# AMOEBA advanced potential energies workshop

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# Why Tinker?

Tinker is a freely available molecular modelling package, primarily authored by Jay Ponder's lab at the University of Washington in St Louis. It has utilities for all sorts of biomolecular simulations, and includes parameters for many of the standard biomolecular force fields, but for learning how to use AMOEBA simulations it has 3 main advantages:

- 1. It's the canonical AMOEBA code other software packages may be faster (more on that later!) but their implementations of AMOEBA will follow that of Tinker
- As the canonical code, Tinker includes the latest AMOEBA parameter and methodology developments, for testing and investigating the effects of polarisation in different systems
   It's free!

In this first tutorial we'll introduce the basic Tinker utilities for setting up and running a molecular dynamics simulation with AMOEBA, along with the file formats Tinker uses for topology, coordinates, and parameters of simulated systems. We'll be using a simple Ala<sub>3</sub> peptide in water as our system of interest, but the concepts are similar for any biomolecule.



# Setting up Ala<sub>3</sub>

# Logging in

Log into Lyceum2, and type 1s. You should see a folder called 'amoebaworkshop'. Change into it:

### cd amoebaworkshop

Inside there will be many more folders, each corresponding to different exercises (you can see these if you type 1s again). For now, change into the 'Exercise\_1' folder, and list the files inside:

```
cd Exercise_1
ls
```

Of course, the first thing we need to set up a molecular dynamics simulation is an initial structure of the system we want to simulate. This might be, for example, an X-ray crystal structure of a protein from the PDB, properly protonated and solvated. In this case we've provided an initial file, AAA\_solv.pdb. This is a linear Ala<sub>3</sub> peptide, which for the sake of speed we've already created and solvated in a box with a 10.0 Å buffer of water on all sides. There are many software packages that can perform this sort of set up, and of course Tinker has its own utilities too (see side note 1).

# File formats

Tinker makes use of two main file formats to run its simulations:

1. The 'xyz'

### Side note 1 - Tinker protein set up

Tinker has its own utilities for creating linear peptide and nucleic acid chains using the protein and nucleic tools, which you can run interactively to build initial structures in Tinker format. These can then be solvated with the xyzedit program (for details see exercise 2). However, for the most part, you might be interested in using initial structures (e.g. crystal structures) that already exist - in which case pdbxyz is the way to go. Pdbxyz is aware of all standard amino acids, nucleic acids, water and mono/divalent ion names and will convert and assign atom types to Tinker format, adding hydrogen atoms to crystal structures where necessary. Problems with pdbxyz are most often caused by discrepancies between atom names in different programs - take care with this in your own simulations.

file: Contains the coordinates and topology of all the atoms in the system

2. **The 'key' file:** Contains the simulation conditions and input options, along with parameters for atoms, or paths to find them

We'll start by making a basic key file. Open up a new file called PDB\_to\_xyz.key in your favourite text editor - I use vi (e.g. type vi PDB\_to\_xyz.key). Add the following into the empty text file, and save it:

```
parameters
/home/***Enter_your_username_here***
/tinker/params/amoebapro13.prm
a-axis 36.281
b-axis 33.399
c-axis 30.222
```

The first line (be sure to enter the temporary Southampton username you've been given correctly...) gives Tinker the location of the parameter file we want to use for this simulation. Tinker is packaged with a variety of force fields, here we'll use the Amoeba protein parameters of 2013 (named amoebapro13.prm in the Tinker 'params' directory).

The next three lines (starting a-axis, b-axis & c-axis) are the X, Y & Z box dimensions in Ångstrom of the periodic system. We've taken these straight from the CRYST records at the top of the PDB file.

# **Converting a PDB to Tinker format**

Now we have a basic key file we can use it to create a Tinker xyz file from our initial PDB file. Tinker has a built-in program to do this, called pdbxyz. Back in the terminal window, type:

```
pdbxyz AAA_solv.pdb -k PDB_to_xyz.key
ls
```

You'll see that two new files have been created, AAA\_solv.seq and AAA\_solv.xyz. Have a look at the .seq file first, again in your favourite text editor. You'll see it just contains the line "1 ALA ALA ALA". This is the peptide sequence that pdbxyz has interpreted and converted. Next, open up AAA\_solv.xyz. It'll have lines that look like this:



As you can see, the xyz file contains the coordinates, topology, and atom type information Tinker needs to describe the system under investigation. We can now use this file to actually start some simulation...

# **Energy minimisation**

# Defining simulation conditions in a key file

The first step in any MD protocol is to relax the initial structure to remove any steric clashes or poor configurations arising from our setup/solvation procedure. Tinker includes multiple different minimisation algorithms, but here we'll use the program minimize with the rapid but simple steepest descent algorithm.

Start by making a new copy of both our xyz file and key file, for tidiness' sake:

cp PDB\_to\_xyz.key AAA\_min.key
cp AAA\_solv.xyz AAA\_min.xyz

Then, edit AAA\_min.key so it contains the following (the comments, starting with #, aren't necessary):

```
parameters /home/***Enter_your_username_here***
/tinker/params/amoebapro13.prm
a-axis 36.281
b-axis 33.399
c-axis 30.222
                            # No. of parallel CPU
openmp-threads 4
threads to use
cutoff 9.0
                            # Nonbonded interactions
direct cutoff
ewald
                            # Switch on PME
ewald-cutoff 7.0
                            # PME real space cutoff
(overrides the 9A above)
                            # Switch on analytical
vdw-correction
long-range vdW correction
polar-eps 0.01
                            # Dipole convergence
criterion in RMS Debye/atm
neighbor-list
                            # Use pairwise neighbor
list for calculating
                            # nonbonded interactions
(improves speed)
                            # Maximum number of
maxiter 2000
minimisation steps
steepest-descent
                            # Use the SD
minimisation algorithm
```

### printout 100 print out energies

We've added some keywords defining the type of simulation we want to perform - a steepest descent minimisation of maximum 2000 steps, with a 9.0 Å vdW cutoff, 7.0 Å real space electrostatics cutoff, and PME and an analytical correction for long range electrostatics and vdW respectively. Additionally we've specified (with the polar-eps keyword) that at every step we wish our induced dipoles to be iterated until the RMS change in atomic dipoles is less than 0.01 Debye/atom. This is a fairly *loose* convergence criterion, and is at the limits of conserving energy in an AMOEBA MD simulation, but we use it here for speed. In practice it's preferred to use a convergence criterion of 1e-5 Debye/atom or below.

# Running a minimisation job

To run this minimisation we'll need to submit a job to the Lyceum cluster. A script to do so has been ready prepared for you, called AAA\_min.x. Open it up, and you'll see the command line used to run a Tinker minimisation, along with some comments describing how it's used:

```
# Command for using Tinker minimize:
# minimize [xyz file] -k [key file] [gradient
convergence criterion]
minimize AAA_min.xyz -k AAA_min.key 0.01 >
AAA_min.out
```

The gradient convergence criterion here is NOT the same as the dipole convergence criterion above. It is the RMS gradient in the energy at which we wish to stop the minimisation. If this gradient is reached before the maximum number of minimisation steps have been performed, the minimisation will stop, as the system is well minimised anyway. 0.01 kcal/mol/A per atom is the default value, and we use it here.

Change any filenames in AAA\_min.x to be correct for the xyz file and key file you have, then submit the job to the cluster:

qsub AAA\_min.x

This will take about 3 minutes to run. You can have a look at the output in real time by following AAA\_min.out, e.g. tail -f AAA\_min.out

Once complete, it's time to run some molecular dynamics...

# Heating to 300K - NVT simulations

# Additions to the key file for dynamics

Once your minimisation job is complete, you'll have a new file in your directory: AAA\_min.xyz\_2. This is the final structure of the solvated Ala<sub>3</sub> system after the minimisation. By default, Tinker appends output files with sequential numbers, so if you wanted to start another minimisation from AAA\_min.xyz\_2, the output would be called AAA\_min.xyz\_3. However, this might get very confusing very quickly, so instead we'll rename our output file to something more sensible, and make a new key file for the next simulation:

#### cp AAA\_min.key AAA\_eq\_nvt.key

As you might have guessed our next simulation will involve heating and equilibrating the system at 300 K, under constant volume (NVT) conditions. Open up the AAA\_eq\_nvt.key file and edit it to look like the following. There are only a few changes that need to be made at the bottom of the file:

parameters /home/***Enter_your_username_here*** /tinker/params/amoebapro13.prm				
a-axis 36.281 b-axis 33.399 c-axis 30.222				
openmp-threads 8				
cutoff 9.0 ewald ewald-cutoff 7.0 vdw-correction polar-eps 0.01				
neighbor-list				
printout 500				
thermostat andersen	# Switch on Andersen			
integrator verlet integrator	<pre># Use a velocity-Verlet</pre>			
archive	<pre># Create a single</pre>			
concatenated in sequence	# MD snapshots			

Notice we've removed the 'maxiter' and 'steepest-descent' keywords as we're no longer performing a minimisation. Instead they've been replaced with keywords to set a thermostat and integrator. Finally the 'archive' keyword will concatenate all of our output files (snapshots of the simulation coordinates) into a single trajectory. Otherwise Tinker prints out every frame as a separate, sequentially numbered file, as it did with the minimisation output above.

### How to use 'dynamic', the MD engine

Next, we'll submit an MD job to the Lyceum cluster - the script to do so is called AAA\_eq\_nvt.x. It contains the following command line, which uses the program dynamic, Tinker's main MD engine:

dynamic	Side note 2 - interactive running		
	All Tinker programs, minimize and dynamic included, are also set up to run and receive input interactively. Simply running 'dynamic' from the command line will prompt the user for all required input data, from input filenames to choice of ensemble to temperatures and pressures. This will also happen if only a partial command line is used, e.g. if you want to run an NVT simulation but forget to specify a desired temperature.		
AAA_eq_nvt.xy AAA_eq_nvt.ke 300 > AAA_eq_	z -k y 5000 1.0 0.5 2 nvt.out		

The numbers following the usual xyz and key file inputs correspond to

the following:

- 1. The number of MD steps to perform (here 5000 steps)
- 2. The timestep, in fs (here 1.0 fs)
- 3. The time interval to print out coordinate information, in ps (here 0.5 ps, or 500 steps)
- The type of simulation to run/which ensemble to run in (here 2 = NVT)
- 5. The desired simulation temperature, in K (here 300 K)

Change any of the filenames in AAA\_eq\_nvt.x to be correct for your own system, and submit the job to Lyceum:

qsub AAA\_eq\_nvt.x

This should take about 6-7 minutes to run.

# Equilibrating to 1 atm - NPT simulations

### **Restarting a simulation**

Open up the output file (AAA\_eq\_nvt.out) from the heating simulation. The last few lines should include a section like this:

Average Values for the Dynamics Steps	Last 500 C	ut of 5000
Simulation Time Total Energy (+/- 5 4696)	5.0000 4860.8933-	Picosecond Kcal/mole
Potential Energy	-6761.2395	Kcal/mole
(+/- 31.1456) Kinetic Energy	1900.3462	Kcal/mole
(+/- 28.3130) Temperature	275.99	Kelvin
(+/- 4.11) Pressure	-770.24	Atmosphere
(+/- 533.81) Density (+/- 0.0000)	0.6305	Grams/cc

This file contains the average and instantaneous system energies, temperatures, pressures etc. If this were a real simulation you may wish to visually check equilibration by writing a small script to extract and plot the progress of these properties over time. From the above we can already see this NVT simulation is too short, as the system temperature hasn't quite reached 300 K. We could change the collision frequency of the Andersen thermostat to speed this up (using the 'collision' keyword - see the Tinker manual), but for now we'll carry on with a pressure equilibration just to demonstrate how it works.

Type 1s and you'll see a couple more new files have been created as outputs from the simulation. The first, AAA\_eq\_nvt.arc, is the 'archive' file, containing all the coordinates of snapshots printed out during the trajectory as a single file (remember, this is why we specified the 'archive' keyword in AAA\_eq\_nvt.key). The second, AAA\_eq\_nvt.dyn, is the 'dynamics' file and contains the current atomic positions, velocities, box dimensions and accelerations at the last trajectory snapshot. The dyn file is therefore the way to perform a (non-binary) restart of a Tinker MD simulation:

If dynamic finds a dyn file with the <u>same prefix</u> as the specified starting xyz file, then coordinates, velocities and box dimensions

#### are <u>restarted</u> from the dyn, <u>overriding anything specified in other</u> <u>inputs</u>

So, in theory, we could use exactly the same command line for dynamic as we did above and the simulation would continue for another 5 ps from where it left off, as Tinker would recognise the existence of the dyn file. However, for tidiness' sake we will again make copies of our inputs and start our NPT equilibration from there.

### **Running under constant pressure**

Start by making copies of the key file and dyn file:

cp AAA\_eq\_nvt.key AAA\_eq2\_npt.key
cp AAA\_eq\_nvt.dyn AAA\_eq2\_npt.dyn

And now add the following to the bottom of the AAA\_eq2\_npt.key key file:

```
barostat berendsen
barostat
tau-pressure 1.0
time to 1.0 ps
```

# Switch on Berendsen
# Set pressure coupling

There are other, rigorous, barostats (e.g. Monte-Carlo, Nose-Hoover) available in Tinker, but for simplicity we'll use the Berendsen one here.

Next we'll extract the last frame of the NVT trajectory to use as the input of the NPT simulation. Again, thanks to the dyn file we don't *need* to do this, but it is sensible for record-keeping and illustrates how to manipulate trajectories. The Tinker archive program is used to do this, and we'll use it interactively as suggested in Side Note 2. Type:

archive AAA\_eq\_nvt.arc

...and follow the on-screen instructions. Here we want option 2, to extract an individual frame. You'll then be asked to select the first and last frame to extract, and the stride between them. Here we want the 10th frame only (the last in our 5 ps trajectory), so type 10 10 1 at the prompt. Once complete, press Enter to quit. Type 1s again and you'll notice a new file has been created, AAA\_eq\_nvt.010. This is the 10th frame of the AAA\_eq\_nvt.arc archive file - the one we want. Rename it as the input for our upcoming NPT simulation, **making sure it has the exact same prefix as the dyn file we want to restart from**:

mv AAA\_eq\_nvt.010 AAA\_eq2\_npt.xyz

Finally we're ready to run. The script to submit is AAA\_eq2\_npt.x, inside you'll find the command line:

dynamic AAA\_eq2\_npt.xyz -k AAA\_eq2\_npt.key 5000 1.0 0.5 4 300 1.0 > AAA\_eq2\_npt.out

Notice there are two changes from the NVT simulation. First we've changed from ensemble option 2 (NVT) to ensemble option 4 (NPT). Second, there is a final number on the command line specifying the pressure we want to run at, in atmospheres (here 1.0 atm). Make any filename changes you need to and submit the job:

#### qsub AAA\_eq2\_npt.x

This should again take about 6-7 minutes to run. Once complete, have a look at the outputs again - you should see the box dimensions and density begin to change as the barostat does its job.

# Extras:

# **Visualising output**

Obviously with these short simulation times our system is nowhere near equilibrated just yet. However, we can still take a look at our trajectories to ensure there are no clashes or unphysical structures created during the dynamics so far. VMD is often used for the visualisation of molecular dynamics simulations, and we'll use it here too.

However, VMD currently (version 1.9.2) has a small bug, in that it will only correctly visualise Tinker archive files that **don't contain any periodic box information**, as older versions of Tinker did not include box dimensions in their archive files. If the periodic boundaries are not crucial for your analysis, you can strip the trajectories of box information with a simple sed command, e.g.

sed '/90.000000\ \ 90.000000\ \ 90.000000/d'
AAA\_eq2\_npt.arc > AAA\_eq2\_stripped.arc

If you wish, strip the trajectories of their box information, copy the archive files (\*.arc) back to your local machine using WinSCP as detailed in the introduction, and visualise in VMD using the file type 'Tinker'.

# **Analysing simulations**

Tinker comes with a whole suite of programs for analysing Tinker xyz or trajectory files, the most useful of which is probably the 'analyze' program. You can run analysis interactively simply by feeding it an input structure and key file (e.g. analyze AAA\_eq2\_npt.arc -k AAA\_eq2\_npt.key) and following the on screen instructions. Alternatively, many other analysis packages, such as MDTraj, Amber cpptraj, or Parmed, are Tinker-aware and can read and manipulate Tinker coordinate files to varying extents. With a little scripting, this allows a diverse range of analysis of AMOEBA simulations beyond what's available in Tinker. More on that, however, in Exercise 3.

# Comments

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