Contact Ion Pairs in the Bulk Affect Anion Interactions with Poly(N-isopropylacrylamide)

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ABSTRACT: Salt effects on the solubility of uncharged polymers in aqueous solutions are usually dominated by anions, while the role of the cation with which they are paired is often ignored. In this study, we examine the influence of three aqueous metal iodide salt solutions (LiI, NaI, and CsI) on the phase transition temperature of poly(N-isopropylacrylamide) (PNIPAM) by measuring the turbidity change of the solutions. Weakly hydrated anions, such as iodide, are known to interact with the polymer and thereby lead to salting-in behavior at low salt concentration followed by salting-out behavior at higher salt concentration. When varying the cation type, an unexpected salting-out trend is observed at higher salt concentrations, Cs⁺ > Na⁺ > Li⁺. Using molecular dynamics simulations, it is demonstrated that this originates from contact ion pair formation in the bulk solution, which introduces a competition for iodide ions between the polymer and cations. The weakly hydrated cation, Cs⁺, forms contact ion pairs with I⁻ in the bulk solution, leading to depletion of CsI from the polymer–water interface. Microscopically, this is correlated with the repulsion of iodide ions from the amide moiety.

INTRODUCTION

Since the end of the 19th century, it has been known that the addition of salts to water affects the properties of non-electrolytes dissolved in the solution. Anions have, for example, the ability to precipitate macromolecules such as polymers and proteins from an aqueous solution. Their propensity to do so is ranked according to the Hofmeister series. The salting-out order of the anionic Hofmeister series is CO₃²⁻ > SO₄²⁻ > S₂O₅²⁻ > H₂PO₄⁻ > F⁻ > Cl⁻ > Br⁻ > NO₃⁻ > I⁻ > ClO₄⁻ > SCN⁻. Anions are usually classified as weakly hydrated (right hand side of the series) or strongly hydrated (left hand side of the series). Weakly hydrated anions partition to nonpolar environments such as air–water and polymer–water interfaces, leading to moderately increased polymer solubility (salting-in behavior). Strongly hydrated anions interact repulsively with and are depleted from the polymer surface instead (salting-out behavior).

Hofmeister effects have commonly been investigated with poly(N-isopropylacrylamide) (PNIPAM), a thermoresponsive water-soluble polymer broadly used as a model for polymers that exhibit a coil–globule transition above their lower critical solution temperature (LCST). Many studies have focused on the effect of the anion in combination with sodium ions, which act as charge balancing counter ions. Indeed, Hofmeister phenomena are stronger for anions than for cations. Despite the focus on anions, a typical consensus cationic Hofmeister series exists in the literature, and the salting-out order for proteins is as follows: N(CH₃)₄⁺ > NH₄⁺ > Cs⁺ > Rb⁺ > K⁺ > Na⁺ > Li⁺ > Ca²⁺ > Mg²⁺. This series is not as well understood as the anionic Hofmeister series. Interestingly, the order is opposite to the anionic series. That is, strongly hydrated cations (the right-hand side of the series) lead to salting-in, and weakly hydrated cations (the left-hand side of the series) lead to salting-out behavior.

In this study, we report ion-specific effects on the LCST of PNIPAM in salt solutions consisting of the weakly hydrated iodide ion in combination with the alkali metal ions (Li⁺, Na⁺, and Cs⁺). The addition of iodide induces salting-in behavior (swelling of the polymer chain) at low salt concentration due to direct ion binding. This is followed by salting-out behavior (collapse of the polymer chain) at higher concentration where solvation of the metal cations and anions in the bulk solution effectively causes the polymer to precipitate out. We show that the LCST behavior (the rate of change with salt concentration) of PNIPAM in metal iodide salt solutions depends on the specific nature of the cation. By employing all-atom molecular dynamics (MD) simulations, this ion-specific behavior is elucidated at the molecular level.

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**EXPERIMENTAL METHODS**

**Materials.** LiI (99.9% purity), NaI (99.5% purity), and CsI (99.9% purity) were all purchased from Sigma-Aldrich. PNIPAM, with a molecular weight of 186,800 g/mol and a polydispersity of 2.63, was purchased from Polymer Source, Inc. A fixed amount of each salt was dissolved with PNIPAM in nitrogen-purged water (used to avoid iodine formation) to obtain solutions at each desired salt concentration (25–1000 mM).

**Phase Transition Temperature Measurements.** The change in turbidity was measured in solutions containing 10 mg/mL PNIPAM and various concentrations of LiI, NaI, and CsI to determine the LCST. An automated melting point apparatus (MPA 100 Optimal, Standford Research Systems) with digital image processing software was used. A ramp rate of 1 °C/min was used to measure the light scattering intensity as a function of temperature. More specifically, the LCST was determined from the onset of the light scattering increase relative to the flat and low intensity baseline observed at colder temperatures. Details concerning the LCST measurements have been described before.

**MOLECULAR DYNAMICS SIMULATIONS**

**System Setup and Simulation Details.** All-atom simulations were carried out using the MD package GROMACS 2018. The simulated systems were composed of a PNIPAM 20-mer chain (isotactic-rich with 60% meso diad content) in aqueous salt solutions of LiI, NaI, and CsI. The setup featured a stretched polymer chain extending through the periodic z-boundary. By connecting the head and tail, the chain had no end groups and can be considered as virtually infinite. This setup has been reported before. By means of the GROMACS pull code, a collapsed structure of the PNIPAM chain was pulled apart generating an elongated chain. The contour length of the 20-mer, $L_c = 5.32$ nm, together with the desired elongation of $\lambda = L_c/L_0 = 0.88$ defined a box size in the z-direction of $L_z = 4.6839$ nm. The box dimensions in the x- and y-directions were both initially 6.5 nm. The computational cost was minimized with this size, while interactions between periodic images were still prevented. A modified OPLS-AA force field was used for the PNIPAM chain. LINCS was utilized to constrain all bonds up to a fourth-order expansion. The chain was solvated in water using the SPC/E potential with the SETTLE constraint algorithm keeping the internal geometry of the water molecules rigid. After energy minimization and equilibration, two water molecules at a time were replaced with an anion (I$^-$) and a cation (Li$^+$, Na$^+$, or Cs$^+$) until a salt concentration of 1 m (6216 water molecules, 112 cations, and 112 anions) was achieved. While molality (m, moles/liter solution) was used to describe the concentration in the simulations, molarity (M, moles/liter solution) was used to describe the concentration in the simulations, molarity (M, moles/kg water) was used to describe the concentration in the simulations, molarity (M, moles/liter solution) was used to describe the concentration in the simulations.

**Interaction between Different Atoms within the PNIPAM Chain.** Interactions between different atoms within the PNIPAM chain as well as polymer–ion and polymer–water interactions were described with the geometric combination rule. The Lorentz–Berthelot combination rule was used to describe ion–ion and ion–water interactions. Furthermore, an additional scaling factor of 0.9 was used for the dispersion interaction strength, $\epsilon_{ij}$ between sodium ions (i) and iodide ions (j) according to the original force field.

The systems were energy-minimized followed by a 2 ns NVT equilibration run using a velocity-rescaling thermostat. Two consecutive NPT equilibration runs were subsequently performed. This started with a 2 ns equilibration using the Berendsen barostat and the velocity-rescaling thermostat. Next, a 3 ns equilibration was performed using the Parrinello–Rahman barostat and the Nose–Hoover thermostat. Finally, production runs of 100 ns were performed using the same barostat and thermostat as the last equilibration run. Coupling times of $\tau_p = 2$ ps and $\tau_T = 1$ ps were used for the barostats and thermostats, respectively, for all runs. All simulations were performed at 1 bar and 300 K. A semi-isotropic pressure coupling scheme in the x- and y-dimensions was applied with compressibilities of $\kappa_x = 4.5 \times 10^{-5}$ bar$^{-1}$, and $\kappa_y = 0$ bar$^{-1}$ was applied in the z-dimension to maintain the stretched PNIPAM chain. Periodic boundary conditions were applied in all three directions. Furthermore, the particle mesh Ewald (PME) method was used with a Fourier spacing of 0.12 nm, PME order of 4, and a real space cutoff of 1.4 nm to treat the long-range electrostatic interactions. A cutoff radius of 1.4 nm was used for van der Waals interactions. No long-range pressure and energy corrections were applied. For the neighbor list, a cutoff distance of 1.4 nm was used and was updated every 0.002 ps. Configurations were saved every 1 ps, and an integration time step of 0.002 ps was used as well.

Based on the general simulation details described above, four sets of simulations (I–IV) were conducted with the PNIPAM chain and one set without it (V). Setup I corresponds to PNIPAM solvated in different salt solutions with all other parameters according to the description above. Setups II and III include modifications of the PNIPAM chain solvated in different salt solutions. Setup II restraints PNIPAM. That is, a position restraint with a force constant of 1000 kJ/(mol nm$^2$) was applied to all atoms of the polymer chain in all three dimensions. Setup III considers a “nonpolar” PNIPAM, i.e., a PNIPAM 20-mer with all partial atomic charges set equal to zero. Setup IV includes PNIPAM solvated in modified CsI solutions, i.e., using Lennard-Jones size parameters, $\sigma_{ij} = \lambda_i \sigma_{i,j} / 2$, between cations (i) and anions (j) scaled with factors $\lambda_i$ of 1.4 and 1.8, respectively. For setup V, bulk simulations of 1.0 m LiI, NaI, and CsI were performed. A cubic box with dimensions of $4 \times 4 \times 4$ nm$^3$ with 2089 water molecules, 38 anions, and 38 cations was used. Force fields and simulation parameters were the same as for the systems containing the polymer chain. Production runs were performed for 50 ns. If not stated otherwise, setup I was used.

**Validation of the Lithium Iodide Force Field.** The Li$^+$ force field was comprised of a nonpolarizable Li$^+$ ion combined with the iodide parameters taken from the NaI and CsI models. This force field was validated against the osmotic coefficient, and the procedure has been described before. However, a flat-bottom position restraint was used with a harmonic force to confine the ions. A cubic box, $5.4 \times 5.4 \times 5.4$ nm$^3$, filled with water, Li$^+$, and I$^-$ ions was simulated using the GROMACS 2019 package. Three different concentrations were examined: 0.2 (water molecules = 280, Li$^+$ = 19, and I$^-$ = 19), 0.6 (water molecules = 5200, Li$^+$ = 56, and I$^-$ = 56), and 1.0 m (water molecules = 5130, Li$^+$ = 93, and I$^-$ = 93). Energy minimization was performed followed by
a 0.25 ns NVT equilibration and a 1.5 ns NPT equilibration. Next, the simulation box was extended in the z-dimension to a total size of 10.8 nm, and the cubic box was placed in the middle. The new larger box was filled with water molecules leading to total numbers of 10,860, 10,820, and 10,810 water molecules, respectively, for the three concentrations. Another energy minimization was carried out followed by a 0.25 ns NVT equilibration and a 1.5 ns NPT equilibration. Afterward, production runs of 40 ns were performed. This was long enough to generate converged osmotic coefficients. The flat-bottom position restraints acted in the z-dimension. A semi-isotropic pressure coupling scheme was therefore applied, and only the x- and y-dimensions of the box were adjusted. Compressibilities of \( \kappa_{x,y} = 4.5 \times 10^{-5} \text{ bar}^{-1} \) and \( \kappa_z = 0 \text{ bar}^{-1} \) were used. All other simulation settings were the same as for the simulations including the PNIPAM chain.

Lithium–iodide Lennard-Jones interactions were scaled to achieve agreement with the experimental osmotic coefficients. This approach has been used before to effectively take polarization effects into account for nonpolarizable forcefields and thereby avoid ion clustering.\(^{22,33}\) A reasonable scaling of \( \varepsilon_{ij} \) between cations (i) and anions (j) did not affect the osmotic coefficients significantly. Instead, \( \sigma_j \) was scaled according to \( \sigma_j = \frac{\lambda_i (\sigma_i + \sigma_j)}{2} \), using a scaling factor \( \lambda_i = 0.93 \). This scaling resulted in osmotic coefficients \( \phi = 0.96 \pm 0.12 \) (0.966), \( \phi = 1.01 \pm 0.11 \) (1.022), and \( \phi = 1.08 \pm 0.05 \) (1.080) for the 0.2, 0.6, and 1.0 m LiI solutions, respectively. The values in parentheses are the corresponding experimentally measured values.\(^{34}\)

**RESULTS**

**Phase Transition Temperatures and Ion Affinities.** Figure 1 shows the phase transition temperature of PNIPAM

![Figure 1. Lower critical solution temperature, \( T_c \), of PNIPAM (10 mg/mL) in LiI, NaI, and CsI solutions as a function of salt concentration, \( c_s \). The symbols are data points representing an average of three measurements. The error bars were calculated as sample standard deviations and are smaller than the size of the data points when not seen. The lines are guides to the eyes.](https://dx.doi.org/10.1021/acs.jpcb.0c11076)

To understand the LCST trend, all-atom MD simulations were employed. Figure 2 shows the preferential binding coefficient for indistinguishable ions (i.e., all cations and anions), \( \Gamma_{23} \), for LiI (blue line), NaI (red line), and CsI (orange line) salt solutions (1 m) as a function of the closest distance to the polymer surface. The subscript 2 stands for the polymer, and the subscript 3 for the ions, while the subscript 1 stands for water. \( \Gamma_{23} \) specifies the relation between the number of ions in the vicinity of the polymer and the statistical number in the bulk solution. A negative \( \Gamma_{23} \) value indicates ion depletion, while a positive \( \Gamma_{23} \) value indicates the favorable partitioning of ions to the polymer–water interface. LiI showed the least negative \( \Gamma_{23} \) value indicating the weakest depletion of ions, while CsI showed the most negative \( \Gamma_{23} \) value and thereby the strongest depletion of the three salts considered. The slopes observed in the LCST curves at higher salt concentration (Figure 1) are proportional to the preferential binding coefficients (Figure 2) under the assumption that \( \Gamma_{23} \) expressed per unit of solvent-accessible surface area, does not depend on the conformation of the chain.\(^{36}\) Thus, the simulation data for PNIPAM in LiI, NaI, and CsI aqueous salt solutions are in qualitative agreement with the experiments.

**Polymer–Ion Interactions.** Interaction energies were calculated in order to elucidate the effect of cations on iodide’s preferential interaction with PNIPAM, polymer–ion, polymer–water, and monomer–solution (including all ions and water molecules) interactions. These are presented in Table 1. PNIPAM–anion interactions are more favorable than PNIPAM–cation interactions. This is reflected in the radial distribution functions (RDFs) between the polymer backbone and the anions, cations, and all indistinguishable ions. The RDFs are shown in Figure 3 for all cases. The behavior of anions and cations can be seen both in the peak heights of the RDFs and in the cumulative number for the hydration shell of the 20-mer PNIPAM chain (distances up to \( r = 0.922 \text{ nm} \)). The numbers of anions in the hydration shell of the 20-mer PNIPAM chain are 5.1, 4.8, and 3.9 for LiI, NaI, and CsI,
Table 1. Total Interaction Energies (Sum of van der Waals and Coulombic Interactions) between the 20-mer PNIPAM Chain and the Iodide, the Three Different Cations, and Water Molecules for the Three Salt Solutions (1 m) \(^{14}\)

<table>
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<tbody>
<tr>
<td>LiI</td>
<td>1.45 ± 0.07</td>
<td>–0.8 ± 0.2</td>
<td>–0.417 ± 0.003</td>
<td>–142 ± 1</td>
</tr>
<tr>
<td>NaI</td>
<td>1.36 ± 0.08</td>
<td>–0.6 ± 0.2</td>
<td>–0.418 ± 0.004</td>
<td>–141 ± 1</td>
</tr>
<tr>
<td>CsI</td>
<td>1.12 ± 0.08</td>
<td>–0.09 ± 0.07</td>
<td>–0.429 ± 0.006</td>
<td>–140 ± 2</td>
</tr>
</tbody>
</table>

“...The energies were obtained by post processing the MD-simulated trajectories using the rerun option in GROMACS, where the electrostatic energies were computed using a reaction field with a dielectric constant of 78. The average interaction energies of PNIPAM with one anion, one cation, and one water molecule are reported. Monomer–solution (including all ions and water molecules) energies represent the total interaction energy between one monomer and the solution (including all water molecules and ions). The given errors are the standard deviation of the mean, \(\sigma/\sqrt{N}\), using sample standard deviation, \(\sigma\) and \(N = 10\) blocks.”

respectively. The numbers of cations are 4.5, 3.8, and 2.9, respectively. Furthermore, the PNIPAM–anion, PNIPAM–cation, and PNIPAM–water interactions are cation-specific. PNIPAM–anion interactions follow the order LiI > NaI > CsI, and the PNIPAM–cation interactions follow the order LiI ≈ NaI > CsI. The PNIPAM–water interactions follow the order CsI > NaI ≈ LiI (Table 1). These results agree with the relative ordering between the abovementioned number of ions in the PNIPAM hydration shell for each salt type. That is, more favorable PNIPAM–ion interactions (at the cost of PNIPAM–water interactions) lead to more ions in the polymer hydration shell.

Spatial probability density maps were created to investigate the spatial position of the ions around the PNIPAM side chain. A distinct preferable position (yellow area) was seen for iodide ions close to the amide NH group and the terminal methyl groups (insets in Figure 3a). The spatial probability density for the iodide ions was, however, affected by the type of cation. The probability at a certain threshold (see details in figure text) is significantly lower for CsI solution (orange box) than for LiI solution (blue box). A distinct preferable position (blue area) was also seen for lithium ions close to the amide oxygen (blue box in the inset of Figure 3b). This was not observed for cesium ions (orange box in the inset of Figure 3b) at the same threshold (see details in figure text), indicating weaker PNIPAM–Cs\(^+\) interactions. Furthermore, the probability of finding iodide ions close to the polymer side chain was higher than for the respective cation. These results are in agreement with the observed weak, but favorable, interactions between the amide oxygen and cations, between the amide NH group and iodide ions, and between the two terminal methyl groups and the iodide ions (not presented in Table 1). The interaction sites and the difference between strongly and weakly hydrated cations are in line with earlier studies of other (macro)molecules.\(^{38–42}\) However, it has been experimentally shown that an NH moiety is not necessary for anion binding.\(^ {15}\)

The abovementioned interactions between part of the PNIPAM chain and ions include an electrostatic contribution. To obtain further insight into the role of these electrostatic...
interactions, MD simulations of a “nonpolar” PNIPAM molecule without partial atomic charges were performed in LiI, NaI, and CsI solutions (1 m). The absence of favorable ion–polymer electrostatic interactions correlates with vanishing ion-specific binding to the polymer. This can be seen in Figure 4, where all three salts are depleted from the polymer surface to more or less the same extent (compare the gray lines). When the partial charges are reintroduced (blue, red, and orange lines in Figure 4), the ion-specific salting-out series (CsI > NaI > LiI), observed in the experimental data (Figure 1), is recovered. Significantly, the LiI salt is less depleted when partial atomic charges on the polymer are reintroduced. This occurs due to favorable electrostatic interactions of iodide with the amide NH and lithium with the amide oxygen. The difference (PNIPAM chain vs “nonpolar” PNIPAM chain) was larger for LiI than for NaI. This originated from LiI interacting the strongest with the polymer chain and thereby being affected the most in the absence of partial atomic charges. By contrast, CsI is more depleted when the partial atomic charges on the polymer are reintroduced. This is likely caused by favorable interactions of CsI ion pairs with the “nonpolar” PNIPAM surface. The “nonpolar” PNIPAM chains demonstrate that electrostatic interactions play a role in the ions’ preferable affinity for the PNIPAM chain.

Thermodynamic Implications of Ion Pairing. Next, cation–anion pair formation in the bulk solutions was investigated (setup V). Specifically, the fraction of contact ion pairs (CIPs) in 1 m bulk solutions was investigated by integrating the cation–anion RDFs up to the first minimum. The fractions of CIPs were 0.82 for CsI, 0.11 for NaI, and 0.07 for LiI. To investigate the influence of ion pairing in the bulk on polymer–ion interactions, MD simulations of PNIPAM chains in CsI solutions with modified ion-pairing ability (scaled cation–anion Lennard-Jones size parameter) were performed. A larger scaling factor, $\lambda_{fi}$, generated less ion pairing. Moreover, it led to less depletion from the PNIPAM chain. This can be seen by comparing the black and gray lines with the orange line in Figure 5.

To quantify the role of ion pairs on a global level, Kirkwood–Buff integrals (KBIs) and nonideality factors were calculated. The dependence of the LCST on the molar salt concentration ($c_3$) reflects how the chemical potential of the PNIPAM chain ($\mu_2$) depends on $c_3$. This dependency is provided by the Wyman–Tanford/Kirkwood–Buff relation $^{35}$

$$\left(\frac{\partial \mu_2}{\partial c_3}\right)_{p,T} = \frac{-RT \Gamma_{23}}{1 + c_3(G_{33} - G_{31})}$$

(1)

where $G_{33}$ and $G_{31}$ denote the salt–salt and salt–water KBIs, respectively, $R$ denotes the gas constant, $T$ the temperature, and $p$ the pressure. This thermodynamic relation shows that the chemical potential of the polymer increases with salt concentration when $\Gamma_{23} > 0$ (the denominator is always positive), leading to a negative slope for the LCST (Figure 1) at high salt concentration. The role of ion pairing on a global level should be reflected by the term in the denominator of this equation, $G_{33} - G_{31}$, which accounts for the nonideality of the bulk electrolyte solution. This term is reported in Table 2, together with the nonideality factor, $1 + c_3(G_{33} - G_{31})^{-1}$, and the preferential binding factors. Owing to ion pairing in bulk solution, $G_{33} - G_{31}$ is larger for CsI than for NaI (i.e., the nonideality factor is smaller, and, in particular, less than unity). The nonideality factor therefore attenuates the difference between the values of the chemical potential derivatives ($\partial \mu_2 / \partial c_3$)$_{p,T}$ of CsI and NaI obtained if only the values of $\Gamma_{23}$.
Table 2. Preferential Binding Coefficients, $G_{33}$, Corresponding to a PNIPAM 20-mer with Three Different Salts at 1 m (Setup I), Difference between Salt–Salt, $G_{33}$, and Salt–Water, $G_{31}$, Kirkwood–Buff Integrals (KBIs), and the Nonideality Factor, $[1 + c (G_{33} - G_{31})]^{-1}$, for the Three Salt Solutions at 1 m (Setup V)\(^a\)

<table>
<thead>
<tr>
<th>salt</th>
<th>preferential binding coefficient $G_{33}$</th>
<th>difference in KBIs $G_{33} - G_{31}$ [nm(^3)]</th>
<th>nonideality factor $[1 + c (G_{33} - G_{31})]^{-1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>LiI</td>
<td>$-0.8 \pm 0.2$</td>
<td>$-0.130$</td>
<td>1.14</td>
</tr>
<tr>
<td>NaI</td>
<td>$-3.6 \pm 0.5$</td>
<td>$-0.095$</td>
<td>1.10</td>
</tr>
<tr>
<td>CsI</td>
<td>$-9.2 \pm 0.4$</td>
<td>0.535</td>
<td>0.67</td>
</tr>
</tbody>
</table>

“Preferential binding coefficients and KBIs were calculated from converged running interval values between 1.5 and 2.0 nm. The KBIs were calculated with an RDF correction and a Krüger volume correction\(^{31}\) (see eq 8 in ref 44). The given errors are the standard deviation of the mean, $\sigma/\sqrt{N}$, using sample standard deviation, $\sigma$, and N = 5 blocks and propagation of uncertainty for the preferential binding coefficients. Errors for the difference in KBIs and nonideality factor were calculated as sample standard deviation using propagation of uncertainty and are smaller than the reported accuracy.

(Figure 2 and Table 2) of these two salts were considered and ideal solution behavior (i.e., equal salt–salt and salt–water affinities; $G_{33} = G_{31}$) would be assumed.

**DISCUSSION**

Stronger salting-out behavior might be expected when $\Gamma^-$ is combined with a high-charge-density cation (e.g., Li\(^+\)), and weaker salting-out behavior might be expected when combined with a low-charge-density cation (e.g., Cs\(^+\)). This expectation is based on the differences in cation hydration free energies, which indicate how easily a cation sheds its hydration shell when it approaches the polymer surface. Therefore, Li\(^+\) would be expected to be more depleted than the weakly hydrated Cs\(^+\), leading to a stronger salting-out behavior. Anions have been shown to follow this behavior.\(^{8,46-49}\) The LCST of PNIPAM in the presence of metal iodide salts shows instead the salting-out order Cs\(^+\)>Na\(^+\)>Li\(^+\) (Figure 1). This indicates that ion exclusion is subtle and charge density considerations alone cannot explain the Hofmeister effects for cations. The unexpected behavior of Li\(^+\) in view of its charge density has drawn attention before.\(^{50-52}\)

The nonlinear LCST behavior with salt concentration (Figure 1) is indicative of polymer–ion interactions.\(^{36}\) Another indication is that the observed difference in the LCST data for the different salts shows a discrepancy with ion partitioning to the air–water interface where Li\(^+\), Na\(^+\), and Cs\(^+\) are all excluded to about the same extent.\(^{53}\) While electrostatic polymer–ion interactions are crucial for the binding of ions to specific sites on the PNIPAM surface and lead to ion specific features in the preferential binding coefficients (Figure 4), the partitioning of ions to the hydration shell of the “nonpolar” PNIPAM chain instead resembles the behavior of the air–water interface. This is in line with earlier studies showing that polar groups draw cations, which are usually excluded from an air–water interface,\(^{54}\) into the interfacial region.\(^{55}\)

PNIPAM–iodide interactions are stronger than PNIPAM–cation interactions, but the cation type has an impact on all polymer–ion interactions and thereby on the dependence of the LCST on salt concentration (Figure 3 and Table 1). It should be noted that this is contrary to earlier studies that found stronger cation than anion interactions with PNIPAM. In that case, MD simulations found direct cation interactions and indirect anion interactions.\(^{56,57}\) The MD simulations provided in the current work, however, are more consistent with both LCST data and previous spectroscopic studies. In fact, iodide ions interact favorably with nonpolar surfaces, while cations need specific binding sites such as an amide oxygen. This should explain the greater iodide affinity for the polymer shown in our study. The underlying mechanisms of cation-specific effects on the PNIPAM–iodide interactions proposed in this study are depicted in Figure 6. Various experiments and MD analyses show that aqueous salt solutions do not form statistical mixtures (equally distributed ions) but rather show ion pairing and ion clustering.\(^{58}\) The more weakly hydrated the cation is, the larger is the propensity for the weakly hydrated iodide to form CIPs with it. This is in line with the law of matching water affinities.\(^{59}\) In an aqueous CsI solution, CIPs are formed due to both ions being weakly hydrated. This leads to charge neutrality for some ion pairs and a loss of the possibility for iodide ions to electrostatically interact with the amide NH group on the polymer (the left side of Figure 6). The ions are thereby repelled from the polymer–water interface, and the LCST is strongly suppressed (salting-out of PNIPAM). The degree to which ion pairing is decisive for the partitioning of ions to the polymer–water interface was demonstrated in this study (Figure 5). When Cs\(^+\) and I\(^-\) were prevented from forming electrically neutral CIPs in the simulations, weaker salt depletion occurred. In an aqueous LiI solution, hardly any CIPs were formed. Instead, the iodide ions could independently move to the polymer–water interface and interact electrostatically with the amide NH group (the right side of Figure 6). Lithium ions followed to avoid charge separation. Lithium ions also showed weakly favorable interactions with the polymer chain themselves (Table 1), namely, with the amide oxygen.

Concurrent interactions of iodide (Figure 3a) and lithium ions (Figure 3b) with PNIPAM lead to a smaller suppression of the LCST in LiI solution than in CsI solution. However, both the relative differences in the preferential binding coefficients (Figure 2) and the large differences in the number of CIPs observed in bulk solution were not reflected in the LCST curves (Figure 1), in particular not for NaI and CsI. This was due to a compensation of two effects. While ion pairing drove iodide away from the polymer and caused a decrease of $\Gamma_{23}$, it further caused a decrease in the effective concentration (activity) of the salt as expressed by the larger...
value of the denominator in eq 1 for CsI than for NaI (Figure 2 and Table 2). Ion pairing in bulk solution should therefore have two thermodynamic implications: (i) it leads to a weaker iodide interaction with the polymer, contributing to salting out; (ii) the corresponding weaker salt–water interaction attenuates the role of solvent-excluded volume60 (an effective force related to the entropy of the solvent that drives polymer compaction and hydrophobic interaction), contributing to salting-in. The observed LCST behavior with increasing concentration of iodide salts (Figure 1) is thus determined by the balance of these two effects.

Salting-in behavior is observed at low salt concentration for all salts due to the free energy favorable partitioning of weakly hydrated iodide ions to the polymer surface. At higher salt concentrations, the probability for ion pairing in the bulk is higher. This leads to a competition between PNIPAM–iodide interactions and iodide–cation interactions in the bulk. The latter dominates for weakly hydrated cations (e.g., Cs+) since they readily form CIPs and they do not show preferable interaction with the polymer. This explains why the driving force to compact the polymer chain (salting-out behavior) sets in at a lower concentration for CsI than for LiI.

The nonadditive and dependent behavior for how ions affect polymer solubility explained in this study updates earlier ideas53,61 about simple additive and independent ion-specific interactions. In addition to polymer–ion interactions, ion–ion interactions, and not only individual ion interactions, are of importance for the understanding of ion-specific effects.62 However, the effect of ion pairing on the LCST of polymers will probably depend on the type of ion pair and the identity of the cations and anions involved. The effects in this work apply to salts containing a weakly hydrated anion. For salts containing a more strongly hydrated anion, such as chloride, the occurrence of solvent-shared ion pairs together with weak, but favorable, polymer–cation interactions instead causes a mitigation of chloride’s salting-out effect when paired with a strongly hydrated cation.63

CONCLUSIONS

We have shown that the LCST behavior of PNIPAM in iodide salt solutions is affected by the specific combination of the anion and the cation. Electrostatic interactions of anions and cations with the amide group play a role in ion interactions with PNIPAM, and the type of cation affects the PNIPAM–iodide interaction. A weakly hydrated cation (e.g., Cs+) results in weaker iodide affinity, while a strongly hydrated cation (e.g., Li+) results in stronger iodide affinity. This speaks against simple additivity and provides yet another example where observed changes in polymer solubility are due to nonadditive effects. The polymer–iodide interaction is inversely correlated with anion–cation ion pairing in the bulk. As we move from the strongly hydrated cation, Li+, to the weakly hydrated cation, Cs+, the affinity for CIP formation with iodide ions in the bulk increases. Such charge neutralization leads to a larger loss in the electrostatic interactions with the polymer chain, explaining the greater decrease in the LCST for iodide salts containing Cs+ than containing Li+.

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Notes

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