What a difference a methyl group makes – probing choline–urea molecular interactions through urea structure modification†

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There is a lack of fundamental knowledge on deep eutectic solvents, even for the most extensively studied mixtures, such as the mixture of choline chloride and urea, which prevents a judicious choice of components to prepare new solvents. The objective of this work is to study and understand the fundamental interactions between choline chloride and urea that lead to the experimentally observed melting temperature depression. To do so, the structure of urea was strategically and progressively modified, in order to block certain interaction centres, and the solid–liquid equilibrium data of each new binary system was experimentally measured. Using this approach, it was concluded that the most important interaction between choline chloride and urea occurs through hydrogen bonding between the chloride anion and the amine groups. Any blockage of these groups severely hampers the melting point depression effect. Raman spectroscopy and DFT calculations were utilized to study in more detail this hydrogen bonding and its nuances.

Introduction

The so-called deep eutectic solvents (DES) have emerged at the beginning of this century as possible substitutes for hazardous organic solvents. They are usually defined as a mixture of a hydrogen bond acceptor (HBA) and a hydrogen bond donor (HBD) that form strong hydrogen bonds, leading to melting temperatures substantially lower than those predicted by assuming an ideal liquid mixture or those presented by the individual pure components. Cholinium chloride (ChCl) is one of the most recurring substances used to prepare DES and a large set of experimental solid–liquid equilibrium data on mixtures containing ChCl is already available in the literature, with most of the DES prepared presenting low toxicity and being readily biodegradable.

Even though the deep eutectic solvent composed of urea and choline chloride was identified as a green solvent in 2003, its structure and molecular interactions, along with microscopic phenomena such as solid–liquid equilibria (SLE), still hold some mysteries. A recent study by van den Bruinhorst and co-workers compares the currently available SLE data for the cholinium chloride/urea system, pointing out that the liquidus temperatures reported by Abbott and co-workers, who measured freezing temperatures, are consistently lower than the melting temperatures reported by other authors due to the phenomenon of supercooling, while Meng et al. attribute the observed discrepancies to the presence of water. Even though not shown in the SLE diagrams reported, XRD data suggest the formation of a 1:2 cholinium chloride–urea cocrystal, whose structure is yet to be explored.

Several experimental and theoretical studies have been devoted to studying the liquid structure and main interactions in the cholinium chloride–urea eutectic, generally pointing to a complex mesh of “overall similar interactions between all constituents” as expressed in Zahn’s hypothesis “similia similibus solvuntur” (“like dissolves like”). Ashworth and co-workers used DFT calculations to assess intermolecular interactions among the cation, anion and hydrogen bond donor. They argue that urea–choline contacts might compete with cholinium–urea interactions since individual OH···Cl hydrogen bonds rank as stronger than N–H···Cl contacts, which show up as medium-strength interactions, surpassed by OH···O=C bonds. This view is only partially supported by the neutron diffraction study of Hammond and colleagues, performed at 303 K, describing a “radially layered sandwich structure” where...
the cholinium cation’s –OH and urea’s –NH moieties strongly interact with the chloride anion while maintaining softer cholinium/C1/C1/C1 urea and urea/C1/C1/C1 urea contacts. Using the same technique, Gilmore and co-workers showed that at 338 K the OH···Cl correlation is weakened in favour of forming OH···OH contacts, while the urea···Cl interaction remains dominant. Charge transfer between the chloride anion and the hydrogen bond donor was initially proposed by Carriazo et al. as constituting the main driver for the melting point depression, yet this view has since been disputed by Zahn and co-workers’ ab initio molecular dynamics simulation, suggesting that charge transfer occurs mainly between the cation and the anion. Another ab initio study, performed by Wagle and colleagues, proposes that the melting points of three cholinium chloride based DES correlate with the bond order of the cholinium···chloride interaction but not with the bond order of the HBD···chloride contacts. Migliorati and colleagues compared eutectic mixtures based on urea and either cholinium chloride or butyltrimethylammonium chloride, finding that while the urea···chloride interaction is relatively favoured in the latter this does not translate into a more pronounced melting point depression. It is then clear that the contribution of the HBD···anion interaction to the melting point depression remains controversial. Our own past vibrational spectroscopy study of the cholinium chloride/urea system failed to unambiguously isolate the vibrational fingerprint of the urea···chloride interaction so that its importance to the melting point depression could not be fully ascertained.

Better knowledge of the molecular interactions of cholinium chloride in deep eutectic solvents would lead to a wiser and more rigorous choice of components to prepare new DES and, thus, is of paramount importance. The main objective of the present work is to understand the fundamental interactions between cholinium chloride and urea by strategically modifying the structure of urea to block specific interaction sites. For that purpose, the solid–liquid equilibrium phase diagrams of five binary mixtures composed of cholinium chloride and urea, thiourea, methylurea, 1,3-dimethylurea or 1,1-dimethylurea were measured and analysed. Raman spectroscopy was used as an auxiliary tool to corroborate our findings. The objective of using thiourea is to probe the importance of the carbonyl group in the cholinium–urea interaction, whilst the methylurea series is used to assess the role of urea’s amine groups in said interaction.

**Experimental details**

**Chemicals**

Pertinent properties for the substances experimentally used in this work are reported in Table 1. Their chemical structures are depicted in Fig. 1. Due to its high hygroscopicity, cholinium chloride was dried under vacuum (0.1 Pa and 298 K) for at least 72 h before use. The remaining compounds were stored at room temperature and used as received. The water content of each compound was measured using a Metrohm 831 Karl–Fischer coulometer with the analyte Hydranal from Riedel-de Haeßn and found to be lower than 600 ppm for cholinium chloride and lower than 10 ppm for the remaining substances.

**Solid–liquid equilibrium phase diagrams**

All samples were prepared inside a dry argon glovebox chamber, weighing each amount of individual pure component inside a
vial using an analytical balance (ALS 220-4N from Kern) with an accuracy of ±0.002 g. For each system its entire composition range was covered. The vials were then heated under stirring, until complete fusion, and then recrystallized. After this heat cycle treatment, the mixtures obtained presented (i) a paste like consistency or (ii) completely solid recrystallization. Depending on the physical state of the final mixture, different melting temperature measurement methods were applied. In case (i), a visual method using an oil bath was applied, where the mixtures were inserted in an oil bath and gradually heated until complete melting. The temperature was controlled with a PT100 probe that was previously calibrated. For case (ii), the solid mixtures were ground inside the glove box chamber and the powder was filled into a glass capillary. The melting point was determined using a thermocouple connected to a multimeter. Raman spectroscopy was performed with a WITec alpha300 RAS+ spectrometer operating at 532 nm as the excitation source with the power set at 25 mW. The samples were enclosed in capped NMR vials using an analytical balance (ALS 220-4N from Kern) with an accuracy of ±0.002 g. For each system its entire composition range was covered. The vials were then heated under stirring, until complete fusion, and then recrystallized. After this heat cycle treatment, the mixtures obtained presented (i) a paste like consistency or (ii) completely solid recrystallization. Depending on the physical state of the final mixture, different melting temperature measurement methods were applied. In case (i), a visual method using an oil bath was applied, where the mixtures were inserted in an oil bath and gradually heated until complete melting. The temperature was controlled with a PT100 probe that was previously calibrated. For case (ii), the solid mixtures were ground inside the glove box chamber and the powder was filled into a glass capillary. The melting point was determined using a thermocouple connected to a multimeter.

Regardless of the method used, all measurements were repeated at least three times, possessing an estimated reproducibility of ±1.4 K (see the ESI†).

It has been shown in the literature that some deep eutectic solvents, specifically malonic acid-based systems, may undergo decomposition during their preparation using the heating cycle described above. However, for all systems studied, no decomposition was detected in any of the mixtures prepared. Moreover, all solid–liquid equilibrium data obtained are consistent, without any indication of component decomposition.

**Raman spectroscopy**

Raman spectra were collected using a WITec alpha300 RAS+ (WITec, Ulm, Germany) Raman microscope. An Nd:YAG laser operating at 532 nm was used as the excitation source with the power set at 25 mW. The samples were enclosed in capped NMR tubes, previously filled inside a dry argon glovebox chamber, and placed within a homemade heating apparatus of the Harney–Millier type. The temperature inside the sample holder was monitored using a thermocouple connected to a multimeter. Raman spectra of the eutectic mixtures based on urea, methylurea and 1,1-dimethylurea, along with their interaction pair with the 6-311+G(d,p) basis set in the gas phase. In addition, the complete series was also computed with the 6-31+G(d,p) basis set optimized in the gas phase and later embedded in the generic ionic liquid solvation model, SMD-GIL. Prior to the DFT calculations, the conformational space was systematically explored with molecular dynamics 100 ps runs with a tight binding novel Hamiltonian. Frequencies were calculated to ensure that all optimized geometries were real minima (no negative eigenvalues). CM5 point charges based on the Hirshfeld partition scheme were computed for all minima. All DFT calculations were performed with the Gaussian 09 Rev A.02 package. Interaction energies ($E_{\text{int}}$) were calculated by subtracting the electronic energy without zero-point correction of the constituents ($E_{\text{M1}}$ and $E_{\text{M2}}$) from that of the interacting pair ($E_{\text{Pair}}$) in the gas phase or embedded in a solvent continuum:

$$E_{\text{int}} = E_{\text{Pair}} - E_{\text{M1}} - E_{\text{M2}}.$$  (1)

In addition to the point charges, the electrostatic potential mapped onto the electron density was also computed for the urea family. The results obtained are presented in the ESI† as visual aids. They were calculated using TURBOMOLE (software package ‘TmoleX†), employing the 6-311++G(d,p) basis set in the gas phase and the M06-2X level of theory.

**Thermodynamic framework**

The solid–liquid equilibrium curve of a component within a eutectic-type mixture, when no solid-phase transitions are present, can be described using the following equation:

$$\ln(x_i \cdot \gamma_i) = \frac{A_{\text{hi}}}{R} \left( \frac{1}{T_{m,i}} - \frac{1}{T} \right) + \frac{A_{\text{m}}C_{\text{pi}}}{R} \left( \frac{T_{m,i}}{T} - \ln \frac{T_{m,i}}{T} - 1 \right)$$  (2)

where $x_i$ is the mole fraction of component $i$ in the liquid phase, $\gamma_i$ its activity coefficient, $A_{\text{hi}}$ its melting enthalpy, $T_{m,i}$ its melting temperature and $A_{\text{m}}C_{\text{pi}}$ its heat capacity change upon melting, whilst $R$ is the ideal gas constant and $T$ is the absolute temperature of the system. Since the contribution of the last term present in eqn (2) is typically negligible and the value of $A_{\text{m}}C_{\text{pi}}$ is usually not available for the compounds used in this work, eqn (2) is well approximated by:

$$\ln(x_i \cdot \gamma_i) = \frac{A_{\text{hi}}}{R} \left( \frac{1}{T_{m,i}} - \frac{1}{T} \right)$$  (3)

Eqn 3 is useful in two ways: it can be used to predict the SLE phase diagram of a mixture considering its liquid phase as ideal by setting $\gamma_i = 1$, or it can be used to calculate the activity coefficients of the components in a mixture using available experimental SLE data. As such, solid–liquid phase diagrams can be used to assess the liquid phase non-ideality of mixtures and, thus, the specific interactions between their components.

**Results and discussion**

The five SLE phase diagrams measured are depicted in Fig. 2 along with the ideal liquid phase curves and the experimental activity coefficients, calculated using eqn (3) and the melting properties in Table 1. Detailed experimental data are reported in Table S1 of the ESI†. Since Abbott et al. already reported the freezing temperatures for the system cholinium chloride/urea, in a small composition range, and for the remaining systems,
in a 2:1 mole proportion, these data are also included. An extensive comparison with literature data for the system cholinium chloride/urea is shown in Fig. S1 in the ESI.† The main reason for the discrepancies observed is related to the high hygroscopicity of cholinium chloride as shown by Meng et al.16 This is the main reason for the careful preparation of all mixtures in a water-free environment as described in the Experimental section.

We first examine the systems cholinium chloride/urea and cholinium chloride/thiourea, the sole difference being the replacement of urea’s carbonyl group by thiourea’s thionyl group. The melting curves and activity coefficients (calculated using eqn (3)) of the components in both systems were merged into Fig. 3 to simplify comparison and analysis.

Fig. 3a reveals an almost identical behaviour in the melting curve of cholinium chloride when mixed with urea or thiourea. In turn, this corresponds to a very similar behaviour of its activity coefficients in both systems, as Fig. 3c shows. Both urea and thiourea also present a remarkably similar behaviour when mixed with cholinium chloride. This may not be easy to see in Fig. 3b but is clear in Fig. 3d, which depicts similar activity
coefficients for both substances. Taking all this into account, it can be safely assumed that the main interactions between cholinium chloride and urea do not involve its carbonyl group, i.e. the interactions between urea and cholinium chloride are not dominated by the carbonyl group interactions in the mixture.

Fig. 4 presents the atomic charges for the urea family, calculated as described in the computational details section (a comparison between the C2 and D2h conformations of urea and thiourea is included in Fig. S2 of the ESI†). The sulphur atom provides thiourea with a larger negative surface than urea, albeit with a more disperse (less intense) charge (see Fig. S3 for a visual aid, ESI†). On the other hand, the electrostatic field generated by the protons is very similar for both molecules. As such, and considering that the swap of an oxygen for a sulphur does not significantly alter the interactions between urea and cholinium chloride, it can be safely assumed that cholinium chloride’s main interactions with urea are via the positive hydrogen atoms in the amine groups, which may form a hydrogen bond with the chloride anion of cholinium chloride.

The interaction energies for the chloride anion with thiourea or urea and for cholinium chloride with two molecules of thiourea or two molecules of urea, embedded in a generic liquid solvent continuum, were computed. Thiourea compounds are stabilized versus urea in excess of 16 and 24 kJ mol⁻¹ respectively. This arises from the substantial differences in the charges of the carbonyl and the thionyl groups for both molecules, and/or enhanced dispersion interactions with the larger, thus more polarizable, sulphur atom. The fact that the thermodynamic behaviour of thiourea and urea, when mixed with cholinium chloride, is essentially similar means that neither of these interactions should play a fundamental role in the previously reported deviations from ideality.

After analysing the effect of the exchange of the oxygen of urea for the sulphur of thiourea, which did not seem to induce significant differences in the behaviour of the SLE phase diagrams of these compounds, the next step is to structurally modify the amine groups of urea by methylation. The results are summarized in Fig. 5, which shows cholinium chloride combined with urea, methylurea or 1,3-dimethylurea. In this series the amine groups of urea are progressively blocked by methylation.

Fig. 5a clearly shows a trend in the behaviour of cholinium chloride with the blockage of the amine groups in urea. Initially, cholinium chloride behaves ideally when mixed with urea. However, when mixed with methylurea, cholinium chloride reveals positive deviations from ideality, which are further enhanced in the 1,3-dimethylurea system. A complementary trend is seen in Fig. 5d, where urea shows the largest negative deviations from ideality, while 1,3-dimethylurea has an almost ideal behaviour. This supports the hypothesis, presented from the analysis of the urea and thiourea systems, that the main interactions between cholinium chloride and urea would occur between the chloride and the protons at the amine groups. As they get replaced by
methyl groups (Fig. S4, ESI†), cholinium chloride loses any energetic advantage to interact with the urea family and presents positive deviations from ideality. A similar behaviour was observed by us in previous work, where cholinium chloride presented positive deviations from ideality when mixed with salts not able to act as hydrogen bond donors.7

Raman spectroscopy provides further confirmation that the chloride anion prefers to interact with urea’s N–H moieties and that the interaction becomes less favourable as the N–H are blocked by methylation. The strength of the N–H ·· ·Cl− interactions can be inferred from the frequency of the NH stretching modes as depicted in Fig. 6, where the upper panels show the sum Raman spectra of the pure liquid urea-family component and the beta polymorph of cholinium chloride while the lower panels display the spectra of the real eutectic mixtures in the liquid state. The broad spectral profiles above 3100 cm−1 result from a sum of contributions including cholinium’s O–H stretching mode and urea’s and alkylurea’s N–H symmetric and asymmetric stretching modes plus combination/overtone modes.54–56 A peak deconvolution procedure was applied to estimate the maxima of the symmetric N–H stretching contributions, highlighted in blue. As a general trend, the symmetric N–H stretch appears at lower wave-numbers in the spectra of the eutectic mixtures relative to the spectra of the pure components. In the latter, the N–H stretch contributions stem only from N–H ·· ·O=C contacts while the spectra of the real eutectic mixtures feature both N–H ·· ·O=C and N–H ·· ·Cl− interactions (since N–H ·· ·O–H contacts are expected to be negligible31). Being so, a possible explanation for the decrease in the N–H frequency is that upon forming the eutectic mixture a portion of the N–H ·· ·O=C contacts are disrupted by the formation of stronger N–H ·· ·Cl− contacts.

The difference in frequency between the pure and mixture spectra can then be taken as a measure of the strength of N–H ·· ·Cl− contacts in the different eutectic systems, with the urea-based eutectic showing the strongest deviation, while the 1,3-dimethylurea system features the lowest frequency shift. Such a tendency is evident in the plot of the N–H stretching frequency shift against the N–H ·· ·Cl− energy of interaction (estimated by DFT as described in the Experimental section), depicted in Fig. 7. This supports the idea that for eutectic mixtures based on cholinium chloride and urea, methylurea or 1,3-dimethylurea, the melting point depression is mostly ruled by the strength of the N–H ·· ·Cl− interactions, which become weaker as the degree of methylation increases.

Finally, the system cholinium chloride/1,1-dimethylurea is analysed. Fig. 8 shows a comparison between the systems cholinium chloride + urea, 1,3-dimethylurea or 1,1-dimethylurea.
Cholinium chloride presents positive deviations from ideality when mixed with 1,1-dimethylurea, as expressed by its activity coefficients, which are slightly higher than for the system cholinium chloride/1,3-dimethylurea, as shown in Fig. 8c. This suggests that fully blocking an amine group of urea (1,1-dimethylurea, see Fig. S5 for a visual aid, ESI†), instead of partially blocking its two amine groups (1,3-dimethylurea), further undermines the ability of cholinium chloride to interact with urea. This can be explained by a preference of the chloride anion to interact with the two bottom protons of the urea molecule, in a similar organisation to the head-to-tail arrangement of urea in its pure form, since these protons establish a continuum positive area between them 57 as opposed to two positive surfaces separated by a more neutral charged zone.

Contrary to the trend of cholinium chloride, Fig. 8d reveals that 1,1-dimethylurea presents negative deviations from ideality, ...
which are larger than those presented by 1,3-dimethylurea. This was not initially expected, since the increase of the cholinium chloride activity coefficient was expected to correlate with an increase of the 1,1-dimethylurea activity coefficient,
following the trend observed for the previous systems studied in this section.

The computed atomic charges for urea, methylurea, 1,3-dimethylurea, and 1,1-dimethylurea previously depicted in Fig. 4 allow for the understanding of electrostatic differences induced in urea by different methylation patterns. The addition of methyl groups results only in local effects on the molecular electrostatic field. The charges on nitrogen show the largest oscillations, from $-0.64$ from unfunctionalized urea, to around $-0.50$ for one methyl group to $-0.37$ for two methyl groups. The remaining charges show little variation. The oxygen charges remain mostly unchanged with a value near $-0.45$ and the central carbon atom and the four HBD protons show similar positive charges from $+0.34$ to $+0.36$. This indicates that the carbonyl group should not play a determinant role in the changes observed by methylation, as already suggested by the thiourea results. In addition, the electrostatic part of the hydrogen bonding should be quite similar for all protons.

In order to further understand the variation of N–H⋯Cl$^-$ contacts a Non-Covalent Interaction (NCI) analysis was performed, which, based on subtle differences of the electronic density, graphically reveals the different intermolecular interactions.\textsuperscript{58,59} These results are reported in Fig. 9.

In the NCI analysis, larger binding interactions are found as larger green (dispersive) and blue (HB) NCI volumes. First, the fact that thiourea and urea show very similar NCI with the chloride anion confirms, once more, that a very similar N–H⋯Cl$^-$ interaction is the reason why the activity coefficients for both molecules, when mixed with ChCl, are similar. The other volumes show only subtle differences, with the exception of 1,1-dimethylurea, which, since it has only one HBD proton available, shows a clearly reduced NCI volume and consequently should have the largest energy penalty. This was confirmed by the interaction energies, which were found to show a similar trend to the depressions of the melting point. Namely, excluding thiourea, the most stable pair was urea, followed by methylurea, with a slightly superior interaction energy, 1,3-dimethylurea (+5 kJ mol$^{-1}$) and 1,1-dimethylurea (+16 kJ mol$^{-1}$). This extended correlation further supports the importance of the N–H⋯Cl$^-$ interactions in the melting point depressions.

Another clarification to the unexpected behaviour of 1,1-dimethylurea can be found looking at its pure component interactions. As mentioned above, the preferred type of interaction of urea in its pure crystalline state takes place between the oxygen of the carbonyl group and the two “trans”-protons, one from each amine group, in what is known as a head-to-tail interaction.\textsuperscript{57} In the liquid phase, the head-to-tail motif is likely to remain predominant, as evidenced in the Raman spectrum shown in Fig. 7a, where N–H stretching modes stemming from head-to-tail contacts (3300–3400 cm$^{-1}$) appear significantly more intense than those arising from the alternative centrosymmetric dimer arrangement (3150–3250 cm$^{-1}$).\textsuperscript{38} By fully blocking one of the amine groups as in 1,1-dimethylurea this interaction is no longer possible, making this compound more prone to interact with the chloride anion, leading to unexpected large negative deviations from ideality. This interpretation can be corroborated and quantified using the vaporisation enthalpy of the urea-family, which, considering there is no association in the vapour phase, is an indicator of how strong the pure component interactions are in the liquid state. This value can be estimated by subtracting the melting enthalpy of the compound from its sublimation enthalpy. The results reported in Table 2 clearly show that the hypothetical liquid state of 1,1-dimethylurea is less energetically favourable when compared with the other ureas.

Conclusions

In this work, the solid–liquid equilibrium phase diagrams of cholinium chloride mixtures with urea and four modified ureas were experimentally measured. The second component of these systems was carefully chosen to represent a small modification in the structure of urea, corresponding to the selective blockage of its interaction centres. The results obtained were correlated against density functional theory-based computer models, which allowed us to conclude that the main interactions between urea and cholinium chloride are the hydrogen bonding between the chloride anion and two protons of urea, one from each amine group.

This work sheds light on the nature of the negative deviations from ideality presented by urea in cholinium chloride/urea mixtures, experimentally confirming the results of neutron diffraction by some authors\textsuperscript{31,32} regarding the dominant interactions in this system. It is still not clear, however, how the

Table 2 Melting and sublimation enthalpies for some of the urea-family compounds studied in this work, along with the resulting vaporization enthalpy, calculated from the difference between them

<table>
<thead>
<tr>
<th>Compound</th>
<th>$\Delta_{m}h$ [kJ mol$^{-1}$]</th>
<th>$\Delta_{v}h$ [kJ mol$^{-1}$]</th>
<th>$\Delta_{s}h$ [kJ mol$^{-1}$]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Urea</td>
<td>14.6$^{40}$</td>
<td>94.8$^{60}$</td>
<td>80.0</td>
</tr>
<tr>
<td>Methylurea</td>
<td>12.5$^{41}$</td>
<td>96.9$^{60}$</td>
<td>84.4</td>
</tr>
<tr>
<td>1,1-Dimethylurea</td>
<td>29.11$^{42}$</td>
<td>94.3$^{60}$</td>
<td>65.6</td>
</tr>
<tr>
<td>1,3-Dimethylurea</td>
<td>13.62$^{43}$</td>
<td>87.6$^{60}$</td>
<td>74.0</td>
</tr>
</tbody>
</table>

![Fig. 9](image_url)
second component can be tailored to induce negative deviations from ideality in cholinium chloride. This would be very useful since it would enhance the melting temperature depression effect and is under investigation in our group.

Conflicts of interest

There are no conflicts to declare.

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