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# Probing the structure of human tRNA<sub>3</sub><sup>Lys</sup> in the presence of ligands using docking, MD simulations and MSM analysis†

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The tRNA<sub>3</sub><sup>Lys</sup>, which acts as a primer for human immunodeficiency virus type 1 (HIV-1) reverse transcription, undergoes structural changes required for the formation of a primer–template complex. Small molecules have been targeted against tRNA<sub>3</sub><sup>Lys</sup> to inhibit the primer–template complex formation. The present study aims to understand the kinetics of the conformational landscape spanned by tRNA<sub>3</sub><sup>Lys</sup> in apo form using molecular dynamics simulations and Markov state modeling. The study is taken further to investigate the effect of small molecules like 1,4T and 1,5T on structural conformations and kinetics of tRNA<sub>3</sub><sup>Lys</sup>, and comparative analysis is presented. Markov state modeling of tRNA<sub>3</sub><sup>Lys</sup> apo resulted in three metastable states where the conformations have shown the non-canonical structures of the anticodon loop. Based on analyses of ligand–tRNA<sub>3</sub><sup>Lys</sup> interactions, crucial ion and water mediated H-bonds and free energy calculations, it was observed that the 1,4-triazole more strongly binds to the tRNA<sub>3</sub><sup>Lys</sup> compared to 1,5-triazole. However, the MSM analysis suggest that the 1,5-triazole binding to tRNA<sub>3</sub><sup>Lys</sup> has brought rigidity not only in the binding pocket (TΨC arm, D–TΨC loop) but also in the whole structure of tRNA<sub>3</sub><sup>Lys</sup>. This may affect the easy opening of primer tRNA<sub>3</sub><sup>Lys</sup> required for HIV-1 reverse transcription.

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## Introduction

The human immunodeficiency virus type 1 (HIV-1) requires human tRNA<sub>3</sub><sup>Lys</sup> for its reverse transcription process. The 3' terminal 18 nucleotides of tRNA<sub>3</sub><sup>Lys</sup> bind to the Primer Binding Site (PBS) of the viral genome (vRNA) in a complementary manner and form a reverse transcription primer–template complex with the help of a chaperone protein called nucleocapsid (NC).<sup>1–10</sup> The formation of the vRNA/tRNA primer–template complex requires the unfolding of the 3D structure of the primer tRNA<sub>3</sub><sup>Lys</sup> and opening of base-pairing in the acceptor and TΨC arms. Annealing between the tRNA and vRNA of the reverse transcription initiation complex is proposed to be achieved by the mature NC.<sup>11–15</sup> NMR studies showed that the NC protein may not be required for the formation of the primer–template complex; instead, it may help in accelerating the process. The unwinding process could be initiated by the unpaired 3' CCA end of the tRNA<sub>3</sub><sup>Lys</sup> and the unpaired bases from the acceptor and TΨC stem junction.<sup>16–20</sup>

As per the Isel *et al.*, the entire reverse transcription initiation of HIV-1 using tRNA<sub>3</sub><sup>Lys</sup> as primer takes place *via* multi step

process. Initially the tRNA<sub>3</sub><sup>Lys</sup> gets opened by breaking of U66–U67 interaction by the HIV-1 NC.<sup>17,21</sup> Prior to the entire acceptor stem opening, the CCA 3' end of tRNA<sub>3</sub><sup>Lys</sup> gets annealed to the PBS of viral RNA. Next, antiPAS/PAS interactions initiates the opening of TΨC-arm as well as D-arm.<sup>22</sup> These findings were supported by Tisne *et al.* with their in tube heat-annealing experiment of tRNA<sub>3</sub><sup>Lys</sup> and viral RNA. They have observed two weak peaks corresponding to the U66 and U67 imino protons at 15 °C. They have also reported about the PBS and anti-PBS hybridization at 25 °C. Apart from this, they have found that one crucial tertiary interaction between T54 and A58 in TΨC loop remains stable even at 60 °C.<sup>17,21,22</sup>

The zinc fingers of NC are also involved in destabilization of the tertiary interactions between the D loop and TΨC loop.<sup>16,21</sup> The reverse transcriptase (RT) enzyme of HIV-1 binds to the primer–template complex and forms the actual reverse transcription initiation complex. The anticodon of tRNA<sub>3</sub><sup>Lys</sup> and A-loop of viral RNA interaction are also important to build the correct functional complex.<sup>1–7</sup> The modified bases present at the anticodon region are critical for maintaining the stability of the complex system.<sup>23–25</sup> The studies regarding the requirement of tRNA<sub>3</sub><sup>Lys</sup> for HIV-1 reverse transcription initiation and the detailed process have been thoroughly discussed in a review paper.<sup>22</sup>

A number of experiments and computational studies have been carried out to target tRNA<sub>3</sub><sup>Lys</sup> to inhibit the HIV-1 reverse transcription either by small molecules, or by antisense

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oligomers.<sup>22,26–29</sup> Several molecular dynamics simulations studies of tRNA molecules have been carried out to understand the effect of modified bases, folding, dynamics, binding of ligands, allostery *etc.*<sup>30–35</sup>

One of the established strategies in drug discovery in recent years is the fragment-based approach.<sup>36–38</sup> The goal of this method is to build drug leads in pieces, through the identification of moderately binding fragments that are either expanded or linked together. Experimental studies have been reported to identify the small molecules targeting tRNA<sub>3</sub><sup>Lys</sup>, the HIV-1 reverse transcription primer, in a fragment-based approach. The small molecules targeting the primer tRNA<sub>3</sub><sup>Lys</sup> to destabilize tRNA/vRNA complex have been investigated in a therapeutic angle.<sup>39</sup> Binding of a library of compounds to the different tRNAs like tRNA<sub>3</sub><sup>Lys</sup>, tRNA<sup>met</sup>, tRNA<sup>phe</sup>, was monitored in a TROSY

experiment which showed improved affinity and specificity for the D-arm region for tRNA<sub>3</sub><sup>Lys</sup>.<sup>39</sup> Many studies have been focused on the peptides targeted to the D arm of tRNA<sub>3</sub><sup>Lys</sup> and their effect on initiation of reverse transcription.<sup>39–41</sup> It has been reported that diaminocyclopentanol (DACP) and kynuramine are millimolar binders of the target tRNA<sub>3</sub><sup>Lys</sup>.<sup>39,40</sup> These two fragments were evolved and connected *via* a 1,2,3-triazole moiety leading to a second generation of compounds. The ligation of two molecules *via* a 1,2,3-triazole moiety is a very popular strategy in chemical biology and has been extensively used.<sup>42–44</sup> In order to study in depth, the influence of the linker on the binding of DACP and kynuramine compounds, 1,4-triazole (1,4T) moiety is changed as its isomer 1,5-triazole (1,5T). This modification is expected to change the orientation between the two fragments without modifying the chemical nature or size of the molecule.

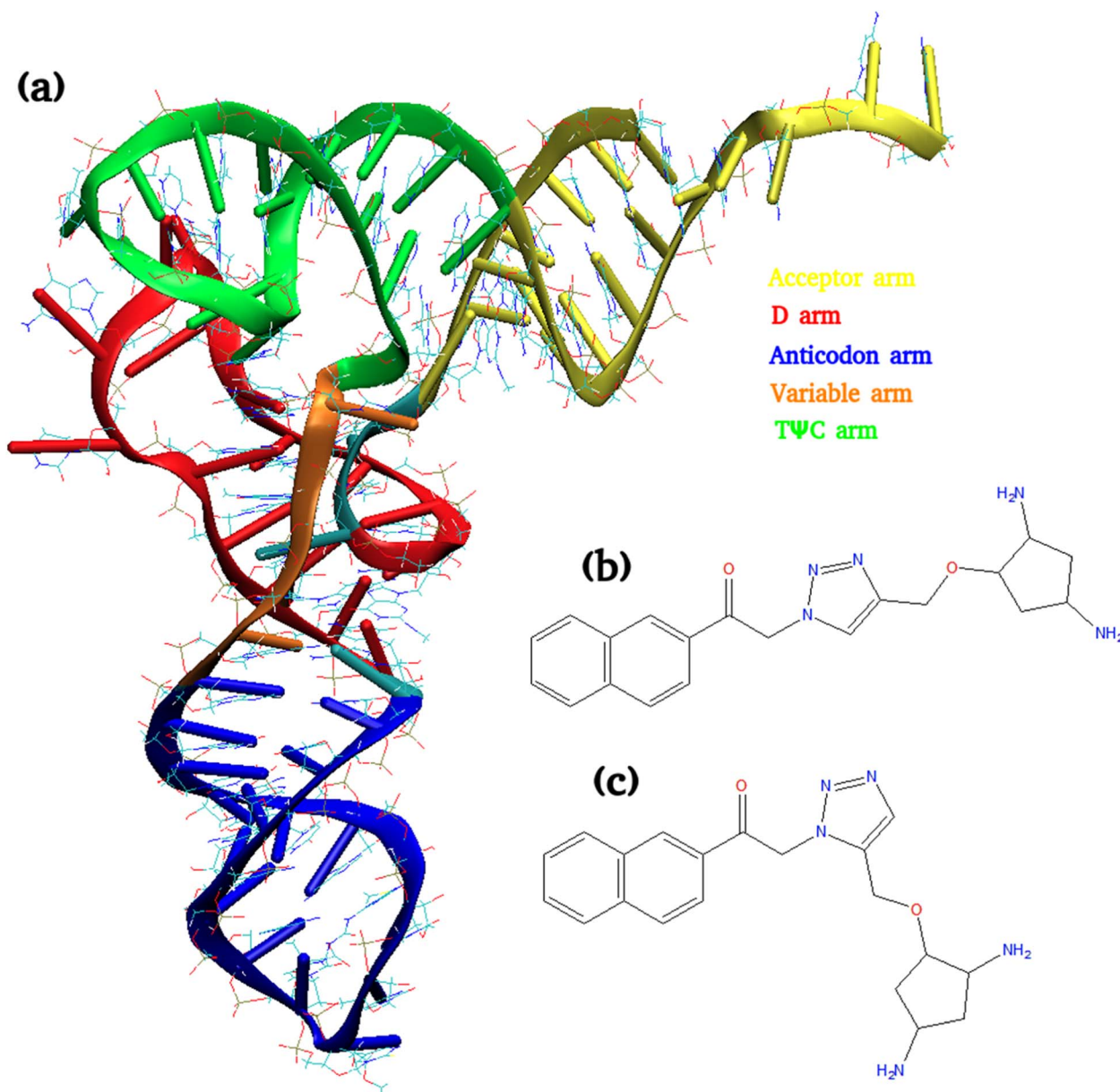


Fig. 1 The structures of (a) tRNA<sub>3</sub><sup>Lys</sup> (b) 1,4T ligand (c) 1,5T ligand.



The structures of these two ligands are shown in Fig. 1. The NMR and experimental studies have been earlier reported to understand the differences in the binding affinities of 1,4T and 1,5T towards tRNA<sub>3</sub><sup>Lys</sup> and they show how the orientation of the two fragments can have a substantial impact on the location of the binding sites on the RNA target. The structural insight into the binding of ligands on the target RNA at atomic level using computational methods may help in understanding and designing novel molecules. The detailed information about the location of binding sites and the interactions of these ligands with the residues of target RNA may throw more light in designing fragment based novel ligands against tRNA<sub>3</sub><sup>Lys</sup>.<sup>45</sup>

Here, an attempt has been made to explore the conformational ensemble of tRNA<sub>3</sub><sup>Lys</sup> in the apo form as well as in the presence of ligands. The two ligands (1,4T and 1,5T) docked with tRNA<sub>3</sub><sup>Lys</sup> have been used as a probe to gain structural insight of the tRNA<sub>3</sub><sup>Lys</sup> using molecular simulations. Multiple simulations for ligand free and ligand bound systems of tRNA<sub>3</sub><sup>Lys</sup> have been carried out to explore conformational space of tRNA<sub>3</sub><sup>Lys</sup>. The trajectory analysis has been done for RMSD, RMSF, H-bond interactions, MM-GBSA free energy, tRNA–ligand interactions. It is realized that to analyse the drug molecules, only binding affinities are not important but kinetics is also a crucial factor in determining the drug efficacy.<sup>46,47</sup> Therefore, Markov State Modeling (MSM) was performed to probe the kinetics of apo tRNA<sub>3</sub><sup>Lys</sup> and ligand bound tRNA<sub>3</sub><sup>Lys</sup> systems. Markov state model is a statistical model which can be built on an ensemble of short trajectories sampled from different regions of the free energy landscape of the system.<sup>48–51</sup> MSM networks describe long-timescale dynamics and equilibrium properties. Also, MSM analysis provides deeper insights to the conformational ensemble of biomolecules at atomic resolution.<sup>52,53</sup>

The kinetically connected metastable states were obtained using Markov State modeling of MD simulation trajectories of tRNA<sub>3</sub><sup>Lys</sup> systems. This may help to understand conformational rearrangement undergone by tRNA<sub>3</sub><sup>Lys</sup> to adopt the ligand in the D–TΨC loop junction. Later, a comparative analysis of all the three systems apo, 1,4T and 1,5T ligand bound tRNA<sub>3</sub><sup>Lys</sup> have been presented here.

## Methods

### Start structures and parameters

The X-ray crystal structure of the tRNA<sub>3</sub><sup>Lys</sup> solved at 3.3 Å [PDB id 1FIR]<sup>54</sup> which is available with the PDB database was used as the starting structure. The ligands 1,4-triazole and 1,5-triazole were built as reported in literature<sup>45</sup> by using the molecular visualization and building tool Gaussview.<sup>55</sup> The parameters for the nonstandard/modified nucleotide bases were calculated for earlier work<sup>26</sup> based on the following methodology which was used for ligand molecules as well. The parameters for the two ligand molecules and modified bases present in tRNA<sub>3</sub><sup>Lys</sup> were derived from AMBER force field ff12SB. Partial atomic charges were calculated by performing single point energy calculations at Hartree–Fock level using 6-311G\* basis set to obtain

electrostatic potential charges. Electronic structure calculations were performed with the Gaussian03 package.<sup>56</sup>

### Molecular docking

The crystal structure of the tRNA<sub>3</sub><sup>Lys</sup> is considered as a receptor and the 1,4T and 1,5T molecules are considered as ligands in the docking methodology. Since, considering multiple conformations for the target structure is computationally expensive, the target tRNA<sub>3</sub><sup>Lys</sup> is considered as a single static structure for docking and multiple conformations are considered for both the ligands. The receptor file was used to select spheres in the receptor in the process of receptor preparation by the DOCK6 program.<sup>57</sup> Hydrogen atoms were removed and the tRNA<sub>3</sub><sup>Lys</sup> was prepared using UCSF Chimera.<sup>58</sup> The solvent-accessible surface of the tRNA<sub>3</sub><sup>Lys</sup> binding site was calculated using a probe radius of 1.4 Å by using the DMS program in the DOCK6 software package. Receptor spheres were generated using the program SPHGEN. Spheres covering the hotspot were selected within 10 Å from the positions of the heavy atoms of the critical residues in the D–TΨC interacting region taken from literature.<sup>45</sup> The grid box that enclosing the selected spheres was generated with an extra 5 Å added in each dimension. Ligand flexibility was employed during the docking process using the DOCK6 module with output presented as grid scores. Docking of both the ligands against different tRNA crystal structures and MSM representative structures of each macrostate, apart from tRNA<sub>3</sub><sup>Lys</sup> was also carried out.

### Molecular dynamics simulations of docked complexes

The molecular dynamics (MD) simulations were carried out using AMBER12 (ref. 59) program with ff12SB force field.<sup>60</sup> The highest grid score (most negative) docking conformations generated by DOCK6 were taken as initial conformations for MD simulations. The two systems (tRNA<sub>3</sub><sup>Lys</sup>–1,4T and tRNA<sub>3</sub><sup>Lys</sup>–1,5T) with their respective ligands having high grid score and one apo system (tRNA<sub>3</sub><sup>Lys</sup> apo) considered as starting structures for MD simulations. The topology parameters of all systems were created by using the XLEAP module of AmberTools. All the systems were neutralized by adding Mg<sup>2+</sup> ions and solvated with a TIP3P<sup>61</sup> water box. Energy minimization was carried out for all the systems by using the steepest descent method for 5000 steps. Then temperature ramping was carried out at a temperature range of 50 K to 300 K for 100 ps, by using constraints of 100 kcal mol<sup>−1</sup>. Equilibration was done by gradually reducing the constraints for 500 ps. The equilibration protocol of Cheatham was followed.<sup>62</sup> Simulations were performed under periodic boundary conditions by employing the Particle Mesh Ewald<sup>63</sup> technique to account for long range electrostatics. MD integration was carried out using a 2.0 fs time step, employing the SHAKE algorithm<sup>64</sup> on all the hydrogen atoms and a nonbonded cutoff of 10.0 Å. The pair list was updated every 100 steps. Constant pressure (1 atm) and temperature (300 K) was maintained throughout the production simulation run. The systems were allowed to equilibrate under production run conditions for 1 ns before collection of data over 100 ns simulation time. Three sets of simulations for each complex system



were carried out for 100 ns and the total 900 ns trajectories were used for the analysis.

### Data analysis

All the trajectories were analyzed using the PTRAJ module of AMBER12. The trajectories were collected at every 10 ps snapshots over 100 ns of simulation data. Trajectories and structures were visualized using VMD<sup>65</sup> and UCSF Chimera.<sup>58</sup> PTRAJ module was used for RMSD, hydrogen bonding, ion and water mediated interactions. The criterion for hydrogen bonding was set at 3.5 Å distance between electron donor atom and hydrogen of electron acceptor atom and 120° angle cut off. The area under the curve was calculated for H-bond plots. The MMGBSA module of AMBER12 was used for calculating the free energy. The water mediated interactions were calculated for all the trajectories by the CPPTRAJ module of AMBER.

### MM-GBSA calculations

The Molecular Mechanics/Generalized Born Surface Area (MM/GBSA) and Molecular Mechanics/Poisson-Boltzmann Surface Area (MM/PBSA) methods calculate binding free energies for macromolecules by combining molecular mechanics calculations and continuum solvation models.

$$\Delta G_{\text{bind}} = \Delta H - T\Delta S = \langle \Delta E_{\text{gas}} \rangle + \langle \Delta G_{\text{sol}} \rangle - T\langle \Delta S \rangle$$

$$\Delta \Delta G_{\text{bind}} = \Delta G_{\text{complex}} - (\Delta G_{\text{lig}} + \Delta G_{\text{rec}})$$

where  $\Delta E_{\text{gas}}$  is a molecular mechanics energy,  $\Delta G_{\text{sol}}$  is the solvation energy calculated either by solving Poisson's equation or by using the Generalized Born solvation model. The  $\Delta S$  is the entropy contribution to the free energy  $\Delta G_{\text{bind}}$  of the molecule. The snapshots were generated at 1 ns intervals from the trajectories of 100 ns length. These 100 snapshots were used to calculate the binding free energy of all the complexes using the MM/GBSA method.

### MSM analysis

Main objective of MSM analysis is to obtain long time kinetic information from short trajectories of MD simulation.<sup>49,50</sup> This way it can capture slow structural changes from the pool of conformational ensembles generated by MD trajectories. Conformational transitions are essential to the functional role of biomolecules. These are rare events which can be captured by constructing a transition probability matrix on discretised conformational state space.<sup>53,69</sup> To investigate three dimensional conformational landscape and kinetics of the metastable states of tRNA<sub>3</sub><sup>Lys</sup> apo and ligand bound tRNA<sub>3</sub><sup>Lys</sup>, Markov state modeling based analysis was performed on MD trajectory data of each system independently. To build Markov state models PyEMMA2.5.4 package was used.<sup>66</sup> First, for feature selection VAMP<sup>67</sup> score was calculated by choosing various features which are listed in the PyEMMA featurization module. VAMP score was calculated for features like residue mean distance, minimum RMSD and phosphate atom pair distance *etc.* The highest VAMP score was observed for the feature set of all phosphate atom pair

distances. The VAMP score calculation is shown in ESI (Fig. S1†). After selecting features, time independent component analysis (tICA)<sup>68,69</sup> was performed on the coordinates of all phosphate atom pairs. Which gave slow linear subspace by those input coordinates and a dimension reduction was achieved by projecting all coordinates on the 10 slowest tICA components. Where 90% of total kinetic variance is retained by these 10 components. First two tICA components for all the three systems were shown in ESI Fig. S2.† Later, this ten dimensional projected data was clustered into 200 microstates using a regular space clustering method. The lag time, the time scale at which the model is Markovian, was calculated by examining the implied time scales at different lag times. At a given lag time  $\tau$  the implied time scale can be calculated as

$$t_i = \frac{-\tau}{\ln[\lambda_i(\tau)]}$$

where,  $t_i$  is the implied time scale and  $\lambda$  is an eigenvalue of the transition matrix  $T(t)$ .

Lag time is the time interval where the conformational space is optimally discretized and each discretised state space represents the microstate of the system. System progresses dynamically through these discrete states at time points separated by lag time. Lag time interval is obtained such that the collective variables or features on which the discretised conformational space is projected are uncorrelated and their auto covariance is maximum at given lag time. The Markovian property of the MSM model is identified at a certain lag time  $\tau$  above which the implied time scale  $t$  remains constant for further lag time. The lag was determined to be 5 ns for tRNA<sub>3</sub><sup>Lys</sup> apo system, 6 ns for tRNA<sub>3</sub><sup>Lys</sup>-1,4T system and 4 ns for tRNA<sub>3</sub><sup>Lys</sup>-1,5T system (ESI Fig. S3†). The microstates were built using this lag time. Further, Markovian property was assessed by the Chapman Kolmogorov test. Markov states or macrosates are obtained by lumping together kinetically relevant microstates. Chapman Kolmogorov (CK) test gives the optimal number of macrosates that best explains the system's dynamical process. The CK test is performed to see how best the Markovianity of the state space is achieved *i.e.* transition from one state to another microstate, just depends on the state at time  $t$  and not the past transition history of the states. The 200 microstates were coarse grained into 3 macro states using Perron cluster cluster analysis (PCCA).<sup>70</sup> The number of transitions between coarse grained macrostate at an interval of a certain lag time is counted and the count matrix is then symmetrized and normalized to obtain the transition probability matrix ( $T$ ). The mean first passage time (MFPT) between each pair of states in the coarse-grained model was calculated using the MFPT calculation module given in Pyemma.

### MSM validation for both apo and ligand bound tRNA<sub>3</sub><sup>Lys</sup>

The validation of the MSMs for the 300 ns of MD simulation data for the tRNA<sub>3</sub><sup>Lys</sup> apo and 600 ns MD data of ligand bound tRNA<sub>3</sub><sup>Lys</sup> were Markovian in nature was done by performing Chapman Kologmorov test and lag time calculation for each of the three simulated systems separately. In MSM validation, a requirement for Markovian behavior is that the Markov state



model of MD trajectory data should satisfy the Chapman–Kolmogorov equation and the implied timescales remain constant at different lag times. The timescale validation was done by observing the implied timescale plot at different lag times. The implied timescales for the 200-microstate model remain unchanged after a lag time of  $\sim 5$  ns (ESI Fig. S4†), which was thus used to construct the microstate MSM. Further to validate the number of macrostates, the MSM probabilities of microstates in a given macrostate at a given lag time was compared with probabilities directly calculated by trajectory data. These Chapman–Kolmogorov tests of estimated and predicted probabilities of microstates were performed for all the three systems tRNA<sub>3</sub><sup>Lys</sup>, tRNA<sub>3</sub><sup>Lys</sup>-1,4T and tRNA<sub>3</sub><sup>Lys</sup>-1,5T. The test was observed to be satisfying at 3 macrostates for all the three simulated systems (ESI Fig. S5†). Then two hundred representative structures were obtained from each of the three macrostates using PCCA. The three dimensional and two dimensional structural properties of these representative conformations from each macrostate of tRNA<sub>3</sub><sup>Lys</sup> apo system were calculated. Kinetics of transitions within the conformations of these states were obtained by mean first-passage time (MFPTs) calculations.

## Results

### Identification of binding location and orientation of ligands

One of the objectives of the present study is to identify the binding location of the two ligands (1,4T and 1,5T) on tRNA<sub>3</sub><sup>Lys</sup> target molecule and understand the dynamic behavior and stable interactions of ligands with tRNA<sub>3</sub><sup>Lys</sup>. We docked these two ligands on to a tRNA<sub>3</sub><sup>Lys</sup> by selecting the D and TΨC regions as binding pocket and sorted the ligand poses based on their grid scores. The top five best grid scores are given in Table 1 and the best pose for each complex are shown in Fig. 2. The docking results show only the static orientation of bound ligands. In order to further explore the efficiency of these ligands as effective tRNA<sub>3</sub><sup>Lys</sup> inhibitors, molecular dynamics (MD) simulations and binding free energy calculations were carried out.

To understand the binding specificity of these two ligands against tRNA<sub>3</sub><sup>Lys</sup>, docking of these two ligands on different tRNAs like tRNA<sup>Phe</sup>, tRNA<sup>Asp</sup> have been carried out by considering the crystal structures. The selected tRNAs are from different sources namely, yeast, *E. coli*, synthetic, and their pdb ids are 1EHZ, 1VTQ, 3L0U, 4TNA and 3TRA respectively.<sup>71–75</sup> The best docking scores of 1,4T and 1,5T ligands against selected

tRNAs along with tRNA<sub>3</sub><sup>Lys</sup> (1FIR) have been given in Table 2 and ESI Fig. S6.† Both the ligands showed best binding scores with tRNA<sub>3</sub><sup>Lys</sup> as compared to other tRNAs.

From the docking results, the best pose for each ligand bound complex systems (tRNA<sub>3</sub><sup>Lys</sup>-1,4T and tRNA<sub>3</sub><sup>Lys</sup>-1,5T) along with a control system tRNA<sub>3</sub><sup>Lys</sup> were selected for MD simulations. The aim of the MD simulations was to explore the structural dynamics of the tRNA<sub>3</sub><sup>Lys</sup> apo as well as ligand-bound tRNA<sub>3</sub><sup>Lys</sup>.

### Intra and intermolecular interactions that affect the stability of the complexes

To find out the overall stability of the system, the hydrogen bonds were calculated for each system and compared. This stability was evaluated by calculating hydrogen bonding percentages (HB%) which is defined as the time over a hydrogen bond. A hydrogen bond is considered where the distance between donor and acceptor atoms is  $< 3.5$  Å and the angle between the donor atom, hydrogen attached to donor and the acceptor atom is  $< 120^\circ$ . These H-bonds include all intermolecular and intra-molecular interactions which satisfy the above criteria. The H-bonds are plotted here by taking the number of H-bonds on X-axis and their residence time on Y-axis as shown in Fig. 3. In case of the first trajectory set (R1) the tRNA<sub>3</sub><sup>Lys</sup> system was showing the highest residence time for a higher number of H-bonds compared to the other two systems. But in the case of second and third trajectory sets (R2) and (R3) the tRNA<sub>3</sub><sup>Lys</sup>-1,4T system was showing the highest residence time for a higher number of H-bonds compared to other two systems. Whereas the tRNA<sub>3</sub><sup>Lys</sup>-1,5T system was always showing low residence time for less number of H-bonds in all the trajectory sets. These observations for tRNA<sub>3</sub><sup>Lys</sup>-1,4T system can be attributed to greater stability of the system. The area under the curve was also calculated for these H-bond plots and given in ESI Table S7.† The values of the area under the curve for the tRNA<sub>3</sub><sup>Lys</sup>-1,4T system was higher as compared to tRNA<sub>3</sub><sup>Lys</sup> system and tRNA<sub>3</sub><sup>Lys</sup>-1,5T system for two sets of trajectories.

The intermolecular interactions between the ligand and tRNA<sub>3</sub><sup>Lys</sup> were also calculated as they may also play an important role in the complex stability and functionality of the system. The detailed list of H-bond interactions between ligand and tRNA<sub>3</sub><sup>Lys</sup> for the two complexes tRNA<sub>3</sub><sup>Lys</sup>-1,4T system and tRNA<sub>3</sub><sup>Lys</sup>-1,5T system were given in Table 3. In the case of tRNA<sub>3</sub><sup>Lys</sup>-1,4T system, the 1,4T ligand interacts with G52, G53 and U60 residues of tRNA<sub>3</sub><sup>Lys</sup> through H-bonds. The 1,4T ligand forms strong and stable H-bonds with G53 and U60 for more than 70% of simulation time (Fig. 4a). The particular H-bond interaction of the 1,4T ligand with U60 residue tRNA<sub>3</sub><sup>Lys</sup> was observed in all the three sets of simulations indicating that it might be one of the crucial interactions. In the tRNA<sub>3</sub><sup>Lys</sup>-1,5T system, the 1,5T ligand interacts with A50 residue of tRNA<sub>3</sub><sup>Lys</sup> through H-bonds (Fig. 4b). The 1,5T ligand forms H-bonds with A50 which are stable for not more than 50% of simulation time as given in Table 3. It was also observed that the aromatic ring of 1,4T ligand formed stacking interactions with 5MC49 (5-methylcytosine) residue of tRNA<sub>3</sub><sup>Lys</sup> as shown in Fig. 4c and d. Since

Table 1 The ligand binding grid scores of top 5 ranked ligand poses for both the complex systems

	tRNA <sub>3</sub> <sup>Lys</sup> -1,4T system	tRNA <sub>3</sub> <sup>Lys</sup> -1,5T system
	Energy (kcal mol <sup>-1</sup> )	Energy (kcal mol <sup>-1</sup> )
Rank 1	-48.92	-46.58
Rank 2	-47.65	-44.91
Rank 3	-45.38	-44.14
Rank 4	-44.76	-42.42
Rank 5	-43.72	-42.35



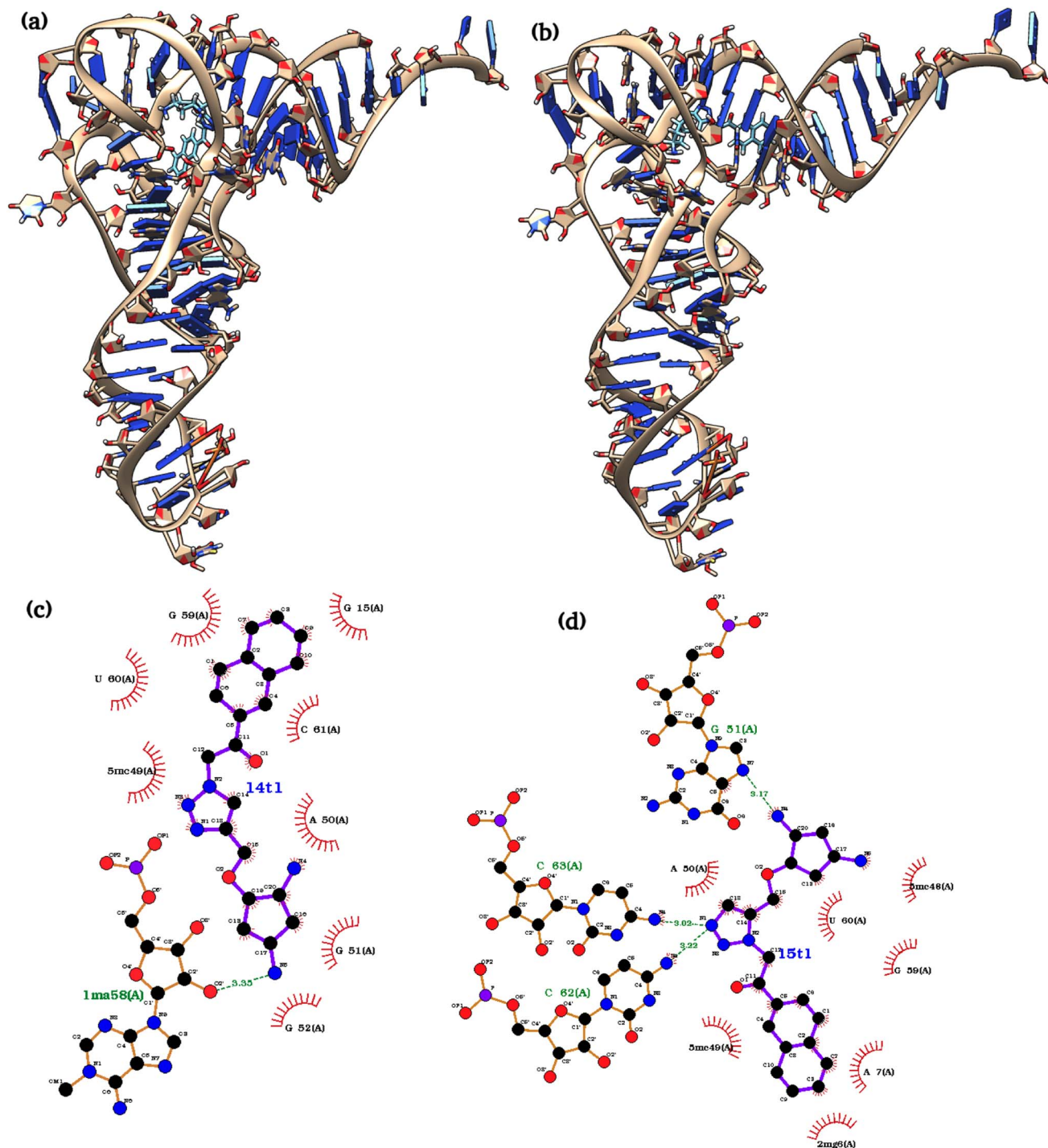


Fig. 2 The structures of best docking poses of both ligands (a) tRNA<sub>3</sub><sup>Lys</sup>-1,4T system (b) tRNA<sub>3</sub><sup>Lys</sup>-1,5T system (c) ligplot of 1,4T interactions (d) ligplot of 1,5T interactions.

the intermolecular interactions are found to be high for the tRNA<sub>3</sub><sup>Lys</sup>-1,4T system as compared to tRNA<sub>3</sub><sup>Lys</sup>-1,5T system, which may infer that the 1,4T ligand shows strong binding to the tRNA<sub>3</sub><sup>Lys</sup>. The additional stacking interactions found for tRNA<sub>3</sub><sup>Lys</sup>-1,4T system may also support the greater stability of the 1,4T ligand binding. The complete list of ligand interactions with the tRNA<sub>3</sub><sup>Lys</sup> are given in ESI Table S8.†

#### Identification of ion and water mediated interactions in the complexes

Apart from the intermolecular interactions between ligand and tRNA<sub>3</sub><sup>Lys</sup>, the water mediated and ion mediated interactions were also identified in the systems. It has been identified from the MD simulations of tRNA<sub>3</sub><sup>Lys</sup>-1,4T system that one of the Mg<sup>2+</sup> ions was able to maintain stable interaction with the N5



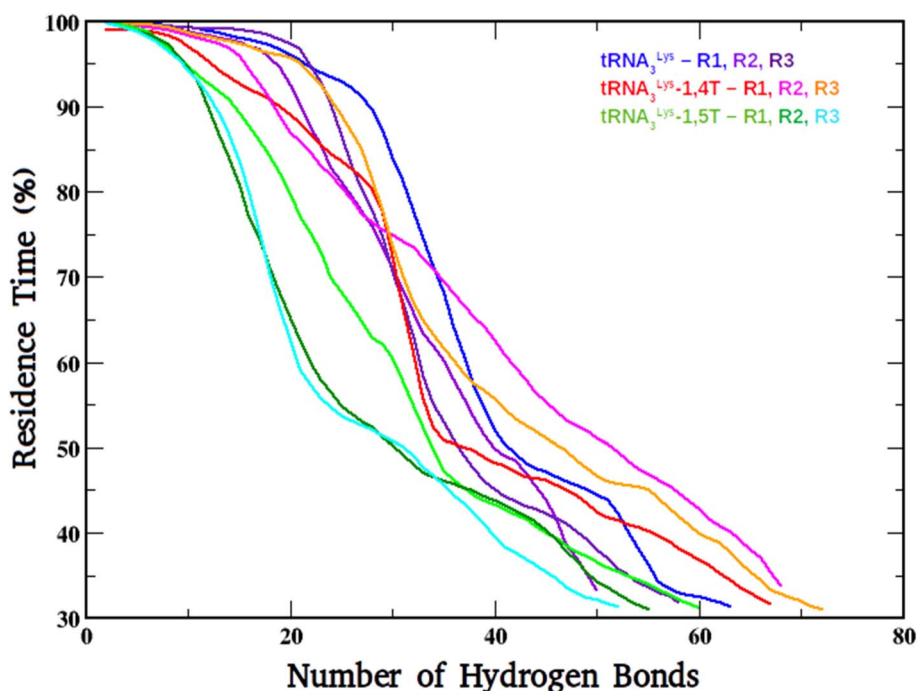
**Table 2** The ligand binding grid scores of 1,4T and 1,5T ligands against different tRNA crystal structures and MSM-cluster representative structures

tRNA structure	1,4T ligand	1,5T ligand
ID	Energy (kcal mol <sup>-1</sup> )	Energy (kcal mol <sup>-1</sup> )
1EHZ	-43.38	-45.98
1VTQ	-47.69	-46.37
3LOU	-45.26	-45.66
4TNA	-43.55	-45.64
3TRA	-43.30	-43.44
1FIR	-52.68	-45.60
PCCA1	-45.21	-45.70
PCCA2	-46.30	-44.85
PCCA3	-43.50	-44.60

atom of 1,4T ligand molecule with a 2 Å distance as shown in Fig. 5. The same Mg<sup>2+</sup> ion was able to maintain H-bond interaction with residue G59 of tRNA<sub>3</sub><sup>Lys</sup>. It also shows a weak interaction with the residue U60 of tRNA<sub>3</sub><sup>Lys</sup>. This particular Mg<sup>2+</sup> ion mediated interaction was in coordination with the ligand and the G59, U60 residues of TΨC arm which may strengthen the complex. In the case of tRNA<sub>3</sub><sup>Lys</sup>-1,5T system, it has been seen that one of the Mg<sup>2+</sup> ions was able to maintain stable interaction with the N5 atom of 1,5T ligand molecule with a 2 Å distance. It also shows a weak interaction with the residue 5MC48 (5-methylcytosine) of tRNA<sub>3</sub><sup>Lys</sup>. The particular interactions of these two ligands (1,4T and 1,5T) with the Mg<sup>2+</sup> ions have been found to be quite stable. The maintenance of stable interaction of Mg<sup>2+</sup> ions with particular residues may also

help the overall stability of the specific TΨC loop or whole structure. The distance of few Mg<sup>2+</sup> ions with the bound ligand and tRNA<sub>3</sub><sup>Lys</sup> for combined trajectories (last 50 ns of each run of respective trajectories joined together to make combined trajectory of 150 ns) have been given in ESI Fig. S9.†

The water molecules which are present in proximity interact with specific residues and may play an important role in the stability or functionality of the system. According to Roh *et al.*<sup>76</sup> higher density of water molecules makes the tRNA more flexible. In simulations, the number of water molecules in the lower hydration shell was determined using the criterion that the water oxygen atoms were within 3.4 Å of non-hydrogen atoms of the tRNA<sub>3</sub><sup>Lys</sup>. The distance criteria for the upper hydration shell was 5 Å. We have also calculated lower and upper hydration shells specifically for ligands. It was observed that the tRNA<sub>3</sub><sup>Lys</sup>-1,4T system showed slightly higher hydration number compared to tRNA<sub>3</sub><sup>Lys</sup>-1,5T system. However, in case of hydration shell around the ligands, the 1,4T ligand show more number of water molecules compare to 1,5T ligand as shown in ESI Fig. S10.† The residues which are involved in water mediated interactions and have their residence time above 40% are given in ESI Table S11.† In the case of the tRNA<sub>3</sub><sup>Lys</sup> apo system, the residue 2MG6 (2*N*-methyl guanosine) interacts with a water molecule for the highest simulation time of 60.12%. The residues of tRNA<sub>3</sub><sup>Lys</sup> form a maximum of 12 interactions with water molecules above 40% of simulation time in a trajectory. In all the three trajectories of tRNA<sub>3</sub><sup>Lys</sup>, five water mediated interactions above 50% of simulation time have been observed. In the case of the tRNA<sub>3</sub><sup>Lys</sup>-1,4T system, the residue C13 interacts with the two water molecules for the highest simulation time of 77.85% and 76.13% respectively. The residues of tRNA<sub>3</sub><sup>Lys</sup> form maximum

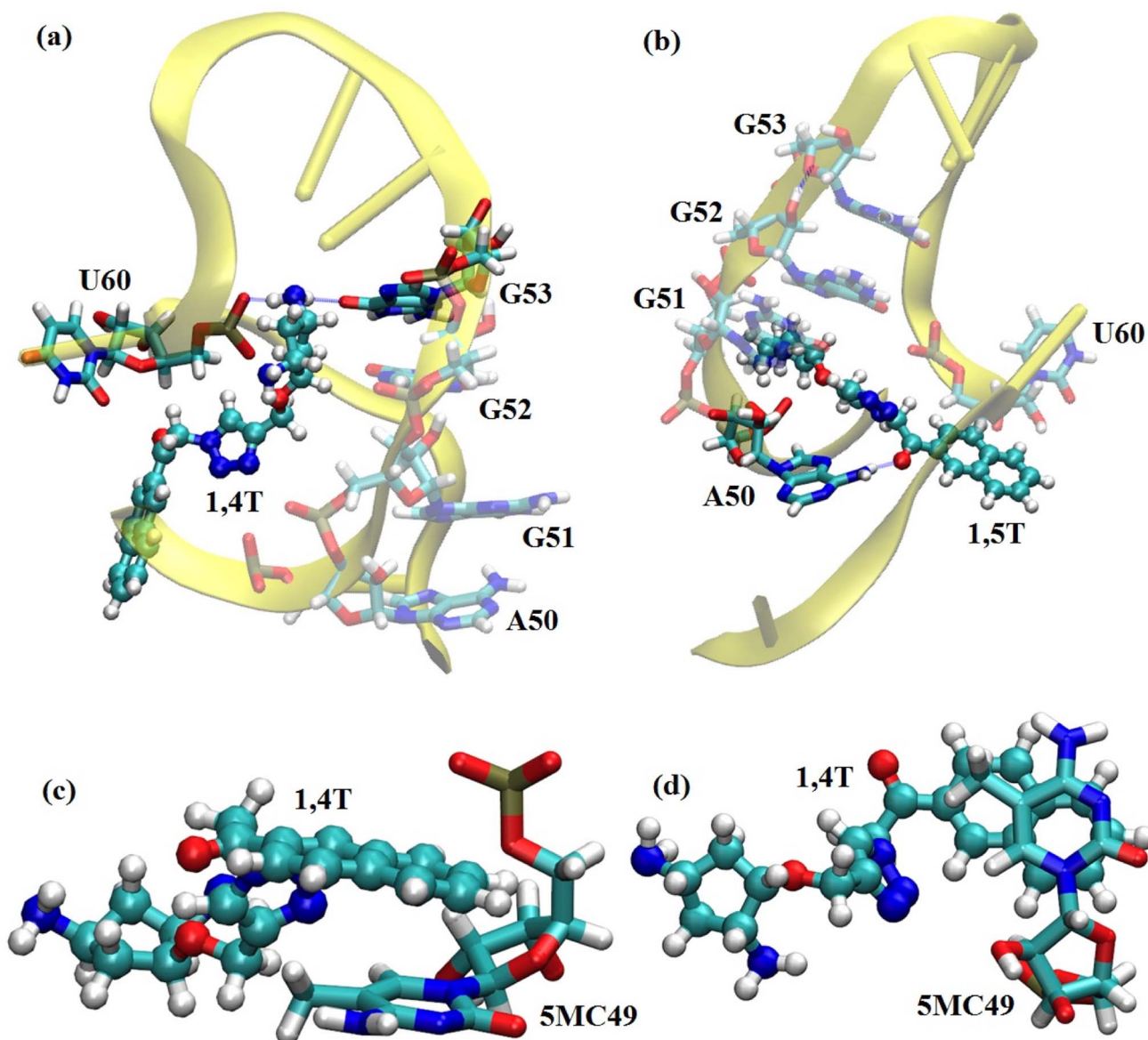


**Fig. 3** The H-bonding plot of tRNA<sub>3</sub><sup>Lys</sup> system, tRNA<sub>3</sub><sup>Lys</sup>-1,4T system and tRNA<sub>3</sub><sup>Lys</sup>-1,5T systems for all the three simulation runs (R1, R2, R3).



**Table 3** H-bond interactions between the tRNA<sub>3</sub><sup>Lys</sup> and ligand for both the systems (tRNA<sub>3</sub><sup>Lys</sup>-1,4T system and tRNA<sub>3</sub><sup>Lys</sup>-1,5T system) above 40% of residence time

Residues	H-bond	tRNA <sub>3</sub> <sup>Lys</sup> -1,4T system			tRNA <sub>3</sub> <sup>Lys</sup> -1,5T system		
		Traj1	Traj2	Traj3	Traj1	Traj2	Traj3
A50-Lig	N6-H62...O1	—	—	—	—	46.77%	—
A50-Lig	N6-H62...O1	—	—	—	—	45.98%	—
A50-Lig	N6-H62...N1	—	—	—	—	—	44.58%
G52-Lig	O6...H23-N5	—	—	54.37%	—	—	—
G53-Lig	O6...H23-N5	91.07%	—	—	—	—	—
U60-Lig	O2P...H22-N5	77.92%	75.15%	56.94%	—	—	—

**Fig. 4** (a) The tRNA<sub>3</sub><sup>Lys</sup> and 1,4T ligand intermolecular interactions. (b) The tRNA<sub>3</sub><sup>Lys</sup> and 1,5T ligand intermolecular interactions. The stacking interaction between 1,4T ligand and 5MC49 residue of tRNA<sub>3</sub><sup>Lys</sup> side view (c) and top view (d).

14 interactions with water molecules above 40% of simulation time in a trajectory for tRNA<sub>3</sub><sup>Lys</sup>-1,4T system. In all the three trajectories of tRNA<sub>3</sub><sup>Lys</sup>-1,4T system, six water mediated

interactions above 50% of simulation time have been observed. In the case of tRNA<sub>3</sub><sup>Lys</sup>-1,5T system, the residue PSU27 (pseudouridine) is showing interaction with the water molecule for



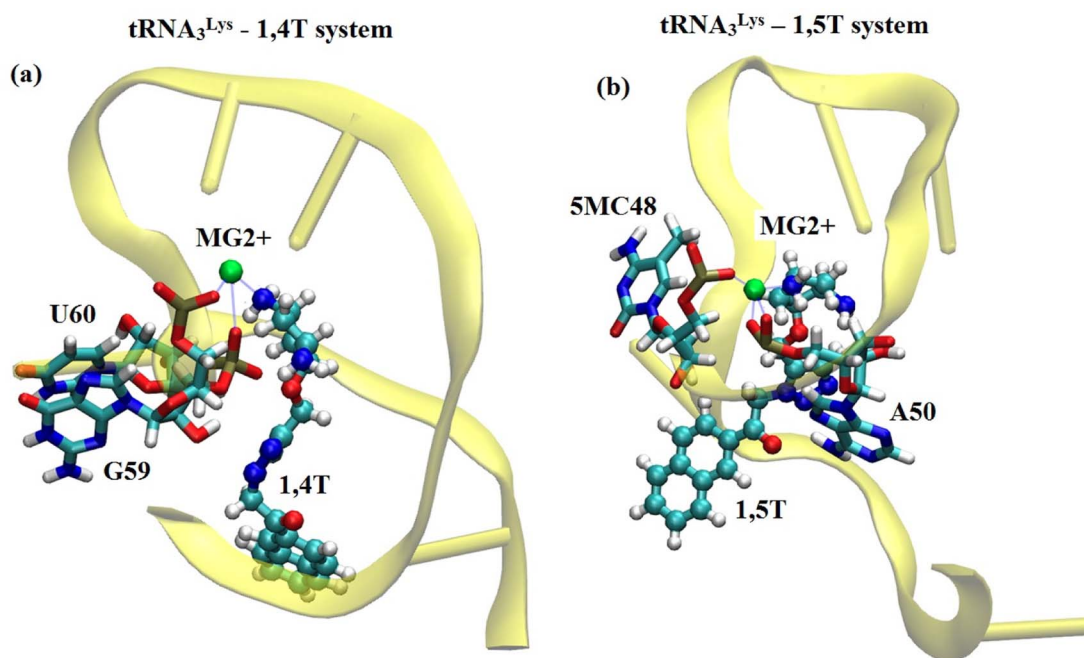


Fig. 5 The ion mediated interactions of 1,4T and 1,5T ligands and various residues of respective complexes. (a) The structures of  $Mg^{2+}$  ion interactions with 1,4T ligand and other residues (G59, U60) of  $tRNA_3^{Lys}$ . (b) The structures of  $Mg^{2+}$  ion interactions with 1,5T ligand and other residues (5MC48, A50) of  $tRNA_3^{Lys}$ .

the highest simulation time of 58.19%. The residues of  $tRNA_3^{Lys}$  form maximum 11 interactions with water molecules above 40% of simulation time in a trajectory for  $tRNA_3^{Lys}$ -1,5T system. In all the three trajectories of  $tRNA_3^{Lys}$ -1,5T system, five water mediated interactions above 50% of simulation time have been observed. The water mediated interactions for the three complex systems are shown in Fig. 6.

It has been observed from the water mediated interactions that majority of the water molecules are interacting with T $\Psi$ C arm residues for all the three systems as given in ESI Table S11.† In case of the  $tRNA_3^{Lys}$  apo system, out of 5 water mediated interactions three are with D-arm and one for each with T $\Psi$ C arm and anticodon arm were formed. In case of  $tRNA_3^{Lys}$ -1,4T system, out of 8 water mediated interactions four for each with D-arm and T $\Psi$ C arm were formed. No water mediated interactions were formed with anticodon residues for the  $tRNA_3^{Lys}$ -1,4T system. In case of  $tRNA_3^{Lys}$ -1,5T system, out of 8 water mediated interactions 2, 3 and 3 interactions formed with D-arm, T $\Psi$ C arm and anticodon arm respectively.

### Structural stability/binding free energy of the three complexes

The binding free energies were calculated for the two ligands with  $tRNA_3^{Lys}$  by MMGBSA method. These calculated free energy values for both the ligand bound complexes  $tRNA_3^{Lys}$ -1,4T system and  $tRNA_3^{Lys}$ -1,5T system were compared and shown in Fig. 7. In both the complexes, the  $tRNA_3^{Lys}$  was considered as a receptor and 1,4T and 1,5T were considered as ligands for their respective systems in the MMGBSA free energy calculations. The average  $\Delta G$  values for three trajectory sets of  $tRNA_3^{Lys}$ -1,4T system are  $-64.58 \text{ kcal mol}^{-1}$ ,  $-57.18 \text{ kcal mol}^{-1}$

and  $-65.68 \text{ kcal mol}^{-1}$  respectively. The average  $\Delta G$  values for three trajectory sets of  $tRNA_3^{Lys}$ -1,5T system are  $-33.90 \text{ kcal mol}^{-1}$ ,  $-24.04 \text{ kcal mol}^{-1}$  and  $-35.02 \text{ kcal mol}^{-1}$  respectively. However, the average  $\Delta G$  values vary for different trajectories of the same complex but the difference between both the complexes are maintaining consistent  $\Delta G$  values. These results indicate that the 1,4T ligand may strongly bind to the  $tRNA_3^{Lys}$ . All the MMGBSA free energy components for both the ligand bound systems were given in ESI Table S12.† It has been observed that dihedral angle energies and van der Waals energies are the major contributions for overall free energy of these systems.

### Exploring conformational flexibility and kinetics using MSM analysis

MSM analysis was performed to understand global conformational landscape as well as local conformational landscape of  $tRNA_3^{Lys}$  apo and 1,4T and 1,5T ligand bound  $tRNA_3^{Lys}$  systems. MSM analysis was performed on the  $tRNA_3^{Lys}$  apo system and then the 1,4T and 1,5T bound  $tRNA_3^{Lys}$  systems. The trajectories were analysed by MSM to understand local and global kinetics of transitions in the conformations of  $tRNA_3^{Lys}$  apo system as well as perturbations in the conformations induced by ligands.

### MSM structural insights to $tRNA_3^{Lys}$ apo system

To quantify global structural variations among the macrostates obtained by MSM analysis, an average RMSD of 200 conformations from each macrostate of  $tRNA_3^{Lys}$  apo system was calculated. The three average values of RMSD of the whole structure representing the three macrostate conformations of



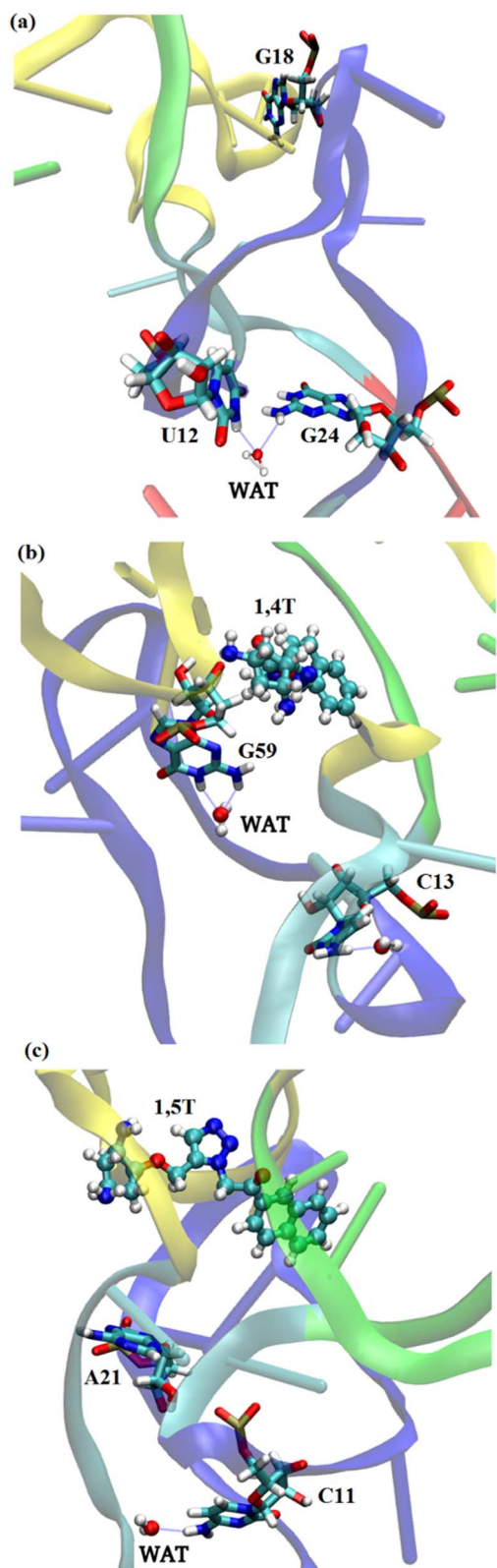


Fig. 6 The water mediated interactions of various residues of all the three complexes (a)  $\text{tRNA}_3^{\text{Lys}}$  system (b)  $\text{tRNA}_3^{\text{Lys}}\text{-1,4T}$  system (c)  $\text{tRNA}_3^{\text{Lys}}\text{-1,5T}$  system.

$\text{tRNA}_3^{\text{Lys}}$  apo systems were compared with the average RMSD values of the MSM macrostates of 1,4T and 1,5T systems. The average RMSD values and standard deviations are given in Table 4. RMSD of  $\text{tRNA}_3^{\text{Lys}}$  apo, MSM macrostate conformations were observed to be having wide range of values (8.38 Å, –1.33 Å) and standard deviation values ( $\sim 1.5$  Å) indicate that apo  $\text{tRNA}_3^{\text{Lys}}$  MSM conformations belonging to different macrostate possess more degrees of freedom and flexibility. To assess the flexibility in the whole  $\text{tRNA}_3^{\text{Lys}}$  structure and quantify the local variations which were contributing maximum to overall conformational degrees of freedom, RMSD of different regions of  $\text{tRNA}_3^{\text{Lys}}$  apo system were calculated *viz* D loop, T $\Psi$ C loop and anticodon regions. These region-specific RMSD values were calculated for the MSM macrostate of  $\text{tRNA}_3^{\text{Lys}}$  apo and  $\text{tRNA}_3^{\text{Lys}}\text{-1,4T}$  and  $\text{tRNA}_3^{\text{Lys}}\text{-1,5T}$  systems. As compared to the pdb structure (PDB id 1FIR) the conformations drawn from second (green) and third (blue) MSM macrostates of  $\text{tRNA}_3^{\text{Lys}}$  apo have shown deviation of the anticodon loop. It was also captured in RMSD values which were varying in the range (1 Å to 6 Å). The RMSD for anticodon regions of conformations from 1<sup>st</sup> MSM macrostate (red) were observed to be conserved with respect to pdb structure. However, D loop RMSD values in case of  $\text{tRNA}_3^{\text{Lys}}$  apo system were seen to have high fluctuations for conformations belonging to the third (blue) macrostate and in consequence this may indicate the distortion of the secondary structure base pairing in D arm. RMSD for the T $\Psi$ C loop of the three macrostates of  $\text{tRNA}_3^{\text{Lys}}$  apo showed fluctuation in the range of 2 Å to 4 Å indicating that T $\Psi$ C loop is conserved in all the three macrostate conformations.

Earlier NMR studies have reported<sup>77</sup> that the  $\text{tRNA}_3^{\text{Lys}}$  apo structure has a complex three dimensional structural landscape. The MSM analysis reported here has shown that there were majorly three macrostate conformations which dominate the three dimensional conformational space of  $\text{tRNA}_3^{\text{Lys}}$  apo. These macrostates were verified using the Chapman Kolmogorov test. Conformations belonging to distinct macrostates were further analysed by calculating base pair (secondary and tertiary interactions) distances from four main regions of the  $\text{tRNA}_3^{\text{Lys}}$  apo system, *viz* acceptor arm, D arm, T $\Psi$ C arm and anticodon arm. The base pairs distances which are responsible for formation of secondary and tertiary structures and reported in NMR studies<sup>77</sup> were calculated for all the 200 conformations belonging to each of the three macrostates of  $\text{tRNA}_3^{\text{Lys}}$  apo. The G6–U67 pair which is responsible for formation of secondary structure of the acceptor arm was found to be fluctuating with a broad range of values  $\sim 3$  Å to 15 Å which indicates that acceptor arm in  $\text{tRNA}_3^{\text{Lys}}$  apo has diverse range of conformation in all the three macrostates (ESI Fig. S13<sup>†</sup>). The D arm intra interactions namely U8–A14, C13–G22 and A14–A21 were calculated (Fig. 8) to check the stability of secondary structure in all the three macrostate conformations of  $\text{tRNA}_3^{\text{Lys}}$  apo system. Although, U8–A14 and C13–G22 interaction distances were observed to be conserved, A14–A21 distance was observed to be varying in the range of 10 Å to 14 Å in all the three macrostates, which indicates that D loop structure spans wide range of conformations for A14–A21 interaction. The tertiary interactions across D and T $\Psi$ C loops of  $\text{tRNA}_3^{\text{Lys}}$  which are known to



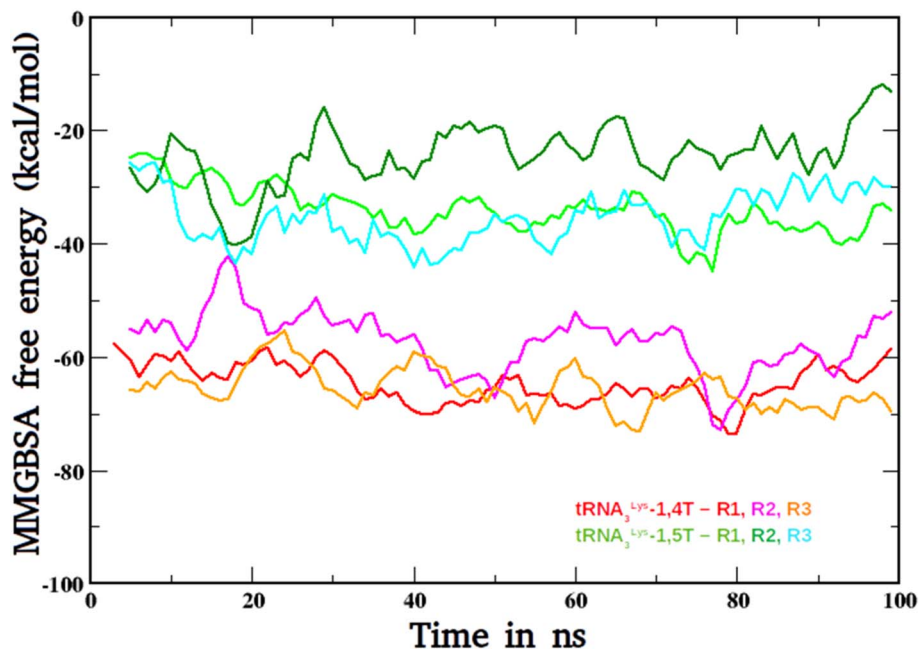


Fig. 7 The MMGBSA-free energy plots of tRNA<sub>3</sub><sup>Lys</sup>-1,4T system and tRNA<sub>3</sub><sup>Lys</sup>-1,5T systems for all the three simulation runs (R1, R2, R3).

Table 4 The RMSD values of MSM metastable state structures of three complexes

RMSD	MSM metastable state	Average RMSD (Å)	Standard deviation (Å)	Minimum RMSD (Å)	Maximum RMSD (Å)
tRNA <sub>3</sub> <sup>Lys</sup>	PCCA1	5.091	1.514	1.57	8.384
	PCCA2	4.826	1.476	1.523	8.339
	PCCA3	4.135	1.66	1.339	8.223
tRNA <sub>3</sub> <sup>Lys</sup> - 1,4T	PCCA1	5.456	0.495	4.446	6.798
	PCCA2	5.57	0.49	4.329	6.754
	PCCA3	5.572	0.573	3.853	6.766
tRNA <sub>3</sub> <sup>Lys</sup> - 1,5T	PCCA1	5.652	0.319	4.403	6.5
	PCCA2	5.203	0.385	4.526	6.251
	PCCA3	5.532	0.25	4.483	6.287

form tertiary structure, mainly U16–G59, G18–Ψ55 and G19–C56 (Fig. 9) were observed to be greater than 12 Å in all the three microstates of tRNA<sub>3</sub><sup>Lys</sup> apo system.

The residue–residue interaction distances across D and TΨC loop indicate the opening of tertiary structure and flexibility of TΨC loop. The major structural difference in 3D conformations segregated by MSM macrostates were observed in the anticodon arm as shown in Fig. 10.

To quantify anticodon structural insights of MSM macrostates, U33–A37 interaction distance which lies in the anticodon loop of tRNA<sub>3</sub><sup>Lys</sup> was calculated. It was observed that the first macrostate (red) of tRNA<sub>3</sub><sup>Lys</sup> apo maintains this distance as that of PDB crystal structure. On the other hand, the remaining two MSM states of tRNA<sub>3</sub><sup>Lys</sup> apo systems (green and blue structures) have shown that more than 90% of conformers have a wide range of variation in the U33–A37 interaction distance. This may indicate that the anticodon loop of tRNA<sub>3</sub><sup>Lys</sup> apo attains noncanonical conformational metastable states. Apart from

anticodon loop structural deviations, the D loop and TΨC loop interaction (C11–G45) was shown wide range of variation ~9 Å to 14 Å (ESI Fig. S14<sup>†</sup>) in all the three MSM metastable states.

The mean time required for the tRNA<sub>3</sub><sup>Lys</sup> apo system to reach a given ending macrostate *j* from a different starting macrostate *i*, is known as the mean first passage time (MFPT for *i* to *j*). The MFPT times were calculated for three macrostates obtained by MSM analysis. The value of MFPT is inversely proportional to the rate of transitions between any two macrostates and the difference in MFPT of any two macrosates indicates difference in heights of free energy barriers of those states. The transition for tRNA<sub>3</sub><sup>Lys</sup> apo ranges from 1 ns to 260 ns and has an average value of 111.80 ns for all 3 possible transitions. MFPT for all the macrostates of tRNA<sub>3</sub><sup>Lys</sup> apo system are shown in Fig. 11. Transitions from state-1 to state-3 and from state-2 to state-3 occurred most quickly which were approximately 1 ns for both the paths. The slowest transition was observed from state-3 to state-1 which was measured to be 260 ns.



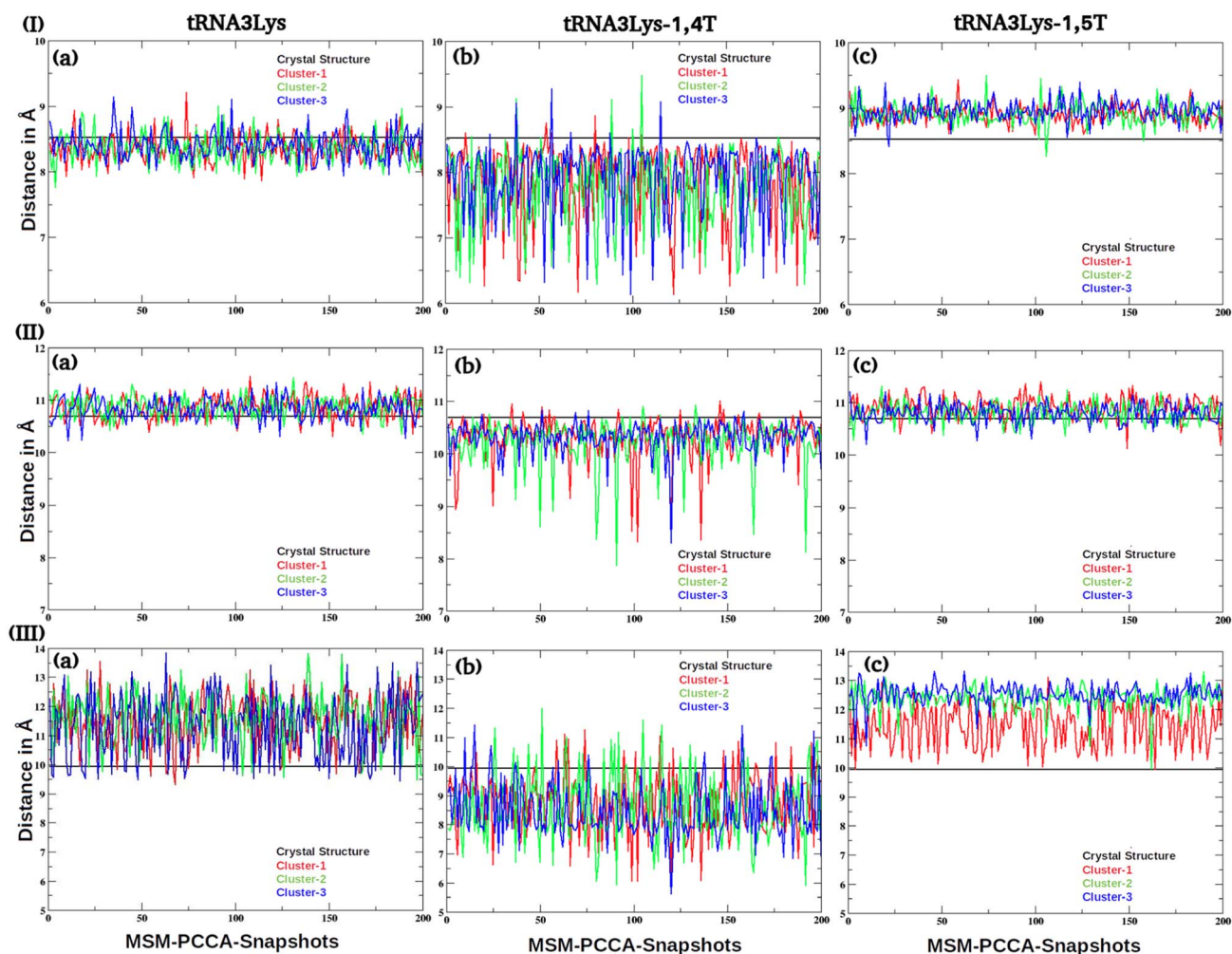


Fig. 8 The inter residue distances of (I) U8–A14, (II) C13–G22 and (III) A14–A21 base pairs were given for (a)  $\text{tRNA}_3^{\text{Lys}}$  (b)  $\text{tRNA}_3^{\text{Lys}}-1,4\text{T}$  and (c)  $\text{tRNA}_3^{\text{Lys}}-1,5\text{T}$  complexes.

Docking of 1,4T and 1,5T ligands against three representative structures, one from each MSM macrostate was carried out. The binding location and the docking scores have been given in ESI Fig. S15† and Table 2.

### MSM structural insights to ligand bound $\text{tRNA}_3^{\text{Lys}}$

To investigate the effect of 1,4T bound and 1,5T bound ligands on  $\text{tRNA}_3^{\text{Lys}}$  conformations, the MSM analysis for 1,4T and 1,5T ligand bound systems were performed on MD trajectory data of each system. While constructing MSM metastable states the ligands were removed from the trajectory data to focus on structural variations induced in the three dimensional  $\text{tRNA}_3^{\text{Lys}}$  conformational landscape, by each ligand.

The presence of ligands 1,4T or 1,5T was observed to be affected by decreasing the range of conformations explored by  $\text{tRNA}_3^{\text{Lys}}$  apo. The RMSD values for both the ligand systems were calculated for the whole system as well as region wise. The range of RMSD values of 1,4T system and 1,5T ligand bound systems were observed to be reduced ( $\sim 4 \text{ \AA}$  to  $6 \text{ \AA}$ ) i. e., approximately within  $\sim 2 \text{ \AA}$  span of values as shown in Table 4.

Whereas the  $\text{tRNA}_3^{\text{Lys}}$  apo system was shown to have quite a wider range of RMSD values (with the range of  $6 \text{ \AA}$ ).

To understand region wise effect of ligands on RMSD values, RMSD of anticodon arm, D loop and T $\Psi$ C loop were calculated for all the conformations of MSM macrostates of both the ligand bound systems. The anticodon arm of the  $\text{tRNA}_3^{\text{Lys}}-1,4\text{T}$  system in all the three metastable states have shown a range of values ( $\sim$ min  $1.5 \text{ \AA}$  to max  $5.5 \text{ \AA}$ ) which was lesser than the  $\text{tRNA}_3^{\text{Lys}}$  apo anticodon RMSD range. However, D loop and T $\Psi$ C loop RMSD of all the three macrostates conformers of 1,4T system have shown  $\sim 1 \text{ \AA}$  increase in the range of values as compared to  $\text{tRNA}_3^{\text{Lys}}$  apo system. On the other hand, in the case of the 1,5T system, RMSD of the whole structure was seen to be significantly reduced ( $\sim 6.5 \text{ \AA} \pm 2 \text{ \AA}$ ) as compared to  $\text{tRNA}_3^{\text{Lys}}$  apo system. This global reduction in RMSD values was also observed in RMSD of the anticodon arm of the 1,5T system where it was seen to be significantly reduced in conformations from the first macrostate (red). The anticodon RMSD values for conformations in second (green) and third (blue) have more range of values indicating fluctuations in secondary structure of



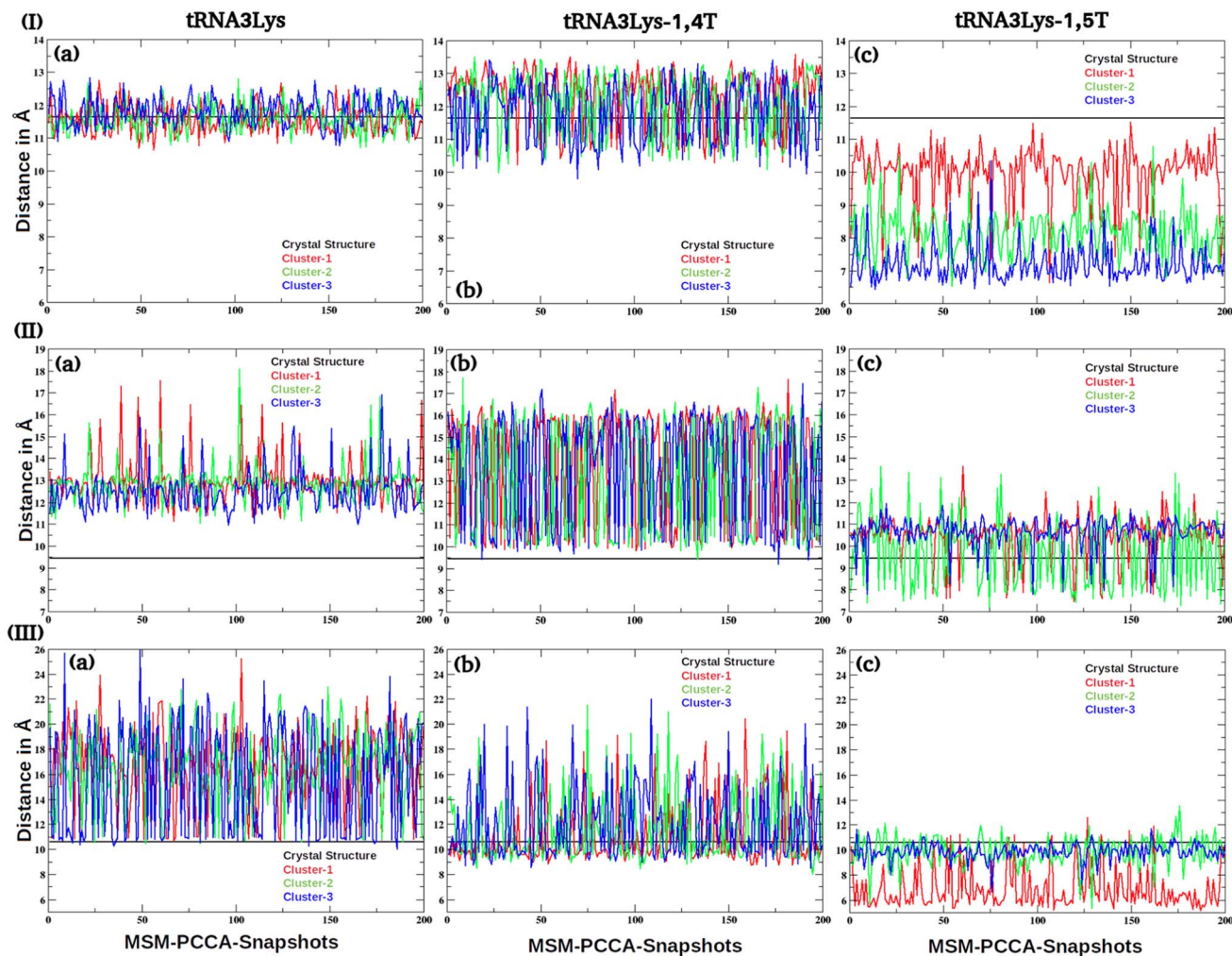


Fig. 9 The inter residue distances of (I) U16–G59, (II) G18– $\Psi$ 55 and (III) G19–C56 interactions were given for (a)  $\text{tRNA}_3^{\text{Lys}}$  (b)  $\text{tRNA}_3^{\text{Lys}}-14,\text{T}$  and (c)  $\text{tRNA}_3^{\text{Lys}}-1,5\text{T}$  complexes.

the anticodon region of the 1,5T system. In case of D arm, conformers from third (blue) MSM macrostate of 1,5T system have shown maximum fluctuations in RMSD values while conformers from first (red) and second (green) MSM macrostate have been observed to be significantly conserved as compared to the pdb structure.

The acceptor arm interaction G6–U67 was observed to be having a fluctuating wide range of values in a 1,4T ligand bound system which was similar to apo  $\text{tRNA}_3^{\text{Lys}}$  system. However, in case of 1,5T bound system variations of G6–U67 interaction distance (ESI Fig. S13<sup>†</sup>) was observed to be significantly reduced in conformers from first (red) and second (green) MSM macrostates whereas conformers in 3<sup>rd</sup> macrostate have similar kind of fluctuation range as compared to  $\text{tRNA}_3^{\text{Lys}}$  apo. The interactions U8–A14, C13–G22 and A14–A21 play an important role in secondary structure formation of the D loop, were calculated in ligand bound systems. The interactions U8–A14 and C13–G22 were found to be maintained in 1,4T and 1,5T MSM macrostate conformers (Fig. 8). However, the average distance between A14–A21 was observed to be reduced by 3 Å in

the 1,4T ligand bound system as compared with  $\text{tRNA}_3^{\text{Lys}}$  apo (Fig. 8). Whereas, the same interaction (A14–A21) was shown to have wider range of variation in conformers from 1st macrostate (red) of 1,5T and the other two macrostate (blue and green) conformers have shown average increase in distance ( $\sim 13$  Å). Therefore, the D loop secondary structure was found to be stable in the 1,4T bound system and disturbed due to increased distance of the A14–A21 interaction in the 1,5T system. The tertiary structure interactions across the D–T $\Psi$ C loops, U16–G59, G18– $\Psi$ 55 and G19–C56 were calculated to understand how the three dimensional structural landscape of  $\text{tRNA}_3^{\text{Lys}}$  gets perturbed in the presence of ligands. In the 1,4T system, the G18– $\Psi$ 55 residue distance was observed to have large range of variation ( $\sim 10$  Å to 16.5 Å) in all the three MSM macrostate conformers (Fig. 9). In the 1,5T system too, similar range of varying G18– $\Psi$ 55 distance was observed in only second MSM macrostate represented in green. Although, conformers belonging to first (red) and third (blue) macrostate have shown reduced fluctuation of G18– $\Psi$ 55 distance. Another tertiary structure forming base pair G19–C56 distance in the 1,4T



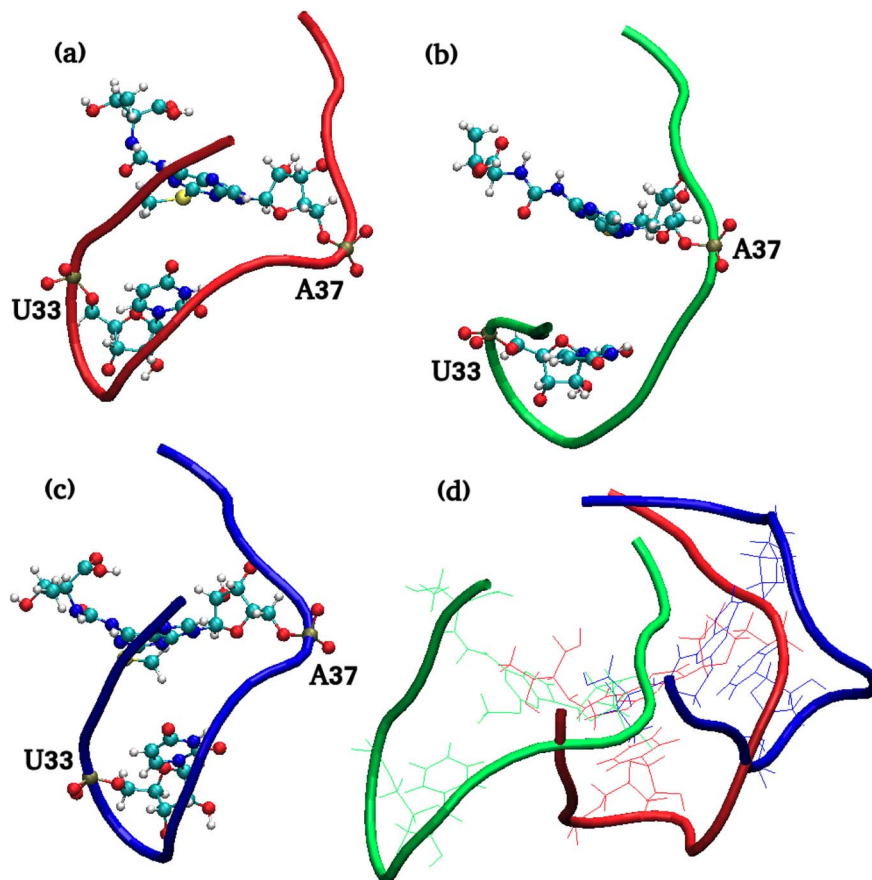


Fig. 10 MSM macrostate structures of anticodon loop of (a)  $\text{tRNA}_3^{\text{Lys}}$  (b)  $\text{tRNA}_3^{\text{Lys}}\text{-14T}$  (c)  $\text{tRNA}_3^{\text{Lys}}\text{-15T}$  and (d) 3 structures superimposed. Residues 33 and 37 are shown in CPK style.

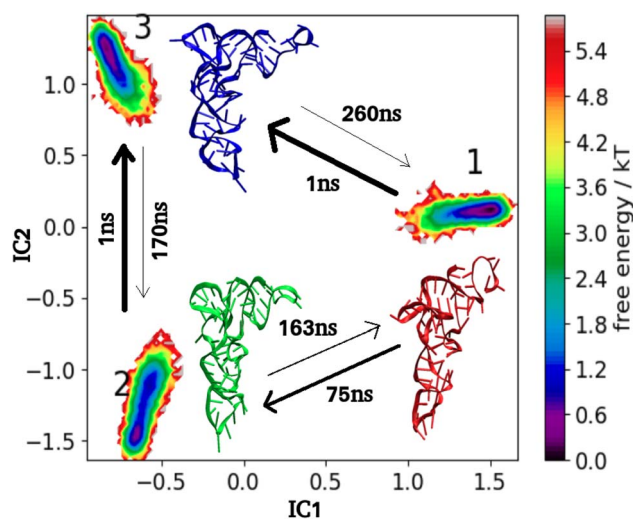


Fig. 11 MSM macrostates and MFPT rates for  $\text{tRNA}_3^{\text{Lys}}$  apo. The arrows show MFPT in ns. The metastable representative structures 1, 2 and 3 are shown in red, green and blue colors respectively.

system was observed to be fluctuating in the range of (9 Å to 19 Å) indicating instability of tertiary structure and widening of D-T $\Psi$ C loop. However, in the 1,5T system the average distance

between G19–C56 pairs was maintained in second (green) and third (blue) conformers and was observed to be closer to starting pdb structure distance  $\sim 10.7$  Å (Fig. 9). The conformers belonging to 1st MSM macrostate have shown reduction in average G19–C56 distance which is around 6.4 Å. This may indicate that the 1,5T ligand strengthens the tertiary structure across the D and T $\Psi$ C loop as compared to 1,4T ligand. The anticodon loop interaction U33–A37 was observed to be lower by  $\sim 2$  Å in all the three macrostates of the 1,4T system. Whereas, the U33–A37 interaction pair in the 1,5T system was observed to be following the canonical loop structure in conformers belonging to the first (red) and third (blue) macrostate. But the 90% of conformers from the second macrostate (green) of the 1,5T system were found to be possessing a wide range of fluctuations (in the range of 6 to 13 Å). As compared to  $\text{tRNA}_3^{\text{Lys}}$  apo system, the D loop and T $\Psi$ C loop interaction (C11–G45) in 1,4T system was observed with range of variation  $\sim 11$  Å to 14 Å (ESI Fig. S14 $\dagger$ ) in all the three MSM metastable states. On the other hand, in the case of the 1,5T system this tertiary structure interaction was seen to be significantly reduced ( $\sim 9$  Å) which is 3 Å less than the PDB structure (ESI Fig. S14 $\dagger$ ).

To quantify the effect of 1,4T and 1,5T ligands, on the overall kinetics and transition pathway of  $\text{tRNA}_3^{\text{Lys}}$ , the mean first passage time (MFPT) matrix was calculated for both the



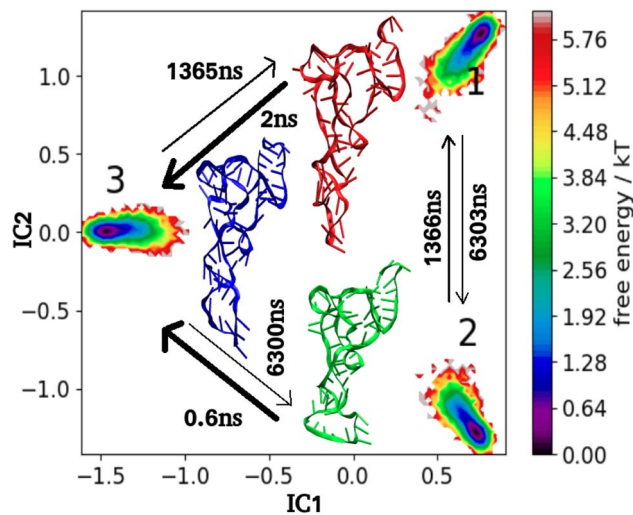


Fig. 12 MSM macrostates and MFPT rates for  $\text{tRNA}_3^{\text{Lys}}-1,4\text{T}$  system. The arrows show MFPT in ns. The metastable representative structures 1, 2 and 3 are shown in red, green and blue colors respectively.

systems. MFPT for 1,4T is shown in Fig. 12 and 1,5T is shown in Fig. 13. The change in distribution of transition of flux can be understood by transition time between any state  $i$  and state  $j$ . In the 1,4T system transitions from state-1 to state-3 (2 ns) and from state-2 to state-3 (0.6 ns) occurred most quickly. The slowest transition was seen from state 1 to state 2 (6303 ns). As compared to  $\text{tRNA}_3^{\text{Lys}}$  apo system, 1,4T system showed a significantly wide range of MFPT transitions (0.6 ns to 6303 ns). This may indicate that the 1,4T ligand leads to large free energy barriers among the macrostates of  $\text{tRNA}_3^{\text{Lys}}$ . However, in the case of a 1,5T system the range of MFPT (10 ns to 300 ns) reveals that the energy barrier between the macrostates of  $\text{tRNA}_3^{\text{Lys}}$  was reduced due to the 1,5T ligand.

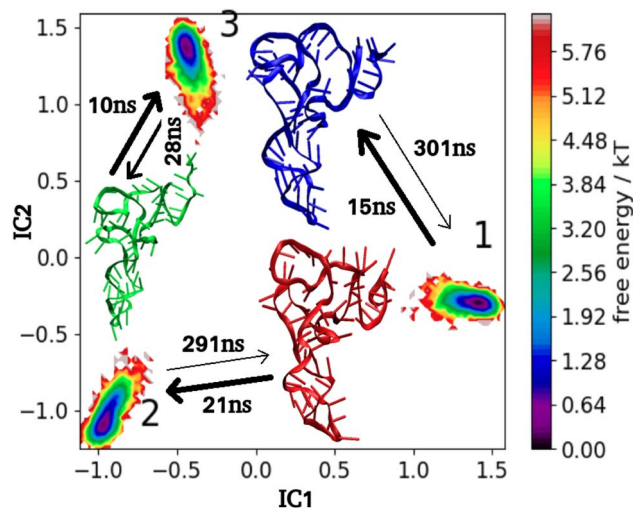


Fig. 13 MSM macrostates and MFPT rates for  $\text{tRNA}_3^{\text{Lys}}-1,5\text{T}$  system. The arrows show MFPT in ns. The metastable representative structures 1, 2 and 3 are shown in red, green and blue colors respectively.

## Discussion

Several studies pointed at the structural versatility of the  $\text{tRNA}_3^{\text{Lys}}$  and its role in the HIV-1 reverse transcription initiation complex, which might adopt different two- and three-dimensional structures.<sup>22</sup> Many experiments have been carried out on the chemical compounds that can bind to the tRNA and restrict the conformational freedom.<sup>36-41,45</sup> One of the approaches is the fragment-based strategies in the field of RNA-ligand discovery which may provide potential lead compounds for antiviral drug development.<sup>39</sup>

The *in silico* docking of two ligands (1,4T and 1,5T) against the  $\text{tRNA}_3^{\text{Lys}}$ , shows the ligand interactions with A50 and C63 residues of the  $\text{tRNA}_3^{\text{Lys}}$  respectively. The NMR spectroscopic chemical shift mapping experiments revealed two possible binding sites (one in the T $\Psi$ C-arm of  $\text{tRNA}_3^{\text{Lys}}$  and other near to the D-arm) on  $\text{tRNA}_3^{\text{Lys}}$ . The residues involved in T $\Psi$ C-arm are C49, A50, G51, G52, G53, T54, C61, C62, C63, U64 and G65. The residues involved in D-arm are U8, G10, C11, U12, C13, A14, G22, A23 and G24.<sup>40</sup> The MD simulations results also show that the stable binding interactions between the ligand and the  $\text{tRNA}_3^{\text{Lys}}$ . The experimental dissociation constant values were determined by Tisne *et al.*<sup>45</sup> for both the ligands 1,4T and 1,5T were found to be 1.8  $\mu\text{M}$  and 3.6  $\mu\text{M}$  respectively. According to these experimental results the 1,4T ligand binds 2 fold strongly to the  $\text{tRNA}_3^{\text{Lys}}$  compared to 1,5T ligand. The MMGBSA free energy calculations of simulation trajectories also show that the 1,4T ligand binds strongly to the  $\text{tRNA}_3^{\text{Lys}}$  compared to the 1,5T ligand. However, the binding location of 1,5T ligand has shown preference for T $\Psi$ C loop than D arm. It was observed from the MD simulations that the 1,4T ligand was able to bind  $\sim 2$  times more strongly than 1,5T ligand at T $\Psi$ C arm with the help of direct H-bond interactions. It was also found that this 1,4T ligand was forming  $\text{Mg}^{2+}$  ion mediated interactions as well as indirect interactions with the T $\Psi$ C arm residues. In case of  $\text{tRNA}_3^{\text{Lys}}-1,4\text{T}$  system, the water molecules are also able to interact with T $\Psi$ C arm residues, close to the 1,4T ligand binding site along with the D arm residues. The starting structures for the simulations are top scored docked poses for each system. In the MD simulations of systems, the water residues are settled in the D arm and may not allow the ligand to bind in the D arm. The D arm residues are interacting strongly with water and may not be allowing 1,4T ligands to interact directly with them. It may be a long-time scale event in the range of milliseconds to seconds to displace water molecules and interact with the D arm by the 1,4T ligand. It may require very long time-scale simulations of multiple docking poses, which is a highly expensive computational process.

Apart from binding strength of the ligand, it is also important to probe the change in structural dynamics due to the ligand. The RMSD of the anticodon loop of  $\text{tRNA}_3^{\text{Lys}}-1,5\text{T}$  system is low compared to other two systems in all three runs of simulations as shown in ESI Fig. S16.<sup>†</sup> It has been observed that the 1,5T ligand bound to the  $\text{tRNA}_3^{\text{Lys}}$  reduces the flexibility of the anticodon region as well. In order to understand the structural dynamics and kinetics of ligand bound systems, Markov state modelling was performed. Markov state modelling



of 300 ns simulations of tRNA<sub>3</sub><sup>Lys</sup> apo system was performed to understand conformational dynamics and functional kinetics. MSM analysis of tRNA<sub>3</sub><sup>Lys</sup> apo was compared independently with MSM results of tRNA<sub>3</sub><sup>Lys</sup> bound with 1,4T and 1,5T ligands. Markov state modeling of tRNA<sub>3</sub><sup>Lys</sup> apo MD trajectory data suggests three principal metastable conformations. These conformations were found to have variations mainly in the anticodon loop of the tRNA<sub>3</sub><sup>Lys</sup>. The experimental study by Benas *et al.* from where the pdb structure (1FIR) was obtained, had observed the canonical form of anticodon loop tRNA<sub>3</sub><sup>Lys</sup>. This observation was attributed to the crystal packing of two tRNAs that are closely associated head-to-tail so that the CCA end of each one interacts with the anticodon of the other one. They have predicted the formation of noncanonical anticodon loop conformation in case of free tRNA<sub>3</sub><sup>Lys</sup>.<sup>54</sup> The MSM results are also suggesting noncanonical conformation of the anticodon loop of apo-tRNA<sub>3</sub><sup>Lys</sup>. Durant *et al.* have reported the canonical conformation of the anticodon loop of tRNA<sub>3</sub><sup>Lys</sup> by incorporating hypermodifications at 34th and 37th positions.<sup>78</sup>

As revealed by NMR studies, the acceptor region was found to be flexible and D-TΨC tertiary structure was observed to be opened up. The transition between metastable states takes few nanoseconds in tRNA<sub>3</sub><sup>Lys</sup> apo and those transition rates become slower to microseconds in tRNA<sub>3</sub><sup>Lys</sup>-1,4T system, whereas the order of transitions were observed to be same (~ns) in tRNA<sub>3</sub><sup>Lys</sup>-1,5T system. MSM analysis of the tRNA<sub>3</sub><sup>Lys</sup>-1,4T system has shown that the 1,4T ligand strengthens the secondary structure in the D arm of tRNA<sub>3</sub><sup>Lys</sup>. However, the residue pairs G18-Ψ55 and G19-C56, were found to be widened up in tRNA<sub>3</sub><sup>Lys</sup>-1,4T system. On the contrary MSM analysis suggests that these tertiary structures forming residue pair distances have reduced due to 1,5T ligand resulting in closing the D-TΨC arm of tRNA<sub>3</sub><sup>Lys</sup>. These findings have been reconfirmed by MSM analysis of only the D-TΨC loop of all the three systems. MSM analysis suggests four metastable states for D-TΨC trajectories of apo tRNA<sub>3</sub><sup>Lys</sup> as well as tRNA<sub>3</sub><sup>Lys</sup>-1,4T system (ESI Fig. S17†). However, MSM of only D-TΨC loop has resulted in three macrostates of tRNA<sub>3</sub><sup>Lys</sup>-1,5T system indicating that reduction in flexibility and closing of D-TΨC loop (ESI Fig. S17-S19†).

It has been known that, in order to form the reverse transcription initiation complex, the tRNA<sub>3</sub><sup>Lys</sup> needs to undergo large-scale conformational rearrangement which requires breaking of acceptor and TΨC stem base pairing.<sup>60</sup> Although the binding studies indicate that 1,4T ligand has stronger binding properties than 1,5T ligand, the rigidity of acceptor and TΨC stem was obtained by 1,5T ligand. This property of the 1,5T ligand may prove to indirectly inhibit the initiation of reverse transcription complex. The computational insight of these ligands binding to tRNA<sub>3</sub><sup>Lys</sup> may help experimentalists to understand interesting and stable interactions, and the dynamics of ligand binding.

## Conclusions

Computational probing of tRNA<sub>3</sub><sup>Lys</sup> in the presence of ligands has been carried out using molecular docking, molecular dynamics simulations and MSM analysis. The conformational freedom of tRNA<sub>3</sub><sup>Lys</sup> structure has been analysed using MSM

analysis. The three stable states obtained for tRNA<sub>3</sub><sup>Lys</sup> with restricted flexibility of tRNA<sub>3</sub><sup>Lys</sup> in the presence of ligands. Different tertiary interactions were also probed in the analysis to understand structural effects in tRNA<sub>3</sub><sup>Lys</sup>. The conformational flexibility, tertiary interactions contribution to various structural effects and other water mediated interactions may guide the future aspects of drug design for HIV-1, where tRNA<sub>3</sub><sup>Lys</sup> acts as a primer for HIV's reverse transcription.

## Author contributions

Mallikarjunachari Uppuladinne conceptualized, conceived the research work and wrote the manuscript. Archana Achalere performed MSM analysis and wrote the manuscript. Uddhavesht Sonavane and Rajendra Joshi edited and reviewed the final manuscript.

## Conflicts of interest

The authors declare no conflict of interest.

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