Accelerated Vibrational Energy Relaxation of Water in Alkaline Environments

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latter vibrations. Here, we investigate whether a similar effect can occur in alkaline solutions. Recent theoretical studies have shown that hydroxide ions possess a different hydration structure compared to protons. We use ultrafast mid-IR pump–probe spectroscopy to investigate the vibrational relaxation dynamics of the OD stretch excitations of HDO molecules dissolved in aqueous solutions of hydroxide ions of different concentrations.

■ EXPERIMENTAL METHODS

Aqueous hydroxide solutions were prepared by diluting a commercially available solution of 50 wt % NaOH (Sigma Aldrich) in deionized water (5.5 μS/m, Milli-Q) at concentrations ranging between 1 and 6 M. In addition, D₂O (99.96% D, Eurisotop) was added to obtain an OD/OH ratio of 4% in all studied samples. The samples were kept between two 2 mm thick CaF₂ windows separated by Teflon spacers of 25 μm for 1–3 M NaOH solutions and 10 μm at higher concentrations. The path length did not affect the relaxation dynamics.

We perform polarization-resolved transient absorption measurements using femtosecond mid-infrared light pulses with a repetition rate of 1 kHz and centered at a frequency of 2510 cm⁻¹ that are resonant with the OD stretch vibration of HDO molecules in aqueous solution. These pulses are generated via parametric down-conversion of Yb:KGW laser pulses in a KTA crystal. At 2510 cm⁻¹, the generated pulses have an energy of 12 μJ, a bandwidth of 120 cm⁻¹, and a pulse duration of 200 fs. At 3390 cm⁻¹, the pulse energy is 25 μJ, the bandwidth is 90 cm⁻¹, and the pulse duration is 280 fs. A ZnSe window at an incident angle of 45° is used to create the pump (transmission), while the reflected beam is sent over a second ZnSe window to generate the probe and reference pulses. The pump pulse is sent over a delay stage to vary the pump–probe time delay and sent through a λ/2 plate to set its polarization at π/4 with respect to the polarization of the probe. A rotating wire grid polarizer is used after the sample to selectively block the probe.
measure the polarization component of the probe that is parallel or perpendicular to the pump polarization.

We measure the pump-induced transient absorption changes ($\Delta \alpha$) as a function of time delay between the pump and the probe. At $t = 0$, the pump pulse excites OD stretches from the ground state to the first excited state, $|0\rangle \rightarrow |1\rangle$. This excitation leads to a reduced density of absorbers in the ground state and stimulated emission out of the excited state, both contributing to an absorption decrease (bleach). The excitation also leads to absorption at lower frequencies, associated with the $|1\rangle \rightarrow |2\rangle$ transition. The vibrational relaxation of the excited state eventually results in heating of the sample, which induces an absorption change that decays on a time scale (tens to hundreds of picoseconds) much slower than that of the pump–probe experiment.

The experimentally obtained parallel and perpendicular transient absorption signals are used to construct the isotropic transient response

$$\Delta \alpha_{\text{iso}}(\omega, t) = \frac{1}{3} \left[ \Delta \alpha_{\parallel}(\omega, t) + 2 \Delta \alpha_{\perp}(\omega, t) \right]$$

where $\omega$ and $t$ are the probe frequency and time delay, respectively. The isotropic signal provides information on the rate of vibrational energy relaxation and is insensitive to molecular reorientation.

**RESULTS**

**Linear IR Absorption.** Figure 1 displays the linear absorption spectra of solutions of 8% HDO and different concentrations of NaOH in H$_2$O. The spectra show a band centered at 2510 cm$^{-1}$ that is assigned to the OD stretch vibration of HDO molecules that are hydrogen-bonded to neighboring water molecules. With increasing OH$^-$ concentration, a broad background absorption rises, which is attributed to the OH stretch vibrations of H$_2$O molecules donating strong hydrogen bonds to OH$^-$. The spectral responses of the OD/OH stretch vibrations are negligibly affected by Na$^+$ ions.

**Isotropic Transient Spectra.** Figure 2 shows isotropic transient absorption spectra for neat HDO/H$_2$O and HDO dissolved in 2 M (middle panel) and 5 M (lower panel) NaOH/H$_2$O solutions. All solutions contain 8% HDO/D$_2$O. The isotropic transient spectra show a pump-induced bleaching signal at around 2510 cm$^{-1}$ due to the excitation of the OD stretch vibration of the HDO molecules. At frequencies <2420 cm$^{-1}$, the upper panel also shows the high-frequency wing of the induced $|1\rangle \rightarrow |2\rangle$ absorption. The lower panels show that the amplitude of the bleaching signal of the OD stretch vibration decreases upon the addition of NaOH because an increasing part of the energy of the excitation pulse is absorbed by OH vibrations of H$_2$O molecules that are hydrogen-bonded to OH$^-$. In addition, with increasing OH$^-$ concentration, the OD groups of an increasing fraction of the HDO molecules will be directly hydrogen-bonded to OH$^-$, which shifts their absorption to lower frequency. As a result, these OD groups no longer contribute to $\sigma_{\text{OD}}$, thus decreasing the amplitude of this spectral component in comparison to $\sigma_{\text{OH}}$ and $\sigma_{\text{therm}}$.

To elucidate the transient spectral response of the excited OH stretch vibrations of H$_2$O solvating OH$^-$, we measure the transient absorption spectra of NaOH/H$_2$O solutions in a concentration range of 1–5 M without the addition of D$_2$O (see the Supporting Information). In these experiments, we only probe the vibrational relaxation dynamics of the ion-associated OH groups. The signal can be well described using a two-step relaxation model, in which the excited OH vibration relaxes rapidly within the solvation structure of the OH$^-$ ion, leading to a locally heated hydration complex with an associated intermediate-state spectrum $\sigma_I$. This locally hot state equilibrates with the surroundings due to heat diffusion,
thus creating a global hot ground state $\sigma_{\text{hgs}}$. This relaxation mechanism has been observed in a previous study on heat diffusion and molecular reorientation in aqueous hydroxide solutions.17

From a least-square fit of the isotropic data, we find that the two subsequent relaxation processes take place with time constants of $0.15 \pm 0.1$ and $0.5 \pm 0.1$ ps, both independent of the concentration. The uncertainty in a time constant of $0.15$ ps is relatively large as a result of the relatively long pulse duration of $0.2$ ps. The spectral signature $\sigma_{\text{OH}}$ of the excited state does not depend on concentration. The spectral shape of the intermediate state $\sigma_1$ is very similar to that of the hot ground state $\sigma_{\text{hgs}}$. The main difference is that $\sigma_1$ has a much larger amplitude.

**Spectral Decomposition and Population Dynamics.**

Based on these observations, we consider the transient absorption signal to be composed of the responses of two excited states and a thermal spectrum $\sigma_{\text{therm}}$. This latter spectrum $\sigma_{\text{therm}}$ represents both the spectral response $\sigma_1$ of the intermediate state of the relaxation of the OH vibration of H$_2$O molecules solvating OH$^-$ and the spectral response $\sigma_{\text{hgs}}$ of the final hot ground state. We describe the responses of the intermediate state and the hot ground state with a single spectral component $\sigma_{\text{therm}}$ because these states have very similar spectral shapes. The time evolution of $\sigma_{\text{therm}}$ thus represents the population dynamics of both the intermediate state and the hot ground states. The transient absorption signal can thus be described as a linear combination of three states with associated transient spectra $\sigma_j$ and time-dependent populations $N_j$

$$\Delta \alpha_{\text{tot}} = \sum_{j=1}^{3} N_j(t) \cdot \sigma_j(\omega)$$  \hspace{1cm} (2)

To determine the relaxation dynamics of the OD stretch vibration, we perform a spectral decomposition of the transient absorption spectra at all delay times and all concentrations. In this analysis, we fix the shape of the independently measured concentration-independent $\sigma_{\text{OH}}$ band, and we fit the shape of the thermal spectrum $\sigma_{\text{therm}}$ at each studied OH$^-$ concentration to the transient absorption spectrum measured at long delay times ($>30$ ps). We treat the spectral shape of the excited OD vibration $\sigma_{\text{OD}}$ as a global fit parameter across all OH$^-$ concentrations. The global fit yields the spectral shape of $\sigma_{\text{OD}}$ and the time-dependent population dynamics $N_j(t)$ of all three states without any assumption about the relaxation mechanism. The fitting procedure is described in detail in the Supporting Information. For a 2 M NaOH/H$_2$O solution, the $\chi^2$-minimization process yields the spectral components shown in Figure 3 and the population dynamics displayed in Figure 4. The solid red line in Figure 4 shows the dynamics of the thermal state, which represents the decay of the intermediate state of the relaxation of the OH vibration of H$_2$O molecules solvating OH$^-$ at short delay times, and the ingrowing signal of the hot ground state at late delay times ($>2$ ps). The solid curves displayed in Figure 2 show that the spectral components shown in Figure 3 fit our experimental data well, even at the high OH$^-$ concentration of 5 M.

In Figure 5, we show the normalized population dynamics of the excited OD vibration for five different NaOH concentrations. We observe that the relaxation becomes faster and increasingly nonexponential with increasing ion concentration. This ion-induced change of the relaxation can be explained by

**Figures:**

- Figure 3. Spectral decomposition of the transient absorption spectrum of a solution of HDO and 2 M NaOH in H$_2$O. The spectra $\sigma_{\text{OD}}$ and $\sigma_{\text{therm}}$ have been fixed in a least-squares optimization process of eq 2 to the data, while the spectral shape $\sigma_{\text{OD}}$ of the OD vibrations is fitted as a global parameter across all OH$^-$ concentrations.

- Figure 4. Fitted population dynamics of the different spectral components of Figure 3 of a solution of HDO and 2 M NaOH in H$_2$O. The population dynamics of the thermal spectrum represents both the decay of the intermediate state of the relaxation of the OH vibration of H$_2$O molecules solvating OH$^-$ and the ingrowing signal of the hot ground state.

- Figure 5. Time-evolving population probability of excited OD stretch vibrations at five different NaOH concentrations. The markers show the temporal decomposition described in the main text. The solid lines show the least-square fit of eq 5, which results in 1.78 ps for the OD vibrational lifetime and 2.98 Å for the Förster radius. The dashed lines represent two single-exponential decays with time constants of 1.75 and 1.2 ps. The difference between the latter curve and the data illustrates that the observed decay is nonexponential.
three possible scenarios: (i) the addition of OH\(^-\) to the solution leads to inhomogeneous broadening of the OD absorption band with a faster relaxation rate at low OD frequencies than at high OD frequencies, thus yielding a nonexponential decay for \(\sigma\) \(OD\); (ii) hydroxide ions affect the binding dynamics of the hydrogen-bond network, which in turn changes overall the vibrational relaxation process; (iii) the spectral overlap between the OD band and the hydroxide continuum enables resonant Förster energy transfer from excited OD vibrations to unexcited hydroxide complexes. To investigate scenario (i), we investigated the frequency dependence of the relaxation of the OD stretch vibration (see Figure S4 in the Supporting Information). We observe this relaxation to be uniform over the OD absorption band, thus ruling out the possibility that the nonexponential relaxation would originate from a frequency dependence of the OD relaxation rate.

To discriminate between scenarios (ii) and (iii) as origins of the ion-induced increase in the relaxation rate, we performed the same experiment on \(H, D\)-reversed isotope solutions, i.e., \(H_2O\) diluted in \(D_2O\). We thus measure the transient spectral response of solutions of HDO and OD\(^-\) in \(D_2O\), following the excitation of the OH stretch vibration of the HDO molecules near 3400 cm\(^{-1}\). Since the OH stretch vibration band does not overlap with the deuterioxide-induced continuum, as the inset in Figure 6 shows, the possibility of Förster energy transfer can be ruled out for these samples.

We find that for all studied OD\(^-\) concentrations, the vibrationally excited OH groups that are directly hydrogen-bonded to OD\(^-\) ions relax with a time constant \(T_r\) of 0.27 ± 0.05 ps and that the intermediate state relaxes with a time constant 0.9 ± 0.3 ps. The OH stretch vibrations of the HDO molecules that are not directly hydrogen-bonded to OD\(^-\) ions relax with a time constant that shows negligible dependence on the OD\(^-\) concentration, as illustrated in Figure 6. At all concentrations, the relaxation can be well described with an exponential function with a decay time constant of 0.75 ps. This relaxation time is similar to the previously reported time constant of 0.7 ps for the excited OH stretch vibration of HDO in neat \(HDO/D_2O\).19

In a previous study of the reorientation dynamics of the OD and OH vibrations of HDO molecules in aqueous hydroxide solutions, we did not explicitly include the vibrational relaxation of the OH groups that form hydrogen bonds with OD\(^-\), and we only considered the thermal effect following the relaxation of these vibrations.20 As a result, the observed relaxation rate represented the average relaxation dynamics of all OH vibrations, and this averaged relaxation becomes faster with increasing OD\(^-\) concentration, as the contribution of OH groups directly bonded to OD\(^-\) increases. In the present study, we investigate the OH stretch relaxation over a larger spectral range and we include the vibrational relaxation of the OH groups that form hydrogen bonds with OD\(^-\). We thus observe that for both the OH groups hydrating OD\(^-\) and the OH groups that are not hydrating OD\(^-\), the vibrational relaxation rate shows negligible dependence on the OD\(^-\) concentration. This means that there is no acceleration of the relaxation of the OH of HDO in OD\(^-\)/\(D_2O\) solutions with increasing OD\(^-\) concentration, only the relative contributions of the two OH species change.

It is interesting that for the OH vibration of HDO in OD\(^-\)/\(D_2O\), we observe a fast relaxation component in the red wing of the OH absorption band, associated with OH groups that directly hydrate OD\(^-\) ions, whereas for the OD vibration of HDO in OH\(^-\)/\(H_2O\) solutions, we do not see a similar component associated with OD groups directly hydrating OH\(^-\) (see Figure S4 in the Supporting Information). This observation suggests that these latter OD groups have a similar vibrational relaxation time constant as the OD groups that are further away from the OH\(^-\) ions. A possible explanation for this difference between OD and OH vibrations is that the relaxation mechanisms differ, with the result that the relaxation of the OH vibration could be much more sensitive to the local hydrogen-bond strength than the relaxation of the OD vibration.

The absence of the concentration dependence of the relaxation of the OH vibrations of HDO molecules further away from OH\(^-\) shows that the change of the hydrogen-bond network resulting from the addition of hydroxide ions has a negligible effect on the vibrational relaxation. Hence, the observed acceleration of the vibrational relaxation for the OD vibrations of HDO in OH\(^-\)/\(H_2O\) solutions can only be explained with the third scenario, i.e., Förster energy transfer from the excited OD vibration to OH vibrations of \(H_2O\) molecules hydrating the OH\(^-\) ions. This energy transfer is enabled by the spectral overlap of the OD stretch absorption spectrum and the wing of the broad and strong absorption band of the OH stretch vibrations of \(H_2O\) molecules hydrating OH\(^-\) ions.

![Figure 6. Time-evolving population probability of excited OH stretch vibrations at five different NaOD concentrations. The markers show the temporal decomposition described in the main text. The solid line represents a monoeponential decay with a time constant of 0.75 ps, similar to that in the neat HDO/D_2O solvent.](https://doi.org/10.1021/acs.jpcb.1c02730)
The observed uniform relaxation of the OD vibration at different frequencies (see Figure S4 in the Supporting Information) is also consistent with a mechanism in which the nonexponential character of this relaxation is caused by Förster energy transfer, as the coupling with the very broad absorption of hydrated OH\(^-\) complexes will not strongly depend on the OD frequency. In principle, the reverse Förster process from hydroxide complexes to OD vibrations can also occur. However, this interaction will have a negligible effect since vibrationally excited OH groups of H\(_2\)O molecules hydrating OH\(^-\) ions quickly relax with a time constant <200 fs.

**Förster Energy Transfer.** The comparison between Figures 5 and 6 shows that Förster energy transfer to the hydrated OH\(^-\) complexes serves as an effective additional relaxation channel for the excited OD vibrations. In the following, we will model the effect of this additional channel using the Förster formalism.

Within the Förster model, the rate of energy transfer between two dipoles is given by

\[
k_{\text{FI}}(r) = \frac{1}{T_{1}} \left( \frac{R_{0}}{r} \right)^{6}
\]

where \(T_{1}\) is the intrinsic lifetime of the excited state and \(r\) is the distance between the two interacting oscillators. The Förster radius \(R_{0}\) determines the distance at which 50% of the excitation energy is transferred via the Förster mechanism within the lifetime \(T_{1}\) of the excitation. It thus follows that the Förster process becomes significant if the mean donor–acceptor distance is comparable or smaller than \(R_{0}\), as will be the case at high OH\(^-\) concentrations. Using eq 3 and assuming a radially uniform distribution of receiving oscillators, the survival probability of donors is given by

\[
S(t, \rho) = \exp\left(-4\pi\rho \int_{a_{0}}^{\infty} \text{d}r r^{2} [1 - e^{-k_{\text{FI}}(r) t}] \right)
\]

where \(\rho\) is the density number of OH\(^-\) ions and \(a_{0}\) is the minimal approach distance, which is the excluded volume of a water molecule (in the limit of \(a_{0} = 0\), we recover the original formula reported by Förster). This expression can be used under the condition that the shapes of the absorption spectra of the donor and acceptor, i.e., of the OD vibration of HDO (\(\sigma_{\text{OD}}\)) and the OH vibrations of H\(_2\)O hydrating OH\(^-\) (\(\sigma_{\text{OH}}\)), do not significantly change when the concentration of OH\(^-\) increases. Indeed, we find strong evidence that the shapes of these spectra hardly change. For \(\sigma_{\text{OH}}\), this can be seen in the inset of Figure 1 and in Figure S1 in the Supporting Information where we show the difference absorption spectra obtained by subtracting the absorption spectrum of H\(_2\)O/D\(_2\)O from the spectra of OH\(^-\)/H\(_2\)O/D\(_2\)O for different OH\(^-\) concentrations. The inset of Figure 1 and the normalized difference spectra of Figure S1 show that the shape of the low-frequency tail of the absorption spectrum of H\(_2\)O molecules hydrating OH\(^-\) is independent of the concentration of OH\(^-\). For the OD vibration of HDO (\(\sigma_{\text{OD}}\)), we find that we can accurately model the transient absorption spectra using the same shape of \(\sigma_{\text{OD}}\) at all studied OH\(^-\) concentrations. This latter observation also implies that the dipole–dipole (Förster) coupling of the OD vibrations and the OH vibrations does not lead to strong mixing of these vibrations and that the perturbative Förster approach remains valid even at high OH\(^-\) concentrations.

Using the Förster model to describe the data implies that we assume that all of the acceptors have the same absorption spectrum and that the acceptors are statistically distributed. The absorption spectrum of the OH stretch vibrations of H\(_2\)O will be inhomogeneously broadened, meaning that different acceptors have different spectra. However, it is not a priori clear whether the species for which the spectra better overlap with the OD band of HDO are closer or further away from the energy donating the OD stretch vibration of HDO, so these effects may well average out. Probably the strongest approximation in using the Förster expression for energy transfer in aqueous environments is that the distribution of distances and relative angles of donors and acceptors is assumed to be statistical. In view of the hydrogen-bond structure of water and hydroxide solutions, this assumption is not valid for short donor–acceptor distances. Hence, the Förster model is expected to break down at high concentrations where short donor–acceptor distances are important.

Assuming that the intrinsic vibrational relaxation process and the Förster energy transfer are independent of each other, the time dependence of the excited population is given by

\[
N(t, \rho) = \exp\left(-\frac{t}{T_{1}} - 4\pi\rho \int_{a_{0}}^{\infty} \text{d}r r^{2} [1 - e^{-k_{\text{FI}}(r) t}] \right)
\]

We fit this expression to the experimental data shown in Figure 5 with \(T_{1}\) and \(R_{0}\) as global parameters. We fix \(a_{0}\) to the previously reported value of 1.5 Å for OD vibrations in aqueous solutions.\(^{10}\) The resulting fits are represented by the solid curves in Figure 5. The intrinsic vibrational lifetime \(T_{1}\) of the OD vibration obtained from the fit is 1.78 ± 0.05 ps, which agrees well with the results of previous studies.\(^{5,21}\) We find a Förster radius of 2.8 ± 0.2 Å, which is similar to the characteristic intermolecular distance of liquid water.\(^{22,23}\)

**DISCUSSION**

We find that the hydration complexes of OH\(^-\) can act as antennae receiving energy from nearby excited vibrations. Because of their broad, nearly continuous absorption spectrum, we can expect that hydrated hydroxide ions can quench the energy of any IR-active vibration between 1000 and 3400 cm\(^{-1}\), like amide and carbonyl vibrations (~1700 cm\(^{-1}\)) that are ubiquitous in (bio)chemistry. As such, hydrated hydroxide ions can act as effective dissipators of excess vibrational energy in aqueous chemical reactions.

Our findings are in line with previous observations for protonated aqueous solutions.\(^{10}\) The hydrated proton shows a similar continuum absorption band as the hydrated hydroxide ion and is also observed to accept energy from excited OD vibrations, thereby accelerating their relaxation.\(^{10}\) For the coupling between OD vibrations and hydrated protons, a larger Förster radius of ~4 Å was found, showing that hydrated protons are even more efficient acceptors of vibrational energy than hydrated hydroxide ions. This difference in the efficiency of Förster energy transfer between acidic and basic solutions is consistent with the stronger absorption of low-frequency OH stretch vibrations in acidic solutions compared to the absorption of these vibrations in basic solutions, at corresponding H\(^+\)/OH\(^-\) concentrations. This difference in absorption strength indicates that the number density and/or transition dipole of low-frequency OH species is higher in acidic solutions than in basic solutions, thus providing more
CONCLUSIONS

We studied the vibrational relaxation mechanism of OD stretch excitations of HDO molecules in NaOH/H₂O solutions using femtosecond mid-infrared transient absorption spectroscopy. We decomposed the measured transient spectra into spectral components corresponding to the excited OD stretch vibration of the HDO molecules, the excited OH stretch vibrations of H₂O molecules hydrating OH⁻ ions, and the hot ground state that results after the vibrational relaxation is completed. This analysis yielded the population dynamics of all three states. With increasing concentration of OH⁻ ions, the excited OH stretch vibrations of hydrated OH⁻ complexes showed that hydrated OH⁻ complexes can serve as effective antennae and dissipators of excess vibrational energy in aqueous chemical reactions.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jpcb.1c02730. Linear absorption spectra; transient absorption signals of solutions of NaOH in H₂O; detailed description of the fitting procedure (PDF)

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Notes

The authors declare no competing financial interest.

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