EC solvent mixture. The high value of the solvation number in 100 wt.% EC may be due to the primary solvation layer together with some contribution from the secndary solvation layer.

An Estimation of the Volume of Restructured Water Shells around Hydrophobic Solutes

**Ü. HALDNA and L. ORASTE** 

Department of Chemistry, Tartu State University, Tartu, Estonian S.S.R., U.S.S.R.

Water is known to be a largely associated liquid. The presence of non-electrolytes or large organic ions, tends to strengthen the hydrogen bonds between the water molecules near the large hydrophobic solute, and a cage or 'iceberg' is effectively formed around them. This model of the hydrophobic hydration of alkyl groups is widely accepted but the thickness of restructured water shells around hydrophobic solutes has not been determined. In order to estimate the volume of restructured water per mole hydrophobic solute we used the differential conductometry of carefully thermostatted (0.001 °C) solutions [1, 2]. The experimental procedure consists of measuring the specific electrical conductance  $(\chi)$  of a strong electrolyte solution before and after adding a small amount of hydrophobic solute. Provided the equivalent conductance of the strong electrolyte ions in the restructured water shell is zero [3] the volume of the restructured water shell is given by [2, 3].

$$V_{s,o} = \frac{\Delta \chi \cdot 10^3}{\chi \cdot c} \text{ [cm^3/mol]}$$

where  $\Delta \chi$  is the change in  $\chi$ , corrected for the increase in volume and c is the molar concentration of the strong electrolyte.

We have attempted to measure the  $V_{s,o}$ -values in aqueous solutions of perchloric acid for 16 alkylammonium ions. The results obtained can be summarized as follows:

$$V_{s,o} = n(CH_3) \cdot V(CH_3) + n(CH_2) \cdot V(CH_2) + n(NH_3^*) \cdot V(NH_3^*)$$

where n(i) is the number of groups i in the ion studied and V(i) is the specific volume of hydration shell of i-group. In 10–20% HClO<sub>4</sub>(w/w), V(CH<sub>3</sub>)  $\approx$ V(CH<sub>2</sub>)  $\approx$  40 ± 10 [cm<sup>3</sup>/n(i) mol] and V(NH<sub>3</sub><sup>+</sup>)  $\approx$ -48 ± 10 [cm<sup>3</sup>/n(i) mol]. The negative value for  $V(NH_3^*)$  reflects the negative hydration of  $-NH_3^*$ -group.

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## Structure of Aliphatic Alcohols in Electrical Double Layer on Electrode Surface

U. PALM and M. VÄÄRTNŎU

Laboratory of Electrochemistry, Tartu State University, Tartu 202400, Estonian S.S.R., U.S.S.R.

The structure of solvent in the field of the electrical double layer exerts significant influence on the inner layer properties on adsorption of ions and molecules. Some theoretical models considering differently the interaction of solvent dipoles in the inner layer have recently been elaborated [1-3]. On the basis of the experimental data obtained by us on bismuth electrode in alcoholic solutions, mainly in ethanol, the analysis of the validity of the cluster models [1-3] for the description of the solvent structure in the inner layer has been carried out in this paper. Cluster model was chosen since alcohols belong to the group of associated liquids. According to cluster model solvent exists on the electrode surface in the form of separate chemisorbed molecules with constant orientation and of associates of molecules (clusters) the orientation of which depends upon electrode charge q and the composition of which varies with temperature T.

By use of the equations of the cluster model [1-3] the electrode charge has been calculated for several temperatures in the interval from -15 to 50 °C. The results of the theoretical calculations were compared with the experimental data in the form of the plots of the inner layer integral capacity  $K_{02}$  against q at several T. Experimental values of  $K_{02}$  were found from the differential capacity of bismuth in alcoholic solutions of LiClO<sub>4</sub> at various temperatures.

On the basis of the calculations it was established, that at T < 25 °C the ethanol molecules exist in the inner layer mainly in the form of two-dimensional double associates due to the hydrogen bonding