




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# Effect of monovalent salt concentration and peptide secondary structure in peptide-micelle binding†

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Recently, we reported a cationic 14 residue peptide LL-14 (LKWLKLLKWLKLL) with salt-sensitive broad-spectrum antimicrobial potency. However, the mechanism of its salt (NaCl) sensitivity remained unclear. In this study, we have reported computational (~14.2  $\mu$ s of MD) and experimental (CD, fluorescence) investigations to examine the salt-sensitivity and the role of peptide secondary structure on LL-14 binding to simple membrane mimetic (SDS, DPC) systems. LL-14 was shown to adopt a random coil ( $P^C$ ) conformation in water and  $\alpha$ -helical conformation ( $P^H$ ) in the peptide:SDS micelle complex, accompanied by tryptophan burial, using both simulations and experiments. Simulations successfully deconvoluted the LL-14:micelle binding event in terms of secondary structure (random coil  $P^C$  versus helix  $P^H$ ) and gave atomic insight into the initial and final LL-14:SDS complexes. Electrostatics drove the N-terminus (L1 and K2) of LL-14 ( $P^C$  or  $P^H$ ) to bind the SDS micellar surface, initiating complex formation. LL-14 in amphipathic  $P^H$  conformation bound faster and buried deeper into the SDS micelle relative to  $P^C$ . Increasing NaCl concentration incrementally delayed LL-14:micelle binding by shielding the overall charges of the interacting partners. LL-14 binding to the SDS micelle was significantly faster relative to that of the zwitterionic DPC micelle due to electrostatic reasons. Cationic  $\alpha$ -helical amphipathic peptides (with positively charged N-terminus) with low salt-ion concentration seemed to be ideal for faster SDS binding.

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## Introduction

Antimicrobial peptides (AMPs) have drawn significant attention as potential antimicrobial and therapeutic agents.<sup>1–6</sup> The rapid rise of antibiotic resistance in microbes and the development of multi-drug resistant bacteria have resulted in a therapeutic deadlock.<sup>7–11</sup> Additionally, the slow development of new classes of antibiotics, have augmented to the complexity of the problem. This necessitates the immediate development of alternate classes of therapeutic molecules that can combat antimicrobial infections. AMPs are small polypeptides (~10–50 amino acids long) present in various kingdoms of life starting from microbes, insects, crustaceans, plants, animals and humans which are endowed with the ability to fight antimicrobial infections.<sup>12</sup> Their presence in nature for millions of years, as the first line of host-defence against microbial

infections caused by bacteria, fungi, yeast, viruses *etc.*, indicate slow development of resistance against them. Additionally, AMPs are biocompatible and upon degradation forms amino acid residues in comparison to the toxic metabolites generated by the other classes of compounds. Thus AMPs are promising as an alternative to conventional antibiotics. AMPs have variable sequences, secondary structures and modes of action (majority of AMPs are membrane-active). However, despite the huge potential of the AMPs as alternate antimicrobial therapeutics, there has only been limited commercial translation of these molecules till date. This is primarily owing to the salt sensitivity of the antimicrobial potency of the AMPs and their short serum stability due to susceptibility to protease degradation. Researchers around the world are trying to develop synthetic AMPs which can circumvent these inherent problems of natural AMPs to develop robust economically viable therapeutics. Additionally, despite computational and experimental advancement, the rational design of selective potent AMPs remains limited<sup>13</sup> due to the lack of molecular details of peptide-membrane interactions. Understanding the mechanism of interaction between AMPs and the microbial membranes in atomic terms is the key to designing new efficient membrane-active AMPs.

MD simulations partially fill up the gap by offering an atomistic understanding of the same, using simple models.<sup>14–32</sup>

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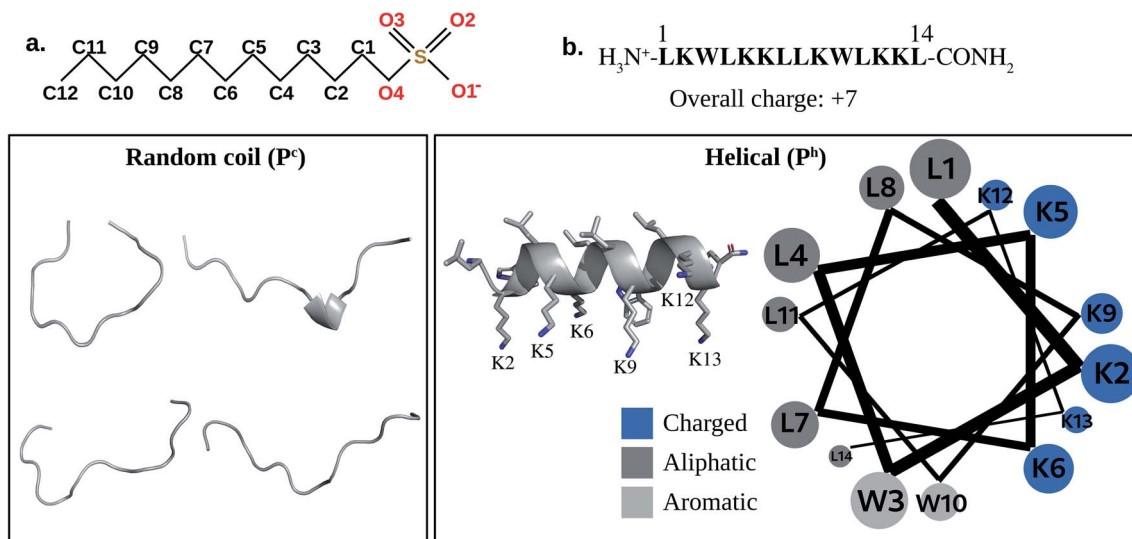


Fig. 1 Structure of (a) Sodium Dodecyl Sulphate (SDS) (b) positively charged LL-14 (–N terminal as  $\text{–NH}_3^+$  and C-terminal is capped as  $\text{–CONH}_2$ ). The random-coil ( $P^c$ ) and  $\alpha$ -helical structure of LL-14 are highlighted in the box. Helical wheel representation of the  $\alpha$ -helical LL-14 with amino acid sequence highlighting the amphipathicity.

It has been understood that electrostatics is one of the major driving forces that lead to the microbial membrane affinity of the cationic AMPs.<sup>33</sup> MD simulation studies on AMP-membrane mimetic systems offer insights into attributes like antimicrobial potency, cytotoxicity, specificity *etc.*<sup>25,34–36</sup> Experimental investigations including circular dichroism (CD), fluorescence spectroscopy, nuclear magnetic resonance (NMR), isothermal calorimetry (ITC), microscopy (FESEM, CLSM *etc.*), biochemical assays *etc.* have been used extensively in gaining insights into the mechanism of antimicrobial activity of AMPs.<sup>37</sup> Thus, the combination of computational and experimental investigations may accelerate the discovery of therapeutic AMPs.

No systematic investigation has been done to date for understanding the salt sensitivity of the AMPs. Though some salt tolerant AMPs are reported in the literature, they are mostly results of serendipitous discovery rather than of rational design. Most AMPs lose their activity in the presence of physiological salt concentrations which severely limits the commercialization of AMPs. For example, various peptides *viz.*,  $\beta$ -defensins (except for  $\beta$ -defensin III) present in humans, are prone to salt-induced inactivation.<sup>38–40</sup> In our present study, we have tried to understand the reason for the salt sensitivity of a synthetic AMP, LL-14 that was earlier reported by our group.<sup>41</sup> Cationic fourteen residue long membranolytic LL-14 (Fig. 1), was shown to have high activity (MIC<sub>99</sub>  $\sim$ 1–10  $\mu\text{M}$ ) against several strains of ESKAPE pathogens and fungal strains.<sup>41</sup> However, the activity of LL-14 was salt-sensitive against several microbes (*P. aeruginosa*, *S. aureus*, *C. neoformans*) leading to a partial loss of activity at physiological NaCl concentrations.

Combining molecular simulations and experimental studies, we have attempted to understand the salt sensitivity and the role of peptide secondary structural features on the antimicrobial potency of LL-14 using simple membrane mimetic micelles. Fluorescence and CD experiments confirmed that the free peptide was unstructured in water and attained  $\alpha$ -

helical ( $P^h$ ) conformation only after micelle binding. All-atom classical MD simulations were performed with LL-14 peptide (in two different conformations: (A) random coil or unstructured- $P^c$ , and (B) amphipathic  $\alpha$ -helical- $P^h$ ) in presence and absence of membrane mimetic system (SDS micelles: mimicking negatively charged prokaryotic membranes) in water with varying monovalent salt (NaCl) concentrations. MD simulations suggested that free LL-14 in water preferred random coil ( $P^c$ ) conformation in lines with fluorescence and CD experiments. Simulations provided atomic insight into the initial binding complex LL-14( $P^c$ ):SDS micelles. However, peptide conformational change in response to micelle binding (LL-14( $P^c$ ):SDS  $\rightarrow$  LL-14( $P^h$ ):SDS) was not observed within the time-scale of our MD simulations. To gain insight into the LL-14( $P^h$ ):SDS complex, simulations were performed to study  $\alpha$ -helical ( $P^h$ ) peptide binding to SDS and the resulting LL-14( $P^h$ ):SDS complex agreed with our conclusions from fluorescence experiment. Although, free LL-14 in the  $\alpha$ -helical ( $P^h$ ) peptide was not experimentally characterized and unrealistic, but the study of  $P^h$  binding to SDS was important for two reasons: (1) structural information about final LL-14( $P^h$ ):SDS complex was obtained, and (2) comparison of the initial binding events in terms of the peptide secondary structure ( $P^c$  binding to SDS *versus*  $P^h$  binding to SDS) was possible. The results indicated that salt (NaCl) systematically delayed the initial SDS binding event and secondary structure was crucial for initial binding ( $P^h$  binds faster than  $P^c$ ).

## Methods

### Molecular dynamics setup

**Modelling the peptides (unstructured-coil- $P^c$  and  $\alpha$ -helical- $P^h$ ).** Experimentally verified anti-microbial peptide LL-14 (Fourteen residue, sequence LKWLKLLKWLKLL, Fig. 1) was selected and modelled using PyMOL.<sup>42</sup> The N- and C-termini of



the peptide were  $-\text{NH}_3^+$  and  $-\text{CONH}_2$ , respectively. Two initial models of the peptide were prepared, a linear-chain ( $\text{P}^c$ ) and  $\alpha$ -helical ( $\text{P}^h$ ) conformation. N-terminal and the side-chain of lysine residues were positively charged, resulting in an overall +7 charge of the peptide (Fig. 1). The MD setup of free peptide in water included overlaid water-box (dimension  $\sim 10 \times 10 \times 10 \text{ nm}^3$ ) and counter-ions. The overall charge of the simulation box was zero. MD model included more than 98 000 atoms, consisting of  $\sim 32\,665$  water molecules and 7 chloride ions (Table S1†).

**Modelling of membrane-mimetic systems (SDS and DPC micelles).** Micelles were modelled using CHARM-GUI Membrane Builder.<sup>43</sup> A total of 60 molecules of SDS were used to model the micelles. The experimental value of aggregation number<sup>44</sup> was close to 60 and was adopted previously<sup>25,31,45,46</sup> for computer simulations of SDS micelle. Solvation was done by overlaying a water-box of dimension  $\sim 14 \times 14 \times 14 \text{ nm}^3$ . A total of 60  $\text{Na}^+$  ions were added for neutralizing the simulation box. Additional 138 and 278 ions ( $\text{Na}^+$  and  $\text{Cl}^-$  each) were added to model 0.5% and 1.0% w/v salt environment respectively. A total of  $\sim 260\,000$  atoms were considered for MD simulations of SDS in water. The final snap of the post-equilibrated production dynamics was selected for the peptide-micelle binding study. DPC micelle was modelled in the same way as SDS.

**Peptide-SDS/DPC micelle setup.** The final snapshot of free micelle in water was selected and the center of mass of the peptide ( $\text{P}^c$  or  $\text{P}^h$ ) was placed more than 50 Å away from the centre of mass of micelle (Fig. S1a†). The waters overlapping with the peptide were deleted and counter-ions ( $\text{Cl}^-$ ) were added to ensure charge neutralization. Simulations were carried out for different monovalent salt ( $\text{NaCl}$ , at 0.0% w/v, 0.5% w/v (85.5  $\text{mmol l}^{-1}$ ) and 1.0% w/v (171  $\text{mmol l}^{-1}$ )) concentrations with various peptide-micelle relative orientations (see “Sampling and convergence of MD simulations” section below for detail). Simulations were performed in the cubic water box of dimension  $14 \times 14 \times 14 \text{ nm}^3$  centred at the micelle. More than 260 000 atoms were considered for MD simulations (Table S1†).

**Simulation specification.** GROMACS software<sup>47</sup> was used to perform MD simulations. Peptide, micelle, water and ions were simulated using standard CHARMM36 force field.<sup>48,49</sup> TIP3P water model<sup>50</sup> was used to describe water. Covalent bonds involving hydrogen atoms were constrained using the LINCS algorithm.<sup>51,52</sup> Neutral simulation box (including peptide, micelle, ions, waters) was first energy minimized (step size of 0.01 nm and  $\sim 50\,000$  steps) by the steepest descent algorithm. After minimization, the resulting structure was subjected to 200 ps equilibration (that included molecular dynamics with 100 ps NVT followed by 100 ps NPT ensemble). Initial velocities were assigned from a Maxwell velocity distribution at 300 K. After attaining equilibration, MD was performed using NPT ensemble. During equilibration, temperature was controlled using modified Berendsen thermostat (V-rescale,<sup>53</sup> coupling with a time constant of 0.1 ps at 300 K). Simulations were done with 2 fs time-step, 300 K temperature and 1 bar pressure. Modified Berendsen thermostat<sup>53,54</sup> and Parrinello–Rahman algorithm<sup>55</sup> (for pressure coupling with the compressibility of  $4.5 \times 10^{-5} \text{ bar}^{-1}$  and a time constant of 2 ps) were used to

maintain the temperature and pressure of the simulation system. Periodic boundary conditions were used in all simulations. Long-range electrostatic interactions were treated using the particle-mesh Ewald algorithm.<sup>56</sup> Short-range van der Waals interactions were truncated at 1.2 nm. Structures were collected for analysis at an interval of every 10 ps.

**Sampling and convergence of MD simulations.** Key objective of MD simulations of a single peptide binding to a membrane mimetic system was to capture the essence of AMP: membrane binding by identifying the (1) initial and final peptide:micelle complex at atomic resolution, (2) time-scale of the binding event, and (3) binding induced change in the dynamics and structure. The convergence of the reported data must be supported by various independent MD replicas differing in the initial distance and orientations of the peptide relative to micelle and in their initial velocities. However, this approach was known to be limited by convergence issues.<sup>16,20–22,24</sup> Magainin peptides in POPC bilayer simulations were reported not to be converged up to 20 ns MD.<sup>22</sup> Simulations of the helical (ovispirin) and  $\beta$ -sheet (protegrin-1) AMPs in presence of lipid micelles showed that the final state of the complex depended upon the starting orientation of the peptide relative to the micelle.<sup>18,19,57</sup> On the contrary, cell-penetrating peptide penetratin in POPC and POPG bilayers, was successful in achieving convergence, *i.e.*, converged final state from different replicas with varying starting conformations.<sup>29</sup>

MD simulations were performed with the peptides ( $\text{P}^c$ ,  $\text{P}^h$ ) in the aqueous phase, initially placed away from the micelle surface in three different orientations relative to the micelle surface (Fig. S1a†). Each simulation model (Setup A, Setup B, and Setup C, Fig. S1a†) was subjected to multiple independent MD simulations, using randomized initial velocities (rep1, rep2..., Table S2†) at various salt ( $\text{NaCl}$ ) concentrations (Table S2†). 5 independent replicas were considered for the calculation of temporal averages. First direct contact between the LL-14 and SDS micelle was defined as the distance between the positively charged nitrogen of LL-14 and the oxygen head group of the SDS, and fixed to  $\sim 0.35 \text{ nm}$  (cut-off distance for hydrogen bonds in proteins).<sup>58</sup> Analysing multiple independent MD trajectories, we estimated the distance ( $d_{\text{COM}}$ ) between the centre of mass of LL-14 ( $\text{P}^c$  or  $\text{P}^h$ ) and the centre of mass of SDS micelle as  $3.47 \pm 0.01 \text{ nm}$ , which represented the first contact (Fig. S1b†) distance between them. It should be noted that the choice of centre-of-mass cut-off distance as 3.47 nm for describing first LL-14:SDS contact was the best estimate with a standard deviation of 0.01 nm. Simulations of free peptides ( $\text{P}^c$ ,  $\text{P}^h$ ) and the free micelles (SDS, DPC) in the water were performed. Each replica was subjected to post-equilibrated MD simulations for a minimum of 10 ns to a maximum of 6  $\mu\text{s}$  (Table S2†). Total  $\sim 14.2 \mu\text{s}$  simulation was considered for analysis which confirmed convergence of our reported data.

**Circular dichroism (CD) experiments.** CD spectroscopy was performed to gain insight into the secondary structural content of LL-14 in presence and absence of SDS micelle. Curious to look into the stability of the LL-14:micelle complex, we performed CD of the system in the presence of a chaotropic reagent Urea. CD spectra were recorded on a Jasco J-815



spectropolarimeter (Tokyo, Japan) calibrated with ( $\pm$ )10-camphorsulfonic acid for optical rotation. The spectra were recorded from 250 nm to 190 nm using 1 mm path length, in a Suprasil quartz cuvette at a scan rate of 100 nm min<sup>-1</sup>, interval of 0.5 nm and time constant of 1 s and taking an accumulation of three scans at 298 K. Observed ellipticity value ( $\theta$  in mdeg) was plotted against the wavelength (in nm). In all the cases peptide, SDS, and urea concentrations were 50  $\mu$ M, 30 mM, and 8 M respectively in the final solution. Baselines were corrected by using respective solvent systems.

**Fluorescence experiments.** The intrinsic Trp fluorescence property of LL-14 peptide was used to analyse interaction of LL-14 with SDS micelles in absence and presence of salt (150 mM NaCl, physiological concentration). Tryptophan side-chains of LL-14 were excited at 280 nm and emission was recorded from 295–540 nm in presence of aqueous solution of SDS micelles with or without 150 mM NaCl using Hitachi F-7000 FL spectrometer at 25 °C. The experiments were performed at 50  $\mu$ M LL-14 concentration and 30 mM SDS, well above the critical micelle concentration of SDS (CMC = 8.5 mM).

## Results

### Structural and dynamical properties of LL-14 and SDS micelle

MD simulations of the free peptide (P<sup>c</sup> or P<sup>h</sup>) and micelle in water provided clues about their size, shape, dynamics. Simulations of free peptide in water starting with two different conformations (*i.e.*, extended:P<sup>c</sup> or  $\alpha$ -helical:P<sup>h</sup>) showed that (1) P<sup>c</sup> remained as random coil all along the MD trajectory (up to 1  $\mu$ s, see Mov1\_Pc\_in\_Water.mpg), (2) P<sup>h</sup> (helix) conformation started melting after 800 ns (Mov2\_Ph\_in\_Water.mpg). The results showed that free peptide preferred random coil conformation in water (Fig. 1). Distance of selected atoms to the centre of mass (COM) of SDS micelle (Fig. 2a) indicated that counterion (Na<sup>+</sup>) distribution was severely affected upon peptide binding. The average distance between Na<sup>+</sup> and the COM of SDS-micelle increased upon peptide binding and was strikingly large in the case of P<sup>c</sup> binding relative to P<sup>h</sup>. The results suggested deeper binding of P<sup>h</sup> into the SDS micelle core, leading to a closer approach of the counter-ions relative to P<sup>c</sup> binding.

The radial distribution functions (RDF: oxygen atom of water molecules around the sulphur atoms of the SDS micelle) calculated from MD simulation indicated that the first coordination sphere of water was located 0.38 nm away from the sulphur atom of the micelle (Fig. 2b). Beyond the first coordination shell, the water distribution was unstructured and relatively less dense than the bulk. Peptide binding displaced water from the first coordination sphere, resulting in a dip of the RDF peak (Fig. 2b). The effective radius<sup>32,59,60</sup> or size ( $R_s$ ) of the micelle can be defined as the average of two distances, (a) the centre of mass of the micelle to the sulphur atoms of the head group (1.98 nm, Fig. 2a and b) head group sulphur to the first peak of water oxygen radial distribution function (0.38 Å, Fig. 2b) minus the radius of water (0.14 nm). Our estimated effective radius of the micelle was 2.22 nm, which was in excellent agreement with the X-ray scattering experiment<sup>60</sup> (2.23 nm). Micelle radius<sup>32,59</sup> is also defined as:

$$R_s = \sqrt{\frac{5}{3}}R_g \quad (1)$$

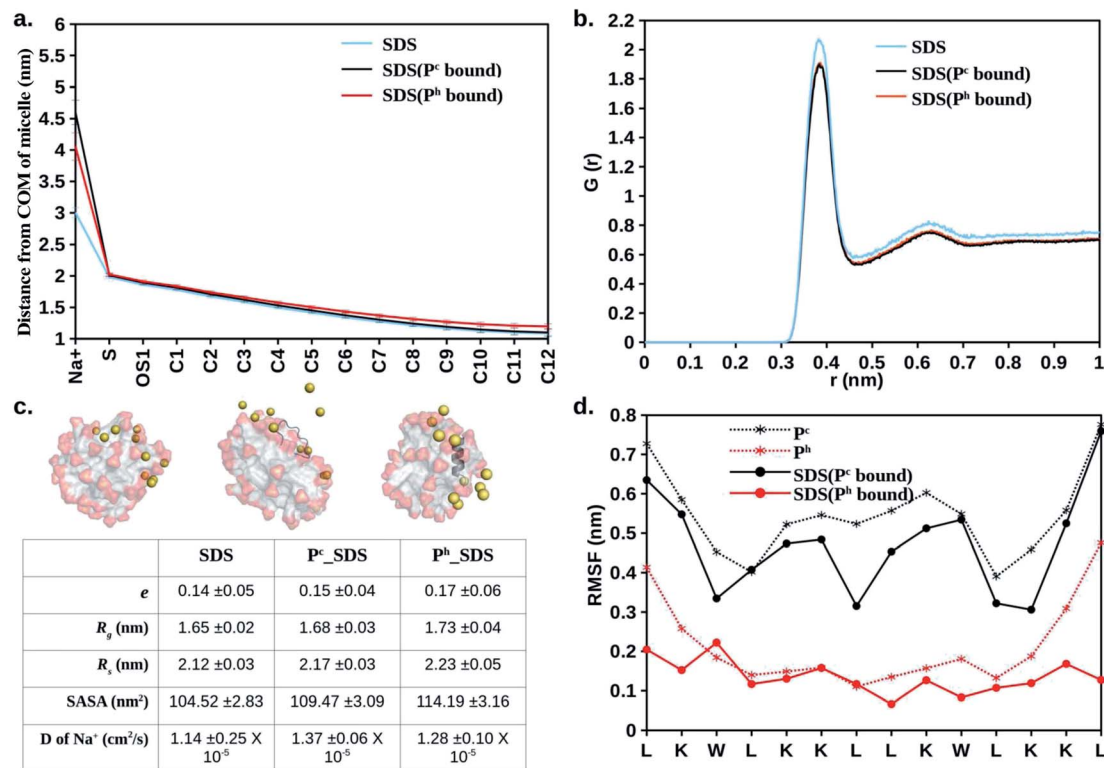
where  $R_g$  is the trajectory average radius of gyration of the micelle, computed with gromacs tool *gmx gyrate*. The MD averaged radius of gyration,  $R_g$ , of the free SDS micelle was 1.65 nm, which was very close to the previous reports (1.54–1.62 nm).<sup>25,32,61</sup> The shape of the micelle can be characterized by the eccentricity defined as,<sup>62</sup>

$$e = 1 - \frac{I_{\min}}{I_{\text{avg}}} \quad (2)$$

where  $I_{\min}$  is the smallest moment of inertia among  $x$ ,  $y$ , or  $z$  axis and  $I_{\text{avg}}$  is the average of moments of inertia along the  $x$ ,  $y$ , and  $z$  axes. A perfect sphere is characterized with  $e = 0$ . Therefore, the deviation from a spherical shape can be quantified by estimating the eccentricity; lower the value, the more spherical is the micelle. The eccentricity of SDS micelle as a function of time suggested that the micelle was not spherical (eccentricity of a sphere  $\neq 0$ ), but the shape was stable during the course of MD simulations (Fig. S2†). The MD averaged (last 20 ns of 50 ns MD) value of eccentricity and  $\frac{I_{\min}}{I_{\text{avg}}}$  (of free SDS micelle in water, Fig. 2c) were  $0.14 \pm 0.05$  and  $1.31 \pm 0.12$  respectively (close to previous studies:  $\frac{I_{\min}}{I_{\text{avg}}}$  ranging between 1.02–1.13).<sup>25,32,63</sup> The eccentricity of the micelle was found to be independent of the salt concentration (Table S3†).

Quantification of solvent accessible surface area (SASA)<sup>64</sup> also gave insight into the structural properties of the micelle. From each MD snapshot, the ions, water, peptide were first removed. Then a spherical probe molecule with the radius of 0.14 nm (mimicking water) was rolled over the surface of the micelle. The area formed by the centre of the probe quantified the total accessible surface area (SASA). The average value of the solvent-accessible surface area (SASA) of the peptide-free micelle was 104.5 nm<sup>2</sup> (Fig. 2c), which differed significantly relative to a perfect sphere of radius 2.22 nm having a SASA of 61.9 nm<sup>2</sup>. The surface roughness of the SDS micelle and its non-spherical nature resulted in nearly double the SDS surface area relative to that of a perfect sphere. SASA of the SDS increased significantly (Fig. 2c) in response to interaction with helical peptide (P<sup>h</sup>), indicating severe structural distortion and solvent exposure. MD simulations with helical (P<sup>h</sup>) and coiled (P<sup>c</sup>) peptide in SDS micelle revealed that the former penetrated the SDS micelles, whereas, the latter lied on the SDS surface (Fig. 2c). Peptide binding displaced the counter ions from the micelle surface to the bulk and increased the average distance between the counter-ions (Na<sup>+</sup>) and COM of SDS micelle (Fig. 2a). P<sup>c</sup> acted as a barrier (by lying over the SDS surface, Fig. 2c) between the counter ions (Na<sup>+</sup>) and SDS surface, increasing the bulk Na<sup>+</sup> concentration resulting in the largest average distance between the SDS and Na<sup>+</sup> ions (Fig. 2a). Penetration of the peptide (P<sup>h</sup>) into SDS micelle resulted in a lower average distance between Na<sup>+</sup> and COM of SDS (Fig. 2c and a). The flexibility of the peptide chain was estimated by calculating the root mean square fluctuation (RMSF) of the





**Fig. 2** Average structural and dynamical parameters obtained from simulation. (a) Distance of selected atoms (Fig. 1a) from the centre of mass of the SDS micelle. (b) Oxygen (H<sub>2</sub>O) to sulphur (SDS) radial distribution function. (c) Representative snapshots of free SDS, P<sup>c</sup> bound SDS, and P<sup>h</sup> bound SDS. Na<sup>+</sup> (Yellow sphere), peptide (dark grey), SDS (aliphatic: light grey, sulphate: red). Waters and hydrogens not shown for clarity. Eccentricity (*e*), radius of gyration (*R<sub>g</sub>*), micelle radius (*R<sub>s</sub>*), solvent accessible surface area (SASA), diffusion coefficient (*D*) of Na<sup>+</sup> are given. (d) Residue-wise root mean square fluctuation (RMSF) of the heavy atoms of LL-14 in the absence and presence of SDS micelle. Error bars (standard deviation) are shown either as vertical bars (a) or reported after ± (c).

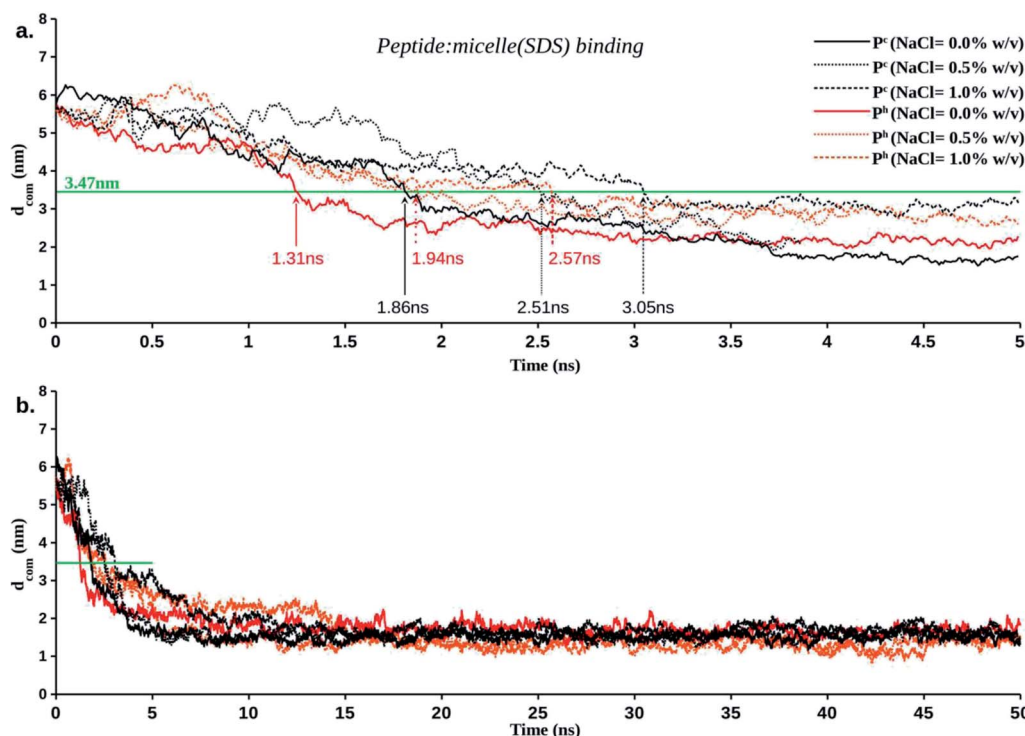
heavy-atoms of the amino acid residues in the absence and presence of micelle (Fig. 2d). RMSF plots revealed the following features: (1)  $\alpha$ -helical peptide P<sup>h</sup> had smaller RMSF relative to its random-coil-like P<sup>c</sup> analogue. Fluctuations of the amino acid residues were expected to be more pronounced in the random-coil like conformations and less pronounced in the structurally rigid  $\alpha$ -helical conformations. (2) A noticeable reduction in the flexibility (particularly in the -N and -C terminal of LL-14) was observed in the case of P<sup>h</sup> peptide binding to SDS. Micelle induced loss of LL-14 flexibility was less pronounced for P<sup>c</sup> relative to P<sup>h</sup> analogue. Thus, the secondary structure of the peptide turned out to play a key role in the structure and dynamics of LL-14:SDS complexes. Secondary structural content of the peptide was estimated (Fig. S2†) using GROMACS analysis tool (*gmx do\_dssp*) that uses DSSP software.<sup>65</sup> No significant change in the peptide conformation was observed in response to SDS binding. Peptide (P<sup>h</sup>/P<sup>c</sup>) more or less retain its conformation (helical/Coil) after SDS micelle binding during the course of our simulations.

### Salt effect in peptide-micelle (SDS) binding

The distance between the centre of mass of the peptide and the centre of mass of the micelle (*d<sub>COM</sub>*) was plotted as a function of time (Fig. 3). Distance *d<sub>COM</sub>* ~3.47 nm indicated first direct contact (see Method) between peptide and micelle. The results

revealed several remarkable features (Fig. 3). Firstly, the temporal decay of *d<sub>COM</sub>* was much faster for  $\alpha$ -helical (P<sup>h</sup>) LL-14 relative to its unstructured random-coil-like (P<sup>c</sup>) conformation. This highlighted the importance of secondary structure in peptide-micelle binding kinetics. Secondly, the presence of monovalent salt drastically delayed the peptide-micelle interaction. The salt-induced time delay was independent of the structure of the peptide (helical, P<sup>h</sup> or random-coil P<sup>c</sup>), indicating a general feature of the salt effect in peptide-micelle binding. An increase in salt concentration (0.0% → 0.5% → 1.0% w/v) systematically delayed the peptide-micelle binding. Thirdly, LL-14 bound to SDS micelle in a rapid (initial contact within 4 ns of MD) and irreversible manner. Fourthly, the distance between the centre of mass of the peptide and micelle was more or less constant after 10 ns, indicating structural convergence of the peptide-micelle complex. The robustness of the above features was confirmed from multiple trajectories (Table S2†), that differed in their initial velocities and relative positions. It should be noted that a single peptide in an explicit cubic water box of dimension ~14 nm referred to a concentration of ~605  $\mu$ M in the MD simulation model. Thus, the concentration of peptide in MD was much larger than the MIC against several ESKAPE pathogens (1–10 M) and in CD experiments (100  $\mu$ M).<sup>41</sup> High concentration of peptide resulted in faster binding of peptide to micelle (within 5 ns for various MD





**Fig. 3** Temporal averaged distance between the centre of mass of a LL-14 and the centre of mass of the SDS micelle ( $d_{COM}$ ) as a function of time. (a) Initial binding event. Peptide: micelle initial contact distance is represented as green line at  $d_{COM} \sim 3.47$  nm. (b) Plateau of  $d_{COM}$  vs. time plot (after 10 ns) indicate structural convergence of the final peptide: micelle complex. Error bars are not shown for clarity (see Fig. S3† for detail).

models). Using a larger water-box could be an alternative for lowering the peptide concentration, but would incur huge computational cost, rendering it impractical. The systematic salt-induced delay in the interaction time was a robust feature observed from several MD studies (confirmed from multiple simulations of various MD models, see method). Thus it can be claimed that the general principle of salt-induced delay was captured by our simulation setup with moderate computational cost.

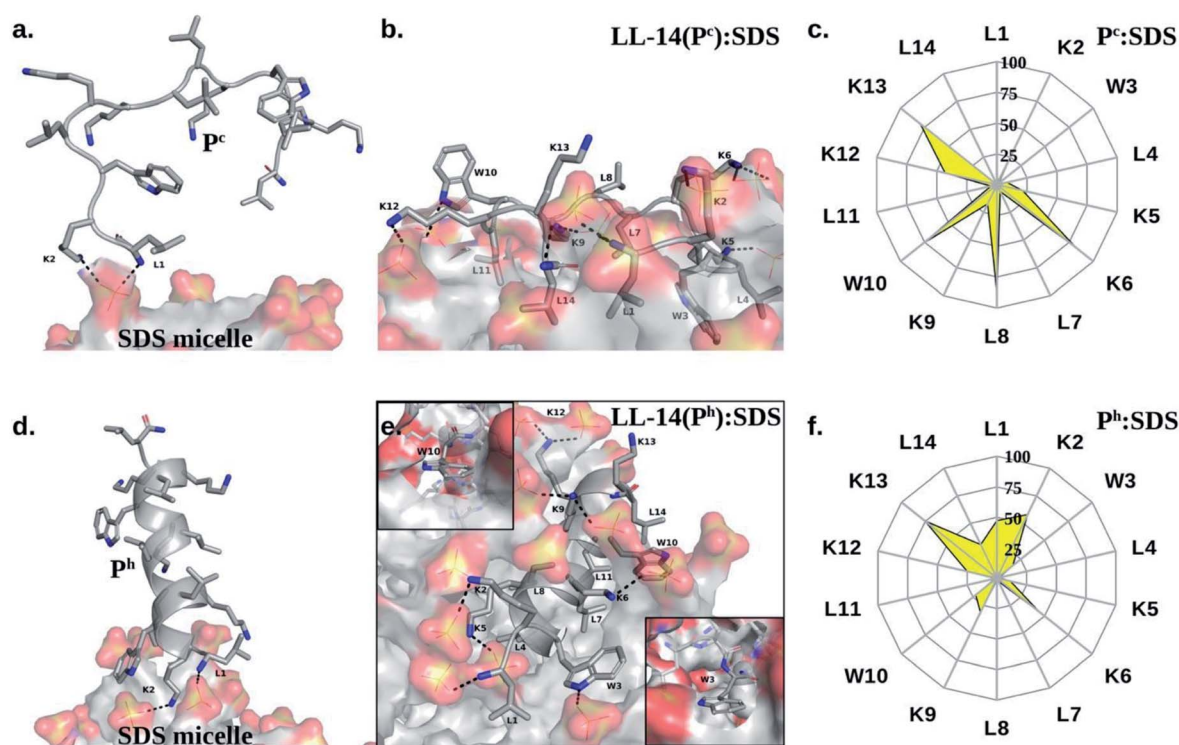
### Structural insight into the sequential peptide-micelle binding event from MD simulations

Several prominent structural features were observed from MD simulations. Firstly, initial LL-14:SDS contact was formed by involving the positively charged N-terminal of the peptide and the negatively charged micelle surface (Fig. 4a and d). We considered initial contact when the distance between the positively charged nitrogen of the peptide and the oxygen head group of the micelle was 0.35 nm. Secondly, in the finally converged LL-14:SDS complex, the helical peptide ( $P^h$ ) was engulfed by the micelle (except at the terminals), whereas the random-coil-like peptide ( $P^c$ ) lied over the surface of the micelle (Fig. 4b and e). We found that the peptide-micelle binding followed a sequential order. First, electrostatic interaction between negatively charged SDS surface with the positive charge of peptide brought them close together. Distance plot between positively charged nitrogen's of the peptide and the centre of mass of SDS (Fig. S4†) during the MD trajectories, including

visual examination of MD snaps confirmed that the N-terminal, and the side-chains of K2 of LL-14 played a key role in establishing the initial contact (Fig. 4a and d). Next, additional favourable electrostatic interactions (involving other positively charged lysine's and phosphate head group of SDS) and hydrophobic interactions (involving aliphatic/aromatic side chains of the peptide and aliphatic chain of micelle) led to a stable peptide-SDS complex (Fig. 4b and e).

The peptide backbone of  $P^c$  satisfied its hydrogen bonding requirement by forming an H-bond with waters or sulphates of the SDS micelle or both (Fig. S5†). Out of the two tryptophan's (W3 and W10), only one tryptophan (W3) of  $P^c$  was buried in the hydrophobic aliphatic pocket of SDS (Fig. 4b and c). Side-chain of four residues (K6, L8, W10, and K13) of  $P^c$  showed more than 50% solvent exposure (Fig. 4b and c) while being complexed with micelle. In  $P^h$ , the backbone was involved in secondary structure formation (*i.e.*, helix), thus, unavailable for interaction with water or SDS.  $P^h$  was mostly buried inside the micelle, except for residues K2, K13 (Fig. 4e and f). Note, both the tryptophans (W3, W10) were buried into the hydrophobic core of the SDS micelle in the  $P^h$ :SDS complex (Fig. 4e and f). Solvent exposure of  $P^c$  in  $P^c$ :micelle complex was much larger relative to  $P^h$  in  $P^h$ :micelle complex (Fig. 4c and f). We estimated the hydration of the hydrophobic part of the SDS micelle in  $P^c$ :micelle and  $P^h$ :micelle complexes. Hydration of the hydrophobic parts ( $P^c$ :micelle =  $87.1 \pm 8.6$ ,  $P^h$ :micelle =  $98.6 \pm 8.7$ ) was estimated by averaging the number of water molecules (from the last 20 ns of 50 ns of MD trajectory) within the commonly





**Fig. 4** Structures from the peptide: micelle (SDS) binding event, with distinct secondary structures of the peptide ( $P^c$  or  $P^h$ ) as initial condition. (a and d): peptide: SDS-micelle initial contact. Micelle surface forms direct interactions with the  $\text{NH}_3^+$  of the N-terminal (L1) and side-chain of K2 of the LL-14 peptide both in the (a)  $P^c$  and (d)  $P^h$  conformations. (b and e) Final structure of the peptide: micelle complex. Interaction network is shown as dotted line for (b)  $P^c$ :micelle and (e)  $P^h$ :micelle complex respectively. For clarity, van der Waals radii of the micelle was reduced by 50% in (e). The local environment around W3 and W10 are explicitly shown without van der Waals reduction in the in-set (e). (c and f) Residue-wise solvent exposure (in percentage) of LL-14 in the final complex was estimated as: (SASA of amino acid residue in SDS:LL-14 complex/SASA of the amino acid residue in LL-14)  $\times$  100. Trajectory averaged percent solvent exposure (yellow) is shown in the net-plot with contours of constant percent exposure. Waters and hydrogens not shown for clarity in the structures.

used cut-off of 0.35 nm (ref. 25) of the non-polar moiety (C1–C12 of Fig. 1a) of each SDS molecule. The higher water exposure of the lipid portion of the micelle in the  $P^h$ :micelle relative to  $P^c$ :micelle complex, indicated higher hydrophobic destabilization of the micelle in the former.

Notably, irrespective of the secondary structural nature of the peptide ( $P^c$  or  $P^h$ ), electrostatic interaction between the positively charged N-terminal of the peptide and the negatively charged anionic head group of the micelle lipid was the driving force for the initial binding (Fig. 4a and d). Structural convergence in terms of the observed atomic interaction network in the final complex (Fig. 4b and e) was confirmed from multiple independent MD simulations (Fig. S1a and Table S2<sup>†</sup>). Simulations with various peptide ( $P^h$  or  $P^c$ ) orientation relative to the micelle surface were found to be converged to the same final structure of the peptide:micelle complex (Fig. 4b and e) and independent of the salt concentration (Fig. S6<sup>†</sup>). Reasonable sampling ( $>14 \mu\text{s}$  of MD) and converged structure of the final peptide:micelle complex indicated that the reported MD structures of the peptide:micelle complex represented the true minima on the free energy hypersurface.

MD simulation with a neutral DPC micelle (zwitterionic lipid that mimics mammalian membrane) showed that  $P^c$  was able to

bind DPC micelle surface after 870 ns of dynamics (Mov3\_Pc\_DPC.mpg), the delay being attributed to the obvious electrostatic reasons. The results highlighted the importance of favourable electrostatic interactions in facilitating peptide:micelle interaction as reported in earlier studies.<sup>33</sup> Though LL-14 peptide was found to be cytotoxic,<sup>41</sup> MD studies suggested that the interaction of LL-14 with the mammalian membranes was much slower compared to that with the microbial membranes. A detailed structural analysis of the peptide:DPC binding is outside the scope of this paper, and will be reported elsewhere.

The driving force for the initial LL-14:SDS binding event was investigated by plotting the electrostatic (Coulomb) and Lennard-Jones (van der Waals is the attractive term) energy as a function of time (Fig. 5). The results showed that electrostatic energy profile, decayed faster for  $P^h$  relative to  $P^c$ , whereas there was a little difference in the decay of Lennard-Jones energy for  $P^h$  and  $P^c$  during the initial binding event (*i.e.*, within 2 ns of MD, see Fig. 2a), highlighting the electrostatic origin for the faster binding of  $P^h$  relative to  $P^c$  as observed in Fig. 3. Though both the “electrostatic” and “van der Waals” energies were favourable (negative sign, Fig. 5), the former was the primary contributor to the overall interaction potential energy. The



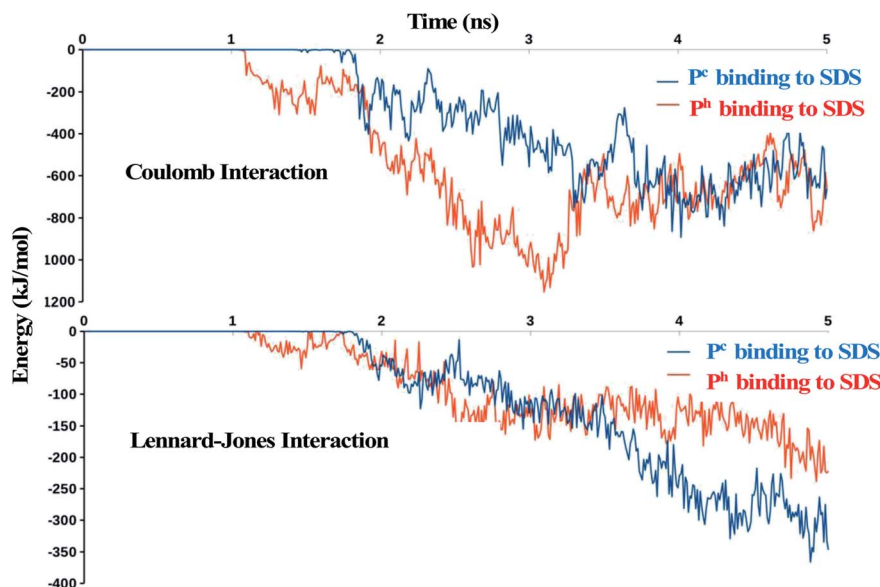


Fig. 5 Electrostatic and Lennard-Jones interaction energy as a function of time during the initial stage of peptide: micelle (SDS) binding. The peptides in distinctly different conformations ( $P^c$  or  $P^h$ ) were subjected to MD simulation. Long-range cut-off of 1.4 nm was used to compute the energies.

favourable electrostatic term was attributed to the interactions between the positively charged side-chains of peptide and negatively charged SDS surface whereas, the favourable Lennard-Jones potential energy indicated favourable van der Waals interactions involving placement of hydrophobic residues of the peptide (Trp, Leu, aliphatic chain of Lys) into the aliphatic core of SDS micelle.

### Secondary structural insight from CD experiments

CD experiments (Fig. 6a) confirmed that free LL-14 in water adopted a random coil like ( $P^c$ ) conformation with a negative peak at 200 nm. The result directly corroborated with the helix

melting of free LL-14 observed in the MD simulation (Mov2\_Ph\_in\_Water.mpg). However, in the presence of SDS micelles, a positive peak at 195 nm and two negative peaks at 208 nm and 222 nm were observed (characteristic features of significant helical content), which indicated helical conformation of LL-14 in the SDS micelle bound state. Thus, CD experiments suggested SDS micelle induced conformational change of LL-14 peptide ( $P^c$  free in water  $\rightarrow$   $P^h$  bound to SDS). Furthermore, the stability of the LL-14:SDS complex was investigated by CD experiments in the presence of chaotropic reagent urea. Experiments were performed either by adding urea into the LL-14:SDS complex, or by adding LL-14 into the solution of urea

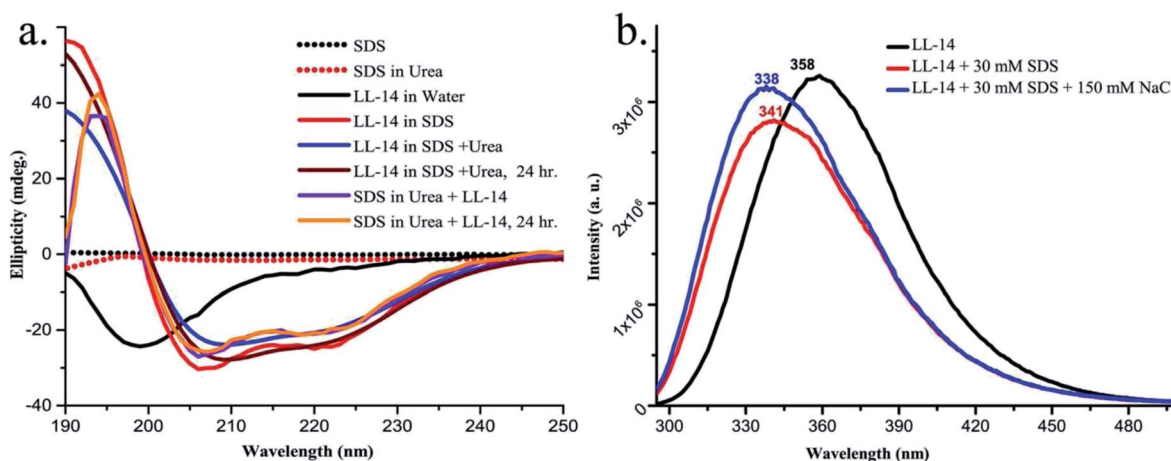


Fig. 6 (a) CD spectra of free SDS micelle (as control, broken line) and LL-14 in various conditions (presence or absence of SDS micelle and urea, solid lines). CD spectra indicate characteristic SDS induced LL-14 conformational change from random coil to helix ( $P^c$ : Solid black  $\rightarrow$   $P^h$ : Solid colours) even in the presence of urea. (b) Tryptophan Fluorescence emission spectra of free LL-14 (black), LL-14 in presence of SDS (red), and LL-14 in presence of SDS and salt (red). Blue shift in LL-14 emission observed upon addition of SDS micelle in presence or absence of salt (NaCl).



and SDS micelle. Results confirmed that LL-14 in complex with SDS retained its helical conformation ( $P^h$  bound to SDS) even in presence of urea and was independent of the order of addition of urea in the experiments.

### Fluorescence studies and tryptophan burial in the final peptide:micelle complex

LL-14 binding to SDS micelle was investigated by monitoring the intrinsic fluorescence emission maxima of the tryptophan residues of LL14 in the presence and absence of SDS micelle (Fig. 6b). The free LL-14 peptide in water, upon being excited at  $\lambda_{\text{max,ex}} = 280$  nm showed an excitation emission with  $\lambda_{\text{max,em}}$  at 358 nm. Upon addition of 30 mM of SDS micellar solution (above CMC = 8.5 mM) to the peptide,  $\lambda_{\text{max,em}}$  of LL-14 shifted to 341 nm. This large blue shift of  $\lambda_{\text{max,em}}$  of about 17 nm in the SDS micellar system indicated the change in the microenvironment of tryptophan residues. In other words, the blue shift in the presence of SDS micelles suggested the peptide-micelle binding with the insertion of the tryptophan side chains into the hydrophobic core of the micelle. A blue shift (20 nm,  $\lambda_{\text{max,em}}$  at 338 nm) was also observed in presence of 150 mM NaCl, thus indicating a salt-independent feature of LL-14 binding to SDS micelle. The more or less similar value of the blue shift (20 nm/17 nm in presence/absence of NaCl) indicated that tryptophan burial into the SDS hydrophobic core was independent of salt. Presence of a single peak at the shifted position (338 nm/341 nm in presence/absence of NaCl) indicated that both the tryptophan residues (W3 and W10) of LL-14 were in the similar environment in the micelle bound state. MD structures of the converged LL-14( $P^h$ ):SDS micellar complex, also showed burial of both the tryptophan residues (W3 and W10) in the SDS micelles thus corroborating the experimental observations. Thus, the fluorescence data indirectly proved that the structure of the final peptide:micelle complex was salt-independent and similar to the LL-14( $P^h$ ):SDS micellar complex observed from MD simulations.

## Discussion

The mechanism of membrane-active AMP induced cell-death is a multi-step process that entails AMP binding to membrane, membrane induced secondary structure formation of the peptide (coil  $\rightarrow$  helix is most common), local peptide aggregation at the membrane and subsequent membrane destabilization leading to cell lysis.<sup>17</sup> Unfortunately, peptide aggregation and cell lysis are still non-tractable by current computational power due to inherent complexity and large time-scale associated with the event. However, all-atom classical MD simulations are useful for investigating the AMP binding to simple membrane mimetic systems and can shed light on the general aspects (*viz.*, structure, dynamics and kinetics of peptide:micelle binding) of AMP induced cell death. These information would be useful in designing peptide antimicrobials with improved activity and selectivity against pathogenic microbes.

Broad-spectrum cationic antimicrobial peptide<sup>41</sup> LL-14 (Fig. 1) was selected and its binding to membrane mimetic

systems was investigated in atomic details using MD simulations and experiments (CD, fluorescence). Simulations of free LL-14 in water showed that the helix ( $P^h$ ) conformation melted after  $\sim 800$  ns of MD, and LL-14 preferred random-coil ( $P^c$ ) conformation. CD experiments also revealed that free LL-14 adopted a random coil ( $P^c$ ) structure in water. Fast LL-14:SDS binding (within 4 ns of MD) (Fig. 3) relative to slow helix melting ( $\sim 800$  ns), allowed us to study LL-14:SDS binding event considering the secondary structure of the peptide (*i.e.*,  $P^c$  versus  $P^h$  binding to SDS, despite the fact that  $P^h$  was not the preferred conformation of the free peptide in water). The results suggested that the secondary structure of the peptide played a crucial role in SDS micelle binding kinetics (Fig. 3) and insertion (Fig. 4). Helical ( $P^h$ ) LL-14 established faster initial contact with SDS and was deeply inserted into the micelle in the final stage of binding relative to its coil analogue ( $P^c$ ). Thus, the secondary structure of peptide, was crucial for the fine-tuning of the speed of the initial peptide:micelle binding event. Electrostatic interaction between the negatively charged head-group of SDS and positively charged nitrogen's of peptide (particularly N-terminal and side-chain of K2) was the key factor for interaction. This was followed by further hydrophobic stabilization (placement of aliphatic/aromatic side-chains of the amino acids into the aliphatic interior of the micelle) and additional electrostatic interactions. The settled peptide ( $P^h$ ) was found to be under the micelle surface ("hot-dog" like geometry), while  $P^c$  was found to be lying on the micelle surface. MD structure of the final converged LL-14:SDS complex showed burial of both the tryptophan residues (W3, W10) of LL-14 into the hydrophobic core of the SDS micelles in the case of the  $P^h$ -SDS complex (Fig. 4e), contrary to that of the  $P^c$ -micelle complex, where only one of the tryptophan side chains was buried (Fig. 4b).

CD experiments suggested SDS induced helical conformation of the peptide in the final LL-14:SDS complex (Fig. 6). The peptide retained its helical conformation ( $P^h$ ) in the LL-14:SDS complex even after 24 hours of urea exposure, suggesting the stability of the complex. Furthermore, LL-14 addition to urea containing SDS micellar solution also supported  $P^h$  conformation of LL-14. Thus, SDS induced LL-14 conformational change ( $P^c \rightarrow P^h$ ) was a consistent and robust feature. We observed a blue shift in the emission maxima of the tryptophan fluorescence (Fig. 6), in the presence of the SDS micellar system, suggesting the burial of the side chains of tryptophan in the hydrophobic inner core of the micelles. The observation of distinct single peak indicated similar environment around both the Tryptophan's (W3 and W10) of LL-14 upon micelle binding. Thus, fluorescence experiments corroborated with the observed final MD structures of LL-14( $P^h$ ):SDS complex (Fig. 4e).

It is worth mentioning that, we did not observe  $P^c \rightarrow P^h$  conformational change in response to micelle binding during our finite simulation time-scale (even extending the simulation up to 6  $\mu$ s, see Mov4\_Pc\_SDS.mpg). Thus, the micelle induced LL-14 folding was not computationally tractable and certainly limited by the time scale and simplified description of the system. Nonetheless, MD simulation of  $P^c$  binding to SDS provided atomic insight into the thermodynamically hidden



(experimentally unresolved) high energy state, *i.e.*, LL-14(P<sup>c</sup>):SDS complex. Moreover, MD simulation of P<sup>h</sup> binding to SDS was successful in providing atomic model (*i.e.*, LL-14(P<sup>h</sup>):SDS complex, Fig. 4e) for the experimentally characterized state (Fig. 6). A large activation barrier between the high energy (LL-14(P<sup>c</sup>):SDS complex) and low energy (*i.e.*, LL-14(P<sup>h</sup>):SDS complex) state might hinder the observation of micelle induced LL-14 conformational change (P<sup>c</sup> → P<sup>h</sup>) during the course of MD simulations. Interestingly, MD simulations not only provided atomic insight into the various stages (*viz.*, initial (Fig. 4b), final (Fig. 4e) complex) of peptide binding and/insertion process but was also useful in understanding the kinetic aspects (Fig. 3). An increase in salt (NaCl) concentration shielded the charges of both the SDS micelle and peptide resulting in a delayed binding process (Fig. 3). The final structure of the peptide-micelle complex was well converged and independent of the salt concentration. Thus, although the salt delayed the forward-rate ( $k_{on}$ ) of the LL-14:SDS initial binding event, the atomic interaction network observed in the final peptide-micelle complex was independent of the salt concentration. Fluorescence experiments showed a similar amount of blue shift in the  $\lambda_{max}$  of the Trp fluorescence emission in the presence of SDS both in the absence and presence of salt, suggesting similar amount of Trp side chain burial in the LL14:SDS complex. This was completely in alignment with the findings from the simulation (Fig. 4e). Though, the difference in the interaction times of the peptide with the SDS micelles was small (~1 ns), it should be remembered that the simulations were performed at a much higher concentration (~6 × MIC) due to the constraint in the computational expenditure. In spite of the absolute value of the delay time, our results definitely showed the trend of the effect of salt ions. It may be hypothesized that slowing down of the initial peptide-micelle binding process might facilitate microbial progeny growth, leading to the impaired activity of LL-14 in the presence of salt. It should be noted, that our argument did not underestimate the crucial effect of hydrophobic stabilization in peptide-micelle/membrane interactions. It was an important component of antimicrobial effect, but came into play only after the initial peptide-micelle binding event. The present results also emphasized the importance of terminals of the peptide in rendering the activity of the AMPs, consistent with the previous experimental study.<sup>33</sup> We had demonstrated earlier that the conversion of the free C-terminal to the amidated analogue in a small synthetic AMP P4, improved the antimicrobial potency many folds. Thus,  $\alpha$ -helical conformation of any cationic peptide and lower salt (NaCl) concentration would be ideal conditions for fast peptide-micelle (SDS) binding.

MD studies based on simple micelle models certainly have limitations in addressing issues like the effect of various other salts (monovalent, divalent *etc.*), importance of peptide-lipid ratios, role of proteins embedded in the bacterial membrane, inadequate representation of microbial lipids *etc.* Nevertheless, systematic MD simulations emphasized the importance of (1) the secondary structure and the N-terminal of LL-14 for establishing fast initial contact with SDS, and (2) salt in delaying the peptide-micelle association.

Thus the above discussions indicated that: (i) peptide-SDS micelle binding was sequential and electrostatics drove the initial binding followed by van der Waals stabilization. (ii) Presence of salt (NaCl) delayed the initial peptide:micelle contact by shielding the overall charge of the interacting partners. (iii) Peptide in  $\alpha$ -helical conformation (P<sup>h</sup>) bound faster to the SDS micelle than its random-coil analogue (P<sup>c</sup>). Thus, peptide:micelle binding kinetics depended on both the salt concentration and the nature of the peptide (P<sup>c</sup> or P<sup>h</sup>). (iv) The converged MD structure of the final LL-14(P<sup>h</sup>):SDS complex (Fig. 4e) corroborated with the fluorescence experiments. The computational analysis complemented our experimental studies by providing structural and dynamical insight into the experimentally unresolved/unrealized high-energy states, like, free P<sup>h</sup> in water and final P<sup>c</sup>:SDS complex (Fig. 4b). LL-14 binding to DPC micelle (zwitterionic eukaryotic membrane mimic) was observed to be much slower relative to SDS micelle binding.

The CD and fluorescence experiments were performed in the SDS micellar environment (above CMC concentration) at pH = 7. SDS is highly acidic with a  $pK_a = -1.5$ .<sup>66</sup> Thus, it is unlikely to alter the protonation state of the SDS molecule (negatively charged) around pH = 7. Thus, MD simulations were performed for charged SDS micelle at the same pH. However, the critical micelle concentration (CMC) of the SDS depends on the pH (ref. 67) and the salt concentration.<sup>68</sup> Self-assembly behaviour (micelle-bilayer) of lauric acid (fatty acid) was investigated extensively by Morrow *et al.*, employing continuous constant pH molecular dynamics (CpHMD) simulations.<sup>69,70</sup> It was shown that lauric acid self-assembly was pH dependent (low pH favoured bilayer and high pH favoured micelle). Although, lauric acid is different from SDS as the former contain -COOH group whereas the later contain -SO<sub>4</sub> group, but the results were of general importance to the surfactant assembly. Vila Viçosa *et al.* investigated<sup>71</sup> the titration curve of oleic acid (carboxylic acid with 18 carbons) bilayer using extended CpHMD method (CpHMD-L). The computed titration curves were in good agreement with the experimental data. The adopted methodology (CpHMD-L) is very promising for studying more realistic pH sensitive lipid bilayers. Similar to SDS, the 14-mer AMP (LKWLKLLKWLKLL) considered in this study contained titratable lysine side-chains and amino N-terminal. The  $pK_a$  of the side-chain of lysine (amino acid) and N-terminal of Leucine (amino acid) are 10.53 and 9.6 respectively.<sup>72</sup> At neutral pH = 7 the titratable residues (of the free peptide) were expected to be positively charged, thus, simulations were performed with +7 charged 14-mer peptide. However, large shifts in  $pK_a$  values (~5 units) of lysine residues buried inside a protein interior was reported by Isom *et al.*<sup>73</sup> Titration events were shown to alter the AMP:micelle binding event.<sup>74-79</sup> In the present study however, salt-bridge interaction was reported to be formed involving negatively charged SDS surface and NH<sub>3</sub><sup>+</sup> tips of peptide. The ion-pair (SO<sub>4</sub><sup>-</sup>:NH<sub>3</sub><sup>+</sup>) might alter the charge state to form (SO<sub>4</sub>H:NH<sub>2</sub>) in the anhydrous hydrophobic core, investigating which is beyond the scope of the present study. Thus, continuous constant pH molecular dynamics simulations might be



useful for understanding the role of pH in the mechanism of peptide:micelle binding.

## Conclusion

In this paper we have successfully established the link between the salt concentration (as well as secondary-structure of peptide) and the speed of peptide:micelle binding, and achieved structural convergence of the final peptide:micelle complex that was complemented by experimental observations. Salt (NaCl) induced slowing down of the kinetics of the peptide micelle interaction might be one of the possible reasons for the salt sensitivity of the AMPs in general. A detailed thermodynamic analysis (binding affinities at various salt concentrations) of peptide:micelle interaction will be reported in a separate study. We expect that the observations reported here will be relevant for other cationic peptides and micelle mimetic systems in general.

## Abbreviations

AMP	Anti-microbial peptide
CD	Circular dichroism
CMC	Critical micelle concentration
DPC	Dodecylphosphocholine
MD	Molecular dynamics
NMR	Nuclear magnetic resonance
PDB	Protein data bank
PME	Particle mesh Ewald
RMSD	Root mean square deviation
RMSF	Root mean square fluctuation
SDS	Sodium dodecyl sulphate
Trp	Tryptophan

## Conflicts of interest

The authors declare no conflict of interest.

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