**PyConSolv 1.0.6.3**

**User Manual**

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# Acknowledgement and Contributions

**When using PyConSolv, please cite the following publication**:

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**Principal contributors to the current codes**:

Hannah Schmitz (RWTH AACHEN, TU Wien)

Maren Podewitz (TU Wien)

Paul Csepan (TU Wien)

Radu Alexandru Talmazan (TU Wien)

**For more information, please visit:** [**https://github.com/PodewitzLab/PyConSolv**](https://github.com/PodewitzLab/PyConSolv)

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# Introduction

PyConSolv is a python-based software package meant to simplify the setup for molecular dynamics (MD) simulations for transition metal containing catalysts, as well as analyse the resulting trajectories and generate conformers. It interfaces and relies on freely available software packages to execute a state-of-the-art parametrization process.

# Installation

## Software Requirements

To function, PyConSolv requires the following software packages to be installed on your system:

* [ORCA 5.0.x](https://orcaforum.kofo.mpg.de/index.php)1 (Tested with ORCA 5.0.4)
* [AmberTools](https://ambermd.org/AmberTools.php)2 (Tested with version 20-22)
* [Multiwfn](http://sobereva.com/multiwfn/)3 (Tested with version 3.8)

## Python Requirements

To install PyConSolv on your system, Python 3.10 is needed, as well as the following requirements:

* numpy >= 1.23.1
* pandas >= 1.4.3
* matplotlib >= 3.5.3
* rdkit >= 2022.09.1
* parmed >= 4.2.0

## Setting up the python environment

It is recommended to create a virtual environment for PyConSolv, to avoid compatibility problems. Installation is performed as shown in code snipped 1, below:



Code snippet 1 Environment creation.

# Usage

To begin the parametrization, an XYZ (XMol format) structure of your complex should be prepared and located in an empty folder. In case an explicit solvent or a counterion which has not been pre-parametrized is required ([see solvent keyword](#_Keywords_for_the)), an empty folder should be created and an XYZ structure of your desired structure should be placed inside. The recommended folder structure looks as shown in figure 1, below.



Figure 1 initial folder structure example

Once the desired folder structure is complete, PyConSolv can be started on the input structure.

## Command line usage

The PyConSolv parametrization process can be started by calling the pyconsolv script. For the execution of the script an XYZ input file must be provided.

A number of optional parameters can be passed to the function. For details of the available keywords, please see the [keword](#_Keywords) section. While default values are provided, it is recommended to change the QM optimization method, as well as the basis set and dispersion corrections to match your needs. Likewise, it is very important to assign a suitable total charge for your complex.

A critical aspect is the solvent. One of the pre-parametrized solvents can be used or, as mentioned above, any user-provided solvent. A simple example is provided in code snipped 2.



Code snippet 2 Simple example on how to run pyconsolv from the command line.

## Usage within python

The ConfGen module must be imported from the PyConSolv package to make the parametrization functions available. To begin the parametrization process, execute the run function of the PyConSolv object.

A number of parameters can be passed to the run function. For details of the available keywords, please see the [keword](#_Keywords) section. While default values are provided, it is recommended to change the QM optimization method, as well as the basis set and dispersion corrections to match your needs. Likewise, it is very important to assign a suitable total charge for your complex.

A critical aspect is the solvent. One of the pre-parametrized solvents can be used or, as mentioned above, any user-provided solvent. A simple example is provided in code snipped 3.



Code snippet 3 Simple example of parametrization for a neutral compound, while changing settings for the QM method and requesting acetonitrile as a solvent. In this case, the QM method will be DFT with the BP86 functional, def2-SVP basis set, D4 dispersion corrections, using implicit solvation with acetonitrile. The calculation will be run on 8 CPU cores.

The output from PyConSolv will guide the user through the parametrization progress. There are a few steps which must take place for the parametrization, which are explained under [methodology](#_Program_structure).

While the parametrization is running, a more complex folder structure is created, with each critical step being performed in a separate folder, to allow the user to investigate the output of each step, in detail, as well as use the input/output files for other tasks, unrelated to PyConSolv. The final folder structure looks as shown in figure 2. For a detailed explanation of the content of the files, please consult the [methodology](#_Program_structure_and) section.



Figure 2 Folder structure and important files after parametrization is complete

# Methodology

## Program structure and files

PyConsolv follow a workflow that consists of multiple steps:

1. Folder structure setup

During this step, the folder structure required for PyConSolv is created and the input file is checked for compatibility. If required, the input file is re-organized in such a way as to allow for an error-free parametrization, by grouping the ligand structures together. ORCA input files are also created.

1. Geometry optimization and frequency calculation

Structure optimization and a subsequent frequency calculation is performed using ORCA

1. Fragment splitting

The individual fragments are detected and split up in preparation for parametrization using a depth first search algorithm.

1. Fragment charge assignment

A UI window pops up with a Lewis structure representation of each individual substructure, where the user must provide a charge for the ligand/metal.

1. Generation of initial forcefield parameter files

Each ligand is parametrized using antechamber, with the previously provided total charge.

1. RESP charge calculation

Using Multiwfn 3.8, the RESP charges are [calculated](#_Charge_Fitting).

1. MCPB.py execution

MCPB.py input file is created and the script is then executed, in order to determine the metal parameters.

1. Solvent box construction

A solvent box is constructed, using the desired solvent and counterion (if applicable).

1. Equilibration setup

An equilibration [workflow](#_AmberMD) is prepared, along with all the needed input files. Using the GPU accelerated implementation of pmemd (Amber), the system is then equilibrated.

1. Simulation setup

The equilibrated system is copied to the simulation folder and some files required for a final analysis are created. A script is provided for starting the simulation using pmemd (Amber)

1. Analysis

The analysis of a simulation can be performed by calling the pyconsolv script with the -a option and providing the simulation basename as input (e.g. “pyconsolv sim-01 -a”)

## Charge Fitting

Multiwfn 3.81 is used to compute the charges for the parametrizations, as follows.

Nuclear + Electronic Merz-Kollman RESP1 type atomic charge mode is used with 6 points/Å2. All atoms are used for fitting 4 layers, with the scaling factors: 1.4,1.6,1.8,2.0. Automatic radii are utilized for fitting, missing radii are taken from UFF and scaled by 1/1.2. For fitting, the tightness parameter is 0.1, the restraint strengths for the 2 stages are set to 0.0005 and 0.0010, respectively. The maximum number of iterations is 50 and the convergence threshold is set to 10-6. Charge equivalence constraints are enabled for CH2 and CH3 groups. No charge constraint settings are used and the connectivity is determined automatically.

These settings can be changed by the end user, by stopping the parametrization and modifying the multiwfn.input file created in the orca frequency calculation folder, followed by re-running pyconsolv.

## MCPB.py

The MCPB.py module of AmberTools is used in the parametrization process. With the help of it, the parameters for the metal center are created, using the Seminario method, by taking the force constants from the hessian calculated for the system at hand.

The end result is the creation of a force-field modification file and structure files which can be imported into tleap for the solvation of the structure.

## AmberMD

By default, PyConSolv is built with Amber support in mind. Not only relying on the MCPB.py module from AmberTools for parametrization, but also on the Amber MD engine for equilibration and MD simulations.

Once the parametrization step is complete an automatic equilibration procedure is started, as detailed below:

1. The entire simulation box is restrained using cartesian restraints (restraint weight = 1000 kcal mol-1 Å-2), with the exception of the hydrogen atoms, which undergo an energy minimization for 500 cycles.
2. The solute heavy atoms remain restrained and the whole system is minimized for 1000 cycles (restraint weight = 1000 kcal mol-1 Å-2).
3. The system is equilibrated under constant volume to achieve a temperature of 300K, using Langevin dynamics with the collision frequency of 2.0 ps-1, a timestep of 0.001 ps and the SHAKE algorithm for hydrogen bonds.
4. The solute heavy atoms remain restrained (restraint weight = 1000 kcal mol-1 Å-2) and the whole system is equilibrated at constant pressure (1 atm), to achieve a reasonable system density and eliminate the voids created by the automated solvent placement in tleap. This is achieved using Langevin dynamics with the collision frequency of 2.0 ps-1, a timestep of 0.001 ps and the SHAKE algorithm for hydrogen bonds.
5. The system is subsequently cooled under constant volume to achieve a temperature of 100K, using Langevin dynamics with the collision frequency of 2.0 ps-1, a timestep of 0.001 ps and the SHAKE algorithm for hydrogen bonds.
6. A stepwise reduction of the restraints takes place, from 1000 to 0 kcal mol-1 Å-2, minimizing at each step (1000, 500, 200, 100, 50, 20, 10, 5, 4, 3, 2, 1, 0.5 kcal mol-1 Å-2)
7. The system is heated up under constant volume to achieve a temperature of 300K, using Langevin dynamics with the collision frequency of 2.0 ps-1, a timestep of 0.001 ps and the SHAKE algorithm for hydrogen bonds.
8. A final constant pressure (1 atm) equilibration takes place, using Langevin dynamics with the collision frequency of 2.0 ps-1, a timestep of 0.001 ps and the SHAKE algorithm for hydrogen bonds.

The equilibrated simulation box is taken as the starting point for the classical MD simulation. This takes place under constant pressure (1 atm, Berendsen barostat) with isotropic position scaling, a pressure relaxation time of 2.0 ps with Langevin dynamics, SHAKE for the hydrogen atoms and a timestep of 0.001 ps.

## Clustering

Three clustering methods are currently supported: Kmeans, DBscan and Hierarchical. For each of these methods, the user is required to provide some input, such as number of clusters or thresholds for cluster determination. For the clustering, an interface to CPPTRAJ4 is called.

## Solvents

The following solvents are natively supported by PyConSolv: Acetone, Acetonitrile, Ammonia, Benzene, Tetrachloromethane, Dichloromethane, Chloroform, Cyclohexane, DMF, DMSO, Ethanol, Hexane, Methanol, 1-Octanol, Pyridine, THF, Toluene, Water.

These (with the exception of water) have been parametrized for use with PyConSolv, and verified that they approach experimental densities.

# Keywords

## Keywords for PyConSolv

Parametrization options:

* **-c, charge**  : charge of the complete system; type int; default 0
* **-m, method** : ORCA 5 method line; type string; default 'PBE0'
* **-b, basis** : Basis set for ORCA calculations; type string; default 'def2-SVP'
* **-d, dsp** : Dispersion corrections; type string; default 'D4'
* **-p, cpu** : number of CPU cores to be used; type int; default 12
* **-s, solvent** : solvent to be used for MD simulation; type string; default 'Water'.
* **-mult** : multiplicity of the system, default 1
* **-noopt** : perform a single point calculation instead of a geometry optimization
* **-box** : set the solvent box size, default 10
* **-e, --engine** : choose the simulation engine to run the equilibration and simulation with. Options are amber and gromacs, amber is default.
* **-rst, --restraint** : perform a restrained simulation

Analysis options:

* -**a , --analyze** : analyze a simulation
* **-mask, --mask** : atomid mask for clustering
* **-cluster, --cluster** : clustering method
* **-v, --version** : show program's version number and exit
* -**nosp**  : skip single point calculations for clusters
* **-qmmm** : use a qmmm approach to determine cluster energy ranking

The method, basis and dsp keywords conform to the ORCA 5 naming conventions. In theory any input that would be suitable for an orca optimization can be entered here.

Solvents can be either one of the pre-parametrized ones: Water, Acetonitrile, Acetone, Benzene, Cyclohexane, Chloroform, CCl4, CH2Cl2, DMF, DMSO, Ethanol, Hexane, Methanol, Ammonia, Octanol, THF, Toluene or can take the value 'custom'. When the value 'custom' is passed to pyconsolv, a solvent parametrization dialogue is started. Here, the user is required to input the absolute path to the location of the custom solvent XYZ file and subsequently provide information about the permittivity and refraction index of the solvent.

## Keywords for the pyconsolv.restart file

Once PyConSolv is running, a pyconsolv.restart file is created in the folder where your input XYZ file is located. This file contains a keyword, which represents the last successful step of the parametrization. The values taken can be setup, orca, antechamber, frcmod, multiwfn, mcpb, tleap, equilibration or DONE. If you want to restart the parametrization at a specific point, you may edit this file and restart the PyConSolv process.

# Tutorial

## Parametrization and simulation

This is a basic tutorial for running PyConSolv from the command line.

The first step is to call the “pyconsolv” command with the appropriate parameters for an input file in an empty directory e.g.:



PyConSolv will show a greeting message and show the folder where the calculations will take place, as shown in figure 3, below.

Figure 3 PyConSolv greeting message and calculation folder location.

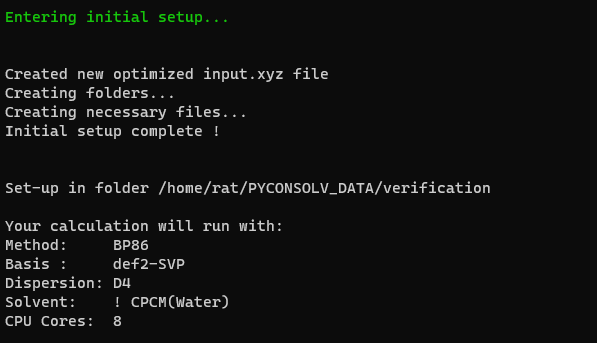
Following the greeting message, more information is provided, regarding the supplied input (see figure 4)

Figure 4 Folder structure is created and the input options are reflected in the output message

Once setup is complete, a geometry optimization is performed on the input structure, followed by a frequency calculation (figure 5).

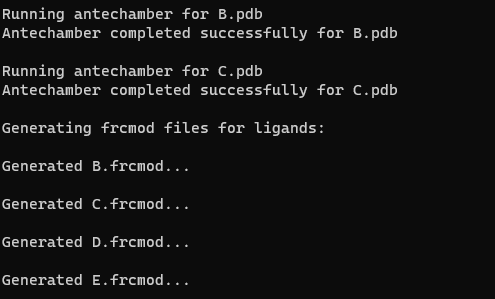


Figure 5 The charges are assigned to each fragment and the program proceeds to the generation of frcmod files.

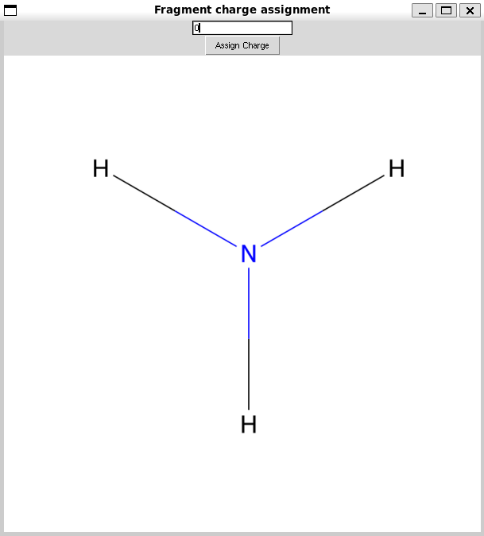


Figure 6 Charge assignment dialogue. The user is asked to input the appropriate charge for each fragment.

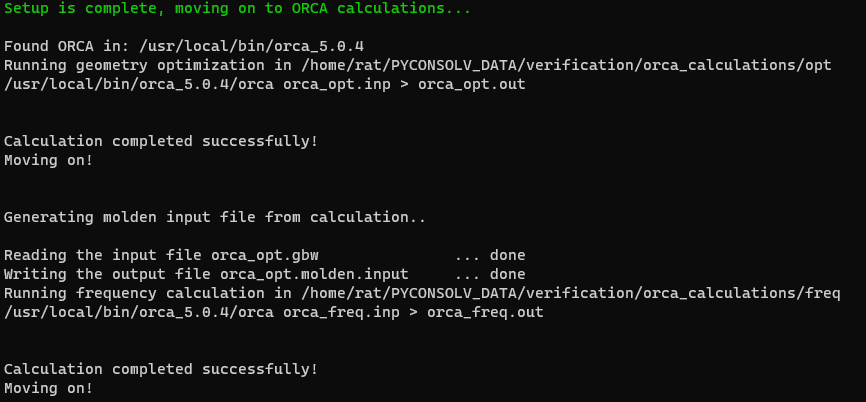


Figure 7 ORCA calculations are performed automatically.

Following the frequency calculation, a pop-up window is displayed to the user, asking for charge assignment for each fragment (figure 6).

Once the fragment charges have been assigned, a charge map file is written in the [MCPB\_setup](#_Program_structure) folder for future reference (figure 7).

Once the charge maps file was written, the force field modification files (frcmod files) are created for each fragment, and the MCPB.py input file is created.

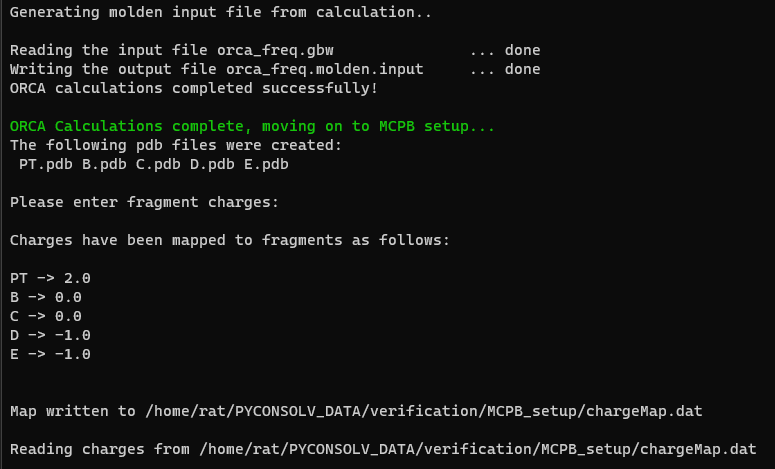


Figure 8 Force field modification files are created.

Once the frcmod files have been created, the RESP charge calculations are started. This is performed with Multiwfn, which requires a molden formatted file as input. This is created automatically from the orca output, followed by a check for ECPs, which have to be marked for correct processing with Multiwfn (figure 9).

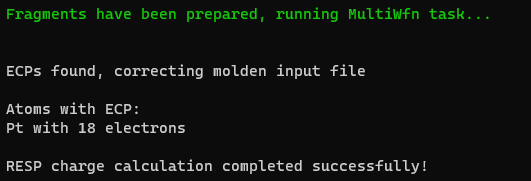


Figure 9 RESP charge calculation, preceded by an ECP presence check in the molden file.

With the RESP charge calculation complete, the MCPB.py parametrization is started and the MCPB.py generated tleap file is corrected for missing bond and correctness, followed by the modification for the appropriate solvent and box size (figure 10). The box size is chosen automatically, such that the closest box edge is 15 Å away from the solute. This is adjusted automatically, in case the box is still too small, during the equilibration.

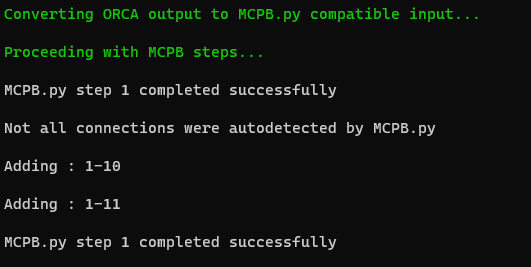


Figure 10 MCPB.py parametrization is performed.

Figure 11 System equilibration is started.

The equilibration is performed and the user is informed of the success at the end of the equilibration step (figure 11). Here the next suggested steps are provided to the user (figure 12)

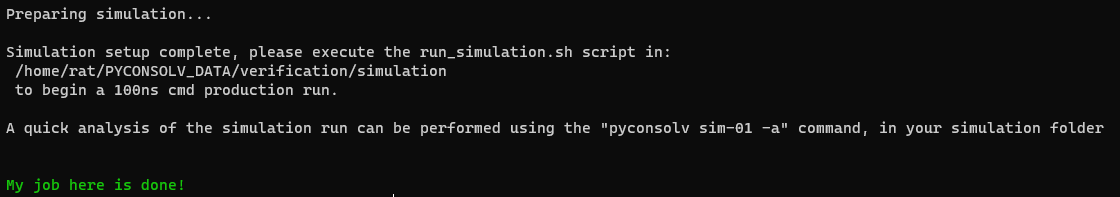


Figure 12 Equilibration is completed and next steps are suggested.

## Analysis and clustering

The clustering module of PyConSolv can be called with the “-a ” option on any simulation. To begin the analysis, the basename of the simulation file must be provided. The default basename for a simulation started with PyConSolv is sim-01. In this case, the analysis can be started with:



The user is the asked to provide atom indexes for aligning the simulation (figure 13), followed by options regarding the clustering method.

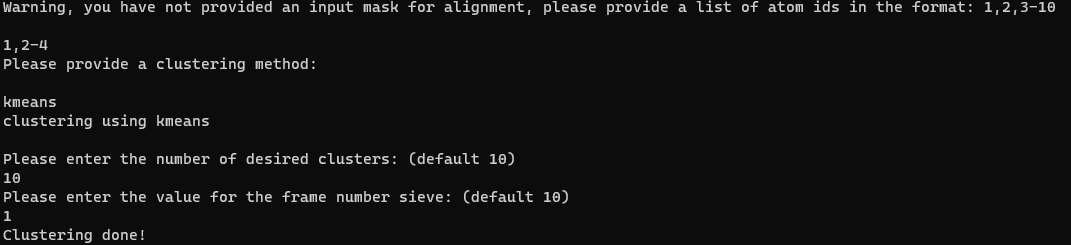
Once the clustering is done, single point energy calculations are automatically performed, at the same level of theory as the one used for parametrization. Finally, the energetic ranking of the conformers is printed out (figure 14), as well as saved in the “cluster\_ranking.dat” output file.

Figure 13 Interactive dialogue for the analysis and clustering of a simulation.

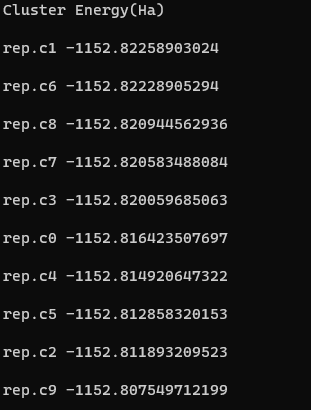


Figure 14 Cluster energy ranking using DFT single point energies.

## Change of force constants

If you want to change force constants for dihedral angles during the process, this needs to be done before running the simulation and after the fragmentation and parametrization is done. This especially useful for double bonds with metals, which might not be correctly identified as such.

1. In the MCPB\_setup folder. First identify the dihedral angle, that needs to change by looking at a visualization of the molecule. The potential energy function (see below) can then be changed in the “LIG\_mcbpy.frcmod” file to the needed value.

The definition of the potential is as follows:

“A-B-C-D 3 2.00 0.00 3.0”,

Where:

* The first term A-B-C-D is the dihedral angle
* The second term 3 is the divider
* The third term (2.00) is Vbarrier in kcal/mol
* The fourth term (0.00) is the phase
* The last term (3.0) is the periodicity

An example for the C3-Y5-M1-Y1 dihedral can be seen below:



Figure 15 Dihedral potential terms in frcmod file

To determine the appropriate values the following equation:

(Please check out the Amber Manual 2023, **15.1.6. Bonded Interaction Terms,** p.278 for more details). A similar procedure can also be done for bonds and angles.

1. Execute the “tleap -f LIG\_tleap.in”
2. Edit the pyconsolv.restart file so that it contains only the “tleap” keyword
3. Restart pyconsolv with the original command

## Clustering with QMMM

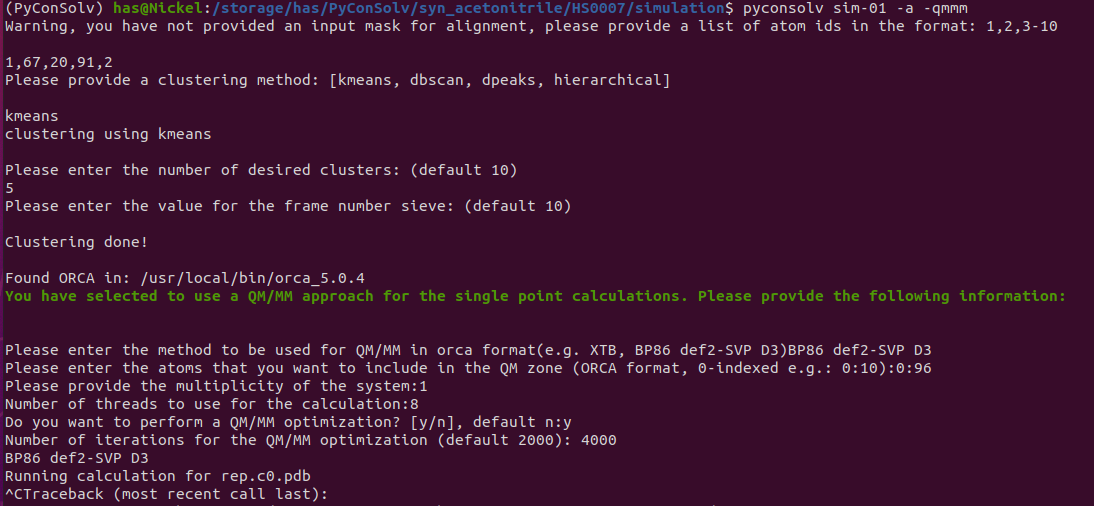
1. Start the analysis by typing pyconsolv “sim-01” –a –qmmm and fill out the questions according to your wishes (Figure 16).

Figure 16 PyConSolv QMMM analysis example.

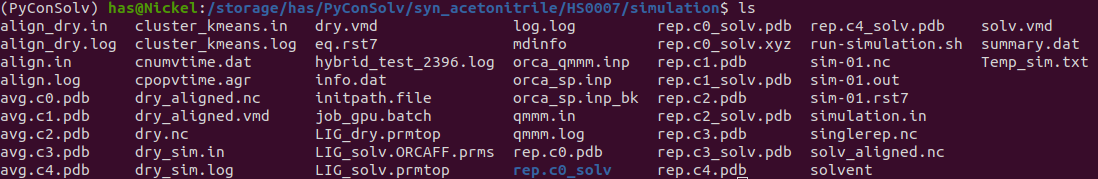
1. The following files can be found after the clustering (Figure 17). In summary.dat the properties of the clusters created can be found. In cnumvtime the frames, which belong to certain clusters over time are plotted. In cpopvtime the change of the population against time is shown.

Figure 17 Files generated after clustering with QMMM.