Dielectric relaxation of nonaqueous ionic liquid microemulsions: polarization, microstructure, and phase transition†

Cancan Zhang, Zhen Zhen, Liyan Ma and Kongshuang Zhao*

Two nonaqueous ionic liquid (IL) microemulsions (toluene/TX-100/[bmim][PF_6] and [bmim][BF_4]/TX-100/benzene) were studied by dielectric spectroscopy covering a wide frequency range (40 Hz to 110 MHz). A unique relaxation was observed in the radio frequency (RF) range. By methodically analyzing the dependence of relaxation parameters on the ILs content, the microstructures of the microemulsions were identified. Additionally, based on the interfacial polarization theory and Einstein equation, the mechanism of the relaxation caused by the fluctuation of IL anions along the TX-100 PEO chain was confirmed, what’s more, according to the dependence of the dc conductivity of the microemulsions on IL concentration, it was concluded that the hydrophilicity of the IL in the nonaqueous IL microemulsions may play a crucial role in the electrical conduction mechanism: our analysis results suggest that a dynamic percolation process occurs in the toluene/TX-100/[bmim][PF_6] system in which IL is hydrophobic, while a static percolation happens in benzene/TX-100/[bmim][BF_4] where IL is hydrophilic. The otherness of relaxation time provides evidence that there is a possible coupling effect between IL and TX-100. Moreover, there are hints that all of the disparities, such as relaxation time, percolation type, ion migration rate and the size of different micro zones, may just stem from the different hydrophobicity of the two kinds of IL.

1. Introduction

Due to their unique physicochemical properties, such as environmental friendliness, low volatility, wide electrochemical window, low flammability, high thermal stability, and wide liquid range, ILs have been receiving more and more attention in recent years.1–6

Recently, many researchers have been attempting to use ILs to prepare microemulsions, since in 2004 Han et al. first found an uncommon microemulsion system where [bmim][BF_4] in the fashion of nanoscale droplets was dispersed in a continuous hydrocarbon solvent,7,8 which soon afterwards was investigated by Eastoe et al. and Sarkar et al.9,10 respectively. Therefrom, the study of nonaqueous microemulsions with ILs instead of water has become an active research area.11–17 A large number of studies have shown that the nonaqueous microemulsion systems have generous obvious advantages, for example, compared with a similar aqueous solution, the nonaqueous systems often show a large stable region in isotropic solution.17,18 In addition, due to their characteristics, ILs can provide micro environments for some special reactions. Although the solubility of polar solutes in ILs is still limited, it can be surmounted through the introduction of micelles or IL/O microemulsions which can provide a hydrocarbon chain region. In this context, it is pretty significant to study the ternary microemulsions which ILs participate in constructing.

The copious efforts of many researchers on a wide variety of nonaqueous microemulsions with diverse oil phase have brought to light their potential applicabilities in massive practical applications (such as chemical reaction, separation, nanoparticles and porous materials preparation etc.).19–26 which greatly widens the application range of the traditional microemulsions. Nevertheless, from what we know, there is still no more comparative reports about microemulsions with different ILs phase, which may make a thorough inquiry of the microstructure, especially the polarity and formation mechanism of the micro structure and provide more basic information for the application of nonaqueous IL-based microemulsions.

Dielectric spectroscopy (DS), as a powerful tool to explore the polarization and dynamics inside materials, have been early used to study the dynamic and structural properties of colloidal systems, such as emulsion and particle dispersion.7,27–29 On the other hand, percolation can not only bring considerable current interest in practical application, but provide us a very good way to judge the microstructure transition in a microemulsion.7,27,28 Most of these studies have focused on promulgating the dynamics, the percolation process and the interaction between...
the droplets. For example, Buchner et al.\textsuperscript{39} conducted the dielectric measurement on water/octane/C\textsubscript{12}E\textsubscript{8} from which they determined the dynamic percolation process induced by the temperature and obtained the percolation temperature. Asami\textsuperscript{38} studied the temperature dependence of the dielectric properties of water/toluene/TX-100 microemulsion, and gained favorable construction on the percolation and dielectric spectrum data with the interfacial polarization theory. Cametti\textsuperscript{39} measured the temperature dependent dielectric spectrum for AOT/decane/water microemulsion, and obtained the percolation temperature from the static permittivity and the DC conductivity data. They attributed the low-frequency dielectric relaxation to the polarization of surfactant head group from the water–surfactant interface. Z. Chen and R. Nozaki\textsuperscript{32} found a new relaxation in 10\textsuperscript{7} to 10\textsuperscript{8} Hz through comparing water/TX-100/[bmim][PF\textsubscript{6}] and TX-100/water mixtures with various \(\phi\textsubscript{w}\), and put it down to the hopping of IL cationic/anionic species between the anionic/cationic sites. And their studies above are about aqueous microemulsion. To the best of our knowledge, the dielectric study about the percolation of nonaqueous microemulsion and their comparisons is very little. Among them, Gao et al. used the relationship between conductivity and percolation to estimate the effect of water on the IL microemulsion.\textsuperscript{44} We recently reported the dielectric behavior of nonaqueous IL microemulsions and explained the percolation phenomena of systems combining percolation theory.\textsuperscript{44,45} And these works firstly systematically discussed the solubility and polarity of ILs, the interaction between IL and TX-100, and percolation phenomena in the B.C. phase. Meanwhile, some characteristics of soft materials in such microemulsion systems have been explained. In that way, whether other IL-based microemulsion systems also have a similar dynamic behavior and formation mechanism, especially, how about their percolation behavior? Because percolation problem is a significant criterion of the structure and phase transition, as well as an important issue in the dielectric study of microemulsions. In this study, we select the toluene/TX-100/[bmim][PF\textsubscript{6}] (Han 2005) and benzene/TX-100/[bmim][BF\textsubscript{4}] (Zheng 2007) two representative systems. Through measuring their dielectric behavior in radio frequency, their phase transition processes and percolation phenomena were discussed, and the relaxation time of the ionic fluctuation and migration, the electric conduction model and the difference of ion transport rate caused by the distinct hydrophobicity of ILs were proffered.

2. Experimental and methods

2.1 Materials and preparations of IL microemulsions

The IL [bmim][PF\textsubscript{6}] and [bmim][BF\textsubscript{4}] (purity > 99%) were both purchased from Shanghai Cheng Jie Chemical Co. Ltd., China. The content of residual chloride and water in these two ILs were both less than 800 ppm, and less than 1000 ppm respectively. TX-100 (\(\rho\)-1,1,3,3-tetramethylbutyl)phenoxypolyoxyethylene polyglycol) (reagent grade) was obtained from Amresco Chemical Inc. America. Benzene and toluene were produced by Beijing Chemical Reagent Factory, China, and were analytical reagent grade. The structural formula of TX-100 and IL [bmim][PF\textsubscript{6}], [bmim][BF\textsubscript{4}] are all shown in Fig. 1. Before used, they were dried under vacuum for 12 hours at 80 °C to remove water.

The sample compositions of dielectric measurements are shown in the isothermal triangular phase diagram,\textsuperscript{36,37} as Fig. 2, presents the measured phase points which make up the path of experiment: Fig. 2a denotes the ternary system in which the weight concentration of IL is increased to the weight ratio of IL/TX-100 3 : 2; when the weight ratio of benzene/TX-100 is fixed as 3 : 2; Fig. 2b shows the ternary system in which the weight concentration of IL is increased to the weight ratio of IL/TX-100 1 : 1, when the weight ratio of toluene/TX-100 is fixed as 3 : 1. These two paths both span the microstructure concentration range of the microemulsion monophase region, so it can be used to study the transformation of the system mesoscopic order. The points taking from Fig. 2 determines the quality ratio of the three components in the system and their specific quantity respectively needed was calculated by per sample 3 g, and weighed with the electronic analysis balance, then mixed. The prepared sample was put into a sealed glass bottle which was clean and dry. Light grey area in Fig. 2 represents IL packages oil area (O/IL) in which there are a mass of microemulsion spheres. Whereas the bright grey area delegates the oil coats ILs region (IL/O), there are a great deal of microemulsion spheres. There is a B.C. structure during the transition of the two kinds of mesoscopic structures.

2.2 Dielectric measurements

The low-frequency dielectric measurement of the two microemulsion systems were performed on a 4294A precision impedance analyzer (Agilent Technologies, made in Japan) that allows a continuous frequency measurement from 40 Hz to 110 MHz. A dielectric measurement cell with concentrically cylindrical platinum electrodes\textsuperscript{38} was employed and connected to the impedance analyzer by a 16047E spring clip fixture (Agilent Technologies, made in Japan). The original data capacitance \(C\) and conductance \(G\), which was corrected by the measured parameters such as cell constant \(C_0\), the stray capacitance \(C_s\) and the residual inductance \(L_r\) (obtained by using ethanol, water, and KCl solution) according to Schwab’s method. The permittivity \(\varepsilon\) and the conductivity \(\kappa\) were calculated according to \(\varepsilon = (C - C_s)/C_0\) and \(\kappa = G/C_0\) \((\varepsilon_0 = 8.854 \times 10^{-12} \text{ F m}^{-1})\) is
3. Results and discussion

3.1 The concentration dependence of dielectric spectrum

From Fig. 3, there is a remarkable dielectric relaxation around $10^7$ to $10^8$ Hz. It’s easy to observe that both the dielectric permittivity of vacuum) [the detailed description is provided in the part A of the ESI†].

2.3 Determination of relaxation parameters

Dielectric relaxation can be characterized by a series of dielectric parameters, which can be obtained by fitting the experimental data with the Havriliak–Negami equation.

$$
\varepsilon^\prime(\omega) = \varepsilon^\prime - j\varepsilon^\prime\prime = \varepsilon_b + \sum_i \frac{\Delta\varepsilon_i}{(1 + (j\omega\tau_i)^n)^\beta}
$$

where $\omega$ is the angular frequency, $\varepsilon_b$ is the high-frequency limit of relative permittivity, $i$ is the number of relaxation, $\Delta\varepsilon$ is the relaxation intensity, $\omega = (\kappa - \kappa_i)/\varepsilon\omega_0$, $\kappa_i$ is the low-frequency limit of conductivity, $\tau_i$ is the angular frequency, $\alpha (0 < \alpha \leq 1)$ and $\beta (0 < \beta \leq 1)$ both are parameters related to the distribution of relaxation time (the detailed description is showed in the part B of the ESI†).

3.2 Relaxation mechanism

Due to containing a hydrophilic chain composed of approximately 10 EO units, TX-100 can be bonded by water molecules through hydrogen bonding. The combination between water and TX-100 plays a crucial role in deciding phase structure, which also, in a large extent, determines the physicochemical properties of the microemulsion TX-100/water/IL. The dielectric behavior of TX-100/water mixtures has been studied by Asami in detail. An obvious relaxation was observed in the vicinity of 1 GHz, which was accredited to the bulk water and bound water. After that, the dielectric measurements of ILs microemulsions which constructed by IL and TX-100 with water or oil were also respectively performed as mentioned above. In these measurements, researchers observed another relaxation of higher radio frequency $10^7$ to $10^8$ Hz. Nevertheless, just as what have been used to interprete the relaxation of the IL/TX-100/water system, many possible relaxation mechanisms may contribute to this relaxation. There is a system the IL/TX-100/cyclohexane in our previous study, quite similar to this present work. We have formerly ascribed the relaxation found in that work to the counterion polarization at the micelle interface. In our present results, the relaxation in $10^7$ to $10^8$ Hz frequency range has been likewise observed and its relaxation intensity increase with increasing the concentration of IL [bmm][PF6] or [bmm][BF4] (see Fig. 3a and b). Besides, it can be seen from Table 1 that the changes of the relaxation parameters with concentration of ILs were segmented (details will be discussed later), which means that at least one relaxation mechanism should be responsible for this relaxation. In general, this relaxation is due to the aggregation or recombination, the tangential or radial diffusion of the interfacial ions, and the fluctuations caused by the thermal motion of the aggregates. The kinetic relaxation time of micellar is generally in 0.6 μs, and thermal motion and interfacial
polarization are respectively about 15 µs, 160 µs. These relaxation times are far greater than the observed values. Therefore, with a rough estimate, these three mechanisms didn’t mainly contribute to the observed the relaxation.

In order to explain our results, we proposed a model shown in Fig. 4. In the two systems, the OE units on the TX-100 chain have electrostatic interaction on the cations [bmim]+ of ILs, which limits the diffusion of cations. Further, due to the electrostatic attraction from the limited cations, the anions [BF4]−/[PF6]− are as well forced to distribute around the cations. Nevertheless, the movement of anions isn’t fully fettered by the weak electrostatic attraction. Therefore, anions still have a diffusion coefficient D.

Before and after the application of electric field, in order to balance the distribution, the anions fluctuate within the PEO chain length and produce the relaxation. Assuming that the fluctuation length 〈x〉 is the length of the PEO chain (see Fig. 4D), the time that this relaxation process needed can be calculated according to the Einstein displacement equation:

$$\tau = \langle x \rangle^2 / 2 D \cdot$$

(2)

For the system benzene/TX-100/[bmim][BF4], putting limit molar conductivity $\Lambda_m^0([\text{bmim}][\text{BF}_4], 298 \text{ K}) = 9.37 \text{ S m}^{-1} \text{ M}^{-1}$, $\lambda_\text{i}([\text{bmim}], 298 \text{ K}) = 5.17 \text{ S m}^{-1} \text{ M}^{-1}$, reported in the literature, into independent ionic motion law formula:

$$\Lambda_m^0 = \nu \lambda_+ + \nu \lambda_-$$

(3)

where $\nu$ is the ionic charge, $\lambda$ is the molar conductivity of the ion infinite dilution. The value of $\lambda_\text{i}([\text{BF}_4]^{-}, 298 \text{ K})$ can be calculated from the formula (3), namely, $4.20 \text{ S m}^{-1} \text{ M}^{-1}$, and then take it into the following formula:

$$D = (RT/F^2)[\nu/\lambda_i]$$

(4)

where $R$, $F$, and $Z$ are the gas constant, Faraday constant, the number of ions, respectively, thus the infinite dilution diffusion coefficient of $[\text{BF}_4]^{-}$, $D_{\infty}([\text{BF}_4]^{-}, 298 \text{ K}) = 1.12 \times 10^{-9} \text{ m}^2 \text{ s}^{-1}$ was obtained. Then taking the TX-100 PEO chain length 20 Amy$^{41}$ and $D_{\infty}([\text{BF}_4]^{-}, 298 \text{ K}) = 1.12 \times 10^{-9} \text{ m}^2 \text{ s}^{-1}$ into the formula (2), the relaxation time of 1.78 ns was calculated. In the same way, the relaxation time of the toluene/TX-100/[bmim][PF6] system, whose value is 1.32 ns, can also be calculated, with $D_{\infty}([\text{PF}_6]^{-}, 298 \text{ K}) = 1.515 \times 10^{-7} \text{ m}^2 \text{ s}^{-1}$ (ref. 41) and the 20 Amy PEO chain length of the TX-100 [ref. 42] from formula (2).

According to Fig. 5a and Table 1, we can find that the experimental values of the relaxation time for the benzene/TX-100/[bmim][BF4] system are between 1.330 ns and 4.272 ns which is in the same order of magnitude with its theoretical value 1.78 ns. Similarly, the experimental values of the relaxation time for toluene/TX-100/[bmim][PF6] system are between 0.313 ns and 4.743 ns (Fig. 5b and Table 1) which is also in the same order of its theoretical 1.32 ns. Accordingly, there is

| $C_{[\text{bmim}][\text{BF}_4]}$ (wt) | $\tau_{[\text{bmim}][\text{BF}_4]}$ (ns) | $\kappa_{[\text{bmim}][\text{BF}_4]}$ (S m$^{-1}$) | $C_{[\text{bmim}][\text{PF}_6]}$ (wt) | $\tau_{[\text{bmim}][\text{PF}_6]}$ (ns) | $\kappa_{[\text{bmim}][\text{PF}_6]}$ (S m$^{-1}$) |
|------------------|------------------|------------------|------------------|------------------|------------------|
| 0.018 | 4.27 ± 0.07 | 0.0007 ± 0.0005 | 0.041 | 4.74 ± 0.10 | 0.0025 ± 0.0004 |
| 0.028 | 4.13 ± 0.05 | 0.0018 ± 0.0006 | 0.067 | 4.13 ± 0.12 | 0.0066 ± 0.0004 |
| 0.047 | 3.59 ± 0.06 | 0.0063 ± 0.0004 | 0.098 | 3.59 ± 0.09 | 0.0173 ± 0.0006 |
| 0.069 | 3.13 ± 0.07 | 0.0121 ± 0.0008 | 0.128 | 2.06 ± 0.05 | 0.0380 ± 0.0008 |
| 0.095 | 2.56 ± 0.08 | 0.0288 ± 0.0007 | 0.153 | 1.79 ± 0.08 | 0.0454 ± 0.0006 |
| 0.136 | 1.79 ± 0.09 | 0.0524 ± 0.0006 | 0.196 | 0.35 ± 0.11 | 0.0645 ± 0.0004 |
| 0.162 | 1.67 ± 0.10 | 0.0876 ± 0.0005 | 0.214 | 0.32 ± 0.14 | 0.0731 ± 0.0007 |
| 0.187 | 1.56 ± 0.08 | 0.1023 ± 0.0007 | 0.229 | 0.31 ± 0.10 | 0.0833 ± 0.0005 |
| 0.208 | 1.45 ± 0.11 | 0.1154 ± 0.0008 | 0.244 | 0.36 ± 0.08 | 0.0866 ± 0.0003 |
| 0.238 | 1.33 ± 0.09 | 0.1328 ± 0.0006 | 0.265 | 0.39 ± 0.05 | 0.0913 ± 0.0004 |
| 0.344 | 1.34 ± 0.06 | 0.1421 ± 0.0007 | 0.314 | 0.61 ± 0.07 | 0.0947 ± 0.0005 |
| 0.394 | 1.76 ± 0.05 | 0.1256 ± 0.0008 | 0.356 | 1.67 ± 0.06 | 0.0861 ± 0.0006 |
| 0.450 | 2.08 ± 0.07 | 0.1172 ± 0.0009 | 0.390 | 2.17 ± 0.05 | 0.0819 ± 0.0005 |
| 0.499 | 2.37 ± 0.05 | 0.0999 ± 0.0006 | 0.432 | 3.13 ± 0.08 | 0.0755 ± 0.0006 |
| 0.548 | 3.13 ± 0.08 | 0.0871 ± 0.0004 | 0.471 | 4.13 ± 0.11 | 0.0534 ± 0.0004 |
reason to believe that the relaxation is caused by the fluctuations of the IL anions in the TX-100 hydrophilic chain.

For comparison, the relaxation time $\tau$ for the two IL-based microemulsions was plotted as a function of the weight fraction of ILs in Fig. 5. As can be seen that whether for benzene/TX-100/[bmim][BF$_4$] or toluene/TX-100/[bmim][PF$_6$], the values of $\tau$ were divided into three regions (at the vicinity of 0.1 and 0.28 for benzene/TX-100/[bmim][BF$_4$]; at about 0.18 and 0.3 for toluene/TX-100/[bmim][PF$_6$]). It is worth mentioning that the two inflection points arrowed in Fig. 5 were in amazing consistency with the points P1 and P2 in their phase diagram (Fig. 2), and the three regions correspond with the IL/O, B.C. and O/IL micro area in the phase diagram, respectively. This indicates that the dielectric parameter can be an indicator which can monitor the phase transition and provide certain information on the transition.

It was also seen from Fig. 5 that when ILs concentration $C_{\text{IL}}$ increase near B.C. region, the values of observed $\tau$ close to the calculated values, while the calculated values were always lower than the observed values in O/IL or IL/O micro region. This might be explained as: in the O/IL and IL/O micro regions, ions fluctuated in a confined space. Because in the micelle/reverse micelle shell, there is the ineluctable electrostatic interaction between the ions and fixed charges on the adjacent chains, and the calculated values were obtained under the assumption that the ions drifted freely in the free space. While for the B.C. micro region, although there may be different microstructures for these two different systems, the ion migration was promoted in B.C. of both systems. Because there was no micelles/reverse micelles shell for benzene/TX-100/[bmim][BF$_4$] in B.C. (see Fig. 4B) so that ions in aqueous channel can fluctuate freely; and for toluene/TX-100/[bmim][PF$_6$] (Fig. 4C), the exchange between molecule clusters also accelerated the ion migration. As a result, the experiment values in B.C. of both systems were far closer to the calculated values compared to the above two micro areas.

### 3.3 Electric conduction mechanism

The ions of the ILs played the electric conduction role in the present studied microemulsions. From the model in Fig. 4D, it can be seen that, it is the anions surrounding the TX-100 molecular chain that fluctuated under the action of an alternating electric field. The low frequency conductivity, namely, the $10^5$ to $10^7$ Hz frequency band in the conductivity spectrum (Fig. 6), is ascribed to the migration of anions and cations in the bulk solution along the direction of the electric field (the low frequency band below $10^5$ Hz was from the electrode polarization (EP)\textsuperscript{[a]}). From Fig. 6, we can roughly see that the change of the conductivity $\kappa$ with $C_{\text{IL}}$ is non-linear, i.e. not monotonically increase or decrease. To analyze this variation trend in detail, we plotted the low-frequency conductivity $\kappa_l$ (the values of $\kappa_l$ were drawn from 1 MHz to avoid the influence of EP) against the $C_{\text{IL}}$ for the two IL-based microemulsions in Fig. 7.

From Fig. 7, it can be seen that $\kappa_l$ shows different change trends in the different micro regions of both two systems. Besides, it’s worth mentioning that the inflection points of the two microemulsions are unanimous with the points P1, P2 of the phase diagram shown in Fig. 2. This accordance means that discontinuity of conductivity indicated the changes in the internal structure of the microemulsions. And according to the conductivity, we can get explanations about the mechanism of the microscopic phase transition: the conductivity was small, but increased linearly with the increase of the $C_{\text{IL}}$ in the IL/O region. This is because that although the IL was encapsulated in the microemulsion droplets, due to the large conductivity of the IL, the conductivity of the system was also influenced strongly by the IL. In the B.C. region, since the ILs formed their own special transportation pipeline which enhanced the conductive ability, $\kappa_l$ appears sharp increase at the junction between IL/O and B.C. region for benzene/TX-100/[bmim][BF$_4$] microemulsion (Fig. 7a). While $\kappa_l$ of toluene/TX-100/[bmim][PF$_6$] kept increase trend (Fig. 7b). In the O/IL region, owing to the gradually increasing viscosity of the system, although the IL increased, the migration of IL was constricted. Consequently, the low-frequency conductivity of the systems decreased.

In addition, the areas of the B.C. of the two microemulsion systems in both Fig. 5 and 7, were similar-sized. While their IL/O and O/IL micro regions were different from one another. Regarding to this difference, we can consider from the difference of the hydrophobicity of the ILs in the two microemulsions: [bmim][BF$_4$] is hydrophilic and [bmim][PF$_6$] is hydrophobic. In this way, with the increase of $C_{\text{IL}}$, [bmim][BF$_4$] was easier to form a core with the hydrophilic surfactant TX-100, so the area of O/IL was larger than IL/O. In contrast, as a result of the hydrophobiability of [bmim][PF$_6$], the difference...
between [bmim][PF₆] and toluene was not large, so the IL/O area and O/IL area was roughly the same.

3.4 The percolation in IL-based microemulsions

The uprush of $k_l$ in B.C. region of Fig. 7 reminds us of percolation. In order to obtain the percolation threshold $C_p$, we plotted $d k_l / d C_{IL}$ against $C_{IL}$ as inset in Fig. 7. And its inflection point gives the percolation threshold $C_p$ for benzene/TX-100/[bmim][BF₄] and $C_p$ for toluene/TX-100/[bmim][PF₆] microemulsion, which are pointed out by the arrows. Generally, there are two kinds of theories to explain the percolation process of the low-frequency conductivity of the microemulsion: the static percolation theory comes down the percolation to the appearance of the B.C. structure of oil and water. Dynamic percolation theory holds that it's the interaction between water-drops that causes the formation of percolation clusters, and ions are transported by the exchange of clusters. Both the static and dynamic percolation theory agree that when it closes to the percolation transition concentration (percolation threshold), the relation between DC conductivity and the concentration of the hydrophilic phase follows the scaling law:

$$k_l \propto |C_p - C|^{-s}, \quad C_p > C$$

$$k_l \propto |C - C_p|^\mu, \quad C_p < C$$

where the values of $s$ are predicted to be 0.745 and 1.246 respectively in the static and dynamic percolation theory, and $\mu$ are both 1.9 (ref. 46 and 47) in the two theories. In order to determine the type of percolation of the two tested systems, we take the points near the two sides of the percolation threshold to plot the logarithm of DC conductivity against concentration gradients, and make a straight line respectively. So both the values of $s$ and $\mu$ can be obtained from taking the slopes, as shown in Table 2. The values of index $s$ obtained were 0.503 and 1.039 respectively. That means: for the benzene/TX-100/[bmim][BF₄] system, in the B.C. range, ions were directly into the continuous phase TX-100 constructed through the pipeline, which was called the static percolation process (model graph shown in Fig. 4B). While for toluene/TX-100/[bmim][PF₆] system, the ions reached the other side by the ion exchange between the molecular clusters. That was so-called dynamic process (model graph shown in Fig. 4C).

From the dynamic perspective of the mechanism, the ion migration rate is in proportion to the absolute value of $C_p - C$ which imply that the smaller the absolute value of $C_p - C$ was, far closer to equilibrium, the lower ion transfer rate was. According to the results of our experiments, there is reason to believe that the ions transfer rate of benzene/TX-100/[bmim][BF₄] system was faster than toluene/TX-100/[bmim][PF₆] system. And we can interpret it in the light of percolation model (dynamic or static). In the benzene/TX-100/[bmim][BF₄] system...
Table 2  Comparison of experimental data and theoretical data of percolation threshold

| Entries | Static percolation | Dynamic percolation | Benzene/TX-100/ [bmim][BF₄] | Toluene/TX-100/ [bmim][PF₆] |
|---------|-------------------|---------------------|-----------------------------|----------------------------|
| s       | 0.7               | 1.2                 | 0.503                       | 1.039                      |
| μ       | 1.9               | 1.9                 | 0.609                       | 0.809                      |

(Fig. 4B), because of the hydrophilicity of [bmim][BF₄], they increasingly penetrated to the core of the reverse micelles of TX-100 which urged the system gradually to form a continuous phase just like an ion channel. That's what the static percolation process called, thus the ion migration rate was much faster. While in toluene/TX-100/[bmim][PF₆] system (Fig. 4C), [bmim] [PF₆] is hydrophobic which make it less easily enter the core of the micelles. In addition, IL ions participate in the formation of the molecular clusters of oil phase with toluene. As a result, the ions conducted charges by exchanging charges between molecular clusters, namely dynamic percolation process, so that ion migration rate become relatively slower. Regarding the values of μ of the two systems were somewhat low, this may be because those factual factors didn’t perfectly conform to the assumptions, such as the too large viscosity of the system.

4. Concluding remarks

In emulsion or microemulsion system, the huge oil–water interface results in a relaxation which called the interfacial polarization, relying on the ratio of the electrical properties of both oil and aqueous phase. More specifically, the permittivity of water is close to 80, while that of the benzene and toluene are less than 10. Accordingly, interfacial polarization occurs because of the large differences in electrical properties between the oil phase and water phase. However, the permittivity of IL is just close to 10 similarly to that of oil phase, as a result that the interfacial polarization become less marked and even disappearing. Instead, a relaxation caused by ions fluctuating on the molecular chains of TX-100 was observed. This conclusion was confirmed by the relaxation time obtained with Einstein equation and comparing with our experimental values.

The experiment paths are divided into three distinct regions by the IL concentration dependence of relaxation time or low frequency conductivity. The inflection points were in amazing accordance with the phase transition points in the corresponding phase diagram. In addition, the relaxation parameters and electrical parameters also reveal the microstructure of all regions. All these above strongly confirmed that dielectric analysis can be used as a powerful indicator to find phase change process.

The different of anions of ILs resulted in the different relaxation time, although the ions fluctuate similarly along the same TX-100 molecular chain. This mainly stemmed from the difference on the hydrophilicity of the ILs between the two microemulsion systems: [bmim][BF₄] is hydrophilic, while the [bmim][PF₆] is hydrophobic. From micromosaic point of view, with the change of IL’s concentration, there are three microdomains, i.e., IL/O, intermediate transition region B.C., and O/IL. And our analyzing results suggested diverse microscopic physical picture to show how the ion migrated through the B.C. of two systems. In the benzene/TX-100/[bmim][BF₄] microemulsion, the ions in B.C. directly entered into the continuous phase TX-100 constructed through the pipeline, which was so-called the static percolation process. While for the toluene/TX-100/[bmim][PF₆], ions transported in a dynamic process: ions reached the other side through the ionic exchange between the surfactant molecule clusters.

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