Ion transfer across the water|trifluorotoluene interface

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A B S T R A C T

This work introduces trifluorotoluene as an alternative organic solvent for studying charge transfer across liquid|liquid interfaces. In comparison with the most popular organic solvents in this field, 1,2-dichloroethane and nitrobenzene, trifluorotoluene is less toxic since it is not considered a carcinogen agent, and offers a larger potential window. When comparing the standard Gibbs energies of ion partition from water to trifluorotoluene with those reported at the water|1,2-dichloroethane and water|nitrobenzene interfaces, linear relationships were found, making easy to extrapolate the Gibbs energy of other ions from these empiric correlations. 1,2-dichloroethane and nitrobenzene can be efficiently replaced by trifluorotoluene for studying charge transfer at liquid|liquid interfaces.

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1. Introduction

Even though 1,2-dichloroethane (DCE) [1–8] and nitrobenzene (NB) [9–12] are among the most popular solvents for studying charge transfer across liquid|liquid interfaces, they are classified as highly toxic solvents and carcinogen agents [13–18].

Other solvents such as 2-nitrophenyl octyl ether (NPOE), 2-heptanone [19,20], 2-octanone [20], 1,6-dichlorohexane [21,22], 1,4-dichlorobutane [22], acetoephone [23], dichlorobenzene [24], and even lately room temperature ionic liquids (RTIL) [25–29] have been found to be suitable, but are either expensive or difficult to prepare.

Trifluorotoluene (TFT) is a colourless and free-flowing liquid with a relatively low toxicity and price, considered as an efficient substitute for dichloromethane and similar solvents in organic synthesis [30,31]. It is not a carcinogen agent according to its MSDS safety data sheet. Comparing the physical properties of DCE and TFT, the dielectric constant and density of TFT (9.2 at 20 °C and 1.19 g/mL, respectively) are only slightly smaller than those of DCE (10.3 at 20 °C and 1.25 g/mL), whilst the boiling point of TFT (102 °C) is 18 °C higher than that of DCE, suggesting that TFT could be a potential candidate to replace DCE in studies at liquid|liquid interfaces.

In this work, the standard transfer potential (Δφ θ) of different ions transferred across the water|TFT interface were determined and compared with those reported at the water|DCE and water|NB interfaces. The relationship between the standard Gibbs energies of the partition (ΔG θ,w→f,), when comparing the three interfaces is linear, making easy to extrapolate the Gibbs energy of other ions from such empiric correlations.

2. Experimental

2.1. Chemicals and reagents

All solvents and chemicals were used as received without further purification. Bis(triphenylphosphoranylidine)ammonium chloride (BACl 98%), lithium tetrakis(pentafluorophenyl)-borate ethyl etherate (LiTB purum), tetramethylammonium chloride (TMCl 98%), tetrabutylammonium bromide (TBABr 99%) and sodium tetraphenylborate (NaTPB, 99.5%) were purchased from Fluka. α,α,α-Trifluorotoluene (TFT 99% +), tetraethylammonium chloride (TEACl 99%) and tetrabromo- and tetrabromocinnamylammonium (TBrPAb 98%) were purchased from Acros. Tetraphenylarsonium chloride monohydrate (TPASCl 99%) was purchased from ABCR. 1,2-Dichloroethane (DCE, grade HPLC) was purchased by Applichem. Tetraphenylarsonium tetrabromide (TPASTPB) was prepared as reported in Ref. [33]. Bis(triphenylphosphoranylidine)ammonium tetrakis(pentafluorophenyl)-borate (BATB) was prepared as reported in Ref. [8]. All the aqueous solutions were prepared in ultra pure water (18.2 MΩ cm−1). The solubility of TPASTPB in TFT was found to be rather low, thus, in order to dissolve enough salt to see reliable transfer of ions, a saturated TPASTPB solution was prepared, sonicated for 15 min and filtrated to remove the insoluble salt. Clear signals of the transfer of TPAS+ and TPB− were observed by cyclic voltammetry.

2.2. Ion transfer voltammetry measurements at the water|TFT interface

Ion transfer voltammetry measurements at the water|TFT interface were performed in a four-electrode cell with two reference electrodes to polarise the interface and two platinum counter electrodes to provide the current. A commercial potentiostat (PGSTAT 30, Metrohm, CH) was used. The external potential was applied by means of two silver/silver chloride (Ag/AgCl) reference electrodes,
which were connected to the aqueous and the organic phases, respectively, by means of a Luggin capillary. The composition of the electrochemical cells is illustrated in Scheme 1. The origin of the Galvani potential difference $\Delta o_{\phi}$ was estimated by using the extra-thermodynamic TATB assumption [32].

3. Results and discussion

Taking into account the standard transfer potential of the ions forming the supporting electrolytes in both phases (see cell I, Scheme 1; $BA^+$, $TB^-$, $Li^+$ and $Cl^-$), the cyclic voltammograms in Fig. 1 are limited by the transfer of $Li^+$ in the positive side of the scale and by $Cl^-$ at the negative part [8]. In spite of the low solubility of the electrolyte TPAsTPB in TFT, the $\Delta o_{\phi}^{1/2}$-scale was calibrated by using the extra-thermodynamic TATB assumption in which the standard Gibbs energy of partition of TPAsTPB can be split into two equal parts for the cation and the anion because the radii of both ions are very similar; $\Delta G_0^{w-o} (TPA^+) = \Delta G_0^{w-o} (TPB)$ or $\Delta o_{\phi}^{1/2} (TPA^+) = -\Delta o_{\phi}^{1/2} (TPB^-)$ [32]. Thus, the zero of the $\Delta o_{\phi}^{1/2}$-scale is located in the middle of the half-wave potentials of both ions (Fig. 1a). TPropA$^+$ was added also to the aqueous phase (see cell II, Scheme 1) in order to determine its $\Delta o_{\phi}^{1/2}$ and use it as a secondary reference in forward voltammograms made at the water|TFT interface. Fig. 1a shows the calibrated voltammogram of the supporting electrolyte at the water|TFT and the water|DCE interfaces. The potential window obtained at the water|TFT interface is around 400 mV and 100 mV larger toward the positive and the negative side, respectively, than the potential window obtained at the water|DCE interface.

The usual analogous series of ammonium salts, TMA$^+$, TEA$^+$, TPropA$^+$ and TBA$^+$, was studied at the water|TFT interface (cells IV–VI, Scheme 1) in order to calibrate the whole potential window and determine if there is any important deviation with respect to the behaviour found at the water|DCE and water|NB interfaces for the same series (Fig. 1c). As expected from the behaviour observed in DCE the scan rate dependence of the transfer of TPropA$^+$ and TMA$^+$ (see cell IV, Scheme 1) when tested from 10 mV/s to 50 mV/s, showed that the process is controlled by diffusion. The same behaviour was found for the other cations. The calibration of the potential window was made systematically from TPropA$^+$ whose $\Delta o_{\phi}^{1/2}$ was already determined using the cell III (Scheme 1), then, using the cell IV the $\Delta o_{\phi}^{1/2}$ of TMA$^+$ was calibrated, which was in turn used to determine the $\Delta o_{\phi}^{1/2}$ of TBA$^+$ using the cell V, and finally the $\Delta o_{\phi}^{1/2}$ of TBA$^+$ was used to calibrate the transfer of TEA$^+$ using the cell VI.

![Fig. 1](https://example.com/f1.png)

Fig. 1. (a) Calibration of the potential window at the water|TFT interface using the TATB assumption. Scan rate: 10 mV/s, SE: Supporting Electrolyte. (b) Comparison of the potential windows obtained at the water|TFT and water|DCE interfaces, scan rate: 20 mV/s. (c) Determination of the $\Delta o_{\phi}^{1/2}$ of TMA$^+$, TEA$^+$, and TBA$^+$ at the water|TFT interface. Scan rate: 10 mV/s for cells IV and V and 30 mV/s for cell VI.
Eq. (1) was used to estimate the standard potentials \( \Delta G^0_{\text{w,TFT}} \) at the water|TFT interface from the half-wave potential values, \( \Delta G^0_{\text{w,1/2}} \), obtained experimentally,

\[
\Delta G^0_{\text{w,1/2}} = \Delta G^0_{\text{f}} + \frac{RT}{2F} \ln \left( \frac{D_m}{D_f} \right) + \frac{RT}{2F} \ln \left( \frac{\gamma_m}{\gamma_f} \right)
\]  

(1)

where \( D_m, D_f \) and \( \gamma_m, \gamma_f \) are the diffusion and activity coefficients for the aqueous and the organic phases, respectively. The ratio \( D_m/D_f \) was estimated by using Walden’s rule \( \eta_mD_m = \eta_fD_f \), with \( \eta \) the viscosity. The viscosity of water and TFT is equal to 0.0091 P and 0.00038 P (at 25 °C), respectively, and the constant \( \frac{RT}{2F} \ln \left( \frac{\gamma_m}{\gamma_f} \right) \) was calculated to be ±41.3 mV for univalent ions.

The activity coefficients were calculated by using the extended Debye–Hückel equation (Eq. (3)) [32], where \( \mu \) is the ionic strength and \( \alpha \) is the ion size, which for the ions studied in this work can be assumed as 0.3 nm in the aqueous phase and 0.6 nm in the organic phase [32].

\[
\log(\gamma_i) = -\frac{A\mu_1/2}{1 + B\alpha_1/2}
\]  

(2)

The constants \( A \) and \( B \) were estimated by using Eqs. (3) and (4) [34], respectively. The obtained values were \( A = 12.71 \text{L}^{1/2} \text{mol}^{-1/2} \) and \( B = 9.60 \text{mol}^{-1/2} \text{nm}^{-1} \) for TFT and \( A = 0.51 \text{L}^{1/2} \text{mol}^{-1/2} \) and \( B = 3.3 \text{mol}^{-1/2} \text{nm}^{-1} \) for water.

\[
A = 1.8245 \times 10^{-6}
\]  

(3)

\[
B = 502.904
\]  

(4)

Therefore, the constant \( \frac{RT}{2F} \ln \left( \frac{\gamma_m}{\gamma_f} \right) \) was calculated to be ±35.6 mV for univalent ions.

Finally, the \( \Delta G^0_{\text{w,1/2}} \) determined experimentally and the corrected \( \Delta G^0_{\text{w,1/2}} \) and \( \Delta G^0_{\text{w,TFT}} \) are tabulated in Table 1 along with \( \Delta G^0_{\text{w,DCE}} \) and \( \Delta G^0_{\text{w,NB}} \) reported for water|DCE and water|NB interfaces.

\[
\Delta G^0_{\text{w,TFT}} \rightarrow \Delta G^0_{\text{w,DCE}} \rightarrow \Delta G^0_{\text{w,NB}}
\]

(5)

The \( \Delta G^0_{\text{w,1/2}} \) determined experimentally and the corrected \( \Delta G^0_{\text{w,1/2}} \) and \( \Delta G^0_{\text{w,TFT}} \) are tabulated in Table 1 along with \( \Delta G^0_{\text{w,DCE}} \) and \( \Delta G^0_{\text{w,NB}} \) reported for water|DCE and water|NB interfaces. \( \Delta G^0_{\text{w,DCE}} \) and \( \Delta G^0_{\text{w,NB}} \) obtained at the water|TFT interface are quite similar to those reported for the water|DCE interface, which in some extent validates the data obtained in this work, considering that the dielectric constant of TFT is only slightly smaller than that of DCE.

Fig. 2 shows the relationship between the \( \Delta G^0_{\text{w,TFT}} \) obtained at the water|TFT interface and those reported at the water|DCE and water|NB interfaces. In both cases linear relationships were found, in agreement with previous works for similar solvents [32]. The slopes of the correlation curves are close to the unity in both cases; 1.0 and 1.3 when comparing with DCE and NB respectively, and the intercepts are -0.48 (TFT vs DCE) and 12.2 (TFT vs NB) indicating that the \( \Delta G^0_{\text{w,TFT}} \) are larger than those reported for the transfer to NB. Those values also correlate rather well with those found by other authors for similar comparisons [32] and suggest that the \( \Delta G^0_{\text{w,DCE}} \) and \( \Delta G^0_{\text{w,NB}} \) determined at the water|DCE and water|NB interfaces, could be used to estimate \( \Delta G^0_{\text{w,TFT}} \) at the water|TFT interface by using these empirical relationships [32].

4. Conclusion

TFT was found to be a good substitute for DCE and NB for studying charge transfer at liquid|liquid interfaces, with the advantage of having lower toxicity (TFT is not considered a carcinogen agent) and price and offering a larger potential window.

Acknowledgements

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