

# Introduction

## Contents

- 1. **Configuration**
- 2. **Simulation**
- 3. **Applications**
- 4. **Additional Modules**
- [Solvate Suite Homepage](#)

Welcome to the comprehensive ***Solvate Suite User Guide***! The *Solvate Suite* is a modular and integrated toolkit designed to streamline the preparation and processing of molecular systems with explicit solvents, bridging molecular dynamics (MD) simulations with quantum chemistry calculations. This package was conceived to meet the needs of researchers working at the interface between classical mechanics and electronic structure methods, especially in contexts that require a more realistic modeling of solvated environments, such as reaction mechanisms in solution, spectroscopic studies, and thermochemical properties.

With a flexible and highly customizable approach, *Solvate Suite* allows you to:

- Create initial configurations for AIMD/BOMD simulations with *ORCA* or classical MD simulations with *GROMACS*;
- Generate solvation boxes with various solvents and force fields (OPLS-AA, GROMOS, CHARMM, and AMBER) from diverse sources (*GROMACS* database, LigParGen, ATB, or *Q-Force*);
- Integrate data from molecular dynamics simulations with input files for electronic structure calculations (semiempirical, DFT, MP2, etc.);
- Automate repetitive steps such as snapshot extraction, and input preparation for software like *Gaussian*, *ORCA*, *xTB*, and *GROMACS*;
- Perform statistical and structural analyses of solvent environments around target species.

This manual was written to guide users — from beginners to advanced — through the efficient use of *Solvate Suite*'s features. Throughout the chapters, you will find step-by-step instructions, practical examples, useful tips, and conceptual explanations that help you understand not only the “how”, but also the “why” of each step.

Get ready to accelerate your computational workflow and explore complex systems with greater control, reproducibility, and speed.

## Citation

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# 1. Configuration

## SETTING UP MD SIMULATIONS

In the **first stage**, *Solvate Suite* provides optimized protocols for setting up the initial configuration of simulation boxes, enabling AIMD/BOMD simulations with *ORCA* or classical MD simulations with *GROMACS*. The construction of the initial simulation box is handled by the **PACKS** module, through the iterative use of the *PackMol* program. In the case of classical MD simulations, this process includes creating or adjusting force field parameters and building the topology input file. The generation of force field parameters is managed by the **GMDFF** module, which produces parameter files (itp) and applies specific adjustments to the topology file (top) based on various input formats, such as files generated by *Gaussian*, *ORCA*, *xTB*, or *GROMACS*.

### Note

Although the **PACKS** module manages the use of **GMDFF**, enabling the generation of force field parameters during the packing process, it is recommended to run the **GMDFF** module beforehand. Regardless, the **PACKS** module will verify the consistency of the parameters at the end of the packing procedure.

The main program can control the use of the core modules, allowing uniform execution throughout all stages of the suite's use. For example, the suite's first stage involves creating and configuring the simulation box. This is done using the following basic command structure:

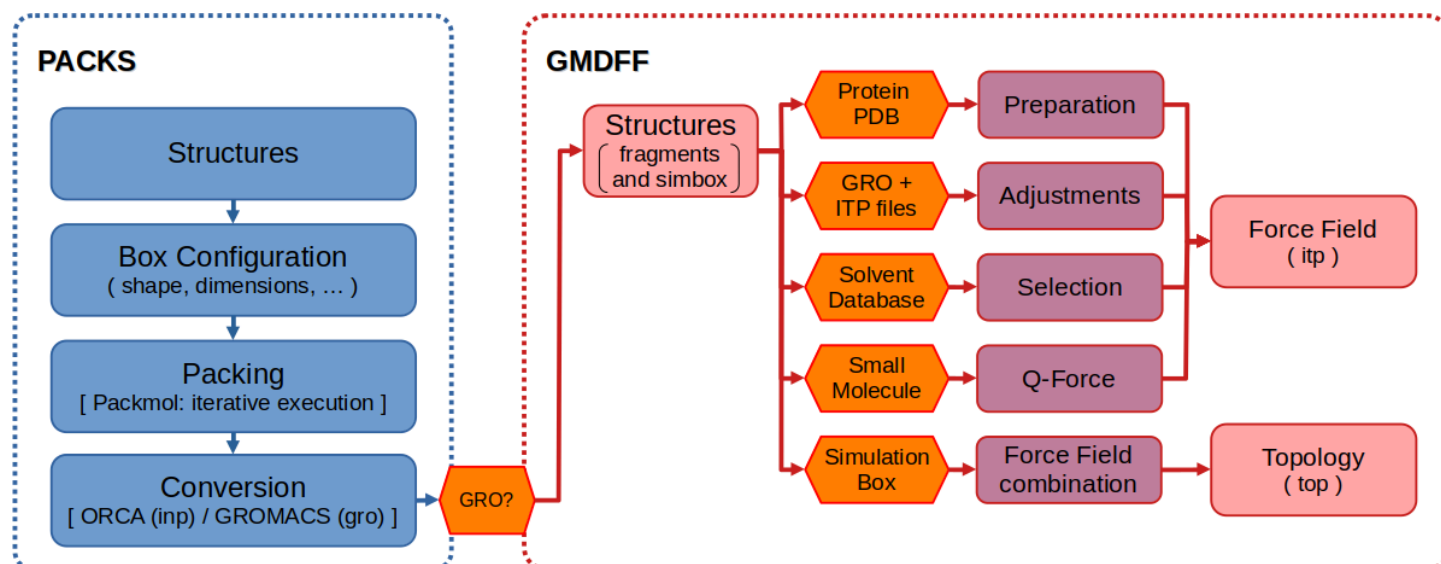
```
solvate <solute.ext> <solvent.ext> [{-options}]
```

where *ext* is one of the extensions recognized by *Solvate Suite* (com/log [*Gaussian*], inp/out [*ORCA*], xyz/xtb [*xTB*], gro/pdb [*GROMACS*]), *inputN* the files with the structures of the solute and solvent molecules, and “-options” a set of options for the configuration of the box to be created.

### **Note**

The list of options can be seen from the applications menu, which is accessible from the command line by typing the module name, without any other parameters.

## STAGE STRUCTURE



## 1.1. PACKS

### 1.1.1. Simulation Box Creation

The **PACKS** module mediates the creation of the simulation box, through the iterative use of the *PackMol* program. The module can be run from the main program or directly from the command:

```
packs <input1.ext> <input2.ext> {...} [<-options>]
```

where *ext* is one of the extensions recognized by *Solvate Suite*, *inputN* the files with the structures of the solute and solvent molecules, and “-options” a set of options for the configuration of the box to be created. The program accepts input and output file formats of *Gaussian* (com, gjf and log), *ORCA* (inp and out), xTB (xyz and xtb), *GROMACS* (gro) and *PDB* (pdb), common formats in Computational Chemistry.

**Note**

Simply type `packs` in the terminal and press *enter* to see the full list of execution options for the module.

The most common use of this module is to create an initial configuration for a system with a single solute molecule surrounded by a number of solvent molecules of a single type:

```
packs <solute.ext> <solvent.ext> -mol N
```

where *N* corresponds to the number of solvent molecules. In the case of ionic solutes, it is possible to include counter-ions with the command:

```
packs <solute.ext> <solvent.ext> -mol N -cnt couterion.ext
```

The program determines the number of counterions needed to neutralize the charge in the simulation box. In these cases, the number *N* of molecules includes the incorporated counterions. The *Solvate* program also makes it possible to create simulation boxes with solvent mixtures, in which case it is necessary to inform the proportion between the molecules:

```
packs <solute.ext> <solvent1.ext> <solvent2.ext> ... -pro [N1:N2:N3...]
```

The initial box can also be created based on the molar concentration *M* of the solute (a situation in which there is more than one solute molecule in the box):

```
packs <solute.ext> <solvent.ext> -cco M
```

or based on molal concentration  $M$ :

```
packs <solute.ext> <solvent.ext> -bco M
```

The box density  $D$  is automatically adjusted based on the density of the solvent used (if it is part of the program's database), or defined manually by the user:

```
packs <solute.ext> <solvent.ext> -mol N -den D
```

The list of parameterized solvents is obtained from the command:

```
packs -lst
```

Any of the parameterized solvents can be provided as input to the program in the simulation box creation step.

Several additional options for adjusting packaging parameters are available. The complete list can be accessed from the help menu of the PACKS module.

## 1.1.2. Execution Options

```
packs <solute> [{<solvent>}] [{<-options>}]
```

```

-----
<solute> = File with solute geometry (.com/log/inp/out/xyz/xtb/gro/pdb)
<solvent> = File with solvent geometry (.com/log/inp/out/xyz/xtb/gro/pdb)
-----
-mol = Number of solvent molecules in the packing step. [Default: 0]
-npk = Number of packing layers. [Default: 2]
-cel = Dimensions of the simulation cell (in nanometers). [Default: Not]
-gap = Gap between solute and cell edges (in nanometers). [Default: Not]
-bco = Molal concentration of the simulation cell (in mol·kg-1). [Default: Not]
-cco = Molar concentration of the simulation cell (in mol·L-1). [Default: Not]
-pro = Proportion between solvent molecules. [N1:N2:...]. [Default: Not]
-den = Density of the simulation cell (in g·mL-1). [Default: 1.0]
-cnt = File with counterion to be added for charge neutrality. [Default: Nor]
-shp = Packaging format (sphere|cube|box). [Default: Sph]
-fmt = Type of generated file (inp|gro). [Default: Aut]
-tol = Minimum distance tolerated in packing (in Å). [Default: Not]
-rep : Resubmit the packing procedure with new parameters. [Default: No]
-for : Run in forced mode (no warning messages). [Default: No]
-nor : Run in normal mode (conventional packing configuration). [Default: No]
-ite : Run in iterative mode (to adjust tolerance). [Default: No]
-inc : Insert external force field into generated topology. [Default: No]
-fft = Force field parameters. [OPL|GRO|CHM|AMB|LPG|QFC|ATB] [Default: OPL]
-bak : Save original input files. [Default: No]
-svt = Solvent to be used in C-PCM/ALPB solvation model. [Default: Aut]
-lst : Lists the set of solvents in the database. [Default: No]
-chk : Checks the consistency of simulation box settings. [Default: No]
-----
(*) Configurable options in automated mode. (†) GROMACS only

```

### 1.1.3. Quickstart Examples

- Example #1: [Creating a box for ORCA with a solute and water molecules](#)
- Example #2: [Creating a box for ORCA with a solute and solvent mixture](#)
- Example #3: [Creating a box for GROMACS with a solute and water molecules](#)
- Example #4: [Creating a box for GROMACS with a solute and a solvent mixture](#)

## 1.2. GMDFF

### 1.2.1. MD Force Fields

The **GMDFF** module mediates the creation of force field parameters (itp files), as well as making special adjustments for the construction of the topology (top file). The simplest usage consists of the

following command:

```
gmdff (input.ext)
```

where *ext* is one of the extensions recognized by the *Solvate Suite* (com/log [*Gaussian*], inp/out [*ORCA*], xyz/xtb [*xTB*], gro/pdb [*GROMACS*]).

**Note**

Simply type `gmdff` in the terminal and press *enter* to see the full list of execution options for the module.

If the simulation box is prepared for the *GROMACS* program (which is determined by the extension of the input files) the program automatically runs the GMDFF module, which prepares the force field files as well as the system topology file. This means that, in the general case, it is possible to create the simulation box and its topology file from a single command line. In these cases, however, it is recommended to run the GMDFF module first, in order to prepare all the necessary force field files required for executing the PACKS module. This is done using the following command structure:

```
gmdff (input.ext) [{-options}]
```

where *ext* is one of the extensions recognized by Solvate Suite (com/log [*Gaussian*], inp/out [*ORCA*], xyz/xtb [*xTB*], gro/pdb [*GROMACS*]), *input* is the file with the structure of the solute or solvent molecule, and “-options” a set of options for the configuration of the force field to be created.

There are 4 scenarios in which this module can be used:

- In the case of biomolecule PDB files, the program performs structure preparation (removal of heteroatoms and addition of hydrogens, without adjusting protonation based on pH, which may require additional preparations to be carried out by the user), followed by conversion to the *GROMACS* format (gro file) and creation of the force field parameters in a single input (itp file).
- In the case of force fields obtained by external servers (such as LigParGen and ATB) the module removes any residual charges, a necessary step for building simulation boxes that are, in fact, electrically neutral. This is done by homogeneously compensating for any residual charge between all atoms in the structure.

- In the case of internally parameterized solvent molecules, the program selects the appropriate parameters and writes them to a force field input (itp file), as well as converting the input file to the appropriate format (gro file).
- In the case of small molecules, the module sequentially executes the *Q-Force* program to generate parameters from a quantum chemical calculation.

**Note**

In the case of using the *Q-Force* program, the user can generate a default settings file with the `-chk` option. The edited file can then be read by the program with the `-cfg` option. Alternatively, the user can provide his own settings file with the `-cfg <file.ini>` command.

For *Q-Force*, it is possible to check whether the options are appropriate to start the force field creation:

```
gmdff <input.ext> -chk
```

In this case, a configuration file (ini) is generated with a default set of execution options, which can be edited by the user to apply custom configurations. This file can then be read using the following command:

```
gmdff <input.ext> -cfg
```

Alternatively, the user can provide a custom settings file via the following command:

```
gmdff <input.ext> -cfg config.ini
```

Several force field parameters (OPLS/AA, GROMOS, CHARMM and AMBER) for a wide range of solvents are part of the program database, coming from several sources (GROMACS repository, LigParGen, ATB and *Q-Force*). The complete list of parameters can be displayed from the command:

```
gmdff -lst
```



It is possible to display only the parameters of a specific force field, such as OPLS/AA:

```
gmdff -lst opls
```

Once you have identified the desired parameters, you can obtain them from the command:

```
gmdff {solvent_forcefield}
```

For example, the TIP3P parameters in the OPLS/AA force field are obtained from:

```
gmdff tip3p_oplsaa.wtr
```

Finally, if the input file matches that of the simulation box, the program combines the force field files into a single and independent topology file. This results in only two files for starting the simulation: the simulation box (gro file) and the topology (top file, with all the force field parameters).

## 1.2.2. Execution Options

```
gmdff {molecule.ext} [{-options}]
```

```

-----
.ext = Input file extension (.com/log/inp/out/xyz/xtb/pdb/gro)
+ -----
  • Q-Force Options
  -chr = Molecular charge. [Default: 0]
  -mul = Molecular multiplicity. [Default: 1]
  -pcm = Solvent for C-PCM model. [Default: None]
  -chk : Verify generated configuration file before submit. [Default: No]
  -cfg = Execute with user defined input configurations. [Default: No]
  -prc = Number of processors to be used. [Default: 4]
  -mem = Amount of memory in GB to be used. [Default: 8]

  • Additional Options
  -fld = Force field parameters. [OPL|GRO|CHM|AMB|LPG|QFC|ATB] [Default: OPL]
  -wtr = Water model (conditional). [SPC|SPCE||TIP3P|TIP4P|TIP5P] [Default: SPC]
  -lst : Provides the complete list of parameterized solvents. [Default: No]
  -for : Execute in forced mode. [Default: No]
-----
Total CPUs: 12 (max. MPI-procs.) :: System Memory: 5 GB
-----
(†) Specifies the model to use if the input file corresponds to a water molecule.
(*) Repositories: https://traken.chem.yale.edu/ligpargen
                  https://atb.uq.edu.au
                  https://cgenff.com

```

### 1.2.3. Quickstart Examples

- Example #1: [Preparing a protein PDB file](#)
- Example #2: [Creating the Q-Force force field parameters](#)
- Example #3: [Handling force field parameters from external servers](#)
- Example #4: [Handling force field parameters from internal database](#)

## 2. Simulation

### EXECUTING MD SIMULATIONS

In the **second stage**, the *Solvate Suite* provides a group of modules to manage the MD simulations. From the **MDRUN** module, the simulations with the *ORCA* and *GROMACS* programs are executed using the same set of command lines. In the case of *ORCA*, the simulation execution options are set directly in the input (inp file) at run time on the command line. In the case of *GROMACS*, the options are adjusted and saved in its own input settings (mdp file). In the latter case, there are more

simulation execution options, such as the type of barostat and thermostat, time constant for pressure and temperature coupling, isothermal compressibility, and saving frequency, among others. In any case, the simulation properties can be extracted and analyzed from the **DATAS** module, which has resources for statistical and graphical analysis.

#### **Note**

The modular design allows users to run each step of the workflow separately, enabling detailed control and progress evaluation at any point in the simulation.

The second stage of using the suite consists of executing and analyzing the simulation. The execution can be done using the following command structure:

```
solvate <simbox.ext> -mdrun [{-options}]
```

where *ext* is one of the extensions recognized by *Solvate Suite* (inp [*ORCA*], or gro [*GROMACS*]), *simbox* is the simulation box file in its initial configuration, and “-options” a set of options for the simulation control. After each step performed (minimization, equilibration, and production), the data can be analyzed using the following command structure:

```
solvate <simbox.ext> -datas [{-options}]
```

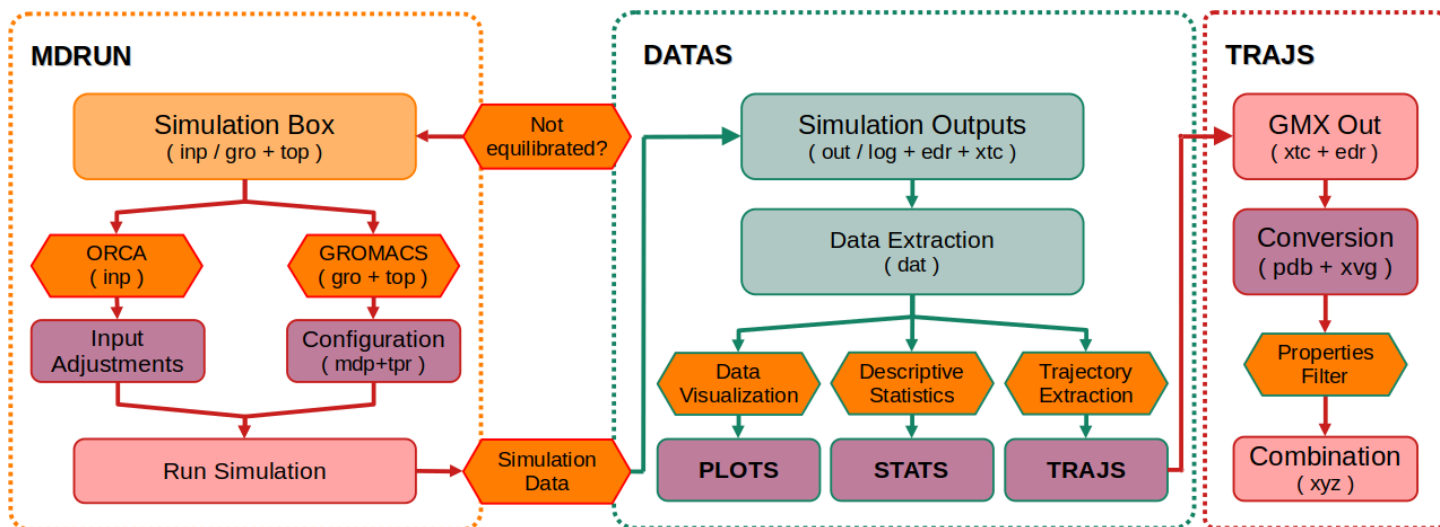
where “-options” is a set of options for the simulation data analysis.

Several options for adjusting simulation parameters, as well as analyzing properties and extracting data, are available for the *ORCA* and *GROMACS* programs. The complete list can be accessed in the help menu of the MDRUN and DATAS modules.

#### **Note**

The list of options can be seen from the applications menu, which is accessible from the command line by typing the module name, without any other parameters.

## **STAGE STRUCTURE**



## 2.1. MDRUN

### 2.1.1. MD Execution

The **MDRUN** module manages the execution of simulations, through a unified interface for the *ORCA* and *GROMACS* programs. The module can be run from the main program or directly from the command:

```
mdrun <simbox.ext> [{-options}]
```

where *ext* is one of the extensions recognized by *Solvate Suite* (**inp** [*ORCA*] or **gro** [*GROMACS*]), and “-options” a set of the configurations for the simulation execution.

#### **Note**

Simply type **mdrun** in the terminal and press *enter* to see the full list of execution options for the module.

### **Note**

In the case of using the *GROMACS* program, the user can generate a default settings file with the `-chk` option. The edited file can then be read by the program with the `-cfg` option. Alternatively, the user can provide his own settings file with the `-cfg <file.mdp>` command.

For *GROMACS*, it is possible to check whether the options automatically configured after packaging are appropriate to start the simulation:

```
mdrun (simbox.gro) -chk
```

In this case, a configuration file (mdp) is generated with a default set of execution options, which can be edited by the user to apply custom configurations. This file can then be read using the following command:

```
mdrun (simbox.gro) -cfg
```

Alternatively, the user can provide a custom settings file via the following command:

```
mdrun (simbox.gro) -cfg config.mdp
```

Before starting the simulation, it is necessary to carry out an initialization step for the initial box, which includes energy minimization and box optimization. For example, to initialize the simulation box at a pressure of 1.0 bar and a temperature of 298.15 K, you can use the following command:

```
mdrun (simbox.gro) -ini -prs 1.0 -tmp 298.15
```

After initialization, temperature equilibration can be performed in an NVT ensemble. For example, running 1.0 ns for equilibration in the NVT ensemble can be executed with the following command:

```
mdrun (simbox.gro) -nvt -tmp 298.15 -tot 1.0
```

and the pressure equilibration, in an NPT ensemble, with:

```
mdrun (simbox.gro) -npt -prs 1.0 -tmp 298.15 -tot 1.0
```

The equilibration steps involve saving the trajectory every 100 steps (in a trr file) to conserve disk space.

#### **Note**

To assess the equilibrium, the properties acquired during the simulation can be analyzed using the **DATAS** module.

After achieving equilibration, the production step can proceed. In this step, the trajectory is saved at each step of the simulation (in an xtc file). Production is run with the following command:

```
mdrun (simbox.gro) -prd -prs 1.0 -tmp 298.15 -tot 5.0
```

#### **Note**

The trajectory extraction procedure can be performed using the **DATAS** module, which has several extraction filters available.

Finally, after extraction, it is possible to remove several files that are unnecessary for the final step of the microsolvation treatment, which can be done with the command:

```
mdrun (simbox.gro) -cls
```

## 2.1.2. Execution Options

```
mdrun (file.ext) [{-options}]
```

## ORCA/GROMACS

auto : Execute the MD simulation with automatic settings.

» Description

ORC: [Default: -tot = 50.0 ps (prod.) | -dts = 1.0 fs]

GMX: [Default: -tot = 2.50 ns (prod.) | -dts = 2.0 fs]

+ -----

» Automation Options:

-auto : Start MD simulation with automatic settings.

-init : Init. energy minimization with simulation box optimization procedure.

-equi : Exec. MD equilibration procedure (with auto restart).

-prod : Exec. MD production procedure (with auto restart).

-bomd : Run a BOMD simulation at GFN2-xTB level.

» Procedure Options:

-min : Execute an energy minimization of the simulation box.

-opt : Execute an optimization of the simulation box.

-nvt : Run an equilibration simulation on NVT ensemble.

-npt : Run an equilibration simulation on NPT ensemble.

-prd : Run a production simulation on NVT/NPT ensemble.

» Settings Options:

-met = MD simulation method (GFF|GF1|GF2). [Default: GFF]

-prs = Pressure for the MD simulation (in bar). [Default: 1.0]

-tcp = Time constant for pressure coupling (in ps). [Default: 0.5]

-bar = Type of barostat pressure coupling. [Default: Ber]

\* BRDS = Berendsen | CRES = C-rescale | PARR = Parrinello | MTTK

-tmp = Temperature for MD simulation (in K). [Default: 298]

-tct = Time constant for temperature coupling (in ps). [Default: 0.5]

-ter = Type of thermostat temperature coupling. [Default: V-r]

\* BRDS = Berendsen | VRES = V-rescale | NOSE = Nose-Hoover | ADSN = Ande

-kap = Isothermal compressibility value (in bar<sup>-1</sup>). [Default: 1.0]

-nst = Simulation steps to update neighbor list. [Default: 100]

-dmp = Number of steps for dump positions. [Default: 1]

-dts = Simulation time step (in fs). [Default: 1.0]

-tot = Simulation additional (ORCA) or total (GMX) time (in ps). [Default: Aut]

-nrs = Number of simulation run steps. [Default: Aut]

» Additional Options:

-prc = Number of parallel CPU processes to run. [Default: 4]

-chk : Check generated files before submit. [Default: No]

-cfg = Execute with user defined input configurations. [Default: No]

-fix : Performs MD without atom constraints. [Default: No]

-nhr : Exec. with no H-bonds. restraints (for production steps). [Default: No]

-clr : Cleaning unnecessary files (after extraction). [Default: No]

-cpt : Compact output files. [Default: No]

-ext : Extract output files. [Default: No]

-for : Execute in forced mode (suppressing upper to 2 warnings). [Default: No]

(\*) Configurable options in automated mode. (●) ORCA available. (■) GROMACS available

## 2.1.3. Quickstart Examples

- Example #1: [Submitting a simulation with ORCA](#)
- Example #2: [Submitting a simulation with GROMACS](#)

## 2.2. DATAS

### 2.2.1. MD Analysis

The **DATAS** module performs graphical and statistical analysis of simulated properties, through a unified interface for the *ORCA* and *GROMACS* programs. The module can be run from the main program or directly from the command:

```
datas (simbox.ext) [{-options}]
```

where *ext* is one of the extensions recognized by *Solvate Suite* (out [*ORCA*], or log [*GROMACS*]), and “-options” a set of the configurations for the simulation execution.

#### **Note**

Simply type **datas** in the terminal and press *enter* to see the full list of execution options for the module.

To assess the equilibrium, the properties acquired during the simulation can be analyzed with the following basic command structure:

```
datas (simbox.ext)
```

Properties such as pressure, temperature, density, and potential energy are evaluated graphically and statistically (descriptive statistics include the calculation of standard error, standard deviation and drift of properties; RMSD and RDF analysis are also available).

After a production step, simulation data extraction can be performed with trajectory truncation. Several options are available, including the ability to perform RMSD and RDF analyses of the



truncated trajectory. The simplest procedure is to run the command:

```
datas <simbox.ext> -ext
```

It is possible to apply a set of filters to the extraction of configurations. For example, it is possible to extract configurations whose pressure is around the target pressure  $P$ , within a narrow window of  $\pm D$ :

```
datas <simbox.ext> -ext -ptr [ $P:D$ ]
```

## 2.2.2. DATAS Options

```
datas <file.ext> [-dat [-opt]][-plt [-opt]] [-trj [-opt]]
```

```

-----
auto : Execute an automated properties extraction from simulation files.
      » Description
      Properties extraction + statistical and graphical analysis
      » NOTE: this option is recommended in queue systems.
      + -----
-dat : Generate a dat file only. [for GMX: -min|-opt|-nvt|-npt]
-cpt : Generate a dat file in compact format.
-dot : Convert dots to commas.
-psa : Performs statistical analysis.
-for : Force new property data extraction.
      + -----
-rms : RMSD analysis for SLT molecular residue (full trajectory).
-rdf : RDF analysis for center of masses of ref SLT and sel SVT residues.
      = [I:J] : RDF analysis for ref SLT/I and sel SVT/J labels.
-pdf : PDF analysis for center of masses of ref SLT and sel SVT residues.
      = [I:J] : PDF analysis for ref SLT/I and sel SVT/J indexes.
-sma : Calculate simple moving average at frequency 5 to the extracted RDF data.
-tim = Final simulation interval (in ps) to extract RDF data (def.: 100 ps).
-cnv = Cumulative numerical value to be highlighted in the RDF/PDF analysis.
-shw : Visualize the performed SMA analysis.
      + -----
-plt : Generate graphs for params vs simulation time.
-stp : Generate graphs for params vs simulation step.
      + -----
-trj : Analyze the trajectory file.
-ext : Extract a truncated trajectory file.
-ctr = [I:J] : Skip I equilibration steps and extract J production steps.
-btr = [B:X] : Select frames from B trajectory blocks into X% final steps.
-ptr = [P:D] : Select configurations around pressure P with tolerance  $\pm D$  (in bar).
-prd : Select configuration production steps only and the corresponding properties.
-grp : Extract short-ranged non-bonded potential energies.
-nbx : Do not make corrections for "fragmented" molecules near the box edges.
-----
(*) Configurable options in automated mode. (●) ORCA available. (■) GROMACS available.
(†) RDF = Radial Distribution Function. (from TRJ.XTC file)
(‡) PDF = Pair Distribution Function. (from TRY.XYZ file)

```

## 2.2.3. MD Statistics

The **STATS** module performs statistical analysis of simulated properties, through a unified interface for the *ORCA* and *GROMACS* programs. It can be used from the command:

```
stats {simbox.ext} [{-options}]
```

where *ext* is one of the extensions recognized by *Solvate Suite* (out [*ORCA*], or log [*GROMACS*]), and “-options” a set of the configurations for the simulation execution.

#### **Note**

Simply type `stats` in the terminal and press *enter* to see the full list of execution options for the module.

## 2.2.4. STATS Options

```
stats <datafile.dat> [{-options}]
```

```
-----  
-cpt : Read a generated dat file in compact format.  
-for : Force new property data extraction.  
-----
```

## 2.2.5. MD Graphics

The `PLOTS` module performs graphical analysis of simulated properties, through a unified interface for the *ORCA* and *GROMACS* programs. It can be used from the command:

```
plots <simbox.ext> [{-options}]
```

where *ext* is one of the extensions recognized by *Solvate Suite* (out [*ORCA*], or log [*GROMACS*]), and “-options” a set of the configurations for the simulation execution.

#### **Note**

Simply type `plots` in the terminal and press *enter* to see the full list of execution options for the module.

## 2.2.6. PLOTS Options

```
plots <inputs.ext> [{-options}]
```

```
-----  
Formatted plots:
```

```
+ -----  
-tim : Plot graphs for params vs simulation time.  
-stp : Plot graphs for params vs simulation step.  
-cnv = Cumulative value to be highlighted in RDF analysis.  
+ -----
```

```
General plots:
```

```
+ -----  
-csv : Load an unformatted general plot.  
-plt : Plot y versus x as lines (default).  
-sct : A scatter plot of y vs x with varying marker size/color.  
-bar : Make a bar plot.  
-stm : Create a stem plot.  
+ -----  
-sma : Simple moving average to the extracted RDF data.  
+ -----
```

```
Saving plots:
```

```
+ -----  
-bak : Save generated graphs (without iterative interface).  
-----
```

```
(*) ext = dat, rms, rdf, csv, nmr
```

## 2.2.7. MD Trajectories

The **TRAJS** module performs the extraction of simulated trajectory from *GROMACS* program. It can be used from the command:

```
trajs <simbox.gro> [{-options}]
```

### Note

Simply type **trajs** in the terminal and press *enter* to see the full list of execution options for the module.

## 2.2.8. TRAJ Options

trajs (trajectory.gro/xtc) [{-options}]

```
-----
auto : Execute an automated trajectory extraction from simulation files.
      » Description
      btrj = 1000 blocks with 10% extraction
      » NOTE: this option is recommended in queue systems.
      + #1: Individual Options - (no data analysis) -----
-itr = I      : Extracts position structure #I from the trajectory.
-ntr = N      : Extract N equally spaced MD steps for truncated trajectory.
-str = M      : Extract equally spaced frames at every M steps from trajectory.
      + #2: Combinable Options - (with data analysis) -----
-ctr = [I:J]  : Skip I equilibration steps and extract J production steps.
-btr = [B:X]  : Select frames from B trajectory blocks into X% final steps.
-ptr = [P:D]  : Select configurations around pressure P with tolerance ±D (in bar)
      + #3: Output file formats - (GROMACS only) -----
-xyz : Output in XYZ file format. [default]
-pdb : Output in PDB file format.
-gro : Output in GRO file format.
      + #4: Other Options -----
-prd : Select configuration production steps only and the corresponding properties.
-grp : Extract short-ranged non-bonded potential energies.
-nbx : Don't make corrections for "fragmented" molecules near the box edges.
-bak : Save intermediate PDB and XVG files for alternative conversions.
-deb : Save intermediate XVG file and create a reconstructed ENT file.
-for : Force new extraction and override previous PDB file.
-----
(*) Configurable options in automated mode. (●) ORCA available. (■) GROMACS available.
(†) Structure position, not configuration step.
```

## 2.2.9. Quickstart Examples

- Example #1: [Data analysis from ORCA simulation](#)
- Example #2: [Data analysis from GROMACS simulation](#)

# 3. Applications

## PERFORMING APPLICATIONS

In the **third stage**, following trajectory extraction, the truncated trajectory file *simbox.trj.xyz* is generated. The **MACRO** (under development) and **MICRO** modules can be used for the treatment of biomolecules or for microsolvation analysis, respectively.

## PERFORMING THE MICROSOLVATION TREATMENT

In the first step of the microsolvation treatment, the module selects  $N$  low-energy clusters, each composed of  $M$  solvent molecules. The value of  $M$  may be: (1) determined from a prior radial distribution function (RDF) analysis using the DATAS module; (2) defined based on the solute-to-solvent ratio at a user-specified concentration; or (3) calculated automatically, using a cutoff distance corresponding to the diameter of a solvation sphere. The extraction is performed using the following command structure:

```
solvate {simbox.trj.xyz} -micro [{-options}]
```

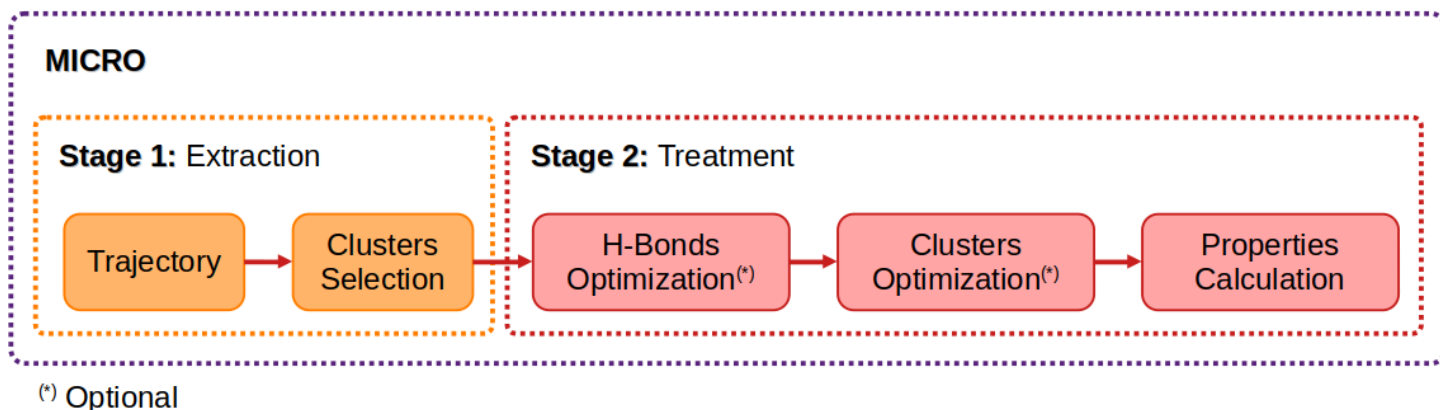
After this extraction, the selected clusters are stored in the file *solute.trj.xyz*. In the second step, this file serves as input to optimize the hydrogen bonds within the microsolvation clusters. Subsequently, all microsolvation clusters can be optimized (with constraints, at the xTB level). Finally, the desired properties can be calculated based on user-specified configurations. This procedure is carried out using the following command structure:

```
solvate {solute.trj.xyz} -micro [{-options}]
```

### Note

Various processing options for cluster treatment are available. The complete list can be accessed via the help menu of the MICRO module by executing the module name without any arguments on the command line. These procedures can be fully automated using the “-auto” option.

## STAGE STRUCTURE



## 3.1. MACRO

Under development...

## 3.2. MICRO

### 3.2.1. Clusters Extraction

The **MICRO** module manages the extraction of microsolvation clusters. It is possible to select  $N$  low-energy clusters, each containing  $M$  solvent molecules. The value of  $M$  can be determined from a previous RDF analysis (using the DATAS module), defined based on the solute-solvent ratio (at a user-specified concentration), or calculated automatically (based on a cutoff distance corresponding to the diameter of the sphere surrounding the solvent molecule). The extraction is performed using the following command:

```
micro {simbox.trj.xyz} -str  $N$  -nsv  $M$ 
```

The extraction based on molal concentration  $C$  is performed using the command:

```
micro {simbox.trj.xyz} -str  $N$  -bco  $C$ 
```

The extraction based on the number  $S$  of solvation shells (estimated from the solvent radius) is performed using the command:

```
micro {simbox.trj.xyz} -str N -sol S
```

**Note**

Simply type `micro` in the terminal and press *enter* to see the full list of execution options for the module.

### 3.2.2. Clusters Treatment

The `MICRO` module can also perform property calculations based on the microsolvation cluster treatment. This includes the optimization of hydrogen bonds within the clusters, constrained optimization of the entire clusters at the xTB level, and subsequent property evaluation with configurations set by the user. These calculations are executed using the following command:

```
micro {solute.trj.xyz} -opt -hbd -key "keywords"
```

**Note**

Simply type `micro` in the terminal and press *enter* to see the full list of execution options for the module.

### 3.2.3. Execution Options

```
micro {trajectory.trj.xyz} [{-options}]
```



#### MICROSOLVATION CLUSTERS EXTRACTION

```
auto : Execute an automated microsolvation clusterization from simulation files
  » Description
    bCon = 1 mol·kg-1 (for setting the number of solvent molecules)
    Nstr = 1% of trajectory structures (1000 min, 10000 max)
    cTRJ = All production steps
  + -----
-str = Number of clustered structures selected trajectory.      [Default: 100]
-nsv = Number of solvent molecules in the clustering process.   [Default: 0]
-sol = Number of solvent layers in the clustering process.      [Default: 1]
-bco = Molal conc. of the microsolvation cluster (in mol·kg-1). [Default: 0.6]
-cfg = N° of MD equilibration steps [:N° of MD production steps]. [Default: 0:6]
-chr = Value of microsolvated solute charge.                   [Default: 0]
-mlt = Value of microsolvated solute multiplicity.             [Default: 1]
-prs = Value of simulation pressure (in bar).                   [Default: 1.6]
-tmp = Value of simulation temperature (in K).                  [Default: 298]
-svt = Solvent to be used in ALPB solvation model.             [Default: Aut]
-rtn : Retain the original orientation of the clusters.         [Default: No]
-psv : Preserve the original trajectory file.                   [Default: No]
-for : Execute in forced mode.                                  [Default: No]
  + -----
-prc = Number of parallel CPU processes to run.                [Default: 8]
-mem = Amount of memory to be used (in GB).                    [Default: 16]
```

micro (solute.trj.xyz) [{-options}]

#### MICROSOLVATION CLUSTERS TREATMENT

```
-opt : Optimize microsolvation cluster structures (FCA procedure). [Default: No]
-hbd : Optimize hydrogen bonding in the microsolvation cavity.     [Default: No]
-key = "List of keywords" for final calculation (with ORCA).       [Default: M06]
-for : Execute in forced mode.                                      [Default: No]
  + -----
-prc = Number of parallel CPU processes to run.                    [Default: 8]
-mem = Amount of memory to be used (in GB).                        [Default: 16]
```

(\*) Configurable options in automated mode.

### 3.2.4. Quickstart Examples

- Example #1: [Extracting clusters from a simulation with ORCA](#)

- Example #2: [Extracting clusters from a simulation with GROMACS](#)

## 4. Additional Modules

### 4.1. File Management

#### 4.1.1. OUTIN

- Manipulation of input/output files

```
outin <file_inp.ext> <file_out.ext> [{-options}]
```

---

## FILE FORMATS

---

.ext: com : Gaussian (read/write)[gjf]  
inp : ORCA (read/write)  
xyz : xTB (read/write)  
gro : GROMACS (read/write)  
pdb : PDB (read/write)  
log : Gaussian (read)  
out : ORCA (read)  
xtb : xTB (read)  
xtc : GROMACS (read)  
trj : XYZ/GRO/PDB (read)  
+ -----  
cpt : Data Comp (write)[Compress calculated data]  
dat : Calc Data (write)[Extract calculated data (option: nogeom)]  
sup : Supl Data (write)[Extract supplementary data]  
sep : Sepa Data (write)[Separate input/output data]  
spl : Splt Data (write)[Split input/output data]  
div : Divd Data (write)[Divide input/output data]  
job : Jobs Exec (write)[Divide input/output jobs]  
frq : Calc Frqs (write)[Extract vibrational frequency data]  
ifr = Imag Dist (write)[Imaginary mode distortion (factor of distortion)]  
+ -----  
slt : SLTs Extn (write)[Solute extraction from trajectory frames]  
svt : SVTs Extn (write)[Solvent extraction from trajectory frames]  
pcf = PChg File (write)[Solvent as point charges][ = Scale factor for aton]

---

---

## CONVERTION OPTIONS

---

-key = "List of keywords" (for Gaussian or ORCA).  
-add = "List of keywords/additional" (for Gaussian).  
-svt = Include solvent type for implicit (with ALPB or PCM solvation method).  
-nmr : Include configuration for NMR calculation (for ORCA).  
+ -----  
-itr = Extract structure n° N of a trajectory.  
-ntr = Number N of equally spaced MD steps for truncated trajectory.  
-str = Extract equally spaced frames at every M steps from trajectory.  
-ctr = Skipe I equilibration and extract J production MD steps (auto mode).  
-btr = Select frames from I trajectory sections into J% final steps.  
-ptr = Select frames aroud pressure P with tolerance  $\pm D$  (in bar).  
-nsm = Number of solvent molecules in the clustering process.  
-bmc = Molal conc. of the microsolvation cluster (in mol·kg<sup>-1</sup>).  
-grp : Extract short-ranged non-bonded potential energies.  
-nbx : Don't make corrections for "fragmented" molecules near the box edges.  
+ -----  
-prs = Include pressure configuration for thermodynamic calculation.  
-tmp = Include temperature configuration for thermodynamic calculation.  
-fcc = Final concentration after compression (in mol·L<sup>-1</sup>).  
-frq = Scale factor for frequencies (for zero-point energy correction).  
-qha = Quasi-Harmonic Approximation (with optional cutoff frequency).

-----  
(\*) I,J = Integers; N,P,D = Real. (○) xTB available. (●) ORCA available. (■) GROMACS

## 4.1.2. FILES

- Input/output file Management

```
-----  
files <solute+solvent_molN.ext> [<-options>]  
  
PARAMETERS  
    input.ext [Output LOG/OUT/XTB file]  
  
OPTIONS  
-cpt          [Compact the intermediate files] [Default: No] [-cp]  
-ext          [Extract the intermediate files] [Default: No] [-ex]  
-----
```

## 4.1.3. SOLVS

- List of available solvents and ions for packaging

```
-----  
solvs [<solvent>] [<-option>]  
  
OPTIONS/DEFAULTS  
-field T      [Force field parameters]  
    ► ALL = All force fields  
    ● OPL = OPLS-AA/L all-atom force field  
    ○ LPG = OPLS-AA/L (from LigParGen)  
    ○ QFC = OPLS-AA/L (from Q-Force)  
    ● GRO = GROMOS96 54a7 force field  
    ○ ATB = GROMOS96 54a7 (from ATB)  
    ● CHM = CHARMM36 all-atom force field  
    ● AMB = AMBER-GS force field  
-----
```

## 4.1.4. MDPAR

- List of available configuration files for MD simulations

```

-----
mdpar [<-step N>]

OPTIONS/DEFAULTS
  -step N    [Step for extracting MDP parameters]
             ► 0/ALL = List all MDP files
             ○ 1/OPT = Box Optimization
             ○ 2/MIN = Energy Minimization
             ● 3/NVT = NVT Equilibration
             ● 4/NPT = NPT Equilibration
             ● 5/PRD = NPT Production
-----

```

## 4.2. Biomolecular Treatment

### 4.2.1. PDBIO

- PDB processor for Biomolecular systems

```

-----
pdbio <input.pdb> [<-options>]

PARAMETERS
  input.pdb [Input PDB file]
  output.gro/itp [Output GRO/ITP file]

OPTIONS/DEFAULTS
  -ffp T    [Force field parameters] [Default: OPLS]
            * GRO = GROMOS96 54a7 force field
            * OPL = OPLS-AA/L all-atom force field

ADDITIONAL
  -pdb      [Archives PDB input]
  -gro      [Generate GRO geometry file]
  -itp      [Produces ITP forcefield file]
  -ion      [Don't do ITP files of ionic species]
  -lap      [Override PDB input file]
  -cls      [Wipe all PDB comment lines]
  -bak      [Create a backup of auxiliary files]
-----

```

## 4.2.2. PDB2X

- PDB cleaner module

```
-----  
pdb2x <input.pdb> [<-options>]  
  
PARAMETERS  
    input.pdb [Input PDB file]  
    output.ent [Output ENT file]  
-----
```

## 4.2.3. IONIC

- PDB ions forcefield constructor

```
-----  
ionic <input.pdb> [<-options>]  
  
PARAMETERS  
    input.pdb [Input PDB file]  
    output.gro/itp [Output GRO/ITP file]  
-----
```

## 4.2.4. GROUT

- GRO files processor module

```
-----  
grout <input.gro/itp> [<-options>]  
  
PARAMETERS  
    input.gro/itp [Input GRO/ITP file]  
    output.gro/itp [Output GRO/ITP file]  
  
OPTIONS  
    -gro          [Archives GRO input/output]  
    -itp          [Produces a concatenated ITP forcefield file]  
    -pdb          [Converts a PDB file to GRO input geometry]  
-----
```

## 4.2.5. GRO2X

- GRO cleaner module

```
-----  
gro2x <input.gro/itp> [<-options>]  
  
PARAMETERS  
  input.gro/itp [Input GRO/ITP file]  
  output.gro/itp [Output GRO/ITP file]  
-----
```

## 4.2.6. GROTC

- GRO total charge calculator

```
-----  
grotc <input.gro>  
  
PARAMETERS  
  input.gro      [Input GRO file]  
-----
```

---

## 4.3. Microsolvation Treatment

### 4.3.1. CONFS

- Conformational analysis automator

```

-----
confs <input.ext> [<-options>]

PARAMETERS
    input.ext [Input COM/LOG/INP/OUT/XYZ/XTB/GRO/PDB file]

OPTIONS
    -xff      [Use GFN-FF in conformational analysis (default)]      [-xf]
    -xtb      [Use GFN2-xTB in conformational analysis (slower)]     [-xt]
    -key T    [Keywords for final optimization («xTB Opt Freq»)]     [-kw]
    -svt T    [Include implicit solvation configuration]              [-sv]
    -prs N    [Pressure for thermochemistry calculation (in atm)]    [-ps]
    -tmp N    [Temperature for thermochemistry calculation (in K)]   [-tp]
    -con N    [Final concentration after compression (in mol·L-1)]   [-fc]
    -min N    [minimum Boltzmann weight for free energy averaging]   [-mp]
    -prc P    [Number of processors to be used]                       [-pc]
    -mem M    [Amount of memory to use in GB]                        [-mm]
    -deb      [Check generated inp file before submit]                [-db]
    -for      [Execute in forced mode]                                [-fm]

OUTPUT FORMATS
    -inp      [Output file in ORCA format (default)]                  [-in]
    -gro      [Output file in GROMACS format]                          [-gr]
    -pdb      [Output file in PDB format (for LigParGen)]             [-pd]
    -xyz      [Output file in XYZ format (for LigParGen)]             [-xy]
    -sep      [Split the output file into its conformers]             [-sp]
-----

NOTE: This is an interface to CREST program.

```

## 4.3.2. BONDS

- Hydrogen bonds analyzer



```

-----
bonds <input.ext> [<-options>]

PARAMETERS
    input.ext [Input COM/LOG/INP/OUT/XYZ/XTB/GRO/PDB file]

OPTIONS
    -xff          [Use GFN-FF in conformational analysis (default)]      [-xf]
    -xtb          [Use GFN2-xtb in conformational analysis (slower)]      [-xt]
    -key T        [Keywords for final optimization («xtb Opt Freq»)]      [-kw]
    -svt T        [Include explicit solvent and PCM configuration]         [-sv]
    -prs N        [Pressure for thermochemistry calculation (in atm)]      [-ps]
    -tmp N        [Temperature for thermochemistry calculation (in K)]     [-tp]
    -con N        [Final concentration after compression (in mol·L-1)]      [-fc]
    -min N        [minimum Boltzmann weight for free energy averaging]     [-mp]
    -prc P        [Number of processors to be used]                       [-pc]
    -mem M        [Amount of memory to use in GB]                         [-mm]
    -deb          [Check generated inp file before submit]                 [-db]
    -for          [Execute in forced mode]                                 [-fm]

PROCEDURES
    -nci N        [Use CREST/NCI procedure with N svt mol]                [-nc]
    -hbd N        [Use CREST/HBD procedure with N svt mol] (default)      [-hb]
-----
NOTE: This is an interface to CREST and ORCA programs.

```

### 4.3.3. MSRUN

- Microssolvation Aggregator

```

-----
msrun <simbox.trj.xyz> [<-options>](a)
msrun <solute.trj.xyz> [<-options>](b)

PARAMETERS
<simbox.ext> [Input simbox TRJ.XYZ file]
<solute.ext> [Input solute TRJ.XYZ file]

PROCEDURES
-auto          [Execute an automated microsolvation optimization]
-gcc C         [GCALC module configuration. ( Ex.: -gc "-ini -opt -max 100 -fca" )]
-que C         [QUEUE module configuration. ( Ex.: -qc "-key "#ONIOM(M062X/6-311+G(d

OPTIONS
-prc N         [Number of parallel CPU processes to run] [Default:
-mem N         [Amount of memmory to be used (in GB)] [Default:
-----
(a) From extracted tractory file (with TRajs). [Complete procedure by default]
(b) From extracted clusters file (with MICRO). [Partial procedure, without MICRO]
(*) Each module can be run independently.
(†) The "-key" option can be passed separately.

```

## 4.3.4. GCALC

- Gibbs Free Energy Calculator

```

-----
gcalc <solute.trj.xyz> [<-options>]

PARAMETERS
<solute.ext> [Input solute TRJ.XYZ file]

PROCEDURES
-auto          [Execute an automated microsolvation optimization]
                » NOTE: this option is recommended in queue systems.
                » HINT: -auto          [starts processing 100 clusters]
                  : -auto N          [starts processing N clusters]
                  : -auto +M         [add M clusters to the previous analysis]

STEPS CONTROL
• Stage #1: Clusters Characterization
-ini           [Execute clusters characterization and selection]          [Default:
                Characterization for structures selection.
                Procedure performed at xTB-GFF level.

• Stage #2: Clusters Treatment
-opt           [Optimization and free energy calculation] (FSA|FCA|FCO)    [Default:
                Calculation after structures selection
                Procedure performed at xTB-GFN2 level combined with DFT

» FSA: [F]ull [S]olutes [A]ccommodated geometry (with SPH).1 «xTB level»
» FCA: [F]ull [C]luster [A]ccommodated geometry (with SPH).1 «xTB level»
» FCO: [F]ull [C]luster [O]ptimized geometry (with CHM).2 «xTB level»

ADDITIONAL OPTIONS
• Selection Options
-max M         [Maximum number of structures into energy window]          [Default:
-add M         [Add M clusters to GFE procedure from INI step.             [Default:

• System Options
-chr N         [Value of microsolvated solute charge]                    [Default:
-mlt N         [Value of microsolvated solute multiplicity]              [Default:
-prs P         [Pressure for thermochemistry calculation (in atm)]        [Default:
-tmp T         [Temperature for thermochemistry calculation (in K)]       [Default:

• Solvation Options
-svt S         [Solvent type (for ALPB/CPCM solvation method)]           [Default:
-nis           [No implicit solvation]                                    [Default:

• Configuration Options
-prc N         [Number of parallel CPU processes to run]                 [Default:
-mem M         [Amount of memmory to be used (in GB)]                   [Default:

• Additional Options
-key T         ["List of keywords" (for ONIOM program)]                  [Default:
-fcc N         [Final concentration after compression]                  [Default:
-hbd           [Include H-bonds at extrapolation (for ORCA final step).   [Default:
-nmr           [Config. for NMR calculation (for ORCA final step).       [Default:
-for           [Force reexecution stages.                                [Default:

```

```

-fil      [Filter high energy clusters (above 100.0 kcal·mol-1).      [Default:
+ -----
-dat      [Extract data for selected stage.                        [Default:
-plt      [Plot extracted data.                                    [Default:
-bak      [Save generated graphs (without iterative interface).    [Default:
+ -----
-exc [i,..][List of structural clusters to exclude. (inp) [i,j,k,...] [Default:
-clr [i,..][List of optimized clusters to clean.      (out) [i,j,k,...] [Default:
+ -----
-del N    [Delete specified previous optimization]                [Default:
-ext N    [Extract specified previous calculation]                [Default:
-ren T    [Rename specified previous description] "DSC" / [OLD:NEW] [Default:
-chk      [Check all previous calculations] (= spec. to extract)   [Default:
-log      [Summary of performed calculations]                    [Default:
+ -----
-out T    [Force reading specified ONIOM output file]              [Default:
          [Force reading default ONIOM output file]                [Default:
+ -----
-bnd      [Execute in BONDS mode]                                  [Default:
-----
(*) Configurable options in automated mode.
(▪) Try "-sN" to view stage N only.
(†) The "ith" cluster identified from trj.xyz input file.
(‡) The "ith" cluster identified through graphical analysis.
(1) SPH = Single Point Hessian.
(2) CHM = Conventional Hessiam Method.

```

## 4.3.5. QUEUE

- QUEUE System Manager

```

-----
queue [<-options>]

OPTIONS
-inf          [Show current ONIOM calculation configuration]
-log          [Show current ONIOM calculation results]
-run          [Perform the calculation procedure]
              + -----
-key T        [Complete list of keywords for ONIOM calculation configuration]
-chr N        [Charge value of the microsolvation cluster]
-mlt N        [Multiplicity value of the microsolvation cluster]
-des T        [Optional output configuration description]
-fcc N        [Final concentration after compression]                [Default:
-con          [Force concatenation of results over warnings]
-clr          [Clean up run folder after archiving results]
              + -----
-srm          [Execute calculation in a serial mode]
-for          [Execute calculation in forced mode]
              + -----
-bnd          [Execution for BONDS module]                            [Default:
-gcc          [Execution for GCALC module]                            [Default:
              + -----
-prc N        [Number of parallel CPU processes to run]                [Default:
-mem N        [Amount of memmory to be used (in GB)]                  [Default:
-----

```

## 4.3.6. ONIOM

- ONIOM Program

```

-----
oniom <input.gjf> [<-options>]

PARAMETERS
<input.gjf> [File with GaussView format for Solvate/Microsolvation calculation]

OPTIONS
-opt          [Execute geometry optimization in forced mode]                [Default:
-ext          [Execute ONIOM extrapolation in forced mode]                [Default:
              + -----
-srm          [Execute calculation in a serial mode]                        [Default:
-for          [Execute optimization and extrapolation in forced mode]      [Default:
-deb          [No extrapolation execution: only tests]                    [Default:
              + -----
-prc N        [Number of parallel CPU processes to run]                    [Default:
-mem N        [Amount of memmory to be used (in GB)]                      [Default:
-----

```

## 4.4. Multiple Purposes

### 4.4.1. CHECK

- Monitoring of procedures progress

```
-----  
check <input.ext> [<compt.ext>] [<-options>]  
  
PARAMETERS  
    input.ext [Input file]  
    compt.ext [Compacted file]  
              [Extensions: COM/INP/XYZ/LOG/OUT/XTB/TMP]  
  
OPTIONS  
    -ext      [Extract compacted internal file]  
              (ex.: check infile.zip cptfile.cpt -ext)  
    -del      [Deletes compacted internal file][zip only]  
              (ex.: check infile.zip cptfile.cpt -del)  
    -ren      [Renames compacted internal file][zip only]  
              (ex.: check infile.zip oldfile.cpt -ren newfile.cpt)  
-----
```

### 4.4.2. CLEAN

- Cleaner for temporary files

### 4.4.3. COMPT

- Compression of output files

```
-----  
compt <output.log/out/xtb> [-chk]  
  
PARAMETERS  
    output.ext [Output LOG/OUT/XTB file]  
-----
```

#### 4.4.4. DEPLY

- Full deployment Python dependencies check

#### 4.4.5. IMAGS

- Imaginary mode distortion

```
-----  
imgs <output.log/out/xtb> [-img fator]  
  
PARAMETERS  
  output.ext [Output LOG/OUT/XTB file]  
-----
```

#### 4.4.6. INFOS

- Displaying system information

#### 4.4.7. MOLAP

- Structural alignment for RMSD minimization

```
-----  
molap <reference.ext> <structure.ext> [<trajectory.trj.xyz>] [<-options>]  
  
PARAMETERS  
  files.ext [Input  COM/LOG/INP/OUT/XYZ/XTB/GRO/PDB file]  
  
OPTIONS  
  -bak      [Save modified "structure" file with minimized RMSD (at system or  
  -ref      [Save modified "structure" file with minimized RMSD (at "referenc  
  -str      [Save modified "structure" file with minimized RMSD (at "structur  
  -slt      [Reorientation relative to the central solute fragment]  
-----
```

#### 4.4.8. RMSDS

- Calculation of structural RMSD and solute energy

```

-----
rmsds <reference.ext> <structure.ext> [<preoptimization.ext>] [<trajectory.trj.x
PARAMETERS
    input.ext [Output COM/LOG/INP/OUT/XYZ/XTB/GRO/PDB file]

OPTIONS
-min          [Maximum structural superposition for RMSD minimization]
-bak          [Save RMSD information in the input "structure" file]
-for          [Force new calculation and override previous output]
-----

```

### 4.4.9. SCALC

- Chemical shifts calculator

```

-----
scalc <input.trj.xyz> [<-options>]

PARAMETERS
    input.trj.xyz [Output TRJ file]

OPTIONS
    -ato A      [Atomic symbol of the species to be analyzed] [-at]
    -ref N      [Reference value for chemical shifts (in ppm)] [-rf]
    -eqv [L]    [List of equivalent atoms. (note: -eqv=[A1=A2:B1=B2=B3...])] [-eq
+-----+
    -out F      [Force reading specified ONIOM output file] [-of
    -out        [Force reading default ONIOM output file] [-of
+-----+
    -hbd        [Use BONDS data] [-hb
    -gcc        [Use GCALC data] [-gc
    -plt        [Plot the NMR spectrum] [-pt
-----
* Note : Requires "gcc" file, and optionally "pcm" and "exp" files.

```

### 4.4.10. SOLVX

- Solvent identifier in clusters



```

-----
solvx <file.trj.xyz>

PARAMETERS
  file.trj.xyz [Output TJR.XYZ file]
-----

```

## 4.4.11. **STAND**

- Modify input to standard orientation

```

-----
stand <input.ext> &

PARAMETERS
  input.ext [Input GJF/COM/INP/XYZ/GRO]
-----

```

## 4.4.12. **STATX**

- Full statistical analysis directly from GMX program

```

-----
statx <output.log>

PARAMETERS
  output.log [Output LOG file]
-----

```

## 4.4.13. **SUBST**

- Modification of texts in editable files

```

-----
subst <input.ext> "text1" "text2"

PARAMETERS
  input.ext [Extensions: all]
-----

```

## 4.4.14. SUMMY

- Extraction of properties from output files

```
-----  
summy <output.log/out/xtb>  
  
PARAMETERS  
    output.ext  [Output LOG/OUT/XTB file]  
  
OPTIONS/DEFAULTS  
    -prs N      [Pressure configuration for thermodynamic calculation]  
    -tmp N      [Temperature configuration for thermodynamic calculation]  
    -fcc N      [Final concentration after compression (in mol·L-1)]  
    -frq N      [Scale factor for frequencies (for zero-point energy correction)]  
    -qha <N>    [Quasi-Harmonic Approximation (with optional cutoff frequency)]  
-----
```

## 4.4.15. SUPPY

- Extraction of supplementary data from output files

```
-----  
suppy <output.log/out/xtb>  
  
PARAMETERS  
    output.ext  [Output LOG/OUT/XTB file]  
  
OPTIONS/DEFAULTS  
    -prs N      [Pressure configuration for thermodynamic calculation]  
    -tmp N      [Temperature configuration for thermodynamic calculation]  
    -fcc N      [Final concentration after compression (in mol·L-1)]  
    -frq N      [Scale factor for frequencies (for zero-point energy correction)]  
    -qha <N>    [Quasi-Harmonic Approximation (with optional cutoff frequency)]  
-----
```

## 4.4.16. SVIEW

- Visualization of input/output files

```

-----
svview <input.ext> [<-options>] &

PARAMETERS
    input.ext  [Input file] (*)

GENERAL OPTIONS
    -try I      [Generate a try file with N truncated steps (.trj file)]
    -vmd        [Force file open with VMD program]
    -cpt        [Perform compression of input/output files for GVW program]

JMOL OPTIONS
    -noc        [Run without input conversion]

VMD OPTIONS
    -cpk        [Show solute with VDW representation]
    -vdw        [Show solute with VDW representation]
    -lico       [Activate Licorice for solvent shell]
    -dyna       [Activate DynamicBonds for solvent shell]
    -surf       [Activate QuickSurf for solvent shell]
    -text       [Execute in a text mode]
-----
(*) Conversions: COM/GJF/LOG/INP/OUT/XYZ/XTB/PDB/GRO/TRJ

```

## 4.5. Queuing Systems

### 4.5.1. SUBMIT

- Run the program in the system queue

```
-----
submit <program> <input.ext> <-options> <-ntasks I>

PARAMETERS
  program    [gsn/orca/xtb/gmx program]
  input.ext  [Input GJF/COM/INP/XYZ/GRO file]

OPTIONS/DEFAULTS
  -options   [Program specific options]
  -queue T   [Name of available queue]
  -ntasks I  [Number of processors]      [-prc]
  -qmemor I  [Amount of memory (in GB)][-mem]
  -preserve  [Preserve the submission job file]
  -debug     [Check generated job before submit]

AVAILABLE SYSTEM BATCH
  »Local Machine

WARNING
  This is an open-source script that may require
  adjustments depending on the available queue
  system
-----
```

## 4.6. Program Interfaces

### 4.6.1. runORCA

- *ORCA* program interface

### 4.6.2. runXTB

- *xTB* program interface

### 4.6.3. runCREST

- *CREST* program interface

#### 4.6.4. **runGMX**

- *GROMACS* program interface

#### 4.6.5. **runBABEL**

- *oBabel* program interface

#### 4.6.6. **runJMOL**

- *JMol* program interface

#### 4.6.7. **runVMD**

- *VMD* program interface

#### 4.6.8. **runPACKMOL**

- *PackMol* program interface

#### 4.6.9. **runQFORCE**

- *Q-Force* program interface