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Spectroscopy on triphenylamine and its van der Waals complexes

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Both vibrationally and rotationally resolved spectra of the $S_1 - S_0$ transition in jet-cooled triphenylamine (TPA) around 340-320 nm are reported. Medium resolution spectra (0.5-1.0 cm$^{-1}$ resolution) are recorded using (1 + 1)-resonance enhanced multi photon ionization (REMPI) with mass selective time-of-flight (TOF) detection in a pulsed molecular beam apparatus. The origin of the $S_1 - S_0$ transition is at 29520.7 cm$^{-1}$, higher than halfway to the ionization potential (IP) found at 6.89 eV. A vibrational progression in the symmetric torsion mode (114 cm$^{-1}$) as well as in the symmetric C-N stretching mode (280 cm$^{-1}$) is observed in the electronic spectra. The spectrum of the most abundant isomer of the TPA–Ar (TPA–Kr) complexes is blue-shifted by 211 cm$^{-1}$ (216 cm$^{-1}$) with respect to the spectrum of the free TPA molecule. High-resolution (the resolution mainly being determined by the natural linewidth of the transition, i.e. 36 MHz) spectra are recorded using laser induced fluorescence (LIF) in a cw molecular beam apparatus. Individual rotational transitions are resolved and the spectrum shows unambiguously that TPA is a symmetric top molecule. The rotational constant $B''$ in the $S_0$ state of TPA is equal to $B'' = 403.7$ ± 0.5 MHz. Upon $S_1 - S_0$ excitation both $B$ and $C$ increase with $7.4$ ± 0.1 MHz and $2.8$ ± 0.1 MHz, respectively. The spectrum of the blue-shifted TPA–Ar isomer is the spectrum of a symmetric top molecule as well, and therefore the Ar atom has to be located on the $C_3$ symmetry axis, either on top of or underneath the umbrella formed by the phenyl rings. It appears that when Ar or Kr forms a complex with TPA, the first Ar, Kr, atom goes preferentially in a position on the $C_3$ symmetry axis of TPA, a position which causes an abnormal blue-shift of the spectrum. With the first rare-gas atom located in this special position, the second rare-gas atom is forced into a “normal” position, i.e. above one of the phenyl rings, causing a normal red-shift with respect to the TPA–Ar complex.

1. Introduction

A spectroscopic investigation of triphenylamine (TPA) is of special interest for the understanding of the interaction between the lone pair electron of the N atom and the $\pi$ electrons of the adjoining aromatic phenyl groups ($\pi\pi$ conjugation). The interplay between $\pi\pi$ conjugation and steric hindrance becomes more important in going from aniline via diphenylamine to triphenylamine; the character of the lone pair electron in TPA will be strongly influenced by the molecular conformation. Up to now several experiments have been performed to determine the geometrical structure of TPA in the gas phase [1], in the liquid phase [2–5], as well as in solid films [6], but a definite structure for TPA has not yet emerged from these measurements. The geometrical structure of the TPA cation in the gas phase and in solid films has also only been determined indirectly via a comparison of measured and calculated photoelectron spectra [7].

In the present work both vibrationally and rotationally resolved spectra of jet-cooled TPA are reported. The geometrical structure of TPA in the $S_0$ and in the $S_1$ state is deduced from the measured rotational constants.

Molecular beam techniques have been used extensively to study van der Waals complexes consisting of organic aromatic hydrocarbons bonded to rare-gas atoms. Such heteroclusters have provided an interdisciplinary testing ground for the energetics and the dynamics related to intermolecular potential, cluster growth and solvent perturbation [8,9]. The microscopic solvent shifts of these heteroclusters are generally to lower energy, are linearly dependent on the polarizability of the rare-gas atom and are additive.
for rare-gas atoms in equivalent structural positions. Different structural isomers have different spectral shifts. A solvent shift towards higher energy has been reported for fluorobenzene [10], phenyl acetylene [11], para-xylene [12] and more recently for aniline [13,14]: the cluster of each of these molecules with three Ar atoms shows a blue-shift with respect to the cluster with two Ar atoms.

In the present work we report spectra of various isomers of the TPA-Ar, Kr vdW complexes. The spectra of the most abundant TPA-Ar, Kr isomers show a large anomalous blue-shift with respect to the spectrum of the free TPA molecule. The structure of these blue-shifted vdW complexes is determined, and the origin of the blue-shift is discussed.

2. Vibrationally resolved (1+1)-REMPI spectroscopy

2.1. Experimental setup

The search for the \( S_1 \rightarrow S_0 \) transition in TPA was made using pulsed lasers and a pulsed molecular beam apparatus at the IBM Almaden Research Center. The vapour pressure of TPA at room temperature is about \( 10^{-5} \) Torr, and laser vaporization followed by supersonic cooling in a rare-gas expansion was used to reproduce an intense pulsed beam of jet-cooled TPA. The laser desorption jet-cooling apparatus has been described in detail recently [15], and only a short experimental description will be given here.

A KrF excimer laser (approximately 50 \( \mu J \) in a 0.25 mm diameter spot) is used to desorb the TPA from a piece of fritted glass placed very close (within two nozzle diameters) to the orifice of a supersonic jet. The fritted glass acts as a matrix for the TPA and yields a steady supply of TPA to the surface. The supersonic jet is formed by expanding 8-10 atm of rare gas (Ar or Kr) through a 0.5 mm nozzle. The laser desorbed molecules are entrained in the jet and their internal degrees of freedom are cooled by multiple collisions in the expansion region. Rotational temperatures down to 5 K are obtained in this way. The jet-cooled desorbed molecules obtain the same velocity as the supersonically expanding rare gas and arrive in the detection chamber after having passed through a 1 by 4 mm slit skimmer.

In the detection chamber the laser desorbed TPA molecules are detected via resonant one photon \( S_1 \rightarrow S_0 \) laser excitation followed by one photon ionization, either by the same or by a longer wavelength laser. Nd: YAG pumped pulsed dye lasers with a typical bandwidth of 0.4 cm\(^{-1}\) are used for excitation and ionization. The ions are formed between the extraction plates of a Wiley-McLaren type linear time-of-flight (TOF) mass spectrometer. A mass resolution \( M/\Delta M \) of 300-400 is obtained, enough to separate TPA (245 amu) from its \(^{13}\)C containing isotope. Every single laser pulse yields a complete TOF mass-spectrum, which is displayed on a digital oscilloscope and stored in a PC. With the PC, gates are set over various mass ranges, and wavelength spectra of several molecules (or molecular fragments) can be measured simultaneously.

2.2. The \( S_1 \rightarrow S_0 \) transition of TPA

The origin of the lowest allowed singlet transition in TPA is found around 339 nm; the corresponding transitions in aniline and diphenylamine have their respective origins at 294 [16] and 308 nm [15]. In the upper part of fig. 1, the (1+1)-REMPI spectrum of jet-cooled TPA is shown. A tunable, pulsed dye laser with a fluence of 1.0 mJ/cm\(^2\) is used to induce the \( S_1 \rightarrow S_0 \) transition in TPA, and to subsequently ionize from the \( S_1 \) state. Positive ions of the parent mass (245 amu) are detected. Under these conditions, and even when two orders of magnitude higher laser fluences are used, no fragmentation of the TPA upon ionization is observed.

In the TPA spectrum shown in fig. 1a, two vibrational progressions are visible, corresponding to vibrational modes at 114 and 280 cm\(^{-1}\). Both of the progressions are very harmonic. These are the only two low-frequency modes that appear strongly when a low excitation fluence is used. Although not indicated in the figure, other slightly weaker fundamental modes can be recognized in the low-fluence excitation spectrum around 383, 583, 715 and 742 cm\(^{-1}\). All these modes combine again with the 114 and 280 cm\(^{-1}\) vibrations.

When higher laser fluences are used for the resonant excitation step, a number of extra peaks appear. A (1+1)-REMPI spectrum of jet-cooled TPA recorded with a laser fluence of 100 mJ/cm\(^2\) is shown in fig. 2. The stronger peaks in the spectrum are all
Fig. 1. Vibrationally resolved (1+1)-REMPI spectrum of jet-cooled TPA (a), TPA–Ar (b) and TPA–Kr (c). In all cases the parent ion is mass selectively detected. In the spectrum of TPA, two vibrational progressions are indicated (values in cm\(^{-1}\)). In the lower panels the shift of the van der Waals complexes relative to the free TPA is indicated (in cm\(^{-1}\)).

saturated by the excitation laser (they should thus be equally intense) and clipped by the detection electronics to better show the weaker features. The peaks at 114 and 280 cm\(^{-1}\) are indicated again to aid in comparing this spectrum to the one shown in fig. 1a. At low frequency there are extra peaks at 59, 131, 137 and at 143 cm\(^{-1}\). All these weaker peaks combine again with the 114 and 280 cm\(^{-1}\) fundamental. The two weak peaks indicated with an asterisk in fig. 1 show the same saturation behaviour as the strong fundamentals. These peaks have always approximately 1% of the intensity of the main peaks, also when low excitation fluences are used. Although these peaks are 23 and 32 cm\(^{-1}\) to the blue of the TPA origin, we believe them to be hot bands, originating from vibrationally excited TPA, as will be discussed in section 4.

In an experiment in which two frequency doubled pulsed dye lasers pumped with two independent Nd:YAG lasers were used, we measured the ionization potential (IP) of TPA. A low-fluence dye laser (1.0 mJ/cm\(^2\)) was used to resonantly excite TPA from the \(S_0\) to the \(S_1\) state, whereas a second dye laser with two orders of magnitude higher fluence was used to ionize TPA from the \(S_1\) state. A wavelength scan of the second dye laser, showing the onset of ionization, is shown in fig. 3. In fig. 3a the onset of ionization is shown for jet-cooled TPA, resonantly excited to the vibrationless level in the electronically excited \(S_1\) state. The onset of ionization is seen around 385 nm, at 25968 cm\(^{-1}\) to be more precise. If TPA is resonantly excited to the 114 cm\(^{-1}\) vibration in the \(S_1\) state, one would expect the onset of ionization to appear when the ionizing laser has an 114 cm\(^{-1}\) lower frequency, i.e. around 386.7 nm. In fig. 3b this spectrum is shown and there is only a 10 cm\(^{-1}\) red-shift of the strongest onset of the IP; however, this indicates that the Franck-Condon factors for the transition from the \(S_1\) state of the neutral to the ground state of the positive ion are more or less diagonal for this mode; from the one quantum excited 114 cm\(^{-1}\) vibration in the TPA \(S_1\) state preferentially TPA ground state ions that have one quantum excited in the corresponding vibration are produced. Moreover, it can be deduced from the spectrum in fig. 3b that the frequency of the vibration in the ion that corresponds to the vibration of 114 cm\(^{-1}\) in the neutral is approximately 10 cm\(^{-1}\) lower, i.e. around 104 cm\(^{-1}\). In fig. 3c something similar is seen when TPA is resonantly excited to the 280 cm\(^{-1}\) vibration in the \(S_1\) state and the ionization laser is scanned over the ionization onset. Although now a weak onset of ionization is observed around 389 nm, as expected, it can be concluded from the strong step in the ionization continuum around 385 nm that again preferen-
tially ions that have one quantum of the corresponding excited vibration are formed. The frequency of this vibration in the ion is just a few cm$^{-1}$ larger than 280 cm$^{-1}$.

The apparent IP of TPA, as deduced from the spectra shown in fig. 3 and from the absolute frequency of the origin of the $S_1 \rightarrow S_0$ transition (29521 cm$^{-1}$), is at 6.880 eV, in good agreement with a previously estimated value [17]. It should be noted that the measurements of the IP are performed in an electric field of approximately 300 V/cm, and the actual field-free value for the IP will be several thousandths of an eV higher [18].

From a measurement of the double resonance TPA-ion signal as a function of the time delay between the two pulsed lasers an upper limit of 6 ns is deduced for the lifetime of the TPA $S_1$ state; an exact value for the lifetime can not be determined in this way since the pulse length of each of the dye lasers is already 4–5 ns.

2.3. Spectra of TPA–Ar, Kr van der Waals complexes

We have reported previously that under slightly different desorption conditions, complexes between the laser desorbed molecules and the carrier gas can be produced, internally cooled and mass selectively detected [19]. $(1+1)$-REMPI spectra of TPA–Ar and TPA–Kr obtained that way are shown in figs. 1b and 1c, respectively. Both spectra are measured at the parent mass, i.e. at 285 amu for TPA–Ar and at 329 amu for TPA–Kr, $^{84}$Kr being the most abundant Kr isotope. In these spectra the intensity of the strongest peaks is about 200–500 times smaller than of that in the TPA spectrum. The most striking observation in both the spectrum of TPA–Ar and of TPA–Kr is both the size and the sign of the spectral shift with respect to the TPA origin. Never before has a blue-shift been observed in the complex of an aromatic molecule with either one Ar or one Kr atom, let alone a blue-shift of over 200 cm$^{-1}$.

One has to be careful in assigning the correct spectral features to the specific size cluster, even if mass selective detection is applied; higher TPA–Ar$_n$ clusters will fragment and give rise to a signal on the TPA–Ar$_m$ mass ($m < n$) as well. There are two approaches commonly used to overcome this problem. In the first approach care is taken not to detect the fragments of larger clusters, although they are still formed [20,21]. In the other approach, soft ionization (i.e. bringing the molecule just barely above the ionization limit) is applied to strongly limit the fragmentation process itself [14]. For this latter approach two different lasers would have to be used for TPA, as one-color $(1+1)$-REMPI of TPA can bring the ion about 3500 cm$^{-1}$ above the IP. In our measurements of the cluster spectra we recorded the spectra of TPA–Ar$_n$, Kr$_n$ for $n=0–4$ simultaneously, thereby being able to trace the origin of the various lines. All the spectral features observed in figs. 1b and 1c are due to the TPA complexed with one Ar, Kr, respectively. Although there was severe fragmentation of these complexes.

Fig. 2. Vibrationally resolved $(1+1)$-REMPI spectrum of jet-cooled TPA using 100 mJ/cm$^2$ for excitation and ionization. The stronger peaks in the spectrum are all off scale. Clear structure is observed up to at least 1000 cm$^{-1}$ above the $S_1$ origin.
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Fig. 3. Wavelength dependence of the TPA ion signal showing the onset of ionization after resonant excitation of TPA to (a) the vibrationless level in the $S_1$ state; (b) the 114 cm$^{-1}$ vibration in the $S_1$ state; and (c) the 280 cm$^{-1}$ vibration in the $S_1$ state.

A  

B  

C  

WAVELENGTH (nm)  

ION INTENSITY

(shifted position for an aromatic molecule complexed to an Ar atom, we tried very hard to optimize on this signal. There was no way, however, to get clearly resolved single sharp lines. Sharp structure does appear 211 cm$^{-1}$ blue-shifted from the TPA origin. We believe this to be the origin of one of the possible, and apparently one of the most stable, TPA–Ar isomers. The TPA fundamental modes at 114 and 280 cm$^{-1}$ are seen in the spectrum of the Ar complex as well, and from this a lower limit of 280 cm$^{-1}$ can be deduced for the binding energy of this TPA–Ar isomer in the $S_1$ state.

There is another strong peak at 239 cm$^{-1}$ to the blue of the TPA origin, which might be the origin of another isomer of TPA–Ar. Again, the main vibrational modes in the TPA molecule are built on top of this. One might also argue that this extra peak is due to a vdW Ar–TPA stretching mode at 28 cm$^{-1}$. In that case, however, more of these modes should appear with comparable intensity, but they do not. In addition, the relative intensity of the 211 and 239 cm$^{-1}$ shifted peaks changes with changing laser-desorption/jet-cooling conditions, i.e. with changing production conditions for the van der Waals complexes. Some extra peaks that we do observe at a slightly large frequency are attributed to vdW stretching modes of the TPA–Ar complex.

One might argue that the structure observed for the TPA–Ar complex to the blue of the TPA origin is due to a complex that is actually 69 cm$^{-1}$ red shifted, and of which we only see the TPA vibrational progression ($-69+280=211$). Although there is absolutely no signal visible at the thus predicted origin, we can also find strong experimental evidence against this conjecture by looking at the TPA–Kr complex. Kr has a larger polarizability than Ar (by a factor 1.5) and consequently a larger red-shift of the hypothetical origin (and of the TPA modes built on top of this) should be visible. Instead, a somewhat larger blue-shift (216 cm$^{-1}$) is observed for TPA–Kr than for TPA–Ar, ruling out this hypothesis. All in all the spectrum of TPA–Kr looks very similar to the TPA–Ar spectrum. There is some sign of a red-shifted isomer (somewhat further red-shifted than the TPA–Ar complex, as expected), but the strongest signal is from a 216 cm$^{-1}$ blue-shifted isomer. Again, the two strongest TPA modes are seen in the Kr complex as well. There is a set of two peaks visible that are ap-

(which made the spectra of these complexes appear just as strong at the TPA mass as at the parent mass of the complex) distinctly different spectral features were observed at the higher cluster masses.

About 30 cm$^{-1}$ red-shifted from the TPA origin, the TPA–Ar spectrum, displayed in fig. 1b, shows a broad structure. As this is the position where we expected the origin of the $S_1 \rightarrow S_0$ transition in the TPA–Ar complex to appear, i.e. it is the "normal" red-
approximately 21.5 and 26 cm\(^{-1}\) blue-shifted from the origin of this TPA–Kr complex. It might well be that one of them is actually the origin of another TPA–Kr isomer, while the other is due to a vdW TPA–Kr stretching mode.

We also measured the spectrum of the TPA–Ar\(_2\) complex, and this spectrum is shown in fig. 4, albeit on a different wavelength scale. The origin of the strongest TPA–Ar\(_2\) isomer was found at 29708 cm\(^{-1}\), about 24 cm\(^{-1}\) red-shifted with respect to the strongest TPA–Ar origin, but still with an overall blue-shift of 187 cm\(^{-1}\) with respect to the TPA origin. As indicated in the figure, the fundamental vibrations of TPA are seen in the spectrum of the Ar\(_2\) complex as well. Again, there are some extra peaks at low frequencies, 22 and 28 cm\(^{-1}\) further to the blue of the aforementioned TPA–Ar\(_2\) origin, which might either be due to vdW vibrations in the complex or due to other TPA–Ar\(_2\) isomers.

A first conclusion from these measurements is that when Ar or Kr forms a complex with TPA, the first Ar, Kr, atom goes preferentially in a “special” position, a position which causes an abnormal blue-shift of the spectrum. With the first rare-gas atom sitting in this special position, the second rare-gas atom is forced into a “normal” position, i.e. above one of the phenyl rings, causing a normal red-shift with respect to the TPA–Ar complex.

### 3. High-resolution LIF spectroscopy

#### 3.1. Experimental setup

In the Nijmegen laboratory high-resolution spectroscopy was performed to determine both the geometrical structure of TPA itself as well as that of the TPA–Ar complexes. Again, details of both the molecular beam machine as well as of the narrow bandwidth UV radiation source have been given elsewhere [22], and only a short description is given here.

TPA is heated in a quartz oven to approximately 200°C (10 Torr vapour pressure) and is expanded supersonically together with 0.6–1.0 atm Ar through a 0.15 mm nozzle, kept at a slightly higher temperature. The molecular beam is skimmed twice and enters a differentially pumped LIF detection chamber about 30 cm away from the beam orifice. There the molecular beam is crossed perpendicularly with the weakly focused UV laser beam. The TPA molecules are resonantly excited from the S\(_0\) to the S\(_1\) state and the total fluorescence back to the ground state is detected. The UV radiation is obtained by frequency doubling in a LiIO\(_3\) crystal inside the cavity of a single-frequency ring dye laser, operating on DCM. By pumping the dye laser with 6 W of an Ar ion laser (all lines), 0.2–0.5 mW of tunable radiation in the 339–336 nm range with an effective bandwidth of about 3.0 MHz is obtained. A single mode laser scan can be made over 50 GHz, enough to record the whole envelope of a single rovibronic band in the S\(_1\)–S\(_0\) electronic transition of TPA in a single scan. The TPA spectra are recorded together with the transmission peaks of a pressure and temperature stabilized interferometer with a free spectral range (in the UV) of 148.25 ± 0.05 MHz. The absolute frequency of the transmission peaks of the interferometer, and thereby the absolute frequency of the TPA lines, is determined by the simultaneous recording of the I\(_2\) absorption spectrum in a cell at the fundamental laser frequency. Although the I\(_2\) absorption spectrum is relatively sparse in this region, an accuracy better than 0.1 cm\(^{-1}\) can still be obtained for the absolute frequency, the error being mainly due to the determination of the centre of the broad and not always symmetric I\(_2\)-lines. Relative line positions are measured to an accuracy of 10 MHz.
3.2. The rotationally resolved spectrum of TPA

In the top portion of fig. 5 the measured high-resolution spectrum of the origin of the TPA $S_1 \rightarrow S_0$ transition is shown. The absolute frequency of the origin (0.0 on the scale in the figure) is at $\nu_{\text{vac}} = 29520.7$ cm$^{-1}$. The signal intensity of the strongest peak is around 5000 counts/s, whereas there is a continuous background of about 200 counts/s due to scattered laser radiation. This high-resolution spectrum is clearly that of a parallel transition of a symmetric top molecule. There is a strong Q-branch, blue degraded, an R-branch and a P-branch with a band head. The lower portion of fig. 5 shows a fit to the data. The formulas for a rigid oblate symmetric top molecule are used and these fit the data perfectly well. The constants we used are the rotational constant $B''$ in the $S_0$ state and $\Delta B$ and $\Delta C$, the differences of constants in the $S_1$ and $S_0$ state. The rotational constant determined by the moment of inertia around the symmetry axis of TPA, the c-axis, cannot be determined in a parallel transition of a symmetric top molecule. The values that we used in the fit are given in table 1. It should be noted that the values of $\Delta B$ and $\Delta C$ are much more precise than the value of $B''$; the inaccuracy in the latter constant is due to a slight drift of the interferometer during the scan of the spectrum and is the standard deviation in the value of $B''$ obtained on various days, scanning in either frequency direction. For the $C''$ constant a value of 230 MHz was assumed, based on an estimate of the geometrical structure of TPA, as will be discussed later. The exact value for this constant is not critical anyway; it only influences to a minor extent the intensities of the various lines. A perpendicular transition, which is intrinsically weak in the TPA $S_1 \rightarrow S_0$ band, has to be measured to determine the absolute value of $C''$ in TPA. To obtain the fit shown in fig. 5b, a rotational temperature of 4.0 K was assumed, a value that we typically find for the rotational temperature in large aromatic molecules using this molecular beam apparatus. J-values up to $J = 100$ were included in the fit; to reproduce the observed band head in the P-branch more accurately even higher J-values have to be included.

In fig. 6 an enlargement of a part of the experimental (upper) and fitted (lower) R-branch of TPA is shown. Individual J, K lines are resolved. In the fit a Lorentzian lineshape with a full width at half maximum (fwhm) of 40 MHz is assumed. The contribution to the linewidth due to residual Doppler broadening in the molecular beam (i.e. the Gaussian

![Image of the rotationally resolved spectrum of TPA]

Fig. 5. High-resolution LIF spectrum of the origin of the $S_1 \rightarrow S_0$ transition in jet-cooled TPA. The experimentally observed spectrum (a) can be fitted into the smallest detail, using the formulas for a rigid oblate symmetric top molecule, as shown in (b). Note the band head in the P-branch for large $J''$, $K''$ values.

### Table 1

<table>
<thead>
<tr>
<th>Rotational constants (MHz)</th>
<th>TPA</th>
<th>TPA–Ar</th>
</tr>
</thead>
<tbody>
<tr>
<td>$B''$</td>
<td>403.7 ± 0.5</td>
<td>280 ± 5</td>
</tr>
<tr>
<td>$C''$</td>
<td>230$^{a)}$</td>
<td>230$^{a)}$</td>
</tr>
<tr>
<td>$\Delta B = B'' - B''$</td>
<td>7.4 ± 0.1</td>
<td>-8.5 ± 0.5</td>
</tr>
<tr>
<td>$\Delta C = C'' - C''$</td>
<td>2.8 ± 0.1</td>
<td>2.8$^{b)}$</td>
</tr>
</tbody>
</table>

$^{a)}$ Calculated value for a TPA geometry with $\phi = 50.0^\circ$ and $\angle \text{CNC} = 120.0^\circ$.

$^{b)}$ Kept fixed to the value as obtained for TPA.
contribution to the linewidth) is known to be 12 MHz. The largest contribution to the observed linewidth therefore has to come from lifetime broadening. Deconvolution of the experimentally observed Voigt profile yields a Lorentzian contribution to the linewidth of around 36 MHz, implying a lifetime for individual $J', K'$ levels in the vibrationless $S_1$ state of TPA of $4.5 \pm 0.5$ ns.

In the fit to the experimental data the spin statistical weights have not been taken into account, as they are very close to one with five fermions present in each phenyl ring. No splitting is observed due to the isomeric right-hand rotating and left-hand rotating forms of TPA.

In addition to the origin of the $S_1 \rightarrow S_0$ transition we also measured the transitions to the 114 and 280 cm$^{-1}$ vibrations under high resolution. We found the same rotational constants within the quoted error bars. As the spectrum of the transition to either one of these vibrations is also that of a parallel transition of TPA, both vibrations are due to symmetric modes.

### 3.3. High-resolution spectra of TPA–Ar

To determine the geometrical structure of especially the blue-shifted TPA–Ar complex we measured its spectrum under high resolution, as had been done previously for other vdW clusters between aromatic molecules and rare-gas atoms in the same molecular beam apparatus [23,24]. The best experimental spectrum which we managed to obtain reproducibly for the 211 cm$^{-1}$ blue-shifted TPA–Ar isomer is shown in the upper part of fig. 7. Although it looks hopeless at first sight, there is some clear structure visible that we managed to simulate, as shown in the lower portion of the figure. Most importantly, the observed TPA–Ar spectrum can be simulated using the formulas for a rigid symmetric top.

In our simulation we assume the Ar atom to be located on the $c$-axis of the TPA molecule. Then neither $C''$ nor $\Delta C$ will change in going from TPA to TPA–Ar. Furthermore, we assume the individual lines to have a Lorentzian shape with a width (fwhm) of 40 MHz, just as for the free TPA. A rotational temperature of 5.0 K is taken for the simulation of the
TPA–Ar complex. This leaves only two constants that have to be varied, $B''$ and $\Delta B$, of which $B''$ can be reasonably estimated from the calculated position of the Ar atom on the c-axis in the minimum energy configuration, as will be discussed in section 4.3. Because the observed blue-shift indicates that the TPA–Ar complex is less bound in the excited state than in the ground state, one expects the N–Ar distance to increase upon $S_1\rightarrow S_0$ excitation, producing a negative value of $\Delta B$.

Having only these two parameters to vary, the simulated spectrum shown in fig. 7b is obtained with $B'' = 280$ MHz and $\Delta B = -8.5$ MHz, as also indicated in table 1. Since $\Delta B$ is negative, the R-branch shows a band head, whereas the Q-branch runs into the P-branch. The apparently single sharp lines around the origin are due to a coincidental bunching of a large number of lines, and no individual $J''$, $K''$ assignments can be made. The absolute frequency of the origin of the spectrum of this TPA–Ar isomer is 29731.2 cm$^{-1}$, so the spectrum of this isomer is blue-shifted by 210.5 cm$^{-1}$ with respect to the transition in TPA. The total number of lines included in the simulated part of the spectrum that is shown is about 5000.

Up to now we did not succeed in measuring the high-resolution spectrum of the (possibly) TPA–Ar isomer that is even 28 cm$^{-1}$ further blue-shifted nor of the less-structured red-shifted TPA–Ar isomer. Both in view of the low signal to noise ratio on the TPA–Ar complex that we did observe, as well as in view of the fact that the production conditions in the cw beam machine are very different from those in the laser-desorption jet-cooling apparatus, this is not too surprising.

4. Discussion

4.1. The geometrical structure of TPA

As already mentioned in section 1 there is some ambiguity in the literature on the geometrical structure of free TPA in the gas phase. The high-resolution spectra reported here give a straight answer to this question: within our present accuracy of 0.1 MHz, two of the rotational constants for TPA are identical, implying a symmetric top structure for TPA. This means that the TPA molecule has at least C$_3$ symmetry. For the rest of the discussion we will therefore use the geometry of TPA as depicted in fig. 8.

The three phenyl rings are assumed to be flat and identical to each other, with fixed C–C (1.379 Å) and C–H (1.070 Å) distances and $\angle$CCC angles and $\angle$CCH angles of exactly 120°. The values for the bondlengths are the averaged values as obtained from an X-ray crystallographic study of TPA in the solid [6]. Then there are only three other parameters left that are allowed to vary, i.e. the C–N distance (taken as 1.419 Å in the $S_0$ state of TPA), the angle $\phi$ by which the phenyl groups are rotated around the C–N bond and the $\angle$CNC angle. The angles are defined such that a fully planar geometry of TPA means $\phi = 0$ and $\angle$CNC = 120°. If $\angle$CNC is exactly 120° TPA...
has additional C₂ symmetry axes through each of the
three phenyl rings, yielding a D₃ symmetry for the
molecule. In the literature up to now no definite val-
ues for either φ or ∠CNC have been obtained, al-
though averaging of the existing literature values
would give a φ somewhere between 15° and 60° and
a ∠CNC angle that is quite large, between 120° and
114°.

The S₁→S₀ transition in TPA is probably best de-
scribed as a π*→n transition, which means that if TPA
is electronically excited from S₀ to S₁, one of the lone
pair electrons on the N atom is moved towards the
phenyl rings. Therefore, upon excitation a change is
mainly expected in the bonds and the angles involv-
ing the N atom. More specifically, the C−N bonds are
expected to shorten and to get stiffer upon S₁→S₀
excitation.

The vibrationally resolved electronic S₁→S₀ spec-
trum of TPA, shown in fig. 1a, shows a progression
in two low-frequency modes, indicating a substantial
change in the geometry of TPA upon excitation. The
280 cm⁻¹ mode, which we know to be symmetric
from its high-resolution spectrum, is most likely due
to the symmetric N−phenyl stretch mode. This mode
has been observed at 290 cm⁻¹ in Raman spectra of
TPA in a CS₂ solution [2]. More recently, the fre-
cuency of this mode has been calculated in the S₀ state
as well as in the ground state of the positive ion of
TPA to be around 300 cm⁻¹ [25]. A long progres-
sion in this mode is in full agreement with the ex-
pected shortening of the C−N bonds upon excitation.
The other low-frequency vibrational mode at 114
cm⁻¹ is also symmetric and is most likely due to the
symmetric torsion of the phenyl rings around the C−
N bond. The frequency of this mode has also been
calculated [25], and there is a large difference be-
tween the calculated frequency of this mode in the S₀
state of the neutral (74 cm⁻¹) and in the ground state
of the positive ion (104 cm⁻¹). From the IP mea-
surements as described in section 2.2 we know that
the frequency of the mode corresponding to the 114
cm⁻¹ vibration in the S₁ state is around 104 cm⁻¹ in
the ground state of the ion. Apparently the frequency
for the symmetric torsion mode increases substam-
tially in going from S₀ to S₁, indicating an increased
stiffness in the C−N bonds in the excited state. The
observed Franck–Condon intensity pattern in the
progression of this mode is in agreement with a change
in the angle φ of 2° to 3° in going from S₀ to S₁. A
Franck–Condon analysis does not give the sign of this
change.

In the recent theoretical calculation [25] two low-
frequency modes are found for TPA in the S₀ state
around 20–25 cm⁻¹. In the ground state of the ion
these modes are at a much larger frequency, around
45–50 cm⁻¹. It is expected that the frequency of the
corresponding modes in the S₁ state of TPA is close
to the frequency of those in the ion. Under our exper-
imental conditions these low-frequency modes are
certainly populated to some extent and transitions
from these nonsymmetric low-frequency modes in the
S₀ state to the corresponding modes in the S₁ state
will be blue-shifted with respect to the TPA origin.
The 23 and 32 cm⁻¹ blue-shifted lines that are ob-
served in the TPA spectra are therefore probably hot-
bands, as mentioned in section 2.2.

The rotational constants for TPA, as determined
from the high-resolution spectroscopy of TPA and as
given in table 1, can be fitted to several geometries.
If it is assumed that only the C−N bondlength and the
angle φ change upon excitation, then ΔB and ΔC in-
dicate a decrease of the angle φ of 2° to 3° and a de-
crease in the C−N bond length of 0.03 Å in going from
S₀ to S₁. This change in φ is in perfect agreement with
the observed Franck–Condon intensity distribution
in the progression for the symmetric torsion mode.
The change in C−N bond length is very moderate,
compared with for instance the situation in aniline
[14]. The absolute value of B" can be fitted to a set
of (φ, ∠CNC) angles, as indicated in table 2. For all
sets the changes in φ and the C−N bond length are as
mentioned before. As seen in table 2, if the N atom is
in the plane of the three adjacent C atoms the angle φ
is 50.0°. A more pyramidal structure of TPA is only
in agreement with the measured B" value if at the
same time the angle φ increases, i.e. if at the same
time the phenyl rings turn towards a more staggered
geometry. Instead of the ∠CNC often the “umbrella-
angle” Θ, i.e. the angle by which the C−N bonds are
bent out of the xy-plane, is used. These angles are re-
lated via: cos(∠CNC) = 1 − 1/3 cos² Θ.

4.2. The geometry of the various TPA−Ar, Kr isomers

For the 210.5 cm⁻¹ blue-shifted TPA−Ar isomer
the Ar is located on the C₃ axis, as is unambiguously
In the first column, various possible \((\phi, \angle \text{CNC})\) combinations that yield a rotational constant along the \(b\)-axis that is in agreement with the experimentally found value for \(B^*\) are given. In the electronic ground state of TPA, \(\phi\) and \(\angle \text{CNC}\) are the only variables; for all the bond lengths as well as for all the other angles, values as obtained from X-ray analysis of solid TPA are taken. In the second and third column the calculated minimum value for the TPA–Ar van der Waals potential on the \(z\)-axis as well as the distance of the Ar atom to the N nucleus are given for an Ar atom above and underneath the umbrella, respectively. In the last column the measured value for the Ar position, i.e. the position as deduced from the measured \(B^*\) value for the complex, is indicated.

<table>
<thead>
<tr>
<th>Possible ((\phi, \angle \text{CNC})) combinations (deg.)</th>
<th>TPA–Ar vdW min. above umbrella (cm(^{-1}), Å)</th>
<th>TPA–Ar vdW min. under umbrella (cm(^{-1}), Å)</th>
<th>Ar position ((\pm 0.10 \text{ Å}))</th>
<th>Ar position (Å, Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(50.0, 120.0)</td>
<td>-438, +4.22</td>
<td>-438, -4.22</td>
<td>+4.01, -4.01</td>
<td>+4.01, -4.01</td>
</tr>
<tr>
<td>(54.5, 119.0)</td>
<td>-391, +4.24</td>
<td>-420, -4.56</td>
<td>+3.74, -4.28</td>
<td>+3.74, -4.28</td>
</tr>
<tr>
<td>(59.0, 118.0)</td>
<td>-359, +4.34</td>
<td>-400, -4.78</td>
<td>+3.63, -4.39</td>
<td>+3.63, -4.39</td>
</tr>
<tr>
<td>(64.0, 117.0)</td>
<td>-331, +4.44</td>
<td>-383, -4.98</td>
<td>+3.54, -4.48</td>
<td>+3.54, -4.48</td>
</tr>
<tr>
<td>(70.0, 116.0)</td>
<td>-307, +4.56</td>
<td>-367, -5.18</td>
<td>+3.47, -4.55</td>
<td>+3.47, -4.55</td>
</tr>
<tr>
<td>(80.0, 115.0)</td>
<td>-284, +4.69</td>
<td>-331, -5.38</td>
<td>+3.41, -4.62</td>
<td>+3.41, -4.62</td>
</tr>
</tbody>
</table>

The shift of TPA–Ar relative to TPA is about 2.5% of the shift of TPA relative to benzene.

4.3. Calculations on the minimum energy TPA–Ar complex geometries

To rationalize the position of the Ar atom on the \(C_3\) axis in TPA, as well as to rationalize the apparent existence of various TPA–Ar isomers, the potential energy surface for an Ar atom around a TPA molecule in its electronic ground state is calculated. For this, atom–atom potential of the Lennard-Jones type are summed over all the atoms in TPA, as prescribed by Ondrechen et al. [26]. For the ground state of TPA interacting with an Ar atom, we use the same C–Ar, H–Ar and N–Ar interaction parameters as used in the calculation of aniline–Ar, cluster geometries [13]. The geometry of TPA in the electronic ground state is taken as discussed in section 4.1 and as shown in the high-resolution spectrum of this complex. There are of course two positions possible, one underneath and one above the umbrella; these positions are equivalent if \(\angle \text{CNC}\) equals 120°. The positions for the Ar atom on the \(C_3\) axis as deduced from the measured \(B^*\) value for the TPA–Ar complex are indicated in the last column of Table 2. The exact positions are of course strongly dependent on the choices that are made for the angles \(\phi, \angle \text{CNC}\). In all cases, however, the magnitude and sign of \(\Delta B\) for TPA–Ar imply that in the electronically excited state of the complex the Ar atom has moved considerably further away from the N atom, approximately by 0.20 ± 0.02 Å.

Qualitatively, the fact that the Ar atom moves further away from the N atom upon electronic excitation is in agreement with the observed blue-shift of the spectrum; both facts indicate that the Ar atom is less bound in the \(S_1\) state than in the \(S_0\) state.

The question remains, however, which effect causes this abnormal blue-shift. As the electronic transition in TPA is probably best described as a \(\pi^*\)–\(\pi\) transition, there is a larger electron density on the \(C_3\) axis, due to the lone pair electrons of the N nucleus, in the ground state of the TPA molecule than in the electronically excited state. Therefore a stronger overlap of the Ar atom with the electron density of TPA is expected in the ground state than in the electronically excited state. If attractive forces between the Ar atom and the TPA molecule are more important than the repulsive contributions, this would explain the sign of the observed electronic shift of the TPA–Ar complex. An explanation analogous to the one given by Hermine et al. to explain the blue-shift for one of the observed aniline–Ar, isomers [14] holds for the TPA–Ar complex; the Ar atom sitting on the \(C_3\) axis slightly hinders the electron density on the N nucleus to be hybridized with the \(\pi\) system of the phenyl rings as much as it would like to, i.e. as much as it is in the free TPA. This then causes a kind of a chemical shift in the complexed TPA, bringing its spectrum more in the direction of the lowest singlet–singlet transition in benzene, this is, to the blue. It should be noted that the shift of TPA–Ar relative to TPA is about 2.5% of the shift of TPA relative to benzene.
For all the different ($\phi$, $\angle$ CNC) combinations shown in table 2, the two-dimensional minimum energy surface for an Ar atom attached to the TPA molecule is calculated. The plane defined by the N atom and the three adjacent C atoms (if $\angle$ CNC is 120.0°) is taken to be the xy-plane. The z-axis is along the C$_3$ symmetry axis of TPA, positive z-values indicating a position above the umbrella. In (more or less) planar molecules, a two-dimensional picture of the binding energy versus position of the Ar atom relative to the molecular framework is often given by taking a cut through the potential energy surface at a certain height above the molecule. In the case of TPA this would not make any sense, as the molecule is not planar at all. Therefore, the two-dimensional contour plots shown for two different sets of possible ($\phi$, $\angle$ CNC) angles in fig. 9, are 2D projections of the 3D minimum energy surface on the xy-plane. The minimum energy values in a certain point ($x$, $y$) are calculated along the z-coordinate, from $z = \infty$ to $z = 0$ and from $z = 0$ to $z = -\infty$. The values for the energy minima thus obtained are plotted in a contour plot in the left and right column of fig. 9, respectively. Different values of $z$ belong to each of the points in the figure. Contour lines are drawn in 40 cm$^{-1}$ intervals, and the contour lines at $-320$ cm$^{-1}$ are indicated in each of the plots.

A first inspection of fig. 9 shows that it is indeed energetically favourable for the Ar atom to sit on the C$_3$ symmetry axis of the TPA molecule, although there

![Fig. 9. Topographic plots of the TPA-Ar potential energy surface for two possible ($\phi$, $\angle$ CNC) combinations. For the combination A (50.0°, 120.0°) and B (59.0°, 118.0°) the minimum energy for the TPA-Ar complex for a certain value of ($x$, $y$) is calculated as a function of $z$, for z-values from $+\infty$ to $0$ (left column) and for z-values from $0$ to $-\infty$. So the left column shows contour plots for positive z-values (positions above the umbrella) whereas in the right column the corresponding TPA-Ar potential energy surface for negative z-values is projected on a 2D plane. Contour lines are 40 cm$^{-1}$ apart.](image-url)
are also some other, and even deeper, minimum energy configurations possible. In fig. 9a the minimum energy on the C₃ symmetry axis is $-438 \text{ cm}^{-1}$ and this value is obtained for a $z$-value of $z = \pm 4.22 \text{ Å}$. The even deeper off-axis minima, indicated with number (1), (2) and (3), belong to $z$-values closer to the $xy$-plane. The minimum numbered (1) is for a $z$-value around $z = +1.0 \text{ Å}$ and the minimum numbered (3) is an equivalent minimum on the other side of the TPA, so for $z = -1.0 \text{ Å}$; since in fig. 9a $\angle \text{CNC}$ is equal to $120^\circ$, positions above and below the TPA are identical. The equivalent minima numbered (2) are in the $z=0$ plane and these act as “gateways” for the Ar atom to move to the other side of the molecule. A closer inspection of the contour plots learns that the minimum energy surface is critically dependent on the chosen TPA geometry. In going from the flat geometry (fig. 9a) towards a more umbrella-like geometry (fig. 9b) several features change. First of all, as $\angle \text{CNC}$ decreases, the “top” and “bottom” of TPA are no longer identical. The minimum on the symmetry axis for both positive and negative values of $z$ becomes less deep, distinctly more so for the minimum on top of the umbrella (positive $z$-values) than for the one underneath. (Note that the first contour line away from the $C_3$ axis in fig. 9b is at $-320$ and at $-360 \text{ cm}^{-1}$ for the left and right column, respectively.) Secondly, the various minima between the phenyl rings gradually merge together and get deeper and deeper, much deeper than the minima on the $C_3$ axis. For $(\phi, \angle \text{CNC})$ combinations further down in table 2, first the minimum on the $C_3$ axis above the umbrella, but the eventually also the minimum on the $C_3$ axis underneath the umbrella, completely disappears.

Care should be taken in the interpretation of these contour plots. The Lennard-Jones model is too simplistic to describe the complicated vdW interactions. The atom–atom interaction parameters have been used extensively and have been optimized for a number of aromatics, but in practically all of these systems the rare-gas atom is located more or less above the centre of a benzene ring. It has been pointed out recently [27] that in this case the simple Lennard-Jones atom–atom potential model works remarkably well, but that this is no longer true when the rare-gas atom is further off to the side of the benzene plane. In the case of TPA–Ar, minima are found either on the symmetry axis, i.e. far away from the centrelines through the phenyl rings, or above one of the phenyl rings. For the latter minima the interaction with one of the rings is probably described quite correctly, but this is certainly not the case for the interaction with the other two rings. Therefore, care should be taken especially when the relative depths of the various minima for the TPA–Ar complex are compared, since now binding to different positions in the TPA molecule is compared.

The position of the various minima is more likely to be correct, however, and it is clear from the calculations that, generally speaking, only two types of TPA–Ar complexes will be possible, i.e. either with the Ar atom on the $C_3$ symmetry axis or with the Ar atom above one of the phenyl rings. Experimentally we do observe the isomer with the Ar on the $C_3$ axis to be the most abundant (this is the blue-shifted complex), although there is a sign of the presence of the other type of TPA–Ar complex as well (the slightly red-shifted complex).

For each set of $(\phi, \angle \text{CNC})$ indicated in table 2, the depth of the calculated potential energy minima on the C₃ axis as well as the $z$-coordinate for which these minima appear are given in the second and third column. The measured Ar–N distance, i.e. the Ar–N distance inferred from the high-resolution LIF spectrum of the 211 cm⁻¹ blue-shifted TPA–Ar isomer, is given in the last column of table 2. Of course, two positions are always possible, and the fitted Ar–N distance is dependent on the chosen TPA geometry. Best agreement between the calculated and observed Ar position is obtained if a planar TPA geometry is assumed, although a slightly bent TPA geometry cannot be ruled out.

5. Conclusions

The rotationally resolved spectrum of the $S_1 - S_0$ transition of jet-cooled TPA is the spectrum of a symmetric top molecule. The identical phenyl rings are rotated around the C–N bonds out of the planar geometry over an angle $\phi$ of about $50^\circ$. The $\angle \text{CNC}$ angle is close to or equal to $120^\circ$ for gas phase TPA. Upon electronic $S_1 - S_0$ excitation, there is only a minor change in the geometry of the molecule; the C–N bond length decreases by approximately 2% whereas
The phenyl rings rotate over 2–3° towards a flatter geometry.

The spectrum of the most abundant TPA–Ar complex shows an abnormal blue-shift with respect to the spectrum of the free TPA molecule. The high-resolution spectrum of this blue-shifted complex is again that of a symmetric top molecule, indicating that the Ar atom is located on the C₃ symmetry axis of TPA. The spectrum of another possible TPA–Ar complex shows a “normal” red-shift, and in this case the Ar atom is most likely located above one of the phenyl rings.

The detailed interaction mechanism responsible for the observed blue-shift of the TPA–Ar complex is not understood yet. The TPA molecule, being a symmetric top molecule, should be perfectly suited to study the effect of an aromatic-molecule–rare-gas interaction in which the rare-gas atom is bound to a less common place of the aromatic molecule. Moreover, the data presented here can be used to test various (empirical) models for the vdW interactions.

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References