1 Introduction

Antagonistic salts, which consist of hydrophilic cations and hydrophobic anions, have recently been found to induce mesoscale assembly in binary aqueous solvents.\textsuperscript{1,2} Already with a few mM of added salt mesoscale structures were observed, and increasing the salt concentration caused the appearance of an ordered lamella-like phase.\textsuperscript{1-3} These observations were initially assigned to the hydrophilic and hydrophobic nature of cations and anions, respectively, with the antagonistic salt behaving essentially as a surfactant.\textsuperscript{3} In contrast, no solvent ordering is observed upon adding simple inorganic salts. For a recent review on the topic, see ref. 4.

The mechanism of the aforementioned assembled structures is still under debate. Some theoretical models emphasize the interpretation in terms of cation and anion preferential solubility, with the solvent ordering being caused by the unequal partitioning of hydrophilic and hydrophobic ions in water- and organic-rich phases.\textsuperscript{5,6} On the other hand, theoretical studies of antagonistic salts exhibiting no ion size asymmetry do not find mesoscale solvent ordering.\textsuperscript{5} Indeed, an alternative model-independent prediction indicates that such solvent structures can only be generated if the added salt exhibits significant asymmetry of ion size.\textsuperscript{6} Note that all previous experiments were conducted on a system containing antagonistic salt, which exhibits ion size asymmetry.

In this paper, we focus on the effect of ion size asymmetry on mesoscale ordering in binary aqueous solvents. For this purpose we have carried out a small-angle X-ray scattering (SAXS) experiment in a mixture of water and 2,6-dimethylpyridine (denoted as 2,6-DMP) by adding 10 mM of quaternary ammonium bromide. The SAXS data were collected for a salt series ranging from the symmetrical short-chained tetraethylammonium bromide to the longer-chained tetraheptylammonium bromide. Most importantly, our experimental results are the first to show that a systematic increase of ion size asymmetry leads to a gradual build-up of mesoscale order in binary aqueous solvents.

The reason for choosing a series containing symmetrical tetra-\textit{n}-alkylammonium salts is dictated by their appealing properties. First, the cations become larger as the number of carbon atoms increases; since they are built in a symmetrical manner (four hydrocarbon chains), adding more carbons gives a bulky ion. Second, it is well known that their properties are linked with the structure and number of hydrocarbons.\textsuperscript{5} Therefore, we expected the data to depend on the length of alkyls. The choice of salts with the same anion results in a neat series ranging from the symmetrical short-chained tetraethylammonium bromide to the longer-chained tetraheptylammonium bromide. Most importantly, our experimental results are the first to show that a systematic increase of ion size asymmetry leads to a gradual build-up of mesoscale order in binary aqueous solvents.
Moreover, we collected data from our binary solvent containing the antagonistic sodium tetraphenylborate (Sigma-Aldrich, purity ≥ 99.5%) as a reference. All chemicals were used as received, without further purification.

### 3 Results and discussion

The phase diagram of the neat mixture of water and 2,6-DMP is well known, exhibiting an immiscibility loop with a lower critical temperature $T_c \approx 307$ K at a volume fraction $\phi \approx 0.3$ of 2,6-DMP. Furthermore, the addition of ions usually alters the phase diagram. Hydrophilic inorganic salts enlarge the two-phase region and shift the critical point, in contrast to antagonistic salts which lead to the shrinking and eventual disappearance of the two-phase region as a function of the added salt concentration. The latter behavior is in line with the addition of ionic surfactants such as sodium dodecyl sulfate (SDS) and dioctyl sodium sulfosuccinate (often denoted as AOT).

Let us start by considering the effect of introducing quaternary ammonium salts into the water–2,6-DMP solvent. In Fig. 1 we present the experimental phase diagram as obtained following ref. 15, using a temperature step of 0.5 K and an equilibration time of 20 minutes. Strikingly, there is no effect of 10 mM tetraalkylammonium bromide on the lower critical point location of the water–2,6-DMP mixture, in stark contrast to the salts mentioned above. This finding suggests that the quaternary ammonium salts introduced here may be proven to be a highly useful model system to study salt-induced mesoscale ordering in binary aqueous solvents. In particular, the absence of a shift of the lower critical point facilitates our systematic study on the effect of ion-size asymmetry in salt-induced mesoscale solvent ordering. In the rest of this study, we will therefore focus on the critical composition $\phi = 0.3$ and 10 mM added salt.

The sample with 2,6-DMP content of $\phi = 0.1$ and 10 mM of added tetraheptylammonium bromide (C7) was turbid already at room temperature, directly after mixing all components. This is in contrast to mixtures with $\phi \geq 0.20$, in which C7 dissolves completely. In order to verify this observation, the preparation was repeated with the same result. The limited solubility of C7 in the solvent provides a possible explanation for the turbidity of the sample. Adding 10 mM C7 to the mixture with a low fraction of 2,6-DMP could cause spontaneous emulsification, in line with the previous observations of oil-in-water emulsions containing symmetrical alkylammonium ions.

Having verified that 10 mM of quaternary ammonium salts do not alter the phase diagram (except C7 at small $\phi$), we next turn to the main objective of our study – solvent ordering induced by ion size asymmetry. For this purpose we have carried out a SAXS experiment at beamline I911-4 of the MAX-lab (Lund, Sweden), using an X-ray wavelength of $\lambda = 0.91$ Å. We kept the samples in thin-walled, 1.5 mm-diameter glass capillaries, and we recorded the scattered X-rays 1.9 m behind the sample using the two-dimensional pixel detector PILATUS 1 M. The X-ray beam focused onto the detector had a size of 0.22 × 0.3 mm at the sample position. We collected data at several temperatures below $T_c$ in the range of $\Delta T = T_c - T \approx 4 - 9$ K and we used an exposure time of 300 s for each sample. For more specific beamline details, we refer the reader to ref. 17.

In order to exemplify the SAXS experiment, we compare in Fig. 2 data obtained upon adding either 10 mM hydrophilic tetraethylammonium bromide or tetraheptylammonium bromide. The data are presented for a temperature of 303 K in order to emphasize the characteristic feature of solvent ordering, and we have subtracted a temperature-independent background consisting of a linear combination of recorded water and 2,6-DMP data with a 2,6-DMP volume fraction $\phi = 0.3$ (see the ESI for details). The neat solvent can be described by the classical Ornstein–Zernike (OZ) model, $I(q) = I(0)/(1 + q^2 \xi^2)$, where $I(q)$ is the scattered intensity as a function of scattering vector modulus $q$ and $\xi$ the bulk correlation length describing the concentration fluctuations. The latter is strongly temperature dependent and close to the critical point it follows the three-dimensional Ising universality class: $\xi = \xi_0 (\Delta T/T_c)^{\nu}$, with the critical exponent $\nu \approx 0.63$, $\Delta T = T_c - T$, and $\xi_0$ a sample-dependent constant. For the system composed of water and 2,6-DMP the constant has previously been found to be $\xi_0 = 0.20 \pm 0.02$ nm, and for the salt-containing mixtures studied here we observe $\xi \approx 1.7$ nm at 298 K, in reasonable agreement with the neat case considering our accuracy in Fig. 1. First, we discuss the case of adding C2. A small amount of C2 does not induce ordering of the solvent, as is clear from the linear behavior in the OZ plot of Fig. 2(b). In this respect C2 behaves like simple inorganic salts. Second, if we change the salt to C7, it shows up as a concentration-dependent shoulder in the SAXS intensity of panel (a) and as a non-linearity in panel (c). This behavior is similar to that when adding the antagonistic sodium tetraphenylborate NaBPh$_4$, and we thus expect this...
shoulder to be further enhanced when increasing the temperature towards the critical point.\textsuperscript{3,13} We emphasize that the C7 data of Fig. 2 can no longer be described by the OZ model.

Let us focus on the case of C7. In order to describe the experimental data properly and to capture the mesoscale structure, one needs to employ a generalized OZ model,\textsuperscript{5,6,8}

\[
I(q) = \frac{I(0)}{1 + \left|1 - g^2/(1 + \kappa^{-2}q^2)\right|q^2}\quad(1)
\]

In comparison with the classical OZ model, additional parameters are introduced. First, the generalized OZ relation depends on the solution’s ionic strength via the Debye screening length $\kappa^{-1}$. Assuming the mixture’s dielectric constant $\varepsilon_{\text{mix}} = \phi\varepsilon_{\text{2,6-DMP}} + (1 - \phi)\varepsilon_{\text{water}}$ with $\varepsilon_{\text{2,6-DMP}} \approx 7$ and $\varepsilon_{\text{water}} \approx 80$, we obtain $\kappa^{-1} \approx 2.59$ nm for a 10 mM solution with 1:1 valency.

Second, an interaction contrast $g$ is introduced, i.e., a contrast between anion–solvent and cation–solvent interactions. We can relate $g$ to the salt-induced mesoscale ordering in the solvent;\textsuperscript{5,6,8} with increasing $g$ the solvent becomes more ordered, and for $g > 1$ a shoulder (or peak) appears in the structure factor. Note that in the limit of $g = 0$, and hence no interaction contrast, eqn (1) yields the classical OZ model.

It is worth noting that for low salt concentrations (as studied here) the different proposed theoretical models lead to the same functional form of the structure factor, although the underlying mechanisms differ; while some models\textsuperscript{5,6} assign the solvent ordering to the asymmetric solvation of ions, exhibiting surfactant-like behavior,\textsuperscript{6} the only model-independent prediction emphasizes asymmetric steric ion–solvent interactions.\textsuperscript{8}

In Fig. 3 we show the modulus of the interaction contrast $|g|$ versus the size of the cation, as obtained by fitting the generalized OZ model of eqn (1) to the SAXS data. In order to exemplify our sensitivity to the value of $|g|$, we also show in Fig. 2(a)
For the salt series, the steric cation–solvent interactions are more significant than the anion–solvent interactions, which results in the observed gradual build-up of the solvent ordering. This is in qualitative agreement with model-independent theoretical predictions. These results are of high importance in the fields of soft matter and colloidal science, where similar ordering in the so-called pre-Ouzo region is applied for designing stable surfactant-free microemulsions and for mesoscale solubilization using hydrodopoles instead of surfactants.

Finally, we note that more pronounced mesoscale solvent ordering has been observed upon increasing the concentration of antagonistic salts. Here we have focused on a low salt concentration, where no salt-induced shift of the lower critical point occurs, thus facilitating our systematic study as a function of ion-size asymmetry. At this low salt concentration we observe only the initial build-up of mesoscale ordering. In the future we foresee extending these studies to larger salt concentrations and different solvent compositions, where more pronounced mesoscale solvent ordering is expected.

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