Hydrogen Bond Reorganization and Vibrational Relaxation in Water Studied with Ultrafast Infrared Spectroscopy

by

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Abstract

Water consists of an extended hydrogen bond network that is constantly evolving. More than just a description of the time averaged structure is necessary to understand any process that occurs in water. In this thesis we study the dynamic regime, which involves fluctuations and rearrangements that occur on the tens of femtoseconds to picosecond time scale. The dynamic regime involves hydrogen bond breaking and forming which interlaces with translations and reorientation and ultimately large-scale reorganization. Our experimental technique is ultrafast infrared spectroscopy of the OX stretch (where X = H, D, or T) of isotopically dilute water. The OX stretch frequency is sensitive to its environment, and loss of frequency correlation provides a picture of local and collective hydrogen bond dynamics. With pump-probe experiments we are also able to measure vibrational relaxation and reorientational dynamics of water.

We present the first infrared linear absorption spectrum of the OT stretch of isotopically dilute tritiated water and compare line shape parameters to the other water isotopologues to provide further evidence that electric field fluctuations properly describe line broadening of the infrared spectrum of water. Measurement of the infrared spectrum of tritiated water is the first step towards an experiment that may be capable of directly monitoring the relative geometry between two water molecules during a hydrogen bond switch. We calculate the change in electric field and transition dipole coupling during an idealized hydrogen switch to determine the correlated frequency shifts one might observe in such an experiment.

To test the proposed vibrational relaxation pathway in isotopically dilute water, we present the first pump-probe of tritiated water and the temperature-dependent lifetime of deuterated water (or HOD in H₂O). For the OT and OH stretch, our experimental findings agree with the proposed mechanism in which the vibrational energy first relaxes to the intramolecular bend. However, evidence from the temperature-dependent measurements of the OD stretch show multiple pathways may be in competition that have different dependencies on temperature. Our results call for further experimental and theoretical studies.
We acquire temperature-dependent 2D IR and pump-probe anisotropy measurements of the OD stretch of HOD in H₂O in order to test if spectral diffusion, which reports on hydrogen bond rearrangements, and reorientation are correlated in water. We compare the temperature dependence of the picosecond decay to a number of measures of structural relaxation and find similar activation barrier heights and slightly non-Arrhenius behavior, which suggests that the reaction coordinate for hydrogen bond reorganization in water is collective.

Thesis Supervisor: Andrei Tokmakoff
Title: Professor of Chemistry
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Chapter 1

Introduction

Set against the shimmering black background of space, the Earth appears as a bright blue marble. Indeed, it is the abundance of water on the surface that gives Earth its blue appearance. Water covers approximately 70% of the Earth’s surface and the human body is made up of 55 to 78% water. Water is intrinsic to life and its unique properties are attributed to an attractive force first suggested at the beginning of last century, the hydrogen bond. The importance of the hydrogen bond has been revealed through the years, but this has not been matched with an equal understanding. Although a number of proposals of the structure of water have been made through the years, a time averaged description is not sufficient in understanding the properties of this important liquid. Dynamic properties that previously could not be explored due to experimental limitations can now be understood with ultrafast infrared spectroscopy. The dynamic regime, which is on the order of tens of femtoseconds to picoseconds, involves hydrogen bond breaking and forming which interlaces with translation and reorientation and ultimately large-scale structural rearrangement. The dynamic regime, which plays an intrinsic role in chemistry and biology, will be better understood with a motion picture of hydrogen bond rearrangement in water.

The first recorded suggestion for an attractive intermolecular force between a hydrogen and an electronegative atom was made in 1902. Alfred Werner proposed that an ammonium salt has a configuration in which a proton linked the ammonia
molecule and the ion. He symbolized the interaction \((\text{H}_3\text{N}...\text{H})\text{X}\) and called it Nebenvalenz, or minor valence [8]. For reference, this observation was made two years after Max Planck introduces quantum theory to explain thermal radiation and 29 years before James Chadwick discovered the neutron. Within the first half of the century of Werner’s suggestion, the time-averaged radial distribution function of water was determined by x-ray diffraction [9, 10]. The first observation of the sensitivity of infrared spectroscopy to hydrogen bonding was made in 1936 [11]. To quote Pimental and McClellan, infrared spectroscopy is, ”...the most sensitive, the most characteristic, and one of the most informative manifestations of the H bond” [12]. Time-averaged experiments have lead to convenient geometrical definitions of the hydrogen bond, for instance, ”two water molecules are chosen as being hydrogen bonded only if their interoxygen distance is less than 3.5 Å, and simultaneously the O-H...O angle is less than 30°” [13]. However, to understand any process that occurs in water, more than just structure-based definition of the hydrogen bond network is required. The dynamics in water are important to a number of processes including the high mobility transport of \(\text{H}^+\) and \(\text{OH}^-\), dynamic solvent effects, electrochemistry, and in biology, the transport and conformational changes of macromolecules.

Dynamic information includes an understanding of how structural rearrangements occur in water. For instance, how does an initial hydrogen bond break and what becomes of the broken hydrogen bond? What is the time scale between the initial rupture of a hydrogen bond and the formation of a new hydrogen bond? Also, how many molecules are important for the description of the hydrogen bond switch? Let us consider a scenario in which (1) a water molecule that is initially hydrogen bonded has enough thermal energy to rupture a hydrogen bond, (2) with Brownian motion the water molecule undergoes diffusive motion until (3) the molecule is in a favorable geometry for hydrogen bond formation with a new partner. In this example the switching event occurs on a slower time scale than the fastest intermolecular motions in water and only one water molecule is important for the description. An experimental technique that has both the necessary temporal resolution and structural sensitivity to the motions just described is essential for a description of the structural
reorganization of water.

Ultrafast infrared spectroscopy is the tool we will use to explore the evolving structure of water. Ted Maiman invented the first operating laser in the middle of 1960 [14] and Ali Javan was credited with the first gas laser later that year [15]. In chemistry and physics, the laser has been an invaluable tool in understanding fundamental properties of molecules and macromolecules. Since the advent of the laser, the duration of the shortest pulse has been whittled down towards a single-cycle. What drives the quest for shorter pulses? Elementary chemical processes in gases and liquids take place on time scales ranging from femtoseconds to picoseconds. The fluctuations of the hydrogen bonding network of water are unusually fast compared to other simple liquids. The dynamics of the hydrogen bond network of water can only be captured with pulses shorter than the fastest events. Likewise, photographing more than just a blur when a balloon bursts or a bullet impacts an apple requires a minimum "shutter speed". Ultrafast spectroscopy, which interrogates the optical response of a system in the time-domain, is the "camera" that can capture the dynamics of water. In this thesis I present my part in unveiling the enigmatic beauty of the evolving hydrogen bond network of water.

1.1 Hypotheses of the Structure of Water

Defining the structure of any liquid is a complex problem since the instantaneous structure is disordered and the liquid is undergoing continuos reorganization. The time-averaged structure of water has been debated through the years [16], beginning even before the recognition of the hydrogen bond. In one view, the "mixture" model, a number of stable configurations exist in water. In the other, "continuum" model, the environment undergoes fluctuations about an average structure. In 1892, Roentgen proposed the first mixture model in which water consisted of a "bulky" ice-like component and a less bulky "normal liquid" component [17]. Frank and Wen suggested the "flickering cluster" mixture model, in which water consists of non-hydrogen bonded water molecules and hydrogen-bonded clusters. Hydrogen bonding is highly coopera-
tive in this model, which predicts that if one hydrogen bond breaks a cluster dissolves [18]. In 1962, Nemethy and Scheraga interpreted the radial distribution function in terms of a two component mixture model that consisted of ice-like water and water molecules with broken hydrogen bonds [19]. The interstitial model is similar in that the average structure of ice remains upon melting, but non-hydrogen bonded water molecules occupy the cavities of the lattice structure [20]. Mixture models were extended to include multiple structural components: four-, three-, and two-coordinated hydrogen bonded entities [21]. Models that can be considered "continuum" models, in that they consist of, on average, four-coordinated hydrogen bonded molecules with differing degrees of distortion, were proposed by Pople [22] and Bernal and Fowler [23]. Although the aforementioned mixture and continuum models have implications for the time-dependence of water structure (see below), they were developed to describe measures of the average structure.

Eisenberg and Kauzmann recognized that time scale must be considered when defining the structure of water [16]. In other words, the term "structure" has a different definition depending on the time interval of the measurement. They define three different structures depending on the exposure time, which are separated by the average period of a vibration, or $\tau_V$, and the average time between two displacements, or $\tau_D$. The first structure is the instantaneous structure, or I-Structure, which is the structure that will be observed if the time exposure is short compared to $\tau_V$. To observe the vibrationally averaged structure, or V-Structure, the time exposure is longer than $\tau_V$ but shorter than $\tau_D$. Finally, the last structure is the diffusionally averaged structure, or D-Structure, where the exposure time is longer than $\tau_D$ [16].

Time averaged experiments, such as x-ray diffraction, report on the D-Structure of liquid water. The ultrafast infrared experiments that will be described in this thesis report on the V-Structure and, therefore, provide a picture of intermolecular motion in water on a longer time scale than the vibrational period of the stretch ($\sim10$ and 13 fs for the OH and OD stretch, respectively). Not surprisingly, experimental techniques sensitive to different time scales have lead to debate in the literature on the structure of water. For instance, recently x-ray absorption spectroscopy (XAS) and nonresonant
x-ray Raman scatter (XRS) at the oxygen K-edge, techniques sensitive to distortions within the first coordination shell on time scales shorter than the vibrational period, or the *I-Structure*, have been measured for liquid water [24]. The authors concluded that water molecules are not on average in a tetrahedral geometry, rather each water molecule is in two hydrogen-bonded configurations with one strong donor and one strong acceptor. However, theoretical interpretations of the data have shown that the results are consistent with a unimodal density of tetrahedral liquid water [25].

Two models were developed circa 1980 that considered not only the time averaged structure, but also dynamics: the random network model [26] and the correlated-site percolation model [27]. In the percolation model, liquid water is above the percolation threshold and, therefore, consists of a extended hydrogen bond network. However, there is correlation between neighboring sites in hydrogen-bond coordination number. For instance, four coordinated water molecules, $f_4$, clump together to form patches of low density. In this model, transport is dominated by the mobile fraction, which is water molecules with $f_0$ and $f_1$ coordination. However, the model does not include a description of how hydrogen bond rearrangements occur. The random network model proposed by Sceats and Rice [26] was an extension of the model proposed by Pople [22] and, therefore, predicts that on average water is four coordinated. The authors recognized that it is useful to view two different time scales for water: the inertial motion that includes librations and hindered translations and a slower time scale that includes diffusion and other relaxation processes.

Around the same time, Stillinger and Weber proposed analyzing both the thermodynamics and dynamics of water in terms of inherent structures [28]. In this treatment, the potential energy surface consists of a number of wells, or "inherent structures", and there are distortional displacements from the wells due to vibrations. Fundamental structural rearrangement can be viewed not as changes in instantaneous structures, or *I-Structures*, but rather by transitions between inherent structures with the vibrational components removed. For larger systems, inherent structures are determined by quenching along a trajectory. Within the inherent structure framework, Ohmine et al. determined the moment ratio, which reports the average displacement.
of the molecules in a system, in a transition between inherent structures to determine if motion in water is collective, or delocalized [29]. For a system of 216 molecules, and upon averaging over 400 inherent structure transitions, they found ~20 molecules move simultaneously indicating molecular motions are quite collective in water [29]. Sciortino et al. performed another interesting study using inherent structures to understand the effect of defects on molecular mobility in liquid water [30]. They found that at low temperatures five-coordinated water molecules provide low-energy paths to transition between different tetrahedral local arrangements [30].

A simple question arises: what do the different structural models predict for the nature of a broken hydrogen bond? Figure 1-1 shows a pictoral free energy surfaces depicting two possibilities. In this figure, HB and NHB refer to water molecules in a stable and broken or highly-strained hydrogen bond, respectively. For continuum models, the average structure is a four coordinated water molecule and therefore the broken hydrogen bond is a transition state between switching hydrogen bonding partners, as shown in the lower blue curve. Defining a reaction coordinate for hydrogen bond switching would only be reasonable for the continuum model. If a water molecule is anything but, on average, four coordinated the broken hydrogen bond or a highly strained configuration would be stable, which is implied by the mixture model and shown as the green curve. In other words, a broken hydrogen bond would persist beyond the fastest intermolecular motions of water. Ultrafast infrared experiments and molecular dynamics (MD) simulations performed by the Tokmakoff group, which will be discussed below, have shown strong evidence that the nature of the broken hydrogen bond more closely follows the implications of the continuum model.

There have been recent theoretical developments in the temporal description of a broken hydrogen bond and the mechanism of hydrogen bond switching. With molecular dynamics (MD) simulations with the SPC/E model for water [31], Laage and Hynes found reorientation occurs not by small diffusive steps but rather through large amplitude angular jumps that exchange a hydrogen bond with a small contribution from the diffusive motion of the hydrogen bond frame [1, 32]. The mechanism is shown in Figure 1-2. The green lines indicate hydrogen bonds between the three
Figure 1-1: Two possibilities for the free energy surface of water

Water molecules pictured and the blue lines indicate hydrogen bonds with water molecules not shown. The first and second hydrogen bond acceptors, $O^a$ and $O^b$, become overcoordinated and undercoordinated, respectively, due to fluctuations in the hydrogen bond network. $O^b$ approaches $H^*$ from the second solvation shell as $O^a$ moves out of the first solvation shell (Figure 1-2A) leading to the $R_{O^aO^a}=R_{O^bO^b}$ conformation (Figure 1-2B) where the $H^*$ jump takes place. On average the jump of $H^*$ from $O^a$ to $O^b$ occurs within 250 fs over an angle of 60°. The molecular jump mechanism agrees with the blue curve shown in the pictorial free energy surfaces in Figure 1-1. Interestingly, since at the bifurcation angle the water molecule that acts as the donor is five-coordinated, the mechanism also agrees with the theoretical work of Sciortino et al. [30]. Laage and Hynes also looked at the temperature dependence of reorientation and found that the large amplitude jumps and diffusive motion both contribute at every temperature, but large amplitude jumps are dominant [32]. Finally, they generated the free energy surface as a function of the change in the initial and final acceptor H-bond coordination number, the change in the distance from the rotating water and the initial and final acceptors, and the angular coordinate of the rotating water molecule. They found the translational motion of the first and second hydrogen bond acceptors had the largest contribution to the free energy barrier [32].
The mechanism of hydrogen bond switching has also been studied with a "first principle" approach that combines \textit{ab initio} based water potential with explicit quantum treatment of molecular motion [33]. The computational method included path-integral molecular dynamics [34] to calculate structural quantities and centroid molecular dynamics [35] to calculate dynamic and spectroscopic properties with the \textit{ab initio} based TTM3-F water potential [36]. Paesani first shows the applicability of the model by calculating temperature-dependent linear and non-linear infrared spectra of the OD stretch of HOD in H$_2$O. Calculation of the molecular mechanism of hydrogen bond switching indicated that, on average, the molecules undergo a large angular jump, agreeing with the mechanism proposed by Laage and Hynes [1, 32]. However, a large number of different pathways were observed, indicating that a simple description of molecular motion was not applicable [33].

1.2 Infrared Spectroscopy of Hydrogen Bonded Liquids

Ultrafast mid-infrared spectroscopy has both the time and structural sensitivity necessary to study structural rearrangement in water. A red-shift in the fundamental and higher harmonics of the stretching mode, $\nu_s$, upon hydrogen bond formation was first observed around 1930 for a number of molecules that contain an intramolecular hydrogen bond [37, 38]. In 1932, a shift in $\nu_s$ with concentration and temperature
was observed in solutions of alcohols [39]. In water, the OH stretch of isotopically
dilute HOD in D$_2$O red shifts from 3707.47 cm$^{-1}$ in the gas phase [40] to $\sim$3310 cm$^{-1}$
in hexagonal ice, a $\sim$400 cm$^{-1}$ shift! In liquid water the peak of the OH stretch
absorption band is $\sim$100 cm$^{-1}$ blue shifted from the value for ice, indicating that the
hydrogen bond strength is intermediate between ice and gas phase but more similar
to ice. We show the linear absorption spectrum of isotopically dilute HOD in H$_2$O
and HOD in D$_2$O from 1000 to 4000 cm$^{-1}$ at room temperature in Figure 1-3 with the
stretch ($\nu_S$), bend ($\delta$), and combination bands within this spectral range indicated.

![Figure 1-3: Linear absorption spectrum of isotopically dilute HOD in H$_2$O (green)
and HOD in D$_2$O (blue) in the mid-infrared region with the transitions indicated in
blue for D$_2$O, green for H$_2$O, and black for HOD. The spectra are not offset.](image)

There have been many attempts to link the shift in frequency with hydrogen
bonding to structure and thermodynamic properties. Only parameters that correlate
with the vibrational frequency will be elucidated with infrared spectroscopy. Rundle
and Parasol first noticed a relation between $\nu_S$ and R$_{OO}$, the O...O distance between
adjacent water molecules, for hydrogen bonded solids [41]. Novak assembled a com-
prehensive list of $\nu_S$ and R$_{OO}$ for strong, medium, and weak hydrogen bonded salts
[42]. This correlation is based on Badger’s rule, which defines a relationship between
the force constant for stretching and the equilibrium intramolecular bond length [43].
Wall and Hornig used the experimentally determined relationship between $\omega_{OH}$ and
R$_{OO}$ from hydrogen bonded crystals to recast the Raman spectrum of isotopically
dilute water in terms of $R_{OO}$ [44]. The peak corresponded to $R_{OO} = 2.85 \text{ Å}$, which agreed within 0.04 Å with the first peak in $g_{OO}$ from x-ray diffraction [44].

Approximately 35 years later, the correlation between $\omega_{OH}$ and $R_{OO}$ was tested for liquid water with mixed quantum-classical molecular dynamics (MD) simulations [45, 46, 47, 48]. The correlation on the red side was decent but dispersion was observed on the blue side (where red and blue refer to low and high frequency compared to band center, respectively). In the study by Lawrence and Skinner, the authors also decompose the frequency distribution into molecules with and without hydrogen bonds, determined by the geometric criteria given earlier. They found that although the distributions overlap, water molecules without a hydrogen bond clearly absorb on the blue side of the line width [45, 46].

Eaves et al. explored the correlation between $\omega_{OH}$ and a number of structural parameters: $R_{OO}$, $\alpha$ (the angle between the intramolecular O-H bond and $R_{OO}$), tetrahedrality, and the electric field and reported the correlation coefficient, $\rho$ [46]. In this treatment, $\rho = 1$ indicates perfect correlation, $\rho = -1$ is perfectly anti-correlated, and $\rho = 0$ indicates no correlation. They found $\omega_{OH}$ and $\cos(\alpha)$ ($\rho = -0.47$) are less correlated than $\omega_{OH}$ and $R_{OO}$ ($\rho = 0.77$), and there is even less correlation between $\omega_{OH}$ and the tetrahedrality ($\rho = -0.25$). However, the correlation between $\omega_{OH}$ and the electric field from the hydrogen bonded neighbor, $E_0$, is quite good ($\rho = 0.89$) and if all the molecules in the simulation are included in the electric field calculation, $E$, the correlation is near unity [46]. From ab initio calculations, Hermansson first observed a relationship between $\nu_S$ and the electric field for OH\textsuperscript{-}, LiOH-H\textsubscript{2}O, and various water-ion/neutral clusters and crystals [49, 50, 51, 52]. In 1998, Buch and co-workers determined an empirical relationship between the shift in vibrational frequency and the electric field projected in the direction of the OH bond from experimental H\textsubscript{2}O cluster studies [53] and in 2004 Corcelli determined a linear relationship between the electric field and frequency by performing ab initio calculations on clusters selected from MD simulations of liquid water [54]. The relationship has been refined by the Skinner group and it is found to be quadratic [55, 56]. These findings cast the system bath Hamiltonian in terms of the Stark shift Hamiltonian, which implies vibrational
dephasing is a result of electric field fluctuations [46]. However, since the electric field depends on both $R_{OO}$ and $\alpha$, $\omega_{OH}$ is related to the local structure and, therefore, the vibrational frequency is a viable tool to study local dynamics [46].

1.3 Recent Experimental and Theoretical Results

1.3.1 Frequency Fluctuations

An isotopically dilute system provides a probe of local structural dynamics, as resonant intermolecular transfer is turned off, and it imitates neat water for any process largely dominated by electrostatics. Spectral diffusion of the OH stretch of HOD in D$_2$O was initially reported in 1991 by Graener et al. with spectral hole burning experiments performed with relatively narrow-band mid-IR pulses [57, 58]. In a spectral hole burning experiment, a pump pulse excites a narrow spectral hole in the absorption band and a weak probe pulse monitors the spectral hole as it broadens and shifts towards band center as a result of spectral diffusion. However, there is an inherent time-bandwidth trade-off in that spectrally narrow pulses have relatively long pulse durations. The initial experiments performed with 1+ ps pulses lead the authors to conclude that the OH stretch absorption band contained several sub-bands, suggesting that the structure of water is better described with a mixture model [57, 58]. Later spectral hole burning experiments with femtosecond pulses found a $\sim$1 ps correlation time and did not find evidence for sub-bands [59, 60, 61].

Stenger et al. reported the first photon-echo measurements of isotopically dilute water in 2001 [62]. Photon echo experiments are capable of discerning static, or inhomogenous, and dynamic, or homogenous, line broadening mechanisms and are sensitive to frequency fluctuations. Contributions to homogenous broadening, denoted as $T_2$, are population relaxation ($T_1$), pure dephasing ($T_2^*$), and reorientation ($\tau_{OR}$). Population relaxation is an inherent property of individual molecules. Pure dephasing and reorientation are both ensemble averaged effects, in that randomization occurs through interactions with other molecules. Inhomogeneous broadening is
a consequence of the presence of distribution of frequencies, or, configurations. Conventional first order, or linear spectroscopy, is not capable of discerning between these broadening mechanisms. This quickly diminishes the hopes to extract structural information from a linear absorption spectrum and has provided fuel for the debate on the dynamic structure of water. For example, the observation of an isosbestic point in the temperature-dependent Raman spectrum lead to arguments for the "mixture" model [63]. As an aside, Smith et al. showed that an isosbestic point dose not imply a multi-component mixture [64].

Stenger et al. performed a three-pulse photon echo experiment on HOD in D$_2$O with the second and third pulses time coincident [62]. The first pulse creates a coherence and the second pulse excites a population grating, which is instantly probed with the time coincident third pulse. The experiment is primarily sensitive to the pure dephasing time, $T_2^*$. The authors found $T_2^*=90$ fs, which is in between the dephasing time expected for purely homogeneous (66 fs) and purely inhomogeneous broadening (132 fs). The significant amount of homogeneous broadening suggests there are very rapid spectral fluctuations, which were estimated to be on a $\sim 30$ fs time scale [62]. Yeremenko et al. [65] performed a heterodyne detected two-photon echo experiment on HOD in D$_2$O. The goal of the measurement was to track the transition frequency fluctuation correlation function, $C_\omega(\tau)=\langle \delta\omega(\tau)\delta\omega(0) \rangle$, to understand the structural fluctuations of the hydrogen-bonded network. The two-point frequency correlation function is exact if frequency fluctuations are Gaussian. The authors modeled their results with a bi-exponential frequency correlation function with a 130 fs and 900 fs time constant attributed to hydrogen bond breaking and rearrangement of the hydrogen bond network, respectively [65].

Stenger et al. reported the first three-pulse echo peak shift (3PEPS) measurement of HOD in D$_2$O [66]. With 90 fs pulses the authors found the spectral diffusion contained a slow component on the order of 5-15 ps [66]. However, Yeremenko et al. found interference from the background D$_2$O greatly affects photon echo experiments at long time delays [67]. The Tokmakoff group measured the three-pulse photon echo peak shift of HOD in D$_2$O with 50-fs pulses spanning the line width of the OH
The temporally short pulses ensure that the fastest dynamics are captured. In three-pulse photon echo peak shift measurements, the time $\tau_1$, or the time between the first and second pulses, that maximizes the signal for each $\tau_2$ is denoted as $\tau_1^\ast$. The decay of $\tau_1^\ast$ with $\tau_2$ indicates loss of inhomogeneity due to spectral diffusion. The signal is integrated over $\tau_3$ so the measurement is an average over all microscopic environments. The experimental peak shift displayed a fast decay within 75 fs, a recurrence at 150 fs followed by a 1.2 ps exponential decay. With a response function formalism [70], the peak shift was self-consistently modeled with the vibrational lifetime, anisotropy decay, and linear absorption spectrum to extract the two-point frequency correlation function, $C_\omega(\tau)$. The best fit resulted in slightly different amplitudes and timescales with a fast decay in 50 fs, a recurrence at 180 fs, and 1.4 ps exponential decay [69]. We show the best fit correlation function in Figure 1-4. The difference lies within the assumptions made to relate the two-point correlation function to the peak shift measurement.

Figure 1-4: (blue) Fit to the integrated pump-probe (PP) anisotropy for the OH stretch of HOD in D$_2$O with the decay ascribed to the librational motion indicated. (red) Best fit frequency correlation function to the 3PEPS measurement with the recurrence due to the hydrogen bond stretch indicated.
In order to interpret the experimental results, one must first understand how structural dynamics are related to the frequency correlation function. As we discussed in Section 1.2, with MD simulations Eaves et al. found that the frequency of the stretch correlates nearly perfectly with the projection of the electric field from all the molecules in the simulation on the proton of interest \( (E) \) \[46\]. The electric field is dominated by the hydrogen bond partner \( (E_0) \). In order to understand the dynamics that drive vibrational dephasing, the time correlation function (TCF) of \( \omega_{OH} \) and the proposed structure parameters should be compared. At early times, the TCF of \( \omega_{OH}, E, E_0, \) and \( R_{OO} \) display similar behavior, including a recurrence in a similar time period, and the TCF of \( E \) follows that of \( \omega_{OH} \) nearly perfectly. The beat is not present in the TCF of the entire electric field minus that of the hydrogen bonded partner. Therefore, the local electric field fluctuation from the hydrogen bonded partner produces the beat and the molecular origin is the intermolecular hydrogen bond stretch. Interestingly, the long time decays of all the structural parameters studied show a similar behavior on a time scale greater than 200 fs, even those that were not similar to the early time behavior of \( C_\omega(\tau) \) and had poor correlation with \( \omega_{OH} \). This suggests that large scale, or collective, rearrangements dominate on the longer timescales and a picture of local dynamics is lost as one moves into the kinetic regime (see Figure 1-4) \[68, 46\].

The broadband pulses in the aforementioned photon echo experiments ensure the fastest motions are captured. However, all microscopic environments contribute to the integrated signal. Two dimensional infrared spectroscopy (2D IR) is a technique capable of providing information on spectrally distinct environments while retaining fast temporal resolution. Put another way, the previous homodyne experiments provided a glimpse into local structural dynamics averaged over all hydrogen-bonded environments and heterodyned 2D IR measurements can report on the local structural dynamics in spectrally distinct hydrogen-bonded environments. 2D IR is a third order technique in which three pulses create a third order nonlinear polarization that radiates a signal field. For well-separated pulses, the first pulse creates a coherence and after the evolution time, \( \tau_1 \), the second pulse creates a population state. In the
Figure 1-5: (Top) Model frequency correlation function, which is a single exponential with a 500 fs time constant. (Bottom) Four model 2D surfaces calculated with the model frequency correlation function at the waiting times: $\tau_1 = 0, 500, 1000, \text{ and } 2000$ fs (indicated with red circles on the frequency correlation function).

pump-probe geometry, which we will describe in detail in the next chapter, the third pulse, which arrives after a waiting time, $\tau_2$, relative to the second pulse, acts also as the local oscillator that heterodynes the signal field. $\tau_1$ is numerically Fourier transformed to create the $\omega_1$ axis and the signal and local oscillator are dispersed with a grating to create the $\omega_3$ axis. Example 2D surfaces are shown in Figure 1-5 for a system with peak absorption at 2500 cm$^{-1}$ and 160 cm$^{-1}$ line width at full-width half maximum (FWHM). In the example the dynamics are the same across the line shape. Each 2D surface displays a positive $\nu=1\leftarrow0$ peak and a negative $\nu=2\leftarrow1$ peak.
that is anharmonically shifted in $\omega_3$. The line shape as a function of waiting time, $\tau_2$, relays spectral diffusion. The 2D surfaces shown in Figure 1-5 were calculated at four waiting times: $\tau_2 = 0$ fs, 500 fs, 1 ps, and 2 ps. The time points are marked on the frequency correlation function for the model system, which is a single exponential with a 500 fs correlation time. At $\tau_2 = 0$ fs, the system retains memory of the initial excitation frequency, and therefore there is positive correlation along the diagonal. With increasing waiting time frequency memory is lost and the peaks appear uncorrelated. The homodyne photon echo peak shift can be recovered from the absolute value of the 2D IR spectra projected onto the $r_1$ axis. The decrease in amplitude with $\tau_2$ is a result of population relaxation and molecular reorientation. The projection on the $\omega_3$ axis is equivalent to the dispersed pump probe. For waiting times longer than the lifetime, $T_1$, thermalization from population relaxation needs to be accounted for in water.

The 2D IR spectrum of HOD in D$_2$O as a function of waiting time has been measured by the Tokmakoff group [71, 72, 73]. Even at $\tau_2 = 0$ an asymmetry between the red and blue sides is evident. For an inhomogeneous system, the anti-diagonal width ($\omega_1 + \omega_3 = 2\Omega$) is related to the homogeneous line width [33]. The broader anti-diagonal width on the blue side indicates quicker dephasing relative to the red side. The waiting time dependence was quantified with the first moment of the positive signal distribution for slices in $\omega_1$, which is shown in Figure 5-12. A marked difference is observed as the blue side decays monotonically and reaches an asymptotic value in 100 fs and on the red side an oscillation is observed at $\sim$130 fs followed by a slower decay to band center. This observation, along with others, resulted in the conclusion that hydrogen bonds are broken only fleetingly, or, the NHB is a transient species that appears due to fluctuations about a hydrogen bond or in concerted switching of hydrogen bonded partners. The transient behavior suggests that hydrogen bond switching is concerted. The observations from the ultrafast measurements have been corroborated with molecular dynamics simulations [71, 73, 74].

The nonlinear experiments described thus far have been on the OH stretch of HOD in D$_2$O. In 2001, Kropman et al. performed a spectral hole burning experiment
on the OD stretch of HOD in H$_2$O and found a frequency correlation time of 400 fs [75]. Laenen et al. also performed a spectral hole burning experiment and found not only multiple sub-bands but a 2 ps correlation time [76]. Asbury et al. acquired a waiting time series of 2D IR surfaces of the OD stretch of HOD in H$_2$O [77]. The authors characterized spectral diffusion by determining the dynamic line width, or, the full width at half maximum (FWHM) of a slice along $\omega_1$ taken at the maximum in $\omega_3$, at each waiting time. Similar to the treatment by Fecko et al. [69] the 2D surfaces were modeled with a response function formalism to determine the best fit frequency correlation function, $C_\omega(\tau)$. The decay constants of the best fit triexponential correlation function were 48 fs, 400 fs, and 1.4 ps with the amplitudes 41%, 15%, and 44%, respectively [77]. The long time decay is in reasonable agreement with the best-fit correlation function determined for the OH stretch of HOD in D$_2$O [69], however, the recurrence in the correlation function from the hydrogen bond stretch was not experimentally observed for the OD stretch.

Theoretical modeling has been indispensable for interpreting experimental results and providing a microscopic picture of the dynamics in water. As discussed in Section 1.2, the stretch frequency is correlated to the electric field of the surrounding

Figure 1-6: First moment as a function of waiting time, $\tau_2$ of the positive signal distribution of $\omega_1$ slices on the red and blue side of the OH stretch 2D IR line shape.
With the development of an empirical relationship between the frequency and electric field, frequencies can be efficiently calculated "on the fly" during molecular dynamics (MD) simulations. Skinner and coworkers have been instrumental in developing and refining an approach that combines electronic structure methods and molecular dynamics simulations (ES/MD) to study linear and nonlinear infrared spectroscopy of water [54, 7, 78, 55, 56]. Initial work by Corcelli et al. [54] based the approach to determine an empirical relationship between frequency and electric field upon similar approaches for water clusters [53, 79, 80] and N-methylacetamide (NMA) in water [81, 82, 83, 84]. Here we will present a basic outline of the approach. During a MD simulation of a water model, for instance the SPC/E model, clusters are randomly selected and the electric field at the proton of interest projected along the intramolecular bond is determined and the frequency is directly calculated by generating the one-dimensional potential energy curve and solving the relevant Schrödinger equation. Clusters are chosen from a MD simulation and not optimized in order to better estimate geometries that would be found in the liquid. Initial work found an approximately linear relationship between the \textit{ab initio} frequency and electric field [54], which can be used with MD simulation to calculate the frequency correlation function and distribution of frequencies. This approach can also be applied to determine an empirical relationship between the electric field and transition dipole or molecular polarizability for calculations of the infrared and Raman spectra. The empirical relationship has been refined by the Skinner group and a quadratic relationship between the frequency and electric field has been found to be more accurate [55, 56].

The frequency correlation function has been calculated with MD simulations using a number of water models [46, 85, 86]. To obtain better statistics, Corcelli et al. performed a calculation with pure H$_2$O, rather than isotopically dilute water, such that every OH bond in the simulation could be treated as the OD vibration of interest for the purpose of comparing to experimental results on the OD stretch of HOD in H$_2$O. They found the frequency correlation function calculated with the fixed charge SPC/E and TIP4P [87] models displayed an oscillation due to the hydrogen bond stretch but the polarizable SPC/FQ [88] model did not. However, after fitting the
calculated frequency correlation functions with triexponentials, they found the long time decay of the SPC/FQ correlation function agreed much better with experiment: 1.45 ps for the SPC/FQ model compared to 900 fs for the SPC/E model. Schmidt et al. calculated the infrared and Raman line shape and observables from third-order experiments with MD simulations with the SPC/E, TIP4P, SPC/FQ, TIP4P/FQ [88], TIP5P/E [89], and Dang-Chang [90] water models and determined that the SPC/E model gave the best overall agreement with experiment [86].

1.3.2 Vibrational Relaxation

Pump-probe (or transient absorption) experiments are capable of measuring population relaxation and molecular reorientation. The first two interactions come from the first pulse, or pump, therefore $\tau_1 = 0$. The signal is measured as a function of $\tau_2$, the delay between the pump and probe pulse, and the measurement detects pump-induced transmission changes of a weak probe pulse. The signal is emitted in the direction of the probe pulse and, as in FTIR measurements, the decrease in transmission is due to the interference between the probe pulse and out of phase emitted signal. The vibrational lifetime of an excited vibration can be measured by the decay of the magic angle pump probe, or $S_{iso}$, which is free from molecular reorientation. Magic angle refers to the relative polarization between the linearly polarized pump and probe pulses. $S_{iso}$ can also be constructed from the separately measured parallel and perpendicular pump-probes, as: $S_{iso}(\tau_2) = (S_{zzzz} + 2S_{zzzy})/3$. Since multiple vibrational levels are accessible, the dispersed pump probe displays a positive peak from the $\nu=1\leftarrow0$ ground state bleach and stimulated emission and an anharmonically shifted negative peak from the $\nu=2\rightarrow1$ induced absorption. The signal can be integrated (homodyne) or dispersed into frequency components.

In isotopically dilute water, the lifetime of the OH stretch of HOD in D$_2$O has been reported to be 0.5 to 1 ps [91, 92, 76, 93], which is shorter than the measured lifetime of the OD stretch of HOD in D$_2$O that has been reported to be 1.45 to 2 ps [92, 75, 76, 94, 95]. The lifetime of the bend, $\delta$ of HOD in D$_2$O, was found to be 380 fs [96]. Temperature-dependent measurements of the lifetime of the OH stretch
[97, 98, 99] and OD stretch [100, 101, 102] have found the lifetime generally increases with increasing temperature. Theoretical studies of vibrational relaxation of the OH stretch of HOD in D₂O have found the dominant pathway for relaxation is through the overtone of the HOD bend a lifetime of 7.7 ps [103], 2.7 ps [104], and 2.3 ps [105]. A theoretical study of vibrational relaxation of the OD stretch of HOD in H₂O performed similarly to the theoretical investigations of the OH stretch, found the primary pathway was through the fundamental of the bend (19.05 ps) or directly to the ground state (5.04 ps) depending on if the solvent was treated as rigid or flexible, respectively [106]. For both the OH and OD stretch the calculated lifetime is considerably longer than the measured lifetime. Vibrational relaxation of νₛ of pure H₂O [107, 108, 109], δ of pure H₂O [109, 110, 111], the combination band, or νₗ + δ, of pure H₂O [112], the high frequency portion of νₗ of pure H₂O, and νₛ of pure D₂O [113] have also been measured to understand energy flow in pure water.

1.3.3 Reorientation

Reorientation in water can be studied with time dependent pump-probe anisotropy determined from the parallel and perpendicular pump-probes:

\[ R(\tau_2) = \frac{(S_{zzzz} - S_{zyzy})}{(S_{zzzz} + 2S_{zyzy})}. \]  

(1.1)

The anisotropy is proportional the second rank Legendre polynomial orientational correlation function: \( R(\tau_2) = (2/5)C_R(\tau_2) \), where \( C_R(\tau_2) = \langle P_2[\hat{\mu}(t)\cdot\hat{\mu}(0)] \rangle \), \( P_2 \) is the second order Legendre polynomial, and \( \hat{\mu} \) is the transition dipole. Other measurements sensitive to reorientation include dielectric relaxation (DR) [6], THz time domain spectroscopy (THz-TDS) [5], NMR [4, 114], optical Kerr effect (OKE) spectroscopy [3], and quasi-elastic neutron scattering (QENS) [115]. With the proper assumptions, DR and THz-TDS are related the first order Legendre polynomial and NMR, OKE, and QENS are related to the second order Legendre polynomial. If reorientation occurs diffusively, the relationship between the first and second order decay time is: \( \tau_{R,2} = \tau_{R,1}/3 \).
The infrared pump-probe anisotropy has been measured for the OH stretch of HOD in D$_2$O [116, 117, 118, 93]. The pump-probe anisotropy measured by Loparo et al. with $\sim$50 fs pulses, which was integrated over the $\nu=1\leftrightarrow0$ peak, showed to two distinct decay components. The decay was fit with a biexponential with the time constants 60 fs and 3 ps [93]. The fast, or inertial, decay was ascribed to the libration, or hindered rotation, and slow decay to diffusive reorientation. The fit to the integrated anisotropy is shown in Figure 1-4 along with the best fit frequency correlation function determined by Fecko et al. [69]. The pump-probe anisotropy of the OD stretch of HOD in H$_2$O shows a long time decay of $\sim$2.5 ps [119, 120, 121, 95].

Frequency resolved pump probe anisotropy of isotopically dilute water has been measured to study the rotational dynamics of water molecules initially excited in different environments [119, 121]. Bakker et al. measured the pump-probe anisotropy of the OD stretch of HOD in H$_2$O pumping at the center and blue wing of the line width [119]. They find that when pumping at the center of the line width the anisotropy decay is independent of frequency. However, when pumping on the blue wing, the anisotropy of the molecules on the red side of the line width drops to 0.2 by the initial time point, 200 fs. This observation suggests that molecules initially excited on the blue wing must reorient to return to the red side of the line width [119]. Moilenan et al. measured the dispersed pump-probe (DPP) anisotropy of the OD stretch of HOD in H$_2$O [121]. Although spectral information is obscured within the first 200 fs due to non-resonant effects, the authors extrapolate to 100 fs and find there is a greater initial drop in anisotropy on the blue side of the line width ($\sim$0.36 at 2500 cm$^{-1}$ compared to $\sim$0.32 at 2625 cm$^{-1}$) and after 200 fs, the anisotropy decay is frequency independent. This observation agrees with MD simulations performed by Lawrence and Skinner, which show there is a correlation between frequency and reorientation [45]. The authors found a larger amplitude inertial reorientation in the orientational correlation function for molecules absorbing on the blue side of the line width [45], which corresponds to molecules in the NHB configuration.
1.4 Thesis Outline

Previous studies of the hydrogen bond rearrangements of water have concluded that hydrogen bonds are broken only fleetingly as hydrogen bonding partners rearrange. This observation has been made experimentally in our group with a waiting time series of two-dimensional infrared (2D IR) surfaces of the OH stretch of HOD in D$_2$O. 2D IR is a structural sensitive technique that offers ultrafast time resolution. The fleeting nature of broken hydrogen bonds observed experimentally agrees with the molecular jump mechanism for hydrogen bond rearrangement proposed with molecular dynamics simulations. However, this agreement is only circumstantial and further experiments are necessary to understand the relationship between reorientation and hydrogen bond rearrangements. In this thesis we present first and third-order experiments of isotopically dilute water aimed to understand vibrational population relaxation, reorientational motion, and the relationship between reorientation and hydrogen bond reorganization in water measured with the frequency correlation function. We also use temperature as an experimental handle in the measurement of all three phenomena.

In Chapter 2 we detail our method for generating ultrafast mid-infrared pulses and performing third-order experiments. Traditionally, 2D measurements have been acquired in the background free "boxcar" geometry, in which the three input fields are at three corners of a box and, to conserve momentum, the signal is emitted in the fourth corner. However, the third order experiments presented in this thesis were acquired in the pump-probe geometry, in which a collinear pump pair creates a vibrationally excited state and the third interaction is also the local oscillator. We will therefore detail the pump-probe interferometer and discuss aspects unique to the data collection geometry.

In Chapter 3 we present the first infrared linear absorption spectrum and in Chapter 4 the first non-linear measurement of radioactive isotopically dilute HOT in H$_2$O. Experiments with isotopically dilute water provide a local picture of hydrogen bond dynamics but information on the environment is inferred, as dephasing is controlled
by system-bath interactions. Monitoring the waiting time dependence of the cross peak, which requires the interaction of two spectrally distinct species, in 2D IR measurements of isotopically dilute HOD and HOT in H₂O is a way to directly study the relative motion between two water molecules in solution. As a step in the direction we measured the first linear absorption spectrum of tritiated water. We compare the linear absorption of the OT stretch, or \( \omega_{OT} \), with \( \omega_{OD} \) and \( \omega_{OH} \) of isotopically dilute water to test the reduced mass scaling relationships for frequency, line width, and extinction coefficient. Harmonic scaling relationships are found to be inadequate, and deviations are discussed in terms of electrical and mechanical anharmonicity. The inverse scaling of the line width with the square root of reduced mass provide further evidence that electric field fluctuations properly describe the line broadening in water. In the chapter we also present model calculations of the correlated frequency shifts of \( \omega_{OT} \) and \( \omega_{OD} \) relevant for the proposed experiment by calculating on site frequencies and off-diagonal coupling during an idealized hydrogen bond switch. Finally, we present Gaussian calculations used to determine the intermolecular coupling between \( \omega_{OT} \) and \( \omega_{OD} \).

In Chapter 4 we present an investigation into the vibrational relaxation pathway in isotopically dilute water. Ideally vibrational relaxation would be studied with pulses broad enough to span the entire mid-infrared with the time-resolution to capture even the initial energy transfer. Before this experiment is realized we study the proposed relaxation pathway by comparing the vibrational lifetime between isotopes and comparing the temperature-dependent lifetime to temperature-dependent shifts in frequency of the linear absorption spectrum. We find evidence that the fundamental of the bend is the dominant accepting mode for \( \omega_{OH} \) and \( \omega_{OT} \). However, for \( \omega_{OD} \) the temperature dependence of the lifetime indicates that there may be additional pathways in competition that have different dependencies on temperature.

In Chapter 5 we use temperature-dependent ultrafast infrared spectroscopy of dilute HOD in H₂O to study the picosecond reorganization of the hydrogen bond network of liquid water. Temperature-dependent 2D IR, pump-probe, and linear absorption measurements are self-consistently modeled with a response function formal-
ism that includes the effects of spectral diffusion, population lifetime, reorientational motion, and non-equilibrium heating of the local environment upon vibrational relaxation. Over the range of 278 to 345 K, we find the time scales of spectral diffusion and reorientational relaxation decrease from approximately 2.4 to 0.7 ps and 4.6 to 1.2 ps, respectively, which corresponds to the barrier heights of 3.4 and 3.7 kcal/mol, respectively. We compare the temperature dependence of the time scales to a number of measures of structural relaxation and find similar effective activation barrier heights and slightly non-Arrhenius behavior, which suggests that the reaction coordinate for the hydrogen bond rearrangement in water is collective. Frequency and orientational correlation functions calculated with molecular dynamics (MD) simulations over the same temperature range support our interpretations.

In Chapter 5 we also present dispersed pump-probe and 2D IR anisotropy measurements of the OD stretch of HOD in H₂O to test proposed molecular jump mechanism of hydrogen bond switching. The proposed mechanism predicts that water molecules initially excited in a weak or broken hydrogen bond will rapidly rotate in order to form a hydrogen bond. Rotational dynamics of water molecules initially excited in different hydrogen bond configurations can be resolved by spectrally dispersing the signal. Preliminary measurements show that molecules initially excited on the blue side of the line shape display faster reorientational motion. Our predictions from previous experimental results and the implications of the proposed mechanism also suggest that the long time anisotropy decay will be uniform across the absorption band since broken hydrogen bonds do not persist, or, water molecules originally tagged on the blue side of the line shape will have statistically sampled the entire absorption band within the first couple hundred femtoseconds. Preliminary measurements show a similar long time decay across the band. Finally, in the 2D measurement we find the anisotropy along the diagonal on the blue side of the line shape and in the off-diagonal region decays faster than along the diagonal on the red-side. This observation indicates that water molecules must reorient in order to switch hydrogen bond partners.
Chapter 2

Experimental Methods

2.1 Introduction

We use ultrafast infrared spectroscopy to study the intermolecular motions and vibrational relaxation in water. In this chapter we present our method for generating and characterizing ultrafast infrared pulses. We first generate and amplify 800 nm with a Ti:sapphire oscillator and regenerative amplifier. We create near-infrared pulses with optical parametric amplification with which we create mid-infrared pulses with difference frequency generation. We describe our method for acquiring third-order measurements in the pump-probe geometry and the pulse diagnostics that are acquired prior to and concurrently with data acquisition. Finally, we present our method for data processing, which is unique to the pump-probe geometry.

2.2 Generation of Ultrafast Mid-Infrared Pulses

2.2.1 800 nm Pulse Generation

We generate sub 100 fs mid-infrared (IR) pulses in a home-built optical parametric amplifier (OPA) pumped with 800 nm at 1 kHz (1.05 mJ, 40 fs). A schematic of the set-up is shown in Figure 2-1. A home-built Ti:Sapphire oscillator pumped with a Spectra-Physics Millenia Nd:YLF laser generates a pulse at 86 MHz with ~40 nm of
bandwidth (FWHM) and a center wavelength of ~790 nm. An example spectrum of
the oscillator output is shown in Figure 2-2b. After passing through a KMLabs ultra-
fast pulse stretcher, the pulse is amplified in a home-built Ti:Sapphire regenerative
amplifier [122] pumped by a Spectra-Physics Empower Nd:YAG laser. The pulse is
then compressed in a KMLabs ultrafast compressor before entering the OPA. Both
the stretcher and compressor consist of a grating pair. Figure 2-2 shows a frequency
and time resolved spectrum of the output pulse from the amplifier recorded by sec-
ond harmonic generation (SHG) frequency resolved optical gating (FROG) with the
commercial system, Grenouille [123]. The oscillator spectrum is peaked at 794 nm
with a pulse width 43.8 nm at full width half maximum (FWHM). The Grenouille
measures a pulse centered at 800 nm with 25.7 nm FWHM. The pulse has negligible
third order dispersion. The FWHM in time is 54.44 fs, which gives a pulse length of
38.88 fs.

Figure 2-1: Schematic of the oscillator, stretcher, amplifier, compressor, and OPA
lay-out (not drawn to scale). The filled black circles indicate periscopes.

2.2.2 Mid-Infrared Generation

From the 800 nm we generate ultrafast mid-infrared pulses with non-linear opti-
cal transformations. From Maxwell’s equations and the constitutive relations for a
nonmagnetic material we can derive the wave equation for nonlinear polarization
\[(a) \text{SHG FROG measured with Grenouille. (b) Spectrum of 800 nm after the oscillator (blue), amplifier (green), and measured with the Grenouille (red).}\]

(assuming \(\nabla \cdot \mathbf{E} = 0\))

\[\left(\nabla^2 - \mu_0 \sigma \frac{\partial}{\partial t} - \mu_0 \varepsilon \frac{\partial^2}{\partial t^2}\right)E = \mu_0 \frac{\partial^2}{\partial t^2} P^{NL}. \quad (2.1)\]

Inserting the dispersion relation \(k^2 = \mu_0 \varepsilon \omega^2\), and applying the slowing varying envelope approximation yields

\[\frac{\partial E}{\partial z} + \alpha E + \frac{n}{c} \frac{\partial E}{\partial t} = \frac{-i \omega \mu_0 c}{2n} (e \cdot p) P^{NL} e^{i(k-k_0)z}, \text{ where } \alpha = \frac{\mu_0 \sigma c}{2n}. \quad (2.2)\]

In a lossless material and within the steady state approximation the second two terms on the left hand side disappear. If we consider second harmonic generation and assume the pumping field, \(E(\omega_1)\), is undepleted we can directly integrate the equation from \(z = 0\) to \(L\). With the boundary condition \(E(\omega_2, z = 0) = 0\) this integration yields the intensity

\[I(\omega_2, L) = \frac{2 \omega_2^2 d_{ref}^2}{n_0^3 c^3 \varepsilon_0} L^2 I(\omega_1)^2 \left[\frac{\sin \Delta k L/2}{\Delta k L/2}\right]^2, \text{ where } \Delta k = k(\omega_2) - 2k(\omega_1), \quad (2.3)\]

where \(d_{ref}\) is a scalar calculated from the nonlinear coefficients, \(d_{il}\). Notice that assuming the pump field is undepleted results in an intensity that will increase without bound with the length of the crystal. \(\Delta k\) is referred to as the phase mismatch and
efficient nonlinear processes require phase matching, or $\Delta k = 0$.

In optical parametric generation (OPG) the idler and signal are created from quantum noise or parametric fluorescence. Here one intense pump wave at $\omega_3$ generates two waves at $\omega_1$ and $\omega_2$ (idler and signal, respectively) with the convention $\omega_3 > \omega_2 > \omega_1$. If, in addition to $\omega_3$, a weak seed pulse at $\omega_1$ or $\omega_2$ is present the process becomes optical parametric amplification. Compared to optical parametric amplification, OPG needs high pump intensities that can only be achieved with ultrashort laser pulses and/or by using a resonator for multiple amplification. The last possibility is termed optical parametric oscillation (OPO). In our application ultrafast mid-infrared pulses are created with optical parametric amplification followed by difference frequency generation (DFG), where two waves at frequencies $\omega_3$ and $\omega_2$ generate a field at $\omega_1$, or $\omega_1 = \omega_3 - \omega_2$.

The two coupled wave equations that describe OPG (assuming $E(\omega_3)$ is undepleted) are

\begin{align}
\left( \frac{\partial}{\partial z} + \alpha \right) E_1 &= -i \frac{\omega_1 d_{\text{eff}}}{n_1 c} E_3 E_2^* e^{-i \Delta k z} \\
\left( \frac{\partial}{\partial z} + \alpha \right) E_2 &= -i \frac{\omega_1 d_{\text{eff}}}{n_2 c} E_3 E_1^* e^{-i \Delta k z}
\end{align}

(2.4) (2.5)

where $\Delta k = k_3 - k_2 - k_1$ and $\alpha_1$ and $\alpha_2$ have been set to $\alpha$. Solving the equations yields the eigenvalues

\begin{align}
\gamma'_{\pm} &= -\alpha \pm \sqrt{\gamma^2 - \left( \frac{\Delta k}{2} \right)^2} \\
\gamma^2 &= \frac{\omega_1 \omega_2 d_{\text{eff}}^2 |E_3|^2}{n_1 n_2 c^2 I_3} = \frac{2 \omega_1 \omega_2 d_{\text{eff}}^2 I_3}{n_1 n_2 n_3 \epsilon_0 c^3 I_3}.
\end{align}

(2.6)

$\gamma$ is known as parametric gain and the equation written as $\gamma'_{\pm} = -\alpha \pm g$ introduces the variable $g$, defined as total gain. To have effective energy exchange between the pump and signal pulses, the group velocity mismatch (GVM) must be small to avoid temporal walk-off. This condition sets a lower limit on the duration of the pump.
pulse \cite{124},

\[ \tau_3 \geq L \left| \frac{1}{\nu_2} - \frac{1}{\nu_3} \right| \] \hspace{1cm} (2.7)

where \( L \) is the length of the crystal. \( \nu_2 \) and \( \nu_3 \) are the group velocities of the signal (or idler) and pump, respectively. To obtain the shortest pulse possible the gain bandwidth must be as large as possible. An approximation of the parametric gain bandwidth for an OPG/OPA is given by \cite{124},

\[ \Delta \nu = \frac{0.53}{c \left[ 1/\nu_2 - 1/\nu_3 \right]} \sqrt{\frac{\gamma}{L}} \] \hspace{1cm} (2.8)

where \( c \) is the speed of light. This expression is derived for when \( \omega_3 \) is narrow band, which is not valid for femtosecond pulses. Generally a numerical treatment is necessary to take into consideration depletion, saturation effects, and GVM \cite{125}.

**β-Barium Borate (BBO)** is a common crystal used in the generation of near-IR pulses due to its high damage threshold, high non-linear coefficients, and commercial availability, amongst other advantages. One drawback is the transparency range, shown in Figure 2-3, as wavelengths above 2.6 \( \mu \)m cannot be generated with high efficiency for typical lengths of a few millimeters. An important requirement is for the crystal to have a UV band edge that prevents two photon absorption at high pump intensities of 800 nm. BBO satisfies this requirement since it is transparent at 400 nm. The angular dependence of the indices of refraction, \( n_o \) and \( n_e \), are described by the Sellmeier equations. For BBO they are \cite{126}

\[ n_o^2 = 2.7359 + \frac{0.01878}{\lambda^2 - 0.01822} - 0.01354\lambda^2 \] \hspace{1cm} (2.9)

\[ n_e^2 = 2.3753 + \frac{0.01224}{\lambda^2 - 0.01667} - 0.01516\lambda^2 \] \hspace{1cm} (2.10)

where \( \lambda \) is in microns. Since \( n_o > n_e \), BBO is a negative uniaxial crystal. \( \omega_3 \) must be extraordinary to achieve phasematching. The two possibilities for phasematching are then type-I, where both \( \omega_1 \) and \( \omega_2 \) are ordinary, and type-II, where \( \omega_1 \) (or \( \omega_2 \)) is extraordinary and the other is ordinary. Type-II phasematching has the advantage
that the signal and idler can be separated after the first pass. As can be seen in equation 2.6, the parametric gain is dependent on \( d_{\text{eff}} \). For BBO,

\[
d_{\text{eff},I} = d_{31} \sin \theta - d_{22} \cos(\theta) \sin(3\phi) \tag{2.11}
\]

\[
d_{\text{eff},II} = d_{22} \cos^2(\theta) \cos(3\phi) \tag{2.12}
\]

where \( d_{22} = 2.2 \) and \( d_{31} = 0.08 \) pm/V [126]. The bandwidth (fs/mm) is estimated from the group velocity mismatch between the pump and signal (idler) and calculated with the equation,

\[
\frac{t_p}{L} = 2\left(\frac{1}{\nu_g(\omega_3)} - \frac{1}{\nu_g(\omega_2(1))}\right). \tag{2.13}
\]

The group velocity is calculated by,

\[
\frac{1}{\nu_g} = \frac{n}{c} \left(1 - \frac{\lambda \, dn}{n \, d\lambda}\right). \tag{2.14}
\]

Table 2-1 gives the signal and idler wavelengths, phasematching angle, \(|d_{\text{eff}}|\), gain, and bandwidth (BW) for a 3 mm crystal estimated from the GVM for mid-infrared pulses generated at 4, 5, and 6 \( \mu \)m. The calculation is performed for type-II phase-matching in which the \( \omega_1 \) is extraordinary and \( \omega_2 \) is ordinary. The parametric gain, in \( \text{cm}^{-1} \), can be obtained from the gain given in the table by multiplying by the square root of the pump intensity (in W/cm\(^2\)).

<table>
<thead>
<tr>
<th>Target mid-IR ( \lambda ):</th>
<th>4 ( \mu )m</th>
<th>5 ( \mu )m</th>
<th>6 ( \mu )m</th>
</tr>
</thead>
<tbody>
<tr>
<td>signal [( \mu )m]</td>
<td>1.33</td>
<td>1.38</td>
<td>1.42</td>
</tr>
<tr>
<td>idler [( \mu )m]</td>
<td>2.00</td>
<td>1.90</td>
<td>1.85</td>
</tr>
<tr>
<td>( \theta ) [degrees]</td>
<td>26.2</td>
<td>26.6</td>
<td>26.9</td>
</tr>
<tr>
<td>( d_{\text{eff}} ) [pm/V]</td>
<td>1.57</td>
<td>1.55</td>
<td>1.54</td>
</tr>
<tr>
<td>( \gamma' ) [W(^{-1/2})]</td>
<td>7.90x10(^{-5})</td>
<td>7.89x10(^{-5})</td>
<td>7.87x10(^{-5})</td>
</tr>
<tr>
<td>BW [cm(^{-1})], ( \omega_2 )</td>
<td>281</td>
<td>270</td>
<td>260</td>
</tr>
<tr>
<td>BW [cm(^{-1})], ( \omega_1 )</td>
<td>409</td>
<td>350</td>
<td>326</td>
</tr>
</tbody>
</table>

Table 2.1: Parameters for optical parametric amplification with a 3 mm BBO crystal

Similar coupled differential equations for wave propagation as those expressed for
optical parametric amplification can be written for DFG. The phase mismatch is $\Delta k = k_3 - k_1 - k_2$ and the solution assuming $\omega_3$ is undepleted is:

$$I_2(L) = I_2(0)cosh^2\gamma z$$
$$I_1(L) = \frac{\omega_1}{\omega_2} I_2(0)sinh^2\gamma z$$

where $\gamma$ is the gain defined above.

A common crystal for the generation of mid-infrared light is AgGaS$_2$ (AGS) through difference frequency generation (DFG) [127, 128, 129, 130]. The transmission curve for AgGaS$_2$ is shown in Figure 2-3. The Sellmeier equations for AgGaS$_2$ are:

$$n_o^2 = 5.7975 + \frac{0.2311}{\lambda^2 - 0.0688} - 0.00257\lambda^2$$
$$n_e^2 = 5.5436 + \frac{0.2230}{\lambda^2 - 0.0946} - 0.00261\lambda^2.$$

AgGaS$_2$ is also a negative uniaxial material and setting $\omega_1$ to the mid-infrared wavelength allows for three types of phasematching. $\omega_3$ is extraordinary in all three types. In Type-I phasematching $\omega_1$ and $\omega_2$ are ordinary and in Type IIa(b) $\omega_1$ is extraordinary (ordinary) and $\omega_2$ is ordinary (extraordinary). The effective nonlinear coefficients
are:

\[ d_{eff,I} = d_{36} \sin(\theta) \sin(2\phi) \quad (2.19) \]

\[ d_{eff,II} = d_{36} \sin(2\theta) \sin(2\phi). \quad (2.20) \]

where \( d_{36} = 13 \text{ pm/V} \) [131]. The phasematching angle, \( d_{eff} \), gain, and bandwidth at 4, 5, and 6 \( \mu \text{m} \) for type-II phasematching of a 0.5-mm AgGaS\(_2\) are given in Table 2-2.

<table>
<thead>
<tr>
<th>Target mid-IR ( \lambda )</th>
<th>4 ( \mu \text{m} )</th>
<th>5 ( \mu \text{m} )</th>
<th>6 ( \mu \text{m} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \theta ) [degrees]</td>
<td>51.8</td>
<td>45.5</td>
<td>41.8</td>
</tr>
<tr>
<td>( d_{eff} ) [pm/V]</td>
<td>14.8</td>
<td>15.3</td>
<td>15.3</td>
</tr>
<tr>
<td>( \gamma' ) [W(^{-1/2})]</td>
<td>2.43( \times )10(^{-4})</td>
<td>2.30( \times )10(^{-4})</td>
<td>2.13( \times )10(^{-4})</td>
</tr>
<tr>
<td>BW [cm(^{-1})], ( \omega_1 )</td>
<td>287</td>
<td>324</td>
<td>373</td>
</tr>
</tbody>
</table>

Table 2.2: Parameters for difference frequency generation with a 0.5 mm AgGaS\(_2\).

We generate mid-infrared light with white light seeded two stage optical parametric amplification followed by difference frequency generation (DFG). A schematic of the OPA is shown in Figure 2-4. A beamsplitter transmits 1% of the 800 nm to create a white light continuum in a sapphire plate, which mixes with a second portion of the 800 nm (9%) in a 3 mm Type-II BBO crystal (\( \theta=26^\circ, \phi=0^\circ \)) to create a higher frequency (signal) and lower frequency (idler) pulse. The two 800 nm beamsplitters are shown in green in Figure 2-4. After the first pass, a dichroic reflects the signal into a beam dump and transmits the idler, which passes back through the BBO with the third portion of 800 nm (90%). The dichroics, which are shown in cyan, are highly reflective at 1.2 to 1.46 \( \mu \text{m} \) and highly transmissive at 1.64 to 2.1 \( \mu \text{m} \). The idler is the seed in the second pass to create higher power signal and idler. Finally, the mid-IR is created through difference frequency generation of the signal and idler in a 0.5 mm AgGaS\(_2\) crystal. The OPA is tunable between \( \lambda_{max} \approx 4-7 \) microns (see Figure 2-5). Notice the absorption lines in the red-shifted spectrum from the absorption by water vapor and the relatively narrow CO\(_2\) absorption at \(~2370 \text{ cm}^{-1}\) in the green and blue spectra. The water vapor lines and CO\(_2\) absorption are marked at the top of the figure.
The spectra of the signal and idler used for generating 4 μm pulses are shown in Figure 2-6. The spectrum is taken after the mid-IR/signal/idler pass through 1 mm of Ge rotated to Brewster's angle used for HeNe overlap. Although Ge greatly decreases the amount of signal and idler, they are still present (~7% and ~23% of the total light, respectively). Ideally, if dispersion can be matched, one would use a thicker piece or additional piece of Ge (or Si) to further decrease the amount of signal and idler (see the section on dispersion). The wavelength axis is from 20 to 1.67 μm since the detector sensitivity is peaked at 8 μm and is therefore not sensitive at 1.33 μm (the wavelength of the signal). Although only the 2 μm idler peak is expected within this wavelength range, there is intensity at multiple wavelengths for both the signal and idler. We placed a piece of anti-reflection (AR) coated silicon before the monochromator to see if the extra peaks are real or from higher order diffraction off...
Figure 2-5: Mid-IR pulses generated with the OPA by tuning the BBO and AGS crystals.

Figure 2-6: Spectra of the signal (blue) and idler (red) used for generating 4 μm pulses with (right) and without (left) an AR-coated piece of Si (dashed) before the monochromator.

the grating. The transmittance of the Si is shown as a gray dashed line on the right-hand side of the figure. The peaks in the idler spectrum at 4 and 6 μm disappear with the Si showing that the peaks are not real. The peak at 6 μm in the signal spectrum disappears with the Si but the peaks at 8 μm and 4 μm remain, showing that another non-linear process is occurring to create these wavelengths. However,
the 8 \(\mu m\) peak may be second order diffraction from the 4 \(\mu m\) peak.

2.3 Pump-probe interferometer

2.3.1 General Set-Up

The interferometer is designed to acquire non-linear measurements in the pump-probe geometry [132]. We use two beamsplitters to split the mid-infrared into three pulses. A third beamsplitter recombines the collinear pulse pair that creates the population state. The third pulse acts as the probe and local oscillator. The interferometer set-up must also allow for pulse diagnostics to be quickly and easily acquired. It is advantageous to make the interferometer as small as possible so that fluctuations from air currents, temperature gradients, etc. are minimized. Finally, second order dispersion, which will be discussed in the next section, must be matched in order for pulses to be transform limited.

The layout of the interferometer is shown in Figure 2-7. The grey dashed line outlines the plexiglas and aluminum box and the black line outlines the 2'x2' breadboard upon which the majority of the interferometer is built. We use beamsplitters to split the infrared into three parts to acquire data in the pump-probe geometry in which a collinear pulse pair creates the population state and the third pulse acts as the probe and local oscillator. The beamsplitters, which consist of a thin layer of Ge on KBr substrate, are polarization sensitive. If the infrared is p(s)-polarized, \(~70\%/30\%\) (\(~45\%/55\%\)) is transmitted/reflected. We chose to split the beams \(~50/50\) at each beamsplitter so we use a turning periscope between the HeNe overlap and interferometer to rotate the polarization. The first beamsplitter transmits the probe beam (cyan, \(~45\%) and reflects the beam that is split into the two pump beams (green and blue), which are recombined prior to the sample (\(~14\% each). Since the beamsplitters absorb an appreciable amount of the HeNe, the interferometer was designed so that each of the three beams are transmitted once through a beamsplitter. With the retroreflectors, the probe beam is lowered by 0.5” towards the table and the two pump
beams are raised by 0.5" so that the beams are displaced vertically by 1". In order to avoid added dispersion, we use reflective optics (parabolics) to focus the beams into the sample position and re-collimate after the sample. Given that the effective focal length of the parabolics is 3", the crossing angle of the pump and probe at the sample is ~19°. The signal is spectrally dispersed after the sample with either a 40, 75, or 100-mm/groove grating to a 64-channel N2 cooled MCT detector. In the pump-probe geometry, a transient absorption measurement can be acquired simply by blocking the non-chopped pump beam (top trace in Figure 2-8).

Pulse delays are controlled with a pair of AR-coated ZnSe wedges mounted to mechanical stages (Newport). The stages control the pulse delay by changing the amount of material the beam transverses by translating the wedge face perpendicular to the beam path [133]. Each wedge pair is aligned antiparallel so there is negligible beam displacement. The placement of the wedge pair is shown in the upper left hand corner of Figure 2-7. The wedges are designed to have 0.1 fs delay per μm travel and a total optical path length delay of 5000 fs, which requires a 5 cm wedge. However, since the beams are not infinitely small, an extra 1 cm in length is added. The required wedge angle can be calculated from

$$\theta = \arctan\left(\frac{R \times c}{\Delta n}\right),$$

(2.21)

where R is the resolution (0.1 fs/μm), c is the speed of light, and $\Delta n = n_{ZnSe} - n_{air}$. The refractive index of ZnSe varies from 2.4339 to 2.4201 from 3.8 to 7.4 μm. The ZnSe wedges are 2 cm in height, 6 cm in length and have a 1.2° wedge. Data can be acquired either by step scanning, which implies there is an input equilibration time for each time point, or by fast scanning the wedges at constant velocity. The velocity is determined by:

$$V = \frac{\Delta t \times f}{R \times N_s},$$

(2.22)

where $\Delta t$ is the input time step, R is the resolution (time/length), f is the repetition rate of the laser (1/time), and $N_s$ is the number of shots to average for each time
point. The total time for a scan is then the total distance (determined from the total time window and resolution) divided by the velocity.
Figure 2-7: Interferometer.
2.3.2 Pulse Diagnostics

As mentioned in the previous section, the design of the beamsplitter set-up allows for pulse diagnostics to be performed relatively easily. The red line in Figure 2-7 traces the path the infrared and HeNe travel for HeNe overlap. After sending the beam into the interferometer we want to ensure the beams are tightly focused and overlapped at the sample position. We check the overlap and beam diameter with a pinhole at the sample position by focusing the beams into a room temperature MCT with a second parabolic after the sample (see Figure 2-8). In this configuration we also acquire autocorrelations either by second harmonic generation with a AgGaS₂ crystal or by collecting scatter though a small aperture pinhole [134]. The later is preferred since the interferometric measurement is not sensitive to the polarization of the pulses. Also, assuming the stages are calibrated, the array can be calibrated simply by removing the second parabolic to allow the scatter to enter the array. The replica of the pump pair created off the back side of the recombining beamsplitter is directed into the single channel monochromator. A pulse spectrum can be acquired simply by blocking one of the two pump beams and stepping the monochromator in frequency.

In the pump-probe geometry, the replica of the pump pair can be used to perform diagnostics simultaneously with data collection (in real time). We calibrate the stage scanned to create $\tau_1$ by directing the replica of the pulse pair through a monochromator set to a frequency near the peak of the spectrum. An example of the stage calibration trace collected during data collection is shown in Figure 2-8. We assume the frequency the monochromator is correct in order to correct the $\tau_1$-axis by Fourier transform of the time trace. Finally, a portion of the pump pair replica can be directed into a room temperature MCT detector to collect the real-time field autocorrelation. An example time trace is shown in Figure 2-8. With real time acquisition of the zero timing between the two pump beams, collecting third order measurements in the pump-probe geometry is, in principle, free of any pulse timing ambiguity during $\tau_1$ and $\tau_3$. 


2.3.3 Pump-Probe Contribution from the Pump Beams

In the pump-probe geometry, with the signal dispersed, the total signal can be written as [135]:

\[
S(\omega_3, \tau_1, \tau_2) \propto |E_3(\omega_3) + i\omega_3[P^{(1)}(\omega_3) + P^{(3)}(\omega_3, \tau_2, \tau_1) \\
+ P_{ppa}^{(3)}(\omega_3, \tau_2, \tau_1) + P_{ppb}^{(3)}(\omega_3, \tau_2, \tau_1)]|^2
\]  
(2.23)

In this equation, \(E_3\) is the probe field, \(P^{(1)}\) is the free induction decay from the third pulse, \(P^{(3)}\) is the desired third order response that requires the interaction of all three pulses, and \(P_{ppa/b}^{(3)}\) are the PP signals between each pump beam and the probe. The a/b subscripts denote the pump beams a and b marked in Figure 2-7. In the current set-up pump beam b is chopped. In Figure 2-9 we show the order of the pulses impinging on the sample with and without the chopper closed when pump b is the stationary beam and pump a is scanned. The total signal with the chopper open
is:

$$S^o(\omega_3, \tau_1, \tau_2) \propto |E_3(\omega_3) + i\omega_3[P^{(1)}(\omega_3) + P^{(3)}(\omega_3, \tau_2, \tau_1) + P_{ppa}(\omega_3, \tau_2) + P_{ppb}(\omega_3, \tau_2)]|^2$$

(2.24)

and with the chopper closed it is:

$$S^*(\omega_3, \tau_1, \tau_2) \propto |E_3(\omega_3) + i\omega_3[P^{(1)}(\omega_3) + P^{(3)}(\omega_3, \tau_2, \tau_1)]|^2.$$  

(2.25)

If we make the safe assumption that $|E_3| \gg |P|$ we find:

$$S^o - S^* \approx i\omega_3E_3(\omega_3)[P^{(3)}(\omega_3, \tau_1, \tau_2) + P^{(3)}_{ppb}(\omega_3, \tau_2)]$$

(2.26)

Figure 2-9: Pulse ordering with the chopper open and closed.

In Figure 2-10 we compare scanning the unchopped pump beam (stage a) and the chopped pump beam (stage b) at three wavelengths with the sample HOD in H2O at $\tau_2=120$ fs. The PP between pump pulse b and the probe is constant with $\tau_1$ when pump pulse a is scanned. The PP signal corresponds to a PP acquired at the waiting time: $P_{ppb}(\omega_3, \tau_2=120$ fs). However, if pump pulse b is scanned the pump-probe signal depends on $\tau_1$: $P_{ppa}(\omega_3, \tau_2=120$ fs+$\tau_1$). Therefore, if only one pump pulse is chopped,
it is better to scan the non-chopped beam since the PP appears as a constant off-set and will be removed by Fourier transform to create the excitation, or $\omega_1$, dimension. The frequency slice in Figure 2-10a/c corresponds to a bleach/stimulated emission in the PP signal that is positive/negative at early time and decays with increased waiting time. The frequency slice in Figure 2-10d has very low PP signal and shows complicated behavior when pump $b$ is scanned. Finally, in Figure 2-10b we compare the difference between the red and cyan trace in Figure 2-10a when stage calibration has (blue) and has not (green) been performed for each of the stages. Although both stages are off only by 1 fs over 200 fs (in the opposite manner, so 2 fs total) this can still be quite important for interferometric measurements.

![Figure 2-10](image)

Figure 2-10: (a,c,d) Comparison of scanning the chopped (pump b) vs. non-chopped (pump a) beam. (b) Difference between the traces obtained by scanning each of the pumps with stage calibration (blue) and without stage calibration (green).
2.3.4 Second-Order Dispersion, or GVD

In \( \text{H}_2\text{O} \), the broad libration, or hindered rotation, band is peaked at 680 cm\(^{-1} \), which corresponds to a period of 50 fs. To follow the dynamics, we desire a probe that is sensitive to structure with a "shutter speed" faster than the fastest intermolecular motions. Ideally we will generate transform limited pulses, or, pulses without second (or higher order) dispersion. The spectral phase of the electric field can be expanded in a Taylor expansion about a center frequency \([136]\):

\[
\phi(\omega) = \phi(\omega_0) + \phi'(\omega_0)(\omega - \omega_0) + \frac{1}{2} \phi''(\omega_0)(\omega - \omega_0)^2 + \frac{1}{6} \phi'''(\omega - \omega_0)^3 + ... \quad (2.27)
\]

where \( \phi' \), \( \phi'' \), \( \phi''' \) are derivatives of the phase with respect to the frequency and are known as the group delay, second-order dispersion (or group velocity dispersion (GVD)), and third-order dispersion (TOD), respectively. The group delay is defined as \( T(\omega) = \partial \phi(\omega)/\partial \omega \), and the Taylor expansion gives:

\[
T(\omega) = \phi'(\omega_0) + \phi''(\omega_0)(\omega - \omega_0) + \frac{1}{2} \phi'''(\omega_0)(\omega - \omega_0)^2 + ... \quad (2.28)
\]

Therefore when the GVD is non-zero the pulse will have a linear frequency chirp. Normal, or positive, dispersion occurs when higher frequency components travel slower than lower frequency components. The pulse is then positively chirped since frequency will increase with time. Anomalous, or negative dispersion, is the opposite situation. Demirdöven and coworkers demonstrated that the use of optical materials with GVD of opposite signs can successfully compress mid-IR pulses to 75 fs at 5 \( \mu \text{m} \) with 6 \( \mu \text{J} \) of energy and that the approach can be used throughout the mid-infrared until third-order dispersion becomes important \([130]\).

In Figure 2-11 we show the GVD per millimeter of material from 3.5 to 7 \( \mu \text{m} \) for the materials in the current interferometer. The GVD is calculated with the material dispersion equation \([136]\) and Sellmeier equations \([137]\). In Table 2-3 we list the thickness of the materials and the GVD at 4 and 6 \( \mu \text{m} \). Since the dispersion of all materials generally increases with decreasing wavelength, we replaced the 3 mm
Figure 2-11: GVD of the materials in the interferometer.

<table>
<thead>
<tr>
<th>Material</th>
<th>T (mm)</th>
<th>4 μm</th>
<th>6 μm</th>
</tr>
</thead>
<tbody>
<tr>
<td>DFG AgGaS₂</td>
<td>0.5</td>
<td>2</td>
<td>-19</td>
</tr>
<tr>
<td>HeNe Ge</td>
<td>4 (4), 3 (6)</td>
<td>1140</td>
<td>2181</td>
</tr>
<tr>
<td>lens KBr</td>
<td>2</td>
<td>-6</td>
<td>-136</td>
</tr>
<tr>
<td>bandpass Si</td>
<td>1</td>
<td>378</td>
<td></td>
</tr>
<tr>
<td>beamsplitter KBr</td>
<td>3</td>
<td>-10</td>
<td>-203</td>
</tr>
<tr>
<td>wedges ZnSe</td>
<td>4 to 6.4</td>
<td>300 to 479</td>
<td>-544 to -870</td>
</tr>
<tr>
<td>waveplate MgF₂</td>
<td>2.5</td>
<td>-1038</td>
<td></td>
</tr>
<tr>
<td>polarizer ZnSe</td>
<td>1</td>
<td>75</td>
<td>-136</td>
</tr>
<tr>
<td>window CaF₂</td>
<td>1</td>
<td>-280</td>
<td>-1110</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td></td>
<td>561 to 740</td>
<td>33 to -293</td>
</tr>
</tbody>
</table>

Table 2.3: Second order dispersion at 4 and 6 μm (in fs²).

piece of Ge used for HeNe overlap with a 1 mm piece upon tuning from 6 μm to 4 μm. The 1 mm piece of anti-reflection coated Si (spectrum shown on the right in Figure 2-6) is used at 4 μm to absorb signal and idler in addition to the Ge. Each of the three beams are transmitted only once through a KBr beamsplitter (3 mm thick). The thickness of a single wedge varies from 2 to 3.2 mm. Therefore, the total thickness of the wedge pair can range from 4 to 6.4 mm. At 6 μm we did not use waveplates. Rather, we rotated the polarization of the probe between parallel and perpendicular relative to the pump with a turning periscope. Finally, the 1 mm CaF₂
window refers to the front window of the sample cell. At 4 μm the total dispersion at the sample from all the material in the interferometer ranges from 561 to 740 fs$^2$. We found adding a 3 mm piece of CaF$_2$ decreased the chirp (3 mm = -840 fs$^2$). At 6 μm the dispersion at the sample ranges from 33 to -293 fs$^2$ and we found additional material was not necessary.

2.4 Data processing

The nonlinear experiments described in this thesis evaluate the interaction of matter with three infrared pulses. The third-order polarization, $P^{(3)}$, is expressed as a triple convolution of the response function $R$ with the incident fields, $E$:

$$ P^{(3)}(t) = \int_0^\infty dt_3 \int_0^\infty dt_2 \int_0^\infty dt_1 R(t_3, t_2, t_1)E(t - t_3) \times E(t - t_3 - t_2)E(t - t_3 - t_2 - t_1). $$  

(2.29)

In this equation $t_1$ and $t_2$ are the time periods between the field-matter interactions and $t_3$ is the time between the last interaction and the detection at time $t$. The incident fields have a defined envelope, polarization, wave vector, frequency, and phase. The frequency is near resonant with both the $\nu=1\leftarrow 0$ peak and anharmonically shifted $\nu=2\leftarrow 1$ peak of interest. The tensorial response function is a thrice-nested commutator of the transition dipole operator, $\mu$, evaluated at each field-matter interaction:

$$ R(t_3, t_2, t_1) = (i/\hbar)^3 [[[\mu(t_3 + t_2 + t_1), \mu(t_2 + t_1)], \mu(t_1)], \mu(0)]. $$  

(2.30)

The response function evolves under the material Hamiltonian and therefore it inherently contains microscopic information on the system of interest. Expanding the commutator results in eight separate terms, but only four are unique since the other four are complex conjugates. Until this point no simplifying assumptions have been applied.

The ensuing assumptions have previously proved useful in modeling third order spectroscopy. The response function is simplified by assuming the vibrational and
rotational degrees of freedom are separable and is then written as a product of the two contributions. The vibrational response function includes terms dependent on the transition dipole, eigenstate energy, and evolution under the Hamiltonian. To describe the evolution under the Hamiltonian, Sung and Silbey derived expressions for the nonlinear dephasing functions in terms of two-point energy gap correlation functions between the system eigenstates, which assumes the second cumulant approximation [138]. The cumulant approximation is exact if frequency fluctuations are Gaussian. The authors derived the theory for four wave mixing experiments of a multilevel system coupled to a harmonic bath. The energy gap correlation functions reflect the change in the system energy upon coupling to a bath mode. The orientational response function projects the polarization of the incident electric field onto the distribution of molecular coordinates and includes transition dipole unit vector and Legendre polynomials derived for a spherical rotor.

Double-sided Feynman diagrams pictorially relay the possible contributions to the signal. The pathways are termed Liouville pathways as the Liouville-von Neuman equation describes how the density matrix evolves in time under the quantum mechanical Hamiltonian. The initial pathways, which involve permutations of three time periods on the original eight Feynman diagrams, can be decreased by invoking physical arguments including, but not limited to, the rotating wave approximation and consideration of the phase matching direction. In nonlinear electronic spectroscopy four diagrams represent the possible contributions for the two level system. In nonlinear vibrational spectroscopy the multilevel structure must be taken into account. For water, a three level system, eight Feynman diagrams remain outside the regime of overlapping pulses. Four are rephasing, \( S_- \), in that the oscillation frequencies in the third time period are conjugate, or nearly conjugate, to those in the first time period. The ability to rephase relies on frequency correlation between the first and third time periods, and frequencies evolve during the second time period. The non-rephasing \( S_+ \) contributions involve frequencies that are the same, or nearly the same, in the first and third time periods. 2D IR measurements are commonly acquired in the box-car geometry. In this geometry the rephasing and non-rephasing signals, emitted
in the \( k_{\text{sig}} = -k_1 + k_2 + k_3 \) and \( k_{\text{sig}} = k_1 - k_2 + k_3 \) phasematched directions, respectively, are measured separately. After Fourier transform to the frequency domain, the spectra are summed to yield the purely absorptive line shape [139].

2.4.1 Extracting Complex Correlation, Rephasing and Non-Rephasing Spectra

In the pump-probe geometry \( k_1 \) and \( k_2 \) are indistinguishable and, therefore, the real part of the 2D IR rephasing \( (S_-) \) and non-rephasing \( (S_+) \) pathways are collected [140]. However, utilizing the Kramers-Kronig assumption we can extract the complex correlation, rephasing and non-rephasing 2D information. The data is collected as a function of \( \tau_1 \) and \( \omega_3 \), since a grating performs the Fourier transform when dispersing the signal onto a multi-channel array. The signal can be written as:

\[
S = [S_-(\tau_1, \omega_3, \phi_1 = 0, \phi_3 = 0) + S_+(\tau_1, \omega_3, \phi_1 = 0, \phi_3 = 0)]. \tag{2.31}
\]

where \( S_- \) and \( S_+ \) represent the rephasing and non-rephasing contributions to the nonlinear signal, respectively. A single sided, one-dimensional Fourier transform is performed in \( \tau_1 \) to create the \( \omega_1 \) axis and produce the 2D IR spectrum. The real \( (S') \) and imaginary \( (S'') \) components of this spectrum can be written as:

\[
S' = [S_-(\omega_1, \omega_3, \phi_1 = 0, \phi_3 = 0) + S_+(\omega_1, \omega_3, \phi_1 = 0, \phi_3 = 0)] \tag{2.32}
\]

\[
S'' = [S_-(\omega_1, \omega_3, \phi_1 = 90, \phi_3 = 0) + S_+(\omega_1, \omega_3, \phi_1 = 90, \phi_3 = 0)]. \tag{2.33}
\]

(1) \( S' \) and \( S'' \) are inverse Fourier transformed in the \( \omega_3 \) dimension, (2) causality is applied by setting the time-domain signal for \( \tau_3 < 0 \) to zero, (3) and the signal is
Fourier transformed with respect to $\tau_3$ to obtain complex 2D spectra:

$$\text{Re}(S') = [S_-(\omega_1, \omega_3, \phi_1 = 0, \phi_3 = 0) + S_+(\omega_1, \omega_3, \phi_1 = 0, \phi_3 = 0)]$$ (2.34)

$$\text{Im}(S') = [S_-(\omega_1, \omega_3, \phi_1 = 0, \phi_3 = 90) + S_+(\omega_1, \omega_3, \phi_1 = 0, \phi_3 = 90)]$$ (2.35)

$$\text{Re}(S'') = [S_-(\omega_1, \omega_3, \phi_1 = 90, \phi_3 = 0) + S_+(\omega_1, \omega_3, \phi_1 = 90, \phi_3 = 0)]$$ (2.36)

$$\text{Im}(S'') = [S_-(\omega_1, \omega_3, \phi_1 = 90, \phi_3 = 90) + S_+(\omega_1, \omega_3, \phi_1 = 90, \phi_3 = 90)]$$ (2.37)

Re($S'$) and Im($S'$) are shown in Figure 2-12a and 2-12b, respectively, for a 2D surface of the OD stretch of HOD in H$_2$O at $\tau_2 = 600$ fs at 286 K. At this point the complex correlation surface can be constructed from Re($S'$) and Im($S'$) [135]. The absolute value, $S_{\text{Abs}} = |\text{Re}(S') + i\text{Im}(S')|$, and phase representation, $S_{\phi} = \arctan(\text{Im}(S')/\text{Re}(S'))$, are shown in Figure 2-12c and 2-12d, respectively.

![Figure 2-12](image)

Figure 2-12: (a) Real, (b) Imaginary, (c) Absolute Value, and (d) Phase representations of a 2D surface of HOD in H$_2$O at 286 K and $\tau_2 = 600$ fs.

In order to separate the rephasing and non-rephasing pathways we will rewrite
Figure 2-13: (a) Re(S_−), (b) Im(S_−), (c) S_−, (d) Re(S_+), (e) Im(S_+), and (f) S_+ representations of a 2D surface of HOD in H_2O at 286 K and τ_2 = 600 fs.

The terms in the above equations as:

\[
\text{Re}(S') = \left[ S_-(\omega_1, \omega_3, \phi_1 = 0, \phi_3 = 0) + S_+(\omega_1, \omega_3, \phi_1 = 0, \phi_3 = 0) \right] \quad (2.38) \\
\text{Im}(S') = \left[ S_-(\omega_1, \omega_3, \phi_1 = 90, \phi_3 = 0) - S_+(\omega_1, \omega_3, \phi_1 = 90, \phi_3 = 0) \right] \quad (2.39) \\
\text{Re}(S'') = \left[ S_-(\omega_1, \omega_3, \phi_1 = 90, \phi_3 = 0) + S_+(\omega_1, \omega_3, \phi_1 = 90, \phi_3 = 0) \right] \quad (2.40) \\
\text{Im}(S'') = \left[ -S_-\left(\omega_1, \omega_3, \phi_1 = 0, \phi_3 = 0\right) + S_+\left(\omega_1, \omega_3, \phi_1 = 0, \phi_3 = 0\right) \right] \quad (2.41)
\]

given the following relationships:

\[
S_-(\omega_1, \omega_3, \phi_1 = 0, \phi_3 = 0) = -S_-(\omega_1, \omega_3, \phi_1 = 90, \phi_3 = 90) \quad (2.42) \\
S_-(\omega_1, \omega_3, \phi_1 = 90, \phi_3 = 0) = S_-\left(\omega_1, \omega_3, \phi_1 = 0, \phi_3 = 90\right) \quad (2.43) \\
S_+(\omega_1, \omega_3, \phi_1 = 0, \phi_3 = 0) = S_+\left(\omega_1, \omega_3, \phi_1 = 90, \phi_3 = 90\right) \quad (2.44) \\
S_+(\omega_1, \omega_3, \phi_1 = 90, \phi_3 = 0) = -S_+\left(\omega_1, \omega_3, \phi_1 = 0, \phi_3 = 90\right) \quad (2.45)
\]

Now we can separate the rephasing (S_-) and non-rephasing (S_+) pathways, for instance, by subtracting Im(S'') from Re(S') to obtain the real part of the rephasing.
pathway. Explicitly we have:

\[
\begin{align*}
\text{Re}(S') - \text{Im}(S'') & = S_-(\omega_1, \omega_3, \phi_1 = 0, \phi_3 = 0) = \text{Re}(S_-) \\
\text{Re}(S'') + \text{Im}(S') & = S_-(\omega_1, \omega_3, \phi_1 = 90, \phi_3 = 0) = \text{Im}(S_-) \\
\text{Re}(S') + \text{Im}(S'') & = S_+(\omega_1, \omega_3, \phi_1 = 0, \phi_3 = 0) = \text{Re}(S_+) \\
\text{Re}(S'') - \text{Im}(S') & = S_+(\omega_1, \omega_3, \phi_1 = 90, \phi_3 = 0) = \text{Im}(S_+) 
\end{align*}
\]

These terms are shown in Figure 2-13a, 2-13b, 2-13d, and 2-13e, respectively. Finally in Figure 2-13c and 2-13f we show the complex rephasing and non-rephasing surfaces. We normalize both surfaces by the maximum of the rephasing surface to show that the rephasing pathway still dominates at this waiting time and temperature indicating spectral diffusion is not yet complete [141, 72].

### 2.4.2 Cross Phase Modulation

In pump-probe spectroscopy, an intense pump pulse can modulate the refractive index of the solvent or sample window causing a time-dependent phase modulation of the probe. This is referred to as cross phase modulation (xpm) and, since the total energy of the probe remains constant, it can only be observed if the probe is spectrally dispersed after the sample. Kovalenko et al. experimentally and theoretically explored non-resonant contributions to femtosecond visible pump-supercontinuum-probe (PSCP) spectroscopy [142]. In their treatment, they define the pump pulse as a nonchirped Gaussian

\[
E_1(t) = a_1 \exp \left[ -t^2 / 2 \tau_1^2 + i \Omega_1 t \right] 
\]

where \(a_1\) is the amplitude, \(\Omega_1\) is the central frequency, and \(\tau_1\) is the pulse duration. They consider a probe pulse with a linear chirp, written as:

\[
E_2(t) = \exp \left( -t^2 / 2 \tau_2^2 + i (\Omega_2 t + \beta t^2) \right),
\]

\(\tau_2\)
where $\beta$ is the chirp rate. They find the differential instantaneous electronic signal can be written as:

$$
\Delta D_e(\omega_2, \tau_d) = -\frac{12 \omega_2 |a_1|^2 \sigma_e}{\sqrt{\eta}} \text{Im}\{\exp[-\frac{[t_d + t_o(\omega_2)]^2}{\eta \tau_1^2}] + \frac{t_o(\omega_2) t_d + t_o(\omega_2)}{\beta \tau_2^2 \eta} + \frac{(t_o(\omega_2))^2}{2 \beta \tau_2^2} \frac{1}{\tau_1^2 \eta} - i \phi/2]\}
$$

(2.52)

where $\sigma_e$ is the amplitude, $t_d$ is the delay between the pump and the probe pulse, $\eta = (1+2\xi^2/\alpha)$, $\alpha = 1-i2\beta \tau_2^2$, $\xi = \tau_2/\tau_1$, $\phi = \arctan(\text{Im} \eta / \text{Re} \eta)$, and $t_o(\omega_2) \approx (\omega_2-\Omega_2)/2\beta$. 

Figure 2-14: (a) Dispersed pump-probe for a pump-linearly chirped probe (left) and pump-no chirp probe (right). (b) The amplitude of the xpm at $t_p = 0$ fs. (c) Slices of the xpm in frequency at $\Omega_2-50$ cm$^{-1}$ (red), $\Omega_2$ (green), and $\Omega_2+50$ cm$^{-1}$ (blue).
When \(2\beta\tau_2^2 \gg 1\), one has \(|\eta| \approx 1\), \(\phi \approx 1/\beta\tau_1^2\), and the signal becomes

\[
\Delta D_e(\omega_2, \tau_d) \approx -\frac{12\omega_2}{|\eta|} \frac{a_1^2 \sigma_e}{\sqrt{\tau_1^2}} \exp[-\frac{(t_d + t_0(\omega_2))^2}{\tau_1^2}] \\
\times \sin\left[\frac{1}{2\beta\tau_1^2} \frac{[t_d + t_0(\omega_2)]^2}{\beta\tau_1^4} - \frac{t_0(\omega_2)[t_d + t_0(\omega_2)]}{\beta\tau_2^2 \tau_1^2}\right].
\]

(2.53)

The authors did not consider the case when both the pump and probe have chirp, which, if the probe pulse has chirp, would be the case in our experiments. For a non-chirped probe pulse, \(\beta = 0\), and the expression for the differential instantaneous electronic signal is:

\[
\Delta D_e(\omega_2, \tau_d) = -\frac{12\omega_2}{|\eta|} \frac{a_1^2 \sigma_e}{\sqrt{\tau_1^2}} \exp\left[\frac{\xi^4\tau_1^2(\omega_2 - \Omega_2)}{1 + 2\xi^2}\right] \\
\times \sin\left[\frac{2\xi^2}{1 + 2\xi^2} t_d(\omega_2 - \Omega_2)\right]
\]

(2.54)

In Figure 2-14 we show an example of cross phase modulation when the probe pulse is either linearly chirped (equation 2.53) or is not chirped (equation 2.54). In both cases \(\tau_1 = \tau_2 = 45\) fs and \(\Omega_2 = 2508\) cm\(^{-1}\). When the probe pulse is not chirped the xpm is sinusoidal about \(t_p = 0\) fs. Therefore the amplitude is zero at \(t_p = 0\) for all probe frequencies (Figure 2-14b). At the peak of the probe pulse the xpm is zero (Figure 2-14c) and flips sign on either side of \(\Omega_2\). When the probe pulse is linearly chirped (\(\beta = 1 \times 10^{-2}\) fs\(^{-2}\) in this example) the xpm is not zero at \(\Omega_2\). Rather there is positive amplitude peaked at \(t_p = 0\) fs and symmetric negative side lobes. As \(\omega_2\) is tuned away from \(\Omega_2\) the side lobes become asymmetric (Figure 2-14c) and the maximum shifts from \(t_p = 0\) fs (Figure 2-14b).

In Figure 2-15 we compare experimentally measured cross phase modulation with the model calculations. The experimental xpm is taken with H\(_2\)O at the sample position between two 1 mm thick CaF\(_2\) windows with a 6 \(\mu\)m spacer. Modeling the experimental data with equation 2.54 and the same input values as Figure 2-14, largely reproduces the observed xpm, although the asymmetry about \(t_p\) is not reproduced. When \(\omega_2 \approx \Omega_2\) and the xpm arising from an unchirped probe pulse is \(\sim 0\),
Figure 2-15: Experimental xpm (blue) compared to model calculations with linearly chirped probe (green) and unchirped probe (red).

the experimental trace resembles model xpm with a linearly chirped pulse. However, the width is not properly reproduced. \( \tau_1, \tau_2, \) and \( \Omega_2 \) are set by fitting the experimental xpm with equation 2.54 and not adjusted when applying equation 2.53. Also, adjusting \( \beta \) only affected the number of oscillations, not the width of the envelope. The above equations assume the sample is an infinitely thin, therefore propagation affects are neglected. Ekvall et al. [143] theoretically and experimentally studied the cross phase modulation in a piece of fused silica in visible pump-supercontinuum-probe spectroscopy. They considered propagation effects by including group velocity mismatch between the pump and probe and first and second order dispersion of the white light continuum probe and numerically solved the coupled wave propagation equations. The model calculations reproduce the asymmetry about \( t_p \) we observe experimentally (Figure 4 in ref. [143]), which they attribute to propagation effects.

As far as we are aware, the effect of cross phase modulation in 2D IR measurements has not been addressed experimentally or theoretically. To get an idea of how the xpm will appear in 2D measurements we multiplied the spectra in Figure 2-14a and 2-14b
by a sine function oscillating at $\Omega_2$ in $\tau_1$ then Fourier transformed. The resulting 2D spectra are shown in Figure 2-16. If there is linear chirp the 2D surface shows a doublet that is negative on the red side of $\Omega_2$ and positive on the blue side. If there is no chirp the 2D surface shows a clover shape with a node at $\omega_1 = \omega_3 = \Omega_2$. Cross-phase modulation will only be important when the pulses are overlapping.

2.5 Acknowledgments

We would like to thank Lauren DeFlores, Kevin Jones, and Matt DeCamp for their helpful insights and thoughtful discussions.
Chapter 3

Infrared Spectroscopy of Tritiated Water

3.1 Introduction

Studies of hydrogen bond dynamics of water with ultrafast non-linear infrared spectroscopy have commonly used isotopically dilute solutions such as HOD in D₂O [58, 62, 68, 67, 71] or HOD in H₂O [77, 120]. The anharmonic OH or OD stretching vibration is localized in these mixtures, and particularly sensitive to hydrogen bond configurations, as manifested by the fluctuating electric field experienced by the proton or deuteron [68, 46, 78]. However, with only one spectroscopically observable species, information on the environment is inferred indirectly. Using experiments on HOD in D₂O, we recently concluded that hydrogen bond switching involves a concerted motion of no less than three molecules: the hydrogen bond donor and the initial and final hydrogen bond acceptor molecules [73]. A more direct test of this process requires a spectroscopic probe of correlated motion between hydrogen-bonded molecules. This could potentially be observed using correlation spectroscopy between OD and OT stretches in H₂O, exploiting both the coupling between species and sensitivity of the vibrational frequency to hydrogen bond configuration. As a step in this direction, we report on the infrared absorption spectrum of ωOT of HOT in H₂O. Later in the chapter we present a calculation to model what will be observed
in such an experiment. In the calculation, the diagonal frequency modulation from the intermolecular electric field and off-diagonal transition dipole coupling (TDC) are determined during an idealized, concerted hydrogen bond switch. Finally, intermolecular coupling between the OD and OT stretch is determined by performing \textit{ab initio} frequency calculations on water isotopologue dimer combinations.

\section{3.2 Linear Absorption of Tritiated Water}

The high radioactivity of tritium has precluded previous liquid infrared studies, although a Raman spectrum of the OT stretch for 10\% T\textsubscript{2}O in H\textsubscript{2}O (\sim 60 Ci) has been reported [144]. Line shape parameters for the absorption spectrum of \(\omega_{\text{OT}}\) are reported, including the frequency at maximum absorbance (\(\omega^*\)), integrated intensity (\(I\)), molar absorptivity (\(\varepsilon\)), and full width half maximum (\(\Delta\omega\)). Comparison of the linear infrared absorption spectra of the three isotopes, \(\omega_{\text{OT}}:\omega_{\text{OD}}:\omega_{\text{OH}}\), provides information on the physical effects and interactions that influence the red-shift, broadening, and intensity gain of the OX stretch upon forming the hydrogen bond. Within the simplest harmonic approximations, the line shape parameters should remain constant for all isotopologes when scaled by reduced mass \(\mu_R\) as \(\omega^*\sqrt{\mu_R}, I\sqrt{\mu_R}, \varepsilon\mu_R,\) and \(\Delta\omega\sqrt{\mu_R}\). These scaling relationships are investigated in an attempt to test the generality of electric field models for the hydride stretching line shape and predictions of the importance of mechanical and electrical anharmonicity to the line shape.

\subsection{3.2.1 Experimental Methods}

Dilute tritiated water, H\textsubscript{3}T in H\textsubscript{2}O, was obtained from Sigma (W3260, CAS\# 13670-17-2) and used without further purification. The extent of labeling is 5 \(\pm 0.5\) Ci g\textsuperscript{-1} or 0.2 \(\pm 0.02\) M given the half-life 4500 \(\pm 8\) days for \(^3\text{H}\) [145]. FTIR spectra of tritiated water and reference deionized H\textsubscript{2}O were obtained with a spectrometer (Mattson Infinity Gold) whose beam was routed externally to tightly focus its beam to the sample position using 10 cm parabolic mirrors. This was done to avoid contamination of the spectrometer and decrease the amount of tritiated water used. The sample
cell consisted of 120 μm thick, 9 mm ID adhesive spacers (Secure Seal™ Imaging Spacers, Grace Biolabs) placed between 1 mm thick, 2" dia. CaF₂ windows. The spacer accommodated side-by-side sample and reference wells. Each well was filled by approximately 8 μL of sample (~40 mCl of ³H) and translation stages provided reproducible movement between the reference and sample.

FTIR spectra of HOD in H₂O and D₂O (Cambridge Isotope Laboratories, CAS# 7789-20-0) were collected with the same spectrometer. The sample cell consisted of 3 mm thick CaF₂ windows (Pike Technologies, Inc.) and a 50 μm teflon spacer. For both ωOH and ωOD a dilution series of solutions were measured to separate absorption contributions from the three species present (HOD, H₂O and D₂O). The solutions were kept dilute to avoid intermolecular interactions. Spectra were obtained by iteratively solving

\[ A = d(ε_{HOD}C_{HOD} + ε_{H₂O}C_{H₂O} + ε_{D₂O}C_{D₂O}), \]  

using the equilibrium constant 3.82 for the reaction H₂O + D₂O ⇌ 2HOD [146].

### 3.2.2 Results

Figure 3-1 shows the infrared absorption spectrum of the OT stretch in terms of the molar absorptivity, ε(ω) [10³ cm² mol⁻¹]. The absorption line shape has a peak at \( ω^* = 2121 \text{ cm}^{-1} \), and line width of \( Δω = 127 \text{ cm}^{-1} \) (FWHM), and a qualitatively similar shape to the other HOX isotopologues.

Comparison of the line shape parameters of ωOT, ωOD, and ωOH is made in Figure 3-2 and Table 3-1. The scaling dependence of the absorption coefficient with isotopic substitution is predicted from the expression that relates it to the Fourier transform of the dipole-dipole correlation function [147]

\[ ε(ω) = \frac{2π}{3nℏc}ω(1 - e^{−ℏω}) \int_{−∞}^{∞} dt e^{iωt} \langle \vec{μ}(t) \cdot \vec{μ}(0) \rangle. \]  

As a point of reference, for a harmonic oscillator, the transition frequency (ω =
\[ \sqrt{\frac{k}{\mu_R}} \] and square of the transition dipole are inversely proportional to the square root of the reduced mass. Given that the transition dipole operator is linear in the system coordinate, \( Q \), the mass weighting of the matrix element arises from \( Q = \frac{\hat{x}(h^2/\mu_R k)^{1/4}}{2\pi} \). Figure 3-2a shows the molar extinction coefficient for each mode using a reduced frequency axis \( \omega \sqrt{\mu_R} \) in which the reduced mass for the OX stretch is estimated from \( \mu_R = m_X m_O/(m_X + m_O) \). To within 1%, this form gives the same results as modified diatomic approximations [148, 54]. The reduced mass scaling does not lead to coincident peak maxima. To account for anharmonicity of the OX stretch present before hydrogen bonding, we compare the scaling to the case where the frequency is normalized relative to the gas phase OX frequencies, \( \omega_g \). In this case (Figure 3-2b), the normalized bands align much better and showing only slight deviations in peak maxima (OH:OD:OT = 0.9184:0.9210:0.9224).

Equation 3.2 can also be used to predict the line width scaling with reduced mass for interactions of a system with a harmonic bath. Using the Condon approximation and assuming Gaussian statistics, the dipole correlation function is often expressed in terms of the fluctuating energy gap arising from a system-bath interaction

\[
\langle \vec{\mu}(t) \cdot \vec{\mu}(0) \rangle = |\mu_0|^2 e^{-i\omega_0 t} \exp\left(-\frac{1}{\hbar^2} \int_0^t dt'' \int_0^{t''} dt' \langle H_{SB}(t'')H_{SB}(t') \rangle \right). \tag{3.3}
\]
Figure 3-2: (A) The stretching band of each of the three isotopologues of water ($\omega_{OH}$ for HOD in D$_2$O, $\omega_{OD}$ for HOD in H$_2$O, and $\omega_{OT}$ for HOT in H$_2$O) with the frequency axis scaled by the square root of the reduced mass of each local mode. (B) The stretching band with the frequency axis scaled by the gas phase frequencies and intensities normalized to compare $\Delta \omega$. (C) Peak intensities scaled by the reduced mass of each isotope to compare $\varepsilon$.

where $H_{SB}$ refers to the system-bath interaction Hamiltonian. The reduced mass scaling of the line width depends on the details of $H_{SB}$ and the dynamics. In the static case, if $H_{SB}$ is linear in the OX coordinate, an approximation that is valid for this system [46], the line width should scale inversely as $\sqrt{\mu_R}$ [69]. This observa-
tion predicts superimposed line shapes in Figure 3-2b. The correspondence is close; however, we find $\Delta \omega_{OH}$ is 17% larger than $\Delta \omega_{OD}$ on the scaled axis and 24% larger than $\Delta \omega_{OT}$. The preferential narrowing of the higher masses suggests that this may be a motional narrowing effect that results from the same spectral dynamics applied to different frequency distributions. The line shape of the OX stretch in isotopically dilute water only deviates slightly from Gaussian. The asymmetry, calculated by the variation of half-widths at half-height, $(\Delta \omega_{1/2,\text{blue}} - \Delta \omega_{1/2,\text{red}})/\Delta \omega$, is $\sim 2.5\%$ for the three isotopes.

<table>
<thead>
<tr>
<th></th>
<th>OH</th>
<th>OD</th>
<th>OT</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\mu_R$</td>
<td>0.9481</td>
<td>1.7889</td>
<td>2.5377</td>
</tr>
<tr>
<td>$\omega_g$ (cm$^{-1}$)</td>
<td>3707.47</td>
<td>2723.68</td>
<td>2299.77</td>
</tr>
<tr>
<td>$\omega_i$ (cm$^{-1}$)</td>
<td>3405±2</td>
<td>2509±2</td>
<td>2121±4</td>
</tr>
<tr>
<td>$I$ (cm$^2$mol$^{-1}$)</td>
<td>4922±183</td>
<td>3786±97</td>
<td>2321±238</td>
</tr>
<tr>
<td>$\Delta \omega$ (cm$^{-1}$)</td>
<td>254±4</td>
<td>160±4</td>
<td>127±8</td>
</tr>
<tr>
<td>$\varepsilon$ (10$^3$cm$^2$mol$^{-1}$)</td>
<td>60±2</td>
<td>52±1</td>
<td>33±3</td>
</tr>
</tbody>
</table>

Table 3.1: Comparison of the absorption line shape parameters for the three isotopologues of water ($\omega_{OH}$ for HOD in D$_2$O, $\omega_{OD}$ for HOD in H$_2$O, and $\omega_{OT}$ for HOT in H$_2$O)

The largest deviations from reduced mass scaling are observed for the molar absorptivity, $\varepsilon(\omega)$. Equation 3.2 predicts an inverse scaling of $\varepsilon$ with $\mu_R$, which is tested on the reduced axes in Figure 3-2c. For this parameter $\mu_OD\varepsilon_{OD}$ is larger than $\mu_{OH}\varepsilon_{OH}$ by 40 ± 4% and $\mu_{OT}\varepsilon_{OT}$ is larger than $\mu_{OH}\varepsilon_{OH}$ by 32 ± 13%. A reduced mass scaling of $1/\sqrt{\mu_R}$ leads to somewhat better, although still unsatisfying, agreement. In Table 3-1, we also report the integrated areas calculated by

$$I = \int_{\text{band}} \frac{\varepsilon(\omega)}{\omega} d\omega.$$  \hspace{1cm} (3.4)

The integration limits correspond to molar absorptivities at one tenth of the peak absorptivity to avoid interference of the overtone of the HOD bend centered at approximately 2950 cm$^{-1}$. Since $I$ is approximately a product of $\varepsilon$ and $\Delta \omega$, it will not be discussed further.
3.2.3 Discussion

A considerable amount of literature has been devoted to describing isotope substitution experiments for the OX stretch IR spectroscopy, and the failings of simple reduced mass scaling in the observables. Haas and Hornig found that the line width of $\omega_{OH}$ is four times greater than $\omega_{OH}$ in isotopically dilute ice and proposed proton tunneling to explain the observation [149]. Bournay and Marchal [150, 151, 152] invoked both anharmonic coupling to the hydrogen bond stretch and electrical anharmonicity to account for the anomalous intensity scaling in carboxylic acid dimers. With three isotopes, the present results provide unique opportunity to test the linearity of certain scaling relationships. Anharmonicity is seen to be clearly important, as judged by the inadequacy of harmonic reduced mass scaling relationships for $\omega^*$ and $\varepsilon$. The amplitude scaling offers no simple explanation; however, the $\omega^*$ scaling clearly originates in the anharmonicity of the OX stretch. This can be understood from the eigenvalues of a Morse oscillator (second order expansion), which indicate that

$$\frac{\omega_{OX}}{\omega_{OH}} = \rho(1 - 2\rho\chi_e^H)/(1 - 2\chi_e^H).$$

(3.5)

Here $\chi_e^H$ is the anharmonicity of the OH stretch and $\rho = \sqrt{\mu_R^{OH}/\mu_R^{OX}}$. With the gas phase anharmonicity, this expression gives close agreement to the observed $\omega^*$.

The approximate inverse $\sqrt{\mu_R}$ scaling of line width provides more evidence for the recently proposed line broadening mechanism. A series of recent studies have used models based on molecular dynamics (MD) simulations to establish a linear correlation between OX stretching frequency and electric field projected onto the OX coordinate for $\omega_{OH}$ (HOD in D$_2$O) and $\omega_{OD}$ (HOD in H$_2$O) [46, 78, 69, 7]. In simulations of one model, which describes the spectroscopy as a linear interaction of the anharmonic OX Morse potential with the intermolecular degrees of freedom [46], we found the same $\sqrt{\mu_R}$ scaling with line width [69]. The remaining small differences in line width could arise from any of a number of sources. The most likely candidate is motional narrowing, which is seen using the Kubo model as an example.
Given a distribution of OX frequencies $\langle \Delta^2 \rangle$ and a correlation time $\tau_c$, the linewidth is predicted to scale as $\langle \Delta^2 \rangle^{1/2}$ in the inhomogeneous limit and $\langle \Delta^2 \rangle \tau_c$ in the homogeneous limit. Since $\langle \Delta^2 \rangle$ scales as $1/\mu_R$ and $\tau_c$ is invariant with isotope, linewidth scalings between $\sqrt{\mu_R}$ and $\mu_R$ are expected. Additionally, Fermi resonance between the OH stretch and the overtone of the HOD bend is a possible contributor to the enhanced red broadening of the OH stretch in D$_2$O. Anharmonic coupling between the OH stretch and hydrogen bond stretch could also play a role. It is unlikely that this difference results from differences in vibrational population lifetime or reorientational dynamics. Some of these effects are incorporated in the theoretical studies of the OH and OD line shapes by Corcelli and Skinner [54]. Using a combined electronic structure/molecular dynamics approach that correlates electric field with OX frequency, they report a linewidth ratio, $\Delta \omega_{OH} : \Delta \omega_{OD}$, of 1.36 and 1.71 for simulations with the TIP4P [87] and SPC/E [31] water model, respectively.

Electrical anharmonicity, or nonlinearity of the transition dipole moment in the OX coordinate, is a potential explanation for anomalous amplitude or line width variations. The non-Condon effect, which relates the increase of the OH transition dipole moment to an increased hydrogen bonding interaction, is another nonlinearity that has already been shown important for describing the IR-Raman non-coincidence effect [78], and for quantitative modeling of OX stretch nonlinear IR spectroscopy [7, 156]. Its importance is apparent from the 20-fold increase of infrared absorption upon going from the gas to condensed phase. It also explains [7] differences in the reported frequency time-correlation functions of HOD in D$_2$O [68, 69, 93] and HOD in H$_2$O [77, 157, 158]. The non-Condon effect will skew an otherwise symmetric line shape due to frequency dependence of the transition dipole. All three line shapes are similarly asymmetric with a deviation from a Gaussian line shape on the red side. A linear proportionality between the OH transition dipole moment and OH frequency can be used to argue that this effect should not influence the reduced mass scaling of the line shape [7]. We have argued that a nonlinear proportionality is more realistic [156], and it is unclear how this will depend on the OX reduced mass.

It is difficult to explain what leads to the relatively low transition moment for the
OH stretch relative to the OT and OD. One intriguing possibility is the increased hydrogen bond strength for the higher mass species, and the difference in solvent (H₂O vs. D₂O). Studies on H₂O-D₂O mixtures in ice [159, 160] and clusters [161], and the ion-molecule pair I⁻·HDO [162] show the deuteron preferentially hydrogen bonds due to the difference in zero point energy. We investigated whether H₂O or D₂O forms stronger hydrogen bonds through line shape parameters of the HOD bend in both H₂O and D₂O. The peak wavelength is shifted to the red (1460 to 1457 cm⁻¹) and narrower (84 to 79 cm⁻¹) in H₂O. This corresponds to the HOD in D₂O bend parameters at a higher temperature (~40°C). Indeed it is evident that HOD forms stronger hydrogen bonds in D₂O.

### 3.3 Model Calculations of Correlated Frequency Shifts

Previous ultrafast infrared experiments have employed a low concentration of HOD in D₂O/H₂O to study local structural changes in the hydrogen bonded environment of water. 2D IR measurements revealed heterogeneous dynamics in spectrally distinct environments and determined that hydrogen bonds are broken only fleetingly [71, 73]. The transient behavior suggests a mechanism for hydrogen bond switching that involves at least three molecules. The mechanism is an outstanding question that has recently been addressed theoretically [1, 32]. Information on the hydrogen bonded partner can be inferred in experiments on HOD in D₂O/H₂O but a direct spectroscopic signature of the partner has yet to be exploited. Extending to a third isotope offers this possibility. The dynamics of the hydrogen bond environment can be studied with a two-color polarization selective two-dimensional infrared (2DIR) experiment on a low concentration of HOD and HOT in H₂O. The measured signal will not be present without the involvement of multiple distinct molecules. If coupling is dominated by electrostatics, intermolecular coupling strength will be dependent on the relative geometry of the hydrogen bonded oscillators. The experiment is there-
fore capable of directly revealing how motion of the two molecules interlaces with hydrogen bond rearrangements. A clear understanding of how correlated geometric fluctuations map onto changes in frequency is absolutely necessary to derive information from this experiment. This section presents a simple calculation to stipulate on what an experiment may show and how information can be extracted to address the mechanism of hydrogen bond switching.

The linear absorption spectra of HOT in H$_2$O and HOD in H$_2$O are shown in Figure 3-3, the peaks are separated by 380 cm$^{-1}$, or $\sim$240 cm$^{-1}$ between the blue (red) side of the FWHM of $\omega_{OT}$ ($\omega_{OD}$). This system extends the three levels of HOD to six levels. The six levels are the ground state, two $\nu=1\leftrightarrow0$ transitions, two $\nu=2\leftrightarrow1$ transitions, and the combination state. The six level systems Rh(CO)$_2$acac (RDC) in chloroform and hexane [163] and N-methylacetamide (NMA) in various solvents [164, 165] have been previously studied in the Tokmakoff group with 2D IR. In a 2D IR spectrum, spectral information is shown in the peak positions, amplitudes, and line shapes. The anharmonic vibrational potential determines the peak positions. In general, peaks along the diagonal are the $\nu=1\leftrightarrow0$ and $\nu=2\leftrightarrow1$ transitions of the individual modes and cross peaks involve coupling between modes. Amplitudes of the cross peaks reflect the relative magnitudes and orientations of the transition dipoles. These studies successfully determined the relative orientation of the modes and magnitude of the coupling with polarization selective 2D IR. A major difference is that these experiments studied mechanical coupling, or through bond coupling, of normal modes and the proposed experiment will study local modes coupled primarily electrostatically. However, there is some degree of charge transfer upon forming a hydrogen bond in which the lone pair orbitals of the O atom donate to the $\sigma^*_{OH}$ antibonding orbital (see [166] and references within). The goal remains similar in that the cross peaks will be utilized in a time series of 2D IR spectra to observe local structure change during a hydrogen bond switching. The background free cross peaks only appear in the case that the OD and OT oscillators are within approximately one solvation shell of each other.

In the model calculation the local modes are the diagonal elements of the system
Figure 3-3: The OD stretch of 1% HOD in H₂O and the OT stretch of 0.3% HOT in H₂O. The maximum of the OT stretch is scaled by the ratio of the molar absorptivities. See Table 3-1.

Hamiltonian and the off-diagonal terms are the couplings between modes. To construct the Hamiltonian, I. Begin with the gas phase potential for the OX and OY stretch:

\[ H = \begin{bmatrix} \omega_{OX}^{GAS} \\ \omega_{OY}^{GAS} \end{bmatrix} \]  \hspace{1cm} (3.6)

II. Include the modulation to the diagonal terms from solvation, and:

\[ H = \begin{bmatrix} \omega_{OX}^{GAS} + \Delta \omega_{OX} \\ \omega_{OY}^{GAS} + \Delta \omega_{OY} \end{bmatrix} \]  \hspace{1cm} (3.7)

III. Include coupling between modes:

\[ H = \begin{bmatrix} \omega_{OX}^{GAS} + \Delta \omega_{OX} & F_{OX,OY} \\ F_{OY,OX} & \omega_{OY}^{GAS} + \Delta \omega_{OY} \end{bmatrix} \]  \hspace{1cm} (3.8)

In the calculation, the diagonal terms are modulated by the intermolecular electric field and off-diagonal coupling is computed with transition dipole coupling. First we will show the change in electric field and TDC with geometry for a hydrogen bond
dimer. Then the full Hamiltonian will be diagonalized to calculate frequencies during an idealized, concerted hydrogen bond switch.

### 3.3.1 Diagonal Elements: Electric Field

<table>
<thead>
<tr>
<th>Vibration</th>
<th>Empirical Relationship</th>
</tr>
</thead>
</table>
| OH        | $\omega_{10} = 3806.1 \text{ cm}^{-1} - 10792 \text{ cm}^{-1}/\text{a.u. E}$  
|           | $x_{10} = 0.093316 \text{ Å} - 7.3478 \times 10^{-6} \text{ Å/cm}^{-1} \omega_{10}$ |
| OD        | $\omega_{10} = 2792.7 \text{ cm}^{-1} - 7559.5 \text{ cm}^{-1}/\text{a.u. E}$  
|           | $x_{10} = 0.081787 \text{ Å} - 9.266 \times 10^{-6} \text{ Å/cm}^{-1} \omega_{10}$ |
| OT        | $\omega_{10} = 2326.4 \text{ cm}^{-1} - 6596.4 \text{ cm}^{-1}/\text{a.u. E}$  
|           | $x_{10} = 0.062853 \text{ Å} - 6.1355 \times 10^{-6} \text{ Å/cm}^{-1} \omega_{10}$ |
| ALL       | $\mu'/\mu_g' = 1.377 + 53.03 \text{ a.u.}^{-1} \text{E}$ |

Table 3.2: Empirical relationships using the SPC/E model relating the electric field to the frequency and transition dipole for each of the three isotopes. The relationships for OH and OD are from [7] and the relationship for OT was derived by reduced mass scaling.

Corcelli and Skinner [78, 7] performed density functional calculations on clusters of water molecules, generated from liquid-state configurations, to empirically relate a quantity easily extracted from MD simulations to the transition frequency and transition dipole. Correlating the frequency to the electrostatic potential has been a popular technique, but the authors found that for water the electric field is superior [167]. They used linear regression to correlate the electric field from the solvent along the OH (or OD) stretch of the HOD molecule calculated at the H (or D) atom of interest with the *ab initio* frequencies. The relationships used in the model calculation, with the values for OT derived by scaling by the reduced mass, are given in Table 3-2. Auer and Skinner [55, 56] have since reported a quadratic dependence on the electric field; this would be simple to include in the calculations. The electric field at the proton of interest is calculated by:

$$E = \hat{u} \times \sum_{i=1}^{3n} \frac{q_i \hat{r}_{iX}}{r_{iX}^2} \quad (3.9)$$

where $n$ is the total number of molecules and $\hat{u}$ is a unit vector along the intramolecular
bond of interest. The partial charges match the SPC/E model; a three site model with a partial charge of -0.8476 at the oxygen and +0.4238 at each of the protons. As an aside, the SPC/E potential was parameterized to reproduce the experimental room-temperature density and heat of vaporization. However, the model has been shown to be a good choice for understanding dynamics experiments [86]. Each molecule is in the tetrahedral geometry (109.47°) with an intramolecular bond length of 0.97 Å.

Figure 3-4: The change in electric field with geometry at the proton circled in red for each atom of the adjacent water molecule. The total electric field is also shown ($E_{\text{total}} = E_{H1} + E_{H2} + E_O$). The electric field is in atomic units.

Figure 3-4 shows the change in electric field with geometry for a pair of water molecules as $R_{OO}$ ranges between 2.5 and 4 Å and $\alpha$, the angle between the intramolecular bond of the donating HB proton and $R_{OO}$, is varied from 0-90°. The
electric field from each of the three atoms of the hydrogen bond acceptor and the total electric field are shown. The oxygen has the largest contribution. Given the total electric field at $\alpha = 0^\circ$ and $R_{OO} = 2.8\text{Å}$, the average length of a hydrogen bond in water, the frequency for an OD stretch is 2559 cm$^{-1}$, which is $\sim$50 cm$^{-1}$ shifted to the blue from the experimental value. Decreasing $R_{OO}$ by 0.3Å, to 2.5Å, red-shifts the frequency by 156 cm$^{-1}$. A standard geometric criteria for determining if two water molecules are hydrogen bonded is $R_{OO}$ is less than 3.5Å and $\alpha$ is less than 30$^\circ$ [13]. The value for $R_{OO}$ is meant to include the range of the water molecule’s first coordination shell and it is determined by the minimum in the oxygen-oxygen radial distribution function [168]. The angle component was estimated from Debye-Waller factors [169] and it is meant to represent the librational amplitude that will break a hydrogen bond. At $\alpha = 30^\circ$ and $R_{OO} = 3.5\text{Å}$ the OD stretch frequency is 2743 cm$^{-1}$, $\sim$19 cm$^{-1}$ greater than the experimental gas phase value.

### 3.3.2 Off-Diagonal Elements: TDC

The off-diagonal coupling is determined with transition dipole coupling (TDC). TDC will break down if the intramolecular bond distance is on the order of the separation of the dipoles, which are approximated as point dipoles. The TDC is calculated by:

$$F_{XY} = \frac{\partial^2 V_{Ox,Oy}}{\partial Q_{Ox} \partial Q_{Oy}} = \frac{\left(\frac{\partial^2 \phi_{Ox}}{\partial Q_{Ox}^2}\right) \cdot \left(\frac{\partial^2 \phi_{Oy}}{\partial Q_{Oy}^2}\right) - 3\left[\left(\frac{\partial^2 \phi_{Ox}}{\partial Q_{Ox}^2}\right) \cdot n_{Ox,Oy}\left[\left(\frac{\partial^2 \phi_{Oy}}{\partial Q_{Oy}^2}\right) \cdot n_{Ox,Oy}\right]\right]}{R_{Ox,Oy}^3}$$

(3.10)

Figure 3-5 shows the coupling in cm$^{-1}$ calculated by TDC between the transition dipole of the donating proton and each transition dipole of the acceptor with the same geometric change as the electric field calculation above. The transition dipoles are placed at the protons, for convenience, and have a value of 0.5 D/Å. Recently, Auer and Skinner found that placing the dipole along the bond 0.58Å from the oxygen gave the best comparison between intermolecular couplings determined from ab initio calculations in dimers and TDC [56]. This would be simple to incorporate. Finally, the matrix element between the ground and first excited state, $x_{10}$, is set to 1Å, which is exaggerated. Under these conditions the coupling with both dipoles at $\alpha =$
0° and \( R_{OO} = 2.5 \text{Å} \) is -54 cm\(^{-1}\). At \( \alpha = 30° \) and \( R_{OO} = 3.5 \text{Å} \), the TDC is -5 and -29 cm\(^{-1}\) with the transition dipole involving H1 and H2, respectively (see Figure 3-4 for proton number assignments). For comparison, Auer and Skinner [56] calculated the TDC with SPC/E MD simulation and found the average of the largest intermolecular coupling frequencies over many chromophores to be -28 cm\(^{-1}\).

Figure 3-5: Change in TDC with geometry between the transition dipole of the proton circled in red in Figure 3-4 and each of the two transition dipoles of the adjacent water molecule. The TDC is in cm\(^{-1}\).

3.3.3 Model Calculation of Concerted HB Switch

In the model calculation, three molecules are involved in an idealized concerted hydrogen bond switch. The geometric change is similar to that described above. An HOD molecule is undergoing a hydrogen bond switch in which the D atom is first hydrogen bonded to an H\(_2\)O molecule, then subsequently an HOT molecule. Of course, the isotopes can be interchanged. Here D, A1, and A2 refer to donor, first acceptor, and second acceptor, respectively. The switching is concerted in that \( R_{O(D)O(A1)} \) increases from 2.5 to 4 Å as \( \alpha \) increases from 0-90° (or \( \alpha \) decreases from 90-0° with respect to \( R_{O(D)O(A2)} \) and \( R_{O(D)O(A2)} \) decreases from 4 to 2.5 Å). Figure 3-6a shows the initial configuration. Figure 3-6b shows the total electric field at D along with the electric field from each atom of the first acceptor. The second acceptor is the
mirror image reflected over the bifurcation angle. The total electric field is maximum when the deuterium is linearly hydrogen bonded to an acceptor and minimum at the bifurcation angle, 45°, where the distances between the deuterium and the oxygens of the first and second acceptor are equal. Notice the non-hydrogen bonded species contributes very little to the electric field. Figure 3-6c shows the TDC under the same conditions. Again, the second acceptor is the mirror image. Although the hydrogen bonded partner has stronger coupling with the donor while in a linear HB, there is still a fair amount of coupling with the non-hydrogen bonded water molecule (-54 cm\(^{-1}\) with the hydrogen bonded partner compared to ±16 cm\(^{-1}\) with the second acceptor at the initial configuration: \(R_{O(D)O(A1)} = 2.5\text{Å}, R_{O(D)O(A2)} = 4\text{Å},\) and \(\alpha = 0°\)).

![Figure 3-6: Change in electric field and TDC during an idealized concerted HB switch.](image)

(a) Initial Configuration

(b) Total Electric Field [a.u.]

(c) TDC [cm\(^{-1}\)]

\[R_{DO(A2)-DO(A1)} \text{ [Å]}\]

Figure 3-6: Change in electric field and TDC during an idealized concerted HB switch. (a) The initial configuration from the top and side. The total electric field (b) and TDC (c) at the deuterium (donor) due to oxygen (red) and each hydrogen (cyan) of the first acceptor. The dashed blue indicates a broken HB with the first acceptor according to standard geometric criteria.

In the calculation, each proton, beyond the proton undergoing the hydrogen bond
switch, has a dummy water molecule hydrogen bonded so that the calculated frequency better reproduces the condensed phase experimental value. The dummy molecule maintains a linear HB during the switch with $R_{OO} = 2.8 \text{Å}$. Intramolecular coupling is included for the first acceptor, $\text{H}_2\text{O}$, which was estimated as $-50 \text{ cm}^{-1}$ from the splitting of peaks of the isolated molecule. In the condensed phase this may not be rigorous since symmetry is broken by the fluctuating solvent. Auer and Skinner [56] determined an empirical relationship between intramolecular coupling and electric field through \textit{ab initio} cluster calculations on $\text{H}_2\text{O}$ and found a broad distribution with the mean coupling of $-27.1 \text{ cm}^{-1}$. The intramolecular coupling for HOD and HOT in the calculation is set to zero.

To calculate frequencies during the switch we diagonalize the 6x6 Hamiltonian:

$$
H = \begin{bmatrix}
\omega_D & 0 & F_{D,H(T)} & F_{D,T} & F_{D,H(1)} & F_{D,H(2)} \\
0 & \omega_{H(D)} & F_{H(D),H(T)} & F_{H(D),T} & F_{H(D),H(1)} & F_{H(D),H(2)} \\
F_{H(T),D} & F_{H(T),H(D)} & \omega_{H(T)} & 0 & F_{H(T),H(1)} & F_{H(T),H(2)} \\
F_{T,D} & F_{T,H(D)} & 0 & \omega_T & F_{T,H(1)} & F_{T,H(2)} \\
F_{H(1),D} & F_{H(1),H(D)} & F_{H(1),H(T)} & F_{H(1),T} & \omega_{H(1)} & intra \\
F_{H(2),D} & F_{H(2),H(D)} & F_{H(2),H(T)} & F_{H(2),T} & intra & \omega_{H(2)}
\end{bmatrix}
$$

Figure 3-7a displays the frequencies of the OD and OT stretch during the hydrogen bond switch for $\mu = 0 \text{ D/Å}$ (solid) and $\mu = 0.5 \text{ D/Å}$ (dashed and dot-dashed). The dipole moments are constant and equal for all bonds. The OD stretch frequency changes most drastically as this is the only proton switching hydrogen bond partners and, as expected, it moves from the red to the blue and back to the red during the switch. Note that with intermolecular coupling turned on, $\omega_{OD}$ is blue shifted when it is hydrogen bonded to HOT relative to $\text{H}_2\text{O}$. $\omega_{OD}$ is 2407 cm$^{-1}$ and 2419 cm$^{-1}$ when in a linear HB with $\text{H}_2\text{O} (R_{OO} = 2.5 \text{ Å})$ and HOT, respectively. Although the site energy for $\omega_{OT}$ is fixed by the dummy molecule accepting a hydrogen bond from OT, $\omega_{OT}$ changes during the hydrogen bond switch due to the change in off-diagonal coupling. Figure 3-7b shows the correlation between $\omega_{OD}$ and $\omega_{OT}$. Note that there is no correlation when the coupling is turned off. With intermolecular coupling, $\omega_{OD}$
and $\omega_{OT}$ are positively correlated while in a hydrogen bond, which will result in a diagonally elongated cross peak parallel to the diagonal. Polarization selective 2D IR experiments can measure the average angle between the dipoles. The value is expected to be near $70.53^\circ$ where the frequencies are correlated. This value can be evaluated in a time series to see how relative orientation changes as the molecule undergoes hydrogen bond breaking and reforming.

This simple calculation does not take into consideration the non-Condon effect. Since the transition dipole increases with the strengthening of the hydrogen bond [156, 7] the non-Condon effect is expected to cause a larger disparity in intermolecular coupling between HB and NHB species. This can easily be included by using the relationships in Table 3-2. There are two main experimental difficulties that need to be considered. The first issue is dilution. This experiment requires an HOD and HOT molecule to be within two solvation shells of one another. If instead two HOD(T) water molecules are near each other resonant energy transfer will dominate. The second related issue is strength of coupling. The coupling must be strong enough such that the inverse time would be faster than the dynamics of interest. Otherwise, the experiment will only be able to report on the kinetics of energy transfer, however difficulties concerning the vibrational lifetime and hot ground state need to be kept
under consideration.

### 3.4 Gaussian Calculations

We performed B3LYP/6-311++G** density functional calculations with Gaussian98 on water isotopologue dimer combinations to determine the intermolecular coupling between $\omega_{OD}$ and $\omega_{OT}$. The calculations were performed with water as the solvent, which applies a constant dielectric field, and the dimer geometry was optimized. The intramolecular bond lengths of the hydrogen bond accepting water molecule are 0.9629 Å. However, the intramolecular bond lengths of the hydrogen bond donating water molecule are unequal; the donating OX bond length is relatively lengthened to 0.971 Å and the free OX bond length is shortened to 0.9615 Å. Also, the angle between the protons of the accepting water molecules is 105.8098° compared to 103.9830° for the donating water molecule. $R_{OO}$ is 2.9004 Å and the intramolecular OX bond is tilted off the line that connects the two oxygens by 5.6985°. In other words, the calculation does not predict a perfectly linear hydrogen bond.

Vibrational frequencies were calculated for four dimer pairs: (a) HOD-H$_2$O, (b) HOD-HOT, (c) HOT-H$_2$O, and (d) HOT-HOD, where the atom in bold is the hydrogen bond donor. The spectra for dimers (b) and (d) are shown in Figure 3-8. In agreement with the trend observed between gas and condensed phase vibrational frequencies, the OX stretch red-shifts and the HOX bend blue-shifts upon forming a hydrogen bond. For example, the OT stretch and HOT bend shifts from 2356.1099 and 1334.0667 cm$^{-1}$ to 2263.3100 and 1358.2596 cm$^{-1}$, respectively, between dimer (b) and (d). Likewise the OD stretch and HOD bend shifts from 2808.4353 and 1417.9195 cm$^{-1}$ to 2697.5080 and 1430.0400 cm$^{-1}$, respectively, between dimer (b) and (d). Although the shift in OX stretch frequency is similar between the two isotopes, the HOT bend blue-shifts by more than 2x relative to the HOD bend upon forming a hydrogen bond. Also, the intensity of $\omega_{OD}$ increases by a factor of 6.1988 upon forming a hydrogen bond compared to 5.6528 for $\omega_{OT}$. Furthermore, if an H$_2$O is added to dimer (b) hydrogen bonded to the free OH stretch of HOD, the OD
stretch blue-shifts to 2721.6995 cm\(^{-1}\) (0.9689 Å) relative to dimer (b) but the HOD bend further blue-shifts to 1470.3225 cm\(^{-1}\) (104.1452°). Also, the HOD hydrogen bond becomes more linear in the trimer with the angle decreasing from 5.6985 to 3.6420°.

![Figure 3-8: Mid-infrared spectra of dimers (b) (blue) and (d) (red); also, dimer (b) with additional H\(_2\)O (dashed cyan).](image)

With the four dimer combinations we can estimate the intermolecular coupling between \(\omega_{OD}\) and \(\omega_{OT}\). The diagonal elements are \(\omega_{OD}\) and \(\omega_{OT}\) when hydrogen bonded to H\(_2\)O and the off-diagonal element is the coupling between modes. Diagonalizing the matrix gives \(\omega_{OD}\) and \(\omega_{OT}\) when hydrogen bonded to HOT and HOD, respectively. From the calculation, the OD(T) stretch blue(red)-shifts from 2697.5743 to 2697.5080 cm\(^{-1}\) (2663.4638 to 2263.3100 cm\(^{-1}\)) when the H\(_2\)O hydrogen bond acceptor is replaced by HOT(D). Given these frequencies, the coupling is -8.1778 cm\(^{-1}\).

### 3.5 Conclusions

The infrared absorption spectrum of the OT stretch in tritiated water has been reported, making comparison of this line shape to \(\omega_{OD}\) and \(\omega_{OH}\) spectra. The inadequacy of harmonic scaling relationships in predicting resonance frequencies, amplitudes and line widths is an indication that these approximations, commonly used in the recent simulations of water vibrational spectroscopy, need to be re-visited. The
results suggest that a more nuanced description of electrical and mechanical anharmonicity in the interactions of the OX stretch with the electromagnetic field and intermolecular motions will be necessary to obtain a quantitative understanding of the spectroscopy. Comparison of ultrafast IR spectroscopy on these isotopologues can be used to obtain this insight.

Studies of the HOD in D₂O system with broadband infrared non-linear techniques have provided insight into the fastest timescales of motion in water. 2D IR measurements revealed frequency dependent spectral diffusion. The results imply that the non-hydrogen bond species are transient, which leads to the hypothesis that hydrogen bond switching is a many body problem. This mechanism, however, cannot be determined with the aforementioned system. Extending 2D IR measurements to directly probe two spectrally distinct species provides a route to elucidate the mechanism. Model calculations reveal that \( \omega_{OD} \) and \( \omega_{OT} \) are correlated when they are hydrogen bonded. Polarization selective waiting time dependent 2D IR measurements on this system may reveal the relative structural change during a hydrogen bond switch. An important quantity to consider in the feasibility of the experiment is the intermolecular coupling. With \textit{ab initio} calculations, we found the coupling to be \( \sim 8 \) cm\(^{-1}\).

### 3.6 Acknowledgments

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Chapter 4

Vibrational Relaxation in Isotopically Dilute Water

4.1 Introduction

Understanding vibrational energy redistribution in solution is important for a number of reasons. First, reaction rates and product distributions can be quite different for reactions occurring in the gas and condensed phases. For instance, if vibrational energy redistribution is important in a reaction, the solvent may alter the rate by providing an efficient pathway for energy transfer. This may be an important factor in photoisomerization, proton and electron transfer, and some photodissociation reactions [170, 171]. An understanding of vibrational relaxation processes in solution is an essential piece in the development of a predictive theory of reaction dynamics. Second, comparing the experimental lifetime to the calculated lifetime provides a sensitive test of the adequacy of potential energy surfaces. Finally, we will show that vibrational relaxation and spectral diffusion occur on a similar time scale in water. A clear picture of the vibrational lifetime and pathway for vibrational relaxation will be important since energy dissipation to the low frequency modes has a non-negligible affect on our spectral measurements.

In order to characterize the vibrational relaxation pathway(s) in isotopically dilute water we will compare previous measurements of the temperature dependence of the
lifetime of the OH stretch of HOD in D$_2$O, or T$_{1,OH}$, with the temperature dependence of the lifetime of the OD stretch of HOD in H$_2$O, or T$_{1,OD}$, and the room temperature lifetime of the OT stretch of HOT in H$_2$O, or T$_{1,OT}$. The intramolecular vibrational modes of water are anharmonically coupled. Vibrational relaxation may include a number of downhill or uphill intramolecular pathways with the solvent taking up or contributing excess energy, respectively. What would we expect to see if instead the intramolecular modes were orthogonal? The transition frequencies of $\omega_{OH}$, $\omega_{OD}$, and $\omega_{OT}$ scale largely by reduced mass (see chapter 3). If the dominant relaxation pathway is to the ground vibrational level, one would expect $T_{1,OH} > T_{1,OD} > T_{1,OT}$ since $\omega_{OH} > \omega_{OD} > \omega_{OT}$. Also, one may expect the lifetime to decrease with increased temperature since there will be increased occupancy of the lower energy-accepting bath modes. Neither of these trends have been observed experimentally, however.

An early theoretical investigation of the vibrational relaxation of the OH stretch of HOD in D$_2$O suggested that the intermolecular hydrogen bond stretch may be the dominant accepting mode ($\sim$180 cm$^{-1}$) [172]. In other words, an initially intact hydrogen bond ruptures upon vibrational relaxation. The vibrational lifetime was calculated via a Fermi Golden Rule approach and found to be related to the shift from the gas phase absorption frequency by a power law dependence: $T_{1,OH} = a/(\delta \omega_{OH})^{1.8}$, where a is a constant. The relationship predicts the lifetime will decrease as the frequency red-shifts. Increased hydrogen bond strength softens the OH intramolecular potential, which results in a red-shift of $\omega_{OH}$. With increased temperature there are increased fluctuations in the hydrogen bond network, which results in a blue-shift in the absorption band. The proposed mechanism, then, predicts the lifetime will increase with increased temperature.

Woutersen and coworkers measured the the vibrational lifetime of a 500 $\mu$m layer of a 1:500 solution of HOD in D$_2$O within the range of 30-363 K [97]. In the entire temperature range for ice there is no significant change in $T_{1,OH}$, which is approximately 380 fs. At the melting point, $T_{1,OH}$ suddenly increases from $\sim$380 to $\sim$750 fs and increases by an additional $\sim$20% at 363 K. The observed trend could be reasonably described by the proposed vibrational predissociation mechanism. However,
the possibility that the increase in lifetime with temperature may be the result of an increasing energy gap between the OH stretch and different inter- or intramolecular accepting modes could not be ruled out.

More recent theoretical work by Rey and Hynes [103] and Lawrence and Skinner [104, 105] found that the overtone of the HOD bend was the dominant relaxation pathway for the OH stretch, which also agrees with the experimentally observed temperature dependence of the lifetime. As the temperature is increased, hydrogen bond fluctuations increase and the stretch red-shifts while the bend blue-shifts toward their respective gas phase values. If the bend anharmonicity is constant with temperature the overtone will shift in the same manner as the bend. This will decreases the overlap integral between the overtone of the HOD bend and OH stretch, which results in a decrease in coupling and a longer lifetime.

Dlott and co-workers performed the only experiments on isotopically dilute water that directly probed vibrations beyond the stretch after excitation of the stretch using a mixed IR/Raman technique with ~1.5 ps time resolution [92]. They conclude that the primary route of energy relaxation is through a pair of HOD bend vibrations facilitated by intermoleculer energy transfer to D$_2$O. However, it is possible the overtone of the bend was not observed given that the lifetime of the fundamental of the HOD bend in D$_2$O is 390 fs [96]. Lindner et al. performed two-color IR-pump probe (PP) experiments on the bend and stretch vibrations of pure H$_2$O with ~200 fs time resolution [109]. Albeit a very different species as far as vibrational relaxation is concerned, they too found the vibrational energy of the photo-excited H$_2$O stretch is deposited to a pair of bending vibrations.

A theoretical study on the OD stretch of HOD in H$_2$O, performed with a similar method as the calculations on the OH stretch [103, 104, 105], found the dominant pathway was either through the fundamental of the HOD bend or directly to the ground state depending on if the bath was treated as rigid or flexible, respectively [106]. The room temperature [95, 94] and temperature dependent [100] lifetime of the OD stretch in HOD in H$_2$O has been measured. Steinel et al. originally reported T$_{1,OD} = 1.45$ ps for the lifetime at the room temperature [94]. More recently, Fayer
and co-workers used a different method to analyze the dispersed pump probes (DPPs) and reported a lifetime of 1.7 ps [95]. The longer lifetime of the OD stretch compared to the OH stretch agrees with the proposed mechanism for vibrational relaxation given that the frequency difference between the OH stretch and the HOD bend overtone is $\sim 500 \text{ cm}^{-1}$ and the difference between the OD stretch and the HOD bend fundamental, the closest intramolecular mode that is lower in energy, is $\sim 1050 \text{ cm}^{-1}$. Tielrooij and Bakker [100] measured the temperature-dependence and found the lifetime increased from approximately 1.7 to 2.2 ps from 274 to 343 K. However from 274 K to room temperature the lifetime is constant within error.

The most direct method to elucidate the vibrational relaxation pathway in water is narrowband (or broadband) pump-broadband probe experiments. In this context, narrowband refers to a pulse that spans the entire spectral region of the vibration of interest ($\nu=1\leftarrow 0$ and the anharmonically shifted $\nu=2\leftarrow 1$) and the broadband probe spans the entire mid-infrared. Also, in this idealized experiment, the time resolution would be fast enough to detect even the initial energy transfer. This experiment has yet to be realized in practice, however there has been progress in this direction [173].

One way to test the proposed vibrational relaxation pathway is to compare the lifetime of different water OX stretches (where X = H, D, and T). In this chapter we report the lifetime of the OT stretch of dilute HOT in H$_2$O and compare the relaxation rate to the other isotopes. In Figure 4-1 we show a diagram of the peak positions of the intramolecular and intermolecular modes of HOD in D$_2$O (left), HOD in H$_2$O (middle), and HOT in H$_2$O (right). For $\omega_{OT}$ (2120 cm$^{-1}$ [174]), the nearest intramolecular vibration that is lower in energy is the HOT bend. From the Gaussian calculations described in chapter 3 we can estimate the frequency of the HOT bend. The frequency of the HOD bend in a trimer in which the central HOD molecule is hydrogen bonded to two H$_2$O molecules is 1470 cm$^{-1}$ and the experimental value in the condensed phase is 1450 cm$^{-1}$. If we replace the HOD with a HOT the frequency of the HOT bend from the calculation is 1400 cm$^{-1}$. If we scale this by comparing the calculated HOD bend frequency with the experimental value the HOT bend frequency is $\sim 1380 \text{ cm}^{-1}$. The frequency difference between $\omega_{OT}$ and $\delta_{HOT}$ is then $\sim 740 \text{ cm}^{-1}$. 
Since the energy difference is in between the energy difference between $\omega_{OH} - 2\delta_{HOD} \approx 500 \text{ cm}^{-1}$ and $\omega_{OD} - \delta_{HOD} \approx 1050 \text{ cm}^{-1}$, one would expect $T_{1,OT}$ to also be in between 700 - 740 fs [93, 98] and 1.45 - 1.7 ps [95, 94, 100].

Another handle we have to test the proposed vibrational relaxation mechanism is temperature. Since changing the temperature alters the fluctuations in the hydrogen bond network in water and the vibrational frequencies are sensitive to hydrogen bonding, vibrational bands shift with changing temperature. Vibrational modes are uniquely dependent on the degree of hydrogen bonding and, therefore, have unique dependencies on temperature. By changing the temperature we can tune the energy gap between modes and compare the experimentally measured change in lifetime with the predicted change. In this chapter we will present the temperature dependent lifetime of $\omega_{OD}$ of HOD in H$_2$O measured with dispersed pump probes and compare to changes in vibrational frequencies with temperature.

Figure 4-1: Energy level diagram of three isotopologues of water: HOD (cyan) in D$_2$O (green), HOD in H$_2$O (blue), and HOT (red) in H$_2$O.
4.2 Dispersed Pump Probe of Tritiated Water

The sample conditions for transient absorption measurement of ω_{OT} of HOT in H\textsubscript{2}O are the same as described in Chapter 3 for acquiring the absorption spectrum. The concentrations of HOT and the solvent H\textsubscript{2}O are approximately 0.2 M and 55 M, respectively. Given the molar absorptivities \cite{112, 174} the ratio of the optical density of the combination band to ω_{OT} is ~30. Therefore, absorption from the background H\textsubscript{2}O is a complication in the transient absorption measurement. Also, although heterodyned detected third order measurements and linear absorption measurements are proportional to the fourth and second power of the transition dipole, respectively, third order signal from the combination band may contribute to the third order signal at this concentration of HOT \cite{112}. The pump-probe was measured with the interferometer described in Chapter 2 and the pulse spectrum is shown in Figure 4-2. It is peaked at ~2045 cm\textsuperscript{-1} with a pulse width of ~220 cm\textsuperscript{-1} at full width half maximum (FWHM) and pulse length of ~85 fs. In Figure 4-2 we show the linear absorption spectrum of the OT stretch of HOT in H\textsubscript{2}O with the background H\textsubscript{2}O subtracted and the absorption of the H\textsubscript{2}O combination band plotted against the molar absorptivity. The combination band is the combination of the bend (1650 cm\textsuperscript{-1}) and libration (peak at ~680 cm\textsuperscript{-1} and FWHM from ~420 to 860 cm\textsuperscript{-1}). The peak of the combination band is ~2130 cm\textsuperscript{-1} and the linewidth is approximately 300 cm\textsuperscript{-1} (FWHM) \cite{112}.

In Figure 4-3 we show slices in time of the experimental transient absorption measurement, a model pump-probe of ω_{OT}, and the absorption spectrum of the combination band of H\textsubscript{2}O. The model pump-probe is a difference of two Gaussians with one shifted by the anharmonicity (ω\textsubscript{10}−ω\textsubscript{21}=Δ):

\[
S(\omega) = \exp[-(\omega - \Omega)^2/\sigma^2] - \exp(-(\omega - (\Omega - \Delta))^2/\sigma^2).
\]  

(4.1)

In this equation, Ω is the central frequency and σ is related to the line width. In this model the line width of the ν=2→1 is the same as the ν=1→0, however, it has been suggested that the width of the ν=2→1 is considerably larger than the fundamental
The anharmonicity of the $\omega_{OT}$, $\Delta = 130$ cm$^{-1}$, was estimated by reduced mass scaling from the anharmonicity of $\omega_{OD}$ (160 cm$^{-1}$ [77]) and $\omega_{OH}$ (250 cm$^{-1}$ [60]). In the $\nu=2\leftarrow1$ region the experimentally measured transient absorption appears quite similar to the model. The induced absorption decays towards zero as the waiting time is increased. In the $\nu=1\leftarrow0$ region, however, the transient absorption remains largely flat with positive amplitude near the zero crossing shifting slightly towards lower frequency with increased waiting time, reminiscent of the shift observed due to the thermally shifted ground state (TSGS, discussed in the last section of this chapter). Also, there is a slight dip in the positive amplitude near the peak of the combination band. It appears as though absorption by the background H$_2$O is non-negligible. Also, the experiment was performed in the zzzz polarization geometry for maximum signal. The decay is, therefore, sensitive to vibrational relaxation and reorientation. Nonetheless we can get an idea of the vibrational lifetime by fitting the decay of the signal in the $\nu=2\leftarrow1$ region. Fitting a mono-exponential past the cross-phase modulation (discussed in Chapter 2) results in a decay of $\sim850$ fs, which will be faster than $T_{1,OT}$ due to the contribution from reorientation. Indeed the decay time is in between $T_{1,OH}$ and $T_{1,OD}$, and therefore agrees with the proposed
relaxation mechanism in which the dominant relaxation pathway is through the closest intramolecular mode that is lower in energy.

Figure 4-3: Slices in time of the transient absorption of HOT in H₂O (circles), the model pump-probe of ω₀T (blue) and the linear absorption of H₂O (green).

In Figure 4-4a we show a model dispersed pump-probe of ω₀T and combination band of H₂O. The anharmonicity of the combination band is 300 cm⁻¹ [112]. It is interesting to note that the anharmonicity is roughly equal to the line width at FWHM for all the transitions. In Figure 4-4a the frequencies marked at 1993 (blue), 2008 (green), 2023 (red), and 2038 cm⁻¹ (cyan) correspond to induced absorption for ω₀T and bleach/stimulated emission for the combination band of H₂O. The zero-crossing of the ω₀T PP is near 2054 cm⁻¹ (magenta) and at 2070 cm⁻¹ (gold) both the PP of ω₀T and the combination band show a bleach/stimulated emission. Slices in frequency of the PP measurement (Szzzz) of HOT in H₂O is shown in Figure 4-4b for the frequencies marked in Figure 4-4a. The slices at 1993-2038 cm⁻¹ begin negative and eventually cross to positive values at longer waiting times, which is most likely due to the TSGS (which will be discussed in the last section of this chapter). At 2054 and 2070 cm⁻¹ the transient absorption is positive and remains largely flat over the time frame of the experiment. From the experimental slices in frequency we can be confident we are mainly sensitive to the transient absorption of ω₀T, not the
combination band of H\textsubscript{2}O.

![Graph of linear absorption](image)

Figure 4-4: (a) Model pump-probe of $\omega_{OT}$ (blue) and the combination band of water (green) with the frequencies in (b) marked with lines. (b) Slices in frequency of the experimental pump-probe of HOT in H\textsubscript{2}O.

### 4.3 Temperature Dependent Lifetime of $\omega_{OD}$

#### 4.3.1 Temperature Dependent Linear Absorption

We report the temperature dependence of the linear absorption of HOD in H\textsubscript{2}O to predict the temperature dependence of $T_{1,OD}$ given the proposed mechanism in which relaxation occurs primarily through the fundamental of the HOD bend. In Figure 4-5 we show the change in linear absorption with temperature for HOD in H\textsubscript{2}O without (a) and with (b) the background H\textsubscript{2}O subtracted. The maximum of $\omega_{OD}$, the HOD bend, and the H\textsubscript{2}O combination band is shown as black dashed lines and the spectra at the two temperature extrema are shown in the top row of the figure. The HOD bend red-shifts and narrows and $\omega_{OD}$ blue-shifts and broadens with increasing temperature. In this temperature range the maximum of the H\textsubscript{2}O combination band $(\nu_L + \delta)$ red shifts by 70 cm\textsuperscript{-1} from 2156 cm\textsuperscript{-1} at 277 K to 2086 cm\textsuperscript{-1} at 348 K. The intense peak at $\sim$1650 cm\textsuperscript{-1} in Figure 4-5a is the H\textsubscript{2}O bend and the broad featureless absorption that extends from the low frequency side of the spectrum is the librational band.
Figure 4-5: Temperature dependent linear absorption of HOD in H₂O without (a) and with (b) the background H₂O subtracted. (Top) Linear absorption at the lowest and highest temperature of the contour plot. (Bottom) Dashed lines indicate the change in peak maxima with temperature for (a) δ_{HOD}, H₂O δ+ν₇, and ω_{OD} (left to right) and (b) δ_{HOD} (left) and ω_{OD} (right).

Finally, the low intensity but relatively broad peak at ~1850 cm⁻¹ in Figure 4-5b is the combination band (ν₇ + δ) of HOD.

In Figure 4-6 we show the change in peak position (a) and line width (FWHM) (b) for ω_{OD} (blue) and δ_{HOD} (green). The change in frequency is relative to the value at 308 K, which is 2514 cm⁻¹ (1454 cm⁻¹) and 166 cm⁻¹ (83 cm⁻¹) for the peak and FWHM of ω_{OD} (δ_{HOD}), respectively. ω_{OD} and δ_{HOD} have the opposite trend in both frequency and FWHM with temperature and although the peak frequency of ω_{OD} is more sensitive to temperature the FWHM of δ_{HOD} changes more drastically with temperature. Equations describing the temperature dependencies are given in Table 4-1. In Figure 4-6c we show the energy gap between ω_{OD} and δ_{HOD} with temperature at the peak of each transition (cyan) and at the FWHM (red; red side of ω_{OD} minus the blue side of δ_{HOD}). The energy gap broadens from 1038 cm⁻¹ (919 cm⁻¹) to 1086 cm⁻¹ (963 cm⁻¹) at the peak (FWHM) between 277 to 348 K. In both cases the change with temperature is linear and the overall change is ~5%. In Figure 4-6d
Figure 4-6: (a) Change in peak frequency with temperature for $\omega_{OD}$ (blue) and $\delta_{HOD}$ (green) relative to the value at 308 K (2514 and 1454 cm$^{-1}$, respectively). (b) Change in line width (FWHM) relative to the value at 308 K (166 and 83 cm$^{-1}$, respectively). Difference in wavenumbers between (c) $\omega_{OD}$ and $\delta_{HOD}$ and (d) $2\delta_{HOD}$ and $\omega_{OD}$ at the peak of each transition (cyan) and closest gap between the FWHM (red).

we show the energy gap between $\omega_{OD}$ and the overtone of $\delta_{HOD}$ ($2\delta_{HOD}$) at the peak (cyan) and FWHM (red; red side of $2\delta_{HOD}$ minus the blue side of $\omega_{OD}$) assuming the anharmonicity is zero and the line width is the same for the fundamental and overtone of the bend, which are most likely not valid assumptions. Nonetheless, the energy gap narrows from 422 cm$^{-1}$ (236 cm$^{-1}$) to 363 cm$^{-1}$ (209 cm$^{-1}$) between 277 to 348 K at the peak (FWHM) of the two transitions. Note that 209 cm$^{-1}$ is less than kT at 348 K. The change with temperature at the peak is quite linear although the change at FWHM is slightly quadratic and the overall change is $\sim$14% and $\sim$11% at the peak and FWHM, respectively.

If the OD stretch decays to the closest intramolecular mode that is lower in energy one may expect a linear increase in $T_{1,OD}$ with increased temperature since the energy gap between $\omega_{OD}$ and $\delta_{HOD}$ linearly increases with temperature. However, although
Temperature Dependence

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Expression</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peak, $\omega_{OD}$</td>
<td>$0.5199 \text{ cm}^{-1}/K \times T[K] + 2354 \text{ cm}^{-1}$</td>
</tr>
<tr>
<td>FWHM, $\omega_{OD}$</td>
<td>$1.601 \times 10^{-3} \text{ cm}^{-1}/K^2 \times T^2[K^2] + 1.142 \text{ cm}^{-1}/K \times T[K] - 33.69 \text{ cm}^{-1}$</td>
</tr>
<tr>
<td>Peak, $\delta_{HOD}$</td>
<td>$-0.1497 \text{ cm}^{-1}/K \times T[K] + 1502 \text{ cm}^{-1}$</td>
</tr>
<tr>
<td>FWHM, $\delta_{HOD}$</td>
<td>$-0.2882 \text{ cm}^{-1}/K \times T[K] + 171.9 \text{ cm}^{-1}$</td>
</tr>
</tbody>
</table>

Table 4.1: Temperature dependencies of the line shape parameters of $\omega_{OD}$ and $\delta_{HOD}$ and the energy gap between the stretch and fundamental and overtone of the bend

the energy gap between $\omega_{OD}$ and $\delta_{HOD}$ increases with increasing temperature, the energy gap between the $\omega_{OD}$ and the $2\delta_{HOD}$ decreases. Also, the energy gap between $\omega_{OD}$ and $2\delta_{HOD}$ is $\sim 2kT$ or less over the entire temperature range. Therefore, the lifetime may show a non-trivial dependence with temperature or even decrease with increased temperature if relaxation to the overtone of the bend is the dominant pathway.

4.3.2 Temperature Dependent Dispersed Pump Probes

In Figure 5-1 we show the linear absorption of HOD in H$_2$O at five temperatures with (a) and without (b) the background H$_2$O subtracted. A 5% solution of D$_2$O in H$_2$O was held in a home-built brass sample cell between 1 mm CaF$_2$ windows with a 6 $\mu$m teflon spacer. The temperature of the sample cell was controlled with a water-cooled chiller. The condensed phase spectra red shifts by $\sim 200 \text{ cm}^{-1}$ from the gas phase ($2723.68 \text{ cm}^{-1}$ [159]) due to hydrogen bonding. The $\sim 160 \text{ cm}^{-1}$ line width (FWHM) primarily reflects a distribution of hydrogen bond strengths [155]. From 278 to 345 K the peak blue shifts from 2499 to 2534 cm$^{-1}$ (0.53 cm$^{-1}$/K), indicating an increase in hydrogen bond fluctuations, and the FWHM broadens from 159 to 180 cm$^{-1}$. However, the total area decreases with increasing temperature due to the non-Condon effect [156, 7]. In the same temperature range the HOD bend, which can be seen in Figure 5-1b, narrows and red-shifts from 1460 to 1450 cm$^{-1}$. 

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Figure 4-7: Temperature-dependent linear absorption measurements of dilute HOD in H2O with (a) and without (b) the solvent H2O subtracted. The spectra are offset along the absorbance axis.

Transient absorption measurements at the five temperatures shown in Figure 5-1 are shown in Figure 4-8. Experimental conditions will be described in detail in the next chapter. Briefly, the mid-infrared pulses were centered at ~2520 cm⁻¹, the pulse width was ~230 cm⁻¹ (FWHM), and the pulse duration was ~80 fs. The transient absorption measurements are collected in the Szzzz and Szzzy polarizations and the magic angle signal is constructed as: $S_{iso} = (S_{zzzz} + 2S_{zzzy})/3$. Effects from the TSGS (discussed in the next section) are apparent within experimental time window (~3.65 ps). As the delay between the pump and probe increases, the bleach shifts towards lower frequency and a negative feature grows in above the bleach. The absorption from atmospheric CO2 at ~2320 to 2360 cm⁻¹ significantly decreases the signal from the ν=2<–1 transition. The DPPs at the five temperature appear quite similar beyond the expected shift to higher frequency with increased temperature. Slices at the frequency with maximum intensity, normalized to the value at $t = 200$ fs, are shown in the last subplot in Figure 4-8. The transient absorption appears quite similar between the five temperatures, with a possible slight increase in $T_{1,OD}$ with increasing temperature. The thermally shifted ground state (TSGS) must be accounted for, however, before $T_{1,OD}$ can be determined.

Bakker and co-workers have reported the temperature dependent transient ab-
Figure 4-8: Transient absorption measurements of HOD in H$_2$O at 278, 286, 295, 323, and 345 K and slices in frequency with maximum intensity at each of the five temperatures.

Absorption of $\omega_{OD}$ in isotopically dilute water [100]. To determine $T_{1,OD}$ they used a fit model that includes decay from $\nu=1$ to an intermediate level, which they suggest is the fundamental of the HOD bend, before relaxation to the low frequency modes [120]. The transient absorption is written as:

$$
\Delta \alpha (\omega, t) = \Delta \alpha_{\text{end}}(\omega) \left[ \frac{T^*}{T_1 - T^*} e^{-t/T^*} - \frac{T_1}{T_1 - T^*} e^{-t/T_1} + 1 \right]
+ N_1(0) \Delta \alpha_{\text{init}}(\omega) e^{-t/T_1}
$$

(4.2)

where $\Delta \alpha_{\text{init}}(\omega)$ and $\Delta \alpha_{\text{end}}(\omega)$ denote the spectral contribution from the initial excitation, which includes $\nu=1\rightarrow0$ and $\nu=2\rightarrow1$, and TSGS, respectively. The shape of the two spectral components are independent of time delay. $T_1$ is the vibrational
lifetime of $\omega_{OD}$, $T^*$ is the lifetime of the intermediate state, and $N_1(0)$ denotes the initial population in the $\nu=1$ level. In Figure 4-9 we show the result of applying the fit model at 295 K starting at $t=200$ fs. The model (green) fits the data (blue) very well. In Figure 4-9a we show a frequency slice at $\omega = 2522$ cm$^{-1}$, which is in the $\nu=1\leftarrow 0$ region. At early delays, $\Delta \alpha_{\text{init}}$ is the dominant contribution to $\Delta \alpha$. As the delay is increased, $\Delta \alpha_{\text{end}}$ grows in causing the trace to decay to a positive offset at this frequency. In Figure 4-9c we show a frequency slice at $\omega = 2380$ cm$^{-1}$, which is in the $\nu=2\leftarrow 1$ region. Although the trace begins negative, it crosses to positive value at $\sim 2$ ps due to the growth of the TSGS. In Figure 4-9b and d we show slices in time at $t=1$ ps and 3.65 ps, respectively. At 1 ps, $\Delta \alpha_{\text{init}}$ is dominant but at 3.65 ps, $\Delta \alpha_{\text{end}}$ is the dominant contribution. Again, in this fit model the spectral shape of $\Delta \alpha_{\text{init}}$ and $\Delta \alpha_{\text{end}}$ are independent of delay; only the relative contribution of the two spectral components is time dependent.

![Figure 4-9](image_url)

Figure 4-9: Experimental transient absorption at 295 K (blue) with the results from the fit (green) and $\Delta \alpha_{\text{init}}$ (red dashed) and $\Delta \alpha_{\text{end}}$ (cyan dashed) contributions shown separately at (a) $\omega = 2522$ cm$^{-1}$, (b) $\omega = 2380$ cm$^{-1}$, (c) $t = 1$ ps, and (d) $t = 3.65$ ps.
In Figure 4-10 we show both $T_1$ and $T^*$ from the fit model for the five temperatures in this study. The uncertainty is $\pm 75$ fs. Tielrooij and Bakker [100] measured the temperature-dependent vibrational lifetime and found it increased from approximately 1.7 to 2.2 ps from 274 to 343 K. In the current study we find the vibrational lifetime increases from 1.4 to 2.0 ps from 278 to 345 K. These lifetimes are somewhat faster than those of Tielrooij but agree with the lifetime first reported by Fayer and co-workers [94]. Also, the intermediate lifetimes in the current study increase relatively slightly from 450 to 590 fs across the temperature range compared to the decrease from $\sim 900$ to $350$ fs from 274 to 343 K in the study performed by Tielrooij and Bakker [100]. In both studies the isotropic pump-probe is constructed from the $S_{zzzz}$ and $S_{zzyy}$ transient absorption measurements acquired with the pump $45^\circ$ relative to the probe. An important difference is that the measurements in the former study are taken to $t = 10+$ ps but only to $t = 3.65$ ps in the current study. Measuring to longer delay times is ideal. An interesting similarity between the two measurements, however, is that the vibrational lifetime of the OD stretch is largely unchanged from the lowest temperature to room temperature. We found that fitting the integrated isotropic transient absorption assuming that the TSGS grows in at the same rate as the decay of the excited state resulted in similarly decent fits within our experimental time window. The resulting lifetimes were slightly shorter but followed the same trend. This observation does not rule out that vibrational energy decays through an intermediate state, it only implies that decay from the intermediate is very fast. Given this, we will not discuss $T^*$ further.

Although the energy gap between $\omega_{OD}$ and the fundamental and overtone of the bend show, to first approximation, a linear dependence with temperature, the lifetime has a non-trivial temperature dependence. $T_{1,OD}$ remains largely the same from 278 K to 295 K, then increases by $\sim 40\%$ by 345 K. The overall increase in lifetime above room temperature shows that relaxation to the fundamental of the bend is an important pathway. However, the similar lifetimes at room temperature and below suggests that multiple relaxation pathways with different temperature dependencies may be in competition. The theoretical study, which was performed at room temper-
Figure 4-10: $T_1$ and $T^*$ for $\omega_{OD}$ of HOD in H$_2$O with temperature from the fit model.

Figure 4-11: Linear absorption coefficient, $K$ ($=4\pi\nu k$), of H$_2$O (1, black) and D$_2$O (2, red) from [2]. Copyright 2009 American Institute of Physics.

ature, considered the rate constants for energy transfer from $\omega_{OD}$ to $\delta_{HOD}$, $\delta_{H_2O}$, and $\omega_{H_2O}$ with excess energy taken up or contributed by the rotations and translations [106]; but it did not consider a pathway through $2\delta_{HOD}$. The energy gap between $\omega_{OD}$ and $\delta_{HOD}$ is more than 2x greater than $\omega_{OD}$ and $2\delta_{HOD}$ at all temperatures and although the energy gap between $\omega_{OD}$ and $\delta_{HOD}$ increases by $\sim$5%, it decreases by $\sim$14% (or $\sim$11% at FWHM) between $\omega_{OD}$ and $2\delta_{HOD}$. The broad librational band of H$_2$O, which is peaked at $\sim$680 cm$^{-1}$, or a combination of low frequency modes are
capable of contributing the excess energy. In Figure 4-11 we show the linear absorption spectrum of H$_2$O showing the continuous absorption from 0 to 2000+ cm$^{-1}$. The overtone of the bend is just one suggestion for a competitive pathway that may cause the non-trivial temperature dependence. An experimental study monitoring the deposition of energy into the HOD bend upon excitation of the OD stretch at multiple temperatures and a temperature dependent theoretical study of the vibrational lifetime would be helpful in determining if there is a change in dominant mechanism with temperature.

4.4 Thermally Shifted Ground State in Pump-Probe Measurements

Non-equilibrium heating upon vibrational relaxation in water was first observed as a blue-shift in frequency and decrease in absorption cross section of the OH stretch in pump-probe (PP) experiments of HOD in D$_2$O [175]. Energy deposited in the stretch decays and eventually populates the low frequency modes of the originally excited molecule and surrounding solvent. The excitation of these modes appears as heating since the fluctuations of the hydrogen bond network increase, which decreases the degree of hydrogen bonding. The "thermally shifted ground state" (TSGS) prevents the equilibrium ground state hole from refilling, which gives rise to a bleach that persists until thermal diffusion out of the spectral window. Since the signal arising from the TSGS and equilibrium ground state are out of phase, at waiting times well past the vibrational lifetime the dispersed pump-probe resembles a difference spectrum between linear spectra separated by ~1 K [72].

In Figure 4-12 we explore the effect of the TSGS on transient absorption measurements. The dispersed pump probes are calculated for the OD stretch of HOD in H$_2$O with a response function formalism we will describe in detail in the next chapter. In the current calculation the frequency correlation function is real, which excludes any effect due to the vibrational Stokes shift. In the calculation, the peak of the funda-
Figure 4-12: DPP of $\omega_{OD}$ without (a) and with (b) the TSGS. Slices in (c) frequency and time (d) without (solid) and with (dashed) the TSGS.

The anharmonic is 2508 cm$^{-1}$, the line width is 166 cm$^{-1}$ (FWHM), and the anharmonicity, $\omega_{10}-\omega_{21}=\Delta$, is 162 cm$^{-1}$ [77]. The lifetime of the initially excited state is $T_1 \approx 1.45$ ps and the TSGS contribution grows in as a bi-exponential with the time scales $T_1$ and $T^* \approx 540$ fs, where $T^*$ is the lifetime of an intermediate state (see Equation 4.2). In Figure 4-12a we show the dispersed pump-probe without including the TSGS. The positive amplitude is from the ground state bleach and stimulated emission of the $\nu=1 \leftrightarrow 0$ transition. The anharmonically shifted negative amplitude is due to the induced absorption of the $\nu=2 \leftrightarrow 1$ transition. We include the TSGS for the DPP calculation shown in Figure 4-12b. The bleach persists past the vibrational lifetime and negative amplitude appears above the bleach. In Figure 4-12c and Figure 4-12d we compare slices in frequency and time, respectively, between the calculated DPPs without (solid) and with (dashed) the TSGS. Figure 4-12c shows that if the TSGS is not included all three frequency slices decay to zero. However, when the TSGS is
included the frequency slices that begin negative can decay to a positive offset (red) and vice versa (blue). Figure 4-12d shows that if the TSGS is not included the zero crossing between the $\nu=1\rightarrow 0$ and $\nu=2\rightarrow 1$ peaks does not shift with waiting time. At t=0 fs there is no difference when the TSGS is included in the calculation (the dashed blue lies on top of solid blue). However, at longer waiting times the signal persists past the vibrational lifetime, positive amplitude shifts towards lower frequency, and the DPP dips to negative values on the high-frequency side of the line width.

4.5 Conclusion

In this chapter we studied the vibrational relaxation of isotopically dilute water. At room temperature, the lifetime of $\omega_{OH}$ is 700-740 fs [93, 98] and the lifetime of $\omega_{OD}$ is 1.45 to 1.7 ps [98, 95, 100]. We measured the dispersed pump probe of $\omega_{OT}$ in the zzzz polarization geometry and found a decay time of $\sim$850 fs. Therefore we find $T_{1,OD} > T_{1,OT} > T_{1,OH}$ which agrees with the proposed mechanism for vibrational relaxation, in which the stretch decays through the closest intramolecular mode that is lower in energy, since $\omega_{OD} - \delta_{HOD} (\sim 1050 \text{ cm}^{-1}) > \omega_{OT} - \delta_{HOT} (\sim 740 \text{ cm}^{-1}) > \omega_{OH} - 2\delta_{HOD} (\sim 500 \text{ cm}^{-1})$. We also presented the temperature dependence of the linear absorption of HOD in H$_2$O and $T_{1,OD}$ as a step in testing the temperature dependence of the proposed mechanism. We found that although the linear absorption changes monotonically with temperature, $T_{1,OD}$ shows a non-trivial behavior. At room temperature and below, the lifetime is the same within error and then it increases above room temperature. Since the energy gap between $\omega_{OD}$ and $\delta_{HOD}$ increases with increasing temperature the behavior above room temperature suggests that relaxation to the fundamental of the bend is an important pathway. However the behavior at room temperature and below suggests that there may be additional competing pathways that have different dependencies on temperature. Broadband experiments and further calculations are still necessary to definitely elucidate the vibrational relaxation pathway in isotopically dilute water.
Chapter 5

Hydrogen Bond Rearrangements in Water

5.1 Introduction

Despite years of extensive study and continuous improvements in experimental and theoretical methods, the femtosecond-to-picosecond dynamics that describe changes to the hydrogen bonding structure of water continue to be difficult to describe. Being a dense medium, the liquid intermolecular dynamics involve the collective motion of large numbers of molecules, for which translational and rotational degrees of freedom are strongly intertwined. This complexity is heightened in the case of the water due to its locally structured hydrogen bond network. The details of structural relaxation in water have been studied by molecular dynamics (MD) simulations beginning with the original work of Rahman and Stillinger [176]. Intermolecular motions and the interconversion between structures have been analyzed in a number of ways including through groups, chains, polyhedra, clathrate structures, inherent structures, and instantaneous normal modes [177, 28]. No correlations between local structural variables, such as hydrogen bond participation, exist between configurational states separated by more than a few picoseconds [13]. Therefore, the dynamics of structural relaxation processes are not readily described in terms of structure on molecular length scales.
Recent experimental and theoretical studies of water have provided new perspectives on the local dynamics of hydrogen bonding, describing in detail how the hydrogen bond distances and angular motions change during the course of collective structural reorganizations [71, 1, 178]. With ultrafast infrared spectroscopy and MD simulations Eaves et al. observed that water molecules in broken or strained hydrogen bond configurations return to a stable hydrogen bond within the fastest intermolecular motions suggesting that hydrogen bond switching is a concerted process [71]. Laage and Hynes performed MD simulations and found the dominant mechanism of reorientation is through large amplitude angular jumps with a contribution from diffusive reorientation of the intermolecular O...O framework [1]. In these studies, molecular motions that have been identified as important for reorganization include concerted hydrogen bond switching, large angle jumps, translation within the first and second solvation shells, and diffusive reorientation of a hydrogen bonded pair of water molecules.

It is important to recognize that local geometric variables are just different parameters that capture the motion along a collective reaction coordinate. Different motions may be projected from the collective dynamics and certain coordinates may project more favorably than others. However, long time relaxation phenomena of internal variables should all decay in a correlated fashion [68]. Absolute time scales from different experiments are difficult to compare since the observables are uniquely sensitive to relaxation, yet the temperature dependence should be identical if rearrangements occur along a collective reaction coordinate.

In this chapter we report on the temperature dependence of spectral diffusion and reorientation of the OD stretch of dilute HOD in H₂O to determine if the relaxation of these different measures of water structural change are correlated. In order to quantify the temperature-dependent time scales we self-consistently model the pump-probe anisotropy, vibrational relaxation, and spectral diffusion with a response function formalism that includes the solvent H₂O stretch and combination band and the thermally shifted ground state (TSGS) of all three modes. We find a similar barrier height for both spectral diffusion and reorientation along with a number of
other measurements that are sensitive to collective and single molecule relaxation of water. We also observe a similar behavior for spectral diffusion and reorientation calculated with MD simulations with the SPC/E [31] and SPC/FQ [88] model of water. Finally, we find the frequency dependence of the pump-probe and 2D anisotropy suggests that water molecules in broken or strained hydrogen bond configurations reorient more quickly than band center to form a stable hydrogen bond.

5.2 Experimental Methods

Temperature-dependent Fourier transform infrared (FTIR) absorption spectra of dilute HOD in H₂O were collected at 1 cm⁻¹ resolution with a Nicolet 380 FTIR spectrometer from Thermo-Electron Corporation. The sample was contained between 1 mm thick CaF₂ windows with a path length of 6 μm set by a Teflon spacer for both linear and nonlinear measurements. The temperature of the home built brass sample cell was controlled with a water-cooled chiller, Neslab RTE-7.

Sub-100 fs infrared pulses were generated with a home-built system consisting of an amplified Ti:sapphire pulse (800 nm, 1 kHz, 1 W) pumping an optical parametric amplifier (OPA). The system was described in detail in Chapter 2. Here we will briefly mention the salient features. A small portion of 800 nm (~1%) creates white light in a sapphire plate. After the white light seed and a portion of the 800 nm (~9%) pass through a 3 mm β-Barium Borate (BBO) crystal, the idler (2 μm) seeds a second pass through the BBO with the remaining 800 nm. Mid-infrared light is created by difference frequency generation of the signal (1.33 μm) and idler in a 0.5 mm AgGaS₂ crystal. The spectrum was centered at 2500 cm⁻¹ with a FWHM of ~230 cm⁻¹. Although the infrared pulse was contained in a purged box there was still residual CO₂ creating a dip in the spectrum. The pulses were compressed to ~70 fs for the 2D measurements and ~80 fs for the pump-probe (PP) measurements. Cross phase modulation (XPM) (described in Chapter 2) obscures spectral information within approximately two times the pulse width. Pulse delays were controlled with a pair of anti-reflection (AR) coated ZnSe wedges mounted to stages that translate the wedge
face perpendicular to the beam path. The ZnSe wedges are 6 cm in length and have a 1.2° wedge.

We measured temperature-dependent two-dimensional infrared (2D IR) surfaces in the zzzz polarization geometry to characterize spectral diffusion. We used KBr beam splitters to split the mid-infrared pulse into three parts to acquire surfaces in the pump-probe geometry in which a collinear pulse pair creates a vibrational excitation, and the third pulse acts as the probe and local oscillator [132]. The measured signal is the real part of the rephasing plus non-rephasing pathways [140]. The $\tau_1$-axis was created by step-scanning the non-chopped pump beam in 2 fs steps to 600 fs. Scanning the non-chopped pump beam removes the contribution to the total signal of the $\tau_1$-dependent PP between the probe and one of the pump beams. The PP between the probe and the other pump beam is independent of $\tau_1$ and is, therefore, removed by numerical Fourier transform to create the excitation, or $\omega_1$, dimension. The probe beam is spectrally dispersed after the sample with a 75-grooves/mm grating onto a 64-channel liquid nitrogen cooled MCT array. The $\sim 6 \text{ cm}^{-1}$ resolution in $\omega_3$ gives a detection axis with 400 cm$^{-1}$ bandwidth. In addition to the acquisition of the 2D signal, the interferometric autocorrelation and interference fringes for stage calibration are simultaneously measured from the replica of the collinear pump pair from the backside of the recombining beam splitter [132]. A small reflection is directed into a room temperature MCT to measure the interference between the pulses that constrains the relative pulse timing to the step size. The remainder is passed through a monochromater set to 2500 cm$^{-1}$ with a resolution of 1 cm$^{-1}$, and the interference is collected with a single channel MCT detector and Fourier transformed to determine the correction to the $\tau_1$-axis [132].

We used polarization-selective transient absorption to study temperature-dependent reorientational dynamics and vibrational relaxation. In the pump-probe geometry a pump-probe can be measured simply by blocking the non-chopped pump beam. A MgF$_2$ $\lambda/2$-waveplate (Alphalas) is placed in the pump path to rotate the polarization 45° relative to the probe. The polarization of the pump and probe were selected with ZnSe wire-grid polarizers (Molelectron). There are two polarizers after the sample.
The analyzing polarizer is rotated between parallel (zzzz) and perpendicular (zzyy) relative to the pump. The second polarizer is set parallel to the probe such that the polarization-sensitive monochromator grating sees the same polarization for zzzz and zzyy measurements. The pump-probe measurements were acquired by fast scanning the delay between the pump and probe and averaging the scans. The isotropic signal, which is free of rotations and therefore measures vibrational relaxation, is constructed from the zzzz and zzyy measurements as:

\[
S_{iso}(\tau_2) = \frac{S_{zzzz}(\tau_2) + 2S_{zzyy}(\tau_2)}{3}.
\] (5.1)

The anisotropy, which measures reorientational dynamics, is constructed as:

\[
C_R(\tau_2) = \frac{S_{zzzz}(\tau_2) - S_{zzyy}(\tau_2)}{S_{zzzz}(\tau_2) + 2S_{zzyy}(\tau_2)}.
\] (5.2)

### 5.3 Experimental Results

Temperature-dependent linear absorption of the OD stretch of HOD in H\textsubscript{2}O at the five temperatures in this study (278, 286, 295, 323, and 345 K) with the background H\textsubscript{2}O subtracted are shown in Figure 5-1a and without subtraction in Figure 5-1b. The condensed phase spectra red shifts by \(\sim 215\) cm\(^{-1}\) from the gas phase (2723.68 cm\(^{-1}\) [159]) due to hydrogen bonding. The \(\sim 160\) cm\(^{-1}\) full-width half maximum (FWHM) primarily reflects a distribution of hydrogen bond strengths [155]. From 278 to 345 K the peak blue shifts from 2499 to 2534 cm\(^{-1}\) (0.53 cm\(^{-1}\)/K), indicating an increase in hydrogen bond fluctuations, and the FWHM broadens from 159 to 180 cm\(^{-1}\). However, the total area decreases with increasing temperature due to the non-Condon effect [156, 7]. In the same temperature range the HOD bend, which can be seen in Figure 5-1b, narrows and red shifts from 1460 to 1450 cm\(^{-1}\). The model calculations, which will be described below, include the combination band \((\nu_L + \delta)\) and stretch of the H\textsubscript{2}O bath. From 278 to 345 K the H\textsubscript{2}O stretch red shifts from 3389 to 3433 cm\(^{-1}\) (0.66 cm\(^{-1}\)/K) and the combination band blue shifts from 2134 to 2075 cm\(^{-1}\) with a weak quadratic dependence.
Figure 5-1: Temperature-dependent linear absorption of dilute HOD in H$_2$O with (a) and without (b) the solvent H$_2$O subtracted. The traces are offset along the absorbance axis. The fit from our spectral model at each temperature is shown in dark gray.

Isotropic transient absorption at each of the five temperatures is shown in Figure 5-2a. The temperature dependence of the lifetime is discussed in detail in Chapter 4. However, the important details will be mentioned here. The isotropic PP is calculated from the zzzz and zzyy transient absorption measurements with Equation 5.1. The dispersed PP is integrated over 75\% of the maximum in frequency (of a slice taken at $\tau_2 = 200$ fs to avoid XPM) to take into account the change in center frequency with temperature. Thermalization of the low frequency modes occurs upon vibrational relaxation of the OD stretch. This causes a blue shift in the frequency of the stretch that prevents refilling of the ground state bleach. At long waiting times the thermally shifted ground state (TSGS) appears as a difference spectrum between linear spectra separated by $\sim 1$ K [72]. The TSGS has been included in a fit model by Rezus and Bakker [120] that also includes relaxation through an intermediate state. The fit model does not consider effects from the vibrational Stokes shift. The experimental vibrational lifetimes found by applying the fit model starting at $\tau_2 = 400$ fs for each of the five temperatures are shown in Figure 5-3.

Normalized temperature-dependent pump-probe anisotropies are shown in Figure 5-2b. Before calculating the pump-probe anisotropy, the bi-exponential growth
Figure 5-2: (a) Isotropic transient absorption (offset) and (b) anisotropy (normalized at 200 fs) of dilute HOD in H$_2$O at each of the five experimental temperatures plotted as a function of delay time between the pump and probe. The best fit from our spectral model normalized at 200 fs for both (a) and (b) is shown in dark gray. (c) Phase line slope over $\omega_1 = \omega_3 = 2470$ to $2530$ cm$^{-1}$ determined from the temperature and waiting time dependent 2D IR with the best fit from our model shown in gray. The experimental phase line slope and fits are offset by 0.075 per temperature as the temperature is decreased from 345 K.
of the TSGS, determined by fitting the isotropic PP, is subtracted from the zzzz and zzyy transient absorption measurements. This approach assumes the TSGS does not depend on the polarization, which has been shown to be a reasonable assumption for the OD stretch [119]. The dispersed anisotropies are integrated over a frequency range set by the maximum in frequency of a slice in the isotropic signal at \( \tau_2 = 200 \) fs. Although XPM obscures spectral information within the first \( \sim 150 \) fs, we fit the anisotropies with a bi-exponential beginning at 200 fs. A bi-exponential decay has previously been observed for the anisotropy decay of the OH stretch of HOD in D
\textsubscript{2}O measured with shorter pulses [93]. The inertial decay is set to 50 fs for all temperatures, the inertial time scale measured for the OH stretch of HOD in D
\textsubscript{2}O at room temperature [93], and the initial value is set to 0.4. We do not observe a clear trend between the inertial amplitude and temperature in the bi-exponential fits. MD simulations predict the inertial amplitude will increase with increasing temperature but experiments have reported that there is a change in frequency dependence of the inertial component with temperature, which complicates a straightforward relationship [179, 121]. The long time decay of the anisotropy, shown in Figure 5-3, becomes faster with increasing temperature. We found very similar values for the long-time decay when fitting a single exponential beginning at 200 fs without constraining the initial value.

2D IR surfaces at each of the five temperatures at three waiting times are shown in Figure 5-4. Since multiple vibrational levels are accessible within the bandwidth of our pulse, the 2D surface displays a positive peak from ground state bleach (\( \nu=1\leftarrow 0 \)) and an anharmonically shifted negative peak from excited state absorption (\( \nu=2\leftarrow 1 \)). At early waiting times (\( \tau_2 < \tau_C \), where \( \tau_C \) represents the correlation time), the peaks are diagonally elongated indicating frequency correlation between the first and third time periods. In the top row of Figure 5-4 we show 2D surfaces at \( \tau_2 = 160 \) fs for each of the five temperatures. Notice at 278 K the pear shape of the positive peak where the anti-diagonal line width is narrow on the red side and relatively broad on the blue side (where red and blue refer to low and high frequencies relative to band center, respectively). The asymmetry across the line width forms the basis of a previous
Figure 5-3: Temperature-dependent lifetime, \( T_1 \), (red triangles), picosecond decay of the integrated pump-probe anisotropy (green squares) and picosecond decay of the best fit frequency correlation function from the fit model (blue circles).

Publication in which we analyzed the waiting time dependence of the line shape for the OH stretch of dilute HOD in D\(_2\)O and found that strained or broken hydrogen bonds exist only fleetingly [71, 73]. Note that as the temperature is increased, but as the waiting time is kept constant, the asymmetry across the line shape decreases and the anti-diagonal line width becomes generally broader.

We also plot the phase representation of the surfaces. We detail how we extract the complex surface from the measured real surface in Chapter 2. At early waiting times the phase lines are tilted toward the diagonal. At \( \tau_2 = 400 \) fs (second row of Figure 5-4) the frequencies initially excited have more time to diffuse through the line shape. As a result the peaks are more symmetric, or appear more homogeneous, and the phase lines have tilted toward \( \omega_1 \) at all temperatures. Notice, however, at low temperature the line shape is more inhomogeneous relative to high temperature. At long waiting times (\( \tau_2 > \tau_C \)), sufficient spectral diffusion occurs during the waiting time such that peaks appear round due to loss of correlation. Also, an additional negative feature appears above the diagonal upon sufficient vibrational relaxation due to the TSGS. In the third row of Figure 5-4 we show the temperature-dependent 2D surfaces at
3.2 ps. Note the surfaces are generally round and there is additional negative feature above the diagonal, particularly visible for the three lowest temperatures, that acts to narrow the positive amplitude along $\omega_3$. Finally, note that there appears to be a residual inhomogeneity, especially apparent at higher temperatures in the phase representation, on the red side of the spectrum.

We used a number of metrics to characterize the waiting time dependence of spectral diffusion [141], including the dynamic line width [77], slope of the node [180, 72], and the center line slope [181]. The dynamic line width (DLW) is the line width (FWHM) of a slice along $\omega_1$ at the maximum in $\omega_3$. The DLW is shown at 278, 295, and 345 K for one data set in Figure 5-5. Due to inhomogeneous broadening the FWHM is narrowed when $\tau_2 < \tau_C$ and broadens towards the asymptotic value, set either by the line width of the absorption band or the bandwidth of the pulse, as the waiting time is increased.

The slope of the node between the $\nu=1\leftarrow 0$ and $\nu=2\leftarrow 1$ peaks of the absorptive 2D spectrum is shown in Figure 5-6 at the three temperatures for the same data set. The slope decreases as the waiting time is increased, however, the slope of the node will only be well defined if the $\nu=2\leftarrow 1$ has sufficient amplitude. If the node is ill-defined the slope will be ill-defined, which can be seen in the trace for the highest temperature (345 K).

The center line slope (CLS) for the three temperatures is shown in Figure 5-7. The CLS is similar to the slope of the node except it is the slope of a line connecting the maxima along $\omega_1$ of the $\nu=1\leftarrow 0$ peak. It is an attractive metric in that it can still be used after the $\nu=2\leftarrow 1$ peak has decayed, however, the maximum of the peak is not as well defined as the node, in that it is gradually sloping.

After retrieving the complex correlation spectrum and the rephasing and non-rephasing surfaces on the basis of the Kramers-Kronig assumption (see Chapter 2) we also considered the normalized difference of the amplitude of the rephasing to non-rephasing surfaces, or inhomogeneity index [141, 72], the slope of the imaginary node, and the phase line slope [182]. In Figure 5-8 we show the inhomogeneity index, $I$, defined as $I = (A_-A_+)/A_+ + A_+$, where $A_+$ is the integral over the non-
Figure 5-4: Experimental absorptive and phase 2D surfaces of dilute HOD in H2O at each of the five temperatures at three waiting times: \( \tau_2 = 160 \) fs (top), 400 fs (middle) and 3.2 ps (bottom). The contours in the absorptive 2DIR spectra are at 12% intervals relative to the maximum. The phase spectra are plotted from \(-\pi/2\) to \(\pi/2\) and are windowed by 10% of the absolute value spectra.

rephasing and rephasing surfaces, respectively. Frequency memory is reflected in unequal weighting between the rephasing and non-rephasing pathways, with more amplitude in the rephasing pathway. Therefore, as the waiting time is increased, and frequency memory is lost, the inhomogeneity index will decrease.
Figure 5-5: Dynamic line width (DLW) for time ($\tau_2 = 40$ fs to 4 ps) and temperature-dependent (278, 295, and 345 K) 2D surfaces of the OD stretch of HOD in H$_2$O (right). 2D surfaces at 295 K and $\tau_2 = 200$ fs (top) and 2 ps (bottom) with the DLW marked in cyan (left).

Figure 5-6: Slope of the node (Real) for time ($\tau_2 = 40$ fs to 4 ps) and temperature-dependent (278, 295, and 345 K) absorptive 2D surfaces of the OD stretch of HOD in H$_2$O (right). 2D surfaces at 295 K and $\tau_2 = 200$ fs (top) and 2 ps (bottom) with the node marked in cyan (left).

We show the slope of the $\phi=0$ phase line, which is equivalent to the slope of the node of the imaginary surface for the three temperatures in Figure 5-9. Again, this metric is similar to the slope of the node of the real surface and CLS, but it is
superior in that it is viable at all waiting times and the node is better defined than the maximum of the peak.

Finally, we show the phase line slope (PLS) for the three temperatures in Figure 5-10. The PLS is the average of the slopes of the phase lines contained within a box with
Figure 5-9: Slope of the $\phi=0$ phase line, or node of the imaginary surface, for time ($\tau_2 = 40$ fs to 4 ps) and temperature-dependent (278, 295, and 345 K) 2D surfaces of the OD stretch of HOD in H$_2$O (right). 2D phase surfaces at 295 K and $\tau_2 = 200$ fs (top) and 2 ps (bottom) with the $\phi=0$ phase line marked in cyan (left).

Figure 5-10: Phase line slope (PLS) for time ($\tau_2 = 40$ fs to 4 ps) and temperature-dependent (278, 295, and 345 K) 2D surfaces of the OD stretch of HOD in H$_2$O (right). 2D phase surfaces at 295 K and $\tau_2 = 200$ fs (top) and 2 ps (bottom) with $\omega_1 = \omega_3 = 2470$ to 2530 cm$^{-1}$ marked in magenta (left).

the bounds set to $\omega_1 = \omega_3 = 2470$ to 2530 cm$^{-1}$, although the range is arbitrary. For instance, one can study the heterogeneity of spectral diffusion by selecting different frequency regions within the line shape. This is our preferred metric since it provides superior signal to noise.
All metrics generally show the same trend between temperatures in that the decay is faster with increasing temperature yet there is an offset apparent at longer waiting times that is larger with higher temperatures. We compare the normalized metrics at 295 K in Figure 5-11 (normalized at the first recorded waiting time, $\tau_2 = 40$ fs). For the DLW we subtract the last value before normalization. Ideally we would subtract that asymptotic value, so it is not surprising that this metric varies the most from the other metrics. Upon comparison, we find the CLS is the noisiest metric. In Figure 5-2c we show our preferred metric, the phase line slope, $\tan(\phi)$, over the same frequencies as Figure 5-10 for the five temperatures in this study averaged over multiple data sets. We also determined the phase line slope within a box with the bounds set to 90% of the maximum in $\omega_1$ and $\omega_3$ for each surface [182]. Both treatments show a very similar behavior. The phase line slope decreases with increasing temperature. However, there is an offset that appears larger with increasing temperature.

![Figure 5-11: Comparison of the metrics at room temperature normalized at the initial waiting time, $\tau_2 = 40$ fs.](image-url)
5.4 Spectroscopic Model

In order to quantify the temperature-dependence of spectral diffusion, reorientational dynamics, and vibrational relaxation, we employ a widely used nonlinear response function formalism to self-consistently model the FTIR, PP and 2D IR measurements [70]. The model is similar to one we used to study the OH stretch of HOD in D$_2$O [69]. The response function, $S$, describes the vibrational dynamics of OD dephasing ($S_\omega$), vibrational population relaxation ($S_{\text{pop}}$), and reorientation ($Y$) as separable contributions to the vibronic response function, in addition to non-equilibrium heating effects that arise from vibrational relaxation of the OD stretch:

$$S = Y(S_\omega S_{\text{pop},\omega} + S_{\text{TSGS}} S_{\text{pop},TSGS}).$$

Dephasing is treated as fluctuations of the $\nu=0, 1,$ and 2 system eigenstates that result from coupling to a harmonic bath characterized by a bath time-correlation function, $C_\omega(t)$. The remaining components are treated as phenomenological rate processes.

This model makes several approximations that assume the various dynamics can be treated as homogeneous processes. We use the Condon and second-order cumulant approximations [70, 183], which state that the transition dipole moment is independent of bath, and that the frequency fluctuations are purely Gaussian. Both approximations have been shown to be invalid for water on time scales short compared to spectral diffusion [156, 7, 73, 78, 184]. Rotation and vibrational relaxation are treated as Markovian processes that are independent of vibrational dephasing, although evidence indicates that this is a poor assumption for the same short time scales. We proceed with this analysis since the emphasis of this study is on relaxation kinetics on time-scales longer than the correlation time for spectral diffusion. On these long time-scales structural reorganization of the hydrogen bond network has effectively scrambled correlations among the different relaxation processes.

The timescale on which the assumptions become valid can be deduced from an analysis of heterogeneity within the 2D IR line shape. In Figure 5-12a we show the waiting time-dependence of the first moment of the positive signal distribution for
slices in $\omega_1$ on the red and blue side of the line shape at 295 K [72]. The decrease in
the first moment on both sides of the line width at longer waiting times is due to the
growth of the TSGS. For $\tau_2 \leq 200$ fs, there is a significant difference between first
moment on the red and blue side, with a fast decrease on the blue side and a slight
recurrence at $\sim 120$ fs on the red side. A recurrence, or beat, has been observed in
previous PS and 2D IR measurements of the OH stretch [68] and MD simulations of
the OH and OD stretch (for certain water models) [86]. After approximately 500 fs
the dynamics become more similar across the line width. At 278 K (Figure 5-12b),
similar observations are made, but the difference across the line width is somewhat
larger. Also, the recurrence on the red side of the line width is more pronounced at the
lower temperature. These observations indicate that our model properly handles the
measurement of relaxation time-scales for waiting times longer than the OD frequency
correlation time.

Figure 5-12: First moment of the positive amplitude for slices in $\omega_1$ at 2454 cm$^{-1}$
(red) and 2551 cm$^{-1}$ (blue) at the temperatures 295 K (a) and 278 K (b). The insets
show the short time behavior for each of the two temperatures.

5.4.1 Vibrational Dephasing

To model the vibrational dephasing response function $S_{\omega}$, we include rephasing (-)
and non-rephasing (+) pathways that contribute to third-order nonlinear experiments
for pulses separated in time. Within the Condon approximation, and combining the
dehphasings and vibrational population relaxation $S_\omega S_{\text{pop}}$, the response functions are

$$S_-(\tau_3, \tau_2, \tau_1) = |\mu_{10}|^2 \exp\left[-\frac{\tau_3}{T_1}\right]\exp[i(\omega_{10})\tau_1 - \frac{\tau_1}{2T_1}]$$
$$\times \left\{ |\mu_{10}|^2 \exp[-i(\omega_{10})\tau_3 - \frac{\tau_3}{2T_1}]\left[F_{0101}^{(3)}(\tau_3, \tau_2, \tau_1) + F_{0101}^{(4)}(\tau_3, \tau_2, \tau_1)\right] \right. \\
- \left. |\mu_{21}|^2 \exp[-(\omega_{21})\tau_3 - \frac{3\tau_3}{2T_1}]\left[F_{0121}^{(2)}(\tau_3, \tau_2, \tau_1)\right]^*\right\}, \quad (5.4)$$

$$S_+(\tau_3, \tau_2, \tau_1) = |\mu_{10}|^2 \exp\left[-\frac{\tau_3}{T_1}\right]\exp[-i(\omega_{10})\tau_1 - \frac{\tau_1}{2T_1}]$$
$$\times \left\{ |\mu_{10}|^2 \exp[-i(\omega_{10})\tau_3 - \frac{\tau_3}{2T_1}]\left[F_{0101}^{(1)}(\tau_3, \tau_2, \tau_1) + F_{0101}^{(2)}(\tau_3, \tau_2, \tau_1)\right] \right. \\
- \left. |\mu_{21}|^2 \exp[-(\omega_{21})\tau_3 - \frac{3\tau_3}{2T_1}]\left[F_{0121}^{(4)}(\tau_3, \tau_2, \tau_1)\right]^*\right\}. \quad (5.5)$$

In these expressions $\omega_{10}, \omega_{21}$ and $\mu_{10}, \mu_{21}$ denote the transition frequency and tran-
sition dipole of the $\nu=1\rightarrow 0$ and $\nu=2\rightarrow 1$ transitions, respectively. The room tem-
perature anharmonicity is fixed at $\Delta = \omega_{10} - \omega_{21} = 162 \text{ cm}^{-1}$ for the OD stretch [77] and is taken to be independent of temperature. We assume harmonic scaling of the transition dipole, therefore $\mu_{21} = 2\mu_{10}$. $F_{abcd}$ are the dephasing functions, in which the indices refer to the vibrational 0, 1 and 2 states. $T_1$ is the vibrational lifetime, which is determined by fitting the isotropic transient absorption. We include population relaxation during the coherence periods with the timescale determined by: $\Gamma_{ab} = (1/2)(\Gamma_{aa} + \Gamma_{ba})$, where $T_1 = 1/\Gamma_{11}$. We assume the population relaxation scales with quantum number, $\Gamma_{aa} = a\Gamma_{11}$, therefore $T_1$ is the only necessary input.

Under the second-order cumulant approximation, the dephasing functions in Equa-
tion 5.5 can be expressed in terms of line shape functions, \( g \) \[138\]:

\[
-ln F^{(1)}_{0101}(\tau_3, \tau_2, \tau_1) = g_{11}(\tau_1) + g_{11}(\tau_2) + g_{11}(\tau_3) - g_{11}(\tau_1 + \tau_2)
- g_{11}(\tau_2 + \tau_3) + g_{11}(\tau_1 + \tau_2 + \tau_3),
\]

\[
-ln F^{(2)}_{0101}(\tau_3, \tau_2, \tau_1) = g_{11}(\tau_1) + g_{11}^*(\tau_2) + g_{11}^*(\tau_3) - g_{11}(\tau_1 + \tau_2)
- g_{11}(\tau_2 + \tau_3) + g_{11}(\tau_1 + \tau_2 + \tau_3),
\]

\[
-ln [F^{(3)}_{0101}(\tau_3, \tau_2, \tau_1)]^* = g_{11}(\tau_1) - g_{11}(\tau_2) + g_{11}(\tau_3) + g_{11}(\tau_1 + \tau_2)
+ g_{11}(\tau_2 + \tau_3) - g_{11}(\tau_1 + \tau_2 + \tau_3),
\]

\[
-ln [F^{(4)}_{0101}(\tau_3, \tau_2, \tau_1)]^* = g_{11}(\tau_1) - g_{11}^*(\tau_2) + g_{11}(\tau_3) + g_{11}(\tau_1 + \tau_2)
+ g_{11}^*(\tau_2 + \tau_3) - g_{11}(\tau_1 + \tau_2 + \tau_3),
\]

\[
-ln F^{(2)}_{0121}(\tau_3, \tau_2, \tau_1) = g_{11}(\tau_1) + g_{11}^*(\tau_2) + g_{11}^*(\tau_3) - g_{11}(\tau_2 + \tau_3)
- g_{11}(\tau_1 + \tau_2) + g_{11}(\tau_1 + \tau_2 + \tau_3) - g_{11}(\tau_2 + \tau_3)
+ g_{11}(\tau_1 + \tau_2 + \tau_3) - g_{11}(\tau_1 + \tau_2 + \tau_3),
\]

\[
-ln [F^{(4)}_{0121}(\tau_3, \tau_2, \tau_1)]^* = g_{11}(\tau_1) - g_{11}^*(\tau_2) + g_{11}^*(\tau_2) + g_{11}(\tau_3) - g_{12}(\tau_3)
- g_{21}(\tau_3) + g_{22}(\tau_3) + g_{11}(\tau_1 + \tau_2) - g_{21}(\tau_1 + \tau_2) - g_{11}(\tau_1 + \tau_2 + \tau_3)
+ g_{11}(\tau_1 + \tau_2 + \tau_3) - g_{12}(\tau_2 + \tau_3).
\]

The line shape functions are defined in terms of two-point frequency correlation functions:

\[
g_{pq}(t) = \int_0^t d\tau \int_0^{\tau} d\tau' C_{pq}(\tau - \tau')
\]

where

\[
C_{pq}(\tau - \tau') = \langle \delta \omega_{p0}(\tau) \delta \omega_{q0}(\tau') \rangle.
\]

The indices \( p \) and \( q \) refer to OD vibrational energy levels. In practice, we define one correlation function \( C_\omega(t) = C_{11}(t) \), and use harmonic scaling relations to relate correlation function for transitions involving \( \nu = 0, 1, \) and 2: \( C_{22}(t) = 2C_{21}(t) = \ldots \)
Motivated by a number of observations, we use a complex frequency correlation function in the present work [70]. First, librational motion plays an important role in the short time dynamics, and it is unclear if the high-temperature limit applies to librational frequencies. Also, harmonic bath models make explicit predictions about the temperature dependence of spectroscopic observables; for instance, the absorption line width is linear in T in the high temperature limit. Also, the Stokes shift, which accounts for the response of the bath to OD excitation, has been predicted to be small but a relevant contribution to the spectroscopy. A complex exponential correlation function can be used to properly account for these effects. Computational modeling of HOD in H$_2$O indicated that a classical correlation function is satisfactory for linear measurements, but quantum effects may become important for nonlinear measurements, especially for strongly inhomogeneously broadened systems [185]. Overall, the analysis with a complex correlation function did not differ greatly from a real correlation function, but it does identify interesting aspects of the data that are discussed below and in the following sections.

Similar to prior studies of HOD in H$_2$O by Fayer and co-workers [77], a tri-exponential was chosen as the functional form of the frequency correlation function [70]:

$$C_\omega(t) = \sum_{j=1}^{3} A_j \left[ \frac{1}{\tau_{C,j}} \coth(\frac{\hbar}{2kT\tau_{C,j}}) - i \frac{1}{\tau_{C,j}} \right] \exp\left( -\frac{t}{\tau_{C,j}} \right)$$

(5.9)

In this equation $\tau_{C,j}$ is the correlation time for the $j$th exponential relaxation component and $A_j$ is its amplitude. For this model, the reorganization energy is $\lambda = \sum_j A_j$, and $2\lambda$ is the Stokes shift. For a correlation function comprised of exponentials an approximation for the line shape function can be written as [70]:

$$g(t) = \sum_j A_j [\tau_{C,j} \coth(\frac{\hbar}{2kT\tau_{C,j}}) - i\tau_{C,j}] \exp\left( -\frac{t}{\tau_{C,j}} \right) + \frac{t}{\tau_{C,j}} - 1.$$  

(5.10)

Simulations discussed below show that the inertial decay of the frequency corre-
lation function has a weak temperature dependence. Therefore, we fixed the inertial 
decay time scale at $\tau_{C,1} = 50$ fs for all five temperatures, however, we found it nec-
essary to vary the amplitude to match experiment. To add constraints, we initially 
attempted to keep the intermediate timescale constant at $\tau_{C,2} = 400$ fs, the inter-
mediate timescale found by Fayer and co-workers [77], but we found the fit of the 
two highest temperatures, 323 and 345 K, unsatisfactory. At these temperatures the 
intermediate timescale was set to 200 fs. Therefore, only three parameters of the 
frequency correlation function are independent: the timescale of the long-time decay, 
$\tau_{C,3}$, and the amplitudes of the inertial and intermediate-time decays. A recurrence 
was not included in the correlation function since it is a relatively minor effect and 
we are primarily interested in the long-time behavior. In addition to the OD stretch 
of HOD, we also use this formalism to treat the weak background absorption from 
the H$_2$O stretch and libration-bend combination band (see Section S.1 in this chapter 
for details).

5.4.2 Rotational Response

For the case that vibrational and orientational response functions are separable and 
reorientational motion is treated as a Markovian process, the nonlinear orientational 
correlation functions can be written in terms of $C_{R,n} = \langle P_n[\hat{\mu}(t) \cdot \hat{\mu}(0)] \rangle$, where $P_n$ is 
the $n$th order Legendre polynomial and $\hat{\mu}$ is the transition dipole unit vector [186]. 
The orientational responses are then expressed as:

$$Y_{zzzz}(\tau_3, \tau_2, \tau_1) = \frac{1}{9} C_{R,1}(\tau_1)[1 + \frac{4}{5} C_{R,2}(\tau_2)] C_{R,1}(\tau_3),$$

$$Y_{zzyy}(\tau_3, \tau_2, \tau_1) = \frac{1}{9} C_{R,1}(\tau_1)[1 - \frac{2}{5} C_{R,2}(\tau_2)] C_{R,1}(\tau_3).$$

(5.11)

For pump-probe measurements, the isotropic response $Y_{iso} = (Y_{zzzz} + 2Y_{zzyy})/3$ is ob-
served to be independent of orientational dynamics during $\tau_2$, whereas the anisotropy 
is proportional to the second order orientational correlation function: $C_R(\tau_2) = (2/5)C_{R,2}(\tau_2)$. 
Guided by observations from the MD simulations below and previous experimental 
results [93], we model the anisotropy and second rank rotational correlation function
as a biexponential:

\[ C_R(t, T) = (1 - A_R(T))exp(-\frac{t}{\tau_{\text{inertial}}}) + A_R(T)exp(-\frac{t}{\tau_R(T)}). \]  

(5.12)

An inertial reorientational response with a time-scale of \( \tau_{\text{inertial}} = 50 \) fs is taken to be independent of temperature, but the amplitude \( A_R \) and reorientational correlation time \( \tau_R \) are allowed to vary. For the purpose of describing 2D IR experiments, we include orientational relaxation during \( \tau_1 \) and \( \tau_3 \), which requires a first rank orientational correlation function, \( C_{R,1} \). We relate \( C_{R,1} \) and \( C_{R,2} \) by scaling their orientational correlation times, but leave the inertial time scale unchanged. Based on recent MD simulations of water [32], we use a scaling of \( \tau_{R,1}/\tau_{R,2} = 2.3 \), although results are found to be indistinguishable within the model from results using the value of 3.0 predicted for small-angle diffusive reorientation. The linear spectrum, 2D IR and pump-probes are sensitive to the scaling; the pump-probe anisotropy, however, is not sensitive to the scaling within the model.

### 5.4.3 Thermal Effects

The model includes a third order response from the TSGS for all three modes. The response can be written as:

\[ S_{\pm}^{\text{TSGS}} = A^{\text{TSGS}} | \mu_{10} |^4 N^{\text{TSGS}}(\tau_2)exp\left(\mp i(\omega_{10})\tau_1 - i(\omega_{10})\tau_3\right) \]

\[ \times exp\left( -g_{11}^{(1)}(\tau_1) - g_{11}(\tau_3)\right) [1 - | \Delta \mu_{10} |^2 exp(-i\Delta \omega \tau_3)]. \]  

(5.13)

In this equation \( \Delta \mu_{10} \) denotes the decrease in transition dipole strength, \( \Delta \mu_{10} = (\mu_{10} - \mu_{10}^{\text{TSGS}})/\mu_{10} \), and \( \Delta \omega \) the blue-shift in frequency, \( \Delta \omega = \omega_{10} - \omega_{10}^{\text{TSGS}} \), induced upon vibrational relaxation. This formalism assumes an uncorrelated TSGS [66, 69, 187], in which frequency memory is lost upon vibrational relaxation. Temperature-dependent linear absorption spectra show that the frequency of the OD stretch changes linearly with temperature. Therefore, for all temperatures, \( \Delta \omega \) is set to the frequency change for 1 K increase in temperature, 0.53 cm\(^{-1}\). \( A^{\text{TSGS}} \), the intensity of the thermally
induced response, and $\Delta \mu_{10}$ are determined by fitting both the integrated isotropic PP at $\tau_2 > 1.5$ ps and the 2D surface at $\tau_2 = 4$ ps projected along $\omega_3$.

$N^{TSGS}(\tau_2)$ represents the growth of the thermal response with waiting time determined by fitting the integrated isotropic PP measurements. The integrated isotropic PP is fit with the model developed by Bakker [120]:

$$N^{TSGS}(\tau_2) = 1 + \frac{1}{T_1 - T^*}[T^*\exp(-\tau_2/T^*) - T_1\exp(-\tau_2/T_1)].$$

In this equation $T^*$ represents the lifetime of an intermediate state, which has been suggested to be the fundamental of the HOD bend [100]. Yeremenko and co-workers found an important contribution arising from the solvent D$_2$O in experiments on the OH stretch of HOD in D$_2$O [67, 187]. However, this effect is small for absorptive 2D IR and PP measurements, and is more important for measurements sensitive to dispersive changes, such as photon echo and transient grating [187]. Thermally induced shifts are also included for the background H$_2$O stretch and combination band, although the effect is rather small in the region of the OD stretch (see Section S.1 in this chapter).

### 5.4.4 Modeling Results

To apply the model to the OD stretch of HOD in H$_2$O, we first fit the experimental integrated isotropic PP and PP anisotropy at each temperature to determine $T_1$, $N^{TSGS}(\tau_2)$, and $C_R(t)$. We make an initial guess for $C_\omega(t)$ and solve for $\lambda$ by fitting the line width of the linear absorption measurement. The isotropic PP and 2D IR surfaces are calculated and compared to experiment to determine $A^{TSGS}$ and $\Delta \mu_{10}$. The 2D surface is a two dimensional Fourier transform of $Y(S_\omega S_{pop,\omega} + S_{TSGS}S_{pop,TSGS})$. The complex surface is determined from the calculated 2D surfaces with the same method used for experimental 2D surfaces. The parameters for the frequency correlation function are adjusted at each temperature to best fit the experimental phase line slope over the same frequency range as the experiment. We allow for an arbitrary offset to the phase line slope generated from the model when comparing to the experimental
phase line slope. Although we are interested in the long time behavior we fit the entire waiting time range to obtain the most reliable information. A self-consistent fitting of all the data provides enough constraints to quantitatively extract the longest correlation times, lifetimes, and reorganization energy at any temperature.

The reorganization energy, $\lambda$, is found by fitting the FWHM of the linear spectrum. The "full" treatment of calculating the linear spectrum includes a complex frequency correlation function, reorientation, and vibrational relaxation. The absorption line width is calculated by:

$$\sigma(\omega) \propto |\mu_{10}|^2 \int_0^\infty dt \exp[i(\omega - \omega_{10})t - \frac{t}{2T_1} - g_{11}(t)]C_{R,1}(t).$$  \hspace{1cm} (5.15)

In Figure 5-13 we show the experimental FTIR of HOD in H$_2$O at 295 K, with the background H$_2$O subtracted, along with the results of the "full" treatment using the best-fit frequency correlation function. Since we are using the cumulant approximation, the model spectrum is Gaussian. With the same input $\omega_{10}$ and $C_\omega(0)$, we calculate the linear spectrum first without including reorientation, then additionally without including vibrational relaxation. The spectrum narrows from 166 to 152 cm$^{-1}$ when reorientation is not included and an additional 2 cm$^{-1}$ when vibrational relaxation is not included. Therefore, reorientation is more important but it only accounts for $\sim$8% of the overall width ($\sim$10% for reorientation + vibrational relaxation). We also show the result of only considering the real part of the frequency correlation function. The peak frequency is red-shifted by $\sim$3 cm$^{-1}$ at this temperature when using a complex correlation function since the imaginary part introduces a Stokes shift.

The linear spectra calculated with the model for each of the five experimental temperatures are shown as solid lines in Figure 5-1. The reorganization energy, $\lambda$, and the complex initial value of the frequency correlation function, $C_\omega(0)$, for each temperature are given in Table 5-1. We find that at a given temperature the magnitude of the red-shift was most sensitive to the amplitude of the inertial decay (an increased amplitude resulted in an increased red-shift). Experimental and input fre-
Figure 5-13: Experimental linear absorption of the OD stretch of HOD in H$_2$O with the background H$_2$O subtracted (dashed gray), along with spectra calculated with the spectral model using the "full" treatment (solid blue), using $C_\omega(t)$ with R(t) and $T_1$ (dot dashed green), using $C_\omega(t)$ without including R(t) (solid red), and using $C_\omega(t)$ without including R(t) or $T_1$ (dot dashed cyan)

The amplitude of the inertial decay increases from 41% to 71% from 278 to 345 K. The amplitude of the intermediate and long-time decays both decrease with increasing temperature. The long-time decay decreases with increasing temperature, which is in agreement with our previous study [182]. The experimental phase line slopes and best fits with the model, including the offset, are shown as solid lines in Figure 5-2c.

Temperature and waiting time-dependent 2D surfaces generated from the model with the best-fit frequency correlation functions are shown in Figure 5-14. The apparent asymmetry is in small part due to including the H$_2$O combination band and stretch but mainly due to windowing by the experimental pulse spectrum after the 2D surface is calculated. The 2D surfaces from the model are successful in capturing the average dynamics with waiting time and temperature. The model 2D IR surfaces agree with the experimental surfaces best at lower temperatures. As the temperature
Figure 5-14: Absorptive and phase 2D surfaces calculated with the model at each of the five experimental temperatures at three waiting times: $\tau_2 = 160$ fs (top), 400 fs (middle) and 3.2 ps (bottom). The contours in the absorptive 2DIR spectra are at 12% intervals relative to the maximum. The phase spectra are plotted from $-\pi/2$ to $\pi/2$ and are windowed by 10% of the absolute value spectra.
is increased the residual inhomogeneity on the red side of the line shape, which is not captured with the model, becomes more important.

![Diagram](image)

Figure 5-15: Absorptive and phase 2D surfaces calculated with the model at $\tau_2 = 2$ ps and 278 K with (a) the "full" treatment, (b) using $C'_\omega(t)$, (c) not including the TSGS, and (d) not including the background H$_2$O.

In order to understand the influence of a complex correlation function on the calculated time scales, we calculate the 2D surface and phase line slope with the best-fit correlation function at $\tau_2 = 2$ ps for four cases: (a) the full treatment, (b) including only the real part of the correlation function, (c) not including the TSGS, and (d) not including the response of the bath H$_2$O. Cases (b)-(d) are the same as (a) except for where noted (for case (b), however, the input $\omega_{10}$ was necessarily different). Figure 5-15 and Figure 5-16 show the 2D surface and phase line slope, respectively, for each of the four cases for 278 K (Figure 5-15) and 278 K and 345 K (Figure 5-16). We also show the phase line slope for the full treatment but without windowing the 2D surfaces by the pulse spectrum. As can be seen in both Figure 5-15 and 5-16 including the solvent H$_2$O has little to no effect on the calculated dynamics within this model. The contribution from the H$_2$O stretch may become more significant if the non-Condon effect were included. As can be seen in Figure 5-15b and 5-16 including the imaginary part of the correlation function makes a non-negligible difference with
a larger effect at low temperature. The effect of the TSGS, shown in Figure 5-15c and 5-16, is non-negligible and the contribution becomes more important with increased waiting time. Since the TSGS is uncorrelated, a homogeneously broadened doublet (negative above the diagonal) grows in with waiting time. Comparing Figure 5-15a and 5-15c shows the negative contribution from the TSGS above the diagonal acts to narrow the positive peak along $\omega_3$. The phase lines of a homogeneously broadened peak are parallel to $\omega_1$ indicating that $\omega_3$ is independent of $\omega_1$. As a result, including the TSGS acts to decrease the phase line slope, as can be seen in Figure 5-16. Finally, The effect of windowing by the experimental spectrum on the calculated dynamics is shown in Figure 5-16. The experimental spectrum makes a non-negligible difference on the dynamics, which as been observed before for the OD stretch of HOD in H$_2$O with similar experimental conditions [77].

![Figure 5-16](image)

Figure 5-16: Phase line slope calculated with the spectral model for (a) 278 K and (b) 345 K with the "full" treatment (blue), using $C'_w(t)$ (green), not including the TSGS (red), not including the background H$_2$O (dot dashed cyan), and not windowing by the experimental spectrum (magenta).

After determination of the best-fit frequency correlation function the dispersed isotropic pump-probe is calculated using the isotropic orientational response function. The integrated isotropic signal from the model is compared to the experimental traces in Figure 5-2a. In order to compare the model and experimental anisotropy the third order response, without including the TSGS response, is calculated with the zzzz and
zzzy orientational response functions. The comparison is shown in Figure 5-2b.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>278 K</th>
<th>286 K</th>
<th>295 K</th>
<th>323 K</th>
<th>345 K</th>
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<td>1.5</td>
<td>1.4</td>
<td>1.6</td>
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</tr>
<tr>
<td>(T^* \text{ [fs]})</td>
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<td>450</td>
<td>540</td>
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<td>590</td>
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<td>(C_R(t))</td>
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<td>((0.64) \times 10^{-7})</td>
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<td>((A_1/\lambda) \tau_1 \text{ [fs]})</td>
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</tbody>
</table>

Table 5.1: Summary of the temperature dependent parameters contained within our spectral model for the OD stretch of HOD in H\(_2\)O.

### 5.5 Discussion

Over the past decade, the combination of ultrafast infrared spectroscopy and MD simulations of isotopically dilute water have developed an increasingly detailed molecular picture of the structural dynamics of water on the femtosecond to picosecond time scale. Since the frequency at which water absorbs is sensitive to its local hydrogen bonding environment, monitoring frequency evolution provides a picture of the evolving structure. On the picosecond time scale, theoretical work has shown that a number of relaxation mechanisms, including density and polarization fluctuations on length scales longer than the molecular diameter, occur on the same time scale as spectral diffusion suggesting that the picosecond time scale can not be assigned to
local molecular motions but rather to collective rearrangements [68, 46]. Although absolute time scales cannot be compared between experiments that are sensitive to different relaxation phenomena, if the structural evolution is collective the temperature dependence of these measurements will be correlated.

The inertial decay of the pump-probe anisotropy has been attributed to librations, or hindered rotations, of the OX (where X = H or D) transition dipole, and long time decay to collective reorientation [93]. The observed temperature-dependence of the long time relaxation agrees with previous studies using methods related to (with proper assumptions) the first Legendre polynomial: dielectric relaxation [6] and THz time domain spectroscopy [5]; and the second Legendre polynomial: NMR [4, 114] and OKE [3]. Tielrooij and Bakker measured the temperature dependence of the reorientation time of the OD stretch of HOD in H2O with ultrafast infrared spectroscopy and found it decreased from 4.8 to 0.97 ps from 274 to 343 K [100], which is in reasonable agreement with our study. Assuming the long time decay follows Arrhenius behavior, \(1/\tau_c = A \exp[-E_a/RT]\), we find a barrier height of 3.7 kcal/mol compared to 4.1 kcal/mol found by Petersen and Bakker [188]. The Arrhenius plot in Figure 5-17 shows that the trend is nearly linear with a slight quadratic dependence.

Previous measurements of the spectral diffusion of isotopically dilute water have assigned the long-time decay to collective reorganization of the liquid structure [68, 69]. Using the long-time decay from the temperature-dependent frequency correlation functions determined with the model (2.4 ps for 278 K to 0.7 ps for 345 K) and assuming Arrhenius behavior we find a barrier height of 3.4 kcal/mol. As can be seen in Figure 5-17 the trend is linear, with a slight quadratic dependence in an opposite manner as the reorientation. We also determined the barrier height for the integrated real part of the correlation functions and found 2.9 kcal/mol with a trend that is quite linear. Finally, we also characterized the temperature-dependent spectral diffusion by fitting the phase slope decay with a number of functional forms and determining the correlation time either from the decay directly, integrating the normalized function, or from the 1/e period and found a barrier height of 3.4 ± 0.5 kcal/mol and a trend more similar to the reorientation [182]. The differences in barrier height and trend
Figure 5-17: Arrhenius plot of the picosecond decay of the temperature-dependent best-fit frequency correlation function ($C_\omega(t)$, open circles) determined with the spectral model and anisotropy ($C_R(t)$, open squares) along with previous temperature-dependent measures of reorientational relaxation in water: optical Kerr effect [3] (OKE, filled circles), NMR [4] (crosses), terahertz time domain spectroscopy [5] (THz-TDS, asterisks), and dielectric relaxation [6] (DR, plusses). We also show the result of applying the Stokes-Einstein-Debye equation (SED, line).

should be taken as the uncertainty in our determination of the picosecond spectral diffusion time scale.

The similar temperature dependence for both these dynamics experiments (with activation energies of 3.4 kcal/mol for spectral diffusion and 3.7 kcal/mol for reorientation) suggests a common mechanism. Recent theoretical work by Laage and Hynes [1, 32], found that reorientation occurred through large amplitude angular jumps that exchanged a hydrogen bond with a small contribution from the diffusive motion of the hydrogen bond frame. This would suggest exactly what is observed in the current study since exchanging hydrogen bonds causes a significant change in frequency [73]. The authors also looked at the temperature dependence of reorientation and found that the large amplitude jumps and diffusive motion both contribute at every temperature but large amplitude jumps are dominant [32].
In Figure 5-17 we show an Arrhenius plot that includes previous temperature-dependent studies of the first and second order orientational correlation functions along with the results of the current study. The barrier height is similar for all the experiments included in the figure, and a similar quadratic dependence is observed. Results from quasi-elastic neutron scattering (QENS) experiments have not been included since recent theoretical work called for the reassessment of reorientation times of water determined from QENS experiments [189]. The barrier heights from the plot range from 3.9 to 4.7 kcal/mol over liquid phase temperatures where 4.7 kcal/mol is from the dielectric relaxation (DR) study, which samples the higher slope portion of the curve. The ratio between $\tau_{R,2}$, measured in the current study with pump-probe anisotropy, and $\tau_{R,1}$, measured with THz time domain spectroscopy (THz-TDS) [5] is $2.7 \pm 0.1$ over the temperature range in the current study and there is no clear temperature dependence. We also plot the result of calculating $\tau_{R,2}$ derived from the Stokes-Einstein-Debye (SED) equation, $\tau_{R,2}=4\pi\eta(T)R^3/3kT$, where $\eta(T)$ is the temperature-dependent viscosity [190] and $R$ is the hydrodynamic radius [16]. For both quantities we use the values for H$_2$O. The Stokes-Einstein-Debye (SED) equation assumes the reorientation is diffusive, which predicts that the ratio between $C_{R,1}(t)$ and $C_{R,2}(t)$ is 3. We do not observe this ratio experimentally and reorientation dominated by diffusive steps disagrees with the theoretical work by Laage and Hynes [1, 32]. However, the temperature dependence of $\tau_2$ calculated with the SED equation shows a similar barrier height (4.3 kcal/mol) and quadratic behavior. Although we are comparing experiments that are each uniquely sensitive to the evolving structure of the liquid we observe a similar barrier and for the most part a similar quadratic behavior. This observation again points to the collective nature of the picosecond relaxation in water.

In previous studies, the non-Arrhenius behavior of thermodynamic and transport properties of water have been attributed to relaxation processes characteristic of supercooled liquids. Speedy and Angell found the temperature dependence of the isothermal compressibility, expansivity, isobaric heat capacity, diffusion coefficient, viscosity, dielectric relaxation, and $^{17}O$ spin relaxation rate followed power law
behavior. A plot of relaxation rates against \( \log(T/T_s-1) \) using a homogeneous nucleation temperature \( T_s = 228 \text{ K} \) was linear [191, 192]. It has been suggested that water undergoes a fragile-to-strong transition at this temperature [193] and the relaxation processes subsequently assume an Arrhenius behavior until the glass formation temperature [194]. The Vogel-Fulcher-Tammann law, which has also been used to model relaxation rates in supercooled water, predicts a linear relationship between the log of the temperature dependence of the property and \( 1/(T-T_c) \) [114]. Although several empirical relationships have been established, the microscopic explanation for the non-Arrhenius behavior in terms of molecular behavior is unclear, and indicates that a need for further studies at this level.

5.6 Molecular Dynamics Simulations

5.6.1 Fixed Charge SPC/E

We calculate temperature-dependent OD vibrational dephasing and orientational correlation functions with MD simulations to support our conclusions on the collective nature of hydrogen bond rearrangement in water. We used the fixed point SPC/E water potential [31], which was used to illustrate concerted hydrogen bond switching and to calculate electric field induced vibrational frequency shifts [1, 46]. The method only differs from previous MD simulations performed in our group in that HOD is placed in a bath of H$_2$O rather than D$_2$O and the OD stretching frequency is calculated [68, 71]. Trajectories were calculated at four temperatures (273, 298, 310, and 330 K) with the density chosen to match experiments. The OD frequency correlation functions, \( C_\omega(\tau)=(\delta \omega_{OD}(0)\delta \omega_{OD}(\tau)) \), and OD bond reorientational correlation functions, \( C_2(\tau)=(P_2(\hat{r}_{OD}(0)\cdot\hat{r}_{OD}(\tau))) \), are shown in Figure 5-18a and 5-18b, respectively. For \( C_\omega(\tau) \), the sub-100 fs decays are indistinguishable, but the picosecond tails show relaxation times that decrease from roughly 1.4 ps at 273 K to 450 fs at 330 K. Similarly for \( C_2(\tau) \), beyond the early librational time scales, the correlation functions decay faster with increased temperature, from 4.5 ps at 273 K to 1.4 ps at
Figure 5-18: Normalized temperature-dependent correlation functions for (a) the OD stretch vibrational frequency, $C_\omega(\tau)$, and (b) OD bond reorientation, $C_2(\tau)$, from MD simulations with the SPC/E water model. (c) Arrhenius plots of correlation times in femtoseconds obtained from the simulation.

330 K. An Arrhenius plot (Figure 5-18c) for the temperature-dependent correlation times gives a barrier to frequency randomization of $E_{a,\omega} = 3.6$ kcal/mol and a barrier of $E_{a,\text{rot}} = 3.6$ kcal/mol for reorientation. The latter number is in agreement with the 3.5 kcal/mol value previously calculated by Laage and Hynes for the OH stretch [32]. These values are similar to what we observe experimentally (3.4 kcal/mol for spectral diffusion and 3.7 kcal/mol for reorientation) and the same quadratic behavior is observed for $C_2$. 

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5.6.2 Polarizable SPC/FQ

The frequency and orientational correlation functions calculated with the SPC/FQ water model [88] were provided by Steve Corcelli (University of Notre Dame) and James Skinner (University of Wisconsin Madison). The MD simulations of neat H$_2$O using the electronically polarizable SPC-FQ model have been described previously. Here we will summarize the most salient details. As in the SPC/E model, in the SPC/FQ model the geometry of the individual water molecules is held rigid with an OH bond length of 1 Å and an HOH angle of 109.47°. Unlike the SPC/E model, SPC/FQ models electronic polarizability by allowing its three atomic centered point charges to fluctuate based on the instantaneous environment of the water molecule. In principle, the charges could be calculated self-consistently at each MD time step, but in practice the charges are regarded as dynamical variables in an extended Lagrangian and propagated in time [88]. This elegant approach results in a substantial computational savings. There are three reasons why the SPC/FQ model was chosen for this work on the temperature-dependent ultrafast IR spectroscopy of HOD/H$_2$O: (1) Previous studies at room temperature have shown that the SPC/FQ model exhibits rotational dynamics that are in better agreement with experiment than other commonly employed non polarizable water models (e.g. TIP4P [87] and SPC/E [31]) [88]. (2) At room temperature the longest time scale for the decay of the frequency fluctuation time correlation function is also in better agreement with experiment than TIP4P and SPC/E [85]. (3) The IR absorption and Raman scattering spectra computed for SPC/FQ are in excellent agreement with experiment from 10 to 90°C [78].

We calculated trajectories of 128 SPC/FQ H$_2$O molecules at five temperatures in cubic boxes whose sizes were chosen to mimic the experimental number density at 283, 303, 323, 343, and 363 K. The 1 ns trajectories were collected in the NVE ensemble with a 0.5 fs time step, where the charge degrees of freedom were maintained at 5 K by rescaling the charge velocities every 1000 steps. A running average of the temperature was monitored to ensure that it did not deviate outside of ±1.5 K of
the target temperature. Other details regarding the MD simulations are contained in Reference [78].

Two quantities were computed from the trajectories: the OH bond orientational correlation function, \( C_R = \langle P_2[\hat{u}_{OH}(0) \cdot \hat{u}_{OH}(t)] \rangle \), where \( P_2 \) is the 2nd Legendre polynomial and \( \hat{u}_{OH}(t) \) is a unit vector that points along an OH bond, and the normalized electric field fluctuation time correlation function, \( C_E(t) = \langle \delta E_{OD}(0) \delta E_{OD}(t) \rangle / \langle (\delta E_{OD})^2 \rangle \), where \( E_{OD}(t) \) is the magnitude of the electric field due to the surrounding water molecules along an OH bond of interest and \( \delta E_{OD}(t) = E_{OD}(t) - \langle E_{OD} \rangle \) represents the fluctuation of \( E_{OD}(t) \) from its equilibrium value, \( \langle E_{OD} \rangle \). \( C_R(t) \) was computed for the 256 independent OH bonds in the simulations. Although the simulations were of neat \( \text{H}_2\text{O} \), for the purposes of computing \( C_E(t) \) each of the bonds in the simulation were assumed to be the OD stretch of interest, which assumes that the dynamics of a single HOD molecule in water are similar to that of an \( \text{H}_2\text{O} \) molecule. The electric fields along each OH bond were then computed using the instantaneous charges on each of the other 127 water molecules in the simulation, where the effects of long ranged electrostatics were incorporated using an approximation to the Ewald sum [195].

The rationale for computing the normalized electric field fluctuating correlation function is that, within certain approximations, \( C_E(t) \) is equal to the normalized frequency fluctuation correlation function, \( C_\omega(t) \), which can be extracted from the ultrafast 2D IR measurements on HOD in \( \text{H}_2\text{O} \). The key assumption needed to connect \( C_E(t) \) with \( C_\omega(t) \) is that the instantaneous OD vibrational frequency of interest can be linearly related to the electric field along the bond, \( \omega_{OD} = a + bE_{OD} \). Previous density functional theory calculations on 100 HOD-(\( \text{H}_2\text{O} \))\(_n\) clusters containing between 4 and 9 water molecules extracted from a room temperature MD simulation have empirically established a linear relationship (\( a = 2745.8 \text{ cm}^{-1} \) and \( b = 4870.3 \text{ cm}^{-1}/\text{au} \)) between the OD stretch frequency of HOD with the electric field due to the surrounding solvent along its bond. It is important to note, as a caveat, that more recent work on more and larger clusters, albeit for different water models, have found that quadratic relationships between the electric field and vibrational frequency
Figure 5-19: Normalized temperature-dependent correlation functions for (a) the OD stretch vibrational frequency, $C_\omega(t)$, and (b) the bond reorientation, $C_R(t)$, from MD simulations with the SPC/FQ water model. (c) Arrhenius plot of the correlation times in femtoseconds obtained from exponential fits to the picosecond decay.

are more appropriate [55, 56]. Nevertheless, for discerning trends in the long time decay of the frequency fluctuation correlation function as a function of temperature, invoking the approximately linear relationship available from the previous literature is reasonable.

The results for the frequency time correlation functions, $C_\omega(t)$, and OD bond
orientation correlation functions, $C_{R,2}(t)$ for five temperatures in the range from 283 to 363 K are shown in Figure 5-19. The functions show the same trend as observed in the experiment: the long time decay is faster with increasing temperature. Also, similar to the results using the SPC/E model, we find that the inertial dynamics are roughly independent of temperature, although the amplitude does increase with temperature for the orientational correlation functions. We fit the picosecond decay with single exponentials and the time constants are shown in an Arrhenius plot in Figure 5-19c. The behavior is clearly nonlinear, which is similar, although more pronounced, than the experimental orientational relaxation and other observables (see Figure 5-17). Interestingly, the temperature-dependent trends for the two relaxation processes are similar within a scaling factor. We find the activation energy is larger for the polarizable SPC/FQ model than the activation energy determined with the fixed-charge SPC/E model [182]: 6.4 kcal/mol for $C_{R,2}(t)$ and 5.0 kcal/mol for $C_\omega(t)$.

Using temperature-dependent experimental NMR measurements of specific molecule-fixed unit vectors and MD simulations, Ropp et al. showed that rotational motion in water is anisotropic [196]. We show the calculated orientational correlation function for the OD bond vector (bond), out of plane vector (perp), $C_2$ symmetry axis, and H-H intramolecular axis in Figure 5-20. Although the time scales differ, the non-Arrhenius behavior is the same for all the calculated orientational correlation functions.

5.7 Concluding Remarks

By using ultrafast infrared spectroscopy to measure the temperature-dependent picosecond decay of reorientation and spectral diffusion of the OD stretch of dilute HOD in H$_2$O, and comparing the non-Arrhenius behavior to previous measures of relaxation processes in water, we have found that hydrogen bond rearrangement in water is collective. To determine the picosecond decay we self-consistently model 2D IR, pump-probe, and linear absorption measurements with a response function formalism that includes the effects of spectral diffusion, population lifetime, and reorientation
motion and non-equilibrium heating upon vibrational relaxation. Prior experimental and theoretical work has shown that hydrogen bond switching is concerted and involves large angle reorientation [71, 1]. This work provides evidence that translation and reorientation are intertwined with hydrogen bond rearrangement in the collective reorganization of the liquid. MD simulations of temperature-dependent OD vibrational and orientational correlation functions are used to support our conclusions. It remains a question what the correlation length for reorganization is, which collective variables may be best suited to describe or study hydrogen bond rearrangement, and the microscopic origin of the non-Arrhenius behavior from the liquid to supercooled regions of water.

5.8 Acknowledgments

We would like to thank Krupa Ramasesha, Sean Roberts, and Joel Eaves for the temperature-dependent MD simulations using the SPC/E water model. We would like to thank Krupa Ramasesha for her help acquiring the 2D IR and pump-probe anisotropy measurements.
We explicitly include the background absorption of the combination band and stretch of the bath H₂O in our model to explore the origin of the long time offset observed in the phase line slope. The H₂O combination band absorbs in the OD stretch region and there is a relatively small amount of absorption from the H₂O stretch. However, linear absorption and heterodyned third order measurements scale with the transition dipole to the second and fourth power, respectively. The molar absorptivity of the H₂O combination band and stretch are \( \sim 3.5 \) and \( \sim 100 \text{ M}^{-1} \text{ cm}^{-1} \) at the peak, respectively [112]. Given the sensitivity of the transition dipole on environment, with an increasing transition dipole with hydrogen bonding, it is not unreasonable for the H₂O stretch to contribute to the third order signal in the region of the OD stretch [197, 112].

Since H₂O was explicitly included in the model, all of the parameters discussed for the OD stretch had to be defined for the H₂O combination band and stretch. The transition frequencies, \( \omega_{10} \), and transition dipoles, \( \mu_{10} \), are determined by fitting the temperature-dependent linear absorption of pure H₂O. The fits are shown in Figure 5-1b. The room temperature anharmonicities used to determine \( \omega_{21} \) are 300 cm\(^{-1} \) for the H₂O combination band [112] and 250 cm\(^{-1} \) for the H₂O stretch [109]. The anharmonicity is held constant for all temperatures. Room temperature vibrational lifetime, \( T_1 \), and intermediate state lifetime, \( T^* \), are 140 fs and 840 fs for the H₂O combination band [112] and 240 fs and 740 fs for the stretch [109], respectively, which are held constant for all temperatures. It is assumed that the H₂O background shares the same orientational and frequency correlation function as the OD stretch at each temperature. The change in transition dipole, \( \Delta \mu \), and intensity of the TSGS response, \( A^{TSGS} \), for the H₂O combination band and stretch are set to values previously found for the OH stretch of HOD in D₂O at room temperature and held constant for all temperatures [198]. The temperature-dependent parameters are given in Table 2 and the temperature-independent parameters are given in Table 3.

In Figure S-21 we show 2D surfaces calculated with the model at 278 K at \( \tau_2 = 0 \text{ fs} \),
Table 2: Summary of the temperature-dependent parameters contained within our spectral model for the stretch ($\nu_S$) and combination band ($\nu_{L+\delta}$) of H$_2$O.

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<td>$\Delta\omega$ [cm$^{-1}$] (with 1 K)</td>
<td>-0.72</td>
<td>-0.75</td>
<td>-0.80</td>
<td>-0.92</td>
<td>-1.03</td>
</tr>
</tbody>
</table>

H$_2$O $\nu_S$

| $\omega_{10}$ [cm$^{-1}$] (from FTIR) | 3389 | 3395 | 3401 | 3419 | 3433 |
| $\omega_{10}$ [cm$^{-1}$] (input model) | 3403 | 3409 | 3415 | 3432 | 3446 |
| FWHM [cm$^{-1}$] | 393 | 390 | 392 | 401 | 409 |

Table 3: Summary of the parameters contained within our spectral model for the stretch ($\nu_S$) and combination band ($\nu_{L+\delta}$) of H$_2$O that do not depend on temperature.

<table>
<thead>
<tr>
<th></th>
<th>H$<em>2$O $\nu</em>{L+\delta}$</th>
<th>H$_2$O $\nu_S$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\Delta$ [cm$^{-1}$]</td>
<td>300</td>
<td>250</td>
</tr>
<tr>
<td>$\Delta\omega$ [cm$^{-1}$]</td>
<td>See Table 2</td>
<td>0.66</td>
</tr>
<tr>
<td>FWHM [cm$^{-1}$]</td>
<td>300</td>
<td>See Table 2</td>
</tr>
<tr>
<td>$T_1$ [fs]</td>
<td>140</td>
<td>240</td>
</tr>
<tr>
<td>$T^*$ [fs]</td>
<td>840</td>
<td>740</td>
</tr>
<tr>
<td>$\Delta\mu$</td>
<td>0.9986</td>
<td>0.9986</td>
</tr>
<tr>
<td>$A_{TSGS}$</td>
<td>62</td>
<td>62</td>
</tr>
</tbody>
</table>
600 fs, and 4 ps. The first and second columns show the results of the calculation with only the response of the H₂O combination band and stretch, respectively. Although the lifetimes of the H₂O stretch and combination band are relatively short (\( T_1 < 300 \) fs) the modes still contribute at long waiting times due to the TSGS. The H₂O stretch blue-shifts and the combination band red-shifts with increasing temperature; therefore, the signal from the TSGS is positive in the region of the OD stretch for both modes. Although the peak of the H₂O stretch is shifted from the OD stretch \( \sim 2.4 \) x further than the combination band at 278 K, we find the H₂O stretch has a larger contribution to the signal, albeit still small, near the peak of the OD stretch \( (\omega_1, \omega_3 \sim 2500 \text{ cm}^{-1}) \) at all waiting times. In the third column, we show the result when all three modes are included and the fourth column shows the result of windowing by the experimental spectrum after Fourier transform. The experimental spectrum decreases the amplitude of the \( \nu=2 \rightarrow 1 \) signal and the TSGS; and the residual atmospheric CO₂ additionally absorbs from \( \sim 2320 \) to 2360 cm\(^{-1}\).

.2 Vibrational Stokes Shift

The spectroscopic model used in this study assumes the OD stretch is coupled to a thermally occupied harmonic bath [70]. The complex correlation function is proportional to the coupling parameter, \( \lambda \), which is determined by fitting the FWHM of the linear absorption spectrum. At room temperature \( \lambda = 2.7 \times 10^{-3} \text{ fs}^{-1} \), which represents a Stokes shift of 14 cm\(^{-1}\). If the input frequency and orientational correlation functions and lifetime are kept constant, and only the temperature is changed, the FWHM calculated with the model at the four other experimental temperatures are within \( \sim 1\% \) of experimental values. However, the approximation predicts a linear relationship between temperature and FWHM and a quadratic relationship is observed experimentally (see Chapter 4). If the nonlinear measurements are calculated with the same input \( \lambda \), frequency and orientational correlation functions, and lifetime, but with changing temperature, there is a negligible difference in dynamics. Therefore, the correlation functions must be adjusted at each temperature to match.
Figure -21: Absorptive 2D surfaces calculated with our spectral model at 278 K at \( \tau_2 = 0 \) fs (first row), 600 fs (second row), and 4 ps (third row). The calculation includes only the H\(_2\)O combination band (first column), only the H\(_2\)O stretch (second column), the H\(_2\)O combination band and stretch and OD stretch (third column), and all three modes windowed by the experimental spectrum.

The experiment, which results in a different \( \lambda \) for each temperature (see Table 5.1).

The model developed by Rezus and Bakker to fit vibrational transient absorption constructs the time-dependent signal from two time-independent spectral line shapes [120]. The model does not include spectral shifts that would arise from a vibrational Stokes shift. In 1999, Woutersen and Bakker reported PP experiments on the OH stretch of HOD in D\(_2\)O that probed dynamics on the red and blue side of the line shape upon pumping on the blue and red side, respectively [60]. The authors use a Brownian oscillator model to fit the experimental traces and report a Stokes shift of 74 cm\(^{-1}\) [60]. With MD simulations, Lawrence and Skinner calculated a Stokes shift of 57 cm\(^{-1}\) for the same system [155]. However, with a response function formalism, Yeremenko reproduced the results of Woutersen and Bakker without including a Stokes shift in the model; the observed behavior arises from the spectrally narrow pump and probe.
pulses interacting within the pulse duration [187]. The existence of a dynamical Stokes shift in the vibrational spectroscopy of water has yet to be verified experimentally. The Stokes shift will be reflected in dispersed pump probe measurements as a time-dependent shift of the zero crossing between the $\nu=1\leftarrow 0$ and $\nu=2\leftarrow 1$ peaks. We observe a 27 cm$^{-1}$ red-shift of the zero-crossing by 150 fs for all temperatures in the current study. However, this may very likely be an artifact of cross-phase modulation. However, a Stokes shift falls out of the response function formulism used to self-consistently model the linear and non-linear measurements.

![Video](image)

Figure -22: Dispersed pump-probe calculated with the model at 278 K with (a) $C_\omega(t)$ without including the TSGS, (b) $C_\omega(t)$ without including the TSGS, and (c) $C_\omega(t)$ including the TSGS.

Since the TSGS also causes a shift in the zero-crossing we calculate the dispersed pump-probe (DPP) with and without the TSGS. In Figure S-22, we show DPPs calculated from the model at 278 K for three cases: (a) only including the real part of $C_\omega(t)$, or $C'_\omega(t)$, and not including the TSGS, (b) using the complex $C_\omega(t)$ but not including the TSGS, and (c) using the complex $C_\omega(t)$ and including the TSGS. In Figure S-22a the $\nu=1\leftarrow 0$ and $\nu=2\leftarrow 1$ peaks decay at the same rate with no time-dependent shift in the zero crossing. As can be seen in Figure S-22b, there is a shift in the zero-crossing with increased delay, even when the TSGS is not included. The zero crossing red-shifts by 9.1 cm$^{-1}$ from $\tau_2 = 0$ to 400 fs and an additional 5.9 cm$^{-1}$ between 400 fs and 4 ps. When the TSGS is included, the zero crossing red-shifts an
additional 1.3 cm⁻¹ between \( \tau_2 = 0 \) to 400 fs and from \( \tau_2 = 400 \) fs to 4 ps the zero crossing shifts a total of 62 cm⁻¹. The effect of the population relaxation induced TSGS can clearly be seen in Figure S-22c: there is a persistence of the positive peak to longer waiting times and the appearance of a negative peak at higher frequencies [175, 94].

.3 Frequency-Dependent Reorientation

With MD simulations, Laage and Hynes found the primary mechanism of reorientation in water is through a large angle jump concurrent with switching a hydrogen bond partner [1, 32]. The mechanism leads to predictions for what will be observed in a frequency-resolved anisotropy measurement since the frequency at which a molecule absorbs is sensitive to its environment. Correlation between frequency and reorientation has been observed with MD simulations of the OH stretch of HOD in D₂O, which found a larger amplitude inertial reorientation in the orientational correlation function for molecules absorbing on the blue side of the line width [45]. The 2D anisotropy has also been calculated with MD simulations for the OH stretch of HOD in D₂O [74]. The study found that the anisotropy of molecules that are initially in broken or strained hydrogen bonds (NHB) drops from 0.4 to 0.25 within 300 fs, whereas there is little change in anisotropy for molecules initially in stable hydrogen bonds (HB) within that time scale. Also, the drop in anisotropy in the off-diagonal region of the 2D surface is similar to the diagonal region for NHB indicating that molecules reorient in order to switch from NHB to HB and vice versa [74].

Bakker et al. measured the pump-probe anisotropy of the OD stretch of HOD in H₂O pumping at the center and blue wing of the line width [119]. They find that when pumping at the center of the line width the anisotropy decay is independent of frequency. However, when pumping on the blue wing, the anisotropy of the molecules on the red side of the line width drops to 0.2 by the initial time point, 200 fs. This observation suggests that molecules initially excited on the blue wing must reorient to return to the red side of the line width [119]. Moilenan et al. measured the pump-
probe anisotropy of the OD stretch of HOD in H₂O at three temperatures: 1, 25, and 65°C [121]. Although spectral information is obscured within the first 200 fs due to non-resonant effects, the authors extrapolate to 100 fs and find, at room temperature, there is a greater initial drop in anisotropy on the blue side of the line width (~0.36 at 2500 cm⁻¹ compared to ~0.32 at 2625 cm⁻¹) and after 200 fs, the anisotropy decay is frequency independent. At 65°C the difference across the line width at 100 fs is exaggerated compared to room temperature and at 1°C they observe little difference across the line width [121]. Finally, Skinner et al. calculated the temperature dependence of the pump-probe anisotropy with MD simulations and found that although there is a larger initial drop in anisotropy at higher temperature, the frequency dependence, in which the anisotropy drops to a lower value on the blue side of the line width upon inertial reorientation, is independent of temperature [179].

In this section we will present the room temperature pump-probe and waiting time dependent 2D anisotropy of the OD stretch of HOD in H₂O. We find a larger initial drop in anisotropy on the blue side of the line width for both measurements. We will also show the effect of non-resonant effects on the pump-probe anisotropy, which must be treated with care when extracting information on the inertial decay.

Figure -23: The calculated anisotropy at room temperature from our spectral model with the effect of pulse convolution (left). The frequency-dependence of the calculated anisotropy at room temperature from our spectral model including the effect of pulse convolution and cross phase modulation (right).
First we will show the effect of the pulse and non-resonant effects, explicitly cross phase modulation, on pump-probe anisotropy. In the model calculation, we begin with the $S_{zzzz}$ and $S_{zzyy}$ pump-probes calculated with our spectral model at room temperature and convolve a Gaussian pulse with different pulse lengths with both polarization geometries before determining the anisotropy. We show the result in Figure S-23 for 40, 80 and 160 fs pulses. The anisotropy still begins at 0.4 but the inertial decay, 50 fs without pulse convolution, appears slower when the pulse is included. In the current study the pulse width is $\sim$80 fs (shown in red), which will obscure the anisotropy until $\sim$200 fs. Next we include cross phase modulation from a fit to the experimentally measured cross phase modulation (XPM) assuming the pulse is not chirped (see Chapter 2). The result is shown in Figure S-23(right). There is no frequency-dependence included in the model, however, XPM introduces an apparent difference across the frequency range until $\sim$80 fs. Additionally, the anisotropy does not begin at 0.4. The XPM has the smallest effect near the peak frequency of the pulse and becomes more important with detuning between the peak of the pulse and probe frequency. In other words, if one were to take a slice at 50 fs the value of the anisotropy would appear progressively smaller as the probe frequency is detuned from the peak of the pulse.

![Figure -24: Slices in frequency (left) and $t = 200$ fs (right) of the dispersed pump-probe anisotropy acquired at room temperature.](image)
In Figure S-24 we show the experimental dispersed pump-probe anisotropy of the OD stretch of HOD in H₂O acquired at room temperature. Similar to what we observe in the model calculation when the XPM is included, the anisotropy does not begin at 0.4. Nonetheless, if we take a slice at t = 200 fs, where pulse effects should no longer be important, we find that the anisotropy has dropped to a considerably lower value on the blue side of the line width. This observation agrees with the molecular jump mechanism for reorientation [1, 32] and previous MD simulations of the frequency dependence of the anisotropy [45]. Also, although the frequency dependence of the initial drop is more severe compared to the previous measurement [121], we notice the measurements agree in that the long time decay is identical over the frequency range.

![Figure S-25](image)

Figure S-25: Absolute value 2D anisotropy of dilute HOD in H₂O acquired at $\tau_2 = 100, 250, 300, 400, 1000$ fs.

In Figure S-25 we show the absolute value 2D anisotropy of HOD in H₂O at the waiting times, $\tau_2 = 100, 250, 300, 400, 1000$ fs. The absolute value 2D anisotropy, which is calculated from absolute value 2D surfaces acquired in the parallel and
perpendicular geometry, is used to avoid divergence in the anisotropy determined from the absorptive surfaces due to the node between the \( \nu=1\leftarrow 0 \) and \( \nu=2\leftarrow 1 \) peaks. At the earliest waiting time, \( \tau = 100 \) fs, the anisotropy is higher at \( \omega_1 = \omega_3 = \) blue, which disagrees with the trend observed for the following waiting times. However, at 100 fs pulse effects are most likely still important. Past 100 fs the anisotropy is higher on the red side of the diagonal relative to both the blue side of the diagonal and the off-diagonal regions. This observation is in reasonable agreement with what has been calculated with MD simulations [74] and what is suggested by the molecular jump mechanism for reorientation [1]. In Figure S-26 we show slices in the absolute value anisotropy at \( \omega_1 = 2475 \) (left) and 2550 cm\(^{-1}\) (right). On the red side of the line width the anisotropy is slightly lower at \( \omega_3 = \) blue and remains largely unchanged for the first four waiting times. At 1 ps the anisotropy drops to \(-0.22\). On the blue side of the line width the anisotropy is again slightly lower on the \( \omega_3 = \) blue side and drops faster than the red side. The results of the 2D anisotropy measurements need to be confirmed experimentally and analyzed in more detail to extract quantitative information on the correlation between frequency shifts and reorientation.

![Figure S-26](image)

Figure -26: Slices in the 2D absolute value anisotropy with waiting time at \( \omega_1 = 2475 \) cm\(^{-1}\) (left) and \( \omega_1 = 2550 \) cm\(^{-1}\) (right)
Appendix A

Pump-Probes and 2D IR of Pure D₂O

In this thesis we presented linear and non-linear measurements of dilute HOT and HOD in H₂O. An isotopically dilute system provides a probe of local structural dynamics as resonant intermolecular transfer is turned off and non-resonant intermolecular transfer will occur on a longer time scale than spectral diffusion. Dynamics of the isotopically dilute system imitates neat water for any process largely dominated by electrostatics. However, studying pure water and the effects of delocalization is an interesting problem in of itself. We have, therefore, performed preliminary experiments on pure D₂O. In the gas phase, the symmetric ($\nu_1$) and antisymmetric ($\nu_3$) stretch of D₂O absorb at 2669.40 and 2787.92 cm⁻¹, respectively [199]. In the condensed phase the degeneracy between the local modes is largely broken, given the different local environments, although there is opportunity for coupling to nearby oscillators. Band center shifts to 2500 cm⁻¹ and the FWHM is 325 cm⁻¹, which is ~2.7x broader than the frequency splitting in the gas phase. Delocalization has been shown to affect both linear and non-linear measurements of pure water [200, 56]. For instance, Auer and Skinner recently calculated the IR and Raman spectra of H₂O with a mixed quantum classical theoretical method and found the peak at 3250 cm⁻¹ in the VV Raman spectrum arises from collective interactions [56]. The calculation of the line shapes included both intramolecular coupling and intermolecular coupling.
estimated by transition dipole coupling. They found the instantaneous distribution of local mode frequencies of HOD in D$_2$O and pure H$_2$O was very similar and, when the coupling was turned off, the relatively broad IR line shape calculated for H$_2$O resembled the line shape for the isotopically dilute system. In Figure A-1 we show the normalized linear absorption spectrum of pure H$_2$O (blue), pure D$_2$O (green), and the OD stretch of HOD in H$_2$O (red) scaled in frequency by their respective gas phase frequencies (2723.68 cm$^{-1}$ for the OD stretch of HOD [159]). For H$_2$O and D$_2$O the average of the symmetric and antisymmetric gas phase frequencies was used (for H$_2$O $\nu_1 = 3657.05$ cm$^{-1}$ and $\nu_3 = 3755.93$ cm$^{-1}$ [199]). Indeed the line shape for H$_2$O and D$_2$O is broader than the isotopically dilute system. It is interesting to note that the line shape of D$_2$O is broader than H$_2$O. This is the opposite trend then what is expected for motional narrowing and what has been observed for the isotopically dilute system [174].

Figure A-1: Normalized linear absorption spectrum of H$_2$O (blue), D$_2$O (green), and the OD stretch of HOD in H$_2$O (red) scaled in frequency by their respective gas phase frequencies.

Miller and co-workers measured the room temperature and temperature dependent 2D IR and pump-probes of pure H$_2$O [200, 107]. At room temperature they found spectral diffusion occurred within 50-100 fs, which lead to the suggestion that spectral diffusion was dominated by librational motion. They found that as temper-
ature was decreased to 274 K frequency correlations persisted beyond ~200 fs. The relatively fast ~80 fs anisotropy decay, which was attributed to resonant intermolecular energy transfer, was unaffected by temperature. Finally, it was found that the lifetime increased from 180 fs at 274 K to 300 fs at 340 K. Paarman et al. used a fully anharmonic nonadiabatic simulation procedure, which explicitly treated intermolecular coupling as well as fluctuations and anharmonicities of transition frequencies and dipole moments, to calculate the pure H₂O 2D IR and anisotropy measurements [201]. They found the fast dephasing observed in the 2D IR measurements was mainly a result of intramolecular mixing between states in the two-dimensional anharmonic OH stretch potential. Intermolecular coupling was necessary match the fast anisotropy decay, but they found 12 cm⁻¹ coupling sufficient [201]. For pure D₂O, only pump probes have been reported to this point, which show a 400 fs lifetime [113]. Here we present 2D IR measurements and pump-probes (Szzzz) for pure D₂O. Indeed we observe that the dynamics are very fast and, given the broad absorption line shape of D₂O, our pulse width limits the spectral information.

In Figure A-2a we show the dispersed pump-probe of pure D₂O acquired in the zzzz polarization geometry. We held D₂O between two CaF₂ windows without a spacer. The optical density was ~1. We acquired dispersed pump-probes by fast scanning the delay between the pump and probe, then averaging the scans. Slices in frequency are shown in Figure A-2c. Based on the previous measurement of the lifetime of the OD stretch of D₂O the decay observed within the first ~500 fs corresponds to decay from the excited state. The positive (negative) amplitude to the red (blue) of ~2600 cm⁻¹ that grows in with delay between the pump and probe is most likely due to the TSGS, which appears to have a much larger effect in pure D₂O compared to isotopically dilute water. In Figure A-2b and Figure A-2d we show slices in time. Also, in Figure A-2b we show the absorption spectrum of D₂O as the dashed gray line. Since our pulse width (~230 cm⁻¹) is narrower than the absorption band the transient absorption signal is narrowed. The delay times in Figure A-2b correspond to the waiting times of the 2D surfaces we will show shortly. In Figure A-2d the delay times are 0 (blue), 100 fs (green), 200 fs (red), 400 fs (cyan), 800 fs (magenta), 1.6 ps
It is surprising to see the signal at long time surpasses the signal at $t = 0$ fs. If we fit a mono-exponential the frequency slice that corresponds to the zero crossing between the positive and negative amplitude of the TSGS, the magenta line shown Figure A-2a, we find a decay time of 355 fs. This time scale is faster than the reported $T_1$, which is expected since molecular rotations contribute to the decay given that the measurement is in the $zzzz$ polarization geometry.

Figure A-2: (a) DPP of pure $D_2O$ in the $zzzz$ polarization, (b) Slices in frequency of the DPP, (c) Slices in time of the DPP with the $D_2O$ spectrum shown in dashed gray, and (d) Slices in time to longer delay times.

In Figure A-3 we show 2D IR surfaces for pure $D_2O$ taken in the $zzzz$ polarization geometry in the pump-probe geometry by step-scanning the $\tau_1$ axis in 2 fs steps to 600 fs. The sample conditions were the same as the PP experiments. Again, the response is spectrally narrowed by our pulse spectrum. The $\nu=1\leftarrow0$ peak is very broad, and although it is singly peaked, the line shape is not symmetric. At the two earliest waiting times there appears to be off-diagonal broadening that disappears by $\sim120$-160 fs. This is most likely due to non-resonant response. From $T = 120$ - 200 fs the spectra look very similar, although the $\nu=2\leftarrow1$ peak is decaying. At $T = 240$
fs the $\nu=1\rightarrow0$ peak appears to have broadened along $\omega_1$. Since the dynamics in pure water are relatively very fast it would be best to repeat the experiments with shorter pulses in time and, therefore, a broader spectral range.

Figure A-3: 2D IR of pure $\text{D}_2\text{O}$ at six waiting times, $T = 40, 80, 120, 160, 200$, and 240 fs.
Appendix B

Linear Absorption of Isotopically Dilute Ice Ih

The vibrational frequency of the proton stretch in water is sensitive to its environment. This link stems from the studies of Novak [42], who found a strong correlation between the interatomic oxygen-oxygen distance of hydrogen bonded partners, $R_{OO}$, and $\omega_{OH}$ in hydrogen bonded solids, which was corroborated with molecular dynamics (MD) simulations [68, 46]. Tokmakoff and co-workers used two dimensional infrared spectroscopy (2D IR), a technique capable of distinguishing dynamics of spectrally distinct environments while maintaining fast time resolution, to study dynamics across the line width of the OH stretch of HOD in D$_2$O [71, 73]. They found an asymmetric line shape that indicated water molecules originally absorbing at non-hydrogen bonded (NHB) frequencies, which include both broken and strained hydrogen bonds, return to hydrogen bonded (HB) frequencies within the fastest intermolecular time scales in water [71, 73]. How would the line shape appear in an experiment where the NHB species are largely removed? This experiment can be realized with 2D IR of ice.

All crystalline phases of ice follow the ice rules set forth Bernal and Fowler in 1933 [23], which are: (1) water molecules in ice are intact, i.e. two hydrogens are covalently bonded to each oxygen, neglecting the small fraction of ionized molecules (this rule has the exception ice X), (2) each water molecule is orientated such that two
intramolecular OH bonds are directed approximately toward two of the four nearest neighboring oxygen atoms, and (3) only one hydrogen atom lies approximately along the line joining adjacent oxygen atoms. Weaknesses, or Bjerrum defects, in the ice crystal occur when the ice rules are disobeyed. A L ("leere") defect occurs when two adjacent oxygen atoms have no intervening proton (O...O), and a D ("doppel") defect occurs when there are two hydrogen atoms between adjacent oxygens (O-H...H-O). Other defects may be caused by the presence of H$_3$O$^+$ and OH$^-$ ions. Hexagonal ice (ice Ih) is the form of all natural snow and ice on Earth, as evidenced in the sixfold symmetry in ice crystals grown from water vapor, or snow flakes. The average structure belongs to the P63/mmc centrosymmetric space group. The structure is shown in Figure B-1. It consists of puckered layers perpendicular to the c-crystal axis containing water molecules in a chair form hexagonal ring. There are also boat form hexagonal rings formed by three molecules in one layer and three in an adjacent layer. The O...O distance for H$_2$O is approximately 2.75 Å [202], compared to 2.98 Å for the gas phase dimer [203]. As Pauling noticed in 1935 [204], even under the ice rules there are six possible orientations for every water, or put another way, the hydrogens are disordered. This is true for ice Ih and this give rise to a zero point energy of 3.41 J mol$^{-1}$ K$^{-1}$ [205].

![UNIT CELL](image)

Figure B-1: Structure of Ice Ih [206]
We show the linear absorption of H$_2$O and dilute HOD in H$_2$O in Figure B-2. The sample is held in a brass sample cell between CaF$_2$ windows with a 6 $\mu$m spacer and the temperature is controlled by water/glycerol circulation. The temperature is monitored with a thermocouple taped to the top of the brass sample cell. The red trace was taken near room temperature (25°C) and the dark blue trace is ice (which forms at $\sim$-4°C, according to the thermocouple). The peak wavelength, line width (FWHM), and peak absorption of the OD stretch, with the background H$_2$O subtracted, are shown in Figure B-3. Starting at room temperature, the OD stretch red-shifts from 2509 cm$^{-1}$ to 2444 cm$^{-1}$, the peak absorbance increases by $\sim$4.7x (0.24 to 1.15), and the FWHM decreases by $\sim$4.1x (165 to 40 cm$^{-1}$) upon freezing. The red-shift in frequency is expected given that R$_{OO}$ in ice Ih is $\sim$2.75 Å (H$_2$O) compared to $\sim$2.85 Å (HOD in D$_2$O) [207] on average at room temperature. In crystalline ice the oxygen atoms are ordered, therefore, one would expect a large decrease in the static distribution of frequencies and correspondingly a decrease in the line width. In the liquid phase the line shape is mainly inhomogeneously broadened [155]. For instance, a simple calculation shows that the lifetime of $\omega_{OD}$ in water contributes $\sim$3.5 cm$^{-1}$ to the 165 cm$^{-1}$ line width. The lifetime of $\omega_{OD}$ in ice has yet to be measured, however, the lifetime of $\omega_{OH}$ in isotopically dilute ice is 380 fs [97]. The FWHM of $\omega_{OH}$ in the ice form is $\sim$50 cm$^{-1}$ compared to $\sim$265 cm$^{-1}$ in the liquid phase (not shown). Therefore the lifetime contributes $\sim$25% to the line width in ice compared to $\sim$3% in the liquid phase (given a 700 fs lifetime [93]). Li and Skinner calculated the Raman and IR spectrum for isotopically dilute HOD in H$_2$O and D$_2$O [208]. They found the spectra could be understood as a convolution of Gaussian inhomogeneous broadening due to proton disorder and Lorentzian homogeneous broadening due to motional narrowing from thermal fluctuations [208].

Given that the oxygens in ice are ordered, a lack of inhomogeneous broadening, or even very fast dephasing compared to the width of the static distribution of frequencies, will produce a small, if any, photon echo. If this is the case, the 2D IR line shape of ice will be symmetric in $\omega_1$ and $\omega_3$ at all waiting times. However, disorder of the protons will most likely introduce inhomogeneity, as seen in the calculations of
Figure B-2: Linear absorption of H₂O (upper) and dilute HOD in H₂O (lower) in the liquid and ice phase. The peaks labeled in the upper (lower) spectra correspond to H₂O (HOD) transitions.

the linear absorption [208]. With the constraints of the lattice, spectral diffusion may very well be slower for ice than water. If this is the case, there will be elongation along the diagonal that may persist past ~1 ps. Li and Skinner calculated the frequency correlation function for HOD in D₂O at 245 K [208] and found an initial decay faster than water to negative correlation, a recurrence at 145 fs from the hydrogen bond stretch, and several subsequent beats. This correlation function indicates that within 145 fs the peak will appear homogeneously broadened.

We attempted to acquire a 2D IR measurement of ice in the same manner as the linear absorption measurements were acquired but scatter from the ice dominated the signal. Sample preparation will need to be carefully considered before this experiment is realized.
Figure B-3: (a) Peak wavelength, (b) peak absorbance, (c) line width (FWHM), and (d) linear absorption of $\omega_{OD}$ of HOD in H$_2$O with the background H$_2$O subtracted.
Appendix C

Urea

Osmolytes are small organic molecules that affect the stability of proteins. Stabilizing osmolytes include trimethylamine N-oxide (TMAO), glycerol, and sugars. Urea is a known denaturant of proteins but the mechanism of denaturation is unknown. Urea may disrupt the hydrogen bond network of water surrounding the protein allowing for the solvation of hydrophobic groups. In this case, urea is considered a structure-breaker of water and a chaotropic solute under the Hofmeister series nomenclature. Alternatively, it may interact directly with the side chains and/or protein backbone through electrostatic and/or van der Waals forces. The indirect and direct interaction mechanisms are not mutually exclusive and the dominant mechanism may shift as the protein undergoes conformational changes.

In this appendix we present linear absorption and 2D IR measurements of urea in water and urea and N-methylacetamide (NMA) in water as part of a study of the mechanism of denaturation. Urea association has been studied through infrared spectroscopy by means of principle component analysis (PCA) and 2D correlation spectroscopy [209]. The authors found that the maximum efficiency of urea dimerization occurs at ~2 M, after which urea forms higher aggregates [209]. Ultrafast pump-probe experiments probing water directly upon addition of urea have shown that urea has a negligible effect on the hydrogen bond dynamics of water [210]. There has not been an ultrafast nonlinear infrared experiment performed that directly probes the vibrations of urea. Cross peaks in a 2D measurement arise from coupling, whether
through space or through bond, between spectrally distinct species. Therefore, polarization selective 2D IR measurements of different isotopic species of urea are capable of directly probing if urea forms dimers in water and, if so, the geometry of the dimer. Likewise, 2D IR measurements of urea and spectrally distinct peptides and proteins are capable of providing information on the direct mechanism of denaturation.

Figure C-1: (a) Linear absorption of deuterated $^{12}$C-urea in D$_2$O as a function of concentration normalized to the carbonyl stretch. The background D$_2$O is subtracted. (b) Difference spectrum between each concentration and the 125 mM solution and a ball and stick representation of a urea molecule [oxygen (red), carbon (black), nitrogen (blue), and deuterium (white)].

The concentration dependent linear absorption spectrum of deuterated $^{12}$C-urea in D$_2$O is shown in Figure C-1a and a ball and stick representation of a urea molecule in shown in Figure C-1b. The peak at $\sim$1600 cm$^{-1}$ is primarily composed of carbonyl stretch and the peak at $\sim$1500 cm$^{-1}$ is primarily composed of C-N stretch. As the concentration is increased, additional peaks appear to the red and blue of
Figure C-2: Shift in peak of carbonyl stretch of urea with concentration.

the C-N stretch at \(~1405 \text{ cm}^{-1}\) and \(~1545 \text{ cm}^{-1}\), respectively. The carbonyl stretch blue-shifts with increasing concentration. The frequency of the peak maximum with concentration, which shows a cubic behavior, is shown in Figure C-2. In Figure C-1b we show the difference spectrum between each concentration and the \(C = 125 \text{ mM}\) solution. The blue-shift of the carbonyl stretch is seen along with the decrease in intensity of the C-N stretch at \(~1500 \text{ cm}^{-1}\) and increase at \(~1405 \text{ cm}^{-1}\) and \(~1545 \text{ cm}^{-1}\). It appears as the additional peaks that grow in with concentration are a result of coupling between associated urea molecules.

In Figure C-3 we show waiting time dependent 2D IR surfaces of 1 M \(^{12}\text{C}-\text{urea}\) in \(\text{D}_2\text{O}\) acquired in the ZZYY polarization geometry. The surfaces were acquired in the pump-probe geometry with the mid-infrared pulse centered at \(~1575 \text{ cm}^{-1}\) with \(~450 \text{ cm}^{-1}\) of band width (FWHM). There are peaks along the diagonal from the carbonyl and C-N stretch and the intermediate peak at \(~1545 \text{ cm}^{-1}\). The cross peaks between the carbonyl and C-N stretch are present at the earliest waiting time, \(\tau_2 = 75 \text{ fs}\), and grow in with increasing waiting time. Also, the C-N stretch becomes more intense with increasing waiting time relative to the carbonyl stretch due to the downhill flow of vibrational energy. The cross peaks are more intense relative to the diagonal peaks in the ZZYY polarization geometry compared to the ZZZZ polarization geometry (not shown) indicating that the transition dipoles are closer to 90° than 0°. In agreement with the concentration dependent linear absorption measurements, there is not an
Figure C-3: Waiting time dependent 2D IR surfaces of 1 M $^{12}$C-urea in D$_2$O acquired in the ZZYY polarization geometry.

intermediate peak observed between the carbonyl and main C-N stretching modes in a waiting time dependent series of 2D IR surfaces of 100 mM $^{12}$C-urea in D$_2$O acquired in the ZZYY polarization geometry (not shown).

We can directly confirm the association of urea in water by identifying cross peaks in a 2D IR measurement between isotopically distinct species, namely $^{12}$C-urea and $^{13}$C-urea. In Figure C-4 we show the linear absorption of 2.5 M $^{12}$C-urea and $^{13}$C-urea in D$_2$O. The carbonyl and C-N stretches of $^{13}$C-urea are red-shifted from the stretches of $^{12}$C-urea by approximately 45 and 35 cm$^{-1}$, respectively (determined at 250 mM). In Figure C-5 we show concentration dependent 2D IR surfaces of $^{12}$C-urea and $^{13}$C-urea in D$_2$O acquired at $\tau_2 = 100$ fs in the ZZYY polarization geometry. The cross peaks between the intramolecular carbonyl and C-N stretch of each species is clearly seen. On the blue side of the cross peak between the $^{12}$C-urea carbonyl and C-
Figure C-4: Linear absorption of 2.5 M $^{12}$C-urea and $^{13}$C-urea in D$_2$O (blue). Linear absorption of $^{13}$C-urea and d7-NMA in D$_2$O (green). The amide I and II modes of NMA are indicated. The molecule is a ball and stick representation of NMA [oxygen (red), carbon (black), nitrogen (blue), and deuterium (white)].

N stretch there is additional amplitude that appears with increasing concentration. This additional amplitude may arise from association between isotopically distinct species. However, it may also arise from coupling to the additional intermediate peak that grows in with increasing concentration for each species. In either case, it appears to arise from association between urea molecules.

We additionally acquired a waiting time series of 2D IR surfaces of 2.5 M $^{12}$C-urea and $^{13}$C-urea in D$_2$O in the ZZYY and ZZZZ polarization geometries. The surfaces are shown in Figure C-6 for five waiting times in the ZZYY polarization geometry and at $\tau_2 = 300$ fs for the ZZZZ polarization geometry. With increasing waiting time a ridge grows in between the C-N stretches of $^{12}$C-urea and $^{13}$C-urea. This may be direct evidence for association between isotopically distinct species. The surface acquired in the ZZZZ polarization geometry suggests that if association is occurring, the transition dipoles between molecules are less likely to be aligned parallel to one another.

NMA is a small molecule representation of a single unit of the protein backbone that is ideal to study how urea may interact with a protein, if it indeed denatures by
direct interaction. In Figure C-4 we show the linear absorption spectrum of $^{13}$C-urea and d7-NMA in D$_2$O and in Figure C-7 we show 2D IR surfaces of $^{13}$C-urea and d7-NMA in D$_2$O in the ZZYY and ZZZZ polarization geometries at two waiting times. The peaks along the diagonal from red to blue frequency correspond to the C-N stretch of $^{13}$C-urea, the amide II mode of d7-NMA (C-N stretch and N-D bend), the carbonyl stretch of $^{13}$C-urea, and the amide I mode of d7-NMA (primarily carbonyl stretch). The cross peaks between intramolecular modes are clearly seen in the ZZYY polarization geometry, but there is no indication of intermolecular coupling, even at low contours. The surfaces acquired in the ZZZZ polarization geometry indicate that the intramolecular transition dipoles are aligned closer to $90^\circ$ than $0^\circ$.

In conclusion, linear absorption measurements of a single isotope of urea appear to indicate association of urea molecules in water. Calculations of frequency shifts upon dimerization, or upon oligimer formation, would be helpful in confirming this
Figure C-6: 2D IR surfaces of 2.5 M $^{12}$C-urea and $^{13}$C-urea in D$_2$O acquired at five waiting times in the ZZYY polarization geometry and $\tau_2 = 300$ fs in the ZZZZ polarization geometry.

Additionally, the increasing intensity of a ridge between the C-N stretches of $^{12}$C-urea and $^{13}$C-urea with increasing waiting time observed in 2D IR measurements acquired in the ZZYY polarization geometry is an initial direct confirmation of the association of urea molecules. Linear absorption and 2D IR measurements of urea in an apolar solvent and/or film would help in the characterization of associated urea molecules. 2D IR surfaces of $^{13}$C-urea and d7-NMA in D$_2$O initially indicate that the molecules do not associate in water. Further tests of how urea may interact directly with a protein can include measurements of urea and polyglycine (model of protein backbone), urea and polytryptophan, polyalanine, or polyisoleucine (models of a hydrophobic side chain), and urea and polyasparagine (model of a hydrophilic side chain). Finally, linear and nonlinear measurements of urea and a protein would
Figure C-7: 2D IR surfaces of $^{13}$C-urea and d7-NMA in D$_2$O in the ZZYY and ZZZZ polarization geometries at two waiting times.

be most interesting.

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Bibliography


