
MRChem Documentation

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MRChem is a numerical real-space code for molecular electronic structure calculations within the self-consistent field (SCF) approximations of quantum chemistry (Hartree-Fock and Density Functional Theory). The code is divided in two main parts: the [MultiResolution Computation Program Package](#) (MRCPP), which is a general purpose numerical mathematics library based on multiresolution analysis and the multiwavelet basis which provide low-scaling algorithms as well as rigorous error control in numerical computations, and the MultiResolution Chemistry (MRChem) program that uses the functionalities of MRCPP for computational chemistry applications.

The code is being developed at the [Hylleraas Centre for Quantum Molecular Sciences](#) at [UiT - The Arctic University of Norway](#).

The code is under active development, and the latest stable releases as well as development versions can be found on [GitHub](#).

**CHAPTER
ONE**

FEATURES IN MRCHEM-1.1:

- **Wave functions:**

- **Kohn-Sham DFT**
 - * Spin-polarized
 - * Spin-unpolarized
 - * LDA, GGA and hybrid functionals
- **Hartree-Fock**
 - * Restricted closed-shell
 - * Unrestricted
- **Explicit external fields**
 - * Electric field
- **Solvent effects**
 - * Cavity-free PCM
 - * Poisson-Boltzmann PCM
 - * Linearized Poisson-Boltzmann PCM

- **Properties:**

- Ground state energy
- Dipole moment
- Quadrupole moment
- Polarizability
- Magnetizability
- NMR shielding constant
- Geometric derivative

- **Parallel implementation:**

- Shared memory (OpenMP): ~20 cores
- Distributed memory (MPI): ~1000 procs
- Hybrid scheme (MPI + OpenMP): ~10 000 cores

- **Current size limitations:**

- ~2000 orbitals on ~100 high-end compute nodes (128 core/256GiB mem)
- ~100 orbitals on a single high-memory (1TB) compute node

CHAPTER
TWO

UPCOMING FEATURES:

- **Wave functions:**
 - Meta-GGAs
 - ZORA Hamiltonian
 - Periodic Boundary Conditions
 - External magnetic field
- **Properties:**
 - Optical rotation
 - Spin-spin coupling constant
 - Hyperfine coupling constant
 - Magnetically induced currents
 - Hyperpolarizability
 - Geometry optimization
- **Performance:**
 - Reduced memory footprint
 - Improved DFT scaling and performance

2.1 Installation

2.1.1 Build prerequisites

- Python-3.9 (or later)
- CMake-3.14 (or later)
- GNU-5.4 or Intel-17 (or later) compilers (C++14 standard)

Hint: We have collected the recommended modules for the different Norwegian HPC systems under `tools/<machine>.env`. These files can be sourced in order to get a working environment on the respective machines, and may also serve as a guide for other HPC systems.

C++ dependencies

The MRChem program depends on the following C++ libraries:

- Input handling: [nlohmann/json-3.6](#)
- Multiwavelets: [MRCPP-1.4](#)
- Linear algebra: [Eigen-3.4](#)
- DFT functionals: [XCFun-2.0](#)

All these dependencies will be downloaded automatically at configure time by CMake, but can also be linked manually by setting the variables:

```
MRCPP_DIR=<path_to_mrcpp>/share/cmake/MRCPP  
XCFun_DIR=<path_to_xcfun>/share/cmake/XCFun  
Eigen3_DIR=<path_to_eigen3>/share/eigen3/cmake  
nlohmann_json_DIR=<path_to_nlohmann_json>
```

Python dependencies

Users only need a Python3 interpreter, which is used for configuration (`setup` script) as well as launching the program (`mrchem` script).

Developers will need some extra Python packages to update the input parser and build the documentation locally with Sphinx.

We **strongly** suggest not to install these Python dependencies globally, but rather to use a local virtual environment. We provide a `Pipfile` for specifying the Python dependencies. We recommend using [Pipenv](#), since it manages virtual environment and package installation seamlessly. After installing it with your package manager, run:

```
$ pipenv install --dev
```

to create a virtual environment with all developer packages installed.

The environment can be activated with:

```
$ pipenv shell
```

Alternatively, any Python command can be run within the virtual environment by doing:

```
$ pipenv run python -c "print('Hello, world')"
```

2.1.2 Obtaining and building the code

The latest development version of MRChem can be found on the `master` branch on GitHub:

```
$ git clone https://github.com/MRChemSoft/mrchem.git
```

The released versions can be found from Git tags vX.Y.Z under the `release/X.Y` branches in the same repository, or a zip file can be downloaded from [Zenodo](#).

By default, all dependencies will be **fetched** at configure time if they are not already available.

Configure

The `setup` script will create a directory called `<build-dir>` and run CMake. There are several options available for the setup, the most important being:

```
--cxx=<CXX>
    C++ compiler [default: g++]

--omp
    Enable OpenMP parallelization [default: False]

--mpi
    Enable MPI parallelization [default: False]

--type=<TYPE>
    Set the CMake build type (debug, release, relwithdeinfo, minsizerel) [default: release]

--prefix=<PATH>
    Set the install path for make install [default: '/usr/local']

--cmake-options=<STRING>
    Define options to CMake [default: '']

-h --help
    List all options
```

The code can be built with four levels of parallelization:

- no parallelization
- only shared memory (OpenMP)
- only distributed memory (MPI)
- hybrid OpenMP + MPI

Note: In practice we recommend the **shared memory version** for running on your personal laptop/workstation, and the **hybrid version** for running on a HPC cluster. The serial and pure MPI versions are only useful for debugging.

The default build is *without* parallelization and using GNU compilers:

```
$ ./setup --prefix=<install-dir> <build-dir>
```

To use Intel compilers you need to specify the `--cxx` option:

```
$ ./setup --prefix=<install-dir> --cxx=icpc <build-dir>
```

To build the code with shared memory (OpenMP) parallelization, add the `--omp` option:

```
$ ./setup --prefix=<install-dir> --omp <build-dir>
```

To build the code with distributed memory (MPI) parallelization, add the `--mpi` option *and* change to the respective MPI compilers (`--cxx=mpicxx` for GNU and `--cxx=mpiicpc` for Intel):

```
$ ./setup --prefix=<install-dir> --omp --mpi --cxx=mpicxx <build-dir>
```

When dependencies are fetched at configuration time, they will be downloaded into `<build-dir>/_deps`. For the example of MRCPP, sources are saved into the folders `<build-dir>/_deps/mrcpp_sources-src` and built into `<build-dir>/_deps/mrcpp_sources-build`.

Note: If you compile the MRCPP library manually as a separate project, the level of parallelization **must be the same** for MRCPP and MRChem. Similar options apply for the MRCPP setup, see mrcpp.readthedocs.io.

Build

If the CMake configuration is successful, the code is compiled with:

```
$ cd <build-dir>
$ make
```

Test

A test suite is provided to make sure that everything compiled properly. To run a collection of small unit tests:

```
$ cd <build-dir>
$ ctest -L unit
```

To run a couple of more involved integration tests:

```
$ cd <build-dir>
$ ctest -L integration
```

Install

After the build has been verified with the test suite, it can be installed with the following command:

```
$ cd <build-dir>
$ make install
```

This will install *two* executables under the <install-path>:

```
<install-path>/bin/mrchem      # Python input parser and launcher
<install-path>/bin/mrchem.x    # MRChem executable
```

Please refer to the [User's Manual](#) for instructions for how to run the program.

Hint: We have collected scripts for configure and build of the hybrid OpenMP + MPI version on the different Norwegian HPC systems under tools/<machine>.sh. These scripts will build the current version under build-{\$version}, run the unit tests and install under install-{\$version}, e.g. to build version v1.0.0 on Fram:

```
$ cd mrchem
$ git checkout v1.0.0
$ tools/fram.sh
```

The configure step requires internet access, so the scripts must be run on the login nodes, and it will run on a single core, so it might take some minutes to complete. The scripts will *not* install the [Python dependencies](#), so this must be done manually in order to run the code.

2.2 User's Manual

The MRChem program comes as two executables:

```
<install-path>/bin/mrchem           # Python input parser and launcher
<install-path>/bin/mrchem.x         # MRChem main executable
```

where the former is a Python script that reads and validates the *user input file* and produces a new *program input file* which is then passed as argument to the latter, which is the actual C++ executable.

The input and output of the program is thus organized as *three* separate files:

File extension	Description	Format
.inp	User input file	GETKW/JSON
.json	Program input/output	JSON
.out	User output file	Text

The name of the user input file can be anything, as long as it has the .inp extension, and the corresponding .json and .out files will get the same name prefix. The JSON program file will get both an "input" and an "output" section. This "input" section is rather detailed and contains very implementation specific keywords, but it is automatically generated by the mrchem script, based on the more generic keywords of the user input file. The mrchem script will further launch the mrchem.x main executable, which will produce the text output file as well as the "output" section of the JSON in/out file. The contents of all these files will be discussed in more detail in the sections below.

2.2.1 Running the program

In the following we will assume to have a valid user input file for the water molecule called h2o.inp, e.g. like this

```
world_prec = 1.0e-4

WaveFunction {
    method = B3LYP
}

Molecule {
$coords
O  0.0000  0.000 -0.125
H -1.4375  0.000  1.025
H  1.4375  0.000  1.025
$end
}
```

To run the calculation, pass the file name (without extension) as argument to the mrchem script (make sure you understand the difference between the .inp, .json and .out file, as described in the previous section):

```
$ mrchem h2o
```

This will under the hood actually do the following two steps:

```
$ mrchem h2o.inp > h2o.json
$ mrchem.x h2o.json > h2o.out
```

The first step includes input validation, which means that everything that passes this step is a well-formed computation.

Dry-running the input parser

The execution of the two steps above can be done separately by dry-running the parser script:

```
$ mrchem --dryrun h2o
```

This will run only the input validation part and generate the `h2o.json` program input, but it will *not* launch the main executable `mrchem.x`. This can then be done manually in a subsequent step by calling:

```
$ mrchem.x h2o.json
```

This separation can be useful for instance for developers or advanced users who want to change some automatically generated input values before launching the actual program, see *Input schema*.

Printing to standard output

By default the program will write to the text output file (`.out` extension), but if you rather would like it printed in the terminal you can add the `--stdout` option (then no text output file is created):

```
$ mrchem --stdout h2o
```

Reproducing old calculations

The JSON in/out file acts as a full record of the calculation, and can be used to reproduce old results. Simply pass the JSON file once more to `mrchem.x`, and the "output" section will be overwritten:

```
$ mrchem.x h2o.json
```

User input in JSON format

The user input file can be written in JSON format instead of the standard syntax which is described in detail below. This is very convenient if you have for instance a Python script to generate input files. The water example above in JSON format reads (the `coords` string is not very elegant, but unfortunately that's just how JSON works...):

```
{
  "world_prec": 1.0e-4,
  "WaveFunction": {
    "method": "B3LYP"
  },
  "Molecule": {
    "coords": "0  0.0000  0.000 -0.125\nH -1.4375  0.000  1.025\nH  1.4375  0.000  1.025\nH"
  }
}
```

which can be passed to the input parser with the `--json` option:

```
$ mrchem --json h2o
```

Note: A *user input file* in JSON format must **NOT** be confused with the JSON in/out file for the `mrchem.x` program. The file should still have a `.inp` extension, and contain all the same keywords which have to be validated and translated by the `mrchem` script into the `.json` *program input file*.

Parallel execution

The MRChem program comes with support for both shared memory and distributed memory parallelization, as well as a hybrid combination of the two. In order to activate these capabilities, the code needs to be compiled with OpenMP and/or MPI support (`--omp` and/or `--mpi` options to the CMake `setup` script, see [Installation](#) instructions).

Shared memory OpenMP

For the shared memory part, the program will automatically pick up the number of threads from the environment variable `OMP_NUM_THREADS`. If this variable is *not* set it will usually default to the maximum available. So, to run the code on 16 threads (all sharing the same physical memory space):

```
$ OMP_NUM_THREADS=16 mrchem h2o
```

Note that this is the number of threads will be set by `OMP_NUM_THREADS` only if the code is compiled without MPI support, see below.

Distributed memory MPI

In order to run a program in an MPI parallel fashion, it must be executed with an MPI launcher like `mpirun`, `mpiexec`, `srun`, etc. Note that it is only the main executable `mrchem.x` that should be launched in parallel, **not** the `mrchem` input parser script. This can be achieved *either* by running these separately in a dry-run (here two MPI processes):

```
$ mrchem --dryrun h2o
$ mpirun -np 2 mrchem.x h2o.json
```

or in a single command by passing the launcher string as argument to the parser:

```
$ mrchem --launcher="mpirun -np 2" h2o
```

This string can contain any argument you would normally pass to `mpirun` as it will be literally prepended to the `mrchem.x` command when the `mrchem` script executes the main program.

Hint: For best performance, it is recommended to use shared memory *within* each **NUMA** domain (usually one per socket) of your CPU, and MPI across NUMA domains and ultimately machines. Ideally, the number of OpenMP threads should be between 8-20. E.g. on hardware with two sockets of 16 cores each, scale the number of MPI processes by the size of the molecule, typically one process per ~5 orbitals or so (and definitely not *more* than one process per orbital). The actual number of threads will be set automatically regardless of the value of `OMP_NUM_THREADS`.

Job example (Betzy)

This job will use 4 compute nodes, with 12 MPI processes on each, and the MPI process will use up to 15 OpenMP threads. 4 MPI process per node are used for the “Bank”. The Bank processes are using only one thread, therefore there is in practice no overallocation. It is however important that `bank_size` is set to be at least $4 \times 4 = 16$ (it is by default set, correctly, to one third of total MPI size, i.e. $4 \times 12 / 3 = 16$). It would also be possible to set 16 tasks per node, and set the bank size parameter accordingly to $8 \times 4 = 32$. The flags are optimized for the OpenMPI (foss) library on Betzy (note that H2O is a very small molecule for such setup!).

```
#!/bin/bash -l
#SBATCH --nodes=4
#SBATCH --tasks-per-node=12

export UCX_LOG_LEVEL=ERROR

~/my_path/to/mrchem --launcher='mpirun --rank-by node --map-by socket --bind-to numa' h2o
```

--rank-by node

Tells the system to place the first MPI rank on the first node, the second MPI rank on the second node, until the last node, then start at the first node again.

--map-by socket

Tells the system to map (group) MPI ranks according to socket before distribution between nodes. This will ensure that for example two bank cores will access different parts of memory and be placed as the 16th thread of a numa group.

--bind-to numa

Tells the system to bind cores to one NUMA (Non Uniform Memory Access) group. On Betzy memory configuration groups cores by groups of 16, with cores in the same group having the same access to memory (other cores will have access to that part of the memory too, but slower). That means that a process will only be allowed to use one of the 16 cores of the group. (The operating system may change the core assigned to a thread/process and, without precautions, it may be assigned to any other core, which would result in much reduced performance). The 16 cores of the group may then be used by the threads initiated by that MPI process.

Advanced option: Alternatively one can get full control of task placement using the Slurm workload manager by replacing `mpirun` with `srun` and setting explicit CPU masks as:

--distribution=cyclic:cyclic The first cyclic will put the first rank on the first node, the second rank on the second node etc. The second cyclic distribute the ranks withing the nodes.

More examples can be found in the [mrchem-examples](#) repository on GitHub.

Parallel pitfalls

Warning: Parallel program execution is not a black box procedure, and the behavior and efficiency of the run depends on several factors, like hardware configuration, operating system, compiler type and flags, libraries for OpenMP and MPI, type of queing system on a shared cluster, etc. Please make sure that the program runs correctly on *your* system and is able to utilize the computational resources before commencing production calculations.

Typical pitfalls for OpenMP

- Not compiling with correct OpenMP support.
- Not setting number of threads correctly.
- **Hyper-threads:** the round-robin thread distribution might fill all hyper-threads on each core before moving on to the next physical core. In general we discourage the use of hyper-threads, and recommend a single thread per physical core.
- **Thread binding:** all threads may be bound to the same core, which means you can have e.g. 16 threads competing for the limited resources available on this single core (typically two hyper-threads) while all other cores are left idle.

Typical pitfalls for MPI

- Not compiling with the correct MPI support.
- Default launcher options might not give correct behavior.
- **Process binding:** if a process is bound to a core, then all its spawned threads will also be bound to the same core. In general we recommend binding to socket/NUMA.
- **Process distribution:** in a multinode setup, all MPI processes might land on the same machine, or the round-robin procedure might count each core as a separate machine.

How to verify a parallel MRChem run

- In the printed output, verify that MRCPP has actually been compiled with correct support for MPI and/or OpenMP:

```
-----  
MRCPP version      : 1.2.0  
Git branch         : master  
Git commit hash    : 686037cb78be601ac58b  
Git commit author  : Stig Rune Jensen  
Git commit date    : Wed Apr 8 11:35:00 2020 +0200  
  
Linear algebra     : EIGEN v3.3.7  
Parallelization    : MPI/OpenMP  
-----
```

- In the printed output, verify that the correct number of processes and threads has been detected:

MPI processes	:	(no bank)	2
OpenMP threads	:		16
Total cores	:		32

- Monitor your run with `top` to see that you got the expected number of `mrchem.x` processes (MPI), and that they actually run at the expected CPU percentage (OpenMP):

PID	USER	PR	NI	VIRT	RES	SHR	S	%CPU	%MEM	TIME+	COMMAND
9502	stig	25	5	489456	162064	6628	R	1595,3	2,0	0:14.50	mrchem.x
9503	stig	25	5	489596	162456	6796	R	1591,7	2,0	0:14.33	mrchem.x

- Monitor your run with `htop` to see which core/hyper-thread is being used by each process. This is very useful to get the correct binding/pinning of processes and threads. In general you want one threads per core, which means that every other hyper-thread should remain idle. In a hybrid MPI/OpenMP setup it is rather common that each MPI process becomes bound to a single core, which means that all threads spawned by this process will occupy the same core (possibly two hyper-threads). This is then easily detected with `htop`.
- Perform dummy executions of your parallel launcher (`mpirun`, `srun`, etc) to check whether it picks up the correct parameters from the resource manager on your cluster (SLURM, Torque, etc). You can then for instance report bindings and host name for each process:

```
$ mpirun --print-rank-map hostname
```

Play with the launcher options until you get it right. Note that Intel and OpenMPI have slightly different options for their `mpirun` and usually different behavior. Beware that the behavior can also change when you move from single- to multinode execution, so it is in general not sufficient to verify you runs on a single machine.

- Perform a small scaling test on e.g. 1, 2, 4 processes and/or 1, 2, 4 threads and verify that the total computation time is reduced as expected (don't expect 100% efficiency at any step).

2.2.2 User input file

The input file is organized in sections and keywords that can be of different type. Input keywords and sections are **case-sensitive**, while *values* are **case-insensitive**.

```
Section {
    keyword_1 = 1                      # int
    keyword_2 = 3.14                     # float
    keyword_3 = [1, 2, 3]                 # int array
    keyword_4 = foo                      # string
    keyword_5 = true                     # boolean
}
```

Valid options for booleans are `true/false`, `on/off` or `yes/no`. Single word strings can be given without quotes (be careful of special characters, like slashes in file paths). A complete list of available input keywords can be found in the [User input reference](#).

Top section

The main input section contain four keywords: the relative precision ϵ_{rel} that will be guaranteed in the calculation and the size, origin and unit of the computational domain. The top section is not specified by name, just write the keywords directly, e.g

```
world_prec = 1.0e-5          # Overall relative precision
world_size = 5                # Size of domain 2^{world_size}
world_unit = bohr             # Global length unit
world_origin = [0.0, 0.0, 0.0] # Global gauge origin
```

The relative precision sets an upper limit for the number of correct digits you are expected to get out of the computation (note that $\epsilon_{rel} = 10^{-6}$ yields μ Ha accuracy for the hydrogen molecule, but only mHa accuracy for benzene).

The computational domain is always symmetric around the origin, with *total* size given by the `world_size` parameter as $[2^n]^3$, e.i. `world_size = 5` gives a domain of $[-16, 16]^3$. Make sure that the world is large enough to allow the molecular density to reach zero on the boundary. The `world_size` parameter can be left out, in which case the size will be estimated based on the molecular geometry. The `world_unit` relates to **all** coordinates given in the input file and can be one of two options: `angstrom` or `bohr`.

Note: The `world_size` will be only approximately scaled by the `angstrom` unit, by adding an extra factor of 2 rather than the appropriate factor of ~1.89. This means that e.g. `world_size = 5` ($[-16, 16]^3$) with `world_unit = angstrom` will be translated into $[-32, 32]^3$ bohrs.

Precisions

MRChem uses a smoothed nuclear potential to avoid numerical problems in connection with the $Z/|r - R|$ singularity. The smoothing is controlled by a single parameter `nuc_prec` that is related to the expected error in the energy due to the smoothing. There are also different precision parameters for the *construction* of the Poisson and Helmholtz integral operators.

```
Precisions {
    nuclear_prec = 1.0e-6          # For construction of nuclear potential
    poisson_prec = 1.0e-6          # For construction of Poisson operators
    helmholtz_prec = 1.0e-6        # For construction of Helmholtz operators
}
```

By default, all precision parameters follow `world_prec` and usually don't need to be changed.

Printer

This section controls the format of the printed output file (.out extension). The most important option is the `print_level`, but it also gives options for number of digits in the printed output, as well as the line width (defaults are shown):

```
Printer {
    print_level = 0                # Level of detail in the printed output
    print_width = 75               # Line width (in characters) of printed output
    print_prec = 6                  # Number of digits in floating point output
}
```

Note that energies will be printed with *twice* as many digits. Available print levels are:

- `print_level=-1` no output is printed
- `print_level=0` prints mainly properties
- `print_level=1` adds timings for individual steps
- `print_level=2` adds memory and timing information on OrbitalVector level
- `print_level=3` adds details for individual terms of the Fock operator
- `print_level=4` adds memory and timing information on Orbital level
- `print_level>=5` adds debug information at MRChem level
- `print_level>=10` adds debug information at MRCPP level

MPI

This section defines some parameters that are used in MPI runs (defaults shown):

```
MPI {  
    bank_size = -1                      # Number of processes used as memory bank  
    omp_threads = -1                     # Number of omp threads to use  
    numerically_exact = false           # Guarantee MPI invariant results  
    share_nuclear_potential = false     # Use MPI shared memory window  
    share_coulomb_potential = false      # Use MPI shared memory window  
    share_xc_potential = false          # Use MPI shared memory window  
}
```

The memory bank will allow larger molecules to get though if memory is the limiting factor, but it will be slower, as the bank processes will not take part in any computation. For calculations involving exact exchange (Hartree-Fock or hybrid DFT functionals) a memory bank is **required** whenever there's more than one MPI process. A negative bank size will set it automatically based on the number of available processes. For pure DFT functionals on smaller molecules it is likely more efficient to set `bank_size = 0`, otherwise it's recommended to use the default. If a particular calculation runs out of memory, it might help to increase the number of bank processes from the default value.

The number of threads to use in OpenMP can be forced using the `omp_threads` flag. For MPI runs, it is strongly advised to leave the default, as the optimal value can be difficult to guess. The environment variable `OMP_NUM_THREADS` is not used for MPI runs.

The `numerically_exact` keyword will trigger algorithms that guarantee that the computed results are invariant (within double precision) with respect to the number of MPI processes. These exact algorithms require more memory and are thus not default. Even when the numbers are *not* MPI invariant they should be correct and identical within the chosen `world_prec`.

The `share_potential` keywords are used to share the memory space for the particular functions between all processes located on the same physical machine. This will save memory but it might slow the calculation down, since the shared memory cannot be “fast” memory (NUMA) for all processes at once.

Basis

This section defines the polynomial MultiWavelet basis

```
Basis {
    type = Interpolating          # Legendre or Interpolating
    order = 7                      # Polynomial order of MW basis
}
```

The MW basis is defined by the polynomial order k , and the type of scaling functions: Legendre or Interpolating polynomials (in the current implementation it doesn't really matter which type you choose). Note that increased precision requires higher polynomial order (use e.g. $k = 5$ for $\epsilon_{rel} = 10^{-3}$, and $k = 13$ for $\epsilon_{rel} = 10^{-9}$, and interpolate in between). If the `order` keyword is left out it will be set automatically according to

$$k = -1.5 * \log_{10}(\epsilon_{rel})$$

The Basis section can usually safely be omitted in the input.

Molecule

This input section specifies the geometry (given in `world_unit` units), charge and spin multiplicity of the molecule, e.g. for water (coords must be specified, otherwise defaults are shown):

```
Molecule {
    charge = 0                      # Total charge of molecule
    multiplicity = 1                 # Spin multiplicity
    translate = false                # Translate CoM to world_origin
$coords
O  0.0000    0.0000    0.0000      # Atomic symbol and coordinate
H  0.0000    1.4375    1.1500      # Atomic symbol and coordinate
H  0.0000   -1.4375    1.1500      # Atomic symbol and coordinate
$end
}
```

Since the computational domain is always cubic and symmetric around the origin it is usually a good idea to `translate` the molecule to the origin (as long as the `world_origin` is the true origin).

WaveFunction

Here we give the wavefunction method, environment used (for solvent models) and whether we run spin restricted (alpha and beta spins are forced to occupy the same spatial orbitals) or not (method must be specified, otherwise defaults are shown):

```
WaveFunction {
    method = <wavefunction_method>      # Core, Hartree, HF or DFT
    restricted = true                    # Spin restricted/unrestricted
    environment = pcm                  # Environment (pcm, pcm-pb, pcm-lpb) defaults to
    ↵none
}
```

There are currently four methods available: Core Hamiltonian, Hartree, Hartree-Fock (HF) and Density Functional Theory (DFT). When running DFT you can *either* set one of the default functionals in this section (e.g. `method = B3LYP`), *or* you can set `method = DFT` and specify a “non-standard” functional in the separate DFT section (see below). See [User input reference](#) for a list of available default functionals.

The solvent model implemented is a cavity free PCM, described in [gerez2023]. In this model we have implemented the Generalized Poisson equation solver, keyword `pcm`, a Poisson-Boltzmann solver, keyword `pcm-pb` and a Linearized Poisson-Boltzmann solver, keyword `pcm-lpb`. Further details for the calculation have to be included in the PCM section, see :ref: *User input reference* for details.

Note: Restricted open-shell wavefunctions are not supported.

DFT

This section can be omitted if you are using a default functional, see above. Here we specify the exchange-correlation functional used in DFT (functional names must be specified, otherwise defaults are shown)

```
DFT {
    spin = false                                # Use spin-polarized functionals
    density_cutoff = 0.0                         # Cutoff to set XC potential to zero
$functionals
<func1>    1.0                                # Functional name and coefficient
<func2>    1.0                                # Functional name and coefficient
$end
}
```

You can specify as many functionals as you want, and they will be added on top of each other with the given coefficient. Both exchange and correlation functionals must be set explicitly, e.g. `SLATERX` and `VWN5C` for the standard LDA functional. For hybrid functionals you must specify the amount of exact Hartree-Fock exchange as a separate functional `EXX` (`EXX 0.2` for B3LYP and `EXX 0.25` for PBE0 etc.). Option to use spin-polarized functionals or not. Unrestricted calculations will use spin-polarized functionals by default. The XC functionals are provided by the `XCFun` library.

Properties

Specify which properties to compute. By default, only the ground state SCF energy as well as orbital energies will be computed. Currently the following properties are available (all but the dipole moment are `false` by default)

```
Properties {
    dipole_moment = true                         # Compute dipole moment
    quadrupole_moment = false                    # Compute quadrupole moment
    polarizability = false                       # Compute polarizability
    magnetizability = false                      # Compute magnetizability
    nmr_shielding = false                        # Compute NMR shieldings
    geometric_derivative = false                 # Compute geometric derivative
    plot_density = false                         # Plot converged density
    plot_orbitals = []                           # Plot converged orbitals
}
```

Some properties can be further specified in dedicated sections.

Warning: The computation of the molecular gradient suffers greatly from numerical noise. The code replaces the nucleus-electron attraction with a smoothed potential. This can only partially recover the nuclear cusps, even with tight precision. The molecular gradient is only suited for use in geometry optimization of small molecules and with tight precision thresholds.

Polarizability

The polarizability can be computed with several frequencies (by default only static polarizability is computed):

```
Polarizability {
    frequency = [0.0, 0.0656]           # List of frequencies to compute
}
```

NMRSshielding

For the NMR shielding we can specify a list of nuclei to compute (by default all nuclei are computed):

```
NMRSshielding {
    nuclear_specific = false            # Use nuclear specific perturbation operator
    nucleus_k = [0,1,2]                 # List of nuclei to compute (-1 computes all)
}
```

The `nuclear_specific` keyword triggers response calculations using the nuclear magnetic moment operator instead of the external magnetic field. For small molecules this is not recommended since it requires a separate response calculation for each nucleus, but it might be beneficial for larger systems if you are interested only in a single shielding constant. Note that the components of the *perturbing* operator defines the *row* index in the output tensor, so `nuclear_specific = true` will result in a shielding tensor which is the transpose of the one obtained with `nuclear_specific = false`.

Plotter

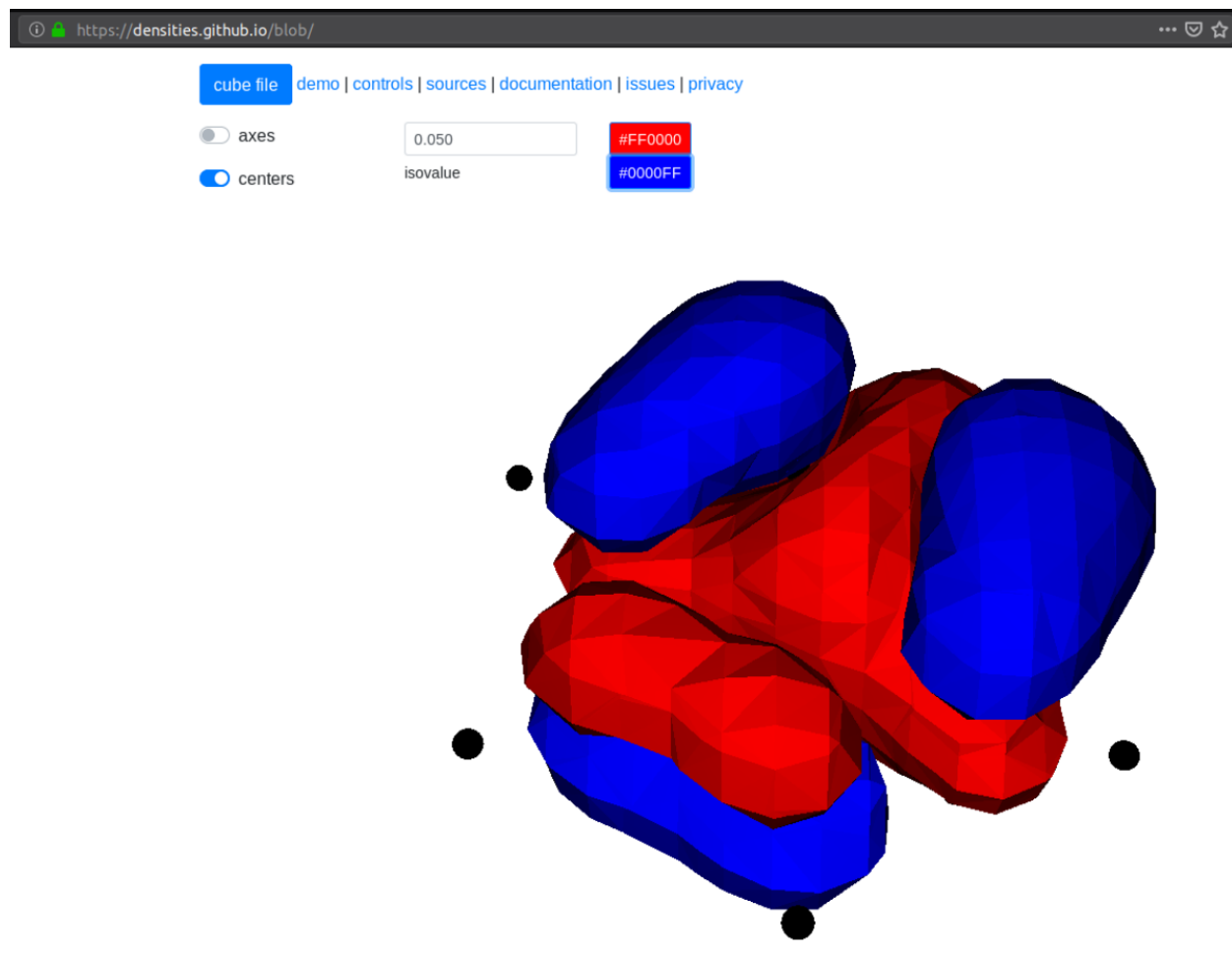
The `plot_density` and `plot_orbitals` properties will use the Plotter section to specify the parameters of the plots (by default you will get a cube plot on the unit cube):

```
Plotter {
    path = plots                         # File path to store plots
    type = cube                           # Plot type (line, surf, cube)
    points = [20, 20, 20]                  # Number of grid points
    O = [-4.0,-4.0,-4.0]                  # Plot origin
    A = [8.0, 0.0, 0.0]                   # Boundary vector
    B = [0.0, 8.0, 0.0]                   # Boundary vector
    C = [0.0, 0.0, 8.0]                   # Boundary vector
}
```

The plotting grid is computed from the vectors O, A, B and C in the following way:

1. `line` plot: along the vector A starting from O, using `points[0]` number of points.
2. `surf` plot: on the area spanned by the vectors A and B starting from O, using `points[0]` and `points[1]` points in each direction.
3. `cube` plot: on the volume spanned by the vectors A, B and C starting from O, using `points[0]`, `points[1]` and `points[2]` points in each direction.

The above example will plot on a 20x20x20 grid in the volume $[-4,4]^3$, and the generated files (e.g. `plots/phi_1_re.cube`) can be viewed directly in a web browser by `blob`, like this benzene orbital:



SCF

This section specifies the parameters for the SCF optimization of the ground state wavefunction.

SCF solver

The optimization is controlled by the following keywords (defaults shown):

```
SCF {
    run = true                                # Run SCF solver
    kain = 5                                  # Length of KAIN iterative subspace
    max_iter = 100                            # Maximum number of SCF iterations
    rotation = 0                               # Iterations between diagonalize/localize
    localize = false                           # Use canonical or localized orbitals
    start_prec = -1.0                          # Dynamic precision, start value
    final_prec = -1.0                          # Dynamic precision, final value
    orbital_thrs = 10 * world_prec           # Convergence threshold orbitals
    energy_thrs = -1.0                         # Convergence threshold energy
}
```

If `run = false` no SCF is performed, and the properties are computed directly on the initial guess wavefunction.

The `kain` (Krylov Accelerated Inexact Newton) keyword gives the length of the iterative subspace accelerator (similar to DIIS). The `rotation` keyword gives the number of iterations between every orbital rotation, which can be either localization or diagonalization, depending on the `localize` keyword. The first two iterations in the SCF are always rotated, otherwise it is controlled by the `rotation` keyword (usually this is not very important, but sometimes it fails to converge if the orbitals drift too far away from the localized/canonical forms).

The dynamic precision keywords control how the numerical precision is changed throughout the optimization. One can choose to use a lower `start_prec` in the first iterations which is gradually increased to `final_prec` (both are equal to `world_prec` by default). Note that lower initial precision might affect the convergence rate.

In general, the important convergence threshold is that of the orbitals, and by default this is set one order of magnitude higher than the overall `world_prec`. For simple energy calculations, however, it is not necessary to converge the orbitals this much due to the quadratic convergence of the energy. This means that the number of correct digits in the total energy will be saturated well before this point, and one should rather use the `energy_thrs` keyword in this case in order to save a few iterations.

Note: It is usually not feasible to converge the orbitals *beyond* the overall precision `world_prec` due to numerical noise.

Initial guess

Several types of initial guess are available:

- `core` and `sad` requires no further input and computes guesses from scratch.
- `chk` and `mw` require input files from previous MW calculations.
- `cube` requires input files computed from other sources.

The `core` and `sad` guesses are computed by diagonalizing the Hamiltonian matrix using a Core or Superposition of Atomic Densities (SAD) Hamiltonian, respectively. The matrix is constructed in a small AO basis with a given “zeta quality”, which should be added as a suffix in the keyword. Available AO bases are hydrogenic orbitals of single `sz`, double `dz`, triple `tz` and quadruple `qz` zeta size.

The SAD guess can also be computed in a small GTO basis (3-21G), using the guess type `sad_gto`. In this case another input keyword `guess_screen` becomes active for screening in the MW projection of the Gaussians. The screening value is given in standard deviations. Such screening will greatly improve the efficiency of the guess for large systems. It is, however, not recommended to reduce the value much below 10 StdDevs, as this will have the *opposite* effect on efficiency due to introduction of discontinuities at the cutoff point, which leads to higher grid refinement. `sad_gto` is usually the preferred guess both for accuracy and efficiency, and is thus the default choice.

The `core` and `sad` guesses are fully specified with the following keywords (defaults shown):

```
SCF {
    guess_prec = 1.0e-3                      # Numerical precision used in guess
    guess_type = sad_gto                      # Type of initial guess (chk, mw, cube, core_XX, ...
    ↪sad_XX)
    guess_screen = 12.0                        # Number of StdDev before a GTO is set to zero_
    ↪(sad_gto)
}
```

Checkpointing

The program can dump checkpoint files at every iteration using the `write_checkpoint` keyword (defaults shown):

```
SCF {
    path_checkpoint = checkpoint          # Path to checkpoint files
    write_checkpoint = false              # Save checkpoint files every iteration
}
```

This allows the calculation to be restarted in case it crashes e.g. due to time limit or hardware failure on a cluster. This is done by setting `guess_type = chk` in the subsequent calculation:

```
SCF {
    guess_type = chk                      # Type of initial guess (chk, mw, cube, core_XX, ...
    ↵sad_XX)
}
```

In this case the `path_checkpoint` must be the same as the previous calculation, as well as all other parameters in the calculation (Molecule and Basis in particular).

Write orbitals

The converged orbitals can be saved to file with the `write_orbitals` keyword (defaults shown):

```
SCF {
    path_orbitals = orbitals            # Path to orbital files
    write_orbitals = false              # Save converged orbitals to file
}
```

This will make individual files for each orbital under the `path_orbitals` directory. These orbitals can be used as starting point for subsequent calculations using the `guess_type = mw` initial guess:

```
SCF {
    guess_prec = 1.0e-3                # Numerical precision used in guess
    guess_type = mw                    # Type of initial guess (chk, mw, cube, core_XX, ...
    ↵sad_XX)
}
```

Here the orbitals will be re-projected onto the current MW basis with precision `guess_prec`. We also need to specify the paths to the input files:

```
Files {
    guess_phi_p = initial_guess/phi_p      # Path to paired MW orbitals
    guess_phi_a = initial_guess/phi_a      # Path to alpha MW orbitals
    guess_phi_b = initial_guess/phi_b      # Path to beta MW orbitals
}
```

Note that by default orbitals are written to the directory called `orbitals` but the `mw` guess reads from the directory `initial_guess` (this is to avoid overwriting the files by default). So, in order to use MW orbitals from a previous calculation, you must either change one of the paths (`SCF.path_orbitals` or `Files.guess_phi_p` etc), or manually copy the files between the default locations.

Note: The `mw` guess must not be confused with the `chk` guess, although they are similar. The `chk` guess will blindly read

in the orbitals that are present, regardless of the current molecular structure and computational setup (if you run with a different computational domain or MW basis type/order the calculation will crash). The `mw` guess will re-project the old orbitals onto the new computational setup and populate the orbitals based on the *new* molecule (here the computation domain and MW basis do *not* have to match).

Response

This section specifies the parameters for the SCF optimization of the linear response functions. There might be several independent response calculations depending on the requested properties, e.g.

```
Polarizability {
    frequency = [0.0, 0.0656]           # List of frequencies to compute
}
```

will run one response for each frequency (each with three Cartesian components), while

```
Properties {
    magnetizability = true            # Compute magnetizability
    nmr_shielding = true             # Compute NMR shieldings
}
```

will combine both properties into a single response calculation, since the perturbation operator is the same in both cases (unless you choose `NMRShielding.nuclear_specific = true`, in which case there will be a different response for each nucleus).

Response solver

The optimization is controlled by the following keywords (defaults shown):

```
Response {
    run = [true,true,true]           # Run response solver [x,y,z] direction
    kain = 5                         # Length of KAIN iterative subspace
    max_iter = 100                   # Maximum number of SCF iterations
    localize = false                 # Use canonical or localized orbitals
    start_prec = -1.0                # Dynamic precision, start value
    final_prec = -1.0                # Dynamic precision, final value
    orbital_thrs = 10 * world_prec  # Convergence threshold orbitals
}
```

Each linear response calculation involves the three Cartesian components of the appropriate perturbation operator. If any of the components of `run` is `false`, no response is performed in that particular direction, and the properties are computed directly on the initial guess response functions (usually zero guess).

The `kain` (Krylov Accelerated Inexact Newton) keyword gives the length of the iterative subspace accelerator (similar to DIIS). The `localize` keyword relates to the unperturbed orbitals, and can be set independently of the `SCF.localize` keyword.

The dynamic precision keywords control how the numerical precision is changed throughout the optimization. One can choose to use a lower `start_prec` in the first iterations which is gradually increased to `final_prec` (both are equal to `world_prec` by default). Note that lower initial precision might affect the convergence rate.

For response calculations, the important convergence threshold is that of the orbitals, and by default this is set one order of magnitude higher than the overall `world_prec`.

Note: The quality of the response property depends on both the perturbed as well as the unperturbed orbitals, so they should be equally well converged.

Initial guess

The following initial guesses are available:

- `none` start from a zero guess for the response functions.
- `chk` and `mw` require input files from previous MW calculations.

By default, no initial guess is generated for the response functions, but the `chk` and `mw` guesses work similarly as for the SCF.

Checkpointing

The program can dump checkpoint files at every iteration using the `write_checkpoint` keyword (defaults shown):

```
Response {
    path_checkpoint = checkpoint          # Path to checkpoint files
    write_checkpoint = false               # Save checkpoint files every iteration
}
```

This allows the calculation to be restarted in case it crashes e.g. due to time limit or hardware failure on a cluster. This is done by setting `guess_type = chk` in the subsequent calculation:

```
Response {
    guess_type = chk                      # Type of initial guess (none, chk, mw)
}
```

In this case the `path_checkpoint` must be the same as the previous calculation, as well as all other parameters in the calculation (Molecule and Basis in particular).

Write orbitals

The converged response orbitals can be saved to file with the `write_orbitals` keyword (defaults shown):

```
Response {
    path_orbitals = orbitals            # Path to orbital files
    write_orbitals = false              # Save converged orbitals to file
}
```

This will make individual files for each orbital under the `path_orbitals` directory. These orbitals can be used as starting point for subsequent calculations using the `guess_type = mw` initial guess:

```
Response {
    guess_prec = 1.0e-3                 # Numerical precision used in guess
    guess_type = mw                     # Type of initial guess (chk, mw, cube, core_XX, ...
    ↵sad_XX)
}
```

Here the orbitals will be re-projected onto the current MW basis with precision `guess_prec`. We also need to specify the paths to the input files (only X for static perturbations, X and Y for dynamic perturbations):

```
Files {
    guess_X_p = initial_guess/X_p          # Path to paired MW orbitals
    guess_X_a = initial_guess/X_a          # Path to alpha MW orbitals
    guess_X_b = initial_guess/X_b          # Path to beta MW orbitals
    guess_Y_p = initial_guess/Y_p          # Path to paired MW orbitals
    guess_Y_a = initial_guess/Y_a          # Path to alpha MW orbitals
    guess_Y_b = initial_guess/Y_b          # Path to beta MW orbitals
}
```

Note that by default orbitals are written to the directory called `orbitals` but the `mw` guess reads from the directory `initial_guess` (this is to avoid overwriting the files by default). So, in order to use MW orbitals from a previous calculation, you must either change one of the paths (`Response.path_orbitals` or `Files.guess_X_p` etc), or manually copy the files between the default locations.

2.2.3 User input reference

- Keywords without a default value are **required**.
- Default values are either explicit or computed from the value of other keywords in the input.
- Sections where all keywords have a default value can be omitted.
- Predicates, if present, are the functions run to validate user input.

Keywords

`world_prec`

Overall relative precision in the calculation.

Type float

Predicates

- `1.0e-10 < value < 1.0`

`world_size`

Total size of computational domain given as `2**world_size`. Always cubic and symmetric around the origin. Negative value means it will be computed from the molecular geometry.

Type int

Default -1

Predicates

- `value <= 10`

`world_unit`

Length unit for *all* coordinates given in user input. Everything will be converted to atomic units (bohr) before the main executable is launched, so the JSON input is *always* given in bohrs.

Type str

Default bohr

Predicates

- `value.lower() in ["bohr", "angstrom"]`

world_origin

Global gauge origin of the calculation.

Type List[float]

Default [0.0, 0.0, 0.0]

Predicates

- len(value) == 3

Sections**Precisions**

Define specific precision parameters.

Keywords**exchange_prec**

Precision parameter used in construction of Exchange operators. Negative value means it will follow the dynamic precision in SCF.

Type float

Default -1.0

helmholtz_prec

Precision parameter used in construction of Helmholtz operators. Negative value means it will follow the dynamic precision in SCF.

Type float

Default -1.0

poisson_prec

Precision parameter used in construction of Poisson operators.

Type float

Default user['world_prec']

Predicates

- 1.0e-10 < value < 1.0

nuclear_prec

Precision parameter used in smoothing and projection of nuclear potential.

Type float

Default user['world_prec']

Predicates

- 1.0e-10 < value < 1.0

Printer

Define variables for printed output.

Keywords**print_level**

Level of detail in the written output. Level 0 for production calculations, negative level for complete silence.

Type int

Default 0

print_mpi
Write separate output from each MPI to file called
`<file_name>-<mpi-rank>.out`.

Type bool

Default False

print_prec

Number of digits in property output (energies will get twice this number of digits).

Type int

Default 6

Predicates

- $0 < \text{value} < 10$

print_width

Line width of printed output (in number of characters).

Type int

Default 75

Predicates

- $50 < \text{value} < 100$

print_constants

Print table of physical constants used by MRChem.

Type bool

Default False

Plotter

Give details regarding the density and orbital plots. Three types of plots are available, line, surface and cube, and the plotting ranges are defined by three vectors (A, B and C) and an origin (O): line: plots on line spanned by A, starting from O. surf: plots on surface spanned by A and B, starting from O. cube: plots on volume spanned by A, B and C, starting from O.

Keywords

path

File path to plot directory.

Type str

Default plots

Predicates

- $\text{value}[-1] \neq '/'$

type

Type of plot: line (1D), surface (2D) or cube (3D).

Type str

Default cube

Predicates

- $\text{value.lower()} \in ['line', 'surf', 'cube']$

points

Number of points in each direction on the cube grid.

Type List[int]

Default [20, 20, 20]

Predicates

- all(p > 0 for p in value)
- not (user['Plotter']['type'] == 'line' and len(value) < 1)
- not (user['Plotter']['type'] == 'surf' and len(value) < 2)
- not (user['Plotter']['type'] == 'cube' and len(value) < 3)

O

Origin of plotting ranges.

Type List[float]

Default [0.0, 0.0, 0.0]

Predicates

- len(value) == 3

A

First boundary vector for plot.

Type List[float]

Default [1.0, 0.0, 0.0]

Predicates

- len(value) == 3

B

Second boundary vector for plot.

Type List[float]

Default [0.0, 1.0, 0.0]

Predicates

- len(value) == 3

C

Third boundary vector for plot.

Type List[float]

Default [0.0, 0.0, 1.0]

Predicates

- len(value) == 3

MPI

Define MPI related parameters.

Keywords

numerically_exact

This will use MPI algorithms that guarantees that the output is invariant wrt the number of MPI processes.

Type bool

Default False

shared_memory_size

Size (MB) of the MPI shared memory blocks of each shared function.

Type int

Default 10000

share_nuclear_potential

This will use MPI shared memory for the nuclear potential.

Type bool

Default False

share_coulomb_potential

This will use MPI shared memory for the Coulomb potential.

Type bool

Default False

share_xc_potential

This will use MPI shared memory for the exchange-correlation potential.

Type bool

Default False

bank_size

Number of MPI processes exclusively dedicated to manage orbital bank.

Type int

Default -1

omp_threads

Force the number of OpenMP threads.

Type int

Default -1

Basis

Define polynomial basis.

Keywords**order**

Polynomial order of multiwavelet basis. Negative value means it will be set automatically based on the world precision.

Type int

Default -1

type

Polynomial type of multiwavelet basis.

Type str

Default interpolating

Predicates

- value.lower() in ['interpolating', 'legendre']

Derivatives

Define various derivative operators used in the code.

Keywords

kinetic

Derivative used in kinetic operator.

Type str

Default abgv_55

h_b_dip

Derivative used in magnetic dipole operator.

Type str

Default abgv_00

h_m_pso

Derivative used in paramagnetic spin-orbit operator.

Type str

Default abgv_00

zora

Derivative used ZORA potential.

Type str

Default abgv_00

Molecule

Define molecule.

Keywords

charge

Total charge of molecule.

Type int

Default 0

multiplicity

Spin multiplicity of molecule.

Type int

Default 1

Predicates

- value > 0

translate

Translate coordinates such that center of mass coincides with the global gauge origin.

Type bool

Default False

coords

Coordinates in xyz format. Atoms can be given either using atom symbol or atom number

Type str

WaveFunction

Define the wavefunction method.

Keywords

method

Wavefunction method. See predicates for valid methods. hf, hartreefock and hartree-fock all mean the same thing, while lda is an alias for svwn5. You can set a non-standard DFT functional (e.g. varying the amount of exact exchange) by choosing dft and specifying the functional(s) in the DFT section below.

Type str

Predicates

- value.lower() in ['core', 'hartree', 'hf', 'hartreefock', 'hartree-fock', 'dft', 'lda', 'svwn3', 'svwn5', 'pbe', 'pbe0', 'bpw91', 'bp86', 'b3p86', 'b3p86-g', 'blyp', 'b3lyp', 'b3lyp-g', 'olyp', 'kt1', 'kt2', 'kt3']

restricted

Use spin restricted wavefunction.

Type bool

Default True

relativity

Set method for relativistic treatment. ZORA for fully self-consistent ZORA potential, by default including all potentials (V_nuc, J, V_xc) but this can be overwritten in the ZORA section. nzora is shortcut for nuclear-ZORA, i.e. only V_nuc is included (this keyword takes precedence over keywords in the ZORA section).

Type str

Default none

Predicates

- value.lower() in ['none', 'zora', 'nzora']

environment

Set method for treatment of environment. none for vacuum calculation. PCM for Polarizable Continuum Model, which will activate the PCM input section for further parametrization options. The PB and LPB variants add the Poisson-Boltzmann and Linearized Poisson-Boltzmann solvers, respectively.

Type str

Default none

Predicates

- `value.lower() in ['none', 'pcm', 'pcm_pb', 'pcm_lpb']`

nuclear_model

Type of nucleus model. Point-like (numerical smoothing): HFYGB (default), parabola or minimal. Finite models (physical smoothing): Gaussian or Homogeneous sphere Finite models are derived from nuclear RMS radius, Visscher (1997)

Type str**Default** point_like**Predicates**

- `value.lower() in ['point_like', 'point_parabola', 'point_minimal', 'finite_gaussian', 'finite_sphere']`

ZORA

Define required parameters for the ZORA Hamiltonian.

Keywords**include_nuclear**

Include the nuclear potential V_{nuc} in the ZORA potential.

Type bool**Default** True**include_coulomb**

Include the Coulomb potential J in the ZORA potential.

Type bool**Default** True**include_xc**

Include the XC potential V_{xc} in the ZORA potential.

Type bool**Default** True**DFT**

Define the exchange-correlation functional in case of DFT.

Keywords**density_cutoff**

Hard cutoff for passing density values to XCFun.

Type float**Default** 0.0**functionals**

List of density functionals with numerical coefficient. E.g. for PBE0 EXX 0.25, PBEX 0.75, PBEC 1.0, see XCFun documentation <<https://xcfun.readthedocs.io/>>_.

Type str**Default** `` ``**spin**

Use spin separated density functionals.

Type bool

Default not(user['WaveFunction']['restricted'])

Properties

Provide a list of properties to compute (total SCF energy and orbital energies are always computed).

Keywords

dipole_moment

Compute dipole moment.

Type bool

Default True

quadrupole_moment

Compute quadrupole moment. Note: Gauge origin dependent, should be used with `translate = true` in Molecule.

Type bool

Default False

polarizability

Compute polarizability tensor.

Type bool

Default False

magnetizability

Compute magnetizability tensor.

Type bool

Default False

nmr_shielding

Compute NMR shielding tensor.

Type bool

Default False

plot_density

Plot converged electron density.

Type bool

Default False

plot_orbitals

Plot converged molecular orbitals from list of indices, negative index plots all orbitals.

Type List[int]

Default []

geometric_derivative

Compute geometric derivative.

Type bool

Default user['GeometryOptimizer']['run']

ExternalFields

Define external electromagnetic fields.

Keywords**electric_field**

Strength of external electric field.

Type List[float]

Default []

Predicates

- len(value) == 0 or len(value) == 3

Polarizability

Give details regarding the polarizability calculation.

Keywords**frequency**

List of external field frequencies.

Type List[float]

Default [0.0]

NMRShielding

Give details regarding the NMR shileding calculation.

Keywords**nuclear_specific**

Use nuclear specific perturbation operator (h_m_pso).

Type bool

Default False

nucleus_k

List of nuclei to compute. Negative value computes all nuclei.

Type List[int]

Default [-1]

Files

Defines file paths used for program input/output. Note: all paths must be given in quotes if they contain slashes “path/to/file”.

Keywords**guess_basis**

File name for GTO basis set, used with gto guess.

Type str

Default initial_guess/mrchem.bas

guess_gto_p

File name for paired orbitals, used with gto guess.

Type str

Default initial_guess/mrchem.mop

guess_gto_a

File name for alpha orbitals, used with gto guess.

Type str

Default initial_guess/mrchem.moa

guess_gto_b

File name for beta orbitals, used with gto guess.

Type str

Default initial_guess/mrchem.mob

guess_phi_p

File name for paired orbitals, used with mw guess. Expected path is
``<path_orbitals>/phi_p_scf_idx_<0...Np>_<re/im>.mw

Type str

Default initial_guess/phi_p

guess_phi_a

File name for alpha orbitals, used with mw guess. Expected path is
``<path_orbitals>/phi_a_scf_idx_<0...Na>_<re/im>.mw

Type str

Default initial_guess/phi_a

guess_phi_b

File name for beta orbitals, used with mw guess. Expected path is
``<path_orbitals>/phi_b_scf_idx_<0...Nb>_<re/im>.mw

Type str

Default initial_guess/phi_b

guess_x_p

File name for paired response orbitals, used with mw guess. Expected path is
``<path_orbitals>/x_p_rsp_idx_<0...Np>_<re/im>.mw

Type str

Default initial_guess/x_p

guess_x_a

File name for alpha response orbitals, used with mw guess. Expected path is
``<path_orbitals>/x_a_rsp_idx_<0...Na>_<re/im>.mw

Type str

Default initial_guess/X_a

guess_x_b

File name for beta response orbitals, used with mw guess. Expected path is
``<path_orbitals>/x_b_rsp_idx_<0...Nb>_<re/im>.mw

Type str

Default initial_guess/X_b

guess_y_p

File name for paired response orbitals, used with mw guess. Expected path is
``<path_orbitals>/y_p_rsp_idx_<0...Np>_<re/im>.mw

Type str**Default** initial_guess/Y_p**guess_y_a**

File name for alpha response orbitals, used with mw guess. Expected path is
``<path_orbitals>/y_a_rsp_idx_<0...Na>_<re/im>.mw

Type str**Default** initial_guess/Y_a**guess_y_b**

File name for beta response orbitals, used with mw guess. Expected path is
``<path_orbitals>/y_b_rsp_idx_<0...Nb>_<re/im>.mw

Type str**Default** initial_guess/Y_b**guess_cube_p**

File name for paired orbitals, used with cube guess. Expected path is
``<path_orbitals>/phi_p_scf_idx_<0...Np>_<re/im>.cube where *Np* is the
number of orbitals and *re/im* denote real or imaginary parts.

Type str**Default** initial_guess/phi_p**guess_cube_a**

File name for alpha orbitals, used with cube guess. Expected path is
``<path_orbitals>/phi_a_scf_idx_<0...Na>_<re/im>.cube where *Np* is the
number of orbitals and *re/im* denote real or imaginary parts.

Type str**Default** initial_guess/phi_a**guess_cube_b**

File name for beta orbitals, used with cube guess. Expected path is
``<path_orbitals>/phi_b_scf_idx_<0...Nb>_<re/im>.cube where *Np* is the
number of orbitals and *re/im* denote real or imaginary parts.

Type str**Default** initial_guess/phi_b**guess_cube_x_p**

File name for paired response orbitals, used with cube guess. Expected path
is ``<path_orbitals>/x_p_rsp_<0,1,2>_<0...Np>_<re/im>.cube where 0,1,2
indicate the index of the components of the response vectors. *Np* is the number
of orbitals and *re/im* denote real or imaginary parts.

Type str**Default** initial_guess/x_p**guess_cube_x_a**

File name for alpha response orbitals, used with cube guess. Expected path
is ``<path_orbitals>/x_a_rsp_<0,1,2>_<0...Na>_<re/im>.cube where 0,1,2
indicate the index of the components of the response vectors. *Np* is the number
of orbitals and *re/im* denote real or imaginary parts.

Type str

Default initial_guess/x_a**guess_cube_x_b**

File name for beta response orbitals, used with cube guess. Expected path is ``<path_orbitals>/x_b_rsp_<0,1,2>_<0...Nb>_<re/im>.cube where 0,1,2 indicate the index of the components of the response vectors. Np is the number of orbitals and *re/im* denote real or imaginary parts.

Type str**Default initial_guess/x_b****guess_cube_y_p**

File name for paired response orbitals, used with cube guess. Expected path is ``<path_orbitals>/y_p_rsp_<0,1,2>_<0...Np>_<re/im>.cube where 0,1,2 indicate the index of the components of the response vectors. Np is the number of orbitals and *re/im* denote real or imaginary parts.

Type str**Default initial_guess/y_p****guess_cube_y_a**

File name for alpha response orbitals, used with cube guess. Expected path is ``<path_orbitals>/y_a_rsp_<0,1,2>_<0...Na>_<re/im>.cube where 0,1,2 indicate the index of the components of the response vectors. Np is the number of orbitals and *re/im* denote real or imaginary parts.

Type str**Default initial_guess/y_a****guess_cube_y_b**

File name for beta response orbitals, used with cube guess. Expected path is ``<path_orbitals>/y_b_rsp_<0,1,2>_<0...Nb>_<re/im>.cube where 0,1,2 indicate the index of the components of the response vectors. Np is the number of orbitals and *re/im* denote real or imaginary parts.

Type str**Default initial_guess/y_b****cube_vectors**

Directory where cube vectors are stored for mrchem calculation.

Type str**Default cube_vectors/****SCF**

Includes parameters related to the ground state SCF orbital optimization.

Keywords**run**

Run SCF solver. Otherwise properties are computed on the initial orbitals.

Type bool**Default True****max_iter**

Maximum number of SCF iterations.

Type int

Default 100

kain

Length of KAIN iterative history.

Type int

Default 5

rotation

Number of iterations between each diagonalization/localization.

Type int

Default 0

localize

Use canonical or localized orbitals.

Type bool

Default False

energy_thrs

Convergence threshold for SCF energy.

Type float

Default -1.0

guess_prec

Precision parameter used in construction of initial guess.

Type float

Default 0.001

Predicates

- $1.0e-10 < \text{value} < 1.0$

guess_screen

Screening parameter used in GTO evaluations, in number of standard deviations. Every coordinate beyond N StdDev from the Gaussian center is evaluated to zero. Note that too aggressive screening is counter productive, because it leads to a sharp cutoff in the resulting function which requires higher grid refinement. Negative value means no screening.

Type float

Default 12.0

start_prec

Incremental precision in SCF iterations, initial value.

Type float

Default -1.0

final_prec

Incremental precision in SCF iterations, final value.

Type float

Default -1.0

guess_type

Type of initial guess for ground state orbitals. `chk` restarts a previous calculation which was dumped using the `write_checkpoint` keyword. This will load MRA and electron spin configuration directly from the checkpoint files, which are thus required to be identical in the two calculations. `mw` will start from final orbitals in a previous calculation written using the `write_orbitals` keyword. The orbitals will be re-projected into the new computational setup, which means that the electron spin configuration and MRA can be different in the two calculations. `gto` reads precomputed GTO orbitals (requires extra non-standard input files for basis set and MO coefficients). `core` and `sad` will diagonalize the Fock matrix in the given AO basis (SZ, DZ, TZ or QZ) using a Core or Superposition of Atomic Densities Hamiltonian, respectively. `cube` will start from orbitals saved in cubefiles from external calculations.

Type str**Default** sad_gto**Predicates**

- value.lower() in ['mw', 'chk', 'gto', 'core_sz', 'core_dz', 'core_tz', 'core_qz', 'sad_sz', 'sad_dz', 'sad_tz', 'sad_qz', 'sad_gto', 'cube']

write_checkpoint

Write orbitals to disk in each iteration, file name <path_checkpoint>/phi_scf_idx_<0..N>. Can be used as `chk` initial guess in subsequent calculations. Note: must be given in quotes if there are slashes in the path “path/to/checkpoint”.

Type bool**Default** False**path_checkpoint**

Path to checkpoint files during SCF, used with `write_checkpoint` and `chk` guess.

Type str**Default** checkpoint**Predicates**

- value[-1] != '/'

path_orbitals

Path to where converged orbitals will be written in connection with the `write_orbitals` keyword. Note: must be given in quotes if there are slashes in the path “path/to/orbitals”.

Type str**Default** orbitals**Predicates**

- value[-1] != '/'

write_orbitals

Write final orbitals to disk, file name <path_orbitals>/phi_<p/a/

b>_scf_idx_<0..Np/Na/Nb>. Can be used as **mw** initial guess in subsequent calculations.

Type bool

Default user['GeometryOptimizer']['use_previous_guess']

orbital_thrs

Convergence threshold for orbital residuals.

Type float

Default 10 * user['world_prec']

Response

Includes parameters related to the response SCF optimization.

Keywords

run

In which Cartesian directions to run response solver.

Type List[bool]

Default [True, True, True]

max_iter

Maximum number of response iterations.

Type int

Default 100

kain

Length of KAIN iterative history.

Type int

Default 5

property_thrs

Convergence threshold for symmetric property. Symmetric meaning the property computed from the same operator as the response perturbation, e.g. for external magnetic field the symmetric property corresponds to the magnetizability (NMR shielding in non-symmetric, since one of the operators is external magnetic field, while the other is nuclear magnetic moment).

Type float

Default -1.0

start_prec

Incremental precision in SCF iterations, initial value.

Type float

Default -1.0

final_prec

Incremental precision in SCF iterations, final value.

Type float

Default -1.0

guess_prec

Precision parameter used in construction of initial guess.

Type float

Default 0.001

Predicates

- 1.0e-10 < value < 1.0

guess_type

Type of initial guess for response. none will start from a zero guess for the response functions. chk restarts a previous calculation which was dumped using the write_checkpoint keyword. mw will start from final orbitals in a previous calculation written using the write_orbitals keyword. The orbitals will be re-projected into the new computational setup.

Type str

Default none

Predicates

- value.lower() in ['none', 'chk', 'mw', 'cube']

write_checkpoint

Write perturbed orbitals to disk in each iteration, file name <path_checkpoint>/<X/Y>_rsp_<direction>_idx_<0..N>. Can be used as chk initial guess in subsequent calculations.

Type bool

Default False

path_checkpoint

Path to checkpoint files during SCF, used with write_checkpoint and chk guess.

Type str

Default checkpoint

Predicates

- value[-1] != '/'

write_orbitals

Write final perturbed orbitals to disk, file name <path_orbitals>/<X/Y>_<p/a/b>_rsp_<direction>_idx_<0..Np>/Na/Nb>. Can be used as mw initial guess in subsequent calculations.

Type bool

Default False

path_orbitals

Path to where converged orbitals will be written in connection with the write_orbitals keyword.

Type str

Default orbitals

Predicates

- `value[-1] != '/'`

orbital_thrs

Convergence threshold for orbital residuals.

Type float

Default 10 * user['world_prec']

localize

Use canonical or localized unperturbed orbitals.

Type bool

Default user['SCF']['localize']

PCM

Includes parameters related to the computation of the reaction field energy of a system in an environment within the Polarizable Continuum Model.

Sections**SCRF**

Parameters for the Self-Consistent Reaction Field optimization.

Keywords**max_iter**

Max number of iterations allowed in the nested procedure.

Type int

Default 100

dynamic_thrs

Set the convergence threshold for the nested procedure. `true` will dynamically tighten the convergence threshold based on the absolute value of the latest orbital update as. When the orbitals are close to convergence (`mo_residual < world_prec*10`) the convergence threshold will be set equal to `world_prec`. `false` uses `world_prec` as convergence threshold throughout.

Type bool

Default True

density_type

What part of the total molecular charge density to use in the algorithm. `total` uses the total charge density. `nuclear` uses only the nuclear part of the total charge density. `electronic` uses only the electronic part of the total charge density.

Type str

Default total

Predicates

- `value.lower() in ['total', 'nuclear', 'electronic']`

kain

Number of previous reaction field iterates kept for convergence acceleration during the nested procedure.

Type int**Default** user['SCF']['kain']**Solvent**

Parameters for the Self-Consistent Reaction Field optimization.

Sections**Permittivity**

Parameters for the permittivity function.

Keywords**epsilon_in**

Permittivity inside the cavity. 1.0 is the permittivity of free space, anything other than this is undefined behaviour.

Type float**Default** 1.0**formulation**

Formulation of the Permittivity function. Currently only the exponential is available.

Type str**Default** exponential**Predicates**

- value.lower() in ['exponential']

Sections**epsilon_out**

Parameters for the continuum solvent outside the cavity.

Keywords**nonequilibrium**

Whether to use the nonequilibrium formulation of response, *i.e.* use the dynamic permittivity for the calculation of the response reaction field. Defaults to false.

Type bool**Default** False**static**

Static permittivity outside the cavity. This is characteristic of the solvent used.

Type float**Default** 1.0

dynamic

Dynamic permittivity outside the cavity. This is characteristic of the solvent used and relevant only in response calculations. Defaults to the same value as *epsilon_static*.

Type float

Default

user['PCM']['Solvent']['Permittivity']['epsilon_out'][

DebyeHuckelScreening

Parameters for the Debye-Hückel screening factor

Keywords**ion_strength**

Ionic strength of the electrolyte in mol/L. This represents the concentration of the ions in the bulk solvent.

Type float

Default 1.0

ion_radius

Amount with which the vdw-radius of the atoms will be increased. The screening factor will have an area of effect that is often going to be larger than the vdw-cavity, but centered in the same atoms.

Type float

Default 0.0

ion_width

Width of the transition between the solute and the ion accessible part.

Type float

Default 0.2

formulation

formulation of the debye-huckel screening factor. Currently only the variable factor is implemented.
variable: implement the screening functions as $k = (1 - C_{\text{ion}})k_{\text{out}}$

Type str

Default variable

Predicates

- `value.lower() in ['variable']`

Cavity

Define the interlocking spheres cavity.

Keywords

mode

Determines how to set up the interlocking spheres cavity.
atoms: centers are taken from the molecular geometry, radii taken from tabulated data (van der Waals radius), and rescaled using the parameters alpha, beta and sigma ($R_i <- \alpha * R_i + \beta * \sigma$). Default spheres can be modified and/or extra spheres added, using the \$spheres section, see documentation. **explicit**: centers and radii given explicitly in the spheres block.

Type str

Default atoms

Predicates

- value.lower() in ['atoms', 'explicit']

spheres

This input parameter affects the list of spheres used to generate the cavity. In all cases, values for the radius, the radius scaling factor (alpha), the width (sigma), and the width scaling factor (beta) can be modified. If they are not specified their global default values are used. In **atoms** mode, we *modify* the default list of spheres, built with centers from the molecular geometry and radii from internal tabulated van der Waals values. To *substitute* a sphere, include a line like: `` \$spheres i R [alpha] [beta] [sigma] \$end `` to specify that the *i* atom in the molecule (0-based indexing) should use radius *R* instead of the pre-tabulated vdW radius. To *add* a sphere, include a line like: `` \$spheres x y z R [alpha] [beta] [sigma] \$end ` to specify that a sphere of radius ` *R* should be added at position (*x*, *y*, *z*). Spheres added in this way are not aware of their parent atom, if any. They will **not** contribute to the molecular gradient. In **explicit** mode, we *build* the complete sphere list from scratch. You can add a line like: `` \$spheres x y z R [alpha] [beta] [sigma] \$end `` to specify that a sphere of radius *R* should be added at position (*x*, *y*, *z*). Spheres added in this way are not aware of their parent atom, if any. They will **not** contribute to the molecular gradient. Alternatively, you can specify a line like: `` \$spheres i R [alpha] [beta] [sigma] \$end `` to specify that the *i* atom in the molecule (0-based indexing) should use radius *R*. Spheres added in this way are aware of their parent atom. They will contribute to the molecular gradient.

Type str

Default ````

alpha

Scaling factor on the radius term for the cavity rescaling (R_i)

$\leftarrow \alpha * R_i + \beta * \sigma\right). Only used for the default vdW radii in *atoms* mode, not if explicit $spheres are given.$

Type float

Default 1.1

beta

Scaling factor on the boundary width term for the cavity rescaling ($R_i \leftarrow \alpha * R_i + \beta * \sigma$). Only used for the default vdW radii in *atoms* mode, not if explicit \$spheres are given.

Type float

Default 0.5

sigma

Width of cavity boundary, smaller value means sharper transition. **This quantity has dimensions of length. The default value is in atomic units.**

Type float

Default 0.2

GeometryOptimizer

Includes parameters related to the internal geometry optimization using the SQNM (Stabilized Quasi-Newton Method) for noisy PES. Geometry optimizations require accurate forces. Consider setting world_prec to 1e-5 to 1e-7. Convergence issues can usually be solved by increasing the precision of the SCF calculation. If that does not work, try setting the initial step size manually.

Keywords

run

Run optimizer. Otherwise single point energy/properties are computed.

Type bool

Default False

use_previous_guess

Start each SCF from the converged orbitals from the previous geometry step. The guess_type will change to “mw” after the first iteration, and the intermediate orbitals will be stored in the “orbitals” directory. If toggled off, start over using the same initial guess method as in the first iteration.

Type bool

Default False

init_step_size

Initial step size. For systems with hard bonds (e.g. C-C) use a value between and 1.0 and 2.5. If a system only contains weaker bonds a value up to 5.0 may speed up the convergence. Use a small negative number (should be between -0.1 and -0.5) for an automatic guess. The optimal step size is the inverse of the largest eigenvalue of the hessian matrix.

Type float

Default -0.5

minimal_step_size

Minimal step size. It rarely makes sense to change it.

Type float

Default 0.01

max_history_length

Maximum length of history list. Energies and forces from the previous n geometry optimization iterations are used to estimate the hessian matrix. Use a value between 2 and 20. A lower value makes the SQNM algorithm behave more like steepest descent and slows down convergence. But it can handle more noise in the energies and forces. It rarely makes sense to change it.

Type int

Default 10

subspace_tolerance

Lower limit on linear dependencies of basis vectors in history listSubspace tolerance. Use a number between 1e-9 and 1e-1. A high subspace tolerance slows down convergence but improves numerical stability when the energies and forces contain a lot of noise. It rarely makes sense to change it.

Type float

Default 0.001

max_iter

Maximum number of geometry optimization iterations.

Type int

Default 100

max_force_component

The geometry optimization stopps when the absolute value of all force components is smaller than this keyword. A value between 1e-3 and 1e-4 is tight enough for most applications.

Type float

Default 0.005

Constants

Physical and mathematical constants used by MRChem

Keywords

hartree2simagnetizability

Conversion factor for magnetizability from atomic units to SI units (unit: J T^-2). Affected code: Printed value of the magnetizability property.

Type float

Default 78.9451185

light_speed

Speed of light in atomic units (unit: au). Affected code: Relativistic Hamiltonians (ZORA, etc.)

Type float

Default 137.035999084

angstrom2bohrs

Conversion factor for Cartesian coordinates from Angstrom to Bohr (unit: Å^-1). Affected code: Parsing of input coordinates, printed coordinates

Type float

Default 1.8897261246257702

hartree2kjamol

Conversion factor from Hartree to kJ/mol (unit: kJ mol^-1). Affected code: Printed value of energies.

Type float

Default 2625.4996394798254

hartree2kcalmol

Conversion factor from Hartree to kcal/mol (unit: kcal mol^-1). Affected code: Printed value of energies.

Type float

Default 627.5094740630558

hartree2ev

Conversion factor from Hartree to eV (unit: ev). Affected code: Printed value of energies.

Type float

Default 27.211386245988

hartree2wavenumbers

Conversion factor from Hartree to wavenumbers (unit: cm^-1). Affected code: Printed value of frequencies.

Type float

Default 219474.6313632

fine_structure_constant

Fine-structure constant in atomic units (unit: au). Affected code: Certain magnetic interaction operators.

Type float

Default 0.0072973525693

electron_g_factor

Electron g factor in atomic units (unit: au). Affected code: Certain magnetic interaction operators.

Type float**Default** -2.00231930436256**dipmom_au2debye**

Conversion factor for dipoles from atomic units to Debye (unit: ?). Affected code: Printed value of dipole moments.

Type float**Default** 2.5417464739297717**boltzmann_constant**

Boltzmann constant in (unit: J K⁻¹). Affected code: Value of the Debye-Huckel screening parameter in the Poisson-Boltzmann equation.

Type float**Default** 1.380649e-23**elementary_charge**

Elementary charge in (unit: C). Affected code: Value of the Debye-Huckel screening parameter in the Poisson-Boltzmann equation.

Type float**Default** 1.602176634e-19**e0**

Permittivity of free space (unit: F m⁻¹). Affected code: Value of the Debye-Huckel screening parameter in the Poisson-Boltzmann equation.

Type float**Default** 8.8541878128e-12**N_a**

Avogadro constant (unit: mol⁻¹). Affected code: Value of the Debye-Huckel screening parameter in the Poisson-Boltzmann equation.

Type float**Default** 6.02214076e+23**meter2bohr**

conversion factor from meter to Bohr radius (unit: m⁻¹). Affected code: Value of the Debye-Huckel screening parameter in the Poisson-Boltzmann equation.

Type float**Default** 18897261246.2577**Elements**

list of elements with data

Sections

h

data of element

Keywords

vdw-radius

radius of element

Type float

Default 1.2

covalent

covalent value element

Type float

Default 0.32

Z

z-value of element

Type int

Default 1

mass

mass of element

Type float

Default 1.00794

symbol

symbol of element

Type str

Default H

bpt

bpt of element

Type float

Default 20.268

mpt

mpt of element

Type float

Default 14.025

density

density of element

Type float

Default 0.0899

volume

volume of element

Type float

Default 14.4

name
name of element
Type str
Default Hydrogen

debye
debye of element
Type float
Default 110.0

a
a of element
Type float
Default 3.75

crystal
crystal of element
Type str
Default HEX

cpera
cpera of element
Type float
Default 1.731

conf
conf of element
Type str
Default 1s1

r_rms
r_rms of element
Type float
Default 2.6569547399e-05

he
data of element

Keywords

vdw-radius
radius of element
Type float
Default 1.4

covalent
covalent value element
Type float

Default 0.93

Z

z-value of element

Type int

Default 2

mass

mass of element

Type float

Default 4.002602

symbol

symbol of element

Type str

Default He

bpt

bpt of element

Type float

Default 4.215

mpt

mpt of element

Type float

Default 0.95

density

density of element

Type float

Default 0.1787

volume

volume of element

Type float

Default 0.0

name

name of element

Type str

Default Helium

debye

debye of element

Type float

Default -26.0

a

a of element

Type float
Default 3.57

crystal
crystal of element

Type str
Default HEX

cpera
cpera of element

Type float
Default 1.633

conf
conf of element

Type str
Default 1s2

r_rms
r_rms of element

Type float
Default 3.5849373401e-05

li
data of element

Keywords

vdw-radius
radius of element

Type float
Default 1.81

covalent
covalent value element

Type float
Default 1.23

Z
z-value of element

Type int
Default 3

mass
mass of element

Type float
Default 6.941

symbol
symbol of element

Type str
Default Li

bpt
bpt of element

Type float
Default 1615.0

mpt
mpt of element

Type float
Default 453.7

density
density of element

Type float
Default 0.53

volume
volume of element

Type float
Default 13.1

name
name of element

Type str
Default Lithium

debye
debye of element

Type float
Default 400.0

a
a of element

Type float
Default 3.49

crystal
crystal of element

Type str
Default BCC

cpera
cpera of element

Type float
Default 0.0

conf

conf of element

Type str**Default** 1s2_2s1**r_rms**

r_rms of element

Type float**Default** 4.0992133976e-05**be**

data of element

Keywords**vdw-radius**

radius of element

Type float**Default** 1.53**covalent**

covalent value element

Type float**Default** 0.9**Z**

z-value of element

Type int**Default** 4**mass**

mass of element

Type float**Default** 9.012182**symbol**

symbol of element

Type str**Default** Be**bpt**

bpt of element

Type float**Default** 2745.0**mpt**

mpt of element

Type float**Default** 1560.0

density
density of element

Type float

Default 1.85

volume
volume of element

Type float

Default 5.0

name
name of element

Type str

Default Beryllium

debye
debye of element

Type float

Default 1000.0

a
a of element

Type float

Default 2.29

crystal
crystal of element

Type str

Default HEX

cpera
cpera of element

Type float

Default 1.567

conf
conf of element

Type str

Default 1s2_2s2

r_rms
r_rms of element

Type float

Default 4.3632829651e-05

b
data of element

Keywords

vdw-radius
radius of element
Type float
Default 1.92

covalent
covalent value element
Type float
Default 0.82

Z
z-value of element
Type int
Default 5

mass
mass of element
Type float
Default 10.811

symbol
symbol of element
Type str
Default B

bpt
bpt of element
Type float
Default 4275.0

mpt
mpt of element
Type float
Default 2300.0

density
density of element
Type float
Default 2.34

volume
volume of element
Type float
Default 4.6

name
name of element
Type str

Default Boron

debye
debye of element
Type float
Default 1250.0

a
a of element
Type float
Default 8.73

crystal
crystal of element
Type str
Default TET

cpera
cpera of element
Type float
Default 0.576

conf
conf of element
Type str
Default 1s2_2s2_2p1

r_rms
r_rms of element
Type float
Default 4.5906118608e-05

c
data of element

Keywords

vdw-radius
radius of element
Type float
Default 1.7

covalent
covalent value element
Type float
Default 0.77

Z
z-value of element
Type int

Default 6**mass**

mass of element

Type float**Default 12.011****symbol**

symbol of element

Type str**Default C****bpt**

bpt of element

Type float**Default 4470.0****mpt**

mpt of element

Type float**Default 4100.0****density**

density of element

Type float**Default 2.62****volume**

volume of element

Type float**Default 4.58****name**

name of element

Type str**Default Carbon****debye**

debye of element

Type float**Default 1860.0****a**

a of element

Type float**Default 3.57****crystal**

crystal of element

Type str

Default DIA

cpera

cpera of element

Type float

Default 0.0

conf

conf of element

Type str

Default 1s2_2s2_2p2

r_rms

r_rms of element

Type float

Default 4.6940079496e-05

n

data of element

Keywords

vdw-radius

radius of element

Type float

Default 1.55

covalent

covalent value element

Type float

Default 0.75

Z

z-value of element

Type int

Default 7

mass

mass of element

Type float

Default 14.00674

symbol

symbol of element

Type str

Default N

bpt

bpt of element

Type float
Default 77.35

mpt
mpt of element

Type float
Default 63.14

density
density of element

Type float
Default 1.251

volume
volume of element

Type float
Default 17.3

name
name of element

Type str
Default Nitrogen

debye
debye of element

Type float
Default -79.0

a
a of element

Type float
Default 4.039

crystal
crystal of element

Type str
Default HEX

cpera
cpera of element

Type float
Default 1.651

conf
conf of element

Type str
Default 1s2_2s2_2p3

r_rms

r_rms of element

Type float

Default 4.8847128967e-05

o

data of element

Keywords

vdw-radius

radius of element

Type float

Default 1.52

covalent

covalent value element

Type float

Default 0.73

Z

z-value of element

Type int

Default 8

mass

mass of element

Type float

Default 15.9994

symbol

symbol of element

Type str

Default 0

bpt

bpt of element

Type float

Default 90.18

mpt

mpt of element

Type float

Default 50.35

density

density of element

Type float

Default 1.429

volume
volume of element

Type float

Default 14.0

name
name of element

Type str

Default Oxygen

debye
debye of element

Type float

Default -46.0

a
a of element

Type float

Default 6.83

crystal
crystal of element

Type str

Default CUB

cpera
cpera of element

Type float

Default 0.0

conf
conf of element

Type str

Default 1s2_2s2_2p4

r_rms
r_rms of element

Type float

Default 5.0580178957e-05

f
data of element

Keywords

vdw-radius
radius of element

Type float

Default 1.47

covalent
covalent value element
Type float
Default 0.72

Z
z-value of element
Type int
Default 9

mass
mass of element
Type float
Default 18.9984032

symbol
symbol of element
Type str
Default F

bpt
bpt of element
Type float
Default 84.95

mpt
mpt of element
Type float
Default 53.48

density
density of element
Type float
Default 1.696

volume
volume of element
Type float
Default 17.1

name
name of element
Type str
Default Fluorine

debye
debye of element
Type float

Default 0.0

a
a of element

Type float

Default 0.0

crystal
crystal of element

Type str

Default MCL

cpera
cpera of element

Type float

Default 0.0

conf
conf of element

Type str

Default 1s2_2s2_2p5

r_rms
r_rms of element

Type float

Default 5.2927138943e-05

ne
data of element

Keywords

vdw-radius
radius of element

Type float

Default 1.54

covalent
covalent value element

Type float

Default 0.71

Z
z-value of element

Type int

Default 10

mass
mass of element

Type float

Default 20.1797

symbol

symbol of element

Type str

Default Ne

bpt

bpt of element

Type float

Default 27.096

mpt

mpt of element

Type float

Default 24.553

density

density of element

Type float

Default 0.901

volume

volume of element

Type float

Default 16.7

name

name of element

Type str

Default Neon

debye

debye of element

Type float

Default 63.0

a

a of element

Type float

Default 4.43

crystal

crystal of element

Type str

Default FCC

cpera

cpera of element

Type float

Default 0.0

conf

conf of element

Type str

Default 1s2_2s2_2p6

r_rms

r_rms of element

Type float

Default 5.3654104231e-05

na

data of element

Keywords

vdw-radius

radius of element

Type float

Default 2.27

covalent

covalent value element

Type float

Default 1.54

Z

z-value of element

Type int

Default 11

mass

mass of element

Type float

Default 22.989768

symbol

symbol of element

Type str

Default Na

bpt

bpt of element

Type float

Default 1156.0

mpt

mpt of element

Type float
Default 371.0

density
density of element
Type float
Default 0.97

volume
volume of element
Type float
Default 23.7

name
name of element
Type str
Default Sodium

debye
debye of element
Type float
Default 150.0

a
a of element
Type float
Default 4.23

crystal
crystal of element
Type str
Default BCC

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default [Ne]3s1

r_rms
r_rms of element
Type float
Default 5.5699159416e-05

mg
data of element

Keywords

vdw-radius
radius of element

Type float

Default 1.73

covalent
covalent value element

Type float

Default 1.36

Z
z-value of element

Type int

Default 12

mass
mass of element

Type float

Default 24.305

symbol
symbol of element

Type str

Default Mg

bpt
bpt of element

Type float

Default 1363.0

mpt
mpt of element

Type float

Default 922.0

density
density of element

Type float

Default 1.74

volume
volume of element

Type float

Default 13.97

name
name of element
Type str
Default Magnesium

debye
debye of element
Type float
Default 318.0

a
a of element
Type float
Default 3.21

crystal
crystal of element
Type str
Default HEX

cpera
cpera of element
Type float
Default 1.624

conf
conf of element
Type str
Default [Ne]3s2

r_rms
r_rms of element
Type float
Default 5.6341070732e-05

al
data of element

Keywords

vdw-radius
radius of element
Type float
Default 1.84

covalent
covalent value element
Type float
Default 1.18

Z
z-value of element
Type int
Default 13

mass
mass of element
Type float
Default 26.981539

symbol
symbol of element
Type str
Default Al

bpt
bpt of element
Type float
Default 2793.0

mpt
mpt of element
Type float
Default 933.25

density
density of element
Type float
Default 2.7

volume
volume of element
Type float
Default 10.0

name
name of element
Type str
Default Aluminum

debye
debye of element
Type float
Default 394.0

a
a of element
Type float

Default 4.05

crystal

crystal of element

Type str

Default FCC

cpera

cpera of element

Type float

Default 0.0

conf

conf of element

Type str

Default [Ne]3s2_3p1

r_rms

r_rms of element

Type float

Default 5.8165765928e-05

si

data of element

Keywords

vdw-radius

radius of element

Type float

Default 2.1

covalent

covalent value element

Type float

Default 1.11

Z

z-value of element

Type int

Default 14

mass

mass of element

Type float

Default 28.0855

symbol

symbol of element

Type str

Default Si

bpt
bpt of element

Type float

Default 3540.0

mpt
mpt of element

Type float

Default 1685.0

density
density of element

Type float

Default 2.33

volume
volume of element

Type float

Default 12.1

name
name of element

Type str

Default Silicon

debye
debye of element

Type float

Default 625.0

a
a of element

Type float

Default 5.43

crystal
crystal of element

Type str

Default DIA

cpera
cpera of element

Type float

Default 0.0

conf
conf of element

Type str
Default [Ne]3s2_3p2

r_rms
r_rms of element

Type float
Default 5.8743802504e-05

p
data of element

Keywords

vdw-radius
radius of element

Type float
Default 1.8

covalent
covalent value element

Type float
Default 1.06

Z
z-value of element

Type int
Default 15

mass
mass of element

Type float
Default 30.97362

symbol
symbol of element

Type str
Default P

bpt
bpt of element

Type float
Default 550.0

mpt
mpt of element

Type float
Default 317.3

density
density of element

Type float

Default 1.82

volume

volume of element

Type float

Default 17.0

name

name of element

Type str

Default Phosphorus

debye

debye of element

Type float

Default 0.0

a

a of element

Type float

Default 7.17

crystal

crystal of element

Type str

Default CUB

cpera

cpera of element

Type float

Default 0.0

conf

conf of element

Type str

Default [Ne]3s2_3p3

r_rms

r_rms of element

Type float

Default 6.0399312923e-05

s

data of element

Keywords

vdw-radius

radius of element

Type float
Default 1.8

covalent
covalent value element

Type float
Default 1.02

Z
z-value of element

Type int
Default 16

mass
mass of element

Type float
Default 32.066

symbol
symbol of element

Type str
Default S

bpt
bpt of element

Type float
Default 717.75

mpt
mpt of element

Type float
Default 388.36

density
density of element

Type float
Default 2.07

volume
volume of element

Type float
Default 15.5

name
name of element

Type str
Default Sulfur

debye
debye of element

Type float

Default 0.0

a
a of element

Type float

Default 10.47

crystal
crystal of element

Type str

Default ORC

cpera
cpera of element

Type float

Default 0.0

conf
conf of element

Type str

Default [Ne]3s2_3p4

r_rms
r_rms of element

Type float

Default 6.0927308666e-05

cl
data of element

Keywords

vdw-radius
radius of element

Type float

Default 1.75

covalent
covalent value element

Type float

Default 0.99

Z
z-value of element

Type int

Default 17

mass
mass of element
Type float
Default 35.4527

symbol
symbol of element
Type str
Default Cl

bpt
bpt of element
Type float
Default 239.1

mpt
mpt of element
Type float
Default 172.16

density
density of element
Type float
Default 3.17

volume
volume of element
Type float
Default 22.7

name
name of element
Type str
Default Chlorine

debye
debye of element
Type float
Default 0.0

a
a of element
Type float
Default 6.24

crystal
crystal of element
Type str

Default ORC

cpera
cpera of element

Type float

Default 0.0

conf
conf of element

Type str

Default [Ne]3s2_3p5

r_rms
r_rms of element

Type float

Default 6.2448101115e-05

ar
data of element

Keywords

vdw-radius
radius of element

Type float

Default 1.88

covalent
covalent value element

Type float

Default 0.98

Z
z-value of element

Type int

Default 18

mass
mass of element

Type float

Default 39.948

symbol
symbol of element

Type str

Default Ar

bpt
bpt of element

Type float

Default 87.3

mpt
mpt of element
Type float
Default 83.81

density
density of element
Type float
Default 1.784

volume
volume of element
Type float
Default 28.5

name
name of element
Type str
Default Argon

debye
debye of element
Type float
Default 85.0

a
a of element
Type float
Default 5.26

crystal
crystal of element
Type str
Default FCC

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default [Ne]3s2_3p6

r_rms
r_rms of element

Type float

Default 6.4800211825e-05

k

data of element

Keywords

vdw-radius

radius of element

Type float

Default 2.75

covalent

covalent value element

Type float

Default 2.03

Z

z-value of element

Type int

Default 19

mass

mass of element

Type float

Default 39.0983

symbol

symbol of element

Type str

Default K

bpt

bpt of element

Type float

Default 1032.0

mpt

mpt of element

Type float

Default 336.35

density

density of element

Type float

Default 0.86

volume

volume of element

Type float
Default 45.46

name
name of element

Type str
Default Potassium

debye
debye of element

Type float
Default 100.0

a
a of element

Type float
Default 5.23

crystal
crystal of element

Type str
Default BCC

cpera
cpera of element

Type float
Default 0.0

conf
conf of element

Type str
Default [Ar]4s1

r_rms
r_rms of element

Type float
Default 6.4346167051e-05

ca
data of element

Keywords

vdw-radius
radius of element

Type float
Default 2.31

covalent
covalent value element

Type float
Default 1.91

z
z-value of element

Type int
Default 20

mass
mass of element

Type float
Default 40.078

symbol
symbol of element

Type str
Default Ca

bpt
bpt of element

Type float
Default 1757.0

mpt
mpt of element

Type float
Default 1112.0

density
density of element

Type float
Default 1.55

volume
volume of element

Type float
Default 29.9

name
name of element

Type str
Default Calcium

debye
debye of element

Type float
Default 230.0

a
a of element

Type float

Default 5.58

crystal
crystal of element

Type str

Default FCC

cpera
cpera of element

Type float

Default 0.0

conf
conf of element

Type str

Default [Ar]4s2

r_rms
r_rms of element

Type float

Default 6.4800211825e-05

sc
data of element

Keywords

vdw-radius
radius of element

Type float

Default -1.0

covalent
covalent value element

Type float

Default 1.62

Z
z-value of element

Type int

Default 21

mass
mass of element

Type float

Default 44.95591

symbol
symbol of element
Type str
Default Sc

bpt
bpt of element
Type float
Default 3104.0

mpt
mpt of element
Type float
Default 1812.0

density
density of element
Type float
Default 3.0

volume
volume of element
Type float
Default 15.0

name
name of element
Type str
Default Scandium

debye
debye of element
Type float
Default -359.0

a
a of element
Type float
Default 3.31

crystal
crystal of element
Type str
Default HEX

cpera
cpera of element
Type float

Default 1.594

conf

conf of element

Type str

Default [Ar]3d1_4s2

r_rms

r_rms of element

Type float

Default 6.6963627201e-05

ti

data of element

Keywords

vdw-radius

radius of element

Type float

Default -1.0

covalent

covalent value element

Type float

Default 1.45

Z

z-value of element

Type int

Default 22

mass

mass of element

Type float

Default 47.88

symbol

symbol of element

Type str

Default Ti

bpt

bpt of element

Type float

Default 3562.0

mpt

mpt of element

Type float

Default 1943.0

density
density of element
Type float
Default 4.5

volume
volume of element
Type float
Default 10.64

name
name of element
Type str
Default Titanium

debye
debye of element
Type float
Default 380.0

a
a of element
Type float
Default 2.95

crystal
crystal of element
Type str
Default HEX

cpera
cpera of element
Type float
Default 1.588

conf
conf of element
Type str
Default [Ar]3d2_4s2

r_rms
r_rms of element
Type float
Default 6.818557748e-05

v
data of element

Keywords

vdw-radius
radius of element

Type float

Default -1.0

covalent
covalent value element

Type float

Default 1.34

Z
z-value of element

Type int

Default 23

mass
mass of element

Type float

Default 50.9415

symbol
symbol of element

Type str

Default V

bpt
bpt of element

Type float

Default 3682.0

mpt
mpt of element

Type float

Default 2175.0

density
density of element

Type float

Default 5.8

volume
volume of element

Type float

Default 8.78

name
name of element

Type str
Default Vanadium

debye
debye of element

Type float
Default 390.0

a
a of element

Type float
Default 3.02

crystal
crystal of element

Type str
Default BCC

cpera
cpera of element

Type float
Default 0.0

conf
conf of element

Type str
Default [Ar]3d3_4s2

r_rms
r_rms of element

Type float
Default 6.935761683e-05

cr
data of element

Keywords

vdw-radius
radius of element

Type float
Default -1.0

covalent
covalent value element

Type float
Default 1.18

Z
z-value of element

Type int
Default 24

mass
mass of element

Type float
Default 51.9961

symbol
symbol of element

Type str
Default Cr

bpt
bpt of element

Type float
Default 2945.0

mpt
mpt of element

Type float
Default 2130.0

density
density of element

Type float
Default 7.19

volume
volume of element

Type float
Default 7.23

name
name of element

Type str
Default Chromium

debye
debye of element

Type float
Default 460.0

a
a of element

Type float
Default 2.88

crystal
crystal of element

Type str

Default BCC

cpera
cpera of element

Type float

Default 0.0

conf
conf of element

Type str

Default [Ar]3d5_4s1

r_rms
r_rms of element

Type float

Default 6.9738057221e-05

mn
data of element

Keywords

vdw-radius
radius of element

Type float

Default -1.0

covalent
covalent value element

Type float

Default 1.17

Z
z-value of element

Type int

Default 25

mass
mass of element

Type float

Default 54.93085

symbol
symbol of element

Type str

Default Mn

bpt
bpt of element
Type float
Default 2335.0

mpt
mpt of element
Type float
Default 1517.0

density
density of element
Type float
Default 7.43

volume
volume of element
Type float
Default 1.39

name
name of element
Type str
Default Manganese

debye
debye of element
Type float
Default 400.0

a
a of element
Type float
Default 8.89

crystal
crystal of element
Type str
Default CUB

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str

Default [Ar]3d5_4s2

r_rms

r_rms of element

Type float

Default 7.0850896638e-05

fe

data of element

Keywords

vdw-radius

radius of element

Type float

Default -1.0

covalent

covalent value element

Type float

Default 1.17

Z

z-value of element

Type int

Default 26

mass

mass of element

Type float

Default 55.847

symbol

symbol of element

Type str

Default Fe

bpt

bpt of element

Type float

Default 3135.0

mpt

mpt of element

Type float

Default 1809.0

density

density of element

Type float

Default 7.86

volume
volume of element
Type float
Default 7.1

name
name of element
Type str
Default Iron

debye
debye of element
Type float
Default 460.0

a
a of element
Type float
Default 2.87

crystal
crystal of element
Type str
Default BCC

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default [Ar]3d6_4s2

r_rms
r_rms of element
Type float
Default 7.1212829817e-05

co
data of element

Keywords

vdw-radius
radius of element
Type float

Default -1.0
covalent
 covalent value element
Type float
Default 1.16
Z
 z-value of element
Type int
Default 27
mass
 mass of element
Type float
Default 58.9332
symbol
 symbol of element
Type str
Default Co
bpt
 bpt of element
Type float
Default 3201.0
mpt
 mpt of element
Type float
Default 1768.0
density
 density of element
Type float
Default 8.9
volume
 volume of element
Type float
Default 6.7
name
 name of element
Type str
Default Cobalt
debye
 debye of element

Type float

Default 385.0

a

a of element

Type float

Default 2.51

crystal

crystal of element

Type str

Default HEX

cpera

cpera of element

Type float

Default 0.0

conf

conf of element

Type str

Default [Ar]3d7_4s2

r_rms

r_rms of element

Type float

Default 7.2273420879e-05

ni

data of element

Keywords

vdw-radius

radius of element

Type float

Default -1.0

covalent

covalent value element

Type float

Default 1.15

Z

z-value of element

Type int

Default 28

mass

mass of element

Type float
Default 58.69

symbol
symbol of element

Type str
Default Ni

bpt
bpt of element

Type float
Default 3187.0

mpt
mpt of element

Type float
Default 1726.0

density
density of element

Type float
Default 8.9

volume
volume of element

Type float
Default 6.59

name
name of element

Type str
Default Nickel

debye
debye of element

Type float
Default 375.0

a
a of element

Type float
Default 3.52

crystal
crystal of element

Type str
Default FCC

cpera
cpera of element

Type float

Default 0.0

conf
conf of element

Type str

Default [Ar]3d8_4s2

r_rms
r_rms of element

Type float

Default 7.1923970253e-05

cu
data of element

Keywords

vdw-radius
radius of element

Type float

Default -1.0

covalent
covalent value element

Type float

Default 1.17

Z
z-value of element

Type int

Default 29

mass
mass of element

Type float

Default 63.546

symbol
symbol of element

Type str

Default Cu

bpt
bpt of element

Type float

Default 2836.0

mpt
mpt of element
Type float
Default 1357.6

density
density of element
Type float
Default 8.96

volume
volume of element
Type float
Default 7.1

name
name of element
Type str
Default Copper

debye
debye of element
Type float
Default 315.0

a
a of element
Type float
Default 3.61

crystal
crystal of element
Type str
Default FCC

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default [Ar]3d10_4s1

r_rms
r_rms of element
Type float

Default 7.3633018675e-05

zn

data of element

Keywords**vdw-radius**

radius of element

Type float

Default -1.0

covalent

covalent value element

Type float

Default 1.25

Z

z-value of element

Type int

Default 30

mass

mass of element

Type float

Default 65.39

symbol

symbol of element

Type str

Default Zn

bpt

bpt of element

Type float

Default 1180.0

mpt

mpt of element

Type float

Default 692.73

density

density of element

Type float

Default 7.14

volume

volume of element

Type float

Default 9.2

name
name of element

Type str

Default Zinc

debye
debye of element

Type float

Default 234.0

a
a of element

Type float

Default 2.66

crystal
crystal of element

Type str

Default HEX

cpera
cpera of element

Type float

Default 0.0

conf
conf of element

Type str

Default [Ar]3d10_4s2

r_rms
r_rms of element

Type float

Default 7.3963875193e-05

ga
data of element

Keywords

vdw-radius
radius of element

Type float

Default 1.87

covalent
covalent value element

Type float

Default 1.26

Z
z-value of element
Type int
Default 31

mass
mass of element
Type float
Default 69.723

symbol
symbol of element
Type str
Default Ga

bpt
bpt of element
Type float
Default 2478.0

mpt
mpt of element
Type float
Default 302.9

density
density of element
Type float
Default 5.91

volume
volume of element
Type float
Default 11.8

name
name of element
Type str
Default Gallium

debye
debye of element
Type float
Default 240.0

a
a of element

Type float
Default 4.51

crystal
crystal of element

Type str
Default ORC

cpera
cpera of element

Type float
Default 0.0

conf
conf of element

Type str
Default [Ar]3d10_4s2_4p1

r_rms
r_rms of element

Type float
Default 7.5568424848e-05

ge
data of element

Keywords

vdw-radius
radius of element

Type float
Default 2.11

covalent
covalent value element

Type float
Default 1.22

Z
z-value of element

Type int
Default 32

mass
mass of element

Type float
Default 72.61

symbol
symbol of element

Type str
Default Ge

bpt
bpt of element

Type float
Default 3107.0

mpt
mpt of element

Type float
Default 1210.4

density
density of element

Type float
Default 5.32

volume
volume of element

Type float
Default 13.6

name
name of element

Type str
Default Germanium

debye
debye of element

Type float
Default 360.0

a
a of element

Type float
Default 5.66

crystal
crystal of element

Type str
Default DIA

cpera
cpera of element

Type float
Default 0.0

conf
conf of element
Type str
Default [Ar]3d10_4s2_4p2

r_rms
r_rms of element
Type float
Default 7.7097216161e-05

as
data of element

Keywords

vdw-radius
radius of element
Type float
Default 1.85

covalent
covalent value element
Type float
Default 1.2

Z
z-value of element
Type int
Default 33

mass
mass of element
Type float
Default 74.92159

symbol
symbol of element
Type str
Default As

bpt
bpt of element
Type float
Default 876.0

mpt
mpt of element
Type float
Default 1081.0

density
density of element
Type float
Default 5.72

volume
volume of element
Type float
Default 13.1

name
name of element
Type str
Default Arsenic

debye
debye of element
Type float
Default 285.0

a
a of element
Type float
Default 4.13

crystal
crystal of element
Type str
Default RHL

cpera
cpera of element
Type float
Default 54.16

conf
conf of element
Type str
Default [Ar]3d10_4s2_4p3

r_rms
r_rms of element
Type float
Default 7.7394645153e-05

se
data of element

Keywords

vdw-radius
radius of element
Type float
Default 1.9

covalent
covalent value element
Type float
Default 1.16

Z
z-value of element
Type int
Default 34

mass
mass of element
Type float
Default 78.96

symbol
symbol of element
Type str
Default Se

bpt
bpt of element
Type float
Default 958.0

mpt
mpt of element
Type float
Default 494.0

density
density of element
Type float
Default 4.8

volume
volume of element
Type float
Default 16.45

name
name of element
Type str

Default Selenium

debye
debye of element
Type float
Default -150.0

a
a of element
Type float
Default 4.36

crystal
crystal of element
Type str
Default HEX

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default [Ar]3d10_4s2_4p4

r_rms
r_rms of element
Type float
Default 7.8843427408e-05

br
data of element

Keywords

vdw-radius
radius of element
Type float
Default 1.83

covalent
covalent value element
Type float
Default 1.14

Z
z-value of element
Type int

Default 35

mass

mass of element

Type float

Default 79.904

symbol

symbol of element

Type str

Default Br

bpt

bpt of element

Type float

Default 332.25

mpt

mpt of element

Type float

Default 265.9

density

density of element

Type float

Default 3.12

volume

volume of element

Type float

Default 23.5

name

name of element

Type str

Default Bromine

debye

debye of element

Type float

Default 0.0

a

a of element

Type float

Default 6.67

crystal

crystal of element

Type str
Default ORC

cpera
cpera of element

Type float
Default 0.0

conf
conf of element

Type str
Default [Ar]3d10_4s2_4p5

r_rms
r_rms of element

Type float
Default 7.8558604038e-05

kr
data of element

Keywords

vdw-radius
radius of element

Type float
Default 2.02

covalent
covalent value element

Type float
Default 1.12

Z
z-value of element

Type int
Default 36

mass
mass of element

Type float
Default 83.8

symbol
symbol of element

Type str
Default Kr

bpt
bpt of element

Type float
Default 119.8

mpt
mpt of element

Type float
Default 115.78

density
density of element

Type float
Default 3.74

volume
volume of element

Type float
Default 38.9

name
name of element

Type str
Default Krypton

debye
debye of element

Type float
Default -73.0

a
a of element

Type float
Default 5.72

crystal
crystal of element

Type str
Default FCC

cpera
cpera of element

Type float
Default 0.0

conf
conf of element

Type str
Default [Ar]3d10_4s2_4p6

r_rms
r_rms of element
Type float
Default 7.9959560033e-05

rb
data of element

Keywords

vdw-radius
radius of element

Type float
Default 3.03

covalent
covalent value element

Type float
Default 2.16

Z
z-value of element

Type int
Default 37

mass
mass of element

Type float
Default 85.4678

symbol
symbol of element

Type str
Default Rb

bpt
bpt of element

Type float
Default 961.0

mpt
mpt of element

Type float
Default 312.64

density
density of element

Type float
Default 1.53

volume
volume of element
Type float
Default 55.9

name
name of element
Type str
Default Rubidium

debye
debye of element
Type float
Default -56.0

a
a of element
Type float
Default 5.59

crystal
crystal of element
Type str
Default BCC

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default [Kr]5s1

r_rms
r_rms of element
Type float
Default 8.0233033713e-05

sr
data of element

Keywords

vdw-radius
radius of element
Type float
Default 2.49

covalent
covalent value element
Type float
Default 1.91

Z
z-value of element
Type int
Default 38

mass
mass of element
Type float
Default 87.62

symbol
symbol of element
Type str
Default Sr

bpt
bpt of element
Type float
Default 1650.0

mpt
mpt of element
Type float
Default 1041.0

density
density of element
Type float
Default 2.6

volume
volume of element
Type float
Default 33.7

name
name of element
Type str
Default Strontium

debye
debye of element
Type float

Default -147.0

a

a of element

Type float

Default 6.08

crystal

crystal of element

Type str

Default FCC

cpera

cpera of element

Type float

Default 0.0

conf

conf of element

Type str

Default [Kr]5s2

r_rms

r_rms of element

Type float

Default 8.1040799081e-05

y

data of element

Keywords

vdw-radius

radius of element

Type float

Default -1.0

covalent

covalent value element

Type float

Default 1.62

Z

z-value of element

Type int

Default 39

mass

mass of element

Type float

Default 88.90585

symbol
symbol of element

Type str

Default Y

bpt
bpt of element

Type float

Default 3611.0

mpt
mpt of element

Type float

Default 1799.0

density
density of element

Type float

Default 4.5

volume
volume of element

Type float

Default 19.8

name
name of element

Type str

Default Yttrium

debye
debye of element

Type float

Default -256.0

a
a of element

Type float

Default 3.65

crystal
crystal of element

Type str

Default HEX

cpera
cpera of element

Type float
Default 1.571

conf
conf of element

Type str
Default [Kr]4d1_5s2

r_rms
r_rms of element

Type float
Default 8.1305968993e-05

zr
data of element

Keywords

vdw-radius
radius of element

Type float
Default -1.0

covalent
covalent value element

Type float
Default 1.45

Z
z-value of element

Type int
Default 40

mass
mass of element

Type float
Default 91.224

symbol
symbol of element

Type str
Default Zr

bpt
bpt of element

Type float
Default 4682.0

mpt
mpt of element

Type float
Default 2125.0

density
density of element
Type float
Default 6.49

volume
volume of element
Type float
Default 14.1

name
name of element
Type str
Default Zirconium

debye
debye of element
Type float
Default 250.0

a
a of element
Type float
Default 3.23

crystal
crystal of element
Type str
Default HEX

cpera
cpera of element
Type float
Default 1.593

conf
conf of element
Type str
Default [Kr]4d2_5s2

r_rms
r_rms of element
Type float
Default 8.156915998e-05

nb

data of element

Keywords**vdw-radius**

radius of element

Type float

Default -1.0

covalent

covalent value element

Type float

Default 1.34

Z

z-value of element

Type int

Default 41

mass

mass of element

Type float

Default 92.90638

symbol

symbol of element

Type str

Default Nb

bpt

bpt of element

Type float

Default 5017.0

mpt

mpt of element

Type float

Default 2740.0

density

density of element

Type float

Default 8.55

volume

volume of element

Type float

Default 10.87

name
name of element
Type str
Default Niobium

debye
debye of element
Type float
Default 275.0

a
a of element
Type float
Default 3.3

crystal
crystal of element
Type str
Default BCC

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default [Kr]4d4_5s1

r_rms
r_rms of element
Type float
Default 8.2347219223e-05

mo
data of element

Keywords

vdw-radius
radius of element
Type float
Default -1.0

covalent
covalent value element
Type float
Default 1.3

Z
z-value of element
Type int
Default 42

mass
mass of element
Type float
Default 95.94

symbol
symbol of element
Type str
Default Mo

bpt
bpt of element
Type float
Default 4912.0

mpt
mpt of element
Type float
Default 2890.0

density
density of element
Type float
Default 10.2

volume
volume of element
Type float
Default 9.4

name
name of element
Type str
Default Molybdenum

debye
debye of element
Type float
Default 380.0

a
a of element
Type float

Default 3.15

crystal

crystal of element

Type str

Default BCC

cpera

cpera of element

Type float

Default 0.0

conf

conf of element

Type str

Default [Kr]4d5_5s1

r_rms

r_rms of element

Type float

Default 8.3607614434e-05

tc

data of element

Keywords

vdw-radius

radius of element

Type float

Default -1.0

covalent

covalent value element

Type float

Default 1.27

Z

z-value of element

Type int

Default 43

mass

mass of element

Type float

Default -98.0

symbol

symbol of element

Type str

Default Tc
bpt
 bpt of element
Type float
Default 4538.0

mpt
 mpt of element
Type float
Default 2473.0

density
 density of element
Type float
Default 11.5

volume
 volume of element
Type float
Default 8.5

name
 name of element
Type str
Default Technetium

debye
 debye of element
Type float
Default 0.0

a
 a of element
Type float
Default 2.74

crystal
 crystal of element
Type str
Default HEX

cpera
 cpera of element
Type float
Default 1.604

conf
 conf of element

Type str
Default [Kr]4d5_5s2

r_rms
r_rms of element
Type float
Default 8.3607614434e-05

ru
data of element

Keywords

vdw-radius
radius of element
Type float
Default -1.0

covalent
covalent value element
Type float
Default 1.25

Z
z-value of element
Type int
Default 44

mass
mass of element
Type float
Default 101.07

symbol
symbol of element
Type str
Default Ru

bpt
bpt of element
Type float
Default 4423.0

mpt
mpt of element
Type float
Default 2523.0

density
density of element

Type float

Default 12.2

volume

volume of element

Type float

Default 8.3

name

name of element

Type str

Default Ruthenium

debye

debye of element

Type float

Default -382.0

a

a of element

Type float

Default 2.7

crystal

crystal of element

Type str

Default HEX

cpera

cpera of element

Type float

Default 1.584

conf

conf of element

Type str

Default [Kr]4d7_5s1

r_rms

r_rms of element

Type float

Default 8.4585397905e-05

rh

data of element

Keywords

vdw-radius

radius of element

Type float
Default -1.0

covalent
covalent value element

Type float
Default 1.25

Z
z-value of element

Type int
Default 45

mass
mass of element

Type float
Default 102.9055

symbol
symbol of element

Type str
Default Rh

bpt
bpt of element

Type float
Default 3970.0

mpt
mpt of element

Type float
Default 2236.0

density
density of element

Type float
Default 12.4

volume
volume of element

Type float
Default 8.3

name
name of element

Type str
Default Rhodium

debye
debye of element
Type float
Default -350.0

a
a of element
Type float
Default 3.8

crystal
crystal of element
Type str
Default FCC

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default [Kr]4d8_5s1

r_rms
r_rms of element
Type float
Default 8.4825835954e-05

pd
data of element
Keywords

vdw-radius
radius of element
Type float
Default -1.0

covalent
covalent value element
Type float
Default 1.28

Z
z-value of element
Type int
Default 46

mass
mass of element
Type float
Default 106.42

symbol
symbol of element
Type str
Default Pd

bpt
bpt of element
Type float
Default 3237.0

mpt
mpt of element
Type float
Default 1825.0

density
density of element
Type float
Default 12.0

volume
volume of element
Type float
Default 8.9

name
name of element
Type str
Default Palladium

debye
debye of element
Type float
Default 275.0

a
a of element
Type float
Default 3.89

crystal
crystal of element
Type str

Default FCC

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default [Kr]4d10_5s0

r_rms
r_rms of element
Type float
Default 8.5537941156e-05

ag
data of element

Keywords

vdw-radius
radius of element
Type float
Default -1.0

covalent
covalent value element
Type float
Default 1.34

Z
z-value of element
Type int
Default 47

mass
mass of element
Type float
Default 107.8682

symbol
symbol of element
Type str
Default Ag

bpt
bpt of element
Type float

Default 2436.0

mpt
mpt of element
Type float
Default 1234.0

density
density of element
Type float
Default 10.5

volume
volume of element
Type float
Default 10.3

name
name of element
Type str
Default Silver

debye
debye of element
Type float
Default 215.0

a
a of element
Type float
Default 4.09

crystal
crystal of element
Type str
Default FCC

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default [Kr]4d10_5s1

r_rms
r_rms of element

Type float

Default 8.5772320442e-05

cd

data of element

Keywords

vdw-radius

radius of element

Type float

Default -1.0

covalent

covalent value element

Type float

Default 1.48

z

z-value of element

Type int

Default 48

mass

mass of element

Type float

Default 112.411

symbol

symbol of element

Type str

Default Cd

bpt

bpt of element

Type float

Default 1040.0

mpt

mpt of element

Type float

Default 594.18

density

density of element

Type float

Default 8.65

volume

volume of element

Type float
Default 13.1

name
name of element

Type str
Default Cadmium

debye
debye of element

Type float
Default 120.0

a
a of element

Type float
Default 2.98

crystal
crystal of element

Type str
Default HEX

cpera
cpera of element

Type float
Default 1.886

conf
conf of element

Type str
Default [Kr]4d10_5s2

r_rms
r_rms of element

Type float
Default 8.7373430179e-05

in
data of element

Keywords

vdw-radius
radius of element

Type float
Default 1.93

covalent
covalent value element

Type float
Default 1.44

z
z-value of element

Type int
Default 49

mass
mass of element

Type float
Default 114.82

symbol
symbol of element

Type str
Default In

bpt
bpt of element

Type float
Default 2346.0

mpt
mpt of element

Type float
Default 429.76

density
density of element

Type float
Default 7.31

volume
volume of element

Type float
Default 15.7

name
name of element

Type str
Default Indium

debye
debye of element

Type float
Default 129.0

a
a of element

Type float

Default 4.59

crystal
crystal of element

Type str

Default TET

cpera
cpera of element

Type float

Default 1.076

conf
conf of element

Type str

Default [Kr]4d10_5s2_5p1

r_rms
r_rms of element

Type float

Default 8.7596760865e-05

sn
data of element

Keywords

vdw-radius
radius of element

Type float

Default 2.17

covalent
covalent value element

Type float

Default 1.41

Z
z-value of element

Type int

Default 50

mass
mass of element

Type float

Default 118.71

symbol
symbol of element
Type str
Default Sn

bpt
bpt of element
Type float
Default 2876.0

mpt
mpt of element
Type float
Default 505.06

density
density of element
Type float
Default 7.3

volume
volume of element
Type float
Default 16.3

name
name of element
Type str
Default Tin

debye
debye of element
Type float
Default 170.0

a
a of element
Type float
Default 5.82

crystal
crystal of element
Type str
Default TET

cpera
cpera of element
Type float

Default 0.546

conf

conf of element

Type str

Default [Kr]4d10_5s2_5p2

r_rms

r_rms of element

Type float

Default 8.8694413774e-05

sb

data of element

Keywords

vdw-radius

radius of element

Type float

Default 2.06

covalent

covalent value element

Type float

Default 1.4

Z

z-value of element

Type int

Default 51

mass

mass of element

Type float

Default 121.75

symbol

symbol of element

Type str

Default Sb

bpt

bpt of element

Type float

Default 1860.0

mpt

mpt of element

Type float

Default 904.0

density
density of element
Type float
Default 6.68

volume
volume of element
Type float
Default 18.23

name
name of element
Type str
Default Antimony

debye
debye of element
Type float
Default 200.0

a
a of element
Type float
Default 4.51

crystal
crystal of element
Type str
Default RHL

cpera
cpera of element
Type float
Default 57.1

conf
conf of element
Type str
Default [Kr]4d10_5s2_5p3

r_rms
r_rms of element
Type float
Default 8.8910267995e-05

te
data of element

Keywords

vdw-radius
radius of element

Type float

Default 2.06

covalent
covalent value element

Type float

Default 1.36

Z
z-value of element

Type int

Default 52

mass
mass of element

Type float

Default 127.6

symbol
symbol of element

Type str

Default Te

bpt
bpt of element

Type float

Default 1261.0

mpt
mpt of element

Type float

Default 722.65

density
density of element

Type float

Default 6.24

volume
volume of element

Type float

Default 20.5

name
name of element

Type str
Default Tellurium

debye
debye of element

Type float
Default -139.0

a
a of element

Type float
Default 4.45

crystal
crystal of element

Type str
Default HEX

cpera
cpera of element

Type float
Default 1.33

conf
conf of element

Type str
Default [Kr]4d10_5s2_5p4

r_rms
r_rms of element

Type float
Default 9.0801452955e-05

i
data of element

Keywords

vdw-radius
radius of element

Type float
Default 1.98

covalent
covalent value element

Type float
Default 1.33

Z
z-value of element

Type int
Default 53

mass
mass of element

Type float
Default 126.90447

symbol
symbol of element

Type str
Default I

bpt
bpt of element

Type float
Default 458.4

mpt
mpt of element

Type float
Default 386.7

density
density of element

Type float
Default 4.92

volume
volume of element

Type float
Default 25.74

name
name of element

Type str
Default Iodine

debye
debye of element

Type float
Default 0.0

a
a of element

Type float
Default 7.27

crystal
crystal of element

Type str

Default ORC

cpera
cpera of element

Type float

Default 0.0

conf
conf of element

Type str

Default [Kr]4d10_5s2_5p5

r_rms
r_rms of element

Type float

Default 9.018104029e-05

xe
data of element

Keywords

vdw-radius
radius of element

Type float

Default 2.16

covalent
covalent value element

Type float

Default 1.31

Z
z-value of element

Type int

Default 54

mass
mass of element

Type float

Default 131.29

symbol
symbol of element

Type str

Default Xe

bpt
bpt of element
Type float
Default 165.03

mpt
mpt of element
Type float
Default 161.36

density
density of element
Type float
Default 5.89

volume
volume of element
Type float
Default 37.3

name
name of element
Type str
Default Xenon

debye
debye of element
Type float
Default -55.0

a
a of element
Type float
Default 6.2

crystal
crystal of element
Type str
Default FCC

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str

Default [Kr]4d10_5s2_5p6

r_rms

r_rms of element

Type float

Default 9.1209776425e-05

cs

data of element

Keywords

vdw-radius

radius of element

Type float

Default 3.43

covalent

covalent value element

Type float

Default 2.35

Z

z-value of element

Type int

Default 55

mass

mass of element

Type float

Default 132.90543

symbol

symbol of element

Type str

Default Cs

bpt

bpt of element

Type float

Default 944.0

mpt

mpt of element

Type float

Default 301.55

density

density of element

Type float

Default 1.87

volume
volume of element
Type float
Default 71.07

name
name of element
Type str
Default Cesium

debye
debye of element
Type float
Default -40.0

a
a of element
Type float
Default 6.05

crystal
crystal of element
Type str
Default BCC

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default [Xe]6s1

r_rms
r_rms of element
Type float
Default 9.1412392742e-05

ba
data of element

Keywords

vdw-radius
radius of element
Type float

Default 2.68
covalent
 covalent value element
Type float
Default 1.98
Z
 z-value of element
Type int
Default 56
mass
 mass of element
Type float
Default 137.327
symbol
 symbol of element
Type str
Default Ba
bpt
 bpt of element
Type float
Default 2171.0
mpt
 mpt of element
Type float
Default 1002.0
density
 density of element
Type float
Default 3.5
volume
 volume of element
Type float
Default 39.24
name
 name of element
Type str
Default Barium
debye
 debye of element

Type float
Default -110.0

a
a of element

Type float
Default 5.02

crystal
crystal of element

Type str
Default BCC

cpera
cpera of element

Type float
Default 0.0

conf
conf of element

Type str
Default [Xe]6s2

r_rms
r_rms of element

Type float
Default 9.2410525664e-05

la
data of element

Keywords

vdw-radius
radius of element

Type float
Default -1.0

covalent
covalent value element

Type float
Default 1.69

Z
z-value of element

Type int
Default 57

mass
mass of element

Type float
Default 138.9055

symbol
symbol of element

Type str
Default La

bpt
bpt of element

Type float
Default 3730.0

mpt
mpt of element

Type float
Default 1193.0

density
density of element

Type float
Default 6.7

volume
volume of element

Type float
Default 20.73

name
name of element

Type str
Default Lanthanum

debye
debye of element

Type float
Default 132.0

a
a of element

Type float
Default 3.75

crystal
crystal of element

Type str
Default HEX

cpera
cpera of element
Type float
Default 1.619

conf
conf of element
Type str
Default [Xe]5d1_6s2

r_rms
r_rms of element
Type float
Default 9.2607247118e-05

hf
data of element

Keywords

vdw-radius
radius of element
Type float
Default -1.0

covalent
covalent value element
Type float
Default 1.44

Z
z-value of element
Type int
Default 72

mass
mass of element
Type float
Default 178.49

symbol
symbol of element
Type str
Default Hf

bpt
bpt of element
Type float
Default 4876.0

mpt

mpt of element

Type float

Default 2500.0

density

density of element

Type float

Default 13.1

volume

volume of element

Type float

Default 13.6

name

name of element

Type str

Default Hafnium

debye

debye of element

Type float

Default 0.0

a

a of element

Type float

Default 3.2

crystal

crystal of element

Type str

Default HEX

cpera

cpera of element

Type float

Default 1.582

conf

conf of element

Type str

Default [Xe]4f14_5d2_6s2

r_rms

r_rms of element

Type float

Default 9.9970978172e-05

ta

data of element

Keywords**vdw-radius**

radius of element

Type float

Default -1.0

covalent

covalent value element

Type float

Default 1.34

Z

z-value of element

Type int

Default 73

mass

mass of element

Type float

Default 180.9479

symbol

symbol of element

Type str

Default Ta

bpt

bpt of element

Type float

Default 5731.0

mpt

mpt of element

Type float

Default 3287.0

density

density of element

Type float

Default 16.6

volume

volume of element

Type float

Default 10.9

name
name of element
Type str
Default Tantalum

debye
debye of element
Type float
Default 225.0

a
a of element
Type float
Default 3.31

crystal
crystal of element
Type str
Default BCC

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default [Xe]4f14_5d3_6s2

r_rms
r_rms of element
Type float
Default 0.00010013585755

w
data of element

Keywords

vdw-radius
radius of element
Type float
Default -1.0

covalent
covalent value element
Type float

Default 1.3

Z
z-value of element
Type int
Default 74

mass
mass of element
Type float
Default 183.85

symbol
symbol of element
Type str
Default W

bpt
bpt of element
Type float
Default 5828.0

mpt
mpt of element
Type float
Default 3680.0

density
density of element
Type float
Default 19.3

volume
volume of element
Type float
Default 9.53

name
name of element
Type str
Default Tungsten

debye
debye of element
Type float
Default 310.0

a
a of element

Type float
Default 3.16

crystal
crystal of element

Type str
Default BCC

cpera
cpera of element

Type float
Default 0.0

conf
conf of element

Type str
Default [Xe]4f14_5d4_6s2

r_rms
r_rms of element

Type float
Default 0.0001006268807

re
data of element

Keywords

vdw-radius
radius of element

Type float
Default -1.0

covalent
covalent value element

Type float
Default 1.28

Z
z-value of element

Type int
Default 75

mass
mass of element

Type float
Default 186.207

symbol
symbol of element

Type str
Default Re

bpt
bpt of element

Type float
Default 5869.0

mpt
mpt of element

Type float
Default 3453.0

density
density of element

Type float
Default 21.0

volume
volume of element

Type float
Default 8.85

name
name of element

Type str
Default Rhenium

debye
debye of element

Type float
Default 416.0

a
a of element

Type float
Default 2.76

crystal
crystal of element

Type str
Default HEX

cpera
cpera of element

Type float
Default 1.615

conf
conf of element
Type str
Default [Xe]4f14_5d5_6s2

r_rms
r_rms of element
Type float
Default 0.00010111259523

os
data of element

Keywords

vdw-radius
radius of element
Type float
Default -1.0

covalent
covalent value element
Type float
Default 1.26

Z
z-value of element
Type int
Default 76

mass
mass of element
Type float
Default 190.2

symbol
symbol of element
Type str
Default Os

bpt
bpt of element
Type float
Default 5285.0

mpt
mpt of element
Type float
Default 3300.0

density
density of element
Type float
Default 22.4

volume
volume of element
Type float
Default 8.49

name
name of element
Type str
Default Osmium

debye
debye of element
Type float
Default -400.0

a
a of element
Type float
Default 2.74

crystal
crystal of element
Type str
Default HEX

cpera
cpera of element
Type float
Default 1.579

conf
conf of element
Type str
Default [Xe]4f14_5d6_6s2

r_rms
r_rms of element
Type float
Default 0.00010191070333

ir
data of element

Keywords

vdw-radius
radius of element
Type float
Default -1.0

covalent
covalent value element
Type float
Default 1.27

Z
z-value of element
Type int
Default 77

mass
mass of element
Type float
Default 192.22

symbol
symbol of element
Type str
Default Ir

bpt
bpt of element
Type float
Default 4701.0

mpt
mpt of element
Type float
Default 2716.0

density
density of element
Type float
Default 22.5

volume
volume of element
Type float
Default 8.54

name
name of element
Type str

Default Iridium

debye
debye of element
Type float
Default 430.0

a
a of element
Type float
Default 3.84

crystal
crystal of element
Type str
Default FCC

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default [Xe]4f14_5d7_6s2

r_rms
r_rms of element
Type float
Default 0.00010206865731

pt
data of element

Keywords

vdw-radius
radius of element
Type float
Default -1.0

covalent
covalent value element
Type float
Default 1.3

Z
z-value of element
Type int

Default 78

mass

mass of element

Type float

Default 195.08

symbol

symbol of element

Type str

Default Pt

bpt

bpt of element

Type float

Default 4100.0

mpt

mpt of element

Type float

Default 2045.0

density

density of element

Type float

Default 21.4

volume

volume of element

Type float

Default 9.1

name

name of element

Type str

Default Platinum

debye

debye of element

Type float

Default 230.0

a

a of element

Type float

Default 3.92

crystal

crystal of element

Type str
Default FCC

cpera
cpera of element

Type float
Default 0.0

conf
conf of element

Type str
Default [Xe]4f14_5d10_6s0

r_rms
r_rms of element

Type float
Default 0.00010238293593

au
data of element

Keywords

vdw-radius
radius of element

Type float
Default -1.0

covalent
covalent value element

Type float
Default 1.34

Z
z-value of element

Type int
Default 79

mass
mass of element

Type float
Default 196.96654

symbol
symbol of element

Type str
Default Au

bpt
bpt of element

Type float
Default 3130.0

mpt
mpt of element

Type float
Default 1337.58

density
density of element

Type float
Default 19.3

volume
volume of element

Type float
Default 10.2

name
name of element

Type str
Default Gold

debye
debye of element

Type float
Default 170.0

a
a of element

Type float
Default 4.08

crystal
crystal of element

Type str
Default FCC

cpera
cpera of element

Type float
Default 0.0

conf
conf of element

Type str
Default [Xe]4f14_5d10_6s1

r_rms
r_rms of element
Type float
Default 0.00010269507292

hg
data of element

Keywords

vdw-radius
radius of element

Type float
Default -1.0

covalent
covalent value element

Type float
Default 1.49

Z
z-value of element

Type int
Default 80

mass
mass of element

Type float
Default 200.59

symbol
symbol of element

Type str
Default Hg

bpt
bpt of element

Type float
Default 630.0

mpt
mpt of element

Type float
Default 234.28

density
density of element

Type float
Default 13.53

volume
volume of element

Type float

Default 14.82

name
name of element

Type str

Default Mercury

debye
debye of element

Type float

Default 100.0

a
a of element

Type float

Default 2.99

crystal
crystal of element

Type str

Default RHL

cpera
cpera of element

Type float

Default 70.75

conf
conf of element

Type str

Default [Xe]4f14_5d10_6s2

r_rms
r_rms of element

Type float

Default 0.00010346628039

tl
data of element

Keywords

vdw-radius
radius of element

Type float

Default 1.96

covalent
covalent value element
Type float
Default 1.48

Z
z-value of element
Type int
Default 81

mass
mass of element
Type float
Default 204.3833

symbol
symbol of element
Type str
Default Tl

bpt
bpt of element
Type float
Default 1746.0

mpt
mpt of element
Type float
Default 577.0

density
density of element
Type float
Default 11.85

volume
volume of element
Type float
Default 17.2

name
name of element
Type str
Default Thallium

debye
debye of element
Type float

Default 96.0

a

a of element

Type float

Default 3.46

crystal

crystal of element

Type str

Default HEX

cpera

cpera of element

Type float

Default 1.599

conf

conf of element

Type str

Default [Xe]4f14_5d10_6s2_6p1

r_rms

r_rms of element

Type float

Default 0.00010392291259

pb

data of element

Keywords

vdw-radius

radius of element

Type float

Default 2.02

covalent

covalent value element

Type float

Default 1.47

Z

z-value of element

Type int

Default 82

mass

mass of element

Type float

Default 207.2

symbol

symbol of element

Type str

Default Pb

bpt

bpt of element

Type float

Default 2023.0

mpt

mpt of element

Type float

Default 600.6

density

density of element

Type float

Default 11.4

volume

volume of element

Type float

Default 18.17

name

name of element

Type str

Default Lead

debye

debye of element

Type float

Default 88.0

a

a of element

Type float

Default 4.95

crystal

crystal of element

Type str

Default FCC

cpera

cpera of element

Type float

Default 0.0

conf

conf of element

Type str

Default [Xe]4f14_5d10_6s2_6p2

r_rms

r_rms of element

Type float

Default 0.0001043751113

bi

data of element

Keywords

vdw-radius

radius of element

Type float

Default 2.07

covalent

covalent value element

Type float

Default 1.46

Z

z-value of element

Type int

Default 83

mass

mass of element

Type float

Default 208.98037

symbol

symbol of element

Type str

Default Bi

bpt

bpt of element

Type float

Default 1837.0

mpt

mpt of element

Type float
Default 544.52

density
density of element
Type float
Default 9.8

volume
volume of element
Type float
Default 21.3

name
name of element
Type str
Default Bismuth

debye
debye of element
Type float
Default 120.0

a
a of element
Type float
Default 4.75

crystal
crystal of element
Type str
Default RHL

cpera
cpera of element
Type float
Default 57.23

conf
conf of element
Type str
Default [Xe]4f14_5d10_6s2_6p3

r_rms
r_rms of element
Type float
Default 0.00010452487744

po
data of element

Keywords

vdw-radius
radius of element

Type float

Default 1.97

covalent
covalent value element

Type float

Default 1.46

Z
z-value of element

Type int

Default 84

mass
mass of element

Type float

Default -209.0

symbol
symbol of element

Type str

Default Po

bpt
bpt of element

Type float

Default 1235.0

mpt
mpt of element

Type float

Default 527.0

density
density of element

Type float

Default 9.4

volume
volume of element

Type float

Default 22.23

name
name of element
Type str
Default Polonium

debye
debye of element
Type float
Default 0.0

a
a of element
Type float
Default 3.35

crystal
crystal of element
Type str
Default SC

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default [Xe]4f14_5d10_6s2_6p4

r_rms
r_rms of element
Type float
Default 0.00010452487744

at
data of element

Keywords

vdw-radius
radius of element
Type float
Default 2.02

covalent
covalent value element
Type float
Default 1.45

Z
z-value of element
Type int
Default 85

mass
mass of element
Type float
Default 210.0

symbol
symbol of element
Type str
Default At

bpt
bpt of element
Type float
Default 610.0

mpt
mpt of element
Type float
Default 575.0

density
density of element
Type float
Default 0.0

volume
volume of element
Type float
Default 0.0

name
name of element
Type str
Default Astatine

debye
debye of element
Type float
Default 0.0

a
a of element
Type float

Default 0.0

crystal

crystal of element

Type str

Default `''''`

cpera

cpera of element

Type float

Default 0.0

conf

conf of element

Type str

Default [Xe]4f14_5d10_6s2_6p5

r_rms

r_rms of element

Type float

Default 0.0001046741666

rn

data of element

Keywords

vdw-radius

radius of element

Type float

Default 2.2

covalent

covalent value element

Type float

Default 1.43

Z

z-value of element

Type int

Default 86

mass

mass of element

Type float

Default -222.0

symbol

symbol of element

Type str

Default Rn
bpt
 bpt of element
Type float
Default 211.0
mpt
 mpt of element
Type float
Default 202.0
density
 density of element
Type float
Default 9.91
volume
 volume of element
Type float
Default 50.5
name
 name of element
Type str
Default Radon
debye
 debye of element
Type float
Default 0.0
a
 a of element
Type float
Default 0.0
crystal
 crystal of element
Type str
Default FCC
cpera
 cpera of element
Type float
Default 0.0
conf
 conf of element

Type str
Default [Xe]4f14_5d10_6s2_6p6

r_rms
r_rms of element

Type float
Default 0.00010642976299

fr
data of element

Keywords

vdw-radius
radius of element

Type float
Default 3.48

covalent
covalent value element

Type float
Default 2.5

Z
z-value of element

Type int
Default 87

mass
mass of element

Type float
Default -223.0

symbol
symbol of element

Type str
Default Fr

bpt
bpt of element

Type float
Default 950.0

mpt
mpt of element

Type float
Default 300.0

density
density of element

Type float

Default 0.0

volume

volume of element

Type float

Default 0.0

name

name of element

Type str

Default Francium

debye

debye of element

Type float

Default 0.0

a

a of element

Type float

Default 0.0

crystal

crystal of element

Type str

Default BCC

cpera

cpera of element

Type float

Default 0.0

conf

conf of element

Type str

Default [Rn]7s1

r_rms

r_rms of element

Type float

Default 0.00010657317899

ra

data of element

Keywords

vdw-radius

radius of element

Type float
Default 2.83

covalent
covalent value element

Type float
Default 2.4

Z
z-value of element

Type int
Default 88

mass
mass of element

Type float
Default 226.025

symbol
symbol of element

Type str
Default Ra

bpt
bpt of element

Type float
Default 1809.0

mpt
mpt of element

Type float
Default 973.0

density
density of element

Type float
Default 5.0

volume
volume of element

Type float
Default 45.2

name
name of element

Type str
Default Radium

debye
debye of element

Type float

Default 0.0

a
a of element

Type float

Default 0.0

crystal
crystal of element

Type str

Default ^``

cpera
cpera of element

Type float

Default 0.0

conf
conf of element

Type str

Default [Rn]7s2

r_rms
r_rms of element

Type float

Default 0.000107000871

ac
data of element

Keywords

vdw-radius
radius of element

Type float

Default -1.0

covalent
covalent value element

Type float

Default 2.2

Z
z-value of element

Type int

Default 89

mass
mass of element
Type float
Default 227.028

symbol
symbol of element
Type str
Default Ac

bpt
bpt of element
Type float
Default 3473.0

mpt
mpt of element
Type float
Default 1323.0

density
density of element
Type float
Default 10.07

volume
volume of element
Type float
Default 22.54

name
name of element
Type str
Default Actinium

debye
debye of element
Type float
Default 0.0

a
a of element
Type float
Default 5.31

crystal
crystal of element
Type str

Default FCC

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default [Rn]6d1_7s2

r_rms
r_rms of element
Type float
Default 0.00010714259349

rf
data of element

Keywords

vdw-radius
radius of element
Type float
Default -1.0

covalent
covalent value element
Type float
Default 0.0

Z
z-value of element
Type int
Default 104

mass
mass of element
Type float
Default -257.0

symbol
symbol of element
Type str
Default Rf

bpt
bpt of element
Type float

Default 0.0

mpt
mpt of element
Type float
Default 0.0

density
density of element
Type float
Default 0.0

volume
volume of element
Type float
Default 0.0

name
name of element
Type str
Default Rutherfordium

debye
debye of element
Type float
Default 0.0

a
a of element
Type float
Default 0.0

crystal
crystal of element
Type str
Default ^``^

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default 4-5s

r_rms
r_rms of element

Type float
Default 0.0001117320442

db
data of element

Keywords

vdw-radius
radius of element

Type float
Default -1.0

covalent
covalent value element

Type float
Default 0.0

z
z-value of element

Type int
Default 105

mass
mass of element

Type float
Default -262.0

symbol
symbol of element

Type str
Default Db

bpt
bpt of element

Type float
Default 0.0

mpt
mpt of element

Type float
Default 0.0

density
density of element

Type float
Default 0.0

volume
volume of element

Type float
Default 0.0

name
name of element

Type str
Default Dubnium

debye
debye of element

Type float
Default 0.0

a
a of element

Type float
Default 0.0

crystal
crystal of element

Type str
Default ````

cpera
cpera of element

Type float
Default 0.0

conf
conf of element

Type str
Default 40s

r_rms
r_rms of element

Type float
Default 0.00011186082063

sg
data of element

Keywords

vdw-radius
radius of element

Type float
Default -1.0

covalent
covalent value element

Type float
Default 0.0

z
z-value of element

Type int
Default 106

mass
mass of element

Type float
Default -263.0

symbol
symbol of element

Type str
Default Sg

bpt
bpt of element

Type float
Default 0.0

mpt
mpt of element

Type float
Default 0.0

density
density of element

Type float
Default 0.0

volume
volume of element

Type float
Default 0.0

name
name of element

Type str
Default Seaborgium

debye
debye of element

Type float
Default 0.0

a
a of element

Type float

Default 0.0

crystal
crystal of element

Type str

Default ''''

cpera
cpera of element

Type float

Default 0.0

conf
conf of element

Type str

Default 0.9s

r_rms
r_rms of element

Type float

Default 0.00011198926979

bh
data of element

Keywords

vdw-radius
radius of element

Type float

Default -1.0

covalent
covalent value element

Type float

Default 0.0

Z
z-value of element

Type int

Default 107

mass
mass of element

Type float

Default -262.0

symbol
symbol of element
Type str
Default Bh

bpt
bpt of element
Type float
Default 0.0

mpt
mpt of element
Type float
Default 0.0

density
density of element
Type float
Default 0.0

volume
volume of element
Type float
Default 0.0

name
name of element
Type str
Default Bohrium

debye
debye of element
Type float
Default 0.0

a
a of element
Type float
Default 0.0

crystal
crystal of element
Type str
Default ^~^

cpera
cpera of element
Type float

Default 0.0

conf

conf of element

Type str

Default 2ms

r_rms

r_rms of element

Type float

Default 0.00011186082063

hs

data of element

Keywords

vdw-radius

radius of element

Type float

Default -1.0

covalent

covalent value element

Type float

Default 0.0

Z

z-value of element

Type int

Default 108

mass

mass of element

Type float

Default -264.0

symbol

symbol of element

Type str

Default Hs

bpt

bpt of element

Type float

Default 0.0

mpt

mpt of element

Type float

Default 0.0

density
density of element
Type float
Default 0.0

volume
volume of element
Type float
Default 0.0

name
name of element
Type str
Default Hassium

debye
debye of element
Type float
Default 0.0

a
a of element
Type float
Default 0.0

crystal
crystal of element
Type str
Default ^``^

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default ^``^

r_rms
r_rms of element
Type float
Default 0.0001122451946

mt
data of element

Keywords

vdw-radius
radius of element

Type float

Default -1.0

covalent
covalent value element

Type float

Default 0.0

Z
z-value of element

Type int

Default 109

mass
mass of element

Type float

Default -266.0

symbol
symbol of element

Type str

Default Mt

bpt
bpt of element

Type float

Default 0.0

mpt
mpt of element

Type float

Default 0.0

density
density of element

Type float

Default 0.0

volume
volume of element

Type float

Default 0.0

name
name of element

Type str
Default Meitnerium

debye
debye of element

Type float
Default 0.0

a
a of element

Type float
Default 0.0

crystal
crystal of element

Type str
Default ````

cpera
cpera of element

Type float
Default 0.0

conf
conf of element

Type str
Default 5ms

r_rms
r_rms of element

Type float
Default 0.00011237267433

110
data of element

Keywords

vdw-radius
radius of element

Type float
Default -1.0

covalent
covalent value element

Type float
Default 0.0

Z
z-value of element

Type int
Default 110

mass
mass of element

Type float
Default -269.0

symbol
symbol of element

Type str
Default 110

bpt
bpt of element

Type float
Default 0.0

mpt
mpt of element

Type float
Default 0.0

density
density of element

Type float
Default 0.0

volume
volume of element

Type float
Default 0.0

name
name of element

Type str
Default (recent_disc.)

debye
debye of element

Type float
Default 0.0

a
a of element

Type float
Default 0.0

crystal
crystal of element

Type str

Default `'''`

cpera
cpera of element

Type float

Default `0.0`

conf
conf of element

Type str

Default `'''`

r_rms
r_rms of element

Type float

Default `-1.0`

111
data of element

Keywords

vdw-radius
radius of element

Type float

Default `-1.0`

covalent
covalent value element

Type float

Default `0.0`

Z
z-value of element

Type int

Default `111`

mass
mass of element

Type float

Default `-272.0`

symbol
symbol of element

Type str

Default `111`

bpt
bpt of element
Type float
Default 0.0

mpt
mpt of element
Type float
Default 0.0

density
density of element
Type float
Default 0.0

volume
volume of element
Type float
Default 0.0

name
name of element
Type str
Default (recent_disc.)

debye
debye of element
Type float
Default 0.0

a
a of element
Type float
Default 0.0

crystal
crystal of element
Type str
Default ***

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str

Default 4/1000s

r_rms
r_rms of element

Type float

Default -1.0

112

data of element

Keywords

vdw-radius
radius of element

Type float

Default -1.0

covalent
covalent value element

Type float

Default 0.0

Z
z-value of element

Type int

Default 112

mass
mass of element

Type float

Default -277.0

symbol
symbol of element

Type str

Default 112

bpt
bpt of element

Type float

Default 0.0

mpt
mpt of element

Type float

Default 0.0

density
density of element

Type float

Default 0.0

volume
volume of element
Type float
Default 0.0

name
name of element
Type str
Default (recent_disc.)

debye
debye of element
Type float
Default 0.0

a
a of element
Type float
Default 0.0

crystal
crystal of element
Type str
Default ^``^

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default 280μs

r_rms
r_rms of element
Type float
Default -1.0

ce
data of element

Keywords

vdw-radius
radius of element
Type float

Default -1.0
covalent
 covalent value element
Type float
Default 1.65

Z
 z-value of element
Type int
Default 58

mass
 mass of element
Type float
Default 140.115

symbol
 symbol of element
Type str
Default Ce

bpt
 bpt of element
Type float
Default 3699.0

mpt
 mpt of element
Type float
Default 1071.0

density
 density of element
Type float
Default 6.78

volume
 volume of element
Type float
Default 20.67

name
 name of element
Type str
Default Cerium

debye
 debye of element

Type float
Default -139.0

a
a of element

Type float
Default 5.16

crystal
crystal of element

Type str
Default FCC

cpera
cpera of element

Type float
Default 0.0

conf
conf of element

Type str
Default [Xe]4f2_5d0_6s2

r_rms
r_rms of element

Type float
Default 9.2803027311e-05

pr
data of element

Keywords

vdw-radius
radius of element

Type float
Default -1.0

covalent
covalent value element

Type float
Default 1.65

Z
z-value of element

Type int
Default 59

mass
mass of element

Type float
Default 140.90765

symbol
symbol of element

Type str
Default Pr

bpt
bpt of element

Type float
Default 3785.0

mpt
mpt of element

Type float
Default 1204.0

density
density of element

Type float
Default 6.77

volume
volume of element

Type float
Default 20.8

name
name of element

Type str
Default Praseodymium

debye
debye of element

Type float
Default -152.0

a
a of element

Type float
Default 3.67

crystal
crystal of element

Type str
Default HEX

cpera
cpera of element
Type float
Default 1.614

conf
conf of element
Type str
Default [Xe]4f3_5d0_6s2

r_rms
r_rms of element
Type float
Default 9.2997877424e-05

nd
data of element

Keywords

vdw-radius
radius of element
Type float
Default -1.0

covalent
covalent value element
Type float
Default 1.64

Z
z-value of element
Type int
Default 60

mass
mass of element
Type float
Default 144.24

symbol
symbol of element
Type str
Default Nd

bpt
bpt of element
Type float
Default 3341.0

mpt
mpt of element
Type float
Default 1289.0

density
density of element
Type float
Default 7.0

volume
volume of element
Type float
Default 20.6

name
name of element
Type str
Default Neodymium

debye
debye of element
Type float
Default -157.0

a
a of element
Type float
Default 3.66

crystal
crystal of element
Type str
Default HEX

cpera
cpera of element
Type float
Default 1.614

conf
conf of element
Type str
Default [Xe]4f4_5d0_6s2

r_rms
r_rms of element
Type float

Default 9.3576955934e-05

pm

data of element

Keywords**vdw-radius**

radius of element

Type float

Default -1.0

covalent

covalent value element

Type float

Default 1.63

Z

z-value of element

Type int

Default 61

mass

mass of element

Type float

Default -145.0

symbol

symbol of element

Type str

Default Pm

bpt

bpt of element

Type float

Default 3785.0

mpt

mpt of element

Type float

Default 1204.0

density

density of element

Type float

Default 6.475

volume

volume of element

Type float

Default 22.39

name
name of element
Type str
Default Promethium

debye
debye of element
Type float
Default 0.0

a
a of element
Type float
Default 0.0

crystal
crystal of element
Type str
Default 

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default [Xe]4f5_5d0_6s2

r_rms
r_rms of element
Type float
Default 9.3768193375e-05

sm
data of element

Keywords

vdw-radius
radius of element
Type float
Default -1.0

covalent
covalent value element
Type float

Default 1.62

Z

z-value of element

Type int

Default 62

mass

mass of element

Type float

Default 150.36

symbol

symbol of element

Type str

Default Sm

bpt

bpt of element

Type float

Default 2064.0

mpt

mpt of element

Type float

Default 1345.0

density

density of element

Type float

Default 7.54

volume

volume of element

Type float

Default 19.95

name

name of element

Type str

Default Samarium

debye

debye of element

Type float

Default 166.0

a

a of element

Type float
Default 9.0

crystal
crystal of element

Type str
Default RHL

cpera
cpera of element

Type float
Default 23.22

conf
conf of element

Type str
Default [Xe]4f6_5d0_6s2

r_rms
r_rms of element

Type float
Default 9.5082839751e-05

eu
data of element

Keywords

vdw-radius
radius of element

Type float
Default -1.0

covalent
covalent value element

Type float
Default 1.85

Z
z-value of element

Type int
Default 63

mass
mass of element

Type float
Default 151.965

symbol
symbol of element

Type str
Default Eu

bpt
bpt of element

Type float
Default 1870.0

mpt
mpt of element

Type float
Default 1090.0

density
density of element

Type float
Default 5.26

volume
volume of element

Type float
Default 28.9

name
name of element

Type str
Default Europium

debye
debye of element

Type float
Default -107.0

a
a of element

Type float
Default 4.61

crystal
crystal of element

Type str
Default BCC

cpera
cpera of element

Type float
Default 0.0

conf

conf of element

Type str

Default [Xe]4f7_5d0_6s2

r_rms

r_rms of element

Type float

Default 9.5267329183e-05

gd

data of element

Keywords

vdw-radius

radius of element

Type float

Default -1.0

covalent

covalent value element

Type float

Default 1.61

Z

z-value of element

Type int

Default 64

mass

mass of element

Type float

Default 157.25

symbol

symbol of element

Type str

Default Gd

bpt

bpt of element

Type float

Default 3539.0

mpt

mpt of element

Type float

Default 1585.0

density
density of element
Type float
Default 7.89

volume
volume of element
Type float
Default 19.9

name
name of element
Type str
Default Gadolinium

debye
debye of element
Type float
Default -176.0

a
a of element
Type float
Default 3.64

crystal
crystal of element
Type str
Default HEX

cpera
cpera of element
Type float
Default 1.588

conf
conf of element
Type str
Default [Xe]4f7_5d1_6s2

r_rms
r_rms of element
Type float
Default 9.6177915369e-05

tb
data of element

Keywords

vdw-radius
radius of element
Type float
Default -1.0

covalent
covalent value element
Type float
Default 1.59

Z
z-value of element
Type int
Default 65

mass
mass of element
Type float
Default 158.92534

symbol
symbol of element
Type str
Default Tb

bpt
bpt of element
Type float
Default 3496.0

mpt
mpt of element
Type float
Default 1630.0

density
density of element
Type float
Default 8.27

volume
volume of element
Type float
Default 19.2

name
name of element
Type str

Default Terbium

debye
debye of element
Type float
Default -188.0

a
a of element
Type float
Default 3.6

crystal
crystal of element
Type str
Default HEX

cpera
cpera of element
Type float
Default 1.581

conf
conf of element
Type str
Default [Xe]4f9_5d0_6s2

r_rms
r_rms of element
Type float
Default 9.6357719009e-05

dy
data of element

Keywords

vdw-radius
radius of element
Type float
Default -1.0

covalent
covalent value element
Type float
Default 1.59

Z
z-value of element
Type int

Default 66

mass

mass of element

Type float

Default 162.5

symbol

symbol of element

Type str

Default Dy

bpt

bpt of element

Type float

Default 2835.0

mpt

mpt of element

Type float

Default 1682.0

density

density of element

Type float

Default 8.54

volume

volume of element

Type float

Default 19.0

name

name of element

Type str

Default Dysprosium

debye

debye of element

Type float

Default -186.0

a

a of element

Type float

Default 3.59

crystal

crystal of element

Type str
Default HEX

cpera
cpera of element

Type float
Default 1.573

conf
conf of element

Type str
Default [Xe]4f10_5d0_6s2

r_rms
r_rms of element

Type float
Default 9.6892647152e-05

ho
data of element

Keywords

vdw-radius
radius of element

Type float
Default -1.0

covalent
covalent value element

Type float
Default 1.58

Z
z-value of element

Type int
Default 67

mass
mass of element

Type float
Default 164.93032

symbol
symbol of element

Type str
Default Ho

bpt
bpt of element

Type float
Default 2968.0

mpt
mpt of element

Type float
Default 1743.0

density
density of element

Type float
Default 8.8

volume
volume of element

Type float
Default 18.7

name
name of element

Type str
Default Holmium

debye
debye of element

Type float
Default -191.0

a
a of element

Type float
Default 3.58

crystal
crystal of element

Type str
Default HEX

cpera
cpera of element

Type float
Default 1.57

conf
conf of element

Type str
Default [Xe]4f11_5d0_6s2

r_rms
r_rms of element
Type float
Default 9.6892647152e-05

er
data of element

Keywords

vdw-radius
radius of element

Type float
Default -1.0

covalent
covalent value element

Type float
Default 1.57

Z
z-value of element

Type int
Default 68

mass
mass of element

Type float
Default 167.26

symbol
symbol of element

Type str
Default Er

bpt
bpt of element

Type float
Default 3136.0

mpt
mpt of element

Type float
Default 1795.0

density
density of element

Type float
Default 9.05

volume
volume of element
Type float
Default 18.4

name
name of element
Type str
Default Erbium

debye
debye of element
Type float
Default -195.0

a
a of element
Type float
Default 3.56

crystal
crystal of element
Type str
Default HEX

cpera
cpera of element
Type float
Default 1.57

conf
conf of element
Type str
Default [Xe]4f12_5d0_6s2

r_rms
r_rms of element
Type float
Default 9.7943009317e-05

tm
data of element

Keywords

vdw-radius
radius of element
Type float
Default -1.0

covalent
covalent value element
Type float
Default 1.56

Z
z-value of element
Type int
Default 69

mass
mass of element
Type float
Default 168.93421

symbol
symbol of element
Type str
Default Tm

bpt
bpt of element
Type float
Default 2220.0

mpt
mpt of element
Type float
Default 1818.0

density
density of element
Type float
Default 9.33

volume
volume of element
Type float
Default 18.1

name
name of element
Type str
Default Thulium

debye
debye of element
Type float

Default -200.0

a

a of element

Type float

Default 3.54

crystal

crystal of element

Type str

Default HEX

cpera

cpera of element

Type float

Default 1.57

conf

conf of element

Type str

Default [Xe]4f13_5d0_6s2

r_rms

r_rms of element

Type float

Default 9.811562674e-05

yb

data of element

Keywords

vdw-radius

radius of element

Type float

Default -1.0

covalent

covalent value element

Type float

Default 1.74

Z

z-value of element

Type int

Default 70

mass

mass of element

Type float

Default 173.04
symbol
symbol of element
Type str
Default Yb
bpt
bpt of element
Type float
Default 1467.0
mpt
mpt of element
Type float
Default 1097.0
density
density of element
Type float
Default 6.98
volume
volume of element
Type float
Default 24.79
name
name of element
Type str
Default Ytterbium
debye
debye of element
Type float
Default -118.0
a
a of element
Type float
Default 5.49
crystal
crystal of element
Type str
Default FCC
cpera
cpera of element

Type float

Default 0.0

conf

conf of element

Type str

Default [Xe]4f14_5d0_6s2

r_rms

r_rms of element

Type float

Default 9.8968651305e-05

lu

data of element

Keywords

vdw-radius

radius of element

Type float

Default -1.0

covalent

covalent value element

Type float

Default 1.56

Z

z-value of element

Type int

Default 71

mass

mass of element

Type float

Default 174.967

symbol

symbol of element

Type str

Default Lu

bpt

bpt of element

Type float

Default 3668.0

mpt

mpt of element

Type float
Default 1936.0

density
density of element

Type float
Default 9.84

volume
volume of element

Type float
Default 17.78

name
name of element

Type str
Default Lutetium

debye
debye of element

Type float
Default -207.0

a
a of element

Type float
Default 3.51

crystal
crystal of element

Type str
Default HEX

cpera
cpera of element

Type float
Default 1.585

conf
conf of element

Type str
Default [Xe]4f14_5d1_6s2

r_rms
r_rms of element

Type float
Default 9.9137288835e-05

th

data of element

Keywords**vdw-radius**

radius of element

Type float**Default -1.0****covalent**

covalent value element

Type float**Default 1.65****Z**

z-value of element

Type int**Default 90****mass**

mass of element

Type float**Default 232.0381****symbol**

symbol of element

Type str**Default Th****bpt**

bpt of element

Type float**Default 5061.0****mpt**

mpt of element

Type float**Default 2028.0****density**

density of element

Type float**Default 11.7****volume**

volume of element

Type float**Default 19.9**

name
name of element
Type str
Default Thorium

debye
debye of element
Type float
Default 100.0

a
a of element
Type float
Default 5.08

crystal
crystal of element
Type str
Default FCC

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default [Rn]6d2_7s2

r_rms
r_rms of element
Type float
Default 0.00010784503195

pa
data of element

Keywords

vdw-radius
radius of element
Type float
Default -1.0

covalent
covalent value element
Type float
Default 0.0

Z
z-value of element
Type int
Default 91

mass
mass of element
Type float
Default 231.03588

symbol
symbol of element
Type str
Default Pa

bpt
bpt of element
Type float
Default 0.0

mpt
mpt of element
Type float
Default 0.0

density
density of element
Type float
Default 15.4

volume
volume of element
Type float
Default 15.0

name
name of element
Type str
Default Protactinium

debye
debye of element
Type float
Default 0.0

a
a of element
Type float

Default 3.92

crystal

crystal of element

Type str

Default TET

cpera

cpera of element

Type float

Default 0.825

conf

conf of element

Type str

Default [Rn]5f2_6d1_7s2

r_rms

r_rms of element

Type float

Default 0.00010770535752

u

data of element

Keywords

vdw-radius

radius of element

Type float

Default -1.0

covalent

covalent value element

Type float

Default 1.42

Z

z-value of element

Type int

Default 92

mass

mass of element

Type float

Default 238.0289

symbol

symbol of element

Type str

Default U

bpt
bpt of element
Type float
Default 4407.0

mpt
mpt of element
Type float
Default 1405.0

density
density of element
Type float
Default 18.9

volume
volume of element
Type float
Default 12.59

name
name of element
Type str
Default Uranium

debye
debye of element
Type float
Default -210.0

a
a of element
Type float
Default 2.85

crystal
crystal of element
Type str
Default ORC

cpera
cpera of element
Type float
Default 0.0

conf
conf of element

Type str
Default [Rn]5f3_6d1_7s2

r_rms
r_rms of element
Type float
Default 0.00010867476102

np
data of element

Keywords

vdw-radius
radius of element
Type float
Default -1.0

covalent
covalent value element
Type float
Default 0.0

Z
z-value of element
Type int
Default 93

mass
mass of element
Type float
Default 237.048

symbol
symbol of element
Type str
Default Np

bpt
bpt of element
Type float
Default 0.0

mpt
mpt of element
Type float
Default 910.0

density
density of element

Type float
Default 20.4

volume
volume of element

Type float
Default 11.62

name
name of element

Type str
Default Neptunium

debye
debye of element

Type float
Default -188.0

a
a of element

Type float
Default 4.72

crystal
crystal of element

Type str
Default ORC

cpera
cpera of element

Type float
Default 0.0

conf
conf of element

Type str
Default [Rn]5f4_6d1_7s2

r_rms
r_rms of element

Type float
Default 0.00010853744903

pu
data of element

Keywords

vdw-radius
radius of element

Type float
Default -1.0

covalent
covalent value element

Type float
Default 0.0

Z
z-value of element

Type int
Default 94

mass
mass of element

Type float
Default -244.0

symbol
symbol of element

Type str
Default Pu

bpt
bpt of element

Type float
Default 3503.0

mpt
mpt of element

Type float
Default 913.0

density
density of element

Type float
Default 19.8

volume
volume of element

Type float
Default 12.32

name
name of element

Type str
Default Plutonium

debye
debye of element
Type float
Default -150.0

a
a of element
Type float
Default 0.0

crystal
crystal of element
Type str
Default MCL

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default [Rn]5f6_6d0_7s2

r_rms
r_rms of element
Type float
Default 0.00010949065967

am
data of element
Keywords

vdw-radius
radius of element
Type float
Default -1.0

covalent
covalent value element
Type float
Default 0.0

Z
z-value of element
Type int
Default 95

mass
mass of element
Type float
Default -243.0

symbol
symbol of element
Type str
Default Am

bpt
bpt of element
Type float
Default 2880.0

mpt
mpt of element
Type float
Default 1268.0

density
density of element
Type float
Default 13.6

volume
volume of element
Type float
Default 17.86

name
name of element
Type str
Default Americium

debye
debye of element
Type float
Default 0.0

a
a of element
Type float
Default 0.0

crystal
crystal of element
Type str

Default 

cpera
cpera of element

Type float

Default 0.0

conf
conf of element

Type str

Default [Rn] 5f7_6d0_7s2

r_rms
r_rms of element

Type float

Default 0.00010935561268

cm
data of element

Keywords

vdw-radius
radius of element

Type float

Default -1.0

covalent
covalent value element

Type float

Default 0.0

Z
z-value of element

Type int

Default 96

mass
mass of element

Type float

Default -247.0

symbol
symbol of element

Type str

Default Cm

bpt
bpt of element

Type float

Default 0.0

mpt
mpt of element
Type float
Default 1340.0

density
density of element
Type float
Default 13.511

volume
volume of element
Type float
Default 18.28

name
name of element
Type str
Default Curium

debye
debye of element
Type float
Default 0.0

a
a of element
Type float
Default 0.0

crystal
crystal of element
Type str
Default ^``^

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default [Rn]5f7_6d1_7s2

r_rms
r_rms of element

Type float

Default 0.00010989359973

bk

data of element

Keywords

vdw-radius

radius of element

Type float

Default -1.0

covalent

covalent value element

Type float

Default 0.0

Z

z-value of element

Type int

Default 97

mass

mass of element

Type float

Default -247.0

symbol

symbol of element

Type str

Default Bk

bpt

bpt of element

Type float

Default 0.0

mpt

mpt of element

Type float

Default 0.0

density

density of element

Type float

Default 0.0

volume

volume of element

Type float
Default 0.0

name
name of element

Type str
Default Berkelium

debye
debye of element

Type float
Default 0.0

a
a of element

Type float
Default 0.0

crystal
crystal of element

Type str
Default ````

cpera
cpera of element

Type float
Default 0.0

conf
conf of element

Type str
Default [Rn]5f8_6d1_7s2

r_rms
r_rms of element

Type float
Default 0.00010989359973

cf
data of element

Keywords

vdw-radius
radius of element

Type float
Default -1.0

covalent
covalent value element

Type float
Default 0.0

z
z-value of element

Type int
Default 98

mass
mass of element

Type float
Default -251.0

symbol
symbol of element

Type str
Default Cf

bpt
bpt of element

Type float
Default 0.0

mpt
mpt of element

Type float
Default 900.0

density
density of element

Type float
Default 0.0

volume
volume of element

Type float
Default 0.0

name
name of element

Type str
Default Californium

debye
debye of element

Type float
Default 0.0

a
a of element
Type float
Default 0.0

crystal
crystal of element
Type str
Default ''''

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str
Default [Rn]5f10_6d0_7s2

r_rms
r_rms of element
Type float
Default 0.00011042580946

es
data of element

Keywords

vdw-radius
radius of element
Type float
Default -1.0

covalent
covalent value element
Type float
Default 0.0

Z
z-value of element
Type int
Default 99

mass
mass of element
Type float
Default -252.0

symbol
symbol of element

Type str

Default Es

bpt
bpt of element

Type float

Default 0.0

mpt
mpt of element

Type float

Default 0.0

density
density of element

Type float

Default 0.0

volume
volume of element

Type float

Default 0.0

name
name of element

Type str

Default Einsteinium

debye
debye of element

Type float

Default 0.0

a
a of element

Type float

Default 0.0

crystal
crystal of element

Type str

Default ^``^

cpera
cpera of element

Type float

Default 0.0

conf

conf of element

Type str

Default [Rn]5f11_6d0_7s2

r_rms

r_rms of element

Type float

Default 0.00011055797721

fm

data of element

Keywords

vdw-radius

radius of element

Type float

Default -1.0

covalent

covalent value element

Type float

Default 0.0

Z

z-value of element

Type int

Default 100

mass

mass of element

Type float

Default -257.0

symbol

symbol of element

Type str

Default Fm

bpt

bpt of element

Type float

Default 0.0

mpt

mpt of element

Type float

Default 0.0

density

density of element

Type float

Default 0.0

volume

volume of element

Type float

Default 0.0

name

name of element

Type str

Default Fermium

debye

debye of element

Type float

Default 0.0

a

a of element

Type float

Default 0.0

crystal

crystal of element

Type str

Default ^``^

cpera

cpera of element

Type float

Default 0.0

conf

conf of element

Type str

Default [Rn]5f12_6d0_7s2

r_rms

r_rms of element

Type float

Default 0.00011121362374

md

data of element

Keywords

vdw-radius
radius of element

Type float

Default -1.0

covalent
covalent value element

Type float

Default 0.0

Z
z-value of element

Type int

Default 101

mass
mass of element

Type float

Default -258.0

symbol
symbol of element

Type str

Default Md

bpt
bpt of element

Type float

Default 0.0

mpt
mpt of element

Type float

Default 0.0

density
density of element

Type float

Default 0.0

volume
volume of element

Type float

Default 0.0

name
name of element

Type str
Default Mendelevium

debye
debye of element

Type float
Default 0.0

a
a of element

Type float
Default 0.0

crystal
crystal of element

Type str
Default ````

cpera
cpera of element

Type float
Default 0.0

conf
conf of element

Type str
Default [Rn]5f13_6d0_7s2

r_rms
r_rms of element

Type float
Default 0.00011134373034

“no”
data of element

Keywords

vdw-radius
radius of element

Type float
Default -1.0

covalent
covalent value element

Type float
Default 0.0

Z
z-value of element

Type int
Default 102

mass
mass of element

Type float
Default -259.0

symbol
symbol of element

Type str
Default "No"

bpt
bpt of element

Type float
Default 0.0

mpt
mpt of element

Type float
Default 0.0

density
density of element

Type float
Default 0.0

volume
volume of element

Type float
Default 0.0

name
name of element

Type str
Default Nobelium

debye
debye of element

Type float
Default 0.0

a
a of element

Type float
Default 0.0

crystal
crystal of element

Type str

Default `---`

cpera
cpera of element

Type float

Default `0.0`

conf
conf of element

Type str

Default `[Rn]5f14_6d0_7s2`

r_rms
r_rms of element

Type float

Default `0.00011147350119`

lr
data of element

Keywords

vdw-radius
radius of element

Type float

Default `-1.0`

covalent
covalent value element

Type float

Default `0.0`

z
z-value of element

Type int

Default `103`

mass
mass of element

Type float

Default `-260.0`

symbol
symbol of element

Type str

Default `Lr`

bpt
bpt of element
Type float
Default 0.0

mpt
mpt of element
Type float
Default 0.0

density
density of element
Type float
Default 0.0

volume
volume of element
Type float
Default 0.0

name
name of element
Type str
Default Lawrencium

debye
debye of element
Type float
Default 0.0

a
a of element
Type float
Default 0.0

crystal
crystal of element
Type str
Default ***

cpera
cpera of element
Type float
Default 0.0

conf
conf of element
Type str

Default [Rn] 5f14_6d1_7s2
r_rms
r_rms of element
Type float
Default 0.00011186082063

2.2.4 Running MRChem with QCEngine

MRChem >=1.0 can be used as a computational engine with the [QCEngine](#) program executor. QCEngine can be useful for running calculations on large sets of molecules and input parameters. The results are collected in standardised [QCSchema format](#), which makes it easy to build post-processing pipelines and store data according to Findability, Accessibility, Interoperability, and Reuse (FAIR) of digital assets principles. Furthermore, QCEngine provides different geometry optimization drivers that can use the molecular gradient computed by MRChem for structural optimization.

Installation

The easiest way is to install both QCEngine and MRChem in a Conda environment using the precompiled version:

```
conda create -n mrchem-qcng mrchem qcengine qcelemental geometric optking pip -c conda-forge
conda activate mrchem-qcng
python -m pip install -U pyberny
```

It is also possible to use your own installation of MRChem: just make sure that the installation folder is in your PATH.

Note: If you want to use the precompiled, MPI-parallel version of MRChem with OpenMPI, install `mrchem=*-*openmpi*` instead of just `mrchem`. A binary package compiled against MPICH is also available: `mrchem=*-*mpich*`.

Single compute

Calculations in QCEngine are defined in Python scripts. For example, the following runs MRChem to obtain the energy of water:

```
import qcelemental as qcel
import qcengine as qcng

mol = qcel.models.Molecule(geometry=[[0, 0, 0], [0, 1.5, 0], [0, 0, 1.5]],
                             symbols=["O", "H", "H"],
                             connectivity=[[0, 1, 1], [0, 2, 1]])
print(mol)

computation = {
    "molecule": mol,
    "driver": "energy",
    "model": {"method": "HF"},
    "keywords": {"world_prec": 1.0e-3},
```

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```
}
```

```
ret = qcng.compute(computation, "mrchem")
```

```
print(f"E_HF = {ret.return_result} Hartree")
```

You can save this sample as *mrchem-run-hf.py* and execute it with:

```
python mrchem-run-hf.py
```

Which will print to screen:

```
Molecule(name='H2O', formula='H2O', hash='b41d0c5')
```

```
E_HF = -75.9789291596064 Hartree
```

Note that:

1. The molecule is specified, in Angstrom, using a `QCElemental` object.
2. The computation is described using a Python dictionary.
3. The `driver` selects the kind of calculation you want to run with MRChem. Available drivers are: - `energy`, for single-point energy calculations. - `gradient`, for evaluation of the molecular gradient at a given geometry. - `properties`, for the calculation of molecular properties.
4. The `model` selects the wavefunction: HF for Hartree-Fock and any of the DFT functionals known to MRChem for a corresponding DFT calculation.
5. The `keywords` key in the dictionary accepts a dictionary of MRChem options. Any of the options in the usual input file are recognized.

Once you have a dictionary defining your computation, you can run it with:

```
ret = qcng.compute(computation, "mrchem")
```

You can reuse the same dictionary with *multiple* computational engine, e.g. other quantum chemistry programs that are recognized as executors by QCEngine. The return value from the `compute` function contains all data produced during the calculation in QCSchema format including, for example, the execution time elapsed. The full JSON output produced by MRChem is also available and can be inspected in Python as:

```
mrchem_json_out = ret.extras["raw_output"]["output"]
```

The full, human-readable input is saved as the `stdout` property of the object returned by `compute`.

Parallelism

QCEngine allows you to exploit available parallel hardware. For example, to use 20 OpenMP threads in your MRChem calculation you would provide an additional task configuration dictionary as a `task_config` argument to `compute`:

```
ret = qcng.compute(
    computation,
    "mrchem",
    task_config={"ncores": 20})
```

You can inspect how the job was launched by printing out the provenance dictionary:

```
print(ret.extras["raw_output"]["output"]["provenance"])
```

```
{
  "creator": "MRChem",
  "mpi_processes": 1,
  "routine": "/home/roberto/miniconda3/envs/mrchem-qcng/bin/mrchem.x",
  "total_cores": 1,
  "version": "1.1.0",
  "ncores": 12,
  "nnodes": 1,
  "ranks_per_node": 1,
  "cores_per_rank": 12,
  "total_ranks": 1
}
```

It is also possible to run MPI-parallel and hybrid MPI+OpenMP jobs. Assuming that you installed the MPICH version of the MRChem MPI-parallel Conda package, the basic `task_config` argument to `compute` would look like:

```
task = {
  "nnodes": 1, # number of nodes
  "ncores": 12, # number of cores per task on each node
  "cores_per_rank": 6, # number of cores per MPI rank
  "use_mpiexec": True, # launch with MPI
  "mpiexec_command": "mpiexec -n {total_ranks}", # the invocation of MPI
}
```

This task configuration will launch a MPI job with 2 ranks on a single node. Each rank has access to 6 cores for OpenMP parallelization. The provenance dictionary now shows:

```
{
  "creator": "MRChem",
  "mpi_processes": 2,
  "routine": "mpiexec -n 2 /home/roberto/miniconda3/envs/mrchem-qcng/bin/mrchem.x",
  "total_cores": 12,
  "version": "1.1.0",
  "ncores": 12,
  "nnodes": 1,
  "ranks_per_node": 2,
  "cores_per_rank": 6,
  "total_ranks": 2
}
```

The `mpiexec_command` is a string that will be interpolated to provide the exact invocation. In the above example, MRChem will be run with:

```
mpiexec -n 2 /home/roberto/miniconda3/envs/mrchem-qcng/bin/mrchem.x
```

The following interpolation parameters are understood by QC Engine when creating the MPI invocation:

- `{nnodes}`: number of nodes.
- `{cores_per_rank}`: number of cores to use for each MPI rank.
- `{ranks_per_node}`: number of MPI ranks per node. Computed as `ncores // cores_per_rank`.
- `{total_ranks}`: total number of MPI ranks. Computed as `nnodes * ranks_per_node`.

More complex MPI invocations are possible by setting the appropriate `mpiexec_command` in the task configuration. For usage with a scheduler, such as SLURM, you should refer to the documentation of your computing cluster and the documentation of QC Engine.

Geometry optimizations

Running geometry optimizations is just as easy as single compute. The following example optimizes the structure of water using the SVWN5 functional with MW4. The `geometric` package is used as optimization driver, but `pyberny` or `optking` would also work.

Warning: The computation of the molecular gradient can be affected by significant numerical noise for MW3 and MW4, to the point that it can be impossible to converge a geometry optimization. Using a tighter precision might help, but the cost of the calculation might be prohibitively large.

```
import qcelemental as qcel
import qcengine as qcng

mol = qcel.models.Molecule(
    geometry=[
        [ 0.29127930, 3.00875625, 0.20308515],
        [-1.21253048, 1.95820900, 0.10303324],
        [ 0.10002049, 4.24958115,-1.10222079]
    ],
    symbols=["O", "H", "H"],
    fix_com=True,
    fix_orientation=True,
    fix_symmetry="c1")

opt_input = {
    "keywords": {
        "program": "mrchem",
        "maxiter": 70
    },
    "input_specification": {
        "driver": "gradient",
        "model": {
            "method": "SVWN5",
        },
        "keywords": {
            "world_prec": 1.0e-4,
            "SCF": {
                "guess_type": "core_dz",
            }
        }
    },
    "initial_molecule": mol,
}

opt = qcng.compute_procedure(
    opt_input,
    "geometric",
```

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```

task_config={"ncores": 20}

print(opt.stdout)

print("==> Optimized geometry <==")
print(opt.final_molecule.pretty_print())

print("==> Optimized geometric parameters <==")
for m in [[0, 1], [0, 2], [1, 0, 2]]:
    opt_val = opt.final_molecule.measure(m)
    print(f"Internal degree of freedom [{m}] = {opt_val:.3f}")

```

Running this script will print all the steps taken during the structural optimization. The final printout contains the optimized geometry:

```
Geometry (in Angstrom), charge = 0.0, multiplicity = 1:
```

Center	X	Y	Z
O	-4.146209038013	2.134923126314	-3.559202294678
H	-4.906566693905	1.536801624016	-3.587431156799
H	-4.270830051398	2.773072094238	-4.275607223691

and the optimized values of bond distances and bond angle:

```

Internal degree of freedom [0, 1] = 1.829
Internal degree of freedom [0, 2] = 1.828
Internal degree of freedom [1, 0, 2] = 106.549

```

2.2.5 Program input/output file

Input schema

```

"input": {
  "schema_name": string,                                # Name of the input schema
  "schema_version": int,                               # Version of the input schema
  "molecule": {                                         # Section for Molecule specification
    "charge": int,                                     # Total molecular charge
    "multiplicity": int,                             # Total spin multiplicity
    "coords": array[                                     # Array of atoms
      {
        "atom": string,                                # Atomic symbol
        "xyz": array[float]                            # Nuclear Cartesian coordinate
      }
    ],
    "cavity": {                                         # Array of cavity spheres
      "spheres": array[                                 # (one entry per sphere)
        {
          "center": array[float],                      # Cartesian coordinate of sphere center
          "radius": float,                            # Radius of cavity sphere
          "alpha": float,                           # Scaling factor of radius
        }
      ],
    }
  }
}

```

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```

        "beta": float
        "sigma": float
    }
],
}
},
"mpi": {
    "bank_size": int,
    "omp_threads": int,
    "numerically_exact": bool,
    "shared_memory_size": int
},
"mra": {
    "basis_type": string,
    "basis_order": int,
    "max_scale": int,
    "min_scale": int,
    "boxes": array[int],
    "corner": array[int]
},
"printer": {
    "file_name": string,
    "print_level": int,
    "print_mpi": bool,
    "print_prec": int,
    "print_width": int
},
"scf_calculation": {
    "fock_operator": {
        "kinetic_operator": {
            "derivative": string
        },
        "nuclear_operator": {
            "proj_prec": float,
            "smooth_prec": float,
            "shared_memory": bool
        },
        "coulomb_operator": {
            "poisson_prec": float,
            "shared_memory": bool
        },
        "exchange_operator": {
            "poisson_prec": float,
            "screen": bool
        },
        "reaction_operator": {
            "poisson_prec": float,
            "kain": int,
        }
    },
    "procedure": {
        "max_iter": int,
    },
    "SCRF procedure": {
        "optimizer": string,
    }
},
}
}

# Scaling factor of width
# Width of cavity boundary

# Section for MPI specification
# Number of MPI ranks in memory bank
# Number of omp threads
# Guarantee MPI invariant results
# Size (MB) of MPI shared memory blocks

# Section for MultiResolution Analysis
# Basis type (interpolating/legendre)
# Polynomial order of basis
# Maximum level of refinement
# Minimum level of refinement (root scale)
# Number of root boxes
# Translation of first root box

# Section for printed output
# Name of output file
# Amount of printed output
# Use separate output file for each MPI
# Number of digits for printed output
# Line width of printed output

# Section for SCF specification
# Contributions to Fock operator
# Add Kinetic operator to Fock
# Type of derivative operator

# Add Nuclear operator to Fock
# Projection prec for potential
# Smoothing parameter for potential
# Use shared memory for potential

# Add Coulomb operator to Fock
# Build prec for Poisson operator
# Use shared memory for potential

# Add Exchange operator to Fock
# Build prec for Poisson operator
# Use screening in Exchange operator

# Add Reaction operator to Fock
# Precision for Poisson operator
# Length of KAIN history in nested SCRF

# Maximum number of iterations in nested
# Use density or potential in KAIN solver

```

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```

    "dynamic_thrs": bool,
    "threshold"
        "density_type": string,
    "electronic"
        "epsilon_in": float,
        "epsilon_out": float,
        "formulation": string
    },
    "xc_operator": {
        "shared_memory": bool,
        "xc_functional": {
            "spin": bool,
            "cutoff": float,
            "functionals": array[
                {
                    "coef": float,
                    "name": string
                }
            ]
        }
    },
    "zora_operator": {
        "include_nuclear": bool,
        "include_coulomb": bool,
        "include_xc": bool
    },
    "external_operator": {
        "electric_field": array[float],
        "r_0": array[float]
    }
},
"initial_guess": {
    "type": string,
    "prec": float,
    "zeta": int,
    "method": string,
    "localize": bool,
    "restricted": bool,
    "relativity": string,
    "screen": float,
    "file_chk": string,
    "file_basis": string,
    "file_gto_a": string,
    "file_gto_b": string,
    "file_gto_p": string,
    "file_phi_a": string,
    "file_phi_b": string,
    "file_phi_p": string,
    "file_CUBE_a": str,
    "file_CUBE_b": str,
    "file_CUBE_p": str
},

```

Use static or dynamic convergence
Type of charge density [total, nuclear,
Permittivity inside the cavity
Permittivity outside the cavity
Formulation of the permittivity function
Add XC operator to Fock
Use shared memory for potential
XC functional specification
Use spin separated functional
Cutoff value for small densities
Array of density functionals
Numerical coefficient
Functional name
Include V_nuc in ZORA potential
Include V_coul in ZORA potential
Include V_xc in ZORA potential
Add external field operator to Fock
Electric field vector
Gauge origin for electric field
Initial guess specification
Type of initial guess
Precision for initial guess
Zeta quality for AO basis
Name of method for initial energy
Use localized orbitals
Use spin restricted orbitals
Name of relativistic method
Screening used in GTO evaluations
Path to checkpoint file
Path to GTO basis file
Path to GTO MO file (alpha)
Path to GTO MO file (beta)
Path to GTO MO file (paired)
Path to MW orbital file (alpha)
Path to MW orbital file (beta)
Path to MW orbital file (paired)
Path to CUBE orbital file (alpha)
Path to CUBE orbital file (beta)
Path to CUBE orbital file (paired)

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```

"scf_solver": {                                     # SCF solver specification
    "kain": int,                                 # Length of KAIN history
    "max_iter": int,                            # Maximum number of iterations
    "method": string,                           # Name of electronic structure method
    "relativity": string,                      # Name of relativistic method
    "rotation": int,                            # Iterations between localize/diagonalize
    "localize": bool,                           # Use localized orbitals
    "checkpoint": bool,                         # Save checkpoint file
    "file_chk": string,                        # Name of checkpoint file
    "start_prec": float,                       # Start precision for solver
    "final_prec": float,                        # Final precision for solver
    "helmholtz_prec": float,                   # Precision for Helmholtz operators
    "orbital_thrs": float,                     # Convergence threshold orbitals
    "energy_thrs":float                         # Convergence threshold energy
},
"properties": {                                    # Collection of properties to compute
    "dipole_moment": {                          # Collection of dipole moments
        "id (string)": {                        # Unique id: 'dip-${number}'
            "precision": float,                # Operator precision
            "operator": string,              # Operator used for property
            "r_0": array[float]             # Operator gauge origin
        }
    },
    "quadrupole_moment": {                      # Collection of quadrupole moments
        "id (string)": {                        # Unique id: 'quad-${number}'
            "precision": float,                # Operator precision
            "operator": string,              # Operator used for property
            "r_0": array[float]             # Operator gauge origin
        }
    },
    "geometric_derivative": {                  # Collection of geometric derivatives
        "id (string)": {                        # Unique id: 'geom-${number}'
            "precision": float,                # Operator precision
            "operator": string,              # Operator used for property
            "smooth_prec": float           # Smoothing parameter for potential
        }
    }
},
"plots": {                                       # Collection of plots to perform
    "density": bool,                           # Plot converged densities
    "orbitals": array[int],                    # List of orbitals to plot
    "plotter": {                                # Section specifying plotting parameters
        "path": string,                         # Path to output files
        "type": string,                         # Type of plot (line, surf or cube)
        "points": array[int],                  # Number of points in each direction
        "O": array[float],                      # Plotting range origin
        "A": array[float],                      # Plotting range A vector
        "B": array[float],                      # Plotting range B vector
        "C": array[float]                       # Plotting range C vector
    }
},
"rsp_calculations": {                           # Collection of response calculations
}

```

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```

id (string): {                                     # Response id: e.g. 'ext_el-${frequency}'
  "dynamic": bool,                                # Use dynamic response solver
  "frequency": float,                             # Perturbation frequency
  "perturbation": {                               # Perturbation operator
    "operator": string                            # Operator used in response calculation
  },
  "components": array[                           # Array of perturbation components
    {
      "initial_guess": {                         # (one per Cartesian direction)
        "type": string,                          # Initial guess specification
        "prec": float,                           # Type of initial guess
        "file_chk_x": string,                   # Precision for initial guess
        "file_chk_y": string,                   # Path to checkpoint file for X
        "file_x_a": string,                     # Path to checkpoint file for Y
        "file_x_b": string,                     # Path to MW file for X (alpha)
        "file_x_p": string,                     # Path to MW file for X (beta)
        "file_y_a": string,                     # Path to MW file for X (paired)
        "file_y_b": string,                     # Path to MW file for Y (alpha)
        "file_y_p": string,                     # Path to MW file for Y (beta)
        "file_y_p": string                      # Path to MW file for Y (paired)
      },
      "rsp_solver": {                           # Response solver specification
        "kain": int,                            # Length of KAIN history
        "max_iter": int,                        # Maximum number of iterations
        "method": string,                       # Name of electronic structure method
        "checkpoint": bool,                     # Save checkpoint file
        "file_chk_x": string,                  # Name of X checkpoint file
        "file_chk_y": string,                  # Name of Y checkpoint file
        "orth_prec": float,                    # Precision for orthogonalization
        "start_prec": float,                   # Start precision for solver
        "final_prec": float,                   # Final precision for solver
        "helmholtz_prec": float,              # Precision for Helmholtz operators
        "orbital_thrs": float,                 # Convergence threshold orbitals
        "property_thrs": float                # Convergence threshold property
      }
    }
  ],
  "properties": {                                 # Collection of properties to compute
    "polarizability": {                         # Collection of polarizabilities
      id (string): {                           # Unique id: 'pol-${frequency}'
        "precision": float,                   # Operator precision
        "operator": string,                  # Operator used for property
        "r_0": array[float]                  # Operator gauge origin
      }
    },
    "magnetizability": {                        # Collection of magnetizabilities
      id (string): {                           # Unique id: 'mag-${frequency}'
        "frequency": float,                  # Perturbation frequency
        "precision": float,                  # Operator precision
        "dia_operator": string,             # Operator used for diamagnetic property
        "para_operator": string,            # Operator used for paramagnetic property
        "derivative": string,               # Operator derivative type
        "r_0": array[float]                  # Operator gauge origin
      }
    }
  }
]

```

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```

        }
    },
    "nmr_shielding": {
        "id": (string): {
            "precision": float,
            "dia_operator": string,
            "para_operator": string,
            "derivative": string,
            "smoothing": float,
            "r_0": array[float],
            "r_K": array[float]
        }
    },
    "fock_operator": {
        "coulomb_operator": {
            "poisson_prec": float,
            "shared_memory": bool
        },
        "exchange_operator": {
            "poisson_prec": float,
            "screen": bool
        },
        "xc_operator": {
            "shared_memory": bool,
            "xc_functional": {
                "spin": bool,
                "cutoff": float,
                "functionals": array[
                    {
                        "coef": float,
                        "name": string
                    }
                ]
            }
        }
    },
    "unperturbed": {
        "prec": float,
        "localize": bool,
        "fock_operator": {
            "kinetic_operator": {
                "derivative": string
            },
            "nuclear_operator": {
                "proj_prec": float,
                "smooth_prec": float,
                "shared_memory": bool
            },
            "coulomb_operator": {
                "poisson_prec": float,
                "shared_memory": bool
            }
        }
    }
},
# Collection of NMR shieldings
# Unique id: 'nmr-${nuc_idx}${atom_symbol}'
# Operator precision
# Operator used for diamagnetic property
# Operator used for paramagnetic property
# Operator derivative type
# Operator smoothing parameter
# Operator gauge origin
# Nuclear coordinate

# Contributions to perturbed Fock operator
# Add Coulomb operator to Fock
# Build prec for Poisson operator
# Use shared memory for potential

# Add Exchange operator to Fock
# Build prec for Poisson operator
# Use screening in Exchange operator

# Add XC operator to Fock
# Use shared memory for potential
# XC functional specification
# Use spin separated functional
# Cutoff value for small densities
# Array of density functionals

# Numerical coefficient
# Functional name

# Section for unperturbed part of response
# Precision used for unperturbed system
# Use localized unperturbed orbitals
# Contributions to unperturbed Fock operator
# Add Kinetic operator to Fock
# Type of derivative operator

# Add Nuclear operator to Fock
# Projection prec for potential
# Smoothing parameter for potential
# Use shared memory for potential

# Add Coulomb operator to Fock
# Build prec for Poisson operator
# Use shared memory for potential

```

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```

    },
    "exchange_operator": {
        "poisson_prec": float,
        "screen": bool
    },
    "xc_operator": {
        "shared_memory": bool,
        "xc_functional": {
            "spin": bool,
            "cutoff": float,
            "functionals": array[
                {
                    "coef": float,
                    "name": string
                }
            ]
        }
    },
    "external_operator": {
        "electric_field": array[float],
        "r_0": array[float]
    }
},
"constants": {
    "angstrom2bohrs": float,
    "dipmom_au2debye": float,
    Debye
        "electron_g_factor": float,
        "fine_structure_constant": float,
        "hartree2ev": float,
        "hartree2kcalmol": float,
        "hartree2kjmol": float,
        "hartree2simagnetizability": float,
        "hartree2wavenumbers": float,
        "light_speed": float
    }
}

```

Physical constants used throughout MRChem
Conversion factor from Angstrom to Bohr
Conversion factor from atomic units to debye
Electron g factor in atomic units
Fine-structure constant in atomic units
Conversion factor from Hartree to eV
Conversion factor from Hartree to kcal/mol
Conversion factor from Hartree to kJ/mol
Conversion factor from Hartree to J T^-2
Conversion factor from Hartree to cm^-1
Speed of light in vacuo in atomic units

Output schema

```

"output": {
    "success": bool,
    succeeded
    "schema_name": string,
    "schema_version": int,
    "provenance": {
        obtained
    }
}

```

Whether all requested calculations succeeded
Name of the output schema
Version of the output schema
Information on how the results were obtained

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```

"creator": string,                                # Program name
"version": string,                               # Program version
"nthreads": int,                                 # Number of OpenMP threads used
"mpi_processes": int,                            # Number of MPI processes used
"total_cores": int,                             # Total number of cores used
"routine": string,                             # The function that generated the output
},
"properties": {                                    # Collection of final properties
    "charge": int,                                # Total molecular charge
    "multiplicity": int,                           # Total spin multiplicity
    "center_of_mass": array[float],                # Center of mass coordinate
    "geometry": array[ [ {                         # Array of atoms
        "symbol": string,                          # (one entry per atom)
        "xyz": array[float] } ] ],                  # Atomic symbol
                                                # Cartesian coordinate
],
"orbital_energies": {                            # Collection of orbital energies
    "spin": array[string],                        # Array of spins ('p', 'a' or 'b')
    "energy": array[float],                      # Array of energies
    "occupation": array[int],                   # Array of orbital occupations
    "sum_occupied": float,                      # \sum_i occupation[i]*energy[i]
},
"scf_energy": {                                    # Collection of energy contributions
    "E_kin": float,                             # Kinetic energy
    "E_nn": float,                            # Classical nuclear-nuclear interaction
    "E_en": float,                            # Classical electron-nuclear interaction
    "E_ee": float,                            # Classical electron-electron interaction
    "E_next": float,                           # Classical nuclear-external field
},
interaction: {                                     # Classical electron-external field
    "E_eext": float,
},
interaction: {                                     # Hartree-Fock exact exchange energy
    "E_x": float,                            # DFT exchange-correlation energy
    "E_xc": float,                           # Sum of electronic contributions
    "E_el": float,                            # Sum of nuclear contributions
    "E_nuc": float,                           # Sum of all contributions
    "E_tot": float,                           # Electronic reaction energy
    "Er_el": float,                           # Nuclear reaction energy
    "Er_nuc": float,                           # Sum of all reaction energy contributions
},
"dipole_moment": {                                # Collection of electric dipole moments
    id (string): {                            # Unique id: 'dip-${number}'
        "r_0": array[float],                    # Gauge origin vector
        "vector": array[float],                 # Total dipole vector
        "vector_el": array[float],              # Electronic dipole vector
        "vector_nuc": array[float],             # Nuclear dipole vector
        "magnitude": float,                   # Magnitude of total vector
    }
},
"quadrupole_moment": {                            # Collection of electric quadrupole moments
    id (string): {                            # Unique id: 'quad-${number}'}
}
}

```

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```

    "r_0": array[float],                                # Gauge origin vector
    "tensor": array[float],                             # Total quadrupole tensor
    "tensor_el": array[float],                         # Electronic quadrupole tensor
    "tensor_nuc": array[float]                          # Nuclear quadrupole tensor
  }
},
"polarizability": {
  id (string): {
    "frequency": float,                            # Unique id: 'pol-${frequency}'
    "r_0": array[float],                           # Gauge origin vector
    "tensor": array[float],                         # Full polarizability tensor
    "isotropic_average": float                     # Diagonal average
  }
},
"magnetizability": {
  id (string): {
    "frequency": float,                            # Unique id: 'mag-${frequency}'
    "r_0": array[float],                           # Perturbation frequency
    "tensor": array[float],                         # Gauge origin vector
    "tensor_dia": array[float],                    # Full magnetizability tensor
    "tensor_para": array[float],                  # Diamagnetic tensor
    "isotropic_average": float                   # Paramagnetic tensor
  }
},
"nmr_shielding": {
  id (string): {
    "r_0": array[float],                           # Collection of NMR shielding tensors
    "r_K": array[float],                           # Unique id: 'nmr-${nuc_idx}+${atom_symbol}'
    "tensor": array[float],                         # Gauge origin vector
    "tensor_dia": array[float],                    # Nuclear coordinate vector
    "tensor_para": array[float],                  # Full NMR shielding tensor
    "diagonalized_tensor": array[float],          # Diamagnetic tensor
    "isotropic_average": float,                   # Paramagnetic tensor
    "anisotropy": float,                          # Diagonalized tensor used for (an)isotropy
    "anisotropy": float,                          # Diagonal average
    "anisotropy": float                          # Anisotropy of tensor
  }
},
"geometric_derivative": {
  id (string): {
    "electronic": array[float],                  # Collection of geometric derivatives
    "electronic_norm": float,                     # Unique id: 'geom-${number}'
    "geoemtric derivative":                      # Electronic component of the geometric_
    "nuclear": array[float],                      # Norm of the electronic component of the_
    "geoemtric derivative":                      # Nuclear component of the geometric_
    "nuclear_norm": float,                       # Norm of the nuclear component of the_
    "geometric derivative":                      # Geometric derivative
    "total": array[float],                        # Norm of the geometric derivative
    "total_norm": float
  }
},
"scf_calculation": {                                # Ground state SCF calculation
}

```

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```

"success": bool,
"initial_energy": {
    "E_kin": float,
    "E_nn": float,
    "E_en": float,
    "E_ee": float,
    "E_next": float,
},
"interaction": {
    "E_eext": float,
},
"interaction": {
    "E_x": float,
    "E_xc": float,
    "E_el": float,
    "E_nuc": float,
    "E_tot": float,
    "Er_el": float,
    "Er_nuc": float,
    "Er_tot": float
},
},
"scf_solver": {
    "converged": bool,
    "wall_time": float,
    "cycles": array[
        {
            "energy_total": float,
            "energy_update": float,
            "mo_residual": float,
            "wall_time": float,
            "energy_terms": {
                "E_kin": float,
                "E_nn": float,
                "E_en": float,
                "E_ee": float,
                "E_next": float,
            }
        }
    ]
},
"rsp_calculations": {
    id (string): {
        "success": bool,
        "initial_energy": {
            "E_kin": float,
            "E_nn": float,
            "E_en": float,
            "E_ee": float,
            "E_next": float,
        },
        "interaction": {
            "E_eext": float,
        },
        "interaction": {
            "E_x": float,
            "E_xc": float,
            "E_el": float,
            "E_nuc": float,
            "E_tot": float,
            "Er_el": float,
            "Er_nuc": float,
            "Er_tot": float
        },
        "details": {
            "converged": bool,
            "wall_time": float,
            "cycles": array[
                {
                    "energy_total": float,
                    "energy_update": float,
                    "mo_residual": float,
                    "wall_time": float,
                    "energy_terms": {
                        "E_kin": float,
                        "E_nn": float,
                        "E_en": float,
                        "E_ee": float,
                        "E_next": float,
                    }
                }
            ]
        }
    }
},
"rsp_calculations": {
    id (string): {
        "success": bool,
        "initial_energy": {
            "E_kin": float,
            "E_nn": float,
            "E_en": float,
            "E_ee": float,
            "E_next": float,
        },
        "interaction": {
            "E_eext": float,
        },
        "interaction": {
            "E_x": float,
            "E_xc": float,
            "E_el": float,
            "E_nuc": float,
            "E_tot": float,
            "Er_el": float,
            "Er_nuc": float,
            "Er_tot": float
        },
        "details": {
            "converged": bool,
            "wall_time": float,
            "cycles": array[
                {
                    "energy_total": float,
                    "energy_update": float,
                    "mo_residual": float,
                    "wall_time": float,
                    "energy_terms": {
                        "E_kin": float,
                        "E_nn": float,
                        "E_en": float,
                        "E_ee": float,
                        "E_next": float,
                    }
                }
            ]
        }
    }
},
"rsp_calculations": {
    id (string): {
        "success": bool,
        "initial_energy": {
            "E_kin": float,
            "E_nn": float,
            "E_en": float,
            "E_ee": float,
            "E_next": float,
        },
        "interaction": {
            "E_eext": float,
        },
        "interaction": {
            "E_x": float,
            "E_xc": float,
            "E_el": float,
            "E_nuc": float,
            "E_tot": float,
            "Er_el": float,
            "Er_nuc": float,
            "Er_tot": float
        },
        "details": {
            "converged": bool,
            "wall_time": float,
            "cycles": array[
                {
                    "energy_total": float,
                    "energy_update": float,
                    "mo_residual": float,
                    "wall_time": float,
                    "energy_terms": {
                        "E_kin": float,
                        "E_nn": float,
                        "E_en": float,
                        "E_ee": float,
                        "E_next": float,
                    }
                }
            ]
        }
    }
}
}

```

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2.2.6 Running a geometry optimization

In the following we will assume to have a valid user input file for the water molecule called `h2o.inp`, e.g. like this

```

world_prec = 1.0e-6
world_unit = angstrom

WaveFunction {
    method = lda
}

Molecule {
$coords
O          0.00000      0.00000      0.11779
H          0.00000      0.75545     -0.47116
H          0.00000     -0.75545     -0.47116
$end
}

```

A geometry optimization can be run by adding **GeometryOptimizer** section to any normal .inp file and setting the run keyword to **true**:

```
GeometryOptimizer{  
    run = true  
}
```

This will start a geometry optimization with the default settings.

Obtaining accurate forces

In the above H₂O input example the `world_prec` parameter is chosen really small. This is necessary to get accurate forces. If a looser precision it is chosen, the geometry optimization may not converge. Pay attention to the warning:

```
WARNING: Noise in force is larger than 0.2 times the largest force component!!!
```

```
Geometry optimization convergence cannot be guaranteed!!!
```

This is printed when the noise level is too high. Usually, geometry optimizations will not converge when this warning is printed. In that case, either tighten the `world_prec`, `orb_thrs` (or both) or loosen the convergence criterion of the geometry optimization.

Pre-relax input geometries

Running high precision multi resolution wavelet calculations is computationally expensive. It is therefore not advisable to use an input geometry with high forces, a small `world_prec` and start the simulation. An optimized workflow would look something like this:

1. Optimize the geometry with a gaussian basis set. This can be done with a number of gaussian basis set codes
2. Use inaccurate forces (`world_prec ~ 1e-4`) and a rather loose convergence criterion (`max_force_component ~ 1e-2`) for the geometry optimization for a pre-relaxation with MRChem.
3. Do a tight geometry optimization (`max_force_component ~ 5e-4`) and with an accurate MRChem calculation (`world_prec ~ 1e-6`)

Reuse orbitals

For tight geometry optimizations where the input structure is already close to the local minimum (using cheaper pre-relaxations), it makes sense to use the orbitals from the geometry optimization iteration i for the start of iteration $i+1$. This feature can be enabled by setting:

```
use_previous_guess = true
```

Choosing an initial step size

If there are some problems in the first couple of geometry optimization iterations (energy and force norm increasing) the initial step size should be chosen manually. If a conservative choice (`init_step_size ~ 0.8`) does not solve the problem, the problem is usually in the input geometry (wrong units, unphysical, ...) or in the potential energy surface (too much noise, error in the DFT input section, ...).

Convergence problem can be analyzed by visualizing the optimization trajectory and plots of the energy and force norm versus the geometry optimization iterations.

2.3 Programmer's Manual

2.3.1 Classes and functions reference

Chemistry

Classes for the chemistry overlay

Environment

Classes for the solvent environment overlay

Cavity

class **Cavity** : public mrcpp::RepresentableFunction<3>

Interlocking spheres cavity centered on the nuclei of the molecule.

The *Cavity* class represents the following function Fosso-Tande2013

$$C(\mathbf{r}) = 1 - \prod_{i=1}^N (1 - C_i(\mathbf{r}))$$

$$C_i(\mathbf{r}) = 1 - \frac{1}{2} \left(1 + \operatorname{erf} \left(\frac{|\mathbf{r} - \mathbf{r}_i| - R_i}{\sigma_i} \right) \right)$$

where \mathbf{r} is the coordinate of a point in 3D space, \mathbf{r}_i is the coordinate of the i -th nucleus, R_i is the radius of the i -th sphere, and σ_i is the width of the transition between the inside and outside of the cavity. The transition has a sigmoidal shape, such that the boundary is a smooth function instead of sharp boundaries often seen in other continuum models. This function is 1 inside and 0 outside the cavity.

The radii are computed as:

$$R_i = \alpha_i R_{0,i} + \beta_i \sigma_i$$

where:

- $R_{0,i}$ is the atomic radius. By default, the van der Waals radius.
- α_i is a scaling factor. By default, 1.1
- β_i is a width scaling factor. By default, 0.5
- σ_i is the width. By default, 0.2 bohr

Public Functions

Cavity(const std::vector<mrcpp::Coord<3>> &coords, const std::vector<double> &R, const std::vector<double> &alphas, const std::vector<double> &betas, const std::vector<double> &sigmas)

Initializes the members of the class and constructs the analytical gradient vector of the *Cavity*.

inline **Cavity**(const std::vector<mrcpp::Coord<3>> &coords, const std::vector<double> &R, double sigma)

Initializes the members of the class and constructs the analytical gradient vector of the *Cavity*.

This CTOR applies a single width factor to the cavity and **does** not modify the radii. That is, in the formula:

$$R_i = \alpha_i R_{0,i} + \beta_i \sigma_i$$

for every atom i , $\alpha_i = 1.0$ and $\beta_i = 0.0$.

double **evalf**(const mrcpp::Coord<3> &r) const override

Evaluates the value of the cavity at a 3D point r.

Parameters

r – coordinate of 3D point at which the *Cavity* is to be evaluated at.

Returns

double value of the *Cavity* at point r

inline std::vector<mrcpp::Coord<3>> **getCoordinates**() const

Returns *centers*.

inline std::vector<double> **getOriginalRadii**() const

Returns *radii_0*.

inline std::vector<double> **getRadii**() const

Returns *radii*.

inline std::vector<double> **getRadiiScalings**() const

Returns *alphas*.

inline std::vector<double> **getWidths**() const

Returns *sigmas*.

inline std::vector<double> **getWidthScalings**() const

Returns *betas*.

void **printParameters**() const

Print parameters.

Protected Attributes

std::vector<double> **radii_0**

Contains the *unscaled* radius of each sphere in #Center.

std::vector<double> **alphas**

The radius scaling factor for each sphere.

`std::vector<double> betas`

The width scaling factor for each sphere.

`std::vector<double> sigmas`

The width for each sphere.

`std::vector<double> radii`

Contains the radius of each sphere in #Center. $R_i = \alpha_i R_{0,i} + \beta_i \sigma_i$.

`std::vector<mrcpp::Coord<3>> centers`

Contains each of the spheres centered on the nuclei of the Molecule.

Related

```
auto gradCavity(const mrcpp::Coord<3> &r, int index, const std::vector<mrcpp::Coord<3>> &centers, const
                std::vector<double> &radii, const std::vector<double> &widths) -> double
```

Constructs a single element of the gradient of the *Cavity*.

This constructs the analytical partial derivative of the *Cavity* C with respect to x , y or z coordinates and evaluates it at a point \mathbf{r} . This is given for x by

$$\frac{\partial C(\mathbf{r})}{\partial x} = (1 - C(\mathbf{r})) \sum_{i=1}^N -\frac{(x - x_i) e^{-\frac{s_i^2(\mathbf{r})}{\sigma^2}}}{\sqrt{\pi} \sigma \left(0.5 \operatorname{erf}\left(\frac{s_i(\mathbf{r})}{\sigma}\right) + 0.5 \right) |\mathbf{r} - \mathbf{r}_i|}$$

where the subscript i is the index related to each sphere in the cavity, and s is the signed normal distance from the surface of each sphere.

Parameters

- **r** – The coordinates of a test point in 3D space.
- **index** – An integer that defines the variable of differentiation (0->x, 1->y and 2->z).
- **centers** – A vector containing the coordinates of the centers of the spheres in the cavity.
- **radii** – A vector containing the radii of the spheres.
- **width** – A double value describing the width of the transition at the boundary of the spheres.

Returns

A double number which represents the value of the differential (w.r.t. x, y or z) at point \mathbf{r} .

Permittivity

```
class Permittivity : public mrchem::StepFunction
```

Permittivity function related to a substrate molecule and a solvent continuum. The *Permittivity* class represents the following function Fosso-Tande2013.

$$\epsilon(\mathbf{r}) = \epsilon_{in} \exp \left(\left(\log \frac{\epsilon_{out}}{\epsilon_{in}} \right) (1 - C(\mathbf{r})) \right)$$

where \mathbf{r} is the coordinate of a point in 3D space, C is the #cavity function of the substrate, and ϵ_{in} and ϵ_{out} are the dielectric constants describing, respectively, the permittivity inside and outside the #cavity of the substrate.

Public Functions

Permittivity(std::shared_ptr<*Cavity*> cavity, double epsilon_in, double epsilon_out, std::string formulation)

Standard constructor. Initializes the #cavity, #epsilon_in and #epsilon_out with the input parameters.

Parameters

- **cavity** – interlocking spheres of *Cavity* class.
- **epsilon_in** – permittivity inside the #cavity.
- **epsilon_out** – permittivity outside the #cavity.
- **formulation** – Decides which formulation of the *Permittivity* function to implement, only exponential available as of now.

double **evalf**(const mrcpp::Coord<3> &r) const override

Evaluates *Permittivity* at a point in 3D space with respect to the state of #inverse.

Parameters

r – coordinates of a 3D point in space.

Returns

$\frac{1}{\epsilon(\mathbf{r})}$ if #inverse is true, and $\epsilon(\mathbf{r})$ if #inverse is false.

DHScreening

class **DHScreening** : public mrchem::StepFunction

Square of the Debye-Huckel Screening parameter.

This is used for the Poisson-Boltzmann solver. The *DHScreening* function is defined as

$$\kappa^2(\mathbf{r}) = \begin{cases} \kappa_{out}^2 & \text{if } \mathbf{r} \notin \Omega_{ion} \\ 0.0 & \text{if } \mathbf{r} \in \Omega_{ion} \end{cases}$$

This can be parametrized a number of ways. The one used here is

$$\kappa^2(\mathbf{r}) = (1 - C_{ion}(\mathbf{r}))\kappa_{out}^2$$

Where $C_{ion}(\mathbf{r})$ is the ion accessible *Cavity* function.

Public Functions

DHScreening(std::shared_ptr<*Cavity*> cavity_ion, double kappa_out, const std::string &formulation)

Standard constructor. Initializes the #cavity_ion and #kappa_out with the input parameters.

#kappa_out is given by

$$\kappa = \sqrt{\frac{2000I_0e^2N_aI_0}{\epsilon_{out}\epsilon_{in}k_BT}}$$

where N_a is the Avogadro constant, e is the elementary charge, I_0 is the concentration of the ions, k_B is the Boltzmann constant, T is the temperature, ϵ_{out} is the permittivity of the solvent and ϵ_{in} is the permittivity of free space.

Parameters

- **cavity_ion** – interlocking spheres of *Cavity* class.
- **kappa_out** – value of the screening function outside the #cavity_ion.
- **formulation** – Decides which formulation of the *DHScreening* function to implement, only continuous screening function available.

double **evalf**(const mrcpp::Coord<3> &r) const override

Evaluates *DHScreening* at a point in 3D space.

Parameters

r – coordinates of a 3D point in space.

Returns

Value at point r.

Private Members

std::string **formulation** = {"Continuous Screening Function"}

Formulation of the *DHScreening* function. Only linear variable is used now.

GPESolver**class GPESolver**

Solves the Generalized Poisson equation iteratively.

The Generalized Poisson equation is given by

$$\nabla \cdot (\epsilon(\mathbf{r}) \nabla V(\mathbf{r})) = -4\pi\rho(\mathbf{r})$$

where $\epsilon(\mathbf{r})$ is the permittivity, $V(\mathbf{r})$ is the total electrostatic potential and $\rho(\mathbf{r})$ is the molecular charge density defined as:

$$\rho(\mathbf{r}) = \rho_{el}(\mathbf{r}) + \rho_{nuc}(\mathbf{r})$$

where ρ_{el} is the electronic charge density and ρ_{nuc} is the nuclear charge density. The Generalized Poisson equation is solved iteratively through a set of micro-iteration on each SCF-iteration by application of the Poisson operator \mathcal{P} :cite:Fosso-Tande2013

$$V_R(\mathbf{r}) = \mathcal{P} \star [\rho_{eff}(\mathbf{r}) - \rho(\mathbf{r}) + \gamma_s(\mathbf{r})]$$

where $\gamma_s(\mathbf{r})$ is the surface charge distribution describing the polarization at the surface, $\rho_{eff}(\mathbf{r})$ is the effective charge density given by $\frac{\rho(\mathbf{r})}{\epsilon(\mathbf{r})}$ and $V_R(\mathbf{r})$ is the reaction potential.

We utilize a so-called dynamic threshold to more easily converge the reaction potential. This is done by setting the convergence threshold of the micro-iterations to the MO update of the previous SCF iteration, unless the MO update is small enough (once the quality of the MOs is good enough, we use the default convergence threshold). Another optimization used is that we utilize the previous SCF converged Reaction potential as an initial guess for the next micro-iterations. These procedures are investigated and explained in :cite:gerez2023

Subclassed by *mrchem::PBESolver*

Public Functions

double setConvergenceThreshold(double prec)

Sets the convergence threshold for the micro-iterations, used with dynamic thresholding.

will check if the MO update is small enough (ten times as big) wrt. to the scf convergence threshold, if so, it will use the default convergence threshold. If not, it will use the MO update as the convergence threshold.

Parameters

prec – value to set the convergence threshold to

Returns

the current convergence threshold.

auto computeEnergies(const Density &rho_el) -> std::tuple<double, double>

Computes the energy contributions from the reaction potential.

We compute the reaction energy through the following integral:

$$E_R = \frac{1}{2} \int \rho_{el}(\mathbf{r}) V_R(\mathbf{r}) d\mathbf{r} + \frac{1}{2} \int \rho_{nuc}(\mathbf{r}) V_R(\mathbf{r}) d\mathbf{r}$$

Each term represents the electronic and nuclear contributions to the reaction energy, respectively. We compute each term separately, and return a tuple containing both.

Parameters

rho_el – the electronic charge density

Returns

a tuple containing the electronic and nuclear energy contributions

Protected Functions

void computeDensities(const Density &rho_el, Density &rho_out)

computes density wrt. the density_type variable

The total charge density is given by the sum of the electronic and nuclear charge densities:

$$\rho(\mathbf{r}) = \rho_{el}(\mathbf{r}) + \rho_{nuc}(\mathbf{r})$$

where ρ_{el} is the electronic charge density and ρ_{nuc} is the nuclear charge density. The nuclear charge density is stored in the class variable rho_nuc, while we compute the electronic charge density from the molecular orbitals. The class variable *density_type* decides the density which will be computed in rho_out, options are **total**, **electronic** and **nuclear**.

Parameters

• **Phi** – the molecular orbitals

• **rho_out** – Density function in which the density will be computed.

virtual void computeGamma(mrcpp::ComplexFunction &potential, mrcpp::ComplexFunction &out_gamma)

Computes the surface charge distribution due to polarization at the solute-solvent boundary.

The surface charge distribution is given by

$$\gamma_s(\mathbf{r}) = \frac{\log \frac{\epsilon_{in}}{\epsilon_{out}}}{4\pi} (\nabla C(\mathbf{r}) \cdot \nabla V(\mathbf{r}))$$

where ϵ_{in} is the permittivity inside the cavity and ϵ_{out} is the permittivity outside the cavity.

Parameters

- **potential** – Potential used to compute $\nabla V(\mathbf{r})$
- **out_gamma** – ComplexFunction in which the surface charge distribution will be computed.

```
mrcpp::ComplexFunction solvePoissonEquation(const mrcpp::ComplexFunction &ingamma, const Density &rho_el)
```

Iterates once through the Generalized Poisson equation to compute the reaction potential.

Constructs the effective charge density $\rho_{eff}(\mathbf{r})$ and the Poisson operator \mathcal{P} as:

$$V_R(\mathbf{r}) = \mathcal{P} \star [\rho_{eff}(\mathbf{r}) - \rho(\mathbf{r}) + \gamma_s(\mathbf{r})]$$

where $\gamma_s(\mathbf{r})$ is the surface charge distribution describing the polarization at the surface, $\rho_{eff}(\mathbf{r})$ is the effective charge density given by $\frac{\rho(\mathbf{r})}{\epsilon(\mathbf{r})}$ and $V_R(\mathbf{r})$ is the reaction potential.

Parameters

- **ingamma** – the surface charge distribution
- **Phi** – the molecular orbitals

Returns

the reaction potential

```
void accelerateConvergence(mrcpp::ComplexFunction &dfunc, mrcpp::ComplexFunction &func, KAIN &kain)
```

Uses KAIN to accelerate convergece of the reaction potential.

Parameters

- **dfunc** – the current update of the reaction potential
- **func** – the current reaction potential
- **kain** – the KAIN object

```
void runMicroIterations(const mrcpp::ComplexFunction &V_vac, const Density &rho_el)
```

Iterates through the application of the Poisson operator to Solve the Generalized Poisson equation.

Iterating through the application of the Poisson operator is done through a set of micro-iterations, where the convergence threshold is set to the MO update of the previous SCF iteration. The micro-iterations are done through the following steps:

- Compute the total potential as $V(\mathbf{r}) = V_{vac}(\mathbf{r}) + V_R(\mathbf{r})$
- Compute the surface charge distribution $\gamma_s(\mathbf{r})$ with *computeGamma*
- Compute a new reaction potential $V_R(\mathbf{r})$ by application of the Poisson operator with *solvePoissonEquation*
- Calculate the update of the reaction potential as $\Delta V_R(\mathbf{r}) = V_R(\mathbf{r}) - V_R^{old}(\mathbf{r})$
- Accelerate convergence of the reaction potential through KAIN
- Update the reaction potential as $V_R(\mathbf{r}) = V_R^{old}(\mathbf{r}) + \Delta V_R(\mathbf{r})$
- Check if the reaction potential has converged, if not, repeat from step 1.

Parameters

- **V_vac** – the vacuum potential
- **Phi_p** – the molecular orbitals

```
mrcpp::ComplexFunction &solveEquation(double prec, const Density &rho_el)
```

Setups and computes the reaction potential through the microiterations.

An initial guess of the reaction potential is computed with the following steps:

- i. Set the total potential as $V(\mathbf{r}) = V_{vac}(\mathbf{r})$
- ii. Compute the surface charge distribution $\gamma_s(\mathbf{r})$ from this potential
- iii. Iterate once through the application of the Poisson operator to return the initial guess of the reaction potential $V_R(\mathbf{r})$

the method then runs the micro-iterations through `runMicroIterations` and returns the converged reaction potential. If this is not the first SCF iteration, the previous converged reaction potential is used as an initial guess for the micro-iterations.

Parameters

- **V_vac** – the vacuum potential
- **Phi_p** – the molecular orbitals

Returns

The converged reaction potential for the current SCF iteration

```
void resetComplexFunction(mrcpp::ComplexFunction &function)
```

Frees the memory used by the FunctionTrees of the input Complexfunction and reallocates them.

This is done to avoid memory leaks when the ComplexFunction is used in the micro-iterations.

Parameters

- **function** – the ComplexFunction to reset

Protected Attributes

SCRFDensityType **density_type**

Decides which density we will use for computing the reaction potential, options are **total**, **electronic** and **nuclear**.

PBESolver

```
class PBESolver : public mrchem::GPESolver
```

Solves the Poisson-Boltzmann equation iteratively.

The Poisson-Boltzmann equation is solved iteratively using the SCRF procedure outlined in [GPESolver](#). The Poisson-Boltzmann equation models the electrostatic potential in a solvent with electrolytes. The general equation for electrolyte solutions is given by

$$\nabla \cdot \epsilon \nabla V_{tot} = -4\pi (\rho_{el} + \rho_{nuc} + \rho_{ext})$$

where V_{tot} is the total electrostatic potential, ϵ is the permittivity function of the solvent, ρ_{el} is the electronic charge density, ρ_{nuc} is the nuclear charge density and ρ_{ext} is the external charge density. In the general form for the Poisson-Boltzmann equation, the external charge density is approximated by assuming a boltzmann distribution of the ions.

$$\rho_{ext} = \sum_i^{N_{ion}} q_i e I_{0,i} \exp\left(-\frac{q_i e V_{tot}}{k_B T}\right)$$

where $I_{0,i}$ is the concentration of the i-th ion species, q_i is the charge of the i-th ion species, k_B is the Boltzmann constant and T is the temperature. In this implementation we assume a 1:1 ($I_{0,0} = I_{0,1}$) electrolyte solution of ions of same opposite charges ($z_i = +1, -1$). This simplifies the external density to

$$\rho_{ext} = -2eI_0 \sinh\left(\frac{eV_{tot}}{2k_B T}\right)$$

where I_0 is the concentration of the ions. We can plug this into the first equation (and massage terms a bit) to arrive at the Poisson-Boltzmann equation for 1:1 electrolyte solution

$$\nabla^2 V_R = -4\pi \frac{1-\epsilon}{\epsilon} (\rho_{el} + \rho_{nuc}) + \gamma_s - \kappa^2 \sinh(V_{tot})$$

where γ_s is the surface charge density, κ is obtained from the *DHScreening* class and V_R is the reaction potential.

Subclassed by *mrchem::LPBESolver*

Protected Functions

```
virtual void computeGamma(mrcpp::ComplexFunction &potential, mrcpp::ComplexFunction &out_gamma)
    override
```

constructs the surface chage distribution and adds it to the PB term

Method follows the implementation in *GPESolver::computeGamma*, but adds the PB term to the surface charge distribution.

Parameters

- **potential** – [in] the potential to compute ∇V from
- **out_gamma** – [out] the ComplexFunction in which to store the result

```
virtual void computePBTerm(mrcpp::ComplexFunction &V_tot, const double salt_factor,
    mrcpp::ComplexFunction &pb_term)
```

Computes the PB term.

The PB term is computed as $\kappa^2 \sinh(V_{tot})$ and returned.

Parameters

- **V_tot** – [in] the total potential
- **salt_factor** – [in] the salt factor deciding how much of the total concentration to include in the PB term
- **pb_term** – [out] the ComplexFunction in which to store the result

Protected Attributes

DHScreening **kappa**

the *DHScreening* object used to compute the PB term κ

LPBESolver

```
class LPBESolver : public mrchem::PBESolver
```

Solves the Linearized Poisson-Boltzmann equation iteratively.

The Linearized Poisson-Boltzmann equation is solved iteratively using the SCRF procedure outlined in *GPE-Solver* and *PBESolver*. The linearized Poisson-Boltzmann equation is a further simplification of the Poisson-Boltzmann equation, outlined in *PBESolver*, where the PB term is expanded and only the linear term is included. This is a good approximation for low ionic strength solutions. The linearized Poisson-Boltzmann equation is given by

$$\nabla^2 V_R = -4\pi \frac{1-\epsilon}{\epsilon} (\rho_{el} + \rho_{nuc}) + \gamma_s - \kappa^2 V_{tot}$$

where γ_s is the surface charge density, κ is obtained from the *DHScreening* class and V_R is the reaction potential.

Protected Functions

```
virtual void computePBTerm(mrcpp::ComplexFunction &V_tot, const double salt_factor,  
                           mrcpp::ComplexFunction &pb_term) override
```

Computes the PB term.

The PB term is computed as $\kappa^2 V_{tot}$ and returned.

Parameters

- **V_tot** – [in] the total potential
- **salt_factor** – [in] the salt factor deciding how much of the total concentration to include in the PB term
- **pb_term** – [out] the ComplexFunction in which to store the result

Initial Guess

Classes providing the initial guess of the orbitals

Properties

Classes for the calculation of molecular properties

Quantum Mechanical Functions

Classes to handle quantum mechanical functions such as electronic density, molecular orbitals.

QMOperators

The classes that implement quantum mechanical operators

QMPotential

class QMPotential

Operator defining a multiplicative potential.

Inherits the general features of a complex function from mrcpp::ComplexFunction and implements the multiplication of this function with an Orbital. The actual function representing the operator needs to be implemented in the derived classes, where the *re and *im FunctionTree pointers should be assigned in the setup() function and deallocated in the clear() function.

XCOoperator

class XCOoperator

DFT Exchange-Correlation operator containing a single *XCPotential*.

This class is a simple TensorOperator realization of

XCPotential

class XCPotential

Exchange-Correlation potential defined by a particular (spin) density.

The XC potential is computed by mapping of the density through a XC functional, provided by the XCFun library. There are two ways of defining the density:

- 1) Use `getDensity()` prior to `setup()` and build the density as you like.
- 2) Provide a default set of orbitals in the constructor that is used to compute the density on-the-fly in `setup()`.

If a set of orbitals has NOT been given in the constructor, the density MUST be explicitly computed prior to `setup()`. The density will be computed on-the-fly in `setup()` ONLY if it is not already available. After `setup()` the operator will be fixed until `clear()`, which deletes both the density and the potential.

LDA and GGA functionals are supported as well as two different ways to compute the XC potentials: either with explicit derivatives or gamma-type derivatives.

ReactionPotential

class ReactionPotential : public mrchem::QMPotential

class containing the solvent-substrate interaction reaction potential obtained by solving

$$\Delta V_R = -4\pi \left(\rho \frac{1 - \epsilon}{\epsilon} + \gamma_s \right)$$

where ρ is the total molecular density of a solute molecule, ϵ is the *Permittivity* function of the continuum and γ_s is the surface charge distribution.

Subclassed by `mrchem::ReactionPotentialD1`, `mrchem::ReactionPotentialD2`

Public Functions

```
explicit ReactionPotential(std::unique_ptr<GPESolver> scrf, std::shared_ptr<mrchem::OrbitalVector>
                           Phi = nullptr, bool mpi_share = false)
```

Initializes the *ReactionPotential* class.

Parameters

- **scrf** – A *GPESolver* instance which contains the parameters needed to compute the *ReactionPotential*.
- **Phi** – A pointer to a vector which contains the orbitals optimized in the SCF procedure.

```
inline void updateMOResidual(double const err_t)
```

Updates the solver.mo_residual member variable. This variable is used to set the convergence criterion in the dynamic convergence method.

Protected Attributes

```
std::unique_ptr<GPESolver> solver
```

A *GPESolver* instance used to compute the *ReactionPotential*.

```
std::shared_ptr<OrbitalVector> orbitals
```

Unperturbed orbitals defining the ground-state electron density for the SCRF procedure.

SCF Solver

Classes for the resolution of the SCF equations of HF and DFT

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