

IN-SILICO STRUCTURAL ANALYSIS AFFECTING THERMOSTABILITY IN
RECOMBINANT PSYCHROPHILIC CHITINASE
(CHI II) FROM *Glaciozyma antarctica* PI12

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The day has finally come. When I can say it out loud
'I made it'

To *mama & bapak*,
this is for you

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ABSTRACT

Cold-adapted enzymes are significant with structure flexibility and high catalytic activity at low temperature. High structural flexibility could be due to combination of several features such as weak intramolecular bonds, decreased compactness of hydrophobic core and reduced number of proline and arginine residues. However, to compensate the structural flexibility, cold-adapted enzymes are also thermolabile which causes them to be easily inactivated at elevated temperature. Therefore, it would be more interesting and beneficial if more stable cold-adapted enzymes are produced to fulfill the industrial needs. In this study, a novel cold-adapted chitinase (CHI II) from *Glaciozyma antarctica* PI12 was rationally designed to improve their thermostability thus make them more resistant to increased temperature. Four CHI II mutants were designed through rational design named as A157Q, I134P, mutant Loop and Y257R by manipulating the structural hydrophobicity, introduction of proline in the loop regions, introduction of arginine salt bridges and loop shortening. Mutant Loop was designed by removing 9 residues in loop regions thus makes loop involved became shorter. Stability of all mutants was first predicted through a computational approach where all structures were subjected to 10 ns molecular dynamics simulation at three temperatures; 273 K, 288 K and at 300 K. Based on the simulation, it was found that mutants I134P, mutant Loop and Y257R exhibited structural stability at 300 K. This conclusion was made based on low and stable root-mean square deviation (RMSD) value at 300 K in comparison to RMSD values at 288 K and 273 K. Low RMSD values indicated mutant structure experienced low structural deviation throughout the simulation. Besides, this observation is correlated with reduction of structure compactness (radius of gyration), reduced solvent accessible surface area and increased numbers of hydrogen and salt bridges. However, mutant A157Q experienced structure destabilization at 300 K. Substitution of helix-preferred residue, alanine with a thermolabile residue, glutamine had caused A157Q structure become loosely packed at 300 K indicating a thermal denaturation. To support the theoretical model, CHI II and all mutants were then cloned into *Pichia pastoris* expression vector pPICZαC and expressed in *P. pastoris* (GS115).

ABSTRAK

Enzim tahan sejuk adalah dikenali dengan struktur yang fleksibel serta aktiviti bermangkai yang tinggi pada suhu rendah. Fleksibiliti struktur adalah disebabkan oleh kombinasi beberapa ciri seperti ikatan intramolekul yang lemah, penurunan kepadatan teras hidrofobik dan pengurangan sisa prolina dan arginina. Walau bagaimanapun, untuk menebus kembali fleksibiliti strukturnya, enzim tahan sejuk bersifat termolabil yang menyebabkannya mudah untuk ternyahaktif pada suhu lampau tinggi. Oleh itu, ianya sangat menarik dan bermanfaat jika enzim tahan sejuk yang lebih stabil dapat dihasilkan bagi memenuhi keperluan industri. Dalam kajian ini, enzim kitinase tahan sejuk (CHI II) dari organisma *Glaciozyma antarctica* PI12 telah direkabentuk secara rasional untuk meningkatkan tahap termostabilitinya dan menjadikan CHI II lebih tahan kepada peningkatan suhu. Empat mutan CHI II telah di rekabentuk melalui rekabentuk rasional yang dinamakan sebagai A157Q, I134P, Gelung mutan dan Y257R dengan mengubah suai kehidrofobikan struktur, memperkenalkan prolina di kawasan gelung, memperkenalkan titian garam arginina serta pemendekan gelung. Gelung mutan telah direkabentuk dengan pemotongan 9 sisa di kawasan gelung menyebabkan gelung yang terlibat menjadi semakin pendek. Kestabilan kesemua mutan diramalkan terlebih dahulu melalui pendekatan pengkomputeran di mana kesemua struktur mutan tertakluk kepada simulasi 10 ns dinamik molekul yang dijalankan pada tiga suhu iaitu pada 273 K, 288 K dan 300 K. Berdasarkan simulasi yang dijalankan, mutan I134P, gelung mutan dan Y257R menunjukkan nilai sisihan punca min kuasa dua (RMSD) yang rendah dan stabil pada suhu 300 K, jika dibandingkan dengan nilai RMSD pada 288 K dan 273 K. Nilai RMSD yang rendah menggambarkan struktur mutan telah mengalami sisihan berstruktur rendah keseluruhan simulasi dijalankan. Sebaliknya, pemerhatian ini berkorelasi dengan penurunan kepadatan struktur (jejari legaran), penurunan luas permukaan boleh capai pelarut dan peningkatan ikatan hidrogen serta titian garam. Walau bagaimanapun, mutan A157Q mengalami penurunan kestabilan struktur pada 300 K. Penggantian sisa pilihan heliks, iaitu alanina dengan sisa termolabil, glutamina telah menyebabkan struktur A157Q menjadi longgar pada suhu 300 K menandakan penyahasian terma. Untuk menyokong model teori ini, CHI II dan kesemua mutan telah diklon ke dalam vektor ekspresi *Pichia pastoris* pPICZ α C dan dinyatakan dalam *P. pastoris* (GS115).

TABLE OF CONTENT

CHAPTER	TITLE	PAGE
	DECLARATION	ii
	DEDICATION	iii
	ACKNOWLEDGEMENT	iv
	ABSTRACT	v
	ABSTRAK	vi
	TABLE OF CONTENTS	vii
	LIST OF TABLES	xi
	LIST OF FIGURES	xii
	LIST OF SYMBOLS AND ABBREVIATIONS	xv
	LIST OF APPENDICES	xix
1	INTRODUCTION	1
	1.1 Background of study	1
	1.2 Problem statements	3
	1.3 Objectives	4
	1.4 Scopes of study	4
2	LITERATURE REVIEW	5
	2.1 Psychrophiles, a Cold-Loving Microbes	5
	2.2 Cold Adapted Enzymes	6
	2.3 Thermostability Improvement through rational Design	10

2.3.1	Proline Substitution to Improve Protein Thermostability	12
2.3.2	Improving Thermostability by Loop Shortening	13
2.3.3	Introduction of Salt Bridges to Improve Protein Thermostability	13
2.3.4	Substitution of Surface Alanine Residue to a More Hydrophilic Residues	14
2.4	Thermostability Studies Through Molecular Dynamic (MD) Simulation	14
2.5	Chitin	16
2.6	Chitinase, a Chitinolytic Enzyme	18
2.6.1	Chitinase Subfamily and Classifications	19
2.6.2	Functional Domain of Fungal Chitinases	23
2.6.3	Roles of Chitinases in Various Organisms	25
2.7	A novel Chitinase (CHI II) From <i>Glaciozyma antarctica</i> P112	26
2.8	Chitinases and Their Applications	28
2.8.1	Bioconversion of Chitin in Production of Single Cell Proteins	28
2.8.2	Enzymatic Hydrolysis in Production of Chitooligosaccharides	29
2.8.3	Biocontrol Agents	30
2.9	Heterologous Expression in <i>Pichia pastoris</i>	30
3	MATERIALS AND METHODS	33
3.1	Introduction	33
3.2	Source of Sequences	35
3.3	Tools and Software	35
3.4	Mutation Design Strategy	38
3.5	Construction of A157Q, I134P and Y257R 3D Models	38
3.6	Homology Modeling of Mutant Loop	38
3.7	Molecular Dynamics (MD) Simulation	39

3.7.1	Preparation Stage	40
3.7.2	Production Stage	40
3.7.3	Trajectories Analysis for CHI II and Mutants	41
3.7.3.1	Root Mean Square Deviation (RMSD)	41
3.7.3.2	Root Mean Square Fluctuation (RMSF)	41
3.7.3.3	Radius of Gyration	41
3.7.3.4	Analysis of Secondary Structure for CHI II and it Mutants	42
3.7.3.5	Solvent Accessible Surface Area (SASA)	42
3.7.3.6	Analysis of Hydrogen Bonds and Salt Bridges	42
3.8	Cloning of CHI II and It Mutants into <i>Pichia pastoris</i> Expression System	42
3.9	Expression of Recombinant CHI II in <i>Pichia Pastoris</i> strains	45
3.10	Screening of Mutant Expression	45
3.10.1	SDS PAGE Analysis	45
3.10.2	Western Blotting	46
3.10.3	Chitinase Assay	47
4	RESULTS AND DISCUSSION	49
4.1	Mutants Design Strategy	49
4.1.1	Introduction of Salt Bridge in Mutant Y257R	49
4.1.2	Loop Shortening in Mutant Loop	52
4.1.3	Introduction of Proline in Loop Regions for Mutant I134P	55
4.1.4	Substitution of Alanine to Glutamine in Mutant A157Q	56
4.2	Construction of Mutants 3D Models	57
4.2.1	Construction of A157Q, I134P and Y257R 3D Models Using PyMol	58

4.2.2	Homology Modeling of Mutant Loop	59
4.3	Molecular Dynamics (MD) Simulation of CHI II and Mutants	64
4.4	Comparative Analysis of MD Simulation Trajectories	65
4.4.1	Root Mean Square Deviation (RMSD)	65
4.4.2	Root Mean Square Fluctuation (RMSF)	71
4.4.3	Radius of Gyration	75
4.4.4	Analysis of Secondary Structure for CHI II and Its Mutants	79
4.4.5	Solvent Accessible Surface Area (SASA)	84
4.4.6	Analysis of Hydrogen Bonds	85
4.4.7	Analysis of Salt Bridges	87
4.5	Cloning of CHI II and Its Mutants into <i>Pichia pastoris</i>	92
4.6	Expression of CHI II Mutants into <i>P. Pastoris</i> GS115	96
5	CONCLUSION	100
5.1	Conclusions	100
5.2	Recommendations	101
	REFERENCES	102
	APPENDIX A-F	119

LIST OF TABLES

TABLE NO	TITLE	PAGE
3.1	List of tools and software used in this study	36
4.1	Model evaluation of CHI II and mutant A157Q, I134P and Y257R	59
4.2	The alignment search result against PDB database from various servers	60
4.3	The best model generated by Modeller 9.10 and evaluations	61
4.4	Evaluation of mutant Loop model (model 84) before and after energy minimization (EM)	62
4.5	Average radius of gyration for CHI II and all mutants at 273 K, 288 K and 300 K.	78
4.6	The average values of total solvent accessible surface area of CHI II and all mutants at three temperatures	85
4.7	The average number of hydrogen bonds in CHI II and mutants at 273 K, 288 K and 300 K	87
4.8	The average number of salt bridges in CHI II and mutants at three temperatures.	88
4.9	The list of salt bridges that are conserved in CHI II and mutants at 273 K, 288 K and 300 K.	90

LIST OF FIGURES

FIGURE NO	TITLE	PAGE
2.1	Factors contributed to the structural flexibility of cold-adapted enzymes	8
2.2	The structures of amino acids arginine, proline and glycine	9
2.3	Dry chitin flakes and the arrangement of chitin polymer	16
2.4	The arrangement of chitin microfibrils in three polymorph forms	17
2.5	Illustration of cleavage pattern of chitinolytic enzymes	19
2.6	The classification of chitinase GH18 and GH19 subfamilies and subgroup	21
2.7	The domain organization of fungal chitinases in GH18 family	25
2.8	The predicted structure of CHI II by using threading method.	28
3.1	Illustration of operational framework of this study	34
3.2	Three stages of MD simulation consists of preparation, production and analysis stages	39
3.3	The overview of OE-PCR strategy to obtain a mutant's gene	43
4.1	The location of three salt bridges found in CHI I structures	51
4.2	Multiple sequence alignment showed the mutation site of Y257R	52
4.3	The structure alignment between CHI II and its template 1ITX	53
4.4	Multiple sequence alignment showed the mutation site of mutant Loop	55
4.5	Multiple sequence alignment showed the mutation site or I134P	56

4.6	Multiple sequence alignment showed the mutation site of A157Q	57
4.7	Ramachandran plot of mutant Loops model (a) before energy minimization and (b) after energy minimization	63
4.8	RMSD overlay of CHI II and four mutant at 273 K	67
4.9	RMSD overlay of CHI II and four mutants at 288 K	68
4.10	RMSD overlay of CHI II and four mutants at 300 K	69
4.11	An overview of CHI II and mutants structure snapshots in 5 ns interval at 300 K simulation	70
4.12	The overlay RMSF graph of CHI II and all mutants at 273 K	73
4.13	The overlay RMSF graph of CHI II and all mutants at 288 K	74
4.14	The overlay RMSF graph of CHI II and all mutants at 300 K	75
4.15	The overlay Rg graph of CHI II and all mutants at 273 K	76
4.16	The overlay Rg graph of CHI II and all mutants at (a) 288 K and (b) 300 K	77
4.17	Secondary structure assignment for CHI II and mutants at 273 K	81
4.18	Secondary structure assignment for CHI II and mutants at 288 K	82
4.19	Secondary structure assignment for CHI II and mutants at 300 K	83
4.20	The interaction maps between Arg57 and Asp205 which involved carboxylate group of Asp205 and guanidium group of Arg257.	91
4.21	The formation and breaking of salt bridge Asp205 and Arg257 which was introduced in Y257R at three temperatures.	91
4.22	PCR product of first PCR and second PCR reaction for all mutants.	93
4.23	Linearization of pPICZ α C and A157Q constructed vector	94
4.24	Formation of A157Q-GS115 colonies on YPD and YPDS plate (a) YPDS plate with 100 μ g/ml of zeocin, (b) YPD agar with 500 μ g/ml of zeocin, (c) YPD agar with 1000 μ g/ml of zeocin and (d) YPD agar with 2000 μ g/ml of zeocin	95

4.25	PCR colony for all mutant gene integration into <i>P. pastoris</i> GS115.	96
4.26	SDS-PAGE analysis of CHI II and mutant A157Q	97
4.27	SDS-PAGE analysis of mutant Loop, I134P and Y257R.	98

LIST OF SYMBOLS AND ABBREVIATIONS

%	-	percent
*	-	asterisk
.top	-	Topology file
~	-	About
>	-	Greater
≥	-	Greater or equal to
°	-	Degree
°C	-	degree celcius
½	-	Half
3D	-	Three dimensional
A	-	Alanine
Å	-	Angstrom
Arg or R	-	Arginine
Asn	-	Asparagine
Asp or D	-	Aspartic acid/aspartate
BD	-	Binding domain
<i>C. congregates</i>	-	<i>Coprinellus congregates</i>
<i>C. immitis</i>	-	<i>Coccidioides immitis</i>
CAZy	-	Carbohydrate Active Enzyme database
CBD	-	Chitin binding domain
CID	-	Chitinase insertion domain
Cl ⁻	-	Chloride ion
Cys	-	Cysteine
Cα	-	Carbon alpha

DNA	-	Deoxyribonucleic acid
DOPE	-	Discrete optimized protein energy
DSSP	-	Dictionary of secondary structure prediction
EC	-	enzyme commission
F	-	Phenylalanine
<i>G. antarctica</i>	-	<i>Glaciozyma Antarctica</i>
GH	-	Glycosyl hydrolase
GlcN	-	Glucosamine
GlcNAc	-	N-acetylglucosamine
Glu or E	-	Glutamic acid/glutamate
Glu or Q	-	Glutamine
Gly or G	-	Glycine
GPI	-	Glycosylphosphatidylinositol
H	-	Hydrogen
<i>H. atroviridis</i>	-	<i>Hypocrea atroviridis</i>
His or H	-	Hisitidne
ID	-	Identifier
Ile or I	-	Isoleucine
K	-	Kelvin
kDa	-	Kilodalton
kJ/mol	-	kilojoule per mol
L	-	Leucine
Lys or K	-	Lysine
LysM	-	Lysin motif
MD	-	Molecular dynamics
MSA	-	Multiple sequence alignment
N	-	Nitrogen
N		Asparagine
Na ⁺	-	Sodium ion
Nm	-	Nanometer
Ns	-	Nanoseconds
O	-	Oxygen
PDB	-	Protein data bank

pKa	-	Acid dissociation constant
PME	-	Particle mesh ewald
Pro or P	-	Proline
Rg	-	Radius of gyration
RMSD	-	Root mean square deviation
RMSF	-	Root mean square fluctuation
<i>S. cerevisiae</i>	-	<i>Saccharomyces cerevisiae</i>
SASA	-	Solvent accessible surface area
SCP	-	Single cell protein
Ser or S	-	Serine
sp.	-	Species
SPC	-	Simple point charge
T	-	Threonine
<i>T. atroviride</i>	-	<i>Trichoderma atroviride</i>
<i>T. aurantiacus</i>	-	<i>Thermoascus aurantiacus</i>
<i>T. lanuginosus</i>	-	<i>Thermomyces lanuginosus</i>
TIM	-	Triosephosphate isomerase
T _m	-	Midpoint temperature
Tyr or Y	-	Tyrosine
V	-	Valine
VMD	-	Visual molecular dynamic
α	-	Alpha
β	-	Beta
γ	-	Gamma
φ	-	Phi
ψ	-	Psi
μg/μl	-	Microgram per microliter
μl	-	Microliter
μmol	-	Micromole
BMGY	-	Buffered glycerol complex medium
BMMY	-	Buffered methanol complex medium
Bp	-	Base pair
EDTA	-	Ethylenediaminetetraacetic acid

LB	-	Luria Bertani
M	-	Molarity
ml	-	Milliliter
mM	-	Millimole
OE	-	Overlapping extension
<i>P. pastoris</i>	-	<i>Pichia pastoris</i>
PCR	-	Polymerase chain reaction
U	-	Unit activity
V	-	Voltage
v/v	-	Volume over volume
YPD	-	Yeast extract peptone dextrose
YPDS	-	Yeast extract peptone dextrose sorbitol
zeo ^R	-	Zeocin resistance

LIST OF APPENDICES

APPENDIX	TITLE	PAGE
A	Tutorial for MD Simulation	119
B	Routine and Media	123
C	Primer Sequences	128
D	pPICZ α C Vector Map	129
E	Nucleotide and Protein Sequences of CHI II and All mutants	130
F	Sequencing Result	132

CHAPTER 1

INTRODUCTION

1.1 Background of Study

Earth's surface is dominated by low temperature regions including polar region, mountains top and oceans which cover 80% of the biosphere where temperatures never exceed 5°C. Low temperatures are known to place severe physicochemical constraints on cellular function by negatively influencing cell integrity, water viscosity, solute diffusion rate, membrane fluidity, enzyme kinetics and macromolecular interactions (Marx *et al.*, 2004). However, despite most other species cannot grow at this low temperature regions, an extremophiles known as psychrophiles are able to survive and inhabit this region. Then the question arises: how can psychrophiles survive, let alone thrive under this harsh conditions? The answer is, psychrophiles evolved and adapted to their environments by developing unique mechanisms to keep their cellular components stable and active.

The ability of psychrophiles to survive in cold regions is therefore dependent on numbers of adaptive strategies to successfully counteract those low temperature constraints (D'Amico *et al.*, 2006). One of the strategies is by producing the cold-adapted enzymes that able to perform their catalysis efficiently under these extreme environmental conditions (D'Amico *et al.*, 2006). For these reasons, cold-adapted enzymes have been considered as biotechnological potential due to their ability to perform catalysis at low temperatures thus offering advantages in the

environmental application and energy savings in industrial processes (Gianese *et al.*, 2001).

While other enzymes are subject to cold denaturation and suffered the loss of activity at low temperatures, cold-adapted enzymes are resistant to cold denaturation with efficient catalytic activity. Their survival is correlated with their structural flexibility that was believed as a compensation for the freezing effect in cold habitats (Johns & Somero 2004). Structures flexibility of cold-adapted enzymes is the result of combination of several features such as increased numbers of hydrophobic side chains that are exposed to the solvent, a decrease in the compactness of hydrophobic core, a higher number of glycine and lysine residues, a reduced number of proline and arginine residues and weakening of intramolecular bonds (Rodrigues & Tiedje 2008). However, because of their structural flexibility, cold-adapted enzymes become less stable and also thermolabile which cause them to denature at elevated temperature (Siddiqui & Cavicchioli 2006).

Therefore, cold-adapted enzymes are often engineered either through rational design or directed evolution to improve its thermostability. Thermostability is defined as improved long-term survival under mild conditions and increased ability to remain active under harsh industrial condition but still retains its catalytic efficiency (Wijma *et al.*, 2013). In this study, a cold-adapted chitinase named as CHI II was used as the subject understudied. Chitinase (EC 3.2.2.14) are categorized under glycosyl hydrolases (GH) family and can be found in wide range of organisms such as bacteria, fungi, yeasts, plants and mammals. Capabilities of chitinase to hydrolyse chitin to a low molecular weight chitooligomers cause them to have broad potential in industrial, agricultural and medicinal functions (Dahiya *et al.*, 2006; Liu *et al.*, 2013; Patil *et al.*, 2000; Park & Kim 2010; Khan *et al.*, 2015).

CHI II was previously isolated from *Glaciozyma antarctica* P112 and its three-dimensional (3D) structures had been modeled by Ramli *et al.* (2011, 2012). Based on the structure analysis and primary sequence analysis, several characteristics related to cold adaptations were found in CHI II. CHI II was identified to have less number of salt bridges and arginine residues, increase in surface hydrophobicity and reduced

number of hydrogen bonds (Ramli *et al.* 2012). These characteristics were proved to be related to structural flexibility of CHI II which causes CHI II to be thermolabile and could not withstand elevated temperature and harsh environment. In addition to the wide potential of chitinases in industrial application and biotechnological application, it is best for CHI II to be engineered to improve their thermostability. Based on the information obtain from previous study on amino acid affecting thermostability of cold adapted chitinase and based on comparison studies between mesophilic, psychrophilic and thermophilic enzymes, rational design was used to design CHI II mutants (Mavromatis *et al.*, 2003; Siddiqui & Cavicchioli 2006). Therefore, four mutants of CHI II will be designed through rational design and the effect of the mutation will be studied using an *in-silico* approach. In particular, the mutant's structure stability will be studied through molecular dynamic (MD) simulation at three temperatures: 273 K, 288 K and 300 K. This is an indicator of CHI II mutants performance as it reflects the ability of the mutants to perform under conditions relevant to an industrial process where enzyme is continually affected by temperature elevation.

1.2 Problem Statement

About 80 000 metric tons biomass waste of marine invertebrate were produced every year and it was predicted that the oceans will be depleted of chitin if this insoluble biomass is not converted into simple and recyclable material (Patil *et al.*, 2000). Capabilities of cold-adapted chitinases to have high catalytic efficiency and high flexibility (low stability) at low temperatures allows them to offer several novel opportunities in industrial application. Because of their inherent flexible structure, cold-adapted chitinase was correlated to be thermolabile as their reaction rates decrease when the temperature increases. Hence, this condition becomes a limitation for cold-adapted chitinase to be used in industrial application. Thus, production of cold-adapted enzymes chitinase with desired thermostability become an important aspect of industrial application which could also help to overcome chitin depletion. This can be achieved through mutagenesis of cold-adapted chitinase to improve its thermostability without compromising its structurally dependent cold-adapted properties (Cesarini *et al.*, 2012).

1.3 Objectives

The main objective of this study is to analyse the effect of amino acids substitution, loop shortening and introduction of the salt bridge in the non-catalytic region on CHI II thermostability through *in-silico* approach.

1.4 Scopes of Study

The scope of this study are:

- a) Design four CHI II mutants through rational design.
- b) Construction of four mutants three-dimensional (3D) structures using mutagenesis plugin in PyMOL and homology modeling by using Modeller.
- c) Performing the Molecular Dynamics (MD) simulation of CHI II and its four mutants at three different temperatures; 273 K, 288 K and 300 K.
- d) Performing comparative trajectories analysis on CHI II and its mutants to study the effect of mutation on CHI II thermostability.

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APPENDIX A

APPENDIX A: TUTORIAL FOR MD SIMULATION

1) Process the pdb file with `pdb2gmx`

```
pdb2gmx -ignh -f inputabf.pdb -o abf.pdb -p abf.top -water spce
```

`Pdb2gmx` command converts the `pdb` file to a `gromacs` file and write the topology.

Points to ponder:

- What is the total mass of your protein?
- What is the total charge of your protein?
- Open the topology file (`abf.top`) using `Gedit`, see how the force define protein

2) Set-up box for simulation

```
editconf -bt cubic -f abf.pdb -o abf_bsolv.pdb -d 2.0
```

`Editconf` specify the simulation box. ‘-d’ sets the dimension of the box 2.0 nm (20A).

It should at no less than 0.9 nm for most system.

3) Solvate the box

```
genbox -cp abf_bsolv.pdb -cs spc216.gro -o abf_bion.pdb -p abf.top
```

Points to ponder:

- Does the size of your box change after the solvation?
- How many SOL molecules were added into your simulation box?
- Any change to your topology file?

4) Neutralize the protein system

```
grompp -f ion.mdp -c abf_bion.pdb -p abf.top -o ion.tpr -maxwarn 5
```

```
genion -s ion.tpr -o abf_b4em.pdb -nname CL (-pname NA) -nn 2 (-np 2) -p abf.top  
-g ion.log
```

Type “13” for SOL

Points to ponder:

- How many NA and CL had been added into the system?
- Is there any charge in the topology file?
- What do the flags used along the genion?

5) Energy Minimization

```
grompp -f em.mdp -c abf_b4em.pdb -p abf.top -o em.tpr -maxwarn 5
```

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

Points to ponder:

- What do `-v` and `-deffnm` mean?
- How many steps does the system take to converge?
- How many output files are there?

The output will be `em.gro`. If the F_{\max} did not converge, repeat the step by changing the input and output file (`em.mdp` file did not change). Take output from first step (`em.gro`) as an input for second step (after `-c` command). The second step output will be `em2.tpr` (after `-o` command). Run the simulation by naming `em2` at `mdrun` step.

6) Position Restrained Molecular Dynamics (equilibration)

```
grompp -f pr.mdp -c em.gro -p abf.top -o pr.tpr -maxwarn 5
```

Once the `pr.tpr` is generated successfully, run the position restrained MD

```
mdrun -v -deffnm pr
```

Points to ponder:

- Is there any note/warning when `grompp` is pre-processing the `pr.mdp`?

7) Convert Gromacs File

```
editconf -f file.gro -o file.pdb
```

8) First Evaluation

- First evaluate the system and see the water molecules had been equilibrated or not
- Compute the RMSD of the protein backbone and plot several graphs


```
g_rms -f pr.trr -s pr.tpr -o rmsd_pr
```

 - Examine using GRACE. `xmgrace rmsd_pr.svg`
 - For least square fit and RMSD calculation, select group 4 (Backbone)

- The program will generate a plot for RMSD over time
- c) Examine the temperature:
 - g_energy -f pr.edr -o temperature_pr*
 - Select '14' (Temperature)
 - Examine using GRACE. *xmgrace temperature_pr.xvg*
- d) Use *g_energy* to plot *density_pr.xvg* and *pressure_pr.xvg*, use *xmgrace* command to plot the graph.
- e) System had been equilibrated and may proceed to the production stage when:
 - a. The temperature plot stabilized/constant at 300K
 - b. The average reading for *density_pr.xvg* and *pressure_pr.xvg* are 1000 kg/m³ and 1.05 bar respectively

9) Production Stage

```
grompp -f md.mdp -c pr.gro -p abf.top -o md.tpr -maxwarn 5
mdrun -v -deffnm md
```

10) Trajectories analysis

Time evolving coordinates of a system are called trajectories. Trajectory files (*.trr) are normally binary files that contain several sets of coordinates for the system.

- a) Compress the trajectory
 - trjconv -f md.trr -s md.tpr -o md.xtc -pbc nojump*
- b) Analyse the energy output (same for potential energy, kinetic energy and total energy)
 - g_energy -f md.edr -o xxx* and plot *xmgrace -nxy xxx.xvg*,
- c) Measure radius of gyration and select '4' (Backbone).
 - g_gyrate -f md.trr -s md.tpr -o abf_gyrate.xvg*
- d) Measure RMSD of the structure by and select '4' (Backbone).
 - g_rms -s md.tpr -f md.trr -dt 10 -o md_rmsd.xvg*
- e) Compare RMSD to the NMR structure and select '4' (Backbone)
 - g_rms -s em.tpr -f md.trr -o abf_rmsd.xvg*
- f) RMS fluctuation of atom positions and select '3' (C-alpha).
 - g_rmsf -s md.tpr -f md.trr -b 200 -e 1000 -o abf_rmsf*
- g) RMSF to compute average structures and select '1' (Protein).

g_rmsf -s md.tpr -f md.trr -b 800 -e 1000 -o abf_xvg.pdb

- h) Analyse the secondary structure of model by and select '1' (Protein).

do_dssp -s md.tpr -f md.trr -o abf_ss.xpm -dt 10