CASSANDRA

Computational Atomistic Simulation Software At Notre Dame for Research Advances

User Manual 1.1

Written by:

Edward J. Maginn, Jindal K. Shah, Eliseo Marin-Rimoldi, Sandip Khan, Ryan Gotchy Mullen, Neeraj Rai, Thomas Rosch, Andrew Paluch

Preface and Disclaimer

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This user manual is distributed along with the Cassandra software to aid in setting up various input files required for carrying out a Cassandra Monte Carlo simulation. Every effort is made to release the most updated and complete version of the manual when a new version of the software is released. To report any inconsistencies, errors or missing information, or to suggest improvements, send email to Edward Maginn (ed@nd.edu).

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Chapter 1

Introduction

Cassandra is an open source Monte Carlo package capable of simulating any number of molecules composed of rings, chains, or both. It can be used to simulate compounds such as small organic molecules, oligomers, aqueous solutions and ionic liquids. It handles a standard "Class I"-type force field having fixed bond lengths, harmonic bond angles and improper angles, a CHARMM or OPLS-style dihedral potential, a Lennard-Jones 12-6 potential and fixed partial charges. It does *not* treat flexible bond lengths. Cassandra uses OpenMP parallelization and comes with a number of scripts, utilities and examples to help with simulation setup.

Cassandra is capable of simulating systems in the following ensembles:

- Canonical (NVT)
- Isothermal-Isobaric (NPT)
- Grand canonical (μVT)
- Constant volume Gibbs (NVT-Gibbs)
- Constant Pressure Gibbs (NPT- Gibbs)

1.1 Distribution

Cassandra is distributed as a tar file cassandra.tar. You can unpack the distribution by running the command

> tar -xvf cassandra.tar

Upon successful unpacking of the archive file, the Cassandra directory will have a number of subdirectories. Please refer to the README file in the main Cassandra directory for a detailed information on each of the subdirectories.

Documentation - Contains this user guide and a document showing how molecular connectivity files are generated.

Examples - Contains example input files and short simulations of various systems in the above ensembles.

MCF - Molecular connectivity files for a number of molecules. These can be used as the basis for generating your own MCF files for molecules of interest.

Scripts - Useful scripts to set up simulation input files.

Src - Cassandra source code.

Chapter 2

Force Field

2.1 Bonds

Cassandra is designed assuming all bond lengths are fixed. If you wish to utilize a force field developed with flexible bond lengths, we recommend that you either use the nominal or "equilibrium" bond lengths of the force field as the fixed bond lengths specified for a Cassandra simulation or carry out an energy minimization of the molecule with a package that treats flexible bond lengths and utilize the bond lengths obtained from the minimization.

Table 2.1: Cassandra units for bonds

Parameter	Symbol	Units
Bond length	1	Å

2.2 Angles

Cassandra supports two types of bond angles:

- 'fixed' : The angle declared as fixed is not perturbed during the course of the simulation.
- 'harmonic' : The bond angle energy is calculated as

$$E_{\theta} = K_{\theta}(\theta - \theta_0)^2 \tag{2.1}$$

where the user must specify K_{θ} and θ_0 . Note that a factor of 1/2 is **not used** in the energy calculation of a bond angle. Make sure you know how the force constant is defined in any force field you use.

Table 2.2: Cassandra units for angles

Parameter	Symbol	Units
Nominal bond angle Bond angle force constant	$ heta_0 \ K_ heta$	$\frac{\mathrm{degrees}}{\mathrm{K/rad^2}}$

2.3 Dihedrals

Cassandra can handle four different types of dihedral angles:

• 'OPLS': The functional form of the dihedral potential is

$$E_{\phi} = a_0 + a_1 \left(1 + \cos(\phi) \right) + a_2 \left(1 - \cos(2\phi) \right) + a_3 \left(1 + \cos(3\phi) \right) \tag{2.2}$$

where a_0 , a_1 , a_2 and a_3 are specified by the user.

• 'CHARMM': The functional form of the potential is

$$E_{\phi} = a_0 (1 + \cos(a_1 \phi - \delta)) \tag{2.3}$$

where a_0 , a_1 and δ are specified by the user.

 \bullet 'harmonic': The dihedral potential is of the form:

$$E_{\phi} = K_{\phi}(\phi - \phi_0)^2 \tag{2.4}$$

where K_{ϕ} and ϕ_0 are specified by the user.

• 'none': There is no dihedral potential between the given atoms.

2.4 Impropers

Improper energy calculations can be carried out with the following two options:

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Table 2.3: Cassandra units for dihedrals

Functional Form	Parameter	Units
OPLS	a_0, a_1, a_2, a_3	kJ/mol
CHARMM	$a_0 \\ a_1 \\ \delta$	kJ/mol dimensionless degrees
harmonic	$K_{\phi} \ \phi_0$	K/rad ² degrees

- 'none': The improper energy is set to zero for the improper angle.
- 'harmonic': The following functional form is used to calculate the energy due to an improper angle

$$E_{\psi} = K_{\psi} \left(\psi - \psi_0 \right)^2 \tag{2.5}$$

where K_{ψ} and ψ_0 are specified by the user.

Table 2.4: Cassandra units for impropers

Parameter	Symbol	Units
Force constant	K_{ψ}	K/rad^2
Improper	ψ_0	degrees

2.5 Nonbonded

The nonbonded interactions between two atoms i and j are due to repulsion-dispersion interactions and electrostatic interactions (if any).

2.5.1 Repulsion-Dispersion Interactions

The repulsion-dispersion interactions can take one of the following forms:

• Lennard-Jones 12-6 potential (LJ):

$$\mathcal{V}(r_{ij}) = 4\epsilon_{ij} \left[\left(\frac{\sigma_{ij}}{r_{ij}} \right)^{12} - \left(\frac{\sigma_{ij}}{r_{ij}} \right)^{6} \right]$$
 (2.6)

where ϵ_{ij} and σ_{ij} are the energy and size parameters set by the user. For unlike interactions, different combining rules can be used, as described elsewhere. Note that this option only evaluates the energy up to a specified cutoff distance. As described below, analytic tail corrections to the pressure and energy can be specified to account for the finite cutoff distance.

• Cut and shift potential:

$$\mathcal{V}(r_{ij}) = 4\epsilon_{ij} \left[\left(\frac{\sigma_{ij}}{r_{ij}} \right)^{12} - \left(\frac{\sigma_{ij}}{r_{ij}} \right)^{6} \right] - 4\epsilon_{ij} \left[\left(\frac{\sigma_{ij}}{r_{cut}} \right)^{12} - \left(\frac{\sigma_{ij}}{r_{cut}} \right)^{6} \right]$$
 (2.7)

where ϵ_{ij} and σ_{ij} are the energy and size parameters set by the user and r_{cut} is the cutoff distance. This option forces the potential energy to be zero at the cutoff distance. For unlike interactions, different combining rules can be used, as described elsewhere.

• Cut and switch potential:

$$\mathcal{V}(r_{ij}) = 4\epsilon_{ij} \left[\left(\frac{\sigma_{ij}}{r_{ij}} \right)^{12} - \left(\frac{\sigma_{ij}}{r_{ij}} \right)^{6} \right] f \tag{2.8}$$

The factor f takes the following values:

$$f = \begin{cases} 1.0 & r_{ij} \le r_{on} \\ \frac{(r_{off}^2 - r_{ij}^2)(r_{off}^2 - r_{on}^2 + 2r_{ij}^2)}{(r_{off}^2 - r_{on}^2)^3} & r_{on} < r_{ij} < r_{off} \\ 0.0 & r_{ij} \ge r_{off} \end{cases}$$
(2.9)

where ϵ_{ij} and σ_{ij} are the energy and size parameters set by the user. This option smoothly forces the potential to go to zero at a distance r_{off} , and begins altering the potential at a distance of r_{on} . Both of these parameters must be specified by the user. For unlike interactions, different combining rules can be used, as described elsewhere.

• Mie potential (generalized form of LJ):

$$\mathcal{V}(r_{ij}) = \left(\frac{n}{n-m}\right) \left(\frac{n}{m}\right)^{\frac{m}{n-m}} \epsilon_{ij} \left[\left(\frac{\sigma_{ij}}{r_{ij}}\right)^n - \left(\frac{\sigma_{ij}}{r_{ij}}\right)^m \right]$$
 (2.10)

where ϵ_{ij} and σ_{ij} are the energy and size parameters and n and m are the repulsive and attractive exponents set by the user. This option allows for the use of a generalized LJ potential (for LJ, n=12 and m=6). Note that this option only evaluates the energy up to a specified cutoff distance. Both n and m can take on separate integer or float values set by the user. For unlike interactions, different combining rules can be used, as described elsewhere.

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• Mie cut and shift potential:

$$\mathcal{V}(r_{ij}) = \left(\frac{n}{n-m}\right) \left(\frac{n}{m}\right)^{\frac{m}{n-m}} \epsilon_{ij} \left[\left(\frac{\sigma_{ij}}{r_{ij}}\right)^n - \left(\frac{\sigma_{ij}}{r_{ij}}\right)^m \right] - \left(\frac{n}{n-m}\right) \left(\frac{n}{m}\right)^{\frac{m}{n-m}} \epsilon_{ij} \left[\left(\frac{\sigma_{ij}}{r_{cut}}\right)^n - \left(\frac{\sigma_{ij}}{r_{cut}}\right)^m \right]$$
(2.11)

where ϵ_{ij} and σ_{ij} are the energy and size parameters and n and m are the repulsive and attractive exponents set by the user. This option forces the potential energy to be zero at the cutoff distance (i.e. setting n = 12 and m = 6 provides the same potential as the LJ cut and shift option). For unlike interactions, different combining rules can be used, as described elsewhere.

• Tail corrections: If the Lennard-Jones potential is used, standard Lennard-Jones tail corrections are used to approximate the long range dispersion interactions

Table 2.5: Cassandra units for repulsion-dispersion interactions

Symbol	Units
ϵ/k_B	K
σ	Å
	,

2.5.2 Electrostatics

Electrostatic interactions are given by Coulomb's law

$$\mathcal{V}_{elec}(r_{ij}) = \frac{1}{4\pi\epsilon_0} \frac{q_i q_j}{r_{ij}}.$$
(2.12)

where q_i and q_j are partial charges specified by the user and placed on atomic positions given by r_i and r_j . In a simulation, the electrostatic interactions are calculated using either an Ewald summation or a direct summation using the minimum image convention. Note that the total energy that is printed out in the property file is extensive. Consequently, to obtain intensive energies, the printed energies must divided by the total number of molecules in the system.

Table 2.6: Cassandra units for coulombic interactions

Parameter	Symbol	Units
Charge	q	e

2.5. NONBONDED

Table 2.7: Summary of Cassandra units for input variables

Bond length	1	Å		
A	ngles			
Nominal bond angle	$ heta_0$	degrees		
Bond angle force constant	$\overset{\circ}{K_{ heta}}$	K/rad^2		
Dihed	lral angle			
OPLS	a_0, a_1, a_2, a_3	kJ/mol		
CHARMM	a_0	kJ/mol		
	a_1	dimensionless		
	δ	degrees		
harmonic	K_{ϕ}	K/rad^2		
	ϕ_0	degrees		
Impro Force constant Improper angle	per angle $K_{\psi} \ \psi_0$	$ m K/rad^2$ degrees		
Non	bonded			
Energy parameter	ϵ/k_B	K		
Collision diameter	σ	Å		
Charge	q	e		
Simulation	n Parameters			
Simulation box length		Å		
Volume		$ m \AA^3$		
Distances		Å		
Rotational width		degrees		
Temperature		K		
Pressure		bar		
Chemical potential		kJ/mol		
Energy		kJ/mol		

Chapter 3

Cassandra Basics

3.1 Flow Diagram

A flow diagram that overviews the setup for a Cassandra simulation is displayed on figure 3.1. This diagram employs two automation scripts located in the /Scripts/ directory: mcfgen.py and library_setup.py. These scripts are particularly useful when simulating large molecules. For details about how to use them, please refer to sections 5.2 and 5.3 of this user guide, and to the README files located in the subdirectories inside the directory /Scripts/.

3.2 Cassandra Simulation Setup

Once a system is identified, setting up a Cassandra simulation from scratch requires preparation of the following files.

- A molecular connectivity file (MCF) (*.mcf) containing the molecular connectivity information on bonds, angles, dihedrals, impropers and whether the molecule is composed of fragments. For information on the MCF file, please refer to section 4.2.
- An input file (*.inp) (See section 4.1)
- If the molecule is composed of fragments, then a fragment library file for each of the fragments is required. For instructions on how to generate these files, please refer to the section 5.3.

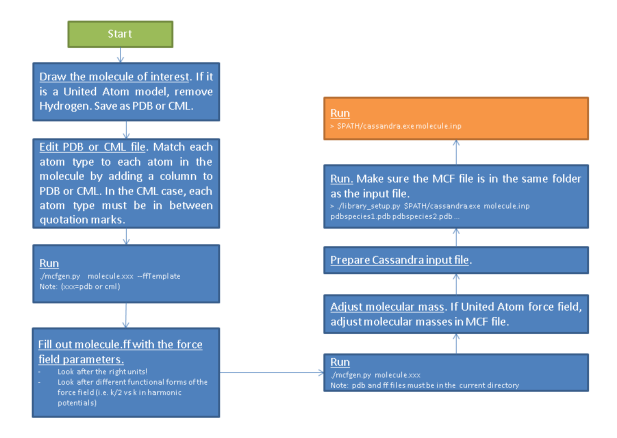


Figure 3.1: Flow diagram representing a typical setup of a Cassandra simulation

MCF files for united-atom models of methane, isobutane, dimethylhexane, cyclohexane and diethylether are provided in the MCF directory. Input files for NVT, NPT, GCMC and GEMC ensembles are located in the Examples directory which also contains fragment library files for a number of molecules simulated in these ensembles.

3.3 Cassandra File Preparation

3.3.1 MCF File

One MCF file is required for each unique species in a simulation. A species is defined as a collection of atoms associated with each other through bonds. Thus a molecule is a species as is an ion. If you wanted to simulate sodium sulfate, you would need separate MCF files for the sodium ion and the sulfate ion. MCF files can be created manually or by using the scripts provided with the code, as described in the section 5.2. Instructions for generating an MCF file can also be found in the Scripts/MCF_Generation/README file. We will collect MCF files submitted to us by users and will post them on the Cassandra website http://cassandra.nd.edu. If you have an MCF file you would like us to post, send it to ed@nd.edu.

3.3.2 Input File

An input file is required for a Cassandra simulation. The input file specifies conditions for the simulation and various keywords required for the simulation in a given ensemble. Please refer to Chapter 4.1 for further details.

3.3.3 Fragment Library Generation

Cassandra makes use of reservoir sampling schemes to sample correctly and efficiently various coupled intramolecular degrees of freedom associated with branch points and rings. Details may be found in our publication [?]. The idea is to decompose the entire molecule into fragments that are either branch points or ring groups, each coupled to other fragments via a single dihedral angle. Thus, the total number of fragments of a molecule is the sum of branch points and ring groups in the molecule. The neighboring fragments are connected by two common atoms present in each of the fragments. Note that the ring group contains all the ring atoms and those directly bonded to the ring atoms. For each fragment identified, Cassandra runs a pre-simulation in the gas phase to sample the intramolecular degrees of freedom. A library of a large number of these conformations are stored for use in an actual simulation.

The gas phase library generation has been automated with the script library_setup.py located in the Scripts/Frag_Library_Setup directory. Use the following command for generating the fragment library.

> python path_to_Scripts/Frag_Library_Setup/library_setup.py cassandra_executable input_filename mol1.pdb (mol1.cml) mol2.pdb (mol2.cml) ...

where input_filename is the name of the input file for the actual simulation and mol1.pdb mol2.pdb ... or mol1.cml mol2.cml ... correspond to the names of the pdb (or cml) files used to generated the MCF files. Make sure that if a file does not exist in the current working directory, its path relative to the current working directory is specified.

3.4 Running a Simulation

To launch a Cassandra simulation, run the following command:

> cassandra_executable input_filename

The executable will read input_filename and execute the instructions. Make sure that the required files (MCF, fragment library files) are located in the directories as given in the input file.

3.5 Restarting a Simulation

Restarting a simulation requires either a checkpoint file (*.chk produced by Cassandra) or a configuration file obtained from xyz files generated from a previous simulation. For the set up of these simulations, a script in Scripts/Read_Old is provided. Detailed instructions are contained in the README file in this directory.

3.6 Cassandra Output Files

Cassandra generates several output files which can be used for later analysis. All have as a prefix the Run_Name specified in the input file. See Chapter 4.1 for details. The type of output is specified by the file name suffix. The following are generated:

- Log file (*.log): Contains basic information on what the run is, timing information and reports the various parameters specified by the user. A complete copy of the input file is reproduced. Other important information includes the move acceptance rates. You can use the log file to keep track of what conditions were simulated.
- Coordinate file (*.box#.xyz): For each box in the system, a set of xyz coordinates are written out with a frequency specified by the user (Coord_Freq). The file has as a header the number of atoms in the box. Following this, the atomic coordinates of molecule 1 of species 1 are written, then the coordinates of molecule 2 of species 1 are written, etc. After all the coordinates of the molecules of species 1 are written, the coordinates of the molecules of species 2 are written, etc. You can use this file to do all your structural analysis and post processing.

Note that if you generate your initial configuration using the make_config command, the first "snapshot" of the coordinate file will contain the initial configuration of all the species in the system for a given box. You can use this configuration to check on whether the initial configuration is reasonable, or use it as an input to other codes. Note that the initial configuration will be generated using a configurational biased scheme, so it may be a better starting configuration that if you used other methods.

- Checkpoint file (*.chk): A checkpoint file is written every Coord-Freq steps. This can be used to restart a simulation from this point using all of the same information as the run that was used to generate the checkpoint file. To do this, you must use the checkpoint restart option (see Chapter 4.1). It will basically pick up where the simulation left off, using the same random number seed, maximum displacements, etc. This is useful in case your job crashes and you want to continue running a job. You can also use the checkpoint file to start a new simulation using the configuration of the checkpoint file as an initial configuration and the optimized maximum displacements. To do this, use the script read_old.py. You will need to set a new random number seed if you do this. See the documentation in Chapter 4.1 for more details.
- H-matrix file (*.box#.H): This file is written to every Coord_Freq MC steps. The first line is the box volume in angstrom³. The next three lines are the box coordinates in angstrom in an H-matrix form. Since Cassandra only supports cubic boxes at the moment, this is just a diagonal and symmetric matrix, but is included here for later versions that will enable non-orthogonal boxes. After this, a blank line is written. The next line is the box number, and the final line(s) is(are) the species ID and number of molecules for that species in this box. If there are three species, there will be three lines. This output is repeated every Coord_Freq times. This file allows you to compute the density of the box during constant pressure simulations.
- Property file (*.box#.prp#): This file lists the instantaneous thermodynamic and state properties for each box. Note that you can have more than one property file (hence the# after 'prp') and more than one box (also why there is a # after 'box'). The user specifies which properties are to be written and in what order, and these are then reproduced in this file. The file is written to every Prop_Freq steps. A header is written to the first two lines to designate what each property is. You may use this file to compute thermodynamic averages.

Chapter 4

Files Required to Run Cassandra

4.1 Simulation Input File

This is a required file that is given as an argument to the Cassandra executable. You must generate this by hand, but you can use the input files in the Examples directory as a guide. The input file contains a number of keywords that define simulation parameters and thermodynamic state point for the simulation. A keyword is identified by a '#' while comments follow a '!'. Note that the order of the keywords in the input file is immaterial, but the format of the arguments of the keyword are important and are explained below, along with a complete listing of all keywords.

4.1.1 Run Name

Run_Name - Simulation run name Character*120

The run name is specified on the next line following the keyword. This name is used as a prefix for all the files produced by the simulation. For example,

Run_Name
dee.out

Cassandra will then use dee.out as prefix for all output files created.

4.1.2 Simulation Type

Sim_Type Character*120

Sets the ensemble (and thus the suite of moves) of a Cassandra simulation. Currently the following ensembles are supported:

- NVT_MC (Canonical ensemble)
- NVT_MIN (Canonical ensemble in which minimization is carried out)
- NPT_MC (Isothermal-isobaric ensemble)
- GCMC (Grand canonical ensemble)
- GEMC (Gibbs ensemble)
- NVT_MC_Fragment (Canonical ensemble simulation of a non-ring fragment)
- NVT_MC_Ring_Fragment (Canonical ensemble simulation of a ring fragment)

Note that NVT_MC_Fragment and NVT_MC_Ring_Fragment are used only for the fragment generation and are not used in the normal simulation. For example,

```
# Sim_Type
NPT_MC
```

will run an NPT MC simulation.

4.1.3 Number of species

Nbr_Species Integer

Total number of species in the simulation. For ionic systems, each ion is counted as a separate species. For example, for a mixture of two species, use the following:

```
# Nbr_Species
```

2

4.1.4 VDW Style

VDW_Style

Character(i,1) Character(i,2) Real(i,3) Real(i,4)/Logical(i,4)

This keyword specifies the functional form of repulsion dispersion interactions to be used and if tail corrections are added for the box i. One line is required for each box. The options for Character(i,1) are "LJ" for a Lennard-Jones 12-6 potential or "None" if you wish to turn off all repulsion-dispersion interactions. Character(i,2) and Real(i,3) are specified only if Character(i,1) is set to "LJ". Character(i,2) describes the truncation scheme used for the Lennard-Jones potential. Options are "cut", "cut_tail", "cut_switch", "cut_shift", "mie", and "mie_cut_shift". Refer to Chapter 2 for the functional forms. The other parameters Real(i,3) and Real(i,4)/Logical(i,4) depend on the selection of Character(i,2) as described below:

cut: This option cuts the potential off at the distance specified by (Real(i,3)). The fourth parameter is omitted.

For example, to simulate one box with a 14 Å cutoff specify the following:

```
# VDW_Style LJ cut 14.0
```

Similarly, for a two box simulations such as used in the Gibbs ensemble where both boxes have a 14 Å cutoff, use the following:

```
# VDW_Style
LJ cut 14.0
LJ cut 14.0
```

cut_tail: This options cuts the potential off at a distance corresponding to (Real(i,3)) and applies analytic tail corrections to the energy and pressure. An optional fourth argument (Logical(i,4)) can be set to 'TRUE' or 'true', in which case Real(i,3) is ignored and the cutoff distance is always set to half of the simulation box length. The cutoff will change during the course of the simulation when attempting volume moves. This option is provided to enable reproduction of literature simulations that use a cut off distance of half the simulation box length, but its use is highly discouraged.

For example, to simulate one box with a 14 Å cutoff using tail corrections, specify the following:

```
# VDW_Style
LJ cut_tail 14.0
```

For a two box simulation where the first box has a 14 Å cutoff and the second one has a 20 Å cutoff, use the following:

```
# VDW_Style
LJ cut_tail 14.0
```

LJ cut_tail 20.0

cut_switch: This option cuts the potential off and smoothly brings the potential to zero using a spline. The potential is cutoff and the spline turned on at a distance specified by Real(i,3) (r_{on} in Eq 2.8) and the potential goes to zero at a distance specified by Real(i,4) (r_{off} in Eq 2.8).

For example, a one box simulation using the cut_switch option could be specified as follows:

```
# VDW_Style
LJ cut_switch 12.0 14.0
```

In this case, the Lennard-Jones potential would end at 12.0 Å and be smoothly taken to zero at 14.0 Å. Obviously, $r_{on} < r_{off}$ or Real(i,3) < Real(i,4).

cut_shift: This option cuts the potential off at a distance specified by Real(i,3) and shifts the entire potential so that at this distance the potential is zero. The fourth parameter Real(i,4)/Logical(i,4) is omitted. The functional form of this potential is given in eq 2.7.

To perform a two box simulation with a cut_shift option in which both boxes have a 10.5 Å cutoff, use the following:

```
# VDW_Style
LJ cut_shift 10.5
LJ cut_shift 10.5
```

mie: This option cuts the potential off at a distance specified by Real(i,3). It must be used with the keyword "# Mie_Nonbond" (see section 4.1.7) which specifies exponents n and m for each species. The fourth parameter Real(i,4)/Logical(i,4) is omitted. The functional form of this potential is given in eq 2.10.

For example, a one box simulation using the mie option can be specified as follows: # VDW_Style LJ mie 14

mie_cut_shift: This option cuts the potential off at a distance specified by Real(i,3) and shifts the entire potential so that at this distance the potential is zero. It must be used with the keyword "# Mie_Nonbond" (see section 4.1.7) which specifies exponents n and m for each species. The fourth parameter Real(i,4)/Logical(i,4) is omitted. The functional form of this potential is given in eq 2.11.

A two box simulation using the mie_cut_shift option with a 14 Å cutoff can be specified as follows:

```
# VDW_Style
LJ mie_cut_shift 14
LJ mie_cut_shift 14
```

Note: For all options, cutoff distances must be less than or equal to the shortest edge length of a simulation box.

4.1.5 Charge Style

Charge_Style Character(i,1) Character(i,2) Real(i,3) Real(i,4)

Cassandra allows the use of fixed partial charges on atomic centers using a Coulomb potential of the form given in Eq 2.12. To specify this option for box i, set Character(i,1) to "coul". For this option, Character(i,2) can be set to either "Ewald" if you want to use an Ewald sum to compute Coulombic interactions or it can be set to "cut", in which case the Coulombic interactions will be cut off and the long range interactions ignored. For the Ewald option, Real(i,3) is the real space cutoff distance and Real(i,4) specifies the accuracy of the Ewald summation. A reasonable value for the accuracy is 10^{-5} . Note that the number of reciprocal vectors for the Ewald summation is determined in the code based on the accuracy parameter. For more details, see the paper by Fincham [?].

For example,

```
# Charge_Style
coul Ewald 12.0 1E-5
```

will use the Ewald sum for a single box. The real space cutoff will be 12 Å and the accuracy will be 10^{-5} . If you have two boxes, like in a Gibbs ensemble calculation, then you could use the following:

```
# Charge_Style
coul Ewald 12.0 1E-5
coul Ewald 30.0 1E-5
```

This will use an Ewald sum for both boxes. In the first box, the real space cutoff will be 12 Å while in the second box a larger cutoff of 30 Å will be used. Note: When performing Gibbs ensemble simulations of vapor-liquid equilibria, the vapor box is often much larger than the liquid box. In this case, you will want to use a longer real space cutoff for the larger vapor box to avoid using too many reciprocal space vectors. Also note that the real space cutoffs must always be less than or equal to half of the shortest edge length of a simulation box.

If you do not wish to use a Coulomb potential (for example, your model has no partial charges), you still must specify # Charge_Style. In this case, set Character(i,1) to "NONE". If "NONE" is selected for Character(i,1) then Character(i,2), Real(i,3) and Real(i,4) are omitted.

For example,

```
# Charge_Style
NONE
```

should be used if you have no partial charges and are simulating a single box. A two box simulation with no partial charges would be specified as

```
# Charge_Style
NONE
NONE
```

Note: If the cutoff in VDW_Style is set to half of the simulation box length, any cutoff distance specified in the Charge_Style section will default to the half of the simulation box length. In the case of Ewald summation, however, the accuracy will be the same as Real(i,4).

4.1.6 Intramolecular Scaling

```
# Intra_Scaling
Real(i,1) Real(i,2) Real(i,3) Real(i,4)
Real(i,5) Real(i,6) Real(i,7) Real(i,8)
```

This keyword sets the intramolecular scaling for 1-2, 1-3, 1-4 and 1-N interactions within a given species. 1-2 means interactions between a given atom 1 and another atom 2 directly bonded to it, 1-3 means interactions between atom 1 and other atoms 3 separated from atom 1 by exactly two bonds, etc. The first line corresponds to the VDW scaling: Real(i,1) Real(i,2) Real(i,3) Real(i,4) apply to 1-2, 1-3, 1-4 and 1-N interactions, where N corresponds to all atoms separated from atom 1 by more than three bonds. The second line corresponds to the Coulomb scaling: Real(i,5) Real(i,6) Real(i,7) Real(i,8) apply to 1-2, 1-3, 1-4 and 1-N interactions. These lines are repeated for each species in the simulation. Note that intramolecular scaling applies to all the boxes in the simulation.

For example,

```
# Intra_Scaling
0.0 0.0 0.5 1.0
0.0 0.0 0.5 1.0
```

would turn off 1-2 and 1-3 interactions, would scale the VDW and Coulombic interactions for 1-4 atoms by 50% and would use full interactions for all other atom pairs in the species.

If you had two species in the simulation and wanted the same intramolecular scaling as above, you would specify

```
# Intra_Scaling
0.0 0.0 0.5 1.0
0.0 0.0 0.5 1.0
0.0 0.0 0.5 1.0
0.0 0.0 0.5 1.0
```

In the absence of the # Intra_Scaling keyword, default values of 0.0, 0.0, 0.5 and 1.0 will be used for 1-2, 1-3, 1-4 and 1-N for VDW and Coulomb interactions for all the species in the simulation. If # Charge_Scaling is set to "NONE", you must still specify the Intra_Scaling for Coulombic interactions, but no interactions will

be computed.

4.1.7 Mie Nonbond

```
# Mie_Nonbond Int(i,1) Int(j,2) Real(i,j,3) Real(i,j,4) * (N species * N species) lines
```

This keyword should only be used if VDW_Style is set to "mie" or "mie_cut_shift". Int(i,1) and Int(j,2) specify the index for each species pair combination. Real(i,j,3) and Real(i,j,4) are the corresponding n and m exponents, respectively.

For example, a system with two species interacting with a LJ potential can be specified as follows:

```
# Mie_Nonbond
1 1 12 6
1 2 12 6
```

2 1 12 6

2 2 12 6

Note that this keyword sensitive to the correct species pair ordering. Also note that exponents for species pairs i and j, along with j and i, are both specified. For N species in a system, (N * N) lines should be specified.

Likewise, for a system including three species, with species 1 interacting with n = 9 and m = 6 only with itself and all other pair combinations with n = 12 and m = 4, the following lines can be specified:

```
# Mie_Nonbond
```

1 1 9 6

1 2 12 4

1 3 12 4

2 1 12 4

2 2 12 4

2 3 12 4

3 1 12 4 3 2 12 4

3 3 12 4

4.1.8 Mixing Rule

Mixing_Rule

Character

Sets the method by which Lennard-Jones interactions between unlike atoms are calculated. Acceptable options are "LB" for Lorentz-Berthelot and "geometric" for geometric. If this keyword is missing, "LB" is used as default.

4.1.9 Starting Seed

Seed_Info Integer(1) Integer(2)

Inputs for the starting random number seeds for the simulation. Note that Cassandra uses a random number generator proposed by L'Ecuyer [?], which takes five seeds to calculate a random number, out of which three are defined internally while two Integer(1) and Integer(2) are supplied by the user. When a 'checkpoint' file is used to restart a simulation (see # Start_Type below), the user supplied seeds will be overwritten by those present in the checkpoint file. If # Start_Type is set to 'read_old', then the seeds specified in the input file are used.

As an example,

Seed_Info 1244432 8263662

is an acceptable way of specifying the seeds. Note that two independent simulations can be run using the same input information if different seeds are used. If two simulations having exactly the same input information and the same seeds are run, the results will be identical.

4.1.10 Minimum Cutoff

Rcutoff_Low Real(1)

Sets the minimum distance Real(1) in Å such that any MC move bringing two sites closer than this distance will be immediately rejected. It avoids numerical problems associated with random moves that happen to place atoms very close to one another such that they will have unphysically strong repulsion or attraction. This distance must be less than the intramolecular distance of all atoms in a species which are not bonded to one another. For models that use dummy sites without explicitly defining bonds between dummy and atomic sites of the molecules (for example, the TIP4P water model), it is important that the minimum distance is set to be less than the shortest distance between any two sites on the molecule. For most systems, 1 Å seems to work OK, but for models with dummy sites, a shorter value may be required.

4.1.11 Pair Energy Storage

Pair_Energy Logical(1)

Cassandra can use a time saving feature in which the energies between molecules are stored and used during energy evaluations after a move, thereby saving a loop over all molecules. This requires more memory, but it can be faster. The default is to not use this feature. If you wish to use this, set Logical(1) to 'TRUE' or 'true'.

4.1.12 Molecule Files

Molecule_Files Character(i,1) Integer(i,2)

This specifies the name of the molecular connectivity file (*.mcf) and the maximum total number of molecules of a given species specified by this MCF file. A separate line is required for each species present in the simulation. Character(i,1) is the name of the MCF file for species i. Integer(i,2) denotes the maximum number of molecules expected for the species.

For example

```
# Molecule_Files
butane.mcf 100
hexane.mcf 20
octane.mcf 5
```

specifies that there are three different species, and the MCF files state the names of the files where information on the three species can be found. Species 1 is butane, species 2 is hexane and species 3 is octane. There can be a maximum of 100 butane molecules, 20 hexane molecules and 5 octane molecules in the total system. The maximum number of molecules specified here will be used to allocate memory for each species, so do not use larger numbers than are needed.

4.1.13 Simulation Box

Box_Info Integer(1) Character(i,2) Real(i,3)

This keyword sets parameters for the simulation boxes. Integer(1) specifies the total number of boxes

in the simulation. For now, Gibbs ensemble simulations must have two boxes. Character(i,2) is the shape of the i^{th} simulation box. Right now, the only supported box shape in Cassandra is cubic, so use the keyword "CUBIC". Real(i,3) is the length of the box edges for a cubic box in Å. Information for additional boxes is provided in an analogous fashion and is separated from the previous box by a blank line. For a two box simulation, box information is given as:

```
# Box_Info
2
CUBIC
30.0
CUBIC
60.0
```

This will construct two cubic boxes. The first will be 30 X 30 X 30 cubic \mathring{A} and the second will be 60 X 60 X 60 cubic \mathring{A} .

4.1.14 Temperature

Temperature_Info $Real(1) Real(2) \dots$

The temperature setpoint in Kelvin Real(1) Real(2) ... for simulation boxes 1, 2, ...

4.1.15 Pressure

Pressure_Info

 $Real(1) Real(2) \dots$

The pressure setpoint in bar Real(1) Real(2) ... for simulation boxes 1, 2, If the simulation type does not require an input pressure (e.g., NVT), this command will be ignored.

4.1.16 Chemical Potential

Chemical_Potential_Info

 $Real(1) Real(2) \dots$

The chemical potential setpoint (shifted by a species-specific constant) in kJ/mol of each insertable species. Each chemical potential will be assigned in the order species appear in the Molecule_Files section. This keyword is only used for grand canonical simulations. See Eq. (6.31) for more information.

4.1.17 Move Probabilities

Move_Probabilities This section specifies the probabilities associated with different types of MC moves to be performed during the course of simulation. Please ensure that the move probabilities add up to 1.0. An error will be generated if this is not the case. All the headers are optional but an error will be produced if a move is required (for example, volume fluctuations in an NPT simulation) and the move is not specified.

Translation

Prob_Translation

Real(1)

Real(i,j) *One line required for each box, and one value required for each species on each line.

Real(1) is the probability of performing a center of mass translation move. For each box i, the maximum displacement of species j is specified by Real(i,j).

For example, if you have two species and two boxes, you would specify the translation probability as

Prob_Translation

0.25

2.0 2.5

10.0 11.0

This will tell Cassandra to attempt center of mass translations 25% of the total moves. For box 1, the maximum displacement will be 2.0 Å for species 1 and 2.5 Å for species 2. For box 2, the maximum displacement for species 1 will be 10.0 Å and it will be 11.0 Å for species 2. Note that attempted moves will occur with equal probability for a give box, but attempts for a species are proportional to their mole fraction in the box.

Rotation

Prob_Rotation

Real(1)

Real(i,j) *One line required for each box, and one value required for each species on each line.

The probability of performing a rotation move is specified by Real(1) while Real(i,j) denotes the maximum rotational width for species j in box i in degrees. If you are only simulating spherical molecules (such as Lennard-Jones particles), then do not use this keyword. If you are simulating a multi-species system where some of the species have rotational degrees of freedom and some species are spheres, then specify an appropriate value of Real(i,j) for the species having rotational degrees of freedom, and set Real(i,j) equal to zero for the spherical species. Linear molecules are a special case, where rotation is handled in Eulerian space. If you have a linear molecule such as carbon monoxide, specify any non-zero value for Real(i,j). Cassandra

will properly sample the rotational degrees of freedom but will not use the value set by Real(i,j). Note that spherical molecules are not considered when choosing which species to perform a rotational move on.

For example, if you are simulating two species in two boxes and if the first species has rotational degrees of freedom while the second is spherical, you would specify the rotational probability as

Prob_Rotation 0.25 30.0 0.0 180.0 0.0

This will tell Cassandra to perform rotational moves 25% of the total moves. For box 1, the maximum rotational width will be 30° for species 1 and 0.0° for species 2. For box 2, the maximum rotational width will be 180° for species 1 and 0.0° for species 2. Note that, since the maximum rotational width of species 2 is set to 0° in both boxes, no rotational moves will be attempted on species 2.

For a single box simulation with three species such that the first species has rotational degree of freedom, the second is a linear molecule and the third species is spherical, you would specify

Prob_Rotation 0.25 30.0 10.0 0.0

This will tell Cassandra to attempt rotational move 25% of the total moves. The maximum rotational width for species 1 is 30° and that for species 2 is 10.0° . Since the species 2 is a linear molecule, its rotation will be attempted in Eulerian angles and Cassandra will not use this value. Since the rotational width is set equal to 0° for species 3, no rotational moves will be attempted for this species.

Regrowth

Prob_Regrowth

Real(1)

 $Real(i,2)^*$ One for each species

A regrowth move consists of deleting part of the molecule randomly and then regrowing the deleted part via a configurational bias algorithm. This can result in relatively substantial conformational changes for the molecule, but the cost of this move is higher than that of a simple translation or rotation. The probability of attempting a regrowth move is specified by Real(1) while Real(i,2) specifies the relative probability of performing this move on species i. For monatomic species, Real(i,2) is set to zero. Note that the user needs to ensure that the relative probabilities add up to 1 otherwise Cassandra will display an error and quit.

For example, if you are simulating three species of which the first species is monatomic, you would specify the following:

Prob_Regrowth

0.3

0.0 0.7 0.3

This will tell Cassandra to attempt regrowth move 30% of the total moves. The relative probabilities of performing regrowth moves on species 1, 2 and 3 are 0.0, 0.7 and 0.3 respectively.

Volume

Prob_Volume

Real(1)

Real(2,i) * One line required for each box except for GEMC-NVT

Sets the probability of volume displacement moves. This flag is required for NPT, GEMC-NPT and GEMC-NVT simulations. Do not specify for any other simulation type. Real(1) is the relative probability of attempting a box volume change. Note that volume changes are bold and expensive moves and should be attempted infrequently. This probability should normally not exceed 0.05 and values from 0.01-0.03 are typical. Real(2,i) specifies the maximum volume displacement in \mathring{A}^3 for box i. If you are simulating a two box system, a value of Real(2,i) is required for each box on separate lines. Note that the exception to this is for a GEMC-NVT simulation, where there are two boxes but the volume moves are coupled. In this case, only a single value of Real(2,i) is specified. The default is to change the box volume by random amounts up to the maximum value specified by Real(2,i). For example, if you are simulating a liquid with a single box in the NPT ensemble, you would specify the following:

Prob_Volume

0.02

300

This will tell Cassandra to attempt volume moves 2% of the total moves. The box volume would be changed by random amounts ranging from -300 ų to +300 ų. For a liquid box 20 Å per side, this would result in a maximum box edge length change of about 0.25 Å, which is a reasonable value. Larger volume changes should be used for vapor boxes. If you wish to perform a GEMC-NPT simulation, you might specify the following:

Prob_Volume

0.02

300

5000

This will tell Cassandra to attempt volume moves 2% of the total moves. The first box volume (assumed here to be smaller and of higher density, such as would occur if it were the liquid box) would be changed by random amounts ranging from -300 Å³ to +300 Å³. The second box volume would be changed by random amounts ranging from -5000 Å³ to +5000 Å³. As with all move probabilities, you can experiment with making larger or smaller moves. Note that if the # Run_Type is set to 'Equilibration', Cassandra will attempt

to optimize the magnitude of the volume change to achieve about 50% acceptance rates.

Insertion and Deletion Moves

Prob_Insertion

Real(1) Character(i,1)Character(i,2)

This flag is set only for GCMC simulations. Real(1) sets the probability of attempting insetion moves. If there is more than one species, each is chosen for an insertion attempt with equal probability. Character(i,1) and Character(i,2) control the manner with which the insertions are carried out for each species i. Right now, Cassandra only uses one insertion method (a reservoir sampling approach). Later versions will have other options. So for now, the only option is to set Character(i,1) equal to 'insertion method'. If the species can be inserted or deleted, set Character(i,2) equal to 'reservoir'. If the species is 'non-volatile' and should not be inserted or deleted or should stay in its original box, then set Character(i,2) equal to 'none'. Then whichever box that species starts in, it will remain there for the whole simulation. You must repeat these flags for each species i. For example, if you are performing a GCMC simulation with two species that can be inserted, you might specify the following

Prob_Insertion 0.1 insertion method reservoir insertion method reservoir

This will tell Cassandra to attempt insertions 10% of the total moves and both species will be inserted using the reservoir insertion method.

Prob_Insertion 0.1 insertion method reservoir insertion method none

This will tell Cassandra to attempt insertions 10% of the total moves. Only species 1 will be inserted, while species 2 will not get inserted.

Prob_Deletion Real (1)

Sets the relative probability of deletion during the course of a simulation. Each exchangeable species is

randomly chosen and a deletion move is attempted on a randomly chosen molecule of this species. If a species has an insertion method 'none', no attempt is made to delete it. You must specify the same deletion probability as the insertion probability to satisfy microscopic reversibility. If you fail to do this, Cassandra will give an error and quit.

Prob_Swap

Real(1)

Character(i,1)

Character(i,2)

This keyword is set only for a GEMC simulation to enable transfer of species between two boxes. Real(1) sets the relative probability of attempting transfer of a molecule from one box to the other. During the swap, the donor and receiving boxes are chosen randomly. The species chosen for transfer is selected according to its overall mole fraction which is calculated only for the species that can be exchanged between boxes. Thus, species that are "non-volatile" are not included while computing the mole fractions. A molecule is then chosen randomly for transfer.

Similar to the # Prob_Insertion section, Character(i,1) and Character(i,2) describe the manner in which the swap is carried out for each species i. At present, the only option is to set Character(i,1) equal to 'insertion method'. If the species can be swapped, set Character(i,2) equal to 'reservoir'. If the species is not to be transferred between boxes, then set Character(i,2) to 'none'. Then whichever box that species starts in, it will remain in that box for the whole simulation. These flags are to be repeated for each species i. For example, while performing a GEMC simulation for three species the first two of which are exchanged while the third is not, you might specify the following:

Prob_Swap 0.1 insertion method reservoir insertion method reservoir insertion method none

This will tell Cassandra to attempt swap moves 10% of the total moves. Attempts will be made to transfer species 1 and 2 between available boxes while molecules of species 3 will remain in the boxes they are present in at the start of the simulation.

Flip Move

Prob_Ring

Real(1) Real(2)

This keyword is used when flip moves are to be attempted to sample bond angles and dihedral angles in a ring fragment. For more details on this move, see our publication [?]. Note that this keyword is used only in input files that generate configuration libraries of ring moieties. The input file of the actual simula-

tion would involve the "# Prob_Regrowth" keyword. The relative probability of attempting a flip move is specified by Real(1) while the maximum angular displacement in degrees for the move is given by Real(2). For example, if the flip is to be attempted 30% of the time and the maximum angular displacement for the move is 20° specify the following:

Prob_Ring
0.30 20.0

Done_Probability_Info

This is a required keyword that marks the end of the section for specifying move probabilities. It must occur after # Move_Probability and all the move probabilities must be specified between these two keywords. Once Cassandra reads # Done_Probability_info, it checks to make sure the probabilities sum to unity. If not, an error will be given.

4.1.18 Start Type

Start_Type Character(1)

This keyword specifies whether Cassandra generates an initial configuration or uses a previously generated configuration to start a simulation. *Character(1)* takes one of the three options: 'make_config', 'checkpoint' or 'read_old' and it determines what configuration is used to start a simulation.

When 'make_config' is used as the start type, Cassandra will generate an initial configuration. With this option, additional information is required on the number of molecules of each species in every box and is specified as follows:

make_config

Integer(j,k) * One line for each species and one entry on each line for each box

where Integer(j,k) represents the number of molecules of species j in box k. Thus, for example, to generate an initial configuration for two species in two boxes such that the numbers of molecules of species 1 in box 1 and 2 are 100 and 50 respectively and those for species 2 are 75 and 25 respectively, the input file must contain the following:

Start_Type
make_config
100 50
75 25

During the course of a simulation, Cassandra periodically generates a checkpoint file (*.chk) containing information about the total number of translation, rotation and volume moves along with the random number seeds and the coordinates of all the molecules and their box number at the time the file is written. Cassandra provides the capability of restarting from this state point in the event that a simulation crashes

or running a production simulation from an equilibrated configuration. For this purpose, in addition to the 'checkpoint' keyword, additional information in the form of the name of the checkpoint file Character(2) is required in the following format:

checkpoint

Character(2)

For example, to continue simulations from a checkpoint file 'methane_vle_T148.chk', you might specify:

```
# Start_Type
checkpoint
methane_vle_T148.chk
```

Cassandra also provides a 'read_old' option to make use of just the coordinates of molecules to start a simulation. For example, a configuration generated at a lower temperature may be used to jump start a simulation at a higher temperature. When the 'read_old' option is used, additional information in the form of the file names Character(k,3) is required as shown below:

$read_old$

Character(k,3) * One line for each box

For example, to start a GEMC simulation simulation using the configurations of the two boxes, you might specify:

```
# Start_Type
read_old
box1.readold
box2.readold
```

This will tell Cassandra to use the configurations of the two boxes stored in box1.readold and box2.readold to start a simulation. Note that configurations of the boxes can be easily extracted from the checkpoint file using the utility read_old.py provided in Scripts/Read_Old.

4.1.19 Run Type

```
# Run_Type
Character(1) Integer(2) Integer(2)
```

This keyword is used to specify whether a given simulation is an equilibration or a production run. For an equilibration run, the maximum translational, rotational and volume widths (for an NPT or a GEMC simulation) are adjusted to achieve 50% acceptance rates. During a production run, the maximum displacement width for different moves are held constant.

Depending on the type of the simulation, Character(1) can be set to either "Equilibration" or "Production".

For an Equilibration run, Integer(1) denotes the number of MC steps performed for a given thermal move before the corresponding maximum displacement width is updated. Integer(2) is the number of MC volume moves after which the volume displacement width is updated. This number is optional if no volume moves are performed during a simulation (for example in an NVT or a GCMC simulation). When the run type is set to Production, the MC moves refer to the frequency at which the acceptance ratios for various moves will be computed and output to the log file. These acceptance rates should be checked to make sure proper sampling is achieved.

For an NPT equilibration run in which the widths of the thermal move are to be updated after 100 MC moves and maximum volume displacements after 10 volume moves, specify the following:

```
# Run_Type
Equilibration 100 10
```

For an NVT production run in which the acceptance ratios of various thermal moves are printed to the log file after every 250 MC steps of a given thermal move, use the following:

```
# Run_Type
Production 250
```

4.1.20 Frequency

Simulation_Length_Info

Units Character(1)
Prop_Freq Integer(2)
Coord_Freq Integer(3)
MCsteps or Stop Integer(4)
Done_Simulation_Length_Info

This section specifies the frequency at which thermodynamic properties and coordinates are output to a file. Character(1) determines the method by which the simulation is terminated and data is output. If Character(1) is to 'Minutes', then the simulation stops after the specified time in minutes. The format for this option is given below:

Units Minutes

Prop_Freq Integer(2) Coord_Freq Integer(3) Stop Integer(4)

With this option, thermodynamic quantities are output every Integer(2) minutes, coordinates are written to the disk every Integer(3) minutes and the total simulation time is specified in minutes by Integer(4). For example, to run a simulation for 60 minutes such that thermodynamic quantities are written every minute and the coordinates are output every 10 minutes, use the following:

Simulation Length_Info
Units Minutes
Prop_Freq 1
Coord_Freq 10
Stop 60
Done_Simulation_Length_Info

Similar to # Move_Probabilities section, the end of the simulation length section always includes the # Done_Simulation_Length_Info line.

Simulations can also be run for a given number of MC steps. To enable this feature, *Character(1)* is set to 'Steps'. Additional information is required and is given in the following format:

Units Steps

Prop_Freq Integer(2) Coord_Freq Integer(3) MCsteps Integer(4)

With this option, thermodynamic quantities are output every Integer(2) MC steps, coordinates are written at a frequency of Integer(3) MC steps and the simulation terminates after Integer(4) steps. Note that an MC step is defined as a single MC move, regardless of type and independent of system size.

To run a simulation of 50,000 steps such that thermodynamic quantities are printed every 100 MC steps and coordinates are output every 10,000 steps, use the following:

Simulation Length_Info
Units Steps
Prop_Freq 100
Coord_Freq 10000
MCsteps 50000
Done_Simulation_Length_Info

4.1.21 Average

Average_Info Integer(1)

This section specifies how thermodynamic quantities are output. At present, Cassandra writes instantaneous values of thermodynamic quantities at a frequency given by either *Nthermofreq* or *thermofreq* in the # Frequency_Info section. *Integer(1)* is set to 1 for this purpose. Later versions of Cassandra will have the ability to output block averages as well. Thus, you will specify the following section in your input file:

```
# Average_Info
1
```

4.1.22 Property Output

Property_Info Integer(i) Character(i,j) * One line for each property

This section provides information on the properties that are output. More than one section is allowed for multiple boxes. In this case, each section is separated by a blank line. Integer(i) is the identity of the box for which the properties are desired. Character(i,j) is the property that is to be output. Each property is specified on a separate line. At present, the acceptable entries include:

Energy_Total: Total energy of the system (Extensive) in kJ/mol

Energy_LJ: Lennard-Jones energy of the system in kJ/mol Energy_Elec: Electrostatic energy of the system in kJ/mol

 $\label{lem:energy_intra} \textbf{Energy_Intra}: \ \ \textbf{Total intramolecular energy of the system including bonded and non-bonded interactions in kJ/mol}$

Enthalpy: Enthalpy of the system (Extensive) kJ/mol

Pressure: Pressure of the system in bar Volume: Volume of the system in $Å^3$ Nmols: Number of molecules of a species Density: Density of a species in $\#/\mathring{A}^3$

For example, if you would like total energy, volume and pressure of a one box system to be written, you may specify the following:

Property_Info 1
Energy_Total
Volume
Pressure

For a GEMC-NVT simulation, total energy and density of all the species in box 1 and total energy, density of all the species in box 2 along with the pressure may be output using the following format:

Property_Info 1
Energy_Total
Density

Property_Info 2
Energy_Total
Density
Pressure

4.1.23 Fragment Files

Fragment_Files

Character(i,j) Integer(i,j) * One line for each fragment i in species j

In this section, information about the fragment library is specified. Character(i,j) gives the location of the fragment library of fragment i in species j; Integer(i,j) is the corresponding integer id specifying the type of the fragment. This section is automatically generated by library_setup.py. However, if there is a need to change this section, follow the example given below.

For a simulation involving two species of which the first one contains three distinct fragments and species 2 has two identical fragments, this section might look like:

```
# Fragment_Files
frag_1_1.dat 1
frag_2_1.dat 2
frag_3_1.dat 3
frag_1_2.dat 4
frag_1_2.dat 4
```

This will tell Cassandra to use the files frag_1_1.dat, frag_2_1.dat and frag_3_1.dat for the three fragments of species 1. Since species 2 has two identical fragment, Cassandra will use the same fragment library frag_1_2.dat for these fragments.

4.1.24 File Info

File_Info

Character(1)

This section is used only while generating a fragment library. Cassandra will use the filename specified in *Character(1)* to store different conformations of the fragment being simulated. Once again, this section is automatically handled by library_setup.py. However, if the user wishes to modify this part, use the following template:

```
# File_Info
frag.dat
```

This will tell Cassandra to store the fragment library in the file named frag.dat.

4.1.25 CBMC parameters

```
# CBMC_Info
kappa_ins Integer(1)
kappa_rot Integer(2)
kappa_dih Integer(3)
rcut_cbmc Real(i,4) * Number of entries equal to number of simulation boxes
```

Cassandra utilizes a configurational bias methodology based on reservoir sampling [?]. This section sets a number of parameters required for biased insertion/deletion (refer to the sections # Prob_Insertion, # Prob_Deletion and # Prob_Swap) and configurational regrowth (# Prob_Regrowth section) of molecules. For a biased insertion, a fragment is chosen at random and given a random orientation. A number of trial positions are generated for the center-of-mass of the fragment. One of the trial positions is then selected randomly based on the Boltzmann weight of the energy of the trial position. The number of trial insertion positions is given by Integer(1).

Once a trial position for the insertion is chosen, rotational bias may be applied by generating a number of trial orientations. Integer(2) specifies the number of such trial orientations. This feature will be implemented in later versions of Cassandra and any value for Integer(2), at present, is ignored. To avoid any confusion, Integer(2) is set to 0.

After the biased placement of the first fragment, additional fragments directly bonded to the first fragment are placed. Each of these fragments undergoes a number of trial orientations with respect to the fragment to which it is added. *Integer(3)* controls the number of such orientations that are generated.

For all the trials, energy of the partially grown molecule with itself and surrounding molecules is to be calculated. For this purpose, a short cutoff is used. Real(i,4) specifies the cutoff distance in Å for each of the boxes in a simulation. A short cutoff is fast, but might miss some overlaps. You can experiment with this value to optimize it for your system.

For a GEMC simulation in which 12 candidate positions are generated for biased insertion/deletion, 10 trials for biased dihedral angle selection and the cutoff for biasing energy calculation is set to 5.0 Å in box 1 and 6.5 Å in box 2, this section would look like:

```
# CBMC_Info
kappa_ins 12
kappa_rot 0
kappa_dih 10
rcut_cbmc 5.0 6.5
```

4.2. MCF FILE

4.2 MCF File

A Molecular Connectivity File (MCF) defines the information related to bonds, angles, dihedrals, impropers fragments and non bonded interactions for a given species. One MCF file is required for each species present in the system. The information contained in this file involves the force field parameters, atoms participating in each of the interactions and the functional form used in each potential contribution. The keywords are preceded by a '#' and comments follow a '!'. Similarly to the input file, the order of the keywords is not important. A complete list of the keywords is provided below.

Note that parameters for all of the following keywords must be separated by spaces only. Do not use the tab character.

Note that MCF files are generated by the script mcfgen.py automatically. The following description is provided for the users who wish to modify the MCF file or build the MCF file on their own.

4.2.1 Atom Info

Atom_Info

Integer(1)

Integer(2) Character(3)*6 Character(4)*2 Real(5) Real(6) Character(7)*20 Real(8) Real(9) Character(10)

This keyword specifies the information for non-bonded interactions. It is a required keyword in the MCF file. If not specified, the code will abort. The inputs are specified below:

- Integer(1): Total number of atoms in the species.
- Integer(2): Atom index.
- Character(3)*6: Atom type up to 6 characters. This string of characters should be unique for each interaction site in the system, i.e. do not use the same atom type for two atoms in the same (or different) species unless the (pseudo)atoms have the same atom types.
- Character(4)*2: Atom element name up to 2 characters.
- Real(5): Mass of the atom in amu. Note that for united atom models, this would be the mass of the entire pseudoatom.
- Real(6): Charge on the atom.
- Character(7): Specifies functional form for VDW interactions to be used in the simulation. This must match what is given for # VDW_Style (subsection 4.1.4) in the input file. At present only 'LJ' style is permitted.

- Real(8): The energy parameter in K.
- Real(9): Collision diameter (σ) in Å.
- Character(10): Set to 'ring' only if a given atom is part of a ring fragment. Note that a ring fragment is defined as those atoms that belong to the ring (e.g. in cyclohexane, all the six carbons) and any atom directly bonded to these ring atoms (e.g. in cyclohexane, all the hydrogens). In other words, all of the ring and exoring atoms are given the ring flag. For atoms that are not part of rings, leave this field blank.

Note that for species with a single fragment, the branch point atom is listed as the first atom.

For example, for a united atom pentane model:

```
# Atom_Info
5
1 C1_s1 C 15.0107 0.0 LJ 98.0 3.75
2 C2_s1 C 14.0107 0.0 LJ 46.0 3.95
3 C3_s1 C 14.0107 0.0 LJ 46.0 3.95
4 C4_s1 C 14.0107 0.0 LJ 46.0 3.95
5 C5_s1 C 15.0107 0.0 LJ 98.0 3.75
```

The number below the keyword # Atom_Info specifies a species with 5 interaction sites, consistent with a united atom pentane model. The first column specifies the atom ID of each of the pseudo atoms. The second and third columns provide the atom type and atom name, respectively. The fourth column represents the atomic mass of each pseudoatom. Note that the mass of C1_s1 is 15.0107 for this united atom model, as it involves a carbon and three hydrogen atoms. The same applies for all other interaction sites. The fifth column contains the partial charges placed on each of these pseudoatoms. The sixth, seventh and eighth columns contain the repulsion-dispersion functional form, the energy parameter and the collision diameter respectively. In this case, the usual Lennard-Jones functional form is used. Note that none of these atoms used the flag 'ring', as no rings are present in this molecule.

For a molecule containing rings, for example cyclohexane:

```
# Atom_Info
6
1 C1_s1 C 14.0107 0.0 LJ 52.5 3.91 ring
2 C2_s1 C 14.0107 0.0 LJ 52.5 3.91 ring
3 C3_s1 C 14.0107 0.0 LJ 52.5 3.91 ring
4 C4_s1 C 14.0107 0.0 LJ 52.5 3.91 ring
5 C5_s1 C 14.0107 0.0 LJ 52.5 3.91 ring
6 C6_s1 C 14.0107 0.0 LJ 52.5 3.91 ring
```

Note the flag 'ring' was appended as the last column for this cyclic molecule.

Finally, for the SPC/E water model:

4.2. MCF FILE

```
# Atom_Info
3
1 01_s1 0 16.00 -0.8476 LJ 78.20 3.1656
2 H2_s1 H 1.000 0.4238 LJ 0.0 0.0
3 H3_s1 H 1.000 0.4238 LJ 0.0 0.0
```

This is a molecule with a single fragment. Therefore, the branch point atom must be specified as the first atom in the list. In this case, oxygen is the branch point and thus its atom ID is 1.

4.2.2 Bond Info

```
# Bond_Info
Integer(1)
Integer(i,2) Integer(i,3) Integer(i,4) Character(i,5) Real(i,6) Real(i,7)
```

This section provides information on the number of bonds in a molecule and atoms involved in each bond along with its type. It is a required keyword in the MCF file. If not specified, the code will abort. The inputs are specified below:

- Integer(1): Total number of bonds in the species. From the next line onwards, the bonds are listed sequentially and information for each bond is included on a separate line.
- Integer(i,2): Index of the i^{th} bond.
- Integer(i,3) Integer(i,4): IDs of the atoms participating in the bond.
- Character(i,5): Type of the bond. At present only 'fixed' is permitted.
- Real(i,6): Specifies the bond length for a particular bond in Å.

Note that at present, Cassandra simulations can be carried out only for fixed bond length systems.

For example, for the water model SPC/E, the # Bond_Info section is the following:

```
# Bond_Info
2
1 1 2 fixed 1.0
2 1 3 fixed 1.0
```

In the above example, two bonds are specified whose fixed length is set to 1.0 Å.

4.2.3 Angle Info

```
# Angle Info
Integer(1)
Integer(i,2) Integer(i,3) Integer(i,4) Integer(i,5) Character(i,6) Real(i,7) Real(i,8)
```

The section lists the information on the angles in the species. It is a required keyword in the MCF file. If not specified, the code will abort.

- Integer(1): Number of angles in the species.
- Integer(i,2): Index of the i^{th} angle.
- Integer(i,3) Integer(i,4) Integer(i,5): IDs of the atoms participating in the i^{th} angle. Note that Integer(i,4) is the ID of the central atom.
- Character(i,6): Type of the angle. Currently, Cassandra supports 'fixed' and 'harmonic' (Eq. section 2.1) angles. For the 'fixed' option, Real(i,7) is the value of the angle and Real(i,8) is ignored by the code if specified. In the case of 'harmonic' potential type, Real(i,7) specifies the harmonic force constant (K/rad^2) while Real(i,8) is the nominal bond angle (in degrees).

For example, for a united atom pentane molecule with flexible angles, this section is the following:

```
# Angle_Info
3
1 1 2 3 harmonic 31250.0 114.0
2 2 3 4 harmonic 31250.0 114.0
3 3 4 5 harmonic 31250.0 114.0
```

In the above example, the three angles between the pseudoatoms found in the pentane model are specified. The three angles have an harmonic potential, whose force constant is equal and is set to 31250.0 K/rad². Finally, the equilibrium angle for these angles is 114.0° .

An example for SPC/E water model with fixed angles is provided below:

```
# Angle_Info
1
1 2 1 3 fixed 109.47
```

This model has only one angle that is set to 109.47°. Note that this angle is fixed, so there is no force constant.

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4.2.4 Dihedral Info

Dihderal_Info

Integer(1)

 $Integer(i,2) \ Integer(i,3) \ Integer(i,4) \ Integer(i,5) \ Integer(i,6) \ Character(i,7) \ Real(i,8) \ Real(i,9) \ Real(i,10) \ Real(i,11)$

This section of the MCF file lists the number of dihedral angles and associated information for a given species. It is a required keyword in the MCF file. If not specified, the code will abort.

- Integer(1): Lists the number of dihedral angles.
- Integer(i,2): Index of the i^{th} dihedral angle.
- Integer(i,3): Integer(i,6) IDs of the atoms in the i^{th} dihedral angle.
- Character(i,7): Dihedral potential type. Acceptable options are 'OPLS', 'CHARMM', 'harmonic' and 'none'. If 'OPLS' dihedral potential type is selected, then the real numbers Real(i,8) Real(i,11) are the coefficients in the Fourier series (see Eq. 2.2). The units are in kJ/mol. For the 'CHARMM' dihedral potential type, three additional parameters are specified: a_0, a_1 and δ (section 2.3). If 'harmonic' dihedral potential type is used, then two additional parameters, K_{phi} and ϕ_0 (section Eq. 2.4), are specified. For the 'none' dihedral potential type, no additional parameters are necessary.

For example, for a united atom pentane molecule using an OPLS dihedral potential type, the dihedrals are specified as follows:

```
# Dihedral_Info
2
1 1 2 3 4 OPLS 0.0 2.95188 -0.5670 6.5794
2 2 3 4 5 OPLS 0.0 2.95188 -0.5670 6.5794
```

In this model two dihedral angles are specified by atoms 1,2,3,4 and 2,3,4,5. This model uses an OPLS functional form and thus four parameters are provided after the OPLS flag.

4.2.5 Fragment Info

Fragment_Info

```
Integer(1)
```

```
Integer(i,2) \ Integer(i,3) \ Integer(i,4) \ Integer(i,5) \ \dots \ Integer(i,2+Integer(i,3))
```

This section defines the total number of fragments in a given species. It is an optional keyword. However, if the species is composed of fragments, then this section must be specified. The inputs are specified below:

- Integer(1): Total number of fragments.
- Integer(i,2): Index of the i^{th} fragment.
- Integer(i,3): Number of atoms in the i^{th} fragment.
- Integer(i,4) ... Integer(i,2+integer(i,3)): List of the atom IDs in the fragment. The first atom ID is that for the branch point atom. Atom ordering for the remaining atoms must match the order of atoms in the fragment library files.

For example, for a pentane united atom model:

```
# Fragment_Info
3
1 3 2 1 3
2 3 3 2 4
3 3 4 3 5
```

This specifies three fragments. Each of these fragments has three atoms. The first atom specified for each of the fragments is the branch point atom.

4.2.6 Fragment Connectivity

```
# Fragment_Connectivity
Integer(1)
Integer(i,2) Integer(i,3) Integer(i,4)
```

The section lists the fragment connectivity - which fragment is bonded to which other fragment. It is a required keyword if **Fragment_Info** is specified.

- Integer(1): total number of fragment connections.
- Integer(i,2): index of the i^{th} fragment connectivity.
- Integer(i,3) Integer(i,4): fragment IDs participating in the connectivty.

For example, for a pentane united atom model:

```
# Fragment_Connectivity
2
1 1 2
2 2 3
```

In this example, there are three fragments, therefore, two fragment connectivities must be specified. Note that fragment 1 is connected to fragment 2 and fragment 2 is connected to fragment 3.

Chapter 5

Utilities

5.1 Input File GUI

This script provides a graphical user interface with which the user may create input files for use in simulations. The GUI is written in Python and uses the wxWidgets for Python module, which may be found at:

http://wxpython.org/download.php

5.2 Generate an MCF file

The script mcf_gen.py is a tool that aims to ease the setup of MCF files from scratch (see section 4.2 to learn more about MCF files), as the generation of these files by hand can be error prone. In this section, a pentane MCF file will be generated to demonstrate the use of this tool. The Transferable Potentials for Phase Equilibria (TraPPE) force field will be used to represent the pentane molecular interactions. This force field involves a pairwise-additive 12-6 Lennard-Jones potential to represent the dispersion-repulsion interactions. Additionally, bond angles and dihedral angles are represented through harmonic and OPLS functional forms, respectively. Bond lengths are kept constant. The force field mathematical expression becomes

$$U = \sum_{angles} K_{\theta}(\theta - \theta_{0})^{2} + \sum_{dihedrals} \frac{1}{2} K_{1}[1 + cos(\phi)] + \frac{1}{2} K_{2}[1 - cos(2\phi)] + \frac{1}{2} K_{3}[1 + cos(3\phi)] + \frac{1}{2} K_{4}[1 - cos(4\phi)] + \sum_{i} \sum_{i>j} 4\epsilon_{ij} \left[\left(\frac{\sigma_{ij}}{r_{ij}}\right)^{12} - \left(\frac{\sigma_{ij}}{r_{ij}}\right)^{6} \right]$$

First, generate (or obtain) a PDB file or a CML file. To generate a PDB or CML file, software such as Gaussview or Avogadro can be used. Alternatively, PDB files can be downloaded from the internet (e.g. www.rcsb.org). In this example, a pentane PDB file using the program Gaussview v5.08 will be generated, as shown below.

```
        BEMARK
        1 File created by GaussView 5.0.8

        HETATM
        1 C
        0 - 4.277 0.355 0.000
        C

        HETATM
        2 C
        0 - 3.763 1.081 1.257
        C

        HETATM
        3 C
        0 - 2.223 1.078 1.259
        C

        HETATM
        5 C
        0 - 1.710 1.805 2.516
        C

        HETATM
        5 C
        0 - 0.170 1.802 2.517
        C

        CONECT
        1 2 CONECT 2 1 3 CONECT 3 2 4
        CONECT 3 2 4
        CONECT 4 3 5

        CONECT 4 3 5 CONECT 5 4
        4 5 CONECT 5 4
        C
        C
```

Append a column containing the atom types.

Avogadro v1.1.1 can also be used to generate CML files. The procedure is analogous to the one previously presented for Gaussview.

Modify the pentane united atom CML file. Note that the atom type is appended as a last column between quotation marks.

In the terminal, run the following command:

```
>python mcfgen.py pentane.pdb --ffTemplate
```

This command will create an .ff file. The first three sections of the FF file are displayed next. Do not modify these.

```
atomtypes
2
begin atom-atomtype
1 CH3
2 CH2
3 CH2
4 CH2
5 CH3
end atom-atomtype
dihedraltype OPLS
```

The force field parameters for non-bonded (not shown), bonded, angle, dihedral (not shown) and coulombic interactions (not shown) must be entered next to the corresponding keyword. For example, the angle type CH3 CH2 CH2 has an angle of 114.0. This value must be placed next to the "Angle" keyword.

bonded CH2 CH2 Length 1.54 Constant fixed

angles CH3 CH2 CH2 Angle 114.0 Constant 31250.0

For more examples of filled ff files, please refer to the examples contained in the /Scripts/MCF_Generation/directory. Using the filled .ff file, run:

> python mcfgen.py pentane.pdb

Check the file newly created pentane.mcf for any possible errors. This example can be found in the directory $/Scripts/MCF_Generation/PDB/$

5.3 Generate library of fragment configurations

The goal of the script library_setup.py is to automate the generation of fragment libraries. As a starting point, the script requires the simulation input file, and the MCF and PDB files for each of the species. To run this script, type

> library_setup.py \$PATH\$/cassandra.exe input_file.inp pdbfilespecies1.pdb pdfilespecies2.pdb

This script will create the necessary files to create the fragment libraries. It will also run Cassandra to generate these libraries, whose location will be at

/species?/frag?/frag?.inp

where '?' refers to the species number, for example, species 1, species 2 etc.

Note that the script overwrites the section of the input file where needed (i.e. # Frag_Info) with the aforementioned directory locations.

Chapter 6

Implementing the Metropolis Acceptance Criteria

All Monte Carlo moves are implemented in Cassandra to preserve detailed balance between each pair of microstates m and n

$$\Pi_{mn} \alpha_{mn} p_m = \Pi_{nm} \alpha_{nm} p_n \tag{6.1}$$

where Π_{mn} is the probability of accepting the move from microstate m to microstate n, α_{mn} is the probability of attempting the move that will form n from m, and p_m is the probability of m in the ensemble of interest.

In Cassandra, detailed balance is enforced via the Metropolis criterion

$$\Pi_{mn} = \min\left(1, \frac{\alpha_{nm}}{\alpha_{mn}} \frac{p_n}{p_m}\right) \tag{6.2}$$

The ratio in Eq. (6.2) will often involve an exponential, e.g. $e^{-\beta\Delta U}$. To preserve precision in the energy calculation, the acceptance probability is computed

$$\Pi_{mn} = \exp\left\{-\max\left[0, \ln\left(\frac{\alpha_{mn}}{\alpha_{nm}}\frac{p_m}{p_n}\right)\right]\right\}$$
(6.3)

The logarithm, defined in code as ln_pacc, is tested in the function accept_or_reject() which is defined in file accept_or_reject.f90. If ln_pacc is greater than 0 and less than a maximum numerical value, Π_{mn} is computed and compared to a random number.

Code 6.1: accept_or_reject.f90

```
47
      accept = .FALSE.
48
      IF (ln_pacc <= 0.0_DP) THEN</pre>
49
50
          accept = .TRUE.
51
52
      ELSE IF ( ln_pacc < max_kBT) THEN
53
54
55
         pacc = DEXP(-ln_pacc)
56
          IF ( rranf() <= pacc ) THEN</pre>
57
58
59
             accept = .TRUE.
60
61
          END IF
62
63
      END IF
```

6.1 Canonical Monte Carlo

In the canonical ensemble, the number of molecules N, the volume V and temperature T are all constant. The position, orientation and conformation of a semi-flexible molecule with fixed bond-lengths containing M atoms is given by a 2M+1-dimensional vector \mathbf{q} . The positions, orientations and conformations of all N molecules are denoted \mathbf{q}^N .

The probability of observing microstate m with configuration \mathbf{q}_m^N is

$$p_m = \frac{e^{-\beta U(\mathbf{q}_m^N)}}{Z(N, V, T)} d\mathbf{q}^N$$
(6.4)

where β is the inverse temperature $1/k_BT$, U is the potential energy, the differential volume $d\mathbf{q}^N$ is included to make p_m dimensionless and Z is the configurational partition function

$$Z(N, V, T) = \int e^{-\beta U(\mathbf{q}^N)} d\mathbf{q}^N.$$
 (6.5)

The integral is over all N(2M + 1) degrees of freedom. The ratio of microstate probabilities follows from Eq. (6.4)

$$\frac{p_m}{p_n} = \frac{e^{-\beta U(\mathbf{q}_m^N)} d\mathbf{q}^N / Z(N, V, T)}{e^{-\beta U(\mathbf{q}_n^N)} d\mathbf{q}^N / Z(N, V, T)}$$

$$= e^{\beta (U_n - U_m)} = e^{\beta \Delta U} \tag{6.6}$$

Table 6.1: Variable symbols and code names for translating and rotating a molecule

Symbol	Code name
β	beta(this_box)
ΔU	$delta_e$

The configurational partition function Z and differential volume $d\mathbf{q}^N$ both cancel, leaving only the ratio of Boltzmann factors.

New configurations are generated by attempting moves that translate, rotate and regrow a randomly selected molecule.

6.1.1 Translating a Molecule

A molecule is translated by moving its center of mass in each Cartesian direction by a random amount chosen from the uniform distribution on the interval $[-\delta r_{max}, \delta r_{max}]$. The maximum displacement δr_{max} must be given in the input file. The translation move is symmetric in forward and reverse directions. That is, either microstate n can be formed from microstate m and vice versa by moving one molecule within δr_{max} in each Cartesian direction, or microstate n cannot be formed at all. As a result, $\alpha_{mn} = \alpha_{nm}$.

The acceptance probability for a translation move follows from Eq. (6.6)

$$\ln\left(\frac{\alpha_{mn}}{\alpha_{nm}}\frac{p_m}{p_n}\right) = \ln\left(\frac{p_m}{p_n}\right) = \beta\Delta U \tag{6.7}$$

In Cassandra, the translation move is implemented in the subroutine Translate defined in translate.f90. The relevant lines from version 1.1 are quoted below. The variable names in the translate.f90 code are identified with the symbols from Eq. (6.7) in Table 6.1.

Code 6.2: translate.f90

```
274 ln_pacc = beta(this_box) * delta_e
275 accept = accept_or_reject(ln_pacc)
```

6.1.2 Rotating a Molecule

A linear molecule is rotated differently than a nonlinear molecule. A molecule is identified as linear if it is composed of 2 atoms or if all the angles are rigid with a bond angle of 180° . If the molecule is linear:

1. Pick three random angles: ϕ on $[-\pi, \pi]$, $\cos(\theta)$ on [-1,1], and ψ on $[-\pi, \pi]$.

2. With the origin at the molecule's center of mass, rotate by ϕ around z, rotate by θ around x', and rotate by ψ around z', as shown in Fig. ??.

Even though three angles are randomly chosen, the probability of the resulting orientation is $d\cos(\theta)d\phi/4\pi$.

If the molecule is nonlinear:

- 1. Randomly select an axis: x, y, or z.
- 2. Choose a random angular displacement $\delta\theta$ from $[-\delta\theta_{max}, \delta\theta_{max}]$. $\delta\theta_{max}$ must be given in the input file.
- 3. Rotate the molecule around a vector parallel to the selected axis and through its center of mass by $\delta\theta$.

In either case, the rotation move is symmetric, $\alpha_{mn} = \alpha_{nm}$, and the acceptance criteria is given by Eq. (6.7). The rotation move is implemented in subroutine Rotate defined in rotate.f90.

```
Code 6.3: rotate.f90
```

```
261 ln_pacc = beta(this_box) * delta_e
262 accept = accept_or_reject(ln_pacc)
```

6.1.3 Regrowing a Molecule

Internal degrees of freedom in flexible molecules are sampled by deleting one or more fragments from the molecule and replacing the deleted fragments with conformations from a reservoir of fragment conformations. If the molecule consists of only a single fragment (e.g., water, all atom methane, united atom propane, all atom cyclohexane), the entire molecule is deleted and replaced as follows:

- 1. Randomly select a molecule i with uniform probability 1/N, record its center-of-mass position and delete it.
- 2. Select a molecular conformation with Boltzmann probability $e^{-\beta U(\mathbf{q}_{int,n}^{(i)})}/Z_{int}$, where $\mathbf{q}_{int,n}^{(i)}$ are the internal bond or improper angles of molecule i in microstate n and Z_{int} is the configurational partition function over internal degrees of freedom (see Eq. (6.13)).
- 3. Pick three random angles: ϕ on $[-\pi, \pi]$, $\cos(\theta)$ on [-1,1], and ψ on $[-\pi, \pi]$. Rotate the molecule as shown in Fig. ??. The probability of the resulting orientation is $d\mathbf{q}_{rot}/Z_{rot}$, which for a nonlinear molecule is $d\cos(\theta)d\phi d\psi/8\pi^2$.
- 4. Place the molecule with the selected conformation and orientation at the same center-of-mass position as the deleted molecule.

Regrowing a monoatomic particle has no effect. Regrowing a linear molecule is the same as rotating it. The overall probability α_{mn} of regrowing a molecule with the selected orientation and conformation is

$$\alpha_{mn} = \frac{1}{N} \frac{d\mathbf{q}_{rot}}{Z_{rot}} \frac{e^{-\beta U(\mathbf{q}_n^{(i)})} d\mathbf{q}_{int}}{Z_{int}}$$
(6.8)

where $\mathbf{q}_n^{(i)}$ denotes the position, orientation and conformation of molecule i in microstate n and $U(\mathbf{q}_n^{(i)})$ is the potential energy of the isolated molecule i, i.e. the intramolecular potential energy. The reverse probability α_{nm} is identical except for the intramolecular potential energy $U(\mathbf{q}_m^{(i)})$ of molecule i in microstate m. Using Eqs. (6.6) and (6.8), the acceptance criteria for the regrowth of a single fragment molecule is

$$\ln\left(\frac{\alpha_{mn}}{\alpha_{nm}}\frac{p_m}{p_n}\right) = \beta\left[\left(U(\mathbf{q}_n^N) - U(\mathbf{q}_m^N)\right) - \left(U(\mathbf{q}_n^{(i)}) - U(\mathbf{q}_m^{(i)})\right)\right]$$

$$= \beta\Delta U - \beta\Delta U_{int}^{(i)} = \beta\Delta U_{inter}^{(i)}$$
(6.9)

Only the change in the intermolecular potential energy between molecule i and the other N-1 molecules contributes to the acceptance criteria. The code that implements Eq. (6.9) is shown in Code 6.7 in Section 6.3.3.

If the molecule consists of more than one fragment (e.g., all atom ethane, all atom toluene, united atom butane), a bond is cut and the severed fragments are regrown using Configurational Bias Monte Carlo (CBMC). See Section 6.3.3 for more details.

6.1.4 Canonical Partition Function

In Sections 6.1.1-6.1.2, the microstate probability is normalized by the configuration partition function Z because the only relevant degrees of freedom are configurational. In other ensembles, the full partition function Q appears, integrated over both configuration space \mathbf{q}^N and momenta space \mathbf{p}_q^N

$$Q(N, V, T) = \frac{1}{h^{N(2M+1)}N!} \int e^{-\beta H(\mathbf{p}_q^N, \mathbf{q}^N)} d\mathbf{p}_q^N d\mathbf{q}^N$$
(6.10)

where the 2M+1 momenta \mathbf{p}_q are conjugate to the generalized coordinates \mathbf{q} . The momenta and configuration integrals are separable, and the single molecule momenta integrals are all identical.

$$Q(N, V, T) = \frac{1}{N!} \left[\int e^{-\beta U(\mathbf{q}^N)} d\mathbf{q}^N \right] \left[\frac{1}{h^{2M+1}} \int e^{-\beta K(\mathbf{p}_q)} d\mathbf{p}_q \right]^N$$
$$= \frac{1}{N!} Z(N, V, T) \left[\frac{Q(1, V, T)}{Z(1, V, T)} \right]^N$$
(6.11)

where Q(1, V, T) is the partition function of a single molecule in a box. The center of mass integrals for a single molecule are separable from the integrals over rotational and internal degrees of freedom:

$$Q(1, V, T) = Q_{com}Q_{rot+int} = V\Lambda^{-3}Q_{rot+int}$$

$$(6.12)$$

where Λ is the de Broglie wavelength of the molecule and the rotational and internal momenta integratals in $Q_{rot+int}$ are not separable since the moments of inertia will depend on the conformation adopted by the molecule. The configurational partition function is further separable into center of mass (translational), orientational and internal degrees of freedom:

$$Z(1, V, T) = V Z_{rot} Z_{int} \tag{6.13}$$

where the volume V is the translational partition function and Z_{rot} equals 4π for a linear molecule and $8\pi^2$ for a nonlinear molecule.

6.2 Isothermal-Isobaric Monte Carlo

In the isothermal-isobaric ensemble, the number of particles N, the pressure P and temperature T are all constant while the volume V and energy E fluctuate. The partition function is

$$\Delta(N, P, T) = \int e^{-\beta PV} Q(N, V, T) dV$$
(6.14)

Note that Q is dimensionless and Δ has dimensions of volume. The probability of the system having volume V is

$$p(V) = \frac{Q(N, V, T)e^{-\beta PV}}{\Delta(N, P, T)}dV$$
(6.15)

The probability of observing microstate m with configuration \mathbf{q}_m^N and volume V_m is

$$p_{m} = \frac{e^{-\beta U(\mathbf{q}_{m}^{N})} d\mathbf{q}_{m}^{N}}{Z(N, V_{m}, T)} \frac{Q(N, V_{m}, T) e^{-\beta P V_{m}} dV}{\Delta(N, P, T)}$$

$$= \frac{e^{-\beta U_{m} - \beta P V_{m}}}{\Delta(N, P, T)} \left(\frac{Q(1, V_{m}, T)}{Z(1, V_{m}, T)} d\mathbf{q}_{m}\right)^{N} dV$$

$$(6.16)$$

where the differential volume element $d\mathbf{q}_m^N$ has subscript m because it depends on the volume V_m . The ratio of microstate probabilities is

$$\frac{p_m}{p_n} = e^{\beta(U_n - U_m) + \beta P(V_n - V_m)} \left(\frac{d\mathbf{q}_m}{d\mathbf{q}_n}\right)^N = e^{\beta \Delta U + \beta P \Delta V} \left(\frac{d\mathbf{q}_m}{d\mathbf{q}_n}\right)^N \tag{6.17}$$

6.2.1 Scaling the Volume

In Cassandra, new volumes are sampled as follows:

- 1. Pick a random volume ΔV with uniform probability from the interval $[-\delta V_{max}, \, \delta V_{max}]$. The trial volume is $V + \Delta V$.
- 2. Scale the box lengths uniformly.
- 3. Scale the center of mass of each molecule uniformly.

The probability of selecting ΔV is the same as selecting $-\Delta V$ which makes scaling the volume symmetric, $\alpha_{mn} = \alpha_{nm}$. Scaling the configurations changes the differential element $d\mathbf{q}_m^N$ surrounding configuration \mathbf{q}_m^N . Only the molecular centers of mass change, so we can separate $d\mathbf{q}$ into 3 center of mass coordinates $d\mathbf{r}_{com}$ and 2M-2 orientational and internal coordinates $d\mathbf{q}_{rot+int}$. The scaled center of mass positions are held constant, making $d\mathbf{r}_{com} = V d\mathbf{s}_{com}$. The acceptance probability for a volume scaling move is

$$\ln\left(\frac{\alpha_{mn}}{\alpha_{nm}}\frac{p_m}{p_n}\right) = \ln\left(\frac{p_m}{p_n}\right) = \beta\Delta U + \beta P\Delta V + N\ln\left(\frac{V_m}{V_n}\right)$$
(6.18)

The volume scaling move is implemented in subroutine Volume_Change defined in volume_change.f90.

Code 6.4: volume_change.f90

Symbol	Code name
β	beta(this_box)
ΔU	$ m delta_e$
P	$pressure(this_box)$
ΔV	$ m delta_volume$
N	$total_molecules$
V_n	box_list(this_box)%volume
V_m	box_list_old%volume

Table 6.2: Variable symbols and code names for volume scaling move.

6.3 Grand Canonical Monte Carlo

In the grand canonical ensemble, the chemical potential μ , the volume V and temperature T are held constant while the number of molecules N and energy E fluctuate. The partition function is

$$\Xi(\mu, V, T) = \sum_{N=0}^{\infty} Q(N, V, T) e^{\beta \mu N}$$
(6.19)

The probability of the system having N molecules is

$$p(N) = \frac{Q(N, V, T)e^{\beta\mu N}}{\Xi(\mu, V, T)}$$

$$(6.20)$$

The probability of observing microstate m with N_m molecules and configuration $\mathbf{q}_m^{N_m}$ is

$$p_{m} = \frac{e^{-\beta U(\mathbf{q}_{m}^{N_{m}})} d\mathbf{q}^{N_{m}}}{Z(N_{m}, V, T)} \frac{Q(N_{m}, V, T) e^{\beta \mu N_{m}}}{\Xi(\mu, V, T)}$$

$$= \frac{e^{-\beta U_{m} + \beta \mu N_{m}}}{\Xi(\mu, V, T)} \left[\frac{Q(1, V, T)}{Z(1, V, T)} d\mathbf{q} \right]^{N_{m}}$$
(6.21)

Note that Eq. (6.21) does not contain the factorial $N_m!$ that accounts for indistinguishable particles. In a simulation, particles *are* distinguishable: they are numbered and specific particles are picked for MC moves. The ratio of microstate probabilities is

$$\frac{p_m}{p_n} = e^{\beta \Delta U - \beta \mu \Delta N} \left[\frac{Q(1, V, T)}{Z(1, V, T)} d\mathbf{q} \right]^{-\Delta N}$$
(6.22)

Alternatively, Eq. (6.22) can be recast to use the fugacity f instead of the chemical potential μ . The relationship between μ and f is

$$\mu = -k_B T \ln \left(\frac{Q(1, V, T)}{N} \right) = -k_B T \ln \left(\frac{Q(1, V, T)}{\beta f V} \right)$$
(6.23)

Inserting Eq. (6.23) into Eq. (6.22) yields

$$\frac{p_m}{p_n} = e^{\beta \Delta U} \left[\frac{\beta f V}{Z(1, V, T)} d\mathbf{q} \right]^{-\Delta N}$$
(6.24)

Fluctuations in the number of molecules are achieved by inserting and deleting molecules. A successful insertion increases the number of molecules from N to N+1, i.e. $\Delta N=1$. A successful deletion decreases the number of molecules from N to N-1, i.e. $\Delta N=-1$.

Random insertions and deletions (see Section 6.6) in the liquid phase typically have very high ΔU due to core overlap and dangling bonds, respectively, making the probability of acceptance very low. Instead, insertions in Cassandra are attempted using Configurational Bias Monte Carlo.

6.3.1 Inserting a Molecule with Configurational Bias Monte Carlo

In Configurational Bias Monte Carlo (CBMC), the molecular conformation of the inserted molecule is molded to the insertion cavity. First, the molecule is parsed into fragments such that each fragment is composed of (a) a central atom and the atoms directly bonded to it (see Fig. ??), or (b) a ring of atoms and all the atoms directly bonded to them. Then, a position, orientation and molecular conformation of the molecule to be inserted are selected via the following steps:

- 1. Select the order in which each fragment of the (N+1)th molecule will be placed. The probability of the resulting sequence is p_{seq} . (See example in Table. 6.3.)
 - (a) The first fragment i is chosen with uniform probability $1/N_{frag}$.
 - (b) Subsequent fragments must be connected to a previously chosen fragment and are chosen with the uniform probability $1/N_{cnxn}$, where the number of connections $N_{cnxn} = \sum_{ij} \delta_{ij} h_i (1 h_j)$ is summed over all fragments i and j. h_i is 1 if fragment i has been previously chosen and 0 otherwise. δ_{ij} is 1 if fragments i and j are connected and 0 otherwise.
- 2. Select a conformation for fragment i with Boltzmann probability $e^{-\beta U(\mathbf{q}_{frag_i})}d\mathbf{q}_{frag_i}/Z_{frag_i}$, where \mathbf{q}_{frag_i} are the internal degrees of freedom (angles and/or impropers) associated with fragment i.

Table 6.3: Possible sequences and probabilities for inserting the fragments of the all-atom model of propane shown in Fig. ??.

Sequence	p_{seq}
1 2 3	1/3
2 1 3	1/6
2 3 1	1/6
3 2 1	1/3

- 3. Select an orientation with uniform probability $d\mathbf{q}_{rot}/Z_{rot}$.
- 4. Select a coordinate for the center of mass (COM) of fragment i:
 - (a) Select κ_{ins} trial coordinates \mathbf{r}_k , each with uniform probability $d\mathbf{r}/V$. Since one of the trial coordinates will be selected later, the individual probabilities are additive. The probability of the collection of trial coordinates is $\kappa_{ins}d\mathbf{r}/V$.
 - (b) Compute the change in potential energy ΔU_k^{ins} of inserting fragment i at each position \mathbf{r}_k into configuration \mathbf{q}_m^N .
 - (c) Select one of the trial coordinates with probability $e^{-\beta\Delta U_k^{ins}}/\sum_k e^{-\beta\Delta U_k^{ins}}$.
- 5. For each additional fragment j:
 - (a) Select a fragment conformation with Boltzmann probability $e^{-\beta U(\mathbf{q}_{frag_j})}d\mathbf{q}_{frag_j}/Z_{frag_j}$
 - (b) Select the first of κ_{dih} trial dihedrals ϕ_0 with uniform probability from the interval $[0, \frac{2\pi}{\kappa_{dih}})$. Additional trial dihedrals are equally spaced around the unit circle, $\phi_k = \phi_{k-1} + 2\pi/\kappa_{dih}$. The probability of selecting ϕ_0 is $\kappa_{dih}d\phi/2\pi$.
 - (c) Compute the change in potential energy ΔU_k^{dih} of attaching fragment j to the growing molecule with each dihedral ϕ_k .
 - (d) Select one of the trial dihedrals with probability $e^{-\beta\Delta U_k^{dih}}/\sum_k e^{-\beta\Delta U_k^{dih}}$.

The overall probability α_{mn} of attempting the insertion with the selected position, orientation and conformation is

$$\alpha_{mn} = p_{seq} \frac{d\mathbf{q}_{rot}}{Z_{rot}} \frac{\kappa_{ins} d\mathbf{r}}{V} \frac{e^{-\beta \Delta U_k^{ins}}}{\sum_k e^{-\beta \Delta U_k^{ins}}} \times \begin{bmatrix} \prod_{i=1}^{N_{frag}} \frac{e^{-\beta U(\mathbf{q}_{frag_i})} d\mathbf{q}_{frag_i}}{Z_{frag_i}} \end{bmatrix} \begin{bmatrix} \prod_{j=1}^{N_{frag}-1} \frac{\kappa_{dih} d\phi}{2\pi} \frac{e^{-\beta \Delta U_k^{dih}}}{\sum_k e^{-\beta \Delta U_k^{dih}}} \end{bmatrix}$$

$$= p_{seq} p_{bias} \frac{e^{-\beta U(\mathbf{q}_{frag})} d\mathbf{q}}{V Z_{rot} Z_{frag} \Omega_{dih}}$$

$$(6.25)$$

where $Z_{frag} = \prod_i Z_{frag_i}$ is the configurational partition function over degrees of freedom internal to each fragment, $U(\mathbf{q}_{frag}) = \sum_i U(\mathbf{q}_{frag_i})$ is the summed potential energy of each of the (disconnected) fragments, $\Omega_{dih} = (2\pi)^{N_{frag}-1}$ and p_{bias} is

$$p_{bias} = \frac{\kappa_{ins} \ e^{-\beta \Delta U_k^{ins}}}{\sum_k e^{-\beta \Delta U_k^{ins}}} \left[\prod_{j=1}^{N_{frag}-1} \frac{\kappa_{dih} \ e^{-\beta \Delta U_k^{dih}}}{\sum_k e^{-\beta \Delta U_k^{dih}}} \right]$$
(6.27)

Note that the term $VZ_{rot}Z_{frag}\Omega_{dih}$ in the denominator of Eq. (6.26) differs from $Z(1, V, T) = VZ_{rot}Z_{int}$.

In the reverse move, 1 of the N+1 particles is randomly selected for deletion. The probability α_{nm} of picking the molecule we just inserted is

$$\alpha_{nm} = \frac{1}{N+1} \tag{6.28}$$

Combining Eqs. (6.26) and (6.28) with Eq. (6.22) or Eq. (6.24) gives the acceptance probability for a CBMC insertion move

$$\ln\left(\frac{\alpha_{mn}}{\alpha_{nm}}\frac{p_m}{p_n}\right) = \beta\left[\Delta U - U(\mathbf{q}_{frag,n}^{(N+1)})\right] - \beta\mu' + \ln\left(\frac{(N+1)\Lambda^3}{V}\right) + \ln\left(p_{seq}p_{bias}\right)$$
(6.29)

$$= \beta \left[\Delta U - U(\mathbf{q}_{frag,n}^{(N+1)}) \right] + \ln \left(\frac{N+1}{\beta f'V} \right) + \ln \left(p_{seq} p_{bias} \right)$$
(6.30)

where μ' and f' are, respectively, a shifted chemical potential and a skewed fugacity,

$$\mu' = \mu + k_B T \ln \left(Q_{rot+int} \frac{Z_{frag} \Omega_{dih}}{Z_{int}} \right)$$

$$f' = f \frac{Z_{frag} \Omega_{dih}}{Z_{int}}$$

$$(6.31)$$

$$f' = f \frac{Z_{frag} \Omega_{dih}}{Z_{int}} \tag{6.32}$$

All of the terms in Eqs. (6.31) and (6.32) are intensive. GCMC simulations using Eqs. (6.29) and (6.30) will converge to the same average density regardless of the simulation volume V. However, the values of μ' or f' that correspond to the converged density will not match tabulated values of μ or f computed from experimental data.

Note that the term Z^{IG}/Ω from Macedonia et al. would be equivalent to $Z_{int}/\Omega_{frag}\Omega_{dih}$ in the nomenclature used here. The configurational partition function of the disconnected fragments integrates over a Boltzmann factor, $Z_{frag} = \int e^{-\beta U(\mathbf{q}_{frag})} d\mathbf{q}_{frag}$, whereas the term $\Omega_{frag} = \int d\mathbf{q}_{frag}$ does not.

	, <u> </u>
\mathbf{Symbol}	Code name
β	beta(this_box)
ΔU	$delta_e$
$U(\mathbf{q}_{frag})$	E_angle + nrg_ring_frag_tot
p_{seq}	P_seq
p_{bias}	P_bias
μ'	species_list(is)%chem_potential
N	$nmols(is,this_box)$
V	box_list(this_box)%volume
Λ	species_list(is)%de_broglie(this_box)
f'	species_list(is)%fugacity

Table 6.4: Variable symbols and code names for inserting a molecule

In Cassandra, the insertion move is implemented in the subroutine Insertion in insertion.f90. The relevant lines from version 1.1 are quoted below. The variable names in the insertion.f90 code are identified with symbols in Table 6.4.

Code 6.5: insertion.f90

```
441
      ln_pacc = beta(this_box) * (delta_e - E_angle - nrg_ring_frag_tot)
      ln_pacc = ln_pacc + DLOG(P_seq * P_bias) &
447
                         + DLOG(REAL(nmols(is,this_box)+1,DP)) &
448
449
                         - DLOG(box_list(this_box)%volume)
450
451
      IF(lchempot) THEN
452
          ! chemical potential is input
         ln_pacc = ln_pacc - species_list(is)%chem_potential * beta(this_box) &
453
454
                    + 3.0_DP*DLOG(species_list(is)%de_broglie(this_box))
      ELSE
455
456
          ! fugacity is input
         ln_pacc = ln_pacc - DLOG(species_list(is)%fugacity) &
457
458
                            - DLOG(beta(this_box)) &
459
      END IF
460
461
      accept = accept_or_reject(ln_pacc)
```

6.3.2 Deleting a Molecule that was Inserted via Configurational Bias Monte Carlo

The probability α_{mn} of choosing a molecule to delete is

$$\alpha_{mn} = \frac{1}{N} \tag{6.33}$$

The probability of the reverse move α_{nm} requires knowledge of the sequence and biasing probabilities p_{seq} and p_{bias} that would have been used to place the molecule if it was being inserted. p_{seq} and p_{bias} can be calculated using the following procedure:

- 1. Select the fragment order using the same procedure for inserting a molecule. The probability of the resulting sequence is p_{seq} .
- 2. The first fragment in the sequence is fragment j. Calculate the intramolecular potential energy of fragment j's current conformation, $U(\mathbf{q}_{frag_j})$. The probability of this conformation is Boltzmann $e^{-\beta U(\mathbf{q}_{frag_j})}d\mathbf{q}_{frag_j}/Z_{frag_j}$.
- 3. The probability of the fragment's current orientation is $d\mathbf{q}_{rot}/Z_{rot}$.
- 4. Calculate the weight of the fragment's current center of mass (COM) coordinates:
 - (a) Compute the interaction potential energy ΔU^{ins} between fragment j and the other N-1 molecules.
 - (b) Select $\kappa_{ins} 1$ trial coordinates \mathbf{r}_k , each with uniform probability $d\mathbf{r}/V$.
 - (c) Calculate the weight of the fragment's current COM amongst the trial coordinates, $e^{-\beta\Delta U^{ins}}/\sum_k e^{-\beta\Delta U^{ins}_k}$.
- 5. For each additional fragment j:
 - (a) Calculate the intramolecular potential energy of fragment j's current conformation, $U(\mathbf{q}_{frag_j})$. The weight of this conformation in the Boltzmann distribution is $e^{-\beta U(\mathbf{q}_{frag_j})} d\mathbf{q}_{frag_i}/Z_{frag_i}$.
 - (b) Calculate the interaction potential energy ΔU^{dih} between fragment j, on the one hand, and fragments i through j-1 and the other N-1 molecules.
 - (c) Calculate the current dihedral ϕ_0 of fragment j. Compute the interaction potential energy ΔU_k^{dih} at $\kappa_{dih} 1$ trial dihedrals $\phi_k = \phi_{k-1} + 2\pi/\kappa_{dih}$.
 - (d) Compute the weight of ϕ_0 amongst the trial dihedrals, $e^{-\beta\Delta U^{dih}}/\sum_k e^{-\beta\Delta U^{dih}_k}$.

The overall probability α_{nm} is

$$\alpha_{nm} = p_{seq} \ p_{bias} \ \frac{e^{-\beta U(\mathbf{q}_{frag})} d\mathbf{q}}{V Z_{rot} Z_{frag} \Omega_{dih}}. \tag{6.34}$$

The acceptance criteria for deleting a molecule inserted via CBMC is

$$\ln\left(\frac{\alpha_{mn}}{\alpha_{nm}}\frac{p_m}{p_n}\right) = \beta\left[\Delta U + U(\mathbf{q}_{frag,m}^{(i)})\right] + \beta\mu' + \ln\left(\frac{V}{N\Lambda^3}\right) - \ln\left(p_{seq}p_{bias}\right)$$
(6.35)

$$= \beta \left[\Delta U + U(\mathbf{q}_{frag,m}^{(i)}) \right] + \ln \left(\frac{\beta f'V}{N} \right) - \ln \left(p_{seq} p_{bias} \right)$$
 (6.36)

In Cassandra, the deletion move is implemented in the subroutine Deletion in deletion.f90. The relevant lines are quoted below. The variable names in deletion.f90 code are identified with symbols in Table 6.5.

Code 6.6: deletion.f90

```
ln_pacc = beta(this_box) * (delta_e + E_angle + nrg_ring_frag_tot)
334
340
      ln_pacc = ln_pacc - DLOG(P_seq * P_bias) &
341
                          - DLOG(REAL(nmols(is,this_box),DP)) &
                         + DLOG(box_list(this_box)%volume)
342
343
344
      IF(lchempot) THEN
345
         ! chemical potential is input
         ln_pacc = ln_pacc + beta(this_box) * species_list(is)%chem_potential &
346
347
                  - 3.0_DP*DLOG(species_list(is)%de_broglie(this_box))
348
      ELSE
349
         ! fugacity is input
350
         ln_pacc = ln_pacc + DLOG(species_list(is)%fugacity) &
351
                             + DLOG(beta(this_box)) &
352
      END IF
353
354
      accept = accept_or_reject(ln_pacc)
```

6.3.3 Regrowing a Molecule with Configurational Bias Monte Carlo

Regrowing a molecule that has more than one fragment is a combination deletion and insertion move. Starting from microstate m:

- 1. Randomly select a molecule with uniform probability 1/N.
- 2. Randomly select a bond to cut on the selected molecule with uniform probability $1/N_{bonds}$.
- 3. Delete the fragments on one side of the bond or the other with equal probability. The number of deleted fragments is N_{del} .

Symbol	Code name
β	beta(this_box)
ΔU	$delta_e$
$U(\mathbf{q}_{frag})$	E_angle + nrg_ring_frag_tot
p_{seq}	P_seq
p_{bias}	P_bias
μ'	species_list(is)%chem_potential
N	$nmols(is,this_box)$
V	box_list(this_box)%volume
Λ	species_list(is)%de_broglie(this_box)
f'	species_list(is)%fugacity

Table 6.5: Variable symbols and code names for deleting a molecule

4. Reinsert the deleted fragments using the CBMC procedures for selecting the order of inserting the fragments, choosing a fragment conformation, and a connecting dihedral value (see Section 6.3.1).

The overall probability α_{mn} of attempting to regrow the molecule with the selected conformation is

$$\alpha_{mn} = \frac{p_{seq}}{NN_{bonds}} \left[\prod_{j=1}^{N_{del}} \frac{e^{-\beta U(\mathbf{q}_{frag_{j}}^{(i)})} d\mathbf{q}_{frag_{j}}}{Z_{frag_{j}}} \right] \left[\prod_{j=1}^{N_{del}} \frac{\kappa_{dih} d\phi}{2\pi} \frac{e^{-\beta \Delta U_{k}^{dih}}}{\sum_{k} e^{-\beta \Delta U_{k}^{dih}}} \right]$$

$$= \frac{p_{seq}}{NN_{bonds}} \frac{e^{-\beta U(\mathbf{q}_{del,n}^{(i)})} d\mathbf{q}}{Z_{del} \Omega_{del}} p_{forward}$$
(6.37)

where $Z_{del} = \prod_i Z_{frag_j}$ is the configurational partition function over degrees of freedom internal to the deleted fragments, $U(\mathbf{q}_{del,n}^{(i)}) = \sum_j U(\mathbf{q}_{frag_j})$ is the summed potential energy of each deleted fragment with the conformations in microstate n, $\Omega_{del} = (2\pi)^{N_{del}}$ and $p_{forward}$ is the biasing probability

$$p_{forward} = \prod_{j=1}^{N_{del}} \frac{\kappa_{dih} e^{-\beta \Delta U_k^{dih}}}{\sum_k e^{-\beta \Delta U_k^{dih}}}$$
(6.38)

The reverse move is identical except for the potential energy of the deleted fragments $U(\mathbf{q}_{del,m}^{(i)})$ in microstate m and the biasing probability $p_{reverse}$ which will depend on the values of the connecting dihedrals. Using Eqs. (6.6) and (6.37), the acceptance criteria is:

$$\ln\left(\frac{\alpha_{mn}}{\alpha_{nm}}\frac{p_m}{p_n}\right) = \beta\left[\left(U(\mathbf{q}_n^N) - U(\mathbf{q}_{del,n}^{(i)})\right) - \left(U(\mathbf{q}_m^N) - U(\mathbf{q}_{del,m}^{(i)})\right)\right] + \ln\left(\frac{p_{forward}}{p_{reverse}}\right)$$
(6.39)

v	0 0
Symbol	Code name
β	beta(this_box)
$U(\mathbf{q}_n^N) - U(\mathbf{q}_{del,n}^{(i)})$ $U(\mathbf{q}_m^N) - U(\mathbf{q}_{del,m}^{(i)})$	delta_e_n - nrg_ring_frag_forward
$U(\mathbf{q}_m^N) - U(\mathbf{q}_{del,m}^{(i)})$	delta_e_o - nrg_ring_frag_reverse
$p_{forward}$	P_forward
$p_{reverse}$	P_reverse

Table 6.6: Variable symbols and code names for regrowing a molecule

Eq. (6.39) is implemented in subroutine cut_N_grow() in file cutNgrow.f90.

```
Code 6.7: cutNgrow.f90
```

6.4 Gibbs Ensemble Monte Carlo

The Gibbs Ensemble Monte Carlo method is a standard technique for studying phase equilibria of pure fluids and mixtures. It is oftentimes the method of choice to study vapor-liquid equilibria due to its intuitive physical basis. In Cassandra, the NVT and NPT versions of the Gibbs Ensemble (GEMC-NVT and GEMC-NPT) are implemented. The GEMC-NVT method is suitable for simulating vapor liquid equilibria of pure systems, since pure substances require the specification of only one intensive variable (temperature) to completely specify a state of two phases. By contrast, mixtures require the specification of an additional degree of freedom (pressure). Thus, in the GEMC-NPT method, the pressure is specified in addition to temperature.

In section 6.4.1, the acceptance rules for the NVT version will be derived and references to their implementation in Cassandra will be presented. For information about the implementation of the NPT version, please refer to section 6.4.2

6.4.1 Gibbs Ensemble-NVT

In the GEMC-NVT method, the system is comprised by two boxes, one representing a vapor phase (denoted by superscript v) and the other representing a liquid phase (denoted by superscript l). These boxes are in thermal contact with a heat bath that maintains them at constant temperature, thus ensuring thermal

equilibrium. To achieve phase equilibrium, the boxes are allowed to exchange volume and particles under the constraint of constant total volume $(V^t = V^l + V^v)$ and constant number of particles $(N^t = N^l + N^v)$. The partition function of this system is

$$Q(N^{t}, V^{t}, T) = \frac{1}{\Lambda^{3N^{t}} N^{t}!} \sum_{N^{l}}^{N^{t}} \frac{N^{t}!}{N^{l}! N^{v}!} \int_{0}^{V^{t}} dV^{l} (V^{l})^{N^{l}} (V^{v})^{N^{v}} \int d\mathbf{s}^{N^{l}} e^{-\beta U(\mathbf{s}^{N^{l}})} \int d\mathbf{s}^{N^{v}} e^{-\beta U(\mathbf{s}^{N^{v}})}$$
(6.40)

Where Λ is the de Broglie wavelength, \mathbf{s}^{N^l} and \mathbf{s}^{N^v} are the scaled coordinates of the atoms in the two phases, $U\left(\mathbf{s}^{N^l}\right)$ and $U\left(\mathbf{s}^{N^v}\right)$ are the potential energies of the liquid and the vapor box, respectively. The microstate probability distribution is

$$p_m \propto (V^l)^{N^l} (V^v)^{N^v} e^{-\beta U \left(\mathbf{s}^{N^l}\right) - \beta U \left(\mathbf{s}^{N^v}\right)}$$

$$\tag{6.41}$$

Note that the particle number factorials are not included in equation 6.41, as particles are distinguishable in a simulation (see also equation 6.21).

In the GEMC-NVT technique, thermal equilibrium is attained by performing translation, rotation and regrowth moves. The acceptance rules for these moves can be extracted from the probability distribution 6.41, and their functional form is exactly the same as that presented in sections 6.1.1, 6.1.2, 6.1.3 and 6.3.3, namely

$$\ln\left(\frac{\alpha_{mn}}{\alpha_{nm}}\frac{p_m}{p_n}\right) = -\beta\Delta U \tag{6.42}$$

Exchange of volume between the two boxes is carried out to achieve pressure equilibrium. Assuming volume is being transfered from box A to box B, the acceptance rule is also derived from equation 6.41 and yields

$$\ln\left(\frac{\alpha_{mn}}{\alpha_{nm}}\frac{p_m}{p_n}\right) = -\beta\Delta U^A - \beta\Delta U^B + N^A \ln\left(\frac{V_m^A}{V_n^A}\right) + N^B \ln\left(\frac{V_m^B}{V_n^B}\right)$$
(6.43)

This acceptance rule is implemented in the file gemc_nvt_volume.f90 as follows

Symbol	Code name
β^A	beta(box1)
eta^B	beta(box2)
ΔU^A	$ m delta_e_1$
ΔU^B	$delta_e_2$
N^A	$tot_mol_box_1$
N^B	$tot_mol_box_2$
V_m^A	box_list(box1)%volume
V_m^B	box_list(box2)%volume
V_n^A	box_list_old_1%volume
V^B	box_list_old_2%volume

Table 6.7: Variable symbols and code names for the volume scaling move in the GEMC-NVT method.

Code 6.8: gemc_nvt_volume.f90

Finally, exchange of particles is conducted between the two boxes to attain equality of chemical potentials. If the particle is transferred from box A into box B, the acceptance rule for this move is

$$\ln\left(\frac{\alpha_{mn}}{\alpha_{nm}}\frac{p_m}{p_n}\right) = \ln\left(\frac{\alpha_{mn}}{\alpha_{nm}}\right)\ln\left(\frac{V^B}{V^A}\right) - \beta\Delta U^A - \beta\Delta U^B$$
(6.44)

Note that in contrast to the volume move, the particle swap is not a symmetric move, as it might involve configurational biased regrowths. For the case of transfering a molecule from box A into box B, the forward probability α_{nm} is calculated as follows

- 1. Pick box A
- 2. Pick a molecule from box A out of the \mathcal{N}_n^A possible molecules
- 3. Insert molecule in box B using protocol presented in section 6.3.1

The attempt probability of generating configuration m is

$$\alpha_{nm} = \frac{1}{2} \frac{1}{N_n^A} p_{seq_{nm}} \frac{d\mathbf{q}_{rot}}{Z_{rot}} \frac{\kappa_{ins} V^B d\mathbf{s}}{V^B} \left[\frac{e^{-\beta \Delta U_k^{ins}}}{\sum_k e^{-\beta \Delta U_k^{ins}}} \right]^B \times \left[\prod_{i=1}^{N_{frag}} \frac{e^{-\beta U(\mathbf{q}_{frag_i})} d\mathbf{q}_{frag_i}}{Z_{frag_i}} \right]_{nm} \left[\prod_{j=1}^{N_{frag}-1} \frac{\kappa_{dih} d\phi}{2\pi} \frac{e^{-\beta \Delta U_k^{dih}}}{\sum_k e^{-\beta \Delta U_k^{dih}}} \right]_{nm}$$

$$= \frac{1}{2} \frac{1}{N_n^A} p_{seq_{nm}} p_{bias_{nm}} \frac{e^{-\beta U(\mathbf{q}_{frag_{nm}})} d\mathbf{q}_{rot} d\mathbf{s} \prod_{i=1}^{N_{frag}} d\mathbf{q}_{frag_{i,nm}} \prod_{j=1}^{N_{frag}-1} d\phi_{nm}}{Z_{rot} Z_{frag} \Omega_{dih}}$$

$$(6.45)$$

Once the new configuration m has been generated, the reverse probability α_{mn} must be calculated. This represents the probability of generating the old state n, pretending that the current state is m. The procedure is similar as before

- 1. Pick box B
- 2. Pick the swapped molecule from box B of the $N_n^B + 1$ possible molecules
- 3. Insert molecule back in box A using protocol presented in section 6.3.1

The attempt probability of generating the old configuration n pretending we are in the new configuration m is

$$\alpha_{mn} = \frac{1}{2} \frac{1}{N_n^B + 1} p_{seq_{mn}} p_{bias_{mn}} \frac{e^{-\beta U(\mathbf{q}_{frag_{mn}})} d\mathbf{q}_{rot} d\mathbf{s} \prod_{i=1}^{N_{frag}} d\mathbf{q}_{frag_{i,mn}} \prod_{j=1}^{N_{frag} - 1} d\phi_{mn}}{Z_{rot} Z_{frag} \Omega_{dih}}$$
(6.47)

The acceptance rule now becomes

$$\ln\left(\frac{\alpha_{mn}}{\alpha_{nm}}\frac{p_m}{p_n}\right) = \ln\left(\frac{p_{bias_{mn}}}{p_{bias_{nm}}}\frac{N_n^A}{N_n^B + 1}\frac{V^B}{V^A}\right) + \beta U(\mathbf{q}_{frag_{nm}}) - \beta U(\mathbf{q}_{frag_{mn}}) - \beta \Delta U^A - \beta \Delta U^B$$
 (6.48)

Note that the terms p_{seq} cancelled out, as the same fragment regrowth sequence is used in the forward and reverse moves. The implementation of the particle swap move is found in the file gemc_particle_transfer.f90 as follows

Symbol	Code name
β^A	beta(box_out)
β^B	beta(box_in)
ΔU^A	-delta_e_out
ΔU^B	delta_e_in
$U(\mathbf{q}_{frag_{nm}})$	e_angle_in + nrg_ring_frag_in
$U(\mathbf{q}_{frag_{mn}})$	e_angle_out + nrg_ring_frag_out
V_m^A	box_list(box_out)%volume
V_m^B	$box_list(box_in)\%volume$
$p_{bias_{nm}}$	P_forward
p_{bias}	P_reverse

Table 6.8: Variable symbols and code names for the particle transfer move in the GEMC-NVT method.

Code 6.9: gemc_particle_transfer.f90

```
505
    delta_e_in_pacc = delta_e_in
506
    delta_e_out_pacc = delta_e_out
518
    delta_e_in_pacc = delta_e_in_pacc - e_angle_in - nrg_ring_frag_in
519
    delta_e_out_pacc = delta_e_out_pacc - e_angle_out - nrg_ring_frag_out
526
    ln_pacc = beta(box_in)*delta_e_in_pacc - beta(box_out)*delta_e_out_pacc
527
528
    ln_pacc = ln_pacc - DLOG(box_list(box_in)%volume) &
                         + DLOG(box_list(box_out)%volume) &
529
530
                         - DLOG(REAL(nmols(this_species,box_out),DP)) &
531
                         + DLOG(REAL(nmols(this_species,box_in) + 1, DP))
532
    ln_pacc = ln_pacc + DLOG(P_forward / P_reverse)
533
    accept = accept_or_reject(ln_pacc)
534
```

6.4.2 Gibbs Ensemble-NPT

In the GEMC-NPT method, the system is comprised of two boxes. These boxes are allowed to exchange particles to attain equality of chemical potential. In contrast to the GEMC-NVT method, the volume of each box is allowed to fluctuate independently from the other box. As a result, the total volume of the system is not constant and the pressure must be specified as a simulation parameter in addition to the temperature. This is consistent with the Gibbs phase rule, which requires the specification of two intensive variables (e.g. pressure and temperature) to fully specify a state with two phases.

The probability distribution of this ensemble is very similar to its NVT counterpart (see equation 6.41),

$$p_m \propto (V^l)^{N^l} (V^v)^{N^v} e^{-\beta U \left(\mathbf{s}^{N^l}\right) - \beta U \left(\mathbf{s}^{N^v}\right) - \beta P \left(V^l + V^v\right)}$$

$$\tag{6.49}$$

Thermal equilibration moves can be derived from the above probability distribution. They involve the usual translation, rotation and regrowth moves. The functional form of the acceptance rule is the same as those exposed in sections 6.1.1, 6.1.2, 6.1.3, 6.3.3. Similarly, particle swap moves are carried out in the same way as it was exposed in section 6.4.1. Finally, volume moves are carried out by picking one of the boxes and performing a volume change using the protocol exposed in section 6.2.1.

6.5 Multicomponent Systems

Sections 6.1-6.4 have each been developed for pure component systems. The Monte Carlo moves and acceptance criteria for multicomponent systems are straightforward extensions of the pure component moves. The only modification needed to translate, rotate and regrow molecules is to first select a species. In these moves, a species is selected randomly in proportion to its mole fraction N_i/N . When inserting and deleting a molecule, the mole fractions of each species change. In these cases, a species in a multicomponent system is selected instead with uniform probability $1/N_{species}$. In either case, species selection is symmetric for both forward and reverse moves and so cancels from the acceptance criterion.

6.6 Appendix

6.6.1 Inserting a Molecule Randomly

To insert a molecule, a position, orientation and molecular conformation must each be selected. The probability of inserting the new molecule at a random location is $d\mathbf{r}/V$, where $d\mathbf{r}$ is a Cartesian volume element of a single atom. The probability of choosing the molecule orientation is $d\mathbf{q}_{rot}/Z_{rot}$, which for a linear molecule is $d\cos(\theta)d\phi/4\pi$ and for a nonlinear molecule is $d\cos(\theta)d\phi/4\pi^2$. The probability of the molecule conformation only depends on the remaining 2M-5 internal bond angles, dihedral angles and improper angles \mathbf{q}_{int} . A thermal ensemble of configurations is Boltzmann distributed $e^{-\beta U(\mathbf{q}_{int})}/Z_{int}$. The overall probability α_{mn} is

$$\alpha_{mn} = \frac{d\mathbf{r}}{V} \frac{d\mathbf{q}_{rot}}{Z_{rot}} \frac{e^{-\beta U(\mathbf{q}_{int,N+1,n})}}{Z_{int}} d\mathbf{q}_{int} = \frac{e^{-\beta U(\mathbf{q}_{N+1,n})}}{Z(1,V,T)} d\mathbf{q}.$$
(6.50)

where we have used Eq. (6.13) to recover Z(1, V, T) and recognized that only internal degrees of freedom contribute to the potential energy of the isolated N+1th molecule in microstate n, $U(\mathbf{q}_{N+1,n})=U(\mathbf{q}_{int,N+1,n})$.

For a point particle with no rotational or internal degrees of freedom, α_{mn} reduces to $d\mathbf{r}/V$. For molecules with internal flexibility, a library of configurations distributed according to $e^{-\beta U(\mathbf{q}_{int})}/Z_{int}$ can be generated from a single molecule MC simulation. In the reverse move, 1 of the N+1 particles is randomly selected for deletion. The probability α_{nm} of picking the molecule we just inserted is

$$\alpha_{nm} = \frac{1}{N+1} \tag{6.51}$$

The acceptance probability for a random insertion move is

$$\ln\left(\frac{\alpha_{mn}}{\alpha_{nm}}\frac{p_m}{p_n}\right) = \beta\left[\Delta U - U(\mathbf{q}_{N+1})\right] - \beta\mu + \ln\left(\frac{N+1}{Q(1,V,T)}\right)$$
(6.52)

where $U(\mathbf{q}_{N+1})$ is the intramolecular potential energy of the inserted molecule. Q(1, V, T) is typically not known a priori, nor is it easily estimated. Substituting Eq. (6.12) into Eq. (6.52) and absorbing $Q_{rot+int}$ into a shifted chemical potential μ'

$$\mu' = \mu - k_B T \ln(Q_{rot+int}) \tag{6.53}$$

gives the acceptance criteria for inserting a molecule

$$\ln\left(\frac{\alpha_{mn}}{\alpha_{nm}}\frac{p_m}{p_n}\right) = \beta\left[\Delta U - U(\mathbf{q}_{N+1})\right] - \beta\mu' + \ln\left(\frac{(N+1)\Lambda^3}{V}\right). \tag{6.54}$$

The terms absorbed into μ' are intensive and therefore GCMC simulations using Eq. (6.54) will converge to a specific average density. However, the value of μ' that corresponds to the converged density will *not* match tabulated values of μ computed from experimental data.

Substituting Eq. (6.23) into Eq. (6.52) gives

$$\ln\left(\frac{\alpha_{mn}}{\alpha_{nm}}\frac{p_m}{p_n}\right) = \beta \left[\Delta U - U(\mathbf{q}_{N+1})\right] + \ln\left(\frac{N+1}{\beta f V}\right)$$
(6.55)

where no terms have been absorbed into the fugacity f. Note also that the partition function has completely been eliminated from the acceptance criteria.

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6.6.2 Deleting a Molecule Inserted Randomly

The probability α_{mn} of choosing a molecule to delete is

$$\alpha_{mn} = \frac{1}{N} \tag{6.56}$$

The probability α_{nm} of inserting that molecule back in is

$$\alpha_{nm} = \frac{e^{-\beta U(\mathbf{q})}}{Z(1, V, T)} d\mathbf{q}$$
(6.57)

The acceptance probability for deleting a molecule inserted randomly is

$$\ln\left(\frac{\alpha_{mn}}{\alpha_{nm}}\frac{p_m}{p_n}\right) = \beta\left[\Delta U + U(\mathbf{q}_N)\right] + \beta\mu' + \ln\left(\frac{V}{N\Lambda^3}\right)$$
(6.58)

$$= \beta \left[\Delta U + U(\mathbf{q}_N) \right] + \ln \left(\frac{\beta f V}{N} \right)$$
 (6.59)

Note that in ΔU is defined differently in Eqs. (6.54) and (6.55) than in Eqs. (6.58) and (6.59). In the former, the new configuration has more molecules, $\Delta U = U(\mathbf{q}_n^{N+1}) - U(\mathbf{q}_m^N)$. In the latter, the new configuration has fewer molecules, $\Delta U = U(\mathbf{q}_n^{N-1}) - U(\mathbf{q}_m^N)$.