Infrared Spectroscopy of Hydrated Bicarbonate Anion Clusters: \( \text{HCO}_3^-(\text{H}_2\text{O})_{1-10} \)

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Abstract: Infrared multiple photon dissociation spectra are reported for \( \text{HCO}_3^-(\text{H}_2\text{O})_{1-10} \) clusters in the spectral range of \( 600-1800 \text{ cm}^{-1} \). In addition, electronic structure calculations at the MP2/6-311+G(d,p) level have been performed on the \( n = 1-8 \) clusters to identify the structure of the low-lying isomers and to assign the observed spectral features. General trends in the stepwise solvation motifs of the bicarbonate anion can be deduced from the overall agreement between the calculated and experimental spectra. The most important of these is the strong preference of the water molecules to bind to the negatively charged \( \text{CO}_2 \) moiety of the \( \text{HCO}_3^- \) anion. However, a maximum of four water molecules interact directly with this site. The binding motif in the most stable isomer of the \( n = 4 \) cluster, a four-membered ring with each water forming a single H-bond with the \( \text{CO}_2 \) moiety, is retained in all of the lowest-energy isomers of the larger clusters. Starting at \( n = 6 \), additional solvent molecules are found to form a second hydration layer, resulting in a water–water network bound to the \( \text{CO}_2 \) moiety of the bicarbonate anion. Binding of a water to the hydroxyl group of \( \text{HCO}_3^- \) is particularly disfavored and apparently does not occur in any of the clusters investigated here. Similarities and differences with the infrared spectrum of aqueous bicarbonate are discussed in light of these trends.

1. Introduction

The bicarbonate anion, \( \text{HCO}_3^- \), is a ubiquitous species in aqueous chemistry. Because bicarbonate is a weak base, it plays a central role in the acid–base equilibrium formed when \( \text{CO}_2 \) is dissolved in water:

\[
\text{CO}_2 + \text{H}_2\text{O} \leftrightarrow \text{H}_2\text{CO}_3 \leftrightarrow \text{HCO}_3^- + \text{H}^+ \leftrightarrow \text{CO}_3^{2-} + 2\text{H}^+
\]

(1)

This species is thus important in processes such as pH homeostasis in oceans,\(^1\) the formation of external skeletons by calcifying organisms,\(^2\) the transport of \( \text{CO}_2 \) between the metabolizing tissues and the lungs,\(^3\) and the regulation of blood pH. The nucleophilic attack of the hydroxide anion on \( \text{CO}_2 \):

\[
\text{OH}^- + \text{CO}_2 \rightarrow \text{HCO}_3^- \quad (2)
\]

is a benchmark for the study of solvation effects on chemical reaction dynamics.\(^4\)–\(^9\) While this reaction is predicted to proceed without a reaction barrier in the gas-phase,\(^4,6,10\) a substantial enthalpic barrier of 13.0 ± 0.6 kcal/mol has been reported in aqueous solution.\(^1,11,12\) Solvent interactions with the reactants, products, and complex along the reaction coordinate are thus important in understanding the thermodynamics and kinetics of this reaction in the aqueous phase. In this paper, we report gas-phase infrared spectra of the \( \text{HCO}_3^- \cdot (\text{H}_2\text{O})_{1-10} \) clusters. These spectra, in combination with the accompanying electronic structure calculations, provide a detailed probe of bicarbonate anion microsolvation.

The gas-phase thermochemical properties of the bare bicarbonate anion have been studied via collision-induced dissociation and proton transfer reactions.\(^13\) The \( \text{HCO}_3^- \) anion has been studied by Raman and infrared spectroscopy in aqueous solution\(^14\)–\(^18\) and in the crystalline phase.\(^17,19\) Analysis of

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these spectra is complicated by possible counterion interactions and formation of bicarbonate dimers. Rudolph and co-workers have studied the infrared and Raman spectra of very dilute solutions of KHCO₃ and KDCO₃ and concluded that the hydrated HCO₃⁻ anion possess a C₃ symmetry, in contrast to the Cₛ (planar) symmetry expected in the gas-phase.¹⁷,¹⁸ They also reported density functional theory (DFT) calculations on the HCO₃⁻·(H₂O)ₙ₋₁ clusters.¹⁷ While these bulk studies probe the structure of the HCO₃⁻ in solution, they do not provide detailed information on the microscopic structure of the solvation shells or the specific interactions between the solvent molecules and the anion. On the other hand, infrared spectroscopy of size-selected ion clusters²¹–²⁵ has emerged in recent years as a very powerful technique for probing microscopic ion hydration.

Here, we use vibrational action spectroscopy of gas-phase clusters to probe the microhydration of the HCO₃⁻ anion by studying the effect of stepwise hydration on its spectroscopy, structure, and energetics. We measure infrared multiple photon dissociation spectra of mass-selected HCO₃⁻·(H₂O)$_n$ clusters at low temperature and in the absence of a counterion. The spectra are acquired in the spectral region spanning 550–1800 cm⁻¹, which includes five of the characteristic vibrational modes of the anion as well as the libration and bending modes of the solvent molecules. The spectra are analyzed with the help of simulated linear absorption spectra from electronic structure calculations. This combination of experiment and theory has proved very useful in recent studies²⁶,²⁷ on the microhydration of the SO₄²⁻ and NO₃⁻ anions.

2. Experimental Details

Infrared multiple photon photodissociation (IRMPD) experiments²⁶,²⁹ were carried out on a previously described ring electrode trap/time-of-flight mass spectrometer,³⁰ which was temporarily installed at the “Free Electron Laser for Infrared eXperiments” (FELIX) user facility³¹ in the FOM Institute Rijnhuizen (The Netherlands). HCO₃⁻·(H₂O)$_n$ ions were produced by electrospray of a 5 mM solution of sodium bicarbonate (NaHCO₃) in a 1:1 mixture of water and acetonitrile using a modified Waters z-spray source. The negative ions were transferred into a high vacuum system, sampled by a 3 mm diameter skimmer, and focused into a radio frequency (RF) linear ion guide. Ions of interest were then mass selected in a quadrupole mass filter, reflected by 90° using an electrostatic quadrupole ion deflector, and focused into a linear, ring electrode RF ion trap. The ion trap was filled with He buffer gas and cooled to 12 K by means of a closed cycle He cryostat in order to cool the ions by collisions. The ions were accumulated and cooled in the trap for 195 ms. They were then brought into the extraction region of a time-of-flight (TOF) mass spectrometer and irradiated with the pulsed radiation from FELIX prior to the application of high voltage pulses to the TOF plates. FELIX macropulses were used at a repetition rate of 5 Hz, a pulse length of 5 µs, a laser wavelength bandwidth of ∼0.25% rms over the range from 550 to 1800 cm⁻¹, and typical pulse energies of up to 50 mJ.

3. Results

IRMPD spectra of HCO₃⁻·(H₂O)$_n$ clusters, with n = 1–10 are shown in Figure 1. These spectra were generated by summing the yield of the various fragment ions as a function of the photodissociation laser wavelength and reflect the absorption spectrum of the parent ion. This signal was nearly background-free, in contrast to monitoring depletion of the parent ion signal. The only fragmentation pathways observed involved loss of one or more water molecules. The loss of more than one water molecule might have been due in part to fragment absorption as the different cluster sizes have similar infrared absorption bands. No fragmentation was observed for the bare HCO₃⁻ anion.

The experimental IRMPD spectra comprise six bands labeled A–F in Figure 1. Peak B, which appears as a small shoulder for the n = 1 cluster, red-shifts from 860 cm⁻¹ to 825 cm⁻¹ upon addition of the second water. This peak subsequently blue-shifts by only 16 cm⁻¹ from n = 3–10. Peak C blue-shifts from 887 cm⁻¹ to 986 cm⁻¹ as the number of waters increases. Peak D moves from 1177 cm⁻¹ to 1205 cm⁻¹ between HCO₃⁻·(H₂O)$_n$ and HCO₃⁻·(H₂O)$_2$ and is not observed for larger clusters. Peak

![Figure 1](https://example.com/figure1.png)

**Figure 1.** Experimental IRMPD spectra of HCO₃⁻·(H₂O)$_n$ ions with n = 1–10. The total fragment ion yield is plotted as a function of irradiation wavelength (cm⁻¹).

structures of HCO\textsuperscript{3-}

Geometry optimizations were performed for a large number of initial computational limitations, only the n to particular solvation motifs of the HCO\textsuperscript{3-} mental frequencies for HCO\textsuperscript{3-} employing the B3LYP hybrid exchange-correlation functional\textsuperscript{33,34}

in the spectra of HCO\textsuperscript{3-}

Figure 2.


(35) Scott, A. P.; Radom, L.


herein were derived from the MP2/6-311\textsuperscript{G(d,p)} basis set. Vibration analysis was performed on all the optimized geometries to ensure that the structures corresponded to local minima. The lowest-energy isomers were then reoptimized with the second-order Moller–Plesset (MP2) method using the same 6-311+G(d,p) basis set. The MP2 structures were found to be very similar to those obtained with DFT, but the energetic ordering of the various isomers often changed for the clusters with n ≥ 3. In some instances, small differences in structures between isomers were observed, for instance an H-atom pointing up or down, in which case the lowest-energy form was used, but for the purposes of our study these were not considered to be distinct isomers. Simulated infrared spectra were derived from the MP2/6-311+G(d,p) scaled harmonic frequencies (scaling factor: 0.970)\textsuperscript{35} and linear absorption intensities. For comparison with experimental results, stick spectra from the simulations were convoluted using a Gaussian line shape function with a 15 cm\textsuperscript{-1} full width at half-maximum (fwhm). All the energetics presented herein were derived from the MP2/6-311+G(d,p) values and include the scaled zero-point energy correction.

4. Computational Details

Electronic structure calculations were performed using the Gaussian 03 program\textsuperscript{32} in order to assign the experimental spectra to particular solvation motifs of the HCO\textsuperscript{3-} anion. Because of computational limitations, only the n = 1–8 clusters were studied. Geometry optimizations were performed for a large number of initial structures of HCO\textsuperscript{3-}-(H\textsubscript{2}O)\textsubscript{n} clusters using density functional theory, employing the B3LYP hybrid exchange-correlation functional\textsuperscript{33,34} and the 6-311+G(d,p) basis set. Vibration analysis was performed on all the optimized geometries to ensure that the structures corresponded to local minima. The lowest-energy isomers were then reoptimized with the second-order Moller–Plesset (MP2) method using the same 6-311+G(d,p) basis set. The MP2 structures were found to be very similar to those obtained with DFT, but the energetic ordering of the various isomers often changed for the clusters with n ≥ 3. In some instances, small differences in structures between isomers were observed, for instance an H-atom pointing up or down, in which case the lowest-energy form was used, but for the purposes of our study these were not considered to be distinct isomers. Simulated infrared spectra were derived from the MP2/6-311+G(d,p) scaled harmonic frequencies (scaling factor: 0.970)\textsuperscript{35} and linear absorption intensities. For comparison with experimental results, stick spectra from the simulations were convoluted using a Gaussian line shape function with a 15 cm\textsuperscript{-1} full width at half-maximum (fwhm). All the energetics presented herein were derived from the MP2/6-311+G(d,p) values and include the scaled zero-point energy correction.

Table 1. Peak Positions (in cm\textsuperscript{-1}) in the HCO\textsuperscript{3-}·(H\textsubscript{2}O)\textsubscript{1-10} Experimental IR Spectra\textsuperscript{a}

<table>
<thead>
<tr>
<th>HCO\textsuperscript{3-}·(H\textsubscript{2}O)\textsubscript{n}</th>
<th>band A</th>
<th>band B</th>
<th>band C</th>
<th>band D</th>
<th>band E</th>
<th>band F</th>
</tr>
</thead>
<tbody>
<tr>
<td>ν\textsubscript{a}(a')</td>
<td>OH str.</td>
<td>3752 cm\textsuperscript{-1}</td>
<td>ν\textsubscript{a}(a')</td>
<td>CO\textsubscript{2} asym str.</td>
<td>1734 cm\textsuperscript{-1}</td>
<td>ν\textsubscript{a}(a')</td>
</tr>
<tr>
<td>ν\textsubscript{a}(a')</td>
<td>COOH bend</td>
<td>1711 cm\textsuperscript{-1}</td>
<td>ν\textsubscript{a}(a')</td>
<td>C–OH stretch</td>
<td>873 cm\textsuperscript{-1}</td>
<td>ν\textsubscript{a}(a')</td>
</tr>
<tr>
<td>ν\textsubscript{a}(a')</td>
<td>CO wag.</td>
<td>655 cm\textsuperscript{-1}</td>
<td>ν\textsubscript{d}(a')</td>
<td>CO \textsubscript{out-of-plane}</td>
<td>787 cm\textsuperscript{-1}</td>
<td>ν\textsubscript{a}(a')</td>
</tr>
</tbody>
</table>

\textsuperscript{a} Band assignments are referenced to HCO\textsuperscript{3-} vibrational modes in Figure 2. \textsuperscript{b} From ref 17. \textsuperscript{c} From ref 16. \textsuperscript{d} From ref 36.

E, which is the most intense feature in most of the spectra, blue-shifts from 1288 to 1353 cm\textsuperscript{-1} as the cluster size increases. Peak F is a broader feature whose individual components are partially resolved for only a few cluster sizes. In contrast to the other bands, it generally red-shifts as the number of water molecules increases. Finally, the peak labeled A is a broad, unresolved feature between 600 and 800 cm\textsuperscript{-1} that appears only in the spectra of HCO\textsuperscript{3-}·(H\textsubscript{2}O)\textsubscript{n} and larger clusters. Experimental frequencies for HCO\textsuperscript{3-}·(H\textsubscript{2}O)\textsubscript{1-10} are summarized in Table 1.

5. Analysis

The six bands observed in the IRPMD spectra can be readily assigned on the basis of previous infrared studies of HCO\textsuperscript{3-} in bulk water\textsuperscript{17} and calculated vibrational frequencies of the bare and solvated anion. The normal modes of HCO\textsuperscript{3-} are shown in Figure 2, along with their calculated and scaled gas-phase frequencies at the MP2/6-311+G(d,p) level of theory. The small peaks B and C are assigned to the CO\textsubscript{3} out-of-plane mode (ν\textsubscript{7}) and C–OH stretching mode (ν\textsubscript{9}), respectively. Peak D is assigned to the COH bending mode (ν\textsubscript{8}), whereas the more intense peak E is assigned to the CO\textsubscript{2} symmetric stretch (ν\textsubscript{6}). Peak F is assigned to the CO\textsubscript{2} asymmetric stretch (ν\textsubscript{2}) of the anion core as well as the various water bending modes.\textsuperscript{36} These assignments are summarized in Table 1. Finally, peak A is located in the region where the CO\textsubscript{2} bending mode (ν\textsubscript{6}) is expected. However, this mode has very weak infrared absorption,\textsuperscript{17} and thus band A is instead assigned to the various water librational modes.\textsuperscript{36}

The following sections summarize the theoretical and experimental results for each individual cluster. For each cluster up to n = 8, the structures, energetics, and simulated linear absorption spectra of the three lowest-lying isomers are shown together with the corresponding experimental IRPMD spectra in Figures 3–10. The atom and position labeling used in this section and in section 6 is shown at the bottom of Figure 2. 5.1. HCO\textsuperscript{3-}·(H\textsubscript{2}O). In Figure 3, the lowest-energy isomer, labeled 1-I, is a C\textsubscript{2v} structure in which the water forms two H-bonds with the negatively charged CO\textsubscript{3} moiety of the HCO\textsuperscript{3-} anion. The 1-II isomer lies 3.4 kJ/mol higher and consists of a doubly H-bonded water in which the hydroxyl group oxygen

\textsuperscript{32} Frisch, M. J.; et al. Gaussian 03, Revision C.02; Gaussian, Inc.: Wallingford CT, 2004.


\textsuperscript{34} Becke, A. D. J. Chem. Phys. 1993, 98, 5648.

\textsuperscript{35} Scott, A. P.; Radom, L. J. Phys. Chem. 1996, 100, 16502.

\textsuperscript{36} NIST Mass Spec Data Center; Stein, S. E., Director; Infrared Spectra. In NIST Chemistry WebBook. NIST Standard Reference Database Number 69; Linstrom, P. J., Mallard, W. G., Eds.; National Institute of Standards and Technology; Gaithersburg, MD, http://webbook.nist.gov, retrieved November 2, 2009.)
of the OH and one oxygen of the CO$_2$ act as acceptors. In this isomer, the water molecule is located slightly out of the HCO$_3^-$ plane. In the third isomer, 1-III, the water forms one H-bond with the CO$_2$ moiety and acts as an acceptor for a second H-bond with the OH group. This isomer lies 10.8 kJ/mol above 1-I. No minima were found for isomers with a singly H-bonded water at either the B3LYP/6-311+G(d,p) or MP2/6-311+G(d,p) level of theory.

The simulated infrared spectra of 1-I agrees best with experiment. This isomer yields calculated positions of peaks C$-$F which are in excellent agreement with the experimental values. However, the simulated spectrum of 1-I includes a peak at 719 cm$^{-1}$, corresponding to the water libration mode, that is not observed experimentally. The absence of features in the 550$-$800 cm$^{-1}$ region of the IRMPD spectra of the small HCO$_3^-$·(H$_2$O)$_n$ clusters is a general feature that will be discussed in section 6. The presence of isomer 1-II can be eliminated on the basis of feature F. In the simulated spectra of 1-II, the water bending (1692 cm$^{-1}$) and asymmetric CO$_2$ stretching (1746 cm$^{-1}$) modes form two distinct features, while only a single peak is observed in the IRMPD spectra. For isomer 1-III, peak D occurs at higher energy than in the experimental spectra and the libration of the singly H-bonded water. Also, in 2-III (as in 1-III), the water at the $\gamma$ position induces a blue-shift of peak D with respect to the experimental frequency.

5.2. HCO$_3^-$·(H$_2$O)$_2$. Figure 4 shows that in the lowest energy isomer of the $n = 2$ cluster, 2-I, the waters are at the $\alpha$ and $\beta$ sites, and each forms two H-bonds with the HCO$_3^-$ anion. The 2-II isomer lies higher in energy by 4.4 kJ/mol above 2-I and has one water molecule at each of the three sites. The 2-III isomer has two waters in the $\alpha$ position and one in the $\gamma$ position. Note that isomer 2-III was found to be the global minimum at the B3LYP level, in accordance with the previous study of Rudolph et al.$^{17}$ Once again, the simulated spectrum of the lowest-energy isomer offers the best agreement with the experimental spectra. The simulated spectra of 3-II and 3-III exhibit a splitting of peak C that is not observed experimentally. This splitting arises from two linear combinations of the C$-$OH stretch and libration of the singly H-bonded water. However, since the intensities of the libration bands are uncertain (see section 6), the presence of isomer 3-III cannot be ruled out. In 3-II, the calculated position of peak D at 1277 cm$^{-1}$ is noticeably higher than the experimental frequency, 1196 cm$^{-1}$. It thus appears that isomer 3-II does not contribute to the experimental spectrum.
Figure 5. Experimental IRMPD spectrum of HCO$_3^-$·(H$_2$O)$_3$ (top panel), with the simulated linear absorption spectra, structures, and relative energies (kJ/mol) of the three lowest-energy isomers.

Figure 6. Experimental IRMPD spectrum of HCO$_3^-$·(H$_2$O)$_4$ (top panel), with the simulated linear absorption spectra, structures, and relative energies (kJ/mol) of the three lowest-energy isomers.

Figure 7. Experimental IRMPD spectrum of HCO$_3^-$·(H$_2$O)$_5$ (top panel), with the simulated linear absorption spectra, structures, and relative energies (kJ/mol) of the three lowest-energy isomers.

5.4. HCO$_3^-$·(H$_2$O)$_4$. Figure 6 shows results for the $n = 4$ cluster. In the lowest-energy isomer, 4-I, all four waters are at the $\alpha$ site. Each forms one H-bond with the HCO$_3^-$ anion and another with an adjacent water, resulting in a four-membered ring. Isomer 4-II, lying 2.3 kJ/mol above 4-I, has a three-membered ring in the $\alpha$ position, similar to 3-I, and the fourth water in the $\beta$ position. Isomer 4-III has two singly H-bonded waters in the $\alpha$ position, while the two remaining waters are at the $\beta$ and $\gamma$ sites. This isomer lies 5.4 kJ/mol above 4-I.

The simulated spectra of 4-I and 4-II are very similar and in good agreement with experiment. The position and intensity of peak F is in slightly better agreement with the simulated spectrum of 4-I but the presence of 4-II cannot be ruled out. For isomer 4-III, the calculated position of peak D is blue-shifted with respect to the experimental value to such an extent that it appears as a shoulder on peak E in the simulation. Isomer 4-III was found to be the minimum energy structure at the B3LYP level which is again not consistent with the experimental observations.

5.5. HCO$_3^-$·(H$_2$O)$_5$. For the $n = 5$ cluster, shown in Figure 7, the three lowest-energy structures each have four waters in the $\alpha$ position forming a single H-bond with the HCO$_3^-$ anion, similar to the ring arrangement in 4-I. In the lowest-energy isomer, labeled 5-I, the fifth water molecule is in the $\beta$ position, while it is at the $\gamma$ site in the 5-II isomer. In the 5-III isomer, the fifth solvent molecule does not bind directly to the HCO$_3^-$ anion but rather to two other water molecules, thus forming a five-membered hydrogen-bonded network in the $\alpha$ position. Isomers 5-II and 5-III are 2.3 and 3.1 kJ/mol above the 5-I structure, respectively.

The simulated spectra of 5-I and 5-III are in reasonable agreement with the experiment, but both overestimate the splitting of peak F. The spectrum of 5-I reproduces the experimental spectrum better around peaks B and C. Isomer 5-II can be ruled out on the basis of the doublet between peaks B and C, that is not observed experimentally, and the predicted position of peak D which is blue-shifted with respect to the experimental value.

5.6. HCO$_3^-$·(H$_2$O)$_6$. Experimental and simulated spectra from the three calculated lowest-lying isomers of HCO$_3^-$·(H$_2$O)$_6$ are shown in Figures 8, 9, and 10. For each of these three clusters, the calculated lowest-energy structure
is based on 5-I; the additional water molecules extend the water–water network at the α site and form a second solvation layer. Most of the other low-lying isomers in this range have one water binding in the γ position. For the \( n = 6 \) and \( n = 7 \) clusters, we can rule out these isomers as the major contributors to the experimental spectra on the basis of the position of band D, just as in the smaller clusters. However, for \( n = 6 \), we observe a weak absorption at \( \sim 1290 \text{ cm}^{-1} \) which might indicate a small amount of 6-II. In the \( n = 8 \) experimental spectrum, band D is not clearly observed, but there is a weak feature at \( \sim 1200 \text{ cm}^{-1} \) which corresponds to the calculated position of band D in isomer 8-I. The calculated shape of band F for 8-I is also in better agreement with the experimental spectrum than the other isomers. Thus, we conclude that 8-I is probably the main isomer present in the experiment.

6. Discussion

For some clusters, comparing the experimental and calculated spectra does not always identify a single isomer as being responsible for the spectrum. It is particularly difficult to distinguish between the isomers with water binding in the α or β position, because they typically have very similar spectra in the energy range studied here. However, the overall good agreement between the calculated spectra and the experiment indicates that our binding motifs are reasonable. Clear trends emerge from the calculated stepwise solvation motifs presented here. The most striking of these is the strong preference of the water molecules for the α position. This trend is consistent with the classical picture of the bicarbonate anion in which the negative charge is shared by the O(1) and O(2) oxygen atoms. However, we find that four waters at most can directly interact with the HCO\(_3\) anion in the α position. The binding motif in isomer 4-I, in which each of the four waters in the α position forms a single H-bond with the CO\(_2\) moiety, is retained in all the most stable isomers of the larger clusters.

Our calculations found a maximum of one water molecule bound to a bicarbonate site other than the α position in the most stable isomers. It is preferentially located at the β site, forming H-bonds with both the O(2) and O(3) oxygen atoms of the bicarbonate anion. The isomers having water molecules in the γ position were systematically found to be higher in energy.
Moreover, these isomers always yielded a frequency for peak D, the $\nu_4$ COH bend, that was blue-shifted relative to the experimental spectra.

For clusters with $n > 5$, five water molecules at most were found to interact directly with the bicarbonate anion. Starting at $n = 6$, the additional solvent molecules formed a second hydration shell, extending the water–water network located at the $\alpha$ site. Finally, in all the isomers presented here, the water molecules were located within or close to the plane of the bicarbonate anion. Isomers with water located on top of the HCO$_3^-$ core were significantly higher in energy. This result differs from the microsolvation motifs found for the similarly planar NO$_3^-$ anion. For NO$_3^-$($H_2O$)$_n$ cluster with $n = 4–6$, the most stable isomers included a four-membered water ring located above the NO$_3^-$ molecular plane.$^{26}$

The monotonic frequency shifts observed in the IRMPD spectra as a function of cluster size are well-reproduced by the calculations. In order to understand these solvation-induced frequency shifts, we have to examine the effect of hydration on the HCO$_3^-$ anion geometry. In the bare anion, the C–O(1), C–O(2), and C–O(3) bond lengths are calculated to be 1.254 Å, 1.238 Å, and 1.452 Å, respectively, at the MP2/6-311+G(d,p) level. These bond lengths are consistent with the C–O(3) having single bond character, while the C–O(1) and C–O(2) have ~1.5 bond character owing to sharing of the negative charge. The slightly longer C–O(1) bond, with respect to C–O(2), indicates that the interaction between the hydrogen and O(1) stabilizes the resonance structure in which the negative charge is on the O(1) atom. Upon the addition of a water in the $\alpha$ position, (isomer 1-I) the C–O(3) bond length is reduced to 1.424 Å. On the other hand, the C–O(1) and C–O(2) bonds are slightly elongated to 1.259 Å and 1.242 Å, respectively. In addition, the O(1)CO(2) angle is slightly reduced, from 135.0° to 131.2°. Thus, the interaction of water at the $\alpha$ position increases the strength of the C–O(3) bond while weakening the C–O(1) and C–O(2) bonds. These trends continue as more water molecules are added at the $\alpha$ position. For example, in isomer 8-I, the C–O(1), C–O(2), and C–O(3) bond lengths are calculated to be 1.258 Å, 1.251 Å, and 1.393 Å, respectively.

In the experimental spectra, bands C and E display the largest frequency shifts upon sequential addition of water molecules. They are blue-shifted by 99 cm$^{-1}$ and 65 cm$^{-1}$, respectively, as the cluster size increases from 1 to 10 water molecules. These two bands are assigned to the $v_3$ and $v_5$ modes, which primarily consist of C–O(3) stretching motion. The large blue-shift observed for these bands is thus consistent with strengthening of the C–O(3) bond induced by the water interactions in the $\alpha$ position. The frequency of the $v_5$ mode, which mainly involves C–O(1) and C–O(2) stretching motion, cannot be directly derived from our spectra because of the overlap with the bending mode of the water in band F. However, from the overall shifts of band F, we estimate that the frequency of this mode is red-shifted by about 50 cm$^{-1}$ as the cluster size increases from 1 to 10 water molecules. This trend is again consistent with the water interactions in the $\alpha$ position which elongate the C–O(1) and C–O(2) bonds. The frequency of the $v_3$ mode (band D), the CO(3)H bending motion, is blue-shifted by only 30 cm$^{-1}$ with increasing cluster size. This is consistent with our calculations which show that no water molecule directly interacts with the H of the HCO$_3^-$ core in the cluster size range studied here. The small blue-shift of band D might be from shortening of the C–O(3) bond and increased charge on the CO$_2$ moiety which would perturb the interaction between the H and O(1) atoms.

The only band which does not show a monotonic frequency shift upon solvation is peak B, assigned to the CO$_3$ out-of-plane mode. It red-shifts from 860 cm$^{-1}$ to 825 cm$^{-1}$ upon addition of the second water and subsequently blue-shifts by only 16 cm$^{-1}$ from $n = 3–10$. The calculation does not reproduce this non-monotonic behavior of peak B in the smaller clusters. It is possible that this discrepancy is due to anharmonic effects because in the solvated clusters, the CO$_3$ out-of-plane mode of the bicarbonate anion also involves water libration motions. It is also possible that peak B in the HCO$_3^-$($H_2O$)$_n$ experimental spectrum, where it appears as a small shoulder, does not correspond to the CO$_3$ out-of-plane mode of isomer 1-I, but rather the C–OH stretching mode (feature C) of another isomer such as 1-II.

Some of the calculated absorption bands are not observed in the experimental spectra, most notably the absorption bands in the 600–800 cm$^{-1}$ spectral region for clusters with $n < 6$ (labeled as peak A in the larger clusters). These bands are from various water librational modes. Their absence in the experimental spectra is similar to what was observed for the NO$_3^-$($H_2O$)$_n$ clusters$^{26}$ and can originate from several effects. For example, the number of photons required to reach the dissociation continuum is 3 times larger at 600 cm$^{-1}$ than at 1800 cm$^{-1}$. This renders the low-frequency modes considerably more difficult to observe in our IRMPD spectra than those at higher frequencies. However, peaks B and C appear in our spectra with intensities close to their calculated linear absorptions, so frequency alone cannot explain the absence of peak A in small cluster sizes.

Structural changes that accompany IR multiphoton absorption offer a more likely origin for the absence of low-frequency librational bands. For example, in the 1-I isomer, the absorption of a few photons at the water librational mode frequency is not sufficient to break both H-bonds, but can break a single H-bond. However, the resulting singly bonded water cannot absorb any additional photons, because the absorption frequency of its librational mode has changed and is now out of resonance with the irradiation wavelength. This interpretation is supported by our experimental spectra in which the water libration band (band A) appears only for clusters with six or more water molecules, since HCO$_3^-$($H_2O$)$_6$ is the smallest cluster in which the lowest-lying isomer (isomer 6-I) contains a singly bonded water molecule. These clusters are thus more likely to dissociate rather than isomerize upon absorption of infrared photons. Moreover, the intensity of band A doubles in HCO$_3^-$($H_2O$)$_6$ for which the lowest-energy isomer (7-I) contains two singly bonded water molecules.

In the cluster size range studied here, the frequencies of all the observed absorption bands monotonically shift toward their aqueous values as the number of water molecule is increased. With the exception of band D, all the absorption bands are within 30 cm$^{-1}$ of their position in aqueous solution by the $n = 10$ cluster. In the smaller clusters, band D is found to be more red-shifted than the other bands with respect to the aqueous frequencies. In the $n = 7$ cluster, the largest cluster in which this band is clearly observed, band D is still 115 cm$^{-1}$ below its aqueous frequency of 1320 cm$^{-1}$. This difference can be easily understood from the calculated vibrational spectra of the various isomers. Our calculations found that cluster isomers with a water at the $\gamma$ site were systematically higher in energy.
However, it can be seen from the calculated spectra of those isomers that a water molecule at the $\gamma$ position induces a $\sim 100$ cm$^{-1}$ blue-shift of band D. In aqueous solution, at least one water molecule is certainly found in this position. It would be of interest to determine how large a cluster has to be before the lowest-energy structure has a water at the $\gamma$ site.

Finally, it is worthwhile to comment on the insights that the present study may yield on the structure of the fully hydrated bicarbonate anion. Careful examination of the polarization of the Raman modes in very dilute solutions of KHCO$_3$ and KDCO$_3$ led Rudolph and co-workers to conclude that the hydrated HCO$_3^-$ anion has a $C_1$ symmetry in contrast to the $C_s$ symmetry expected in the gas-phase.$^{17,18}$ In the clusters considered here, the calculated lowest-lying isomers all have a HCO$_3^-$ anion with $C_1$ (planar) symmetry. However, the isomers having a water in the $\gamma$ position (such as 8-III) have a HCO$_3^-$ core with a $C_1$ symmetry, with a O(1)CO(3)H dihedral angle of about 2°. Thus, the HCO$_3^-$ core may well adopt a geometry with $C_1$ symmetry in larger clusters and bulk water where the ion is fully solvated.

7. Conclusion

Infrared spectra are reported for the HCO$_3^-$($H_2O$)$_n$, $n = 1$–10 clusters in the 600–1800 cm$^{-1}$ spectral range. Electronic structure calculations using DFT and MP2 have been performed on the $n = 1$–8 species to determine the structure of the lowest-lying isomers. Comparison of the experimental and calculated spectra does not always enable unambiguous identification of a single isomer for each cluster. Nonetheless, the overall good agreement between the calculated spectra and experiment allows us to determine general trends in the stepwise solvation motifs of the bicarbonate anion. The most important of these is the strong preference of water molecules to bind to the negatively charged CO$_2$ moiety (the $\alpha$ site) of the HCO$_3^-$ anion. However, a maximum of four waters can directly interact with the anion at this position. The binding motif in the most stable isomer of $n = 4$, which consists of a four-membered ring with each water forming a single H-bond with the CO$_2$ moiety, is retained in all the most stable isomers of the larger clusters. Starting at $n = 6$, the additional solvent molecules are found to form a second hydration shell, extending the water–water network located at the $\alpha$ site of the bicarbonate anion. We also find that binding at the hydroxyl group of HCO$_3^-$ is disfavored for all clusters studied here; this result accounts for the largest discrepancy between the cluster spectra and the IR spectrum of aqueous bicarbonate.

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