# **GROMACS Introductory Tutorial**

Gromacs ver 3.3.1

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## **GROMACS** Tutorial for Solvation Study of Spider Toxin Peptide.

Yu, H., Rosen, M. K., Saccomano, N. A., Phillips, D., Volkmann, R. A., Schreiber, S. L.: Sequential assignment and structure determination of spider toxin omega-Aga-IVB. *Biochemistry* 32 pp. 13123 (1993)

GROMACS is a high-end, high performance research tool designed for the study of protein dynamics using classical molecular dynamics theory. [1] This versatile software package is Gnu public license free software. You may download it from <a href="http://www.gromacs.org">http://www.gromacs.org</a>. GROMACS runs on linux, unix, and on Windows (a recent development).

**Synopsis**. In this tutorial, you will study a toxin isolated from the venom of the funnel web spider. Venom toxins have been used in the past to identify cation channels. Calcium ion channels regulate the entry of this ion into cells. Nerve signals are highly governed by the balance of ions in neuronal cells. It is hypothesized that exposed positively charged residues in venoms like the spider toxin here bind favorably to the negatively charged entrance of the cell's ion channel. The spider toxin in this tutorial has its positively charged residues pointing predominantly to one side of the peptide. The ion channel is blocked, resulting in blocked nerve signal leading to paralysis and ultimately to death (presumably via respiratory failure).

We will study this peptide toxin using explicit solvation dynamics. First, we will compare an in vacuo model to a solvated model. We will solvate the peptide in a water box followed by equilibration using Newton's laws of motion. We will compare and contrast the impact of counterions in the explicit solvation simulation. We will seek answers to the following questions:

Is the secondary structure stable to the dynamics conditions?

Are the side chains of the positively charged residues predominantly displaced to one side of the peptide structure? Do the counterions hold these positively charged residues in place or do they move around?

What role does water play in maintaining the structure of proteins?

**Note**: You will generate gromacs (\*.gro) structure files in this tutorial. To view these files, you must use VMD (Download from: <a href="http://www.ks.uiuc.edu/Research/vmd/">http://www.ks.uiuc.edu/Research/vmd/</a>). In addition, you should obtain a copy of the GROMACS user manual at <a href="http://www.gromacs.org">http://www.gromacs.org</a>.

Download 1OMB.PDB from the Protein Data Bank (http://www.rcsb.org/pdb/).

It is advisable to use DeepView (Download DeepView from <a href="http://www.expasy.ch/spdbv/">http://www.expasy.ch/spdbv/</a>) to preview the file if you know that your structure may be disordered (i.e. residues with missing side chains). DeepView will replace any missing side chains (However, beware as Deep View will mark those rebuilt side chains with a strange control character that can only be removed manually using a

text editor!). There are no missing side chains in this pdb file, so we will not worry about that in this exercise.

Create a subdirectory in your unix account called "fwspider". Within this new directory, create the following directories: "invacuo", "wet", and "ionwet". Use sftp to copy the 1OMB.pdb file to the fwspider subdirectories (place a copy in each subdirectory within the fwspider directory). (IMPORTANT! Whenever you sftp a text file to a unix system from Windows, be sure to convert the file to a unix text file. Do this using the **to\_unix** command (e.g. **to\_unix** filename filename converts filename to a unix text file. In RedHat Linux, use the **dos2unix** command. Text editors in Windows like MS Word add control characters that may cause errors in unix programs.)

Process the pdb file with pdb2gmx. The pdb2gmx (to view the command line options for this command just type pdb2gmx –h; In fact, to get help for any command in Gromacs just use the –h flag) command converts your pdb file to a gromacs file and writes the topology for you. This file is derived from an NMR structure which contains hydrogen atoms. Therefore, we use the –ignh flag to ignore hydrogen atoms in the pdb file. The –ff flag is used to select the forcefield (G43a1 is the Gromos 96 FF, a united atom FF). The –f flag reads the pdb file. The –o flag outputs a new pdb file (various file formats supported) with the name you have given it in the command line. The –p flag is used for output and naming of the topology file. The topology file is very important as it contains all of the forcefield parameters (based upon the forcefield that you select in the initial prompt) for your model. Recent studies have revealed that the SPC/E water model [2] performs best in water box simulations. [3] Use the spce water model as it is better for use with the long-range electrostatics methods. So, we use the –water flag to specify this model.

pdb2gmx -ignh -ff G43a1 -f 1OMB.pdb -o fws.pdb -p fws.top -water spce

Setup the Box for your simulations

editconf -bt cubic -f fws.pdb -o fws.pdb -d 0.9

What you have done in this command is specify that you want to use a simple cubic box. The  $-\mathbf{d}$  0.9 flag sets the dimensions of the box based upon setting the box edge approx 0.9 nm (i.e. 9.0 angstroms) from the molecule(s) periphery. Ideally you should set  $-\mathbf{d}$  at no less than 0.85 nm for most systems. [4]

[Special Note: **editconf** may also be used to convert gromacs files (\*.gro) to pdb files (\*.pdb) and vice versa. For example: **editconf** – **f** file.gro – **o** file.pdb converts file.gro to the pdb file file.pdb]

You may use the files generated from the this step to begin your in vacuo simulation. For the in vacuo work just move ahead to the energy minimization step followed by the molecular dynamics step (No position restrained dynamics necessary for in vacuo work. Why?).

Solvate the Box

**genbox** -cp fws.pdb -cs spc216.gro -o fws b4em.pdb -p fws.top

The **genbox** command generates the water box based upon the dimensions/box type that you had specified using **editconf**. In the command above, we specify the spc water box. The **genbox** program will add the correct number of water molecules needed to solvate your box of given dimensions.

# Setup the energy minimization.

Use the em.mdp file. Gromacs uses special \*.mdp files to setup the parameters for every type of calculation that it performs. Look into the contents of this file. It specifies a steepest descents minimization to remove bad van der Waals contacts. Edit the file and change **nsteps** to 400. If the minimization fails to converge, re-submit with nsteps = 500. (The minimization should converge in less than 400 steps; however, different platforms respond differently.) To re-submit the job, you will need to re-run grompp. (Note: the path to the c pre-processor may be different on your machine. Use the **which** command to locate [i.e. which cpp]!)

## Content of em.mdp:

```
title
                       FWS
                     /usr/bin/cpp; the c pre-processor
срр
define
                    = -DFLEXIBLE
                    = none
constraints
integrator
                      steep
                    = 0.002
                             ; ps !
                    = 400
nsteps
nstlist
                    = 10
ns type
                    = grid
rlist
                    = 1.0
coulombtype
                    = PME
rcoulomb
                    = 1.0
vdwtype
                    = cut-off
rvdw
                    = 1.4
                    = 0.12
fourierspacing
fourier nx
                  = 0
fourier ny
                  = 0
fourier_nz
pme_order
ewald_rtol
optimize_fft
                  = 0
                  = 4
                  = 1e-5
                        = yes
        Energy minimizing stuff
;
;
                       1000.0
emtol
emstep
                       0.01
```

#### Important aspects of the em.mdp file:

**title** – The title can be any given text description (limit 64 characters; keep it short and simple!) **cpp** – location of the pre-processor

**define** – defines to pass to the pre-processor. –DFLEXIBLE will tell grompp to include the flexible SPC water model instead of the rigid SPC into your topology. This allows steepest descents to minimize further

**constraints** – sets any constraints used in the model.

**integrator** – *steep* tells grompp that this run is a steepest descents minimization. Use *cg* for conjugate gradient.

**dt** – not necessary for minimization. Only needed for dynamics integrators (like **md**).

**nsteps** – In minimization runs, this is just the maximum number of iterations.

**nstlist** – frequency to update the neighbor list. nstlist = 10 (updates the list every 10 steps).

**rlist** – cut-off distance for short-range neighbor list.

**coulombtype** – tells gromacs how to model electrostatics. PME is particle mesh ewald method (please see the Gromacs user manual for more information).

**rcoulomb** – distance for the coulomb cut-off

vdwtype – tells Gromacs how to treat van der Waals interactions (cut-off, Shift, etc.)

rvdw – distance for the LJ or Buckingham potential cut-off

EM Stuff

**emtol** – the minimization converges when the max force is smaller than this value (in units of kJ mol<sup>-1</sup> nm<sup>-1</sup>)

emstep – initial step size (in nm).

Now process the files with **grompp**. **grompp** is the pre-processor program (the gromacs pre-processor "grompp" Get it! Sigh!). **grompp** will setup your run for input into **mdrun**.

**grompp** – **f** em.mdp – **c** fws b4em.pdb – **p** fws.top – **o** fws em.tpr

The -f flag in grompp is used to input the parameter file (\*.mdp). The -c flag is used to input the coordinate file (the pdb file, \*.pdb); -p inputs the topology and -o outputs the input file (\*.tpr) needed for mdrun.

<u>Using genion and the tpr file to add ions</u>. You may use the tpr file generated here to add counterions to your model to neutralize any net charge. Our model has a net charge of + 2.00. Therefore, we want to add two chloride ions. Copy the fws\_em.tpr file that you used for your explicit solvated model to your "ionwet" subdirectory. In addition, copy your fws.top and posre.itp files from your explicit solvation model to your ionwet subdirectory. Use the genion command to add the chloride ions.

genion -s fws em.tpr -o fws ion.pdb -nname CL- -nn 2 -g fws ion.log

Where **–nname** is the negative ion name (CL- for the Gromos G43a1 force field; see the ions.itp file for specifics wrt force field), **-nn** is the number of negative ions to add. The **–g** flag gives a name to the output log for genion.

When you process this command, you will be prompted to provide a continuous group of solvent molecules, which should be Group 12 (SOL). Type 12 then <enter>. You will notice that two solvent molecules have been replaced by Cl ions. Now you must make the following additions/corrections to your fws.top file:

Add

#include "ions.itp" (Note: This should be present by default already in version 3.2 & greater)

after the include statement for the force field. Subtract 2 from the total number of SOL molecules and add a line for 2 Cl molecules in the [ molecules ] section at the end of the file. You will also need to modify the temperature coupling scheme for your pr md.mdp and md.mdp files.

Now, you will use fws\_ion.pdb in place of fws\_b4em.pdb for generating the actual input tpr using grompp for energy minimization.

<u>Remember</u>: If you added the chloride ions, you will need to run the grompp step again. First remove the old fws\_em.tpr file, then run the next grompp command below. We added chloride ions in our model to neutralized the overall net charge of the model.

```
grompp -f em.mdp -c fws ion.pdb -p fws.top -o fws em.tpr
```

Run the energy minimization in background (Use the ampersand & at the end of the command).

```
mdrun -v -deffnm em
```

( **-deffnm** tells gromacs the default filename to use for the run. This is a very useful shortcut.) Use the **tail** command to check on the progress of the minimization.

```
tail -15 em.log
```

When the minimization is complete, you should see the following summary in your log file indicating that steepest descents converged. Use **tail -50** em.log

```
Steepest Descents converged to Fmax < 1000 in 245 steps Potential Energy = -2.0551497e+05 Maximum force = 8.9138922e+02 on atom 106 Norm of force = 5.0448242e+03
```

Setup the position-restrained molecular dynamics. What is position-restrained MD? You are restraining (or partially freezing, if you will) the atom positions of the macromolecule while letting the solvent move in the simulation. This is done to "soak" the water molecules into the macromolecule. The relaxation time of water is  $\sim 10$  ps. Therefore, we need to do dynamics in excess of 10 ps for position-restrained work. This example uses a 20.0 ps time scale (at least an order of magnitude greater. Larger models (large proteins/lipids) may require longer preequilibration periods of 50 or 100 ps or more. The settings below will work well for the Gromacs force field. Consult the Gromacs manual for optimal settings for other force fields (e.g. the

GROMOS 96 force field nstlist = 10 and rvdw = 1.4 are recommended). Under **coulombtype**, PME stands for "Particle Mesh Ewald" electrostatics. [5, 6] PME is the best method for computing long-range electrostatics (gives more reliable energy estimates especially for systems where counterions like Na $^+$ , Cl $^-$ , Ca $^{2+}$ , etc. are used). Due to the nature of this particular protein having exposed charged residues with a net +2 charge to the system, it is beneficial for us to use PME. It is even more beneficial for us to use counterions to balance the charge and set the system to net neutral. The **all-bonds** option under **constraints** applies the Linear Constraint algorithm for fixing all bond lengths in the system (important to use this option when dt > 0.001 ps). [7] Study the mdp file below.

## Content of pr.mdp.

```
title
                        FWS
                    = 10
warnings
                    = /usr/bin/cpp
срр
define
                    = -DPOSRES
constraints integrator
                  = all-bonds
                    = md
dt
                    = 0.002 ; ps!
                   = 10000 ; total 20.0 ps.
nsteps
nstcomm
nstxout
nstvout
                   = 1
                   = 250
                               ; collect data every 0.5 ps
                   = 1000
                    = ()
nstfout
                    = 10
nstlog
nstenergy
                    = 10
                   = 10
nstlist
ns_type
rlist
                  = grid
                   = 1.0
coulombtype
rcoulomb
vdwtype
rvdw
                 = PME
                    = 1.0
                    = cut-off
                    = 1.4
fourierspacing = 0.12
fourier_nx = 0
fourier ny
                 = 0
fourier_ny = 0
fourier_nz = 0
pme_order = 4
ewald_rtol = 1e-5
optimize_fft = yes
; Berendsen temperature coupling is on
Tcoupl = berendsen
tau_t = 0.1 0.1
tc-grps = protein no
ref t = 300 300
                             0.1
                    = protein non-protein
                    = 300 300
ref t
; Pressure coupling is on
Pcoupl = parrinello-rahman tau p = 0.5
                    = 0.5
tau p
compressibility = 4.5e-5
ref p = 1.0
ref p
; Generate velocites is on at 300 K.
gen_vel = yes
gen temp
                    = 300.0
gen seed
                   = 173529
```

Important things to know about the mdp file.

The –DPOSRE in the **define** statement tells Gromacs to perform position restrained dynamics.

The **constraints** statement is as previously discussed. *all-bonds* sets the LINCS constraint for all bonds. [7]

The **integrator** tells gromacs what type of dynamics algorithm to use (another option is "sd" for stochastic dynamics).

dt is the time step (we have selected 2 fs; however, this must be entered in units of ps!).

**nsteps** is the number of steps to run (just multiply nsteps X dt to compute the length of the simulation).

**nstxout** tells gromacs how often to collect a "snapshot" for the trajectory. (e.g. nstxout = 250 with dt = 0.002 would collect a snapshot every 0.5 ps)

**coulombtype** selects the manner in which Gromacs computes electrostatic interactions between atoms. (PME is particle mesh ewald; there are other options like cut-off).

**rcoulomb** and **rvdw** are the cutoffs (in nm; 1.0 nm = 10.0 angstroms) for the electrostatic and van der Waals terms.

The temperature coupling section is very important and must be filled out correctly.

**Tcoupl** = berendsen [8] (type of temperature coupling; another option is nose-hoover)

**tau\_t** = Time constant for temperature coupling (units = ps). You must list one per tc\_grp in the order in which tc\_grps appear.

**tc\_grps** = groups to couple to the thermostat (every atom or residue in your model must be represented here by appropriate index groups).

ref\_t = reference temperature for coupling (i.e. the temperature of your MD simulation in degrees K). You must list one per tc\_grp in the order in which tc\_grps appear.

When you alter the temperature, don't forget to change the **gen\_temp** variable for velocity generation.

**pcoupl** – is the Parrinello-Rahman barostat.

**pcoupltype** – isotropic means that the "box" will expand or contract evenly in all directions (x, y, and z) in order to maintain the proper pressure. <u>Note</u>: Use semiisotropic for membrane simulations. **tau p** – time constant for coupling (in ps).

**compressibility** – this is the compressibility of the solvent used in the simulation in bar<sup>-1</sup> (the setting above is for water at 300 K and 1 atm).

**ref**  $\mathbf{p}$  – the reference pressure for the coupling (in bar) (1 atm  $\sim 0.983$  bar).

Pre-process the pr.mdp file.

```
grompp –f pr.mdp –c em.gro –p fws.top –o pr.tpr
```

## nohup mdrun -deffnm pr &

Use the **tail** command to check the pr.log file.

The md.mdp parameter file looks very similar to the pr.mdp file. There are several differences. The define statement is not necessary as we are not running as position restrained dynamics.

<u>Content of md.mdp for explicit solvation (Special Note</u>: For *in vacuo*, remove the entries for "sol" in the temperature coupling section. For the simulation that includes the counterions, add an entry for the ions in the temperature coupling section.)

```
title
                     FWS
                     /usr/bin/cpp
cpp
constraints
                  = all-bonds
integrator
                 = md
                 = 0.002 ; ps!
nsteps
                 = 25000 ; total 50 ps.
nsteps
nstcomm
nstxout
                 = 1
                 = 500 ; collect data every 1 ps
nstvout
nstfout
                  = 0
                  = 0
                 = 10
nstlist
ns_type
rlist
                 = grid
                 = 1.0
coulombtype
rcoulomb
vdwtype
rvdw
                = PME
= 1.0
                = cut-off
= 1.4
rvdw
fourierspacing = 0.12
fourier_nx = 0
fourier_ny = 0
fourier_nz = 0
pme_order = 4
; Berendsen temperature coupling is on
Tcoupl = berendsen
tau t
                   = 0.1 0.1
                   = protein non-protein
tc-grps
                    = 300
ref t
                                  300
; Pressure coupling is on
Pcoupl = parrinello-rahman tau p = 0.5
tau p
                = 4.5e-5
compressibility
ref p
                 = 1.0
; Generate velocites is on at 300 K.
gen vel = yes
```

gen_temp		300.0 173529			
gen_seed	_	173329			

**grompp** – **f** md.mdp – **c** pr.gro – **p** fws.top – **o** md.tpr

# nohup mdrun -deffnm md &

Use the tail command to check the md.log file.

You may compress the trajectory using **trjconv** to save on disk space.

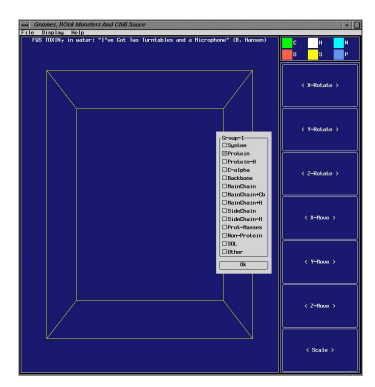
**trjconv** – **f** md.trr – **o** md.xtc

Once you have made the \*.xtc file, you may delete the \*.trr file.

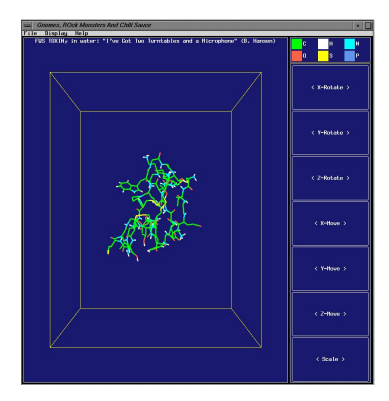
Use **ngmx** to view the trajectory (you may also download to your PC and use VMD to view the trajectory).

**ngmx** −**f** md.trr (or md.xtc) −**s** md.tpr

When the viewer comes up, you will see a dialog box with a number of selections. Check the box labeled "Protein", then click <u>Ok</u>.



ngmx's initial startup dialog.



Selecting "Protein" permits us to view the protein only without interference from the  $\sim$ 3,000 water molecules that occupy the rest of the box.

Use **X-Rotate** to rotate the box up and down (left mouse click goes up; right mouse click goes down). Use **Y-Rotate** to rotate the box from left to right or vice versa (left mouse click rotates to the left and right mouse click rotates to the right). **Scale** at the bottom permits you to zoom in and zoom out (use left mouse button to zoom in and right mouse button to zoom out).

To view another group in the model go to **Display > Filter ...** and the initial dialog box will come up again and you may opt to view some other index group (e.g. the backbone).

To view the animation of the MD trajectory go to **Display > Animate**. Controls used for the playback of the trajectory appear at the bottom of the window. Use the center arrow button to view one time step at time. Use the forward double arrow to play the whole trajectory, and use the "pause" button to the right to stop the trajectory playback. Use the double arrow to the left to reset the animation.

Unfortunately, the save as pdb option under the File menu in ngmx does not work. Therefore... *The best way to view the trajectory and select snapshots to save as \*.PDB files is to use visual molecular dynamics (VMD)*. (Download from: <a href="http://www.ks.uiuc.edu/Research/vmd/">http://www.ks.uiuc.edu/Research/vmd/</a> It's free to academics and it runs on unix and windows!).

#### **Analysis**

One of the major advantages of Gromacs (other than the fact that it is GNU public domain and free!) is the robust set of programs available for analyzing the trajectories. We will discuss a few of the most important analytic tools in the Gromacs arsenal here.

## **Groups**

# make\_ndx

The make\_ndx program is useful for generating groups (ID tags for specific atoms or residues that you may want to analyze). Gromacs generates default groups which may be adequate for your work as is. However, if you need to do more in depth analysis, use make\_ndx to tag specific items in your model. See the manual for more information.

How to use **make\_ndx** to setup index (ndx) files. Use make\_ndx to identify particular groups that you might want to freeze during a simulation or gather special energy information. Let's look at an example where we want to freeze the N and C terminal amino acids of a protein. Always use make\_ndx to create an index group for use with the **grompp** program.

In this sample case, we have a coordinate file of a collagen triple helix, post position restrained dynamics, where we want to freeze the N & C terminal for production run. We must first identify the residue #'s in the coordinate file (In our case clg\_b4md.pdb) that identify the N & C termini. The command line is simple enough ...

# make\_ndx -f clg\_b4md.pdb -o clg\_ter.ndx

You will see the following output (we left out the descriptive info at the beginning) followed by a command prompt (>).

```
Reading structure file
Going to read 0 old index file(s)
Analysing residue names:
Opening library file /usr/share/gromacs/top/aminoacids.dat
There are: 2194 OTHER residues
There are: 108 PROTEIN residues
There are: 0 DNA residues
Analysing Protein...
Analysing Other...

O System : 7365 atoms
1 Protein : 765 atoms
2 Protein-H : 687 atoms
3 C-alpha : 108 atoms
4 Backbone : 324 atoms
5 MainChain : 435 atoms
6 MainChain+Cb : 507 atoms
7 MainChain+H : 477 atoms
8 SideChain : 288 atoms
9 SideChain-H : 252 atoms
10 Prot-Masses : 765 atoms
11 Non-Protein : 6600 atoms
12 DRG : 21 atoms
13 SOL : 6579 atoms
14 Other : 6600 atoms
```

```
nr: group ! 'name' nr name 'splitch' nr Enter: list groups
'a': atom & 'del' nr 'splitres' nr 'l': list residues
't': atom type | 'keep' nr 'splitat' nr 'h': help
'r': residue 'res' nr 'chain' char
"name": group 'case': case sensitive 'q': save and quit
```

Use the 'r' command to enter the list of residue numbers that represent the N & C termini of the triple helix.

```
> r 1 36 37 72 73 108
15 r 1 36 37 72 73 108 : 51 atoms
```

**Note:** You may also use a dash to specify a residue number range (e.g. to specify residues 1 to 36 use > r 1-36)

The default name  $(r_1_36_37_72_73_108)$  giving to the new index group that you have just created is cumbersome. Lets rename it using the name command. We will use the index group # (15) in the command.

```
(15) in the command.
> name 15 Terminal
> v
Turned verbose on
```

```
      0 System
      : 7365 atoms

      1 Protein
      : 765 atoms

      2 Protein-H
      : 687 atoms

      3 C-alpha
      : 108 atoms

      4 Backbone
      : 324 atoms

      5 MainChain
      : 435 atoms

      6 MainChain+Cb
      : 507 atoms

      7 MainChain+H
      : 477 atoms

      8 SideChain
      : 288 atoms

      9 SideChain-H
      : 252 atoms

      10 Prot-Masses
      : 765 atoms

      11 Non-Protein
      : 6600 atoms

      12 DRG
      : 21 atoms

      13 SOL
      : 6579 atoms

      14 Other
      : 6600 atoms

      15 Terminal
      : 51 atoms
```

```
nr : group ! 'name' nr name 'splitch' nr Enter: list groups
'a': atom & 'del' nr 'splitres' nr 'l': list residues
't': atom type | 'keep' nr 'splitat' nr 'h': help
'r': residue 'res' nr 'chain' char
"name": group 'case': case sensitive 'q': save and quit
```

>

We used the 'v' command to verify that our name change was successful. To finish up, use the 'q' command to save and quit and you're done.

Now how do we freeze the groups? Easy, add the following lines to your md.mdp file

Remember, you must include the new index file when you use this md.mdp file to create your run input file (tpr) using grompp. Use the –n flag to grompp. For example ...

```
grompp -f md.mdp -c pr.gro -p clg.top -n clg ter.ndx -o md.tpr
```

# **Properties**

## g confrms

To compare the final structure to the original PDB file obtained from the PDB use g\_confrms (for a description use g\_confrms -h). This program takes two structures and performs a least squares fit.

```
g confrms -f1 1OMB.pdb -f2 md.gro -o fit.pdb
```

You will be prompted to select a group (select the Backbone (group 4) both times). The program reports an RMS deviation and produces an output file (fit.pdb). The output file will show the two structures superimposed on each other.

#### g covar

Use to compute the covariance matrix (see manual for more info). May also be used to compute an average structure from a dynamics trajectory. For example to compute the average of the last 200 ps of a 1 ns dynamics run use...

```
g covar – f traj.xtc – s topol.tpr – b 801 – e 1000 – av traj avg.pdb
```

<u>Warning</u> – Average structures tend to be crude. Minimization is recommended.

#### g\_energy

Use this program to plot energy data, pressure, volume, density, etc.

```
g energy –f md.edr –o fws pe.xvg
```

You will be prompted for the specific data that you want to include in the output file (\*.xvg). The output file is a type of spreadsheet file that can be read using Xmgr or Grace. This file is a text file and can be read into Microsoft Excel; however, you will need to do some light editing of the file.

For example in our case above (your results may be different), we see the following ...

Select the terms you want from the following list

G96Angle	Proper-Dih.	Improper-Dih.	LJ-14	Coulomb-14
LJ-(SR)	LJ-(LR)	Coulomb-(SR)	Coulrecip.	Potential
Kinetic-En.	Total-Energy	Temperature	Pressure-(bar)	Box-X
Box-Y	Box-Z	Volume	Density-(SI)	pV
Vir-XX	Vir-XY	Vir-XZ	Vir-YX	Vir-YY
Vir-YZ	Vir-ZX	Vir-ZY	Vir-ZZ	Pres-XX-(bar)
Pres-XY-(bar)	Pres-XZ-(bar)	Pres-YX-(bar)	Pres-YY-(bar)	Pres-YZ-(bar)
Pres-ZX-(bar)	Pres-ZY-(bar)	Pres-ZZ-(bar)	#Surf*SurfTen	Pcoupl-Mu-XX
Pcoupl-Mu-YY	Pcoupl-Mu-ZZ	Mu-X	Mu-Y	Mu-Z
T-Protein	T-SOL	T-CL-	Lamb-Protein	Lamb-SOL
Lamb-CL-				

For potential energy, type "Potential" <enter> Hit the <enter> key again

We get a summary with average PE and RMSD in kJ/mol

```
Last frame read 500 time 100.000
```

Statistics over 50001 steps [ 0.0000 thru 100.0000 ps ], 1 data sets

Energy	Average	RMSD	Fluct.	Drift	Tot-Drift
Potential	-192661	311.547	311.501	0.18468	18.4684

To read \*.xvg files into Grace use the following command:

## xmgrace -nxy fws pe.xvg

You may obtain Grace from <a href="http://plasma-gate.weizmann.ac.il/Grace/">http://plasma-gate.weizmann.ac.il/Grace/</a>. Grace runs on Linux and in Unix only. <a href="https://grace.or.or.org/">If you do not have Grace or Xmgr, just import or read the \*.xvg file into MS Excel as a space delimited file.</a>

# g\_gyrate

Use **g\_gyrate** to measure the radius of gyration. This quantity gives a measure of the "compactness" of the structure. This gives a measure of the mass of the atom(s) relative the center of mass of the molecule.

g gyrate -f md.trr -s md.tpr -o fws gyrate.xvg

## g rms and g rmsdist

These programs are useful for measuring root mean square deviations in the structure. Use **g\_rms** to evaluate the deviation of the structure from the original starting structure over the course of the simulation. (The **-dt** 10 option tells the program to write every tenth frame)

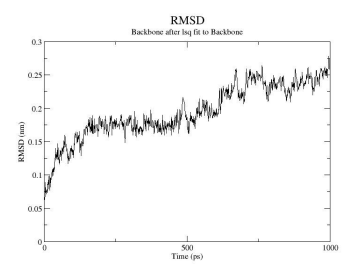
To compare rmsd to the NMR structure use the following command ...

Select group 4 (Backbone) for the least squares fit. The program will generate a plot of rmsd over time (rmsd.xvg) which you may import into Excel as a space delimited file.

#### g rmsf

Computes root mean square fluctuation of atom positions. Like **g\_covar** this command may be used to compute average structures. For example, to calculate the average of the last 500 ps of a 2 ns (2000 ps) dynamics simulation, use the following...

Select a range where you observe some equilibrium from your RMSD plot (computed using **g rms**). For example ...



The example above is the result of a 1 ns simulation (your results might be different). This simulation still needs to be run even longer as it has not completely equilibrated. In the example above, we might use the 200 to 500 ps range for computing an average structure as we observe a leveling off in deviation from the reference (initial structure). The command would be ...

Select group 1 "Protein" when prompted.

<u>Warning</u> – Average structures tend to be crude. Minimization is recommended.

Suggested em.mdp files setup for an *in vacuo* minimization. Perform steepest descents first followed by conjugate gradient. <u>CAUTION!</u> You may need to run **pdb2gmx** to setup a new set of topology files for your minimization especially if you have isolated a specific group in computing your average structure (as opposed to the entire system).

```
Steepest Descents in vacuo
     User kerrigan (236)
;
     Input file
;
;
                    = /usr/bin/cpp
срр
constraints
integrator
nsteps
                   = none
                  = steep
                  = 400
     Energy minimizing stuff
;
                    = 1000
emtol
emstep
                   = 0.01
nstcomm
ns_type
morse
                   = 1
                 - 1
= grid
                    = no
morse
coulombtype = Shift
vdw_type = Shift
rlist - ...
rcoulomb = 1.2
= 1.2
rcoulomb_switch = 1.0
rvdw_switch = 1.0
epsilon r
                    = 6.0
Tcoupl
                    = no
Pcoupl
                    = no
gen vel = no
```

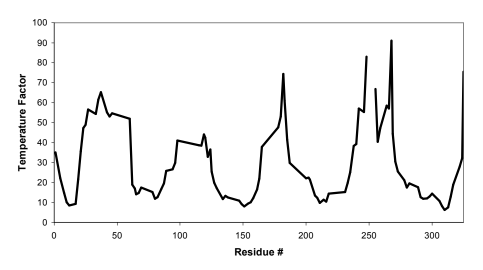
```
; Conjugate gradient with morse potential
; in vacuo
; User kerrigan (236)
;
; Input file
;
cpp = /usr/bin/cpp
constraints = none
integrator = cg
nsteps = 3000
;
; Energy minimizing stuff
;
emtol = 100
emstep = 0.01
```

nstcgsteep	=	1000
nstcomm	=	1
morse	=	yes
coulombtype	=	Shift
vdw type	=	Shift
ns type	=	grid
rlist	=	1.4
rcoulomb	=	1.2
rvdw	=	1.2
rcoulomb switch	=	1.0
rvdw switch	=	1.0
epsilon r	=	6.0
Tcoupl	=	no
Pcoupl	=	no
gen_vel	=	no

The **g\_rmsf** command may also be used to compute temperature factors. The computed *temperature factors* can be compared to x-ray crystal structure temperature factors.

g rmsf-s md.tpr-f md.xtc-o rmsf.xvg-oq fws bfac.pdb

Use the "Backbone" group once again.



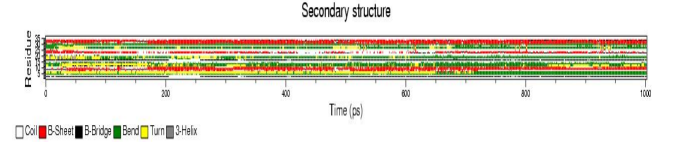
# do dssp

Use the do\_dssp command to analyze the secondary structure of your model. You must have the dssp program (<a href="http://swift.cmbi.ru.nl/gv/dssp/">http://swift.cmbi.ru.nl/gv/dssp/</a>) installed (in /usr/local/bin). [9]

do dssp -s md.tpr -f md.xtc -o fws ss.xpm

Select Group 1 (Protein) for the calculation. Use the xpm2ps program to convert the xpm file to eps format. Then use the ImageMagick convert program to convert the eps file to png or whatever desired image format.

xpm2ps -f fws\_ss.xpm -o fws\_ss.eps
convert fws\_ss.eps fws\_ss.png



Where residue # is on the y axis and time (ps) is on the x-axis. As seen from the original NMR structure below ...

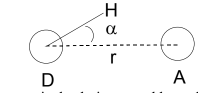


From the dssp plot above, we see 3 regions of red for the 3 sheets (displayed in yellow in the structure image to the left). The one short region in the middle is the least stable. The image to the left was prepared using the Pymol program. (http://pymol.sourceforge.net/)

# g hbond

Use **g\_hbond** to measure the number of hydrogen bonds, hbond distances & angles between molecules or groups through the course of the simulation.

**g hbond –f** md.trr –**s** md.tpr –**num** fws hnum.xvg



Geometrical relations used by g\_hbond.

The defaults in Gromacs 3.3.1 are:

 $r \le 0.35nm$  $\alpha \le 30^{\circ}$ 

Use the  $-\mathbf{r}$  and  $-\mathbf{a}$  flags to set other limits. By default,  $g_{\text{h}}$  bond analyzes the donor-acceptor  $(r_{DA})$  distance). You may change this behaviour by setting the  $-\mathbf{da}$  flag to "no". Setting da to no instructs  $g_{\text{h}}$  bond to analyze the  $r_{HA}$  distance.

See the Gromacs manual for more options.

## g saltbr

Use **g\_saltbr** to analyze salt bridges between residues in your simulation. The program outputs a set of \*.xvg files, which give distances between minus/minus, charged residues plus/minus (the most interesting) and plus/plus charged residues.

g saltbr -f md.trr -s md.tpr

How To Save – specific time points from a trajectory as \*.PDB files:

To get a specific frame (3000 ps in this example) instead of the whole trajectory as a pdb file use additionally the **-dump** option, e.g.

trjconv -f traj.xtc -s file.tpr -o time 3000ps.pdb -dump 3000

Perform other analysis tasks as recommended by your instructor.

# Appendix.

How to restart a crashed run.

**tpbconv** -**s** prev.tpr -**f** prev.trr -**e** prev.edr -**o** restart.tpr

**mdrun -s** restart.tpr **-deffnm** myrestart (the **-deffnm** option sets the default filename for all file options to mdrun)

How to extend a run.

tpbconv -f traj.trr -s topol.tpr -e ener.edr -o tpxout.tpr -time \$VALUE -until \$VALUE

Where \$VALUE = ps (e.g. if you wanted to extend a 2 ns run to 5 ns, then \$VALUE = 5000)

How to setup a parallel run

**grompp** -**np** # -**f** md.mdp -**c** pr.gro -**p** fws.top -**o** md.tpr

Where the **-np** flag is used to designate the number of nodes for parallel execution.

Then use mdrun mpi to run the parallel job. For example on the UMDNJ SunFire use ...

mprun –np #/products/gromacs/sparc-sun-solaris2.8/bin/mdrun mpi –deffnm md

Use the above command in your Sun Grid Engine batch submission script!

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