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# From DFT to MLFT

*Use of FPLO and Quanty to perform  
ab-initio ligand field multiplets calculations*

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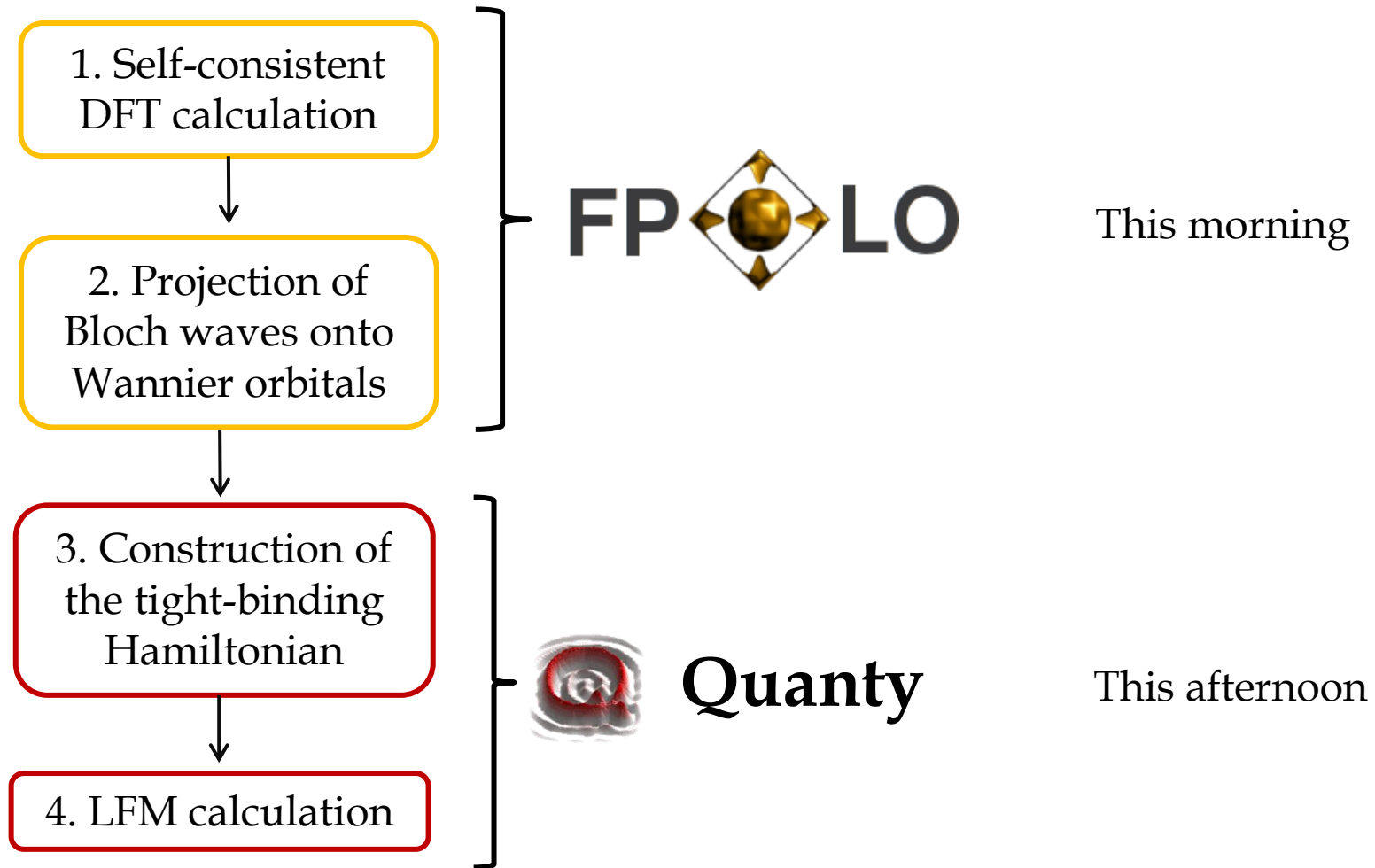


UNIVERSITÄT  
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Workshops in Heidelberg, Germany  
Hands on spectroscopy calculations of quantum materials  
September 27th 2018

# “*Ab-initio*” Ligand Field Multiplet: procedure



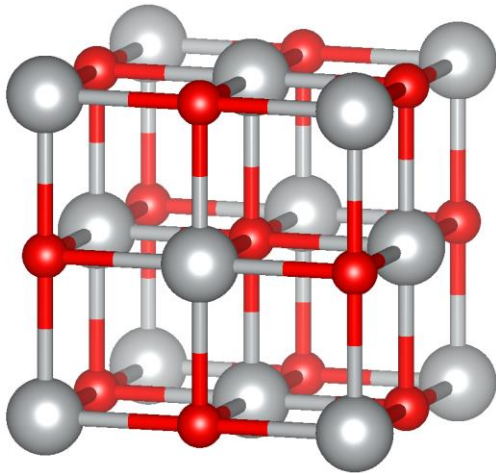
# Example I: NiO

NiO: a strongly correlated oxide

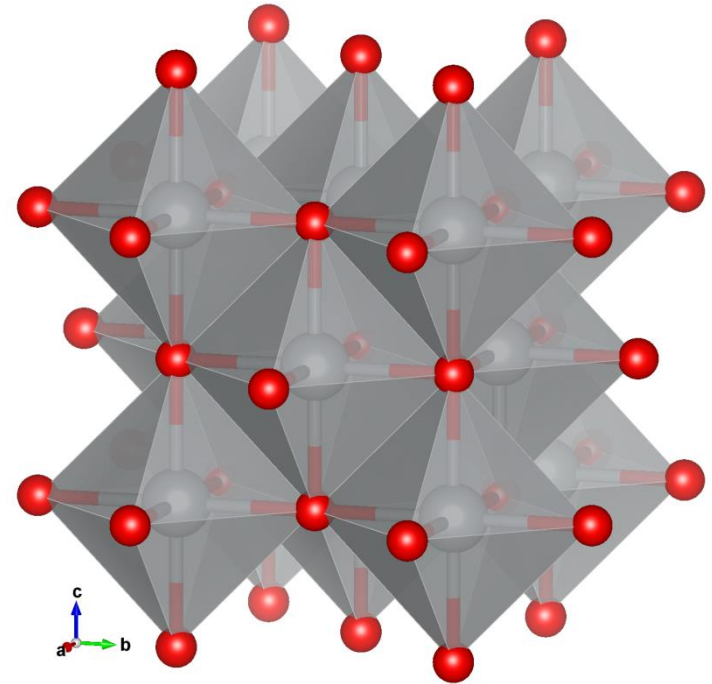
Rock salt structure

Space group  $Fm\bar{3}m$

$a=4.178 \text{ \AA}$

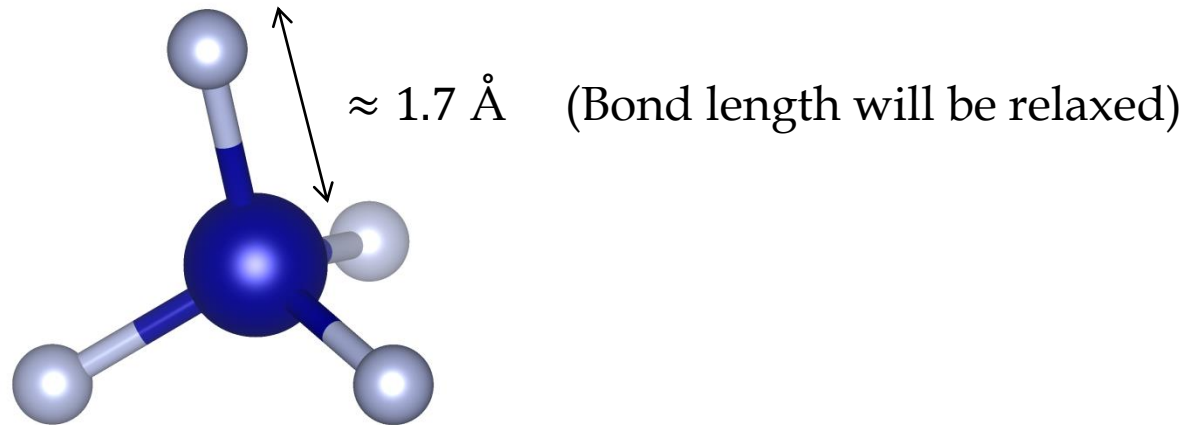


NiO<sub>6</sub> octahedra



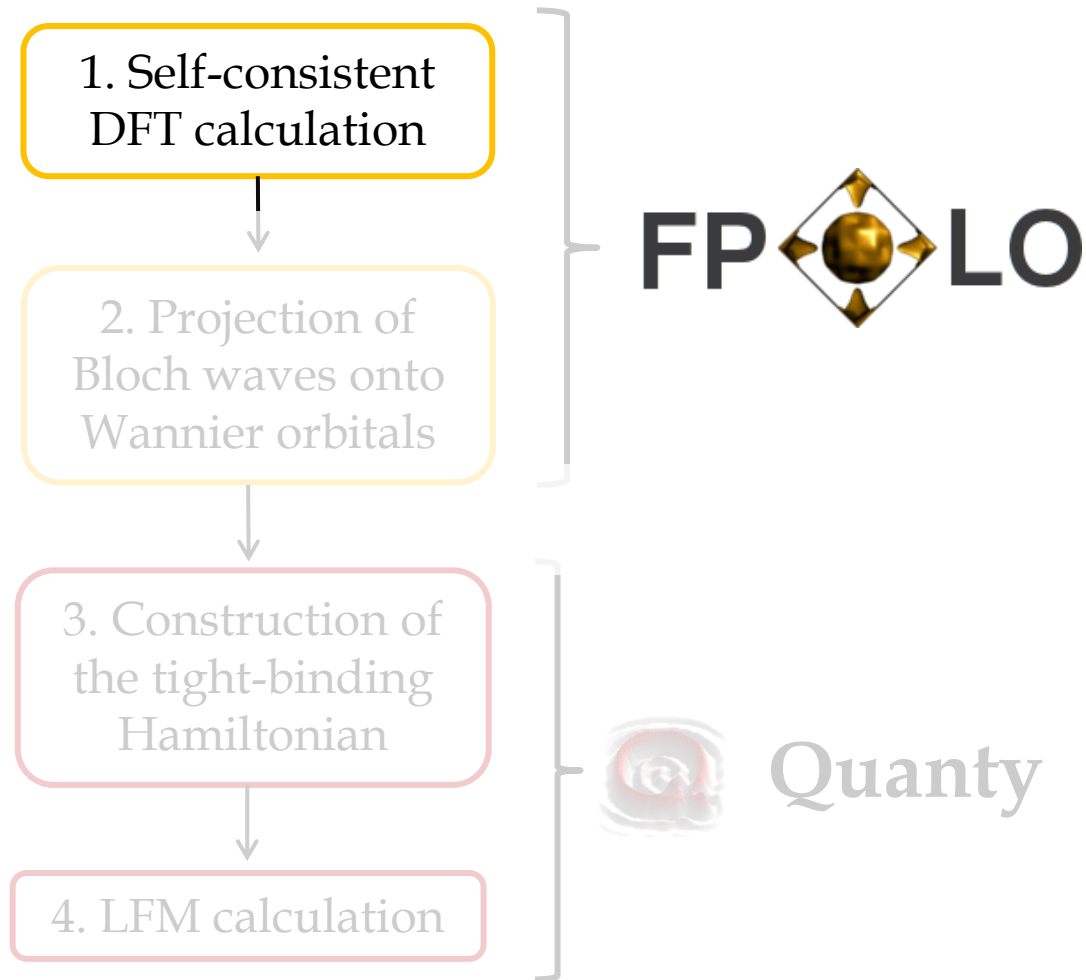
# Example II: molecule $\text{CrF}_4$

$\text{CrF}_4$ : Tetrahedral molecule



# Part 1: Density functional theory calculation : bands, DOS, Wannier functions

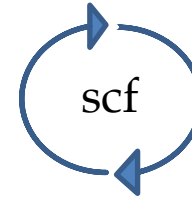
# “Ab-initio” Ligand Field Multiplet: procedure





## Full-potential local-orbital

- Kohn-Sham DFT - Self consistent



Basis: local orbitals at site **s** in the cell at **T**  
(non orthogonal)

$$u_{n,\mathbf{k}}(r) = \frac{1}{\sqrt{N}} \sum_{\mathbf{T}, \mathbf{s}, L} \phi_{\mathbf{s}, L}(\mathbf{r} - \mathbf{T} - \mathbf{s}) c_{n, \mathbf{k}}^{\mathbf{s}, L} e^{i\mathbf{k} \cdot (\mathbf{T} + \mathbf{s})}$$

- Solution of the **exact problem** for valence orbitals (frozen core electrons)

K. Koepnick and H. Eschrig, *Phys. Rev. B* **59**, 1743 (1999).

H. Eschrig, K. Koepnick, and I. Chaplygin, *J. Solid State Chemistry* **176**, 482 (2003).

# fedit

FPLO includes an input editor: `fedit`

```
$ cd TutorialDFTtoMLFT/NiO && mkdir FPLOcalc && cd FPLOcalc
```

```
$ fedit14.00-49-x86_64
```



Numbers vary with the version you have installed

```
fpl014.00-49-x86_64 OUTPUT
e(x)it
[...]
by K. Koepernik, A.Ernst and H.Eschrig (2003)
relativistic version by Ingo Opahle
LSDA+U by Igor Chaplygin

Any publication of results obtained by this program
has to include the citation:

K.Koepernik and H.Eschrig, Phys. Rev. B 59, 1743 (1999)

Any publication of CPA results obtained by this program
additionally has to include the citation:

K. Koepernik, B. Velicky, R. Hayn and H. Eschrig,
Phys. Rev. B 55, 5717 (1997)

main version: 14.00
sub version: M-CPA
release      : 49

compiled with ifort 1400
compiled with external LAPACK libraries

date       : Fri Sep 14 10:25:37 2018
host      : lin64n

File =.sym does not exist, will create it!
File =.sym created!
File =.in does not exist, will create it!
TERMINATION: Normal : File =.in created!
-----STDERR:

STATUS: OK (14.00-49:M-CPA)
```

Type

X



# fedit

## Main menu

```
MAIN MENU
[ ] (Q)uit/save (+) Symmetry (H)elp
-----
GENERAL DATA
(S)pin sorts : 1 (I)nitial polarization : [ ]
(K)-mesh subdivision : 12 12 12 (O)ccupied bands : -1
(N)umber of iterat. : 30 (A)ccuracy of density : 1.e-6
(T)otal energy calc. : [X] A(C)curacy of Etot : 1.e-8
Conver(G)ence condit : Density (-) Options : ...
(R)elativistic : scalar relativistic
(V)xc-version : Perdew Wang 92 (LSDA)
(F)inite nucleus : Point charge
xc-field str(E)ngth : 1.0
(W) fixed spin mom. : [ ] (Y) spin moment : 1.0
-----
RELATIVISTIC SETTINGS
Q(U)antization-axis : 0 0 1
-----
OTHERS
(>) verbosity level : more information
-----
STATUS: OK (14.00-49:M-CPA)
```

Type

+

to enter symmetry menu

Red keys: select another menu or exit

H opens a help screen

Blue keys: select an entry

# fedit

## Symmetry

```
SYMMEYRY MENU
[ ] e(X)it (+) Update (H)elp

(C)ompound      : NiO
s(T)tructure type : Crystal
(S)pacegroup    : FM3M (225)
(U)nit of length : Angstroem
(L)attice constants : 4.178 4.178 4.178
(A)xis angles   : 90. 90. 90.

Subgroup (G)enerators :

(N)umber of atoms : 2

Wyckoff positions

(1) -th atom sort : Ni      0. 0. 0.
(2) -th atom sort : 0       0.5 0.5 0.5
```

Fill in the lattice structure and chemical composition

Type



to update



Do not forget !!!

then type



to come back to the main menu

# 1<sup>st</sup> scf calculation

Main menu

```
MAIN MENU
[ ] (Q)uit/save (+) Symmetry (H)elp

GENERAL DATA
(S)pin sorts : 1 (I)nitial polarization : [ ]
(K)-mesh subdivision : 10 10 10 (O)ccupied bands : -1
(N)umber of iterat. : 30 (A)ccuracy of density : 1.e-6
(T)otal energy calc. : [X] A(C)curacy of Etot : 1.e-8
Conver(G)ence condit : Density (-) Options : ...
(R)elativistic : scalar relativistic
(V)xc-version : Perdew Wang 92 (LSDA)
(F)inite nucleus : Point charge
xc-field str(E)ngth : 1.0
(W) fixed spin mom. : [ ] (Y) spin moment : 1.0


RELATIVISTIC SETTINGS
Q(U)antization-axis : 0 0 1

OTHERS
(>) verbosity level : more information

STATUS: OK ( 14.00-49:M-CPA )

STATUS: OK ( 14.00-49:M-CPA )
```

Set the number of k-points to 10x10x10

Type   to quit fedit

Run FPLO:

\$ `fplo14.00-49-x86_64 | tee out.scflow`



fplo automatically uses the files created by fedit

# 1<sup>st</sup> scf calculation

Output (out.scflow) : Iteration progress, Population Analysis, Energy

= .dens: electron density; if present in the folder it is used as input by FPLO

Iteration progress:

```
$ grep 'last deviation' out.scflow
```

```
SCF: iteration 0 dimension 0 last deviation u= 0.00E+00
SCF: iteration 1 dimension 1 last deviation u= 0.24E+00
SCF: iteration 2 dimension 2 last deviation u= 0.23E+00
SCF: iteration 3 dimension 3 last deviation u= 0.81E-01
SCF: iteration 4 dimension 1 last deviation u= 0.28E-01
SCF: iteration 5 dimension 2 last deviation u= 0.55E-02
SCF: iteration 6 dimension 3 last deviation u= 0.29E-02
SCF: iteration 7 dimension 1 last deviation u= 0.13E-02
SCF: iteration 8 dimension 2 last deviation u= 0.18E-03
SCF: iteration 9 dimension 3 last deviation u= 0.73E-04
SCF: iteration 10 dimension 1 last deviation u= 0.47E-04
SCF: iteration 11 dimension 2 last deviation u= 0.49E-05
SCF: iteration 12 dimension 1 last deviation u= 0.26E-05
SCF: iteration 13 dimension 1 last deviation u= 0.86E-06
CONVERGED
```

# Display structure

FPLO includes a graphical interface: XFLO

```
$ xfplo =.in
```

The screenshot displays the XFLO 14.00 graphical interface. The main window, titled 'XFLO 14.00: Untitled', shows a 3D model of a crystal structure with green spheres (Ni) and red spheres (O) arranged in a cubic lattice. A pink sphere is also visible. The interface includes a menu bar (Activity, File, Input, Plot, Tools, Run, Settings, Window, Help) and a toolbar with icons for file operations and calculations.

A secondary window, titled 'XFLO 14.00: Measure distances', is open on the right. It displays a table of atomic positions and distances. The table has columns: No, Elm, Site, Wyck, and Pos. The data is as follows:

No	Elm	Site	Wyck	Pos
13	Ni	1	1	( 7.89527575022322 , 7.89527575022322 , 7.89527575022322
27	O	2	2	( 3.94763787511161 , 7.89527575022322 , 7.89527575022322
dif				( -3.94763787511161 , 0 , 0
dist 1st 2nd				= 3.94764
				= 2.089

Below the table, there is a 'Clear' button and a 'Save' button. A text box contains the following information:

```
site 2
difvec -3.947637875111609 -0.0000000000000002 -0.0000000000000002
```

At the bottom of the window, a message states: 'Click any sequence of atoms ... you will find out, how it works. 'ESC' for reset. 'M' for aborting.'

# 2<sup>nd</sup> scf calculation

```
$ fedit14.00-49-x86_64
```

Main menu

```
MAIN MENU
[ ] (Q)uit/save (+) Symmetry (H)elp

-----
GENERAL DATA
(S)pin sorts : 1 (I)nitia polarization : [ ]
(K)-mesh subdivision : 40 40 40 (O)ccupied bands : -1
(N)umber of iterat. : 30 (A)ccuracy of density : 1.e-9
(T)otal energy calc. : [X] A(C)curacy of Etot : 1.e-8
Conver(G)ence condit : Density
(-) Options : ...
(R)elativistic : scalar relativistic
(V)xc-version : Perdew Wang 92 (LSDA)
(F)inite nucleus : Point charge
xc-field str(E)ngth : 1.0
(W) fixed spin mom. : [ ] (Y) spin moment : 1.0

-----
RELATIVISTIC SETTINGS
Q(U)antization-axis : 0 0 1

-----
OTHERS
(>) verbosity level : more information

-----
STATUS: OK (14.00-49:M-CPA)
```

Increase the number of k-points to 40x40x40  
and decrease the accuracy of density to  $10^{-9}$

Type  to enter the Options menu

# 2<sup>nd</sup> scf calculation

## Options

```
e(X)it                                     OPTIONS                                     (H)elp
-----
Options
(0) CALC_DOS                               : [ ] (1) NOT_USED                               : [ ]
(2) FULLBZ                                 : [ ] (3) CALC_PLASMON_FREQ                         : [ ]
(4) EMPTY_LATTICE_TEST                     : [ ] (5) NO_DOS                                       : [ ]
(6) PLOT_REALFUNC                         : [ ] (7) PLOT_BASIS                                  : [X]
(8) TEST_LOI                              : [ ] (9) TEST_DIAGNO                                : [ ]
(A) TEST_SYMMETRIZATION                   : [ ] (B) TEST_HS_SYM                                : [ ]
(C) PROT_PRINT_BASIS                      : [ ] (D) PROT_MAKELATTICE                           : [ ]
(E) PROT_STRUCTURE_PRNT                   : [ ] (F) PROT_TCI                                    : [ ]
(G) NOT_USED                              : [ ] (I) NOT_USED                                    : [ ]
(J) NOT_USED                              : [ ] (K) NO_SYMMETRYTEST                            : [X]
(L) NO_POTENTIAL                          : [ ] (M) NOT_USED                                    : [ ]
(N) NO_CORE                               : [ ] (O) NOT_USED                                    : [ ]
(P) NO_POPANALYSIS                        : [ ] (Q) NO_LOI                                      : [ ]
(R) NO_BASIS                              : [ ] (S) NO_EFCH                                      : [ ]
(T) NOT_USED                              : [ ] (U) NOT_USED                                    : [ ]
(V) NOT_USED                              : [ ]

STATUS: OK                                     ( 14.00-49:M-CPA)
```

Type **7** to save the basis orbitals to disk

Type **x** **Q** **y** to quit

# 2<sup>nd</sup> scf calculation

Run FPLO:

```
$ fplo14.00-49-x86_64 | tee out.scf
```

```
+fcor.sort.spin: radial part of the core orbitals  
+fval.sort.spin: radial part of the valence orbitals  
+fkor.sort.spin: kinetic core functions  
+fkval.sort.spin: kinetic valence functions
```



# Band structure and DOS calculation



Do it after a converged scf (will use =.dens)

```
$ fedit14.00-49-x86_64
```

Type **space** **B** to enter Bandplot menu

Type **B** to compute bands

Type **W** to compute weights

Type **X** **Q** **Y** to quit

```
$ fplot14.00-49-x86_64 | tee out.band
```

```
BANDPLOT
e(X)it
(B)andstructure plot      : [X]
(R)ead sym-points        : [X]
(S)teps between sym-points : 50

DOS/AKBL/BANDS
Number of e(-)pts (non-CPA) : 1000
(P)lot IDOS      : [ ]
plot n(E)t DOS  : [ ]

(L)ower energy bound [eV] relative to E_f : -20.0
(U)pper energy bound [eV] relative to E_f : 20.0

Restr(I)ct bands to window : [ ]

Local (D)OS sites      :
Output +(C)oeff file : [ ]

BAND WEIGHTS/FAT BANDS
Weights def (F)ile      :
(W)eights                : [X]
(T)ransform quant. axis  : [ ]
X-(A)xis                 : 1.0 0.0 0.0
(Z)-axis                 : 0.0 0.0 1.0

(N)umber of sym-points   : 9

No.   Label      k-point
-----
(1)   : $~G      : 0 0 0
(2)   : X         : 1 0 0
(3)   : W         : 1 1/2 0
(4)   : K         : 3/4 3/4 0
(5)   : $~G      : 0 0 0
(6)   : L         : 1/2 1/2 1/2
(7)   : W         : 1 1/2 0
(8)   : U         : 1 1/4 1/4
(9)   : X         : 1 0 0

STATUS: OK (14.
```

# Band structure and DOS calculation

+band/+bweights Band structure and band weights

+dos.sort\* Projected DOS

```
$ head -n 1 +dos.sort001.n100[1-7]
```

```
==> +dos.sort001.n1001 <==  
# sort =      1  nl = 3s spin =      1
```

```
==> +dos.sort001.n1002 <==  
# sort =      1  nl = 3p spin =      1
```

```
==> +dos.sort001.n1003 <==  
# sort =      1  nl = 4s spin =      1
```

```
==> +dos.sort001.n1004 <==  
# sort =      1  nl = 5s spin =      1
```

```
==> +dos.sort001.n1005 <==  
# sort =      1  nl = 3d spin =      1
```

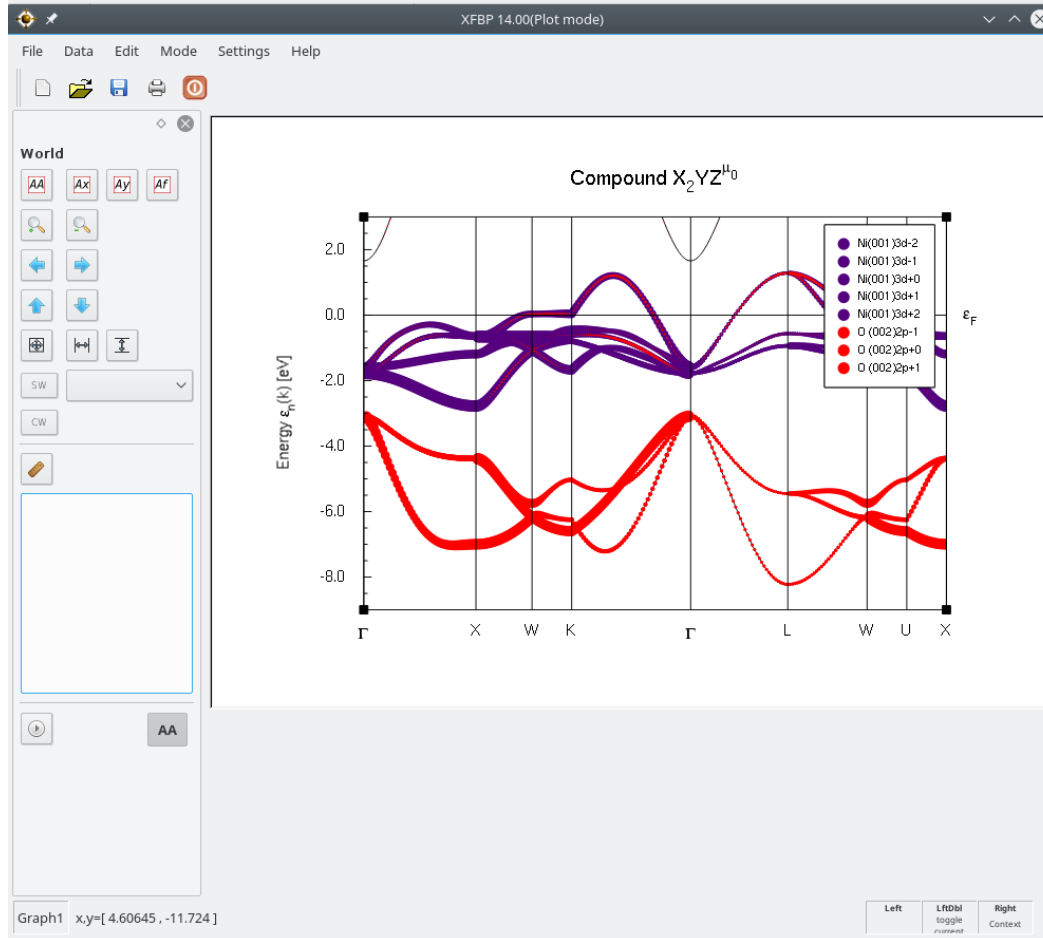
```
==> +dos.sort001.n1006 <==  
# sort =      1  nl = 4d spin =      1
```

```
==> +dos.sort001.n1007 <==  
# sort =      1  nl = 4p spin =      1
```

# Band structure and DOS plot

FPLO includes a plotting program: XFBP

\$ xfbp +bweights



● Ni 3d

● O 2p

(weights added by double clicking on the graph)

Non magnetic DFT calculations → NiO described as conductor

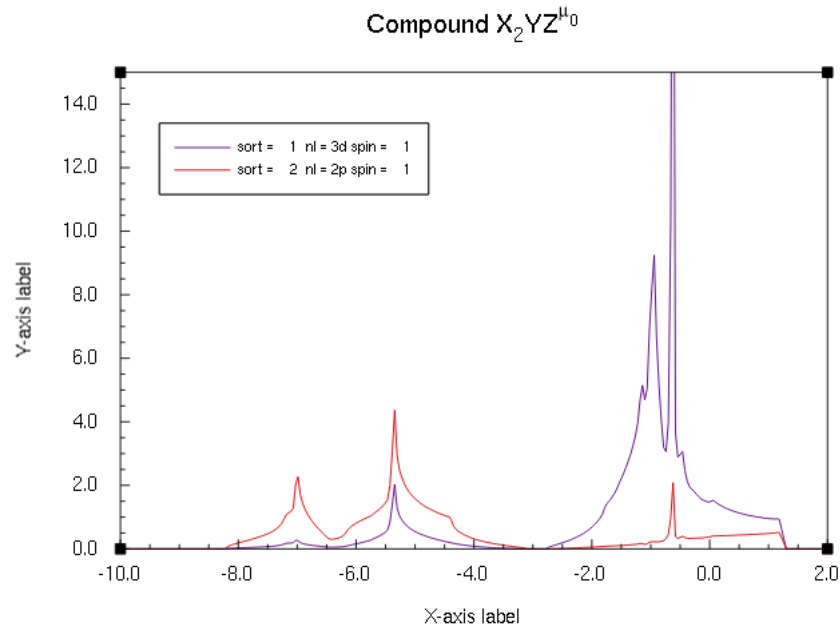
Failing of DFT to describe highly correlated materials

# Band structure and DOS plot

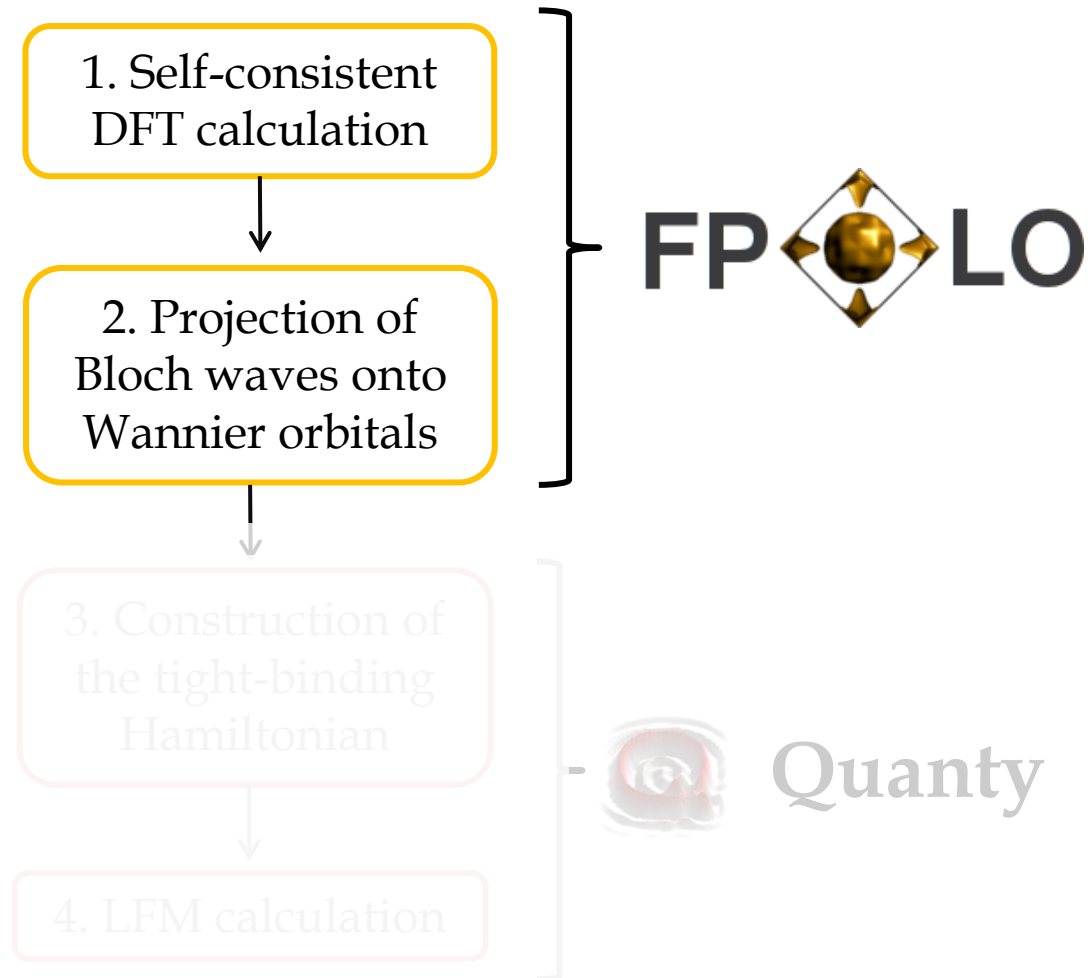
FPLO includes a plotting program: XFBP

```
$ xfbp +dos.sort001.n1005 +dos.sort002.n1004
```

DOS (states/eV) *vs* Energy (eV)



# “Ab-initio” Ligand Field Multiplet: procedure

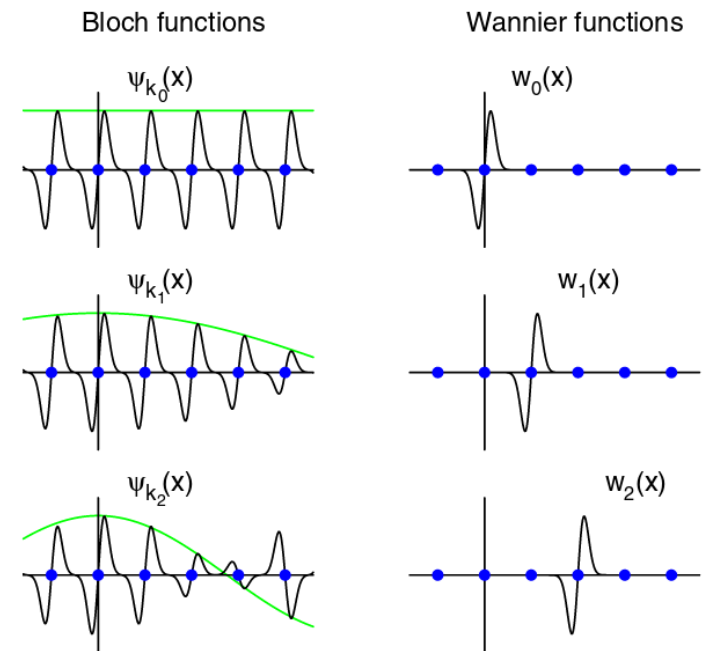


# Wannier orbitals

Wannier functions:

- set of orthonormal localized functions
- span the same space as the Bloch functions

$$W_{\mu}(\mathbf{r} - \mathbf{T}) = \frac{V}{(2\pi)^3} \int_{BZ} d\mathbf{k} e^{-i\mathbf{k} \cdot \mathbf{T}} \sum_n \underbrace{\psi_n^{\mathbf{k}}(\mathbf{r})}_{\text{Kohn-Sham functions}} \underbrace{U_{n\mu}^{\mathbf{k}}}_{U^{\mathbf{k}}: \text{Unitary matrix}}$$



Marzari et al. Rev. Mod. Phys., Vol. 84, No. 4

Projections onto Wannier functions can be interfaced with any band structure method

FPLO allows to obtain highly localized atom-like Wannier functions  
( $\neq$  maximally localized)

# Wannier functions with FPLO

The Wannier module of FPLO is activated if:

- A file `=.wandef` is present in the folder
- The keyword `doit` is found in the file

```
$ cp ../=.wandef
```

```
$ fplo14.00-49-x86_64 | tee out.wandef
```

Compute the projection matrix (still in  $k$  space)

```
$ fplo14.00-49-x86_64 | tee out.wan
```

Compute Wannier functions

# Wannier functions with FPLO

File = .wandef

```
doit
```

```
---- real space grid for pictures of WFs -----
```

```
WF_grid_basis conv
```

```
WF_grid_directions
```

```
2 0 0
```

```
0 2 0
```

```
0 0 2
```

} Output of the WFs on the real space  
grid (visualization)

```
WF_grid_subdivision 30 30 30
```

} Real space grid size. You can set it to 1 1 1  
if you do not need to plot the orbitals  
(save time and memory)

```
-----  
ham_cutoff 18
```

```
WF_ham_threshold 0
```

```
WF_coeff_threshold 0
```

```
WF_write_coeff_stats on
```

```
ham_write_t_stats on
```

In the output only hopping elements  $|t| > \text{WF\_ham\_threshold}$  are  
written, for atoms distances less than `ham_cutoff`.

Only contributions of the FPLO orbitals to the WFs, which are larger than  
`WF_coeff_threshold` are written in `+WF_coefficients`

```
----- ham export grid -----
```

```
k_grid_basis prim
```

```
k_grid_directions
```

```
1 0 0
```

```
0 1 0
```

```
0 0 1
```

```
k_grid_subdivision 1 1 1
```

```
k_grid_incl_periodic_points off
```

} Output of the WFs on the reciprocal  
space grid



# Wannier functions with FPLO

File = .wandef

----- NiO -----

```
wandef
on
  name Ni 3dxy
  emin -9
  emax +1.5
  de 0.0
  contrib
    site 1
    difvec 0 0 0
    xaxis 1 0 0
    zaxis 0 0 1
    orb 3d-2
    fac 1
```

One WF  
projector

Energy window: 1 from emin to emax, falling like a gaussian of width de

Orbital sitting at site 1

difvec must be non zero if several contributions in the Wannier function

xaxis zaxis: local coordinate system

Orbital 3d-2 →  
fac is the relative weight of each contrib

-2	-1	0	1	2
xy	yz	z <sup>2</sup>	xz	x <sup>2</sup> -y <sup>2</sup>

```
wandef
on
  name Ni 3dyz
  emin -9
  emax +1.5
  de 0.0
  contrib
    site 1
    difvec 0 0 0
    xaxis 1 0 0
    zaxis 0 0 1
    orb 3d-1
    fac 1
```

...



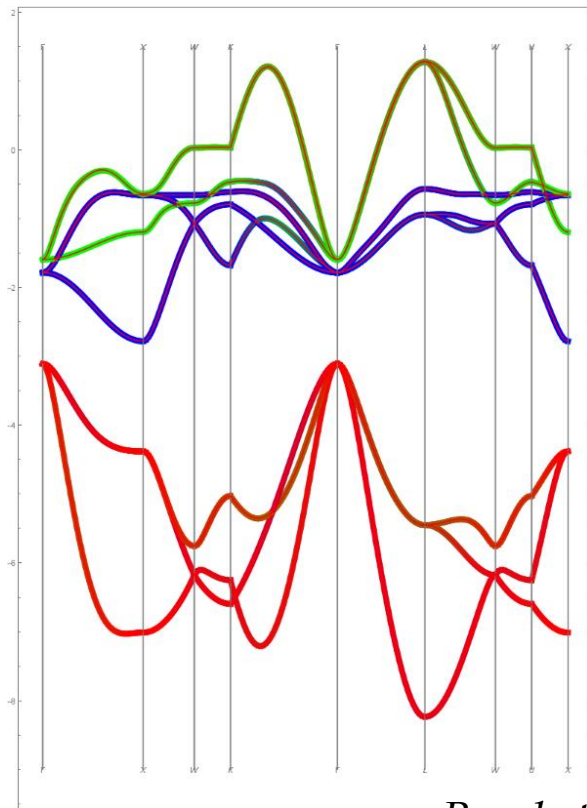
There can be more sites than Wyckoff positions  
(check scf output)

Define Wannier orbitals for each of them

# Check band structure and plot orbitals



If you have mathematica open file **NiO\_wannier.nb** in folder **NiO**



Band structure calculated

- (i) with the full basis set (red lines) and
- (ii) with the Wannier-functions basis set.

Green: Ni 3d  $e_g$

Blue: Ni 3d  $t_{2g}$

Red: O 2p

→ superimposed

*Band structure plot will be implemented in Quanta in a near future*

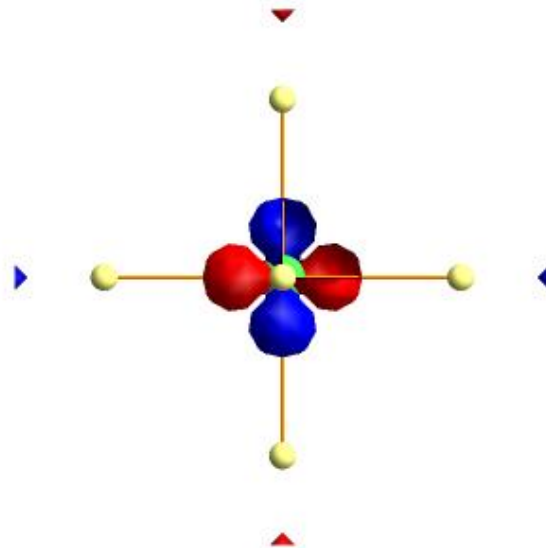
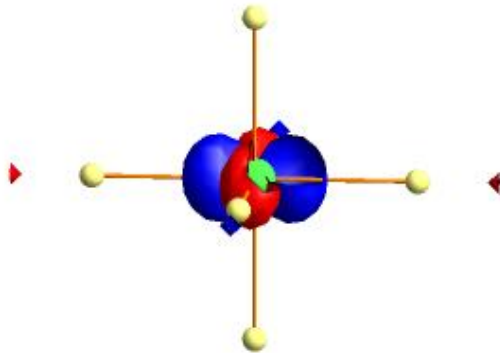
# Check band structure and plot orbitals



If you have mathematica open file **NiO\_wannier.nb** in folder **NiO**

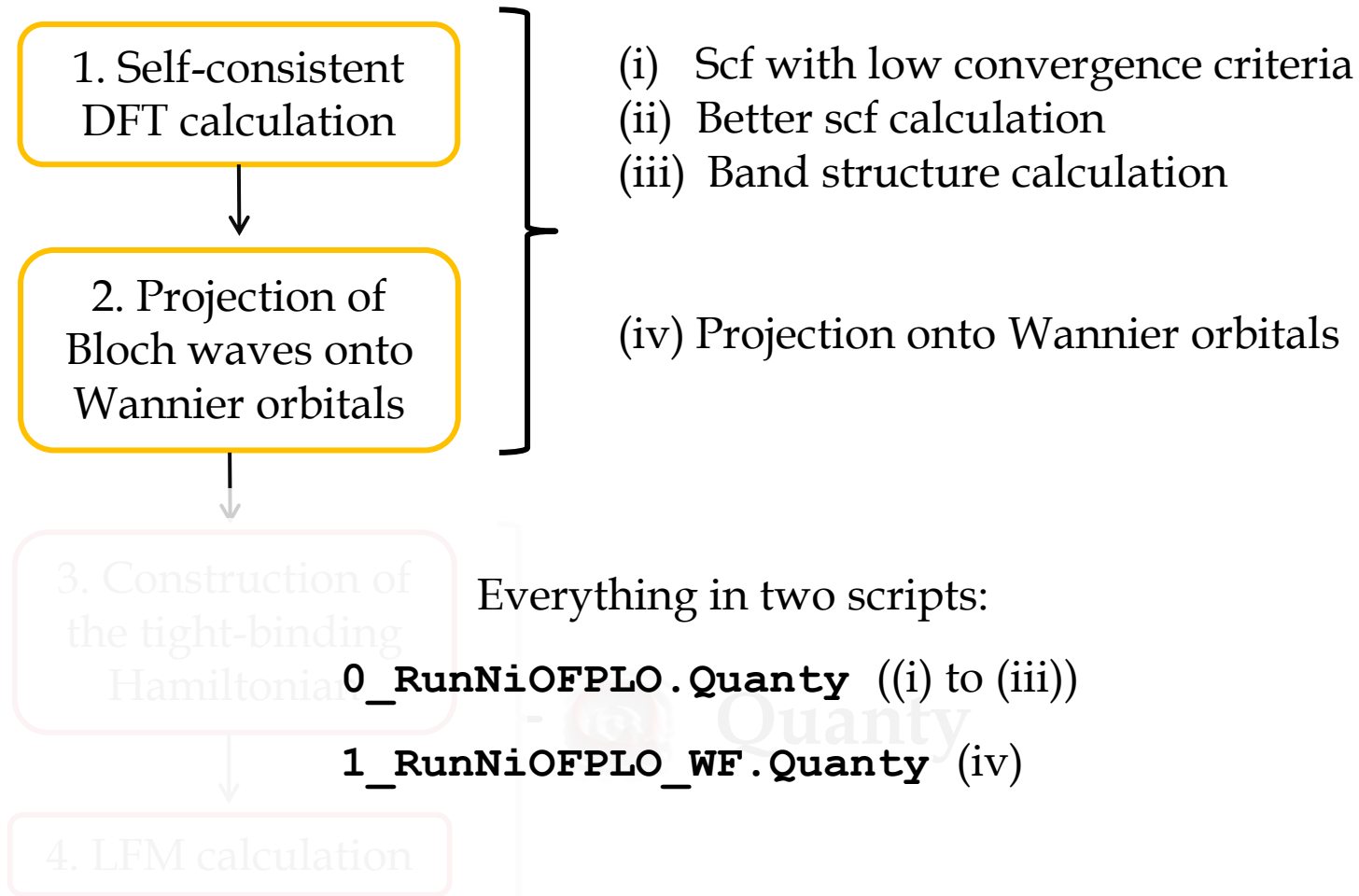
Wannier orbitals:

e.g. Ni  $3d$   $x^2-y^2$



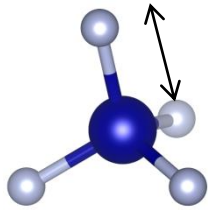
# Script method

\$ cd Scripts



Scripts could be written in any language  
(bash, python...)

# Example II: molecule $\text{CrF}_4$



Unknown bond length  $\rightarrow$  needs to be optimized

Procedure:

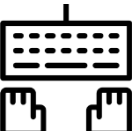
- (i) Optimization of atomic positions
- (ii) Well converged scf calculation
- (iii) Band structure calculation
- (iv) Projection onto Wannier orbitals

Use the script for NiO as a starting point:

```
$ cp NiO/Scripts/0_RunNiOFPLO.Quanty TutorialCrF4
$ cd TutorialCrF4
$ mv 0_RunNiOFPLO.Quanty 0_RunCrF4FPLO.Quanty } Rename file
```

# Example II: molecule $\text{CrF}_4$

- Modify file 0\_RunCrF4FPLO.Quanty



```
# title
@c@CrF4
# enter spacegroup select box - 215
@s@
@215@
@x@
# structure type - molecule
@t@
@m@
@x@
# lattice constants;
@l@ 4.178 4.178 4.178
# set axis angles
@a@ 90. 90. 90.
# setup Wyckoff positions
@n@2
# Now, give list of all Wyckoff positions.
@1@ Cr @ 0. 0. 0.
@2@ F @ 1. 1. 1.
```

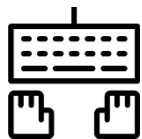
Not needed for a molecule

# Example II: molecule CrF<sub>4</sub>

- Modify file 0\_RunCrF4FPLO.Quanty



In the first FEditMenuOptions (below the structure)



```
# Set maximum number of iterations to 200
@n@200
# Set convergence criterion to Density OR energy
@g@
@1@
@x@
# activate structure optimization
# (The space before the 'f' opens the alternative menu
bar.)
```

```
@ f@
@f@
@2@
@x@
# leave forces menu
@x@
```

Lines to be added for  
structural optimization

# Example II: molecule $\text{CrF}_4$

- Modify file `0_RunCrF4FPLO.Quantity`



In the second `FEditMenuOptions`



```
# deactivate site geometry optimization again
# (The space before the 'f' opens the alternative menu
bar.)
@ f@
@f@
@1@
@x@
# leave forces menu
@x@
```

- \$ `Quantity 0_RunCrF4FPLO.Quantity`



# Example II: molecule $\text{CrF}_4$

```
$ cd DFT
```

```
$ ls
```

```
+band      +dos.sort002      +dos.total.l003  +fcor.002.4      +fkcor.002.4      +fval.001.2      +log      +points
=.dens      +dos.sort002.nl001 +dos.total.l004  +fedit           +fkval.001.1      +fval.001.3      out.band      +run
+dos.sort001 +dos.sort002.nl002 +dos.total.l005  +fedithelp       +fkval.001.2      +fval.001.4      out.fedit     =.sym.template
+dos.sort001.nl001 +dos.sort002.nl003 +fcor.001.1      +fkcor.001.1      +fkval.001.3      +fval.002.1      out.feditband =.sym
+dos.sort001.nl002 +dos.sort002.nl004 +fcor.001.2      +fkcor.001.2      +fkval.001.4      +fval.002.2      out.feditkmesh =.sym.bak
+dos.sort001.nl003 +dos.sort002.nl005 +fcor.001.3      +fkcor.001.3      +fkval.002.1      +fval.002.3      out.scf        =.sym.bak.forces
+dos.sort001.nl004 +dos.sort002.nl006 +fcor.001.4      +fkcor.001.4      +fkval.002.2      +fval.002.4      out.scflow     +symmetry
+dos.sort001.nl005 +dos.total          +fcor.002.1      +fkcor.002.1      +fkval.002.3      =.in              =.pipe         +symminfo
+dos.sort001.nl006 +dos.total.l001     +fcor.002.2      +fkcor.002.2      +fkval.002.4      =.in.bak          =.pipeband     +vatom.001
+dos.sort001.nl007 +dos.total.l002     +fcor.002.3      +fkcor.002.3      +fval.001.1       =.in.bak.forces   =.pipekmesh    +vatom.002
```

**=.sym** contains information on the optimized structure



Do not manipulate this file (written by fedit)

Optimized position of F

F: 0.9826012122 0.9826012122 0.9826012122

# Example II: molecule $\text{CrF}_4$

```
$ cd DFT
```

```
$ ls
```

```
+band      +dos.sort002      +dos.total.l003      +fcor.002.4      +fkcor.002.4      +fval.001.2      +log      +points
=.dens      +dos.sort002.nl001      +dos.total.l004      +fedit      +fkval.001.1      +fval.001.3      out.band      +run
+dos.sort001      +dos.sort002.nl002      +dos.total.l005      +fedithelp      +fkval.001.2      +fval.001.4      out.fedit      =.str_template
+dos.sort001.nl001      +dos.sort002.nl003      +fcor.001.1      +fkcor.001.1      +fkval.001.3      +fval.002.1      out.feditband      =.sym
+dos.sort001.nl002      +dos.sort002.nl004      +fcor.001.2      +fkcor.001.2      +fkval.001.4      +fval.002.2      out.feditkmesh      =.sym.bak
+dos.sort001.nl003      +dos.sort002.nl005      +fcor.001.3      +fkcor.001.3      +fkval.002.1      +fval.002.3      out.scf      =.sym.bak.forces
+dos.sort001.nl004      +dos.sort002.nl006      +fcor.001.4      +fkcor.001.4      +fkval.002.2      +fval.002.4      out.scflow      +symmetry
+dos.sort001.nl005      +dos.total      +fcor.002.1      +fkcor.002.1      +fkval.002.3      =.in      =.pipe      +symminfo
+dos.sort001.nl006      +dos.total.l001      +fcor.002.2      +fkcor.002.2      +fkval.002.4      =.in.bak      =.pipeband      +vatom.001
+dos.sort001.nl007      +dos.total.l002      +fcor.002.3      +fkcor.002.3      +fval.001.1      =.in.bak.forces      =.pipekmesh      +vatom.002
```

In **out.scf**, the sites are listed

```
Number of sites :    5
Number of real sites :    5
Number of nonempty real sites :    5
No.  Element  WPS  CPA-Block    X          Y          Z
  1   Cr       1      1      0.00000000000000000000  0.00000000000000000000  0.00000000000000000000
  2   F       2      2      1.856847181144710      1.856847181144710      1.856847181144710
  3   F       2      3     -1.856847181144710      1.856847181144710     -1.856847181144710
  4   F       2      4      1.856847181144710     -1.856847181144710     -1.856847181144710
  5   F       2      5     -1.856847181144710     -1.856847181144710      1.856847181144710
```



The 4 F sites must be listed in the `=.wandef` files,  
even if they correspond to the same Wyckoff position

# Example II: molecule $\text{CrF}_4$

- Have a look at the file `1_RunCrF4FPLO_WF.Quanty`

```
WanDef=[ [
```

```
...
```

```
wandef
```

```
on
```

```
    name Cr_3d-2
```

```
    emin -10
```

```
    emax  10
```

```
    de 0
```

```
    contrib
```

```
        site 1
```

```
        difvec 0 0 0
```

```
        xaxis 1 0 0
```

```
        zaxis 0 0 1
```

```
        orb 3d-2
```

```
        fac  1
```

For Cr atom 5 similar definitions:

3d-2 3d-1 3d0 3d1 3d2

```
...]]
```

# Example II: molecule $\text{CrF}_4$

- Have a look at the file `1_RunCrF4FPLO_WF.Quanty`

For F atoms, the script can include a loop over the sites to avoid writing 4 times the same thing:

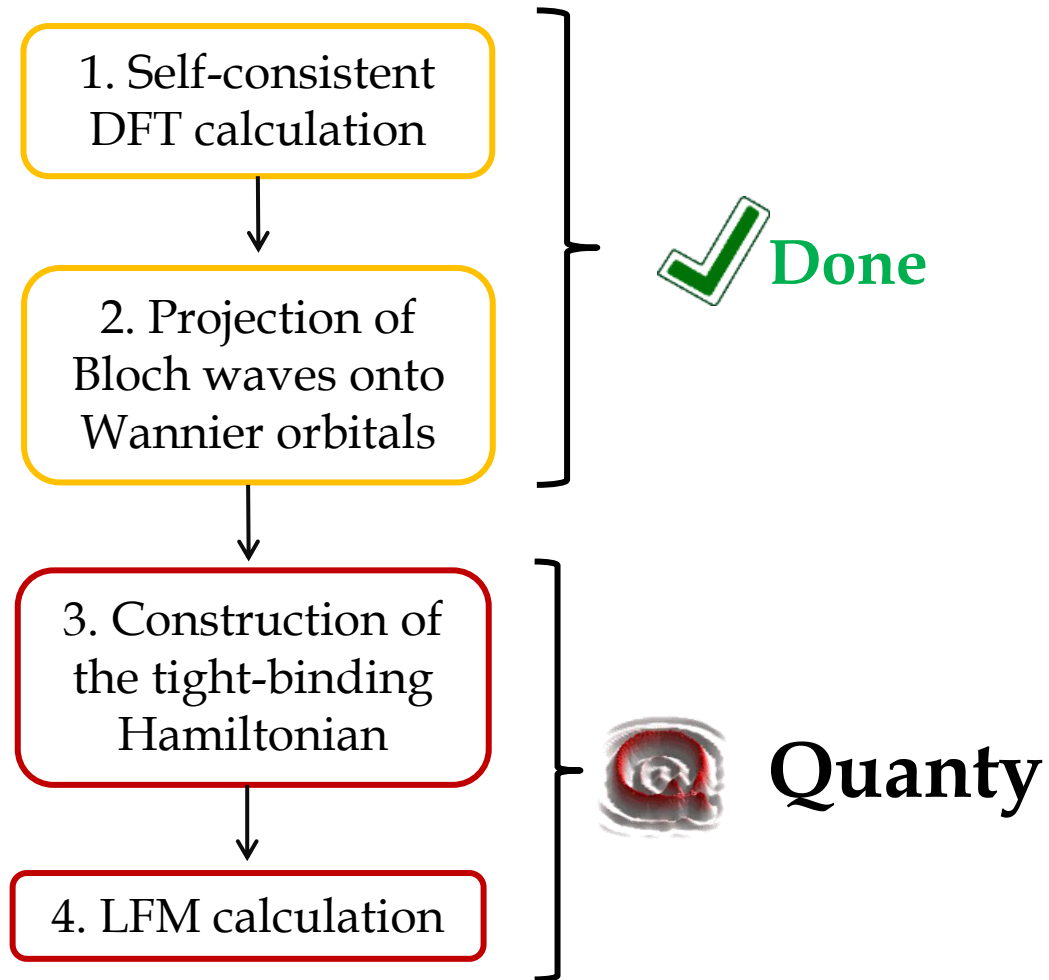
```
for i=1,4 do
WanDef = WanDef..[[

wundef
on      name F_]]..i..[[_3p-1
      emin -10
      emax  10
      de 0
      contrib
          site ]]..(i+1)..[[
          difvec 0 0 0
          xaxis 1 0 0
          zaxis 0 0 1
          orb 3p-1
          fac    1

...]]
```

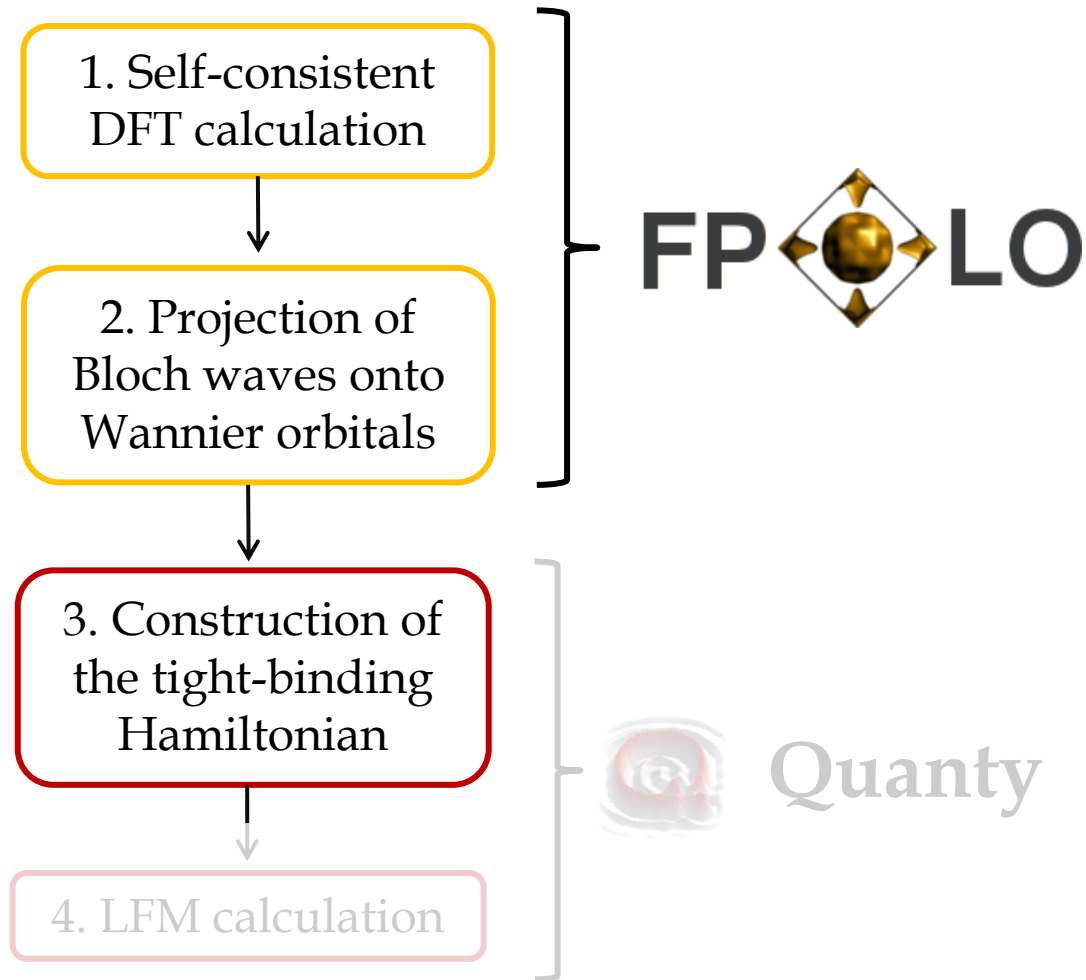
- `$ Quanty 0_RunCrF4FPLO.Quanty`

# Script method



## Part 2: Tight binding Hamiltonian and LFM calculation

# “*Ab-initio*” Ligand Field Multiplet: procedure



# Example I: NiO - Ground state calculation

View the script `2_groundstate.Quanty` in folder `NiO/LFMcalc`



# Tight-Binding Hamiltonian

- Read the file out.wan and

```
FPLOOut = FileReadDresdenFPLO("DFT/out.wan")
```

- Create the tight binding Hamiltonian for the crystal

```
TB = TightBindingDefFromDresdenFPLO(FPLOOut)
```



```
print(TB)
```

Display all the hopping parameters in the crystal

# Tight-Binding Hamiltonian

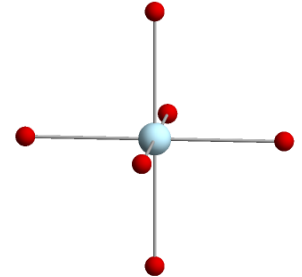
- Choice of a cluster of atoms

Find all atoms inside a sphere of radius 4 centered at the origin:

```
NewCluster =  
FindAllAtomsInsideSphere(TB.Atoms, TB.Cell, {0, 0, 0}, 4)
```

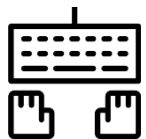
=> 5 Ni *d* and 6\*3 O *p*

Ni surrounded by 6 O  
3.95 Bohr away



- Construction of the matrix  $H_{TB}$   $(H_{TB})_{\mu'\mu} = \langle W_{\mu'} | H_{\text{DFT}} | W_{\mu} \rangle$

```
HDFTLarge, ClusterTB = CreateClusterHamiltonian(TB, {"open",  
NewCluster}, {"AddSpin", true}))
```



```
print(HDFTLarge)
```

Operator form

```
print(OperatorToMatrix(Hamiltonian))
```

Matrix form  
(23x23 matrix)

$$H = \left( \begin{array}{c|c} 3d \text{ orbitals} & 3d - L \text{ coupling} \\ \hline 3d - L \text{ coupling} & L \text{ orbitals} \end{array} \right)$$

# Linear combination of ligand orbitals

- Compute rotation matrix

Some linear combinations of ligand p orbitals do not interact with the metal  $d$  orbitals  $\Rightarrow$  Possibly drastic reduction of the basis set size

In  $O_h$  symmetry 5 ligand orbitals (two  $e_g$  and three  $t_{2g}$ ) instead of 18 orbitals

Symmetry Adapted Linear Combination (see Ballhausen - Introduction fo Ligand Field Theory - p.154)

0.	0.	0.	0.	0.	0.5	-0.5	0.	0.	0.5	0.	0.	0.	0.	-0.5	0.	0.	0.
0.5	0.	0.	0.	0.5	0.	0.	0.	0.	0.	0.	0.	0.	-0.5	0.	-0.5	0.	0.
0.	-0.58	0.	0.29	0.	0.	0.	0.	-0.29	0.	0.	0.29	-0.29	0.	0.	0.	0.58	0.
0.	0.	0.5	0.	0.	0.	0.	-0.5	0.	0.	0.5	0.	0.	0.	0.	0.	0.	-0.5
0.	0.	0.	0.5	0.	0.	0.	0.	0.5	0.	0.	-0.5	-0.5	0.	0.	0.	0.	0.

In Quantity, the block band diagonalize algorithm computes the required rotation matrix in general symmetry:

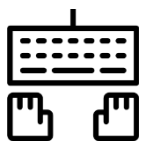
```
RTB, T = BlockBandDiagonalize(ClusterTB, [{"Ni", {0, 0, 0}}])
```

Tight binding  
object

Rotation matrix

Tight binding  
object

Starting block  
(Ni  $d$  block will remain unchanged)

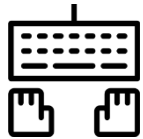


`Print(T)`

# Linear combination of ligand orbitals

- Construction of the rotated tight-binding Hamiltonian

```
HDFT, ClusterTB, IndexFPLO = CreateClusterHamiltonian(RTB,  
{"open", RTB.Atoms}, [{"AddSpin", true},  
{"ReturnTBIndicesDict", true}])
```



```
print(HDFT)
```

```
print(OperatorToMatrix(HDFT))    20x20 matrix (with spin)
```

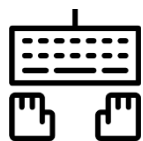
The number of orbitals was reduced from 46 to 20

⇒ drastic reduction of the number of possible states from  $\binom{46}{44}=1035$  to  $\binom{20}{18}=190$

# Linear combination of ligand orbitals

- Construction of the rotated tight-binding Hamiltonian

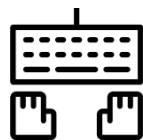
```
HDFT, ClusterTB, IndexFPLO = CreateClusterHamiltonian(RTB,  
{"open", RTB.Atoms}, [{"AddSpin", true},  
{"ReturnTBIndicesDict", true}])
```



**print(IndexFPLO)** List the automatically generated indices

- Generate more convenient indices.

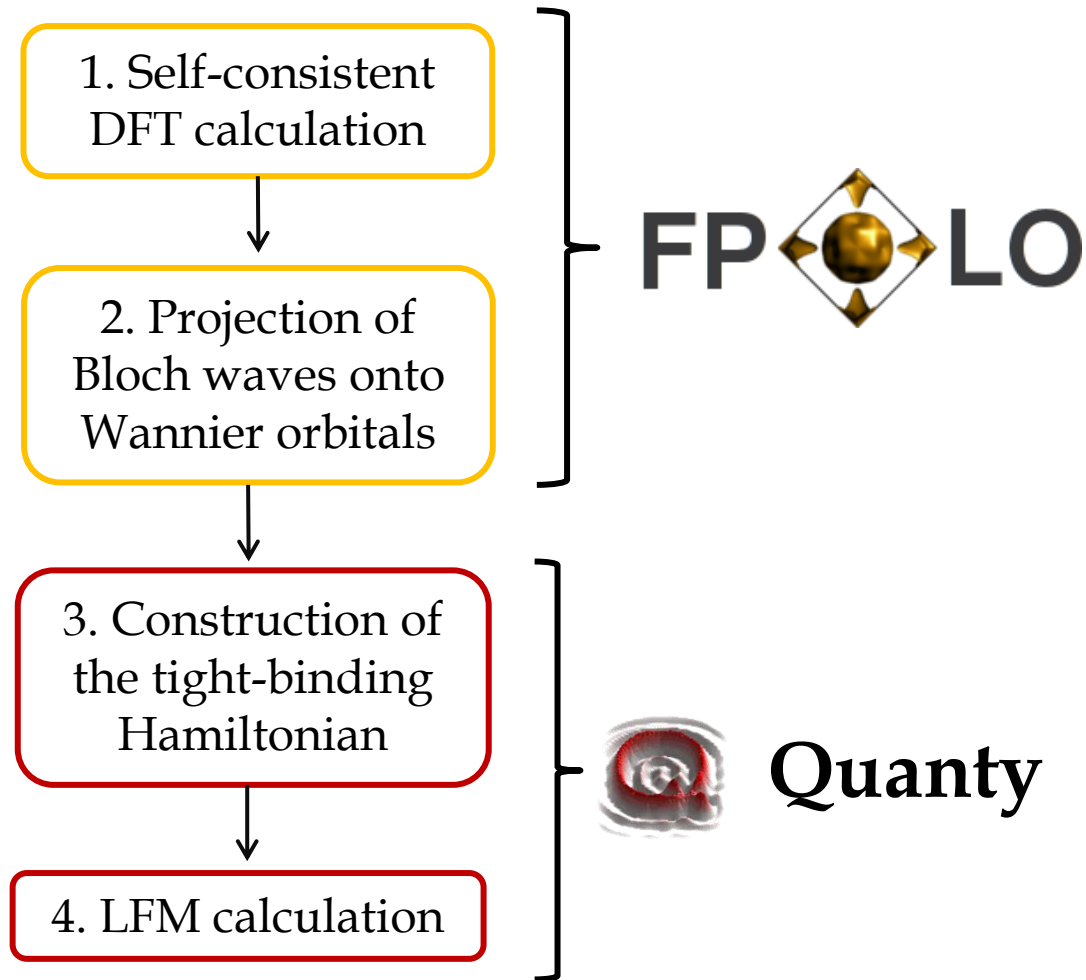
```
Index, NFermi =  
CreateAtomicIndicesDict({"Ni_3d", "Ligand_d"})
```



**print(Index)** List the groups of indices

*e.g.* Ni\_3d\_Up: 3d orbitals of Ni with spin up  
use `Index["Ni_3d_Up"]` to refer to those

# “*Ab-initio*” Ligand Field Multiplet: procedure



# Hamiltonian

$$\begin{aligned} \text{Hamiltonian} = & \text{HDFT} \\ & -F0dd * \text{OppF0MFDFT} - F2dd * \text{OppF2MFDFT} - F4dd * \text{OppF4MFDFT} \\ & + F0dd * \text{OppF0} + F2dd * \text{OppF2} + F4dd * \text{OppF4} \\ & + Bz * (2 * \text{OppSz} + \text{OppLz}) + \text{zeta\_3d} * \text{Oppldots} \end{aligned}$$

# Hamiltonian

```
Hamiltonian = HDFT  
-F0dd*OppF0MFDFT-F2dd*OppF2MFDFT-F4dd*OppF4MFDFT  
+ F0dd * OppF0 + F2dd * OppF2 + F4dd * OppF4  
+ Bz * (2*OppSz + OppLz) + zeta3d * OppIdots
```

**Coulomb operator:**  $F_{0dd} \text{Opp}F_{0\_3d} + F_{2dd} \text{Opp}F_{2\_3d} + F_{4dd} \text{Opp}F_{4\_3d}$

```
OppF0 =  
Rotate(NewOperator("U",10,Index["Ni_3d_Up"],Index["Ni_3d_Dn"],{1,0,0}),YtoZMatrix("d"))  
OppF2 =  
Rotate(NewOperator("U",10,Index["Ni_3d_Up"],Index["Ni_3d_Dn"],{0,1,0}),YtoZMatrix("d"))  
OppF4 =  
Rotate(NewOperator("U",10,Index["Ni_3d_Up"],Index["Ni_3d_Dn"],{0,0,1}),YtoZMatrix("d"))
```

↖  
Operators are rotated  
to a basis of tesseral harmonics

↑  
Within Ni *d* shell

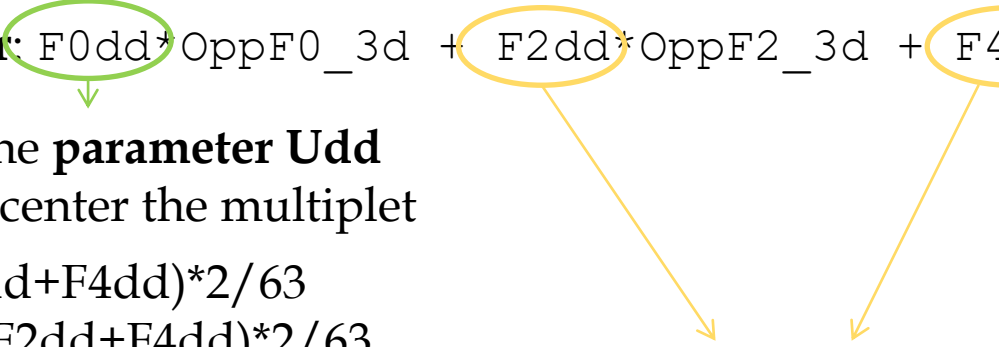
↑  
F<sub>0</sub>, F<sub>2</sub> or F<sub>4</sub>



# Hamiltonian

```
Hamiltonian = HDFT  
-F0dd*OppF0MFDFT-F2dd*OppF2MFDFT-F4dd*OppF4MFDFT  
+ F0dd * OppF0 + F2dd * OppF2 + F4dd * OppF4  
+ Bz * (2*OppSz + OppLz) + zeta_3d * OppIdots
```

**Coulomb operator:**  $F0dd * OppF0\_3d + F2dd * OppF2\_3d + F4dd * OppF4\_3d$



Determined from the **parameter Udd**  
which refers to the center the multiplet

$$Udd = F0dd + (F2dd + F4dd) * 2/63$$
$$\Rightarrow F0dd = Udd - (F2dd + F4dd) * 2/63$$

Computed from the DFT radial wave-functions

```
slaterIntegrals = GetSlaterIntegrals({"3d"}, correlatedRadialFunctions)  
F2dd = slaterIntegrals["3d 3d 3d 3d"][2] * EnergyUnits.Ha.value  
F4dd = slaterIntegrals["3d 3d 3d 3d"][4] * EnergyUnits.Ha.value
```

# Hamiltonian

```
Hamiltonian = HDFT
              -F0dd*OppF0MFDFT-F2dd*OppF2MFDFT-F4dd*OppF4MFDFT
              + F0dd * OppF0 + F2dd * OppF2 + F4dd * OppF4
              + Bz * (2*OppSz + OppLz) + zeta_3d * Oppldots
```

## Double counting corrections

Coulomb interactions are included in the Kohn-Sham potential. Addition of the full Coulomb interaction leads to a "double counting" problem. A correction is needed:

- (i) Determine DFT density matrix of the  $d$  shell (5x5 matrix)
- (ii) Calculate the Coulomb operator **within mean-field approximation** including self interaction (in KS-DFT the electron interacts with the entire electron density which includes its own density):

```
OppF0MFDFT =
MeanFieldOperator(OppF0, rhod, [{"AddDFTSelfInteraction", true}])
OppF2MFDFT =
MeanFieldOperator(OppF2, rhod, [{"AddDFTSelfInteraction", true}])
OppF4MFDFT =
MeanFieldOperator(OppF4, rhod, [{"AddDFTSelfInteraction", true}])
```

# Hamiltonian

$$\begin{aligned} \text{Hamiltonian} = & \text{HDFT} \\ & -F0dd * \text{OppF0MFDT} - F2dd * \text{OppF2MFDT} - F4dd * \text{OppF4MFDT} \\ & + F0dd * \text{OppF0} + F2dd * \text{OppF2} + F4dd * \text{OppF4} \\ & + Bz * (2 * \text{OppSz} + \text{OppLz}) + \text{zeta\_3d} * \text{Oppldots} \end{aligned}$$

External magnetic field and spin orbit coupling within the Ni  $3d$  shell

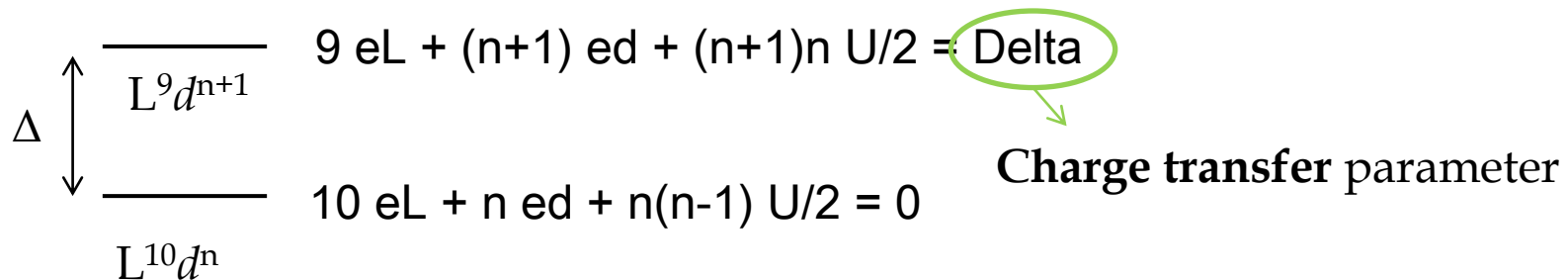


You can add an exchange field  $H_z * \text{OppSz}$

# Hamiltonian

$$\begin{aligned} \text{Hamiltonian} = & \text{HDFT} \\ & -F0dd * \text{OppF0MF DFT} - F2dd * \text{OppF2MF DFT} - F4dd * \text{OppF4MF DFT} \\ & + F0dd * \text{OppF0} + F2dd * \text{OppF2} + F4dd * \text{OppF4} \\ & + Bz * (2 * \text{OppSz} + \text{OppLz}) + \text{zeta\_3d} * \text{Oppldots} \end{aligned}$$

- Set on-site energies:



$$\begin{array}{l} \Delta \begin{array}{c} \text{---} \\ | \\ \text{---} \end{array} \begin{array}{l} L^9 d^{n+1} \\ L^{10} d^n \end{array} \end{array}$$

$$\begin{aligned} 9 eL + (n+1) ed + (n+1)n U/2 &= \Delta \\ 10 eL + n ed + n(n-1) U/2 &= 0 \end{aligned}$$

Charge transfer parameter

$$ed = (10 * \Delta - nd * (19 + nd) * Udd / 2) / (10 + nd)$$

$$eL = nd * ((1 + nd) * Udd / 2 - \Delta) / (10 + nd)$$

```
OperatorSetTrace(Hamiltonian, ed, Index["Ni_3d"])
```

```
OperatorSetTrace(Hamiltonian, eL, Index["Ligand_d"])
```

# Compute eigenvectors

- Compute eigenvectors

```
StartRestrictions = {NFermi, 0,  
{DeterminantString(NFermi, Index["Ni_3d"]), nd, nd},  
{DeterminantString(NFermi, Index["Ligand_d"]), 10, 10}}
```

```
psiList = Eigensystem(Hamiltonian, StartRestrictions, Npsi)
```

```
$ Quantity 2_groundstate.Quantity
```

⇒ Print the energy and expectation values of several operators  
for the `Npsi` first eigenstates

# Example I: NiO - XAS Calculation

```
$ cp 2_groundstate.Quanty 3_XAScalc.Quanty
```

and modify `3_XAScalc.Quanty`

Or copy `3_XAS.Quanty` from `../Scripts`

- Add the Ni 2p shell in the index list

Change line:

Index, NFermi =

CreateAtomicIndicesDict({"Ni\_3d", "Ligand\_d", "Ni\_2p"},  
{ {"Ni", {"Ni\_2p", "Ni\_3d"}} })

Add line:

HDFT.NF = NFermi

The number of fermionic states is now 26



# Example I: NiO - XAS Calculation

- Define the  $2p$ - $3d$  Coulomb repulsion operator

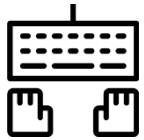


Add lines:

```
OppUpdF0 = Rotate(NewOperator("U", NFermi,  
Index["Ni_2p_Up"],Index["Ni_2p_Dn"],  
Index["Ni_3d_Up"],Index["Ni_3d_Dn"], {1,0},  
{0,0}),YtoZMatrix({"Ni_3d","Ligand_d","Ni_2p"}))  
OppUpdF2 = Rotate(NewOperator("U", NFermi,  
Index["Ni_2p_Up"],Index["Ni_2p_Dn"],  
Index["Ni_3d_Up"],Index["Ni_3d_Dn"], {0,1},  
{0,0}),YtoZMatrix({"Ni_3d","Ligand_d","Ni_2p"}))  
OppUpdG1 = Rotate(NewOperator("U", NFermi,  
Index["Ni_2p_Up"],Index["Ni_2p_Dn"],  
Index["Ni_3d_Up"],Index["Ni_3d_Dn"], {0,0},  
{1,0}),YtoZMatrix({"Ni_3d","Ligand_d","Ni_2p"}))  
OppUpdG3 = Rotate(NewOperator("U", NFermi,  
Index["Ni_2p_Up"],Index["Ni_2p_Dn"],  
Index["Ni_3d_Up"],Index["Ni_3d_Dn"], {0,0},  
{0,1}),YtoZMatrix({"Ni_3d","Ligand_d","Ni_2p"}))
```

# Example I: NiO - XAS Calculation

- Compute  $2p$ - $3d$  Slater integrals



Add line:

Upd = 8.5

Add line:

```
correlatedRadialFunctions =  
ReadFPLOBasisFunctions({"2p"}, "DFT/+fcor.001.1")
```

Change line:

```
slaterIntegrals =  
GetSlaterIntegrals({"2p", "3d"}, correlatedRadialFunctions)
```

Add lines:


```
F2pd = slaterIntegrals["3d 2p 3d 2p"][2] *  
EnergyUnits.Ha.value  
G1pd = slaterIntegrals["2p 3d 3d 2p"][1] *  
EnergyUnits.Ha.value  
G3pd = slaterIntegrals["2p 3d 3d 2p"][3] *  
EnergyUnits.Ha.value  
F0pd = Upd + (1/15)*G1pd + (3/70)*G3pd
```



# Example I: NiO - XAS Calculation

- Construct a larger rotation matrix

Change line:

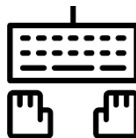
 ~~YtoZdouble = YtoZMatrix({"Ni\_3d", "Ligand\_d"})~~

YtoZtriple = YtoZMatrix({"Ni\_3d", "Ligand\_d", "Ni\_2p"})

Replace **all** YtoZdouble by YtoZtriple

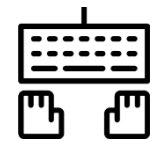
- Change the definition of the `OppN_Ni` to include  $2p$  states

Change line:

 `NewOperator("Number", NFermi, Index["Ni"], Index["Ni"],  
{1,1,1,1,1,1,1,1,1,1,1,1,1,1,1})`

# Example I: NiO - XAS Calculation

- Define electric dipole transition operators for several polarization vectors



Add lines:

```
t=math.sqrt(1/2)
```

```
Akm = {{1,-1,t},{1, 1,-t}}
```

```
TXASx = NewOperator("CF", NFermi, Index["Ni_3d_Up"],  
Index["Ni_3d_Dn"], Index["Ni_2p_Up"], Index["Ni_2p_Dn"],  
Akm)
```

```
Akm = {{1,-1,t*I},{1, 1,t*I}}
```

```
TXASy = NewOperator("CF", NFermi, Index["Ni_3d_Up"],  
Index["Ni_3d_Dn"], Index["Ni_2p_Up"], Index["Ni_2p_Dn"],  
Akm)
```

```
Akm = {{1,0,1}}
```

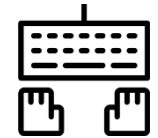
```
TXASz = NewOperator("CF", NFermi, Index["Ni_3d_Up"],  
Index["Ni_3d_Dn"], Index["Ni_2p_Up"], Index["Ni_2p_Dn"],  
Akm)
```

```
TXASr = t*(TXASx - I * TXASy)
```

```
TXASl =-t*(TXASx + I * TXASy)
```

# Example I: NiO - XAS Calculation

- Define a Hamiltonian including  $p$ - $d$  interactions and  $2p$  spin orbit coupling



Add line:

```
zeta_2p = 11.51
```

Add lines:

```
XASHamiltonian = HDFT - F0dd * OppF0MFDFT - F2dd *  
OppF2MFDFT - F4dd * OppF4MFDFT  
+ F0dd * OppF0 + F2dd * OppF2 + F4dd * OppF4  
+ Bz * (2*OppSz + OppLz) + zeta_3d * Oppldots_3d  
+ zeta_2p * Oppldots_2p  
+ F0pd * OppUpdF0 + F2pd * OppUpdF2 + G1pd *  
OppUpdG1 + G3pd * OppUpdG3
```



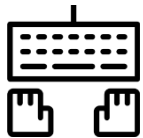
You can add an exchange field  $H_z * OppSz$

# Example I: NiO - XAS Calculation

- Add on-site energies

$\Delta$ 
 $\begin{array}{c} \text{---} \\ \updownarrow \\ \text{---} \end{array}$ 
 $2p^6L^9d^{n+1}$ 
 $6\text{ ep} + 9\text{ eL} + (n+1)\text{ ed} + (n+1)n\text{ Udd}/2 + 6(n+1)\text{ Upd} = \text{Delta}$

$2p^6L^{10}d^n$ 
 $2p^5L^{10}d^{n+1}$ 
 $5\text{ ep} + 10\text{ eL} + (n+1)\text{ ed} + (n+1)(n)\text{ Udd}/2 + 5(n+1)\text{ Upd} = 0$ 
 $6\text{ ep} + 10\text{ eL} + n\text{ ed} + n(n-1)\text{ Udd}/2 + 6n\text{ Upd} = 0$



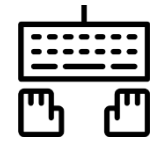
Add lines:

$$\begin{aligned} \text{epfinal} &= (10 * \text{Delta} + (1 + \text{nd}) * (\text{nd} * \text{Udd} / 2 - (10 + \text{nd}) * \text{Upd})) / (16 + \text{nd}) \\ \text{edfinal} &= (10 * \text{Delta} - \text{nd} * (31 + \text{nd}) * \text{Udd} / 2 - 90 * \text{Upd}) / (16 + \text{nd}) \\ \text{eLfinal} &= ((1 + \text{nd}) * (\text{nd} * \text{Udd} / 2 + 6 * \text{Upd}) - (6 + \text{nd}) * \text{Delta}) / (16 + \text{nd}) \end{aligned}$$

```
OperatorSetTrace(XASHamiltonian, epfinal, Index["Ni_2p"])
OperatorSetTrace(XASHamiltonian, edfinal, Index["Ni_3d"])
OperatorSetTrace(XASHamiltonian, eLfinal, Index["Ligand_d"])
```

# Example I: NiO - XAS Calculation

- Compute eigenstates of Hamiltonian



Change line:

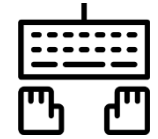
`Npsi = 3`

Change line:

```
StartRestrictions = {NFermi, 0,  
{DeterminantString(NFermi, Index["Ni_3d"]), nd, nd},  
{DeterminantString(NFermi, Index["Ligand_d"]), 10, 10},  
{DeterminantString(NFermi, Index["Ni_2p"]), 6, 6}}
```

# Example I: NiO - XAS Calculation

- Compute the spectra for the  $N_{\text{psi}}$  lowest eigenstates



Add lines:

```
XASSpectra = CreateSpectra(XASHamiltonian, {TXASz, TXASr,  
TXASl}, psiList, {"Emin",-15}, {"Emax",25}, {"NE",2000},  
{"Gamma",0.1}))
```

Broadening

```
XASSpectra.Broaden(0.4, {{-3.7, 0.45}, {-2.2, 0.65}, { 0.0,  
0.65}, { 1.0, 2.00}, { 6 , 2.00}, { 8 , 0.80}, {13.2,  
0.80}, {14.0, 0.90}, {16.0, 0.90}, {17.0, 2.00}})
```

Isotropic spectra for the ground state

```
XASIsoSpectra = Spectra.Sum(XASSpectra,{1,0,0, 1,0,0,  
1,0,0}))
```

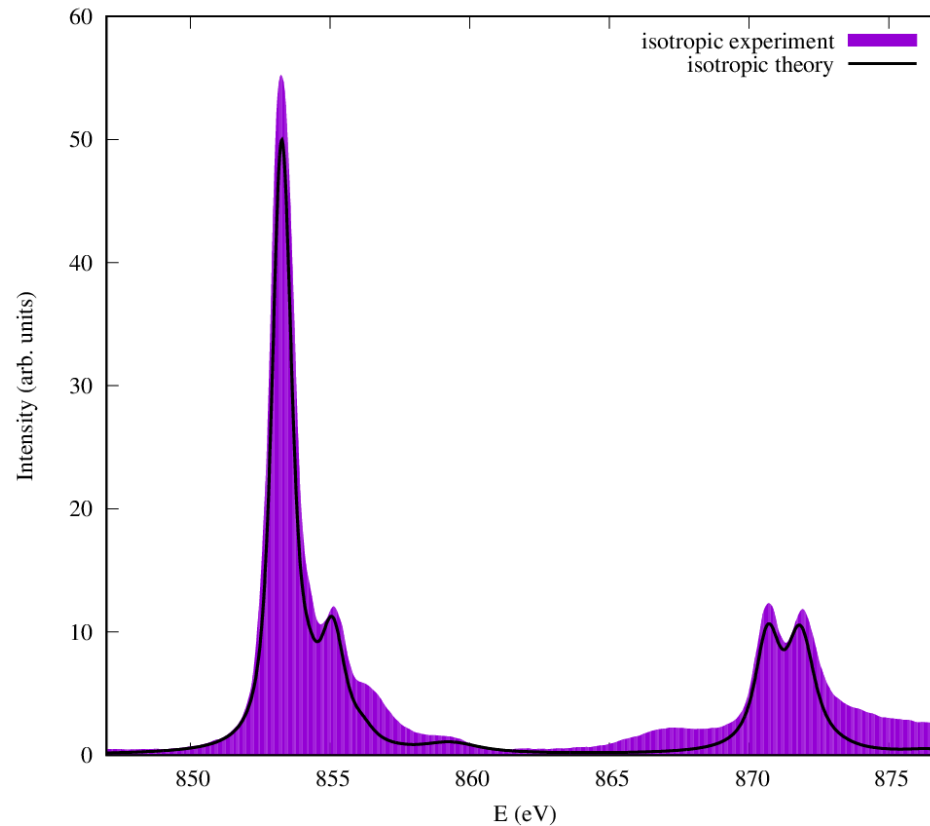
Write files

```
XASSpectra.Print({"file","XASSpec.dat"})  
XASIsoSpectra.Print({"file","XASIsoSpec.dat"})
```

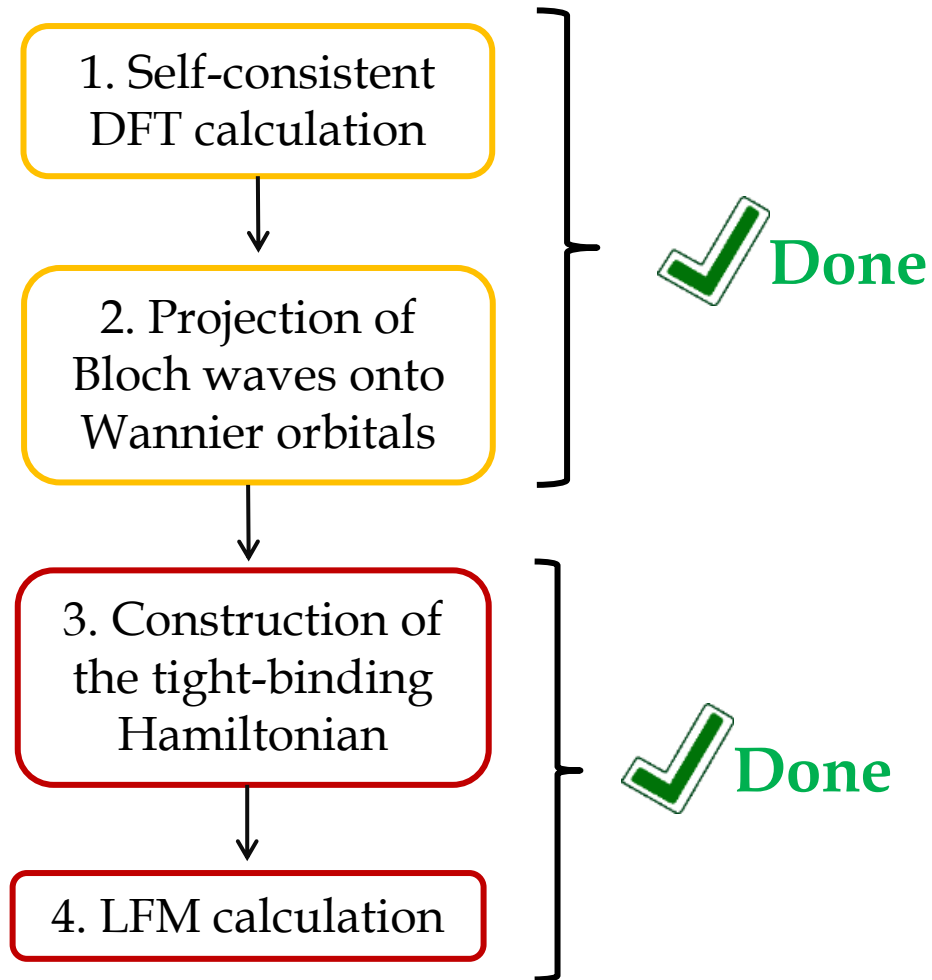
# Example I: NiO - XAS Calculation

```
$  Quanty 3_XAS.Quanty
```

```
$  gnuplot XASSpec.gnuplot  
    (generate.ps files)
```

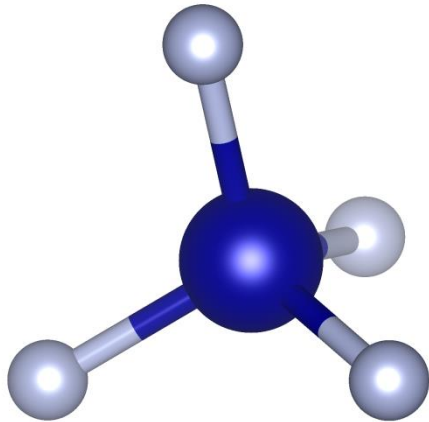


# Script method





# Example II: $\text{CrF}_4$ – Ground state calculation



Changes to make from NiO groundstate calculation are listed in  
**fromNiOtoCrF4** in folder **CrF4/LFMcal**