

Ion–water hydrogen-bond switching observed with 2D IR vibrational echo chemical exchange spectroscopy

David E. Moilanen, Daryl Wong, Daniel E. Rosenfeld, Emily E. Fenn, and M. D. Fayer¹

Department of Chemistry, Stanford University, Stanford, CA 94305

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The exchange of water hydroxyl hydrogen bonds between anions and water oxygens is observed directly with ultrafast 2D IR vibrational echo chemical exchange spectroscopy (CES). The OD hydroxyl stretch of dilute HOD in H₂O in concentrated (5.5 M) aqueous solutions of sodium tetrafluoroborate (NaBF₄) displays a spectrum with a broad water-like band (hydroxyl bound to water oxygen) and a resolved, blue shifted band (hydroxyl bound to BF₄⁻). At short time (200 fs), the 2D IR vibrational echo spectrum has 4 peaks, 2 on the diagonal and 2 off-diagonal. The 2 diagonal peaks are the 0–1 transitions of the water-like band and the hydroxyl-anion band. Vibrational echo emissions at the 1–2 transition frequencies give rise to 2 off-diagonal peaks. On a picosecond time scale, additional off-diagonal peaks grow in. These new peaks arise from chemical exchange between water hydroxyls bound to anions and hydroxyls bound to water oxygens. The growth of the chemical exchange peaks yields the time dependence of anion–water hydroxyl hydrogen bond switching under thermal equilibrium conditions as $T_{aw} = 7 \pm 1$ ps. Pump-probe measurements of the orientational relaxation rates and vibrational lifetimes are used in the CES data analysis. The pump-probe measurements are shown to have the correct functional form for a system undergoing exchange.

2D IR spectroscopy | hydration of ions | hydrogen bond dynamics | ion hydrogen bonds chemical exchange | ionic solutions

Water interacting with ions occurs in a wide variety of systems ranging from ocean salt water to water interacting with charged amino acids at the surfaces of proteins (1). The properties of pure liquid water are determined by the nature of its hydrogen bond network. A water molecule can have as many as 4 hydrogen bonds with other water molecules, forming an approximately tetrahedral structure. The pure water hydrogen bond network is constantly evolving with a range of time scales from tens of femtoseconds to picoseconds (2–5). Hydrogen bonds are continually forming and breaking through concerted hydrogen bond rearrangements (6). These dynamical processes can be observed on the time scale they occur in considerable detail by using ultrafast infrared spectroscopy. Measurements of spectral diffusion, described in terms of the frequency–frequency correlation function (FFCF), by using ultrafast 2D IR vibrational echo spectroscopy (3, 7, 8) as well as other ultrafast IR techniques (4, 5) have determined the multiple time scales for the hydrogen bond dynamics. The slowest time component of the FFCF (1.7 ps) is associated with the randomization of the hydrogen bond network through concerted hydrogen bond rearrangements. The orientational relaxation time of pure water (2.6 ps) (2, 5) is also assigned to concerted hydrogen bond rearrangement via jump reorientation (6).

In aqueous salt solutions the structure of water is modified in the vicinity of the ions as the water oxygens preferentially solvate the cations and the water hydroxyls solvate the anions (9, 10). The structures of the hydration shells around individual ions, charged groups of large molecules, or charged amino acids on the surfaces of proteins are determined by ion–dipole interactions

between water molecules and the charged group (11, 12). These interactions will influence both the structure and dynamics of water in the proximity of ions.

In pure water, the hydroxyl stretching IR absorption band is very broad because of the distribution of strengths and numbers of hydrogen bonds (13, 14). When salt is added, hydrogen bonding to the ions causes shifts of the hydroxyl stretch frequency (15–17). The stretching frequency is sensitive primarily to the type of anion and, to a lesser extent, the cation. Addition of NaCl, NaBr, and NaI cause blue shifts of the spectrum that increase as the anion becomes bigger (17). For these salts, the spectra are broad and the shifts are not large. The hydroxyl–anion spectra overlap substantially with the spectrum of hydroxyls bound to water oxygens so that the hydroxyl–anion spectra are not resolved from the hydroxyl–water oxygen spectrum.

Here, we present the results of ultrafast 2D IR vibrational echo chemical exchange experiments on an aqueous solution of the anion BF₄⁻ from the salt NaBF₄. For this anion, the stretch spectrum of hydroxyls bound to BF₄⁻ is substantially shifted to the blue, and the band is relatively narrow (16). The hydroxyl–anion peak is readily observable on the high-frequency side of the broad hydroxyl–oxygen band (see Fig. 1). The observed splitting of the hydroxyl–anion peak from the hydroxyl–water peak makes the experiments discussed here possible.

Ionic hydration and the dynamics of water in salt solutions have been studied extensively by NMR (10, 18), neutron diffraction (19), computer simulations (20–22), and infrared spectroscopy (8, 15, 17, 23). A great deal is known about the structure of ionic solvation shells from diffraction experiments. However, unraveling the dynamics of the solvation shell and the back and forth exchange of waters hydrogen bonded to ions and to water molecules is a much more difficult experimental problem.

A number of studies using ultrafast infrared experiments performed on the water hydroxyl stretching mode have been directed at determining the influence of salts on water dynamics and the rates of hydrogen bond switching between water and anions (8, 24). These experiments (8, 24) and related theory (20–22) have provided insights into water–ion dynamics. However, unambiguous experimental determination of the hydrogen bond switching rate is difficult because the 2 types of hydroxyls are not usually spectroscopically resolved. Below, we report the direct observation of exchange of hydroxyl hydrogen bonds between water and BF₄⁻ anions using ultrafast 2D IR vibrational echo chemical exchange spectroscopy (25).

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¹To whom correspondence should be addressed. E-mail: fayer@stanford.edu.

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axis in Figs. 2 and 3). The vibrational echo signal reads out information about the final structures of all labeled species by their frequencies ω_m .

There are 2 types of time periods in the experiment. The times between pulses 1 and 2 and between pulse 3 and the vibrational echo pulse are called coherence periods. During these periods, the OD vibrations are in coherent superpositions of 2 vibrational states. Fast frequency fluctuations of the vibrational oscillators induced by fast structural fluctuations of the system cause dynamic dephasing, which is one contribution to the line shapes in the conventional absorption spectrum. During the period T_w between pulses 2 and 3, called the population period, an oscillator is in a vibrational eigenstate. Slower structural fluctuations of the system cause the OD frequency to evolve in time. This time evolution of the frequency, termed spectral diffusion, contributes to the 2D line shapes. As discussed in the Introduction, during the time period T_w , chemical exchange causes off-diagonal peaks to grow in.

There are several factors that contribute to the time dependence of the 2D IR spectra. Vibrational population relaxation and orientational relaxation cause all of the peaks to decay. Whereas population relaxation will eventually cause all peaks to decay to zero, orientational relaxation reduces the amplitudes of all of the peaks but not to zero. Spectral diffusion causes all peaks to change in shape but preserves the peak volumes. Chemical exchange causes the diagonal peaks to partially decay. Only the off-diagonal chemical exchange peaks grow in amplitude. By determining the time dependence of the peak volumes, spectral diffusion is eliminated from the determination of the chemical exchange rate. The population relaxation and orientational relaxation are measured by polarization-selective pump-probe experiments and used in the data analysis as discussed below (27).

Analysis of the data following the methods of Kwak *et al.* (27) yields the exchange rate. The number of $hw \rightarrow ha$ exchanges is equal to the number of $ha \rightarrow hw$ exchanges because the system is in equilibrium. To determine the exchange times, it is necessary to know the equilibrium constant, which is the ratio R of hydroxyls bound to anions to hydroxyls bound to water. X-ray diffraction studies of NaBF_4 at almost the same concentration used here (7 waters per NaBF_4) indicate that ≈ 4 water molecules (4 hydroxyls of 14) hydrate a BF_4^- (32).

The equilibrium constant R can also be determined by a comparison of the linear IR spectrum to the 2D IR spectrum. In the IR absorption experiments (see Fig. 1), the areas of the hw band and the ha band were determined. If the transition dipoles, μ_{hw} and μ_{ha} were the same for the 2 species, then the ratio of the areas of the bands would give R . However, the transition dipoles are not the same, and the areas of these bands are proportional to $C_{hw}\mu_{hw}^2$ and $C_{ha}\mu_{ha}^2$, where the C s are the concentrations. The volumes of the hw band and the ha band on the diagonal of the 2D IR spectrum (see Fig. 2) are proportional to $C_{hw}\mu_{hw}^4$ and $C_{ha}\mu_{ha}^4$, respectively. The method for the determination of the volumes is discussed below. Combining the 2 types of measurements yields R and the ratio of the square of the transition dipoles. Within experimental error, we find that $R = 0.3$, the same as determined from the X-ray experiments (32), and $\mu_{hw}^2/\mu_{ha}^2 = 1.7$.

Fig. 1 displays the OD stretch spectrum of HOD in water and with increasing concentrations of NaBF_4 . The value of n in Fig. 1 denotes the number of water molecules per molecule of NaBF_4 . As the concentration of the salt increases, a peak develops on the high-frequency edge of the water spectrum. Its peak position is independent of concentration. This peak is attributed to ODs bonded to the BF_4^- anions (16). In addition, the water peak shifts to the blue with increasing concentration but essentially maintains its shape. At the concentration used in the CES experiments, 5.5 M, the shift is 19 cm^{-1} . At this high concentration, there is no bulk water in the sample. By bulk water, we mean

water that is many solvent shells from an ion. At 5.5 M, which is 7 water molecules per NaBF_4 , all of the water molecules will be associated with a cation, an anion, or both. At the highest concentration, fitting the 2 bands gives the peak positions and widths, which are: $2,526 \text{ cm}^{-1}$ (165 cm^{-1}) and $2,646 \text{ cm}^{-1}$ (45 cm^{-1}), where the number in parentheses is the full width at half maximum.

Fig. 2 displays contour plots of 2D IR vibrational echo data at $T_w = 200 \text{ fs}$, which is short compared with the chemical exchange time. For clarity, the contours occur at intervals of 10% of the maximum. The data used in the analysis has much finer gradations. There are 4 bands, 2 going in the positive direction (labeled +) and 2 going in the negative direction (labeled -). The narrow positive peak to the upper right labeled ha_{01} arises from the ha 0-1 transition. Almost directly below it is the peak going in the negative direction produced by vibrational echo emission at the 1-2 transition of ha , labeled ha_{12} . This peak is slightly displaced to the right because of a small amount of overlap with the broad hw 0-1 transition going in the positive direction. The difference in the center frequencies of the ha 0-1 and 1-2 bands along the ω_m axis (vertical axis) is the anharmonic shift of the OD hydroxyl stretch for hydroxyls bound to anions. The shift is 90 cm^{-1} . Below the broad 0-1 hw band going in the positive direction (labeled hw_{01}) is the corresponding broad band going in the negative direction for the 1-2 transition of the OD hw stretch labeled hw_{12} . The anharmonic shift is 140 cm^{-1} .

Chemical exchange will cause 4 additional off-diagonal peaks to grow in, 2 of which are going in the positive direction, arising from the 0-1 diagonal transitions, and 2 that are going in the negative direction and come from the 1-2 transitions. Fig. 3a displays the 2D spectrum at $T_w = 4 \text{ ps}$ in which the growth of the chemical exchange peaks is clear. Only the higher-frequency region of the 2D spectrum is presented, making it easier to see the nature of the data. This portion of the spectrum is the part used in the data analysis discussed below. The IR pulses used in the experiments have a center frequency of $2,550 \text{ cm}^{-1}$ to efficiently pump the 0-1 transition of hw and ha . The laser intensity is relatively weak in the hw 1-2 region, reducing the data quality at long times. In Fig. 3a, the exchange peaks are clearly visible. The most evident chemical exchange peak is the 0-1 $hw \rightarrow ha$ peak, labeled A in Fig. 3a. The spectrum shown in Fig. 3a contains 2 other exchange peaks. The 1-2 $hw \rightarrow ha$ peak is labeled B. Because the 1-2 $hw \rightarrow ha$ peak is negative, relatively narrow along the ω_m axis and extended along the ω_r axis, it eats away a strip from the diagonal 0-1 hw band. This is seen very clearly by comparing the hw peak in Fig. 3a to its short-time counterpart in Fig. 2. The last exchange peak in Fig. 3a is the 0-1 $ha \rightarrow hw$ peak, labeled C. Like peak A, it is going in the positive direction and is manifested as a reduction in the bottom portion of the 1-2 ha peak going in the negative direction. The shapes of the diagonal and off-diagonal CES peaks have been explicated theoretically and experimentally (27, 38).

As discussed above, spectral diffusion causes the shapes of the bands to change with increasing T_w , but spectral diffusion does not change the peak volumes (27). Therefore, to determine the $hw \leftrightarrow ha$ chemical exchange rate, the peak volumes of the 6 peaks that occur in data like that presented in Fig. 3a are determined for a series of T_w s. A detailed theoretical study in which CES data were analyzed by using a full diagrammatic perturbation theory of the spectral diffusion (27) in addition to chemical exchange gave the same results as the method of fitting the peak volumes (25). The peaks are fit simultaneously with 6 2D Gaussian functions. The positions, widths, volumes, and tilts of the Gaussians are varied to minimize the square of the residuals. Although there are many parameters, there are also many constraints. For example, the diagonal 0-1 ha peak and the off-diagonal 1-2 ha peak should be identical in volume except for sign (positive vs. negative) and the shift along the ω_m axis, and

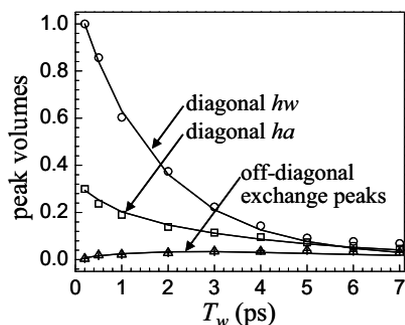


Fig. 4. Peak volumes as a function of T_w of the diagonal and chemical exchange peaks from data like those shown in Fig. 3. The solid curves are the result of to the exchange kinetic model.

the off-diagonal chemical exchange $0-1\ hw \rightarrow ha$ peak should be identical to the off-diagonal chemical exchange $0-1\ ha \rightarrow hw$ peak. In practice, experimental error results in such constraints not being met perfectly, but any serious deviation is an indication that the fit has settled into a false minimum. Fig. 3*b* is the result of fitting the data in Fig. 3*a*. It is clear that the fitting model reproduces the data with substantial accuracy.

Once the peak volumes are obtained for the series of 2D spectra over a range of T_w , it is necessary to relate the volumes to populations. The populations associated with the diagonal $0-1\ hw$ and ha peaks are proportional to their volumes times μ_{hw}^4 and μ_{ha}^4 , respectively. Assuming the harmonic approximation, the off-diagonal $1-2\ ha$ peak also scales by μ_{ha}^4 . The 3 off-diagonal chemical exchange peaks in Fig. 3 all scale as $\mu_{hw}^2\mu_{ha}^2$. The peak volumes are scaled by these factors by using the factor $\mu_{hw}^2/\mu_{ha}^2 = 1.7$, which was determined as described above. The time dependence of the $ha\ 0-1$ and $1-2$ peaks should be the same (25, 27). The time dependence of all 3 chemical exchange off-diagonal peaks should be the same (25, 27). Therefore, all of the dynamics are reflected in the subset of peaks, the diagonal $0-1\ ha$ peak, the diagonal $0-1\ hw$ peak, and the off-diagonal $0-1$ chemical exchange $hw \rightarrow ha$ peak. These are plotted in Fig. 4. The data are normalized to the shortest time point of the hw peak.

To obtain the chemical exchange rate, it is necessary to fit the data in Fig. 4 with a model that includes the vibrational population decay rates of hw and ha , and the orientational relaxation rates of hw and ha . In analyzing the data, the procedures applied previously are used (25, 27). However, there are 2 complications. In earlier experiments on, for example, the formation and dissociation of phenol-benzene π hydrogen bonded solute-solvent complexes in the mixed solvent of benzene and carbon tetrachloride (25), it was possible to make independent determinations of the population and orientational decay rates. In the current experiments, it is not possible to make the equivalent measurements, so the exchange data and measured vibrational lifetime decays are fit simultaneously by using the same kinetic model. The advantage of this fitting method is that it provides an internally consistent confirmation that the 2 very different types of measurements provide the same information about the underlying molecular dynamics of the system.

Polarization selective pump-probe experiments were used to determine the apparent lifetime and orientational relaxation rate for ha at the frequency of the $0-1$ transition of ha , $2,646\ \text{cm}^{-1}$. The apparent lifetime and orientational relaxation rate for hw were determined at the frequency of the $1-2$ transition of hw , $2,365\ \text{cm}^{-1}$, rather than at the $0-1$ transition, because the $1-2$ transition of ha overlaps with the $0-1$ transition of hw . By using the equilibrium constant R and the measured values as an initial guess, the data in Figs. 4 and 5 were simultaneously fit to a system

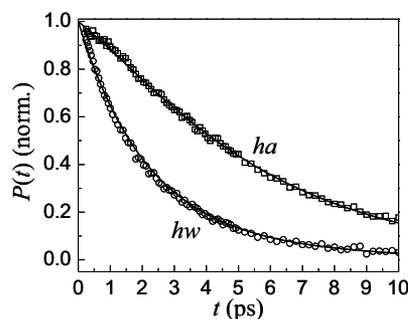


Fig. 5. Vibrational population decays for ha and hw . The solid curves are the result of simultaneously fitting the population decays and the peak volumes shown in Fig. 4 to the exchange kinetic model. The model accurately reproduces the highly nonexponential shape of the ha data.

of coupled differential equations (27). In the fitting, the only adjustable parameters are the 2 vibrational lifetimes and the exchange rate for the process, $ha \rightarrow hw$. To reduce the number of adjustable parameters, the orientational times were not varied in the fitting because, as discussed below, the time scale for reorientation is almost the same for the 2 species, so exchange has very little effect on the measured values. It was found that if the exchange data in Fig. 4 were fit with the $ha \rightarrow hw$ exchange rate as a single adjustable parameter by using an initial guess for the lifetimes as fixed input parameters, the resulting exchange rate was fairly close to the value found from the simultaneous fits, but the quality of the fits was not nearly as good. The simultaneous fitting is necessary to obtain the correct vibrational lifetimes and correct functional forms of the population decays as well as to obtain high quality fits through all 3 datasets in Fig. 4. The simultaneous fits give an exchange rate, vibrational lifetimes, and orientational relaxation times that are internally consistent with the data measured by 2D IR CES and polarization selective pump-probe spectroscopy.

The fits to the exchange model are shown as the solid curves in Figs. 4 and 5. The quality of the fits in both Figures is excellent. In Fig. 4, the fit reproduces the time dependence of the diagonal and chemical exchange peaks. The exchange time for $ha \rightarrow hw$ is $T_{aw} = 7 \pm 1$ ps. The exchange time for $hw \rightarrow ha$, T_{wa} , is related to T_{aw} by the equilibrium constant. Because the system is in equilibrium, $R = k_{wa}/k_{aw}$, where $k_{aw} = 1/T_{aw}$ and $k_{wa} = 1/T_{wa}$ are the rate constants for exchange and R is the equilibrium constant. Using $R = 0.3$ gives $T_{wa} = 24$ ps. This value will be dependent on the concentration of salt because $hw \rightarrow ha$ can only occur if a water molecule is very close to an anion. Therefore, the exchange dynamics are better characterized by the time constant, $T_{aw} = 7$ ps, which may be relatively insensitive to salt concentration as discussed below.

Fig. 5 shows the vibrational population relaxation for hw and ha with the results of the exchange model fit as solid curves. The process of exchange modifies the vibrational population decays such that the experimentally measured decays do not provide the true lifetimes. Initially, the data are fit with single exponential decays to extract a first guess for the lifetimes. The hw decay is substantially faster than the ha decay. Because the system is in equilibrium, at $t = 0$, equal population is exchanged between the 2 species. However, as time proceeds, the hw population decays more than the ha population, and the exchange process will serve to bolster the population of hw , making the apparent lifetime longer than the true lifetime. The converse is true for ha . As discussed below, the shapes of the population decay curves, particularly the ha curve, are substantially influenced by the chemical exchange. Note that the number of molecules undergoing exchange is always constant, but the fraction of excited molecules of each species depends on the vibrational lifetime of that species.

The hw decay is close to exponential because the intrinsic lifetime is much shorter than the exchange time. A single exponential fit to the hw population decay gives the apparent time constant, $\tilde{\tau}_{hw} = 2.5$ ps. The tilde indicates an apparent lifetime. However, the ha decay is far from exponential, and it is remarkable how well the exchange model reproduces its nonexponential shape. At early times, before significant exchange has occurred, the ha population decay is fairly close to the true lifetime. A single exponential fit to the ha lifetime decay from 0.2 ps to 2 ps gives a time constant of $\tilde{\tau}_{ha} = 8$ ps. In Fig. 5, it can be seen that the ha decay during the first picosecond is slower and then becomes steeper as the exchange process shortens the apparent vibrational lifetime. A fit beginning at 3 ps (after exchange has begun to play a role) gives an apparent time constant of $\tilde{\tau}_{ha} = 4.7$ ps. Simultaneously fitting the 2D IR CES data and the pump probe data yields the true values for the lifetimes of $\tau_{hw} = 2.2 \pm 0.1$ ps and $\tau_{ha} = 9.4 \pm 1$ ps. The true values are consistent with the exchange process increasing the apparent lifetime of the fast component and decreasing the apparent lifetime of the slow component.

Measurements of the anisotropy following procedures described previously (2) yields values of $\tau_r^{ha} = 5.0$ ps and $\tau_r^{hw} = 4.1$ ps where τ_r^{ha} and τ_r^{hw} are the long time orientational correlation times for ha and hw , respectively. The orientational dynamics of ha are biexponential with a fast component of 0.7 ps due to fast orientational motion within an intact hydrogen bonding configuration (39). The biexponential nature of the orientational dynamics of ha is well described by a wobbling-in-a-cone model (39–41), and its effects on the exchange process were included in the fitting described above. Because of space limitations, the orientational relaxation data are not shown here. Because the values for the long time components of the orientational relaxation are quite similar, they are changed negligibly by exchange.

The time constants for orientational motion are faster than the time constant for chemical exchange, $T_{aw} = 7 \pm 1$ ps. The orientational relaxation times and the chemical exchange times are not directly comparable because the orientational decay time is the decay of the second Legendre polynomial correlation function. Nonetheless, it is reasonable to expect that orientational relaxation will be faster than chemical exchange. In pure water, orientational relaxation is modeled as jump reorientation. Here, the jumps can be from ha to hw and hw to ha , as well as ha to ha , and hw to hw . The first 2 types of jumps produce both chemical exchange and orientational relaxation, whereas the last 2 produce only orientational relaxation. In addition, it has been proposed that the bonding of a water hydroxyl to BF_4^- is not spatially specific to the fluorines (42). Hydrogen bond rearrangements affecting the other hydroxyl might cause the ha to move on the surface of the anion, producing orientational relaxation. Thus, there are more pathways contributing to orientational relaxation than to chemical exchange, in accord with the observation of faster reorientation. Interestingly, the orientational relaxation times are less than a factor of 2 slower than the orientational relaxation time of pure water, $\tau_r = 2.6$ ps. Thus, the dynamics of water in this concentrated salt solution are not tremendously slower than in pure water.

Previously, 2D IR vibrational echo experiments and pump-probe experiments were used to study water dynamics in NaBr solutions at several concentrations (8, 41). It was found that the concentration dependence of the FFCF and the orientational relaxation were identical. These results and their agreement with MD simulations (43) were interpreted as reflecting the exchange time. However, in this earlier experimental study, the exchange was not measured directly, and the additional pathways that could lead to reorientation or spectral diffusion without exchange were not considered. In accord with the results reported here, the orientational relaxation times were slower than that of pure water but only by approximately a

factor of 2. Based on the results obtained here, it is likely that the NaBr $ha \leftrightarrow hw$ exchange is somewhat slower than suggested previously (8, 41).

In the concentrated solution studied here, the solvation shells of the cations and anions are strongly overlapped, and it is quite possible that there is no second solvation shell of water molecules. It is important to note that a chemical exchange event does not require that a water molecule leaves the first solvation shell of the anion. All that is necessary for an exchange event to be observed is that a hydrogen bond rearrangement occurs so that $ha \rightarrow hw$ or $hw \rightarrow ha$. On a molecular level, this may occur if an HOD initially donating an OD hydrogen bond to BF_4^- undergoes an exchange event and ends up donating its OH hydrogen bond to the same BF_4^- . Although this mechanism has been observed in simulations of bulk water (44), it is much more common for the donating water molecule to move out of the first solvation shell of the initial acceptor (21, 22). The data reported here demonstrate that the time for hydrogen bond switching in this concentrated solution is slower than in bulk water, but only by a relatively small factor. If solvation shell exchange is governed by hydrogen bond switching, it is reasonable to expect that the exchange time reported here for $ha \rightarrow hw$, $T_{aw} = 7 \pm 1$ ps, is representative of the time scale for a water molecule to leave the solvation shell of an anion in more dilute solutions.

Two-color pump probe IR studies have suggested that the dynamics of water in the solvation shell of ions are substantially slower than in pure water (24). In contrast, NMR studies of water solvating ions (10, 18) and proteins (1) suggested that the dynamics are not drastically different from those of pure water. Recent MD simulations (21, 22) and ultrafast 2D IR experiments (8, 41) also suggest that water dynamical processes do not slow drastically in the proximity of ions. The current 2D IR CES experiments provide a direct measurement of the $hw \leftrightarrow ha$ chemical exchange. Combined with the pump-probe experiments and the application of the exchange model to the dynamics measured with both CES and pump-probe, significant details of water dynamics in the proximity of ions have been explicated.

Concluding Remarks

Chemical exchange in the solvation shells of BF_4^- in concentrated aqueous solutions of NaBF_4 was measured with a combination of 2D IR CES and polarization selective pump-probe spectroscopy. The exchange events consist of hydroxyls switching hydrogen bonding partners between BF_4^- anions (ha) and water oxygens (hw). A kinetic model, which includes the effects of the different vibrational lifetimes and orientational relaxation of the 2 species, was fit to the exchange data and pump probe data. The agreement between the model and the data (Figs. 4 and 5) is excellent and the time constant for chemical exchange, $ha \rightarrow hw$ is $T_{aw} = 7 \pm 1$ ps. The experiments show that the time scale for a hydroxyl to switch from being hydrogen bonded to an anion to being bonded to a water oxygen is only several times longer than the time scales for hydrogen bond rearrangement dynamics in pure water. These results have broad implications, suggesting a relatively short residence time for water near many charged solutes including the charged amino acids at the surfaces of proteins.

Materials and Methods

The CES experiments are performed on the OD hydroxyl stretch of dilute (5%) HOD in $\text{H}_2\text{O}/\text{NaBF}_4$ at 298 K. The concentration of NaBF_4 used in the CES experiments is 5.5 M. A Ti:Sapphire oscillator/regenerative amplifier pump an OPA and difference frequency stage to produce ≈ 70 -fs IR pulses.

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