# OPENCAP-MD manual

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## 1 Directory Structure of OPENCAP-MD

Below is the structured view of the OPENCAP-MD package.

## 2 Description of Files and Directories

### **Top-Level Directories**

• optimizer/: Contains scripts for optimizing molecular structures using different quantum chemistry packages. Available interfaces are:

```
optimizer_MOLCAS.py: Geometry optimizer for OpenMolcas optimizer_columbus.py: Geometry optimizer for Columbus
```

- utils/: Includes various utility functions for handling molecular data and parsing input/output files.
- sharc-interface/: Provides a MOLCAS-OPENCAP interface with SHARC AIMD package. Also used by optimizer\_MOLCAS.py.
- examples/: Contains example cases for running OpenMolcas and Columbus based geometry optimizations.

#### 2.1 OpenMolcas specific files

The files needed for OpenMolcas based optimization is structured as shown in 2.1. For reference see examples/openmolcas directory.

```
OPENCAP-MD
|-- examples
    |-- openmolcas
         |-- run-opencapmd.sh
         |-- geom
         |-- QM
              |-- CC-PVTZ-3P
              |-- MOLCAS.template
              |-- OPENCAPMD.gbs
              |-- MOLCAS.resources
              |-- runQM.sh
              Figure 2: Directory structure of OpenMolcas based calculation.
```

• geom: geometry file in SHARC or Columbus format. The columns in the following correspond to atom symbol, atomic number, x, y, z coordinates in bohr and lastly the atomic masses.

С	6.0	-0.033968	-0.214849	-0.021765	12.00000000
Н	1.0	-0.093819	0.027585	1.992555	1.00782504
Н	1.0	1.770523	0.027597	-0.918910	1.00782504
C	6.0	-2.094619	-0.788029	-1.341295	12.00000000
Н	1.0	-2.034768	-1.030463	-3.355614	1.00782504
Н	1.0	-3.899110	-1.030473	-0.444149	1.00782504
Figure 3: Geometry file structure for geom input.					

- QM/: A directory containing input files for SHARC\_MOLCAS\_OPENCAP.py interface (the file structure is similar to that of SHARC\_MOLCAS.py interface.).
  - CC-PVTZ-3P: Basis set file used for calculations only if a modified basis is used which is not a standard choice from molcas basis library. The basis file should have molcas basis-library file structure. See the file examples/openmolcas/QM/CC-PVTZ-3P for reference. This file should have a header as following

```
#Hamiltonian NRH
#Nucleus UNK
#Contraction CCC
#AllElectron AE_
```

and individual atom specific header should be renamed by the basis file name with proper count of uncontracted and contracted basis count in the header.

(see header in Fig. 4 /C.cc-pVTZ-3P.Dunning.10s8p2d1f.4s6p2d1f.).

```
/C.cc-pVTZ-3P.Dunning.10s8p2d1f.4s6p2d1f.
T.H. Dunning. J. Chem. Phys. 90 (1989) 1007-1023. doi:10.1063/1.456153
CARBON (10s, 8p, 2d, 1f) \rightarrow [4s, 6p, 2d, 1f]
6.0
* s-type functions
10
8236.00000000
1235.00000000
280.80000000
79.27000000
25.59000000
8.99700000
3.31900000
0.90590000
0.36430000
0.12850000
0.0005310000 \ -0.0001130000 \ \ 0.0000000000 \ \ 0.0000000000
0.0041080000 \ -0.0008780000 \ \ 0.0000000000 \ \ 0.0000000000
. . .
          Figure 4: Custom basis file for SHARC_MOLCAS_OPENCAP.py
```

```
* /X.OPENCAPMD.Diffuse.2s5p2d.2s5p2d.
* 2s5p2d on the ghost atom
* GHOST (2s,5p,2d) -> [2s,5p,2d]
0.00
    2
2
0.05135
0.025675
1.0000000E+00 0.00000000E+00
0.0000000E+00 1.00000000E+00
5 5
0.0806
0.05373
0.03582
0.02388
0.01592
1.0000000E+00 0.00000000E+00
                                  0.000000000E+00 0. 0.
0.0000000E+00 1.00000000E+00
                                  0.000000000E+00 0. 0.
0.0000000E+00 0.00000000E+00
                                 1.000000000E+00 0. 0.
0. 0. 0. 1. 0.
0. 0. 0. 0. 1.
   2
2
0.159
0.0795
1.0000000E+00 0.00000000E+00
0.0000000E+00 1.00000000E+00
Figure 5: Custom basis file for SHARC_MOLCAS_OPENCAP.py for ghost atom in COM
of current geometry.
```

— MOLCAS.template: Template file for SHARC\_MOLCAS\_OPENCAP.py. This is also a file standard to SHARC. This file contains keyword to run the SHARC\_MOLCAS\_OPENCAP.py interface. A sample file with explanation of keywords are given in the following section (Fig. 6). Keywords are not case sensitive.

```
basis cc-pvtz-3p
# Basis set chosen (in openmolcas-library format)
# In this example this is not a standard basisset
# available in molcas library
BASLIB /usr/ethene/SA15-3e20o-CCT3PA/QM/
# Basis file location for openmolcas input file
# if default basisset is not used
# In this example MOLCAS will seek for CC-PVTZ-3P file in BASLIB.
# This modified basisfile name is case sensitive.
ras2 20
# CAS orbital size
nactel 3
# No. of active electrons
inactive 7
# Inactive orbitals
roots 15
# Total no. of states in state averaging
charge -1
# Charge of the system
CAP_ETA 1771.25
# Fixed eta value
# (This value is scaled by 1E-5 in actual calculation.)
act_states 3
# Active state to follow
# (resoance state from initial calculation.)
cap_x 7.62
cap_y 4.20
cap_z 6.02
# CAP onset along all directions in atomic units.
calc_neutral
# Calculate neutral energy
# (if absent, take lowest root as a dummy neutral energy)
track_all
# Keyword for wf overlap
# Use RASSI module of openmolcas to calculate
# Other option is track_wfoverlap (the default choice).
natorb
# Print natural orbitals (real part of complex density)
# Need OPENCAPMD.gbs basis file in pyscf format
# Should be in QM/ directory
            Figure 6: MOLCAS.template file with necessary keywords.
```

- **OPENCAPMD.gbs**: Gaussian basis set file written in pyscf basis input format (needed if natorb is invoked in MOLCAS.template file.) The file may look like the Fig. 7.

```
spherical
***
H O
S 4 1.00
                         1.968500D-02
1.379770D-01
4.781480D-01
5.012400D-01
1.301000D+01
1.962000D+00
4.446000D-01
1.22000D-01
S 1 1.00
1.220000D-01 1.000000D+00
P 1 1.00
7.27000D-01
                                         1.0000000
****
С
          0
    9 1.00
6.665000D+03
                                       6.920000D-04
1.00000D+03
                                      5.329000D-03
                                      2.707700D-02
2.280000D+02

      2.250000D+02
      2.707700D-02

      6.471000D+01
      1.017180D-01

      2.106000D+01
      2.747400D-01

      7.495000D+00
      4.485640D-01

      2.797000D+00
      2.850740D-01

      5.215000D-01
      1.520400D-02

      1.596000D-01
      -3.191000D-03

. . . .
***
              Figure 7: A sample file snapshot of OPENCAPMD.gbs file read by pyscf.
```

MOLCAS.resources: Resource file for OpenMolcas jobs. This is also a standard file in SHARC.
 A sample file looks like the following. Keywords are not case sensitive.

```
memory 8800
# Memory per core (in MB), (not TOTAL Memory)
# If 256 GB Memory is distributed over 16 core,
# this means 16000 is the value to be used.
molcas $MOLCAS
# MOLCAS variable, can also use absolute path as well.
scratchdir /scratch/usr/ethene/3pA/
# Scratch directory
ncpu 1
# No. of cpu for EACH parallel run (NOT total cpus)
driver /usr/molcas_build/pymolcas
# pymolcas module path
WFOVERLAP /usr/WfOverlap/bin/wfoverlap.x
# WF-OVERLAP directory with the binary file (needed)
ncpu_avail 28
# Total no. of cpu for the job
delay 0.01
# Delay between parallel jobs (not needed strictly)
rasorb
# Read casscf orbitals (MOLCAS.RasOrb) files as guess
# for consecutive steps in geometry optimization cycles.
# (other option is scforb for SCF orbital guesses.)
optimizer geometric
# Optimizer chosen (Default is geometric)
            Figure 8: MOLCAS.resources file with necessary keywords.
```

- runQM.sh: Script to execute SHARC\_MOLCAS\_OPENCAP.py interface (default to SHARC). If not present, then it will be created by optimizer\_MOLCAS.py. A sample file looks like following (similar to SHARC-QC software interfaces).

```
#!/bin/sh
export SPATH=/opencapmd/sharc-interface/SHARC_MOLCAS_OPENCAP.py
cd QM || exit
"$SPATH" QM.in >> QM.log 2>> QM.err
exit $?
```

- run-opencapmd.sh: Bash script to execute OpenCAP-MD with OpenMolcas. The optimizer is called via the following command.

```
# Change to the path of the opencap-md
export OPTIMIZER_PATH=/path_TO_opencapmd/optimizer/
export OPTIMIZER="$OPTIMIZER_PATH/optimizer_MOLCAS.py"
$OPTIMIZER 'pwd' > logfile
```

optimizer\_MOLCAS.py takes command line argument of the directory path where the job is supposed to run.

• Outputs created: The output files created during geometry optimizations.

- iteration\_details: Logs iteration details from the optimization process.
- geometric.log, opt\_log.out, berny.log: A logfile generated by the geometric, optking, pyberny optimizers respectively.
- $-\mathbf{QM}/\mathbf{QM.log}$ : Log file from SHARC\_MOLCAS\_OPENCAP.py execution.
- QM/PCAP.out: Complex and neutral reference energies generated during the optimization.

## 2.2 Additional keywords

Keyword	Description
no_capgrad_correct	If user wants to skip the CAP gradient calculation.
ghost	This keyword is followed by a string with basis function count (e.g. Os.3p.3d for a 3p-3d function on ghost atom or 2s.2p for a 2s-2p basis on ghost atom). The keyowrd adds ghost atom in center of mass (COM) of the current geometry. This demands presence of a basis set in molcas format in QM/X.basis file. See fig. 5 for the QM/X.basis file for custom basis file structure.
scforb	If scf orbitals are used as guess for next iteration.
track_wfoverlap	If WF-OVERLAP package is used to calculate wave-function overlaps.
screening	Keyword followed by a float value (say, 0.0001) which works as a thershold to remove gradient and coupling jobs that have insignificant contributions.

Table 1: Description of some additional MOLCAS.template keywords

Keyword	Description
scforb	If SCF orbitals are used as guess for next iteration.
always_guess	If at each step new SCF guess is used.
debug	If all electronic structure calculation files are stored for debugging purpose (requires significant disk-space).
optimizer	three options: geometric, optking, pyberny

Table 2: Description of some additional MOLCAS.resources keywords

## 2.3 Special constraints/parameters for optimizers

#### 2.3.1 geometric, pyberny

The keywords in this following section are optimizer specific and can be added to MOLCAS.resources file and the optimizer will adapt accordingly.

Threshold keyword	Description
gradientmax	Maximum gradient in a.u
gradientrms	RMS gradient in a.u
stepmax	Maximum displacement in a.u
steprms	RMS displacement in a.u
deltae	Energy convergence in a.u

Table 3: Description of optimizer specific MOLCAS.resources threshold keywords

#### 2.3.2 optking

Create a file optking.params in run directory where optking optimizer specific keywords are invoked (see Psi4 OptKing Module Documentation). A sample input may look like Fig. 9

```
g_convergence qchem
ADD_AUXILIARY_BONDS True
opt_coordinates both
INCLUDE_OOFP False

Figure 9: A sample file for optking.params.
```

These above keywords are specific for optking module. There are several ways the same convergence parameters shown in Table 3 can be invoked for optking by changing the g\_convergence keyword in optking.params. A detailed logistics and available options are in optking convergence algorithm document.

#### 2.4 Columbus specific files

For reference please look at the file structure needed in the examples/columbus directory as shown in Fig. 10.

• COLUMBUS.template: Input template for Columbus based optimization.

```
nstates 5
# Total no of states in MR-CIS/MCSCF calculation calculation
eta_opt 0.00848
# optimized eta chosen (Scaled by 1.0)
# (In atomic units.)
act_state 2
# Active state to track
cap_x 2.77
cap_y 2.77
cap_z 4.88
\# cap onset in x, y and z directions
# in atomic units
natorb
# Print natural orbitals in DIAG basis
# (OPENCAPMD.gbs basis file is needed in pyscf format)
maxsqnorm 0.998001
# Maximum square norm as a cut-off.
\mbox{\# c^2} sum limits printing of all CSFs and Slater determinants
# (Applicable for WF-OVERLAP calculation only).
symmetry
# If point group Symmetry is invoked.
```

### 2.5 Additional keywords

Keyword	Description
calculation	Followed by a string. For SA-CASSCF calculation use mcscf. Default is MR-CIS, hence this keyword is not needed.
track	Followed by a string. Tracking method during geometry optimization.
	(When symmetry is on) two options are there: civecs and dens, which are done by CI-vector overlap and attachment-detachment densities, respectively. Default is dens.
	(When symmetry is off $(C_1)$ ) WF-OVERLAP is the default choice and this keyword is redundant.
no_capgrad_correct	If user wants to skip the CAP gradient calculation.
screening	Keyword followed by a float value (say, 0.0001) which works as a thershold used to remove gradient and coupling jobs that have insignificant contributions.
ghost_atom	Add ghost atom in center of mass (COM). Not available if symmetry is on.

 $\begin{tabular}{ll} Table 4: Description of some additional {\tt COLUMBUS.template} & keywords \\ \end{tabular}$ 

• COLUMBUS.resourcses: Input resource file for Columbus simulations. The keywords are not case sensitive.

```
WFOVERLAP/usr/wf-overlap/bin/wfoverlap.x
# wf-overlap binary file location

memory 240000
# Total Memory available

ncpu 16
# Total cpu available

optimizer optking
# Optimizer chosen optking (if Symmetry is on)

inputdir /usr/OPENCAPMD/N2/MRCIS/colfiles
# Input files for COLUMBUS run
# (If inputdir key is absent, default is 'copyfiles/' directory
# in the working directory.)
```

Keyword	Description
columbus	Columbus absolute path. If absent, the script will try to read \$COLUMBUS variable from command line.
scratchdir	Scratch directory where parallel gradient and coupling calculations are performed, this is disk-space intensive. If absent, the path is SAVEDIR/scratch in current working directory with same file-tree structure.
savedir	Absolute path where all Columbus electronic structure package runs it calculation. If absent, SAVEDIR is prepared in current work directory. Within this directory, folders from each iteration is present: sp_0, sp_1, sp_2, sp_3 and so on. Each folder will contain all Columbus run files and needed for debugging and restart.
restart	Followed by an integer value, that is the iteration number. This demands that savedir is accessible. Otherwise the restart will fail. The user is asked to resubmit a new job with the last geometry in a new directory.
optimizer	Optimizers available. Default is optking. Additionally you can use optking.params for more parameter control of the optimizer.
	(When symmetry is on) Use optking.
	(When symmetry is off $(C_1)$ ) Use geometric
debug	All Columbus single point calculation directories are saved (with Columbus WORK directory). Disk-space intensive.

Table 5: Description of some additional  ${\tt COLUMBUS.resources}$  keywords

• run-opencapmd.sh: Execution script for Columbus calculations. Same to that of OpenMolcas specific optimizer submission.

```
# Change to the path of the opencap-md
export OPTIMIZER_PATH=/path_TO_opencapmd/optimizer/
export OPTIMIZER="$OPTIMIZER_PATH/optimizer_columbus.py"
$OPTIMIZER 'pwd' > logfile
```

• optking.params: Configuration file for optimization settings specific to optking optimizer. See section 2.3.2.

```
colfiles/
|-- mcdrtin.*, mcscfin
# (and all MCSCF related files)
|-- ANION/
|--|-- ciudgin
# (MR-CI related files or MC-SCF restart files)
|--|- ABACUSIN, ABACUSIN.NAD (# and all other files)
# (These gradient and NAC specific files should be present)
|-- NEUTRAL/
Figure 11: Directory structure of Columbus files specific for projected-CAP calculation
```

• colfiles/: Contains all Columbus-related input \$COLUMBUS/colinp generated output files (not expanded here for brevity, see examples/columbus/colfiles for reference). User has to create the reference files that will be used for different iterations. These absolute path containing these files should be given in the inputdir keyword in the COLUMBUS.resources file. The main structure of the columbus is shown in 2.5 file tree.

The inputdir should have the MCSCF files generated by the user using \$COLUMBUS/colinp. For reference, the Columbus specific control.run file looks like the following.

```
niter = 1
hermit
scf
mcscf
```

Within the main inputdir, there are two main directories: ANION and NEUTRAL.

The ANION folder should have files to calculate MR-CIS energies, transition moments, gradients and couplings. The set up of ANION folder should have ABACUSIN.NAD and ABACUSIN for couplings and gradients respectively. Even though slope and nadcoupl is present in control.run file, these will be calculated in parallel after pciudg keyword is executed. The initial MO for this calculation is from mcscf run in the previous directory. For reference, the Columbus specific control.run file looks like the following.

```
niter = 1
hermit
ciudgmom
mcscf
pciudg
slope
nadcoupl
```

The NEUTRAL folder should have files for MR-CI calculation for neutral state. The initial MO for this calculation is from mcscf run in the previous directory.

niter = 1 hermit ciudg

#### 2.6 Working principle of optimizer\_columbus.py

This module is heavily dependent on *apriori* generation of Columbus iinput files using \$COLUMBUS/colinp. Given that the colfiles/ are present in the work directory.

- Changes geom file in each iteration (run in savedir/sp\_0 for iteration 0)
- Executes \$COLUMBUS/geom.unik.x to get symmetry unique atoms and coordinates in geom.unik file.
- Changes the daltaoin file with new symmetry unique atoms and coordinates. (Needed for \$COLUMBUS/dalton.x for AO integral generation)
- Perform SA-CASSCF calculation in the savedir/sp\_0 (say, for iteration 0) directory. Copy the MO guess (mocoef) in ANION, NEUTRAL folder.
- Run the Columbus job in ANION folder and generate the energies and transition moments for anionic states. Make sure that you use the parallelized module \$COLUMBUS/pciudg.x instead of \$COLUMBUS/ciudg.x to save a lot of computation time for MR-CI calculations.
  - After the energies and 1-RDMs are generated, gradients and couplings are calculated in parallel (using a combination of \$COLUMBUS/cigrd.x ,\$COLUMBUS/tran.x and \$COLUMBUS/dalton.x as prescribed in \$COLUMBUS/runc script).
- Once ANION folder execution is done, switch to NEUTRAL folder and calculate neutral energy.
- Finally, when all jobs are successful Projected CAP files needed for OpenCAP is generated and OPENCAP-MD is called to calculate complex energies and CAP-gradient augmented gradients.
- Iteration specific energy and gradients are fed to the optimizer.