# **Qbox User Guide**

version 1.63.7

2017-03-14

http://qboxcode.org

# 1 Introduction

Qbox is a First-Principles Molecular Dynamics code. It can be used to compute the electronic structure of atoms, molecules, solids and liquids within the Density Functional Theory (DFT) formalism. It can also be used to perform first-principles molecular dynamics (FPMD) simulations using forces computed within DFT. Qbox computes the solutions of the Kohn-Sham equations using a plane-wave basis set and norm-conserving pseudopotentials. It can perform constant-temperature and/or constant-pressure simulations.

# 1.1 Recent features

Release 1.63.7 is a bugfix release. See <a href="http://qboxcode.org/trac/wiki/ReleaseNotes">http://qboxcode.org/trac/wiki/ReleaseNotes</a> .

Release 1.63.4 includes the implementation of the B3LYP functional for spin-polarized systems and the set velocity command that sets the velocity of an atom. Previously released features include:

- the partial\_charge command computing the amount of electronic charge located inside a sphere of a given radius centered on an arbitrary atom (1.63.2).
- the iter\_cmd variable used to specify the name of a command or a script that will be executed periodically during a simulation. The iter\_cmd\_period variable defines the interval between executions of the iter\_cmd script, i.e. the iter\_cmd script is executed every iter\_cmd\_period iterations. This can be used to generate plot files for animations (1.63.2). Note that the move command cannot currently be used as part of the iter\_cmd script since synchronization of atomic positions is not performed.
- The Harris-Foulkes functional used to compute the value of etotal\_int during self-consistent iterations. This provides more accurate values of etotal\_int when a combination of self-consistent and non self-consistent iterations is used (e.g. nitsef > 0 and nite>0) (1.63.2)

Qbox can use the SG15 library of Optimized Norm-Conserving Vanderbilt (ONCV) pseudopotentials available at <a href="http://www.quantum-simulation.org">http://www.quantum-simulation.org</a> ONCV potentials were built to reproduce all-electron calculations with high accuracy. Note: using ONCV potentials requires the updated XML Schema file species.xsd provided in the xml directory of the distribution.

# 2 Qbox distribution

Qbox is distributed in source form under the GPL license. For source distribution see <a href="http://qboxcode.org">http://qboxcode.org</a>. Qbox has been built on several Linux platforms including CentOS-7, CentOS-6, Fedora, Ubuntu, BlueGene/P, BlueGene/Q, Cray XE6, Cray XC30, Dell PowerEdge C8220 (TACC stampede).

Information on how to build Qbox is available at <a href="http://qboxcode.org">http://qboxcode.org</a>.

# 3 Basic Qbox operation

# 3.1 Starting Qbox

On most platforms, Qbox is started using the mpirun command. On a Linux platform using the mpich implementation of MPI, the command is

```
mpirun -np <ntasks> qbox exec input file > output file
```

Depending on your installation of MPI, the arguments of the mpirun command may differ. The input file contains a list of commands that are read and executed by Qbox. The output file created by Qbox is a valid and well formed XML document. It can be used to extract various diagnostic information after the run is completed. During a Qbox run, the output file can be used to track the progress of the simulation.

# 3.2 Qbox commands

When used in interactive mode, Qbox prints a prompt

```
[qbox]
```

and waits for the user to type a command. When used in batch mode, Qbox reads input from an input file. Output is written on standard output (stdout) and can be redirected to a file. A Qbox input file consists of a sequence of Qbox commands with one command per line. Qbox commands are

```
compute the angle formed by three atoms
angle
atom
                define an atom
bisection
                apply recursive subspace bisection to wave functions
compute_mlwf
                compute maximally localized Wannier functions
<u>constraint</u>
                manage constraints on atomic positions
distance
                compute the distance between two atoms
extforce
                add external forces on atoms
fold in ws
                fold atoms into the Wigner-Seitz cell
help
                print a short message about the use of a command
<u>kpoint</u>
                add or remove k-points
                print a list of currently defined atoms
list atoms
                print a list of currently defined atomic species
<u>list_species</u>
load
                load a sample from a file previously saved
move
                move atoms
partial charge compute the amount of charge in atom-centered spheres.
                generate a plot file with atoms and/or charge density
plot
                print the value of a Obox variable
print
<u>quit</u>
                exit Qbox
randomize_r
randomize_v
randomize_wf
                add random noise to the atomic positions
                add random noise to the atomic velocities
                add random noise to the wavefunction coefficients
                rescale all atomic velocities
<u>rescale v</u>
reset vcm
                set the velocity of the center of mass to zero
                initialize the random number generator
rseed
                run MD or electronic optimization steps
run
                save a sample on a file for later use
save
set
                assign a value to a Qbox variable
set velocity
                set the velocity of an atom
species
                define a new atomic species
status
                print a summary of the current state
                impose a strain on the sample
strain
torsion
                compute the dihedral angle defined by four atoms
!(shell escape) execute a shell command
```

Commands and their syntax are described in detail in the Section "Qbox commands". If a command is read on input and is not in the above list, Qbox interprets it as the name of an input

script and attempts to open a file having that name in the current working directory. If the file can be opened and is readable, Qbox starts interpreting each line of that file as its input. Qbox scripts can be nested. At the end of a script, Qbox returns to the previous script level and continues to read commands. At the end of the topmost level script, Qbox exits.

Unix commands can be issued within a Qbox input sequence using a shell escape "!" character at the beginning of a line. For example, the line

```
[gbox] !date
```

invokes the Unix date command.

Comments can be inserted in Qbox input by inserting a '#' character at the beginning of each comment line

```
[qbox] # print the list of all atoms
[qbox] list atoms
```

Comments can also be added at the end of a command line by inserting a "#" character where the comment starts

```
[qbox] list atoms # get a list of all atoms
```

### 3.3 Qbox variables

Calculation parameters such as the plane-wave energy cutoff are specified using Qbox variables. Qbox variables can be set using the set command. Their value can be printed using the print command. For example the command

```
set ecut 24
```

sets the variable ecut to the value 24. This causes the plane wave basis set to be resized to include all plane waves with a kinetic energy not exceeding 24 Rydberg. Other Qbox variables can be set similarly. The Qbox variables are

```
coefficient of HF exchange in PBE0
alpha PBE0
                      ionic dynamics control variable
atoms dyn
                      bisection levels in Hartree-Fock exchange
blHF
                      bisection threshold in Hartree-Fock exchange
btHF
cell
                      dimensions of the unit cell
cell dyn
                      unit cell dynamics control variable
                      control of allowed unit cell motions
cell lock
<u>cell mass</u>
                      fictitious mass of the unit cell
<u>charge mix coeff</u>
                      mixing coefficient for charge density update
<u>charge mix ndim</u>
                      Anderson dimension for charge density mixing
charge mix rcut
                      Kerker screening for charge density update
                      debug parameters (not for normal use)
debuq
<u>dt</u>
                      time step (a.u.)
                      plane wave energy cutoff (Ry)
<u>ecut</u>
                      energy cutoff of the preconditioner (Ry)
ecutprec
                      energy cutoff of the confinement potential
ecuts
                      applied electric field
<u>e field</u>
                      fictitious electronic mass (for CP dynamics)
emass
ext stress
                      externally applied stress (GPa)
```

```
fermi temp
                      Fermi temperature (K)
iter cmd
                     script executed every iter cmd period steps
iter cmd period
                     number of steps between iter cmd executions
                     number of empty states
nempty
                     net charge of the system
<u>net charge</u>
                     maximum size of process grid columns
nrowmax
                     number of spin degrees of freedom
nspin
                     algorithm used to compute dipole and quadrupole
polarization
                     dimensions of the reference unit cell
ref cell
scf tol
                     tolerance criterion for SCF iterations
stress
                     stress control variable
<u>thermostat</u>
                     thermostat control variable
th temp
                     thermostat temperature (K)
th time
                     thermostat time constant (a.u.)
th width
                     thermostat width (K)
wf diag
                     wavefunction diagonalization control variable
wf dyn
                     wavefunction dynamics control variable
                      exchange-correlation control variable
<u>XC</u>
```

Obox variables have a default value. Refer to the Section "Obox variables" for details.

# 3.4 Structure of a Qbox script

A Qbox script starts with commands defining the sample being simulated. For example, the sequence of commands

```
set cell 20 0 0 0 20 0 0 0 20 species silicon silicon.xml atom Sil silicon 3.000 0.000 0.000 atom Si2 silicon 0.000 2.000 0.000 atom Si3 silicon -3.000 0.000 0.000 atom Si4 silicon 0.000 -2.000 0.000
```

defines a sample in a cubic unit cell of size 20 a.u. (Bohr radii). The sample consists of a 4-atom silicon cluster. The definition of the species "silicon" is given in the file silicon.xml. A species definition document contains all the information needed to describe an atomic species, including the pseudopotential used to represent the electron-ion interaction. Species can also be defined by a URI as in the following command

```
species carbon http://fpmd.ucdavis.edu/potentials/C/C HSCV PBE-1.0.xml
```

When a URI is used to define a species, Qbox downloads the species definition from the corresponding web site. Note that this requires web access from the computer on which Qbox is running. This access may not be available on the compute nodes of some clusters. If web access is not available on compute nodes, the species definition file must be downloaded before starting Qbox using e.g. the wget command

```
wget http://fpmd.ucdavis.edu/potentials/C/C HSCV PBE-1.0.xml
```

which creates a local copy of the file C\_HSCV\_PBE-1.0.xml. The species command can then be invoked in a Qbox script using the local file

```
species carbon C HSCV PBE-1.0.xml
```

Species definition documents are XML documents that follow the syntax defined by the

# 4 Examples of use of Qbox

The test directory of the Qbox distribution contains examples of electronic structure calculations and molecular dynamics simulations.

# 4.1 Electronic structure of a Si₄ cluster

The following example shows how to use Qbox to compute the electronic ground state of a silicon 4-atom cluster. Approximate input atomic coordinates are used. The electronic ground state is computed using 200 iterations of the preconditioned steepest descent algorithm with Anderson acceleration (PSDA).

```
# example: electronic structure of si4
set cell 20 0 0 0 20 0 0 0 20
species silicon silicon.xml
atom Si1 silicon 3.500 0.000 0.000
atom Si2 silicon 0.000 2.000 0.000
atom Si3 silicon -3.000 0.000 0.000
atom Si4 silicon 0.500 -2.000 0.000
set ecut 12
set wf_dyn PSDA
set ecutprec 4
randomize_wf
run 0 200
```

# 4.2 Structure optimization

The following example shows how to optimize the structure of a Si<sub>4</sub> cluster. The electronic ground state is computed using 200 iterations of the preconditioned steepest descent algorithm (PSDA). The structure is then optimized using 50 steps of the steepest descent with Anderson acceleration (SDA) algorithm. Five steps of electronic optimization are performed before each ionic step.

```
# example: structure optimization of Si4
set cell 20 0 0 0 20 0 0 0 20
species silicon silicon.xml
atom Sil silicon 3.500 0.000 0.000
atom Si2 silicon 0.000 2.000 0.000
atom Si3 silicon -3.500 0.000 0.000
atom Si4 silicon 0.000 -2.000 0.000
set ecut 12
set wf dyn PSDA
set ecutprec 4
randomize wf
run 0 200
# start structure optimization
set atoms dyn SDA
set dt 100
run 50 5
```

# 4.3 Molecular dynamics simulation

The following example shows how to perform a Born-Oppenheimer molecular dynamics simulation. The electronic ground state is first computed using 200 iterations of the preconditioned steepest descent

algorithm (PSDA). A Born-Oppenheimer MD is then done by running 50 ionic steps with 5 electronic iterations at each ionic step.

```
# example: molecular dynamics of si4
set cell 20 0 0 0 20 0 0 0 20
species silicon silicon.xml
atom Si1 silicon 3.500 0.000 0.000
atom Si2 silicon 0.000 2.000 0.000
atom Si3 silicon -3.500 0.000 0.000
atom Si4 silicon 0.000 -2.000 0.000
set ecut 12
set wf dyn PSDA
set ecutprec 4
randomize wf
run 0 200
# start molecular dynamics
set atoms dyn MD
set dt 40
run 50 5
```

# 4.4 Saving and restarting a simulation

The following example shows how to save the simulation sample on a file at the end of a simulation and how to restart from the saved file. In the first script, the electronic ground state is computed using 200 iterations of the preconditioned steepest descent algorithm (PSDA). A Born-Oppenheimer MD is then done by performing 50 ionic steps with 5 electronic iterations at each ionic step. The sample is saved at the end of the first script on file si4.xml. In the second script, the sample is loaded from the restart file and 50 additional MD steps are performed.

```
# script 1: electronic ground state and 50 MD steps
set cell 20 0 0 0 20 0 0 0 20
species silicon silicon.xml
atom Si1 silicon 3.500 0.000 0.000 atom Si2 silicon 0.000 2.000 0.000
atom Si3 silicon -3.500 0.000 0.000
atom Si4 silicon 0.000 -2.000 0.000
set ecut 12
set wf dyn PSDA
set ecutprec 4
randomize wf
run 0 200
# perform 10 MD steps
set atoms dyn MD
set dt 40
run 50 5
# save the sample
save si4.xml
# script 2: restart from saved sample and
# perform 50 MD steps
load si4.xml
set wf dyn PSDA
set ecutprec 4
```

```
set atoms_dyn MD
run 50 5
# save the sample
save si4.xml
```

# 4.5 Changing the plane wave energy cutoff on the fly

The size of sample files can become large when simulating large systems. Although Qbox uses a parallel I/O algorithm to save sample data, the process of writing data to a file can be slow, especially on platforms that do not offer a parallel file system. This problem can be alleviated by saving the sample with reduced resolution for wavefunctions. The value of the variable ecut can be changed using the set ecut command at any time during the simulation. When the value of ecut changes, Qbox interpolates wavefunctions onto the plane wave basis defined by the new value. The size of the sample file can be reduced by changing the value of ecut to a small value before using the save command. When loading a sample that was saved with reduced resolution, the original resolution can be restored by first redefining the value of ecut and then reoptimizing the wavefunctions. The ability to change energy cutoff on the fly is also useful to accelerate a ground state calculation by starting the calculation at low cutoff and gradually increasing the cutoff to the final desired value. This procedure can be compared to the full approximation scheme of the multigrid method.

# 4.6 Using Qbox in client-server mode

Qbox can be used in client-server mode. This means that Qbox can run on one computer as a server that "listens" to commands being sent from another computer. Commands can be sent by a user, but also by another program (the "client", or "driver"). This allows for extensions of the functionality of Qbox in which the client and the server interact to achieve a result that is not easy to get with the conventional approach of submitting a fixed input file to Qbox. A Qbox driver (or "Qbox application") program sends commands to Qbox by writing them to a file. Qbox reads the commands, executes them, and produces an output file. The client code can then read the output file, process it, and send the next commands to Qbox. Synchronization of this process is implemented through the creation (or destruction) of a file (the "lock" file) that signals that the output (or input) file is ready for use.

Qbox is run in client-server mode by invoking it as follows

```
mpirun -np <ntasks> qbox exec -server input file output file
```

The sequence of operations in client-server mode is described below:

#### Obox:

- 1) Create a new file output file.
- 2) Read commands from input\_file and execute them until the end of file is reached. Write the output on output\_file. After executing the last command in input\_file, close output\_file and create a file named input\_file.lock to signal that Qbox is ready for more commands.
- 3) Wait for the file input file.lock to be removed by the driver.
- 4) Go to 1)

### Client:

- 1) Wait for the file input file.lock to appear.
- 2) Open the file output file and read its contents.
- 3) Analyze the contents and decide what course of action to take.

- 4) Open the file input file and overwrite it with new commands.
- 5) Close input file.
- 6) Remove input file.lock to signal that the file input file is ready.
- 7) Go to 1)

Note that in server mode, Qbox only exits when it reads the quit command. When reaching the end of file in input, Qbox does not exit but creates the lock file and then waits for the client to remove it. It then starts reading the input file again.

The contents of output\_file form a valid XML document, i.e. The XML header is repeated every time the output file is rewritten. This allows one to process each output file using XML tools, such as e.g. XSLT scripts.

A client can manage multiple copies of Qbox running in server mode. In that case, multiple copies of Qbox must be launched with different input and output file names. For example:

```
mpirun -np <ntasks> qbox_exec -server in0 out0 &
mpirun -np <ntasks> qbox_exec -server in1 out1 &
mpirun -np <ntasks> qbox_exec -server in2 out2 &
```

The client must then write on the files in0, in1, in2, and read output from the files out0, out1, out2. This allows for simulations involving multiple samples, such as replica exchange dynamics, nudged-elastic-band (NEB) simulations, or path-integral molecular dynamics.

# 5 Qbox commands

# angle

**NAME** 

angle --compute the value of the angle defined by the positions of three atoms

#### **SYNOPSIS**

angle [-pbc] atom1 atom2 atom3

### DESCRIPTION

The angle command prints the value of the angle formed by the three atoms given as arguments. The names *atom1*, *atom2* and *atom3* must refer to the names of atoms currently defined in the sample. If the -pbc option is used, the positions of the atoms are interpreted as those of the nearest atom replica, taking into account periodic boundary conditions.

### RELATED INFORMATION

list atoms, distance, torsion

### atom

**NAME** 

atom --add an atom to the current sample

#### **SYNOPSIS**

atom name species x y z [vx vy vz]

### **DESCRIPTION**

The atom command adds an atom to the current sample. The *name* argument can be any character string but must differ from all the other names of atoms in the current sample. The *species* argument must refer to an atomic species previously defined using the <u>species</u> command. The position of the atom is specified by its coordinates *x*, *y*, *z* in atomic units (Bohr). Optionally, the velocity of the atom can be specified by its components *vx*, *vy*, *vz* in atomic units (Bohr/atomic-unit-of-time). (One atomic-unit-of-time is 0.02418885 fs).

#### RELATED INFORMATION

list\_atoms, species

### bisection

**NAME** 

bisection –perform recursive subspace bisection of the wave functions

#### SYNOPSIS

bisection lx ly lz threshold

### DESCRIPTION

The bisection command transform electronic wave functions according to recursive subspace bisection, as described in [F. Gygi, Phys. Rev. Lett. 102, 166406 (2009)]. The resulting wave functions are maximally localized in rectangular domains defined by recursively subdividing the unit cell lx, ly, and lz times in the x, y, and z directions respectively. The lx, ly, lz arguments define the level of recursion used in the x, y, z directions. The threshold argument is used to print information about the bisected wave functions. For each wave function, the bisection command prints the value of the localization vector, the size of the wave function and the number of non-zero overlaps with other functions (for the given value of the threshold). The localization vector is a binary vector in which a pair of bits is associated with each of the bisecting planes of the recursive bisection process, starting at the least significant bit. Within each pair of bits, a value of 1 signifies that the wave function has significant amplitude on one side of the corresponding bisecting plane. Thus the bit pattern "11" signifies that the wave function is extended across the corresponding bisecting plane, while the bit patterns "01" and "10" signify that the wave function is localized on one side only of the bisecting plane. The bit values are defined by the amount of electronic charge located on either side of the bisecting plane. The value is 1 if the amount of charge is larger than the threshold. At the end of the bisection command, wave functions are transformed so as to be maximally localized in the subdomains defined by the bisection planes. The effect of recursive bisection can then be inspected using the plot command.

#### RELATED INFORMATION

# compute\_mlwf

**NAME** 

compute\_mlwf -compute maximally localized Wannier functions

**SYNOPSIS** 

compute mlwf

# **DESCRIPTION**

The compute\_mlwf command transforms the current wave functions into maximally localized Wannier functions following the algorithm in Computer Physics Communications 155, p. 1 (2003) and a one-sided iterative Jacobi algorithm for simultaneous diagonalization. The position of Wannier centers and the corresponding spreads are printed on output. The value of the electronic, ionic and total dipole are printed. The iterative methods stops when the decrease of the spread is sufficiently small between two iterations, or when a maximum number of iterations is reached. In that latter case, the compute\_mlwf command can be issued again to try to improve the convergence of the spread minimization. After execution of the compute\_mlwf command, the wave functions are maximally localized.

#### RELATED INFORMATION

### constraint

#### NAME

constraint -- manage constraints on atomic positions

#### **SYNOPSIS**

constraint define position constraint\_name atom1
constraint define distance constraint\_name atom1 atom2 distance [velocity]
constraint define angle constraint\_name atom1 atom2 atom3 angle [velocity]
constraint define torsion constraint\_name atom1 atom2 atom3 atom4 angle [velocity]
constraint list
constraint delete constraint\_name
constraint enforce
constraint set constraint\_name value [velocity]

### DESCRIPTION

The constraint command is used to manage the constraint set. Constraints can be of the following types: position, distance, angle or torsion. Constraints are added to the constraint set using the "constraint define" command. They can be removed from the set using the "constraint delete" command. A list of constraints can be printed using the "constraint list" command. Some constraints have an associated value that can be modified using the "constraint set" command. The atom names used in the constraint command must refer to atoms previously defined using the atom command. All constraints have a name, which allows for selective removal of constraints and for individual modification of the constraint values.

constraint define position constraint name atom1

Define a position constraint on atom *atom1*. This locks the atom into its current position. constraint define distance *constraint name atom1 atom2 distance [velocity]* 

Define a distance constraint on atoms *atom1* and *atom2*. If a *velocity* argument is given, the value of the distance will change at each time step of the simulation at a rate specified by the *velocity* argument. The *velocity* must be given in [Bohr/(a.u. time)].

constraint define angle constraint\_name atom1 atom2 atom3 angle [velocity]

Define an angle constraint on atoms *atom1*, *atom2* and *atom3*. If a *velocity* argument is given, the value of the angle will change at each time step of the simulation at a rate specified by the *velocity* argument. The *velocity* must be given in [degree/(a.u. time)].

constraint define torsion constraint name atom1 atom2 atom3 atom4 angle [velocity]

Define a torsion (or dihedral) constraint on atoms *atom1*, *atom2*, *atom3* and *atom4*. If a *velocity* argument is given, the value of the angle will change at each time step of the simulation at a rate specified by the *velocity* argument. The *velocity* must be given in [degree/(a.u. time)].

constraint list

Print a list of all currently defined constraints.

constraint delete constraint name

Remove the constraint *constraint\_name* from the constraint set.

constraint enforce

Modify atomic positions so as to enforce all constraints using the SHAKE algorithm. constraint set *constraint\_name value [velocity]* 

Modify the value of a constraint, and optionally its velocity. This applies to the distance, angle and torsion constraints only, for which the value is distance, angle and angle, respectively.

#### RELATED INFORMATION

list atoms, angle, distance, torsion

### distance

#### NAME

distance [-pbc] --print the distance between two atoms

### **SYNOPSIS**

distance atom1 atom2

#### DESCRIPTION

The distance command prints the value of the distance separating two atoms. If the -pbc option is used, the positions of the atoms are interpreted as those of the nearest atom replica, taking into account periodic boundary conditions.

#### RELATED INFORMATION

angle, torsion

# extforce

#### NAME

extforce -add, modify or delete external forces on atoms

### **SYNOPSIS**

```
extforce define atomic extforce_name atom fx fy fz extforce define pair extforce_name atom1 atom2 f extforce define global extforce_name fx fy fz extforce set extforce_name fx fy fz extforce set extforce_name f extforce list extforce delete extforce_name
```

### **DESCRIPTION**

The extforce command is used to define, modify or delete external forces acting on specific atoms. The "define", "set", "list" and "delete" subcommands modiy the set of external forces as detailed below for each choice of parameters.

```
extforce define atomic extforce name atom fx fy fz
```

This command defines an external force named extforce\_name acting on atom atom. The force has components fx,fy,fz. The force components must be given in atomic units (Hartree/Bohr).

```
extforce define pair extforce name atom1 atom2 f
```

This command defines a pair force named extforce\_name acting only on the atoms atom1 and atom2. The magnitude of the pair force is f and must be given in atomic units (Hartree/Bohr). A positive value of f defines an attractive force between atom1 and atom2.

extforce define global extforce name fx fy fz

This command defines a global external force named extforce\_name acting on all atoms. The force has components fx,fy,fz. The force components must be given in atomic units (Hartree/Bohr).

extforce set extforce name fx fy fz

This command modifies the components of the force associated with the external force extforce name. This syntax applies to the "atomic" and "global" forces only.

extforce set *extforce* name f

This command modifies the magnitude of the force associated with the external force extforce name. This syntax applies to the "pair" force only.

extforce list

This command prints a list of all external forces.

extforce delete extforce name

This command removes the external force extforce\_name from the set of external forces.

#### RELATED INFORMATION

# fold\_in\_ws

**NAME** 

fold in ws -fold all atoms within the Wigner-Seitz cell

**SYNOPSIS** 

fold in ws

#### DESCRIPTION

The fold\_in\_ws command moves atoms by multiples of the unit cell lattice vectors so that all atoms are within the Wigner-Seitz cell.

### RELATED INFORMATION

# help

**NAME** 

help --print a brief help message about a command SYNOPSIS

1 1 5

help [command]

**DESCRIPTION** 

The help command prints a short description of the *command* given as an argument. When used without arguments, prints a list of valid commands.

RELATED INFORMATION

# **kpoint**

### NAME

kpoint –define and manage the set of k-points used in the calculation of the electronic structure.

#### **SYNOPSIS**

kpoint list kpoint add kx ky kz weight kpoint delete kx ky kz

# **DESCRIPTION**

The kpoint command is used to add and delete k-points to the set of k-points used in the electronic structure calculation. The kpoint is defined by a vector on the basis of the reciprocal lattice vectors. If reciprocal lattice vectors are  $\mathbf{b_1}$   $\mathbf{b_2}$  and  $\mathbf{b_3}$  the k-point defined by the numbers (kx, ky, kz) on the command line is  $kx * \mathbf{b_1} + ky * \mathbf{b_2} + kz * \mathbf{b_3}$ . For example, the X point of the Brillouin Zone for an FCC lattice is specified as kx=0.5, ky=0.5, kz=0.0.

The list of all currently defined k-points can be printed using the command kpoint list. The sum of the *weight* arguments must add up to 1.0. This is currently not checked by Qbox. Some k-points can be defined with zero weight. In that case, the electronic wavefunctions and eigenvalues are computed at these points, but they are not included in the calculation of the charge density.

By default, Qbox starts with a k-point set containing a single k-point: k=(0,0,0) (the  $\Gamma$  point) with a weight of 1.0. When defining a k-point set, it is necessary to first *delete* the  $\Gamma$  point before defining other k-points. This is due to two possible reasons:

- 1) The desired k-point set does not contain the  $\Gamma$  point.
- 2) The desired k-point set contains the  $\Gamma$  point, but with a weight different from 1.0. In this case, the  $\Gamma$  point must be deleted and then added in order to be defined with the correct weight. The only way to change the weight of a k-point is to delete it and define it again.

Qbox does not perform any symmetrization of the charge density to reduce the number of k-points to the irreducible wedge of the Brillouin Zone. The full set of k-points must be defined (except for the k, -k symmetry, i.e. If k is included in the set, then -k need not be included).

Modifying the k-point set erases all wavefunctions. It is not possible to modify the k-point set after running a calculation or after loading a sample without resetting the wavefunctions.

Note: When deleting a kpoint, the arguments kx, ky, kz are compared to the coordinates of the kpoints currently defined. Since comparisons of floating point numbers are unreliable, the kpoint delete command will delete any k-point located within a radius of  $10^{-6}$  of the vector (kx, ky, kz). Similarly, when adding a new k-point, the kpoint add command will exit without defining the new k-point and print a warning message if a previously defined k-point is located within a radius of  $10^{-6}$  of the new kpoint.

# **EXAMPLE**

Define a set of 8 k-points for a simple cubic or orthorhombic cell. This k-point set is equivalent to doubling the cell in all three directions

```
kpoint delete 0 0 0
# Gamma point
kpoint add 0.0 0.0 0.0 0.125
# X point
kpoint add 0.5 0.0 0.0 0.125
```

```
kpoint add 0.0 0.5 0.0 0.125
kpoint add 0.0 0.0 0.5 0.125
# R point
kpoint add 0.5 0.5 0.5 0.125
# M point
kpoint add 0.5 0.5 0.0 0.125
kpoint add 0.5 0.0 0.5 0.125
kpoint add 0.5 0.0 0.5 0.125
```

Define the *X* and *L* points in the Brillouin Zone of a FCC lattice, with zero weight:

```
# a1 = (a/2, a/2, 0)
# a2 = (0, a/2, a/2)
# a3 = (a/2, 0, a/2)
# b1 = (2*pi/a) ( 1.0, 1.0, -1.0)
# b2 = (2*pi/a) (-1.0, 1.0, 1.0)
# b3 = (2*pi/a) ( 1.0, -1.0, 1.0)
# X point
# X = (2*pi/a) ( 1.0, 0.0, 0.0 ) = 0.5 * ( b1 + b3 )
kpoint 0.5 0.0 0.5 0.0000
# L point
# L = (2*pi/a) ( 0.5, 0.5, 0.5 ) = 0.5 * ( b1 + b2 + b3 )
kpoint add 0.5 0.5 0.5 0.0000
```

# **RELATED INFORMATION**

# list\_atoms

**NAME** 

list atoms --print a list of atoms currently defined in the sample

# **SYNOPSIS**

list atoms

### DESCRIPTION

The list\_atoms command prints a list of all atoms currently defined.

### **RELATED INFORMATION**

list\_species, atom

# list\_species

**NAME** 

list species --print a list of all species currently defined

#### **SYNOPSIS**

list species

#### DESCRIPTION

The list\_species command prints a list of all species currently defined. For each species, the parameters of the corresponding pseudopotential are printed.

### RELATED INFORMATION

list atoms, species

#### load

NAME

load --load a sample from an XML sample document

### **SYNOPSIS**

load URI

#### DESCRIPTION

The load command loads a simulation sample defined in an XML document provided by the *URI* argument. The *URI* can be a local file, in which case Qbox will open and read the file. If *URI* is a *URL* (e.g. <a href="http://www.quantum-simulation.org/examples/samples/ch4.xml">http://www.quantum-simulation.org/examples/samples/ch4.xml</a>) Qbox will download the document from the corresponding web site. Note that loading a URL remotely only works if the nodes on which Qbox is running have web access. This may not be the case on some parallel computers in which compute nodes do not have web access.

Qbox sample documents must conform to the XML Schema definition provided at <a href="http://www.quantum-simulation.org">http://www.quantum-simulation.org</a>

#### RELATED INFORMATION

save

#### move

NAME

move --change the position of an atom

# **SYNOPSIS**

move *atom* to *x y z* move *atom* by *dx dy dz* 

# DESCRIPTION

The move command moves an atom to an absolute position specified by x, y, z or by a relative displacement specified by dx, dy, dz. Positions or displacements must be given in atomic units (Bohr).

### RELATED INFORMATION

# plot

#### NAME

plot –generate a plot file with atoms and/or charge density, orbitals or local potential

#### **SYNOPSIS**

```
plot filename
plot -density filename
plot -wf n filename
plot -wfs nmin nmax [-spin {1|2}] filename
plot -vlocal [-spin {1|2}] filename
```

### DESCRIPTION

The plot command creates a plot file to be viewed with another rendering program such as VMD or XcrysDen. The type of output file generated depends on the arguments given to the plot command.

- plot filename
  - This command generates an xyz file containing atomic positions only.
- plot -density [-spin {1|2}] *filename*This command generates a file in *cube* format containing the atomic positions and the total charge density. If the -spin option is used, the density of the first (1) or second (2) spin is used.
- plot -wf n [-spin {1|2}] *filename*This command generates a file in *cube* format containing the *n*-th wave function. If the -spin option is used, the wave function of the first (1) or second (2) spin is used.
- plot -wfs nmin nmax [-spin {1|2}] filename
  Generate a file in cube format containing the sum of the squares of the amplitudes of the wavefunctions nmin to nmax inclusive. If the -spin option is used, the wave functions of the first (1) or second (2) spin are used.
- plot -vlocal [-spin {1|2}] filename
   This command generates a file in *cube* format containing the local potential
   (Vlocal+VHartree+Vxc). If the -spin option is used, the potential for the first (1) or second
   (2) spin is used.

The plot command is currently only working for wave functions at the  $\Gamma$  point.

#### RELATED INFORMATION

# partial\_charge

### **NAME**

partial\_charge -compute the amount of electronic charge in a sphere centered on an atom

#### **SYNOPSIS**

```
partial charge [-spin {1|2}] name radius
```

### DESCRIPTION

The partial charge command computes the amount of electronic charge contained in a sphere of

radius *radius* centered on atom *name*. The *radius* value must be specified in atomic units (Bohr). When using the -spin option, the charge density of the given spin is computed. If nspin=2 and the -spin option is not used, the total charge is computed.

### RELATED INFORMATION

# print

**NAME** 

print --print the current value of a Qbox variable

#### **SYNOPSIS**

print variable

# **DESCRIPTION**

The print command prints the current value a Qbox variable. For a list of variables, see the section "Qbox variables".

# **RELATED INFORMATION**

<u>set</u>

# quit

**NAME** 

quit --exit Qbox

### **SYNOPSIS**

quit

### **DESCRIPTION**

The quit command exits Qbox without saving any information about the sample. To save the current sample, see the <u>save</u> command.

# **RELATED INFORMATION**

<u>save</u>

# randomize\_r

NAME

randomize r --add a random perturbation to atomic positions

### **SYNOPSIS**

#### DESCRIPTION

The randomize\_r command adds random numbers to the coordinates of the atomic positions. The random displacements are drawn from a normal distribution scaled by the *amplitude* argument.

#### RELATED INFORMATION

randomize v

# randomize\_v

### **NAME**

randomize\_v –initialize atomic velocities with random values from a Maxwell Boltzmann distribution

#### **SYNOPSIS**

randomize v temperature

# **DESCRIPTION**

The randomize\_v command initializes atomic velocities with random numbers drawn from a Maxwell-Boltzmann distribution. The *temperature* argument determines the temperature of the distribution.

### RELATED INFORMATION

randomize r

# randomize\_wf

#### **NAME**

randomize\_wf --add a random perturbation to electronic wavefunctions

#### **SYNOPSIS**

randomize wf [amplitude]

### DESCRIPTION

The randomize\_wf command adds random numbers to the Fourier coefficients of the electronic wave function. The *amplitude* argument can be used to change the intensity of the perturbation. The randomize\_wf command is used at the beginning of an electronic structure calculation when the symmetry of the atomic coordinates is high. In such situations, the iterative algorithms used to compute the electronic ground state can converge to saddle points of the energy functional instead of true minima. Using randomize\_wf introduces a slight symmetry breaking which is sufficient to avoid high-symmetry saddle points.

# RELATED INFORMATION

# rescale\_v

### NAME

rescale v –rescale atomic velocities

### **SYNOPSIS**

rescale\_v scaling factor

### **DESCRIPTION**

The rescale\_v command rescales the velocity of all atoms by a common factor defined by the *scaling factor* argument.

### RELATED INFORMATION

# reset vcm

**NAME** 

reset vcm --reset the velocity of the center of mass to zero

### **SYNOPSIS**

reset vcm

### **DESCRIPTION**

The reset\_vcm command modifies the velocity of all atoms so as to ensure that the velocity of the center of mass is zero. The current value of the velocity of the center of mass is printed by the <u>status</u> command

### RELATED INFORMATION

### rseed

### **NAME**

rseed –initialize the random number generator

### **SYNOPSIS**

rseed [seed]

### **DESCRIPTION**

The rseed command initializes the random number generator. Qbox uses random numbers in the implementation of the <u>randomize\_wf</u> command and in stochastic thermostats (LOWE, ANDERSON, BDP). If a seed argument is provided, it is used to initialize the random number generator. If no seed is provided the time function is used to generate a seed. Note: when running

multiple instances of Qbox in client-server mode, and if the rseed command is not used, all instances may be initialized with the same seed if they are started at the same time. This problem can be avoided by using the rseed command for each instance with a different argument, or by starting all instances in a staggered way using a delay of a few seconds.

### run

#### **NAME**

run --update electronic wavefunctions and/or atomic positions and/or unit cell

### **SYNOPSIS**

```
run [-atomic_density] niter
run [-atomic_density]niter nitscf
run [-atomic_density]niter nitscf nite
```

#### DESCRIPTION

The run command starts a simulation in which atomic positions and/or electronic states are updated. The algorithms used to update the atomic positions and electronic states are determined by the variables atoms dyn and wf dyn. The parameters are defined as follows

- -atomic\_density The first self-consistent iteration is started using a charge density made of a superposition of atomic charge densities.
- *niter* The number of ionic steps to be performed, i.e. steps during which atomic positions are updated. This number can be zero if only electronic wavefunction updates are desired.
- *nitscf* The maximum number of self-consistent electronic iterations. The charge density is updated at the beginning of each self-consistent iteration. Iterations may be skipped if the energy has reached convergence within the <u>scf\_tol</u> tolerance criterion.
- *nite* The number of electronic iterations performed between updates of the charge density The run command can be used in the following ways
  - run niter
    - Perform *niter* ionic steps. Before each ionic step, the electronic states are updated once, and the charge density is updated (i.e. both parameters *nitscf* and *nite* default to 1). Using "run 0" computes the total energy without modifying the electronic wavefunction.
  - run niter nitscf
    - Perform *niter* ionic steps. Before each ionic step, the charge density is updated *nitscf* times. Before each update of the charge density, the electronic states are updated once (i.e. the parameter *nite* defaults to 1.
  - run niter nitscf nite
    - Perform *niter* ionic steps. Before each ionic step, the charge density is updated *nitscf* times. Before each charge density update, the electronic states are updated *nite* times.

# Example 1

run 0 100

is used to perform 100 self-consistent electronic optimization steps. Only electronic optimization is performed. This the most common way of computing the electronic ground state.

### Example 2

run 50 10

is used to perform 50 ionic steps, with 10 charge density updates before each ionic step. Before each charge density update, the electronic states are updated once.

# Example 3

run 50 5 10

is used to performed 50 ionic steps. The charge density is updated 5 times before each ionic step. The electronic states are updated 10 times before each charge density update.

If the variable <u>cell\_dyn</u> is not set to LOCKED, the unit cell parameters are updated each time the atomic positions are updated. In that case, the <u>stress</u> variable must be set to ON so that the stress tensor is computed before the cell parameters are updated.

Atomic positions are updated before the charge density and electronic states are updated. As a consequence, the atomic positions and the electronic structure are consistent at the end of a run command.

#### RELATED INFORMATION

atoms dyn, wf dyn, scf tol

#### save

NAME

save --save the current sample into a file

#### **SYNOPSIS**

save [-serial] [-text] [-atomsonly] [-no wfv] filename

# **DESCRIPTION**

The save command saves the current sample into a file in XML format. The format used conforms to the XML Schema defined at <a href="http://www.quantum-simulation.org">http://www.quantum-simulation.org</a>. The information saved includes the dimensions of the unit cell, atomic positions and velocities, the electronic wave function, and optionally the time derivative of the wave function. If the -serial option is used, the data is saved from task 0 only and no attempt is made to use optimized I/O functionality. If the -text option is used, the wave function coefficients are written in formatted text form rather than encoded in base64 format in the sample file. If the -atomsonly option is used, only the <atomset>"element is written.">http://www.quantum-simulation.org</a>.

### RELATED INFORMATION

load

#### set

**NAME** 

set --assign a value to a Qbox variable

### **SYNOPSIS**

set variable value [value ...]

#### DESCRIPTION

The set command assigns the value(s) *value* to the variable *variable*. Some variables (e.g. <u>cell</u>) are multivalued, in which case the set command requires multiple arguments.

#### RELATED INFORMATION

print

# set\_velocity

**NAME** 

set velocity --set the velocity of an atom

#### **SYNOPSIS**

set velocity atom name  $\{vx | *|-\} \{vy | *|-\} \{vz | *|-\}$ 

#### DESCRIPTION

The set\_velocity command sets the velocity of an atom to vx,vy,vz. If one or more of the arguments is '\*', the corresponding component of the velocity is unchanged. If an argument is '-', the corresponding component of the velocity changes sign.

The set command assigns the value(s) *value* to the variable *variable*. Some variables (e.g. <u>cell</u>) are multivalued, in which case the set command requires multiple arguments.

**Example:** set\_velocity C18 0.01 \* - will set the component vx to 0.01, leave the component vy unchanged and change the sign of the component vz.

#### RELATED INFORMATION

move, list atoms

# species

**NAME** 

species --define a new atomic species and add it to the list of currently known species

### **SYNOPSIS**

species name URI

### DESCRIPTION

The species command defines a new atomic species under the name *name*. The newly defined species is added to the list of currently known species. The definition is read from the given *URI*. If the *URI* is a local file, Qbox opens and reads it. If the *URI* is a *URL* (e.g.

<a href="http://www.quantum-simulation.org/examples/species/hydrogen\_pbe.xml">http://www.quantum-simulation.org/examples/species/hydrogen\_pbe.xml</a>) Qbox downloads the species definition from the corresponding web site. The species definition document must conform to the species XML Schema definition given at <a href="http://www.quantum-simulation.org">http://www.quantum-simulation.org</a>

If the species *name* is already defined, the definition is replaced with that of the *URI* argument.

### RELATED INFORMATION

# list species, atom

# status

**NAME** 

status --print status of the current sample

**SYNOPSIS** 

status

# **DESCRIPTION**

The status command prints a brief summary of the characteristics of the current sample.

### RELATED INFORMATION

### strain

**NAME** 

strain –apply strain on the system

# **SYNOPSIS**

strain [-atomsonly] [-inverse] uxx uyy uzz uxy uyz uxz

# **DESCRIPTION**

The strain command modifies the shape of the unit cell and modifies the atomic positions to impose a strain defined by the components of the symmetric strain tensor u. Using the -inverse flag causes the inverse transformation to be applied. Using -atomsonly changes the postitions of atoms without affecting the unit cell

# **RELATED INFORMATION**

# torsion

**NAME** 

torsion --print the value of the torsion angle (dihedral) defined by the positions of four atoms

### **SYNOPSIS**

torsion [-pbc] atom1 atom2 atom3 atom4

### **DESCRIPTION**

The torsion command prints the value of the angle (dihedral) defined by the four atoms given as arguments. The names *atom1*, *atom2* atom3 and *atom4* must refer to the names of atoms currently defined in the sample. If the -pbc option is used, the positions of the atoms are interpreted as those of the nearest atom replica, taking into account periodic boundary conditions.

# RELATED INFORMATION

list atoms, angle, distance

# ! (shell escape)

**NAME** 

! (shell escape) --execute a Unix command from withing Qbox

# **SYNOPSIS**

! command [arguments]

# **DESCRIPTION**

The! command executes the command given as an argument in a Unix shell.

**RELATED INFORMATION** 

# 6 Qbox variables

# alpha\_PBE0

**NAME** 

alpha\_PBE0 -coefficient of Hartree-Fock exchange in the PBE0 xc functional

### DESCRIPTION

The alpha\_PBE0 variable defines the coefficient multiplying the Hartree-Fock exchange energy in the PBE0 exchange-correlation functional. The default value is 0.25.

#### **ALLOWED VALUES**

non-negative real numbers

#### RELATED INFORMATION

<u>xc</u>

# atoms\_dyn

**NAME** 

atoms dyn --atom dynamics control variable

### DESCRIPTION

The atoms\_dyn variable determines the algorithm used to update atomic positions during a simulation. The following values are allowed:

- LOCKED: Ionic forces are not computed and atomic positions are not updated. This is the default.
- SD: Steepest Descent. Ionic forces are computed and atomic positions are updated using the steepest descent algorithm

$$\vec{R}(t+dt) = \vec{R}(t) + \frac{dt^2}{m}\vec{F}(t)$$

where F is the force, m is the ionic mass and dt is the time step (see variable  $\underline{dt}$ ).

- SDA: Steepest Descent with Acceleration. A line minimization algorithm is used to find a
  minimum satisfying the Wolfe conditions before changing the descent direction. Using this
  algorithm requires a full calculation of the electronic ground state at each ionic step, i.e. nitscf>
  1 in the run command.
- CG: Conjugate Gradient. A line minimization algorithm is used to find a minimum satisfying the Wolfe conditions before changing the descent direction. The Polak-Ribiere formula is used to update descent directions. Using this algorithm requires a full calculation of the electronic ground state at each ionic step, i.e. *nitscf* > 1 in the run command. Using this algorithm requires a full calculation of the electronic ground state at each ionic step, i.e. *nitscf* > 1 in the run command.
- MD: Molecular dynamics. Ionic forces are computed and ionic positions are updated using the Verlet algorithm

$$\vec{R}(t+dt) = 2\vec{R}(t) - \vec{R}(t-dt) + \frac{dt^2}{m}\vec{F}(t)$$

This algorithm can be used to perform either Born-Oppenheimer molecular dynamics (using *nitscf* > 1 in the run command) or Car-Parrinello molecular dynamics (*nitscf*=1). See also the wf\_dyn variable.

• BMD: Blocked Molecular dynamics. This algorithm updates ionic positions according to the MD algorithm (Verlet), but velocities are reset to zero every time the total energy increases. This algorithm is used to optimize geometry. It requires a full calculation of the electronic ground state at each ionic step, i.e. *nitscf* > 1 in the run command.

### ALLOWED VALUES

LOCKED, SD, SDA, CG, MD, BMD

#### RELATED INFORMATION

dt

### **bIHF**

**NAME** 

blHF -bisection levels for Hartree-Fock exchange computation

### DESCRIPTION

The blHF variable consists of three integer values  $lx\ ly\ lz$  that specify the number of recursive levels of bisection that must be performed on wave functions in the x, y, and z directions when computing the Hartree-Fock exchange energy. The values of blHF are only used if the variable btHF is non-zero.

#### **ALLOWED VALUES**

positive integers

### RELATED INFORMATION

btHF, bisection

#### **btHF**

**NAME** 

btHF -bisection threshold for Hartree-Fock exchange computation

# **DESCRIPTION**

The btHF variable defines the threshold used in the recursive bisection performed on wave functions when computing the Hartree-Fock exchange energy. If btHF is zero, no bisection is performed during Hartree-Fock exchange calculations. The default value of btHF is zero.

#### **ALLOWED VALUES**

floating point numbers in [0,1]

# RELATED INFORMATION

blHF, bisection

#### cell

**NAME** 

cell --unit cell parameters

# **DESCRIPTION**

The cell variable contains the coordinates of the three lattice vectors  $\vec{a}_1, \vec{a}_2, \vec{a}_3$  defining the unit cell. The unit cell is defined using the <u>set</u> command with the following arguments

set cell 
$$a_{1x} a_{1y} a_{1z} a_{2x} a_{2y} a_{2z} a_{3x} a_{3y} a_{3z}$$

The lattice vectors  $\vec{a}_1, \vec{a}_2, \vec{a}_3$  form the columns of the 3x3 lattice parameter matrix A

$$A = \begin{pmatrix} a_{11} & a_{12} & a_{13} \\ a_{21} & a_{22} & a_{32} \\ a_{31} & a_{32} & a_{33} \end{pmatrix}$$

The default value of all cell parameters is zero.

### **ALLOWED VALUES**

positive real numbers

#### RELATED INFORMATION

ref cell

# cell\_dyn

NAME

cell dyn --cell dynamics control variable

### **DESCRIPTION**

The cell\_dyn variable determines the algorithm used to update the unit cell during a simulation. The following values are allowed:

- LOCKED: The unit cell is not updated. This is the default.
- SD: Steepest Descent. The unit cell is updated using the computed stress tensor and the steepest descent algorithm

$$a_{ij}(t+dt) = a_{ij}(t) + \frac{dt^2}{m_{\Omega}} \Omega \left[ \sigma(t) - \sigma^{ext}(t) \right] A(t)$$

where  $\sigma(t)$  is the stress tensor,  $\sigma^{ext}(t)$  is the externally applied stress (variable <u>ext\_stress</u>),  $\Omega$  is the volume of the unit cell,  $m_{\Omega}$  is the mass of the unit cell (variable <u>cell\_mass</u>), A is the 3x3 lattice parameter matrix (see <u>cell</u> variable) and dt is the time step (variable <u>dt</u>).

• CG: Conjugate Gradient. The unit cell and the atomic positions are updated using the computed stress tensor and the atomic forces using a conjugate gradient algorithm. Note that both the unit cell and atomic positions are optmized simultaneously independently of the value of

### atoms dyn.

# ALLOWED VALUES LOCKED, SD, CG

### RELATED INFORMATION

cell, ref\_cell, cell\_lock

# cell lock

NAME

cell lock --cell dynamics constraints control variable

#### DESCRIPTION

The cell\_lock variable is used to restrict the possible changes of the unit cell parameters. The allowed values are

- OFF: No restriction on the unit cell parameters are enforced. This is the default.
- A: The lattice vector  $\vec{a}_1$  is fixed.
- B: The lattice vector  $\vec{a}_2$  is fixed.
- C: The lattice vector  $\vec{a}_3$  is fixed.
- AB: The lattice vectors  $\vec{a}_1$  and  $\vec{a}_2$  are fixed.
- AC: The lattice vectors  $\vec{a}_1$  and  $\vec{a}_3$  are fixed.
- BC: The lattice vectors  $\vec{a_2}$  and  $\vec{a_3}$  are fixed.
- ABC: All lattice vectors are fixed. This is equivalent to <u>cell dyn</u> = LOCKED.
- S: The shape of the unit cell is preserved. Lattice vectors can change length but not direction.
- AS: The lattice vector  $\vec{a}_1$  is fixed and the shape of the unit cell is preserved.
- BS: The lattice vector  $\vec{a}_2$  is fixed and the shape of the unit cell is preserved.
- CS: The lattice vector  $\vec{a}_3$  is fixed and the shape of the unit cell is preserved.
- ABS: The lattice vectors  $\vec{a}_1$  and  $\vec{a}_2$  are fixed and the shape of the unit cell is preserved.
- ACS: The lattice vectors  $\vec{a}_1$  and  $\vec{a}_3$  are fixed and the shape of the unit cell is preserved.
- BCS: The lattice vectors  $\vec{a}_2$  and  $\vec{a}_3$  are fixed and the shape of the unit cell is preserved.
- R: The aspect ratio of the unit cell is preserved. All lattice vectors are rescaled by the same constant.

# **ALLOWED VALUES**

OFF, A, B, C, AB, AC, BC, ABC, S, AS, BS, CS, ABS, ACS, BCS, R

### RELATED INFORMATION

cell, cell\_dyn, cell\_mass

# cell\_mass

NAME

cell mass -- mass of the unit cell

#### DESCRIPTION

The cell\_mass variable is the mass of the unit cell in atomic units (in which the mass of a carbon atom is 12.0). The default value is 10000.

#### ALLOWED VALUES

positive real values

#### RELATED INFORMATION

cell, cell dyn

# charge mix coeff

**NAME** 

charge mix coeff --charge density mixing coefficient

### DESCRIPTION

The charge density mixing coefficient is used to update the electronic charge density during self-consistent iterations (see <a href="mailto:charge\_mix\_ndim">charge\_mix\_ndim</a>). The default value is 0.5. Smaller values can be used to accelerate the convergence of self-consistent iterations in metallic systems.

#### ALLOWED VALUES

real values in the interval [0,1]

### RELATED INFORMATION

charge mix rcut, charge mix ndim

# charge\_mix\_ndim

**NAME** 

charge mix ndim –dimension of Anderson acceleration of charge mixing

#### DESCRIPTION

The charge\_mix\_ndim parameter n determines how many previous charge density corrections are used in the Anderson acceleration of the charge mixing scheme. The new input charge density is computed according to

$$\rho_{\rm in}^{(k+1)} = \overline{\rho} + \alpha \overline{f}$$

where  $\alpha$  is the charge mixing coefficient (<u>charge mix coeff</u>) and  $\overline{f}$  is the least-squares residual in the subspace spanned by the *n* previous charge density corrections

$$\bar{f} = \delta \rho^{(k)} + \sum_{i=1}^{n} \theta_i (\delta \rho^{(k-i)} - \delta \rho^{(k)})$$

where  $\delta \rho^{(k)} = \rho_{out}^{(k)} - \rho_{in}^{(k)}$  is the difference between the output and input charge density of the self-

consistent iteration k, and

$$\overline{\rho} = \rho_{\mathrm{in}}^{(k)} + \sum_{i=1}^{n} \theta_{i} (\rho_{\mathrm{in}}^{(k-i)} - \rho_{\mathrm{in}}^{(k)})$$

The coefficients  $\theta_i$  are chosen so as to minimize  $\|\overline{f}\|_2^2$  in a row-weighted least squares sense. The weight used in the LS calculation is the Kerker screening function

$$w(\mathbf{G}) = \frac{\mathbf{G}^2 + q_0^2}{\mathbf{G}^2}$$

where  $q_0 = 2\pi/r_c$  and  $r_c$  is the charge mixing cutoff radius (<u>charge\_mix\_rcut</u>). The default value of n is 3.

# SPECIAL VALUES

Using n=0 leads to simple mixing:  $\rho_{in}^{(k+1)} = \rho_{in}^{(k)} + \alpha \left(\rho_{out}^{(k)} - \rho_{in}^{(k)}\right)$ 

# **ALLOWED VALUES**

non-negative integers

#### RELATED INFORMATION

charge mix rcut, charge mix coeff

# charge\_mix\_rcut

**NAME** 

charge mix rcut --charge mixing cutoff radius

### **DESCRIPTION**

The charge\_mix\_rcut variable is used to define the range of the Coulomb interaction in the Kerker screening function used in the charge density mixing scheme (see <a href="charge mix">charge mix</a> ndim).

The default value of charge mix rcut is 10 a.u.

If charge mix rcut is set to zero, no screening is used.

### **ALLOWED VALUES**

positive real numbers

# RELATED INFORMATION

<u>charge mix ndim</u>, <u>charge mix ndim</u>

# debug

NAME

debug --debug parameters

**DESCRIPTION** 

The debug variable is used to pass debug parameters to Qbox. It is not intended for normal use.

### **ALLOWED VALUES**

character strings

### RELATED INFORMATION

### dt

**NAME** 

dt --simulation time step

# **DESCRIPTION**

The dt variable is the simulation time step in atomic units of time (1 a.u. of time = 0.02418885 fs). The default value is 3 a.u.

### **ALLOWED VALUES**

non-negative real numbers

#### RELATED INFORMATION

atoms dyn, wf dyn, cell dyn

# ecut

NAME

ecut --plane-wave basis energy cutoff

# **DESCRIPTION**

The ecut variable defines the size of the plane wave basis used to define the electronic wavefunctions. It must given in Rydberg units. The wavefunction plane wave basis consists of all plane waves having a kinetic energy smaller than  $E_{cut}$ . The charge density and the total potential are described using a larger basis set that includes all plane waves with a kinetic energy smaller than  $4E_{cut}$ . The default value of ecut is zero, in which case the plane wave basis contains one basis function—the plane wave of wavevector  $\mathbf{G} = 0$ .

#### **ALLOWED VALUES**

non-negative real numbers

### RELATED INFORMATION

# ecutprec

NAME

### **DESCRIPTION**

The ecutprec variable defines the energy cutoff used in the preconditioner for electronic structure optimization. Corrections to the electronic wavefunctions are preconditioned in Fourier space using a diagonal preconditioning matrix K whose elements are defined by

$$k(G) = \begin{cases} k_{ij} = \delta_{ij} k(G) \\ \frac{1}{2 E_{cut}^{prec}} & \frac{1}{2} G^2 < E_{cut}^{prec} \\ \frac{1}{2} E & \frac{1}{2} E \ge E_{cut}^{prec} \end{cases}$$

The value of ecutprec must be given in Rydberg units. Preconditioning is only used if the <u>wf\_dyn</u> variable is set to either PSD PSDA or JD. If ecutprec = 0, an automatic preconditioner is used. The default value of ecutprec is zero (automatic preconditioning).

### **ALLOWED VALUES**

non-negative real numbers smaller than or equal to ecut.

### RELATED INFORMATION

ecut, wf dyn

### ecuts

**NAME** 

ecuts --energy cutoff for stress confinement potential

### **DESCRIPTION**

The ecuts variable defines the energy cutoff used in a confinement potential in Fourier space. The confinement potential is used when computing the stress tensor with variable cell size in order to ensure constant resolution as the unit cell changes size. The confinement energy is

$$E_{conf} = \frac{1}{2} \sum_{n,G} |c_{nG}|^2 G^2 f(G)$$

where

where
$$f(g) = f_s \left( 1 - \frac{1}{1 + \exp((G^2/2 - E_{cut}^s)/\sigma_s)} \right)$$

$$f_s = 2 \text{ and } \sigma_s = \frac{1}{2}.$$

The default value of ecuts is zero, in which case the confinement potential is not used. For a detailed description of the use of confinement potentials in constant pressure simulations, see 1. P. Focher, G. L. Chiarotti, M. Bernasconi, *et al.* Structural Phase-Transformations Via 1St-Principles Simulation, Europhys. Lett. **26** (5): 345-351 (1994).

2. M. Bernasconi, G. L. Chiarotti, P. Focher, *et al.* First-Principle Constant-Pressure Molecular-Dynamics, Journal Of Physics And Chemistry Of Solids **56** (3-4): 501-505 (1995).

### **ALLOWED VALUES**

positive real numbers smaller than or equal to ecut

### RELATED INFORMATION

stress, cell\_dyn

# e field

**NAME** 

e field --applied electric field (displacement field)

### DESCRIPTION

The e\_field variable consists of three numbers that define the applied electric field (displacement field, or induction)  $\vec{D} = (D_x, D_y, D_z)$ . The field must be given in atomic units (1 Hartree/(electron Bohr) =  $5.1422 \times 10^{11}$  V/m. The polarization variable must be set to define the algorithm used to compute the electric dipole and quadrupole in the presence of the applied field. The default value is zero.

#### ALLOWED VALUES

real numbers

#### RELATED INFORMATION

polarization

### emass

NAME

emass --fictitious electronic mass for Car-Parrinello simulations

### **DESCRIPTION**

The emass variable defines the fictitious electronic mass used in Car-Parrinello simulations. The default value is zero, in which case the fictitious electronic mass used in the calculation is  $m_e = 2 E_{cut} dt^2$ . The value of emass is only relevant if the variable wf\_dyn is set to MD (i.e. if the wave function dynamics is Car-Parrinello). It is ignored otherwise.

### **ALLOWED VALUES**

positive real values

#### RELATED INFORMATION

wf dyn

# ext\_stress

NAME

ext stress --external stress

### **DESCRIPTION**

The ext\_stress variable determines the value of the externally applied stress. The ext\_stress variable must be set using the following syntax

set ext stress  $\sigma_{xx} \sigma_{yy} \sigma_{zz} \sigma_{xy} \sigma_{yz} \sigma_{xz}$ 

where the values of the elements of the stress tensor must be given in GPa units. The external stress can be positive or negative. The default value is zero.

### ALLOWED VALUES

real numbers

#### RELATED INFORMATION

stress

# fermi\_temp

**NAME** 

fermi temp -- Fermi temperature for fractionally occupied states

# **DESCRIPTION**

The fermi\_temp variable determines the value of the Fermi temperature used in the calculation of occupation factors for fractionally occupied states. The value must be given in Kelvin. The default value is zero.

#### **ALLOWED VALUES**

non-negative real numbers

### RELATED INFORMATION

nempty

# iter\_cmd

**NAME** 

iter cmd --command or script to be executed every iter cmd period ionic steps

### DESCRIPTION

The iter\_cmd variable defines a command or script that is executed every iter\_cmd\_period ionic steps. If a single command must be executed, the full command (with arguments) can be used as the value of iter\_cmd. If multiple commands are used, they must be written in a local script file, and the name of the script is used as the value of iter\_cmd.

### **ALLOWED VALUES**

string

### RELATED INFORMATION

iter cmd period

# iter\_cmd\_period

**NAME** 

iter cmd period --number of ionic steps between executions of iter cmd

#### DESCRIPTION

The iter\_cmd\_period variable defines the number of ionic steps that separate successive executions of the command (or script) defined by iter cmd. The default value is 1.

### **ALLOWED VALUES**

positive integers

### RELATED INFORMATION

iter cmd

# nempty

**NAME** 

nempty --number of empty electronic states

### DESCRIPTION

The nempty variable determines the number of electronic states that are included in the calculation in addition to the number of states needed to accommodate the total number of electrons. If nempty is non-zero, the eigenvalues and eigenvectors of the Kohn-Sham hamiltonian are computed at each electronic iteration and the charge density is recomputed from the eigenvectors using a Fermi distribution. The default value is of nempty is zero.

#### **ALLOWED VALUES**

non-negative integers

#### RELATED INFORMATION

fermi\_temp

# net\_charge

**NAME** 

net\_charge --net charge of the system

### DESCRIPTION

The net\_charge variable is used to control the total amount of electronic charge in the calculation. If net\_charge = 0, the total number of electrons is determined from the sum of the valence charges given in the species definition files and the system is neutral. If net\_charge = -1, an extra electron is added to the system. If net\_charge = 1, an electron is removed from the system. The default value is zero.

# **ALLOWED VALUES**

integers

#### RELATED INFORMATION

### nrowmax

NAME

nrowmax --maximum number of process grid rows

### **DESCRIPTION**

The nrowmax variable determines the shape of the process grid used in parallel calculations. It is used to optimize performance when large numbers of parallel tasks are used. The default value is 32. Setting the value of nrowmax erases all information about the current wave functions.

# **ALLOWED VALUES**

positive integers

#### RELATED INFORMATION

# nspin

NAME

nspin --number of spin degrees of freedom

### DESCRIPTION

The nspin variable is used to determine the number of spin components of the wave function. The default value is 1.

# **ALLOWED VALUES**

1 or 2

### RELATED INFORMATION

# polarization

**NAME** 

polarization -- algorithm used to compute the dipole and quadrupole moments

#### DESCRIPTION

The polarization variable is used to determine the algorithm used to compute the electric dipole and quadrupole. The allowed values are

•	OFF	The dipole and quadrupole are not computed. This is the default.
•	MLWF	The electronic dipole is defined as the center of charge of
		maximally localized Wannier functions (MLWF).
•	MLWF_REF	The electronic dipole is defined as the center of charge of
		maximally localized Wannier functions (MLWF) with the
		refinement correction proposed by M. Stengel and N. Spaldin, Phys.
		Rev. B73, 075121 (2006).
•	MLWF_REF_Q	The electronic dipole is defined as for MLWF_REF and the
		quadrupole moment is computed.
•	BERRY	The electronic dipole is computed using the definition based on
		the Berry phase as defined by I. Souza et al, Phys. Rev. Lett. 89,
		117602 (2002).

The ionic, electronic and total dipole  $\vec{d}$  are computed and printed in units of (electron Bohr). Note that the total dipole is only defined modulo an additive constant multiple of the lattice vectors in a periodic system. A dipole expressed in (electron Bohr) can be converted to (Debye) using the relation 1 (electron Bohr) = 2.5417462 (Debye). Conversely, dipoles expressed in (Debye) can be converted to (electron Bohr) using the relation 1 (Debye) = 0.39343031 (electron Bohr).

If an external field is applied, i.e. the displacement field  $\vec{D} = (D_x, D_y, D_z)$  defined by the <u>e\_field</u> variable is non-zero, the calculation of the electronic ground state takes into account the presence of the external field. The electric enthalpy is included in the total enthalpy.

The polarizability tensor  $\alpha_{ij}$  of a molecule is defined as

$$\alpha_{ij} = \frac{\delta d_i}{\delta D_i}$$

and is defined in units of (Bohr³),  $\delta d_i$  in (electron Bohr), and  $\delta D_j$  in (Hartree/(electron Bohr)). The polarizability tensor  $\alpha_{ij}$  of a molecule can be computed using calculations of the total dipole induced by small changes  $\delta D_j$  around  $\vec{D}{=}0$ .

The change in dipole  $\delta d_i$  due to a small change in applied field  $\delta D_j$  is computed using the centered finite difference expression

$$\delta d_i = \frac{1}{2} \left[ d_i (+\delta D_j) - d_i (-\delta D_j) \right]$$

The polarizability tensor can then be expressed as

$$\alpha_{ij} = \frac{d_i(+\delta D_j) - d_i(-\delta D_j)}{2\delta D_i}$$

The full polarization tensor can be obtained using 6 calculations of the dipole (2 evaluations of  $\delta d_i(\delta D_j)$  in 3 directions). Note that since the polarization due to a finite field includes linear and higher order terms, this calculation must be repeated with decreasing values of  $\delta D_j$  in order to obtain an accurate value of the linear polarizability.

In solids, the average polarization (dipole per unit volume) is defined as

$$\vec{P} = \frac{\vec{d}}{\Omega}$$

where  $\Omega$  is the unit cell volume. The susceptibility tensor  $\chi_{ij}$  and the dielectric tensor  $\epsilon_{ij}$  are related by  $\epsilon = I + \chi$  and the polarization is

$$\vec{P} = \chi \vec{E} = \chi \epsilon^{-1} \vec{D} = (I - \epsilon^{-1}) \vec{D}$$

A small change in applied field  $\delta D$  induces a change of polarization  $\delta \vec{P}$ 

$$\delta \vec{P} = (I - \epsilon^{-1}) \delta \vec{D}$$

If a sufficiently large unit cell is used, an estimate of the inverse dielectric tensor can be computed using the finite-difference expression

$$\left(\epsilon^{-1}\right)_{ij} = \delta_{ij} - \frac{\delta P_i(+\delta D_j) - \delta P_i(-\delta D_j)}{2\delta D_j}$$

or, in terms of the dipole moment,

$$(\epsilon^{-1})_{ij} = \delta_{ij} - \frac{\delta d_i (+\delta D_j) - \delta d_i (-\delta D_j)}{2\Omega \delta D_i} .$$

The quadrupole moment tensor is computed if the MLWF\_REF\_Q value is used. It is defined as

 $Q_{ij} = e \int_{\Omega} x_i x_j \rho(x) d^3 x$  in units of (e Bohr²). Note that this may differ from other definitions used in the literature, which may include a factor 1/2 in the above definition. The traceless quadrupole tensor defined as  $Q - (\frac{1}{3} Tr Q) I$  is also computed and printed. Some definitions used in the literature include an extra multiplicative factor of 3, or 3/2.

The quadrupole moment is sometimes expressed in units of (Debye Å), related to (e Bohr²) by the expressions 1 (Debye Å) = 0.743476 (e Bohr²) and 1(e Bohr²) = 1.3450336 (Debye Å). The NIST Computational Chemistry Database <a href="http://cccbdb.nist.gov">http://cccbdb.nist.gov</a> provides quadrupole moments of molecules in (Debye Å). A comparison of Qbox results with NIST values can be made using the relation:

$$Q^{NIST}$$
 (Debye Å) = (3/2) 1.3450336  $Q^{Qbox}$  (e Bohr<sup>2</sup>).

Born effective charges are defined as the derivatives of ionic forces with respect to the applied field

$$Z_{ij}^{\alpha} = \frac{\delta F_i^{\alpha}}{\delta D_i}$$

where  $F_i^{\alpha}$  is the *i*th component of the force on atom  $\alpha$ . Born effective charges can be similarly computed using finite difference expressions.

As of release 1.62.3 the calculation of the polarization is only implemented at the  $\Gamma$  point of the Brillouin zone, for systems having no spin polarization.

#### ALLOWED VALUES

OFF, MLWF, MLWF REF, MLWF REF Q, BERRY

### RELATED INFORMATION

e field

# ref cell

**NAME** 

ref cell --reference unit cell

### **DESCRIPTION**

The ref\_cell variable determines the size of the reference unit cell. The reference unit cell is used in constant pressure calculations to ensure constant resolution of the basis set as the unit cell changes size. The unit cell must always be enclosed in the reference unit cell during a constant pressure calculation.

#### ALLOWED VALUES

positive real numbers

### RELATED INFORMATION

cell

# scf tol

**NAME** 

scf tol-tolerance for convergence of SCF iterations

### **DESCRIPTION**

The scf\_tol variable determines the energy tolerance criterion for convergence of SCF iterations. The unit is (a.u.) (Hartree). The default value is zero. The maximum number of SCF iterations is determined by the second argument of the run command: "run niter nitscf nite". If scf\_tol is set to a non-zero value and if the energy changes by less than scf\_tol in three successive SCF iterations, the remaining SCF iterations are skipped. If scf\_tol is zero, the number of SCF iterations is nitscf.

### ALLOWED VALUES

non-negative real numbers

#### RELATED INFORMATION

<u>run</u>

#### stress

NAME

stress --stress calculation control variable

### **DESCRIPTION**

The stress variable determines whether the stress tensor is calculated. The possible values are ON and OFF. The default is OFF.

#### **ALLOWED VALUES**

ON, OFF

#### RELATED INFORMATION

ext stress, cell dyn

### thermostat

**NAME** 

thermostat --thermostat control variable

#### DESCRIPTION

The thermostat variable determines what type of thermostat is used for constant temperature simulations. The choices are

- OFF: No thermostat is used. This is the default.
- SCALING: Scaling of velocities. At each MD step, the velocities of all atoms are rescaled as  $v_i = v_i (1 \eta dt)$

where

$$\eta = \frac{1}{\tau} \tanh \frac{T - T_{ref}}{\Lambda}$$

- $\tau$  is the thermostat time constant (variable <u>th\_time</u>), T is the instantaneous temperature computed from the ionic kinetic energy,  $T_{ref}$  is the thermostat reference temperature (variable <u>th\_temp</u>), and  $\Delta$  is the thermostat temperature width (variable <u>th\_width</u>).
- ANDERSEN: Andersen thermostat. The atoms are subjected to random collisions with particles drawn from a Maxwell distribution of velocities with temperature  $T_{\text{ref}}$ . The collision frequency is  $1/\tau$  where  $\tau$  is given by the variable th time (see H. C. Andersen, J. Chem. Phys. 72, 2384 (1980)).
- LOWE: Lowe thermostat. Pairs of atoms are subjected to random collisions with a particle drawn from a Maxwell distribution of velocities with temperature T<sub>ref</sub>. The collision frequency is 1/τ where τ is given by the variable th\_time. The Lowe thermostat is similar to the Andersen thermostat but conserves total momentum (see C. P. Lowe, Europhys. Lett. 47, 145 (1999)).

• BDP: Bussi-Donadio-Parrinello thermostat. The stochastic thermostat described by Bussi, Donadio and Parrinello in J. Chem. Phys. **126**, 014101 (2007) is used. The parameter τ used in the BDP paper is the value of the variable th\_time. The value of the variable th\_width is ignored. When using the BDP thermostat, the value of <econst> printed on output corresponds to the conserved energy described in the above paper.

### ALLOWED VALUES

OFF, SCALING, ANDERSON, LOWE, BDP

#### RELATED INFORMATION

th temp, th time, th width,

# th\_temp

**NAME** 

th temp --thermostat reference temperature

# **DESCRIPTION**

The th\_temp variable determines the thermostat reference temperature. The default is zero. See thermostat.

# **ALLOWED VALUES**

non-negative real numbers

#### RELATED INFORMATION

th time, th width, thermostat

# th\_time

NAME

th time --thermostat time constant

#### DESCRIPTION

The th\_time variable determines the thermostat time constant. The time constant is a measure of the time over which the thermostat adjusts the temperature. See thermostat for details. The value of th\_time must be given in atomic units of time. The default value is 5000 a.u. (~ 120 fs).

# **ALLOWED VALUES**

positive real numbers

#### RELATED INFORMATION

th temp, th width, thermostat

# th width

NAME

th\_width --thermostat temperature window

DESCRIPTION

The th\_width determines the value of the thermostat temperature width. See <u>thermostat</u> details. The value of th\_width must be given in Kelvin units. The default value is 100 K.

**ALLOWED VALUES** 

positive real numbers

### RELATED INFORMATION

th temp, th time, thermostat

# wf\_diag

**NAME** 

wf diag --diagonalization control variable

#### DESCRIPTION

The wf\_diag variable determines whether eigenvectors and/or eigenvalues of the hamiltonian are computed after each optimization of the electronic structure. The choices are

- T: True. Eigenvalues and eigenvectors are computed. Eigenvalues are printed on output in eV units.
- F: False. Eigenvalues and eigenvectors are not computed. This is the default.
- EIGVAL: Eigenvalues only. The eigenvalues are computed and printed, but eigenvectors are not computed.
- MLWF: Maximally localized Wannier Functions (MLWFs) are computed.
- MLWFC: The position of MLWF centers is computed and printed but MLWFs are not computed.

If empty states are added to the calculation (variable  $\underline{\text{nempty}} > 0$ ) the eigenvalues and eigenvectors are always computed.

**ALLOWED VALUES** 

T, F, EIGVAL, MLWF, MLWFC

RELATED INFORMATION

# wf\_dyn

**NAME** 

wf dyn -wave function dynamics control variable

### DESCRIPTION

The wf\_dyn variable determines which algorithm is used to update the wave functions during electronic structure optimization. The choices are

 $\bullet$  SD: Steepest Descent. This the default. Wavefunctions are updated as follows:  $\psi^{(k+1)} = \psi^{(k)} - \alpha \, H \, \psi^{(k)}$ 

where

$$\alpha = \begin{cases} \frac{1}{2E_{cut}} & m_e = 0\\ \frac{dt^2}{m_e} & m_e > 0 \end{cases}$$

and  $m_e$  is the fictitious electronic mass (variable emass). By default,  $m_e$ =0. (Note that the SD algorithm is very inefficient and is not used in practice. It is provided for debugging purposes).

• PSD: Preconditioned Steepest Descent. Wavefunctions are updated as follows:  $\psi^{(k+1)} = \psi^{(k)} - \alpha K P_{\perp} H \psi^{(k)}$ 

where K is a preconditioning matrix

$$k(G) = \begin{cases} k_{ij} = \delta_{ij} k(G) \\ \frac{1}{2E_{cut}^{prec}} & \frac{1}{2}G^2 < E_{cut}^{prec} \\ \frac{1}{2}E & \frac{1}{2}E \ge E_{cut}^{prec} \end{cases}$$

and  $P_{\perp} = I - \sum_{i} |\Psi_{i}\rangle\langle\Psi_{i}|$  is a projector on the orthogonal complement of the subspace spanned by all wavefunctions. The preconditioning matrix depends on the value of  $E_{cut}^{prec}$  (variable ecutprec).

- PSDA: Preconditioned Steepest Descent with Anderson acceleration. Wavefunctions are updated as with the PSD option, and convergence is accelerated by the Anderson scheme (D. G. Anderson, JACM 12, No 4, pp. 547-560 (1965)).
- JD: Jacobi-Davidson. Wavefunctions are updated using a preconditioned Jacobi-Davidson algorithm.
- MD: Molecular Dynamics. Wavefunctions are updated using the Car-Parrinello scheme  $\psi_i^{(k+1)} = 2 \psi_i^{(k)} \psi_i^{(k-1)} \frac{dt^2}{m_a} H \psi_i^{(k)} + \sum_j \Lambda_{ij} \psi_j^{(k)}$

where holonomic constraints are used to enforce orthogonality.

LOCKED. Wavefunctions are not updated.

#### ALLOWED VALUES

SD, PSD, PSDA, JD, MD, LOCKED

RELATED INFORMATION

ecutprec

#### XC

**NAME** 

xc --exchange-correlation functional control variable

### **DESCRIPTION**

The xc variable determines which exchange-correlation functional is used in the electronic structure calculation. The choices are

• LDA: Local Density Approximation, Ceperley-Alder data. This is the default.

- VWN: Local Density Approximation, parameterized by Vosko, Wilk and Nusair.
- PBE: Perdew-Burke-Ernzerhof GGA functional.
- PBE0: Hybrid density functional [C. Adamo and V. Barone, JCP 110, 6158 (1999)].
- B3LYP: Three-parameter Becke-Lee-Yang-Parr hybrid density functional.

# ALLOWED VALUES

LDA, VWN, PBE, PBE0, B3LYP

RELATED INFORMATION