



# **Scripting Manual**

## ***Amsterdam Modeling Suite 2023.1***

**[www.scm.com](http://www.scm.com)**

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## GETTING STARTED

To conveniently use the scripting tools of the Amsterdam Modeling Suite you need to set some shell [environment variables](#) and add the `AMSBIN` folder to your `PATH`. This can be done by sourcing file `amsbashrc.sh`, which is located in the Amsterdam Modeling Suite installation directory.

### 1.1 Linux

Note: if you followed the [Linux Quickstart installation Guide](#), the `amsbashrc.sh` should be automatically sourced when you start up a new terminal, and you can ignore the following steps.

- Start up a terminal
- Source the `amsbashrc.sh` with the following command (note: you should replace `path_to_installation_directory` with the actual path to your AMS installation directory:

```
. path_to_installation_directory/amsbashrc.sh
```

- To test that you properly sourced the `amsbashrc.sh` file you can type the following command, which should yield the help message for the [amsprep](#) (page 3) command line tool:

```
amsprep -h
```

### 1.2 MacOS

- From the AMSjobs GUI module, click on the **Help** dropdown menu and select **Terminal**. This will open a new terminal with all necessary environment variables already set and the `AMSBIN` folder already added to the `PATH`.

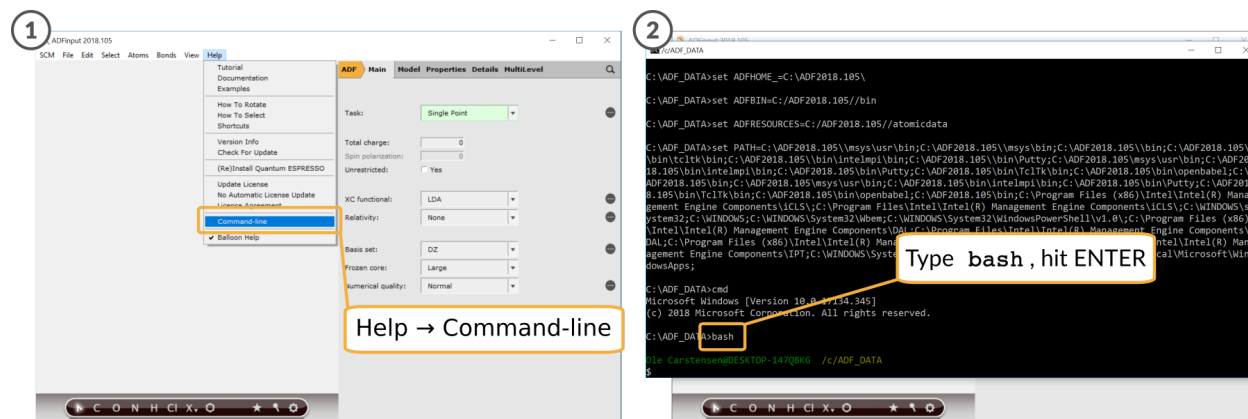
Alternatively, you can follow the Linux steps.

### 1.3 Windows

Every Windows installation of AMS2019 and newer, as well as older ADF versions, come with a pre-configured command line. The easiest way to access the command line is via the Help menu of the graphical user interface:

Go to **Help** → **Command-line**

Inside the command line window, type `bash` and hit ENTER (alternative: type `sh`)



The advantage of calling the command line from the GUI is that you will find yourself in the current working directory right away. In situations in which the GUI is not available, it is also possible to use the pre-configured command line directly:

Double click the file `ams_command_line.bat` in your AMS installation directory (e.g. `C:/AMS2021.101`)

## 1.4 Notes on Python

Python scripts should be executed using the python3 interpreter shipped with Amsterdam Modeling Suite:

```
$AMSBIN/amspython scriptname.py
```

Further information can be found here: *Python Stack in Amsterdam Modeling Suite* (page 21).

## COMMAND LINE TOOLS

- **amsprep**: prepare an ADF job from a script (or command line).
- **amsreport**: get information (including images) from an ADF result file (for use in your script, or to generate an HTML or tab-separated report).
- **pkf**, **cpkf**, **dmpkf**, **udmpkf**: the KF utilities, which are command-line utilities to process KF files.

## 2.1 AMSprep: generate (multiple) ADF jobs

AMSprep allows one to generate input files for the different programs of the Amsterdam Modeling Suite by means of console commands. As such AMSprep can be used to run the same type of calculation on a series of different chemical systems. Another important example are automatic checks of the convergence of the results with respect to the computational parameters e.g. by varying input settings such as basis set choice or numerical integration accuracy while recomputing the same system.

AMSprepare (\$AMSBIN/amsprep) generates a job script from a template .ams file. Such a template file can either be produced by AMSinput or simply be found among the default templates included. These default templates are identical to those present in AMSinput.

Two **examples** are presented here to demonstrate the capabilities of AMSprep:

- In [BakersetSP](#) you will see how to use amsprep to run a particular job for a test set of molecules. The individual molecular structures are provided as xyz-files which contain no ADF specific information. AMSreport is used to collect the values of the bonding energies resulting from these calculations.
- In [ConvergenceTestCH4](#) you will see how to use AMSprep to test convergence of the bonding energy with respect to the basis set and the numerical integration grid.

The options of AMSprep are listed when running the module without further command line arguments, or with the -h flag:

```
% amsprep -h
AMSprepare (amsprep) generates a job script from a .ams file (the template),
with user specified changes to input options / method / system.

Usage: amsprep -t template.ams [-m molecule.(ams|adf|xyz|mol|t21)] [-z charge] [-s-
↪spin]

                                [-runtype SinglePoint|GeometryOptimization|Frequencies]
                                [-gradientonly]
                                [-q quality] [-zlmfit quality] [-kspace quality]
                                [-lattice v1.x v1.y v1.z ...]
                                [-i integration] [-b basis] [-c core] [-r relativity]
                                [-basiscacheid id]
```

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```

[-x xcpotential] [-e xcenergy] [-bondsonly]
[-dftbmodel DFTB|SCC-DFTB|DFTB3] [-dftbparameters dir]
[-dftbdispersion None|Default|D3-BJ|D2|ULG|UFF]
[-logfile logfile] [-j jobname] [-a amsfile]
[-dist "atom1 atom2 distance ..."]
[-angle "atom1 atom2 atom3 angle ..."]
[-dihed "atom1 atom2 atom3 atom4 angle ..."]
[-atomtype "atom type ..."]
[-structure "atom structure ..."]
[-pointcharges file]
[-efield "Ex Ey Ez"]
[-rxforcefield fname] [-rxniter n] [-rxnrstep n] [-
→rxtstep T]
[-rxmethod method] [-rxmdtemp T] [-rxmdpres p]
[-region "name at1 at2... "]
[-fragments prefix] [-onejob]
[-g "key value"]

```

Start with a job template, adjust it for this particular job, and write the resulting  
→job  
to standard output. Values specified should match exactly the values as you would  
→specify  
using AMSinput, also for menu choices.

#### TEMPLATE

-t: the .ams file (saved by AMSinput) to be used as template, defining the whole job  
All other options override values from this job

Instead of a .ams file, you may also specify the name of one of the standard  
→templates  
as defined in AMSinput: "Single Point", Frequencies, "Geometry Optimization", etc  
A special option for energy and gradients  
for the current geometry: EG (see also -gradientsonly)

Some shortcuts: SP, EG, GO, FREQ, optionally prefixed by  
→(ADF|BAND|DFTB|UFF|MOPAC)-  
For example: ADF-FREQ, BAND-SP, DFTB-GO, MOPAC-EG

Some ReaxFF shortcuts: REAXFF-EG for a single ReaxFF iteration

#### CHANGES TO TEMPLATE

-m: the molecule to use, element types and coordinates  
This can be taken from anything that AMSinput can import,  
for example .ams, .mol, xyz or .t21 files

The -m flag may be repeated, each molecule added will be in its own region  
This may be used for fragment calculations, but it does not work with .ams files

If you specify an .sdf file, you can select which frames to import:

conformers.sdf#1-10	loop over the first 10 frames
conformers.sdf#e2.0	loop over all frames with energy below 2.0 (units as in the file, wrt the lowest energy of all frames in the file, energies from comment lines)
conformers.sdf#1-10e2.0	loop over the first 10 frames, and use only those with energy below 2.0
conformers.sdf	use the first frame of the sdf file

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If you specify a .t21 file, you can select which frames or range of frames to

```

→import:
    ajob.t21#ircf3           3rd frame in the IRC forward path
    ajob.t21#ircb2         2nd frame in the IRC backward path
    ajob.t21#h7            7th frame in the history
    ajob.t21#lt8           8th frame in the LT path
    ajob.t21#ircf3-10      IRCForward frame 3, 4, ... 10
    ajob.t21#ircf          IRCForward all frames, starting at 1
    ajob.t21#ircf0-       IRCForward all frames, starting at 0
                          (original geometry, before first step)

```

If you specify a .cry file, the compound to import may be specified:

```

$AMSHOME/atomicdata/Molecules/Crystals/Cubic/CsCl.cry#MgTl

```

When looping, all resulting jobs will be joined together, the jobname and ams

```

→files
    get the frame sequence number appended after an _
    When looping only one -m flag may be specified

```

-xyz: use xyz coordinates from specified file, not touching anything else  
it is applied after -t and -m  
the elements and number of atoms should match  
currently works with KF and xyz files

-smiles: use smiles to describe the molecule

-irc: when using IRC frames in the -m flag, revert the backwards order

-dist: change the distance between atom1 and atom2 to the specified distance  
the arguments must be enclosed in quotes, and may be repeated for multiple

```

→distances

```

-angle: change the angle (atom1, atom2, atom3) to the specified angle  
the arguments must be enclosed in quotes, and may be repeated for multiple

```

→angles

```

-dihed: change the dihedral (atom1, atom2, atom3, atom4) to the specified angle  
the arguments must be enclosed in quotes, and may be repeated for multiple

```

→angles

```

-atomtype: set the type (element) of atom to type  
the arguments must be enclosed in quotes, and may be repeated for multiple

```

→types

```

-structure: add a structure just as if using the structure tool in AMSinput  
atom is the selected atom, structure is the name of the structure file  
the arguments must be enclosed in quotes, and may be repeated for multiple

```

→changes

```

-liststructures: list available structure files for use with -structure, and exit

-runtype: run type (SinglePoint,GeometryOptimization,Frequencies)

-gradientonly: after calculating the gradients, stop  
works also for excited state gradients if requested in your template

-z: charge (real number)

-s: spin (integer), if not zero this implies an unrestricted calculation

-q: quality (Basic, Normal, Good, VeryGood or Excellent), default for Becke/ZlmFit

-i: integration (integer)

-i: Becke integration (Basic, Normal, Good, VeryGood or Excellent)

-i: teVelde integration (integer)

-zlmfit: ZlmFit quality (Basic, Normal, Good, VeryGood or Excellent)

-kspace: KSpace quality (GammaOnly, Basic, Normal, Good, VeryGood or Excellent)

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```

-lattice: lattice vectors first three numbers for the first vector, next for the
↳second etc
    The dimension follows from the number of vectors
-b: basis type (SZ, DZ, DZP, TZ, TCP, TZ2P, QZ4P)
-c: core type (None, Small, Medium, Large)
-basiccacheid id: refer to t21 files from previous runs prefixed with this id
-r: relativistic level (None, Scalar, Spin-Orbit), using ZORA
-x: XC potential during SCF, one from the options available in AMSinput:
    LDA,
    GGA:BP, GGA:BLYP, GGA:PW91, GGA:mpW, GGA:PBE, GGA:RPBE, GGA:revPBE, GGA:mPBE,
    GGA:OLYP, GGA:OPBE,
    Model:SAOP, Model:LB94,
    Hartree-Fock,
    Hybrid:B3LYP, Hybrid:B3LYP*, Hybrid:B1LYP, Hybrid:KMLYP, Hybrid:O3LYP,
↳Hybrid:X3LYP,
    Hybrid:BHandH, Hybrid:BHandHLYP, Hybrid:B1PW91, Hybrid:MPW1PW, Hybrid:MPW1K,
    Hybrid:PBE0, Hybrid:OPBE0
-e: XC energy after SCF (Default, LDA+GGA_METAGGA, LDA+GGA+METAGGA+HYBRIDS)
-pointcharges: file, file with point charges, one point charge per line (ADF only)
    x y z charge, xyz in Angstrom, charge in elementary units (+1 for a
↳proton)
-efield: Ex Ey Ez the electric field vector (in Hartree/(e Bohr))
-k: replace any key, the single argument will be broken into:
    the key, the replacement value, and END for a block key
    all separated by spaces. To insert a return, add a |
    When the key is not found, it is added just before the ATOMS key
    The -k key may be repeated, and is applied at the end, replacing even earlier
↳changes

-dftbmodel DFTB|SCC-DFTB|DFTB3: select the DFTB model
-dftbparameters dir: select the directory with DFTB parameters
-dftbdispersion [None|Default|D3-BJ|D2|ULG|UFF]: dispersion option to use, default is
↳None

-rxforcefield fname: the ReaxFF force field file
-rxniter n: number of ReaxFF iterations
-rxnrstep n: number of non-reactive iterations (out of the total number of iterations)
-rxtstep T: the time step used in the MD simulation
-rxmethod string: the simulation type: Velocity Verlet + Berendsen|NPT|NVE
-rxmdtemp T: the thermostat temperature
-rxmdpres p: the required pressure

-region name at1 at2 ...: make a region with specified name and atoms, may be repeated
    The atom numbers at1 at2 refer to input order, after geometry modifications,
↳start at 1
    Use at1-at2 to refer to all atoms between at1 and including at2
    If the region key is present all regions already present are deleted
-fragments prefix: set up a fragment calculation, prefix fragment run/job scripts
↳with prefix
    if this key is present fragment run/job scripts will be saved
↳(unless -onejob)
    if a job script is requested, the fragment job names will be
↳prefix.fragname.job
    -onejob: for fragment jobs, concatenate the fragment jobs and final job into one on
↳stdout
-g "key value": set any key to the specified value (note key value within quotes)
    key: internal name in AMSinput for some option, see bin/amsinput.tcl/tpl/
↳Defaults.tpl

```

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```

        value: set gin(key) to the specified value
-nochain: unset chain option (used internally by chain jobs)

OUTPUT
-bondsonly: only the bonds as generated by the GUI will be exported (the GUIBONDS_
↳block)
-logfile: force the specified logfile to be used in the run script
-j: produce a fully runnable job (as the .job files from AMSjobs),
    using the specified jobname.
    The job script produces files like jobname.out, jobname.t21 etc. Several job_
↳scripts can simply
    be concatenated, the results will be stored in different files using th jobname_
↳parameter
    the default is a simple run script (the .run file from AMSinput, files are left_
↳as they are)
-a: save a .ams file that matches the run script, except for the -k arguments
    (they are listed in the user input field)
    amsfile is the name of the AMSinput, including the .ams extension (required)

Example: calculate gradients for a molecule in file mymol.xyz
        amsprep -t GO -m mymol.xyz -k "stopafter ggrads"

Example: calculate gradients for a molecule in file mymol.xyz, using good quality_
↳integration and fit:
        amsprep -t GO -q Good -m mymol.xyz -k "stopafter ggrads"

Example: calculate DFTB frequencies for a molecule in file mymol.xyz
        amsprep -t DFTB-FREQ -m mymol.xyz

```

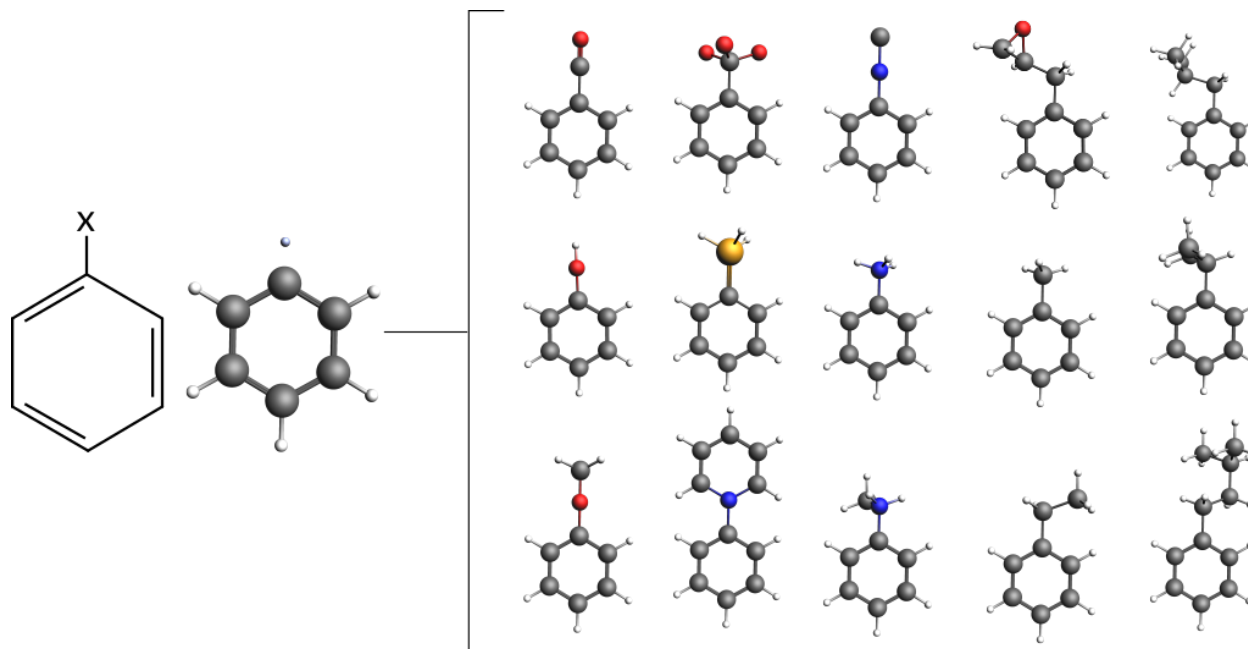
### 2.1.1 Additional Notes

CRSprep represents a scripting solution which is exclusively oriented towards generating input files for the COSMO-RS program.

## 2.2 Tutorial: Generate structures for substituent effects screening

### Overview

Screening substituent patterns of a base compound is a common task in computer aided materials design. In the following short tutorial we demonstrate how you can use amsprep to automatize the replacement of substituents with just a few lines of simple shell scripting.



Contents:

- The library of substituents in AMSinput
- Exchanging substituents with AMSprep
- Combining AMSprep and AMSreport in shell script

### The substituent library in AMSinput

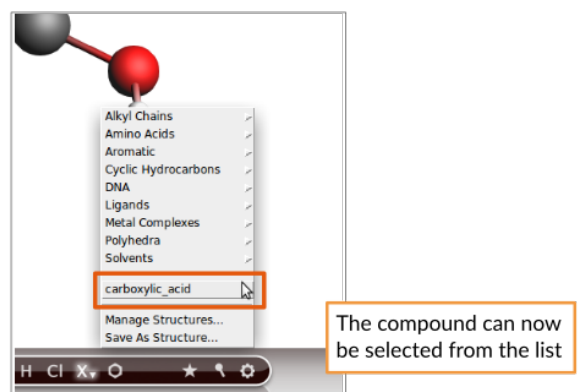
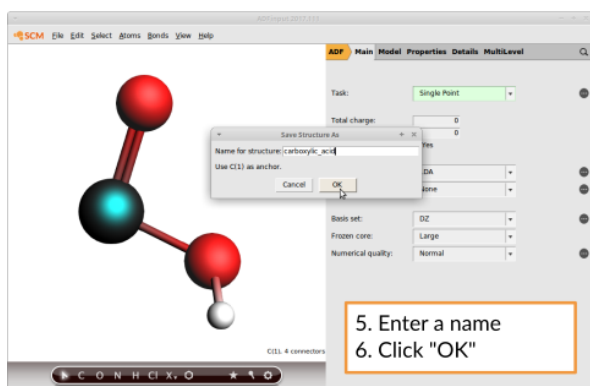
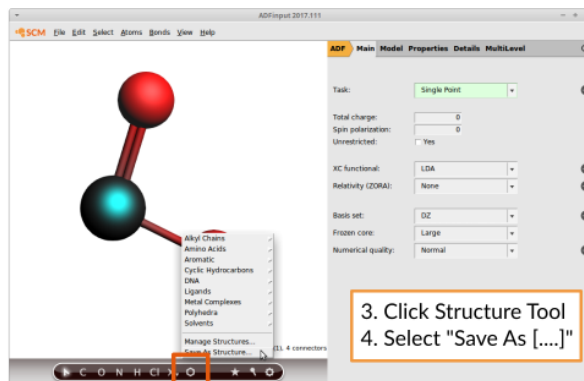
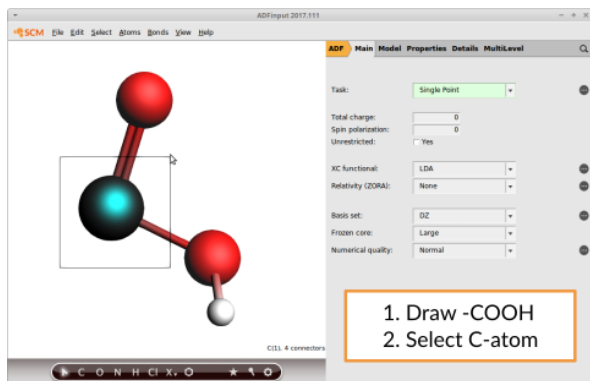
AMSinput comes with a customizable library of common substituents that we can use for our screening purposes right away. It can be accessed via the structure builder tool in AMSinput:

Note how the entries are organized. For example the isocyanide functional group (“NC”) can be found in “Ligands”.

Its also possible to add your own compounds: Simply draw the structure of interest and select the atom which will serve

as an anchor.

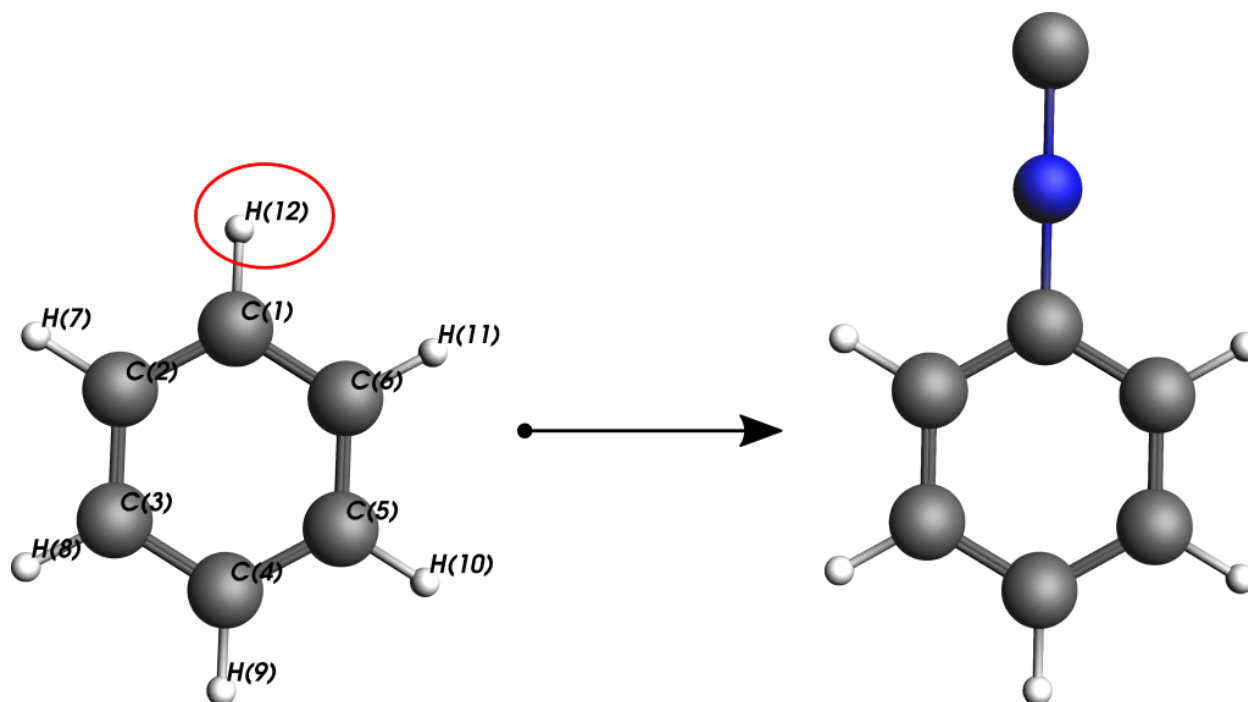
This is how you would add a carboxylic acid group:



AMSinput will always consider the atom that lies in the origin of the coordinate system to be the anchor. If you use the Save As Structure command this will be done for you. More information, including another example, can be found in the [Building Molecules GUI tutorial](#).

### Exchanging substituents with amsprep

Let us consider a simple benzene ring as the base compound:



The `amsprep` command to exchange Hydrogen atom #12 with an isocyanide group (“NC”) and create a runfile for a UFF geometry optimization is:

```
"$AMSBIN/amsprep" -t UFF-GO -m benzene.xyz -structure "12 Ligands/NC.ams" > "benzene_
↳NC.run"
```

Remember that the “CN” group was located in the “Ligands” menu hence “Ligands/NC.ams”. In case the path contains whitespace, you need to escape the whitespace as in this example

```
"$AMSBIN/amsprep" -t UFF-GO -m benzene.xyz -structure "12 Alkyl\ Chains/Ethyl.ams" >
↳"ethyl_benzene.run"
```

When using custom substituents, e.g. the hydroxylic\_acid in the above example, a full path need to be provided to `amsprep`. The path is displayed when clicking on the Structure Tool in AMSinput and selecting “Manage your structures”. On an ubuntu linux system the path is “/home/[your\_username]/.scm\_gui/Structures” and the command to use your own structures becomes:

```
"$AMSBIN/amsprep" -t UFF-GO -m benzene.xyz -structure "12 /home/[your_username]/.scm_
↳gui/Structures/carboxylic_acid.ams" > "benzoic_acid.run"
```

### Bringing it all together

The following few lines of shell script demonstrate how to automatically exchange the substituents on a benzene ring, run a UFF optimization on the new structure and extract the optimized geometry with `amsreport`.

```
#!/bin/sh
#
# copy the file benzene.xyz from the ADF compounds database
#
cp "$AMSHOME/atomicdata/Molecules/ADF/Benzene.xyz" .
#
# loop through different substituents
#
```

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```

for ligand in CN CO CO3 NC NH2 NH2CH3 NH3 OC OCH3 OH PH3 Pyridine; do
#
# prepare the coordinates and the UFF calculation
#
"$AMSBIN/amsprep" -t UFF-GO -m Benzene.xyz -structure "12 Ligands/$ligand.ams" >
↪"benzene_$ligand.run"
#
# run UFF GeoOpt
#
sh "./benzene_$ligand.run"
#
# extract the optimized geometry via amsreport
#
"$AMSBIN/amsreport" uff.rkf SDF > "benzene_$ligand.mol"
#
# rename the generic UFF output file
#
mv uff.rkf "benzene_$ligand.rkf"
done

```

### Running the script

Linux and Mac: Copy and paste the above into a file called `substituents_script` and execute it in the command line

```
sh substituents_script
```

Windows: Just use the pre-configured shell `ams_command_line.bat` shipped with ADF to run the same command as the Linux and Mac users.

## 2.3 AMSreport: generate reports

The utility AMSreport (`$AMSBIN/amsreport`) allows to retrieve the results (including images) computed from the binary output files of either ADF, BAND, ReaxFF, DFTB, UFF, or MOPAC. For ADF this is the `.t21` file (TAPE21). It can also be the `.runkf` file from BAND, the `.rxkf` file from ReaxFF or the `.rkf` file from DFTB, MOPAC or UFF.

The selected results are printed out via standard output or, alternatively, either written to a tab separated file or an HTML file. When creating a new output file AMSreport will also generate a line with headers identifying the information. Images are generated using the ADF-GUI.

Also individual KF variables can be retrieved from the file as shown by the following example, which illustrates how to obtain the bonding energy from a `.t21` file.

```
amsreport job.t21 BondingEnergy
```

Also high-quality pictures of orbitals can be obtained as shown below.

```
amsreport job.t21 HOMO LUMO+1 -v "-grid Fine" -v "-antialias" -v "-bgcolor #ffffff"
```

The options of AMSreport are listed when running the module without further command line arguments. At present the following command line options are available

**-h** prints the help screen.

**Hint:** If used with the name of a valid KF file in the command line the `-h` option lists the names of all data blocks present in that file. It is strongly encouraged to use this option to retrieve the names of the options available in a given situation.

```
amsreport -h job.t21
```

- i** specifies the input file (.t21 etc). If the specified input file is not present ADF tries to find a valid input file based on the information in the matching .ams file or the most recent available binary output file.
- usefile** specifies the input file like `-i` but without attempting to find a matching file if the specified input file does not exist. Typically `-usefile` is used to avoid reading data from the result file.
- I <pattern>** glob files, and run over all matching result files
- o** the name of the html file in which the output of AMSreport will be stored. The output will be printed to standard output if this option is absent.
- plain** print only output data from AMSreport without any labels and/or units. The same can be achieved by setting the environment variable `SCM_AMSREPORT_PLAIN` to yes.
- noplain** print output data with tab separators, labels, and units. Used to override the aforementioned variable `SCM_AMSREPORT_PLAIN`.
- v** command line to pass to amsview (without filenames) to generate images. The image will be generated by AMSview stored in a directory with a name based on the result file, and with extension .jpgs. The result file will contain a path to the image file (directly, or in an IMG tag) After the `-v` the arguments must be listed, with proper quoting. Repeat the `-v` flag for multiple arguments. The individual `-scmgeometry`, `-bgcolor`, `-zoom`, `-viewplane`, `-antialias` and `-grid` options will be collected and applied to all view options.

Some shortcuts are predefined (HOMO, HOMO+1, LUMO, Molecule, Density, Potential) and some additional useful flags include

`-scmgeometry` (default 200x200) `-bgcolor` (default #220000), `-zoom` (default 1.0) `-viewplane` (default {1 2 5}) `-antialias` (off when not present, especially useful with light bgcolors) `-grid` (Coarse when not present, Medium when specified, or value after flag if a value is present)

**examples** HOMO-1 LUMO+1 `-v "-viewplane {0 0 1}" -v "-grid Fine" -v "-antialias"`

- r** Specifies the result to be retrieved by AMSreport from the binary output file. If this command is omitted all unspecified command line arguments but the first (denotes input file name) will be considered as arguments for this flag.

If `-r` is present, the desired result is specified as a string either in form of its preset name (see below) or via a section%variable pair (see the [KF utilities documentation](#)). The `-r` flag (or arguments without flag) may be repeated for multiple results. Additional details can be specified after the variable name, separated by "#". For example

**range** "variable#index" or "variable#firstindex:lastindex", index starts at 1

**format** TclTk format string, e.g. 8.3f or 12.6g

**examples** prints a formatted table of the coordinates

```
-r "Geometry%xyz#12.4f##3"
```

prints a formatted table for the first two atoms only

```
-r "Geometry%xyz#1:9#12.4f##3"
```

coordinates of the first two atoms in one line

```
-r "Geometry%xyz#12.4f#1:9"
```



print just the first coordinate

```
-r "Geometry%xyz#1"
```

print the bond energy

```
-r "Energys%Bond Energy"
```

While any proper KF variable can be accessed via a “section%variable” construct, the following predefined keys are available for the KF files resulting from the various programs of the Amsterdam Modeling Suite.

#### ADF-specific ``-r`` presets for .t21 files

**orient\*** affine transform (3x4) from input to internal ADF orientation, format after #

**iorient\*** affine transform (3x4) from internal ADF to input orientation, format after #

**title** title of the calculation

**type** calculation type (single point, geometry optimization, ...)

**weight** molecular weight

**symmetry** molecular symmetry

**natoms** number of atoms

**integration** integration accuracy

**integration-min** minimum integration accuracy

**integration-max** maximum integration accuracy

**scfstatus** SCF convergence status

**charge** the requested charge

**charges** shorthand for Voronoi, Hirshfeld and Mulliken charges

**voronoi** Voronoi deformation charges

**hirshfeld** Hirshfeld fragment charges, atomic fragment definition required

**mdc** All available MDC atom charges

**mdc-m** MDC-M charges

**mdc-d** MDC-D charges

**mdc-q** MDC-Q charges

**mulliken** Mulliken charges

**bondorders** Mayer bond orders

**nmr** NMR shieldings

**nmr-shieldings** NMR shieldings

**nmr-shielding-tensor** NMR shielding tensor

**nmr-j-coupling-tensor** NMR j coupling tensor

**nmr-k-coupling-tensor** NMR k coupling tensor

**nmr-j-coupling-constant** NMR j coupling constant

**nmr-k-coupling-constant** NMR k coupling constant

**dipolev\*** dipole vector

**dipole** dipole moment (length of dipole vector)

**quadrupole** quadrupole tensor

**orbital-info** orbital info (energy, occupation and label), format for energy after #, range after # with HOMO or LUMO for example:

```
orbital-info#HOMO, orbital-info#HOMO-1,  
orbital-info#HOMO-2:LUMO+2, orbital-info#HOMO#12.8f
```

**orbital-e\*** orbital energies, format and range after # as in orbital-info

**orbital-o\*** orbital occupations, format and range after # as in orbital-info

**orbital-l\*** orbital labels, format and range after # as in orbital-info

**homo-lumo-gap\*** HOMO-LUMO gap, format after #

**atomlabels** name of atoms with sequence number, starting at 0

**atomlabels-from0** name of atoms with sequence number, starting at 0

**atomlabels-from1** name of atoms with sequence number, starting at 1

**nstep** number of steps in history / LT / IRC data, type (h,lt,ircf,ircb) after #

**spin** the requested spin polarization

**step** use coordinates from history / LT / IRC data, step number after # with h for history, lt for LT, ircf/ircb for forward/backward IRC if no letter after #, history data will be used (if not, last step will be used) for example:

```
step#23 (or step#h23), step#lt4, step#ircf3
```

**geometry, geometry-a\*, geometry-b\*** geometry (element type and coordinates), in input order, in angstrom or bohr (default)

**sdf** geometry in SDF format

**bgf** geometry in BGF format

**distance** distance between two atoms, in angstrom. Input separated by #

labels (optional): include atom labels in output

format (optional): format field

atom numbers, starting at 1, in input order

examples

```
distance#2#3, distance#labels#2#3, distance#-8.3f#5#8,  
distance#labels#8.4f#1#2, distance#2#3#4#5, distance#labels#1#2#3#4
```

**angle** angle between three atoms, in degrees. Input see distance, but with three atoms per angle

**dihedral** dihedral between four atoms, in degrees. Input see distance, but with four atoms per dihedral

**hessian\*** Hessian matrix (from GeoOpt%Hessian\_CART), fmt and nperline options after #

**gradients\*** gradients with respect to nuclear displacements (from GeoOpt%Gradients), fmt and nperline options after #

**energies\*** all available energies (bonding up to xc, with labels), fmt option after #

**bonding** total bonding energy

**pauli** total pauli repulsion

**steric** total steric interaction

**orbital** total orbital interaction

**electrostatic** electrostatic energy

**kinetic** kinetic energy

**coulomb** electrostatic (steric and orbital interaction) energy

**xc** exchange-correlation energy

**dispersion** dispersion energy

**frequencies\*** IR Frequencies, format, nperline and range (n, or n:n, start at 1) after #

**frequint\*** IR Intensities, format, nperline and range (n, or n:n, start at 1) after #

**freqlabel\*** IR Frequencies label (symmetry), format, nperline and range (n, or n:n, start at 1) after #

**normalmode\*** normal modes (mass weighted), format, nperline and range (n, or n:n, start at 1) after #

**zeropoint\*** zero-point energy

**excitation\*** Excitation energies, format, nperline and range (n, or n:n, start at 1) after #

**oscillatorstrength\*** Oscillator strengths for the excitation energies format, nperline and range (n, or n:n, start at 1) after #

**excitlabel\*** Excitation labels (symmetry), format, nperline and range (n, or n:n, start at 1) after #

**BAND specific ``-r`` presets for .runkf files**

**natoms** number of atoms

**geometry, geometry-a\*, geometry-b\*** geometry (element type and coordinates), in input order, in angstrom or bohr (default)

**sdf** geometry in SDF format

**bgf** geometry in BGF format

**distance** distance between two atoms, in angstrom. Input separated by #

labels (optional): include atom labels in output

format (optional): format field

atom numbers, starting at 1, in input order

examples

```
distance#2#3, distance#labels#2#3, distance#-8.3f#5#8,
distance#labels#8.4f#1#2, distance#2#3#4#5, distance#labels#1#2#3#4
```

**angle** angle between three atoms, in degrees. #4 Input see distance, but with three atoms per angle

**dihedral** dihedral between four atoms, in degrees. Input see distance, but with our atoms per dihedral

**atomlabel, atomlabel-from0** name of atoms with sequence number, starting at 0

**atomlabel-from1** name of atoms with sequence number, starting at 1

**ReaxFF specific presets for .rxkf files**

**natoms** number of atoms

**geometry, geometry-a\*, geometry-b\*** geometry (element type and coordinates), in input order, in angstrom or bohr (default)

**distance** distance between two atoms, in angstrom. Input separated by #

labels (optional): include atom labels in output

format (optional): format field

atom numbers, starting at 1, in input order

examples

```
distance#2#3, distance#labels#2#3, distance#-8.3f#5#8,
distance#labels#8.4f#1#2, distance#2#3#4#5, distance#labels#1#2#3#4
```

**angle** angle between three atoms, in degrees. #4 Input see distance, but with three atoms per angle

**dihedral** dihedral between four atoms, in degrees. Input see distance, but with four atoms per dihedral

**atomlabel, atomlabel-from0** name of atoms with sequence number, starting at 0

**atomlabel-from1** name of atoms with sequence number, starting at 1

**rx-frame n options** information for a particular reaxff frame. Note the spaces, you will need to quote this key.

```
n: frame number 0, 1, 2, ... (is not the ReaxFF step number)

options: combination of the following (if omitted, all will be reported)

  nframes: total number of frames

  step: the ReaxFF step number for the specified frame

  nats: number of atoms

  xyz: the xyz coordinates

  names: element names (C, H etc) for each atom in the same order as the
  ↪coordinates

  neighbors: bond information

  cell: cell information
```

example

```
amsreport water.rxxf "rx-frame 20 step xyz cell"
```

**pdbtrajectory** the trajectory information (including molecule details) as a sequence of PDB models due to limitations of the PDB format to less than 100000 atoms and it will not be a standard conforming PDB file

**pdbtrajectory-**(nobonds|usepdbinfo)

```
nobonds: as pdbtrajectory, but no bond info (CONNECT records)

usepdbinfo: as pdbtrajectory, but use pdb residue info from first step instead of
  ↪reaxff mol info

xmol: the trajectory information (only element, xyz) in xmol format
```

(continues on next page)

(continued from previous page)

```

gro: trajectory as .gro file (xyz and velocities) options after a - sign:

  m : print list of molecule names and formulas only

  x : allow xyz only frames (missing velocities)

  f : add forces if available

  tf : add the time step, f is a floating point number that is the time per step in
↳ps

examples: gro-x, gro-f, gro-xf, gro-ft0.0001, gro-xt0.001, etc.

Special features for ReaxFF parameter optimization: a geo file in biograph format can
↳be converted from a DFT result file using the bfg option above.

```

**example** Input file: geo (biograph format)

-rxtrainset: run over frames in the input file (should be a bgf BIOGRAPH file), put all charges, bonds and angles in the trainset.in (on stdout).

Input file: ffield (reaxff force field file). The source ffield file determines which atoms, bonds etc are present.

-ffield-min: generate ffield file with all values replaced by min values

-ffield-max: generate ffield file with all values replaced by max values

-ffield-bool: generate ffield file with all values replaced by bool values

-minmax filename: use data from filename for min and max values,

format: see RxParRange.txt in atomicdata/ForceFields/ReaxFF

**General presets for .rkf files**

**natoms** number of atoms

**geometry**, **geometry-a\***, **geometry-b\*** geometry (element type and coordinates), in input order, in angstrom or bohr (default)

**sdf** geometry in SDF format

**bgf** geometry in BGF format

**distance** distance between two atoms, in angstrom. Input separated by #

labels (optional): include atom labels in output

format (optional): format field

atom numbers, starting at 1, in input order

examples

```

distance#2#3, distance#labels#2#3, distance#-8.3f#5#8,
distance#labels#8.4f#1#2, distance#2#3#4#5, distance#labels#1#2#3#4

```

**angle** angle between three atoms, in degrees. #4 Input see distance, but with three atoms per angle

**dihedral** dihedral between four atoms, in degrees. Input see distance, but with our atoms per dihedral

**hessian\*** Hessian matrix (from GeoOpt%Hessian\_CART), fmt and nperline options after #

**gradients\*** gradients with respect to nuclear displacements (from GeoOpt%Gradients), fmt and nperline options after #

**energies** all available energies (bonding up to xc, with labels), fmt option after #

### 2.3.1 Additional notes

- SDF and BGF records can be produced from from ANY file that can be read by AMSinput.
- KFReader is a free (LGPL) alternative to AMSreport. The C sources are [available in our download section](http://www.scm.com/Downloads/KFReader-20140106.zip) (<http://www.scm.com/Downloads/KFReader-20140106.zip>) and can be modified for more specific needs.

## 2.4 KF command line utilities

There are four utility programs for manipulating files in the so-called Keyed File (KF) format from the command shell. Two of them convert KF files from binary to ASCII and vice versa. See the pkf and dmpkf utilities for a description of the ASCII format of a kf file. Such a readable version of a KF file can be useful to inspect its contents in detail.

All programs from the package will convert a KF file to the binary format native to this platform if necessary. In such a case, the original file will be renamed to a file with tilde “~” appended to its name and a message will printed on the standard output.

The KF software was developed at the Vrije Universiteit Amsterdam as a general-purpose package for storing data and re-accessing it via keyword-driven procedures.

### pkf

```
pkf file1 { file2 ... fileN }
```

pkf prints a summary of the contents of the kf files file1... fileN.

All variables are listed by name, type (integer, real, character, logical), and size (number of array elements) and bundled into named sections.

To put the results in an ASCII file for later inspection:

```
pkf file > ascii_result
```

Each section on the file contains an index of its variables and their associated values. All data are organized in blocks. Each section may have any number of index blocks and any number of data blocks (this depends simply on the amount of data to be stored in such a block). In addition there is one special section, the SuperIndex, which is an index of all sections on the file.

The output of pkf consists of:

- General information about the file (name of the file, internally used unit numbers during processing the file...)
- A summary of the SuperIndex, hence an index of blocks in the file and the associated sections.
- A summary: total numbers of blocks associated with the different types of blocks.
- For each section a list of its variables. For each variable in the list the following is displayed
  - The variable name.
  - Its length, i.e. the storage requirements of the variable within the file.
  - Its ‘used’ size, hence the file storage associated with the variable (in units off 8 Bytes for double precision real numbers, 4 for integers, etc.).

- The number of actual elements within the variable (for real, integer, and logical data types) or the number of characters in a string.
- The (logical) index of the data block it is stored in.
- The off-set of the data within its data block.
- Its value or the first element of an array variable, respectively.

**cpkf**

```
cpkf file1 file2 {key1 .. keyn}
```

cpkf copies the sections and/or variables key1 .. keyn from file1 to file2.

If a referenced section or variable already exists on file2 it is overwritten, else it is created. Sections and variables which are already present on file2 but which are not referenced in the command are not affected.

If no sections and/or variables are explicitly mentioned at all the copying is carried out for all sections and variables on file1.

As a side effect of this operation any ‘holes’ eventually present in the original due to the formal deletion of obsolete sections and variables are not copied. Note that the KF file is not rearranged upon deletion of data. Rather only the corresponding entries in the index tables are removed in this case. During the copying process the data is however rearranged for optimum storage efficiency and the resulting file copy may therefore be smaller than the corresponding original.

Skipping specific sections during the copying process can be manually controlled as follows:

```
cpkf file1 file2 -rm section1 ...
```

In this form, all sections will be copied except for the ones specified on the command line, thus effectively removing them from the file.

To copy and rename a section:

```
cpkf file1 file2 "section_name --rename new_section_name"
```

**dmpkf**

A utility to extract information from a KF file and make it available in ASCII format:

```
dmpkf file {key1 .. keyn}
```

dmpkf prints the sections and/or variables from the file file indicated by key1 .. keyn on standard output. The complete file is printed if no sections or variables are specified.

The format to be used for the individual keys:

```
Sec%Var
```

where Var the variable of interest present in section Sec. The complete section is dumped if no variable name is specified.

By redirecting the result to another file a human readable output is obtained:

```
dmpkf file > ascii_result
```

The output contains for each printed variable:

- One line with the name of the section it belongs to;
- One line with the name of the variable itself;
- One line with three integers:

- The amount of space reserved for the variable on the file which is, however, relevant for programs operating with KF files only;
- The amount of data associated with the variable: for reals, integers, logicals: the number of such elements; for strings: the number of characters;
- An integer code for the data type of the variable: 1=integer, 2=real, 3=character, 4=logical;
- The values of the variable (on as many lines as necessary): for scalar variables only one value, for arrays as many values as the array contains.

### **udmpkf**

A utility to put information read from standard input into a KF file:

```
udmpkf file
```

udmpkf reads an ASCII file in the format created by dmpkf from standard input and creates the binary KF file therefrom. If such a KF file is already present the sections and variables in the input file are appended to the existing KF file. Whenever a section or variable already exists in target file it will be overwritten. Other data on the target file are not affected.

The combination of dmpkf and udkmpkf makes it easy to modify KF files with a normal text editor:

```
dmpkf TAPE21 > t21_ASCII
```

After the desired modifications within t21\_ASCII this file may be reconverted into a binary KF file:

```
udmpkf < t21_ASCII TAPE21_new
```

Also note that dmpkf and udkmpkf only require a single argument here, respectively, as “< t21\_ASCII” passes the content of the edited file via the standard input.



## PYTHON STACK IN AMSTERDAM MODELING SUITE

### 3.1 General

The Amsterdam Modeling Suite includes a python stack based on the [Enthought Python Distribution](https://www.enthought.com/products/epd/) (<https://www.enthought.com/products/epd/>). In the AMS2023 release, the python version is 3.8.12.

AMS version	Python version
AMS2023	3.8.12
AMS2022	3.6.9
AMS2021	3.6.9
AMS2020	3.6.9

This python stack is completely separate from any other python installations on the system.

All programs within the Amsterdam Modeling Suite launch python via a special command, `amspython`. In previous versions of the Amsterdam Modeling Suite, the command was called `startpython`, but starting with AMS2020, the python launcher is called `amspython`.

### 3.2 Included modules

Some of the included modules are:

- `numpy` (1.21.2) and `scipy` (1.8.0): Big modules with a lot of functionality for math and science, more information on the [SciPy website](https://www.scipy.org/) (<https://www.scipy.org/>).
- `IPython` (7.22.0): An improved interactive python shell, more information can be found on the [iPython website](https://ipython.org/) (<https://ipython.org/>). Can be started with:

```
$AMSBIN/amsipython
```

- `ase` (3.22.1): ASE (Atomic Simulation Environment) is a python module for atomistic simulations, more information in the [ASE documentation](#) (page 25).
- `matplotlib` (3.5.1): A library for plotting data in 2D, more information on the [Matplotlib website](https://matplotlib.org/) (<https://matplotlib.org/>). We do not ship an interactive backend for matplotlib, so make sure to set a [non-interactive backend](https://matplotlib.org/faq/howto_faq.html#generate-images-without-having-a-window-appear) ([https://matplotlib.org/faq/howto\\_faq.html#generate-images-without-having-a-window-appear](https://matplotlib.org/faq/howto_faq.html#generate-images-without-having-a-window-appear)) when using it. For example the Agg backend for PNGs:

```
import matplotlib
matplotlib.use('Agg')
```

- `pip` (21.3.1): The recommended tool for installing packages from the Python Package Index (PyPI). The [pip documentation](https://pip.pypa.io/en/stable/) (https://pip.pypa.io/en/stable/) explains in detail how to use this tool, but for the Python stack shipped with the Amsterdam Modeling Suite all `pip` commands need to be prefixed with `$AMSBIN/amspython -m`:

```
$AMSBIN/amspython -m pip list
$AMSBIN/amspython -m pip show scipy
$AMSBIN/amspython -m pip search rotate-backups
$AMSBIN/amspython -m pip install rotate-backups
```

- `flexmd`: A module for running MD simulations with adaptive QM/MM regions. Details can be found in the [FlexMD documentation](#) (page 27).
- `plams`: PLAMS (Python Library for Automating Molecular Simulation) is a collection of tools that aim at providing powerful, flexible and easily extendable Python interface to molecular modeling programs. It takes care of input preparation, job execution, file management and output processing as well as helps with building more advanced data workflows. See the [PLAMS tutorials](#) and [PLAMS documentation](#) for more information.
- `autografs`: AuToGraFS stands for Automatic Topological Generator for Framework Structures. Information and examples can be found in the [AuToGraFS documentation](#) (page 39).

### 3.3 Using other modules with the AMS Python Stack

You can extend the the AMS Python Stack with other modules. You can use `pip` (see above) to install additional modules if they are available on the Python Package Index (PyPI (https://pypi.python.org/pypi)). This is the recommended way of installing packages:

```
$AMSBIN/amspython -m pip install name_of_package
```

You can also do this for your own packages that you develop, for example:

```
$AMSBIN/amspython -m pip install -e .
```

Alternatively, you can add the location of the source to the `SCM_PYTHONPATH` variable to make the module available in the AMS Python Stack. To avoid collisions with other python installations on the system, we unload `PYTHONPATH` and `PYTHONHOME` from the environment when launching the ADF Python Stack and put the content of `SCM_PYTHONPATH` into `PYTHONPATH`.

---

**Hint:** If you for some reason have to use the `PYTHONPATH` variable and are unable to use `SCM_PYTHONPATH`, you can modify `$AMSBIN/amspython` and `$AMSBIN/amsipython` to not have it cleared when starting python.

---

### 3.4 Install and run Jupyter Lab (Jupyter Notebooks)

#### 3.4.1 Install Jupyter Lab

**AMS2023 or later:** Install Jupyter lab in the [Package Manager](#) (GUI: SCM → Packages, Command-line: `$AMSBIN/amspackages gui`).

**AMS2022:**

```
"$AMSBIN/amspython" -m pip install -U jupyter_client jupyterlab
```

---

**Note:** On Windows, you may first need to upgrade pip:

```
"$AMSBIN/amspython" -m pip install -U pip
```

---

## 3.4.2 Run Jupyter Lab

To launch:

```
"$AMSBIN/amspython" -m jupyterlab
```

## 3.5 Python virtual environment

### 3.5.1 Default python virtual environment

Starting with AMS2020, the `amspython` command by default checks for a python virtual environment inside the user's home directory. If it does not find a virtual environment, it will create one in the following location (these are the default values for the `SCM_PYTHONDIR` environment variable):

- Windows: `$USERPROFILE/.scm/python`
- Mac: `$HOME/Library/Application Support/SCM/python`
- Linux: `$HOME/.scm/python`

`amspython` then launches the python binary located inside the virtual environment.

If you install additional packages via `pip` (see above), they will be installed into the virtual environment's `site-packages`.

### 3.5.2 Virtual environments for different AMS versions

The virtual environment is tied to the major release version of AMS, which is reflected in the name of the virtual environment directory (e.g. `AMS2020.1.venv`).

Thus, upgrading from `AMS2020.101` to `AMS2020.102` will automatically let you use all python packages that you installed into the `AMS2020.101` virtual environment.

To use previously installed python packages when upgrading from `AMS2020.101` to e.g. `AMS2020.301` or `AMS2021.101`, simply copy or rename `AMS2020.1.venv` to `AMS2020.3.venv` or `AMS2021.1.venv` in the same directory.

If you have several installations of different major releases of the Amsterdam Modeling Suite on the same computer, multiple virtual environments will also be created, one for each installation.

### 3.5.3 Changing the location of the python virtual environment

Set the `SCM_PYTHONDIR` environment variable to a directory in which the virtual environment will be installed. If you use the graphical user interface (GUI), this environment variable can be changed in the GUI preferences. If you use the command line, set it in your `amsbashrc.sh`. If you use both the GUI and the command line, you should change it in both places.

### 3.5.4 Disabling the virtual environment

Follow the steps for changing the location of the python virtual environment, but set `SCM_PYTHONDIR` to be empty.

### 3.5.5 Uninstalling the virtual environment

Just delete the directory containing the virtual environment. This will also delete any packages that you have installed into it.

### 3.5.6 Reinstalling the virtual environment

To force a reinstallation of the virtual environment, even if it already exists, run `$AMSBIN/amspython --install_venv`. This will not remove any packages inside the virtual environment.

## 3.6 Useful commands

- Start a python shell:

```
$AMSBIN/amspython
```

- Install new packages via pip:

```
$AMSBIN/amspython -m pip install name_of_package
```

- Find out the location of an installed package (e.g. numpy):

```
$AMSBIN/amspython -c 'import numpy; print(numpy.__file__)'
```

- Find out which python binary is launched by amspython:

```
$AMSBIN/amspython -c 'import sys; print(sys.executable)'
```

- Disable the virtual environment (one-time):

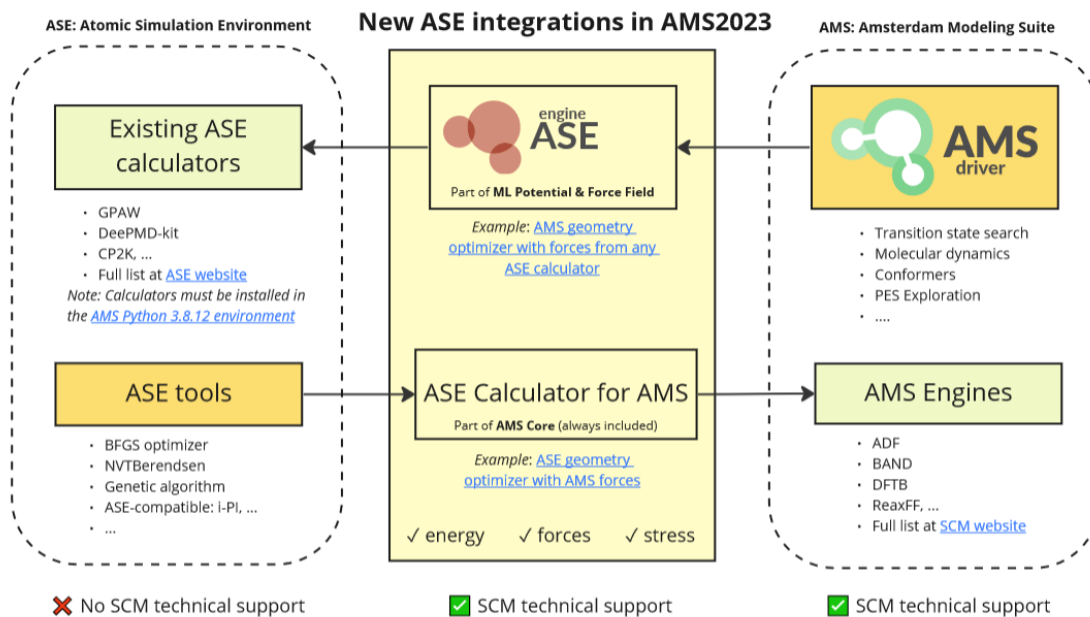
```
SCM_PYTHONDIR='' $AMSBIN/amspython
```

## ASE

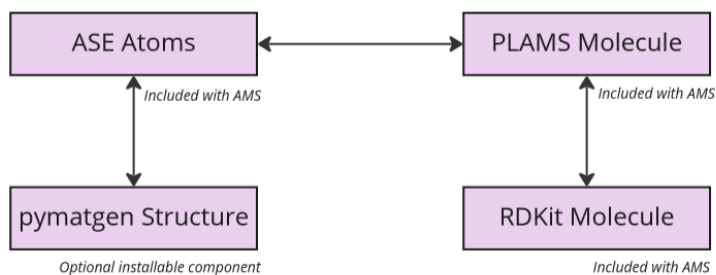
The [Atomic Simulation Environment \(ASE\)](https://wiki.fysik.dtu.dk/ase/) (<https://wiki.fysik.dtu.dk/ase/>) tool collection suite was designed as a flexible, easy-to-use, and customizable approach for the manipulation of quantum chemical models as well as for setting up and running the calculations required and for the analysis of the final results.

The development of ASE was originally started at the Technical University of Denmark but received also significant contributions from a large, international community. While ASE is [available](https://gitlab.com/ase/ase/tree/3.13.0/) (<https://gitlab.com/ase/ase/tree/3.13.0/>) under a GNU LGPL license, a modified version of this library is shipped together with the Amsterdam Modeling Suite.

PLAMS provides an interface to use [AMS as an ASE Calculator](#), while [Engine ASE](#) allows [ASE Calculators](#) (<https://wiki.fysik.dtu.dk/ase/ase/calculators/calculators.html#supported-calculators>) to be used within the AMS driver.



### Chemical systems in Python



**Note:** On Windows machines the developers of the ASE library provide only a partial support, see [the ASE website](https://wiki.fysik.dtu.dk/ase/download.html#installation-on-windows) (<https://wiki.fysik.dtu.dk/ase/download.html#installation-on-windows>) for further details.

## FLEXMD

FlexMD is a python library for Flexible multi-scale Molecular Dynamics simulation, developed by Rosa E. Buló, Christoph Jacob, Stefano Borini, Tao Jiang, Jelle Boereboom, Stanislav Simko and Hans van Schoot.

### 5.1 Basic philosophy and intended usage

We present a flexible python library for molecular dynamics, specialized in multi-scale simulations in a broad sense. At its core, the library interfaces the *Atomistic Simulation Environment (ASE)* (page 25) [*I* (page 34)] molecular dynamics modules with a wide range of molecular mechanics and electronic structure codes. As such, it allows simple dynamics using forces computed with any energy/gradient evaluator provided by the ADF package.

Additionally, FlexMD allows the partitioning of a system into regions described at different resolution, with the aim of running multi-scale (hybrid) force calculations. Besides the traditional, rigid, multi-scale partitioning, FlexMD includes different schemes for Adaptive Multi-scale Molecular Dynamics. Such simulations allow the resolution of a particle to change according to its distance from a predefined active site, which is a necessity for successful multi-scale description of diffusive systems such as chemical reactions in solution.

Finally, the library couples the dynamics to rare events techniques, either implemented in FlexMD itself, or accessible through an interface with the PLUMED library for free energy calculations [*7* (page 35)], opening the possibility for evaluation on time-scales beyond the reach of standard molecular dynamics simulations.

The FlexMD package is designed to make simulation options possible that are not available natively in the ADF package. Its flexible nature makes it very versatile, but comes at a cost. This cost might be completely negligible in most simulations, but it can be very high in some cases (usually when combining only cheap methods such as forcefields).

The intended users for the FlexMD package are those with some Unix/Linux experience and a basic understanding of the [Python Programming Language](http://python.org/) (<http://python.org/>). The user is also supposed to have a basic understanding of the various methods he wishes to combine. For example, if metadynamics is supposed to be combined with ADF, FlexMD expects the user to have knowledge about DFT calculations and the usage of Collective Variables. Finally, as with every computational method, the user should monitor the FlexMD performance, both in accuracy and speed.

### 5.2 FlexMD functionality summary

#### Molecule

##### Input/output

- Reads and writes PDB and XYZ files
- Reads and writes topology data (in CHARMM format)
- Reads and writes force field data (on CHARMM format)

### Analysis

- Extracts geometry data

### Drawing functionality

- Adds atoms and bonds
- Changes bond-lengths, angles and torsions
- Cuts fragments
- Cuts solvent boxes and droplets
- Performs rotations and translations, to fit bonds to axes and planes

### Periodic functionality

- Adds periodic images
- Wraps molecules into periodic box

### Water specific

- Finds hydrogen bonds
- Finds shortest water bridge connecting H-donor and acceptor

## Energy and force calculations

### Standard

- ADF
- DFTB
- REAXFF
- UFF
- MOPAC
- NAMD
- Lennard-Jones force fields

### Multi-scale

- QM/MM, mechanical embedding: Combines all the codes above
- Hybrid: More flexible than QM/MM. Combines different force calculations by summing or subtracting the energies and forces. The standard calculations (above) can therefore be combined with:
  - Metadynamics
  - Plumed (external code that computes free energy data)
  - Constraints
- Adaptive QM/MM (for chemistry in solution)
  - Difference-based Adaptive Solvation (DAS)
  - Sorted Adaptive Partitioning (SAP)
  - Buffered-Core (BC)
  - Flexible Inner Region Ensemble Separator (FIRES)

## Molecular Dynamics



- Uses ASE as the molecular dynamics driver for all above methods
- Analyses trajectories

## 5.3 Introduction

FlexMD is a python package providing molecular dynamics (MD) simulations using the energy evaluation methods made available by the ADF suite. A set of example scripts can be found in the examples/scmlib directory of a standard ADF installation.

FlexMD can be accessed interactively by running `amspython`, followed by a standard python import command for the package `scm.flexmd`. The python help function can be used to obtain detailed documentation about all FlexMD classes. In the following example, an inquiry of one class (the `MDMolecule` class) can be performed.

```
$ amspython
from scm import flexmd
help(flexmd.MDMolecule)
```

To leave the interactive help, press `q`. The help function can also be used to list the contents of the FlexMD package:

```
$ amspython
from scm import flexmd
help(flexmd)
```

Python can also give the help documentation as plain text:

```
$ amspython
from scm import flexmd
import pydoc
print pydoc.render_doc(flexmd.ForceJob, "Help on %s")
```

## 5.4 Molecular Dynamics

FlexMD defines the molecular system under study through the **MDMolecule** class: an instantiation of this class holds all information about the molecular system to be simulated, such as coordinates, topology, and force field parameters (if needed). An **MDMolecule** object can be initialized from a PDB or XYZ file, by specifying its path at object creation.

An interface to energy evaluators is provided by specialized **ForceJob** classes, acting as wrappers around the ADF suite of programs. A **ForceJob** requires an **MDMolecule** object to be specified at creation. The resulting **ForceJob** object can either be used directly by the Atomistic Simulation Environment (ASE) [1 (page 34)] library as a calculator object (see examples/scmlib/ASE\_emt\_h2o) or with the **ASEMDPropagator** class, which provides methods for running an MD time step using ASE classes. Internally, the propagator sets up the required ASE objects, passes the **ForceJob** object to them, and retrieves the new positions and velocities. An additional protagonist, an **MDManager** class instance, coordinates the MD simulation by running the MD steps with the **ASEMDPropagator** object and writing trajectory information to file.

During creation of an **MDManager** object, a directory 'QMMD' is created, which contains a file `TRAJEC00.DCD` holding the geometries along the trajectory, a file `FTRAJEC00.DCD` holding the forces along the trajectory, and finally a file `ENERGY00.dat` holding the potential and kinetic energy, as well as the temperature throughout the evaluation. To extract the geometries from the trajectory file, the **DCDFile** class is available, providing methods to read and write geometries to and from a trajectory file in DCD format. The **MDManager** is also responsible for handling restart of a previous MD evaluation: if a 'QMMD' directory is already present at script invocation, the new output files will be assigned the number subsequent to the highest numbered files in that directory. In addition, provided the previous run terminated normally, the restart will continue from the final geometry and velocity of the previous run.

The ADF package contains different electronic structure methods of varying degrees of accuracy and speed. The best-known methods are the ADF Density Functional Theory (DFT) code itself, and the BAND DFT code for periodic systems. FlexMD provides an interface toward both programs. For the interface with ADF, FlexMD makes use of classes from PyADF [2 (page 34)], a scripting framework for efficient quantum chemistry calculations. In addition to ADF and BAND, several semi-empirical methods are included in the ADF suite, such as DFTB and the NDDO type schemes available in the MOPAC package [3 (page 34)]. The ADF suite also provides classical mechanics methods, such as the reactive force field ReaxFF and the simple force field UFF. Interfaces to all of these methods are available in FlexMD. A simple example of a python script for MD using the **UFFForceJob** class for UFF calculations can be found in the examples directory, under `examples/scmlib/flexmd_uff_h2o`.

To increase the flexibility of FlexMD, an interface towards force calculations using the NAMD2.8 classical molecular dynamics package is provided (`examples/scmlib/namd_h2o`). NAMD2.8 is not distributed with the ADF suite, but it is available from a third party to be downloaded and installed (<http://www.ks.uiuc.edu/Development/Download/download.cgi>).

## 5.5 Multi-scale Molecular Dynamics

The design of the **ForceJob** class allows for flexible extension of its behavior, while at the same time keeping the client code unaware of its nature: it can either act as a simple wrapper for ADF programs, or it can be a more complex orchestrating class, combining simpler **ForceJob** classes to implement multi-scale strategies. One application of this extensible design can be found in the **QMMMForceJob** object, which combines a QM and an MM method in an IMOMM-type scheme (mechanical embedding). The **QMMMForceJob** object is assigned two other **ForceJob** objects, the first representing the high-resolution calculation (QM), while the other represents the low resolution (MM). Both **ForceJob** objects contain an **MDMolecule** object for the full molecular system. The selection of the QM-region is handled by the **QMMMForceJob**, which contains the information about the part of the molecule that constitutes the QM region. When forces are requested from the **QMMMForceJob**, the following behavior is orchestrated: first, a MM force calculation is performed on the full system; then, the QM-region is selected, a QM calculation is executed solely for that region, and energy and forces are added to those from the full system MM calculation. Finally, an MM calculation is computed for the small QM-region, and the energy and forces are subtracted, yielding the final result, returned to the invoker. In symbols:

$$EQM/MM(\text{Full}) = EMM(\text{Full}) + EQM(\text{QMRegion}) - EMM(\text{QMRegion})$$

The **QMMMForceJob** handles periodic boundary conditions if the low-level (MM) method supports this feature (i.e. NAMD). Whether the periodic interaction of the QM region with itself is handled at high or low resolution depends on the method used for the QM calculation. An example of QM/MM MD calculations can be found in the examples directory `examples/scmlib/qmmm_dftbUFF_2h2o`. The **QMMMForceJob** allows the use of link atoms when the QM boundary cuts through covalent bonds. However, this feature comes at the price of an increased script complexity. An example of a QMMM link-atom MD simulation is provided in the examples directory, under `examples/scmlib/qmmm_linkatom_dftbNAMD_glutamate`.

For more complex multi-scale calculations the **HybridForceJob** class can be used. This class allows the combination of a large set of different **ForceJobs**, each of them describing either the same, or different molecular systems. Each **ForceJob** can either involve a calculation on the full **MDMolecule** object it contains, or restricted to a specified region of the corresponding molecule. The forces from each contributing **ForceJob** can either be added or subtracted from the total force according to user preference, as specified at construction of the **HybridForceJob** object.

In order to perform QM/MM simulations on chemical reactivity in solution, it is important that the description of the solvent molecules can change on the fly, as the molecules move towards or away from the reactive region. To facilitate this, an **AdaptiveQMMMForceJob** class is available to provide adaptive QM/MM simulations using several available schemes, as described by Bulo et al.[4 (page 34)] and P. Fleurat-Lessard et al.[8 (page 35)] In these schemes, the description of the diffusing molecules changes gradually from QM to MM and vice versa, based on the distance of those molecules to a predefined reactive site. Various schemes are available for assigning the QM and MM character of the molecules. The class contains a **QMMMForceJob** object, as well as a partitioning object that assigns the partial QM and MM character to the molecules. An examples python script for such an adaptive QM/MM simulation, using the DAS [4 (page 34)] method, is provided in the examples directory, `examples/scmlib/adqmmm_mopacscmUFF_h2o`.

## 5.6 Biased Molecular Dynamics

Constraints can be added to a simulation using the derived **ForceJob** class **WallJob**. The constraint is in the form of a large one-dimensional Gaussian on the potential energy surface, along a predefined Collective Variable (CV). Examples of CV's are the distance between two atoms, the coordination number of two atoms, but also more complex quantities such as the minimum distance between two sets of atoms, or the distance of an atom to a hydroxide ion. The Collective Variables can be specified through the **CollectiveVariable** class. Derived **CollectiveVariable** classes are available to specify sums or multiples of other **CollectiveVariable** objects.

Regular MD calculations are limited in the time-scales achievable with current hardware. The order of such time-scales is much smaller than what is required for chemical reactions. To overcome this problem, two rare-events methods have been implemented directly into the library: metadynamics [5 (page 34)] and umbrella sampling [6 (page 35)]. Both these methods involve biasing the simulations along a CV. An example of a metadynamics input can be found in the examples directory in `examples/scmlib/metadynamics_emt_h2o`.

For a wider range of rare-events methods, FlexMD also offers an interface with the PLUMED library for free energy calculations [7 (page 35)]. To use this, a PLUMED input file is required, and for this we refer to the PLUMED manual. An example of a FlexMD input script using PLUMED can be found in the examples directory in `examples/scmlib/plumed_emt_h2o`.

## 5.7 Working with FlexMD

It is recommended to read the sections *Introduction* (page 29) and *Molecular Dynamics* (page 29) before working with FlexMD. Basic understanding of the [Python Programming Language](http://python.org/) (<http://python.org/>) is also required. The Python website hosts documentation and a [tutorial](http://docs.python.org/2/tutorial/) (<http://docs.python.org/2/tutorial/>) that can be used to learn Python.

The performance of the FlexMD package is difficult to predict because it depends on system size, the type of ForceJobs used and how these ForceJobs are combined. It is advised to first test the overhead of the FlexMD package for your system before running large simulations. When ab initio forces are involved, the overhead should not give a significant performance penalty. However, it may become a bottleneck when your system only uses cheap forcefields.

### 5.7.1 Creating a molecule object

FlexMD can be run through the interactive python interpreter in the ADF package. To start it, run: `$AMSBIN/amspython` in a terminal, followed by:

```
from scm import flexmd
```

Note that it is also possible, and usually more convenient, to write your FlexMD code in a file and then to execute this file. To do this, type all the commands you would use in the interactive interpreter in a file, and then enter `$AMSBIN/amspython myFlexMDjob.py` in a terminal (after changing to the directory where the file was stored of course).

Most FlexMD jobs will start with importing FlexMD and creating an MDMolecule object. This can be done by starting from a geometry in xyz or pdb format, or by manually adding the atoms in the FlexMDjob.py file. Geometries can be generated in the ADF GUI, and then be exported to xyz file. For more details on the MDMolecule object, run `$AMSBIN/amspython, import flexmd and call help(flexmd.mdmolecule)`.

```
from scm import flexmd
myMol = flexmd.MDMolecule('myGeometryFile.xyz')
```

Some ForceJobs require the system to be periodic. If we create an MDMolecule object from a pdb file that includes periodic information, the periodic boundary conditions are automatically imported. If the information is not there, we can add it to the MDMolecule object:

```
myMol = flexmd.pdb.set_box([50.0,25.0,100.0])
```

Info on `set_box` (and other functions, such as `set_cellvectors`, and `write_pdb`) can be found using `help(flexmd.pdbmolecule)`.

It is also possible to write the info in the `MDMolecule` object to a `pdb` file. to do so, call `pdb.write_pdb('mypdbfile.pdb')` on the `myMol` object:

```
myMol.pdb.write_pdb('mypdbfile.pdb', box=True)
```

### 5.7.2 Creating a ForceJob

To specify what type of forces we want to use in the MD simulation, a `ForceJob` must be created. FlexMD has a number of `ForceJobs` (see `PACKAGE CONTENTS` in `help(flexmd)`), most of them with examples in `$AMSHOME/examples/scmlib`. The `ForceJobs` can be combined into a single `ForceJob` using `flexmd.hybrid_ForceJob`. As an example, we combine a `reaxff_ForceJob` with a `metadynamicsjob` and a `walljob`:

```
from scm import flexmd
myMol = flexmd.MDMolecule('myGeometryFile.xyz')
myMol = flexmd.pdb.set_box([50.0,25.0,100.0])
    # setup our reaxff ForceJob and attach the forcefield file
    # (place the ff file in the same dir as the script and the xyz!)
myReaxffForceJob = flexmd.ReaxffForceJob(molecule=myMol)
myReaxffForceJob.settings.set_ff_filename('reax_forcefield_file.ff')

    # next we define the collective variable: the distance between atom 1 and 2
myCvs = [flexmd.DistCV([1,2])]
    # create a set of metadynamics properties, using the CV
mtdSettings = flexmd.MetadynamicsSettings(cvs=myCvs, widths=[0.30], height=0.25 )
    # create the metadynamics job by combining the molecule, settings (with CV)
    # and the number of md steps between depositing metadynamics hills.
myMetadynamicsjob = flexmd.MetadynamicsJob( myMol, settings=mtdSettings, nstep=150 )

    # add a wall to prevent the two atoms from drifting more than 10 Angstrom away.
myWalljob = flexmd.WallJob(molecule=myMol, cvs=myCvs, cntrs=[10.0], widths=[1.0],
↪ heights=[500.0])

    # combine the forces into a hybrid job that will be used for the MD
myForceJob = flexmd.HybridForceJob( [[myReaxffForceJob,'+'], [theMetadynamicsjob,'+'],
↪ [theWalljob,'+']], myMol )
```

Note that all the `ForceJobs` require some special input and settings, and that these settings can be applied both before and after defining the `ForceJob`. For the `reaxffForceJob`, we first define the `ForceJob`, and add the forcefield parameters file afterwards. For the metadynamics job we reverse this, and first create a `metadynamicsJobSettings` object, which is then used in the creation of the metadynamics job. For more detailed info on the different `ForceJobs` and their inputs, see the help function by calling `help` on a `ForceJob`, for example: `help(flexmd.ReaxffForceJob)` or `help(flexmd.WallJob)`. Also remember that other examples of `ForceJobs` can be found in `$AMSHOME/examples/scmlib`.

### 5.7.3 Creating and running the MD job

Before the simulation can be set in motion, a propagator is needed. The `propagatorJob` controls simulation settings such as temperature and timestep size. FlexMD uses the Atomistic Simulation Environment (ASE) [1 (page 34)] for this. The `MDPropagatorJob` object is created just like the other objects in FlexMD:

```
# do this after importing flexmd and creating a ForceJob.
# it creates the MDPropagator job, with some settings
myMDJob = flexmd.ASEMDPropagatorJob( ForceJob=myForceJob )
myMDJob.settings.set_tempcontrol( True, nhfreq=2, maxdef=50.0 )
myMDJob.settings.set_temperature(300.0)
myMDJob.settings.set_timestep( 0.02 )
```

For more details on the `ASEMDPropagatorJob`, view its help page: `help(flexmd.ASEMDPropagatorJob)`, or take a look at the `MDSettings` object: `help(flexmd.MDSettings)`.

The `propagatorJob` can be used to create an `MDManager` object:

```
# create an MD manager
myManager = flexmd.MDManager( mdjob=myMDJob)
```

The manager object is now in control of the MD simulation, and we can use it to run the simulation for a number of steps:

```
# tell the MD manager to run the simulation
myManager.run( ncycles = 2500 )
```

Note that the number of steps here should be increased a lot if metadynamics effects are to be observed, but it is always wise to first run a small number of steps to check if everything works. Some information will be printed during the simulation, depending on the settings of the components used. The manager will also create some folders in the working directory, and store the data produced by the simulation in there.

The full `flexmd` jobfile should now look something like this:

```
from scm import flexmd
myMol = flexmd.MDMolecule('myGeometryFile.xyz')
myMol = flexmd.pdb.set_box([50.0,25.0,100.0])
# setup our reaxff ForceJob and attach the forcefield file
# (place the ff file in the same dir as the script and the xyz!)
myReaxffForceJob = flexmd.ReaxffForceJob(molecule=myMol)
myReaxffForceJob.settings.set_ff_filename('reax_forcefield_file.ff')

# next we define the collective variable: the distance between atom 1 and 2
myCvs = [flexmd.DistCV([1,2])]
# create a set of metadynamics properties, using the CV
mtdSettings = flexmd.MetadynamicsSettings(cvs=myCvs, widths=[0.30], height=0.25 )
# create the metadynamics job by combining the molecule, settings (with CV)
# and the number of md steps between depositing metadynamics hills.
myMetadynamicsjob = flexmd.MetadynamicsJob( myMol, settings=mtdSettings, nstep=150 )

# add a wall to prevent the two atoms from drifting more than 10 Angstrom away.
myWalljob = flexmd.WallJob(molecule=myMol, cvs=myCvs, cntrs=[10.0], widths=[1.0], ↵
↵heights=[500.0])

# combine the forces into a hybrid job that will be used for the MD
myForceJob = flexmd.HybridForceJob( [[myReaxffForceJob,'+'], [theMetadynamicsjob,'+'], ↵
↵ [theWalljob,'+']], myMol )

# do this after importing flexmd and creating a ForceJob.
```

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```

# it creates the MDPPropagator job, with some settings
myMDJob = flexmd.ASEMDPropagatorJob( ForceJob=myForceJob )
myMDJob.settings.set_tempcontrol( True, nhfreq=2, maxdef=50.0 )
myMDJob.settings.set_temperature(300.0)
myMDJob.settings.set_timestep( 0.02 )

# create an MD manager
myManager = flexmd.MDManager( mdjob=myMDJob)
# tell the MD manager to run the simulation
myManager.run( ncycles = 2500 )

```

This job can be resumed by just running the script again, FlexMD should pick up on the coordinates and velocities from the QMMD folder. To do this, simply run the job again, and a new set of trajectory and energy files should appear in the QMMD folder. If you wish to specify which files FlexMD restarts from, you can use the `set_restartnum()` function on `myMDJob.settings`:

```

# add this below the "myMDJob.settings.set_timestep( 0.02 )" line
# FlexMD will restart from the files with index numbers 1 lower than you pass to
↪set_restartnum
# This will cause FlexMD to restart from QMMD/TRAJEC00.dcd and QMMD/TRAJEC00.dcd
myMDJob.settings.set_restartnum(1)

```

## 5.8 Required Citations

When you publish results in the scientific literature that were obtained with programs of the ADF package, you are required to include references to the program package with the appropriate release number, and a few key publications.

**For calculations with FlexMD:** 1. FlexMD 2017, SCM, T. Jiang, H. van Schoot, S. Simko, C. R. Jacob, S. Borini, R. E. Buló, *A python library for flexible multi-scale molecular dynamics simulations*. <http://www.scm.com>

### 5.8.1 External programs and Libraries

[Click here](#) for the list of programs and/or libraries used in the ADF package. On some platforms optimized libraries have been used and/or vendor specific MPI implementations.

## 5.9 References

1. S.R. Bahn, K.W. Jacobsen, *An object-oriented scripting interface to a legacy electronic structure code*. *Comput. Sci. Engin.* 4, 56-66 (2002) (<https://doi.org/10.1109/5992.998641>). The Atomistic Simulation Environment website and documentation (<https://wiki.fysik.dtu.dk/ase/>)
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7. M. Bonomi, D. Branduardi, G. Bussi, C. Camilloni, D. Provasi, P. Raiteri, D. Donadio, F. Marinelli, F. Pietrucci, R.A. Broglia, M. Parrinello, *PLUMED: A portable plugin for free-energy calculations with molecular dynamics*. *Comp. Phys. Comm.* 180, 1961-1972 (2009) (<https://doi.org/10.1016/j.cpc.2009.05.011>). *PLUMED website and documentation* (<http://www.plumed-code.org/>)
8. P. Fleurat-Lessard, C. Michel, R. E. Bulo, *Energy extrapolation schemes for adaptive multi-scale molecular dynamics simulations*. *J. Chem. Phys.* 137, 174111 (2012) (<https://doi.org/10.1063/1.4739743>)





## PLAMS

PLAMS (Python Library for Automating Molecular Simulation) is a collection of tools that aim at providing powerful, flexible and easily extendable Python interface to molecular modeling programs. It takes care of input preparation, job execution, file management and output processing as well as helps with building more advanced data workflows.

- [PLAMS tutorials](#)
- [PLAMS documentation](#)

### 6.1 Required Citations

When you publish results in the scientific literature that were obtained with programs of the ADF package, you are required to include references to the program package with the appropriate release number, and a few key publications.

**For calculations with PLAMS:** PLAMS, written by Michał Handzlik, <http://www.scm.com>

#### 6.1.1 External programs and Libraries

[Click here](#) for the list of programs and/or libraries used in the ADF package. On some platforms optimized libraries have been used and/or vendor specific MPI implementations.



## AUTOGRAFS

The [Automatic Topological Generator for Framework Structures](http://pubs.acs.org/doi/abs/10.1021/jp507643v) (<http://pubs.acs.org/doi/abs/10.1021/jp507643v>) is a software for the design and high throughput generation of framework structures such as Metal-Organic Frameworks (MOFs), Zeolites, Covalent-Organic Frameworks (COFs), or any product of reticular chemistry. The underlying library for chemical objects makes extensive use of the *ASE* (page 25) environment shipped with the Amsterdam Modeling Suite.

AuToGraFS was developed in the group of Prof. Dr. Heine at Jacobs University Bremen by Dr. M. Addicoat and D. Coupry. While a console application is [available](https://github.com/maddicoat/AuToGraFS) (<https://github.com/maddicoat/AuToGraFS>) under a GNU LGPL license, the version shipped together with the Amsterdam Modeling Suite was heavily modified into a fully importable python library for easy scripting capabilities. The software was also fully integrated into the GUI, and some native functions were developed for added functionality in this particular environment. A database of building units often used in existing MOFs is also provided, to cover basic needs.

The rest of this section is dedicated to a documentation and a demonstration of the usage of both the scripting capabilities and GUI tools from the Automatic Topological Generator for Framework Structures.

### 7.1 General AuToGraFS Scripting concepts

AuToGraFS relies on the concept of underlying nets common to many structures. These nets, or topologies can be found in the [RCSR](http://rcsr.anu.edu.au/) (<http://rcsr.anu.edu.au/>) and [EPINET](http://epinet.anu.edu.au/) (<http://epinet.anu.edu.au/>) databases. Once implemented, AuToGraFS will follow these blueprints to generate a framework from building units of compatible geometries. Since these geometries are ideal objects, and most chemical structures are distorted, an optimization step using *UFF* is necessary for the building units to “snap into place”. Correct handling of bonding information and *UFF* atom types are crucial to the generation of correct structures.

#### 7.1.1 Components of AuToGraFS

##### The Fragment class

This is a slightly modified version of the native ASE Atoms object, designed to hold essential data for *UFF* postprocessing. This englobes the *uff* atom types, and the bonding information.

```
from scm.autografs.fragment import Fragment
from ase import Atoms

line = Fragment(ase.Atoms("X2", positions=...),
                mmtypes=["H_", "H_"],
                bonds = [[0,0,1],[0,0,1],[1,1,0]],
                shape="linear",
                unit=None,
                name="a_line_has_no_name")
```

Where `mmtypes` is a list of the UFF atom types symbols, ordered as the corresponding atoms in the structure, and `bonds` is a symmetric `numpy array` (<https://docs.scipy.org/doc/numpy-1.10.0/reference/generated/numpy.array.html>) of size equal to the number of atoms. The  $i,j$  elements in the `bonds` array are the bond order between atom  $i$  and atom  $j$  of the structure. These bond orders should follow the UFF conventions (1.5 for aromatic, 0.5 for coordination bonds, 0.001 for hydrogen bonds...). The shape holds information for AuToGraFS about the connectivity of the fragment. Each dummy atom, with symbol "X" in ASE, represents a point of connection between two fragments. In the above example, the shape is clearly "linear". This is used to check compatibility between a fragment and a given topology. The unit of a fragment is set internally by AuToGraFS and is of no interest here. The name of a fragment is used only for logging purposes. Individual building blocks of a framework are manipulated through the `Fragment` class. It is possible to export a fragment from the GUI using `edit -> framework -> export fragment`. If the UFF atom types are not specified, an automatic typing script will take care of it.

### The Model class

This is a container class that holds in one place the topology of a framework and the correctly positioned fragments. Most of the postprocessing methods found in AuToGraFS are Model methods.

```
from scm.autografs import *

model = autografs.Model(...)

# view the framework in adfinput. set clean to False to keep dummy atoms.
model.view(clean=True, verbose=True)

# write the name.ams and name.run files
model.write(name="framework", clean=True, verbose=True)

# returns a Fragment, with correct bonding information and mmtypes
# if indices is True, it also returns the index of the corresponding fragment in the_
↳model for each atom
atoms = model.get_atoms(self, clean=False, indices=False)

# returns a supercell of the framework as a Fragment
supercell = model * (2,2,2)

# deletes a fragment in the framework. if no index is given, the fragment will be_
↳chosen at random.
# the defects will cap with hydrogen the empty spaces. returns a Fragment
# here, a linker at random will be deleted
model.insert_defect(indices=None, centers=0, linkers=1)

# flip a linear object
model.flip_fragment(index=the_fragment_index)

# rotates a linear object around its axis by angle in degrees
model.rotate_fragment(index=the_fragment_index, angle=85.0)

# Will add a functional group to the selected fragment on a random hydrogen
# if and only if the hydrogen is connected to a carbon.
# for more precise handling, use the GUI for now.
model.functionalize_fragment(functional_group="NH2", index=the_fragment_index)
```

## The Autografs class

This class is the builder in itself. Given a path to a database of building units in .inp format (exportable via the GUI), it will generate any valid framework from a topology name, a center and linker name or objects, and optionally a pillar name or object. The linkers, centers and pillars objects have to be passed as Fragment .

```

from scm.autografs import *
from random import choice

# create the generator and update the database of building units
mofgen = autografs.Autografs(refresh_database=True, verbose=True, path="path/to/my/
↳fragments/")

# choose a topology from the available one given no center or linkers
topologies = mofgen.get_available_topologies(center=None, linker=None)
topology = choice(topologies)

# choose a center in the list of available centers for the topology
centers = mofgen.get_available_centers(topology=topology, linker=None)
center = choice(centers)

# choose a linker in the list of available linkers for the topology and center
linkers = mofgen.get_available_linkers(topology=topology, center=center)
linker = choice(linkers)

# modify the linker by changing all hydrogens to fluorine
linker_structure = mofgen.database[linker]
linker_symbols = linker_structure.get_chemical_symbols()
linker_structure.set_chemical_symbols(["F" if l=="H" else l for l in linker_symbols])

# generate the framework using multiprocessing tools and automatic scaling of unit_
↳cell
framework = mofgen.make(lable="my_framework", topology=topology, center=center,
↳linker=linker, pillar=None, form_factor=None, parallel=True)

# visualize in adfinput
framework.view()

```

do not forget to run a quick UFF optimization after generation to optimize lattice and fragment into an optimal configuration.

### 7.1.2 About the databases of building units

AuToGraFS uses both a binary database format, and structures in the DemonNano file format from which the binary is compiled. To ask AuToGraFS to use a custom directory containing .inp files, simply pass the directory to the path argument when initializing the constructor. The “.inp” file format should be formatted like the following example.

```

Data: SBUtype = linker
Data: shape = linear
Data: name = benzene
GEOMETRY CARTESIAN
C      0.00000000    -1.39103633    0.00000000    MMTYPE=C_R    QMMM=MM BOND=2/1.
↳5:6/1.5:12/1.0
C      -1.20606758    -0.69440959    0.00000000    MMTYPE=C_R    QMMM=MM BOND=1/1.
↳5:3/1.5:8/1.0
C      -1.20606758    0.69440959    -0.00000000    MMTYPE=C_R    QMMM=MM BOND=2/1.
↳5:4/1.5:10/1.0

```

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C	-0.00000000	1.39103633	0.00000000	MMTYPE=C_R	QMMM=MM BOND=3/1.
↪5:5/1.5:11/1.0					
C	1.20606758	0.69440959	0.00000000	MMTYPE=C_R	QMMM=MM BOND=4/1.
↪5:6/1.5:7/1.0					
C	1.20606758	-0.69440959	-0.00000000	MMTYPE=C_R	QMMM=MM BOND=1/1.
↪5:5/1.5:9/1.0					
H	2.02806392	1.16933783	0.00000000	MMTYPE=H_	QMMM=MM BOND=5/1.0
H	-2.02806392	-1.16933783	0.00000000	MMTYPE=H_	QMMM=MM BOND=2/1.0
H	2.02806392	-1.16933783	0.00000000	MMTYPE=H_	QMMM=MM BOND=6/1.0
H	-2.02806392	1.16933783	0.00000000	MMTYPE=H_	QMMM=MM BOND=3/1.0
X	0.00000000	2.09099413	0.00000000	MMTYPE=H_	QMMM=MM BOND=4/1.0
X	0.00000000	-2.09099413	0.00000000	MMTYPE=H_	QMMM=MM BOND=1/1.0
END					

### 7.1.3 Using the overhauled Atom Typer

Both AuToGraFS and the GUI now use a new python library for the assignment of UFF types to atoms and generation of the bonding matrix. The process goes as follows:

```

from scm.autografs import atomtyper

# instantiate the typer on a readable molecule file (e.g: "mol.cif")
typer = atomtyper.MolTyper("path/of/fileToType")

# choose the UFF library. options are uff and uff4mof
library = read_db("uff4mof")

# actual typing is done here
typer.type_mol(library)

# prints the bond matrix as a numpy array
# item i, j of matrix is the bond order between atoms i and j
print typer.get_guibonds()

# print a list of UFF types in order
print typer.get_mmtypes()

# write a SCM-UFF input file with correct info
typer.structure.write(name="mytypedmol")

```

## 7.2 AuToGraFS Examples

the following examples work with all calculators of the Amsterdam Modeling Suite. Any python script using these libraries should be launched using the “\$AMSBIN/amspython” binary.

### 7.2.1 Generation of all available pillared SURMOF

```
# Required imports
from scm.autografs import *
from itertools import combinations_with_replacement

# create the generator object
mofgen = autografs.Autografs(path="./my_database/")

# the SURMOF family of frameworks is
topology = "pcu"

# get all available linkers, center, pillars for the pcu topology
centers = mofgen.get_available_centers(topology=topology)
for center in centers:
    # the pcu topology accepts up to three different linkers.
    # here, we'll use two and a pillar
    linkers = mofgen.get_available_linkers(topology=topology, center=center)
    for linker in combinations_with_replacement(linkers, 2):
        pillars = mofgen.get_available_pillars(topology=topology, center=center)
        for pillar in pillars:
            label = "{0}-{1}-{2}".format(center, linker,pillar)
            # generate this particular framework
            framework = mofgen.make(label=label,
                                   topology=topology,
                                   center=center,
                                   linker=linker,
                                   pillar=pillar)

            # write the UFF inpufile under "label.run" and "label.ams"
            framework.write()
```

### 7.2.2 Generation of a defectuous UIO-66 MOF from custom files

```
# Required imports
from scm.autografs import *
from random import gauss

# get the building units as Fragment objects
center = utils.read_inp("UIO66_center.inp")
linker = utils.read_inp("UIO66_linker.inp")

# instantiate the generator
mofgen = autografs.Autografs()

# generate the MOF
mof = mofgen.make(label="UIO66", topology="bcu", center=center, linker=linker)

# get the 3*3*3 supercell
```

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```
supercell = mof * (3,3,3)

# introduce 10 defects in a gaussian distribution around the center of the
↳supercell
# the connections are capped with hydrogen.
indices = []
mu      = len(supercell)/2
sigma   = len(supercell)/4
number_of_defects = 10
while len(indices) < number_of_defects:
    index = int(abs(gauss(mu, sigma)))
    if (index not in indices) and index < len(supercell):
        indices.append(index)
supercell.insert_defect(indices = indices)

# view resulting framework in adfinput
supercell.view()
```

### 7.2.3 Generation of conformers in the IRMOF-5

```
# Required imports
from scm.autografs import *
from itertools import combinations_with_replacement

# instantiate the generator
mofgen = autografs.Autografs()

# generate the MOF
mof = mofgen.make(label="IRMOF-5", topology="pcu", center="mof5", linker="benzene")

# get the indices of the benzenes in the mof
benzenes = mof.get_linkers()
choice_angles = [0.0, 45.0, 90.0]

# generate all possible combinations
iteration = 0
for angles in combinations_with_replacement(choice_angles, 3):
    conformer = mof.copy()
    for benzene, angle in zip(benzenes, angles):
        conformer.rotate_fragment(index=benzene, angle=angle)
    conformer.write(name="conformer-{}".format(iteration))
    iteration += 1
```



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