



DMRG-CASPT2 with Molcas-CheMPS2 interface

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Log in Dirac

- Connect your laptop to eduroam
- All the information about the accounts is put on https://quantchem.kuleuven.be/owncloud

```
userid: password:
```

- Register using the URLs in b_network_registration_vouchers.txt
- Connect to the B-zone using the pulse-client



Log in Dirac

Connect to dirac, password provided

```
ssh tccmx@dirac.quantchem.kuleuven.be
git clone https://github.com/quanp/polyene
ssh node87
x=2,3,etc.
```

- Molcas-CheMPS2 installed at /home/tccm1/source
- interactively run molcas.sh on node87 (64 CPUs)



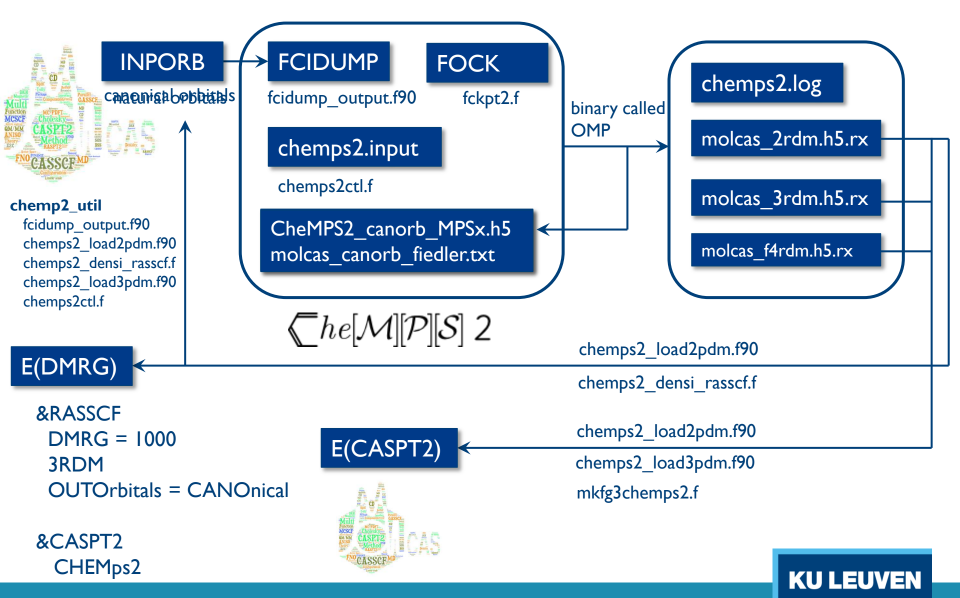
Molcas-CheMPS2 main features

CheMPS2

- ✓ DMRG-SCF with PCM
- ✓ SS- and SA- with DMRG-SCF calculations
- ✓ DMRG-CASPT2 with pseudocanonical basis
 - ✓ Not bad choice for compact molecules, e.g. TM complexes
 - ✓ Construction of F.4-RDM is simplified
 - × Not the optimal type of orbital for DMRG
- ✓ DMRG-CASPT2 with non-pseudocanonical basis implemented



DMRG-CASPT2 calculation with Molcas-CheMPS2



Installation

Cmake > 3.0

HDF5

https://support.hdfgroup.org/HDF5

must be configured with Fortran, Fortran 2003, and C++ enabled:

```
CC=icc CXX=icpc FC=ifort ./configure --enable-fortran
--enable-fortran2003 --enable-cxx
```



Installation

CheMPS2

https://github.com/SebWouters/CheMPS2

only Open Multi-Processing (OpenMP) is supported:

```
CC=icc CXX=icpc FC=ifort
CMAKE_INCLUDE_PATH=/path/to/hdf5/include
CMAKE_LIBRARY_PATH=/path/to/hdf5/lib cmake .. -DMKL=ON
-DWITH_MPI=OFF
```



Installation

Molcas

parallel Molcas supported

```
FC=ifort CC=icc CXX=icpc

cmake .. -DCMAKE_BUILD_TYPE=Release -DMPI=ON -DLINALG=MKL

-DMKLROOT=/path/to/intel/mkl -DMPI_C_COMPILER=mpicc

-DMPI_Fortran_COMPILER=mpif90 -DMPI_CXX_COMPILER=mpicxx

-DOPENMOLCAS_DIR=/path/to/OpenMolcas -DHDF5=ON

-DHDF5_LIBRARIES=/path/to/hdf5/lib

-DHDF5_INCLUDE_DIRS=/path/to/hdf5/include

-DCHEMPS2=ON -DCHEMPS2_DIR=/path/to/CheMPS2/bin
```



verification

molcas verify 850 851 --keep



Input:

```
&RASSCF

DMRG = 1000

3RDM
```

Dependencies:

- RUNFILE, INPORB
- molcas_natorb_fiedler.txt, CheMPS2_natorb_MPSx.h5 (x=0,1,...)
- molcas_canorb_fiedler.txt, CheMPS2_canorb_MPSx.h5
 (x=0,1,...)



Log

```
++ DMRG sweep specifications:
Number of renormalized basis
                                                              (DMRG)
                                           1000
Number of root(s) required
                                                              (CIROot)
                                           1
Maximum number of sweeps
                                          8
                                                              (MXSWeep)
Maximum number of sweeps in RDM
                                          40
                                                              (MXCAnonical)
Threshold for restarting
                                          0.500E-02
                                                              (CHBLb)
Minimum Davidson tolerance
                                                              (DAVTolerance)
                                          0.100E-06
DMRG convergence threshold
                                          0.500E-08
Noise prefactor
                                          0.500E-01
                                                              (NOISe)
Restart from previous calculation
                                                              (CHREstart)
Calculate 3-RDM and F.4-RDM
                                           Т
                                                              (3RDM)
Restart scheme in 3-RDM and F.4-RDM
                                          0
                                                              (DMREstart)
Root chosen for geometry opt.
                                           1
```

Output files

- log file
- chemps2.input chemps2.log chemps2.log.total
- FCIDUMP_CHEMPS2
- molcas_natorb_fiedler.txt
 molcas_canorb_fiedler.txt
- CheMPS2_natorb_MPSx.h5CheMPS2_canorb_MPSx.h5
- molcas_2rdm.h5.rx
- molcas_3rdm.h5.rx
- molcas_f4rdm.h5.rx
- FOCK_CHEMPS2



```
Input:
&CASPT2
CHEMps2
```

Dependencies:

- RUNFILE, JOBIPH
- molcas_2rdm.h5.rx
- molcas_3rdm.h5.rx
- molcas_f4rdm.h5.rx



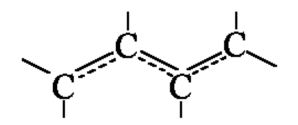
Log: very similar to conventional CASPT2 calculations

```
Multi-State initialization phase begins for group 1
CHEMPS2> MKRPTORB assumes PSEUDOCANONICAL orbitals!
Compute HO matrices for state
HO matrices have been computed.
CASPT2 EQUATION SOLUTION
Total nr of CASPT2 parameters:
 Before reduction:
 After reduction:
                   5586
The contributions to the second order correlation energy in atomic units.
IT.
                                                          BJAT
                                                                                   RNORM
    -0.001863
            -0.207844
                                                                                  0.001367
     -0.001866
            -0.004760
                      0.000000
                             -0.046475
                                     -0.025995
                                               -0.001016
                                                        -0.001073
                                                                -0.126819
                                                                         -0.208004
                                                                                  0.000145
     -0.001866
            -0.004761
                             -0.046475
                                      -0.025996
                                                        -0.001073
                                                                -0.126819
                                                                                  0.000017
                      0.000000
                                               -0.001016
                                                                         -0.208006
FINAL CASPT2 RESULT:
   Reference energy:
                     -191.8232886709
   E2 (Non-variational):
                      -0.2080040379
   E2 (Variational):
                      -0.2080040493
   Total energy:
                     -192.0312927202
   Residual norm:
                       0.0000006831
   Reference weight:
                       0.91044
```



Examples

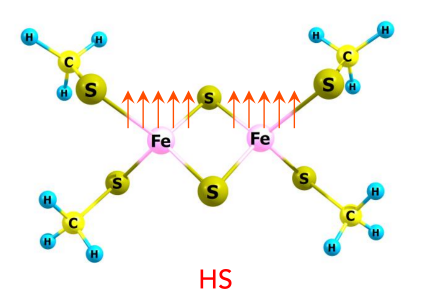
- DMRG calculations with C_nH_{n+2}
- Active space CAS(n,n) n = 4, 6, 8, 10



https://github.com/quanp/polyene



Larger example: [Fe₂S₂(SCH₃)₄]²⁻



C_{2h}
Minimal ANO-RCC basis sets
Low CD
INPORB: RASSCF orbitals

DAVTolerance = 1.0E-5 DMRG = 500 or 2000



Active space of $[Fe_2S_2(SCH_3)_4]^{2-}$

