

Hybrid Manual

Amsterdam Modeling Suite 2025.1

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CHAPTER

INTRODUCTION

The Hybrid engine allows the user to set up multi-scale or multi-layered simulations by combining the results of a set of other AMS engines for different regions of the molecular system. The interface is like that of any other AMS engine, with a range of settings as input, and a potential energy and gradients as the main output. As such the Hybrid engine can be used in any AMS application such as geometry optimization, molecular dynamics, etc.

The Hybrid engine can be used for a wide range of multi-scale simulations, involving many different layers of the molecular system and many different levels of accuracy. The most commonly used multi-scale setup is a combined quantum mechanical / molecular mechanical (QM/MM) scheme. Such a QM/MM scheme divides the system into two different regions; (i) a QM region that is described by a QM engine such as ADF or DFTB, and (ii) an MM region that is described by an MM engine such as the ForceField engine. Within the QM/MM definition the user can choose between mechanical embedding and electrostatic embedding, and the choice determines which sub-options and which sub-engines are available.

In the following, first the broad use of the Hybrid engine for a flexible linear combination of engine results is outlined. Second, the QM/MM feature with its two available options (mechanical embedding and electrostatic embedding) is described in detail.

See also:

Tutorial: Inorganic linker in organic framework

1.1 Linear Combination of Energy Terms

Longtime users of the SCM software may know this functionality as QUILD. It can be used to set up a multi-layer computation according to the ONIOM scheme developed by Morokuma, but it has applicability beyond only layered setups. In general, this flexible and straight-forward feature provides a linear combination of results from different engines computed for different regions of the molecular system.

$$E = \sum_{i}^{N} w_i \ E^{\text{engine(i)}}(\text{region(i)})$$

The engines may be all the same, and the regions can overlap or even be identical. Each individual energy is scaled by a weight. Most commonly, the weights will sum to one (the energy is a weighted average of its components), but this is not a requirement of the engine.

There are three main reasons to use such a linear combination of energy terms: to speed up a calculation, make it more accurate, or do some state averaging.

- 1) When the molecular system is large, efficiency may be improved by a layered setup, describing only a small region of interest with an accurate engine, while interaction with the environment is described with a more efficient engine.
- 2) DFT results can sometimes be improved by applying different functionals to different (or the same) regions.

3) Experimental results may be more accurately represented by a weighted average over several spin or excited states.

In a three-layered setup (Figure: active region of interest A in yellow, immediate surroundings B in pink, and long-range environment E in blue), the ONIOM energy expression contains five different terms, and uses three different engines (engines 1, 2, 3, in decreasing order of accuracy).

$$E = E^{\operatorname{engine}(3)}(A+B+C) - E^{\operatorname{engine}(3)}(A+B) + E^{\operatorname{engine}(2)}(A+B) - E^{\operatorname{engine}(2)}(A) + E^{\operatorname{engine}(1)}(A)$$



For systems like DNA, a GGA functional may best describe the intramolecular interactions (Figure: regions A and B), while the hydrogen bonds between regions are better described with an LDA functional. The corresponding energy expression contains five terms, and uses two different engine-setups (engines 1 and 2 for GGA and LDA respectively).

$$E = E^{\operatorname{engine}(1)}(\mathbf{A}) + E^{\operatorname{engine}(1)}(\mathbf{B}) + E^{\operatorname{engine}(2)}(\mathbf{A}+\mathbf{B}) - E^{\operatorname{engine}(2)}(\mathbf{A}) - E^{\operatorname{engine}(2)}(\mathbf{B})$$



The Linear Combination of Energy Terms feature is enabled with the Energy block in the Hybrid engine input. Because of its simplicity the feature works with any combination of sub-engines.

Example

In this two-layer example, a water molecule at the DFTB level is embedded in an environment of a single water molecule at the UFF level. The energy is a linear combination of three different energy terms, with respective weights w = (1, 1, -1).

$$E = E^{\rm uff}(\rm QM+MM) + E^{\rm dftb}(\rm QM) - E^{\rm uff}(\rm QM)$$

```
Task GeometryOptimization

System

Atoms

0 -1.8782 0.0294 -0.7574 region=QM

H -0.9986 0.2961 -0.3861 region=QM

H -1.8623 -0.9560 -0.6510 region=QM
```

(continues on next page)

(continued from previous page)

```
0 0.0121 -1.3731 0.5074 region=MM
       H 0.8930 -1.7879 0.3172 region=MM
       H -0.5625 -2.1395 0.7656 region=MM
    End
End
Engine Hybrid
   Energy
      Term region=QM EngineId=DFTB
                                        factor=1.0
      Term region=* EngineId=ForceField factor=1.0
      Term region=QM EngineId=ForceField factor=-1.0
   End
   Engine DFTB
       Model GFN1-xTB
   EndEngine
   Engine ForceField
       Type UFF
    EndEngine
EndEngine
```

Observe the following

- Regions are defined as usual via the System block
- The energy expression is defined in the Energy block, with multiple occurrences of the Term subkey
- The Hybrid engine block contains subblocks of other engines that are to be used
- The (sub) engines are referred to by an *EngineId* (page 11), usually the name of the engine, in this case DFTB and ForceField.

1.2 QM/MM

The Linear Combination of Energy Terms feature is very flexible, and can combine a multitude of layers and levels of accuracy. However, the most common multi-scale scheme is a two-layer scheme, with the active region of interest described at high accuracy, and the environment, which presumably has less impact on the desired outcome, described at lower accuracy. While the definition of high and low accuracy can vary (QM/QM', MM/MM', and even MM/QM combinations are possible), most often the high accuracy description is at electronic resolution (QM), while at low-accuracy the smallest particles described are atoms (MM).



1.2.1 Mechanical embedding

The mechanical embedding option is a short-cut to a subset of the possibilities provided by the flexible Linear Combination of Energy Terms described above. In this two-layer setup, the active region A is described with accurate engine (1), while the environment E (as well as the interactions between the two regions) is described with a more efficient engine (2). A subtractive scheme is used to compute the multi-scale energy and gradients. The energy expression involves a linear combination of only three energy terms.

$$E = E^{\text{engine}(2)}(A+E) + E^{\text{engine}(1)}(A) - E^{\text{engine}(2)}(A)$$

As a result of this subtractive setup, any parameters required by engine(2) need to be provided for the full system. However, if engine(2) is the force field engine (which in QM/MM it will often be), then the corresponding energy terms are comprised of atom-pair interaction energies only. As a result, all contributions in energy-term 2 will cancel against the identical contributions in energy-term 1, which means that arbitrary parameters can be selected for region A energy contributions. The only remaining parameters of interest for engine(2) are those that describe the interaction inside region E and between regions A and E.

Example

As in the previous example, a water molecule at the DFTB level is embedded in an environment of a single water molecule at the UFF level. The computation is completely equivalent to the example for the Linear Combination of Energy Terms feature, with the simplified input as the only difference.

```
Task GeometryOptimization
System
   Atoms
       0 -1.8782 0.0294 -0.7574 region=QM
       H -0.9986 0.2961 -0.3861 region=QM
       H -1.8623 -0.9560 -0.6510 region=QM
       0 0.0121 -1.3731 0.5074 region=MM
       H 0.8930 -1.7879 0.3172 region=MM
       H -0.5625 -2.1395 0.7656 region=MM
    End
End
Engine Hybrid
    QMMM
      Embedding Mechanical
      mmEngineID ForceField
      qmEngineID DFTB
       qmRegion QM
    End
    Engine DFTB
       Model GFN1-xTB
   EndEngine
    Engine ForceField
       Type UFF
   EndEngine
EndEngine
```

The (electrostatic) interactions between regions are described only at the lowest level of accuracy, so that the electron density in region A is not polarized by the charges in region B.

Supported engines

Like the Linear Combination of Energy Terms feature, the mechanical embedding option goes beyond QM/MM in that any combination of engines is implemented (QM/QM', MM/MM', MM/QM).

1.2.2 Electrostatic embedding

Electrostatic embedding is the more common application of two-layer QM/MM, where active region A is assumed to have an electron density, and this density is polarized by the point charges in environment region E.



A purely subtractive energy expression is no longer possible, as the interaction energy is not determined by a single description (QM or MM), but by a combination of both.

$$E = E^{\mathrm{QM}}(\mathrm{A}) + E^{\mathrm{int}}(\mathrm{A}|\mathrm{E}) + E^{\mathrm{MM}}(\mathrm{E})$$

In practice, atomic point charges from the lower-level engine (usually an MM force field), are passed to the higher level QM engine, which computes the interaction of the electron density in region A with the point charges from region E.

$$E^{\text{int/elstat}}(\mathbf{A}|\mathbf{E}) = \int \rho^{\mathbf{A}}(\boldsymbol{r}) \ V^{\mathbf{E}}(\boldsymbol{r}) d\boldsymbol{r} = \int \rho^{\mathbf{A}}(\boldsymbol{r}) \sum_{i}^{N_{\mathbf{E}}} \ \frac{q_{i}^{\mathbf{E}}}{|\boldsymbol{r} - \boldsymbol{r}_{i}^{\mathbf{E}}|} d\boldsymbol{r}$$

Here $V^{\rm E}$ is the potential from the E-region point charges, $\rho_{\rm A}$ is the charge density of the A-region sub-system, which is affected by the positions of the E-region point charges, $N_{\rm E}$ is the number of E-region atoms, $q_i^{\rm E}$ is the point charge of E-region atom *i*, and $r_i^{\rm E}$ is the position of E-region atom *i*.

The remaining non electrostatic interactions are calculated at the MM level by a subtractive scheme.

$$E^{\text{int/nonelstat}}(\mathbf{A}|\mathbf{E}) = E_{\text{nonelstat}}^{\text{MM}}(\mathbf{A}+\mathbf{E}) - E_{\text{nonelstat}}^{\text{MM}}(\mathbf{A}) - E_{\text{nonelstat}}^{\text{MM}}(\mathbf{E})$$

The nonelectrostatic energy $E_{\text{nonelstat}}^{\text{MM}}$ is obtained by simply setting the MM point charges to zero.

The final QM/MM energy consists of five terms, computed with two different engines.

- 1) The MM energy of region E.
- 2) The QM energy of region A, including interaction with point charges region E.
- 3) The MM energy of full system without electrostatics.
- 4) The MM energy of region E without electrostatics.
- 5) The MM energy of region A without electrostatics.

$$E = E^{MM}(E) + E^{QM}(A(V^{E})) + E^{MM}_{\text{nonelstat}}(A+E) - E^{MM}_{\text{nonelstat}}(A) - E^{MM}_{\text{nonelstat}}(E)$$

Currently, only non-polarizable forcefields are supported, so that the MM charges q_i^E are not affected by the QM potential.

Example

Again, a water molecule at the DFTB level is embedded in an environment of a single water molecule at the UFF level, this time using electrostatic embedding.

```
Task GeometryOptimization
System
   Atoms
       0 -1.8782 0.0294 -0.7574 region=QM
       H -0.9986 0.2961 -0.3861 region=QM
       H -1.8623 -0.9560 -0.6510 region=QM
       0 0.0121 -1.3731 0.5074 region=MM
       H 0.8930 -1.7879 0.3172 region=MM
       H -0.5625 -2.1395 0.7656 region=MM
   End
End
Engine Hybrid
    QMMM
      Embedding Electrostatic
      mmEngineID ForceField
      qmEngineID DFTB
      qmRegion QM
   End
   Engine DFTB
       Model GFN1-xTB
   EndEngine
   Engine ForceField
       Type UFF
   EndEngine
EndEngine
eor
```

Supported engines

In the electrostatic embedding QM/MM setup of the Hybrid engine, the sub-engines involved can have one of two "roles": the QM role and the MM role.

Engines supporting the QM role:

- adf
- band
- dftb

Engines supporting the MM role:

forcefield

1.3 Capping Atoms

Whenever the boundary between two regions crosses a covalent bond (or better put, whenever the Hybrid engine notices that a sub-engine is assigned a system with dangling bonds), capping atoms - or link atoms - are assigned to satisfy the valence of the boundary atoms (see Figure below). By default, the engine adds hydrogen as capping atom, though a different (single) element can be selected by the user.



The capping atoms are added according to the AddRemove methodology [1 (page 103)], in which the capping atoms follow the position of the real atoms in the total system. By default, the capping atoms are positioned along the vector of the dangling covalent bond, and at a distance that corresponds to the sum of the covalent radii of the capping atom and the boundary atom to which is has been attached. An alternative option can be selected that places the capping atom again along the vector of the dangling bond, but at a distance to the connected boundary atom that is a fraction of the boundary atoms in the full system.

Capping can be disabled (via the *Energy* (page 12) or *QMMM* (page 14) block) and options can be set in the *Capping* (page 16) block. An element as well as force field atom-type and charge can be assigned to the capping atom. When more than one capping atom is present in the system, they will all have the same element, type, and charge.

1.3.1 Position of the Boundary

By default two sanity checks are performed on user-defined boundaries, and the following choices are not accepted by the engine.

- A boundary across bonds with a bond order larger than 1.25. The user can prevent this by using the AllowHigh-BondOrders subkey, see the *Capping* (page 16) documentation.
- A (QM) sub-region with more than one capping atom representing the same MM-boundary atom. The hybrid engine will not accept this, unless the user overrides it, see the CheckCapping subkey of *Capping* (page 16).

The GUI (amsinput) has an option (Make selection cappable, from the Select menu) to automatically extend a selection, so that the two checks will pass.

1.3.2 QM/MM Partitioning Examples

In a QM/MM simulation the basic question is how to partition the system into the QM and MM regions A and E. When studying an active site of a catalyst, for example, one must decide where to put the QM/MM boundary. Putting the boundary too close to the reaction center will jeopardize the chemical realism of the model. On the other hand, if one places the boundary too far away, the computational expense of the QM calculation may become problematic. Each system is different in this respect and the user must perform the proper tests to validate the appropriateness of the QM/MM partitioning used. We strongly suggest that the reader examines the literature on QM/MM methods and understands the basic limitations of the approach.

Below we give examples of the QM/MM partitioning choices that are by default prohibited by the engine. For comparison, we also give some representative examples of QM/MM partitioning that the engine does allow. In the examples, the region enclosed in the dotted polygon represents the A-region and the atoms labeled with 'LI' are the MM boundary atoms to be replaced by a capping - or link - atom.

In the examples in Figure 1, the boundary crosses double, triple, or aromatic bonds, so that a simple capping atom cannot satisfy the valence of the QM fragment and the electronic structure of the QM sub-system becomes drastically different from that of the full system.



Figure 1 Examples of partitioning that should not be used because the link bonds are double or aromatic bonds.

Next, figure 2a depicts examples of partitioning that are not allowed because the MM boundary atom has a covalent bond to more than one QM atom. An MM boundary atom can only be bonded to one QM atom. Figure 2b shows the opposite, which is allowed. In other words, one QM atom can be bonded to more than one MM boundary atom. Note that there is no limit to the number of capping atoms that can be placed, just that each MM boundary atom can only be bonded to one QM atom.

a NOT ALLOWED



Figure 2 a) Examples of partitioning that are not allowed because the MM boundary atom has a covalent bond to more than one QM atom. b) The allowed reverse of the examples shown in (a). An MM boundary atom can **only** bond to one QM atom.

Then, figure 3 provides some representative examples of partitioning that the program does allow. Example **a** shows a typical solute-solvent QM/MM partitioning where there are no link bonds at all. Example **b** depicts two separate molecules each possessing a QM and a MM region. We emphasize that any number of molecules and link bonds can be used. Example **c** seems very similar to the earlier example in Figure 1. The difference is that the ring in Figure 3cd is not aromatic and consequently the link bonds in example **d** cross *single* bonds. Example **d** shows a single molecule, with two QM regions separated by an MM region. For this example, two equivalent pedagogic representations of the sample partitioning are displayed. Example **e** is a representative organometallic complex.



Figure 3 Representative examples of QM/MM partitioning that can be used in the Hybrid engine.

1.3.3 Electrostatic Embedding

Electrostatic embedding involving capping atoms can, without additional measures, result in unphysical behavior, such as nuclear fusion or extreme repulsion. The problem is the point charge of the MM boundary atom, which is located very close to the QM capping atom. To remedy this, our default implementation sets the charge at the position of the MM boundary atom to zero, and redistributes it over the remaining MM atoms while keeping the total charge (in the MM region) constant. While this avoids plain pathological behavior, the user should note that this alters the dipole moment of the MM region, which may have consequences for the behavior of the system.

In the electrostatic embedding setup, the sub-region E that is passed to the MM-engine is by default not provided with capping atoms, since the MM engine can handle an un-capped system. To compute the electrostatic embedding energy for a system with capping atoms, the energy expression needs to be slightly adjusted.

- 1) The MM energy of region E, without capping atoms.
- 2) The QM energy of capped A-region (A_C) , including interaction with point charges region E.
- 3) The MM energy of full system without electrostatics.
- 4) The MM energy of region E without electrostatics and without capping atoms.
- 5) The MM energy of capped A-region (A_C) without electrostatics.

$$E = E^{MM}(E) + E^{QM}(A_C(V^E)) + E^{MM}_{\text{nonelstat}}(A+E) - E^{MM}_{\text{nonelstat}}(A_C) - E^{MM}_{\text{nonelstat}}(E)$$

CHAPTER

TWO

HYBRID ENGINE OPTIONS

2.1 Sub-engines and EngineIDs

Inside the Hybrid engine input block one or more sub-engine blocks can be defined. These have exactly the same format as regular engine blocks. There is, however, one extra feature: the EngineId. Optionally, a unique name can be added to the engine definition as an extra string, serving as an identifier. By default the identifier is simply the engine name. The extra string allows the user to select the same engine multiple times, each time with different settings.

This is an example, where we use the same engine (ADF) with two different spin polarizations, using results corresponding to the lowest energy.

```
Engine Hybrid
   Energy # we want results from the engine that yields the lowest energy to be this.
\leftrightarrowengine's result
       DynamicFactors UseLowestEnergy
       # The user-supplied factors are irrelevant here so we omit them
       Term EngineId=adf-singlet Region=*
       Term EngineId=adf-triplet Region=*
    End
    Engine ADF adf-singlet
                              # here adf-singlet is the EngineID
      SpinPolarization 0
      Unrestricted False
   EndEngine
   Engine ADF adf-triplet
      SpinPolarization 2
      Unrestricted True
   EndEngine
EndEngine
```

EngineIDs are for instance used in the technical example *QUILD* (page 88), which tests that EngineIDs are case-insensitive.

Engine header

Engine

Type Block

Recurring True

Description

The input for the computational (sub) engine. The header of the block determines the type of the engine. An optional second word in the header serves as the EngineID, if not present it defaults to the engine name. Currently it is not allowed to have a Hybrid engine as a sub engine.

2.2 Linear Combination of Energy Terms

The block Energy triggers a QUILD-like setup, allowing the energy to be defined as a linear combination of energy terms. Each energy term can be computed with a different engine.

See the *basic QUILD example* (page 2). As you can see capping can be enabled per energy term, and the user can set a charge per term (for the corresponding region).

```
Energy

DynamicFactors [Default | UseLowestEnergy | UseHighestEnergy]

Term

Charge float

EngineID string

Factor float

Region string

UseCappingAtoms Yes/No

End

End
```

Energy

Туре

Block

Description

This block is there to construct the energy.

DynamicFactors

Type

Multiple Choice

Default value Default

Options

[Default, UseLowestEnergy, UseHighestEnergy]

GUI name

Adjust factors

Description

Default - use factors as set in the corresponding Term blocks;

UseLowestEnergy - set all factors to 0 except for that of the engine with the lowest energy, which is set to 1;

UseHighestEnergy - set all factors to 0 except for that of the engine with the highest energy, which is set to 1.

The last two options make sense only for non-QMMM hybrid calculation (that is, if the QMMM block is not present) and only when using engines whose energies can be compared directly.

Term

Type Block

Recurring

True

Description

This block is there to construct the energy term. Can have multiple occurrences

Charge

Type Float

Default value 0.0

Description

Net charge to be used for this energy term.

EngineID

Type String

Description Identifier for the engine

Factor

Type Float

Default value 1.0

Description

Region

Type String

Description Identifier for the region

UseCappingAtoms

Туре

Bool

Default value Yes

Description

Whether to use capping for broken bonds

2.3 QM/MM

The alternative to the Energy block is the QMMM block, which triggers a two-layer computation. The embedding type can be selected with the Embedding key, for which mechanical or electrostatic can be selected. The former option triggers a specific linear combination of three energy terms, and can therefore also be set up using the Energy block (see *basic QUILD example* (page 2)).

See the *basic electrostatic embedding example* (page 5). Capping can be disabled, and charges can be set for the QM and MM regions.

```
QMMM
Embedding [Mechanical | Electrostatic]
MMCharge float
MMEngineID string
QMCharge float
QMEngineID string
QMRegion string
UseCappingAtoms Yes/No
End
```

Lina

QMMM

Туре

Block

Description

This block is there to identify the QMMM engines.

Embedding

Туре

Multiple Choice

Default value

Electrostatic

Options

[Mechanical, Electrostatic]

Description

Determines how the QM region is embedded into the MM region.

Mechanical embedding embedding can also be achieved using the Energy%Terms keywords, but the common case of a two region mechanical QM/MM embedding is easier to set up using this keyword.

MMCharge

Туре

Float

Default value 0.0

Description

Net charge to be used for the MM region.

MMEngineID

Туре

String

Description

Identifier for the MM engine

QMCharge

Type Float

Default value 0.0

Description

Net charge to be used for the QM region.

QMEngineID

Type String

Description Identifier for the QM engine

QMRegion

Type String

Description

Identifier for the QM region. The rest of the system is considered the MM region.

UseCappingAtoms

Type Bool

Default value Yes

Description

Whether to use capping for broken bonds.

2.4 Committee

It is also possible to run the hybrid engine in Committee mode. When using the hybrid engine as a committee, the average is taken of each engine, meaning that all the regions and factors must be the same. Additionally, the spread of results of the different engines is understood as an uncertainty. The standard deviation of the different engine energies and forces are computed and reported as the uncertainty.

```
Committee
Enabled Yes/No
End
```

Committee

Type Block

Description

Settings for using the hybrid engine as a committee. The factors and region for each engine must be the same. When committee is enabled the standard deviation is also reported as the uncertainty.

Enabled

Type Bool

Default value

No

Description Enable committee

2.5 Capping

Whether of not capping is enabled is set inside the Energy and QMMM blocks. If enabled then the user can influence the position and type of the capping atom with the Capping sub-block.

Capping

```
AllowHighBondOrders Yes/No
AtomicInfoForCappingAtom string
CappingElement string
CheckCapping Yes/No
Distance float
Option [Fractional | Fixed]
End
```

Capping

Туре

Block

Description

This block is about capping details. Capping occurs with hydrogen atoms when a bond is broken between an atom inside the region and one outside.

AllowHighBondOrders

Туре

Bool

Default value

No

Description

Allows capping of interregional aromatic, double and triple bonds. This is normally not a good idea, since the capping is done with hydrogen atoms.

AtomicInfoForCappingAtom

Туре

String

Default value

ForceField.Type=H_ForceField.Charge=0.0

Description

The AtomicInfo for the capping atoms. Typically a string like ForceField.Type=X much like forcefield info is entered in the System block for normal atoms.

CappingElement

Туре

String

Default value

Η

Description

The element to be used for capping. The hydrogen atom has the advantage that it is very small.

CheckCapping

Туре

Bool

Default value

Yes

Description

The same outside atom can be involved in multiple capping coordinate definitions. This is not a good idea, and this will not be accepted by using this check.

Distance

Туре

Float

Default value -1.0

Description

A negative value means automatic. In that case the sum of covalent radii is used

Option

Type Multiple Choice

Default value Fixed

Options

[Fractional, Fixed]

GUI name

Capping option

Description

The capping atom is always along the broken bond vector.

The bond distance between the capping atom and the two atoms are obtained from covalent radii, let us call them D1H and D2H.

With option=Fractional the capping is on the bond vector with the fraction D1H/(D1H+D2H).

With the Fixed option it at the distance D1H from atom 1. A distance of zero always means the coordinate of the inside atom.

For a specific application of QM/MM with capping atoms see this example (page 86).

2.6 Restarts

In a molecular dynamics run or geometry optimization, the geometries at subsequent steps are often very similar. Generally, efficiency can be gained by providing the engines with information from the previous step ("restart"), as this might speed up the SCF or charge equilibration procedure, if applicable. To the forcefield engine, this might avoid re-loading of the database, guessing bonds, etc. at every step. By default all sub-engines are provided with restart information. It can be switched off with the RestartSubEngines key.

RestartSubEngines Yes/No

RestartSubEngines

Type Bool

Default value Yes

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Description

Save all the results of the subengines and pass those in a next geometry step or MD step.

2.7 Charges per region

The user can specify charges per region associated with each energy term.

Depending on the setup the charges can be set with the Energy%Term%Charge or the QMMM%mmCharge and QMMM%mmCharge keys.

For a QM engine the charge for a region determines the number of electrons in the region defined in the energy term. For the ForceField engine, charges are specified per atom, and they should add up to the charge specified for the region.

2.7.1 Linear Combination of Energy Terms

When the Linear Combination of Energy Terms feature is selected, but using the Energy block, the energy is a linear combination of independent calculations. It is in this spirit that the total charge is considered to be

$$Q = \sum_{i}^{N} w_i \ Q^{\text{engine}(i)} \big(region(i) \big)$$

In the QMMM setup the total charge is the sum of the charge of the mm region and the qm region.

When Energy%DynamicFactors is set to anything other than Default the factors specified in the Term blocks are not used.

2.7.2 Electrostatic Embedding

In general, the charges for sub-regions should be consistent with the charge specified for the total system. However, in the case of an electrostatic embedding computation with capping atoms, the sum of charges of the subsystems used in the computation of the five energy terms is allowed to deviate from the total system charge.

$$E = E^{MM}(E) + E^{QM}(A_C(V^E)) + E^{MM}_{\text{nonelstat}}(A+E) - E^{MM}_{\text{nonelstat}}(A_C) - E^{MM}_{\text{nonelstat}}(E)$$

The MM region that will be passed to the MM engine (term 1) will often have a fractional charge, due to un-capped dangling bonds. The fractional charge of the QM-region however (term 2), should be corrected by the capping atom

charges, to yield a chemical system that optimally resembles the full system. As a result, the sub-region charges do not need to add up to the total charge of the system.

CHAPTER

THREE

PDB2ADF: TRANSFORM PDB FILE TO QMMM INPUT FILE

The pdb2adf utility program was written by Marcel Swart.

3.1 Overview

3.1.1 General description

Starting from the ADF2005.01 version the utility pdb2adf is available in the official release. Previously this utility could be found on the contributed software page. Starting from AMS2020 the default is to make an amsified ADF input, which can not be used with previous versions. One can get the old style input if the environment variable SCM_PDB2ADF is set to OLD (only to be used with ADF<=2019). One can get the NEWQMMM style input input if the environment variable SCM_PDB2ADF is set to NEW (only to be used with ADF<=2019).

The pdb2adf utility was written to read a PDB file (http://www.pdb.org), which contains the atomic coordinates of a protein structure, and transform it into an ADF inputfile, particularly for use with QMMM calculations. It can also be used for setting up a solvent shell around a solute molecule.

The PDB files are generally used for protein structures, and are formatted according to certain rules, see: http://www.wwpdb.org/docs.html, and the part about the official PDB format below.

For every residue/molecule present in the PDB file, there should be a fragment file available, either in the general AMS library (\$AMSHOME/atomicdata/pdb2adf directory), or in the local directory where the pdb2adf program is being called. Fragment files in the local directory take higher priority than those in the general AMS library. The fragment files are formatted, based loosely on AMBER parameter files, and contain information about the residues; e.g., the atoms present, with their general and forcefield atomnames, atomic charges, connections to other atoms for creating their positions when not found on the PDB file, etc.; see part about fragment files below. Available in the AMS library are fragment files for amino acid residues, including those at the N- or C-terminal residue, three solvents (water, methanol, chloroform), some ions that are present frequently in protein structures (copper, fluoride), etc.

Also present in the AMS library are solvent box files that can be used to place a layer of solvents surrounding the protein, or a solute. Available are the three solvents mentioned above.

After reading the PDB and corresponding fragment files, the program tries to figure out which atoms are missing, and will add those; it uses the information provided on the fragment files to do so. For certain amino acid residues, there are several protonation states possible, e.g. histidine can be protonated at the N-delta position, at the N-epsilon position, or on both. The default option is to choose the fully charged option for aspartate (Asp), glutamate (Glu), lysine (Lys) residues, and decide for each histidine (His) and cysteine (Cys) residue individually what the protonation state should be. In those individual cases, the distances of neighboring molecules/residues are given that may help determine the protonation state. See the protein example below.

After all that is setup properly, a list is given with residue names/numbers, from which you can choose those that should be placed in the QM system; afterwards, for each of the selected QM residues, a choice should be made where to cut-off the QM part. The most appropriate point to cut-off seems to be at the C-alpha position, except when dealing with a

proline (Pro). The latter residue is cyclic, e.g. the sidechain is connected to the C-alpha carbon ! For that residue, it may be better to include the C-alpha, H-alpha, and backbone carbonyl group of the preceding residue in the QM part.

The program will try to use to replace the ".pdb" extension of the PDB file by ".pdb2adf" for the AMS inputfile to be made; for convenience, the program also writes out an ".p2a.pdb" file with the complete system as it being made by the program. This file can then be visualized by conventional viewer programs (such as iMol, VMD, Molekel, AMSview) for visual inspection if everything has been carried out correctly.

Given below are two examples, one for the application of a protein, the other how to set up a solvent shell run.

3.1.2 Things to notice

- The QMMM implementation in AMS2020 is new. It uses the AMS Hybrid engine.
- By default pdb2adf makes an AMS Hybrid Engine input format.
- The pdb2adf program uses AMBER parameter files, and is setup to work with the AMBER force field, version AMBER95, which is designed for and works well for biosystems.
- For questions, remarks, contact: support@scm.com.

For ADF<=2019 only:

- The old style QMMM input format is used if the environment variable SCM_PDB2ADF is set to OLD.
- The NEWQMMM style QMMM input format is used if the environment variable SCM_PDB2ADF is set to NEW.

Columns	Data Type	Field	Definition
1 - 6	Record name	'ATOM' or 'HETATM'	
7 - 11	Integer	serial	Atom serial number.
13 - 16	Atom	name	Atom name.
17	Character	altLoc	Alternate location indicator.
18 - 20	Residue name	resName	Residue name.
22	Character	chainID	Chain identifier.
23 - 26	Integer	resSeq	Residue sequence number.
27	AChar	iCode	borderleft for insertion of residues.
31 - 38	Real(8.3)	Х	Orthogonal coordinates for X in Angstroms.
39 - 46	Real(8.3)	у	Orthogonal coordinates for Y in Angstroms.
47 - 54	Real(8.3)	Z	Orthogonal coordinates for Z in Angstroms.
55 - 60	Real(6.2)	occupancy	Occupancy.
61 - 66	Real(6.2)	tempFactor	Temperature factor.
73 - 76	LString(4)	segID	Segment identifier, left-justified.
77 - 78	LString(2)	element	Element symbol, right-justified.
79 - 80	LString(2)	charge	Charge on the atom.

3.1.3 Official PDB format

Typical examples from PDB-files:

[1		2		3	4	5	6	7	8
12345	6789012	345	6789012	23456789	90123456789	0123456	789012345	6789012345	6789012345	567890
ATOM	76	0	GLY	A9	6.671	55.354	35.873	1.00 14.7	5 A	
ATOM	77	Ν	ASN	A10	6.876	53.257	36.629	1.00 16.0	9 A	

(continues on next page)

									(commaca nom previous page)
ATOM	62	0	GLY A	9	6.791	55.214	35.719	1.00 15.61	4AZU 153
ATOM	63	Ν	ASN A	10	6.892	53.135	36.555	1.00 12.64	4AZU 154

The pdb2adf utility is flexible, and should be able to read most PDB files, even those with incomplete or erroneous line formats. From every ATOM/HETATM line, it tries to read:

- atom number
- atom name
- residuename
- · chain identifier
- residue number
- X,Y,Z coordinates

Hints for proper formatting:

- · always group together atoms that belong to one residue
- always give the atom name on columns 13-16
- when specifying a chain-id use only letters (or a blank)

3.1.4 Contents of fragment file

Given below is the contents of the fragment file for water. The first line is a comment line, the only important parameter is the NOCONNECT keyword, which indicates that the program should not try to make any connections to other residues/molecules. Then follow three lines, that define the orientation in space of the residue; they are not used for general fragments, but are relevant and important for amino acid residues and DNA nucleotides. Finally, for each atom in the molecule, there should be a line with its number in the fragment; its name to be used in PDB files; the AMBER forcefield atomtype; a dummy atomname; connections and coordinates (bond, angle, dihedral angle) to other atoms in the molecule that can be used to give the position of the atom if it is not present in the PDB file; the atomic charge; and after the exclamation mark (!) the connections to other atoms in this fragment, or other fragments in case of amino acid residues/DNA nucleotides. The current version does not use the latter connections yet, but the next version will probably use them.

НОН	Water m	olect	ule NC	CONNEC	CT						
1	DUMM	DU	М	0	0	0	0.0000	0.0000	0.0000		
2	DUMM	DU	М	1	0	0	1.4490	0.0000	0.0000		
3	DUMM	DU	М	2	1	0	1.5220	111.1000	0.0000		
4	0	OW	0	0	0	0	0.0000	0.0000	0.0000	-0.8340	! 🖬
⇔ 5	6										
5	H1	ΗW	Н	4	0	0	0.9572	0.0000	0.0000	0.4170	! 🖬
-→4											
6	H2	ΗW	Н	4	5	0	0.9572	104.5200	0.0000	0.4170	! 🖬
-→ 4											

3.1.5 Contents of solvent box files

The first line is a comment line, followed by a line with the total number of atoms in the solvent box and the dimensions of the box (in Angstroms); then for each atom in the box, the atom name, which must match the PDB atomname, and the Cartesian coordinates, again in Angstroms.

3.2 Usage of pdb2adf

3.2.1 Short description

The program works interactively, and should be straightforwardly to use. However, for some of the stages in the output a short description is given below.

```
P D B 2 A D F - program
version 2008.01
Written by: Marcel Swart, 2008
This program uses AMBER parameter files
see: http://amber.scripps.edu
```

Do you want a logfile to be written (Y/n) $\ensuremath{\text{?}}$

This option exists to create a logfile of what pdb2adf does. However, it should normally be used only for debugging purposes.

```
Ignoring atom on line:
ATOM 974 OH LYS A 128 -10.073 42.775 15.690 1.00 38.79 5AZU1065
```

This is a warning that the atom on that particular line is ignored, should normally occur only few times (less than ten). Depends also on how well the PDB file follows the PDB format rules.

```
Data Processed:
Nat: 2519
Nmol: 196
NChains: 1
```

Information about what has been read on the PDB file: the total number of atoms (Nat), number of molecules/residues (Nmol) and number of protein chains (Nchains).

Please wait, making connection tables

At this point, the connections between the atoms are being made by looking at atom distances. It may take a while, depending on the size of the system.

Do you want to make separate files for each chain (Y/n) ?

You have the option to make different inputfiles for different protein chains, but you can also make one inputfile for all of them together.

```
Found the following terminal amino acid residues : (C-term) 128 (N-term) 1 Do you want to use these as terminal residues (Y/n) ?
```

Info is given about the C- and N-terminal residue of each chain. Reported for making sure they are chosen correctly. Note, if the C- and N-terminal residues are connected (rarely the case probably), enter N here.

```
Multiple AMBER options for HIS :

0 Decide every time differently

1 HID Histidine Delta Hydrogen

2 HIE Histidine Epsilon Hydrogen

3 HIP Histidine E & D Hydrogens

Suggested option: 0
```

For a number of residues (His, Glu, Asp, Lys and Cys) there is more than one option available in the AMBER95 force field, depending on the protonation state (His, Glu, Asp and Lys) or the existence of a sulphur bridge/connection to a metal atom (Cys). The default is to choose a different option for the His and Cys residues, and use one option for Glu, Asp and Lys (fully charged). However, if wanted you can make a choice for all residues.

```
Multiple AMBER options for CYS
                               3 (
                                     3) :
 1 CYS
          Cysteine (SH)
          Deprotonated Cysteine (S-)
 2
     СҮМ
 3
     СҮХ
          Cystine (S-S bridge)
 Connections and Nearest Atoms for SG CYS
                                         3 SG ( P2A # 41 PDB#
                                                                  20)
     Dist P2A Nr PDB Nr Label
                                             Near
                                                     Dist P2A Nr PDB Nr
                                                                         Label
1
     1.82
              38
                  19 CB CYS
                                   3 CB
                                              1
                                                      3.79
                                                             2382
                                                                     980
                                                                         0 ...
→HOH 151 O
     2.02 461 193 SG CYS 26 SG
                                              2
                                                     3.80
                                                               22
                                                                     0 HC 🗖
2
       2
→GLN
                                              3
                                                     4.04
                                                             2391
                                                                     983 0 ..
→HOH 154 O
                                              4
                                                      4.15
                                                              509
                                                                     206
                                                                         0 . .
→GLN
       28 O
                                              5
                                                      4.18
                                                              522
                                                                       0 HA 🖵
→ PHE
       29
Suggestion: 3
```

The options for Cys3 are given, with information about the atoms bonded to the SG sulphur atom (on the left), as well as the closest five non-bonded atoms (on the right). This information may help you decide which choice to make for this particular residue. Also given (on the bottom) is the suggested choice, which is based, in this case, on the presence of a sulphur bridge.

```
Multiple AMBER options for HIS
                              46 (
                                    46) :
          Histidine Delta Hydrogen
 1
     HID
 2
     HIE
          Histidine Epsilon Hydrogen
     HIP
          Histidine E & D Hydrogens
 3
                                         46 ND1 ( P2A # 844 PDB# 347 )
 Connections and Nearest Atoms for ND HIS
    Dist P2A Nr PDB Nr Label
                                             Near Dist P2A Nr PDB Nr Label
1
    1.37 843 346 CG HIS
                                   46 CG
                                              1
                                                      2.62
                                                            2166
                                                                     0
                                                                          Н1 🗕
→MET 121
2 1.33
          846
                    349 CE HIS
                                   46 CE1
                                               2
                                                      3.23
                                                             2080
                                                                      863
                                                                          ND 🗖
→HIS 117 ND1
     2.04
3
          2318
                    959 CU CU
                                  130 CU
                                              3 HB
                                                      3.33
                                                             2163
                                                                      900
                                                                          S 🗖
→MET 121 SD
                                               4
                                                      3.40
                                                              2164
                                                                      901 CT 🖵
→MET 121 CE
                                                              2082
                                               5
                                                      3.57
                                                                      865 CE 🗖
→HIS 117 CE1
 Connections and Nearest Atoms for NE HIS
                                         46 NE2 ( P2A # 848 PDB# 350 )
     Dist P2A Nr PDB Nr Label
                                              Near
                                                      Dist P2A Nr PDB Nr Label
                                                                   (continues on next page)
```

											(continue	d from pre	evious	page)
1	1.32		846	349	CE	HIS	46	CE1	1 HB	2.70	162	67	0	
→ASN	10	0												
2	1.37		850	348	CD	HIS	46	CD2	2	2.83	814	0	Н1	_
→MET	44													
									3	3.23	2166	0	H1	_
∽ МЕТ	121													
									4	3.52	822	332	0	_
<u></u> мет	44	0												
									5	3.74	813	334	СТ	<u> </u>
∽ МЕТ	44	CG												
Sugges	stion:	2												

For His residues, the information is given for both the delta- and the epsilon nitrogen atoms. Also indicated (by HB) is the presence of a hydrogen bond with another atom. The definition used here is that two atoms are hydrogen bonded if they are both non-carbon/non-hydrogen atoms, and the distance between them is less than the sum of the van der Waals radii of the atoms. It is a simple definition, but seems to be effective. In this case, as the N(delta) is bonded to copper, the proton should be attached to the N(epsilon).

Making choice for which molecules should be QM, which MM

Now we come to the part where the division in the QM and MM systems is made.

```
Residues belonging to chain 0
Option Molecule Option Molecule
                                Option Molecule Option Molecule
→Option Molecule
   1: ALA 1
                  28: GLN 28
                                  55: ASP 55
                                                  82: ALA 82
→109: ALA 109
                                   56: LYS 56
   2: GLN 2
                  29: PHE 29
                                                   83: HIS
                                                           83
→110: TYR 110
etc
```

All molecules/residues belonging to chain 0 are given, with an option number.

Give option number of molecules to be put **in** QM region (**or** 'c' to **continue**): Note: by specifying a negative number a molecule **is** removed **from the** QM region

Here you are asked to enter the option numbers of the residues you want to put in the QM system.

Putting GLY 45 **in** QM region Putting HIS 46 **in** QM region

In this case, Gly45 and His46 have been put in the QM system.

```
Make a choice for the QM/MM treatment of GLY 45
0: Put completely in QM region
1: Cut off at C-alpha (put NH in QM region, CO in MM region)
2: Cut off at C-alpha (put NH in MM region, CO in QM region)
3: Cut off at C-alpha (put NH and CO in MM region)
4: Cut off at C-alpha (put NH and CO in QM region, sidechain in MM region)
5: Put only part of sidechain in QM region
Suggestion: 2
Give choice:
```

A choice should be made for where to cut-off the QM system. Normally this is done at the C(alpha) position, and you should simply choose the Suggestion.

So	lvent	molecules		(SOL/HOH)		belonging		to	this	chain:							
	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17.
\hookrightarrow	18	19	20														
	21	22	23	24	25	26	27	28	29	30	31	32	33	34	35	36	37
\hookrightarrow	38	39	40														
	41	42	43	44	45	46	47	48	49	50	51	52	53	54	55	56	57 .
\hookrightarrow	58	59	60														
	61	62	63	64	65	66											
Giv	ve the	e num	ber of	f the	mole	cule	to be	put	in (QM regi	on	(or 'c'	to	contin	ue):		

Also water molecules can be put in the QM system.

Box Shape options: 1 Spherical box 2 Cubic box Make a choice:

Type of box to be used.

```
Maximum atomic distance (Angs) from center 25.62
Give boxsize (def.: 28.62 Angs)
```

Size of box to be used to put a layer of solvent molecules around the system. Max. dist. is the maximal distance of any protein atom from the center of mass of the protein. Usually you should choose a boxsize at least 6 Angstrom larger (so at least two solvent molecules are surrounding the system).

•••

```
Using BOXSIZE value of 30.0000
Adding atoms for box 1 Added (Box):
                                                 0 Excl. (1): 648 Excl.
                                     0 (Total):
→ (2):
        0
Adding atoms for box 2 Added (Box):
                                     9 (Total):
                                                 9 Excl. (1): 639 Excl.
↔ (2): 0
Adding atoms for box 63 Added (Box):
                                     3 (Total): 7635 Excl. (1): 645 Excl.
↔ (2): 0
Adding atoms for box 64 Added (Box):
                                      0 (Total): 7635 Excl. (1): 648 Excl.
↔ (2): 0
Writing inputfile for chain
                         1
```

A total amount of 7635 atoms (2545 water molecules) has been added.

```
Inputfile(s) written, everything processed, work has been done.
Thank you for using the PDB2ADF program.
Normal ending of PDB2ADF program
______
```

ADF inputfile(s) have been written, the PDB-file has been processes. Everything is done.

3.2.2 An example on protein structure

The idea of this example is to make an adf-input file using a PDB of an azurin (1DYZ.pdb.txt). The result of this example should be that in the adf-input file the active site of azurin (Figure 1) is in the QM part, and the rest of the protein is in the MM part, and that the solvent water is added (in a box), which is also in the MM part.



Figure 1: the active site of azurin

Usage of pdb2adf

The program works interactively. Given below in **bold** are the parts that the user has to type. In cases where the user agrees with the suggestion given by the program, the user can press the Enter key indicated with **Enter**.

```
P D B 2 A D F - program
version 2008.01
Written by: Marcel Swart, 2008
This program uses AMBER parameter files
see: http://amber.scripps.edu
```

Please give name of PDB-file

1DYZ.pdb.txt

Do you want a logfile to be written (Y/n) ?

Enter

```
read fragments
Data Processed:
Nat: 2519
Nmol: 196
NChains: 1
Please wait, making connection tables
```

(continues on next page)

(continued from previous page)

```
Now finding nearby atoms
Assigning chain ID to all residues
Completing residues for which only option is available
Found the following terminal amino acid residues : (C-term) 129 (N-term) 1
Do you want to use these as terminal residues (Y/n) ?
```

Enter

```
Refinding nearby atoms (including atoms added {\tt in} residue completion)
```

Multiple AMBER options **for** HIS : 0 Decide every time differently 1 HID Histidine Delta Hydrogen 2 HIE Histidine Epsilon Hydrogen 3 HIP Histidine E & D Hydrogens Suggested option: 0

Enter

```
Using 0: Decide every time differently
```

```
Multiple AMBER options for GLU :

0 Decide every time differently

1 GLU Glutamic acid (COO-)

2 GLH Neutral Glutamic acid (COOH)
```

Suggested option: 1

Enter

```
Using 17 GLU Glutamic acid (COO-)

Multiple AMBER options for ASP :

0 Decide every time differently

1 ASP Aspartic acid (COO-)

2 ASH Neutral Aspartatic acid (COOH)

Suggested option: 1
```

Enter

```
Using 18 ASP Aspartic acid (COO-)

Multiple AMBER options for LYS :

0 Decide every time differently

1 LYS Charged Lysine (NH3+)

2 LYN Neutral Lysine (NH2)

Suggested option: 1
```

Enter

Using 19 LYS Charged Lysine (NH3+)

(continues on next page)

(continued from previous page)

```
Multiple AMBER options for CYS :

0 Decide every time differently

1 CYS Cysteine (SH)

2 CYM Deprotonated Cysteine (S-)

3 CYX Cystine (S-S bridge)

Suggested option: 0
```

Enter

```
Using 0: Decide every time differently
Making Choices for Chain 0
Multiple AMBER options for CYS 3 ( 3) :
1 CYS Cysteine (SH)
 2 CYM Deprotonated Cysteine (S-)
 3 CYX Cystine (S-S bridge)
 Connections and Nearest Atoms for SG CYS 3 SG ( P2A # 41 PDB# 20 )
  Dist P2A Nr PDB Nr Label Near Dist P2A Nr PDB Nr Label
   1.82 38 19 CB CYS 3 CB
1
                                  1
                                        3.79 2382 980 O 🗖
→HOH 151 O
                                  2 3.80 22 0 HC .
2 2.02 461 193 SG CYS 26 SG
⇔GLN 2
                                   3
                                       4.04 2391
                                                   983 O 🗖
→HOH 154 O
                                   4
                                              509
                                        4.15
                                                    206 O 🗖
→GLN 28 O
                                   5
                                        4.18 522 0 HA
→PHE 29
Suggestion: 3
```

Enter

```
Multiple AMBER options for CYS 26 ( 26) :
 1 CYS Cysteine (SH)
 2 CYM Deprotonated Cysteine (S-)
 3 CYX Cystine (S-S bridge)
  Connections and Nearest Atoms for SG CYS 26 SG ( P2A # 461 PDB# 193 )

        Dist
        P2A Nr
        PDB Nr
        Label
        Near
        Dist
        P2A Nr
        PDB Nr
        Label

        1.82
        458
        192
        CB
        CYS
        26
        CB
        1
        3.41
        522
        0
        HA

1
⇔PHE 29
2 2.02 41 20 SG CYS 3 SG 2 3.43 411 168 0 ...
→ASP 23 O
                                                                 3.60 2322
                                                            3
                                                                                       960 O 🗖
→HOH 131 O
                                                            4
                                                                     3.91
                                                                             403
                                                                                       169 CB 🗖
→ASP 23 CB
                                                            5
                                                                     4.15 387
                                                                                         0 HC 🗖
→VAL 22
Suggestion: 3
```

Enter

Multiple AMBER options for HIS 32 (32) :															
1	1 HID Histidine Delta Hydrogen														
2	HIE	Histidine Epsilon Hydrogen													
3	HIP	IP Histidine E & D Hydrogens													
Connections and Nearest Atoms for ND HIS 32 ND1 (P2A # 581 PDB# 244)															
Dist P2A Nr PDB Nr Label Near Dist P2A Nr PDB Nr Lab															
1	1.39	580	243	CG	HIS	32	CG	1	3.41	545	0	HC 🗖			
⇔THR	30														
2	1.33	583	246	CE	HIS	32	CE1	2	3.43	76	33	0 🖵			
→ALA	5	0													
								3	3.58	90	40	ОН 🗖			
⇔THR	6	OG1													
								4	3.99	91	0	но 🗖			
⇔THR	6														
								5	4.17	68	0	H 💶			
⇔ALA	5														
Coni	nectio	ns and N	learest A	toms	for	NE HI	S 32	NE2 (P2A	# 585	PDB# 2	247)				
	Dist	P2A Nr	PDB Nr	Lab	el			Near	Dist	P2A Nr	PDB Nr	Label			
1	1.31	583	246	CE	HIS	32	CE1	1	2.86	544	0	HC 👝			
⇔THR	30														
2	1.37	587	245	CD	HIS	32	CD2	2	3.00	545	0	HC 🗖			
⇔THR	30														
								3	3.14	1677	0	НО 💶			
⇔SER	94														
								4	3.42	542	229	CT 💶			
⇔THR	30	CG2													
								5	3.65	1676	688	ОН 🗖			
⇔SER	94	OG													
Sugge	stion:	1													

3

Multi	Multiple AMBER options for HIS 35 (35) :														
1	1 HID Histidine Delta Hydrogen														
2	HIE Histidine Epsilon Hydrogen														
3	3 HIP Histidine E & D Hydrogens														
Coni	Connections and Nearest Atoms for ND HIS 35 ND1 (P2A # 649 PDB# 271)														
	Dist	P2A Nr	PDB Nr	Labe	el			Near	Dist	P2A Nr	PDB Nr	Label			
1	1.38	648	270	CG	HIS	35	CG	1	2.46	682	0	Н 💶			
⇔GLY	37														
2	1.32	651	273	CE	HIS	35	CE1	2	2.69	1604	0	H1 💶			
⇔GLY	89														
								3	3.31	681	282	N 💶			
⇔GLY	37	N													
								4	3.56	1602	653	CT 💶			
⇔GLY	89	CA													
								5	3.67	152	0	H1 💶			
⇔ASN	10														
Coni	nectio	ns and N	earest A	toms	for	NE HI	S 35	NE2 (P2A	# 653	PDB# 2	74)				
	Dist	P2A Nr	PDB Nr	Labe	el			Near	Dist	P2A Nr	PDB Nr	Label			
1	1.33	651	273	CE	HIS	35	CE1	1 HB	2.91	822	332	0 🖬			
⇔MET	44	0													
2	1.37	655	272	CD	HIS	35	CD2	2	3.24	814	0	Н1 💶			

(continues on next page)

(continued from previous page) →MET 44 3.24 850 3 348 CD 🗖 →HIS 46 CD2 0 H1 🗕 4 3.34 1593 ⇔GLY 88 5 3.75 848 350 NE 🗕 →HIS 46 NE2 Suggestion: 2

3

Multiple AMBER options for HIS 46 (46) :															
1	HID Histidine Delta Hydrogen														
2	HIE	E Histidine Epsilon Hydrogen													
3	HIP	HIP Histidine E & D Hydrogens													
Coni	Connections and Nearest Atoms for ND HIS 46 ND1 (P2A # 844 PDB# 347)														
	Dist P2A Nr PDB Nr Label Near Dist P2A Nr PDB Nr La														
1	1.37	843	346	CG	HIS	46	CG	1	2.62	2166	0	H1 💶			
⇔MET	121														
2	1.33	846	349	CE	HIS	46	CE1	2	3.23	2080	863	ND 💶			
⇔HIS	117	ND1													
3	2.04	2318	959	CU	CU	130	CU	3 HB	3.33	2163	900	S 💶			
⇔MET	121	SD													
								4	3.40	2164	901	CT 🗖			
⇔MET	121	CE													
								5	3.57	2082	865	CE 💶			
⇔HIS	117	CE1													
Coni	nectio	ns and N	learest A	toms	for	NE HI	S 4	6 NE2 (P2A	# 848	PDB# 3	350)				
	Dist	P2A Nr	PDB Nr	Lab	el			Near	Dist	P2A Nr	PDB Nr	Label			
1	1.32	846	349	CE	HIS	46	CE1	1 HB	2.70	162	67	0 💶			
→ASN	10	0													
2	1.37	850	348	CD	HIS	46	CD2	2	2.83	814	0	H1 🖵			
⇔MET	44														
								3	3.23	2166	0	H1 🖵			
→MET	121														
								4	3.52	822	332	0 💶			
→MET	44	0													
								5	3.74	813	334	CT 🗖			
⇔MET	44	CG													
Sugges	stion:	2													

Enter

Multiple AMBER options for HIS 83 (83) : 1 HID Histidine Delta Hydrogen 2 HIE Histidine Epsilon Hydrogen 3 HIP Histidine E & D Hydrogens Connections and Nearest Atoms for ND HIS 83 ND1 (P2A # 1494 PDB# 613) Dist P2A Nr PDB Nr Label Near Dist P2A Nr PDB Nr Label 1.39 1493 612 CG HIS 83 CG 1 2.67 1317 0 HC 🗕 1 →VAL 73 2 1.33 1496 615 CE HIS 83 CE1 2 3.63 1315 542 CT 🗖 →VAL 73 CG2 3 3.74 0 HC 🖵 1310

(continues on next page)
										(cont	inued from pro	evious page)
⇔VAL	73											
								4	3.82	1316	0	HC 💶
\hookrightarrow VAL	73											
	7.0							5	3.86	1313	0	НС 🗕
⇔VAL	13											
Cont	nection	ns and N	earest A	toms	for	NE HT	S 83	NE2 (P2A	# 1498	PDB# 6	516)	
	Dist	P2A Nr	PDB Nr	Lab	el			Near	Dist	P2A Nr	PDB Nr	Label
1	1 32	1496	615	CE	HIS	83	CE1	1	3 0 9	1313	0	HC
 	73	T 1 7 0	010		1110	00	CUI	1	5.05	1010	0	
	1 3 8	1500	614	CD	UTC	63	CD2	2	3 11	1317	0	чс
ے ۲77 T	1.50	1000	014	CD	111.5	0.0	CDZ	2	J.44	1017	0	
↔ V AL	15							2	2 00	0.005	0.0.1	0
	4 5 0	<u>^</u>						3	3.88	2385	981	0 _
→НОН	152	0								4044		~ -
								4	3.93	1311	541	CT L
⇔VAL	73	CG1										
								5	4.03	1309	540	CT 🗖
⇔VAL	73	CB										
Sugges	stion:	2										

3

Multiple AMBER options for CYS 112 (112) :												
1	CYS	Cystein	e (SH)									
2	CYM	Deproto	nated Cy	stei	ne (S	5-)						
3	CYX	Cystine	ystine (S-S bridge)									
Conr	Connections and Nearest Atoms for SG CYS 112 SG (P2A # 2001 PDB# 828)											
	Dist	P2A Nr	PDB Nr	Lab	el			Near	Dist	P2A Nr	PDB Nr	Label
1	1.82	1998	827	СВ	CYS	112	CB	1	2.53	858	0	H 💶
→ASN	47											
2	2.14	2318	959	CU	CU	130	CU	2	2.65	2023	0	Н 💶
⇔PHE	114											
								3	3.00	2028	0	HC 💶
⇔PHE	114											
								4	3.29	868	0	H 💶
⇔ASN	47											
								5	3.39	2027	0	HC 💶
⇔PHE	114											
Sugges	stion:	2										

Enter

```
Multiple AMBER options for HIS 117 ( 117) :
1 HID Histidine Delta Hydrogen
 2 HIE Histidine Epsilon Hydrogen
 3 HIP Histidine E & D Hydrogens
Connections and Nearest Atoms for ND HIS 117 ND1 ( P2A # 2080 PDB# 863 )
Dist P2A Nr PDB Nr Label
1 1.37 2079 862 CG HIS 117 CG
                                     Near Dist P2A Nr PDB Nr Label
                                        1
                                               2.82 2028 0 HC _
→PHE 114
2 1.34 2082 865 CE HIS 117 CE1 2 3.23 844 347 ND -
→HIS 46 ND1
                                                              0 НА 🗖
3 1.99 2318
                 959 CU CU 130 CU
                                         3
                                               3.26 2031
⇔PHE 114
                                         4
                                                3.27 832
                                                             340 O 🗖
                                                           (continues on next page)
```

(continued from previous page) \hookrightarrow GLY 45 0 349 CE 🖵 5 3.43 846 →HIS 46 CE1 Connections and Nearest Atoms for NE HIS 117 NE2 (P2A # 2084 PDB# 866) Near Dist P2A Nr PDB Nr Label Dist P2A Nr PDB Nr Label 865 CE HIS 117 CE1 1 1.31 2082 1 2.57 209 0 H1 🖵 →MET 13 2 1.37 2086 864 CD HIS 117 CD2 2 2.65 2031 0 HA 🗕 ⇔PHE 114 3 HB 2.74 2406 988 O 🗖 →HOH 159 O 4 3.34 2030 841 CA 🗕 →PHE 114 CD1 5 3.41 204 0 H1 🗆 →MET 13 Suggestion: 2

Enter

Making Choices for Chain 1												
Completing residues with multiple options available, and solvent molecules												
Checking positions of newly added atoms												
Making choice for which molecules should be QM, which MM												
Residues belonging to chain 0												
Option Molecule Option Molecule Option Molecule _												
⇔Option Mo	olecule											
1: ALA	1	28:	GLN	28	55:	ASP	55	82:	ALA	82	-	
→109: ALA	109											
2: GLN	2	29:	PHE	29	56:	LYS	56	83:	HIS	83	-	
⇔110: TYR	110	20.	TUD	2.0	E 7 .	CT N	57	0.4 -	TUD	0.4		
3: CIS	3 111	30:	IHK	30	57:	GLN	57	84:	IHK	84	-	
↔III. FRE		31.	MFT	31	50.	אד א	5.0	85.	TVC	85		
4. GLU	112	51.	1412-1	JI	50.	АЦА	50	0	112	0.5	-	
→112. CIS 5. ALA	5	32.	HIS	32	59.	VAT.	59	86•	VAT.	86		
→113: SER	113	02.		01			0.5			00		
6: THR	6	33:	LEU	33	60:	ALA	60	87:	ILE	87		
→114: PHE	114											
7: VAL	7	34:	LYS	34	61:	THR	61	88:	GLY	88	_	
⊶115: PRO	115											
8: GLU	8	35:	HIS	35	62:	ASP	62	89:	GLY	89	_	
⇔116: GLY	116											
9: SER	9	36:	VAL	36	63:	GLY	63	90:	GLY	90		
⇔117: HIS	117											
10: ASN	10	37:	GLY	37	64:	MET	64	91:	GLU	91		
⇔118: TRP	118											
11: ASP	11	38:	LYS	38	65:	GLY	65	92:	SER	92	-	
→119: ALA	119											
12: ALA	12	39:	MET	39	66:	ALA	66	93:	ASP	93	-	
→120: MET	120											

									(cont	inued from	previous page)
13: MET	13	40:	ALA	40	67:	GLY	67	94:	SER	94	
→121: MET	121										
14: GLN	14	41:	LYS	41	68:	LEU	68	95:	VAL	95	_
→122: LYS	122										
15: TYR	15	42:	VAL	42	69:	ALA	69	96:	THR	96	
→123: GLY	123										
16: ASN	16	43:	ALA	43	70:	GLN	70	97:	PHE	97	
\leftrightarrow 124: THR	124										
17: VAL	17	44:	MET	44	71:	ASP	71	98:	ASP	98	
→125: LEU	125										
18: LYS	18	45:	GLY	45	72:	TYR	72	99:	VAL	99	
⊶126: LYS	126										
19: GLU	19	46:	HIS	46	73:	VAL	73	100:	SER	100	<u> </u>
⊶127: LEU	127										
20: ILE	20	47:	ASN	47	74:	LYS	74	101:	LYS	101	<u> </u>
↔128: GLY	128										
21: VAL	21	48:	LEU	48	75:	ALA	75	102:	ILE	102	
→129: SER	129										
22: VAL	22	49:	VAL	49	76:	GLY	76	103:	ALA	103	
⇔130: CU	130										
23: ASP	23	50:	LEU	50	77:	ASP	77	104:	ALA	104	
24: LYS	24	51:	THR	51	78:	THR	78	105:	GLY	105	
25: SER	25	52:	LYS	52	79:	ARG	79	106:	GLU	106	
26: CYS	26	53:	ASP	53	80:	VAL	80	107:	ASN	107	
27: LYS	27	54:	ALA	54	81:	ILE	81	108:	TYR	108	
Give option	number of	molecu	lles	to be	put in QM	regi	on (or	'c' to c	ontin	ue):	
Note: by sp	ecifying a	negat	ive n	umber	a molecule	e is :	removed	from th	e QM	region	

45 46 112 117 121 130

PuttingGLY45inQMregionPuttingHIS46inQMregionPuttingCYS112inQMregionPuttingHIS117inQMregionPuttingMET121inQMregionPuttingCU130inQMregion

Give option number of molecules to be put **in** QM region (**or** 'c' to **continue**): Note: by specifying a negative number a molecule **is** removed **from the** QM region

с

Make a choice for the QM/MM treatment of GLY 45

 Put completely in QM region
 Cut off at C-alpha (put NH in QM region, CO in MM region)
 Cut off at C-alpha (put NH in MM region, CO in QM region)
 Cut off at C-alpha (put NH and CO in MM region)
 Cut off at C-alpha (put NH and CO in QM region, sidechain in MM region)
 Put only part of sidechain in QM region

Suggestion: 2
Give choice:

Enter

```
Make a choice for the QM/MM treatment of HIS 46

    Put completely in QM region
    Cut off at C-alpha (put NH in QM region, CO in MM region)
    Cut off at C-alpha (put NH in MM region, CO in QM region)
    Cut off at C-alpha (put NH and CO in MM region)
    Cut off at C-alpha (put NH and CO in QM region)
    Cut off at C-alpha (put NH and CO in QM region)
    Put only part of sidechain in QM region

Suggestion: 1
Give choice:
```

Enter

```
Make a choice for the QM/MM treatment of CYS 112

    Put completely in QM region
    Cut off at C-alpha (put NH in QM region, CO in MM region)
    Cut off at C-alpha (put NH in MM region, CO in QM region)
    Cut off at C-alpha (put NH and CO in MM region)
    Cut off at C-alpha (put NH and CO in QM region)
    Cut off at C-alpha (put NH and CO in QM region)
    Put only part of sidechain in QM region

Suggestion: 3
Give choice:
```

Enter

```
Make a choice for the QM/MM treatment of HIS 117

    Put completely in QM region
    Cut off at C-alpha (put NH in QM region, CO in MM region)
    Cut off at C-alpha (put NH in MM region, CO in QM region)
    Cut off at C-alpha (put NH and CO in MM region)
    Cut off at C-alpha (put NH and CO in QM region, sidechain in MM region)
    Put only part of sidechain in QM region

Suggestion: 3
Give choice:
```

Enter

```
Make a choice for the QM/MM treatment of MET 121
0: Put completely in QM region
1: Cut off at C-alpha (put NH in QM region, CO in MM region)
2: Cut off at C-alpha (put NH in MM region, CO in QM region)
3: Cut off at C-alpha (put NH and CO in MM region)
4: Cut off at C-alpha (put NH and CO in QM region, sidechain in MM region)
5: Put only part of sidechain in QM region
Suggestion: 3
Give choice:
```

Enter

```
Make a choice for the QM/MM treatment of CU 130
0: Put completely in QM region
1: Put only part of molecule in QM region
Suggestion: 0
Give choice:
```

Enter

```
Total formal charge on molecule CU 130
                                    2.0000
Solvent molecules (SOL/HOH) belonging to this chain:
 1 2 3 4 5 6 7 8 9 10 11
18 19 20
                                                            16
                                            12
                                                 13
                                                    14 15
                                                                17_
→ 18
  21
      22 23 24 25 26 27 28 29
                                     30
                                        31
                                             32
                                                 33
                                                     34
                                                       35
                                                             36
                                                                37_
↔ 38
     39
         40
 41
     42 43 44 45 46 47 48 49 50 51 52 53 54 55 56 57.
→ 58 59 60
 61 62 63 64 65 66
Give the number of the molecule to be put in QM region (or 'c' to continue):
```

с

```
Residues belonging to chain 1
```

```
Do you want to add solvent to your system (Y/n) \ensuremath{\text{?}}
```

Enter

Solvent (box) available: 1: HOH HOH Water molecule 2: MOH MOH Methanol molecule 3: CHL Chloroform molecule

1

```
Reading contents of solvent box p2abox.HOH
Box Shape options:
1 Spherical box
2 Cubic box
Make a choice:
```

1

```
Writing inputfile for chain 0
Using total charge 1.0 and total spin 1.0
Maximum atomic distance (Angs) from center 25.62
Give boxsize (def.: 28.62 Angs)
```

30.0

```
Using BOXSIZE value of 30.0000
Adding atoms for box 1 Added (Box): 0 (Total): 0 Excl. (1): 648 Excl.
↔ (2): 0
                                      9 (Total): 9 Excl. (1): 639 Excl.
Adding atoms for box 2 Added (Box):
↔ (2): 0
Adding atoms for box 3 Added (Box):
                                      3 (Total): 12 Excl. (1):
                                                                645 Excl.
\leftrightarrow (2): 0
Adding atoms for box 4 Added (Box):
                                      0 (Total): 12 Excl. (1): 648 Excl.
↔ (2): 0
Adding atoms for box
                    5 Added (Box):
                                       6 (Total): 18 Excl. (1): 642 Excl.
                                                               (continues on next page)
```

→ (2):	:	0											
Adding	atoms	for	box	6	Added	(Box):	228	(Total):	246	Excl.	(1):	420	Excl.
→ (2): Adding	atoms	for	box	7	Added	(Box):	219	(Total):	465	Excl.	(1):	429	Excl.
→ (2) :		0											
Adding	atoms	for	box	8	Added	(Box):	9	(Total):	474	Excl.	(1):	639	Excl.
Adding	atoms	for	box	9	Added	(Box):	0	(Total):	474	Excl.	(1):	648	Excl.
↔ (2):		0	,	4.0			005	(600			100	
Adding \rightarrow (2):	atoms:	tor D	xod	10	Added	(Box):	225	(Total):	699	Excl.	(1):	423	EXCI.
Adding	atoms	for	box	11	Added	(Box):	216	(Total):	915	Excl.	(1):	432	Excl.
\rightarrow (2):	: atoms) for	hov	12	Added	(Boy) ·	6	(Total).	921	Fyel	(1) •	642	Fyel
\rightarrow (2):	:	0	DOX	12	Added	(DOA).	0	(10001).	<i>J</i> ∠⊥	LACI:	(⊥)•	210	LACT .
Adding	atoms	for	box	13	Added	(Box):	0	(Total):	921	Excl.	(1):	648	Excl.
→ (2) Adding	atoms	for	box	14	Added	(Box):	6	(Total):	927	Excl.	(1):	642	Excl.
↔ (2) :		0	,	4.5		(=)	10	(_ 1		60 G	
Adding \rightarrow (2):	atoms:	tor D	box	15	Added	(Box):	12	(Total):	939	Excl.	(1):	636	Excl.
Adding	atoms	for	box	16	Added	(Box):	0	(Total):	939	Excl.	(1):	648	Excl.
→ (2): Adding	: atoms) for	box	17	Added	(Box):	12	(Total):	9.5.1	Excl.	(1):	636	Excl.
→ (2) :	:	0	2011	- /	maada	(2011) •	10	(10001).	501	2	(-/•	000	20011
Adding	atoms.	for	box	18	Added	(Box):	210	(Total):	1161	Excl.	(1):	438	Excl.
Adding	atoms	for	box	19	Added	(Box):	219	(Total):	1380	Excl.	(1):	429	Excl.
\rightarrow (2):	:	0 5	la e	2.0	ک ما ما م	(D) -	2	(Tatal)	1202	Tree 1	(1) -	CAE	Dec a l
\rightarrow (2):	atoms	101 0	xod	20	Added	(BOX):	3	(IOLAI):	1303	EXCI.	(⊥):	645	EXCI.
Adding	atoms	for	box	21	Added	(Box):	216	(Total):	1599	Excl.	(1):	417	Excl.
→ (2): Adding	atoms	b for	box	22	Added	(Box):	381	(Total):	1980	Excl.	(1):	3	Excl.
→ (2)	26	4				(-) -		, , -					
Adding	atoms	for	box	23	Added	(Box):	261	(Total):	2241	Excl.	(1):	3	Excl.
Adding	atoms	for	box	24	Added	(Box):	183	(Total):	2424	Excl.	(1):	423	Excl.
\rightarrow (2):	4: 4:	2	hor	25	Nddod	(Dev)	100	(Totol).	2612	Eval	(1)	126	Eval
\leftrightarrow (2):	acoms : 3	3	XOG	20	Added	(BOX):	109	(IOLAI):	2013	EXCI.	(1):	420	EXCI.
Adding	atoms	for	box	26	Added	(Box):	186	(Total):	2799	Excl.	(1):	3	Excl.
→ (2): Adding	atoms	for	box	27	Added	(Box):	351	(Total):	3150	Excl.	(1):	3	Excl.
→ (2) :	29	4											
Adding	atoms	for	box	28	Added	(Box):	222	(Total):	3372	Excl.	(1):	420	Excl.
Adding	atoms	for	box	29	Added	(Box):	9	(Total):	3381	Excl.	(1):	639	Excl.
\rightarrow (2):	: atomo	for	hov	30	Addad	(Boy) .	162	(Total).	35/3	Fyel	(1) •	120	Fyel
\rightarrow (2)	: 5	7	DOX	50	Audeu	(DOA) :	TOZ	(10041).	5545	HACT.	(±)•	729	DACT .
Adding	atoms	for	box	31	Added	(Box):	219	(Total):	3762	Excl.	(1):	426	Excl.
→ (∠) Adding	atoms	for	box	32	Added	(Box):	6	(Total):	3768	Excl.	(1):	642	Excl.
\rightarrow (2)		0	1.				-		0.0.0.4			640	D
Adding \rightarrow (2):	atoms:	lor D	xod	33	Added	(Box):	6	(Total):	3//4	EXC1.	(1):	642	EXC1.

(continues on next page)

											(continued	from pre	vious page)
Adding → (2):	atoms	for	box	34	Added	(Box):	219	(Total):	3993	Excl.	(1):	426	Excl.
Adding ↔ (2):	atoms	for	box	35	Added	(Box):	216	(Total):	4209	Excl.	(1):	432	Excl.
Adding ↔ (2):	atoms	for	box	36	Added	(Box):	6	(Total):	4215	Excl.	(1):	642	Excl.
Adding ↔ (2):	atoms	for	box	37	Added	(Box):	219	(Total):	4434	Excl.	(1):	426	Excl.
Adding ↔ (2):	atoms 363	for	box	38	Added	(Box):	279	(Total):	4713	Excl.	(1):	6	Excl.
Adding → (2):	atoms	for 7	box	39	Added	(Box):	231	(Total):	4944	Excl.	(1):	0	Excl.
Adding	atoms	for	box	40	Added	(Box):	195	(Total):	5139	Excl.	(1):	432	Excl.
Adding (2) :	atoms	for	box	41	Added	(Box):	231	(Total):	5370	Excl.	(1):	414	Excl.
Adding ↔ (2):	atoms	for	box	42	Added	(Box):	324	(Total):	5694	Excl.	(1):	0	Excl.
Adding ↔ (2):	atoms 234	for	box	43	Added	(Box):	408	(Total):	6102	Excl.	(1):	6	Excl.
Adding ↔ (2):	atoms	for	box	44	Added	(Box):	204	(Total):	6306	Excl.	(1):	435	Excl.
Adding ↔ (2):	atoms	for	box	45	Added	(Box):	6	(Total):	6312	Excl.	(1):	642	Excl.
Adding ↔ (2):	atoms	for	box	46	Added	(Box):	177	(Total):	6489	Excl.	(1):	435	Excl.
Adding ↔ (2):	atoms	for	box	47	Added	(Box):	219	(Total):	6708	Excl.	(1):	429	Excl.
Adding	atoms	for	box	48	Added	(Box):	6	(Total):	6714	Excl.	(1):	642	Excl.
Adding (2) :	atoms	for	box	49	Added	(Box):	0	(Total):	6714	Excl.	(1):	648	Excl.
Adding → (2):	atoms	for	box	50	Added	(Box):	3	(Total):	6717	Excl.	(1):	645	Excl.
Adding	atoms	for	box	51	Added	(Box):	6	(Total):	6723	Excl.	(1):	642	Excl.
Adding ↔ (2):	atoms (for	box	52	Added	(Box):	0	(Total):	6723	Excl.	(1):	648	Excl.
Adding → (2):	atoms	for	box	53	Added	(Box):	9	(Total):	6732	Excl.	(1):	639	Excl.
Adding	atoms	for	box	54	Added	(Box):	222	(Total):	6954	Excl.	(1):	426	Excl.
Adding ↔ (2):	atoms	for	box	55	Added	(Box):	213	(Total):	7167	Excl.	(1):	426	Excl.
Adding	atoms	for	box	56	Added	(Box):	6	(Total):	7173	Excl.	(1):	642	Excl.
Adding	atoms	for	box	57	Added	(Box):	3	(Total):	7176	Excl.	(1):	645	Excl.
Adding (2)	atoms	for	box	58	Added	(Box):	219	(Total):	7395	Excl.	(1):	423	Excl.
Adding (2)	atoms	for	box	59	Added	(Box):	219	(Total):	7614	Excl.	(1):	429	Excl.
Adding (2)	atoms	for	box	60	Added	(Box):	6	(Total):	7620	Excl.	(1):	642	Excl.
Adding	atoms	for	box	61	Added	(Box):	0	(Total):	7620	Excl.	(1):	648	Excl.
Adding	atoms	for	box	62	Added	(Box):	12	(Total):	7632	Excl.	(1): (con	636 tinues or	Excl.

```
↔ (2):
        0
Adding atoms for box 63 Added (Box):
                                      3 (Total): 7635 Excl. (1): 645 Excl.
↔ (2):
        0
Adding atoms for box 64 Added (Box):
                                      0 (Total): 7635 Excl. (1): 648 Excl.
↔ (2):
        0
Total spin 1.0
Writing inputfile for chain
                          1
There are no atoms in this chain, ignoring it
Inputfile(s) written, everything processed, work has been done.
Thank you for using the PDB2ADF program.
_____
Normal ending of PDB2ADF program
```

Contents of the 1DYZ.pdb2adf file generated by pdb2adf

The file is not given completely, since it contains more than 9000 atoms.

```
#! /bin/sh
$AMSBIN/ams << eor</pre>
System
 Charge
       1.0
 Atoms
        -1.1930 25.6890 17.1840 region=MM ForceField.Charge=.141400
  Ν
↔ForceField.Type=N3 ! 1 ALA 1 N
       -0.3133 25.1929 17.1970 region=MM ForceField.Charge=.199700
  Н
↔ForceField.Type=H ! 2 ALA 1 H1
       -1.3738 25.1438 18.0148 region=MM ForceField.Charge=.199700
  Н
↔ForceField.Type=H ! 3 ALA 1 H2
  Н
       -1.5170 24.8559 16.7138 region=MM ForceField.Charge=.199700
↔ForceField.Type=H ! 4 ALA 1 H3
  С
        -1.4820 27.1340 16.8960 region=MM ForceField.Charge=.096200
→ForceField.Type=CT ! 5 ALA
                                1 CA
       -2.1350 27.2082 16.0264 region=MM ForceField.Charge=.088900
  Н
→ForceField.Type=HP ! 6 ALA
                                1 HA
      -2.1950 27.7860 18.0880 region=MM ForceField.Charge=-.059700
  С
→ForceField.Type=CT ! 7 ALA
                                1 CB
      -1.5602 27.7210 18.9717 region=MM ForceField.Charge=.030000
  Н
↔ForceField.Type=HC ! 8 ALA 1 HB1
  H -2.3971 28.8331 17.8627 region=MM ForceField.Charge=.030000
→ForceField.Type=HC ! 9 ALA
                                1 HB2
  Н
      -3.1350 27.2677 18.2776 region=MM ForceField.Charge=.030000
→ForceField.Type=HC ! 10 ALA
                               1 HB3
  С
       -0.1820 27.8790 16.5880 region=MM ForceField.Charge=.616300
→ForceField.Type=C ! 11 ALA
                              1 C
 0 0.8890 27.4920 17.0690 region=MM ForceField.Charge=-.572200 _
↔ForceField.Type=0 ! 12 ALA 1 0
 N -0.2890
                 28.9420 15.7940 region=MM ForceField.Charge=-.415700 _
↔ForceField.Type=N ! 13 GLN 2 N
  . . .
```

Н 11.6901	6.5638	30.5231 region=MM ForceField.Charge=.271900	
→ForceField.Type=H	! 690	GLY 45 H	
C 11.3760	8.5410	29.7530 region=QM ForceField.Charge=025200	—
⊶ForceField.Type=CT	! 691	GLY 45 CA	
Н 10.9114	9.3322	30.3413 region=QM ForceField.Charge=.069800	—
⊶ForceField.Type=H1	! 692	GLY 45 HA2	
Н 12.4602	8.6423	29.8009 region=QM ForceField.Charge=.069800	
↔ForceField.Type=H1	! 693	GLY 45 HA3	
C 10.9630	8.7450	28.3090 region=QM ForceField.Charge=.597300	_
↔ForceField.Type=C	! 694	GLY 45 C	
0 10.8510	7.7910	27.5300 region=QM ForceField.Charge=567900	_
↔ForceField.Type=0	! 695	GLY 45 O	
N 10.6890	9.9800	27.9260 region=QM ForceField.Charge=415700	—
→ForceField.Type=N	! 696	HIS 46 N	
H 10.7572	10./382	28.5898 region=QM ForceField.Charge=.2/1900	-
→ForceField.Type=H	! 69/	HIS 46 H	
C 10.2900	10.2500	26.5530 region=QM ForceField.Charge=058100	-
↔ForceField.lype=Cl	! 698	HIS 46 CA	
H IU.SSI/	9.3991	25.9240 region=QM ForceField.Charge=.136000	—
→Forcerieid.iype=Hi	: 699 10 E120	HIS 40 HA	
C 0.///U	10.5120	28.4440 region-QM Forcerieid.charge007400	
→Forcerieid.iype=ci	11 2472	HIS 40 CB	
H 0.5050	11.34/3	27.0095 region-QM forcerieid.charge050700	
	10 7532	25 /118 region=OM ForceField Charge 036700	
n 0.JZZ	10.7552	23.4116 Tegion-QM forcerteid.charge036700	
	9 3590	26 8430 region=OM ForceField Charge= 186800	
ForceField Type=CC	1 703	HIS A6 CC	-
N 8 0710	8 0910	26 3490 region=OM ForceField Charge=- 543200	
Serverield Type=NB	1 704	HIS 46 ND1	
C 7 1230	7 3010	26 8370 region=OM ForceField Charge= 163500	
ServeField.Type=CR	1 705	HIS 46 CE1	
н 7.0894	6.2496	26.5516 region=OM ForceField.Charge=.143500	
↔ForceField.Tvpe=H5	! 706	HIS 46 HE1	
N 6.3580	8.0230	27.6330 region=QM ForceField.Charge=279500	_
⊶ForceField.Type=NA	! 707	HIS 46 NE2	
н 5.5568	7.6742	28.1395 region=QM ForceField.Charge=.333900	_
⊶ForceField.Type=H	! 708	HIS 46 HE2	
C 6.8210	9.3110	27.6620 region=QM ForceField.Charge=220700	
⊶ForceField.Type=CW	! 709	HIS 46 CD2	
Н 6.3141	10.0588	28.2719 region=QM ForceField.Charge=.186200	<u>ـ</u>
→ForceField.Type=H4	! 710	HIS 46 HD2	
C 10.9790	11.4950	26.0450 region=MM ForceField.Charge=.597300	
⇔ForceField.Type=C	! 711	HIS 46 C	
C 11.0290	8.8020	20.9600 region=QM ForceField.Charge=.035000	
⊶ForceField.Type=CT	! 1648	CYS 112 CA	
Н 11.3902	9.8061	21.1823 region=QM ForceField.Charge=.048000	_
⊶ForceField.Type=H1	! 1649	CYS 112 HA	
C 10.0620	8.3640	22.0630 region=QM ForceField.Charge=736000	
⊶ForceField.Type=CT	! 1650	CYS 112 CB	
Н 9.2477	9.0845	22.1402 region=QM ForceField.Charge=.244000	—
⊶ForceField.Type=H1	! 1651	CYS 112 HB3	
Н 9.6557	7.3817	21.8218 region=QM ForceField.Charge=.244000	—
↔ForceField.Type=H1	! 1652	CYS 112 HB2	
S 10.8340	8.2410	23.7100 region=QM ForceField.Charge=736000	-
↔ForceField.Type=SH	! 1653	CYS 112 SG	
		(continu	ues on next page)

C 10.1650	3.3080	22.4340 region=QM	ForceField.Charge=058100	—
↔ForceField.Type=CT	! 1710	HIS 117 CA		
Н 9.2929	2.7403	22.7584 region=QM	ForceField.Charge=.136000	-
→ForceField.Type=H1	! 1/11	HIS 11/HA	EargeEigld Charges 007400	
C IU.I/SU	4.6030	HIS 117 CB	rorcerieid.charge007400	.
Н 11.1220	5.1220	23.1143 region=OM	ForceField.Charge=.036700	
→ForceField.Type=HC	! 1713	HIS 117 HB2		
н 9.3551	5.2459	22.9418 region=QM	ForceField.Charge=.036700	
\rightarrow ForceField.Type=HC	! 1714	HIS 117 HB3		
C 10.0160	4.3980	24.7440 region=QM	ForceField.Charge=.186800	
→ForceField.Type=CC	! 1715	HIS 117 CG		
N 9.7040	5.4090	25.6080 region=QM	ForceField.Charge=543200	
→ForceField.Type=NB	! 1716	HIS 117 ND1		
C 9.6570	4.9300	26.8540 region=QM	Forcefield.Charge=.163500	—
→roicerieid.iype-ck	: 1/1/ 5 5952	27 6851 region=OM	ForceField Charge= 1/3500	
Serverield Type=H5	1718	HTS 117 HE1	roreer retu. charge=.145500	-
N 9.9280	3.6450	26.8000 region=OM	ForceField.Charge=279500	
→ForceField.Type=NA	! 1719	HIS 117 NE2		
н 9.9617	3.0260	27.5974 region=QM	ForceField.Charge=.333900	_
→ForceField.Type=H	! 1720	HIS 117 HE2	-	
C 10.1580	3.2710	25.4990 region=QM	ForceField.Charge=220700	
\rightarrow ForceField.Type=CW	! 1721	HIS 117 CD2		
Н 10.3982	2.2340	25.2644 region=QM	ForceField.Charge=.186200	
↔ForceField.Type=H4	! 1722	HIS 117 HD2		
C 6.0350	6.2800	19.5280 region=QM	ForceField.Charge=023700	
→ForceField.Type=CT	! 1778	MET 121 CA		
H 4.9702	6.5113	19.5559 region=QM	Forcefield.Charge=.088000	-
G 6 6730	: 1779 6 7710	20 8330 region=0M	ForceField Charge= 03/200	
→ForceField.Type=CT	1780	MET 121 CB	roreer retu. charge=.034200	
н 7.7511	6.6157	20.7919 region=OM	ForceField.Charge=.024100	
→ForceField.Type=HC	! 1781	MET 121 HB2		
н 6.4641	7.8329	20.9631 region=QM	ForceField.Charge=.024100	<u> </u>
\hookrightarrow ForceField.Type=HC	! 1782	MET 121 HB3		
C 6.1560	6.0500	22.0720 region=QM	ForceField.Charge=.001800	
↔ForceField.Type=CT	! 1783	MET 121 CG		
Н 5.0693	6.1257	22.1101 region=QM	ForceField.Charge=.044000	—
→ForceField.Type=H1	! 1784	MET 121 HG2		
H 6.4453	5.0000	22.0292 region=QM	ForceField.Charge=.044000	—
G 6 7760	6 6970	23 61/0 region=0M	ForceField Charge=- 273700	
-ForceField Type=S	1786	MET 121 SD	rorcerieia.charge= .273700	
C 6.0690	8.3070	23.6050 region=OM	ForceField.Charge=053600	
→ForceField.Tvpe=CT	! 1787	MET 121 CE		
н 4.9825	8.2271	23.5709 region=QM	ForceField.Charge=.068400	_
→ForceField.Type=H1	! 1788	MET 121 HE1		
Н 6.3654	8.8396	24.5086 region=QM	ForceField.Charge=.068400	
→ForceField.Type=H1	! 1789	MET 121 HE2		
Н 6.4202	8.8537	22.7299 region=QM	ForceField.Charge=.068400	-
→ForceField.Type=H1	! 1790	MET 121 HE3		
CU 9.5640	/.3450	25.1/50 region=QM	ForceField.Charge=2.000000	-
↔rorcerieid.Type=CU	: 1915	CU 130 CU		
··· 0 31 1328	34 4612	22 6903 region=MM	ForceField Charge=- 834000	
→ForceField.Type=OW	! 9746	НОН 2545 О	1010011014. Onarge .001000	_

```
31.8908 34.5740 22.1167 region=MM ForceField.Charge=.417000
   Η
↔ForceField.Type=HW ! 9747 HOH 2545 H1
  Н
        30.6706 35.2981 22.6446 region=MM ForceField.Charge=.417000
                                                                         ↔ForceField.Type=HW ! 9748 HOH 2545 H2
 End
 BondOrders
      1
             5 1.0
            2 1.0
       1
       1
             3 1.0
             4 1.0
       1
       5
             7 1.0
       5
           11 1.0
       5
            6 1.0
       7
            8 1.0
       7
             9 1.0
       7
           10 1.0
           12 1.0
      11
           13 1.0
      11
           15 1.0
      13
      13
            14 1.0
      15
            17 1.0
            28 1.0
      15
      15
            16 1.0
      17
            20 1.0
      17
           18 1.0
      17
           19 1.0
     . . .
    9746
         9747 1.0
    9746 9748 1.0
 End
End
Task GeometryOptimization
GeometryOptimization
MaxIterations 100
 Convergence Gradients=1e-3
End
Engine Hybrid
 QMMM QMRegion=QM QMEngineID=ADF MMEngineID=ForceField
 Capping
  AtomicInfoForCappingAtom ForceField.Type=H1 ForceField.Charge=0.0
 End
 Engine ADF
   Title QM/MM calculation setup by pdb2adf: M.Swart et al., 2020
   Symmetry NOSYM
   Eprint
     SFO NOEIG NOOVL
   End
   XC
     GGA BP86
   End
```

```
Basis
     type TZP
     core small
   End
   SCF
     diis ok=0.01
     Converge 1.0e-5 1.0e-5
     Iterations 99
   End
   Unrestricted
   SpinPolarization
                     1.0
 EndEngine
 Engine ForceField
   Type Amber95
   ForceFieldFile $AMSHOME/atomicdata/ForceFields/amber95.ff
 EndEngine
EndEngine
eor
```

3.2.3 An example on solvent shell run

The idea of this example is to make an adf-input file using a PDB file of water (hoh.pdb.txt), in the solvent methanol. The water molecule in the adf-input file should be in the QM part, and the solvent methanol (in a box) is in MM part.

Contents of the hoh.pdb file

```
TITLE
       PDB-FILE CORRESPONDING TO pdb2adf-GENERATED ADF-INPUTFILE
REMARK
       Written by M. Swart, March 2005
HETATM
      1 H1 HOH 1 1.716 26.282 11.239 1.00 0.00
                                                              1DYZ H
                            2.439 25.795 11.634 1.00 0.00
        20
              HOH
                     1
                                                              1DYZ O
HETATM
HETATM 3 H2 HOH
                     1
                           3.140 26.440 11.729 1.00 0.00
                                                              1DYZ H
END
```

Usage of pdb2adf

The program works interactively. Given below in **bold** are the parts that the user has to type. In cases where the user agrees with the suggestion given by the program, the user can press the Enter key indicated with **Enter**.

```
P D B 2 A D F - program
version 2008.01
Written by: Marcel Swart, 2008
This program uses AMBER parameter files
see: http://amber.scripps.edu
Please give name of PDB-file
```

hoh.pdb.txt

```
Do you want a logfile to be written (Y/n) ?
```

Enter

```
read fragments
Data Processed:
   a Processed:
Nat: 3
                 1
   Nmol:
NChains:
                  0
Please wait, making connection tables
Now finding nearby atoms
Assigning chain ID to all residues
Completing residues for which only option is available
Refinding nearby atoms (including atoms added in residue completion)
          Making Choices for Chain
                                      0
      _ _ _ _ _ _ _ _ _ _ _ _ _ _ _
Completing residues with multiple options available, and solvent molecules
Checking positions of newly added atoms
Making choice for which molecules should be QM, which MM
Residues belonging to chain 0
Solvent molecules (SOL/HOH) belonging to this chain:
   1
Give the number of the molecule to be put in QM region (or 'c' to continue):
```

1

Putting HOH 1 in QM region

Give the number of the molecule to be put in QM region (or 'c' to continue):

с

Do you want to add solvent to your system (Y/n) ?

Enter

Solvent (box) available: 1: HOH HOH Water molecule 2: MOH MOH Methanol molecule 3: CHL CHL Chloroform molecule

2

```
Reading contents of solvent box p2abox.MOH
Box Shape options:
1 Spherical box
2 Cubic box
Make a choice:
```

1

```
Writing inputfile for chain 0
Using total charge 0.0 and total spin 0.0
Maximum atomic distance (Angs) from center 0.92
Give boxsize (def.: 15.00 Angs)
```

14.0

Using BOXSIZE value of	14.	0000								
Adding atoms for box	1	Added	(Box):	84	(Total):	84	Excl.	(1):	660	Excl.
↔ (2): 6										
Adding atoms for box	2	Added	(Box):	102	(Total):	186	Excl.	(1):	642	Excl.
↔ (2): 6										
Adding atoms for box	3	Added	(Box):	102	(Total):	288	Excl.	(1):	642	Excl.
↔ (2): 6										
Adding atoms for box	4	Added	(Box):	108	(Total):	396	Excl.	(1):	642	Excl.
↔ (2): 0										
Adding atoms for box	5	Added	(Box):	120	(Total):	516	Excl.	(1):	630	Excl.
↔ (2): 0										
Adding atoms for box	6	Added	(Box):	96	(Total):	612	Excl.	(1):	654	Excl.
↔ (2): 0										
Adding atoms for box	7	Added	(Box):	108	(Total):	720	Excl.	(1):	642	Excl.
↔ (2): 0										
Adding atoms for box	8	Added	(Box):	102	(Total):	822	Excl.	(1):	642	Excl.
↔ (2): 6										
Inputfile(s) written, e	every	thing p	processed	, wor	rk has been	done	•			
Thank you for using the	e PDB2	2ADF pr	ogram.							
Normal ending of PDB2ADF program										

Contents of the hoh.pdb2adf file generated by pdb2adf

The file is not given completely, since it contains more than 800 atoms.

```
#! /bin/sh
$AMSBIN/ams << eor
System
 Charge 0.0
 Atoms
       2.4390 25.7950 11.6340 region=QM ForceField.Charge=-.834000
  0
↔ForceField.Type=OW ! 1 HOH 1 O
 H 1.7160 26.2820 11.2390 region=QM ForceField.Charge=.417000
↔ForceField.Type=HW ! 2 HOH 1 H1
  H 3.1400 26.4400 11.7290 region=QM ForceField.Charge=.417000
↔ForceField.Type=HW ! 3 HOH 1 H2
  C -10.0667 22.2493 11.7437 region=MM ForceField.Charge=.116600
                                                                ____
↔ForceField.Type=CT ! 4 MOH 1 C1
  H -10.2077 21.5053 10.9597 region=MM ForceField.Charge=.037200
                                                                ↔ForceField.Type=H1 ! 5 MOH 1 HC1
  H -10.5047 21.8683 12.6667 region=MM ForceField.Charge=.037200
↔ForceField.Type=H1 ! 6 MOH 1 HC2
```

Н -10.5167	23.2103	11.4977 region=MM	ForceField.Charge=.037200	_
→ForceField.Type=H1	! 7	MOH 1 HC3		
U -8.7387	22.3983	MOH 1 01	ForceFleId.Charge=649/00	-
H -8.3007	22.6943	11.2607 region=MM	ForceField.Charge=.421500	
→ForceField.Type=HC	! 9	MOH 1 HO1	1010011010.0nargo .121000	
C -0.2827	19.0253	2.2847 region=MM	ForceField.Charge=.116600	_
←ForceField.Type=CT	! 10	MOH 2 C1		
н -0.5357	18.2063	2.9567 region=MM	ForceField.Charge=.037200	_
↔ForceField.Type=H1	! 11	MOH 2 HC1		
Н 0.7633	19.2913	2.4407 region=MM	ForceField.Charge=.037200	-
\rightarrow ForceField.Type=HI	: 14 10 8753	2 5107 region=MM	ForceField Charge 037200	
→ForceField.Tvpe=H1	! 13	MOH 2 HC3	roreer reru.enarge=.037200	
0 -0.4997	18.6373	0.9467 region=MM	ForceField.Charge=649700	
→ForceField.Type=OH	! 14	МОН 2 01	-	
Н 0.1123	17.9313	0.7287 region=MM	ForceField.Charge=.421500	<u> </u>
⊶ForceField.Type=HC	! 15	MOH 2 HO1		
C 6.1721	28.5021	18.9485 region=MM	ForceField.Charge=.116600	<u> </u>
→ForceField.lype=Cl	27 9/31	MOH 137 CI 18 8355 region=MM	ForceField Charge 037200	
ForceField Type=H1	27.9431 I 821	MOH 137 HC1	roicerieid.chaige037200	<u> </u>
н 6.3621	29.4771	19.3985 region=MM	ForceField.Charge=.037200	_
→ForceField.Type=H1	! 822	MOH 137 HC2		
н 5.4711	27.9401	19.5645 region=MM	ForceField.Charge=.037200	<u> </u>
→ForceField.Type=H1	! 823	MOH 137 HC3		
0 5.5611	28.7181	17.7095 region=MM	ForceField.Charge=649700	<u> </u>
→ForceField.Type=OH	! 824	MOH 137 01	Ferrestield Charges 421500	
H J.2031	27.0021	17.3935 region=MM	ForceFleid.Charge=.421500	
End	: 025	MOII 157 IIO1		
BondOrders				
1 2 1.0				
1 3 1.0				
4 5 1.0				
4 6 1.0				
4 7 1.0				
8 9 1.0				
10 11 1.0				
10 12 1.0				
10 13 1.0				
10 14 1.0				
14 15 1.0				
•••				
820 823 1 0				
820 824 1.0				
824 825 1.0				
End				
End				
Task GeometryOptimiz	ation			

```
GeometryOptimization
 MaxIterations 100
 Convergence Gradients=1e-3
End
Engine Hybrid
 QMMM QMRegion=QM QMEngineID=ADF MMEngineID=ForceField
 Capping
   AtomicInfoForCappingAtom ForceField.Type=H1 ForceField.Charge=0.0
 End
 Engine ADF
   Title QM/MM calculation setup by pdb2adf: M.Swart et al., 2020
   Symmetry NOSYM
   Eprint
     SFO NOEIG NOOVL
   End
   ХC
     GGA BP86
    End
   Basis
     type TZP
     core small
   End
   SCF
     diis ok=0.01
     Converge 1.0e-5 1.0e-5
     Iterations 99
   End
  EndEngine
 Engine ForceField
   Type Amber95
   ForceFieldFile $AMSHOME/atomicdata/ForceFields/amber95.ff
 EndEngine
EndEngine
eor
```

CHAPTER

FOUR

EXAMPLES

Examples of the Hybrid engine are scattered over the examples directory. Here we pick a few.

We do not repeat here all functionality that is available from the AMS driver level, see the AMS Examples.

4.1 Example: QMMM with various forcefields

Download qmmm_water.run

```
#!/bin/sh
# This example shows you how you can use the forcield engine in a qmmm setup
# Both the regions and the atom typing and charges (if any) go via te AMS system block
\# UFF
# ===
AMS_JOBNAME=uff $AMSBIN/ams <<eor
Task GeometryOptimization
System
   Atoms
       0 -1.8782 0.0294 -0.7574 region=QM
       H -0.9986 0.2961 -0.3861 region=QM
       H -1.8623 -0.9560 -0.6510 region=QM
       0 0.0121 -1.3731 0.5074 region=MM
       H 0.8930 -1.7879 0.3172 region=MM
       H -0.5625 -2.1395 0.7656 region=MM
   End
End
Engine Hybrid
   QMMM qmRegion=QM qmEngineID=DFTB mmEngineID=ForceField
   Engine DFTB
    Model GFN1-xTB
   EndEngine
   Engine ForceField
       Type UFF
   EndEngine
EndEngine
```

```
(continued from previous page)
eor
# Amber
# =====
AMS_JOBNAME=amber $AMSBIN/ams <<eor
Task GeometryOptimization
System
  Atoms
     0 -1.8782 0.0294 -0.7574 region=QM ForceField.Charge=-0.8340 ForceField.
⇔Type=OW
      H-0.9986 0.2961 -0.3861 region=QM ForceField.Charge=0.4170 ForceField.
→Type=HW
       H -1.8623 -0.9560 -0.6510 region=QM ForceField.Charge=0.4170 ForceField.
→Type=HW
       0 0.0121 -1.3731 0.5074 region=MM ForceField.Charge=-0.8340 ForceField.
→Type=OW
       H 0.8930 -1.7879 0.3172 region=MM ForceField.Charge=0.4170 ForceField.
→Type=HW
       H -0.5625 -2.1395 0.7656 region=MM ForceField.Charge=0.4170 ForceField.
\hookrightarrowType=HW
   End
   BondOrders
       1 2 1.0
        1 3 1.0
       4 5 1.0
        4 6 1.0
   End
End
Engine Hybrid
   QMMM qmRegion=QM qmEngineID=DFTB mmEngineID=ForceField
   Engine DFTB
      Model GFN1-xTB
   EndEngine
   Engine ForceField
       Type Amber95
       ForceFieldFile $AMSHOME/atomicdata/ForceFields/amber95.ff
   EndEngine
EndEngine
eor
# Tripos
# =====
AMS_JOBNAME=tripos $AMSBIN/ams <<eor
Task GeometryOptimization
System
   Atoms
       0 -1.8782 0.0294 -0.7574 region=QM ForceField.Charge=-0.8340 ForceField.
                                                                        (continues on next page)
```

```
(continued from previous page)
\leftrightarrowType=0.3
       H -0.9986 0.2961 -0.3861 region=QM ForceField.Charge=0.4170 ForceField.
⇔Type=H
       H -1.8623 -0.9560 -0.6510 region=QM ForceField.Charge=0.4170 ForceField.
⇔Type=H
       0 0.0121 -1.3731 0.5074 region=MM ForceField.Charge=-0.8340 ForceField.
\rightarrowType=0.3
       H 0.8930 -1.7879 0.3172 region=MM ForceField.Charge=0.4170 ForceField.
⇔Type=H
      H -0.5625 -2.1395 0.7656 region=MM ForceField.Charge=0.4170 ForceField.
⇔Type=H
   End
   BondOrders
       1 2 1.0
        1 3 1.0
        4 5 1.0
        4 6 1.0
   End
End
Engine Hybrid
   QMMM qmRegion=QM qmEngineID=DFTB mmEngineID=ForceField
   Engine DFTB
      Model GFN1-xTB
   EndEngine
   Engine ForceField
       Type Tripos5.2
       ForceFieldFile $AMSRESOURCES/ForceFields/tripos5.2.ff
   EndEngine
EndEngine
eor
```

4.2 Example: Mechanical embedding QUILD

Download QUILD_water.run

#!/bin/sh
This example shows you how you can use the forcefield with mechanical embedding
AMS_JOBNAME=uff \$AMSBIN/ams < <eor< td=""></eor<>
Task GeometryOptimization
System
Atoms
0 -1.8782 0.0294 -0.7574 region=QM
H -0.9986 0.2961 -0.3861 region=QM
H -1.8623 -0.9560 -0.6510 region=QM
0 0.0121 -1.3731 0.5074 region=MM
H 0.8930 -1.7879 0.3172 region=MM

```
H -0.5625 -2.1395 0.7656 region=MM
   End
End
Engine Hybrid
   Energy
      Term region=QM EngineId=DFTB
                                       factor=1.0
      Term region=* EngineId=ForceField factor=1.0
      Term region=QM EngineId=ForceField factor=-1.0
   End
   Engine DFTB
     Model GFN1-xTB
   EndEngine
   Engine ForceField
       Type UFF
   EndEngine
EndEngine
eor
```

4.3 Example: Hybrid engine with charged regions

Download HybridWithCharges.run

```
#!/bin/sh
# not needed, just slighly faster
export NSCM=1
report=report.txt
printf "Here we treat H3O+ as qm and OH- as the MM region (Optimizing without regions_
→gives two H2O molecules)\n" > $report
printf "We do this with both mechanical and electrostatic embedding\n" >> $report
printf "\n%15s %10s %10s %10s\n" "embedding" "charge" "d(0-0)" "charges" >> $report
for charge in 0.0 1.0
do
if [ "$charge" = "0.0" ]; then
chargeOinOHm=0.0
chargeHinOHm=0.0
chargeOinH3Op=0.0
chargeHinH3Op=0.0
else
chargeOinOHm=-1.123
chargeHinOHm=0.123
chargeOinH3Op=-0.5
chargeHinH3Op=0.5
fi
```

```
export AMS_JOBNAME=quild.charge=$charge
rm -rf $AMS_JOBNAME.results
"$AMSBIN/ams" << eor
Task GeometryOptimization
Properties Charges=yes
GeometryOptimization
Convergence Gradients=1.0e-6
End
System
   Atoms
       0 -1.527946410885647 -0.2107366711137158 -0.0008116899510243671 region=QM
↔ ForceField.Charge=$chargeOinH3Op
      H -0.8459142126057956 0.3517312394359257 0.4094504676540848
                                                                   region=QM _
↔ ForceField.Charge=$chargeHinH3Op
       H -1.834953147575289 0.1051014241823828 -0.8704652381864062
                                                                   region=OM ...
→ForceField.Charge=$chargeHinH3Op
       H -1.328032016244278 -1.164422847242489 0.02894848344144469
                                                                   region=QM 🔒
→ForceField.Charge=$chargeHinH3Op
      0 0.6370858511871781 -0.3378071707560572 -0.0006181020627287671 region=MM

→ForceField.Charge=$chargeOinOHm
       H 1.318474396634582 0.2241299231185073 0.4092568796869673
                                                                   region=MM 🔒
↔ForceField.Charge=$chargeHinOHm
   End
   GuessBonds True
End
Engine Hybrid
   Energy
      Term Factor=1.0 Region=* EngineID=ForceField
      Term Factor=-1.0 Region=QM EngineID=ForceField
                                                     Charge=$charge
      Term Factor=1.0 Region=QM EngineID=DFTB
                                                      Charge=$charge
   End
   Engine DFTB
       Model GFN1-xTB
   EndEngine
   Engine ForceField
   EndEngine
EndEngine
eor
ddd=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#1#5`
eee=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -k AMSResults%Charges#5.3f`
→$eee >> $report
export AMS_JOBNAME=qmmm.charge=$charge
```

```
rm -rf $AMS_JOBNAME.results
"$AMSBIN/ams" << eor
Properties Charges=yes
Task GeometryOptimization
GeometryOptimization
 Convergence Gradients=1.0e-6
End
System
   Atoms
       0 -1.527946410885647 -0.2107366711137158 -0.0008116899510243671 region=QM
                                                                                 —
→ForceField.Charge=$chargeOinH3Op
       H -0.8459142126057956 0.3517312394359257 0.4094504676540848
                                                                     region=QM
→ForceField.Charge=$chargeHinH3Op
       H -1.834953147575289 0.1051014241823828 -0.8704652381864062
                                                                     region=QM
                                                                                 ↔ ForceField.Charge=$chargeHinH3Op
       H -1.328032016244278 -1.164422847242489 0.02894848344144469
                                                                     region=OM
                                                                                 —
→ForceField.Charge=$chargeHinH3Op
       0 0.6370858511871781 -0.3378071707560572 -0.0006181020627287671 region=MM
→ForceField.Charge=$chargeOinOHm
       H 1.318474396634582 0.2241299231185073 0.4092568796869673
                                                                     region=MM
                                                                                 ____
↔ForceField.Charge=$chargeHinOHm
   End
   GuessBonds True
End
Engine Hybrid
   QMMM QMRegion=QM QMEngineID=DFTB MMEngineID=ForceField QMCharge=$charge MMCharge=-
⇔$charge
   Engine DFTB
      Model GFN1-xTB
   EndEngine
   Engine ForceField
   EndEngine
EndEngine
eor
ddd=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#1#5`
eee=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -k AMSResults%Charges#5.3f
⇔$ddd $eee >> $report
done
printf "\n* Using charges shortens the O-O distance\n" >> $report
printf "* In this case the results (mechanical vs. electrostatic) are quite similar.
\hookrightarrow as apparently the OH does not polarize the QM region much\n" >> $report
```

```
(continues on next page)
```

```
echo "start of report"
cat $report
echo "end of report"
report=report2.txt
printf "\nNow we add an extra OH- to the mm region and get a total charge of -1\n" >
⇔$report
printf "We do this with mechanical and electrostatic embedding\n" >> $report
printf "We look at two distances: d(01-05) and d(01-07) \n" >> $report
printf "Atom 01 is in the H3O+ and atoms 05 and 07 are in the two OH- molecules\n" >>
printf "\n%15s %10s %10s %10s %10s \n" "embedding" "optim" "d(1-5)" "d(1-7)"

→ "energy" >> $report

charge=1.0
chargeOinOHm=-1.123
chargeHinOHm=0.123
chargeOinH3Op=-0.5
chargeHinH3Op=0.5
for embedding in mechanical electrostatic
do
for optim in FIRE # Quasi-Newton
do
export AMS_JOBNAME=embedding=$embedding.optim=$optim
rm -rf $AMS_JOBNAME.results
"$AMSBIN/ams" << eor
Task GeometryOptimization
Properties Charges=yes
GeometryOptimization
 Method $optim
 MaxIterations 3000
 Convergence Gradients=1.0e-6
End
System
   Atoms
       0 0.9019652567984636 -1.133079116834755 0.01338426553857459 region=QM
→ForceField.Charge=$chargeOinH3Op
       H 0.1122251167578682 -1.036551903399635 0.5668491423154995
                                                                      region=QM 🔒
←ForceField.Charge=$chargeHinH3Op
```

```
(continued from previous page)
       H 1.037136681303829 -0.2320347366030556 -0.3773644469587724
                                                                       region=QM 🔒
→ForceField.Charge=$chargeHinH3Op
       H 1.678241221654873 -1.266912785246295 0.5779953693196539
                                                                       region=QM 🔒
←ForceField.Charge=$chargeHinH3Op
       0 -1.130580450693341 0.6009421414132099 -0.02453852439122078 region=MM -
→ForceField.Charge=$chargeOinOHm
       H -1.671378074377012 1.410809444490273 -0.2141830902463049
                                                                        region=MM 🗕
↔ForceField.Charge=$chargeHinOHm
       O 3.346891191122751 −0.05485781804516161 0.01059240308504993 region=MM _
→ForceField.Charge=$chargeOinOHm
       H 4.099773764135065 0.5660034244354222 -0.1683355405307263
                                                                       region=MM 🔒
↔ForceField.Charge=$chargeHinOHm
   End
   BondOrders
        1 3 1.0
        1 4 1.0
        2 1 1.0
        5 6 1.0
         7 8 1.0
   End
   Charge -1.0
End
Engine Hybrid
   QMMM qmRegion=QM qmCharge=1.0 mmCharge=-2.0 qmEngineID=dftb mmEngineID=forcefield.
→Embedding=$embedding
   Engine DFTB
       Model GFN1-xTB
   EndEngine
    Engine ForceField
   EndEngine
EndEngine
eor
d15=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#1#5`
d17=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#1#7`
eee=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -k "AMSResults%Energy"`
printf "%15s %15s %10s %10s %10.4f\n" $embedding $optim $d15 $d17 $eee >> $report
done
done
printf "\n* Very flat PES as function of these two distances\n" >> $report
printf "\n* Electrostatic embeddiding gives a bit shorter distances\n" >> $report
echo "start of report"
cat $report
echo "end of report"
```

4.4 Example: Loading MM charges for regions

In this example we consider an OH- with an H3O+ fragment. As the charges on the fragments are kept fixed, the formation of two water molecules is avoided.

First we "estimate" the charges for the two fragments with a DFTB calculation.

These charges are then loaded for the correct regions in the total system. Observe that this is done in the System block, see the System definition section of the AMS manual.

We do this first for a QUILD-like setup (mechanical embedding), and next for a QMMM calculation with electrostatic coupling.

Download LoadCharges.run

```
#!/bin/sh
# Here we treat H3O+ as qm and OH- as the MM region (Optimizing without regions gives_
→two H2O molecules)
# We do this with a QUILD setup (mechanical embedding) and electrostatic embedding_
\leftrightarrow (OMMM)
# We obtain the charges from a DFTB calculation
# In this case the results (QUILD vs. QMMM) are quite similar as apparently the OH_
\rightarrow does not polarize the QM region much
report=report.txt
echo "method distance charges" > $report
# first we do two DFTB calculations on the two fragments
export AMS_JOBNAME=H2O+.dftb
rm -rf $AMS_JOBNAME.results
"$AMSBIN/ams" << eor
Task SinglePoint
Properties Charges=yes
GeometryOptimization
 Convergence Gradients=1.0e-6
End
System
   Atoms
        0 -1.527946410885647 -0.2107366711137158 -0.0008116899510243671
        H -0.8459142126057956 0.3517312394359257 0.4094504676540848
        H -1.834953147575289 0.1051014241823828 -0.8704652381864062
        H -1.328032016244278 -1.164422847242489 0.02894848344144469
   End
   Charge 1.0
    GuessBonds True
End
Engine DFTB
```

```
Properties Charges=yes
```

"\$AMSBIN/ams" << eor

Task SinglePoint

export AMS_JOBNAME=OH-.dftb

rm -rf \$AMS_JOBNAME.results

EndEngine

eor

```
GeometryOptimization
Convergence Gradients=1.0e-6
End
System
   Atoms
       0 0.6370858511871781 -0.3378071707560572 -0.0006181020627287671
       H 1.318474396634582 0.2241299231185073 0.4092568796869673
   End
   Charge -1.0
   GuessBonds True
End
Engine DFTB
EndEngine
eor
# Now we run it in a QUILD-like setup (mechanical embedding)
export AMS_JOBNAME=quild
rm -rf $AMS_JOBNAME.results
"$AMSBIN/ams" << eor
Task GeometryOptimization
Properties Charges=yes
GeometryOptimization
 Convergence Gradients=1.0e-6
End
System
   Atoms
       0 -1.527946410885647 -0.2107366711137158 -0.0008116899510243671 region=QM
       H -0.8459142126057956 0.3517312394359257 0.4094504676540848
                                                                      region=QM
       H -1.834953147575289 0.1051014241823828 -0.8704652381864062
                                                                        region=QM
       H -1.328032016244278 -1.164422847242489 0.02894848344144469
                                                                        region=QM
        0 0.6370858511871781 -0.3378071707560572 -0.0006181020627287671 region=MM
```

```
(continues on next page)
```

```
(continued from previous page)
        H 1.318474396634582 0.2241299231185073 0.4092568796869673
                                                                         region=MM
   End
   GuessBonds True
    LoadForceFieldCharges region=QM file=H2O+.dftb.results
    LoadForceFieldCharges region=MM file=OH-.dftb.results
End
Engine Hybrid
   Energy
      Term Factor=1.0 Region=* EngineID=ForceField
      Term Factor=-1.0 Region=QM EngineID=ForceField
                                                          Charge=1.0
      Term Factor=1.0 Region=QM EngineID=DFTB
                                                          Charge=1.0
   End
   Engine DFTB
       Model GFN1-xTB
   EndEngine
    Engine ForceField
   EndEngine
EndEngine
eor
ddd=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#1#5`
eee=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -k AMSResults%Charges#5.3f`
echo "quild $charge $ddd $eee" >> $report
# Now we run it in a QMMM-like setup
export AMS_JOBNAME=qmmm
rm -rf $AMS JOBNAME.results
"$AMSBIN/ams" << eor
Properties Charges=yes
Task GeometryOptimization
GeometryOptimization
 Convergence Gradients=1.0e-6
End
System
   Atoms
       0 -1.527946410885647 -0.2107366711137158 -0.0008116899510243671 region=QM
       H -0.8459142126057956 0.3517312394359257 0.4094504676540848 region=QM
       H -1.834953147575289 0.1051014241823828 -0.8704652381864062
                                                                         region=QM
       H -1.328032016244278 -1.164422847242489 0.02894848344144469
                                                                         region=QM
        0 0.6370858511871781 -0.3378071707560572 -0.0006181020627287671 region=MM
        H 1.318474396634582 0.2241299231185073 0.4092568796869673
                                                                         region=MM
   End
```

```
GuessBonds True
   LoadForceFieldCharges region=QM file=H2O+.dftb.results
    LoadForceFieldCharges region=MM file=OH-.dftb.results
End
Engine Hybrid
   QMMM QMRegion=QM QMEngineID=DFTB MMEngineID=ForceField QMCharge=1.0 MMCharge=-1.0
   Engine DFTB
      Model GFN1-xTB
   EndEngine
   Engine ForceField
   EndEngine
EndEngine
eor
ddd=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#1#5`
eee=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -k AMSResults%Charges#5.3f
           $charge $ddd $eee" >> $report
echo "qmmm
echo "start of report"
cat $report
echo "end of report"
```

4.5 Example: Molecular gun with the hybrid engine

In this example we are going to really stretch the use of the Hybrid Engine, and shoot bullets (treated with a QM engine) at a surface described at the MM level.

The choice of bullets are HF molecules and the target is a two dimensional BN sheet, that looks like a graphene sheet, with half of the C atoms turned into N and the other half into B atoms. In a BN sheet the atoms have of course a small charge, which we pre calculate with a QM engine (DFTB).

It is important to understand the role of bonds in this example, because the number of atoms is not constant during the simulations, as bullets are fired (and hence appear), ricochet off the surface and hence disappear after a while. They may as well stick to, or penetrate into the surface, but this is beyond the hybrid engine concept.

In this example there are fixed bond orders withing the target and within the bullets. This is because we specify Guess-Bonds in the two system blocks (target and bullet). When a bullet is added its bonds are automatically added. The hybrid engine itself will never guess bonds and always use what is specified on input. No bonds are ever formed between the bullet and the surface (the QM and MM regions).

Download HybridGun.run

```
#!/bin/sh
# In this example we use the hybrid engine in a molecular gun MD application,...
⇔shooting HF molecules at a BN surface
```

(continued from previous page) # The BN slab represents the MM region and the "bullets" are the QM region # The regions are defined in the xyz files using end of line strings (atom attributes) # First we do two dftb calculations to get a guess of the charges to be used by the $_$ →force field. STRUCTDIR=\$AMSHOME/examples/Hybrid/HybridGun/molecules export AMS_JOBNAME=BNSlab.dftb rm -rf \$AMS_JOBNAME.results \$AMSBIN/ams << eor</pre> Task SinglePoint System GeometryFile \$STRUCTDIR/BNSlab.xyz End Engine DFTB EndEngine eor export AMS_JOBNAME=HF.dftb rm -rf \$AMS JOBNAME.results \$AMSBIN/ams << eor</pre> Task SinglePoint System GeometryFile \$STRUCTDIR/HF.xyz End Engine DFTB EndEngine eor # now we can run our MD simulation using both mechanical and electrostatic embedding for embedding in mechanical electrostatic do # because electrostatic embedding is more expensive we limit here the number of steps steps=1400 if [\$embedding = electrostatic] then steps=300 fi

```
export AMS_JOBNAME=SinkBox.embedding=$embedding
rm -rf $AMS_JOBNAME.results
$AMSBIN/ams << eor</pre>
Task MolecularDynamics
System
   GeometryFile $STRUCTDIR/BNSlab.xyz
   GuessBonds true
   LoadForceFieldCharges file=BNSlab.dftb.results
End
System H2
  GeometryFile $STRUCTDIR/HF.xyz
  GuessBonds true
  LoadForceFieldCharges file=HF.dftb.results
End
RNGSeed -1341016088 83513668 1764626453 -87803069 -1149690266 1963370818 -1393571175_
→1985130742
MolecularDynamics
   NSteps $steps
   Trajectory
       SamplingFreq 20
   End
   InitialVelocities
       Temperature 300
   End
   AddMolecules
       System H2
       Frequency 159
       CoordsBox 0 3 0 8.57 6 7
       VelocityDirection 0.45752820 0 -0.5540656
       Velocity 0.07
       Rotate Yes
       MinDistance 3.0
   End
   Preserve
       Momentum No
       AngularMomentum No
   End
   RemoveMolecules
     Formula *
     Frequency 101
     SinkBox FractionalCoordsBox="0 1 0 1 8 1000"
  End
End
Constraints
```

(continues on next page)

```
Atom 1
End
Engine Hybrid
QMMM qmRegion=qm mmEngineID=ForceField qmEngineID=dftb embedding=$embedding
Engine dftb
EndEngine
Engine ForceField
NonBondedCutoff 50 [Bohr]
EndEngine
EndEngine
eor
done
```

4.6 Example: The effect of specifying atom types, or not

Whether or not you specify the ForceField.Type on input (via atom attributes) makes a difference for the hybrid engine using a ForceField sub engine.

If you do, then for all regions these atom types will be used.

If you do not specify them, then for each region independently the atom typing will done automatically (if possible).

```
Download AtomAttributes.run
```

```
#!/bin/bash
export NSCM=1
# Here we show the role played by the (ForceField.Type) atom attribute
# on purpose we specify a strange type for the carbon atoms "C_2" (nonsensical for_
→any of the regions)
# we also show that capitalization does not matter for the keys ForceField.Charge and \_
\hookrightarrow Type.
# For the values, such as C_R the case matters
# In the first run we specify in the input ForceField.Type
# The result is that for all regions this will be used, and the type for C is always_
\hookrightarrow C-2
# ___
# region mol. C-type
#
  MM CH3 C_2
#
        С2Н6 С_2
#
  *
  QM
        CH4 C_2
#
#-
```

```
(continued from previous page)
export AMS_JOBNAME=type=inp.cap=none
rm -rf $AMS JOBNAME.results
$AMSBIN/ams <<eor</pre>
Task GeometryOptimization
Properties Gradients=yes
System
  Atoms
      C -1.559601 -1.234340 0.000641 region=QM forcefield.charge=-0.27 forcefield.
\rightarrowtype=C_2
       H -1.898371 -0.303860 0.503021 region=QM ForceField.Charge=0.09
                                                                           ForceField.
→Type=H_
       H -2.034545 -2.108050 0.494609 region=QM ForceField.Charge=0.09
                                                                           ForceField.
→Type=H_
       H -1.869847 -1.205955 -1.065139 region=QM ForceField.Charge=0.09
                                                                           ForceField.
→Type=H_
       C -0.047661 -1.348892 0.094039 region=MM ForceField.Charge=-0.27 ForceField.
\rightarrowType=C_2
       H 0.427282 -0.475182 -0.399929 region=MM ForceField.Charge=0.09
                                                                            ForceField.
\hookrightarrowType=H_
       H 0.291107 -2.279373 -0.408341 region=MM ForceField.Charge=0.09
                                                                           ForceField.
\hookrightarrowType=H_
       H 0.262583 -1.377277 1.159819 region=MM ForceField.Charge=0.09
                                                                           ForceField.
→Type=H_
   End
   GuessBonds True
End
Engine Hybrid
   QMMM qmRegion=QM qmEngineID=DFTB mmEngineID=ForceField
   Engine DFTB
   EndEngine
  Engine ForceField
  EndEngine
EndEngine
eor
# Not specifying the types, they will be guessed independently for all regions
#
# region mol. C-type
# _____
# MM CH3 C_R
# * C2H6 C_3
# QM CH4 C_3
#--
export AMS_JOBNAME=type=none.cap=none
```

```
rm -rf $AMS_JOBNAME.results
$AMSBIN/ams <<eor</pre>
Task GeometryOptimization
Properties Gradients=yes
System
   Atoms
       C -1.559601 -1.234340 0.000641 region=QM forcefield.charge=-0.27
       H -1.898371 -0.303860 0.503021 region=QM ForceField.Charge=0.09
       H -2.034545 -2.108050 0.494609 region=QM ForceField.Charge=0.09
       H -1.869847 -1.205955 -1.065139 region=QM ForceField.Charge=0.09
       C -0.047661 -1.348892 0.094039 region=MM ForceField.Charge=-0.27
       H 0.427282 -0.475182 -0.399929 region=MM ForceField.Charge=0.09
       H 0.291107 -2.279373 -0.408341 region=MM ForceField.Charge=0.09
       H 0.262583 -1.377277 1.159819 region=MM ForceField.Charge=0.09
   End
    GuessBonds True
End
Engine Hybrid
   QMMM qmRegion=QM qmEngineID=DFTB mmEngineID=ForceField
   Engine DFTB
   EndEngine
   Engine ForceField
   EndEngine
   GuessAttributesOnce False
EndEngine
eor
# The last two runs are mostly a technical test
# We change the capping setup, but that influences only the capping atom
export AMS_JOBNAME=type=inp.cap=inp
rm -rf $AMS_JOBNAME.results
$AMSBIN/ams <<eor</pre>
Task GeometryOptimization
Properties Gradients=yes
System
   Atoms
       C -1.559601 -1.234340 0.000641 region=QM forcefield.charge=-0.27 forcefield.
\rightarrowtype=C_2
```

H -1.898371 -0.303860 0.503021 region=QM ForceField.Charge=0.09	ForceField.
→Type=H_ H -2.034545 -2.108050 0.494609 region=QM ForceField.Charge=0.09	ForceField.
→Type=H_	ForceField
→Type=H_	roncerneid.
C -0.047661 -1.348892 0.094039 region=MM ForceField.Charge=-0.27	ForceField.
H 0.427282 -0.475182 -0.399929 region=MM ForceField.Charge=0.09	ForceField.
→Type=H_ H 0.291107 -2.279373 -0.408341 region=MM ForceField.Charge=0.09	ForceField.
→Type=H_ H 0.262583 -1.377277 1.159819 region=MM ForceField.Charge=0.09	ForceField.
←Type=H_ End	
GuessBonds True	
End	
QMMM qmRegion=QM qmEngineID=DFTB mmEngineID=ForceField	
Engine DFTB	
EndEngine	
Engine ForceField EndEngine	
Capping	
CappingElement Li	
AtomicInfoForCappingAtom ForceField.Type=Li End	
EndEngine	
eor	
export AMS_JOBNAME=type=none.cap=inp	
rm -rf \$AMS_JOBNAME.results	
\$AMSBIN/ams < <eor< td=""><td></td></eor<>	
Task GeometryOptimization	
Properties Gradients=yes	
System	
Atoms	
C -1.559601 -1.234340 0.000641 region=QM forcefield.charge=-0.27 H -1 898371 -0 303860 0 503021 region=OM ForceField Charge=0 09	
H -2.034545 -2.108050 0.494609 region=QM ForceField.Charge=0.09	
H -1.869847 -1.205955 -1.065139 region=QM ForceField.Charge=0.09	
C -0.047661 -1.348892 0.094039 region=MM ForceField.Charge=-0.27	
H 0.427282 -0.475182 -0.399929 region=MM ForceField.Charge=0.09	
H 0.262583 -1.377277 1.159819 region=MM ForceField.Charge=0.09	

```
End
    GuessBonds True
End
Engine Hybrid
   QMMM qmRegion=QM qmEngineID=DFTB mmEngineID=ForceField
   Engine DFTB
   EndEngine
   Engine ForceField
   EndEngine
   Capping
        CappingElement Li
        AtomicInfoForCappingAtom ForceField.Type=Li
   End
   GuessAttributesOnce False
EndEngine
eor
```

4.7 Example: The role of specifying the atom types

Now we look at a Propanenitrile molecule, the QM region is highlighted.



We run this with and without specifying the atom types on input. In principle this makes a difference for the MM type for atom "C(3)" in the MM sub calculation on atoms 3,7,8, and 9. If specified it will be C_3 (as it is in the whole Propanenitrile molecule), but if not it will be guessed as C_R. In practice there is no effect for this calculation.

Let us have a look at the report generated by the example, that pretty much explains what is done

Download report Propanenitrile.txt

We first check how bad the MM method is compared to the QM method for some distances. ${ \rightarrow } \text{in the QM}$ region
Here are the distance C(1)-C(2) C(1)-N(4) C(2)-H(5)	distances qm 1.456 1.147 1.095	(Angstrom) mm 1.467 1.157 1.110	as obtained with a QM and an MM method err(mm) 0.011 0.010 0.015					
Can we get be	etter resul	ts for the	QM region with the hybrid engine?					
Even though UFF has automatic atom typing, it still matters (in principle) whether we_ → specify it on input or not * Without typing for each region the types are automatically guessed * With typing the types are always as on input (for all regions)								
The only dif:	ference is	in the C t	ype for the MM region.					
Here are the →explicit ty	distances yping	(Angstrom)	as obtained with a QM and an Hybrid method without.					
distance	qm	hybrid	err(hybrid)					
C(1)-C(2)	1.456	1.456	0.000					
C(1)-N(4)	1.147	1.147	0.000					
C(2)-H(5)	1.095	1.092	-0.003					
Here are the →explicit ty	distances yping	(Angstrom)	as obtained with a QM and an Hybrid method with $\!\!\!\!\!\!$					
distance	qm	hybrid	err(hybrid)					
C(1)-C(2)	1.456	1.456	0.000					
C(1)-N(4)	1.147	1.147	0.000					
C(2)-H(5)	1.095	1.092	-0.003					
Here are some observations for this example * The hybrid engine does better than pure MM * The subtle issue whether or not we specify the types has negligible effect.								

```
Download Propanenitrile.run
```

```
#!/bin/sh
export NSCM=1
for engine in dftb forcefield
do
export AMS_JOBNAME=$engine
rm -rf $AMS_JOBNAME.results
"$AMSBIN/ams" << eor
Task GeometryOptimization
System
    Atoms
        C -0.02116 1.01286 0.0 region=qm
        C 0.01258 -0.45034 0.0 region=qm
        C 1.44394 -1.0175 0.0
        N -0.03362 2.17616 0.0 region=qm</pre>
```

```
(continued from previous page)
       H -0.54281 -0.80179 0.88302 region=qm
       H -0.54281 -0.80179 -0.88302 region=qm
       H 1.40659 -2.11445 0.0
       H 1.99584 -0.68766 -0.88907
       Н 1.99584 -0.68766 0.88907
   End
    GuessBonds true
End
Engine $engine
EndEngine
eor
done
report=report.txt
printf "\nWe first check how bad the MM method is compared to the QM method for some_
→distances in the QM region\n" > $report
bond1="C(1)-C(2)"
bond2="C(1)-N(4)"
bond3="C(2)-H(5)"
bond4="C(2)-C(3)"
aaa1qm=`$AMSBIN/amsreport dftb.results/dftb.rkf -r distance#1#2`
bbb1qm=`$AMSBIN/amsreport dftb.results/dftb.rkf -r distance#1#4`
ccc1qm=`$AMSBIN/amsreport dftb.results/dftb.rkf -r distance#2#5`
ddd1qm=`$AMSBIN/amsreport dftb.results/dftb.rkf -r distance#2#3`
aaa1mm=`$AMSBIN/amsreport forcefield.results/forcefield.rkf -r distance#1#2`
bbb1mm=`$AMSBIN/amsreport forcefield.results/forcefield.rkf -r distance#1#4`
ccc1mm=`$AMSBIN/amsreport forcefield.results/forcefield.rkf -r distance#2#5`
ddd1mm=`$AMSBIN/amsreport forcefield.results/forcefield.rkf -r distance#2#3
errmma=`echo "$aaa1mm- $aaa1qm" | bc`
errmmb=`echo "$bbb1mm- $bbb1qm" | bc`
errmmc=`echo "$ccc1mm- $ccc1qm" | bc`
errmmd=`echo "$ddd1mm- $ddd1qm" | bc`
printf "\nHere are the distances (Angstrom) as obtained with a QM and an MM method\n".
↔>> $report
printf "%10s %10s %10s %10s \n" "distance" "qm" "mm" "err(mm) ">> $report
printf "%10s %10.3f %10.3f %10.3f \n" $bond1 $aaa1qm $aaa1mm $errmma >> $report
printf "%10s %10.3f %10.3f %10.3f\n"
                                       $bond2 $bbb1qm $bbb1mm $errmmb >> $report
printf "%10s %10.3f %10.3f %10.3f \n" $bond3 $ccc1qm $ccc1mm $errmmc >> $report
printf "\nCan we get better results for the QM region with the hybrid engine?\n" >>
-→$report
printf "\nEven though UFF has automatic atom typing, it still matters (in principle)_
\rightarrow whether we specify it on input or not\n" >> $report
printf " * Without typing for each region the types are automatically guessed\n" >>
⇔$report
```

```
(continued from previous page)
```

```
printf " * With typing the types are always as on input (for all regions) \n" >>
⇔Śreport
printf "\nThe only difference is in the C type for the MM region.\n" >>$report
export AMS_JOBNAME=hybrid.types=no
rm -rf $AMS JOBNAME.results
"$AMSBIN/ams" << eor
Task GeometryOptimization
System
   Atoms
       C -0.02116 1.01286 0.0 region=qm
       C 0.01258 -0.45034 0.0 region=qm
        C 1.44394 -1.0175 0.0
        N -0.03362 2.17616 0.0 region=qm
        H -0.54281 -0.80179 0.88302 region=qm
        H -0.54281 -0.80179 -0.88302 region=qm
        H 1.40659 -2.11445 0.0
       Н 1.99584 -0.68766 -0.88907
       н 1.99584 -0.68766 0.88907
    End
   GuessBonds true
End
Engine Hybrid
   QMMM qmRegion=qm qmEngineID=dftb mmEngineID=forcefield
   Engine DFTB
   EndEngine
   Engine ForceField
   EndEngine
EndEngine
eor
aaa1hybrid=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#1#2`
bbb1hybrid=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#1#4`
ccc1hybrid=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#2#5`
ddd1hybrid=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#2#3`
errhybrida=`echo "$aaa1hybrid- $aaa1qm" | bc`
errhybridb=`echo "$bbb1hybrid- $bbb1qm" | bc
errhybridc=`echo "$ccc1hybrid- $ccc1qm" | bc`
errhybridd=`echo "$ddd1hybrid- $ddd1qm" | bc
printf "\nHere are the distances (Angstrom) as obtained with a QM and an Hybrid.
→method without explicit typing\n" >> $report
printf "%10s %10s %10s %10s \n" "distance" "qm" "hybrid" "err(hybrid) ">> $report
printf "%10s %10.3f %10.3f %10.3f \n" $bond1 $aaa1qm $aaa1hybrid $errhybrida >>
                                                                          (continues on next page)
```

```
(continued from previous page)
⇔$report
                                               $bbb1qm $bbb1hybrid $errhybridb >>
printf "%10s %10.3f %10.3f %10.3f\n"
⇔$report
printf "%10s %10.3f %10.3f %10.3f\n"
                                      $bond3 $ccc1qm $ccc1hybrid $errhybridc >>
⇔$report
export AMS_JOBNAME=hybrid.types=yes
rm -rf $AMS_JOBNAME.results
"$AMSBIN/ams" << eor
Task GeometryOptimization
System
   Atoms
                               region=qm
       C -0.02116 1.01286 0.0
                                    region=qm
       C 0.01258 -0.45034 0.0
       C 1.44394 -1.0175 0.0
                                    region=mm
                               region=qm
       N -0.03362 2.17616 0.0
       H -0.54281 -0.80179 0.88302 region=qm
       H -0.54281 -0.80179 -0.88302 region=qm
       H 1.40659 -2.11445 0.0 region=mm
       H 1.99584 -0.68766 -0.88907 region=mm
       H 1.99584 -0.68766 0.88907 region=mm
   End
   GuessBonds true
    LoadForceFieldAtomTypes File=forcefield.results
End
Engine Hybrid
   QMMM qmRegion=qm qmEngineID=dftb mmEngineID=forcefield
   Engine DFTB
   EndEngine
   Engine ForceField
   EndEngine
EndEngine
eor
aaa1hybrid=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#1#2`
bbb1hybrid=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#1#4`
ccc1hybrid=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#2#5`
ddd1hybrid=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#2#3`
errhybrida=`echo "$aaa1hybrid- $aaa1qm" | bc`
errhybridb=`echo "$bbb1hybrid- $bbb1qm" | bc
errhybridc=`echo "$ccc1hybrid- $ccc1qm" | bc`
errhybridd=`echo "$ddd1hybrid- $ddd1qm" | bc
printf "\nHere are the distances (Angstrom) as obtained with a QM and an Hybrid_
\rightarrowmethod with explicit typing\n" >> $report
```

```
printf "%10s %10s %10s %10s \n" "distance" "qm" "hybrid" "err(hybrid)">> $report
printf "%10s %10.3f %10.3f %10.3f \n" $bond1 $aaa1qm $aaa1hybrid $errhybrida >>
$report
printf "%10s %10.3f %10.3f %10.3f \n" $bond2 $bbb1qm $bbb1hybrid $errhybridb >>
$report
printf "%10s %10.3f %10.3f %10.3f \n" $bond3 $ccc1qm $ccc1hybrid $errhybridc >>
$report
printf "\nHere are some observations for this example\n" >>$report
printf " * The hybrid engine does better than pure MM\n" >>$report
printf " * The subtle issue whether or not we specify the types has negligible_
$effect.\n" >>$report
echo "begin report"
cat $report
echo "end report"
```

4.8 Example: Mixing DFT functionals

We consider a system of two weakly bonded molecules, namely NH3 and N2. We will use a GGA for the intra molecular interactions and LDA for the intermolecular one.

We look at two bond lengths, an N-H bond within the NH3 molecule, this is the "intra" bond.

The other is the bond from the N in NH3 to an N atom in the N2 molecule: the "inter" bond.

First we run the whole system with LDA and GGA, and finally with the hybrid engine.

The result for the hybrid calculation is that the "inter" bond has the value of the GGA calculation, whereas the "intra" one is equal to the LDA calculated one.

The energy expression used in the hybrid calculation is

 $E_{\mu}^{\text{hybrid}} = E_{\mu}^{\text{LDA/*}} + E_{\mu}^{\text{GGA/NH3}} - E_{\mu}^{\text{LDA/NH3}} + E_{\mu}^{\text{GGA/N2}} - E_{\mu}^{\text{LDA/N2}}$

Remember that the region * indicates the whole system, i.e. NH3 + N2.

Download MixingDFTFunctionals.run

```
system=AmmoniaN2
export AMS_JOBNAME=$system.lda
rm -rf $AMS_JOBNAME.results
$AMSBIN/ams<<EOF</pre>
Task GeometryOptimization
GeometryOptimization
    Convergence Gradients=1e-5
end
System
  Atoms
          -1.57871800-0.046611000.00000000 region=one-2.158621000.13639600-0.80956500 region=one-2.158621000.136396000.80956500 region=one-0.849471000.658193000.00000000 region=one
   Ν
    Η
    Η

        -0.84947100
        0.65819300
        0.00000000 region=one

        1.57871800
        0.04661100
        0.00000000 region=two

        1.03629999
        -1.31580113
        -0.10254699 region=two

     Η
     Ν
    Ν
  End
 GuessBonds True
end
Engine adf
  Basis Type=$bas
EndEngine
EOF
dInter=`$AMSBIN/amsreport $AMS_JOBNAME.results/adf.rkf -r distance#$intera1#$intera2`
dIntra=`$AMSBIN/amsreport $AMS_JOBNAME.results/adf.rkf -r distance#$intraa1#$intraa2`
echo "lda $dInter $dIntra" >> $report
export AMS_JOBNAME=$system.gga
rm -rf $AMS_JOBNAME.results
$AMSBIN/ams<<EOF</pre>
Task GeometryOptimization
GeometryOptimization
     Convergence Gradients=1e-5
end
LoadSystem
File $system.lda.results/adf.rkf
End
```

```
Engine adf
  xc gga=pbe
  Basis Type=$bas
EndEngine
EOF
dInter=`$AMSBIN/amsreport $AMS_JOBNAME.results/adf.rkf -r distance#$intera1#$intera2`
dIntra=`$AMSBIN/amsreport $AMS_JOBNAME.results/adf.rkf -r distance#$intraa1#$intraa2`
echo "gga $dInter $dIntra" >> $report
export AMS_JOBNAME=$system.hybrid
rm -rf $AMS_JOBNAME.results
$AMSBIN/ams<<EOF
Task GeometryOptimization
GeometryOptimization
   Convergence Gradients=1e-5
end
LoadSystem
 File $system.lda.results/adf.rkf
End
Engine Hybrid
   Energy
       Term Factor=1.0 Region=* EngineID=adf-lda
       Term Factor=-1.0 Region=one EngineID=adf-lda
       Term Factor=1.0 Region=one EngineID=adf-gga
       Term Factor=-1.0 Region=two EngineID=adf-lda
       Term Factor=1.0 Region=two EngineID=adf-gga
   End
   Engine adf adf-lda
       Basis Type=$bas
   EndEngine
   Engine adf adf-gga
      xc qqa=pbe
       Basis Type=$bas
   EndEngine
EndEngine
EOF
dInter=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#$intera1#
→$intera2`
```

```
(continued from previous page)
dIntra=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#$intraa1#
$intraa2`
echo "hybrid $dInter $dIntra" >> $report
echo "start of report"
cat $report
echo "end of report"
```

4.9 Example: Using capping atoms in a periodic system

Here we look at a polyethylene-like (infinite) chain, the carbons being alternatingly substituted with B and N atoms.



We will use a bunch of different QM systems in a QMMM setup, and check what happens. Inevitably we need to break B-N bonds, and hence capping atoms are used. (In the picture the QM region according to variant one is shown.) We also show the results obtained for the system with the pure QM and MM methods.

Let us have a look at the report generated by the example, that pretty much explains what is done

Download report PeriodicCapping.txt



					× 1	107	
	variation	QM at	oms				
	var1	B(1),H	H(2)				
	var2	2 B(5),H(6)					
	var3	N(3),H(4)					
	var4	N(7),H	H(8)				
	var5	B(1),H(2),N(3),H	H(4)				
	var6	B(5),H(6),N(7),H	H(8)				
Vari	ation one i	s equivalent to va	ariation two, and	d variation 3 shou	ld be equivalent	- 	
∽Wi	th variatio	n 4					
Vari	ation five	is equivalent to v	variation six				
Here	are the di	stances (Angstrom)	as obtained wit	th a QM and an MM	method		
di	stance	qm mm					
B(1) -H (2)	1.182 1.185					
B(5) - H (6)	1.182 1.185					
N (3) -H (4)	1.007 1.045					
N(/)-H(8)	1.007 1.045					
B(I) - N(3)	1.431 1.508					
B(S) - N (/)	1.431 1.508					
Norr	the three the	hubmid onging with	a corrected a transiet	iona for the OM m	aton		
NOW	we try the	nybria engine witi	i several variat.	tons for the QM re	egion		
THO	capping mot	hode are tried as	woll				
IWO	capping met	nous are crieu as	weit.				
	wariation	capping	enerau	B(1)-H(2)	B(5)-H(6)		
. N (3) -H (A)	N(7) - H(8)	B(1) = N(3)	B(5) - N(7)	D(3) II(0)	-	
	var1	fixed	-2 901499	1 184	1 185		
\hookrightarrow	1.045	1.045	1,508	1.508	1.100		
	var1	fractional	-2.787165	1.198	1.182		
\hookrightarrow	1.044	1.044	1.673	1.505		_	
	var2	fixed	-2.901499	1.185	1.184		
\hookrightarrow	1.045	1.045	1.508	1.508			
	var2	fractional	-2.787165	1.182	1.198	_	
\hookrightarrow	1.044	1.044	1.505	1.673			
	var3	fixed	-4.791110	1.184	1.184		
\hookrightarrow	0.993	1.044	1.508	1.507			
	var3	fractional	-4.733046	1.184	1.184	_	
\hookrightarrow	0.997	1.045	1.657	1.506			
	var4	fixed	-4.791110	1.184	1.184	_	
\hookrightarrow	1.044	0.993	1.507	1.508			
	var4	fractional	-4.733046	1.184	1.184		
\hookrightarrow	1.045	0.997	1.506	1.657			
	var5	fixed	-6.741093	1.189	1.187		
\hookrightarrow	1.003	1.045	1.390	1.489			
	var5	fractional	-6.648313	1.198	1.183	<u> </u>	
\hookrightarrow	1.003	1.045	1.405	1.505			
	var6	fixed	-6.741093	1.187	1.189	<u> </u>	
\hookrightarrow	1.045	1.003	1.489	1.390			
	var6	fractional	-6.648313	1.183	1.198	-	
\hookrightarrow	1.045	1.003	1.505	1.405			
Here	are some o	bservations					
	* generall	y the fixed cappir	ng seems a bit be	etter			
Here	are some r	emarks			<i>i</i>		
					(continues on ne	ext page)	

```
* Starting from the initial very bad structure the fixed capping fails_

→ completely for variant 5 and 6

    (not if you use as qm engine band and as mm engine dftb)

    * A reasonable starting geometry can avoid strange collapses

    * The more the two engines disagree about the capped QM region, the stronger the_

    → capping forces
```

Download PeriodicCapping.run

```
#!/bin/sh
export NSCM=1
report=report.txt
STRUCTDIR=$AMSHOME/examples/Hybrid/PeriodicCapping/systems
# ensure that not a comma is used for decimals in the printf function
LC_NUMERIC=en_US.UTF-8
export AMS_JOBNAME=reference
rm -rf $AMS_JOBNAME.results
$AMSBIN/ams<<EOF
Task GeometryOptimization
GeometryOptimization OptimizeLattice=yes Method=FIRE MaxIterations=300
System
  GeometryFile $STRUCTDIR/var1.xyz
  GuessBonds true
end
Engine DFTB
EndEngine
EOF
aaa1qm=`$AMSBIN/amsreport $AMS_JOBNAME.results/dftb.rkf -r distance#1#2`
aaa2gm=`$AMSBIN/amsreport $AMS JOBNAME.results/dftb.rkf -r distance#5#6`
bbb1qm=`$AMSBIN/amsreport $AMS_JOBNAME.results/dftb.rkf -r distance#3#4`
bbb2qm=`$AMSBIN/amsreport $AMS_JOBNAME.results/dftb.rkf -r distance#7#8`
ccc1qm=`$AMSBIN/amsreport $AMS_JOBNAME.results/dftb.rkf -r distance#1#3`
ccc2qm=`$AMSBIN/amsreport $AMS_JOBNAME.results/dftb.rkf -r distance#5#7`
printf "We optimize the lattice and test several distances\n" > $report
printf "\nThe system can be cut in several variations into a a QM and an MM part_
→breaking a B-N bond\n" >>$report
printf "\n%15s %20s\n" "variation" "QM atoms" >>$report
printf "%15s %20s\n" "var1" "B(1),H(2)" >>$report
printf "%15s %20s\n" "var2" "B(5),H(6)" >>$report
printf "%15s %20s\n" "var3" "N(3),H(4)" >>$report
printf "%15s %20s\n" "var4" "N(7),H(8)" >>$report
```

```
(continued from previous page)
printf "%15s %20s\n" "var5" "B(1),H(2),N(3),H(4)" >>$report
printf "%15s %20s\n" "var6" "B(5),H(6),N(7),H(8)" >>$report
printf "\nVariation one is equivalent to variation two, and variation 3 should be_
\rightarrowequivalent with variation 4\n" >>$report
printf "\nVariation five is equivalent to variation six\n" >>$report
export AMS_JOBNAME=cheap
rm -rf $AMS JOBNAME.results
$AMSBIN/ams<<EOF
Task GeometryOptimization
GeometryOptimization OptimizeLattice=yes Method=FIRE MaxIterations=300
System
  GeometryFile $STRUCTDIR/var1.xyz
  GuessBonds true
end
Engine ForceField
EndEngine
EOF
aaa1mm=`$AMSBIN/amsreport $AMS JOBNAME.results/forcefield.rkf -r distance#1#2`
aaa2mm=`$AMSBIN/amsreport $AMS_JOBNAME.results/forcefield.rkf -r distance#5#6`
bbb1mm=`$AMSBIN/amsreport $AMS_JOBNAME.results/forcefield.rkf -r distance#3#4`
bbb2mm=`$AMSBIN/amsreport $AMS_JOBNAME.results/forcefield.rkf -r distance#7#8`
ccc1mm=`$AMSBIN/amsreport $AMS_JOBNAME.results/forcefield.rkf -r distance#1#3`
ccc2mm=`$AMSBIN/amsreport $AMS_JOBNAME.results/forcefield.rkf -r distance#5#7`
printf "\nHere are the distances (Angstrom) as obtained with a QM and an MM method\n".
→>> $report
printf "%10s %10s %10s \n" "distance" "qm" "mm" >> $report
printf "%10s %10.3f %10.3f\n" "B(1)-H(2)" $aaa1qm $aaa1mm >> $report
printf "%10s %10.3f %10.3f\n"
                             "B(5)-H(6)" $aaa2qm $aaa2mm >> $report
printf "%10s %10.3f %10.3f\n" "N(3)-H(4)" $bbb1qm $bbb1mm >> $report
printf "%10s %10.3f %10.3f\n" "N(7)-H(8)" $bbb2qm $bbb2mm >> $report
                              "B(1)-N(3)" $ccc1qm $ccc1mm >> $report
printf "%10s %10.3f %10.3f\n"
printf "%10s %10.3f %10.3f\n"
                               "B(5)-N(7)" $ccc2qm $ccc2mm >> $report
printf "\nNow we try the hybrid engine with several variations for the QM region\n" >>

→ $report

printf "\nTwo capping methods are tried as well.\n" >>$report
↔ "energy" "B(1)-H(2)" "B(5)-H(6)" "N(3)-H(4)" "N(7)-H(8)" "B(1)-N(3)" "B(5)-N(7)" >
→> $report
for system in var1 var2 var3 var4 var5 var6
do
```

(continued from previous page) # This calc in only needed to start from a reasonable guess export AMS_JOBNAME=\$system.cheap rm -rf \$AMS_JOBNAME.results \$AMSBIN/ams<<EOF</pre> Task GeometryOptimization GeometryOptimization OptimizeLattice=yes Method=FIRE MaxIterations=300 System GeometryFile \$STRUCTDIR/\$system.xyz GuessBonds true end Engine ForceField EndEngine EOF for embedding in electrostatic do for capping in fixed fractional do export AMS_JOBNAME=\$system.embedding=\$embedding.capping=\$capping.go rm -rf \$AMS_JOBNAME.results \$AMSBIN/ams<<EOF</pre> Task GeometryOptimization GeometryOptimization OptimizeLattice=yes Method=FIRE MaxIterations=100 LoadSystem File \$system.cheap.results End Engine Hybrid Capping AllowHighBondOrders=true Option=\$capping QMMM qmRegion=qm qmEngineID=dftb mmEngineID=ForceField Embedding=\$embedding Engine Band EndEngine Engine DFTB EndEngine

```
Engine ForceField
   EndEngine
EndEngine
EOF
aaa1=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#1#2
aaa2=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#5#6`
bbb1=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#3#4`
bbb2=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#7#8`
ccc1=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#1#3`
ccc2=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#5#7`
xxx=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -k "AMSResults%Energy"`
printf "%15s %15s %15.6f %15.3f %15.3f %15.3f %15.3f %15.3f %15.3f %15.3f %15.3f %
↔$capping $xxx $aaa1 $aaa2 $bbb1 $bbb2 $ccc1 $ccc2 >> $report
done
done
done
printf "\nHere are some observations\n" >>$report
printf " * generally the fixed capping seems a bit better\n" >>$report
printf "\nHere are some remarks\n" >>$report
printf " * Starting from the initial very bad structure the fixed capping fails_
→completely for variant 5 and 6\n" >>$report
printf "
             (not if you use as qm engine band and as mm engine dftb) \n" >>$report
printf "
            * A reasonable starting geometry can avoid strange collapses\n" >>$report
printf "
           ^{\star} The more the two engines disagree about the capped QM region, the
→stronger the capping forces\n" >>$report
echo "begin report"
cat $report
echo "end report"
```

4.10 Example: Using capping atoms in a periodic system with charges

Now we look at a BN chain and use charges for the MM calculation.



Let us have a look at the report generated by the example, that pretty much explains what is done

Download report PeriodicCappingWithCharges.txt

We optimize the lattice and test several distances We divide the system in such a way that there are two equivalent, and hence neutral. ⇔regions. Here are the distances (Angstrom) as obtained with a QM and an MM method distance qm mm err(mm) B-H 1.182 1.181 -0.001 1.007 1.041 0.034 N-H B-N 1.431 1.498 0.067 Of course the force field results do not exactly match the QM results, the error- \rightarrow displayed in the last column Now we try the hybrid engine, can we improve the bonds in the QM region? We start from the geometry calculated with the (cheap) forcefield In this table we show the errors in bond lengths (in the QM region) of the hybrid- \rightarrow method with respect to the QM method

	embedding	capping	energy	В-Н	N-H				
\hookrightarrow	B-N	C 1 1	6 045045	0.004	0 007				
	mechanical	Ilxed	-6.845015	0.004	-0.00/	-			
\hookrightarrow	-0.049		6 546965	0.040	0.000				
	mechanical	fractional	-6./4636/	0.013	-0.006	-			
\hookrightarrow	-0.034								
el	ectrostatic	fixed	-6.748753	0.002	-0.004	-			
\hookrightarrow	-0.040								
el	ectrostatic	fractional	-6.652196	0.010	-0.003	-			
\hookrightarrow	-0.024								
Here	are some obs	ervations							
* the B-H distance is a bit worse than with a plain forcefield, especially with.									
⇔fractional capping									
\star the N-H distance is much better than with the plain forcefield									
* the B-N distance is a bit better than with the plain forcefield, now too short.									
↔ Fractional capping works best.									
* Electrostatic embedding is doing slightly better than mechanical embedding,.									
\rightarrow the biggest improvement is on the B-N bond									

Download PeriodicCappingWithCharges.run

```
#!/bin/bash
export NSCM=1
report=report.txt
STRUCTDIR=$AMSHOME/examples/Hybrid/PeriodicCapping/systems
# ensure that not a comma is used for decimals in the printf function
LC_NUMERIC=en_US.UTF-8
export AMS_JOBNAME=reference
rm -rf $AMS_JOBNAME.results
$AMSBIN/ams<<EOF</pre>
Task GeometryOptimization
GeometryOptimization OptimizeLattice=yes Method=FIRE MaxIterations=300
System
  GeometryFile $STRUCTDIR/var5.xyz
  GuessBonds true
end
Engine DFTB
EndEngine
EOF
aaa1qm=`$AMSBIN/amsreport $AMS_JOBNAME.results/dftb.rkf -r distance#1#2`
bbb1qm=`$AMSBIN/amsreport $AMS_JOBNAME.results/dftb.rkf -r distance#3#4`
ccc1qm=`$AMSBIN/amsreport $AMS_JOBNAME.results/dftb.rkf -r distance#1#3`
```

```
(continued from previous page)
printf "We optimize the lattice and test several distances\n" > $report
printf "\nWe divide the system in such a way that there are two equivalent, and hence.
→neutral regions.\n" >>$report
export AMS_JOBNAME=cheap
rm -rf $AMS JOBNAME.results
$AMSBIN/ams<<EOF
Task GeometryOptimization
GeometryOptimization OptimizeLattice=yes Method=FIRE MaxIterations=300
System
   GeometryFile $STRUCTDIR/var5.xyz
   LoadForceFieldCharges File=reference.results
  GuessBonds true
end
Engine ForceField
   NonBondedCutoff 50 [Bohr]
EndEngine
EOF
aaa1mm=`$AMSBIN/amsreport $AMS JOBNAME.results/forcefield.rkf -r distance#1#2`
bbb1mm=`$AMSBIN/amsreport $AMS_JOBNAME.results/forcefield.rkf -r distance#3#4`
ccc1mm=`$AMSBIN/amsreport $AMS_JOBNAME.results/forcefield.rkf -r distance#1#3`
errmma=`echo "$aaa1mm- $aaa1qm" | bc`
errmmb=`echo "$bbb1mm- $bbb1qm" | bc
errmmc=`echo "$ccc1mm- $ccc1qm" | bc`
printf "\nHere are the distances (Angstrom) as obtained with a QM and an MM method\n"...
→>> $report
printf "%10s %10s %10s %10s \n" "distance" "qm" "mm" "err(mm)">> $report
printf "%10s %10.3f %10.3f %10.3f \n" "B-H" $aaa1qm $aaa1mm $errmma >> $report
printf "%10s %10.3f %10.3f %10.3f \n" "N-H" $bbb1am $bbb1mm $errmmb >> $report
printf "%10s %10.3f %10.3f %10.3f \n" "B-N" $ccc1qm $ccc1mm $errmmc >> $report
printf "\nOf course the force field results do not exactly match the QM results, the.
\rightarrowerror displayed in the last column\n" >> $report
printf "\nNow we try the hybrid engine, can we improve the bonds in the QM region?\n".
↔>> $report
printf "\nWe start from the geometry calculated with the (cheap) forcefield\n" >>
-→$report
printf "\nIn this table we show the errors in bond lengths (in the QM region) of the_
↔ hybrid method with respect to the QM method\n" >> $report
printf "\n%15s %15s %15s %15s %15s %15s \n" "embedding" "capping" "energy" "B-H" "N-H
                                                                          (continues on next page)
```

Chapter 4. Examples

```
↔" "B-N" >> $report
for system in var5
do
for embedding in mechanical electrostatic
do
for capping in fixed fractional
do
export AMS_JOBNAME=$system.embedding=$embedding.capping=$capping.go
rm -rf $AMS_JOBNAME.results
$AMSBIN/ams<<EOF</pre>
Task GeometryOptimization
Properties Gradients=yes
GeometryOptimization OptimizeLattice=yes Method=FIRE MaxIterations=100
LoadSystem
File cheap.results
End
Engine Hybrid
        Capping AllowHighBondOrders=true Option=$capping
 QMMM qmRegion=qm qmEngineID=dftb mmEngineID=ForceField Embedding=$embedding
   Engine Band
   EndEngine
   Engine DFTB
   EndEngine
   Engine ForceField
       NonBondedCutoff 50 [Bohr]
   EndEngine
EndEngine
EOF
aaa1=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#1#2`
bbb1=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#3#4`
ccc1=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -r distance#1#3`
xxx=`$AMSBIN/amsreport $AMS_JOBNAME.results/hybrid.rkf -k "AMSResults%Energy"`
# printf "%15s %15s %15s %15.6f %15.3f %15.3f %15.3f \n" $embedding $system $capping
→$xxx $aaa1 $bbb1 $ccc1 >> $report
erra=`echo "$aaa1- $aaa1qm" | bc
errb=`echo "$bbb1- $bbb1qm" | bc
errc=`echo "$ccc1- $ccc1qm" | bc
```

```
printf "%15s %15.6f %15.3f %15.3f %15.3f n" $embedding $capping $xxx $erra $errb
done
done
done
printf "\nHere are some observations\n" >>$report
printf " * the B-H distance is a bit worse than with a plain forcefield,
\leftrightarrowespecially with fractional capping\n" >>$report
printf "
            * the N-H distance is much better than with the plain forcefield \n" >>
-→$report
printf "
           * the B-N distance is a bit better than with the plain forcefield, now_
→too short. Fractional capping works best.\n" >>$report
printf " * Electrostatic embedding is doing slightly better than mechanical.
→embedding, the biggest improvement is on the B-N bond\n" >>$report
echo "begin report"
cat $report
echo "end report"
```

4.11 Example: QMMM with capping atoms

This is an example of QMMM using capping atoms. Capping atoms are added automatically when bonds are broken (between the QM and MM region). Because the amber forcefield is used the AtomicInfoForCappingAtom needs to be set, as the default type "H" is not an AMBER type.

Download QMMM_Butane.run

```
#! /bin/sh
"$AMSBIN/ams" <<eor
Task GeometryOptimization
GeometryOptimization
   Convergence Gradients=1E-4
End
System
   Atoms
       С
          0.0000 0.0000 0.0000 region=QM ForceField.Charge=0.0 ForceField.
→Type=CT
       Н 1.0910 0.0000 0.0000
                                    region=QM ForceField.Charge=0.0 ForceField.
\leftrightarrowTvpe=HC
       H -0.3598 1.0300 0.0000 region=QM ForceField.Charge=0.0 ForceField.
\leftrightarrowType=HC
       C -0.5021 -0.7074 -1.2586 region=QM ForceField.Charge=0.0 ForceField.
→Type=CT
       H -0.1397 -1.7383 -1.2662 region=QM ForceField.Charge=0.0 ForceField.
\hookrightarrowType=HC
       H -1.5949 -0.7053 -1.2662 region=QM ForceField.Charge=0.0 ForceField.
\rightarrowType=HC
       H -0.1307 -0.1841 -2.1433 region=QM ForceField.Charge=0.0 ForceField.
```

 \rightarrow Type=HC C -0.5195 -0.7318 1.2374 region=MM ForceField.Charge=0.0 ForceField. ⇔Type=CT C -0.0376 -0.0530 2.5227 region=MM ForceField.Charge=0.0 ForceField. →Type=CT H 1.0524 -0.0554 2.5580 region=MM ForceField.Charge=0.0 ForceField. →Type=HC H-0.3994 0.9752 2.5580 region=MM ForceField.Charge=0.0 ForceField. \hookrightarrow Type=HC H -0.4230 -0.5961 3.3860 region=MM ForceField.Charge=0.0 ForceField. \rightarrow Type=HC H -0.1631 -1.7630 1.2286 region=MM ForceField.Charge=0.0 ForceField. \leftrightarrow Type=HC H -1.6105 -0.7355 1.2286 region=MM ForceField.Charge=0.0 ForceField. →Type=HC End BondOrders 1 2 1.0 1 3 1.0 1 8 1.0 1 4 1.0 8 14 1.0 8 13 1.0 8 9 1.0 4 5 1.0 4 6 1.0 4 7 1.0 9 12 1.0 9 11 1.0 9 10 1.0 End End Engine Hybrid QMMM QMRegion QM QMEngineID ADF MMEngineID ForceField End Capping AtomicInfoForCappingAtom ForceField.Type=HC ForceField.Charge=0.0 End Engine ADF Basis Type DZP End Relativitv Level None End EndEngine Engine ForceField Type Amber95 EndEngine EndEngine

eor

4.12 Example*: Mechanical embedding (QUILD)

This is a fairly technical test, testing the case insensitivity of region and engine names. Furthermore gradients are tested.

Download qmmm_water.run

```
#!/bin/sh
hybridenginename=hybrid  # this is a temporary hack
report=report.txt
echo "Start of the report" > $report
echo "-----" >> $report
echo "Start of the region names test" >> $report
echo "-----" >> $report
export AMS_JOBNAME=regionnamestest
rm -rf $AMS_JOBNAME.results
$AMSBIN/ams <<eor</pre>
Task SinglePoint
Properties Gradients=yes
System
 Atoms
    C 0.0 0.0 0.0 region=a
    0 1.13 0.0 0.0 region=A
    0 -1.13 0.0 0.0 region=B
  End
  BondOrders
   1 2 2.0
      1 3 2.0
 End
End
Engine Hybrid
  Energy
     Term factor=1.0region=*engineID=ForceFieldTerm factor=1.0region=AengineID=dftB-2Term factor=-1.0region=aengineID=ForceField
  End
  Capping
     AllowHighBondOrders True # Because we cut through double bonds here ...
     AtomicInfoForCappingAtom ForceField.Type=H_ # Remove ForceField.Charge_
→because we do not specify charges for other atoms
```

```
End
  Engine ForceField
  EndEngine
  Engine dftb dftb-1
   Model GFN1-xTB
  EndEngine
  Engine dftb dFtb-2
   Model SCC-DFTB
   ResourcesDir QUASINANO2015
  EndEngine
  Engine mopac
  EndEngine
EndEngine
eor
echo "Energy for region names test" >> $report
$AMSBIN/amsreport $AMS_JOBNAME.results/$hybridenginename.rkf -k "AMSResults%Energy" 💶
↔>> $report
                      -----" >> $report
echo "-----
echo "End of the region names test" >> $report
echo "-----" >> $report
echo "-----" >> $report
echo "Start of the gradient test" >> $report
          -----" >> $report
echo "-----
# strange capitalization is on purpose
for capping in fractionAl fiXed
do
for num in true false
do
export AMS_JOBNAME=gradtest.capping=$capping.num=$num
rm -rf $AMS JOBNAME.results
$AMSBIN/ams <<eor</pre>
Task SinglePoint
Properties Gradients=yes
EngineDebugging IgnoreGradientsRequest=$num NeverQuiet=false
NumericalDifferentiation NuclearStepSize=1.0e-4
System
 Atoms
  C 0.0 0.0 0.0 region=a
                                                                     (continues on next page)
```

```
0 1.13 0.0 0.0 region=a
    0 -1.13 0.0 0.0 region=b
 End
 BondOrders
   1 2 2.0
      1 3 2.0
 End
End
Engine Hybrid
 Capping
    Option $capping
    AllowHighBondOrders True # Because we cut through double bonds here ...
    AtomicInfoForCappingAtom ForceField.Type=H_ # Remove ForceField.Charge_
↔ because we do not specify charges for other atoms
  End
  Energy
    Term factor=1.0 region=*
                                  engineID=ForceField
     Term factor=1.0 region=a
                                 engineID=dftb-2
engineID=ForceField
    Term factor=-1.0 region=a
  End
  Engine ForceField
  EndEngine
  Engine dftb dftb-1
  Model GFN1-xTB
  EndEngine
  Engine dftb dftb-2
   Model SCC-DFTB
   ResourcesDir QUASINANO2015
  EndEngine
  Engine mopac
  EndEngine
EndEngine
eor
echo "gradients for capping=$capping num=$num:" >> $report
$AMSBIN/amsreport $AMS_JOBNAME.results/$hybridenginename.rkf -k "AMSResults%Gradients#
→#3" >> $report
done
done
echo "-----" >> $report
echo "End of the gradient test" >> $report
echo "-----" >> $report
echo "-----" >> $report
echo "Start of the singleton test" >> $report
echo "-----" >> $report
echo "Calculate the same system twice with the same engine but with different.
                                                                    (continues on next page)
```

```
(continued from previous page)
```

```
⇔settings. The energy should be non-zero" >> $report
for engine in dftb band adf mopac
do
export AMS_JOBNAME=testsingleton.engine.$engine
rm -rf $AMS_JOBNAME.results
$AMSBIN/ams << eor</pre>
Task SinglePoint
System
  GeometryFile $AMSHOME/atomicdata/Molecules/TestMols/Acetamide.xyz
End
Engine Hybrid
  AllowSanityCheckWarnings true
  Energy
     Term factor=1.0 region=*
                                       engineID=$engine-1
     Term factor=-1.0 region=*
                                       engineID=$engine-2
  End
  Engine band band-1
   xc gga=pbe
  EndEngine
  Engine band band-2
     basis type=SZ
  EndEngine
  Engine adf adf-1
    xc gga=pbe
  EndEngine
  Engine adf adf-2
    basis type=SZ
  EndEngine
  Engine mopac mopac-1
    Model PM6
  EndEngine
  Engine mopac mopac-2
    Model PM7
  EndEngine
  Engine DFTB dftb-1
    Model GFN1-xTB
  EndEngine
  Engine DFTB dftb-2
     Model SCC-DFTB
     ResourcesDIR QUASINAN02015
  EndEngine
EndEngine
eor
echo "Hopfully nonzero energy for engine=$engine" >> $report
$AMSBIN/amsreport $AMS_JOBNAME.results/$hybridenginename.rkf -k "AMSResults%Energy"
↔>> $report
                                                                          (continues on next page)
```

```
done
echo "------" >> $report
echo "End of the singleton test" >> $report
echo "------" >> $report
cat $report
```

4.13 Example: pdb2adf transforms a PDB file to a QMMM input file

Download pdb2adf.run

```
#! /bin/sh
# This example shows how to use the utility pdb2adf,
# which creates an amsified ADF input file (ADF>=2020) from a PDB file,
# for a subsequent QM/MM calculation using ADF.
# First create the PDB file
# __
cat << eor > chymotrypsin.pdb
HEADER COMPLEX (SERINE PROTEASE/INHIBITOR) 12-MAR-97 1AFQ
        CRYSTAL STRUCTURE OF BOVINE GAMMA-CHYMOTRYPSIN COMPLEXED
TITLE
TITLE
        2 WITH A SYNTHETIC INHIBITOR
REMARK
      Adaptation of original PDB file by M. Swart, March 2005
REMARK
      only coordinates of GAMMA-CHYMOTRYPSIN are kept;
REMARK
REMARK rest has been deleted.
REMARK
ATOM
        1 N CYSA 1
                             13.717 20.021 22.754 1.00 13.46 PROA N
        2 CA CYS A 1
                             14.211 18.932 23.617 1.00 13.34
                                                                   PROA C
ATOM
ATOM
        3 C CYS A 1
                             13.597 19.033 25.005 1.00 13.34
                                                                   PROA C
ATOM
        4 O CYSA 1
                             12.953 20.026 25.329 1.00 13.48
                                                                   PROA O
ATOM
        5 CB CYS A 1
                             15.734 19.018 23.753 1.00 13.44
                                                                   PROA C
        6 SG CYS A 1
                             16.298 20.647 24.361 1.00 13.30
                                                                   PROA S
ATOM
                             13.801 17.985 25.813 1.00 13.44
         7 N GLY A 2
ATOM
                                                                   PROA N
         8 CA GLY A 2
                             13.369 17.952 27.214 1.00 13.65
                                                                    PROA C
ATOM
                              11.904 18.088 27.631 1.00 13.87
ATOM
         9 C
               GLY A 2
                                                                    PROA C
ATOM
        10 O
               GLY A 2
                             11.669 18.375 28.799 1.00 13.63
                                                                    PROA O
ATOM
        11 N
               VAL A 3
                             10.947 17.887 26.732 1.00 14.18
                                                                    PROA N
ATOM
       12 CA VAL A 3
                             9.559 17.968 27.090 1.00 14.86
                                                                   PROA C
ATOM
       13 C VAL A 3
                             8.875 16.684 26.624 1.00 15.04
                                                                  PROA C
ATOM
       14 O VAL A 3
                             8.529 16.546 25.452 1.00 14.91
                                                                  PROA O
ATOM
       15 CB VAL A 3
                             8.861 19.211 26.437 1.00 15.00
                                                                  PROA C
       16 CG1 VAL A 3
                             7.403 19.299 26.880 1.00 15.08
ATOM
                                                                   PROA C
ATOM
       17 CG2 VAL A 3
                             9.585 20.486 26.805 1.00 15.27
                                                                   PROA C
       18 N PRO A 4
                             8.754 15.691 27.519 1.00 15.39
ATOM
                                                                   PROA N
       19 CA PRO A 4
                             8.121 14.407 27.206 1.00 16.09
                                                                    PROA C
ATOM

        20
        C
        PRO A
        4
        6.675
        14.535
        26.769
        1.00
        16.35

        21
        O
        PRO A
        4
        5.957
        15.387
        27.275
        1.00
        16.43

                                                                    PROA C
ATOM
ATOM
                                                                    PROA O
```

ATOM	22	СВ	PRO	А	4	8.219	13.635	28.527	1.00 15.91	PROA	С
ATOM	23	CG	PRO	А	4	9.369	14.297	29.244	1.00 16.02	PROA	С
ATOM	24	CD	PRO	А	4	9.166	15.742	28.928	1.00 15.56	PROA	С
ATOM	25	Ν	ALA	А	5	6.262	13.690	25.827	1.00 16.87	PROA	N
ATOM	26	CA	ALA	А	5	4.874	13.703	25.351	1.00 17.53	PROA	С
ATOM	27	С	ALA	А	5	4.020	13.055	26.437	1.00 17.86	PROA	С
ATOM	28	0	ALA	А	5	2.862	13.413	26.643	1.00 17.81	PROA	0
ATOM	29	СВ	ALA	А	5	4.740	12.936	24.027	1.00 17.34	PROA	С
ATOM	30	Ν	ILE	А	6	4.615	12.104	27.143	1.00 18.37	PROA	N
ATOM	31	CA	ILE	А	6	3.941	11.419	28.239	1.00 18.99	PROA	С
ATOM	32	С	ILE	А	6	4.553	11.993	29.514	1.00 19.59	PROA	С
ATOM	33	0	ILE	А	6	5.726	11.771	29.807	1.00 19.17	PROA	0
ATOM	34	CB	ILE	А	6	4.190	9.909	28.190	1.00 19.29	PROA	С
ATOM	35	CG1	ILE	А	6	3.631	9.335	26.886	1.00 19.66	PROA	С
ATOM	36	CG2	ILE	А	6	3.552	9.232	29.399	1.00 19.38	PROA	С
ATOM	37	CD1	ILE	А	6	3.977	7.887	26.674	1.00 20.59	PROA	С
ATOM	38	Ν	GLN	А	7	3.760	12.742	30.265	1.00 20.52	PROA	N
ATOM	39	CA	GLN	А	7	4.262	13.374	31.468	1.00 21.69	PROA	С
ATOM	40	С	GLN	А	7	4.683	12.459	32.597	1.00 22.02	PROA	С
ATOM	41	0	GLN	А	7	3.954	11.535	32.978	1.00 21.89	PROA	0
ATOM	42	CB	GLN	А	7	3.259	14.392	31.997	1.00 22.77	PROA	С
ATOM	43	CG	GLN	А	7	3.369	15.749	31.349	1.00 24.86	PROA	С
ATOM	44	CD	GLN	А	7	2.467	16.774	32.004	1.00 25.86	PROA	С
ATOM	45	OE1	GLN	А	7	1.660	17.417	31.337	1.00 27.24	PROA	0
ATOM	46	NE2	GLN	Α	7	2.601	16.934	33.325	1.00 26.82	PROA	N
ATOM	47	Ν	PRO	Α	8	5.898	12.675	33.125	1.00 22.41	PROA	N
ATOM	48	CA	PRO	Α	8	6.345	11.830	34.231	1.00 22.84	PROA	С
ATOM	49	С	PRO	А	8	5.524	12.215	35.459	1.00 23.50	PROA	С
ATOM	50	0	PRO	Α	8	5.069	13.359	35.575	1.00 23.62	PROA	0
ATOM	51	СВ	PRO	Α	8	7.821	12.219	34.389	1.00 22.58	PROA	С
ATOM	52	CG	PRO	Α	8	7.864	13.641	33.894	1.00 22.61	PROA	С
ATOM	53	CD	PRO	Α	8	6.972	13.583	32.678	1.00 22.34	PROA	С
ATOM	54	Ν	VAL	Α	9	5.267	11.244	36.323	1.00 24.00	PROA	N
ATOM	55	CA	VAL	A	9	4.516	11.478	37.543	1.00 24.57	PROA	С
ATOM	56	С	VAL	Α	9	5.471	11.122	38.665	1.00 24.99	PROA	С
ATOM	57	0	VAL	A	9	5.927	9.982	38.759	1.00 24.82	PROA	0
ATOM	58	CB	VAL	A	9	3.273	10.580	37.613	1.00 24.64	PROA	C
ATOM	59	CG1	VAL	A	9	2.596	10.725	38.969	1.00 24.88	PROA	C
ATOM	60	CG2	VAL	A	9	2.308	10.935	36.488	1.00 24.78	PROA	C
ATOM	61	N	LEU	A	10	5.827	12.119	39.464	1.00 25.78	PROA	N
ATOM	62	CA	LEU	A	10	6.752	11.921	40.568	1.00 26.56	PROA	C
ATOM	63	С	LEU	A	10	6.043	11.958	41.914	1.00 26.89	PROA	C
ATOM	64	O	LEU	A	10	5.187	12.847	42.105	1.00 27.07	PROA	0
ATOM	65	CB	LEU	A	10	1.857	12.973	40.501	1.00 26.93	PROA	C
ATOM	66	CG CD1	LEU	A	10	8.721	14.055	39.255	1.00 27.39	PROA	C
ATOM	60	CDI	LEU	A	10	9.351	11 601	30.769	1 00 27.66	PROA	C
ATOM	60	CDZ OVT	LEU	A	10	9.700	11.001	39.000	1.00 27.46	PROA	
AIOM	70	UXI	LEU	A	10	0.329	11.000	42.743	1.00 27.55	PROA	0
TER	10		LEU	A	TO						
END											
eor											
#											
π											
# then r	מ מווי	roar	am +	2	reate ID	F input f	File				
# The program works interactively. The input described here are answers to the											
- mesti	ons	in wo	110.		CIUCLIVE.	-y. 1118	input de	.serrbed	nere are answers		
, gueses	0110									<i>.</i>	

```
(continued from previous page)
# that were asked interactively.
# In cases where the user agrees with the suggestion given by the program,
# the user can press the **Enter** key, which is shown here with an empty line.
# _____
→ - - -
$AMSBIN/pdb2adf << eor</pre>
chymotrypsin.pdb
3 4 5
С
5
3 4 15 16
С
Υ
1
1
17.5
eor
# -
# Questions asked were:
# Q1: Please give name of PDB-file
# A1: chymotrypsin.pdb
# Q2: Do you want a logfile to be written (Y/n) ?
# A2: Enter
#
 Q3: ..
#
      Found the following terminal amino acid residues : (C-term) 10 (N-term)
\rightarrow 1
     Do you want to use these as terminal residues (Y/n) ?
#
# A3: Enter
# Q4: Multiple AMBER options for CYS :
#
      0 Decide every time differently
#
       1 CYS Cysteine (SH)
       2 CYM Deprotonated Cysteine (S-)
#
       3 CYX Cystine (S-S bridge)
#
#
     Suggested option: 0
# A4: Enter
  Q5: Multiple AMBER options for CYS 1 ( 1) :
#
#
      . . .
#
      Suggestion: 1
  A5: Enter
#
  06: ...
#
#
      Option Molecule Option Molecule Option Molecule _
→ Option Molecule
                            4: PRO 4
                                               7: GLN 7 10: LEU 10
    1: CYS 1
#
#
          2: GLY 2
                            5: ALA 5
                                               8: PRO 8
          3: VAL 3
                            6: ILE 6
                                               9: VAL
                                                         9
#
     Give option number of molecules to be put in QM region (or 'c' to continue):
#
#
     Note: by specifying a negative number a molecule is removed from the QM region
#
  A6: 3 4 5
```

```
Q7: ...
      Give option number of molecules to be put in QM region (or 'c' to continue):
#
#
      Note: by specifying a negative number a molecule is removed from the QM region
#
  A7: C
  Q8: Make a choice for the QM/MM treatment of VAL
                                                    3
#
       0: Put completely in QM region
#
#
       1: Cut off at C-alpha (put NH in QM region, CO in MM region)
#
       2: Cut off at C-alpha (put NH in MM region, CO in QM region)
       3: Cut off at C-alpha (put NH and CO in MM region)
#
#
       4: Cut off at C-alpha (put NH and CO in QM region, sidechain in MM region)
#
       5: Put only part of sidechain in QM region
#
      Suggestion: 2
#
      Give choice:
#
 A8: 5
#
  Q9: Atoms belonging to molecule VAL 3
                       6: HB MM
#
      1: N MM
                                                           11: CG2 MM
    16:0 MM
\rightarrow
        2: H MM
#
                                  7: CG1 MM
                                                           12: HG21 MM
         3: CA
                 MM
                                   8: HG11 MM
#
                                                            13: HG22 MM
         4: HA
                                  9: HG12 MM
                                                            14: HG23 MM
#
                 MM
         5: CB MM
#
                                  10: HG13 MM
                                                            15: C MM
#
      Give option number of atoms to be put in QM region (or 'c' to continue):
      (Note: a range can be entered as 3-21, while a negative number removes an atom)
#
# A9: 3 4 15 16
# Q10: ...
#
      Give option number of atoms to be put in QM region (or 'c' to continue):
#
      (Note: a range can be entered as 3-21, while a negative number removes an atom)
# A10: C
# 011: Make a choice for the OM/MM treatment of PRO 4
      . . .
      Suggestion: 2
# A11: Enter
# Q12: Make a choice for the QM/MM treatment of ALA 5
      Suggestion: 1
# A12: Enter
# Q13: Do you want to add solvent to your system (Y/n) ?
# A13: Y
# Q14: Solvent (box) available:
#
        1: HOH HOH Water molecule
         2: MOH
                   MOH Methanol molecule
#
#
        3: CHL
                   CHL Chloroform molecule
# A14: 1
# Q15: Box Shape options:
      1 Spherical box
#
       2 Cubic box
#
# A15: 1
# Q16: Give boxsize (def.: 16.71 Angs)
# A16: 17.5
# --
→ - - -
# _____
# now for checking correctness
# --
echo ""
```

```
echo "ADF inputfile made by pdb2adf"
echo ""
```

cat chymotrypsin.pdb2adf

rm chymotrypsin.pdb chymotrypsin.pdb.log chymotrypsin.p2a.pdb chymotrypsin.pdb2adf

CHAPTER

FIVE

KEYWORDS

5.1 Links to manual entries

hybrid:

- *Capping* (page 16)
- Committee (page 15)
- *Energy* (page 12)

- *Engine* (page 11)
- *QMMM* (page 14)
- RestartSubEngines (page 18)

5.2 Summary of all keywords

5.2.1 Engine Hybrid

AllowSanityCheckWarnings

Туре

Bool

Default value No

Description

Sanity checks will be performed on the setup. If this option is on, only warnings are printed. If not the program will stop on warnings.

Capping

Туре

Block

Description

This block is about capping details. Capping occurs with hydrogen atoms when a bond is broken between an atom inside the region and one outside.

AllowHighBondOrders

Type Bool

Default value No

Description

Allows capping of interregional aromatic, double and triple bonds. This is normally not a good idea, since the capping is done with hydrogen atoms.

AtomicInfoForCappingAtom

Туре

String

Default value

ForceField.Type=H_ForceField.Charge=0.0

Description

The AtomicInfo for the capping atoms. Typically a string like ForceField.Type=X much like forcefield info is entered in the System block for normal atoms.

CappingElement

Туре

String

Default value

Η

Description

The element to be used for capping. The hydrogen atom has the advantage that it is very small.

CheckCapping

Туре

Bool

Default value

Yes

Description

The same outside atom can be involved in multiple capping coordinate definitions. This is not a good idea, and this will not be accepted by using this check.

Distance

Type Float

Default value

-1.0

Description

A negative value means automatic. In that case the sum of covalent radii is used

Option

Туре

Multiple Choice

Default value Fixed

Fixed

Options

[Fractional, Fixed]

GUI name

Capping option

Description

The capping atom is always along the broken bond vector.

The bond distance between the capping atom and the two atoms are obtained from covalent radii, let us call them D1H and D2H.

With option=Fractional the capping is on the bond vector with the fraction D1H/(D1H+D2H).

With the Fixed option it at the distance D1H from atom 1. A distance of zero always means the coordinate of the inside atom.

Committee

Туре

Block

Description

Settings for using the hybrid engine as a committee. The factors and region for each engine must be the same. When committee is enabled the standard deviation is also reported as the uncertainty.

Enabled

Type Bool

Default value No

Description Enable committee

Energy

Туре

Block

Description

This block is there to construct the energy.

DynamicFactors

Туре

Multiple Choice

Default value Default

Options

[Default, UseLowestEnergy, UseHighestEnergy]

GUI name

Adjust factors

Description

Default - use factors as set in the corresponding Term blocks;

UseLowestEnergy - set all factors to 0 except for that of the engine with the lowest energy, which is set to 1;

UseHighestEnergy - set all factors to 0 except for that of the engine with the highest energy, which is set to 1.

The last two options make sense only for non-QMMM hybrid calculation (that is, if the QMMM block is not present) and only when using engines whose energies can be compared directly.

Term

Type Block

Recurring

True

Description

This block is there to construct the energy term. Can have multiple occurrences

Charge

Type Float

Default value

0.0

Description Net charge to be used for this energy term.

EngineID

Type String

Description Identifier for the engine

Factor

Type Float

Default value 1.0

Description

Region

Type String

Description Identifier for the region

UseCappingAtoms

Туре

Bool

Default value

Yes

Description

Whether to use capping for broken bonds

Engine

Туре

Block

Recurring True

Description

The input for the computational (sub) engine. The header of the block determines the type of the engine. An optional second word in the header serves as the EngineID, if not present it defaults to the engine name. Currently it is not allowed to have a Hybrid engine as a sub engine.

GuessAttributesOnce

Туре

Bool

Default value

Yes

Description

If any ForceField subengines are defined, and if automatic atom typing is possible, then the atom typing is done at the level of the Hybrid engine, and not of the ForceField subengines. This ensures that the same atom types and charges are used in each subsystem, so that pair energy terms that should cancel, will cancel. If set to False, then for each energy term the atom types and charges of the subsystem will be determined separately.

QMMM

Туре

Block

Description

This block is there to identify the QMMM engines.

Embedding

Type Multiple Choice

Default value

Electrostatic

Options

[Mechanical, Electrostatic]

Description

Determines how the QM region is embedded into the MM region.

Mechanical embedding embedding can also be achieved using the Energy%Terms keywords, but the common case of a two region mechanical QM/MM embedding is easier to set up using this keyword.

MMCharge

Туре

Float

Default value 0.0

Description

Net charge to be used for the MM region.

MMEngineID

Type String

Description

Identifier for the MM engine

QMCharge

Type Float

Tiout

Default value 0.0

0.0

Description

Net charge to be used for the QM region.

QMEngineID

Type String

Description Identifier for the QM engine

QMRegion

Type

String

Description Identifier for the QM region. The rest of the system is considered the MM region.

UseCappingAtoms

Type Bool

Default value

Yes

Description

Whether to use capping for broken bonds.

RestartSubEngines

Type Bool

Default value

Yes

Description

Save all the results of the subengines and pass those in a next geometry step or MD step.

TweakRequestForSubEngines

Туре

Bool

Default value

Yes

Description

Only request what is really needed, gradients and charges.

CHAPTER

SIX

REFERENCES

1. M. Swart, AddRemove: A new link model for use in QM/MM studies. International Journal of Quantum Chemistry 91, 177 (2003) (https://doi.org/10.1002/qua.10463)