# **Simulation with Amber**

Naf Guo 2025 Ontion 2: Getting source code in tar format

# https://ambermd.org/GetAmber.php

从2024年的某一天开始,大善人amber学术免费了! 我们需要下载amber24和ambertools24

Option 2.	octaining soc	noc oode iii t	al Ioiiiiat		
					s distributed in source C++ and Fortran9
"Download" e4232d79e1e Note: Installa	button. The file b69f679db75c2 tion instructions	e size you shou le6ac2906.	of the <u>Amber 2</u>	out <b>541 MB</b> , a	ing form and click the and the md5sum of the
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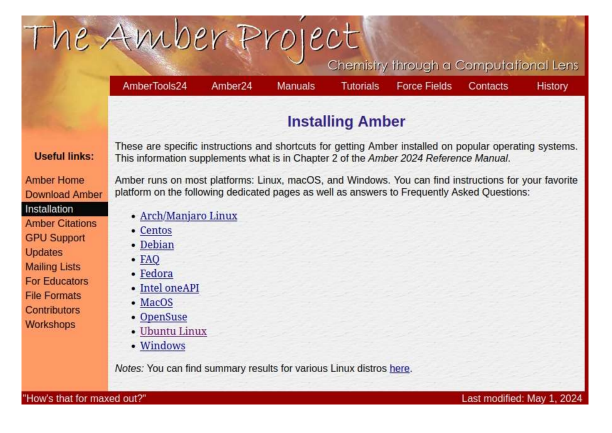
Getting Amber24 for non-com	mercia	al us	е					
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The downloaded file will be 0e9a01f2f2d5f0c1ebc75d2bdd5cde12.	about	173	MB,	and	have	an	md5sum	of
Please note: You must also download an Installation instructions are in Chapter 2 instructions for many systems may be found	2 of the	e Amb	er 2024	Refer	ence M	anual.		

Ambertools24的安装包

Amber24的安装包

# **Install Amber24**

https://ambermd.org/Installation.php



经过gromacs的洗礼以后,我们对怎么安装软件应该有一定的熟练度了, amber官方提供了不同操作系统下的安 装教程。

## **Install Amber24 on Ubuntu**

#### 首先使用apt安装依赖

```
Ubuntu 18.04 - Ubuntu 20.04

The following command should install all necessary packages to compile Amber on any Ubuntu version in the range above:

apt -y update
apt -y install tcsh make \
gcc gfortran \
flex bison patch bc wget \
xorg-dev libz-dev libbz2-dev

If you want to install Amber in parallel, you can use OpenMPI through:

apt -y install openmpi-bin libopenmpi-dev openssh-client
```

### 之后使用amber提供的run\_cnake脚本安装amber

#### Building with cmake

We highly recommend that you refer to Chapter 2 of the Amber 2024 Reference Manual for detailed instructions on how to install Amber 24.

Since Amber20, the build system has move to cmake. A script called run\_cmake is available in the amber24\_src/build directory. For most users, the options chosen in this script should be OK.

The installation of Amber is performed in two steps: cmake configuration, then building and install:

cd amber24 src/build

- # optional: edit the run\_cmake script to make any needed changes;
- # most users should not need to do this
- run cmake
- # Next, build and install the code:

make install

#### 对ubuntu24.04

sudo apt install tcsh make gcc gfortran flex bison patch bc wget xorg-dev libz-dev libbz2-dev

tar xvfj AmberTools24.tar.bz2

tar xvfj Amber24.tar.bz2

解压缩

```
# Assume this is Linux:

cmake $AMBER_PREFIX/amber24_src \
    -DCMAKE_INSTALL_PREFIX=$AMBER_PREFIX/amber24 \
    -DCOMPILER=GNUL \
    -DDPI=FALSE_-DCUDA=FALSE_DINSTALL_TESTS=TRUE \
    -DDOWNLOAD_MINICONDA=TRUE \
    2>&1 | tee cmake.log
```

之后极其重要的一步:修改build文件夹内的run\_cmake脚本,修改DCUDA后为TRUE以编译支持GPU加速的pmemd

./run\_cmake

make install -j 4

source /home/path-to-amber-dir/amber24/amber.sh

# **Amber Tutorials**

Amber具有丰富的官方教程

https://ambermd.org/tutorials/

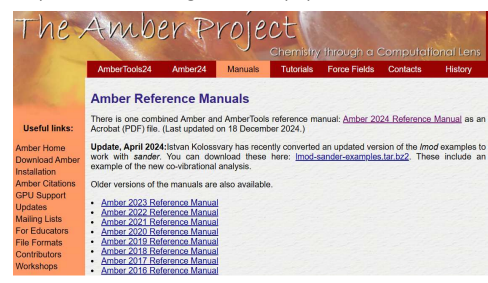


#### 几个与模拟强相关的教程

https://ambermd.org/tutorials/basic/tutorial7/index.php https://ambermd.org/tutorials/basic/tutorial8/index.php https://ambermd.org/tutorials/basic/tutorial13/index.php https://ambermd.org/tutorials/basic/tutorial14/index.php

#### Amber具有详细的文档

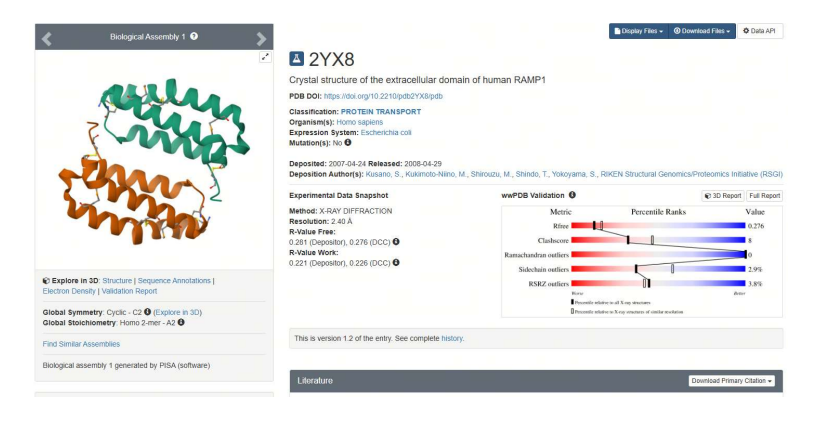
https://ambermd.org/Manuals.php



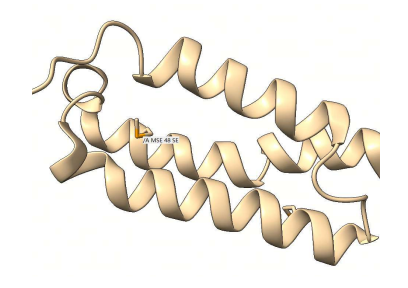
# **Download PDB**

# https://ambermd.org/tutorials/basic/tutorial7/index.php

我们就按照官方教程,下载2YX8作为测试体系



# **Protein Preparation**



此结构有一个有趣的地方,就是它有二个残基是MSE,所谓MSE,就是把甲硫氨酸MET里的硫(S)换成了硒(Se)

需要手动修改回MET

MOTA	167	0D2	ASP A	47	10.174	7.560	35.935	1.00 54.62	0
HETATM	168	N	MSE A	48	6.674	9.818	33.156	1.00 45.09	N
HETATM	169	CA	MSE A	48	6.049	10.959	32.490	1.00 44.83	С
HETATM	170	С	MSE A	48	4.687	10.621	31.903	1.00 44.85	С
HETATM	171	0	MSE A	48	3.782	11.457	31.884	1.00 44.46	0
HETATM	172	CB	MSE A	48	6.968	11.508	31.392	1.00 42.34	С
HETATM	173	CG	MSE A	48	8.206	12.210	31.932	1.00 37.42	С
HETATM	174	SE	MSE A	48	7.764	13.636	33.161	1.00 33.09	SE
HETATM	175	CE	MSE A	48	7.589	15.051	31.873	1.00 30.88	С
ATOM	176	N	GLU A	49	4.548	9.390	31.429	1.00 45.27	N

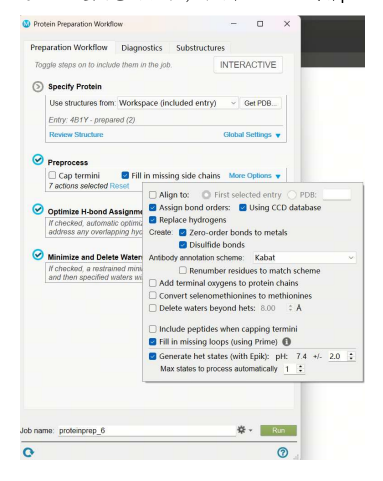


MOTA	167	0D2	ASP	A	47	10.174	7.560	35.935	1.00 54.62	0
MOTA	168	N	MET	A	48	6.674	9.818	33.156	1.00 45.09	N
MOTA	169	CA	MET	A	48	6.049	10.959	32.490	1.00 44.83	C
MOTA	170	C	MET	A	48	4.687	10.621	31.903	1.00 44.85	C
MOTA	171	0	MET	A	48	3.782	11.457	31.884	1.00 44.46	0
MOTA	172	CB	MET	Α	48	6.968	11.508	31.392	1.00 42.34	C
MOTA	173	CG	MET	A	48	8.206	12.210	31.932	1.00 37.42	C
MOTA	174	SD	MET	A	48	7.764	13.636	33.161	1.00 33.09	S
MOTA	175	CE	MET	A	48	7.589	15.051	31.873	1.00 30.88	C
MOTA	176	N	GLU	A	49	4.548	9.390	31.429	1.00 45.27	N

https://ambermd.org/tutorials/basic/tutorial7/index.php

# **Protein Preparation**

之后就是熟悉的流程,使用maestro或者pdbfixer进行蛋白准备



此处需要记录一下准备好的蛋白带电量,比如-3

https://htmlpreview.github.io/?https://github.com/openmm/pdbfixer/blob/master/Manual.html



pdbfixer 4b1y.pdb --output protein.pdb --keep-heterogens=none --add-residues --verbose

# **Protein Preparation**

之后记得处理质子化状态

使用maestro分配的质子化状态

https://github.com/Sept-naf/gromacs-tutorials/blob/main/maestro2amber.py

python maestro2amber.py maestro\_format.pdb amber\_readable.pdb

使用pdb2pqr分配质子化状态

https://pdb2pqr.readthedocs.io/en/latest/index.html

pip install pdb2pqr -i https://pypi.tuna.tsinghua.edu.cn/simple

对一个已经使用maestro或者pdbfixer处理好了的pdb文件

pdb2pqr input.pdb output.pdb --ffout AMBER --with-ph 7.4

此时得到的文件就带有amber力场格式的pH7.4下的质子化状态

# **Build Simulation System**

到这里蛋白处理好了, 可以搭建模拟体系了, amber搭建体系需要用tleap

# tleap

使用tleap之后会进入tleap的界面

(base) naf@yu-carve-hard-jade:~/Documents/tutorials/amber\$ tleap

- -I: Adding /home/naf/software/amber24/dat/leap/prep to search path.
- -I: Adding /home/naf/software/amber24/dat/leap/lib to search path.
- -I: Adding /home/naf/software/amber24/dat/leap/parm to search path.
- -I: Adding /home/naf/software/amber24/dat/leap/cmd to search path.

Welcome to LEaP!

(no leapro in search path)

通过左边进入tleap的提示是不是已经可以猜测amber力场参数文件都在哪找了

# https://ambermd.org/tutorials/pengfei/index.php

Structure File	Library Files	Parameter Files
Molecules Atom Names	Units (Residues) Atom Names	
	Atom Types Charges	Atom Types
Connections	Connections	
Coordinates	Coordinates	2712000-0-000
	Masses	Masses
	Bonded Params	Bonded Params
	Nonbond Params	Nonbond Params

Topology	Coordinates
Units (Residues)	
Atom Names	
Atom Types	
Charges	
Connections	
	Coordinates
Masses	
Bonded Params	
Nonbond Params	

Amber官方提供了leap的教学,它将 PDB与力场参数文件组织起来,变成能 进行模拟的拓扑文件和坐标文件

# **Build Simulation System**

## tleap

## 在tleap内

source leaprc.protein.ff19SB 导入蛋白, 水和离子的力场参数 source leaprc.water.opc 导入做好了蛋白质准备的,处理好了质子 ramp=loadpdb 2YX8 prepared.pdb 化状态的蛋白, 且将其命名为ramp bond ramp.27.SG ramp.82.SG 手动连接所有的二硫键 bond ramp.40.SG ramp.72.SG bond ramp.57.SG ramp.104.SG 由于前面知道蛋白带负电, 这里用 addlons2 ramp Na+ 0 Na+中和电荷:对正电体系则用CI-给ramp添加水盒子 solvateOct ramp OPCBOX 10.0 addlonsRand ramp Na+ 19 Cl- 19 添加离子, 使体系盐浓度为0.15M, 计算方法在下一页 存储拓扑文件prmtop和坐标inpcrd SaveAmberParm ramp RAMP.prmtop RAMP.inpcrd

# Calculating Salt in an Explicit Water System

https://ambermd.org/tutorials/basic/tutorial8/index.php

Scaling up box by a factor of 1.413369 to meet diagonal cut criterion

Solute vdw bounding box:

45.400 31.108 30.346

Total bounding box for atom centers: 73.667 73.667 73.667

(box expansion for 'iso' is 55.9%)

Solvent unit box:

18.865 18.478 19.006

The number of boxes: x= 4 y= 4 z= 4

Volume: 208141.839 A^3 (oct)

Total mass 110188.212 amu, Density 0.879 g/cc

Added 5569 residues.

2. Convert the volume of the system in Å<sup>3</sup> to liters using the following conversion factors.

Be sure to distribute the exponents when performing the calculations.

$$208141.839\,\mathring{\mathrm{A}}^{3}x\frac{1\,m^{3}}{\left(10^{10}\mathring{\mathrm{A}}\right)^{3}}\,x\,\frac{(10^{2}cm)^{3}}{1\,m^{3}}\,x\,\frac{1\,mL}{1\,cm^{3}}\,x\,\frac{1\,L}{10^{3}\,mL} = 2.08141839\,x\,10^{-22}L$$

3. Determine how many chloride ions are present in one liter of solution at a concentration of 150 mM.

$$\frac{150 \ mmol}{1 \ L} \ x \ \frac{1 \ mol}{10^{3} mmol} \ x \ \frac{6.022 \ x \ 10^{23} \ Cl^{-} \ atoms}{1 \ mol} = 9.03 \ x \ 10^{22} Cl^{-} \ atoms/L$$

4. Determine how many chloride ions are needed in the system.

Multiply the volume of the box by the concentration of Cl<sup>-</sup> ions and round to the nearest whole number.

$$2.08141839 \times 10^{-22} L \times 9.03 \times 10^{22} \frac{Cl^{-} atoms}{L} = 18.8 Cl^{-} ions \approx 19 Cl^{-} ions$$

在加水盒子时能看到整个体系的体积, 记录下来

把体积单位从立方埃换算成升

计算对这么多升溶液,需要多少离子能达到0.15M

计算具体需要的离子数目

总结其实就是Volume乘以一个确定的数值,然后四舍五入(或者四舍六入五成双)

208141.839 \* 9.033e-05

### More detail

> charge ramp

Total unperturbed charge:

Total perturbed charge:

我们已经做好模拟前的准备了,但是,出于学习的考虑,再更进一步前,我们看几条leap命令

```
> desc ramp.27.SG
MOTA
            Normal
                        Perturbed
             SG
                            SG
Name:
Type:
              S
             -0.0984
                            0.000
Charge:
Polarization: 0.0000
                           0.000
                      (not affected by pert)
Element: S
Atom flags: (decimal 196608 hex 0x30000)
       posfxd n posblt n posdrwn n selected n
       pert n notdisp n touched Y posknwn Y
       internal n needsmin n needsbuild n
Atom position: 14.367393, -8.553670, -4.209354
Atom velocity: 0.000000, 0.000000, 0.000000
 Bonded to .R<NCYX 27>.A<CB 7> by a single bond.
 Bonded to .R<CYX 82>.A<SG 8> by a single bond.
```

0.000000

0.000000

```
> check ramp
Checking 'ramp'....
Warning: Close contact of 1.169 angstroms between nonbonded atoms HH and HE2
----- .R<TYR 66>.A<HH 15> and .R<HIE 97>.A<HE2 13>
Checking parameters for unit 'ramp'.
Checking for bond parameters.
Checking for angle parameters.
check: Warnings: 20977
Unit is OK.
```

charge命令, 检查体系电量

desc命令, 检查体系部分的情况

check命令,检查体系力场是否完整

# **Save PDB**

Amber有一个非常讨厌的地方,就是它会对你的体系残基重新编号,从1开始

> savepdb ramp out.pdb
Writing pdb file: out.pdb
 printing CRYST1 record to PDB file with box info

Warning: Converting N-terminal residue name to PDB format: NCYX -> CYX
Warning: Converting C-terminal residue name to PDB format: CSER -> SER

将体系存一份PDB

#### 打开你会发现,蛋白残基范围变成了1-81

CRYST1	73.	069	65.678	5	8.496 90.0	00.00	90.00	P 1	er veran	ATOM	1277	N	SER	81	-10.719	14.186	7.108	1.00	0.00
MOTA	1	N	CYX	1	17.613	-8.356	-4.929	1.00	0.00	ATOM	1278	Н	SER	81	-9.994	13.541		1.00	0.00
MOTA	2	H1	CYX	1	17.543	-8.149	-5.916	1.00	0.00	ATOM	1279	CA	SER	81	-11.750	14.550	8.095	1.00	0.00
ATOM	3	H2	CYX	1	18.588	-8.478	-4.692	1.00	0.00	ATOM	1280	HA	SER	81	-12.735	14.529	7.629	1.00	0.00
MOTA	4	H3	CYX	1	17.048	-7.693	-4.418	1.00	0.00	ATOM	1281	CB	SER	81	-11.738	13.545	9.276	1.00	0.00
MOTA	5	CA	CYX	1	16.985	-9.672	-4.689	1.00	0.00	ATOM	1282	HB <sub>2</sub>	SER	81	-12.538	13.834	9.957	1.00	0.00
ATOM	6		CYX	1		-10.354	-4.272	1.00	0.00	ATOM	1283	HB3	SER	81	-11.949	12.558	8.866	1.00	0.00
MOTA	7	CB	CYX	1	15.798	-9.562	-3.698	1.00	0.00	ATOM	1284	OG	SER	81	-10.506	13.511	9.980	1.00	0.00
MOTA	8		CYX	1		-10.559	-3.464	1.00	0.00	ATOM	1285	HG	SER	81	-10.563	12.873	10.695	1.00	0.00
MOTA	9	0.0000000	CYX	1	16.133	-9.074	-2.782	1.00	0.00	ATOM	1286	C	SER	81	-11.518	15.982	8.552	1.00	0.00
MOTA	10	SG	CYX	1	14.367	-8.554	-4.209	1.00	0.00	ATOM	1287	0	SER	81	-12.267	16.544	9.350	1.00	0.00
ATOM	11	C	CYX	1		-10.329	-6.039	1.00	0.00	ATOM	1288	0XT	SER	81	-10.554	16.622	8.135	1.00	0.00
MOTA	12	0	CYX	1	17.022	-9.899	-7.113	1.00	0.00	TER									

体系准备好了,可以退出tleap,正式开始模拟了,首先仍然是能量最小化

pmemd -O -i 1min.in -o 1min.out -p RAMP.prmtop -c RAMP.inpcrd -r 1min.rst7 -inf 1min.info -ref RAMP.inpcrd -x mdcrd.1min



#### 1min.in

```
minimization of solvent
&cntrl

imin = 1, maxcyc = 1000,

ncyc = 20, ntx = 1,

ntwe = 0, ntwr = 500, ntpr = 50,

ntc = 2, ntf = 2, ntb = 1, ntp = 0,

cut = 10.0,

ntr=1, restraintmask = ':1-81',

restraint_wt = 100.,

ioutfm=1, ntxo=2,

/
```

- ntr=1代表要对体系加限制
- restraintmask = ':1-81'代表对残基1-81加限制,注意这是 amber重排序过的编号,可以在上一页存出的pdb里确认
- restraint\_wt = 100 限制大小100 kcal/mol\*Ang^-2.

NSTEP = 648000 TIME(PS) =

1-4 NB =

Etot = -89860.5401 EKtot =

EELEC = -120003.9513 EHBOND =

0.0000 IMP =

256.8588 1-4 EEL =

能量最小化以后是体系加热

pmemd.cuda -O -i 2heat.in -o 2heat.out -p RAMP.prmtop -c 1min.rst7 -r 2heat.rst7 -inf 2heat.info -ref 1min.rst7 -x mdcrd.2heat

使用pmemd.cuda, 这是GPU加速的版本 上一步得到的坐标, 这一步的输入坐标

#### 2heat.in

```
&cntrl
imin = 0, nstlim = 2000000, dt = 0.001,
irest = 0, ntx = 1, ig = -1,
tempi = 100.0, temp0 = 298.0,
ntc = 2, ntf = 2, tol = 0.00001,
ntwx = 10000, ntwe = 0, ntwr = 1000, ntpr = 1000,
cut = 8.0, iwrap = 0,
ntt = 3, gamma_ln=1., ntb = 1, ntp = 0,
nscm = 0,
ntr=1, restraintmask=':1-81', restraint_wt=100.0
nmropt=1,
ioutfm=1, ntxo=2,
/
&wt TYPE="TEMPO", istep1=0, istep2=1000000, value1=100., value2=298., /
&wt TYPE="END", /
```

2ns

EAMBER (non-restraint) = -100582.4742NMR restraints: Bond = 0.000 Angle = Current Timing Info Total steps: 1000000 | Completed: 648000 ( 64.8%) | Remaining: Average timings for last 213000 steps: Elapsed(s) = 59.92 Per Step(ms) = 0.28 ns/day = 307.15 seconds/ns = 281.29 Average timings for all steps: Elapsed(s) = 179.10 Per Step(ms) =0.28 ns/day = 312.61 seconds/ns = Estimated time remaining: 1.6 minutes.

648.000 TEMP(K) = 228.63 PRESS =

2626.6844 VDWAALS =

0.0000 RESTRAINT =

15575.7389

10302.7109 EPtot 359.9953 DIHED

0.0000 CMAP

ntb=1, 恒体积

从0到1000000步将温度从100K升到298K

NPT平衡

pmemd.cuda -O -i 3md.in -o 3md.out -p RAMP.prmtop -c 2heat.rst7 -r 3md.rst7 -inf 3md.info -ref 2heat.rst7 -x mdcrd.3md

#### 3md.in

```
&cntrl

imin = 0, nstlim = 1000000, dt = 0.001,

irest = 1, ntx = 5, ig = -1,

temp0 = 298.0,

ntc = 2, ntf = 2, tol = 0.00001,

ntwx = 10000, ntwe = 0, ntwr = 1000, ntpr = 1000,

cut = 8.0, iwrap = 0,

ntt = 3, gamma_ln=1.0, ntb = 2, ntp = 1, barostat = 2,

nscm = 0,

ntr=1, restraintmask=':1-81', restraint_wt=100.

ioutfm=1, ntxo=2,

/
```

1ns

ntb=2, 恒压 ntp=1, 各向同性压浴 barostat=2, 蒙特卡洛压浴 pmemd.cuda -O -i 4md.in -o 4md.out -p RAMP.prmtop -c 3md.rst7 -r 4md.rst7 -inf 4md.info -ref 3md.rst7 -x mdcrd.4md

#### 4md.in

```
&cntrl

imin = 0, nstlim = 1000000, dt = 0.001,

irest = 1, ntx = 5, ig = -1,

temp0 = 298.0,

ntc = 2, ntf = 2, tol = 0.00001,

ntwx = 10000, ntwe = 0, ntwr = 1000, ntpr = 1000,

cut = 8.0, iwrap = 0,

ntt = 3, gamma_ln=1.0, ntb = 2, ntp = 1,

nscm = 0, barostat = 2,

ntr=1, restraintmask=':1-81', restraint_wt=10.

ioutfm=1, ntxo=2,

/
```

1ns

放松限制到10

pmemd -O -i 5min.in -o 5min.out -p RAMP.prmtop -c 4md.rst7 -r 5min.rst7 -inf 5min.info -ref 4md.rst7 -x mdcrd.5min

#### 5min.in

```
Minimization of everything excluding backbone &cntrl
imin = 1, maxcyc = 1000,
ncyc = 30, ntx = 1,
ntwe = 0, ntwr = 500, ntpr = 50,
ntc = 2, ntf = 2, ntb = 1, ntp = 0,
cut = 8.0,
ntr=1, restraintmask="@CA,N,C", restraint_wt=10.
ioutfm=1, ntxo=2,
/
```

之前约束了残基1-81的所有原子,此时只约束名为CA,N,C的原子,由于我们的体系里除了蛋白就是水和离子,所以这里实际上只约束了蛋白质的主链原子

# Simulating System: NPT平衡6

pmemd.cuda -O -i 6md.in -o 6md.out -p RAMP.prmtop -c 5min.rst7 -r 6md.rst7 -inf 6md.info -ref 5min.rst7 -x mdcrd.6md

#### 6md.in

```
&cntrl

imin = 0, nstlim = 1000000, dt = 0.001,

irest = 0, ntx = 1, ig = -1,

tempi = 298.0, temp0 = 298.0,

ntc = 2, ntf = 2, tol = 0.00001,

ntwx = 10000, ntwe = 0, ntwr = 1000, ntpr = 1000,

cut = 8.0, iwrap = 0,

ntt = 3, gamma_ln=1.0, ntb = 2, ntp = 1,

nscm = 0, barostat = 2,

ntr=1, restraintmask="@CA,N,C", restraint_wt=10.

ioutfm=1, ntxo=2,

/
```

1ns

NPT

pmemd.cuda -O -i 7md.in -o 7md.out -p RAMP.prmtop -c 6md.rst7 -r 7md.rst7 -inf 7md.info -ref 6md.rst7 -x mdcrd.7md

#### 7md.in

```
&cntrl

imin = 0, nstlim = 1000000, dt = 0.001,

irest = 1, ntx = 5, ig = -1,

temp0 = 298.0,

ntc = 2, ntf = 2, tol = 0.00001,

ntwx = 10000, ntwe = 0, ntwr = 1000, ntpr = 1000,

cut = 8.0, iwrap = 0,

ntt = 3, gamma_ln=1.0, ntb = 2, ntp = 1,

nscm = 0, barostat = 2,

ntr=1, restraintmask="@CA,N,C", restraint_wt=1.

ioutfm=1, ntxo=2,

/
```

1ns

放松约束到1

pmemd.cuda -O -i 8md.in -o 8md.out -p RAMP.prmtop -c 7md.rst7 -r 8md.rst7 -inf 8md.info -ref 7md.rst7 -x mdcrd.8md

#### 8md.in

```
&cntrl

imin = 0, nstlim = 1000000, dt = 0.001,

irest = 1, ntx = 5, ig = -1,

temp0 = 298.0,

ntc = 2, ntf = 2, tol = 0.00001,

ntwx = 10000, ntwe = 0, ntwr = 1000, ntpr = 1000,

cut = 8.0, iwrap = 0,

ntt = 3, gamma_ln=1.0, ntb = 2, ntp = 1,

nscm = 0, barostat = 2,

ntr=1, restraintmask="@CA,N,C", restraint_wt=0.1

ioutfm=1, ntxo=2,

/
```

1ns

放松约束到0.1

pmemd.cuda -O -i 9md.in -o 9md.out -p RAMP.prmtop -c 8md.rst7 -r 9md.rst7 -inf 9md.info -ref 8md.rst7 -x mdcrd.9md

#### 9md.in

```
&cntrl

imin = 0, nstlim = 1000000, dt = 0.001,

irest = 1, ntx = 5, ig = -1,

temp0 = 298.0,

ntc = 2, ntf = 2, tol = 0.00001,

ntwx = 10000, ntwe = 0, ntwr = 1000, ntpr = 1000,

cut = 8.0, iwrap = 0,

ntt = 3, gamma_ln=1.0, ntb = 2, ntp = 1,

nscm = 1000, barostat = 2,

ioutfm=1, ntxo=2,

/
```

1ns

无约束平衡

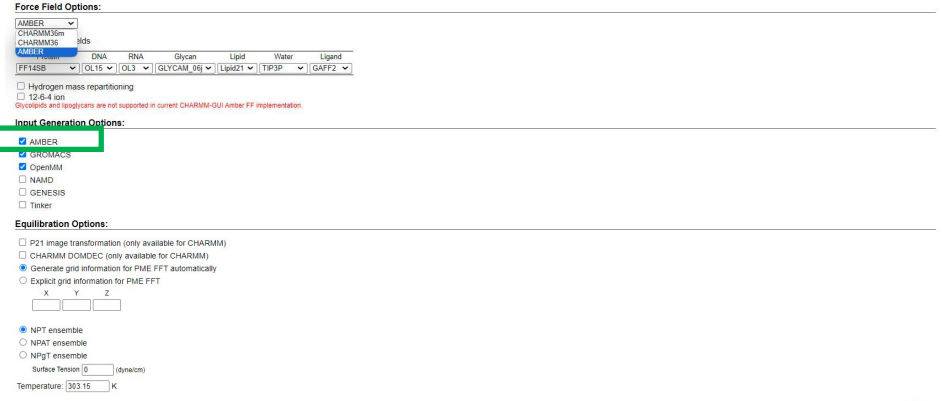
pmemd.cuda -O -i md.in -o md.out -p RAMP.prmtop -c 9md.rst7 -r md.rst7 -inf md.info -ref 9md.rst7 -x mdcrd.md

#### md.in

```
Explicit solvent molecular dynamics constant pressure 50 ns MD &cntrl
imin=0, irest=1, ntx=5,
ntpr=50000, ntwx=50000, ntwr=50000, nstlim=25000000,
dt=0.002, ntt=3, tempi=300,
temp0=300, gamma_ln=1.0, ig=-1,
ntp=1, ntc=2, ntf=2, cut=9,
ntb=2, iwrap=1, ioutfm=1,
/
```

# **Membrane Protein Simulation?**

#### Charmm-gui



Next Step: Next Step: