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# **easyInterface**

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## EXAMPLE GALLERIES

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easyInterface is a library to interface crystallographic calculators to front end applications, JuPyter notebooks and scripting interfaces.

The code of the project is on Github: [easyInterface](#)



## FEATURES OF EASYINTERFACE

Boom!





## PROJECTS USING EASYINTERFACE

easyInterface is currently being used in the following projects:

- [easyDiffraction](#) - A scientific software for modelling and analysis of the neutron diffraction data



## INSTALLATION

### 3.1 Install via pip

You can do a direct install via pip by using:

```
$ pip install easyInterface
```

### 3.2 Install as an easyInterface developer

You can get the latest development source from our [Github repository](#). You need `setuptools` installed in your system to install `easyInterface`. For example, you can do:

```
$ git clone https://github.com/easyDiffraction/easyInterface
$ cd easyInterface
$ pip install -r requirements.txt
$ pip install -e .
```

### 3.3 Main Contents

#### 3.3.1 Introduction to easyInterface

Here we can see some examples of `easyInterface` in action

#### 3.3.2 Scripted Examples

This section gathers examples which don't produce any figures. These examples show the basic features of `easyInterface`.

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**Note:** Click [here](#) to download the full example code or to run this example in your browser via Binder

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#### Creating a interface

This demonstrates an example of how to load an example and create a interface to a `crispy` calculator. Information about the project is then displayed.

```
import os

from easyInterface.Diffraction.Calculators import CryspyCalculator
from easyInterface.Diffraction.Interface import CalculatorInterface

main_rcif = os.path.join('examples', 'Fe3O4_powder-1d_neutrons-pol_5C1(LLB)', 'main.
↪cif')
calculator = CryspyCalculator(main_rcif)

interface = CalculatorInterface(calculator)

print(interface.project_dict)

print(interface.phasesIds())

print(interface.getPhase(interface.phasesIds()[0]))
```

**Total running time of the script:** ( 0 minutes 0.000 seconds)

---

**Note:** Click [here](#) to download the full example code or to run this example in your browser via Binder

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### Creating a QT interface

This demonstrates an example of how to load an example and create a QT interface to a cryspy calculator. Information about the project is then displayed.

```
import os

from easyInterface.Diffraction.Calculators import CryspyCalculator
from easyInterface.Diffraction.QtInterface import QtCalculatorInterface

main_rcif = os.path.join('examples', 'Fe3O4_powder-1d_neutrons-pol_5C1(LLB)', 'main.
↪cif')
calculator = CryspyCalculator(main_rcif)

interface = QtCalculatorInterface(calculator, None)

print(interface.project_dict)

print(interface.phasesIds())

print(interface.getPhase(interface.phasesIds()[0]))
```

**Total running time of the script:** ( 0 minutes 0.000 seconds)

---

**Note:** Click [here](#) to download the full example code or to run this example in your browser via Binder

---

## Performing a fit

This demonstrates an example of how to load an example and create an interface to a crspsy calculator and then fit a value.

```
import os
from easyInterface.Diffraction.Calculators import CrspsyCalculator
from easyInterface.Diffraction.Interface import CalculatorInterface

main_rcif = os.path.join('examples', 'PbSO4_powder-1d_neutrons-unpol_D1A(ILL)', 'main.
↪cif')
calculator = CrspsyCalculator(main_rcif)

interface = CalculatorInterface(calculator)

print(interface.project_dict)

phase_ids = interface.phasesIds()

print(phase_ids)

phase = interface.getPhase(phase_ids[0])
phase['phasename'] = 'PbSO5'
interface.addPhase(phase)
interface.removePhase('PbSO5')

phase = interface.getPhase(phase_ids[0])
interface.setPhaseValue(phase_ids[0], ['atoms', 'Pb', 'fract_x'], 0.18)
interface.setPhases(phase)
print(phase)

interface.setPhaseRefine(phase_ids[0], ['atoms', 'Pb', 'fract_x'], True)

calc = interface.getCalculations()
print(calc)

res = interface.refine()

print(res)
```

Total running time of the script: ( 0 minutes 0.000 seconds)

## 3.3.3 Interface and Calculators

### easyInterface Interface

**class** easyInterface.Diffraction.Interface.CalculatorInterface (*calculator*)

Interface to calculators in the *easyInterface.Diffraction.Calculator* class.

**addExperimentDefinition** (*exp\_path*)

Add an experiment to be simulated from a cif file. Note that this will not have any crystallographic phases associated with it.

**Parameters** *exp\_path* (str) – Path to a experiment file (.cif)

**addPhase** (*phase*)

Add a new phases from a cif file to the list of existing crystal phases.

**Parameters** `phase` (*Phase*) –

**addPhaseDefinition** (*phase\_path*)

Add a new phases from a cif file to the list of existing crystal phases.

**Parameters** `phase_path` (*str*) – Path to a phase definition file (*.cif*)

Example:

```
interface = CalculatorInterface(calculator)
phase_path = '~/Experiments/new_phase.cif'
interface.addPhaseDefinition(phase_path)
```

**asCifDict** ()

...

**Return type** *dict*

**asDict** ()

Return data dict.

**Return type** *dict*

**experimentsCount** ()

Returns number of experiments in the project.

**Return type** *int*

**experimentsIds** ()

Returns labels of the experiments in the project.

**Return type** *list*

**property final\_chi\_square**

Calculates the final chi squared of the simulation. Where the final chi squared is the chi squared divided by the number of data points.

**Return type** *float*

**Returns** Final chi squared

**getPhase** (*phase\_name*)

Returns a phase from the project dictionary by name if one is supplied. If the phase name is none then all phases are returned. If the phase name does not exist **KeyError** is thrown.

**Parameters** `phase_name` (*Optional[str]*) – Name of the phase to be returned or None for all phases

**Return type** *Phase*

**Returns** Copy of the project dictionaries phase object with name `phase_name`

**Raises** **KeyError** – The supplied key is not a valid phase name

**phasesCount** ()

Returns number of phases in the project.

**Return type** *int*

**phasesIds** ()

Returns labels of the phases in the project.

**Return type** *list*

**refine** ()

refinement ...

**Return type** dict

**saveCifs** (*save\_dir*)

Write project cif files (*main.cif*, *experiments.cif* and *phases.cif*) to a user supplied directory. This contains all information needed to recreate the project dictionary.

**Parameters** **save\_dir** (str) – Directory to where the project cif files should be saved.

**Returns** None

**setExperiment** (*experiment*)

Set phases (sample model tab in GUI)

**setExperimentDefinition** (*exp\_path*)

Parse the relevant phases file and update the corresponding model

**setExperiments** (*experiments=None*)

Set experiments (Experimental data tab in GUI)

**setPhase** (*phase*)

Set phases (sample model tab in GUI)

**setPhaseDefinition** (*phase\_path*)

Parse a phases cif file and replace existing crystal phases

**Parameters** **phase\_path** (str) – Path to new phase definition file (*.cif*)

Example:

```
interface = CalculatorInterface(calculator)
phase_path = '~/Experiments/phases.cif'
interface.setPhaseDefinition(phase_path)
```

**setPhases** (*phases=None*)

Set phases (sample model tab in GUI)

**setProjectFromCalculator** ()

Sets the project dictionary from the calculator given on initialisation. Calling this function will regenerate the project dictionary and changes may be lost.

**updatePhases** ()

Synchronise the phases in project dictionary by queering the calculator object.

**Returns** None

**writeExpCif** (*save\_dir*)

Write the *experiments.cif* where all experiments in the project dictionary are saved to file. This includes the instrumental parameters and which phases are in the experiment/s

**Parameters** **save\_dir** (str) – Directory to where the experiment cif file should be saved.

**Returns** None

**writeMainCif** (*save\_dir*)

Write the *main.cif* where links to the experiments and phases are stored and other generalised project information.

**Parameters** **save\_dir** (str) – Directory to where the main cif file should be saved.

**Returns** None

**writePhaseCif** (*save\_dir*)

Write the *phases.cif* where all phases in the project dictionary are saved to file. This cif file should be compatible with other crystallographic software.

**Parameters** `save_dir` (str) – Directory to where the phases cif file should be saved.

**Returns** None

### easyInterfaces Project Dictionary

```
class easyInterface.Diffraction.Interface.ProjectDict (interface, app, calculator,  
                                                    info, phases, experiments,  
                                                    calculations)
```

This class deals with the creation and modification of the main project dictionary.

```
classmethod default ()
```

Create a default and empty project dictionary

**Return type** *ProjectDict*

**Returns** Default project dictionary with undo/redo functionality

```
classmethod fromPars (experiments, phases, calculations)
```

Create a main project dictionary from phases and experiments.

**Parameters**

- **experiments** (Union[*Experiments*, *Experiment*, List[*Experiment*]]) – A collection of experiments to be compared to calculations
- **phases** (Union[*Phases*, *Phase*, List[*Phase*]]) – A Collection of crystallographic phases to be calculated

**Return type** *ProjectDict*

**Returns** Project dictionary with undo/redo

### easyInterface Cryspy Calculator

```
class easyInterface.Diffraction.Calculators.CryspyCalculator (main_rcif_path=None)
```

```
asCifDict ()
```

...

**Return type** dict

```
getPhases ()
```

Set phases (sample model tab in GUI)

**Return type** *Phases*

```
refine ()
```

refinement ...

```
setExperiments (experiments)
```

Set experiments (Experimental data tab in GUI)

```
setObjFromProjectDicts (phases, experiments)
```

Set all the cryspy parameters from project dictionary

```
setPhases (phases)
```

Set phases (sample model tab in GUI)



### 3.3.4 Container Classes

#### Phase Classes

**class** easyInterface.Diffraction.DataClasses.PhaseObj.Atom.**ADP** (*u\_11*, *u\_22*,  
*u\_33*, *u\_12*, *u\_13*,  
*u\_23*)

Data store for Atom site anisotropic displacement parameters

**class** easyInterface.Diffraction.DataClasses.PhaseObj.Atom.**Atom** (*atom\_site\_label*,  
*type\_symbol*,  
*scat\_length\_neutron*,  
*fract\_x*, *fract\_y*,  
*fract\_z*, *oc-*  
*cupancy*,  
*adp\_type*,  
*U\_iso\_or\_equiv*,  
*ADp*, *MSp*)

Storage for details about an atom

**classmethod default** (*atom\_site\_label*)

Default constructor for an atom given a unique name in the phase

**Parameters** *atom\_site\_label* (*str*) – The atoms unique name in the phase

**Return type** *Atom*

**Returns** Default atom with a given name

**classmethod fromPars** (*atom\_site\_label*, *type\_symbol*, *scat\_length\_neutron*, *fract\_x*, *fract\_y*,  
*fract\_z*, *occupancy*, *adp\_type*, *U\_iso\_or\_equiv*, *ADp=None*, *MSp=None*)

Atom constructor from parameters

**Parameters**

- **atom\_site\_label** (*str*) – The unique name of the atom in the phase
- **type\_symbol** (*str*) – The type of atom
- **scat\_length\_neutron** (*float*) – Neutron scattering length
- **fract\_x** (*float*) – X position
- **fract\_y** (*float*) – Y position
- **fract\_z** (*float*) – Z position
- **occupancy** (*float*) – Site occupancy
- **adp\_type** (*str*) – ADP type code
- **U\_iso\_or\_equiv** (*float*) – Isotropic atomic displacement parameter

**Return type** *Atom*

**Returns** Fully formed atom data store

**classmethod fromXYZ** (*atom\_site\_label*, *type\_symbol*, *x*, *y*, *z*)

Construct an atom from name, type and position

**Parameters**

- **atom\_site\_label** (*str*) – The atoms unique name in the phase
- **type\_symbol** (*str*) – The type of atom

- **x** (float) – X position
- **y** (float) – Y position
- **z** (float) – Z position

**Return type** *Atom*

**Returns** Atom with name type and position filled in

**class** easyInterface.Diffraction.DataClasses.PhaseObj.Atom.**Atoms** (*atoms*)  
Container for multiple atoms

**class** easyInterface.Diffraction.DataClasses.PhaseObj.Atom.**MSP** (*MSPTYPE*, *chi\_11*,  
*chi\_22*, *chi\_33*,  
*chi\_12*, *chi\_13*,  
*chi\_23*)  
Data store for Atom site magnetic susceptibility parameters

**class** easyInterface.Diffraction.DataClasses.PhaseObj.Cell.**Cell** (*length\_a*,  
*length\_b*,  
*length\_c*, *angle\_alpha*,  
*angle\_beta*,  
*angle\_gamma*)  
Container for crystallographic unit cell parameters

**classmethod default** ()  
Default constructor for a crystallographic unit cell

**Return type** *Cell*

**Returns** Default crystallographic unit cell container

**classmethod fromPars** (*length\_a*, *length\_b*, *length\_c*, *angle\_alpha*, *angle\_beta*, *angle\_gamma*)  
Constructor of a crystallographic unit cell when parameters are known

**Parameters**

- **length\_a** (float) – Unit cell length a
- **length\_b** (float) – Unit cell length b
- **length\_c** (float) – Unit cell length c
- **angle\_alpha** (float) – Unit cell angle alpha
- **angle\_beta** (float) – Unit cell angle beta
- **angle\_gamma** (float) – Unit cell angle gamma

**Return type** *Cell*

**Returns**

**class** easyInterface.Diffraction.DataClasses.PhaseObj.Phase.**Phase** (*name*, *space-group*, *cell*,  
*atoms*, *sites*)  
Container for crystallographic phase information

**classmethod default** (*name*)  
Default constructor for a crystallographic phase with a given name

**Return type** *Phase*

**Returns** Default empty phase with a name

**class** easyInterface.Diffraction.DataClasses.PhaseObj.Phase.**Phases** (*phases*)  
 Container for multiple phases

**renamePhase** (*old\_phase\_name*, *new\_phase\_name*)  
 Easy method of renaming a phase

**Parameters**

- **old\_phase\_name** (*str*) – phase name to be changed
- **new\_phase\_name** (*str*) – new phase name

**Return type** NoReturn

**class** easyInterface.Diffraction.DataClasses.PhaseObj.SpaceGroup.**SpaceGroup** (*crystal\_system*,  
*space\_group\_name\_H*  
*space\_group\_IT\_num*  
*ori-*  
*gin\_choice*)

## Data Classes

**class** easyInterface.Diffraction.DataClasses.DataObj.Calculation.**BraggPeaks** (*bragg\_peaks*)  
 Container for multiple calculations

**class** easyInterface.Diffraction.DataClasses.DataObj.Calculation.**CalculatedPattern** (*x*,  
*y\_calc*,  
*y\_diff\_lower*  
*y\_diff\_upper*)

Storage container for a calculated pattern

**class** easyInterface.Diffraction.DataClasses.DataObj.Calculation.**Calculation** (*name*,  
*bragg\_peaks*,  
*cal-*  
*cu-*  
*lated\_pattern*,  
*lim-*  
*its*)

Storage container for calculations

**class** easyInterface.Diffraction.DataClasses.DataObj.Calculation.**Calculations** (*calculations*)  
 Container for multiple calculations

**class** easyInterface.Diffraction.DataClasses.DataObj.Calculation.**CrystalBraggPeaks** (*name*,  
*h*,  
*k*,  
*l*,  
*ttheta*)

Generator for HKL reflections and corresponding two theta.

**class** easyInterface.Diffraction.DataClasses.DataObj.Calculation.**Limits** (*y\_obs\_lower=-*  
*inf*,  
*y\_obs\_upper=inf*,  
*y\_diff\_upper=inf*,  
*y\_diff\_lower=-*  
*inf*,  
*x\_calc=None*,  
*y\_calc=None*)

Generator for limits of a dataset

```
class easyInterface.Diffraction.DataClasses.DataObj.Experiment.Background (ttheta,  
in-  
ten-  
sity)  
  
Data store for the background data parameters  
  
classmethod default ()  
    Default constructor for a background point  
  
    Return type Background  
  
    Returns Default background data object  
  
classmethod fromPars (ttheta, intensity)  
    Constructor for background when two theta and intensity are known  
  
    Parameters  
        • ttheta (float) – Two Theta angle in degrees  
        • intensity (float) – Value for intensity  
  
    Return type Background  
  
    Returns Background data dict  
  
class easyInterface.Diffraction.DataClasses.DataObj.Experiment.Backgrounds (backgrounds)  
    Store for a collection of background points  
  
class easyInterface.Diffraction.DataClasses.DataObj.Experiment.Experiment (name,  
wave-  
length,  
off-  
set,  
phase,  
back-  
ground,  
res-  
o-  
lu-  
tion,  
mea-  
sured_pattern)  
  
Experimental details data container  
  
classmethod default (name)  
    Default constructor for an Experiment  
  
    Parameters name (str) – What the experiment should be called  
  
    Return type Experiment  
  
    Returns Default empty experiment  
  
classmethod fromPars (name, wavelength, offset, scale, background, resolution, mea-  
sured_pattern)  
    Constructor of experiment from parameters  
  
    Parameters  
        • name (str) – What the experiment should be called  
        • wavelength (float) – Experimental wavelength
```

- **offset** (float) – Experimental offset
- **scale** (float) – Scale parameter
- **background** (*Backgrounds*) – Background model
- **resolution** (*Resolution*) – Resolution model
- **measured\_pattern** (*MeasuredPattern*) – The Measured data

**Return type** *Experiment*

**Returns** Experiment from parameters

**class** easyInterface.Diffraction.DataClasses.DataObj.Experiment.**ExperimentPhase** (*name*, *scale*)

Storage container for the Experimental Phase details

**classmethod default** (*name*)

Default experimental phase data container

**Return type** *ExperimentPhase*

**Returns** Default experimental phase data container

**classmethod fromPars** (*name*, *scale*)

Parameter initialised experimental phase data container

**Return type** *ExperimentPhase*

**Returns** Set experimental phase data container

**class** easyInterface.Diffraction.DataClasses.DataObj.Experiment.**ExperimentPhases** (*experiment\_phases*)

Storage of multiple phase markers associated with experiments

**class** easyInterface.Diffraction.DataClasses.DataObj.Experiment.**Experiments** (*experiments*)

Container for multiple experiments

**class** easyInterface.Diffraction.DataClasses.DataObj.Experiment.**MeasuredPattern** (*x*, *y\_obs*, *sy\_obs*, *y\_obs\_up=None*, *sy\_obs\_up=None*, *y\_obs\_down=None*, *sy\_obs\_down=None*)

Storage container for measured patterns

**classmethod default** (*polarised=False*)

Default constructor for measured data container.

**Parameters** **polarised** (bool) – Should the container be initialised as a polarised data container?

**Returns** Empty data container

**property isPolarised**

Is the measured data of a polarised type?

**Return type** bool

**Returns** True if it is from a polarised measurement, false otherwise

**property y\_obs\_lower**

Lower data confidence bound.

**Return type** list

**Returns** value of lower confidence bound

**property y\_obs\_upper**

Upper data confidence bound.

**Return type** list

**Returns** value of upper confidence bound

```
class easyInterface.Diffraction.DataClasses.DataObj.Experiment.Resolution(u,  
                                                                    v,  
                                                                    w,  
                                                                    x,  
                                                                    y)
```

Data store for the resolution parameters

**classmethod default**()

Default constructor for the resolution dict

**Return type** *Resolution*

**Returns** Default resolution dict

**classmethod fromPars**(u, v, w, x, y)

Constructor when resolution parameters are known

**Parameters**

- **u** (float) – resolution parameter u
- **v** (float) – resolution parameter v
- **w** (float) – resolution parameter w
- **x** (float) – resolution parameter x
- **y** (float) – resolution parameter y

**Return type** *Resolution*

**Returns** Resolution dictionary with values set

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