

Isotope effect reveals non-cooperative water dynamics in salt solutions

May 28 2018

Water molecules surrounding ions behave in a much less cooperative way than they do in bulk water. This follows from a study on the isotopedependent dielectric response of salt solutions, which has just been published in *Physical Review Letters* by researchers from the Amsterdam research institutes HIMS and AMOLF. Their results lead to an update of Nobel-laureate Onsager's 40-year old theory for the response of salt solutions to electric fields, and enables a reliable determination of hydration numbers that play a key role in chemistry and biophysics.

Solutions of ions in water are ubiquitous in physics, chemistry and biology, and the complicated way in which ions influence the hydrogenbond network of water has been the subject of intensive experimental and computational research. Ions in <u>solution</u> interact with the water <u>molecules</u> surrounding them, thereby reducing the dielectric polarizability of an ionic solution as compared to that of neat water. This reduction arises from two effects: the rotational immobilization of water molecules directly binding to the ions (static contribution), and the longrange dynamical response of water molecules to the moving ions (kinetic contribution).

The separation of these two contributions to the reduction of the dielectric polarizability has always been a challenge in research on ionic solutions. Currently, the standard approach is to use the theoretical expression for the kinetic contribution derived by Onsager and his co-worker Hubbard, which was published in a paper in 1977 that has become a classic in the field.



Putting the model to the test

Researchers from the University of Amsterdam's Van 't Hoff Institute for Molecular Sciences and the Amsterdam-based AMOLF research institute have now developed an experimental method to determine the kinetic contribution. In this method, they compare the dielectric polarizability of a solution of ions in ordinary water (H2O) with the response of the same ions dissolved in heavy water (D2O). As the static contribution to the reduction of the <u>dielectric</u> polarizability will be the same in H2O and D2O, this comparison provides direct information on the kinetic contribution. In this way they could test Hubbard and Onsager's model, and found that the experimentally observed kinetic contribution is much less than theoretically predicted. The researchers explain the discrepancy from a reduced cooperativity of the motion of water molecules surrounding ions—in contrast to the theory that assumes a uniform degree of cooperativity for all <u>water</u> molecules, independent of their surroundings.

The team proposes a simple modification of Hubbard and Onsager's theory to take the locally reduced cooperativity into account. This modification makes it possible to determine ionic hydration numbers in a reliable manner, and avoids the unphysical (negative!) hydration numbers that are sometimes obtained when using the original Hubbard-Onsager theory. The results will thus be of practical use, since hydration numbers are commonly used in chemistry and biophysics to characterize salt solutions.

More information: Roberto Cota et al. Evidence for Reduced Hydrogen-Bond Cooperativity in Ionic Solvation Shells from Isotope-Dependent Dielectric Relaxation, *Physical Review Letters* (2018). DOI: <u>10.1103/PhysRevLett.120.216001</u>



Provided by University of Amsterdam

Citation: Isotope effect reveals non-cooperative water dynamics in salt solutions (2018, May 28) retrieved 25 April 2024 from https://phys.org/news/2018-05-isotope-effect-reveals-non-cooperative-dynamics.html

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