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- 1. Introduction (what is GOMC?)
- 2. How to get the software
- 3. Building the code
  - a. Required compilers, libraries, programs (this includes psfgen and tcl which are needed to build the PSF files)
  - b. Supported operating systems
  - c. Serial
  - d. GPU
- 4. Input file formats
  - a. PDB, PSF, parameter, control file (define each part of the files here)
- 5. Output file formats
  - a. Working with VMD
- 6. System setup
  - a. Building initial coordinates with Packmol
  - b. Creating a PSF file
  - c. Creating the control file
- 7. Running the simulation
- 8. Examples
- 9. How to get help/technical support.
- 10. Summary of algorithms used in the code

# **Tutorial Overview**

This document will introduce a new user to how to download, compile and run the GOMC molecular simulation code. A working knowledge of statistical physics is recommended as a perquisite to understanding this tutorial.

To demonstrate the capabilities of the code, the user is guide through the process of downloading and compiling a GOMC executable. That executable is then used to perform saturated vapor liquid equilibria (VLE) studies on systems of pure isobutane (R600a), a branched alkane that is seeing increasing use as a refrigerant/propellant.

## http://en.wikipedia.org/wiki/Isobutane

The Transferable Potentials for Phase Equilibria (TraPPE) united atom (UA) forcefield is used to describe the molecular geometry constraints and the intermolecular interactions.

# Introduction (what is GOMC?)

Monte Carlo (MC) simulations are a kind of simulations that are driven by stochastic processes. The "GO" stands for GPU Optimized, as this code was intended to run optimally on modern graphics process hardware.

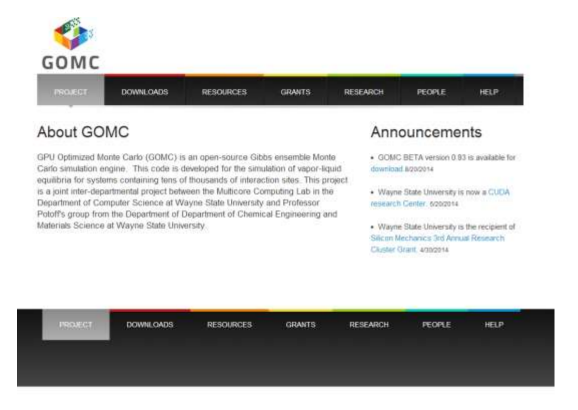
More specifically, this engine includes serial and GPU-optimized (multi-threaded) codes designed to run Markov chain Boltzmann sampling of chemical systems -- effectively sets of points defined by topological maps and interaction algorithms in a simulation box. From statistical mechanics we know this is one way to sample phase space and model chemical systems.

GOMC currently supports both canonical and Gibbs ensemble simulations. Support for GCMC and GEMC-NPT will be added shortly. GOMC uses widely used simulations file types (PDB, CHARMM-style parameter file, PSF). GOMC includes configurational bias algorithms for both linear and branched systems. Support for cyclic molecules and charged systems is being added.

# How to get the software

Currently the latest public code builds, the project logo, manual, and other resources can be obtained via the website:

gomc.eng.wayne.edu



The code can be found under the download tab, just below and to the right of the logo. When new betas (or eventually release builds) are announced, they will replace the prior code under the downloads tab. An announcement will be posted on the front page to notify users.

Currently version control is handled through an internal SVN system maintained by the developers at Wayne State University. The posted builds are "frozen" versions of the code that have been validated for a number of systems and ensembles. Eventually the project will be hosted as a Git repository to allow for greater collaboration and a faster means of spotting new releases.

# **Platform and Software Requirements**

# **Supported Operating Systems**

GOMC officially supports **Windows 7**, **8**, and most modern distributions of **Linux** (see the next section). This software may compile on recent versions of **OS X**, but that platform is not officially supported.

# **Required Software Prequisites**

GOMC has some mild software requirements, which are widely available for Linux operating systems. Required software are:

- 1. C++03 Compliant Compiler
  - a. Linux/OS X
    - i. icc (Intel c++ compiler)

Type...

icc --version

...in a terminal. If gives a version number 4.4 or later, you're all set. If it's older than 4.4 (released in 2009), we recommend upgrading.

In Linux, the Intel compiler will generally produce the fastest serial executables (when running on Intel Core processors).

ii. g++ (GNU GCC)

Type...

g++ --version

...in a terminal. If gives a version number 4.4 or later, you're all set. If it's older than 4.4 (released in 2009), we recommend upgrading.

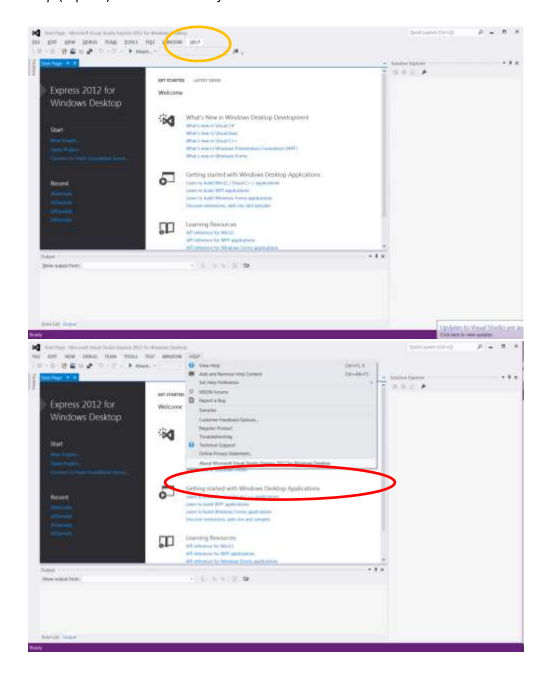
### b. Windows

i. Visual Studio

Microsoft's Visual Studio 2010 or later is recommended.

Check version:

Help (top tab) → About Microsoft Visual Studio



# 2. cmake (if compiling on Linux)

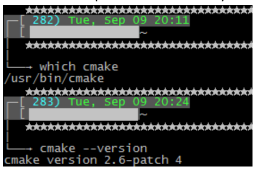
To check if cmake is installed

which cmake

To check version number:

cmake --version

Here's an example from one of our systems:



# 3. nvcc/CUDA libs

The GPU builds of the code requires NVIDIA's CUDA 6.0 or newer...

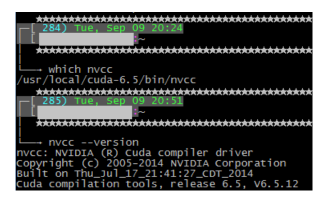
To check if nvcc is installed

which nvcc

To check version number:

nvcc --version

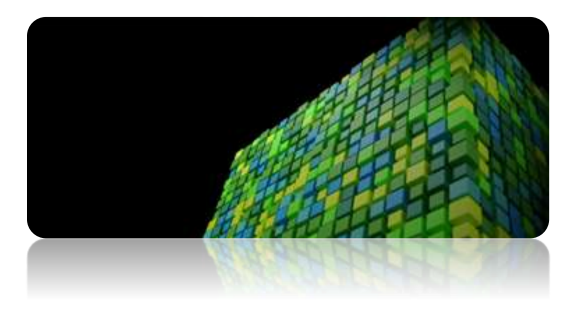
Here's an example from one of our systems:



CUDA is viewed as an essential requirement, but is not used to compile the serial code, which

can be compiled on systems without CUDA.

To download CUDA visit NVIDIA's webpage: https://developer.nvidia.com/cuda-downloads



CUDA is required to compile the GPU executable in both Windows and Linux. Please refer to CUDA Developer webpages to select an appropriate version for the desired platform.

To install CUDA in Linux *root/sudo* privileges are generally required. In Windows, administrative access is required.

# **Highly Recommended Software Tools**

# **NOTE:**

These programs are used in this manual and are generally to be considered <u>necessary</u> for its examples.

# **VMD**

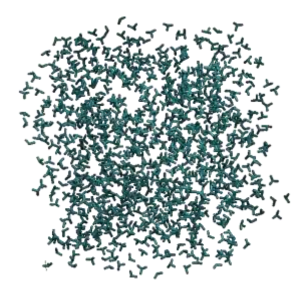
VMD (Visual Molecular Dynamics) is a 3-D visualization and manipulation engine for molecular systems written in C-language. VMD is distributed and maintained by the University of Illinois at Urbana-Champaign. Its source and binaries are available free to download. It comes with a robust scripting engine capable of running python and tcl scripts.

More info can be found out here:

http://www.ks.uiuc.edu/Research/vmd/

GOMC uses the same fundamental file types – PDB (coordinates) and PSF (topology) as VMD, although it uses some special tricks to obey certain rules of those file formats.

One useful purpose of VMD is visualization of your systems.



(A system of united atom isobutane molecules is seen above.)

The most critical part of VMD, though is a tool called PSFGen. PSFGen uses a tcl or python script to generate a PDB and PSF file for a system of one or more molecules. It is perhaps the most convenient way to generate a compliant PSF file.

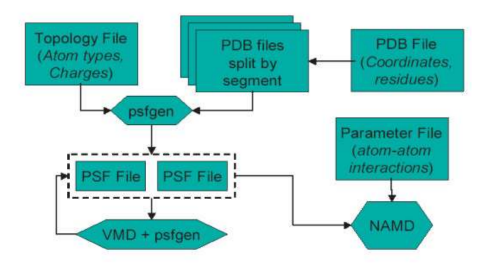


FIGURE: An overview of the PSFGen file generation process and its relationship to VMD/NAMD

To read more about PSFGen, please see:

Plugin homepage @ UIUC

http://www.ks.uiuc.edu/Research/vmd/plugins/psfgen/

"Generating a Protein Structure File (PSF)", part of the NAMD Tutorial from UIUC <a href="http://www.ks.uiuc.edu/Training/Tutorials/namd/namd-tutorial-html/node6.html">http://www.ks.uiuc.edu/Training/Tutorials/namd/namd-tutorial-html/node6.html</a>

In-Depth Overview [PDF]

http://www.ks.uiuc.edu/Research/vmd/plugins/psfgen/ug.pdf

#### **Packmol**

Packmol is a molecule packing tool created by José Mario Martínez, a professor of mathematics at the State University of Campinas, Brazil. It is written in Fortran and is free to download. More information is available on its homepage:

http://www.ime.unicamp.br/~martinez/packmol/

To compile it a Fortran language compiler is needed, such as gfortran. Many Linux distributions no longer automatically come with Fortran compilers, so this may need to be installed.

Packmol allows a specified number of molecules to be packed at defined separating distances within a certain region of space. Packmol's limitations include that it is unaware of topology – it treats each molecule or group of molecules it's packing as a rigid set of points.

## **WARNING**

Another more seriously limitation is that it is not aware of periodic boundary conditions (PBC). As a result, when using packmol to pack PDBs for GOMC, it is recommended to pack to a box 2 to 3 Angstroms smaller than the simulation box size. This prevents hard overlaps over the periodic boundary.

# Other Useful Software Tools

### **Grace**

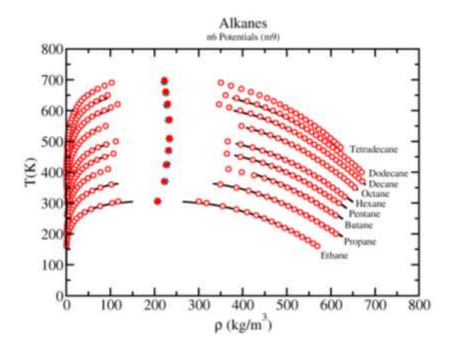
Grace is a piece of graphing software written and maintained by the Weizmann Institute of Science's Plasma Laboratory (Rehovot, Israel). Mostly used in Linux, it can also be compiled in Windows, although the developers warn it may be missing some functionality.



In-depth information and the source can be found on the project page, here:

# http://plasma-gate.weizmann.ac.il/Grace/

When compiled, Grace's executable in Linux is typically named "xmgrace". This tool allows the production of high quality, precise line and dot graphs. This makes it ideal for visualizing much of the thermodynamic data from the GOMC engine. Here is an example of the results of simulations of saturated VLE densities of linear alkanes produced with Grace:



# Cygwin

Cygwin provides Microsoft Windows users with a Unix-like environment and command-line interface. It offers Windows-compatible ports of common Linux applications. It's one option to assist in building and visualizing systems in Windows.

# https://cygwin.com/



The software is free and open source, licensed under the GNU General Public License version 3. Its primary maintainers are Red Hat Inc. and NetApp. One of the most impressive abilities of Cygwin is its ability to launch a full Windows-compatible X-server Window, which allows convenient visualization of Linux app GUIs. It is compatible with the Grace graphing software. In practice, this package behaves most analogously to a Linux virtual machine in Windows.

# **Compiling GOMC**

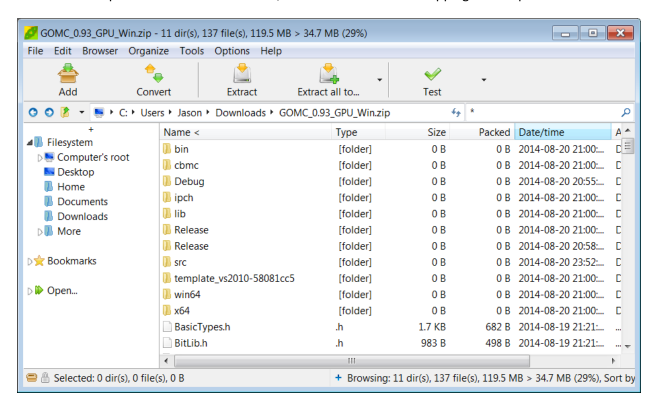
# Extracting the Code

GOMC is distributed as a compressed folder which contains the source and build system. To compile the code after downloading it, the first step is to extract the compressed build folder.

In Windows the folder for the GPU code is compressed using a standard \*.zip file format. To unzip simply use a utility like Peazip:

# http://peazip.sourceforge.net/

Here's an example of what the downloaded, code looks like when unzipping in Peazip...



In Linux the GPU and Serial codes are compressed using gzip and tar (\*.tar.gz). To extract, simply move to the desire folder and type:

tar –xzvf <file name>.tar.gz

Compiling the Code

GPU Code:

#### Compilation on Windows:

Once the code is extracted, to compile it on Windows, you need to load the project into Visual Studio. To do that, open the extracted folder, and then double click on the solution file of the desired Visual Studio version. After the solution is opened in Visual Studio, you can go to the "Build" menu and select "Build solution" to compile the code. You can compile either with release mode or debug mode by selecting the desired mode from the "Solution Configuration" drop box.

To run the project, simply click the run button or hit F5 on the keyboard.

## Compilation on Linux:

To compile the GPU code on Linux, just go into the directory of the project, and then type

```
make
```

You can configure the "makefile" file to choose different C compilers, select the desired compute capability, and to configure many more compilation flags.

The default compute capability is set to 3.0. To change the compute capability, go to the GENCODE\_FLAGS option, and set it to one of the compute capability flags that are defined in the file.

```
# CUDA code generation flags
GENCODE_SM10 := -gencode arch=compute_10,code=sm_10
GENCODE_SM20 := -gencode arch=compute_20,code=sm_21
GENCODE_SM30 := -gencode arch=compute_30,code=sm_30
GENCODE_SM35 := -gencode arch=compute_35,code=sm_35
GENCODE_FLAGS := $(GENCODE_SM30)
```

To run the program, run the executable "GOMC.out". To run this, the system's LD\_LIBRARY\_PATH will need to be configured to support CUDA (more on this later).

#### Serial Code:

Compilation on Windows:

See GPU "Compilation on Windows" section and follow an identical procedure for the released serial code. See README for instructions on how to use the CMake-GUI to build the configuration and solution files necessary for the Windows build.

Compilation on Linux:

In Linux the GPU code uses a simple makefile. Enter the directory and type

make all

... which will use the Makefile to compile a GPU-compatible executable called "GOMC.out". To run this the system's LD\_LIBRARY\_PATH will need to be configured to support CUDA (more on this later).

For the serial code – which uses cmake for compilation – simply go to the base directory and type:

```
./metamake.sh
```

This cmake script will create a directory named "bin". Enter this directory:

cd bin

...and type:

make

Two executables – GOMC\_Serial\_GEMC (Gibbs ensemble) and GOMC\_Serial\_NVT (NVT ensemble) will be produced. By default the distribution compiles in release mode. To compile in debug mode (if you're using the code as a developer) open the file "CMakeCache.txt", while still in the "bin" folder. This file contains information used by cmake to build the executables. To compile in debug mode, simply change the value after "CMAKE\_BUILD\_TYPE:STRING=" from "Release" to "Debug"...

```
# This is the CMakeCache file.
# For build in directory: [REDACTED]/Serial code/bin
# It was generated by CMake: /usr/bin/cmake
# You can edit this file to change values found and used by cmake.
# If you do not want to change any of the values, simply exit the editor.
# If you do want to change a value, simply edit, save, and exit the editor.
# The syntax for the file is as follows:
# KEY:TYPE=VALUE
# KEY is the name of a variable in the cache.
# TYPE is a hint to GUI's for the type of VALUE, DO NOT EDIT TYPE!.
# VALUE is the current value for the KEY.
*****************
# EXTERNAL cache entries
******************
//Path to a program.
CMAKE AR: FILEPATH=/usr/bin/ar
//Choose the type of build, options are: None Debug Release RelWithDebInfo
// MinSizeRel
CMAKE BUILD TYPE:STRING=Release
//Enable/Disable color output during build.
CMAKE COLOR MAKEFILE: BOOL=ON
//CXX compiler.
CMAKE CXX COMPILER: FILEPATH=/usr/bin/c++
```

To...

```
# This is the CMakeCache file.
# For build in directory: [REDACTED]/Serial code/bin
# It was generated by CMake: /usr/bin/cmake
# You can edit this file to change values found and used by cmake.
# If you do not want to change any of the values, simply exit the editor.
# If you do want to change a value, simply edit, save, and exit the editor.
# The syntax for the file is as follows:
# KEY:TYPE=VALUE
# KEY is the name of a variable in the cache.
# TYPE is a hint to GUI's for the type of VALUE, DO NOT EDIT TYPE!.
# VALUE is the current value for the KEY.
##########################
# EXTERNAL cache entries
##############################
//Path to a program.
CMAKE AR:FILEPATH=/usr/bin/ar
//Choose the type of build, options are: None Debug Release RelWithDebInfo
// MinSizeRel.
CMAKE BUILD TYPE:STRING=Debug
//Enable/Disable color output during build.
CMAKE COLOR MAKEFILE: BOOL=ON
//CXX compiler.
CMAKE CXX COMPILER:FILEPATH=/usr/bin/c++
```

# And retype the command:

### make

The outputted executables should now be compiled with debugger symbols.

You can also swap the compiler by modifying the "CMAKE\_CXX\_COMPILER" variable. For more information refer to the Cmake documentation.

# **Input File Formats**

A typical GOMC GEMC directory has the following:

- GOMC executable
- in.dat (proprietary control file)
- One (NVT ensemble) or two PDB files (Gibbs ensemble)
- One (NVT ensemble) or two PSF files (Gibbs ensemble)
- A CHARMM-style parameter file
- An exotic-style parameter file (may be empty, but must be present)

#### PDB

The PDB file stores coordinates for the simulation. The file format is widely adopted...

- Protein Databank (PDB) Files (plural: PDB files)
- Open format, well-documented
- Fixed-width format (hence white space is significant)
- Up to 13.5m page views a month; up to 55.8m FTP requests per month
- Used by NAMD, GROMACS, CHARMM, ACEMD, Amber

An overview of the PDB standard can be found here:

# http://www.wwpdb.org/docs.html

The advantage of PDB files are their ubiquity and thorough documentation. Disadvantages include limited fixed point floating precision for coordinates, unused space, and proprietary implementations creating inconsistencies.

One PDB file is required per box, so for NVT ensemble simulations one file is expected, for Gibbs ensemble two files are required.

GOMC recognizes the following keywords in PDB files:

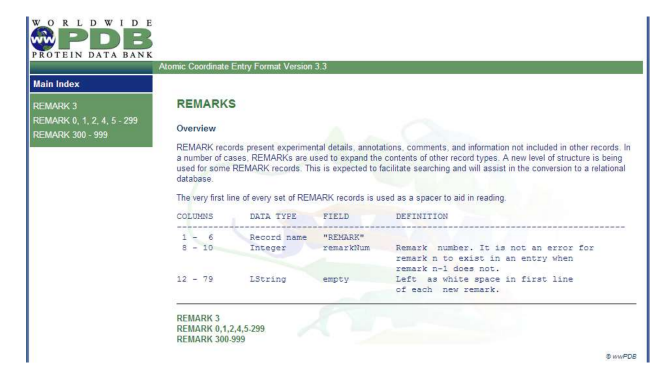
- ❖ REMARK
- CRYST1
- ATOM
- ❖ END

Currently **REMARK** is ignored (formerly it was used to store proprietary information in frames, e.g. step number, etc. Note, packmol typically leaves the following remark:

# REMARK original generated coordinate pdb file

...at the top of the file. Note this is another example of an inconsistency with the spec. As of the PDB

v3.30 specification the REMARK entry contains an identifying integer, which is supposed to occupy lines 8-10.



A file generated by packmol has "ori" in this position. Hence you may see future codes that are incompatible with this legacy kind of remarks.

Note also that the spaces 7 and 11 are not reserved; hence they may be used in proprietary specifications.

**CRYST1** can be used to store the cell dimensions, which can also be put as a tag in the proprietary control file.

http://www.wwpdb.org/documentation/format33/sect8.html#CRYST1

# Crystallographic and Coordinate Transformation Section

This section describes the geometry of the crystallographic experiment and the coordinate system transformations.

### CRYST1

#### Overview

The CRYST1 record presents the unit cell parameters, space group, and Z value. If the structure was not determined by crystallographic means, CRYST1 simply provides the unitary values, with an appropriate REMARK.

#### Record Format

COLUMNS		INS	DATA TYPE	FIELD	DEFINITION		
1		6	Record name	"CRYST1"	0-		
7	-	15	Real(9.3)	a	a (Angstroms).		
16	-	24	Real (9.3)	b	b (Angstroms).		
25	-	33	Real (9.3)	С	c (Angstroms).		
34	-	40	Real (7.2)	alpha	alpha (degrees).		
41	-	47	Real (7.2)	beta	beta (degrees).		
48	-	54	Real (7.2)	gamma	gamma (degrees).		
56	-	66	LString	sGroup	Space group.		
67	-	70	Integer	z	Z value.		

#### Details

- If the entry describes a structure determined by a technique other than X-ray crystallography, CRYST1 contains a = b = c = 1.0, alpha = beta = gamma = 90 degrees, space group = P.1, and Z = 1.
- The Hermann-Mauguin space group symbol is given without parenthesis, e.g., P 43 21 2. Please note that the screw
  axis is described as a two digit number.
- The full International Table's Hermann-Mauguin symbol is used, e.g., P 1 21 1 instead of P 21.
- For a rhombohedral space group in the hexagonal setting, the lattice type symbol used is H.
- The Z value is the number of polymeric chains in a unit cell. In the case of heteropolymers, Z is the number of occurrences of the most populous chain.

As an example, given two chains A and B, each with a different sequence, and the space group P 2 that has two equipoints in the standard unit cell, the following table gives the correct Z value.

Here's an example...

#### REMARK 8 **REMARK** 8901234567890123456789012345678901234567890123456789012345678901234567 890 71.490 71.490 71.490 90.00 90.00 CRYST1 90.00 P 1 1 C1 ISB A 24.378 36.667 45.645 0.00 0.00 ATOM ISB C

This file indicates a box size of 71.49 Angstroms per side. The '90.00' entries, indicate the cell is cubic.

# NOTE:

Non-cubic cells are not yet supported in this code.

The main entry in the PDB file are **ATOM** entries. The keyword "ATOM" is always followed by two spaces. An entry has a number of fields....

#### **Coordinate Section**

The Coordinate Section contains the collection of atomic coordinates as well as the MODEL and ENDMDL records.

#### **ATOM**

#### Overview

The ATOM records present the atomic coordinates for standard amino acids and nucleotides. They also present the occupancy and temperature factor for each atom. Non-polymer chemical coordinates use the HETATM record type. The element symbol is always present on each ATOM record; charge is optional.

Changes in ATOM/HETATM records result from the standardization atom and residue nomenclature. This nomenclature is described in the Chemical Component Dictionary (ftp://ftp.wwpdb.org/pub/pdb/data/monomers).

#### Record Format

COLUMNS	DATA TYPE	FIELD	DEFINITION
1 - 6	Record name	"ATOM "	
7 - 11	Integer	serial	Atom serial number.
13 - 16	Atom	name	Atom name.
17	Character	altLoc	Alternate location indicator.
18 - 20	Residue name	resName	Residue name.
22	Character	chainID	Chain identifier.
23 - 26	Integer	resSeq	Residue sequence number.
27	AChar	iCode	Code for insertion of residues.
31 - 38	Real(8.3)	x	Orthogonal coordinates for X in Angstroms.
39 - 46	Real(8.3)	У	Orthogonal coordinates for Y in Angstroms.
47 - 54	Real(8.3)	Z	Orthogonal coordinates for Z in Angstroms.
55 - 60	Real(6.2)	occupancy	Occupancy.
61 - 66	Real(6.2)	tempFactor	Temperature factor.
77 - 78	LString(2)	element	Element symbol, right-justified.
79 - 80	LString(2)	charge	Charge on the atom.

#### Details

- · ATOM records for proteins are listed from amino to carboxyl terminus.
- · Nucleic acid residues are listed from the 5' to the 3' terminus.
- Alignment of one-letter atom name such as C starts at column 14, while two-letter atom name such as FE starts at column 13.
- · Atom nomenclature begins with atom type.
- · No ordering is specified for polysaccharides.
- · Non-blank alphanumerical character is used for chain identifier.
- The list of ATOM records in a chain is terminated by a TER record.
- . If more than one model is present in the entry, each model is delimited by MODEL and ENDMDL records.
- AltLoc is the place holder to indicate alternate conformation. The alternate conformation can be in the entire polymer
  chain, or several residues or partial residue (several atoms within one residue). If an atom is provided in more than one
  position, then a non-blank alternate location indicator must be used for each of the atomic positions. Within a residue,
  all atoms that are associated with each other in a given conformation are assigned the same alternate position indicator.
  There are two ways of representing alternate conformation- either at atom level or at residue level (see examples).
- For atoms that are in alternate sites indicated by the alternate site indicator, sorting of atoms in the ATOM/HETATM list
  uses the following general rules:
  - In the simple case that involves a few atoms or a few residues with alternate sites, the coordinates occur one after the other in the entry.
  - In the case of a large heterogen groups which are disordered, the atoms for each conformer are listed together.
- Alphabet letters are commonly used for insertion code. The insertion code is used when two residues have the same numbering. The combination of residue numbering and insertion code defines the unique residue.
- . If the depositor provides the data, then the isotropic B value is given for the temperature factor.
- If there are neither isotropic B values from the depositor, nor anisotropic temperature factors in ANISOU, then the default value of 0.0 is used for the temperature factor.
- Columns 79 80 indicate any charge on the atom, e.g., 2+, 1-. In most cases, these are blank.
- For refinements with program REFMAC prior 5.5.0042 which use TLS refinement, the values of B may include only the TLS contribution to the isotropic temperature factor rather than the full isotropic value.

The key parameters are the coordinates x, y, and z. The precision in these is limited to eight whole decimal digits and three fractional decimal digits.

Other important entries are the residue name, atom name, and chainID. Numbering is important primarily in that it represents an inconvenience in packing/loading large systems. Looking at the previous example:

#### REMARK <u>2\_\_\_\_3\_\_\_4\_\_\_5\_\_\_6\_\_\_7\_\_\_</u> REMARK 890123456789012345678901234567890123456789012345678901234567 890 71,490 71.490 90.00 CRYST1 71.490 90.00 90.00 P 1 ATOM 1 C1 ISB A 24.378 36.667 45.645 0.00 0.00 ISB C

The atom name is "C1" and residue name is "ISB". The PSF file (next section) contains a lookup table of atoms. These contain the atom name from the PDB and the name of the atom kind in the parameter file it corresponds to. As multiple different atom names will all correspond to the same parameter, these can be viewed "atom aliases" of sorts. The chain letter (in this case 'A') is sometimes used when packing a number of PDBs into a single PDB file.

# A few important Notes/Warnings on Undocument PDB Format Conventions:

- While it is only explicitly stated in some other sections of the PDB file, the general convention observed by most codes is to right align, when padding with white space.
- Some codes (including PSFGen/VMD) use the 21 unused character to add a fourth letter to the residue (molecule name). This extension is currently support, but is unofficial and hence may change in the future.
- VMD requires a constant number of ATOMs in a multi-frame PDB (multiple records terminated by "END" in a single file). To compensate for this all atoms from all boxes in the system are written to the outputted PDBs from this code.
- For atoms not currently in a box, the coordinates are set to <0.00, 0.00, 0.00>
- The occupancy is commonly just set to "1.00" and is left unused by many codes. We recycle this legacy parameter by using it to denote the box a particle is in, in our outputted PDBs (box  $0 \rightarrow$ occupancy=0.00; box  $1 \rightarrow$  occupancy=1.00)
- As the x, y, and z coordinates are fixed point with only three digits of precision, the energy values you get when restarting may be mildly different, particularly for bonded interactions due to roundoff in the coordinates. This will eventually be remedied by the implementation of a full-precision trajectory (e.g. DCD) file.
- The "ISB" entry in columns 73-75 is not an official part of the PDB standard. This is a proprietary entry called "Segname", which has been embraced by NAMD and some other codes.

\_\_\_\_\_

# Pending Improvement

Note, currently a box is limited to holding up to 9,999 residues, or 99,999 atom entries, if sticking to decimal integer numbering. Like NAMD/X-PLOR, an upcoming build of the code will add support for using hexadecimal in the molecule numbering if 9,999 residues (see: NAMD) and using the alphanumeric overflow defined in the X-PLOR spec.

# This will allow for a maximum of:

9,999 (standard uint) + 999 x 26 (overflow, alphanumeric)  $\rightarrow$  35,973 molecules (residue #s) 99,999 (standard uint) + 9,999 x 26 (overflow, alphanumeric)  $\rightarrow$  359,973 ATOM entries

Past either of these and an "overflow" solution is hit, where it will print stars to the respective field. This file should still be readable, but is consider unsafe in terms of portability, as it will work for GOMC and NAMD, but may fail for other codes.

\_\_\_\_\_

A frame in the PDB file is terminated with the keyword END.

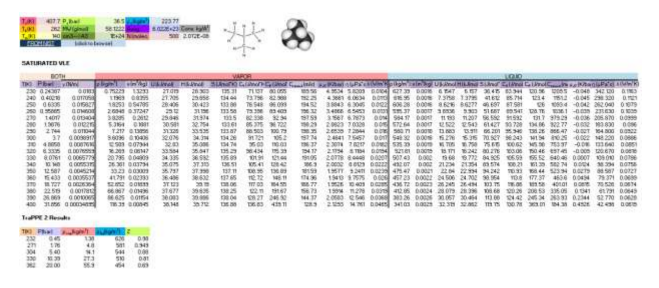
With that overview of the format in mind, here's how a PDB file is typically built.

First, a single molecule PDB is obtained. In this example, the QM software package Gaussian was used to draw the molecule, which was then edited by hand to adhere to the PDB spec properly. The end result is a PDB for a single molecule:

REMARK	1 File	created	by	GaussView 5	5.0.8	
ATOM	1 C1	ISB	1	0.911	-0.313	0.000
C						
ATOM	2 C2	ISB	1	1.424	-1.765	0.000
C						
ATOM	3 C3	ISB	1	-0.629	0.313	0.000
C						
ATOM	4 C4	ISB	1	1.424	0.413	-1.257
C						
END						

Next packings are calculated, to place the simulation in a region of vapor-liquid coexistence. There are a couple of ways to do this in Gibbs ensemble:

- 1. Pack both boxes to a single "middle" density, which is an average of the liquid and vapor densities.
- 2. Same as 1, but add a modest amount to axis of one box (e.g. 10-30 A). This technique can be handy in the constant pressure Gibbs ensemble.
- 3. Pack one box to the predicted liquid density and the other to the vapor density.



A good reference for getting the information needed to estimate packing is the NIST Web Book database of pure compounds:

# http://webbook.nist.gov/chemistry/

After packing is determined, a basic pack can be performed with a packmol script. Here is one example:

tolerance 3.0
filetype pdb
output STEP2\_ISB\_packed\_BOX\_0.pdb

structure isobutane.pdb number 1000 inside box 0. 0. 0. 70.20 70.20 70.20 end structure

Packmol scripts are typically saved with the extension \*.inp, so this might be named "pack\_isobutane.inp".

To run the script, we type:

./packmol < pack\_isobutane.inp

A more efficient way than packing individual simulations by hand is to embed a packmol script in a larger script, which reads in a table of comma or space separated values for the packing box sizes, as calculated via a spreadsheet.

Here's an example snippet from tcl:

When packing large files, packmol will adjust for this by incrementing the chain letter. This will cause issues with PSFGen. The solution is to use "grep" in Linux (or a similar method in Windows) to separate the PDB into several separate PDB files, one per chain. Each collection of residues of a specific kind will typically end up in a file together using this methodology.

The PDB output packmol is fed into PSFGen, along with a topology file. PSFGen will output a final PDB, which will be nearly identical to the packmol generated file. The only observable difference is generally the addition of a "segname", the aforementioned field VMD/NAMD insert into an unspecified set of columns in **ATOM** entries. The two PDB files generated by PSFGen (or one if an NVT ensemble simulation is being prepped) will serve as the coordinates files for a new simulation.

#### **PSF File**

The PSF file stores the topology, mass, charges, and atom identities of molecules in the system.

- Protein Structure File (PSF)
- Space-separated file
- Used by NAMD, CHARMM, X-PLOR

The PSF file is not as robustly documented as the PDB format, but a basic description of it can be found here:

### http://www.ks.uiuc.edu/Training/Tutorials/namd/namd-tutorial-win-html/node24.html

The PSF file is generally laid out in a series of sections. At the top of each section there is typically a line with a numeric value. This value lists the number of entries in that section (lines can contain multiple entries; a dihedral, for example has two quadruplet entries of atom indices per line). Note, that outside

the remarks and atom section this number is typically smaller than the number of lines by a factor of 2 to 4.

PSF files always start with the string "PSF" on their first line.

GOMC reuses PSF reading code from NAMD, hence it should have much of the same flexibility and limitations. By section, the segments of a PSF file are:

- TITLE: remarks on the file
- BONDS: the bonds (if applicable) in molecules
- ANGLE: the bonds (if applicable) in molecules
- DIHEDRAL: the bonds (if applicable) in molecules
- IMPROPER: the bonds (if applicable) in molecules
- (other sections such as cross terms)

The code currently skips the title section and reads in the bonds, angles, dihedrals and impropers.

## A few important Notes/Warnings:

- The PSF file format is a highly redundant file format. For example it repeats identical topology of thousands of molecules of a common kind, in some cases. GOMC follows the same approach as NAMD, allowing this excess information externally and compiling it in the code.
- Other sections (e.g. cross terms) are unsupported or legacy, hence are ignored.
- Following the restrictions of VMD, the order of the PSF atoms must match the order in the PDB file.
- Improper entries are read and stored, but are not currently used. Support will eventually be added for this.

The PSF file is typically generated using PSFGen. It is convenient to make a script to do this. For example the script:

```
psfgen << ENDMOL
topology ./Top_Branched_Alkanes.inp
segment ISB {
    pdb ./STEP2_ISB_packed_BOX_0.pdb
    first none
    last none
}
coordpdb ./STEP2_ISB_packed_BOX_0.pdb ISB
writepsf ./STEP3_START_ISB_sys_BOX_0.psf
writepdb ./STEP3_START_ISB_sys_BOX_0.pdb</pre>
```

Typically one script is run per box to generate a finalized PDB/PSF for that box. The script requires one additional file, the NAMD-style topology file. While GOMC does not directly read or interact with this file, it's typically used to generate the PSF and hence is consider one of the integral file types. It will be briefly discussed in the following section.

As with the packmol script, its often useful to pack the PSFGen script in a tcl or python script. Here's an example:

```
#(20) Pack copies of base model in first box.
                         #(20) Pack copies of base model in first box.
exec packmol << "
    tolerance 3.0
    filetype pdb
    output STEP2_${compound_shorthand}_packed_BOX_0.pdb</pre>
                               structure STEP1_${compound_shorthand}.pdb
number ${num_per_box}
inside box 1. 1. 1. ${pack_l}. ${pack_l}.
end structure"
                         #(21) Pack copies of base model in second box.
                         exec packmol
                               tolerance 3.0
filetype pdb
output STEP2_${compound_shorthand}_packed_BOX_1.pdb
                              structure STEP1_${compound_shorthand}.pdb
number ${num_per_box}
inside box 1. 1. 1. ${pack_v}. ${pack_v}. ${pack_v}.
end structure"
                         #(22) Run tcl script to build the final PDB/PSF for box 0
                         exec psfgen << "
topology ./Top_Branched_Alkanes.inp
                               segment ${compound_shorthand} {
   pdb ./step2_${compound_shorthand}_packed_BOX_0.pdb
   first none
                                      last none
                               coordpdb ./STEP2_${compound_shorthand}_packed_BOX_0.pdb ${c\
ompound_shorthand}
                               writepsf ./STEP3_START_${compound_shorthand}_sys_BOX_0.psf
writepdb ./STEP3_START_${compound_shorthand}_sys_BOX_0.pdb
                         #(23) Run tcl script to build the final PDB/PSF for box 1
                         exec psfgen << " '
topology ./Top_Branched_Alkanes.inp</pre>
                               segment ${compound_shorthand} {
   pdb ./STEP2_${compound_shorthand}_packed_BOX_1.pdb
   first none
                                      last none
                               coordpdb ./STEP2_${compound_shorthand}_packed_BOX_1.pdb ${c\}
ompound_shorthand}
                               writepsf ./STEP3_START_${compound_shorthand}_sys_BOX_1.psf
writepdb ./STEP3_START_${compound_shorthand}_sys_BOX_1.pdb
```

Here's a peek at how the generated PSF file looks for a packed isobutane system (abridged):

**PSF** 

3 !NTITLE
REMARKS original generated structure x-plor psf file
REMARKS topology ./Top\_Branched\_Alkanes.inp
REMARKS segment ISB { first NONE; last NONE; auto angles dihedrals }

4000 !NATOM
1 ISB 1 ISB C1 CH1 0.000000 13.0190

2	ISB	1	ISB	C2	СН3	0.000000	) :	15.0350		0
3	ISB	1	ISB	C3	СН3	0.000000	) :	15.0350		0
4	ISB	1	ISB	C4	СН3	0.000000		15.0350		0
	ISB	2	ISB	C1	CH1	0.000000		13.0190		0
6	ISB	2	ISB	C2	СН3	0.000000		15.0350		0
	ISB	2	ISB	C3	СНЗ	0.000000		15.0350		0
8	ISB	2	ISB	C4	СН3	0.000000		15.0350		0
9	ISB	3	ISB	C1	CH1	0.000000		13.0190		0
10	ISB	3	ISB	C2	СНЗ	0.000000	) :	15.0350		0
3997		1000	ISB	C1	CH1	0.000000		13.0190		0
	ISB		ISB	C2	CH3	0.000000		15.0350		0
3999		1000		C3	CH3	0.000000		15.0350		0
4000	ISB	1000	ISB	C4	СН3	0.000000	) :	15.0350		0
3000	!NBC	MD: ha	onds							
	: NDC	2		1	3	1	4	5	6	
1 5 9		7		5	3 8	9	10	5 9	11	
9		12	1		14	13	15	13	16	
17		18	1		19	17	20	21	22	
21 25		23 28	2	9	24 30	25 29	26 31	25 29	27 32	
33		34	3		35	33	36	37	38	
		JT	,	,	33	33	30	31	30	
39	977	3980		3981	3982	3981	3983	3981	3984	
3985		986	398		3987	3985	3988	3989	3990	
3989		991	398		3992	3993	3994	3993	3995	
3993	3	3996	399	/	3998	3997	3999	3997	4000	
3000	INTH	IETA: a	angle	S						
2		1		4	2	1	3	3	1	4
6		5		8	6	5	7	7	5	4 8 12
10		9	1		10	9	11	11	9	12
14		13	1		14	13	15	15	13	16
18 22		17	2		18	17	19	19	17	20
26		21 25	2		22 26	21 25	23 27	23 27	21 25	24 28
20		23		O	20	23	21	<i>L1</i>	23	20

O !NPHI: dihedrals

0 !NIMPHI: impropers

# **Topology File**

The topology is a whitespace separated file format which contains a list of atoms (with their masses), plus a list of residue information (charges, composition, and topology). It is essentially a non-redundant lookup table equivalent to the PSF file.

This is followed by a series of residues, which tell PSFGen what's bonded to what. Each residue is comprised of four key elements:

- A header beginning with the keyword **RESI** with the residue name and net charge
- A body with multiple **ATOM** entries (not to be confused with the PDB-style entries of the same name) which list the partial charge on the particle and what kind of atom each named atom in a specific molecule/residue is.
- A section of lines starting with the word **BOND** containing pairs of bonded atoms (typically up to 3 per line)
- A closing section with instructions for PSFGen.

Here's an example of a residue definition for isobutane:

RESI ISB	0.00 !	isobutane	- Trappe
GROUP			
ATOM C1 CH1	0.00 !	C3	
ATOM C2 CH3	0.00 !	C2-C1	
ATOM C3 CH3	0.00 !	C4	
ATOM C4 CH3	0.00 !		
BOND C1 C2 C1	C3 C1 C4		
PATCHING FIRS	NONE LAST	NONE	

Here's a full parameter file prepared to pack a system of isobutane:

```
*
* Custom top file -- branched alkanes
*
```

```
1 1
         CH3
                 15.035 C
MASS
       1
MASS
          CH1
                 13.019 C
                0.00 ! isobutane - Trappe
RESI ISB
GROUP
ATOM C1 CH1
                0.00 !
                            C3
ATOM C2 CH3
                0.00 ! c2-c1
                0.00 !
ATOM C3 CH3
                            C4
                0.00 !
ATOM C4 CH3
BOND C1 C2 C1 C3 C1 C4
PATCHING FIRS NONE LAST NONE
```

### **END**

Note the keyword **END** must be used to terminate this file and keywords related to the autogeneration process must be placed near the top of the file, after the **MASS** definitions

More in-depth information can be found in the following links...

http://www.ks.uiuc.edu/Training/Tutorials/science/topology/topology-tutorial.pdf

"NAMD Tutorial: 4. Examining the Topology File"

http://www.ks.uiuc.edu/Training/Tutorials/science/topology/topology-html/node4.html

"Developing Topology and Parameter Files"

http://www.ks.uiuc.edu/Training/Tutorials/science/forcefield-tutorial/forcefield-html/node6.html

"NAMD Tutorial: 25. Topology Files"

http://www.ks.uiuc.edu/Training/Tutorials/namd/namd-tutorial-win-html/node25.html

...courtesy of UIUC

<sup>&</sup>quot;Topology Tutorial" (PDF, in-depth)

## Parameter File(s):

Currently GOMC uses a single parameter file. However the user must specify from two choices of the kind of parameter file:

- CHARMM (Chemistry at HARvard Molecular Mechanics) compatible parameter file
- "exotic" parameter file

If the parameter file type is not specified or if the chosen file is missing an error will be thrown.

Both forcefield file options are whitespace separated files, with sections preceded by a tag. When a known tag (representing a kind of molecular interaction in the model) is encountered, reading of that section of the forcefield begins. Comments (anything after a \* or !) and whitespace are ignored. Reading concludes when the end of the file is reached or another section tag is encountered.

# CHARMM format parameter file

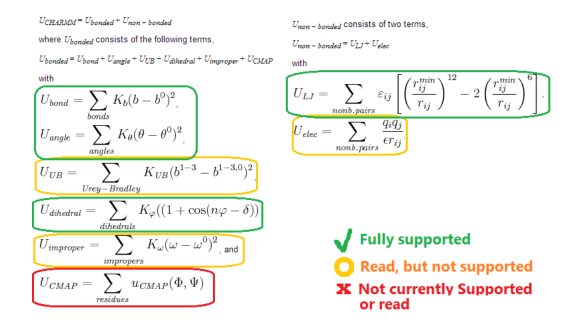
CHARMM contains a widely used model used in Monte Carlo and molecular dynamics simulations for describing energies. It is intended to be compatible with other codes that use this format, such as NAMD.

For a general overview of the CHARMM forcefield, see:

http://www.charmmtutorial.org/index.php/The Energy Function

(Cont'd on next page)

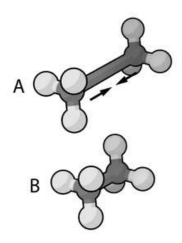
#### Here's the basic CHARMM contributions:



As seen above, the following are recognized, read and used:

# BONDS

- Quadratic expression describing bond stretching based on bond length (b) in Angstrom
- Typically is ignored as bonds are rigid for Monte Carlo sims. To specify that it is to be ignored put a very large value i.e. "99999999999" for Kb.



[Image Courtesy of Wikimedia Commons]

# ANGLES

 $\circ$  Describes the conformational behavior of an angle ( $\vartheta$ ) between three atoms , one of which is shared branch point to the other two.

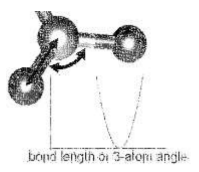


Image Courtesy of Wikimedia Commons

• **DIHEDRALS** Describes crankshaft like rotation behavior about a central bond in a series of three consecutive bonds (rotation is given as  $\varphi$ )

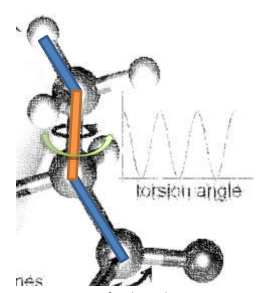


Image Courtesy of Wikimedia Commons

NONBONDED This section should only be used if the NONBONDED\_MIE section is not being
used in the exotic parameter file. This section describes 12-6 (Lennard-Jones) nonbonded
interactions.

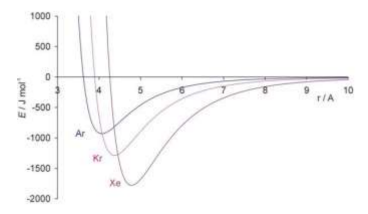


Image Courtesy of the Univ. of Bristol

Currently supported sections of the CHARMM compliant file include **BONDS**, **ANGLES**, **DIHEDRALS**, **IMPROPERS**, **NONBONDED**. Other sections such as **CMAP** are not currently read or supported.

**BONDS** (aka "bond stretching) are one key sections of the CHARMM-compliant file. Units for the  $K_b$  variable in this section are in kcal/mol (for  $K_b$ ); the  $b_0$  section (which represents the default bond length for that kind of pair) is measured in Angstroms.

### **NOTE:**

The Kb value may appear odd, but this is because a larger value corresponds to a more rigid bond. As Monte Carlo forcefields (e.g. TraPPE) typically treat molecules as rigid constructs, Kb is set to a large value -- 9999999999

Next comes an **ANGLES** (bond bending) section.  $\theta$  and  $\theta_0$  are commonly measured in degrees and  $K_\theta$  is measured in kcal/mol/K. These values are often given in the literature in Kelvin (K). To convert Kelvin to kcal/mol/K, multiply by the Boltzmann constant --  $k_b$ , 0.0019872041 kcal/mol.

Here is an example of what is necessary for isobutane:

```
ANGLES
!V(angle) = Ktheta(Theta - Theta0)**2
!V(Urey-Bradley) = Kub(S - S0)**2
!Ktheta: kcal/mole/rad**2
!Theta0: degrees
!Kub: kcal/mole/A**2 (Urey-Bradley)
!s0: A
!
! Ktheta (kcal/mol) = Ktheta (K) * Boltz. const.
!atom types
                    Ktheta
                                  Theta0
                                           Kub(?) SO(?)
CH3 CH1 CH3
                    62.100125
                                  112.00 ! Trappe 2
```

Some CHARMM **ANGLES** section entries include Urey-Bradley potentials ( $K_{ub}$ ,  $b_{ub}$ ), in addition to the standard quadratic angle potential. The constants related to this potential function are currently read in, but the logic has not been added to calculate this potential function. Support for this potential function will be added in later versions of the code.

The final major bonded interactions section of the CHARMM compliant parameter file are the **DIHEDRALS**. Each dihedral is composed of a dihedral series of 1 or more terms. Often there are 4 to 6 terms in a dihedral. Angles (for the dihedrals' deltas) are given in degrees.

(Since isobutane has no dihedral, here's the parameters pertaining to 2,3-dimethylbutane.)

```
DIHEDRALS
!v(dihedral) = Kchi(1 + cos(n(chi) - delta))
!Kchi: kcal/mole
!n: multiplicity
!delta: degrees
!
! Kchi (kcal/mol) = Kchi (K) * Boltz. const.
                                      delta
!atom types
                    Kchi
                                                          description
                                 n
                   -0.498907
                                        0.0
    CH1 CH1 X
                                 0
                                                          ! Trappe 2
                                        0.0
X
    CH1 CH1 X
                    0.851974
                                 1
                                                           Trappe 2
    CH1 CH1 X
                   -0.222269
                                 2
                                      180.0
X
                                                          ! Trappe 2
    CH1 CH1 X
                    0.876894
                                        0.0
                                                          ! Trappe 2
```

### NOTE:

The code allows the use of 'X' to indicate ambiguous positions on the ends. This is useful as this kind are often determined solely by the two middle atoms in the middle of the dihedral, according to the literature

**IMPROPERS** – energy parameters used to describe out-of-plane rocking are currently read in, but unused. The section is often blank. If it becomes necessary, algorithms to calculate the improper energy will need to be added.

The final section of the CHARMM style parameter file is the **NONBONDED**. In order to use TraPPE this section of the CHARMM compliant file will be critical. Here's an example with our isobutane potential models.

```
NONBONDED nbxmod 5 atom cdiel fshift vatom vdistance vfswitch -
!cutnb 14.0 ctofnb 12.0 ctonnb 10.0 eps 1.0 e14fac 1.0 wmin 1.5
!V(Lennard-Jones) = Eps,i,j[(Rmin,i,j/ri,j)**12 - 2(Rmin,i,j/ri,j)**6]
!epsilon: kcal/mole, Eps,i,j = sqrt(eps,i * eps,j)
!Rmin/2: A, Rmin,i,j = Rmin/2,i + Rmin/2,j
! Rmin = sig * (2^{(1/6)}) / 2 ; eps (kcal/mol) = eps (K) * Boltz.
const.
                                Boltzmann = 0.0019872041 kcal / (mol *
K)
!atom ignored
                 epsilon
                              Rmin/2
                                           ignored eps.1-4
                                                             Rmin/2,1-
CH3
       0.0
                -0.194745992
                              2.10461634058 ! Trappe 1
CH1
       0.0
                -0.019872040
                              2.62656119304 ! Trappe 2
Control File
End
```

### NOTE:

Beware, the  $R_{min} \neq \sigma$ .  $R_{min}$  is the distance to the x-intercept (where interaction energy goes from being repulsive to positive.  $\sigma$  is the distance to the deepest part of the well, where the attraction is maximum. To  $\sigma\sigma$  convert to  $R_{min}$ , simply multiply  $R_{min}$  by 0.56123102415, and flag it with a negative sign.

Here's the finished Exotic file for isobutane (basically, empty, as we have no dihedrals, but with comments left in, to prepare for future reuse):

```
Parameter file for GO-MC
*
DIHEDRALS_POWER_SERIES
!V(dihedral) = Kchi * cos(chi - delta)^n
!Kchi: kcal/mole
!n: power
!delta: shift
!atom types
                   Kchi
                                delta
                                                    description
DIHEDRALS_QUAD_CIS_TRANS
!V(dihedral) = Kchi(1 + cos(n(chi) - delta))
! Kchi: kcal/mole
!n: 0 - cis; 1 - trans
!delta: shift
                                 delta
                                                     description
!atom types
                   Kchi
NONBONDED_MIE
!v(mie) = 4*eps*((sig_ij/r_ij)^n-(sig_ij/r_ij)^6)
!atom
        eps
                     siq_ij
                                n
                                     description
```

And here's the finished CHARMM format Parameter file for CHARMM isobutane.

```
* Parameter file for GO-MC
* Contains parameters for isobutane (Trappe 2 FF)
BONDS
!V(bond) = Kb(b - b0)**2
!Kb: kcal/mole/A**2
!b0: A
! Kb (kcal/mol) = Kb (K) * Boltz. const.; (9999999999 if no
stretching)
               Кb
                               b0
                                         description
!atom type
                               1.540
               999999999
CH3 CH1
                                       ! Trappe 2
ANGLES
!V(angle) = Ktheta(Theta - Theta0)**2
!V(Urey-Bradley) = Kub(S - S0)**2
!Ktheta: kcal/mole/rad**2
!Theta0: degrees
!Kub: kcal/mole/A**2 (Urey-Bradley)
!s0: A
! Ktheta (kcal/mol) = Ktheta (K) * Boltz. const.
                    Ktheta
!atom types
                                 Theta0
                                          Kub(?) SO(?)
CH3 CH1 CH3
                    62.100125
                                 112.00 ! Trappe 2
DIHEDRALS
!v(dihedral) = Kchi(1 + cos(n(chi) - delta))
!Kchi: kcal/mole
!n: multiplicity
!delta: degrees
! Kchi (kcal/mol) = Kchi (K) * Boltz. const.
!atom types
                    Kchi
                                     delta
                                                         description
                                n
NONBONDED nbxmod 5 atom cdiel fshift vatom vdistance vfswitch -
!cutnb 14.0 ctofnb 12.0 ctonnb 10.0 eps 1.0 e14fac 1.0 wmin 1.5
!V(Lennard-Jones) = Eps,i,j[(Rmin,i,j/ri,j)**12 - 2(Rmin,i,j/ri,j)**6]
```

```
!epsilon: kcal/mole, Eps,i,j = sqrt(eps,i * eps,j)
!Rmin/2: A, Rmin,i,j = Rmin/2,i + Rmin/2,j
!
 Rmin = sig * (2^{(1/6)}) / 2 ; eps (kcal/mol) = eps (K) * Boltz.
const.
                                 Boltzmann = 0.0019872041 kcal / (mol *
K)
                 epsilon
!atom
       ignored
                               Rmin/2
                                                     eps,1-4
                                                               Rmin/2,1-
                                            ignored
CH3
       0.0
                -0.194745992
                              2.10461634058 ! Trappe 1
                -0.019872040
                              2.62656119304 ! Trappe 2
CH1
       0.0
```

Note, the dihedrals section contains no values for this file, but was inserted none the less to demonstrate an empty section, similar to the unused sections of the exotic file. Alternatively empty tagged sections can simply be left off (as is done for **IMPROPERS** in this case).

### "Exotic" parameter file

The exotic file is intended for use with nonstandard/specialty models of molecular interaction which are not included in CHARMM standard. Currently a single custom interaction is included:

NONBONDED\_MIE n-6 potential. The Lennard-Jones potential (12-6) is a subset of this
potential. Manipulating the repulsive exponent gives you another means of tuning your
forcefield.

Otherwise the Exotic file reuses the same geometry section headings -- BONDS / ANGLES / DIHEDRALS / etc. The only difference in these sections versus in the CHARMM format forcefield file is that the energies are in Kelvin ('K'), the unit most commonly found for parameters in Monte Carlo chemical simulation literature. This precludes the need to convert to kcal/mol, the energy unit used in CHARMM.

The most frequently used section of the exotic files in the Mie potential section, **NONBONDED\_MIE**.

Here are the parameters that are used to simulate alkanes.

```
NONBONDED_MIE
!v(mie) = 4*eps*((sig_ij/r_ij)^n-(sig_ij/r_ij)^6)
                             sig_ij
3.740
3.783
3.990
!atom
                                                    description
                                                      Potoff, et al.
Potoff, et al.
Potoff, et al.
CH4
           161.00
                                             14
                                                                            '09
                                                                            '09
CH3
           121.25
                                             16
CH2
            61.00
                                                                            '09
                                             16
```

Note the exotic file uses  $\sigma$ , *not* the  $R_{min}$  used by CHARMM, although the units (Angstroms) are the same. The energy in the exotic file are expressed in Kelvin (K), as this is the standard convention in the literature.

### Control File (in.dat):

The control file is GOMC's proprietary input file. It contains key settings. The settings generally fall under three categories:

- Input/Simulation Setup
- System Settings for During Run
- Output Settings

### **Input/Simulation Setup**

In this section input file names are listed. In addition if you want to restart your simulation or use integer seed for running your simulation you need to modify this section according to your purpose.

InState: Determines whether to restart, and if so what step to restart from.

(WARNING: Restarts are not currently supported)

- Value 1: <BOOLEAN> true if restart, false otherwise
- Value 2: <ULONG> step to restart from

Example:

- PRNG: Dictates how to start the pseudo-random number generator (PRNG).
  - o Value 1: <STRING >
    - RANDOM: Randomizes Mersenne Twister PRNG with random bits based on the system time.

Example:

 RESTART: Used for restarting a previous simulation, this option loads in the Mersenne Twister's state from a saved file.

(WARNING: Restarts are not currently supported)

### Example:

INTSEED: This option "seeds" the Mersenne Twister PRNG with a standard integer. When the same integer is used, the generated PRNG stream should be the same every time, which is helpful in tracking down bugs.

### Example:

- o Value 2: ULONG or UINT: If "INTSEED" command is used (see above example).
- InParam: Provides the name of the parameter file to use for the simulation
  - Value 1: <STRING> Defines the name of the parameter file.

### Example:

- **FFKind:** Tells what format the above specified forcefield is using. If this tag is not specified the system assumes an exotic format is being used.
  - Value 1: <STRING> Defines the parameter file format

### Possible values:

- CHARMM
- Exotic (default option)

### Example:

- InPDB: Gives the filenames of the PDB files (coordinates) for each box in the system (one entry for NVT ensemble, two for GEMC/GCMC)
  - Value 1: <STRING> Defines pdb file name of first box.
  - Value 2 (GEMC and GCMC only): <STRING> Defines pdb file name of second box.
     (Note: This box is the reservoir in GCMC.)

### Example (NVT ensemble):

- InPSF: Gives the filenames of the PSF (topology) files for each box in the system (one entry for NVT ensemble, two for GEMC/GCMC)
  - Value 1: <STRING> Defines psf file name of first box.
  - Value 2 (GEMC and GCMC only): <STRING> Defines psf file name of second box.

### System Settings for During Run Setup

This section contains all the variables not involved with the output of data during the simulation or with the reading of input files at the start of the simulation. In other words, it contains settings related to the moves, the thermodynamic constants (based on choice of ensemble), and the length of the simulation.

Note that some tags or entries for tags are only used in certain ensembles (e.g. Gibbs ensemble). These cases are denoted with colored text.

GEMC: (FOR Gibb ensemble runs only) Defines what type of Gibbs Ensemble simulation you
want to run.

If neglected in Gibbs ensemble, it simply defaults to constant volume (NVT) Gibbs ensemble

- Value 1: <STRING> -- allows you to pick between isovolumetric ("NVT") and isobaric ("NPT") Gibbs ensemble simulations
  - NVT: Run simulation w/ constant mole number, volume and temperature.

### Example:

NPT: Run simulation w/ constant mole number, pressure and temperature.

Example:

- Value 2: <DOUBLE> -- Constant pressure in bars to run simulation at, if NPT ensemble is selected (see above example).
- System: This section defines the parameters that are required to calculate energy.
  - Value 1: <BOOLEAN> whether to use long range corrections (values: "true" or "false")
  - Value 2: <DOUBLE> constant temperature of simulation in Kelvin
  - Value 3: <DOUBLE> the distance to truncate the Lennard-Jones potential at Example:

### Note/Warning:

This isn't fully CHARM compliant. In CHARMM the 1-4 interaction scaling is supposed to be handled by having a separate 1-4 parameter in the nonbonded section. In the future the code's support for the NONBONDED\_MIE and NONBONDED sections of the forcefield will be updated to incorporate these parameters. The constant will then be transititioned to an electrostatics 1-4 scaling terms, along the lines of what NAMD currently does.

- **Steps**: Intervals for key step events
  - Value 1: <ULONG> total number of steps to run for (one move is performed for each step) (cycles = this value / number of molecules in the system)
  - Value 2: <ULONG> estimate of the number of steps necessary to equilibrate the system; averaging will begin at this step.
  - Value 3: <ULONG> the number of steps per adjustment to the maximum constants associated with each move (e.g. maximum distance in xyz to displace, the maximum volume in A<sup>3</sup> to swap, etc.)

### Example:

- **ChemPot** (FOR grand canonical (GC) ensemble runs only): Chemical potential to run the simulation at.
  - Value 1: <STRING> The resname to apply this chemical potential w.r.t.
  - Value 2: <DOUBLE>— The chemical potential value in degrees Kelvin (should be negative).

### Note:

For binary systems include multiple copies of the tag, one per residue kind.

### Example:

- Percent: Fractional percentage of each move (should add up to 1.0)
  - Value 1: <DOUBLE> % displace
  - Value 2: <DOUBLE> % rotate
  - Value 3: <DOUBLE> (FOR Gibbs ensemble runs only) % of volume swaps
  - o Value 4: <DOUBLE> (FOR Gibbs and GC ensemble runs only) % of molecule swaps

### Example:

- BOXDIM: Defines the axis lengths of simulation box. This tag may occur multiple times (It occurs once for NVT, but twice for Gibbs ensemble or GC ensemble).
  - Value 1: <INTEGER> sets box number (first box is box '0')
  - Value 2: <DOUBLE> x-axis length in Angstroms
  - Value 3: <DOUBLE> y-axis length in Angstroms
  - Value 4: <DOUBLE> z-axis length in Angstroms

### Example (NVT ensemble):

- CBMCNonbonded: (FOR Gibbs and GC ensemble runs only, currently) Number of CBMC trials to perform during certain stages of the regrowth
  - O Value 1: <INTEGER> Number of initial insertion sites to try
  - Value 2: <INTEGER> Number of trials for subsequent regrown positions in topology

### Example:

- **CBMCBonded:** (FOR Gibbs and GC ensemble runs only, currently) Number of CBMC trials to perform for geometry (per the coupled-decoupled CBMC scheme):
  - O Value 1: <INTEGER> Number of trials per angle
  - O Value 2: <INTEGER> Number of trials per dihedral.

### Example:

### **Output Controls**

This section contains all the values that control output in the control file. For example certain variables control the naming of files dumped of the block averaged thermodynamic variables of interest, the PDB files, etc.

- OutNameUnique: Unique name for simulation used to name the block output files.
  - o Value 1: <STRING> unique phrase to identify this system

### Example:

- OutState: controls output of PDB file (coordinates) and PRNG state.
  - Value 1: <BOOLEAN> –"true" enables dumping these files; "false" disables dumping.
  - Value 2: <ULONG> steps per dump event

### Example:

- OutPDB: Name of the output PDB, if PDB dumping is enabled (one files for NVT, two files for Gibbs ensemble or GC ensemble).
  - Value 1: <STRING> name of PDB file to output to for first box.
  - Value 2: <STRING> (FOR Gibb ensemble or GC ensemble runs only) name of output pdb file for second box

### Note/Warning:

The PDB file contains an entry for every ATOM, in all boxes read in. This allows VMD (which requires a constant number of atoms to properly parse frames (with a bit of help). Atoms that are not currently in a specific box are given the coordinate 0.00, 0.00, 0.00. The occupancy value corresponds to the box a molecule is currently in (e.g. 0.00 for box 0; 1.00 for box 1).

- OutPSF, OutSeed: Names of PSF and seed (PRNG state/frame) files (respectively).
  - Value 1: <STRING> File name

### Example:

### Note/Warning:

The PSF file contains an entry for every ATOM, in all boxes read in. It also contains all the topology for every molecule in both boxes, corresponding to the merged PDB format.

- OutConsole: Controls the output to STDIO ("the console") of messages such as acceptance statistics, averages, and run timing info.
  - Value 1: <BOOLEAN> "true" enable message printing; "false" disables dumping.

Value 2: <ULONG> – number of steps per print

### Example:

# enable, frequency

OutConsole true 10000

- OutBlockAverage, OutFluctuation, OutHistogram: Controls the
  output of block averages, fluctuation logs, and histograms (respectively). Block averages are
  averages of thermodynamic values of interest for chunks of the simulation (for post processing
  of averages or std. dev. In those values); fluctuations are a log of instantaneous values for those
  quantities; histograms are a binned listing of observation frequency for a specific
  thermodynamic variable. In this code they also control the output of a file containing
  energy/particle samples, commonly used in GCMC simulations for histogram reweighting
  purposes.
  - Value 1: <BOOLEAN> "true" enable message printing; "false" disables it.
  - o Value 2: <ULONG> number of steps per file output

### Example:

- OutHistSettings: Controls the output of the energy/particle sample file and the distribution file for particle counts, commonly referred to as the "histogram" output.
  - Value 1: <STRING> Short phrase to use in the name of the binned histogram for particle distribution.
  - Value 2: <STRING> Short phrase to use in the name of the energy/particle count sample file.
  - Value 3: <UINT> Run number to use in the above file names.
  - O Value 4: <CHAR> Run letter to use in above file names.
  - Value 5: <UINT> the number of steps per histogram sample.

### Example:

##############	########	########	########	########	##########
#					
#	hist.	sample	Run	Run	Steps Per
#	name	name	number	letter	Hist
Sample					
##############	#########	########	########	########	#########
#					
OutHistSettings	dis	his	1	a	200

 OutEnergy\*\*\*, OutPressure\*\*\*, OutMolNumber\*\*, OutDensity\*\*, OutVolume\*: Enables/disables for specific kinds of file output for tracked thermodynamic quantities

- Value 1: <BOOLEAN> "true" enable message output of block averages of this tracked parameter (and in some cases such as entry, components); "false" disables it.
- Value 2: <BOOLEAN> "true" enable message output of a fluctuation log for this tracked parameter (and in some cases such as entry, components); "false" disables it.
- Value 3: <BOOLEAN> "true" enable message output of a histogram of observation frequency for values of this tracked parameter (and in some cases such as entry, components); "false" disables it.

### Example:

###############	#########	########	######
# enable: blk av	vg., fluct	t., hist	
###############	##########	#######	#####
OutEnergy	true	true	true
OutPressure	true	true	true
OutMolNum	true	true	true
OutDensity	true	true	true
OutVolume	true	true	true

### Note/Warning:

Fluctuation logs and histograms are not currently supported. However, they are included here for future support. As a result, these tags can be ignored at present, but will be necessary for future functionality. Unlike the previous enable section, these values do require three entries per line.

Also, more tracked parameters, such as move acceptance, move constant values, chemical potential, sampled geometry, etc. may be added in the future.

Here's a sample of how a typical NVT ensemble control file might be expected to look:

```
#______
   GOMC beta 0.97 - NVT Control example
###
  INPUT
####################################
#GEMC type
      Pressure (bar)
GEMC
   NVT
####################################
# enable, step
true 1000000
InState
# kind {RESTART, RANDOM, INTSEED}
PRNG
   RANDOM
# charmm param, exotic param
Par_Vrabec_Argon.inp
# kind of forcefield
FFKind
    CHARMM
# pdb file
InPDB STEP3_START_AR_sys.pdb
# psf file
InPDB STEP3_START_AR_sys.psf
###
 SYSTEM
#Temp (K) #cutoff (A)
# SYSTEM VARS #LRC
true 0.85
System
             10
#tiil Equil
# STEPS PER...
     #total
                #Per adjustment
Steps
     1000000
           10000000
                1000
# MOVE %
        #disp. #rot.
1.00 0.00
Percent
```

```
# box #, x, y, z
BoxDim 0 40.00 40.00 40.00
# statistics filename add
OutNameUnique T_120_K
##############################
# enable, frequency
##############################
OutState true 1000000
##############################
# Box
###############################
OutPDB AR_T_120_K_Run_1.pdb
# (PSF, seed)
OutPSF
     AR_T_120_K_Run_1.psf
     AR_T_120_K_Run_1_seedOut.dat
OutSeed
####################################
OutConsole
       true
# enable, frequency
########################
# enable, frequency
############################
OutBlockAverage true
              100000
OutFluctuation
          false
              10000
OutHistogram
          false 100000
# enable: blk avg., fluct., hist.
OutEnergy
          true
               true
                    true
OutPressure
          true
               true
                    true
```

Here's a sample of how a typical Gibbs ensemble control file might be expected to look:

```
GOMC beta 0.97 - Gibbs ensemble Control example
INPUT
###################################
# enable, step
#####################################
InState
    true
       1000000
PRNG
   RANDOM
# charmm param, exotic param
Par_TraPPE_BranchedAlkanes.inp
InParam
# kind of forcefield
FFKind
    CHARMM
# pdbs for 1st and 2nd box
InPDB STEP3_START_ISB_sys_BOX_0.pdb STEP3_START_ISB_sys_BOX_1.pdb
# psfs for 1st and 2nd box
InPSF STEP3_START_ISB_sys_BOX_0.psf STEP3_START_ISB_sys_BOX_1.psf
###
  SYSTEM
#LRC
          #Temp (K) #cutoff (A)
# SYSTEM VARS
                     #1-4 Inter
Svstem
       true
          330.00
               10
                     0.0
# STEPS PER...
      #total
            #tiil Equil
                  #Per adjustment
Steps
      1000000
            10000000
                  1000
# MOVE %
         #disp. #rot. #vol.
                  #mol.
0.626 0.10 0.005 0.269
Percent
BoxDim 0 71.49 71.49 71.49
BoxDim 1
    71.49 71.49 71.49
```

```
CBMCNonbonded 10
              8
####################################
# arowth tr.
           ang
                dih
######################################
CBMCBonded
           100
                20
OUTPUT
# statistics filename add
OutNameUnique T_330.00_K
##########################
# PDB box 0, 1
###############################
OutPDB ISB_T_330.00_K_Run_1_MIE_BOX_0.pdb
ISB_T_330.00_K_Run_1_MIE_BOX_1.pdb
# (merged PSF, seed)
ISB_T_330.00_K_Run_1_MIE_merged.psf
ISB_T_330.00_K_Run_1_MIE_seedOut.dat
OutPSF
OutSeed
###################################
# enable, frequency
###################################
OutConsole true
              10000
##################################
# enable, frequency
##############################
                 100000
OutBlockAverage true
OutFluctuation
            false
                 10000
                 100000
OutHistogram
            false
# enable, frequency
hist.
                  sample
                        Run
                              Run
                                    Steps Per
                  name
                        number
                              letter
                                    Hist Sample
           name
OutHistSettings dis
                  his
                        1
                                    200
                              a
OutBlockAverage
           true
                 100000
# enable: blk avg., fluct., hist.
OutEnergy
            true
                  true
                       true
OutPressure
            true
                  true
                       true
OutMolNum
            true
                  true
                       true
OutDensity
            true
                  true
                       true
OutVolume
            true
                  true
                       true
```

#################################

first

# growth tr.

nth

Here's a sample of how a typical grand canonical ensemble control file might be expected to look:

```
GOMC beta 0.97 - Grand canonical ensemble Control example
 -----
======= ----- INPUT ------
###################
# enable, step
####################
        1000000
InState
     false
# kind {RESTART, RANDOM, INTSEED}
RANDOM
# charmm param, exotic param
FFKind
     Exotic
     Par_Vrabec01_Noble_Gases_Exotic.inp
InParam
# pdbs for 1st and 2nd box
InPDB STEP3_START_AR_vap_BOX_0.pdb
STEP3_START_AR_reservoir_BOX_1.pdb
# pdb x2 (new sys) psf x1 (restart)
InPSF STEP3_START_AR_vap_BOX_0.psf
STEP3_START_AR_reservoir_BOX_1.psf
##############################
# seed file (if restart)
##############################
#InSeed
     seedIn.dat
######
     ------ SYSTEM ------
# SYSTEM VARS #LRC #Temp (K) #cutoff (A) #1-4 Inter.
true 140.00
                     0.0
                10
#tiil Equil
# STEPS PER...
      #total
                   #Per adjustment
1000
      1000000
            500000
Steps
#disp. #rot. #mol.
# MOVE %
0.300 0.00 0.700
Percent
```

```
Mol. Name Chem. Pot.
ChemPot AR
                   -968
######################
# box #, x, y, z
###########################
BoxDim 0 25.00 25.00 25.00
BoxDim 1 25.00 25.00 25.00
###################################
# growth tr. first
###################################
CBMCNonbonded 10
# growth tr.
            ang
                  dih
#####################################
CBMCBonded 100 20
##############################
# statistics filename add
#############################
OutNameUnique T_140_K_u_968_r1a_v
##############################
# enable, frequency
##############################
OutState true
              100000
##############################
# Box 0, Box 1 PDBs
###############################
OutPDB AR_T_140_K_u_968_r1a_v_B0x_0.pdb
AR_T_140_K_u_968_r1a_r_BOX_1.pdb
##############################
# (merged PSF, seed)
###############################
OutPSF AR_T_140_K_u_968_r1a_merged.psf
OutSeed AR_T_140_K_u_968_r1a_seedOut.dat
#############################
# enable, frequency
############################
OutConsole true 100000
#############################
# enable, frequency
###########################
OutBlockAverage true
                   100000
OutFluctuation
             true
                   100000
OutHistogram
             true
                   100000
hist.
                    sample Run
                                  Run
                                         Steps Per
             name
                    name
                           number
                                  letter
                                         Hist Sample
OutHistSettings dis
                    his
                          1
                                         200
####################################
# enable: blk avg., fluct., hist.
```

### true OutEnergy true true OutPressure true true true OutMolNum true true true OutDensity true true true

## **GOMC's Output Files, Terminal Output**

GOMC currently supports several kinds of output:

- ❖ STDIO ("console") Output
- File Output
  - o PDB
  - o PSF
  - o Block Averages

### Console Output

A variety of useful information relating to variable averages, acceptance rates, file I/O messages warnings, and other kinds of information is printed formatted to the STDIO, which in Linux will typically be displayed in the terminal. This output can be redirected into a log file in Linux using the '>' operator

The first section of this console output typically includes some intro info relating to the forcefield read. This output is important as it may contain text relating to issues encountered if there was an error in the current run (e.g. a bad parameter, etc.

Next some info on the system's starting configuration will print:

For most of the run info on the variables and move statistics will be printed, e.g.:

```
STEP 9999 of 40000000 (0.0249975% done)
         === BOX 0 ===
Volume: 361095 A^3
Tot # mol: 946
Molecule Type ISB
Number: 946 ;
Density: 0.252856 g/ml
Energy (in K): total: -491491
inter: -713432 ; inter (tail corr.): -74880.3
intra (bonded): 296821 ; intra (nonbonded): 0
Displace (Box 0) -- tries: 3329; # Accept: 1366; % Accept : 41.0333; Max Amt.: 0.919088
Rotate (Box 0) -- tries: 420; # Accept: 241; % Accept : 57.381; Max Amt.: 3.12267
Molecule Transfer (Box 0 -> Box 1) -- tries: 1265; # Accept: 562; % Accept : 44.4269
Volume Transfer (Box 0 -> Box 1) -- tries: 82; # Accept: 54; % Accept : 65.8537; Max Amt.: 6834.38
        === BOX 1 ===
Volume: 369650 A^3
Tot # mol: 1054
Molecule Type ISB
Number: 1054 ;
Density: 0.275204 g/ml
Energy (in K): total: -619525
inter: -855439 ; inter (tail corr.): -90802.5
intra (bonded): 326716 ; intra (nonbonded): 0
Displace (Box 1) -- tries: 3226; # Accept: 1293; % Accept : 40.0806; Max Amt.: 0.879687
Rotate (Box 1) -- tries: 463; # Accept: 275; % Accept : 59.3952; Max Amt.: 3.14159
Molecule Transfer (Box 1 -> Box 0) -- tries: 1215; # Accept: 508; % Accept : 41.8107
Volume Transfer (Box 1 -> Box 0) -- tries: 0; # Accept: 0; % Accept : 0; Max Amt.: 300
         === SYSTEM ===
Energy (in K): total: -1.11102e+06
inter: -1.56887e+06 ; inter (tail corr.): -165683
intra (bonded): 623538 ; intra (nonbonded): 0
Steps/sec. : 178.955
```

### Note:

It's important to watch the acceptance rates and adjust the move percentages and CBMC trial amounts to get the desired rate of move acceptance.

At the end of the run timing information and other wrapup info will print, such as any failures encountered in the energetic consistency checks:

```
Simulation Time (total): 207330sec.
GOMC Serial Version 0.96
Completed at: Sat Sep 6 13:56:26 2014
On hostname:
```

### PDB and PSF Files

The PDB and PSF output (merging of atom entries) has already been mentioned/explained in previous sections. To recap: the PDB file's **ATOM** entries' occupancy is used to represent the box the molecule is in in the current frame. All molecules are listed in order they were read in. i.e. if box 0 has  $1..N_1$  molecules and box 1 has  $1..N_2$  molecules, then all of the molecules in box 0 are listed first, then all the molecules in box 1, i.e.  $1..N_1$ ,  $N_1+1..N_1+N_2$ . PDB frames are written as standard PDBs to consecutive file frames.

To visualize you'll need to open the PDB and PSF outputted files in VMD and use the command:

```
pbc join connected
```

In the TKConsole in order to get the current frame to obey the periodicity. Some peculiarities are still be observed, so please review the PDB file if anything looks odd (e.g. bonds will sometimes not show up in later frames). The GOMC team is in the process of improving VMD compatibility; but this may require changes to VMD to allow the files outputted to be properly read.

For Gibbs ensemble, the same PSF will be used to visualize either box, i.e.

```
vmd\ ISB\_T\_330.00\_K\_Run\_0\_MIE\_BOX\_0.pdb\ ISB\_T\_330.00\_K\_Run\_0\_MIE\_merged.pdb
```

...for visualizing the first box and...

```
vmd\ ISB\_T\_330.00\_K\_Run\_0\_MIE\_BOX\_1.pdb\ ISB\_T\_330.00\_K\_Run\_0\_MIE\_merged.pdb
```

... for visualizing the second box.

### **Block Output Files**

GOMC tracks a number of thermodynamics variables of interest during the simulation, some independent, others dependent on each other. The variables are tracked by this module:

- Density (g/ml)
- Energy
  - o Intermolecular
  - intramolecular bonded

- Intramolecular nonbonded
- tail corrections
- total interactions
- Virial coefficient
- Intermolecular
- tail corrections
- Pressure (bar)
- total

It is convenient to average the block file using simple shell utilities in Linux:

```
alias avg 'awk -v start_ln=\!:2 -v column=\!:3 '"'"BEGIN {n=0; sum=0; std_dev=\

0} NR>start_ln { sum+=$column; result[n]=$column; n++} END {
avg=sum/n; sum=0; \

for(x=1;x<=n;x++){ sum+=((result[x]-avg)^2)}; std_dev=sqrt(sum/n); 
print avg" "\

std_dev }'"'" \!:1'

alias avgd 'awk -v start_ln=\!:2 -v column=\!:3 '"'"BEGIN {n=0; 
sum=0; std_dev\

=0} NR>$start_ln { result[n]=$column*1000; sum+=result[n]; n++} END {
avg=sum/n\

; sum=0; for(x=1;x<=n;x++){ sum+=((result[x]-avg)^2)}; 
std_dev=sqrt(sum/n); pri\

nt avg" "std_dev }'"'" \!:1'</pre>
To call this, make sure its directory is in your path and type "./
```

On the command file the name of the file goes first.

```
avg Blk_Pressure_T_330.00_K_Run_0_MIE.dat 100 2
```

This should display the average, skipping the first 100 lines of the file. Adjust the # of skipped lines as appropriate based on when the system is found to be equilibrated, and adjust the fourth parameter for the desired box (with 1 being box 0, 2 being box 1).

# **Putting it all Together: Running a GOMC Simulation**

To recap the previous examples, a simulation of isobutane will be completed for a single temperature point on the saturated vapor liquid coexistence curve. Please refer to the attached reference directories if you get stuck.

Files for this simulation will be put in the empty folder "1-Isobutane-UserGenerated". If errors are encountered during the building of the system please refer to the folders "2-Isobutane-Reference/1\_ScriptsComplete/" and "1-Isobutane-Reference/2\_FullyBuilt/" for the proper files and results at various stages of the build.

The general plan for running the simulation is:

- 1. Build GOMC (if not done already)
- 2. Copy GOMC GEMC executable to build directory
- 3. Create scripts, PDB, and topology file to build the system, plus in.dat file and parameter files to prepare for runtime
- 4. Build finished PDBs and PSFs using the simulation.
- 5. Run the simulation in the terminal.
- 6. Analyze the output.

Please complete steps 1 and 2 and then traverse to the directory, which should now have a single file "GOMC\_Serial\_GEMC" in it.

Next, six files need to be made:

- PDB file for isobutane
- Topology file describing isobutane residue
- Two \*.inp packmol scripts to pack two system boxes
- Two TCL scripts to input into PSFGen to generate the final configuration

### isobutane.pdb:

REMARK	1 File	created	by	GaussView 5	.0.8	
ATOM	1 C1	ISB	1	0.911	-0.313	0.000
C						
ATOM	2 C2	ISB	1	1.424	-1.765	0.000
C						
ATOM	3 C3	ISB	1	-0.629	-0.313	0.000
C						
ATOM	4 C4	ISB	1	1.424	0.413	-1.257
C						
END						

```
Top_Branched_Alkane.inp:
* Custom top file -- branched alkanes
1
  1
MASS
       1 CH3
                 15.035 C
MASS
       3 CH1
                 13.019 C !
RESI ISB
                0.00 ! isobutane - Trappe
GROUP
ATOM C1 CH1
                0.00 !
                            C3
ATOM C2 CH3
                0.00 ! c2-c1
                0.00 !
ATOM C3 CH3
                            C4
                0.00 !
ATOM C4 CH3
BOND C1 C2 C1 C3 C1 C4
PATCHING FIRS NONE LAST NONE
END
pack_box_0.inp:
tolerance 3.0
filetype pdb
output STEP2_ISB_packed_BOX_0.pdb
structure isobutane.pdb
number 1000
inside box 0. 0. 0. 68.00 68.00 68.00
end structure
pack_box_1.inp:
tolerance 3.0
filetype pdb
output STEP2_ISB_packed_BOX_1.pdb
structure isobutane.pdb
number 1000
inside box 0. 0. 0. 68.00 68.00 68.00
end structure
```

```
build_box_0.inp:
psfgen << ENDMOL
topology ./Top_Branched_Alkane.inp
segment ISB {
    pdb ./STEP2_ISB_packed_BOX_0.pdb
    first none
    last none
}
coordpdb ./STEP2_ISB_packed_BOX_0.pdb ISB
writepsf ./STEP3_START_ISB_sys_BOX_0.psf
writepdb ./STEP3_START_ISB_sys_BOX_0.pdb
build box 1.inp:
psfgen << ENDMOL
topology ./Top_Branched_Alkane.inp
segment ISB {
    pdb ./STEP2_ISB_packed_BOX_1.pdb
    first none
    last none
}
coordpdb ./STEP2_ISB_packed_BOX_1.pdb ISB
writepsf ./STEP3_START_ISB_sys_BOX_1.psf
writepdb ./STEP3_START_ISB_sys_BOX_1.pdb
```

These files can be created with a standard Linux or Windows text editor. Please also copy a packmol executable into the working directory. The resulting folder can be compared against the "Expected\_Results/Build\_Scripts\_and\_Model" directory in the sample code folder.

Once those files are created, run:

```
./packmol < pack_box_0.inp
./packmol < pack_box_1.inp</pre>
```

This will create the intermediate PDBs. Please examine them for correctness (the results obtained can be compared against the "Expected Results/PDB Intermediates" directory in the sample code folder.

Then run the PSFGen scripts to finish the system

```
./build_box_0.inp
./build_box_1.inp
```

Note, you probably will have to first make your scripts executable using:

```
chmod a+x build_box_*.inp
```

This will create the intermediate PDBs. Please examine them for correctness (the results obtained can be compared against the "Expected\_Results/Complete\_PDB\_and\_PSF" directory in the sample code folder.

To run the code a few additional things will be needed:

- ❖ A GOMC Gibbs ensemble executable
- A control file
- Parameter files.

Enter the control file (in.dat) in the text editor

```
GOMC beta 0.94 - Gibbs ensemble Control example
INPUT
# enable. step
##################################
        1000000
InState
     true
INTSEED
PRNG
# charmm param, exotic param
Par_CHARMM_TraPPE_2_FF.inp
                  Par_Exotic_TraPPE_2_FF.inp
# pdbs for 1st and 2nd box
InPDB STEP3_START_ISB_sys_BOX_0.pdb STEP3_START_ISB_sys_BOX_1.pdb
# psfs for 1st and 2nd box
```

```
InPSF STEP3_START_ISB_sys_BOX_0.psf STEP3_START_ISB_sys_BOX_1.psf
SYSTEM
# SYSTEM VARS
       #LRC
           #Temp (K) #cutoff (A)
                      #1-4 Inter.
true
           330.00
                10
                      0.0
#tiil Equil #Per adjustment
# STEPS PER... #total
10000000
       1000000
                   1000
# MOVE %
         #disp. #rot. #vol. #mol.
0.626 0.10 0.005 0.269
Percent
BoxDim 0 71.49 71.49 71.49
BoxDim 1
     71.49 71.49 71.49
# growth tr. first nth
###############################
Growth
      53
        8
OUTPUT
# statistics filename add
OutNameUnique T_330.00_K
#############################
# PDB box 0, 1
###############################
OutPDB ISB_T_330.00_K_Run_1_MIE_BOX_0.pdb
ISB_T_330.00_K_Run_1_MIE_BOX_1.pdb
# (merged PSF, seed)
ISB_T_330.00_K_Run_1_MIE_merged.psf
    ISB T 330.00 K Run 1 MIE seedOut.dat
# enable, frequency
OutConsole true 10000
# enable, frequency
```

```
OutBlockAverage
                       100000
               true
# enable: blk avg., fluct., hist.
OutEnergy
                true
                        true
                               true
OutPressure
                true
                        true
                               true
OutMolNum
                true
                        true
                               true
OutDensity
                               true
                true
                        true
OutVolume
                        true
                               true
                true
Exotic parameters for isobutane:
* Parameter file for GO-MC
*
DIHEDRALS_POWER_SERIES
!V(dihedral) = Kchi * cos(chi - delta)^n
!Kchi: kcal/mole
!n: power
!delta: shift
!atom types
                 Kchi
                         n
                             delta
                                               description
DIHEDRALS_QUAD_CIS_TRANS
!V(dihedral) = Kchi(1 + cos(n(chi) - delta))
! Kchi: kcal/mole
!n: 0 - cis; 1 - trans
!delta: shift
!atom types
                 Kchi
                          n
                              delta
                                                description
NONBONDED_MIE
!v(mie) = 4*eps*((sig_ij/r_ij)^n-(sig_ij/r_ij)^6)
!atom
                                  description
       eps
                   sig_ij
                             n
```

CHARMM format Parameter file for CHARMM isobutene:

```
* Parameter file for GO-MC
```

```
* Contains parameters for isobutane (Trappe 2 FF)
BONDS
!V(bond) = Kb(b - b0)**2
!Kb: kcal/mole/A**2
!b0: A
! Kb (kcal/mol) = Kb (K) * Boltz. const.; (9999999999 if no
stretching)
               Кb
                               b0
                                         description
!atom type
CH3 CH1
               999999999
                               1.540
                                       ! Trappe 2
ANGLES
!V(angle) = Ktheta(Theta - Theta0)**2
!V(Urey-Bradley) = Kub(S - S0)**2
!Ktheta: kcal/mole/rad**2
!Theta0: degrees
!Kub: kcal/mole/A**2 (Urey-Bradley)
!s0: A
! Ktheta (kcal/mol) = Ktheta (K) * Boltz. const.
!atom types
                    Ktheta
                                 Theta0
                                          Kub(?) SO(?)
CH3 CH1 CH3
                    62.100125
                                 112.00 ! Trappe 2
DIHEDRALS
!V(dihedral) = Kchi(1 + cos(n(chi) - delta))
!Kchi: kcal/mole
!n: multiplicity
!delta: degrees
! Kchi (kcal/mol) = Kchi (K) * Boltz. const.
                                     delta
!atom types
                    Kchi
                                n
                                                         description
NONBONDED nbxmod 5 atom cdiel fshift vatom vdistance vfswitch -
!cutnb 14.0 ctofnb 12.0 ctonnb 10.0 eps 1.0 e14fac 1.0 wmin 1.5
!V(Lennard-Jones) = Eps,i,j[(Rmin,i,j/ri,j)**12 - 2(Rmin,i,j/ri,j)**6]
!epsilon: kcal/mole, Eps,i,j = sqrt(eps,i * eps,j)
!Rmin/2: A, Rmin,i,j = Rmin/2,i + Rmin/2,j
```

```
! Rmin = sig * (2^{(1/6)}) / 2 ; eps (kcal/mol) = eps (K) * Boltz.
const.
                                  Boltzmann = 0.0019872041 \text{ kcal / (mol *)}
K)
!atom
       ignored
                  epsilon
                                Rmin/2
                                              ignored
                                                                  Rmin/2,1-
                                                        eps,1-4
CH3
       0.0
                 -0.194745992
                                2.10461634058 ! Trappe 1
CH1
       0.0
                 -0.019872040
                                2.62656119304 ! Trappe
```

Once these four files have been added to the output directory the simulation should be ready to run. The files and directory can be compared to "Expected\_Results/Ready\_To\_Run" directory in the sample code folder.

Assuming the code is named GOMC\_Serial\_GEMC run using:

```
./GOMC_Serial_GEMC > out_ISB_T_330.00_K_RUN_0.log &
```

Progress can be monitored with the tail command:

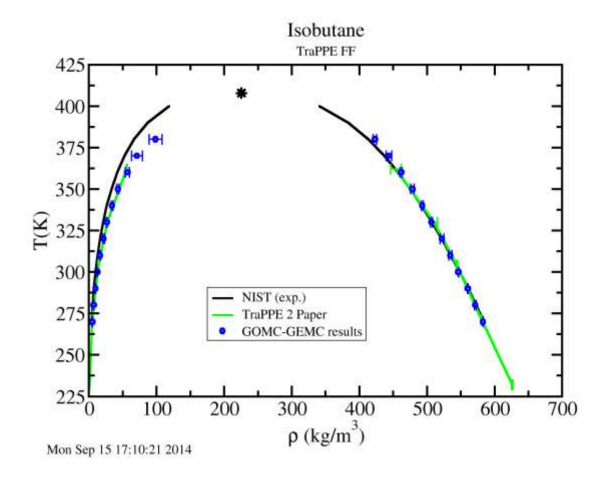
```
tail -f out_ISB*log
```

When the run is done, please compare the results to those found in

"Expected\_Results/Simulation\_Output" directory in the sample code folder. Use the "avg" alias, added in the previous section to determine the equilibrated block averages.

Congratulations, a single phase coexistence point on the saturated vapor liquid curve has been examined using GOMC operating in the Gibbs ensemble.

This process can be repeated additional times to produce new results (saving the old results aside) to produce averages. It is recommended that at least three simulations be run per point, although for larger/longer simulations sometimes it is appropriate to run only a single simulation.



Repeating this process for multiple temperatures will allow you to obtain the following results. This data is contained in the folder "Expected\_Results/TraPPE\_Isobutane\_VLE\_Graphs".

# NVT Ensemble Gibbs Ensemble Configurational Bias

**Summary of Algorithms in the Code** 

# **How to Get Help/Technical Support**

**GPU Algorithms/Optimizations** 

Please send an email to gomc@eng.wayne.edu