ZEKE-PFI spectroscopy of 1:1 complexes of sodium with water and ammonia

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Abstract

ZEKE-PFI (zero kinetic energy pulsed field ionization) photoelectron spectra of the Na(H\textsubscript{2}O), Na(D\textsubscript{2}O), Na(NH\textsubscript{3}), and Na(ND\textsubscript{3}) complexes are reported. Spectra of all four complexes were obtained by single-photon ionization, and, for the Na(NH\textsubscript{3}) and Na(ND\textsubscript{3}) complexes, by two-color (\textit{i} + \textit{i}') photoionization as well, with the \textit{A} \textsuperscript{2} \textit{E} state serving as the intermediate resonance. Improved values for the ionization energies (IE) and intermolecular vibrational frequencies of the complexes were determined. The single-photon ZEKE-PFI spectra show transitions only between states of the same vibrational symmetry, in accord with the selection rule for allowed electronic transitions. Some of the two-color ZEKE-PFI spectra, however, show strong transitions between states of different vibrational symmetry which we attribute to vibronic coupling in the intermediate state.

1. Introduction

Alkali metal ions solvated by water and other polar molecules have received considerable attention [1-5] due to their importance in many chemical and biological systems, the central nervous system being a prime example. An accurate description of the manner in which the alkali ion-solvent and solvent-solvent interactions change within the solvation shells as a function of ionic radius is required to explain phenomena such as ion selectivity in membrane channels [6], but adequately detailed information is difficult to obtain from condensed phase measurements. Consequently, many researchers have studied gas phase clusters comprised of a metal ion attached to a few solvent molecules, since they form more accessible models of solution behavior. For clusters of the type Na\textsuperscript{+}(H\textsubscript{2}O)\textsubscript{n} and Na\textsuperscript{+}(NH\textsubscript{3})\textsubscript{n}, the binding energies of the solvent molecules to the clusters have been measured as a function of \textit{n} [7,8], and, for \textit{n} \leq 6, these data agree well with the results of numerous theoretical studies [9-11]. However, in the case of Na\textsuperscript{+}(NH\textsubscript{3})\textsubscript{n} clusters there is less agreement on the number of solvent molecules required to fill the first solvation shell, with estimates ranging from 4 to 6 having been obtained by various methods [8,11,12].

While quantitative intermolecular potential energy surfaces (IPESs) can be obtained from high-resolution spectra of the cluster intermolecular vibrational modes, as has been demonstrated in the case of hydrogen-bonded dimers [13], the absence of such data has thus far prevented the determination of the IPESs of Na\textsuperscript{+}(H\textsubscript{2}O)\textsubscript{n} and Na\textsuperscript{+}(NH\textsubscript{3})\textsubscript{n} clusters. Indeed, the only measurements of intermolecular vibrational modes of such clusters were made by Hertel
and co-workers, who determined the sodium ion-molecule stretching vibrational frequencies for the 1:1 complexes of Na+ with water and ammonia from photoionization efficiency (PIE) spectra of the neutral dimers [14,15].

Neutral clusters composed of an alkali metal atom and solvent molecules are also of interest in their own right because of their utility in the study of electron solvation [16-20]. In this context, it is interesting to note that the ionization energies (IE) of clusters with \( n \leq 20 \) determined from PIE spectra reveal that while the Na(NH\(_3\))\(_n\) IEs decreased monotonically with \( n \), those for Na(H\(_2\)O)\(_n\) decreased until \( n = 4 \), whereupon they reached the bulk value and remained essentially constant [21]. Misaizu et al. found the same trends in the IEs of clusters of cesium atoms with water or ammonia [22].

Theoretical studies have offered two possible explanations for the different IE behavior of sodium-water and sodium-ammonia clusters. Using local spin density functional theory, Barnett and Landman [23] found that the Na atoms in Na(H\(_2\)O)\(_n\) clusters with \( n \geq 4 \) are surrounded by H\(_2\)O molecules and that the Na 3s electron is partially removed from the atom and delocalized in a 'surface Rydberg-like state.' They claim that delocalization would not occur to the same extent in Na(NH\(_3\))\(_n\) clusters with \( n \leq 30 \), and that the cluster IEs will change until this size is reached. In an ab initio MO study, Hashimoto and Morokuma [10,11] found that the Na atom is surrounded by NH\(_3\) molecules and that the 3s electron is partially delocalized in large clusters, but in large Na(H\(_2\)O)\(_n\) clusters they predict the Na atom is attached to the surface of a water cluster with the 3s electron located outside the water cluster and the IE largely unaffected by the addition of water molecules. Unfortunately, no direct structural evidence for either the neutral or ionized clusters has been obtained, and much more vibration-rotation data is needed to characterize their IPESs.

As our initial step toward a general description of alkali metal-solvent interactions, we have conducted a study of 1:1 complexes of sodium with water or ammonia in the gas phase using ZEKE-PFI (zero kinetic energy-pulsed field ionization) photoelectron spectroscopy. It is well documented that the IEs of neutral clusters and the rovibrational energy levels of the ionized clusters can be measured more accurately with ZEKE-PFI spectroscopy than with energy-resolved photoelectron or PIE spectroscopies [24-27]. From our ZEKE-PFI spectra we have obtained improved IEs for the Na(H\(_2\)O) and Na(NH\(_3\)) dimers and improved sodium ion-molecule stretching vibrational frequencies for the ionized complexes. We also report here the first measurement of the intermolecular bending vibrational frequencies for the Na\(^+\)(NH\(_3\)) and Na\(^+\)(ND\(_3\)) complexes.

2. Experimental

Neutral complexes were formed and interrogated in a three-stage, differentially pumped vacuum chamber which included a 'pick-up' cluster source, a ZEKE-PFI spectrometer, and a reflectron time-of-flight (TOF) mass spectrometer. The 'pick-up' [14,15] source consisted of a pulsed supersonic jet which contained the water or ammonia molecules and a sodium beam that intersected the supersonic jet at right angles. The sodium atoms, after undergoing several collisions, were entrained in the jet, and a small fraction of them formed clusters. The sodium beam was produced by heating a sodium-filled stainless steel oven with a 3 mm diameter aperture to 630 K. The oven's aperture was placed approximately 5 mm below and 5 mm downstream of the orifice of the pulsed valve (General Valve Series 9, 0.8 mm nozzle diameter). For complexes with water, a mixture of 1 atm of water vapor and 1.5 atm of argon was expanded through the valve, while for ammonia complexes, ammonia at a pressure of 3 atm was expanded next. After passing through a 2 mm diameter skimmer placed 50 mm away from the pulsed valve orifice, the molecular beam travelled 48 cm before reaching the ZEKE-PFI spectrometer's ionization region. The ZEKE-PFI spectrometer, which was fabricated from the modified ionization source of a commercial TOF mass spectrometer (R.M. Jordan Co.), consisted of two mesh grids used for extracting the electrons, a third grid that provided additional acceleration, a 2-μm flight tube, and a microchannel plate detector. Ionization of the high-n Rydberg states was accomplished by a pulsed electric field applied between the extraction grids 2.5 μs after the excitation laser intersected the cluster beam, with a typical magnitude of 4 V/cm and a risetime of 80
ns. The post-extraction acceleration grid and the μ-metal tube were held at a potential of 9 V. In the single-photon experiments, the complexes were probed with the frequency-doubled output of a dye laser (Lambda Physik FL 3002) pumped by a Nd:YAG laser (Continuum YG681). In the two-color experiments, the fundamental outputs of two FL3002 dye lasers which had bandwidths of \( \approx 0.2 \text{ cm}^{-1} \) were used. The wavelengths of the dye lasers were calibrated with a wavemeter (Burleigh WA-4500A) and corrected to their vacuum values. The ZEKE-PFI signal was amplified and then recorded on a digital oscilloscope (LeCroy 9400A). The ZEKE-PFI peak heights were not corrected for the variations of laser power with wavelength.

3. Results and discussion

Single-photon ZEKE-PFI spectra of Na(H\(_2\)O) and Na(D\(_2\)O) are presented in Fig. 1, while those for Na(NH\(_3\)) and Na(ND\(_3\)) are shown in Fig. 2. We associate the first and most intense peak in each spectrum with the \( \Delta v = 0 \) transition from the neutral complex's vibronic ground state to the vibronic ground state of the cation. This transition is denoted by \( 0^0_+ \). Here the notation \( \nu' \nu'' \) is such that the leading number refers to the vibrational mode involved in the transition, with the subscript and superscript denoting the vibrational quantum number in the lower and upper state, respectively. Quantum numbers of cations are followed by a plus symbol. The vibronic ground states of the neutral and ionized Na(NH\(_3\)) and Na(ND\(_3\)) complexes have \( \text{A}_1 \) symmetry in the \( \text{C}_3v \) point group [28], while the vibronic ground states of the neutral and ionized Na(H\(_2\)O) and Na(D\(_2\)O) complexes have \( \text{A}_1 \) symmetry in the \( \text{C}_2v \) point group.

Adiabatic ionization energies of \( 35323 \pm 10 \text{ cm}^{-1} \) for Na(H\(_2\)O), \( 35249 \pm 10 \text{ cm}^{-1} \) for Na(D\(_2\)O), \( 34435 \pm 10 \text{ cm}^{-1} \) for Na(NH\(_3\)), and \( 34368 \pm 10 \text{ cm}^{-1} \) for Na(ND\(_3\)) are derived from the spectra. While ZEKE-PFI spectroscopy is capable of giving more accurate IEs, the uncertainties of these values are large because of as yet unresolved rotational structure in the peaks and because no correction has been made for the lowering of the IE by the extraction field. The field-induced shift of the IE is denoted by \( \Delta E \) and is given in cm\(^{-1}\) by \( \Delta E = aF^{1/2} \) where \( F \) is the electric field in V/cm and \( a \) is a constant that generally falls between 2 and 6 [27]. However, we have not accurately determined \( a \) for the complexes in this study. The adiabatic IEs for Na(H\(_2\)O) and Na(NH\(_3\)) are, nevertheless, in excellent agreement with values obtained from PIE spectra by Schulz et al. [14] and Nitsch et al. [15] and with calculated values [10,11,23], while the adiabatic IEs of the deuterated complexes are shifted \( \approx 70 \text{ cm}^{-1} \) to lower frequency because of the changes in their zero point vibrational energies.

We assign the other peaks in these single-photon ZEKE-PFI spectra to transitions involving the sodium ion–molecule stretching and bending vibrational modes. The Na(NH\(_3\)) and Na(ND\(_3\)) complexes each
have three vibrational modes of $a_1$ symmetry and three vibrational modes of $e$ symmetry [29]. The sodium–ammonia stretching mode is the lowest frequency mode of $a_1$ symmetry and is denoted by $\nu_3$, while the doubly degenerate sodium–ammonia bending mode is the lowest frequency mode of $e$ symmetry and is denoted by $\nu_6$. The Na(H$_2$O) and Na(D$_2$O) complexes also have six vibrational modes, three of which have $a_1$ symmetry. The sodium–water stretching mode is the lowest frequency mode of $a_1$ symmetry and is again denoted by $\nu_3$. There are two non-degenerate sodium–water bending modes, one with $b_1$ symmetry and the other of $b_2$ symmetry, which are the lowest frequency non-totally symmetric modes in the complexes [9]. They are denoted by $\nu_{3/6}$, but we will hereafter refer to the bending modes only as $\nu_6$ since, on account of spectral resolution or transition strength, only single peaks associated with the bending vibration were observed in our spectra, and no distinction between $\nu_3$ and $\nu_6$ could be made.

Following the $0^+_0$ peak, each spectrum shows a short progression in $\nu_3$. The frequencies of $\nu_3$ are found to be $300 \pm 5$ cm$^{-1}$ for Na$^+(H_2O)$, $296 \pm 5$ cm$^{-1}$ for Na$^+(D_2O)$, $305 \pm 5$ cm$^{-1}$ for Na$^+(NH_3)$, and $286 \pm 5$ cm$^{-1}$ for Na$^+(ND_3)$. The Na$^+(H_2O)$ and Na$^+(NH_3)$ stretching frequencies are in excellent agreement with the values found by Schulz et al. [14] and Nitsch et al. [15], and the Na$^+(H_2O)$ frequency is also very close to that calculated previously [9,23]. A summary of all the vibrational frequencies measured in this study may be found in Table 1. Unresolved rotational structure in the ZEKE-PFI peaks and the poorer signal-to-noise ratios of the higher peaks in the stretching mode progressions prevented us from measuring accurate anharmonicities. The lengths of the progressions in $\nu_3$ indicate that the complexes experience significant but not extremely large contractions of the sodium ion–molecule bond lengths upon ionization, also in agreement with theory. The calculated Na–O bond lengths for the Na(H$_2$O) and Na$^+(H_2O)$ complexes are 2.355 and 2.23 Å, respectively [10], while the Na(NH$_3$) and Na$^+(NH_3)$ bond lengths are predicted to be 2.381 and 2.365 Å [11].

The strongest peaks in the spectra in Figs. 1 and 2 are most likely produced by the ionization of neutral complexes in their vibronic ground states. There are, however, weaker peaks that may plausibly be attributed to the ionization of vibrationally excited neutral complexes. In each spectrum, the $0^+_0$ peak and the peaks in the progression in $\nu_3$ are each followed by two very weak peaks at higher energy. We assign these peaks to two sets of hot bands, one

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### Table 1

<table>
<thead>
<tr>
<th>IE (cm$^{-1}$)</th>
<th>Na(H$_2$O)</th>
<th>Na(D$_2$O)</th>
<th>Na(NH$_3$)</th>
<th>Na(ND$_3$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\nu_3$ (cm$^{-1}$), ion</td>
<td>$35323$</td>
<td>$35249$</td>
<td>$34435$</td>
<td>$34368$</td>
</tr>
<tr>
<td>$\nu_3$ (cm$^{-1}$), neutral</td>
<td>$207$</td>
<td>$206$</td>
<td>$215$</td>
<td>$196$</td>
</tr>
<tr>
<td>$\nu_3$ (cm$^{-1}$), neutral</td>
<td>--</td>
<td>--</td>
<td>$468$</td>
<td>$340$</td>
</tr>
<tr>
<td>$\nu_6$ (cm$^{-1}$), neutral</td>
<td>--</td>
<td>--</td>
<td>$278$</td>
<td>$200$</td>
</tr>
</tbody>
</table>

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* $\nu_6$ (ion) $\sim$ $\nu_6$(neutral) $= 240$ cm$^{-1}$.
* $\nu_6$ (ion) $\sim$ $\nu_6$(neutral) $= 190$ cm$^{-1}$. 
set produced by the ionization of neutral complexes from the $\nu_3$ state, the other produced by complexes with $\nu_6$ initially populated.

The small peaks that appear $\approx 100$ cm$^{-1}$ above the $0_0^{0+}$ peaks of all four complexes probably arise from $\Delta \nu = 0$ transitions that originate from the $\nu = 1$ level of $\nu_3$. If this assignment is correct, the energy difference between the $3_1^{1+}$ hot band and the $0_0^{0+}$ peak is determined by the difference between the frequency of $\nu_3$ in the neutral complex and the frequency of $\nu_3$ in the ionized complex. The frequencies of $\nu_3$ in the ionized complexes, listed in Table 1, are $\approx 300$ cm$^{-1}$, while in the neutral Na(H$_2$O) and Na(NH$_3$) complexes $\nu_3$ has a frequency of $\approx 185$ cm$^{-1}$ [29]. The $3_1^{1+}$ transition should therefore appear $\approx 100$ cm$^{-1}$ higher in energy than the $0_0^{0+}$ transition, as observed. The shift is roughly the same for the deuterated complexes because isotopic substitution does not cause a large change in the pseudo-diatomic stretching reduced mass. A progression in $\nu_3$ is built upon the $3_1^{1+}$ false origin.

The Na(H$_2$O) spectrum exhibits a weak peak 240 cm$^{-1}$ above the $0_0^{0+}$ peak, while the Na(NH$_3$) spectrum shows a weak peak 190 cm$^{-1}$ above its $0_0^{0+}$ origin. We attribute these peaks to $\Delta \nu = 0$ transitions, but now originating from the $\nu = 1$ level of $\nu_6$. Such an assignment is supported by the Na(D$_2$O) and Na(ND$_3$) data, in which these $6_1^{0+}$ hot bands are located only $\approx 190$ and $\approx 140$ cm$^{-1}$ above the respective $0_0^{0+}$ peaks. This isotopic reduction of the energy difference between the $6_1^{0+}$ and $0_0^{0+}$ transitions, by almost a factor of $1/\sqrt{2}$, is expected for a vibrational mode involving primarily the motion of the hydrogen atoms. In the Na(NH$_3$) and Na(ND$_3$) spectra, only a single peak associated with $\nu_6$ is expected, since $\nu_6$ is doubly degenerate. On the other hand, the Na(H$_2$O) and Na(D$_2$O) spectra might be expected to show one peak for each of the two non-degenerate intermolecular bending vibrational modes. The spectra in Fig. 1 hint at the existence of a second very weak peak that could be assigned as a false origin associated with another bending mode, but the signal-to-noise ratios were too poor to clarify this matter. Progressions in $\nu_3$ are then built upon the $6_1^{0+}$ false origins.

In order to measure the frequency of $\nu_6$ for the Na$^+(\text{NH}_3)$ and Na$^+(\text{ND}_3)$ complexes, which could not be obtained from the single-photon ZEKE-PFI spectra alone, we performed ZEKE-PFI experiments using resonant two-color $(1 + 1')$ photoionization via the $\tilde{A}^2E$ state. The $\tilde{A}^2E$ excited electronic state of Na(NH$_3$) has been studied theoretically by Greer et al. [28] and experimentally by Nitsch et al. [29]. A spectrum containing several vibrational bands of the Na(ND$_3$) $\tilde{A}^2E \leftrightarrow \tilde{X}^2A_1$ transition, obtained by resonant two-color $(1 + 1')$ photoionization spectroscopy, is shown in Fig. 3a. The peaks are assigned to the following modes: $0_0^{0+}$ at 12250 cm$^{-1}$, $6_1^{0+}$ at 12470 cm$^{-1}$, $3_1^{0+}$ at 12550 cm$^{-1}$, and $6_1^{0+}3_1^{0+}$ at 12750 cm$^{-1}$. The $6_1^{0+}$ and $3_1^{0+}$ peaks are well resolved in this spectrum, but were not resolved in Nitsch et al.'s spectrum of the $\tilde{A}^2E \leftrightarrow \tilde{X}^2A_1$ transition of Na(NH$_3$) because the frequencies of $\nu_6$ and $\nu_3$ in this state
and isotomer are almost the same. This near degeneracy led Nitsch et al. to assign peaks at higher energy to a progression in $\nu_6$, but the $6_b^1\cdot 3_b^0$ transition in our spectrum of Na(ND$_3$) shows that there is a progression in $\nu_3$, built on the $6^1$ level, not a progression based solely on the $\nu_6$ vibrational mode.

The $6_b^0$ transition would be a forbidden transition if the Born–Oppenheimer and Condon approximations were rigorously satisfied. In such a case, the transition moment integral $M$ is given by

$$M = \langle \psi_e^- | er | \psi_e^+ \rangle \langle \psi_v^- | \psi_v^+ \rangle,$$

where $\psi_e^+$ and $\psi_v^+$ are the electronic wavefunctions for the initial and final states, respectively, $\psi_e^-$ and $\psi_v^-$ are the corresponding vibrational wavefunctions, and $er$ is the electric dipole moment operator. The integral containing the product of the vibrational wave functions is zero by symmetry, because $e \otimes a_1$ does not contain the $a_1$ symmetry element. However, the $6_b^0$ transition may become allowed by vibronic coupling, in which case the transition moment integral cannot be separated into electronic and vibrational components, and is given instead by

$$M = \langle \psi_e^- | er | \psi_e^- \rangle,$$

where $\psi_e^-$ and $\psi_e^+$ are the vibronic wavefunctions for the initial and final states, respectively.

If the vibronic coupling is strong and the $\tilde{A}$ state of Na(NH$_3$) does indeed possess $E$ symmetry, the $6_b^0$ transition can become allowed through the Jahn–Teller effect. Excitation of an $e$ vibrational mode in an $E$ electronic state produces vibronic states having $E$, $A_1$, and $A_2$ symmetries. The transition from the $A_1$ ground state to an $E$ vibronic state is a perpendicular transition, while the transition to the $A_1$ vibronic state is a parallel transition and the $A_2 \leftrightarrow A_1$ transition is forbidden. Only the $E \leftrightarrow A_1$ transition will have significant intensity, however, because it can borrow intensity from the transitions to the $\tilde{A}$ state that are allowed in the absence of vibronic coupling, which are perpendicular transitions [30].

Greer et al. predict that the second excited electronic state of Na(NH$_3$) has $A_1$ symmetry and lies only 2400 cm$^{-1}$ above the $\tilde{A}$ state. The experimental work of Nitsch et al. suggests that there is indeed an excited electronic state only 800 cm$^{-1}$ above the $\tilde{A}$ state. Since the symmetries of the lowest excited electronic states of this complex have not been deter-

mined experimentally, the possibility exists that the $\tilde{A}$ state has $A_1$ symmetry and that the next highest level is of $E$ symmetry. If the $\tilde{A}$ state does have $A_1$ symmetry, the $6_b^0$ transition can become allowed not by the Jahn–Teller effect but by Herzberg–Teller coupling with the nearby electronic state of $E$ symmetry [31]. In this case, the $0^0$ band would be a parallel band, and the $6_b^0$ band would be a perpendicular band because it borrows intensity from a perpendicular transition to an electronic state of $E$ symmetry.

Which vibronic coupling mechanism is operative in this case must be determined from the rotational structure of the bands. Band profiles of the $0^0$ and $6_b^1$ transitions of Na(ND$_3$) are shown in Fig. 3b and c. The band contours are strikingly different. The $\tilde{A} \leftarrow X 0^0$ band rises very sharply to a peak and then tails off smoothly. Such a band contour is characteristic of a parallel transition of a prolate symmetric top, with $P$ branch head formation resulting from shrinkage of the Na–N bond. The $6_b^1$ band, however, shows several sharp peaks spaced by 3.4 cm$^{-1}$, which is roughly one-half of the peak spacing observed in the Na(NH$_3$) spectrum (not shown). This spacing is very close to the rotational constant of ammonia about its $C_3$ axis. It is also roughly an order of magnitude greater than the $B$ rotational constants of these complexes derived from the calculated structures. Such a pattern is characteristic of a perpendicular transition of a symmetric top molecule for which $A \gg B$.

If the $\tilde{A} \leftarrow X 0^0$ transition is parallel and the $6_b^0$ transition is perpendicular, the Herzberg–Teller vibronic coupling mechanism would then be responsible for the $6_b^1$ transition. However, such a conclusion should not be based on unresolved rotational structure, since the structure of perpendicular bands can sometimes look like that of parallel bands. This happens because the peak spacing in a perpendicular band depends on the total vibronic angular momentum, which has an electronic component due to the degenerate electronic state and a vibrational component when the degenerate vibration is excited in that state [30]. Since the $0^0$ level has no vibrational angular momentum, the peak spacing in a perpendicular $0^0$ band will be different than in the $6_b^0$ band. A famous example of this type of behavior is the $C \leftarrow X$ transition of CH$_3$I, which shows such
pseudo-parallel bands [32]. Complete rotational resolution, then, will be required to settle the issue of the electronic state ordering and the vibronic coupling mechanism operative in the Na(NH₃) complex.

Two-color excitation through the vibronic levels of the A state has been used to further characterize the IPESs of the ionized complexes, particularly through the isolation of ν₆. Due to the Δν = 0 propensity rule which generally holds in the photoionization of molecules having similar geometries in their neutral and ionized forms [33], the vibrational mode that is excited in the intermediate electronic state will usually show the strongest peak in the ZEKE-PFI spectrum. A two-color ZEKE-PFI spectrum of Na(ND₃) taken via the 0° level of the A state is shown in Fig. 4a. The strongest peak results from ionization to the cation ground state (the 0°⁺ transition). This Δν = 0 behavior is expected, since the calculated geometry of the A state of the Na(NH₃) complex is very close to that of the Na⁺(NH₃) complex. Two weaker peaks appear 290 and 340 cm⁻¹ higher in energy, and are assigned to the 3⁺₁ and 6⁺₁ transitions, respectively. It should be noted that the 3⁺₁ transition is much weaker than in the single-photon ZEKE-PFI spectrum, and this probably indicates that the Na–N bond is shorter in the A state than in the ground state of the neutral complex, as Greer et al. have suggested.

Two-color ZEKE-PFI spectra taken via the 6¹ level in the A state of the Na(ND₃) and Na(NH₃) complexes are shown in Fig. 4b and c, respectively. The assignment of the Na(ND₃) 6⁺₁ transition in Fig. 4a is supported by the spectrum in Fig. 4b, which shows a strong peak for the 6⁺₁ transition, a strong feature 340 ± 10 cm⁻¹ above the 6⁺₁ peak corresponding to the 6⁺₁ transition, and a weak peak 680 cm⁻¹ above the 6⁺₁ peak that we assign to the 6⁺₁ transition. The spectrum of Na(NH₃) in Fig. 4c was obtained by tuning the excitation laser to the low-frequency side of the spectral feature formed by the superposition of the 6⁺₀ and 3⁺₀ bands in the Na(NH₃) A ← X transition. It shows a strong 6⁺₁ peak and a strong peak 468 ± 10 cm⁻¹ higher in energy, which we assign as the 6⁺₁ transition. The intensity of the 6⁺₁ peak is greater than that of the 6⁺₁ peak, seemingly in violation of the Δν = 0 propensity rule. This deviation from expected behavior is partly due to a loss of intensity in the ionization laser beam as the blue edge of the laser dye's tuning curve was approached. However, it is significant that the 6⁺₁ transition appears with intensity comparable to that of the 6⁺₁ transition.

The ratio of the frequencies of the ν₆ vibrational modes of the Na⁺(ND₃) and Na⁺(NH₃) complexes is 0.726, which is approximately what one would expect for this bending mode. The Na⁺(NH₃) bending frequency is also reasonably close to the bending frequencies calculated for the Na⁺(H₂O) complex [9]. The increased 'stiffness' in the bending coordinate relative to the stretching coordinate results from the directionality of the ion–dipole force which is the main contributor to the bonding in this complex [29].

Combining these results with the changes in the ν₆ modes derived from the hot bands observed in the single-photon ZEKE-PFI spectra, frequencies for ν₆
ZEKE-PFI spectra of Na(NH$_3$)$_3$ and Na(ND$_3$)$_3$ taken via the 6$^1$3$^1$ level of the Σ state are shown in Fig. 5b and c. The Na(NH$_3$)$_3$ spectrum shows a 6$^1_3$+ 3$^0_+$ peak, a 6$^0_3$+ 3$^1_+$ peak, a 6$^0_0$+ 3$^2_+$ peak, and a peak 770 cm$^{-1}$ above the 6$^1_3$+ 3$^1_+$ peak that we assign to the 6$^1_1$+ 3$^1_+$ transition. The 6$^1_1$+ 3$^0_+$ transition appears with very little intensity. The Na(ND$_3$)$_3$ spectrum shows a peak for the 6$^0_3$+ 3$^1_+$ transition, a peak for the 6$^0_0$+ 3$^1_+$ transition 280 cm$^{-1}$ above it, and a third peak 624 cm$^{-1}$ above 6$^0_0$+ 3$^0_+$ peak that corresponds to the 6$^1_1$+ 3$^1_+$ transition. These spectra confirm our assignments of the intermediate states.

The most startling aspect of the ZEKE-PFI spectra taken via the 6$^1$3$^1$ level is the violation of the Δν = 0 propensity rule by strong transitions to ion vibrational states of different symmetry. The spectra show the expected Δν = 0 transition to the 6$^1$3$^1$+ level which has Σ symmetry, but also strong transitions to the origin and the 3$^1_1$+ level, which have Α symmetry. Interestingly, the peak for the transition to the 6$^1_1$+ level is weak or non-existent. Similar behavior is observed in the spectra taken via the 6$^1$ level, shown in Fig. 4b and c, which show the Δν = 0 peaks for the transition to the 6$^1$3$^1$+ level as well as strong transitions to the ion ground states. In contrast, the ZEKE-PFI spectra shown in Figs. 4 and 5a, taken via states of Α$_1$ vibrational symmetry that possess no excitation of ν$_6$, show only very weak transitions to states of Σ vibrational symmetry. Thus, the selection rule for allowed electronic transitions which forbids interactions between states of different vibrational symmetry (see Eq. (1)) is most closely followed in the single-photon ZEKE-PFI spectra, which show only transitions between states having the same vibrational symmetry.

We attribute the transitions between states of different vibrational symmetry to strong vibronic coupling in the Σ state, which our spectra of the Σ state have shown to exist. We note that vibronic coupling in the intermediate electronic state has been found to be responsible for transitions between states of different vibrational symmetry in two-color ZEKE-PFI spectra of other molecules [34,35]. It is possible that the transitions could result from vibronic coupling in the cation, but this would require either the existence of low-lying excited electronic states in the cation or degeneracy of the ground electronic state of the cation. To our knowledge,

in the neutral Na(NH$_3$)$_3$ and Na(ND$_3$)$_3$ complexes are found to be 278 ± 20 and 200 ± 20 cm$^{-1}$, respectively. It is somewhat surprising that the bending frequency for the neutral Na(NH$_3$)$_3$ complex is higher than that for the stretching mode, since the calculations that have been done on similar species to date have found that the stretching frequencies are higher [19]. The measurement of the vibrational frequencies of other alkali metal–molecule complexes should shed considerable light on the nature of the bond interactions in these neutral complexes.

The assigned frequency of ν$_6$ for the Na$_3^+$-(NH$_3$)$_3$ complex is supported by the spectrum presented in Fig. 5a, taken via the 3$^1_1$ level of the Σ state. The strongest peak in the spectrum is assigned to the 3$^1_1$+ transition, and weak peaks that we assign to the 3$^0_0$+, 6$^0_0$+ 3$^0_+$, and 3$^0_1$+ transitions are evident. The 6$^0_0$+ 3$^0_+$ peak is 470 cm$^{-1}$ above the 3$^0_1$+ peak.

Fig. 5. Two-color ZEKE-PFI spectra of (a) Na(NH$_3$)$_3$ via the 3$^1_1$ level of its Σ state (b) Na(NH$_3$)$_3$ via the 6$^1$3$^1$ level of its Σ state (c) Na(ND$_3$)$_3$ via the 6$^1$3$^1$ level of its Σ state.
neither condition is satisfied in the Na+\((\text{NH}_3)\) complex. Further understanding of vibronic coupling in this complex, as well as the structural parameters of complexes of sodium ions with ammonia, water and other molecules, can be obtained from rotationally resolved two-color ZEKE-PFI spectra which are currently being taken in our laboratory.

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