



Scientific Computing & Modelling

BAND-GUI Reference Manual

**ADF Program System
Release 2008.01**

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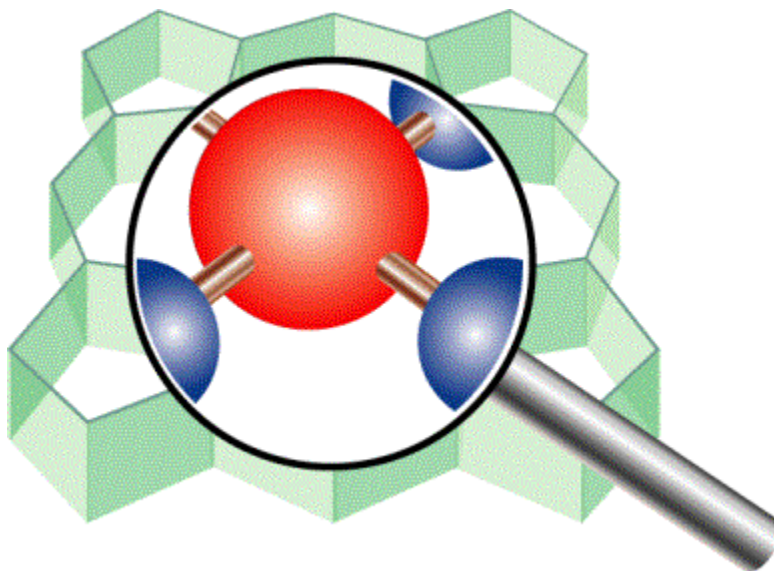


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Introduction

This document will provide a basic reference manual of the BAND-GUI. BAND-GUI is the Graphical User Interface for the ADF-BAND package.

If you are new to the BAND-GUI we advise you to study the BAND-GUI tutorial.

The BAND-GUI modules

The BAND-GUI consists of several modules:

ADFjobs

This utility (`$ADFBIN/adfjobs`) manages your jobs: run a job on your local machine or on remote machines. It also serves as a interface to all files belonging to your job, and it servers as a convenient launcher of the other ADF-GUI modules.

BANDinput

A utility program (`$ADFBIN/bandinput`), which enables users to easily create BAND jobs. You can use BANDinput to define your periodic structure (geometry), and to set details of your BAND job using an easy-to-use graphical user interface. BANDinput will generate the complete job script for you. This script takes care of running BAND.

ADFview

A simple program (`$ADFBIN/adfview`) that displays volume data, such as the electron deformation density and orbitals.

BANDstructure

This program (`$ADFBIN/bandstructure`) shows a plot of the electronic band structure

ADFmovie

This program (`$ADFBIN/adfmovie`) shows the progress of a geometry optimization

ADFTail

A minor ADF-GUI utility (`$ADFBIN/adftail`) that will just show the contents of a text file, updating when the text file grows (like the UNIX `tail -f` command). It is used to monitor the 'logfile'. The progress of an BAND (and ADF) calculation is always written to this file.

ADFoutput

A Basic Output Browser (`$ADFBIN/adfoutput`) for the output generated by BAND (and ADF).

The SCM (logo) menu

All BAND-GUI modules have the SCM menu on the left-hand side, on most systems represented by a small SCM logo. You can easily switch between the different modules of the BAND-GUI using this menu:

- Input: activate BANDinput
- View: activate ADFview

- Band Structure: activate BANDstructure
- Movie: activate ADFmovie
- Logfile: activate ADFtail
- Output: activate BOB

When you use the SCM menu while some file is connected to the current BAND-GUI module, the selected BAND-GUI module will be activated showing data belonging to the same calculation. The title bar of any BAND-GUI module shows which file is connected, if any. Thus, you can easily switch between viewing the logfile, output, input, orbitals, etc, all belonging together.

When you use the SCM menu when no file is connected (the title bar just shows the name of the module), the selected BAND-GUI module will be started without file.

On most platforms you can tear off the SCM menu by selecting the dashed line at the top of the menu.

Automatic bug reports

Please enable the automatic mailing of bug reports. These reports contain detailed information about the internal state of the BAND-GUI module having some problem, but do not contain personal information. Information about the structure of your molecule **is** included.

You can enable this by setting the global environment variable `SCM_ERROR_MAIL`, for example in your login script:

```
SCM_ERROR_MAIL=errors@scm.com
export SCM_ERROR_MAIL
```

This will ensure that the bug report will be sent to the e-mail address you specify. If you specify your own e-mail address you can see exactly what kind of information is contained in the error report.

Your comments and bug reports are very welcome. Please send them to `support@scm.com`, or use the 'Feedback' menu command from the 'Help' menu of the BANDGUI modules.

Environment Variables

The BANDGUI is normally installed as part of the ADF package. In the following table the environment variables that are specific for the BANDGUI are listed:

Name	Meaning	Default Value
<code>SCM_ERROR_MAIL</code>	e-mail address for error reports	no error e-mail sent
<code>SCM_GUIRC</code>	location of the preferences file	<code>\$HOME/.scm_guiirc</code>
<code>SCM_TPLDIR</code>	location of the templates directory	none (no extra templates loaded)
<code>SCM_STRUCTURES</code>	location of the structures directory	none (no extra structures loaded)
<code>SCM_RESULTDIR</code>	location of the results directory	none (current directory used)
<code>DISPLAY</code>	X-window display to use	required (for all X11 programs) except on Windows

Environment Variables

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SCM_ERROR_MAIL	e-mail address for error reports	no error e-mail sent
SCM_GUIRC	location of the preferences file	\$HOME/.scm_guiirc
SCM_TPLDIR	location of the templates directory	none (no extra templates loaded)
SCM_RESULTDIR	location of the results directory	none (current directory used)
DISPLAY	X-window display to use	required (for all X11 programs) except on Windows

Mouse Interaction

Rotate, Translate and Zoom

In the modules that provide a 3D view of your periodic structure (currently BANDinput, ADFview) you can rotate, translate and zoom using the mouse.

Drag with the mouse: press a mouse button, and move it while holding it down. A one-button mouse Click is the same as a Left Click. Which mouse button, and which modifier key you press at the same time, determines what will happen:

Rotate	Left
Rotate in-plane	Ctrl-Left
Translate	Middle, or Alt-Left
Zoom	Right, or Command-Left (drag up or down)

The rotate, translate and zoom operations change how you look at the molecule. They do not change the coordinates. When you Right or command-Left without dragging the mouse, a popup menu will appear.

In BANDinput operating with the mouse on the selection will move the selection only. In that case the geometry of your molecule (and thus the coordinates) will change. Zooming the selection will move it perpendicular to the screen. You operate on the selection by starting the drag operation with the mouse above a selected object.

In the View menu you can select either 'Mouse as trackball' or 'Mouse as joystick'. If 'Mouse as trackball' is selected, you need to drag with the mouse (move the mouse with a button pressed down). If 'Mouse as joystick' is selected you just need to press and keep the button pressed down. The direction of movement etc will depend on the position of the mouse with respect to the center of the 3D view area. Note that 'Mouse as joystick' disables some pop-up menus.

Selecting

In the modules that provide a 3D view of your molecule (currently BANDinput, ADFview) you can make selections using the mouse.

Click on an object: make a new selection with it

Click in space: clear selection

Shift-Click on object: add or remove it from the selection

Shift-Drag in space: add all objects within the rectangle to the selection

Pop-up menus: Select all similar elements or all bonds by using the pop-up menus

In some modules there are additional ways to select objects using menu commands. Furthermore, one can select atoms from the list in the coordinates window.

Pop-up menus

In many cases pop-up menus are attached to objects (atoms, bonds, empty space, peaks, levels, ...) that allow you to do something with that particular object. To get the pop-up menu (a one-button mouse button is the same as a Left mouse button):

Press and hold the Right mouse button on the object: the pop-up appears almost directly.

Press and hold the Left mouse button on the object: it will take a small time for the pop-up to appear.

If a pop-up menu is available it will appear.

BAND-GUI modules

ADFjobs

See the [ADF-GUI ADFjobs documentation](#).

BANDinput

Introduction

BANDinput is a utility program (\$ADFBIN/bandinput), which enables ADF-BAND users to easily create BAND jobs. You can use BANDinput to define your molecule (geometry), and to set details of your BAND job using an easy-to-use graphical user interface. BANDinput will generate the complete job script for you. This script takes care of running BAND programs as required. You can also use BANDinput to run these script files on your local machine in the background.

Starting BANDinput

If you have installed the ADF package correctly, the bandinput command is located in \$ADFBIN.

If \$ADFBIN is included in your PATH environment variable, you can start the ADFinput program with the following command:

```
bandinput [filename]
```

The filename is optional. BANDinput only handles files that were created by BANDinput before, which have a .band extension.

An alternative method to start BANDinput: select the Input command from the SCM menu.

Under windows you can start BANDinput by simply double-clicking the icon on the desktop.

Menu Commands

File menu

New

Same as quitting BANDinput and starting again without specifying a file name.

Open...

Open an existing BANDinput file (with .band extension).

Import Coordinates...

Use this menu command to import the geometry of your molecule from file.

You can import coordinates from a .runkf file generated by BAND, a .t21 file generated by ADF, from a .band file as saved by BANDinput, or from a text file (for example, a .mol or a .xyz file).

If you import coordinates from a .t21 file note that the extension must be .t21. A name like 'TAPE21' will not work. Also, bond information and lattice vectors are not present, just the coordinates are imported and bonds will be guessed (and no distinction will be made between different kinds of bonds).

Importing from a .runkf, or .band file is straight-forward: coordinates, bond information and lattice vectors are present so you will get exactly what you saved.

Importing from a .mol file will also give you both the coordinates and bond information of the file, but not the lattice vectors.

Importing from a text file is rather flexible: BANDinput needs three real numbers next to each other. These will be interpreted as x, y and z coordinate. One additional integer or the abbreviation of an element is also needed to identify the kind of atom.

To be recognized as real, the real number must contain a '.' (dot), and at least one digit before or after the dot. Real numbers with exponents (E or D) are **not** recognized.

If an integer is used to specify the element (the nuclear charge), it may **not** contain a '.' (dot).

Bond information and lattice vectors are not imported, even if present in your file (unless importing from a .band file). After you have imported some coordinates, BANDinput tries to guess the bonds between the imported atoms. This might not be very accurate.

Z-matrix import is not available.

Directly after the 'Import Coordinates...' command the newly imported atoms are selected. This makes it easy to reposition them with respect to other atoms that may already be present, remove the automatically guessed bonds, or use other operations on the newly imported atoms and bonds.

Export Coordinates...

Export the current geometry (unit cell only) as a simple space separated list of element name and xyz coordinates. The number of lines will match the number of atoms. No bond information is written to the text file.

You will be prompted to specify a file name.

Save

Save the current state of what is present in BANDinput. If you have not saved before, BANDinput will ask you to specify a file name.

Not only the .band file will be saved, but also a matching .run file which is a run script corresponding to your input.

Save As...

Save the current state of what is present in ADFinput in a file with a name of your choice.

Not only the .band file will be saved, but also a matching .run file which is a run script corresponding to your input.

Revert...

After opening a .band file with BANDinput and making some changes, you can use the 'Revert...' command to undo all your changes. It is the same as quitting without saving, and opening the same file again.

Save Picture...

Save a picture of your periodic structure (only the drawing area with your structure, no input options) in a file.

The format used to save your picture is determined by the extension of the file name you specify. If you do not specify a known extension, it will use the standard picture format as specified using the 'Default Picture Format' menu.

Default Picture Format

Use the submenu to select the format to use when saving a picture.

Note that this is just the default to use, the user specifies the format with the extension. When the extension is not recognized (or when no extension is specified) the default format will be used.

Run

Start the BAND program as selected in all the input options.

This is done by telling ADFJobs to execute the saved run script. If you have made changes in BANDinput, you will first be asked to save the changes. Just as the Save menu command, this will also save the run script (with the .run extension). Next the run script is executed on your local machine, and ADFtail is started to monitor the progress of the calculation using the logfile.

Preferences...

The Preferences... menu command gives access to the 'Preferences' input panel. You may also select this panel from the pull-down menu on top of the input panels.

This panel is used to set a number of preferences for the ADF-GUI and BAND-GUI, not only for BANDinput.

To save the preferences, click on the Save button in the Preferences panel.

Quit

Stop BANDinput, ask you to save changes if you made any.

Edit menu

Undo

Undo the last operation. It uses a stack of many operations, so you can use the Undo command repeatedly.

Redo

If you have undone some operation, you can use Redo to do it again.

Cut

Make a copy of the current selection, and next delete the original.

Cut, Copy and Paste work within text fields, and with atoms and bonds in the drawing area.

Copy

Make a copy of the current selection.

Cut, Copy and Paste work within text fields, and with atoms and bonds in the drawing area.

Paste

Paste the current copy.

Cut, Copy and Paste work within text fields, and with atoms and bonds in the drawing area.

Clear

Delete the current selection.

Group

The currently selected atoms and bonds will be grouped together. Once grouped, if you select any of the group members the whole group will be selected.

You may nest groups if you wish, the original group structure will be remembered.

Ungroup

Remove the grouping of the currently selected items.

If you ungroup a nested group, only the top grouping will be removed and you will recover your original groups.

Set origin

Translate all atoms such that the selected atom will be the new origin. If nothing is selected, the center of all atoms will be the new origin. If more than one atom is selected, the center of the selected atoms will be the new origin.

Bond Lengths constrained

When creating new atoms bonded to existing atoms, BANDinput will constrain the bond length to the textbook value.

If you do not wish this to happen, select this menu command to toggle this behavior.

Add Bond

Create a bond (if possible depending on the number of free connectors) between two selected atoms.

Add Hydrogen

The 'Add Hydrogen' menu command will add hydrogen atoms to your structure until every connector is connected. The number of connectors and the number of lone pairs determine the geometry. For example, the Oxygen atom has four connectors and two lone pairs in a tetrahedral arrangement.

The hydrogens will only be added to selected atoms, or to your whole molecule if no atoms are selected.

Remove Hydrogen

The hydrogen atoms will be removed from your molecule.

If you have selected part of your molecule, only hydrogen atoms in your selection will be removed.

Fuse Atoms

Fuse atoms that are very close together (in 3D space) to a single atom. If atom types differ, one of the types will randomly be chosen.

Select All

Select all atoms and bonds in your structure.

Select Molecule

Select all atoms and bonds that are somehow connected to the current selection.

Select Connected

Select all atoms that are directly connected (by one bond) to the current selection.

Guess Bonds

Roughly guess what atoms are bonded together, depending on the distance between the atoms compared with text book values. The bond type is currently always set to single, so you probably need to make some changes.

Only BANDinput uses the bond information. The BAND program itself does not use any of the bond information.

Bonds will only be guessed for the selected atoms, leaving the remaining part of your molecule unchanged. If nothing is selected, bonds will be guessed for your whole molecule.

Remove Bonds

Remove all bonds in the current selection, or in your whole molecule if nothing is selected.

View menu

Reset View

When you use the 'Reset View' menu command, the translation, zoom and rotation settings will be adjusted such that the entire molecule is visible. The center of your molecule will be the new rotation center.

Use it for example when you translate the molecule 'out of view'.

View direction

With the sub menus you can choose for standard view orientations along the x, y, and z axis.

Fly to selection

Zoom to the center of the current selection (or of the whole molecule if nothing is selected). The zooming occurs 'real life', as if you are flying to that point. The zoom point will be the new center of rotation.

Align screen

First select three atoms defining a plane (thus, not three atoms on a line ...).

Next, use the 'Align Screen' command to rotate your molecule such that the plane defined is parallel to the screen.

Mouse as

Determine how your mouse works:

Trackball: you rotate etc by dragging around the screen. This is the default, and most intuitive, mode.

Joystick: you rotate etc by pressing down a mouse button off-center, and keep it pressed down. Your molecule will rotate, translate etc in the direction of your mouse button (with respect to the center of the drawing area). Since the press-and-keep-down conflicts with pop-up menus, they will be disabled.

Anti-alias

Use the anti-alias technique to improve the quality of the pictures. Especially sharp edges will look smoother.

It works very well, but is rather time-consuming to calculate. As a result everything will be very very slow. So use it only if you want to prepare a high-quality picture for a presentation (and have plenty of time ...).

Molecule Resolution

Set the resolution of the molecule display (the number of triangles used to represent the atomic spheres and the bonds). If you have a big structure and a low-end graphics card you can speed up display by choosing 'Low'. For high-end graphic cards you might not see a significant difference.

Background

Select the color of the background.

Periodic

If 'Show Periodic' is selected more than 1 unit cell is visualized. The number of repeated cells can be '(-1,1)' the default, or a bit more '(-2,2)'. If 'Show Lattice Vectors' is selected the axes of the lattice vectors are

drawn. With 'Parallel Perspective' you can toggle whether or not a vanishing point is used for the perspective.

Model menu

The 'Model' menu gives access to the first group of input panels on your right-hand side. They roughly specify your system.

You can also use the pull-down menu at the top right side (above the panels) to get to these panels.

Properties menu

The 'Properties' menu gives access to the second group of input panels on your right-hand side.

Details menu

The 'Details' menu gives access to the third group of input panels on your right-hand side. You can use these to specify many details that are typically not needed. Expert users will get access to more BAND options with these panels. You can use the 'User Input' Menu to add BAND options that are not available (yet) in the BAND-GUI directly.

You can also use the pull-down menu at the top right side (above the panels) to get to these panels.

Help menu

The help menu provides an easy way to get to information about the BAND-GUI. It will start a browser on your local machine, and connect to the SCM web site to get information.

Buttons and Tools

The button bar contains a number of tools and buttons:



From left to right the meaning is

Button	Description	Keyboard shortcut
Select Tool	Used to select, and to rotate, translate or zoom (with modifier keys or mouse buttons).	Esc
Atom Tools	C-tool, O-tool, N-tool, H-tool, Cl-tool: atom-tools used to <ul style="list-style-type: none">• create a C, O, N, H or Cl atom,• change an existing atom,• create bonds between existing atoms.	C, O, N, H, Cl, S & P
Element Tool	A tool used to <ul style="list-style-type: none">• create any atom (from the periodic table that will pop-up),• change an existing atom,	

	• create bonds between existing atoms.	
Top view (z)	Change to the default view along the z-axis	
Side view (x)	Change to the default view along the x-axis	
Side view (y)	Change to the default view along the y-axis	
Perspective	Toggle between drawing with and without perspective	
Periodic display	Toggle between showing a single unit cell and showing a couple of them.	

Getting and changing geometry details

If you select 2, 3 or 4 atoms some geometry information will be presented at the bottom of the screen:

- 2 atoms: distance
- 3 atoms: angle
- 4 atoms: dihedral angle
- 5 atoms: angle between two planes

Often you can also change the information displayed. The geometry of your molecule will be updated accordingly. This is not always possible: ring structures make it impossible since BANDinput does not know how to change other bonds and angles. In such a case you might temporarily remove a bond, fine-tune your geometry, and finally recreate the bond you removed. The planes of the plane angle are defined through (in order of selection) atoms 1, 2 & 3 and atoms 3, 4 & 5. The order of selection is important in all cases.

Keyboard shortcuts

Many tools and menu commands have keyboard shortcuts associated.

For menu command shortcuts, the shortcut is listed in the menu. On most UNIX systems (including Linux) you need to use the control key together with some letter. On a Macintosh (running locally) you need to use the command key together with a letter.

The following table lists the other keyboard shortcuts. Just press the indicated key without any modifier keys:

Key	Function
Esc	Select-tool (end other tool)
C	C-tool
O	O-tool
N	N-tool
H	H-tool
P	P-tool
S	S-tool
backspace	Delete selection
delete	Delete selection

Presets and Defaults

All input options have default values. However, the default values depend on the main task you have chosen, and on further properties you may select to calculate. BANDinput uses presets that are simply a collection of input values to be used together as defaults.

BAND only has a few presets, Accurate and Fast, which set the basis set and (K-space) integration accuracy. You may also define your own templates.

Use a Preset

Select the preset you want to use from the Preset menu, located in the main input options area.

All or some input values will change to the values specified in the preset you select.

Revert to preset values

If you want to undo your changes and get back to the default values as specified in the current template, simply select the template again from the Template menu.

Color Code

The input fields use a color coding to warn you they have been modified:

- No special color: the field has its original default value.
- Yellow: the field has been changed by the user (only).
- Green: the field has been changed by the preset (only).
- Red: the field has been changed by the preset, and next by the user.

The pull-down menu that you use to switch between panels uses a similar color-coding to point you to fields that have been changed:

- No special color: all fields in the panel have their original default value.
- Yellow: some fields have been changed by the user.
- Green: some fields has been changed by the preset.
- Red: some fields has been changed by the preset, and some (possibly also) by the user.

Make your own presets

It is very easy to make your own presets, collecting all or a few default values for the typical jobs you like to perform.

When the `SCM_TPLDIR` environment variable has been set, BANDinput will look for user-defined presets (when starting up) in the directory `$SCM_TPLDIR`.

Thus, to use your own presets first set up your preset directory if you have not already done so:

- make a directory in which to store your presets
- set the `SCM_TPLDIR` environment variable to point to this directory

Next, in BANDinput (restart it if it was already running):

- select the preset to start with (or None if you wish to start with an empty preset)
- edit all the fields as you would like them to be stored in a preset.
- select 'Save as (Full) Preset' from the Preset menu
- specify a file name, ending with `!.tpl`
- click on Save.

If you now check your Preset menu you will find a new entry.

The name of the preset is the file name you have chosen, but without the .tpl extension.

The difference between a Full Preset and a 'normal' Preset is that a Full Preset will save all input options, and a 'normal' preset will save only the yellow or green fields (options that have been changed by the user or by the active template).

If you wish to store only fields that you have changed yourself in the preset, make sure you start with the None preset.

Defaults

The default values that are shown when you start BANDinput are generated as follows:

- Use the Defaults preset supplied by SCM
- Use the Defaults preset that the user has defined, if any
- Use the 'None' preset

Thus, you can change the defaults by saving a preset called Defaults.tpl in your SCM_TPLDIR.

Molecule Editor Tricks

Selecting

You can make or change a selection using the mouse or using menu commands.

Making a selection with the mouse: see [Selecting](#)

Making a selection with a menu command: see [Edit Menu](#)

Pop-up menus: Select all similar elements or all bonds by using the pop-up menus.

Delete an atom

Select the atom (click on it), and press the Backspace key.

Delete a bond

Select the bond (click on it), and press the Backspace key.

Delete the selection

Make your selection, and press the Backspace key.

Make a bond

Take one of the atom tools.

Next click once on the first atom you want to connect. You will enter the bonding mode (the line to the mouse position from the atom you just clicked on will be your visual cue for the bonding mode).

Next click on the atom you want to make the bond to.

The bond will be created (if possible), and you will revert to the normal select mode.

Make a bond, alternative method

Select the two atoms that you wish to be bonded together.

Select the 'Add Bond' command from the 'Edit' menu or press ctrl-l.

Change the bond type

Select the bond or bonds to change. Use the pop-up menu and choose the required bond type from the Bond type pull-down menu.

Move an atom (possibly perpendicular to the screen)

First select the atom that you want to move.

Next, translate (middle mouse button, or alt left mouse button), but start with the mouse on the atom that you want to move.

If you wish to move the atom perpendicular to the screen: use the right mouse button (or command left mouse button) and move the mouse up or down). This is equivalent to zooming.

Rotate or translate the selection

First make your selection.

Next rotate, translate or zoom as usual, but start with the mouse in the selection. So if you click and drag the selection, ONLY the selection will be rotated or translated. If you click and drag anywhere else the whole molecule will be rotated or translated (actually, only your viewpoint).

If you 'zoom' the selection you are really moving the selected objects perpendicular to the screen (in or out the screen).

Run Script

Save your input using the Save or Save As ... menu commands.

Your input will be saved in a file with the name you specify, and at the same time a run script will be saved. It has the same name, but with '.run' appended.

To execute your calculation execute this run script.

You can also use the 'Run' command from the 'File' menu: it will save the run script first, and then execute it on your local machine, and start ADFtail to follow the execution (showing the logfile).

By default RUNKF (result data) is saved.

If the environment variable SCM_RESULTDIR has been set, the run script will change into that directory. Next it will run from there, and all result files will be stored in that directory.

If the environment variable SCM_RESULTDIR has not been set, the run script will execute in the directory where it is started, and the result files will also be located in that place.

Input options remarks

Empty fields

Some input fields do not have a value from the template. In those cases BANDinput does not specify the value, but leaves the value to be determined by the BAND program.

Coordinates

The coordinates panel shows the coordinates corresponding to the molecule visible in the molecule display. You may edit the coordinate values here as well.

User Input

You can use the User Input field to specify any kind of text. The text will be appended without any change to the end of the BAND input. This way, you may access some keys that are not (yet) available in BANDinput.

ADFview

See the [ADF-GUI ADFview documentation](#).

ADFview is used by the BAND-GUI as well as by the ADF-GUI. ADFview uses the .runkf file, which is the extension of a BAND result file.

BANDstructure

Introduction

BANDstructure shows you a plot of the band structure along a series of lines of high symmetry. The line elements are separated by vertical gray bars.

Starting BANDstructure

If you have installed the ADF package correctly, BANDstructure is located in \$ADFBIN.

If \$ADFBIN is included in your PATH environment variable, you can start the BANDstructure program with the following command:

```
bandstructure [filename]
```

The filename is optional. BANDstructure only handles RUNKF files with extension .runkf .

There is a second possibility to start BANDstructure if one of the GUI modules has already been started. All BAND-GUI modules have the SCM menu on the left-hand side, on most systems represented by a small SCM logo. Using this SCM menu ADFlevels can be started by selecting the 'Band structure' command.

Menu commands

File: Open...

Close the current project and select a new RUNKF file to open.

File: Save Postscript...

Save the current diagram as a postscript file.

File: Export to Gnuplot...

Save the current diagram as an input for the gnuplot program.

File: Quit

Stop and quit BANDstructure.

Options:Show K-points table

Shows information of all the k-points in a table.

Options:Show all K-labels

Normally only the label of the begin and end point are shown. Enabling this option, it shows also the labels of the intermediate points, which may become very crowded.

Options:Show Fermi energy

Show or hide the Fermi level indicated with a red line.

Help

Provide quick access to the documentation on the SCM website.

Pop-up menus

You activate pop-up menus by pressing the (left) mouse button on an object, and keep the mouse button pressed down for some time until the menu pops up.

ADFmovie

See the [ADF-GUI ADFmovie documentation](#).

For BAND it can handle .logfile and .runkf files, and is used to display the progress of a geometry optimization.

ADFtail

See the [ADF-GUI ADFtail documentation](#).

ADFtail is used by the BAND-GUI as well as by the ADF-GUI. ADFtail opens the .logfile file, and update it while the text file grows. The progress of an BAND (or ADF) calculation is always written to this file.

ADFoutput

See the [ADF-GUI ADFoutput documentation](#).

ADFoutput is used by the BAND-GUI as well as by the ADF-GUI. ADFoutput opens the .out file, which is the output file generated by a BAND (or ADF) calculation.