TBKOSTER

TIGHT-BINDING MAGNETIC MOLECULAR DYNAMICS FOR EVERYONE

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Abstract

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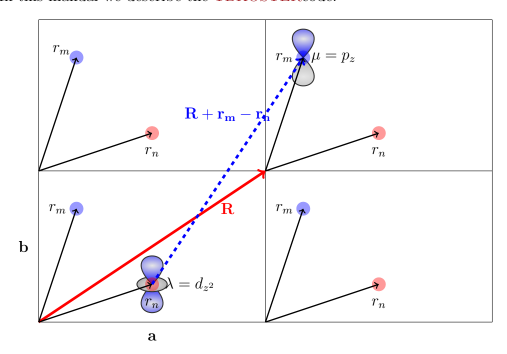
^{*}Supported by CEA

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Preface

In this manual we describe the TBKOSTERcode.



The manual contains:

- \bullet Background theory for magnetic Tight-Binding with several atoms per unit-cell, periodic boundary conditions and overlaps.
- Installation guide
- \bullet Instructions to run the code
- TBKOSTER process and post-process options.
- Description of input and output files.

CHAPTER

Background theory

- General TB formalism with several atoms per unit-cell, non-collinear magnetism, overlap integrals and periodic boundary conditions is described thoroughly in file TB.pdf.
- NRL TB Hamiltonian is described in the review paper: An efficient magnetic tight-binding method for transition metals and alloys C. Barreteau, D. Spanjaard, MC Desjonqueres Comptes Rendus Physique 17 406-429 (2016).
- Magnetic Force Theorem (MFT) and its application to the calculation of magneto-crystalline anisotropy can be found in: *Magnetocrystalline anisotropy energy of Fe(001) and Fe(110) slabs and nanoclusters: A detailed local analysis within a tight-binding model* D. Li, A. Smogunov, C. Barreteau, F. Ducastelle, and D. Spanjaard Phys. Rev. B **88** 214413 (2013).
- **TB** spin dynamics is described in *Spin dynamics from a constrained magnetic tight-binding model* Ramon Cardias, Cyrille Barreteau, Pascal Thibaudeau, and Chu Chun Fu Phys. Rev. B **103**, 235436 (2021).

Installation

The first step is to download the latest release of TBKOSTER from its github repository. To proceed, you have to check that git is installed:

\$ locate git

Then, clone the distant repository to your local one:

\$ git clone https://github.com/araven/TBKOSTER.git

In order to build this documentation, a LaTeX distribution is mandatory, including some external packages such as pdflatex, bibtex and htlatex.

2.1 Linux

Preferred method: on Ubuntu 16.04 and later, install gfortran and cmake to compile TBKOSTER. Be sure to get the mandatory related dependencies of these packages.

 $\$ sudo apt install cmake gfortran clang doxygen graphviz \ libblas-dev liblapack-dev libomp-dev texlive-latex-base \ texlive-latex-extra tex4ht

Be sure to update to cmake release 3.9 or higher. In order to build the code, to the root of TBKOSTER directory:

2.2. MacOS

```
$ if ! test -d linux; then mkdir linux; fi
$ cd linux
$ cmake ...
```

To get access to the OpenMP implementation, then update your packages with openmp support.

If you want to use Intel compiler, MKL and Intel Lapack libraries, you have to tell this to cmake as, in sequential:

```
$ BLA_VENDOR=Intel10_64lp_seq FC=ifort cmake ...
```

Be sure that the variable MKLROOT is set accordingly.

In order to get good numerical performance, you have to produce a Release version as :

```
$ cmake -DCMAKE_BUILD_TYPE=Release ...
```

You can combine all these options. If you prefer to prepare an installation with a given installed Lapack library and gfortran try:

```
$ BLA_VENDOR=OpenBLAS FC=gfortran cmake ...
```

2.2 MacOS

Preferred method: on MacOS 10.11 and later, install the cmake, lapack and gfortran with llvm support, with the ports subsystem (http://www.macports.org)

```
\ sudo port install cmake gcc6 libgcc6 gcc_select \ llvm-3.9 llvm_select lapack libomp
```

In order to build the code, to the root of TBKOSTER directory:

```
$ if ! test -d macos; then mkdir macos; fi
$ cd macos
$ cmake ...
```

For both Linux and MacOS platform, you can invoke cmake with Release or Debug option in order to deploy these releases. Simply try

To get access to the OpenMP implementation, then update your ports with openmp package. You can easily change your settings with

```
$ port select —summary
$ port select —set llvm mp-llvm -3.9
$ port select —set gcc mp-gcc6
```

In order to get good numerical performance, you may use the OpenBLAS library and produce a Release version as:

```
$ port install openblas
$ BLA_VENDOR=OpenBLAS cmake -DCMAKE_BUILD_TYPE=Release ...
```

2.3 MS Windows

For Windows earlier than 10 release 1709, you have to consider the following method: download and follow the instructions to install MSYS2 software distro and building platform (http://www.msys2.org/). Open an MSYS console and first upgrade the whole system:

```
$ pacman -Syu
get the compilation toolchain :
$ pacman -S mingw-w64-x86_64-toolchain
get the gfortran compiler and lapack library :
$ pacman -S mingw-w64-x86_64-gcc-libgfortran
$ pacman -S mingw-w64-x86_64-openblas
```

get the cmake program to control the software compilation process using simple platform and compiler independent configuration files, and to generate native makefiles and workspaces that can be used in the compiler environment of your choice:

\$ pacman —S cmake

In order to build the code, open a MSYS MINGW 64-bit console. To the root of TBKOSTER directory :

```
$ if ! test -d win; then mkdir win; fi
$ cd win
$ cmake -G"MinGW-Makefiles" -DCMAKESH="CMAKESH-NOTFOUND" ...
$ mingw32-make
```

In order to run TBKOSTER.exe, be sure you have the TERM and TER-MINFO environment variables up to date into your .bashrc file:

6 2.3. MS Windows

```
$ export TERM=xterm
```

\$ export TERMINFO=/c/Program Files/msys/mingw64/share/terminfo

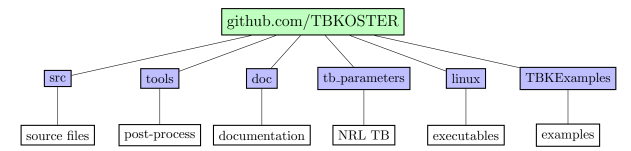
No OpenMP implementation for MS Windows has been tested.

For Windows higher than 10 release 1709, you simply have to activate the optional Windows SubSystem for Linux, download the Ubuntu Package from the Microsoft Marketplace and follow the instructions of the Linux section of this manual.



Running the code

3.1 Directories



3.2 linux directory

The "linux" directory should contain...

3.3 running TBKOSTER

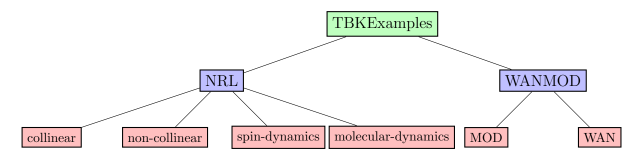
TBKOSTERneeds an in_master.txt file to run. in_master.txt contains all the parameters of the calculation. This file will be described in details in Input File Command Reference section.

CHAPTER

Getting Started

To get started we recommend to first run the various examples that are provided within the TBKOSTER git repository.

In the TBKOSTER directory one can find a TBKExamples directory organized as follow;



In each sub-directory one can find a README.md file and a list of examplesxx directories each containing a jobs.sh file. This jobs.sh can be executed by the command ./jobs.sh



Input File Command Reference

5.1 input files

$in_master.txt$

The file in_master.txt is the main input file of TBKOSTER.

namelist	Variable	Description
&calculation		
	pre_processing	type of pre-processing calculation:
	(char)	'txt2xyz' \rightarrow write geometry in xyz format,
	1 1	directory for saving pre-processing calculation:
	(char)	'txt2xyz' \rightarrow default value,

processing (char) type of processing calculation: $\operatorname{'scf'} \to \mathbf{self\text{-}consistent},$ 'md' \rightarrow molecular-dynamics. 'sd' \rightarrow spin-dynamics. type of post-processing calculation (always prepost_processing (char) ceded by a scf calculation): 'band'→ band structure calculation, 'dos' \rightarrow density of states calculation. 'forces' \rightarrow forces calculation. 'mft' \rightarrow magnetic force theorem. &units energy (char) Energy unit: $'hau' \rightarrow Hartree atomic units$, 'rau'→ Rydberg atomic units, 'ev' \rightarrow electronvolts. length (char) Length unit: 'hau'→ Hartree atomic units, 'rau'→ Rydberg atomic units, 'nm' \rightarrow nanometer.

'ang' \rightarrow angstrom.

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time (char) time unit:

'hau' \rightarrow Hartree atomic units,

'rau' \rightarrow Rydberg atomic units,

'fs' \rightarrow femtoseconds.

mass (char) Mass unit:

'hau' \rightarrow Hartree atomic units,

'rau' \rightarrow Rydberg atomic units,

'rau' \rightarrow Rydberg atomic units,

'g/mol' \rightarrow g/mol.

&element ne (int) Number of different elements in the system. symbol(i) (char) Symbol of the elements in the system i = $1, \cdots, ne$ no(i) (int) number of orbitals of element $i = 1, \dots, ne$ (optional) o(i,1:no(i)) (int) list of orbitals $i = 1, \dots, ne$ input charge of element $i = 1, \dots, ne$ q(i) (real) s input charge of element $i = 1, \dots, ne$ q_s(i) (real) p input charge of element $i = 1, \dots, ne$ q_p(i) (real) d input charge of element $i = 1, \dots, ne$ q_d(i) (real) u_lcn(i) (real) U (in eV) value of element $i = 1, \dots, ne$ for local charge neutrality u_lcn_d(i) (real) Ud (in eV) value of element $i = 1, \dots, ne$ for local d charge neutrality Stoner parameter of element $i = 1, \dots, ne$ for of i_stoner_d(i) (real) d orbitals for magnetic systems. xi_so_p(i) (real) spin-orbit coupling constant of element i = $1, \dots, ne$ for p orbitals

	<pre>xi_so_d(i) (real)</pre>	spin-orbit coupling constant of element $i = 1, \dots, ne$ for d orbitals
&element_t	tb	
	tb_type (char)	type of TB calculation
		$'$ nrl' \rightarrow NRL TB ,
		'mod' \rightarrow model TB (input=mod.dat),
		'wan' \rightarrow Wannier TB (input=hr.dat).
	filename(i) (char)	file with the NRL TB parameters of element $i = 1, \dots, ne$.
&lattice		
	v_factor (real) v(1:3,3) (real)	Multiplication factor applied to the lattice vectors. 3 translation vectors $\mathbf{a} = v(1,:), \mathbf{b} = v(2,:), \mathbf{c} = v(3,:),$
&atom		
	ns (int)	Type of magnetic system.
		$'1' \rightarrow$ non magnetic system,
		$2' \rightarrow$ collinear spin,
		$'4' \rightarrow$ non-collinear spin.
	<pre>na (int) k_spiral(3) (real) ntag (int)</pre>	Number of atoms in the system. spiral vector for spin-spiral calculations (defaut $k_spiral=(0,0,0)$). Number of tags to give names to differents atoms.
	<pre>stag(i) (int) tag(i) (char) pbc(3) (int)</pre>	Number of atoms of tag $i=1,\dots,ntag$. tag of atom $i=1,\dots,ntag$. number of unit-cell in the three periodic directions to search for neighbours. if $pbc(i)=0$ then there is no periodicity in direction i .

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	r_coord (char)	type of coordinate for atom positions,
		' direct ' \rightarrow in fractions of a, b and c coordinate,
		'cartesian' $\rightarrow xyz$ coordinates,
	x(i,3) (real) m_coord (char)	coordinates of atom $i = 1, \dots, na$, type of coordinate for magnetism,
		'spherical' $\rightarrow m$, θ and ϕ coordinate,
		'cartesian' $\rightarrow xyz$ oordinates,
	m_listing (char)	type of magnetic assignation,
		$\mathbf{by_atom'} \rightarrow \mathbf{by} \ \mathbf{atom},$
		'by_tag' \rightarrow by tag,
	m(i,3) (real)	magnetic coordinates of atom $i=1,\cdots,na$ or $i=1,\cdots,ntag$
	<pre>lambda_pen_listing (char)</pre>	type of magnetic assignation,
	(Char)	$\mathbf{by_atom'} \rightarrow \mathbf{by} \ \mathbf{atom},$
		'by_tag' \rightarrow by tag,
	lambda_pen(i) (real)	magnetic penalization factor of atom $i=1,\cdots,na$ or $i=1,\cdots,ntag$
&mesh	(-	
	type (char)	type of k mesh
		$\mathbf{p}' \rightarrow \mathbf{p}' \rightarrow \mathbf{p}$ monkhorst pack mesh,
		'path' \rightarrow path in k-space,
		'list' \rightarrow list of k-vectors.

	$x_{\tt coord}\ ({\rm char})$	type of k mesh
		' direct ' \rightarrow in fractions of a , b and c coordinate,
		'cartesian' $\rightarrow xyz$ coordinates,
	<pre>gx(3) (int) dx(3) (int) nxs (int) gxs (int) xs(1:nxs,3) (int) xs_label(1:nxs) (int) nx (int) x(1:nx,3) (int)</pre>	$gx = (n_a, n_b, n_c), n_i$ integer, when type=mp. $dx = (d_a, d_b, d_c) d_i = 0, 1$ shift when type=mp. number of symmetry point when type=path. number of points between two consecutive symmetry-point when type=path. coordinate of the symmetry points when type=path label (G,X,M,K etc) of the symmetry points when type=path number of k points when type=list. coordinate of the k points when type=list
&hamilto	onian_tb	
	e_e_interaction (char)	type of electronic interaction ${}'\mathbf{stoner'} \! \to \mathrm{monkhorst\ pack\ mesh},$ ${}'\mathrm{ujb'} \! \to \mathrm{TB+U(J,B)},$
	$\begin{array}{c} {\tt m_penalization} \\ {\rm (char)} \end{array}$	type of magnetic penalization $'$ none $'$ \rightarrow no penalization
		'r' \rightarrow penalization of the amplitude of the spin moment $m(i)$ for each atom.
		$r, \theta' \rightarrow \text{penalization of the amplitude of the spin moment } m(i) \text{ and on the } \theta(i) \text{ angle for each atom.}$
		$r, \theta, \phi' \rightarrow \text{penalization of the amplitude of the spin moment } m(i) \text{ and on the } \theta(i) \text{ and } \phi(i) \text{ angles for each atom.}$
		$\theta' \rightarrow \text{penalization of the the } \theta(i) \text{ angle for each atom.}$

 ${}^{{}^{\backprime}}\!\theta,\!\phi{}^{{}^{\backprime}}\!\!\rightarrow$ penalization of the $\theta(i)$ and $\phi(i)$ an-

 $\phi' \rightarrow \text{penalization of the } \phi(i) \text{ angles for each } \phi' \rightarrow \text{penalization of the } \phi(i)$

gles for each atom.

atom.

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&energy		
	smearing (char)	type of smearing
		$\mathbf{mp'} \rightarrow \mathbf{Methfessel}$ Paxton,
		'fd' \rightarrow Fermi Dirac,
		'mv' \rightarrow Marzari-Venderbilt,
		$g' \rightarrow Gaussian,$
	<pre>degauss (real) fixed_fermi_level (logical)</pre>	electronic broadeding
		'false' \rightarrow Fermi level determined by number of electrons in the system,
		'true' \rightarrow Fermi level fixed to a given value en_f_ff1,
	<pre>en_f_ffl (entier)</pre>	value of the fixed "Fermi level" if fixed_fermi_level=true
	fixed_spin_moment (logical)	
	(logical)	$^{\prime}\mathbf{false}^{\prime}\mathbf{ ightarrow}\;,$
		'true' \to Fermi spin moment calculation, the total magnetization being equal to ${\tt m_fsm},$
	m_fsm (entier)	value of the fixed spin moment when fixed_spin_moment=true
&mixing		
	type (char)	type of smearing
		' $\mathbf{broyden'} \rightarrow \mathbf{Broyden\ mixing}$
		'linear' \rightarrow linear mixing,

	<pre>alpha (real) n_init (int) n_hist (int)</pre>	mixing coefficient $0 < \alpha < 1$ first step of Broyden mixing (default 1) number of history steps of Broyden mixing (default 50)
&scf		
	verbose (logical)	type of smearing
		'false'→ minimum writing in out_log.txt file 'true'→ verbose writing in out_log.txt file,
	<pre>delta_en (real) delta_q (real) ni_min (int) ni_max (int)</pre>	energy crtiterium for scf calculation charge crtiterium for scf calculation miniumum number of iteration (default 2) maximum number of iteration (default 50)

post-processing

a) band

band/in_band.txt

namelist	Variable	Description
&band		
	proj (char)	type of projection
		'none' \rightarrow no projection,
		'site' \rightarrow atomic site projection,
		'spin' \rightarrow spin projection (only if $ns=4$) to plot spin-textures.
		'orbit' \rightarrow orbit projection (only if $ns=4$) to plot orbit-textures.
		'spin,orbit' \rightarrow spin and orbit projection (only if $ns=4$) to plot spin and orbit-textures.

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i_min (int)	lowest band index to include in the calculation
	$(\text{default } i_m in = 1)$
$\mathtt{i_max}\ (\mathrm{int})$	largest band index to include in the calculation
	$(\text{default } i_m ax = n m ax)$
${ t na_band} \ ({ t int})$	number of atomic sites on which projection should
	be done (default na_band=0).
ia_band(1:na_band)	index of the atomic sites.

band/in_mesh.txt

Note that for band-structure calculation, although the type of mesh is by default type=mp, band structure should be performed either with verb+type+=path or verb+type+=list.

- path is used for "traditional" band-structure plotting
- list is used for 2D plot of spin, orbit or velocity textures.

namelist	Variable	Description
	type (char)	type of k mesh
		$\mathbf{p}' \rightarrow \mathbf{p}' \rightarrow \mathbf{p}$ monkhorst pack mesh,
		'path' \rightarrow path in k-space,
		'list' \rightarrow list of k-vector,
	x_coord (char)	type of k mesh
		' direct ' \rightarrow in fractions of a , b and c coordinate,
		'cartesian' $\rightarrow xyz$ coordinates,

gx(3) (int) dx(3) (int) nxs (int)	$gx = (n_a, n_b, n_c), n_i$ integer, when type=mp. $dx = (d_a, d_b, d_c) d_i = 0, 1$ shift when type=mp. number of symmetry point when type=path.
	v v 1
$\mathtt{gxs}\ (\mathrm{int})$	number of points between two consecutive
	symmetry-point when type=path.
xs(1:nxs,3) (int)	coordinate of the symmetry points when
	type=path
xs_label(1:nxs)	label (G,X,M,K etc) of the symmetry points
(int)	when type=path
$\mathtt{nx}\ (\mathrm{int})$	number of k points when type=list.
x(1:nx,3) (int)	coordinate of the k points when type=list

b) PDOS

dos/in_dos.txt

namelist	Variable	Description
&dos		
	nen (int)	number of energy points
	${\tt na_dos}\ ({ m int})$	number of atomic sites on which the pdos will be
		projected
	$ia(1:na_dos)$ (int)	index of the sites.
	${\tt en_min} \; ({ m real})$	lower bound of energy window.
	en_max (real)	upper bound of energy window.

namelist	Variable	Description
&energy		•
	smearing (char)	smearing type
		$\mathbf{mp'} \rightarrow \mathbf{Mathfessel}$ Paxton,
		'mv' \rightarrow Marzari Vanderbilt,
		'fd' \rightarrow derivative of Fermi-Dirac.
		$g' \rightarrow Gaussian.$
	${\tt degauss} ({\rm real})$	broadening

dos/in_mesh.txt

In dos/in_mesh.txt all the options of &mesh are available however for PDOS one usually only use a MP k-point grid. Hence we have only listed below the most common parameters.

namelist	Variable	Description
	type (char)	type of k mesh
		$\mathbf{p}' \rightarrow \mathbf{p}' \rightarrow \mathbf{p}$ monkhorst pack mesh,
		'path' \rightarrow path in k-space,
		'list' \rightarrow list of k-vectors.
	gx(3) (int) dx(3) (int)	$gx = (n_a, n_b, n_c), n_i$ integer, when type=mp. $dx = (d_a, d_b, d_c) d_i = 0, 1$ shift when type=mp.

c) MFT (magnetic force theorem)

The magnetic force theorem is used in two contexts: to evaluate the magnetocrystalline anisotropy or to compare the total energy of different magnetic configurations.

namelist	Variable	Description
&mft		
	calc (char)	type of calculation
		'mae' \rightarrow magnetic anisotropy calculation,
		'mconfig' \rightarrow calculation of various magnetic non-collinear configurations,
	type (char)	type of mesh in (θ, ϕ)
		'mesh' \rightarrow regular mesh over the full spherical coordinates,
		'list' \rightarrow list of several (θ, ϕ) angles,
		'path' \rightarrow path between various (θ, ϕ) angles,
	<pre>na_mft (int)</pre>	number of site on which the energy is projected (default na_mft=0)
	<pre>ia(1:na_mfr) (int)</pre>	index of the sites, if na_mft=0 no need to specify all the indices.
	nxa (int)	• calc=mae&type=mesh: (nxa,nxa) mesh over the spherical coordinates,
		• calc=mae&type=path: nxa is the number
		 points between two angles, calc=mae&ype=list: nxa is the number of angles in the list,
		• calc=mfonfig: nxa is the number of magnetic configurations.

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<pre>mconfig(na,nxa,3)</pre>	define explicitly the nxa magnetic configurations
(real)	over the na atoms of system
$\mathtt{nangle}\;(\mathrm{int})$	in case type=path; nangle is the number of angles
	(2) to generate the path.
<pre>angle_xs(nxa,3)</pre>	list of angles in spherical coordinates
(real)	

mft/in_energy.txt

namelist	Variable	Description
&energy		
	${ t smearing} \ ({ t char})$	smearing type
		$\mathbf{p}' \rightarrow \mathbf{Mathfessel}$ Paxton,
		'mv' \rightarrow Marzari Vanderbilt,
		'fd' \rightarrow derivative of Fermi-Dirac.
		$g' \rightarrow Gaussian.$
	degauss (real)	broadening

mft/in_mesh.txt

In mesh/in_mesh.txt all the options of &mesh are available however for MFT one only use a MP k-point grid. Hence we have only listed below the most common parameters.

namelist	Variable	Description
	type (char)	type of k mesh
		$'mp' \rightarrow monkhorst pack mesh,$
		'path' \rightarrow path in k-space,
		'list' \rightarrow list of k-vectors.
	gx(3) (int) dx(3) (int)	$gx = (n_a, n_b, n_c), n_i$ integer, when type=mp. $dx = (d_a, d_b, d_c) d_i = 0, 1$ shift when type=mp.

5.2 output files

All the output files (out_xxxx.txt) of TBKOSTER are in namelist format (except out_log.txt), hence they can be re-used as input files for further calculations.

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$out_log.txt$

out_log.txt is the main output file of TBKOSTER. The first lines of out_log.txt contains a summary of all the parameters (including the defaults) used for the calculation. Then, if verbosity=.true. TBKOSTER print out in out_log.txt the charge and various quantities at each scf iteration.

out_charge.txt

out_charge.txt contains:

- the mulliken charge q_mul(1:na,1:3,0:ns-1)
- and net charge rho_net(1:na,1:1:no,1:no,1:ns)

Importantly out_charge.txt can be copied in in_charge.txt whenever one needs a restart, for instance when a calculation is not fully converged.

post-process output

a) txt2xyz

This option is used when one want to visualize the evolution of the magnetic configuration during the scf cycle. This is particularly useful in the case of non-collinear magnetism. This option necessitate the setting of verbose=.true. in &scf. out_atom_tb_xx.txt files are saved during the scf iterations. The txt2xyz option transforms the .txt file into .xyz that contain position and magnetization of atoms in the unit-cell. These files can be visualized via avogadro software to generate a movie showing the convergence process.

b) dos

The main output file is dos/out_dos.txt file. It contains in namelist format:

- dos_tot(1:nE,1:ns): Total density of states
- dos_'orb' (1:na_dos,1:nE,1:ns): Projected density of states on different sites, different orbitals (orb=s,px, py etc..) and different spin index.

c) band

The main output file is band/out_band.txt file. It contains in namelist format:

• en_k(1:nh,1:nk,1:ns): Eigenvalues nh: size of Hamiltonian, nk: number of k points.

If proj=site

• w_band_site(1:nh,1:nk,1:na_band,1:norb,1:ns): weight of wavefunction on different sites, different orbitals and different spin index.

If proj=spin

• w_band_spin(1:nh,1:nk,0:na_band,1:4): average value of the Pauli matrices $\sigma_x, \sigma_y, \sigma_z$ (and its norm) for band index ih, k point ik, site ia_band.

If proj=orbit

• w_band_orb(1:nh,1:nk,0:na_band,1:4): average value of the orbital matrices Lx, L_y , L_z (and its norm) for band index ih, k point ik, site ia_band.

d) mft

The main output file is mft/out_mft_xxx.txt file. It contains for each magnetic configuration xxx in namelist format:

- mconfig(1:na,1:2): Magnetic configuration (θ, ϕ) for each atom of the unit-cell
- mft_'orb'(1:na_mft): band energy projected on the selected sites for each orbitals (orb=s,px,py etc..).
- mft_sum: Sum of the band energies projected on the selected sites of the unit-cell
 - mft_tot: Total band energy, it should be equal to mft_sum if nsite=na

tools output

The various tools read the output files of TBKOSTERin namelist format and reorganize them in a format directly readable by plotting software (xmgrace, gnuplot etc..)

a) pdos.x

pdos.x generates the following files:

- dos_tot.dat: Total density of states
- pdos-s.dat, pdos-p.dat, pdos-d.dat and pdos-spd.dat: PDOS

b) bands.x

bands.x generates different types of files depending on the options for the projection (proj) and for the type of mesh.

mesh=path

If proj=none

• band.dat: band structure along the k path

If proj=site

- band.dat: band structure along the k path
- band_weight_site_orb.dat: band structure along the k PATH and associated weight on the defined sites.

If proj=spin and/or proj=orb

• band_weight_spinorb.dat: band structure along the k PATH and associated amplitude of the spin and/or orbit.

mesh=list

Note that the "list" k-points are usually generated in cartesian coordinates via the build_kpoints.f90 program. "list" mesh option is used to generate 2D maps

If proj=spin and/or proj=orb

bands.x generates two files:

- fermi.dat: 2D maps for Fermi surface
- fermi.dat: 2D maps for "spin" or "orbit" vector field.

c) mft.x

There are two type of MFT calculations: calc= mae or calc=mconfig.

calc=mae

mft.x reads the out_mft_xxx.txt files and creates several files:

If type=path

- mae_angle.dat: MAE along the (θ, ϕ) path where the first angle of the path is taken as the reference zero energy.
- mae_atom.dat: MAE energy decomposed on the different sites, and evaluated as the difference between the first and last magnetic orientation.

 \bullet mae.xyz: Same as mae_atom.dat but in (x,y,z) coordinates.

If type=mesh

• mae_angle.dat: MAE along the (θ, ϕ) mesh where the first angle of the path is taken as the reference zero energy.

calc=mconfig

mft.x reads the out_mft_xxx.txt files and creates one summary file:

• mft_config.dat which contains the list of MFT band energies for the various configurations.

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