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Liquefying Compounds by Forming Deep Eutectic Solvents: A Case-Study for Organic Acids and Alcohols

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Abstract

The criterion to distinguish a simple eutectic mixture from a deep eutectic solvent (DES) lies in the deviations to thermodynamic ideality presented by the components in the system. In this work, the current knowledge of the molecular interactions in types III and V DES is explored to liquefy a set of three fatty acids and three fatty alcohols, here used as model compounds for carboxyl and hydroxyl containing solid compounds.

This work shows that thymol, a stronger than usual hydrogen bond donor, is able to form deep eutectic solvents of type V with the fatty alcohols studied. This is particularly interesting since these DES formed are hydrophobic. Regarding type III DES, the results suggest that the prototypical DES hydrogen bond acceptor, cholinium chloride, is unable to induce negative deviations to ideality in the model molecules studied. By substituting choline with tetramethylammonium chloride it is shown that the choline hydroxyl group is responsible for the difficulty in forming choline-based deep eutectic solvents and that its absence induces strong negative deviations to ideality in the alkylammonium side. Finally, it is demonstrated that tetrabutylammonium chloride acts as a chloride donning agent, causing significant negative deviations to ideality in both fatty acids and alcohols, and leading to the formation of deep eutectic solvents of type III.

1. Introduction

Deep eutectic solvents (DES) are a novel class of solvents^{1,2} that emerged from the seminal works of Abbott and co-workers,^{2,3} where it was reported that binary mixtures of cholinium chloride and urea or a series of carboxylic acids (malonic acid, phenylpropanoid acid or phenylacetic acid) are liquid at room temperature. Putting it simply, and despite some questionable exceptions also presented by Abbott et al.^{4,5} using liquid components, eutectic solvents (ES) could be defined as liquid mixtures prepared from pure solid compounds. The liquid phase of ES arises due to the melting point depression of the system, which depends on the melting temperature and enthalpy of its components as well as their intermolecular interactions. The greater the negative deviations to thermodynamic ideality of both components, the greater the melting temperature depression, with the prefix *deep* here attributed to systems showing negative deviations to ideality.⁶

Deep eutectic solvents have found applications in a variety of scientific areas, such as extraction and separation,⁷⁻¹² reaction chemistry,¹³⁻¹⁶ analytical chemistry,¹⁷ biodiesel synthesis^{18,19} and electrochemistry.²⁰ Interestingly, cholinium chloride-based DES are by far the most studied in the literature,^{1-3,6-21} even though cholinium chloride typically displays a thermodynamically ideal behavior in these systems.^{6,22} Its attractiveness as a DES component stems from its low toxicity,²³ high biodegradability²⁴ and low melting enthalpy (4.3 kJ/mol),²⁵ causing it to present a steep solid-liquid equilibrium line and, thus, a sharp melting temperature depression effect.

The non-ideality of a liquid mixture arises from the establishment of intermolecular interactions of different strength than the interactions the compounds establish with themselves in their pure liquid phases. Since negative deviations to ideality are the basis for classifying an ES as *deep*, our research group has been focused on understanding the interactions between DES components, aiming at rationalizing the design of true deep eutectic solvents.

Recently we proposed a new type of DES (type V) based on non-ionic substances.²⁶ Despite the structural similarity of its components, the system thymol/menthol presents large negative deviations to ideality and is an example of this new class of DES. Due to the electron withdrawing effect of the aromatic resonance of thymol, its hydroxyl group becomes more positive, with the oxygen becoming a poorer hydrogen bond acceptor and its proton becoming a better hydrogen bond donor when compared with the hydrogen bonding capability of typical alcohols (such as menthol). Hence, thymol and menthol establish a stronger hydrogen bond

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3 network than those present in the liquid phase of the pure components. We have also identified
4 mixtures in which tetrabutylammonium chloride ($[N_{4,4,4,4}]Cl$) behaves as a chloride donning
5 agent,²⁷ analogous to the mechanism of formation of type I DES. An example is the system
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7 $[N_{1,1,1,1}]Cl/[N_{4,4,4,4}]Cl$ with $[N_{1,1,1,1}]Cl$ standing for tetramethylammonium chloride, which
8 presents large negative deviations to ideality²⁷ due to a synergetic share of the available
9 chloride ions, in which the coordination number of chloride anions around the $[N_{1,1,1,1}]^+$ cation
10 increases (when compared to its pure liquid phase) and decreases around the $[N_{4,4,4,4}]^+$ cation
11 (in comparison to its pure liquid phase). This phenomenon, as will be shown here, can be
12 explored to design novel type III deep eutectic solvents.
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19 The objective of this work is to show that the current understanding of the molecular
20 mechanisms behind DES formation, with emphasis on DES of types III and V, allows for an
21 informed choice of *liquefying agents*, i.e. compounds that induce negative deviations to ideality
22 in other solid compounds, originating a liquid mixture. Fatty acids (and carboxylic acids in
23 general) are both hydrogen bond donors and acceptors, with the possibility of forming dimers
24 in their pure liquid phases. Likewise, fatty alcohols are also both good hydrogen bond donors
25 and acceptors. Their simple structure, minimizing the possibility of other specific interactions
26 that could create difficulties in the interpretation of the results, and their melting points above
27 room temperature, makes them attractive model compounds for this study.
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35 In this work, a set of experimental solid-liquid equilibrium data taken from the literature,
36 complemented with new data herein measured, is used to rationalize the liquefaction of fatty
37 acids or alcohols using the deep eutectic solvent approach. Fatty acids (tetradecanoic acid,
38 hexadecanoic acid and octadecanoic acid) and fatty alcohols (tetradecan-1-ol, hexadecan-1-ol
39 and octadecan-1-ol) are here used as model compounds to be liquefied and menthol, thymol,
40 cholinium chloride (ChCl), $[N_{1,1,1,1}]Cl$ or $[N_{4,4,4,4}]Cl$ as compounds to induce the liquefaction.
41 The identification of the dominant molecular interactions present on these systems will help
42 understand the behavior of a plethora of other systems, and will pave the way to the design of
43 novel DES that are not just ideal mixtures of compounds forming, in some cases, eutectic
44 solvents.
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2. Methods

2.1 Chemicals

The substances experimentally used in this work are reported in Table 1, along with their supplier and purity (mass %). Due to their hygroscopicity, $[N_{1,1,1,1}]Cl$ and $[N_{4,4,4,4}]Cl$ were dried before use by stirring in vacuum (0.1 Pa) at room temperature (298 K), for at least 72 hours. The remaining compounds were used as received. The water content of the substances was measured using a Metrohm 831 Karl Fischer coulometer, with the analyte Hydranal Coulomat AG from Riedel-de-Haen.

Table 1. List of substances experimentally used in this work along with their CAS number, supplier, mass purity and water content.

Substance	CAS Number	Supplier	Purity	Water Content /ppm
$[N_{1,1,1,1}]Cl$	75-57-0	Sigma-Aldrich	97 %	2957
$[N_{4,4,4,4}]Cl$	1112-67-0	Sigma-Aldrich	97 %	5758
Thymol	89-83-8	TCI	>99 %	185
(-)-Menthol	2216-51-5	Acros Organics	99.5 %	79
Tetradecanoic Acid	544-63-8	Acros Organics	99 %	0
Hexadecanoic Acid	57-10-3	Aldrich	99 %	13
Octadecanoic Acid	57-11-4	Sigma-Aldrich	99 %	12
Tetradecan-1-ol	112-72-1	Alfa Aesar	98 %	66
Hexadecan-1-ol	36653-82-4	Alfa Aesar	98 %	6
Octadecan-1-ol	112-92-5	Sigma	99 %	18

2.2 Solid-Liquid Phase Diagram Measurement

To study liquefaction by forming type V DES, the solid-liquid equilibrium (SLE) phase diagram of the binary systems composed of thymol or (-)-menthol (menthol) and tetradecan-1-ol (tetradecanol), hexadecan-1-ol (hexadecanol) or octadecan-1-ol (octadecanol) were experimentally measured. Moreover, to study liquefaction by forming type III DES, the SLE phase diagram of the binary systems composed of $[N_{1,1,1,1}]Cl$ and tetradecanol, hexadecanol or octadecanol, and composed of $[N_{4,4,4,4}]Cl$ and tetradecanoic acid, hexadecanoic acid, octadecanoic acid, tetradecanol, hexadecanol or octadecanol were also measured in this work.

For each binary system, mixtures were prepared covering the entire composition range. The mass of each component was weighted using an analytical balance (model ALS 220-4N from Kern) with a readability of 0.1 mg. After preparation, the samples were heated under stirring

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3 until fusion and recrystallized to obtain a homogeneous solid mixture. Then, the samples were
4 crushed with a mortar and pestle and glass capillaries were filled with the resulting powder.
5 The melting temperature of the sample in each glass capillary, taken as the temperature at which
6 complete fusion is observed, was measured using the melting point device model M-565 from
7 Büchi, with a temperature resolution of 0.1 K and a temperature gradient of 0.1 K/min. This
8 procedure was repeated at least three times. For systems composed of either [N_{1,1,1,1}]Cl or
9 [N_{4,4,4,4}]Cl, the sample and capillary preparation were performed inside a dry-argon glove box
10 to prevent atmospheric humidity contamination.
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14 In the cases where the melting temperature of the mixture was below room temperature, the
15 recrystallization technique was not possible. Thus, approximately 5 mg of these samples
16 weighed using a micro-analytical balance AD6 (PerkinElmer, USA, precision = 2 × 10⁻⁶ g),
17 were tightly sealed in aluminum pans and their melting temperature measured using differential
18 scanning calorimetry (DSC). A Hitachi DSC7000X model, working at atmospheric pressure,
19 coupled with a cooling system, was used to perform the measurements. The cooling and heating
20 rates were respectively 5 K/min and 2 K/min and each temperature was taken as the peak
21 temperature upon heating. The DSC equipment was previously calibrated with several
22 standards (heptane, octane, decane, 4-nitrotoluene, naphthalene, benzoic acid, diphenylacetic
23 acid, indium, tin, caffeine, lead, zinc, potassium nitrate, water, and anthracene) with purities
24 higher than 99 wt%.
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37 The experimental methodologies described above have already been checked for the SLE phase
38 diagram measurement of DES by comparing the measured data with other sources and
39 methods^{28,29} as well as applying thermodynamic consistency tests.²⁵ The foundations of these
40 methodologies are extensively described by Hefter and Tomkins.³⁰
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45 2.3 Modelling

46 Solid-liquid equilibrium,³¹⁻³³ when no solid-solid transitions are present, the components
47 crystallize as pure substances, assuming also constant heat capacity change upon melting, and
48 neglecting the difference between melting and triple point properties, is described by:
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$$52 \ln(x_i \cdot \gamma_i) = \frac{\Delta_m h_i}{R} \cdot \left(\frac{1}{T_{m,i}} - \frac{1}{T} \right) + \frac{\Delta_m C_{P,i}}{R} \cdot \left(\frac{T_{m,i}}{T} - \ln \frac{T_{m,i}}{T} - 1 \right) \quad (1)$$

53 where x_i is the mole fraction of component i , γ_i is its activity coefficient in the liquid phase, Δ_m
54 h_i is its melting enthalpy, $T_{m,i}$ is its melting temperature, $\Delta_m C_{P,i}$ is its heat capacity change
55 upon melting, R is the ideal gas constant and T is the absolute temperature of the system. Given
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3 the small temperature difference between the melting temperature of the mixtures herein
4 studied and those of the pure compounds the heat capacity term is negligible,^{31,32} with
5 Equation 1 simplifying to:
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$$\ln(x_i \cdot \gamma_i) = \frac{\Delta_m h_i}{R} \cdot \left(\frac{1}{T_{m,i}} - \frac{1}{T} \right) \quad (2)$$

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12 The activity coefficients of the components in a eutectic solvent can be calculated from the
13 experimental solid-liquid phase diagram by rearranging Equation 2:
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$$\gamma_i = \frac{\exp\left[\frac{\Delta_m h_i}{R} \cdot \left(\frac{1}{T_{m,i}} - \frac{1}{T}\right)\right]}{x_i} \quad (3)$$

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21 In this work, the activity coefficients were calculated using Equation 3 and the melting
22 properties listed in Table S1. Even for the cases where the activity coefficients were already
23 reported in the literature, they were herein recalculated using the melting properties of Table S1
24 to perform consistent comparisons between all systems.
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30 **3. Discussion**

31 **3.1 Liquefying acids and alcohols by forming type V DES**

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33 Due to the non-ionic nature of type V DES, the interactions established between their
34 components are simpler to understand and discuss, and our analysis starts with these systems.
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36 The solid-liquid phase diagram of menthol and the model fatty acids, taken from the
37 literature,³⁴ are depicted in Figure 1, along with the corresponding activity coefficients
38 estimated from Eq. (3) as described in section 2.3.
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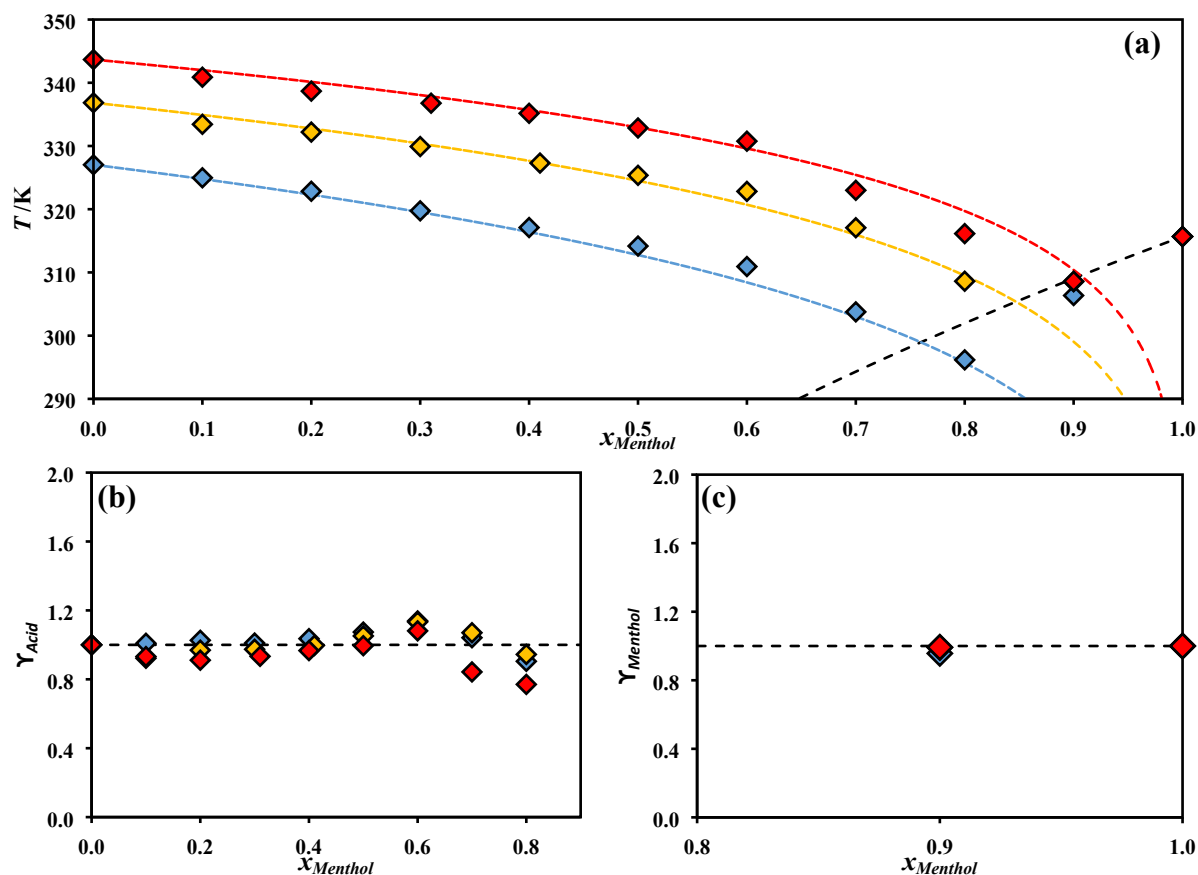


Figure 1. Solid-liquid phase diagrams³⁴ (a) and corresponding activity coefficients of fatty acids (b) and menthol (c), for the systems composed of menthol and tetradecanoic acid (\blacklozenge), hexadecanoic acid (\blacklozenge) or octadecanoic acid (\blacklozenge). Dashed lines represent the ideal liquidus lines (--- menthol, --- tetradecanoic acid, --- hexadecanoic acid and --- octadecanoic acid). Dotted lines represent thermodynamic ideality ($\gamma=1$).

Figure 1 shows that both fatty acids and menthol behave ideally when mixed, with only mild deviations to ideality presented by the acids for high menthol concentration. Therefore, the molecular interactions established in the mixture are similar and of the same strength to those present in the pure component liquid phases (hydrogen bonding between hydroxyl groups of similar nature). As such, and even though the system menthol/tetradecanoic acid presents a eutectic temperature around room temperature (ca. 296 K), menthol fails at forming deep eutectic solvents with the organic acids studied.

The solid-liquid phase diagram of menthol and the model fatty alcohols were measured in this work, as described in section 2.2, and are depicted in Figure 2, along with the corresponding activity coefficients.

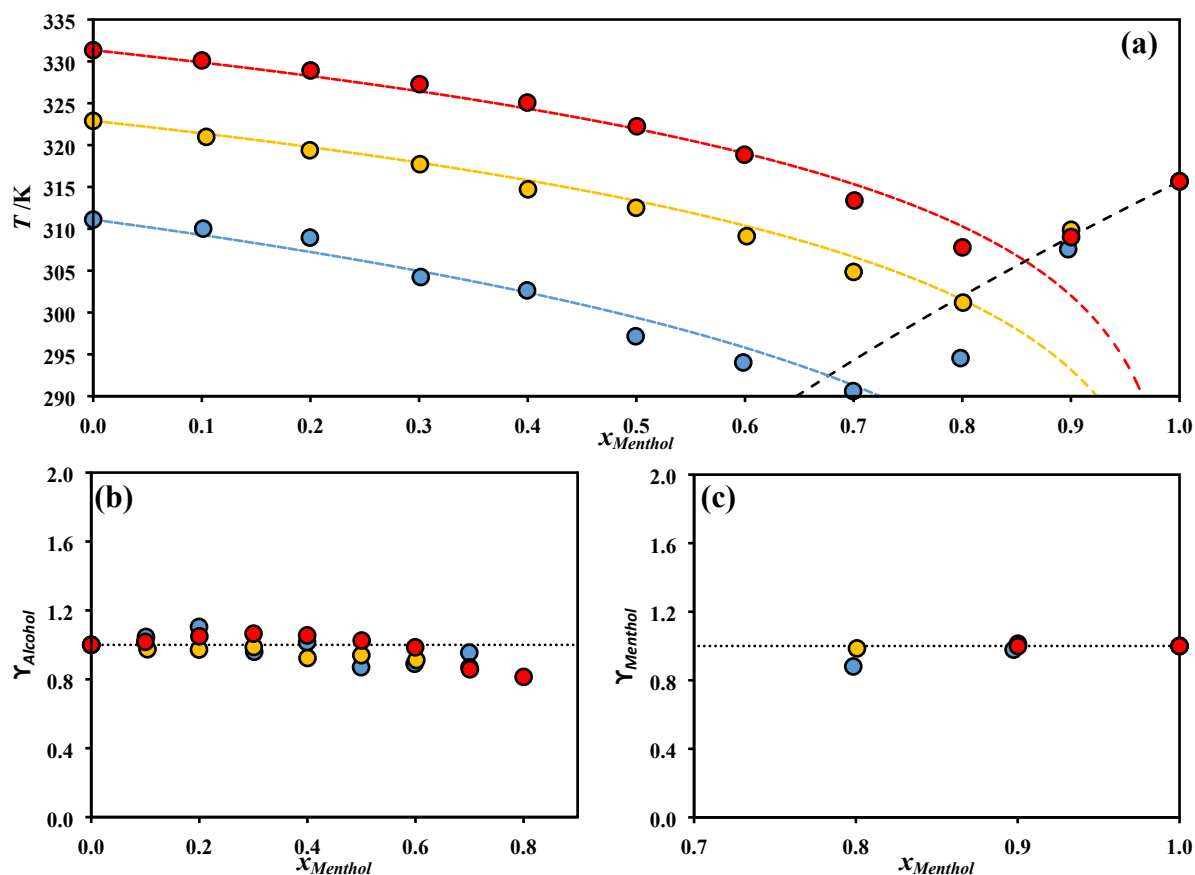


Figure 2. Solid-liquid phase diagrams (a) and corresponding activity coefficients of fatty alcohols (b) and menthol (c), for the systems composed of menthol and tetradecanol (\bullet), hexadecanol (\bullet) or octadecanol (\bullet). Dashed lines represent the ideal liquidus lines (--- menthol, --- tetradecanol, --- hexadecanol and --- octadecanol). Dotted lines represent thermodynamic ideality ($\gamma=1$).

Similar to the menthol/fatty acid systems depicted in Figure 1, Figure 2 reveals that mixtures of menthol and fatty alcohols behave ideally. Again, the molecular interactions between menthol and the fatty alcohols present in these systems are comparable in strength to those found in the liquid phases of the pure components.

The results reported in Figures 1 and 2 reassert that the existence of hydrogen bonds between the two components of a mixture does not guarantee the presence of negative deviations to ideality and, thus, the formation of deep eutectic solvents. For this to happen, the interactions established between the components must be stronger than those present in the pure compound liquid phases. Within the framework of type V DES, these results show that aliphatic alcohols such as menthol, being hydrogen bond donors and acceptors of average strength able to establish strong interactions with themselves in their pure liquid phases, are not useful to liquefy other substances with hydroxyl and/or carboxyl groups by causing deviations to

thermodynamic ideality. This conclusion is in line to our previous work, where it was shown that menthol was not responsible for the significant negative deviations to ideality present in the thymol/menthol system.²⁶ It is also in good agreement, and helps explaining, previous reports of SLE phase diagrams of menthol forming ideal liquid mixtures with various other compounds^{34–39}.

Having discussed the effect of menthol on organic acids and alcohols, the focus now shifts to thymol (see Figure S2 for both structures). Unlike menthol, thymol has a hydrogen bond capability unbalance: it is a stronger hydrogen bond donor and a weaker hydrogen bond acceptor than other hydroxyl-containing non-phenolic compounds.²⁶ The solid-liquid phase diagram of thymol and the model fatty acids, taken from the literature,³⁴ are depicted in Figure 3, along with the corresponding activity coefficients.

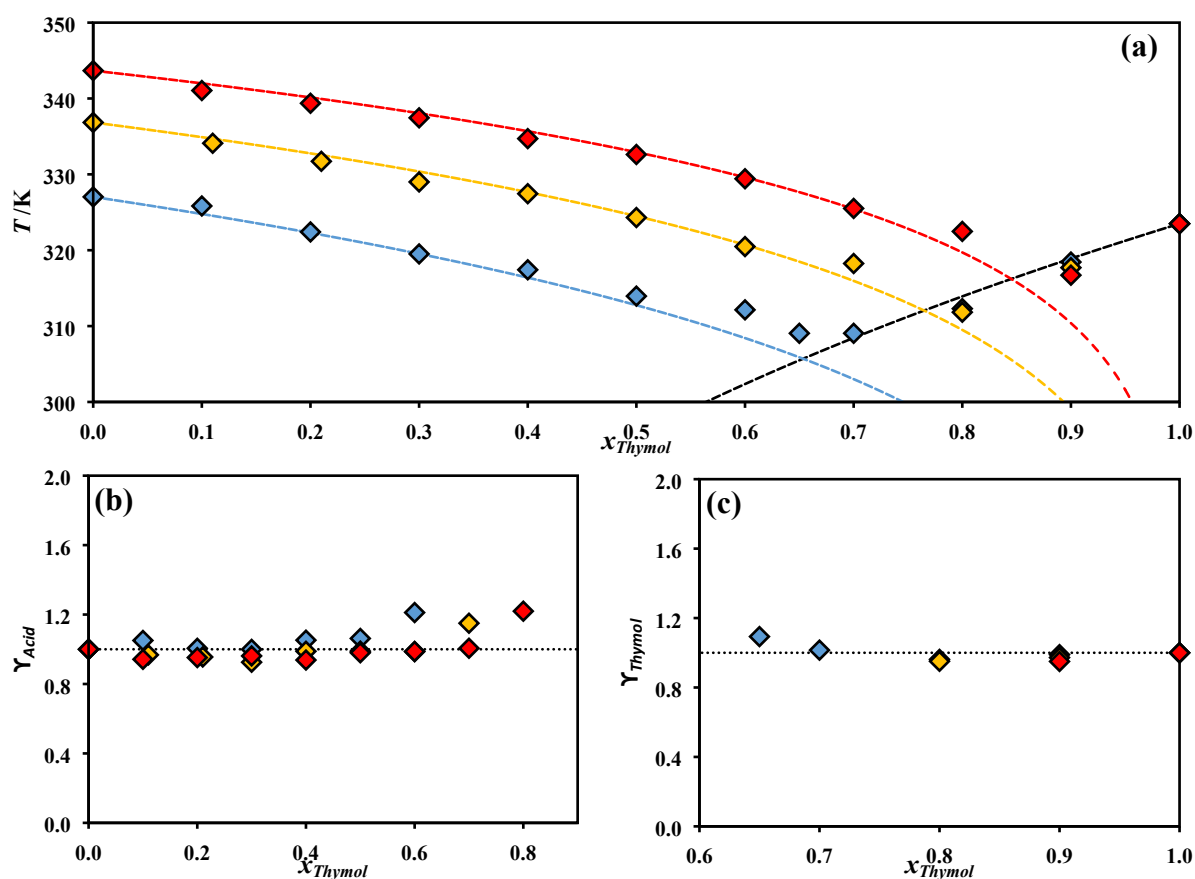


Figure 3. Solid-liquid phase diagrams³⁴ (a) and corresponding activity coefficients of fatty acids (b) and thymol (c), for the systems composed of thymol and tetradecanoic acid (\blacklozenge), hexadecanoic acid (\blacklozenge) or octadecanoic acid (\blacklozenge). Dashed lines represent the ideal liquidus lines (--- thymol, --- tetradecanoic acid, --- hexadecanoic acid and --- octadecanoic acid). Dotted lines represent thermodynamic ideality ($\gamma=1$).

Despite its hydrogen bond donating strength, Figure 3 shows that thymol cannot induce negative deviations to ideality in the model fatty acids and, thus, cannot form deep eutectic solvents with these substances. In fact, and unlike menthol/acid systems, the fatty acids studied present small positive deviations to ideality for high thymol concentration. Since the principal structural difference between thymol and menthol is that the former is a stronger hydrogen bond donor and a weaker hydrogen bond acceptor, the different activity coefficient results from Figures 1b and 3b suggest that fatty acids are acting as the hydrogen bond donors and thymol/menthol as the hydrogen bond acceptors. Menthol being a better hydrogen bond acceptor than thymol²⁶ induces the small negative deviations seen in Figure 1b.

The solid-liquid phase diagram of thymol with the model fatty alcohols were measured in this work, as described in section 2.2, and are depicted in Figure 4, together with the corresponding activity coefficients.

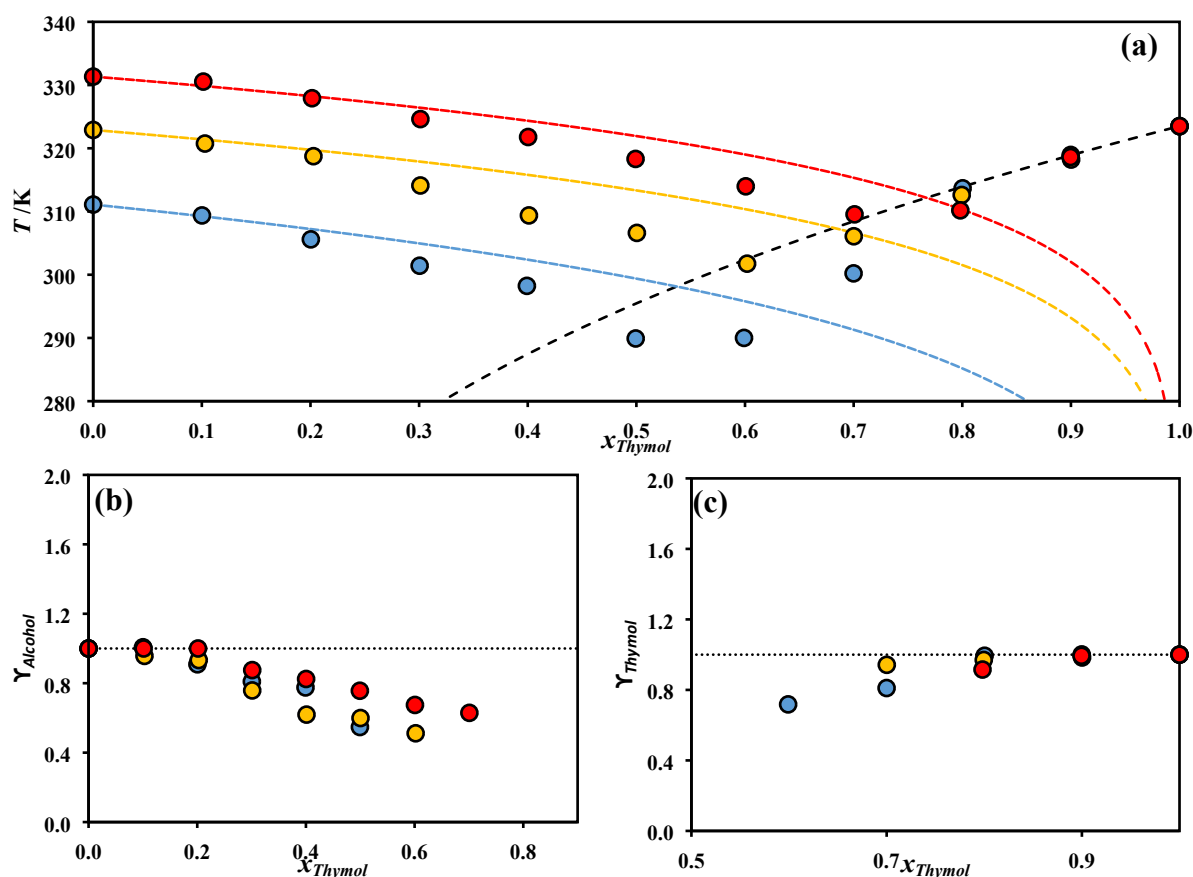


Figure 4. Solid-liquid phase diagrams (a) and corresponding activity coefficients of fatty alcohols (b) and thymol (c), for the systems composed of thymol and tetradecanol (\bullet), hexadecanol (\circ) or octadecanol (\bullet). Dashed lines represent the ideal liquidus lines (--- thymol, --- tetradecanol, --- hexadecanol and --- for octadecanol). Dotted lines represent thermodynamic ideality ($\gamma=1$).

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3 The phase diagrams depicted in Figure 4 demonstrate the usefulness of thymol as a liquefying
4 agent by forming DES of type V. Because thymol is a stronger hydrogen bond donor than the
5 fatty alcohols it can induce large deviations to ideality on the aliphatic alcohols by forming
6 hydrogen bonds that are stronger than those present in the pure compounds, as also shown in a
7 previous work for the mixture thymol-menthol,²⁶ leading to deep eutectic solvents of type V.
8 Hydrophobic deep eutectic solvents, proposed for the first time by van Osch et al.,⁴⁰ expand
9 the applicability of DES since most previously reported in the literature are choline-based,
10 hence hydrophilic. In this respect, the new DES reported in Figure 4 are important since they
11 are formed by hydrophobic substances, echoing the importance of type V DES, and the
12 informed selection of the compounds of the eutectic solvent mixture, in the preparation of novel
13 hydrophobic solvents.
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23 **3.2 Liquefying alcohols and acids by forming type III DES**

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25 After showing how the type V DES framework can be used in the liquefaction of organic
26 alcohols, yet failing, with the liquefying agents studied, at forming deep eutectic solvents with
27 the fatty acids, type III DES are now discussed. We start by analyzing the prototypical DES-
28 forming component: cholinium chloride. It has been previously proposed that the
29 thermodynamic behavior of the prototypical cholinium chloride/urea DES is governed by a
30 hydrogen bond between two protons of urea and a chloride anion,⁴¹ which is a behavior
31 reminiscent of the chloride transfer in the type I DES-analogous ionic liquid mixtures
32 mentioned in the introduction section.²⁷
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39 To investigate if cholinium chloride is able to form DES with fatty acids, Figure 5 depicts their
40 binary solid-liquid phase diagrams,²⁸ and the corresponding activity coefficients.
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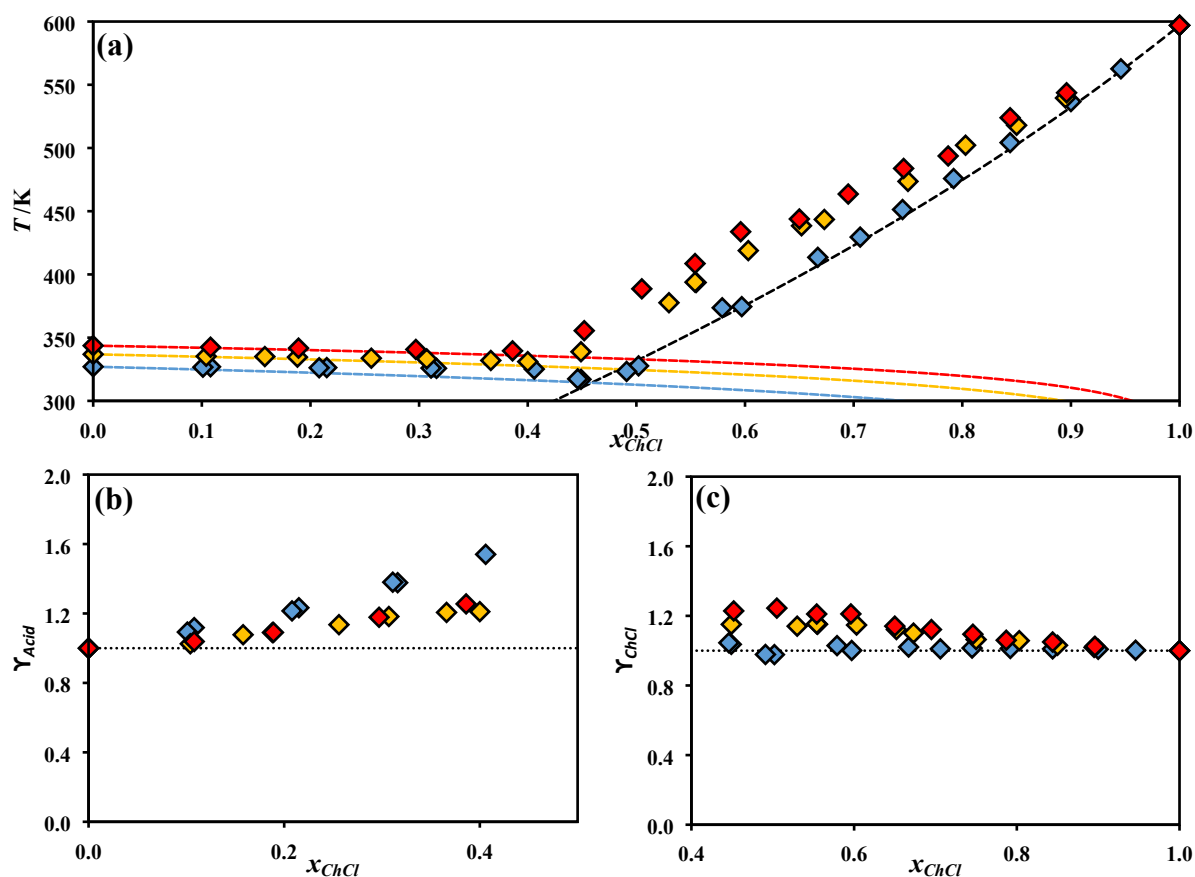


Figure 5. Solid-liquid phase diagrams²⁸ (a) and corresponding activity coefficients of fatty acids (b) and ChCl (c), for the systems composed of ChCl and tetradecanoic acid (\diamond), hexadecanoic acid (\diamond) or octadecanoic acid (\diamond). Dashed lines represent the ideal liquidus lines (--- ChCl, --- tetradecanoic acid, --- hexadecanoic acid and --- octadecanoic acid). Dotted lines represent thermodynamic ideality ($\gamma=1$).

Excluding the ChCl/tetradecanoic acid system in which cholinium chloride behaves ideally, Figure 5 reveals that both cholinium chloride and fatty acids present significant positive deviations to ideality when mixed. To investigate if, unlike the ChCl/acid systems, cholinium chloride is able to form DES with fatty alcohols, Figure 6 depicts the solid-liquid phase diagrams of these systems, taken from the literature,²⁸ along with the corresponding activity coefficients.

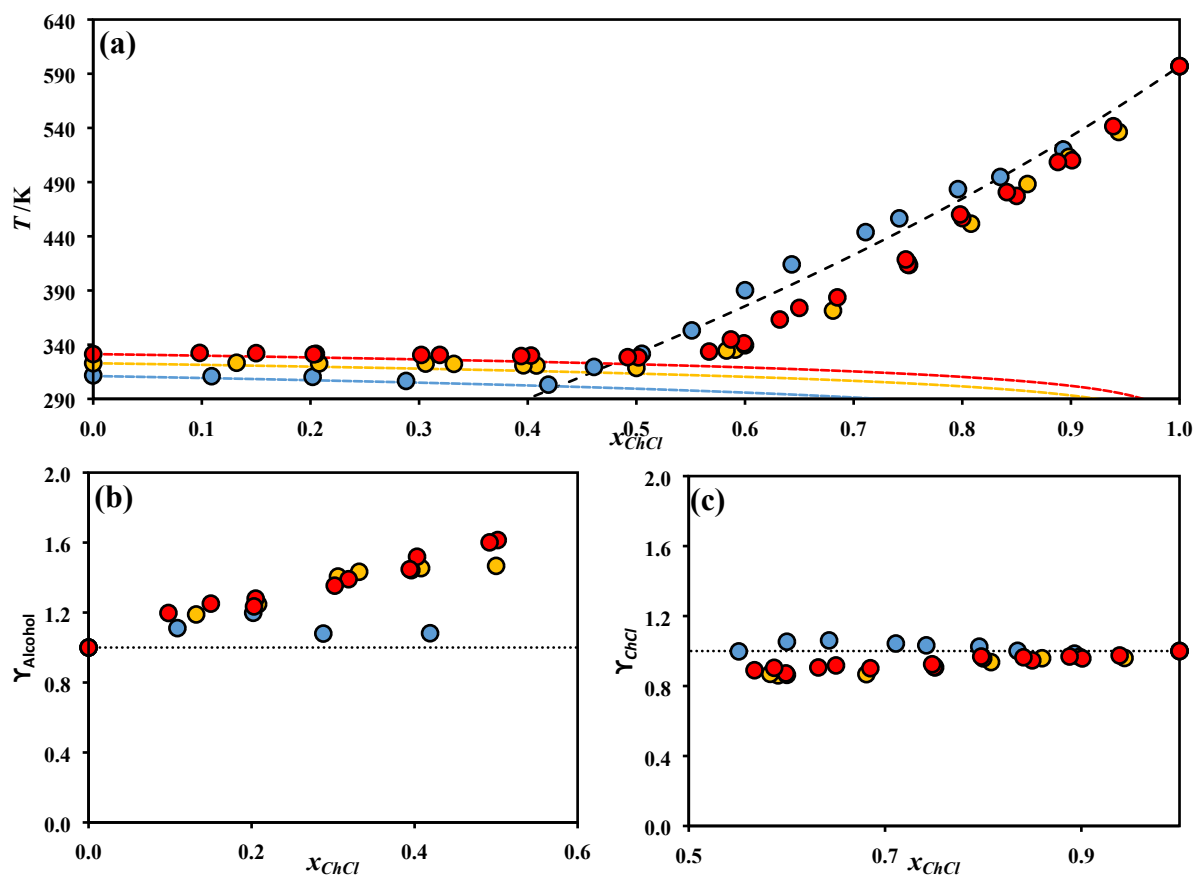


Figure 6. Solid-liquid phase diagrams²⁸ (a) and corresponding activity coefficients of fatty alcohols (b) and ChCl (c), for the systems composed of ChCl and tetradecanol (●), hexadecanol (●) or octadecanol (●). Dashed lines represent the ideal liquidus lines (--- ChCl, --- tetradecanol, --- hexadecanol and --- octadecanol). Dotted lines represent thermodynamic ideality ($\gamma=1$).

The results reported in Figure 6 are similar to those shown in Figure 5, with the fatty alcohols presenting large positive deviations to ideality. However, cholinium chloride now shows either an ideal behavior or small negative deviations to ideality. These results confirm that cholinium chloride does not present a particular ability to establish new, and stronger than in pure, intermolecular interactions in mixtures.

Since these results show that cholinium chloride cannot form deep eutectic solvents with the fatty acids or alcohols studied, $[N_{1,1,1,1}]\text{Cl}$ will be now considered. This is an interesting compound to discuss since it is structurally similar to cholinium chloride, with the sole difference being the hydroxyethyl group of the cholinium cation replaced by a methyl group. Thus, by studying $[N_{1,1,1,1}]\text{Cl}$ -based systems the effect of the choline hydroxyl group can be probed whilst checking for DES formation by charge transfer. Figure 7 depicts the solid-liquid

phase diagrams of $[N_{1,1,1,1}]Cl$ and the model fatty acids, taken from the literature,²⁹ along with the corresponding activity coefficients.

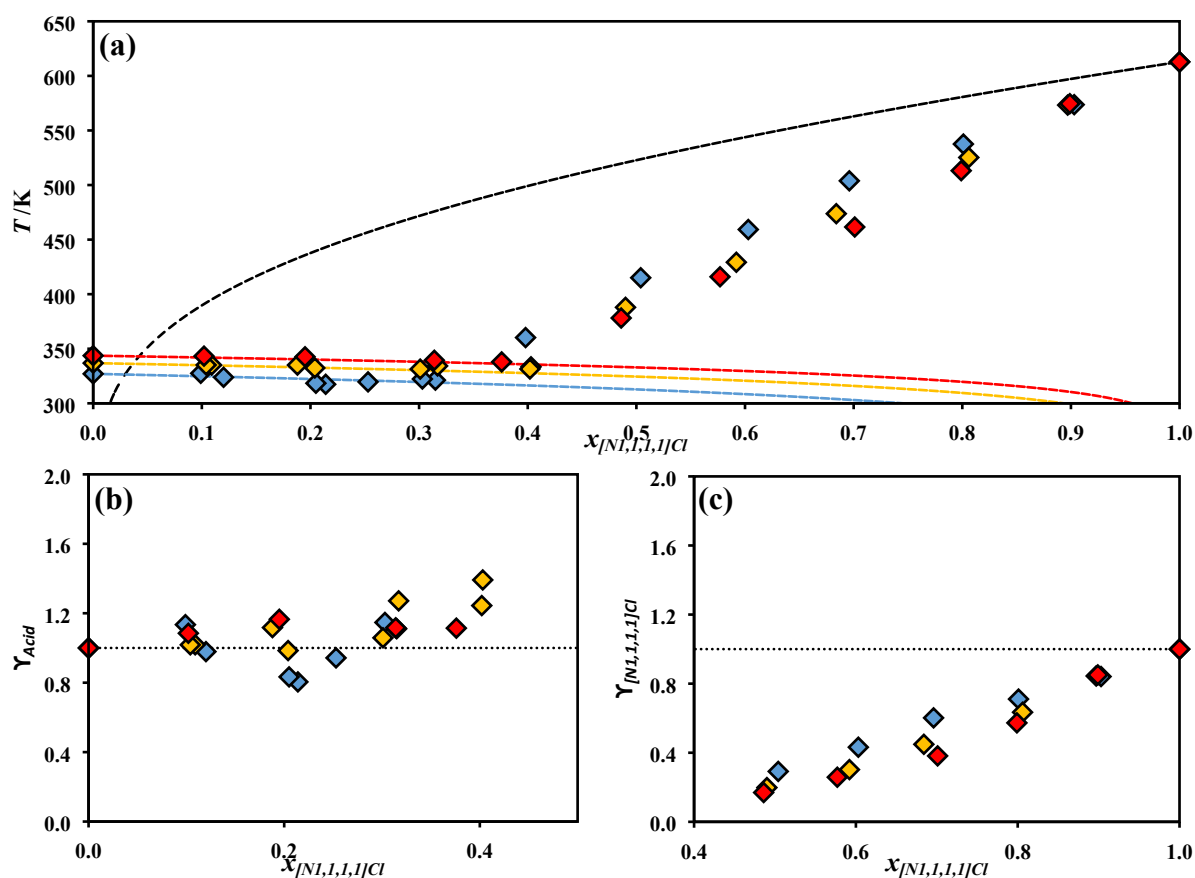


Figure 7. Solid-liquid phase diagrams²⁹ (a) and corresponding activity coefficients of fatty acids (b) and $[N_{1,1,1,1}]Cl$ (c), for the systems composed of $[N_{1,1,1,1}]Cl$ and tetradecanoic acid (\blacklozenge), hexadecanoic acid (\blacklozenge) or octadecanoic acid (\blacklozenge). Dashed lines represent the ideal liquidus lines (--- $[N_{1,1,1,1}]Cl$, --- tetradecanoic acid, --- hexadecanoic acid and --- octadecanoic acid). Dotted lines represent thermodynamic ideality ($\gamma=1$).

The impact of the hydroxyl group present in cholinium chloride is quite significant, as the comparison of Figures 5 and 7 reveal. While cholinium chloride presented large positive deviations to ideality when mixed with the model acids, $[N_{1,1,1,1}]Cl$ presents the opposite behavior, *i.e.* large negative deviations to ideality. Despite this change in behavior of the liquefying agent, the model fatty acids still show positive deviations to ideality identical to those found in the ChCl-based systems. This suggests that the hydroxyl group endows cholinium chloride with stronger interactions in its pure phase, namely hydrogen bonding between the chloride anion and the highly positive hydroxyl group. By removing this effect in $[N_{1,1,1,1}]Cl$, the hydrogen bonding with the fatty acids becomes more favorable than its interactions in the pure liquid phase, resulting in the negative deviations seen.

To check if the absence of hydroxyl group in $[N_{1,1,1,1}]Cl$ also affects the fatty alcohol systems, Figure 8 depicts the solid-liquid phase equilibrium of $[N_{1,1,1,1}]Cl$ and the fatty alcohols, along with the corresponding activity coefficients.

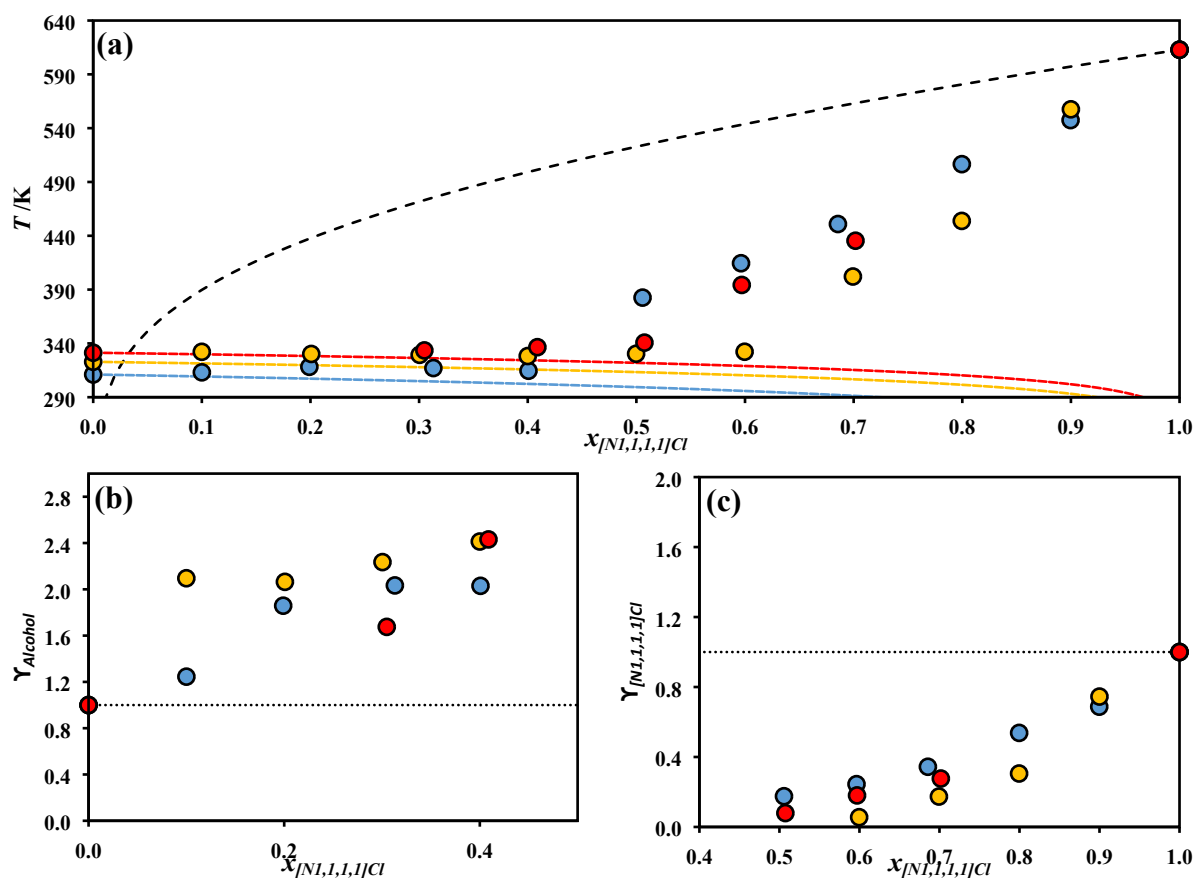


Figure 8. Solid-liquid phase diagrams (a) and corresponding activity coefficients of fatty alcohols (b) and $[N_{1,1,1,1}]Cl$ (c), for the systems composed of $[N_{1,1,1,1}]Cl$ and tetradecanol (\bullet), hexadecanol (\bullet) or octadecanol (\bullet). Dashed lines represent the ideal liquidus lines (--- $[N_{1,1,1,1}]Cl$, --- tetradecanol, --- hexadecanol and --- octadecanol). Dotted lines represent thermodynamic ideality ($\gamma=1$).

As observed for the systems of $[N_{1,1,1,1}]Cl$ and fatty acids, the absence of a hydroxyl group in the tetramethylammonium cation results in very large deviations to ideality. Working under the assumption that the chloride anion is a stronger hydrogen bond acceptor than the hydroxyl groups of the fatty acids and alcohols studied (this will be later proven using $[N_{4,4,4,4}]Cl$, $COOH--Cl$ or $COH--Cl$ interactions should lead to negative deviations to ideality presented by the acids/alcohols. On the contrary, the acids/alcohols in these systems show positive deviations to ideality suggesting that $COOH--Cl$ or $COH--Cl$ interactions are not occurring.

Having seen that cholinium chloride is unable to form deep eutectic solvents with the model acids and alcohols studied due to its hydroxyl group and that $[N_{1,1,1,1}]Cl$ produces highly

asymmetrical systems with regards to activity coefficients, $[N_{4,4,4,4}]Cl$ is now discussed. $[N_{4,4,4,4}]Cl$ has been shown to behave as a chloride donating agent, with its cation being able to form apolar domains depleted in chloride anions.²⁷ In this work, the solid-liquid phase diagrams of the systems composed by $[N_{4,4,4,4}]Cl$ and the model fatty acids were measured and are depicted in Figure 9.

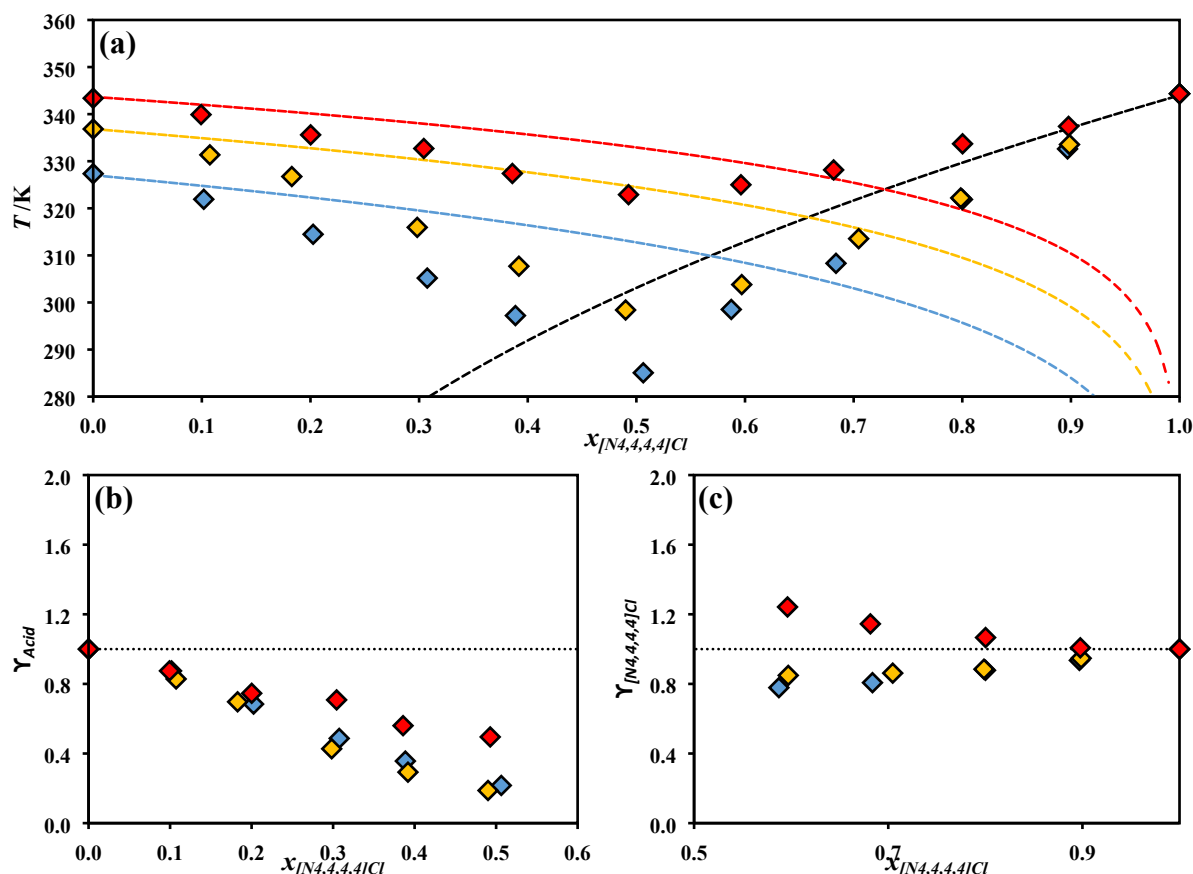


Figure 9. Solid-liquid phase diagrams (a) and corresponding activity coefficients of fatty acids (b) and $[N_{4,4,4,4}]Cl$ (c), for the systems composed of $[N_{4,4,4,4}]Cl$ and tetradecanoic acid (\blacklozenge), hexadecanoic acid (\blacklozenge) or octadecanoic acid (\blacklozenge). Dashed lines represent the ideal liquidus lines (--- $[N_{4,4,4,4}]Cl$, --- tetradecanoic acid, --- hexadecanoic acid and --- octadecanoic acid). Dotted lines represent thermodynamic ideality ($\gamma=1$).

The results reported in Figure 9 are extremely interesting. On the contrary to what was observed for all systems previously discussed, for the first time negative deviations to ideality are present in the carboxylic acids. The $[N_{4,4,4,4}]Cl$, however, presents either a near-ideal behavior or even weak positive deviations to ideality. This behavior is reminiscent of that previously observed in the $[N_{1,1,1,1}]Cl/[N_{4,4,4,4}]Cl$ system,²⁷ supporting the idea that chloride anions are being transferred from the $[N_{4,4,4,4}]$ cation to the carboxylic acid head, engaging in stronger hydrogen bonds than those present in the pure acid liquid phases.

Bearing in mind the success of $[N_{4,4,4,4}]Cl$ in inducing negative deviations to ideality in the model acids studied, the alcohol-based systems are now analyzed. The solid-liquid phase diagrams of these systems were measured in this work and are depicted in Figure 10, along with the corresponding activity coefficients. As can be seen, the behavior of $[N_{4,4,4,4}]Cl$ mixed with fatty alcohols is quite similar to that observed with fatty acids. Again, $[N_{4,4,4,4}]Cl$ is able to induce negative deviations to ideality in the alcohols, while retaining a near-ideal behavior or showing weak positive deviations to ideality.

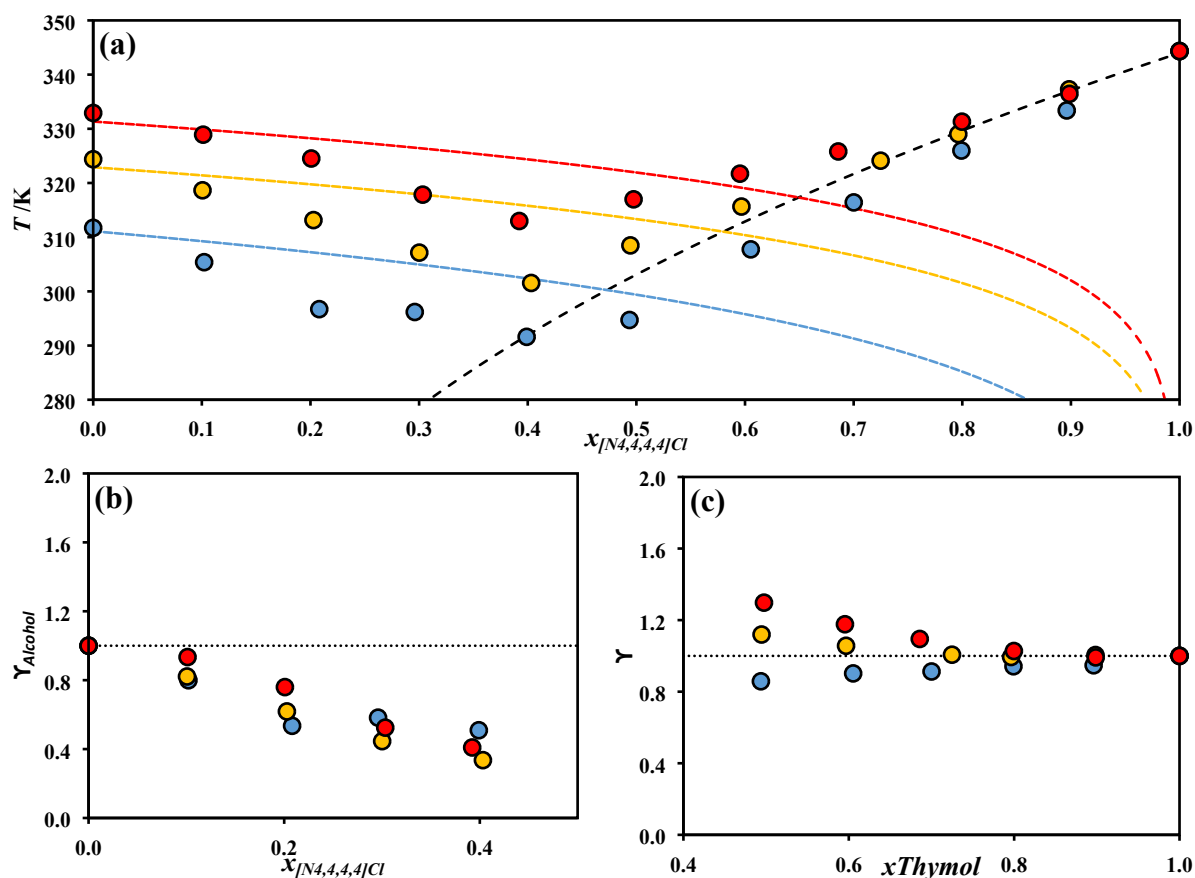


Figure 10. Solid-liquid phase diagrams (a) and corresponding activity coefficients of fatty alcohols (b) and $[N_{4,4,4,4}]Cl$ (c), for the systems composed of $[N_{4,4,4,4}]Cl$ and tetradecanol (\bullet), hexadecanol (\circ) or octadecanol (\circ). Dashed lines represent the ideal liquidus lines (--- $[N_{4,4,4,4}]Cl$, --- tetradecanol, --- hexadecanol and --- octadecanol). Dotted lines represent thermodynamic ideality ($\gamma=1$).

The thermodynamic behavior of the $[N_{4,4,4,4}]Cl$ -based systems and the $[N_{1,1,1,1}]Cl$ -based systems is essentially opposite. $[N_{1,1,1,1}]Cl$ formed highly asymmetrical deep eutectic solvents with the model fatty acids/alcohols studied, presenting severe negative deviations to ideality itself but failing to induce them in the other component. Contrary to this behavior, $[N_{4,4,4,4}]Cl$ induces strong negative deviations to ideality in the organic molecules but shows itself weak

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3 positive deviations to ideality. Such a drastic difference in thermodynamic behavior suggests
4 that the molecular interactions are fundamentally different in those two sets of systems.
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7 Based on the results of our previous work,²⁷ given the weak interaction between the cation and
8 anion in $[N_{4,4,4,4}]Cl$, it is expected that this compound donates chloride anions to the organic
9 acids and alcohols, establishing hydrogen bonds that are stronger than those present in the
10 acids/alcohols pure liquid phases. Since the cation of $[N_{1,1,1,1}]Cl$ is much more densely charged,
11 it is unable to similarly donate chloride anions, functioning as the receiving end of chloride
12 anions in the $[N_{1,1,1,1}]Cl/[N_{4,4,4,4}]Cl$ system as previously shown.²⁷ Considering that $[N_{1,1,1,1}]Cl$
13 shows severe negative deviations to ideality and considering that the acids and alcohols
14 presented severe positive deviations to ideality (which indicates that the new interactions
15 established in the mixture are weaker than those present in their pure liquid phase), it is
16 concluded that the cation of $[N_{1,1,1,1}]Cl$ is interacting with the negative zones of the organic
17 molecules (lone pairs of the oxygen atoms), further stabilizing itself by withdrawing additional
18 negative charge but preventing the hydrogen bonding between the organic molecules, leading
19 to the asymmetrical deviations to ideality seen in these systems.
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33 4. Conclusions

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35 Throughout this work, the feasibility of forming deep eutectic solvents based on fatty alcohols
36 or acids was studied using different liquefying agents. This work reiterates the importance of
37 studying the solid-liquid phase diagrams of *deep* eutectic solvents in order to understand their
38 molecular interactions, the origin of their deviations to ideality and rationalizing the search for
39 new DES.
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44 Regarding type V DES, it was shown that thymol induces negative deviations to ideality in
45 organic alcohols, allowing the design of new hydrophobic deep eutectic solvents. Furthermore,
46 it was shown that menthol forms ideal mixtures with the fatty acids or alcohols studied, as
47 expected due to the similarity of the interactions established in these mixtures to those
48 established in the pure components liquid phases, confirming that the existence of hydrogen
49 bonding networks in eutectic solvents does not guarantee negative deviations to ideality and,
50 thus, does not mean that the eutectic solvent is *deep*.
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57 In terms of type III DES, it was shown that cholinium chloride is unable to form deep eutectic
58 solvents with either fatty acids or alcohols. Using $[N_{1,1,1,1}]Cl$, it was shown that the ideal
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3 behavior of cholinium chloride in the systems studied stems from its hydroxyethyl group,
4 which allows for strong cation-anion interactions in its pure, hypothetical, liquid phase. Thus,
5 although due to its low melting enthalpy and *green* properties, cholinium chloride can be seen
6 as a generic eutectic solvent-forming compound, it is however unable to form deep eutectic
7 solvents with organic acids or alcohols. Finally, it was shown that [N_{4,4,4,4}]Cl forms deep
8 eutectic solvents with the fatty acids and alcohols studied.
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11 Within the *liquefying-agent* concept, this work demonstrates the usefulness of thymol (type V
12 DES) and [N_{4,4,4,4}]Cl (type III DES) in forming deep eutectic solvents with organic acids and
13 alcohols, leading to the liquefaction of these substances. More specifically, liquid mixtures
14 below room temperature (298 K) were obtained for tetradecanol when mixed with either
15 thymol or [N_{4,4,4,4}]Cl, and for tetradecanoic and hexadecanoic acids when mixed with
16 [N_{4,4,4,4}]Cl.
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19 An understanding of the interactions between liquefying agents and organic substances allows
20 for the design of novel DES of importance to green chemistry. This work highlights some of
21 these interactions and liquefying agents that can induce them, but further investigation is still
22 required to compile a full body of these favorable interactions for classes of compounds other
23 than acids or alcohols.
24

25 **Supporting Information**

26 The Supporting Information is available free of charge at: <https://pubs.acs.org/doi/???/???>.

27 SI includes melting properties, detailed experimental results and chemical structures. Table S1:
28 melting properties. Tables S2: detailed experimental results. Figures S1-S2: chemical
29 structures.
30
31

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