

#### From modelling to McStas and more

A short guide

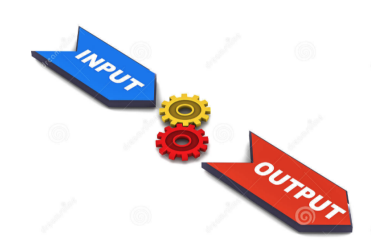
E. Farhi, ILL







# **Data formats**





#### McStas Data formats



McStas uses a number of input data formats, all text based.

- •Lau (Single\_crystal, PowderN, Isotropic\_Sqw)
- Laz (PowderN, Isotropic\_Sqw)
- •Sqw (Isotropic\_Sqw)
- •qSq (Isotropic\_Sqw) mostly for liquids
- •nxs (Sample\_nxs) this is *not* a NeXus/HDF
- •PDB (SANSPDB)

The question is how to generate these files...



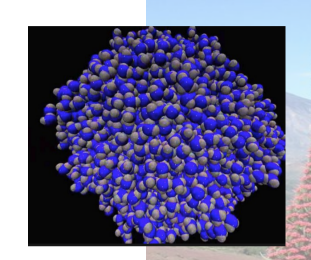


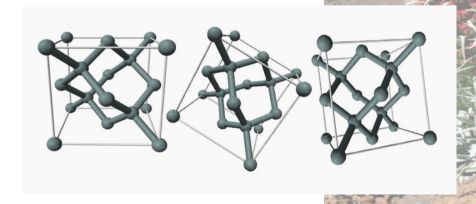


### **Material modelling formats**

- MD (classical or DFT)
  - trajectories, many data formats
- Lattice dynamics:
  - PhonoPy FORCE\_SETS

**–** ...



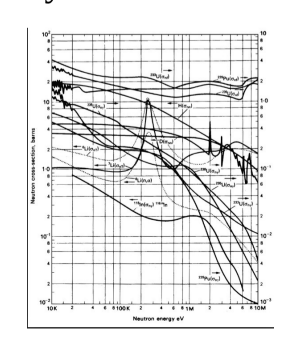






#### **Nuclear data base formats**

- ENDF, JEFF, JENDL, ...
  - all use a standard format for a set of materials. Most are monoatomic / incoherent
- ACE
  - Same as ENDF, but in binary for MCNP







# **Experiments**

- Intensity(d)  $\rightarrow$  F(Q)  $\rightarrow$  S(Q) and Laz
- Intensity(HKL)  $\rightarrow$  F2(HKL)  $\rightarrow$  Lau
- Intensity(theta, ToF)  $\rightarrow S(q, w)$
- Often saved as text and/or NeXus





## **Tools**

MDANSE2018

- iFit
- Mantid
- Python/ASE (IO)
- MDANSE/nMoldyn (converters)
- Scripts of all kinds
  - OpenBABEL, ...
- cif2hkl
- bash







### **Material DataBases**

- ICSD <a href="http://icsd.ill.fr/icsd/index.html">http://icsd.ill.fr/icsd/index.html</a>
- COD <a href="http://crystallography.net">http://crystallography.net</a>
- <a href="https://www.materialsproject.org/">https://www.materialsproject.org/</a>
- PhononDB <a href="http://phonondb.mtl.kyoto-u.ac.jp/">http://phonondb.mtl.kyoto-u.ac.jp/</a>
- Nomad <a href="https://repository.nomad-coe.eu/">https://repository.nomad-coe.eu/</a>

Mostly provide structures...



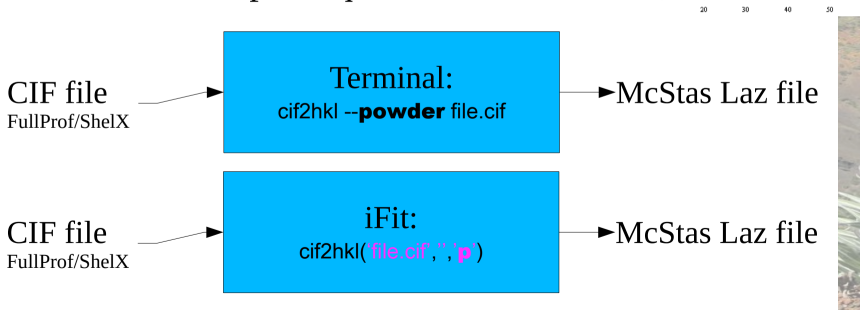


### McStas: structure: powder

• For powders, the pure elastic diffraction is easy to

prepare: LAZ

- PowderN
- Single\_crystal(powder=1, ...)
- Isotropic\_Sqw







#### McStas: structure: SX

• For single crystals, the pure elastic diffraction is

easy to prepare: LAU

- PowderN
- Single\_crystal
- Isotropic\_Sqw







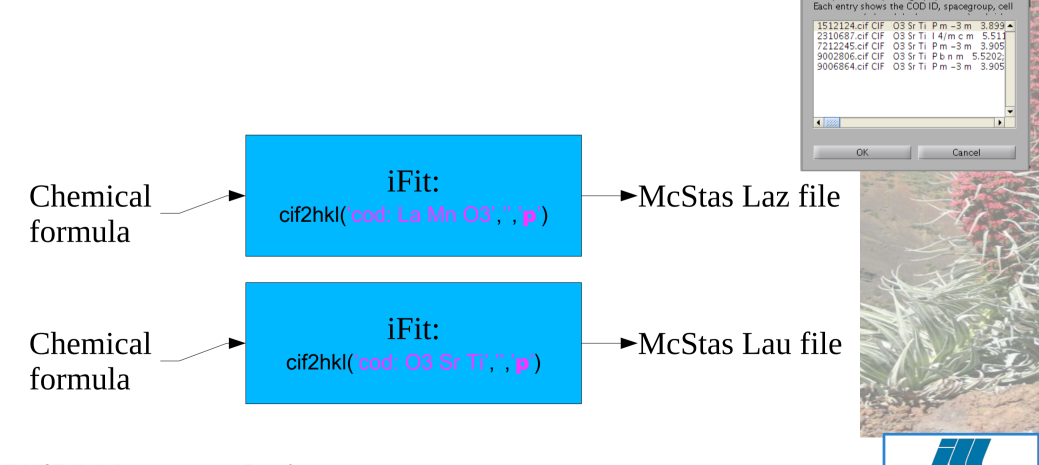
cif2hkl: Crystallog

Here are the entries for O3 Sr Ti

from the Crystallography Open Database (COD) <a href="http://www.crystallography.net">http://www.crystallography.net</a>.

#### McStas: structure: COD

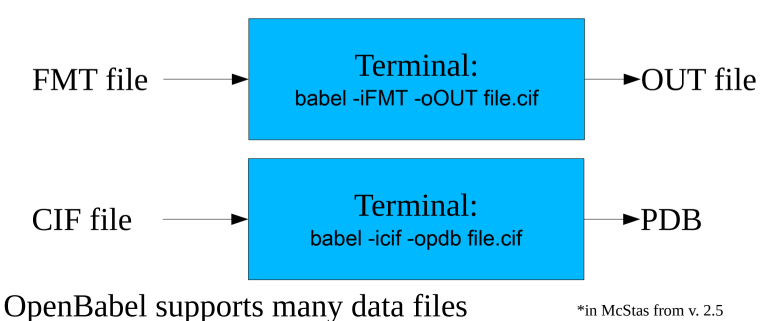
- Cif2hkl from iFit can input chemical formula
- Requires internet connection to COD





#### McStas: structure: others

- Other McStas structure file formats
  - SANSPDB and similars: get from Data Bases
  - Sample\_nxs: manual edit only (nxs, text)
  - Isotropic\_Sqw: manual edit only (qSq, text)
  - NCrystal\_sample\*: (ncmat,nxs,lau)



NEUTRONS FOR SOCIETY



### McStas: dyn from ToF spec

- Import and correct data with e.g. LAMP, Mantid
- Integrate radially (|q|) and compute S(q, W)
- Export to e.g. HDF/NeXus or other
- Import then export with iFit:
  - sqw=iData\_Sqw2D('file')
  - sqw.saveas('file.sqw','mcstas')
- Can/should always edit manually (text)

```
# weight   4.002 in [g/mol]
# density   0.4784 in [g/cm^3]
# sigma_abs   0.00747 absorption scattering cross section in [barn]
# sigma_coh   1.34   coherent scattering cross section in [barn]
# sigma_inc   0   incoherent scattering cross section in [barn]
# Temperature 2   in [K]
# classical   0   experimental, contains Bose factor
```



### McStas: dyn from exp. DOS

- Import and correct data with e.g. LAMP, Mantid
- Estimate g(w) with e.g. LAMP, Mantid, MUPHOCOR
- Export to e.g. HDF/NeXus or other
- Import then export with iFit:
  - g=iData\_vDOS('file')
  - inc=g.incoherent;
  - inc.saveas('inc.sqw', 'mcstas')
- Can/should always edit manually (text)





### McStas: dyn from nuclear DB

- Import and export with iFit:
  - sqw=iData\_Sqw2D('tsl.endf')
  - saveas(sqw(1), 'inc.sqw', 'mcstas')
- Can/should always edit manually (text)

```
# weight   4.002 in [g/mol]
# density   0.4784 in [g/cm^3]
# sigma_abs   0.00747 absorption scattering cross section in [barn]
# sigma_coh   1.34   coherent scattering cross section in [barn]
# sigma_inc   0    incoherent scattering cross section in [barn]
# Temperature 2   in [K]
# classical   0   experimental, contains Bose factor
```





### McStas: dyn from MD

- Import trajectory into MDANSE (converters)
- Double click the trajectory to allow analysis
- Use Plugins:
  - Analysis/Scattering/DCSF and DISF
  - Analysis/Scattering/Structure Factor
  - Analysis/Dynamics/Density of States
- Import then export with iFit:
  - sqw=iData\_Sqw2D('DCSF.nc')
  - sqw.saveas('coh.sqw','mcstas')
- Can/should always edit manually (text)





### McStas: dyn from nuclear DB

- Import and export with iFit:
  - sqw=iData\_Sqw2D('tsl.endf')
  - saveas(sqw(1), 'inc.sqw', 'mcstas')
- Can/should always edit manually (text)

```
# weight   4.002 in [g/mol]
# density   0.4784 in [g/cm^3]
# sigma_abs   0.00747 absorption scattering cross section in [barn]
# sigma_coh   1.34   coherent scattering cross section in [barn]
# sigma_inc   0    incoherent scattering cross section in [barn]
# Temperature 2   in [K]
# classical   0   experimental, contains Bose factor
```





#### **McStas: dyn from Lattice Dynamics**

- Using PhonoPy:
  - iFit: s4d=sqw\_phonons('phonopy\_directory')
- Computing on the fly:
  - iFit: s4d=sqw\_phonons('cod: La Mn O3')
  - p=powder(s4d)
  - d=iData\_Sqw2D(iData(p, [], 0:10, 0:100))
  - d.saveas('coh.sqw','mcstas')
- Other:
  - Import in iFit, convert to Sqw2D, export.





### **Neutron events**

- McStas handle 'rays' as packets of neutron events.
- Summing up builds intensity and distributions.
- McStas can read/write 'events' from other software:
  - MCNP TRAC and SSW
  - TRIPOLI
  - MCPL
- McStas can generate events as well for Mantid via NeXus/HDF files.



### McStas automation:



### bash

- Can use cif2hkl and mcrun commands:
  - cif2hkl 'file' --out 'file.laz' --powder
  - mcrun –dir=test instr Powder=file.laz
  - mcplot.pl -psc test

[~]\$ bash







### McStas automation: iFit

• iFit is a generic infrastructure which gathers data sets handling, fitting, S(q,w) models, and McStas hooks.

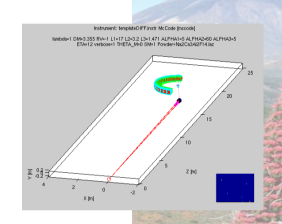






### McStas iFit: build

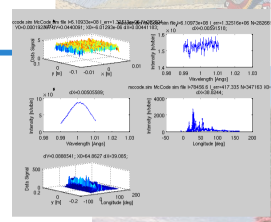
- McStas can be controlled from within iFit.
- Open Matlab/iFit
- Create the McStas model with:
  - model = mccode('instr')
- Plot the geometry with:
  - plot(model) % has contextual menus
- Edit the instrument and re-compile
  - edit(model)



#### MDANSE2018

### McStas iFit: eval

- Run with (default 1e6 event and pars)
  - data = iData(model, [], nan);
  - subplot(model) % plot monitors, contextual menus
- Specify parameters
  - data = iData(model, 'lambda=2.36; coh=Cu.laz')
- Do a scan:
  - data = iData(model, 'lambda= $[1.2 \ 2.4 \ 3.6]$ ')
- Change neutron events #
  - model.UserData.options.ncount = 1e7;





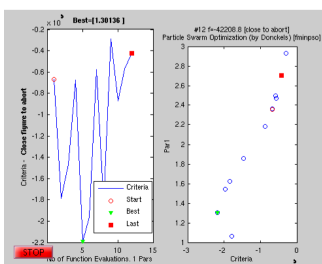


# McStas iFit: optim

- Fix all parameters but *lambda*, Maximize model value:
  - mlock(model, 'all'); munlock(model, 'lambda')
  - xlim(model, 'lambda', [1 3]); % bounds
  - fmax(model, 'lambda=2.36', '', nan)

fmax(model, 'lambda=2.36', 'OutputFcn=fminplot',

nan)







### McStas iFit: advanced

- You can add McStas models:
  - model=mccode('instr1')+mccode('instr2') + ...
- For instance:
  - Instr1: structure
  - Instr2: spin-wave
  - Instr3: phonons
  - Instr4: incoherent
- The different models can also be assembled as a set of samples chosen at execution in a single McStas instrument.

