Surface active ionic liquid induced conformational transition in aqueous medium of hemoglobin†

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The interaction of human hemoglobin (Hb) with surface active ionic liquids (SAILs), 1-dodecyl-3-methylimidazolium chloride [C12mim][Cl] and 1-hexyl-3-methylimidazolium dodecylsulfate [C12mim][SDS], has been studied in aqueous medium through various techniques like surface tension, UV-vis spectroscopy, fluorescence spectroscopy, isothermal titration calorimetry (ITC) and dynamic light scattering (DLS). The interactional behavior of SAILs toward Hb at the air/solution interface is investigated and various interfacial and thermodynamic parameters have also been calculated. The conformational changes in Hb upon interacting with SAILs have been illustrated from UV-visible measurements in combination with fluorescence spectroscopy. These results indicate that at lower concentration the [C12mim][SDS] monomer forms stronger Hb–[C12mim][SDS] monomer complexes as compared to [C12mim][Cl], whereas at higher concentration [C12mim][Cl] denatures Hb more and induces the release of heme from the hydrophobic pocket of Hb. The enthalpy changes were also investigated by using isothermal titration calorimetry (ITC). The dynamic light scattering (DLS) measurements revealed the effect of SAILs on the hydrodynamic diameter (Dh) of Hb.

1. Introduction

Hemoglobin (Hb) the main component of red blood cells is an iron containing oxygen transporter and fundamental part of vertebrate erythrocytes. It plays an important role in transporting oxygen from lungs to different tissues.1–3 It transports H+, HCO3− (bicarbonate) 2,3-bisphosphoglycerate and carbon dioxide, and maintains the pH of blood.4–6 Hemoglobin, with a molecular weight 67 kDa, has a tetrameric structure consisting of four heme prosthetic groups and four polypeptide chains (two α chains and two β chains), which are held together by a number of noncovalent interactions such as hydrogen bonds, van der Waals interactions, electrostatic interactions, and hydrophilic/hydrophobic forces.6–7 The α subunit contains 141 amino acid residues and the β subunit contains 146 amino acid residues. There are two major conformations of Hb (R and T); oxygen binds to both states but has relatively higher affinity for the R state.8 The deformation of Hb is associated with many diseases like leukemia, anemia, porphyria, sickle cell anemia and hemolyis.9–10

The field of protein–surfactant interactions is of preponderant importance and widely studied using many techniques owing to their applications in biological, medicine, pharmaceutica, cosmetics, emulsifiers, paints and coatings.11–14 Proteins are frequently utilized in conjugation with various surfactants to control and improve the interfacial tension as well as bulk properties of proteins. In addition surfactants have also been found to induce structural changes in thermodynamically stable conformations of proteins which in turn affect functional activities of proteins. Thus such conformational alterations in proteins due to clashes between surfactants and proteins make the in vitro protein–surfactant studies relevant. There are several reports in literature where surfactants seems to improve interfacial and bulk properties of proteins and also induce conformational alterations of proteins in terms of folding and unfolding, fibril formation of proteins, self aggregation of proteins and affects the biological activity of proteins.15–17 Sarrion et al. have studied the binding affinity of dimeric surfactants to calf thymus DNA and demonstrated the effect of spacer length of dimeric surfactants on DNA–surfactants interactions employing spectroscopy, (UV-visible and fluorescence) atomic force microscopy and various other techniques.18 The changes in secondary structure of bovine liver catalase in the presence of gemini surfactants and unfolding of protein have also been investigated by Akram et al.19 Gebicka and Banasiak have explored the conformational changes in methemoglobin upon binding with anionic surfactants sodium dodecyl sulphate (SDS) and sodium bis(2-ethylhexyl) sulfosuccinate (AOT) and reported the formation of pentacoordinated species in micellar media.20 Cationic surfactant hexadecyltrimethylammonium bromide (HTAB) has been reported to induce alterations in terms of unfolding in Hb using spectroscopy measurements.21 The hydrophobic adsorption

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mechanism of two different surfactants, hydrogenated (sodium octanoate (SO)) and fluorinated (sodium perfluorooctanoate (SPFO)) on hemoglobin and number of adsorption sites on Hb have been studied by Prieto et al.\textsuperscript{22}

Over the past few years, the field of surfactant chemistry has anticipated the replacement of conventional surfactants with a new class of surfactants named as surface active ionic liquids (SAILs) due to their tunable physicochemical properties and innate surface activity.\textsuperscript{25–28} A variety of SAILs have been investigated in the literature which are mainly based on imidazolium, pyridinium, morpholinium and pyrrolidinium cations but the imidazolium based SAILs are of specific interest due to their better antimicrobial activity.\textsuperscript{27–31} In these days, the dynamics of binding of SAILs to proteins represents more active area of investigation. There are numerous studies where the interactions of gelatin with room temperature ionic liquids and concluded the effect of hydrophobicity of ionic liquids on the conformation of gelatin.\textsuperscript{32} Miller et al. have studied the thermal stability and conformational changes in myoglobin in the presence of ionic liquids and reported that myoglobin undergoes destabilization and unfolding in the presence of ionic liquids.\textsuperscript{33} Shu et al. have demonstrated the folding–unfolding behaviour of bovine serum albumin in ionic liquids solutions and reported that probing of protein–ionic liquids interactions is method dependent.\textsuperscript{34}

Till now, the interactions of SAILs and blood plasma protein hemoglobin have been scarcely investigated.\textsuperscript{35–40} Venkatesu et al. have studied the conformational stability of Hb in the presence of varying concentrations of ionic liquids using spectroscopy and molecular docking techniques.\textsuperscript{36} In another report they have investigated the interactions and effect of ionic liquids on the stability/destability of Hb using various techniques.\textsuperscript{39} Thus, the scarcity of such studies in the literature prompted us to carry out a detailed analysis of interaction phenomena in the Hb–SAIL systems because such studies are of prime importance in the field of biotechnological areas. Therefore, with an aim to scrutinize the interactions present between hemoglobin and SAILs herein we have studied the conformational alterations in hemoglobin mediated by SAILs: 1-dodecyl-3-methyl imidazolium chloride [C\textsubscript{12}mim][Cl] and 1-hexyl-3-methylimidazolium dodecylsulfate [C\textsubscript{6}mim][SDS] at pH 6.

In brief we have studied the interactions between hemoglobin and SAILs using a variety of techniques. The tensiometry measurements have been exploited to get better insights into the interaction process and to explore interfacial properties of SAILs in the presence of Hb. The UV-visible spectroscopy, fluorescence spectroscopy and isothermal titration calorimetry were employed to understand the spectroscopic and thermodynamic aspects of Hb–SAILs complex respectively. Dynamic light scattering (DLS) measurements were also carried out to exemplify the variations in hydrodynamic diameter (\(D_h\)) of Hb in the presence of SAILs and the results obtained from all various techniques matches very well. The mode of interaction of both SAILs with Hb is different. The observed results can be considered of special interest toward understanding of Hb–SAILs colloidal chemistry as well as for designing formulations for industrial applications.

2. Experimental

2.1. Materials

Human hemoglobin as lyophilized powder, stored at 2–8 °C was purchased from Sigma Aldrich and used as received. The surface active ionic liquid, 1-dodecyl-3-methylimidazolium chloride [C\textsubscript{12}mim][Cl] and 1-hexyl-3-methylimidazolium dodecylsulfate [C\textsubscript{6}mim][SDS] were synthesized and characterized using \(^1\)H NMR as per the procedures mentioned elsewhere.\textsuperscript{45} The synthetic procedure of SAILs has been discussed in Annexure S1 of ESI\textsuperscript{†} and the structures of the SAILs are given in Scheme 1. AR-grade potassium dihydrogen phosphate and dipotassium hydrogen phosphate were purchased from Merck, India. All the solutions used in the work were prepared in phosphate buffer (5 mM and pH 6) using analytical balance with a precision of ±0.0001 g.

2.2. Methods

2.2.1. Surface tension measurements. The surface tension values (\(\gamma\)) were measured using a Du Nouy ring tensiometer (Kruss Easy Dyne Tensiometer) from Kruss Gmbh (Hamburg, Germany) equipped with thermostat employing ring detachment method at 298.15 K. The aqueous solution of [C\textsubscript{12}mim][Cl] and [C\textsubscript{6}mim][SDS] prepared in buffer were added into Hb solution and stirred for 2–3 min for complete mixing. The resultant solutions were kept for 5 min for equilibration prior to measurement. The data was collected thrice with accuracy ±0.1 mN m\textsuperscript{−1}.

2.2.2. Fluorescence measurements. The steady state fluorescence measurements were performed on Hitachi F-4600 fluorescence spectrophotometer using a 10 mm path length quartz cuvette at 298.15 ± 0.1 K. The titrations were performed by adding concentrated stock solutions of [C\textsubscript{12}mim][Cl] and [C\textsubscript{6}mim][SDS] directly into the quartz cuvette containing 2 mL of 5 \(\mu\)M Hb solution. Changes in the intrinsic fluorescence of Hb were analyzed at an excitation wavelength (\(\lambda_\text{ex}\)) of 280 nm.
and the emission spectra of Hb was recorded in the wavelength range of 300–500 nm wavelength.

2.2.3. UV-visible measurements. The absorption spectra were recorded on a UV-1800 Shimadzu UV-visible spectrophotometer with a quartz cuvette with a path length of 1 cm. The absorbance of pure Hb in the presence and absence of [C₆mim][Cl] and [C₁₂mim][SDS] were recorded at 298.15 K in the range of 200–800 nm.

2.2.4. Isothermal titration calorimetry (ITC) measurements. Calorimetric titrations were performed by using a MicroCal IT200 microcalorimeter at 298.15 K. The sample cell was filled with 200 µL of a solution of 5 µM Hb and the syringe was filled with 40 µL of SAILs solutions and 2 µL aliquots were added into cell. The parameters like time of addition and duration between each addition were monitored by the software provided with the instrument.

2.2.5. Dynamic light scattering (DLS) measurements. DLS measurements were carried out to determine the changes in the size of Hb upon interaction with [C₁₂mim][Cl] and [C₆mim][SDS] using Malvern NanoZS zeta-sizer equipped with 632.8 nm He–Ne laser in backscattering mode at a scattering angle of 173°. The temperature 298.15 K was maintained by in-built temperature controller having an accuracy of ±0.1 K. All the samples prior to measurements were properly filtered from 0.2 µm filters to avoid interference from dust particles.

2.2.6. Turbidity measurements. The turbidity measurements were performed on OAKTON T-100 turbidity meter. The sample cell was filled with 10 mL 5 µM Hb solution and titrated with concentrated solution of SAILs with continuous stirring.

3. Results and discussion

3.1. Surface tension measurements

The tensiometry profiles of SAILs, [C₆mim][SDS] and [C₁₂mim][Cl] in the presence and absence of 5 µM Hb in buffer (pH 6.0) have been shown in Fig. 1(a) and (b) respectively. The γ value decrease linearly for SAILs in buffer with increasing concentration of SAILs before reaching critical micelle concentration (cmc) and after that a nearly constant value is obtained up to the completion of monolayer formation at γcmc. The lower γcmc (27.1) value of [C₆mim][SDS] is an evidence that [C₆mim][SDS] is more densely packed and has better surface activity as compared to [C₁₂mim][Cl] (γcmc = 31.0). The marked decrease in cmc of [C₆mim][SDS] is due to the presence of both [C₆mim]⁺ ions which provide hydrophobicity and 12 carbon long alkyl chain (SDS) as counterion which reduces the hydration and enhances counterion binding. The 6 carbon alkyl chain of [C₆mim]⁺ interact with 12 carbon alkyl chain of anion via hydrophobic interactions similar to that of mixed micelles thus [C₆mim][SDS] possess lower cmc value. The population of [C₆mim][SDS] is more at air–water interface due to effective screening offered by counterion between head groups, van der Waals interaction between alkyl chains of cation and anion along with H-bond and electrostatic interaction between sulphate and imidazolium group. In the presence of Hb, different tensiometric profiles were obtained for SAILs (shown in Fig. 1(a) and (b)). The γ value for Hb was found to be low (50 mN m⁻¹) which suggested that Hb is firmly surface active and get adsorbed at air–water interface. With successive addition of [C₆mim][SDS], γ value decreases sharply up to C₁ after that it decreases at a slow rate up to C₂ followed by again steep decrease up to C₃ to attain saturation (Fig. 1(a)). Initially at lower concentration of [C₆mim][SDS] the sharp decrease in surface tension with lower slope indicates the initial complexation of [C₆mim][SDS] with Hb which leads to the formation of surface active Hb–[C₆mim][SDS] monomer complex. At pH 6 (below isoelectric point) Hb display positive charge. The interactions are mainly electrostatic in nature in dilute concentration region as the positively charged [C₆mim]⁺ ions of SAIL interacts with negatively charged amino acid residues present on the protein along with the hydrophobic interactions between alkyl chain and hydrophobic moieties of Hb. After C₁ the γ value decreases slowly due to the formation of highly surface active Hb–[C₆mim][SDS] monomer complexes which gets adsorbed and accumulated at air–water interface. After C₂, further increase in concentrations of [C₆mim][SDS] sudden sharp decrease in γ value has been observed. More
added [C₆mim][SDS] occupy free sites at interface thus reduces γ up to C₃. In region C₁–C₂ Hb undergoes conformational changes and Hb–[C₆mim][SDS] monomer complex transformed into Hb–[C₆mim][SDS] aggregates complex. The formation of highly surface active Hb–[C₆mim][SDS] aggregate complex is driven by electrostatic interactions though the role of hydrophobic interactions between alkyl chain of [C₆mim][SDS] and hydrophobic parts of Hb cannot be ruled out. Some earlier reports have also suggested the formation of complexes between proteins and ionic liquids.⁴⁶–⁴⁷ After C₃, more addition of [C₆mim][SDS] does not change γ value and saturation arises due to the formation of free micelles along with Hb–[C₆mim][SDS] aggregates. Various interfacial and thermodynamic parameters such as surface excess concentration (τₘₐₓ), minimum area per molecule (Aₘᵢₙ), surface pressure (πₑₘₑₜ), standard Gibb's free energy of micellization (∆Gₘₑₜ) and standard Gibb's free energy of adsorption (∆Gₐₙₑ) were evaluated using relevant equations (Annexure III of ESFI) and provided in Table 1. The τₘₐₓ value for Hb + [C₆mim][SDS] system is higher in comparison to Hb free [C₆mim][SDS] system which implies that mixture has better surface activity than pure [C₆mim][SDS]. As expected opposite trend has been observed in Aₘᵢₙ values indicating the better compactness due to densely packed molecules in the presence of Hb as compared in buffer. The negative values of both ∆Gₘₑₜ and ∆Gₐₙₑ clearly indicates the feasibility of both micellization and adsorption phenomenon.

In case of Hb + [C₁₂mim][Cl] system the behaviour of γ is quite different as compared to Hb + [C₆mim][SDS] system. As clear from Fig. 1(b) the γ value of [C₁₂mim][Cl] in buffer decreases linearly and showed a minima near cmc after that γ attain constant region. Whereas in the presence of Hb tensiometric profile exhibit three breaks points owing to Hb–[C₁₂mim][Cl] interactions. Initially at low concentration of [C₁₂mim][Cl], γ decreases sharply upto C₁ due to the formation of surface active Hb–[C₁₂mim][Cl] monomer complex at interface owing to the presence of mainly electrostatic interactions along with hydrophobic interactions. The change in slope is less in Hb–[C₁₂mim][Cl] system when compared to Hb–[C₆mim][SDS] system indicating the presence of more strong electrostatic interactions in [C₁₂mim][Cl] monomers and Hb. In region C₁–C₂ a plateau region is formed. It must be considered that beginning concentrations of plateau is the region where Hb–[C₁₂mim][Cl] monomer complex starts to transform into Hb–[C₁₂mim][Cl] aggregate complex and ending concentration is the region where the aggregates continued to bind with protein sites that restrict the transfer of SAILs molecules to the interface to maintain constant γ value. Guillot et al. has reported the formation of flat region in the aqueous solution of a poly-electrolyte, carbosymylecellulose, and a cationic surfactant, dodecyltrimethylammonium bromide (DTAB).⁴⁸ On reaching C₂ the formed aggregate complex collapsed and invaded into bulk from interface which leads to rise in γ value to maximum. Above C₃ γ remains almost constant and further addition of SAIL leads to the formation of free micelles. The calculated interfacial and thermodynamic parameters for [C₆mim][Cl] in the absence and presence of Hb have also been computed and given in Table 1. The lower τₘₐₓ and higher Aₘᵢₙ of [C₁₂mim][Cl] as compared to [C₆mim][SDS] indicates the lower surface activity of [C₁₂mim][Cl]. For Hb–[C₆mim][SDS] system, there is no slope change prior to reach cmc (C₂) hence τₘₐₓ and Aₘᵢₙ values were not determined for mixture. The more negative value of ∆Gₘₑₜ in pure SAILs solutions when compared to Hb–SAIL system indicate the greater feasibility of micellization of SAILs in the absence of Hb. The value of various transition concentrations (C₁, C₂ and C₃) for Hb–[C₆mim][SDS]/[C₁₂mim][Cl] system observed from different techniques are given in Table 2.

3.2. Spectroscopic study (UV-vis absorption and fluorescence measurements)

To study the change in protein conformation upon binding with ligands UV-vis spectroscopy and fluorescence spectroscopy methods are good to be adopted.⁴⁹,⁵⁰ Both are very simple but efficacious techniques to analyze the binding of SAILs [C₁₂mim][Cl] and [C₆mim][SDS] with Hb. The isoelectric point of Hb is 6.8 and we have studied the interaction of SAILs with

### Table 1

<table>
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<tr>
<th>System</th>
<th>cmc (mM)</th>
<th>γₑₘₑₜ (mN m⁻¹)</th>
<th>πₑₘₑₜ (mN m⁻¹)</th>
<th>10⁶τₘₐₓ (mol m⁻²)</th>
<th>Aₘᵢₙ (Å²)</th>
<th>∆Gₘₑₜ (kJ mol⁻¹)</th>
<th>∆Gₐₙₑ (kJ mol⁻¹)</th>
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<td>44.99</td>
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<td>34.3</td>
<td>15.7</td>
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### Table 2

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<th>Flu</th>
<th>UV</th>
<th>DLS</th>
<th>Turbidity</th>
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<tr>
<td>C₁</td>
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<td>C₃</td>
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<td>1.43</td>
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<tr>
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<tr>
<td>C₁</td>
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Hb at pH 6 where Hb displays positive charge. The UV-vis spectra of Hb shows several peaks, one at 276 nm due to phenyl group of Trp and tyrosine residue, 406 nm (Soret band, due to $\pi \rightarrow \pi^*$ transition), 500, 536, 576 and 630 nm (due to oxy band or Q band and ligand to metal charge transfer spectra). The allowed $\pi \rightarrow \pi^*$ transitions are mainly due to heme group of protein which is embedded in the hydrophobic pocket formed by protein backbone.\textsuperscript{51} Thus appearance of Soret band at 405 nm confirms the folded or native structure of Hb. Any disturbance or changes in Soret band reveals the reduction in $\alpha$-helix content of Hb.\textsuperscript{52} Fig. 2(a) and (b) shows the UV-vis absorption spectra of Hb with increasing concentration of [C$_6$ mim][SDS] and [C$_{12}$ mim][Cl] respectively whereas Fig. 3(a) and (b) shows the variation of absorption as a function of SAILs concentration. The expanded region considering the absorbance peaks at 495, 535, 575 and 630 nm have been shown in Fig. S2(a) and (b) in ESI.\textsuperscript{†} With the addition of [C$_6$ mim][SDS], absorption peak of Soret band at 405 nm decreases linearly up to C1 without any shift in $\lambda_{\text{max}}$ (Fig. S3\textsuperscript{†}). These results indicates that initially [C$_6$ mim][SDS] interact with backbone of Hb through weak electrostatic interactions, hydrogen bonding and hydrophobic interactions. Once the C$_1$ reached, the turbidity appears which results into increment of UV-vis absorption upto C$_1'$ followed by decrement upto C$_0$. The appearance of turbidity is the result of charge neutralisation of oppositely charged [C$_6$ mim][SDS] and amino acid residues of Hb. Once the C$_1$ reached, the $\lambda_{\text{max}}$ also shifts and a new peak starts to form at 415 nm, the main characteristic of hemichrome.\textsuperscript{53} The formation of new band at 415 nm clearly indicates the oxidation of oxyhemoglobin to methemoglobin (metHb) and latter converted into hemichrome.\textsuperscript{54} The autooxidation process is not a simple but it is associated with the dissociative mechanism with the loss of superoxide anion (O$_2^-$) from oxyHb and leads to the formation of 5-coordinated intermediate. This vacant position in 5-coordinated Fe$^{3+}$ complex is thus occupied by either hydroxyl ion or by water molecules (from surrounding medium) to form metHb.\textsuperscript{55} But in the presence of surface active ionic liquids the metHb gets converted to hemichrome. The potentiality of interactions present between Hb and SAILs bring a comprehensive change in surrounding environment of
the heme pocket and induce the conversion of met-Hb to hemichrome by breaking hydrogen bond between distal histidine imidazole and water molecule/hydroxyl ions. With increasing concentration of \([\text{C}_6\text{mim}]\text{[SDS]}\) after \(C_2\) the absorption decreases but \(\lambda_{\text{max}}\) remains unchanged unlike Hb–\([\text{C}_{12}\text{mim}]\text{[Cl]}\) system (discussed ahead) which reveals that addition of more \([\text{C}_6\text{mim}]\text{[SDS]}\) leads to formation of \([\text{C}_6\text{mim}]\text{[SDS]}\)–Hb aggregates via strong electrostatic and hydrophobic interactions. With careful observation it was observed that turbidity appears only up to \(C_1\) after that it disappears with sharp linear decrement in UV–vis absorption up to \(C_2\). The disappearance of turbidity clearly indicates that when \(C_2\) reached the hydrophobic interactions between hydrophobic tail of \([\text{C}_6\text{mim}]\text{[SDS]}\) and Hb dominates over the electrostatic interactions and they stabilized the hemichrome formation. These results were contrary to earlier reported Hb–SDS system where SDS micelles induce the heme release and solubilise them in micellar core whereas in \([\text{C}_6\text{mim}]\text{[SDS]}\) molecules the long hydrophobic chain forms network like structure thus prevent the release of heme molecules.\(^{36}\) These results were also confirmed from the changes in the fluorescence intensity upon addition of \([\text{C}_6\text{mim}]\text{[SDS]}\) as shown in Fig. 4(a). The fluorescence of proteins arises from Trp, tyrosine and phenylalanine present in protein. In hemoglobin the intrinsic fluorescence primarily arises from 3 Trp residues (\(\alpha214\text{Trp}, \beta215\text{Trp} \) and \(\beta216\text{Trp}\)).\(^{37}\) The hemoglobin exhibit weak fluorescence due to the quenching of Trp fluorescence by heme group of porphyrin ring by radiationless energy transfer.\(^{38}\) In presence of \([\text{C}_6\text{mim}]\text{[SDS]}\) the fluorescence intensity of Hb increases might be because of two main reasons. First, is the increased hydrophobicity around heme group and second could be the prevention of quenching effect owing to non-radiative energy transfer from Trp residue to heme group. The heme group is initially embedded in the hydrophobic cavity of the protein but with the addition of SAIL the hydrophobic tail of \([\text{C}_6\text{mim}]\text{[SDS]}\) penetrate into hydrophobic cavity of Hb and heme group is exposed which leads to the fluorescence enhancement. The fluorescence intensity increase slowly up to \(C_1\) which indicates the presence of interactions between Hb–\([\text{C}_6\text{mim}]\text{[SDS]}\). After \(C_1\) the fluorescence intensity increases and mixture become turbid indicates that Hb–\([\text{C}_6\text{mim}]\text{[SDS]}\) aggregates formed owing to strong electrostatic interactions between oppositely charged groups but once \(C_2\) reached turbidity disappears and blue shift in \(\lambda_{\text{max}}\) have also been observed. The fluorescence intensity sharply increases after \(C_2\) and continues to increase up till \(C_3\) reached after which saturation comes. These results reveal that after the disappearance of turbidity hydrophobic interactions dominates over electrostatic interactions. The unchanged \(\lambda_{\text{max}}\) suggest that hemichrome is not further changed and stabilized by \([\text{C}_6\text{mim}]\text{[SDS]}\) micelles. The sharp rise in fluorescence intensity is up to \(C_3\) after that it becomes constant as free micelles formed and Hb surface get saturated with formed micelles. As the turbid nature of Hb–\([\text{C}_6\text{mim}]\text{[SDS]}\) solution was monitored by naked eye hence turbidity measurements were further performed. The variation of turbidity of Hb as a function of concentration of \([\text{C}_6\text{mim}]\text{[SDS]}\) and \([\text{C}_{12}\text{mim}]\text{[Cl]}\) has been provided as Fig. S4(a) and (b) respectively and the transitions extracted from the plot matches well with the transitions obtained from other techniques.

The UV–vis and variation in fluorescence intensity plot of Hb in the presence of \([\text{C}_{12}\text{mim}]\text{[Cl]}\) show different results as shown in Fig. 2(b), 3(b) and 4(b). The fluorescence emission spectra of Hb with varying concentration of \([\text{C}_{12}\text{mim}]\text{[Cl]}\) has been shown as Fig. S5.† As shown in Fig. 2(b) and 3(b) the absorption of Soret band of Hb decreases linearly up to \(C_1\) without any change shift in \(\lambda_{\text{max}}\) indicates that initially \([\text{C}_{12}\text{mim}]\text{[Cl]}\) interact with Hb via weak forces. The decrease in absorption is continues up to \(C_1\) after that the decrement in absorption peak get slow. When the concentration of \([\text{C}_{12}\text{mim}]\text{[Cl]}\) reached above \(C_1\) the absorption of Soret band decreases gently along with red shift from 405 nm to 413 nm. This change in Soret band after \(C_1\) confirms that geometric distortion arise due to dielectric and electrostatic interactions of proteins and changes the symmetry in heme environment. These results were also in accordance with the fluorescence measurements results.

![Fig. 4](image-url)  
**Fig. 4** Variation of fluorescence emission intensity of Hb (5 µM) in PBS buffer as a function of SAIL concentration for (a) \([\text{C}_6\text{mim}]\text{[SDS]}\) and (b) \([\text{C}_{12}\text{mim}]\text{[Cl]}\) when excited at 280 nm.
(Fig. 4(b)) where at lower concentration of [C_{12}mim][Cl] (below C_1) the fluorescence intensity of Hb increases which has been attributed to conformational changes in Hb on interacting with SAIL molecules. On partial unfolding of Hb, the distance between Trp residue and heme group is increased which shield the nonradiative energy transfer from Trp to heme moiety. With further increasing [C_{12}mim][Cl] concentration the intensity decreases due to the formation of Hb–[C_{12}mim][Cl] aggregate complex guided by enhanced hydrophobic interactions between [C_{12}mim][Cl] tail and hydrophobic moieties of Hb. After C_2 addition of more [C_{12}mim][Cl] increases fluorescence intensity due to increased hydrophobicity around heme group and then reaches upto C_3 after that the addition of more SAIL leads to the saturation of Hb in aqueous medium. Whereas in case of UV-vis, once C_2 reached the absorbance decreases very slightly but blue shift of Soret band has been observed, latter shifts from 413 nm to 404 nm. The transitions in χ_{max} of Soret band from 405 nm to 413 nm and from 413 nm to 404 nm at higher concentration of [C_{12}mim][Cl] indicate the transformation of aquomethHb to hemichrome and hemicr–heme monomer which is solubilised in the [C_{12}mim][Cl] system. The negative value of ΔG indicates the feasibility and spontaneity of Hb–SAILs complex formation.

3.3. Quantitative evaluation of Hb–SAILs interaction

In view of tensiometry, UV-vis spectroscopy and fluorescence measurements it was concluded that at lower concentration of SAILs (before C_1) Hb–[C_{6}mim][SDS] and Hb–[C_{12}mim][Cl] monomer complex were formed and guided by electrostatic and hydrophobic interactions. Thus for quantitative estimation of the binding of SAILs to the Hb the absorbance data is used in the Benesi–Hildebrand equation which is given as follows:  
\[ \frac{1}{A_0 - A} = \frac{1}{K_a (A_1 - A_0)} \frac{1}{[\text{SAIL}]^2} + \frac{1}{A_1 - A_0} \]  
where A_0, A and A_1 are the absorbance in the absence, at intermediate Hb–SAILs complex concentration and at infinite concentration of SAILs respectively and K_a is the binding constant. The plot of 1/A–A versus 1/[SAIL] gives straight line when n = 1 indicating 1 : 1 stoichiometry for these complexes (Fig. 5). The stoichiometry was further confirmed using Job’s plot (discussed in Annexure III†). The binding constant further used to calculate free energy change (ΔG) for Hb–[C_{6}mim][SDS] and Hb–[C_{12}mim][Cl] complexes using eqn (2) and provided in Table 3.

\[ \Delta G = -RT \ln K_a \]  
where R and T are universal gas constant and temperature respectively. It can be observed from the K_a values that binding constant is stronger for [C_{6}mim][SDS] + Hb system as compared to [C_{12}mim][Cl] due to difference in their hydrophobicity. It must be kept in mind that the binding constant determined from B–H equation gives the quantitative detail of interactions present in monomer region only. Thus in Hb–SAIL monomer complex both electrostatic and hydrophobic forces are present, but electrostatic forces are strong in Hb–[C_{6}mim][SDS] system as compared to Hb–[C_{12}mim][Cl] system. The negative value of ΔG indicates the feasibility and spontaneity of Hb–SAILs complex formation.

**Fig. 5** Benesi–Hildebrand plot for binding constant determination using changes in absorbation spectra of Hb (5 μM) in the presence of varying concentration of SAILs.
Table 3  Estimated binding constant ($K_b$), Gibbs free energy change ($\Delta G$) and correlation coefficients ($R_c$) for Hb–[C6mim][SDS] and Hb–[C12mim][Cl] systems from UV-visible measurements

<table>
<thead>
<tr>
<th>System</th>
<th>$K_b$ ($\times 10^4$ M$^{-1}$)</th>
<th>$\Delta G$ (kJ mol$^{-1}$)</th>
<th>$R_c$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hb–[C6mim][SDS]</td>
<td>16.93</td>
<td>-24.13</td>
<td>0.9989</td>
</tr>
<tr>
<td>Hb–[C12mim][Cl]</td>
<td>0.22</td>
<td>-13.40</td>
<td>0.9997</td>
</tr>
</tbody>
</table>

3.4. Isothermal titration calorimetry (ITC) measurements

The ITC experiments were performed further to explain the enthalpy changes in binding of Hb with SAILs in premicellar and postmicellar regions. The enthalpogram of [C6mim][SDS] and [C12mim][Cl] aggregation in the absence and presence of Hb at 298.15 K have been shown in Fig. 6(a) and (b) respectively. To interpret the actual heat changes upon interaction of [C6mim][SDS] with Hb the enthalpogram of [C6mim][SDS] in buffer has been subtracted from [C6mim][SDS] binding to Hb and shown in Fig. 6(a). The difference in heat changes due to binding of [C6mim][SDS] with Hb clearly demonstrates the presence of interactions between them. The subtracted enthalpogram has been divided into 4 regions and unlike Hb–[C6mim][SDS] system, Hb–[C12mim][Cl] system has less heat changes. In region I, when the concentration of [C12mim][Cl] is below C1 the observed endothermic heat changes were attributed to the initial binding of [C12mim][Cl] monomers via weak electrostatic and hydrophobic interactions. The hydrophobic chain of [C12mim][Cl] interacts with the hydrophobic residues present on the surface of Hb thus favours the hydrophobic interactions mainly. When [C12mim][Cl] content is above C1, the dH values changes with increasing binding of [C12mim][Cl] aggregates to Hb. In region II the enthalpy changes decreases endothermically and were attributed to conformational changes in Hb. The [C12mim][Cl] molecules on binding to Hb causes the geometric distortion and exposed the charged amino acid residues and negatively charged residues which are now free to bind with positively charged head group of [C12mim][Cl] via electrostatic interactions. The spectroscopic measurement also reveals that after C2 hemichrome is formed and denaturation of Hb occurs. Thus addition of [C12mim][Cl] makes the hemoglobin denatured and micelles like aggregates formed at C2 (12.3 mM). In region III when the [C12mim][Cl] concentration is above C2 the endothermic heat changes were observed upto C3 as denatured Hb has more hydrophobic chains which are exposed thus [C12mim][Cl] induces the heme group to be released from hemoglobin.

Fig. 6  Variation of the observed enthalpy ($\Delta H$) with the SAIL concentration for the titration of (a) [C6mim][SDS] and (b) [C12mim][Cl] into PBS buffer and Hb in PBS buffer.
and subsequently solubilised in the micellar core. After C₁ the unchanged or nearly constant heat changes values have been observed due to the saturation of Hb.

3.5. Dynamic light scattering (DLS) measurements

DLS measurements were used to investigate the change in size of Hb upon interacting with SAILs. In the absence of SAILs the apparent size of Hb in buffer is 6 nm indicates that protein exist in the form of random coil due to strong hydrophobic interactions among polypeptide chains. The variation in size of Hb in the presence and absence of SAILs has been shown in Fig. 7(a) and (b). In the presence of [C₆mim][SDS] the hydrodynamic diameter (Dₗ) changes from 6 nm to 7.5 nm indicate that some kind of interactions take place between [C₆mim][SDS] and Hb. Further with increasing concentration of [C₆mim][SDS] the Dₗ value reaches 419 nm indicating the formation of complex between Hb and [C₆mim][SDS] monomers governed by strong electrostatic and hydrophobic interactions. With further increasing [C₆mim][SDS] concentration, the Dₗ value sharply increases and reaches up to C₁ due to the formation of larger Hb [C₆mim][SDS] aggregates or coacervates which makes the solution turbid. After C₁ the size sharply decreases from 889 nm to 380 owing to either disintegration of large sized aggregates or resolubilization of aggregates and the consequence is the disappearance of turbidity. The size of the mixed aggregates continues to decrease till C₃ reached because more addition of SAIL induces repulsive interactions which destabilize and breakdown the mixed aggregates. At higher concentration well above C₅, Dₗ slightly varies indicating the stabilization of mixed aggregates governed by hydrophobic forces between alkyl chain of [C₆mim][SDS] and hydrophobic patches on Hb.

The variation of hydrodynamic diameter (Dₗ) of Hb versus [C₁₂mim][Cl] concentration has been shown in Fig. 7(b). With the addition of [C₁₂mim][Cl] the Dₗ value of Hb suddenly increased upto 666 nm indicating that initially hydrophobic chain of [C₁₂mim][Cl] interact with the hydrophobic moiety of Hb which results into the formation of large Hb–[C₁₂mim][Cl] monomer complex. At C₁ the monomer complex transformed into Hb–[C₁₂mim][Cl] aggregate complex but with the more addition of [C₁₂mim][Cl] the large sized Hb–[C₁₂mim][Cl] aggregates disintegrate into relatively small sized aggregates thus Dₗ values decreases sharply upto 195 nm and after that it decreases slowly upto C₂. In concentration region C₁–C₂ conformational changes occurs in Hb and folded structure of Hb can be expected to form with the loss of superoxide anion with the release of heme monomer. These results are also justified from UV-visible results where pentacoordinated species is formed with the release of heme. After C₂ with further addition of [C₁₂mim][Cl] hydrophobic interactions increases which induces the heme monomer to leave the hydrophobic pocket of protein thus size increases but only upto C₃. Once C₃ reached the Dₗ value remains unchanged due to the saturation of Hb with the addition of more [C₁₂mim][Cl].

4. Conclusions

In conclusion, in this work we have studied the comparative studies on the interactions of Hb with [C₆mim][SDS] and [C₁₂mim][Cl] with the aid of various techniques. The surface tension measurements have been performed to study the interactional behavior at the air–solution interface and the results have been discussed in detail giving various transitional concentrations and evaluating τₘₐₓ and Aₗₘᵢₙ values. The binding mechanism and structural alterations in Hb induced by SAILs have been thoroughly studied using spectroscopic measurements. The results based on the experiments confirms the formation of hemichrome in Hb–SAIL systems but the ejection of heme from Hb hydrophobic pocket to micellar core takes place only in Hb–[C₁₂mim][Cl] system, whereas the long alkyl chain of [C₆mim][SDS] forms network like structure and prevent the release of heme molecule thus hemichrome remains stabilized. The addition of [C₆mim][SDS] to Hb leads to aggregation of protein which is evidenced from appearance of turbidity and appearance of larger aggregates in DLS measurements. Despite these investigations, ITC measurements were also performed to investigate the enthalpy changes and exact

Fig. 7 Variation of the hydrodynamic diameter (Dₗ) as a function of SAIL concentration (a) [C₆mim][SDS] and (b) [C₁₂mim][Cl] in Hb (5 μM) solution in PBS buffer.
nature of molecular interactions present in Hb–SAIL system. As protein–surfactant systems are consistent with the biological membranes so gives the opportunity to explore surface active ionic liquids to understand various biomembrane-protein functions. Our findings provide an important insight into the Hb–SAIL interactions essential for determining their future use as excipients in pharmaceutical formulations containing proteins. These results may also show the potential utility of Hb–SAILs interactions in biotechnological processes and pharmaceutical formulations.

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