# **Bonsu - The Interactive Phase Retrieval Suite Documentation**

Release 3.4.2

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# **CONTENTS**

1	Installation       1.1 Installation on Mac OS X        1.2 Installation on Windows        1.3 Installation on Linux/Unix	1 1 1 2
2	Introduction 2.1 Overview	5 5 5 5
3		7 7 10 10
4	4.1 History of the Interactive Phase Retrieval Suite	11 11 11
5	5.1 Library Reference - Visualisation Tools 5.2 Library Reference - Import Tools 5.3 Library Reference - Export Tools 5.4 Library Reference - Functions 5.5 Library Reference - Phasing Algorithms	13 13 15 17 17 23 31
6	6.1 Core concepts	33 33 35 36 37
7	Indices and tables	45
Py	thon Module Index	47
Ind	lex	49

### INSTALLATION

# 1.1 Installation on Mac OS X

If a native Python environment is used, ensure that all dependencies are installed, ideally from the Python Package Index using pip. They are listed below.

- wxPython
- NumPy
- VTK
- Pillow
- H5Py

A stand alone python distribution such as Anaconda Python or Enthought Canopy will likely include all of the above prerequisites.

It is then possible to install Bonsu from the Python Package Index using the following command:

\$ pip install bonsu

# 1.2 Installation on Windows

Download 64-bit Python  $\geq$  3.7 for Microsoft Windows from the Python website and follow the installation instructions. Ensure that pip is included in the installation. Prerequisite binaries are available for download in the Python Package Index. They are also listed below.

- wxPython
- NumPy
- VTK
- Pillow
- H5Py

Prerequisites are installed using the pip command found in the PythonScript folder. For example:

\$ pip.exe install wxpython

**Note:** Prebuilt VTK packages  $\geq 7.0$  in PyPI generally requre a graphics card with OpenGL 3 support.

Alternatively, it is possible to install a stand alone python distribution such as Anaconda Python or Enthought Canopy. This will likely include all of the above prerequisites.

It is then possible to install Bonsu from the Python Package Index using the following command

```
$ pip.exe install bonsu
```

# 1.3 Installation on Linux/Unix

# 1.3.1 System Preparation

Before Bonsu can be installed from source, a number of software packages are required. They are namely:

- Python
- wxPython
- FFTW with threading
- NumPy (FFTW aware)
- VTK (compiled with python bindings)
- Python Imaging Library (Pillow)
- H5Py
- A reasonably new c<sup>++</sup>compiler such as gcc

As most Linux systems and services require Python natively, it is likely that Python is already provided in your distribution. Also, many distributions package all of the above prerequisites. Please check your distributions repositories.

It is also possible to install a stand alone python distribution such as that provided by Anaconda or Enthought. This will include most of the above prerequisites. In order to compile and use any additional packages (such as FFTW) into the stand alone distribution, a number of environmental variables should be set. In Anaconda Python, ensure that you create and activate your local environment:

```
$ conda create --name myenv
$ conda activate myenv
$ export PREFIX="/path/to/conda/myenv/"
```

### 1.3.2 Note on Building FFTW

To build FFTW into a stand alone python distribution such as that provided by Enthought , use the following sequence of commands:

```
$ ./configure --prefix=${PREFIX} --enable-threads --enable-shared
$ make
$ make install
```

### 1.3.3 Installing Bonsu

Source code for Bonsu is available for download here.

#### See also:

https://github.com/bonsudev/bonsu

Unpack the source code with the following:

```
$ tar -xvf Bonsu-{x}.tar.gz
```

where x is the current version.

Enter the newly created directory and first build the source:

### \$ python setup.py build

If this is successful you will now have a fully prepared built source tree for installation. To install the package type the following (super user privileges may be required):

```
$ python setup.py install
```

To install the package to a non-default location (perhaps where super user privileges are not required) use the following command:

```
$ python setup.py install --home=/my/install/path
```

The --home option defines the installation base directory. Files are installed to the following directories under the installation base as follows

Type of file	e of file   Installation directory	
modules	dules home/lib/python	
scripts	pts home/bin	
data	home	
C headers	headers home/include/python/distname	

To make Python find the distributions installed with this scheme, it is necessary to modify Python's search path(s) to include the above.

To start the program, execute the following from the command line:

### \$ bonsu &

Note: For Linux users, a link to the program should be visible in the applications menu.

### INTRODUCTION

# 2.1 Overview

Bonsu, the interactive phase retrieval suite, is the first phase retrieval software package for phase retrieval with realtime visualisation in both two and three dimensions. It is complete with an inventory of algorithms and routines for data manipulation and reconstruction.

Bonsu is open-source, is designed around the python language (with c bindings) and is largely platform independent. It is also able to handle data in formats such as:

- 1. SPE
- 2. HDF5
- 3. VTK
- 4. Numpy
- 5. Image formats including PNG, JPG, TIFF, PPM

# 2.2 Arrays

Bonsu uses NumPy array format (http://numpy.org) internally for manipulating and outputting raw data. Double-precision complex NumPy arrays are used for phase retrieval. NumPy is part of the 'Scientific Computing Tools For Python' or SciPy package (http://scipy.org) available in Python. There are a number of standard tools made available for array conversion, manipulation and viewing. Tools for cropping, padding, binning, support array creation and mask array creation are provided. Arrays can be exported in both NumPy array and VTK format.

For further details see Array Creation and Storage.

# 2.3 Algorithms

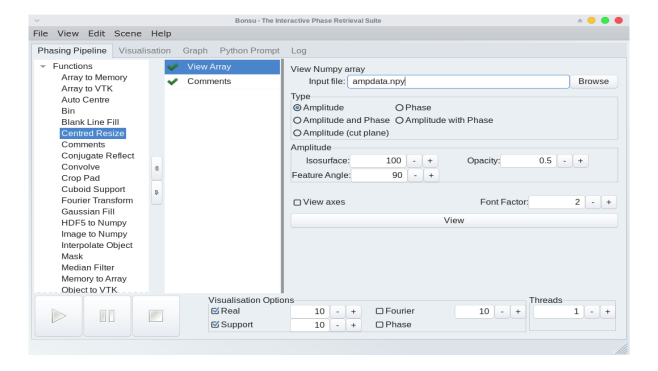
Bonsu comes complete with a number of *algorithms* for phase retrieval. They include Fienup's hybrid inputoutput (HIO) (Fienup, 1982), HIO with positivity constraint, phase-constrained HIO (Harder et al., 2010), hybrid projection reflection (Bauschke et al., 2003), relaxed average alternating reflection (Luke, 2005) and the error reduction algorithm (Gerchberg & Saxton, 1972). In addition, a shrink-wrap algorithm that is able to make use of either of the above algorithms is also provided (Marchesini et al., 2003; Newton et al., 2010). Phase retrieval algorithms are implemented in the c language for fast execution and seamlessly integrated into the application.

For a complete list see *Library Reference – Phasing Algorithms*.

### **GENERAL USAGE**

The following describes the general usage procedure when interacting with the graphical user interface. If you would like to script with Bonsu, there are at least two options available. The first option is to use the *Python Script function* which permits arbitrary operations on arrays. This is useful when combining custom operations on arrays with native functions. The second option is to import Bonsu into a Python script. The procedure for doing so is described in detail *here*.

# 3.1 User Interface



The graphical user interface is divided into five tabs which are namely Phasing Pipeline, Visualisation, Graph, Python Prompt and Log. Algorithms and tools are manipulated and executed on the Phasing Pipeline tab. During algorithmic phase retrieval, real-time images of the object during reconstruction are shown on the Visualisation tab. Information regarding the statistical error during phase retrieval is displayed on the Graph tab. The Log tab provides the user with any information the various algorithms and tools return during execution. The Python Prompt is, as it suggests, a fully functioning interactive python shell.

When loading and saving files, the current working directory where the program was loaded is used as the default path. It is possible to change the working directory by selecting  $Edit \rightarrow Current \ Working \ Directory$  from the main

menu.

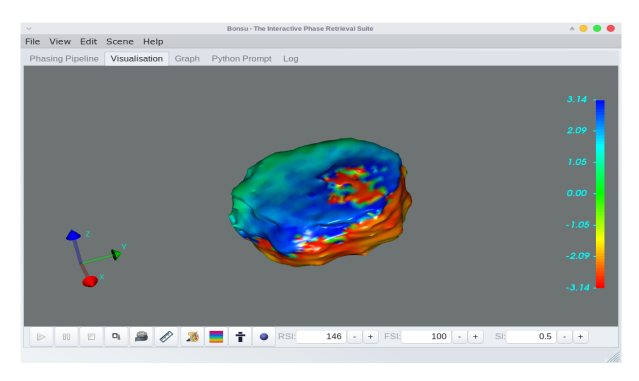
The remainder of this section describes in further detail the use of each graphical interface tab.

# 3.1.1 Phasing Pipeline

Most user interaction occurs in the Phasing Pipeline tab. This tab is divided into three columns. From left to right, the first shows a drop-down menu divided into three sections where various array functions, phasing algorithms and posterior phasing operations can be found. For a complete list of available operations and algorithms, please see the *Library Reference*. Once selected from the drop-down menu (with a carriage return or mouse double click), an instance of the selected item appears in the centre column (or pipeline), where it can be selected in the pipeline and properties modified for that instance. The properties appear in the right-hand column. Check boxes to the immediate left of each item in the pipeline provide the option to exempt that item from execution without removing the item from the pipeline. By simply selecting the check so that a cross is displayed, the item will be ignored during execution of the pipeline. This is often useful for testing purposes. Once items have been added to the pipeline, it is possible to save the contents of the pipeline and all custom settings by simply choosing Save from the menu. At this point, the user is prompted for a file name. See *Saving Sessions* for further details.

Below the three columns are the control buttons Start, Pause and Stop. These buttons will, respectively, execute all items in the pipeline sequentially in the order that they appear, pause the execution of an algorithm or stop the execution of an algorithm. The remaining checkboxes control the type of visualization that appears on the 'Visualisation' tab and the maximum number of threads used by the Fourier transform algorithm. The options include no visualization, real-space amplitude, real-space amplitude with phase, Fourier-space amplitude and Fourier-space amplitude with phase.

### 3.1.2 Visualisation



In order to view real-time visualisation, either real-space or Fourier-space must be selected from the visualisation options on the Phasing Pipeline tab. This is the default. The Visualisation tab is where a rendered scene of the reconstruction will appear. From here the user can interact fully with and manipulate an object generated with either a view function (e.g. *View Array* or *View Object*) or an algorithm during reconstructed

(see *Library Reference – Phasing Algorithms*). Either a two-dimensional or three-dimensional object is displayed depending on the dimensions of the data. In both cases, pressing the "p" keyboard button will display the current co-ordinates of the mouse cursor and print these coordinates to the *Log*.

It is also possible to undock the rendered scene by selecting  $View \to Visualisation \to Undock$  from the main menu. This will open the current visualisation in a new window. Selecting  $View \to Visualisation \to Dock$  returns the rendered scene back to the Visualisation tab.

Below the rendered scene, there are a number of buttons for manipulating the visualisation. The control buttons Start, Pause and Stop are identical in operation to those on the Phasing Pipeline tab and are present for convenience. To save images of the object for further use, select the save button. After this, no prompt for a file name will appear. Instead, the file will be saved to the working directory with the present date and time used as a filename string. The Measure scene button can be used to perform measurements on the rendered object and to view the orientation widget. The Scene background button will open a window that can be used to modify the background colour of the rendered scene. The Lookup table button will open a window which can be used to change the colour map for the amplitude and phase of the rendered object. The Data range button will open a window that will allow the user to adjust the data range used for the colour map. The Isosurface button will display three number fields for the Real space isosurface (RSI), Fourier space isosurface (FSI) and Support isosurface (SI).

### 3.1.3 Visualisation: Animation

A feature for animating the rendered scene is made available by selecting the Animate button. From here a window appears with various options for scene animation. The axis about which the rotation occurs can be selected by choosing a value greater than zero. Either a single axis or a composite of the three rectilinear (x,y,z) axes can be selected. If multiple axis are chosen, their vector sum is used as the final axis of rotation. The rendered scene is generally rotated about the centre of mass.

Options for setting the angle of increment and total number of rotation steps are also available. It is also possible to save an image for each step by selecting save scene and providing a file name. Indices of the animation step will be appended to the filename automatically.

### 3.1.4 Visualisation: Measurement

Features for performing measurements on the rendered scene are made available by selecting the Measure scene button on the Visualisation tab. A window appears which shows a number of tabs for measurement. When enabled, the Line Scan tab provides a feature for arbitrary direction line scans measurements in both two and three dimensions. A line will appear in the render window with handles at either end for manipulation. As the line is stretched and moved, the corresponding line scan through the data appears on the window tab. Information on the line properties are also displayed on the *Log* tab. To save the data, select Save Data from the window tab. No prompt for a file name will appear. Instead, the file will be saved in csv format to the working directory with the present date and time used as a filename string.

When enabled, the Angle tab provides a feature for measuring arbitrary angle in both two and three dimensions. In order to begin using this feature, the user must first select three points using the mouse cursor in the render scene. These points will become the vertices of the angular measurement.

3.1. User Interface 9

# 3.1.5 **Graph**

During phase reconstruction, the Graph tab displays the reconstruction residual error plotted on a graph with dynamic axes. From here the graph can be paused (independently of the reconstruction process) and the data saved to a text file. No prompt for a file name will appear. Instead, the file will be saved in csv format to the working directory with the present date and time used as a filename string. It is also possible to save the reconstruction residual error information automatically after the phase retrieval process, by appending *Save Residual* to the phasing pipeline.

# 3.1.6 Python Prompt

The 'Python Prompt' tab enables the user to perform additional tasks through the Python interpreter. This might be the execution of additional Python scripts or additional array operations with standard Numpy tools. Data arrays stored in memory using the memoryn keywords (see *Array Creation and Storage* below) are accessible as variables of the same name and can be manipulated as NumPy arrays. Changes made to memory variable arrays will affect subsequent use of that array in the phasing pipeline.

# 3.1.7 Log

The 'Log' tab allows the user to view additional information on the program's operation and any graph data values in text form. From here the log data can be saved to a text file. No prompt for a file name will appear. Instead, the file will be saved in txt format to the working directory with the present date and time used as a filename string.

# 3.2 Array Creation and Storage

NumPy array format is used internally for data manipulation. NumPy arrays loaded from file are converted to double-precision complex format in row-major order (c style). NumPy arrays are also saved in this format by default.

When a large number of array operations are performed, it is often desirable to store the result in memory at each intermediate step. This can help to reduce disk usage and improve overall performance at the expense of additional memory usage. Bonsu provides a way to store and access arrays in memory, that is accessible to all operations and algorithms. By using either of the keywords memory0, memory1, ..., memoryn (where n is a positive integer) as a string in place of a NumPy file path, it is possible to save to memory an array manipulated by any phasing algorithm or operation that uses NumPy arrays. The array is then accessible by using the corresponding keyword in place of the NumPy file path when attempting to load an array. It is also possible to directly visualise the array using *View Array*.

In addition to the above, keywords memorysequence and memorycoords can be used to view or use *sequence data* and *Co-ordinate Transformation* arrays respectively. This is useful for using *View Object* exclusively with arrays in memory.

# 3.3 Saving a Session

It is possible to save the contents of the pipeline and all customised settings by simply choosing Save from the program menu. Pipeline data is saved in '.fin' file format. Similarly, to recover a pipeline, simply select open from the program menu and locate the saved '.fin' file. Bonsu also allows pipeline entries to be appended. For example, opening a '.fin' file twice will append the list of tasks twice into the pipeline. To clear the pipeline, simply select New from the menu.

**CHAPTER** 

**FOUR** 

# **HISTORY AND LICENCE**

# 4.1 History of the Interactive Phase Retrieval Suite

The Interactive Phase Retrieval Suite was first conceived in 2011 by Marcus Newton with the major design goal of providing a complete set of tools for phase reconstruction and visualisation. It was intended as the successor to previous *command line only* tools which were widely in use at the time. It aimed to be encompassing and placed emphasis on maintaining reproducibility on various platforms. It achieved this largely by providing the means to easily share encapsulated data analysis parameters.

The name *Bonsu* means whale or large marine mammals of the Cetacea group, in the Akan language of western African. This was chosen as a sentiment to the "eye-like" focus of the sound beam that certain cetacean mammals emit during sonar echo location.

# 4.2 Licence

All releases of the Interactive Phase Retrieval Suite are Open Source (see http://www.opensource.org/ for the Open Source Definition). Bonsu, the Interactive Phase Retrieval Suite, is distributed under the GNU General Public License. A copy of the GNU General Public License is provided with the program. Also see http://www.gnu.org/licenses/.

# LIBRARY REFERENCE

# 5.1 Library Reference - Visualisation Tools

### 5.1.1 Comments

Input comments into the pipeline.

### 5.1.2 Nexus File Viewer for I16

Diamond Light Source Nexus File Viewer for Beamline I16

# 5.1.3 View Array

Render an image of the two or three-dimensional input Numpy array to the Visualisation tab.

### Parameters:

**Input file** User provided path string to the input NumPy file.

### Type

- Amplitude An iso-surface of the array amplitude.
- Phase An interactive scalar cut plane of the array phase.
- Amplitude and Phase An iso-surface of the array amplitude and an interactive scalar cut plane of the array phase.
- Amplitude with Phase An iso-surface of the array amplitude with the array phase mapped onto the iso-surface.
- Amplitude (cut plane) An interactive scalar cut plane of the array amplitude.

# Amplitude

- Iso-surface Scalar value for the rendered iso-surface.
- Opacity Opacity of the isosurface when viewing Amplitude and Phase.
- Feature angle Minimum angle between two surface normals of the rendered object.

#### Phase

- Max Maximum phase.
- Min Minimum phase.

Origin Origin point of the interactive scalar cut plane.

Normal Normal vector to the interactive scalar cut plane.

**Spacing** Spacing of the input array indices.

View Axes Optionally view the axes of the array.

# 5.1.4 View Object

Render an object of the three-dimensional input Numpy array with it's corresponding co-ordinates array to the *Visualisation tab*.

#### Parameters:

**Input file** User provided path string to the input NumPy file.

**Co-ord's File** User provided path string to the corresponding co-ordinates NumPy array file. Alternatively, the *memory array* keyword memorycoords provides access to the internally stored coordinate array. Generating a coordinate array is achieved using the *Coordinate transformation* operation.

### **Type**

- Amplitude An iso-surface of the object amplitude.
- **Phase** An interactive scalar cut plane of the object phase.
- **Amplitude and Phase** An iso-surface of the object amplitude and an interactive scalar cut plane of the object phase.
- Amplitude with Phase An iso-surface of the object amplitude with the object phase mapped onto the iso-surface.
- Amplitude (cut plane) An interactive scalar cut plane of the object amplitude.

### Amplitude

- **Iso-surface** Scalar value for the rendered iso-surface.
- Opacity Opacity of the isosurface when viewing Amplitude and Phase.
- Feature angle Minimum angle between two surface normals of the rendered object.

### Phase

- Max Maximum phase.
- Min Minimum phase.

**Origin** Origin point of the interactive scalar cut plane.

Normal Normal vector to the interactive scalar cut plane.

View Axes Optionally view the axes of the object.

# 5.1.5 View Support

Render an image of a three-dimensional input Numpy data array and its Support array to the Visualisation tab.

### **Parameters**:

**Support file** User provided path string to the NumPy array file.

Data array file User provided path string to the NumPy array file.

### **Isosurface options**

- Support iso-surface Scalar value for the rendered iso-surface on the interval [0,1].
- **Opacity** Opacity of the support isosurface.
- Data file iso-surface Scalar value for the rendered iso-surface.
- Feature angle Minimum angle between two surface normals of the rendered object.

View Axes Optionally view the axes of the array.

# 5.1.6 View VTK Array

Render an input VTK array to the Visualisation tab.

#### **Parameters:**

**Input file** User provided path string to the input NumPy file.

### Type

- Amplitude An iso-surface of the array amplitude.
- Phase An interactive scalar cut plane of the array phase.
- Amplitude (cut plane) An interactive scalar cut plane of the array amplitude.

### **Amplitude**

- Iso-surface Scalar value for the rendered iso-surface.
- Feature angle Minimum angle between two surface normals of the rendered object.

#### Phase

- Max Maximum phase.
- Min Minimum phase.

**Origin** Origin point of the interactive scalar cut plane.

Normal Normal vector to the interactive scalar cut plane.

View Axes Optionally view the axes of the array.

# 5.2 Library Reference - Import Tools

# 5.2.1 Array to Memory

Provides a means to load a NumPy array from disk directly into a memory by using either of the 10 keywords memory0, memory1, ..., memory9 as a string in place of a NumPy file path. See *Array Creation and Storage* for furthr details.

### Parameters:

**Input file** User provided path string to the input NumPy file.

Output File User provided choice of memory keyword.

### 5.2.2 Empty Array

Creates an empty new array. If any array is supplied in the (xyz) from array field, the dimensions of this array will be used for the empty array. Otherwise, the specified dimensions are used.

### Parameters:

Output file User provided path string for the output NumPy file.

- (xyz) from array Optional user provided path string for NumPy file used to provide the support array dimensions.
- (xyz) from dimensions {x,y,z} Dimensions of the support array. This is overridden if the (xyz) from array field is used.

# 5.2.3 HDF5 to Numpy

Provides a means to browse and import data within a Hierarchical Data Format (HDF) file. HDF5 is supported via the python H5Py package.

#### Parameters:

**Input HDF file** User provided path string to the input HDF file.

HDF key path Comma separated list of HDF keys found within the input HDF file.

Output File User provided path string for the output NumPy file.

# 5.2.4 Image to Numpy

Provides a means to import image data into NumPy format. Accepted formats include Portable Network Graphics (PNG), JPEG, Portable PixMap (PPM) and Tagged Image File (TIF) format. Output Numpy arrays are three dimensional with unit length in the third dimension.

#### **Parameters**:

**Input Image file** User provided path string to the input image file.

Output File User provided path string for the output NumPy file.

### 5.2.5 Load Co-ordinates

Initialises or replaces the internally stored Co-ordinates data with the supplied array.

### **Parameters**:

**Input file** User provided path string to the input NumPy file.

# 5.2.6 Load PSF

Initialises or replaces the internally stored Point Spread Function data with the supplied array. This is typically used before a *partial coherence* phase retrieval algorithm commences.

### **Parameters:**

**Input file** User provided path string to the input NumPy file.

### 5.2.7 SPE to Numpy

Provides a means to import data from SPE format into NumPy format. During conversion, additional information is provided and appears in the Log.

### Parameters:

Input SPE file User provided path string to the input SPE file.

Output File User provided path string for the output NumPy file.

# 5.3 Library Reference - Export Tools

# 5.3.1 Array to VTK

Convert an input NumPy array into VTK file format. Uses either the amplitude or phase of the array.

#### Parameters:

**Input file** User provided path string to the input NumPy file.

Output File User provided path string for the output VTK file.

**Type** Use either the amplitude or the phase of the complex input file in order to construct a VTK array.

# 5.3.2 Memory to Array

Provides a means to save to disk a NumPy array stored in memory. See Array to Memory for furthr details.

#### **Parameters**:

**Input file** User provided path string to the input NumPy file.

Output File User provided choice of memory keyword.

# 5.3.3 Object to VTK

Convert an input NumPy array and it's corresponding co-ordinates array into VTK file format. Uses either the amplitude or phase of the array.

### **Parameters:**

**Input file** User provided path string to the input NumPy file.

**Co-ord's File** User provided path string to the corresponding co-ordinates NumPy file. Generating coordinates files is achieved using the *Coordinate transformation* operation.

Output File User provided path string for the output VTK file.

Type Use either the amplitude or the phase of the complex input file in order to construct a VTK array.

# 5.4 Library Reference - Functions

### 5.4.1 Auto Centre

Centres the input NumPy array by placing the voxel with the largest amplitude at the centre of the array. The output NumPy array will necessarily increase in dimensions when compared to the input if centring is required.

### **Parameters:**

**Input file** User provided path string to the input NumPy file.

Output File User provided path string for the output NumPy file.

### 5.4.2 Bin

Creates an array in which each voxel results from the grouping and addition of neighbouring voxels of the input file. The dimensions used (in each  $\{x,y,z\}$  direction) for each voxel group is dictated by the Bin dimensions parameter.

#### **Parameters:**

**Input file** User provided path string to the input NumPy file from which bin groups are defined.

Output File User provided path string for the output bin NumPy file.

**Bin dimensions** {x,y,z} Dimensions used for grouping voxels.

# 5.4.3 Blank Line Fill Array

Applies an average filter to the amplitude of non-zero voxels of the input array within the specified region.

### **Parameters**:

**Input file** User provided path string to the input NumPy file.

ROI path Region of interest within the input HDF file.

Output File User provided path string for the output NumPy file.

Filter kernel dimensions Dimensions of the averging kernel.

### 5.4.4 Centred Resize

Centres the input NumPy array by placing the voxel with the largest amplitude at the centre of the array and then resizes the cetered array to the specified dimensions.

#### **Parameters:**

Input file User provided path string to the input NumPy file.

Output File User provided path string for the output NumPy file.

**Dimensions**  $\{x,y,z\}$  Dimensions of the output array.

# 5.4.5 Conjugate Reflect

Takes the complex conjugate and reflects the coordinates of the input NumPy file.

### Parameters:

**Input file** User provided path string to the input NumPy file.

Output File User provided path string for the output NumPy file.

### 5.4.6 Convolve

Performs a discrete convolution of the two input NumPy files.

### Parameters:

First Input file User provided path string to the first input NumPy file.

**Second Input file** User provided path string to the second input NumPy file.

Output File User provided path string for the output NumPy file.

# 5.4.7 Crop Pad

Provides a means to crop and zero pad a NumPy array. The input array is first cropped and subsequently zero padded according to the specified parameters.

#### Parameters:

**Input file** User provided path string to the input NumPy file.

Output File User provided path string for the output NumPy file.

**Crop dimensions: Start:**  $\{i,j,k\}$  Number of indices by which to crop the array along each  $\{i,j,k\}$  direction. Cropping direction begins from the start of the array (and counts forward) for each  $\{i,j,k\}$  direction.

**Crop dimensions: End: {i,j,k}** Number of indices by which to crop the array along each {i,j,k} direction. Cropping direction begins from the end of the array (and counts backward) for each {i,j,k} direction.

**Pad dimensions: Start:**  $\{i,j,k\}$  Number of indices by which to pad the array along each  $\{i,j,k\}$  direction. Padding direction begins from the start of the array for each  $\{i,j,k\}$  direction.

**Pad dimensions:** End: {i,j,k} Number of indices by which to pad the array along each {i,j,k} direction. Padding direction begins from the end of the array for each {i,j,k} direction.

# 5.4.8 Cuboid Support

Creates a *support* array for use in phase retrieval. Voxels in this array take a value of 1 + 0j inside the *support* and zero everywhere else. The support is automatically centred within the array.

If any array is supplied in the (xyz) from array field, the dimensions of this array will be used for the *support* array. Otherwise, the specified dimensions will be used.

#### **Parameters**:

Support file User provided path string for the output support NumPy file.

(xyz) from array Optional user provided path string for NumPy file used to provide the support array dimensions.

(xyz) from dimensions {x,y,z} Dimensions of the support array. This is overridden if the (xyz) from array field is used.

Support size {x,y,z} Dimensions of the support. This must be smaller than the size of the array.

# **5.4.9 Polyhedron Support**

Creates a *support* array for use in phase retrieval using the specified planes to form the surface. The vector formed by each terminal and initial point coordinate pair is the normal vector to a surface that encloses the support. Voxels in this array take a value of 1 + 0j inside the *support* and zero everywhere else.

If any array is supplied in the (xyz) from array field, the dimensions of this array will be used for the *support* array. Otherwise, the specified dimensions will be used.

#### **Parameters:**

Support file User provided path string for the output support NumPy file.

(xyz) from array Optional user provided path string for NumPy file used to provide the support array dimensions.

(xyz) from dimensions {x,y,z} Dimensions of the support array. This is overridden if the (xyz) from array field is used.

**Initial Point Coordinates** List of initial point coordinates of vector normals to each surface. One per line with comma separated ordinates. Parentheses are ignored.

**Terminal Point Coordinates** List of terminal point coordinates of vector normals to each surface. One per line with comma separated ordinates. Parentheses are ignored.

### 5.4.10 Fourier Transform

Performs a Fast Fourier transform (FFT) on the input NumPy file in the specified direction.

#### **Parameters:**

**Input file** User provided path string to the input NumPy file.

Output File User provided path string for the output NumPy file.

To

- Fourier Space Transform to Fourier space.
- Real Space Transform to Real space.

### 5.4.11 Gaussian Fill

Fill the input NumPy file with a Gaussian function.

#### **Parameters**:

**Input file** User provided path string to the input NumPy file.

Output File User provided path string for the output NumPy file.

**Sigma** Standard Deviation of the Gaussian function.

# 5.4.12 Interpolate Object

Interpolates an input NumPy array and it's corresponding co-ordinates onto regular grid NumPy array with dimensions dictated by Array grid size. Shepards famous algorithm is used for the interpolation.

#### **Parameters**:

**Input file** User provided path string to the input NumPy file.

**Co-ord's File** User provided path string to the corresponding co-ordinates NumPy file. Generating coordinates files is achieved using the *Coordinate transformation* operation.

Output File User provided path string for the output NumPy file.

**Array grid size** Dimensions of the output NumPy array.

**Array Bounds: Start:** {x,y,z} Start bounding coordinates. Each value must be smaller than the End bounding coordinates. If all values are zero, they are ignored and the digault bounds are used instead.

**Array Bounds: End:** {x,y,z} End bounding coordinates. Each value must be larger than the Start bounding coordinates. If all values are zero, they are ignored and the digault bounds are used instead.

**Interpolation range** Fraction of the input array used for interpolating each point. Larger values approaching unity will significantly slow down the interpolation process.

### 5.4.13 Affine Transform

Performs an affine (linear) transformation on the input coordinates file.

#### **Parameters**:

**Input co-ord's file** User provided path string to the corresponding co-ordinates NumPy file. Generating coordinates files is achieved using the *Coordinate transformation* operation.

Output co-ord's file User provided path string for the output co-ordinates NumPy file.

**Translate:** {x,y,z} Translation of coordinates in each orthogonal direction. Units are equal to those of the original coordinates.

Scale: {x,y,z} Scaling of each orthogonal axis.

**Rotate:** {x,y,z} Rotation about each orthogonal axis. Units are in degrees.

### 5.4.14 Mask

Creates a mask array for use in phase retrieval. Voxels in this array take a value of 1+0j if the corresponding voxel of the input file is inside the range of Minimum Value to Maximum Value. Voxels outside the same range are set to zero.

#### **Parameters:**

**Input file** User provided path string to the input NumPy file from which Minimum Value and Maximum Value are obtained.

Output File User provided path string for the output mask NumPy file.

**Maximum Value** Values above (equal or below) Maximum Value in the input file will result in a corresponding voxel value of zero (1+0j) in the mask.

**Minimum Value** Values below (equal or above) Minimum Value in the input file will result in a corresponding voxel value of zero (1+0j) in the mask.

### 5.4.15 Median Filter

Applies a median filter to the amplitude of the input array. Voxels are only replaced in the array if they deviate from the median filtered equivalent by an amount greater than the Normal deviation.

### **Parameters**:

**Input file** User provided path string to the input NumPy file.

Output File User provided path string for the output NumPy file.

Filter kernel dimensions Dimensions of the median filtering kernel.

**Normal deviation** Median filter voxel replace cut-off value. This value is compared element-wise to the normalised difference in amplitude of voxels from the input array and from the median filtered equivalent, i.e. the magnitude of (median filtered array - array)/ array. Voxels that exceed this cut-off are replaced with their median-filtered equivalent.

# 5.4.16 Python Script

Execute python script. *Memory arrays* are exposed to this function.

# 5.4.17 Rotate Support

Rotates a binary array (such as a support array) by the specified angle in degrees normal to the specified axis. This operation is only intended for binary arrays.

### Parameters:

**Input file** User provided path string to the input NumPy file.

**Output File** User provided path string for the output NumPy file.

**Axis** Axis normal to the rotation. 1,2 and 3 correspond to axis x,y and z.

Angle Angle of rotation in degrees.

### 5.4.18 Scale Array

Scales the input array by a given amount. Scaling is performed element-wise.

#### **Parameters**:

**Input file** User provided path string to the input NumPy file.

**Output File** User provided path string for the output NumPy file.

**Scale factor** Factor by which each voxel of the array is scaled.

# 5.4.19 Sum or Subtract Array

Takes the sum or difference of two input arrays and places the result in the output array.

### Parameters:

**Input file 1** User provided path string to the input NumPy file.

Input file 2 User provided path string to the input NumPy file.

Output File User provided path string for the output NumPy file.

Add or Subtract Choose whether to add or subtract array 2 from array 1.

### 5.4.20 Threshold Data

Creates an array using data from the input file for which voxels outside the range of Minimum Value to Maximum Value are set to zero.

#### Parameters:

Input file User provided path string to the input NumPy file from which Minimum Value and Maximum Value are obtained.

Output File User provided path string for the output mask NumPy file.

**Maximum Value** Values above Maximum Value in the input file will result in a corresponding voxel value of zero in the output file.

**Minimum Value** Values below Minimum Value in the input file will result in a corresponding voxel value of zero in the output file.

# **5.4.21 Transpose Array**

Transposes an array from dimensions  $\{x,y,z\}$  to dimensions  $\{z,y,x\}$ .

**Parameters**:

**Input file** User provided path string to the input NumPy file.

Output File User provided choice of memory keyword.

### 5.4.22 Voxel Replace

Replaces all voxels with indices in the range of Start dimensions and End dimensions with the specified complex value.

**Parameters**:

**Input file** User provided path string to the input NumPy file.

Output File User provided path string for the output NumPy file.

**Start dimensions:** {i,j,k} Starting indices from which the replacement will begin.

**End dimensions:** {i,j,k} Final indices for which the replacement will end.

**Complex Value** Real and imaginary parts of a complex number used repeatedly to replace voxel data in the input file.

# 5.4.23 Wrap Data

Reorders the data of the input NumPy file into wrap-around order.

Parameters:

**Input file** User provided path string to the input NumPy file.

Output File User provided path string for the output NumPy file.

**Wrap Direction** If an array dimension has an odd number of elements, a Forward followed by a Reverse wrap is required to obtain the original array.

# 5.5 Library Reference – Phasing Algorithms

When coherent diffraction measurements are performed phase information is lost and in general we obtain the reciprocal space intensity distribution  $I(\mathbf{k}) = |\hat{\rho}_0(\mathbf{k})|^2 = |\hat{\rho}_0(\mathbf{k})e^{i\phi(\mathbf{k})}|^2$ , where  $\hat{\rho}(\mathbf{k})$  is the complex density in Fourier-space. We wish to recover the complex real-space density  $\rho(\mathbf{r})$ . This is achieved using iterative reconstruction algorithms that traverse back and forth between direct and Fourier space while applying a constraint at each turn.

Generally, phase retrieval algorithms define a support region in real-space for which the amplitude of the object density is unrestricted. Outside this region the amplitude of the density  $\rho(\mathbf{r})$  is minimised in some way. A Fourier-space constraint is also defined  $(P_{FS})$  and requires the objects amplitude to be proportional in some way to the original measurement on some set  $\mathcal{M}$  such that  $P_{FS}|\hat{\rho}(\mathbf{k})|e^{i\phi(\mathbf{k})}=|\hat{\rho}_0(\mathbf{k})||e^{i\phi(\mathbf{k})}$ , for all  $\mathbf{k}$  in  $\mathcal{M}$ .

It is possible to execute more than one of algorithms in the *phasing pipeline* Algorithms executed in this way will share reconstruction data, hereafter referred to as sequence data. It is also possible, unless otherwise stated, for algorithms to share support data. This is achieved by leaving blank the path string field for the support NumPy file. Such a feature is desirable for algorithms that modify the support array such as the shrink-wrap method.

The following describes the available algorithms for phase retrieval.

### 5.5.1 Random Start

Randomises the phase of the internally stored sequence data. This is typically used before phase retrieval commences. If this is not used, the sequence data is filled with zeros.

#### **Parameters**:

**Amp max** Specifies the amplitude of the array that is randomised.

### 5.5.2 Array Start

Replaces the internally stored sequence data with the supplied array. This is typically used before phase retrieval commences.

#### Parameters:

**Input file** User provided path string to the input NumPy file.

# 5.5.3 Hybrid Input-Output (HIO)

Fienup's hybrid input-output (HIO) algorithm. See here for more details.

### Parameters:

**Exp amp** User provided path string to the input experimental data NumPy file.

**Square Root Exp amp** Causes the Exp amp array to be square rooted before phase retrieval commences. This is necessary if the array contains intensity measurements.

Support User provided path string to the support data NumPy file.

Beta Relaxation parameter for the HIO algorithm.

**Iterations** Total number of iterations for which the algorithm will perform.

### **5.5.4 HIO Mask**

Fienup's hybrid input-output (HIO) algorithm with the addition of a Fourier space constraint *mask*. See here for more details.

### Parameters:

Exp amp User provided path string to the input experimental data NumPy file.

**Square Root Exp amp** Causes the Exp amp array to be square rooted before phase retrieval commences. This is necessary if the array contains intensity measurements.

**Support** User provided path string to the *support* data NumPy file.

Mask User provided path string to the mask data NumPy file.

**Beta** Relaxation parameter for the HIO algorithm.

**Iterations** Total number of iterations for which the algorithm will perform.

### 5.5.5 HIO Plus

Fienup's hybrid input-output (HIO) algorithm with non-negativity constraint and with the addition of a Fourier space constraint *mask*. See here for more details.

#### **Parameters**:

**Exp amp** User provided path string to the input experimental data NumPy file.

**Square Root Exp amp** Causes the Exp amp array to be square rooted before phase retrieval commences. This is necessary if the array contains intensity measurements.

**Support** User provided path string to the *support* data NumPy file.

Mask User provided path string to the mask data NumPy file.

Beta Relaxation parameter for the HIO algorithm.

**Iterations** Total number of iterations for which the algorithm will perform.

# 5.5.6 Phase Constrained HIO (PCHIO)

Fienup's hybrid input-output (HIO) algorithm with phase constraint and with the addition of a Fourier space constraint *mask*. See here for more details.

#### Parameters:

**Exp amp** User provided path string to the input experimental data NumPy file.

**Square Root Exp amp** Causes the Exp amp array to be square rooted before phase retrieval commences. This is necessary if the array contains intensity measurements.

**Support** User provided path string to the *support* data NumPy file.

Mask User provided path string to the mask data NumPy file.

Beta Relaxation parameter for the HIO algorithm.

**Iterations** Total number of iterations for which the algorithm will perform.

Phase Max Voxels with phase greater than Phase max are penalised.

**Phase Min** Voxels with phase lesser than Phase min are penalised.

### **5.5.7 Phase Gradient Constrained HIO (PGCHIO)**

Fienup's hybrid input-output (HIO) algorithm with phase gradient constraint in the Q-vector direction with the addition of a Fourier space constraint *mask*. See here for more details.

Warning: Co-ordinate transformation of the object or Q-vector may be required.

### Parameters:

**Exp amp** User provided path string to the input experimental data NumPy file.

**Square Root Exp amp** Causes the Exp amp array to be square rooted before phase retrieval commences. This is necessary if the array contains intensity measurements.

**Support** User provided path string to the *support* data NumPy file.

Mask User provided path string to the mask data NumPy file.

Beta Relaxation parameter for the HIO algorithm.

**Iterations** Total number of iterations for which the algorithm will perform.

Phase Max Voxels with phase gradient greater than Phase max are penalised, in the direction the Q-vector.

Phase Min Voxels with phase gradient lesser than Phase min are penalised, in the direction the Q-vector.

 $Q\{x,y,z\}$  Q-vector, in real space coordinates. Only the vectors direction is significant.

# 5.5.8 Error Reduction (ER)

Error Reduction (ER) algorithm. See here for more details.

### Parameters:

**Exp amp** User provided path string to the input experimental data NumPy file.

**Square Root Exp amp** Causes the Exp amp array to be square rooted before phase retrieval commences. This is necessary if the array contains intensity measurements.

**Support** User provided path string to the *support* data NumPy file.

**Iterations** Total number of iterations for which the algorithm will perform.

### 5.5.9 ER Mask

Error Reduction (ER) algorithm with the addition of a Fourier space constraint *mask*. See here for more details.

#### **Parameters**

**Exp amp** User provided path string to the input experimental data NumPy file.

**Square Root Exp amp** Causes the Exp amp array to be square rooted before phase retrieval commences. This is necessary if the array contains intensity measurements.

**Support** User provided path string to the *support* data NumPy file.

**Mask** User provided path string to the mask data NumPy file.

**Iterations** Total number of iterations for which the algorithm will perform.

## 5.5.10 Phase Only ER (POER)

Phase Only Error Reduction (POER) algorithm with the addition of a Fourier space constraint *mask*. See here for more details.

### Parameters:

**Exp amp** User provided path string to the input experimental data NumPy file.

**Square Root Exp amp** Causes the Exp amp array to be square rooted before phase retrieval commences. This is necessary if the array contains intensity measurements.

Support User provided path string to the *support* data NumPy file.

Mask User provided path string to the mask data NumPy file.

**Iterations** Total number of iterations for which the algorithm will perform.

# 5.5.11 Relaxed Average Alternating Reflection (RAAR)

Relaxed Average Alternating Reflection (RAAR) algorithm. See here for more details.

#### **Parameters:**

**Exp amp** User provided path string to the input experimental data NumPy file.

**Square Root Exp amp** Causes the Exp amp array to be square rooted before phase retrieval commences. This is necessary if the array contains intensity measurements.

**Support** User provided path string to the *support* data NumPy file.

Mask User provided path string to the mask data NumPy file.

Beta Relaxation parameter for the RAAR algorithm.

**Iterations** Total number of iterations for which the algorithm will perform.

# 5.5.12 Hybrid Projection Reflection (HPR)

Hybrid Projection Reflection (HPR) algorithm. See here for more details.

#### **Parameters:**

**Exp amp** User provided path string to the input experimental data NumPy file.

**Square Root Exp amp** Causes the Exp amp array to be square rooted before phase retrieval commences. This is necessary if the array contains intensity measurements.

Support User provided path string to the support data NumPy file.

Mask User provided path string to the mask data NumPy file.

**Beta** Relaxation parameter for the HPR algorithm.

**Iterations** Total number of iterations for which the algorithm will perform.

# 5.5.13 Compressed Sensing HIO (CSHIO)

Compressed Sensing HIO (CSHIO) algorithm. See here for more details.

### **Parameters**:

Exp amp User provided path string to the input experimental data NumPy file.

**Square Root Exp amp** Causes the Exp amp array to be square rooted before phase retrieval commences. This is necessary if the array contains intensity measurements.

Support User provided path string to the support data NumPy file.

Mask User provided path string to the mask data NumPy file.

**Beta** Relaxation parameter for the HIO algorithm.

**Iterations** Total number of iterations for which the algorithm will perform.

**p-norm** p-norm of the Lebesgue space.

**Epsilon** Positive relaxation parameter  $(\epsilon)$  for the weighted p-norm.

**Epsilon min** Minimum value that epsilon can take.

**Divisor** Amount by which epsilon is divided when the following condition is met:  $\|\rho^{(n)}(\mathbf{r})\|_2 - \|\rho^{(n-1)}(\mathbf{r})\|_2\| < \frac{\sqrt{\epsilon}}{n}$ 

**eta** Parameter  $(\eta)$  in the divisor condition above.

# 5.5.14 HIO Mask with Partial Coherence Optimisation (HIO Mask PC)

HIO Mask with Partial Coherence Optimisation (HIO Mask PC) algorithm. See here for more details.

#### **Parameters:**

**Exp amp** User provided path string to the input experimental data NumPy file.

**Square Root Exp amp** Causes the Exp amp array to be square rooted before phase retrieval commences. This is necessary if the array contains intensity measurements.

**Support** User provided path string to the *support* data NumPy file.

**Mask** User provided path string to the mask data NumPy file.

Beta Relaxation parameter for the HIO algorithm.

**Iterations** Total number of iterations for which the algorithm will perform (not including R-L iterations).

**Iterations preceding R-L optimisation** Interval of iterations after which Richardson-Lucy optimisation commences in successive cycles.

**R-L iterations** Number of R-L iterations performed at each cycle.

Interval between R-L optimisation Number of iterations between each Richardson-Lucy optimisation cycle.

**Initial PSF HWHM** The initial point spread function (PSF) optimised using the R-L algorithm is a normal Lorentzian function  $(\frac{1}{\pi} \frac{a}{x^2 + a^2})$ . The user can specify the HWHM of its Fourier transform  $(e^{-a|x|})$ .

**Zero fill end dimensions of PSF** After optimising the PSF with the R-L algorithm, voxels upto a distance {i,j,k} from the perimeter of the PSF array are set to zero. This can improve stability of the algorithm.

**Reset PSF** Resets the PSF to a normal Lorentzian function before the next R-L optimisation cycle.

# 5.5.15 ER Mask with Partial Coherence Optimisation (ER Mask PC)

ER Mask with Partial Coherence Optimisation (ER Mask PC) algorithm. See here for more details.

### Parameters:

**Exp amp** User provided path string to the input experimental data NumPy file.

**Square Root Exp amp** Causes the Exp amp array to be square rooted before phase retrieval commences. This is necessary if the array contains intensity measurements.

**Support** User provided path string to the *support* data NumPy file.

Mask User provided path string to the mask data NumPy file.

Beta Relaxation parameter for the ER algorithm.

Iterations Total number of iterations for which the algorithm will perform (not including R-L iterations).

**Iterations preceding R-L optimisation** Interval of iterations after which Richardson-Lucy optimisation commences in successive cycles.

**R-L iterations** Number of R-L iterations performed at each cycle.

Interval between R-L optimisation Number of iterations between each Richardson-Lucy optimisation cycle.

**Initial PSF HWHM** The initial point spread function (PSF) optimised using the R-L algorithm is a normal Lorentzian function  $(\frac{1}{\pi} \frac{a}{x^2 + a^2})$ . The user can specify the HWHM of its Fourier transform  $(e^{-a|x|})$ .

**Zero fill end dimensions of PSF** After optimising the PSF with the R-L algorithm, voxels upto a distance {i,j,k} from the perimeter of the PSF array are set to zero. This can improve stability of the algorithm.

**Reset PSF** Resets the PSF to a normal Lorentzian function before the next R-L optimisation cycle.

# 5.5.16 HPR Mask with Partial Coherence Optimisation (HPR Mask PC)

HPR Mask with Partial Coherence Optimisation (HPR Mask PC) algorithm. See here for more details.

#### **Parameters:**

**Exp amp** User provided path string to the input experimental data NumPy file.

**Square Root Exp amp** Causes the Exp amp array to be square rooted before phase retrieval commences. This is necessary if the array contains intensity measurements.

**Support** User provided path string to the *support* data NumPy file.

**Mask** User provided path string to the mask data NumPy file.

Beta Relaxation parameter for the HPR algorithm.

**Iterations** Total number of iterations for which the algorithm will perform (not including R-L iterations).

**Iterations preceding R-L optimisation** Interval of iterations after which Richardson-Lucy optimisation commences in successive cycles.

**R-L iterations** Number of R-L iterations performed at each cycle.

**Interval between R-L optimisation** Number of iterations between each Richardson-Lucy optimisation cycle.

**Initial PSF HWHM** The initial point spread function (PSF) optimised using the R-L algorithm is a normal Lorentzian function  $(\frac{1}{\pi} \frac{a}{x^2 + a^2})$ . The user can specify the HWHM of its Fourier transform  $(e^{-a|x|})$ .

**Zero fill end dimensions of PSF** After optimising the PSF with the R-L algorithm, voxels upto a distance {i,j,k} from the perimeter of the PSF array are set to zero. This can improve stability of the algorithm.

**Reset PSF** Resets the PSF to a normal Lorentzian function before the next R-L optimisation cycle.

# 5.5.17 RAAR Mask with Partial Coherence Optimisation (RAAR Mask PC)

RAAR Mask with Partial Coherence Optimisation (RAAR Mask PC) algorithm. See here for more details.

# Parameters:

**Exp amp** User provided path string to the input experimental data NumPy file.

**Square Root Exp amp** Causes the Exp amp array to be square rooted before phase retrieval commences. This is necessary if the array contains intensity measurements.

**Support** User provided path string to the *support* data NumPy file.

Mask User provided path string to the mask data NumPy file.

Beta Relaxation parameter for the RAAR algorithm.

Iterations Total number of iterations for which the algorithm will perform (not including R-L iterations).

**Iterations preceding R-L optimisation** Interval of iterations after which Richardson-Lucy optimisation commences in successive cycles.

**R-L iterations** Number of R-L iterations performed at each cycle.

Interval between R-L optimisation Number of iterations between each Richardson-Lucy optimisation cycle.

**Initial PSF HWHM** The initial point spread function (PSF) optimised using the R-L algorithm is a normal Lorentzian function  $(\frac{1}{\pi} \frac{a}{x^2 + a^2})$ . The user can specify the HWHM of its Fourier transform  $(e^{-a|x|})$ .

**Zero fill end dimensions of PSF** After optimising the PSF with the R-L algorithm, voxels upto a distance  $\{i,j,k\}$  from the perimeter of the PSF array are set to zero. This can improve stability of the algorithm.

**Reset PSF** Resets the PSF to a normal Lorentzian function before the next R-L optimisation cycle.

# 5.5.18 Reweighted 2D Saddle Point Optimisation (SO2D)

Reweighted 2D Saddle Point Optimisation (SO2D) algorithm. See here for more details.

#### **Parameters:**

**Exp amp** User provided path string to the input experimental data NumPy file.

**Square Root Exp amp** Causes the Exp amp array to be square rooted before phase retrieval commences. This is necessary if the array contains intensity measurements.

**Support** User provided path string to the *support* data NumPy file.

Mask User provided path string to the mask data NumPy file.

Iterations Total number of iterations for which the algorithm will perform.

Step optimisation iterations Total number of iterations performed when optimising the step length.

**Max step size** Maximum size for the total step length  $(\sqrt{\alpha^2 + \beta^2})$ .

**Initial Beta** Relaxation parameter for the initial step  $(\beta)$ .

Max step increment Maximum amount by which the step can be incremented.

**Min step increment** Minimum amount by which the step can be incremented.

**Exit Ratio** Value below  $|\psi|/|\psi_0|$  will halt step length optimisation.

**Reset Ratio** Value above  $|\psi|/|\psi_0|$  will reset step length to that of the initial value (initial  $\beta$ ).

**Exit Error** Value below  $(\psi^{(n+1)} - \psi^{(n)})/\psi^{(n)}$  will halt step length optimisation.

# 5.5.19 Shrink Wrap

Shrink Wrap algorithm. See here for more details.

### **Parameters**:

**Exp amp** User provided path string to the input experimental data NumPy file.

**Square Root Exp amp** Causes the Exp amp array to be square rooted before phase retrieval commences. This is necessary if the array contains intensity measurements.

Support User provided path string to the *support* data NumPy file.

Mask User provided path string to the mask data NumPy file.

**Beta** Relaxation parameter for the HIO algorithm.

**Iterations** Total number of iterations for which the algorithm will perform.

Cycle length Interval of iterations after which the support is updated.

Sigma Standard deviation of the Gaussian smoothing function for the support.

**Threshold** Fractional value below which sequence data is not used when creating the new support.

Algorithm Algorithm to use when performing the shrink wrapped phase retrieval.

# 5.6 Library Reference - Phasing Operations

### 5.6.1 Save Sequence

Save sequence data to a NumPy file.

**Parameters**:

**Output file** User provided path string for the output NumPy file.

### 5.6.2 Save Support

Save Support data to a NumPy file.

**Parameters:** 

Output file User provided path string for the output NumPy file.

### 5.6.3 Save Residual

Save Residual data to a NumPy file.

**Parameters:** 

**Output file** User provided path string for the output CSV file.

### 5.6.4 Save PSF

Saves the internally stored Point Spread Function data to a NumPy file.

**Parameters:** 

Output file User provided path string for the ouput NumPy file.

### 5.6.5 Co-ordinate Transformation

Perform a co-ordinate transformation on either sequence data or an input array. The transformation is derived using the principles as demonstrated by Pfeifer, M.A. . The generated co-ordinate array is stored internally and is accessible for *viewing* with the *memory array* keyword memorycoords.

### **Parameters**:

Transform from Source from which to perform the transformation against.

**Transform type** Transform in real or reciprocal space.

**Input data** User provided path string to the input NumPy file.

Output amp file User provided path string for the output amplitude VTK file.

Output phase file User provided path string for the output phase VTK file.

**Rocking curve type** Theta  $(\theta)$  or Phi  $(\phi)$  rocking curve measurement.

Array binning Amount by which the array was binned. See the section on binning for further details.

**2theta**  $2\theta$  angle.

**d theta** Increment angle for  $\theta$  rocking curve measurement.

**phi**  $\phi$  angle.

**d phi** Increment angle for  $\phi$  rocking curve measurement.

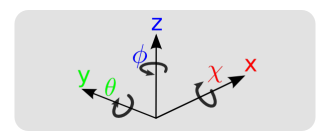
Pixel  $\{x,y\}$  (microns) Detector pixel dimensions, in microns.

wavelength (nm) Wavelength of light used in experiment. Units of nanometres.

**Arm length (m)** Distance from the sample to the centre of diffraction pattern at the detector. Units of metres.

CCD x-axis flip Reverse the direction of detector x-axis readout. Required for some PI CCD detectors.

**Experimental geometry:** 



Curved arrows show the direction of rotation of object (as opposed to the axis). The beam is assumed to travel along the positive x direction.

### 5.6.6 Save Co-ordinates

Save Co-ordinates data to a NumPy file.

**Output file** User provided path string for the output NumPy file.

**Note:** A co-ordinate transformation must occur prior to this operation.

## **SCRIPTING WITH BONSU**

## 6.1 Core concepts

Bonsu provides an object-oriented scripting interface to each algorithm for ease and interoperability with additional Python modules. To begin using this interface, import the phasing module as follows:

```
>>> from bonsu import phasing
```

Once imported, the phasing module provides access to all phase retrieval algorithms. For a complete list see *Library Reference – Phasing Algorithms*.

Each algorithm will take a number of NumPy arrays as input. Input data is loaded and prepared in wrap around order. It is therefore also necessary to utilise the *Wrap Data* function. Note that the data must be stored in a "C"-ordered double precision complex Numpy array:

The data mask in wrap around order is also needed:

```
>>> mask = numpy.load("mask.npy")
>>> mask[:] = phasing.WrapArray(mask)
```

An array to contain the reconstructed object is also needed and can be created in a number of ways, with data initialised as needed, i.e. random data or initial starting data from a previous reconstruction.

```
>>> seqdata = numpy.empty_like(expdata)
>>> seqdata[:] = 1.0 # initialise as 1.0.
```

To utilise the *HIO Mask* algorithm, create an instance as follows:

```
>>> hio = phasing.HIOMask()
```

Then initialise the remaining parameters for 100 iterations:

```
>>> hio.SetStartiter(0)
>>> hio.SetNumiter(100)
>>> hio.SetNumthreads(3)
>>> hio.SetBeta(0.9)
>>> hio.SetExpdata(expdata)
>>> hio.SetSupport(support)
>>> hio.SetMask(mask)
>>> hio.SetSeqdata(seqdata)
```

To begin reconstruction we first prepare the algorithm as follows:

```
>>> hio.Prepare()
```

Execution then follows:

```
>>> hio.Start()
```

In a threaded or non-blocking use of the algorithm, it is possible the pause, resume and stop the reconstruction using the following methods:

```
>>> hio.Pause()
>>> hio.Resume()
>>> hio.Stop()
```

The output of the reconstruction is saved as follows:

```
>>> numpy.save("output.npy", seqdata)
```

Each algorithm can use Shrink Wrap to modify the support during reconstruction. Prefix the algorithm name with SW to utilise the Shrink Wrap version of the algorithm. For example, to use the Shrink Wrap version of the *HIO Mask* algorithm, import and instantiate the following:

```
>>> swhio = phasing.SWHIOMask()
```

Then initialise the remaining parameters as follows:

```
>>> swhio.SetStartiter(0)
>>> swhio.SetNumiter(100)
>>> swhio.SetNumthreads(3)
>>> swhio.SetExpdata(expdata)
>>> swhio.SetSupport(support)
>>> swhio.SetMask(mask)
>>> swhio.SetSeqdata(seqdata)
>>> swhio.Prepare()
```

Methods unque to the Shrink Wrap implementation that require initialisation are as follows:

```
>>> swhio.SetSWCyclelength(20)
>>> swhio.SetSWThreshold(0.2)
>>> swhio.SetSWSigma(3.0)
```

To begin reconstruction we also first prepare the Shrink Wrap algorithm as follows:

```
>>> swhio.SWPrepare()
```

Execution then follows:

```
>>> swhio.SWStart()
```

## 6.2 Custom Algorithms

Bonsu provides tools to build custom projection algorithms. To build a custom algorithm, create a script that imports an abstract algorithm class and subclasses the DoIter and RSCons methods. DoIter is called at each iteration of the algorithm. It will apply the real-space and Fourier-space constraints. RSCons is a method called by DoIter to apply the real-space constraint.

```
from bonsu.phasing.abstract import PhaseAbstract
class MyAlgorithm(PhaseAbstract):
        My custom HIO algorithm
        def DoIter(self):
                self.rho_m1[:] = self.seqdata[:]
                fftw_stride(self.seqdata,self.seqdata,self.plan,FFTW_TORECIP,1)
                self.SetRes()
                self.SetAmplitudes()
                fftw_stride(self.seqdata,self.seqdata,self.plan,FFTW_TOREAL,1)
                n = self.citer_flow[0]
                self.residual[n] = self.res/self.sos
                amp = numpy.absolute(self.seqdata)
                sos1 = numpy.sum(amp*amp)
                self.RSCons()
                amp = numpy.absolute(self.segdata)
                sos2 = numpy.sum(amp*amp)
                norm = sqrt(sos1/sos2)
                self.seqdata[:] = norm*self.seqdata[:]
                self.citer_flow[0] += 1
        def RSCons(self):
                realsupport = numpy.real(self.support)
                nosupport = realsupport < 0.5</pre>
                self.seqdata[nosupport] = self.rho_m1[nosupport] - self.
⇔seqdata[nosupport] * self.beta
```

The custom algorithm is then instantiated as before:

```
myalg = MyAlgorithm()
```

Initialisation of parameters proceeds as before. To begin reconstruction prepare the algorithm as follows:

```
myalg.PrepareCustom()
```

Execution then follows as before:

```
myalg.Start()
```

## 6.3 Concurrent Phase Retrieval

Bonsu provides tools to reconstruct atomic displacement field information (and subsequently strain field information) via concurrent phase reconstruction using diffraction pattern data that is obtained from multiple Bragg reflections of a single crystal. The method used is described in detail in the following publication: Newton, Phys. Rev. B 102, 014104 (2020).

The concurrent phase retrieval method can in principle utilise any projection algorithm implemented using the *Custom Algorithm interface*. The following example illustrates how to use utilise the method using the standard Abstract Algorithm class (HIO). Please note that diffraction pattern data should undergo *co-ordinate transformation* and *interpolation* onto a regular grid prior to commencement of the reconstruction process. Please also note that no checking of the data type (numpy.cdouble) and consistency in the array dimensions is performed.

```
from bonsu.phasing.concurrent import PhaseConcurrent
from bonsu.phasing.abstract import PhaseAbstract
from bonsu.phasing.wrap import WrapArray
# Specify number of diffraction patterns / Q-vectors
NQ = 4
# Instantiate class object
d = PhaseConcurrent()
# Specify class object of phase retrieval algorithm used for reconstruction
# Define your own and use here, if preferred.
d.SetBaseClass(PhaseAbstract)
# Set number of concurrent datasets / Q-vectors
d.SetNQ(NQ)
# Set Q-vectors
d.SetQ(0, [1,0,0])
d.SetQ(1, [1,1,0])
d.SetQ(2, [0,1,0])
d.SetQ(3, [0,0,1])
# Define names of files
expnames = ["diffamp0.npy","diffamp1.npy","diffamp2.npy","diffamp3.npy"]
masknames = ["mask0.npy","mask1.npy","mask2.npy","mask3.npy"]
supportname = "support.npy"
# Use names above to load arrays into list of length NQ
amps = [WrapArray(numpy.load(name)) for name in expnames]
masks = [WrapArray(numpy.load(name)) for name in masknames]
supports = [numpy.load(supportname)]*NQ
seqdatas = [numpy.ones_like(amp) for amp in amps]
# Class methods below are the same as those in the phase retrieval algorithm
# class but will instead take a list of length NQ.
d.SetExpdata(amps)
d.SetSupport(supports)
d.SetMask(masks)
d.SetSeqdata(seqdatas)
d.SetBeta([0.9]*NQ)
# Set iterations
d.SetStartiter(0)
d.SetNumiter(100)
# Number of FFTW threads for each concurrent reconstruction.
```

(continues on next page)

(continued from previous page)

```
d.SetNumthreads(1)

# Prepare
d.Prepare()
# Start
d.Start()

# Save reconstructed phase for each Q-vector
# Filenames are automatically indexed.
d.SaveSequences("output.npy")

# Save displacement field
d.SaveDField("dfield.npy")
```

## 6.4 Class Reference

## 6.4.1 Abstract Classes

```
class bonsu.phasing.abstract.PhaseAbstract(parent=None)
     Bases: object
     Phasing Base Class
     Prepare()
          Prepare algorithm.
     PrepareCustom()
          Prepare algorithm using FFTW python interface.
     SetSOS()
          Set Sum of Squares.
     GetSOS()
          Get Sum of Squares.
     SetStartiter(startiter)
          Set the starting iteration number.
     SetNumiter(numiter)
          Set the number of iterations of the algorithm to perform.
     SetNumthreads(nthreads)
          Set the number of FFTW threads.
     SetSeqdata(seqdata)
          Set the reconstruction data array.
     GetSeqdata()
          Get the reconstruction data array.
     SetExpdata(expdata)
          Set the raw experimental amplitude data array.
     GetExpdata()
          Get the raw experimental amplitude data array.
     SetSupport(support)
          Set the support array.
     GetSupport()
```

Get the support array.

#### SetMask(mask)

Set the mask data array.

#### GetMask()

Set the mask data array.

#### SetBeta(beta)

Set beta relaxation parameter.

#### GetBeta()

Get beta relaxation parameter.

#### Pause()

Pause the reconstruction process.

#### Resume()

Resume the reconstruction process.

#### Start()

Start the reconstruction process.

#### Stop()

Stop the reconstruction process.

## class bonsu.phasing.abstract.PhaseAbstractPC(parent=None)

Bases: bonsu.phasing.abstract.PhaseAbstract

Phasing Partial Coherence Base Class

#### SetPSF(psf)

Set point-spread function from data array.

#### GetPSF()

Get point-spread function from data array.

#### LorentzFillPSF()

Create a point-spread function with Lorentz distribution. This will use the HWHM specified using SetGammaHWHM.

#### SetNumiterRLpre(niterrlpre)

Set the number of iterations before RL optimisation.

#### GetNumiterRLpre()

Get the number of iterations before RL optimisation.

#### SetNumiterRL(niterrl)

Set the number of RL iterations.

#### GetNumiterRL()

Get the number of RL iterations.

#### SetNumiterRLinterval(niterrlinterval)

Set number of iterations between each RL optimisation cycle.

## GetNumiterRLinterval()

Get number of iterations between each RL optimisation cycle.

## **SetGammaHWHM**(gammaHWHM)

Set the half-width half-maximum of the Lorentz distribution. This is utilised in the LorentzFillPSF method.

#### GetGammaHWHM()

Get the half-width half-maximum of the Lorentz distribution.

#### SetPSFZeroEnd(ze)

Voxels upto a distance of [i,j,k] from the perimeter of the PSF array are set to zero. This can improve stability of the algorithm.

#### GetPSFZeroEnd()

Get zero voxels perimeter size.

#### Prepare()

Prepare algorithm.

#### class bonsu.phasing.ShrinkWrap.ShrinkWrap

Shrink Wrap algorithm.

#### SetSWCyclelength(cycle)

Set interval of iterations after which the support is updated.

#### GetSWCyclelength()

Get interval of iterations after which the support is updated.

#### SetSWSigma(sigma)

Set standard deviation of the Gaussian smoothing function for the support.

#### GetSWSigma()

Get standard deviation of the Gaussian smoothing function for the support.

#### SetSWThreshold(thresh)

Set fractional value below which sequence data is not used when creating the new support.

#### GetSWThreshold()

Get fractional value below which sequence data is not used when creating the new support.

#### SWPrepare()

Prepare shrink wrap algorithm.

#### SWStart()

Start the shrink wrap reconstruction process.

## 6.4.2 Hybrid Input-Output Classes

## class bonsu.phasing.HIO.HIO(parent=None)

Bases: bonsu.phasing.abstract.PhaseAbstract

Fienup's hybrid input-output (HIO) algorithm.

#### class bonsu.phasing.HIO.SWHIO

Bases: bonsu.phasing.HIO.HIO, bonsu.phasing.ShrinkWrap.ShrinkWrap

Fienup's hybrid input-output (HIO) algorithm. Shrink wrapped.

## class bonsu.phasing.HIO.HIOMask(parent=None)

Bases: bonsu.phasing.abstract.PhaseAbstract

Fienup's hybrid input-output (HIO) algorithm with the addition of a Fourier space constraint mask.

#### class bonsu.phasing.HIO.SWHIOMask

Bases: bonsu.phasing.HIO.HIOMask, bonsu.phasing.ShrinkWrap.ShrinkWrap

Fienup's hybrid input-output (HIO) algorithm with the addition of a Fourier space constraint mask. Shrink wrapped.

## class bonsu.phasing.HIO.HIOPlus(parent=None)

Bases: bonsu.phasing.abstract.PhaseAbstract

Fienup's hybrid input-output (HIO) algorithm with non-negativity constraint and with the addition of a Fourier space constraint mask.

#### class bonsu.phasing.HIO.SWHIOPlus

Bases: bonsu.phasing.HIO.HIOPlus, bonsu.phasing.ShrinkWrap.ShrinkWrap

Fienup's hybrid input-output (HIO) algorithm with non-negativity constraint and with the addition of a Fourier space constraint mask. Shrink wrapped.

#### class bonsu.phasing.HIO.PCHIO(parent=None)

Bases: bonsu.phasing.abstract.PhaseAbstract

Fienup's hybrid input-output (HIO) algorithm with phase constraint and with the addition of a Fourier space constraint mask.

#### SetMaxphase(max)

Set phase maximum.

#### GetMaxphase()

Get phase maximum.

#### SetMinphase(min)

Set phase minimum.

#### GetMinphase()

Get phase minimum.

#### class bonsu.phasing.HIO.SWPCHIO

Bases: bonsu.phasing.HIO.PCHIO, bonsu.phasing.ShrinkWrap.ShrinkWrap

Fienup's hybrid input-output (HIO) algorithm with phase constraint and with the addition of a Fourier space constraint mask. Shrink wrapped.

## class bonsu.phasing.HIO.PGCHIO(parent=None)

Bases: bonsu.phasing.abstract.PhaseAbstract

Fienup's hybrid input-output (HIO) algorithm with phase gradient constraint in the Q-vector direction with the addition of a Fourier space constraint mask.

#### SetMaxphase(max)

Set phase maximum.

#### GetMaxphase()

Get phase maximum.

#### SetMinphase(min)

Set phase minimum.

#### GetMinphase()

Get phase minimum.

#### SetQ(q)

Set Q-vector tuple

#### GetQ()

Get Q-vector tuple

#### Prepare()

Prepare algorithm.

## ${\bf class}\ {\tt bonsu.phasing.HIO.SWPGCHIO}$

Bases: bonsu.phasing.HIO.PGCHIO, bonsu.phasing.ShrinkWrap.ShrinkWrap

Fienup's hybrid input-output (HIO) algorithm with phase gradient constraint in the Q-vector direction with the addition of a Fourier space constraint mask. Shrink wrapped.

## class bonsu.phasing.HIO.HIOMaskPC(parent=None)

Bases: bonsu.phasing.abstract.PhaseAbstractPC

HIO Mask with Partial Coherence Optimisation.

#### class bonsu.phasing.HIO.SWHIOMaskPC

Bases: bonsu.phasing.HIO.HIOMaskPC, bonsu.phasing.ShrinkWrap.ShrinkWrap

HIO Mask with Partial Coherence Optimisation. Shrink wrapped.

#### Start()

Start the reconstruction process.

#### 6.4.3 Error Reduction Classes

#### class bonsu.phasing.ER.ER(parent=None)

Bases: bonsu.phasing.abstract.PhaseAbstract

Error Reduction (ER) algorithm.

#### class bonsu.phasing.ER.SWER

Bases: bonsu.phasing.ER.ER, bonsu.phasing.ShrinkWrap.ShrinkWrap

Error Reduction (ER) algorithm. Shrink wrapped.

#### class bonsu.phasing.ER.ERMask(parent=None)

Bases: bonsu.phasing.abstract.PhaseAbstract

Error Reduction (ER) algorithm with the addition of a Fourier space constraint mask.

#### class bonsu.phasing.ER.SWERMask

Bases: bonsu.phasing.ER.ERMask, bonsu.phasing.ShrinkWrap.ShrinkWrap

Error Reduction (ER) algorithm with the addition of a Fourier space constraint mask. Shrink wrapped.

#### class bonsu.phasing.ER.POER(parent=None)

Bases: bonsu.phasing.abstract.PhaseAbstract

Phase Only Error Reduction (POER) algorithm with the addition of a Fourier space constraint mask.

#### class bonsu.phasing.ER.ERMaskPC(parent=None)

Bases: bonsu.phasing.abstract.PhaseAbstractPC

ER Mask with Partial Coherence Optimisation algorithm.

#### class bonsu.phasing.ER.SWERMaskPC

Bases: bonsu.phasing.ER.ERMaskPC, bonsu.phasing.ShrinkWrap.ShrinkWrap

ER Mask with Partial Coherence Optimisation algorithm. Shrink wrapped.

#### Start()

Start the reconstruction process.

### 6.4.4 Hybrid Project-Reflection Classes

#### class bonsu.phasing.HPR.HPR(parent=None)

Bases: bonsu.phasing.abstract.PhaseAbstract

Hybrid Projection Reflection (HPR) algorithm.

#### class bonsu.phasing.HPR.SWHPR

Bases: bonsu.phasing.HPR.HPR, bonsu.phasing.ShrinkWrap.ShrinkWrap

Hybrid Projection Reflection (HPR) algorithm. Shrink wrapped.

## class bonsu.phasing.HPR.HPRPC(parent=None)

Bases: bonsu.phasing.abstract.PhaseAbstractPC

HPR Mask with Partial Coherence Optimisation algorithm.

#### class bonsu.phasing.HPR.SWHPRPC

Bases: bonsu.phasing.HPR.HPRPC, bonsu.phasing.ShrinkWrap.ShrinkWrap

HPR Mask with Partial Coherence Optimisation algorithm. Shrink wrapped.

#### Start()

Start the reconstruction process.

## 6.4.5 Relaxed Average Alternating Reflection Classes

## class bonsu.phasing.RAAR.RAAR(parent=None)

Bases: bonsu.phasing.abstract.PhaseAbstract

Relaxed Average Alternating Reflection (RAAR) algorithm.

#### class bonsu.phasing.RAAR.SWRAAR

Bases: bonsu.phasing.RAAR.RAAR, bonsu.phasing.ShrinkWrap.ShrinkWrap

Relaxed Average Alternating Reflection (RAAR) algorithm. Shrink wrapped.

#### class bonsu.phasing.RAAR.RAARPC(parent=None)

Bases: bonsu.phasing.abstract.PhaseAbstractPC

RAAR Mask with Partial Coherence Optimisation algorithm.

#### class bonsu.phasing.RAAR.SWRAARPC

Bases: bonsu.phasing.RAAR.RAARPC, bonsu.phasing.ShrinkWrap.ShrinkWrap

RAAR Mask with Partial Coherence Optimisation algorithm. Shrink wrapped.

#### Start()

Start the reconstruction process.

### 6.4.6 CSHIO Classes

## class bonsu.phasing.CSHIO.CSHIO(parent=None)

Bases: bonsu.phasing.abstract.PhaseAbstract

Compressed Sensing HIO (CSHIO) algorithm.

#### SetPnorm(p)

Set p-norm of the Lebesgue space.

#### GetPnorm()

Get p-norm of the Lebesgue space.

## SetEpsilon(epsilon)

Set positive relaxation parameter (epsilon) for the weighted p-norm.

#### GetEpsilon()

Get positive relaxation parameter (epsilon) for the weighted p-norm.

### SetEpsilonmin(epsilonmin)

Set minimum value that epsilon can take.

#### GetEpsilonmin()

Set minimum value that epsilon can take.

#### SetDivisor(d)

Set amount by which epsilon is divided when constraint is met.

#### GetDivisor()

Get amount by which epsilon is divided when constraint is met.

#### SetEta(eta)

Set parameter in the divisor condition.

#### GetEta()

Get parameter in the divisor condition.

## Prepare()

Prepare algorithm.

#### class bonsu.phasing.CSHIO.SWCSHIO

Bases: bonsu.phasing.CSHIO.CSHIO, bonsu.phasing.ShrinkWrap.ShrinkWrap

Compressed Sensing HIO (CSHIO) algorithm. Shrink wrapped.

#### 6.4.7 SO2D Classes

#### class bonsu.phasing.SO2D.SO2D(parent=None)

Bases: bonsu.phasing.abstract.PhaseAbstract

#### SetNumsoiter(numsoiter)

Set total number of iterations performed when optimising the step length.

#### GetNumsoiter()

Get total number of iterations performed when optimising the step length.

#### SetReweightiter(reweightiter)

Set iteration after which reweighting is applied. A negative value implies no reweighting.

#### GetReweightiter()

Get iteration after which reweighting is applied. A negative value implies no reweighting.

#### **SetDtaumax**(*dtaumax*)

Set maximum amount by which the step can be incremented.

#### GetDtaumax()

Get maximum amount by which the step can be incremented.

#### SetDtaumin(dtaumin)

Set minimum amount by which the step can be incremented.

#### GetDtaumin()

Get minimum amount by which the step can be incremented.

#### **SetTaumax**(taumax)

Set maximum size for the total step length.

#### GetTaumax()

Get maximum size for the total step length.

#### **SetPsiexitratio**(psiexitratio)

Set value below |psi|/|psi\_0| that will halt step length optimisation.

## GetPsiexitratio()

Get value below |psi|/|psi\_0| that will halt step length optimisation.

#### SetPsiexiterror(psiexiterror)

Set value below  $(psi^{(n+1)} - psi^{(n)})/psi^{(n)}$  that will halt step length optimisation.

#### GetPsiexiterror()

Get value below (psi^(n+1) - psi^(n))/psi^(n) that will halt step length optimisation.

## SetPsiresetratio(psiresetratio)

Set value above <a href="mailto:psi\_0">[psi\_0]</a> that will reset the step length to that of the initial value.

#### GetPsiresetratio()

Get value above |psi|/|psi\_0| that will reset the step length to that of the initial value.

#### Prepare()

Prepare algorithm.

### class bonsu.phasing.SO2D.SWSO2D

 $Bases:\ bonsu.phasing.SO2D.SO2D,\ bonsu.phasing.Shrink \verb|Wrap.Shrink|| Wrap.Shrink Wrap.Shri$ 

Bonsu - The Interactive Phase Retrieval Suite Documentation, Release 3.4.2			

## **CHAPTER**

# **SEVEN**

# **INDICES AND TABLES**

- genindex
- search

Bonsu - The Interactive Phase Retrieval Suite Documentation, Release 3.4.2	
· · · · · · · · · · · · · · · · · · ·	_

## **PYTHON MODULE INDEX**

## b

```
bonsu.phasing.abstract, 37
bonsu.phasing.CSHIO, 42
bonsu.phasing.ER, 41
bonsu.phasing.HIO, 39
bonsu.phasing.HPR, 41
bonsu.phasing.RAAR, 42
bonsu.phasing.ShrinkWrap, 39
bonsu.phasing.SO2D, 43
```

# **INDEX**

A	С		
Affine Transform, 21	Centred Resize, 18		
Axis scaling, 21	Input file, 18		
Input co-ord's file, 21	Output File, 18		
Output co-ord's file, 21	Co-ordinate Transformation, 31		
Rotation, 21	2theta, 31		
Translation, 21	Arm length $(m)$ , 31		
Array Start, 24	Array binning, 31		
Input file, 24	CCD x-axis flip, 31		
Array to Memory, 15, 17	d phi, 31		
Input file, 15, 17	d theta, 31		
Output File, 15, 17	Input data, 31		
Array to VTK, 17	Output amp file, 31		
Input file, 17	Output phase file, 31		
Output File, 17	phi, 31		
Type, 17	Pixel {x,y} (microns), 31		
Auto Centre, 17	Rocking curve type, 31		
Input file, 17	Transform from, 31		
Output File, 17	Transform type, 31		
D	wavelength $(nm)$ , 31		
В	Comments, 13		
Bin, 18	Input file, 13		
Bin dimensions, 18	Output File, 13		
Input file, 18	Compressed Sensing HIO (CSHIO), 27		
Output File, 18	Beta, 27		
Blank Line Fill, 18	Divisor, 27		
Input file, 18	Epsilon, 27		
Output File, 18	Epsilon min, 27		
bonsu.phasing.abstract	Eta, 27		
module, 37	Exp amp, 27		
bonsu.phasing.CSHIO	Iterations, 27		
module, 42			
bonsu.phasing.ER	p-norm, 27		
module, 41	Relax modulus constraint, 27		
bonsu.phasing.HIO	Square root exp amp, 27		
module, 39	Support, 27		
bonsu.phasing.HPR	Conjugate Reflect, 18		
module, 41	Input file, 18		
bonsu.phasing.RAAR	Output File, 18		
module, 42	Convolve, 18		
bonsu.phasing.ShrinkWrap	First Input file, 18		
module, 39	Output File, 18		
bonsu.phasing.SO2D	Second Input file, 18		
module, 43	Crop Pad, 19		
	Crop dimensions: End, 19		
	Crop dimensions: Start, 19		

Input file, 19	Output File, 20
Output File, 19	Sigma, 20
Pad dimensions: End, 19	GetBeta() (bonsu.phasing.abstract.PhaseAbstract
Pad dimensions: Start, 19	method), 38
CSHIO (class in bonsu.phasing.CSHIO), 42	GetDivisor() (bonsu.phasing.CSHIO.CSHIO
Cuboid Support, 19	method), 42
<pre>(xyz) from array, 19 (xyz) from dimensions, 19</pre>	GetDtaumax() (bonsu.phasing.SO2D.SO2D method), 43
Support file, 19	<pre>GetDtaumin() (bonsu.phasing.SO2D.SO2D method),</pre>
Support size, 19	43
D	GetEpsilon() (bonsu.phasing.CSHIO.CSHIO method), 42
	GetEpsilonmin() (bonsu.phasing.CSHIO.CSHIO
Dock Visualisation, 9	method), 42
E	GetEta() (bonsu.phasing.CSHIO.CSHIO method), 42
Empty Array, 15	GetExpdata() (bonsu.phasing.abstract.PhaseAbstract
(xyz) from array, 15	method), 37
(xyz) from dimensions, 15	${\tt GetGammaHWHM()}\ (bonsu.phasing.abstract.Phase Abstract PC$
Output file, 15	method), 38
ER (class in bonsu.phasing.ER), 41	GetMask() (bonsu.phasing.abstract.PhaseAbstract
ER Mask, 26	method), 38
Exp amp, 26	GetMaxphase() (bonsu.phasing.HIO.PCHIO
Iterations, 26	method), 40
Mask, 26	GetMaxphase() (bonsu.phasing.HIO.PGCHIO
Square root exp amp, 26	method), 40
Support, 26	GetMinphase() (bonsu.phasing.HIO.PCHIO
ER Mask with Partial Coherence	method), 40
Optimisation (ER Mask PC), 28	GetMinphase() (bonsu.phasing.HIO.PGCHIO
Beta, 28	method), 40
Exp amp, 28	GetNumiterRL() (bonsu.phasing.abstract.PhaseAbstractPC
Initial PSF HWHM, 28	method), 38
Interval between R-L optimisation, 28	GetNumiterRLinterval()
Iterations, 28	(bonsu.phasing.abstract.PhaseAbstractPC
Iterations, 20 Iterations preceding R-L optimisation,	method), 38
28	${\tt GetNumiterRLpre()}\ (bonsu.phasing.abstract.Phase Abstract PC and the property of the pro$
Mask, 28	method), 38
R-L iterations, 28	GetNumsoiter() (bonsu.phasing.SO2D.SO2D
Reset PSF, 28	method), 43
Square root exp amp, $28$	GetPnorm() (bonsu.phasing.CSHIO.CSHIO method),
Support, 28	42
Zero fill end dimensions of PSF, 28	GetPSF() (bonsu.phasing.abstract.PhaseAbstractPC
ERMask (class in bonsu.phasing.ER), 41	method), 38
ERMaskPC (class in bonsu.phasing.ER), 41	GetPSFZeroEnd() (bonsu.phasing.abstract.PhaseAbstractPC
Error Reduction (ER), 26	method), 38
Exp amp, 26	GetPsiexiterror() (bonsu.phasing.SO2D.SO2D
Iterations, 26	method), 43
Square root exp amp, 26	GetPsiexitratio() (bonsu.phasing.SO2D.SO2D
Support, 26	method), 43
_	GetPsiresetratio() (bonsu.phasing.SO2D.SO2D
F	method), 43
Fourier Transform, 20	GetQ() (bonsu.phasing.HIO.PGCHIO method), 40
Direction, 20	GetReweightiter() (bonsu.phasing.SO2D.SO2D
Input file, 20	method), 43
Output File, 20	${\tt GetSeqdata()}\ (bonsu.phasing.abstract.Phase Abstract$
output 1116, 20	method), 37
G	GetSOS() (bonsu.phasing.abstract.PhaseAbstract
	method), 37
Gaussian Fill, 20	GetSupport() (bonsu.phasing.abstract.PhaseAbstract
Input file, 20	

```
method), 37
                                                     Iterations preceding R-L optimisation,
GetSWCyclelength()
                                                         29
                                                     Mask, 29
        (bonsu.phasing.ShrinkWrap.ShrinkWrap
        method), 39
                                                     R-L iterations, 29
GetSWSigma() (bonsu.phasing.ShrinkWrap.ShrinkWrap
                                                     Reset PSF, 29
        method), 39
                                                     Square root exp amp, 29
GetSWThreshold() (bonsu.phasing.ShrinkWrap.ShrinkWrap Support, 29
        method), 39
                                                     Zero fill end dimensions of PSF, 29
GetTaumax() (bonsu.phasing.SO2D.SO2D method), HPRPC (class in bonsu.phasing.HPR), 41
                                                 Hybrid Input-Output (HIO), 24
        43
                                                     Beta, 24
Н
                                                     Exp amp, 24
HDF5 to Numpy, 16
                                                     Iterations, 24
                                                     Square root exp amp, 24
    HDF key path, 16
    Input HDF file, 16
                                                     Support, 24
    Output File, 16
                                                 Hybrid Projection Reflection (HPR), 27
                                                     Beta, 27
HIO (class in bonsu.phasing.HIO), 39
                                                     Exp amp, 27
HIO Mask, 24
    Beta, 24
                                                     Iterations, 27
                                                     Mask, 27
    Exp amp, 24
                                                     Square root exp amp, 27
    Iterations, 24
                                                     Support, 27
    Mask, 24
    Square root exp amp, 24
    Support, 24
HIO Mask with Partial Coherence
                                                 Image to Numpy, 16
        Optimisation (HIO Mask PC), 28
                                                     Input Image file, 16
    Beta, 28
                                                     Output File, 16
    Exp amp, 28
                                                 Interpolate Object, 20
    Initial PSF HWHM, 28
                                                     Array Bounds: End, 20
    Interval between R-L optimisation, 28
                                                     Array Bounds: Start, 20
    Iterations, 28
                                                     Array grid size, 20
    Iterations preceding R-L optimisation,
                                                     Co-ord's File, 20
        28
                                                     Input file, 20
    Mask, 28
                                                     Interpolation range, 20
    R-L iterations, 28
                                                     Output File, 20
    Reset PSF, 28
                                                 L
    Square root exp amp, 28
    Support, 28
                                                 Load Co-ordinates, 16
    Zero fill end dimensions of PSF, 28
                                                     Input file, 16
HIO Plus, 25
                                                 Load PSF, 16
    Beta, 25
                                                     Input file, 16
    Exp amp, 25
                                                 LorentzFillPSF() (bonsu.phasing.abstract.PhaseAbstractPC
    Iterations, 25
                                                         method), 38
    Mask, 25
    Square root exp amp, 25
                                                 M
    Support, 25
                                                 Mask, 21
HIOMask (class in bonsu.phasing.HIO), 39
                                                     Input file, 21
HIOMaskPC (class in bonsu.phasing.HIO), 40
                                                     Maximum Value, 21
HIOPlus (class in bonsu.phasing.HIO), 39
                                                     Minimum Value, 21
HPR (class in bonsu.phasing.HPR), 41
                                                     Output File, 21
HPR Mask with Partial Coherence
                                                 Median Filter, 21
        Optimisation (HPR Mask PC), 29
                                                     Input file, 21
    Beta, 29
                                                     Output File, 21
    Exp amp, 29
                                                 Memory Arrays
    Initial PSF HWHM, 29
                                                     Array Creation, 10
    Interval between R-L optimisation, 29
                                                 module
    Iterations, 29
                                                     bonsu.phasing.abstract, 37
                                                     bonsu.phasing.CSHIO, 42
```

bonsu.phasing.ER, 41 bonsu.phasing.HIO, 39	Prepare() (bonsu.phasing.abstract.PhaseAbstractPC method), 39
bonsu.phasing.HPR, 41 bonsu.phasing.RAAR, 42	Prepare() (bonsu.phasing.CSHIO.CSHIO method), 42
bonsu.phasing.ShrinkWrap, 39	Prepare() (bonsu.phasing.HIO.PGCHIO method), 40
bonsu.phasing.SO2D, 43	Prepare() (bonsu.phasing.SO2D.SO2D method), 43
N	PrepareCustom() (bonsu.phasing.abstract.PhaseAbstract method), 37
Nexus Viewer I16,13	Python Script, 22
0	Input file, 22 Output File, 22
	output Fire, 22
Object to VTK, 17	R
Co-ord's File, 17	RAAR (class in bonsu.phasing.RAAR), 42
<pre>Input file, 17 Output File, 17</pre>	RAAR Mask with Partial Coherence
Type, 17	Optimisation (RAAR Mask PC), 29
Type, 17	Beta, 29
P	Exp amp, 29
	Initial PSF HWHM, 29
Pause() (bonsu.phasing.abstract.PhaseAbstract	Interval between R-L optimisation, 29
method), 38	Iterations, 29
PCHIO (class in bonsu.phasing.HIO), 39	Iterations preceding R-L optimisation,
PGCHIO ( <i>class in bonsu.phasing.HIO</i> ), 40 Phase Constrained HIO, 25	29
	Mask, 29
Beta, 25	R-L iterations, 29
Exp amp, 25 Iterations, 25	Reset PSF, 29
Mask, 25	Square root exp amp, 29
Phase Max, 25	Support, 29
Phase Min, 25	Zero fill end dimensions of PSF, 29
Square root exp amp, 25	RAARPC (class in bonsu.phasing.RAAR), 42
Support, 25	Random Start, 24
Phase Gradient Constrained HIO, 25	Amp max, 24
Beta, 25	Relaxed Average Alternating Reflection
Exp amp, 25	(RAAR), 27
Iterations, 25	Beta, 27
Mask, 25	Exp amp, 27
Phase Max, 25	Iterations, 27
Phase Min, 25	Mask, 27
Square root exp amp, 25	Square root exp amp, 27
Support, 25	Support, 27
Phase Only ER(POER), 26	Resume() (bonsu.phasing.abstract.PhaseAbstract
Exp amp, 26	<pre>method), 38 Reweighted 2D Saddle Point Optimisation</pre>
Iterations, 26	(SO2D), 30
Mask, 26	Exit Error, 30
Square root exp amp, 26	Exit Ratio, 30
Support, 26	Exp amp, 30
PhaseAbstract (class in bonsu.phasing.abstract), 37	Initial beta, 30
PhaseAbstractPC (class in bonsu.phasing.abstract), 38	Iterations, 30
	Mask, 30
POER (class in bonsu.phasing.ER), 41	Max step increment, 30
Polyhedron Support, 19 (xyz) from array, 19	Max step size, 30
(xyz) from dimensions, 19	Min step increment, 30
Initial Point Coordinates, 19	Reset Ratio, 30
Support file, 19	Square root exp amp, 30
Terminal Point Coordinates, 19	Step optimisation iterations, 30
Prepare() (bonsu.phasing.abstract.PhaseAbstract	Support, 30
method), 37	Rotate Support, 22

Angle, 22		SetNumsoiter()	(bonsu.phasing.SO2D.SO2D
Axis, 22		method), 43	
Input file, 22			su.phasing.abstract.PhaseAbstract
Output File, 22		method), 37	
S		SetPnorm() (bonsu.pha	ssing.CSHIO.CSHIO method),
Save Co-ordinates,	32	SetPSF() (bonsu.phas method), 38	ing.abstract.PhaseAbstractPC
Output file, 32			su.phasing.abstract.PhaseAbstractP0
Save PSF, 31 Input file, 31		method), 38	su.phusing.ubsiruci.i huseribsirucii
Save Residual, 31		SetPsiexiterror()	(bonsu.phasing.SO2D.SO2D
Output file, 31		method), 43	
Save Sequence, 31		SetPsiexitratio()	(bonsu.phasing.SO2D.SO2D
Output file, 31		method), 43	
Save Support, 31		SetPsiresetratio()	(bonsu.phasing.SO2D.SO2D
Output file, 31		method), 43	
Scale Array, 22			HIO.PGCHIO method), 40
Input file, 22		SetReweightiter()	(bonsu.phasing.SO2D.SO2D
Output File, 22		<i>method</i> ), 43	I I I DI AI
Scale factor, 2	2		hasing.abstract.PhaseAbstract
sequence data, 23		method), 37	Large all the Alexand
	phasing.abstract.PhaseAbstract	SetSOS() (bonsu.p. method), 37	hasing.abstract.PhaseAbstract
method), 38	d a compagnition		ı.phasing.abstract.PhaseAbstract
SetDivisor()	(bonsu.phasing.CSHIO.CSHIO	method), 37	i.pnasing.abstract.FnaseAbstract
method), 42	1 : 6030 6030 4 1	* * * * * * * * * * * * * * * * * * * *	hasing.abstract.PhaseAbstract
43	.phasing.SO2D.SO2D method),	method), 37	nasing.aosiraci.1 nascriosiraci
	.phasing.SO2D.SO2D method),	SetSWCyclelength()	
43	.pnasing.502D.502D memoa),		g.ShrinkWrap.ShrinkWrap
SetEpsilon()	(bonsu.phasing.CSHIO.CSHIO	method), 39	
method), 42	( · · · · · · · · · · · · · · · · · · ·	SetSWSigma()(bonsu.p.	hasing.ShrinkWrap.ShrinkWrap
SetEpsilonmin()	(bonsu.phasing.CSHIO.CSHIO	method), 39	
method), 42			nsu.phasing.ShrinkWrap.ShrinkWrap
SetEta() (bonsu.phas	ing.CSHIO.CSHIO method), 42	method), 39	
SetExpdata() (bonsu. method), 37	phasing.abstract.PhaseAbstract	SetTaumax() (bonsu.pa	hasing.SO2D.SO2D method),
SetGammaHWHM()(bon	su.phasing.abstract.PhaseAbstrac	Shrink Wrap, 30	
method), 38	1 0	Algorithm, 30	
SetMask() (bonsu.	phasing.abstract.PhaseAbstract	Beta, 30	
method), 37		Cycle length, 30	
SetMaxphase()	(bonsu.phasing.HIO.PCHIO	Exp amp, 30	
method), 40		Iterations, 30	
SetMaxphase()	(bonsu.phasing.HIO.PGCHIO	Mask, 30	
method), 40		Sigma, 30	20
SetMinphase() <i>method</i> ), 40	(bonsu.phasing.HIO.PCHIO	Square root exp Support, 30	amp, 30
SetMinphase()	(bonsu.phasing.HIO.PGCHIO	Threshold, 30	
method), 40			onsu.phasing.ShrinkWrap), 39
	phasing.abstract.PhaseAbstract	SO2D (class in bonsu.pha SPE to Numpy, 16	sing.SO2D), 43
method), 37	I · I · · · · · · · · · · · · · · · · ·		16
	su.phasing.abstract.PhaseAbstrac	Output File, 16	10
method), 38	(a) ()	_	hasing.abstract.PhaseAbstract
SetNumiterRLinterv	a1() 1g.abstract.PhaseAbstractPC	method), 38	
method), 38	iz.wosiiwci.i iwseAvsiiwcii C		g.ER.SWERMaskPC method),
	bonsu.phasing.abstract.PhaseAbs		
method), 38	2 2		phasing.HIO.SWHIOMaskPC
/,		method), 40	

Start() (bonsu.phasing.HPR.SWHPRPC method), 41	Cut plane normal, 14
Start() (bonsu.phasing.RAAR.SWRAARPC method), 42	Cut plane origin, 14 Input file, 14
Stop() (bonsu.phasing.abstract.PhaseAbstract	Phase options, 14
method), 38	Type, 14
Sum or Subtract Array, 22	View Support, 14
Add or Subtract, 22	Data array file, 14
Input file, 22	Support file, 14
Output File, 22	View VTK Array, 15
support, 23	Amplitude options, 15
SWCSHIO (class in bonsu.phasing.CSHIO), 42	Axes, 15
SWER (class in bonsu.phasing.ER), 41	Cut plane normal, 15
SWERMask (class in bonsu.phasing.ER), 41	Cut plane origin, 15
SWERMaskPC (class in bonsu.phasing.ER), 41	Input file, 15
SWHIO (class in bonsu.phasing.HIO), 39	Phase options, 15
SWHIO (class in bonsu.phasing.HIO), 39	Type, 15
SWHIOMASKPC (class in bonsu.phasing.HIO), 40	Voxel Replace, 23
SWHIOHASKI C (class in bonsu.phasing.HIO), 40	Complex Value, 23
SWHPR (class in bonsu.phasing.HPR), 41	End dimensions, 23
SWHPRPC (class in bonsu.phasing.HPR), 41	Input file, 23
SWPCHIO (class in bonsu.phasing.HIO), 40	Output File, 23
SWPCHIO (class in bonsu.phasing.HIO), 40 SWPGCHIO (class in bonsu.phasing.HIO), 40	Start dimensions, 23
* * *	Start dimensions, 23
SWPrepare() (bonsu.phasing.ShrinkWrap.ShrinkWrap method), 39	W
SWRAAR (class in bonsu.phasing.RAAR), 42	Wrap Data, 23
SWRAARPC (class in bonsu.phasing.RAAR), 42	Input file, 23
SWSO2D (class in bonsu.phasing.SO2D), 43	Output File, 23
SWStart() (bonsu.phasing.ShrinkWrap.ShrinkWrap	Wrap Direction, 23
method), 39	
Т	
Threshold Data, 22	
Input file, 22	
Maximum Value, 22	
Minimum Value, 22	
Output File, 22	
Transpose Array, 23	
Input file, 23	
Output File, 23	
output Tire, 25	
U	
Undock Visualisation, 9	
V	
View Array, 13	
Amplitude options, 13	
Array spacing, 13	
Axes, 13, 14	
Cut plane normal, 13	
Cut plane origin, 13	
Input file, 13	
Isosurface options, 14	
Phase options, 13	
Type, 13	
View Object, 14	
Amplitude options, 14	
Axes, 14	
Co-ord's File 14	