Range of ion specific effects in the hydration of ions

Safr Merchant and Dilip Asthagiri
Department of Chemical and Biomolecular Engineering and The Institute of NanoBioTechnology, Johns Hopkins University, Baltimore, MD 21218

(Dated: January 27, 2013)

Within the quasichemical approach, the hydration free energy of an ion is decomposed into a chemical term accounting for ion specific ion-water interactions within the coordination sphere and nonspecific contributions accounting for packing (excluded volume) and long range interactions. The change in the chemical term with a change in the radius of the coordination sphere is the compressive force exerted by the bulk solvent medium on the surface of the coordination sphere. For the Na\(^{+}\), K\(^{+}\), F\(^{-}\), and Cl\(^{-}\) ions considered here this compressive force becomes equal for similarly charged ions for coordination radii of about 3.9 Å, not much larger than a water molecule. These results show that ion specific effects are short ranged and arise primarily due to differences in the local ion-water interactions.

Keywords: Hofmeister effect, potential distribution theorem, Monte Carlo, simulations

Numerous aqueous phase processes including, for example, the stability of biological macromolecules and colloidal suspensions, are strongly influenced by the chemical type of the inorganic ions dissolved in the aqueous phase. However, despite their ubiquity, a molecular level understanding of ion specific effects remains elusive. Ideas such as distortion of hydrogen bond network of the bulk solvent, balance between ion-water and water-water interactions, and influence of ion-specific dispersion forces have been proposed to rationalize these effects.

Understanding how the ion affects the solvent matrix remains of first interest in understanding ion specific effects. Recent spectroscopic measurements on salt solutions suggest that the ion only affects hydrogen bonds of water molecules within its first hydration shell. An earlier theory and simulation study by us also showed that only a small subset of water molecules in the first hydration shell of the ion are sensitive to the type of the ion. These results taken together suggest that ion specific ion-solvent interactions are limited only to the ion’s local neighborhood. In this Communication, we address the question of how far in the liquid the chemical type of the ion is felt and if the point at which ion-specificity is lost is also the point where continuum models of hydration begin to take hold.

To address these questions, we first define a coordination sphere of radius r around the ion to separate the ion interaction with water molecules within the coordination shell from the longer-range interaction of the ion with the solvent outside the coordination sphere. With such a spatial partitioning, the hydration free energy of the ion, \(\mu_{\text{ex}}\), can be written as

\[ \mu_{\text{ex}} = k_{\text{B}}T \ln x_{0}(r) - k_{\text{B}}T \ln p_{0}(r) + \mu_{\text{ex}}^{\text{outer}}(r) \]  (1)

where \(k_{\text{B}}\) is Boltzmann’s constant and \(T\) is the temperature, \(k_{\text{B}}T \ln x_{0}(r)\) is the chemical contribution to the hydration free energy, \(-k_{\text{B}}T \ln p_{0}(r)\) is the packing contribution, and \(\mu_{\text{ex}}^{\text{outer}}(r)\) is the contribution to the hydration free energy due to ion interaction with the solvent outside the coordination shell. \(x_{0}(r)\) is the probability of observing no water molecules within the coordination sphere of radius r around the ion; thus \(k_{\text{B}}T \ln x_{0}(r)\) is the free energy gained by allowing water molecules to flood the empty coordination sphere, justifying the identification of this term with local, chemical interactions. \(p_{0}(r)\) is the probability of observing an empty coordination sphere, but in the absence of the ion: \(-k_{\text{B}}T \ln p_{0}(r)\) is the free energy required to create a cavity of radius r in the bulk solvent. \(\mu_{\text{ex}}^{\text{outer}}(r)\) is the free energy gained in placing the ion in an empty cavity: it accounts for the interaction of the ion with the bulk medium outside the coordination sphere. For a sufficiently large \(r\), we expect \(\mu_{\text{ex}}^{\text{outer}}(r)\) to be composed of a large number of small non-specific contributions. Since Eq. (1) is a tautology, the point where \(\mu_{\text{ex}}^{\text{outer}}(r)\) becomes insensitive to ion type is also the range to which \(k_{\text{B}}T \ln x_{0}(r)\) captures all the ion-specific effects.

To identify the range of ion specific effects, we studied ion-water systems under NVT conditions using Metropolis Monte Carlo simulations. Water was modeled using the SPC/E model and parameters for Na\(^{+}\), K\(^{+}\), F\(^{-}\), Cl\(^{-}\) ions were obtained from an earlier study. The chemical term was estimated by growing the coordination sphere in steps of 0.05 Å. At any \(r\), the probability of observing zero (0) water molecules in a coordination sphere of radius \(r + 0.05\) Å was obtained. From this data, the entire \(x_{0}(r)\) versus \(r\) profile was reconstructed. A similar strategy was followed for \(p_{0}(r)\) versus \(r\), but with a step size of 0.1 Å. At each \(r\) value, \(3 \times 10^{5}\) sweeps of equilibrium were followed by \(3 \times 10^{5}\) sweeps of production. Each sweep comprised one attempted move of each particle in the system. The maximum allowed translation and rotation were adjusted during the equilibration phase to target an acceptance ratio of 0.3. In the production phase these maximum displacement values were held fixed. Configurations are saved every 10 sweeps for analysis.

From Fig. I we observe that for \(r > 3.9\) Å the change in the chemical term depends only on the charge of the ion and is independent of its chemical nature. Fig. I emphasizes that, for the ions studied here, ion-specific
The left hand side of the above equation is the compressive force of the bulk on the coordination sphere of the ion relative to the compressive force on an ion-free coordination sphere.

For a sufficiently large \( r \), we expect the interaction of the ion with the bulk medium outside the coordination sphere to be well-described by a Gaussian. In this case, for an ion of charge \( q \), we have

\[
\mu_{\text{outer}}^{\text{ex}} = q \langle \phi \rangle_0 - \frac{\beta}{2} q^2 \langle (\phi - \langle \phi \rangle_0)^2 \rangle_0,
\]

where \( \langle \ldots \rangle_0 \) denotes averaging in the presence of an uncharged ion, \( \langle \phi \rangle_0 \) is the potential at the center of an uncharged ion, and \( \langle (\phi - \langle \phi \rangle_0)^2 \rangle_0 \) is the fluctuation in the electrostatic potential. By defining \( 1/r_{\text{Born}} = \beta \langle (\phi - \langle \phi \rangle_0)^2 \rangle_0 / 2 \), it is readily seen that the fluctuation contribution is just the Born model of hydration. (For clarity, in Eq. 3 we do not explicitly indicate the corrections due to electrostatic self-interaction, as these depend solely on \( q \) and not on the ion type.)

In the radius range of interest we find that \( \partial \langle \phi \rangle_0 / \partial r \) is about a \( k_B T/\AA \). (A similar estimate is obtained based on the data in Ref. 21 for a cavity.) Neglecting this contribution, we thus expect that asymptotically

\[
k_B T \partial \ln x_0 / \partial r \sim -q^2 \left( 1 - \frac{1}{\epsilon} \right),
\]

where \( \epsilon \) is the dielectric constant for water.

Figure 2 shows that for \( r > 3.9 \) \( \AA \) the relative compressive force between similarly charged ions becomes equal, but is different for anions and cations. The large \( r \) value is also different from the value obtained using the Born model, where the Born-radius is set equal to the coordination radius.

The discrepancy in the asymptotic limit between simulation data and that based on the Born model is not particularly surprising, since the coordination radius is not the same as the Born radius. For example, the position of the water molecule is denoted by the position of its oxygen atom and thus the coordination shell strictly excludes only the oxygen atoms of the water molecules but not their corresponding hydrogen atoms, which can enter the coordination sphere. Since this effect will be more pronounced for the anions than the cations, we expect that the equivalent Born radius for anions to be smaller than that for the cations. What is interesting, however, is the observation that by empirically reducing the cavity radius by 1.3 \( \AA \) for anions and 0.5 \( \AA \) for cations (Fig. 3), the relative compressive force for both anions and cations is similar for large coordination radii. Further, the large-\( r \) curvature of the radially translated curves compares reasonably well with that based on the Born model of hydration. (We maintain the difference of 0.8 \( \AA \), as earlier studies on such shifts find a sim-
FIG. 3: The data in Fig. 2 have been shifted along $r$ by $-1.3$ Å for the anions and by $-0.5$ Å for the cations. With this empirical shift, the compressive force $F = k_B T \frac{\partial \ln \rho(r)}{\partial r}$, Eq. 2, follows the trend predicted by the Born model (Eq. 4).

In summary, for the ions studied here, we find that a coordination radius of about 3.9 Å, not much larger than the size of a water molecule, is sufficient to account for all ion specific ion-water interactions. Our results are in accordance with experimental results that suggest that the influence of the ion on hydrogen bonding between water molecules is minimal for water molecules beyond the first hydration shell. The short-range of ion specific ion-water interactions further suggests that any framework for modeling ion-specific effects needs to acknowledge the molecular characteristics of local ion-water interactions.

* Corresponding author: Fax: +1-410-516-5510; Email: dilipa@jhu.edu

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