Quantum-State Controlled Chemical Reactions of Ultracold Potassium-Rubidium Molecules


How does a chemical reaction proceed at ultralow temperatures? Can simple quantum mechanical rules such as quantum statistics, single-particle scattering, and quantum threshold laws provide a clear understanding of the molecular reactivity under a vanishing collision energy? Starting with an optically trapped near-quantum-degenerate gas of polar $^{40}$K$^{87}$Rb molecules prepared in their absolute ground state, we report experimental evidence for exothermic atom-exchange chemical reactions. When these fermionic molecules were prepared in a single quantum state at a temperature of a few hundred nanokelvin, we observed p-wave-dominated quantum threshold collisions arising from tunneling through an angular momentum barrier followed by a short-range chemical reaction with a probability near unity. When these molecules were prepared in two different internal states or when molecules and atoms were brought together, the reaction rates were enhanced by a factor of 10 to 100 as a result of s-wave scattering, which does not have a centrifugal barrier. The measured rates agree with predicted universal loss rates related to the two-body van der Waals length.

Scientific interest in precisely understanding the fundamental aspects of chemical reactions and controlling their dynamic processes has stimulated pioneering work on molecular beams to study state-to-state reactions using molecular alignment, velocity selections, and collisional broadening measurement. However, substantial motional energies remained in earlier work, and thermal statistical averages were a necessary ingredient. By preparing a molecular ensemble’s translational degrees of freedom in the quantum regime, we expect to develop fundamental insights into how chemical reaction processes may be precisely guided by quantum mechanics. Reaction dynamics at vanishingly low energies remain a fascinating and yet unexplored scientific realm (6). Under unprecedented energy resolution, each step of a complex reaction may be analyzed on the basis of single quantum states and single reaction channels. For example, we can study how reactivity is dictated by the quantum statistics of the molecule as a whole.

That chemical reactions could occur at ultralow temperatures seems at first glance counterintuitive. However, ultracold collisions, where particles scatter only in the partial wave with lowest angular momentum, are governed by quantum statistics and quantum threshold behaviors described by the Bethe-Wigner laws (7–9). In this regime, particles are represented by their de Broglie wavelength, which increases with reduced temperature. This wave nature of particles replaces our intuitive and classical picture of collisions. The wave manifestation of tunneling through reaction or angular momentum barriers may play a dominant role in dynamics, and scattering resonances can have dramatic effects on reactions (10). In addition, any barrierless chemical reactions will always take place when two reactants are sufficiently close together (11). In this case, chemical reaction rates will be determined to a large extent by collisional properties at large intermolecular separations, and thus by how the two partners approach each other. Once their separation reaches a characteristic length scale ($\sim 10^{-10}$ m), a chemical reaction happens with a near unity probability. Therefore, chemical reactions can be surprisingly efficient even at ultracold temperatures. Indeed, this model for barrierless reactions predicts loss rates that are universal in the sense that they do not depend on the details of the short-range interactions, but instead can be estimated using only knowledge of the long-range interactions (12).

Like the case of collisions of ultracold atoms, the study of ultracold chemical reactions will play a fundamental role in advancing the field of molecular quantum gases. For example, understanding and manipulating collisions of atoms at ultralow temperatures ($)1\mu K$ has been crucial for the realization of quantum degenerate gases (13–15), Fermi superfluids that provide opportunities to explore the underlying connection between superconductivity and Bose-Einstein condensation (16), neutral atom–based systems for quantum information science (17–19), and strongly correlated quantum gases (20, 21). Ultracold molecules undergo a more diverse set of collisional processes, with distinct inelastic collision mechanisms arising from chemical reactions, in addition to the traditional state-changing collisions seen with ultracold atoms and highly vibrationally excited molecules (22). Furthermore,
A prerequisite for exploring ultracold chemical reactions is a gaseous molecular sample that is sufficiently dense, ultracold, and suitable for precise control of specific quantum states (6). The starting point for this work is an ultracold trapped gas of fermionic $^{40}{\text{K}}^{87}{\text{Rb}}$ molecules prepared in a single hyperfine level of the rovibronic ground state ($N = 0, v = 0$ of $X\Sigma^+$) (29, 30). The optical trap depth is $-k_B T_{\text{trap}} = 10 \times 10^{-12}$ cm$^{-3}$, where $k_B$ is Boltzmann’s constant. The molecules are produced using a single step of two-photon Raman transfer from extremely weakly bound molecules at a magnetic field of $\sim 545.9$ G. The coherent transfer is efficient and does not heat the gas, resulting in a gas of rovibronic ground-state molecules with an average number density of $10^{12}$ to $10^{13}$ cm$^{-3}$ and a translational temperature of a few hundred nanokelvin. At this ultralow temperature, even the tiny molecular hyperfine-state energy splittings are much larger than the translational energy. Manipulation of the hyperfine states hence becomes extremely important in exploring possible collision channels. In addition, complete control over the internal quantum state of the molecules permits direct observation of the role of quantum statistics in determining the molecular interactions.

Table 1. Summary of the relevant molecular energetics involved in possible chemical reactions. The binding energies are given with respect to the threshold energy for free atoms in the absence of a magnetic field. The $^{87}{\text{Rb}}_2$ and $^{40}{\text{K}}_2$ binding energies include isotope shifts from the data in the respective references. The trimer binding energies are unknown. Calculations of trimer binding energies are needed and will be important for future experiments on any bi-alkali species.

<table>
<thead>
<tr>
<th>Molecule</th>
<th>$\nu = 0$ binding energy (D$_\nu$)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>$^{87}{\text{Rb}}_2$</td>
<td>3965.8 (±0.4) cm$^{-1}$</td>
<td>(45)</td>
</tr>
<tr>
<td>$^{40}{\text{K}}^{87}{\text{Rb}}$</td>
<td>4180.417 cm$^{-1}$</td>
<td>(29)</td>
</tr>
<tr>
<td>$^{40}{\text{K}}_2$</td>
<td>4450.389 (±0.004) cm$^{-1}$</td>
<td>(46)</td>
</tr>
<tr>
<td>$K_2{\text{Rb}}$</td>
<td>Unknown</td>
<td></td>
</tr>
<tr>
<td>$K_2{\text{Rb}}_2$</td>
<td>Unknown</td>
<td></td>
</tr>
</tbody>
</table>

To probe the quantum nature of ultracold molecular collisions and chemical reactions, we begin by preparing the KRb molecules in a single nuclear spin state of the rovibronic ground state.

**Fig. 2.** Inelastic collisions between spin-polarized (indistinguishable) or different spin-state (distinguishable) fermionic molecules in the rovibronic ground state of $^{40}{\text{K}}^{87}{\text{Rb}}$. (A) Sample data showing the time dependence of the molecule number density. Here the molecules are prepared in a single hyperfine state, $|{-4, 1/2}\rangle$, and the molecular density decays slowly with a rate coefficient of $3.3 (±0.7) \times 10^{-12}$ cm$^{-3}$ s$^{-1}$ at $T = 250$ nK. (B) Loss rate coefficient versus temperature. The loss rate increases linearly with temperature for spin-polarized molecules, which collide via $p$-wave (inset in (A)) at low temperature. Data were taken for molecules prepared in either $|{-4, 1/2}\rangle$ (solid circles) or the lowest-energy state $|{-4, 3/2}\rangle$ (open triangles). A linear fit (solid line) to the $|{-4, 1/2}\rangle$ data yields the temperature-dependent loss rate to be $1.2 (±0.3) \times 10^{-5}$ cm$^{-3}$ s$^{-1}$ at $T = 250$ nK. For the $|{-4, 3/2}\rangle$ case, where the collisional loss can only be due to chemically reactive scattering, the loss rate is similar. The dotted and dashed lines are theoretical predictions from the QT model and MQDT, respectively. When the molecules are prepared in a mixture of the $|{-4, 1/2}\rangle$ and $|{-4, 3/2}\rangle$ states (solid squares), we observe a temperature-independent decay rate that is 10 to 100 times that for the spin-polarized case. The error bars represent 1 SD of the decay rate coefficients arising from fluctuations of the molecular density, temperature, and fitting uncertainty of the two-body loss curves.

**Fig. 3.** A summary of the essential role that quantum statistics and quantum measurements play in determining the rates of inelastic collisions. Our experimental observations confirm a universal loss mechanism.

References

1. The starting point for this work is an ultracold trapped gas of fermionic $^{40}{\text{K}}^{87}{\text{Rb}}$ molecules prepared in a single hyperfine level of the rovibronic ground state ($N = 0, v = 0$ of $X\Sigma^+$) (29, 30). The optical trap depth is $-k_B T_{\text{trap}} = 10 \times 10^{-12}$ cm$^{-3}$, where $k_B$ is Boltzmann’s constant. The molecules are produced using a single step of two-photon Raman transfer from extremely weakly bound molecules at a magnetic field of $\sim 545.9$ G. The coherent transfer is efficient and does not heat the gas, resulting in a gas of rovibronic ground-state molecules with an average number density of $10^{12}$ to $10^{13}$ cm$^{-3}$ and a translational temperature of a few hundred nanokelvin. At this ultralow temperature, even the tiny molecular hyperfine-state energy splittings are much larger than the translational energy. Manipulation of the hyperfine states hence becomes extremely important in exploring possible collision channels. In addition, complete control over the internal quantum state of the molecules permits direct observation of the role of quantum statistics in determining the molecular interactions.

2. A precise measurement of the hyperfine structure and the manipulation of individual hyperfine state populations in the $X\Sigma^+$ ground state of $^{40}{\text{K}}^{87}{\text{Rb}}$ were reported in (30). The $X\Sigma^+$ state has zero total electronic angular momentum, so that the hyperfine structure is basically the Zeeman effect of the nuclear spins $I^K = 4$ and $I^{87}\text{Rb} = 3/2$ (12, 31) at the applied magnetic field. The hyperfine structure is derived in Fig. 1, where a total of 36 states are labeled by their projections of the individual nuclear spins, $m_{87}\text{Rb}$ and $m_K$. For the current study, we produce molecules either in a single spin state ($m_{87}\text{Rb}^2 m_K^2$) or in a mixture of two spin states. The hyperfine states used are an excited state $|{-4, 1/2}\rangle$ and the lowest-energy spin state $|{-4, 3/2}\rangle$; these are marked by the two ellipses in Fig. 1. The $|{-4, 1/2}\rangle$ state is populated directly by the two-photon Raman transfer starting from the weakly bound molecules (29, 30). The spin state of these molecules can be further manipulated coherently. For example, the entire $|{-4, 1/2}\rangle$ population can be transferred into the lowest hyperfine state, $|{-4, 3/2}\rangle$, using two successive $\pi$ pulses through a rotationally excited $N = 1$ intermediate level. The $N = 1$ level has strong nuclear electric quadrupole interactions that couple rotations with nuclear spins, enabling nuclear spin flips (30, 31). We can probe molecules in any particular hyperfine state by reversing the entire transfer process and putting the population back into the initial weakly bound state. We then use high–signal-to-noise ratio absortion imaging to measure the molecular gas number and temperature.

3. The ability to control molecular internal states including the electronic, vibrational, rotational, and nuclear spin degrees of freedom, and in particular the possibility of preparing them in the lowest-energy state, allows us to probe inelastic collisions in a way that limits unwanted loss mechanisms and reduces ambiguities in the identification of possible chemical reaction channels. This is crucial because it is difficult in the ultracold gas experiment to find probes for the direct observation of reaction products. In Table 1, we consider the binding energies for various types of molecules made from different combinations of $^{40}{\text{K}}$ and $^{87}{\text{Rb}}$ atoms. These energy estimates allow us to assess whether a specific two-body reaction process is endothermic or exothermic. We note that all results reported here were obtained in the absence of any external electric field, and hence the effective molecular dipole moment in the lab frame is zero.

4. To probe the quantum nature of ultracold molecular collisions and chemical reactions, we begin by preparing the KB molecules in a single nuclear spin state of the rovibronic ground state.
All unpaired atoms that remain after the initial stage of the molecular creation process are selectively removed from the optical trap via resonant light scattering (32). From an argument based on the energetics summarized in Table 1, the molecule-molecule collisions have a possible exothermic chemical reaction, namely KrB + KrB → K₂ + Rb₂, which releases ~10⁻¹⁰ cm⁻³ of kinetic energy. The reactions KrB + KrB → K₂ + Rb + Rb and KrB + Rb + Rb → KrB + K + K could also be exothermic. All of these reactions require breaking and making molecular bonds. If the KrB molecules are prepared in an excited hyperfine state, spin relaxation to a lower hyperfine state provides an additional inelastic scattering mechanism.

The quantum statistics of the molecules plays an essential role in collisions at a temperature of a few hundred nanokelvin, where collisions with large-impact parameters and correspondingly large centrifugal barriers are frozen out and the collisions are typically dominated by a single large partial wave with orbital angular momentum quantum number $L = 0$ (s-wave) or $L = 1$ (p-wave). Our KrB molecules are fermions, and therefore the total wave function describing a KrB + KrB collision is antisymmetric with respect to molecular exchange. For spin-polarized molecules all prepared in exactly the same internal quantum state, the p-wave is the lowest-energy symmetry-allowed collision channel. The height of the centrifugal barrier for the $L = 1$ KrB-KrB collisions is $\hbar^2 \times 24 \mu K (33, 34)$. This barrier height is more than an order of magnitude larger than $\hbar^2 T$, where $T$ is the translational temperature of the molecular gas. Thus, collisions of spin-polarized molecules are expected to proceed predominantly via tunneling through the p-wave barrier. Note that $T$ is greater than 1.4 times the Fermi energy. If two molecules make it through the barrier to short range, chemical reactions or hyperfine state-changing collisions can take place, leading to a loss of the entrance channel population. We note that even a single nuclear spin flip corresponds to a released quantity of energy that is above the trap depth, and this would contribute to loss of trapped molecules.

The quantum nature of the collisions can be seen in the temperature dependence of loss rates. The Bethe-Wigner threshold law predicts that the p-wave inelastic/reactive collision rate should be linear in temperature ($\propto T$). To look for this behavior, we first prepared spin-polarized molecules in the single hyperfine state $|\uparrow, 1/2\rangle$ for various values of $T$ ranging from 200 to 900 nK (35). The temperature is measured from the expansion energy of the molecular gas after releasing it from the optical trap. For each initial temperature, we observed the time-dependent molecular loss (36) and extracted a two-body loss rate $\beta$ (which is twice the collisional event rate) by fitting the measured decay of the molecular gas density $n$ versus time $t$ (Fig. 2A) to

$$\frac{dn}{dt} = -\beta n^2 - an$$  (1)

Here, the first term on the right accounts for number loss and the measured $\beta$ can be compared to theoretical predictions. The second term accounts for density change due to heating of the trapped gas during the measurement. Within a single measurement, we observe an increase in temperature that is at most 30%. In the analysis for each data set, we fit the measured temperature to a linear heating rate and obtain a constant slope $c$. In Eq. 1, we then use $a = (3/2)[c/(T + c)]$, where $T$ is the initial temperature. At our lowest temperature of 250 nK, the heating was 7 (2) ms (37). Using this assumption, we can compare the predicted p-wave threshold law to that measured for molecules in the $|\uparrow, 1/2\rangle$ state.

To understand the loss rates, we use two models: a simple quantum threshold model (QT), and a model that uses the formalism of multichannel quantum defect theory (MQDT). In the QT model, the loss rate for collisions with energy equal to or above the height of the p-wave barrier is determined by the Langevin capture rate (37), which assumes that the probability for chemical reactions and/or hyperfine state-changing collisions is unity. For all collision energies below the height of the p-wave barrier, we assume in this model that the loss follows the Bethe-Wigner threshold laws (7, 8). Using this assumption, we obtain a simple analytical expression for the p-wave loss rate coefficient of two indistinguishable molecules, which scales linearly with $T$ (38):

$$\beta = \frac{\pi}{4} \left( \frac{3^{1/4} \mu^3 \sigma^3}{\hbar^4} \right)^{1/4} k \hbar T$$  (2)

where $\mu$ is the reduced mass and $\hbar$ is Planck’s constant divided by 2π. Using a van der Waals dispersion coefficient of $C_6 = 16130$ a.u. for KrB-KrB with an uncertainty of ±10% (33, 34), the slope of the rate coefficient is predicted to be 1.5 (±0.1) $\times 10^{-5}$ cm³ s⁻¹ K⁻¹, which agrees well with the experimental measurement.

In the second model, the loss rate coefficient is found directly by calculating the
quantum tunneling rate through the p-wave barrier (39). This calculation gives $\beta = 0.8 \times (0.1) \times 10^{-3} \text{ cm}^2 \text{s}^{-1} \text{ K}^{-1}$, which agrees with the experiment within mutual uncertainties. This $\beta$ can also be derived analytically from the properties of the long-range potential to give $\beta = (11.48 \pi) \left( k_0 T/h \right)$, where $\pi = 0.4778 \left( 2\mu C_6/h^2 \right)^{1/4} = 6.3 \text{ nm}$ is the characteristic length of the van der Waals potential (12). This fully quantum calculation can be put in the same form as the QT model (Eq. 2) and gives a $\beta$ that is smaller by a factor of 0.528. With these simple theories, the agreement with our molecule-molecule collisional loss measurements suggests that the chemical reaction rates are strongly influenced by the long-range interactions. This observation opens intriguing control possibilities because the long-range interaction can be controlled by selecting quantum states and tuning collision energies via applied electric and magnetic fields.

Reaction rates should be markedly different if molecules are prepared in a mixture of different hyperfine states as s-wave scattering becomes allowed. We measured the inelastic collision rates for rovibronic ground-state molecules that were prepared in a roughly 50-50 incoherent mixture of the two hyperfine spin states $| -4, 3/2 \rangle$ and $| -4, 1/2 \rangle$. The time-dependent number density of trapped molecules was measured for both spin states. We observed the same loss rate for both states, consistent with loss due to collisions between distinguishable molecules in different spin states. The rate coefficient is determined to be $1.9 \times (0.4) \times 10^{-10} \text{ cm}^3 \text{s}^{-1}$, independent of temperature (solid squares in Fig. 2B). In comparison to our measurements for p-wave collisions between spin-polarized molecules, the s-wave collision rate between molecules in different hyperfine states is higher by a factor of 10 to 100 for a similar temperature range.

The MQDT model can also be used to estimate collision rates where the dominant collision channel is s-wave. Here, the relevant length scale is determined by the inter-molecular van der Waals potential without any angular momentum barrier (40). We assume that when the molecules approach each other within the van der Waals length $\bar{r}$, chemical reactions take place and remove these entrant molecules with a near-unity probability. The universal loss rate coefficient, $\beta = (2h/\mu)^{1/2} \bar{r}$ (12), predicts a $\beta$ value of $0.8 \times 10^{-16} \text{ cm}^2 \text{s}^{-1}$, which is a factor of 2 lower than the experimentally observed value. This difference suggests that short-range physics may play some role in the loss dynamics. An enhancement in the rate coefficient (up to the energy-dependent unitarity limit, which is $4 \times 10^{-16} \text{ cm}^2 \text{s}^{-1}$ for a gas at 400 nK) is possible if there is a partial reflection of the colliding species back into the entrance channel (39). The reflected amplitude interferes with the incoming amplitude and can either increase or decrease the rate coefficient from its “universal” value. Additional theory and experiment are needed to explore this possibility.

In addition to molecule-molecule reactions, the relatively long lifetime of a pure gas of spin-polarized molecules in the optical trap (~1 s) affords time to look for chemical reactions between atoms and molecules. The MQDT model described above can also predict the atom-molecule reaction rates determined by the long-range physics for the universal loss mechanism. To prepare the atom-molecule mixture, we control the atom density by selectively removing or heating unpaired atoms after the initial molecule creation (41). For these experiments, we typically work with an atom number about 5 to 15 times the molecule number. All atoms and molecules are prepared in their lowest-energy states at 545.9 G. Specifically, K atoms are in their $|F = 9/2, m_F = -9/2 \rangle$ state, Rb in $|F = 1, m_F = 1 \rangle$, and KRb in $|F = 4, 3/2 \rangle$. Here $F$ is the total atomic spin and $m_F$ is the spin projection. We performed two separate experiments, one with K and KRb and the second with Rb and KRb, at a temperature less than 1 µK. In both cases, the background of atoms in the other, undesired, species was less than 1000, corresponding to a density below $5 \times 10^{-10}$ cm$^{-3}$. Because both the atoms and the molecules are prepared in their lowest-energy states, trap loss due to inelastic spin-changing collisions is not possible. However, from Table 1, we do expect loss from chemical reactions for the exothermic reaction $\text{K} + \text{KRb} \rightarrow \text{K}_2 + \text{Rb}$, whereas the endothermic $\text{Rb} + \text{KRb} \rightarrow \text{Rb}_2 + \text{K}$ should be forbidden.

For each experiment, we measured the time dependence of the trapped molecule population. Typical molecular loss curves are shown in Fig. 3A. To extract the inelastic collision rate, we assume that the atom number density is constant. (This is approximately true because the number of atoms is much larger than the number of molecules.) The trapped molecule number should then decay as

$$\frac{d}{dt}N_{\text{molecule}} = -\beta \times N_{\text{molecule}} \times n_{\text{atom}}$$

where $N_{\text{molecule}}$ is the number of molecules, $n_{\text{atom}}$ is the atomic density, and $\beta$ is the inelastic rate coefficient. We can then extract $\beta$ via an exponential fit, $\exp(-\beta \times n_{\text{atom}} \times t)$. As can be seen in Fig. 3B, in general we find that the molecules are lost from the trap at a much faster rate when K atoms are present than when a similar density of Rb is present.

To see if the molecule loss arises from atom-molecule collisions, we measure the dependence of the loss rate on the atom gas density. For the case of KRb + K, we observe a clear linear dependence on the atomic density and extract an inelastic collision rate coefficient of $1.7 \pm (0.3) \times 10^{-10} \text{ cm}^3 \text{s}^{-1}$. This rate coefficient (due to chemical reactions) can again be predicted from the van der Waals length characterizing the long-range part of the potential between the collision partners. Using methods similar to those used for KRb + KRb, Kotochigova has calculated $C_6$ for KRb + K to be 7020 (±700) a.u. (34), which gives a predicted reaction rate of $1.1 \times 10^{-10} \text{ cm}^3 \text{s}^{-1}$. Again we find that the MQDT value is close but somewhat lower than the measured value; this result further solidifies the importance of long-range quantum scattering in the inelastic process.

In contrast, for KRb + Rb collisions, the density dependence of the loss rate is not obvious (Fig. 3B). Nevertheless, we again fit the dependence as linear and obtain an upper limit for the rate coefficient of $0.13 \pm (0.04) \times 10^{-10} \text{ cm}^3 \text{s}^{-1}$, which is one order of magnitude smaller than what we measure for KRb + K. Our measurement is consistent with the fact that for KRb + Rb, there is no two-body chemical reaction pathway. A possible mechanism for the residual nonzero rate coefficient for KRb + Rb would be collisions of ground-state molecules with undetected molecules in high-lying vibrationally excited states. These contaminant molecules could be produced by inelastic collisions of Rb atoms with our weakly bound KRb molecules (42) before the two-photon Raman process that produces ground-state molecules. Another possible loss mechanism is Rb + Rb + KRb three-body inelastic collisions. Further experiments will be needed to check these possibilities. We note that this suppressed loss rate is observed only when both KRb and Rb are prepared in their lowest-energy internal states at 545.9 G. If either the molecules or the atoms are in an excited hyperfine state, then the observed inelastic rate coefficient rises again to the order of $10^{-10} \text{ cm}^3 \text{s}^{-1}$. Therefore, for practical reasons, removing Rb atoms is important for creating a long-lived sample of ground-state KRb molecules.

Together the studies presented here show that we have observed barrierless chemical reactions in the short range, with the rates determined by long-range scattering dynamics dictated by quantum statistics, angular momentum barriers, and threshold laws. We see that a change as seemingly insignificant as flipping a single nuclear spin dramatically changes the rate of molecular collisions. A pure gas of spin-polarized KRb molecules is long-lived in an optical trap (surviving for a time on the order of 1 s) whereas a spin mixed KRb sample decays 10 to 100 times as quickly. Our results clearly show that chemical reactions can proceed with high rates in the ultracold regime.

References and Notes
Low-Frequency Modes of Aqueous Alkali Halide Solutions: Glimpsing the Hydrogen Bonding Vibration

Ismael A. Heisler and Stephen R. Meech*

The solvation of ions in aqueous media is a fundamental process in biology and chemistry. Here, we report direct time-domain observations of the hydrogen bond vibrational mode formed between a halide ion (chloride, bromide, or iodide) and the surrounding water molecules. The frequency of the hydrogen bond mode is sensitive to both the atomic weight and the concentration of the ion. The peak frequencies fall in the 125 to 175 wave-number range, a spectral region of the spectra of the OH···X bonds are drawn. However, the H-bond mode is not well characterized, and a more complete description of it will help to refine the analysis and simulation of aqueous solvation dynamics.

Here, we report direct time-domain experimental observations of the spectra of the OH···X hydrogen bond through measurements of the low-frequency isotropic Raman response. To access the low-frequency region of the spectrum, where H-bond modes are expected to appear, we probed in the time domain the decay of polarization induced in the sample by an ultrafast pump pulse. In a two-beam pump-probe configuration, this experiment measures the transient optical Kerr effect, which, when combined with heterodyne detection and Fourier transform analysis, yields the Raman spectral density with excellent signal-to-noise ratio in the 0- to 500-cm⁻¹ region. In the present experiments, we exploited a four-beam transient grating diffractive optic geometry, described in detail by Goodno et al. (17). Essentially, pump and probe beams are focused onto a diffractive optic element to fluctuations in the length of the OD···X hydrogen bond (H-bond), where X is the anion. The intermediate (subpicosecond) and longer (picosecond) relaxation components, which increase with increasing halide ion concentration, reflect the transition from local H-bond fluctuations to global reorganization and the reorganization of H-bond networks, respectively. From these studies of the OD mode, inferences on the structure and dynamics of the OD···X H-bonds are drawn. However, the H-bond mode itself is not well characterized, and a more complete description of it will help to refine the analysis and simulation of aqueous solvation dynamics.

1. To remove selective amounts of K, we partially transfer the population of K atoms from |F = 9/2, m_F = –9/2, –5/2⟩ using two successive radio-frequency (rf) pulses. We then clean out K atoms in the 9/2, –9/2 state using resonant light as in (32). Once these atoms are heated out of the trap, we again use rf pulses to spin-flip all K |9/2, –5/2⟩ atoms back down to |9/2, –9/2⟩. To selectively vary the density of RB, we perform a complete spin flip from |1⟩, 1⟩ to |2⟩, 2⟩ and then turn on resonant light for a duration between 1 and 10 μs. The resonant light both removes RB atoms and heats the remaining gas. After the resonant light pulse, RB atoms are transferred back to the lowest hyperfine state |1⟩, 2⟩. Then we apply an additional resonant light pulse to remove any residual RB in |2⟩, 2⟩, thus ensuring that all RB atoms are in the |1⟩, 1⟩ state.
7. Supported by NSF Innovations in Measurement Science—Ultracold Stable Molecules, NSF Physics Frontier Center at JILA, U.S. Department of Energy, Air Force Office of Scientific Research MURI on Ultracold Molecules, and a NSF graduate fellowship (B.N.) and by Office of Naval Research grant N00014091P20041 (P.S.). We thank S. Kotochigova for providing her calculations of the van der Waals dispersion coefficients that are relevant to our experiments.
8. November 2009; accepted 28 December 2009 10.1126/science.1184121

School of Chemistry, University of East Anglia, Norwich NR4 7J, UK.
*To whom correspondence should be addressed. E-mail: s.meech@uea.ac.uk

The frequency of the hydrogen bond mode is sensitive to both the atomic weight and the concentration of the ion. The peak frequencies fall in the 125 to 175 wave-number range, a spectral region of the spectra of the OH···X bonds are drawn. However, the H-bond mode is not well characterized, and a more complete description of it will help to refine the analysis and simulation of aqueous solvation dynamics.

Here, we report direct time-domain experimental observations of the spectra of the OH···X hydrogen bond through measurements of the low-frequency isotropic Raman response. To access the low-frequency region of the spectrum, where H-bond modes are expected to appear, we probed in the time domain the decay of polarization induced in the sample by an ultrafast pump pulse. In a two-beam pump-probe configuration, this experiment measures the transient optical Kerr effect, which, when combined with heterodyne detection and Fourier transform analysis, yields the Raman spectral density with excellent signal-to-noise ratio in the 0- to 500-cm⁻¹ region. In the present experiments, we exploited a four-beam transient grating diffractive optic geometry, described in detail by Goodno et al. (17). Essentially, pump and probe beams are focused onto a diffractive optic element to fluctuations in the length of the OD···X hydrogen bond (H-bond), where X is the anion. The intermediate (subpicosecond) and longer (picosecond) relaxation components, which increase with increasing halide ion concentration, reflect the transition from local H-bond fluctuations to global reorganization and the reorganization of H-bond networks, respectively. From these studies of the OD mode, inferences on the structure and dynamics of the OD···X H-bonds are drawn. However, the H-bond mode itself is not well characterized, and a more complete description of it will help to refine the analysis and simulation of aqueous solvation dynamics.

Here, we report direct time-domain experimental observations of the spectra of the OH···X hydrogen bond through measurements of the low-frequency isotropic Raman response. To access the low-frequency region of the spectrum, where H-bond modes are expected to appear, we probed in the time domain the decay of polarization induced in the sample by an ultrafast pump pulse. In a two-beam pump-probe configuration, this experiment measures the transient optical Kerr effect, which, when combined with heterodyne detection and Fourier transform analysis, yields the Raman spectral density with excellent signal-to-noise ratio in the 0- to 500-cm⁻¹ region. In the present experiments, we exploited a four-beam transient grating diffractive optic geometry, described in detail by Goodno et al. (17). Essentially, pump and probe beams are focused onto a diffractive optic element to fluctuations in the length of the OD···X hydrogen bond (H-bond), where X is the anion. The intermediate (subpicosecond) and longer (picosecond) relaxation components, which increase with increasing halide ion concentration, reflect the transition from local H-bond fluctuations to global reorganization and the reorganization of H-bond networks, respectively. From these studies of the OD mode, inferences on the structure and dynamics of the OD···X H-bonds are drawn. However, the H-bond mode itself is not well characterized, and a more complete description of it will help to refine the analysis and simulation of aqueous solvation dynamics.

Here, we report direct time-domain experimental observations of the spectra of the OH···X hydrogen bond through measurements of the low-frequency isotropic Raman response. To access the low-frequency region of the spectrum, where H-bond modes are expected to appear, we probed in the time domain the decay of polarization induced in the sample by an ultrafast pump pulse. In a two-beam pump-probe configuration, this experiment measures the transient optical Kerr effect, which, when combined with heterodyne detection and Fourier transform analysis, yields the Raman spectral density with excellent signal-to-noise ratio in the 0- to 500-cm⁻¹ region. In the present experiments, we exploited a four-beam transient grating diffractive optic geometry, described in detail by Goodno et al. (17). Essentially, pump and probe beams are focused onto a diffractive optic element to fluctuations in the length of the OD···X hydrogen bond (H-bond), where X is the anion. The intermediate (subpicosecond) and longer (picosecond) relaxation components, which increase with increasing halide ion concentration, reflect the transition from local H-bond fluctuations to global reorganization and the reorganization of H-bond networks, respectively. From these studies of the OD mode, inferences on the structure and dynamics of the OD···X H-bonds are drawn. However, the H-bond mode itself is not well characterized, and a more complete description of it will help to refine the analysis and simulation of aqueous solvation dynamics.

Here, we report direct time-domain experimental observations of the spectra of the OH···X hydrogen bond through measurements of the low-frequency isotropic Raman response. To access the low-frequency region of the spectrum, where H-bond modes are expected to appear, we probed in the time domain the decay of polarization induced in the sample by an ultrafast pump pulse. In a two-beam pump-probe configuration, this experiment measures the transient optical Kerr effect, which, when combined with heterodyne detection and Fourier transform analysis, yields the Raman spectral density with excellent signal-to-noise ratio in the 0- to 500-cm⁻¹ region. In the present experiments, we exploited a four-beam transient grating diffractive optic geometry, described in detail by Goodno et al. (17). Essentially, pump and probe beams are focused onto a diffractive optic element to fluctuations in the length of the OD···X hydrogen bond (H-bond), where X is the anion. The intermediate (subpicosecond) and longer (picosecond) relaxation components, which increase with increasing halide ion concentration, reflect the transition from local H-bond fluctuations to global reorganization and the reorganization of H-bond networks, respectively. From these studies of the OD mode, inferences on the structure and dynamics of the OD···X H-bonds are drawn. However, the H-bond mode itself is not well characterized, and a more complete description of it will help to refine the analysis and simulation of aqueous solvation dynamics.

Here, we report direct time-domain experimental observations of the spectra of the OH···X hydrogen bond through measurements of the low-frequency isotropic Raman response. To access the low-frequency region of the spectrum, where H-bond modes are expected to appear, we probed in the time domain the decay of polarization induced in the sample by an ultrafast pump pulse. In a two-beam pump-probe configuration, this experiment measures the transient optical Kerr effect, which, when combined with heterodyne detection and Fourier transform analysis, yields the Raman spectral density with excellent signal-to-noise ratio in the 0- to 500-cm⁻¹ region. In the present experiments, we exploited a four-beam transient grating diffractive optic geometry, described in detail by Goodno et al. (17). Essentially, pump and probe beams are focused onto a diffractive optic element to fluctuations in the length of the OD···X hydrogen bond (H-bond), where X is the anion. The intermediate (subpicosecond) and longer (picosecond) relaxation components, which increase with increasing halide ion concentration, reflect the transition from local H-bond fluctuations to global reorganization and the reorganization of H-bond networks, respectively. From these studies of the OD mode, inferences on the structure and dynamics of the OD···X H-bonds are drawn. However, the H-bond mode itself is not well characterized, and a more complete description of it will help to refine the analysis and simulation of aqueous solvation dynamics.