attribute), 192 info (ImplicitSolver attribute), 194 info (ScipySolver attribute), 195

initialize() (ConstantInterrupts method), 249 initialize() (FixedInterrupts method), 250 initialize() (InteractivePlotTracker method), 248 initialize() (InterruptsBase method), 251 initialize() (MaterialConservationTracker method), 257 initialize() (PlotTracker method), 259 initialize() (ProgressTracker method), 260 initialize() (RealtimeInterrupts method), 252 initialize() (RuntimeTracker method), 261 initialize() (StorageTracker method), 200 initialize() (TrackerBase method), 245 initialize() (TrackerCollection method), 246 initialized (in module pde.tools.mpi), 229 insert () (DataFieldBase method), 63 instancemethod() (hybridmethod method), 228 integral (DataFieldBase property), 63 integral (ScalarField property), 83 integral (Tensor2Field property), 87 integral (VectorField property), 92 integrals (FieldCollection property), 77 integrate() (GridBase method), 149 InteractivePlotTracker (class in pde.trackers.interactive), 247 interface\_width (AllenCahnPDE attribute), 171 interpolate() (DataFieldBase method), 63 interpolate\_to\_grid() (DataFieldBase method), 64 interpolate\_to\_grid() (FieldCollection method), 77 interpolate\_to\_grid() (ScalarField method), 83 interpolate\_to\_grid() (VectorField method), 92 InterruptsBase (class in pde.trackers.interrupts), 250 is\_available() (Movie class method), 263 is complex (FieldBase property), 72 is\_main (in module pde.tools.mpi), 230 is sde (PDEBase property), 173 is\_zero (ScalarExpression property), 220 items() (MovieStorage method), 205 items() (StorageBase method), 198 items() (StorageView method), 201 iter\_mirror\_points() (CartesianGrid method), 159 (CylindricalSymGrid iter\_mirror\_points() method), 164 iter\_mirror\_points() (GridBase method), 150

# J

# Κ

KPZInterfacePDE (class in	pde.pdes.kpz_interface),
177	
KuramotoSivashinskyPDE	(class in
pde.pdes.kuramoto_siva	shinsky), 178

# L

label (FieldBase property), 72		
labels ( <i>FieldCollection property</i> ), 78		
laplace() (ScalarField method), 83		
laplace() (VectorField method), 93		
length (CylindricalSymGrid property)	, 165	
<pre>link_value() (ConstBCBase method</pre>	<i>l</i> ), 111	
LivePlotTracker (class in pde.trac	kers.tracke	ers), 256
LogarithmicInterrupts	(class	in
pde.trackers.interrupts), 251		
low (BoundaryAxisBase attribute), 100		

# Μ

•••		
magnitude (DataFieldBase prope	erty), <mark>64</mark>	
magnitudes (FieldCollection pro	perty), 78	3
major_axis (SphericalCoordinate	tes attribu	te), 132
<pre>make_adjacent_evaluator     105</pre>	() ( <i>BCE</i>	Base method),
make_adjacent_evaluator	()	(Con-
stBC1stOrderBase method	<i>l</i> ), 108	
make_adjacent_evaluator	()	(Con-
stBC2ndOrderBase metho	od), 110	
make_adjacent_evaluator	() (	ExpressionBC
<i>method</i> ), 115		
<pre>make_array_constructor(</pre>	) (ii	n module
pde.tools.numba), 232		
<pre>make_cell_volume_compil</pre>	ed()	(GridBase
<i>method</i> ), 150		
<pre>make_colored_noise()</pre>	(in	module
pde.tools.spectral), 242		
<pre>make_derivative()</pre>	(in	module
pde.grids.operators.comm	on), 135	
<pre>make_derivative2()</pre>	(in	module
pde.grids.operators.comm	on), 135	
<pre>make_divergence()</pre>	(in	module
pde.grids.operators.cartes	<i>ian</i> ), 133	
<pre>make_divergence()</pre>	(in	module
pde.grids.operators.cylind	lrical_sym	<i>ı</i> ), 136
<pre>make_divergence()</pre>	(in	module
pde.grids.operators.polar_	_sym), 13	9
<pre>make_divergence()</pre>	(in	module
pde.grids.operators.spheri		
<pre>make_dot_operator() (Data</pre>	FieldBase	e method), 64
<pre>make_general_poisson_so</pre>	lver()	(in module
pde.grids.operators.comm	on), 135	
<pre>make_ghost_cell_sender(</pre>	) ( <i>BCB</i> )	ase method),
106		

106
<pre>make_ghost_cell_setter() (Boundaries method),</pre>
98 make wheet call catter() ( $Poundam Aris Page$
<pre>make_ghost_cell_setter() (BoundaryAxisBase method), 100</pre>
<pre>make_ghost_cell_setter() (UserBC method),</pre>
125
<pre>make_gradient() (in module</pre>
pde.grids.operators.cartesian), 133
<pre>make_gradient() (in module</pre>
pde.grids.operators.cylindrical_sym), 137
<pre>make_gradient() (in module</pre>
pde.grids.operators.polar_sym), 139
<pre>make_gradient() (in module</pre>
pde.grids.operators.spherical_sym), 142
<pre>make_gradient_squared() (in module</pre>
pde.grids.operators.cylindrical_sym), 137
<pre>make_gradient_squared() (in module</pre>
pde.grids.operators.polar_sym), 139
<pre>make_gradient_squared() (in module</pre>
pde.grids.operators.spherical_sym), 142
<pre>make_inserter_compiled() (GridBase method),</pre>
<pre>make_integrator() (GridBase method), 150</pre>
<pre>make_interpolator() (DataFieldBase method), 65</pre>
<pre>make_laplace() (in module</pre>
pde.grids.operators.cartesian), 134
<pre>make_laplace() (in module</pre>
pde.grids.operators.cylindrical_sym), 137
<pre>make_laplace() (in module</pre>
<pre>make_laplace() (in module pde.grids.operators.polar_sym), 140</pre>
pde.grids.operators.polar_sym), 140 make_laplace() (in module
pde.grids.operators.polar_sym), 140make_laplace()(inmodulepde.grids.operators.spherical_sym), 142
<pre>pde.grids.operators.polar_sym), 140 make_laplace() (in module     pde.grids.operators.spherical_sym), 142 make_laplace_from_matrix() (in module</pre>
<pre>pde.grids.operators.polar_sym), 140 make_laplace() (in module     pde.grids.operators.spherical_sym), 142 make_laplace_from_matrix() (in module     pde.grids.operators.common), 136</pre>
<pre>pde.grids.operators.polar_sym), 140 make_laplace() (in module     pde.grids.operators.spherical_sym), 142 make_laplace_from_matrix() (in module</pre>
<pre>pde.grids.operators.polar_sym), 140 make_laplace() (in module     pde.grids.operators.spherical_sym), 142 make_laplace_from_matrix() (in module     pde.grids.operators.common), 136 make_modify_after_step() (PDEBase method),     173 make_movie() (ScalarFieldPlot method), 265</pre>
<pre>pde.grids.operators.polar_sym), 140 make_laplace() (in module     pde.grids.operators.spherical_sym), 142 make_laplace_from_matrix() (in module     pde.grids.operators.common), 136 make_modify_after_step() (PDEBase method),     173 make_movie() (ScalarFieldPlot method), 265 make_normalize_point_compiled() (GridBase</pre>
<pre>pde.grids.operators.polar_sym), 140 make_laplace() (in module     pde.grids.operators.spherical_sym), 142 make_laplace_from_matrix() (in module     pde.grids.operators.common), 136 make_modify_after_step() (PDEBase method),     173 make_movie() (ScalarFieldPlot method), 265 make_normalize_point_compiled() (GridBase     method), 151</pre>
<pre>pde.grids.operators.polar_sym), 140 make_laplace() (in module     pde.grids.operators.spherical_sym), 142 make_laplace_from_matrix() (in module     pde.grids.operators.common), 136 make_modify_after_step() (PDEBase method),     173 make_movie() (ScalarFieldPlot method), 265 make_normalize_point_compiled() (GridBase     method), 151 make_operator() (GridBase method), 151</pre>
<pre>pde.grids.operators.polar_sym), 140 make_laplace() (in module     pde.grids.operators.spherical_sym), 142 make_laplace_from_matrix() (in module     pde.grids.operators.common), 136 make_modify_after_step() (PDEBase method),     173 make_movie() (ScalarFieldPlot method), 265 make_normalize_point_compiled() (GridBase     method), 151 make_operator() (GridBase method), 151 make_operator_no_bc() (GridBase method), 152</pre>
<pre>pde.grids.operators.polar_sym), 140 make_laplace() (in module     pde.grids.operators.spherical_sym), 142 make_laplace_from_matrix() (in module     pde.grids.operators.common), 136 make_modify_after_step() (PDEBase method),     173 make_movie() (ScalarFieldPlot method), 265 make_normalize_point_compiled() (GridBase     method), 151 make_operator() (GridBase method), 151</pre>
<pre>pde.grids.operators.polar_sym), 140 make_laplace() (in module     pde.grids.operators.spherical_sym), 142 make_laplace_from_matrix() (in module     pde.grids.operators.common), 136 make_modify_after_step() (PDEBase method),     173 make_movie() (ScalarFieldPlot method), 265 make_normalize_point_compiled() (GridBase     method), 151 make_operator() (GridBase method), 152 make_outer_prod_operator() (VectorField</pre>
<pre>pde.grids.operators.polar_sym), 140 make_laplace() (in module     pde.grids.operators.spherical_sym), 142 make_laplace_from_matrix() (in module     pde.grids.operators.common), 136 make_modify_after_step() (PDEBase method),     173 make_movie() (ScalarFieldPlot method), 265 make_normalize_point_compiled() (GridBase     method), 151 make_operator() (GridBase method), 152 make_outer_prod_operator() (VectorField     method), 93</pre>
<pre>pde.grids.operators.polar_sym), 140 make_laplace() (in module     pde.grids.operators.spherical_sym), 142 make_laplace_from_matrix() (in module     pde.grids.operators.common), 136 make_modify_after_step() (PDEBase method),     173 make_movie() (ScalarFieldPlot method), 265 make_normalize_point_compiled() (GridBase     method), 151 make_operator() (GridBase method), 152 make_outer_prod_operator() (VectorField     method), 93 make_pde_rhs() (PDEBase method), 173</pre>
<pre>pde.grids.operators.polar_sym), 140 make_laplace() (in module     pde.grids.operators.spherical_sym), 142 make_laplace_from_matrix() (in module     pde.grids.operators.common), 136 make_modify_after_step() (PDEBase method),     173 make_movie() (ScalarFieldPlot method), 265 make_normalize_point_compiled() (GridBase     method), 151 make_operator() (GridBase method), 151 make_operator_no_bc() (GridBase method), 152 make_outer_prod_operator() (VectorField     method), 93 make_poisson_solver() (in module </pre>
<pre>pde.grids.operators.polar_sym), 140 make_laplace() (in module     pde.grids.operators.spherical_sym), 142 make_laplace_from_matrix() (in module     pde.grids.operators.common), 136 make_modify_after_step() (PDEBase method),     173 make_movie() (ScalarFieldPlot method), 265 make_normalize_point_compiled() (GridBase     method), 151 make_operator() (GridBase method), 151 make_operator_no_bc() (GridBase method), 152 make_outer_prod_operator() (VectorField     method), 93 make_pde_rhs() (PDEBase method), 173 make_operator_scartesian), 134</pre>
<pre>de.grids.operators.polar_sym), 140 make_laplace() (in module     pde.grids.operators.spherical_sym), 142 make_laplace_from_matrix() (in module     pde.grids.operators.common), 136 make_movie() (ScalarFieldPlot method), 265 make_normalize_point_compiled() (GridBase     method), 151 make_operator() (GridBase method), 152 make_operator_no_bc() (GridBase method), 152 make_outer_prod_operator() (VectorField     method), 93 make_pde_rhs() (PDEBase method), 134 make_poisson_solver() (in module     pde.grids.operators.cartesian), 134 make_poisson_solver() (in module     pde.grids.operators.cylindrical_sym), 138 make_poisson_solver() (in module</pre>
<pre>pde.grids.operators.polar_sym), 140 make_laplace() (in module     pde.grids.operators.spherical_sym), 142 make_laplace_from_matrix() (in module     pde.grids.operators.common), 136 make_modify_after_step() (PDEBase method),     173 make_movie() (ScalarFieldPlot method), 265 make_normalize_point_compiled() (GridBase     method), 151 make_operator() (GridBase method), 151 make_operator_no_bc() (GridBase method), 152 make_outer_prod_operator() (VectorField     method), 93 make_poisson_solver() (in module     pde.grids.operators.cylindrical_sym), 138 make_poisson_solver() (in module     pde.grids.operators.polar_sym), 140</pre>
<pre>de.grids.operators.polar_sym), 140 make_laplace() (in module     pde.grids.operators.spherical_sym), 142 make_laplace_from_matrix() (in module     pde.grids.operators.common), 136 make_movie() (ScalarFieldPlot method), 265 make_normalize_point_compiled() (GridBase     method), 151 make_operator() (GridBase method), 152 make_operator_no_bc() (GridBase method), 152 make_outer_prod_operator() (VectorField     method), 93 make_pde_rhs() (PDEBase method), 134 make_poisson_solver() (in module     pde.grids.operators.cartesian), 134 make_poisson_solver() (in module     pde.grids.operators.cylindrical_sym), 138 make_poisson_solver() (in module</pre>

<pre>make_serializer() (in module pde.tools.cache), 212</pre>
<pre>make_stepper() (AdaptiveSolverBase method), 189</pre>
<pre>make_stepper() (ExplicitMPISolver method), 194</pre>
<pre>make_stepper() (ScipySolver method), 187, 195</pre>
<pre>make_stepper() (SolverBase method), 190</pre>
<pre>make_tensor_divergence() (in module</pre>
pde.grids.operators.cartesian), 134
<pre>make_tensor_divergence() (in module</pre>
pde.grids.operators.cylindrical_sym), 138
<pre>make_tensor_divergence() (in module</pre>
pde.grids.operators.polar_sym), 140
make_tensor_divergence() (in module
pde.grids.operators.spherical_sym), 143
make_tensor_double_divergence() (in module
<pre>pde.grids.operators.spherical_sym), 143 make_unserializer() (in module pde.tools.cache),</pre>
212
make_vector_gradient() (in module
pde.grids.operators.cartesian), 134
make_vector_gradient() (in module
pde.grids.operators.cylindrical_sym), 138
make_vector_gradient() (in module
pde.grids.operators.polar_sym), 141
make_vector_gradient() (in module
pde.grids.operators.spherical_sym), 144
make_vector_laplace() (in module
pde.grids.operators.cartesian), 135
<pre>make_vector_laplace() (in module</pre>
pde.grids.operators.cylindrical_sym), 138 make_virtual_point_evaluator() (BCBase
pde.grids.operators.cylindrical_sym), 138
<pre>pde.grids.operators.cylindrical_sym), 138 make_virtual_point_evaluator() (BCBase</pre>
<pre>pde.grids.operators.cylindrical_sym), 138 make_virtual_point_evaluator() (BCBase method), 106</pre>
<pre>pde.grids.operators.cylindrical_sym), 138 make_virtual_point_evaluator() (BCBase     method), 106 make_virtual_point_evaluator() (Con-     stBC1stOrderBase method), 109 make_virtual_point_evaluator() (Con-</pre>
<pre>pde.grids.operators.cylindrical_sym), 138 make_virtual_point_evaluator() (BCBase method), 106 make_virtual_point_evaluator() (Con- stBC1stOrderBase method), 109 make_virtual_point_evaluator() (Con- stBC2ndOrderBase method), 110</pre>
<pre>pde.grids.operators.cylindrical_sym), 138 make_virtual_point_evaluator() (BCBase     method), 106 make_virtual_point_evaluator() (Con-     stBC1stOrderBase method), 109 make_virtual_point_evaluator() (Con-</pre>
<pre>pde.grids.operators.cylindrical_sym), 138 make_virtual_point_evaluator() (BCBase     method), 106 make_virtual_point_evaluator() (Con-     stBC1stOrderBase method), 109 make_virtual_point_evaluator() (Con-     stBC2ndOrderBase method), 110 make_virtual_point_evaluator() (Expres- </pre>
<pre>pde.grids.operators.cylindrical_sym), 138 make_virtual_point_evaluator() (BCBase</pre>
<pre>pde.grids.operators.cylindrical_sym), 138 make_virtual_point_evaluator() (BCBase</pre>
<pre>pde.grids.operators.cylindrical_sym), 138 make_virtual_point_evaluator() (BCBase     method), 106 make_virtual_point_evaluator() (Con-     stBC1stOrderBase method), 109 make_virtual_point_evaluator() (Con-     stBC2ndOrderBase method), 110 make_virtual_point_evaluator() (ExpressionBC method), 115 make_virtual_point_evaluator() (UserBC     method), 126 mapping_jacobian() (CoordinatesBase method),</pre>
<pre>pde.grids.operators.cylindrical_sym), 138 make_virtual_point_evaluator() (BCBase</pre>
<pre>pde.grids.operators.cylindrical_sym), 138 make_virtual_point_evaluator() (BCBase</pre>
<pre>pde.grids.operators.cylindrical_sym), 138 make_virtual_point_evaluator() (BCBase</pre>
<pre>pde.grids.operators.cylindrical_sym), 138 make_virtual_point_evaluator() (BCBase     method), 106 make_virtual_point_evaluator() (Con-     stBC1stOrderBase method), 109 make_virtual_point_evaluator() (Con-     stBC2ndOrderBase method), 110 make_virtual_point_evaluator() (ExpressionBC method), 115 make_virtual_point_evaluator() (UserBC     method), 126 mapping_jacobian() (CoordinatesBase method),     128 MaterialConservationTracker (class in     pde.trackers.trackers), 256 max_value (FFmpegFormat property), 224</pre>
<pre>pde.grids.operators.cylindrical_sym), 138 make_virtual_point_evaluator() (BCBase     method), 106 make_virtual_point_evaluator() (Con-     stBC1stOrderBase method), 109 make_virtual_point_evaluator() (Con-     stBC2ndOrderBase method), 110 make_virtual_point_evaluator() (ExpressionBC method), 115 make_virtual_point_evaluator() (UserBC     method), 126 mapping_jacobian() (CoordinatesBase method),     128 MaterialConservationTracker (class in     pde.trackers.trackers), 256 max_value (FFmpegFormat property), 224 mean (OnlineStatistics attribute), 225</pre>
<pre>pde.grids.operators.cylindrical_sym), 138 make_virtual_point_evaluator() (BCBase     method), 106 make_virtual_point_evaluator() (Con-     stBC1stOrderBase method), 109 make_virtual_point_evaluator() (Con-     stBC2ndOrderBase method), 110 make_virtual_point_evaluator() (ExpressionBC method), 115 make_virtual_point_evaluator() (UserBC     method), 126 mapping_jacobian() (CoordinatesBase method),     128 MaterialConservationTracker (class in     pde.trackers.trackers), 256 max_value (FFmpegFormat property), 224 mean (OnlineStatistics attribute), 225 MemoryStorage (class in pde.tracge.memory), 202</pre>
<pre>pde.grids.operators.cylindrical_sym), 138 make_virtual_point_evaluator() (BCBase</pre>
<pre>pde.grids.operators.cylindrical_sym), 138 make_virtual_point_evaluator() (BCBase</pre>
<pre>pde.grids.operators.cylindrical_sym), 138 make_virtual_point_evaluator() (BCBase     method), 106 make_virtual_point_evaluator() (Con-     stBC1stOrderBase method), 109 make_virtual_point_evaluator() (Con-     stBC2ndOrderBase method), 110 make_virtual_point_evaluator() (ExpressionBC method), 115 make_virtual_point_evaluator() (UserBC     method), 126 mapping_jacobian() (CoordinatesBase method),     128 MaterialConservationTracker (class in     pde.trackers.trackers), 256 max_value (FFmpegFormat property), 224 mean (OnlineStatistics attribute), 225 MemoryStorage (class in pde.storage.memory), 202 metric() (CoordinatesBase method), 128 MixedBC (class in pde.grids.boundaries.local), 119 module     pde, 59     pde.fields, 59</pre>
<pre>pde.grids.operators.cylindrical_sym), 138 make_virtual_point_evaluator() (BCBase</pre>

pde.fields.scalar,80 pde.fields.tensorial,85 pde.fields.vectorial,89 pde.grids,94 pde.grids.base,144 pde.grids.boundaries,95 pde.grids.boundaries.axes,97 pde.grids.boundaries.axis,99 pde.grids.boundaries.local, 102 pde.grids.cartesian, 157 pde.grids.coordinates, 127 pde.grids.coordinates.base, 127 pde.grids.coordinates.bipolar, 130 pde.grids.coordinates.bispherical, 130 pde.grids.coordinates.cartesian, 131 pde.grids.coordinates.cylindrical, 131 pde.grids.coordinates.polar, 132 pde.grids.coordinates.spherical, 132 pde.grids.cylindrical, 161 pde.grids.operators, 133 pde.grids.operators.cartesian, 133 pde.grids.operators.common, 135 pde.grids.operators.cylindrical\_sym, 136 pde.grids.operators.polar\_sym, 139 pde.grids.operators.spherical\_sym, 141 pde.grids.spherical, 165 pde.pdes, 170 pde.pdes.allen\_cahn, 170 pde.pdes.base, 171 pde.pdes.cahn\_hilliard, 175 pde.pdes.diffusion, 176 pde.pdes.kpz\_interface, 177 pde.pdes.kuramoto sivashinsky, 178 pde.pdes.laplace, 180 pde.pdes.pde, 181 pde.pdes.swift\_hohenberg, 183 pde.pdes.wave, 184 pde.solvers, 185 pde.solvers.adams\_bashforth, 188 pde.solvers.base, 188 pde.solvers.controller, 190 pde.solvers.crank\_nicolson, 192 pde.solvers.explicit, 192 pde.solvers.explicit\_mpi, 193 pde.solvers.implicit, 194 pde.solvers.scipy, 195 pde.storage, 195 pde.storage.base, 196 pde.storage.file, 201 pde.storage.memory, 202

```
pde.storage.movie, 204
    pde.tools, 207
    pde.tools.cache,207
    pde.tools.config,213
    pde.tools.cuboid, 214
    pde.tools.docstrings, 217
    pde.tools.expressions, 217
    pde.tools.ffmpeq, 223
    pde.tools.math, 225
    pde.tools.misc, 226
   pde.tools.mpi, 229
    pde.tools.numba, 231
    pde.tools.output, 233
    pde.tools.parameters, 234
    pde.tools.parse_duration,237
    pde.tools.plotting,237
    pde.tools.spectral, 242
    pde.tools.typing,243
   pde.trackers, 243
    pde.trackers.base, 244
    pde.trackers.interactive, 247
    pde.trackers.interrupts, 249
    pde.trackers.trackers, 253
    pde.visualization, 262
    pde.visualization.movies, 262
    pde.visualization.plotting, 264
module_available() (in module pde.tools.misc), 228
Movie (class in pde.visualization.movies), 262
movie() (in module pde.visualization.movies), 263
movie_multiple()
                            (in
                                        module
       pde.visualization.movies), 263
movie_scalar() (in module pde.visualization.movies),
        264
MovieStorage (class in pde.storage.movie), 204
mpi allreduce() (in module pde.tools.mpi), 230
mpi_recv() (in module pde.tools.mpi), 230
mpi send() (in module pde.tools.mpi), 230
mutable (Cuboid property), 216
```

# Ν

name (AdamsBashforthSolver attribute), 185, 188 name (ConsistencyTracker attribute), 254 name (CrankNicolsonSolver attribute), 187, 192 name (ExplicitMPISolver attribute), 194 name (ExplicitSolver attribute), 187, 192 name (ImplicitSolver attribute), 187, 194 name (InteractivePlotTracker attribute), 248 name (LivePlotTracker attribute), 256 name (MaterialConservationTracker attribute), 257 name (OperatorInfo attribute), 156 name (PrintTracker attribute), 259 name (ProgressTracker attribute), 260 name (ScipySolver attribute), 188, 195 name (SteadyStateTracker attribute), 262 names (BCBase attribute), 106 names (CurvatureBC attribute), 113 names (DirichletBC attribute), 114 names (ExpressionBC attribute), 116 names (ExpressionDerivativeBC attribute), 117 names (ExpressionMixedBC attribute), 118 names (ExpressionValueBC attribute), 119 names (MixedBC attribute), 120 names (NeumannBC attribute), 121 names (NormalCurvatureBC attribute), 122 names (NormalDirichletBC attribute), 123 names (NormalMixedBC attribute), 124 names (NormalNeumannBC attribute), 124 names (UserBC attribute), 126 napari\_add\_layers() (in module pde.tools.plotting), 240 module napari\_process() (in pde.trackers.interactive), 248 napari\_viewer() (in module pde.tools.plotting), 240 NapariViewer (class in pde.trackers.interactive), 248 nested\_plotting\_check (class in pde.tools.plotting), 240 NeumannBC (class in pde.grids.boundaries.local), 121 next () (ConstantInterrupts method), 249 next () (FixedInterrupts method), 250 next () (InterruptsBase method), 251 next () (LogarithmicInterrupts method), 251 next () (RealtimeInterrupts method), 252 noise\_realization() (PDEBase method), 174 normal (BCBase attribute), 106 normal (NormalCurvatureBC attribute), 122 normal (NormalDirichletBC attribute), 123 normal (NormalMixedBC attribute), 124 normal (NormalNeumannBC attribute), 125 NormalCurvatureBC (class in pde.grids.boundaries.local), 121 NormalDirichletBC (class in pde.grids.boundaries.local), 122 normalize\_point() (GridBase method), 152 NormalMixedBC (class in pde.grids.boundaries.local), 123 NormalNeumannBC (class in pde.grids.boundaries.local), 124 num\_axes (GridBase attribute), 152 num\_cells (GridBase property), 152 numba\_dict() (in module pde.tools.numba), 232 numba\_environment() (in module pde.tools.numba), 232 numba\_type (GridBase property), 152 number() (in module pde.tools.misc), 228 number\_array() (in module pde.tools.misc), 228

# 0

objects\_equal() (in module pde.tools.cache), 212

ol\_flat\_idx() (in module pde.tools.numba), 232 ol\_mpi\_allreduce() (in module pde.tools.mpi), 230 ol\_mpi\_recv() (in module pde.tools.mpi), 230 ol\_mpi\_send() (in module pde.tools.mpi), 231 OnlineStatistics (class in pde.tools.math), 225 OperatorFactory (class in pde.tools.typing), 243 OperatorInfo (class in pde.grids.base), 155 operatorS (GridBase attribute), 153 OperatorType (class in pde.tools.typing), 243 outer\_product() (VectorField method), 93 OutputBase (class in pde.tools.output), 233

# Ρ

packages\_from\_requirements() (in module pde.tools.config), 214 parallel\_run (in module pde.tools.mpi), 231 Parameter (class in pde.tools.parameters), 235 Parameterized (class in pde.tools.parameters), 235 parameters (PlotReference attribute), 238 parameters\_default (Parameterized attribute), 236 parse\_duration() (in module pde.tools.parse\_duration), 237 parse\_interrupt() (in module pde.trackers.interrupts), 252 parse\_number() (in module pde.tools.expressions), 2.2.2 parse\_version\_str() (in module pde.tools.config), 214pde module, 59 PDE (class in pde.pdes.pde), 181 pde.fields module, 59 pde.fields.base module, 60 pde.fields.collection module,74 pde.fields.scalar module, 80 pde.fields.tensorial module,85 pde.fields.vectorial module,89 pde.grids module,94 pde.grids.base module, 144 pde.grids.boundaries module, 95 pde.grids.boundaries.axes module, 97 pde.grids.boundaries.axis module, 99 pde.grids.boundaries.local

module, 102 pde.grids.cartesian module, 157 pde.grids.coordinates module, 127 pde.grids.coordinates.base module, 127 pde.grids.coordinates.bipolar module, 130 pde.grids.coordinates.bispherical module, 130 pde.grids.coordinates.cartesian module, 131 pde.grids.coordinates.cylindrical module, 131 pde.grids.coordinates.polar module, 132 pde.grids.coordinates.spherical module, 132 pde.grids.cylindrical module, 161 pde.grids.operators module, 133 pde.grids.operators.cartesian module, 133 pde.grids.operators.common module, 135 pde.grids.operators.cylindrical\_sym module, 136 pde.grids.operators.polar\_sym module, 139 pde.grids.operators.spherical\_sym module, 141 pde.grids.spherical module, 165 pde.pdes module, 170 pde.pdes.allen\_cahn module, 170 pde.pdes.base module, 171 pde.pdes.cahn\_hilliard module, 175 pde.pdes.diffusion module, 176 pde.pdes.kpz\_interface module, 177 pde.pdes.kuramoto\_sivashinsky module, 178 pde.pdes.laplace module, 180 pde.pdes.pde module, 181

pde.pdes.swift\_hohenberg

module, 183 pde.pdes.wave module, 184 pde.solvers module, 185 pde.solvers.adams bashforth module.188 pde.solvers.base module.188 pde.solvers.controller module, 190 pde.solvers.crank\_nicolson module, 192 pde.solvers.explicit module, 192 pde.solvers.explicit\_mpi module, 193 pde.solvers.implicit module, 194 pde.solvers.scipy module, 195 pde.storage module, 195 pde.storage.base module, 196 pde.storage.file module, 201 pde.storage.memory module, 202 pde.storage.movie module, 204 pde.tools module, 207 pde.tools.cache module, 207 pde.tools.config module, 213 pde.tools.cuboid module, 214 pde.tools.docstrings module, 217 pde.tools.expressions module, 217 pde.tools.ffmpeg module, 223 pde.tools.math module, 225 pde.tools.misc module, 226 pde.tools.mpi module, 229 pde.tools.numba module, 231 pde.tools.output

module, 233 pde.tools.parameters module, 234 pde.tools.parse\_duration module, 237 pde.tools.plotting module.237 pde.tools.spectral module, 242 pde.tools.typing module, 243 pde.trackers module, 243 pde.trackers.base module, 244 pde.trackers.interactive module, 247 pde.trackers.interrupts module, 249 pde.trackers.trackers module, 253 pde.visualization module, 262 pde.visualization.movies module, 262 pde.visualization.plotting module, 264 PDEBase (class in pde.pdes.base), 171 periodic (BCBase property), 106 periodic (Boundaries property), 99 periodic (BoundaryAxisBase property), 100 periodic (GridBase property), 153 PeriodicityError, 156 pix\_fmt\_data (FFmpegFormat attribute), 224 pix fmt file (FFmpegFormat attribute), 224 plot() (CartesianGrid method), 159 plot () (DataFieldBase method), 65 plot() (FieldBase method), 73 plot () (FieldCollection method), 78 plot () (GridBase method), 153 plot () (SphericalSymGridBase method), 169 plot\_components() (Tensor2Field method), 87 plot\_interactive() (FieldBase method), 73 module plot\_interactive() (in pde.visualization.plotting), 266 module plot\_kymograph() (in pde.visualization.plotting), 267 plot\_kymographs() (in module pde.visualization.plotting), 268 plot\_magnitudes() module (in pde.visualization.plotting), 268 plot\_on\_axes() (in module pde.tools.plotting), 241 plot\_on\_figure() (in module pde.tools.plotting), 241 PlotReference (class in pde.tools.plotting), 238

PlottingContextBase (class in pde.tools.plotting), 239 PlotTracker (class in pde.trackers.trackers), 257 point\_from\_cartesian() (GridBase method), 153 point to cartesian() (GridBase method), 153 PolarCoordinates (class in pde.grids.coordinates.polar), 132 PolarSymGrid (class in pde.grids.spherical), 165 pos\_diff() (CoordinatesBase method), 128 pos\_from\_cart() (CoordinatesBase method), 129 pos\_to\_cart() (CoordinatesBase method), 129 preserve\_scalars() (in module pde.tools.misc), 229 PrintTracker (class in pde.trackers.trackers), 259 progress\_bar\_format (SteadyStateTracker attribute), 262 ProgressTracker (class in pde.trackers.trackers), 259 project () (ScalarField method), 84 PYTHONPATH, 5

# R

radius (CylindricalSymGrid property), 165 radius (SphericalSymGridBase property), 169 random colored() (DataFieldBase class method), 66 random\_harmonic() (DataFieldBase class method), 67 random\_normal() (DataFieldBase class method), 67 random seed() (in module pde.tools.numba), 232 random uniform() (DataFieldBase class method), 68 rank (BoundaryAxisBase property), 100 rank (DataFieldBase attribute), 68 rank (ExpressionBase property), 219 rank (in module pde.tools.mpi), 231 rank (ScalarField attribute), 84 rank (Tensor2Field attribute), 88 rank (TensorExpression property), 221 rank (VectorField attribute), 94 rank\_in (OperatorInfo attribute), 156 rank\_out (OperatorInfo attribute), 156 RankError, 74 real (FieldBase property), 73 RealtimeInterrupts (class in pde.trackers.interrupts), 251 register\_operator() (GridBase class method), 153 registered\_boundary\_condition\_classes() (in module pde.grids.boundaries.local), 126 registered\_boundary\_condition\_names() (in module pde.grids.boundaries.local), 126 registered\_operators() (in module pde.grids.base), 156 registered\_solvers (SolverBase attribute), 190 registered\_solvers() (in module pde.solvers), 188 replace\_in\_docstring() (in module pde.tools.docstrings), 217 run () (Controller method), 186, 191

RuntimeTracker (class in pde.trackers.trackers), 260

## S

- save() (Movie method), 263
- savefig() (ScalarFieldPlot method), 266
- ScalarExpression (class in pde.tools.expressions), 219
- ScalarField (class in pde.fields.scalar), 80
- ScalarFieldPlot (*class in pde.visualization.plotting*), 264

```
scale_factors() (CoordinatesBase method), 129
```

- ScipySolver (class in pde.solvers), 187
- ScipySolver (class in pde.solvers.scipy), 195
- SerializedDict (class in pde.tools.cache), 208
- set\_ghost\_cells() (BCBase method), 106
- set\_ghost\_cells() (Boundaries method), 99
- set\_ghost\_cells() (BoundaryAxisBase method),
  100
- set\_ghost\_cells() (ConstBC1stOrderBase method),
  109
- set\_ghost\_cells() (ConstBC2ndOrderBase method), 110

```
set_ghost_cells() (DataFieldBase method), 68
```

```
set_ghost_cells() (ExpressionBC method), 116
```

- set\_ghost\_cells() (UserBC method), 126
- setter() (classproperty method), 227
- shape (ExpressionBase property), 219
- shape (GridBase property), 154
- shape (ScalarExpression attribute), 220
- shape (StorageBase property), 198
- shape (TensorExpression property), 221
- show() (*BasicOutput method*), 233 show() (*JupyterOutput method*), 233
- show () (*OutputBase method*), 233
- show\_parameters() (*Parameterized method*), 236
- sigma\_auto\_scale (SmoothData1D attribute), 226
- size (Cuboid property), 216
- size (in module pde.tools.mpi), 231
- skipUnlessModule() (in module pde.tools.misc), 229
- slice() (CartesianGrid method), 160
- slice() (CylindricalSymGrid method), 165
- slice() (GridBase method), 154
- slice() (ScalarField method), 84
- slice() (UnitGrid method), 161
- smooth() (DataFieldBase method), 69
- smooth() (FieldCollection method), 79
- SmoothData1D (class in pde.tools.math), 225
- solve() (PDEBase method), 174

SolverBase (class in pde.solvers.base), 189 SphericalCoordinates (class pde.grids.coordinates.spherical), 132 SphericalSymGrid (class in pde.grids.spherical), 166 SphericalSymGridBase (class in pde.grids.spherical), 166 sphinx\_display\_parameters() (in module pde.tools.parameters), 236 split\_mpi() (FieldBase method), 73 start\_writing() (FileStorage method), 202 start\_writing() (MemoryStorage method), 203 start\_writing() (MovieStorage method), 205 start\_writing() (StorageBase method), 198 state (*CartesianGrid property*), 160 state (CylindricalSymGrid property), 165 state (GridBase property), 154 state (SphericalSymGridBase property), 169 state (UnitGrid property), 161 state\_serialized (GridBase property), 154 SteadyStateTracker (class in pde.trackers.trackers), 261 storage (Storage Tracker attribute), 199 StorageBase (class in pde.storage.base), 196 StorageTracker (class in pde.storage.base), 199 StorageView (class in pde.storage.base), 200 supports\_update (JupyterPlottingContext attribute), 238 supports\_update (PlottingContextBase attribute), 239 surface\_area (Cuboid property), 216 SwiftHohenbergPDE (class in pde.pdes.swift\_hohenberg), 183 symmetrize() (Tensor2Field method), 88

# Т

t\_range (Controller property), 186, 191 Tensor2Field (class in pde.fields.tensorial), 85 TensorExpression (class in pde.tools.expressions), 220 time next action (TrackerCollection attribute), 246 times (DataTracker attribute), 254 times (*FileStorage property*), 202 times (MovieStorage attribute), 206 times (StorageBase attribute), 198 times (Storage View property), 201 to\_cartesian() (UnitGrid method), 161 to\_dict() (Config method), 213 to\_file() (DataTracker method), 255 to\_file() (FieldBase method), 73 to\_scalar() (DataFieldBase method), 69 to\_scalar() (ScalarField method), 85 to\_scalar() (Tensor2Field method), 88 to\_scalar() (VectorField method), 94 to\_subgrid() (BCBase method), 107 to\_subgrid() (ConstBCBase method), 112

to subgrid() (ExpressionBC method), 116 in to\_subgrid() (MixedBC method), 120 to subgrid() (UserBC method), 126 trace() (Tensor2Field method), 88 tracker() (MovieStorage method), 206 tracker() (StorageBase method), 198 tracker action times (TrackerCollection attribute), 246 TrackerBase (class in pde.trackers.base), 244 TrackerCollection (class in pde.trackers.base), 245 trackers (TrackerCollection attribute), 245 transform() (GridBase method), 154 transpose() (Tensor2Field method), 89 typical\_discretization (GridBase property), 155

# U

uniform cell volumes (GridBase attribute), 155 uniform\_discretization() (in module pde.grids.operators.common), 136 UnitGrid (class in pde.grids.cartesian), 160 unserialize\_attributes() (DataFieldBase class method), 69 (FieldBase unserialize attributes() class method), 74 unserialize\_attributes() (FieldCollection class method), 80 update() (DictFiniteCapacity method), 207 update() (NapariViewer method), 248 update() (ScalarFieldPlot method), 266 UserBC (class in pde.grids.boundaries.local), 125

# V

```
value (ConstBCBase property), 112
value (ScalarExpression property), 220
value (TensorExpression property), 221
value_is_linked (ConstBCBase attribute), 112
variables (PDE attribute), 181
vec to cart() (CoordinatesBase method), 129
VectorField (class in pde. fields. vectorial), 89
vertices (Cuboid property), 216
view_field() (StorageBase method), 199
VirtualPointEvaluator (class in pde.tools.typing),
         243
volume (CartesianGrid property), 160
volume (Cuboid property), 216
volume (CylindricalSymGrid property), 165
volume (GridBase property), 155
volume (SphericalSymGridBase property), 169
volume_factor() (CoordinatesBase method), 129
volume from radius()
                                             module
                                   (in
         pde.grids.spherical), 169
```

# W

WavePDE (class in pde.pdes.wave), 184

write\_mode (StorageBase attribute), 199
writeable (FieldBase property), 74