

LIO User Guide

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Version Developers

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Introduction

What is LIO?

Welcome to the LIO project! LIO is a library that can perform electronic structure calculations using density functional theory.

Instalation

If you are reading this manual, you probably already have a version of LIO ready to compile. If you don't, or if you want to make sure you have the most most up-to-date version of the code, all you need to do is either download it the git repository online or use git to clone a copy.

For the first option, go to https://github.com/MALBECC/lio and click on the green button that says clone or download and click on Download ZIP.

For the second one, you can directly run the following command:

```
{\tt git clone https://github.com/MALBECC/lio.git} \quad .
```

Pre-requisites

In addition to an UNIX-like OS, fortran and c++ compilers, LIO depends on LAPACK LAPACK and BLAS libraries for linear algebra calculations. In addition, $CUDA\ 6.5$

or higher is required for GPU calculations, which unleash LIO's true potential. As of the writing of this manual, LIO has not yet been tested with CUDA 9.0.

In addition, *libxc* is required for its usage, although said library is CPU-only. This is entirely optional as LIO can run without libxc, using only the PBE functional.

Compilation

By default, LIO compiles with GPU options enabled. It is highly recommended to specify the GPU architecture as a compilation option, since the compiler performs additional enhancements. After compilation, LIOHOME environment variable should be set to the current LIO installation directory:

For a CPU-only compilation, use:

If INTEL compilers are present, they can be used by setting the *intel* option to 1, or to 2 if INTEL MKL usage is also desired.

make
$$intel=2$$

The following is a list of available compilation and their meanings. They can be used in any combination possible (including several GPU architectures for greater compatibility). For example, the default LIO compilation could be written as:

make
$$cuda=2 sm30=1 sm52=1 sm61=1$$

Variable	Description
cuda 0, 1, 2	default cuda=2 Sets the level of GPU dependencies. cuda=0 means a CPU-only compilation, =1 enables GPU, and =2 includes CUBLAS usage for linear algebra operations.
intel 0, 1, 2	default intel=0 Sets the usage of INTEL compilers. intel=0 means only GNU compilers, =1 means INTEL, and =2 uses INTEL MKL instead of BLAS/LAPACK routines.

Table 1.1: Compilation options

Variable	Description
precision	default precision=0
0, 1	Sets level of precision in XC calculations. By default LIO uses mixed precision; precision=1 sets everything in double precision. This is specially useful when attempting high-precision geometry optimizations.
analytics	$default\ analytics{=}0$
0, 1, 2, 3	Setting analytics = 1 specifies a profiling compilation, while analytics = 2 and 3 set higher levels of debugging information (for usage with gdb, for example).
$\mathrm{sm}\mathrm{X}\mathrm{X}$	$default \ sm30{=}1 \ sm52{=}1 \ sm61{=}1$
0, 1	Sets a specific GPU architecture for compilation. Available options are sm30, sm35 (Kepler, CUDA \geq 5.0), sm50, sm52 (Maxwell and GeForce 980, CUDA \geq 6.5), sm60 and sm61 (Pascal and GeForce 1080, CUDA \geq 8.0).

Table 1.1: Compilation options

MD-Engine Interfacing

LIO can be linked with AMBER, our own GROMACS fork, our own HYBRID code for QM/MM calculations.

In all three cases, the LIOHOME environment variable should be set to the current LIO installation directory. In addition to this section, please refer to each of the software packages' installation manual for further clarifications.

In order to compile AMBER with LIO, AMBER should be compiled after LIO with the following options set:

```
export AMBERHOME=/dir/to/amber/
./configure -lio -noX11 -netcdfstatic gnu
make clean
make install
```

For a GROMACS-LIO compilation, GROMACS should be compiled after LIO with the following options:

```
cd gromacs_compilation_directory/
cmake gromacs_src_dir/ -DGMX_QMMM=1

-DGMX_QMMM_PROGRAM="lio" -DLIO_LINK_FLAGS="-L/usr/lib
-L/usr/lib64 -L$LIOHOME/g2g -L$LIOHOME/lioamber -lg2g
-llio-g2g" -DGMX_GPU=0 -DGMX_THREAD_MPI=0

make
make install
```

Running LIO

In order to run LIO as a QM stand-alone program, you must first set the LIOHOME environment variable to that of the current LIO installation path (if it was not set before), and add /liosolo folder to the current excecutable paths.

```
export LIOHOME=/dir/to/lio/
export PATH=$PATH:$LIOHOME/liosolo
```

Then, LIO can be exceduted as follows:

```
liosolo -i input file -c coords.xyz
```

Where input_file contains the input options and coords.xyz contains the coordinates in XYZ format. For more options, please refer to Chapter 2.

When running LIO with AMBER, you only need to specify 'qm_theory=extern' in the &qmmm section of the code (please refer to the AmberTools manual for further &qmmm options), in addition to specifying both the QM system and its charge. When running with GROMACS, you only need to specify the QM system and its charge. In both cases, additional LIO options will be read from a 'lio.in' file in the working directory.

Tips and tricks - Optimizing your runs

There are several ways to optimize your production runs, and it is highly recommended to fine tune these settings if the same system or very similar systems are going to run for extended periods of time. Please see the corresponding sections in this manual for a more detailed explanation of these variables.

The first set to tune is rmax and rmaxs, which are related to the integral cut-offs used in both Coulomb and fitting set integrals. The *higher* the value of rmaxs and the lower the value of rmax, the calculations will run faster. Keep in mind, however, that rmax should never be lower that rmaxs, and that tuning those values for a faster calculation will certainly result in a loss of precision. Therefore, the idea is to decrease rmax and increase rmaxs until the difference in energies and forces stops being negligible.

Next is the set of options available for the GPU library which performs the exchange-correlation calculations. The option $max_function_exponent$ indicates the maximum exponent considered for a fuction in a point of the grid; it should be tweaked taking into account the aforementioned criterium: the faster the calculation is performed, the lesser precision is achieved. There are also three other options whose optimal values depend on the GPU architecture available: $little_cube_size$, $min_points_per_cube$, and $sphere_radius$. These can be tweaked for maximum speed without worrying about the resulting precision.

General Settings

In this chapter we will describe the basic settings for running the code. These will be necessary or useful for any of the features of the code that you might be interested in using.

General System Description

Variable	Description
natom	integer = 0
	Sets the number of atoms only in the QM region. This is
	ignored when calling LIO from another MM engine.
nsol	integer = 0
	Sets the number of "solvent" atoms, meaning atoms in the
	MM region. This is also ignored when calling LIO externally.
$_{ m charge}$	integer = 0
	Sets the total charge for the QM region.
OPEN	logical = .false.
	Indicates whether an open-shell calculation should be per-
	formed.
nUnp	integer = 0
-	Table 2.1: Conoral setup

Table 2.1: General setup

Variable	Description
	Indicates the number of unpaired electrons for an open-shell calculation, in a closed-shell calculation this variable is ignored. Keep in mind that the number of unpaired electrons is NOT the multiplicity, rather nUnp = Mult+1.

Table 2.1: General setup

Common Inputs and Outputs

Variable	Description
style	logical = .false. Activates a more stylish version of the output.
verbose	 integer = 0 Sets verbose level for output. 0 = Nothing is printed save for a small LIO start message. 1 = only the input namelist and the final convergence results are printed. 2 = information for each iteration is added. 3 = GPU module information is added. 4 = additional miscellaneous information is printed.
write XYZ	logical = .false. Writes and xyz file containing the QM region, using the file-name specified in fcoord. This file is written in each of the MD steps when LIO is used with AMBER/GROMACS/HY-BRID.
fCoord	character * 20 =' qm.xyz' Output file containing the QM region.
timers	 integer = 0 Sets the timer level for benchmarking and profiling. 0 = no timers. 1 = only timers in certain subroutines. 2 = full timer summary.
dBug	logical = .false. Checks for NaNs in Fock and Rho matrices.

Table 2.2: Common inputs and outputs

Variable	Description
gaussian_convert	logical = .false. Reads an electron density from Gaussian 09 as starting guess. This feature is still experimental.
print_coeffs	logical = .false. Prints coefficients and energies for each molecular orbital.
VCInp	logical = .false. Reads a MO coefficient restart from frestartin.
$\operatorname{restart_freq}$	$integer=1$ Writes a MO coefficient restart every restart_freq iterations into frestart.
frestart	char * 20 = 'restart.out' Name of the output density or coefficient restart file.
frestartin	char * 20 = 'restart.in' Name of the input density or coefficient restart file.
$\operatorname{rst} \operatorname{\underline{-dens}}$	integer = 0 When rst_dens >= 1, the restart read by VCInp must be a density matrix (not coefficient matrix) restart. Likewise, when rst_dens = 2, the restart written with restart_freq is a density matrix restart (instead of a coefficient matrix restart).

Table 2.2: Common inputs and outputs

Properties Calculations

Variable	Description
writeForces	logical = .false. Performs forces calculation and prints the result to a file named "forces".
dipole	logical = .false. Performs dipole calculation and prints the result to a file named "dipole".

Table 2.3: Common inputs and outputs

Variable	Description
$\operatorname{mulliken}$	logical = .false.
	Performs a Mulliken population analysis and outputs the re-
	sult to a file called "mulliken".
lowdin	logical = .false.
	Performs a Lowdin population analysis and outputs the re-
	sult to a file called "mulliken".
fukui	logical = .false.
	Calculates the condensed-to-atoms Fukui function for the
	system, calculating the spin-polarized version in open-shell
	systems.

Table 2.3: Common inputs and outputs

GPU Options

These options affect the calculations in the GPU module. For the exchange correlation integrals, the integration grid is separated into cubes and spheres, with the smaller cubes (less points per cube) being calculated in CPU while the rest is calculated in GPU. Most of these options should be tweaked for optimal performance in a given system.

Variable	Description
gpu_level integer	default = 4 Determines which calculations are performed by the GPU, only available when compiled with cuda > 0. 0/1 = only exchange-correlation integrals. 2 = adds QM/MM interaction energies and gradients. 3 = adds Coulomb energies and gradients. 4 = adds nuclear attraction energies and gradients. 5 = adds Coulomb basis fitting energies and gradients.
$ooknote{max_function_exponent} integer$	$default = 10$ Ignores functions with $ exponent > max_function_exponent$. This is only for the exchange-correlation calculations.

Table 2.4: GPU Module Options

Variable	Description
little_cube_size double precision	default = 8.0d0 Small cube-type point group size (in Angstrom).
$egin{array}{c} \mathbf{min_points_per_cube} \ integer \end{array}$	default = 1 Minimum number of grid points in a cube.
$\begin{array}{c} \mathbf{assign_all_functions} \\ logical \end{array}$	default = .false. Calculate all functions (ignores max_function_exponent). This is intended only as a debug option and its usage is not recommended.
sphere_radius double precision	default = 0.6d0 Radius of the sphere-type point groups. 0 means there are no sphere-type groups, 1 means all points are contained in sphere-type groups.
${\bf remove_zero_weights} \\ logical$	default = .true. Discard functions for those whose weight is zero (.false. option only remains as a debug option).
$rac{ ext{energy_all_iterations}}{logical}$	default = .false. Calculate Exc energy in all SCF iterations. Usually, XC energy is only calculated in the final step in order to accelerate calculations.
free_global_memory double precision	default = 0.0d0 Fraction of global GPU memory available for the calculation (1 means 100%).

Table 2.4: GPU Module Options

Model Hamiltonian

Density Functional Theory

Variable	Description
iexch	integer = 9 Identifies the exchange-correlation potential to use with the calculation when not using libxc. Iexch=9 is the only option currently available.
use_libxc	logical = .false. Activates the use of libxc version of the XC potential.
${\it ex_functional_id}$	integer = none Exchange functional to use with libxc. Please refer to libxc for the desired functional.
${\it ec_functional_id}$	integer = none Correlation functional to use with libxc. Please refer to libxc for the desired functional.
int_basis	$logical = .false.$ If true, looks for the internal basis indicated in variables $basis_set$ and $fitting_set$ (defaults are $'DZVP'$ and $'DZVPCoulombFitting'$. If false, an external basis file must be provided in the keyword $basis_set$.
basis_set	char * 20 =' DZVP'

Table 3.1: DFT Hamiltonian

Variable	Description
	Name of the basis set used in the calculation when <i>int_basis</i> is set to true. If not, it is the name of the file containing the custom basis data.
fitting_set	char*100 =' DZVPCoulombFitting' Name of the fitting set used in the calculation when int_basis is set to true.
n_ghosts	integer = 0 Number of ghost atoms. Ghost atoms are considered for the basis functions of the system, but they are considered as having zero electrons and zero nuclear charge.
$ghost_atoms$	$integer=0$ A list containing the atom indeces for those considered ghost_atoms. For example, $ghost_atoms=1,2$ considers the first and second atoms as ghosts.
rmax	double=16.0d0 Maximum exponent in 3-center integrals. If exponent is greater than $rmax$, the current term is ignored.
rmaxs	double = 5.0d0 If the exponent in 3-center integral is within $rmax$ and $rmaxs$, calculation is performed using double precision. If it is between 0 and $rmaxs$, the calculation is performed in single precision. This is ignored when using the AINT module.
iGrid	integer=2 Grid type when iterating through SCF. Available values are 1 (less dense) and 2 (more dense).
iGrid2	integer=2 Grid type for the final density calculation in SCF. Available values are 1 (less dense) and 2 (more dense).

Table 3.1: DFT Hamiltonian

Convergence Options

Variable	Description
initial_guess	integer = 0 Selects the method for calculating a starting guess. When set to 0, the initial guess comes from the 1-electron integrals, while setting it to 1 performs an Aufbau-like initial guess.
nMax	integer = 100 Maximum number of SCF iteration steps.
told	double $precision = 1.0d - 6$ Criterium for the maximal square deviation of the density matrix to consider that the convergence has been achieved.
etold	double $precision = 1.0d0$ Criterium for the maximal energy difference to consider that the convergence has been achieved.
DIIS	logical = .true. Uses DIIS algorithm for convergence. If disabled, it tries to converge using a damping factor between iterations.
nDIIS	integer = 30 Number of matrices considered for DIIS. Only change this if you know what you are doing.
gold	$\label{eq:convergence} \begin{split} &double\ precision = 1.0d + 1\\ &\text{Determines the weight of the previous density matrix in the}\\ &\text{linear combination with the new one when using the damping}\\ &\text{convergence method (DIIS} = .\text{false.} \ / \ \text{hybrid_converg} \). \end{split}$
hybrid_converg	 logical = .false. Uses the hybrid convergence algorithm: it starts using the damping factor, and after a threshold is met, changes to DIIS. This method usually gives the best results in cases which are difficult to converge.
$\operatorname{good} _\operatorname{cut}$	double $precision = 1.0d - 3$ Sets the threshold to start DIIS when activating hybrid convergence (hybrid_converg = 1). When the mean square deviation for the density matrix between two iteration steps reaches this threshold, DIIS is activated.

Table 3.2: BO-MD useful settings.

External Electric Fields

Effective Core Potentials

Single Atom Potentials

Tight Binding Interface

Ground State Calculations

Single-point and Born-Oppenheimer Molecular Dynamics

Single-point calculations consist in finding the ground state density that minimizes the energy for a given nuclei distribution. Once found, the program can also calculate the force field of that density for the MD-engine to move the nuclei.

Variable	Description
initial_guess	integer=0 Selects the method for calculating a starting guess. This is only useful for the first MD step, since after that the starting guess is the electron density from the previous step.
nMax	integer = 100 Maximum number of SCF iteration steps.
told	double $precision = 1.0d - 6$ Criterium for density matrix convergence.
etold	double $precision = 1.0d0$ Criterium for energy convergence.
DIIS	logical = .true. Uses DIIS algorithm for convergence.
hybrid_converg	logical = .false. Table 4.1: BO-MD useful settings

Table 4.1: BO-MD useful settings.

Variable	Description
	Uses the hybrid convergence algorithm.
$\operatorname{good} _\operatorname{cut}$	double $precision = 1.0d - 3$ Sets the threshold to start DIIS when activating hybrid convergence.
VCInp	logical = .false. Reads a MO coefficient restart.
$\operatorname{restart_freq}$	$integer = 1$ Writes a MO coefficient restart every restart_freq steps.
frestart	char * 20 = 'restart.out' Name of the output restart file.
frestartin	char * 20 = 'restart.in' Name of the input restart file.
$\operatorname{rst_dens}$	<pre>integer = 0 rst_dens = 1 reads a density matrix restart, while rst_dens = 2 both reads and writes a density matrix restart.</pre>
basis_set	char * 20 =' DZVP' Name of the basis set used in the calculation.

Table 4.1: BO-MD useful settings.

Geometry optimizations

Geometry optimizations or energy minimization is the process of finding an atomic arrangement in space where the force on each atom is acceptably close to zero.

Implemenation

LIO has a simple steepest-descent algorithm. The idea is to move the system in the force direction, at a λ step value.

$$\vec{r}_{new}^i = \vec{r}^i + \lambda \vec{F}^i \tag{4.1}$$

Without a linear search algorithm λ is obtained as $\frac{steep_size}{|\vec{F}_{max}|}$. If the energy decreases with the movement, the step is accepted; but if the energy increases with the step, the steep is rejected and λ is reduced. Each accepted move increases step size a 20% and each rejected move decreases step size a 50%.

In a linear search algorithm the system scans the energy as function of λ and predicts the best value of λ to move the system in the gradient direction.

Best λ in lineal search algorithm is obtained by a quadratic function ajusted using minimum energy of the scan and previous and next points.

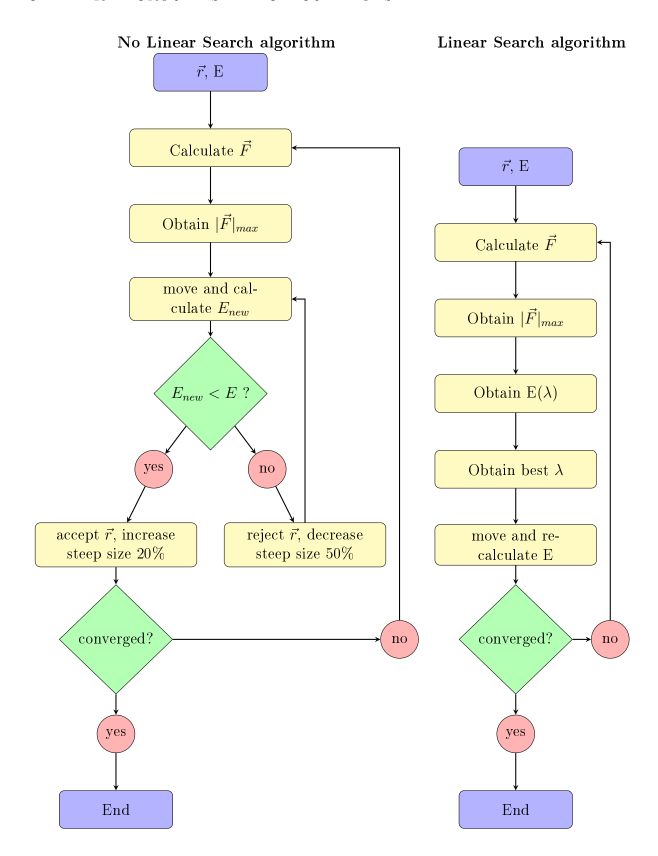
Using geometry optimizations

Adding steep=t in LIO input enables geometry optimization (steepest descent, lineal search by default). Convergence criteria are set by Force_cut and Energy_cut (5E-4 Hartree/bohr and 1E-4 Hartree by Default). The number of minimization steeps is set by n_min_steps (500 by default) and initial distance steep is set by minimization_steep (by default 0.05 bohr)

It is highly advisable to compile LIO in double precision in order to minimise the error in exchange-correlation forces (precision=1). Outputs of geometry optimizations are traj.xyz (atoms coordinates in each steepes descent movement) and optimization.out (steep, energy and others). If verbose=true optimization.out includes the energy of each linear search point.

Examples

Examples of geometry optimization are made in lio/test/13 geom optim.



Restraints

LIO may add an extra potential term to the Hamiltonian in order to restrain the distance between specified pairs of atoms.

Implemenation

The implementation is a simple harmonic potential over a generalized coordinate r.

$$U = \frac{1}{2}k[r - l_0]^2 \tag{4.2}$$

r may be defined as a weighted combination of distances between pairs of atoms.

$$r = \sum_{i} \sum_{j>i} w_{ij} |\vec{r_i} - \vec{r_j}| \tag{4.3}$$

In this formulation the force over an atom l is:

$$\vec{F}_{l} = -k[r - l_{0}] \sum_{i} \sum_{j>i} w_{ij} \frac{\vec{r}_{ij}}{r_{ij}} \eta_{ijl}$$
(4.4)

Where η_{ijl} is defined as:

$$\eta_{ijl} = \begin{cases} 1 & \text{if } l = i \\ -1 & \text{if } l = j \\ 0 & \text{in other case} \end{cases}$$

Using Restraints

The number of pairs of atoms to be added in the restraint potential(s) is defined by setting the variable number_restr, and a list of distance restrains have to be added to in an additional lio.restrain file. For example:

Columns a_i and a_j contain the atom numbers in the QM system to be restrained, while the index number determines which distances contribute to a same generalized reaction coordinate. The remaining columns are the force constants (k), weights of that distance in the generalized coordinate (w_{ij}) and equilibrium positions in atomic units (l_0) .

Examples

1)In lio.in:

 $number_restr = 1$

in lio.restrain:

$$a_i$$
 a_j index k w_{ij} l_0
1 2 0 0.1 1.0 7.86

Potential added to system:

$$U = \frac{1}{2}0.1 \left[1.0 |\vec{r_1} - \vec{r_2}| - 7.86 \right]^2 \tag{4.5}$$

2)In lio.in:

number restr = 2

in lio.restrain:

$$a_i$$
 a_j index k w_{ij} l_0
1 2 0 0.1 1.0 7.86
3 4 0 0.1 -1.0 7.86

Potential added to system:

$$U = \frac{1}{2}0.1 \left[1.0 |\vec{r_1} - \vec{r_2}| - 1.0 |\vec{r_3} - \vec{r_4}| - 7.86 \right]^2$$
 (4.6)

3)In lio.in:

 $number_restr = 4$

in lio.restrain:

Potential added to system:

$$U = \frac{1}{2}0.1 \left[1.0 |\vec{r_1} - \vec{r_2}| - 1.0 |\vec{r_3} - \vec{r_4}| - 7.86 \right]^2 + \frac{1}{2}0.3 \left[3.5 |\vec{r_1} - \vec{r_3}| - 2.2 |\vec{r_7} - \vec{r_8}| + 2.31 \right]^2$$
(4.7)

Electron Dynamics

Real Time TD-DFT

Electronic transport

Ehrenfest Dynamics

Post-Processing Tools

TD-Analize: Electronic Spectra

CubeGen: Orbital and Density Visualization

Reference Section

This section contains a quick reference for all of LIO's input variables and commandline options. For more detailed descriptions, please refer to the previous chapters.

Command line options

Variable	Description
-i file_name character*20	default = 'lio.in' Name of the input file containing LIO options.
-c crd_file.xyz character*20	default = 'qm.xyz' Name of thte XYZ file containing coordinates.
-b basis_file character*20	default = 'basis' A file containing the basis set and fitting set data, only used when int_basis=f.
-v logical	default = .false. Sets verbose level to 4.

Table 7.1: Command Line

Keywords - General Setup

Variable	Description
natom	default=0
integer	Number of QM atoms in the system.
nsol	default=0
integer	Number of classical atoms in the system.
charge	default=0
integer	Total charge of the QM system.
open	default = .false.
logical	Perform an open-shell calculation.
nunp	default=0
integer	Number of unpaired electrons for open-shell calculations.
style	default = .false.
logical	Activates a formatted version of the output.
\mathbf{fcoord}	default = 'qm.xyz'
character *20	Name of the output file for the coordinates of the QM system.
writexyz	de fault = .false.
logical	Writes an xyz file containing the QM system.
verbose	default=1
integer	Determines the amount of information printed, from 0 (nothing) to 5 (everything).
timers	default=0
integer	Activates timers $(=1 \text{ or } =2)$.
dbug	de fault = .false.
logical	Checks for NaNs.
write Forces	de fault = .false.
logical	Writes final forces to an output file.

Table 7.2: General Setup

Variable	Description
dipole logical	default = .false. Calculates and prints dipole moment.
$egin{aligned} \mathbf{mulliken} \ logical \end{aligned}$	default = .false. Performs a Mulliken Population Analysis.
$egin{aligned} \mathbf{lowdin} \ logical \end{aligned}$	default = .false. Performs a Lowdin Population Analysis.
fukui logical	default = .false. Calculates atomic Fukui function.
$\begin{array}{c} \mathbf{gaussian_convert} \\ logical \end{array}$	default = .false. Reads a Gaussian09 density matrix.
$egin{array}{c} \mathbf{print_coeffs} \ logical \end{array}$	default = .false. Prints MO coefficients in AO basis.

Table 7.2: General Setup

Keywords - GPU Options

Variable	Description
$egin{aligned} \mathbf{gpu_level} \ integer \end{aligned}$	default = 4 Determines which calculations are performed by the GPU. (0 = only XC, 5 = everything).
$egin{array}{c} \mathbf{max_function_exponent} \\ integer \end{array}$	$default = 10$ Ignore functions with $ exponent > max_function_exponent.$
$egin{aligned} & ext{little_cube_size} \ & double\ precision \end{aligned}$	default = 8.0d0 Small cube-type point group size.
$egin{array}{c} \mathbf{min_points_per_cube} \ integer \end{array}$	default = 1 Minimum number of grid points in a cube.
$\begin{array}{c} \mathbf{assign_all_functions} \\ logical \end{array}$	$default = .false.$ Calculate all functions (ignores $max_function_exponent$).
sphere_radius double precision	default = 0.6d0 Proportion of points contained in sphere-type groups (from 0 to 1).
$egin{array}{c} \mathbf{remove_zero_weights} \ logical \end{array}$	default = .true. Discard functions for those whose weight is zero.
$rac{ ext{energy_all_iterations}}{logical}$	default = .false. Calculate Exc energy in all SCF iterations.
free_global_memory double precision	default = 0.0d0 Fraction of global GPU memory available for the calculation (1 means $100%$).

Table 7.3: GPU Module Options

Keywords - DFT Hamiltonian

Variable	Description
iexch integer	default = 9 Identifies the exchange-correlation potential to use with the calculation when not using libxc. Iexch=9 is the only option currently available.
$egin{array}{c} {f use_libxc} \ logical \end{array}$	default = .false. Activates the use of libxc version of the XC potential.
$rac{\mathbf{ex_functional_id}}{integer}$	default = ? Exchange functional to use with libxc.
$egin{array}{l} \mathbf{ec_functional_id} \ integer \end{array}$	default = ? Correlation functional to use with libxc.
int_basis logical	default = .true. If true, looks for the internal basis indicated in variables basis_set and fitting_set; if false, an external basis file must be provided.
$egin{array}{c} \mathbf{basis_set} \\ \mathit{character*20} \end{array}$	default='DZVP' Name of the basis set used, or the name of the custom basis set file.
$\begin{array}{c} \mathbf{fitting_set} \\ character*100 \end{array}$	default = 'DZVP Coulomb Fitting' Name of the fitting set used in the calculation.
$rac{\mathbf{n}_{-}\mathbf{ghosts}}{integer}$	default = 0 Number of ghost atoms.
$rac{ extbf{ghost_atoms}}{integer}$	default = 0 A list containing the ghost atom indeces.
$egin{array}{c} \mathbf{rmax} \ double \ precision \end{array}$	default = 16.0d0 Maximum exponent in 3-center integrals.
$egin{array}{c} \mathbf{rmaxs} \\ double \ precision \end{array}$	default = 5.0d0 Maximum exponent for 3-center integrals in single precision.

Table 7.4: DFT Hamiltonian

Variable	Description
${f iGrid} \ integer$	default = 2 Grid type when iterating through SCF.
iGrid2 integer	default=2 Grid type for final energy calculation in SCF.

Table 7.4: DFT Hamiltonian

Keywords - Effective Core Potentials

Variable	Description
ECPMode logical	$default = .false. \ Activate effective core potentials.$
$\begin{array}{c} \mathbf{ECPTypes} \\ integer \end{array}$	default = 0 Number of atoms with ECP.
$\begin{array}{c} \mathbf{tipeECP} \\ character *30 \end{array}$	default = 'NOT-DEFINED' Type of ECP used.
$egin{aligned} \mathbf{ZListECP} \ integer \end{aligned}$	default = 0 Array with Z of atoms with ECP enabled.
$egin{aligned} \mathbf{cutECP} \ logical \end{aligned}$	default = .true. Enables cuts for ECP integrals.
$egin{array}{c} {f cut 2_0} \ double\ precision \end{array}$	default = 15.d0 Cut value for 2-center ECP integrals.
cut3_0 double precision	default = 12.d0 Cut value for 3-center ECP integrals.
$rac{\mathbf{ECP_debug}}{logical}$	default = .false. Enables ECP debug mode.
$egin{aligned} \mathbf{local}_{nonlocal} \ integer \end{aligned}$	default=0 Calculates only local terms (when $=1$) or only non-local terms (when $=2$).
$rac{ extbf{ECP_full_range_}}{logical}$	$egin{aligned} \mathbf{int} & efault = .false. \ & ext{Enables full-range integral calculations.} \end{aligned}$
$egin{array}{c} \mathbf{verbose_ECP} \ integer \end{array}$	default = 0 Controls ECP verbose levels.
$egin{aligned} \mathbf{fock_ECP_read} \ logical \end{aligned}$	default = .false. Enables restart read in ECP.

Table 7.5: Effective Core Potentials

Variable	Description
$egin{aligned} \mathbf{fock_ECP_write} \ logical \end{aligned}$	default = .false. Enables restart write in ECP.
fullTimer_ECP logical	default = .false. Enables full timers in ECP.

Table 7.5: Effective Core Potentials

Keywords - DFTB Embedding

Variable	Description
$rac{{f dftb_calc}}{logical}$	default = .false. Activates the TB embedding of the system.
$egin{aligned} \mathbf{MTB} \ integer \end{aligned}$	default = 0 TODO Size of the two tight-binding subatrices.
$rac{\mathbf{end} \mathbf{bTB}}{integer}$	default = 0 TODO Index matrix size.
$rac{\mathbf{start_tdtb}}{integer}$	default = 0 TODO Initial time step for evolution of diagonal TB terms (????).
$rac{\mathbf{end_tdtb}}{integer}$	default = 0 TODO Final time step for evolution of diagonal TB terms (????).
alfaTB double precision	default = UNSET Manually sets the on-site energies (diagonal values) for the TB part of the Hamiltonian.
$\begin{array}{c} \textbf{betaTB} \\ \textit{double precision} \end{array}$	default = UNSET Manually sets the hopping terms for the TB part of the Hamiltonian (ie, the non-diagonal nearest neighbour terms for TB - TB interactions).
gammaTB double precision	default = UNSET Manually sets the hopping terms for the interaction between TB atoms and DFT atoms.
$egin{array}{c} ext{Vbias}_ ext{TB} \ double\ precision \end{array}$	default = UNSET Sets a bias for the on-site energies to simulate electrodes.
${f TBload} \ logical$	$default = .false. \ { m TODO}.$
${f TBsave} \ logical$	$default = .false. \ { m TODO}.$

Table 7.6: DFTB Embedding

Keywords - Fields and Biases

Variable	Description
field logical	default = .false. Use an external field (perturbation in TD).
$egin{aligned} \mathbf{a0} \ double\ precision \end{aligned}$	default = 1.0d3 A dividing factor in electric field calculations.
epsilon double precision	default = 1.0d0 Relative permitivity of the medium.
Fx, Fy, Fz double precision	default=0.05d0 The value of the external electric field in the x, y and z directions.
$rac{ extbf{nfields_iso}}{integer}$	default = 0 Number of shape-isotropic fields. If $nfields_iso > 0$, the inputs in $field_iso_file$ are ignored
field_iso_file character*20	default = 'field.in' Isotropic fields input file.
$rac{ extbf{nfields_aniso}}{integer}$	default = 0 Number of shape-anisotropic fields. If $nfields_aniso > 0$, the inputs in $field_aniso_file$ are ignored
$egin{aligned} \mathbf{field_aniso_file} \\ character * 20 \end{aligned}$	default = 'field.in' Anisotropic fields input file.
$rac{ extbf{fockbias}_{ extbf{is}} extbf{is}_{ extbf{active}}}{logical}$	$default = .false. \ { m TODO}.$
fockbias_is_shaped	$egin{aligned} default = .false. \ TODO. \end{aligned}$
$fockbias_readfile$ $character*80$	default = 'atombias.in' Atomic bias input file.
fockbias_timeamp0 double precision	$default = UNSET \ { m TODO}.$

Table 7.7: Fields and Biases

Variable	Description
$egin{aligned} \mathbf{fockbias_timefall} \ double \ precision \end{aligned}$	default = UNSET TODO.
$egin{aligned} \mathbf{fockbias_timegrow} \ double \ precision \end{aligned}$	$default = \mathit{UNSET} \ \mathrm{TODO}.$

Table 7.7: Fields and Biases

Keywords - Self Consistent Field

Variable	Description
$rac{ ext{initial_guess}}{integer}$	default = 0 Method for generating the initial guess for the SCF.
$egin{aligned} \mathbf{nMax} \ integer \end{aligned}$	default = 100 Maximum number of SCF steps.
$egin{array}{c} \mathbf{told} \ double \ precision \end{array}$	default = 1.0d-6 Tolerance threshold for density matrix convergence.
${\bf Etold}\\ double\ precision$	default = 1.0d0 Tolerance threshold for energy convergence.
DIIS logical	default=.true. Use DIIS convergence accelerator if true, or damping convergence accelerator if false.
$\begin{array}{c} \mathbf{nDIIS} \\ integer \end{array}$	default = 30 Number of DIIS convergence iterations.
$\begin{array}{c} \textbf{gold} \\ \textit{double precision} \end{array}$	default = 1.0d1 Proportion of old matrix to use when using the damping mixture (gold = X means that it will use an 1:X new to old proportion).
$rac{\mathbf{hybrid}}{logical}$	default = .false. Use Hybrid convergence accelerator.
$egin{array}{c} \mathbf{good_cut} \ double\ precision \end{array}$	default=1.0d-5 Tolerance threshold for damped convergence, switch to DIIS afterwards.
veinp logical	default = .false. Reads the molecular orbital coefficients from $frestart$ and uses that as the starting guess for the first SCF cycle.
$egin{array}{c} \mathbf{rst_dens} \ integer \end{array}$	default = 0 rst_dens = 1 reads a density matrix restart, while rst_dens = 2 both reads and writes a density matrix restart.

Table 7.8: Self Consistent Field

Variable	Description
frestartin character*20	default = 'restart.in' Filename for the input containing the molecular orbital coefficients to be used as starting guess when $vcinp = .true.$
$frestart \\ character*20$	default = 'restart.out' Filename for the output containing the molecular orbital coefficients.
$egin{array}{c} \mathbf{restart_freq} \ integer \end{array}$	default=0 Indicates the frequency for writing the restart: it will do so every set number of calls to LIO (that is, number of steps of nuclear moves performed by the MD-engine.

Table 7.8: Self Consistent Field

Keywords - Geometry Optimization

Variable	Description
steep logical	default = .false. Activate steepest descent algorithm for geometry optimization.
Force_cut double precision	default=5.0d-4 Convergence criteria in forces (Hartree/bohr) for geometry optimization.
Energy_cut double precision	default=1.0d-4 Convergence criteria in energy (Hartree) for geometry optimization.
minimzation_steep double precision	default = 0.05d0 Initial distance steep (bohr).
$rac{ extbf{n}_{-} extbf{min}_{-} extsf{steeps}}{integer}$	default = 500 Maximum number of geometry optimization steps.
$egin{aligned} ext{lineal} ext{search} \ logical \end{aligned}$	default = .true. Enable lineal search algorithm.
$egin{aligned} \mathbf{n} _\mathbf{points} \ integer \end{aligned}$	default = 5 Number of points scaned for lineal search.
$rac{\mathbf{number_restr}}{integer}$	default = 0 Number of distance restraints used.

Table 7.9: Geometry Optimization

Keywords - Real Time TD-DFT

Variable	Description
timeDep integer	default = 0 Use RT-TD-DFT when timeDep = 1.
tdStep double precision	default = 2.0d-5 Timestep for TD-DFT (in atomic units).
$\begin{array}{c} \mathbf{ntdStep} \\ integer \end{array}$	default = 0 Total number of TD-DFT steps.
$\begin{array}{c} \textbf{propagator} \\ integer \end{array}$	default = 1 RT-TD-DFT propagator (1 = Verlet, 2 = Magnus).
$egin{aligned} \mathbf{NBCH} \ integer \end{aligned}$	default = 10 Number of $[\rho]$. Fock ⁿ commutators in Magnus.
${f tdrestart} \ logical$	$default = .false.$ Reads an input restart for TD (named td_in.restart).
$rac{\mathbf{td}_{-}\mathbf{rst}_{-}\mathbf{freq}}{integer}$	$default = 500$ Write the TD restart every td_rst_freq steps.
$egin{aligned} \mathbf{td}_{-}\mathbf{do}_{-}\mathbf{pop} \ integer \end{aligned}$	default=0 Number of step stride in which the pop will be written. (0 means it is never written).
$egin{aligned} \mathbf{writeDens} \ logical \end{aligned}$	default = .false. Writes electronic density to an output file after having finished the calculations.

Table 7.10: Real Time TD-DFT

Keywords - Transport

Variable	Description
$rac{\mathbf{transport_calc}}{logical}$	$default = .false. \ { m TODO}.$
$egin{array}{c} \mathbf{generate_rho0} \ logical \end{array}$	$default = .false. \ { m TODO}.$
$egin{aligned} \mathbf{gate_field} \ logical \end{aligned}$	$default = .false. \ { m TODO}.$
$rac{ extbf{driving_rate}}{ ext{double precision}}$	$default = UNSET \ { m TODO}.$
$egin{aligned} \mathbf{pop_drive} \ integer \end{aligned}$	$default = UNSET \ { m TODO}.$
$egin{array}{c} \mathbf{save_charge_freq} \ integer \end{array}$	$default = UNSET \ { m TODO}.$
$egin{array}{c} \mathbf{nbias} \ integer \end{array}$	$default = UNSET \ { m TODO}.$

Table 7.11: Transport

Keywords - Ehrenfest

Variable	Description
$rac{ extbf{ndyn_steps}}{integer}$	default = 0 Number of nuclear movement steps.
$egin{aligned} \mathbf{rsto_nfreq} \ integer \end{aligned}$	default = 0 Frequency (in steps) in which the restart is printed. (A value of 0 means only written in the end)
$egin{array}{c} \mathbf{rsto_saves} \ logical \end{array}$	$default = .false. \ { m TODO}.$
$egin{array}{c} \mathbf{rsti_loads} \ logical \end{array}$	$default = .false. \ { m TODO}.$
$rac{ extbf{nullify}_{-} extbf{forces}}{logical}$	default = .false. Returns 0 for all forces to the MD engine.
$rac{ extbf{nullify}_{-} extbf{forces}}{logical}$	default = .false. Returns 0 for all forces to the MD engine.
$egin{array}{c} \mathbf{eefld_on} \ logical \end{array}$	$default = .false. \ { m TODO}.$
$egin{array}{c} \mathbf{eefld_timegih} \\ logical \end{array}$	$default = .false. \ { m TODO}.$
$egin{array}{c} \mathbf{eefld_timegfh} \ logical \end{array}$	$default = .false. \ { m TODO}.$
eefld_ampx/ampy/a dn p n u $lt=2.D$ -5 $double\ precision$ TODO.	
$egin{array}{c} \mathbf{eefld_timeamp} \ double\ precision \end{array}$	$default = 2.D$ -5 ${ m TODO}$.
eefld_timepos double precision	default = 2.D-5 TODO.

Table 7.12: Ehrenfest

Variable	Description
eefld_wavelen double precision	$default = 2.D ext{-}5 \ ext{TODO}.$

Table 7.12: Ehrenfest

Keywords - CubeGen

Variable	Description
$egin{array}{c} \mathbf{cube_dens} \ logical \end{array}$	default = .false. Prints the electronic density.
$egin{aligned} \mathbf{cubeGen_only} \ logical \end{aligned}$	default = .false. Avoid running SCF, only do cube Gen from a restart.
$egin{array}{c} \mathbf{cube_res} \ integer \end{array}$	default = 40 Number of voxels per dimension (resolution).
$egin{array}{c} \mathbf{cube_sel} \ integer \end{array}$	default = 0 Select only a particular orbital for printing $(0 = all)$.
$rac{ extbf{cube_dens_file}}{character*20}$	default = 'dens.cube' File containing the electronic density.
$egin{array}{c} \mathbf{cube_orb} \ logical \end{array}$	default = .false. Prints orbital shapes.
$egin{array}{c} \mathbf{cube_sqrt_orb} \ logical \end{array}$	default = .false. Prints the orbitals' square root.
cube_orb_file character*20	default = 'orb.cube' File containing the orbital shapes.
$egin{array}{c} \mathbf{cube_elec} \ logical \end{array}$	default = .false. Prints the electric field.
${f cube_elec_file} \ character*20$	default = 'field.cube' File containing the electrical field.

Table 7.13: CubeGen