The infrared spectrum of the benzene–Ar cation

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The infrared (IR) absorption spectra of the jet-cooled C6H6 and C6D6 cations, complexed with Ar, are measured throughout the 450–1500 cm−1 region via IR-laser-induced vibrational dissociation spectroscopy. The IR spectrum of the C4H6–Ar cation is dominated by a Fermi resonance between the IR active ν11 mode and two components of the combination mode of the lowest frequency modes ν6 and ν16. A stringent upper limit of 316 cm−1 is found for the value of the dissociation limit D0 of the neutral C4D6–Ar complex. © 1999 American Institute of Physics.

The benzene cation with its doubly degenerate X2E∗ ground state is arguably the most prototypical example of a Jahn–Teller distorted system. According to Jahn and Teller,1 the coupling of nuclear motion with the electronic degeneracy leads to a distortion of the molecule along certain normal coordinates so that the potential minimum is no longer at the symmetrical position. Consequently, there has been considerable interest to unravel the vibrational properties and to deduce information on this Jahn–Teller distorted structure for the benzene cation.2 For many of the halogenated benzenoid cations gas-phase laser-induced fluorescence (LIF) studies have been applied and have yielded a wealth of information and improved understanding on the Jahn–Teller effect as such, as in these studies the Jahn–Teller active modes are directly accessible.3,4 Its low fluorescence quantum yield has prohibited such gas-phase LIF studies on the benzene cation. Photoelectron spectroscopy5 and threshold-electron (zero electron kinetic energy, ZEKE)6–8 and threshold-ion detection (mass analyzed threshold ionization, MATI) schemes9–11 have been successfully used, however, to measure and assign several vibrational modes in the benzene cations.

In this communication we report the first gas-phase IR absorption spectra of the C6H6 and C6D6 cations, complexed with Ar, throughout the 450–1500 cm−1 region. The spectra are measured via IR-laser-induced vibrational dissociation of the laser prepared cationic van der Waals complexes. The experimental detection scheme also allows one to determine an upper limit for the dissociation energy of these weakly bonded complexes. As the rare gas atom in the ionic complex is located above the plane of the benzene molecule, on the symmetry axis, the van der Waals complex retains a sufficiently high symmetry to also be Jahn–Teller distorted, and the IR spectrum of the ionic complex is expected to resemble closely that of the bare benzene cation. This is substantiated by the good agreement between the observed MATI spectra for bare benzene and those of its argon complexes.10

The experimental setup and the detection method that is used are similar to those described previously in our work reporting on the IR spectra of aniline–Ar,13 naphthalene–Ar,14 and pyrazine–Ar15 and are only briefly given here.

The experiments are performed using the free electron laser for infrared experiments (FELIX)16 as a source of intense pulsed infrared radiation. FELIX produces IR radiation that is continuously tunable over the 40–2000 cm−1 range. The light output consists of macropulses of about 4 μs duration containing up to 100 mJ of energy. Each macropulse consists of a train of micropulses that are 0.3–5 ps long and 1 ns apart. In the present experiments, the macropulse repetition rate is 10 Hz, and the bandwidth of the laser is typically set to 0.5%–1.0% of the central frequency. In the molecular beam apparatus, a pulsed valve backed with 2 bar Ar premixed with a fraction of a percent of C6H6 (C6D6), releases gas pulses of typically 50 μs duration. About 4 cm downstream, the molecular beam is skimmed upon entering a differentially pumped linear time-of-flight (TOF) mass spectrometer. The molecules in the beam interact with incoming UV laser beams as well as with the collimated FELIX beam (approximately 2–3 mm diameter) at the crossing point of the mutually perpendicular molecular beam axis, laser beam axis, and TOF-tube axis. A scheme of the experimental detection method that is used is given in Fig. 1. The van der Waals complexes under study are ionized via two-color (1+1′)-resonance enhanced multiphoton ionization (REMPI) via the 61(S1) state under field-free conditions, using the appropriate UV laser wavelengths.12,17 The energy of the ionizing photon is chosen such as to bring the van der Waals complex just barely above the ionization threshold. With an excess energy of less than 10 cm−1, even the lowest van der Waals mode of the complex cannot be excited.18 Therefore, ions will only be prepared in the ground vibrational level, and will be about as cold as the neutral benzene–Ar complexes that they originate from. Shortly after production, the complex ions are irradiated with the macropulse of FELIX. The maximum energy density of FELIX used in these ex-
FIG. 1. Energy-level scheme for $\text{C}_6\text{H}_6$ and for $\text{C}_6\text{H}_6^–\text{Ar}$, indicating the experimental method used to record the IR spectrum of the complex cation.

The observed IR absorption spectrum of the benzene–Ar complex is expected as a single strong peak around 683 cm$^{-1}$ but has never been assigned. The corresponding mode in the C$\text{D}_6$–Ar complex is consequently safely assigned to the $v_{11}$ mode, which is of electronic-vibrational $E$ symmetry result, which are all IR active from the vibrationless level in the ion. Of these, two $A$ modes and two $E$ modes are in the high-frequency C–H stretching region, whereas the $E_{1u}$ component of the $v_{16}$ mode, known from ZEKE and MATI experiments to be located at 306 cm$^{-1}$ (264 cm$^{-1}$) for the C$\text{H}_6$ (C$\text{D}_6$) cation, is outside of our scan region at the low frequency side.

In the electronic ground state of neutral benzene, the $a_{2u}$ mode and the three $e_{1u}$ modes are IR active. When the vibrational symmetry is combined with the $E_{1g}$ electronic symmetry of the ground state of the cation, the number of possible vibrational modes is doubled. In particular, six ungerade modes of electronic-vibrational $A$ symmetry and ten ungerade modes of electronic-vibrational $E$ symmetry result, which are all IR active from the vibrationless level in the ion. Of these, two $A$ modes and two $E$ modes are in the high-frequency C–H stretching region, whereas the $E_{1u}$ component of the $v_{16}$ mode, known from ZEKE and MATI experiments to be located at 306 cm$^{-1}$ (264 cm$^{-1}$) for the C$\text{H}_6$ (C$\text{D}_6$) cation, is outside of our scan region at the low frequency side.

In the electronic ground state of the neutral molecule the IR active $a_{2u}$ $v_{11}$ mode [out-of-plane C–H(D) bending] is found at 671 cm$^{-1}$ (503 cm$^{-1}$) for C$\text{H}_6$ (C$\text{D}_6$). The strong IR active mode observed at 485 cm$^{-1}$ in the spectrum of the C$\text{D}_6$–Ar complex is consequently safely assigned to this $v_{11}$ mode, which is of electronic-vibrational $E_{1u}$ symmetry in the C$\text{D}_6$ cation. This mode has been observed before at 488 cm$^{-1}$ as a weak peak in ZEKE and MATI spectra, but has never been assigned. The corresponding mode in the C$\text{H}_6$–Ar complex is expected as a single strong peak around 650 cm$^{-1}$. In the spectrum, however, two almost equally strong peaks are observed at 631 and 698 cm$^{-1}$ instead, thus forming a textbook example of a Fermi resonance. To get a Fermi resonance with the $v_{11}$ $E_{1u}$ mode, a nearby combination mode of the same symmetry is required, for which the combination of the linearly Jahn–Teller active $e_{2g}$ mode $v_6$ and the quadratically Jahn–Teller active $e_{2u}$ mode $v_{16}$ is the
only candidate. From a two-mode \( \nu_6, \nu_{16} \) Jahn–Teller analysis, two components of the \( \nu_6, \nu_{16} \) combination mode with the correct symmetry are expected in this region, so there actually is a three level interaction possible. Assuming that the modes at 631, 661, and 698 cm\(^{-1}\) belong to the Fermi triplet, and taking the same interaction parameter between the \( \nu_{11} \) mode and either component of the combination band, the deperturbed frequency of the \( \nu_{11} \) mode is found at 665 cm\(^{-1}\), whereas the two \( E_{1u} \) components of the \( \nu_6, \nu_{16} \) combination mode are found at 656 and 669 cm\(^{-1}\). The interaction parameter that is found to describe the observed spectral structure correctly is 23 cm\(^{-1}\).

It is interesting to note that all three components of the Fermi triplet have also been observed in ZEKE and MATI spectroscopy on \( \text{C}_6\text{H}_6 \) at the same (within 2 cm\(^{-1}\)) frequency positions.\(^6\)\(^,\)\(^8\)\(^,\)\(^11\) Moreover, the almost completely mixed outer two components from this triplet indeed appear with the same intensity in these spectra. The two lowest frequency components of this triplet had been assigned to the two doubly degenerate components of the \( \nu_6, \nu_{16} \) combination band,\(^6\)\(^,\)\(^8\)\(^,\)\(^11\) but it was already noted that these modes would then be significantly shifted away from their expected frequencies.\(^11\) The highest frequency component of the Fermi triplet had not been assigned. The deperturbed values we find for the frequencies of the two components of the \( \nu_6, \nu_{16} \) combination band match the expected frequencies\(^11\) very well, and provide support to the two-mode Jahn–Teller model as presented by Goode, Hofstein, and Johnson.\(^11\)

Knowing that this Fermi interaction is in effect in the \( \text{C}_6\text{H}_6 \) cation, it also has to be present in the deuterated species. Due to a much larger frequency difference between the interacting levels, the effect of the interaction is less pronounced, though it nevertheless gives some IR intensity to the \( \nu_6, \nu_{16} \) combination bands seen at 602 and 619 cm\(^{-1}\). A similar deperturbation analysis on the triplet of modes observed at 485, 602, and 619 cm\(^{-1}\) yields deperturbed frequencies for the \( \nu_{11} \) mode of 494 cm\(^{-1}\) and for the \( E_{1u} \) components of the \( \nu_6, \nu_{16} \) combination mode of 598 and 615 cm\(^{-1}\), again in excellent agreement with calculations.\(^11\) An interaction parameter of 25 cm\(^{-1}\) is found to describe the observations most accurately.

There is another weak peak observed at 678 cm\(^{-1}\) in the spectrum of the \( \text{C}_6\text{D}_6 – \text{Ar} \) cation. This peak coincides exactly with the \( E_{1u} \) component of the Jahn–Teller active mode \( \nu_6 \) in the \( \text{C}_6\text{D}_6 \) cation,\(^7\)\(^,\)\(^8\)\(^,\)\(^11\) which might show up weakly in our spectrum due to the breaking of gerade–ungerade symmetry in the complex. Other possible low frequency ungerade modes of \( A \) and \( E \) symmetry that this feature could be assigned to are the components of the combination mode of the \( \text{b}_{1g} \) mode \( \nu_{11} \) at 418 cm\(^{-1}\) (344 cm\(^{-1}\) for \( \text{C}_6\text{D}_6^{10} \))\(^,\)\(^8\)\(^,\)\(^11\) with the \( \nu_{16} \) mode, but these are expected to be above 700 cm\(^{-1}\) for \( \text{C}_6\text{D}_6 \). The peak observed at 588 cm\(^{-1}\) in the spectrum of the deuterated complex might be attributed to the lowest frequency component of this \( \nu_4, \nu_{16} \) combination band, expected to be around 592 cm\(^{-1}\).\(^8\)

For the higher frequency peaks in either of the observed spectra assignment is only possible when a comparison can be made to calculated spectra. Preliminary calculations using the Becke3LYP method with Dunning’s D95(d,p) basis set as implemented in GAUSSIAN 98, and assuming a compressed \( D_{2x} \) distorted structure for the cation, have already yielded remarkable agreement with the observed IR spectra. However, high quality calculated IR spectra for the benzene cation in \( D_{6h} \) symmetry are required to make a meaningful comparison. As both frequency positions and line intensity information are available from the IR absorption spectra shown here, it is expected that it will be possible at that point to assign many of the ungerade fundamental modes in the cation unambiguously, even though the possible occurrence of Fermi interactions is likely to complicate the spectra throughout.

For the strong IR resonance at 485 cm\(^{-1}\) in the \( \text{C}_6\text{D}_6 – \text{Ar} \) cation, the absolute value for the fraction of IR-laser-induced dissociation products is measured as a function of IR-laser intensity, and the results are shown in the inset to Fig. 2. If absorption of a single IR photon leads to the appearance of benzene ion dissociation products, the benzene ion branching ratio will have a \( (1 – \exp(–\sigma I)) \) dependence on the IR-laser intensity \( I \), with \( \sigma \) the absorption cross section for the corresponding single photon transition. In particular, for low values of the product \( \sigma I \) the yield of benzene ions will depend linearly on the IR-laser intensity. If the absorption of more than one IR photon is required to overcome the dissociation limit, a higher-than-linear intensity dependence is expected for low values of \( \sigma I \). From the measured fractional dissociation curve it is concluded that the vibrational level at 485 cm\(^{-1}\) is located above the dissociation limit of the ionic \( \text{C}_6\text{D}_6 – \text{Ar} \) complex. This information combined with the 169 cm\(^{-1}\) redshift of the ionization potential of deuterated benzene upon complexation with an Ar atom\(^12\) puts an upper limit on the value of the dissociation limit \( D_0 \) of neutral \( \text{C}_6\text{D}_6 – \text{Ar} \) in its electronic ground state of \( D_0 < 316 \) cm\(^{-1}\). This is a more stringent upper limit than the presently accepted experimental upper limit of 340 cm\(^{-1}\).\(^10\) and still some 15 cm\(^{-1}\) below the value of \( D_0 \) found in recent highly accurate \textit{ab initio} calculations on the ground state of the benzene–Ar complex.\(^20\)\(^,\)\(^21\)

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