# Conformational sampling of enzyme dynamics: Triosephosphate Isomerase

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Referee: Prof. Dr. Kai Tittmann Co-referee: Prof. Dr. Burkhard Morgenstern Date of the oral examination: 17.08.2012 To my parents, my dear brother, friends, and family.

"We never are right, we can only be sure we are wrong" R.P. Feynman

## **List of Abbreviations**

TIM	Triosephosphate Isomerase
CTIM	Closed TIM
OTIM	Open TIM
CS	Crystal simulations
APO	Apo TIM
CS-O	CS with OTIM
CS-C	CS with CTIM
CS-2PG	2PG ligated CS with CTIM
GAFF	General Amber forcefield
GAP	Glyceraldehyde-3-Phosphate
DHAP	Dihydroxyacetone Phosphate
EDT1	Enediolate-1
EDT2	Enediolate-2
2PG	2-Phosphoglycerate
G3P	Glyecerol-3-phosphate
PGH	Phosphoglycolohydroxamate
IPP	2-(N-Formyl-N-hydroxy)-amino-ethylphosphonate
RT	Residence time
RMSD	Root mean square deviation
PCA	Principal Component analysis
MD	Molecular dynamics
QM	Quantum mechanics
HF	Hartree-Fock
MgCl <sub>2</sub>	Magnesium chloride
NaCl	Sodium chloride
PME	Particle-mesh Ewald
NH	Nośe-Hoover
PR	Parrinello-Rahman

LJ	Lennard-Jones
LINCS	Linear constraint solver
NPT	N=Number of particles; P=Pressure; T=Temperature
NPT	N=Number of particles; V=Volume; T=Temperature
NMR	Nuclear Magnetic Resonance
SSNMR	Solid state NMR
US	Umbrella sampling
k <sub>ex</sub>	Exchange rate between open and closed conformations
k <sub>cat</sub>	Catalytic rate
k <sub>m</sub>	Michaelis constant
fs	femtoseconds
ns	nanosecond
ps	picosecond
μs	microsecond
$\Delta G$	Free energy difference

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## Prologue

Proteins are fascinating biomolecules which carry out wide variety of functions in various biological processes. One among many of these functions is enzyme catalysis. A protein which acts as a catalyst in a biochemical reaction is called an enzyme. During this process of enzyme catalysis, an enzyme may (Triosephosphate Isomerase, RNase A, Adenylate kinase, Cyclophilin-A, Monoclonal antibody SPE7) or may not undergo a structural change (Carbonic anhydrase)<sup>1,2</sup>.

Enzymes are highly specific with respect to a substrate and accelerate the rate of a reaction compared to the rate in the solution. It is believed that enzymes achieve this high acceleration rate by lowering the free energy of the activation i.e. the transition state is more stable in the enzyme compared to the solution. Role of dynamics in enzyme catalysis has been subject of serious debate and its influence on the actual mechanism has not been understood that well<sup>1-4</sup>. But any mutation in the enzyme structure which limits the conformations that can be sampled by the enzyme, may lead to a disruption in the actual reaction mechanism the enzyme is proficient in.

The general opinion is that enzymes adopt multiple conformations to facilitate the reaction they catalyze<sup>1,2,4</sup>. There are two models discussing the conditions of a conformational change in an enzyme. *Induced fit* model describes that an enzyme exists predominantly in *non-catalytic* conformation and undergoes a conformational change from *non-catalytic* to *catalytic conformation* only in the presence of the specific substrate<sup>5</sup>. Whereas conformational selection suggests that an enzyme samples both *catalytic* and *non-catalytic* conformations even in the absence of the substrate and binding of the substrate preferentially increases the population of the *catalytic* conformation<sup>6,7</sup>. To better understand how an enzyme works, it is very important to identify the various substates that an enzyme.

Our study is aimed at understanding the structure-dynamics-function relationship of Triosephosphate Isomerase, one such enzyme which undergoes conformational changes during the enzyme catalysis, using Molecular dynamics (MD) simulations. Since 1958 various scientific studies on TIM have elucidated structure<sup>8-16</sup>, reaction mechanism<sup>17-34</sup>, loop-6 transition between open and closed state <sup>12,30,35-41</sup> and the factors affecting or influencing

the transition<sup>39-42</sup>. These studies have provided glimpses of individual events of what essentially is a dynamic process.

The different conformations of TIM suggest that for each task i.e. substrate binding, catalysis and product release, the protein adopts to a specific conformation suited for that particular task. We are interested in studying the sequence of events from the binding of the ligand to the release of the product by means of various MD simulation techniques.

## **1** Introduction

Glycolysis involves conversion of six carbon compound glucose, to two molecules of three carbon compound pyruvate, harvesting two molecules of Adenosine triphosphate (ATP). The fourth reaction of this metabolic pathway involves conversion of Fructose 1, 6-Bisphosphate to a ketone compound Dihydroxyacetone Phosphate (DHAP) and an aldehyde compound Glyceraldehyde 3-Phosphate (GAP). GAP proceeds further to form one pyruvate molecule, whereas DHAP is converted to GAP by TIM in the fifth step of glycolysis, which in turn forms the second pyruvate. Interconversion of DHAP to GAP and GAP to DHAP is catalyzed by TIM in both the directions<sup>43</sup> (reaction mechanism discussed in section 1.4) at a catalytic rate ( $k_{cat}$ ) of 10<sup>3</sup> s<sup>-1</sup> and 10<sup>4</sup> s<sup>-1</sup> respectively.

## 1.1 TIM structure

TIM was first crystallized by Phillips et al. in 1975<sup>8</sup>. TIM is active as a dimer (Figure 1.1) in most of the organisms but tetramers were also reported in thermophillic organisms<sup>11,13,15,43</sup>. Mutations in the dimer interface lead to the formation of the monomers which exhibit a 1000 fold lower catalytic rate when compared with the wild type dimer<sup>44</sup>. Hence to counter diseases like malaria and African sleeping sickness various studies have concentrated on developing an inhibitor which binds at the dimer interface and disrupts the quaternary structure<sup>45-51</sup>. Each monomer is composed of approximately 250 amino acids. The tertiary structure of TIM is composed of 8  $\alpha$  and 8 parallel  $\beta$  sheets interconnected by loops (Figure 1.1). This ( $\beta\alpha$ )<sub>8</sub> fold also known as TIM-barrel fold, is shared by more than 10% of known enzyme structures<sup>52</sup>.



Figure 1.1: Structure of TIM dimer with DHAP in the active site. Loop-6 (red), loop-7 (green) and loop-8 (orange) envelope the active site. Loop-5 (purple) interacts with N-terminus residues of loop-6. In inset residues of the active site which are highly conserved.

The active site of TIM is enveloped by three distinct loops (Figure 1.1) loop-6, loop-7, loop-8. Residues H94, E166 are the proposed catalytic residues involved in the proton shuttling between C1 and C2 carbon atoms of the substrates. G172 (loop-6), S212 (loop-7), G233 and G234 (loop-8) provide hydrogen bonds to the phosphate moiety of the substrate and stabilize it (Figure 1.1). N10 and K12 interact electrostatically with the substrate (Figure 1.1). Residues that are involved in catalysis and that are interacting by means of hydrogen bonds or coulombic interactions are conserved throughout all the sequences<sup>43</sup>.



Figure 1.2: a) Fit of OTIM (green) structure on CTIM (blue). Major structural difference is seen at loop-6 (open loop-6 in red). In inset open (green) and closed (blue) loop-5. b) N-terminus, C-terminus and tip of open loop-6. In our simulations, to monitor hinge regions dynamics W169-S212 and V176-S212 c-alpha (green spheres) distances are monitored.

Loop-6 N-terminus (167-PVWA-170), tip (171-IGTG-174) residues are much more conserved than the C-terminus (175-KVA-177) residues (Figure 1.2b). Wang et al.<sup>41</sup> reported that there is an evolutionary link between the sequence of N-terminus segment loop-6 (PVW) and loop-7. The typical loop-6 N-terminus sequence is '167PVW' with a corresponding loop-7 sequence '209YGGS'. But in archaebacteria loop-6 N-terminus sequence is '167PPE' and typical loop-7 sequence being '209TGAG/209CGAG'. A substitution of archaebacterial

loop-7 sequence into chicken TIM, with unmodified loop-6, leads to an impaired enzymatic activity<sup>41</sup>.

There are three structural differences between open TIM (OTIM: 5TIM-A) and closed TIM (CTIM: 6TIM-B) structures (Figure 1.2a). The major structural difference between OTIM and CTIM is seen in loop-6. Backbone RMSD of 5TIM-A with respect to 6TIM-B excluding loop-6 is 0.034nm. The second structural difference is seen in loop-7 where three residues undergo a change in backbone phi ( $\Phi$ ; backbone atoms C'-N-C $\alpha$ -C''), psi ( $\Psi$ ; backbone atoms N-C $\alpha$ -C-N'') dihedral angles between OTIM and CTIM (Figure 1.3c). There is also a minute structural change seen in loop-5 between OTIM and CTIM (Figure 1.2a). Carboxylate moiety of E128 from loop-5 forms a hydrogen bond with epsilon hydrogen of loop-6 W169 in closed state (Figure 1.4a). This hydrogen bond is absent in open loop-6. Excluding loop-6 and loop-7, rest of the protein is highly identical between OTIM and CTIM. The role of loop-6, loop-7 and residues involved in the enzyme catalysis are further discussed below.

## 1.2 Loop-6

Loop-6 is composed of 11 aminoacids (Consensus sequence: "PVWAIGTGKTA"). N-terminus of loop-6 ('PVW') is rich in hydrophobic aminoacids whereas tip ("AIGTG") of loop-6 and C-terminus ("KTA") are hydrophilic. N-terminus of loop-6 interacts with the loop-5 whereas tip of the loop-6 and C-terminus are solvent exposed. Loop-6 acts as a lid on the active site. Depending upon the orientation of loop-6 upon the active site, TIM has two distinct conformations; open and closed (Figure 1.3). We have used distance between residues W169-S212 and V176-S212 as a criterion to distinguish between OTIM and CTIM (Figure 1.3 a, b). The reasons for this will be further discussed in section 1.5.



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Figure 1.3: Closed (a) and open (b) conformations of TIM based on distance between W169-S212 (Closed: 0.95nm, Open: 1.16nm) and V176-S212 (Closed: 0.51nm, Open: 0.87nm). Distance to loop-6 (red) and loop-7(red). c) Loop-7 in open (green) and closed conformation (blue). N-terminus distance is shown in blue and C-terminus in red.

A mutation study by Pompliano et al.<sup>20</sup> where loop-6 residues '171IGTG' were deleted, suggested that closure of loop-6 during the enzyme catalysis is assumed to stabilize the enediol phosphate intermediate as well as the transition state<sup>20</sup>. In the mutant TIM they found a six fold higher formation of methylglyoxal and inorganic phosphate compared to the wild type enzyme<sup>20</sup>. The so formed methylglyoxal is toxic to the cell<sup>53</sup>.

#### 1.2.1 NMR studies on loop-6

As stated earlier, dynamic nature of loop-6 has been investigated by several scientific studies. W169 present in the N-terminus of loop-6 was used as an indicator of the loop-6 conformation in the studies conducted by Ann McDermott group<sup>35-38</sup>. There is change in the environment of the W169 between the two conformations. In open loop-6, epsilon hydrogen of W169 is involved in a hydrogen bond with hydroxyl group of Y165. In closed loop-6, epsilon hydrogen of W169 forms hydrogen bond with carboxylate moiety of E128 (Figure 1.4). Upon closure of loop-6 W169 swings by 30°-40° (Figure 1.4c). Patrick Loria group<sup>39,41</sup> in their solution NMR relaxation dispersion experiments monitored the dynamics of loop-6 N-terminus V168 and C-terminus K175, T178 residues.



Figure 1.4: a, b) W169 forms hydrogen bond with E128 in CTIM and with Y165 in OTIM.

Solid state NMR (SSNMR) study by Williams et al.<sup>35</sup> in 1995 of apo and Glycerol-3-phosphate (G3P) ligated TIM reported that loop-6 motion is not ligand gated and occurs at a

rate of  $10^4$  s<sup>-1</sup>. In SSNMR, the obtained deuterium line shapes are simulated with ligand concentration, W169 angle and loop-6 populations as parameters. Based on the crystallographic data Williams et al.<sup>35</sup> assumed that in their NMR experiments, in apo state open loop-6 is the major conformation and in the ligated TIM closed loop-6 is the major conformation. Using the exchange rates and population ratios obtained from the simulated line shapes, barrier heights and free energy difference between open and closed conformations of apo TIM and G3P, 2PG (2-Phosphoglycerate) ligated TIM was calculated (Figure 1.5). The reported free energy difference between the major conformation and the minor conformation ranges between 1.2 kcal.mol<sup>-1</sup> and 1.8 kcal.mol<sup>-1</sup>. If the binding free energy between open and CTIM's, then it would be expected that in apo state, OTIM is favoured by -1.8 kcal.mol<sup>-1</sup> and in ligated TIM with G3P/2PG closed state is favoured by -4.5/-6.8 kcal.mol<sup>-1</sup> with respect to apo OTIM<sup>35</sup>.



Figure 1.5: Estimates of free energy difference between open and CTIM based on data from Williams et al.<sup>35</sup>. For G3P and 2PG the binding free energies were calculated from the binding constant and were added to the free energy estimates from the line shape simulations.

T-jump relaxation studies by Desamero et.al<sup>38</sup> in 2002 and solid, solution state NMR studies by Rozovsky et al.<sup>36</sup> in 2003, reported that loop-6 opening rate in presence of G3P is of the

same order as the catalytic rate of TIM which is  $10^4 \text{ s}^{-1}$ . Desamero et.al<sup>38</sup> further suggested that loop-6 opening rate depends upon the ligand present in the active site. It has to be noted that the opening and closing rates reported by the above studies are the k<sub>close</sub> and k<sub>open</sub> rates (Figure 1.6). It was difficult to measure the k<sub>ex</sub> in apo TIM sample using fluorinated W168 in solution state NMR studies because no change in chemical shift is observed and Rozovsky et al.<sup>36</sup> suggested that it might be because of low population of closed loop-6. It is speculated though that in apo state loop-6 motion is in the regime of  $10^5 \text{ s}^{-1} - 10^6 \text{ s}^{-1.37}$ . k<sub>on</sub> and k<sub>off</sub> rates were proposed to be  $10^6 \text{ s}^{-1}$ .<sup>37</sup>



Figure 1.6: Scheme of kinetic states used in NMR experiments by Rozovsky et al.<sup>36,37</sup>.

By mutational studies and using TROSY NMR spin relaxation experiments, Berlow et al.<sup>39</sup> has shown that N and C-terminal hinges of loop-6 are coupled. Y209F mutation in loop-7 destabilizes the closed conformation of loop-6 and increases the population of open state because of the loss of hydrogen bond between C-terminus loop-6 residue A176 and Y209. The effect of this mutation is seen in the dynamics of both N and C-terminus hinges. Surprisingly they found different populations distributions for N and C-terminal hinges, which might imply that N and C-terminus hinges of loop-6 are uncoupled and they suggest that more experiments are required to confirm this "novel" view of loop-6 motion<sup>39</sup>.

Further NMR studies by Wang et al.<sup>41</sup> where chicken loop-7 (209YGGS) was replaced by archaeal loop-7 sequence (209TGAG), reported a 240 fold loss of activity for TIM. The affect of this loop-7 mutation on the exchange rate ( $k_{ex}$ ) between open and closed conformations of loop-6 for V167 and T177 residues was identical (wild type: 9000s<sup>-1</sup>; loop-7 mutant: 18000s<sup>-1</sup>) and based on this data they suggested that in the absence of the proper corresponding loop-7 sequence, concerted motion between the N and C-terminal hinges of loop-6 is affected. Wang et al.<sup>41</sup> also calculated the height of the activation barriers for loop-6 closure for V168 and T177 in wild type enzyme (V167: 13.7±1.8 kcal.mol<sup>-1</sup>; T177: 15.7±0.98 kcal.mol<sup>-1</sup>) and as well as the loop-7 mutant (V167: 1.4±0.38 kcal.mol<sup>-1</sup>; T177: 6.26±2.6 kcal.mol<sup>-1</sup>)<sup>41</sup>.

The barrier heights are calculated by measuring the change in contribution of chemical exchange process to the relaxation rate ( $R_{ex}$ ) by varying temperature. Though the effect on

the obtained  $k_{ex}$  rate in the loop-7 mutant was identical ( $k_{ex}$  wild type: 9000 s<sup>-1</sup>, loop-7 mutant: 18,000 s<sup>-1</sup>) on N and C-terminus hinges, the height of the activation barriers was different and Wang et al.<sup>41</sup> pointed out that this difference may be either because of difference in population rations for N and C-terminus hinges in open and closed states or the effect of temperature on the population of V167 and T177 in open and closed states is different in the mutant<sup>41</sup>.

#### **1.2.2** Theoretical studies on loop-6

There were also several theoretical studies conducted to study loop-6's dynamics. Joseph et al.<sup>54</sup> in 1990 were the first to perform a molecular dynamics (MD) simulation of TIM and they reported that closure of loop-6 starts after binding of substrate which initiates displacement of E166 into swung-in conformation followed by closure of loop-6. These simulations were 100 ps long and only selected part of the proteins were allowed to move<sup>54</sup>.

Later in 1993 Wade et al.<sup>55</sup> performed a Brownian dynamics (BD) simulation on TIM where the entire protein was restrained excluding the loop-6 was modelled as a linear chain and each residue of loop-6 was represented by a sphere. These simulations lasted for 70 ns to 100 ns and suggested that loop-6 acts as a gate on the active site and does not influence the rate of diffusion of ligands into the active site.

In 1998 Derreumaux et al.<sup>56</sup> using high temperature langevin dynamics simulation demonstrated that the hydrogen bond between Y209 and A176 is the last hydrogen bond to break for opening of loop-6 and like previous studies confirmed that loop-6 motion is a rigid body motion<sup>56</sup>. A combined stochastic boundary MD (SBMD) simulation and solution NMR of Massi et al.<sup>57</sup> reported that loop-6 moves as a rigid body. The SBMD approach involves simulating only a particular region of interest of protein which in this case was all residues within 1.6 nm of A176 and this system was simulated for 10 ns<sup>57</sup>.

## 1.3 Loop-7

Loop-7 (Figure 1.7a) is located above the active site and is composed of six amino acids ('YGGSVN'). Phi ( $\Phi$ ) and psi ( $\Psi$ ) backbone dihedral angles of loop-7 residues G210, G211 and S212 differ between open and CTIM structures (Figure 1.7). It was suggested that the flip of G210 $\Psi$  into closed conformation induces conformational change of E166 from 'swung-out' to 'swung-in' reactive conformation<sup>43</sup>. S212 amide group is involved in a hydrogen bond formation with the phosphate moiety of the bound substrate.

Y209 of loop-7 is involved in two hydrogen bonds, one with A177 of loop-6 (Figure 1.7b) stabilizing the closed state of loop-6. The second hydrogen bond is formed between the hydroxyl group of Y209 and carbonyl oxygen of G211 (Figure 1.7b). This interaction stabilizes the G211 of loop-7 in the closed state<sup>10</sup>. Mutagenesis<sup>42,58</sup> and NMR studies<sup>39</sup> of Y209F mutant suggested that loss of hydrogen bond between the Y209 and A177 results in an increased population of the open state with a 2000 fold decrease in the catalytic activity. Loop-7 as stated above in section 1.2 was suggested to bring about concerted motions between N and C-terminus hinges.



Figure 1.7: a) Loop-7 open (green) and closed (blue) conformations. b) Hydrogen bonds formed by Y209 with G211 of loop-7 and A177 of loop-6 in closed state.

## **1.4 Catalytic mechanism**

Residues K12, H94 and E166 are involved in the catalysis of interconversion of DHAP to GAP and vice versa. To explain the reaction mechanism, many experimental<sup>17-20,22,25,26,31-34</sup> and computational<sup>21,23,24,27-30</sup> studies were carried out and two possible mechanisms were proposed. The classical and the criss-cross mechanism (Figure 1.8). In both the mechanisms the first step is abstraction of the C1-R proton in DHAP and C2-R proton with GAP as substrate by the catalytic base E166. This leads to the formation of the intermediary cisenediol transition state<sup>59</sup>.

In the classical mechanism from the second step onwards H94 is thought to play a role in shuttling of protons. Whereas in the criss-cross mechanism, E166 can slide over the cisenediolate plane between the C1 and C2 carbons and thereby shuttle the protons over the carbons. The last step in this process is reprotonation of the enediolate-2 intermediate by E166 leading to product formation.



Figure 1.8: Classical and criss-cross reaction mechanism pathways of TIM catalyzed isomerase reaction from DHAP to GAP. Image taken from Cui.et al.<sup>27</sup>.

Stabilizing the transition state by closure of loop-6 is assumed to be key for TIM's catalytic power of  $10^9 \text{ M}^{-1}\text{s}^{-1}$  when compared with the same reaction in solution<sup>43</sup>. As stated earlier

deletion of loop-6 residues '171IGTG' leads to a six fold formation of methylglyoxal and inorganic phosphate compared to the wild type enzyme<sup>20</sup>. Natural substrate and the transition state are stabilized in the active site by series of hydrogen bonds and electrostatic interactions with N10 and K12. Closure of loop-6 excludes water from the active site lowering down the dielectric constant of the active site. This lower dielectric constant in the active site pocket is thought to enhance the columbic interactions between the charged residues N10, K12 with the substrate and as well as the transition state<sup>60</sup>. Kursula et al.<sup>61</sup> proposed that closure of loop-6 sandwiches the catalytic residue E166 between the hydrogen on C1 carbon of DHAP, when compared with the same reaction in solution<sup>61</sup>.

The hydrogens of the carboxamide group of N10 forms hydrogen bonds with hydroxyl oxygen on the first carbon of DHAP (Figure 1.1). K12 is in the proximity of forming hydrogen bonds with the ketonic oxygen O2 as well as the oxygens of the phosphate moiety. The hydrogen of the hydroxyl group on the first carbon forms a hydrogen bond with the carboxylate oxygen (OE1/OE2) of E166 (Figure 1.1). The amide hydrogen's of G172, S212, G233 and G234 can form hydrogen bonds with the oxygen's of the phosphate moiety (Figure 1.1).

### **1.5** Aim of this study

There is a widespread interest in designing drugs to inhibit TIM, to counter malaria and African sleeping sickness<sup>45-51</sup> and also designing novel enzymes based on TIM's scaffold <sup>44,62-64</sup>. Prior to that, it is important to understand step by step process of the conformational changes of loop-6 and loop-7 and the factors that drive these conformational changes. NMR and fluorescence spectroscopic studies on loop-6 have revealed insight into the exchange rates ( $k_{ex}$ ), between encounter complex OTIM and ligated (G3P/2PG) CTIM. The  $k_{ex}$  for apo TIM between open and closed conformations is difficult to measure by experiments because of skewed populations of loop-6 in closed conformation<sup>36</sup>. Previous computational studies did not simulate the entire protein and spanned only 100 ns, which is not enough to capture a complete picture of such a slow (microsecond to millisecond) conformational change process.

Atomistic microsecond MD simulations of entire protein with and without ligand will help us to identify various conformational changes in TIM at atomic detail. Our study is aimed at addressing the following questions on TIM.

- What is the nature of dynamics of loop-6? Is it a rigid body motion or does N and C-terminus hinges of loop-6 move in uncorrelated fashion?
- What factors contribute to motion of loop-6? Is closed conformation sampled in apo TIM? Or is the closed conformation of loop-6 ligand induced?
- How does loop-7 influence dynamics of loop-6? Does it really bring about a concerted motion of loop-6?
- What is the nature of loop-7 conformational change? Influence of loop-7 dynamics on loop-6 has been studied but process of loop-7 conformational was not studied till now.
- Are there any other factors that influence the loop-6 and loop-7 conformational change?
- What is the free energy difference (ΔG) between open and closed conformations of loop-6 and loop-7 and what is the effect of various ligands on the ΔG?
- TIM catalyzes the reaction in both the directions i.e. binding of GAP to active site will form DHAP and vice versa. And an important question here is how does the product formed at the end of the reaction released, since the product formed at the end of previous reaction is the substrate for the next reaction?
- What changes happen in the active site upon the binding of ligand?

## 2 Methodology

## 2.1 Molecular Dynamics

Molecular dynamics (MD) is very useful tool to study the evolution of a system's dynamics. The first ever molecular dynamics simulation of a protein was reported in 1977 by McCammon et al.<sup>65</sup> and the first simulation of a protein with solvent was of bovine pancreatic trypsin inhibitor by Gunsteren et al.<sup>66</sup>. MD simulations have been used to study complicated phenomenon like photochemistry of proteins<sup>67,68</sup>, conformational changes in proteins<sup>69,70</sup>, water and ion permeation through channels<sup>71</sup>, and protein folding<sup>72,73</sup>.

The first important assumption in molecular dynamics is the Born-Oppenheimer approximation<sup>74</sup>, which states that electrons being lighter than nuclei adopt to change in nuclear positions instantly and based on this, the interaction of nuclei can be used to describe the physico-chemical properties of the system of interest<sup>75</sup>. To study the dynamics of a complex system like protein the interaction between various atoms are described using a molecular mechanics forcefield. A molecular mechanics forcefield describes the interaction between various atoms using set of bonded and non bonded interactions modeled by analytical expressions (Figure 2.1). There are many flavors of available forcefields, most commonly used ones are CHARMM <sup>76</sup>, AMBER <sup>77</sup>, GROMOS<sup>78</sup>, OPLS<sup>79</sup>.

The forcefield parameters for describing the bonded and non bonded interactions are obtained from either quantum chemical calculations or by fitting to experimental data from experimental techniques like NMR, X-ray, neutron diffraction, infrared and Raman spectroscopy<sup>80</sup>. Each atom is represented by a sphere of mass 'm' and a partial charge 'q'. The sum over all the bonded and non bonded interactions of all the atoms in the system constitutes the potential energy (*V*) of the system (Eq 1.1).

$$V = V_{bond} + V_{angle} + V_{dihedral} + V_{vdw} + V_{coulomb}$$
1.1.

. .



Figure 2.1: Bonded and non-bonded interactions used to describe interactions between atoms in a forcefield.  $k_{ij}$ ,  $k_{\theta}$  are the force constants of the respective bond, angle bending potential and torsional potential.  $r_{ij}$  is the distance between two atoms *i* and *j*.  $\theta_{ijk}$  is the angle between three atoms and the dihedral angles between atoms *i*, *j*, *k* and *l* is represented by  $\theta_{ijkl}$ .  $r_0$  and  $\theta_0$  are the reference bond and angle values.  $q_i$  and  $q_j$  are the charges on atoms *i* and *j*.  $\varepsilon_0$  is the permittivity constant.

The force acting on each atom at each step in a MD simulation is the derivative of the potential energy with respect to the atomic position in x, y and z directions (equation 1.2). These forces when combined with the current positions and velocities are used to generate new positions<sup>75</sup>.

$$F_i = \frac{dV}{dx_i}$$
 1.2.

Using very short time steps of one or few femtoseconds (fs), equations of motion based on Newton's second law are integrated using Verlet leap-frog algorithm<sup>81</sup>. Since the time steps used for numerical integration are very short, the force acting on each atom is assumed to be constant during the time interval of generating new positions. Atoms are then moved to a new position; and an updated set of forces is computed and the process is repeated. In this way trajectories are generated and from these trajectories we can calculate the properties of interest like free energies, diffusion constants of solvent molecules, NMR chemical shifts etc.

The current state of the art computer clusters enable us to run microseconds ( $\mu$ s) to milliseconds (ms) of MD simulations<sup>75</sup>.

#### 2.1.1 Principal component analysis (PCA)

Proteins undergo complicated conformational changes and it is difficult to understand and identify the conformational changes of a protein by looking at a molecular dynamics trajectory. Amadei et.al<sup>82</sup> in 1993 developed PCA where a covariance matrix between all atoms is calculated and from the diagonalization of this matrix we can obtain eigenvectors. An eigenvector is a vector which describes a particular motion in a protein and the corresponding eigenvalue of each eigenvector describes the amplitude of the motion. Eigenvectors are sorted in descending order based on their eigenvalues. Approximately 90% motion of the protein can usually described by using the first 20 eigenvectors depending on the nature of the motion<sup>82</sup>.

#### 2.1.2 Umbrella sampling simulations

It is not only important to understand the dynamic nature of the motions we see in a protein but also to understand the energetics of the process. Depending upon the biomolecular system under investigation and the nature of conformational changes at times even with world class computer facilities it is very difficult if not impossible to see enough transitions between the various states populated by the system.

Some states which are separated by high energy barriers may or may not be accessible in the MD simulations. Even if accessible there may or may not be statistical significant enough transitions to calculate the free energy difference ( $\Delta G$ ) between the two states of interest A and B using the probabilities of system in state A and state B using equation 1.3.

$$\Delta G = -k_B T ln \frac{p_A}{p_B}$$
 1.3.

In a scenario where sampling of a particular state of interest is not sampled, one has to employ enhanced sampling methods like Umbrella sampling (US). In US one defines a reaction coordinate, which can be an eigenvector from PCA or a dihedral angle or angle or distance between two atoms and restricts the system along this reaction coordinate using a series of harmonic potentials. The harmonic potentials which have been used to restrain the system had to be accounted for, before calculating the final free energy which is done using the weighted histogram analysis (WHAM) method<sup>83,84</sup>.

To perform US simulations a reaction coordinate (e.g. eigenvector from PCA) is chosen which describes the property of interest like a conformational change. For e.g. the two reference states are Open (O) and closed states (C) and the eigenvector describes this conformational change from open to closed states. The open and closed structures are projected onto the eigenvector and there eigenvalues are noted for e.g. Open: -1.20 nm Closed: 1.30 nm.

To efficiently calculate the free energy difference between these two states based on WHAM we need to have overlap between these two points. For that purpose we start multiple umbrella simulations with 0.2 nm gap between the successive umbrella windows e.g. -1.50 nm, -1.30 nm, -1.10 nm....1.20 nm, 1.40 nm. Normal MD simulation which either started with open structure or closed structure is projected onto this eigenvector and starting structures for each umbrella simulation with the corresponding eigenvalues are taken from the normal MD simulation. If the simulation has not sampled a particular portion of the reaction coordinate, starting structures are generated by linear interpolation between the two end structures i.e. open and closed state structures.

It has to be noted that given the advantages of US, the design of the reaction coordinate can influence the outcome of free energy i.e. the reaction coordinate should precisely describe the transition from state A to state B and any reaction coordinate which fails to include all the elements nessaccary for the transition may suffer from convergence issues.

#### 2.1.3 Error Estimation

Uncertainties in the calculated observables can be calculated using the block average approach discussed by Hess et al.<sup>85</sup> or by bootstrap method<sup>86</sup>. To estimate the error it is a prerequisite to select uncorrelated data points. We have started multiple trajectories with the same starting structure and if each trajectory is considered independent from the other trajectories in the ensemble we can use the bootstrap approach.

In bootstrapping procedure, where the average ' $A_x$ ' of observable 'x' calculated from each trajectory of the ensemble is bootstrapped to estimate the uncertainty in  $A_x$ .

In bootstrapping using given set of 'N' observables like  $A_1 \dots A_n$ , new 'N' observables are generated  $B_1 \dots B_n$  with replacement i.e. each observable  $A_x$  has the equal probability of being selected and the average of these observables  $C_x$  is calculated. This procedure is repeated 'M' number of times collecting  $C_x \dots C_m$ , where 'M' usually can be anything between 100 to  $10^6$  and the error estimate usually converges after few thousand iterations. The standard deviation of  $C_x$ ..... $C_m$  is the error of the variable of interest. We have verified this for our calculations and after 10000 iterations the error estimate does not change.

#### 2.2 Simulation setup

To study the structure-dynamics-function relationship of TIM, we performed apo and holo TIM molecular dynamics (MD) simulations using GROMACS MD package<sup>87-91</sup>. We used the amber99sb<sup>92</sup> all atom forcefield a variant of the Amber-99<sup>93</sup> potential in which charges were derived using Quantum mechanics (QM) calculations with HF-6/31G\* basis set with improved backbone chain torsion potentials. We used the AMBER ports<sup>94</sup> for implementation of amber99sb in GROMACS. We have used the ion type definitions from the Joung et al.<sup>95</sup>.

Ligated TIM simulations were performed with natural substrate Dihydroxyacetone phosphate (DHAP), inhibitor 2-phosphoglycerate (2PG), reaction intermediates eneldioate-1 (EDT1) and eneldioate-2 (EDT2; Figure 2.2)<sup>27</sup>. To keep the forcefield parameters of ligand molecules compatible with the protein forcefield, General amber forcefield (GAFF)<sup>96</sup> parameters were developed for the ligand molecules which is described in section 2.2.2.1.



Figure 2.2: Natural substrates DHAP and GAP, reaction intermediates EDT1 and EDT2 and the strongest binding inhibitor for TIM 2PG. Except GAP we have used rest of the molecules in our simulations.

#### 2.2.1 Apo TIM simulations

Apo TIM simulations were performed using  $5\text{TIM}^{10}$  (1.83 Å),  $6\text{TIM}^{97}$  (2.20 Å),  $1\text{NEY}^{12}$  (1.20 Å) and  $1\text{R}2\text{R}^{98}$  (1.50 Å) crystal structures. Structures for apo TIM simulations were created by removing the ligand molecules present in the active site. In general all the dimer TIM crystal structures have one chain open and one chain closed. In 5TIM and 6TIM crystal

structures there is no substrate/inhibitor bound to the active site of chain-A because of crystal contacts<sup>10</sup> and hence chain-A is in the open conformation in both 5TIM and 6TIM. Chain-B of 5TIM has a sulfate ion and active site of chain-B of 6TIM has the substrate analogue Glycerol-3-phosphate (G3P). We have used distance between C-alpha atoms of W169 and S212, V176 and S212 to distinguish TIM structures into open and CTIM structures which will be discussed in section 3.1 and based on these distances chain-B of 5TIM and 6TIM are in closed state.

In 1R2R crystal there are two dimers per asymmetric unit i.e. four chains in total. Chains A, C and D are open and chain-B is closed. 1R2R is the only crystal structure apart from the TIM monomeric variant 2VEI where one of the chains (chain-B) is in closed state with an unoccupied active site.

For MD simulations, we wanted to create a TIM structure with both chains open and one structure with both chains closed. So we performed least square fit (LSQ) of backbone of 5TIM chain-A (open) on the chain-B excluding residues K12, H94, E166, loop-6 and loop-7. Same procedure was performed using 6TIM crystal structure where closed chain-B was fitted on open chain-B and excluding the groups mentioned above to create a TIM dimer with both chains closed. The difference between the fitted chains and crystal structures was ~0.03 nm i.e. they are highly identical. These structures would be referred to as fitted-5TIM (F5TIM), fitted-6TIM (F6TIM) and fitted-1R2R (F1R2R). 1NEY is the only crystal structure with the natural substrate DHAP in both the active site pockets. We removed the DHAP molecule from both the active sites to create an apo 1NEY structure.

In all simulations we have chosen cysteines to be neutral (CYN) and lysine's were protonated (LYP). Histidine's were either HIE (hydrogen on epsilon nitrogen) or HID (hydrogen on delta nitrogen) depending on the local hydrogen bonding possibilities. The catalytic residue H94 is with hydrogen at epsilon nitrogen<sup>9,10,23</sup>.

Each crystal structure was placed in a dodecahedron box with a minimum distance of 1nm between the protein and the box walls. The box was solvated using the genbox module and ions were added to neutralize the system. Before the start of the MD simulation, the system of interest is subjected to an energy minimization procedure using steepest descent method until the largest force acting on the system was smaller than 239 kcal.mol<sup>-1</sup>.nm<sup>-1</sup>. Energy minimization is performed to remove any unfavorable interactions (like overlapping Van der

Waals spheres) present in the crystal structure which may result in large forces and create instability in the system.

After energy minimization, system is slowly equilibrated to the chosen reference temperature and pressure values, which in our simulations are 298 K and 1 atm by coupling the system to a thermostat and a barostat. Temperature and pressure equilibration are done in two phases. First the temperature of the system is brought to 298 K in 100 ps by coupling the system to berendsen thermostat<sup>99</sup>. During the temperature and pressure equilibration solvent is allowed to equilibrate and the protein atoms are position restrained using a force constant of 239 kcal.mol<sup>-1</sup>.nm<sup>-2</sup>, to preserve the crystal structure.

The end structure of the temperature equilibration is used for pressure equilibration using berendsen barostat for 1 ns. The pressure equilibrated structure (Figure 2.3a) was used for the production run where the protein is free to evolve at constant temperature and pressure of 298 K and 1 atm. Multiple trajectories were started using random starting velocities. Unless mentioned otherwise, same simulation set up and equilibration protocol was used for all the simulations.

- Five trajectories each of F5TIM and F6TIM were performed for 1 µs each.
- Three simulations of apo 1NEY were performed for 300 ns each.
- Three trajectories with F1R2R structure lasting 100 ns each.
- Simulations of unperturbed 5TIM, 6TIM and 1R2R crystal structures were also performed.

#### 2.2.2 Holo TIM simulations

We have performed holo TIM simulations using F5TIM, F6TIM, 6TIM, 1NEY and 1N55<sup>61</sup> crystal structures. Simulations with DHAP were performed using F6TIM, F5TIM, 6TIM and 1NEY crystal structures. 1N55 is the highest resolution (0.82 Å) structure available for TIM with 2PG bound in the active site pocket. Serena Donnini created the dimer of 1N55 crystal structure by performing a matrix transformation of the crystal coordinates using the BIOMT records provided in 1N55 pdb file. Simulations with inhibitor 2PG and reaction intermediates EDT1, EDT2 were performed with this 1N55 dimer structure. EDT1/EDT2 were placed in the active sites of 1N55 by least square fitting the respective reaction intermediate onto the 2PG present in the crystal structure. For DHAP simulations, the parameterized DHAP was first least square fitted to the substrate analogue G3P present in the chain-B of 6TIM. This

structure was used as the starting structure to create DHAP ligated dimers of F5TIM, F6TIM and 6TIM.

After creating the various starting structures with DHAP/EDT1/EDT2/2PG, the same equilibration procedure was applied which was used for apo TIM simulations with one extra equilibration step we call '*Ligand Equilibration procedure (LEP)*'. In LEP position restraints on both the protein and the ligand were coupled to a lambda ( $\lambda$ ) parameter and as  $\lambda$  is transformed from 0 to 1 the force constant of position restraints decreases from 239 kcal.mol<sup>-1</sup>.nm<sup>-2</sup> to 0 kcal.mol<sup>-1</sup>.nm<sup>-2</sup>, allowing the protein and ligand to equilibrate. The end structure from this equilibration procedure was used for production run.

#### 2.2.2.1 Ligand Parameterization

TIM interacts with multiple ligand molecules. DHAP and GAP are the natural substrates; 2PG is the strongest binding inhibitor for TIM (Table 1). Charge on DHAP is -2 and 2PG, EDT1 and EDT2 have a charge of  $-3^{27,39-41,100,101}$ . EDT1 is the first reaction intermediate when DHAP is converted to GAP, formed after the abstraction of proton from C1 carbon by catalytic base E166. EDT2 is the last reaction intermediate which is awaiting a proton transfer from E166 carboxylate moiety onto the C2 carbon completing the formation of GAP<sup>27</sup>.

As stated earlier General Amber forcefield (GAFF) parameters for these ligand molecules were generated (Figure 10) using Gaussian03<sup>102</sup> and Antechamber<sup>103</sup> package. The structure optimization and charge derivation was done using the basis set mentioned in Table 2. Antechamber<sup>103</sup> tool was used to do the RESP<sup>104</sup> charge fitting for every ligand molecule based on the charges obtained from QM calculation. Generally 6-31g(d)<sup>105,106</sup> basis set with Hartree-Fock (HF) method is used for charge calculation in GAFF forcefield but this basis set placed negative charges on methyl hydrogen's in the ligand. Therefore we tried multiple basis sets with HF and B3LYP<sup>107-109</sup> and B3LYP/6-311G(d)<sup>110,111</sup> gave us the best charges for DHAP, EDT1 and EDT2. We tried the same for 2PG and finally reasonable charges were obtained with basis sets mentioned in Table 2.1.

Ligand	Charge	Binding affinity	Structural optimization	Charge calculation
DHAP	-2	$0.97 \text{ mM}^{20}$	HF/6-31G (d)	B3LYP/6-311G(d)
EDT1	-3		HF/6-31G(d)	B3LYP/6-311G(d)
EDT2	-3		HF/6-31G(d)	B3LYP/6-311G(d)
2PG	-3	$0.015 \text{ mM}^{112}$	HF/6-31g(2d,p)	HF/6-311G(3d2f,p)

Table 2.1: Basis sets used for structure optimization and charge calculation. Binding affinities of natural substrates DHAP and GAP and inhibition constants of inhibitor G3P and 2PG.

\*HF: Hartree-Fock, B3LYP: Becke three parameter hybrid functional<sup>109</sup> with Lee, Yang and Parr correlations functional<sup>107,108</sup>.

To test whether the torsional parameters obtained at QM level hold in Molecular Mechanics (MM) forcefield, we performed a potential energy surface (PES) scan around the dihedrals formed by C1-C2 and C2-C3 of bonds of DHAP molecule at QM level using Gaussian03 package and then repeated the same at MM level using Gromacs 4.0.x and if different were optimized to reproduce the QM potential energy surface. Torsional parameters obtained from DHAP for dihedral angles formed by C1-C2, C2-C3 were used for C1-C2, C2-C3 bonds of EDT1 and EDT2 as well. Parameters for DHAP, EDT1 and EDT2 and 2PG are listed in appendix section 7.1. The parameterized ligand molecule was placed in the active site of the respective crystal structure and this structure was used for further holo TIM simulations.

- Three trajectories each for F5TIM, F6TIM, 6TIM with DHAP and 1N55 with 2PG, 1N55 with EDT1 and 1N55 with EDT2 in the active site were simulated.
- 1N55, EDT1 and EDT2 simulations were simulated for 300 ns each, whereas simulations with DHAP were stopped as soon as DHAP escaped from both the pockets, which was approximately 200 ns to 300 ns.

#### 2.2.3 Crystal simulations

Surprisingly in our apo TIM simulations with F1R2R and unperturbed 1R2R crystal structures, loop-6 opens up in first 20 ns of simulation time. To further explore the affect of crystal environment on dynamics of loop-6 and loop-7 we performed crystal unit cell simulations as described by Cerutti et al.<sup>113,114</sup>. One unit cell of 1R2R crystal structure was simulated to see the effect of the crystal environment on dynamics of loop-6 and loop-7. In 1R2R crystal structure each asymmetric unit is made up of two dimers and four asymmetric units form one unit cell. Each unit cell contains 12 open chains and four closed chains. Symmetry records from the pdb file were used to create a unit cell. Such a unit cell contained

crystal waters and the protein. Two sets of apo crystal simulations were performed. One structure with 1.1 M DMSO and 0.2 M of  $MgCl_2$  salt concentration used in mother liquor was created (Crystal Solution Simulation: CSS) and a second simulation with just solvent (Water Crystal Simulation: WCS). Both systems were neutralized by adding additional ions.

A solvent box of the same dimensions as unit cell was created and the unit cell with protein and crystal waters was embedded into the solvent unit cell using g\_membed<sup>115</sup> tool. The resultant structure was subjected to energy minimization as done for apo and crystal simulations, after which temperature equilibration of 1 ns and pressure equilibration of 20 ns was performed. As suggested by Cerutti et al.<sup>113,114</sup>, after pressure equilibration, volume of the simulation box was calculated and if it differed from the crystal unit cell volume by more than 0.3%, solvent molecules were added or deleted from the system.



Figure 2.3: Normal MD (a) and crystal unit cell (b) simulation box filled with water (red lines), chloride ions (blue spheres) and protein (cyan) in new cartoon representation.

The whole procedure of energy minimization, temperature equilibration and pressure equilibration were performed until the unit cell with required X, Y and Z dimensions were obtained within ~0.3% of the crystal unit cell volume. The final equilibrated structure contained ~102300 atoms and was used for production run (Figure 2.3b). In all production run trajectories unit cell volume does not differ by more than 0.3% of the crystal unit cell volume. We have simulated five trajectories with the same equilibrated structure and random starting velocities. All the open chains are combined into Crystal simulations open ensemble (CS-O) and closed chains into crystal simulations closed ensemble (CS-C).

We have also performed holo crystal simulations with 1N55 crystal structure with 2PG bound to the active site pocket. Starting structure for 2PG crystal simulations (CS-2PG) was subjected to the same protocol as the 1R2R crystal simulations. In 1N55 crystal structure, there is one monomer per asymmetric unit i.e. four monomers per unit cell and we have simulated one unit cell in MD. The 1N55 unit cell structure was subjected to similar equilibration procedure as 1R2R unit cell structure to create the simulation box which differed in volume by no more than ~0.3% of X-ray crystal unit cell. The end structure of volume equilibration was used to start three production run simulations. The volume equilibrated was also subjected to LEP and this structure was used for three more production run simulations.

- Five apo TIM crystal unit cell simulations of 300 ns length, with the corresponding equilibrated structures of WCS and CSS (which were created from 1R2R crystal structure) were performed.
- Six simulations with 2PG ligated 1N55 crystal structure (CS-2PG) three with the volume equilibrated structure and three with the end structure of LEP were performed.
- The reference temperature for unit cell simulations of 1R2R and 1N55 was 291 K<sup>98</sup> and 295 K<sup>61</sup>, the room temperature at which these proteins were crystallized.

TIP3P water model was used for solvent<sup>116</sup>. We have used Particle-mesh Ewald algorithm<sup>117,118</sup> to evaluate the electrostatistics interaction with a cutoff of 1.0 nm. Van der Waals interactions were calculated with a cut-off of 1.4 nm. LINCS<sup>119</sup> algorithm was used to constrain on all-bonds. Nose-Hoover (NH) thermostat<sup>120,121</sup> and Parrinello-Rahman (PR) barostat<sup>122,123</sup> were used for maintaining NPT ensemble for the production run simulations. For unit cell simulations we used Berendsen barostat<sup>124</sup> along with because if its efficiency in maintaining unit cell volume better than PR barostat along with v-rescale thermostat<sup>125</sup>. Coordinates were saved every 20 ps and energies were written every 10 ps.

## **3** Results and Discussion

### 3.1 Analysis of various X-ray structures

Importance of X-ray crystallography cannot be overstated in determining the three dimensional structure of proteins. If a large number of crystal or NMR structures are available for the protein of interest, a thorough analysis of the various available structures can provide wealth of information regarding the conformational properties of the protein.

Of the 126 TIM X-ray structures available, 93 were analyzed using X-ray PDB Analysis Tool (XPATv1.2) written by us which reads a pdb or multiple pdb files and calculates distances or angles or dihedral angles for selected residue(s), as requested by the user. Twenty three structures with resolution greater than 2.5 Å were excluded from the analysis. Phi and psi angles of loop-7 residues Y209 to N214 and loop-6 residues E166 to T178 were calculated to identify the dihedral angles undergoing a change between open and closed states (Table 3).

As discussed in section 1.2, dynamics of V168 from loop-6 N-terminus and C-terminus residues K175 and T178 were monitored in NMR experiments<sup>39,41</sup>. So, to segregate TIM X-ray structures into open and closed states based on N-terminus and C-terminus conformations of loop-6, distance between W169 from N-terminus of loop-6 and loop-7 residue S212 was calculated in all X-ray structures (W169-S212). And to determine loop-6 C-terminus conformation, distance between loop-6 C-terminus residue V176 and S212 (V176-S212) was calculated. Based on W169-S212 and V176-S212 distances (Table 3.1), TIM structures were classified as open TIM (OTIM) when both loop-6 N and C-terminus are open (W169-S212: 1.15 nm, V176-S212: 0.88 nm) and as closed TIM (CTIM) when both N and C-terminus are closed (W169-S212: 0.95 nm, V176-S212: 0.50 nm).

As discussed in section 1.3, loop-7 also exists in open and closed conformations. Loop-7 is segregated into open and closed conformations based on the backbone phi ( $\Phi$ ) and psi ( $\Psi$ ) dihedral angles of residues G210, G211, S212 and V212. Dihedral angles G210 $\Phi$ , G210 $\Psi$ , G211 $\Phi$ , G211 $\Psi$ , S212 $\Phi$ , S212 $\Psi$  and V213 $\Phi$  differ by more than 25° between open and closed conformations (Table 3.1).

		-
Residue	Closed	Open
W169-S212	$0.95 \pm 0.02$	$1.15 \pm 0.03$
V176-S212	$0.50 \pm 0.04$	$0.88 \pm 0.08$
Р167-Ψ	126 ± 8	77 ± 7
К175-Ψ	81 ± 10	$154 \pm 9$
V176-Ф	296 ± 11	237 ± 14
G210-Ф	221 ± 5	248 ± 6
G210-Ψ	151 ± 6	23 ± 7
G211-Φ	128 ± 5	270 ± 6
G211-Ψ	93 ± 5	197 ± 7
S212-Φ	$62 \pm 5$	292 ± 8
S212-Ψ	36 ± 9	115 ± 9
V213-Ф	286 ± 7	286 ± 12

Table 3.1: Average and standard deviation, of backbone dihedral angles  $(\Phi, \Psi)$  of loop-6 and loop-7 residues which differ between open and CTIM's along with the distance between residues W169-S212 and V176-S212. Distances are in nanometers and angles are in degrees.

Majority of X-ray structures (Table 4) in the absence of ligand are in open state (43) and in the presence of ligand are in closed state (73). It must be noted that there are two crystal structures with loop-6 in closed conformation (1R2R-B, 2VEN-A<sup>126</sup>) without the ligand in the active site pocket. Loop-7 of 1R2R-B is also in closed conformation whereas in 2VEN-A, excluding S212 $\Phi$ , entire loop-7 is in open conformation.

Table 3.2: A count of open TIM (OTIM) and closed TIM (CTIM) structures in apo and holo states from analysis of X-ray structures using XPATv1.2. Holo TIM is divided into reactive complex (RC) and encounter complex (EC) states based on ligand orientation in the active site.

<b>Confirmation</b> \Ligand	APO	HOLO	
		RC	EC
OTIM	43	4	5
CTIM	2	73	0

In 1R2R crystal structure, out of four chains A, B, C and D, chain-B is in closed state. There is no ligand in the active site but there is a water molecule (W171) which is present at the location where phosphate moiety of ligand is situated forming a hydrogen bond with S212

amide group<sup>98</sup>. Further studies are necessary to ascertain, if a water molecule located at the phosphate moiety's position can cause flip of loop-7 to closed state. Closed loop-6 of 1R2R-B was reported as a feature of TIM's conformational heterogeneity by Ricardo et.al<sup>98</sup>.

The second crystal structure which has loop-6 in closed conformation is 2VEN. Loop-6 in chain-A of 2VEN dimer adopts closed conformation in the absence of ligand. In chain-A of 2VEN-A electron density for lid region (I172-GT-175G) of loop-6 is missing. In OTIM loop-6 has high B-factors and same was observed for loop-6 in 2VEN-A<sup>126</sup>. But based on loop-6 N and C-terminus distance we would suggest that 2VEN-A as a closed structure. These two closed-apo structures suggest that loop-6 might also exist in closed conformation even in the absence of the ligand, though open state is the preferred conformation in apo TIM.

In our analysis we also came across 14 chains where both loop-6 and loop-7 were in open conformation with ligand occupying the active site (open-holo). To understand the reason behind these structures being in open conformation despite of the occupied active site, we analyzed the ligand orientation. Each structure was visually compared with the encounter complex (EC) structure of 1LZO-B<sup>127</sup> and reactive complex (RC) of 1NEY<sup>128</sup>, 1N55<sup>61</sup> structures (Figure 11a). In 1M7O<sup>48</sup> (3PG: 3-Phosphoglycerate) and 1M7P<sup>48</sup> (G3P) where substrate analogs are bound in RC state, the S96F mutation prevents loop-6 from closure. Rest of the 10 chains had ligand in the encounter complex position. When we analyzed the orientation of ligand in the closed-holo structures we found out that all the closed-holo chains had the ligand in RC orientation.

In the encounter complex we found van der Waal's clashes between the phosphate moiety of the inhibitor and the loop-6 residues I171 and G172 which can prevent loop-6 closure (Figure 11b). The mechanism of loop-7 conformational change and influence of ligand on loop-7 has not been explored which we address in this work in section 3.5.1. As stated earlier, in chain-B of 1R2R crystal structure, a water molecule occupies the space of one of the oxygen's of the phosphate moiety, within hydrogen bonding distance G172 of loop-6 and S212 of loop-7. Looking at the X-ray structures it might be speculated that the electrostatic repulsion between the phosphate moiety and the carbonyl oxygen might drive the flip of G211 $\Psi$  and S212 $\Phi$  to closed conformation and it is quite surprising that a 0.2 nm change in shift of phosphate moiety towards the mouth of active site induces a loop-7 opening.


Figure 3.1: a) 2-Phosphoglycerate (2PG) in reactive (1N55-A) and encounter complex (1LZO-B) structures with active site residues K12, H94 and E166. b) 2PG in encounter complex with closed loop-6 residues I171-G172. Closed loop-6 is from 1N55.

Our analysis also revealed structures where loop-6 N and C-terminus were in different states i.e. both the hinges were neither closed nor open at the same time (Table 5). Crystal structures 2BTM, 2J24<sup>14</sup> are mutant TIM structures whereas 2VEI<sup>126</sup> is a monomeric TIM variant. The P167A mutation in 2J24 induces this conformational heterogeneity in loop-6 whereas in 2BTM<sup>129</sup> the mutations H12N, K13G are not in loop-6. In 2VEI, V233 was mutated to A233 to extend the active site and enable binding of citrate (CIT) molecule. Loop-6 and loop-7 are in closed state, in these CIT bound structures. The mutant structures have decreased catalytic activity<sup>14,129</sup>. 1WYI<sup>130</sup> is a wild type human TIM with no mutations at 2.20Å resolution<sup>130</sup>.

PDB	Lo	Loop-7	
	N-terminus (W169-S212)	C-terminus (V176-S212)	-
1WYI-B-APO	Open	Closed	Open
2BTM-A-PGA	Closed	Open	Closed
2BTM-B-PGA	Closed	Open	Closed
2J24-A-APO	Closed	Open	Open
2J24-B-APO	Closed	Open	Open
2VEI-A-APO	Open	Closed	Open

Table 3.3: X-ray structures with loop-6 N and C-terminus in different states.

To summarize the TIM X-ray structures analysis:

- In crystal structures loop-6 and loop-7 exist in open conformation in the absence of ligand.
- Loop-6 may also exist in closed conformation in the absence of the ligand (2VEN-A and 1R2R-B).
- Apart from chain-B of 1R2R X-ray structure where a water molecule is located in the place of phosphate moiety, loop-7 does not exist in closed conformation when there is no ligand bound to the active site.
- There is only one crystal structure with natural substrate DHAP bound in the RC orientation with both loop-6 and loop-7 in closed conformation.
- Ligand molecules i.e. substrate or inhibitor can exist in RC and EC states in the active site.
- Loop-6 and loop-7 are in open conformation when the ligand is in the EC state.
- Only when a ligand molecule is in the RC state, both loop-6 and loop-7 are seen in closed conformation.

As suggested earlier in section 1.5 in our simulations we will monitor the conformational change of loop-6 and loop-7. Using MD simulations we will also check the influence of ligand's orientation in the active site address on the conformations of loop-6 and loop-7.

# 3.2 Stability of simulations

It is important to verify the stability of simulated trajectories. We have looked at the Root mean square deviation (RMSD) values of all the trajectories. RMSD tells us how much the structures deviate from the starting crystal structure. All the trajectories showed a stable RMSD of less than 0.2 nm. Major structural changes are seen in loop-6 with minor fluctuations in loop-5 and loop-7. Rest of the protein is highly stable.

# 3.3 Ligand residence time

We have simulated OTIM and CTIM structures with natural substrate DHAP, reaction intermediates EDT1 and EDT2 and inhibitor 2PG. In each holo TIM simulation we have calculated the time spent by each ligand in RC, EC and apo orientations which we define as residence time (RT). RT is calculated using Principal component analysis (PCA) and RMSD analysis. To describe the change in orientation of each ligand in the active site pocket from RC to EC orientation, we have used the eigenvector-1 (LEv-1) from PCA (section 2.1.1).

PCA was performed on chain-B of 1LZO where 2PG is sitting at the mouth of the active site which is EC orientation (Figure 3.1a) and 1NEY structure where DHAP is present in RC orientation. LEv-1 describes the ligand transition from RC to EC state.

We also calculated RMSD of the ligand molecule (excluding hydrogen atoms) with respect to the ligand molecule in the starting structure of the simulation by using the protein backbone as the fitting group. When the ligand moves out of EC state into the solution, RMSD of the ligand with respect to the ligand molecule in the starting structure is greater than 0.7 nm. Therefore all the frames of the trajectory where RMSD of the ligand is less than or equal to 0.7 nm, are used to calculate RT. Frames with RMSD greater than 0.7 nm are considered apo.

Based on the eigenvalues of LEv-1 we classify the ligands orientation into RC (-0.50 nm to 0.50 nm) and EC (-0.5 nm to 1.0 nm) states. We have started all our simulations with DHAP, 2PG, EDT1 and EDT2 in the RC orientation. In Figure 3.2 we have plotted the average RT of each ligand molecule over the entire ensemble and the error bars were calculated by bootstrap method which was described in section 2.1.3.

As seen in Figure 3.2 all the ligand molecules spend more simulation time in RC state compared to the EC state. We have placed DHAP in both OTIM and CTIM active site pockets. Irrespective of the starting TIM conformation, DHAP moves out of the pocket in all the simulations. The average RT (time spent in RC+EC orientations) of DHAP with OTIM as the starting structure is 74 ns which is less compared to 121 ns when the simulation starts with CTIM (Figure 3.2).

Inhibitor 2PG, in five out of six chains is bound to the active site pocket for entire 300 ns of simulation. Only in one chain out of the six chains we observe 2PG moving into the solution after 117 ns (data not shown). Contrary to X-ray data, in holo TIM simulations with DHAP and 2PG loop-6 does not stay closed and transiently samples open and closed conformations (which is discussed further in 3.4.1). So we have also performed simulations with the reaction intermediates EDT1 and EDT2 (Figure 1.8).

Similar to 2PG, EDT1 stays inside the active site pocket for entire length of simulation for 300 ns in five out of six chains and only in one chain EDT1 moves into the solution after 157 ns. Whereas in the simulations with EDT2 bound to the active site, we observe EDT2 dissociating into the solution in all our simulations. Average RT of EDT2 (224 ns) is much longer than the natural substrate DHAP and shorter than EDT1 (275 ns; Figure 3.2). In

comparison, 2PG, EDT1 and EDT2 spend more time in the active site compared to the natural substrate DHAP (Figure 3.2).



Figure 3.2: The average residence time (RT) of natural substrate DHAP, reaction intermediates EDT1, EDT2 and inhibitor 2PG in normal MD simulation (2PG) and crystal simulation (CS-2PG) the active site pocket of TIM, in reactive complex (RC) and encounter complex (EC) orientation. Error bars were calculated by bootstrapping the averages as discussed in section 2.1.3.

TIM has achieved maximum theoretical efficiency possible for an enzyme  $(10^9 \text{ M}^{-1}.\text{s}^{-1})$  and is limited by the diffusion of the substrate into the active site<sup>43</sup>. To achieve this turnover rate not only would the enzyme optimize the efficiency of catalytic steps but would also get rid of the product formed in the active site as soon as possible. Taking that view point into consideration it would be of no surprise that DHAP escapes into solution in all our simulations.

2PG is a transition state analogue and the charge of -3 compared to the charge on substrates DHAP and GAP of -2, provides additional electrostatic stabilization in the active site contributing to its higher binding affinity<sup>112</sup>. Hence 2PG should bind very strongly to the enzyme and 2PG stays bound to the active site for the entire 300 ns of the simulation. Enzymes are speculated to stabilize the reaction intermediates more compared to the substrate, and we observe higher RT for reaction intermediates in our simulations compared

to the natural substrate. But to confirm our speculations binding free energy of these ligand molecules in OTIM and CTIM conformations needs to be calculated which is a more reliable measure compared to the RT presented here.

# 3.4 Dynamics of loop-6

Loop-6, as discussed in section 1.2 plays a very pivotal role in TIM's function. The fundamental questions we want to answer about loop-6 are as follows.

- Nature of loop-6 conformational change, is it ligand driven or a spontaneous change driven by thermal fluctuations?
- Is loop-6 conformational change a rigid body motion i.e. both N and C-terminus move together (correlated) or N and C-terminus of loop-6 independent?
- What is the free energy difference between the open and closed loop-6 in both presence and absence of the ligand, and what factors contribute to this free energy difference?
- How does binding of a ligand (substrate/inhibitor/reaction intermediates) affect dynamics of loop-6?

To understand the various aspects of dynamics of loop-6, we performed multiple apo and holo TIM simulations. In our holo TIM simulations we have used the natural substrate DHAP, inhibitor 2PG and reaction intermediates EDT1 and EDT2. All the ligated TIM simulations start with ligand in the RC orientation.

## 3.4.1 Apo and holo TIM simulations

We have performed 16 apo TIM simulations in which there are 13 OTIM chains and 19 CTIM chains. Out of 13 OTIM chains with open loop-6 conformation, in 10 chains, 2 to 14 transitions between open and closed conformations of loop-6 are observed. In three OTIM chains there are no transitions. All the simulations that start with loop-6 in closed conformation, undergo a conformational change to open state in the first 20 ns of simulation time (Figure 3.3d), longest being 150 ns. In seven CTIM chain trajectories no transitions between open and closed loop-6 are observed and 2 to 21 transitions are seen in rest of the 12 CTIM chains. A typical loop-6 N-terminus (W169-S212) and C-terminus (V176-S212) distance plot is shown in Figure 3.3a where we have 20 transitions between open and closed conformations of the 12 CTIM structure.



Figure 3.3: Dynamics of Loop-6 N-terminus (W169-S212) and C-terminus (V176-S212) in a microsecond trajectory which started with an apo CTIM structure (a) and 300 ns long trajectory where EDT1 is bound for entire length of simulations (b). Loop-6 RMSD with respect to closed loop-6 is shown in the third panel. c) Average number of transitions between open and closed conformations of loop-6. For ligated ensembles only that part of the trajectory during which ligand is bound to the active is considered. d) Average time taken by loop-6 N-terminus (W169-S212) and C-terminus (V176-S212) to undergo conformational change from closed to open conformation for the first time in the simulation, when the simulations started with a CTIM structure in various ensembles. Error bars are calculated using bootstrap method described in section 2.1.3. In inset of plot (d) loop-6 with N-terminus (blue), tip (orange) and C-terminus (red), loop-7 (cyan) are shown. Dotted blue line is the N-terminus distance in the X-ray closed structure and dotted green line in X-ray open structure. Dotted cyan line is the RMSD of X-ray open loop-6 versus X-ray closed loop-6.

To check if the closed loop-6 in MD simulations resembles the closed loop-6 of the crystal structure, we have calculated the RMSD of the backbone of loop-6 with respect to loop-6 of the CTIM X-ray structure and OTIM X-ray structure for all our trajectories. (RMSD with respect to X-ray closed loop-6 is shown here in (Figure 3.3a, b). RMSD with respect to both X-ray open loop-6 and X-ray closed loop-6 is shown in appendix section 7.2, Figure 7.1).

The entire protein backbone excluding loop-6 is used as the fitting group. RMSD of X-ray open loop-6 versus X-ray closed loop-6 is 0.39 nm. The RMSD difference observed between the MD loop-6 and X-ray closed loop-6 structure is between 0.15 nm and 0.2 nm (Figure 3.3a, b) suggesting that loop-6 is indeed closed.

Our analysis of TIM X-ray crystal structures has revealed that when ligand is bound in RC orientation loop-6 is in closed conformation. Surprisingly in our holo TIM simulations with DHAP/2PG/EDT1/EDT2 in RC/EC orientation no preference for closed conformation of loop-6 is observed (Figure 3.3b) as expected from the X-ray data. In all the holo TIM simulations that start with CTIM structure, loop-6 undergoes a conformational change to open loop-6 in the first 40ns (Figure 3.3d). In simulations with DHAP/2PG during the simulation when the ligand is bound to the active site in RC orientation, no transitions back to the closed conformation of loop-6 are observed (Figure 3.3c). In simulations with EDT1 and EDT2 in RC orientation we observe one transition each, in two out of five chains in which ligand is bound to the active site (Figure 3.3c).

Loop-6 to and fro samples open and closed conformations in apo TIM simulations (Figure 3.3). There are fewer transitions in holo TIM simulations because of shorter simulation times (Figure 3.3). It might be possible that the time required for closure of loop-6 is a much slower process than 300 ns of simulation time during which the ligand molecules are bound to the active site, to induce closure of loop-6. As expected from X-ray structure data in holo TIM simulations we do not see preference for closed state. Richard and co workers<sup>31</sup> have proposed that the binding energy of the phosphate moiety is used for inducing closed conformation for loop-6 which is not observed in our holo TIM simulations.

# 3.4.2 Effect of crystal environment

In apo and holo TIM ensembles when simulations start with CTIM structure loop-6 undergoes a conformational change to open state in the first 40 ns of simulation (Figure 3.3d). To see if there was an influence of crystal environment on the conformation of loop-6 i.e. may be only in the X-ray crystal, closed conformation of loop-6 is stable. So, we performed both apo and holo crystal unit cell simulations with 1R2R (apo) and 1N55 (holo with 2PG in active site pocket) structures. There is no difference observed in the behaviour of loop-6 in the crystal solution simulation (CSS) and water crystal simulation (WCS) hence we discuss both of them as one ensemble. In crystal simulations CS-C and CS-2PG, time taken for this conformational change is greater than 100 ns (Figure 3.3d) compared to the first 40 ns

in normal MD simulations. Time taken by N-terminus and C-terminus of loop-6 to open is different with N-terminus opening first followed by C-terminus. The reason behind this is addressed in section 3.4.4.

Ensemble	<b>Diffusion constant</b> (x10 <sup>-5</sup> cm <sup>2</sup> .s <sup>-1</sup> )
Apo/holo TIM	5.02
CSS	0.58
WCS	0.78
Experimental value for Water	$2.30^{116}$

Table 3.4: Diffusion constant of solvent in the simulation box of various ensembles.

Diffusion constant of water from MD simulations was calculated using Einstein relation<sup>131</sup> from mean square displacement of water molecules. Solvent dynamics in crystal simulations is 10 times slower than in the normal MD simulations (Table 3.4). Because of the slow solvent dynamics, the dynamics in the crystal simulations is slow (Figure 3.4d). In 9 out of 40 chains, loop-6 stays closed for the entire 300 ns of simulation (Figure 3.4a). In six chains, C-terminus of loop-6 stays closed and N-terminus undergoes transitions between open, closed structures and the opposite is seen in only two chains. Even in the 2PG ligated unit cell simulations though we observe a slower dynamics of loop-6, loop-6 undergoes conformational change from closed state to open state. It is clear that the crystal environment slows down dynamics of loop-6, nonetheless loop-6 does undergo a gradual conformational change in both apo and holo crystal simulations (Figure 3.4b).



Figure 3.4: Dynamics of loop-6 in crystal simulations. (a) Simulation which started with closed loop-6 stays closed for the entire length of simulation and (b) simulation with four transitions between open

and closed conformations of loop-6. Loop-6 RMSD with respect to closed loop-6 is shown in the third panel. Dotted blue line is the N-terminus/C-terminus distance in the X-ray closed structure and dotted green line in X-ray open structure. Dotted cyan line is the RMSD of X-ray open loop-6 versus X-ray closed loop-6.

Diffraction data for 1R2R and 1N55 was collected at 100 K. It is a common practice to flash freeze the protein crystal to less than 125 K to reduce the radiation damage to the crystal<sup>132</sup>. An assumption made prior to flash freezing a protein is that this procedure traps the protein in conformation at the room temperature and does not affect the conformational equilibrium of the protein<sup>133-135</sup>.

Time scale of this procedure is in seconds and there is uncertainty about the effect of flash freezing on conformation of the crystallized protein<sup>136-139</sup>. Halle<sup>136</sup> suggested that during flash freezing procedure, different regions of protein may get trapped in low a enthalpic minimum at different temperatures and the whole structure may not be a representative of structure at one single temperature. Dunlop et al.<sup>137</sup> in 2005 collected diffraction data for six crystals, three at 100 K and three at 293 K and demonstrated that there is a significant difference with respect to side chains and hydrations sites specifically at the surface of the protein.

Our crystal simulations were performed at 291 K. Simulations at 100 K will further slow down the dynamics and if the assumption that conformational equilibrium is not altered by flash freezing, our simulations at 291 K should reproduce the experimental data. To verify this, we have compared the B-factors of C-alpha atoms from crystal open (CS-O) and crystal closed (CS-C) MD simulations with experimental B-factors of chain-A from 1R2R (OTIM Crystal @ 100 K), chain-B from 1R2R (CTIM Crystal @ 100 K), 1R2S (OTIM Crystal @ 298 K) and 1N55 crystal structures for which diffraction data was collected at 100 K, 298 K and 100 K respectively (Figure 3.5a).



Figure 3.5: Average B-factors calculated from MD unit cell simulations of apo 1R2R (a) and 2PG ligated 1N55, CS-2PG (b).

We calculated the spearmen rank correlation coefficient<sup>140</sup> between the B-factors from MD crystal simulations and B-factors from crystal structures and B-factors calculated from normal APO MD simulations. Crystal MD simulations were able to reproduce the trend seen in X-ray crystal structures and show a moderately strong correlation between 0.60 and 0.78 (Table 3.5). Residues 1-10, 38-50, 75-90 reproduce the trend seen in crystal data but have much smaller fluctuations. This may be because they are trapped in some local minimum and require more simulation time to equilibrate.

Protein residues 29 to 32 in CS-O, residues 132 to 142, loop-7 residues G207 and G208 in both CS-O and CS-C have higher B-factors than 1R2R crystal at 100 K but similar to seen at 1R2S. Loop-6 residues 167 to 174 have higher flexibility when compared to crystal data both at 100 K and 291 K. But majority of the protein shows B-factors resembling the 1R2R crystal at 100 K hence higher correlation of greater than 0.7 with 1R2R crystal structure than 1R2S. In both the crystal simulations CS-C, CS-O and normal apo MD simulations fluctuations are seen in the same protein fragments with one exception of residues 69 to 72 (solvent exposed loop) which show higher fluctuations in normal MD simulations (Figure 3.5a).

Table 3.5: Table 3.5: Spearman rank correlation coefficient between B-factors of C-alpha atoms calculated from MD unit cell simulations and crystal structures.

MD ensemble	1R2R open	1R2R closed	1R2S open	Apo MD	
	at 100 K	at 100 K	at 298 K	simulations at	
				298K	
CS-O	0.770	0.664	0.669	0.822	
CS-C	0.712	0.606	0.586	0.744	
1N55 Closed at 100K					
CS-2PG		0.613			

B-factors calculated from 1N55 crystal unit cell simulations (Figure 3.5b) also show higher B-factors than the crystal data especially in the loop-6 residues 167 to 178 and loop-7 residues 211 to 217. Rest of the residues which show deviation from crystal data are present in the dimer interface (68 to 75 and 99 to 104) or are present in solvent exposed loops (29 to 34, 52 to 57 and 149 to 156). As each asymetric unit of the 1N55 unit cell has a monomer higher fluctuation in the dimer interface is expected where there are contacts with the other monomer. Though the temperature in 1R2R and 1N55 unit cell simulations was maintained at 288.3 $\pm$ 0.01 K and 292.2 $\pm$ 0.01 K the agreement with experiment is not exact. Especially we observe deviations in loop-5, loop-6 and loop-7 regions. Loop-5 and loop-7 resemble the crystal data at 298 K.

The reason behind the difference between the MD unit cell simulations and the X-ray data can be following. Aparicio et al.<sup>98</sup> argued that the closed chain in 1R2R crystal structure is represents the "conformational heterogenity" of the protein which means that loop-6 can adopt open and closed conformations in apo TIM as well. Our simulations support this argument as we see transitions between open and closed conformation of loop-6 in both 1R2R and 1N55 crystal unit cell simulations. The time taken for loop-6 to open, starting with a closed conformation is much longer in the crystal environment because of the slow solvent dynamics. Aparicio et al.<sup>98</sup> have ruled out effect of crystal contacts and DMSO on the closed conformations of loop-6 and loop-7 and pointed out to a water (Water 171) which interacts with amide groups of G172 of loop-6 and S212 of loop-7, stabilizing the respective closed conformations.

We have performed simulations at 291 K and 295 K respectively for 1R2R and 1N55 crystal simulations and Aparicio et al.<sup>98</sup> also suggested that effects of crystallization process cannot be overlooked. To verify this hypothesis that flash freezing affects on the conformation of

loop's 5, 6 and 7, multiple annealing simulations of structures from MD simulations at 298 K to 100 K (at which diffreaction data was collection) in microsecond to millisecond time scale starting with open loop-6 and loop-7 have to be performed which is computationally very intensive.

### 3.4.3 Loop-6 N and C-terminus correlation

Berlow et al.<sup>39</sup> and Wang et al.<sup>41</sup> observed different population ratios for N-terminus and Cterminus hinges of loop-6 and suggested requirement of further studies to explore this novel view of loop-6 dynamics. To verify this we divided loop-6 into five conformations based on W169-S212 and V176-S212 distances for loop-6 (Table 3.6). In all apo and holo TIM ensembles population percentage of each conformation of loop-6 in each simulation was calculated.

Conformation of loop-6	Description		
Closed (C)	Both N-terminus & C-terminus are closed		
N-terminus closed (NC)	N-terminus closed & C-terminus open		
C-terminus closed (CC)	N-terminus open & C-terminus closed		
Intermediate (IMD)	Neither closed nor open		
Open (O)	Both N-terminus & C-terminus are open		

Table 3.6: Various conformations of loop-6 based on W169-S212 and V176-S212 distances.

If loop-6 N and C-terminus hinges are correlated and loop-6 moves as a rigid body we would expect only O, IMD and C states to be populated. And in all our simulations both apo, holo and unit cell simulations, not only do we observe O, C and IMD states but also different populations for NC and CC states. In Figure 3.6, loop-6 average population percentage in each conformation of loop-6 is shown for four ensembles. The uncertainties in the population percentages were calculated by boot strap method as described in section 2.1.3.

As stated earlier in sections 3.4.1 and 3.4.2 we do not see any influence of the ligand bound in the active site on the population of loop-6 in a particular conformation. In apo and holo TIM simulations percentage of closed loop-6 conformation is less than 1% and only in CS-C and CS-2PG it is 30 % and 33% (Figure 3.6). The percentage of NC and CC conformation varies between 1% to 10% and the difference can be attributed to the difference in the apo ensemble (23  $\mu$ s) and holo TIM ensemble (1.7  $\mu$ s to 2.7  $\mu$ s) sizes. In the NMR experiments loop-6 was only identified as open (NMR-Open: NMR-O) and closed (NMR-Closed: NMR-C) states. Based on the angular hop of W169 present in the N-terminus of loop-6, which was used as an indicator of loop-6 conformation by McDermott and coworkers<sup>35-38</sup> we have segregated loop-6 five conformations into NMR-O and NMR-C which we will discuss in section 3.4.8.2.



Figure 3.6: Average population percentages of loop-6 in various conformations, in Apo TIM ensemble (APO), apo unit cell simulation started with closed structure (CS-C), CTIM simulation with EDT1 bound in the active site (EDT-1) and 1N55 unit cell simulation ligated with 2PG. The value in square brackets is the percentage of simulation time 2PG or EDT1 is bound in RC orientation. Loop-6 conformations Open (cyan), closed (green), N-terminus is closed (dark blue), C-terminus is closed (red) and neither N-terminus nor C-terminus are open or closed (orange). The average population percentage with error bar is given in the legend for each pie chart.

We also calculated the normalized mutual information<sup>141,142</sup> (NMI) between W169-S212 and V176-S212 distances in all the TIM ensembles (Figure 3.7). On a NMI scale of 0 to 1, 0

means no correlation and 1 means perfect correlation. In all our simulations we observe a consistent NMI of less than 0.2 (Figure 3.7), which would imply that N-terminus and C-terminus of loop-6 are uncorrelated. The reason behind this independent movement is that N-terminus of loop-6 interacts with loop-5 and is obstructed by loop-5 before it can open which is discussed in sections 3.4.4 and 3.4.7. Whereas C-terminus is solvent exposed and is not inhibited by part of the protein in regards to its dynamics.



Figure 3.7: Normalized Mutual Information (NMI) between W168-S212 and V176-S212 distances in various TIM ensembles. On a scale of 0 to 1, 0 means no correlation and 1 means perfect correlation.

#### 3.4.4 Effect of Loop-5 on loop-6 conformation

The effect of loop-5 on dynamics of loop-6 is largely unexplored. A non zero exchange contribution was detected for loop-5 residues G127 and L130 by Berlow et al.<sup>39</sup> suggesting a conformational change for these residues but this was not further investigated<sup>39</sup>. Only a minor structural change is observed in loop-5 between the OTIM and the CTIM X-ray structures with RMSD less than 0.05 nm. Loop-5 residue E128 forms a hydrogen bond with W169 of loop-6 in closed conformation and this hydrogen bond is broken in open loop-6.

To quantify the correlation existing between loop-5 and loop-6 residues in all the ensembles, we used the generalized correlation coefficient ( $r_{MI}$ ) developed by Lange et al.<sup>143</sup> which calculates correlations between atoms based on the atomic fluctuations. Two pairs of atoms have correlated motion will show higher correlation (on a scale of 0 to 1, 0 means no correlation and 1 means correlation) than the atoms whose motion is not coupled. There is a much higher correlation ( $r_{MI}$  between 0.48 and 0.60) between loop-5 residues 127 to 131 and loop-6 N-terminus residues P167, V168 and W169 (Figure 3.8). The correlation between loop-5 residues 127 to 131 and loop-6 residues decreases to 0.16 as we proceed towards loop-6 C-terminus residues (Figure 3.8).



Figure 3.8: Correlation between Loop-5 residues 127 to 137 and loop-6 residues 167 to 178 in apo TIM ensemble (APO), DHAP ligated TIM ensemble (DHAP), 2PG ligated TIM (ensemble) and EDT1 ligated TIM ensemble (EDT1). Loop-6 N-terminus residues are P167, V168 and W169. C-Loop-6 C-terminus residues are K175, V176 and A177. On a scale of 0 to 1, 0 means no correlation and 1 means perfect correlation.

The correlation between loop-5 residues 127 to 131 and loop-6 C-terminus residues 175 to 177 is less than 0.35. And the nature of correlation does not change in presence of DHAP or EDT1 or 2PG or in unit cell simulations (Figure 3.8). Our results show that loop-6 N-terminus hinge is more correlated with loop-5 residues 127 to 131 then loop-6 C-terminus hinge.

We performed three simulations with 6TIM crystal structure with position restraints on residues 1 to 150 (Pres-6TIM). We wanted to freeze motion of the first 150 residues which includes loop-5 (127 to 137). Loop-6 of chain-B in 6TIM is closed in the starting structure and during the simulation residues 172 to 177 adopt open conformation while N-terminus of loop-6 (167 to 169) is hindered by loop-5 and stays closed (Figure 3.9). From our Pres-6TIM simulations it can be seen that change in conformation of loop-5 from closed to open is required for loop-6 N-terminus to adopt open conformation. But in chain-A where loop-5 is arrested in open conformation, loop-6 N-terminus can sample closed conformation.



Figure 3.9: Snapshot of chain-B from the Pres-6TIM simulation where first 150 residues are position restrained in which N-terminus (blue) of loop-6 is closed because loop-5 is also in closed conformation. Solvent exposed tip (orange) and C-terminus (red) of loop-6 are open. Starting structure is in yellow.

### 3.4.5 Loop-6 transition rate

Based on the W169-S212 and V176-S212 distances, average number of transitions in various apo and ligated ensembles including the unit cell simulations was calculated (Figure 3.10). A transition is counted as one jump from open to closed conformation of loop-6 and back to

open state. It has to be noted we do not observe a transition in every trajectory between open and closed conformations of loop-6. In few trajectories we have as many as 20 transitions (Figure 3.4a).

The difference in number of transitions observed between various ensembles should be interpreted cautiously because our simulations have not converged i.e. in trajectories started with the same starting structure, in some trajectories we have seen transitions throughout the trajectory and in some simulations no transitions at all. Based on the transitions we have calculated the exchange rate between  $(k_{ex})$  open and closed conformations can be estimated which is on the order of  $10^6 \text{ s}^{-1}$ .



Figure 3.10: Average number of transitions of loop-6 between open and closed conformations. For ligated ensembles only that part of the trajectory during which ligand is bound in the active is considered.

In the ligated trajectories of DHAP-C and 2PG where we start with closed loop-6 and DHAP-O where we start with open loop-6, we have no transitions between open and closed conformations of loop-6 as long as the ligand is present in the active site. In DHAP-C and 2PG as stated earlier, loop-6 undergoes a conformational change to open conformation in the first 10 ns of the simulation (Figure 3.3d). This must not be interpreted as, when the ligand is bound to the active site, loop-6 does not undergo a conformational change as we do see transitions in EDT1, EDT2 and CS-2PG. We need to perform longer simulations with ligand bound to the active site as we did for apo TIM simulations which were 1 µs long.

### 3.4.6 Loop-6 PCA

Loop-6 undergoes conformational change between open and closed states in apo, holo and unit cell ensembles on microsecond time scale. But as suggested by the X-ray data we do not observe a preference for closed loop-6 when ligand is bound to the active site. The experimental estimate for the free energy difference ( $\Delta G$ ) between open and closed states is given in Table 8. In apo TIM with open loop-6 as major conformation, the  $\Delta G$  between open and closed conformations of loop-6 is 1.23 kcal.mol<sup>-1</sup> to 1.77 kcal.mol<sup>-1</sup>. Based on the X-ray data when the ligand is bound to the active site it is expected that loop-6 is closed with  $\Delta G$  of -1.23 kcal.mol<sup>-1</sup> to -1.88 kcal.mol<sup>-1</sup>.

Table 3.7: Estimates of the free energy difference between open and closed conformations of loop-6 based on population ratios from various NMR experiments.

Apo/holo	Open (p <sub>a</sub> )	Closed (p <sub>b</sub> )	$\Delta G (\text{kcal.mol}^{-1})$	
APO	8/20	1	1.23 to $1.77^{35}$	
2PG/G3P	1	8/20	$-1.23$ to $-1.77^{35}$	
APO (V167)	0.74	0.26	0.62 <sup>39</sup>	
APO (T177)	0.91	0.09	1.37 <sup>39</sup>	
G3P	0.04	0.96	-1.8857	

To calculate the free energy difference between open and closed states of loop-6 from MD simulations first we have to design a reaction coordinate which describes this conformational change. Principal component analysis (PCA) is a very useful tool to identify, isolate and describe the large scale motions present in proteins. PCA was performed on chain-A of 5TIM (open loop-6) and chain-B of 6TIM (closed loop-6) X-ray structures, using the backbone atoms of the entire protein, excluding all the dimer interface loops (loops 1 to 4), loop-5, 7 and 8 along with N (resid 1 to 10) and C (241 to 249) terminus fragments. Eigenvector-1 (Ev-1) described the motion of loop-6 from open to closed conformation. All the TIM ensembles were projected along the Ev-1. In Figure 3.11 to and fro sampling of open and closed states is seen along Ev-1. Using the probability distribution along Ev-1 we can estimate the  $\Delta$ G between the open and closed states of loop-6 in various ensembles from equation 1.4 (Figure 3.12).



Figure 3.11: In panel-1 and panel-2 W169-S212 (blue), V176-S212 (red) distance evolution along time starting with a CTIM trajectory are shown. In the third panel we have plotted the projection of the same trajectory onto Eigenvector-1 (Ev-1, magenta) describing loop-6 motion. In the fourth panel we have the loop-6 (L6) RMSD showing that indeed when loop-6 is closed based on W169-S212, V176-S212 and Ev-1 of loop-6, rmsd is smaller than or equal to 0.2 nm. Dotted blue line is the N-terminus/C-terminus distance in the X-ray closed structure and dotted green line in X-ray open structure. Dotted cyan line is the RMSD of X-ray open loop-6 versus X-ray closed loop-6.

We have combined both Apo-O and Apo-C ensembles to create APO ensemble because we observe similar behavior of loop-6 in both the ensembles. We have done the same for DHAP-O and DHAP-C ensembles. The free energy along the Ev-1 was calculated using the entire ensemble data (Figure 3.12). The error in free energy was calculated by estimating the free energy from each trajectory of the ensemble and bootstrapping the obtained free energy values 10000 times as described in section 2.1.3. The height of the error bars reflects the measure of sampling in MD simulation at that particular point along the Ev-1. As more

1.4.

trajectories sample a particular point, the height of the error bars become smaller and regions where sampling is not sufficient we have larger error bars.

In APO TIM ensemble we observe two minima, the first minimum at OTIM X-ray structure (-1.2 nm, green dotted line) between -0.6 nm and -1.4 nm (Figure 3.12a). A second shallow minimum is observed at the CTIM X-ray structure (1.3 nm, blue dotted line) between 0.9 nm and 1.5 nm. The free energy difference between open and closed loop-6 in APO TIM ensembles is  $1.35\pm0.36$  kcal.mol<sup>-1</sup> which is in the range of experimentally predicted values of 1.23 kcalmol<sup>-1</sup> and 1.77 kcal.mol<sup>-1</sup> (Table 3.7), suggesting that the population distribution of open and closed loop-6 structures is not different from population distribution in NMR experiments<sup>35,37,39</sup>.



Figure 3.12: Loop-6 free energy profiles from various ensembles; apo, DHAP, EDT1, EDT2, 2PG, apo unit cell simulations started with open structure (CS-O), started with closed structure (CS-C) and ligated 2PG unit cell simulation ensembles. Dotted blue line is the reference eigenvalue along eigenvector-1 for the X-ray closed structure and dotted green line for X-ray open structure.

 $\Delta$ G calculated from ligated ensembles of DHAP, 2PG, EDT1, EDT2 is between 3.14±0.38 kcal.mol<sup>-1</sup> to 4.96±0.22 kcal.mol<sup>-1</sup> (Figure 3.12a, b). The difference between the  $\Delta$ G calculated from holo TIM simulations and the NMR holo ensemble can arise because of sampling issue i.e. the 300 ns simulation time for each trajectory might not be long enough to observe the ligand induced closed conformation for loop-6.

The  $\Delta$ G calculated from apo unit cell simulations of CS-O and CS-C which start with OTIM and CTIM structure is 2.4±0.2 kcal.mol<sup>-1</sup> favoring open loop-6 and -2.1±0.33 kcal.mol<sup>-1</sup> favoring closed loop-6 (Figure 3.12c). This is a surprising result especially the  $\Delta$ G from CS-C ensemble which suggests that in the X-ray crystal environment the structures that are trapped in closed conformation have to work against the free energy gradient to reach the open state. For loop-6 to open or close in crystal unit cell, water surrounding the protein has to be displaced and since the solvent dynamics is much slower than in the normal MD simulations, starting conformation of the simulation is favored much more. We do observe sampling at but not any minima at reference CTIM X-ray structure for CS-O ensemble and at the OTIM Xray structure for CS-C ensemble.

CS-2PG simulations start with a CTIM structure with 2PG bound to the active site pocket. The minima for closed loop-6 conformation, is seen between 0.6 nm and 0.9 nm (Figure 3.12c). RMSD analysis of loop-6 of structures from 0.6 nm to 0.9 nm of eigenvector revealed that loop-6 conformation resembles the X-ray closed loop-6 conformation (Figure 3.13). In CS-2PG crystal unit simulations closed loop-6 conformation is favored by  $-5.24\pm0.14$  kcal.mol<sup>-1</sup> (Figure 3.12c).



Figure 3.13: W169-S212 (blue), V176-S212 (red) distance evolution along time starting with a CTIM structure from CS-2PG ensemble trajectory. In the third panel we have plotted the projection of the

same trajectory onto Eigenvector-1 (magenta) which describes loop-6 motion. In the fourth panel we have plotted the loop-6 (L6) RMSD (orange) showing that indeed when loop-6 is closed based on W169-S212, V176-S212 distances and projection of loop-6, rmsd is smaller than or equal to 0.2 nm and when loop-6 is open RMSD of loop-6 resembles X-ray open structure.

Williams et al.<sup>35</sup> suggested that the  $\Delta G$  between the 2PG ligated closed loop-6 and an unligated open conformation is -6.8 kcal.mol<sup>-1</sup> (Figure 1.5) if the binding free energy of 2PG ( $G_{bind} = -6.8 \text{ kcal.mol}^{-1}$ ) is taken into account. Just based on loop-6 population ratios reported by Williams et al.<sup>35</sup> and Massi et al.<sup>57</sup> the  $\Delta G$  in G3P/2PG ligated TIM would be between - 1.23 kcal.mol<sup>-1</sup> and -1.83 kcal.mol<sup>-1</sup>. To calculate the contribution of  $G_{bind}$  on  $\Delta G$  we have to calculate the  $G_{bind}$  of each DHAP/EDT1/EDT2/2PG for open and closed loop-6 conformations, which would be the focus of future work.

#### 3.4.7 Free energy of Loop-6

The  $\Delta$ G calculated from the apo TIM ensembles match the experimental values calculated from NMR population ratios. But the  $\Delta$ G from holo TIM simulations of DHAP, EDT1, EDT2 and 2PG are in disagreement with the NMR data which might be because of sampling issue as suggested earlier in section 3.4.6. To overcome the sampling problem we used enhanced sampling technique called Umbrella sampling (US) which was introduced in section 2.1.2.

We have demonstrated that loop-6 Ev-1 describes the transition of loop-6 between open and closed conformations (Figure 3.11). So it was chosen as the coordinate for performing umbrella sampling simulation (US) to calculate the free energy difference ( $\Delta G$ ) between open and closed conformations. The main reason behind excluding the dimer interface loops 1-4 and loop-5, loop-7 and loop-8 was the assumption that the regions of the protein which are correlated with loop-6 motion will adopt to the conformational change of loop-6 and also to avoid forcing of any region of protein along the reaction coordinate which does not contribute to this conformational change of loop-6.

Starting structures for each simulation were taken from eight individual trajectories with structures corresponding to eigenvalues along the reaction coordinate Ev-1. We performed totally eight US simulations, six apo (F5TIM-B, F6TIM-B, 5TIM-A, 5TIM-B, 6TIM-A and 6TIM-B) and two 2PG ligated simulations, 1N55-A and 1N55-B. We used 2PG in our US simulations over DHAP because of its longer residence time in the active site pocket and availability of NMR experimental data for comparison. As described in section 2.1.2

umbrella windows were created between -1.60 nm and 1.60 nm with an interval of 0.2 nm. The OTIM X-ray structure projection on Ev-1 is at -1.22 nm and CTIM X-ray structure is at 1.29 nm.

All the trajectories from which structures were picked for US simulations, have sampled both open and closed states. If a particular part along the reaction coordinate was not sampled in MD simulation, starting structure for that particular interval was created by linear interpolation. A harmonic restraining potential of 259 kcal.mol<sup>-1</sup> was used for each umbrella window to restrict the each structure at chosen point along the reaction coordinate.

The first 60 ns of each trajectory in US simulation was discarded as equilibration and only data from 60 ns to 110 ns was used to calculate the free energy profiles. US simulations from OTIM-B, CTIM-B, 6TIM-A, 6TIM-B, 5TIM-B and 1N55-A simulations will be referred to as loop-5 open US simulations (L5O) and 5TIM-A, 1N55-B as loop-5 closed US simulations (L5C). As demonstrated earlier in section 3.4.4, loop-5 motion is correlated with loop-6 N-terminus. A profound effect of loop-5 on loop-6 N-terminus conformation is seen in L5O US simulations. In L5C US simulations, in structures along the reaction coordinate from 0.6 nm to 1.4 nm where loop-6 is in closed conformation, loop-5 is also in closed state (Figure 3.14a). But in L5O US simulations, in the region between 0.6 nm to 1.4 nm, loop-5 adopts open conformation which causes loop-6 N-terminus to open (Figure 3.14b). This semi closed loop-6 where N-terminus is open, because loop-5 is open, causes an increase in free energy difference between open and closed loop-6 states (Figure 3.14b).



Figure 3.14: Free energy profiles from Umbrella sampling simulations from a) L5-Closed profiles and b) L5-Open profiles. In panel two and three each point represents the average of

loop-6 RMSD with respect to the closed loop-6 (L6) in CTIM X-ray structure and loop-5 (L5) projection in eigenvector-1 (Ev-1) of loop-5 pca, which shows loop-5 conformational change between open (green) and closed (blue) loop-5. In panels 1 and 3 dotted blue line is the reference eigenvalue along the eigenvector-1 for the X-ray closed structure and dotted green line for the X-ray open structure. In panel-2 dotted green line represents the RMSD of X-ray open loop-6 with respect to X-ray open loop6.

The  $\Delta G$  calculated from apo US simulation ( $\Delta G_{apo}$ ) of 5TIM-A trajectory is 1.15±0.5 kcal.mol<sup>-1</sup> (Figure 3.14a) which is comparable to the experimental value which lies between 1.23 kcalmol<sup>-1</sup> and 1.77 kcal.mol<sup>-1</sup> (Table 8). Whereas the  $\Delta G$  calculated from 2PG ligated 1N55-B US simulation ( $\Delta G_{2PG}$ ) is -3.4±0.2 kcal.mol<sup>-1</sup> (Figure 3.14a) is far from the experimental estimate of -6.8 kcal.mol<sup>-1</sup> and requires longer simulation time for each umbrella. Since we find profound influence of loop-5 on conformation of N-terminus of loop-6, a new reaction coordinate which includes loop-5 as well is required.

# 3.4.8 Comparison with NMR

We have demonstrated that loop-6 transiently samples both open and closed conformations of loop-6 in apo and holo TIM ensembles. We have also defined loop-6 into five conformations and it is essential to segregate these conformations into NMR-Open (NMR-O) and NMR-Closed (NMR-C) since in NMR experiment only open and closed conformations are identifiable. A lot of experimental information is available on the dynamics of loop-6 in form of chemical shifts, relaxation dispersion data. We calculated the NMR chemical shifts and angular hop of W169 indole ring with respect to Z-axis from our MD simulations to compare with NMR data.

### 3.4.8.1 Calculation of Chemical Shifts

Chemical shifts are highly sensitive to the local environment of a protein residue and are used as reporters of change in conformation of a protein<sup>36,39,41</sup>. SHIFTX2<sup>144</sup> is the latest chemical shift prediction program which has three times higher accuracy compared to other available software SHIFTX<sup>145</sup>, SPARTA+<sup>146</sup>, SHIFTS<sup>147,148</sup>, CAMSHIFT<sup>149</sup> and PROSHIFT<sup>150</sup>. SHIFTX2 achieves higher accuracy (three times smaller mean square error) compared to other available programs by combining both structure and sequence based prediction approach.

Berlow et al.<sup>39</sup> from their NMR spin relaxation experiments reported backbone nitrogen and amide hydrogen's chemical shift of V167 present in N-terminus of loop-6, T172 and T177

present in C-terminus of loop-6 in chicken TIM. To test the prediction power of SHIFTX2, we used X-ray structure from Rabbit (1R2R) which was used in MD simulations and chicken (1TPH) TIM structure as control because chicken TIM was used in NMR experiments. Rabbit TIM has 88% overall sequence identity and 100% loop-6 sequence identity with chicken TIM. Chain-B of 1R2R and chain-1 of 1TPH are closed and 1R2R chain-A is an OTIM structure. In NMR experiments the sample was at pH 7.5, deuterated and the data was collected at 298 K. Chemical shifts by SHIFTX2 were calculated using the same conditions.

Backbone nitrogen chemical shifts predicted by SHIFTX2 for residues V167, T172 and T177 which were monitored in the experiment do not differ between OTIM and CTIM structures (Table 3.8). For example chemical shift from NMR experiment for V167 in OTIM is 130.68 ppm but the predicted chemical shift for 1R2R open and CTIM structures is ~125 ppm i.e. open and closed loop-6 cannot be differentiated based on the predicted chemical shifts (Table 3.8). Since the predicted chemical shifts from X-ray OTIM and CTIM crystal structures do not match with the experimental data we do not continue further with this analysis.

Table 3.8: Comparison of SHIFTX2 predicted backbone nitrogen chemical shifts (ppm), for 1R2R and 1TPH crystal structures. Only data from A and B chains of 1R2R and chain-1 of 1TPH are shown. Chain-C and D of 1R2R and chain-2 of 1TPH show similar chemical shifts.

<b>Residue</b> \Structure	NMR-Open	NMR-Closed	1 <b>R2R-A</b>	1R2R-B	1TPH-1
V167	130.68	127.00	123.81	125.00	125.34
T172	108.01	106.43	106.64	106.26	106.58
T177	115.82	112.89	113.85	112.17	111.84

#### 3.4.8.2 W169 angular hop

Ann McDermott and co-workers<sup>35,37</sup> used W169 as an indicator of loop-6 conformational change. The angle between the W169 indole ring and the Z-axis differs by  $\sim 30^{\circ}$  between OTIM (145°) and CTIM (114°) structures. We have calculated the angle between W169 indole ring and the Z-axis of the simulation box for every structure in the MD simulation. Based on loop-6 W169-S212 and V176-S212 distances, we have previously divided loop-6 into five conformations (Table 3.6). We calculated the distribution of W169 angle for each conformation of loop-6 in all ensembles (Figure 3.15).

For Open and IMD (loop-6 is neither closed nor open; Table 3.7) loop-6 conformations, W169 angle corresponds to open W169 angle of 145°, which is seen in OTIM X-ray structure.

When N-terminus of loop-6 is closed or both termini of loop-6 are closed we observe that W169 angle corresponds to CTIM X-ray structure value of 114° (Figure 3.15). W169 is present in N-terminus of loop-6 and it was expected that W169 would be only a reporter of closure of loop-6 N-terminus and entire loop-6. Surprisingly in crystal unit cell simulations and EDT1, EDT2 ligated TIM simulations, W169 angle corresponds to closed state even when C-terminus of loop-6 is closed and N-terminus is open. So we have combined Open, IMD as NMR-Open loop-6 states and C-terminus closed, N-terminus closed and loop-6 closed as NMR-Closed loop-6 states.



Figure 3.15: Distribution of W169 angle with respect to Z-axis for each of the five conformations of loop-6 in apo TIM ensemble (a), EDT2 ensemble (b), apo open unit cell simulation CS-O (c) and 2PG ligated unit cell simulation (d). When both termini of loop-6 are closed (Closed) or N-termini is closed (N-Closed) or C-termini is closed (C-Closed) we observe that W169 angle distribution shifts towards CTIM X-ray W169 angle (114°). W169 angle distribution for loop-6 intermeditae conformation (IMD) and open conformation corresponds to OTIM W169 angle (145°).

For each NMR-O and NMR-C conformation from MD ensembles, we calculated the average W169 angle with respect to Z-axis (Figure 3.16). In our simulations, in both apo and holo TIM

ensembles the difference in W169 angle ( $\Delta$ W169A) between NMR-O and NMR-C conformations is 6° to 10° (Figure 3.16) with k<sub>ex</sub> of 10<sup>6</sup> s<sup>-1</sup>. The  $\Delta$ W169A from apo TIM ensembles matches the SSNMR data from Rozovsky et al.<sup>37</sup> in 2001.

Rozovsky et al.<sup>37</sup> modeled the loop-6 conformational change in apo TIM sample with a low angular motion of 2° for W169 and  $k_{ex}$  of  $10^6 \text{ s}^{-1}$  and in G3P ligated sample with a larger W169 angular motion of 20° to 40° with  $k_{ex}$  of  $10^4 \text{ s}^{-1}$ . Since we do not observe a preference for closed loop-6 in ligated TIM structures, the  $\Delta$ W169A from holo TIM ensembles does not match the expected value reported by Rozovsky et al.<sup>37</sup>.



Figure 3.16: Average W169 angle with respect to Z-axis. On X-axis is the loop-6 population percentage in closed state (a) and open state (b).

Williams et al.<sup>35</sup> proposed a population ratio in the range of 1:8 to 1:20 for minor and major conformations with  $\Delta$ W169A of 20° to 50°. In MD ensembles the population of the major conformation NMR-O is ~90% and of the minor conformation NMR-C is less than 10% except in CS-2PG (NMR-C 75%, NMR-O 25%) and CS-C (NMR-C 60%, NMR-O 40%) ensembles.

If we consider the ensembles with NMR-O/NMR-C populations of greater than 75% the  $\Delta$ W169A value is 26° (Figure 24). This would imply that in SSNMR experiment if a sample has 100 OTIM structures, greater than 75% of the structures have to adopt closed loop-6 to observe a change in the line shape. We observe this in CS-C ensemble which starts with closed loop-6, 40% of the ensemble adopts NMR-O and 60% NMR-C conformation with W169 angles of 130° and 122° (Figure 24) and we would expect that as the population of NMR-O increases W169 angle would reach 145°. In apo TIM ensembles we do not see such a shift in population of loop-6 conformation.

#### 3.4.9 Discussion

Loop-6 has a very important role to play in the reaction mechanism catalyzed by TIM. Based on X-ray structures loop-6 was proposed to exist in open and closed conformations. In apo TIM open loop-6 is the major conformation and in ligated TIM it is the closed loop-6. Closure of TIM loop-6 excludes water from the active site lowering the dielectric constant of which strengthens the electrostatic stabilization of the transition state<sup>20</sup>. Closure of loop-6 was also proposed to clamp the catalytic E166 between the two hydrophobic residues I171 and L231, which increases its basicity.

We find that as expected by Rozovsky et al.<sup>36,37</sup>, in apo TIM loop-6 does sample open and closed conformations at  $10^6$  s<sup>-1</sup> with W169 angular hop of  $6^{\circ}\pm 2^{\circ}$  (experimental value  $2^{\circ}$ ; Figure 3.16). But when the apo TIM simulations (APO) start with a closed TIM structure (CTIM), a conformational change to open loop-6 is seen within the first 10 ns. Once the loop-6 in CTIM structures open up, for the rest of the simulation no difference is seen in loop-6 dynamics between simulations that start with open TIM (OTIM) structure and that start with CTIM structure.

In apo OTIM crystal unit cell simulations (CS-O) and in CTIM crystal unit cell simulations (CS-C), loop-6 like in APO ensemble samples open and closed loop-6 conformations at microsecond time scale. Unlike the normal MD simulations that start with CTIM structure, loop-6 stays closed for greater than 100 ns (Figure 3.3d) with 60% of closed loop-6 conformation whereas the same in APO ensemble is 8% (Figure 3.16a). As expected, this is because of slow solvent dynamics in the crystal unit cell simulations (Table 3.4). On the other hand in CS-O percentage of open and closed loop-6 populations are 92% and 8% as seen in APO ensemble (Figure 3.16b).

From Principal component analysis (PCA) using X-ray 5TIM and 6TIM structures we found eigenvector-1 (Ev-1) which described loop-6 conformational change. Using the distribution of projection along Ev-1, the free energy difference ( $\Delta G$ ) between open and closed loop-6 calculated from APO ensemble ( $1.35\pm0.36$  kcal.mol<sup>-1</sup>; Figure 3.12a) and apo loop-6 umbrella sampling (US) simulation ( $1.15\pm0.5$  kcal.mol<sup>-1</sup>; Figure 3.14a) is within the NMR experimental<sup>35,39</sup> range of 1.23 kcal.mol<sup>-1</sup> to 1.77 kcal.mol<sup>-1</sup> which suggests that the population ratios of the major open and minor closed loop-6 conformation in apo TIM MD (9:1) and NMR (8:1 to 20:1) are identical. The  $\Delta$ G calculated from CS-O (2.4±0.2 kcal.mol<sup>-1</sup>; Figure 3.12c) is higher than the NMR experimental<sup>35</sup> value because of five times lesser transitions compared to the APO ensemble (Figure 3.10). Whereas the  $\Delta$ G calculated from CS-C (-2.1±0.33 kcal.mol<sup>-1</sup>; Figure 3.12c) favors closed loop-6 with a very strong minima observed at CTIM X-ray structure (Figure 3.12c) which implies that the crystal environment stabilizes closed loop-6.

The crystal unit simulations that start with apo 1R2R crystal structure from Rabbit at 100 K in which chains A, C and D are open and only chain-B is closed. The two other crystal structures 1R2T and 1R2S from the same organism but obtained at 85 K and 298 K do not show the same closed loop-6 conformation. Aparicio et al. pointed out to a water molecule which is present in place of the phosphate moiety of the ligand that might stabilize this closed loop-6 and further suggested that effect of flash freezing cannot be overlooked. Effects of flash freezing have been reported for side chain conformations of residues<sup>137,139</sup> but not for such a large conformational change.

In the holo TIM simulations with natural substrate DHAP, inhibitor 2PG, reaction intermediates EDT1 and EDT2 we do not observe any conformational preference for closed loop-6 as expected from the X-ray data. The population ratio of open and closed loop-6 in normal holo TIM simulations is similar to apo TIM simulations with less than 10% of closed loop-6. Only in the 2PG ligated crystal unit cell simulation (CS-2PG) which starts with a CTIM structure, 75% of the ensemble has closed loop-6 and the rest 25% is open loop-6.

Based on NMR studies the expected  $k_{ex}$  between open and closed conformations in G3P ligated TIM sample is  $10^4$  s<sup>-1</sup>. This means that in the presence of the ligand, closure of loop-6 takes much longer time (once in 100 µs) than in the apo TIM structures (once in 1 µs). And our simulations with ligand are only 300 ns long. Though, in loop-7 we observe DHAP induced closed state (starting with an OTIM structure) in 50 ns of simulation time which is discussed in section 3.5. Such a spontaneous effect on conformation of loop-6 when ligand is bound to the active site, is not seen in our MD simulations. It is also possible that ligand induces conformational change from open to closed state in loop-7 first and loop-6 might follow later, which is not seen in our 300 ns long MD simulations.

The  $\Delta G$  calculated from ligated ensembles is between 3.14±0.38 kcal.mol<sup>-1</sup> and 4.96±0.22 kcal.mol<sup>-1</sup> (Figure 3.12 a, b) which is not close to the experimental value of -6.8 kcal.mol<sup>-1</sup>. Only the  $\Delta G$  calculated from CS-2PG (-5.24±0.14 kcal.mol<sup>-1</sup>; Figure 3.12c) is comparable to the experimental value. We performed two 2PG ligated US simulations 1N55-A and 1N55-B

to calculate loop-6 free energy difference while 2PG is bound to the active site. The  $\Delta G$  calculated from 1N55-A ensemble was ~2.7 kcal.mol<sup>-1</sup> and from 1N55-B it was -3.4 kcal.mol<sup>-1</sup> which is an improvement over normal MD simulations, but still not closed to the experimental value. The positive  $\Delta G$  from 1N55-A is because of loop-5 open conformation in the CTIM structures along the reaction coordinate. A new reaction coordinate which also includes loop-5 conformational change is required to calculate more reliable estimates of  $\Delta G$  from apo and holo TIM structures.

TIM catalyzes a reversible isomerization reaction where binding of DHAP results in formation of GAP and vice versa. A ligand induced conformational change for loop-6 as proposed by Richard and co workers and X-ray data raises a very important question. How does TIM initiate opening of loop-6 once the reaction is completed? Donnini et al.<sup>151</sup> has proposed a mechanism where closure of loop-6 stores energy as strain in ring of P167. Upon completion of reaction this energy is used to open loop-6. Such a mechanism also needs an instigator which signals completion of reaction, followed by opening of loop-6 and release of product.

A different proposition can be made where even when the substrate molecules are bound to the active site, loop-6 undergoes a conformational change between open and closed conformations with the same  $k_{ex}$  of apo TIM ( $10^6 \text{ s}^{-1}$ ). In this scenario, when the substrate is bound to the active site, closure of loop-6 will enable the catalysis of the reaction. Since conformation of loop-6 is not ligand induced, spontaneous opening of loop-6 will be followed by release of product and this will not require any trigger for initiating opening of loop-6, once it is closed.

In our simulations we have monitored the dynamics of N-terminus (P167-V168-W169) and C-terminus hinges (K175-V176-A177) of loop-6 using distances between W169 and loop-7 residue S212 (W169-S212), V176 of loop-6 and S212 (V176-S212). Based on these W169-S212 and V176-S212 distances we divided loop-6 into five conformations (Table 3.6). Not only we observe different populations for the N and C-terminus hinges in open and closed conformations but also the correlation analysis between W169-S212 and V176-S212 distances has revealed that N and C-terminus hinges are uncorrelated i.e. loop-6 motion is not rigid.

N-terminus of loop-6 interacts with loop-5 whereas C-terminus of loop-6 is solvent exposed. For N-terminus of loop-6 to open loop-5 has to also adopt open conformation. Hence higher correlation of 0.5 to 0.6 is seen between loop-5 and loop-6 N-terminus compared to the 0.2 seen between N-terminus and C-terminus distances. This effect of loop-5 is much more pronounced in US simulations.

The Ev-1 which was used as the reaction coordinate for US simulations only describes the loop-6 conformational change but not loop-5. The reason for leaving out loop-5 from Ev-1 was that if loop-5 conformational change is strongly correlated with loop-6, loop-5 will adopt to loop-6 conformational change. But in US simulations we observe that when loop-5 is in open conformation, loop-6 N-terminus hinge also adopts open conformation and does not stay closed. This suggests that loop-5 closure is essential to stabilize N-terminus of loop-6 in closed conformation.

To verify this we have performed simulations with 6TIM X-ray structure where chain-B is closed and chain-A is open. The first 150 residues of the protein which includes loop-5 (residues 127 to 137) were position restrained. In these simulations, when loop-5 is restrained in closed conformation, loop-6 N-terminus stays closed but loop-6 C-terminus and tip undergo conformational change to open state. When loop-5 is restrained in open conformation in chain-A, both loop-6 N-terminus and C-terminus sample open, closed conformations.

We do not find any influence of any ligand molecule on conformation of loop-6 irrespective of its reactive complex (RC) or encounter complex (EC) orientation in the active site. From the analysis of X-ray structures we found out that when any ligand molecule is bound in EC state loop-6 is not present in closed conformation. In simulations with DHAP and CS-2PG when ligand is present in EC orientation closed conformation for loop-6 is observed. Phosphate moiety of loop-6 interacts with the tip of loop-6 (A170 to G174). We observe that closed conformation of N-terminus and C-terminus hinges is not hindered by phosphate moiety of ligand as is seen for tip of loop-6 (Figure 3.1). When ligand is in EC orientation, tip of loop-6 opens in our simulations but N-terminus and C-terminus are not affected.

In all the OTIM X-ray structures, loop-5, 6 and 7 are in open conformation and in CTIM structure all the three loops are in closed conformation. In our apo TIM simulations when loop-7 (which will be discussed in 3.5) does not sample the closed conformation, loop-6 undergoes a conformational change between open and closed conformations. Because of this reason it might be possible that the closed conformation sampled in the NMR experiments (both loop-6 and loop-7 are closed) and MD simulations might be different. To confirm this,

NMR studies which can study the conformational changes of loop-5, loop-6 and loop-7 are required.

# 3.5 Dynamics of loop-7

Role of Loop-7 was discussed in section 1.3. The backbone dihedral angles phi (C'-N-C $\alpha$ -C"), psi (N-C $\alpha$ -C- N') of residues G210, G211, S212 and V213 differ between open and closed conformations based on which loop-7 is classified into open and closed states (Table 3.1). Loop-7 is found in open conformation in apo TIM structures and closed conformation is adopted in presence of the ligand molecules (Figure 3.17b). Among all the apo TIM X-ray structures, loop-7 adopts closed conformation only in chain-B of 1R2R crystal structure. A water molecule is present in place of the oxygen of the phosphate moiety of the ligand, stabilizing the closed conformation of G211-psi (G211 $\Psi$ ) and S212-phi (S212 $\Phi$ ) dihedral angles in the chain-B of 1R2R (Figure 3.17b).



Figure 3.17: a) Open (green) and closed (blue) conformations of Loop-7. b) Loop-7 in closed conformation with DHAP bound to the active site. Hydrogen bond between amide group of S212 and phosphate moiety of DHAP along with hydrogen bond between the hydroxyl group of Y209 and carbonyl oxygen of G211 are only possible when G211 $\Psi$  and S212 $\Phi$  dihedral angles flip to closed state. c) In chain-B of 1R2R crystal structure a water molecule (blue oxygen and green hydrogens) is present in place of the oxygen of the phosphate moeity of DHAP. This water stabilizes closed conformation of S212 $\Phi$ . DHAP is shown in superposition with the water molecule.

We have performed multiple apo TIM and holo TIM simulations both in normal solvent and crystal unit cell simulations as described in section 2.2. We monitored the backbone dihedral angles G210-psi (G210 $\Psi$ ), G211-phi (G211 $\Phi$ ), G211 $\Psi$ , S212 $\Phi$ , S212-psi (S212 $\Psi$ ) and V213-phi (V213 $\Phi$ ) in various apo and holo TIM ensembles. G210 $\Psi$ , G211 $\Phi$ , S212 $\Psi$  and V213 $\Phi$  sample open and closed conformations in apo and holo TIM simulations (Figure 3.18a, b). Whereas G211 $\Psi$  and S212 $\Phi$  sample closed conformation only in presence of a

water molecule located at a specific position or when the ligand molecules are bound to the active site (Figure 3.18 b). In this thesis we would like to mainly focus on the dynamics of G210 $\Psi$  and G211 $\Phi$  and its effect on conformation of catalytic residue E166 which is discussed in section 3.5.1 and in section 3.5.2 we discuss the dynamics of dihedral angles G211 $\Psi$  and S212 $\Phi$ .



Figure 3.18: Dynamics of G210 $\Psi$ , G211 $\Phi$ , G211 $\Psi$ , S212 $\Phi$ , S212 $\Psi$  and V212 $\Phi$  dihedral angles in an apo OTIM simulation (a) and DHAP ligated OTIM simulation (b). In both the simulations all the loop-7 dihedral angless start in open conformation. Apart from G211 $\Psi$  and S212 $\Phi$  rest of the dihedral angles sample open and closed conformations in both apo and holo TIM simulations. Closed conformation of G211 $\Psi$  and S212 $\Phi$  is ligand induced and in apo TIM simulations they prefer open conformation. The reference values for X-ray open (green dotted line) and X-ray closed (blue dotted line) conformations for each dihedral angles.

#### 3.5.1 Dynamics of G210Ψ and G211Φ

In both apo and holo TIM ensembles no effect of any ligand molecule is seen on the dynamics of G210 $\Psi$  and G211 $\Phi$  dihedral angles. Casteleijn et al.<sup>14</sup> proposed that, on ligand binding the closed conformation of G210 $\Psi$  induces a change in conformation of catalytic residue E166 from non reactive 'swung-out' to reactive 'swung-in' conformation (Figure 3.19a). The clash between the carbonyl oxygen of G210 and P167 side chain was proposed to initiate this conformational change.

To test this hypothesis, we have monitored the dihedral angles of  $G210\Psi$  and  $G211\Phi$  along with the conformational change of E166. PCA was performed on the chain-A of 5TIM (open; Figure 3.19a) and chain-B of 6TIM (closed; Figure 3.19a). The first eigenvector (Ev-1) from

the PCA described the conformational change of E166 from swung-in to swung-out conformation (Figure 3.19b).



Figure 3.19: a) E166 swung-in (blue) conformation in CTIM and swung-out (green) conformation in OTIM. b) Conformational change from swung-in (blue) to swung-out (green) oritentation of E166-P167 along the first eigenvector from PCA.

We have monitored the dynamics of E166 along with the dihedral angles G210 $\Psi$  and G211 $\Phi$ . G210 $\Psi$  and G211 $\Phi$  are correlated i.e. when G210 $\Psi$  is in open conformation, G211 $\Phi$  is also in open conformation and vice versa (Figure 3.20). Since E166 is present in the beginning of N-terminus of loop-6, we would expect that if E166 conformation is affected by loop-6 conformation then N-terminus of loop-6 will have the dominant effect. But we do not see such an effect of N-terminus on conformation of E166. E166 can sample both open and closed conformations independent of loop-6 N-terminus conformations (Figure 3.20) and same behaviour is seen across all TIM ensembles that we have simulated.

E166 conformation is dependent on the conformations of dihedral angles G210 $\Psi$  and G211 $\Phi$ . In simulations where G210 $\Psi$  and G211 $\Phi$  stay in open conformation for the entire length of the simulation, E166 does not sample the swung-in conformation (data not shown). A deviation from the X-ray closed state for G210 $\Psi$  and G211 $\Phi$  dihedral angles causes a change in the E166 conformation from swung-in to swung-out conformation (Figure 3.20b). To determine the orientation of G210 $\Psi$  and G211 $\Phi$  dihedral angles when E166 is in swung-in or swung-out conformation, we calculated the average of G210 $\Psi$  and G211 $\Phi$  dihedral angles for both swung-in and swung-out conformations of E166. A conservative cut-off of 0.3±0.3 nm for swung-out and -0.3±0.3 nm for swung-in was used to bin the dihedral angles assuming a two state population for E166 conformation.



Figure 3.20: Dynamics of G210 $\Psi$ , G211 $\Phi$  are shown in the first and the second panel. In the third panel, eigenvector-1 (Ev-1) from PCA of X-ray structures which describes the conformational change of E166 between CTIM swung-in conformation (dotted blue line) and OTIM swung-out conformation (green dotted line) is shown. Distance between C-alpha atoms of W169 present in N-terminus of loop-6 and loop-7 residue S212 is shown in fourth panel. In an apo CTIM simulation (a) and EDT1 ligated CTIM structure simulation (b) the change in conformation of G210 $\Psi$  and G21 $\Phi$  also causes a change in E166 conformation along Ev-1. The reference values for X-ray open and X-ray closed conformations in each panel are shown in dotted green line and dotted blue line.

If the conformation of dihedral angles G210 $\Psi$  and G211 $\Phi$  induce the conformational change in E166 conformation as seen from our data, then we would expect that the average value of the dihedral angle over the entire ensemble when E166 is in swung-in conformation should correspond to the X-ray closed value (Table 3.1) and when E166 is in swung-out conformation should correspond to X-ray open value (Table 3.1). Since we observe that dihedral angles G210 $\Psi$  and G211 $\Phi$  are correlated, here we only present the average values calculated for G210 $\Psi$  dihedral angle over the entire ensemble when E166 is in swung-in and swung-out conformations.

In Figure 3.21 along x-axis we have shown the average value of  $G210\Psi$  angle and along yaxis the population percentage of E166 in swung-in (Figure 3.19a) or swung-out conformation (Figure 3.19a) in various TIM ensembles. As we expected above, we indeed
observe that whenever E166 is in swung-in conformation G210 $\Psi$  dihedral angle corresponds to the X-ray closed state (Figure 3.21a) and when E166 is in swung-out conformation dihedral angle of G210 $\Psi$  samples the X-ray open state (Figure 3.21b). The large error bars for the population percentage represent the large variation in the population of E166 in swung-in or swuing-out conformation, suggesting that more sampling is required especially for the TIM ensemble with EDT1 bound to the active site pocket (Figure 3.21b). In CS-2PG simulation the percentage of Swung-in conformation for E166 is ~3% which might account for the deviation of average of G210 $\Psi$  dihedral angle from X-ray open state (Figure 3.21b).



Figure 3.21: Average dihedral angle values of  $G210\Psi$  in various TIM ensembles when E166 is in swung-in conformation (a) and swung-out conformation (b).  $G210\Psi$  average open (green dotted line) and closed (blue dotted line) dihedral angle values from analysis of OTIM and CTIM X-ray structures (see section 3.1). Y-axis shows the population percentage of E166 in swung-in or swung-out conformations over the entire ensemble. Error bars were calculated from bootstrap analysis.

#### 3.5.2 Dynamics of G211 $\Psi$ and S212 $\Phi$

Dihedral angles G211 $\Psi$  and S212 $\Phi$  sample the closed conformation only when the ligand molecules are bound to the active site (Figure 3.18) or when a water molecule is present in place of the oxygen of the phosphate moiety of the substrate. In absence of this water molecule or the ligand, G211 $\Psi$  and S212 $\Phi$  are present in open conformation. The effect of water molecules and ligand on the conformation of G211 $\Psi$  and S212 $\Phi$  are further discussed in section 3.5.2.1 and 3.5.2.2.

#### 3.5.2.1 Effect of water

Apo OTIM and CTIM simulations were started using 5TIM and 6TIM X-ray structures. In apo OTIM simulations, out of 13 chains only in one chain G211 $\Psi$  and S212 $\Phi$  flip from open to closed state (stays closed for 10 ns) and back to open conformation. In apo CTIM simulations, out of 25 chains, in five chains, G211 $\Psi$  and S212 $\Phi$  transiently sample open and closed conformations. When G211 $\Psi$  and S212 $\Phi$  flip from open to closed conformation, a water molecule similar to the one in the 1R2R crystal structure (Figure 3.17b) was found in these apo TIM simulations (Figure 3.22b) which was not present in the beginning of the simulation. In rest of the apo OTIM and apo CTIM simulations, in absence of this water molecule at this particular position only open conformation is sampled by G211 $\Psi$  and S212 $\Phi$  dihedral angles.

The effect of water molecules on G211 $\Psi$  and S212 $\Phi$  conformation is much more profound in apo crystal unit cell simulations which were started with 1R2R crystal structure. Out of 120 open chains, in 23 chains we observe that G211 $\Psi$  and S212 $\Phi$  flip from open to closed state spontaneously. And in five chains G211 $\Psi$  and S212 $\Phi$  flip to closed state and stay closed for 175 ns to 270 ns of simulation time which is not seen in the normal apo OTIM and CTIM MD simulations of 5TIM and 6TIM crystal structures. In these 23 chains, a water molecule which was not present in the beginning of the simulation was found within the hydrogen bonding distance of amide group of S212 stabilizing G211 $\Psi$  and S212 $\Phi$  in closed state as seen in chain-B of the 1R2R X-ray structure (Figure 3.17b).



Figure 3.22: a) Apo crystal simulation where G211 $\Psi$  and S212 $\Phi$  start in open conformation and flip to closed conformation. The reference values for X-ray open and X-ray closed conformations in each panel are shown in dotted green line and dotted blue line. b) The closed conformation of G211 $\Psi$  and S212 $\Phi$  in this apo crystal simulation is stabilized by water molecules. Hydrogen bonds between loop-7 residues and water molecules are shown in dotted red and blue lines.

#### **3.5.2.2 Effect of ligand**

As discussed in section 2.2.2 we performed multiple holo TIM simulations with natural substrate DHAP, inhibitor 2PG, reaction intermediates EDT1 and EDT2 in the active site pocket. DHAP was placed in the active site of both OTIM and CTIM structures whereas rest of the ligand molecules were placed in the active site pocket of CTIM structures. All the ligand molecules in the beginning of the simulation are bound to the active site in reactive complex (RC) orientation (Figure 3.1a).

In section 3.4.1 we have shown that no influence of any ligand molecule is seen on the conformation of loop-6 i.e. even when the ligand molecules are bound to the active site pocket for entire length of simulation (~300 ns), loop-6 does not adopt the closed conformation. But in the holo TIM simulations which start with loop-7 in open conformation when DHAP is bound to the active site, dihedral angles G211 $\Psi$  and S212 $\Phi$  flip from open to closed states spontaneously (Figure 3.21). And when the ligand escapes into the solvent within the first 40 ns from OTIM chains, such an effect is not observed. In section 3.1, we proposed that the flip of G211 $\Psi$  and S212 $\Phi$  into closed state may be caused by the electrostatic repulsion between carbonyl oxygen of G211 and the negatively charged phosphate moiety of the substrate DHAP molecule.

We calculated the coulombic and Van der Waals interaction energy between the carbonyl oxygen of G211 and the amide group of S212 with phosphate moiety of natural substrate DHAP, reaction intermediates EDT1, EDT2 and the inhibitor molecule 2PG (Figure 3.21a). The electrostatic repulsion between the negatively charged phosphate moiety and the carbonyl oxygen of the G211 can be as large is 4 kcal.mol<sup>-1</sup> and this repulsion causes the rotation of G211 $\Psi$  (N-C $\alpha$ -C-N). This rotation around G211 $\Psi$  simultaneously induces S212 $\Phi$  into the closed conformation and brings amide group into the position to form hydrogen bond with phosphate moiety of the substrate. This flip to closed state provides additional -8 kcal.mol<sup>-1</sup> of stabilization energy for the ligand molecule in the active site (Figure 3.21a).



Figure 3.23: a) An OTIM structure which has DHAP bound in the active site pocket for the entire length of the simulation where  $G211\Psi$  and  $S212\Phi$  flip from open to closed conformation. In the third panel RMSD of DHAP calculated with respect to the DHAP in the starting structure of the simulation. In the fourth panel the coulombic interaction energy between the phosphate moiety of DHAP and G211 carbonyl oxygen, S212 amide group is shown. The reference values for X-ray open and X-ray closed conformations in panel-1 and panel-2 are shown in dotted green line and dotted blue line. In panel-3 dotted blue line represents the cut-off RMSD after which ligand molecule is out of the active site in the solvent. b) In OTIM simulation loop-7 adopts a closed conformation in presence of DHAP. X-ray closed loop-7 is shown in transperent blue licorice. Dotted lines represent the hydrogen bonds.

The contribution of van der Waals interaction energy is compared to coulombic interaction energy between G211, S211 and phosphate moiety of ligand molecules is small. When G211 $\Psi$  and S212 $\Phi$  are in open conformation because of the electrostatic repulsion between the phosphate moiety and carbonyl oxygen of G211, both the groups are apart which translates to a favorable Van der Waals interaction of less than -1 kcal.mol<sup>-1</sup>. But when G211 $\Psi$  and S212 $\Phi$  adopt closed conformation because of favorable electrostatic interactions (-8 kcal.mol<sup>-1</sup>) with phosphate moiety of ligand, steric clashes between phosphate oxygen and amide group of S212 increases Van der Waals energy up to ~3 kcal.mol<sup>-1</sup> with average Van der Waals energy less than 1 kcal.mol<sup>-1</sup> (data not shown). The free energy required to cause the flip of G211 $\Psi$  and S212 $\Phi$  from open to closed state remains to be calculated.

#### 3.5.3 Discussion

We have monitored the dynamics of dihedral angles  $G210\Psi$ ,  $G211\Phi$ ,  $G211\Psi$ ,  $S212\Phi$ ,  $S212\Psi$ and  $V212\Phi$  in various apo and holo TIM ensembles. Based on X-ray crystal structures it was proposed that closed conformation of loop-7 is ligand induced<sup>14,152</sup>. In our simulations we find that dihedral angles G210 $\Psi$ , G211 $\Phi$ , S212 $\Psi$  and V212 $\Phi$  undergo transitions between open and closed conformations in apo, holo and unit cell simulations (Figure 3.18) and dihedral angles G211 $\Psi$  and S212 $\Phi$  sample closed state only in presence of a ligand molecule (Figure 3.18b) or a strategically located water molecule (Figure 3.20) as seen in chain-B of 1R2R crystal structure (Figure 3.17b).

As proposed by Casteleijn et al.<sup>14</sup> we also find that conformation of G210 $\Psi$  and G211 $\Phi$  influence conformation of catalytic residue E166. Since we observe transitions between open and closed conformations for G210 $\Psi$  and G211 $\Phi$  in both apo and holo TIM simulations, as suggested by Casteleijn et al.<sup>14</sup> binding of ligand to the active site may not be nessaccary to induce closed conformation for G210 $\Psi$  and G211 $\Phi$ . Depending on the conformation of G210 $\Psi$  and G210 $\Psi$  and G211 $\Phi$ . Depending on the conformation of G210 $\Psi$  and G210 $\Psi$  and G211 $\Phi$ . Depending on the conformation and swung-out conformations in apo, holo and unit cell simulations.

In the criss-cross pathway (Figure 1.8) of the reaction mechanism, catalytic residue E166 removes the proton from C1 carbon of DHAP and moves to C2 carbon to donate the proton to the ketone oxygen<sup>23,27</sup>. In this pathway E166 has to shuttle between the two carbon atoms of the substrate. Alahuhta et al.<sup>16</sup> reported high mobility for the side chain of E166. In the TIM structure complexed with Phosphoglycolohydroxamate (PGH) which is a reaction intermediate analogue, E166 points away from the C2 carbon of the ligand (Figure 2.2) whereas in TIM complex with 2-(N-Formyl-N-hydroxy)-amino-ethylphosphonate (IPP) which mimics the natural substrate GAP, E166 points at C2 atom of the ligand<sup>16</sup>.

If the flip of G210 $\Psi$  from open to closed conformation induces the '*swung-in*' conformation, it would be an advantage to have the G210 $\Psi$  unperturbed by the presence of ligand in the active site. The change in this dihedral angle will aid in E166 movement between the two carbon atoms of the ligand.

Casteleijn et al.<sup>14</sup> also suggested that when G210 $\Psi$  adopts closed conformation, the steric clashes between P167 and carbonyl oxygen of G210 might trigger closure of loop-6. In our apo TIM simulations even when G210 $\Psi$  stays in closed conformation (Figure 3.18), loop-6 conformation is not influenced i.e. loop-6 transiently samples both open and closed conformations.

Sampson et al.<sup>42</sup> measured the barriers for the reaction mechanism in Y209F mutant and wild type. They reported that, there is an increase in the barrier height for the enolization by 5

kcal.mol<sup>-1</sup> to 6 kcal.mol<sup>-1</sup> in the Y209F mutant compared to the wild type. Based on the free energy profile Sampson et al.<sup>42</sup> postulated that closure of loop-6 is affected because of loss of hydrogen bond between Y209 and A176 of loop-6 which in turn affects the reactive conformation of E166 accounting for the rise in barrier for proton transfer.

Before the work of Casteleijn et al.<sup>14</sup> in 2006, it was speculated that closure of loop-6 induced conformational change in E166. But now it is clear that the change in dihedral angles of G210 $\Psi$  and G211 $\Phi$  regulate the conformational change in E166 and conformation of loop-6 does not affect E166. Berlow et al.<sup>39</sup> using NMR experiments in apo TIM has showed that Y209F mutation destabilizes the closed conformation by 0.84 kcal.mol<sup>-1</sup>, which does not account for the large change in the barrier of enolization reported by Sampson et al.<sup>42</sup>.

Based on these new results, an alternative hypothesis can be postulated where Y209F mutation though it affects the stability of the closed loop-6 conformation, might have a larger affect on the conformation of G210 $\Psi$  and G211 $\Phi$  dihedral angles in turn affecting the conformational sampling of E166. This hypothesis has to be tested by performing atomistic MD simulations of Y209F and comparing the populations of E166 in swung-in and swung-out conformation.

Our apo and holo TIM simulations have demonstrated that there is a pronounced influence of ligand and water molecules on the conformation of G211 $\Psi$  and S212 $\Phi$  dihedral angles. Electrostatic repulsion of, 4 kcal.mol<sup>-1</sup> between the negatively charged phosphate moiety of the ligand molecule and the carbonyl oxygen of G211 induces closed conformation for G211 $\Psi$  and S212 $\Phi$  dihedral angles. The change in conformation of these two dihedral angles provides around -8 kcal.mol<sup>-1</sup> of additional stabilization energy to the ligand molecule. As seen in chain-B of 1R2R crystal structure, in our apo open unit cell simulations we also found a water molecule which can stabilize the closed conformation of G211 $\Psi$  and S212 $\Phi$  for 175 ns to 270 ns.

Amyes et al.<sup>31</sup> reported that phosphate moiety of the substrate molecule induces closed conformation for loop-6 which we do not observe in our holo TIM simulations. Rather we suggest that phosphate moiety induces closed state for dihedral angles G211 $\Psi$  and S212 $\Phi$ . To summarize there is no effect of ligand molecules DHAP, EDT1, EDT2 2PG on conformation of dihedral angles G210 $\Psi$ , G211 $\Phi$ , S212 $\Psi$  and V212 $\Phi$  whereas closed conformation of G211 $\Psi$  and S212 $\Phi$  is ligand induced. As seen in 1R2R-B crystal structure, a water molecule can also induce closed conformation for G211 $\Psi$  and S212 $\Phi$ . The free energy required for

loop-7 conformational change between open and closed states, in presence of water and various ligand molecules has to be calculated in future.

To summarize loop-7 might play a dual role in TIM's catalysis. Dihedral angles G210 $\Psi$  and G211 $\Phi$  are not affected by ligand's presence in active site and might help shuttling of protons in the reaction mechanism by regulating the conformation of E166. G211 $\Psi$  and S212 $\Phi$  sample the closed conformation only in presence of a ligand molecule or a strategically placed water molecule. The closed conformation of G211 $\Psi$  and S212 $\Phi$  removes the electrostatic repulsion between the carbonyl oxygen of G211 and the phosphate moiety of the ligand molecules and enables the formation of a hydrogen bond between the amide group of S212 and phosphate moiety's oxygen, anchoring the ligand molecule in the active site.

## **4** Conclusions and Outlook

Triosephosphate Isomerase (TIM) catalyzes the reversible isomerization reaction where Glyceraldehyde 3-phosphate (GAP) is converted to Dihydroxyacetone phosphate (DHAP) and vice versa. The catalytic rate of the reaction  $(k_{cat})$  is  $10^4 \text{ s}^{-1}$  when the substrate is GAP and  $10^3 \text{ s}^{-1}$  when it is DHAP. TIM is a highly efficient enzyme with  $k_{cat}/k_m$  at  $10^9 \text{ M}^{-1}\text{ s}^{-1}$  and the reaction rate is only limited by diffusion of the substrate into the active site. To attain this catalytic efficiency, TIM has to streamline all the factors that contribute to the catalysis. These factors can be any conformational change involved in the catalysis, factors that affect the substrate binding and the product dissociation, as well as the barriers along the steps of the reaction mechanism.

In TIM structures, loop-6 and loop-7 envelope the active site and exist in open and closed conformations. Based on experimental studies it was proposed that the catalysis takes place when both loop-6 and loop-7 are in closed conformation. Loop-6 conformational change has been studied extensively studied by NMR experiments but not by microsecond MD simulations of entire TIM dimer. Dynamics of loop-7 conformational change is unexplored.

In this dissertation we mainly wanted to address the following questions. Is loop-6 and loop-7 conformation ligand gated or is it a natural motion of the protein? What factors affect the conformational changes in loop-6 and loop-7? What is the role of these conformational changes in the reaction catalyzed by TIM? How is the product formed at the end of the reaction released? Is there any influence of dynamics of loop-6 on loop-7 or vice versa? Below, we have summarized the new insights into the dynamics of these two loops (loop-6 and 7) and loop-5 and then we discuss the plausible sequence of events in TIM's catalysis.

## 4.1 Loop-6

In X-ray structures, in the absence of the ligand (apo) TIM loop-6 exists in open conformation and in closed conformation only when a ligand is present i.e. closed conformation of loop-6 is ligand induced. NMR studies have reported that loop-6 undergoes conformational change between open and closed conformations at same exchange rate ( $k_{ex}$ ) as  $k_{cat}$  of the enzyme which is  $10^4 \text{ s}^{-1}$ , when substrate analog G3P or 2PG are bound to the active site and  $k_{ex}$  in the apo enzyme is speculated to be  $10^5$  to  $10^6 \text{ s}^{-1}$ .

In section 3.4 of this thesis we have shown that loop-6 spontaneously undergoes transitions between open and closed conformations in apo TIM structures at  $k_{ex}$  of  $10^6$  s<sup>-1</sup>. As expected

from the X-ray data, no preference for closed loop-6 is observed in our simulations with DHAP/EDT1/EDT2/2PG bound to the active site. Either the closure of loop-6 in presence of these ligand molecules is a much slower process than the simulation length of each trajectory, or loop-6 as in apo TIM simulations does not close even in the when one of these ligand molecules bind to the active site. We also find that closure of N and C-termini is not affected by ligand being in reactive conformation since it is the tip of loop-6 (170-AIGTG-174) that interacts with ligand and tip of loop-6 opens when ligand is in encounter complex orientation.

There is an affect of solvent dynamics on loop-6 conformational change. In the apo crystal unit cell simulations with 1R2R X-ray structure and 2PG ligated crystal unit cell simulations with 1N55 X-ray structure, because of ten times slower diffusion constant of solvent, opening times of loop-6 starting from closed conformation are ten times longer compared to the normal MD simulations. Nonetheless even in crystal the unit cell loop-6 transiently samples open and closed conformations.

The free energy difference ( $\Delta$ G) between open and closed loop-6 conformations calculated from both the normal MD simulations (1.35 kcal.mol<sup>-1</sup>) and the umbrella sampling (US) simulation (1.15 kcal.mol<sup>-1</sup>) is close to the experimental estimate (1.22 kcal.mol<sup>-1</sup> to 1.77 kcal.mol<sup>-1</sup>). Since we do not have any transitions in DHAP/2PG ligated ensemble and very few transitions in EDT1 and EDT2 ligated TIM structures, the  $\Delta$ G calculated from ligated TIM simulations (3.14 kcal.mol<sup>-1</sup> to 4.89 kcal.mol<sup>-1</sup>) is not in agreement with the experimental data (-6.8 kcal.mol<sup>-1</sup>). The  $\Delta$ G calculated from US simulation with 2PG ligated TIM structure (-3.4 kcal.mol<sup>-1</sup>) is an improvement over normal 2PG ligated solvent MD simulations but still requires more simulation time.

Our simulations also strengthen the observation of Berlow et al.<sup>39</sup> and Wang et al.<sup>41</sup> that the population densities of loop-6 N-terminus and C-terminus in open and closed conformations are different. This suggests that loop-6 N and C-terminus do not move as a rigid body as proposed by earlier studies. The reason behind this is that, for N-terminus to open, loop-5 also has to adopt open conformation. Whereas C-terminus is solvent exposed and is not inhibited by any part of protein.

We find that, when loop-5 is restrained in closed conformation, loop-6 C-terminus samples open and closed conformations, whereas loop-6 N-terminus stays closed. When loop-5 is restrained in open conformation, loop-6 N and C-terminus sample both open and closed conformation. This confirms that loop-6 N and C-terminus are independent and that loop-5

conformation influences loop-6 N-terminus conformation. This uncoupled movement of Nterminus and C-terminus hinges of loop-6 and their influence on the reaction mechanism remains to be studied.

Effect of loop-5 conformation is also seen in loop-6 umbrella sampling (US) simulations. A restraining potential of 259 kcal.mol<sup>-1</sup> was used to keep loop-6 closed along the reaction coordinate and when loop-5 adopts open conformation, on which there was no restraining potential applied, N-terminus of loop-6 opens. This further emphasizes importance of loop-5 in regulating conformation of loop-6 N-terminus. In future studies, to calculate loop-6 free energy difference between open and closed conformations, a reaction coordinate which describes both loop-5 and loop-6 conformational change from open to closed conformations should be used.

#### 4.2 Loop-7

Loop-7 is a six residue fragment ('YGGSVN') which also undergoes a conformational change. The backbone phi ( $\Phi$ ), psi ( $\Psi$ ) dihedral angles of G210 $\Psi$ , G211 $\Phi$ , G211 $\Psi$ , S212 $\Phi$ , S212 $\Psi$  and V212 $\Phi$  differ between open and closed conformations. Just like for loop-6, based on X-ray structures it was proposed that loop-7 closed conformation is ligand induced. With one exception i.e. in chain-B of 1R2R crystal structure both loop-6 and loop-7 are closed in the absence of a ligand. A water molecule is present in the active site in the place of a one of the oxygens of the phosphate moiety. This water molecule can form a hydrogen bond with the amide group of S212 stabilizing the closed conformation of loop-7.

In our simulations we rather find that G210 $\Psi$ , G211 $\Phi$ , S212 $\Psi$  and V212 $\Phi$  sample open and closed conformations in both apo and holo TIM simulations. Whereas G211 $\Psi$  and S212 $\Phi$  prefer open conformation in the absence of the ligand and adopt closed conformation only when a ligand molecule is bound to the active site. As seen in the chain-B of 1R2R crystal structure, we also find that in apo TIM simulations a water can also trigger closed conformation for G211 $\Psi$  and S212 $\Phi$ . The role of S212 $\Psi$  and V212 $\Phi$  is not known and was not studied in this work.

We monitored the conformational change of E166 between swung-in and swung-out conformations and G210 $\Psi$  between open and closed states. Our results suggest that G210 $\Psi$  conformation regulates the conformation of E166 in agreement with the proposal of Casteleijn et al.<sup>14</sup>. We propose that a flexible G210 $\Psi$  and G211 $\Phi$ , which are not influenced

by ligand, would be an advantage for the catalytic mechanism for shuttling protons between the carbon atoms by catalytic residue E166. This hypothesis has to be studied by further computational studies.

Y209F mutant affects the catalytic rate and the existing hypothesis suggests that it is because of impairment of loop-6 closure on the active site. It might be possible that Y209F mutation might impair the conformational change of  $G210\Psi$  and thereby affect the E166 conformational change to the reactive conformation. This hypothesis has to be studied by simulations of Y209F TIM mutant.

The change in conformation of G211 $\Psi$  and S212 $\Phi$  leads to stabilization of the ligand in the active site. In the open conformation of G211 $\Psi$  and S212 $\Phi$ , carbonyl oxygen of G211 points towards the negatively charged (-2) phosphate moiety of the substrate and in the closed conformation the amide group of S212 forms a hydrogen bond with the phosphate moiety brings forth. We investigated the mechanism of this conformational change using MD simulations of apo TIM and holo TIM simulations.

In apo TIM simulations G211 $\Psi$  and S212 $\Phi$  prefer the open conformation and sample the closed conformation only when a water molecule is present within hydrogen bonding distance of S212 amide group. This water is similar to the water molecule seen in the chain-B of 1R2R crystal structure. In the absence of this water molecule G211 $\Psi$  and S212 $\Phi$  never sample the closed conformation in apo TIM simulations. Energetics of this process, where water molecule induces closed conformation for G211 $\Psi$  and S212 $\Phi$  is unclear and further studies are required.

We investigated the mechanism of  $G211\Psi$  and  $S212\Phi$  conformational change in presence of a ligand molecule. When  $G211\Psi$  and  $S212\Phi$  are in open conformation, the electrostatic repulsion between the carbonyl oxygen of G211 and the dianionic phosphate moiety of the ligand molecule induces a closed conformation for  $G211\Psi$  and  $S212\Phi$ . In closed state as stated earlier, amide group of S212 and oxygen of the phosphate moiety can form a hydrogen bond and provide additional stabilization interactions to the substrate in the active site.

Amyes et al.<sup>31</sup> suggested that some part of the phosphate binding free energy is used to induce closed conformation for loop-6. Based on our MD simulations we would rather suggest that it is rather utilized in inducing closed conformation for G211 $\Psi$  and S212 $\Phi$  and not for loop-6 because even when ligand molecules (DHAP/EDT1/EDT2/2PG) are bound to

the active site we do not observe closed conformation for loop-6. From our simulations it is quite evident that closed conformation of  $G211\Psi$  and  $S212\Phi$  is induced by either by a ligand molecule or a strategically placed water molecule.

#### **4.3** Plausible sequence of events

Base on previous studies, the existing view of TIM's catalysis and sequence of events that enable the reaction are following. Binding of either of the substrates DHAP/GAP induces closure of loop-7. G210 $\Psi$  and G211 $\Phi$  flip to closed state, triggers swung-in conformation for E166 and closed conformation for P167, which in turn closes loop-6. After the catalysis, loop-7 flips to open conformation followed by opening of loop-6 and release of the product.

From our results we propose the following scenario:

Binding of ligand will induce a closed conformation for  $G211\Psi$  and  $S212\Phi$ , which will an additional hydrogen bond to the substrate. Since the conformational change of  $G210\Psi$  is not influenced by binding of substrate, transitions between open and closed conformations will induce a conformational change in E166. This might help shuttling of protons by E166 between the two carbon atoms of the substrate.

Loop-6 transiently samples open and closed conformations. And when the substrate is bound to the active site, after closure of loop-6, catalysis will take place. If the closed conformation of loop-6 is not ligand induced, no special trigger is required to initiate opening of loop-6 and opening of loop-6 will provide an opportunity for release of the product.

Loop-6 has to stay closed long enough for catalysis to take place without the formation of methylglyoxal which is formed by solvent exposure of the reaction intermediates. It was suggested that the cellular environment where TIM performs this catalysis, mimics the crystal environment because of crowding in the cellular cytoplasm<sup>153-155</sup>. In our crystal simulations, starting from an open conformation, loop-6 undergoes a spontaneous conformational change to closed state, stays closed for greater than 100 ns and samples the open conformation again. Proton transfer occurs in sub picosecond time scales and since loop-6 stays closed for nanoseconds, this time duration should be sufficient for the catalytic process.

The catalytic rate of TIM is  $10^4 \text{ s}^{-1}$  in GAP to DHAP direction and  $10^3 \text{ s}^{-1}$  in GAP to DHAP direction, i.e. to catalyze one molecule of GAP to DHAP time required is 100 µs and for one molecule of DHAP to GAP it is 1 ms. We observe conformational changes in loop-6 and

loop-7 dihedral angles G210 $\Psi$ , G211 $\Phi$ , G211 $\Psi$  and S212 $\Phi$  in microsecond regime. As long as the conformational changes that affect the catalysis are faster than or equal to the diffusion rate of the substrate, than the catalytic rate of the reaction is still limited by the diffusion of the substrate into the active site or product out of the active site. To have a complete understanding of the functioning of TIM, further computational and experimental studies which can probe the affect of each individual conformational change and the affect of it on the barriers of the reaction mechanism are required.

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# 7 Appendix

# 7.1 Ligand Parameters

GAFF<sup>96</sup> forcefield parameters for natural substrate DHAP, reaction intermediates EDT1 and EDT2 and inhibitor 2PG used in TIM simulations

## 7.1.1 Atom names, atom types and partial charges

Ligand molecule	Atom name	Atom type	Atomic charge
DDP	C1	C3D	0.24790
	H11	H1D	0.00400
	H12	H1D	0.00400
	01	O1D	0.71750
	ОН	H2D	0.47580
	C2	CDD	0.49960
	O2	O2D	-0.56360
	C3	C3D	0.23470
	H21	H1D	0.00460
	H22	H1D	0.00460
	O3	OSD	-0.57620
	P1	P1D	1.28070
	O4	O2D	-0.96620
	O5	O2D	-0.96620
	06	O2D	-0.96620
EDT1	C1	C2O	0.18480
	C2	C30	-0.01260
	H1	H1O	0.03820
	H2	H1O	0.03820
	01	OSO	-0.48890
	P1	P50	1.24520
	O2	00	-0.96690
	O3	00	-0.96690
	O4	00	-0.96690
	C3	C2O	-0.35500
	Н3	H4O	0.16250
	O5	00	-0.73360
	O6	ОНО	-0.58920
	H4	НОО	0.41110

Table 7.1: Atom names, atomtypes and partial charges on each atom for vaiorus ligand molecules.

EDT2	01	OE	-0.74580
	C1	C2E	-0.23110
	C2	C3E	0.18850
	H1	H1E	0.01160
	H2	H1E	0.01160
	O2	OSE	-0.51460
	P1	P5E	1.22330
	03	OE	-0.96040
	04	OE	-0.96040
	05	OE	-0.96040
	C3	C2E	0.10500
	H3	H4E	0.00610
	O6	OHE	-0.57680
	H4	HOE	0.40340
2PG	C1	C11	0.14400
	H1	H11	0.01020
	H2	H11	0.01020
	01	OS1	-0.58280
	P1	P11	1.42970
	02	011	-1.02600
	03	011	-1.02600
	04	011	-1.02600
	C2	C12	1.01290
	05	011	-0.97310
	06	011	-0.97310

#### 7.1.2 Atom types and Lennard-Jones parameters

Atom type	Mass (amu)	LJ σ (nm)	LJ ε (kJ mol <sup>-1</sup> )
C3D	12.010	0.3399670	0.4577300
H1D	1.008	0.2471350	0.0656888
P1D	30.970	0.3741770	0.8368000
H2D	1.008	0.0000000	0.0000000
OSD	16.000	0.3000010	0.7112800
CDD	12.010	0.3399670	0.3598240
O1D	16.000	0.3066470	0.8803140
O2D	16.000	0.2959920	0.8786400
H1E	1.008	0.2471350	0.0656888
C3E	12.010	0.3399670	0.4577300
P5E	30.970	0.3741770	0.8368000
C2E	12.010	0.3399670	0.3598240
H4E	1.008	0.2510550	0.0627600
HOE	1.008	0.0000000	0.0000000
OSE	16.000	0.3000010	0.7112800
OHE	16.000	0.3066470	0.8803140
OE	16.000	0.2959920	0.8786400
C3O	12.010	0.3399670	0.4577300
H1O	1.008	0.2471350	0.0656888
P5O	30.970	0.3741770	0.8368000
C2O	12.010	0.3399670	0.3598240
H4O	1.008	0.2510550	0.0627600
OSO	16.000	0.3000010	0.7112800
HOO	1.008	0.0000000	0.0000000
OHO	16.000	0.3066470	0.8803140
00	16.000	0.2959920	0.8786400
C12	12.010	0.3399670	0.3598240
H11	1.008	0.2471350	0.0656888
P11	30.970	0.3741770	0.8368000
011	16.000	0.2959920	0.8786400
OS1	16.000	0.3000010	0.7112800

Table 7.2: Lennard-Jones (LJ) parameters for various atom types. Epsilon ( $\epsilon$ ) is the potential well depth and sigma ( $\sigma$ ) is the distance at which the inter-particle potential is zero.

#### 7.1.3 Bond Parameters

Atom type <i>i</i>	Atom type j	Bond length (nm)	Force constant (kJ.mol <sup>-1</sup> ·nm <sup>-2</sup> )
C3D	H1D	0.10930	281080.0
O1D	H2D	0.09740	309280.0
C3D	O1D	0.14260	262840.0
C3D	CDD	0.15080	274720.0
CDD	O2D	0.12140	542250.0
C3D	OSD	0.14390	252300.0
OSD	P1D	0.16020	286600.0
P1D	O2D	0.14810	408110.0
C3O	H1O	0.10930	281080.0
C2O	H4O	0.10840	291710.0
ОНО	HOO	0.09740	309280.0
C2O	C30	0.15080	274720.0
C2O	C2O	0.13240	493460.0
C2O	00	0.12610	457060.0
C3O	OSO	0.14390	252300.0
OSO	P5O	0.16020	286600.0
P5O	00	0.14810	408110.0
C2O	ОНО	0.13330	355970.0
C3E	H1E	0.10930	281080.0
C2E	H4E	0.10840	291710.0
OHE	HOE	0.09740	309280.0
OE	C2E	0.12610	457060.0
C2E	C3E	0.15080	274720.0
C2E	C2E	0.13240	493460.0
C2E	OHE	0.13330	355970.0
C3E	OSE	0.14390	252300.0
OSE	P5E	0.16020	286600.0
P5E	OE	0.14810	408110.0
C11	H11	0.10930	281080.0
C11	OS1	0.14390	252300.0
C11	C12	0.15080	274720.0
OS1	P11	0.16020	286600.0

Table 7.3: Bond parameters between atom types i and j.

P11	011	0.14810	408110.0
C12	011	0.12140	542250.0

## 7.1.4 Angle parameters

Atom type <i>i</i>	Atom type j	Atom type k	Angle (°)	Force constant (kJ.mol <sup>-1</sup> .rad <sup>-2</sup> )
H11	C11	H11	109.55	328.03
H11	C11	OS1	108.82	425.09
H11	C11	C12	107.66	398.32
C11	OS1	P11	118.00	656.89
C11	C12	011	123.11	569.02
OS1	C11	C12	109.82	569.02
OS1	P11	011	116.09	368.19
011	P11	011	115.80	384.93
011	C12	011	127.33	661.91
C3D	O1D	H2D	108.16	394.13
H1D	C3D	H1D	109.55	328.03
H1D	C3D	01D	109.88	426.77
H1D	C3D	CDD	107.66	398.32
CDD	C3D	H1D	107.66	398.32
H1D	C3D	OSD	108.82	425.09
C3D	CDD	O2D	123.11	569.02
C3D	CDD	C3D	116.05	525.51
O1D	C3D	CDD	109.46	572.37
CDD	C3D	OSD	109.82	569.02
O2D	CDD	C3D	123.11	569.02
C3D	OSD	P1D	118.00	656.89
OSD	P1D	O2D	116.09	368.19
O2D	P1D	O2D	115.80	384.93
C2O	C30	H1O	110.46	393.30
C2O	C2O	H4O	124.68	412.54
H1O	C30	H1O	109.55	328.03
H1O	C30	OSO	108.82	425.09
C2O	ОНО	НОО	108.98	417.56
H4O	C2O	ОНО	116.22	437.65
C2O	C30	OSO	108.88	571.53
C2O	C2O	ОНО	122.28	599.15
C3O	C2O	C20	123.42	538.06

Table 7.4: Angle parameters between atom types i, j and k.

C3O	C20	00	116.88	578.23
C30	OSO	P5O	118.00	656.89
OSO	P5O	00	116.09	368.19
00	P5O	00	115.80	384.93
C2O	C2O	00	130.89	594.13
OE	C2E	H4E	120.31	426.29
C2E	C3E	H1E	110.46	393.30
C2E	C2E	H4E	124.68	412.54
C2E	OHE	HOE	108.98	417.56
H1E	C3E	H1E	109.55	328.03
H1E	C3E	OSE	108.82	425.09
OE	C2E	C2E	130.89	594.13
C2E	C3E	OSE	108.88	571.53
C3E	C2E	C2E	123.42	538.06
C3E	C2E	OHE	118.81	564.00
C3E	OSE	P5E	118.00	656.89
OSE	P5E	OE	116.09	368.19
OE	P5E	OE	115.80	384.93
C2E	C2E	OHE	122.28	599.15

## 7.1.5 Dihedral angles parameters

Table 7.5: Dihedral parameters using improper and proper dihedrals function between atom types i, j, k and l. F<sub>c</sub> is force constant. Improper dihedrals are used to keep planar groups planar. Proper dihedrals are used to define standard dihedral angles.

Atom type	Atom type	Atom type	Atom type	Dihedral	Phase	F <sub>c</sub> (kJ.mol <sup>-</sup>	Multiplici
i	j	k	l	type	(°)	1)	ty
OH3	C33	C33	OH3	improper	300.00	100.00000	2
OH3	C33	C33	OS3	improper	180.00	100.00000	2
C11	011	C12	011	improper	180.00	43.93200	2
LP	LP	ON	NN	improper	180.00	85.57336	2
O1D	C3D	CDD	O2D	improper	111.33	3.07483	2
OS1	C11	C12	011	proper	180.00	4.30500	2
C12	C11	OS1	P11	proper	0.00	1.60387	3
C11	OS1	P11	011	proper	0.00	1.04600	3

Atom	Atom	Atom	Atom						
type i	type j	type k	type <i>l</i>	C <sub>0</sub>	<b>C</b> <sub>1</sub>	<b>C</b> <sub>2</sub>	<b>C</b> <sub>3</sub>	C <sub>4</sub>	C <sub>5</sub>
				0.0000	0.000	0.0000	0.000	0.000	0.000
C3D	CDD	C3D	H1D	0	00	0	00	00	00
							-		
				0.6987	2.096	0.0000	2.794	0.000	0.000
H1D	C3D	O1D	H2D	3	18	0	91	00	00
				0.0000	0.000	0.0000	0.000	0.000	0.000
H1D	C3D	CDD	O2D	0	00	0	00	00	00
				0.0000	0.000	0.0000	0.000	0.000	0.000
H1D	C3D	CDD	C3D	0	00	0	00	00	00
				0.0000	0.000	0.0000	0.000	0.000	0.000
CDD	C3D	O1D	H2D	0	00	0	00	00	00
				0.0000	0.000	0.0000	0.000	0.000	0.000
O2D	CDD	C3D	H1D	0	00	0	00	00	00
							-		
111D	COD	ano	DID	1.6024	4.807	0.0000	6.409	0.000	0.000
HID	C3D	OSD	PID	7	42	0	89	00	00
COD	CDD	COD	0.00	0.0000	0.000	0.0000	0.000	0.000	0.000
C3D	CDD	C3D	OSD	0	00	0	00	00	00
				0.0000	-	-	-	-	7.002
01D	C2D	CDD	010	0.0000	0.023	9.7308	4.022	0.959	/.003
UID	C3D	CDD	02D	0	30	0 0000	43	4/	43
01D	C2D	CDD	C2D	0.0000	0.000	0.0000	0.000	0.000	0.000
	C3D	CDD	CSD	0,0000	0.000	0,0000	0.000	0.000	0.000
CDD	C3D	OSD	DID	0.0000	0.000	0.0000	0.000	0.000	0.000
CDD	CJD	USD		0	00	0	00	00	00
				37 127	8 662	21 609	- 1 183	2 1 1 3	0.968
02D	CDD	C3D	OSD	00	65	90	59	2.113 54	43
020	CDD	0.50	ODD	0.0000	0.000	6 6944	0.000	0,000	0.000
C3D	OSD	P1D	02D	0.0000	00	0.0911	00	00	00
002	0.52	112	010	0.0000	0.000	0.0000	0.000	0.000	0.000
C3D	C3D	CDD	O2D	0	00	0	00	00	00
			-	_		_			
				8.7864	0.000	8.7864	0.000	0.000	0.000
C2O	C2O	OHO	HOO	0	00	0	00	00	00
						-			
				55.647	0.000	55.647	0.000	0.000	0.000
C3O	C2O	C2O	H4O	20	00	20	00	00	00
				0.0000	0.000	0.0000	0.000	0.000	0.000
C2O	C2O	C3O	H10	0	00	0	00	00	00
				0.0000	0.000	0.0000	0.000	0.000	0.000
00	C2O	C3O	H1O	0	00	0	00	00	00
							-		
				1.6024	4.807	0.0000	6.409	0.000	0.000
H10	C30	OSO	P50	7	42	0	89	00	00
					0.000	-	0.000	0.000	0.000
0.0	000	000		55.647	0.000	55.647	0.000	0.000	0.000
00	C20	C20	H4O	20	00	20	00	00	00
H4O	C2O	OHO	HOO	8.7864	0.000	-	0.000	0.000	0.000

Table 7.6: Dihedral parameters defined using Ryckaert-Bellemans function between atom types *i*, *j*, *k* and *l*. Force constants  $C_0$ ,  $C_1$ ,  $C_2$ ,  $C_3$ ,  $C_4$  and  $C_5$  are in kJ.mol<sup>-1</sup>.

				0	00	8.7864	00	00	00
						-			
				9.2048	0.000	9.2048	0.000	0.000	0.000
C2O	H4O	C2O	OHO	0	00	0	00	00	00
				1.6024	4.807	0.0000	- 6.409	0.000	0.000
C2O	C30	OSO	P5O	7	42	0	89	00	00
				55 647	0.000	-	0.000	0.000	0.000
C30	C20	C20	оно	20	0.000	55.647 20	0.000	0.000	0.000
	020	020		0.0000	0.000	6.6944	0.000	0.000	0.000
C30	OSO	P50	00	0	00	0	00	00	00
C20	C20	C30	050	0.0000	0.000	0.0000	0.000	0.000	0.000
0.20	0.20	0.50	030	0	-	-	-	00	00
				37.127	8.662	21.609	1.183	2.113	0.968
00	C20	C30	OSO	00	65	90	59	54	43
				55.647	0.000	- 55.647	0.000	0.000	0.000
00	C2O	C2O	OHO	20	00	20	00	00	00
00	C20	C20	C20	0.0000	0.000	0.0000	0.000	0.000	0.000
00	0.20	0.50	0.20	0	00	-	00	00	00
				55.647	0.000	55.647	0.000	0.000	0.000
C3E	C2E	C2E	H4E	20	00	20	00	00	00
				8 7864	0.000	- 8 7864	0.000	0.000	0.000
C3E	C2E	OHE	HOE	0.7004	00	0.7004	00	00	00
	~~~	~~~	~~~	0.0000	0.000	0.0000	0.000	0.000	0.000
H1E	C3E	C2E	C2E	0	00	0	00	00	00
H1E	C3E	C2E	OHE	0.0000	0.000	0.0000	0.000	0.000	0.000
							-		
	C2E	OSE	DSE	1.6024	4.807	0.0000	6.409	0.000	0.000
	CJE	USE	PJE	/	42	-	89	00	00
				8.7864	0.000	8.7864	0.000	0.000	0.000
C2E	C2E	OHE	HOE	0	00	0	00	00	00
				55.647	0.000	- 55.647	0.000	0.000	0.000
H4E	C2E	C2E	OHE	20	00	20	00	00	00
				0.0040	0.000	-	0.000	0.000	0.000
OE	C2E	H4E	C2E	9.2048	0.000	9.2048	0.000	0.000	0.000
	C2L	IIIL		0	00	-	00	00	00
<b>0</b> E	COL			55.647	0.000	55.647	0.000	0.000	0.000
OE	C2E	C2E	C3E	20	00	- 20	00	00	00
				55.647	0.000	55.647	0.000	0.000	0.000
OE	C2E	C2E	OHE	20	00	20	00	00	00
				1 6024	4 807	0.0000	-	0.000	0.000
C2E	C3E	OSE	P5E	7	42	0.0000	89	0.000	00

				0.0000	0.000	6.6944	0.000	0.000	0.000
C3E	OSE	P5E	OE	0	00	0	00	00	00
				0.0000	0.000	0.0000	0.000	0.000	0.000
OSE	C3E	C2E	C2E	0	00	0	00	00	00
					-	-	-		
				37.127	8.662	21.609	1.183	2.113	0.968
OSE	C3E	C2E	OHE	00	65	90	59	54	43
				0.0000	0.000	0.0000	0.000	0.000	0.000
C2E	C3E	C2E	OHE	0	00	0	00	00	00
							-		
				1.6024	4.807	0.0000	6.409	0.000	0.000
H11	C11	OS1	P11	7	42	0	89	00	00
					-				
				3.6819	4.351	0.0000	1.338	0.000	0.000
H11	C11	C12	O11	2	36	0	88	00	00
				0.0000	0.000	6.6944	0.000	0.000	0.000
C11	OS1	P11	011	0	00	0	00	00	00
				0.0000	0.000	0.0000	0.000	0.000	0.000
OS1	C11	C12	011	0	00	0	00	00	00

#### 7.2 Loop-6 RMSD plots

RMSD of loop-6 from a normal MD CTIM simulation and a CTIM crystal simulation, with respect to open loop-6 from chain-A of X-ray structure 5TIM and closed loop-6 from chain-B of X-ray structure is shown in Figure 7.1 below. RMSD of open loop-6 with respect to closed loop-6 is 0.39 nm. In Figure 7.1 we can see that when both N-terminus (blue) and C-terminus (red) are closed (X-ray closed distance dotted green line) then loop-6 RMSD is close to the X-ray closed structure and around 0.39 nm with respect to X-ray open loop-6 and vice versa (Figure 7.1). In crystal simulation both N-terminus and C-terminus hinge are closed hence RMSD with respect to closed loop-6 (cyan line) is small and RMSD with respect to open loop-6 is greater than or equal to 0.39 nm as it is seen in X-ray structures.



Figure 7.1: Dynamics of N-terminus (W169-S212; blue) and C-terminus (V176-S212; red) hinges of loop-6. In the third panel loop-6 RMSD with respect to closed loop-6 from chain-B of 6TIM X-ray structure (orange) and open loop-6 from chain-A of X-ray structure 5TIM (cyan) are shown. RMSD of open loop-6 versus closed loop-6 is shown in dotted black line. Dotted blue line is the N-terminus/C-terminus distance in the X-ray closed structure and dotter green line in X-ray open structure.

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