Time-resolved terahertz–Raman spectroscopy reveals that cations and anions distinctly modify intermolecular interactions of water

Vasileios Balos, Naveen Kumar Kaliannan, Hossam Elgabarty, Martin Wolf, Thomas D. Kühne and Mohsen Sajadi

The solvation of ions changes the physical, chemical and thermodynamic properties of water, and the microscopic origin of this behaviour is believed to be ion-induced perturbation of water’s hydrogen-bonding network. Here we provide microscopic insights into this process by monitoring the dissipation of energy in salt solutions using time-resolved terahertz–Raman spectroscopy. We resonantly drive the low-frequency rotational dynamics of water molecules using intense terahertz pulses and probe the Raman response of their intermolecular translational motions. We find that the intermolecular rotational-to-translational energy transfer is enhanced by highly charged cations and is drastically reduced by highly charged anions, scaling with the ion surface charge density and ion concentration. Our molecular dynamics simulations reveal that the water–water hydrogen-bond strength between the first and second solvation shells of cations increases, while it decreases around anions. The opposite effects of cations and anions on the intermolecular interactions of water resemble the effects of ions on the stabilization and denaturation of proteins.

Ions are ubiquitous in nature and their solvation is of fundamental importance in chemical and biological reactions. They affect the folding and unfolding of proteins and enzymes, they are responsible for the transmission of neural signals, they affect chemical equilibria and they define the efficiency of electrochemical reactions. In most if not all of these processes, strong interactions between ions and water molecules are believed to play a central role. Due to electrostatic interactions, ions perturb the dynamics and the local structure of the hydrogen-bonding network of water; although the extent of these effects is not yet fully understood. Ions are typically categorized into structure makers and breakers. Relative to the strength of water–water interactions, the structure makers stabilize the water structure due to their high surface charge density (SCD) and the structure breakers, consisting of ions with low SCD, interact weakly with water molecules and promote disorder in the hydrogen-bonding network of water. However, this SCD-based categorization neglects the polarity of ions. Moreover, the distinct solvation mechanisms of cations and anions are not grounded on a comprehensive, molecular-level description.

Extensive theoretical and experimental studies have been conducted to gain microscopic insight into ion solvation. However, due to the complexity of the underlying processes and the selective sensitivity of the employed methods, the pictures that have emerged are still elusive and in many cases contradictory. Some methods are inherently more sensitive to either the effect of anions or that of cations. For instance, ultrafast vibrational spectroscopy, in which the stretch vibration of water’s hydroxyl group is used as a local probe to measure the orientational correlation time of water molecules, is more sensitive to anionic effects. Employing this technique, Bakker and co-workers resolved slow hydrogen-bonding dynamics of water molecules that hydrate anions and found almost no effect on the reorientational dynamics of water molecules surrounding cations. In contrast, dielectric relaxation spectroscopy, which probes the dynamics of the molecular permanent dipole moment, showed only local ionic effects, limited to the water molecules in the first solvation shell of the cations. On the other hand, neutron scattering resolved the extended distortion of the hydrogen-bonding network and showed structural modifications similar to the impact of pressure on water, but with marginal effect on the hydrogen bonds between water molecules. Interestingly, methods that directly interrogate the hydrogen-bonding network dynamics of water, such as terahertz (THz) and low-frequency Raman spectroscopies, turned out to be triumphant in revealing further details of ion solvation. For the sub-1 THz region, Netz and co-workers theoretically predicted and later experimentally resolved the ion–water and ion–ion correlation signatures, connecting the ion mobility to the macroscopic electrolyte conductivity. Using high-frequency THz spectroscopy, Havenith and co-workers resolved resonances due to the vibrational motion of ions inside their water cages. The isotropic optical Kerr effect spectroscopy by Meech and co-workers resolved a hydrogen-bond vibrational mode, formed between anions and their surrounding water molecules. Anisotropic optical Kerr effect spectroscopy by Wynne et al. revealed the jamming of the translational motion of water molecules in strong cationic solutions.

The aforementioned methods measure the ensemble-averaged two-point time-correlation function \( \langle A(0), A(t) \rangle \), with \( t \) being time and \( A \) being the dipole moment vector in THz spectroscopy, the polarizability tensor elements in Raman spectroscopy and the particle density in neutron scattering. More recently, Hamm and co-workers introduced a combined Raman–THz spectroscopy, which measures the three-point cross-correlation of the dipole moment vector and the polarizability tensor \( \Pi \), that is, \( \langle \Pi(t_1), \mu(t_2), \mu(t_3) \rangle \). They demonstrated the heterogeneity of the local structure of...
Articles Nature Chemistry

Thus, by form effectively give rise to a two-time-point response function of the selective polarizability. Although the latter field-matter interactions tem is resolved by an optical–Raman pulse interacting with its col-

Upon excitation of the rotational degrees of freedom of water medium. The effect is monitored by an optical

Here we employ a similar approach and study the nonlinear THz response of aqueous ion solutions (Fig. 1a). In contrast to Hamm’s approach, we induce the nonlinear effect with an intense THz pulse via a two-electric-field interaction with the collective permanent dipole moment of the liquid. The response of the system is resolved by an optical–Raman pulse interacting with its collective polarizability. Although the latter field-matter interactions effectively give rise to a two-time-point response function of the form $\langle \mu(0), \mu(t), J(l) \rangle$, the hybrid nature of the interactions provides microscopic insight into the molecular processes. Thus, by selective excitation of an intermolecular mode/process and probing the response of low-frequency Raman-active intermolecular modes of the liquid, we are able to trace in real time the dissipation of the deposited THz energy into the intermolecular degrees of freedom of water.

Using this technique, we recently demonstrated that in liquid water under ambient conditions, a single-cycle pulse centred at ~1 THz resonantly dumps the majority (85%) of its energy into molecular rotations. The deposited energy is transferred quickly (faster than ~300 fs) to the collective translational motions, due to the strong coupling between the intermolecular rotational and translational dynamics in the hydrogen-bonding network of water (Fig. 1b and Supplementary Fig. 1).

In the current study, we use the latter rotational–translational coupling as an intermolecular probe to interrogate the ion-induced perturbations of the hydrogen-bonding network of water. For a systematic study, we select salts that contain a strongly charged cation or anion. The respective counterions are chosen to be almost ‘neutral’; that is, counterion–water and water–water interaction strengths are comparable. The cations of choice are potassium ($K^+$), sodium ($Na^+$), lithium ($Li^+$) and magnesium ($Mg^{2+}$) with chloride ($Cl^-$) as their counterion. The anions are sulfate ($SO_4^{2-}$) and carbonate ($CO_3^{2-}$) with $Na^+$ as the counterion, and fluoride ($F^-$) with $K^+$ as its counterion, for higher solubility. We have also studied the response of $MgSO_4$ solutions in which both the cation and the anion are strongly charged. The selected ions have a relatively small electric polarizability, a criterion imposed for reducing potential artefacts emanating from direct contribution of the ion polarizability into the measured signals.

**Results**

Experiments are performed in the transient THz Kerr effect (TKE) configuration by which the THz electric-field-induced optical birefringence is measured (Methods). The resulting TKE signals of all solutions are displayed in Fig. 2 and Supplementary Fig. 4. In both panels the signals are compared with that of pure water. In all signals, two main contributions can be discerned: (1) a coherent instantaneous electronic response and (2) a flipped relaxation tail, which contains information on the nuclear response of the solutions. The amplitude of the latter contribution is highly sensitive to the ionic content of the solutions; it is enhanced by the solutions of chloride salts with strong cations and reduced by solutions of sodium salts (and KF) with strong anions. Moreover, the relaxation tail of all signals can be fit mono-exponentially with decay times in the range ~0.5–0.8 ps (Supplementary Figs. 2, 3 and 5). Additionally, we have measured the THz pump power dependence of the TKE response of pure water and $MgCl_2$ solution. As shown in Supplementary Fig. 6, the TKE signals scale with the square of the THz electric field.

In Fig. 3a,b, we compare the TKE signal amplitudes of all the ionic solutions against the salt concentration and the gas-phase SCD, respectively. Interestingly, for strong cations, the signal amplitudes linearly scale with both the salt concentration and the SCD. For strong anions, a similar trend but with an opposite sign is observed. In general, the slope of the linear fit to the signal amplitudes is steeper for anions than for cations, a possible indication of a stronger anionic versus cationic effect in water. Note also that the TKE signal amplitude of LiCl versus salt concentration exhibits a discernible deviation from linearity (Fig. 3a), likely due to the light mass, high charge density and nuclear quantum effects of $Li^+$ (refs. 43,44). Similarly, the smallest and lightest anion, namely $F^-$, manifests a marked deviation from the other anions in Fig. 3b (refs. 34,46).

The calculated THz electric-field-induced polarizability anisotropies $\Delta J(l)$ of $MgCl_2$ (at 1 M, 2 M and 4 M) and $Na_2SO_4$ (at 1 M) are shown in Fig. 4a,b. The results are obtained from the molecular dynamics (MD) trajectories, using the atomic multipole optimized energetics for biomolecular applications (AMOEBA) polarizable force field and the extended-dipole-induced-dipole (XDD) model as described by Torri (refs. 48,49), but modified to include the second hyperpolarizability (Methods). Interestingly, relative to pure water, the $\Delta J(l)$ amplitude of $MgCl_2$ is enhanced, while that of $Na_2SO_4$ is reduced. The amplitude drop of the $Na_2SO_4$ solution is modest and not as large as that observed in the experiment, likely due to limitations originating from the employed XDD model (Methods) or the employed force field. The calculated $\Delta J(l)$ for $NaCl$ (at 2 M and 4 M) replicates the behaviour of $MgCl_2$ (Supplementary Fig. 7).
Δ \Pi \propto \langle \alpha \rangle \Delta \alpha \theta 

In the following, we show how the changes in the TKE signal amplitude and the corresponding simulated quantity $\Delta I(t)$ are related to the modification of water–water intermolecular interactions.

Discussion

In the TKE experiment, due to the action of the pump field polarized along $x$ (Fig. 1a), the probe pulse, polarized at 45° relative to $x$, encounters a transient difference $\Delta n = n_x - n_y$ between the refractive indices along the $x$ and $y$ directions. The resulting birefringence is given by \cite{43, 49}

$$\Delta n \propto \langle \Delta \Pi_{xx} - \Delta \Pi_{yy} \rangle, \quad (1)$$

where $\Delta \Pi_{ij}$ is the pump-induced change in the collective electronic polarizability tensor element $\Pi_{ij}$. Here, $\Pi$ refers to the liquid phase and contains contributions from interactions/collisions between molecules in the condensed phase. The variation $\Delta \Pi$ can, in principle, be written as a sum $\Delta \Pi^{\text{int}} + \Delta \Pi^{\text{col}}$, whose two contributions arise, respectively, from the intrinsic gas-phase molecular polarizability ($\Pi^{\text{int}}$) and the intermolecular interactions and collisions ($\Pi^{\text{col}}$) in the condensed phase\cite{39}.

The $\Delta \Pi^{\text{col}}$ characterizes the degree of anisotropy of the unperturbed $\Pi$ and is usually labelled $\Delta \alpha$ for single molecules. Averaging $\Delta \alpha$ over all molecules according to equation (1) yields an expression for single-molecule rotational birefringence $\Delta n_{\text{rot}}$ that scales with the degree of molecular alignment $\langle P_r(\cos \theta) \rangle = (3 \cos^2 \theta - 1)/2$, with $\theta$ being the angle between the molecular dipole vector and the polarization axis of the THz pump pulse, and the molecular polarizability anisotropy $\Delta \alpha$, that is, $\Delta n_{\text{rot}} \propto \Delta \alpha \langle P_r(\cos \theta) \rangle$. The averaged $\Delta \Pi^{\text{col}}$ makes another contribution to the transient birefringence and arises directly or indirectly from the pump-induced changes in the collision-induced polarizability. Additionally, THz electric-field-induced ionization and charge separation may contribute to the total polarizability of water\cite{11, 12}. However, as demonstrated by Elsässer and co-workers, the latter contributions

---

**Fig. 2** | TKE response of aqueous salt solutions. a. Terahertz-driven transient optical birefringence of a series of salts with strong cations and Cl$^-$ as their counterion, at 4 M concentration. The signal amplitude is cation specific and increases with SCD: MgCl$_2$ > LiCl > NaCl > KCl > H$_2$O. b. The same as a but for strong anions at their maximum concentrations with K$^+$ or Na$^+$ as the counterion. The signal amplitude is anion specific and decreases with SCD: Na$_2$CO$_3$ > Na$_2$SO$_4$ > KF > H$_2$O. MgSO$_4$ salt is composed of a strong cation and a strong anion; hence its birefringence signal at 2 M concentration is added to both panels (dashed blue line). Red arrows indicate the TKE amplitude variation with respect to that of pure water.

**Fig. 3** | TKE amplitude dependence on concentration and SCD. a. The amplitude of the TKE signals, normalized (norm.) to that of pure water, scales linearly with the salt molar concentrations. b. The amplitude of the TKE signals relative to that of pure water scales linearly with respect to the gas-phase SCD of the salts. The TKE amplitude of SO$_4^{2-}$ or CO$_3^{2-}$ at 4 M is obtained by extrapolating the TKE amplitudes to higher concentrations. As MgSO$_4$ bears two highly charged ions, it is not included in b. Error bars indicate the standard deviations of the mean determined from the corresponding TKE signals at negative pump-probe delays. The solid lines are linear fits to the data points.
modulate the isotropic part of the complex dielectric response of water, with negligible impact on its anisotropic response, measured in the TKE experiment\cite{44,45}.

As summarized schematically in Fig. 1b, the THz energy deposited into the rotational degrees of freedom of water is transferred into the translational motion of the neighbouring water molecules\cite{44,45}. In a previous paper\cite{44}, we showed that the decay of the TKE signal's tail reveals the relaxation of the translational motion of water molecules, most likely that of the hydrogen-bond bending mode due to its spectral proximity to the excitation frequency (Supplementary Figs. 1–5). Therefore, given the initial excitation of the rotational dynamics of water with the THz pump pulse, the rotation-to-translation energy transfer causes a coherent increase in the rate of the collisions between water molecules, which can be resolved as a change in the refractive index of the liquid via $\Delta \Pi$. 

In light of the TKE response of pure water, the ion-induced enhancement/weakening of the TKE signals may indicate underlying changes in the strength of the intermolecular water–water hydrogen-bonding interactions, by which the rotational–translational coupling is modified. To examine the soundness of this proposition, we first decompose $\Delta I$ into the contributions $\Delta I^{\Pi}$ and $\Delta I^T$. The corresponding results for MgCl$_2$ at 4 M, NaCl at 4 M and Na$_2$SO$_4$ at 1 M are shown in Fig. 4 and Supplementary Fig. 7. Remarkably, while in MgCl$_2$ and NaCl solutions the collision-induced polarizability anisotropy $\Delta I^T$ makes the dominant contribution to the total polarizability anisotropy, its contribution in the Na$_2$SO$_4$ solution is minor.

Along the same line, the increase/decrease of the collision-induced polarizability may also be observed in the kinetic energy (KE) of the translational motion of water molecules. Thereby, we calculate the transient excess translational KE(t) of water molecules in MgCl$_2$ and Na$_2$SO$_4$ solutions. As shown in Fig. 5, the translational KE(t) of water molecules in MgCl$_2$ solutions is substantially enhanced relative to pure water. As displayed in the inset of the same figure, the enhancement scales linearly with the salt concentration. The KE(t) of the water molecules in NaCl solutions also shows a similar trend (Supplementary Fig. 8). For the Na$_2$SO$_4$ solution, although the expected decrease in the translational KE(t) is not observed, its amplitude is not as large as that of the MgCl$_2$ solution at the same concentration. This parallels our previous observation that the employed parameterization of the polarizable force field underestimates the effects of Na$_2$SO$_4$ in the polarizability calculations.

To connect the latter findings with the local intermolecular interactions in water, we resort to equilibrium ab initio MD (AIMD) simulations and calculate the water–water hydrogen-bond strength. For quantifying the strength of the hydrogen bonds, we employ...
energy decomposition analysis for condensed phase systems based on absolutely localized molecular orbitals (ALMO-EDA) within Kohn–Sham density functional theory (Methods). The hydrogen bonds are categorized into four disjoint classes: those between the first and second solvation shells of an anion; the same for cations; those involving one water molecule, which is simultaneously shared in the first solvation shell of an anion and a cation; and those between the remaining bulk-type water molecules, meaning that these hydrogen bonds are between two water molecules, both of which are not in the first solvation shell of an ion.

As shown in Fig. 6, by comparing the water–water hydrogen-bond strength around ions with that of the bulk-type water, the following conclusions can be drawn: In MgCl₂ solution, the Mg²⁺ ions with their high charge density lead the water–water hydrogen bonds in their immediate surroundings to be 2–3 kJ mol⁻¹ stronger than more distant hydrogen bonds. This is in line with a recent experimental study by Shalit et al. denoting the ability of strong cations such as Mg²⁺ to structure the hydrogen-bonding network of water. On the other hand, in the Na₂SO₄ solution, the SO₄²⁻ ions show an opposite trend, namely the neighbouring hydrogen bonds are slightly weaker. Along the same lines, by analysing the changes in the OH-stretch vibration frequency of water molecules in the first and second solvation shells of CO₃²⁻, Yadav et al. reported the weakening of the water–water hydrogen-bonding strength around CO₃²⁻ (ref. 17). Previous ab initio calculations on solvated clusters have also shown this increase (decrease) of the hydrogen-bond strength between the first and second solvation shells of cations (anions) ¹⁸.

Thereby, the emerging picture thus far is as follows. Ions influence the water–water intermolecular hydrogen-bond strength between the first and second solvation shells. Highly charged cations strengthen this interaction and highly charged anions weaken it. The ion-induced change in the water–water interaction strength alters the intermolecular coupling between the rotational and translational degrees of freedom. As such, the THz energy initially deposited into the intermolecular rotational motion of water funnels into the translational motions, with the efficiency determined by the ionic content of the solution, and influences the collision-induced contribution to polarizability anisotropy, which is captured in our TKE experiment. We now consider the THz response of MgSO₄, a system in which both the anion and the cation are strongly charged, to investigate whether the impact of two strong ions coexisting in a solution is additive or cooperative. As shown in Figs. 2 and 3a and Supplementary Fig. 4, the TKE signal amplitude of MgSO₄ relative to the signal amplitudes of MgCl₂ and Na₂SO₄ displays an intermediate behaviour: in contrast to Na₂SO₄, the TKE signal of MgSO₄ is larger than the signal of pure water, while it is smaller than that of MgCl₂. Moreover, as shown in Fig. 6, the water–water hydrogen-bond strength around Mg²⁺ and SO₄²⁻ in MgSO₄ solution differs from that in MgCl₂ and Na₂SO₄ solutions. For MgSO₄, the average of all ion-mediated contributions is larger than the bulk-type water–water hydrogen-bond strength, signifying its ‘cationic character’.

Notably, this indicates a non-additive but cooperative effect of ions in MgSO₄ solution²⁶,⁵⁹, since in solutions with ‘neutral’ counterions, SO₄²⁻ shows a larger effect compared to Mg²⁺, as seen in Fig. 3a.

In light of these results, the linear concentration dependence of the TKE signal amplitude in strong cationic solutions, holding even at elevated concentrations, may now be understood. In particular, in the extreme case of the MgCl₂ solution at 4 M, all water molecules are consumed in the first solvation shells of ions; hence, there is virtually no free water to form a second hydration shell for each ion. Nevertheless, we still observe a linear trend in the concentration dependence of the experimental, as well as the calculated,
quantities (Fig. 3a and insets in Figs. 4a and 5). We believe that in such solutions, the role of Cl⁻ in water–water interactions is critical. The water–Cl⁻ interaction strength is comparable with that of water–water in bulk (Fig. 6), and the dynamics of water molecules solvating Cl⁻ are minimally perturbed by Cl⁻ (ref. 49). Thereby, there are still abundant water molecules in the system whose dynamics and hydrogen-bond strengths resemble that of bulk, by which the second solvation of Mg²⁺ at high concentrations can be formed.

To summarize, our joint TKE experiment and MD simulations in a large set of aqueous ionic solutions revealed that strong cations enhance the total polarizability anisotropy of water, whereas strong anions weaken this effect, relative to that in pure water. The polarizable classical MD simulations successfully reproduce the experimental results and further show that the collision/interaction-induced polarizability is enhanced in the presence of strong cations and weakened by strong anions. The origin of these effects is attributed by AIMD simulations to the changes in the water–water hydrogen-bond interaction between the first and second solvation shells of ions; strong cations make hydrogen bonds stronger, while strong anions weaken them.

Notably, although ions with high SCD, irrespective of their polarity, interact strongly with their immediate surrounding water molecules and structure water in their first solvation shells (refs 16, 17), the hydrogen-bond strength between the first and second solvation shells depends not only on the SCD but also on the polarity of the ions. This subtle difference distinguishes our findings from the traditional SCD-based categorization of ions into structure makers and breakers, as discussed in the introduction.

Finally, the opposite impact of strong anions and cations on the water–water intermolecular interactions is in line with the impact of ion polarity on the stabilization and denaturation of proteins’ tertiary structure in aqueous solutions (that is, the Hofmeister effect) (refs 18, 19). While strongly hydrated cations denature proteins, mainly by direct preferential binding to the polar amide groups or by pairing with the protein’s negatively charged carboxylate groups (refs 20, 21), the mechanism of protein stabilization by strongly hydrated anions remains elusive. Our findings regarding the weakening of the water–water hydrogen-bond strength in the presence of strong anions may support the mechanism suggested by Collins et al., according to which the anion-induced protein stabilization is water mediated and attributed to the ‘interfacial effects of strongly hydrated anions near the surface of proteins’ (ref. 22). Less efficient energy transfer in water and weaker water–water intermolecular interactions may result in converting the liquid into a ‘less good solvent’ by strong anions, which may drive the proteins to minimize their solvent-accessible surface area by folding (ref. 23).

Online content
Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41557-022-00977-2.

Received: 31 August 2021; Accepted: 16 May 2022;
Published online: 30 June 2022

References
1. Collins, K. D., Neilson, G. W. & Enderby, J. E. Ions in water: characterizing the forces that control chemical processes and biological structure. Biophys. Chem. 128, 95–104 (2007).
2. Marcus, Y. Effect of ions on the structure of water. Pure Appl. Chem. 82, 1889–1899 (2010).
3. Yang, Z. Hofmeister effects: an explanation for the impact of ionic liquids on biocatalysis. J. Biotechnol. 144, 12–22 (2009).
4. Savitchenko, L. P., Poo, M. M. & Rassakov, D. A. Electrodiffusion phenomena in neuroscience: a neglected companion. Nat. Rev. Neurosci. 18, 598–612 (2017).
5. Balos, V. et al. Specific ion effects on an oligopeptide: bidentate binding matters for the guanidinium cation. Angew. Chem. Int. Ed. 58, 332–337 (2019).
6. Ózvegy-Latos-Schtädtler, I. et al. Beyond the Hofmeister series: ion-specific effects on proteins and their biological functions. J. Phys. Chem. B 121, 1997–2014 (2017).
7. Ma, D. & Jan, L. Y. ER transport signals and trafficking of potassium channels and receptors. Curr. Opin. Neurobiol. 12, 287–292 (2002).
8. Oldham, H. B. & Myland, J. C. Fundamentals of Electrochemical Science (Academic Press, 1993).
9. Roosen-Runge, F., Heck, B. S., Zhang, F., Kohlbacher, O. & Schreiber, F. Interplay of pH and binding of multivalent metal ions: charge inversion and reentrant condensation in protein solutions. J. Phys. Chem. B 117, 5777–5787 (2013).
10. Bopp, G. E. et al. Metal oxide surfaces and their interactions with aqueous solutions and microbial organisms. Chem. Rev. 99, 77–174 (1999).
11. Zhao, R., Biesheuvel, P. M., Miedema, H., Bruning, H. & van der Wal, A. Charge efficiency: a functional tool to probe the double-layer structure inside of porous electrodes and application in the modeling of capacitive deionization. J. Phys. Chem. Lett. 1, 205–210 (2010).
12. Jungwirth, P. & Laugé, D. Ion-induced long-range orientational correlations in water: strong or weak, physiologically relevant or unimportant, and unique to water or not? J. Phys. Chem. Lett. 9, 2056–2057 (2018).
13. Chen, Y. et al. Electrolytes induce long-range orientational order and free energy changes in the H-bond network of bulk water. Sci. Adv. 2, e1501891 (2016).
14. Borgia, D., Belloni, L. & Levesque, M. What does second-harmonic scattering measure in diluted electrolytes? J. Phys. Chem. Lett. 9, 3698–3702 (2018).
15. Morita, T., Westh, P., Nishikawa, K. & Koga, Y. How much weaker are the effects of cations than those of anions? The effects of K⁺ and Cs⁺ on the molecular organization of liquid H₂O. J. Phys. Chem. B 118, 8744–8747 (2014).
16. Hribar, B., Southall, N. T., Vlachy, V. & Dill, K. A. How ions affect the structure of water. J. Am. Chem. Soc. 124, 12302–12311 (2002).
17. Remying, R. C. et al. Water lone pair delocalization in classical and quantum descriptions of the hydration of model ions. J. Phys. Chem. B 122, 3519–3527 (2018).
18. Shali, A., Ahmed, S., Savolainen, J. & Hamm, P. Terahertz echoes reveal the inhomogeneity of aqueous salt solutions. Nat. Chem. 9, 273–278 (2017).
19. Zangi, R. Can salting-in/salting-out ions be classified as chaotropes/kosmotropes? J. Phys. Chem. B 114, 643–650 (2010).
20. Soper, A. K. & Weckström, K. Ion solvation and water structure in potassium halide aqueous solutions. Biophys. Chem. 124, 180–191 (2006).
21. Paschek, D. & Ludwig, R. Specific ion effects on water structure and dynamics beyond the first hydration shell. Angew. Chem. Int. Ed. 50, 352–353 (2011).
22. Vila Verde, A., Santer, M. & Lipowsky, R. Solvent-shared pairs of densely charged ions induce intense but short-range supra-additive slowdown of water rotation. Phys. Chem. Chem. Phys. 18, 1918–1930 (2016).
23. Buchner, R., Chen, T. & Hefter, G. Complexity in “simple” electrolyte solutions: ion pairing in MgSO₄(aq). J. Phys. Chem. B 108, 2365–2375 (2004).
24. Buchner, R., Hefter, G. & May, P. M. Dielectric relaxation of aqueous NaCl solutions. J. Phys. Chem. A 103, 1–9 (1999).
25. Conte, P. Effects of ions on water structure: a low-field ‘H T, NMR relaxometry approach. Magn. Reson. Chem. 53, 711–718 (2015).
26. Tielrooij, K., Garcia-Araez, N., Bonn, M. & Bakker, H. J. Cooperativity in ion hydration. Science 328, 1006–1009 (2010).
27. Tielrooij, K. J. et al. Anisotropic water reorientation around ions. J. Phys. Chem. B 115, 12638–12647 (2011).
28. van der Poll, S. T., Tielrooij, K., Hunger, J., Backus, E. H. G. & Bakker, H. J. Femtosecond study of the effects of ions and hydrophobes on the dynamics of water. Faraday Discuss. 160, 171–189 (2013).
29. Turton, D. A., Hunger, J., Hefter, G., Buchner, R. & Wynne, K. Glasslike behavior in aqueous electrolyte solutions. J. Chem. Phys. 128, 161102 (2008).
30. Buchner, R., Capewell, S. G., Hefter, G. & May, P. M. Ion-pair and solvent relaxation processes in aqueous Na₂SO₄ solutions. J. Phys. Chem. B 103, 1185–1192 (1999).
31. Wachter, W., Kunz, W., Buchner, R. & Hefter, G. Is there an anionic Hofmeister effect on water dynamics? Dielectric spectroscopy of aqueous solutions of NaBr, NaI, NaNO₃, NaClO₄, and NaSCN. J. Phys. Chem. A 109, 8675–8683 (2005).
32. Mancinelli, R., Botti, A., Bruni, F., Ricci, M. A. & Soper, A. K. Hydration of sodium, potassium, and chloride ions in solution and the concept of structure maker/breaker. J. Phys. Chem. B 111, 15370–15377 (2007).
33. Rinne, K. F., Gekle, S. & Netz, R. R. Ion-specific solvation water dynamics: single water versus collective water effects. J. Phys. Chem. A 118, 11677–11677 (2014).
34. Rinne, K. F., Gekle, S. & Netz, R. R. Dissecting ion-specific dielectric spectra of sodium-halide solutions into solvation water and ionic contributions. J. Chem. Phys. 141, 214502 (2014).
35. Balos, V. et al. Macroscopic conductivity of aqueous electrolyte solutions scales with ultrafast microscopic ion motions. Nat. Commun. 11, 1611 (2020).

36. Schmidt, D. A. et al. Rattling in the cage: ions as probes of sub-picosecond water. J. Am. Chem. Soc. 131, 18512–18517 (2009).

37. Schwaag, G., Sebastiani, F. & Havenith, M. Ion hydration and ion pairing as probed by THz spectroscopy. Angew. Chem. Int. Ed. 58, 3000–3013 (2019).

38. Heisler, I. A. & Meech, S. R. Low-frequency modes of aqueous alkali halide solutions: glimpsing the hydrogen bonding vibration. Science 327, 857–861 (2010).

39. Heisler, I. A., Mazur, K. & Meech, S. R. Low-frequency modes of aqueous alkali halide solutions: an ultrafast optical Kerr effect study. J. Phys. Chem. B 115, 1863–1873 (2011).

40. Elgabarty, H. et al. Energy transfer within the hydrogen bonding network of water following resonant terahertz excitation. Sci. Adv. 6, aay7674 (2020).

41. Tomaš, R., Jovanović, T. & Bešter-Rogač, M. Viscosity B-coefficient for sodium chloride in aqueous mixtures of 1,4-dioxane at different temperatures. Acta Chim. Slov. 62, 531–537 (2015).

42. Hydrogen, J. Niedermayer, S., Buchner, R. & Hefer, G. Are nanoscale ion aggregates present in aqueous solutions of guanidinium salts? J. Phys. Chem. B 114, 13617–13627 (2010).

43. Sajadi, M., Wolf, M. & Kampfrath, T. Transient birefringence of liquids induced by terahertz electric-field torque on permanent molecular dipoles. Nat. Commun. 8, 14963 (2017).

44. Marcus, Y. The standard partial molar volumes of ions in solution. Part 4. Ionic volumes in water at 0–100 °C. J. Phys. Chem. B 113, 10283–10291 (2009).

45. Wilkins, D. M., Manolopoulos, D. E. & Dang, L. X. Nuclear quantum effects in water exchange around lithium and fluoride ions. J. Phys. Chem. B 114, 9674–9680 (2010).

46. Flores, S. C., Kherb, J., Konelick, N., Chen, X. & Cremer, P. S. The effects of Hofmeister cations at negatively charged hydrophilic surfaces. J. Phys. Chem. C 116, 5730–5734 (2012).

47. Chowdhuri, S. & Chandra, A. Dynamics of halide ion–water hydrogen bonds in aqueous solutions: dependence on ion size and temperature. J. Phys. Chem. B 110, 9674–9680 (2006).

48. Torii, H. Extended dipole-induced dipole mechanism for generating Raman and optical Kerr effect intensities of low-frequency dynamics in liquids. Chem. Phys. Lett. 353, 431–438 (2002).

49. Boyd, R. W. Nonlinear Optics (Academic Press, 2007).

50. Skaf, M. S. & Vechi, S. M. Polarizability anisotropy relaxation in pure and aqueous dimethyl sulfoxide. J. Chem. Phys. 119, 2181–2187 (2003).

51. Ghalgouhi, A., Fingerhut, B. P., Reimann, K., Elsaesser, T. & Woerner, M. Terahertz polaron oscillations of electrons solvated in liquid water. Phys. Rev. Lett. 126, 097401 (2021).

52. Ghalgouhi, A. et al. Field-induced tunneling ionization and terahertz-driven electron dynamics in liquid water. J. Phys. Chem. Lett. 11, 7717–7722 (2020).

53. Zhao, H. et al. Ultrafast hydrogen bond dynamics of liquid water revealed by terahertz-induced transient birefringence. Light Sci. Appl. 9, 136 (2020).

54. Khalilullin, R. Z. & Kühne, T. D. Microscopic properties of liquid water from combined ab initio molecular dynamics and energy decomposition studies. Phys. Chem. Chem. Phys. 15, 15746–15766 (2013).

55. Khalilullin, R. Z., Cobar, E. A., Lochan, R. C., Bell, A. T. & Head-Gordon, M. Unraveling the origin of intermolecular interactions using absolutely localized molecular orbitals. J. Phys. Chem. A 111, 8753–8765 (2007).

56. Watlou, I. et al. The structure of water in the hydration shell of cations from X-ray Raman and small angle X-ray scattering measurements. J. Chem. Phys. 134, 064513 (2011).

57. Yadav, S. & Chandra, A. Structural and dynamical nature of hydration shells of the carbonate ion in water: an ab initio molecular dynamics study. J. Phys. Chem. B 122, 1495–1504 (2018).

58. Urbic, T. Ions increase strength of hydrogen bond in water. Chem. Phys. Lett. 610–611, 159–162 (2014).

59. Vila Verde, A. & Lipowsky, R. Cooperative slowdown of water rotation near densely charged ions is intense but short-ranged. J. Phys. Chem. B 117, 10556–10566 (2013).

60. Laage, D. & Hynes, J. T. Reorientational dynamics of water molecules in anionic hydration shells. Proc. Natl Acad. Sci. USA 104, 11167–11172 (2007).

61. Suresh, S. J., Kapoor, K., Talwar, S. & Rastogi, A. Internal structure of water around cations. J. Mol. Liq. 174, 135–142 (2012).

62. Marcus, Y. The effect of complex anions on the structure of water. J. Solution Chem. 44, 2258–2265 (2015).

63. Marcus, Y. Effect of ions on the structure of water: structure making and breaking. Chem. Rev. 109, 1346–1370 (2009).

64. Smith, J. D., Saykally, R. J. & Geissler, P. L. The effects of dissolved halide anions on hydrogen bonding in liquid water. J. Am. Chem. Soc. 129, 13847–13856 (2007).

65. Stefanovich, E. V., Boldyrev, A. I., Truong, T. N. & Simons, J. Ab initio study of the stabilization of multiply charged anions in water. J. Phys. Chem. B 102, 4205–4208 (1998).

66. Zhang, Y. & Cremer, P. S. Chemistry of Hofmeister anions and osmolytes. Annu. Rev. Phys. Chem. 61, 63–83 (2010).

67. Hofmeister, F. Zur Lehre von der wirkung der salze. Arch. Exp. Pathol. Pharmacol. 24, 395–413 (1888).

68. Okur, H. I., Kherb, J. & Cremer, P. S. Cations bind only weakly to amides in aqueous solutions. J. Am. Chem. Soc. 135, 5062–5067 (2013).

69. Balos, V., Bonn, M. & Hunger, J. Quantifying transient interactions between amide groups and the guanidinium cation. Phys. Chem. Chem. Phys. 17, 28539–28543 (2015).

70. Collins, K. D. Ions from the Hofmeister series and osmolyte: effects on proteins in solution and in the crystallization process. Methods 34, 300–311 (2004).

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.
Methods

Experiment. The THz–Raman experiment is performed in the TKE configuration, whose details are given elsewhere15,94. As shown schematically in Fig. 1a, a linearly polarized THz electric field with a strength of ~2 MV cm\(^{-1}\) pumps the aqueous salt solutions. The liquid samples (thickness of 100 μm) are held between a rear glass window and a 150-mm-thick silicon nitride (SiN) membrane as the entrance window95. The SiN thin window exhibits a negligible Kerr signal and, therefore, eases the challenges for separating the liquid response from that of the window73. The induced transient birefringence, \(\Delta n(t)\), is measured by a temporally delayed and collinearly propagating probe pulse (800 nm, 2 nJ, 8 fs), whose induced linear polarization is set to an angle of 45° relative to the THz electric-field polarization. Due to the pump-induced birefringence, the probe field components polarized parallel (∥) and perpendicular (⊥) to the pump field acquire a phase difference \(\Delta \phi\) when propagating through the sample, thereby resulting in elliptical polarization. The \(\Delta \phi\) is detected with a combination of a quarter-wave plate and a Wollaston prism, from which each split the incoming beam in two perpendicularly polarized beams with power \(P_1\) and \(P_2\). In the limit [\(\Delta \phi << 1\)], the normalized difference \(P_1 - P_2\) fulfills \(P_1 - P_2 \approx \Delta \phi\). The measured phase shift is related to the change in the refractive index \(\Delta n\) of the liquid, that is, \(\Delta \phi = \Delta n \omega c\), where \(c\) is the speed of light.

Samples. All samples were prepared volumetrically by adding Milli-Q water to the weighed amount of salt. The salts were all handled in a glove box to avoid contamination of the sample, \(\omega\), which simplifies the analysis of electrolyte solutions. The liquid samples (thickness of 100 μm) were prepared by dissolving MgCl\(_2\), NaCl and Na\(_2\)CO\(_3\), we reached close to the solubility limit and prepared an additional sample that one needs perturbation strengths that are huge in order to produce a detectable response that is larger than the statistical noise, but this is becoming less important for quantifying the strength of the hydrogen bond—there is no imposed artificial partitioning of the electronic density into separate non-overlapping entities. Charge transfer and polarization effects are naturally accounted for, and finite-size effects are fully included, so that the strength of the electric field varies at each point of space even on submolecular distances. AIMD simulations of aqueous ionic solutions have been found to give reliable results, provided that dispersion effects are carefully treated96.

KE decomposition. In the pFFMD simulations, at each MD snapshot, the translational KE of each molecule is computed from the velocity of the molecular center of mass. The rotational KE is then calculated as the difference between the total molecular KE and the KE of the molecular centre of mass.

Hydrogen-bond ALMO-EDA analysis. For quantifying the strength of the hydrogen bond, we employ ALMO-EDA97 within Kohn–Sham density functional theory. Conceptually, ALMO-EDA decomposes intermolecular interaction energies first by filtering out the frozen electrostatic and polarization effects from the total many-body intermolecular binding energy. The remaining charge transfer contribution is then split into pairwise two-body terms, each corresponding to an individual hydrogen bond or ion–water interaction in the system. These two-body terms are obtained self-consistently under fully periodic boundary conditions. The water–water hydrogen-bond strength is then calculated for all pairs of molecules.

pFFMD. We performed pFFMD simulations on three different aqueous electrolyte solutions—MgCl\(_2\), NaCl and Na\(_2\)SO\(_4\) (1 m, 2 m and 4 m). NaCl (2 m and 4 m) and Na\(_2\)SO\(_4\), (1 m)—plus on pure liquid water (as a reference). Supplementary Table 1 lists the details of the simulated systems. Simulations were performed under fully periodic boundary conditions and a time step of 0.4 fs. For each system, we generated 25,000 non-equilibrium MD trajectories. Initial configurations were uniformly sampled from equilibrated canonical ensemble MD simulations. For each configuration, the thermostat was switched off (microcanonical ensemble) and then a THz electric-field pulse, identical to the one used in our experiment, was applied in the simulation after 0.1 ps. To improve the signal-to-noise ratio, we used a pulse amplitude that is eight times stronger than the experimental one.

Simulations were performed using the Tinker software98, using double precision floating point numbers. The velocity Verlet algorithm was employed to time-propagate the positions and velocities of atoms. The AMOEBA force field parameter set ‘amoebanuc17’ was used99,100. For the sulfate ion, the parameters were taken from ref. 101. A buffered 14-7 potential12 was used to include the effects of a short-range repulsion–dispersion interaction. Short-range interactions were truncated at 9 Å, whereas the smooth particle mesh Ewald12 was employed to treat the long-range electrostatic interactions. The mutual induced dipoles were computed using the conjugate gradients method12,104 with a tolerance of 0.00001 debye. For inclusion of the external electric field \(E\) in energy and force calculations in Tinker, we introduced into the code the electric field term \(\mathcal{F} = k_1 E\) and induced-dipole term \(\mu = k_2 E\), where \(\mu\) is the atomic charge. The constants \(k_1 = 1,185.85\) and \(k_2 = 3,567\) give the forces and dipoles in kilocalories per angstrom per mole and square angstroms, when the electric field \(E\) and polarizability \(\alpha\) are in atomic units and cubic angstroms. The existing implementation of conjugate gradient minimization in Tinker was utilized to compute the mutual induced dipoles that produce the local field at each molecule.

The time-dependent polarizability anisotropy results reported in this work were averaged over all the 25,000 trajectories. For all our analyses involving hydrogen bonds, we have used a standard geometric definition of the hydrogen bond (O–O distance of <3.5 Å and hydrogen bond angle of <30°)105.

AIMD. To study the hydrogen-bond strength between water molecules in aqueous electrolyte solutions, we performed AIMD simulations under field-free conditions for pure liquid water (as reference) and for MgCl\(_2\), NaCl and Na\(_2\)SO\(_4\). Five-atom water system, five independent MD trajectories were simulated. Each trajectory was first equilibrated for 10 ps in the canonical ensemble, followed by a 50 ps production run under the microcanonical ensemble. The AIMD simulations were conducted using the Quickstep module of the CP2K software106. Throughout, the energies and forces were computed using the mixed Gaussian/plane-waves approach107, with the Kohn–Sham orbitals represented by an accurate triple-zeta basis set with sets of polarization functions (TZV2P)106. Plane waves with a cut-off of 400 Ry were used to represent the charge density, whereas the core electrons were described by the Goedecker–Teter–Hutter pseudopotentials108,109. The BLYP exchange–correlation functional was used together with a damped interatomic potential for dispersion interactions ( Grimme-D3)110. Simulations were conducted using a time step of 0.4 fs.

The average hydrogen-bond strengths shown in this work were obtained by performing ALMO-EDA for 500 AIMD snapshots uniformly sampled from the 50 ps production trajectories. The technical details behind ALMO are described elsewhere111. All ALMO-EDA calculations in this work were performed using the ALMO-EDA implementation in the CP2K program112 with the same settings as used in our AIMD simulations.

Calculation of polarizability anisotropy. The polarizability anisotropy was calculated from the ensemble-averaged difference between the xx component of the polarizability tensor, and the average of the remaining two diagonal components, yy and zz (laboratory reference frame, with the z axis being defined by the THz electric field):

\[
\Delta \alpha = \alpha_{xx} - \frac{\alpha_{yy} + \alpha_{zz}}{2}.
\]
The total polarizability of each component can be calculated by summing the permanent ($\alpha_{\text{perm}}$) and induced ($\alpha_{\text{ind}}$) polarizabilities of all the molecules in the system:

$$\alpha_{\text{total}} = \sum_{i=1}^{n} (\alpha_{i}^\text{perm} + \alpha_{i}^\text{ind})$$

where $n$ is the total number of molecules.

The induced polarizability of each molecule was computed using an XDDH mechanism in a self-consistent field manner. The model is extended by the inclusion of both the first and second hyperpolarizabilities. The self-consistent field equation for the first and second hyperpolarizability XDDH mechanisms is given by the following expression:

$$\alpha_{ij}^\text{ind} = \alpha_{ij}^\text{perm} \sum_{j=1}^{n} \left( \beta_{ij} - \gamma_{ij} \right) - \beta_{ij}$$

where $\beta_{ij}$ and $\gamma_{ij}$ are the first and second hyperpolarizability tensors of molecule $i$, respectively; $\gamma_{ij}$ is the local electric-field vector at molecule $i$ created only by all the surrounding molecules (that is, the permanent electric charges and the permanent and induced dipoles of all entities but excluding the external terahertz field) within a 7.5 Å distance from the center of mass of the molecule. We have verified the validity of this cut-off by replicating the simulation box in all directions and increasing the cut-off distance to 15 Å, which gave only a negligible change in anisotropy. The self-consistent field equation was solved iteratively until the solution reached the convergence tolerance of 0.000001 Å$^3$.

The induced dipole moment of each molecule was obtained using the preconditioned conjugate gradient method, with a convergence tolerance of 0.000001 debyes and with the following initial guess, residue, and direction:

Initial guess: $\mu_{i}^\text{initial} = \alpha_{i}^\text{perm}$

$$\mu_{i}^\text{initial} = \sum_{j=1}^{n} \left( \beta_{ij} - \gamma_{ij} \right) - \beta_{ij}$$

Initial residue: $\mu_{i}^\text{initial} = \sum_{j=1}^{n} \left( \beta_{ij} - \gamma_{ij} \right) - \beta_{ij}$

where $\mu_{i}^\text{initial}$ is the external unit vector pointing from molecule $i$ to $j$, $q$ is the molecular charge and $E_{\text{external}}$ is the externally applied field vector.

Parameterization of the dipole moments, polarizabilities and hyperpolarizabilities. The permanent dipole moment of the ions was set to zero, while for water a gas-phase dipole moment of 1.93 debyes was assigned along the molecular bisector. The ionic polarizability parameters from ref. 37 were used to model the permanent polarizabilities of the ions, whereas the polarizability tensor of water was calculated in the gas phase. All the employed parameters are listed in Supplementary Table 2. The values of the dipole moment and polarizability parameters of gas-phase water reported in this work were parameterized using the CP2K program, while the first and second hyperpolarizability parameters of gas-phase water were computed using the DALTON program. For parameterization we used the same settings as described in our AIMD simulations.

Data availability

The raw data underlying all the figures, as well as the full data needed to support, interpret, and extend the research in the article, can be downloaded at https://doi.org/10.5281/zenodo.6514905.

Code availability

The source codes needed to support, interpret, and extend the research in the article can be downloaded at https://doi.org/10.5281/zenodo.6514905. The source code for the calculation of the anisotropy polarizability is available at https://github.com/NaveenKhalan/Dipole-Polarizability.

References

71. Hoffmann, M. C. et al. Terahertz Kerr effect. Appl. Phys. Lett. 95, 131–132 (2009).
72. Balos, V., Bierhance, G., Wolf, M. & Sajadi, M. Terahertz-magnetic-field induced ultrafast Faraday rotation of molecular liquids. Phys. Rev. Lett. 124, 093201 (2020).
73. Sajadi, M., Wolf, M. & Kampfrath, T. Terahertz-field-induced optical birefringence in common window and substrate materials. Opt. Express 23, 28985–28992 (2015).
74. Pluharova, E., Marsalek, O., Schmidt, B. & Jungwirth, P. Ab initio molecular dynamics approach to a quantitative description of ion pairing in water. J. Phys. Chem. Lett. 4, 4177–4181 (2013).
75. Yue, S. & Panagiotopoulos, A. Z. Dynamic properties of aqueous electrolyte solutions from non-polarizable, polarizable, and scaled-charge models. Mol. Phys. 117, 3538–3549 (2019).
76. Ding, Y., Hassanali, A. A. & Parrinello, M. Anomalous water diffusion in salt solutions. Proc. Natl Acad. Sci. USA 111, 3310–3315 (2014).
77. Moučka, F., Nezbeda, I. & Smith, W. R. Molecular force field development for aqueous electrolytes: I. incorporating appropriate experimental data and the inadequacy of simple electrolyte force fields based on Lennard-Jones and point charge interactions with Lorentz–Berthelot rules. J. Chem. Theory Comput. 9, 5076–5085 (2013).
78. Ponder, J. W. et al. Current status of the AMOEBA polarizable force field. J. Phys. Chem. B 114, 2549–2564 (2010).
79. Tommaso, D. D. & De Leeuw, N. H. Structure and dynamics of the hydrated magnesium ion and of the solvated magnesium carbonates: insights from first principles simulations. Phys. Chem. Chem. Phys. 12, 894–901 (2010).
80. Yip, S. (ed.) Handbook of Materials Modeling (Springer, 2005).
81. Hynes, A. T. et al. Tinker 8: software tools for molecular design. J. Chem. Theory Comput. 14, 5273–5289 (2018).
82. Goedecker, S. & Teter, M. Separable dual-space Gaussian pseudopotentials. Phys. Rev. B 54, 145–152 (2005).
83. Lambrecht, D. S., Clark, G. N. I., Head-Gordon, T. & Head-Gordon, M. Exploring the rich energy landscape of sulfate–water clusters $SO_4^{-}$–$H_2O$$_{n}$: an electronic structure approach. J. Phys. Chem. A 115, 11438–11454 (2011).
84. Halgren, T. A. Representation of van der Waals (vdW) interactions in molecular mechanics force fields: potential form, combination rules, and vdW parameters. J. Am. Chem. Soc. 114, 7827–7843 (1992).
85. Essmann, U. et al. A smooth particle mesh Ewald method. J. Chem. Phys. 103, 8577–8593 (1995).
86. Aviat, F. et al. Truncated conjugate gradient: an optimal strategy for the analytical evaluation of the many-body polarization energy and forces in molecular simulations. J. Chem. Theory Comput. 13, 180–190 (2017).
87. Wang, W. & Skeel, R. D. Fast evaluation of polarizable forces. J. Chem. Phys. 123, 164107 (2005).
88. Luzar, A. & Chandler, D. Hydrogen-bond kinetics in liquid water. Nature 379, 55–57 (1996).
89. Hübe, T. D. et al. CP2K: an electronic structure and molecular dynamics software package - Quickstep: efficient and accurate electronic structure calculations. J. Chem. Phys. 152, 194103 (2020).
90. Lippert, G., Hutter, J. & Parrinello, M. A hybrid Gaussian and plane wave density functional scheme. Mol. Phys. 92, 477–488 (1997).
91. VandeVondele, J. & Hutter, J. Gaussian basis sets for accurate calculations on molecular systems in gas and condensed phases. J. Chem. Phys. 127, 114105 (2007).
92. Goedecker, S. & Teter, M. Separable dual-space Gaussian pseudopotentials. Phys. Rev. B 54, 1703–1710 (1996).
93. Krack, M. Pseudopotentials for H to Kr optimized for gradient-corrected exchange-correlation functionals. Theor. Chem. Acc. 114, 145–152 (2005).
94. Grimm, S., Antony, J., Ehrlich, S. & Krieg, H. A consistent and accurate ab initio parameterization of density functional dispersion correction (DFT-D) for the 94 elements H–Pu. J. Chem. Phys. 132, 154104 (2010).
95. Molina, J. J. et al. Ions in solutions: determining their polarizabilities from first-principles. J. Chem. Phys. 134, 014511 (2011).
96. Jungwirth, P., Curtis, J. E. & Tobias, D. J. Polarizability and aqueous solvation of the sulfate dianion. Chem. Phys. Lett. 367, 704–710 (2003).
97. Aidas, K. et al. The Dalton quantum chemistry program system. Wiley Interdiscip. Rev. Comput. Mol. Sci. 4, 269–284 (2014).

Acknowledgements

We thank the Paderborn Center for Parallel Computing for the generous allocation of computing time on the field-programmable-gate-array-based supercomputer ‘Noctua’. This project received funding from the European Research Council under the European Union’s Horizon 2020 research and innovation programme (grant agreement no. 716412) and the Marie Skłodowska-Curie Actions programme MSCA-IF-2020-101030872 (V.B.). H.E. acknowledges funding from the Deutsche Forschungsgemeinschaft (DFG grant EL 815/2).
and T.D.K. The manuscript was written by V.B., M.S. and H.E. with contributions from N.K.K., T.D.K. and M.W. All authors reviewed the results and approved the final version of the manuscript.

**Funding**
Open access funding provided by the Max Planck Society.

**Competing interests**
The authors declare no competing interests.

---

**Additional information**

**Supplementary information** The online version contains supplementary material available at https://doi.org/10.1038/s41557-022-00977-2.

**Correspondence and requests for materials** should be addressed to Vasileios Balos, Hossam Elgabarty or Mohsen Sajadi.

**Peer review information** *Nature Chemistry* thanks Benjamin Fingerhut, Pascale Roy and the other, anonymous, reviewer(s) for their contribution to the peer review of this work.

**Reprints and permissions information** is available at www.nature.com/reprints.