WaveBlocksND Documentation

Release devel

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Reusable building blocks for simulations with semiclassical wavepackets for solving the time-dependent Schrödinger equation.

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1.1.1 Basic numerics

ComplexMath

About the ComplexMath class

The WaveBlocks Project

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Class documentation

The WaveBlocks Project

Some selected functions for complex math.

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ComplexMath.continuate(data, jump=6.283185307179586, reference=0.0)

Make the given data continuous by removing all jumps of size $k \cdot \text{jump}$ but not touching jumps of any other size. There may be arise problems with jumps that are of size nearly $k \cdot \text{jump}$.

Parameters

- **data** – An array with the input data.
- **jump** – The basic size of jumps which will be removed. Default is $2\pi$.
- **reference** (A single float number.) – This value allows the specify the starting point for continuation explicitly. It can be used together with data.

ComplexMath.cont_angle(data, reference=None)

Compute the angle of a complex number not constrained to the principal value and avoiding discontinuities at the branch cut. This function just applies `continuate(.)` to the complex phase.

Parameters

- **data** – An array with the input data.
• **reference** *(A single float number)* – This value allows the specify the starting point for continuation explicitly. It can be used together with `data`.

**ComplexMath**. `cont_sqrt(data, reference=None)`  
Compute the complex square root (following the Riemann surface) yields a result *not* constrained to the principal value and avoiding discontinuities at the branch cut. This function applies ‘continue(.’ to the complex phase and computes the complex square root according to the formula $\sqrt{z} = \sqrt{r} \exp\left(i \cdot \frac{\phi}{2}\right)$.

**Parameters**

- **data** – An array with the input data.
- **reference** *(A single float number)* – This value allows the specify the starting point for continuation explicitly. It can be used together with `data`.

### Grid

**About the Grid class**

The WaveBlocks Project

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### Inheritance diagram

```
Grid.Grid
```

**Class documentation**

```python
class WaveBlocksND.Grid
    This class is an abstract interface to grids in general.
    get_dimension()
        Return the dimension $D$ of the grid.
```

### DenseGrid

**About the DenseGrid class**

The WaveBlocks Project

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Inheritance diagram

```
Grid.Grid → DenseGrid.DenseGrid
```

Class documentation

class WaveBlocksND.DenseGrid
   This class is an abstract interface to dense grids in general.
   
   **get_dimension()**
   Return the dimension $D$ of the grid.

   **is_regular()**
   Answers the question if the grid spacing is regular. (It can still be different along each axis!)

TensorProductGrid

**About the TensorProductGrid class**

The WaveBlocks Project

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Inheritance diagram

```
Grid.Grid → DenseGrid.DenseGrid → TensorProductGrid.TensorProductGrid
```

Class documentation

class WaveBlocksND.TensorProductGrid(limits, number_nodes)
   This class represents a dense tensor product grid. It can have an arbitrary dimension $D$. The grid nodes are enclosed in a hypercubic bounding box. This box can have different limits $min_i$, $max_i$ along each axis $x_i$. In each of these intervals we place $N_i$ grid nodes. Note that the point $max_i$ is not part of the grid. The grid interior is built as the tensor product of all the grid nodes along all the axes.

   **get_axes(axes=None)**
   Returns the one-dimensional grids along the axes.
Parameters **axes** (A single integer or a list of integers. If set to `None` (default) we return the data for all axes.) – The axes for which we want to get the grid.

Returns A list of ndarrays, each having a shape of \((1, ..., N_i, ..., 1)\). We return a list even if it contains just a single element.

**get_dimension()**
Return the dimension \(D\) of the grid.

**get_extensions**(axes=`None`)
Returns the extensions (length of the edges) of the bounding box.

Parameters **axes** (A single integer or a list of integers. If set to `None` (default) we return the extensions for all axes.) – The axes for which we want to get the extensions.

Returns A list of \(|\text{max}_i - \text{min}_i|\) values.

**get_limits**(axes=`None`)
Returns the limits of the bounding box.

Parameters **axes** (A single integer or a list of integers. If set to `None` (default) we return the limits for all axes.) – The axes for which we want to get the limits.

Returns A list of \((\text{min}_i, \text{max}_i)\) ndarrays.

**get_meshwidths**(axes=`None`)
Returns the meshwidths of the grid.

Parameters **axes** (A single integer or a list of integers. If set to `None` (default) we return the data for all axes.) – The axes for which we want to get the meshwidths.

Returns A list of \(h_i\) values or a single value.

**get_nodes**(flat=`True`, split=`False`)
Returns all grid nodes of the full tensor product grid.

Parameters

* **flat** (Boolean, default is `True`.) – Whether to return the grid with a hypercubic \((D, N_1, ..., N_D)\) shape or a flat \((D, \prod_i D N_i)\) shape.

* **split** (Boolean, default is `False`.) – Whether to return the different components, one for each dimension inside a single ndarray or a list with ndarrays, with one item per dimension.

Returns Depends of the optional arguments.

**get_number_nodes**(axes=`None`, overall=`False`)
Returns the number of grid nodes along a set of axes.

Parameters

* **axes** (A single integer or a list of integers. If set to `None` (default) we return the data for all axes.) – The axes for which we want to get the number of nodes.

* **overall** (Boolean, default is `False`) – Compute the product \(\prod_i D N_i\) of the number \(N_i\) of grid nodes along each axis \(i\) specified.

Returns A list of \(N_i\) values or a single value \(N\).

**is_regular()**
Answers the question if the grid spacing is regular. (It can still be different along each axis!)
GridWrapper

About the GridWrapper class

The WaveBlocks Project

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Inheritance diagram

```
Grid.Grid  GridWrapper.GridWrapper
```

Class documentation

class WaveBlocksND.GridWrapper(anarray)

This class constructs a thin layer around an ndarray and wraps it as Grid subclass for API compatibility. The array must have a shape of \((D, N)\) with \(N\) the overall number of nodes.

Rather than using this class, one should try to elliminate the cases where it is used now.

```
get_dimension()

Return the dimension \(D\) of the grid.
```

```
get_nodes(flat=True, split=False)

Returns all grid nodes.
```

Parameters

- **flat** (Boolean, default is True.) – Whether to return the grid with a hypercubic \((D, N_1, ..., N_D)\) shape or a flat \((D, \prod_i^D N_i)\) shape. Note that the hypercubic shape is not implemented!

- **split** (Boolean, default is False.) – Whether to return the different components, one for each dimension inside a single ndarray or a list with ndarrays, with one item per dimension.

Returns Depends of the optional arguments.

```
get_number_nodes(overall=False)

Returns the number of grid nodes.
```

Parameters **overall** (Boolean, default is False) – Compute the product \(N = \prod_i^D N_i\) of the number \(N_i\) of grid nodes along each dimension \(i\) specified.

Returns A list of \(N_i\) values or a single value \(N\).
QuadratureRule

About the QuadratureRule class

The WaveBlocks Project

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Inheritance diagram

```
QuadratureRule.QuadratureRule
```

Class documentation

```python
class WaveBlocksND.QuadratureRule
    This class is an abstract interface to quadrature rules in general.

    get_description()
        Return a description of this quadrature rule object. A description is a dict containing all key-value pairs necessary to reconstruct the current instance. A description never contains any data.

    get_dimension()
        Returns The space dimension $D$ of the quadrature rule.

    get_nodes()
        Returns A two-dimensional ndarray containing the quadrature nodes $\gamma_i$. The array must have a shape of $(D, |\Gamma|)$.

    get_number_nodes()
        Returns The number of quadrature nodes denoted by $|\Gamma|$ that are part of this quadrature rule $\Gamma = (\gamma, \omega)$.

    get_order()
        Returns The order $R$ of the quadrature rule.

    get_weights()
        Returns A two-dimensional ndarray containing the quadrature weights $\omega_i$. The array must have a shape of $(1, |\Gamma|)$.
```

GaussHermiteQR

About the GaussHermiteQR class

The WaveBlocks Project
class WaveBlocksND.GaussHermiteQR(order)
This class implements a Gauss-Hermite quadrature rule.

get_description()
Return a description of this quadrature rule object. A description is a dict containing all key-value pairs necessary to reconstruct the current instance. A description never contains any data.

get_dimension()
Returns The space dimension $D$ of the quadrature rule.

get_nodes()
Returns the quadrature nodes $\gamma_i$.

get_number_nodes()
Returns The number of quadrature nodes denoted by $|\Gamma|$ that are part of this quadrature rule $\Gamma = (\gamma, \omega)$.

get_order()
Returns The order $R$ of the quadrature rule.

get_weights()
Returns the quadrature weights $\omega_i$.

About the TensorProductQR class

The WaveBlocks Project
Class documentation

```python
class WaveBlocksND.TensorProductQR(rules):
    This class implements the construction of high dimensional quadrature rules from one-dimensional ones by
taking tensor products.

    get_description()
    Return a description of this quadrature rule object. A description is a dict containing all key-value pairs
necessary to reconstruct the current instance. A description never contains any data.

    get_dimension()
    Returns The space dimension \( D \) of the quadrature rule.

    get_nodes(flat=True, split=False)
    Returns the quadrature nodes \( \gamma_i \).

    Parameters
    • flat (Boolean, default is True.) – Dummy parameter for API compatibility with Grids.
    • split (Boolean, default is False.) – Dummy parameter for API compatibility with Grids.

    Returns An ndarray containing the quadrature nodes \( \gamma_i \).

    get_number_nodes()
    Returns The number of quadrature nodes denoted by \( |\Gamma| \) that are part of this quadrature rule
\( \Gamma = (\gamma, \omega) \).

    get_order()
    Returns The order \( R \) of the quadrature rule.

    get_weights()
    Returns the quadrature weights \( \omega_i \).
```

MatrixExponential

About the MatrixExponential functions

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Class documentation

The WaveBlocks Project

This file contains several different algorithms to compute the matrix exponential. Currently we have an exponential based on Pade approximations and an Arnoldi iteration method.

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MatrixExponential. **arnoldi** \((A, v0, k)\)

Arnoldi algorithm to compute the Krylov approximation \(H\) of a matrix \(A\).

**Parameters**

- \(A\) – The matrix \(A\) of shape \(N \times N\) to approximate.
- \(v0\) – The initial vector \(v0\) of length \(N\). (Should be in matrix shape \((N, 1)\) for practical reasons.)
- \(k\) – The number \(k\) of Krylov steps performed.

**Returns** A tuple \((V, H)\) where \(V\) is the large matrix of shape \(N \times k\) containing the orthogonal vectors and \(H\) is the small matrix of shape \(k \times k\) containing the Krylov approximation of \(A\).

MatrixExponential. **matrix_exp_arnoldi** \((A, v, factor, k)\)

Compute the solution of \(v' = Av\) via \(k\) steps of a the Arnoldi krylov method.

**Parameters**

- \(A\) – The matrix \(A\) of shape \(N \times N\).
- \(v\) – The vector \(v\) of length \(N\).
- \(factor\) – An additional scalar factor \(\alpha\).
- \(k\) – The number \(k\) of Krylov steps performed.

**Returns** The (approximate) value of \(\exp (-i\alpha A) v\).

MatrixExponential. **matrix_exp_pade** \((A, v, factor)\)

Compute the solution of \(v' = Av\) with a full matrix exponential via Pade approximation.

**Parameters**

- \(A\) – The matrix \(A\) of shape \(N \times N\).
- \(v\) – The vector \(v\) of length \(N\).
- \(factor\) – An additional scalar factor \(\alpha\).

**Returns** The (approximate) value of \(\exp (-i\alpha A) v\)

Utils

**About the utils class**

The WaveBlocks Project

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Class documentation

The WaveBlocks Project

Various small utility functions.

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Utils.meshgrid_nd(arrays)
   Like “meshgrid()” but for arbitrary number of dimensions.

1.1.2 Basic quantum mechanics

WaveFunction

About the WaveFunction class

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Inheritance diagram

```
WaveFunction.WaveFunction
WaveFunction.WaveFunction
```

Class documentation

class WaveBlocksND.WaveFunction(parameters)
   This class represents a vector valued wavefunction $\Psi$ as used in the vector valued time-dependent Schröedinger equation. The state $\Psi$ consists of the components $\psi_0$ to $\psi_{N-1}$.

   get_grid()
      Return the Grid instance representing the grid $\Gamma$. The wavefunction $\Psi$ is evaluated on the grid nodes to get $\Psi(\Gamma)$.

   get_number_components()
      The number of components $\psi_i$ the vector $\Psi$ consists of.

   get_values(components=None)
      Get the wavefunction values $\psi(\Gamma)$ for each component $\psi_i$ of $\Psi$.

      Parameters components (A single integer or a list of integers. If set to None (default) we return the data for all components.) – The components $i$ for which we want to get the wavefunction values $\psi_i$.

      Returns A list of the values $\psi_i$ for all components $i$. 

Chapter 1. Source code documentation
**kinetic_energy** *(kinetic, summed=False)*
Calculate the kinetic energy \( E_{\text{kin}} := \langle \Psi | T | \Psi \rangle \) of the different components \( \psi_i \).

**Parameters**
- **kinetic** *(A KineticOperator instance.)* – The kinetic energy operator \( T(\omega) \).
- **summed** – Whether to sum up the kinetic energies \( E_i \) of the individual components \( \psi_i \). Default is \( False \).

**Returns** A list with the kinetic energies of the individual components or the overall kinetic energy of the wavefunction. (Depending on the optional arguments.)

**norm** *(values=None, summed=False, components=None)*
Calculate the \( L^2 \) norm of the whole vector values wavefunction \( \Psi \) or some individual components \( \psi_i \). The calculation is done in momentum space.

**Parameters**
- **values** – Allows to use this function for external data, similar to a static function.
- **summed** *(Boolean, default is \( False \).)* – Whether to sum up the norms of the individual components.
- **components** *(A single integer or a list of integers. If set to \( None \) (default) we compute the norm for all components.)* – The components \( \psi_i \) of which the norms are calculated.

**Returns** The \( L^2 \) norm of \( \Psi \) or a list of \( L^2 \) norms of the specified components \( \psi_i \).

**potential_energy** *(potential, summed=False)*
Calculate the potential energy \( E_{\text{pot}} := \langle \Psi | V | \Psi \rangle \) of the different components \( \psi_i \).

**Parameters**
- **potential** – The potential energy operator \( V(x) \).
- **summed** – Whether to sum up the potential energies \( E_i \) of the individual components \( \psi_i \). Default is \( False \).

**Returns** A list with the potential energies of the individual components or the overall potential energy of the wavefunction. (Depending on the optional arguments.)

**set_grid** *(grid)*
Assign a new grid \( \Gamma \) to this WaveFunction instance.

*Note:* The user of this class has to make sure that the grid \( \Gamma \) and the wavefunction values \( \Psi(\Gamma) \) are consistent with each other!

**Parameters** **grid** – A new Grid instance.

**set_values** *(values, components=None)*
Assign new wavefunction values \( \psi_i(\Gamma) \) for each component \( i \) of \( \Psi \) to the current WaveFunction instance.

**Parameters**
- **values** *(Each entry of the list has to be an ndarray.)* – A list with the new values of all components we want to change.
- **components** *(A single integer or a list of integers. If set to \( None \) (default) we set the data for all components.)* – The components \( i \) for which we want to set the new wavefunction values \( \psi_i \).

*Note:* This method does NOT copy the input data arrays.

**Raises** ValueError If the list of **values** has the wrong length.
MatrixPotential

About the MatrixPotential class

The WaveBlocks Project

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Inheritance diagram

Class documentation

class WaveBlocksND.MatrixPotential
    This class represents a potential $V(x)$ with $x \in \mathbb{R}^D$. The potential is given as an analytic expression. Some calculations with the potential are supported. For example calculation of eigenvalues $\lambda_i(x)$ and eigenvectors $\nu_i(x)$ and numerical evaluation on a grid $\Gamma$.

    calculate_eigenvalues()
        Calculate the eigenvalue $\lambda_0(x)$ of the potential $V(x)$.
        Raises NotImplementedError This is an abstract base class.

    calculate_eigenvectors()
        Calculate the eigenvectors $\nu_i(x)$ of the potential $V(x)$.
        Raises NotImplementedError This is an abstract base class.

    calculate_exponential(factor=1)
        Calculate the matrix exponential $\exp(\alpha V)$.
        Parameters factor – The prefactor $\alpha$ in the exponential.
        Raises NotImplementedError This is an abstract base class.

    evaluate_at(grid, entry=None)
        Evaluate the potential $V(x)$ elementwise on a grid $\Gamma$.
        Parameters
            • grid – The grid containing the nodes $\gamma_i$ we want to evaluate the potential at.
            • entry – The indices $(i, j)$ of the component $V_{i,j}(x)$ we want to evaluate or None to evaluate all entries.
        Raises NotImplementedError This is an abstract base class.

    evaluate_eigenvalues_at(grid, entry=None)
        Evaluate the eigenvalues $\Lambda(x)$ elementwise on a grid $\Gamma$. 
Parameters

- **grid** – The grid containing the nodes \( \gamma_i \) we want to evaluate the eigenvalues at.
- **entry** – The index \( i \) of the eigenvalue \( \lambda_i(x) \) we want to evaluate or \( \text{None} \) to evaluate all eigenvalues.

Raises **NotImplementedError** This is an abstract base class.

**evaluate_eigenvectors_at** (**grid**, **entry=None**)  
Evaluate the eigenvectors \( \nu_i(x) \) elementwise on a grid \( \Gamma \).

Parameters

- **grid** – The grid containing the nodes \( \gamma_i \) we want to evaluate the eigenvectors at.
- **entry** – The index \( i \) of the eigenvector \( \nu_i(x) \) we want to evaluate or \( \text{None} \) to evaluate all eigenvectors.

Raises **NotImplementedError** This is an abstract base class.

**evaluate_exponential_at** (**grid**)  
Evaluate the exponential of the potential matrix \( V(x) \) on a grid \( \Gamma \).

Parameters **grid** – The grid containing the nodes \( \gamma_i \) we want to evaluate the exponential at.

Raises **NotImplementedError** This is an abstract base class.

**get_dimension** ()  
Return the dimension \( D \) of the potential \( V(x) \). The dimension is equal to the number of free variables \( x_i \) where \( x := (x_1, x_2, ..., x_D) \).

**get_number_components** ()  
Return the number \( N \) of components the potential \( V(x) \) supports. This is equivalent to the number of energy levels \( \lambda_i(x) \).

**MatrixPotential1S**

About the **MatrixPotential1S** class

The WaveBlocks Project

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Inheritance diagram

```
| MatrixPotential.MatrixPotential | MatrixPotential1S.MatrixPotential1S |
```

1.1. WaveBlocks Classes
Class documentation

class WaveBlocksND.MatrixPotential1S(expression, variables, **kwargs)
This class represents a scalar potential $V(x)$. The potential is given as an analytic $1 \times 1$ matrix expression. Some symbolic calculations with the potential are supported.

calculate_eigenvalues()
Calculate the eigenvalue $\lambda_0(x)$ of the potential $V(x)$. In the scalar case this is just equal to the matrix entry $V_{0,0}(x)$. Note: This function is idempotent and the eigenvalues are memoized for later reuse.

calculate_eigenvectors()
Calculate the eigenvector $\nu_0(x)$ of the potential $V(x)$. In the scalar case this is just the value 1. Note: This function is idempotent and the eigenvectors are memoized for later reuse.

calculate_exponential(factor=1)
Calculate the matrix exponential $\exp(\alpha V)$. In the case of this class the matrix is of size $1 \times 1$ thus the exponential simplifies to the scalar exponential function. Note: This function is idempotent.

Parameters

- **factor** (The prefactor $\alpha$ in the exponential.)

calculate_hessian()
Calculate the Hessian matrix $\nabla^2 V$ of the potential $V(x)$ with $x \in \mathbb{R}^D$. For potentials which depend only one variable, this equals the second derivative and $D = 1$. Note that this function is idempotent.

calculate_jacobian()
Calculate the Jacobian matrix $\nabla V$ of the potential $V(x)$ with $x \in \mathbb{R}^D$. For potentials which depend only one variable, this equals the first derivative and $D = 1$. Note that this function is idempotent.

calculate_local_quadratic(diagonal_component=None)
Calculate the local quadratic approximation $U(x)$ of the potential’s eigenvalue $\lambda(x)$. Note that this function is idempotent.

Parameters

- **diagonal_component** (Dummy parameter that has no effect here.)

calculate_local_remainder(diagonal_component=None)
Calculate the non-quadratic remainder $W(x) = V(x) - U(x)$ of the quadratic Taylor approximation $U(x)$ of the potential’s eigenvalue $\lambda(x)$. Note that this function is idempotent.

Parameters

- **diagonal_component** (Dummy parameter that has no effect here.)

evaluate_at(grid, entry=None, as_matrix=False)
Evaluate the potential $V(x)$ elementwise on a grid $\Gamma$.

Parameters

- **grid** (A Grid instance. (Numpy arrays are not directly supported yet.)) – The grid containing the nodes $\gamma_i$ we want to evaluate the potential at.
- **entry** (A python tuple of two integers.) – The indices $(i,j)$ of the component $V_{i,j}(x)$ we want to evaluate or None to evaluate all entries. This has no effect here as we only have a single entry $V_{0,0}$.
- **as_matrix** – Dummy parameter which has no effect here.

Returns A list containing a single numpy ndarray of shape $(1,|\Gamma|)$.

evaluate_eigenvalues_at(grid, entry=None, as_matrix=False)
Evaluate the eigenvalue $\lambda_0(x)$ elementwise on a grid $\Gamma$.

Parameters

- **grid** (A Grid instance. (Numpy arrays are not directly supported yet.)) – The grid containing the nodes $\gamma_i$ we want to evaluate the eigenvalue at.
• **entry** (*A python tuple of two integers.*) – The indices \((i, j)\) of the component \(\Lambda_{i,j}(x)\) we want to evaluate or *None* to evaluate all entries. If \(j = i\) then we evaluate the eigenvalue \(\lambda_i(x)\). This has no effect here as we only have a single entry \(\lambda_0\).

• **as_matrix** – Dummy parameter which has no effect here.

**Returns** A list containing a single numpy ndarray of shape \((N_1, \ldots, N_D)\).

### evaluate_eigenvectors_at(grid, entry=None)
Evaluate the eigenvector \(v_0(x)\) elementwise on a grid \(\Gamma\).

**Parameters**
- **grid** (*A Grid instance. (Numpy arrays are not directly supported yet.)*) – The grid containing the nodes \(\gamma_i\) we want to evaluate the eigenvector at.
- **entry** (*A singly python integer.*) – The index \(i\) of the eigenvector \(v_i(x)\) we want to evaluate or *None* to evaluate all eigenvectors. This has no effect here as we only have a single entry \(v_0\).

**Returns** A list containing the numpy ndarrays, all of shape \((1, |\Gamma|)\).

### evaluate_exponential_at(grid, entry=None)
Evaluate the exponential of the potential matrix \(V(x)\) on a grid \(\Gamma\).

**Parameters**
- **grid** (*A Grid instance. (Numpy arrays are not directly supported yet.)*) – The grid containing the nodes \(\gamma_i\) we want to evaluate the exponential at.

**Returns** The numerical approximation of the matrix exponential at the given grid nodes.

### evaluate_hessian_at(grid, component=None)
Evaluate the potential’s Hessian \(\nabla^2 V(x)\) at some grid nodes \(\Gamma\).

**Parameters**
- **grid** – The grid nodes \(\Gamma\) the Hessian gets evaluated at.
- **component** – Dummy parameter that has no effect here.

**Returns** The value of the potential’s Hessian at the given nodes. The result is an *ndarray* of shape \((D, D)\) if we evaluate at a single grid node or of shape \((|\Gamma|, D, D)\) if we evaluate at multiple nodes simultaneously.

### evaluate_jacobian_at(grid, component=None)
Evaluate the potential’s Jacobian \(\nabla V(x)\) at some grid nodes \(\Gamma\).

**Parameters**
- **grid** – The grid nodes \(\Gamma\) the Jacobian gets evaluated at.
- **component** – Dummy parameter that has no effect here.

**Returns** The value of the potential’s Jacobian at the given nodes. The result is an *ndarray* of shape \((D, 1)\) if we evaluate at a single grid node or of shape \((|\Gamma|, D)\) if we evaluate at multiple nodes simultaneously.

### evaluate_local_quadratic_at(grid, diagonal_component=None)
Numerically evaluate the local quadratic approximation \(U(x)\) of the potential’s eigenvalue \(\lambda(x)\) at the given grid nodes \(\Gamma\). This function is used for the homogeneous case.

**Parameters**
- **grid** – The grid nodes \(\Gamma\) the quadratic approximation gets evaluated at.
- **diagonal_component** – Dummy parameter that has no effect here.
Returns

A list containing the values $V(\Gamma)$, $\nabla V(\Gamma)$ and $\nabla^2 V(\Gamma)$.

evaluate_local_remainder_at (grid, position, diagonal_component=None, entry=None)

Numerically evaluate the non-quadratic remainder $W(x)$ of the quadratic approximation $U(x)$ of the potential's eigenvalue $\lambda(x)$ at the given nodes $\Gamma$.

Parameters

- grid – The grid nodes $\Gamma$ the remainder $W$ gets evaluated at.
- position – The point $q \in \mathbb{R}^D$ where the Taylor series is computed.
- diagonal_component – Dummy parameter that has no effect here.
- entry – Dummy parameter that has no effect here.

Returns

A list with a single entry consisting of an `ndarray` containing the values of $W(\Gamma)$.

The array is of shape $(1, |\Gamma|)$.

get_dimension ()

Return the dimension $D$ of the potential $V(x)$. The dimension is equal to the number of free variables $x_i$ where $x := (x_1, x_2, ..., x_D)$.

get_number_components ()

Return the number $N$ of components the potential $V(x)$ supports. This is equivalent to the number of energy levels $\lambda_i(x)$.

MatrixPotential2S

About the MatrixPotential2S class

The WaveBlocks Project

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Inheritance diagram

Class documentation

class WaveBlocksND.MatrixPotential2S (expression, variables, **kwargs)

This class represents a matrix potential $V(x)$. The potential is given as an analytic $2 \times 2$ matrix expression. Some symbolic calculations with the potential are supported.

calculate_eigenvalues ()

Calculate the two eigenvalues $\lambda_i(x)$ of the potential $V(x)$. We can do this by symbolic calculations. The multiplicities are taken into account. Note: This function is idempotent and the eigenvalues are memoized for later reuse.
calculate_eigenvectors()
Calculate the two eigenvectors $\nu_i(x)$ of the potential $V(x)$. We can do this by symbolic calculations. Note: This function is idempotent and the eigenvectors are memoized for later reuse.

calculate_exponential (factor=1)
Calculate the matrix exponential $\exp(\alpha V)$. In the case of this class the matrix is of size $2 \times 2$ thus the exponential can be calculated analytically for a general matrix. Note: This function is idempotent.

Parameters factor – The prefactor $\alpha$ in the exponential.

calculate_hessian()
Calculate the Hessian matrix $\nabla^2 \lambda(x)$ of the potential’s eigenvalues $\Lambda(x)$ with $x \in \mathbb{R}^D$. For potentials which depend only one variable, this equals the second derivative and $D = 1$. Note that this function is idempotent.

calculate_jacobian()
Calculate the Jacobian matrix $\nabla \lambda(x)$ of the potential’s eigenvalues $\Lambda(x)$ with $x \in \mathbb{R}^D$. For potentials which depend only one variable, this equals the first derivative and $D = 1$. Note that this function is idempotent.

calculate_local_quadratic (diagonal_component=None)
Calculate the local quadratic approximation matrix $U(x)$ of the potential’s eigenvalues in $\Lambda(x)$. This function can be used for the homogeneous case and takes into account the leading component $\chi \in [0, \ldots, N - 1]$. If the parameter $i$ is not given, calculate the local quadratic approximation matrix $U(x)$ of all the potential’s eigenvalues in $\Lambda$. This case can be used for the inhomogeneous case.

Parameters diagonal_component (Integer or None (default)) – Specifies the index $i$ of the eigenvalue $\lambda_i$ that gets expanded into a Taylor series $u_i$.

calculate_local_remainder (diagonal_component=None)
Calculate the non-quadratic remainder matrix $W(x) = V(x) - U(x)$ of the quadratic approximation matrix $U(x)$ of the potential’s eigenvalue matrix $\Lambda(x)$. In the homogeneous case the matrix $U$ is given by $U(x) = \text{diag}([u_0, \ldots, u_{i-1}])$ where in the inhomogeneous case it is given by $U(x) = \text{diag}([u_0, \ldots, u_{N-1}])$.

Parameters diagonal_component (Integer or None (default)) – Specifies the index $i$ of the eigenvalue $\lambda_i$ that gets expanded into a Taylor series $u_i$. If set to None the inhomogeneous case is computed.

evaluate_at (grid, entry=None, as_matrix=True)
Evaluate the potential $V(x)$ elementwise on a grid $\Gamma$.

Parameters
- grid (A Grid instance. (Numpy arrays are not directly supported yet.)) – The grid containing the nodes $\gamma_i$ we want to evaluate the potential at.
- entry (A python tuple of two integers.) – The indices $(i, j)$ of the component $V_{i,j}(x)$ we want to evaluate or None to evaluate all entries.
- as_matrix – Dummy parameter which has no effect here.

Returns A list containing 4 numpy ndarrays of shape $(1, |\Gamma|)$.

evaluate_eigenvalues_at (grid, entry=None, as_matrix=False)
Evaluate the eigenvalues $\lambda_i(x)$ elementwise on a grid $\Gamma$.

Parameters
- grid (A Grid instance. (Numpy arrays are not directly supported yet.)) – The grid $\Gamma$ containing the nodes $\gamma_i$ we want to evaluate the eigenvalues at.
- entry (A python tuple of two integers.) – The indices $(i, j)$ of the component $\Lambda_{i,j}(x)$ we want to evaluate or None to evaluate all entries. If $j = i$ then we evaluate the eigenvalue $\lambda_i(x)$.
• \texttt{as\_matrix} – Whether to include the off-diagonal zero entries of $\Lambda_{i,j}(x)$ in the return value.

**Returns** A list containing the numpy ndarrays, all of shape $(1, |\Gamma|)$.

\subsection*{evaluate\_eigenvectors\_at \texttt{(grid, entry=None)}}
Evaluate the two eigenvectors $\nu_i(x)$ elementwise on a grid $\Gamma$.

**Parameters**

• \texttt{grid} (A \texttt{Grid} instance. (Numpy arrays are not directly supported yet.)) – The grid containing the nodes $\gamma_i$ we want to evaluate the eigenvectors at.

• \texttt{entry} (A singly python integer.) – The index $i$ of the eigenvector $\nu_i(x)$ we want to evaluate or \texttt{None} to evaluate all eigenvectors.

**Returns** A list containing the numpy ndarrays, all of shape $(N, |\Gamma|)$.

\subsection*{evaluate\_exponential\_at \texttt{(grid)}}
Evaluate the exponential of the potential matrix $V(x)$ on a grid $\Gamma$.

**Parameters** \texttt{grid} (A \texttt{Grid} instance. (Numpy arrays are not directly supported yet.)) – The grid containing the nodes $\gamma_i$ we want to evaluate the exponential at.

**Returns** The numerical approximation of the matrix exponential at the given grid nodes. A list contains the exponentials for all entries $(i,j)$, each having the same shape as the grid.

\subsection*{evaluate\_hessian\_at \texttt{(grid, component=None)}}
Evaluate the list of Hessian matrices $\nabla^2 \lambda_i(x)$ at some grid nodes $\Gamma$.

**Parameters**

• \texttt{grid} (A \texttt{Grid} instance. (Numpy arrays are not directly supported yet.)) – The grid nodes $\Gamma$ the Hessian gets evaluated at.

• \texttt{component} – Dummy parameter that has no effect here.

**Returns** The value of the potential’s Hessian at the given nodes. The result is an ndarray of shape $(D,D)$ is we evaluate at a single grid node or of shape $(|\Gamma|, D, D)$ if we evaluate at multiple nodes simultaneously.

\subsection*{evaluate\_jacobian\_at \texttt{(grid, component=None)}}
Evaluate the list of Jacobian matrices $\nabla \lambda_i(x)$ at some grid nodes $\Gamma$.

**Parameters**

• \texttt{grid} (A \texttt{Grid} instance. (Numpy arrays are not directly supported yet.)) – The grid nodes $\Gamma$ the Jacobian gets evaluated at.

• \texttt{component} – Dummy parameter that has no effect here.

**Returns** The value of the potential’s Jacobian at the given nodes. The result is a list of ndarrays each of shape $(D, D)$ is we evaluate at a single grid node or of shape $(|\Gamma|, D)$ if we evaluate at multiple nodes simultaneously.

\subsection*{evaluate\_local\_quadratic\_at \texttt{(grid, diagonal\_component=None)}}
Numerically evaluate the local quadratic approximation matrix $U(x)$ of the potential’s eigenvalues in $\Lambda(x)$ at the given grid nodes $\Gamma$.

**Parameters**

• \texttt{grid} (A \texttt{Grid} instance. (Numpy arrays are not directly supported yet.)) – The grid $\Gamma$ containing the nodes $\gamma$ we want to evaluate the quadratic approximation at.

• \texttt{diagonal\_component} – Specifies the index $i$ of the eigenvalue $\lambda_i$ that gets expanded into a Taylor series $u_i$. 
Returns A list of tuples or a single tuple. Each tuple \((\lambda, J, H)\) contains the the evaluated eigenvalues \(\lambda_i(\Gamma)\), the Jacobian \(J(\Gamma)\) and the Hessian \(H(\Gamma)\) in this order.

evaluate_local_remainder_at \((grid, position, diagonal\_component=None, entry=None)\)
Numerically evaluate the non-quadratic remainder \(W(x)\) of the quadratic approximation \(U(x)\) of the potential’s eigenvalue \(\Lambda(x)\) at the given nodes \(\Gamma\).

Warning: do not set the diagonal\_component and the entry parameter both to None.

Parameters
- \textbf{grid} – The grid nodes \(\Gamma\) the remainder \(W\) gets evaluated at.
- \textbf{position} – The point \(q \in \mathbb{R}^D\) where the Taylor series is computed.
- \textbf{diagonal\_component} (Integer or None (default)) – Specifies the index \(i\) of the eigenvalue \(\lambda_i\) that gets expanded into a Taylor series \(u_i\) and whose remainder matrix \(W(x) = V(x) - \text{diag}([u_i, \ldots, u_i])\) we evaluate. If set to None the inhomogeneous case given by \(W(x) = V(x) - \text{diag}([u_0, \ldots, u_{N-1}])\) is computed.
- \textbf{entry} (A python tuple of two integers.) – The entry \((i, j)\) of the remainder matrix \(W\) that is evaluated.

Returns A list with \(N^2\) ndarray elements or a single \ndarray. Each containing the values of \(W_{i,j}(\Gamma)\). Each array is of shape \((1, |\Gamma|)\).

get\_dimension ()
Return the dimension \(D\) of the potential \(V(x)\). The dimension is equal to the number of free variables \(x_i\) where \(x := (x_1, x_2, \ldots, x_D)\).

get\_number\_components ()
Return the number \(N\) of components the potential \(V(x)\) supports. This is equivalent to the number of energy levels \(\lambda_i(x)\).

MatrixPotentialMS

About the MatrixPotentialMS class

The WaveBlocks Project
@author: R. Bourquin @copyright: Copyright (C) 2010, 2011, 2012 R. Bourquin @license: Modified BSD License

Inheritance diagram

Class documentation

class WaveBlocksND.MatrixPotentialMS (expression, variables, **kwargs)
This class represents a matrix potential \(V(x)\). The potential is given as an analytic \(N \times N\) matrix expression.
All methods use pure numerical techniques because symbolical calculations are unfeasible for 3 or more energy levels.

**calculate_eigenvalues()**
Calculate all the eigenvalues $\lambda_i(x)$ of the potential $V(x)$. We can not do this by symbolic calculations, hence the function has an empty implementation. We compute the eigenvalues by numerical techniques in the corresponding `evaluate_eigenvalues_at` function.

**calculate_eigenvectors()**
Calculate all the eigenvectors $\nu_i(x)$ of the potential $V(x)$. We can not do this by symbolic calculations, hence the function has an empty implementation. We compute the eigenvectors by numerical techniques in the corresponding `evaluate_eigenvectors_at` function.

**calculate_exponential**(factor=\(1\))
Calculate the matrix exponential $\exp(\alpha V)$. In the case of this class the matrix is of size $N \times N$ thus the exponential can not be calculated analytically for a general matrix. We use numerical approximations to determine the matrix exponential. We just store the prefactor $\alpha$ for use during numerical evaluation.

**Parameters**
- **factor** – The prefactor $\alpha$ in the exponential.

**calculate_hessian()**
Calculate the Hessian matrix $\nabla^2 \lambda_i$ of the potential’s eigenvalues $\Lambda(x)$ with $x \in \mathbb{R}^D$. For potentials which depend only one variable, this equals the second derivative and $D = 1$. Note that this function is idempotent.

**calculate_jacobian()**
Calculate the Jacobian matrix $\nabla \lambda_i$ of the potential’s eigenvalues $\Lambda(x)$ with $x \in \mathbb{R}^D$. For potentials which depend only one variable, this equals the first derivative and $D = 1$. Note that this function is idempotent.

**calculate_local_quadratic**(diagonal_component=None)
Calculate the local quadratic approximation matrix $U(x)$ of the potential’s eigenvalues in $\Lambda(x)$. This function can be used for the homogeneous case and takes into account the leading component $\chi \in [0, \ldots, N - 1]$. If the parameter $i$ is not given, calculate the local quadratic approximation matrix $U(x)$ of all the potential’s eigenvalues in $\Lambda$. This case can be used for the inhomogeneous case.

**Parameters**
- **diagonal_component** – Dummy parameter which has no effect here.

**calculate_local_remainder**(diagonal_component=None)
Calculate the non-quadratic remainder matrix $W(x) = V(x) - U(x)$ of the quadratic approximation matrix $U(x)$ of the potential’s eigenvalue matrix $\Lambda(x)$. In the homogeneous case the matrix $U$ is given by $U(x) = \text{diag}([u_i, \ldots, u_i])$ where in the inhomogeneous case it is given by $U(x) = \text{diag}([u_0, \ldots, u_{N-1}])$.

**Parameters**
- **diagonal_component** (Integer or None (default)) – Specifies the index $i$ of the eigenvalue $\lambda_i$ that gets expanded into a Taylor series $u_i$. If set to None the inhomogeneous case is computed.

**evaluate_at**(grid, entry=None, as_matrix=True)
Evaluate the potential $V(x)$ elementwise on a grid $\Gamma$.

**Parameters**
- **grid** (A Grid instance. (Numpy arrays are not directly supported yet.)) – The grid containing the nodes $\gamma_i$ we want to evaluate the potential at.
- **entry** (A python tuple of two integers.) – The indices $(i, j)$ of the component $V_{i,j}(x)$ we want to evaluate or None to evaluate all entries.
- **as_matrix** – Dummy parameter which has no effect here.

**Returns**
A list containing $N^2$ numpy ndarrays of shape $(1, |\Gamma|)$.
evaluate_eigenvalues_at \( (grid, entry=None, as\_matrix=False, sorted=True) \)
Evaluate the eigenvalues \( \lambda_i(x) \) elementwise on a grid \( \Gamma \).

**Parameters**
- grid (A Grid instance. (Numpy arrays are not directly supported yet.)) – The grid containing the nodes \( \gamma_i \) we want to evaluate the eigenvalues at.
- entry (A python tuple of two integers.) – The indices \( (i, j) \) of the component \( \Lambda_{i,j}(x) \) we want to evaluate or None to evaluate all entries. If \( j=i \) then we evaluate the eigenvalue \( \lambda_i(x) \).
- as\_matrix – Whether to include the off-diagonal zero entries of \( \Lambda_{i,j}(x) \) in the return value.

**Returns** A list containing the numpy ndarrays, all of shape \( (1, |\Gamma|) \).

evaluate_eigenvectors_at \( (grid, sorted=True) \)
Evaluate the eigenvectors \( \nu_i(x) \) elementwise on a grid \( \Gamma \).

**Parameters**
- grid (A Grid instance. (Numpy arrays are not directly supported yet.)) – The grid containing the nodes \( \gamma_i \) we want to evaluate the eigenvectors at.

**Returns** A list containing the \( N \) numpy ndarrays, all of shape \( (D, |\Gamma|) \).

evaluate_exponential_at \( (grid) \)
Evaluate the exponential of the potential matrix \( V(x) \) on a grid \( \Gamma \).

**Parameters**
- grid (A Grid instance. (Numpy arrays are not directly supported yet.)) – The grid containing the nodes \( \gamma_i \) we want to evaluate the exponential at.

**Returns** The numerical approximation of the matrix exponential at the given grid nodes. A list contains the exponentials for all entries \( (i, j) \), each having a shape of \( (1, |\Gamma|) \).

evaluate_hessian_at \( (grid, component=None) \)
Evaluate the list of Hessian matrices \( \nabla^2 \lambda_i(x) \) at some grid nodes \( \Gamma \) for one or all eigenvalues.

**Parameters**
- grid (A Grid instance. (Numpy arrays are not directly supported yet.)) – The grid nodes \( \Gamma \) the Hessian gets evaluated at.
- component – The index \( i \) of the eigenvalue \( \lambda_i \).

**Returns** The value of the potential’s Hessian at the given nodes. The result is an ndarray of shape \( (D, D) \) is we evaluate at a single grid node or of shape \( (D, D, |\Gamma|) \) if we evaluate at multiple nodes simultaneously.

evaluate_jacobian_at \( (grid, component=None) \)
Evaluate the list of Jacobian matrices \( \nabla \lambda_i(x) \) at some grid nodes \( \Gamma \) for one or all eigenvalues.

**Parameters**
- grid (A Grid instance. (Numpy arrays are not directly supported yet.)) – The grid nodes \( \Gamma \) the Jacobian gets evaluated at.
- component – The index \( i \) of the eigenvalue \( \lambda_i \).

**Returns** The value of the potential’s Jacobian at the given nodes. The result is a list of ndarray each of shape \( (D, 1) \) is we evaluate at a single grid node or of shape \( (D, |\Gamma|) \) if we evaluate at multiple nodes simultaneously.

evaluate_local_quadratic_at \( (grid, diagonal\_component=None) \)
Numerically evaluate the local quadratic approximation matrix \( U(x) \) of the potential’s eigenvalues in \( \Lambda(x) \) at the given grid nodes \( \Gamma \).
Parameters

- **grid** (A Grid instance. (Numpy arrays are not directly supported yet.)) – The grid Γ containing the nodes γ we want to evaluate the quadratic approximation at.

- **diagonal_component** – Specifies the index i of the eigenvalue λ_i that gets expanded into a Taylor series u_i.

Returns A list of tuples or a single tuple. Each tuple (λ, J, H) contains the the evaluated eigenvalue λ_i(Γ), its Jacobian J(Γ) and its Hessian H(Γ) in this order.

evaluate_local_remainder_at (grid, position, diagonal_component=None, entry=None)
Numerically evaluate the non-quadratic remainder W(x) of the quadratic approximation U(x) of the potential’s eigenvalue Λ(x) at the given nodes Γ.

Warning: do not set the diagonal_component and the entry parameter both to None.

Parameters

- **grid** – The grid nodes Γ the remainder W gets evaluated at.

- **position** – The point q ∈ ℝ^D where the Taylor series is computed.

- **diagonal_component** (Integer or None (default)) – Specifies the index i of the eigenvalue λ_i that gets expanded into a Taylor series u_i and whose remainder matrix W(x) = V(x) − diag([u_i, ..., u_i]) we evaluate. If set to None the inhomogeneous case given by W(x) = V(x) − diag([u_0, ..., u_{N-1}]) is computed.

- **entry** (A python tuple of two integers.) – The entry (i, j) of the remainder matrix W that is evaluated.

Returns A list with N^2 ndarray elements or a single ndarray. Each containing the values of W_{i,j}(Γ). Each array is of shape (1, |Γ|).

get_dimension()
Return the dimension D of the potential V(x). The dimension is equal to the number of free variables x_i where x := (x_1, x_2, ..., x_D).

get_number_components()
Return the number N of components the potential V(x) supports. This is equivalent to the number of energy levels λ_i(x).

1.1.3 Wavepackets

BasisShape

About the BasisShape class

The WaveBlocks Project

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Class documentation

class WaveBlocksND.BasisShape
This class defines the abstract interface to basis shapes. A basis shape is essentially all information and operations related to the set \( K \) of multi-indices \( k \).

Basis shapes must be immutable objects.

contains \( (k) \)
Checks if a given multi-index \( k \) is part of the basis set \( K \).

Parameters
\[ k \text{ (tuple)} \] – The multi-index we want to test.

Raises
NotImplementedError Abstract interface.

get_basis_size \( (\text{extended=False}) \)
Returns the size \( |K| \) of the basis. The size is the number of distinct multi-indices \( k \) that belong to the basis \( K \).

get_description()
Return a description of this basis shape object. A description is a dict containing all key-value pairs necessary to reconstruct the current basis shape. A description never contains any data.

get_dimension()
Returns the dimension \( D \) of the basis shape \( K \). This is defined as the number of components each multi-index \( k = (k_0, \ldots, k_{D-1}) \) has.

get_neighbours \( (k, \text{direction=None}) \)
Returns a list of all multi-indices that are neighbours of a given multi-index \( k \). A direct neighbours is defined as \( (k_0, \ldots, k_d \pm 1, \ldots, k_{D-1}) \forall d \in [0 \ldots D - 1] \).

Parameters
\[ k \text{ (tuple)} \] – The multi-index of which we want to get the neighbours.

\[ \text{direction (int)} \] – The direction \( 0 \leq d < D \) in which we want to find the neighbours \( k \pm e_d \).

Returns
A list containing the pairs \( (d,k') \).

Raises
NotImplementedError Abstract interface.

get_node_iterator \( (\text{mode='lex'}) \)
Returns an iterator to iterate over all basis elements \( k \).

Parameters
\[ \text{mode (string)} \] – The mode by which we iterate over the indices. Default is ‘lex’ for lexicographical order. Supported is also ‘chain’, for the chain-like mode, details see the manual.

Raises
NotImplementedError Abstract interface.
HyperCubicShape

About the HyperCubicShape class

The WaveBlocks Project

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Inheritance diagram

```
BasisShape.BasisShape  -->  HyperCubicShape.HyperCubicShape
```

Class documentation

class WaveBlocksND.HyperCubicShape (limits)

This class implements the hypercubic basis shape which is the full dense basis set. A basis shape is essentially all information and operations related to the set $K$ of multi-indices $k$.

- **contains** ($k$)
  Checks if a given multi-index $k$ is part of the basis set $K$.
  
  **Parameters** $k$ (tuple) – The multi-index we want to test.

- **extend** ()
  Extend the basis shape such that (at least) all neighbours of all boundary nodes are included in the extended basis shape.

- **get_basis_size** (extended=False)
  Returns the size $|K|$ of the basis. The size is the number of distinct multi-indices $k$ that belong to the basis $K$.

- **get_description** ()
  Return a description of this basis shape object. A description is a dict containing all key-value pairs necessary to reconstruct the current basis shape. A description never contains any data.

- **get_dimension** ()
  Returns the dimension $D$ of the basis shape $K$. This is defined as the number of components each multi-index $k = (k_0, \ldots, k_{D-1})$ has.

- **get_limits** ()
  Returns the upper limit $K_d$ for all directions $d$. return: A tuple of the maximum of the multi-index in each direction.

- **get_neighbours** ($k$, selection=None, direction=None)
  Returns a list of all multi-indices that are neighbours of a given multi-index $k$. A direct neighbour is defined as $(k_0, \ldots, k_d \pm 1, \ldots, k_{D-1}) \forall d \in [0 \ldots D - 1]$.

  **Parameters**
• \(k\) (tuple) – The multi-index of which we want to get the neighbours.

• selection (string with fixed values forward, backward or all. The values all is equivalent to the value None (default).) –

• direction (int) – The direction \(0 \leq d < D\) in which we want to find the neighbours \(k \pm e_d\).

Returns A list containing the pairs \((d, k')\).

get_node_iterator (mode='lex', direction=None)
Returns an iterator to iterate over all basis elements \(k\).

Parameters

• mode (string) – The mode by which we iterate over the indices. Default is ‘lex’ for lexicographical order. Supported is also ‘chain’, for the chain-like mode, details see the manual.

• direction (integer) – If iterating in chainmode this specifies the direction the chains go.

HyperbolicCutShape

About the HyperbolicCutShape class

The WaveBlocks Project

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Inheritance diagram

Class documentation

class WaveBlocksND.HyperbolicCutShape \((D, K)\)

This class implements the hyperbolic cut basis shape which is a special type of sparse basis set. A basis shape is essentially all information and operations related to the set \(K\) of multi-indices \(k\). The hyperbolic cut shape in \(D\) dimensions and with sparsity \(K\) is defined as the set

\[
K(D, K) := \{(k_0, \ldots, k_{D-1})|k_d \geq 0 \forall d \in [0, \ldots, D-1] \land \prod_{d=0}^{D-1} (1 + k_d) \leq K\}
\]

contains \((k)\)
Checks if a given multi-index \(k\) is part of the basis set \(K\).

Parameters \(k\) (tuple) – The multi-index \(k\) we want to test.

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The `extend()` method extends the basis shape such that (at least) all neighbours of all boundary nodes are included in the extended basis shape.

The `get_basis_size(extended=False)` method returns the size $|\mathcal{K}|$ of the basis. The size is the number of distinct multi-indices $k$ that belong to the basis $\mathcal{K}$.

The `get_description()` method returns a description of this basis shape object. A description is a `dict` containing all key-value pairs necessary to reconstruct the current basis shape. A description never contains any data.

The `get_dimension()` method returns the dimension $D$ of the basis shape $\mathcal{K}$. This is defined as the number of components each multi-index $k = (k_0, \ldots, k_{D-1})$ has.

The `get_limits()` method returns the upper limit $K$ which is the same for all directions $d$.

The `get_neighbours(k, selection=None, direction=None)` method returns a list of all multi-indices that are neighbours of a given multi-index $k$. A direct neighbour is defined as $(k_0, \ldots, k_d \pm 1, \ldots, k_{D-1}) \forall d \in [0 \ldots D-1]$.

**Parameters**

- `k` *(tuple)* – The multi-index of which we want to get the neighbours.
- `selection` *(string with fixed values forward, backward or all. The values all is equivalent to the value None (default).)*
- `direction` *(integer.)* – If iterating in chainmode this specifies the direction the chains go.

**Returns** A list containing the pairs $(d, k')$.

The `get_node_iterator(mode='lex', direction=None)` method returns an iterator to iterate over all basis elements $k \in \mathcal{K}$.

**Parameters**

- `mode` *(string)* – The mode by which we iterate over the indices. Default is lex for lexicographical order. Supported is also chain, for the chain-like mode, details see the manual.
- `direction` *(integer.)* – If iterating in chainmode this specifies the direction the chains go.

#### LimitedHyperbolicCutShape

**About the LimitedHyperbolicCutShape class**

The WaveBlocks Project

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Class documentation

class WaveBlocksND.LimitedHyperbolicCutShape \((D, K, \text{limits})\)
This class implements the hyperbolic cut basis shape which is a special type of sparse basis set. A basis shape is essentially all information and operations related to the set \(\mathcal{K}\) of multi-indices \(k\). The hyperbolic cut shape in \(D\) dimensions with sparsity \(S\) and limits \(K = (K_0, \ldots, K_{D-1})\) is defined as the set

\[
\mathcal{K}(D,S,K) := \{(k_0, \ldots, k_{D-1})|0 \leq k_d < K_d \forall d \in [0, \ldots, D-1] \land \prod_{d=0}^{D-1} (1 + k_d) \leq S\}
\]

contains \((k)\)
Checks if a given multi-index \(k\) is part of the basis set \(\mathcal{K}\).

Parameters k \((\text{tuple})\) – The multi-index \(k\) we want to test.

extend \((\text{tight=True})\)
Extend the basis shape such that (at least) all neighbours of all boundary nodes are included in the extended basis shape.

Parameters tight – Whether to cut off the long tails.

Param Boolean, default is False

generate_extension \((\text{tight=True})\)
Extend the basis shape such that (at least) all neighbours of all boundary nodes are included in the extended basis shape.

Parameters tight – Whether to cut off the long tails.

Param Boolean, default is False

get_basis_size \((\text{extended=False})\)
Returns the size \(|\mathcal{K}|\) of the basis. The size is the number of distinct multi-indices \(k\) that belong to the basis \(\mathcal{K}\).

generate_description()
Return a description of this basis shape object. A description is a \texttt{dict} containing all key-value pairs necessary to reconstruct the current basis shape. A description never contains any data.

generate_dimension()
Returns the dimension \(D\) of the basis shape \(\mathcal{K}\). This is defined as the number of components each multi-index \(k = (k_0, \ldots, k_{D-1})\) has.

generate_limits()
Returns the upper limit \(K_d\) for all directions \(d\).

Returns A tuple of the maximum of the multi-index in each direction.

generate_neighbours \((\text{k, selection=None, direction=None})\)
Returns a list of all multi-indices that are neighbours of a given multi-index \(k\). A direct neighbour is defined as \((k_0, \ldots, k_d \pm 1, \ldots, k_{D-1})\forall d \in [0 \ldots D - 1] \).

Parameters

\* k \((\text{tuple})\) – The multi-index of which we want to get the neighbours.
• **selection** (string with fixed values `forward`, `backward` or `all`. The values `all` is equivalent to the value `None` (default).) –

• **direction** (*int*) – The direction $0 \leq d < D$ in which we want to find the neighbours $k \pm e_d$.

Returns A list containing the pairs $(d, k')$.

**get_node_iterator** (mode='lex', direction=None)

Returns an iterator to iterate over all basis elements $k \in K$.

Parameters

• **mode** (*string*) – The mode by which we iterate over the indices. Default is `lex` for lexicographical order. Supported is also `chain`, for the chain-like mode, details see the manual.

• **direction** (*integer*) – If iterating in `chainmode` this specifies the direction the chains go.

### Wavepacket

**About the Wavepacket class**

The WaveBlocks Project

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### Inheritance diagram

```
Wavepacket.Wavepacket
```

### Class documentation

**class** `WaveBlocksND.Wavepacket (parameters)`

This class is an abstract interface to wavepackets in general.

**clone**()

Clone the wavepacket. Return a new copy of the wavepacket and make sure that all references between the two wavepackets get broken.

Raises **NotImplementedError** Abstract interface.

**gen_id**()

Generate an (unique) ID per wavepacket instance.

**get_dimension**()

Returns The space dimension $D$ of the wavepacket $\Psi$.

**get_id**()

Return the packet ID of this wavepacket instance. The ID may be used for storing packets in associative lists.
Returns The ID of the current instance.

get_number_components ()
Returns The number $N$ of components the wavepacket $\Psi$ has.

set_id (anid)
Manually set a new ID for the current wavepacket instance.

Parameters anid (int) – The new ID.

HagedornWavepacketBase

About the HagedornWavepacketBase class

The WaveBlocks Project
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Inheritance diagram

```
Wavepacket.Wavepacket ─────── HagedornWavepacketBase.HagedornWavepacketBase
```

Class documentation

class WaveBlocksND.HagedornWavepacketBase (parameters)
This class implements the abstract Wavepacket interface and contains code common to all types of Hagedorn wavepackets.

clone ()
Clone the wavepacket. Return a new copy of the wavepacket and make sure that all references between the two wavepackets get broken.

Raises NotImplementedError Abstract interface.

gen_id ()
Generate an (unique) ID per wavepacket instance.

get_basis_shape (component=None)
Retrieve the basis shapes $K_i$ for each component $i$.

Parameters component (int) – The component $i$ whose basis shape we request. (Default is None which means to return the basis shapes for all components.

Returns The basis shape for an individual component or a list with all shapes.

get_coefficient (component, index)
Retrieve a single coefficient $c_{k}^{i}$ of the specified component $\Phi_i$ of $\Psi$.

Parameters

• component – The index $i$ of the component $\Phi_i$ we want to update.
• **index** (A tuple of \( D \) integers.) – The multi-index \( k \) of the coefficient \( c^i_k \) we want to update.

  **Returns** A single complex number.

**get_coefficient_vector** ()

Retrieve the coefficients for all components \( \Phi_i \) simultaneously.

  **Returns** The coefficients \( c^i \) of all components \( \Phi_i \) stacked into a single long column vector.

**get_coefficients** (\( component=None \))

Returns the coefficients \( c^i \) for some component \( \Phi_i \) of \( \Psi \) or all the coefficients \( c \) of all components.

  **Note:** this method copies the data arrays.

  **Parameters**

  • **component** (int (Default is \( \text{None} \) meaning all)) – The index \( i \) of the component we want to retrieve.

  **Returns** A single ndarray with the coefficients of the given component or a list containing the ndarrays for each component. Each ndarray is two-dimensional with a shape of \((|K_i|, 1)\).

**get_dimension** ()

  **Returns** The space dimension \( D \) of the wavepacket \( \Psi \).

**get_eps** ()

Retrieve the semi-classical scaling parameter \( \varepsilon \) of the wavepacket.

  **Returns** The value of \( \varepsilon \).

**get_id** ()

Return the packet ID of this wavepacket instance. The ID may be used for storing packets in associative lists.

  **Returns** The ID of the current instance.

**get_number_components** ()

  **Returns** The number \( N \) of components the wavepacket \( \Psi \) has.

**get_quadrature** ()

Return the \( \text{Quadrature} \) subclass instance used computing inner products and evaluating brackets.

  **Returns** The current \( \text{Quadrature} \) subclass instance.

**norm** (\( component=None, summed=False \))

Calculate the \( L^2 \) norm \( \langle \Psi|\Psi \rangle \) of the wavepacket \( \Psi \).

  **Parameters**

  • **component** (int or \( \text{None} \).) – The index \( i \) of the component \( \Phi_i \) whose norm is calculated. The default value is \( \text{None} \) which means to compute the norms of all \( N \) components.

  • **summed** (Boolean, default is \( \text{False} \).) – Whether to sum up the norms \( \langle \Phi_i|\Phi_i \rangle \) of the individual components \( \Phi_i \).

  **Returns** The norm of \( \Psi \) or the norm of \( \Phi_i \) or a list with the \( N \) norms of all components. Depending on the values of \( \text{component} \) and \( \text{summed} \).

**set_basis_shape** (\( basis\_shape, component=None \))

Set the basis shape \( K \) of a given component or for all components.

  **Parameters**

  • **basis\_shape** (A subclass of \( \text{BasisShape} \).) – The basis shape for an individual component or a list with all \( N \) shapes.
• **component** *(int)* – The component \(i\) whose basis shape we want to set. (Default is `None` which means to set the basis shapes for all components.

**set_coefficient** *(component, index, value)*
Set a single coefficient \(c_k\) of the specified component \(\Phi_i\) of \(\Psi\).

**Parameters**

- **component** – The index \(i\) of the component \(\Phi_i\) we want to update.
- **index** *(A tuple of \(D\) integers.)* – The multi-index \(k\) of the coefficient \(c_k\) we want to update.
- **value** – The new value of the coefficient \(c_k\).

**Raises ValueError** For invalid indices \(i\) or \(k\).

**set_coefficient_vector** *(vector)*
Set the coefficients for all components \(\Phi_i\) simultaneously.

**Parameters**

- **vector** *(A two-dimensional `ndarray` of appropriate shape.)* – The coefficients of all components as a single long column vector.

**set_coefficients** *(values, component=None)*
Update all the coefficients \(c\) of \(\Psi\) or update the coefficients \(c_i\) of the components \(\Phi_i\) only.

**Parameters**

- **values** *(An `ndarray` of suitable shape or a list of `ndarrays`.* – The new values of the coefficients \(c_i\) of \(\Phi_i\).
- **component** *(int (Default is `None` meaning all))* – The index \(i\) of the component we want to update with new coefficients.

**Raises ValueError** For invalid component indices \(i\).

**set_id** *(anid)*
Manually set a new ID for the current wavepacket instance.

**Parameters**

- **anid** *(int)* – The new ID.

**set_quadrature** *(quadrature)*
Set the `Quadrature` subclass instance used for computing inner products and evaluating brakets.

**Parameters**

- **quadrature** – The new `Quadrature` subclass instance.

### HagedornWavepacket

#### About the HagedornWavepacket Class

The WaveBlocks Project

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Class documentation

class WaveBlocksND.HagedornWavepacket (dimension, ncomponents, eps)

This class represents homogeneous vector valued Hagedorn wavepackets $\Psi$ with $N$ components in $D$ space dimensions.

Parameters

- **dimension** – The space dimension $D$ the packet has.
- **ncomponents** – The number $N$ of components the packet has.
- **eps** – The semi-classical scaling parameter $\varepsilon$ of the basis functions.

Returns An instance of HagedornWavepacket.

static __new__ (S, ...) → a new object with type S, a subtype of T

__delattr__

x.__delattr__('name') <=> del x.name

__format__ ()

default object formatter

__getattribute__

x.__getattribute__('name') <=> x.name

__hash__

x.__hash__() <=> hash(x)

__init__ (dimension, ncomponents, eps)

Initialize a new homogeneous Hagedorn wavepacket.

    Parameters

    - **dimension** – The space dimension $D$ the packet has.
    - **ncomponents** – The number $N$ of components the packet has.
    - **eps** – The semi-classical scaling parameter $\varepsilon$ of the basis functions.

    Returns An instance of HagedornWavepacket.

__reduce__ ()

helper for pickle

__reduce_ex__ ()

helper for pickle

__repr__

x.__repr__() <=> repr(x)

__setattr__

x.__setattr__('name', value) <=> x.name = value

__sizeof__ () → int

size of object in memory, in bytes

__str__ ()

    Returns A string describing the Hagedorn wavepacket $\Psi$. 
static __subclasshook__()  
Abstract classes can override this to customize issubclass(). 
This is invoked early on by abc.ABCMeta.__subclasscheck__(). It should return True, False or Not Implemented. If it returns Not Implemented, the normal algorithm is used. Otherwise, it overrides the normal algorithm (and the outcome is cached).

__weakref__  
list of weak references to the object (if defined)

evaluate_at (grid, component=None, prefactor=False)  
Evaluate the Hagedorn wavepacket $\Psi$ at the given nodes $\gamma$. 

Parameters  
- grid (A class having a get_nodes(...)() method) – The grid $\Gamma'$ containing the nodes $\gamma$.  
- component – The index $i$ of a single component $\Phi_i$ to evaluate. (Defaults to None for evaluating all components.)  
- prefactor (bool, default is False.) – Whether to include a factor of $\frac{1}{\sqrt{\det(Q)}}$.

Returns  A list of arrays or a single array containing the values of the $\Phi_i$ at the nodes $\gamma$.

evaluate_basis_at (grid, component, prefactor=False)  
Evaluate the basis functions $\phi_k$ recursively at the given nodes $\gamma$. 

Parameters  
- grid (A class having a get_nodes(...)() method) – The grid $\Gamma'$ containing the nodes $\gamma$.  
- component – The index $i$ of a single component $\Phi_i$ to evaluate. We need this to choose the correct basis shape.  
- prefactor (bool, default is False.) – Whether to include a factor of $\frac{1}{\sqrt{\det(Q)}}$.

Returns  A two-dimensional ndarray $H$ of shape $(|K_i|, |\Gamma|)$ where the entry $H[\mu(k), i]$ is the value of $\phi_k(\gamma_i)$.

gen_id()  
Generate an (unique) ID per wavepacket instance.

get_basis_shape (component=None)  
Retrieve the basis shapes $K_i$ for each component $i$. 

Parameters component (int) – The component $i$ whose basis shape we request. (Default is None which means to return the basis shapes for all components.

Returns  The basis shape for an individual component or a list with all shapes.

get_coefficient (component, index)  
Retrieve a single coefficient $c^i_k$ of the specified component $\Phi_i$ of $\Psi$. 

Parameters  
- component – The index $i$ of the component $\Phi_i$ we want to update.  
- index (A tuple of $D$ integers.) – The multi-index $k$ of the coefficient $c^i_k$ we want to update.

Returns  A single complex number.

get_coefficient_vector ()  
Retrieve the coefficients for all components $\Phi_i$ simultaneously.
Returns The coefficients $c^i$ of all components $\Phi_i$ stacked into a single long column vector.

get_coefficients (component=None)
Returns the coefficients $c^i$ for some component $\Phi_i$ of $\Psi$ or all the coefficients $c$ of all components.

Note: this method copies the data arrays.

Parameters component (int (Default is None meaning all)) – The index $i$ of the component we want to retrieve.

Returns A single ndarray with the coefficients of the given component or a list containing the ndarrays for each component. Each ndarray is two-dimensional with a shape of $(|K_i|, 1)$.

get_description ()
Return a description of this wavepacket object. A description is a dict containing all key-value pairs necessary to reconstruct the current instance. A description never contains any data.

get_dimension ()
Returns The space dimension $D$ of the wavepacket $\Psi$.

get_eps ()
Retrieve the semi-classical scaling parameter $\varepsilon$ of the wavepacket.

Returns The value of $\varepsilon$.

get_id ()
Return the packet ID of this wavepacket instance. The ID may be used for storing packets in associative lists.

Returns The ID of the current instance.

get_number_components ()
Returns The number $N$ of components the wavepacket $\Psi$ has.

get_parameters (component=None, aslist=False)
Get the Hagedorn parameter set $\Pi$ of the wavepacket $\Psi$.

Parameters

• component – Dummy parameter for API compatibility with the inhomogeneous packets.

• aslist – Return a list of $N$ parameter tuples. This is for API compatibility with inhomogeneous packets.

Returns The Hagedorn parameter set $\Pi = (q, p, Q, P, S)$ in this order.

get_quadrature ()
Return the Quadrature subclass instance used computing inner products and evaluating brakets.

Returns The current Quadrature subclass instance.

norm (component=None, summed=False)
Calculate the $L^2$ norm $\langle \Psi | \Psi \rangle$ of the wavepacket $\Psi$.

Parameters

• component (int or None.) – The index $i$ of the component $\Phi_i$ whose norm is calculated. The default value is None which means to compute the norms of all $N$ components.

• summed (Boolean, default is False.) – Whether to sum up the norms $\langle \Phi_i | \Phi_i \rangle$ of the individual components $\Phi_i$.

Returns The norm of $\Psi$ or the norm of $\Phi_i$ or a list with the $N$ norms of all components. Depending on the values of component and summed.
**set_basis_shape** *(basis_shape, component=None)*

Set the basis shape \( K \) of a given component or for all components.

**Parameters**

- `basis_shape` *(A subclass of BasisShape.)* – The basis shape for an individual component or a list with all \( N \) shapes.
- `component` *(int)* – The component \( i \) whose basis shape we want to set. (Default is `None` which means to set the basis shapes for all components.)

**set_coefficient** *(component, index, value)*

Set a single coefficient \( c^i_k \) of the specified component \( \Phi_i \) of \( :math:Psi' \).

**Parameters**

- `component` – The index \( i \) of the component \( \Phi_i \) we want to update.
- `index` *(A tuple of \( D \) integers.)* – The multi-index \( k \) of the coefficient \( c^i_k \) we want to update.
- `value` – The new value of the coefficient \( c^i_k \).

**Raises** ValueError For invalid indices \( i \) or \( k \).

**set_coefficient_vector** *(vector)*

Set the coefficients for all components \( \Phi_i \) simultaneously.

*Note: This function does NOT copy the input data! This is for efficiency as this routine is used in the innermost loops.*

**Parameters**

- `vector` *(A two-dimensional ndarray of appropriate shape.)* – The coefficients of all components as a single long column vector.

**set_coefficients** *(values, component=None)*

Update all the coefficients \( c \) of \( \Psi \) or update the coefficients \( c^i \) of the components \( \Phi_i \) only.

*Note: this method copies the data arrays.*

**Parameters**

- `values` *(An ndarray of suitable shape or a list of ndarrays.)* – The new values of the coefficients \( c^i \) of \( \Phi_i \).
- `component` *(int (Default is `None` meaning all))* – The index \( i \) of the component we want to update with new coefficients.

**Raises** ValueError For invalid component indices \( i \).

**set_id** *(anid)*

Manually set a new ID for the current wavepacket instance.

**Parameters**

- `anid` *(int)* – The new ID.

**set_parameters** *(Pi, component=None)*

Set the Hagedorn parameters \( \Pi \) of the wavepacket \( \Psi \).

**Parameters**

- `Pi` – The Hagedorn parameter set \( \Pi = (q, p, Q, P, S) \) in this order.
- `component` – Dummy parameter for API compatibility with the inhomogeneous packets.

**set_quadrature** *(quadrature)*

Set the Quadrature subclass instance used for computing inner products and evaluating brakets.

**Parameters**

- `quadrature` – The new Quadrature subclass instance.
HagedornWavepacketInhomogeneous

About the HagedornWavepacketInhomogeneous class

The WaveBlocks Project

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Inheritance diagram

Class documentation

class WaveBlocksND.HagedornWavepacketInhomogeneous (dimension, ncomponents, eps)
This class represents inhomogeneous vector valued Hagedorn wavepackets $\Psi$ with $N$ components in $D$ space dimensions.

__delattr__
   x.__delattr__('name') ==> del x.name

__format__
   default object formatter

__getattribute__
   x.__getattribute__('name') ==> x.name

__hash__
   x.__hash__() ==> hash(x)

__init__ (dimension, ncomponents, eps)
   Initialize a new inhomogeneous Hagedorn wavepacket.

Parameters

- **dimension** – The space dimension $D$ the packet has.
- **ncomponents** – The number $N$ of components the packet has.
- **eps** – The semi-classical scaling parameter $\varepsilon$ of the basis functions.

Returns An instance of HagedornWavepacketInhomogeneous.

static __new__ (S, ...) → a new object with type S, a subtype of T

__reduce__ ()
   helper for pickle

__reduce_ex__ ()
   helper for pickle

__repr__
   x.__repr__() ==> repr(x)
__setattr__
x.__setattr__('name', value) <==> x.name = value

__sizeof__
size of object in memory, in bytes

__str__
Returns A string describing the Hagedorn wavepacket $\Psi$.

static __subclasshook__()
Abstract classes can override this to customize issubclass().
This is invoked early on by abc.ABCMeta.__subclasscheck__(). It should return True, False or Not Implemented. If it returns Not Implemented, the normal algorithm is used. Otherwise, it overrides the normal algorithm (and the outcome is cached).

__weakref__
list of weak references to the object (if defined)

evaluate_at (grid, component=None, prefactor=False)
Evaluate the Hagedorn wavepacket $\Psi$ at the given nodes $\gamma$.

Parameters

- **grid** (A class having a get_nodes(...) () method.) – The grid $\Gamma'$ containing the nodes $\gamma$.

- **component** – The index $i$ of a single component $\Phi_i$ to evaluate. (Defaults to None for evaluating all components.)

- **prefactor** (bool, default is False.) – Whether to include a factor of $\frac{1}{\sqrt{\det(Q)}}$.

Returns A list of arrays or a single array containing the values of the $\Phi_i$ at the nodes $\gamma$.

evaluate_basis_at (grid, component, prefactor=False)
Evaluate the basis functions $\phi_k$ recursively at the given nodes $\gamma$.

Parameters

- **grid** (A class having a get_nodes(...) () method.) – The grid $\Gamma'$ containing the nodes $\gamma$.

- **component** – The index $i$ of a single component $\Phi_i$ to evaluate.

- **prefactor** (bool, default is False.) – Whether to include a factor of $\frac{1}{\sqrt{\det(Q)}}$.

Returns A two-dimensional ndarray $H$ of shape $(|K_i|, |\Gamma|)$ where the entry $H[\mu(k), i]$ is the value of $\phi_k(\gamma_i)$.

gen_id()
Generate an (unique) ID per wavepacket instance.

get_basis_shape (component=None)
Retrieve the basis shapes $K_i$ for each component $i$.

Parameters **component** (int) – The component $i$ whose basis shape we request. (Default is None which means to return the basis shapes for all components.)

Returns The basis shape for an individual component or a list with all shapes.

get_coefficient (component, index)
Retrieve a single coefficient $c_k^i$ of the specified component $\Phi_i$ of $\Psi$.

Parameters
• **component** – The index \( i \) of the component \( \Phi_i \) we want to update.

• **index** (A tuple of \( D \) integers.) – The multi-index \( k \) of the coefficient \( c_k^i \) we want to update.

**Returns** A single complex number.

**get_coefficient_vector()**
Retrieve the coefficients for all components \( \Phi_i \) simultaneously.

**Returns** The coefficients \( c_i^j \) of all components \( \Phi_i \) stacked into a single long column vector.

**get_coefficients(component=None)**
Returns the coefficients \( c_i^j \) for some component \( \Phi_i \) of \( \Psi \) or all the coefficients \( c_i \) of all components.

Note: this method copies the data arrays.

**Parameters**
- **component** (int (Default is `None` meaning all)) – The index \( i \) of the component we want to retrieve.

**Returns** A single ndarray with the coefficients of the given component or a list containing the ndarrays for each component. Each ndarray is two-dimensional with a shape of \((|K_i|, 1)\).

**get_description()**
Return a description of this wavepacket object. A description is a `dict` containing all key-value pairs necessary to reconstruct the current instance. A description never contains any data.

**get_dimension()**

**Returns** The space dimension \( D \) of the wavepacket \( \Psi \).

**get_eps()**
Retrieve the semi-classical scaling parameter \( \varepsilon \) of the wavepacket.

**Returns** The value of \( \varepsilon \).

**get_id()**
Return the packet ID of this wavepacket instance. The ID may be used for storing packets in associative lists.

**Returns** The ID of the current instance.

**get_number_components()**

**Returns** The number \( N \) of components the wavepacket \( \Psi \) has.

**get_parameters(component=None, aslist=False)**
Get the Hagedorn parameter set \( \Pi_i \) of each component \( \Phi_i \) of the wavepacket \( \Psi \).

**Parameters**
- **component** – The index \( i \) of the component \( \Phi_i \) whose parameters \( \Pi_i \) we want to get.
- **aslist** – Dummy parameter for API compatibility with the homogeneous packets.

**Returns** A list with all parameter sets \( \Pi_i \) or a single parameter set. The parameters \( \Pi_i = (q_i, p_i, Q_i, P_i, S_i) \) are always in this order.

**get_quadrature()**
Return the `Quadrature` subclass instance used computing inner products and evaluating brakets.

**Returns** The current `Quadrature` subclass instance.

**norm(component=None, summed=False)**
Calculate the \( L^2 \) norm \( \langle \Psi | \Psi \rangle \) of the wavepacket \( \Psi \).

**Parameters**
• **component** (int or None) – The index \(i\) of the component \(\Phi_i\) whose norm is calculated. The default value is None which means to compute the norms of all \(N\) components.

• **summed** (Boolean, default is False) – Whether to sum up the norms \(\langle \Phi_i | \Phi_i \rangle\) of the individual components \(\Phi_i\).

**Returns** The norm of \(\Psi\) or the norm of \(\Phi_i\) or a list with the \(N\) norms of all components. Depending on the values of component and summed.

**set_basis_shape**(basis_shape, component=None)

Set the basis shape \(K\) of a given component or for all components.

**Parameters**

• **basis_shape** (A subclass of BasisShape.) – The basis shape for an individual component or a list with all \(N\) shapes.

• **component** (int) – The component \(i\) whose basis shape we want to set. (Default is None which means to set the basis shapes for all components.

**set_coefficient**(component, index, value)

Set a single coefficient \(c_{ik}\) of the specified component \(\Phi_i\) of \(\Psi\)

**Parameters**

• **component** – The index \(i\) of the component \(\Phi_i\) we want to update.

• **index** (A tuple of \(D\) integers.) – The multi-index \(k\) of the coefficient \(c_{ik}\) we want to update.

• **value** – The new value of the coefficient \(c_{ik}\).

**Raises ValueError** For invalid indices \(i\) or \(k\).

**set_coefficient_vector**(vector)

Set the coefficients for all components \(\Phi_i\) simultaneously.

Note: This function does NOT copy the input data! This is for efficiency as this routine is used in the innermost loops.

**Parameters** vector (A two-dimensional ndarray of appropriate shape.) – The coefficients of all components as a single long column vector.

**set_coefficients**(values, component=None)

Update all the coefficients \(c_i\) of \(\Psi\) or update the coefficients \(c_{i}\) of the components \(\Phi_i\) only.

Note: this method copies the data arrays.

**Parameters**

• **values** (An ndarray of suitable shape or a list of ndarrays.) – The new values of the coefficients \(c_{ik}\) of \(\Phi_i\).

• **component** (int (Default is None meaning all)) – The index \(i\) of the component we want to update with new coefficients.

**Raises ValueError** For invalid component indices \(i\).

**set_id**(anid)

Manually set a new ID for the current wavepacket instance.

**Parameters** anid (int) – The new ID.

**set_parameters**(Pi, component=None)

Set the Hagedorn parameter set \(\Pi_i\) of each component \(\text{math}’\Phi_i’\) of the wavepacket \(\Psi\).

**Parameters**
• **Pi** *(A single tuple or a list of tuples)* – The parameter sets \( \Pi_i = (q_i, p_i, Q_i, P_i, S_i) \) with its values in this order.

• **component** – The index \( i \) of the component \( \Phi_i \) whose parameters \( \Pi_i \) we want to update.

```python
set_quadrature(quadrature)
```

Set the **Quadrature** subclass instance used for computing inner products and evaluating brackets.

**Parameters**
- `quadrature` – The new **Quadrature** subclass instance.

**Quadrature**

**About the Quadrature class**

The WaveBlocks Project

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**Class documentation**

class WaveBlocksND.Quadrature

This class is an abstract interface to quadratures in general.

```python
build_matrix()
```

Calculate the matrix elements of \( \langle \Psi | f | \Psi \rangle \) for a general function \( f(x) \) with \( x \in \mathbb{R}^D \). Note that the arguments may vary through subclasses!

**Raises** `NotImplementedError` Abstract interface.

```python
get_description()
```

Return a description of this quadrature object. A description is a `dict` containing all key-value pairs necessary to reconstruct the current instance. A description never contains any data.

```python
get_qr()
```

Return the **QuadratureRule** subclass instance used for quadrature.

**Returns** The current instance of the quadrature rule.

```python
quadrature()
```

Performs the quadrature of \( \langle \Psi | f | \Psi \rangle \) for a general function \( f(x) \) with \( x \in \mathbb{R}^D \). Note that the arguments may vary through subclasses!

**Raises** `NotImplementedError` Abstract interface.
set_qr(QR)
Set the QuadratureRule subclass instance used for quadrature.

Parameters
QR – The new QuadratureRule instance.

transform_nodes()
Transform the quadrature nodes such that they fit the given wavepacket. Note that the arguments may vary
through subclasses!

Raises NotImplementedError Abstract interface.

HomogeneousQuadrature

About the HomogeneousQuadrature class

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Inheritance diagram

Class documentation

class WaveBlocksND.HomogeneousQuadrature (QR=None)

build_matrix (packet, operator=None)
Calculate the matrix elements of \( \langle \Psi | f | \Psi \rangle \) for a general function \( f(x) \) with \( x \in \mathbb{R}^D \). The matrix is
computed without including the coefficients \( c_k \).

Parameters
• packet – The wavepacket \( \Psi \).
• operator – A matrix-valued function \( f(q,x) : \mathbb{R} \times \mathbb{R}^D \rightarrow \mathbb{R}^{N \times N} \).

Returns
A square matrix of size \( \sum_i^N |\mathcal{K}_i| \times \sum_j^N |\mathcal{K}_j| \).

get_description()
Return a description of this quadrature object. A description is a dict containing all key-value pairs
necessary to reconstruct the current instance. A description never contains any data.

get_qr()
Return the QuadratureRule subclass instance used for quadrature.

Returns
The current instance of the quadrature rule.

quadrature (packet, operator=None, summed=False, component=None, diag_component=None)
Performs the quadrature of \( \langle \Psi | f | \Psi \rangle \) for a general function \( f(x) \) with \( x \in \mathbb{R}^D \).

Parameters
• **packet** – The wavepacket $\Psi$.

• **operator** – A matrix-valued function $f(x) : \mathbb{R}^D \rightarrow \mathbb{R}^{N \times N}$.

• **summed** (bool, default is False.) – Whether to sum up the individual integrals $\langle \Phi_i | f_{i,j} | \Phi_j \rangle$.

• **component** – Request only the i-th component of the result. Remember that $i \in [0, N^2 - 1]$.

• **diag_component** – Request only the i-th component from the diagonal entries, here $i \in [0, N - 1]$. Note that component takes precedence over diag_component if both are supplied. (Which is discouraged)

**Returns** The value of the braket $\langle \Psi | f | \Psi \rangle$. This is either a scalar value or a list of $N^2$ scalar elements depending on the value of summed.

**set_qr**(QR)

Set the QuadratureRule subclass instance used for quadrature.

**Parameters**

- QR – The new QuadratureRule instance.

**transform_nodes**(Pi, eps, QR=None)

Transform the quadrature nodes $\gamma$ such that they fit the given wavepacket $\Phi [\Pi]$.

**Parameters**

- Pi – The parameter set $\Pi$ of the wavepacket.
- eps – The value of $\epsilon$ of the wavepacket.
- QR – An optional quadrature rule $\Gamma = (\gamma, \omega)$ providing the nodes. If not given the internal quadrature rule will be used.

**Returns** A two-dimensional ndarray of shape ($D, |\Gamma|$) where $|\Gamma|$ denotes the total number of quadrature nodes.

**InhomogeneousQuadrature**

**About the InhomogeneousQuadrature class**

The WaveBlocks Project

@author: R. Bourquin @copyright: Copyright (C) 2010, 2011, 2012 R. Bourquin @license: Modified BSD License

**Inheritance diagram**

![Inheritance diagram]

**Class documentation**

```python
class WaveBlocksND.InhomogeneousQuadrature(QR=None)
```
build_matrix (pacbra, packet=None, operator=None)

Calculate the matrix elements of \( \langle \Psi | f | \Psi' \rangle \) for a general function \( f(x) \) with \( x \in \mathbb{R}^D \). The matrix is computed without including the coefficients \( c_k^i \).

Parameters

- pacbra – The wavepacket \( \Psi \) from the bra with \( N \) components.
- packet – The wavepacket \( \Psi' \) from the ket with \( N' \) components.
- operator – A matrix-valued function \( f(q, x) : \mathbb{R} \times \mathbb{R}^D \rightarrow \mathbb{R}^{N \times N'} \).

Returns A matrix of size \( \sum_i N_i \times \sum_j N'_j \).

calculate (pacbra, packet=None, operator=None)

get_description ()

Return a description of this quadrature object. A description is a dict containing all key-value pairs necessary to reconstruct the current instance. A description never contains any data.

generate_quad_rule ()

Return the QuadratureRule subclass instance used for quadrature.

Returns The current instance of the quadrature rule.

mix_parameters (Pibra, Piket)

Mix the two parameter sets \( \Pi_i \) and \( \Pi_j \) from the bra and the ket wavepackets \( \Phi \[\Pi_i\] \) and \( \Phi' \[\Pi_j\] \).

Parameters

- Pibra – The parameter set \( \Pi_i \) from the bra part wavepacket.
- Piket – The parameter set \( \Pi_j \) from the ket part wavepacket.

Returns The mixed parameters \( q_0 \) and \( Q_{S} \). (See the theory for details.)

quadrature (pacbra, packet=None, operator=None, summed=False, component=None, diag_component=None)

Performs the quadrature of \( \langle \Psi | f | \Psi' \rangle \) for a general function \( f(x) \) with \( x \in \mathbb{R}^D \).

Parameters

- pacbra – The wavepacket \( \Psi \) from the bra with \( N \) components.
- packet – The wavepacket \( \Psi' \) from the ket with \( N' \) components.
- operator – A matrix-valued function \( f(q, x) : \mathbb{R} \times \mathbb{R}^D \rightarrow \mathbb{R}^{N \times N'} \).
- summed (bool, default is False.) – Whether to sum up the individual integrals \( \langle \Phi, f, i,j \rangle \Phi' \).
- component – Request only the i-th component of the result. Remember that \( i \in [0, N \cdot N' - 1] \).
- diag_component – Request only the i-th component from the diagonal entries, here \( i \in [0, N' - 1] \). Note that component takes precedence over diag_component if both are supplied. (Which is discouraged)

Returns The value of the braket \( \langle \Psi | f | \Psi' \rangle \). This is either a scalar value or a list of \( N \cdot N' \) scalar elements depending on the value of summed.

set_qr (QR)

Set the QuadratureRule subclass instance used for quadrature.

Parameters QR – The new QuadratureRule instance.

transform_nodes (Pibra, Piket, eps, QR=None)

Transform the quadrature nodes \( \gamma \) such that they fit the given wavepackets \( \Phi \[\Pi_i\] \) and \( \Phi' \[\Pi_j\] \) best.

1.1. WaveBlocks Classes
Parameters

- **Pibra** – The parameter set $\Pi_i$ from the bra part wavepacket.
- **Piket** – The parameter set $\Pi_j$ from the ket part wavepacket.
- **eps** – The value of $\varepsilon$ of the wavepacket.
- **QR** – An optional quadrature rule $\Gamma = (\gamma, \omega)$ providing the nodes. If not given the internal quadrature rule will be used.

**Returns** A two-dimensional ndarray of shape $(D, |\Gamma|)$ where $|\Gamma|$ denotes the total number of quadrature nodes.

### BasisTransformation

#### About the BasisTransformation class

The WaveBlocks Project

@author: R. Bourquin @copyright: Copyright (C) 2010, 2011, 2012 R. Bourquin @license: Modified BSD License

#### Inheritance diagram

```
BasisTransformation.BasisTransformation
```

#### Class documentation

class WaveBlocksND.BasisTransformation(potential)

This class is the interface definition for general basis transformation procedures. The transformation switches between the canonical basis of the potential $V(x)$ and its eigenbasis $\Lambda(x)$ consisting of the energy levels $\lambda_i(x)$ with $i \in [0, \ldots, N]$.

- **__delattr__**
  - `x.__delattr__('name') <=> del x.name`

- **__format__**
  - `default object formatter`

- **__getattribute__**
  - `x.__getattribute__('name') <=> x.name`

- **__hash__**
  - `x.__hash__() <=> hash(x)`

- **__init__**(potential)
  - Create a new *BasisTransformation* instance for a given potential matrix $V(x)$. 
Parameters potential (A MatrixPotential instance.) – The potential underlying the basis transformation.

\texttt{__new__} (S, ...) → a new object with type S, a subtype of T

\texttt{__reduce__}()

  helper for pickle

\texttt{__reduce_ex__}()

  helper for pickle

\texttt{__repr__}

  \texttt{x.__repr__()} \iff \texttt{repr(x)}

\texttt{__setattr__}

  \texttt{x.__setattr__('name', value)} \iff \texttt{x.name = value}

\texttt{__sizeof__}()

  \texttt{size of object in memory, in bytes}

\texttt{__str__}

  \texttt{x.__str__()} \iff \texttt{str(x)}

\texttt{__subclasshook__}()

  Abstract classes can override this to customize issubclass().

  This is invoked early on by abc.ABCMeta.__subclasscheck__. It should return True, False or NotImplemented. If it returns NotImplemented, the normal algorithm is used. Otherwise, it overrides the normal algorithm (and the outcome is cached).

\texttt{__weakref__}

  list of weak references to the object (if defined)

\texttt{transform_to_canonical} (transformable)

  Do nothing, implement an identity transformation.

\texttt{transform_to_eigen} (transformable)

  Do nothing, implement an identity transformation.

\textbf{BasisTransformationWF}

\textbf{About the BasisTransformationWF class}

The WaveBlocks Project

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Inheritance diagram

Class documentation

```python
class WaveBlocksND.BasisTransformationWF (potential, grid=None)
```
This class implements basis transformations of wavefunctions $\psi(x)$ between the canonical basis of and the basis $\Lambda(x)$ spanned by the eigenvectors $\nu_i(x)$ of the potential $V(x)$.

```python
__delattr__
   x.__delattr__('name') <==> del x.name

__format__()
   default object formatter

__getattribute__
   x.__getattribute__('name') <==> x.name

__hash__
   x.__hash__() <==> hash(x)

__init__ (potential, grid=None)
   Create a new BasisTransformation instance for a given potential matrix $V(x)$.

Parameters

- **potential** (A MatrixPotential instance.) – The potential underlying the basis transformation.
- **grid** (A Grid subclass instance.) – The grid.

```python
static __new__ (S, ...) → a new object with type S, a subtype of T
```

```python
__reduce__()
   helper for pickle

__reduce_ex__()
   helper for pickle

__repr__
   x.__repr__() <==> repr(x)

__setattr__
   x.__setattr__('name', value) <==> x.name = value

__sizeof__ () → int
   size of object in memory, in bytes

__str__
   x.__str__() <==> str(x)
```

```python
static __subclasshook__ ()
   Abstract classes can override this to customize issubclass().
```
This is invoked early on by abc.ABCMeta._subclasscheck__(). It should return True, False or NotImplemented. If it returns NotImplemented, the normal algorithm is used. Otherwise, it overrides the normal algorithm (and the outcome is cached).

__weakref__
list of weak references to the object (if defined)

set_grid(grid)
Set the grid $\Gamma$ containing the nodes $\gamma$ on which the wavefunction $\psi$ was evaluated. The $N$ eigenvectors $\nu_i$ will be evaluated on the same grid nodes.

Parameters grid (A Grid subclass instance.) – The grid

transform_to_canonical(wavefunction)
Transform the evaluated wavefunction $\psi(\Gamma)$ given in the eigenbasis to the canonical basis.

Parameters wavefunction (A WaveFunction instance.) – The wavefunction to transform.

Returns Another WaveFunction instance containing the transformed wavefunction $\psi'(\Gamma)$.

transform_to_eigen(wavefunction)
Transform the evaluated wavefunction $\psi'(\Gamma)$ given in the canonical basis to the eigenbasis.

Parameters wavefunction (A WaveFunction instance.) – The wavefunction to transform.

Returns Another WaveFunction instance containing the transformed wavefunction $\psi(\Gamma)$.

BasisTransformationHAWP

About the BasisTransformationHAWP class

The WaveBlocks Project

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Inheritance diagram

Class documentation

class WaveBlocksND.BasisTransformationHAWP (potential, builder=None)
This class implements basis transformations of Hagedorn wavepackets $\Psi(x)$ between the canonical basis of and the basis $\Lambda(x)$ spanned by the eigenvectors $\nu_i(x)$ of the potential $V(x)$.

__delattr__
x.__delattr__('name') <=> del x.name

__format__()
default object formatter
__getattribute__
   x.__getattribute__('name') <==> x.name

__hash__
   x.__hash__() <==> hash(x)

__init__ (potential, builder=None)
   Create a new BasisTransformationHAWP instance for a given potential matrix \( V(x) \).

   Parameters
   
   - **potential** (A MatrixPotential instance.) – The potential underlying the basis transformation.
   
   - **builder** (A Quadrature subclass instance.) – An object that can compute this matrix.

static __new__ (S, ...) → a new object with type S, a subtype of T

__reduce__ ()
   helper for pickle

__reduce_ex__ ()
   helper for pickle

__repr__
   x.__repr__() <==> repr(x)

__setattr__
   x.__setattr__('name', value) <==> x.name = value

__sizeof__ () → int
   size of object in memory, in bytes

__str__
   x.__str__() <==> str(x)

static __subclasscheck__ ()
   Abstract classes can override this to customize issubclass().

   This is invoked early on by abc.ABCMeta.__subclasscheck__(). It should return True, False or NotImple-
  mented. If it returns NotImplemented, the normal algorithm is used. Otherwise, it overrides the normal
   algorithm (and the outcome is cached).

__weakref__
   list of weak references to the object (if defined)

set_matrix_builder (builder)
   Set the matrix builder. It is responsible for computing the matrix elements \( \langle \phi_i | V_{ij} | \phi_j \rangle \). This matrix is
   used during the basis transformation.

   Parameters builder (A Quadrature subclass instance.) – An object that can compute this
   matrix.

transform_to_canonical (wavepacket)
   Transform the wavepacket \( \Psi \) given in the eigenbasis to the canonical basis.

   Note that this method acts destructively on the given Wavepacket instance. If this is not desired, clone
   the packet before handing it over to this method.

   Parameters wavepacket (A Wavepacket subclass instance.) – The Hagedorn wavepacket to
   transform.

   Returns Another Wavepacket instance containing the transformed wavepacket \( \Psi' \).
transform_to_eigen(wavepacket)
Transform the wavepacket $\Psi'$ given in the canonical basis to the eigenbasis.

Note that this method acts destructively on the given Wavepacket instance. If this is not desired, clone the packet before handing it over to this method.

Parameters wavepacket (A Wavepacket subclass instance.) – The Hagedorn wavepacket to transform.

Returns Another Wavepacket instance containing the transformed wavepacket $\Psi$.

1.1.4 Observables

Observables

About the Observables class

The WaveBlocks Project

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Inheritance diagram

Class documentation

class WaveBlocksND.Observables
This class is the interface definition for general observable computation procedures.

kinetic_energy(ket, T)
Compute the kinetic energy $E_{\text{kin}} := \langle \psi | T | \psi \rangle$.

Parameters

- ket – The object denoted by $\psi$.
- T – The kinetic energy operator $T$.

Raises NotImplementedError Abstract interface.

potential_energy(ket, potential)
Compute the potential energy $E_{\text{pot}} := \langle \psi | V | \psi \rangle$.

Parameters

- ket – The object denoted by $\psi$.
- potential – The potential $V(x)$.  

1.1. WaveBlocks Classes
Raises **NotImplementedError** Abstract interface.

**ObservablesHAWP**

**About the ObservablesHAWP class**

The WaveBlocks Project

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**Class documentation**

```python
class WaveBlocksND.ObservablesHAWP (quadrature=None)
    This class implements observable computation for Hagedorn wavepackets \( \Psi \).

    `apply_gradient`(wavepacket, component)
    Compute the effect of the gradient operator \(-i\varepsilon^2 \nabla_x\) on the basis functions \(\phi(x)\) of a component \(\Phi_i\) of the Hagedorn wavepacket \(\Psi\).

        Parameters
        • `wavepacket` (A HagedornWavepacketBase subclass instance.) – The wavepacket \(\Psi\) containing \(\Phi_i\).
        • `component` (int) – The index \(i\) of the component \(\Phi_i\).

        Returns
        Extended basis shape \(\hat{\mathcal{K}}\) and new coefficients \(c'\).

    `kinetic_energy`(wavepacket, component=None, summed=False)
    Compute the kinetic energy \(E_{\text{kin}} := \langle \Psi|T|\Psi \rangle\) of the different components \(\Phi_i\) of the wavepacket \(\Psi\).

        Parameters
        • `wavepacket` (A HagedornWavepacketBase subclass instance.) – The wavepacket \(\Psi\) of which we compute the kinetic energy.
        • `component` (Integer or None.) – The index \(i\) of the component \(\Phi_i\) whose kinetic energy we want to compute. If set to None the computation is performed for all \(N\) components.
        • `summed` (Boolean) – Whether to sum up the kinetic energies \(E_i\) of the individual components \(\Phi_i\). Default is False.

        Returns
        A list with the kinetic energies of the individual components or the overall kinetic energy of the wavepacket. (Depending on the optional arguments.)
```
potential_energy(wavepacket, potential, component=None, summed=False)

Compute the potential energy \( E_{\text{pot}} := \langle \Psi | V | \Psi \rangle \) of the different components \( \Phi_i \) of the wavepacket \( \Psi \).

**Parameters**
- **wavepacket** (A `HagedornWavepacketBase` subclass instance.) – The wavepacket \( \Psi \) of which we compute the potential energy.
- **potential** – The potential \( V(x) \).
- **component** (Integer or `None`. ) – The index \( i \) of the component \( \Phi_i \) whose potential energy we want to compute. If set to `None` the computation is performed for all \( N \) components.
- **summed** (Boolean) – Whether to sum up the potential energies \( E_i \) of the individual components \( \Phi_i \). Default is `False`.

**Returns** A list with the potential energies of the individual components or the overall potential energy of the wavepacket. (Depending on the optional arguments.)

set_quadrature(quadrature)

Set the quadrature.

**Parameters**
- **quadrature** (A `Quadrature` subclass instance.) – A quadrature for computing the integrals. Quadrature is only used for the computation of the potential energy \( \langle \Psi | V(x) | \Psi \rangle \) but not for the kinetic energy.

1.1.5 Time propagation

**KineticOperator**

**About the KineticOperator class**

The WaveBlocks Project

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Inheritance diagram

```
KineticOperator.KineticOperator
```

**Class documentation**

```python
class WaveBlocksND.KineticOperator(grid, eps=None)

This class represents the kinetic operator \( T \) in Fourier space.
```

```python
__delattr__
x.__delattr__('name') <=> del x.name
```

1.1. WaveBlocks Classes 53
__format__
default object formatter

__getattribute__
x.__getattribute__('name') <==> x.name

__hash__
x.__hash__() <==> hash(x)

__init__(grid, eps=None)
Compute the Fourier transformation of the position space representation of the kinetic operator $T$.

Parameters
- grid – The position space grid $\Gamma$ of which we compute its Fourier transform $\Omega$.
- eps – The semi-classical scaling parameter $\varepsilon$. (optional)

static __new__(S, ...) → a new object with type S, a subtype of T

__reduce__
helper for pickle

__reduce_ex__
helper for pickle

__repr__
x.__repr__() <==> repr(x)

__setattr__
x.__setattr__('name', value) <==> x.name = value

__sizeof__
→ int
size of object in memory, in bytes

__str__
x.__str__() <==> str(x)

static __subclasshook__
Abstract classes can override this to customize issubclass().
This is invoked early on by abc.ABCMeta.__subclasscheck__(). It should return True, False or NotImple-
mented. If it returns NotImplemented, the normal algorithm is used. Otherwise, it overrides the normal
algorithm (and the outcome is cached).

__weakref__
list of weak references to the object (if defined)

calculate_exponential (factor=1.0)
Calculate the exponential $\exp(\alpha \omega \cdot \omega)$ used in the Strang splitting.

Parameters factor – The prefactor $\alpha$. It defaults to 1 but is usually set to $-\frac{i}{2} \varepsilon^2 \tau$ by the caller.

calculate_operator (eps=None)
Calculate the kinetic operator $\hat{T} = \frac{\varepsilon^4}{\tau} \omega \cdot \omega$ in Fourier space.

Parameters eps – The semi-classical scaling parameter $\varepsilon$. It has to be given here or during the
initialization of the current instance.

evaluate_at (grid=None)
Evaluate the kinetic operator $\hat{T} = \frac{\varepsilon^4}{\tau} \omega \cdot \omega$ in Fourier space. This returns an numpy ndarray by using a
specific $\omega$.

Parameters grid – Unused dummy parameter.
**evaluate_exponential_at** *(grid=None)*
Evaluate the exponential \( \exp(\alpha T) \) in Fourier space. This returns a numpy `ndarray` by using a specific \( \omega \). The factor \( \alpha \) can be set by the corresponding method.

**Parameters**
- `grid` – Unused dummy parameter.

**get_fourier_grid_axes**
Return the grid axes of the Fourier space grid \( \Omega \).

---

**Propagator**

**About the Propagator class**

The WaveBlocks Project

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**Inheritance diagram**

```
            Propagator
```

**Class documentation**

```python
class WaveBlocksND.Propagator

Propagators can numerically simulate the time evolution of quantum states \( \psi(x,t) \) as described by the time-dependent Schroedinger equation

\[
i\varepsilon^2 \frac{\partial}{\partial t} \psi(x,t) = H\psi(x,t)
\]

where the semi-classical scaling parameter \( \varepsilon > 0 \) is already included. The Hamiltonian operator \( H \) is defined as

\[
H = T + V(x) = -\frac{\varepsilon^4}{2} \Delta + V(x)
\]

**get_number_components**

- **Returns** The number \( N \) components of \( \psi(x,t) \).
- **Raises** `NotImplementedError` This is an abstract base class.

**get_potential**

- Returns the potential \( V(x) \) used for time propagation.
```

---

1.1. WaveBlocks Classes 55
Returns A MatrixPotential subclass instance.

propagate()
Given the wavefunction ψ at time t, calculate the new :math:ψ' at time t + τ. We do exactly one timestep of size τ here.

Raises NotImplementedError This is an abstract base class.

FourierPropagator

About the FourierPropagator class

This class can numerically propagate given initial values Ψ(x₀, t₀) on a potential hypersurface V(x). The propagation is done with a Strang splitting of the time propagation operator exp(−iε²τH).

get_number_components()
Get the number N of components of Ψ.

Returns The number N.

get_operators()
Get the kinetic and potential operators T(Ω) and V(Γ).

Returns A tuple (T, V) containing two ndarrays.

get_potential()
Returns the potential V(x) used for time propagation.

Returns A MatrixPotential subclass instance.

get_wavefunction()
Get the wavefunction that stores the current data Ψ(Γ).

Returns The WaveFunction instance.

propagate()
Given the wavefunction values Ψ(Γ) at time t, calculate new values Ψ'(Γ) at time t + τ. We perform exactly one single timestep of size τ within this function.
HagedornPropagator

About the HagedornPropagator class

The WaveBlocks Project

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Inheritance diagram

![Inheritance diagram](Propagator.Propagator -> HagedornPropagator.HagedornPropagator)

Class documentation

class WaveBlocksND.HagedornPropagator (parameters, potential, packets=[])  
This class can numerically propagate given initial values $\Psi$ in a potential $V(x)$. The propagation is done for a given set of homogeneous Hagedorn wavepackets neglecting interaction.

add_wavepacket (packet)  
Add a new wavepacket $\Psi$ to the list of propagated wavepackets.

Parameters packet (A tuple ($\Psi$, $\chi$) with $\Psi$ a HagedornWavepacket instance and $\chi$ an integer.) – The new wavepacket $\Psi$ and its leading component $\chi \in [0, N - 1]$.

get_number_components ()  
Returns The number $N$ of components $\Phi_i$ of $\Psi$.

get_potential ()  
Returns the potential $V(x)$ used for time propagation.

Parameters packet (Integer or None) – The index $i$ (in this list) of a single packet $\Psi_i$ that is to be returned. If set to None (default) return the full list with all packets.

Returns A MatrixPotential subclass instance.

get_wavepackets (packet=None)  
Return the wavepackets $\{\Psi_i\}_i$ that take part in the time propagation by the current HagedornPropagator instance.

Parameters packet (Integer or None) – The index $i$ (in this list) of a single packet $\Psi_i$ that is to be returned. If set to None (default) return the full list with all packets.

Returns A list of HagedornWavepacket instances or a single instance.

propagate ()  
Given a wavepacket $\Psi$ at time $t$ compute the propagated wavepacket at time $t + \tau$. We perform exactly one timestep of size $\tau$ here. This propagation is done for all packets in the list $\{\Psi_i\}_i$ and neglects any interaction between two packets.

set_wavepackets (packetlist)  
Set the list $\{\Psi_i\}_i$ of wavepackets that the propagator will propagate.

1.1. WaveBlocks Classes
Parameters packetlist (A list of \((\Psi_i, \chi_i)\) tuples.) – A list of new wavepackets \(\Psi_i\) and their leading components \(\chi_i\) to propagate.

HagedornPropagatorInhomogeneous

About the HagedornPropagatorInhomogeneous class

The WaveBlocks Project

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Inheritance diagram

```
Propagator.Propagator → HagedornPropagatorInhomogeneous.HagedornPropagatorInhomogeneous
```

Class documentation

```csharp
WaveBlocksND.HagedornPropagatorInhomogeneous
alias of WaveBlocksND.HagedornPropagatorInhomogeneous
```

SimulationLoop

About the SimulationLoop class

The WaveBlocks Project

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Inheritance diagram

```
SimulationLoop.SimulationLoop
```

Class documentation

```csharp
class WaveBlocksND.SimulationLoop(parameters)

This class acts as the main simulation loop. It owns a propagator that propagates a set of initial values during a
```
time evolution. It is responsible for preparing the simulation, setting up initial values and store the simulation data with the help of an IOManager instance.

**end_simulation()**
Do the necessary cleanup after a simulation. For example request the IOManager to write the data and close the output files. Shut down the simulation process. :raise Not ImplementedError: This is an abstract base class.

**prepare_simulation()**
Set up a propagator for the simulation loop. Set the potential and initial values according to the configuration. :raise Not ImplementedError: This is an abstract base class.

**run_simulation()**
Run the simulation. This method will implement the central loop running over all timesteps. Inside of this loop it will call the propagate method of its propagator and save the simulation results. :raise Not ImplementedError: This is an abstract base class.

### SimulationLoopFourier

**About the SimulationLoopFourier class**

The WaveBlocks Project

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**Inheritance diagram**

```
SimulationLoop.SimulationLoop → SimulationLoopFourier.SimulationLoopFourier
```

**Class documentation**

**class WaveBlocksND.SimulationLoopFourier (parameters)**
This class acts as the main simulation loop. It owns a propagator that propagates a set of initial values during a time evolution.

**end_simulation()**
Do the necessary cleanup after a simulation. For example request the IOManager to write the data and close the output files.

**prepare_simulation()**
Set up a Fourier propagator for the simulation loop. Set the potential and initial values according to the configuration.

**Raises** ValueError For invalid or missing input data.

**run_simulation()**
Run the simulation loop for a number of time steps.
SimulationLoopHagedorn

About the SimulationLoopHagedorn class

The WaveBlocks Project
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Inheritance diagram

Class documentation
class WaveBlocksND.SimulationLoopHagedorn (parameters)
   This class acts as the main simulation loop. It owns a propagator that propagates a set of initial values during a time evolution.

   end_simulation()
      Do the necessary cleanup after a simulation. For example request the IOManager to write the data and close the output files.

   prepare_simulation()
      Set up a Hagedorn propagator for the simulation loop. Set the potential and initial values according to the configuration.

      Raises ValueError For invalid or missing input data.

   run_simulation()
      Run the simulation loop for a number of time steps.

SimulationLoopHagedornInhomogeneous

About the SimulationLoopHagedornInhomogeneous class

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Inheritance diagram

**Class documentation**

class WaveBlocksND\SimulationLoopHagedornInhomogeneous (parameters)

This class acts as the main simulation loop. It owns a propagator that propagates a set of initial values during a time evolution.

**end_simulation()**

Do the necessary cleanup after a simulation. For example request the IOManager to write the data and close the output files.

**prepare_simulation()**

Set up a Hagedorn propagator for the simulation loop. Set the potential and initial values according to the configuration.

**Raises** ValueError For invalid or missing input data.

**run_simulation()**

Run the simulation loop for a number of time steps.

**1.1.6 Simulation result storage I/O**

**IOManager**

**About the IOManager class**

The WaveBlocks Project

@author: R. Bourquin @copyright: Copyright (C) 2010, 2011, 2012 R. Bourquin @license: Modified BSD License
class WaveBlocksND.IOManager

An IOManager class that can save various simulation results into data files. For storing the data we use the well established HDF5 file format. An IOManager instace abstracts the input and output operations and translates requests into low-level operations.

create_block(blockid=None, groupid='global')

Create a data block with the specified block ID. Each data block can store several chunks of information, and there can be an arbitrary number of data blocks per file.

Parameters blockid – The ID for the new data block. If not given the blockid will be choosen automatically. The block ID has to be unique.

Returns The block ID of the created block.

create_file(parameters, filename='simulation_results.hdf5')

Set up a new :py:class 'IOManager' instance. The output file is created and opened.

Parameters

• parameters – A ParameterProvider instance containing the current simulation parameters. This is only used for determining the size of new data sets.

• filename – The filename (optionally with filepath) of the file we try to create. If not given the default value from GlobalDefaults is used.

create_group(groupid=None)

Create a data group with the specified group ID. Each data group can contain an arbitrary number of data blocks, and there can be an arbitrary number of data groups per file.

Parameters groupid – The ID for the new data group. If not given the group ID will be choosen automatically. The group ID has to be unique.

Returns The group ID of the created group.

finalize()

Close the open output file and reset the internal information.

find_timestep_index(timegridpath, timestep)

Lookup the index for a given timestep. This assumes the timegrid array is strictly monotone.

get_block_ids(groupid=None, grouped=False)

Return a list containing the IDs for all blocks in the current file structure.

Parameters

• groupid – An optional group ID. If given we return only block IDs for blocks which are a member of this group. If it is None we return all block IDs.

• grouped – If True we group the block IDs by their group into lists. This option is only relevant in case the groupid is not given.

group_ids(exclude=[])  

Return a list containing the IDs for all groups in the current file structure.

Parameters exclude – A list of group IDs to exclude. Per default no group is excluded.

group_of_block(blockid)

Return the ID of the group a given block belongs to or None if there is no such data block.

Parameters blockid – The ID of the given block.
**get_number_blocks** *(groupid=None)*

Return the number of data blocks in the current file structure.

**Parameters**

- **groupid** – An optional group ID. If given we count only data blocks which are a member of this group. If it is `None` (default) we count all data blocks.

**get_number_groups**

Return the number of data block groups in the current file structure.

**must_resize** *(path, slot, axis=0)*

Check if we must resize a given dataset and if yes, resize it.

**open_file** *(filename='simulation_results.hdf5')*

Load a given file that contains the results from another simulation.

**Parameters**

- **filename** – The filename (optionally with filepath) of the file we try to load. If not given the default value from `GlobalDefaults` is used.

**split_data** *(data, axis)*

Split a multi-dimensional data block into slabs along a given axis.

**Parameters**

- **data** – The data tensor given.
- **axis** – The axis along which to split the data.

**Returns**

A list of slices.

---

**IOM_plugin_parameters**

About the **IOM_plugin_parameters** class

The WaveBlocks Project

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**Class documentation**

The WaveBlocks Project

IOM plugin providing functions for handling simulation parameter data.

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**IOM_plugin_parameters.add_parameters** *(self, blockid='global')*

Add storage for the simulation parameters.

**Parameters**

- **blockid** – The ID of the data block to operate on.

**IOM_plugin_parameters.delete_parameters** *(self, blockid='global')*

Remove the stored simulation parameters.

**Parameters**

- **blockid** – The ID of the data block to operate on.

**IOM_plugin_parameters.has_parameters** *(self, blockid='global')*

Ask if the specified data block has the desired data tensor.

**Parameters**

- **blockid** – The ID of the data block to operate on.

**IOM_plugin_parameters.load_parameters** *(self, blockid='global')*

Load the simulation parameters.
Parameters **blockid** – The ID of the data block to operate on.

`.save_parameters(self, parameters, blockid='global')`

Save the simulation parameters.

Parameters

- **parameters** – The simulation parameters to store.
- **blockid** – The ID of the data block to operate on.

`.update_parameters(self, parameters, blockid='global')`

Update the parameters by some new values.

Parameters

- **parameters** – The parameters containing updated values.
- **blockid** – The ID of the data block to operate on.

### IOM_plugin_grid

**About the IOM_plugin_grid class**

The WaveBlocks Project

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### Class documentation

The WaveBlocks Project

IOM plugin providing functions for handling grid data.

@author: R. Bourquin @copyright: Copyright (C) 2010, 2011, 2012 R. Bourquin @license: Modified BSD License

`.add_grid(self, parameters, blockid=0)`

Add storage for a tensor product grid.

Parameters

- **parameters** – A `ParameterProvider` instance containing at least the keys `number_nodes` and `dimension`.
- **blockid** – The ID of the data block to operate on.

`.delete_grid(self, blockid=0)`

Remove the stored grid.

Parameters **blockid** – The ID of the data block to operate on.

`.has_grid(self, blockid=0)`

Ask if the specified data block has the desired data tensor.

Parameters **blockid** – The ID of the data block to operate on.

`.load_grid(self, blockid=0)`

Load the grid nodes.

Parameters **blockid** – The ID of the data block to operate on.

`.save_grid(self, gridnodes, blockid=0)`

Save the grid nodes.
Parameters

- **gridnodes** – The grid nodes to store.
- **blockid** – The ID of the data block to operate on.

**IOM_plugin_wavefunction**

**About the IOM_plugin_wavefunction class**

The WaveBlocks Project

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Class documentation

The WaveBlocks Project

IOM plugin providing functions for handling wavefunction data.

@author: R. Bourquin @copyright: Copyright (C) 2010, 2011, 2012 R. Bourquin @license: Modified BSD License

**IOM_plugin_wavefunction.add_wavefunction** *(self, parameters, timeslots=None, blockid=0)*

Add storage for the sampled wavefunction.

**Parameters**

- **parameters** – A ParameterProvider instance containing at least the keys *number_nodes* and *ncomponents*.
- **timeslots** – The number of time slots we need. Can be *None* to get automatically growing datasets.
- **blockid** – The ID of the data block to operate on.

**IOM_plugin_wavefunction.delete_wavefunction** *(self, blockid=0)*

Remove the stored wavefunction.

**Parameters**

- **blockid** – The ID of the data block to operate on.

**IOM_plugin_wavefunction.has_wavefunction** *(self, blockid=0)*

Ask if the specified data block has the desired data tensor.

**Parameters**

- **blockid** – The ID of the data block to operate on.

**IOM_plugin_wavefunction.load_wavefunction** *(self, timestep=None, blockid=0)*

Load the wavefunction values.

**Parameters**

- **timestep** – Load only the data of this timestep.
- **blockid** – The ID of the data block to operate on.

**IOM_plugin_wavefunction.load_wavefunction_timegrid** *(self, blockid=0)*

Load the wavefunction timegrid.

**Parameters**

- **blockid** – The ID of the data block to operate on.

**IOM_plugin_wavefunction.save_wavefunction** *(self, wavefunctionvalues, timestep=None, blockid=0)*

Save the values \( \psi_i \) of a WaveFunction instance.
Parameters

- **wavefunctionvalues** *(A list of ndarrays.)* – A list of the values to save.
- **timestep** – The timestep at which we save the data.
- **blockid** – The ID of the data block to operate on.

**IOM_plugin_fourieroperators**

About the **IOM_plugin_fourieroperators** class

The WaveBlocks Project

@author: R. Bourquin @copyright: Copyright (C) 2010, 2011, 2012 R. Bourquin @license: Modified BSD License

Class documentation

The WaveBlocks Project

IOM plugin providing functions for handling the propagation operators that appear in the Fourier algorithm.

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**IOM_plugin_fourieroperators.add_fourieroperators** *(self, parameters, blockid=0)*

Add storage for the Fourier propagation operators.

Parameters

- **parameters** – A ParameterProvider instance containing at least the keys `ncomponents` and `number_grid_nodes`.
- **blockid** – The ID of the data block to operate on.

**IOM_plugin_fourieroperators.delete_fourieroperators** *(self, blockid=0)*

Remove the stored Fourier operators.

Parameters **blockid** – The ID of the data block to operate on.

**IOM_plugin_fourieroperators.has_fourieroperators** *(self, blockid=0)*

Ask if the specified data block has the desired data tensor.

Parameters **blockid** – The ID of the data block to operate on.

**IOM_plugin_fourieroperators.load_fourieroperators** *(self, blockid=0)*

Load the Fourier operators.

Parameters **blockid** – The ID of the data block to operate on.

**IOM_plugin_fourieroperators.save_fourieroperators** *(self, operators, blockid=0)*

Save the kinetic and potential operator to a file.

Parameters

- **operators** – The operators to save, given as tuple `(T, V)`.
- **blockid** – The ID of the data block to operate on.
IOM_plugin_wavepacket

About the IOM_plugin_wavepacket class

The WaveBlocks Project

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Class documentation

The WaveBlocks Project

IOM plugin providing functions for handling homogeneous Hagedorn wavepacket data.

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IOM_plugin_wavepacket.add_wavepacket (self, parameters, timeslots=None, blockid=0)

Add storage for the homogeneous wavepackets.

Parameters

• parameters – An ParameterProvider instance with at least the keys dimension and ncomponents.

• timeslots – The number of time slots we need. Can be None to get automatically growing datasets.

• blockid – The ID of the data block to operate on.

IOM_plugin_wavepacket.delete_wavepacket (self, blockid=0)

Remove the stored wavepackets.

Parameters

• blockid – The ID of the data block to operate on.

IOM_plugin_wavepacket.has_wavepacket (self, blockid=0)

Ask if the specified data block has the desired data tensor.

Parameters

• blockid – The ID of the data block to operate on.

IOM_plugin_wavepacket.load_wavepacket_basisslices (self, the_hash=None, blockid=0)

Load the basis shapes by hash.

Parameters

• the_hash – The hash of the basis shape whose description we want to load.

• blockid – The ID of the data block to operate on.

IOM_plugin_wavepacket.load_wavepacket_coefficients (self, timestep=None, get_hashes=False, component=None, blockid=0)

Load the wavepacket coefficients.

Parameters

• timestep – Load only the data of this timestep.

• get_hashes – Return the corresponding basis shape hashes.

• component – Load only data from this component.

• blockid – The ID of the data block to operate on.
Load the wavepacket description.

**Parameters**
- **blockid** – The ID of the data block to operate on.

Load the wavepacket parameters.

**Parameters**
- **timestep** – Load only the data of this timestep.
- **blockid** – The ID of the data block to operate on.

Load the wavepacket timegrid.

**Parameters**
- **blockid** – The ID of the data block to operate on.

Save the basis shapes of the Hagedorn wavepacket to a file.

**Parameters**
- **basisshape** – The basis shape of the Hagedorn wavepacket.
- **blockid** – The ID of the data block to operate on.

Save the coefficients of the Hagedorn wavepacket to a file. Warning: we do only save the hash of the basis shapes here! You have to save the basis shape with the corresponding function too.

**Parameters**
- **coefficients** (A list with $N$ suitable ndarrays.) – The coefficients of the Hagedorn wavepacket.
- **basisshapes** (A list with $N$ BasisShape subclass instances.) – The corresponding basis shapes of the Hagedorn wavepacket.
- **timestep** – The timestep at which we save the data.
- **blockid** – The ID of the data block to operate on.

Save the description of this wavepacket.

**Parameters**
- **descr** – The description.
- **blockid** – The ID of the data block to operate on.

Save the parameter set $\Pi$ of the Hagedorn wavepacket $\Psi$ to a file.

**Parameters**
- **parameters** (A list containing the five ndarrays like $(q, p, Q, P, S)$) – The parameter set of the Hagedorn wavepacket.
- **timestep** – The timestep at which we save the data.
- **blockid** – The ID of the data block to operate on.
IOM_plugin_inhomogwavepacket

About the IOM_plugin_inhomogwavepacket class

The WaveBlocks Project

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Class documentation

The WaveBlocks Project

IOM plugin providing functions for handling homogeneous Hagedorn wavepacket data.

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IOM_plugin_inhomogwavepacket.add_inhomogwavepacket (self, parameters, timeslots=None, blockid=0)

Add storage for the inhomogeneous wavepackets.

Parameters parameters – An ParameterProvider instance with at least the keys dimension and ncomponents.

IOM_plugin_inhomogwavepacket.delete_inhomogwavepacket (self, blockid=0)

Remove the stored wavepackets.

IOM_plugin_inhomogwavepacket.has_inhomogwavepacket (self, blockid=0)

Ask if the specified data block has the desired data tensor.

IOM_plugin_inhomogwavepacket.load_inhomogwavepacket_basisshapes (self, the_hash=None, blockid=0)

Load the basis shapes by hash.

IOM_plugin_inhomogwavepacket.save_inhomogwavepacket_basisshapes (self, basisshape, blockid=0)

Save the basis shapes of the Hagedorn wavepacket to a file.

Parameters coefficients – The basis shapes of the Hagedorn wavepacket.

IOM_plugin_inhomogwavepacket.save_inhomogwavepacket_coefficients (self, coefficients, basisshapes, timestep=None, blockid=0)

Save the coefficients of the Hagedorn wavepacket to a file. Warning: we do only save the hash of the basis shapes here! You have to save the basis shape with the corresponding function too.

Parameters

• coefficients (A list with N suitable ndarrays.) – The coefficients of the Hagedorn wavepacket.

• basisshapes (A list with N BasisShape subclass instances.) – The corresponding basis shapes of the Hagedorn wavepacket.

IOM_plugin_inhomogwavepacket.save_inhomogwavepacket_parameters (self, parameters, timestep=None, blockid=0)

Save the parameter set Π of the Hagedorn wavepacket Ψ to a file.
Parameters parameters (A list containing the five ndarrays like \((q, p, Q, P, S)\)) – The parameter set of the Hagedorn wavepacket.

IOM_plugin_norm

About the IOM_plugin_norm class

The WaveBlocks Project

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Class documentation

The WaveBlocks Project

IOM plugin providing functions for handling norm data.

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IOM_plugin_norm.add_norm (self, parameters, timeslots=None, blockid=0)

Add storage for the norms.

Parameters

- parameters – A ParameterProvider instance containing at least the key ncomponents.
- timeslots – The number of time slots we need. Can be None to get automatically growing datasets.
- blockid – The ID of the data block to operate on.

IOM_plugin_norm.delete_norm (self, blockid=0)

Remove the stored norms.

Parameters

- blockid – The ID of the data block to operate on.

IOM_plugin_norm.has_norm (self, blockid=0)

Ask if the specified data block has the desired data tensor.

Parameters

- blockid – The ID of the data block to operate on.

IOM_plugin_norm.load_norm (self, timestep=None, split=False, blockid=0)

Load the norm data.

Parameters

- timestep – Load only the data of this timestep.
- split – Split the data array into one array for each component.
- blockid – The ID of the data block to operate on.

IOM_plugin_norm.load_norm_timegrid (self, blockid=0)

Load the timegrid corresponding to the norm data.

Parameters

- blockid – The ID of the data block to operate on.

IOM_plugin_norm.save_norm (self, norm, timestep=None, blockid=0)

Save the norm of wavefunctions or wavepackets.

Parameters

- norm – The norm values to save.
• **timestep** – The timestep at which we save the data.
• **blockid** – The ID of the data block to operate on.

### IOM_plugin_energy

#### About the IOM_plugin_energy class

The WaveBlocks Project

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Class documentation

The WaveBlocks Project

IOM plugin providing functions for handling energy data.

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#### IOM_plugin_energy.add_energy

```python
IOM_plugin_energy.add_energy(self, parameters, timeslots=None, blockid=0, key=('kin', 'pot'))
```

**Parameters**

- **parameters** – A ParameterProvider instance containing at least the key `ncomponents`.
- **timeslots** – The number of time slots we need. Can be `None` to get automatically growing datasets.
- **blockid** – The ID of the data block to operate on.
- **key** (Tuple of valid identifier strings that are `kin`, `pot` and `tot`. Default is `"kin", "pot"`.) – Specify which energies to save. All are independent.

#### IOM_plugin_energy.delete_energy

```python
IOM_plugin_energy.delete_energy(self, blockid=0)
```

Remove the stored energies

**Parameters**

- **blockid** – The ID of the data block to operate on.

#### IOM_plugin_energy.has_energy

```python
IOM_plugin_energy.has_energy(self, blockid=0, key=('kin', 'pot'))
```

Ask if the specified data block has the desired data tensor.

**Parameters**

- **blockid** – The ID of the data block to operate on.
- **key** (Tuple of valid identifier strings that are `kin`, `pot` and `tot`. Default is `"kin", "pot"`.) – Specify which energies to save. All are independent.

#### IOM_plugin_energy.load_energy

```python
IOM_plugin_energy.load_energy(self, timestep=None, split=False, blockid=0, key=('kin', 'pot'))
```

Load the energy data.

**Parameters**

- **timestep** – Load only the data of this timestep.
- **split** – Split the array into arrays for each component.
- **blockid** – The ID of the data block to operate on.
- **key** (Tuple of valid identifier strings that are `kin`, `pot` and `tot`. Default is `"kin", "pot"`.) – Specify which energies to save. All are independent.
IOM_plugin_energy.load_energy_timegrid(self, blockid=0, key=('kin', 'pot'))
Load the time grid for specified energies.

Parameters
- **blockid** – The ID of the data block to operate on.
- **key** (Tuple of valid identifier strings that are *kin*, *pot* and *tot*. Default is ("kin", "pot".)) – Specify which energies to save. All are independent.

IOM_plugin_energy.save_energy(self, energies, timestep=None, blockid=0, key=('kin', 'pot'))
Save the kinetic and potential energies to a file.

Parameters
- **energies** – A tuple containing the energies. The order is important, it has to match the order in the *key* argument. Per default the order has to be \((E_{\text{kin}}, E_{\text{pot}})\).
- **timestep** – The timestep at which we save the data.
- **blockid** – The ID of the data block to operate on.
- **key** (Tuple of valid identifier strings that are *kin*, *pot* and *tot*. Default is ("kin", "pot".).) – Specify which energies to save. All are independent.

1.1.7 Other classes

GlobalDefaults

About the `GlobalDefaults` class

The WaveBlocks Project

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Class documentation

The WaveBlocks Project

This file contains some global defaults, for example file names for output files. If a `ParameterProvider` instance is asked about a key which it does not know about it tries to look it up here to see if a default value is available.

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ParameterLoader

About the `ParameterLoader` class

The WaveBlocks Project

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Inheritance diagram

```
ParameterLoader.ParameterLoader
```

Class documentation

class WaveBlocksND.ParameterLoader

    load_from_dict(adict)
    Construct a ParameterProvider instance from a common python key-value dict.
    Parameters adict – A plain python dict with key-value pairs.
    Returns A ParameterProvider instance.

    load_from_file(filepath)
    Read the parameters from a configuration file.
    Parameters filepath – Path to the configuration file.
    Returns A ParameterProvider instance.

ParameterProvider

About the ParameterProvider class

The WaveBlocks Project

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Inheritance diagram

```
ParameterProvider.ParameterProvider
```

1.1. WaveBlocks Classes
Class documentation

class WaveBlocksND.ParameterProvider

    compute_parameters()
    Compute some further parameters from the given ones.

    get_parameters()
    Return a copy of the dict containing all parameters. @return: A copy of the dict containing all parameters.
    The dict will be copied.

    get_timemanager()
    Return the embedded I{TimeManager} instance.

    set_parameters(params)
    Overwrite the dict containing all parameters with a newly provided dict with (possibly) changed parameters.
    @param params: A I{ParameterProvider} instance or a dict with new parameters. The values will be
    deep-copied. No old values will remain.

    update_parameters(params)
    Overwrite the dict containing all parameters with a newly provided dict with (possibly) changed parameters.
    @param params: A I{ParameterProvider} instance or a dict with new parameters. The values will be
    deep-copied. Old values are only overwritten if we have got new values.

FileTools

About the FileTools class

The WaveBlocks Project
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Class documentation

The WaveBlocks Project

This file contains various functions for finding and retrieving the files that contain parameter settings and simulation
results. Note: The terms ‘path’ and ‘ID’ are used as synonyms here. Each simulation ID is just the basename of the
path or the configuration file.
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FileTools.compare_by (namea, nameb, pattern, ldel='[', mdel='=', rdel=']', as_string=True)
Compare two simulation IDs with respect to a (numerical) value in the ID.

Parameters

- namea – The first name in the comparison
- nameb – The second name in the comparison
- pattern – The pattern whose (numerical) value is used for sorting
- ldel – Left delimiter of the pattern
- mdel – Middle delimiter of the pattern
- rdel – Right delimiter of the pattern
• **as_string** – Determines if the values for *pattern* get converted to floats

**Returns** A boolean answer if the IDs are the same w.r.t the pattern.

FileTools.*gather_all*(stringlist, pattern)
Collects all simulation IDs which contain a specific pattern from a given list.

**Parameters**

• **stringlist** – A list with the simulation IDs

• **pattern** – The pattern

**Returns** A list of simulation IDs that contain the given pattern.

FileTools.*get_max_by*(stringlist, pattern, ldel='[', mdel='=', rdel=']', as_string=False)
Get the maximum of a list with simulation IDs with respect to a (numerical) value in the ID. This is just a simple convenience function so that the user needs not to remember if the sort order is ascending or descending which plays no role for iteration.

**Parameters**

• **stringlist** – A list with the simulation IDs

• **pattern** – The pattern whose (numerical) value is used for sorting

• **ldel** – Left delimiter of the pattern

• **mdel** – Middle delimiter of the pattern

• **rdel** – Right delimiter of the pattern

• **as_string** – Determines if the values for *pattern* get converted to floats

**Returns** A sorted list of simulation IDs.

FileTools.*get_min_by*(stringlist, pattern, ldel='[', mdel='=', rdel=']', as_string=False)
Get the minimum of a list with simulation IDs with respect to a (numerical) value in the ID. This is just a simple convenience function so that the user needs not to remember if the sort order is ascending or descending which plays no role for iteration.

**Parameters**

• **stringlist** – A list with the simulation IDs

• **pattern** – The pattern whose (numerical) value is used for sorting

• **ldel** – Left delimiter of the pattern

• **mdel** – Middle delimiter of the pattern

• **rdel** – Right delimiter of the pattern

• **as_string** – Determines if the values for *pattern* get converted to floats

**Returns** A sorted list of simulation IDs.

FileTools.*get_number_simulations*(path)
Get the number of simulations at hand below the given path.

**Parameters** path – The path under which we search for a output file.

**Returns** The number of simulations result directories.

FileTools.*get_parameters_file*(path)
Search for a configuration file containing the simulation parameters under a given path. Note that in case there are more than one .py file under the given path we just return the first one found!

1.1. WaveBlocks Classes 75
**Parameters** `path` – The path under which we search for a configuration file.

**Returns** The path (filename) of the configuration file.

**FileTools.get_result_dirs (path)**
Lists all simulations (IDs) that can be found under the given path.

**Parameters** `path` – The filesystem path under which we search for simulations.

**Returns** A list of simulation IDs.

**FileTools.get_results_file (path)**
Search for a file containing the simulation results under a given path. Note that in case there are more than one .hdf5 file under the given path we just return the first one found!

**Parameters** `path` – The path under which we search for a output file.

**Returns** The path (filename) of the output file.

**FileTools.group_by (stringlist, pattern, ldel='[', mdel='=', rdel=']', as_string=True)**
Groups simulation IDs with respect to a pattern.

**Parameters**
- `stringlist` – A list with the simulation IDs
- `pattern` – The pattern used for grouping
- `ldel` – Left delimiter of the pattern
- `mdel` – Middle delimiter of the pattern
- `rdel` – Right delimiter of the pattern
- `as_string` – Determines if the values for `pattern` get converted to floats. Not used here.

**Returns** A list of groups of simulation IDs.

**FileTools.intersect_by (lista, listb, pattern, ldel='[', mdel='=', rdel=']', as_string=True)**
Find the intersection of two lists containing simulation IDs.

**Parameters**
- `lista` – A first list with the simulation IDs
- `listb` – A second list with the simulation IDs
- `pattern` – The pattern whose numerical value is used for sorting
- `ldel` – Left delimiter of the pattern
- `mdel` – Middle delimiter of the pattern
- `rdel` – Right delimiter of the pattern
- `as_string` – Determines if the values for `pattern` get converted to floats

**Returns** A sorted list of simulation IDs.

**FileTools.name_contains (name, pattern)**
Checks if a simulation ID contains a given pattern.

**Parameters**
- `name` – The full simulation ID.
- `pattern` – The pattern in question.

**Returns** A boolean answer.
FileTools.sort_by (stringlist, pattern, ldel='[', mdel='=', rdel=']', as_string=False)

Sorts simulation IDs with respect to a (numerical) value in the ID.

**Parameters**

- **stringlist** – A list with the simulation IDs
- **pattern** – The pattern whose (numerical) value is used for sorting
- **ldel** – Left delimiter of the pattern
- **mdel** – Middle delimiter of the pattern
- **rdel** – Right delimiter of the pattern
- **as_string** – Determines if the values for pattern get converted to floats

**Returns** A sorted list of simulation IDs.
2.1 Citation

For citation of this project please use one of the following bibtex snippets. The code for one-dimensional simulations:

```bibtex
@misc{waveblocks,
  author = {R. Bourquin and V. Gradinaru},
  title = {{WaveBlocks}: Reusable building blocks for simulations with semiclassical wavepackets},
  year = {2010, 2011},
  url = {https://github.com/raoulbq/WaveBlocks},
  howpublished={https://github.com/raoulbq/WaveBlocks}
}
```

The code for multi-dimensional simulations:

```bibtex
@misc{waveblocksnd,
  author = {R. Bourquin and V. Gradinaru},
  title = {{WaveBlocks}: Reusable building blocks for simulations with semiclassical wavepackets},
  year = {2010, 2011, 2012},
  url = {https://github.com/raoulbq/WaveBlocksND},
  howpublished={https://github.com/raoulbq/WaveBlocksND}
}
```
CHAPTER THREE

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