

To run Orca you need to load it: `module load orca`

to run it (in mylly2) `jsub orca H2O.inp`

There are a lot of example file in `/home/kari/CC2-2022-examples`

To see what is in this dir type `ls -l /home/kari/CC2-2022-examples` (ls is the list command)

you can copy the example files to your own directory: `cp /home/kari/CC2-2022-examples/h2o.inp .`
(there is a dot at the end it is your working directory)

- 1) Compare the Mulliken, Lowdin and CHELPG charges of the $n2c3$ molecule in the lecture notes (Chapter 4). Do a PBE calculation and large basis (TZ or QZ, like cc-pVQZ) of the molecule. The coordinates are in the CC2-2022-examples dir. (you can also look the `n2c3-mol.charge.txt` file for Bader charges and PBE0, MP2 values) . Does these charges agree with each other and to the values in the lectures? Find out what is the CHELPG method. (There is also `n2c3-mol-pbe-qz-H.inp` which computes the Hirshfeld charges.)
- 2) Find how to compute the molecular polarization. Compute polarization of some molecules, like water, methanol, methyl-phenol. What can you say of the polarizability tensor.
- 3) Do a vibrational calculation for a water and methanol molecules. Note that you need to optimize the molecule first. What can you say of the IR intensities. (additional: use `orca_mapspc` tool to make the IR plot, note that the upper limit need to be increased to 4500 cm^{-1} . You can use `gnuplot` to plot the spectra. inside `gnuplot` plot 'h2o-PBE-vib.oout.ir.dat' w l)
- 4) Use the `orca_pltvib` tool to visualize some frequency modes. Use the ase gui to visualize the modes.
- 5) Use the polarization to compute the Raman intensities of water and methanol. NOTE that you need to do frequencies as numerical freq (`numfreq` keyword).

Extra: do the benzene vibration calculation compare that the lectures. (Chapter 4)

Extra: Do calculation of water dimer with PBE. Use `ma-def2-TZVP` basis. You may want to use the van der Waals correction (`D3BJ`). What is the binding energy. (for this you need the water monomer energy). Do the frequency calculations for water molecule and dimer. Compare the energy and Free energy. What is the temperature in these calculations.

ase gui can be started with 'module load python'

The instructions of mylly2 are included.

In the first time make your own directory in /home/kari/CC2-2022-results

mkdir /home/kari/CC2-2022-results/ossi (ossi should be your own name)

At end of exercise copy the results to your result dir: cp *out /home/kari/CC2-2022-results/ossi

Orca input library: <https://sites.google.com/site/orcainputlibrary/home>