xrayutilities

version 1.7

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December 08, 2020

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Welcome to xrayutilities's documentation!

If you look for downloading the package go to Sourceforge or GitHub (source distribution) or the Python package index (MS Windows binary).

Read more about xrayutilities below or in Journal of Applied Crystallography 2013, Volume 46, 1162-1170

Installation

The easiest way to install *xrayutilities* is using the *Python package index version* https://pypi.python.org/pypi/xrayutilities and execute

> pip install xrayutilities

If you prefer the installation from sources see the Source Installation below.

Introduction

Mailing list and issue tracker

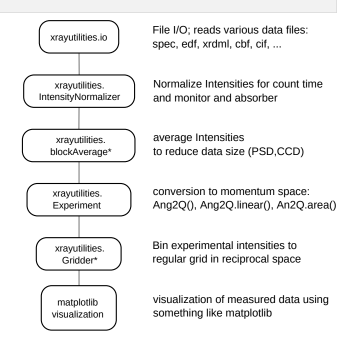
To get in touch with us or report an issue please use the mailing list or the Github issue tracker. When you want to follow announcements of major changes or new releases its recommended to sign up for the mailing list

Overview

xrayutilities is a collection of scripts used to analyze and simulate x-ray diffraction data. It consists of a python package and several routines coded in C. It especially useful for the reciprocal space conversion of diffraction data taken with linear and area detectors. Several models for the simulation of thin film reflectivity and diffraction curves are included.

In the following few concepts of usage for the *xrayutilities* package will be described. First one should get a brief idea of how to analyze x-ray diffraction data with *xrayutilities*. Following that the concept of how angular coordinates of Bragg reflections are calculated is presented. Before describing in detail the installation a minimal example for thin film simulations is shown.

Concept of usage



xrayutilities provides a set of functions to read experimental data from various data file formats. All of them are gathered in the **io**-subpackage. After reading data with a function from the io-submodule the data might be corrected for monitor counts and/or absorption factor of a beam attenuator. A special set of functions is provided to perform this for point, linear and area detectors.

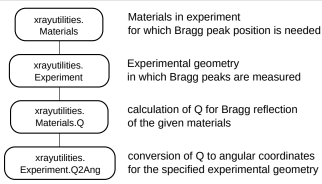
Since the amount of data taken with modern detectors often is too large to be able to work with them properly, a functions for reducing the data from linear and area detectors are provided. They use block-averaging to reduce the amount of data. Use those carefully not to loose the features you are interested in in your measurements.

After the pre-treatment of the data, the core part of the package is the transformation of the angular data to reciprocal space. This is done as described in more detail below using the **experiment**-module. The classes provided within the experiment module provide routines to help performing X-ray diffraction experiments. This includes methods to calculate the diffraction angles (described below) needed to align crystalline samples and to convert data between angular and reciprocal space. The conversion from angular to reciprocal space is implemented very general for various goniometer geometries. It is especially useful in combination with linear and area detectors as described in this article. In standard cases, Users will only need the initialized routines, which predefine a certain goniometer geometry like the popular four-circle and six-circle geometries.

After the conversion to reciprocal space, it is convenient to transform the data to a regular grid for visualization. For this purpose the **gridder**-module has been included into *xrayutilities*. For the visualization of the data in reciprocal space the usage of matplotlib is recommended.

A practical example showing the usage is given below.

Angle calculation using the material classes



Calculation of angles needed to align Bragg reflections in various diffraction geometries is done using the Materials defined in the materials-package. This package provides a set of classes to describe crystal lattices and materials. Once such a material is properly defined one can calculate its properties, which includes the reciprocal lattice points, lattice plane distances, optical properties like the refractive index, the structure factor (including the atomic scattering factor) and the complex polarizability. These atomic properties are extracted from a database included in *xrayutilities*.

Using such a material and an experimental class from the **experiment**-module, describing the experimental setup, the needed diffraction angles can be calculated for certain coplanar diffraction (high, low incidence), grazing incidence diffraction and also special non-coplanar diffraction geometries. In the predefined experimental classes fixed geometries are used. For angle calculation of custom geometries using arbitrary geometries (max. of three free angles) the **q2ang_fit**-module can be used as described in one of the included example files.

hello world

A first example with step by step explanation is shown in the following. It showcases the use of *xrayutilities* to calculate angles and read a scan recorded with a linear detector from spec-file and plots the result as reciprocal space map using matplotlib.

```
1 """
2 Example script to show how to use xrayutilities to read and plot
3 reciprocal space map scans from a spec file created at the ESRF/ID10B
4
5 for details about the measurement see:
6    D Kriegner et al. Nanotechnology 22 425704 (2011)
```

```
7
      http://dx.doi.org/10.1088/0957-4484/22/42/425704
 8 """
 9
10 import os
11
12 import matplotlib.pyplot as plt
13 import numpy
14 import xrayutilities as xu
15
16 # global setting for the experiment
17 sample = "test" # sample name used also as file name for the data file
18 energy = 8042.5 # x-ray energy in eV
19 center_ch = 715.9 # center channel of the linear detector
20 chpdeg = 345.28 # channels per degree of the linear detector
21 roi = [100, 1340] # region of interest of the detector
22 nchannel = 1500 # number of channels of the detector
2.3
24 # intensity normalizer function responsible for count time and absorber
25 # correction
26 normalizer_detcorr = xu.IntensityNormalizer(
27
      "MCA",
      mon="Monitor",
28
29
      time="Seconds",
       absfun=lambda d: d["detcorr"] / d["psd2"].astype(numpy.float))
30
31
32 # substrate material used for Bragg peak calculation to correct for
33 # experimental offsets
34 InP = xu.materials.InP
36 # initialize experimental class to specify the reference directions of your
37 # crystal
38 # 11-2: inplane reference
39 # 111: surface normal
40 hxrd = xu.HXRD(InP.Q(\mathbf{1}, \mathbf{1}, -\mathbf{2}), InP.Q(\mathbf{1}, \mathbf{1}), en=energy)
41
42 # configure linear detector
43 # detector direction + parameters need to be given
44 # mounted along z direction, which corresponds to twotheta
45 hxrd.Ang2Q.init_linear('z-', center_ch, nchannel, chpdeg=chpdeg, roi=roi)
46
47 # read spec file and save to HDF5-file
48 # since reading is much faster from HDF5 once the data are transformed
49 h5file = os.path.join("data", sample + ".h5")
50 try:
51
      s # try if spec file object already exist ("run -i" in ipython)
52 except NameError:
53
       s = xu.io.SPECFile(sample + ".spec", path="data")
54 else:
55
       s.Update()
56 s.Save2HDF5(h5file)
57
58 ################################
59 # InP (333) reciprocal space map
60 omalign = 43.0529 # experimental aligned values
61 ttalign = 86.0733
62 [omnominal, _, _, ttnominal] = hxrd.Q2Ang(
63
       InP.Q(3, 3, 3)) # nominal values of the substrate peak
64
65 # read the data from the HDF5 file
66 # scan number:36, names of motors in spec file: omega= sample rocking, gamma =
```

```
67 # twotheta
68 [om, tt], MAP = xu.io.geth5_scan(h5file, 36, 'omega', 'gamma')
69 # normalize the intensity values (absorber and count time corrections)
70 psdraw = normalizer_detcorr(MAP)
71 # remove unusable detector channels/regions (no averaging of detector channels)
72 psd = xu.blockAveragePSD(psdraw, 1, roi=roi)
73
74 # convert angular coordinates to reciprocal space + correct for offsets
75 [qx, qy, qz] = hxrd.Ang2Q.linear(
76
       om, tt,
77
       delta=[omalign - omnominal, ttalign - ttnominal])
78
79 # calculate data on a regular grid of 200x201 points
80 gridder = xu.Gridder2D(200, 201)
81 gridder(qy, qz, psd)
82 # maplog function limits the shown dynamic range to 8 orders of magnitude
83 # from the maxium
84 INT = xu.maplog(gridder.data.T, 8., 0)
85
86 # plot the intensity as contour plot using matplotlib
88 cf = plt.contourf(gridder.xaxis, gridder.yaxis, INT, 100, extend='min')
89 plt.xlabel(r'$Q_{[11\bar2]}$ ($\mathrm{\AA}^{-1}$)')
90 plt.ylabel(r'Q_{[\bar x]} \ (\bar x_{AA}^{-1})')
91 cb = plt.colorbar(cf)
92 cb.set_label(r"$\log($Int$)$ (cps)")
```

More such examples can be found on the Examples page.

X-ray diffraction and reflectivity simulations

xrayutilties includes a database with optical properties of materials and therefore simulation of reflectivity and diffraction data can be accomplished with relatively little additional input. When the stack of layers is defined along with the layer thickness and material several models for calculation of X-ray reflectivity and dynamical/kinematical X-ray diffraction are provided.

A minimal example for an AlGaAs superlattice structure is shown below. It shows how a basic stack of a superlattice is built from its ingredients and how the reflectivity and dynamical diffraction model are initialized in the most basic form:

```
import xrayutilities as xu
# Build the pseudomorphic sample stack using the elastic parameters
sub = xu.simpack.Layer(xu.materials.GaAs, inf)
lay1 = xu.simpack.Layer(xu.materials.AlGaAs(0.25), 75, relaxation=0.0)
lay2 = xu.simpack.Layer(xu.materials.AlGaAs(0.75), 25, relaxation=0.0)
pls = xu.simpack.PseudomorphicStack001('pseudo', sub+10*(lay1+lay2))
# simulate reflectivity
m = xu.simpack.SpecularReflectivityModel(pls, sample_width=5, beam_width=0.3)
alphai = linspace(0, 10, 1000)
Ixrr = m.simulate(alphai)
# simulate dynamical diffraction curve
alphai = linspace(29, 38, 1000)
md = xu.simpack.DynamicalModel(pls)
Idyn = md.simulate(alphai, hkl=(0, 0, 4))
```

More detailed examples and description of model parameters can be found on the Simulation examples page or in the examples directory.

xrayutilities Python package

xrayutilities is a Python package for assisting with x-ray diffraction experiments. Its the python package included in xrayutilities.

It helps with planning experiments as well as analyzing the data.

Authors:

Dominik Kriegner <dominik.kriegner@gmail.com> and Eugen Wintersberger <eugen.wintersberger@desy.de> for more details see the full API documentation of xrayutilities found here: Examples and API-documentation.

Source Installation

Express instructions

- install the dependencies (Windows: Python(x,y) or WinPython; Linux/Unix: see below for dependencies).
- download xrayutilities from here or use git to check out the latest version.
- open a command line and navigate to the downloaded sources and execute:

```
> pip install
```

which will install *xrayutilities* to the default directory. It should be possible to use it (*import xrayutilities*) from now on in python scripts.

Detailed instructions

Installing xrayutilities is done using Python's setuptools

The package can be installed on Linux, Mac OS X and Microsoft Windows, however, it is mostly tested on Linux/Unix platforms. Please inform one of the authors in case the installation fails!

Required third party software

To keep the coding effort as small as possible *xrayutilities* depends on a large number of third party libraries and Python modules.

The needed runtime dependencies are:

- **Python** the scripting language in which most of *xrayutilities* code is written in. (>= 3.6, for Python 2.7 use *xrayutilities* 1.5.X or older)
- **Numpy** a Python module providing numerical array objects (version >= 1.9)
- **Scipy** a Python module providing standard numerical routines, which is heavily using numpy arrays (version >= 0.13.0)
- h5py a powerful Python interface to HDF5.

For several features optional dependencies are needed:

- Matplotlib a Python module for high quality 1D and 2D plotting (optional, version >= 3.1.0)
- Imfit a Python module for least-squares minimization with bounds and constraints (optionally needed for fitting XRR/XRD data)
- **IPython** although not a dependency of *xrayutilities* the IPython shell is perfectly suited for the interactive use of the *xrayutilities* python package.
- mayavi only used optionally in Crystal.show_unitcell where it produces a superior visualization to otherwise used Matplotlib 3D plots

Additionally, the following Python modules are needed when building *xrayutilities* from source or wanting to test your installation:

- C-compiler Gnu Compiler Collection or any compatible C compiler. On windows you most probably want to use the Microsoft compilers.
- Python development headers
- setuptools build system
- tox needed for running the pre-configured unittest environment, which in principal can also be achieved only by the unittest package (optional)

For building the documention (which you do not need to do) the requirements are:

- sphinx the Python documentation generator
- numpydoc sphinx-extension needed to parse the API-documention
- rst2pdf pdf-generation using sphinx

After installing all required packages you can continue with installing and building the C library.

Building and installing the library and python package

Although the *setup.py* script can be called manually its recommended to always use pip to install *xrayutilities*, which can be done by executing

```
>pip install .
```

or

```
>pip install --prefix=INSTALLPATH
```

in the root directory of the source distribution.

The *-prefix* option sets the root directory for the installation. If it is omitted the libary is installed under the systems default directories (recommended).

Setup of the Python package

You need to make your Python installation aware of where to look for the module. This is usually only needed when installing in non-standard *<install path>* locations. For this case append the installation directory to your *PYTHONPATH* environment variable by

```
>export PYTHONPATH=$PYTHONPATH:<local install path>/lib64/python2.7/site-packages
```

on a Unix/Linux terminal. Or, to make this configuration persistent append this line to your local *.bashrc* file in your home directory. On MS Windows you would like to create a environment variable in the system preferences under system in the advanced tab (Using Python package managers this should be done automatically). Be sure to use the correct directory which might be similar to

```
<local install path>/Lib/site-packages
```

on Windows systems.

Notes for installing on Windows

Since there is no packages manager on Windows the packages need to be installed manual (including all the dependecies) or a pre-packed solution needs two be used. We strongly suggest to use either Anaconda, Python(x,y) or WinPython Python distributions, which include already all of the needed dependencies for installing *xrayutilities*.

One can proceed with the installation of *xrayutilities* directly! The easiest way to do this on windows is to use the binaries distributed on the Python package index or using *pip*, otherwise one can follow the general installation instructions. On Anaconda it can also be done using the conda-forge *xrayutilities* package.

Examples and API-documentation

Examples

In the following a few code-snippets are shown which should help you getting started with *xrayutilities*. Not all of the codes shown in the following will be run-able as stand-alone script. For fully running scripts look in the examples directory in the download found here.

Reading data from data files

The io submodule provides classes for reading x-ray diffraction data in various formats. In the following few examples are given.

Reading SPEC files

Working with spec files in xrayutilities can be done in two distinct ways.

- 1. parsing the spec file for scan headers; and parsing the data only when needed
- 2. parsing the spec file for scan headers; parsing all data and dump them to an HDF5 file; reading the data from the HDF5 file.

Both methods have their pros and cons. For example when you parse the spec-files over a network connection you need to re-read the data again over the network if using method 1) whereas you can dump them to a local file with method 2). But you will parse data of the complete file while dumping it to the HDF5 file.

Both methods work incremental, so they do not start at the beginning of the file when you reread it, but start from the last position they were reading and work with files including data from linear detectors.

An working example for both methods is given in the following.

```
1 import xrayutilities as xu
 2 import os
 4 # open spec file or use open SPECfile instance
 5 try: s
 6 except NameError:
 7
       s = xu.io.SPECFile("sample_name.spec", path="./specdir")
 8
 9 # method (1)
10 s.scan10.ReadData()
11 scan10data = s.scan10.data
12
13 # method (2)
14 h5file = os.path.join("h5dir", "h5file.h5")
15 s.Save2HDF5(h5file) # save content of SPEC file to HDF5 file
16 # read data from HDF5 file
17 [angle1, angle2], scan10data = xu.io.geth5_scan(h5file, [10],
18
                                                     "motorname1",
19
                                                     "motorname2")
```

Seealso

the fully working example hello world

In the following it is shown how to re-parsing the SPEC file for new scans and reread the scans (1) or update the HDF5 file(2)

```
1 s.Update() # reparse for new scans in open SPECFile instance
2
```

Reading EDF files

EDF files are mostly used to store CCD frames at ESRF recorded from various different detectors. This format is therefore used in combination with SPEC files. In an example the EDFFile class is used to parse the data from EDF files and store them to an HDF5 file. HDF5 if perfectly suited because it can handle large amount of data and compression.

```
1 import xrayutilities as xu
2 import numpy
3
4 specfile = "specfile.spec"
5 h5file = "h5file.h5"
6
7 s = xu.io.SPECFile(specfile)
8 s.Save2HDF5(h5file) # save to hdf5 file
9
10 # read ccd frames from EDF files
11 for i in range(1, 1001, 1):
12    efile = "edfdir/sample_%04d.edf" % i
13    e = xu.io.edf.EDFFile(efile)
14    e.ReadData()
15    e.Save2HDF5(h5file, group="/frelon_%04d" % i)
```

Seealso

the fully working example provided in the examples directory perfectly suited for reading data from beamline ID01

Reading XRDML files

Files recorded by Panalytical diffractometers in the .xrdml format can be parsed. All supported file formats can also be parsed transparently when they are saved as compressed files using common compression formats. The parsing of such compressed .xrdml files conversion to reciprocal space and visualization by gridding is shown below:

Seealso

the fully working example provided in the examples directory

Other formats

Other formats which can be read include

- Rigaku .ras files.
- files produces by the experimental control software at Hasylab/Desy (spectra).
- numor files from the ILL neutron facility
- ccd images in the tiff file format produced by RoperScientific CCD cameras and Perkin Elmer detectors.
- files from recorded by Seifert diffractometer control software (.nja)
- support is also provided for reading of cif files from structure databases to extract unit cell parameters as well es read data from those files (pdCIF, ESG files)

See the examples directory for more information and working example scripts.

Angle calculation using Experiment and materials classes

Methods for high angle x-ray diffraction experiments. Mostly for experiments performed in coplanar scattering geometry. An example will be given for the calculation of the position of Bragg reflections.

```
1 import xrayutilities as xu
2 Si = xu.materials.Si  # load material from materials submodule
3
4 # initialize experimental class with directions from experiment
5 hxrd = xu.HXRD(Si.Q(1, 1, -2), Si.Q(1, 1, 1))
6 # calculate angles of Bragg reflections and print them to the screen
7 om, chi, phi, tt = hxrd.Q2Ang(Si.Q(1, 1, 1))
8 print("Si (111)")
9 print("om,tt: %8.3f %8.3f" % (om, tt))
10 om, chi, phi, tt = hxrd.Q2Ang(Si.Q(2, 2, 4))
11 print("Si (224)")
12 print("om,tt: %8.3f %8.3f" % (om, tt))
```

Note that on line 5 the HXRD class is initialized without specifying the energy used in the experiment. It will use the default energy stored in the configuration file, which defaults to CuK-alpha1.

One could also call:

```
hxrd = xu.HXRD(Si.Q(1, 1, -2), Si.Q(1, 1, 1), en=10000) # energy in eV
```

to specify the energy explicitly. The **HXRD** class by default describes a four-circle goniometer as described in more detail here.

Similar functions exist for other experimental geometries. For grazing incidence diffraction one might use

```
gid = xu.GID(Si.Q(1, -1, 0), Si.Q(0, 0, 1))
# calculate angles and print them to the screen
(alphai, azimuth, tt, beta) = gid.Q2Ang(Si.Q(2, -2, 0))
print("azimuth,tt: %8.3f" % (azimuth, tt))
```

There is on implementation of a GID 2S+2D diffractometer. Be sure to check if the order of the detector circles fits your goniometer, otherwise define one yourself!

There exists also a powder diffraction class, which is able to convert powder scans from angular to reciprocal space.

```
1 import xrayutilities as xu
2 import numpy
3
4 energy = 'CuKa12'
5
6 # creating powder experiment
7 xup = xu.PowderExperiment(en=energy)
8 theta = arange(0, 70, 0.01)
9 q = xup.Ang2Q(theta)
```

More information about powdered materials can be obtained from the **PowderDiffraction** class. It contains information about peak positions and intensities

```
1 >>> print(xu.simpack.PowderDiffraction(xu.materials.In))
     Powder diffraction object
 3
     Powder-In (volume: 1, )
 4
 5
     Lattice:
 6
     a1 = (3.252300 \ 0.000000 \ 0.000000), \ 3.252300
     a2 = (0.000000 \ 3.252300 \ 0.000000), \ 3.252300
 7
 8
     a3 = (0.000000 \ 0.0000000 \ 4.946100), \ 4.946100
 9
     alpha = 90.000000, beta = 90.000000, gamma = 90.000000
10
     Lattice base:
     Base point 0: In (49) (0.000000 0.000000 0.000000) occ=1.00 b=0.00
11
12
     Base point 1: In (49) (0.500000 0.500000 0.500000) occ=1.00 b=0.00
13
     Reflections:
14
           h k l | tth
                                 | Q|
                                                          Int (%)
15
                                                  Int
16
          [0, 1, -1] 32.9338 2.312
[0, 0, -2] 36.2964 2.541
17
                                               217.24 100.00
                                                              19.19
18
                                                  41.69
          [-1, 1, 0]
                      39.1392
                                                               31.09
19
                                      2.732
                                                   67.54
         [-1, -1, -2]
20
                        54.4383
                                      3.731
                                                   50.58
                                                                23.28
21
```

If you are interested in simulations of powder diffraction patterns look at section Powder diffraction simulations

Using the Gridder classes

xrayutilities provides Gridder classes for 1D, 2D, and 3D data sets. These Gridders map irregular spaced data onto a regular grid. This is often needed after transforming data measured at equally spaced angular positions to reciprocal space were their spacing is irregular.

In 1D this process actually equals the calculation of a histogram. Below you find the most basic way of using the Gridder in 2D. Other dimensions work very similar.

The most easiest use (what most user might need) is

```
1 import xrayutilities as xu # import Python package
2 g = xu.Gridder2D(100, 101) # initialize the Gridder object, which will
3 # perform Gridding to a regular grid with 100x101 points
4 #===== load some data here =====
5 g(x, y, data) # call the gridder with the data
6 griddata = g.data # the data attribute contains the gridded data.
```

A more complicated example showing also sequential gridding is shown below. You need sequential gridding when you can not load all data at the same time, which is often problematic with 3D data sets. In such cases you need to specify the data range before the first call to the gridder.

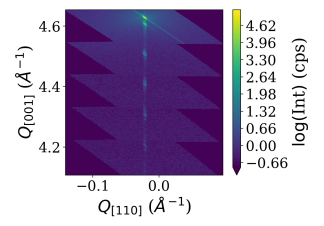
```
1 import xrayutilities as xu # import Python package
2 g = xu.Gridder2D(100, 101) # initialize the Gridder object
3 g.KeepData(True)
4 g.dataRange(1, 2, 3, 4) # (xgrd_min, xgrd_max, ygrd_min, ygrd_max)
5 #===== load some data here =====
6 g(x, y, data) # call the gridder with the data
7 griddata = g.data # the data attribute contains the so far gridded data.
8
9 #===== load some more data here =====
10 g(x, y, data) # call the gridder with the new data
11 griddata = g.data # the data attribute contains the combined gridded data.
```

Gridder2D for visualization

Based on the example of parsed data from XRDML files shown above (Reading XRDML files) we show here how to use the **Gridder2D** class together with matplotlibs contourf.

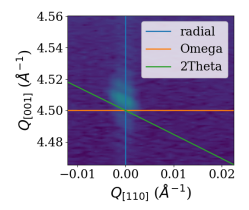
```
1 Si = xu.materials.Si
2 hxrd = xu.HXRD(Si.Q(1, 1, 0), Si.Q(0, 0, 1))
3 qx, qy, qz = hxrd.Ang2Q(om, tt)
4 gridder = xu.Gridder2D(200, 600)
5 gridder(qy, qz, psd)
6 INT = xu.maplog(gridder.data.transpose(), 6, 0)
7 # plot the intensity as contour plot
8 plt.figure()
9 cf = plt.contourf(gridder.xaxis, gridder.yaxis, INT, 100, extend='min')
10 plt.xlabel(r'$Q_{[110]}$ ($\mathrm{\AA^{-1}}$)')
11 plt.ylabel(r'$Q_{[001]}$ ($\mathrm{\AA^{-1}}$)')
12 cb = plt.colorbar(cf)
13 cb.set_label(r"$\log($Int$)$ (cps)")
14 plt.tight_layout()
```

The shown script results in the plot of the reciprocal space map shown below.

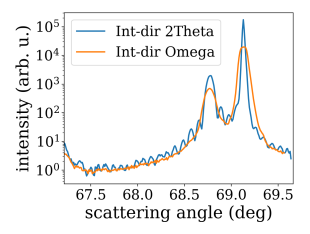


Line cuts from reciprocal space maps

Using the <code>analysis</code> subpackage one can produce line cuts. Starting from the reciprocal space data produced by the reciprocal space conversion as in the last example code we extract radial scan along the crystal truncation rod. For the extraction of line scans the respective functions offer to integrate the data along certain directions. In the present case integration along '2Theta' gives the best result since a broadening in that direction was caused by the beam footprint in the particular experiment. For different line cut functions various integration directions are possible. They are visualized in the figure below.



```
1 # line cut with integration along 2theta to remove beam footprint broadening
 2 qzc, qzint, cmask = xu.analysis.get_radial_scan([qy, qz], psd, [0, 4.5],
 3
                                                    1001, 0.155, intdir='2theta')
 4
 5 # line cut with integration along omega
  qzc_om, qzint_om, cmask_om = xu.analysis.get_radial_scan([qy, qz], psd, [0, 4.5],
 7
                                                    1001, 0.155, intdir='omega')
 8 plt.figure()
 9 plt.semilogy(qzc, qzint, label='Int-dir 2Theta')
10 plt.semilogy(qzc_om, qzint_om, label='Int-dir Omega')
11 plt.xlabel(r'scattering angle (deg)')
12 plt.ylabel(r'intensity (arb. u.)')
13 plt.legend()
14 plt.tight_layout()
```



Seealso

the fully working example provided in the examples directory and the other line cut functions in line cuts

Using the materials subpackage

xrayutilities provides a set of Python classes to describe crystal lattices and materials.

Examples show how to define a new material by defining its lattice and deriving a new material, furthermore materials can be used to calculate the structure factor of a Bragg reflection for an specific energy or the energy dependency of its structure factor for anomalous scattering. Data for this are taken from a database which is included in the download.

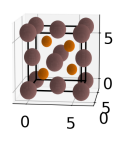
First defining a new material from scratch is shown. This is done from the space group and Wyckhoff positions of the atoms inside the unit cell. Depending on the space group number the initialization of a new SGLattice object expects a different amount of parameters. For a cubic materials only the lattice parameter a should be given while for a triclinic materials a, b, c, alpha, beta, and gamma have to be specified. Its similar for the Wyckoff positions. While some Wyckoff positions require only the type of atom others have some free parameters which can be specified. Below we show the definition of zincblende InP as well as for its hexagonal wurtzite polytype together with a quick visualization of the unit cells. A more accurate visualization of the unit cell can be performed when using show_unitcell() with the Mayavi mode or by using the CIF-exporter and an external tool.

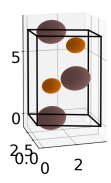
```
1 import matplotlib.pyplot as plt
2 import xrayutilities as xu
3
4 # elements (which contain their x-ray optical properties) are loaded from
5 # xrayutilities.materials.elements
6 In = xu.materials.elements.In
```

```
7 P = xu.materials.elements.P
 8
 9 # define elastic parameters of the material we use a helper function which
10 # creates the 6x6 tensor needed from the only 3 free parameters of a cubic
11 # material.
12 elastictensor = xu.materials.CubicElasticTensor(10.11e+10, 5.61e+10,
13
                                                    4.56e+10)
14 # definition of zincblende InP:
15 InP = xu.materials.Crystal(
16
      "InP", xu.materials.SGLattice(216, 5.8687, atoms=[In, P],
17
                                     pos=['4a', '4c']),
18
       elastictensor)
19
20 # a hexagonal equivalent which shows how parameters change for material
21 # definition with a different space group. Since the elasticity tensor is
22 # optional its not specified here.
23 InPWZ = xu.materials.Crystal(
       "InP(WZ)", xu.materials.SGLattice(186, 4.1423, 6.8013,
25
                                          atoms=[In, P], pos=[('2b', 0),
                                                              ('2b', 3/8.)]))
2.6
27 f = plt.figure()
28 InP.show_unitcell(fig=f, subplot=121)
29 title('InP zincblende')
30 InPWZ.show_unitcell(fig=f, subplot=122)
31 title('InP wurtzite')
```

InP wurtzite

InP zincblende





InP (in both variants) is already included in the xu.materials module and can be loaded by

```
InP = xu.materials.InP
InPWZ = xu.materials.InPWZ
```

Similar definitions exist for many other materials. Alternatively to giving the Wyckoff labels and parameters one can also specify the position of one atom for every unique site within the unit cell. *xrayutilities* will then search the corresponding Wyckoff position of this atom and populate therefore populate all equivalent sites as well. For the example of InP in zincblende form the material definition could also look as shown below. Note that instead of the elements also the elemental symbol as string can be used:

Using the material properties the calculation of the reflection strength of a Bragg reflection can be done as follows

```
1 import xrayutilities as xu
2 import numpy
3
4 # defining material and experimental setup
5 InAs = xu.materials.InAs
```

```
6 energy= 8048 # eV
7
8 # calculate the structure factor for InAs (111) (222) (333)
9 hkllist = [[1, 1, 1], [2, 2, 2], [3, 3, 3]]
10 for hkl in hkllist:
11    qvec = InAs.Q(hkl)
12    F = InAs.StructureFactor(qvec, energy)
13    print(" |F| = %8.3f" % numpy.abs(F))
```

Similar also the energy dependence of the structure factor can be determined

```
1 import matplotlib.pyplot as plt
2
3 energy= numpy.linspace(500, 20000, 5000) # 500 - 20000 eV
4 F = InAs.StructureFactorForEnergy(InAs.Q(1, 1, 1), energy)
5
6 plt.figure(); plt.clf()
7 plt.plot(energy, F.real, '-k', label='Re(F)')
8 plt.plot(energy, F.imag, '-r', label='Imag(F)')
9 plt.xlabel("Energy (eV)"); plt.ylabel("F"); plt.legend()
```

It is also possible to calculate the components of the structure factor of atoms, which may be needed for input into XRD simulations.

```
1 # f = f0(|Q|) + f1(en) + j * f2(en)
2 import xrayutilities as xu
3 import numpy
4
5 Fe = xu.materials.elements.Fe # iron atom
6 Q = numpy.array([0, 0, 1.9], dtype=numpy.double)
7 en = 10000 # energy in eV
8
9 print("Iron (Fe): E: %9.1f eV" % en)
10 print("f0: %8.4g" % Fe.f0(numpy.linalg.norm(Q)))
11 print("f1: %8.4g" % Fe.f1(en))
12 print("f2: %8.4g" % Fe.f2(en))
```

Transformation of SGLattice

SGLattice-objects can be transformed to use a different unit cell setting. This can be used to for example change the origin choice after the material definition or to convert into a totally different setting, e.g. for simulation purposes.

The code below shows the example of the Diamond structure converted between the two different origin choices

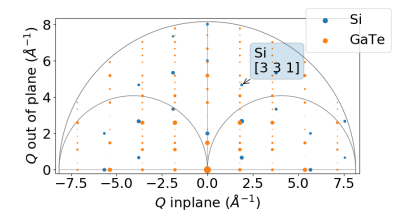
For dynamical diffraction simulations of cubic crystals with (111) surface it might be required to convert the unit cell in a way that a principle axis is pointing along the surface normal. Using an apropriate conversion matrix this is shown for the example of InP

While the built in InP uses the cubic setting with space group F-43m(#216) the converted lattice has rhombohedral space group (in this case R3m(#160)) and converted atomic positions.

Visualization of the Bragg peaks in a reciprocal space plane

If you want to explore which peaks are available and reachable in coplanar diffraction geometry and what their relationship between different materials is *xrayutilities* provides a function which generates a slightly interactive plot which helps you with this task.

The generated plot shows all the existing Bragg spots, their (hkl) label is shown when the mouse is over a certain spot and the diffraction angles calculated by the given **HXRD** object is printed when you click on a certain spot. Not that the primary beam is assumed to come from the left, meaning that high incidence geometry occurs for all peaks with positive inplane momentum transfer.



Calculation of diffraction angles for a general geometry

Often the restricted predefined geometries are not corresponding to the experimental setup, nevertheless *xrayutilities* is able to calculate the goniometer angles needed to reach a certain reciprocal space position.

For this purpose the goniometer together with the geometric restrictions need to be defined and the q-vector in laboratory reference frame needs to be specified. This works for arbitrary goniometer, however, the user is expected to set up bounds to put restrictions to the number of free angles to obtain reproducible results. In general only three angles are needed to fit an arbitrary q-vector (2 sample + 1 detector angles or 1 sample + 2 detector). More goniometer angles can be kept free if some pseudo-angle constraints are used instead.

The example below shows the necessary code to perform such an angle calculation for a custom defined material with orthorhombic unit cell.

```
8
       return 1
9
10 latticeConstants=[5.600, 7.706, 5.3995]
11 SmFeO3 = xu.materials.Crystal("SmFeO3", Pnma(*latticeConstants))
12 # 2S+2D goniometer
13 qconv=xu.QConversion(('x+', 'z+'), ('z+', 'x+'), (0, 1, 0))
14 # [1,1,0] surface normal
15 hxrd = xu.HXRD(SmFeO3.Q(\mathbf{0}, \mathbf{0}, \mathbf{1}), SmFeO3.Q(\mathbf{1}, \mathbf{1}, \mathbf{0}), qconv=qconv)
16
17 hkl=(2, 0, 0)
18 g material = SmFeO3.Q(hkl)
19 q_laboratory = hxrd.Transform(q_material) # transform
20
21 print('SmFeO3: \thkl ', hkl, '\tqvec ', np.round(q_material, 5))
22 print('Lattice plane distance: %.4f' % SmFeO3.planeDistance(hkl))
23
24 #### determine the goniometer angles with the correct geometry restrictions
25 # tell bounds of angles / (min, max) pair or fixed value for all motors.
26 # maximum of three free motors! here the first goniometer angle is fixed.
27 # om, phi, tt, delta
28 bounds = (5, (-180, 180), (-1, 90), (-1, 90))
29 ang, qerror, errcode = xu.Q2AngFit(q_laboratory, hxrd, bounds)
30 print('err %d (%.3g) angles %s' % (errcode, qerror, str(np.round(ang, 5))))
31 # check that qerror is small!!
32 print('sanity check with back-transformation (hkl): ',
         np.round(hxrd.Ang2HKL(*ang,mat=SmFeO3),5))
```

In the example above all angles can be kept free if a pseudo-angle constraint is used in addition. This is shown below for the incidence angle, which when fixed to 5 degree results in the same goniometer angles as shown above. Currently two helper functions for incidence and exit angles (incidenceAngleConst()) and exitAngleConst()) are implemented, but user-defined functions can be supplied.

```
1 aiconstraint = xu.q2ang_fit.incidenceAngleConst
2 bounds = ((0, 90), (-180, 180), (-1, 90), (-1, 90))
3 ang, qerror, errcode = xu.Q2AngFit(
4     q_laboratory, hxrd, bounds,
5     constraints={'type':'eq', 'fun': lambda_a: aiconstraint(a, 5, hxrd)})
```

User-specific config file

Several options of *xrayutilities* can be changed by options in a config file. This includes the default x-ray energy as well as parameters to set the number of threads used by the parallel code and the verbosity of the output.

The default options are stored inside the installad Python module and should not be changed. Instead it is suggested to use a user-specific config file '~/.xrayutilities.conf' or a 'xrayutilities.conf' file in the working directory.

An example of such a user config file is shown below:

```
1 # begin of xrayutilities configuration
2 [xrayutilities]
3
4 # verbosity level of information and debugging outputs
5 # 0: no output
6 # 1: very import notes for users
7 # 2: less import notes for users (e.g. intermediate results)
8 # 3: debuging output (e.g. print everything, which could be interesing)
9 # levels can be changed in the config file as well
10 verbosity = 1
11
12 # default wavelength in Angstrom,
13 wavelength = MoKal # Molybdenum K alphal radiation (17479.374eV)
14
```

```
15 # default energy in eV
16 # if energy is given wavelength settings will be ignored
17 #energy = 10000 #eV
18
19 # number of threads to use in parallel sections of the code
20 nthreads = 1
21 # 0: the maximum number of available threads will be used (as returned by 22 # omp_get_max_threads())
23 # n: n-threads will be used
```

Determining detector parameters

In the following three examples of how to determine the detector parameters for linear and area detectors is given. The procedure we use is in more detail described in this article.

Linear detectors

To determine the detector parameters of a linear detector one needs to perform a scan with the detector angle through the primary beam and aquire a detector spectrum at any point.

Using the following script determines the parameters necessary for the detector initialization, which are:

- · pixelwidth of one channel
- the center channel
- and the detector tilt (optional)

```
1 """
 2 example script to show how the detector parameters
 3 such as pixel width, center channel and detector tilt
 4 can be determined for a linear detector.
 5
 6
 7 import os
 8
 9 import xrayutilities as xu
10
11 # load any data file with with the detector spectra of a reference scan
12 # in the primary beam, here I use spectra measured with a Seifert XRD
13 # diffractometer
14 dfile = os.path.join("data", "primarybeam_alignment20130403_2_dis350.nja")
15 s = xu.io.SeifertScan(dfile)
16
17 ang = s.axispos["T"] # detector angles during the scan
18 spectra = s.data[:, :, 1] # detector spectra aquired
19
20 # determine detector parameters
21 # this function accepts some optional arguments to describe the goniometer
22 # see the API documentation
23 pwidth, cch, tilt = xu.analysis.linear_detector_calib(ang, spectra,
24
                                                         usetilt=True)
```

Area detector (Variant 1)

To determine the detector parameters of a area detector one needs to perform scans with the detector angles through the primary beam and aquire a detector images at any position. For the area detector at least two scans (one with the outer detector and and one with the inner detector angle) are required.

Using the following script determines the parameters necessary for the detector initialization from such scans in the primary beam only. Further down we discuss an other variant which is also able to use additionally detector images recorded at the Bragg reflection of a known reference crystal.

The determined detector parameters are:

- center channels: position of the primary beam at the true zero position of the goniometer (considering the outer angle offset) (2 parameters)
- pixelwidth of the channels in both directions (2 parameters), these two parameters can be replaced by the detector distance (1 parameter) if the pixel size is given as an input
- detector tilt azimuth in degree from 0 to 360
- detector tilt angle in degree (>0deg)
- detector rotation around the primary beam in degree
- outer angle offset, which describes a offset of the outer detector angle from its true zero position

The misalignment parameters as well as the pixel size can be fixed during the fitting.

```
1 """
 2 example script to show the detector parameter determination for area detectors
 3 from images recorded in the primary beam
 5
 6 import os
 7
 8 import xrayutilities as xu
 9
10 en = 10300.0 # eV
11 datadir = os.path.join("data", "wire_") # data path for CCD files
12 # template for the CCD file names
13 filetmp = os.path.join(datadir, "wire_12_%05d.edf.gz")
14
15 # manually selected images
16 # select images which have the primary beam fully on the CCD
17 imagenrs = [2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19,
18
               20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33]
19
20 images = []
21 \text{ ang1} = []
22 \text{ ang} 2 = []
23
24 # read images and angular positions from the data file
25 # this might differ for data taken at different beamlines since
26 # they way how motor positions are stored is not always consistent
27 for imgnr in imagenrs:
28
       filename = filetmp % imgnr
29
       edf = xu.io.EDFFile(filename)
30
      images.append(edf.data)
31
       angl.append(float(edf.header['ESRF_ID01_PSIC_NANO_NU']))
32
       ang2.append(float(edf.header['ESRF_ID01_PSIC_NANO_DEL']))
33
34
35 # call the fit for the detector parameters
36 # detector arm rotations and primary beam direction need to be given.
37 # in total 9 parameters are fitted, however the severl of them can
38 # be fixed. These are the detector tilt azimuth, the detector tilt angle, the
39 # detector rotation around the primary beam and the outer angle offset
40 # The detector pixel size or the detector distance should be kept unfixed to
41 # be optimized by the fit.
42 param, eps = xu.analysis.sample_align.area_detector_calib(
       ang1, ang2, images, ['z+', 'y-'], 'x+',
43
       start=(None, None, 1.0, 45, 0, -0.7, 0),
44
45
      fix=(False, False, True, False, False, False, False),
      wl=xu.en2lam(en))
46
```

A possible output of this script could be

fitted 8.0712e-08 (2,['Parameter param: parameters: epsilon: convergence']) $(cch1, cch2, pwidth1, pwidth2, tiltazimuth, tilt, detrot, outer angle_offset)$ param: 140.07 998.34 4.4545e-05 4.4996e-05 72.0 1.97 -0.792 -1.543 please check the resulting data (consider setting plot=True) detector rotation axis / primary beam direction (given by user): ['z+', 'y-'] / x+ detector pixel directions / distance: z- y+ / 1 detector initialization with: init_area('z-', 'y+', cch1=140.07, cch2=998.34, Nch1=516, pwidth1=4.4545e-05, pwidth2=4.4996e-05, distance=1., detrot=-0.792, tiltazimuth=72.0, tilt=1.543) AND ALWAYS USE an (additional) OFFSET of -1.9741deg in the OUTER DETECTOR ANGLE!

The output gives the fitted detector parameters and compiles the Python code line one needs to use to initialize the detector. Important to note is that the outer angle offset which was determined by the fit (-1.9741 degree in the aboves example) is not included in the initialization of the detector parameters *but* needs to be used in every call to the q-conversion function as offset. This step needs to be performed manually by the user!

Area detector (Variant 2)

In addition to scans in the primary beam this variant enables also the use of detector images recorded in scans at Bragg reflections of a known reference materials. However this also required that the sample orientation and x-ray wavelength need to be fit. To keep the additional parameters as small as possible we only implemented this for symmetric coplanar diffractions.

The advantage of this method is that it is more sensitive to the outer angle offset also at large detector distances. The additional parameters are:

- sample tilt angle in degree
- sample tilt azimuth in degree
- and the x-ray wavelength in Angstrom

```
1 """
 2 example script to show the detector parameter determination for area detectors
 3 from images recorded in the primary beam and at known symmetric coplanar Bragg
 4 reflections of a reference crystal
 5 """
 6
 7 import os
 9 import numpy
10 import xrayutilities as xu
11
12 Si = xu.materials.Si
13
14 datadir = 'data'
15 specfile = "si_align.spec"
16
17 en = 15000 # eV
18 \text{ wl} = \text{xu.en2lam(en)}
19 imgdir = os.path.join(datadir, "si align ") # data path for CCD files
20 filetmp = "si align 12 %04d.edf.gz"
21
22 qconv = xu.QConversion(['z+', 'y-'], ['z+', 'y-'], [1, 0, 0])
23 hxrd = xu.HXRD(Si.Q(1, 1, -2), Si.Q(1, 1, 1), wl=wl, qconv=qconv)
2.4
25 # manually selected images
26
27 s = xu.io.SPECFile(specfile, path=datadir)
28 imagenrs = []
29 for num in [61, 62, 63, 20, 21, 26, 27, 28]:
30
       s[num].ReadData()
       imagenrs = numpy.append(imagenrs, s[num].data['ccd_n'])
31
32
33 # avoid images which do not have to full beam on the detector as well as
```

```
34 # other which show signal due to cosmic radiation
35 avoid_images = [37, 57, 62, 63, 65, 87, 99, 106, 110, 111, 126, 130, 175,
36
                   181, 183, 185, 204, 206, 207, 208, 211, 212, 233, 237, 261,
37
                   275, 290]
38
39 images = []
40 ang1 = [] # outer detector angle
41 ang2 = [] # inner detector angle
42 sang = [] # sample rocking angle
43 hkls = [] # Miller indices of the reference reflections
44
45
46 def hotpixelkill(ccd):
47
48
       function to remove hot pixels from CCD frames
49
       ADD REMOVE VALUES IF NEEDED!
50
51
      ccd[304, 97] = 0
52
      ccd[303, 96] = 0
53
       return ccd
54
55
56 # read images and angular positions from the data file
57 # this might differ for data taken at different beamlines since
58 # they way how motor positions are stored is not always consistent
59 for imgnr in numpy.sort(list(set(imagenrs) - set(avoid images))[::4]):
       filename = os.path.join(imgdir, filetmp % imgnr)
60
61
       edf = xu.io.EDFFile(filename)
62
       ccd = hotpixelkill(edf.data)
63
       images.append(ccd)
64
       angl.append(float(edf.header['motor_pos'].split()[4]))
65
       ang2.append(float(edf.header['motor_pos'].split()[3]))
66
       sang.append(float(edf.header['motor_pos'].split()[1]))
67
       if imgnr > 1293.:
68
           hkls.append((0, 0, 0))
69
       elif imgnr < 139:</pre>
70
           hkls.append((0, 0, numpy.sqrt(27))) # (3,3,3))
71
72
           hkls.append((0, 0, numpy.sqrt(75))) # (5,5,5))
73
74 # call the fit for the detector parameters.
75 # Detector arm rotations and primary beam direction need to be given
76 # in total 8 detector parameters + 2 additional parameters for the reference
77 # crystal orientation and the wavelength are fitted, however the 4 misalignment
78 # parameters of the detector and the 3 other parameters can be fixed.
79 # The fixable parameters are detector tilt azimuth, the detector tilt angle,
80 # the detector rotation around the primary beam, the outer angle offset, sample
81 # tilt, sample tilt azimuth and the x-ray wavelength
82 # Additionally if accurately known the detector pixel size can be given and
83 # fixed and instead the detector distance can be fitted.
84 param, eps = xu.analysis.area_detector_calib_hkl(
85
       sang, ang1, ang2, images, hkls, hxrd, Si, ['z+', 'y-'], 'x+',
86
       start=(None, None, 1.0, 45, 1.69, -0.55, -1.0, 1.3, 60., wl),
87
       fix=(False, False, True, False, False, False, False, False, False, False),
88
       plot=True)
89
90 # Following is an example of the output of the summary of the
91 # area_detector_calib_hkl function
92 # total time needed for fit: 624.51sec
93 # fitted parameters: epsilon: 9.9159e-08 (2,['Parameter convergence'])
```

```
94 # param:
95 # (cch1,cch2,pwidth1,pwidth2,tiltazimuth,tilt,detrot,outerangle_offset,
96 # sampletilt,stazimuth,wavelength)
97 # param: 367.12 349.27 6.8187e-05 6.8405e-05 131.4 2.87 -0.390 -0.061 1.201
98 # 318.44 0.8254
99 # please check the resulting data (consider setting plot=True)
100 # detector rotation axis / primary beam direction (given by user): ['z+', 'y-']
101 # / x+
102 # detector pixel directions / distance: z- y+ / 1
103 # detector initialization with:
104 # init_area('z-', 'y+', cch1=367.12, cch2=349.27, Nch1=516, Nch2=516,
105 # pwidth1=6.8187e-05, pwidth2=6.8405e-05, distance=1., detrot=-0.390,
106 # tiltazimuth=131.4, tilt=2.867)
107 # AND ALWAYS USE an (additional) OFFSET of -0.0611deg in the OUTER
108 # DETECTOR ANGLE!
```

Simulation examples

In the following a few code-snippets are shown which should help you getting started with reflectivity and diffraction simulations using *xrayutilities*. All simulations in *xrayutilities* are for layers systems and currently there are no plans to extend this to other geometries. Note that not all of the codes shown in the following will be run-able as stand-alone scripts. For fully running scripts look in the examples directory in the download found here.

Building Layer stacks for simulations

The basis of all simulations in *xrayutilities* are stacks of layers. Therefore several functions exist to build up such layered systems. The basic building block of all of them is a **Layer** object which takes a material and its thickness in angström as initializing parameter.:

```
import xrayutilities as xu
lay = xu.simpack.Layer(xu.materials.Si, 200)
```

In the shown example a silicon layer with 20 nm thickness is created. The first argument is the material of the layer. For diffraction simulations this needs to be derived from the **Crystal**-class. This means all predefined materials in *xrayutitities* can be used for this purpose. For x-ray reflectivity simulations, however, also knowing the chemical composition and density of the material is sufficient.

A 5 nm thick metallic CoFe compound layer can therefore be defined by:

```
rho_cf = 0.5*8900 + 0.5*7874 # mass density in kg/m^3
mCoFe = xu.materials.Amorphous('CoFe', rho_cf)
lCoFe = xu.simpack.Layer(mat_cf, 50)
```

Note

The Layer object can have several more model dependent properties discussed in detail below.

When several layers are defined they can be combined to a LayerStack which is used for the simulations below.:

```
1 sub = xu.simpack.Layer(xu.materials.Si, float('inf'))
2 lay1 = xu.simpack.Layer(xu.materials.Ge, 200)
3 lay2 = xu.simpack.Layer(xu.materials.SiO2, 30)
4 ls = xu.simpack.LayerStack('Si/Ge', sub, lay1, lay2)
5 # or equivalently
6 ls = xu.simpack.LayerStack('Si/Ge', sub + lay1 + lay2)
```

The last two lines show two different options of creating a stack of layers. As is shown in the last example the substrate thickness can be infinite (see below) and layers can be also stacked by summation. For creation of more complicated superlattice stacks one can further use multiplication:

```
lay1 = xu.simpack.Layer(xu.materials.SiGe(0.3), 50)
lay2 = xu.simpack.Layer(xu.materials.SiGe(0.6), 40)
ls = xu.simpack.LayerStack('Si/SiGe SL', sub + 5*(lay1 + lay2))
```

Pseudomorphic Layers

All stacks of layers described above use the materials in the layer as they are supplied. However, epitaxial systems often adopt the inplane lattice parameter of the layers beneath. To mimic this behavior you can either supply the Layer objects which custom Crystal objects which have the appropriate lattice parameters or use the PseudomorphicStack* classes which to the adaption of the lattice parameters automatically. In this respect the 'relaxation' parameter of the Layer class is important since it allows to create partially/fully relaxed layers.:

```
1 sub = xu.simpack.Layer(xu.materials.Si, float('inf'))
2 buf1 = xu.simpack.Layer(xu.materials.SiGe(0.5), 5000, relaxation=1.0)
3 buf2 = xu.simpack.Layer(xu.materials.SiGe(0.8), 5000, relaxation=1.0)
4 lay1 = xu.simpack.Layer(xu.materials.SiGe(0.6), 50, relaxation=0.0)
5 lay2 = xu.simpack.Layer(xu.materials.SiGe(1.0), 50, relaxation=0.0)
6 # create pseudomorphic superlattice stack
7 pls = xu.simpack.PseudomorphicStack001('SL 5/5', sub+buf1+buf2+5*(lay1+lay2))
```

Note

As indicated by the function name the PseudomorphicStack currently only works for (001) surfaces and cubic materials. Implementations for other surface orientations are planned.

If you would like to check the resulting lattice objects of the different layers you could use:

```
for 1 in pls:
    print(1.material.lattice)
```

Special layer types

So far one special layer mimicking a layer with gradually changing chemical composition is implemented. It consists of several thin sublayers of constant composition. So in order to obtain a smooth grading one has to select enough sublayers. This however has a negativ impact on the performance of all simulation models. A tradeoff needs to found! Below a graded SiGe buffer is shown which consists of 100 sublayers and has total thickness of 1µm.:

Setting up a model

This sectiondescribes the parameters which are common for all diffraction models in *xrayutilties*-simpack. All models need a list of Layers for which the reflected/diffracted signal will be calculated. Further all models have some common parameters which allow scaling and background addition in the model output and contain general information about the calculation which are model-independent. These are

- 'experiment': an **Experiment/HXRD** object which defines the surface geometry of the model. If none is given a default class with (001) surface is generated.
- 'resolution_width': width of the Gaussian resolution function used to convolute with the data. The unit of this parameters depends on the model and can be either in degree or 1/AA.
- '10': is the primary beam flux/intensity
- 'background': is the background added to the simulation after it was scaled by IO

• 'energy': energy in eV used to obtain the optical parameters for the simulation. The energy can alternatively also be supplied via the 'experiment' parameter, however, the 'energy' value overrules this setting. If no energy is given the default energy from the configuration is used.

The mentioned parameters can be supplied to the constructor method of all model classes derived from LayerModel, which applies to all examples mentioned below.:

Reflectivity calculation and fitting

This section shows the calculation and fitting of specular x-ray reflectivity curves as well as the calculation of diffuse x-ray reflectivity curves/maps.

Specular x-ray reflectivity

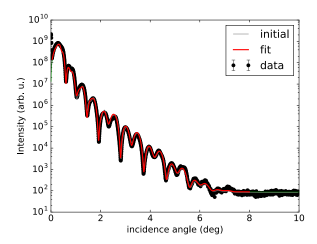
For the specular reflectivity models currently only the Parrat formalism including non-correlated roughnesses is implemented. A minimal working example for a reflectivity calculation follows.:

```
1 # building a stack of layers
2 sub = xu.simpack.Layer(xu.materials.GaAs, float('inf'), roughness=2.0)
3 lay1 = xu.simpack.Layer(xu.materials.AlGaAs(0.25), 75, roughness=2.5)
4 lay2 = xu.simpack.Layer(xu.materials.AlGaAs(0.75), 25, roughness=3.0)
5 pls = xu.simpack.PseudomorphicStack001('pseudo', sub+5*(lay1+lay2))
6
7 # reflectivity calculation
8 m = xu.simpack.SpecularReflectivityModel(pls, sample_width=5, beam_width=0.3)
9 ai = linspace(0, 5, 10000)
10 Ixrr = m.simulate(ai)
```

In addition to the layer thickness also the roughness and density (in kg/m^3) of a Layer can be set since they are important for the reflectivity calculation. This can be done upon definition of the **Layer** or also manipulated at any later stage. Such x-ray reflectivity calculations can also be fitted to experimental data using the **FitModel** class which is shown in detail in the example below (which is also included in the example directory). The fitting is performed using the Imfit Python package which needs to be installed when you want to use this fitting function. This package allows to build complicated models including bounds and correlations between parameters.

```
1 from matplotlib.pylab import *
 2 import xrayutilities as xu
 3 import lmfit
 4 import numpy
 5
 6 # load experimental data
 7 ai, edata, eps = numpy.loadtxt('data/xrr_data.txt'), unpack=True)
 8 ai /= 2.0
 9
10 # define layers
11 # SiO2 / Ru(5) / CoFe(3) / IrMn(3) / AlOx(10)
12 lSiO2 = xu.simpack.Layer(xu.materials.SiO2, inf, roughness=2.5)
13 lRu = xu.simpack.Layer(xu.materials.Ru, 47, roughness=2.8)
14 \text{ rho\_cf} = 0.5*8900 + 0.5*7874
15 mat_cf = xu.materials.Amorphous('CoFe', rho_cf)
16 lCoFe = xu.simpack.Layer(mat_cf, 27, roughness=4.6)
17 lIrMn = xu.simpack.Layer(xu.materials.Ir20Mn80, 21, roughness=3.0)
18 lAl203 = xu.simpack.Layer(xu.materials.Al203, 100, roughness=5.5)
19
20 # create model
21 m = xu.simpack.SpecularReflectivityModel(1SiO2, 1Ru, 1CoFe, 1IrMn, 1Al2O3,
22
                                             energy='CuKa1', resolution_width=0.02,
23
                                             sample_width=6, beam_width=0.25,
24
                                             background=81, I0=6.35e9)
```

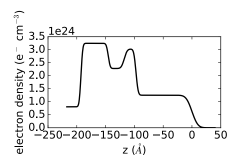
This script can interactively show the fitting progress and after the fitting shows the final plot including the x-ray reflectivity trace of the initial and final parameters.



The picture shows the final plot of the fitting example shown in one of the example scripts.

After building a **SpecularReflectivityModel** is built or fitted the density profile resulting from the thickness and roughness of layers can be plotted easily by:

```
m.densityprofile(500, plot=True) # 500 number of points_
```

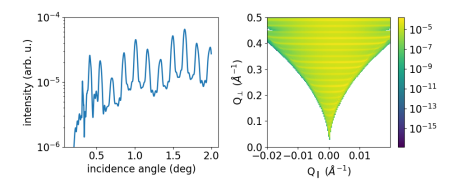


Diffuse reflectivity calculations

For the calculation of diffuse x-ray reflectivity the LayerStack is built equally as shown before. The only difference is that an additional parameter for the lateral correlation length of the roughness can be included: <code>lat_correl</code>. The <code>DiffuseReflectivityModel</code> also takes special parameters which change the vertical correlection length and the way how the diffuse reflectivity is calculated (to be document in more detail). For a Si/Ge superlattice with 5 periods the calculation of the diffuse reflectivity signal at the specular rod is calculated using the <code>simulate()</code> method. A map of the diffuse reflectivity which can be obtained in the coplanar reflection plane can be calculated with the <code>simulate_map()</code> method.

```
1 from matplotlib.pylab import *
 2 import xrayutilities as xu
 3 sub = xu.simpack.Layer(xu.materials.Si, inf, roughness=1, lat_correl=100)
 4 lay1 = xu.simpack.Layer(xu.materials.Si, 200, roughness=1, lat_correl=200)
 5 lay2 = xu.simpack.Layer(xu.materials.Ge, 70, roughness=3, lat_correl=50)
 6 ls = xu.simpack.LayerStack('SL 5', sub+5*(lay2+lay1))
 8 alphai = arange(0.17, 2, 0.001) # for the calculation on the specular rod
 9 qz = arange(0, 0.5, 0.0005) # for the map calculation
10 qL = arange(-0.02, 0.02, 0.0003)
11
12 m = xu.simpack.DiffuseReflectivityModel(ls, sample_width=10, beam_width=1,
13
                                           energy='CuKa1', vert_correl=1000,
14
                                           vert_nu=0, H=1, method=2, vert_int=0)
15 d = m.simulate(alphai)
16 imap = m.simulate_map(qL, qz)
17
18 figure()
19 subplot(121)
20 semilogy(alphai, d, label='diffuse XRR')
21 xlabel('incidence angle (deg)')
22 ylabel('intensity (arb. u.)')
23 ylim(1e-6, 1e-4)
24
25 subplot(122)
26 pcolor(qL, qz, imap.T, norm=mpl.colors.LogNorm())
27 xlabel(r'Q^{\alpha}) = ( \mathrm{AA^{-1}}) 
28 ylabel(r'Q$_\perp$ ($\mathrm{\AA^{-1}}$)')
29 colorbar()
30 tight_layout()
```

The resulting figure shows the simulation result. Currently you have to refer to the docstrings and implementation for further details.



Diffraction calculation

From the very same models as used for XRR calculation one can also perform crystal truncation rod simulations around certain Bragg peaks using various different diffraction models. Depending on the system to model you will have to choose the most apropriate model. Below a short description of the implemented models is given followed by two examples.

Kinematical diffraction models

The most basic models consider only the kinematic diffraction of layers and substrate. Especially the semiinfinite substrate is not well described using the kinematical approximation which results in considerable deviations in close vicinity to substrate Bragg peak with respect to the more acurate dynamical diffraction models.

Such a basic model is employed by:

```
mk = xu.simpack.KinematicalModel(pls, energy=en, resolution_width=0.0001)
Ikin_= mk.simulate(qz,_hkl=(0,_0, 4))
```

A more appealing kinematical model is represented by the **KinematicalMultiBeamModel** class which implements a true multibeam theory is, however, restricted to the use of (001) surfaces and layer thicknesses will be changed to be a multiple of the out of plane lattice spacing. This is necessary since otherwise the structure factor of the unit cell can not be used for the calculation.

It can be employed by:

This model is expected to provide good results especially far away from the substrate peak where the influence of other Bragg peaks on the truncation rod and the variation of the structure factor can not be neglected.

Both kinematical model's **simulate()** method offers two keyword arguments with which basic absorption and refraction correction can be added to the basic models.

Note

The kinematical models can also handle a semi-infinitely thick substrate which results in a diverging intensity at the Bragg peak but provides a basic description of the substrates truncation rod.

Dynamical diffraction models

Acurate description of the diffraction from thin films in close vicinity to the diffraction signal from a bulk substrate is only possible using the dynamical diffraction theory. In **xrayutilities** the dynamical two-beam theory with 4 tiepoints for the calculation of the dispersion surface is implemented. To use this theory you have to supply the **simulate()** method with the incidence angle in degree. Accordingly the 'resolution_width' parameter is also in degree for this model.:

```
md = xu.simpack.DynamicalModel(pls, energy=en, resolution_width=resol)
Idyn = md.simulate(ai, hkl=(0, 0, 4))
```

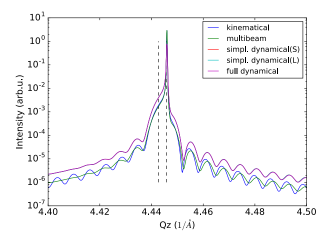
A second simplified dynamical model (SimpleDynamicalCoplanarModel) is also implemented should, however, not be used since its approximations cause mistakes in almost all relevant cases.

The **DynamicalModel** supports the calculation of diffracted signal for 'S' and 'P' polarization geometry. To simulate diffraction data of laboratory sources with Ge(220) monochromator crystal one should use:

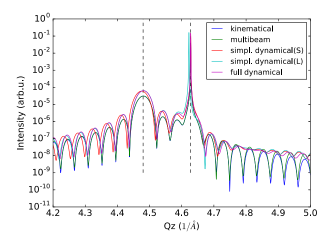
Comparison of diffraction models

Below we show the different implemented models for the case of epitaxial GaAs/AlGaAs and Si/SiGe bilayers. These two cases have very different separation of the layer Bragg peak from the substrate and therefore provide good model system for our models.

We will compare the (004) Bragg peak calculated with different models and but otherwise equal parameters. For scripts used to perform the shown calculation you are referred to the examples directory.



XRD simulations of the (004) Bragg peak of ~100 nm AlGaAs on GaAs(001) using various diffraction models



XRD simulations of the (004) Bragg peak of 15 nm $Si_{0.4}$ $Ge_{0.6}$ on Si(001) using various diffraction models

As can be seen in the images we find that for the AlGaAs system all models except the very basic kinematical model yield an very similar diffraction signal. The second kinematic diffraction model considering the contribution of multiple Bragg peaks on the same truncation rod fails to describe only the ratio of substrate and layer signal, but otherwise results in a very similar line shape as the traces obtained by the dynamic theory.

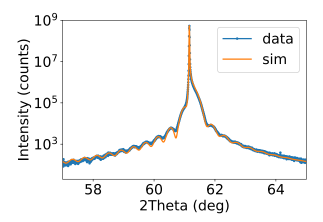
For the SiGe/Si bilayer system bigger differences between the kinematic and dynamic models are found. Further also the difference between the simpler and more sophisticated dynamic model gets obvious further away from the reference position. Interestingly also the multibeam kinematic theory differs considerable from the best dynamic model. As is evident from this second comparison the correct choice of model for the particular system under condideration is crucial for comparison with experimental data.

Fitting of diffraction data

All diffraction models can be embedded into the <code>FitModel</code> class, which is suitable to refine the model parameters. Below (and in the examples directory) a runnable script is shown which shows the fitting for a pseudomorphic InMnAs epilayer on InAs(001). The fitting is performed using the Imfit Python package which needs to be installed when you want to use this fitting function. As one can see below the <code>set_param_hint()</code> function can be used to set up the respective fit parameters including their boundaries and possible correlation with other parameters of the model. It should be equally possible to fit more complex layer structures, however, I expect that one needs to adjust manually the starting parameters to yield something very reasonable. Since this capabilities are rather new please report back any success/problems you have with this via the mailing list.

```
1 import xrayutilities as xu
 2 from matplotlib.pylab import *
 3
 4 # global parameters
 5 wavelength = xu.wavelength('CuKa1')
 6 \text{ offset} = -0.035 # angular offset of the zero position of the data
 7
 8 # set up LayerStack for simulation: InAs(001)/(In,Mn)As(~25 nm)
 9 InAs = xu.materials.InAs
10 InAs.lattice.a = 6.057
11 lInAs = xu.simpack.Layer(InAs, inf)
12 InMnAs = xu.materials.Crystal('InMnAs', xu.materials.SGLattice(
13
       216, 6.050, atoms=('In', 'Mn', 'As'), pos=('4a', '4a', '4c'),
14
       occ=(0.99, 0.01, 1)), cij=InAs.cij)
15 lInMnAs = xu.simpack.Layer(InMnAs, 254, relaxation=0)
16 pstack = xu.simpack.PseudomorphicStack001('list', lInAs, lInMnAs)
17
18 # set up simulation object
19 thetaMono = arcsin(wavelength/(2 * xu.materials.Ge.planeDistance(2, 2, 0)))
20 Cmono = cos(2 * thetaMono)
21 dyn = xu.simpack.DynamicalModel(pstack, I0=1.5e9, background=0,
22
                                    resolution_width=2e-3, polarization='both',
                                    Cmono=Cmono)
23
24 fitmdyn = xu.simpack.FitModel(dyn)
25 fitmdyn.set_param_hint('InMnAs_c', vary=True, min=6.02, max= 6.06)
26 fitmdyn.set_param_hint('InAs_a', vary=True)
27 fitmdyn.set_param_hint('InMnAs_a', expr='InAs_a')
28 fitmdyn.set_param_hint('resolution_width', vary=True)
29 params = fitmdyn.make_params()
30
31 # plot experimental data
32 f = figure(figsize=(7,5))
33 d = xu.io.RASFile('inas_layer_radial_002_004.ras.bz2', path='data'))
34 \text{ scan} = d.\text{scans}[-1]
35 tt = scan.data[scan.scan_axis] - offset
36 semilogy(tt, scan.data['int'], 'o-', ms=3, label='data')
38 # perform fit and plot the result
39 fitmdyn.lmodel.set_hkl((0, 0, 4))
40 ai = (d.scans[-1].data[d.scan.scan_axis] - offset)/2
41 fitr = fitmdyn.fit(d.scans[-1].data['int'], params, ai)
42 print(fitr.fit_report()) # for older lmfit use: lmfit.report_fit(fitr)
```

The resulting figure shows reasonable agreement between the dynamic diffraction simulation and the experimental data.



Powder diffraction simulations

Powder diffraction patterns can be calculated using <code>PowderModel</code>. A specialized class for the definition of powdered materials named <code>Powder</code> exists. The class constructor takes the materials volume and several material parameters specific for the powder material. Among them are <code>crystallite_size_gauss</code> and <code>strain_gauss</code> which can be used to include the effect of finite crystallite size and microstrain. Texture modelled by the March-Dollase pole density offers the <code>preferred_orientation</code> direction parameter as well as a <code>preferred_orientation_factor</code> variable.

The **PowderModel** internally uses **PowderDiffraction** for its calculations which is based on the fundamental parameters approach as implemented and documented here and here.

Several setup specific parameters should be adjusted by a user-specific configuration file or by supplying the appropriate parameters using the *fpsettings* argument of **powderModel**.

If the correct settings are included in the config file the powder diffraction signal of a mixed sample of Co and Fe can be calculated with:

Note that in MS windows you need to encapsulate this code into a dummy function to allow for the multiprocessing module to work correctly. The code then must look like:

```
1 import numpy
 2 import xrayutilities as xu
 3 from multiprocessing import freeze_support
 4
 5 def main():
 6
       tt = numpy.arange(5, 120, 0.01)
 7
       Fe_powder = xu.simpack.Powder(xu.materials.Fe, 1,
 8
                                      crystallite_size_gauss=100e-9)
 9
       Co_powder = xu.simpack.Powder(xu.materials.Co, 5, # 5 times more Co
10
                                      crystallite_size_gauss=200e-9)
11
       pm = xu.simpack.PowderModel(Fe_powder, Co_powder, I0=100)
12
       inte = pm.simulate(tt)
13
       pm.close()
14
15 if __name__ == '__main__':
```

```
16 freeze_support()
17 main()
```

xrayutilities package

Subpackages

xrayutilities.analysis package

Submodules

xrayutilities.analysis.line_cuts module

xrayutilities.analysis.line_cuts.get_arbitrary_line (qpos, intensity, point, vec, npoints, intrange) extracts a line scan from reciprocal space map data along an arbitrary line defined by the point 'point' and propergation vector 'vec'. Integration of the data is performed in a cylindrical volume along the line. This function works for 2D and 3D input data!

Parameters: qpos: list of array-like objects

arrays of x, y (list with two components) or x, y, z (list with three components)

momentum transfers

intensity: array-like

2D or 3D array of reciprocal space intensity with shape equal to the gpos entries

point: tuple, list or array-like

point on the extraction line (2 or 3 coordinates)

vec : tuple, list or array-like

propergation vector of the extraction line (2 or 3 coordinates)

npoints: int

number of points in the output data

intrange : float

radius of the cylindrical integration volume around the extraction line

Returns: qpos, qint : ndarray

line scan coordinates and intensities

used mask: ndarray

mask of used data, shape is the same as the input intensity: True for points which

contributed, False for all others

Examples

```
>>> qcut, qint, mask = get_arbitrary_line([qx, qy, qz], inten, (1.1, 2.2, 0.0), (1, 1, 1), 200, 0.1)
```

xrayutilities.analysis.line_cuts.get_omega_scan (qpos, intensity, cutpos, npoints, intrange, **kwargs) extracts an omega scan from reciprocal space map data with integration along either the 2theta, or radial (omega-2theta) direction. The coplanar diffraction geometry with qy and qz as diffraction plane is assumed. This is consistent with the coplanar geometry implemented in the HXRD-experiment class.

arrays of y, z (list with two components) or x, y, z (list with three components)

momentum transfers

intensity: array-like

2D or 3D array of reciprocal space intensity with shape equal to the qpos entries

cutpos: tuple or list

y/z-position or x/y/z-position at which the line scan should be extracted. this must be

have two entries for 2D data (z-position) and a three for 3D data

npoints: int

number of points in the output data

intrange: float

integration range in along *intdir* in degree. data will be integrated from *-intrange* .. +*intrange*

intdir : {'2theta', 'radial'}, optional

integration direction: '2theta': scattering angle (default), or 'radial': omega-2theta

direction.

wl: float or str, optional

wavelength used to determine angular integration positions

Note:

Although applicable for any set of data, the extraction only makes sense when the data are aligned into the y/z-plane.

Returns: om, omint: ndarray

omega scan coordinates and intensities

used_mask : ndarray

mask of used data, shape is the same as the input intensity: True for points which contributed, False for all others

Examples

```
>>> omcut, omcut_int, mask = get_omega_scan([qy, qz], inten, [2.0, 5.0], 250, intrange=0.1)
```

xrayutilities.analysis.line_cuts.get_qx_scan (qpos, intensity, cutpos, npoints, intrange, **kwargs) extracts a qx scan from 3D reciprocal space map data with integration along either, the perpendicular plane in q-space, omega (sample rocking angle) or 2theta direction. For the integration in angular space (omega, or 2theta) the coplanar diffraction geometry with qy and qz as diffraction plane is assumed. This is consistent with the coplanar geometry implemented in the HXRD-experiment class.

arrays of x, y, z (list with three components) momentum transfers

intensity: array-like

3D array of reciprocal space intensity with shape equal to the qpos entries

cutpos: tuple/list

y/z-position at which the line scan should be extracted. this must be and a tuple/list with the qy, qz cut position

npoints: int

number of points in the output data

intrange: float

integration range in along *intdir*, either in 1/AA (*q*) or degree ('omega', or '2theta'). data will be integrated from *-intrange* .. +*intrange*

intdir: {'q', 'omega', '2theta'}, optional

integration direction: 'q': perpendicular Q-plane (default), 'omega': sample rocking angle, or '2theta': scattering angle.

wl: float or str, optional

wavelength used to determine angular integration positions

Note:

The angular integration directions although applicable for any set of data only makes sense when the data are aligned into the y/z-plane.

Returns: qx, qxint: ndarray

qx scan coordinates and intensities

used_mask : ndarray

mask of used data, shape is the same as the input intensity: True for points which contributed, False for all others

Examples

```
>>> qxcut, qxcut_int, mask = get_qx_scan([qx, qy, qz], inten, [0, 2.0], 250, intrange=0.01)
```

xrayutilities.analysis.line_cuts.get_qy_scan (qpos, intensity, cutpos, npoints, intrange, **kwargs) extracts a qy scan from reciprocal space map data with integration along either, the perpendicular plane in q-space, omega (sample rocking angle) or 2theta direction. For the integration in angular space (omega, or 2theta) the coplanar diffraction geometry with qy and qz as diffraction plane is assumed. This is consistent with the coplanar geometry implemented in the HXRD-experiment class.

arrays of y, z (list with two components) or x, y, z (list with three components)

momentum transfers

intensity: array-like

2D or 3D array of reciprocal space intensity with shape equal to the gpos entries

cutpos: float or tuple/list

x/z-position at which the line scan should be extracted. this must be a float for 2D data

(z-position) and a tuple with two values for 3D data

npoints: int

number of points in the output data

intrange: float

integration range in along *intdir*, either in 1/AA (*q*) or degree ('omega', or '2theta'). data will be integrated from *-intrange* .. +*intrange*

intdir: {'q', 'omega', '2theta'}, optional

integration direction: 'q': perpendicular Q-plane (default), 'omega': sample rocking angle, or '2theta': scattering angle.

wl: float or str, optional

wavelength used to determine angular integration positions

Note:

For 3D data the angular integration directions although applicable for any set of data only makes sense when the data are aligned into the y/z-plane.

Returns: qy, qyint : ndarray

qy scan coordinates and intensities

used_mask : ndarray

mask of used data, shape is the same as the input intensity: True for points which contributed, False for all others

Examples

```
>>> qycut, qycut_int, mask = get_qy_scan([qy, qz], inten, 5.0, 250, intrange=0.02, intdir='2theta')
```

xrayutilities.analysis.line_cuts.get_qz_scan (qpos, intensity, cutpos, npoints, intrange, **kwargs) extracts a qz scan from reciprocal space map data with integration along either, the perpendicular plane in q-space, omega (sample rocking angle) or 2theta direction. For the integration in angular space (omega, or 2theta) the coplanar diffraction geometry with qy and qz as diffraction plane is assumed. This is consistent with the coplanar geometry implemented in the HXRD-experiment class.

arrays of y, z (list with two components) or x, y, z (list with three components)

momentum transfers

intensity: array-like

2D or 3D array of reciprocal space intensity with shape equal to the qpos entries

cutpos: float or tuple/list

x/y-position at which the line scan should be extracted. this must be a float for 2D data

and a tuple with two values for 3D data

npoints: int

number of points in the output data

intrange: float

integration range in along *intdir*, either in 1/AA (*q*) or degree ('omega', or '2theta'). data will be integrated from *-intrange/2* .. +*intrange/2*

intdir: {'q', 'omega', '2theta'}, optional

integration direction: 'q': perpendicular Q-plane (default), 'omega': sample rocking angle, or '2theta': scattering angle.

wl: float or str, optional

wavelength used to determine angular integration positions

Note:

For 3D data the angular integration directions although applicable for any set of data only makes sense when the data are aligned into the y/z-plane.

Returns: qz, qzint : ndarray

qz scan coordinates and intensities

used_mask : ndarray

mask of used data, shape is the same as the input intensity: True for points which contributed, False for all others

Examples

```
>>> qzcut, qzcut_int, mask = get_qz_scan([qy, qz], inten, 3.0, 200, intrange=0.3)
```

xrayutilities.analysis.line_cuts.get_radial_scan (qpos, intensity, cutpos, npoints, intrange,
**kwarqs)

extracts a radial scan from reciprocal space map data with integration along either the omega or 2theta direction. The coplanar diffraction geometry with qy and qz as diffraction plane is assumed. This is consistent with the coplanar geometry implemented in the HXRD-experiment class.

arrays of y, z (list with two components) or x, y, z (list with three components)

momentum transfers

intensity: array-like

2D or 3D array of reciprocal space intensity with shape equal to the qpos entries

cutpos: tuple or list

y/z-position or x/y/z-position at which the line scan should be extracted. this must be

have two entries for 2D data (z-position) and a three for 3D data

npoints: int

number of points in the output data

intrange: float

integration range in along *intdir* in degree. data will be integrated from *-intrange* .. +*intrange*

intdir: {'omega', '2theta'}, optional

integration direction: 'omega': sample rocking angle (default), '2theta': scattering angle

wl: float or str, optional

wavelength used to determine angular integration positions

Note:

Although applicable for any set of data, the extraction only makes sense when the data are aligned into the y/z-plane.

Returns: tt, omttint : ndarray

omega-2theta scan coordinates (2theta values) and intensities

used_mask : ndarray

mask of used data, shape is the same as the input intensity: True for points which contributed, False for all others

Examples

```
>>> ttcut, omtt_int, mask = get_radial_scan([qy, qz], inten, [2.0, 5.0], 250, intrange=0.1)
```

xrayutilities.analysis.line_cuts.get_ttheta_scan (qpos, intensity, cutpos, npoints, intrange,

**kwargs)

extracts a 2theta scan from reciprocal space map data with integration along either the omega or radial direction. The coplanar diffraction geometry with qy and qz as diffraction plane is assumed. This is consistent with the coplanar geometry implemented in the HXRD-experiment class.

arrays of y, z (list with two components) or x, y, z (list with three components)

momentum transfers

intensity: array-like

2D or 3D array of reciprocal space intensity with shape equal to the qpos entries

cutpos: tuple or list

y/z-position or x/y/z-position at which the line scan should be extracted. this must be

have two entries for 2D data (z-position) and a three for 3D data

npoints: int

number of points in the output data

intrange: float

integration range in along intdir in degree. data will be integrated from -intrange ..

+intrange

intdir : {'omega', 'radial'}, optional

integration direction: 'omega': sample rocking angle (default), 'radial': omega-2theta

direction

wl: float or str, optional

wavelength used to determine angular integration positions

Note:

Although applicable for any set of data, the extraction only makes sense when the data are aligned into the y/z-plane.

Returns: tt, ttint: ndarray

2theta scan coordinates and intensities

used_mask : ndarray

mask of used data, shape is the same as the input intensity: True for points which contributed, False for all others

Examples

```
>>> ttcut, tt_int, mask = get_ttheta_scan([qy, qz], inten, [2.0, 5.0], 250, intrange=0.1)
```

xrayutilities.analysis.misc module

miscellaneous functions helpful in the analysis and experiment

xrayutilities.analysis.misc.coplanar_intensity (mat, exp, hkl, thickness, thMono, sample_width=10,
beam width=1)

Calculates the expected intensity of a Bragg peak from an epitaxial thin film measured in coplanar geometry (integration over omega and 2theta in angular space!)

Parameters: mat: Crystal

Crystal instance for structure factor calculation

exp: Experiment

Experimental(HXRD) class for the angle calculation

hkl: list, tuple or array-like

Miller indices of the peak to calculate

thickness: float

film thickness in nm

thMono: float

Bragg angle of the monochromator (deg)

sample_width: float, optional

width of the sample along the beam

beam width: float, optional

width of the beam in the same units as the sample size

Returns: float

intensity of the peak

xrayutilities.analysis.misc.getangles (peak, sur, inp) calculates the chi and phi angles for a given peak

Parameters: peak: list or array-like

hkl for the peak of interest

sur : list or array-like
 hkl of the surface
inp : list or array-like

inplane reference peak or direction

Returns: list

[chi, phi] for the given peak on surface sur with inplane direction inp as reference

Examples

To get the angles for the -224 peak on a 111 surface type

```
>>> [chi, phi] = getangles([-2, 2, 4], [1, 1, 1], [2, 2, 4])
```

xrayutilities.analysis.misc.getunitvector (chi, phi, ndir=(0, 0, 1), idir=(1, 0, 0))

return unit vector determined by spherical angles and definition of the polar axis and inplane reference direction (phi=0)

Parameters: chi, phi : float

spherical angles (polar and azimuthal) in degree

ndir: tuple, list or array-like

polar/z-axis (determines chi=0)

idir: tuple, list or array-like

azimuthal axis (determines phi=0)

xrayutilities.analysis.sample_align module

functions to help with experimental alignment during experiments, especially for experiments with linear and area detectors

xrayutilities.analysis.sample_align.area_detector_calib (angle1, angle2, ccdimages, detaxis, r_i, plot=True, cut_off=0.7, start=(None, None, 1, 0, 0, 0, 0), fix=(False, False, True, False, False, False, False, False), fig=None, wl=None, plotlog=False, nwindow=50, debug=False)

function to calibrate the detector parameters of an area detector it determines the detector tilt possible rotations and offsets in the detector arm angles

Parameters: angle1: array-like

outer detector arm angle

angle2: array-like

inner detector arm angle

ccdimages: array-like

images of the ccd taken at the angles given above

detaxis: list of str

detector arm rotation axis; default: ['z+', 'y-']

r_i : str

primary beam direction [xyz][+-]; default 'x+'

plot: bool, optional

flag to determine if results and intermediate results should be plotted; default: True

cut_off : float, optional

cut off intensity to decide if image is used for the determination or not; default: 0.7 = 70%

start: tuple, optional

sequence of start values of the fit for parameters, which can not be estimated automatically or might want to be fixed. These are: pwidth1, pwidth2, distance, tiltazimuth, tilt, detector_rotation, outerangle_offset. By default (None, None, 1, 0, 0, 0, 0) is used.

fix: tuple of bool

fix parameters of start (default: (False, False, True, False, False, False, False)) It is strongly recommended to either fix the distance or the pwidth1, 2 values.

fig: Figure, optional

matplotlib figure used for plotting the error default: None (creates own figure)

wl: float or str

wavelength of the experiment in Angstrom (default: config.WAVELENGTH) value does not really matter here but does affect the scaling of the error

plotlog: bool

flag to specify if the created error plot should be on log-scale

nwindow: int

window size for determination of the center of mass position after the center of mass of every full image is determined, the center of mass is determined again using a window of size nwindow in order to reduce the effect of hot pixels.

debug: bool

flag to specify that you want to see verbose output and saving of images to show if the CEN determination works

xrayutilities.analysis.sample_align.area_detector_calib_hkl (sampleang, angle1, angle2, ccdimages, hkls, experiment, material, detaxis, r_i, plot=True, cut_off=0.7, start=(None, None, 1, 0, 0, 0, 0, 0, 0, 0); config'), fix=(False, False, True, False, False, False, False, False, False, False), fig=None, plotlog=False, nwindow=50, debug=False)

function to calibrate the detector parameters of an area detector it determines the detector tilt possible rotations and offsets in the detector arm angles

in this variant not only scans through the primary beam but also scans at a set of symmetric reflections can be used for the detector parameter determination. for this not only the detector parameters but in addition the sample orientation and wavelength need to be fit. Both images from the primary beam hkl = (0, 0, 0) and from a symmetric reflection hkl = (h, k, l) need to be given for a successful run.

Parameters: sampleang: array-like

sample rocking angle (needed to align the reflections (same rotation direction as inner detector rotation)) other sample angle are not allowed to be changed during the scans

angle1: array-like

outer detector arm angle

angle2: array-like

inner detector arm angle

ccdimages : array-like

images of the ccd taken at the angles given above

hkls: list or array-like

hkl values for every image

experiment: Experiment

Experiment class object needed to get the UB matrix for the hkl peak treatment

material: Crystal

material used as reference crystal

detaxis: list of str

detector arm rotation axis; default: ['z+', 'y-']

r_i : str

primary beam direction [xyz][+-]; default 'x+'

plot: bool, optional

flag to determine if results and intermediate results should be plotted; default: True

cut off: float, optional

cut off intensity to decide if image is used for the determination or not; default: 0.7 = 70%

start: tuple, optional

sequence of start values of the fit for parameters, which can not be estimated automatically or might want to be fixed. These are: pwidth1, pwidth2, distance, tiltazimuth, tilt, detector_rotation, outerangle_offset, sampletilt, sampletiltazimuth, wavelength. By default (None, None, 1, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0).

fix: tuple of bool

fix parameters of start (default: (False, False, True, False, False, False, False, False, False, False, False, False)) It is strongly recommended to either fix the distance or the pwidth1, 2 values.

fig: Figure, optional

matplotlib figure used for plotting the error default: None (creates own figure)

plotlog: bool

flag to specify if the created error plot should be on log-scale

nwindow: int

window size for determination of the center of mass position after the center of mass of every full image is determined, the center of mass is determined again using a window of size nwindow in order to reduce the effect of hot pixels.

debug: bool

flag to specify that you want to see verbose output and saving of images to show if the CEN determination works

xrayutilities.analysis.sample_align.fit_bragg_peak (om, tt, psd, omalign, ttalign, exphxrd, frange=(0.03, 0.03), peaktype='Gauss', plot=True)

helper function to determine the Bragg peak position in a reciprocal space map used to obtain the position needed for correction of the data. the determination is done by fitting a two dimensional Gaussian (xrayutilities.math.Gauss2d) or Lorentzian (xrayutilities.math.Lorentz2d)

PLEASE ALWAYS CHECK THE RESULT CAREFULLY!

Parameters: om, tt: array-like

angular coordinates of the measurement either with size of psd or of psd.shape[0]

psd: array-like

intensity values needed for fitting

omalign: float

aligned omega value, used as first guess in the fit

ttalign: float

aligned two theta values used as first guess in the fit these values are also used to set the range for the fit: the peak should be within +/-frangeAA^\-1} of those values

exphxrd: Experiment

experiment class used for the conversion between angular and reciprocal space.

frange: tuple of float, optional

data range used for the fit in both directions (see above for details default:(0.03, 0.03) unit: AA\(\frac{1}{}\)

peaktype : {'Gauss', 'Lorentz'}

peak type to fit plot: bool, optional

if True (default) function will plot the result of the fit in comparison with the measurement.

Returns: omfit, ttfit: float

fitted angular values

params: list

fit parameters (of the Gaussian/Lorentzian)

covariance: ndarray

covariance matrix of the fit parameters

xrayutilities.analysis.sample_align.linear_detector_calib (angle, mca_spectra, **keyargs) function to calibrate the detector distance/channel per degrees for a straight linear detector mounted on a detector arm

Parameters: angle: array-like

array of angles in degree of measured detector spectra

mca_spectra: array-like

corresponding detector spectra (shape: (len(angle), Nchannels)

r_i : str, optional

primary beam direction as vector [xyz][+-]; default: 'y+'

detaxis: str, optional

detector arm rotation axis [xyz][+-]; default: 'x+'

Returns: pixelwidth: float

width of the pixel at one meter distance, pixelwidth is negative in case the hit channel number decreases upon an increase of the detector angle

center_channel: float

central channel of the detector

detector_tilt: float, optional

if usetilt=True the fitted tilt of the detector is also returned

Note

L/pixelwidth*pi/180 ~= channel/degree, with the sample detector distance L

The function also prints out how a linear detector can be initialized using

the results obtained from this calibration. Carefully check the results

Other plot: bool

Parameters: flag to specify if a visualization of the fit should be done

usetilt: bool

whether to use model considering a detector tilt, i.e. deviation angle of the pixel direction from orthogonal to the primary beam (default: True)

Seealso

psd_chdeg

low level function with more configurable options

xrayutilities.analysis.sample_align.miscut_calc (phi, aomega, zeros=None, omega0=None,
plot=True)

function to calculate the miscut direction and miscut angle of a sample by fitting a sinusoidal function to the variation of the aligned omega values of more than two reflections. The function can also be used to fit reflectivity alignment values in various azimuths.

Parameters: phi : list, tuple or array-like

azimuths in which the reflection was aligned (deg)

aomega : list, tuple or array-like aligned omega values (deg)

zeros: list, tuple or array-like, optional

angles at which surface is parallel to the beam (deg). For the analysis the angles (aomega - zeros) are used.

omega0: float, optional

if specified the nominal value of the reflection is not included as fit parameter, but is fixed to the specified value. This value is MANDATORY if ONLY TWO AZIMUTHS are given.

plot: bool, optional

flag to specify if a visualization of the fit is wanted. default: True

Returns: omega0 : float

the omega value of the reflection should be close to the nominal one

phi0: float

the azimuth in which the primary beam looks upstairs

miscut: float

amplitude of the sinusoidal variation == miscut angle

xrayutilities.analysis.sample_align.**psd_chdeg** (angles, channels, stdev=None, usetilt=True, plot=True, datap='xk', modelline='--r', modeltilt='-b', fignum=None, mlabel='fit', mtiltlabel='fit w/tilt', dlabel='data', figtitle=True)

function to determine the channels per degree using a linear fit of the function nchannel = center_ch+chdeg*tan(angles) or the equivalent including a detector tilt

Parameters: angles: array-like

detector angles for which the position of the beam was measured

channels: array-like

detector channels where the beam was found

stdev : array-like, optional

standard deviation of the beam position

plot: bool, optional

flag to specify if a visualization of the fit should be done

usetilt: bool, optional

whether to use model considering a detector tilt, i.e. deviation angle of the pixel

direction from orthogonal to the primary beam (default: True)

Returns: pixelwidth: float

the width of one detector channel @ 1m distance, which is negative in case the hit

channel number decreases upon an increase of the detector angle.

centerch: float

center channel of the detector

tilt: float

tilt of the detector from perpendicular to the beam (will be zero in case of usetilt=False)

Note

L/pixelwidth*pi/180 = channel/degree for large detector distance with the sample detector disctance L

Other datap: str, optional

Parameters: plot format of data points

modelline: str, optional
plot format of modelline
modeltilt: str, optional
plot format of modeltilt
fignum: int or str, optional

figure number to use for the plot

mlabel: str

label of the model w/o tilt to be used in the plot

mtiltlabel: str

label of the model with tilt to be used in the plot

dlabel: str

label of the data line to be used in the plot

figtitle: bool

flag to tell if the figure title should show the fit parameters

xrayutilities.analysis.sample_align.psd_refl_align (primarybeam, angles, channels, plot=True) function which calculates the angle at which the sample is parallel to the beam from various angles and detector channels from the reflected beam. The function can be used during the half beam alignment with a linear detector.

Parameters: primarybeam: int

primary beam channel number

angles: *list or array-like* incidence angles

channels: list or array-like

corresponding detector channels

plot: bool, optional

flag to specify if a visualization of the fit is wanted default : True

Returns: float

angle at which the sample is parallel to the beam

Examples

```
>>> psd_refl_align(500, [0, 0.1, 0.2, 0.3], [550, 600, 640, 700])
```

Module contents

xrayutilities.analysis is a package for assisting with the analysis of x-ray diffraction data, mainly reciprocal space maps

Routines for obtaining line cuts from gridded reciprocal space maps are offered, with the ability to integrate the intensity perpendicular to the line cut direction.

xrayutilities.io package

Submodules

xrayutilities.io.cbf module

class xrayutilities.io.cbf.CBFDirectory (datapath, ext='cbf', **keyargs)

Bases: xrayutilities.io.filedir.FileDirectory

Parses a directory for CBF files, which can be stored to a HDF5 file for further usage

class xrayutilities.io.cbf.CBFFile (fname, nxkey='X-Binary-Size-Fastest-Dimension',
nykey='X-Binary-Size-Second-Dimension', dtkey='DataType', path=None)

Bases: object

ReadData ()

Read the CCD data into the .data object this function is called by the initialization

```
Save2HDF5 (h5f, group='/', comp=True)
```

Saves the data stored in the EDF file in a HDF5 file as a HDF5 array. By default the data is stored in the root group of the HDF5 file - this can be changed by passing the name of a target group or a path to the target group via the "group" keyword argument.

Parameters: h5f : file-handle or str

a HDF5 file object or name

group: str, optional

group where to store the data (default to the root of the file)

comp: bool, optional

activate compression - true by default

xrayutilities.io.desy_tty08 module

class for reading data + header information from tty08 data files

tty08 is a system used at beamline P08 at Hasylab Hamburg and creates simple ASCII files to save the data. Information is easily read from the multicolumn data file. the functions below enable also to parse the information of the header

xrayutilities.io.desy_tty08.gettty08_scan (scanname, scannumbers, *args, **keyargs)

function to obtain the angular cooridinates as well as intensity values saved in TTY08 datafiles. Especially usefull for reciprocal space map measurements, and to combine date from several scans

further more it is possible to obtain even more positions from the data file if more than two string arguments with its names are given

Parameters: scanname: str

name of the scans, for multiple scans this needs to be a template string

scannumbers : int, tuple or list

number of the scans of the reciprocal space map

args: str, optional

names of the motors. to read reciprocal space maps measured in coplanar diffraction give:

- omname: the name of the omega motor (or its equivalent)
- ttname: the name of the two theta motor (or its equivalent)

keyargs: dict, optional

keyword arguments are passed on to tty08File

Returns: [ang1, ang2, ...]: list, optional

angular positions of the center channel of the position sensitive detector (numpy.ndarray 1D), omitted if no *arg*s are given

MAP: ndarray

All the data values as stored in the data file (includes the intensities e.g. MAP['MCA']).

Examples

```
>>> [om, tt], MAP = xu.io.gettty08_scan('text%05d.dat', 36, 'omega', 'gamma')
```

class xrayutilities.io.desy_tty08.tty08File (filename, path=None, mcadir=None)

Bases: object

Represents a tty08 data file. The file is read during the Constructor call. This class should work for data stored at beamline P08 using the tty08 acquisition system.

Parameters: filename: str

tty08-filename **mcadir**: str, optional

directory name of MCA files

Read ()

Read the data from the file

ReadMCA ()

xrayutilities.io.edf module

```
class xrayutilities.io.edf. EDFDirectory (datapath, ext='edf', **keyargs)
```

Bases: xrayutilities.io.filedir.FileDirectory

Parses a directory for EDF files, which can be stored to a HDF5 file for further usage

class xrayutilities.io.edf.EDFFile (fname, nxkey='Dim_1', nykey='Dim_2', dtkey='DataType', path='',
header=True, keep_open=False)

Bases: object

Parse ()

Parse file to find the number of entries and read the respective header information

ReadData (nimg=0)

Read the CCD data of the specified image and return the data this function is called automatically when the 'data' property is accessed, but can also be called manually when only a certain image from the file is needed.

Parameters: nimg: int, optional

number of the image which should be read (starts with 0)

Save2HDF5 (h5f, group='/', comp=True)

Saves the data stored in the EDF file in a HDF5 file as a HDF5 array. By default the data is stored in the root group of the HDF5 file - this can be changed by passing the name of a target group or a path to the target group via the "group" keyword argument.

Parameters: h5f: file-handle or str

a HDF5 file object or name

group: str, optional

group where to store the data (default to the root of the file)

comp: bool, optional

activate compression - true by default

property data

xrayutilities.io.fastscan module

modules to help with the analysis of FastScan data acquired at the ESRF. FastScan data are X-ray data (various detectors possible) acquired during scanning the sample in real space with a Piezo Scanner. The same functions might be used to analze traditional SPEC mesh scans.

The module provides three core classes:

- FastScan
- FastScanCCD
- FastScanSeries

where the first two are able to parse single mesh/FastScans when one is interested in data of a single channel detector or are detector and the last one is able to parse full series of such mesh scans with either type of detector

see examples/xrayutilities_kmap_ESRF.py for an example script

```
class xrayutilities.io.fastscan.FastScan (filename, scannr, xmotor='adcX', ymotor='adcY', path=")
Bases: object
```

class to help parsing and treating fast scan data. FastScan is the aquisition of X-ray data while scanning the sample with piezo stages in real space. It's is available at several beamlines at the ESRF synchrotron light-source.

```
grid2D (nx, ny, **kwargs)
```

function to grid the counter data and return the gridded X, Y and Intensity values.

Parameters: nx, ny : int

number of bins in x, and y direction

counter: str, optional

name of the counter to use for gridding (default: 'mpx4int' (ID01))

gridrange: tuple, optional

range for the gridder: format: ((xmin, xmax), (ymin, ymax))

Returns: Gridder2D

Gridder2D object with X, Y, data on regular x, y-grid

motorposition (motorname)

read the position of motor with name given by motorname from the data file. In case the motor is included in the data columns the returned object is an array with all the values from the file (although retrace clean is respected if already performed). In the case the motor is not moved during the scan only one value is returned.

Parameters: motorname: str

name of the motor for which the position is wanted

Returns: ndarray

motor position(s) of motor with name motorname during the scan

parse ()

parse the specifie for the scan number specified in the constructor and store the needed informations in the object properties

retrace_clean()

function to clean the data of the scan from retrace artifacts created by the zig-zag scanning motion of the piezo actuators the function cleans the xvalues, yvalues and data attribute of the FastScan object.

class xrayutilities.io.fastscan.FastScanCCD (*args, **kwargs)

Bases: xrayutilities.io.fastscan.FastScan

class to help parsing and treating fast scan data including CCD frames. FastScan is the aquisition of X-ray data while scanning the sample with piezo stages in real space. It's is available at several beamlines at the ESRF synchrotron light-source. During such fast scan at every grid point CCD frames are recorded and need to be analyzed

getCCD (ccdnr, roi=None, datadir=None, keepdir=0, replacedir=None, nav=[1, 1], filterfunc=None)

function to read the ccd files and return the raw X, Y and DATA values. DATA represents a 3D object with first dimension representing the data point index and the remaining two dimensions representing detector channels

Parameters: ccdnr: array-like or str

array with ccd file numbers of length length(FastScanCCD.data) OR a string with the data column name for the file ccd-numbers

roi: tuple, optional

region of interest on the 2D detector. should be a list of lower and upper bounds of detector channels for the two pixel directions (default: None)

datadir: str, optional

the CCD filenames are usually parsed from the SPEC file. With this option the directory used for the data can be overwritten. Specify the datadir as simple string. Alternatively the innermost directory structure can be automatically taken from the specfile. If this is needed specify the number of directories which should be kept using the keepdir option.

keepdir: int, optional

number of directories which should be taken from the SPEC file. (default: 0)

replacedir: int, optional

number of outer most directory names which should be replaced in the output (default = None). One can either give keepdir, or replacedir, with replace taking preference if both are given.

nav: tuple or list, optional

number of detector pixel which will be averaged together (reduces the date size)

filterfunc: callable

function applied to the CCD-frames before any processing. this function should take a single argument which is the ccddata which need to be returned with the same shape! e.g. remove hot pixels, flat/darkfield correction

Returns: X, Y: ndarray

x, y-array (1D)

DATA : ndarray

3-dimensional data object

getccdFileTemplate (specscan, datadir=None, keepdir=0, replacedir=None)

function to extract the CCD file template string from the comment in the SPEC-file scan-header.

Parameters: specscan: SpecScan

spec-scan object from which header the CCD directory should be extracted

datadir: str, optional

the CCD filenames are usually parsed from the scan object. With this option the directory used for the data can be overwritten. Specify the datadir as simple string. Alternatively the innermost directory structure can be automatically taken from the specifile. If this is needed specify the number of directories which should be kept using the keepdir option.

keepdir: int, optional

number of directories which should be taken from the specscan. (default: 0)

replacedir: int, optional

number of outer most directory names which should be replaced in the output (default = None). One can either give keepdir, or replacedir, with replace taking preference if both are given.

Returns: fmtstr: str

format string for the CCD file name using one number to build the real file name

filenr: int

starting file number

gridCCD (nx, ny, ccdnr, roi=None, datadir=None, keepdir=0, replacedir=None, nav=[1, 1], gridrange=None, filterfunc=None)

function to grid the internal data and ccd files and return the gridded X, Y and DATA values. DATA represents a 4D object with first two dimensions representing X, Y and the remaining two dimensions representing detector channels

Parameters: nx, ny : int

number of bins in x, and y direction

ccdnr: array-like or str

array with ccd file numbers of length length(FastScanCCD.data) OR a string with the data column name for the file ccd-numbers

roi: tuple, optional

region of interest on the 2D detector. should be a list of lower and upper bounds of detector channels for the two pixel directions (default: None)

datadir: str, optional

the CCD filenames are usually parsed from the SPEC file. With this option the directory used for the data can be overwritten. Specify the datadir as simple string. Alternatively the innermost directory structure can be automatically taken from the specifile. If this is needed specify the number of directories which should be kept using the keepdir option.

keepdir: int, optional

number of directories which should be taken from the SPEC file. (default: 0)

replacedir: int, optional

number of outer most directory names which should be replaced in the output (default = None). One can either give keepdir, or replacedir, with replace taking preference if both are given.

nav: tuple or list, optional

number of detector pixel which will be averaged together (reduces the date size)

gridrange: tuple

range for the gridder: format: ((xmin, xmax), (ymin, ymax))

filterfunc: callable

function applied to the CCD-frames before any processing. this function should take a single argument which is the ccddata which need to be returned with the same shape! e.g. remove hot pixels, flat/darkfield correction

Returns: X, Y: ndarray

regular x, y-grid

DATA: ndarray

4-dimensional data object

processCCD (ccdnr, roi, datadir=None, keepdir=0, replacedir=None, filterfunc=None)

function to read a region of interest (ROI) from the ccd files and return the raw X, Y and intensity from ROI.

Parameters: ccdnr: array-like or str

array with ccd file numbers of length length(FastScanCCD.data) OR a string with the data column name for the file ccd-numbers

roi: tuple or list

region of interest on the 2D detector. Either a list of lower and upper bounds of detector channels for the two pixel directions as tuple or a list of mask arrays

datadir: str, optional

the CCD filenames are usually parsed from the SPEC file. With this option the directory used for the data can be overwritten. Specify the datadir as simple string. Alternatively the innermost directory structure can be automatically taken from the specifile. If this is needed specify the number of directories which should be kept using the keepdir option.

keepdir: int, optional

number of directories which should be taken from the SPEC file. (default: 0)

replacedir: int, optional

number of outer most directory names which should be replaced in the output (default = None). One can either give keepdir, or replacedir, with replace taking preference if both are given.

filterfunc: callable, optional

function applied to the CCD-frames before any processing. this function should take a single argument which is the ccddata which need to be returned with the same shape! e.g. remove hot pixels, flat/darkfield correction

Returns: X, Y, DATA: ndarray

x, y-array (1D) as well as 1-dimensional data object

class xrayutilities.io.fastscan.FastScanSeries (filenames, scannrs, nx, ny, *args, **kwargs)

Bases: object

class to help parsing and treating a series of fast scan data including CCD frames. FastScan is the aquisition of X-ray data while scanning the sample with piezo stages in real space. It's is available at several beamlines at the ESRF synchrotron light-source. During such fast scan at every grid point CCD frames are recorded and need to be analyzed.

For the series of FastScans we assume that they are measured at different goniometer angles and therefore transform the data to reciprocal space.

align (deltax, deltay)

Since a sample drift or shift due to rotation often occurs between different FastScans it should be corrected before combining them. Since determining such a shift is not straight-forward in general the user needs to supply the routine with the shifts in order correct the x, y-values for the different FastScans. Such a routine could for example use the integrated CCD intensities and determine the shift using a cross-convolution.

Parameters: deltax, deltay: list

list of shifts in x/y-direction for every FastScan in the data structure

getCCDFrames (posx, posy, typ='real')

function to determine the list of ccd-frame numbers for a specific real space position. The real space position must be within the data limits of the FastScanSeries otherwise an ValueError is thrown

Parameters: posx: float

real space x-position or index in x direction

posy: float

real space y-position or index in y direction

typ: {'real', 'index'}, optional

type of coordinates. specifies if the position is specified as real space coordinate or as

index. (default: 'real')

Returns: list

[[motorpos1, ccdnrs1], [motorpos2, ccdnrs2], ...] where motorposN is from the N-ths FastScan in the series and ccdnrsN is the list of according CCD-frames

get_average_RSM (qnx, qny, qnz, qconv, datadir=None, keepdir=0, replacedir=None, roi=None, nav=(1, 1), filterfunc=None)

function to return the reciprocal space map data averaged over all x, y positions from a series of FastScan measurements. It necessary to give the QConversion-object to be used for the reciprocal space conversion. The QConversion-object is expected to have the 'area' conversion routines configured properly. This function needs to read all detector images, so be prepared to lean back for a moment!

Parameters: qnx, qny, qnz: int

number of points used for the 3D Gridder

qconv: QConversion

QConversion-object to be used for the conversion of the CCD-data to reciprocal

space

roi: tuple, optional

region of interest on the 2D detector. should be a list of lower and upper bounds of detector channels for the two pixel directions (default: None)

nav: tuple or list, optional

number of detector pixel which will be averaged together (reduces the date size)

filterfunc : callable, optional

function applied to the CCD-frames before any processing. this function should take a single argument which is the ccddata which need to be returned with the same shape! e.g. remove hot pixels, flat/darkfield correction

datadir: str, optional

the CCD filenames are usually parsed from the SPEC file. With this option the directory used for the data can be overwritten. Specify the datadir as simple string. Alternatively the innermost directory structure can be automatically taken from the specfile. If this is needed specify the number of directories which should be kept/replaced using the keepdir/replacedir option.

keepdir: int, optional

number of directories which should be taken from the SPEC file. (default: 0)

replacedir: int, optional

number of outer most directory names which should be replaced in the output (default = None). One can either give keepdir, or replacedir, with replace taking preference if both are given.

Returns: Gridder3D

gridded reciprocal space map

get_sxrd_for_qrange (qrange, qconv, datadir=None, keepdir=0, replacedir=None, roi=None, nav=(1, 1),
filterfunc=None)

function to return the real space data averaged over a certain q-range from a series of FastScan measurements. It necessary to give the QConversion-object to be used for the reciprocal space conversion. The QConversion-object is expected to have the 'area' conversion routines configured properly.

Note

This function assumes that all FastScans were performed in the same real space positions, no gridding or aligning is performed!

Parameters: grange: list or tuple

q-limits defining a box in reciprocal space. six values are needed: [minx, maxx, miny,

..., maxz]

qconv: QConversion

QConversion object to be used for the conversion of the CCD-data to reciprocal

space

roi: tuple, optional

region of interest on the 2D detector. should be a list of lower and upper bounds of

detector channels for the two pixel directions (default: None)

nav: tuple or list, optional

number of detector pixel which will be averaged together (reduces the date size)

filterfunc : callable, optional

function applied to the CCD-frames before any processing. this function should take a single argument which is the ccddata which need to be returned with the same shape!

e.g. remove hot pixels, flat/darkfield correction

datadir: str, optional

the CCD filenames are usually parsed from the SPEC file. With this option the directory used for the data can be overwritten. Specify the datadir as simple string. Alternatively the innermost directory structure can be automatically taken from the specfile. If this is needed specify the number of directories which should be kept/replaced using the keepdir/replacedir option.

keepdir: int, optional

number of directories which should be taken from the SPEC file. (default: 0)

replacedir: int, optional

number of outer most directory names which should be replaced in the output (default = None). One can either give keepdir, or replacedir, with replace taking preference if

both are given.

Returns: xvalues, yvalues, data: ndarray

x, y, and data values

grid2Dall (nx, ny, **kwargs)

function to grid the counter data and return the gridded X, Y and Intensity values from all the FastScanSeries.

Parameters: nx, ny : int

number of bins in x, and y direction

counter: str, optional

name of the counter to use for gridding (default: 'mpx4int' (ID01))

gridrange: tuple, optional

range for the gridder: format: ((xmin, xmax), (ymin, ymax))

Returns: Gridder2D

object with X, Y, data on regular x, y-grid

gridRSM (posx, posy, qnx, qny, qnz, qconv, roi=None, nav=[1, 1], typ='real', filterfunc=None, **kwargs)

function to calculate the reciprocal space map at a certain x, y-position from a series of FastScan measurements it is necessary to specify the number of grid-oints for the reciprocal space map and the QConversion-object to be used for the reciprocal space conversion. The QConversion-object is expected to have the 'area' conversion routines configured properly.

Parameters: posx: float

real space x-position or index in x direction

posy: float

real space y-position or index in y direction

qnx, qny, qnz: int

number of points in the Qx, Qy, Qz direction of the gridded reciprocal space map

qconv: QConversion

QConversion-object to be used for the conversion of the CCD-data to reciprocal

space

roi: tuple, optional

region of interest on the 2D detector. should be a list of lower and upper bounds of detector channels for the two pixel directions (default: None)

nav: tuple or list, optional

number of detector pixel which will be averaged together (reduces the date size)

typ: {'real', 'index'}, optional

type of coordinates. specifies if the position is specified as real space coordinate or as index. (default: 'real')

filterfunc: callable, optional

function applied to the CCD-frames before any processing. this function should take a single argument which is the ccddata which need to be returned with the same shape! e.g. remove hot pixels, flat/darkfield correction

UB: ndarray

sample orientation matrix

Returns: Gridder3D

object with gridded reciprocal space map

rawRSM (posx, posy, qconv, roi=None, nav=[1, 1], typ='real', datadir=None, keepdir=0, replacedir=None, filterfunc=None, **kwargs)

function to return the reciprocal space map data at a certain x, y-position from a series of FastScan measurements. It necessary to give the QConversion-object to be used for the reciprocal space conversion. The QConversion-object is expected to have the 'area' conversion routines configured properly.

Parameters: posx: float

real space x-position or index in x direction

posy: float

real space y-position or index in y direction

qconv: QConversion

QConversion-object to be used for the conversion of the CCD-data to reciprocal

space

roi: tuple, optional

region of interest on the 2D detector. should be a list of lower and upper bounds of detector channels for the two pixel directions (default: None)

nav: tuple or list, optional

number of detector pixel which will be averaged together (reduces the date size)

typ: {'real', 'index'}, optional

type of coordinates. specifies if the position is specified as real space coordinate or as index. (default: 'real')

filterfunc : callable, optional

function applied to the CCD-frames before any processing. this function should take a single argument which is the ccddata which need to be returned with the same shape! e.g. remove hot pixels, flat/darkfield correction

UB: array-like, optional

sample orientation matrix

datadir: str, optional

the CCD filenames are usually parsed from the SPEC file. With this option the directory used for the data can be overwritten. Specify the datadir as simple string. Alternatively the innermost directory structure can be automatically taken from the specifile. If this is needed specify the number of directories which should be kept using the keepdir option.

keepdir: int, optional

number of directories which should be taken from the SPEC file. (default: 0)

replacedir: int, optional

number of outer most directory names which should be replaced in the output (default = None). One can either give keepdir, or replacedir, with replace taking preference if both are given.

Returns: qx, qy, qz : ndarray

reciprocal space positions of the reciprocal space map

ccddata: ndarray

raw data of the reciprocal space map

valuelist : ndarray

valuelist containing the ccdframe numbers and corresponding motor positions

read_motors()

read motor values from the series of fast scans

retrace_clean()

perform retrace clean for every FastScan in the series

xrayutilities.io.filedir module

class xrayutilities.io.filedir.FileDirectory (datapath, ext, parser, **keyargs)

Bases: object

Parses a directory for files, which can be stored to a HDF5 file for further usage. The file parser is given to the constructor and must provide a Save2HDF5 method.

```
Save2HDF5 (h5f, group=", comp=True)
```

Saves the data stored in the found files in the specified directory in a HDF5 file as a HDF5 arrays in a subgroup. By default the data is stored in a group given by the foldername - this can be changed by passing the name of a target group or a path to the target group via the "group" keyword argument.

Parameters: h5f: file-handle or str

a HDF5 file object or name

group: str, optional

group where to store the data (defaults to pathname if group is empty string)

comp: bool, optional

activate compression - true by default

xrayutilities.io.helper module

convenience functions to open files for various data file reader

these functions should be used in new parsers since they transparently allow to open gzipped and bzipped files

class xrayutilities.io.helper.xu_h5open (f, mode='r')

Bases: object

helper object to decide if a HDF5 file has to be opened/closed when using with a 'with' statement.

xrayutilities.io.helper.xu_open (filename, mode='rb')

function to open a file no matter if zipped or not. Files with extension '.gz', '.bz2', and '.xz' are assumed to be compressed and transparently opened to read like usual files.

Parameters: filename: str

filename of the file to open (full including path)

mode: str, optional

mode in which the file should be opened

Returns: file-handle

handle of the opened file

Raises: IOError

If the file does not exist an IOError is raised by the open routine, which is not caught

within the function

xrayutilities.io.ill_numor module

module for reading ILL data files (station D23): numor files

class xrayutilities.io.ill_numor.numorFile (filename, path=None)

Bases: object

Represents a ILL data file (numor). The file is read during the Constructor call. This class should work for created at station D23 using the mad acquisition system.

Parameters: filename: str

a string with the name of the data file

Read ()

Read the data from the file

columns = {0: ('detector', 'monitor', 'time', 'gamma', 'omega', 'psi'), 1: ('detector', 'monitor', 'time', 'gamma'), 2: ('detector', 'monitor', 'time', 'omega'), 5: ('detector', 'monitor', 'time', 'psi')}

getline (fid)

ssplit (string)

multispace split. splits string at two or more spaces after stripping it.

```
xrayutilities.io.ill_numor.numor_scan (scannumbers, *args, **kwargs)
```

function to obtain the angular cooridinates as well as intensity values saved in numor datafiles. Especially useful for combining several scans into one data object.

Parameters: scannumbers: int or str or iterable

number of the numors, or list of numbers. This will be transformed to a string and used

as a filename args: str, optional

names of the motors e.g.: 'omega', 'gamma'

kwargs: dict

keyword arguments are passed on to numorFile. e.g. 'path' for the files directory

Returns: [ang1, ang2, ...] : *list*

angular positions list, omitted if no args are given

data: ndarray

all the data values.

Examples

```
>>> [om, gam], data = xu.io.numor_scan(414363, 'omega', 'gamma')
```

xrayutilities.io.imagereader module

class xrayutilities.io.imagereader.ImageReader (nop1, nop2, hdrlen=0, flatfield=None,
darkfield=None, dtype=<class 'numpy.int16'>, byte_swap=False)

Bases: object

parse CCD frames in the form of tiffs or binary data (*.bin) to numpy arrays. ignore the header since it seems to contain no useful data

The routine was tested so far with

- 1. RoperScientific files with 4096x4096 pixels created at Hasylab Hamburg, which save an 16bit integer per point.
- 2. Perkin Elmer images created at Hasylab Hamburg with 2048x2048 pixels.

readImage (filename, path=None)

read image file and correct for dark- and flatfield in case the necessary data are available. returned data = ((image data)-(darkfield))/flatfield*average(flatfield)

Parameters: filename: str

filename of the image to be read. so far only single filenames are supported. The data might be compressed. supported extensions: .tif, .bin and .bin.xz

path : str, optional
 path of the data files

```
class xrayutilities.io.imagereader.PerkinElmer (**keyargs)
```

Bases: xrayutilities.io.imagereader.ImageReader

parse PerkinElmer CCD frames (*.tif) to numpy arrays Ignore the header since it seems to contain no useful data

The routine was tested only for files with 2048x2048 pixel images created at Hasylab Hamburg which save an 32bit float per point.

```
class xrayutilities.io.imagereader.Pilatus100K (**keyargs)
```

Bases: xrayutilities.io.imagereader.ImageReader

parse Dectris Pilatus 100k frames (*.tiff) to numpy arrays Ignore the header since it seems to contain no useful data

```
class xrayutilities.io.imagereader.RoperCCD (**keyargs)
Bases: xrayutilities.io.imagereader.ImageReader
```

parse RoperScientific CCD frames (*.bin) to numpy arrays Ignore the header since it seems to contain no useful data

The routine was tested only for files with 4096x4096 pixel images created at Hasylab Hamburg which save an 16bit integer per point.

class xrayutilities.io.imagereader.TIFFRead (filename, path=None)

Bases: xrayutilities.io.imagereader.ImageReader

class to Parse a TIFF file including extraction of information from the file header in order to determine the image size and data type

The data stored in the image are available in the 'data' property.

xrayutilities.io.imagereader.get_tiff (filename, path=None)
read tiff image file and return the data

Parameters: filename: str

filename of the image to be read. so far only single filenames are supported. The data might be compressed.

path : str, optional
 path of the data file

xrayutilities.io.panalytical_xml module

Panalytical XML (www.XRDML.com) data file parser

based on the native python xml.dom.minidom module. want to keep the number of dependancies as small as possible

class xrayutilities.io.panalytical_xml.xrDMLFile (fname, path=")

Bases: object

class to handle XRDML data files. The class is supplied with a file name and uses the XRDMLScan class to parse the xrdMeasurement in the file

class xrayutilities.io.panalytical_xml.XRDMLMeasurement (measurement, namespace=")

Bases: object

class to handle scans in a XRDML datafile

xrayutilities.io.panalytical_xml.getxrdml_map (filetemplate, scannrs=None, path='.', roi=None) parses multiple XRDML file and concatenates the results for parsing the xrayutilities.io.XRDMLFile class is used. The function can be used for parsing maps measured with the PIXCel 1D detector (and in limited way also for data acquired with a point detector -> see getxrdml_scan instead).

Parameters: filetemplate: str

template string for the file names, can contain a %d which is replaced by the scan number or be a list of filenames

scannrs: int or list, optional

scan number(s)

path : str, optional

common path to the filenames

roi: tuple, optional

region of interest for the PIXCel detector, for other measurements this is not useful!

Returns: om, tt, psd : ndarray

motor positions and data as flattened numpy arrays

Examples

xrayutilities.io.panalytical_xml.getxrdml_scan (filetemplate, *motors, **kwargs)

parses multiple XRDML file and concatenates the results for parsing the xrayutilities.io.XRDMLFile class is used. The function can be used for parsing arbitrary scans and will return the the motor values of the scan motor and additionally the positions of the motors given by in the *motors argument

Parameters: filetemplate: str

template string for the file names, can contain a %d which is replaced by the scan number or be a list of filenames given by the scannrs keyword argument

motors: str

motor names to return: e.g.: 'Omega', '2Theta', ... one can also use abbreviations:

- 'Omega' = 'om' = 'o'
- '2Theta' = 'tt' = 't'
- 'Chi' = 'c'
- 'Phi' = 'p'

scannrs: int or list, optional

scan number(s)

path: str, optional

common path to the filenames

Returns: scanmot, mot1, mot2,..., detectorint : ndarray

motor positions and data as flattened numpy arrays

Examples

xrayutilities.io.pdcif module

class xrayutilities.io.pdcif.pdCIF (filename, datacolumn=None)

Bases: object

the class implements a primitive parser for pdCIF-like files. It reads every entry and collects the information in the header attribute. The first loop containing one of the intensity fields is assumed to be the data the user is interested in and is transferred to the data array which is stored as numpy record array the columns can be accessed by name intensity fields:

- _pd_meas_counts_total
- _pd_meas_intensity_total
- _pd_proc_intensity_total
- _pd_proc_intensity_net
- _pd_calc_intensity_total
- pd calc intensity net

alternatively the data column name can be given as argument to the constructor

Parse ()

parser of the pdCIF file. the method reads the data from the file and fills the data and header attributes with content

class xrayutilities.io.pdcif.pdESG (filename, datacolumn=None)

Bases: xrayutilities.io.pdcif.pdCIF

class for parsing multiple pdCIF loops in one file. This includes especially *.esg files which are supposed to consist of multiple loops of pdCIF data with equal length.

Upon parsing the class tries to combine the data of these different scans into a single data matrix -> same shape of subscan data is assumed

Parse ()

parser of the pdCIF file. the method reads the data from the file and fills the data and header attributes with content

xrayutilities.io.pdcif.remove_comments (line, sep='#')

xrayutilities.io.rigaku_ras module

class for reading data + header information from Rigaku RAS (3-column ASCII) files

Such datafiles are generated by the Smartlab Guidance software from Rigaku.

class xrayutilities.io.rigaku_ras.RASFile (filename, path=None)

Bases: object

Represents a RAS data file. The file is read during the constructor call

Parameters: filename: str

name of the ras-file

path: str, optional

path to the data file

Read ()

Read the data from the file

class xrayutilities.io.rigaku_ras.RASScan (filename, pos)

Bases: object

Represents a single Scan portion of a RAS data file. The scan is parsed during the constructor call

Parameters: filename : str

file name of the data file

pos: int

seek position of the 'RAS_HEADER_START' line

xrayutilities.io.rigaku_ras.getras_scan (scanname, scannumbers, *args, **kwargs)

function to obtain the angular cooridinates as well as intensity values saved in RAS datafiles. Especially useful for reciprocal space map measurements, and to combine date from several scans

further more it is possible to obtain even more positions from the data file if more than two string arguments with its names are given

Parameters: scanname: str

name of the scans, for multiple scans this needs to be a template string

scannumbers: int, tuple or list

number of the scans of the reciprocal space map

args: str, optional

names of the motors. to read reciprocal space maps measured in coplanar diffraction give:

- omname: name of the omega motor (or its equivalent)
- ttname: name of the two theta motor (or its equivalent)

kwargs: dict

keyword arguments forwarded to RASFile function

Returns: [ang1, ang2, ...] : *list*

angular positions are extracted from the respective scan header, or motor positions during the scan. this is omitted if no *args* are given

rasdata: ndarray

the data values (includes the intensities e.g. rasdata['int']).

Examples

```
>>> [om, tt], MAP = xu.io.getras_scan('text%05d.ras', 36, 'Omega',
>>> 'TwoTheta')
```

xrayutilities.io.rotanode_alignment module

parser for the alignment log file of the rotating anode

class xrayutilities.io.rotanode_alignment.RA_Alignment (filename)

Bases: object

class to parse the data file created by the alignment routine (tpalign) at the rotating anode spec installation this routine does an iterative alignment procedure and saves the center of mass values were it moves after each scan. It iterates between two different peaks and iteratively aligns at each peak between two different motors (om/chi at symmetric peaks, om/phi at asymmetric peaks)

Parse ()

parser to read the alignment log and obtain the aligned values at every iteration.

get (key)

keys ()

returns a list of keys for which aligned values were parsed

plot (pname)

function to plot the alignment history for a given peak

Parameters: pname: str

peakname for which the alignment should be plotted

xrayutilities.io.seifert module

a set of routines to convert Seifert ASCII files to HDF5 in fact there exist two posibilities how the data is stored (depending on the use detector):

- 1. as a simple line scan (using the point detector)
- 2. as a map using the PSD

In the first case the data ist stored

```
class xrayutilities.io.seifert.SeifertHeader
Bases: object
```

helper class to represent a Seifert (NJA) scan file header

```
class xrayutilities.io.seifert.SeifertMultiScan (filename, m_scan, m2, path=")
```

Bases: object

Class to parse a Seifert (NJA) multiscan file

```
parse ()
```

```
class xrayutilities.io.seifert.SeifertScan (filename, path=")
```

Bases: object

Class to parse a single Seifert (NJA) scan file

```
parse ()
```

```
xrayutilities.io.seifert.getSeifert_map (filetemplate, scannrs=None, path='.', scantype='map',
Nchannels=1280)
```

parses multiple Seifert *.nja files and concatenates the results. for parsing the xrayutilities.io.SeifertMultiScan class is used. The function can be used for parsing maps measured with the Meteor1D and point detector.

Parameters: filetemplate: str

template string for the file names, can contain a %d which is replaced by the scan

number or be a list of filenames

scannrs: int or list, optional

scan number(s)

path: str, optional

common path to the filenames

scantype: {'map', 'tsk'}, optional

type of datafile: can be either 'map' (reciprocal space map measured with a regular

Seifert job (default)) or 'tsk' (MCA spectra measured using the TaskInterpreter)

Nchannels: int, optional

number of channels of the MCA (needed for 'tsk' measurements)

Returns: om, tt, psd: ndarray

positions and data as flattened numpy arrays

Examples

```
>>> om, tt, psd = xrayutilities.io.getSeifert_map("samplename_%d.xrdml",
>>> [1, 2], path="./data")
```

```
xrayutilities.io.seifert.repair_key (key)
```

Repair a key string in the sense that the string is changed in a way that it can be used as a valid Python identifier. For that purpose all blanks within the string will be replaced by _ and leading numbers get an preceeding _.

xrayutilities.io.spec module

a class for observing a SPEC data file

Motivation:

SPEC files can become quite large. Therefore, subsequently reading the entire file to extract a single scan is a quite cumbersome procedure. This module is a proof of concept code to write a file observer starting a reread of the file starting from a stored offset (last known scan position)

```
class xrayutilities.io.spec.specCmdLine (n, prompt, cmdl, out=")
Bases: object
```

class xrayutilities.io.spec.specfile (filename, path=")

Bases: object

This class represents a single SPEC file. The class provides methodes for updateing an already opened file which makes it particular interesting for interactive use.

Parse ()

Parses the file from the starting at last_offset and adding found scans to the scan list.

```
Save2HDF5 (h5f, comp=True, optattrs={})
```

Save the entire file in an HDF5 file. For that purpose a group is set up in the root group of the file with the name of the file without extension and leading path. If the method is called after an previous update only the scans not written to the file meanwhile are saved.

Parameters: h5f: file-handle or str

a HDF5 file object or its filename

comp: bool, optional

activate compression - true by default

Update ()

reread the file and add newly added files. The parsing starts at the data offset of the last scan gathered during the last parsing run.

class xrayutilities.io.spec.speclog (filename, prompt, path=")

Bases: object

class to parse a SPEC log file to find the command history

Parse ()

class xrayutilities.io.spec.specscan (name, scannr, command, date, time, itime, colnames, hoffset, doffset, fname, imopnames, imopvalues, scan_status)

Bases: object

Represents a single SPEC scan. This class is usually not called by the user directly but used via the SPECFile class.

ClearData ()

Delete the data stored in a scan after it is no longer used.

ReadData ()

Set the data attribute of the scan class.

Save2HDF5 (h5f, group='/', title=", optattrs={}, comp=True)

Save a SPEC scan to an HDF5 file. The method creates a group with the name of the scan and stores the data there as a table object with name "data". By default the scan group is created under the root group of the HDF5 file. The title of the scan group is ususally the scan command. Metadata of the scan are stored as attributes to the scan group. Additional custom attributes to the scan group can be passed as a dictionary via the optattrs keyword argument.

Parameters: h5f : file-handle or str

a HDF5 file object or its filename

group: str, optional

name or group object of the HDF5 group where to store the data

title: str, optional

a string with the title for the data, defaults to the name of scan if empty

optattrs: dict, optional

a dictionary with optional attributes to store for the data

comp: bool, optional

activate compression - true by default

SetMCAParams (mca_column_format, mca_channels, mca_start, mca_stop)

Set the parameters used to save the MCA data to the file. This method calculates the number of lines used to store the MCA data from the number of columns and the

Parameters: mca_column_format: int

number of columns used to save the data

mca_channels: int

number of MCA channels stored

mca start: int

first channel that is stored

mca_stop: int

last channel that is stored

getheader element (key, firstonly=True)

return the value-string of the first appearance of this SPECScan's header element, or a list of all values if firstonly=False

Parameters: specscan: SPECScan

key: str

name of the key to return; e.g. 'UMONO' or 'D'

firstonly: bool, optional

flag to specify if all instances or only the first one should be returned

Returns: valuestring: str

header value (if firstonly=True)

[str1, str2, ...]: list

header values (if firstonly=False)

plot (*args, **keyargs)

Plot scan data to a matplotlib figure. If newfig=True a new figure instance will be created. If logy=True (default is False) the y-axis will be plotted with a logarithmic scale.

Parameters: args: list

arguments for the plot: first argument is the name of x-value column the following pairs of arguments are the y-value names and plot styles allowed are 3, 5, 7,... number of arguments

keyargs: dict, optional

newfig: bool, optional

if True a new figure instance will be created otherwise an existing one will be used

logy: bool, optional

if True a semilogy plot will be done

xrayutilities.io.spec.geth5_scan (h5f, scans, *args, **kwargs)

function to obtain the angular cooridinates as well as intensity values saved in an HDF5 file, which was created from a spec file by the Save2HDF5 method. Especially useful for reciprocal space map measurements.

further more it is possible to obtain even more positions from the data file if more than two string arguments with its names are given

Parameters: h5f: file-handle or str

file object of a HDF5 file opened using h5py or its filename

scans: int, tuple or list

number of the scans of the reciprocal space map

args: str, optional

names of the motors. to read reciprocal space maps measured in coplanar diffraction give:

omname: name of the omega motor (or its equivalent)

ttname: name of the two theta motor (or its equivalent)

kwargs: dict, optional

samplename: str, optional

string with the hdf5-group containing the scan data if ommitted the first child node of h5f.root will be used

rettype: {'list', 'numpy'}, optional

how to return motor positions. by default a list of arrays is returned. when rettype == 'numpy' a record array will be returned.

Returns: [ang1, ang2, ...] : *list*

angular positions of the center channel of the position sensitive detector (numpy.ndarray 1D), this list is omitted if no *args* are given

MAP: ndarray

the data values as stored in the data file (includes the intensities e.g. MAP['MCA']).

Examples

```
>>> [om, tt], MAP = xu.io.geth5_scan(h5file, 36, 'omega', 'gamma')
```

xrayutilities.io.spec.getspec_scan (specf, scans, *args, **kwargs)

function to obtain the angular cooridinates as well as intensity values saved in a SPECFile. Especially useful to combine the data from multiple scans.

further more it is possible to obtain even more positions from the data file if more than two string arguments with its names are given

Parameters: specf : SPECFile

file object

scans: int, tuple or list number of the scans

args: str

names of the motors and counters

rettype: {'list', 'numpy'}, optional

how to return motor positions. by default a list of arrays is returned. when rettype ==

'numpy' a record array will be returned.

Returns: [ang1, ang2, ...] : *list*

coordinates and counters from the SPEC file

Examples

```
>>> [om, tt, cnt2] = xu.io.getspec_scan(s, 36, 'omega', 'gamma',
>>>
```

xrayutilities.io.spectra module

module to handle spectra data

class xrayutilities.io.spectra

Bases: object

Represents a SPECTRA data file. The file is read during the Constructor call. This class should work for data stored at beamlines P08 and BW2 at HASYLAB.

Parameters: filename: str

a string with the name of the SPECTRA file

mcatmp: str, optional

template for the MCA files

mcastart, mcastop: int, optional

start and stop index for the MCA files, if not given, the class tries to determine the start and stop index automatically.

Read ()

Read the data from the file.

ReadMCA ()

Save2HDF5 (h5file, name, group='/', mcaname='MCA')

Saves the scan to an HDF5 file. The scan is saved to a seperate group of name "name". h5file is either a string for the file name or a HDF5 file object. If the mca attribute is not None mca data will be stored to an chunked array of with name mcaname.

Parameters: h5file: file-handle or str

HDF5 file object or name

name: str

name of the group where to store the data

group: str, optional

root group where to store the data

mcaname: str, optional

Name of the MCA in the HDF5 file

Returns: bool or None

The method returns None in the case of everything went fine, True otherwise.

class xrayutilities.io.spectra.SPECTRAFileComments

Bases: dict

Class that describes the comments in the header of a SPECTRA file. The different comments are accessible via the comment keys.

class xrayutilities.io.spectra.SPECTRAFileData

Bases: object

append (col)

class xrayutilities.io.spectra.SPECTRAFileDataColumn (index, name, unit, type)

Bases: object

<code>class xrayutilities.io.spectra.SPECTRAFileParameters () -> new empty dictionary dict(mapping) -> new dictionary initialized from a mapping object's (key, value) pairs dict(iterable) -> new dictionary initialized as if via: $d = \{\}$ for k, v in iterable: $d[k] = v \ dict(**kwargs) -> new \ dictionary initialized with the name=value pairs in the keyword argument list. For example: <math>d[c] = v \ dict(one=1, two=2)$ </code>

Bases: dict

xrayutilities.io.spectra.geth5_spectra_map (h5file, scans, *args, **kwargs)

function to obtain the omega and twotheta as well as intensity values for a reciprocal space map saved in an HDF5 file, which was created from a spectra file by the Save2HDF5 method.

further more it is possible to obtain even more positions from the data file if more than two string arguments with its names are given

Parameters: h5f: file-handle or str

file object of a HDF5 file opened using h5py

scans: int, tuple or list

number of the scans of the reciprocal space map

args: str, optional

arbitrary number of motor names

omname: name of the omega motor (or its equivalent)

• ttname: name of the two theta motor (or its equivalent)

kwargs: dict, optional

mca: str, optional

name of the mca data (if available) otherwise None (default: "MCA")

samplename: str, optional

string with the hdf5-group containing the scan data if omitted the first child node of

h5f.root will be used to determine the sample name

Returns: [ang1, ang2, ...] : *list*

angular positions of the center channel of the position sensitive detector

(numpy.ndarray 1D). one entry for every args-argument given to the function

MAP: ndarray

the data values as stored in the data file (includes the intensities e.g. MAP['MCA']).

Module contents

xrayutilities.materials package

Submodules

xrayutilities.materials.atom module

module containing the Atom class which handles the database access for atomic scattering factors and the atomic mass.

class xrayutilities.materials.atom.Atom (name, num)

Bases: object

property color

f (q, en='config')

function to calculate the atomic structure factor F

Parameters: q: float, array-like

momentum transfer en: float or str, optional

energy for which F should be calculated, if omitted the value from the xrayutilities

configuration is used

Returns: float or array-like

value(s) of the atomic structure factor

£0 (q)

f1 (en='config')

```
£2 (en='config')
```

get_cache (prop, key)

check if a cached value exists to speed up repeated database requests

Returns: bool

True then result contains the cached otherwise False and result is None

result: database value

```
max_cache_length = 1000
```

property radius

set_cache (prop, key, result)

set result to be cached to speed up future calls

property weight

```
xrayutilities.materials.atom.get_key (*args)
generate a hash key for several possible types of arguments
```

xrayutilities.materials.cif module

class xrayutilities.materials.cif.CIFDataset (fid, name, digits)

Bases: object

class for parsing CIF (Crystallographic Information File) files. The class aims to provide an additional way of creating material classes instead of manual entering of the information the lattice constants and unit cell structure are parsed from the CIF file

Parse (fid)

function to parse a CIF data set. The function reads the space group symmetry operations and the basic atom positions as well as the lattice constants and unit cell angles

SGLattice (use_p1=False)

create a SGLattice object with the structure from the CIF file

SymStruct ()

function to obtain the list of different atom positions in the unit cell for the different types of atoms and determine the space group number and origin choice if available. The data are obtained from the data parsed from the CIF file.

class xrayutilities.materials.cif.CIFFile (filestr, digits=4)

Bases: object

class for parsing CIF (Crystallographic Information File) files. The class aims to provide an additional way of creating material classes instead of manual entering of the information the lattice constants and unit cell structure are parsed from the CIF file.

If multiple datasets are present in the CIF file this class will attempt to parse all of them into the the data dictionary. By default all methods access the first data set found in the file.

Parse ()

function to parse a CIF file. The function reads all the included data sets and adds them to the data dictionary.

```
SGLattice (dataset=None, use_p1=False)
```

create a SGLattice object with the structure from the CIF dataset

Parameters: dataset: str, optional

name of the dataset to use. if None the default one will be used.

use_p1: bool, optional

force the use of P1 symmetry, default False

xrayutilities.materials.cif.cifexport (filename, mat)

function to export a Crystal instance to CIF file. This in particular includes the atomic coordinates, however, ignores for example the elastic parameters.

xrayutilities.materials.database module

module to handle the access to the optical parameters database

class xrayutilities.materials.database.DataBase (fname)

Bases: object

Close ()

Close an opend database file.

Create (dbname, dbdesc)

Creates a new database. If the database file already exists its content is delete.

Parameters: dbname: str

name of the database

dbdesc: str

a short description of the database

CreateMaterial (name, description)

This method creates a new material. If the material group already exists the procedure is aborted.

Parameters: name: str

name of the material

description: str

description of the material

GetF0 (q, dset='default')

Obtain the f0 scattering factor component for a particular momentum transfer q.

Parameters: q: float or array-like

momentum transfer

dset: str, optional

specifies which dataset (different oxidation states) should be used

GetF1 (en)

Return the second, energy dependent, real part of the scattering factor for a certain energy en.

Parameters: en: float or array-like

energy

GetF2 (en)

Return the imaginary part of the scattering factor for a certain energy en.

Parameters: en: float or array-like

energy

Open (mode='r')

Open an existing database file.

SetColor (color)

Save color of the element for visualization

Parameters: color: tuple, str

matplotlib color for the element

setF0 (parameters, subset='default')

Save f0 fit parameters for the set material. The fit parameters are stored in the following order: c, a1, b1,......, a4, b4

Parameters: parameters: list or array-like

fit parameters **subset**: str, optional

name the f0 dataset

SetF1F2 (en, f1, f2)

Set f1, f2 values for the active material.

Parameters: en: list or array-like

energy in (eV)

f1 : list or array-like

f1 values

f2: list or array-like

f2 values

SetMaterial (name)

Set a particular material in the database as the actual material. All operations like setting and getting optical constants are done for this particular material.

Parameters: name: str

name of the material

SetRadius (radius)

Save atomic radius for visualization

Parameters: radius: float

atomic radius in Angstrom

SetWeight (weight)

Save weight of the element as float

Parameters: weight: float

atomic standard weight of the element

xrayutilities.materials.database.add_color_from_JMOL (db, cfile, verbose=False)
Read color from JMOL color table and save it to the database.

xrayutilities.materials.database.add_f0_from_intertab (db, itf, verbose=False)
Read f0 data from International Tables of Crystallography and add it to the database.

xrayutilities.materials.database.add_f0_from_xop (db, xop, verbose=False) Read f0 data from f0 xop.dat and add it to the database.

xrayutilities.materials.database.add_f1f2_from_ascii_file (db, asciifile, element, verbose=False)
Read f1 and f2 data for specific element from ASCII file (3 columns) and save it to the database.

xrayutilities.materials.database.add_f1f2_from_henkedb (db, hf, verbose=False) Read f1 and f2 data from Henke database and add it to the database.

xrayutilities.materials.database.add_f1f2_from_kissel (db, kf, verbose=False)
Read f1 and f2 data from Henke database and add it to the database.

xrayutilities.materials.database.add mass from NIST (db, nistfile, verbose=False)

Read atoms standard mass and save it to the database. The mass of the natural isotope mixture is taken from the NIST data!

xrayutilities.materials.database.add_radius_from_wiki (db, dfile, verbose=False)
Read radius from Wikipedia radius table and save it to the database.

xrayutilities.materials.database.createAndFillDatabase (fname, dpath=None, verbose=False) function to create the database and fill it with values from the various source files.

Parameters: fname: str

Filename of the database to be created (including the path)

dpath: str, optional

directory where all the source data files are stored

verbose: bool, optional

flag to determine the verbosity of the script (default: False)

xrayutilities.materials.database.init_material_db (db)

xrayutilities.materials.elements module

xrayutilities.materials.heuslerlib module

implement convenience functions to define Heusler materials.

xrayutilities.materials.heuslerlib.**FullHeuslerCubic225** (X, Y, Z, a, biso=[0, 0, 0], occ=[1, 1, 1]) Full Heusler structure with formula X2YZ. Strukturberichte symbol L2_1; space group Fm-3m (225)

Parameters: X, Y, Z: str or Element

elements

a: float

cubic lattice parameter in Angstroem

biso: list of floats, optional

Debye Waller factors for X, Y, Z elements

occ: list of floats, optional

occupation numbers for the elements X, Y, Z

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.heuslerlib.**FullHeuslerCubic225_A2** (X, Y, Z, a, a2dis, biso=[0, 0, 0], occ=[1, 1, 1])

Full Heusler structure with formula X2YZ. Strukturberichte symbol L2_1; space group Fm-3m (225) with A2-type (W) disorder

Parameters: X, Y, Z: str or Element

elements

a: float

cubic lattice parameter in Angstroem

a2dis: float

amount of A2-type disorder (0: fully ordered, 1: fully disordered)

biso: list of floats, optional

Debye Waller factors for X, Y, Z elements

occ: list of floats, optional

occupation numbers for the elements X, Y, Z

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.heuslerlib.**FullHeuslerCubic225_B2** (X, Y, Z, a, b2dis, biso=[0, 0, 0], occ=[1, 1, 1])

Full Heusler structure with formula X2YZ. Strukturberichte symbol L2_1; space group Fm-3m (225) with B2-type (CsCl) disorder

Parameters: X, Y, Z: str or Element

elements

a: float

cubic lattice parameter in Angstroem

b2dis: float

amount of B2-type disorder (0: fully ordered, 1: fully disordered)

biso: list of floats, optional

Debye Waller factors for X, Y, Z elements

occ: list of floats, optional

occupation numbers for the elements X, Y, Z

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.heuslerlib.**FullHeuslerCubic225_DO3** (X, Y, Z, a, do3disxy, do3disxz, biso=[0, 0, 0], occ=[1, 1, 1])

Full Heusler structure with formula X2YZ. Strukturberichte symbol L2_1; space group Fm-3m (225) with DO_3-type (BiF3) disorder, either between atoms X <-> Y or X <-> Z.

Parameters: X, Y, Z: str or Element

elements

a: float

cubic lattice parameter in Angstroem

do3disxy: float

amount of DO_3-type disorder between X and Y atoms (0: fully ordered, 1: fully disordered)

do3disxz: float

amount of DO_3-type disorder between X and Z atoms (0: fully ordered, 1: fully disordered)

biso: list of floats, optional

Debye Waller factors for X, Y, Z elements

occ: list of floats, optional

occupation numbers for the elements X, Y, Z

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.heuslerlib.**HeuslerHexagonal194** (X, Y, Z, a, c, biso=[0, 0, 0], occ=[1, 1, 1]) Hexagonal Heusler structure with formula XYZ space group P63/mmc (194)

Parameters: X, Y, Z: str or Element

elements

a.c: float

hexagonal lattice parameters in Angstroem

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.heuslerlib.**HeuslerTetragonal119** (X, Y, Z, a, c, biso=[0, 0, 0], occ=[1, 1, 1]) Tetragonal Heusler structure with formula X2YZ space group I-4m2 (119)

Parameters: X, Y, Z: str or Element

elements **a, c**: float

tetragonal lattice parameters in Angstroem

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.heuslerlib.**HeuslerTetragonal139** (X, Y, Z, a, c, biso=[0, 0, 0], occ=[1, 1, 1]) Tetragonal Heusler structure with formula X2YZ space group I4/mmm (139)

Parameters: X, Y, Z: str or Element

elements

a, c: float

tetragonal lattice parameters in Angstroem

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.heuslerlib.**InverseHeuslerCubic216** (X, Y, Z, a, biso=[0, 0, 0], occ=[1, 1, 1])

Full Heusler structure with formula (XY)X'Z structure; space group F-43m (216)

Parameters: X, Y, Z: str or Element

elements

a: float

cubic lattice parameter in Angstroem

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.material module

Classes decribing materials. Materials are devided with respect to their crystalline state in either Amorphous or Crystal types. While for most materials their crystalline state is defined few materials are also included as amorphous which can be useful for calculation of their optical properties.

class xrayutilities.materials.material.alloy (matA, matB, x)

Bases: xrayutilities.materials.material.Crystal

alloys two materials from the same crystal system. If the materials have the same space group the Wyckoff positions within the unit cell will also reflect the alloying.

RelaxationTriangle (hkl, sub, exp)

function which returns the relaxation triangle for a Alloy of given composition. Reciprocal space coordinates are calculated using the user-supplied experimental class

Parameters: hkl : list or array-like

Miller Indices

sub: Crystal, or float

substrate material or lattice constant

exp : Experiment

object from which the Transformation object and ndir are needed

Returns: qy, qz : float

reciprocal space coordinates of the corners of the relaxation triangle

static check_compatibility (matA, matB)

static lattice_const_AB (latA, latB, x, name=")

method to calculated the interpolation of lattice parameters and unit cell angles of the Alloy. By default linear interpolation between the value of material A and B is performed.

Parameters: latA, latB: float or vector

property (lattice parameter/angle) of material A and B. A property can be a scalar or vector.

x: float

fraction of material B in the alloy.

name: str, optional

label of the property which is interpolated. Can be 'a', 'b', 'c', 'alpha', 'beta', or

'gamma'.

property x

class xrayutilities.materials.material.Amorphous (name, density, atoms=None, cij=None)

Bases: xrayutilities.materials.material.Material

amorphous materials are described by this class

chi0 (en='config')

calculates the complex chi_0 values often needed in simulations. They are closely related to delta and beta (n = 1 + $chi_0/2$ + i* $chi_0/2$ vs. n = 1 - delta + i*beta)

delta (en='config')

function to calculate the real part of the deviation of the refractive index from 1 (n=1-delta+i*beta)

Parameters: en: float, array-like or str, optional

energy of the x-rays in eV

Returns: float or array-like

ibeta (en='config')

function to calculate the imaginary part of the deviation of the refractive index from 1 (n=1-delta+i*beta)

Parameters: en: float, array-like or str, optional

energy of the x-rays in eV

Returns: float or array-like

static parseChemForm (cstring)

Parse a string containing a simple chemical formula and transform it to a list of elements together with their relative atomic fraction. e.g. 'H2O' -> [(H, 2/3), (O, 1/3)], where H and O are the Element objects of Hydrogen and Oxygen. Note that every chemical element needs to start with a capital letter! Complicated formulas containing bracket are not supported!

Parameters: cstring: str

string containing the chemical fomula

Returns: list of tuples

chemical element and atomic fraction

xrayutilities.materials.material.Cij2Cijkl(Cij)

Converts the elastic constants matrix (tensor of rank 2) to the full rank 4 cijkl tensor.

Parameters: cij: array-like

(6, 6) cij matrix

Returns: cijkl ndarray

(3, 3, 3, 3) cijkl tensor as numpy array

xrayutilities.materials.material.Cijkl2Cij (cijkl)

Converts the full rank 4 tensor of the elastic constants to the (6, 6) matrix of elastic constants.

Parameters: cijkl ndarray

(3, 3, 3, 3) cijkl tensor as numpy array

Returns: cij: array-like

(6, 6) cij matrix

class xrayutilities.materials.material.crystal (name, lat, cij=None, thetaDebye=None)

Bases: xrayutilities.materials.material.Material

Crystalline materials are described by this class

ApplyStrain (strain)

Applies a certain strain on the lattice of the material. The result is a change in the base vectors of the real space as well as reciprocal space lattice. The full strain matrix (3x3) needs to be given.

Note

NO elastic response of the material will be considered!

property B

GetMismatch (mat)

Calculate the mismatch strain between the material and a second material

HKL (*q)

Return the HKL-coordinates for a certain Q-space position.

Parameters: q: list or array-like

Q-position. its also possible to use HKL(qx, qy, qz).

Q (*hkl)

Return the Q-space position for a certain material.

Parameters: hkl: list or array-like

Miller indices (or Q(h, k, l) is also possible)

StructureFactor (q, en='config', temp=0)

calculates the structure factor of a material for a certain momentum transfer and energy at a certain temperature of the material

Parameters: q: list, tuple or array-like

vectorial momentum transfer

en: float or str, optional

x-ray energy eV, if omitted the value from the xrayutilities configuration is used

temp: float

temperature used for Debye-Waller-factor calculation

Returns: complex

the complex structure factor

StructureFactorForEnergy (q0, en, temp=0)

calculates the structure factor of a material for a certain momentum transfer and a bunch of energies

Parameters: q0: list, tuple or array-like

vectorial momentum transfer

en: list, tuple or array-like energy values in eV

temp: float

temperature used for Debye-Waller-factor calculation

Returns: array-like

complex valued structure factor array

StructureFactorForQ (q, en0='config', temp=0)

calculates the structure factor of a material for a bunch of momentum transfers and a certain energy

Parameters: q: list of vectors or array-like

> vectorial momentum transfers; list of vectores (list, tuple or array) of length 3 e.g.: (Si.Q(0, 0, 4), Si.Q(0, 0, 4.1),...) or numpy.array([Si.Q(0, 0, 4), Si.Q(0, 0, 4.1)])

en0: float or str, optional

x-ray energy eV, if omitted the value from the xrayutilities configuration is used

temp: float

temperature used for Debye-Waller-factor calculation

Returns: array-like

complex valued structure factor array

property a

property a1

property a2

property a3

property alpha

property b

property beta

property c

chemical_composition (natoms=None, with_spaces=False, ndigits=2) determine chemical composition from occupancy of atomic positions.

Parameters: mat : Crystal

> instance of Crystal natoms: int, optional

> > number of atoms to normalize the formula, if None some automatic normalization is attempted using the greatest common divisor of the number of atoms per unit cell. If the number of atoms of any element is fractional natoms=1 is used.

with_spaces: bool, optional

add spaces between the different entries in the output string for CIF combatibility

ndigits: int, optional

number of digits to which floating point numbers are rounded to

Returns: str

representation of the chemical composition

chi0 (en='config')

calculates the complex chi_0 values often needed in simulations. They are closely related to delta and beta (n = 1 + $chi_0/2$ + i* $chi_0/2$ vs. n = 1 - delta + i*beta)

chih (q, en='config', temp=0, polarization='S')

calculates the complex polarizability of a material for a certain momentum transfer and energy

Parameters: q: list, tuple or array-like

momentum transfer vector in (1/A)

en: float or str, optional

x-ray energy eV, if omitted the value from the xrayutilities configuration is used

temp: float, optional

temperature used for Debye-Waller-factor calculation

polarization: {'S', 'P'}, optional sigma or pi polarization

Returns: tuple

(abs(chih_real), abs(chih_imag)) complex polarizability

dTheta (Q, en='config')

function to calculate the refractive peak shift

Parameters: Q: list, tuple or array-like

momentum transfer vector (1/A)

en: float or str, optional

x-ray energy eV, if omitted the value from the xrayutilities configuration is used

Returns: float

peak shift in degree

delta (en='config')

function to calculate the real part of the deviation of the refractive index from 1 (n=1-delta+i*beta)

Parameters: en: float or str, optional

x-ray energy eV, if omitted the value from the xrayutilities configuration is used

Returns: float

property density

calculates the mass density of an material from the mass of the atoms in the unit cell.

Returns: float

mass density in kg/m^3

distances ()

function to obtain distances of atoms in the crystal up to the unit cell size (largest value of a, b, c is the cut-off) returns a list of tuples with distance d and number of occurrence n [(d1, n1), (d2, n2),...]

Note

if the base of the material is empty the list will be empty

environment (*pos, **kwargs)

Returns a list of neighboring atoms for a given position within the unit cell. If the material does not contain any atoms a dummy atom will be placed on the unit cell corners.

Parameters: pos: list or array-like

fractional coordinate in the unit cell

maxdist: float

maximum distance wanted in the list of neighbors (default: 7)

Returns: list of tuples

(distance, atomType, multiplicity) giving distance sorted list of atoms

classmethod fromCIF (ciffilestr, **kwargs)

Create a Crystal from a CIF file. The default data-set from the cif file will be used to create the Crystal.

Parameters: ciffilestr: str, bytes

filename of the CIF file or string representation of the CIF file

kwargs: dict

keyword arguments are passed to the init-method of CIFFile

Returns: Crystal

property gamma

ibeta (en='config')

function to calculate the imaginary part of the deviation of the refractive index from 1 (n=1-delta+i*beta)

Parameters: en: float or str, optional

x-ray energy eV, if omitted the value from the xrayutilities configuration is used

Returns: float

loadLatticefromCIF (ciffilestr)

load the unit cell data (lattice) from the CIF file. Other material properties stay unchanged.

Parameters: ciffilestr: str, bytes

filename of the CIF file or string representation of the CIF file

planeDistance (*hkl)

determines the lattice plane spacing for the planes specified by (hkl)

Parameters: h, k, I: list, tuple or floats

Miller indices of the lattice planes given either as list, tuple or seperate arguments

Returns: float

the lattice plane spacing

Examples

```
>>> xu.materials.Si.planeDistance(0, 0, 4)
1.357760000000001
```

```
or
```

```
>>> xu.materials.Si.planeDistance((1, 1, 1))
3.1356124059796255
```

show_unitcell (fig=None, subplot=111, scale=0.6, complexity=11, linewidth=1.5, mode='matplotlib') visualization of the unit cell using either matplotlibs basic 3D functionality (expect rendering inaccuracies!) or the mayavi mlab package (accurate rendering -> recommended!)

Note

For more flexible visualization consider using the CIF-export feature and use a proper crystal structure viewer.

Parameters: fig: matplotlib Figure, Mayavi Scene, or None, optional

subplot: int or list, optional

subplot to use for the visualization when using matplotlib. This argument of fowarded to the first argument of matplotlibs *add_subplot* function

scale: float, optional

scale the size of the atoms by this additional factor. By default the size of the atoms corresponds to 60% of their atomic radius.

complexity: int, optional

number of steps to approximate the atoms as spheres. Higher values make spheres more accurate, but cause slower plotting.

linewidth: float, optional

line thickness of the unit cell outline

mode: str, optional

defines the plot backend used, can be 'matplotlib' (default) or 'mayavi'.

Returns: figure object of either matplotlib or Mayavi

toCIF (ciffilename)

Export the Crystal to a CIF file.

Parameters: ciffilename: str

filename of the CIF file

class xrayutilities.materials.material.CubicAlloy (matA, matB, x)

Bases: xrayutilities.materials.material.Alloy

ContentBasym (q_inp, q_perp, hkl, sur)

function that determines the content of B in the alloy from the reciprocal space position of an asymmetric peak.

Parameters: q_inp : float

inplane peak position of reflection hkl of the alloy in reciprocal space

q_perp : float

perpendicular peak position of the reflection hkl of the alloy in reciprocal space

hkl: list

Miller indices of the measured asymmetric reflection

sur : list

Miller indices of the surface (determines the perpendicular direction)

Returns: content: float

content of B in the alloy determined from the input variables

list

[a_inplane a_perp, a_bulk_perp(x), eps_inplane, eps_perp]; lattice parameters calculated from the reciprocal space positions as well as the strain (eps) of the layer

ContentBsym (q_perp, hkl, inpr, asub, relax)

function that determines the content of B in the alloy from the reciprocal space position of a symetric peak. As an additional input the substrates lattice parameter and the degree of relaxation must be given

Parameters: q_perp: float

perpendicular peak position of the reflection hkl of the alloy in reciprocal space

hkl: list

Miller indices of the measured symmetric reflection (also defines the surface normal

inpr: list

Miller indices of a Bragg peak defining the inplane reference direction

asub: float

substrate lattice parameter

relax: float

degree of relaxation (needed to obtain the content from symmetric reciprocal space

position)

Returns: content: float

the content of B in the alloy determined from the input variables

xrayutilities.materials.material.CubicElasticTensor (c11, c12, c44)

Assemble the 6x6 matrix of elastic constants for a cubic material from the three independent components of a cubic crystal

Parameters: c11, c12, c44: float

independent components of the elastic tensor of cubic materials

Returns: cij: ndarray

6x6 matrix with elastic constants

xrayutilities.materials.material.HexagonalElasticTensor (c11, c12, c13, c33, c44)

Assemble the 6x6 matrix of elastic constants for a hexagonal material from the five independent components of a hexagonal crystal

Parameters: c11, c12, c13, c33, c44 : float

independent components of the elastic tensor of a hexagonal material

Returns: cij: ndarray

6x6 matrix with elastic constants

class xrayutilities.materials.material.Material (name, cij=None)

Bases: abc.ABC

base class for all Materials. common properties of amorphous and crystalline materials are described by this class from which Amorphous and Crystal are derived from.

absorption_length (en='config')

wavelength dependent x-ray absorption length defined as mu = lambda/(2*pi*2*beta) with lambda and beta as the x-ray wavelength and complex part of the refractive index respectively.

Parameters: en: float or str, optional

energy of the x-rays in eV

Returns: float

the absorption length in um

chi0 (en='config')

calculates the complex chi_0 values often needed in simulations. They are closely related to delta and beta (n = 1 + chi r0/2 + i*chi i0/2 vs. n = 1 - delta + i*beta)

critical_angle (en='config', deg=True)

calculate critical angle for total external reflection

Parameters: en: float or str. optional

energy of the x-rays in eV, if omitted the value from the xrayutilities configuration is

used

deg: bool, optional

return angle in degree if True otherwise radians (default:True)

Returns: float

Angle of total external reflection

abstract delta (en='config')

abstract method which every implementation of a Material has to override

property density

abstract ibeta (en='config')

abstract method which every implementation of a Material has to override

idx_refraction (en='config')

function to calculate the complex index of refraction of a material in the x-ray range

Parameters: en: energy of the x-rays, if omitted the value from the

xrayutilities configuration is used

Returns: n (complex)

property lam

property mu

property nu

xrayutilities.materials.material.**PseudomorphicMaterial** (sub, layer, relaxation=0, trans=None)
This function returns a material whos lattice is pseudomorphic on a particular substrate material. The two materials must have similar unit cell definitions for the algorithm to work correctly, i.e. it does not work for combiniations of materials with different lattice symmetry. It is also crucial that the layer object includes values for the elastic tensor.

Parameters: sub: Crystal

substrate material

layer: Crystal

bulk material of the layer, including its elasticity tensor

relaxation: float, optional

degree of relaxation 0: pseudomorphic, 1: relaxed (default: 0)

trans: Tranform

Transformation which transforms lattice directions into a surface orientated coordinate frame (x, y inplane, z out of plane). If None a (001) surface geometry of a cubic material

is assumed.

Returns: An instance of Crystal holding the new pseudomorphically

strained material.

Raises: InputError

If the layer material has no elastic parameters

xrayutilities.materials.material.WZTensorFromCub (c11ZB, c12ZB, c44ZB)

Determines the hexagonal elastic tensor from the values of the cubic elastic tensor under the assumptions presented in Phys. Rev. B 6, 4546 (1972), which are valid for the WZ <-> ZB polymorphs.

Parameters: c11, c12, c44: float

independent components of the elastic tensor of cubic materials

Returns: cij: ndarray

6x6 matrix with elastic constants

Implementation according to a patch submitted by Julian Stangl

xrayutilities.materials.material.index_map_ij2ijkl(ij)
xrayutilities.materials.material.index_map_ijkl2ij(i,j)

xrayutilities.materials.plot module

xrayutilities.materials.plot.show_reciprocal_space_plane (mat, exp, ttmax=None, maxqout=0.01, scalef=100, ax=None, color=None, show_Laue=True, show_legend=True, projection='perpendicular', label=None) show a plot of the coplanar diffraction plane with peak positions for the respective material. the size of the spots is scaled with the strength of the structure factor

Parameters: mat: Crystal

instance of Crystal for structure factor calculations

exp: Experiment

instance of Experiment (needs to be HXRD, or FourC for onclick action to work correctly). defines the inplane and out of plane direction as well as the sample azimuth

ttmax: float, optional

maximal 2Theta angle to consider, by default 180deg

maxqout: float, optional

maximal out of plane q for plotted Bragg peaks as fraction of exp.k0

scalef: float, or callable, optional

scale factor or function for the marker size. If this is a function it should take only one float argument and return another float which is used as 's' parameter in matplotlib.pyplot.scatter

ax: matplotlib.Axes, optional

matplotlib Axes to use for the plot, useful if multiple materials should be plotted in one plot

color: matplotlib color, optional

show_Laue: bool, optional

flag to indicate if the Laue zones should be indicated

show_legend: bool, optional

flag to indiate if a legend should be shown

projection: 'perpendicular', 'polar', optional

type of projection for Bragg peaks which do not fall into the diffraction plane. 'perpendicular' (default) uses only the inplane component in the scattering plane, whereas 'polar' uses the vectorial absolute value of the two inplane components. See also the 'maxqout' option.

label: None or str, optional

label to be used for the legend. If 'None' the name of the material will be used.

Returns: Axes, plot_handle

xrayutilities.materials.predefined materials module

```
class xrayutilities.materials.predefined materials.AlGaAs (X)
```

Bases: xrayutilities.materials.material.CubicAlloy

class xrayutilities.materials.predefined_materials.SiGe (x)

Bases: xrayutilities.materials.material.CubicAlloy

static lattice_const_AB (latA, latB, x, **kwargs)

method to calculate the lattice parameter of the SiGe alloy with composition Si_{1-x}Ge_x

xrayutilities.materials.spacegrouplattice module

module handling crystal lattice structures. A SGLattice consists of a space group number and the position of atoms specified as Wyckoff positions along with their parameters. Depending on the space group symmetry only certain parameters of the resulting instance will be settable! A cubic lattice for example allows only to set its 'a' lattice parameter but none of the other unit cell shape parameters.

class xrayutilities.materials.spacegrouplattice.SGLattice (sgrp, *args, **kwargs)

Bases: object

lattice object created from the space group number and corresponding unit cell parameters. atoms in the unit cell are specified by their Wyckoff position and their free parameters.

ApplyStrain (eps)

Applies a certain strain on a lattice. The result is a change in the base vectors. The full strain matrix (3x3) needs to be given.

Note

Here you specify the strain and not the stress -> NO elastic response of the material will be considered!

Note

Although the symmetry of the crystal can be lowered by this operation the spacegroup remains unchanged! The 'free_parameters' attribute is, however, updated to mimic the possible reduction of the symmetry.

Parameters: eps: array-like

a 3x3 matrix with all strain components

GetHKL (*args)

determine the Miller indices of the given reciprocal lattice points

GetPoint (*args)

determine lattice points with indices given in the argument

Examples

```
>>> xu.materials.Si.lattice.GetPoint(0, 0, 4) array([ 0. , 0. , 21.72416])
```

```
or
```

```
>>> xu.materials.Si.lattice.GetPoint((1, 1, 1))
array([ 5.43104,  5.43104,  5.43104])
```

GetQ (*args)

determine the reciprocal lattice points with indices given in the argument

UnitCellVolume ()

function to calculate the unit cell volume of a lattice (angstrom^3)

property a

property alpha

property b

base ()

generator of atomic position within the unit cell.

property beta

property c

convert_to_P1()

create a P1 equivalent of this SGLattice instance.

Returns: SGLattice

instance with the same properties as the present lattice, however, in the P1 setting.

equivalent_hkls (hkl)

returns a list of equivalent hkl peaks depending on the crystal system

findsym()

method to return the highest symmetry description of the current material. This method does not consider to change the unit cell dimensions but only searches the highest symmetry spacegroup which with the current unit cell setting can be described. It is therefore not an implementation of FINDSYM [1].

Returns: new SGLattice-instance

a new SGLattice instance is returned with the highest available symmetry description. (see restrictions above)

[1] https://stokes.byu.edu/iso/findsym.php

property gamma

get_allowed_hkl (qmax)

return a set of all allowed reflections up to a maximal specified momentum transfer.

Parameters: qmax: float

maximal momentum transfer

Returns: hklset: set

set of allowed hkl reflections

hkl_allowed (hkl, returnequivalents=False)

check if Bragg reflection with Miller indices hkl can exist according to the reflection conditions. If no reflection conditions are available this function returns True for all hkl values!

Parameters: hkl: tuple or list

Miller indices of the reflection to check

returnequivalents: bool, optional

If True all the equivalent Miller indices of hkl are returned in a set as second return

argument.

Returns: allowed: bool

True if reflection can have non-zero structure factor, false otherwise

equivalents: set, optional

set of equivalent Miller indices if returnequivalents is True

property iscentrosymmetric

returns a boolean to determine if the lattice has centrosymmetry.

isequivalent (hkl1, hkl2)

determining if hkl1 and hkl2 are two crystallographical equivalent pairs of Miller indices. Note that this function considers the effect of non-centrosymmetry!

Parameters: hkl1, hkl2: list

Miller indices to be checked for equivalence

Returns: bool

reflection_conditions()

return string of reflection conditions, both general (from space group) and of Wyckoff positions

property symops

return the set of symmetry operations from the general Wyckoff position of the space group.

transform (mat, origin)

Transform the unit cell with the matrix and origin shift given in the parameters. This function returns a new instance of SGLattice which contains the highest possible symmetry description of the transformed unit cell. After the transformation (see [1]) the findsym method is used to create the new SGLattice instance.

Parameters: mat: (3, 3) list, or ndarray, optional

> transformation matrix of the unit cell. The matrix definition aims to be consistent with what is used on the Bilbao Crystallographic Server [1]. This only defines the linear part, while the origin shift is given by origin.

origin: (3,) list, or ndarray origin shift of the unit cell [1].

[1] https://www.cryst.ehu.es/cgi-bin/cryst/programs/nph-doc-trmat

class xrayutilities.materials.spacegrouplattice.symOp (D, t, m=1)

Bases: object

Class descriping a symmetry operation in a crystal. The symmetry operation is characterized by a 3x3 transformation matrix as well as a 3-vector describing a translation. For magnetic symmetry operations also the time reversal symmetry can be specified (not used in xrayutilities)

property D

transformation matrix of the symmetry operation

```
apply (vec, foldback=True)
apply_axial (vec)
apply_rotation (vec)
combine (other)
static foldback (V)
```

classmethod from_xyz (xyz)

create a SymOp from the xyz notation typically used in CIF files.

Parameters: xyz: str

string describing the symmetry operation (e.g. '-y, -x, z')

property t

translation vector of the symmetry operation

```
xyz (showtimerev=False)
```

return the symmetry operation in xyz notation

class xrayutilities.materials.spacegrouplattice.WyckoffBase (*args, **kwargs)

Bases: list

The WyckoffBase class implements a container for a set of Wyckoff positions that form the base of a crystal lattice. An instance of this class can be treated as a simple container object.

append (atom, pos, occ=1.0, b=0.0) add new Atom to the lattice base

Parameters: atom: Atom

object to be added

pos: tuple or str

Wyckoff position of the atom, along with its parameters. Examples: ('2i', (0.1, 0.2,

0.3)), or '1a'

occ: float, optional

occupancy (default=1.0)

b: float, optional

b-factor of the atom used as $\exp(-b^*q^{**}2/(4^*pi)^{**}2)$ to reduce the intensity of this atom

(only used in case of temp=0 in StructureFactor and chi calculation)

static entry_eq (e1, e2)

compare two entries including all its properties to be equal

Parameters: e1, e2: tuple

tuples with length 4 containing the entries of WyckoffBase which should be compared

index (item)

return the index of the atom (same element, position, and Debye Waller factor). The occupancy is not checked intentionally. If the item is not present a ValueError is raised.

Parameters: item: tuple or list

WyckoffBase entry

Returns: int

static pos_eq (pos1, pos2) compare Wyckoff positions

Parameters: pos1, pos2: tuple

tuples with Wyckoff label and optional parameters

xrayutilities.materials.spacegrouplattice.get_default_sgrp_suf (sgrp_nr)
 determine default space group suffix

xrayutilities.materials.spacegrouplattice.get_possible_sgrp_suf (sgrp_nr)

determine possible space group suffix. Multiple suffixes might be possible for one space group due to different origin choice, unique axis, or choice of the unit cell shape.

Parameters: sgrp_nr: int

space group number

Returns: str or list

either an empty string or a list of possible valid suffix strings

xrayutilities.materials.spacegrouplattice.get_wyckpos (sgrp, atompos) test all Wyckoff positions on every atomic position

Parameters: sgrp: str

space group name

atompos: list

list of atomic positions to identify. All atomic positions are expected to belong to one and

the same Wyckoff position!

Returns: position argument for WyckoffBase.append

xrayutilities.materials.spacegrouplattice.testwp (parint, wp, cifpos, digits=8) test if a Wyckoff position can describe the given position from a CIF file

Parameters: parint: int

telling which Parameters the given Wyckoff position has

wp: str or tuple

expression of the Wyckoff position

cifpos: list, or tuple or array-like

(x, y, z) position of the atom in the CIF file

digits: int

number of digits for which for a comparison of floating point numbers will be rounded to.

By default xu.config.DIGITS is used.

Returns: foundflag: bool

flag to tell if the positions match

pars: array-like or None

parameters associated with the position or None if no parameters are needed

xrayutilities.materials.wyckpos module

<code>class</code> xrayutilities.materials.wyckpos.RangeDict() -> new empty dictionary dict(mapping) -> new dictionary initialized from a mapping object's (key, value) pairs dict(iterable) -> new dictionary initialized as if via: $d = \{\}$ for k, v in iterable: d[k] = v dict(**kwargs) -> new dictionary initialized with the name=value pairs in the keyword argument list. For example: dict(one=1, two=2)

Bases: dict

Module contents

xrayutilities.math package

Submodules

xrayutilities.math.algebra module

module providing analytic algebraic functions not implemented in scipy or any other dependency of xrayutilities. In particular the analytic solution of a quartic equation which is needed for the solution of the dynamic scattering equations.

```
xrayutilities.math.algebra.solve_quartic (a4, a3, a2, a1, a0) analytic solution [1] of the general quartic equation. The solved equation takes the form a \cdot z^4 + a \cdot z^3 + a \cdot z^2 + a \cdot z + a \cdot z
```

Returns: tuple

tuple of the four (complex) solutions of aboves equation.

References

http://mathworld.wolfram.com/QuarticEquation.html

1

xrayutilities.math.fit module

module with a function wrapper to scipy.optimize.leastsq for fitting of a 2D function to a peak or a 1D Gauss fit with the odr package

xrayutilities.math.fit.fit_peak2d (x, y, data, start, drange, fit_function, maxfev=2000)

fit a two dimensional function to a two dimensional data set e.g. a reciprocal space map

Parameters: x, y: array-like

data coordinates (do NOT need to be regularly spaced)

data: array-like

data set used for fitting (e.g. intensity at the data coords)

start: list

set of starting parameters for the fit used as first parameter of function fit_function

drange: list

limits for the data ranges used in the fitting algorithm, e.g. it is clever to use only a small region around the peak which should be fitted: [xmin, xmax, ymin, ymax]

fit_function: callable

function which should be fitted, must be of form accept the parameters $fit_function (x, y, *params) \rightarrow ndarray$

Returns: fitparam: list

fitted parameters

cov: array-like

covariance matrix

xrayutilities.math.fit.gauss_fit (xdata, ydata, iparams=[], maxit=300)

Gauss fit function using odr-pack wrapper in scipy similar to https://github.com/tiagopereira/python_tips/wiki/Scipy%3A-curve-fitting

Parameters: xdata: array-like

x-coordinates of the data to be fitted

ydata: array-like

y-coordinates of the data which should be fit

iparams: list, optional

initial paramters for the fit, determined automatically if not given

maxit: int, optional

maximal iteration number of the fit

Returns: params: list

the parameters as defined in function Gauss1d(x, *param)

sd_params: list

For every parameter the corresponding errors are returned.

itlim: bool

flag to tell if the iteration limit was reached, should be False

xrayutilities.math.fit.linregress(x, y)

fast linregress to avoid usage of scipy.stats which is slow! NaN values in y are ignored by this function.

Parameters: x, y : array-like

data coordinates and values

Returns: p:tuple

parameters of the linear fit (slope, offset)

rsq: float R^2 value

Examples

```
>>> (k, d), R2 = xu.math.linregress(x, y)
```

xrayutilities.math.fit.multPeakFit (x, data, peakpos, peakwidth, dranges=None, peaktype='Gaussian',
returnerror=False)

function to fit multiple Gaussian/Lorentzian peaks with linear background to a set of data

Parameters: x: array-like

x-coordinate of the data

data: array-like

data array with same length as x

peakpos: list

initial parameters for the peak positions

peakwidth: list

initial values for the peak width

dranges: list of tuples

list of tuples with (min, max) value of the data ranges to use. does not need to have the same number of entries as peakpos

peaktype: {'Gaussian', 'Lorentzian'}

type of peaks to be used

returnerror: bool

decides if the fit errors of pos, sigma, and amp are returned (default: False)

Returns: pos: list

peak positions derived by the fit

sigma: list

peak width derived by the fit

amp: list

amplitudes of the peaks derived by the fit

background: array-like

background values at positions x

if returnerror == True:

sd pos: list

standard error of peak positions as returned by scipy.odr.Output

sd_sigma: list

standard error of the peak width

sd_amp: list

standard error of the peak amplitude

xrayutilities.math.fit.multPeakPlot (x, fpos, fwidth, famp, background, dranges=None, peaktype='Gaussian', fig='xu_plot', ax=None, fact=1.0)

function to plot multiple Gaussian/Lorentz peaks with background values given by an array

Parameters: x: array-like

x-coordinate of the data

fpos: list

positions of the peaks

fwidth: list

width of the peaks

famp: list

amplitudes of the peaks

background: array-like

background values, same shape as x

dranges: list of tuples

list of (min, max) values of the data ranges to use. does not need to have the same number of entries as fpos

peaktype : {'Gaussian', 'Lorentzian'}

type of peaks to be used

fig: int, str, or None

matplotlib figure number or name

ax: matplotlib.Axes

matplotlib axes as alternative to the figure name

fact: float

factor to use as multiplicator in the plot

xrayutilities.math.fit.peak_fit (xdata, ydata, iparams=[], peaktype='Gauss', maxit=300,

background='constant', plot=False, func_out=False, debug=False)

fit function using odr-pack wrapper in scipy similar to https://github.com/tiagopereira/python_tips/wiki/Scipy%3A-curve-fitting for Gauss, Lorentz or Pseudovoigt-functions

Parameters: xdata: array_like

x-coordinates of the data to be fitted

ydata: array_like

y-coordinates of the data which should be fit

iparams: list, optional

initial paramters, determined automatically if not specified

peaktype : {'Gauss', 'Lorentz', 'PseudoVoigt', 'PseudoVoigtAsym', 'PseudoVoigtAsym2'},

optional

type of peak to fit

maxit: int, optional

maximal iteration number of the fit

background: {'constant', 'linear'}, optional

type of background function

plot: bool or str, optional

flag to ask for a plot to visually judge the fit. If plot is a string it will be used as figure name, which makes reusing the figures easier.

func out: bool, optional

returns the fitted function, which takes the independent variables as only argument (f(x))

Returns: params: list

the parameters as defined in function *Gauss1d/Lorentz1d/PseudoVoigt1d/PseudoVoigt1dasym*. In the case of linear background one more parameter is included!

sd_params: list

For every parameter the corresponding errors are returned.

itlim: bool

flag to tell if the iteration limit was reached, should be False

fitfunc: function, optional

the function used in the fit can be returned (see func_out).

xrayutilities.math.functions module

module with several common function needed in xray data analysis

xrayutilities.math.functions.Debyel(x)

function to calculate the first Debye function [1] as needed for the calculation of the thermal Debye-Waller-factor by numerical integration

 $D_1(x) = (1/x) \int_0^x t/(\exp(t)-1) dt$

Parameters: x: float

argument of the Debye function

Returns: float

D1(x) float value of the Debye function

References

1 http://en.wikipedia.org/wiki/Debye_function

xrayutilities.math.functions.Gauss1d(x, *p)

function to calculate a general one dimensional Gaussian

Parameters: x: array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the Gaussian [XCEN, SIGMA, AMP, BACKGROUND] for

information: SIGMA = FWHM / (2*sqrt(2*log(2)))

Returns: array-like

the value of the Gaussian described by the parameters p at position x

Examples

Calling with a list of parameters needs a call looking as shown below (note the '*') or explicit listing of the parameters

```
>>> Gauss1d(x,*p)
```

```
>>> Gauss1d(numpy.linspace(0, 10, 100), 5, 1, 1e3, 0)
```

```
xrayutilities.math.functions.Gauss1dArea (*p)
```

function to calculate the area of a Gauss function with neglected background

Parameters: p: list

list of parameters of the Gauss-function [XCEN, SIGMA, AMP, BACKGROUND]

Returns: float

the area of the Gaussian described by the parameters p

```
xrayutilities.math.functions.Gauss1d_der_p (x, *p)
```

function to calculate the derivative of a Gaussian with respect the parameters p for parameter description see Gauss1d

xrayutilities.math.functions.Gauss1d_der_x (x, *p)

```
function to calculate the derivative of a Gaussian with respect to x for parameter description see Gauss1d
```

xrayutilities.math.functions.**Gauss2d** (x, y, *p) function to calculate a general two dimensional Gaussian

Parameters: x, y: array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the Gauss-function [XCEN, YCEN, SIGMAX, SIGMAY, AMP, BACKGROUND, ANGLE]; SIGMA = FWHM / (2*sqrt(2*log(2))); ANGLE = rotation of the

X, Y direction of the Gaussian in radians

Returns: array-like

the value of the Gaussian described by the parameters p at position (x, y)

xrayutilities.math.functions.Gauss2dArea (*p)

function to calculate the area of a 2D Gauss function with neglected background

Parameters: p: list

list of parameters of the Gauss-function [XCEN, YCEN, SIGMAX, SIGMAY, AMP, ANGLE, BACKGROUND]

Returns: float

the area of the Gaussian described by the parameters p

xrayutilities.math.functions.Gauss3d (X, y, z, *p) function to calculate a general three dimensional Gaussian

Parameters: x, y, z : array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the Gauss-function [XCEN, YCEN, ZCEN, SIGMAX, SIGMAY, SIGMAZ, AMP, BACKGROUND];

SIGMA = FWHM / (2*sqrt(2*log(2)))

Returns: array-like

the value of the Gaussian described by the parameters p at positions (x, y, z)

xrayutilities.math.functions.Lorentz1d (x, *p) function to calculate a general one dimensional Lorentzian

Parameters: x: array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the Lorentz-function [XCEN, FWHM, AMP, BACKGROUND]

Returns: array-like

the value of the Lorentian described by the parameters p at position (x, y)

xrayutilities.math.functions.LorentzldArea (*p)

function to calculate the area of a Lorentz function with neglected background

Parameters: p: list

list of parameters of the Lorentz-function [XCEN, FWHM, AMP, BACKGROUND]

Returns: float

the area of the Lorentzian described by the parameters *p*

xrayutilities.math.functions.Lorentz1d_der_p(x, *p)

function to calculate the derivative of a Gaussian with respect the parameters p for parameter description see Lorentz1d

xrayutilities.math.functions.Lorentz1d_der_x(x, *p)

function to calculate the derivative of a Gaussian with respect to x for parameter description see Lorentz1d

xrayutilities.math.functions.Lorentz2d (x, y, *p)

function to calculate a general two dimensional Lorentzian

Parameters: x, y: array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the Lorentz-function [XCEN, YCEN, FWHMX, FWHMY, AMP, BACKGROUND, ANGLE]; ANGLE = rotation of the X, Y direction of the Lorentzian in

radians

Returns: array-like

the value of the Lorentian described by the parameters p at position (x, y)

xrayutilities.math.functions.NormGauss1d (x, *p)

function to calculate a normalized one dimensional Gaussian

Parameters: x: array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the Gaussian [XCEN, SIGMA]; for information: SIGMA = FWHM /

(2*sqrt(2*log(2)))

Returns: array-like

the value of the normalized Gaussian described by the parameters p at position x

xrayutilities.math.functions.NormLorentz1d(x, *p)

function to calculate a normalized one dimensional Lorentzian

Parameters: x: array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the Lorentzian [XCEN, FWHM]

Returns: array-like

the value of the normalized Lorentzian described by the parameters p at position x

xrayutilities.math.functions.PseudoVoigt1d(x, *p)

function to calculate a pseudo Voigt function as linear combination of a Gauss and Lorentz peak

Parameters: x: array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the pseudo Voigt-function [XCEN, FWHM, AMP, BACKGROUND,

ETA]; ETA: 0 ...1 0 means pure Gauss and 1 means pure Lorentz

Returns: array-like

the value of the PseudoVoigt described by the parameters p at position x

xrayutilities.math.functions.**PseudoVoigt1dArea** (*p)

function to calculate the area of a pseudo Voigt function with neglected background

Parameters: p: list

list of parameters of the Lorentz-function [XCEN, FWHM, AMP, BACKGROUND, ETA];

ETA: 0 ... 1 0 means pure Gauss and 1 means pure Lorentz

Returns: float

the area of the PseudoVoigt described by the parameters p

xrayutilities.math.functions.**PseudoVoigt1d_der_p**(x, *p)

function to calculate the derivative of a PseudoVoigt with respect the parameters p for parameter description see PseudoVoigt1d

xrayutilities.math.functions.PseudoVoigt1d der x (x, *p)

function to calculate the derivative of a PseudoVoigt with respect to x

for parameter description see PseudoVoigt1d

xrayutilities.math.functions.**PseudoVoigt1dasym**(x, *p)

function to calculate an asymmetric pseudo Voigt function as linear combination of asymmetric Gauss and Lorentz peak

Parameters: x : array-like

coordinate(s) where the function should be evaluated

list of parameters of the pseudo Voigt-function [XCEN, FWHMLEFT, FWHMRIGHT, AMP, BACKGROUND, ETA]; ETA: 0 ...1 0 means pure Gauss and 1 means pure

Lorentz

array-like Returns:

the value of the PseudoVoigt described by the parameters p at position x

xrayutilities.math.functions.PseudoVoigt1dasym2(x,*p)

function to calculate an asymmetric pseudo Voigt function as linear combination of asymmetric Gauss and Lorentz peak

Parameters: x : naddray

coordinate(s) where the function should be evaluated

p: list

list of parameters of the pseudo Voigt-function [XCEN, FWHMLEFT, FWHMRIGHT, AMP, BACKGROUND, ETALEFT, ETARIGHT]; ETA: 0 ...1 0 means pure Gauss and 1 means pure Lorentz

Returns: array-like

the value of the PseudoVoigt described by the parameters p at position x

xrayutilities.math.functions.**PseudoVoigt2d**(x, y, *p)

function to calculate a pseudo Voigt function as linear combination of a Gauss and Lorentz peak in two dimensions

Parameters: x, y: array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the pseudo Voigt-function [XCEN, YCEN, FWHMX, FWHMY, AMP, BACKGROUND, ANGLE, ETA]; ETA: 0 ... 1 0 means pure Gauss and 1 means pure

Lorentz

array-like Returns:

the value of the PseudoVoigt described by the parameters p at position (x, y)

xrayutilities.math.functions.TwoGauss2d(X, V, *p)

function to calculate two general two dimensional Gaussians

Parameters: x, y : array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the Gauss-function [XCEN1, YCEN1, SIGMAX1, SIGMAY1, AMP1, ANGLE1, XCEN2, YCEN2, SIGMAX2, SIGMAY2, AMP2, ANGLE2, BACKGROUND]; SIGMA = FWHM / (2*sqrt(2*log(2))) ANGLE = rotation of the X, Y direction of the

Gaussian in radians

arrav-like Returns:

the value of the Gaussian described by the parameters p at position (x, y)

xrayutilities.math.functions.heaviside(x)

Heaviside step function for numpy arrays

Parameters: x: scalar or array-like

argument of the step function

Returns: int or array-like

Heaviside step function evaluated for all values of x with datatype integer

xrayutilities.math.functions.kill_spike (data, threshold=2.0, offset=None)

function to smooth **single** data points which differ from the average of the neighboring data points by more than the threshold factor or more than the offset value. Such spikes will be replaced by the mean value of the next neighbors.

Warning

Use this function carefully not to manipulate your data!

Parameters: data: array-like

1d numpy array with experimental data

threshold: float or None

threshold factor to identify outlier data points. If None it will be ignored.

offset: None or float

offset value to identify outlier data points. If None it will be ignored.

Returns: array-like

1d data-array with spikes removed

xrayutilities.math.functions.multPeak1d(x, *args)

function to calculate the sum of multiple peaks in 1D. the peaks can be of different type and a background function (polynom) can also be included.

Parameters: x: array-like

coordinate where the function should be evaluated

args: list

list of peak/function types and parameters for every function type two arguments need to be given first the type of function as string with possible values 'g': Gaussian, 'l': Lorentzian, 'v': PseudoVoigt, 'a': asym. PseudoVoigt, 'p': polynom the second type of arguments is the tuple/list of parameters of the respective function. See documentation of math.Gauss1d, math.Lorentz1d, math.PseudoVoigt1d, math.PseudoVoigt1dasym, and numpy.polyval for details of the different function types.

Returns: array-like

value of the sum of functions at position x

xrayutilities.math.functions.multPeak2d(x, y, *args)

function to calculate the sum of multiple peaks in 2D. the peaks can be of different type and a background function (polynom) can also be included.

Parameters: x, y : array-like

coordinates where the function should be evaluated

args: list

list of peak/function types and parameters for every function type two arguments need to be given first the type of function as string with possible values 'g': Gaussian, 'l': Lorentzian, 'v': PseudoVoigt, 'c': constant the second type of arguments is the tuple/list of parameters of the respective function. See documentation of math.Gauss2d, math.Lorentz2d, math.PseudoVoigt2d for details of the different function types. The constant accepts a single float which will be added to the data

Returns: array-like

value of the sum of functions at position (x, y)

xrayutilities.math.functions.smooth(x, n)

function to smooth an array of data by averaging N adjacent data points

Parameters: x: array-like

1D data array

n:int

number of data points to average

Returns: xsmooth: array-like

smoothed array with same length as x

xrayutilities.math.misc module

xrayutilities.math.misc.center_of_mass (pos, data, background='none', full_output=False)
function to determine the center of mass of an array

Parameters: pos: array-like

position of the data points

data: array-like data values

background : {'none', 'constant', 'linear'}

type of background, either 'none', 'constant' or 'linear'

full_output : bool

return background cleaned data and background-parameters

Returns: float

center of mass position

xrayutilities.math.misc.fwhm_exp (pos, data)

function to determine the full width at half maximum value of experimental data. Please check the obtained value visually (noise influences the result)

Parameters: pos: array-like

position of the data points

data: array-like data values

Returns: float

fwhm value

xrayutilities.math.misc.gcd (lst)

greatest common divisor function using library functions

Parameters: Ist: array-like

array of integer values for which the greatest common divisor should be determined

Returns: gcd: int

xrayutilities.math.transforms module

xrayutilities.math.transforms.ArbRotation (axis, alpha, deg=True)

Returns a transform that represents a rotation around an arbitrary axis by the angle alpha. positive rotation is anti-clockwise when looking from positive end of axis vector

Parameters: axis: list or array-like

rotation axis

alpha: float

rotation angle in degree (deg=True) or in rad (deg=False)

deg: bool

determines the input format of ang (default: True)

Returns: **Transform**

class xrayutilities.math.transforms.AxisToZ (newzaxis)

Bases: xrayutilities.math.transforms.CoordinateTransform

Creates a coordinate transformation to move a certain axis to the z-axis. The rotation is done along the great circle. The x-axis of the new coordinate frame is created to be normal to the new and original z-axis. The new y-axis is create in order to obtain a right handed coordinate system.

class xrayutilities.math.transforms.AxisToZ_keepXY (newzaxis)

Bases: xrayutilities.math.transforms.CoordinateTransform

Creates a coordinate transformation to move a certain axis to the z-axis. The rotation is done along the great circle. The x-axis/y-axis of the new coordinate frame is created to be similar to the old x and y directions. This variant of AxisToZ assumes that the new Z-axis has its main component along the Z-direction

class xrayutilities.math.transforms.CoordinateTransform (V1, V2, V3)

Bases: xrayutilities.math.transforms.Transform

Create a Transformation object which transforms a point into a new coordinate frame. The new frame is determined by the three vectors v1/norm(v1), v2/norm(v2) and v3/norm(v3), which need to be orthogonal!

class xrayutilities.math.transforms.Transform (matrix)

Bases: object

property imatrix

inverse (args, rank=1)

performs inverse transformation a vector, matrix or tensor of rank 4

Parameters: args: list or array-like

object to transform, list or numpy array of shape (..., n) (..., n, n), (..., n, n, n, n, n) where n is the size of the transformation matrix.

rank: int

rank of the supplied object, allowed values are 1, 2, and 4

xrayutilities.math.transforms.XRotation (alpha, deg=True)

Returns a transform that represents a rotation about the x-axis by an angle alpha. If deg=True the angle is assumed to be in degree, otherwise the function expects radiants.

xrayutilities.math.transforms.YRotation (alpha, deg=True)

Returns a transform that represents a rotation about the y-axis by an angle alpha. If deg=True the angle is assumed to be in degree, otherwise the function expects radiants.

xrayutilities.math.transforms.ZRotation (alpha, deg=True)

Returns a transform that represents a rotation about the z-axis by an angle alpha. If deg=True the angle is assumed to be in degree, otherwise the function expects radiants.

xrayutilities.math.transforms.mycross (vec, mat)

function implements the cross-product of a vector with each column of a matrix

xrayutilities.math.transforms.rotarb (vec, axis, ang, deg=True)

function implements the rotation around an arbitrary axis by an angle ang positive rotation is anti-clockwise when looking from positive end of axis vector

Parameters: vec: list or array-like

vector to rotate

axis: list or array-like
rotation axis

ang: float

rotation angle in degree (deg=True) or in rad (deg=False)

deg: bool

determines the input format of ang (default: True)

Returns: rotvec: rotated vector as numpy.array

Examples

```
>>> rotarb([1, 0, 0],[0, 0, 1], 90)
array([ 6.12323400e-17,  1.00000000e+00,  0.00000000e+00])
```

xrayutilities.math.transforms.tensorprod (vec1, vec2) function implements an elementwise multiplication of two vectors

xrayutilities.math.vector module

module with vector operations for vectors of size 3, since for so short vectors numpy does not give the best performance explicit implementation of the equations is performed together with error checking to ensure vectors of length 3.

```
xrayutilities.math.vector.VecAngle ((v1.v2)/(norm(v1)*norm(v2))) alpha = acos((v1.v2)/(norm(v1)*norm(v2)))
```

Parameters: v1, v2 : list or array-like

input vector(s), either one vector or an array of vectors with shape (n, 3)

deg: bool, optional

True: return result in degree, False: in radiants (default: False)

Returns: float or ndarray

the angle included by the two vectors v1 and v2, either a single float or an array with shape (n,)

xrayutilities.math.vector.VecCross (v1, v2, out=None)

Calculate the vector cross product.

Parameters: v1, v2 : list or array-like

input vector(s), either one vector or an array of vectors with shape (n, 3)

out: list or array-like, optional

output vector

Returns: ndarray

cross product either of shape (3,) or (n, 3)

xrayutilities.math.vector.VecDot (v1, v2)

Calculate the vector dot product.

Parameters: v1, v2 : list or array-like

input vector(s), either one vector or an array of vectors with shape (n, 3)

Returns: float or ndarray

innter product of the vectors, either a single float or (n,)

xrayutilities.math.vector.VecNorm(V)

Calculate the norm of a vector.

Parameters: v: list or array-like

input vector(s), either one vector or an array of vectors with shape (n, 3)

Returns: float or ndarray

vector norm, either a single float or shape (n,)

xrayutilities.math.vector.VecUnit (v)

Calculate the unit vector of v.

Parameters: v: list or array-like

input vector(s), either one vector or an array of vectors with shape (n, 3)

Returns: ndarray

unit vector of v, either shape (3,) or (n, 3)

xrayutilities.math.vector.distance (x, y, z, point, vec)

calculate the distance between the point (x, y, z) and the line defined by the point and vector vec

Parameters: x: float or ndarray

x coordinate(s) of the point(s)

y: float or ndarray

y coordinate(s) of the point(s)

z: float or ndarray

z coordinate(s) of the point(s)

point: tuple, list or ndarray

3D point on the line to which the distance should be calculated

vec: tuple, list or ndarray

3D vector defining the propergation direction of the line

xrayutilities.math.vector.getSyntax (Vec)

returns vector direction in the syntax 'x+' 'z-' or equivalents therefore works only for principle vectors of the coordinate system like e.g. [1, 0, 0] or [0, 2, 0]

Parameters: vec : list or array-like

vector of length 3

Returns: str

vector string following the synthax [xyz][+-]

xrayutilities.math.vector.getVector (string)

returns unit vector along a rotation axis given in the syntax 'x+' 'z-' or equivalents

Parameters: string: str

vector string following the synthax [xyz][+-]

Returns: ndarray

vector along the given direction

Module contents

xrayutilities.simpack package

Submodules

xrayutilities.simpack.darwin_theory module

class xrayutilities.simpack.darwin_theory.DarwinModel (qz, qx=0, qy=0, **kwargs)
Bases: xrayutilities.simpack.models.LayerModel

model class inmplementing the basics of the Darwin theory for layers materials. This class is not fully functional and should be used to derive working models for particular material systems.

To make the class functional the user needs to implement the init_structurefactors() and _calc_mono() methods

init_structurefactors()

calculates the needed atomic structure factors

ncalls = 0

simulate (MI)

main simulation function for the Darwin model. will calculate the reflected intensity

Parameters: ml: iterable

monolayer sequence of the sample. This should be created with the function

make_monolayer(). see its documentation for details

class xrayutilities.simpack.darwin_theory.DarwinModelAlGaAs001 (qz, qx=0, qy=0, **kwargs)

Bases: xrayutilities.simpack.darwin_theory.DarwinModelAlloy

Darwin theory of diffraction for Al_x Ga_{1-x} As layers. The model is based on separation of the sample structure into building blocks of atomic planes from which a multibeam dynamical model is calculated.

Alas = <xrayutilities.materials.material.Crystal object>

GaAs = <xrayutilities.materials.material.Crystal object>

aGaAs = 5.65325

classmethod abulk (x)

calculate the bulk (relaxed) lattice parameter of the Al_{x}Ga_{1-x}As alloy

asub = 5.65325

eAl = AI (13)

eAs = As (33)

eGa = Ga(31)

classmethod get_dperp_apar (x, apar, r=1)

calculate inplane lattice parameter and the out of plane lattice plane spacing (of the atomic planes!) from composition and relaxation

Parameters: x: float

chemical composition parameter

apar: float

inplane lattice parameter of the material below the current layer (onto which the present layer is strained to). This value also served as a reference for the relaxation parameter.

r · floot

relaxation parameter. 1=relaxed, 0=pseudomorphic

Returns: dperp: float

perpendicular d-spacing

apar: float

inplane lattice parameter

init_structurefactors (temp=300)

calculates the needed atomic structure factors

Parameters: temp: float, optional

temperature used for the Debye model

static poisson_ratio (x)

calculate the Poisson ratio of the alloy

```
re = 2.8179403262e-05
```

class xrayutilities.simpack.darwin_theory.DarwinModelAlloy (qz, qx=0, qy=0, **kwargs)

Bases: xrayutilities.simpack.darwin_theory.DarwinModel, abc.ABC

extension of the DarwinModel for an binary alloy system were one parameter is used to determine the chemical composition

To make the class functional the user needs to implement the get_dperp_apar() method and define the substrate lattice parameter (asub). See the DarwinModelSiGe001 class for an implementation example.

abstract get_dperp_apar (x, apar, r=1)

calculate inplane lattice parameter and the out of plane lattice plane spacing (of the atomic planes!) from composition and relaxation.

Parameters: x: float

chemical composition parameter

apar: float

inplane lattice parameter of the material below the current layer (onto which the present layer is strained to). This value also served as a reference for the relaxation parameter.

r: float

relaxation parameter. 1=relaxed, 0=pseudomorphic

Returns: dperp: float

apar: float

make_monolayers (S)

create monolayer sequence from layer list

Parameters: s: list

layer model. list of layer dictionaries including possibility to form superlattices. As an example 5 repetitions of a Si(10nm)/Ge(15nm) superlattice on Si would like like:

the dictionaries must contain 't': thickness in A, 'x': chemical composition, and either 'r': relaxation or 'ai': inplane lattice parameter. Future implementations for asymmetric peaks might include layer type 'l' (not yet implemented). Already now any additional property in the dictionary will be handed on to the returned monolayer list.

asub: float

inplane lattice parameter of the substrate

Returns: list

monolayer list in a format understood by the simulate and xGe profile methods

prop profile (ml, prop)

calculate the profile of chemical composition or inplane lattice spacing from a monolayer list. One value for each monolayer in the sample is returned.

Parameters: ml: list

monolayer list created by make_monolayer()

prop: str

name of the property which should be evaluated. Use 'x' for the chemical composition

and 'ai' for the inplane lattice parameter.

Returns: zm: ndarray

z-position, z-0 is the surface

propx: ndarray

value of the property prop for every monolayer

class xrayutilities.simpack.darwin_theory.DarwinModelGaInAs001 (qz, qx=0, qy=0, **kwargs)

Bases: xrayutilities.simpack.darwin_theory.DarwinModelAlloy

Darwin theory of diffraction for Ga_{1-x} In_x As layers. The model is based on separation of the sample structure into building blocks of atomic planes from which a multibeam dynamical model is calculated.

GaAs = <xrayutilities.materials.material.Crystal object>

InAs = <xrayutilities.materials.material.Crystal object>

aGaAs = 5.65325

classmethod abulk (x)

calculate the bulk (relaxed) lattice parameter of the Ga_{1-x}ln_{x}As alloy

asub = 5.65325

eAs = As (33)

eGa = Ga (31)

eIn = ln (49)

classmethod get_dperp_apar (x, apar, r=1)

calculate inplane lattice parameter and the out of plane lattice plane spacing (of the atomic planes!) from composition and relaxation

Parameters: x: float

chemical composition parameter

apar: float

inplane lattice parameter of the material below the current layer (onto which the present layer is strained to). This value also served as a reference for the relaxation parameter.

r: float

relaxation parameter. 1=relaxed, 0=pseudomorphic

Returns: dperp: float

perpendicular d-spacing

apar: float

inplane lattice parameter

init_structurefactors (temp=300)

calculates the needed atomic structure factors

Parameters: temp: float, optional

temperature used for the Debye model

static poisson_ratio (x)

calculate the Poisson ratio of the alloy

```
re = 2.8179403262e-05
```

class xrayutilities.simpack.darwin_theory.DarwinModelSiGe001 (qz, qx=0, qy=0, **kwargs)

Bases: xrayutilities.simpack.darwin_theory.DarwinModelAlloy

model class implementing the Darwin theory of diffraction for SiGe layers. The model is based on separation of the sample structure into building blocks of atomic planes from which a multibeam dynamical model is calculated.

Ge = <xrayutilities.materials.material.Crystal object>

Si = <xrayutilities.materials.material.Crystal object>

aSi = 5.43104

classmethod abulk (x)

calculate the bulk (relaxed) lattice parameter of the alloy

asub = 5.43104

eGe = Ge (32)

esi = Si(14)

classmethod get_dperp_apar (x, apar, r=1)

calculate inplane lattice parameter and the out of plane lattice plane spacing (of the atomic planes!) from composition and relaxation

Parameters: x: float

chemical composition parameter

apar: float

inplane lattice parameter of the material below the current layer (onto which the present layer is strained to). This value also served as a reference for the relaxation parameter.

r: float, optional

relaxation parameter. 1=relaxed, 0=pseudomorphic

Returns: dperp: float

perpendicular d-spacing

apar: float

inplane lattice parameter

init_structurefactors (temp=300)

calculates the needed atomic structure factors

Parameters: temp: float, optional

temperature used for the Debye model

static poisson_ratio (x)

calculate the Poisson ratio of the alloy

re = 2.8179403262e-05

xrayutilities.simpack.darwin_theory.**GradedBuffer** (xfrom, xto, nsteps, thickness, relaxation=1) create a multistep graded composition buffer.

Parameters: xfrom: float

begin of the composition gradient

xto: float

end of the composition gradient

nsteps: int

number of steps of the gradient

thickness: float

total thickness of the Buffer in A

relaxation: float

relaxation of the buffer

Returns: list

layer list needed for the Darwin model simulation

xrayutilities.simpack.darwin_theory.getfirst (iterable, key)

helper function to obtain the first item in a nested iterable

xrayutilities.simpack.darwin_theory.getit (it, key)

generator to obtain items from nested iterable

xrayutilities.simpack.fit module

class xrayutilities.simpack.fit.FitModel (Imodel, verbose=False, plot=False, elog=True, **kwargs)

Bases: object

Wrapper for the Imfit Model class working for instances of LayerModel

Typically this means that after initialization of *FitModel* you want to use make_params to get a *Imfit.Parameters* list which one customizes for fitting.

Later on you can call fit and eval methods with those parameter list.

fit (data, params, x, weights=None, fit_kws=None, **kwargs)

wrapper around Imfit.Model.fit which enables plotting during the fitting

Parameters: data: ndarray

experimental values
params: Imfit.Parameters

list of parameters for the fit, use make_params for generation

x: ndarray

independent variable (incidence angle or q-position depending on the model)

weights: ndarray, optional

values of weights for the fit, same size as data

fit_kws : dict, optional

Options to pass to the minimizer being used

kwargs: dict, optional

keyword arguments which are passed to Imfit.Model.fit

Returns: Imfit.ModelResult

set_fit_limits (xmin=- inf, xmax=inf, mask=None)

set fit limits. If mask is given it must have the same size as the *data* and *x* variables given to fit. If mask is None it will be generated from xmin and xmax.

Parameters: xmin: float, optional

minimum value of x-values to include in the fit

xmax: float, optional

maximum value of x-values to include in the fit

mask: boolean array, optional

mask to be used for the data given to the fit

xrayutilities.simpack.helpers module

xrayutilities.simpack.helpers.coplanar_alphai (qx, qz, en='config') calculate coplanar incidence angle from knowledge of the qx and qz coordinates

Parameters: qx: array-like

inplane momentum transfer component

qz: array-like

out of plane momentum transfer component

en: float or str, optional

x-ray energy (eV). By default the value from the config is used.

Returns: alphai: array-like

the incidence angle in degree. points in the Laue zone are set to 'nan'.

xrayutilities.simpack.helpers.get_qz (qx, alphai, en='config')

calculate the qz position from the qx position and the incidence angle for a coplanar diffraction geometry

Parameters: qx : array-like

inplane momentum transfer component

alphai: array-like

incidence angle (deg)

en: float or str, optional

x-ray energy (eV). By default the value from the config is used.

Returns: array-like

the qz position for the given incidence angle

xrayutilities.simpack.models module

class xrayutilities.simpack.models.DiffuseReflectivityModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.SpecularReflectivityModel

model for diffuse reflectivity calculations

The 'simulate' method calculates the diffuse reflectivity on the specular rod in coplanar geometry in analogy to the SpecularReflectivityModel.

The 'simulate_map' method calculates the diffuse reflectivity for a 2D set of Q-positions. This method can also calculate the intensity for other geometries, like GISAXS with constant incidence angle or a quasi omega/2theta scan in GISAXS geometry.

simulate (alphai)

performs the actual diffuse reflectivity calculation for the specified incidence angles. This method always uses the coplanar geometry independent of the one set during the initialization.

Parameters: alphai: array-like

vector of incidence angles

Returns: array-like

vector of intensities of the reflectivity signal

simulate map (qL, qz)

performs diffuse reflectivity calculation for the rectangular grid of reciprocal space positions define by qL and qz. This method uses the method and geometry set during the initialization of the class.

Parameters: qL: array-like

lateral coordinate in reciprocal space (vector with NqL components)

qz: array-like

vertical coordinate in reciprocal space (vector with Ngz components)

Returns: array-like

matrix of intensities of the reflectivity signal, with shape (len(qL), len(qz))

class xrayutilities.simpack.models.DynamicalModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.SimpleDynamicalCoplanarModel

Dynamical diffraction model for specular and off-specular qz-scans. Calculation of the flux of reflected and diffracted waves for general asymmetric coplanar diffraction from an arbitrary pseudomorphic multilayer is performed by a generalized 2-beam theory (4 tiepoints, S and P polarizations)

The first layer in the model is always assumed to be the semiinfinite substrate indepentent of its given thickness

simulate (alphai, hkl=None, geometry='hi_lo', rettype='intensity')

performs the actual diffraction calculation for the specified incidence angles and uses an analytic solution for the quartic dispersion equation

Parameters: alphai: array-like

vector of incidence angles (deg)

hkl: list or tuple, optional

Miller indices of the diffraction vector (preferable use set_hkl method to speed up repeated calculations of the same peak!)

geometry: {'hi_lo', 'lo_hi'}, optional

'hi_lo' for grazing exit (default) and 'lo_hi' for grazing incidence

rettype: {'intensity', 'field', 'all'}, optional

type of the return value. 'intensity' (default): returns the diffracted beam flux convoluted with the resolution function; 'field': returns the electric field (complex) without convolution with the resolution function, 'all': returns the electric field, ai, af (both in degree), and the reflected intensity.

Returns: array-like

vector of intensities of the diffracted signal, possibly changed return value due the rettype setting!

class xrayutilities.simpack.models.DynamicalReflectivityModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.SpecularReflectivityModel

model for Dynamical Specular Reflectivity Simulations. It uses the transfer Matrix methods as given in chapter 3 "Daillant, J., & Gibaud, A. (2008). X-ray and Neutron Reflectivity"

scanEnergy (energies, angle)

Simulates the Dynamical Reflectivity as a function of photon energy at fixed angle.

Parameters: energies: numpy.ndarray or list

photon energies (in eV).

angle: float

fixed incidence angle

Returns: reflectivity: array-like

vector of intensities of the reflectivity signal

transmitivity: array-like

vector of intensities of the transmitted signal

simulate (alphai)

Simulates the Dynamical Reflectivity as a function of angle of incidence

Parameters: alphai : array-like

vector of incidence angles

Returns: reflectivity: array-like

vector of intensities of the reflectivity signal

transmitivity: array-like

vector of intensities of the transmitted signal

class xrayutilities.simpack.models.KinematicalModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.LayerModel

Kinematical diffraction model for specular and off-specular qz-scans. The model calculates the kinematical contribution of one (hkl) Bragg peak, however considers the variation of the structure factor for different 'q'. The surface geometry is specified using the Experiment-object given to the constructor.

init_chi0()

calculates the needed optical parameters for the simulation. If any of the materials/layers is changing its properties this function needs to be called again before another correct simulation is made. (Changes of thickness does NOT require this!)

simulate (qz, hkl, absorption=False, refraction=False, rettype='intensity')

performs the actual kinematical diffraction calculation on the Qz positions specified considering the contribution from a single Bragg peak.

Parameters: qz: array-like

simulation positions along qz

hkl: list or tuple

Miller indices of the Bragg peak whos truncation rod should be calculated

absorption: bool, optional

flag to tell if absorption correction should be used

refraction: bool, optional

flag to tell if basic refraction correction should be performed. If refraction is True absorption correction is also included independent of the absorption flag.

rettype: {'intensity', 'field', 'all'}

type of the return value. 'intensity' (default): returns the diffracted beam flux convoluted with the resolution function; 'field': returns the electric field (complex) without convolution with the resolution function, 'all': returns the electric field, ai, af (both in degree), and the reflected intensity.

Returns: array-like

return value depends on the setting of *rettype*, by default only the calculate intensity is returned

class xrayutilities.simpack.models.KinematicalMultiBeamModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.KinematicalModel

Kinematical diffraction model for specular and off-specular qz-scans. The model calculates the kinematical contribution of several Bragg peaks on the truncation rod and considers the variation of the structure factor. In order to use a analytical description for the kinematic diffraction signal all layer thicknesses are changed to a multiple of the respective lattice parameter along qz. Therefore this description only works for (001) surfaces.

simulate (qz, hkl, absorption=False, refraction=True, rettype='intensity')

performs the actual kinematical diffraction calculation on the Qz positions specified considering the contribution from a full truncation rod

Parameters: qz: array-like

simulation positions along qz

hkl: list or tuple

Miller indices of the Bragg peak whos truncation rod should be calculated

absorption: bool, optional

flag to tell if absorption correction should be used

refraction: bool, optional,

flag to tell if basic refraction correction should be performed. If refraction is True absorption correction is also included independent of the absorption flag.

rettype: {'intensity', 'field', 'all'}

type of the return value. 'intensity' (default): returns the diffracted beam flux convoluted with the resolution function; 'field': returns the electric field (complex) without convolution with the resolution function, 'all': returns the electric field, ai, af (both in degree), and the reflected intensity.

Returns: array-like

> return value depends on the setting of rettype, by default only the calculate intensity is returned

class xrayutilities.simpack.models.LayerModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.Model, abc.ABC

generic model class from which further thin film models can be derived from

get_polarizations()

return list of polarizations which should be calculated

join polarizations (Is, Ip)

method to calculate the total diffracted intensity from the intensities of S and P-polarization.

abstract simulate ()

abstract method that every implementation of a LayerModel has to override.

class xrayutilities.simpack.models.Model (experiment, **kwargs)

Bases: object

generic model class from which further models can be derived from

convolute resolution (X, y)

convolve simulation result with a resolution function

Parameters: x: array-like

x-values of the simulation, units of x also decide about the unit of the resolution width parameter

y: array-like

y-values of the simulation

Returns: array-like

convoluted y-data with same shape as y

property energy

scale_simulation(y)

scale simulation result with primary beam flux/intensity and add a background.

Parameters: y: array-like

y-values of the simulation

array-like Returns:

scaled y-values

class xrayutilities.simpack.models.ResonantReflectivityModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.SpecularReflectivityModel

model for specular reflectivity calculations CURRENTLY UNDER DEVELOPEMENT! DO NOT USE!

simulate (alphai)

performs the actual reflectivity calculation for the specified incidence angles

Parameters: alphai: array-like

vector of incidence angles

Returns: array-like

vector of intensities of the reflectivity signal

class xrayutilities.simpack.models.SimpleDynamicalCoplanarModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.KinematicalModel

Dynamical diffraction model for specular and off-specular qz-scans. Calculation of the flux of reflected and diffracted waves for general asymmetric coplanar diffraction from an arbitrary pseudomorphic multilayer is performed by a simplified 2-beam theory (2 tiepoints, S and P polarizations)

No restrictions are made for the surface orientation.

The first layer in the model is always assumed to be the semiinfinite substrate indepentent of its given thickness

Note

This model should not be used in real life scenarios since the made approximations severely fail for distances far from the reference position.

set_hkl (*hkl)

To speed up future calculations of the same Bragg peak optical parameters can be pre-calculated using this function.

Parameters: hkl : list or tuple

Miller indices of the Bragg peak for the calculation

simulate (alphai, hkl=None, geometry='hi_lo', idxref=1)

performs the actual diffraction calculation for the specified incidence angles.

Parameters: alphai: array-like

vector of incidence angles (deg)

hkl: list or tuple, optional

Miller indices of the diffraction vector (preferable use set_hkl method to speed up repeated calculations of the same peak!)

geometry: {'hi_lo', 'lo_hi'}, optional

'hi_lo' for grazing exit (default) and 'lo_hi' for grazing incidence

idxref: int, optional

index of the reference layer. In order to get accurate peak position of the film peak you want this to be the index of the film peak (default: 1). For the substrate use 0.

Returns: array-like

vector of intensities of the diffracted signal

class xrayutilities.simpack.models.SpecularReflectivityModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.LayerModel

model for specular reflectivity calculations

densityprofile (nz, plot=False)

calculates the electron density of the layerstack from the thickness and roughness of the individual layers

Parameters: nz: int

number of values on which the profile should be calculated

plot: bool, optional

flag to tell if a plot of the profile should be created

Returns: z: array-like

z-coordinates, z = 0 corresponds to the surface

eprof : array-like
 electron profile

init_cd()

calculates the needed optical parameters for the simulation. If any of the materials/layers is changing its properties this function needs to be called again before another correct simulation is made. (Changes of thickness and roughness do NOT require this!)

simulate (alphai)

performs the actual reflectivity calculation for the specified incidence angles

Parameters: alphai: array-like

vector of incidence angles

Returns: array-like

vector of intensities of the reflectivity signal

xrayutilities.simpack.models.effectiveDensitySlicing (layerstack, step, roughness=0, cutoff=1e-05)

Function to slice a LayerStack into many amorphous sublayers for effective density modelling of X-ray reflectivity of thin and rough multilayers. The resulting LayerStack will consist of perfectly smooth layers with average density/composition resulting from an error-function like transition between the rough layers of the initial stack. At the surface an vacuum layer is automatically added to the initial stack.

Parameters: layerstack: initial LayerStack, can contain only Amorhous layers!

step: thickness (in Angstrom) of the slices in the returned

LayerStack

roughness: roughness of the created sublayers (in Angstrom)

cutoff: layers with relative weights below this value will be ignored

Returns: LayerStack

xrayutilities.simpack.models.startdelta (start, delta, num)

xrayutilities.simpack.mosaicity module

xrayutilities.simpack.mosaicity.mosaic_analytic (qx, qz, RL, RV, Delta, hx, hz, shape) simulation of the coplanar reciprocal space map of a single mosaic layer using a simple analytic approximation

Parameters: qx: array-like

vector of the qx values (offset from the Bragg peak)

qz: array-like

vector of the gz values (offset from the Bragg peak)

RL: float

lateral block radius in Angstrom

RV: float

vertical block radius in Angstrom

Delta: float

root mean square misorientation of the grains in degree

hx: float

lateral component of the diffraction vector

hz: float

vertical component of the diffraction vector

shape: float

shape factor (1..Gaussian)

Returns: array-like

2D array with calculated intensities

xrayutilities.simpack.powder module

This module contains the core definitions for the XRD Fundamental Parameneters Model (FPA) computation in Python. The main computational class is FP_profile, which stores cached information to allow it to efficiently recompute profiles when parameters have been modified. For the user an Powder class is available which can calculate a complete powder pattern of a crystalline material.

The diffractometer line profile functions are calculated by methods from Cheary & Coelho 1998 and Mullen & Cline paper and 'R' package. Accumulate all convolutions in Fourier space, for efficiency, except for axial divergence, which needs to be weighted in real space for I3 integral.

More details about the applied algorithms can be found in the paper by M. H. Mendelhall et al., Journal of Research of NIST 120, 223 (2015) to which you should also refer for a careful definition of all the parameters

class xrayutilities.simpack.powder.FP_profile (anglemode, gaussian_smoother_bins_sigma=1.0,
oversampling=10)

Bases: object

the main fundamental parameters class, which handles a single reflection. This class is designed to be highly extensible by inheriting new convolvers. When it is initialized, it scans its namespace for specially formatted names, which can come from mixin classes. If it finds a function name of the form conv_xxx, it will call this funtion to create a convolver. If it finds a name of the form info_xxx it will associate the dictionary with that convolver, which can be used in UI generation, for example. The class, as it stands, does nothing significant with it. If it finds str_xxx, it will use that function to format a printout of the current state of the convolver conv_xxx, to allow improved report generation for convolvers.

When it is asked to generate a profile, it calls all known convolvers. Each convolver returns the Fourier transform of its convolvution. The transforms are multiplied together, inverse transformed, and after fixing the periodicity issue, subsampled, smoothed and returned.

If a convolver returns None, it is not multipled into the product.

Parameters: max history length: int

the number of histories to cache (default=5); can be overridden if memory is an issue.

length_scale_m : float

length_scale_m sets scaling for nice printing of parameters. if the units are in mm everywhere, set it to 0.001, e.g. convolvers which implement their own str xxx method may use this to format their results, especially if 'natural' units are not meters. Typical is

wavelengths and lattices in nm or angstroms, for example.

add buffer (b)

add a numpy array to the list of objects that can be thrown away on pickling.

Parameters: b : array-like

the buffer to add to the list

Returns: **b**: array-like

return the same buffer, to make nesting easy.

axial_helper (outerbound, innerbound, epsvals, destination, peakpos=0, y0=0, k=0)

the function F0 from the paper. compute k/sqrt(peakpos-x)+y0 nonzero between outer & inner (inner is closer to peak) or k/sqrt(x-peakpos)+y0 if reversed (i.e. if outer > peak) fully evaluated on a specified eps grid, and stuff into destination

Parameters: outerbound: float

the edge of the function farthest from the singularity, referenced to epsvals

innerbound: float

the edge closest to the singularity, referenced to epsvals

epsvals: array-like

the array of two-theta values or offsets

destination : array-like

an array into which final results are summed. modified in place!

peakpos: float

the position of the singularity, referenced to epsvals.

y0: float

the constant offset

k: float

the scale factor

Returns: lower index, upper index: int

python style bounds for region of destination which has been modified.

compute_line_profile (convolver_names=None, compute_derivative=False, return_convolver=False) execute all the convolutions; if convolver_names is None, use everything we have, otherwise, use named convolutions.

Parameters: convolver names: list

a list of convolvers to select. If None, use all found convolvers.

compute_derivative: bool

if True, also return d/dx(function) for peak position fitting

Returns:

a profile_data object with much information about the peak

conv absorption ()

compute the sample transparency correction, including the finite-thickness version

Returns: array-like

the convolver

conv axial()

compute the Fourier transform of the axial divergence component

Returns: array-like

the transform buffer, or None if this is being ignored

conv_displacement ()

compute the peak shift due to sample displacement and the 2theta zero offset

Returns: array-like

the convolver

conv_emission()

compute the emission spectrum and (for convenience) the particle size widths

Returns: array-like

the convolver for the emission and particle sizes

Note

the particle size and strain stuff here is just to be consistent with *Topas* and to be vaguely efficient about the computation, since all of these have the same general shape.

conv_flat_specimen()

compute the convolver for the flat-specimen correction

Returns: array-like

the convolver

conv_global()

a dummy convolver to hold global variables and information. the global context isn't really a convolver, returning *None* means ignore result

Returns: None

always returns None

conv_receiver_slit()

compute the rectangular convolution for the receiver slit or SiPSD pixel size

Returns: array-like

the convolver

conv_si_psd()

compute the convolver for the integral of defocusing of the face of an Si PSD

Returns: array-like

the convolver

conv_smoother ()

compute the convolver to smooth the final result with a Gaussian before downsampling.

Returns: array-like

the convolver

conv_tube_tails()

compute the Fourier transform of the rectangular tube tails function

Returns: array-like

the transform buffer, or None if this is being ignored

full_axdiv_I2 (Lx=None, Ls=None, Lr=None, R=None, twotheta=None, beta=None, epsvals=None)
return the I2 function

Parameters: Lx: float

length of the xray filament

Ls: float

length of the sample

Lr: float

length of the receiver slit

R: float

diffractometer length, assumed symmetrical

twotheta: float

angle, in radians, of the center of the computation

beta: *float* offset angle

epsvals: array-like
array of offsets from center of computation, in radians

Returns: epsvals: array-like

array of offsets from center of computation, in radians

idxmin, idxmax: int

the full python-style bounds of the non-zero region of I2p and I2m

I2p, I2m: array-like

12+ and 12- from the paper, the contributions to the intensity

full_axdiv_I3 (Lx=None, Ls=None, Lr=None, R=None, twotheta=None, epsvals=None, sollerIdeg=None,
sollerDdeg=None, nsteps=10, axDiv=")

carry out the integral of I2 over beta and the Soller slits.

Parameters: Lx: float

length of the xray filament

Ls: float

length of the sample

Lr: float

length of the receiver slit

R: float

the (assumed symmetrical) diffractometer radius

twotheta: float

angle, in radians, of the center of the computation

epsvals: array-like

array of offsets from center of computation, in radians

sollerideg: float

the full-width (both sides) cutoff angle of the incident Soller slit

sollerDdeg: float

the full-width (both sides) cutoff angle of the detector Soller slit

nsteps: int

the number of subdivisions for the integral

axDiv : str not used

Returns: array-like

the accumulated integral, a copy of a persistent buffer _axial

general_tophat (name=", width=None)

a utility to compute a transformed tophat function and save it in a convolver buffer

Parameters: name: str

the name of the convolver cache buffer to update

width: float

the width in 2-theta space of the tophat

Returns: array-like

the updated convolver buffer, or None if the width was None

get_conv (name, key, format=<class 'float'>)

get a cached, pre-computed convolver associated with the given parameters, or a newly zeroed convolver if the cache doesn't contain it. Recycles old cache entries.

This takes advantage of the mutability of arrays. When the contents of the array are changed by the convolver, the cached copy is implicitly updated, so that the next time this is called with the same parameters, it will return the previous array.

Parameters: name: str

the name of the convolver to seek

key: object

any hashable object which identifies the parameters for the computation

format: numpy.dtype, optional

the type of the array to create, if one is not found.

Returns: bool

flag, which is *True* if valid data were found, or *False* if the returned array is zero, and

array, which must be computed by the convolver if flag was False.

get_convolver_information()

return a list of convolvers, and what we know about them. function scans for functions named conv_xxx, and associated info_xxx entries.

Returns: list

list of (convolver_xxx, info_xxx) pairs

get_function_name ()

return the name of the function that called this. Useful for convolvers to identify themselves

Returns: str

name of calling function

get_good_bin_count (count)

find a bin count close to what we need, which works well for Fourier transforms.

Parameters: count: int

a number of bins.

Returns: int

a bin count somewhat larger than count which is efficient for FFT

info_emission = {'group_name': 'Incident beam and crystal size', 'help': 'this should be help information', 'param_info': {'crystallite_size_gauss': ('Gaussian crystallite size fwhm (m)', 1e-06), 'crystallite_size_lor': ('Lorentzian crystallite size fwhm (m)', 1e-06), 'emiss_gauss_widths': ('Gaussian emissions fwhm (m)', (1e-13,)), 'emiss_intensities': ('relative intensities', (1.0,)), 'emiss_lor_widths': ('Lorenztian emission fwhm (m)', (1e-13,)), 'emiss_wavelengths': ('wavelengths (m)', (1.58e-10,))}}

info_global = {'group_name': 'Global parameters', 'help': 'this should be help information', 'param_info': {'d': ('d spacing (m)', 4e-10), 'dominant_wavelength': ('wavelength of most intense line (m)', 1.5e-10), 'twotheta0_deg': ('Bragg center of peak (degrees)', 30.0)}}

classmethod isequivalent (hkl1, hkl2, crystalsystem)

function to determine if according to the convolvers included in this class two sets of Miller indices are equivalent. This function is only called when the class attribute 'isotropic' is False.

Parameters: hkl1, hkl2: list or tuple

Miller indices to be checked for equivalence

crystalsystem: str

symmetry class of the material which is considered

Returns: bool

isotropic = True

length scale m = 1.0

max_history_length = 5

self_clean()

do some cleanup to make us more compact; Instance can no longer be used after doing this, but can be pickled.

set_optimized_window (twotheta_window_center_deg, twotheta_approx_window_fullwidth_deg,
twotheta exact bin spacing deg)

pick a bin count which factors cleanly for FFT, and adjust the window width to preserve the exact center and bin spacing

Parameters: twotheta window center deg: float

exact position of center bin, in degrees

twotheta_approx_window_fullwidth_deg: float

approximate desired width

twotheta_exact_bin_spacing_deg: float

the exact bin spacing to use

set_parameters (convolver='global', **kwargs)

update the dictionary of parameters associated with the given convolver

Parameters: convolver: str

the name of the convolver. name 'global', e.g., attaches to function 'conv_global'

kwargs: dict

keyword-value pairs to update the convolvers dictionary.

set_window (twotheta_window_center_deg, twotheta_window_fullwidth_deg, twotheta_output_points)
move the compute window to a new location and compute grids, without resetting all parameters. Clears convolution history and sets up many arrays.

Parameters: twotheta_window_center_deg: float

the center position of the middle bin of the window, in degrees

twotheta_window_fullwidth_deg: float the full width of the window, in degrees

twotheta_output_points: int

the number of bins in the final output

str_emission()

format the emission spectrum and crystal size information

Returns: str

the formatted information

str_global()

returns a string representation for the global context.

Returns: str

report on global parameters.

class xrayutilities.simpack.powder.PowderDiffraction (mat, **kwargs)

Bases: xrayutilities.experiment.PowderExperiment

Experimental class for powder diffraction. This class calculates the structure factors of powder diffraction lines and uses instances of FP_profile to perform the convolution with experimental resolution function calculated by the fundamental parameters approach. This class uses multiprocessing to speed up calculation. Set config.NTHREADS=1 to restrict this to one worker process.

Calculate (twotheta, **kwargs)

calculate the powder diffraction pattern including convolution with the resolution function and map them onto the twotheta positions. This also performs the calculation of the peak intensities from the internal material object

Parameters: twotheta: array-like

two theta values at which the powder pattern should be calculated.

kwargs: dict

additional keyword arguments are passed to the Convolve function

Returns: array-like

output intensity values for the twotheta values given in the input

Notes

Bragg peaks are only included up to tt cutoff set in the class constructor!

Convolve (twotheta, window_width='config', mode='multi')

convolute the powder lines with the resolution function and map them onto the twotheta positions. This calculates the powder pattern excluding any background contribution

Parameters: twotheta: array-like

two theta values at which the powder pattern should be calculated.

window_width: float, optional

width of the calculation window of a single peak

mode: {'multi, 'local'}, optional

multiprocessing mode, either 'multi' to use multiple processes or 'local' to restrict the calculation to a single process

Note:

Bragg peaks are only included up to tt cutoff set in the class constructor!

Returns: output intensity values for the twotheta values given in the input

close()

correction_factor (ang)

calculate the correction factor for the diffracted intensities. This contains the polarization effects and the Lorentz factor

Parameters: ang: aray-like

theta diffraction angles for which the correction should be calculated

Returns: f: array-like

array of the same shape as ang containing the correction factors

property energy

init_powder_lines (tt_cutoff)

calculates the powder intensity and positions up to an angle of tt_cutoff (deg) and stores the result in the data dictionary whose structure is as follows:

The data dictionary has one entry per line with a unique identifier as key of the entry. The entries themself are dictionaries which have the following entries:

- hkl : (h, k, l), Miller indices of the Bragg peak
- r : reflection strength of the line
- ang : Bragg angle of the peak (theta = 2theta/2!)
- · qpos: reciprocal space position

load_settings_from_config (settings)

load parameters from the config and update these settings with the options from the settings parameter

merge_lines (data)

if calculation is isotropic lines at the same q-position can be merged to one line to reduce the calculational effort

Parameters: data: ndarray

numpy field array with values of 'hkl' (Miller indices of the peaks), 'q' (q-position), and 'r' (reflection strength) as produced by the *reflection_strength* method

Returns: hkl, q, ang, r : array-like

Miller indices, q-position, diffraction angle (Theta), and reflection strength of the

material

reflection_strength (tt_cutoff)

determine structure factors/reflection strength of all Bragg peaks up to tt_cutoff. This function also implements the March-Dollase model for preferred orientation in the symmetric reflection mode. Note that although this means the sample has anisotropic properties the various lines can still be merged together since at the moment no anisotropic crystal shape is supported.

Parameters: tt_cutoff: float

upper cutoff value of 2theta until which the reflection strength are calculated

Returns: ndarray

numpy array with field for 'hkl' (Miller indices of the peaks), 'q' (q-position), and 'r'

(reflection strength) of the Bragg peaks

set_sample_parameters()

load sample parameters from the Powder class and use them in all FP profile instances of this object

set_wavelength_from_params()

sets the wavelenth in the base class from the settings dictionary of the FP_profile classes and also set it in the 'global' part of the parameters

set window (force=False)

sets the calcultion window for all convolvers

property twotheta

update_powder_lines (tt_cutoff)

calculates the powder intensity and positions up to an angle of tt_cutoff (deg) and updates the values in:

- ids: list of unique identifiers of the powder line
- · data: array with intensities
- ang: bragg angles of the peaks (theta=2theta/2!)
- qpos: reciprocal space position of intensities

update_settings (newsettings={})

update settings of all instances of FP_profile

Parameters:

newsettings: dict

which should have its settings changed.

```
property wavelength
  property window_width
xrayutilities.simpack.powder.chunkify(lst, n)
class xrayutilities.simpack.powder.convolver_handler
  Bases: object
  manage the convolvers of on process
  add_convolver (convolver)
  calc (run, ttpeaks)
    calculate profile function for selected convolvers
        Parameters: run: list
                          list of flags of length of convolvers to tell which convolver needs to be run
                      ttpeaks: array-like
                          peak positions for the convolvers
           Returns:
                      list
                          list of profile data result objects
  set_windows (centers, npoints, flag, width)
  update_parameters (parameters)
class xrayutilities.simpack.powder.manager (address=None, authkey=None, serializer='pickle', ctx=None)
  Bases: multiprocessing.managers.BaseManager
class xrayutilities.simpack.powder.profile_data (**kwargs)
  Bases: object
  a skeleton class which makes a combined dict and namespace interface for easy pickling and data passing
  add_symbol (**kwargs)
    add new symbols to both the attributes and dictionary for the class
        Parameters: kwargs: dict
                          keyword=value pairs
xrayutilities.simpack.powdermodel module
class xrayutilities.simpack.powdermodel.PowderModel (*args, **kwargs)
  Bases: object
  Class to help with powder calculations for multiple materials. For basic calculations the Powder class together with
  the Fundamental parameters approach is used.
  close()
  create_fitparameters()
    function to create a fit model with all instrument and sample parameters.
           Returns: Imfit.Parameters
```

dictionary with new settings. It has to include one subdictionary for every convolver

fit (params, twotheta, data, std=None, maxfev=200)
make least squares fit with parameters supplied by the user

Parameters: params: Imfit.Parameters

object with all parameters set as intended by the user

twotheta: array-like

angular values for the fit

data: array-like

experimental intensities for the fit

std: array-like

standard deviation of the experimental data. if 'None' the sqrt of the data will be used

maxfev: int

maximal number of simulations during the least squares refinement

Returns: Imfit.MinimizerResult

plot (twotheta, showlines=True, label='simulation', color=None, formatspec='-', lcolors=[], ax=None, **kwargs) plot the powder diffraction pattern and indicate line positions for all components in the model.

Parameters: twotheta: array-like

positions at which the powder pattern should be evaluated

showlines: bool, optional

flag to decide if peak positions of the components will be shown on the top of the plot

label: str

line label in the plot

color: matplotlib color or None

the color used for the line plot of the simulation

formatspec : str

format specifier of the simulation curve

Icolors: list of matplotlib colors

colors for the line indicators for the various components

ax: matplotlib.axes or None

axes object to be used for plotting, if its given no axes decoration like labels are set.

Further keyword arguments are passed to the simulate method.

Returns: matplotlib.axes object

set_background (btype, **kwargs)
define background as spline or polynomial function

Parameters: btype: {polynomial', 'spline'}

background type; Depending on this value the expected keyword arguments differ.

kwargs: dict

optional keyword arguments

x: array-like, optional

x-values (twotheta) of the background points (if btype='spline')

y: array-like, optional

intensity values of the background (if btype='spline')

p: array-like, optional

polynomial coefficients from the highest degree to the constant term. len of p decides about the degree of the polynomial (if btype='polynomial')

set_lmfit_parameters (Imparams)

function to update the settings of this class during an least squares fit

Parameters: Imparams: Imfit.Parameters

Imfit Parameters list of sample and instrument parameters

set_parameters (params)

set simulation parameters of all subobjects

Parameters: params: dict

settings dictionaries for the convolvers.

simulate (twotheta, **kwargs)

calculate the powder diffraction pattern of all materials and sum the results based on the relative volume of the materials.

Parameters: twotheta: array-like

positions at which the powder pattern should be evaluated

kwargs: dict

optional keyword arguments

background: array-like

an array of background values (same shape as twotheta) if no background is given then the background is calculated as previously set by the set_background function or is 0

further keyword arguments are passed to the Convolve function of of the

PowderDiffraction objects

Returns: array-like

summed powder diffraction intensity of all materials present in the model

xrayutilities.simpack.powdermodel.Rietveld_error_metrics (exp, sim, weight=None, std=None,
Nvar=0, disp=False)

calculates common error metrics for Rietveld refinement.

Parameters: exp: array-like

experimental datapoints

sim : array-like simulated data

weight: array-like, optional

weight factor in the least squares sum. If it is None the weight is estimated from the counting statistics of 'exp'

std: array-like, optional

standard deviation of the experimental data. alternative way of specifying the weight factor, when both are given weight overwrites std!

Nvar: int, optional

number of variables in the refinement

disp: bool, optional

flag to tell if a line with the calculated values should be printed.

Returns: M, Rp, Rwp, Rwpexp, chi2: float

xrayutilities.simpack.powdermodel.plot_powder (twotheta, exp, sim, mask=None, scale='sqrt',
fig='XU:powder', show_diff=True, show_legend=True, labelexp='experiment', labelsim='simulation', formatexp='.-k',
formatsim='-r')

Convenience function to plot the comparison between experimental and simulated powder diffraction data

Parameters: twotheta: array-like

angle values used for the x-axis of the plot (deg)

exp: array-like

experimental data (same shape as twotheta). If None only the simulation and no difference will be plotted

sim: array-like or PowederModel

simulated data or PowderModel instance. If a PowderModel instance is given the plot-method of PowderModel is used.

mask: array-like, optional

mask to reduce the twotheta values to the be used as x-coordinates of sim

scale: {'linear', 'sqrt', 'log'}, optional

string specifying the scale of the y-axis.

fig: str or int, optional

matplotlib figure name (figure will be cleared!)

show_diff: bool, optional

flag to specify if a difference curve should be shown

show_legend: bool, optional

flag to specify if a legend should be shown

labelexp: str

plot label (legend entry) for the experimental data

labelsim: str

plot label for the simulation data

formatexp: str

format specifier for the experimental data

formatsim: str

format specifier for the simulation curve

xrayutilities.simpack.smaterials module

```
class xrayutilities.simpack.smaterials.CrystalStack (name, *args)
  Bases: xrayutilities.simpack.smaterials.LayerStack
  extends the built in list type to enable building a stack of crystalline Layers by various methods.
  check (V)
class xrayutilities.simpack.smaterials.GradedLayerStack (alloy, xfrom, xto, nsteps, thickness,
**kwargs)
  Bases: xrayutilities.simpack.smaterials.CrystalStack
  generates a sequence of layers with a gradient in chemical composition
class xrayutilities.simpack.smaterials.Layer (material, thickness, **kwargs)
  Bases: xrayutilities.simpack.smaterials.SMaterial
  Object describing part of a thin film sample. The properties of a layer are :
                     material: Material (Crystal or Amorhous)
                         an xrayutilties material describing optical and crystal properties of the thin film
                     thickness: float
                         film thickness in Angstrom
class xrayutilities.simpack.smaterials.LayerStack (name, *args)
  Bases: xrayutilities.simpack.smaterials.MaterialList
  extends the built in list type to enable building a stack of Layer by various methods.
  check (v)
class xrayutilities.simpack.smaterials.MaterialList (name, *args)
  Bases: collections.abc.MutableSequence
  class representing the basics of a list of materials for simulations within xrayutilities. It extends the built in list type.
  check (V)
  insert (i, v)
    S.insert(index, value) – insert value before index
class xrayutilities.simpack.smaterials.Powder (material, volume, **kwargs)
  Bases: xrayutilities.simpack.smaterials.SMaterial
  Object describing part of a powder sample. The properties of a powder are:
        Attributes:
                     material: Crystal
                         an xrayutilties material (Crystal) describing optical and crystal properties of the powder
                     volume: float
                         powder's volume (in pseudo units, since only the relative volume enters the calculation)
                     crystallite_size_lor: float, optional
                         Lorentzian crystallite size fwhm (m)
                     crystallite_size_gauss: float, optional
                         Gaussian crystallite size fwhm (m)
                     strain_lor : float, optional
                         extra peak width proportional to tan(theta)
                     strain gauss: float, optional
                         extra peak width proportional to tan(theta)
                     preferred_orientation: tuple, optional
                         HKL of the preferred orientation
                     preferred_orientation_factor : float, optional
                         March-Dollase preferred orientation factor: > 1 for platy crystallits , < 1 for rod-like
                         crystallites, and = 1 for random orientation of crystallites.
```

```
class xrayutilities.simpack.smaterials.PowderList (name, *args)
 Bases: xrayutilities.simpack.smaterials.MaterialList
 extends the built in list type to enable building a list of Powder by various methods.
 check (V)
class xrayutilities.simpack.smaterials.PseudomorphicStack001 (name, *args)
 Bases: xrayutilities.simpack.smaterials.CrystalStack
 generate a sequence of pseudomorphic crystalline Layers. Surface orientation is assumed to be 001 and materials
 must be cubic/tetragonal.
 insert (i, v)
   S.insert(index, value) - insert value before index
 make_epitaxial(i)
 trans = <xrayutilities.math.transforms.Transform object>
class xrayutilities.simpack.smaterials.PseudomorphicStack111 (name, *args)
 Bases: xrayutilities.simpack.smaterials.PseudomorphicStack001
 generate a sequence of pseudomorphic crystalline Layers. Surface orientation is assumed to be 111 and materials
 must be cubic.
 trans = <xrayutilities.math.transforms.CoordinateTransform object>
```

Simulation Material. Extends the xrayutilities Materials by properties needed for simulations property material

Module contents

Bases: object

simulation subpackage of xrayutilities.

This package provides possibilities to simulate X-ray diffraction and reflectivity curves of thin film samples. It could be extended for more general use in future if there is demand for that.

In addition it provides a fitting routine for reflectivity data which is based on Imfit.

class xrayutilities.simpack.smaterials.SMaterial (material, **kwargs)

Submodules

xrayutilities.config module

module to parse xrayutilities user-specific config file the parsed values are provide as global constants for the use in other parts of xrayutilities. The config file with the default constants is found in the python installation path of xrayutilities. It is however not recommended to change things there, instead the user-specific config file ~/.xrayutilities.conf or the local xrayutilities.conf file should be used.

xrayutilities.config.trytomake (obj, key, typefunc)

xrayutilities.exception module

xrayutilities derives its own exceptions which are raised upon wrong input when calling one of xrayutilities functions. none of the pre-defined exceptions is made for that purpose.

```
exception xrayutilities.exception.InputError (msg)
```

Bases: Exception

Exception raised for errors in the input. Either wrong datatype not handled by TypeError or missing mandatory keyword argument (Note that the obligation to give keyword arguments might depend on the value of the arguments itself)

Parameters: expr:str

input expression in which the error occurred

msg: str

explanation of the error

xrayutilities.experiment module

module helping with planning and analyzing experiments. various classes are provided for describing experimental geometries, calculationof angular coordinates of Bragg reflections, conversion of angular coordinates to Q-space and determination of powder diffraction peak positions.

The strength of the module is the versatile QConversion module which can be configured to describe almost any goniometer geometry.

class xrayutilities.experiment.Experiment (ipdir, ndir, **keyargs)

Bases: object

base class for describing experiments users should use the derived classes: HXRD, GID, PowderExperiment

Ang2HKL (*args, **kwargs)

angular to (h, k, l) space conversion. It will set the UB argument to Ang2Q and pass all other parameters unchanged. See Ang2Q for description of the rest of the arguments.

Parameters: args: list

arguments forwarded to Ang2Q

kwargs: dict, optional

optional keyword arguments

B: array-like, optional

reciprocal space conversion matrix of a Crystal. You can specify the matrix B (default identity matrix) shape needs to be (3, 3)

mat: Crystal, optional

Crystal object to use to obtain a B matrix (e.g. xu.materials.Si) can be used as alternative to the B keyword argument B is favored in case both are given

U: array-like, optional

orientation matrix U can be given. If none is given the orientation defined in the Experiment class is used.

dettype: {'point', 'linear', 'area'}, optional

detector type: decides which routine of Ang2Q to call. default 'point'

delta: ndarray, list or tuple, optional

giving delta angles to correct the given ones for misalignment. delta must be an numpy array or list of length 2. used angles are than (om, tt) - delta

wl: float or str, optional

x-ray wavelength in angstroem (default: self. wl)

en: float or str, optional

x-ray energy in eV (default: converted self._wl)

deg: bool, optional

flag to tell if angles are passed as degree (default: True)

sampledis: tuple, list or array-like, optional

sample displacement vector in relative units of the detector distance (default: (0, 0, 0))

Returns: ndarray

H K L coordinates as numpy.ndarray with shape (N, 3) where N corresponds to the number of points given in the input (args)

Q2Ang (qvec)

TiltAngle (q, deg=True)

TiltAngle(q, deg=True): Return the angle between a q-space position and the surface normal.

Parameters: q: list or numpy array with the reciprocal space position

optional keyword arguments:

deg : True/False whether the return value should be in degree or radians (default: True)

Transform (V)

transforms a vector, matrix or tensor of rank 4 (e.g. elasticity tensor) to the coordinate frame of the Experiment class. This is for example necessary before any Q2Ang-conversion can be performed.

Parameters: v: object to transform, list or numpy array of shape

(n,) (n, n), (n, n, n, n) where n is the rank of the transformation matrix

Returns: transformed object of the same shape as v

property energy

property wavelength

class xrayutilities.experiment.FourC (idir, ndir, **keyargs)

Bases: xrayutilities.experiment.HXRD

class describing high angle x-ray diffraction experiments the class helps with calculating the angles of Bragg reflections as well as helps with analyzing measured data

the class describes a four circle (omega, chi, phi, twotheta) goniometer to help with coplanar x-ray diffraction experiments. Nevertheless 3D data can be treated with the use of linear and area detectors. see help self.Ang2Q

class xrayutilities.experiment.GID (idir, ndir, **keyargs)

Bases: xrayutilities.experiment.Experiment

class describing grazing incidence x-ray diffraction experiments the class helps with calculating the angles of Bragg reflections as well as it helps with analyzing measured data

the class describes a four circle (alpha_i, azimuth, twotheta, beta) goniometer to help with GID experiments at the ROTATING ANODE. 3D data can be treated with the use of linear and area detectors. see help self.Ang2Q Using this class the default sample surface orientation is determined by the inner most sample rotation (which is

usually the azimuth motor).

Ang2Q (ai, phi, tt, beta, **kwargs)

angular to momentum space conversion for a point detector. Also see help GID.Ang2Q for procedures which treat line and area detectors

Parameters: ai, phi, tt, beta: float or array-like

sample and detector angles as numpy array, lists or Scalars must be given. All arguments must have the same shape or length. However, if one angle is always the same its enough to give one scalar value.

kwargs: dict, optional

optional keyword arguments

delta: list, tuple or array-like, optional

giving delta angles to correct the given ones for misalignment delta must be an numpy array or list of length 4. Used angles are then ai, phi, tt, beta - delta

UB: array-like, optional

matrix for conversion from (hkl) coordinates to Q of sample used to determine not Q but (hkl) (default: identity matrix)

wl: float or str, optional

x-ray wavelength in angstroem (default: self. wl)

deg: bool, optional

flag to tell if angles are passed as degree (default: True)

Returns: ndarray

reciprocal space positions as numpy.ndarray with shape (N, 3) where N corresponds to the number of points given in the input

Q2Ang (Q, trans=True, deg=True, **kwargs)

calculate the GID angles needed in the experiment the inplane reference direction defines the direction were the reference direction is parallel to the primary beam (i.e. lattice planes perpendicular to the beam)

Note

The behavior of this function is unchanged if the goniometer definition is changed!

Parameters: Q: list, tuple or array-like

array of shape (3) with q-space vector components or 3 separate lists with qx, qy, qz

trans: bool, optional

apply coordinate transformation on Q (default True)

deg: book, optional

(default True) determines if the angles are returned in radians or degrees

Returns: ndarray

a numpy array of shape (4) with four GID scattering angles which are [alpha_i, azimuth, twotheta, beta];

- alpha_i : incidence angle to surface (at the moment always 0)
- azimuth : sample rotation with respect to the inplane reference direction
- twotheta : scattering angle
- beta : exit angle from surface (at the moment always 0)

class xrayutilities.experiment.GISAXS (idir, ndir, **keyargs)

Bases: xrayutilities.experiment.Experiment

class describing grazing incidence x-ray diffraction experiments the class helps with calculating the angles of Bragg reflections as well as it helps with analyzing measured data

the class describes a three circle (alpha_i, twotheta, beta) goniometer to help with GISAXS experiments at the ROTATING ANODE. 3D data can be treated with the use of linear and area detectors. see help self.Ang2Q

Ang2Q (ai, tt, beta, **kwargs)

angular to momentum space conversion for a point detector. Also see help GISAXS.Ang2Q for procedures which treat line and area detectors

Parameters: ai, tt, beta: float or array-like

sample and detector angles as numpy array, lists or Scalars must be given. all arguments must have the same shape or length. Howevver, if one angle is always the same its enough to give one scalar value.

kwargs: dict, optional

optional keyword arguments

delta: list, tuple or array-like, optional

giving delta angles to correct the given ones for misalignment delta must be an numpy array or list of length 3. Used angles are then ai, tt, beta - delta

UB: array-like, optional

matrix for conversion from (hkl) coordinates to Q of sample used to determine not Q but (hkl) (default: identity matrix)

wl: float or str, optional

x-ray wavelength in angstroem (default: self._wl)

deg: bool, optional

flag to tell if angles are passed as degree (default: True)

Returns: ndarray

reciprocal space positions as numpy.ndarray with shape (N, 3) where N corresponds to the number of points given in the input

Q2Ang (Q, trans=True, deg=True, **kwargs)

class xrayutilities.experiment.HXRD (idir, ndir, geometry='hi_lo', **keyargs)

Bases: xrayutilities.experiment.Experiment

class describing high angle x-ray diffraction experiments the class helps with calculating the angles of Bragg reflections as well as helps with analyzing measured data

the class describes a two circle (omega, twotheta) goniometer to help with coplanar x-ray diffraction experiments. Nevertheless 3D data can be treated with the use of linear and area detectors. see help self.Ang2Q

Ang2Q (om, tt, **kwargs)

angular to momentum space conversion for a point detector. Also see help HXRD.Ang2Q for procedures which treat line and area detectors

Parameters: om, tt: float or array-like

sample and detector angles as numpy array, lists or Scalars must be given. All arguments must have the same shape or length. However, if one angle is always the same its enough to give one scalar value.

kwargs: dict, optional

optional keyword arguments

delta: list or array-like

giving delta angles to correct the given ones for misalignment. delta must be an numpy array or list of length 2. Used angles are than om, tt - delta

UB: array-like

matrix for conversion from (hkl) coordinates to Q of sample used to determine not Q but (hkl) (default: identity matrix)

wl: float or str, optional

x-ray wavelength in angstroem (default: self._wl)

deg: bool, optional

flag to tell if angles are passed as degree (default: True)

Returns: ndarray

reciprocal space positions as numpy.ndarray with shape (N, 3) where N corresponds to the number of points given in the input

Q2Ang (*Q, **keyargs)

Convert a reciprocal space vector Q to COPLANAR scattering angles. The keyword argument trans determines whether Q should be transformed to the experimental coordinate frame or not. The coplanar scattering angles correspond to a goniometer with sample rotations ['x+', 'y+', 'z-'] and detector rotation 'x+' and primary beam along y. This is a standard four circle diffractometer.

Note

The behavior of this function is unchanged if the goniometer definition is changed!

Parameters: Q: list, tuple or array-like

array of shape (3) with q-space vector components or 3 separate lists with qx, qy, qz

trans: bool, optional

apply coordinate transformation on Q (default True)

deg: book, optional

(default True) determines if the angles are returned in radians or degrees

geometry: {'hi_lo', 'lo_hi', 'real', 'realTilt'}, optional

determines the scattering geometry (default: self.geometry):

- 'hi_lo' high incidence and low exit
- 'lo_hi' low incidence and high exit
- 'real' general geometry with angles determined by q-coordinates (azimuth); this and upper geometries return [omega, 0, phi, twotheta]
- 'realTilt' general geometry with angles determined by q-coordinates (tilt); returns [omega, chi, phi, twotheta]

refrac: bool, optional

determines if refraction is taken into account; if True then also a material must be given (default: False)

mat: Crystal

Crystal object; needed to obtain its optical properties for refraction correction, otherwise not used

full_output: bool, optional

determines if additional output is given to determine scattering angles more accurately in case refraction is set to True. default: False

fi, fd: tuple or list

if refraction correction is applied one can optionally specify the facet through which the beam enters (fi) and exits (fd) fi, fd must be the surface normal vectors (not transformed & not necessarily normalized). If omitted the normal direction of the experiment is used. Returns: ndarray

full_output=False: a numpy array of shape (4) with four scattering angles which are [omega, chi, phi, twotheta];

- omega: incidence angle with respect to surface
- chi : sample tilt for the case of non-coplanar geometry
- phi : sample azimuth with respect to inplane reference direction
- twotheta : scattering angle/detector angle

full_output=True: a numpy array of shape (6) with five angles which are [omega, chi, phi, twotheta, psi_i, psi_d]

- psi_i : offset of the incidence beam from the scattering plane due to refraction
- pdi_d : offset ot the diffracted beam from the scattering plane due to refraction

class xrayutilities.experiment.NonCOP (idir, ndir, **keyargs)

Bases: xrayutilities.experiment.Experiment

class describing high angle x-ray diffraction experiments. The class helps with calculating the angles of Bragg reflections as well as helps with analyzing measured data for NON-COPLANAR measurements, where the tilt is used to align asymmetric peaks, like in the case of a polefigure measurement.

The class describes a four circle (omega, chi, phi, twotheta) goniometer to help with x-ray diffraction experiments. Linear and area detectors can be treated as described in "help self.Ang2Q"

Ang2Q (om, chi, phi, tt, **kwargs)

angular to momentum space conversion for a point detector. Also see help NonCOP.Ang2Q for procedures which treat line and area detectors

Parameters: om, chi, phi, tt: float or array-like

sample and detector angles as numpy array, lists or Scalars must be given. All arguments must have the same shape or length. However, if one angle is always the same its enough to give one scalar value.

kwargs: dict, optional

optional keyword arguments

delta: list, tuple or array-like, optional

giving delta angles to correct the given ones for misalignment delta must be an numpy array or list of length 4. Used angles are than om, chi, phi, tt - delta

UB: array-like, optional

matrix for conversion from (hkl) coordinates to Q of sample used to determine not Q but (hkl) (default: identity matrix)

wl: float or str, optional

x-ray wavelength in angstroem (default: self._wl)

deg: bool, optional

flag to tell if angles are passed as degree (default: True)

Returns: ndarray

reciprocal space positions as numpy.ndarray with shape (N, 3) where N corresponds to the number of points given in the input

Q2Ang (*Q, **keyargs)

Convert a reciprocal space vector Q to NON-COPLANAR scattering angles. The keyword argument trans determines whether Q should be transformed to the experimental coordinate frame or not.

Note

The behavior of this function is unchanged if the goniometer definition is changed!

Parameters: Q: list, tuple or array-like

array of shape (3) with q-space vector components or 3 separate lists with qx, qy, qz

trans: bool, optional

apply coordinate transformation on Q (default True)

deg: book, optional

(default True) determines if the angles are returned in radians or degrees

Returns:

a numpy array of shape (4) with four scattering angles which are [omega, chi, phi, twotheta];

- omega: incidence angle with respect to surface
- chi : sample tilt for the case of non-coplanar geometry
- phi : sample azimuth with respect to inplane reference direction
- twotheta: scattering angle/detector angle

class xrayutilities.experiment.PowderExperiment (**kwargs)

Bases: xrayutilities.experiment.Experiment

Experimental class for powder diffraction which helps to convert theta angles to momentum transfer space

Q2Ang (qpos, wl=None, deg=True)

Converts reciprocal space values to theta angles

class xrayutilities.experiment.QConversion (sampleAxis, detectorAxis, r_i, **kwargs)

Bases: object

Class for the conversion of angular coordinates to momentum space for arbitrary goniometer geometries and X-ray energy. Both angular scans (where some goniometer angles change during data acquisition) and energy scans (where the energy is varied during acquisition) as well as mixed cases can be treated.

the class is configured with the initialization and does provide three distinct routines for conversion to momentum space for

- point detector: point(...) or __call__()
- linear detector: linear(...)

• area detector: area(...) linear() and area() can only be used after the init_linear() or init_area() routines were called

property UB

area (*args, **kwargs)

angular to momentum space conversion for a area detector the center pixel defined by the init area routine must be in direction of self.r_i when detector angles are zero

the detector geometry must be initialized by the init_area(...) routine

Parameters: args: ndarray, list or Scalars

sample and detector angles; in total *len(self.sampleAxis)* + *len(detectorAxis)* must be given, always starting with the outer most circle. all arguments must have the same shape or length but can be mixed with Scalars (i.e. if an angle is always the same it can be given only once instead of an array)

• sAngles:

sample circle angles, number of arguments must correspond to len(self.sampleAxis)

• dAngles:

detector circle angles, number of arguments must correspond to len(self.detectorAxis)

kwargs: dict, optional

optional keyword arguments

delta: list or array-like, optional

delta angles to correct the given ones for misalignment. delta must be an numpy array or list of len(*args). used angles are then *args - delta

UB: array-like, optional

matrix for conversion from (hkl) coordinates to Q of sample used to determine not Q but (hkl) (default: self.UB)

Nav: tuple or list, optional

number of channels to average to reduce data size e.g. [2, 2] (default: self._area_nav)

roi: list or tuple, optional

region of interest for the detector pixels; e.g. [100, 900, 200, 800] (default: self._area_roi)

wl: float or str, optional

x-ray wavelength in angstroem (default: self._wl)

en: float, optional

x-ray energy in eV (default is converted self._wl). both wavelength and energy can also be an array which enables the QConversion for energy scans. Note that the *en* keyword overrules the *wl* keyword!

deq: bool, optional

flag to tell if angles are passed as degree (default: True)

sampledis: tuple or list or array-like

sample displacement vector in relative units of the detector distance (default: (0, 0, 0))

Returns: reciprocal space position of all detector pixels in a numpy.ndarray of

shape ((*)*(self._area_roi[1] - self._area_roi[0]+1) *

(self._area_roi[3] - self._area_roi[2] + 1), 3) were detectorDir1 is

the fastest varing

property detectorAxis

property handler for _detectorAxis

Returns: list of detector axis following the syntax /[xyz][+-]/

property energy

getDetectorDistance (*args, **kwargs)

obtains the detector distance by applying the detector arm movements. This is especially interesting for the case of 1 or 2D detectors to perform certain geometric corrections.

Parameters: args: list

detector angles. Only detector arm angles as described by the detectorAxis attribute must be given.

kwargs: dict, optional

optional keyword arguments

dim: int, optional

dimension of the detector for which the position should be determined

roi: tuple or list, optional

region of interest for the detector pixels; (default: self._area_roi/self._linear_roi)

Nav: tuple or list, optional

number of channels to average to reduce data size; (default: self._area_nav/self._linear_nav)

deg: bool, optional

flag to tell if angles are passed as degree (default: True)

Returns: ndarray

numpy array with the detector distance

getDetectorPos (*args, **kwargs)

obtains the detector position vector by applying the detector arm rotations.

Parameters: args: list

detector angles. Only detector arm angles as described by the detectorAxis attribute must be given.

kwargs: dict, optional

optional keyword arguments

dim: int, optional

dimension of the detector for which the position should be determined

roi: tuple or list, optional

region of interest for the detector pixels; (default: self._area_roi/self._linear_roi)

Nav: tuple or list, optional

number of channels to average to reduce data size; (default: self._area_nav/self._linear_nav)

deg: bool, optional

flag to tell if angles are passed as degree (default: True)

Returns: ndarray

numpy array of length 3 with vector components of the detector direction. The length of the vector is k.

init_area (detectorDir1, detectorDir2, cch1, cch2, Nch1, Nch2, distance=None, pwidth1=None, pwidth2=None, chpdeg1=None, chpdeg2=None, detrot=0, tiltazimuth=0, tilt=0, **kwargs)

initialization routine for area detectors detector direction as well as distance and pixel size or channels per degree must be given. Two separate pixel sizes and channels per degree for the two orthogonal directions can be given Parameters: detectorDir1: str

direction of the detector (along the pixel direction 1); e.g. 'z+' means higher pixel

numbers at larger z positions

direction of the detector (along the pixel direction 2); e.g. 'x+'

cch1, cch2: float

detectorDir2: str

center pixel, in direction of self.r_i at zero detectorAngles

Nch1, Nch2: int

number of detector pixels along direction 1, 2

distance: float, optional

distance of center pixel from center of rotation

pwidth1, pwidth2: float, optional

width of one pixel (same unit as distance)

chpdeg1, chpdeg2: float, optional

channels per degree (only absolute value is relevant) sign determined through detector Dir1, detector Dir2

detrot : float, optional

angle of the detector rotation around primary beam direction (used to correct misalignments)

tiltazimuth : float, optional

direction of the tilt vector in the detector plane (in degree)

tilt: float, optional

tilt of the detector plane around an axis normal to the direction given by the tiltazimuth

kwargs: dict, optional

optional keyword arguments

Nav: tuple or list, optional

number of channels to average to reduce data size (default: [1, 1])

roi: tuple or list, optional

region of interest for the detector pixels; e.g. [100, 900, 200, 800]

Note

Either distance and pwidth1, pwidth2 or chpdeg1, chpdeg2 must be given!!

Note

the channel numbers run from 0 .. NchX-1

init_linear (detectorDir, cch, Nchannel, distance=None, pixelwidth=None, chpdeg=None, tilt=0, **kwargs) initialization routine for linear detectors detector direction as well as distance and pixel size or channels per degree must be given.

Parameters: detectorDir: str

direction of the detector (along the pixel array); e.g. 'z+'

cch: float

center channel, in direction of self.r_i at zero detectorAngles

Nchannel: int

total number of detector channels

distance: float, optional

distance of center channel from center of rotation

pixelwidth: float, optional

width of one pixel (same unit as distance)

chpdeg: float, optional

channels per degree (only absolute value is relevant) sign determined through

detectorDir

tilt: float, optional

tilt of the detector axis from the detectorDir (in degree)

kwargs: dict, optional

optional keyword arguments

Nav: int, optional

number of channels to average to reduce data size (default: 1)

roi: tuple or list

region of interest for the detector pixels; e.g. [100, 900]

Note

Either distance and pixelwidth or chpdeg must be given!!

Note

the channel numbers run from 0 .. Nchannel-1

linear (*args, **kwargs)

angular to momentum space conversion for a linear detector the cch of the detector must be in direction of self.r_i when detector angles are zero

the detector geometry must be initialized by the init_linear(...) routine

Parameters: args: ndarray, list or Scalars

sample and detector angles; in total *len(self.sampleAxis)* + *len(detectorAxis)* must be given, always starting with the outer most circle. all arguments must have the same shape or length but can be mixed with Scalars (i.e. if an angle is always the same it can be given only once instead of an array)

• sAngles:

sample circle angles, number of arguments must correspond to len(self.sampleAxis)

• dAngles:

detector circle angles, number of arguments must correspond to len(self.detectorAxis)

kwargs: dict, optional

optional keyword arguments

delta: list or array-like, optional

delta angles to correct the given ones for misalignment. delta must be an numpy array or list of len(*args). used angles are then *args - delta

UB: array-like, optional

matrix for conversion from (hkl) coordinates to Q of sample used to determine not Q but (hkl) (default: self.UB)

Nav: int, optional

number of channels to average to reduce data size (default: self._linear_nav)

roi: list or tuple, optional

region of interest for the detector pixels; e.g. [100, 900] (default: self._linear_roi)

wl: float or str, optional

x-ray wavelength in angstroem (default: self._wl)

en: float, optional

x-ray energy in eV (default is converted self._wl). both wavelength and energy can also be an array which enables the QConversion for energy scans. Note that the *en* keyword overrules the *wl* keyword!

deg: bool, optional

flag to tell if angles are passed as degree (default: True)

sampledis: tuple or list or array-like

sample displacement vector in relative units of the detector distance (default: (0, 0, 0))

Returns: reciprocal space position of all detector pixels in a numpy.ndarray of

shape ((*)*(self._linear_roi[1]-self._linear_roi[0]+1) , 3)

point (*args, **kwargs)

angular to momentum space conversion for a point detector located in direction of self.r_i when detector angles are zero

Parameters: args: ndarray, list or Scalars

sample and detector angles; in total *len(self.sampleAxis)* + *len(detectorAxis)* must be given, always starting with the outer most circle. all arguments must have the same shape or length but can be mixed with Scalars (i.e. if an angle is always the same it can be given only once instead of an array)

• sAngles:

sample circle angles, number of arguments must correspond to len(self.sampleAxis)

• dAngles:

detector circle angles, number of arguments must correspond to len(self.detectorAxis)

kwargs: dict, optional

optional keyword arguments

delta: list or array-like, optional

delta angles to correct the given ones for misalignment. delta must be an numpy array or list of len(*args). used angles are then *args - delta

UB: array-like, optional

matrix for conversion from (hkl) coordinates to Q of sample used to determine not Q but (hkl) (default: self.UB)

wl: float or str, optional

x-ray wavelength in angstroem (default: self._wl)

en: float, optional

x-ray energy in eV (default is converted self._wl). both wavelength and energy can also be an array which enables the QConversion for energy scans. Note that the *en* keyword overrules the *wl* keyword!

deg: bool, optional

flag to tell if angles are passed as degree (default: True)

sampledis: tuple or list or array-like

sample displacement vector in relative units of the detector distance (default: (0, 0, 0))

Returns: ndarray

reciprocal space positions as numpy.ndarray with shape (N, 3) where N corresponds to the number of points given in the input

property sampleAxis

property handler for _sampleAxis

Returns: list

sample axes following the syntax /[xyzk][+-]/

transformSample2Lab (vector, *args)

transforms a vector from the sample coordinate frame to the laboratory coordinate system by applying the sample rotations from inner to outer circle.

Parameters: vector: sequence, list or numpy array

vector to transform

args : list

goniometer angles (sample angles or full goniometer angles can be given. If more angles than the sample circles are given they will be ignored)

Returns: ndarray

rotated vector as numpy.array

property wavelength

xrayutilities.gridder module

class xrayutilities.gridder.FuzzyGridder1D (nx)

Bases: xrayutilities.gridder.Gridder1D

An 1D binning class considering every data point to have a finite width. If necessary one data point will be split fractionally over different data bins. This is numerically more effort but represents better the typical case of a experimental data, which do not represent a mathematical point but have a finite width (e.g. X-ray data from a 1D detector).

class xrayutilities.gridder.Gridder

Bases: abc.ABC

Basis class for gridders in xrayutilities. A gridder is a function mapping irregular spaced data onto a regular grid by binning the data into equally sized elements.

There are different ways of defining the regular grid of a Gridder. In xrayutilities the data bins extend beyond the data range in the input data, but the given position being the center of these bins, extends from the minimum to the maximum of the data! The main motivation for this was to create a Gridder, which when feeded with N equidistant data points and gridded with N bins would not change the data position (not the case with numpy.histogramm functions!). Of course this leads to the fact that for homogeneous point density the first and last bin in any direction are not filled as the other bins.

A different definition is used by numpy histogram functions where the bins extend only to the end of the data range. (see numpy histogram, histogram2d, ...)

Clear ()

Clear so far gridded data to reuse this instance of the Gridder

KeepData (bool)

Normalize (bool)

set or unset the normalization flag. Normalization needs to be done to obtain proper gridding but may want to be disabled in certain cases when sequential gridding is performed

property data

return gridded data (performs normalization if switched on)

class xrayutilities.gridder.Gridder1D (nx)
Bases: xrayutilities.gridder.Gridder

dataRange (min, max, fixed=True)

define minimum and maximum data range, usually this is deduced from the given data automatically, however, for sequential gridding it is useful to set this before the first call of the gridder. data outside the range are simply ignored

Parameters: min: float

minimum value of the gridding range

max: float

maximum value of the gridding range

fixed: bool, optional

flag to turn fixed range gridding on (True (default)) or off (False)

savetxt (filename, header=")

save gridded data to a txt file with two columns. The first column is the data coordinate and the second the corresponding data value

Parameters: filename: str

output filename **header**: str, optional

optional header for the data file.

property xaxis

Returns the xaxis of the gridder the returned values correspond to the center of the data bins used by the gridding algorithm

xrayutilities.gridder.axis (min_value, max_value, n)
Compute the a grid axis.

Parameters: min_value : float

axis minimum value

max value: float

axis maximum value

n:int

number of steps

xrayutilities.gridder.delta (min_value, max_value, n)

Compute the stepsize along an axis of a grid.

Parameters: min_value : axis minimum value

max_value: axis maximum value

n: number of steps

class xrayutilities.gridder.npyGridder1D (nx)
Bases: xrayutilities.gridder.Gridder1D

property xaxis

Returns the xaxis of the gridder the returned values correspond to the center of the data bins used by the numpy.histogram function

xrayutilities.gridder.ones (*args)

Compute ones for matrix generation. The shape is determined by the number of input arguments.

xrayutilities.gridder2d module

class xrayutilities.gridder2d.FuzzyGridder2D (nx, ny)

Bases: xrayutilities.gridder2d.Gridder2D

An 2D binning class considering every data point to have a finite area. If necessary one data point will be split fractionally over different data bins. This is numerically more effort but represents better the typical case of a experimental data, which do not represent a mathematical point but have a finite size (e.g. X-ray data from a 2D detector or reciprocal space maps measured with point/linear detector).

Currently only a rectangular area can be considered during the gridding.

class xrayutilities.gridder2d.Gridder2D (nx, ny)

Bases: xrayutilities.gridder.Gridder

SetResolution (nx, ny)

Reset the resolution of the gridder. In this case the original data stored in the object will be deleted.

Parameters: nx: int

number of points in x-direction

ny: int

number of points in y-direction

dataRange (xmin, xmax, ymin, ymax, fixed=True)

define minimum and maximum data range, usually this is deduced from the given data automatically, however, for sequential gridding it is useful to set this before the first call of the gridder. data outside the range are simply ignored

Parameters: xmin, ymin : float

minimum value of the gridding range in x, y

xmax, ymax : float

maximum value of the gridding range in x, y

fixed: bool, optional

flag to turn fixed range gridding on (True (default)) or off (False)

savetxt (filename, header=")

save gridded data to a txt file with two columns. The first two columns are the data coordinates and the last one the corresponding data value.

Parameters: filename: str

output filename **header**: str. optional

optional header for the data file.

property xaxis

property xmatrix

property yaxis

property ymatrix

class xrayutilities.gridder2d.Gridder2DList (nx, ny)

Bases: xrayutilities.gridder2d.Gridder2D

special version of a 2D gridder which performs no actual averaging of the data in one grid/bin but just collects the data-objects belonging to one bin for further treatment by the user

Clear ()

Clear so far gridded data to reuse this instance of the Gridder

property data

return gridded data, in this special version no normalization is defined!

xrayutilities.gridder3d module

```
class xrayutilities.gridder3d.FuzzyGridder3D (nx, ny, nz)
```

Bases: xrayutilities.gridder3d.Gridder3D

An 3D binning class considering every data point to have a finite volume. If necessary one data point will be split fractionally over different data bins. This is numerically more effort but represents better the typical case of a experimental data, which do not represent a mathematical point but have a finite size.

Currently only a quader can be considered as volume during the gridding.

```
class xrayutilities.gridder3d.Gridder3D (nx, ny, nz)
```

Bases: xrayutilities.gridder.Gridder

SetResolution (nx, ny, nz)

dataRange (xmin, xmax, ymin, ymax, zmin, zmax, fixed=True)

define minimum and maximum data range, usually this is deduced from the given data automatically, however, for sequential gridding it is useful to set this before the first call of the gridder. data outside the range are simply ignored

xrayutilities.mpl_helper module

Defines new matplotlib Sqrt scale which further allows for negative values by using the sign of the original value as sign of the plotted value.

```
class xrayutilities.mpl_helper.SqrtAllowNegScale (axis, **kwargs)
 Bases: matplotlib.scale.ScaleBase
 Scales data using a sqrt-function, however, allowing also negative values.
 The scale function:
     sign(y) * sqrt(abs(y))
 The inverse scale function:
     sign(y) * y**2
 class InvertedSqrtTransform (shorthand_name=None)
   Bases: matplotlib.transforms.Transform
   has_inverse = True
   input_dims = 1
   inverted()
     Return the corresponding inverse transformation.
     It holds x == self.inverted().transform(self.transform(x)).
     The return value of this method should be treated as temporary. An update to self does not cause a
     corresponding update to its inverted copy.
   is_separable = True
   output dims = 1
   transform_non_affine (a)
     Apply only the non-affine part of this transformation.
     transform(values)
                                                                              equivalent
                                                                                                      to
     transform_affine(transform_non_affine(values)).
     In non-affine transformations, this is generally equivalent to transform(values). In affine transformations,
     this is always a no-op.
```

```
Parameters: values : array
```

The input values as NumPy array of length input_dims or shape (N x

input_dims).

Returns: array

The output values as NumPy array of length input_dims or shape (N x output_dims), depending on the input.

```
class SqrtTransform (shorthand_name=None)
```

Bases: matplotlib.transforms.Transform

```
has_inverse = True
input_dims = 1
```

inverted()

return the inverse transform for this transform.

```
is_separable = True
output_dims = 1
```

transform_non_affine (a)

This transform takes an Nx1 numpy array and returns a transformed copy.

get_transform()

Return the **Transform** object associated with this scale.

limit_range_for_scale (vmin, vmax, minpos)

Override to limit the bounds of the axis to the domain of the transform. In the case of Mercator, the bounds should be limited to the threshold that was passed in. Unlike the autoscaling provided by the tick locators, this range limiting will always be adhered to, whether the axis range is set manually, determined automatically or changed through panning and zooming.

```
name = 'sqrt'
```

set_default_locators_and_formatters (axis)

Set the locators and formatters of axis to instances suitable for this scale.

```
class xrayutilities.mpl_helper.SqrtTickLocator (nbins=7, symmetric=True)
Bases: matplotlib.ticker.Locator
```

set_params (nbins, symmetric)

Set parameters within this locator.

tick_values (vmin, vmax)

Return the values of the located ticks given vmin and vmax.

Note

To get tick locations with the vmin and vmax values defined automatically for the associated axis simply call the Locator instance:

```
>>> print(type(loc))
<type 'Locator'>
>>> print(loc())
[1, 2, 3, 4]
```

view_limits (dmin, dmax)

Set the view limits to the nearest multiples of base that contain the data

xrayutilities.normalize module

module to provide functions that perform block averaging of intensity arrays to reduce the amount of data (mainly for PSD and CCD measurements

and

provide functions for normalizing intensities for

- count time
- absorber (user-defined function)
- monitor
- flatfield correction

class xrayutilities.normalize.IntensityNormalizer (det=", **keyargs)

Bases: object

generic class for correction of intensity (point detector, or MCA, single CCD frames) for count time and absorber factors the class must be supplied with a absorber correction function and works with data structures provided by xrayutilities.io classes or the corresponding objects from hdf5 files

property absfun

absfun property handler returns the costum correction function or None

property avmon

av_mon property handler

returns the value of the average monitor or None if average is calculated from the monitor field

property darkfield

flatfield property handler

returns the current set darkfield of the detector or None if not set

property det

det property handler

returns the detector field name

property flatfield

flatfield property handler

returns the current set flatfield of the detector or None if not set

property mon

mon property handler

returns the monitor field name or None if not set

property time

time property handler

returns the count time or the field name of the count time or None if time is not set

xrayutilities.normalize.blockAverage1D (data, Nav)

perform block average for 1D array/list of Scalar values all data are used. at the end of the array a smaller cell may be used by the averaging algorithm

Parameters: data: array-like

data which should be contracted (length N)

Nav: int

number of values which should be averaged

Returns: ndarray

block averaged numpy array of data type numpy.double (length ceil(N/Nav))

xrayutilities.normalize.blockAverage2D (data2d, Nav1, Nav2, **kwargs)

perform a block average for 2D array of Scalar values all data are used therefore the margin cells may differ in size

Parameters: data2d: ndarray

array of 2D data shape (N, M)

Nav1, Nav2: int

a field of (Nav1 x Nav2) values is contracted

kwargs: dict, optional

optional keyword argument

roi: tuple or list, optional

region of interest for the 2D array. e.g. [20, 980, 40, 960], reduces M, and M!

Returns: ndarray

block averaged numpy array with type numpy.double with shape (ceil(N/Nav1),

ceil(M/Nav2))

xrayutilities.normalize.blockAverageCCD (data3d, Nav1, Nav2, **kwargs)

perform a block average for 2D frames inside a 3D array, all data are used therefore the margin cells may differ in size

Parameters: data3d: ndarray

array of 3D data shape (Nframes, N, M)

Nav1. Nav2: int

a field of (Nav1 x Nav2) values is contracted

kwargs: dict, optional

optional keyword argument

roi: tuple or list, optional

region of interest for the 2D array. e.g. [20, 980, 40, 960], reduces M, and M!

Returns: ndarray

block averaged numpy array with type numpy.double with shape (Nframes,

ceil(N/Nav1), ceil(M/Nav2))

xrayutilities.normalize.blockAveragePSD (psddata, Nav, **kwargs)

perform a block average for serveral PSD spectra all data are used therefore the last cell used for averaging may differ in size

Parameters: psddata: ndarray

array of 2D data shape (Nspectra, Nchannels)

Nav: int

number of channels which should be averaged

kwargs: dict, optional

optional keyword argument

roi: tuple or list

region of interest for the 2D array. e.g. [20, 980] Nchannels = 980-20

Returns: ndarray

block averaged psd spectra as numpy array with type numpy double of shape (Nspectra

, ceil(Nchannels/Nav))

xrayutilities.q2ang_fit module

Module provides functions to convert a q-vector from reciprocal space to angular space. a simple implementation uses scipy optimize routines to perform a fit for a arbitrary goniometer.

The user is, however, expected to use the bounds variable to put restrictions to the number of free angles to obtain reproducible results. In general only 3 angles are needed to fit an arbitrary q-vector (2 sample + 1 detector angles or 1 sample + 2 detector). More complicated restrictions can be implemented using the Imfit package. (done upon request!)

The function is based on a fitting routine. For a specific goniometer also analytic expressions from literature can be used as they are implemented in the predefined experimental classes HXRD, NonCOP, and GID.

xrayutilities.q2ang_fit.Q2AngFit (qvec, expclass, bounds=None, ormat=array([[1.0, 0.0, 0.0], [0.0, 1.0, 0.0], [0.0, 0.0, 1.0]]), startvalues=None, constraints=[])

Functions to convert a q-vector from reciprocal space to angular space. This implementation uses scipy optimize routines to perform a fit for a goniometer with arbitrary number of goniometer angles.

The user *must* use the bounds variable to put restrictions to the number of free angles to obtain reproducible results. In general only 3 angles are needed to fit an arbitrary q-vector (2 sample + 1 detector angles or 1 sample + 2 detector).

Parameters: qvec: tuple or list or array-like

q-vector for which the angular positions should be calculated

expclass: Experiment

experimental class used to define the goniometer for which the angles should be calculated.

bounds: tuple or list

bounds of the goniometer angles. The number of bounds must correspond to the number of goniometer angles in the expclass. Angles can also be fixed by supplying only one value for a particular angle. e.g.: ((low, up), fix, (low2, up2), (low3, up3))

ormat: array-like

orientation matrix of the sample to be used in the conversion

startvalues : array-like

start values for the fit, which can significantly speed up the conversion. The number of values must correspond to the number of angles in the goniometer of the expclass

constraints: list

sequence of constraint dictionaries. This allows applying arbitrary (e.g. pseudo-angle) contraints by supplying according constraint functions. An entry of the constraints argument must be a dictionary with at least the 'type' and 'fun' set. 'type' can be either 'eq' or 'ineq' for equality or inequality constraints. 'fun' must be a callable function which for 'eq'-constraints returns 0 when the equality condition is fulfilled (see constraints documentation in scipy.optimize.minimize for details). The supplied function will be called with the arguments gonimeter angle list as argument. Typically this means you will have to use a lambda function.

Returns: fittedangles: list

list of fitted goniometer angles

qerror: float

error in reciprocal space

errcode: int

error-code of the scipy minimize function. for a successful fit the error code should be <=2

xrayutilities.q2ang_fit.exitAngleConst (angles, alphaf, xrd)

helper function for an pseudo-angle constraint of the exit angle. Can be used together with the Q2AngFit-routine in the 'constraints' argument. An example use case scenario to fix the exit angle to 1 degree would be: constraints={'type': 'eq', 'fun': lambda a: exitAngleConst(a, 1, xrd)}

Parameters: angles: iterable

fit parameters of Q2AngFit

alphaf: float

the exit angle which should be fixed

xrd: Experiment

the Experiment object to use for qconversion

```
xrayutilities.q2ang_fit.incidenceAngleConst (angles, alphai, xrd)
```

helper function for an pseudo-angle constraint of the incidence angle. Can be used together with the Q2AngFit-routine in the 'constraints' argument. An example use case scenario to fix the incidence angle to 1 degree would be: constraints={'type': 'eq', 'fun': lambda a: incidenceAngleConst(a, 1, xrd)}

Parameters: angles: iterable

fit parameters of Q2AngFit

alphai: float

the incidence angle which should be fixed

xrd: Experiment

the Experiment object to use for gconversion

xrayutilities.utilities module

xrayutilities utilities contains a conglomeration of useful functions which do not fit into one of the other files

xrayutilities.utilities.frac2str (f, denominator_limit=25, fmt='%7.4f') convert a float to a string attempting to represent it as a fraction

Parameters: f: float

floating point number to be represented as string

denominator limit: int

maximal integer used as denominator. If f can't be expressed (within xu.config.EPSILON) by a fraction with a denominator up to this number a floating point string will be returned

fmt: str

format string used in case a floating point representation is needed

Returns: str

xrayutilities.utilities.import_lmfit (funcname='XU')

lazy import function for Imfit

xrayutilities.utilities.import matplotlib pyplot (funcname='XU')

lazy import function of matplotlib.pyplot

Parameters: funcname: str

identification string of the calling function

Returns: flag: bool

the flag is True if the loading was successful and False otherwise.

pyplot

On success pyplot is the matplotlib.pyplot package.

xrayutilities.utilities.import_mayavi_mlab (funcname='XU')

lazy import function of mayavi.mlab

Parameters: funcname: str

identification string of the calling function

Returns: flag: bool

the flag is True if the loading was successful and False otherwise.

mlab

On success mlab is the mayavi.mlab package.

xrayutilities.utilities.maplog (inte, dynlow='config', dynhigh='config')

clips values smaller and larger as the given bounds and returns the log10 of the input array. The bounds are given as exponent with base 10 with respect to the maximum in the input array. The function is implemented in analogy to J. Stangl's matlab implementation.

Parameters: inte: ndarray

numpy.array, values to be cut in range

dynlow: float, optional

10^(-dynlow) will be the minimum cut off

dynhigh: float, optional

10^(-dynhigh) will be the maximum cut off

Returns: ndarray

numpy.array of the same shape as inte, where values smaller/larger than 10^(-dynlow,

dynhigh) were replaced by 10^(-dynlow, dynhigh)

Examples

>>> lint = maplog(int, 5, 2)

xrayutilities.utilities_noconf module

xrayutilities utilities contains a conglomeration of useful functions this part of utilities does not need the config class

class xrayutilities.utilities_noconf.ABC

Bases: object

Helper class that provides a standard way to create an ABC using inheritance.

xrayutilities.utilities_noconf.check_kwargs (kwargs, valid_kwargs, identifier)

Raises an TypeError if kwargs included a key which is not in valid_kwargs.

Parameters: kwargs: dict

keyword arguments dictionary

valid_kwargs: dict

dictionary with valid keyword arguments and their description

identifier: str

string to identifier the caller of this function

xrayutilities.utilities_noconf.clear_bit (f, Offset)

clears the bet at an offset

xrayutilities.utilities_noconf.en2lam (inp)

converts the input energy in eV to a wavelength in Angstrom

Parameters: inp : float or str

energy in eV

Returns: float

wavlength in Angstrom

Examples

>>> wavelength = en2lam(8048)

xrayutilities.utilities_noconf.energy (en)

convert common energy names to energies in eV

so far this works with CuKa1, CuKa2, CuKa12, CuKb, MoKa1

Parameters: en: float, array-like or str

energy either as scalar or array with value in eV, which will be returned unchanged; or

string with name of emission line

Returns: float or array-like

energy in eV

xrayutilities.utilities_noconf.exchange_filepath (orig, new, keep=0, replace=None)

function to exchange the root of a filename with the option of keeping the inner directory structure. This for example includes such a conversion /dir_a/subdir/sample/file.txt -> /home/user/data/sample/file.txt where the

innermost directory name is kept (keep=1), or equally the three outer most are replaced (replace=3). One can either give keep, or replace, with replace taking preference if both are given. Note that replace=1 on Linux/Unix replaces only the root for absolute paths.

Parameters: orig: str

original filename which should have its data root replaced

new: str

new path which should be used instead

keep: int, optional

number of inner most directory names which should be kept the same in the output (default = 0)

replace: int, optional

number of outer most directory names which should be replaced in the output (default = None)

Returns: str

filename string

Examples

```
>>> exchange_filepath('/dir_a/subdir/sam/file.txt', '/data', 1)
'/data/sam/file.txt'
```

xrayutilities.utilities_noconf.exchange_path (orig, new, keep=0, replace=None)

function to exchange the root of a path with the option of keeping the inner directory structure. This for example includes such a conversion /dir_a/subdir/images/sample -> /home/user/data/images/sample where the two innermost directory names are kept (keep=2), or equally the three outer most are replaced (replace=3). One can either give keep, or replace, with replace taking preference if both are given. Note that replace=1 on Linux/Unix replaces only the root for absolute paths.

Parameters: orig: str

original path which should be replaced by the new path

new: str

new path which should be used instead

keep: int, optional

number of inner most directory names which should be kept the same in the output (default = 0)

replace: int, optional

number of outer most directory names which should be replaced in the output (default = None)

Returns: str

directory path string

Examples

```
>>> exchange_path('/dir_a/subdir/img/sam', '/home/user/data', keep=2)
'/home/user/data/img/sam'
```

```
xrayutilities.utilities_noconf.is_valid_variable_name (name)
```

xrayutilities.utilities_noconf.lam2en (inp)

converts the input wavelength in Angstrom to an energy in eV

Parameters: inp : float or str

wavelength in Angstrom

Returns: float

energy in eV

Examples

```
>>> energy = lam2en(1.5406)
```

xrayutilities_utilities_noconf.makeNaturalName (name, check=False)

xrayutilities.utilities_noconf.set_bit (f, offset)

sets the bit at an offset

xrayutilities.utilities_noconf.wavelength(wl)

convert common energy names to energies in eV

so far this works with CuKa1, CuKa2, CuKa12, CuKb, MoKa1

Parameters: wl : float, array-like or str

wavelength; If scalar or array the wavelength in Angstrom will be returned unchanged,

string with emission name is converted to wavelength

Returns: float or array-like

wavelength in Angstrom

Module contents

xrayutilities is a Python package for assisting with x-ray diffraction experiments. Its the python package included in xrayutilities.

It helps with planning experiments as well as analyzing the data.

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xrayutilities.analysis package

Submodules

xrayutilities.analysis.line_cuts module

xrayutilities.analysis.line_cuts.get_arbitrary_line (qpos, intensity, point, vec, npoints, intrange) extracts a line scan from reciprocal space map data along an arbitrary line defined by the point 'point' and propergation vector 'vec'. Integration of the data is performed in a cylindrical volume along the line. This function works for 2D and 3D input data!

Parameters: qpos: list of array-like objects

arrays of x, y (list with two components) or x, y, z (list with three components)

momentum transfers

intensity: array-like

2D or 3D array of reciprocal space intensity with shape equal to the qpos entries

point: tuple, list or array-like

point on the extraction line (2 or 3 coordinates)

vec: tuple, list or array-like

propergation vector of the extraction line (2 or 3 coordinates)

npoints: int

number of points in the output data

intrange: float

radius of the cylindrical integration volume around the extraction line

Returns: qpos, qint: ndarray

line scan coordinates and intensities

used_mask : ndarray

mask of used data, shape is the same as the input intensity: True for points which

contributed, False for all others

Examples

```
>>> qcut, qint, mask = get_arbitrary_line([qx, qy, qz], inten, (1.1, 2.2, 0.0), (1, 1, 1), 200, 0.1)
```

xrayutilities.analysis.line_cuts.get_omega_scan (qpos, intensity, cutpos, npoints, intrange, **kwargs) extracts an omega scan from reciprocal space map data with integration along either the 2theta, or radial (omega-2theta) direction. The coplanar diffraction geometry with qy and qz as diffraction plane is assumed. This is consistent with the coplanar geometry implemented in the HXRD-experiment class.

This function works for 2D and 3D input data in the same way!

Parameters: qpos: list of array-like objects

arrays of y, z (list with two components) or x, y, z (list with three components)

momentum transfers

intensity: array-like

2D or 3D array of reciprocal space intensity with shape equal to the qpos entries

cutpos: tuple or list

y/z-position or x/y/z-position at which the line scan should be extracted. this must be

have two entries for 2D data (z-position) and a three for 3D data

npoints: int

number of points in the output data

intrange : float

integration range in along intdir in degree. data will be integrated from -intrange ..

+intrange

intdir: {'2theta', 'radial'}, optional

integration direction: '2theta': scattering angle (default), or 'radial': omega-2theta

direction.

wl: float or str, optional

wavelength used to determine angular integration positions

Note:

Although applicable for any set of data, the extraction only makes sense when the data

are aligned into the y/z-plane.

Returns: om, omint : ndarray

omega scan coordinates and intensities

used_mask : ndarray

mask of used data, shape is the same as the input intensity: True for points which contributed, False for all others

Examples

```
>>> omcut, omcut_int, mask = get_omega_scan([qy, qz], inten, [2.0, 5.0], 250, intrange=0.1)
```

xrayutilities.analysis.line_cuts.get_qx_scan (qpos, intensity, cutpos, npoints, intrange, **kwargs) extracts a qx scan from 3D reciprocal space map data with integration along either, the perpendicular plane in q-space, omega (sample rocking angle) or 2theta direction. For the integration in angular space (omega, or 2theta) the coplanar diffraction geometry with qy and qz as diffraction plane is assumed. This is consistent with the coplanar geometry implemented in the HXRD-experiment class.

arrays of x, y, z (list with three components) momentum transfers

intensity: array-like

3D array of reciprocal space intensity with shape equal to the qpos entries

cutpos: tuple/list

y/z-position at which the line scan should be extracted. this must be and a tuple/list with the qy, qz cut position

npoints: int

number of points in the output data

intrange: float

integration range in along *intdir*, either in 1/AA (*q*) or degree ('omega', or '2theta'). data will be integrated from *-intrange* .. +*intrange*

intdir: {'q', 'omega', '2theta'}, optional

integration direction: 'q': perpendicular Q-plane (default), 'omega': sample rocking angle, or '2theta': scattering angle.

wl: float or str, optional

wavelength used to determine angular integration positions

Note:

The angular integration directions although applicable for any set of data only makes sense when the data are aligned into the y/z-plane.

Returns: qx, qxint: ndarray

qx scan coordinates and intensities

used_mask : ndarray

mask of used data, shape is the same as the input intensity: True for points which contributed, False for all others

Examples

```
>>> qxcut, qxcut_int, mask = get_qx_scan([qx, qy, qz], inten, [0, 2.0], 250, intrange=0.01)
```

xrayutilities.analysis.line_cuts.get_qy_scan (qpos, intensity, cutpos, npoints, intrange, **kwargs) extracts a qy scan from reciprocal space map data with integration along either, the perpendicular plane in q-space, omega (sample rocking angle) or 2theta direction. For the integration in angular space (omega, or 2theta) the coplanar diffraction geometry with qy and qz as diffraction plane is assumed. This is consistent with the coplanar geometry implemented in the HXRD-experiment class.

arrays of y, z (list with two components) or x, y, z (list with three components)

momentum transfers

intensity: array-like

2D or 3D array of reciprocal space intensity with shape equal to the gpos entries

cutpos: float or tuple/list

x/z-position at which the line scan should be extracted. this must be a float for 2D data

(z-position) and a tuple with two values for 3D data

npoints: int

number of points in the output data

intrange: float

integration range in along *intdir*, either in 1/AA (*q*) or degree ('omega', or '2theta'). data will be integrated from *-intrange* .. +*intrange*

intdir: {'q', 'omega', '2theta'}, optional

integration direction: 'q': perpendicular Q-plane (default), 'omega': sample rocking angle, or '2theta': scattering angle.

wl: float or str, optional

wavelength used to determine angular integration positions

Note:

For 3D data the angular integration directions although applicable for any set of data only makes sense when the data are aligned into the y/z-plane.

Returns: qy, qyint : ndarray

qy scan coordinates and intensities

used_mask : ndarray

mask of used data, shape is the same as the input intensity: True for points which contributed, False for all others

Examples

xrayutilities.analysis.line_cuts.get_qz_scan (qpos, intensity, cutpos, npoints, intrange, **kwargs) extracts a qz scan from reciprocal space map data with integration along either, the perpendicular plane in q-space, omega (sample rocking angle) or 2theta direction. For the integration in angular space (omega, or 2theta) the coplanar diffraction geometry with qy and qz as diffraction plane is assumed. This is consistent with the coplanar geometry implemented in the HXRD-experiment class.

arrays of y, z (list with two components) or x, y, z (list with three components)

momentum transfers

intensity: array-like

2D or 3D array of reciprocal space intensity with shape equal to the qpos entries

cutpos: float or tuple/list

x/y-position at which the line scan should be extracted. this must be a float for 2D data

and a tuple with two values for 3D data

npoints: int

number of points in the output data

intrange: float

integration range in along *intdir*, either in 1/AA (*q*) or degree ('omega', or '2theta'). data will be integrated from *-intrange/2* .. +*intrange/2*

intdir: {'q', 'omega', '2theta'}, optional

integration direction: 'q': perpendicular Q-plane (default), 'omega': sample rocking angle, or '2theta': scattering angle.

wl: float or str, optional

wavelength used to determine angular integration positions

Note:

For 3D data the angular integration directions although applicable for any set of data only makes sense when the data are aligned into the y/z-plane.

Returns: qz, qzint : ndarray

qz scan coordinates and intensities

used_mask : ndarray

mask of used data, shape is the same as the input intensity: True for points which contributed, False for all others

Examples

```
>>> qzcut, qzcut_int, mask = get_qz_scan([qy, qz], inten, 3.0, 200, intrange=0.3)
```

xrayutilities.analysis.line_cuts.get_radial_scan (qpos, intensity, cutpos, npoints, intrange,
**kwarqs)

extracts a radial scan from reciprocal space map data with integration along either the omega or 2theta direction. The coplanar diffraction geometry with qy and qz as diffraction plane is assumed. This is consistent with the coplanar geometry implemented in the HXRD-experiment class.

arrays of y, z (list with two components) or x, y, z (list with three components)

momentum transfers

intensity: array-like

2D or 3D array of reciprocal space intensity with shape equal to the qpos entries

cutpos: tuple or list

y/z-position or x/y/z-position at which the line scan should be extracted. this must be

have two entries for 2D data (z-position) and a three for 3D data

npoints: int

number of points in the output data

intrange: float

integration range in along *intdir* in degree. data will be integrated from *-intrange* .. +*intrange*

intdir: {'omega', '2theta'}, optional

integration direction: 'omega': sample rocking angle (default), '2theta': scattering angle

wl: float or str, optional

wavelength used to determine angular integration positions

Note:

Although applicable for any set of data, the extraction only makes sense when the data are aligned into the y/z-plane.

Returns: tt, omttint : ndarray

omega-2theta scan coordinates (2theta values) and intensities

used_mask : ndarray

mask of used data, shape is the same as the input intensity: True for points which contributed, False for all others

Examples

```
>>> ttcut, omtt_int, mask = get_radial_scan([qy, qz], inten, [2.0, 5.0], 250, intrange=0.1)
```

xrayutilities.analysis.line_cuts.get_ttheta_scan (qpos, intensity, cutpos, npoints, intrange,

**kwargs)

extracts a 2theta scan from reciprocal space map data with integration along either the omega or radial direction. The coplanar diffraction geometry with qy and qz as diffraction plane is assumed. This is consistent with the coplanar geometry implemented in the HXRD-experiment class.

arrays of y, z (list with two components) or x, y, z (list with three components)

momentum transfers

intensity: array-like

2D or 3D array of reciprocal space intensity with shape equal to the qpos entries

cutpos: tuple or list

y/z-position or x/y/z-position at which the line scan should be extracted. this must be

have two entries for 2D data (z-position) and a three for 3D data

npoints: int

number of points in the output data

intrange: float

integration range in along *intdir* in degree. data will be integrated from *-intrange* .. +*intrange*

intdir: {'omega', 'radial'}, optional

integration direction: 'omega': sample rocking angle (default), 'radial': omega-2theta

direction

wl: float or str, optional

wavelength used to determine angular integration positions

Note:

Although applicable for any set of data, the extraction only makes sense when the data are aligned into the y/z-plane.

Returns: tt, ttint: ndarray

2theta scan coordinates and intensities

used_mask : ndarray

mask of used data, shape is the same as the input intensity: True for points which contributed, False for all others

Examples

```
>>> ttcut, tt_int, mask = get_ttheta_scan([qy, qz], inten, [2.0, 5.0], 250, intrange=0.1)
```

xrayutilities.analysis.misc module

miscellaneous functions helpful in the analysis and experiment

xrayutilities.analysis.misc.coplanar_intensity (mat, exp, hkl, thickness, thMono, sample_width=10, beam_width=1)

Calculates the expected intensity of a Bragg peak from an epitaxial thin film measured in coplanar geometry (integration over omega and 2theta in angular space!)

Parameters: mat: Crystal

Crystal instance for structure factor calculation

exp: Experiment

Experimental(HXRD) class for the angle calculation

hkl: list, tuple or array-like

Miller indices of the peak to calculate

thickness: float

film thickness in nm

thMono: float

Bragg angle of the monochromator (deg)

sample_width: float, optional

width of the sample along the beam

beam_width : float, optional

width of the beam in the same units as the sample size

Returns: float

intensity of the peak

xrayutilities.analysis.misc.getangles (peak, sur, inp) calculates the chi and phi angles for a given peak

Parameters: peak: list or array-like

hkl for the peak of interest

sur : list or array-like hkl of the surface inp : list or array-like

inplane reference peak or direction

Returns: list

[chi, phi] for the given peak on surface sur with inplane direction inp as reference

Examples

To get the angles for the -224 peak on a 111 surface type

```
>>> [chi, phi] = getangles([-2, 2, 4], [1, 1, 1], [2, 2, 4])
```

xrayutilities.analysis.misc.getunitvector (chi, phi, ndir=(0, 0, 1), idir=(1, 0, 0))

return unit vector determined by spherical angles and definition of the polar axis and inplane reference direction (phi=0)

Parameters: chi, phi : float

spherical angles (polar and azimuthal) in degree

ndir: tuple, list or array-like

polar/z-axis (determines chi=0)

idir: tuple, list or array-like

azimuthal axis (determines phi=0)

xrayutilities.analysis.sample_align module

functions to help with experimental alignment during experiments, especially for experiments with linear and area detectors

xrayutilities.analysis.sample_align.area_detector_calib (angle1, angle2, ccdimages, detaxis, r_i, plot=True, cut_off=0.7, start=(None, None, 1, 0, 0, 0, 0), fix=(False, False, True, False, Fa

function to calibrate the detector parameters of an area detector it determines the detector tilt possible rotations and offsets in the detector arm angles

Parameters: angle1: array-like

outer detector arm angle

angle2: array-like

inner detector arm angle

ccdimages: array-like

images of the ccd taken at the angles given above

detaxis: list of str

detector arm rotation axis; default: ['z+', 'y-']

r_i : str

primary beam direction [xyz][+-]; default 'x+'

plot: bool, optional

flag to determine if results and intermediate results should be plotted; default: True

cut_off : float, optional

cut off intensity to decide if image is used for the determination or not; default: 0.7 = 70%

start: tuple, optional

sequence of start values of the fit for parameters, which can not be estimated automatically or might want to be fixed. These are: pwidth1, pwidth2, distance, tiltazimuth, tilt, detector_rotation, outerangle_offset. By default (None, None, 1, 0, 0, 0, 0) is used.

fix: tuple of bool

fix parameters of start (default: (False, False, True, False, False, False, False)) It is strongly recommended to either fix the distance or the pwidth1, 2 values.

fig: Figure, optional

matplotlib figure used for plotting the error default: None (creates own figure)

wl: float or str

wavelength of the experiment in Angstrom (default: config.WAVELENGTH) value does not really matter here but does affect the scaling of the error

plotlog: bool

flag to specify if the created error plot should be on log-scale

nwindow: int

window size for determination of the center of mass position after the center of mass of every full image is determined, the center of mass is determined again using a window of size nwindow in order to reduce the effect of hot pixels.

debug: bool

flag to specify that you want to see verbose output and saving of images to show if the CEN determination works

xrayutilities.analysis.sample_align.area_detector_calib_hkl (sampleang, angle1, angle2, ccdimages, hkls, experiment, material, detaxis, r_i, plot=True, cut_off=0.7, start=(None, None, 1, 0, 0, 0, 0, 0, 0, 0); config'), fix=(False, False, True, False, False, False, False, False, False, False), fig=None, plotlog=False, nwindow=50, debug=False)

function to calibrate the detector parameters of an area detector it determines the detector tilt possible rotations and offsets in the detector arm angles

in this variant not only scans through the primary beam but also scans at a set of symmetric reflections can be used for the detector parameter determination. for this not only the detector parameters but in addition the sample orientation and wavelength need to be fit. Both images from the primary beam hkl = (0, 0, 0) and from a symmetric reflection hkl = (h, k, l) need to be given for a successful run.

Parameters: sampleang: array-like

sample rocking angle (needed to align the reflections (same rotation direction as inner detector rotation)) other sample angle are not allowed to be changed during the scans

angle1: array-like

outer detector arm angle

angle2: array-like

inner detector arm angle

ccdimages : array-like

images of the ccd taken at the angles given above

hkls: list or array-like

hkl values for every image

experiment : Experiment

Experiment class object needed to get the UB matrix for the hkl peak treatment

material: Crystal

material used as reference crystal

detaxis: list of str

detector arm rotation axis; default: ['z+', 'y-']

r_i : str

primary beam direction [xyz][+-]; default 'x+'

plot: bool, optional

flag to determine if results and intermediate results should be plotted; default: True

cut off: float, optional

cut off intensity to decide if image is used for the determination or not; default: 0.7 = 70%

start: tuple, optional

sequence of start values of the fit for parameters, which can not be estimated automatically or might want to be fixed. These are: pwidth1, pwidth2, distance, tiltazimuth, tilt, detector_rotation, outerangle_offset, sampletilt, sampletiltazimuth, wavelength. By default (None, None, 1, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0).

fix: tuple of bool

fix parameters of start (default: (False, False, True, False, False, False, False, False, False, False, False, False) It is strongly recommended to either fix the distance or the pwidth1, 2 values.

fig: Figure, optional

matplotlib figure used for plotting the error default: None (creates own figure)

plotlog: bool

flag to specify if the created error plot should be on log-scale

nwindow: int

window size for determination of the center of mass position after the center of mass of every full image is determined, the center of mass is determined again using a window of size nwindow in order to reduce the effect of hot pixels.

debug: bool

flag to specify that you want to see verbose output and saving of images to show if the CEN determination works

xrayutilities.analysis.sample_align.fit_bragg_peak (om, tt, psd, omalign, ttalign, exphxrd, frange=(0.03, 0.03), peaktype='Gauss', plot=True)

helper function to determine the Bragg peak position in a reciprocal space map used to obtain the position needed for correction of the data. the determination is done by fitting a two dimensional Gaussian (xrayutilities.math.Gauss2d) or Lorentzian (xrayutilities.math.Lorentz2d)

PLEASE ALWAYS CHECK THE RESULT CAREFULLY!

Parameters: om, tt: array-like

angular coordinates of the measurement either with size of psd or of psd.shape[0]

psd: array-like

intensity values needed for fitting

omalign: float

aligned omega value, used as first guess in the fit

ttalign: float

aligned two theta values used as first guess in the fit these values are also used to set the range for the fit: the peak should be within +/-frangeAA^{-1} of those values

exphxrd: Experiment

experiment class used for the conversion between angular and reciprocal space.

frange: tuple of float, optional

data range used for the fit in both directions (see above for details default:(0.03, 0.03) unit: AA^{-1})

peaktype : {'Gauss', 'Lorentz'}

peak type to fit plot: bool, optional

if True (default) function will plot the result of the fit in comparison with the measurement.

Returns: omfit, ttfit: float

fitted angular values

params: list

fit parameters (of the Gaussian/Lorentzian)

covariance : ndarray

covariance matrix of the fit parameters

xrayutilities.analysis.sample_align.linear_detector_calib (angle, mca_spectra, **keyargs) function to calibrate the detector distance/channel per degrees for a straight linear detector mounted on a detector arm

Parameters: angle: array-like

array of angles in degree of measured detector spectra

mca_spectra: array-like

corresponding detector spectra (shape: (len(angle), Nchannels)

r_i: str, optional

primary beam direction as vector [xyz][+-]; default: 'y+'

detaxis: str, optional

detector arm rotation axis [xyz][+-]; default: 'x+'

Returns: pixelwidth: float

width of the pixel at one meter distance, pixelwidth is negative in case the hit channel number decreases upon an increase of the detector angle

center_channel: float

central channel of the detector

detector_tilt: float, optional

if usetilt=True the fitted tilt of the detector is also returned

Note

L/pixelwidth*pi/180 ~= channel/degree, with the sample detector distance L

The function also prints out how a linear detector can be initialized using

the results obtained from this calibration. Carefully check the results

Other plot: bool

Parameters: flag to specify if a visualization of the fit should be done

usetilt: bool

whether to use model considering a detector tilt, i.e. deviation angle of the pixel direction from orthogonal to the primary beam (default: True)

Seealso

psd_chdeg

low level function with more configurable options

xrayutilities.analysis.sample_align.miscut_calc (phi, aomega, zeros=None, omega0=None,
plot=True)

function to calculate the miscut direction and miscut angle of a sample by fitting a sinusoidal function to the variation of the aligned omega values of more than two reflections. The function can also be used to fit reflectivity alignment values in various azimuths.

Parameters: phi : list, tuple or array-like

azimuths in which the reflection was aligned (deg)

aomega : list, tuple or array-like aligned omega values (deg)

zeros: list, tuple or array-like, optional

angles at which surface is parallel to the beam (deg). For the analysis the angles (aomega - zeros) are used.

omega0: float, optional

if specified the nominal value of the reflection is not included as fit parameter, but is fixed to the specified value. This value is MANDATORY if ONLY TWO AZIMUTHS are given.

plot: bool, optional

flag to specify if a visualization of the fit is wanted. default: True

Returns: omega0 : float

the omega value of the reflection should be close to the nominal one

phi0: float

the azimuth in which the primary beam looks upstairs

miscut: float

amplitude of the sinusoidal variation == miscut angle

xrayutilities.analysis.sample_align.**psd_chdeg** (angles, channels, stdev=None, usetilt=True, plot=True, datap='xk', modelline='--r', modeltilt='-b', fignum=None, mlabel='fit', mtiltlabel='fit w/tilt', dlabel='data', figtitle=True)

function to determine the channels per degree using a linear fit of the function nchannel = center_ch+chdeg*tan(angles) or the equivalent including a detector tilt

Parameters: angles: array-like

detector angles for which the position of the beam was measured

channels: array-like

detector channels where the beam was found

stdev: array-like, optional

standard deviation of the beam position

plot: bool, optional

flag to specify if a visualization of the fit should be done

usetilt: bool, optional

whether to use model considering a detector tilt, i.e. deviation angle of the pixel

direction from orthogonal to the primary beam (default: True)

Returns: pixelwidth: float

the width of one detector channel @ 1m distance, which is negative in case the hit

channel number decreases upon an increase of the detector angle.

centerch: float

center channel of the detector

tilt: float

tilt of the detector from perpendicular to the beam (will be zero in case of usetilt=False)

Note

L/pixelwidth*pi/180 = channel/degree for large detector distance with the sample detector disctance L

Other datap: str, optional

Parameters: plot format of data points

modelline: str, optional
plot format of modelline
modeltilt: str, optional
plot format of modeltilt
fignum: int or str, optional

figure number to use for the plot

mlabel: str

label of the model w/o tilt to be used in the plot

mtiltlabel: str

label of the model with tilt to be used in the plot

dlabel: str

label of the data line to be used in the plot

figtitle: bool

flag to tell if the figure title should show the fit parameters

xrayutilities.analysis.sample_align.psd_refl_align (primarybeam, angles, channels, plot=True) function which calculates the angle at which the sample is parallel to the beam from various angles and detector channels from the reflected beam. The function can be used during the half beam alignment with a linear detector.

Parameters: primarybeam: int

primary beam channel number

angles: list or array-like incidence angles channels: list or array-like

corresponding detector channels

plot: bool, optional

flag to specify if a visualization of the fit is wanted default : True

Returns: float

angle at which the sample is parallel to the beam

Examples

```
>>> psd_refl_align(500, [0, 0.1, 0.2, 0.3], [550, 600, 640, 700])
```

Module contents

xrayutilities.analysis is a package for assisting with the analysis of x-ray diffraction data, mainly reciprocal space maps

Routines for obtaining line cuts from gridded reciprocal space maps are offered, with the ability to integrate the intensity perpendicular to the line cut direction.

xrayutilities.io package

Submodules

xrayutilities.io.cbf module

```
class xrayutilities.io.cbf.CBFDirectory (datapath, ext='cbf', **keyargs)
```

Bases: xrayutilities.io.filedir.FileDirectory

Parses a directory for CBF files, which can be stored to a HDF5 file for further usage

class xrayutilities.io.cbf.CBFFile (fname, nxkey='X-Binary-Size-Fastest-Dimension',
nykey='X-Binary-Size-Second-Dimension', dtkey='DataType', path=None)

Bases: object

ReadData ()

Read the CCD data into the .data object this function is called by the initialization

```
Save2HDF5 (h5f, group='/', comp=True)
```

Saves the data stored in the EDF file in a HDF5 file as a HDF5 array. By default the data is stored in the root group of the HDF5 file - this can be changed by passing the name of a target group or a path to the target group via the "group" keyword argument.

Parameters: h5f: file-handle or str

a HDF5 file object or name

group: str, optional

group where to store the data (default to the root of the file)

comp: bool, optional

activate compression - true by default

xrayutilities.io.desy_tty08 module

class for reading data + header information from tty08 data files

tty08 is a system used at beamline P08 at Hasylab Hamburg and creates simple ASCII files to save the data. Information is easily read from the multicolumn data file, the functions below enable also to parse the information of the header

xrayutilities.io.desy_tty08.gettty08_scan (scanname, scannumbers, *args, **keyargs)

function to obtain the angular cooridinates as well as intensity values saved in TTY08 datafiles. Especially usefull for reciprocal space map measurements, and to combine date from several scans

further more it is possible to obtain even more positions from the data file if more than two string arguments with its names are given

Parameters: scanname: str

name of the scans, for multiple scans this needs to be a template string

scannumbers: int, tuple or list

number of the scans of the reciprocal space map

args: str, optional

names of the motors, to read reciprocal space maps measured in coplanar diffraction give:

- *omname*: the name of the omega motor (or its equivalent)
- ttname: the name of the two theta motor (or its equivalent)

keyargs: dict, optional

keyword arguments are passed on to tty08File

Returns: [ang1, ang2, ...]: list, optional

> angular positions of the center channel of the position sensitive detector (numpy.ndarray 1D), omitted if no args are given

MAP: ndarray

All the data values as stored in the data file (includes the intensities e.g. MAP['MCA']).

Examples

```
>>> [om, tt], MAP = xu.io.gettty08_scan('text%05d.dat', 36, 'omega',
>>>
                                         'gamma')
```

class xrayutilities.io.desy tty08.tty08File (filename, path=None, mcadir=None)

Bases: object

Represents a tty08 data file. The file is read during the Constructor call. This class should work for data stored at beamline P08 using the tty08 acquisition system.

Parameters: filename: str

tty08-filename

mcadir: str, optional

directory name of MCA files

Read ()

Read the data from the file

ReadMCA ()

xrayutilities.io.edf module

```
class xrayutilities.io.edf.EDFDirectory (datapath, ext='edf', **keyargs)
```

Bases: xrayutilities.io.filedir.FileDirectory

Parses a directory for EDF files, which can be stored to a HDF5 file for further usage

class xrayutilities.io.edf.EDFFile (fname, nxkey='Dim_1', nykey='Dim_2', dtkey='DataType', path=", header=True, keep open=False)

Bases: object

Parse ()

Parse file to find the number of entries and read the respective header information

ReadData (nimg=0)

Read the CCD data of the specified image and return the data this function is called automatically when the 'data' property is accessed, but can also be called manually when only a certain image from the file is needed.

Parameters: nimg: int, optional

number of the image which should be read (starts with 0)

Save2HDF5 (h5f, group='/', comp=True)

Saves the data stored in the EDF file in a HDF5 file as a HDF5 array. By default the data is stored in the root group of the HDF5 file - this can be changed by passing the name of a target group or a path to the target group via the "group" keyword argument.

Parameters: h5f: file-handle or str

a HDF5 file object or name

group: str, optional

group where to store the data (default to the root of the file)

comp: bool, optional

activate compression - true by default

property data

xrayutilities.io.fastscan module

modules to help with the analysis of FastScan data acquired at the ESRF. FastScan data are X-ray data (various detectors possible) acquired during scanning the sample in real space with a Piezo Scanner. The same functions might be used to analyze traditional SPEC mesh scans.

The module provides three core classes:

- FastScan
- FastScanCCD
- FastScanSeries

where the first two are able to parse single mesh/FastScans when one is interested in data of a single channel detector or are detector and the last one is able to parse full series of such mesh scans with either type of detector

see examples/xrayutilities_kmap_ESRF.py for an example script

class xrayutilities.io.fastscan.FastScan (filename, scannr, xmotor='adcX', ymotor='adcY', path=")

Bases: object

class to help parsing and treating fast scan data. FastScan is the aquisition of X-ray data while scanning the sample with piezo stages in real space. It's is available at several beamlines at the ESRF synchrotron light-source.

grid2D (nx, ny, **kwargs)

function to grid the counter data and return the gridded X, Y and Intensity values.

Parameters: nx, ny : int

number of bins in x, and y direction

counter: str, optional

name of the counter to use for gridding (default: 'mpx4int' (ID01))

gridrange: tuple, optional

range for the gridder: format: ((xmin, xmax), (ymin, ymax))

Returns: Gridder2D

Gridder2D object with X, Y, data on regular x, y-grid

motorposition (motorname)

read the position of motor with name given by motorname from the data file. In case the motor is included in the data columns the returned object is an array with all the values from the file (although retrace clean is respected if already performed). In the case the motor is not moved during the scan only one value is returned.

Parameters: motorname: str

name of the motor for which the position is wanted

Returns: ndarray

motor position(s) of motor with name motorname during the scan

parse ()

parse the specifie for the scan number specified in the constructor and store the needed informations in the object properties

retrace_clean()

function to clean the data of the scan from retrace artifacts created by the zig-zag scanning motion of the piezo actuators the function cleans the xvalues, yvalues and data attribute of the FastScan object.

class xrayutilities.io.fastscan.FastScanCCD (*args, **kwargs)

Bases: xrayutilities.io.fastscan.FastScan

class to help parsing and treating fast scan data including CCD frames. FastScan is the aquisition of X-ray data while scanning the sample with piezo stages in real space. It's is available at several beamlines at the ESRF synchrotron light-source. During such fast scan at every grid point CCD frames are recorded and need to be analyzed

getCCD (ccdnr, roi=None, datadir=None, keepdir=0, replacedir=None, nav=[1, 1], filterfunc=None)

function to read the ccd files and return the raw X, Y and DATA values. DATA represents a 3D object with first dimension representing the data point index and the remaining two dimensions representing detector channels

Parameters: ccdnr: array-like or str

array with ccd file numbers of length length(FastScanCCD.data) OR a string with the data column name for the file ccd-numbers

roi: tuple, optional

region of interest on the 2D detector. should be a list of lower and upper bounds of detector channels for the two pixel directions (default: None)

datadir: str, optional

the CCD filenames are usually parsed from the SPEC file. With this option the directory used for the data can be overwritten. Specify the datadir as simple string. Alternatively the innermost directory structure can be automatically taken from the specfile. If this is needed specify the number of directories which should be kept using the keepdir option.

keepdir: int, optional

number of directories which should be taken from the SPEC file. (default: 0)

replacedir: int, optional

number of outer most directory names which should be replaced in the output (default = None). One can either give keepdir, or replacedir, with replace taking preference if both are given.

nav: tuple or list, optional

number of detector pixel which will be averaged together (reduces the date size)

filterfunc : callable

function applied to the CCD-frames before any processing. this function should take a single argument which is the ccddata which need to be returned with the same shape! e.g. remove hot pixels, flat/darkfield correction

Returns: X, Y: ndarray

x, y-array (1D)

DATA : ndarray

3-dimensional data object

getccdFileTemplate (specscan, datadir=None, keepdir=0, replacedir=None)

function to extract the CCD file template string from the comment in the SPEC-file scan-header.

Parameters: specscan: SpecScan

spec-scan object from which header the CCD directory should be extracted

datadir: str, optional

the CCD filenames are usually parsed from the scan object. With this option the directory used for the data can be overwritten. Specify the datadir as simple string. Alternatively the innermost directory structure can be automatically taken from the specifile. If this is needed specify the number of directories which should be kept using the keepdir option.

keepdir: int, optional

number of directories which should be taken from the specscan. (default: 0)

replacedir: int, optional

number of outer most directory names which should be replaced in the output (default = None). One can either give keepdir, or replacedir, with replace taking preference if both are given.

Returns: fmtstr:str

format string for the CCD file name using one number to build the real file name

filenr: int

starting file number

gridCCD (nx, ny, ccdnr, roi=None, datadir=None, keepdir=0, replacedir=None, nav=[1, 1], gridrange=None, filterfunc=None)

function to grid the internal data and ccd files and return the gridded X, Y and DATA values. DATA represents a 4D object with first two dimensions representing X, Y and the remaining two dimensions representing detector channels

Parameters: nx, ny : int

number of bins in x, and y direction

ccdnr: array-like or str

array with ccd file numbers of length length(FastScanCCD.data) OR a string with the data column name for the file ccd-numbers

roi: tuple, optional

region of interest on the 2D detector. should be a list of lower and upper bounds of detector channels for the two pixel directions (default: None)

datadir: str, optional

the CCD filenames are usually parsed from the SPEC file. With this option the directory used for the data can be overwritten. Specify the datadir as simple string. Alternatively the innermost directory structure can be automatically taken from the specifile. If this is needed specify the number of directories which should be kept using the keepdir option.

keepdir: int, optional

number of directories which should be taken from the SPEC file. (default: 0)

replacedir: int, optional

number of outer most directory names which should be replaced in the output (default = None). One can either give keepdir, or replacedir, with replace taking preference if both are given.

nav: tuple or list, optional

number of detector pixel which will be averaged together (reduces the date size)

gridrange: tuple

range for the gridder: format: ((xmin, xmax), (ymin, ymax))

filterfunc: callable

function applied to the CCD-frames before any processing. this function should take a single argument which is the ccddata which need to be returned with the same shape! e.g. remove hot pixels, flat/darkfield correction

Returns: X, Y: ndarray

regular x, y-grid

DATA: ndarray

4-dimensional data object

processCCD (ccdnr, roi, datadir=None, keepdir=0, replacedir=None, filterfunc=None) function to read a region of interest (ROI) from the ccd files and return the raw X, Y and intensity from ROI.

Parameters: ccdnr: array-like or str

array with ccd file numbers of length length(FastScanCCD.data) OR a string with the data column name for the file ccd-numbers

roi: tuple or list

region of interest on the 2D detector. Either a list of lower and upper bounds of detector channels for the two pixel directions as tuple or a list of mask arrays

datadir: str, optional

the CCD filenames are usually parsed from the SPEC file. With this option the directory used for the data can be overwritten. Specify the datadir as simple string. Alternatively the innermost directory structure can be automatically taken from the specifile. If this is needed specify the number of directories which should be kept using the keepdir option.

keepdir: int, optional

number of directories which should be taken from the SPEC file. (default: 0)

replacedir: int, optional

number of outer most directory names which should be replaced in the output (default = None). One can either give keepdir, or replacedir, with replace taking preference if both are given.

filterfunc: callable, optional

function applied to the CCD-frames before any processing. this function should take a single argument which is the ccddata which need to be returned with the same shape! e.g. remove hot pixels, flat/darkfield correction

Returns: X, Y, DATA: ndarray

x, y-array (1D) as well as 1-dimensional data object

class xrayutilities.io.fastscan.FastScanSeries (filenames, scannrs, nx, ny, *args, **kwargs)

Bases: object

class to help parsing and treating a series of fast scan data including CCD frames. FastScan is the aquisition of X-ray data while scanning the sample with piezo stages in real space. It's is available at several beamlines at the ESRF synchrotron light-source. During such fast scan at every grid point CCD frames are recorded and need to be analyzed.

For the series of FastScans we assume that they are measured at different goniometer angles and therefore transform the data to reciprocal space.

align (deltax, deltay)

Since a sample drift or shift due to rotation often occurs between different FastScans it should be corrected before combining them. Since determining such a shift is not straight-forward in general the user needs to supply the routine with the shifts in order correct the x, y-values for the different FastScans. Such a routine could for example use the integrated CCD intensities and determine the shift using a cross-convolution.

Parameters: deltax, deltay: list

list of shifts in x/y-direction for every FastScan in the data structure

getCCDFrames (posx, posy, typ='real')

function to determine the list of ccd-frame numbers for a specific real space position. The real space position must be within the data limits of the FastScanSeries otherwise an ValueError is thrown

Parameters: posx: float

real space x-position or index in x direction

posy: float

real space y-position or index in y direction

typ: {'real', 'index'}, optional

type of coordinates. specifies if the position is specified as real space coordinate or as

index. (default: 'real')

Returns: list

<code>[[motorpos1, ccdnrs1], [motorpos2, ccdnrs2], ...]</code> where motorposN is from the N-ths FastScan in the series and ccdnrsN is the list of according CCD-frames

get_average_RSM (qnx, qny, qnz, qconv, datadir=None, keepdir=0, replacedir=None, roi=None, nav=(1, 1), filterfunc=None)

function to return the reciprocal space map data averaged over all x, y positions from a series of FastScan measurements. It necessary to give the QConversion-object to be used for the reciprocal space conversion. The QConversion-object is expected to have the 'area' conversion routines configured properly. This function needs to read all detector images, so be prepared to lean back for a moment!

Parameters: qnx, qny, qnz : int

number of points used for the 3D Gridder

qconv : QConversion

QConversion-object to be used for the conversion of the CCD-data to reciprocal space

roi: tuple, optional

region of interest on the 2D detector. should be a list of lower and upper bounds of detector channels for the two pixel directions (default: None)

nav: tuple or list, optional

number of detector pixel which will be averaged together (reduces the date size)

filterfunc : callable, optional

function applied to the CCD-frames before any processing. this function should take a single argument which is the ccddata which need to be returned with the same shape! e.g. remove hot pixels, flat/darkfield correction

datadir: str, optional

the CCD filenames are usually parsed from the SPEC file. With this option the directory used for the data can be overwritten. Specify the datadir as simple string. Alternatively the innermost directory structure can be automatically taken from the specfile. If this is needed specify the number of directories which should be kept/replaced using the keepdir/replacedir option.

keepdir: int, optional

number of directories which should be taken from the SPEC file. (default: 0)

replacedir: int, optional

number of outer most directory names which should be replaced in the output (default = None). One can either give keepdir, or replacedir, with replace taking preference if both are given.

Returns: Gridder3D

gridded reciprocal space map

get_sxrd_for_grange (qrange, qconv, datadir=None, keepdir=0, replacedir=None, roi=None, nav=(1, 1),
filterfunc=None)

function to return the real space data averaged over a certain q-range from a series of FastScan measurements. It necessary to give the QConversion-object to be used for the reciprocal space conversion. The QConversion-object is expected to have the 'area' conversion routines configured properly.

Note

This function assumes that all FastScans were performed in the same real space positions, no gridding or aligning is performed!

Parameters: grange: list or tuple

q-limits defining a box in reciprocal space. six values are needed: [minx, maxx, miny,

..., maxz]

qconv: QConversion

QConversion object to be used for the conversion of the CCD-data to reciprocal

space

roi: tuple, optional

region of interest on the 2D detector. should be a list of lower and upper bounds of

detector channels for the two pixel directions (default: None)

nav: tuple or list, optional

number of detector pixel which will be averaged together (reduces the date size)

filterfunc : callable, optional

function applied to the CCD-frames before any processing. this function should take a single argument which is the ccddata which need to be returned with the same shape!

e.g. remove hot pixels, flat/darkfield correction

datadir: str, optional

the CCD filenames are usually parsed from the SPEC file. With this option the directory used for the data can be overwritten. Specify the datadir as simple string. Alternatively the innermost directory structure can be automatically taken from the specfile. If this is needed specify the number of directories which should be kept/replaced using the keepdir/replacedir option.

keepdir: int, optional

number of directories which should be taken from the SPEC file. (default: 0)

replacedir: int, optional

number of outer most directory names which should be replaced in the output (default = None). One can either give keepdir, or replacedir, with replace taking preference if

both are given.

Returns: xvalues, yvalues, data: ndarray

x, y, and data values

grid2Dall (nx, ny, **kwargs)

function to grid the counter data and return the gridded X, Y and Intensity values from all the FastScanSeries.

Parameters: nx, ny : int

number of bins in x, and y direction

counter: str, optional

name of the counter to use for gridding (default: 'mpx4int' (ID01))

gridrange: tuple, optional

range for the gridder: format: ((xmin, xmax), (ymin, ymax))

Returns: Gridder2D

object with X, Y, data on regular x, y-grid

gridrsm (posx, posy, qnx, qny, qnz, qconv, roi=None, nav=[1, 1], typ='real', filterfunc=None, **kwargs) function to calculate the reciprocal space map at a certain x, y-position from a series of FastScan measurements

it is necessary to specify the number of grid-oints for the reciprocal space map and the QConversion-object to be used for the reciprocal space conversion. The QConversion-object is expected to have the 'area' conversion routines configured properly. Parameters: posx: float

real space x-position or index in x direction

posy: float

real space y-position or index in y direction

qnx, qny, qnz: int

number of points in the Qx, Qy, Qz direction of the gridded reciprocal space map

qconv: QConversion

QConversion-object to be used for the conversion of the CCD-data to reciprocal

space

roi: tuple, optional

region of interest on the 2D detector. should be a list of lower and upper bounds of detector channels for the two pixel directions (default: None)

nav: tuple or list, optional

number of detector pixel which will be averaged together (reduces the date size)

typ: {'real', 'index'}, optional

type of coordinates. specifies if the position is specified as real space coordinate or as index. (default: 'real')

filterfunc: callable, optional

function applied to the CCD-frames before any processing. this function should take a single argument which is the ccddata which need to be returned with the same shape! e.g. remove hot pixels, flat/darkfield correction

UB: ndarray

sample orientation matrix

Returns: Gridder3D

object with gridded reciprocal space map

rawRSM (posx, posy, qconv, roi=None, nav=[1, 1], typ='real', datadir=None, keepdir=0, replacedir=None, filterfunc=None, **kwargs)

function to return the reciprocal space map data at a certain x, y-position from a series of FastScan measurements. It necessary to give the QConversion-object to be used for the reciprocal space conversion. The QConversion-object is expected to have the 'area' conversion routines configured properly.

Parameters: posx: float

real space x-position or index in x direction

posy: float

real space y-position or index in y direction

qconv: QConversion

QConversion-object to be used for the conversion of the CCD-data to reciprocal

space

roi: tuple, optional

region of interest on the 2D detector. should be a list of lower and upper bounds of detector channels for the two pixel directions (default: None)

nav: tuple or list, optional

number of detector pixel which will be averaged together (reduces the date size)

typ: {'real', 'index'}, optional

type of coordinates. specifies if the position is specified as real space coordinate or as index. (default: 'real')

filterfunc: callable, optional

function applied to the CCD-frames before any processing. this function should take a single argument which is the ccddata which need to be returned with the same shape! e.g. remove hot pixels, flat/darkfield correction

UB: array-like, optional

sample orientation matrix

datadir: str, optional

the CCD filenames are usually parsed from the SPEC file. With this option the directory used for the data can be overwritten. Specify the datadir as simple string. Alternatively the innermost directory structure can be automatically taken from the specifile. If this is needed specify the number of directories which should be kept using the keepdir option.

keepdir: int, optional

number of directories which should be taken from the SPEC file. (default: 0)

replacedir: int, optional

number of outer most directory names which should be replaced in the output (default = None). One can either give keepdir, or replacedir, with replace taking preference if both are given.

Returns: qx, qy, qz : ndarray

reciprocal space positions of the reciprocal space map

ccddata: ndarray

raw data of the reciprocal space map

valuelist : ndarray

valuelist containing the ccdframe numbers and corresponding motor positions

read_motors()

read motor values from the series of fast scans

retrace_clean()

perform retrace clean for every FastScan in the series

xrayutilities.io.filedir module

class xrayutilities.io.filedir.FileDirectory (datapath, ext, parser, **keyargs)

Bases: object

Parses a directory for files, which can be stored to a HDF5 file for further usage. The file parser is given to the constructor and must provide a Save2HDF5 method.

```
Save2HDF5 (h5f, group=", comp=True)
```

Saves the data stored in the found files in the specified directory in a HDF5 file as a HDF5 arrays in a subgroup. By default the data is stored in a group given by the foldername - this can be changed by passing the name of a target group or a path to the target group via the "group" keyword argument.

Parameters: h5f: file-handle or str

a HDF5 file object or name

group: str, optional

group where to store the data (defaults to pathname if group is empty string)

comp: bool, optional

activate compression - true by default

xrayutilities.io.helper module

convenience functions to open files for various data file reader

these functions should be used in new parsers since they transparently allow to open gzipped and bzipped files

class xrayutilities.io.helper.xu_h5open (f, mode='r')

Bases: object

helper object to decide if a HDF5 file has to be opened/closed when using with a 'with' statement.

xrayutilities.io.helper.xu_open (filename, mode='rb')

function to open a file no matter if zipped or not. Files with extension '.gz', '.bz2', and '.xz' are assumed to be compressed and transparently opened to read like usual files.

Parameters: filename: str

filename of the file to open (full including path)

mode: str, optional

mode in which the file should be opened

Returns: file-handle

handle of the opened file

Raises: IOError

If the file does not exist an IOError is raised by the open routine, which is not caught

within the function

xrayutilities.io.ill_numor module

module for reading ILL data files (station D23): numor files

class xrayutilities.io.ill_numor.numorFile (filename, path=None)

Bases: object

Represents a ILL data file (numor). The file is read during the Constructor call. This class should work for created at station D23 using the mad acquisition system.

Parameters: filename: str

a string with the name of the data file

Read ()

Read the data from the file

columns = {0: ('detector', 'monitor', 'time', 'gamma', 'omega', 'psi'), 1: ('detector', 'monitor', 'time', 'gamma'), 2: ('detector', 'monitor', 'time', 'omega'), 5: ('detector', 'monitor', 'time', 'psi')}

getline (fid)

ssplit (string)

multispace split. splits string at two or more spaces after stripping it.

xrayutilities.io.ill_numor.numor_scan (scannumbers, *args, **kwargs)

function to obtain the angular cooridinates as well as intensity values saved in numor datafiles. Especially useful for combining several scans into one data object.

Parameters: scannumbers: int or str or iterable

number of the numors, or list of numbers. This will be transformed to a string and used

as a filename

args: str, optional

names of the motors e.g.: 'omega', 'gamma'

kwargs: dict

keyword arguments are passed on to numorFile. e.g. 'path' for the files directory

Returns: [ang1, ang2, ...] : *list*

angular positions list, omitted if no args are given

data: ndarray

all the data values.

Examples

>>> [om, gam], data = xu.io.numor_scan(414363, 'omega', 'gamma')

xrayutilities.io.imagereader module

class xrayutilities.io.imagereader.ImageReader (nop1, nop2, hdrlen=0, flatfield=None,
darkfield=None, dtype=<class 'numpy.int16'>, byte_swap=False)

Bases: object

parse CCD frames in the form of tiffs or binary data (*.bin) to numpy arrays. ignore the header since it seems to contain no useful data

The routine was tested so far with

- 1. RoperScientific files with 4096x4096 pixels created at Hasylab Hamburg, which save an 16bit integer per point.
- 2. Perkin Elmer images created at Hasylab Hamburg with 2048x2048 pixels.

readImage (filename, path=None)

read image file and correct for dark- and flatfield in case the necessary data are available. returned data = ((image data)-(darkfield))/flatfield*average(flatfield)

Parameters: filename: str

filename of the image to be read. so far only single filenames are supported. The data might be compressed. supported extensions: .tif, .bin and .bin.xz

path : str, optional
 path of the data files

class xrayutilities.io.imagereader.PerkinElmer (**keyargs)

Bases: xrayutilities.io.imagereader.ImageReader

parse PerkinElmer CCD frames (*.tif) to numpy arrays Ignore the header since it seems to contain no useful data

The routine was tested only for files with 2048x2048 pixel images created at Hasylab Hamburg which save an 32bit float per point.

class xrayutilities.io.imagereader.Pilatus100K (**keyargs)

Bases: xrayutilities.io.imagereader.ImageReader

parse Dectris Pilatus 100k frames (*.tiff) to numpy arrays Ignore the header since it seems to contain no useful data

```
class xrayutilities.io.imagereader.RoperCCD (**keyargs)
```

Bases: xrayutilities.io.imagereader.ImageReader

parse RoperScientific CCD frames (*.bin) to numpy arrays Ignore the header since it seems to contain no useful data

The routine was tested only for files with 4096x4096 pixel images created at Hasylab Hamburg which save an 16bit integer per point.

class xrayutilities.io.imagereader.TIFFRead (filename, path=None)

Bases: xrayutilities.io.imagereader.ImageReader

class to Parse a TIFF file including extraction of information from the file header in order to determine the image size and data type

The data stored in the image are available in the 'data' property.

xrayutilities.io.imagereader.get_tiff (filename, path=None)

read tiff image file and return the data

Parameters: filename : str

filename of the image to be read. so far only single filenames are supported. The data

might be compressed.

path of the data file

path: str, optional

xrayutilities.io.panalytical_xml module

Panalytical XML (www.XRDML.com) data file parser

based on the native python xml.dom.minidom module. want to keep the number of dependancies as small as possible

class xrayutilities.io.panalytical_xml.XRDMLFile (fname, path=")

Bases: object

class to handle XRDML data files. The class is supplied with a file name and uses the XRDMLScan class to parse the xrdMeasurement in the file

class xrayutilities.io.panalytical_xml.XRDMLMeasurement (measurement, namespace=")

Bases: object

class to handle scans in a XRDML datafile

xrayutilities.io.panalytical_xml.getxrdml_map (filetemplate, scannrs=None, path='.', roi=None) parses multiple XRDML file and concatenates the results for parsing the xrayutilities.io.XRDMLFile class is used. The function can be used for parsing maps measured with the PIXCel 1D detector (and in limited way also for data acquired with a point detector -> see getxrdml_scan instead).

Parameters: filetemplate: str

template string for the file names, can contain a %d which is replaced by the scan number or be a list of filenames

scannrs: int or list, optional

scan number(s)

path : str, optional

common path to the filenames

roi: tuple, optional

region of interest for the PIXCel detector, for other measurements this is not useful!

Returns: om, tt, psd: ndarray

motor positions and data as flattened numpy arrays

Examples

```
xrayutilities.io.panalytical_xml.getxrdml_scan (filetemplate, *motors, **kwargs)
```

parses multiple XRDML file and concatenates the results for parsing the xrayutilities.io.XRDMLFile class is used. The function can be used for parsing arbitrary scans and will return the the motor values of the scan motor and additionally the positions of the motors given by in the *motors argument

Parameters: filetemplate: str

template string for the file names, can contain a %d which is replaced by the scan number or be a list of filenames given by the scannrs keyword argument

motors: str

motor names to return: e.g.: 'Omega', '2Theta', ... one can also use abbreviations:

- 'Omega' = 'om' = 'o'
- '2Theta' = 'tt' = 't'
- 'Chi' = 'c'
- 'Phi' = 'p'

scannrs: int or list, optional

scan number(s)

path: str, optional

common path to the filenames

Returns: scanmot, mot1, mot2,..., detectorint : ndarray

motor positions and data as flattened numpy arrays

Examples

xrayutilities.io.pdcif module

class xrayutili ties.io.pdcif.pdCIF (filename, datacolumn=None)

Bases: object

the class implements a primitive parser for pdCIF-like files. It reads every entry and collects the information in the header attribute. The first loop containing one of the intensity fields is assumed to be the data the user is interested in and is transferred to the data array which is stored as numpy record array the columns can be accessed by name intensity fields:

- _pd_meas_counts_total
- _pd_meas_intensity_total
- _pd_proc_intensity_total
- _pd_proc_intensity_net
- _pd_calc_intensity_total

• _pd_calc_intensity_net alternatively the data column name can be given as argument to the constructor

Parse ()

parser of the pdCIF file. the method reads the data from the file and fills the data and header attributes with content

class xrayutilities.io.pdcif.pdESG (filename, datacolumn=None)

Bases: xrayutilities.io.pdcif.pdCIF

class for parsing multiple pdCIF loops in one file. This includes especially *.esg files which are supposed to consist of multiple loops of pdCIF data with equal length.

Upon parsing the class tries to combine the data of these different scans into a single data matrix -> same shape of subscan data is assumed

Parse ()

parser of the pdCIF file. the method reads the data from the file and fills the data and header attributes with content

xrayutilities.io.pdcif.remove_comments (line, sep='#')

xrayutilities.io.rigaku_ras module

class for reading data + header information from Rigaku RAS (3-column ASCII) files

Such datafiles are generated by the Smartlab Guidance software from Rigaku.

class xrayutilities.io.rigaku_ras.RASFile (filename, path=None)

Bases: object

Represents a RAS data file. The file is read during the constructor call

Parameters: filename: str

name of the ras-file

path : str, optional
 path to the data file

Read ()

Read the data from the file

class xrayutilities.io.rigaku_ras.RASScan (filename, pos)

Bases: object

Represents a single Scan portion of a RAS data file. The scan is parsed during the constructor call

Parameters: filename: str

file name of the data file

pos: int

seek position of the 'RAS HEADER START' line

xrayutilities.io.rigaku_ras.getras_scan (scanname, scannumbers, *args, **kwargs)

function to obtain the angular cooridinates as well as intensity values saved in RAS datafiles. Especially useful for reciprocal space map measurements, and to combine date from several scans

further more it is possible to obtain even more positions from the data file if more than two string arguments with its names are given

Parameters: scanname: str

name of the scans, for multiple scans this needs to be a template string

scannumbers: int, tuple or list

number of the scans of the reciprocal space map

args: str, optional

names of the motors, to read reciprocal space maps measured in coplanar diffraction give:

• omname: name of the omega motor (or its equivalent)

ttname: name of the two theta motor (or its equivalent)

kwargs: dict

keyword arguments forwarded to RASFile function

Returns: [ang1, ang2, ...]: list

angular positions are extracted from the respective scan header, or motor positions during the scan. this is omitted if no *args* are given

rasdata: ndarray

the data values (includes the intensities e.g. rasdata['int']).

Examples

```
>>> [om, tt], MAP = xu.io.getras_scan('text%05d.ras', 36, 'Omega',
>>>
```

xrayutilities.io.rotanode_alignment module

parser for the alignment log file of the rotating anode

```
class xrayutilities.io.rotanode_alignment.RA_Alignment (filename)
```

Bases: object

class to parse the data file created by the alignment routine (tpalign) at the rotating anode spec installation this routine does an iterative alignment procedure and saves the center of mass values were it moves after each scan. It iterates between two different peaks and iteratively aligns at each peak between two different motors (om/chi at symmetric peaks, om/phi at asymmetric peaks)

Parse ()

parser to read the alignment log and obtain the aligned values at every iteration.

get (key)

keys ()

returns a list of keys for which aligned values were parsed

plot (pname)

function to plot the alignment history for a given peak

Parameters: pname: str

peakname for which the alignment should be plotted

xrayutilities.io.seifert module

a set of routines to convert Seifert ASCII files to HDF5 in fact there exist two posibilities how the data is stored (depending on the use detector):

- 1. as a simple line scan (using the point detector)
- 2. as a map using the PSD

In the first case the data ist stored

```
class xrayutilities.io.seifert.SeifertHeader
Bases: object
helper class to represent a Seifert (NJA) scan file header

class xrayutilities.io.seifert.SeifertMultiScan (filename, m_scan, m2, path=")
Bases: object
Class to parse a Seifert (NJA) multiscan file

parse ()

class xrayutilities.io.seifert.SeifertScan (filename, path=")
Bases: object
Class to parse a single Seifert (NJA) scan file
```

parse ()

```
xrayutilities.io.seifert.getSeifert_map (filetemplate, scannrs=None, path='.', scantype='map',
Nchannels=1280)
```

parses multiple Seifert *.nja files and concatenates the results. for parsing the xrayutilities.io.SeifertMultiScan class is used. The function can be used for parsing maps measured with the Meteor1D and point detector.

Parameters: filetemplate: str

template string for the file names, can contain a %d which is replaced by the scan

number or be a list of filenames

scannrs: int or list, optional

scan number(s)

path: str, optional

common path to the filenames

scantype: {'map', 'tsk'}, optional

type of datafile: can be either 'map' (reciprocal space map measured with a regular

Seifert job (default)) or 'tsk' (MCA spectra measured using the TaskInterpreter)

Nchannels: int, optional

number of channels of the MCA (needed for 'tsk' measurements)

Returns: om, tt, psd: ndarray

positions and data as flattened numpy arrays

Examples

```
xrayutilities.io.seifert.repair_key (key)
```

Repair a key string in the sense that the string is changed in a way that it can be used as a valid Python identifier. For that purpose all blanks within the string will be replaced by _ and leading numbers get an preceding _.

xrayutilities.io.spec module

a class for observing a SPEC data file

Motivation:

SPEC files can become quite large. Therefore, subsequently reading the entire file to extract a single scan is a quite cumbersome procedure. This module is a proof of concept code to write a file observer starting a reread of the file starting from a stored offset (last known scan position)

```
class xrayutilities.io.spec.specCmdLine (n, prompt, cmdl, out=")
Bases: object
```

class xrayutilities.io.spec.specfile (filename, path=")

Bases: object

This class represents a single SPEC file. The class provides methodes for updateing an already opened file which makes it particular interesting for interactive use.

Parse ()

Parses the file from the starting at last_offset and adding found scans to the scan list.

Save2HDF5 (h5f, comp=True, optattrs={})

Save the entire file in an HDF5 file. For that purpose a group is set up in the root group of the file with the name of the file without extension and leading path. If the method is called after an previous update only the scans not written to the file meanwhile are saved.

Parameters: h5f: file-handle or str

a HDF5 file object or its filename

comp: bool, optional

activate compression - true by default

Update ()

reread the file and add newly added files. The parsing starts at the data offset of the last scan gathered during the last parsing run.

class xrayutilities.io.spec.specLog (filename, prompt, path=")

Bases: object

class to parse a SPEC log file to find the command history

Parse ()

class xrayutilities.io.spec.specscan (name, scannr, command, date, time, itime, colnames, hoffset, doffset, fname, imopnames, imopvalues, scan_status)

Bases: object

Represents a single SPEC scan. This class is usually not called by the user directly but used via the SPECFile class.

ClearData ()

Delete the data stored in a scan after it is no longer used.

ReadData ()

Set the data attribute of the scan class.

Save2HDF5 (h5f, group='/', title=", optattrs={}, comp=True)

Save a SPEC scan to an HDF5 file. The method creates a group with the name of the scan and stores the data there as a table object with name "data". By default the scan group is created under the root group of the HDF5 file. The title of the scan group is ususally the scan command. Metadata of the scan are stored as attributes to the scan group. Additional custom attributes to the scan group can be passed as a dictionary via the optattrs keyword argument.

Parameters: h5f: file-handle or str

a HDF5 file object or its filename

group: str, optional

name or group object of the HDF5 group where to store the data

title: str, optional

a string with the title for the data, defaults to the name of scan if empty

optattrs: dict, optional

a dictionary with optional attributes to store for the data

comp: bool, optional

activate compression - true by default

SetMCAParams (mca_column_format, mca_channels, mca_start, mca_stop)

Set the parameters used to save the MCA data to the file. This method calculates the number of lines used to store the MCA data from the number of columns and the

Parameters: mca_column_format: int

number of columns used to save the data

mca_channels: int

number of MCA channels stored

mca start: int

first channel that is stored

mca_stop: int

last channel that is stored

getheader element (key, firstonly=True)

return the value-string of the first appearance of this SPECScan's header element, or a list of all values if firstonly=False

Parameters: specscan: SPECScan

key: str

name of the key to return; e.g. 'UMONO' or 'D'

firstonly: bool, optional

flag to specify if all instances or only the first one should be returned

Returns: valuestring: str

header value (if firstonly=True)

[str1, str2, ...]: list

header values (if firstonly=False)

plot (*args, **keyargs)

Plot scan data to a matplotlib figure. If newfig=True a new figure instance will be created. If logy=True (default is False) the y-axis will be plotted with a logarithmic scale.

Parameters: args: list

arguments for the plot: first argument is the name of x-value column the following pairs of arguments are the y-value names and plot styles allowed are 3, 5, 7,... number of arguments

keyargs: dict, optional

newfig: bool, optional

if True a new figure instance will be created otherwise an existing one will be used

logy: bool, optional

if True a semilogy plot will be done

xrayutilities.io.spec.geth5_scan (h5f, scans, *args, **kwargs)

function to obtain the angular cooridinates as well as intensity values saved in an HDF5 file, which was created from a spec file by the Save2HDF5 method. Especially useful for reciprocal space map measurements.

further more it is possible to obtain even more positions from the data file if more than two string arguments with its names are given

Parameters: h5f: file-handle or str

file object of a HDF5 file opened using h5py or its filename

scans: int, tuple or list

number of the scans of the reciprocal space map

args: str, optional

names of the motors. to read reciprocal space maps measured in coplanar diffraction give:

omname: name of the omega motor (or its equivalent)

ttname: name of the two theta motor (or its equivalent)

kwargs: dict, optional

samplename: str, optional

string with the hdf5-group containing the scan data if ommitted the first child node of h5f.root will be used

rettype: {'list', 'numpy'}, optional

how to return motor positions. by default a list of arrays is returned. when rettype == 'numpy' a record array will be returned.

Returns: [ang1, ang2, ...] : *list*

angular positions of the center channel of the position sensitive detector

(numpy.ndarray 1D), this list is omitted if no args are given

MAP: ndarray

the data values as stored in the data file (includes the intensities e.g. MAP['MCA']).

Examples

```
>>> [om, tt], MAP = xu.io.geth5_scan(h5file, 36, 'omega', 'gamma')
```

xrayutilities.io.spec.getspec_scan (specf, scans, *args, **kwargs)

function to obtain the angular cooridinates as well as intensity values saved in a SPECFile. Especially useful to combine the data from multiple scans.

further more it is possible to obtain even more positions from the data file if more than two string arguments with its names are given

Parameters: specf : SPECFile

file object

scans: int, tuple or list number of the scans

args: str

names of the motors and counters

rettype: {'list', 'numpy'}, optional

how to return motor positions. by default a list of arrays is returned. when rettype ==

'numpy' a record array will be returned.

Returns: [ang1, ang2, ...] : *list*

coordinates and counters from the SPEC file

Examples

```
>>> [om, tt, cnt2] = xu.io.getspec_scan(s, 36, 'omega', 'gamma',
>>>
```

xrayutilities.io.spectra module

module to handle spectra data

class xrayutilities.io.spectra.spectrafile (filename, mcatmp=None, mcastart=None, mcastop=None)

Bases: object

Represents a SPECTRA data file. The file is read during the Constructor call. This class should work for data stored at beamlines P08 and BW2 at HASYLAB.

Parameters: filename: str

a string with the name of the SPECTRA file

mcatmp: str, optional

template for the MCA files

mcastart, mcastop: int, optional

start and stop index for the MCA files, if not given, the class tries to determine the start and stop index automatically.

Read ()

Read the data from the file.

ReadMCA ()

Save2HDF5 (h5file, name, group='/', mcaname='MCA')

Saves the scan to an HDF5 file. The scan is saved to a seperate group of name "name". h5file is either a string for the file name or a HDF5 file object. If the mca attribute is not None mca data will be stored to an chunked array of with name mcaname.

Parameters: h5file: file-handle or str

HDF5 file object or name

name: str

name of the group where to store the data

group: str, optional

root group where to store the data

mcaname: str, optional

Name of the MCA in the HDF5 file

Returns: bool or None

The method returns None in the case of everything went fine, True otherwise.

class xrayutilities.io.spectra.SPECTRAFileComments

Bases: dict

Class that describes the comments in the header of a SPECTRA file. The different comments are accessible via the comment keys.

class xrayutilities.io.spectra.SPECTRAFileData

Bases: object

append (col)

class xrayutilities.io.spectra.SPECTRAFileDataColumn (index, name, unit, type)

Bases: object

<code>class xrayutilities.io.spectra.SPECTRAFileParameters () -> new empty dictionary dict(mapping) -> new dictionary initialized from a mapping object's (key, value) pairs dict(iterable) -> new dictionary initialized as if via: $d = \{\}$ for k, v in iterable: $d[k] = v \ dict(**kwargs) -> new \ dictionary initialized with the name=value pairs in the keyword argument list. For example: <math>d[c] = v \ dict(one=1, two=2)$ </code>

Bases: dict

xrayutilities.io.spectra.geth5_spectra_map (h5file, scans, *args, **kwargs)

function to obtain the omega and twotheta as well as intensity values for a reciprocal space map saved in an HDF5 file, which was created from a spectra file by the Save2HDF5 method.

further more it is possible to obtain even more positions from the data file if more than two string arguments with its names are given

Parameters: h5f: file-handle or str

file object of a HDF5 file opened using h5py

scans: int, tuple or list

number of the scans of the reciprocal space map

args: str, optional

arbitrary number of motor names

• omname: name of the omega motor (or its equivalent)

• ttname: name of the two theta motor (or its equivalent)

kwargs: dict, optional

mca: str, optional

name of the mca data (if available) otherwise None (default: "MCA")

samplename: str. optional

string with the hdf5-group containing the scan data if omitted the first child node of

h5f.root will be used to determine the sample name

Returns: [ang1, ang2, ...] : *list*

angular positions of the center channel of the position sensitive detector

(numpy.ndarray 1D). one entry for every args-argument given to the function

MAP: ndarray

the data values as stored in the data file (includes the intensities e.g. MAP['MCA']).

Module contents

xrayutilities.materials package

Submodules

xrayutilities.materials.atom module

module containing the Atom class which handles the database access for atomic scattering factors and the atomic mass.

class xrayutilities.materials.atom.Atom (name, num)

Bases: object

property color

f (q, en='config')

function to calculate the atomic structure factor F

Parameters: q: float, array-like

momentum transfer

en: float or str, optional

energy for which F should be calculated, if omitted the value from the xrayutilities

configuration is used

Returns: float or array-like

value(s) of the atomic structure factor

£0 (q)

f1 (en='config')

```
£2 (en='config')
```

get_cache (prop, key)

check if a cached value exists to speed up repeated database requests

Returns: bool

True then result contains the cached otherwise False and result is None

result: database value

 $max_cache_length = 1000$

property radius

set_cache (prop, key, result)

set result to be cached to speed up future calls

property weight

xrayutilities.materials.atom.get_key(*args)

generate a hash key for several possible types of arguments

xrayutilities.materials.cif module

class xrayutilities.materials.cif.CIFDataset (fid, name, digits)

Bases: object

class for parsing CIF (Crystallographic Information File) files. The class aims to provide an additional way of creating material classes instead of manual entering of the information the lattice constants and unit cell structure are parsed from the CIF file

Parse (fid)

function to parse a CIF data set. The function reads the space group symmetry operations and the basic atom positions as well as the lattice constants and unit cell angles

SGLattice (use_p1=False)

create a SGLattice object with the structure from the CIF file

SymStruct ()

function to obtain the list of different atom positions in the unit cell for the different types of atoms and determine the space group number and origin choice if available. The data are obtained from the data parsed from the CIF file.

class xrayutilities.materials.cif.CIFFile (filestr, digits=4)

Bases: object

class for parsing CIF (Crystallographic Information File) files. The class aims to provide an additional way of creating material classes instead of manual entering of the information the lattice constants and unit cell structure are parsed from the CIF file.

If multiple datasets are present in the CIF file this class will attempt to parse all of them into the the data dictionary. By default all methods access the first data set found in the file.

Parse ()

function to parse a CIF file. The function reads all the included data sets and adds them to the data dictionary.

SGLattice (dataset=None, use_p1=False)

create a SGLattice object with the structure from the CIF dataset

Parameters: dataset: str. optional

name of the dataset to use. if None the default one will be used.

use_p1: bool, optional

force the use of P1 symmetry, default False

xrayutilities.materials.cif.cifexport (filename, mat)

function to export a Crystal instance to CIF file. This in particular includes the atomic coordinates, however, ignores for example the elastic parameters.

xrayutilities.materials.database module

module to handle the access to the optical parameters database

class xrayutilities.materials.database.DataBase (fname)

Bases: object

Close ()

Close an opend database file.

Create (dbname, dbdesc)

Creates a new database. If the database file already exists its content is delete.

Parameters: dbname: str

name of the database

dbdesc: str

a short description of the database

CreateMaterial (name, description)

This method creates a new material. If the material group already exists the procedure is aborted.

Parameters: name: str

name of the material

description: str

description of the material

GetF0 (q, dset='default')

Obtain the f0 scattering factor component for a particular momentum transfer q.

Parameters: q: float or array-like

momentum transfer

dset: str, optional

specifies which dataset (different oxidation states) should be used

GetF1 (en)

Return the second, energy dependent, real part of the scattering factor for a certain energy en.

Parameters: en: float or array-like

energy

GetF2 (en)

Return the imaginary part of the scattering factor for a certain energy en.

Parameters: en: float or array-like

energy

Open (mode='r')

Open an existing database file.

SetColor (color)

Save color of the element for visualization

Parameters: color: tuple, str

matplotlib color for the element

setF0 (parameters, subset='default')

Save f0 fit parameters for the set material. The fit parameters are stored in the following order: c, a1, b1,......, a4, b4

Parameters: parameters: list or array-like

fit parameters **subset**: str, optional

name the f0 dataset

SetF1F2 (en, f1, f2)

Set f1, f2 values for the active material.

Parameters: en: list or array-like

energy in (eV)

f1 : list or array-like

f1 values

f2: list or array-like

f2 values

SetMaterial (name)

Set a particular material in the database as the actual material. All operations like setting and getting optical constants are done for this particular material.

Parameters: name: str

name of the material

SetRadius (radius)

Save atomic radius for visualization

Parameters: radius: float

atomic radius in Angstrom

SetWeight (weight)

Save weight of the element as float

Parameters: weight: float

atomic standard weight of the element

xrayutilities.materials.database.add_color_from_JMOL (db, cfile, verbose=False)
Read color from JMOL color table and save it to the database.

xrayutilities.materials.database.add_f0_from_intertab (db, itf, verbose=False)
Read f0 data from International Tables of Crystallography and add it to the database.

xrayutilities.materials.database.add_f0_from_xop (db, xop, verbose=False)
Read f0 data from f0 xop.dat and add it to the database.

xrayutilities.materials.database.add_f1f2_from_ascii_file (db, asciifile, element, verbose=False)
Read f1 and f2 data for specific element from ASCII file (3 columns) and save it to the database.

xrayutilities.materials.database.add_f1f2_from_henkedb (db, hf, verbose=False) Read f1 and f2 data from Henke database and add it to the database.

xrayutilities.materials.database.add_f1f2_from_kissel (db, kf, verbose=False)
Read f1 and f2 data from Henke database and add it to the database.

xrayutilities.materials.database.add mass from NIST (db, nistfile, verbose=False)

Read atoms standard mass and save it to the database. The mass of the natural isotope mixture is taken from the NIST data!

xrayutilities.materials.database.add_radius_from_WIKI (db, dfile, verbose=False)
Read radius from Wikipedia radius table and save it to the database.

xrayutilities.materials.database.createAndFillDatabase (fname, dpath=None, verbose=False) function to create the database and fill it with values from the various source files.

Parameters: fname: str

Filename of the database to be created (including the path)

dpath: str, optional

directory where all the source data files are stored

verbose: bool, optional

flag to determine the verbosity of the script (default: False)

xrayutilities.materials.database.init_material_db (db)

xrayutilities.materials.elements module

xrayutilities.materials.heuslerlib module

implement convenience functions to define Heusler materials.

xrayutilities.materials.heuslerlib.**FullHeuslerCubic225** (X, Y, Z, a, biso=[0, 0, 0], occ=[1, 1, 1]) Full Heusler structure with formula X2YZ. Strukturberichte symbol L2_1; space group Fm-3m (225)

Parameters: X, Y, Z: str or Element

elements

a: float

cubic lattice parameter in Angstroem

biso: list of floats, optional

Debye Waller factors for X, Y, Z elements

occ: list of floats, optional

occupation numbers for the elements X, Y, Z

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.heuslerlib.**FullHeuslerCubic225_A2** (X, Y, Z, a, a2dis, biso=[0, 0, 0], occ=[1, 1, 1])

Full Heusler structure with formula X2YZ. Strukturberichte symbol L2_1; space group Fm-3m (225) with A2-type (W) disorder

Parameters: X, Y, Z: str or Element

elements

a: float

cubic lattice parameter in Angstroem

a2dis: float

amount of A2-type disorder (0: fully ordered, 1: fully disordered)

biso: list of floats, optional

Debye Waller factors for X, Y, Z elements

occ: list of floats, optional

occupation numbers for the elements X, Y, Z

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.heuslerlib.**FullHeuslerCubic225_B2** (X, Y, Z, a, b2dis, biso=[0, 0, 0], occ=[1, 1, 1])

Full Heusler structure with formula X2YZ. Strukturberichte symbol L2_1; space group Fm-3m (225) with B2-type (CsCl) disorder

Parameters: X, Y, Z: str or Element

elements

a: float

cubic lattice parameter in Angstroem

b2dis: float

amount of B2-type disorder (0: fully ordered, 1: fully disordered)

biso: list of floats, optional

Debye Waller factors for X, Y, Z elements

occ: list of floats, optional

occupation numbers for the elements X, Y, Z

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.heuslerlib.**FullHeuslerCubic225_DO3** (X, Y, Z, a, do3disxy, do3disxz, biso=[0, 0, 0], occ=[1, 1, 1])

Full Heusler structure with formula X2YZ. Strukturberichte symbol L2_1; space group Fm-3m (225) with DO_3-type (BiF3) disorder, either between atoms X <-> Y or X <-> Z.

Parameters: X, Y, Z: str or Element

elements

a: float

cubic lattice parameter in Angstroem

do3disxy: float

amount of DO_3-type disorder between X and Y atoms (0: fully ordered, 1: fully disordered)

do3disxz: float

amount of DO_3-type disorder between X and Z atoms (0: fully ordered, 1: fully disordered)

biso: list of floats, optional

Debye Waller factors for X, Y, Z elements

occ: list of floats, optional

occupation numbers for the elements X, Y, Z

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.heuslerlib.**HeuslerHexagonal194** (X, Y, Z, a, c, biso=[0, 0, 0], occ=[1, 1, 1]) Hexagonal Heusler structure with formula XYZ space group P63/mmc (194)

Parameters: X, Y, Z: str or Element

elements

a.c: float

hexagonal lattice parameters in Angstroem

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.heuslerlib.HeuslerTetragonal119 (X, Y, Z, a, c, biso=[0, 0, 0], occ=[1, 1, 1]) Tetragonal Heusler structure with formula X2YZ space group I-4m2 (119)

Parameters: X, Y, Z: str or Element

elements **a, c**: float

tetragonal lattice parameters in Angstroem

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.heuslerlib.**HeuslerTetragonal139** (X, Y, Z, a, c, biso=[0, 0, 0], occ=[1, 1, 1]) Tetragonal Heusler structure with formula X2YZ space group I4/mmm (139)

Parameters: X, Y, Z: str or Element

elements

a, c: float

tetragonal lattice parameters in Angstroem

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.heuslerlib.**InverseHeuslerCubic216** (X, Y, Z, a, biso=[0, 0, 0], occ=[1, 1, 1])

Full Heusler structure with formula (XY)X'Z structure; space group F-43m (216)

Parameters: X, Y, Z: str or Element

elements

a: float

cubic lattice parameter in Angstroem

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.material module

Classes decribing materials. Materials are devided with respect to their crystalline state in either Amorphous or Crystal types. While for most materials their crystalline state is defined few materials are also included as amorphous which can be useful for calculation of their optical properties.

class xrayutilities.materials.material.alloy (matA, matB, x)

Bases: xrayutilities.materials.material.Crystal

alloys two materials from the same crystal system. If the materials have the same space group the Wyckoff positions within the unit cell will also reflect the alloying.

RelaxationTriangle (hkl, sub, exp)

function which returns the relaxation triangle for a Alloy of given composition. Reciprocal space coordinates are calculated using the user-supplied experimental class

Parameters: hkl : list or array-like

Miller Indices **sub**: Crystal, or float

substrate material or lattice constant

exp: Experiment

object from which the Transformation object and ndir are needed

Returns: qy, qz : float

reciprocal space coordinates of the corners of the relaxation triangle

static check_compatibility (matA, matB)

static lattice_const_AB (latA, latB, x, name=")

method to calculated the interpolation of lattice parameters and unit cell angles of the Alloy. By default linear interpolation between the value of material A and B is performed.

Parameters: latA, latB: float or vector

property (lattice parameter/angle) of material A and B. A property can be a scalar or

vector.

x: float

fraction of material B in the alloy.

name: str, optional

label of the property which is interpolated. Can be 'a', 'b', 'c', 'alpha', 'beta', or

'gamma'.

property x

class xrayutilities.materials.material.Amorphous (name, density, atoms=None, cij=None)

Bases: xrayutilities.materials.material.Material

amorphous materials are described by this class

chi0 (en='config')

calculates the complex chi_0 values often needed in simulations. They are closely related to delta and beta ($n = 1 + chi_r0/2 + i*chi_i0/2 vs. n = 1 - delta + i*beta$)

delta (en='config')

function to calculate the real part of the deviation of the refractive index from 1 (n=1-delta+i*beta)

Parameters: en: float, array-like or str, optional

energy of the x-rays in eV

Returns: float or array-like

ibeta (en='config')

function to calculate the imaginary part of the deviation of the refractive index from 1 (n=1-delta+i*beta)

Parameters: en: float, array-like or str, optional

energy of the x-rays in eV

Returns: float or array-like

static parseChemForm (cstring)

Parse a string containing a simple chemical formula and transform it to a list of elements together with their relative atomic fraction. e.g. 'H2O' -> [(H, 2/3), (O, 1/3)], where H and O are the Element objects of Hydrogen and Oxygen. Note that every chemical element needs to start with a capital letter! Complicated formulas containing bracket are not supported!

Parameters: cstring: str

string containing the chemical fomula

Returns: list of tuples

chemical element and atomic fraction

xrayutilities.materials.material.Cij2Cijkl(Cij)

Converts the elastic constants matrix (tensor of rank 2) to the full rank 4 cijkl tensor.

Parameters: cij: array-like

(6, 6) cij matrix

Returns: cijkl ndarray

(3, 3, 3, 3) cijkl tensor as numpy array

xrayutilities.materials.material.Cijkl2Cij (cijkl)

Converts the full rank 4 tensor of the elastic constants to the (6, 6) matrix of elastic constants.

Parameters: cijkl ndarray

(3, 3, 3, 3) cijkl tensor as numpy array

Returns: cij: array-like

(6, 6) cij matrix

class xrayutilities.materials.material.crystal (name, lat, cij=None, thetaDebye=None)

Bases: xrayutilities.materials.material.Material

Crystalline materials are described by this class

ApplyStrain (strain)

Applies a certain strain on the lattice of the material. The result is a change in the base vectors of the real space as well as reciprocal space lattice. The full strain matrix (3x3) needs to be given.

Note

NO elastic response of the material will be considered!

property B

GetMismatch (mat)

Calculate the mismatch strain between the material and a second material

HKL (*q)

Return the HKL-coordinates for a certain Q-space position.

Parameters: q: list or array-like

Q-position. its also possible to use HKL(qx, qy, qz).

Q (*hkl)

Return the Q-space position for a certain material.

Parameters: hkl : list or array-like

Miller indices (or Q(h, k, l) is also possible)

StructureFactor (q, en='config', temp=0)

calculates the structure factor of a material for a certain momentum transfer and energy at a certain temperature of the material

Parameters: q: list, tuple or array-like

vectorial momentum transfer

en: float or str, optional

x-ray energy eV, if omitted the value from the xrayutilities configuration is used

temp: float

temperature used for Debye-Waller-factor calculation

Returns: complex

the complex structure factor

StructureFactorForEnergy (q0, en, temp=0)

calculates the structure factor of a material for a certain momentum transfer and a bunch of energies

Parameters: q0: list, tuple or array-like

vectorial momentum transfer

en : list, tuple or array-like energy values in eV

temp: float

temperature used for Debye-Waller-factor calculation

Returns: array-like

complex valued structure factor array

StructureFactorForQ (q, en0='config', temp=0)

calculates the structure factor of a material for a bunch of momentum transfers and a certain energy

Parameters: q: list of vectors or array-like

vectorial momentum transfers; list of vectores (list, tuple or array) of length 3 e.g.: (Si.Q(0, 0, 4), Si.Q(0, 0, 4.1),...) or numpy.array([Si.Q(0, 0, 4), Si.Q(0, 0, 4.1)])

en0: float or str, optional

x-ray energy eV, if omitted the value from the xrayutilities configuration is used

temp: float

temperature used for Debye-Waller-factor calculation

Returns: array-like

complex valued structure factor array

property a

property a1

property a2

property a3

property alpha

property b

property beta

property c

chemical_composition (natoms=None, with_spaces=False, ndigits=2) determine chemical composition from occupancy of atomic positions.

Parameters: mat : Crystal

instance of Crystal **natoms**: int, optional

number of atoms to normalize the formula, if None some automatic normalization is attempted using the greatest common divisor of the number of atoms per unit cell. If the number of atoms of any element is fractional natoms=1 is used.

with_spaces : bool, optional

add spaces between the different entries in the output string for CIF combatibility

ndigits: int, optional

number of digits to which floating point numbers are rounded to

Returns: str

representation of the chemical composition

chi0 (en='config')

calculates the complex chi_0 values often needed in simulations. They are closely related to delta and beta (n = 1 + $chi_0/2$ + i* $chi_0/2$ vs. n = 1 - delta + i*beta)

chih (q, en='config', temp=0, polarization='S')

calculates the complex polarizability of a material for a certain momentum transfer and energy

Parameters: q: list, tuple or array-like

momentum transfer vector in (1/A)

en: float or str, optional

x-ray energy eV, if omitted the value from the xrayutilities configuration is used

temp: float, optional

temperature used for Debye-Waller-factor calculation

polarization: {'S', 'P'}, optional sigma or pi polarization

Returns: tuple

(abs(chih_real), abs(chih_imag)) complex polarizability

dTheta (Q, en='config')

function to calculate the refractive peak shift

Parameters: Q: list, tuple or array-like

momentum transfer vector (1/A)

en: float or str, optional

x-ray energy eV, if omitted the value from the xrayutilities configuration is used

Returns: float

peak shift in degree

delta (en='config')

function to calculate the real part of the deviation of the refractive index from 1 (n=1-delta+i*beta)

Parameters: en: float or str, optional

x-ray energy eV, if omitted the value from the xrayutilities configuration is used

Returns: float

property density

calculates the mass density of an material from the mass of the atoms in the unit cell.

Returns: float

mass density in kg/m^3

distances ()

function to obtain distances of atoms in the crystal up to the unit cell size (largest value of a, b, c is the cut-off) returns a list of tuples with distance d and number of occurrence n [(d1, n1), (d2, n2),...]

Note

if the base of the material is empty the list will be empty

environment (*pos, **kwargs)

Returns a list of neighboring atoms for a given position within the unit cell. If the material does not contain any atoms a dummy atom will be placed on the unit cell corners.

Parameters: pos: list or array-like

fractional coordinate in the unit cell

maxdist: float

maximum distance wanted in the list of neighbors (default: 7)

Returns: list of tuples

(distance, atomType, multiplicity) giving distance sorted list of atoms

classmethod fromCIF (ciffilestr, **kwargs)

Create a Crystal from a CIF file. The default data-set from the cif file will be used to create the Crystal.

Parameters: ciffilestr: str, bytes

filename of the CIF file or string representation of the CIF file

kwargs: dict

keyword arguments are passed to the init-method of CIFFile

Returns: Crystal

property gamma

ibeta (en='config')

function to calculate the imaginary part of the deviation of the refractive index from 1 (n=1-delta+i*beta)

Parameters: en: float or str, optional

x-ray energy eV, if omitted the value from the xrayutilities configuration is used

Returns: float

loadLatticefromCIF (ciffilestr)

load the unit cell data (lattice) from the CIF file. Other material properties stay unchanged.

Parameters: ciffilestr: str, bytes

filename of the CIF file or string representation of the CIF file

planeDistance (*hkl)

determines the lattice plane spacing for the planes specified by (hkl)

Parameters: h, k, I: list, tuple or floats

Miller indices of the lattice planes given either as list, tuple or seperate arguments

Returns: float

the lattice plane spacing

Examples

```
>>> xu.materials.Si.planeDistance(0, 0, 4)
1.35776000000001
```

```
or
```

```
>>> xu.materials.Si.planeDistance((1, 1, 1))
3.1356124059796255
```

show_unitcell (fig=None, subplot=111, scale=0.6, complexity=11, linewidth=1.5, mode='matplotlib') visualization of the unit cell using either matplotlibs basic 3D functionality (expect rendering inaccuracies!) or the mayavi mlab package (accurate rendering -> recommended!)

Note

For more flexible visualization consider using the CIF-export feature and use a proper crystal structure viewer.

Parameters: fig: matplotlib Figure, Mayavi Scene, or None, optional

subplot: int or list, optional

subplot to use for the visualization when using matplotlib. This argument of fowarded to the first argument of matplotlibs *add_subplot* function

scale: float, optional

scale the size of the atoms by this additional factor. By default the size of the atoms corresponds to 60% of their atomic radius.

complexity: int, optional

number of steps to approximate the atoms as spheres. Higher values make spheres more accurate, but cause slower plotting.

linewidth: float, optional

line thickness of the unit cell outline

mode: str, optional

defines the plot backend used, can be 'matplotlib' (default) or 'mayavi'.

Returns: figure object of either matplotlib or Mayavi

toCIF (ciffilename)

Export the Crystal to a CIF file.

Parameters: ciffilename: str

filename of the CIF file

class xrayutilities.materials.material.CubicAlloy (matA, matB, x)

Bases: xrayutilities.materials.material.Alloy

ContentBasym (q_inp, q_perp, hkl, sur)

function that determines the content of B in the alloy from the reciprocal space position of an asymmetric peak.

Parameters: q_inp : float

inplane peak position of reflection hkl of the alloy in reciprocal space

q_perp : float

perpendicular peak position of the reflection hkl of the alloy in reciprocal space

hkl: list

Miller indices of the measured asymmetric reflection

sur : list

Miller indices of the surface (determines the perpendicular direction)

Returns: content: float

content of B in the alloy determined from the input variables

list

[a_inplane a_perp, a_bulk_perp(x), eps_inplane, eps_perp]; lattice parameters calculated from the reciprocal space positions as well as the strain (eps) of the layer

ContentBsym (q_perp, hkl, inpr, asub, relax)

function that determines the content of B in the alloy from the reciprocal space position of a symetric peak. As an additional input the substrates lattice parameter and the degree of relaxation must be given

Parameters: q_perp: float

perpendicular peak position of the reflection hkl of the alloy in reciprocal space

hkl: list

Miller indices of the measured symmetric reflection (also defines the surface normal

inpr: list

Miller indices of a Bragg peak defining the inplane reference direction

asub: float

substrate lattice parameter

relax: float

degree of relaxation (needed to obtain the content from symmetric reciprocal space

position)

Returns: content: float

the content of B in the alloy determined from the input variables

xrayutilities.materials.material.CubicElasticTensor (c11, c12, c44)

Assemble the 6x6 matrix of elastic constants for a cubic material from the three independent components of a cubic crystal

Parameters: c11, c12, c44 : float

independent components of the elastic tensor of cubic materials

Returns: cij: ndarray

6x6 matrix with elastic constants

xrayutilities.materials.material.HexagonalElasticTensor (c11, c12, c13, c33, c44)

Assemble the 6x6 matrix of elastic constants for a hexagonal material from the five independent components of a hexagonal crystal

Parameters: c11, c12, c13, c33, c44 : float

independent components of the elastic tensor of a hexagonal material

Returns: cij: ndarray

6x6 matrix with elastic constants

class xrayutilities.materials.material.Material (name, cij=None)

Bases: abc.ABC

base class for all Materials. common properties of amorphous and crystalline materials are described by this class from which Amorphous and Crystal are derived from.

absorption_length (en='config')

wavelength dependent x-ray absorption length defined as mu = lambda/(2*pi*2*beta) with lambda and beta as the x-ray wavelength and complex part of the refractive index respectively.

Parameters: en: float or str, optional

energy of the x-rays in eV

Returns: float

the absorption length in um

chi0 (en='config')

calculates the complex chi_0 values often needed in simulations. They are closely related to delta and beta ($n = 1 + chi r0/2 + i^*chi i0/2 vs. n = 1 - delta + i^*beta$)

critical_angle (en='config', deg=True)

calculate critical angle for total external reflection

Parameters: en: float or str. optional

energy of the x-rays in eV, if omitted the value from the xrayutilities configuration is

used

deg: bool, optional

return angle in degree if True otherwise radians (default:True)

Returns: float

Angle of total external reflection

abstract delta (en='config')

abstract method which every implementation of a Material has to override

property density

abstract ibeta (en='config')

abstract method which every implementation of a Material has to override

idx_refraction (en='config')

function to calculate the complex index of refraction of a material in the x-ray range

Parameters: en: energy of the x-rays, if omitted the value from the

xrayutilities configuration is used

Returns: n (complex)

property lam

property mu

property nu

xrayutilities.materials.material.**PseudomorphicMaterial** (sub, layer, relaxation=0, trans=None)
This function returns a material whos lattice is pseudomorphic on a particular substrate material. The two materials must have similar unit cell definitions for the algorithm to work correctly, i.e. it does not work for combiniations of materials with different lattice symmetry. It is also crucial that the layer object includes values for the elastic tensor.

Parameters: sub: Crystal

substrate material

layer: Crystal

bulk material of the layer, including its elasticity tensor

relaxation: float, optional

degree of relaxation 0: pseudomorphic, 1: relaxed (default: 0)

trans: Tranform

Transformation which transforms lattice directions into a surface orientated coordinate frame (x, y inplane, z out of plane). If None a (001) surface geometry of a cubic material

is assumed.

Returns: An instance of Crystal holding the new pseudomorphically

strained material.

Raises: InputError

If the layer material has no elastic parameters

xrayutilities.materials.material.WZTensorFromCub (c11ZB, c12ZB, c44ZB)

Determines the hexagonal elastic tensor from the values of the cubic elastic tensor under the assumptions presented in Phys. Rev. B 6, 4546 (1972), which are valid for the WZ <-> ZB polymorphs.

Parameters: c11, c12, c44: float

independent components of the elastic tensor of cubic materials

Returns: cij: ndarray

6x6 matrix with elastic constants

Implementation according to a patch submitted by Julian Stangl

xrayutilities.materials.material.index_map_ij2ijkl(ij)
xrayutilities.materials.material.index_map_ijkl2ij(i,j)

xrayutilities.materials.plot module

xrayutilities.materials.plot.show_reciprocal_space_plane (mat, exp, ttmax=None, maxqout=0.01, scalef=100, ax=None, color=None, show_Laue=True, show_legend=True, projection='perpendicular', label=None) show a plot of the coplanar diffraction plane with peak positions for the respective material. the size of the spots is scaled with the strength of the structure factor

Parameters: mat: Crystal

instance of Crystal for structure factor calculations

exp: Experiment

instance of Experiment (needs to be HXRD, or FourC for onclick action to work correctly). defines the inplane and out of plane direction as well as the sample azimuth

ttmax: float, optional

maximal 2Theta angle to consider, by default 180deg

maxqout: float, optional

maximal out of plane q for plotted Bragg peaks as fraction of exp.k0

scalef: float, or callable, optional

scale factor or function for the marker size. If this is a function it should take only one float argument and return another float which is used as 's' parameter in matplotlib.pyplot.scatter

ax: matplotlib.Axes, optional

matplotlib Axes to use for the plot, useful if multiple materials should be plotted in one plot

color: matplotlib color, optional

show_Laue: bool, optional

flag to indicate if the Laue zones should be indicated

show_legend: bool, optional

flag to indiate if a legend should be shown

projection: 'perpendicular', 'polar', optional

type of projection for Bragg peaks which do not fall into the diffraction plane. 'perpendicular' (default) uses only the inplane component in the scattering plane, whereas 'polar' uses the vectorial absolute value of the two inplane components. See also the 'maxgout' option.

label: None or str, optional

label to be used for the legend. If 'None' the name of the material will be used.

Returns: Axes, plot_handle

xrayutilities.materials.predefined_materials module

```
class xrayutilities.materials.predefined_materials.AlGaAs (X)
```

Bases: xrayutilities.materials.material.CubicAlloy

class xrayutilities.materials.predefined_materials.SiGe (x)

Bases: xrayutilities.materials.material.CubicAlloy

static lattice_const_AB (latA, latB, x, **kwargs)

method to calculate the lattice parameter of the SiGe alloy with composition Si_{1-x}Ge_x

xrayutilities.materials.spacegrouplattice module

module handling crystal lattice structures. A SGLattice consists of a space group number and the position of atoms specified as Wyckoff positions along with their parameters. Depending on the space group symmetry only certain parameters of the resulting instance will be settable! A cubic lattice for example allows only to set its 'a' lattice parameter but none of the other unit cell shape parameters.

class xrayutilities.materials.spacegrouplattice.SGLattice (sgrp, *args, **kwargs)

Bases: object

lattice object created from the space group number and corresponding unit cell parameters. atoms in the unit cell are specified by their Wyckoff position and their free parameters.

ApplyStrain (eps)

Applies a certain strain on a lattice. The result is a change in the base vectors. The full strain matrix (3x3) needs to be given.

Note

Here you specify the strain and not the stress -> NO elastic response of the material will be considered!

Note

Although the symmetry of the crystal can be lowered by this operation the spacegroup remains unchanged! The 'free_parameters' attribute is, however, updated to mimic the possible reduction of the symmetry.

Parameters: eps: array-like

a 3x3 matrix with all strain components

GetHKL (*args)

determine the Miller indices of the given reciprocal lattice points

GetPoint (*args)

determine lattice points with indices given in the argument

Examples

```
>>> xu.materials.Si.lattice.GetPoint(0, 0, 4) array([ 0. , 0. , 21.72416])
```

```
or
```

```
>>> xu.materials.Si.lattice.GetPoint((1, 1, 1))
array([ 5.43104, 5.43104, 5.43104])
```

GetQ (*args)

determine the reciprocal lattice points with indices given in the argument

UnitCellVolume ()

function to calculate the unit cell volume of a lattice (angstrom^3)

property a

property alpha

property b

base ()

generator of atomic position within the unit cell.

property beta

property c

convert_to_P1()

create a P1 equivalent of this SGLattice instance.

Returns: SGLattice

instance with the same properties as the present lattice, however, in the P1 setting.

equivalent_hkls (hkl)

returns a list of equivalent hkl peaks depending on the crystal system

findsym()

method to return the highest symmetry description of the current material. This method does not consider to change the unit cell dimensions but only searches the highest symmetry spacegroup which with the current unit cell setting can be described. It is therefore not an implementation of FINDSYM [1].

Returns: new SGLattice-instance

a new SGLattice instance is returned with the highest available symmetry description. (see restrictions above)

[1] https://stokes.byu.edu/iso/findsym.php

property gamma

get_allowed_hkl (qmax)

return a set of all allowed reflections up to a maximal specified momentum transfer.

Parameters: qmax: float

maximal momentum transfer

Returns: hklset: set

set of allowed hkl reflections

hkl_allowed (hkl, returnequivalents=False)

check if Bragg reflection with Miller indices hkl can exist according to the reflection conditions. If no reflection conditions are available this function returns True for all hkl values!

Parameters: hkl: tuple or list

Miller indices of the reflection to check

returnequivalents: bool, optional

If True all the equivalent Miller indices of hkl are returned in a set as second return

argument.

Returns: allowed: bool

True if reflection can have non-zero structure factor, false otherwise

equivalents: set, optional

set of equivalent Miller indices if returnequivalents is True

property iscentrosymmetric

returns a boolean to determine if the lattice has centrosymmetry.

isequivalent (hkl1, hkl2)

determining if hkl1 and hkl2 are two crystallographical equivalent pairs of Miller indices. Note that this function considers the effect of non-centrosymmetry!

Parameters: hkl1, hkl2: list

Miller indices to be checked for equivalence

Returns: bool

reflection_conditions()

return string of reflection conditions, both general (from space group) and of Wyckoff positions

property symops

return the set of symmetry operations from the general Wyckoff position of the space group.

transform (mat, origin)

Transform the unit cell with the matrix and origin shift given in the parameters. This function returns a new instance of SGLattice which contains the highest possible symmetry description of the transformed unit cell. After the transformation (see [1]) the findsym method is used to create the new SGLattice instance.

Parameters: mat: (3, 3) list, or ndarray, optional

transformation matrix of the unit cell. The matrix definition aims to be consistent with what is used on the Bilbao Crystallographic Server [1]. This only defines the linear part, while the origin shift is given by origin.

origin : (3,) list, or ndarray
origin shift of the unit cell [1].

[1] https://www.cryst.ehu.es/cgi-bin/cryst/programs/nph-doc-trmat

class xrayutilities.materials.spacegrouplattice.symOp (D, t, m=1)

Bases: object

Class descriping a symmetry operation in a crystal. The symmetry operation is characterized by a 3x3 transformation matrix as well as a 3-vector describing a translation. For magnetic symmetry operations also the time reversal symmetry can be specified (not used in xrayutilities)

property D

transformation matrix of the symmetry operation

```
apply (vec, foldback=True)
apply_axial (vec)
apply_rotation (vec)
combine (other)
static foldback (v)
```

classmethod from_xyz (XYZ)

create a SymOp from the xyz notation typically used in CIF files.

Parameters: xyz: str

string describing the symmetry operation (e.g. '-y, -x, z')

property t

translation vector of the symmetry operation

```
xyz (showtimerev=False)
```

return the symmetry operation in xyz notation

class xrayutilities.materials.spacegrouplattice.WyckoffBase (*args, **kwargs)

Bases: list

The WyckoffBase class implements a container for a set of Wyckoff positions that form the base of a crystal lattice. An instance of this class can be treated as a simple container object.

append (atom, pos, occ=1.0, b=0.0) add new Atom to the lattice base

Parameters: atom: Atom

object to be added

pos: tuple or str

Wyckoff position of the atom, along with its parameters. Examples: ('2i', (0.1, 0.2,

0.3)), or '1a'

occ: float, optional

occupancy (default=1.0)

b: float, optional

b-factor of the atom used as exp(-b*q**2/(4*pi)**2) to reduce the intensity of this atom

(only used in case of temp=0 in StructureFactor and chi calculation)

static entry_eq (e1, e2)

compare two entries including all its properties to be equal

Parameters: e1, e2: tuple

tuples with length 4 containing the entries of WyckoffBase which should be compared

index (item)

return the index of the atom (same element, position, and Debye Waller factor). The occupancy is not checked intentionally. If the item is not present a ValueError is raised.

Parameters: item: tuple or list

WyckoffBase entry

Returns: int

static pos_eq (pos1, pos2) compare Wyckoff positions

Parameters: pos1, pos2: tuple

tuples with Wyckoff label and optional parameters

xrayutilities.materials.spacegrouplattice.get_default_sgrp_suf (sgrp_nr)
 determine default space group suffix

xrayutilities.materials.spacegrouplattice.get_possible_sgrp_suf (sgrp_nr)

determine possible space group suffix. Multiple suffixes might be possible for one space group due to different origin choice, unique axis, or choice of the unit cell shape.

Parameters: sgrp_nr: int

space group number

Returns: str or list

either an empty string or a list of possible valid suffix strings

xrayutilities.materials.spacegrouplattice.get_wyckpos (sgrp, atompos) test all Wyckoff positions on every atomic position

Parameters: sgrp: str

space group name

atompos: list

list of atomic positions to identify. All atomic positions are expected to belong to one and

the same Wyckoff position!

Returns: position argument for WyckoffBase.append

xrayutilities.materials.spacegrouplattice.testwp (parint, wp, cifpos, digits=8) test if a Wyckoff position can describe the given position from a CIF file

Parameters: parint : int

telling which Parameters the given Wyckoff position has

wp: str or tuple

expression of the Wyckoff position

cifpos: list, or tuple or array-like

(x, y, z) position of the atom in the CIF file

digits: int

number of digits for which for a comparison of floating point numbers will be rounded to.

By default xu.config.DIGITS is used.

Returns: foundflag: bool

flag to tell if the positions match

pars: array-like or None

parameters associated with the position or None if no parameters are needed

xrayutilities.materials.wyckpos module

class xrayutilities.materials.wyckpos.RangeDict() -> new empty dictionary dict(mapping) -> new dictionary initialized from a mapping object's (key, value) pairs dict(iterable) -> new dictionary initialized as if via: $d = \{\}$ for k, v in iterable: d[k] = v dict(**kwargs) -> new dictionary initialized with the name=value pairs in the keyword argument list. For example: dict(one=1, two=2)

Bases: dict

Module contents

xrayutilities.math package

Submodules

xrayutilities.math.algebra module

module providing analytic algebraic functions not implemented in scipy or any other dependency of xrayutilities. In particular the analytic solution of a quartic equation which is needed for the solution of the dynamic scattering equations.

```
xrayutilities.math.algebra.solve_quartic (a4, a3, a2, a1, a0) analytic solution [1] of the general quartic equation. The solved equation takes the form a \cdot z^4 + a \cdot z^3 + a \cdot z^2 + a \cdot z + a \cdot z
```

Returns: tuple

tuple of the four (complex) solutions of aboves equation.

References

1

http://mathworld.wolfram.com/QuarticEquation.html

xrayutilities.math.fit module

module with a function wrapper to scipy.optimize.leastsq for fitting of a 2D function to a peak or a 1D Gauss fit with the odr package

xrayutilities.math.fit.**fit_peak2d** (x, y, data, start, drange, fit_function, maxfev=2000) fit a two dimensional function to a two dimensional data set e.g. a reciprocal space map

Parameters: x, y : array-like

data coordinates (do NOT need to be regularly spaced)

data: array-like

data set used for fitting (e.g. intensity at the data coords)

start: list

set of starting parameters for the fit used as first parameter of function fit_function

drange: list

limits for the data ranges used in the fitting algorithm, e.g. it is clever to use only a small region around the peak which should be fitted: [xmin, xmax, ymin, ymax]

fit function: callable

function which should be fitted, must be of form accept the parameters fit_function (x, y, *params) -> ndarray

Returns: fitparam: list

fitted parameters

cov: array-like

covariance matrix

xrayutilities.math.fit.gauss_fit (xdata, ydata, iparams=[], maxit=300)

Gauss fit function using odr-pack wrapper in scipy similar to

https://github.com/tiagopereira/python_tips/wiki/Scipy%3A-curve-fitting

Parameters: xdata: array-like

x-coordinates of the data to be fitted

ydata : array-like

y-coordinates of the data which should be fit

iparams: list, optional

initial paramters for the fit, determined automatically if not given

maxit: int, optional

maximal iteration number of the fit

Returns: params: list

the parameters as defined in function Gauss1d(x, *param)

sd_params : list

For every parameter the corresponding errors are returned.

itlim: bool

flag to tell if the iteration limit was reached, should be False

xrayutilities.math.fit.linregress(x, y)

fast linregress to avoid usage of scipy.stats which is slow! NaN values in y are ignored by this function.

Parameters: x, y : array-like

data coordinates and values

Returns: p:tuple

parameters of the linear fit (slope, offset)

rsq: float R^2 value

Examples

```
>>> (k, d), R2 = xu.math.linregress(x, y)
```

xrayutilities.math.fit.multPeakFit (x, data, peakpos, peakwidth, dranges=None, peaktype='Gaussian',
returnerror=False)

function to fit multiple Gaussian/Lorentzian peaks with linear background to a set of data

Parameters: x: array-like

x-coordinate of the data

data: array-like

data array with same length as x

peakpos: list

initial parameters for the peak positions

peakwidth: list

initial values for the peak width

dranges: list of tuples

list of tuples with (min, max) value of the data ranges to use. does not need to have the

same number of entries as peakpos

peaktype : {'Gaussian', 'Lorentzian'}

type of peaks to be used

returnerror: bool

decides if the fit errors of pos, sigma, and amp are returned (default: False)

Returns: pos: list

peak positions derived by the fit

sigma: list

peak width derived by the fit

amp: list

amplitudes of the peaks derived by the fit

background: array-like

background values at positions x

if returnerror == True:

sd_pos: list

standard error of peak positions as returned by scipy.odr.Output

sd_sigma : list

standard error of the peak width

sd_amp: list

standard error of the peak amplitude

xrayutilities.math.fit.multPeakPlot (x, fpos, fwidth, famp, background, dranges=None,
peaktype='Gaussian', fig='xu_plot', ax=None, fact=1.0)

function to plot multiple Gaussian/Lorentz peaks with background values given by an array

Parameters: x: array-like

x-coordinate of the data

fpos: list

positions of the peaks

fwidth: list

width of the peaks

famp: list

amplitudes of the peaks

background: array-like

background values, same shape as x

dranges: list of tuples

list of (min, max) values of the data ranges to use. does not need to have the same number of entries as fpos

peaktype : {'Gaussian', 'Lorentzian'}

type of peaks to be used

fig: int, str, or None

matplotlib figure number or name

ax: matplotlib.Axes

matplotlib axes as alternative to the figure name

fact: float

factor to use as multiplicator in the plot

xrayutilities.math.fit.peak_fit (xdata, ydata, iparams=[], peaktype='Gauss', maxit=300,

background='constant', plot=False, func_out=False, debug=False)

fit function using odr-pack wrapper in scipy similar to https://github.com/tiagopereira/python_tips/wiki/Scipy%3A-curve-fitting for Gauss, Lorentz or Pseudovoigt-functions

Parameters: xdata: array_like

x-coordinates of the data to be fitted

ydata: array_like

y-coordinates of the data which should be fit

iparams: list, optional

initial paramters, determined automatically if not specified

peaktype : {'Gauss', 'Lorentz', 'PseudoVoigt', 'PseudoVoigtAsym', 'PseudoVoigtAsym2'},

optional

type of peak to fit

maxit: int, optional

maximal iteration number of the fit

background: {'constant', 'linear'}, optional

type of background function

plot: bool or str, optional

flag to ask for a plot to visually judge the fit. If plot is a string it will be used as figure name, which makes reusing the figures easier.

func out: bool, optional

returns the fitted function, which takes the independent variables as only argument (f(x))

Returns: params: list

the parameters as defined in function *Gauss1d/Lorentz1d/PseudoVoigt1d/PseudoVoigt1dasym*. In the case of linear background one more parameter is included!

sd_params: list

For every parameter the corresponding errors are returned.

itlim: bool

flag to tell if the iteration limit was reached, should be False

fitfunc: function, optional

the function used in the fit can be returned (see func_out).

xrayutilities.math.functions module

module with several common function needed in xray data analysis

xrayutilities.math.functions.Debye1(X)

function to calculate the first Debye function [1] as needed for the calculation of the thermal Debye-Waller-factor by numerical integration

 $D_1(x) = (1/x) \int_0^x t/(\exp(t)-1) dt$

Parameters: x: float

argument of the Debye function

Returns: float

D1(x) float value of the Debye function

References

1 http://en.wikipedia.org/wiki/Debye_function

xrayutilities.math.functions.**Gauss1d** (x, *p) function to calculate a general one dimensional Gaussian

Parameters: x: array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the Gaussian [XCEN, SIGMA, AMP, BACKGROUND] for information: SIGMA = FWHM / (2*sqrt(2*log(2)))

Returns: arrav-like

the value of the Gaussian described by the parameters p at position x

Examples

Calling with a list of parameters needs a call looking as shown below (note the '*') or explicit listing of the parameters

```
>>> Gauss1d(x,*p)
```

```
>>> Gauss1d(numpy.linspace(0, 10, 100), 5, 1, 1e3, 0)
```

```
xrayutilities.math.functions.GaussldArea (*p)
```

function to calculate the area of a Gauss function with neglected background

Parameters: p: list

list of parameters of the Gauss-function [XCEN, SIGMA, AMP, BACKGROUND]

Returns: float

the area of the Gaussian described by the parameters p

```
xrayutilities.math.functions.Gauss1d_der_p(x, *p)
```

function to calculate the derivative of a Gaussian with respect the parameters p for parameter description see Gauss1d

```
xrayutilities.math.functions.Gauss1d\_der\_x(x, *p)
```

function to calculate the derivative of a Gaussian with respect to x for parameter description see Gauss1d

xrayutilities.math.functions.Gauss2d(x, y, *p)

function to calculate a general two dimensional Gaussian

Parameters: x, y: array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the Gauss-function [XCEN, YCEN, SIGMAX, SIGMAY, AMP, BACKGROUND, ANGLE]; SIGMA = FWHM / (2*sqrt(2*log(2))); ANGLE = rotation of the

X, Y direction of the Gaussian in radians

Returns: array-like

the value of the Gaussian described by the parameters p at position (x, y)

xrayutilities.math.functions.Gauss2dArea(*p)

function to calculate the area of a 2D Gauss function with neglected background

Parameters: p: list

list of parameters of the Gauss-function [XCEN, YCEN, SIGMAX, SIGMAY, AMP,

ANGLE, BACKGROUND]

Returns: float

the area of the Gaussian described by the parameters p

xrayutilities.math.functions.Gauss3d(x, y, z, *p)

function to calculate a general three dimensional Gaussian

Parameters: x, y, z : array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the Gauss-function [XCEN, YCEN, ZCEN, SIGMAX, SIGMAY,

SIGMAZ, AMP, BACKGROUND];

SIGMA = FWHM / (2*sqrt(2*log(2)))

Returns: array-like

the value of the Gaussian described by the parameters p at positions (x, y, z)

xrayutilities.math.functions.Lorentzld (x, *p)

function to calculate a general one dimensional Lorentzian

Parameters: x : array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the Lorentz-function [XCEN, FWHM, AMP, BACKGROUND]

Returns: array-like

the value of the Lorentian described by the parameters p at position (x, y)

xrayutilities.math.functions.LorentzldArea (*p)

function to calculate the area of a Lorentz function with neglected background

Parameters: p: list

list of parameters of the Lorentz-function [XCEN, FWHM, AMP, BACKGROUND]

Returns: float

the area of the Lorentzian described by the parameters *p*

xrayutilities.math.functions.Lorentz1d_der_p(x, *p)

function to calculate the derivative of a Gaussian with respect the parameters p for parameter description see Lorentz1d

xrayutilities.math.functions.Lorentz1d_der_x(x, *p)

function to calculate the derivative of a Gaussian with respect to x for parameter description see Lorentz1d

xrayutilities.math.functions.Lorentz2d(x, y, *p)

function to calculate a general two dimensional Lorentzian

Parameters: x, y: array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the Lorentz-function [XCEN, YCEN, FWHMX, FWHMY, AMP, BACKGROUND, ANGLE]; ANGLE = rotation of the X, Y direction of the Lorentzian in

radians

Returns: array-like

the value of the Lorentian described by the parameters p at position (x, y)

xrayutilities.math.functions.NormGauss1d(x, *p)

function to calculate a normalized one dimensional Gaussian

Parameters: x: array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the Gaussian [XCEN, SIGMA]; for information: SIGMA = FWHM /

(2*sqrt(2*log(2)))

Returns: array-like

the value of the normalized Gaussian described by the parameters p at position x

xrayutilities.math.functions.NormLorentz1d(x, *p)

function to calculate a normalized one dimensional Lorentzian

Parameters: x: array-like

coordinate(s) where the function should be evaluated

p : list

list of parameters of the Lorentzian [XCEN, FWHM]

Returns: array-like

the value of the normalized Lorentzian described by the parameters p at position x

xrayutilities.math.functions.PseudoVoigt1d(x, *p)

function to calculate a pseudo Voigt function as linear combination of a Gauss and Lorentz peak

Parameters: x: array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the pseudo Voigt-function [XCEN, FWHM, AMP, BACKGROUND,

ETA]; ETA: 0 ...1 0 means pure Gauss and 1 means pure Lorentz

Returns: array-like

the value of the PseudoVoigt described by the parameters p at position x

xrayutilities.math.functions.PseudoVoigt1dArea (*p)

function to calculate the area of a pseudo Voigt function with neglected background

Parameters: p: list

list of parameters of the Lorentz-function [XCEN, FWHM, AMP, BACKGROUND, ETA];

ETA: 0 ... 1 0 means pure Gauss and 1 means pure Lorentz

Returns: float

the area of the PseudoVoigt described by the parameters p

xrayutilities.math.functions.**PseudoVoigt1d_der_p**(X, *p)

function to calculate the derivative of a PseudoVoigt with respect the parameters p for parameter description see PseudoVoigt1d

xrayutilities.math.functions.PseudoVoigt1d der x (x, *p)

function to calculate the derivative of a PseudoVoigt with respect to x

for parameter description see PseudoVoigt1d

xrayutilities.math.functions.**PseudoVoigt1dasym**(x, *p)

function to calculate an asymmetric pseudo Voigt function as linear combination of asymmetric Gauss and Lorentz peak

Parameters: x: array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the pseudo Voigt-function [XCEN, FWHMLEFT, FWHMRIGHT, AMP, BACKGROUND, ETA]; ETA: 0 ...1 0 means pure Gauss and 1 means pure

Lorentz

Returns: array-like

the value of the PseudoVoigt described by the parameters p at position x

xrayutilities.math.functions.PseudoVoigt1dasym2(x, *p)

function to calculate an asymmetric pseudo Voigt function as linear combination of asymmetric Gauss and Lorentz peak

Parameters: x : naddray

coordinate(s) where the function should be evaluated

p: list

list of parameters of the pseudo Voigt-function [XCEN, FWHMLEFT, FWHMRIGHT, AMP, BACKGROUND, ETALEFT, ETARIGHT]; ETA: 0 ...1 0 means pure Gauss and 1 means pure Lorentz

Returns: array-like

the value of the PseudoVoigt described by the parameters p at position x

xrayutilities.math.functions.**PseudoVoigt2d**(x, y, *p)

function to calculate a pseudo Voigt function as linear combination of a Gauss and Lorentz peak in two dimensions

Parameters: x, y : array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the pseudo Voigt-function [XCEN, YCEN, FWHMX, FWHMY, AMP, BACKGROUND, ANGLE, ETA]; ETA: 0 ...1 0 means pure Gauss and 1 means pure

Lorentz

Returns: array-like

the value of the PseudoVoigt described by the parameters p at position (x, y)

xrayutilities.math.functions.**TwoGauss2d**(x, y, *p)

function to calculate two general two dimensional Gaussians

Parameters: x, y : array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the Gauss-function [XCEN1, YCEN1, SIGMAX1, SIGMAY1, AMP1, ANGLE1, XCEN2, YCEN2, SIGMAX2, SIGMAY2, AMP2, ANGLE2, BACKGROUND]; SIGMA = FWHM / (2*sqrt(2*log(2))) ANGLE = rotation of the X, Y direction of the

Gaussian in radians

Returns: array-like

the value of the Gaussian described by the parameters p at position (x, y)

xrayutilities.math.functions.heaviside(x)

Heaviside step function for numpy arrays

Parameters: x: scalar or array-like

argument of the step function

Returns: int or array-like

Heaviside step function evaluated for all values of x with datatype integer

xrayutilities.math.functions.kill_spike (data, threshold=2.0, offset=None)

function to smooth **single** data points which differ from the average of the neighboring data points by more than the threshold factor or more than the offset value. Such spikes will be replaced by the mean value of the next neighbors.

Warning

Use this function carefully not to manipulate your data!

Parameters: data: array-like

1d numpy array with experimental data

threshold: float or None

threshold factor to identify outlier data points. If None it will be ignored.

offset: None or float

offset value to identify outlier data points. If None it will be ignored.

Returns: array-like

1d data-array with spikes removed

xrayutilities.math.functions.multPeak1d(x, *args)

function to calculate the sum of multiple peaks in 1D. the peaks can be of different type and a background function (polynom) can also be included.

Parameters: x: array-like

coordinate where the function should be evaluated

args : list

list of peak/function types and parameters for every function type two arguments need to be given first the type of function as string with possible values 'g': Gaussian, 'l': Lorentzian, 'v': PseudoVoigt, 'a': asym. PseudoVoigt, 'p': polynom the second type of arguments is the tuple/list of parameters of the respective function. See documentation of math.Gauss1d, math.Lorentz1d, math.PseudoVoigt1d, math.PseudoVoigt1dasym, and numpy.polyval for details of the different function types.

Returns: array-like

value of the sum of functions at position x

xrayutilities.math.functions.multPeak2d(x, y, *args)

function to calculate the sum of multiple peaks in 2D. the peaks can be of different type and a background function (polynom) can also be included.

Parameters: x, y : array-like

coordinates where the function should be evaluated

args: list

list of peak/function types and parameters for every function type two arguments need to be given first the type of function as string with possible values 'g': Gaussian, 'l': Lorentzian, 'v': PseudoVoigt, 'c': constant the second type of arguments is the tuple/list of parameters of the respective function. See documentation of math.Gauss2d, math.Lorentz2d, math.PseudoVoigt2d for details of the different function types. The constant accepts a single float which will be added to the data

Returns: array-like

value of the sum of functions at position (x, y)

xrayutilities.math.functions.smooth(x, n)

function to smooth an array of data by averaging N adjacent data points

Parameters: x: array-like

1D data array

n:int

number of data points to average

Returns: xsmooth: array-like

smoothed array with same length as x

xrayutilities.math.misc module

xrayutilities.math.misc.center_of_mass (pos, data, background='none', full_output=False)
function to determine the center of mass of an array

Parameters: pos: array-like

position of the data points

data : array-like data values

background : {'none', 'constant', 'linear'}

type of background, either 'none', 'constant' or 'linear'

full_output : bool

return background cleaned data and background-parameters

Returns: float

center of mass position

xrayutilities.math.misc.fwhm_exp (pos, data)

function to determine the full width at half maximum value of experimental data. Please check the obtained value visually (noise influences the result)

Parameters: pos: array-like

position of the data points

data : array-like data values

Returns: float

fwhm value

xrayutilities.math.misc.gcd (lst)

greatest common divisor function using library functions

Parameters: Ist: array-like

array of integer values for which the greatest common divisor should be determined

Returns: gcd: int

xrayutilities.math.transforms module

xrayutilities.math.transforms.ArbRotation (axis, alpha, deg=True)

Returns a transform that represents a rotation around an arbitrary axis by the angle alpha. positive rotation is anti-clockwise when looking from positive end of axis vector

Parameters: axis: list or array-like

rotation axis

alpha: float

rotation angle in degree (deg=True) or in rad (deg=False)

deg: bool

determines the input format of ang (default: True)

Returns: **Transform**

class xrayutilities.math.transforms.AxisToZ (newzaxis)

Bases: xrayutilities.math.transforms.CoordinateTransform

Creates a coordinate transformation to move a certain axis to the z-axis. The rotation is done along the great circle. The x-axis of the new coordinate frame is created to be normal to the new and original z-axis. The new y-axis is create in order to obtain a right handed coordinate system.

class xrayutilities.math.transforms.AxisToZ_keepXY (newzaxis)

Bases: xrayutilities.math.transforms.CoordinateTransform

Creates a coordinate transformation to move a certain axis to the z-axis. The rotation is done along the great circle. The x-axis/y-axis of the new coordinate frame is created to be similar to the old x and y directions. This variant of AxisToZ assumes that the new Z-axis has its main component along the Z-direction

class xrayutilities.math.transforms.CoordinateTransform (V1, V2, V3)

Bases: xrayutilities.math.transforms.Transform

Create a Transformation object which transforms a point into a new coordinate frame. The new frame is determined by the three vectors v1/norm(v1), v2/norm(v2) and v3/norm(v3), which need to be orthogonal!

class xrayutilities.math.transforms.Transform (matrix)

Bases: object

property imatrix

inverse (args, rank=1)

performs inverse transformation a vector, matrix or tensor of rank 4

Parameters: args: list or array-like

object to transform, list or numpy array of shape (..., n) (..., n, n), (..., n, n, n, n, n) where n is the size of the transformation matrix.

rank: int

rank of the supplied object. allowed values are 1, 2, and 4

xrayutilities.math.transforms.XRotation (alpha, deg=True)

Returns a transform that represents a rotation about the x-axis by an angle alpha. If deg=True the angle is assumed to be in degree, otherwise the function expects radiants.

xrayutilities.math.transforms.YRotation (alpha, deg=True)

Returns a transform that represents a rotation about the y-axis by an angle alpha. If deg=True the angle is assumed to be in degree, otherwise the function expects radiants.

xrayutilities.math.transforms.ZRotation (alpha, deg=True)

Returns a transform that represents a rotation about the z-axis by an angle alpha. If deg=True the angle is assumed to be in degree, otherwise the function expects radiants.

xrayutilities.math.transforms.mycross (vec, mat)

function implements the cross-product of a vector with each column of a matrix

xrayutilities.math.transforms.rotarb (vec, axis, ang, deg=True)

function implements the rotation around an arbitrary axis by an angle ang positive rotation is anti-clockwise when looking from positive end of axis vector

Parameters: vec: list or array-like

vector to rotate

axis: list or array-like
rotation axis

ang: float

rotation angle in degree (deg=True) or in rad (deg=False)

deg: bool

determines the input format of ang (default: True)

Returns: rotvec: rotated vector as numpy.array

Examples

```
>>> rotarb([1, 0, 0],[0, 0, 1], 90)
array([ 6.12323400e-17,  1.00000000e+00,  0.00000000e+00])
```

xrayutilities.math.transforms.tensorprod (vec1, vec2) function implements an elementwise multiplication of two vectors

xrayutilities.math.vector module

module with vector operations for vectors of size 3, since for so short vectors numpy does not give the best performance explicit implementation of the equations is performed together with error checking to ensure vectors of length 3.

```
xrayutilities.math.vector.VecAngle ((v1.v2)/(norm(v1)*norm(v2))) alpha = acos((v1.v2)/(norm(v1)*norm(v2)))
```

Parameters: v1, v2: list or array-like

input vector(s), either one vector or an array of vectors with shape (n, 3)

deg: bool, optional

True: return result in degree, False: in radiants (default: False)

Returns: float or ndarray

the angle included by the two vectors v1 and v2, either a single float or an array with shape (n,)

xrayutilities.math.vector.VecCross (v1, v2, out=None)

Calculate the vector cross product.

Parameters: v1, v2 : list or array-like

input vector(s), either one vector or an array of vectors with shape (n, 3)

out: list or array-like, optional

output vector

Returns: ndarray

cross product either of shape (3,) or (n, 3)

xrayutilities.math.vector.VecDot (v1, v2)

Calculate the vector dot product.

Parameters: v1, v2 : list or array-like

input vector(s), either one vector or an array of vectors with shape (n, 3)

Returns: float or ndarray

innter product of the vectors, either a single float or (n,)

xrayutilities.math.vector.VecNorm(V)

Calculate the norm of a vector.

Parameters: v: list or array-like

input vector(s), either one vector or an array of vectors with shape (n, 3)

Returns: float or ndarray

vector norm, either a single float or shape (n,)

xrayutilities.math.vector.VecUnit (v)

Calculate the unit vector of v.

Parameters: v: list or array-like

input vector(s), either one vector or an array of vectors with shape (n, 3)

Returns: ndarray

unit vector of v, either shape (3,) or (n, 3)

xrayutilities.math.vector.distance (x, y, z, point, vec)

calculate the distance between the point (x, y, z) and the line defined by the point and vector vec

Parameters: x: float or ndarray

x coordinate(s) of the point(s)

y: float or ndarray

y coordinate(s) of the point(s)

z: float or ndarray

z coordinate(s) of the point(s)

point: tuple, list or ndarray

3D point on the line to which the distance should be calculated

vec: tuple, list or ndarray

3D vector defining the propergation direction of the line

xrayutilities.math.vector.getSyntax (Vec)

returns vector direction in the syntax 'x+' 'z-' or equivalents therefore works only for principle vectors of the coordinate system like e.g. [1, 0, 0] or [0, 2, 0]

Parameters: vec: list or array-like

vector of length 3

Returns: str

vector string following the synthax [xyz][+-]

xrayutilities.math.vector.getVector (string)

returns unit vector along a rotation axis given in the syntax 'x+' 'z-' or equivalents

Parameters: string: str

vector string following the synthax [xyz][+-]

Returns: ndarray

vector along the given direction

Module contents

xrayutilities.simpack package

Submodules

xrayutilities.simpack.darwin_theory module

class xrayutilities.simpack.darwin_theory.DarwinModel (qz, qx=0, qy=0, **kwargs)
Bases: xrayutilities.simpack.models.LayerModel

model class inmplementing the basics of the Darwin theory for layers materials. This class is not fully functional and should be used to derive working models for particular material systems.

To make the class functional the user needs to implement the init_structurefactors() and _calc_mono() methods

init structurefactors ()

calculates the needed atomic structure factors

ncalls = 0

simulate (MI)

main simulation function for the Darwin model. will calculate the reflected intensity

Parameters: ml: iterable

monolayer sequence of the sample. This should be created with the function

make_monolayer(). see its documentation for details

class xrayutilities.simpack.darwin_theory.DarwinModelAlGaAs001 (qz, qx=0, qy=0, **kwargs)

Bases: xrayutilities.simpack.darwin_theory.DarwinModelAlloy

Darwin theory of diffraction for Al_x Ga_{1-x} As layers. The model is based on separation of the sample structure into building blocks of atomic planes from which a multibeam dynamical model is calculated.

Alas = <xrayutilities.materials.material.Crystal object>

GaAs = <xrayutilities.materials.material.Crystal object>

aGaAs = 5.65325

classmethod abulk (x)

calculate the bulk (relaxed) lattice parameter of the Al_{x}Ga_{1-x}As alloy

asub = 5.65325

eAl = AI (13)

eAs = As (33)

eGa = Ga(31)

classmethod get_dperp_apar (x, apar, r=1)

calculate inplane lattice parameter and the out of plane lattice plane spacing (of the atomic planes!) from composition and relaxation

Parameters: x: float

chemical composition parameter

apar: float

inplane lattice parameter of the material below the current layer (onto which the present layer is strained to). This value also served as a reference for the relaxation parameter.

parameter

r : float

relaxation parameter. 1=relaxed, 0=pseudomorphic

Returns: dperp: float

perpendicular d-spacing

apar: float

inplane lattice parameter

init_structurefactors (temp=300)

calculates the needed atomic structure factors

Parameters: temp: float, optional

temperature used for the Debye model

static poisson_ratio(x)

calculate the Poisson ratio of the alloy

```
re = 2.8179403262e-05
```

class xrayutilities.simpack.darwin_theory.DarwinModelAlloy (qz, qx=0, qy=0, **kwargs)

Bases: xrayutilities.simpack.darwin_theory.DarwinModel, abc.ABC

extension of the DarwinModel for an binary alloy system were one parameter is used to determine the chemical composition

To make the class functional the user needs to implement the get_dperp_apar() method and define the substrate lattice parameter (asub). See the DarwinModelSiGe001 class for an implementation example.

abstract get_dperp_apar (x, apar, r=1)

calculate inplane lattice parameter and the out of plane lattice plane spacing (of the atomic planes!) from composition and relaxation.

Parameters: x: float

chemical composition parameter

apar: float

inplane lattice parameter of the material below the current layer (onto which the present layer is strained to). This value also served as a reference for the relaxation parameter.

r: float

relaxation parameter. 1=relaxed, 0=pseudomorphic

Returns: dperp: float

apar: float

make_monolayers (S)

create monolayer sequence from layer list

Parameters: s: list

layer model. list of layer dictionaries including possibility to form superlattices. As an example 5 repetitions of a Si(10nm)/Ge(15nm) superlattice on Si would like like:

the dictionaries must contain 't': thickness in A, 'x': chemical composition, and either 'r': relaxation or 'ai': inplane lattice parameter. Future implementations for asymmetric peaks might include layer type 'l' (not yet implemented). Already now any additional property in the dictionary will be handed on to the returned monolayer list.

asub: float

inplane lattice parameter of the substrate

Returns: list

monolayer list in a format understood by the simulate and xGe profile methods

prop profile (ml, prop)

calculate the profile of chemical composition or inplane lattice spacing from a monolayer list. One value for each monolayer in the sample is returned.

Parameters: ml: list

monolayer list created by make_monolayer()

prop: str

name of the property which should be evaluated. Use 'x' for the chemical composition

and 'ai' for the inplane lattice parameter.

Returns: zm: ndarray

z-position, z-0 is the surface

propx : ndarray

value of the property prop for every monolayer

class xrayutilities.simpack.darwin_theory.DarwinModelGaInAs001 (qz, qx=0, qy=0, **kwargs)

Bases: xrayutilities.simpack.darwin_theory.DarwinModelAlloy

Darwin theory of diffraction for Ga_{1-x} In_x As layers. The model is based on separation of the sample structure into building blocks of atomic planes from which a multibeam dynamical model is calculated.

GaAs = <xrayutilities.materials.material.Crystal object>

InAs = <xrayutilities.materials.material.Crystal object>

aGaAs = 5.65325

classmethod abulk (x)

calculate the bulk (relaxed) lattice parameter of the Ga_{1-x}ln_{x}As alloy

asub = 5.65325

eAs = As (33)

eGa = Ga (31)

eIn = ln (49)

classmethod get_dperp_apar (x, apar, r=1)

calculate inplane lattice parameter and the out of plane lattice plane spacing (of the atomic planes!) from composition and relaxation

Parameters: x: float

chemical composition parameter

apar: float

inplane lattice parameter of the material below the current layer (onto which the present layer is strained to). This value also served as a reference for the relaxation parameter.

r: float

relaxation parameter. 1=relaxed, 0=pseudomorphic

Returns: dperp: float

perpendicular d-spacing

apar: float

inplane lattice parameter

init_structurefactors (temp=300)

calculates the needed atomic structure factors

Parameters: temp: float, optional

temperature used for the Debye model

static poisson_ratio (x)

calculate the Poisson ratio of the alloy

```
re = 2.8179403262e-05
```

class xrayutilities.simpack.darwin_theory.DarwinModelSiGe001 (qz, qx=0, qy=0, **kwargs)

Bases: xrayutilities.simpack.darwin_theory.DarwinModelAlloy

model class implementing the Darwin theory of diffraction for SiGe layers. The model is based on separation of the sample structure into building blocks of atomic planes from which a multibeam dynamical model is calculated.

Ge = <xrayutilities.materials.material.Crystal object>

Si = <xrayutilities.materials.material.Crystal object>

aSi = 5.43104

classmethod abulk (x)

calculate the bulk (relaxed) lattice parameter of the alloy

asub = 5.43104

eGe = Ge (32)

esi = Si(14)

classmethod get_dperp_apar (x, apar, r=1)

calculate inplane lattice parameter and the out of plane lattice plane spacing (of the atomic planes!) from composition and relaxation

Parameters: x: float

chemical composition parameter

apar: float

inplane lattice parameter of the material below the current layer (onto which the present layer is strained to). This value also served as a reference for the relaxation parameter.

r: float, optional

relaxation parameter. 1=relaxed, 0=pseudomorphic

Returns: dperp: float

perpendicular d-spacing

apar: float

inplane lattice parameter

init_structurefactors (temp=300)

calculates the needed atomic structure factors

Parameters: temp: float, optional

temperature used for the Debye model

static poisson_ratio (x)

calculate the Poisson ratio of the alloy

re = 2.8179403262e-05

xrayutilities.simpack.darwin_theory.**GradedBuffer** (xfrom, xto, nsteps, thickness, relaxation=1) create a multistep graded composition buffer.

Parameters: xfrom: float

begin of the composition gradient

xto: float

end of the composition gradient

nsteps: int

number of steps of the gradient

thickness: float

total thickness of the Buffer in A

relaxation: float

relaxation of the buffer

Returns: list

layer list needed for the Darwin model simulation

xrayutilities.simpack.darwin_theory.getfirst (iterable, key)

helper function to obtain the first item in a nested iterable

xrayutilities.simpack.darwin_theory.getit (it, key)

generator to obtain items from nested iterable

xrayutilities.simpack.fit module

class xrayutilities.simpack.fit.FitModel (Imodel, verbose=False, plot=False, elog=True, **kwargs)

Bases: object

Wrapper for the Imfit Model class working for instances of LayerModel

Typically this means that after initialization of *FitModel* you want to use make_params to get a *Imfit.Parameters* list which one customizes for fitting.

Later on you can call fit and eval methods with those parameter list.

fit (data, params, x, weights=None, fit_kws=None, **kwargs)

wrapper around Imfit.Model.fit which enables plotting during the fitting

Parameters: data: ndarray

experimental values
params: Imfit.Parameters

list of parameters for the fit, use make_params for generation

x: ndarray

independent variable (incidence angle or q-position depending on the model)

weights: ndarray, optional

values of weights for the fit, same size as data

fit_kws : dict, optional

Options to pass to the minimizer being used

kwargs: dict, optional

keyword arguments which are passed to Imfit.Model.fit

Returns: Imfit.ModelResult

set_fit_limits (xmin=- inf, xmax=inf, mask=None)

set fit limits. If mask is given it must have the same size as the *data* and *x* variables given to fit. If mask is None it will be generated from xmin and xmax.

Parameters: xmin: float, optional

minimum value of x-values to include in the fit

xmax: float, optional

maximum value of x-values to include in the fit

mask: boolean array, optional

mask to be used for the data given to the fit

xrayutilities.simpack.helpers module

xrayutilities.simpack.helpers.coplanar_alphai (qx, qz, en='config') calculate coplanar incidence angle from knowledge of the qx and qz coordinates

Parameters: qx: array-like

inplane momentum transfer component

qz: array-like

out of plane momentum transfer component

en: float or str, optional

x-ray energy (eV). By default the value from the config is used.

Returns: alphai: array-like

the incidence angle in degree. points in the Laue zone are set to 'nan'.

xrayutilities.simpack.helpers.get_qz (qx, alphai, en='config')

calculate the qz position from the qx position and the incidence angle for a coplanar diffraction geometry

Parameters: qx: array-like

inplane momentum transfer component

alphai: array-like

incidence angle (deg)

en: float or str, optional

x-ray energy (eV). By default the value from the config is used.

Returns: array-like

the qz position for the given incidence angle

xrayutilities.simpack.models module

class xrayutilities.simpack.models.DiffuseReflectivityModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.SpecularReflectivityModel

model for diffuse reflectivity calculations

The 'simulate' method calculates the diffuse reflectivity on the specular rod in coplanar geometry in analogy to the SpecularReflectivityModel.

The 'simulate_map' method calculates the diffuse reflectivity for a 2D set of Q-positions. This method can also calculate the intensity for other geometries, like GISAXS with constant incidence angle or a quasi omega/2theta scan in GISAXS geometry.

simulate (alphai)

performs the actual diffuse reflectivity calculation for the specified incidence angles. This method always uses the coplanar geometry independent of the one set during the initialization.

Parameters: alphai: array-like

vector of incidence angles

Returns: array-like

vector of intensities of the reflectivity signal

simulate_map (qL, qz)

performs diffuse reflectivity calculation for the rectangular grid of reciprocal space positions define by qL and qz. This method uses the method and geometry set during the initialization of the class.

Parameters: qL: array-like

lateral coordinate in reciprocal space (vector with NqL components)

qz: array-like

vertical coordinate in reciprocal space (vector with Ngz components)

Returns: array-like

matrix of intensities of the reflectivity signal, with shape (len(qL), len(qz))

class xrayutilities.simpack.models.DynamicalModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.SimpleDynamicalCoplanarModel

Dynamical diffraction model for specular and off-specular qz-scans. Calculation of the flux of reflected and diffracted waves for general asymmetric coplanar diffraction from an arbitrary pseudomorphic multilayer is performed by a generalized 2-beam theory (4 tiepoints, S and P polarizations)

The first layer in the model is always assumed to be the semiinfinite substrate indepentent of its given thickness

simulate (alphai, hkl=None, geometry='hi_lo', rettype='intensity')

performs the actual diffraction calculation for the specified incidence angles and uses an analytic solution for the quartic dispersion equation

Parameters: alphai: array-like

vector of incidence angles (deg)

hkl: list or tuple, optional

Miller indices of the diffraction vector (preferable use set_hkl method to speed up repeated calculations of the same peak!)

geometry : {'hi_lo', 'lo_hi'}, optional

'hi_lo' for grazing exit (default) and 'lo_hi' for grazing incidence

rettype: {'intensity', 'field', 'all'}, optional

type of the return value. 'intensity' (default): returns the diffracted beam flux convoluted with the resolution function; 'field': returns the electric field (complex) without convolution with the resolution function, 'all': returns the electric field, ai, af (both in degree), and the reflected intensity.

Returns: array-like

vector of intensities of the diffracted signal, possibly changed return value due the rettype setting!

class xrayutilities.simpack.models.DynamicalReflectivityModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.SpecularReflectivityModel

model for Dynamical Specular Reflectivity Simulations. It uses the transfer Matrix methods as given in chapter 3 "Daillant, J., & Gibaud, A. (2008). X-ray and Neutron Reflectivity"

scanEnergy (energies, angle)

Simulates the Dynamical Reflectivity as a function of photon energy at fixed angle.

Parameters: energies: numpy.ndarray or list

photon energies (in eV).

angle: float

fixed incidence angle

Returns: reflectivity: array-like

vector of intensities of the reflectivity signal

transmitivity: array-like

vector of intensities of the transmitted signal

simulate (alphai)

Simulates the Dynamical Reflectivity as a function of angle of incidence

Parameters: alphai: array-like

vector of incidence angles

Returns: reflectivity: array-like

vector of intensities of the reflectivity signal

transmitivity: array-like

vector of intensities of the transmitted signal

class xrayutilities.simpack.models.KinematicalModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.LayerModel

Kinematical diffraction model for specular and off-specular qz-scans. The model calculates the kinematical contribution of one (hkl) Bragg peak, however considers the variation of the structure factor for different 'q'. The surface geometry is specified using the Experiment-object given to the constructor.

init_chi0()

calculates the needed optical parameters for the simulation. If any of the materials/layers is changing its properties this function needs to be called again before another correct simulation is made. (Changes of thickness does NOT require this!)

simulate (qz, hkl, absorption=False, refraction=False, rettype='intensity')

performs the actual kinematical diffraction calculation on the Qz positions specified considering the contribution from a single Bragg peak.

Parameters: qz: array-like

simulation positions along qz

hkl: list or tuple

Miller indices of the Bragg peak whos truncation rod should be calculated

absorption: bool, optional

flag to tell if absorption correction should be used

refraction: bool, optional

flag to tell if basic refraction correction should be performed. If refraction is True absorption correction is also included independent of the absorption flag.

rettype: {'intensity', 'field', 'all'}

type of the return value. 'intensity' (default): returns the diffracted beam flux convoluted with the resolution function; 'field': returns the electric field (complex) without convolution with the resolution function, 'all': returns the electric field, ai, af (both in degree), and the reflected intensity.

Returns: array-like

return value depends on the setting of *rettype*, by default only the calculate intensity is returned

class xrayutilities.simpack.models.KinematicalMultiBeamModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.KinematicalModel

Kinematical diffraction model for specular and off-specular qz-scans. The model calculates the kinematical contribution of several Bragg peaks on the truncation rod and considers the variation of the structure factor. In order to use a analytical description for the kinematic diffraction signal all layer thicknesses are changed to a multiple of the respective lattice parameter along qz. Therefore this description only works for (001) surfaces.

simulate (qz, hkl, absorption=False, refraction=True, rettype='intensity')

performs the actual kinematical diffraction calculation on the Qz positions specified considering the contribution from a full truncation rod

Parameters: qz: array-like

simulation positions along qz

hkl: list or tuple

Miller indices of the Bragg peak whos truncation rod should be calculated

absorption: bool, optional

flag to tell if absorption correction should be used

refraction: bool, optional,

flag to tell if basic refraction correction should be performed. If refraction is True absorption correction is also included independent of the absorption flag.

rettype: {'intensity', 'field', 'all'}

type of the return value. 'intensity' (default): returns the diffracted beam flux convoluted with the resolution function; 'field': returns the electric field (complex) without convolution with the resolution function, 'all': returns the electric field, ai, af (both in degree), and the reflected intensity.

Returns: array-like

return value depends on the setting of *rettype*, by default only the calculate intensity is returned

class xrayutilities.simpack.models.LayerModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.Model, abc.ABC

generic model class from which further thin film models can be derived from

get_polarizations()

return list of polarizations which should be calculated

join polarizations (Is, Ip)

method to calculate the total diffracted intensity from the intensities of S and P-polarization.

abstract simulate ()

abstract method that every implementation of a LayerModel has to override.

class xrayutilities.simpack.models.Model (experiment, **kwargs)

Bases: object

generic model class from which further models can be derived from

convolute_resolution (x, y)

convolve simulation result with a resolution function

Parameters: x: array-like

x-values of the simulation, units of x also decide about the unit of the resolution_width parameter

y: array-like

y-values of the simulation

Returns: array-like

convoluted y-data with same shape as y

property energy

scale_simulation(y)

scale simulation result with primary beam flux/intensity and add a background.

Parameters: y: array-like

y-values of the simulation

Returns: array-like

scaled y-values

class xrayutilities.simpack.models.ResonantReflectivityModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.SpecularReflectivityModel

model for specular reflectivity calculations CURRENTLY UNDER DEVELOPEMENT! DO NOT USE!

simulate (alphai)

performs the actual reflectivity calculation for the specified incidence angles

Parameters: alphai: array-like

vector of incidence angles

Returns: array-like

vector of intensities of the reflectivity signal

class xrayutilities.simpack.models.SimpleDynamicalCoplanarModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.KinematicalModel

Dynamical diffraction model for specular and off-specular qz-scans. Calculation of the flux of reflected and diffracted waves for general asymmetric coplanar diffraction from an arbitrary pseudomorphic multilayer is performed by a simplified 2-beam theory (2 tiepoints, S and P polarizations)

No restrictions are made for the surface orientation.

The first layer in the model is always assumed to be the semiinfinite substrate indepentent of its given thickness

Note

This model should not be used in real life scenarios since the made approximations severely fail for distances far from the reference position.

set_hkl (*hkl)

To speed up future calculations of the same Bragg peak optical parameters can be pre-calculated using this function.

Parameters: hkl : list or tuple

Miller indices of the Bragg peak for the calculation

simulate (alphai, hkl=None, geometry='hi_lo', idxref=1)

performs the actual diffraction calculation for the specified incidence angles.

Parameters: alphai: array-like

vector of incidence angles (deg)

hkl: list or tuple, optional

Miller indices of the diffraction vector (preferable use set_hkl method to speed up repeated calculations of the same peak!)

geometry: {'hi_lo', 'lo_hi'}, optional

'hi_lo' for grazing exit (default) and 'lo_hi' for grazing incidence

idxref: int, optional

index of the reference layer. In order to get accurate peak position of the film peak you want this to be the index of the film peak (default: 1). For the substrate use 0.

Returns: array-like

vector of intensities of the diffracted signal

class xrayutilities.simpack.models.SpecularReflectivityModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.LayerModel

model for specular reflectivity calculations

densityprofile (nz, plot=False)

calculates the electron density of the layerstack from the thickness and roughness of the individual layers

Parameters: nz: int

number of values on which the profile should be calculated

plot: bool, optional

flag to tell if a plot of the profile should be created

Returns: z : array-like

z-coordinates, z = 0 corresponds to the surface

eprof : array-like
 electron profile

init_cd()

calculates the needed optical parameters for the simulation. If any of the materials/layers is changing its properties this function needs to be called again before another correct simulation is made. (Changes of thickness and roughness do NOT require this!)

simulate (alphai)

performs the actual reflectivity calculation for the specified incidence angles

Parameters: alphai : array-like

vector of incidence angles

Returns: array-like

vector of intensities of the reflectivity signal

xrayutilities.simpack.models.effectiveDensitySlicing (layerstack, step, roughness=0, cutoff=1e-05)

Function to slice a LayerStack into many amorphous sublayers for effective density modelling of X-ray reflectivity of thin and rough multilayers. The resulting LayerStack will consist of perfectly smooth layers with average density/composition resulting from an error-function like transition between the rough layers of the initial stack. At the surface an vacuum layer is automatically added to the initial stack.

Parameters: layerstack: initial LayerStack, can contain only Amorhous layers!

step: thickness (in Angstrom) of the slices in the returned

LayerStack

roughness: roughness of the created sublayers (in Angstrom)

cutoff: layers with relative weights below this value will be ignored

Returns: LayerStack

xrayutilities.simpack.models.startdelta (start, delta, num)

xrayutilities.simpack.mosaicity module

xrayutilities.simpack.mosaicity.mosaic_analytic (qx, qz, RL, RV, Delta, hx, hz, shape) simulation of the coplanar reciprocal space map of a single mosaic layer using a simple analytic approximation

Parameters: qx: array-like

vector of the qx values (offset from the Bragg peak)

qz: array-like

vector of the qz values (offset from the Bragg peak)

RL: float

lateral block radius in Angstrom

RV: float

vertical block radius in Angstrom

Delta: float

root mean square misorientation of the grains in degree

hx: float

lateral component of the diffraction vector

hz: float

vertical component of the diffraction vector

shape: float

shape factor (1..Gaussian)

Returns: array-like

2D array with calculated intensities

xrayutilities.simpack.powder module

This module contains the core definitions for the XRD Fundamental Parameneters Model (FPA) computation in Python. The main computational class is FP_profile, which stores cached information to allow it to efficiently recompute profiles when parameters have been modified. For the user an Powder class is available which can calculate a complete powder pattern of a crystalline material.

The diffractometer line profile functions are calculated by methods from Cheary & Coelho 1998 and Mullen & Cline paper and 'R' package. Accumulate all convolutions in Fourier space, for efficiency, except for axial divergence, which needs to be weighted in real space for I3 integral.

More details about the applied algorithms can be found in the paper by M. H. Mendelhall et al., Journal of Research of NIST 120, 223 (2015) to which you should also refer for a careful definition of all the parameters

class xrayutilities.simpack.powder.FP_profile (anglemode, gaussian_smoother_bins_sigma=1.0,
oversampling=10)

Bases: object

the main fundamental parameters class, which handles a single reflection. This class is designed to be highly extensible by inheriting new convolvers. When it is initialized, it scans its namespace for specially formatted names, which can come from mixin classes. If it finds a function name of the form conv_xxx, it will call this funtion to create a convolver. If it finds a name of the form info_xxx it will associate the dictionary with that convolver, which can be used in UI generation, for example. The class, as it stands, does nothing significant with it. If it finds str_xxx, it will use that function to format a printout of the current state of the convolver conv_xxx, to allow improved report generation for convolvers.

When it is asked to generate a profile, it calls all known convolvers. Each convolver returns the Fourier transform of its convolvution. The transforms are multiplied together, inverse transformed, and after fixing the periodicity issue, subsampled, smoothed and returned.

If a convolver returns None, it is not multipled into the product.

Parameters: max_history_length: int

the number of histories to cache (default=5); can be overridden if memory is an issue.

length_scale_m : float

length_scale_m sets scaling for nice printing of parameters. if the units are in mm everywhere, set it to 0.001, e.g. convolvers which implement their own str_xxx method may use this to format their results, especially if 'natural' units are not meters. Typical is wavelengths and lattices in nm or angstroms, for example.

add_buffer (b)

add a numpy array to the list of objects that can be thrown away on pickling.

Parameters: b : array-like

the buffer to add to the list

Returns: b: array-like

return the same buffer, to make nesting easy.

axial_helper (outerbound, innerbound, epsvals, destination, peakpos=0, y0=0, k=0)

the function F0 from the paper. compute k/sqrt(peakpos-x)+y0 nonzero between outer & inner (inner is closer to peak) or k/sqrt(x-peakpos)+y0 if reversed (i.e. if outer > peak) fully evaluated on a specified eps grid, and stuff into destination

Parameters: outerbound: float

the edge of the function farthest from the singularity, referenced to epsvals

innerbound: float

the edge closest to the singularity, referenced to epsvals

epsvals: array-like

the array of two-theta values or offsets

destination : array-like

an array into which final results are summed. modified in place!

peakpos: float

the position of the singularity, referenced to epsvals.

y0: float

the constant offset

k: float

the scale factor

Returns: lower_index, upper_index : int

python style bounds for region of destination which has been modified.

compute_line_profile (convolver_names=None, compute_derivative=False, return_convolver=False)
 execute all the convolutions; if convolver_names is None, use everything we have, otherwise, use named convolutions.

Parameters: convolver_names: list

a list of convolvers to select. If None, use all found convolvers.

compute_derivative: bool

if True, also return d/dx(function) for peak position fitting

Returns: object

a profile_data object with much information about the peak

conv_absorption()

compute the sample transparency correction, including the finite-thickness version

Returns: array-like

the convolver

conv axial()

compute the Fourier transform of the axial divergence component

Returns: array-like

the transform buffer, or None if this is being ignored

conv_displacement ()

compute the peak shift due to sample displacement and the 2theta zero offset

Returns: array-like

the convolver

conv_emission()

compute the emission spectrum and (for convenience) the particle size widths

Returns: array-like

the convolver for the emission and particle sizes

Note

the particle size and strain stuff here is just to be consistent with *Topas* and to be vaguely efficient about the computation, since all of these have the same general shape.

conv_flat_specimen()

compute the convolver for the flat-specimen correction

Returns: array-like

the convolver

conv_global()

a dummy convolver to hold global variables and information. the global context isn't really a convolver, returning *None* means ignore result

Returns: None

always returns None

conv_receiver_slit ()

compute the rectangular convolution for the receiver slit or SiPSD pixel size

Returns: array-like

the convolver

conv_si_psd()

compute the convolver for the integral of defocusing of the face of an Si PSD

Returns: array-like

the convolver

conv_smoother()

compute the convolver to smooth the final result with a Gaussian before downsampling.

Returns: array-like

the convolver

conv_tube_tails()

compute the Fourier transform of the rectangular tube tails function

Returns: array-like

the transform buffer, or None if this is being ignored

full_axdiv_I2 (Lx=None, Ls=None, Lr=None, R=None, twotheta=None, beta=None, epsvals=None)
return the I2 function

Parameters: Lx: float

length of the xray filament

Ls: float

length of the sample

Lr: float

length of the receiver slit

R: float

diffractometer length, assumed symmetrical

twotheta: float

angle, in radians, of the center of the computation

beta: *float* offset angle

epsvals : array-like

array of offsets from center of computation, in radians

Returns: epsvals: array-like

array of offsets from center of computation, in radians

idxmin, idxmax: int

the full python-style bounds of the non-zero region of I2p and I2m

I2p, I2m: array-like

12+ and 12- from the paper, the contributions to the intensity

full_axdiv_I3 (Lx=None, Ls=None, Lr=None, R=None, twotheta=None, epsvals=None, sollerIdeg=None,
sollerDdeg=None, nsteps=10, axDiv=")

carry out the integral of I2 over beta and the Soller slits.

Parameters: Lx: float

length of the xray filament

Ls: float

length of the sample

Lr: float

length of the receiver slit

R: float

the (assumed symmetrical) diffractometer radius

twotheta: float

angle, in radians, of the center of the computation

epsvals: array-like

array of offsets from center of computation, in radians

sollerideg: float

the full-width (both sides) cutoff angle of the incident Soller slit

sollerDdeg: float

the full-width (both sides) cutoff angle of the detector Soller slit

nsteps: int

the number of subdivisions for the integral

axDiv : str not used

Returns: array-like

the accumulated integral, a copy of a persistent buffer _axial

general_tophat (name=", width=None)

a utility to compute a transformed tophat function and save it in a convolver buffer

Parameters: name: str

the name of the convolver cache buffer to update

width: float

the width in 2-theta space of the tophat

Returns: array-like

the updated convolver buffer, or None if the width was None

get_conv (name, key, format=<class 'float'>)

get a cached, pre-computed convolver associated with the given parameters, or a newly zeroed convolver if the cache doesn't contain it. Recycles old cache entries.

This takes advantage of the mutability of arrays. When the contents of the array are changed by the convolver, the cached copy is implicitly updated, so that the next time this is called with the same parameters, it will return the previous array.

Parameters: name: str

the name of the convolver to seek

key: object

any hashable object which identifies the parameters for the computation

format: numpy.dtype, optional

the type of the array to create, if one is not found.

Returns: bool

flag, which is *True* if valid data were found, or *False* if the returned array is zero, and *array*, which must be computed by the convolver if *flag* was *False*.

get_convolver_information()

return a list of convolvers, and what we know about them. function scans for functions named conv_xxx, and associated info_xxx entries.

Returns: list

list of (convolver_xxx, info_xxx) pairs

get_function_name ()

return the name of the function that called this. Useful for convolvers to identify themselves

Returns: str

name of calling function

get_good_bin_count (count)

find a bin count close to what we need, which works well for Fourier transforms.

Parameters: count: int

a number of bins.

Returns: int

a bin count somewhat larger than count which is efficient for FFT

info_emission = {'group_name': 'Incident beam and crystal size', 'help': 'this should be help information', 'param_info': {'crystallite_size_gauss': ('Gaussian crystallite size fwhm (m)', 1e-06), 'crystallite_size_lor': ('Lorentzian crystallite size fwhm (m)', 1e-06), 'emiss_gauss_widths': ('Gaussian emissions fwhm (m)', (1e-13,)), 'emiss_intensities': ('relative intensities', (1.0,)), 'emiss_lor_widths': ('Lorenztian emission fwhm (m)', (1e-13,)), 'emiss_wavelengths': ('wavelengths (m)', (1.58e-10,))}}

info_global = {'group_name': 'Global parameters', 'help': 'this should be help information', 'param_info': {'d': ('d spacing (m)', 4e-10), 'dominant_wavelength': ('wavelength of most intense line (m)', 1.5e-10), 'twotheta0_deg': ('Bragg center of peak (degrees)', 30.0)}}

classmethod isequivalent (hkl1, hkl2, crystalsystem)

function to determine if according to the convolvers included in this class two sets of Miller indices are equivalent. This function is only called when the class attribute 'isotropic' is False.

Parameters: hkl1, hkl2: list or tuple

Miller indices to be checked for equivalence

crystalsystem: str

symmetry class of the material which is considered

Returns: bool

isotropic = True

length scale m = 1.0

max_history_length = 5

self_clean()

do some cleanup to make us more compact; Instance can no longer be used after doing this, but can be pickled.

set_optimized_window (twotheta_window_center_deg, twotheta_approx_window_fullwidth_deg,
twotheta exact bin spacing deg)

pick a bin count which factors cleanly for FFT, and adjust the window width to preserve the exact center and bin spacing

Parameters: twotheta window center deg: float

exact position of center bin, in degrees

twotheta_approx_window_fullwidth_deg: float

approximate desired width

twotheta_exact_bin_spacing_deg: float

the exact bin spacing to use

set_parameters (convolver='global', **kwargs)

update the dictionary of parameters associated with the given convolver

Parameters: convolver: str

the name of the convolver. name 'global', e.g., attaches to function 'conv_global'

kwargs: dict

keyword-value pairs to update the convolvers dictionary.

set_window (twotheta_window_center_deg, twotheta_window_fullwidth_deg, twotheta_output_points)
move the compute window to a new location and compute grids, without resetting all parameters. Clears convolution history and sets up many arrays.

Parameters: twotheta_window_center_deg: float

the center position of the middle bin of the window, in degrees

twotheta_window_fullwidth_deg: float the full width of the window, in degrees

twotheta_output_points: int

the number of bins in the final output

str_emission()

format the emission spectrum and crystal size information

Returns: str

the formatted information

str global()

returns a string representation for the global context.

Returns: str

report on global parameters.

class xrayutilities.simpack.powder.PowderDiffraction (mat, **kwargs)

Bases: xrayutilities.experiment.PowderExperiment

Experimental class for powder diffraction. This class calculates the structure factors of powder diffraction lines and uses instances of FP_profile to perform the convolution with experimental resolution function calculated by the fundamental parameters approach. This class uses multiprocessing to speed up calculation. Set config.NTHREADS=1 to restrict this to one worker process.

Calculate (twotheta, **kwargs)

calculate the powder diffraction pattern including convolution with the resolution function and map them onto the twotheta positions. This also performs the calculation of the peak intensities from the internal material object

Parameters: twotheta: array-like

two theta values at which the powder pattern should be calculated.

kwargs: dict

additional keyword arguments are passed to the Convolve function

Returns: array-like

output intensity values for the twotheta values given in the input

Notes

Bragg peaks are only included up to tt cutoff set in the class constructor!

Convolve (twotheta, window_width='config', mode='multi')

convolute the powder lines with the resolution function and map them onto the twotheta positions. This calculates the powder pattern excluding any background contribution

Parameters: twotheta: array-like

two theta values at which the powder pattern should be calculated.

window_width: float, optional

width of the calculation window of a single peak

mode: {'multi, 'local'}, optional

multiprocessing mode, either 'multi' to use multiple processes or 'local' to restrict the calculation to a single process

Note:

Bragg peaks are only included up to tt cutoff set in the class constructor!

Returns: output intensity values for the twotheta values given in the input

close ()

correction_factor (ang)

calculate the correction factor for the diffracted intensities. This contains the polarization effects and the Lorentz factor

Parameters: ang: aray-like

theta diffraction angles for which the correction should be calculated

Returns: f: array-like

array of the same shape as ang containing the correction factors

property energy

init_powder_lines (tt_cutoff)

calculates the powder intensity and positions up to an angle of tt_cutoff (deg) and stores the result in the data dictionary whose structure is as follows:

The data dictionary has one entry per line with a unique identifier as key of the entry. The entries themself are dictionaries which have the following entries:

- hkl : (h, k, l), Miller indices of the Bragg peak
- r : reflection strength of the line
- ang : Bragg angle of the peak (theta = 2theta/2!)
- · qpos: reciprocal space position

load_settings_from_config (settings)

load parameters from the config and update these settings with the options from the settings parameter

merge_lines (data)

if calculation is isotropic lines at the same q-position can be merged to one line to reduce the calculational effort

Parameters: data: ndarray

numpy field array with values of 'hkl' (Miller indices of the peaks), 'q' (q-position), and 'r' (reflection etropeth) as produced by the reflection etropeth method

'r' (reflection strength) as produced by the reflection_strength method

Returns: hkl, q, ang, r: array-like

Miller indices, q-position, diffraction angle (Theta), and reflection strength of the

material

reflection_strength (tt_cutoff)

determine structure factors/reflection strength of all Bragg peaks up to tt_cutoff. This function also implements the March-Dollase model for preferred orientation in the symmetric reflection mode. Note that although this means the sample has anisotropic properties the various lines can still be merged together since at the moment no anisotropic crystal shape is supported.

Parameters: tt_cutoff: float

upper cutoff value of 2theta until which the reflection strength are calculated

Returns: ndarray

numpy array with field for 'hkl' (Miller indices of the peaks), 'q' (q-position), and 'r'

(reflection strength) of the Bragg peaks

set_sample_parameters()

load sample parameters from the Powder class and use them in all FP profile instances of this object

set_wavelength_from_params()

sets the wavelenth in the base class from the settings dictionary of the FP_profile classes and also set it in the 'global' part of the parameters

set window (force=False)

sets the calcultion window for all convolvers

property twotheta

update_powder_lines (tt_cutoff)

calculates the powder intensity and positions up to an angle of tt_cutoff (deg) and updates the values in:

- ids: list of unique identifiers of the powder line
- data: array with intensities
- ang: bragg angles of the peaks (theta=2theta/2!)
- qpos: reciprocal space position of intensities

update_settings (newsettings={})

update settings of all instances of FP_profile

Parameters:

newsettings: dict

```
dictionary with new settings. It has to include one subdictionary for every convolver
                           which should have its settings changed.
  property wavelength
  property window_width
xrayutilities.simpack.powder.chunkify(lst, n)
class xrayutilities.simpack.powder.convolver_handler
  Bases: object
  manage the convolvers of on process
  add_convolver (convolver)
  calc (run, ttpeaks)
    calculate profile function for selected convolvers
        Parameters: run: list
                           list of flags of length of convolvers to tell which convolver needs to be run
                       ttpeaks: array-like
                           peak positions for the convolvers
            Returns:
                      list
                           list of profile data result objects
  set_windows (centers, npoints, flag, width)
  update_parameters (parameters)
class xrayutilities.simpack.powder.manager (address=None, authkey=None, serializer='pickle', ctx=None)
  Bases: multiprocessing.managers.BaseManager
class xrayutilities.simpack.powder.profile_data (**kwargs)
  Bases: object
  a skeleton class which makes a combined dict and namespace interface for easy pickling and data passing
```

xrayutilities.simpack.powdermodel module

Parameters: kwargs: dict

add new symbols to both the attributes and dictionary for the class

keyword=value pairs

add_symbol (**kwargs)

```
class xrayutilities.simpack.powdermodel.PowderModel (*args, **kwargs)
Bases: object
Class to help with powder calculations for multiple materials. For basic calculations the Powder class together with the Fundamental parameters approach is used.

close ()
create_fitparameters ()
function to create a fit model with all instrument and sample parameters.
```

Returns: Imfit.Parameters

fit (params, twotheta, data, std=None, maxfev=200)
make least squares fit with parameters supplied by the user

Parameters: params: Imfit.Parameters

object with all parameters set as intended by the user

twotheta: array-like

angular values for the fit

data: array-like

experimental intensities for the fit

std: array-like

standard deviation of the experimental data. if 'None' the sqrt of the data will be used

maxfev: int

maximal number of simulations during the least squares refinement

Returns: Imfit.MinimizerResult

plot (twotheta, showlines=True, label='simulation', color=None, formatspec='-', lcolors=[], ax=None, **kwargs) plot the powder diffraction pattern and indicate line positions for all components in the model.

Parameters: twotheta: array-like

positions at which the powder pattern should be evaluated

showlines: bool, optional

flag to decide if peak positions of the components will be shown on the top of the plot

label: str

line label in the plot

color: matplotlib color or None

the color used for the line plot of the simulation

formatspec : str

format specifier of the simulation curve

Icolors: list of matplotlib colors

colors for the line indicators for the various components

ax: matplotlib.axes or None

axes object to be used for plotting, if its given no axes decoration like labels are set.

Further keyword arguments are passed to the simulate method.

Returns: matplotlib.axes object

set_background (btype, **kwargs)
define background as spline or polynomial function

Parameters: btype: {polynomial', 'spline'}

background type; Depending on this value the expected keyword arguments differ.

kwargs: dict

optional keyword arguments

x: array-like, optional

x-values (twotheta) of the background points (if btype='spline')

y: array-like, optional

intensity values of the background (if btype='spline')

p: array-like, optional

polynomial coefficients from the highest degree to the constant term. len of p decides about the degree of the polynomial (if btype='polynomial')

set_lmfit_parameters (Imparams)

function to update the settings of this class during an least squares fit

Parameters: Imparams: Imfit.Parameters

Imfit Parameters list of sample and instrument parameters

set_parameters (params)

set simulation parameters of all subobjects

Parameters: params: dict

settings dictionaries for the convolvers.

simulate (twotheta, **kwargs)

calculate the powder diffraction pattern of all materials and sum the results based on the relative volume of the materials.

Parameters: twotheta: array-like

positions at which the powder pattern should be evaluated

kwargs: dict

optional keyword arguments

background: array-like

an array of background values (same shape as twotheta) if no background is given then the background is calculated as previously set by the set_background function or is 0

further keyword arguments are passed to the Convolve function of of the

PowderDiffraction objects

Returns: array-like

summed powder diffraction intensity of all materials present in the model

xrayutilities.simpack.powdermodel.Rietveld_error_metrics (exp, sim, weight=None, std=None,
Nvar=0, disp=False)

calculates common error metrics for Rietveld refinement.

Parameters: exp: array-like

experimental datapoints

sim : array-like simulated data

weight: array-like, optional

weight factor in the least squares sum. If it is None the weight is estimated from the counting statistics of 'exp'

std: array-like, optional

standard deviation of the experimental data. alternative way of specifying the weight factor, when both are given weight overwrites std!

Nvar: int, optional

number of variables in the refinement

disp: bool, optional

flag to tell if a line with the calculated values should be printed.

Returns: M, Rp, Rwp, Rwpexp, chi2: float

xrayutilities.simpack.powdermodel.plot_powder (twotheta, exp, sim, mask=None, scale='sqrt',
fig='XU:powder', show_diff=True, show_legend=True, labelexp='experiment', labelsim='simulation', formatexp='.-k',
formatsim='-r')

Convenience function to plot the comparison between experimental and simulated powder diffraction data

Parameters: twotheta: array-like

angle values used for the x-axis of the plot (deg)

exp: array-like

experimental data (same shape as twotheta). If None only the simulation and no difference will be plotted

sim: array-like or PowederModel

simulated data or PowderModel instance. If a PowderModel instance is given the plot-method of PowderModel is used.

mask: array-like, optional

mask to reduce the twotheta values to the be used as x-coordinates of sim

scale: {'linear', 'sqrt', 'log'}, optional

string specifying the scale of the y-axis.

fig: str or int, optional

matplotlib figure name (figure will be cleared!)

show_diff: bool, optional

flag to specify if a difference curve should be shown

show_legend: bool, optional

flag to specify if a legend should be shown

labelexp: str

plot label (legend entry) for the experimental data

labelsim: str

plot label for the simulation data

formatexp: str

format specifier for the experimental data

formatsim: str

format specifier for the simulation curve

xrayutilities.simpack.smaterials module

```
class xrayutilities.simpack.smaterials.CrystalStack (name, *args)
  Bases: xrayutilities.simpack.smaterials.LayerStack
  extends the built in list type to enable building a stack of crystalline Layers by various methods.
  check (V)
class xrayutilities.simpack.smaterials.GradedLayerStack (alloy, xfrom, xto, nsteps, thickness,
**kwargs)
 Bases: xrayutilities.simpack.smaterials.CrystalStack
  generates a sequence of layers with a gradient in chemical composition
class xrayutilities.simpack.smaterials.Layer (material, thickness, **kwargs)
  Bases: xrayutilities.simpack.smaterials.SMaterial
  Object describing part of a thin film sample. The properties of a layer are:
       Attributes:
                    material: Material (Crystal or Amorhous)
                        an xrayutilties material describing optical and crystal properties of the thin film
                    thickness: float
                        film thickness in Angstrom
class xrayutilities.simpack.smaterials.LayerStack (name, *args)
  Bases: xrayutilities.simpack.smaterials.MaterialList
  extends the built in list type to enable building a stack of Layer by various methods.
  check (V)
class xrayutilities.simpack.smaterials.MaterialList (name, *args)
  Bases: collections.abc.MutableSequence
  class representing the basics of a list of materials for simulations within xrayutilities. It extends the built in list type.
  check (V)
  insert (i, v)
    S.insert(index, value) – insert value before index
class xrayutilities.simpack.smaterials.Powder (material, volume, **kwargs)
  Bases: xrayutilities.simpack.smaterials.SMaterial
  Object describing part of a powder sample. The properties of a powder are:
```

```
Attributes:
                     material: Crystal
                         an xrayutilties material (Crystal) describing optical and crystal properties of the powder
                     volume: float
                         powder's volume (in pseudo units, since only the relative volume enters the calculation)
                     crystallite size lor: float, optional
                         Lorentzian crystallite size fwhm (m)
                     crystallite_size_gauss: float, optional
                         Gaussian crystallite size fwhm (m)
                     strain lor: float, optional
                         extra peak width proportional to tan(theta)
                     strain_gauss: float, optional
                         extra peak width proportional to tan(theta)
                     preferred_orientation: tuple, optional
                         HKL of the preferred orientation
                     preferred_orientation_factor: float, optional
                         March-Dollase preferred orientation factor: > 1 for platy crystallits , < 1 for rod-like
                         crystallites, and = 1 for random orientation of crystallites.
class xrayutilities.simpack.smaterials.PowderList (name, *args)
  Bases: xrayutilities.simpack.smaterials.MaterialList
  extends the built in list type to enable building a list of Powder by various methods.
  check (V)
class xrayutilities.simpack.smaterials.PseudomorphicStack001 (name, *args)
  Bases: xrayutilities.simpack.smaterials.CrystalStack
  generate a sequence of pseudomorphic crystalline Layers. Surface orientation is assumed to be 001 and materials
  must be cubic/tetragonal.
  insert (i, v)
    S.insert(index, value) – insert value before index
 make_epitaxial (i)
  trans = <xrayutilities.math.transforms.Transform object>
class xrayutilities.simpack.smaterials.PseudomorphicStack111 (name, *args)
  Bases: xrayutilities.simpack.smaterials.PseudomorphicStack001
  generate a sequence of pseudomorphic crystalline Layers. Surface orientation is assumed to be 111 and materials
  must be cubic.
  trans = <xrayutilities.math.transforms.CoordinateTransform object>
class xrayutilities.simpack.smaterials.SMaterial (material, **kwargs)
  Bases: object
  Simulation Material. Extends the xrayutilities Materials by properties needed for simulations
  property material
Module contents
```

simulation subpackage of xrayutilities.

This package provides possibilities to simulate X-ray diffraction and reflectivity curves of thin film samples. It could be extended for more general use in future if there is demand for that.

In addition it provides a fitting routine for reflectivity data which is based on Imfit.

xrayutilities

xrayutilities package

Subpackages

xrayutilities.analysis package

Submodules

xrayutilities.analysis.line_cuts module

xrayutilities.analysis.line_cuts.get_arbitrary_line (qpos, intensity, point, vec, npoints, intrange) extracts a line scan from reciprocal space map data along an arbitrary line defined by the point 'point' and propergation vector 'vec'. Integration of the data is performed in a cylindrical volume along the line. This function works for 2D and 3D input data!

Parameters: qpos: list of array-like objects

arrays of x, y (list with two components) or x, y, z (list with three components)

momentum transfers

intensity: array-like

2D or 3D array of reciprocal space intensity with shape equal to the qpos entries

point: tuple, list or array-like

point on the extraction line (2 or 3 coordinates)

vec: tuple, list or array-like

propergation vector of the extraction line (2 or 3 coordinates)

npoints: int

number of points in the output data

intrange : float

radius of the cylindrical integration volume around the extraction line

Returns: qpos, qint : ndarray

line scan coordinates and intensities

used_mask : ndarray

mask of used data, shape is the same as the input intensity: True for points which

contributed, False for all others

Examples

```
>>> qcut, qint, mask = get_arbitrary_line([qx, qy, qz], inten, (1.1, 2.2, 0.0), (1, 1, 1), 200, 0.1)
```

xrayutilities.analysis.line_cuts.get_omega_scan (qpos, intensity, cutpos, npoints, intrange, **kwargs) extracts an omega scan from reciprocal space map data with integration along either the 2theta, or radial (omega-2theta) direction. The coplanar diffraction geometry with qy and qz as diffraction plane is assumed. This is consistent with the coplanar geometry implemented in the HXRD-experiment class.

This function works for 2D and 3D input data in the same way!

Parameters: qpos: list of array-like objects

arrays of y, z (list with two components) or x, y, z (list with three components)

momentum transfers

intensity: array-like

2D or 3D array of reciprocal space intensity with shape equal to the qpos entries

cutpos: tuple or list

y/z-position or x/y/z-position at which the line scan should be extracted. this must be

have two entries for 2D data (z-position) and a three for 3D data

npoints: int

number of points in the output data

intrange: float

integration range in along *intdir* in degree. data will be integrated from *-intrange* .. +*intrange*

intdir : {'2theta', 'radial'}, optional

integration direction: '2theta': scattering angle (default), or 'radial': omega-2theta

direction.

wl: float or str, optional

wavelength used to determine angular integration positions

Note:

Although applicable for any set of data, the extraction only makes sense when the data are aligned into the y/z-plane.

Returns: om, omint : ndarray

omega scan coordinates and intensities

used_mask : ndarray

mask of used data, shape is the same as the input intensity: True for points which contributed, False for all others

Examples

```
>>> omcut, omcut_int, mask = get_omega_scan([qy, qz], inten, [2.0, 5.0], 250, intrange=0.1)
```

xrayutilities.analysis.line_cuts.get_qx_scan (qpos, intensity, cutpos, npoints, intrange, **kwargs) extracts a qx scan from 3D reciprocal space map data with integration along either, the perpendicular plane in q-space, omega (sample rocking angle) or 2theta direction. For the integration in angular space (omega, or 2theta) the coplanar diffraction geometry with qy and qz as diffraction plane is assumed. This is consistent with the coplanar geometry implemented in the HXRD-experiment class.

Parameters: qpos: list of array-like objects

arrays of x, y, z (list with three components) momentum transfers

intensity: array-like

3D array of reciprocal space intensity with shape equal to the qpos entries

cutpos: tuple/list

y/z-position at which the line scan should be extracted. this must be and a tuple/list with the qy, qz cut position

npoints: int

number of points in the output data

intrange: float

integration range in along *intdir*, either in 1/AA (*q*) or degree ('omega', or '2theta'). data will be integrated from *-intrange* .. +*intrange*

intdir: {'q', 'omega', '2theta'}, optional

integration direction: 'q': perpendicular Q-plane (default), 'omega': sample rocking angle, or '2theta': scattering angle.

wl: float or str, optional

wavelength used to determine angular integration positions

Note:

The angular integration directions although applicable for any set of data only makes sense when the data are aligned into the y/z-plane.

Returns: qx, qxint: ndarray

qx scan coordinates and intensities

used_mask : ndarray

mask of used data, shape is the same as the input intensity: True for points which contributed, False for all others

Examples

```
>>> qxcut, qxcut_int, mask = get_qx_scan([qx, qy, qz], inten, [0, 2.0], 250, intrange=0.01)
```

xrayutilities.analysis.line_cuts.get_qy_scan (qpos, intensity, cutpos, npoints, intrange, **kwargs) extracts a qy scan from reciprocal space map data with integration along either, the perpendicular plane in q-space, omega (sample rocking angle) or 2theta direction. For the integration in angular space (omega, or 2theta) the coplanar diffraction geometry with qy and qz as diffraction plane is assumed. This is consistent with the coplanar geometry implemented in the HXRD-experiment class.

This function works for 2D and 3D input data in the same way!

arrays of y, z (list with two components) or x, y, z (list with three components)

momentum transfers

intensity: array-like

2D or 3D array of reciprocal space intensity with shape equal to the qpos entries

cutpos: float or tuple/list

x/z-position at which the line scan should be extracted. this must be a float for 2D data

(z-position) and a tuple with two values for 3D data

npoints: int

number of points in the output data

intrange: float

integration range in along *intdir*, either in 1/AA (*q*) or degree ('omega', or '2theta'). data will be integrated from *-intrange* .. +*intrange*

intdir: {'q', 'omega', '2theta'}, optional

integration direction: 'q': perpendicular Q-plane (default), 'omega': sample rocking

angle, or '2theta': scattering angle.

wavelength used to determine angular integration positions

Note:

For 3D data the angular integration directions although applicable for any set of data only makes sense when the data are aligned into the y/z-plane.

Returns: qy, qyint : ndarray

qy scan coordinates and intensities

used_mask : ndarray

wl: float or str, optional

mask of used data, shape is the same as the input intensity: True for points which contributed. False for all others

Examples

```
>>> qycut, qycut_int, mask = get_qy_scan([qy, qz], inten, 5.0, 250, intrange=0.02, intdir='2theta')
```

xrayutilities.analysis.line_cuts.get_qz_scan (qpos, intensity, cutpos, npoints, intrange, **kwargs) extracts a qz scan from reciprocal space map data with integration along either, the perpendicular plane in q-space, omega (sample rocking angle) or 2theta direction. For the integration in angular space (omega, or 2theta) the coplanar diffraction geometry with qy and qz as diffraction plane is assumed. This is consistent with the coplanar geometry implemented in the HXRD-experiment class.

This function works for 2D and 3D input data in the same way!

arrays of y, z (list with two components) or x, y, z (list with three components)

momentum transfers

intensity: array-like

2D or 3D array of reciprocal space intensity with shape equal to the qpos entries

cutpos: float or tuple/list

x/y-position at which the line scan should be extracted. this must be a float for 2D data

and a tuple with two values for 3D data

npoints: int

number of points in the output data

intrange: float

integration range in along *intdir*, either in 1/AA (*q*) or degree ('omega', or '2theta'). data will be integrated from *-intrange/2* .. *+intrange/2*

intdir: {'q', 'omega', '2theta'}, optional

integration direction: 'q': perpendicular Q-plane (default), 'omega': sample rocking angle, or '2theta': scattering angle.

wl: float or str, optional

wavelength used to determine angular integration positions

Note:

For 3D data the angular integration directions although applicable for any set of data only makes sense when the data are aligned into the y/z-plane.

Returns: qz, qzint : ndarray

qz scan coordinates and intensities

used_mask : ndarray

mask of used data, shape is the same as the input intensity: True for points which contributed, False for all others

Examples

```
>>> qzcut, qzcut_int, mask = get_qz_scan([qy, qz], inten, 3.0, 200, intrange=0.3)
```

xrayutilities.analysis.line_cuts.get_radial_scan (qpos, intensity, cutpos, npoints, intrange,
**kwarqs)

extracts a radial scan from reciprocal space map data with integration along either the omega or 2theta direction. The coplanar diffraction geometry with qy and qz as diffraction plane is assumed. This is consistent with the coplanar geometry implemented in the HXRD-experiment class.

This function works for 2D and 3D input data in the same way!

arrays of y, z (list with two components) or x, y, z (list with three components)

momentum transfers

intensity: array-like

2D or 3D array of reciprocal space intensity with shape equal to the qpos entries

cutpos: tuple or list

y/z-position or x/y/z-position at which the line scan should be extracted. this must be

have two entries for 2D data (z-position) and a three for 3D data

npoints: int

number of points in the output data

intrange: float

integration range in along *intdir* in degree. data will be integrated from *-intrange* .. +*intrange*

intdir: {'omega', '2theta'}, optional

integration direction: 'omega': sample rocking angle (default), '2theta': scattering angle

wl: float or str. optional

wavelength used to determine angular integration positions

Note:

Although applicable for any set of data, the extraction only makes sense when the data are aligned into the y/z-plane.

Returns: tt, omttint : ndarray

omega-2theta scan coordinates (2theta values) and intensities

used_mask : ndarray

mask of used data, shape is the same as the input intensity: True for points which contributed, False for all others

Examples

```
>>> ttcut, omtt_int, mask = get_radial_scan([qy, qz], inten, [2.0, 5.0], 250, intrange=0.1)
```

xrayutilities.analysis.line_cuts.get_ttheta_scan (qpos, intensity, cutpos, npoints, intrange,

**kwargs)

extracts a 2theta scan from reciprocal space map data with integration along either the omega or radial direction. The coplanar diffraction geometry with qy and qz as diffraction plane is assumed. This is consistent with the coplanar geometry implemented in the HXRD-experiment class.

This function works for 2D and 3D input data in the same way!

arrays of y, z (list with two components) or x, y, z (list with three components)

momentum transfers

intensity: array-like

2D or 3D array of reciprocal space intensity with shape equal to the qpos entries

cutpos: tuple or list

y/z-position or x/y/z-position at which the line scan should be extracted. this must be

have two entries for 2D data (z-position) and a three for 3D data

npoints: int

number of points in the output data

intrange: float

integration range in along *intdir* in degree. data will be integrated from *-intrange* .. +*intrange*

intdir: {'omega', 'radial'}, optional

integration direction: 'omega': sample rocking angle (default), 'radial': omega-2theta

direction

wl: float or str, optional

wavelength used to determine angular integration positions

Note:

Although applicable for any set of data, the extraction only makes sense when the data are aligned into the y/z-plane.

Returns: tt, ttint: ndarray

2theta scan coordinates and intensities

used_mask : ndarray

mask of used data, shape is the same as the input intensity: True for points which contributed, False for all others

Examples

```
>>> ttcut, tt_int, mask = get_ttheta_scan([qy, qz], inten, [2.0, 5.0], 250, intrange=0.1)
```

xrayutilities.analysis.misc module

miscellaneous functions helpful in the analysis and experiment

xrayutilities.analysis.misc.coplanar_intensity (mat, exp, hkl, thickness, thMono, sample_width=10, beam_width=1)

Calculates the expected intensity of a Bragg peak from an epitaxial thin film measured in coplanar geometry (integration over omega and 2theta in angular space!)

Parameters: mat: Crystal

Crystal instance for structure factor calculation

exp: Experiment

Experimental(HXRD) class for the angle calculation

hkl: list, tuple or array-like

Miller indices of the peak to calculate

thickness: float

film thickness in nm

thMono: float

Bragg angle of the monochromator (deg)

sample_width: float, optional

width of the sample along the beam

beam_width : float, optional

width of the beam in the same units as the sample size

Returns: float

intensity of the peak

xrayutilities.analysis.misc.getangles (peak, sur, inp) calculates the chi and phi angles for a given peak

Parameters: peak: list or array-like

hkl for the peak of interest

sur : list or array-like
 hkl of the surface
inp : list or array-like

inplane reference peak or direction

Returns: list

[chi, phi] for the given peak on surface sur with inplane direction inp as reference

Examples

To get the angles for the -224 peak on a 111 surface type

```
>>> [chi, phi] = getangles([-2, 2, 4], [1, 1, 1], [2, 2, 4])
```

xrayutilities.analysis.misc.getunitvector (chi, phi, ndir=(0, 0, 1), idir=(1, 0, 0))

return unit vector determined by spherical angles and definition of the polar axis and inplane reference direction (phi=0)

Parameters: chi, phi : float

spherical angles (polar and azimuthal) in degree

ndir: tuple, list or array-like

polar/z-axis (determines chi=0)

idir: tuple, list or array-like

azimuthal axis (determines phi=0)

xrayutilities.analysis.sample_align module

functions to help with experimental alignment during experiments, especially for experiments with linear and area detectors

xrayutilities.analysis.sample_align.area_detector_calib (angle1, angle2, ccdimages, detaxis, r_i, plot=True, cut_off=0.7, start=(None, None, 1, 0, 0, 0), fix=(False, False, True, False, False, False, False), fig=None, wl=None, plotlog=False, nwindow=50, debug=False)

function to calibrate the detector parameters of an area detector it determines the detector tilt possible rotations and offsets in the detector arm angles

Parameters: angle1: array-like

outer detector arm angle

angle2: array-like

inner detector arm angle

ccdimages: array-like

images of the ccd taken at the angles given above

detaxis: list of str

detector arm rotation axis; default: ['z+', 'y-']

r_i : str

primary beam direction [xyz][+-]; default 'x+'

plot: bool, optional

flag to determine if results and intermediate results should be plotted; default: True

cut_off : float, optional

cut off intensity to decide if image is used for the determination or not; default: 0.7 = 70%

start: tuple, optional

sequence of start values of the fit for parameters, which can not be estimated automatically or might want to be fixed. These are: pwidth1, pwidth2, distance, tiltazimuth, tilt, detector_rotation, outerangle_offset. By default (None, None, 1, 0, 0, 0, 0) is used.

fix: tuple of bool

fix parameters of start (default: (False, False, True, False, False, False, False)) It is strongly recommended to either fix the distance or the pwidth1, 2 values.

fig: Figure, optional

matplotlib figure used for plotting the error default: None (creates own figure)

wl: float or str

wavelength of the experiment in Angstrom (default: config.WAVELENGTH) value does not really matter here but does affect the scaling of the error

plotlog: bool

flag to specify if the created error plot should be on log-scale

nwindow: int

window size for determination of the center of mass position after the center of mass of every full image is determined, the center of mass is determined again using a window of size nwindow in order to reduce the effect of hot pixels.

debug: bool

flag to specify that you want to see verbose output and saving of images to show if the CEN determination works

xrayutilities.analysis.sample_align.area_detector_calib_hkl (sampleang, angle1, angle2, ccdimages, hkls, experiment, material, detaxis, r_i, plot=True, cut_off=0.7, start=(None, None, 1, 0, 0, 0, 0, 0, 0, 0); config'), fix=(False, False, True, False, False, False, False, False, False, False), fig=None, plotlog=False, nwindow=50, debug=False)

function to calibrate the detector parameters of an area detector it determines the detector tilt possible rotations and offsets in the detector arm angles

in this variant not only scans through the primary beam but also scans at a set of symmetric reflections can be used for the detector parameter determination. for this not only the detector parameters but in addition the sample orientation and wavelength need to be fit. Both images from the primary beam hkl = (0, 0, 0) and from a symmetric reflection hkl = (h, k, l) need to be given for a successful run.

Parameters: sampleang: array-like

sample rocking angle (needed to align the reflections (same rotation direction as inner detector rotation)) other sample angle are not allowed to be changed during the scans

angle1: array-like

outer detector arm angle

angle2: array-like

inner detector arm angle

ccdimages : array-like

images of the ccd taken at the angles given above

hkls: list or array-like

hkl values for every image

experiment: Experiment

Experiment class object needed to get the UB matrix for the hkl peak treatment

material: Crystal

material used as reference crystal

detaxis: list of str

detector arm rotation axis; default: ['z+', 'y-']

r_i : str

primary beam direction [xyz][+-]; default 'x+'

plot: bool, optional

flag to determine if results and intermediate results should be plotted; default: True

cut off: float, optional

cut off intensity to decide if image is used for the determination or not; default: 0.7 = 70%

start: tuple, optional

sequence of start values of the fit for parameters, which can not be estimated automatically or might want to be fixed. These are: pwidth1, pwidth2, distance, tiltazimuth, tilt, detector_rotation, outerangle_offset, sampletilt, sampletiltazimuth, wavelength. By default (None, None, 1, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0).

fix: tuple of bool

fix parameters of start (default: (False, False, True, False, False, False, False, False, False, False, False, False)) It is strongly recommended to either fix the distance or the pwidth1, 2 values.

fig: Figure, optional

matplotlib figure used for plotting the error default: None (creates own figure)

plotlog: bool

flag to specify if the created error plot should be on log-scale

nwindow: int

window size for determination of the center of mass position after the center of mass of every full image is determined, the center of mass is determined again using a window of size nwindow in order to reduce the effect of hot pixels.

debug: bool

flag to specify that you want to see verbose output and saving of images to show if the CEN determination works

xrayutilities.analysis.sample_align.fit_bragg_peak (om, tt, psd, omalign, ttalign, exphxrd, frange=(0.03, 0.03), peaktype='Gauss', plot=True)

helper function to determine the Bragg peak position in a reciprocal space map used to obtain the position needed for correction of the data. the determination is done by fitting a two dimensional Gaussian (xrayutilities.math.Gauss2d) or Lorentzian (xrayutilities.math.Lorentz2d)

PLEASE ALWAYS CHECK THE RESULT CAREFULLY!

Parameters: om, tt: array-like

angular coordinates of the measurement either with size of psd or of psd.shape[0]

psd: array-like

intensity values needed for fitting

omalign: float

aligned omega value, used as first guess in the fit

ttalign: float

aligned two theta values used as first guess in the fit these values are also used to set the range for the fit: the peak should be within +/-frangeAA^{-1} of those values

exphxrd: Experiment

experiment class used for the conversion between angular and reciprocal space.

frange: tuple of float, optional

data range used for the fit in both directions (see above for details default:(0.03, 0.03) unit: AA^{-1})

peaktype : {'Gauss', 'Lorentz'}

peak type to fit

plot: bool, optional

if True (default) function will plot the result of the fit in comparison with the measurement.

Returns: omfit, ttfit: float

fitted angular values

params: list

fit parameters (of the Gaussian/Lorentzian)

covariance: ndarray

covariance matrix of the fit parameters

xrayutilities.analysis.sample_align.linear_detector_calib (angle, mca_spectra, **keyargs) function to calibrate the detector distance/channel per degrees for a straight linear detector mounted on a detector arm

Parameters: angle: array-like

array of angles in degree of measured detector spectra

mca_spectra: array-like

corresponding detector spectra (shape: (len(angle), Nchannels)

r_i: str, optional

primary beam direction as vector [xyz][+-]; default: 'y+'

detaxis: str, optional

detector arm rotation axis [xyz][+-]; default: 'x+'

Returns: pixelwidth: float

width of the pixel at one meter distance, pixelwidth is negative in case the hit channel number decreases upon an increase of the detector angle

center_channel: float

central channel of the detector

detector_tilt: float, optional

if usetilt=True the fitted tilt of the detector is also returned

Note

L/pixelwidth*pi/180 ~= channel/degree, with the sample detector distance L

The function also prints out how a linear detector can be initialized using

the results obtained from this calibration. Carefully check the results

Other plot: bool

Parameters: flag to specify if a visualization of the fit should be done

usetilt: bool

whether to use model considering a detector tilt, i.e. deviation angle of the pixel direction from orthogonal to the primary beam (default: True)

Seealso

psd_chdeg

low level function with more configurable options

xrayutilities.analysis.sample_align.miscut_calc (phi, aomega, zeros=None, omega0=None,
plot=True)

function to calculate the miscut direction and miscut angle of a sample by fitting a sinusoidal function to the variation of the aligned omega values of more than two reflections. The function can also be used to fit reflectivity alignment values in various azimuths.

Parameters: phi : list, tuple or array-like

azimuths in which the reflection was aligned (deg)

aomega : list, tuple or array-like aligned omega values (deg)

zeros: list, tuple or array-like, optional

angles at which surface is parallel to the beam (deg). For the analysis the angles (aomega - zeros) are used.

omega0: float, optional

if specified the nominal value of the reflection is not included as fit parameter, but is fixed to the specified value. This value is MANDATORY if ONLY TWO AZIMUTHS are given.

plot: bool, optional

flag to specify if a visualization of the fit is wanted. default: True

Returns: omega0 : float

the omega value of the reflection should be close to the nominal one

phi0: float

the azimuth in which the primary beam looks upstairs

miscut: float

amplitude of the sinusoidal variation == miscut angle

xrayutilities.analysis.sample_align.**psd_chdeg** (angles, channels, stdev=None, usetilt=True, plot=True, datap='xk', modelline='--r', modeltilt='-b', fignum=None, mlabel='fit', mtiltlabel='fit w/tilt', dlabel='data', figtitle=True)

function to determine the channels per degree using a linear fit of the function nchannel = center_ch+chdeg*tan(angles) or the equivalent including a detector tilt

Parameters: angles: array-like

detector angles for which the position of the beam was measured

channels: array-like

detector channels where the beam was found

stdev : array-like, optional

standard deviation of the beam position

plot: bool, optional

flag to specify if a visualization of the fit should be done

usetilt: bool, optional

whether to use model considering a detector tilt, i.e. deviation angle of the pixel

direction from orthogonal to the primary beam (default: True)

Returns: pixelwidth: float

the width of one detector channel @ 1m distance, which is negative in case the hit

channel number decreases upon an increase of the detector angle.

centerch: float

center channel of the detector

tilt: float

tilt of the detector from perpendicular to the beam (will be zero in case of usetilt=False)

Note

L/pixelwidth*pi/180 = channel/degree for large detector distance with the sample detector disctance L

Other datap: str, optional

Parameters: plot format of data points

modelline: str, optional
plot format of modelline
modeltilt: str, optional
plot format of modeltilt
fignum: int or str, optional

figure number to use for the plot

mlabel: str

label of the model w/o tilt to be used in the plot

mtiltlabel: str

label of the model with tilt to be used in the plot

dlabel: str

label of the data line to be used in the plot

figtitle: bool

flag to tell if the figure title should show the fit parameters

xrayutilities.analysis.sample_align.psd_refl_align (primarybeam, angles, channels, plot=True) function which calculates the angle at which the sample is parallel to the beam from various angles and detector channels from the reflected beam. The function can be used during the half beam alignment with a linear detector.

Parameters: primarybeam: int

primary beam channel number

angles : list or array-like
incidence angles

channels: list or array-like

corresponding detector channels

plot: bool, optional

flag to specify if a visualization of the fit is wanted default : True

Returns: float

angle at which the sample is parallel to the beam

Examples

```
>>> psd_refl_align(500, [0, 0.1, 0.2, 0.3], [550, 600, 640, 700])
```

Module contents

xrayutilities.analysis is a package for assisting with the analysis of x-ray diffraction data, mainly reciprocal space maps

Routines for obtaining line cuts from gridded reciprocal space maps are offered, with the ability to integrate the intensity perpendicular to the line cut direction.

xrayutilities.io package

Submodules

xrayutilities.io.cbf module

class xrayutilities.io.cbf.CBFDirectory (datapath, ext='cbf', **keyargs)

Bases: xrayutilities.io.filedir.FileDirectory

Parses a directory for CBF files, which can be stored to a HDF5 file for further usage

class xrayutilities.io.cbf.CBFFile (fname, nxkey='X-Binary-Size-Fastest-Dimension',
nykey='X-Binary-Size-Second-Dimension', dtkey='DataType', path=None)

Bases: object

ReadData ()

Read the CCD data into the .data object this function is called by the initialization

```
Save2HDF5 (h5f, group='/', comp=True)
```

Saves the data stored in the EDF file in a HDF5 file as a HDF5 array. By default the data is stored in the root group of the HDF5 file - this can be changed by passing the name of a target group or a path to the target group via the "group" keyword argument.

Parameters: h5f: file-handle or str

a HDF5 file object or name

group: str, optional

group where to store the data (default to the root of the file)

comp: bool, optional

activate compression - true by default

xrayutilities.io.desy_tty08 module

class for reading data + header information from tty08 data files

tty08 is a system used at beamline P08 at Hasylab Hamburg and creates simple ASCII files to save the data. Information is easily read from the multicolumn data file. the functions below enable also to parse the information of the header

xrayutilities.io.desy_tty08.gettty08_scan (scanname, scannumbers, *args, **keyargs)

function to obtain the angular cooridinates as well as intensity values saved in TTY08 datafiles. Especially usefull for reciprocal space map measurements, and to combine date from several scans

further more it is possible to obtain even more positions from the data file if more than two string arguments with its names are given

Parameters: scanname: str

name of the scans, for multiple scans this needs to be a template string

scannumbers: int, tuple or list

number of the scans of the reciprocal space map

args: str, optional

names of the motors. to read reciprocal space maps measured in coplanar diffraction give:

- omname: the name of the omega motor (or its equivalent)
- ttname: the name of the two theta motor (or its equivalent)

keyargs: dict, optional

keyword arguments are passed on to tty08File

Returns: [ang1, ang2, ...]: list, optional

angular positions of the center channel of the position sensitive detector (numpy.ndarray 1D), omitted if no *arg*s are given

MAP: ndarray

All the data values as stored in the data file (includes the intensities e.g. MAP['MCA']).

Examples

```
>>> [om, tt], MAP = xu.io.gettty08_scan('text%05d.dat', 36, 'omega', 'gamma')
```

class xrayutilities.io.desy_tty08.tty08File (filename, path=None, mcadir=None)

Bases: object

Represents a tty08 data file. The file is read during the Constructor call. This class should work for data stored at beamline P08 using the tty08 acquisition system.

Parameters: filename: str

tty08-filename

mcadir: str, optional

directory name of MCA files

Read ()

Read the data from the file

ReadMCA ()

xrayutilities.io.edf module

```
class xrayutilities.io.edf.EDFDirectory (datapath, ext='edf', **keyargs)
```

Bases: xrayutilities.io.filedir.FileDirectory

Parses a directory for EDF files, which can be stored to a HDF5 file for further usage

class xrayutilities.io.edf.EDFFile (fname, nxkey='Dim_1', nykey='Dim_2', dtkey='DataType', path='',
header=True, keep_open=False)

Bases: object

Parse ()

Parse file to find the number of entries and read the respective header information

ReadData (nimg=0)

Read the CCD data of the specified image and return the data this function is called automatically when the 'data' property is accessed, but can also be called manually when only a certain image from the file is needed.

Parameters: nimg: int, optional

number of the image which should be read (starts with 0)

Save2HDF5 (h5f, group='/', comp=True)

Saves the data stored in the EDF file in a HDF5 file as a HDF5 array. By default the data is stored in the root group of the HDF5 file - this can be changed by passing the name of a target group or a path to the target group via the "group" keyword argument.

Parameters: h5f: file-handle or str

a HDF5 file object or name

group: str, optional

group where to store the data (default to the root of the file)

comp: bool, optional

activate compression - true by default

property data

xrayutilities.io.fastscan module

modules to help with the analysis of FastScan data acquired at the ESRF. FastScan data are X-ray data (various detectors possible) acquired during scanning the sample in real space with a Piezo Scanner. The same functions might be used to analze traditional SPEC mesh scans.

The module provides three core classes:

- FastScan
- FastScanCCD
- FastScanSeries

where the first two are able to parse single mesh/FastScans when one is interested in data of a single channel detector or are detector and the last one is able to parse full series of such mesh scans with either type of detector

see examples/xrayutilities_kmap_ESRF.py for an example script

```
class xrayutilities.io.fastscan.FastScan (filename, scannr, xmotor='adcX', ymotor='adcY', path=")
Bases: object
```

class to help parsing and treating fast scan data. FastScan is the aquisition of X-ray data while scanning the sample with piezo stages in real space. It's is available at several beamlines at the ESRF synchrotron light-source.

grid2D (nx, ny, **kwargs)

function to grid the counter data and return the gridded X, Y and Intensity values.

Parameters: nx, ny : int

number of bins in x, and y direction

counter: str, optional

name of the counter to use for gridding (default: 'mpx4int' (ID01))

gridrange: tuple, optional

range for the gridder: format: ((xmin, xmax), (ymin, ymax))

Returns: Gridder2D

Gridder2D object with X, Y, data on regular x, y-grid

motorposition (motorname)

read the position of motor with name given by motorname from the data file. In case the motor is included in the data columns the returned object is an array with all the values from the file (although retrace clean is respected if already performed). In the case the motor is not moved during the scan only one value is returned.

Parameters: motorname: str

name of the motor for which the position is wanted

Returns: ndarray

motor position(s) of motor with name motorname during the scan

parse ()

parse the specifie for the scan number specified in the constructor and store the needed informations in the object properties

retrace_clean()

function to clean the data of the scan from retrace artifacts created by the zig-zag scanning motion of the piezo actuators the function cleans the xvalues, yvalues and data attribute of the FastScan object.

class xrayutilities.io.fastscan.FastScanCCD (*args, **kwargs)

Bases: xrayutilities.io.fastscan.FastScan

class to help parsing and treating fast scan data including CCD frames. FastScan is the aquisition of X-ray data while scanning the sample with piezo stages in real space. It's is available at several beamlines at the ESRF synchrotron light-source. During such fast scan at every grid point CCD frames are recorded and need to be analyzed

getCCD (ccdnr, roi=None, datadir=None, keepdir=0, replacedir=None, nav=[1, 1], filterfunc=None)

function to read the ccd files and return the raw X, Y and DATA values. DATA represents a 3D object with first dimension representing the data point index and the remaining two dimensions representing detector channels

Parameters: ccdnr: array-like or str

array with ccd file numbers of length length(FastScanCCD.data) OR a string with the data column name for the file ccd-numbers

roi: tuple, optional

region of interest on the 2D detector. should be a list of lower and upper bounds of detector channels for the two pixel directions (default: None)

datadir: str, optional

the CCD filenames are usually parsed from the SPEC file. With this option the directory used for the data can be overwritten. Specify the datadir as simple string. Alternatively the innermost directory structure can be automatically taken from the specifile. If this is needed specify the number of directories which should be kept using the keepdir option.

keepdir: int, optional

number of directories which should be taken from the SPEC file. (default: 0)

replacedir: int, optional

number of outer most directory names which should be replaced in the output (default = None). One can either give keepdir, or replacedir, with replace taking preference if both are given.

nav: tuple or list, optional

number of detector pixel which will be averaged together (reduces the date size)

filterfunc : callable

function applied to the CCD-frames before any processing. this function should take a single argument which is the ccddata which need to be returned with the same shape! e.g. remove hot pixels, flat/darkfield correction

Returns: X, Y: ndarray

x, y-array (1D)

DATA: ndarray

3-dimensional data object

getccdFileTemplate (specscan, datadir=None, keepdir=0, replacedir=None)

function to extract the CCD file template string from the comment in the SPEC-file scan-header.

Parameters: specscan: SpecScan

spec-scan object from which header the CCD directory should be extracted

datadir: str, optional

the CCD filenames are usually parsed from the scan object. With this option the directory used for the data can be overwritten. Specify the datadir as simple string. Alternatively the innermost directory structure can be automatically taken from the specfile. If this is needed specify the number of directories which should be kept using the keepdir option.

keepdir: int, optional

number of directories which should be taken from the specscan. (default: 0)

replacedir: int, optional

number of outer most directory names which should be replaced in the output (default = None). One can either give keepdir, or replacedir, with replace taking preference if both are given.

Returns: fmtstr: str

format string for the CCD file name using one number to build the real file name

filenr: int

starting file number

gridCCD (nx, ny, ccdnr, roi=None, datadir=None, keepdir=0, replacedir=None, nav=[1, 1], gridrange=None, filterfunc=None)

function to grid the internal data and ccd files and return the gridded X, Y and DATA values. DATA represents a 4D object with first two dimensions representing X, Y and the remaining two dimensions representing detector channels

Parameters: nx, ny : int

number of bins in x, and y direction

ccdnr: array-like or str

array with ccd file numbers of length length(FastScanCCD.data) OR a string with the data column name for the file ccd-numbers

roi: tuple, optional

region of interest on the 2D detector. should be a list of lower and upper bounds of detector channels for the two pixel directions (default: None)

datadir: str, optional

the CCD filenames are usually parsed from the SPEC file. With this option the directory used for the data can be overwritten. Specify the datadir as simple string. Alternatively the innermost directory structure can be automatically taken from the specifile. If this is needed specify the number of directories which should be kept using the keepdir option.

keepdir: int, optional

number of directories which should be taken from the SPEC file. (default: 0)

replacedir: int, optional

number of outer most directory names which should be replaced in the output (default = None). One can either give keepdir, or replacedir, with replace taking preference if both are given.

nav: tuple or list, optional

number of detector pixel which will be averaged together (reduces the date size)

gridrange: tuple

range for the gridder: format: ((xmin, xmax), (ymin, ymax))

filterfunc: callable

function applied to the CCD-frames before any processing. this function should take a single argument which is the ccddata which need to be returned with the same shape! e.g. remove hot pixels, flat/darkfield correction

Returns: X, Y: ndarray

regular x, y-grid

DATA: ndarray

4-dimensional data object

processCCD (ccdnr, roi, datadir=None, keepdir=0, replacedir=None, filterfunc=None)

function to read a region of interest (ROI) from the ccd files and return the raw X, Y and intensity from ROI.

Parameters: ccdnr: array-like or str

array with ccd file numbers of length length(FastScanCCD.data) OR a string with the data column name for the file ccd-numbers

roi: tuple or list

region of interest on the 2D detector. Either a list of lower and upper bounds of detector channels for the two pixel directions as tuple or a list of mask arrays

datadir: str, optional

the CCD filenames are usually parsed from the SPEC file. With this option the directory used for the data can be overwritten. Specify the datadir as simple string. Alternatively the innermost directory structure can be automatically taken from the specifie. If this is needed specify the number of directories which should be kept using the keepdir option.

keepdir: int, optional

number of directories which should be taken from the SPEC file. (default: 0)

replacedir: int, optional

number of outer most directory names which should be replaced in the output (default = None). One can either give keepdir, or replacedir, with replace taking preference if both are given.

filterfunc: callable, optional

function applied to the CCD-frames before any processing. this function should take a single argument which is the ccddata which need to be returned with the same shape! e.g. remove hot pixels, flat/darkfield correction

Returns: X, Y, DATA: ndarray

x, y-array (1D) as well as 1-dimensional data object

class xrayutilities.io.fastscan.FastScanSeries (filenames, scannrs, nx, ny, *args, **kwargs)

Bases: object

class to help parsing and treating a series of fast scan data including CCD frames. FastScan is the aquisition of X-ray data while scanning the sample with piezo stages in real space. It's is available at several beamlines at the ESRF synchrotron light-source. During such fast scan at every grid point CCD frames are recorded and need to be analyzed.

For the series of FastScans we assume that they are measured at different goniometer angles and therefore transform the data to reciprocal space.

align (deltax, deltay)

Since a sample drift or shift due to rotation often occurs between different FastScans it should be corrected before combining them. Since determining such a shift is not straight-forward in general the user needs to supply the routine with the shifts in order correct the x, y-values for the different FastScans. Such a routine could for example use the integrated CCD intensities and determine the shift using a cross-convolution.

Parameters: deltax, deltay: list

list of shifts in x/y-direction for every FastScan in the data structure

getCCDFrames (posx, posy, typ='real')

function to determine the list of ccd-frame numbers for a specific real space position. The real space position must be within the data limits of the FastScanSeries otherwise an ValueError is thrown

Parameters: posx: float

real space x-position or index in x direction

posy: float

real space y-position or index in y direction

typ: {'real', 'index'}, optional

type of coordinates. specifies if the position is specified as real space coordinate or as

index. (default: 'real')

Returns: list

[[motorpos1, ccdnrs1], [motorpos2, ccdnrs2], ...] where motorposN is from the N-ths FastScan in the series and ccdnrsN is the list of according CCD-frames

get_average_RSM (qnx, qny, qnz, qconv, datadir=None, keepdir=0, replacedir=None, roi=None, nav=(1, 1), filterfunc=None)

function to return the reciprocal space map data averaged over all x, y positions from a series of FastScan measurements. It necessary to give the QConversion-object to be used for the reciprocal space conversion. The QConversion-object is expected to have the 'area' conversion routines configured properly. This function needs to read all detector images, so be prepared to lean back for a moment!

Parameters: qnx, qny, qnz : int

number of points used for the 3D Gridder

qconv: QConversion

QConversion-object to be used for the conversion of the CCD-data to reciprocal space

roi: tuple, optional

region of interest on the 2D detector. should be a list of lower and upper bounds of detector channels for the two pixel directions (default: None)

nav: tuple or list, optional

number of detector pixel which will be averaged together (reduces the date size)

filterfunc : callable, optional

function applied to the CCD-frames before any processing. this function should take a single argument which is the ccddata which need to be returned with the same shape! e.g. remove hot pixels, flat/darkfield correction

datadir: str, optional

the CCD filenames are usually parsed from the SPEC file. With this option the directory used for the data can be overwritten. Specify the datadir as simple string. Alternatively the innermost directory structure can be automatically taken from the specfile. If this is needed specify the number of directories which should be kept/replaced using the keepdir/replacedir option.

keepdir: int, optional

number of directories which should be taken from the SPEC file. (default: 0)

replacedir: int, optional

number of outer most directory names which should be replaced in the output (default = None). One can either give keepdir, or replacedir, with replace taking preference if both are given.

Returns: Gridder3D

gridded reciprocal space map

get_sxrd_for_grange (qrange, qconv, datadir=None, keepdir=0, replacedir=None, roi=None, nav=(1, 1),
filterfunc=None)

function to return the real space data averaged over a certain q-range from a series of FastScan measurements. It necessary to give the QConversion-object to be used for the reciprocal space conversion. The QConversion-object is expected to have the 'area' conversion routines configured properly.

Note

This function assumes that all FastScans were performed in the same real space positions, no gridding or aligning is performed!

Parameters: grange: list or tuple

q-limits defining a box in reciprocal space. six values are needed: [minx, maxx, miny,

..., maxz]

qconv: QConversion

QConversion object to be used for the conversion of the CCD-data to reciprocal

space

roi: tuple, optional

region of interest on the 2D detector. should be a list of lower and upper bounds of

detector channels for the two pixel directions (default: None)

nav: tuple or list, optional

number of detector pixel which will be averaged together (reduces the date size)

filterfunc : callable, optional

function applied to the CCD-frames before any processing. this function should take a single argument which is the ccddata which need to be returned with the same shape!

e.g. remove hot pixels, flat/darkfield correction

datadir: str, optional

the CCD filenames are usually parsed from the SPEC file. With this option the directory used for the data can be overwritten. Specify the datadir as simple string. Alternatively the innermost directory structure can be automatically taken from the specfile. If this is needed specify the number of directories which should be kept/replaced using the keepdir/replacedir option.

keepdir: int, optional

number of directories which should be taken from the SPEC file. (default: 0)

replacedir: int, optional

number of outer most directory names which should be replaced in the output (default = None). One can either give keepdir, or replacedir, with replace taking preference if

both are given.

Returns: xvalues, yvalues, data: ndarray

x, y, and data values

grid2Dall (nx, ny, **kwargs)

function to grid the counter data and return the gridded X, Y and Intensity values from all the FastScanSeries.

Parameters: nx, ny : int

number of bins in x, and y direction

counter: str, optional

name of the counter to use for gridding (default: 'mpx4int' (ID01))

gridrange: tuple, optional

range for the gridder: format: ((xmin, xmax), (ymin, ymax))

Returns: Gridder2D

object with X, Y, data on regular x, y-grid

gridrsm (posx, posy, qnx, qny, qnz, qconv, roi=None, nav=[1, 1], typ='real', filterfunc=None, **kwargs) function to calculate the reciprocal space map at a certain x, y-position from a series of FastScan measurements it is necessary to specify the number of grid-oints for the reciprocal space map and the QConversion-object to be used for the reciprocal space conversion. The QConversion-object is expected to have the 'area' conversion routines configured properly.

Parameters: posx: float

real space x-position or index in x direction

posy: float

real space y-position or index in y direction

qnx, qny, qnz: int

number of points in the Qx, Qy, Qz direction of the gridded reciprocal space map

qconv: QConversion

QConversion-object to be used for the conversion of the CCD-data to reciprocal

space

roi: tuple, optional

region of interest on the 2D detector. should be a list of lower and upper bounds of detector channels for the two pixel directions (default: None)

nav: tuple or list, optional

number of detector pixel which will be averaged together (reduces the date size)

typ: {'real', 'index'}, optional

type of coordinates. specifies if the position is specified as real space coordinate or as index. (default: 'real')

filterfunc: callable, optional

function applied to the CCD-frames before any processing, this function should take a single argument which is the ccddata which need to be returned with the same shape! e.g. remove hot pixels, flat/darkfield correction

UB: ndarray

sample orientation matrix

Returns: Gridder3D

object with gridded reciprocal space map

rawRSM (posx, posy, qconv, roi=None, nav=[1, 1], typ='real', datadir=None, keepdir=0, replacedir=None, filterfunc=None, **kwargs)

function to return the reciprocal space map data at a certain x, y-position from a series of FastScan measurements. It necessary to give the QConversion-object to be used for the reciprocal space conversion. The QConversion-object is expected to have the 'area' conversion routines configured properly.

Parameters: posx: float

real space x-position or index in x direction

posy: float

real space y-position or index in y direction

qconv : QConversion

QConversion-object to be used for the conversion of the CCD-data to reciprocal

space

roi: tuple, optional

region of interest on the 2D detector. should be a list of lower and upper bounds of detector channels for the two pixel directions (default: None)

nav: tuple or list, optional

number of detector pixel which will be averaged together (reduces the date size)

typ: {'real', 'index'}, optional

type of coordinates. specifies if the position is specified as real space coordinate or as index. (default: 'real')

filterfunc: callable, optional

function applied to the CCD-frames before any processing. this function should take a single argument which is the ccddata which need to be returned with the same shape! e.g. remove hot pixels, flat/darkfield correction

UB: array-like, optional

sample orientation matrix

datadir: str, optional

the CCD filenames are usually parsed from the SPEC file. With this option the directory used for the data can be overwritten. Specify the datadir as simple string. Alternatively the innermost directory structure can be automatically taken from the specifile. If this is needed specify the number of directories which should be kept using the keepdir option.

keepdir: int, optional

number of directories which should be taken from the SPEC file. (default: 0)

replacedir: int, optional

number of outer most directory names which should be replaced in the output (default = None). One can either give keepdir, or replacedir, with replace taking preference if both are given.

Returns: qx, qy, qz : ndarray

reciprocal space positions of the reciprocal space map

ccddata: ndarray

raw data of the reciprocal space map

valuelist : ndarray

valuelist containing the ccdframe numbers and corresponding motor positions

read_motors()

read motor values from the series of fast scans

retrace_clean()

perform retrace clean for every FastScan in the series

xrayutilities.io.filedir module

class xrayutilities.io.filedir.FileDirectory (datapath, ext, parser, **keyargs)

Bases: object

Parses a directory for files, which can be stored to a HDF5 file for further usage. The file parser is given to the constructor and must provide a Save2HDF5 method.

```
Save2HDF5 (h5f, group=", comp=True)
```

Saves the data stored in the found files in the specified directory in a HDF5 file as a HDF5 arrays in a subgroup. By default the data is stored in a group given by the foldername - this can be changed by passing the name of a target group or a path to the target group via the "group" keyword argument.

Parameters: h5f: file-handle or str

a HDF5 file object or name

group: str, optional

group where to store the data (defaults to pathname if group is empty string)

comp: bool, optional

activate compression - true by default

xrayutilities.io.helper module

convenience functions to open files for various data file reader

these functions should be used in new parsers since they transparently allow to open gzipped and bzipped files

class xrayutilities.io.helper.xu_h5open (f, mode='r')

Bases: object

helper object to decide if a HDF5 file has to be opened/closed when using with a 'with' statement.

xrayutilities.io.helper.xu_open (filename, mode='rb')

function to open a file no matter if zipped or not. Files with extension '.gz', '.bz2', and '.xz' are assumed to be compressed and transparently opened to read like usual files.

Parameters: filename: str

filename of the file to open (full including path)

mode: str, optional

mode in which the file should be opened

Returns: file-handle

handle of the opened file

Raises: IOError

If the file does not exist an IOError is raised by the open routine, which is not caught

within the function

xrayutilities.io.ill_numor module

module for reading ILL data files (station D23): numor files

class xrayutilities.io.ill_numor.numorFile (filename, path=None)

Bases: object

Represents a ILL data file (numor). The file is read during the Constructor call. This class should work for created at station D23 using the mad acquisition system.

Parameters: filename: str

a string with the name of the data file

Read ()

Read the data from the file

columns = {0: ('detector', 'monitor', 'time', 'gamma', 'omega', 'psi'), 1: ('detector', 'monitor', 'time', 'gamma'), 2: ('detector', 'monitor', 'time', 'omega'), 5: ('detector', 'monitor', 'time', 'psi')}

getline (fid)

ssplit (string)

multispace split. splits string at two or more spaces after stripping it.

```
xrayutilities.io.ill_numor.numor_scan (scannumbers, *args, **kwargs)
```

function to obtain the angular cooridinates as well as intensity values saved in numor datafiles. Especially useful for combining several scans into one data object.

Parameters: scannumbers: int or str or iterable

number of the numors, or list of numbers. This will be transformed to a string and used

as a filename

args: str, optional

names of the motors e.g.: 'omega', 'gamma'

kwargs: dict

keyword arguments are passed on to numorFile. e.g. 'path' for the files directory

Returns: [ang1, ang2, ...] : *list*

angular positions list, omitted if no args are given

data: ndarray

all the data values.

Examples

```
>>> [om, gam], data = xu.io.numor scan(414363, 'omega', 'gamma')
```

xrayutilities.io.imagereader module

class xrayutilities.io.imagereader.ImageReader (nop1, nop2, hdrlen=0, flatfield=None,
darkfield=None, dtype=<class 'numpy.int16'>, byte_swap=False)

Bases: object

parse CCD frames in the form of tiffs or binary data (*.bin) to numpy arrays. ignore the header since it seems to contain no useful data

The routine was tested so far with

- 1. RoperScientific files with 4096x4096 pixels created at Hasylab Hamburg, which save an 16bit integer per point.
- 2. Perkin Elmer images created at Hasylab Hamburg with 2048x2048 pixels.

readImage (filename, path=None)

read image file and correct for dark- and flatfield in case the necessary data are available. returned data = ((image data)-(darkfield))/flatfield*average(flatfield)

Parameters: filename: str

filename of the image to be read. so far only single filenames are supported. The data might be compressed. supported extensions: .tif, .bin and .bin.xz

path : str, optional
 path of the data files

class xrayutilities.io.imagereader.PerkinElmer (**keyargs)

Bases: xrayutilities.io.imagereader.ImageReader

parse PerkinElmer CCD frames (*.tif) to numpy arrays Ignore the header since it seems to contain no useful data

The routine was tested only for files with 2048x2048 pixel images created at Hasylab Hamburg which save an 32bit float per point.

class xrayutilities.io.imagereader.Pilatus100K (**keyargs)

Bases: xrayutilities.io.imagereader.ImageReader

parse Dectris Pilatus 100k frames (*.tiff) to numpy arrays Ignore the header since it seems to contain no useful data

class xrayutilities.io.imagereader.RoperCCD (**keyargs)

Bases: xrayutilities.io.imagereader.ImageReader

parse RoperScientific CCD frames (*.bin) to numpy arrays Ignore the header since it seems to contain no useful data

The routine was tested only for files with 4096x4096 pixel images created at Hasylab Hamburg which save an 16bit integer per point.

class xrayutilities.io.imagereader.TIFFRead (filename, path=None)

Bases: xrayutilities.io.imagereader.ImageReader

class to Parse a TIFF file including extraction of information from the file header in order to determine the image size and data type

The data stored in the image are available in the 'data' property.

xrayutilities.io.imagereader.get_tiff (filename, path=None)
read tiff image file and return the data

Parameters: filename: str

filename of the image to be read. so far only single filenames are supported. The data might be compressed.

path : str, optional
 path of the data file

xrayutilities.io.panalytical_xml module

Panalytical XML (www.XRDML.com) data file parser

based on the native python xml.dom.minidom module. want to keep the number of dependancies as small as possible

class xrayutilities.io.panalytical_xml.xmdLfile (fname, path=")

Bases: object

class to handle XRDML data files. The class is supplied with a file name and uses the XRDMLScan class to parse the xrdMeasurement in the file

class xrayutilities.io.panalytical_xml.XRDMLMeasurement (measurement, namespace=")

Bases: object

class to handle scans in a XRDML datafile

xrayutilities.io.panalytical_xml.getxrdml_map (filetemplate, scannrs=None, path='.', roi=None) parses multiple XRDML file and concatenates the results for parsing the xrayutilities.io.XRDMLFile class is used. The function can be used for parsing maps measured with the PIXCel 1D detector (and in limited way also for data acquired with a point detector -> see getxrdml_scan instead).

Parameters: filetemplate: str

template string for the file names, can contain a %d which is replaced by the scan number or be a list of filenames

scannrs: int or list, optional

scan number(s)

path: str, optional

common path to the filenames

roi: tuple, optional

region of interest for the PIXCel detector, for other measurements this is not useful!

Returns: om, tt, psd: ndarray

motor positions and data as flattened numpy arrays

Examples

```
>>> om, tt, psd = xrayutilities.io.getxrdml_map("samplename_%d.xrdml",
>>> [1, 2], path="./data")
```

xrayutilities.io.panalytical_xml.getxrdml_scan (filetemplate, *motors, **kwargs)

parses multiple XRDML file and concatenates the results for parsing the xrayutilities.io.XRDMLFile class is used. The function can be used for parsing arbitrary scans and will return the the motor values of the scan motor and additionally the positions of the motors given by in the *motors argument

Parameters: filetemplate: str

template string for the file names, can contain a %d which is replaced by the scan number or be a list of filenames given by the scannrs keyword argument

motors: str

motor names to return: e.g.: 'Omega', '2Theta', ... one can also use abbreviations:

- 'Omega' = 'om' = 'o'
- '2Theta' = 'tt' = 't'
- 'Chi' = 'c'
- 'Phi' = 'p'

scannrs: int or list, optional

scan number(s)

path: str, optional

common path to the filenames

Returns: scanmot, mot1, mot2,..., detectorint : ndarray

motor positions and data as flattened numpy arrays

Examples

xrayutilities.io.pdcif module

class xrayutilities.io.pdcif.pdCIF (filename, datacolumn=None)

Bases: object

the class implements a primitive parser for pdCIF-like files. It reads every entry and collects the information in the header attribute. The first loop containing one of the intensity fields is assumed to be the data the user is interested in and is transferred to the data array which is stored as numpy record array the columns can be accessed by name intensity fields:

- _pd_meas_counts_total
- _pd_meas_intensity_total
- _pd_proc_intensity_total
- _pd_proc_intensity_net
- _pd_calc_intensity_total
- pd calc intensity net

alternatively the data column name can be given as argument to the constructor

Parse ()

parser of the pdCIF file. the method reads the data from the file and fills the data and header attributes with content

class xrayutilities.io.pdcif.pdESG (filename, datacolumn=None)

Bases: xrayutilities.io.pdcif.pdCIF

class for parsing multiple pdCIF loops in one file. This includes especially *.esg files which are supposed to consist of multiple loops of pdCIF data with equal length.

Upon parsing the class tries to combine the data of these different scans into a single data matrix -> same shape of subscan data is assumed

Parse ()

parser of the pdCIF file. the method reads the data from the file and fills the data and header attributes with content

```
xrayutilities.io.pdcif.remove comments (line, sep='#')
```

xrayutilities.io.rigaku_ras module

class for reading data + header information from Rigaku RAS (3-column ASCII) files

Such datafiles are generated by the Smartlab Guidance software from Rigaku.

class xrayutilities.io.rigaku_ras.RASFile (filename, path=None)

Bases: object

Represents a RAS data file. The file is read during the constructor call

Parameters: filename: str

name of the ras-file

path: str, optional

path to the data file

Read ()

Read the data from the file

class xrayutilities.io.rigaku_ras.RASScan (filename, pos)

Bases: object

Represents a single Scan portion of a RAS data file. The scan is parsed during the constructor call

Parameters: filename: str

file name of the data file

pos: int

seek position of the 'RAS_HEADER_START' line

xrayutilities.io.rigaku_ras.getras_scan (scanname, scannumbers, *args, **kwargs)

function to obtain the angular cooridinates as well as intensity values saved in RAS datafiles. Especially useful for reciprocal space map measurements, and to combine date from several scans

further more it is possible to obtain even more positions from the data file if more than two string arguments with its names are given

Parameters: scanname: str

name of the scans, for multiple scans this needs to be a template string

scannumbers: int, tuple or list

number of the scans of the reciprocal space map

args: str, optional

names of the motors. to read reciprocal space maps measured in coplanar diffraction give:

- omname: name of the omega motor (or its equivalent)
- ttname: name of the two theta motor (or its equivalent)

kwarqs: dict

keyword arguments forwarded to RASFile function

Returns: [ang1, ang2, ...] : *list*

angular positions are extracted from the respective scan header, or motor positions during the scan. this is omitted if no *args* are given

rasdata : ndarray

the data values (includes the intensities e.g. rasdata['int']).

Examples

```
>>> [om, tt], MAP = xu.io.getras_scan('text%05d.ras', 36, 'Omega',
>>> 'TwoTheta')
```

xrayutilities.io.rotanode alignment module

parser for the alignment log file of the rotating anode

class xrayutilities.io.rotanode_alignment.RA_Alignment (filename)

Bases: object

class to parse the data file created by the alignment routine (tpalign) at the rotating anode spec installation this routine does an iterative alignment procedure and saves the center of mass values were it moves after each scan. It iterates between two different peaks and iteratively aligns at each peak between two different motors (om/chi at symmetric peaks, om/phi at asymmetric peaks)

Parse ()

parser to read the alignment log and obtain the aligned values at every iteration.

get (key)

keys ()

returns a list of keys for which aligned values were parsed

plot (pname)

function to plot the alignment history for a given peak

Parameters: pname: str

peakname for which the alignment should be plotted

xrayutilities.io.seifert module

a set of routines to convert Seifert ASCII files to HDF5 in fact there exist two posibilities how the data is stored (depending on the use detector):

- 1. as a simple line scan (using the point detector)
- 2. as a map using the PSD

In the first case the data ist stored

```
class xrayutilities.io.seifert.SeifertHeader
Bases: object
helper class to represent a Seifert (NJA) scan file header

class xrayutilities.io.seifert.SeifertMultiScan (filename, m_scan, m2, path=")
Bases: object
Class to parse a Seifert (NJA) multiscan file

parse ()

class xrayutilities.io.seifert.SeifertScan (filename, path=")
Bases: object
Class to parse a single Seifert (NJA) scan file
```

parse ()

xrayutilities.io.seifert.getSeifert_map (filetemplate, scannrs=None, path='.', scantype='map',
Nchannels=1280)

parses multiple Seifert *.nja files and concatenates the results. for parsing the xrayutilities.io.SeifertMultiScan class is used. The function can be used for parsing maps measured with the Meteor1D and point detector.

Parameters: filetemplate: str

template string for the file names, can contain a %d which is replaced by the scan

number or be a list of filenames

scannrs: int or list, optional

scan number(s)

path: str, optional

common path to the filenames

scantype: {'map', 'tsk'}, optional

type of datafile: can be either 'map' (reciprocal space map measured with a regular

Seifert job (default)) or 'tsk' (MCA spectra measured using the TaskInterpreter)

Nchannels: int, optional

number of channels of the MCA (needed for 'tsk' measurements)

Returns: om, tt, psd: ndarray

positions and data as flattened numpy arrays

Examples

```
>>> om, tt, psd = xrayutilities.io.getSeifert_map("samplename_%d.xrdml",
>>> [1, 2], path="./data")
```

```
xrayutilities.io.seifert.repair_key (key)
```

Repair a key string in the sense that the string is changed in a way that it can be used as a valid Python identifier. For that purpose all blanks within the string will be replaced by _ and leading numbers get an preceding _.

xrayutilities.io.spec module

a class for observing a SPEC data file

Motivation:

SPEC files can become quite large. Therefore, subsequently reading the entire file to extract a single scan is a quite cumbersome procedure. This module is a proof of concept code to write a file observer starting a reread of the file starting from a stored offset (last known scan position)

```
class xrayutilities.io.spec.specCmdLine (n, prompt, cmdl, out=")
Bases: object
```

class xrayutilities.io.spec.specfile (filename, path=")

Bases: object

This class represents a single SPEC file. The class provides methodes for updateing an already opened file which makes it particular interesting for interactive use.

Parse ()

Parses the file from the starting at last_offset and adding found scans to the scan list.

Save2HDF5 (h5f, comp=True, optattrs={})

Save the entire file in an HDF5 file. For that purpose a group is set up in the root group of the file with the name of the file without extension and leading path. If the method is called after an previous update only the scans not written to the file meanwhile are saved.

Parameters: h5f: file-handle or str

a HDF5 file object or its filename

comp: bool, optional

activate compression - true by default

Update ()

reread the file and add newly added files. The parsing starts at the data offset of the last scan gathered during the last parsing run.

class xrayutilities.io.spec.speclog (filename, prompt, path=")

Bases: object

class to parse a SPEC log file to find the command history

Parse ()

class xrayutilities.io.spec.specscan (name, scannr, command, date, time, itime, colnames, hoffset, doffset, fname, imopnames, imopvalues, scan_status)

Bases: object

Represents a single SPEC scan. This class is usually not called by the user directly but used via the SPECFile class.

ClearData ()

Delete the data stored in a scan after it is no longer used.

ReadData ()

Set the data attribute of the scan class.

Save2HDF5 (h5f, group='/', title=", optattrs={}, comp=True)

Save a SPEC scan to an HDF5 file. The method creates a group with the name of the scan and stores the data there as a table object with name "data". By default the scan group is created under the root group of the HDF5 file. The title of the scan group is ususally the scan command. Metadata of the scan are stored as attributes to the scan group. Additional custom attributes to the scan group can be passed as a dictionary via the optattrs keyword argument.

Parameters: h5f: file-handle or str

a HDF5 file object or its filename

group: str, optional

name or group object of the HDF5 group where to store the data

title: str, optional

a string with the title for the data, defaults to the name of scan if empty

optattrs: dict, optional

a dictionary with optional attributes to store for the data

comp: bool, optional

activate compression - true by default

SetMCAParams (mca_column_format, mca_channels, mca_start, mca_stop)

Set the parameters used to save the MCA data to the file. This method calculates the number of lines used to store the MCA data from the number of columns and the

Parameters: mca_column_format: int

number of columns used to save the data

mca_channels: int

number of MCA channels stored

mca start: int

first channel that is stored

mca_stop: int

last channel that is stored

getheader element (key, firstonly=True)

return the value-string of the first appearance of this SPECScan's header element, or a list of all values if firstonly=False

Parameters: specscan: SPECScan

key: str

name of the key to return; e.g. 'UMONO' or 'D'

firstonly: bool, optional

flag to specify if all instances or only the first one should be returned

Returns: valuestring: str

header value (if firstonly=True)

[str1, str2, ...]: list

header values (if firstonly=False)

plot (*args, **keyargs)

Plot scan data to a matplotlib figure. If newfig=True a new figure instance will be created. If logy=True (default is False) the y-axis will be plotted with a logarithmic scale.

Parameters: args: list

arguments for the plot: first argument is the name of x-value column the following pairs of arguments are the y-value names and plot styles allowed are 3, 5, 7,... number of arguments

keyargs: dict, optional

newfig: bool, optional

if True a new figure instance will be created otherwise an existing one will be used

logy: bool, optional

if True a semilogy plot will be done

xrayutilities.io.spec.geth5_scan (h5f, scans, *args, **kwargs)

function to obtain the angular cooridinates as well as intensity values saved in an HDF5 file, which was created from a spec file by the Save2HDF5 method. Especially useful for reciprocal space map measurements.

further more it is possible to obtain even more positions from the data file if more than two string arguments with its names are given

Parameters: h5f: file-handle or str

file object of a HDF5 file opened using h5py or its filename

scans: int, tuple or list

number of the scans of the reciprocal space map

args: str, optional

names of the motors. to read reciprocal space maps measured in coplanar diffraction give:

omname: name of the omega motor (or its equivalent)

ttname: name of the two theta motor (or its equivalent)

kwargs: dict, optional

samplename: str, optional

string with the hdf5-group containing the scan data if ommitted the first child node of h5f.root will be used

rettype: {'list', 'numpy'}, optional

how to return motor positions. by default a list of arrays is returned. when rettype == 'numpy' a record array will be returned.

Returns: [ang1, ang2, ...] : *list*

angular positions of the center channel of the position sensitive detector (numpy.ndarray 1D), this list is omitted if no *args* are given

MAP: ndarray

the data values as stored in the data file (includes the intensities e.g. MAP['MCA']).

Examples

```
>>> [om, tt], MAP = xu.io.geth5_scan(h5file, 36, 'omega', 'gamma')
```

xrayutilities.io.spec.getspec_scan (specf, scans, *args, **kwargs)

function to obtain the angular cooridinates as well as intensity values saved in a SPECFile. Especially useful to combine the data from multiple scans.

further more it is possible to obtain even more positions from the data file if more than two string arguments with its names are given

Parameters: specf : SPECFile

file object

scans: int, tuple or list number of the scans

args: str

names of the motors and counters

rettype: {'list', 'numpy'}, optional

how to return motor positions. by default a list of arrays is returned. when rettype ==

'numpy' a record array will be returned.

Returns: [ang1, ang2, ...] : *list*

coordinates and counters from the SPEC file

Examples

```
>>> [om, tt, cnt2] = xu.io.getspec_scan(s, 36, 'omega', 'gamma',
>>>
```

xrayutilities.io.spectra module

module to handle spectra data

class xrayutilities.io.spectra.SPECTRAFile (filename, mcatmp=None, mcastart=None, mcastop=None)

Bases: object

Represents a SPECTRA data file. The file is read during the Constructor call. This class should work for data stored at beamlines P08 and BW2 at HASYLAB.

Parameters: filename: str

a string with the name of the SPECTRA file

mcatmp: str, optional

template for the MCA files

mcastart, mcastop: int, optional

start and stop index for the MCA files, if not given, the class tries to determine the start and stop index automatically.

Read ()

Read the data from the file.

ReadMCA ()

Save2HDF5 (h5file, name, group='/', mcaname='MCA')

Saves the scan to an HDF5 file. The scan is saved to a seperate group of name "name". h5file is either a string for the file name or a HDF5 file object. If the mca attribute is not None mca data will be stored to an chunked array of with name mcaname.

Parameters: h5file: file-handle or str

HDF5 file object or name

name: str

name of the group where to store the data

group: str, optional

root group where to store the data

mcaname: str, optional

Name of the MCA in the HDF5 file

Returns: bool or None

The method returns None in the case of everything went fine, True otherwise.

class xrayutilities.io.spectra.SPECTRAFileComments

Bases: dict

Class that describes the comments in the header of a SPECTRA file. The different comments are accessible via the comment keys.

class xrayutilities.io.spectra.SPECTRAFileData

Bases: object

append (col)

class xrayutilities.io.spectra.SPECTRAFileDataColumn (index, name, unit, type)

Bases: object

<code>class xrayutilities.io.spectra.SPECTRAFileParameters () -> new empty dictionary dict(mapping) -> new dictionary initialized from a mapping object's (key, value) pairs dict(iterable) -> new dictionary initialized as if via: $d = \{\}$ for k, v in iterable: $d[k] = v \ dict(**kwargs) -> new \ dictionary initialized with the name=value pairs in the keyword argument list. For example: <math>dict(one=1, two=2)$ </code>

Bases: dict

xrayutilities.io.spectra.geth5_spectra_map (h5file, scans, *args, **kwargs)

function to obtain the omega and twotheta as well as intensity values for a reciprocal space map saved in an HDF5 file, which was created from a spectra file by the Save2HDF5 method.

further more it is possible to obtain even more positions from the data file if more than two string arguments with its names are given

Parameters: h5f: file-handle or str

file object of a HDF5 file opened using h5py

scans: int, tuple or list

number of the scans of the reciprocal space map

args: str, optional

arbitrary number of motor names

omname: name of the omega motor (or its equivalent)

• ttname: name of the two theta motor (or its equivalent)

kwargs: dict, optional

mca: str, optional

name of the mca data (if available) otherwise None (default: "MCA")

samplename: str. optional

string with the hdf5-group containing the scan data if omitted the first child node of

h5f.root will be used to determine the sample name

Returns: [ang1, ang2, ...] : *list*

angular positions of the center channel of the position sensitive detector

(numpy.ndarray 1D). one entry for every args-argument given to the function

MAP: ndarray

the data values as stored in the data file (includes the intensities e.g. MAP['MCA']).

Module contents

xrayutilities.materials package

Submodules

xrayutilities.materials.atom module

module containing the Atom class which handles the database access for atomic scattering factors and the atomic mass.

class xrayutilities.materials.atom.Atom (name, num)

Bases: object

property color

f (q, en='config')

function to calculate the atomic structure factor F

Parameters: q: float, array-like

momentum transfer

en: float or str, optional

energy for which F should be calculated, if omitted the value from the xrayutilities

configuration is used

Returns: float or array-like

value(s) of the atomic structure factor

f0 (q)

f1 (en='config')

```
£2 (en='config')
```

get_cache (prop, key)

check if a cached value exists to speed up repeated database requests

Returns: bool

True then result contains the cached otherwise False and result is None

result: database value

```
max_cache_length = 1000
```

property radius

set_cache (prop, key, result)

set result to be cached to speed up future calls

property weight

```
xrayutilities.materials.atom.get_key (*args)
```

generate a hash key for several possible types of arguments

xrayutilities.materials.cif module

```
class xrayutilities.materials.cif.CIFDataset (fid, name, digits)
```

Bases: object

class for parsing CIF (Crystallographic Information File) files. The class aims to provide an additional way of creating material classes instead of manual entering of the information the lattice constants and unit cell structure are parsed from the CIF file

Parse (fid)

function to parse a CIF data set. The function reads the space group symmetry operations and the basic atom positions as well as the lattice constants and unit cell angles

SGLattice (use_p1=False)

create a SGLattice object with the structure from the CIF file

SymStruct ()

function to obtain the list of different atom positions in the unit cell for the different types of atoms and determine the space group number and origin choice if available. The data are obtained from the data parsed from the CIF file.

```
class xrayutilities.materials.cif.CIFFile (filestr, digits=4)
```

Bases: object

class for parsing CIF (Crystallographic Information File) files. The class aims to provide an additional way of creating material classes instead of manual entering of the information the lattice constants and unit cell structure are parsed from the CIF file.

If multiple datasets are present in the CIF file this class will attempt to parse all of them into the the data dictionary. By default all methods access the first data set found in the file.

Parse ()

function to parse a CIF file. The function reads all the included data sets and adds them to the data dictionary.

```
SGLattice (dataset=None, use_p1=False)
```

create a SGLattice object with the structure from the CIF dataset

Parameters: dataset: str. optional

name of the dataset to use. if None the default one will be used.

use_p1: bool, optional

force the use of P1 symmetry, default False

xrayutilities.materials.cif.cifexport (filename, mat)

function to export a Crystal instance to CIF file. This in particular includes the atomic coordinates, however, ignores for example the elastic parameters.

xrayutilities.materials.database module

module to handle the access to the optical parameters database

class xrayutilities.materials.database.DataBase (fname)

Bases: object

Close ()

Close an opend database file.

Create (dbname, dbdesc)

Creates a new database. If the database file already exists its content is delete.

Parameters: dbname: str

name of the database

dbdesc: str

a short description of the database

CreateMaterial (name, description)

This method creates a new material. If the material group already exists the procedure is aborted.

Parameters: name: str

name of the material

description: str

description of the material

GetF0 (q, dset='default')

Obtain the f0 scattering factor component for a particular momentum transfer q.

Parameters: q: float or array-like

momentum transfer

dset: str, optional

specifies which dataset (different oxidation states) should be used

GetF1 (en)

Return the second, energy dependent, real part of the scattering factor for a certain energy en.

Parameters: en: float or array-like

energy

GetF2 (en)

Return the imaginary part of the scattering factor for a certain energy en.

Parameters: en: float or array-like

energy

Open (mode='r')

Open an existing database file.

SetColor (color)

Save color of the element for visualization

Parameters: color: tuple, str

matplotlib color for the element

setF0 (parameters, subset='default')

Save f0 fit parameters for the set material. The fit parameters are stored in the following order: c, a1, b1,......, a4, b4

Parameters: parameters: list or array-like

fit parameters **subset**: str, optional

name the f0 dataset

SetF1F2 (en, f1, f2)

Set f1, f2 values for the active material.

Parameters: en: list or array-like

energy in (eV)

f1 : list or array-like

f1 values

f2: list or array-like

f2 values

SetMaterial (name)

Set a particular material in the database as the actual material. All operations like setting and getting optical constants are done for this particular material.

Parameters: name: str

name of the material

SetRadius (radius)

Save atomic radius for visualization

Parameters: radius: float

atomic radius in Angstrom

SetWeight (weight)

Save weight of the element as float

Parameters: weight: float

atomic standard weight of the element

xrayutilities.materials.database.add_color_from_JMOL (db, cfile, verbose=False)
Read color from JMOL color table and save it to the database.

xrayutilities.materials.database.add_f0_from_intertab (db, itf, verbose=False)
Read f0 data from International Tables of Crystallography and add it to the database.

xrayutilities.materials.database.add_f0_from_xop (db, xop, verbose=False)
Read f0 data from f0 xop.dat and add it to the database.

xrayutilities.materials.database.add_f1f2_from_ascii_file (db, asciifile, element, verbose=False)
Read f1 and f2 data for specific element from ASCII file (3 columns) and save it to the database.

xrayutilities.materials.database.add_f1f2_from_henkedb (db, hf, verbose=False) Read f1 and f2 data from Henke database and add it to the database.

xrayutilities.materials.database.add_f1f2_from_kissel (db, kf, verbose=False)
Read f1 and f2 data from Henke database and add it to the database.

xrayutilities.materials.database.add mass from NIST (db, nistfile, verbose=False)

Read atoms standard mass and save it to the database. The mass of the natural isotope mixture is taken from the NIST data!

xrayutilities.materials.database.add_radius_from_wiki (db, dfile, verbose=False)
Read radius from Wikipedia radius table and save it to the database.

xrayutilities.materials.database.createAndFillDatabase (fname, dpath=None, verbose=False) function to create the database and fill it with values from the various source files.

Parameters: fname: str

Filename of the database to be created (including the path)

dpath: str, optional

directory where all the source data files are stored

verbose: bool, optional

flag to determine the verbosity of the script (default: False)

xrayutilities.materials.database.init_material_db (db)

xrayutilities.materials.elements module

xrayutilities.materials.heuslerlib module

implement convenience functions to define Heusler materials.

xrayutilities.materials.heuslerlib.**FullHeuslerCubic225** (X, Y, Z, a, biso=[0, 0, 0], occ=[1, 1, 1]) Full Heusler structure with formula X2YZ. Strukturberichte symbol L2_1; space group Fm-3m (225)

Parameters: X, Y, Z: str or Element

elements

a: float

cubic lattice parameter in Angstroem

biso: list of floats, optional

Debye Waller factors for X, Y, Z elements

occ: list of floats, optional

occupation numbers for the elements X, Y, Z

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.heuslerlib.**FullHeuslerCubic225_A2** (X, Y, Z, a, a2dis, biso=[0, 0, 0], occ=[1, 1, 1])

Full Heusler structure with formula X2YZ. Strukturberichte symbol L2_1; space group Fm-3m (225) with A2-type (W) disorder

Parameters: X, Y, Z: str or Element

elements

a: float

cubic lattice parameter in Angstroem

a2dis: float

amount of A2-type disorder (0: fully ordered, 1: fully disordered)

biso: list of floats, optional

Debye Waller factors for X, Y, Z elements

occ: list of floats, optional

occupation numbers for the elements X, Y, Z

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.heuslerlib.**FullHeuslerCubic225_B2** (X, Y, Z, a, b2dis, biso=[0, 0, 0], occ=[1, 1, 1])

Full Heusler structure with formula X2YZ. Strukturberichte symbol L2_1; space group Fm-3m (225) with B2-type (CsCl) disorder

Parameters: X, Y, Z: str or Element

elements

a: float

cubic lattice parameter in Angstroem

b2dis: float

amount of B2-type disorder (0: fully ordered, 1: fully disordered)

biso: list of floats, optional

Debye Waller factors for X, Y, Z elements

occ: list of floats, optional

occupation numbers for the elements X, Y, Z

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.heuslerlib.**FullHeuslerCubic225_DO3** (X, Y, Z, a, do3disxy, do3disxz, biso=[0, 0, 0], occ=[1, 1, 1])

Full Heusler structure with formula X2YZ. Strukturberichte symbol L2_1; space group Fm-3m (225) with DO_3-type (BiF3) disorder, either between atoms X <-> Y or X <-> Z.

Parameters: X, Y, Z: str or Element

elements

a: float

cubic lattice parameter in Angstroem

do3disxy: float

amount of DO_3-type disorder between X and Y atoms (0: fully ordered, 1: fully disordered)

do3disxz: float

amount of DO_3-type disorder between X and Z atoms (0: fully ordered, 1: fully disordered)

biso: list of floats, optional

Debye Waller factors for X, Y, Z elements

occ: list of floats, optional

occupation numbers for the elements X, Y, Z

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.heuslerlib.**HeuslerHexagonal194** (X, Y, Z, a, c, biso=[0, 0, 0], occ=[1, 1, 1]) Hexagonal Heusler structure with formula XYZ space group P63/mmc (194)

Parameters: X, Y, Z: str or Element

elements

a.c: float

hexagonal lattice parameters in Angstroem

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.heuslerlib.**HeuslerTetragonal119** (X, Y, Z, a, c, biso=[0, 0, 0], occ=[1, 1, 1]) Tetragonal Heusler structure with formula X2YZ space group I-4m2 (119)

Parameters: X, Y, Z: str or Element

elements **a, c**: float

tetragonal lattice parameters in Angstroem

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.heuslerlib.**HeuslerTetragonal139** (X, Y, Z, a, c, biso=[0, 0, 0], occ=[1, 1, 1]) Tetragonal Heusler structure with formula X2YZ space group I4/mmm (139)

Parameters: X, Y, Z: str or Element

elements

a, c: float

tetragonal lattice parameters in Angstroem

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.heuslerlib.**InverseHeuslerCubic216** (X, Y, Z, a, biso=[0, 0, 0], occ=[1, 1, 1])

Full Heusler structure with formula (XY)X'Z structure; space group F-43m (216)

Parameters: X, Y, Z: str or Element

elements

a: float

cubic lattice parameter in Angstroem

Returns: Crystal

Crystal describing the Heusler material

xrayutilities.materials.material module

Classes decribing materials. Materials are devided with respect to their crystalline state in either Amorphous or Crystal types. While for most materials their crystalline state is defined few materials are also included as amorphous which can be useful for calculation of their optical properties.

class xrayutilities.materials.material.Alloy (matA, matB, x)

Bases: xrayutilities.materials.material.Crystal

alloys two materials from the same crystal system. If the materials have the same space group the Wyckoff positions within the unit cell will also reflect the alloying.

RelaxationTriangle (hkl, sub, exp)

function which returns the relaxation triangle for a Alloy of given composition. Reciprocal space coordinates are calculated using the user-supplied experimental class

Parameters: hkl : list or array-like

Miller Indices **sub**: Crystal, or float

substrate material or lattice constant

exp: Experiment

object from which the Transformation object and ndir are needed

Returns: qy, qz : float

reciprocal space coordinates of the corners of the relaxation triangle

static check_compatibility (matA, matB)

static lattice_const_AB (latA, latB, x, name=")

method to calculated the interpolation of lattice parameters and unit cell angles of the Alloy. By default linear interpolation between the value of material A and B is performed.

Parameters: latA, latB: float or vector

property (lattice parameter/angle) of material A and B. A property can be a scalar or vector.

x:float

fraction of material B in the alloy.

name: str, optional

label of the property which is interpolated. Can be 'a', 'b', 'c', 'alpha', 'beta', or

'gamma'.

property x

class xrayutilities.materials.material.Amorphous (name, density, atoms=None, cij=None)

Bases: xrayutilities.materials.material.Material

amorphous materials are described by this class

chi0 (en='config')

calculates the complex chi_0 values often needed in simulations. They are closely related to delta and beta (n = 1 + $chi_0/2$ + i* $chi_0/2$ vs. n = 1 - delta + i*beta)

delta (en='config')

function to calculate the real part of the deviation of the refractive index from 1 (n=1-delta+i*beta)

Parameters: en: float, array-like or str, optional

energy of the x-rays in eV

Returns: float or array-like

ibeta (en='config')

function to calculate the imaginary part of the deviation of the refractive index from 1 (n=1-delta+i*beta)

Parameters: en: float, array-like or str, optional

energy of the x-rays in eV

Returns: float or array-like

static parseChemForm (cstring)

Parse a string containing a simple chemical formula and transform it to a list of elements together with their relative atomic fraction. e.g. 'H2O' -> [(H, 2/3), (O, 1/3)], where H and O are the Element objects of Hydrogen and Oxygen. Note that every chemical element needs to start with a capital letter! Complicated formulas containing bracket are not supported!

Parameters: cstring: str

string containing the chemical fomula

Returns: list of tuples

chemical element and atomic fraction

xrayutilities.materials.material.Cij2Cijkl(Cij)

Converts the elastic constants matrix (tensor of rank 2) to the full rank 4 cijkl tensor.

Parameters: cij: array-like

(6, 6) cij matrix

Returns: cijkl ndarray

(3, 3, 3, 3) cijkl tensor as numpy array

xrayutilities.materials.material.Cijkl2Cij (cijkl)

Converts the full rank 4 tensor of the elastic constants to the (6, 6) matrix of elastic constants.

Parameters: cijkl ndarray

(3, 3, 3, 3) cijkl tensor as numpy array

Returns: cij: array-like

(6, 6) cij matrix

class xrayutilities.materials.material.crystal (name, lat, cij=None, thetaDebye=None)

Bases: xrayutilities.materials.material.Material

Crystalline materials are described by this class

ApplyStrain (strain)

Applies a certain strain on the lattice of the material. The result is a change in the base vectors of the real space as well as reciprocal space lattice. The full strain matrix (3x3) needs to be given.

Note

NO elastic response of the material will be considered!

property B

GetMismatch (mat)

Calculate the mismatch strain between the material and a second material

HKL (*q)

Return the HKL-coordinates for a certain Q-space position.

Parameters: q: list or array-like

Q-position. its also possible to use HKL(qx, qy, qz).

Q (*hkl)

Return the Q-space position for a certain material.

Parameters: hkl: list or array-like

Miller indices (or Q(h, k, l) is also possible)

StructureFactor (q, en='config', temp=0)

calculates the structure factor of a material for a certain momentum transfer and energy at a certain temperature of the material

Parameters: q: list, tuple or array-like

vectorial momentum transfer

en: float or str, optional

x-ray energy eV, if omitted the value from the xrayutilities configuration is used

temp: float

temperature used for Debye-Waller-factor calculation

Returns: complex

the complex structure factor

StructureFactorForEnergy (q0, en, temp=0)

calculates the structure factor of a material for a certain momentum transfer and a bunch of energies

Parameters: q0: list, tuple or array-like

vectorial momentum transfer

en: list, tuple or array-like energy values in eV

temp: float

temperature used for Debye-Waller-factor calculation

Returns: array-like

complex valued structure factor array

StructureFactorForQ (q, en0='config', temp=0)

calculates the structure factor of a material for a bunch of momentum transfers and a certain energy

Parameters: q: list of vectors or array-like

> vectorial momentum transfers; list of vectores (list, tuple or array) of length 3 e.g.: (Si.Q(0, 0, 4), Si.Q(0, 0, 4.1),...) or numpy.array([Si.Q(0, 0, 4), Si.Q(0, 0, 4.1)])

en0: float or str, optional

x-ray energy eV, if omitted the value from the xrayutilities configuration is used

temp: float

temperature used for Debye-Waller-factor calculation

Returns: array-like

complex valued structure factor array

property a

property a1

property a2

property a3

property alpha

property b

property beta

property c

chemical_composition (natoms=None, with_spaces=False, ndigits=2) determine chemical composition from occupancy of atomic positions.

Parameters: mat : Crystal

> instance of Crystal natoms: int, optional

> > number of atoms to normalize the formula, if None some automatic normalization is attempted using the greatest common divisor of the number of atoms per unit cell. If the number of atoms of any element is fractional natoms=1 is used.

with_spaces: bool, optional

add spaces between the different entries in the output string for CIF combatibility

ndigits: int, optional

number of digits to which floating point numbers are rounded to

Returns: str

representation of the chemical composition

chi0 (en='config')

calculates the complex chi_0 values often needed in simulations. They are closely related to delta and beta ($n = 1 + chi_r0/2 + i^*chi_i0/2 \text{ vs. } n = 1 - delta + i^*beta$)

chih (q, en='config', temp=0, polarization='S')

calculates the complex polarizability of a material for a certain momentum transfer and energy

Parameters: q: list, tuple or array-like

momentum transfer vector in (1/A)

en: float or str, optional

x-ray energy eV, if omitted the value from the xrayutilities configuration is used

temp: float, optional

temperature used for Debye-Waller-factor calculation

polarization: {'S', 'P'}, optional sigma or pi polarization

Returns: tuple

(abs(chih_real), abs(chih_imag)) complex polarizability

dTheta (Q, en='config')

function to calculate the refractive peak shift

Parameters: Q: list, tuple or array-like

momentum transfer vector (1/A)

en: float or str, optional

x-ray energy eV, if omitted the value from the xrayutilities configuration is used

Returns: float

peak shift in degree

delta (en='config')

function to calculate the real part of the deviation of the refractive index from 1 (n=1-delta+i*beta)

Parameters: en: float or str, optional

x-ray energy eV, if omitted the value from the xrayutilities configuration is used

Returns: float

property density

calculates the mass density of an material from the mass of the atoms in the unit cell.

Returns: float

mass density in kg/m^3

distances ()

function to obtain distances of atoms in the crystal up to the unit cell size (largest value of a, b, c is the cut-off) returns a list of tuples with distance d and number of occurrence n [(d1, n1), (d2, n2),...]

Note

if the base of the material is empty the list will be empty

environment (*pos, **kwargs)

Returns a list of neighboring atoms for a given position within the unit cell. If the material does not contain any atoms a dummy atom will be placed on the unit cell corners.

Parameters: pos: list or array-like

fractional coordinate in the unit cell

maxdist: float

maximum distance wanted in the list of neighbors (default: 7)

Returns: list of tuples

(distance, atomType, multiplicity) giving distance sorted list of atoms

classmethod fromCIF (ciffilestr, **kwargs)

Create a Crystal from a CIF file. The default data-set from the cif file will be used to create the Crystal.

Parameters: ciffilestr: str, bytes

filename of the CIF file or string representation of the CIF file

kwargs: dict

keyword arguments are passed to the init-method of CIFFile

Returns: Crystal

property gamma

ibeta (en='config')

function to calculate the imaginary part of the deviation of the refractive index from 1 (n=1-delta+i*beta)

Parameters: en: float or str, optional

x-ray energy eV, if omitted the value from the xrayutilities configuration is used

Returns: float

loadLatticefromCIF (ciffilestr)

load the unit cell data (lattice) from the CIF file. Other material properties stay unchanged.

Parameters: ciffilestr: str, bytes

filename of the CIF file or string representation of the CIF file

planeDistance (*hkl)

determines the lattice plane spacing for the planes specified by (hkl)

Parameters: h, k, I: list, tuple or floats

Miller indices of the lattice planes given either as list, tuple or seperate arguments

Returns: float

the lattice plane spacing

Examples

```
>>> xu.materials.Si.planeDistance(0, 0, 4)
1.35776000000001
```

```
or
```

```
>>> xu.materials.Si.planeDistance((1, 1, 1))
3.1356124059796255
```

show_unitcell (fig=None, subplot=111, scale=0.6, complexity=11, linewidth=1.5, mode='matplotlib') visualization of the unit cell using either matplotlibs basic 3D functionality (expect rendering inaccuracies!) or the mayavi mlab package (accurate rendering -> recommended!)

Note

For more flexible visualization consider using the CIF-export feature and use a proper crystal structure viewer.

Parameters: fig: matplotlib Figure, Mayavi Scene, or None, optional

subplot: int or list, optional

subplot to use for the visualization when using matplotlib. This argument of fowarded to the first argument of matplotlibs *add_subplot* function

scale: float, optional

scale the size of the atoms by this additional factor. By default the size of the atoms corresponds to 60% of their atomic radius.

complexity: int, optional

number of steps to approximate the atoms as spheres. Higher values make spheres more accurate, but cause slower plotting.

linewidth: float, optional

line thickness of the unit cell outline

mode: str, optional

defines the plot backend used, can be 'matplotlib' (default) or 'mayavi'.

Returns: figure object of either matplotlib or Mayavi

toCIF (ciffilename)

Export the Crystal to a CIF file.

Parameters: ciffilename: str

filename of the CIF file

class xrayutilities.materials.material.CubicAlloy (matA, matB, x)

Bases: xrayutilities.materials.material.Alloy

ContentBasym (q_inp, q_perp, hkl, sur)

function that determines the content of B in the alloy from the reciprocal space position of an asymmetric peak.

Parameters: q_inp: float

inplane peak position of reflection hkl of the alloy in reciprocal space

q_perp : float

perpendicular peak position of the reflection hkl of the alloy in reciprocal space

hkl: list

Miller indices of the measured asymmetric reflection

sur : list

Miller indices of the surface (determines the perpendicular direction)

Returns: content: float

content of B in the alloy determined from the input variables

list

[a_inplane a_perp, a_bulk_perp(x), eps_inplane, eps_perp]; lattice parameters calculated from the reciprocal space positions as well as the strain (eps) of the layer

ContentBsym (q_perp, hkl, inpr, asub, relax)

function that determines the content of B in the alloy from the reciprocal space position of a symetric peak. As an additional input the substrates lattice parameter and the degree of relaxation must be given

Parameters: q_perp: float

perpendicular peak position of the reflection hkl of the alloy in reciprocal space

hkl: list

Miller indices of the measured symmetric reflection (also defines the surface normal

inpr: list

Miller indices of a Bragg peak defining the inplane reference direction

asub: float

substrate lattice parameter

relax: float

degree of relaxation (needed to obtain the content from symmetric reciprocal space

position)

Returns: content: float

the content of B in the alloy determined from the input variables

xrayutilities.materials.material.CubicElasticTensor (c11, c12, c44)

Assemble the 6x6 matrix of elastic constants for a cubic material from the three independent components of a cubic crystal

Parameters: c11, c12, c44 : float

independent components of the elastic tensor of cubic materials

Returns: cij: ndarray

6x6 matrix with elastic constants

xrayutilities.materials.material.HexagonalElasticTensor (c11, c12, c13, c33, c44)

Assemble the 6x6 matrix of elastic constants for a hexagonal material from the five independent components of a hexagonal crystal

Parameters: c11, c12, c13, c33, c44: float

independent components of the elastic tensor of a hexagonal material

Returns: cij: ndarray

6x6 matrix with elastic constants

class xrayutilities.materials.material.Material (name, cij=None)

Bases: abc.ABC

base class for all Materials. common properties of amorphous and crystalline materials are described by this class from which Amorphous and Crystal are derived from.

absorption_length (en='config')

wavelength dependent x-ray absorption length defined as mu = lambda/(2*pi*2*beta) with lambda and beta as the x-ray wavelength and complex part of the refractive index respectively.

Parameters: en: float or str, optional

energy of the x-rays in eV

Returns: float

the absorption length in um

chi0 (en='config')

calculates the complex chi_0 values often needed in simulations. They are closely related to delta and beta (n = 1 + chi r0/2 + i*chi i0/2 vs. n = 1 - delta + i*beta)

critical_angle (en='config', deg=True)

calculate critical angle for total external reflection

Parameters: en: float or str. optional

energy of the x-rays in eV, if omitted the value from the xrayutilities configuration is

used

deg: bool, optional

return angle in degree if True otherwise radians (default:True)

Returns: float

Angle of total external reflection

abstract delta (en='config')

abstract method which every implementation of a Material has to override

property density

abstract ibeta (en='config')

abstract method which every implementation of a Material has to override

idx_refraction (en='config')

function to calculate the complex index of refraction of a material in the x-ray range

Parameters: en: energy of the x-rays, if omitted the value from the

xrayutilities configuration is used

Returns: n (complex)

property lam

property mu

property nu

xrayutilities.materials.material.**PseudomorphicMaterial** (sub, layer, relaxation=0, trans=None)
This function returns a material whos lattice is pseudomorphic on a particular substrate material. The two materials must have similar unit cell definitions for the algorithm to work correctly, i.e. it does not work for combiniations of materials with different lattice symmetry. It is also crucial that the layer object includes values for the elastic tensor.

Parameters: sub: Crystal

substrate material

layer: Crystal

bulk material of the layer, including its elasticity tensor

relaxation: float, optional

degree of relaxation 0: pseudomorphic, 1: relaxed (default: 0)

trans: Tranform

Transformation which transforms lattice directions into a surface orientated coordinate frame (x, y inplane, z out of plane). If None a (001) surface geometry of a cubic material

is assumed.

Returns: An instance of Crystal holding the new pseudomorphically

strained material.

Raises: InputError

If the layer material has no elastic parameters

xrayutilities.materials.material.WZTensorFromCub (c11ZB, c12ZB, c44ZB)

Determines the hexagonal elastic tensor from the values of the cubic elastic tensor under the assumptions presented in Phys. Rev. B 6, 4546 (1972), which are valid for the WZ <-> ZB polymorphs.

Parameters: c11, c12, c44: float

independent components of the elastic tensor of cubic materials

Returns: cij: ndarray

6x6 matrix with elastic constants

Implementation according to a patch submitted by Julian Stangl

xrayutilities.materials.material.index_map_ij2ijkl(ij)
xrayutilities.materials.material.index_map_ijkl2ij(i,j)

xrayutilities.materials.plot module

xrayutilities.materials.plot.show_reciprocal_space_plane (mat, exp, ttmax=None, maxqout=0.01, scalef=100, ax=None, color=None, show_Laue=True, show_legend=True, projection='perpendicular', label=None) show a plot of the coplanar diffraction plane with peak positions for the respective material. the size of the spots is scaled with the strength of the structure factor

Parameters: mat: Crystal

instance of Crystal for structure factor calculations

exp: Experiment

instance of Experiment (needs to be HXRD, or FourC for onclick action to work correctly). defines the inplane and out of plane direction as well as the sample azimuth

ttmax: float, optional

maximal 2Theta angle to consider, by default 180deg

maxqout: float, optional

maximal out of plane q for plotted Bragg peaks as fraction of exp.k0

scalef: float, or callable, optional

scale factor or function for the marker size. If this is a function it should take only one float argument and return another float which is used as 's' parameter in matplotlib.pyplot.scatter

ax: matplotlib.Axes, optional

matplotlib Axes to use for the plot, useful if multiple materials should be plotted in one plot

color: matplotlib color, optional

show_Laue: bool, optional

flag to indicate if the Laue zones should be indicated

show_legend: bool, optional

flag to indiate if a legend should be shown

projection: 'perpendicular', 'polar', optional

type of projection for Bragg peaks which do not fall into the diffraction plane. 'perpendicular' (default) uses only the inplane component in the scattering plane, whereas 'polar' uses the vectorial absolute value of the two inplane components. See also the 'maxqout' option.

label: None or str, optional

label to be used for the legend. If 'None' the name of the material will be used.

Returns: Axes, plot_handle

xrayutilities.materials.predefined_materials module

```
class xrayutilities.materials.predefined_materials.AlGaAs (X)
```

Bases: xrayutilities.materials.material.CubicAlloy

class xrayutilities.materials.predefined_materials.SiGe (x)

Bases: xrayutilities.materials.material.CubicAlloy

static lattice_const_AB (latA, latB, x, **kwargs)

method to calculate the lattice parameter of the SiGe alloy with composition Si_{1-x}Ge_x

xrayutilities.materials.spacegrouplattice module

module handling crystal lattice structures. A SGLattice consists of a space group number and the position of atoms specified as Wyckoff positions along with their parameters. Depending on the space group symmetry only certain parameters of the resulting instance will be settable! A cubic lattice for example allows only to set its 'a' lattice parameter but none of the other unit cell shape parameters.

class xrayutilities.materials.spacegrouplattice.sGLattice (sgrp, *args, **kwargs)

Bases: object

lattice object created from the space group number and corresponding unit cell parameters. atoms in the unit cell are specified by their Wyckoff position and their free parameters.

ApplyStrain (eps)

Applies a certain strain on a lattice. The result is a change in the base vectors. The full strain matrix (3x3) needs to be given.

Note

Here you specify the strain and not the stress -> NO elastic response of the material will be considered!

Note

Although the symmetry of the crystal can be lowered by this operation the spacegroup remains unchanged! The 'free_parameters' attribute is, however, updated to mimic the possible reduction of the symmetry.

Parameters: eps: array-like

a 3x3 matrix with all strain components

GetHKL (*args)

determine the Miller indices of the given reciprocal lattice points

GetPoint (*args)

determine lattice points with indices given in the argument

Examples

```
>>> xu.materials.Si.lattice.GetPoint(0, 0, 4) array([ 0. , 0. , 21.72416])
```

```
or
>>> xu.materials.Si.lattice.GetPoint((1, 1, 1))
```

array([5.43104, 5.43104, 5.43104])

GetQ (*args)

determine the reciprocal lattice points with indices given in the argument

UnitCellVolume ()

function to calculate the unit cell volume of a lattice (angstrom^3)

property a

property alpha

property b

base ()

generator of atomic position within the unit cell.

property beta

property c

convert_to_P1()

create a P1 equivalent of this SGLattice instance.

Returns: SGLattice

instance with the same properties as the present lattice, however, in the P1 setting.

equivalent_hkls (hkl)

returns a list of equivalent hkl peaks depending on the crystal system

findsym()

method to return the highest symmetry description of the current material. This method does not consider to change the unit cell dimensions but only searches the highest symmetry spacegroup which with the current unit cell setting can be described. It is therefore not an implementation of FINDSYM [1].

Returns: new SGLattice-instance

a new SGLattice instance is returned with the highest available symmetry description. (see restrictions above)

[1] https://stokes.byu.edu/iso/findsym.php

property gamma

get_allowed_hkl (qmax)

return a set of all allowed reflections up to a maximal specified momentum transfer.

Parameters: qmax: float

maximal momentum transfer

Returns: hklset: set

set of allowed hkl reflections

hkl_allowed (hkl, returnequivalents=False)

check if Bragg reflection with Miller indices hkl can exist according to the reflection conditions. If no reflection conditions are available this function returns True for all hkl values!

Parameters: hkl: tuple or list

Miller indices of the reflection to check

returnequivalents: bool, optional

If True all the equivalent Miller indices of hkl are returned in a set as second return

argument.

Returns: allowed: bool

True if reflection can have non-zero structure factor, false otherwise

equivalents: set, optional

set of equivalent Miller indices if returnequivalents is True

property iscentrosymmetric

returns a boolean to determine if the lattice has centrosymmetry.

isequivalent (hkl1, hkl2)

determining if hkl1 and hkl2 are two crystallographical equivalent pairs of Miller indices. Note that this function considers the effect of non-centrosymmetry!

Parameters: hkl1, hkl2: list

Miller indices to be checked for equivalence

Returns: bool

reflection_conditions()

return string of reflection conditions, both general (from space group) and of Wyckoff positions

property symops

return the set of symmetry operations from the general Wyckoff position of the space group.

transform (mat, origin)

Transform the unit cell with the matrix and origin shift given in the parameters. This function returns a new instance of SGLattice which contains the highest possible symmetry description of the transformed unit cell. After the transformation (see [1]) the findsym method is used to create the new SGLattice instance.

Parameters: mat: (3, 3) list, or ndarray, optional

transformation matrix of the unit cell. The matrix definition aims to be consistent with what is used on the Bilbao Crystallographic Server [1]. This only defines the linear part, while the origin shift is given by origin.

origin : (3,) list, or ndarray
origin shift of the unit cell [1].

[1] https://www.cryst.ehu.es/cgi-bin/cryst/programs/nph-doc-trmat

class xrayutilities.materials.spacegrouplattice.symOp (D, t, m=1)

Bases: object

Class descriping a symmetry operation in a crystal. The symmetry operation is characterized by a 3x3 transformation matrix as well as a 3-vector describing a translation. For magnetic symmetry operations also the time reversal symmetry can be specified (not used in xrayutilities)

property D

transformation matrix of the symmetry operation

```
apply (vec, foldback=True)
apply_axial (vec)
apply_rotation (vec)
combine (other)
static foldback (v)
```

classmethod from_xyz (XYZ)

create a SymOp from the xyz notation typically used in CIF files.

Parameters: xyz: str

string describing the symmetry operation (e.g. '-y, -x, z')

property t

translation vector of the symmetry operation

```
xyz (showtimerev=False)
```

return the symmetry operation in xyz notation

class xrayutilities.materials.spacegrouplattice.WyckoffBase (*args, **kwargs)

Bases: list

The WyckoffBase class implements a container for a set of Wyckoff positions that form the base of a crystal lattice. An instance of this class can be treated as a simple container object.

append (atom, pos, occ=1.0, b=0.0) add new Atom to the lattice base

Parameters: atom: Atom

object to be added

pos: tuple or str

Wyckoff position of the atom, along with its parameters. Examples: ('2i', (0.1, 0.2,

0.3)), or '1a'

occ: float, optional

occupancy (default=1.0)

b: float, optional

b-factor of the atom used as exp(-b*q**2/(4*pi)**2) to reduce the intensity of this atom

(only used in case of temp=0 in StructureFactor and chi calculation)

static entry_eq (e1, e2)

compare two entries including all its properties to be equal

Parameters: e1, e2: tuple

tuples with length 4 containing the entries of WyckoffBase which should be compared

index (item)

return the index of the atom (same element, position, and Debye Waller factor). The occupancy is not checked intentionally. If the item is not present a ValueError is raised.

Parameters: item: tuple or list

WyckoffBase entry

Returns: int

static pos_eq (pos1, pos2) compare Wyckoff positions

Parameters: pos1, pos2: tuple

tuples with Wyckoff label and optional parameters

xrayutilities.materials.spacegrouplattice.get_default_sgrp_suf (sgrp_nr)
 determine default space group suffix

xrayutilities.materials.spacegrouplattice.get_possible_sgrp_suf (sgrp_nr)

determine possible space group suffix. Multiple suffixes might be possible for one space group due to different origin choice, unique axis, or choice of the unit cell shape.

Parameters: sgrp_nr: int

space group number

Returns: str or list

either an empty string or a list of possible valid suffix strings

xrayutilities.materials.spacegrouplattice.get_wyckpos (sgrp, atompos) test all Wyckoff positions on every atomic position

Parameters: sgrp: str

space group name

atompos: list

list of atomic positions to identify. All atomic positions are expected to belong to one and

the same Wyckoff position!

Returns: position argument for WyckoffBase.append

xrayutilities.materials.spacegrouplattice.testwp (parint, wp, cifpos, digits=8) test if a Wyckoff position can describe the given position from a CIF file

Parameters: parint : int

telling which Parameters the given Wyckoff position has

wp: str or tuple

expression of the Wyckoff position

cifpos: list, or tuple or array-like

(x, y, z) position of the atom in the CIF file

digits: int

number of digits for which for a comparison of floating point numbers will be rounded to.

By default xu.config.DIGITS is used.

Returns: foundflag: bool

flag to tell if the positions match

pars: array-like or None

parameters associated with the position or None if no parameters are needed

xrayutilities.materials.wyckpos module

class xrayutilities.materials.wyckpos.RangeDict() -> new empty dictionary dict(mapping)
-> new dictionary initialized from a mapping object's (key, value) pairs dict(iterable)
-> new dictionary initialized as if via: $d = \{\}$ for k, v in iterable: d[k] = vdict(**kwargs) -> new dictionary initialized with the name=value pairs in the keyword
argument list. For example: dict(one=1, two=2)

Bases: dict

Module contents

xrayutilities.math package

Submodules

xrayutilities.math.algebra module

module providing analytic algebraic functions not implemented in scipy or any other dependency of xrayutilities. In particular the analytic solution of a quartic equation which is needed for the solution of the dynamic scattering equations.

```
xrayutilities.math.algebra.solve_quartic (a4, a3, a2, a1, a0) analytic solution [1] of the general quartic equation. The solved equation takes the form a \cdot z^4 + a \cdot z^3 + a \cdot z^2 + a \cdot z + a \cdot z
```

Returns: tuple

tuple of the four (complex) solutions of aboves equation.

References

http://mathworld.wolfram.com/QuarticEquation.html

1

xrayutilities.math.fit module

module with a function wrapper to scipy.optimize.leastsq for fitting of a 2D function to a peak or a 1D Gauss fit with the odr package

xrayutilities.math.fit.**fit_peak2d** (x, y, data, start, drange, fit_function, maxfev=2000) fit a two dimensional function to a two dimensional data set e.g. a reciprocal space map

Parameters: x, y : array-like

data coordinates (do NOT need to be regularly spaced)

data: array-like

data set used for fitting (e.g. intensity at the data coords)

start : list

set of starting parameters for the fit used as first parameter of function fit_function

drange: list

limits for the data ranges used in the fitting algorithm, e.g. it is clever to use only a small region around the peak which should be fitted: [xmin, xmax, ymin, ymax]

fit_function: callable

function which should be fitted, must be of form accept the parameters $fit_function(x, y, *params) \rightarrow ndarray$

Returns: fitparam: list

fitted parameters

cov: array-like

covariance matrix

xrayutilities.math.fit.gauss_fit (xdata, ydata, iparams=[], maxit=300)

Gauss fit function using odr-pack wrapper in scipy similar to https://github.com/tiagopereira/python_tips/wiki/Scipy%3A-curve-fitting

Parameters: xdata: array-like

x-coordinates of the data to be fitted

ydata: array-like

y-coordinates of the data which should be fit

iparams: list, optional

initial paramters for the fit, determined automatically if not given

maxit: int, optional

maximal iteration number of the fit

Returns: params: list

the parameters as defined in function Gauss1d(x, *param)

sd_params: list

For every parameter the corresponding errors are returned.

itlim: bool

flag to tell if the iteration limit was reached, should be False

xrayutilities.math.fit.linregress(x, y)

fast linregress to avoid usage of scipy.stats which is slow! NaN values in y are ignored by this function.

Parameters: x, y : array-like

data coordinates and values

Returns: p:tuple

parameters of the linear fit (slope, offset)

rsq: float R^2 value

Examples

```
>>> (k, d), R2 = xu.math.linregress(x, y)
```

xrayutilities.math.fit.multPeakFit (x, data, peakpos, peakwidth, dranges=None, peaktype='Gaussian',
returnerror=False)

function to fit multiple Gaussian/Lorentzian peaks with linear background to a set of data

Parameters: x: array-like

x-coordinate of the data

data: array-like

data array with same length as x

peakpos: list

initial parameters for the peak positions

peakwidth: list

initial values for the peak width

dranges: list of tuples

list of tuples with (min, max) value of the data ranges to use. does not need to have the

same number of entries as peakpos

peaktype : {'Gaussian', 'Lorentzian'}

type of peaks to be used

returnerror: bool

decides if the fit errors of pos, sigma, and amp are returned (default: False)

Returns: pos: list

peak positions derived by the fit

sigma: list

peak width derived by the fit

amp: list

amplitudes of the peaks derived by the fit

background: array-like

background values at positions x

if returnerror == True:

sd pos: list

standard error of peak positions as returned by scipy.odr.Output

sd_sigma: list

standard error of the peak width

sd_amp: list

standard error of the peak amplitude

xrayutilities.math.fit.multPeakPlot (x, fpos, fwidth, famp, background, dranges=None,
peaktype='Gaussian', fig='xu_plot', ax=None, fact=1.0)

function to plot multiple Gaussian/Lorentz peaks with background values given by an array

Parameters: x: array-like

x-coordinate of the data

fpos: list

positions of the peaks

fwidth: list

width of the peaks

famp: list

amplitudes of the peaks

background: array-like

background values, same shape as x

dranges: list of tuples

list of (min, max) values of the data ranges to use. does not need to have the same number of entries as fpos

peaktype : {'Gaussian', 'Lorentzian'}

type of peaks to be used

fig: int, str, or None

matplotlib figure number or name

ax: matplotlib.Axes

matplotlib axes as alternative to the figure name

fact : float

factor to use as multiplicator in the plot

xrayutilities.math.fit.peak_fit (xdata, ydata, iparams=[], peaktype='Gauss', maxit=300,

background='constant', plot=False, func_out=False, debug=False)

fit function using odr-pack wrapper in scipy similar to https://github.com/tiagopereira/python_tips/wiki/Scipy%3A-curve-fitting for Gauss, Lorentz or Pseudovoigt-functions

Parameters: xdata: array_like

x-coordinates of the data to be fitted

ydata: array_like

y-coordinates of the data which should be fit

iparams: list, optional

initial paramters, determined automatically if not specified

peaktype : {'Gauss', 'Lorentz', 'PseudoVoigt', 'PseudoVoigtAsym', 'PseudoVoigtAsym2'},

optional

type of peak to fit

maxit: int, optional

maximal iteration number of the fit

background: {'constant', 'linear'}, optional

type of background function

plot: bool or str, optional

flag to ask for a plot to visually judge the fit. If plot is a string it will be used as figure name, which makes reusing the figures easier.

func out: bool, optional

returns the fitted function, which takes the independent variables as only argument (f(x))

Returns: params: list

the parameters as defined in function *Gauss1d/Lorentz1d/PseudoVoigt1d/PseudoVoigt1dasym*. In the case of linear background one more parameter is included!

sd params: list

For every parameter the corresponding errors are returned.

itlim: bool

flag to tell if the iteration limit was reached, should be False

fitfunc: function, optional

the function used in the fit can be returned (see func_out).

xrayutilities.math.functions module

module with several common function needed in xray data analysis

xrayutilities.math.functions.Debyel(x)

function to calculate the first Debye function [1] as needed for the calculation of the thermal Debye-Waller-factor by numerical integration

 $D_1(x) = (1/x) \int_0^x t/(\exp(t)-1) dt$

Parameters: x: float

argument of the Debye function

Returns: float

D1(x) float value of the Debye function

References

1 http://en.wikipedia.org/wiki/Debye_function

xrayutilities.math.functions.Gauss1d(x, *p)

function to calculate a general one dimensional Gaussian

Parameters: x: array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the Gaussian [XCEN, SIGMA, AMP, BACKGROUND] for information: SIGMA = FWHM / (2*sqrt(2*log(2)))

Returns: array-like

the value of the Gaussian described by the parameters p at position x

Examples

Calling with a list of parameters needs a call looking as shown below (note the '*') or explicit listing of the parameters

```
>>> Gauss1d(x,*p)
```

```
>>> Gauss1d(numpy.linspace(0, 10, 100), 5, 1, 1e3, 0)
```

```
xrayutilities.math.functions.Gauss1dArea (*p)
```

function to calculate the area of a Gauss function with neglected background

Parameters: p: list

list of parameters of the Gauss-function [XCEN, SIGMA, AMP, BACKGROUND]

Returns: float

the area of the Gaussian described by the parameters *p*

```
xrayutilities.math.functions.Gauss1d_der_p(x, *p)
```

function to calculate the derivative of a Gaussian with respect the parameters p for parameter description see Gauss1d

xrayutilities.math.functions.Gauss1d_der_x (x, *p)

function to calculate the derivative of a Gaussian with respect to x for parameter description see Gauss1d xrayutilities.math.functions.Gauss2d(x, y, *p) function to calculate a general two dimensional Gaussian Parameters: x, y: array-like coordinate(s) where the function should be evaluated p: list list of parameters of the Gauss-function [XCEN, YCEN, SIGMAX, SIGMAY, AMP, BACKGROUND, ANGLE]; SIGMA = FWHM / (2*sqrt(2*log(2))); ANGLE = rotation of the X, Y direction of the Gaussian in radians array-like Returns: the value of the Gaussian described by the parameters p at position (x, y) xrayutilities.math.functions.Gauss2dArea (*p) function to calculate the area of a 2D Gauss function with neglected background Parameters: p: list list of parameters of the Gauss-function [XCEN, YCEN, SIGMAX, SIGMAY, AMP, ANGLE, BACKGROUND] Returns: float the area of the Gaussian described by the parameters p xrayutilities.math.functions.Gauss3d(x, y, z, *p) function to calculate a general three dimensional Gaussian Parameters: x, y, z : array-like coordinate(s) where the function should be evaluated p: list list of parameters of the Gauss-function [XCEN, YCEN, ZCEN, SIGMAX, SIGMAY, SIGMAZ, AMP, BACKGROUND]; SIGMA = FWHM / (2*sqrt(2*log(2)))Returns: array-like the value of the Gaussian described by the parameters p at positions (x, y, z) xrayutilities.math.functions.Lorentz1d(x, *p) function to calculate a general one dimensional Lorentzian Parameters: x : array-like coordinate(s) where the function should be evaluated p: list list of parameters of the Lorentz-function [XCEN, FWHM, AMP, BACKGROUND] Returns: array-like the value of the Lorentian described by the parameters p at position (x, y) xrayutilities.math.functions.LorentzldArea (*p) function to calculate the area of a Lorentz function with neglected background Parameters: **p**: list list of parameters of the Lorentz-function [XCEN, FWHM, AMP, BACKGROUND] Returns: float the area of the Lorentzian described by the parameters p xrayutilities.math.functions.Lorentz1d_der_p(x, *p) function to calculate the derivative of a Gaussian with respect the parameters p

for parameter description see Lorentz1d

xrayutilities.math.functions.Lorentz1d_der_x(x, *p)

function to calculate the derivative of a Gaussian with respect to x for parameter description see Lorentz1d

xrayutilities.math.functions.Lorentz2d(x, y, *p)

function to calculate a general two dimensional Lorentzian

Parameters: x, y: array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the Lorentz-function [XCEN, YCEN, FWHMX, FWHMY, AMP, BACKGROUND, ANGLE]; ANGLE = rotation of the X, Y direction of the Lorentzian in

radians

Returns: array-like

the value of the Lorentian described by the parameters p at position (x, y)

xrayutilities.math.functions.NormGauss1d(x, *p)

function to calculate a normalized one dimensional Gaussian

Parameters: x: array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the Gaussian [XCEN, SIGMA]; for information: SIGMA = FWHM /

(2*sqrt(2*log(2)))

Returns: array-like

the value of the normalized Gaussian described by the parameters p at position x

xrayutilities.math.functions.NormLorentz1d(x, *p)

function to calculate a normalized one dimensional Lorentzian

Parameters: x: array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the Lorentzian [XCEN, FWHM]

Returns: array-like

the value of the normalized Lorentzian described by the parameters p at position x

xrayutilities.math.functions.PseudoVoigt1d(x, *p)

function to calculate a pseudo Voigt function as linear combination of a Gauss and Lorentz peak

Parameters: x: array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the pseudo Voigt-function [XCEN, FWHM, AMP, BACKGROUND,

ETA]; ETA: 0 ...1 0 means pure Gauss and 1 means pure Lorentz

Returns: array-like

the value of the PseudoVoigt described by the parameters p at position x

xrayutilities.math.functions.PseudoVoigt1dArea (*p)

function to calculate the area of a pseudo Voigt function with neglected background

Parameters: p: list

list of parameters of the Lorentz-function [XCEN, FWHM, AMP, BACKGROUND, ETA];

ETA: 0 ... 1 0 means pure Gauss and 1 means pure Lorentz

Returns: float

the area of the PseudoVoigt described by the parameters p

xrayutilities.math.functions.**PseudoVoigt1d_der_p**(x, *p)

function to calculate the derivative of a PseudoVoigt with respect the parameters p for parameter description see PseudoVoigt1d

 ${\tt xrayutilities.math.functions.} \textbf{PseudoVoigt1d_der_x} \ (x, \ ^*p)$

function to calculate the derivative of a PseudoVoigt with respect to x

for parameter description see PseudoVoigt1d

xrayutilities.math.functions.PseudoVoigtldasym(x, *p)

function to calculate an asymmetric pseudo Voigt function as linear combination of asymmetric Gauss and Lorentz peak

Parameters: x: array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the pseudo Voigt-function [XCEN, FWHMLEFT, FWHMRIGHT, AMP, BACKGROUND, ETA]; ETA: 0 ...1 0 means pure Gauss and 1 means pure

Lorentz

Returns: array-like

the value of the PseudoVoigt described by the parameters p at position x

xrayutilities.math.functions.**PseudoVoigt1dasym2** (x, *p)

function to calculate an asymmetric pseudo Voigt function as linear combination of asymmetric Gauss and Lorentz peak

Parameters: x : naddray

coordinate(s) where the function should be evaluated

p: list

list of parameters of the pseudo Voigt-function [XCEN, FWHMLEFT, FWHMRIGHT, AMP, BACKGROUND, ETALEFT, ETARIGHT]; ETA: 0 ...1 0 means pure Gauss and 1 means pure Lorentz

Returns: array-like

the value of the PseudoVoigt described by the parameters p at position x

xrayutilities.math.functions.**PseudoVoigt2d**(x, y, *p)

function to calculate a pseudo Voigt function as linear combination of a Gauss and Lorentz peak in two dimensions

Parameters: x, y : array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the pseudo Voigt-function [XCEN, YCEN, FWHMX, FWHMY, AMP, BACKGROUND, ANGLE, ETA]; ETA: 0 ...1 0 means pure Gauss and 1 means pure

Lorentz

Returns: array-like

the value of the PseudoVoigt described by the parameters p at position (x, y)

xrayutilities.math.functions.**TwoGauss2d**(x, y, *p)

function to calculate two general two dimensional Gaussians

Parameters: x, y : array-like

coordinate(s) where the function should be evaluated

p: list

list of parameters of the Gauss-function [XCEN1, YCEN1, SIGMAX1, SIGMAY1, AMP1, ANGLE1, XCEN2, YCEN2, SIGMAX2, SIGMAY2, AMP2, ANGLE2, BACKGROUND]; SIGMA = FWHM / (2*sqrt(2*log(2))) ANGLE = rotation of the X, Y direction of the

Gaussian in radians

Returns: array-like

the value of the Gaussian described by the parameters p at position (x, y)

xrayutilities.math.functions.heaviside(x)

Heaviside step function for numpy arrays

Parameters: x: scalar or array-like

argument of the step function

Returns: int or array-like

Heaviside step function evaluated for all values of x with datatype integer

xrayutilities.math.functions.kill_spike (data, threshold=2.0, offset=None)

function to smooth **single** data points which differ from the average of the neighboring data points by more than the threshold factor or more than the offset value. Such spikes will be replaced by the mean value of the next neighbors.

Warning

Use this function carefully not to manipulate your data!

Parameters: data: array-like

1d numpy array with experimental data

threshold: float or None

threshold factor to identify outlier data points. If None it will be ignored.

offset: None or float

offset value to identify outlier data points. If None it will be ignored.

Returns: array-like

1d data-array with spikes removed

xrayutilities.math.functions.multPeak1d(x, *args)

function to calculate the sum of multiple peaks in 1D. the peaks can be of different type and a background function (polynom) can also be included.

Parameters: x: array-like

coordinate where the function should be evaluated

args : list

list of peak/function types and parameters for every function type two arguments need to be given first the type of function as string with possible values 'g': Gaussian, 'l': Lorentzian, 'v': PseudoVoigt, 'a': asym. PseudoVoigt, 'p': polynom the second type of arguments is the tuple/list of parameters of the respective function. See documentation of math.Gauss1d, math.Lorentz1d, math.PseudoVoigt1d, math.PseudoVoigt1dasym, and numpy.polyval for details of the different function types.

Returns: array-like

value of the sum of functions at position x

xrayutilities.math.functions.multPeak2d(x, y, *args)

function to calculate the sum of multiple peaks in 2D. the peaks can be of different type and a background function (polynom) can also be included.

Parameters: x, y: array-like

coordinates where the function should be evaluated

args: list

list of peak/function types and parameters for every function type two arguments need to be given first the type of function as string with possible values 'g': Gaussian, 'l': Lorentzian, 'v': PseudoVoigt, 'c': constant the second type of arguments is the tuple/list of parameters of the respective function. See documentation of math.Gauss2d, math.Lorentz2d, math.PseudoVoigt2d for details of the different function types. The constant accepts a single float which will be added to the data

Returns: array-like

value of the sum of functions at position (x, y)

xrayutilities.math.functions.smooth(x, n)

function to smooth an array of data by averaging N adjacent data points

Parameters: x: array-like

1D data array

n: int

number of data points to average

Returns: xsmooth: array-like

smoothed array with same length as x

xrayutilities.math.misc module

xrayutilities.math.misc.center_of_mass (pos, data, background='none', full_output=False)
function to determine the center of mass of an array

Parameters: pos: array-like

position of the data points

data: array-like data values

background : {'none', 'constant', 'linear'}

type of background, either 'none', 'constant' or 'linear'

full_output : bool

return background cleaned data and background-parameters

Returns: float

center of mass position

xrayutilities.math.misc.fwhm_exp (pos, data)

function to determine the full width at half maximum value of experimental data. Please check the obtained value visually (noise influences the result)

Parameters: pos: array-like

position of the data points

data: array-like data values

Returns: float

fwhm value

xrayutilities.math.misc.gcd (lst)

greatest common divisor function using library functions

Parameters: Ist: array-like

array of integer values for which the greatest common divisor should be determined

Returns: gcd: int

xrayutilities.math.transforms module

xrayutilities.math.transforms.ArbRotation (axis, alpha, deg=True)

Returns a transform that represents a rotation around an arbitrary axis by the angle alpha. positive rotation is anti-clockwise when looking from positive end of axis vector

Parameters: axis: list or array-like

rotation axis

alpha: float

rotation angle in degree (deg=True) or in rad (deg=False)

deg: bool

determines the input format of ang (default: True)

Returns: Transform

class xrayutilities.math.transforms.AxisToZ (newzaxis)

Bases: xrayutilities.math.transforms.CoordinateTransform

Creates a coordinate transformation to move a certain axis to the z-axis. The rotation is done along the great circle. The x-axis of the new coordinate frame is created to be normal to the new and original z-axis. The new y-axis is create in order to obtain a right handed coordinate system.

class xrayutilities.math.transforms.AxisToZ_keepXY (newzaxis)

Bases: xrayutilities.math.transforms.CoordinateTransform

Creates a coordinate transformation to move a certain axis to the z-axis. The rotation is done along the great circle. The x-axis/y-axis of the new coordinate frame is created to be similar to the old x and y directions. This variant of AxisToZ assumes that the new Z-axis has its main component along the Z-direction

class xrayutilities.math.transforms.CoordinateTransform (v1, v2, v3)

Bases: xrayutilities.math.transforms.Transform

Create a Transformation object which transforms a point into a new coordinate frame. The new frame is determined by the three vectors v1/norm(v1), v2/norm(v2) and v3/norm(v3), which need to be orthogonal!

class xrayutilities.math.transforms.Transform (matrix)

Bases: object

property imatrix

inverse (args, rank=1)

performs inverse transformation a vector, matrix or tensor of rank 4

Parameters: args: list or array-like

object to transform, list or numpy array of shape (..., n) (..., n, n), (..., n, n, n, n) where n is the size of the transformation matrix.

rank: int

rank of the supplied object. allowed values are 1, 2, and 4

xrayutilities.math.transforms.XRotation (alpha, deg=True)

Returns a transform that represents a rotation about the x-axis by an angle alpha. If deg=True the angle is assumed to be in degree, otherwise the function expects radiants.

xrayutilities.math.transforms.**YRotation** (alpha, deg=True)

Returns a transform that represents a rotation about the y-axis by an angle alpha. If deg=True the angle is assumed to be in degree, otherwise the function expects radiants.

xrayutilities.math.transforms.ZRotation (alpha, deg=True)

Returns a transform that represents a rotation about the z-axis by an angle alpha. If deg=True the angle is assumed to be in degree, otherwise the function expects radiants.

xrayutilities.math.transforms.mycross (vec, mat)

function implements the cross-product of a vector with each column of a matrix

xrayutilities.math.transforms.rotarb (vec, axis, ang, deg=True)

function implements the rotation around an arbitrary axis by an angle ang positive rotation is anti-clockwise when looking from positive end of axis vector

Parameters: vec: list or array-like

vector to rotate

axis: list or array-like
rotation axis

ang: float

rotation angle in degree (deg=True) or in rad (deg=False)

deg: bool

determines the input format of ang (default: True)

Returns: rotvec: rotated vector as numpy.array

Examples

```
>>> rotarb([1, 0, 0],[0, 0, 1], 90)
array([ 6.12323400e-17,     1.00000000e+00,     0.00000000e+00])
```

xrayutilities.math.transforms.tensorprod (vec1, vec2) function implements an elementwise multiplication of two vectors

xrayutilities.math.vector module

module with vector operations for vectors of size 3, since for so short vectors numpy does not give the best performance explicit implementation of the equations is performed together with error checking to ensure vectors of length 3.

```
xrayutilities.math.vector.VecAngle ((v1.v2)/(norm(v1)*norm(v2)))
alpha = acos((v1.v2)/(norm(v1)*norm(v2)))
```

Parameters: v1, v2 : list or array-like

input vector(s), either one vector or an array of vectors with shape (n, 3)

deg: bool, optional

True: return result in degree, False: in radiants (default: False)

Returns: float or ndarray

the angle included by the two vectors v1 and v2, either a single float or an array with shape (n,)

xrayutilities.math.vector.VecCross (v1, v2, out=None)

Calculate the vector cross product.

Parameters: v1, v2 : list or array-like

input vector(s), either one vector or an array of vectors with shape (n, 3)

out: list or array-like, optional

output vector

Returns: ndarray

cross product either of shape (3,) or (n, 3)

xrayutilities.math.vector.VecDot (v1, v2)

Calculate the vector dot product.

Parameters: v1, v2 : list or array-like

input vector(s), either one vector or an array of vectors with shape (n, 3)

Returns: float or ndarray

innter product of the vectors, either a single float or (n,)

xrayutilities.math.vector.VecNorm(V)

Calculate the norm of a vector.

Parameters: v: list or array-like

input vector(s), either one vector or an array of vectors with shape (n, 3)

Returns: float or ndarray

vector norm, either a single float or shape (n,)

xrayutilities.math.vector.VecUnit (v)

Calculate the unit vector of v.

Parameters: v: list or array-like

input vector(s), either one vector or an array of vectors with shape (n, 3)

Returns: ndarray

unit vector of v, either shape (3,) or (n, 3)

xrayutilities.math.vector.distance (x, y, z, point, vec)

calculate the distance between the point (x, y, z) and the line defined by the point and vector vec

Parameters: x: float or ndarray

x coordinate(s) of the point(s)

y: float or ndarray

y coordinate(s) of the point(s)

z: float or ndarray

z coordinate(s) of the point(s)

point: tuple, list or ndarray

3D point on the line to which the distance should be calculated

vec: tuple, list or ndarray

3D vector defining the propergation direction of the line

xrayutilities.math.vector.getSyntax (Vec)

returns vector direction in the syntax 'x+' 'z-' or equivalents therefore works only for principle vectors of the coordinate system like e.g. [1, 0, 0] or [0, 2, 0]

Parameters: vec: list or array-like

vector of length 3

Returns: str

vector string following the synthax [xyz][+-]

xrayutilities.math.vector.getVector (string)

returns unit vector along a rotation axis given in the syntax 'x+' 'z-' or equivalents

Parameters: string: str

vector string following the synthax [xyz][+-]

Returns: ndarray

vector along the given direction

Module contents

xrayutilities.simpack package

Submodules

xrayutilities.simpack.darwin theory module

```
class xrayutilities.simpack.darwin_theory.DarwinModel (qz, qx=0, qy=0, **kwargs)
```

Bases: xrayutilities.simpack.models.LayerModel

model class inmplementing the basics of the Darwin theory for layers materials. This class is not fully functional and should be used to derive working models for particular material systems.

To make the class functional the user needs to implement the init structurefactors() and calc mono() methods

init_structurefactors()

calculates the needed atomic structure factors

ncalls = 0

simulate (MI)

main simulation function for the Darwin model. will calculate the reflected intensity

Parameters: ml: iterable

monolayer sequence of the sample. This should be created with the function make_monolayer(). see its documentation for details

class xrayutilities.simpack.darwin_theory.DarwinModelAlGaAs001 (qz, qx=0, qy=0, **kwargs)

Bases: xrayutilities.simpack.darwin_theory.DarwinModelAlloy

Darwin theory of diffraction for Al_x Ga_{1-x} As layers. The model is based on separation of the sample structure into building blocks of atomic planes from which a multibeam dynamical model is calculated.

Alas = <xrayutilities.materials.material.Crystal object>

GaAs = <xrayutilities.materials.material.Crystal object>

aGaAs = 5.65325

classmethod abulk (x)

calculate the bulk (relaxed) lattice parameter of the Al_{x}Ga_{1-x}As alloy

asub = 5.65325

eAl = AI(13)

eAs = As (33)

eGa = Ga(31)

classmethod get_dperp_apar (x, apar, r=1)

calculate inplane lattice parameter and the out of plane lattice plane spacing (of the atomic planes!) from composition and relaxation

Parameters: x: float

chemical composition parameter

apar: float

inplane lattice parameter of the material below the current layer (onto which the present layer is strained to). This value also served as a reference for the relaxation parameter.

r: float

relaxation parameter. 1=relaxed, 0=pseudomorphic

Returns: dperp: float

perpendicular d-spacing

apar: float

inplane lattice parameter

init_structurefactors (temp=300)

calculates the needed atomic structure factors

Parameters: temp: float, optional

temperature used for the Debye model

static poisson ratio (x)

calculate the Poisson ratio of the alloy

```
re = 2.8179403262e-05
```

class xrayutilities.simpack.darwin_theory.DarwinModelAlloy (qz, qx=0, qy=0, **kwargs)

Bases: xrayutilities.simpack.darwin_theory.DarwinModel, abc.ABC

extension of the DarwinModel for an binary alloy system were one parameter is used to determine the chemical composition

To make the class functional the user needs to implement the get_dperp_apar() method and define the substrate lattice parameter (asub). See the DarwinModelSiGe001 class for an implementation example.

abstract get_dperp_apar (x, apar, r=1)

calculate inplane lattice parameter and the out of plane lattice plane spacing (of the atomic planes!) from composition and relaxation.

Parameters: x: float

chemical composition parameter

apar: float

inplane lattice parameter of the material below the current layer (onto which the present layer is strained to). This value also served as a reference for the relaxation parameter.

r: float

relaxation parameter. 1=relaxed, 0=pseudomorphic

Returns: dperp: float

apar: float

make_monolayers (S)

create monolayer sequence from layer list

Parameters: s: list

layer model. list of layer dictionaries including possibility to form superlattices. As an example 5 repetitions of a Si(10nm)/Ge(15nm) superlattice on Si would like like:

the dictionaries must contain 't': thickness in A, 'x': chemical composition, and either 'r': relaxation or 'ai': inplane lattice parameter. Future implementations for asymmetric peaks might include layer type 'l' (not yet implemented). Already now any additional property in the dictionary will be handed on to the returned monolayer list.

asub: float

inplane lattice parameter of the substrate

Returns: list

monolayer list in a format understood by the simulate and xGe_profile methods

prop_profile (ml, prop)

calculate the profile of chemical composition or inplane lattice spacing from a monolayer list. One value for each monolayer in the sample is returned.

Parameters: ml: list

monolayer list created by make_monolayer()

prop: str

name of the property which should be evaluated. Use 'x' for the chemical composition

and 'ai' for the inplane lattice parameter.

Returns: zm: ndarray

z-position, z-0 is the surface

propx: ndarray

value of the property prop for every monolayer

class xrayutilities.simpack.darwin_theory.DarwinModelGaInAs001 (qz, qx=0, qy=0, **kwargs)

Bases: xrayutilities.simpack.darwin_theory.DarwinModelAlloy

Darwin theory of diffraction for Ga_{1-x} In_x As layers. The model is based on separation of the sample structure into building blocks of atomic planes from which a multibeam dynamical model is calculated.

GaAs = <xrayutilities.materials.material.Crystal object>

InAs = <xrayutilities.materials.material.Crystal object>

aGaAs = 5.65325

classmethod abulk (x)

calculate the bulk (relaxed) lattice parameter of the Ga_{1-x}ln_{x}As alloy

asub = 5.65325

eAs = As (33)

eGa = Ga (31)

eIn = ln (49)

classmethod get_dperp_apar (x, apar, r=1)

calculate inplane lattice parameter and the out of plane lattice plane spacing (of the atomic planes!) from composition and relaxation

Parameters: x: float

chemical composition parameter

apar: float

inplane lattice parameter of the material below the current layer (onto which the present layer is strained to). This value also served as a reference for the relaxation parameter.

r: float

relaxation parameter. 1=relaxed, 0=pseudomorphic

Returns: dperp: float

perpendicular d-spacing

apar: float

inplane lattice parameter

init_structurefactors (temp=300)

calculates the needed atomic structure factors

Parameters: temp: float, optional

temperature used for the Debye model

static poisson_ratio (x)

calculate the Poisson ratio of the alloy

```
re = 2.8179403262e-05
```

class xrayutilities.simpack.darwin_theory.DarwinModelSiGe001 (qz, qx=0, qy=0, **kwargs)

Bases: xrayutilities.simpack.darwin_theory.DarwinModelAlloy

model class implementing the Darwin theory of diffraction for SiGe layers. The model is based on separation of the sample structure into building blocks of atomic planes from which a multibeam dynamical model is calculated.

Ge = <xrayutilities.materials.material.Crystal object>

Si = <xrayutilities.materials.material.Crystal object>

aSi = 5.43104

classmethod abulk (x)

calculate the bulk (relaxed) lattice parameter of the alloy

asub = 5.43104

eGe = Ge (32)

esi = Si(14)

classmethod get_dperp_apar (x, apar, r=1)

calculate inplane lattice parameter and the out of plane lattice plane spacing (of the atomic planes!) from composition and relaxation

Parameters: x: float

chemical composition parameter

apar: float

inplane lattice parameter of the material below the current layer (onto which the present layer is strained to). This value also served as a reference for the relaxation parameter.

r: float, optional

relaxation parameter. 1=relaxed, 0=pseudomorphic

Returns: dperp: float

perpendicular d-spacing

apar: float

inplane lattice parameter

init_structurefactors (temp=300)

calculates the needed atomic structure factors

Parameters: temp: float, optional

temperature used for the Debye model

static poisson_ratio (x)

calculate the Poisson ratio of the alloy

re = 2.8179403262e-05

xrayutilities.simpack.darwin_theory.**GradedBuffer** (xfrom, xto, nsteps, thickness, relaxation=1) create a multistep graded composition buffer.

Parameters: xfrom: float

begin of the composition gradient

xto: float

end of the composition gradient

nsteps: int

number of steps of the gradient

thickness: float

total thickness of the Buffer in A

relaxation: float

relaxation of the buffer

Returns: list

layer list needed for the Darwin model simulation

xrayutilities.simpack.darwin_theory.getfirst (iterable, key)

helper function to obtain the first item in a nested iterable

xrayutilities.simpack.darwin_theory.getit (it, key)

generator to obtain items from nested iterable

xrayutilities.simpack.fit module

class xrayutilities.simpack.fit.FitModel (Imodel, verbose=False, plot=False, elog=True, **kwargs)

Bases: object

Wrapper for the Imfit Model class working for instances of LayerModel

Typically this means that after initialization of *FitModel* you want to use make_params to get a *Imfit.Parameters* list which one customizes for fitting.

Later on you can call fit and eval methods with those parameter list.

fit (data, params, x, weights=None, fit_kws=None, **kwargs)

wrapper around Imfit. Model. fit which enables plotting during the fitting

Parameters: data: ndarray

experimental values
params: Imfit.Parameters

list of parameters for the fit, use make_params for generation

x: ndarray

independent variable (incidence angle or q-position depending on the model)

weights: ndarray, optional

values of weights for the fit, same size as data

fit kws: dict, optional

Options to pass to the minimizer being used

kwargs: dict, optional

keyword arguments which are passed to Imfit.Model.fit

Returns: Imfit.ModelResult

set_fit_limits (xmin=- inf, xmax=inf, mask=None)

set fit limits. If mask is given it must have the same size as the *data* and *x* variables given to fit. If mask is None it will be generated from xmin and xmax.

Parameters: xmin: float, optional

minimum value of x-values to include in the fit

xmax: float, optional

maximum value of x-values to include in the fit

mask: boolean array, optional

mask to be used for the data given to the fit

xrayutilities.simpack.helpers module

xrayutilities.simpack.helpers.coplanar_alphai (qx, qz, en='config') calculate coplanar incidence angle from knowledge of the qx and qz coordinates

Parameters: qx: array-like

inplane momentum transfer component

qz: array-like

out of plane momentum transfer component

en: float or str, optional

x-ray energy (eV). By default the value from the config is used.

Returns: alphai: array-like

the incidence angle in degree. points in the Laue zone are set to 'nan'.

xrayutilities.simpack.helpers.get_qz (qx, alphai, en='config')

calculate the qz position from the qx position and the incidence angle for a coplanar diffraction geometry

Parameters: qx: array-like

inplane momentum transfer component

alphai: array-like

incidence angle (deg)

en: float or str, optional

x-ray energy (eV). By default the value from the config is used.

Returns: array-like

the gz position for the given incidence angle

xrayutilities.simpack.models module

class xrayutilities.simpack.models.DiffuseReflectivityModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.SpecularReflectivityModel

model for diffuse reflectivity calculations

The 'simulate' method calculates the diffuse reflectivity on the specular rod in coplanar geometry in analogy to the SpecularReflectivityModel.

The 'simulate_map' method calculates the diffuse reflectivity for a 2D set of Q-positions. This method can also calculate the intensity for other geometries, like GISAXS with constant incidence angle or a quasi omega/2theta scan in GISAXS geometry.

simulate (alphai)

performs the actual diffuse reflectivity calculation for the specified incidence angles. This method always uses the coplanar geometry independent of the one set during the initialization.

Parameters: alphai: array-like

vector of incidence angles

Returns: array-like

vector of intensities of the reflectivity signal

simulate_map (qL, qz)

performs diffuse reflectivity calculation for the rectangular grid of reciprocal space positions define by qL and qz. This method uses the method and geometry set during the initialization of the class.

Parameters: qL: array-like

lateral coordinate in reciprocal space (vector with NqL components)

qz: array-like

vertical coordinate in reciprocal space (vector with Ngz components)

Returns: array-like

matrix of intensities of the reflectivity signal, with shape (len(qL), len(qz))

class xrayutilities.simpack.models.DynamicalModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.SimpleDynamicalCoplanarModel

Dynamical diffraction model for specular and off-specular qz-scans. Calculation of the flux of reflected and diffracted waves for general asymmetric coplanar diffraction from an arbitrary pseudomorphic multilayer is performed by a generalized 2-beam theory (4 tiepoints, S and P polarizations)

The first layer in the model is always assumed to be the semiinfinite substrate indepentent of its given thickness

simulate (alphai, hkl=None, geometry='hi_lo', rettype='intensity')

performs the actual diffraction calculation for the specified incidence angles and uses an analytic solution for the quartic dispersion equation

Parameters: alphai: array-like

vector of incidence angles (deg)

hkl: list or tuple, optional

Miller indices of the diffraction vector (preferable use set_hkl method to speed up repeated calculations of the same peak!)

geometry: {'hi_lo', 'lo_hi'}, optional

'hi_lo' for grazing exit (default) and 'lo_hi' for grazing incidence

rettype: {'intensity', 'field', 'all'}, optional

type of the return value. 'intensity' (default): returns the diffracted beam flux convoluted with the resolution function; 'field': returns the electric field (complex) without convolution with the resolution function, 'all': returns the electric field, ai, af (both in degree), and the reflected intensity.

Returns: arrav-like

vector of intensities of the diffracted signal, possibly changed return value due the rettype setting!

class xrayutilities.simpack.models.DynamicalReflectivityModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.SpecularReflectivityModel

model for Dynamical Specular Reflectivity Simulations. It uses the transfer Matrix methods as given in chapter 3 "Daillant, J., & Gibaud, A. (2008). X-ray and Neutron Reflectivity"

scanEnergy (energies, angle)

Simulates the Dynamical Reflectivity as a function of photon energy at fixed angle.

Parameters: energies: numpy.ndarray or list

photon energies (in eV).

angle: float

fixed incidence angle

Returns: reflectivity: array-like

vector of intensities of the reflectivity signal

transmitivity: array-like

vector of intensities of the transmitted signal

simulate (alphai)

Simulates the Dynamical Reflectivity as a function of angle of incidence

Parameters: alphai: array-like

vector of incidence angles

Returns: reflectivity: array-like

vector of intensities of the reflectivity signal

transmitivity: array-like

vector of intensities of the transmitted signal

class xrayutilities.simpack.models.KinematicalModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.LayerModel

Kinematical diffraction model for specular and off-specular qz-scans. The model calculates the kinematical contribution of one (hkl) Bragg peak, however considers the variation of the structure factor for different 'q'. The surface geometry is specified using the Experiment-object given to the constructor.

init_chi0()

calculates the needed optical parameters for the simulation. If any of the materials/layers is changing its properties this function needs to be called again before another correct simulation is made. (Changes of thickness does NOT require this!)

simulate (qz, hkl, absorption=False, refraction=False, rettype='intensity')

performs the actual kinematical diffraction calculation on the Qz positions specified considering the contribution from a single Bragg peak.

Parameters: qz: array-like

simulation positions along qz

hkl: list or tuple

Miller indices of the Bragg peak whos truncation rod should be calculated

absorption: bool, optional

flag to tell if absorption correction should be used

refraction: bool, optional

flag to tell if basic refraction correction should be performed. If refraction is True absorption correction is also included independent of the absorption flag.

rettype: {'intensity', 'field', 'all'}

type of the return value. 'intensity' (default): returns the diffracted beam flux convoluted with the resolution function; 'field': returns the electric field (complex) without convolution with the resolution function, 'all': returns the electric field, ai, af (both in degree), and the reflected intensity.

Returns: array-like

return value depends on the setting of *rettype*, by default only the calculate intensity is returned

class xrayutilities.simpack.models.KinematicalMultiBeamModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.KinematicalModel

Kinematical diffraction model for specular and off-specular qz-scans. The model calculates the kinematical contribution of several Bragg peaks on the truncation rod and considers the variation of the structure factor. In order to use a analytical description for the kinematic diffraction signal all layer thicknesses are changed to a multiple of the respective lattice parameter along qz. Therefore this description only works for (001) surfaces.

simulate (qz, hkl, absorption=False, refraction=True, rettype='intensity')

performs the actual kinematical diffraction calculation on the Qz positions specified considering the contribution from a full truncation rod

Parameters: qz: array-like

simulation positions along qz

hkl: list or tuple

Miller indices of the Bragg peak whos truncation rod should be calculated

absorption: bool, optional

flag to tell if absorption correction should be used

refraction: bool, optional,

flag to tell if basic refraction correction should be performed. If refraction is True absorption correction is also included independent of the absorption flag.

rettype: {'intensity', 'field', 'all'}

type of the return value. 'intensity' (default): returns the diffracted beam flux convoluted with the resolution function; 'field': returns the electric field (complex) without convolution with the resolution function, 'all': returns the electric field, ai, af (both in degree), and the reflected intensity.

Returns: array-like

return value depends on the setting of *rettype*, by default only the calculate intensity is returned

class xrayutilities.simpack.models.LayerModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.Model, abc.ABC

generic model class from which further thin film models can be derived from

get_polarizations()

return list of polarizations which should be calculated

join polarizations (Is, Ip)

method to calculate the total diffracted intensity from the intensities of S and P-polarization.

abstract simulate ()

abstract method that every implementation of a LayerModel has to override.

class xrayutilities.simpack.models.Model (experiment, **kwargs)

Bases: object

generic model class from which further models can be derived from

convolute_resolution (x, y)

convolve simulation result with a resolution function

Parameters: x: array-like

x-values of the simulation, units of x also decide about the unit of the resolution_width parameter

y: array-like

. array-like

y-values of the simulation

Returns: array-like

convoluted y-data with same shape as y

property energy

scale_simulation(y)

scale simulation result with primary beam flux/intensity and add a background.

Parameters: y : array-like

y-values of the simulation

Returns: array-like

scaled y-values

class xrayutilities.simpack.models.ResonantReflectivityModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.SpecularReflectivityModel

model for specular reflectivity calculations CURRENTLY UNDER DEVELOPEMENT! DO NOT USE!

simulate (alphai)

performs the actual reflectivity calculation for the specified incidence angles

Parameters: alphai: array-like

vector of incidence angles

Returns: array-like

vector of intensities of the reflectivity signal

class xrayutilities.simpack.models.SimpleDynamicalCoplanarModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.KinematicalModel

Dynamical diffraction model for specular and off-specular qz-scans. Calculation of the flux of reflected and diffracted waves for general asymmetric coplanar diffraction from an arbitrary pseudomorphic multilayer is performed by a simplified 2-beam theory (2 tiepoints, S and P polarizations)

No restrictions are made for the surface orientation.

The first layer in the model is always assumed to be the semiinfinite substrate indepentent of its given thickness

Note

This model should not be used in real life scenarios since the made approximations severely fail for distances far from the reference position.

set_hkl (*hkl)

To speed up future calculations of the same Bragg peak optical parameters can be pre-calculated using this function.

Parameters: hkl : list or tuple

Miller indices of the Bragg peak for the calculation

simulate (alphai, hkl=None, geometry='hi_lo', idxref=1)

performs the actual diffraction calculation for the specified incidence angles.

Parameters: alphai: array-like

vector of incidence angles (deg)

hkl: list or tuple, optional

Miller indices of the diffraction vector (preferable use set_hkl method to speed up repeated calculations of the same peak!)

geometry: {'hi_lo', 'lo_hi'}, optional

'hi_lo' for grazing exit (default) and 'lo_hi' for grazing incidence

idxref: int, optional

index of the reference layer. In order to get accurate peak position of the film peak you want this to be the index of the film peak (default: 1). For the substrate use 0.

Returns: array-like

vector of intensities of the diffracted signal

class xrayutilities.simpack.models.SpecularReflectivityModel (*args, **kwargs)

Bases: xrayutilities.simpack.models.LayerModel

model for specular reflectivity calculations

densityprofile (nz, plot=False)

calculates the electron density of the layerstack from the thickness and roughness of the individual layers

Parameters: nz: int

number of values on which the profile should be calculated

plot: bool, optional

flag to tell if a plot of the profile should be created

Returns: z: array-like

z-coordinates, z = 0 corresponds to the surface

eprof : array-like
 electron profile

init_cd()

calculates the needed optical parameters for the simulation. If any of the materials/layers is changing its properties this function needs to be called again before another correct simulation is made. (Changes of thickness and roughness do NOT require this!)

simulate (alphai)

performs the actual reflectivity calculation for the specified incidence angles

Parameters: alphai: array-like

vector of incidence angles

Returns: array-like

vector of intensities of the reflectivity signal

xrayutilities.simpack.models.effectiveDensitySlicing (layerstack, step, roughness=0, cutoff=1e-05)

Function to slice a LayerStack into many amorphous sublayers for effective density modelling of X-ray reflectivity of thin and rough multilayers. The resulting LayerStack will consist of perfectly smooth layers with average density/composition resulting from an error-function like transition between the rough layers of the initial stack. At the surface an vacuum layer is automatically added to the initial stack.

Parameters: layerstack: initial LayerStack, can contain only Amorhous layers!

step: thickness (in Angstrom) of the slices in the returned

LayerStack

roughness: roughness of the created sublayers (in Angstrom)

cutoff: layers with relative weights below this value will be ignored

Returns: LayerStack

xrayutilities.simpack.models.startdelta (start, delta, num)

xrayutilities.simpack.mosaicity module

xrayutilities.simpack.mosaicity.mosaic_analytic (qx, qz, RL, RV, Delta, hx, hz, shape) simulation of the coplanar reciprocal space map of a single mosaic layer using a simple analytic approximation

Parameters: qx: array-like

vector of the qx values (offset from the Bragg peak)

qz: array-like

vector of the qz values (offset from the Bragg peak)

RL: float

lateral block radius in Angstrom

RV: float

vertical block radius in Angstrom

Delta: float

root mean square misorientation of the grains in degree

hx: float

lateral component of the diffraction vector

hz: float

vertical component of the diffraction vector

shape: float

shape factor (1.. Gaussian)

Returns: array-like

2D array with calculated intensities

xrayutilities.simpack.powder module

This module contains the core definitions for the XRD Fundamental Parameneters Model (FPA) computation in Python. The main computational class is FP_profile, which stores cached information to allow it to efficiently recompute profiles when parameters have been modified. For the user an Powder class is available which can calculate a complete powder pattern of a crystalline material.

The diffractometer line profile functions are calculated by methods from Cheary & Coelho 1998 and Mullen & Cline paper and 'R' package. Accumulate all convolutions in Fourier space, for efficiency, except for axial divergence, which needs to be weighted in real space for I3 integral.

More details about the applied algorithms can be found in the paper by M. H. Mendelhall et al., Journal of Research of NIST 120, 223 (2015) to which you should also refer for a careful definition of all the parameters

class xrayutilities.simpack.powder.FP_profile (anglemode, gaussian_smoother_bins_sigma=1.0,
oversampling=10)

Bases: object

the main fundamental parameters class, which handles a single reflection. This class is designed to be highly extensible by inheriting new convolvers. When it is initialized, it scans its namespace for specially formatted names, which can come from mixin classes. If it finds a function name of the form conv_xxx, it will call this funtion to create a convolver. If it finds a name of the form info_xxx it will associate the dictionary with that convolver, which can be used in UI generation, for example. The class, as it stands, does nothing significant with it. If it finds str_xxx, it will use that function to format a printout of the current state of the convolver conv_xxx, to allow improved report generation for convolvers.

When it is asked to generate a profile, it calls all known convolvers. Each convolver returns the Fourier transform of its convolvution. The transforms are multiplied together, inverse transformed, and after fixing the periodicity issue, subsampled, smoothed and returned.

If a convolver returns None, it is not multipled into the product.

Parameters: max history length: int

the number of histories to cache (default=5); can be overridden if memory is an issue.

length_scale_m : float

length_scale_m sets scaling for nice printing of parameters. if the units are in mm everywhere, set it to 0.001, e.g. convolvers which implement their own str_xxx method may use this to format their results, especially if 'natural' units are not meters. Typical is wavelengths and lattices in nm or angstroms, for example.

add_buffer (b)

add a numpy array to the list of objects that can be thrown away on pickling.

Parameters: b : array-like

the buffer to add to the list

Returns: b : array-like

return the same buffer, to make nesting easy.

axial_helper (outerbound, innerbound, epsvals, destination, peakpos=0, y0=0, k=0)

the function F0 from the paper. compute k/sqrt(peakpos-x)+y0 nonzero between outer & inner (inner is closer to peak) or k/sqrt(x-peakpos)+y0 if reversed (i.e. if outer > peak) fully evaluated on a specified eps grid, and stuff into destination

Parameters: outerbound: float

the edge of the function farthest from the singularity, referenced to epsvals

innerbound: float

the edge closest to the singularity, referenced to epsvals

epsvals: array-like

the array of two-theta values or offsets

destination: array-like

an array into which final results are summed. modified in place!

peakpos: float

the position of the singularity, referenced to epsvals.

y0: float

the constant offset

k: float

the scale factor

Returns: lower_index, upper_index : int

python style bounds for region of destination which has been modified.

compute_line_profile (convolver_names=None, compute_derivative=False, return_convolver=False)
 execute all the convolutions; if convolver_names is None, use everything we have, otherwise, use named convolutions.

Parameters: convolver names: list

a list of convolvers to select. If None, use all found convolvers.

compute_derivative: bool

if True, also return d/dx(function) for peak position fitting

Returns: object

a profile_data object with much information about the peak

conv_absorption()

compute the sample transparency correction, including the finite-thickness version

Returns: array-like

the convolver

conv_axial()

compute the Fourier transform of the axial divergence component

Returns: array-like

the transform buffer, or None if this is being ignored

conv_displacement ()

compute the peak shift due to sample displacement and the 2theta zero offset

Returns: array-like

the convolver

conv_emission()

compute the emission spectrum and (for convenience) the particle size widths

Returns: array-like

the convolver for the emission and particle sizes

Note

the particle size and strain stuff here is just to be consistent with *Topas* and to be vaguely efficient about the computation, since all of these have the same general shape.

conv_flat_specimen()

compute the convolver for the flat-specimen correction

Returns: array-like

the convolver

conv_global()

a dummy convolver to hold global variables and information. the global context isn't really a convolver, returning *None* means ignore result

Returns: None

always returns None

conv_receiver_slit ()

compute the rectangular convolution for the receiver slit or SiPSD pixel size

Returns: array-like

the convolver

conv_si_psd()

compute the convolver for the integral of defocusing of the face of an Si PSD

Returns: array-like

the convolver

conv_smoother ()

compute the convolver to smooth the final result with a Gaussian before downsampling.

Returns: array-like

the convolver

conv_tube_tails()

compute the Fourier transform of the rectangular tube tails function

Returns: array-like

the transform buffer, or None if this is being ignored

full_axdiv_I2 (Lx=None, Ls=None, Lr=None, R=None, twotheta=None, beta=None, epsvals=None)
return the /2 function

Parameters: Lx: float

length of the xray filament

Ls: float

length of the sample

Lr: float

length of the receiver slit

R: float

diffractometer length, assumed symmetrical

twotheta: float

angle, in radians, of the center of the computation

beta: *float* offset angle

epsvals: array-like
array of offsets from center of computation, in radians

Returns: epsvals: array-like

array of offsets from center of computation, in radians

idxmin, idxmax: int

the full python-style bounds of the non-zero region of I2p and I2m

I2p, I2m: array-like

12+ and 12- from the paper, the contributions to the intensity

full_axdiv_I3 (Lx=None, Ls=None, Lr=None, R=None, twotheta=None, epsvals=None, sollerIdeg=None,
sollerDdeg=None, nsteps=10, axDiv=")

carry out the integral of 12 over beta and the Soller slits.

Parameters: Lx: float

length of the xray filament

Ls: float

length of the sample

Lr: float

length of the receiver slit

R: float

the (assumed symmetrical) diffractometer radius

twotheta: float

angle, in radians, of the center of the computation

epsvals: array-like

array of offsets from center of computation, in radians

sollerideg: float

the full-width (both sides) cutoff angle of the incident Soller slit

sollerDdeg: float

the full-width (both sides) cutoff angle of the detector Soller slit

nsteps: int

the number of subdivisions for the integral

axDiv: str not used

Returns: array-like

the accumulated integral, a copy of a persistent buffer _axial

general_tophat (name=", width=None)

a utility to compute a transformed tophat function and save it in a convolver buffer

Parameters: name: str

the name of the convolver cache buffer to update

width: float

the width in 2-theta space of the tophat

Returns: array-like

the updated convolver buffer, or None if the width was None

get_conv (name, key, format=<class 'float'>)

get a cached, pre-computed convolver associated with the given parameters, or a newly zeroed convolver if the cache doesn't contain it. Recycles old cache entries.

This takes advantage of the mutability of arrays. When the contents of the array are changed by the convolver, the cached copy is implicitly updated, so that the next time this is called with the same parameters, it will return the previous array.

Parameters: name: str

the name of the convolver to seek

key: object

any hashable object which identifies the parameters for the computation

format: numpy.dtype, optional

the type of the array to create, if one is not found.

Returns: bool

flag, which is *True* if valid data were found, or *False* if the returned array is zero, and *array*, which must be computed by the convolver if *flag* was *False*.

get_convolver_information()

return a list of convolvers, and what we know about them. function scans for functions named conv_xxx, and associated info_xxx entries.

Returns: list

list of (convolver_xxx, info_xxx) pairs

get_function_name ()

return the name of the function that called this. Useful for convolvers to identify themselves

Returns: str

name of calling function

get_good_bin_count (count)

find a bin count close to what we need, which works well for Fourier transforms.

Parameters: count: int

a number of bins.

Returns: int

a bin count somewhat larger than count which is efficient for FFT

info_emission = {'group_name': 'Incident beam and crystal size', 'help': 'this should be help information', 'param_info': {'crystallite_size_gauss': ('Gaussian crystallite size fwhm (m)', 1e-06), 'crystallite_size_lor': ('Lorentzian crystallite size fwhm (m)', 1e-06), 'emiss_gauss_widths': ('Gaussian emissions fwhm (m)', (1e-13,)), 'emiss_intensities': ('relative intensities', (1.0,)), 'emiss_lor_widths': ('Lorenztian emission fwhm (m)', (1e-13,)), 'emiss_wavelengths': ('wavelengths (m)', (1.58e-10,))}}

info_global = {'group_name': 'Global parameters', 'help': 'this should be help information', 'param_info': {'d': ('d spacing (m)', 4e-10), 'dominant_wavelength': ('wavelength of most intense line (m)', 1.5e-10), 'twotheta0_deg': ('Bragg center of peak (degrees)', 30.0)}}

classmethod isequivalent (hkl1, hkl2, crystalsystem)

function to determine if according to the convolvers included in this class two sets of Miller indices are equivalent. This function is only called when the class attribute 'isotropic' is False.

Parameters: hkl1, hkl2 : list or tuple

Miller indices to be checked for equivalence

crystalsystem: str

symmetry class of the material which is considered

Returns: bool

isotropic = True

length scale m = 1.0

max_history_length = 5

self_clean()

do some cleanup to make us more compact; Instance can no longer be used after doing this, but can be pickled.

set_optimized_window (twotheta_window_center_deg, twotheta_approx_window_fullwidth_deg,
twotheta exact bin spacing deg)

pick a bin count which factors cleanly for FFT, and adjust the window width to preserve the exact center and bin spacing

Parameters: twotheta_window_center_deg: float

exact position of center bin, in degrees

twotheta_approx_window_fullwidth_deg: float

approximate desired width

twotheta_exact_bin_spacing_deg: float

the exact bin spacing to use

set_parameters (convolver='global', **kwargs)

update the dictionary of parameters associated with the given convolver

Parameters: convolver: str

the name of the convolver. name 'global', e.g., attaches to function 'conv_global'

kwargs: dict

keyword-value pairs to update the convolvers dictionary.

set_window (twotheta_window_center_deg, twotheta_window_fullwidth_deg, twotheta_output_points)
move the compute window to a new location and compute grids, without resetting all parameters. Clears convolution history and sets up many arrays.

Parameters: twotheta_window_center_deg: float

the center position of the middle bin of the window, in degrees

twotheta_window_fullwidth_deg: float the full width of the window, in degrees

twotheta_output_points: int

the number of bins in the final output

str_emission()

format the emission spectrum and crystal size information

Returns: str

the formatted information

str global()

returns a string representation for the global context.

Returns: str

report on global parameters.

class xrayutilities.simpack.powder.PowderDiffraction (mat, **kwargs)

Bases: xrayutilities.experiment.PowderExperiment

Experimental class for powder diffraction. This class calculates the structure factors of powder diffraction lines and uses instances of FP_profile to perform the convolution with experimental resolution function calculated by the fundamental parameters approach. This class uses multiprocessing to speed up calculation. Set config.NTHREADS=1 to restrict this to one worker process.

Calculate (twotheta, **kwargs)

calculate the powder diffraction pattern including convolution with the resolution function and map them onto the twotheta positions. This also performs the calculation of the peak intensities from the internal material object

Parameters: twotheta: array-like

two theta values at which the powder pattern should be calculated.

kwargs: dict

additional keyword arguments are passed to the Convolve function

Returns: array-like

output intensity values for the twotheta values given in the input

Notes

Bragg peaks are only included up to tt_cutoff set in the class constructor!

Convolve (twotheta, window_width='config', mode='multi')

convolute the powder lines with the resolution function and map them onto the twotheta positions. This calculates the powder pattern excluding any background contribution

Parameters: twotheta: array-like

two theta values at which the powder pattern should be calculated.

window_width: float, optional

width of the calculation window of a single peak

mode: {'multi, 'local'}, optional

multiprocessing mode, either 'multi' to use multiple processes or 'local' to restrict the calculation to a single process

Note:

Bragg peaks are only included up to tt cutoff set in the class constructor!

Returns: output intensity values for the twotheta values given in the input

close ()

correction_factor (ang)

calculate the correction factor for the diffracted intensities. This contains the polarization effects and the Lorentz factor

Parameters: ang: aray-like

theta diffraction angles for which the correction should be calculated

Returns: f: array-like

array of the same shape as ang containing the correction factors

property energy

init_powder_lines (tt_cutoff)

calculates the powder intensity and positions up to an angle of tt_cutoff (deg) and stores the result in the data dictionary whose structure is as follows:

The data dictionary has one entry per line with a unique identifier as key of the entry. The entries themself are dictionaries which have the following entries:

- hkl : (h, k, l), Miller indices of the Bragg peak
- r : reflection strength of the line
- ang : Bragg angle of the peak (theta = 2theta/2!)
- · qpos: reciprocal space position

load_settings_from_config (settings)

load parameters from the config and update these settings with the options from the settings parameter

merge_lines (data)

if calculation is isotropic lines at the same q-position can be merged to one line to reduce the calculational effort

Parameters: data : ndarray

numpy field array with values of 'hkl' (Miller indices of the peaks), 'q' (q-position), and 'r' (reflection strength) as produced by the *reflection_strength* method

Returns: hkl, q, ang, r: array-like

Miller indices, q-position, diffraction angle (Theta), and reflection strength of the

material

reflection_strength (tt_cutoff)

determine structure factors/reflection strength of all Bragg peaks up to tt_cutoff. This function also implements the March-Dollase model for preferred orientation in the symmetric reflection mode. Note that although this means the sample has anisotropic properties the various lines can still be merged together since at the moment no anisotropic crystal shape is supported.

Parameters: tt_cutoff: float

upper cutoff value of 2theta until which the reflection strength are calculated

Returns: ndarray

numpy array with field for 'hkl' (Miller indices of the peaks), 'q' (q-position), and 'r'

(reflection strength) of the Bragg peaks

set_sample_parameters()

load sample parameters from the Powder class and use them in all FP_profile instances of this object

set_wavelength_from_params()

sets the wavelenth in the base class from the settings dictionary of the FP_profile classes and also set it in the 'global' part of the parameters

set window (force=False)

sets the calcultion window for all convolvers

property twotheta

update_powder_lines (tt_cutoff)

calculates the powder intensity and positions up to an angle of tt_cutoff (deg) and updates the values in:

- ids: list of unique identifiers of the powder line
- data: array with intensities
- ang: bragg angles of the peaks (theta=2theta/2!)
- qpos: reciprocal space position of intensities

update_settings (newsettings={})

update settings of all instances of FP_profile

Parameters:

newsettings: dict

```
dictionary with new settings. It has to include one subdictionary for every convolver
                           which should have its settings changed.
  property wavelength
  property window_width
xrayutilities.simpack.powder.chunkify(lst, n)
class xrayutilities.simpack.powder.convolver_handler
  Bases: object
  manage the convolvers of on process
  add_convolver (convolver)
  calc (run, ttpeaks)
    calculate profile function for selected convolvers
        Parameters: run: list
                           list of flags of length of convolvers to tell which convolver needs to be run
                       ttpeaks: array-like
                           peak positions for the convolvers
            Returns:
                      list
                           list of profile data result objects
  set_windows (centers, npoints, flag, width)
  update_parameters (parameters)
class xrayutilities.simpack.powder.manager (address=None, authkey=None, serializer='pickle', ctx=None)
  Bases: multiprocessing.managers.BaseManager
class xrayutilities.simpack.powder.profile_data (**kwargs)
  Bases: object
  a skeleton class which makes a combined dict and namespace interface for easy pickling and data passing
  add_symbol (**kwargs)
    add new symbols to both the attributes and dictionary for the class
        Parameters: kwargs: dict
                           keyword=value pairs
xrayutilities.simpack.powdermodel module
class xrayutilities.simpack.powdermodel.PowderModel (*args, **kwargs)
  Bases: object
  Class to help with powder calculations for multiple materials. For basic calculations the Powder class together with
  the Fundamental parameters approach is used.
  close()
  create_fitparameters()
    function to create a fit model with all instrument and sample parameters.
            Returns: Imfit.Parameters
  fit (params, twotheta, data, std=None, maxfev=200)
```

make least squares fit with parameters supplied by the user

Parameters: params: Imfit.Parameters

object with all parameters set as intended by the user

twotheta: array-like

angular values for the fit

data: array-like

experimental intensities for the fit

std: array-like

standard deviation of the experimental data. if 'None' the sqrt of the data will be used

maxfev: int

maximal number of simulations during the least squares refinement

Returns: Imfit.MinimizerResult

plot (twotheta, showlines=True, label='simulation', color=None, formatspec='-', lcolors=[], ax=None, **kwargs) plot the powder diffraction pattern and indicate line positions for all components in the model.

Parameters: twotheta: array-like

positions at which the powder pattern should be evaluated

showlines: bool, optional

flag to decide if peak positions of the components will be shown on the top of the plot

label: str

line label in the plot

color: matplotlib color or None

the color used for the line plot of the simulation

formatspec : str

format specifier of the simulation curve

Icolors: list of matplotlib colors

colors for the line indicators for the various components

ax: matplotlib.axes or None

axes object to be used for plotting, if its given no axes decoration like labels are set.

Further keyword arguments are passed to the simulate method.

Returns: matplotlib.axes object

set_background (btype, **kwargs)

define background as spline or polynomial function

Parameters: btype : {polynomial', 'spline'}

background type; Depending on this value the expected keyword arguments differ.

kwargs: dict

optional keyword arguments

x: array-like, optional

x-values (twotheta) of the background points (if btype='spline')

y: array-like, optional

intensity values of the background (if btype='spline')

p: array-like, optional

polynomial coefficients from the highest degree to the constant term. len of p decides about the degree of the polynomial (if btype='polynomial')

set lmfit parameters (Imparams)

function to update the settings of this class during an least squares fit

Parameters: Imparams: Imfit.Parameters

Imfit Parameters list of sample and instrument parameters

set_parameters (params)

set simulation parameters of all subobjects

Parameters: params: dict

settings dictionaries for the convolvers.

simulate (twotheta, **kwargs)

calculate the powder diffraction pattern of all materials and sum the results based on the relative volume of the materials.

Parameters: twotheta: array-like

positions at which the powder pattern should be evaluated

kwargs: dict

optional keyword arguments

background: array-like

an array of background values (same shape as twotheta) if no background is given then the background is calculated as previously set by the set_background function or is 0

further keyword arguments are passed to the Convolve function of of the

PowderDiffraction objects

Returns: array-like

summed powder diffraction intensity of all materials present in the model

xrayutilities.simpack.powdermodel.Rietveld_error_metrics (exp, sim, weight=None, std=None, Nvar=0, disp=False)

calculates common error metrics for Rietveld refinement.

Parameters: exp: array-like

experimental datapoints

sim : array-like simulated data

weight: array-like, optional

weight factor in the least squares sum. If it is None the weight is estimated from the counting statistics of 'exp'

std: array-like, optional

standard deviation of the experimental data. alternative way of specifying the weight factor, when both are given weight overwrites std!

Nvar: int, optional

number of variables in the refinement

disp: bool, optional

flag to tell if a line with the calculated values should be printed.

Returns: M, Rp, Rwp, Rwpexp, chi2: float

xrayutilities.simpack.powdermodel.plot_powder (twotheta, exp, sim, mask=None, scale='sqrt',
fig='XU:powder', show_diff=True, show_legend=True, labelexp='experiment', labelsim='simulation', formatexp='.-k',
formatsim='-r')

Convenience function to plot the comparison between experimental and simulated powder diffraction data

Parameters: twotheta: array-like

angle values used for the x-axis of the plot (deg)

exp: array-like

experimental data (same shape as twotheta). If None only the simulation and no difference will be plotted

sim: array-like or PowederModel

simulated data or PowderModel instance. If a PowderModel instance is given the plot-method of PowderModel is used.

mask: array-like, optional

mask to reduce the twotheta values to the be used as x-coordinates of sim

scale: {'linear', 'sgrt', 'log'}, optional

string specifying the scale of the y-axis.

fig: str or int, optional

matplotlib figure name (figure will be cleared!)

show_diff: bool, optional

flag to specify if a difference curve should be shown

show_legend: bool, optional

flag to specify if a legend should be shown

labelexp: str

plot label (legend entry) for the experimental data

labelsim: str

plot label for the simulation data

formatexp: str

format specifier for the experimental data

formatsim: str

format specifier for the simulation curve

xrayutilities.simpack.smaterials module

```
class xrayutilities.simpack.smaterials.CrystalStack (name, *args)
Bases: xrayutilities.simpack.smaterials.LayerStack
  extends the built in list type to enable building a stack of crystalline Layers by various methods.

check (v)

class xrayutilities.simpack.smaterials.GradedLayerStack (alloy, xfrom, xto, nsteps, thickness, **kwargs)
Bases: xrayutilities.simpack.smaterials.CrystalStack
  generates a sequence of layers with a gradient in chemical composition
```

class xrayutilities.simpack.smaterials.Layer (material, thickness, **kwargs)

Bases: xrayutilities.simpack.smaterials.SMaterial

Object describing part of a thin film sample. The properties of a layer are :

Attributes: material: Material (Crystal or Amorhous)

an xrayutilties material describing optical and crystal properties of the thin film

thickness: float

film thickness in Angstrom

class xrayutilities.simpack.smaterials.LayerStack (name, *args)

Bases: xrayutilities.simpack.smaterials.MaterialList

```
extends the built in list type to enable building a stack of Layer by various methods.
  check (V)
class xrayutilities.simpack.smaterials.MaterialList (name, *args)
  Bases: collections.abc.MutableSequence
  class representing the basics of a list of materials for simulations within xrayutilities. It extends the built in list type.
  check (V)
  insert (i, v)
    S.insert(index, value) - insert value before index
class xrayutilities.simpack.smaterials.Powder (material, volume, **kwargs)
  Bases: xrayutilities.simpack.smaterials.SMaterial
  Object describing part of a powder sample. The properties of a powder are:
        Attributes:
                     material : Crystal
                         an xrayutilties material (Crystal) describing optical and crystal properties of the powder
                     volume: float
                         powder's volume (in pseudo units, since only the relative volume enters the calculation)
                     crystallite_size_lor: float, optional
                         Lorentzian crystallite size fwhm (m)
                     crystallite_size_gauss: float, optional
                         Gaussian crystallite size fwhm (m)
                     strain lor: float, optional
                         extra peak width proportional to tan(theta)
                     strain_gauss : float, optional
                         extra peak width proportional to tan(theta)
                     preferred_orientation: tuple, optional
                         HKL of the preferred orientation
                     preferred orientation factor: float, optional
                         March-Dollase preferred orientation factor: > 1 for platy crystallits , < 1 for rod-like
                         crystallites, and = 1 for random orientation of crystallites.
class xrayutilities.simpack.smaterials.PowderList (name, *args)
  Bases: xrayutilities.simpack.smaterials.MaterialList
  extends the built in list type to enable building a list of Powder by various methods.
  check (V)
class xrayutilities.simpack.smaterials.PseudomorphicStack001 (name, *args)
  Bases: xrayutilities.simpack.smaterials.CrystalStack
  generate a sequence of pseudomorphic crystalline Layers. Surface orientation is assumed to be 001 and materials
  must be cubic/tetragonal.
  insert (i, v)
    S.insert(index, value) - insert value before index
 make_epitaxial (i)
  trans = <xrayutilities.math.transforms.Transform object>
class xrayutilities.simpack.smaterials.PseudomorphicStack111 (name, *args)
  Bases: xrayutilities.simpack.smaterials.PseudomorphicStack001
```

generate a sequence of pseudomorphic crystalline Layers. Surface orientation is assumed to be 111 and materials must be cubic.

trans = <xrayutilities.math.transforms.CoordinateTransform object>

class xrayutilities.simpack.smaterials.SMaterial (material, **kwargs)

Bases: object

Simulation Material. Extends the xrayutilities Materials by properties needed for simulations

property material

Module contents

simulation subpackage of xrayutilities.

This package provides possibilities to simulate X-ray diffraction and reflectivity curves of thin film samples. It could be extended for more general use in future if there is demand for that.

In addition it provides a fitting routine for reflectivity data which is based on Imfit.

Submodules

xrayutilities.config module

module to parse xrayutilities user-specific config file the parsed values are provide as global constants for the use in other parts of xrayutilities. The config file with the default constants is found in the python installation path of xrayutilities. It is however not recommended to change things there, instead the user-specific config file ~/.xrayutilities.conf or the local xrayutilities.conf file should be used.

xrayutilities.config.trytomake (obj, key, typefunc)

xrayutilities.exception module

xrayutilities derives its own exceptions which are raised upon wrong input when calling one of xrayutilities functions. none of the pre-defined exceptions is made for that purpose.

exception xrayutilities.exception.InputError (msg)

Bases: Exception

Exception raised for errors in the input. Either wrong datatype not handled by TypeError or missing mandatory keyword argument (Note that the obligation to give keyword arguments might depend on the value of the arguments itself)

Parameters: expr:str

input expression in which the error occurred

msg: str

explanation of the error

xrayutilities.experiment module

module helping with planning and analyzing experiments. various classes are provided for describing experimental geometries, calculationof angular coordinates of Bragg reflections, conversion of angular coordinates to Q-space and determination of powder diffraction peak positions.

The strength of the module is the versatile QConversion module which can be configured to describe almost any goniometer geometry.

class xrayutilities.experiment.Experiment (ipdir, ndir, **keyargs)

Bases: object

base class for describing experiments users should use the derived classes: HXRD, GID, PowderExperiment

Ang2HKL (*args, **kwargs)

angular to (h, k, l) space conversion. It will set the UB argument to Ang2Q and pass all other parameters unchanged. See Ang2Q for description of the rest of the arguments.

Parameters: args: list

arguments forwarded to Ang2Q

kwargs: dict, optional

optional keyword arguments

B: array-like, optional

reciprocal space conversion matrix of a Crystal. You can specify the matrix B (default identity matrix) shape needs to be (3, 3)

mat: Crystal, optional

Crystal object to use to obtain a B matrix (e.g. xu.materials.Si) can be used as alternative to the B keyword argument B is favored in case both are given

U: array-like, optional

orientation matrix U can be given. If none is given the orientation defined in the Experiment class is used.

dettype: {'point', 'linear', 'area'}, optional

detector type: decides which routine of Ang2Q to call. default 'point'

delta: ndarray, list or tuple, optional

giving delta angles to correct the given ones for misalignment. delta must be an numpy array or list of length 2. used angles are than (om, tt) - delta

wl: float or str, optional

x-ray wavelength in angstroem (default: self._wl)

en: float or str, optional

x-ray energy in eV (default: converted self._wl)

deg: bool, optional

flag to tell if angles are passed as degree (default: True)

sampledis: tuple, list or array-like, optional

sample displacement vector in relative units of the detector distance (default: (0, 0, 0))

Returns: ndarray

H K L coordinates as numpy.ndarray with shape (N, 3) where N corresponds to the number of points given in the input (args)

Q2Ang (qvec)

TiltAngle (q, deg=True)

TiltAngle(q, deg=True): Return the angle between a q-space position and the surface normal.

Parameters: q: list or numpy array with the reciprocal space position

optional keyword arguments:

deg: True/False whether the return value should be in degree or radians (default: True)

Transform (V)

transforms a vector, matrix or tensor of rank 4 (e.g. elasticity tensor) to the coordinate frame of the Experiment class. This is for example necessary before any Q2Ang-conversion can be performed.

Parameters: v: object to transform, list or numpy array of shape

(n,) (n, n), (n, n, n, n) where n is the rank of the transformation matrix

Returns: transformed object of the same shape as v

property energy

property wavelength

class xrayutilities.experiment.FourC (idir, ndir, **keyargs)

Bases: xrayutilities.experiment.HXRD

class describing high angle x-ray diffraction experiments the class helps with calculating the angles of Bragg reflections as well as helps with analyzing measured data

the class describes a four circle (omega, chi, phi, twotheta) goniometer to help with coplanar x-ray diffraction experiments. Nevertheless 3D data can be treated with the use of linear and area detectors. see help self.Ang2Q

class xrayutilities.experiment.GID (idir, ndir, **keyargs)

Bases: xrayutilities.experiment.Experiment

class describing grazing incidence x-ray diffraction experiments the class helps with calculating the angles of Bragg reflections as well as it helps with analyzing measured data

the class describes a four circle (alpha_i, azimuth, twotheta, beta) goniometer to help with GID experiments at the ROTATING ANODE. 3D data can be treated with the use of linear and area detectors. see help self.Ang2Q

Using this class the default sample surface orientation is determined by the inner most sample rotation (which is usually the azimuth motor).

Ang2Q (ai, phi, tt, beta, **kwargs)

angular to momentum space conversion for a point detector. Also see help GID.Ang2Q for procedures which treat line and area detectors

Parameters: ai, phi, tt, beta: float or array-like

sample and detector angles as numpy array, lists or Scalars must be given. All arguments must have the same shape or length. However, if one angle is always the same its enough to give one scalar value.

kwargs: dict, optional

optional keyword arguments

delta: list, tuple or array-like, optional

giving delta angles to correct the given ones for misalignment delta must be an numpy array or list of length 4. Used angles are then ai, phi, tt, beta - delta

UB: array-like, optional

matrix for conversion from (hkl) coordinates to Q of sample used to determine not Q but (hkl) (default: identity matrix)

wl: float or str, optional

x-ray wavelength in angstroem (default: self._wl)

deg: bool, optional

flag to tell if angles are passed as degree (default: True)

Returns: ndarray

reciprocal space positions as numpy.ndarray with shape (N, 3) where N corresponds to the number of points given in the input

Q2Ang (Q, trans=True, deg=True, **kwargs)

calculate the GID angles needed in the experiment the inplane reference direction defines the direction were the reference direction is parallel to the primary beam (i.e. lattice planes perpendicular to the beam)

Note

The behavior of this function is unchanged if the goniometer definition is changed!

Parameters: Q: list, tuple or array-like

array of shape (3) with q-space vector components or 3 separate lists with qx, qy, qz

trans: bool, optional

apply coordinate transformation on Q (default True)

deg: book, optional

(default True) determines if the angles are returned in radians or degrees

Returns: ndarray

a numpy array of shape (4) with four GID scattering angles which are [alpha_i, azimuth, twotheta, beta];

- alpha_i : incidence angle to surface (at the moment always 0)
- azimuth : sample rotation with respect to the inplane reference direction
- twotheta : scattering angle
- beta : exit angle from surface (at the moment always 0)

class xrayutilities.experiment.GISAXS (idir, ndir, **keyargs)

Bases: xrayutilities.experiment.Experiment

class describing grazing incidence x-ray diffraction experiments the class helps with calculating the angles of Bragg reflections as well as it helps with analyzing measured data

the class describes a three circle (alpha_i, twotheta, beta) goniometer to help with GISAXS experiments at the ROTATING ANODE. 3D data can be treated with the use of linear and area detectors. see help self.Ang2Q

Ang2Q (ai, tt, beta, **kwargs)

angular to momentum space conversion for a point detector. Also see help GISAXS.Ang2Q for procedures which treat line and area detectors

Parameters: ai, tt, beta: float or array-like

sample and detector angles as numpy array, lists or Scalars must be given. all arguments must have the same shape or length. Howevver, if one angle is always the same its enough to give one scalar value.

kwargs: dict, optional

optional keyword arguments

delta: list, tuple or array-like, optional

giving delta angles to correct the given ones for misalignment delta must be an numpy array or list of length 3. Used angles are then ai, tt, beta - delta

UB: array-like, optional

matrix for conversion from (hkl) coordinates to Q of sample used to determine not Q but (hkl) (default: identity matrix)

wl: float or str, optional

x-ray wavelength in angstroem (default: self._wl)

deg: bool, optional

flag to tell if angles are passed as degree (default: True)

Returns: ndarray

reciprocal space positions as numpy.ndarray with shape (N, 3) where N corresponds to the number of points given in the input

Q2Ang (Q, trans=True, deg=True, **kwargs)

class xrayutilities.experiment.HXRD (idir, ndir, geometry='hi_lo', **keyargs)

Bases: xrayutilities.experiment.Experiment

class describing high angle x-ray diffraction experiments the class helps with calculating the angles of Bragg reflections as well as helps with analyzing measured data

the class describes a two circle (omega, twotheta) goniometer to help with coplanar x-ray diffraction experiments. Nevertheless 3D data can be treated with the use of linear and area detectors. see help self.Ang2Q

Ang2Q (om, tt, **kwargs)

angular to momentum space conversion for a point detector. Also see help HXRD.Ang2Q for procedures which treat line and area detectors

Parameters: om, tt : float or array-like

sample and detector angles as numpy array, lists or Scalars must be given. All arguments must have the same shape or length. However, if one angle is always the same its enough to give one scalar value.

kwargs: dict, optional

optional keyword arguments

delta: list or array-like

giving delta angles to correct the given ones for misalignment. delta must be an numpy array or list of length 2. Used angles are than om, tt - delta

UB: array-like

matrix for conversion from (hkl) coordinates to Q of sample used to determine not Q but (hkl) (default: identity matrix)

wl: float or str, optional

x-ray wavelength in angstroem (default: self._wl)

deg: bool, optional

flag to tell if angles are passed as degree (default: True)

Returns: ndarray

reciprocal space positions as numpy.ndarray with shape (N, 3) where N corresponds to the number of points given in the input

Q2Ang (*Q, **keyargs)

Convert a reciprocal space vector Q to COPLANAR scattering angles. The keyword argument trans determines whether Q should be transformed to the experimental coordinate frame or not. The coplanar scattering angles correspond to a goniometer with sample rotations ['x+', 'y+', 'z-'] and detector rotation 'x+' and primary beam along y. This is a standard four circle diffractometer.

Note

The behavior of this function is unchanged if the goniometer definition is changed!

Parameters: Q: list, tuple or array-like

array of shape (3) with q-space vector components or 3 separate lists with qx, qy, qz

trans: bool, optional

apply coordinate transformation on Q (default True)

deg: book, optional

(default True) determines if the angles are returned in radians or degrees

geometry: {'hi_lo', 'lo_hi', 'real', 'realTilt'}, optional

determines the scattering geometry (default: self.geometry):

- 'hi_lo' high incidence and low exit
- 'lo_hi' low incidence and high exit
- 'real' general geometry with angles determined by q-coordinates (azimuth); this and upper geometries return [omega, 0, phi, twotheta]
- 'realTilt' general geometry with angles determined by q-coordinates (tilt); returns [omega, chi, phi, twotheta]

refrac: bool, optional

determines if refraction is taken into account; if True then also a material must be given (default: False)

mat: Crystal

Crystal object; needed to obtain its optical properties for refraction correction, otherwise not used

full_output : bool, optional

determines if additional output is given to determine scattering angles more accurately in case refraction is set to True. default: False

fi, fd: tuple or list

if refraction correction is applied one can optionally specify the facet through which the beam enters (fi) and exits (fd) fi, fd must be the surface normal vectors (not transformed & not necessarily normalized). If omitted the normal direction of the experiment is used.

Returns: ndarray

full_output=False: a numpy array of shape (4) with four scattering angles which are [omega, chi, phi, twotheta];

- omega: incidence angle with respect to surface
- chi : sample tilt for the case of non-coplanar geometry
- phi : sample azimuth with respect to inplane reference direction
- twotheta: scattering angle/detector angle

full_output=True: a numpy array of shape (6) with five angles which are [omega, chi, phi, twotheta, psi_i, psi_d]

- psi_i : offset of the incidence beam from the scattering plane due to refraction
- pdi_d : offset ot the diffracted beam from the scattering plane due to refraction

class xrayutilities.experiment.NonCOP (idir, ndir, **keyargs)

Bases: xrayutilities.experiment.Experiment

class describing high angle x-ray diffraction experiments. The class helps with calculating the angles of Bragg reflections as well as helps with analyzing measured data for NON-COPLANAR measurements, where the tilt is used to align asymmetric peaks, like in the case of a polefigure measurement.

The class describes a four circle (omega, chi, phi, twotheta) goniometer to help with x-ray diffraction experiments. Linear and area detectors can be treated as described in "help self.Ang2Q"

Ang2Q (om, chi, phi, tt, **kwargs)

angular to momentum space conversion for a point detector. Also see help NonCOP.Ang2Q for procedures which treat line and area detectors

Parameters: om, chi, phi, tt: float or array-like

sample and detector angles as numpy array, lists or Scalars must be given. All arguments must have the same shape or length. However, if one angle is always the same its enough to give one scalar value.

kwargs: dict, optional

optional keyword arguments

delta: list, tuple or array-like, optional

giving delta angles to correct the given ones for misalignment delta must be an numpy array or list of length 4. Used angles are than om, chi, phi, tt - delta

UB: array-like, optional

matrix for conversion from (hkl) coordinates to Q of sample used to determine not Q but (hkl) (default: identity matrix)

wl: float or str, optional

x-ray wavelength in angstroem (default: self._wl)

deg: bool, optional

flag to tell if angles are passed as degree (default: True)

Returns: ndarray

reciprocal space positions as numpy.ndarray with shape (N, 3) where N corresponds to the number of points given in the input

Q2Ang (*Q, **keyargs)

Convert a reciprocal space vector Q to NON-COPLANAR scattering angles. The keyword argument trans determines whether Q should be transformed to the experimental coordinate frame or not.

Note

The behavior of this function is unchanged if the goniometer definition is changed!

Parameters: Q: list, tuple or array-like

array of shape (3) with q-space vector components or 3 separate lists with qx, qy, qz

trans: bool, optional

apply coordinate transformation on Q (default True)

deg: book, optional

(default True) determines if the angles are returned in radians or degrees

Returns: ndarray

a numpy array of shape (4) with four scattering angles which are [omega, chi, phi, twotheta];

- · omega: incidence angle with respect to surface
- · chi : sample tilt for the case of non-coplanar geometry
- phi : sample azimuth with respect to inplane reference direction
- twotheta: scattering angle/detector angle

class xrayutilities.experiment.PowderExperiment (**kwargs)

Bases: xrayutilities.experiment.Experiment

Experimental class for powder diffraction which helps to convert theta angles to momentum transfer space

Q2Ang (qpos, wl=None, deg=True)

Converts reciprocal space values to theta angles

class xrayutilities.experiment.QConversion (sampleAxis, detectorAxis, r_i, **kwargs)

Bases: object

Class for the conversion of angular coordinates to momentum space for arbitrary goniometer geometries and X-ray energy. Both angular scans (where some goniometer angles change during data acquisition) and energy scans (where the energy is varied during acquisition) as well as mixed cases can be treated.

the class is configured with the initialization and does provide three distinct routines for conversion to momentum space for

- point detector: point(...) or __call__()
- linear detector: linear(...)

• area detector: area(...) linear() and area() can only be used after the init_linear() or init_area() routines were called

property UB

area (*args, **kwargs)

angular to momentum space conversion for a area detector the center pixel defined by the init_area routine must be in direction of self.r_i when detector angles are zero

the detector geometry must be initialized by the init_area(...) routine

Parameters: args: ndarray, list or Scalars

sample and detector angles; in total *len(self.sampleAxis)* + *len(detectorAxis)* must be given, always starting with the outer most circle. all arguments must have the same shape or length but can be mixed with Scalars (i.e. if an angle is always the same it can be given only once instead of an array)

• sAngles :

sample circle angles, number of arguments must correspond to len(self.sampleAxis)

• dAngles:

detector circle angles, number of arguments must correspond to len(self.detectorAxis)

kwargs: dict, optional

optional keyword arguments

delta: list or array-like, optional

delta angles to correct the given ones for misalignment. delta must be an numpy array or list of len(*args). used angles are then *args - delta

UB: array-like, optional

matrix for conversion from (hkl) coordinates to Q of sample used to determine not Q but (hkl) (default: self.UB)

Nav: tuple or list, optional

number of channels to average to reduce data size e.g. [2, 2] (default: self._area_nav)

roi: list or tuple, optional

region of interest for the detector pixels; e.g. [100, 900, 200, 800] (default: self._area_roi)

wl: float or str, optional

x-ray wavelength in angstroem (default: self._wl)

en: float, optional

x-ray energy in eV (default is converted self._wl). both wavelength and energy can also be an array which enables the QConversion for energy scans. Note that the *en* keyword overrules the *wl* keyword!

deq: bool, optional

flag to tell if angles are passed as degree (default: True)

sampledis: tuple or list or array-like

sample displacement vector in relative units of the detector distance (default: (0, 0, 0))

Returns: reciprocal space position of all detector pixels in a numpy.ndarray of

shape ((*)*(self._area_roi[1] - self._area_roi[0]+1) *

(self._area_roi[3] - self._area_roi[2] + 1), 3) were detectorDir1 is

the fastest varing

property detectorAxis

property handler for _detectorAxis

Returns: list of detector axis following the syntax /[xyz][+-]/

property energy

getDetectorDistance (*args, **kwargs)

obtains the detector distance by applying the detector arm movements. This is especially interesting for the case of 1 or 2D detectors to perform certain geometric corrections.

Parameters: args: list

detector angles. Only detector arm angles as described by the detectorAxis attribute must be given.

kwargs : dict, optional

optional keyword arguments

dim: int, optional

dimension of the detector for which the position should be determined

roi: tuple or list, optional

region of interest for the detector pixels; (default: self._area_roi/self._linear_roi)

Nav: tuple or list, optional

number of channels to average to reduce data size; (default: self._area_nav/self._linear_nav)

deg: bool, optional

flag to tell if angles are passed as degree (default: True)

Returns: ndarray

numpy array with the detector distance

getDetectorPos (*args, **kwargs)

obtains the detector position vector by applying the detector arm rotations.

Parameters: args: list

detector angles. Only detector arm angles as described by the detectorAxis attribute must be given.

kwargs: dict, optional

optional keyword arguments

dim: int, optional

dimension of the detector for which the position should be determined

roi: tuple or list, optional

region of interest for the detector pixels; (default: self._area_roi/self._linear_roi)

Nav: tuple or list, optional

number of channels to average to reduce data size; (default: self._area_nav/self._linear_nav)

deg: bool, optional

flag to tell if angles are passed as degree (default: True)

Returns: ndarray

numpy array of length 3 with vector components of the detector direction. The length of the vector is k.

init_area (detectorDir1, detectorDir2, cch1, cch2, Nch1, Nch2, distance=None, pwidth1=None, pwidth2=None, chpdeg1=None, chpdeg2=None, detrot=0, tiltazimuth=0, tilt=0, **kwargs)

initialization routine for area detectors detector direction as well as distance and pixel size or channels per degree must be given. Two separate pixel sizes and channels per degree for the two orthogonal directions can be given Parameters: detectorDir1: str

direction of the detector (along the pixel direction 1); e.g. 'z+' means higher pixel

numbers at larger z positions

detectorDir2: str

direction of the detector (along the pixel direction 2); e.g. 'x+'

cch1, cch2: float

center pixel, in direction of self.r_i at zero detectorAngles

Nch1, Nch2: int

number of detector pixels along direction 1, 2

distance: float, optional

distance of center pixel from center of rotation

pwidth1, pwidth2: float, optional

width of one pixel (same unit as distance)

chpdeg1, chpdeg2: float, optional

channels per degree (only absolute value is relevant) sign determined through detector Dir1, detector Dir2

detrot : float, optional

angle of the detector rotation around primary beam direction (used to correct misalignments)

tiltazimuth: float, optional

direction of the tilt vector in the detector plane (in degree)

tilt: float, optional

tilt of the detector plane around an axis normal to the direction given by the tiltazimuth

kwargs: dict, optional

optional keyword arguments

Nav: tuple or list, optional

number of channels to average to reduce data size (default: [1, 1])

roi: tuple or list, optional

region of interest for the detector pixels; e.g. [100, 900, 200, 800]

Note

Either distance and pwidth1, pwidth2 or chpdeg1, chpdeg2 must be given!!

Note

the channel numbers run from 0 .. NchX-1

init_linear (detectorDir, cch, Nchannel, distance=None, pixelwidth=None, chpdeg=None, tilt=0, **kwargs) initialization routine for linear detectors detector direction as well as distance and pixel size or channels per degree must be given.

Parameters: detectorDir: str

direction of the detector (along the pixel array); e.g. 'z+'

cch: float

center channel, in direction of self.r_i at zero detectorAngles

Nchannel: int

total number of detector channels

distance: float, optional

distance of center channel from center of rotation

pixelwidth: float, optional

width of one pixel (same unit as distance)

chpdeg: float, optional

channels per degree (only absolute value is relevant) sign determined through

detectorDir

tilt: float, optional

tilt of the detector axis from the detectorDir (in degree)

kwargs: dict, optional

optional keyword arguments

Nav: int, optional

number of channels to average to reduce data size (default: 1)

roi: tuple or list

region of interest for the detector pixels; e.g. [100, 900]

Note

Either distance and pixelwidth or chpdeg must be given!!

Note

the channel numbers run from 0 .. Nchannel-1

linear (*args, **kwargs)

angular to momentum space conversion for a linear detector the cch of the detector must be in direction of self.r_i when detector angles are zero

the detector geometry must be initialized by the init_linear(...) routine

Parameters: args: ndarray, list or Scalars

sample and detector angles; in total *len(self.sampleAxis)* + *len(detectorAxis)* must be given, always starting with the outer most circle. all arguments must have the same shape or length but can be mixed with Scalars (i.e. if an angle is always the same it can be given only once instead of an array)

• sAngles:

sample circle angles, number of arguments must correspond to len(self.sampleAxis)

• dAngles:

detector circle angles, number of arguments must correspond to len(self.detectorAxis)

kwargs: dict, optional

optional keyword arguments

delta: list or array-like, optional

delta angles to correct the given ones for misalignment. delta must be an numpy array or list of len(*args). used angles are then *args - delta

UB: array-like, optional

matrix for conversion from (hkl) coordinates to Q of sample used to determine not Q but (hkl) (default: self.UB)

Nav: int, optional

number of channels to average to reduce data size (default: self._linear_nav)

roi: list or tuple, optional

region of interest for the detector pixels; e.g. [100, 900] (default: self._linear_roi)

wl: float or str, optional

x-ray wavelength in angstroem (default: self._wl)

en: float, optional

x-ray energy in eV (default is converted self._wl). both wavelength and energy can also be an array which enables the QConversion for energy scans. Note that the *en* keyword overrules the *wl* keyword!

deg: bool, optional

flag to tell if angles are passed as degree (default: True)

sampledis: tuple or list or array-like

sample displacement vector in relative units of the detector distance (default: (0, 0, 0))

Returns: reciprocal space position of all detector pixels in a numpy.ndarray of

shape ((*)*(self._linear_roi[1]-self._linear_roi[0]+1) , 3)

point (*args, **kwargs)

angular to momentum space conversion for a point detector located in direction of self.r_i when detector angles are zero

Parameters: args: ndarray, list or Scalars

sample and detector angles; in total *len(self.sampleAxis)* + *len(detectorAxis)* must be given, always starting with the outer most circle. all arguments must have the same shape or length but can be mixed with Scalars (i.e. if an angle is always the same it can be given only once instead of an array)

• sAngles:

sample circle angles, number of arguments must correspond to len(self.sampleAxis)

• dAngles:

detector circle angles, number of arguments must correspond to len(self.detectorAxis)

kwargs: dict, optional

optional keyword arguments

delta: list or array-like, optional

delta angles to correct the given ones for misalignment. delta must be an numpy array or list of len(*args). used angles are then *args - delta

UB: array-like, optional

matrix for conversion from (hkl) coordinates to Q of sample used to determine not Q but (hkl) (default: self.UB)

wl: float or str, optional

x-ray wavelength in angstroem (default: self._wl)

en: float, optional

x-ray energy in eV (default is converted self._wl). both wavelength and energy can also be an array which enables the QConversion for energy scans. Note that the *en* keyword overrules the *wl* keyword!

deg: bool, optional

flag to tell if angles are passed as degree (default: True)

sampledis: tuple or list or array-like

sample displacement vector in relative units of the detector distance (default: (0, 0, 0))

Returns: ndarray

reciprocal space positions as numpy.ndarray with shape (N, 3) where N corresponds to the number of points given in the input

property sampleAxis

property handler for _sampleAxis

Returns: list

sample axes following the syntax /[xyzk][+-]/

transformSample2Lab (vector, *args)

transforms a vector from the sample coordinate frame to the laboratory coordinate system by applying the sample rotations from inner to outer circle.

Parameters: vector: sequence, list or numpy array

vector to transform

args : list

goniometer angles (sample angles or full goniometer angles can be given. If more angles than the sample circles are given they will be ignored)

Returns: ndarray

rotated vector as numpy.array

property wavelength

xrayutilities.gridder module

class xrayutilities.gridder.FuzzyGridder1D (nx)

Bases: xrayutilities.gridder.Gridder1D

An 1D binning class considering every data point to have a finite width. If necessary one data point will be split fractionally over different data bins. This is numerically more effort but represents better the typical case of a experimental data, which do not represent a mathematical point but have a finite width (e.g. X-ray data from a 1D detector).

class xrayutilities.gridder.Gridder

Bases: abc.ABC

Basis class for gridders in xrayutilities. A gridder is a function mapping irregular spaced data onto a regular grid by binning the data into equally sized elements.

There are different ways of defining the regular grid of a Gridder. In xrayutilities the data bins extend beyond the data range in the input data, but the given position being the center of these bins, extends from the minimum to the maximum of the data! The main motivation for this was to create a Gridder, which when feeded with N equidistant data points and gridded with N bins would not change the data position (not the case with numpy.histogramm functions!). Of course this leads to the fact that for homogeneous point density the first and last bin in any direction are not filled as the other bins.

A different definition is used by numpy histogram functions where the bins extend only to the end of the data range. (see numpy histogram, histogram2d, ...)

Clear ()

Clear so far gridded data to reuse this instance of the Gridder

KeepData (bool)

Normalize (bool)

set or unset the normalization flag. Normalization needs to be done to obtain proper gridding but may want to be disabled in certain cases when sequential gridding is performed

property data

return gridded data (performs normalization if switched on)

class xrayutilities.gridder.Gridder1D (nx)
Bases: xrayutilities.gridder.Gridder

dataRange (min, max, fixed=True)

define minimum and maximum data range, usually this is deduced from the given data automatically, however, for sequential gridding it is useful to set this before the first call of the gridder. data outside the range are simply ignored

Parameters: min: float

minimum value of the gridding range

max: float

maximum value of the gridding range

fixed: bool, optional

flag to turn fixed range gridding on (True (default)) or off (False)

savetxt (filename, header=")

save gridded data to a txt file with two columns. The first column is the data coordinate and the second the corresponding data value

Parameters: filename: str

output filename **header**: str, optional

optional header for the data file.

property xaxis

Returns the xaxis of the gridder the returned values correspond to the center of the data bins used by the gridding algorithm

xrayutilities.gridder.axis (min_value, max_value, n)
Compute the a grid axis.

Parameters: min_value : float

axis minimum value

max value: float

axis maximum value

n:int

number of steps

xrayutilities.gridder.delta (min_value, max_value, n)

Compute the stepsize along an axis of a grid.

Parameters: min_value : axis minimum value

max_value: axis maximum value

n: number of steps

class xrayutilities.gridder.npyGridder1D (nx)
Bases: xrayutilities.gridder.Gridder1D

property xaxis

Returns the xaxis of the gridder the returned values correspond to the center of the data bins used by the numpy.histogram function

xrayutilities.gridder.ones (*args)

Compute ones for matrix generation. The shape is determined by the number of input arguments.

xrayutilities.gridder2d module

class xrayutilities.gridder2d.FuzzyGridder2D (nx, ny)

Bases: xrayutilities.gridder2d.Gridder2D

An 2D binning class considering every data point to have a finite area. If necessary one data point will be split fractionally over different data bins. This is numerically more effort but represents better the typical case of a experimental data, which do not represent a mathematical point but have a finite size (e.g. X-ray data from a 2D detector or reciprocal space maps measured with point/linear detector).

Currently only a rectangular area can be considered during the gridding.

class xrayutilities.gridder2d.Gridder2D (nx, ny)

Bases: xrayutilities.gridder.Gridder

SetResolution (nx, ny)

Reset the resolution of the gridder. In this case the original data stored in the object will be deleted.

Parameters: nx: int

number of points in x-direction

ny: int

number of points in y-direction

dataRange (xmin, xmax, ymin, ymax, fixed=True)

define minimum and maximum data range, usually this is deduced from the given data automatically, however, for sequential gridding it is useful to set this before the first call of the gridder. data outside the range are simply ignored

Parameters: xmin, ymin : float

minimum value of the gridding range in x, y

xmax, ymax: float

maximum value of the gridding range in x, y

fixed: bool, optional

flag to turn fixed range gridding on (True (default)) or off (False)

savetxt (filename, header=")

save gridded data to a txt file with two columns. The first two columns are the data coordinates and the last one the corresponding data value.

Parameters: filename: str

output filename **header**: str, optional

optional header for the data file.

property xaxis

property xmatrix

property yaxis

property ymatrix

class xrayutilities.gridder2d.Gridder2DList (nx, ny)

Bases: xrayutilities.gridder2d.Gridder2D

special version of a 2D gridder which performs no actual averaging of the data in one grid/bin but just collects the data-objects belonging to one bin for further treatment by the user

Clear ()

Clear so far gridded data to reuse this instance of the Gridder

property data

return gridded data, in this special version no normalization is defined!

xrayutilities.gridder3d module

```
class xrayutilities.gridder3d.FuzzyGridder3D (nx, ny, nz)
```

Bases: xrayutilities.gridder3d.Gridder3D

An 3D binning class considering every data point to have a finite volume. If necessary one data point will be split fractionally over different data bins. This is numerically more effort but represents better the typical case of a experimental data, which do not represent a mathematical point but have a finite size.

Currently only a quader can be considered as volume during the gridding.

```
class xrayutilities.gridder3d.Gridder3D (nx, ny, nz)
```

Bases: xrayutilities.gridder.Gridder

SetResolution (nx, ny, nz)

dataRange (xmin, xmax, ymin, ymax, zmin, zmax, fixed=True)

define minimum and maximum data range, usually this is deduced from the given data automatically, however, for sequential gridding it is useful to set this before the first call of the gridder. data outside the range are simply ignored

```
Parameters: xmin, ymin, zmin: float
    minimum value of the gridding range in x, y, z
    xmax, ymax, zmax: float
    maximum value of the gridding range in x, y, z
    fixed: bool, optional
    flag to turn fixed range gridding on (True (default)) or off (False)

property xaxis

property ymatrix

property ymatrix

property zaxis

property zmatrix
```

xrayutilities.mpl_helper module

Defines new matplotlib Sqrt scale which further allows for negative values by using the sign of the original value as sign of the plotted value.

```
class xrayutilities.mpl_helper.SqrtAllowNegScale (axis, **kwargs)
 Bases: matplotlib.scale.ScaleBase
 Scales data using a sqrt-function, however, allowing also negative values.
 The scale function:
     sign(y) * sqrt(abs(y))
 The inverse scale function:
     sign(y) * y**2
 class InvertedSqrtTransform (shorthand_name=None)
   Bases: matplotlib.transforms.Transform
   has_inverse = True
   input_dims = 1
    inverted()
     Return the corresponding inverse transformation.
     It holds x == self.inverted().transform(self.transform(x)).
     The return value of this method should be treated as temporary. An update to self does not cause a
     corresponding update to its inverted copy.
   is_separable = True
   output_dims = 1
   transform_non_affine (a)
     Apply only the non-affine part of this transformation.
     transform(values)
                                                         always
                                                                              equivalent
                                                                                                      to
     transform_affine(transform_non_affine(values)).
     In non-affine transformations, this is generally equivalent to transform(values). In affine transformations,
     this is always a no-op.
```

```
Parameters: values : array
```

The input values as NumPy array of length input_dims or shape (N x

input_dims).

Returns: array

The output values as NumPy array of length input_dims or shape (N x output_dims), depending on the input.

```
class SqrtTransform (shorthand_name=None)
```

Bases: matplotlib.transforms.Transform

```
has_inverse = True
input_dims = 1
```

inverted()

return the inverse transform for this transform.

```
is_separable = True
output_dims = 1
```

transform_non_affine (a)

This transform takes an Nx1 numpy array and returns a transformed copy.

get_transform()

Return the **Transform** object associated with this scale.

limit_range_for_scale (vmin, vmax, minpos)

Override to limit the bounds of the axis to the domain of the transform. In the case of Mercator, the bounds should be limited to the threshold that was passed in. Unlike the autoscaling provided by the tick locators, this range limiting will always be adhered to, whether the axis range is set manually, determined automatically or changed through panning and zooming.

```
name = 'sqrt'
```

set_default_locators_and_formatters (axis)

Set the locators and formatters of axis to instances suitable for this scale.

```
class xrayutilities.mpl_helper.SqrtTickLocator (nbins=7, symmetric=True)
Bases: matplotlib.ticker.Locator
```

set_params (nbins, symmetric)

Set parameters within this locator.

tick_values (vmin, vmax)

Return the values of the located ticks given **vmin** and **vmax**.

Note

To get tick locations with the vmin and vmax values defined automatically for the associated axis simply call the Locator instance:

```
>>> print(type(loc))
<type 'Locator'>
>>> print(loc())
[1, 2, 3, 4]
```

view_limits (dmin, dmax)

Set the view limits to the nearest multiples of base that contain the data

xrayutilities.normalize module

module to provide functions that perform block averaging of intensity arrays to reduce the amount of data (mainly for PSD and CCD measurements

and

provide functions for normalizing intensities for

- · count time
- absorber (user-defined function)
- monitor
- flatfield correction

class xrayutilities.normalize.IntensityNormalizer (det=", **keyargs)

Bases: object

generic class for correction of intensity (point detector, or MCA, single CCD frames) for count time and absorber factors the class must be supplied with a absorber correction function and works with data structures provided by xrayutilities.io classes or the corresponding objects from hdf5 files

property absfun

absfun property handler returns the costum correction function or None

property avmon

av_mon property handler

returns the value of the average monitor or None if average is calculated from the monitor field

property darkfield

flatfield property handler

returns the current set darkfield of the detector or None if not set

property det

det property handler

returns the detector field name

property flatfield

flatfield property handler

returns the current set flatfield of the detector or None if not set

property mon

mon property handler

returns the monitor field name or None if not set

property time

time property handler

returns the count time or the field name of the count time or None if time is not set

xrayutilities.normalize.blockAverage1D (data, Nav)

perform block average for 1D array/list of Scalar values all data are used. at the end of the array a smaller cell may be used by the averaging algorithm

Parameters: data: array-like

data which should be contracted (length N)

Nav: int

number of values which should be averaged

Returns: ndarray

block averaged numpy array of data type numpy.double (length ceil(N/Nav))

xrayutilities.normalize.blockAverage2D (data2d, Nav1, Nav2, **kwargs)

perform a block average for 2D array of Scalar values all data are used therefore the margin cells may differ in size

Parameters: data2d : ndarray

array of 2D data shape (N, M)

Nav1, Nav2: int

a field of (Nav1 x Nav2) values is contracted

kwargs: dict, optional

optional keyword argument

roi: tuple or list, optional

region of interest for the 2D array. e.g. [20, 980, 40, 960], reduces M, and M!

Returns: ndarray

block averaged numpy array with type numpy.double with shape (ceil(N/Nav1), ceil(M/Nav2))

xrayutilities.normalize.blockAverageCCD (data3d, Nav1, Nav2, **kwargs)

perform a block average for 2D frames inside a 3D array. all data are used therefore the margin cells may differ in size

Parameters: data3d: ndarray

array of 3D data shape (Nframes, N, M)

Nav1, Nav2: int

a field of (Nav1 x Nav2) values is contracted

kwargs: dict, optional

optional keyword argument

roi: tuple or list, optional

region of interest for the 2D array. e.g. [20, 980, 40, 960], reduces M, and M!

Returns: ndarray

block averaged numpy array with type numpy.double with shape (Nframes, ceil(N/Nav1), ceil(M/Nav2))

xrayutilities.normalize.blockAveragePSD (psddata, Nav, **kwargs)

perform a block average for serveral PSD spectra all data are used therefore the last cell used for averaging may differ in size

Parameters: psddata: ndarray

array of 2D data shape (Nspectra, Nchannels)

Nav: int

number of channels which should be averaged

kwargs: dict, optional

optional keyword argument

roi: tuple or list

region of interest for the 2D array. e.g. [20, 980] Nchannels = 980-20

Returns: ndarray

block averaged psd spectra as numpy array with type numpy.double of shape (Nspectra , ceil(Nchannels/Nav))

xrayutilities.q2ang_fit module

Module provides functions to convert a q-vector from reciprocal space to angular space. a simple implementation uses scipy optimize routines to perform a fit for a arbitrary goniometer.

The user is, however, expected to use the bounds variable to put restrictions to the number of free angles to obtain reproducible results. In general only 3 angles are needed to fit an arbitrary q-vector (2 sample + 1 detector angles or

1 sample + 2 detector). More complicated restrictions can be implemented using the lmfit package. (done upon request!)

The function is based on a fitting routine. For a specific goniometer also analytic expressions from literature can be used as they are implemented in the predefined experimental classes HXRD, NonCOP, and GID.

xrayutilities.q2ang_fit.Q2AngFit (qvec, expclass, bounds=None, ormat=array([[1.0, 0.0, 0.0], [0.0, 1.0, 0.0], [0.0, 0.0, 1.0]]), startvalues=None, constraints=[])

Functions to convert a q-vector from reciprocal space to angular space. This implementation uses scipy optimize routines to perform a fit for a goniometer with arbitrary number of goniometer angles.

The user *must* use the bounds variable to put restrictions to the number of free angles to obtain reproducible results. In general only 3 angles are needed to fit an arbitrary q-vector (2 sample + 1 detector angles or 1 sample + 2 detector).

Parameters: qvec: tuple or list or array-like

q-vector for which the angular positions should be calculated

expclass: Experiment

experimental class used to define the goniometer for which the angles should be calculated.

bounds: tuple or list

bounds of the goniometer angles. The number of bounds must correspond to the number of goniometer angles in the expclass. Angles can also be fixed by supplying only one value for a particular angle. e.g.: ((low, up), fix, (low2, up2), (low3, up3))

ormat: array-like

orientation matrix of the sample to be used in the conversion

startvalues : array-like

start values for the fit, which can significantly speed up the conversion. The number of values must correspond to the number of angles in the goniometer of the expclass

constraints: list

sequence of constraint dictionaries. This allows applying arbitrary (e.g. pseudo-angle) contraints by supplying according constraint functions. An entry of the constraints argument must be a dictionary with at least the 'type' and 'fun' set. 'type' can be either 'eq' or 'ineq' for equality or inequality constraints. 'fun' must be a callable function which for 'eq'-constraints returns 0 when the equality condition is fulfilled (see constraints documentation in scipy.optimize.minimize for details). The supplied function will be called with the arguments gonimeter angle list as argument. Typically this means you will have to use a lambda function.

Returns: fittedangles: list

list of fitted goniometer angles

qerror: float

error in reciprocal space

errcode: int

error-code of the scipy minimize function. for a successful fit the error code should be <=2

xrayutilities.q2ang_fit.exitAngleConst (angles, alphaf, xrd)

helper function for an pseudo-angle constraint of the exit angle. Can be used together with the Q2AngFit-routine in the 'constraints' argument. An example use case scenario to fix the exit angle to 1 degree would be: constraints={'type': 'eq', 'fun': lambda a: exitAngleConst(a, 1, xrd)}

Parameters: angles: iterable

fit parameters of Q2AngFit

alphaf: float

the exit angle which should be fixed

xrd: Experiment

the Experiment object to use for qconversion

```
xrayutilities.q2ang_fit.incidenceAngleConst (angles, alphai, xrd)
```

helper function for an pseudo-angle constraint of the incidence angle. Can be used together with the Q2AngFit-routine in the 'constraints' argument. An example use case scenario to fix the incidence angle to 1 degree would be: constraints={'type': 'eq', 'fun': lambda a: incidenceAngleConst(a, 1, xrd)}

Parameters: angles: iterable

fit parameters of Q2AngFit

alphai: float

the incidence angle which should be fixed

xrd: Experiment

the Experiment object to use for qconversion

xrayutilities.utilities module

xrayutilities utilities contains a conglomeration of useful functions which do not fit into one of the other files

xrayutilities.utilities.frac2str (f, denominator_limit=25, fmt='%7.4f') convert a float to a string attempting to represent it as a fraction

Parameters: f: float

floating point number to be represented as string

denominator_limit : int

maximal integer used as denominator. If f can't be expressed (within xu.config.EPSILON) by a fraction with a denominator up to this number a floating point string will be returned

fmt:str

format string used in case a floating point representation is needed

Returns: str

xrayutilities.utilities.import_lmfit (funcname='XU')

lazy import function for Imfit

xrayutilities.utilities.import_matplotlib_pyplot (funcname='XU') lazy import function of matplotlib.pyplot

Parameters: funcname: str

identification string of the calling function

Returns: flag: bool

the flag is True if the loading was successful and False otherwise.

pyplot

On success pyplot is the matplotlib.pyplot package.

xrayutilities.utilities.import_mayavi_mlab (funcname='XU')

lazy import function of mayavi.mlab

Parameters: funcname: str

identification string of the calling function

Returns: flag: bool

the flag is True if the loading was successful and False otherwise.

mlab

On success mlab is the mayavi.mlab package.

xrayutilities.utilities.maplog (inte, dynlow='config', dynhigh='config')

clips values smaller and larger as the given bounds and returns the log10 of the input array. The bounds are given as exponent with base 10 with respect to the maximum in the input array. The function is implemented in analogy to J. Stangl's matlab implementation.

Parameters: inte: ndarray

numpy.array, values to be cut in range

dynlow: float, optional

10^(-dynlow) will be the minimum cut off

dynhigh: float, optional

10^(-dynhigh) will be the maximum cut off

Returns: ndarray

numpy.array of the same shape as inte, where values smaller/larger than 10^(-dynlow, dynhigh) were replaced by 10^(-dynlow, dynhigh)

Examples

>>> lint = maplog(int, 5, 2)

xrayutilities.utilities_noconf module

xrayutilities utilities contains a conglomeration of useful functions this part of utilities does not need the config class

class xrayutilities.utilities_noconf.ABC

Bases: object

Helper class that provides a standard way to create an ABC using inheritance.

xrayutilities_utilities_noconf.check_kwargs (kwargs, valid_kwargs, identifier)

Raises an TypeError if kwargs included a key which is not in valid_kwargs.

Parameters: kwargs: dict

keyword arguments dictionary

valid_kwargs: dict

dictionary with valid keyword arguments and their description

identifier: str

string to identifier the caller of this function

xrayutilities.utilities_noconf.clear_bit (f, Offset)

clears the bet at an offset

xrayutilities.utilities_noconf.en2lam (inp)

converts the input energy in eV to a wavelength in Angstrom

Parameters: inp: float or str

energy in eV

Returns: float

wavlength in Angstrom

Examples

>>> wavelength = en2lam(8048)

xrayutilities.utilities_noconf.energy (en)

convert common energy names to energies in eV

so far this works with CuKa1, CuKa2, CuKa12, CuKb, MoKa1

Parameters: en : float, array-like or str

energy either as scalar or array with value in eV, which will be returned unchanged; or

string with name of emission line

Returns: float or array-like

energy in eV

xrayutilities_utilities_noconf.exchange_filepath (orig, new, keep=0, replace=None)

function to exchange the root of a filename with the option of keeping the inner directory structure. This for example includes such a conversion /dir_a/subdir/sample/file.txt -> /home/user/data/sample/file.txt where the

innermost directory name is kept (keep=1), or equally the three outer most are replaced (replace=3). One can either give keep, or replace, with replace taking preference if both are given. Note that replace=1 on Linux/Unix replaces only the root for absolute paths.

Parameters: orig: str

original filename which should have its data root replaced

new: str

new path which should be used instead

keep: int, optional

number of inner most directory names which should be kept the same in the output (default = 0)

replace: int, optional

number of outer most directory names which should be replaced in the output (default =

Returns: str

filename string

Examples

```
>>> exchange_filepath('/dir_a/subdir/sam/file.txt', '/data', 1)
'/data/sam/file.txt'
```

xrayutilities_utilities_noconf.exchange_path (orig, new, keep=0, replace=None)

function to exchange the root of a path with the option of keeping the inner directory structure. This for example includes such a conversion /dir_a/subdir/images/sample -> /home/user/data/images/sample where the two innermost directory names are kept (keep=2), or equally the three outer most are replaced (replace=3). One can either give keep, or replace, with replace taking preference if both are given. Note that replace=1 on Linux/Unix replaces only the root for absolute paths.

Parameters: orig: str

original path which should be replaced by the new path

new: str

new path which should be used instead

keep: int, optional

number of inner most directory names which should be kept the same in the output (default = 0)

replace: int, optional

number of outer most directory names which should be replaced in the output (default = None)

Returns: str

directory path string

Examples

```
>>> exchange_path('/dir_a/subdir/img/sam', '/home/user/data', keep=2)
'/home/user/data/img/sam'
```

```
xrayutilities.utilities_noconf.is_valid_variable_name (name)
```

xrayutilities.utilities_noconf.lam2en (inp)

converts the input wavelength in Angstrom to an energy in eV

Parameters: inp : float or str

wavelength in Angstrom

Returns: float

energy in eV

Examples

```
>>> energy = lam2en(1.5406)
```

xrayutilities.utilities_noconf.makeNaturalName (name, check=False)
xrayutilities.utilities_noconf.set_bit (f, offset)
sets the bit at an offset
xrayutilities.utilities_noconf.wavelength (wl)
convert common energy names to energies in eV
so far this works with CuKa1, CuKa2, CuKa12, CuKb, MoKa1

Parameters: wl : float, array-like or str

wavelength; If scalar or array the wavelength in Angstrom will be returned unchanged,

string with emission name is converted to wavelength

Returns: float or array-like

wavelength in Angstrom

Module contents

xrayutilities is a Python package for assisting with x-ray diffraction experiments. Its the python package included in xrayutilities.

It helps with planning experiments as well as analyzing the data.

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xrayutilities.mpl_helper

xrayutilities.normalize

xrayutilities.q2ang_fit

xrayutilities.simpack

xrayutilities.simpack.darwin_theory

xrayutilities.simpack.fit

xrayutilities.simpack.helpers

xrayutilities.simpack.models

xrayutilities.simpack.mosaicity

xrayutilities.simpack.powder

xrayutilities.simpack.powdermodel

xrayutilities.simpack.smaterials

xrayutilities.utilities

xrayutilities_utilities_noconf