User Manual for Monte Carlo Code

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1 Support

Please send questions, bugs, and requests to Tristan Truttmann at tristan.truttmann@gmail.com.

2 Purpose

The MC code was developed as a tool for modeling the physical interaction between energetic compounds and organic adsorbents; however, interaction between any adsorbate and adsorbent can be modeled. The user is required to generate an initial geometry for the system, and then provide appropriate information to the SIESTA quantum chemistry package, as well as the appropriate pseudopotential files. The user is also required to provide simulation information into a preferences file such as the temperature and the size (in atoms) of the ad-molecule. The program will output the geometries and energies of every accepted structure.

3 Dependencies

The code was written for Python 2.6/2.7 and assumes that the standard Python libraries are installed as well as the numpy and matplotlib python libraries. It also assumes that it is running in an environment that has SIESTA 4.0 installed that is callable with the siesta command.

4 Suggested use

The code was designed to be lightweight so that the user can copy the entire software directory to a desired parent directory, and then make the necessary changes to the Siestafiles/ directory and the MC_Pref.py file. The simulation can then be run by either running RunMC.py directly (i.e. ./RunMC.py), or by submitting RunMC.py to a job manager (i.e. qsub RunMC.py). It should be noted that, in order for the program to execute properly, the current working directory at the time of submission must be the directory that contains the RunMC.py file. It should also be noted that the first couple lines of RunMC.py contain PBS directives that should be modified to the user's liking (if she is using a job manager). Job managers that are not based on PBS are not supported, but if the user wants to, she may replace the PBS directories with those of her desired job manager. However, if she does this, she should ensure that the TMPDIR environment variable is set to a scratch path; this prevents excessive writing overhead to the user's home directory (the program will write temporary files to the directory that the job was submitted in if TMPDIR is not defined). The program will write all output files to the output/ directory. If the user runs the

simulation when there are already files in the output directory, the program will move all those files to a directory named archive_output### were ### is an index number. This allows the user to resubmit simulations but keep track of previous program settings and results.

5 Getting Started

To ensure that your machine has the necessary prerequisites, submit the default job as a test using the following instructions. The default job simulates TNT interacting with α -cellulose.

- Copy the software directory to any desired location on your machine. (e.g. cp monte-calo-master /jobs/)
- Move to the cirectory you just created. (e.g. cd /jobs/monte-carlo-master)
- Submit the job if you have a batch manager (i.e. qsub RunMC) or run the script directly (i.e. ./RunMC.py).
- After the job is done running, move to the output directory. Look at the output files. (e.g. cd output && ls)
- Look through your results. You can see the geometries by opening GeomEnsemble.xyz in some molecular viewing program such as Avogadro.

6 Function

This sections explains, intuitively, how the program works. The program aims to model the interaction between a *single* ad-molecule and an adsorbent surface. Further, it does this in a vacuum.

To estimate the binding energy and properly account for thermal energy in the system, a Metropolis Monte Carlo technique is implemented. This involves a random translation and rotation of the ad-molecule followed and preceded by a potential energy calculation using SIESTA. The random move is then accepted or rejected based on the standard Metropolis Monte Carlo acceptance criteria; this is:

$$W = \min\left[1, \exp\left(\frac{-\Delta E}{kT}\right)\right]$$

Here, W is the probability of accepting the move, ΔE is the change in potential energy during the move, k is the Boltzmann constant, and T is the temperature in kelvin. If the move is rejected, the geometry prior to the move is assumed. If the move is accepted, the new geometry is recorded, and another random move starting from the new geometry is taken.

The choice of random moves is important for program performance. Generally speaking, a move that has an acceptance rate of 50% is ideal. For this program, each move has both a translation component, and a rotation component. During the translation component, a random direction, uniformly distributed over the surface of a sphere, is chosen. Then a random translation distance is chosen from a flat distribution between zero and the value of TransLimit. For the rotation component, an axis of rotation is chosen randomly from a uniform distribution over the surface of a sphere. Then a random rotation angle is chosen from a uniform distribution between zero and the value of RhoLimitDeg. The initial values of TransLimit and RhoLimitDeg are imported from

the MC_Pref.py file. If the file does not define these variables, then defaults are imported from MC_Defaults.py. If the user chooses Optimize = False, then these values are kept constant over the simulation. However, if the user chooses Optimize = True, then the these two variables are changed dynamically during the program to reach an acceptance rate near 50%. To do this, every accepted move is followed by increasing the values of both TransLimit and RhoLimitDeg by 5%, and every rejected move is followed by a decrease in these variables by 5%. The program then continues this process until there are EnsembleTartget accepted structures.

7 Files

This section explains the purpose of each file in the MC software. In the software directory, there are, by default, four files and two subdirectories. During operation, there may be an additional three files and several additional directories. These files and directories are summarized in Table 1. The files inside the subdirectories are summarized in Table 2 and Table 3.

Table 1: Summary of the files and subdirectories in the software directory.

$ t MC_Defaults.py$	Contains default preferences that are not defined in the MC_Pref.py file.	
$\texttt{MC_Defaults.pyc}$	A file that python may generate automatically.	
MC_Pref.py	A python file that the user must edit to change preferences of the simulation.	
MC_Pref.pyc	A file that python may generate automatically.	
$\texttt{MC_Util.py}$	The file that contains utilities for the RunMC.py file.	
$\texttt{MC_Util.pyc}$	A file that python may generate automatically.	
RunMC.py	The file that contains the core of the program as well as job management	
	information. The program is called with this script (i.e/RunMC.py)	
SiestaFiles/	A folder that contains necessary information for SIESTA to run.	
archive_output###/	"###" can be any number. These directories are generated to store output	
	files and preferences from previous runs.	
output/	All of the output output files are stored in this directory.	

Table 2: Summary of the files in the SiestaFiles subdirectory.

InputGeom.xyz	A text file in xyz format that stores the initial geometry of the system being
	studied. The user must replace this with her system.
template.fdf	The input file for SIESTA, excluding the geometry block at the end (which
	the program will append automatically). The user must edit this file to her
	specific needs.
.psf	"" are chemical species labels defined in the ChemicalSpeciesLabel block
	of the Siesta Fiels/template.fdf file. The pseudopotential files for SIESTA.
	The users must either generate these files or download them from the internet.

Table 3: Summary of the files in the output subdirectory.

ConfigEnsemble.py	A Python-readable file that stores all the information about every accepted
	MC structure in the system. This includes energy, geometry, lattice informa-
	tion, and the bias from metadynamics or umbrella sampling.
EnergyEnsemble.py	A Python-readable file that stores the energy values at every step in a numpy
	array.
GeomEnsemble.xyz	An xyz file that stores the geometries of every accepted structure.
MC_Pref[archive].py/	A copy of the MC_Pref.py file that was used for the job.
SiestaFiles[archive]/	A copy of the SiestaFiles/ directory that was used for the job.
time log	A file that records important performance information such as parallel
time.log	speedup, acceptance ratio, and time spent in different parts of the program.
output pha	Only exists if submitted as batch jobs. A file that contains the standard output
output.pbs	of the entire job.
aman nha	Only exists if submitted as batch jobs. A file that contains the standard error
error.pbs	out of the entire job.

8 Changing simulation preferences

To submit a custom job, the user is required to do four things:

- Supply an initial geometry to replace the SiestaFiles/InputGeom.xyz file.
- Edit the SiestaFiles/template.fdf file to her needs.
- Supply necessary pseudopotential files as *.psf where "*" are the symbols of each element.
- Edit simulation preferences in the MC_Pref.py file.
- If the user is submitting a batch job, edit the PBS directives at the beginning of RunMC.py. Then run qsub RunMC.py.
- If the user is submitting a regular script, run the command ./RunMC.py.

Further details are provided in the following subsections:

8.1 Edit SiestaFiles/template.fdf

The only difference between the SiestaFiles/template.fdf file and a traditional SIESTA input file is that the former is missing the geometry block. Otherwise, all SIESTA options can be used except options that are not supported by this MC program such as:

- Molecular dunamics are note supported.
- TransSiesta calculations are not supported.
- Minimizations are currently not supported but may be in the future (see Section 9).

• Only the Harris functional is supported, but more functionals will be supported in the future (See Section 10).

Detailed information about the SEISTA input file syntax and options can be found at **their website**. Remember to change the number of atoms and the lattice information.

8.2 Pseudopotential files

Every element in your simulation must have a pseudopotential file. For common functionals, pseudopotentials can often be found in **pseudopotential databases**. For more "exotic" functionals like vdW-DFT, you may have to generate your own pseudopotentials using **the built-in psuedopotential generate** or a third party pseudopotential generator like **OPIUM**.

8.3 Edit MC_Pref.py

The MC_Pref.py file is in Python readable format, so any tricks or shortcuts you know in Python can be used to generate these variables. However, if you spell a variable incorrectly, the program will not read the option and use the program defaults. Therefore, it is very important that you spell the option variables correctly. For every system, the following required options must be changed:

- AdsorbateSize: This variable should be set to the number of atoms in your adsorbate
- SubLatticeGrid: If periodic boundary conditions are specified in the SiestaFiles/template.fdf file and ShepherdOn = True, then this variable should be defined. This is necessary to keep your ad-molecule confined to the smallest possible repeating unit of the adsorbent surface. So if your adsorbent used in you system includes 2 repeating unites in the first vector specified in SiestaFiles/template.fdf, and 3 repeating unites included in the second vector, then you should define SubLatticeGrid = np.array((2,3,1)). Of course, if this confuses you, you can always remove the variable, and the defaults will just limit the system to the unit cell defined in SiestaFiles/template.fdf.

The following options are more ways to tune the calculation to your needs:

- T_Sequence: This variable represents the temperature of the system, or the simulated annealing sequence of temperatures. If you want to run the calculation at a constant temperature, for example 298 K, use T_Sequence = [[1,298]]. However, if you want to run a sequence, use a nested list, where each element is a 2-list where the first value is the number of moves attempted at the temperature specified in the second subelement. The sequence will repeat itself after it is finished. For example, if T_Sequence = [[1000,700],[1000,298]], then the simulation will make 1000 attempts at 700 K, and then 1000 attempts at 298 K, and then repeat the process until the program is complete.
- EnsembleTarget: This specifies the number of accepted structures that must be reached before the program terminates.
- Optimize: This logical variable, when set to True, will optimize the MC step size to approach an acceptance rate of approximately 50%.

- TransLimit: This variable is the maximum translation, in Angstrom, that the ad-molecule will be moved. If Optimize = True, then this value will be automatically adjusted during the program.
- RhoLimitDeg: This variable is the maximum rotation, in degrees, that the ad-molecules will be rotated during a MC step. If Optimize = True, then this value will be automatically adjusted during the program.
- ShepherdOn: This logical variable, if set to True, will prevent the ad-molecule from walking out of the unit cell specified by the lattice info specified in SiestaFiles/template.fdf and SubLatticeGrid.
- NumberOfSiestaCores = This variable specifies the total number of CPU cores to run SIESTA on

9 Coming features

During the fall of 2016, I will be working to implement the following features as an undergraduate honors project. I hope to have all improvements implemented by December 2016:

- Metadynamics and umbrella sampling: The adsorption of energetics onto organic adsorbents is characterized by a potential energy hypersurface with numerous minima separated by high activation barriers. This makes the traditional Metropolis Monte Carlo algorithm, as implemented, insufficient to adequately sample phase space. To overcome this shortcoming, I hope to implement metadynamics and umbrella sampling.
- Covalent interaction relaxation: The current program models the ad-molecule and adsorbent as rigid independent bodies. In coming versions, I will supply the option to relax the geometry of each of these independent bodies at every MC step before calculating the energy used in the acceptance criteria.
- Automatic Ad-molecule placement: I hope to add the option of supplying the admolecule and adsorbent geometries in seperate xyz files, and then have the program automatically place the ad-molecule in the proper place. This will be useful for high-throughput screenings and such.
- Basis set superposition error: Noncovalent binding energy calculations are known to have a large error due to basis set superposition error. I hope to provide an option to correct for this in the future.
- Improved parallelism: The current version supports all parallelism that is supported in SIESTA. However, I may add additional embarrassing parallelism to run on tens to hundreds of nodes if necessary.

If you have requests for additional features, or are waiting for one of the features listed above, please email Tristan Truttmann at tristan.truttmann@gmail.com, so I can be honest with whether I can implement the requested features and on what time frame.

10 Bugs

The following bugs will be worked out in future versions:

• Only the Harris functional is supported so far. This is a top priority to fix. If you really want to fix this, change the search string "siesta: Eharris(eV)" near line 609 in MC_Util.py to the appropriate search string for whatever functional you are using. You have to experiment with SIESTA to find what this keyword is.

11 License

This code is under the MIT License, which allows free distribution, modification, commercial use, and private use of this code as long as the authors are attributed and are not held liable.

12 Acknowledgements

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