Studying of two choline chloride’s deep eutectic solvents in their aqueous mixtures

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Abstract: Deep Eutectic Solvents (DESs) are an interesting category of solvents that recently getting more attention due to their unique properties. DESs proved their reliability in many applications which made them overcoming the traditional solvents. In this work, two deep eutectic solvents choline chloride: urea and choline chloride: glycerol having molar ratio of 1:2 respectively were prepared and characterized. Different water content, 0, 3, 5, 10, 25, 50 and 75 wt. %, were added and the effect was studied. Cyclic voltammetry, viscosity, density and thermogravimetric analysis have been investigated and reported in this research. The viscosity was studied as a function of temperature at normal and high temperatures and as a function of shear rate at low temperature. For our best awareness, the non-Newtonian behavior of the above two DESs is first reported in this work. The shear rate effected the viscosity of the mixtures at low temperatures; thereby, by increasing the applied shear rate, the viscosity of the mixtures decreased at a fixed temperature.

I. INTRODUCTION

Deep Eutectic Solvents (DESs) can be defined as a mixture of a hydrogen bond donor (HBD) and a hydrogen bond accepter (HBA) which can associate with each other to form a eutectic mixture. The final mixture melting point is lower than both of its raw materials separately.

In general, deep eutectic solvents have a very huge depression in their freezing points. They are liquids at temperatures below than 150°C. Most of them are liquids under 70°C, with some even at room temperature. In most cases, deep eutectic solvent can be prepared by mixing one of the quaternary ammonium salts as a hydrogen bond donor with a metal salt as a hydrogen bond acceptor [1].

Andrew Abbott and co-workers introduced the term of “Deep Eutectic solvent” in 2003. They got this name from the idea of synthesizing liquids from a mixture of two high melting point solid raw materials. Multiple researchers subsequently published papers using the term “Deep Eutectic Solvents” [2]. Figure 1 is showing the growth in publications’ number contained “Deep Eutectic Solvents” in their titles, key words, and/or abstracts. The data were obtained using Scopus database and they were updated up to the mid of July 2016.

A very common example of a DES is the mixture of choline chloride and urea. Choline chloride has a melting point of 302 ºC and urea’s melting point is 133 ºC. The mixture melting point is 12 ºC for a molar ratio 1:2, respectively [3].

DESs usually are synthesized by raising the temperature of two solid raw materials while mixing them together. The final high temperature attained depends on the nature of the raw materials, but commonly using higher temperatures and good mixing leads to faster melting [4].

Using Ionic Liquids, many researches have produced nanoparticles from gold, silver, platinum, copper, manganese, titanium, chrome, molybdenum, tungsten, iron, ruthenium, osmium, and their various components such as oxides, halides, etc. By choosing which different anions and cations make up the Ionic Liquids, specific products can be manufactured [5-14].

Deep eutectic solvents have the advantages over ionic liquids of being water-tolerant, biodegradable, less expensive, easier to make, and safer to use. Many deep eutectic solvents are non-toxic compared to ionic liquids [2].

In this work, two common deep eutectic solvents mixtures, Choline Chloride: Urea and Choline Chloride: Glycerol, have been prepared and analyzed in details. Different water contents, 0, 3, 5, 10, 25, 50 and 75 wt. %, were added to both mixtures and the effect was investigated. Cyclic voltammetry, viscosity, density and thermogravimetric analysis properties were tested in this work. The viscosity was studied as a function of shear rate (at low temperatures) and temperature (at higher temperatures). For our best knowledge, this paper is...
the first paper reporting the non-Newtonian behavior for those two DESs at low temperature and study the shear rate effect on the viscosity.

II. EXPERIMENTAL SECTION

2-1 Materials
Choline chloride (≥ 99%) was purchased from Chem-Impex Int’l Inc, USA. Urea (99.4%) and Glycerol (99.5%) were obtained by Fisher Scientific Company, USA. The commercial catalyst of Pt/C (19.7% Pt over Vulcan XC-72ETEK) was obtained by De Nora North America. Deionized water (DI) was obtained by Direct-Q3 UV Millipore device.

2-2 DESs’ Preparation:
Choline chloride was dried at 65 °C and under vacuum for 24 hr to get rid of any possible moisture. The first DES mixture was prepared by mixing the dried choline chloride with urea in molar ratio of 1:2, respectively. But choline chloride with glycerol DES, 1:2 molar ratios, respectively, was prepared by heating glycerol first up to 80 °C and choline chloride was added later. The mixtures were heated at 80 °C and stirred until homogeneous colorless liquids were formed. Finally, DESs mixtures were dried at 80 °C and left under vacuum overnight in order to eliminate any possible moisture before starting characterizations.

2-3 Characterization
Cyclic Voltammetry (CV): The electrochemical test CV, for both dried mixtures, was performed at room temperature using electrochemical workstation model CHI651E. Three electrodes cell with glassy carbon (GC) electrode of 0.196 cm² working electrode was used for this test. The reference electrode was Ag/AgCl and the counter electrode was platinum wire. Alumina slurry of 1µm size was used to polish the GC and it was washed ultrasonically by DI water. The test was occurred using commercial catalyst of ETEK Pt/C. 1mg of Pt/C catalyst was ultrasonically dispersed in 1ml of ethanol for one hour to prepare ink for the test. 10µl of this ink was dropped on the polished GC electrode and dried at room temperature. 10 µl of 0.05% Nafion solution was dropped on the dried catalyst to bind the catalyst on the electrode surface and was left it to dry at room temperature. The potential range of this test was -1V to 1V.

Viscosity: The viscosity was measured using Brook Field DV III Ultra Program Rheometer cone/plate and the attached spindle was #42. The temperature range was from 10 to 80 °C. The viscosity was studied as a function of shear rate at low temperatures and the non-Newtonian behavior was reported. Different water contents, 0%, 3%, 5%, 10%, 25%, 50% and 75% weight ratio, were added to the prepared DESs and the viscosity was measured for the new mixtures. The temperature was controlled using external water circulator ISOTEMP.

Density: The density was measured using 2 ml Gay-Lussac vial at room temperature (22 °C) for the mentioned mixtures above.

Thermogravimetric analysis (TGA): TGA was tested using TA Instruments-Waters LLC device model Q500 and the analyzing temperature range was from10 to 800 °C. The temperature rising rate was 10 °C/min using a platinum pan of 0.379 gm. Nitrogen was used as balance and sample gas and purged in rate of 40 ml/min.

III. RESULT AND DISCUSSION

3-1 Cyclic Voltammetry
In the present study, cyclic voltammetry was tested to figure out the electrochemical behavior for both mixtures as shown in fig2. The dashed curve represented chcl: glycerol DES while the solid one represented chcl: urea mixture. For chcl: urea DES, it can be observed from the forward scan that there are two oxidation waves started at -500 mV and -400 mV. These waves could be attributed to low H⁺ ions concentrations in the DES, H₂O molecules dissociation, impurities which represented by protons, and/or the high hygroscopic of choline chloride. On the other side, no oxidation waves observed for chcl: glycerol DES. It can be seen that chcl: glycerol mixture started to oxidize sharply at potential higher than 700 mV and reached its maximum current density at 920 mV. Whereas, chcl: urea oxidation current increased slightly after the potential of 700 mV which could be attributed to evolution of chlorine gas [15] and reached its maximum current density at 1000 mV.

Couple of reduction peaks are clearly observed at potentials of 320 and -100 mV for chcl: glycerol mixture which could be attributed to water reduction [16]. No reduction peaks observed for chcl: urea DES and the reduction current started to increase sharply at potential lower than -500mV.

3-2 Viscosity
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The viscosity was measured for both mixtures at different water content (0, 3, 5, 10, 25, 50 and 75 wt. %) and different temperatures (10 to 80 °C). At temperatures below 20 °C, these mixtures behaved as Non-Newtonian fluids and the viscosity depended on the shear rate values. That means the viscosity did not have a specific value at a certain temperature because it depended on the shear rate. While at temperatures (above 20 °C) they behaved as Newtonian fluids.

Figure 3 shows the viscosity behavior of chcl: Glycerol of 0 % water content at 10, 12.5, 15 and 17.5 ºC. At 20 ºC and above, this DES started to behave as a Newtonian fluid. However, figure 4 shows the same phenomena for chcl: Urea of 0 % water content mixture at 10, 15 and 20 °C. From the two figures, it can be noted that viscosity decreased with shear rate increasing. This difference was big at low shear rates for the same DES and it started to decrease with shear rate increasing.

From both figures, the shear rate effect decreased with temperature increasing and chcl: urea mixture was more viscous than chcl: glycerol mixture at the same temperature and shear rate. Higher than the mentioned temperatures, 17.5 ºC for chcl: glycerol and 20 ºC for chcl: urea, both DESs mixtures started to behave as Newtonian fluids.

Figure 5 shows the temperature effect on viscosity when the two mixtures behave as Newtonian fluids. It is clear that the viscosity decreased with temperature increasing. Also, whenever the temperature increased, the changing in viscosity value decreased and the curve almost became linear.

3-3 Water content effect on viscosity

The effect of water content on the viscosity was studied in different percentages for the same range of temperatures above. For both mixtures, the water content percentages were 3, 5, 10, 25, 50, and 75 wt.%. For chcl: glycerol mixture, at 3% water content, figure 6 is showing the shear rate effect on viscosity at 10 and 12.5 ºC. While figure 7 is showing the shear rate effect at 10 °C for 10 and 25 % water content. However, for higher temperatures and water contents, the mixture behaved as Newtonian fluid. Whereas, chcl: urea mixture behaved as a Newtonian fluid when water was added in whole temperatures range. So that, the water content effected chcl: urea mixture more than chcl: glycerol mixture.

Figures 8 and 9 show the effect of temperature on viscosity in different water content for both mixtures. Both of them show Newtonian behavior whenever the temperature increases.

The temperature dependence on viscosity was studied for both mixtures in different water content and the data was fitted to Arrhenius equation [17].

\[
\ln(V) = \ln(V_0) + \frac{E_v}{R} \frac{1}{T}
\]  

Where \(V_0\) is a constant, \(R\) is the universal gas constant, \(T\) is the temperature in (K), and \(E_v\) is the activation energy. Table 1 shows the results when equation 1 was applied. It can be noted that the activation energy decreased by increasing the water content for both mixtures, except when the water’s percentage exceeded the DES’s one for chcl: glycerol mixture.

3-4 Density

The density was studied at room temperature (22 ºC) as reported in table 2. It can be noted from that table whenever the water percent increased, the density decreased.

3-5 Thermogravimetric analysis

TGA was studied at range of Room temperature up to 800 ºC. The results show that whenever the water percentage increased, the decomposition rate increased as shown in figures 10 and 11. The reason is because the water started to evaporate before the deep eutectic solvent. Both DESs mixtures at 0% water content started to decompose around 200 ºC.

IV. Conclusion

Binary deep eutectic solvent mixtures consisted of choline: urea and choline chloride: glycerol have synthesized. Physical and electrochemical properties (cyclic voltammetry, viscosity as a function of temperature, shear rate and water content, density and thermogravimetric analysis) for these mixtures were investigated. The findings have implied different properties for both composites. Cyclic voltammetry over Pt/C catalyst showed that choline chloride: urea composite has higher current density than choline chloride: glycerol mixture which implies the better conductivity for the first mixture. The viscosity results showed that choline chloride: urea mixture has higher viscosity than other and both mixtures behaved as a Non-Newtonian fluid at temperatures lower than 20°C and the viscosity became shear rate dependent. Choline chloride: urea mixture was more effected by water content than choline chloride: glycerol composite. The density findings showed that choline chloride: urea composite has higher density and it decreased with increasing the water content. Finally, TGA results showed that both mixtures started to decompose at 200°C and the decomposition rate increased with water content increasing.
ACKNOWLEDGMENTS:
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REFERENCES
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Fig. 1. Number of Publications contained “Deep Eutectic Solvents” in their titles, keywords, and/or abstracts obtained by Scopus database.

Figure 2: Cyclic voltammetry profiles obtained for chcl: urea and chcl: Glycerol DESs using Pt/C catalyst pasted on GC electrode at scan rate: 50 mV s⁻¹.

Figure 3: Viscosity as a function of shear rate for DES (1:2, chcl: glycerol) 0% water content.
Figure 4: Viscosity as a function of shear rate for DES (1:2, chcl: urea) 0% water content.

Figure 5: Viscosity as a function of temperature for choline chcl: glycerol and chcl: urea mixtures, 0% water content.

Figure 6: Viscosity as a function of shear rate for 3% water content in 10 and 12.5 °C for chcl: glycerol mixture.
Figure 7: Viscosity as a function of shear rate for 10 and 25 % water content in 10 and °C for chel: glycerol mixture.

Figure 8: Viscosity as a function of temperature in different water contents for Choline Chloride: Glycerol mixture

Figure 9: Viscosity as a function of temperature in different water contents for Choline Chloride: Urea mixture
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Figure 10: TGA analysis for Choline Chloride: Glycerol mixture in different water contents.

Figure 11: TGA analysis for Choline Chloride: Urea mixture in different water contents.

Table 1. Activation energy relation with viscosity based on Arrhenius equation

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<th>Choline Chloride: Glycerol</th>
<th>Choline Chloride: Urea</th>
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<td>No.</td>
<td>Water %</td>
<td>Ln (V˳)</td>
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Table 2. Densities of DESs at different water content

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<th>Choline Chloride: Urea</th>
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