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Photodissociation UV-VIS spectra of cold protonated azobenzene and 4-(dimethylamino)azobenzene and their benzenediazonium cation fragment

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Abstract:

Gas phase photodissociation electronic spectra of protonated azobenzene (ABH\textsuperscript{+}) and 4-(dimethylamino)azobenzene (dmaABH\textsuperscript{+}) were measured in a cryogenically-cooled ion trap at temperatures of a few tens of Kelvin. Experimental results were complemented with electronic structure calculations in the ground state at the MP2/cc-pVDZ level of theory, and in the low lying excited states using the RI-CC2 method. Calculated energies revealed that only the \textit{trans} isomers of the azonium molecular ions (protonation site on the azo group) will likely exist in the trap at the temperatures achieved in the experiment. The first transition of \textit{trans}-ABH\textsuperscript{+} was found \(\pi\pi\rightarrow\pi\pi^*\) in character, and the absorption band in the spectrum appears strongly red-shifted from that of the neutral molecule. The calculations showed that upon excitation the quasi-planar ground state (S\textsubscript{0}) transforms into a chair-like excited state (S\textsubscript{1}) by twisting the CNNC dihedral angle about 96 degrees. A 41 cm\textsuperscript{-1} active vibrational progression found in the ABH\textsuperscript{+} spectrum may be associated with the twisting of the azo bond. Conversely,
the electronic spectrum of dmaABH\(^+\)exhibits a steep and unstructured S\(_1\)↔S\(_0\)absorption corresponding to a less distorted S\(_1\) state. The next two quasi-degenerate bands in the ABH\(^+\) spectrum evidence sharper onsets and a charge transfer character. Using a second fragmentation laser and an additional He cooling pulse in the trap, it was possible to measure the UV spectrum of cold benzenediazoniumfragments.

1. Introduction:

In the electronic ground state, azobenzene(C\(_6\)H\(_5\)−N=N−C\(_6\)H\(_5\), hereafter denoted AB) can adopt cis and trans configurations defined by the relative orientation of the two phenyl rings attached to the azo group, the trans isomer being approximately 0.6 eV\(^1\) more stable than the cis one, and the trans→cis isomerization barrier in the gas phase is about 1.1 eV.\(^2\) It is well known\(^3,4\) that trans→cis isomerization of AB takes place in the electronic excited state when the molecule is excited at 365 nm, whereas the reverse reaction can be thermally activated in the ground state (in the dark, the half-life of cis-AB in solution is several hours, at room temperature) or induced photochemically by irradiation at 420 nm (lifetime in the ps scale).\(^5\) The photoisomerization mechanism of AB remains a matter of debate despite many years of experimental\(^6,9\) and theoretical\(^10,13\) studies. These results have been interpreted in terms of the contribution of two basic pathways that can in principle take place depending on the excitation scheme used: an inversion mechanism involving an in-plane bending of the azo-ring bonds and an out-of-plane rotation of the NN double bond. The advantageous photochromic properties of AB and its derivative compounds, as well as the ease of attaining optical control of the switching process between two photostable states, motivated research concerning the use of these molecules as molecular devices. A wide variety of complex molecular systems functionalized with ABs that exhibit phototiggerable optical, mechanical and functional changes have been reviewed in the last years.\(^14,17\)

When dissolved in strong acidic aqueous solutions, AB can attach a hydrogen ion to a free electron pair in the azo group forming azonium ions (see Scheme 1a for the particular case of protonated trans-azobenzene, denoted trans-ABH\(^+\)). In azonium ions the positive charge is delocalized by resonance in the aromatic system, lowering the energy of the \(\pi^*\) level. As a result, the \(\pi\pi^*\) absorption band of trans-ABH\(^+\) is shifted ~100 nm to the red with respect to the same transition of AB.\(^18\)
The introduction of additional functional groups on the phenyl rings of AB can dramatically alter the absorption spectra and the photochemical properties. Aminoazobenzenes are of particular interest because these molecules are known to experience tautomeric equilibria in solution, as a result of their multiple protonation sites, and therefore significant pH dependence in their optical absorption spectra. In acidic solutions, protonated aminoazobenzenes (denoted aABH⁺s) may form azonium or ammonium ions depending, respectively, on whether the hydrogen ion is attached to the azo or the amino group. See Scheme 1b for the particular case of protonated trans-4-(dimethylamino)azobenzene (denoted trans-dmaABH⁺). There are two possible structures for the azonium tautomer, depending on the protonation site, indicated in Scheme 1b as (I) when the H atom is next to the unsubstituted ring and (II) when it is attached to the other nitrogen atom. The positive charge in the azonium form (I) is easily delocalized on the aminosubstituted aromatic ring by formation of a quinoid structure and, consequently, the NN bond adopts a hydrazone character. On the other hand, the ammonium tautomer localizes the positive charge on the sp³ nitrogen atom, preserving the azo character of the NN bond. The equilibrium between aABH⁺ azonium and ammonium ions in liquid solution has been extensively studied on account of the distinct UV-VIS absorption spectrum of the different tautomers: i) the visible absorption at \( \lambda_{\text{max}} \approx 500 \text{ nm} \) is assigned to the \( \pi\pi^* \) transition of azonium ions, which is supported by the high \( \pi \) electronic delocalization in the molecule, and ii) the UV band appearing at \( \lambda_{\text{max}} \approx 320 \text{ nm} \) is attributed to the ammonium ions, because this type of tautomer is expected to have a spectrum similar to that of the neutral AB molecule (\( \lambda_{\text{max}} \approx 321 \text{ nm} \)).
Scheme 1. a) Azonium form of protonated trans-azobenzene (trans-ABH⁺). b) Azonium forms (I) and (II), and ammonium form of protonated trans-4-(dimethylamino)azobenzene (trans-dmaABH⁺). Molecular ions studied in the present paper are underlined.

Protonated azo compounds show faster cis→trans thermal isomerization rates, which has been explained in terms of a weaker double bond character of the azo bond due to the strong electron withdrawing effect of the positive charge introduced in the molecule upon protonation.\textsuperscript{20,21} The possibility of modulating the activation barrier for the cis→trans reaction of azo compounds by triggering the local pH has been proposed in the literature\textsuperscript{19,20} as an additional means of controlling a photodevice.

In spite of the abundant research available in the literature on the structural and photophysical properties of protonated azo compounds, no experimental studies have been performed in the gas phase for these molecular ions. Spectroscopic information about isolated ABH⁺ and aABH⁺’s will contribute to clarify the assignments of the bands observed in solution by working in a solvent-free environment, allowing a more reliable comparison of
the experimental data with the calculations. Moreover, an additional advantage of probing isolated ions at very low temperatures in the gas phase is the very low possibility that intramolecular proton transfer reactions take place. Therefore, structural details as the protonation site\textsuperscript{22} in the molecule and the flexibility of the molecular structure upon excitation will also be attainable from the gas phase spectra of the cold ions complemented with ab initio calculations.

In this work we have measured the UV-VIS spectra of isolated cold ABH\textsuperscript{+} and dmaABH\textsuperscript{+} ions via photofragmentation spectroscopy. The ions were prepared and cooled down to few tens of Kelvin into a cryogenic ion trap. Under these conditions and according to ab initio calculations both molecules are protonated on the azo group (azoniumtautomer) and the ions adopt a non-planar configuration (see text). We have also recorded the UV-VIS spectrum of the benzenediazonium cation (C\textsubscript{6}H\textsubscript{5}N\textsubscript{2}\textsuperscript{+}, denoted BD\textsuperscript{+}) one of the most abundant photofragments generated from the studied protonated azocompounds. This cation is a fundamental reaction intermediate with well-known applications to the dye industry, and for which UV-VIS spectroscopic characterization is scarce.

2. Methods:

The experimental set-up has been described in detail in earlier work.\textsuperscript{23,24} Here, we will present a brief summary of the main components of the apparatus and give a short explanation of the technique used to prepare the ions and record the photofragmentation electronic spectra. The protonated parent ion (ABH\textsuperscript{+} or dmaABH\textsuperscript{+}) is produced in an electrospray source from a solution of water/methanol (1:1 in volume) at a concentration of 0.1 mM, containing a trace of acetic acid. The ions formed in the source are injected in a cryogenically-cooled Paul trap through a mass gate. It takes several tens of milliseconds to cool down the ions to a temperature of about 40 K by collisions with cold He atoms pulsed into the trap. After this time, a first laser pulse is triggered and photodissociation of the cold parent ions takes place. Ionic fragments as well as the remaining parent ions can be extracted from the trap, mass-analyzed in a time of flight (TOF) mass spectrometer and their intensities quantified as a function of the laser wavelength. In addition, the ion fragments can be cooled down by introducing a second He pulse in the trap 1 ms after the fragmentation laser pulse, and probed by a second laser pulse triggered after a delay of some tens of milliseconds. Well resolved
electronic spectra of cold daughter ions can be obtained in this way. In order to ensure proper cooling conditions in the trap (product ion temperatures as low as those of parent ions, namely $T<40$ K), a solution of benzylaminewas occasionally electrosprayed in the source and the absence of vibrational hot bands in the benzylium ion ($mlq = 91$) spectrum (produced by photofragmentation of protonated benzylamine) was verified. One or more specific photofragments can be removed from the ion trap by exciting their axial micromotion applying specific auxiliary dipolar radiofrequencies, according to a procedure recently implemented in the set-up.\textsuperscript{25}

Two optical parametric oscillator lasers (EKSPLA model-NT342B) were used in the experiments, operating at a 10 Hz repetition rate and having 8 cm$^{-1}$ of spectral resolution and a scanning step of 0.02 nm. The lasers were focused to a 2 mm$^2$ spot in the trap and the laser power was roughly 5 mW.

Abinitio calculations were carried out with the TURBOMOLE program package,\textsuperscript{26} making use of the resolution-of-the-identity (RI) approximation in order to evaluate the electron-repulsion integrals. Ground state ($S_0$) equilibrium geometries of protonated species were optimized at the MP2 (second order Möller–Plesset perturbation) level of theory using the cc-pVDZ (correlation-consistent polarized valence double-zeta) basis set. Vertical excitation energies, oscillator strengths and equilibrium geometry of the lowest excited singlet states were computed at the CC$^2$\textsuperscript{27} (second order Approximate Coupled Cluster) level, since standard time-dependent DFT (Density Functional Theory) is well-known to produce substantial errors in representing charge transfer (CT) excited states.\textsuperscript{28} Moreover, we have chosen the RI-CC2 method because it proved useful in describing the electronic transitions of similar aromatic ions with nitrogen functional groups.\textsuperscript{29,30} In all the calculations the starting geometries had Cs symmetry and were constrained to distinguish between $\pi$ or $\sigma$ orbitals and $A'$ or $A''$ excited states. In the case of the benzenediazonium ion fragment, adiabatic excitation energies were calculated in addition to the vertical excitation values. For doing this, the energies of the ground and the first two excited states were computed for the optimized $S_1(A')$ and $S_2(A')$ excited state geometries, in Cs symmetry. Additional excited state optimizations were performed for this system and protonated azobenzene, relaxing the condition of maintaining Cs symmetry and allowing non-planar geometries.
3. Results:

3.1 Electronic spectra of ABH+ and dmaABH+ parent ions

Fig. 1a shows the difference mass spectrum (in presence and in absence of the fragmentation laser) obtained upon photodissociation of the parent ion ABH+ ($m/q = 183$, negative peak) at 384.0 nm. The two most abundant ion fragments (positive peaks) correspond to the phenyl cation ($m/q = 77$), formed by heterolytic cleavage of the C–N bond between one of the aromatic rings and the azo group, and BD+ ($m/q = 105$), formed by loss of benzene (B). The CID mass spectrum at low fragmentation energy also yields the phenyl cation as the main fragment, followed by BD+ which is about 5 times less intense.

![Figure 1](image)

**Figure 1.** a) Difference mass spectrum (fragmentation laser on – laser off) of ABH+ ($m/q = 183$, negative peak) at $\lambda = 384.0$ nm. Ion fragment signals (positive peaks) at $m/q = 77$ and 105 correspond respectively to phenyl and BD+ cations. b) Photodissociation electronic spectra of ABH+ obtained from $m/q = 77$ and 105 mass channels. The absorption starts at $\lambda = 480.3$ nm ($m/q = 77$ spectrum) and exhibits a vibrational progression, zoomed in the inset at a higher spectral resolution. Loss of benzene from ABH+ to yield BD+ ($m/q = 105$ spectrum) is restricted to the wavelength range 300–410 nm. c) Electronic spectrum of the conjugated acid of AB in strong acid solution (bulk).
Photodissociation electronic spectra of ABH⁺ are given in Fig. 1b for the mass channels $m/q = 77$ and 105. The absorption starts at $\lambda = 480.3$ nm for the spectrum recorded on $m/q = 77$ evidencing 3 additional electronic transitions: two sharp bands starting at $\lambda = 407.6$ nm and $\lambda = 389.9$ nm, and an UV band starting at $\lambda = 263.3$ nm. Both visible and UV absorptions are also present in the spectrum of the conjugated acid of AB measured in an ethanol-water solution having 65% sulfuric acid\textsuperscript{18} (Fig. 1c).

The first excited state exhibits a low frequency vibrational progression in the vicinity of the band origin that can be analyzed in terms of a 41 cm\textsuperscript{-1} active mode starting most likely at wavelengths 480.3 nm and 479.8 nm (FWHM of peaks = 12 cm\textsuperscript{-1}). In the inset of Fig. 1b there is a zoomed view of this spectral region where the laser was scanned in 0.02 nm steps to achieve the maximal spectral resolution. The transition starting at $\lambda = 480.3$ nm is very weak and cannot be straightforwardly assigned to the 0–0 transition. The absence of a strong 0–0 transition implies a large change in geometry in the excited state, as it will be discussed later. The second spectrum in Fig. 1b reveals that at $\lambda = 410$ nm a new fragmentation channel opens with the production of BD⁺ and then closes at about $\lambda = 300$ nm. In Scheme 3 we propose a mechanism for this fragmentation channel in the excited state, which involves a charge transfer in the molecule followed by an H atom migration.

Photodissociation difference mass spectrum of dmaABH⁺ ($m/q = 226$) at $\lambda = 275.0$ nm is given in Fig. 2a. As in the case of ABH⁺, phenyl ($m/q = 77$) and benzenediazonium ($m/q = 105$) cations are the main photofragments, followed by the dimethylaminophenyl cation ($m/q = 120$) and other minor fragments. Phenyl and dimethylaminophenyl cations likely originate from heterolytic cleavage of the C–N bond at both sides of the azo group, while BD⁺ forms by loss of benzene. In contrast to what is observed for ABH⁺, the photoproduct branching ratio of dmaABH⁺ is invariant as a function of the excitation wavelength and, therefore, the photodissociation electronic spectra in the Fig. 2b results from adding up the signal from all mass channels.
Figure 2. a) Difference mass spectrum (fragmentation laser on – laser off) of dmaABH\(^+\) \((m/q = 226\), negative peak) at \(\lambda = 275.0\) nm. Ion fragment signals (positive peaks) at \(m/q = 77\), 105 and 120 correspond respectively to phenyl, BD\(^+\) and dimethylaminophenyl cations. b) The photodissociation electronic spectra of dmaABH\(^+\) is the same for all fragment mass channels \((m/q 77, 105 \text{ and } 120)\) and therefore, we present the sum of all of them. The absorption starts at \(\lambda = 558.3\) nm, and the band rising is zoomed in the inset. c) Electronic spectrum of the conjugated acid of dmaAB in acid solution (bulk).\(^19\)

The electronic spectrum of dmaABH\(^+\) exhibits a two-band system, one in the visible and the other in a spectral range given by \(\lambda = 240–360\) nm. Considering the width of the bands it remains uncertain how many electronic states are involved in the absorption spectrum. The visible absorption of this ion is shifted about 80 nm to the red in comparison to ABH\(^+\) (the band starts at \(\lambda = 558.3\) nm), which is a direct consequence of the extended conjugation caused by the presence of the electron-donating dma group. The absorption spectrum of a solution of dmaABH\(^+\) in acidulated acetonitrile\(^19\) is shown in Fig. 2c for comparison.

The absorption bands of dmaABH\(^+\) are found broad and do not exhibit a sharp vibrational structure, in contrast to the ABH\(^+\) spectrum. This is shown in the inset of Fig. 2b.
where the onset of the visible band origin was scanned with a higher resolution. Although the onset of the band at \( \lambda = 558.3 \) nm does not appear to be a well-defined 0–0 transition, the absorption rises more steeply (within \( \sim 50 \text{ cm}^{-1} \)) than in the case of ABH\(^+\), as evidenced by comparing the low-energy parts of ABH\(^+\) and dmaABH\(^+\) spectra plotted in Fig. 3. It is not clear whether the broadened spectrum of dmaABH\(^+\) is originating from very low vibrations or from lifetime broadening, which would reveal that the excited state is relatively short-lived as it was found in protonated DNA bases. \(^{32}\) By using a multi-Lorentzian fit, a FWHM of \( \sim 70 \text{ cm}^{-1} \) can be determined for the two bands in the inset of Fig. 2.b, which accounts for an excited state lifetime of about 50 fs, if the broadening is exclusively limited by the lifetime.

\[ 17900 \quad 18000 \quad 18100 \quad 18200 \]

\[ \text{wavenumber (cm}^{-1}) \]

\[ \text{ABH}^+ \quad \text{dmaABH}^+ \]

**Figure 3.** Low-energy part (350 wavenumbers) of ABH\(^+\) and dmaABH\(^+\) spectra normalized by the corresponding absorption band maxima.

### 3.2 Electronic spectrum of the DB\(^+\) fragment
Photolysis of benzenediazonium salts in solution is known\textsuperscript{33} to occur principally via dissociative loss of molecular nitrogen, giving the corresponding phenyl cation as primary product. Likewise, the photofragmentation mass spectrum of isolated BD\textsuperscript{+} ions provides clear evidence for the formation of phenyl cations upon absorption of UV light. BD\textsuperscript{+} ions produced in the trap by photodissociation of ABH\textsuperscript{+} at \( \lambda = 384 \) nm were cooled down to \( T < 40 \) K before making use of a second laser to record the electronic spectrum. As an example, we present in Fig. 4 two mass spectra of BD\textsuperscript{+} ions recorded before (a) and after (b) excitation at \( \lambda = 265 \) nm, where it becomes clear that the phenyl cation is the main photoproduct. The absence of ABH\textsuperscript{+} parent ions and phenyl fragments before excitation was ensured by the use of auxiliary dipolar radiofrequencies\textsuperscript{25} which removed these ions from the trap.

Figure 4. a) Mass spectrum of ABH\textsuperscript{+} photofragments at \( \lambda = 384 \) nm after removal of phenyl and residual parent ions from the trap by applying an auxiliary radiofrequency; under these conditions, BD\textsuperscript{+}(m/q = 105) is the most abundant ion in the trap. b) Photodissociation mass spectrum of BD\textsuperscript{+} ions at 265 nm; the main resulting fragment is the phenyl cation (m/q = 77). c) Photofragmentation spectrum of BD\textsuperscript{+} ions obtained from the m/q = 77 mass channel. The band starting at \( \lambda = 276.0 \) nm is zoomed in the inset. d) Electronic spectrum of benzenediazonium fluoroborate in ethanolic solution (bulk).\textsuperscript{34}
In view of the high efficiency of production of BD⁺ fragments from ABH⁺ ions, we were able to perform a photodissociation experiment as a function of the wavelength of a second laser. The absorption spectrum of cold BD⁺ ions recorded in this way on the phenyl cation \((mql = 77)\) mass channel is shown Fig. 4c. The spectrum consists in two bands, one in the 280–330 nm spectral region starting at \(\lambda = 327.1 \text{ nm}\) and the other one starting at \(\lambda = 276.0 \text{ nm}\). In the inset of Fig. 4c the onset of the stronger band is zoomed at higher resolution (0.02 nm scanning step) with no evidence of sharp vibrational structures, despite the low temperature achieved. For comparison, the spectrum of benzenediazonium fluoroborate in ethanolic solution is given in Fig. 4d.\(^{34}\)

### 3.3 Calculations

Calculations done on ABH⁺ were restricted to ions protonated on one of the two equivalent nitrogen atoms of the azo group (Scheme 1a), which is the most plausible situation. We found that the *trans* isomer of ABH⁺ is the most stable structure, about 0.35 eV (MP2 level of theory, cc-pVDZ basis set) lower in energy than the *cis* homologue, which adopts a twisted conformation upon geometry optimization (Scheme 2a). The same result was obtained using DFT B3LYP/cc-pVDZ level calculations, where the *trans–cis* energy difference results 0.48 eV. Consequently, we focused our attention to the *trans*-ABH⁺ ion, since the population of the *cis* isomer is expected to be negligible in the trap. Compared to the neutral *trans*-AB molecule, for which a planar structure is well established, *trans*-ABH⁺ is slightly twisted out of the plane in the ground state due to steric hindrance between the acidic H atom and the adjacent aromatic ring. A small angle of 18° comes out from our \(S_0\) calculation (Scheme 2a), which is enough to break the Cs symmetry. The lowest frequency vibrational modes calculated for *trans*-ABH⁺ in the ground state are also depicted in Scheme 2a; they can be associated to a torsional movement around the azo bond, an out of the plane butterfly motion and a nearly-in-plane collective molecular bending. In panel b of Scheme 2 we show the optimized twisted structure of the \(S_1\) state of *trans*-ABH⁺ formed by torsion of the CNNC dihedral angle (96° in \(S_1\)), as well as the structure of *cis*-ABH⁺ which maintains its chair-like form in \(S_1\). We have observed in the optimization process that the \(S_1\) state of *trans*-ABH⁺ crosses the ground state through a conical intersection, as it will be discussed later.
Scheme 2. a) MP2-optimized structures of trans-ABH⁺ and cis-ABH⁺ in the S₀ state, and lower frequency vibrations of trans-ABH⁺. b) CC2-optimized structures of trans-ABH⁺ and cis-ABH⁺ in the S₁ state.

The calculated vertical transition energies of the Sᵢ=1−3 excited states, and the corresponding wavelengths and oscillator strengths are given in Table 1 for trans-ABH⁺, as well as the corresponding frontier orbitals. The energies of the band onsets measured in the experiment are also included in the table for comparison. The first excited state (S₁) exhibits a marked ππ* character (bonding π-HOMO to antibonding π*-LUMO transition) and an oscillator strength close to 1 (see Table 1). We have assigned the first band of the ABH⁺ spectrum starting at λ = 480.3 nm to the S₁<–S₀ transition, which represents a strong red shift in comparison to the analogous ππ*<–ππ excitation in neutral AB. The next two
transitions are close in energy, showing an important CT character as it is suggested by the shape of the molecular orbitals involved in the $S_2 \leftrightarrow S_0$ (LUMO$\rightarrow$HOMO$-1$) and $S_3 \leftrightarrow S_0$(LUMO$\rightarrow$HOMO$-2$) transitions.Upon excitation, electron density is transferred from the $\pi$ orbitals of rings 1 or 2 towards the protonated azo group. We have assigned the sharp bands observed at 407.3 nm and 389.9 nm to the $S_2 \leftrightarrow S_0$ and $S_3 \leftrightarrow S_0$ CT transitions, in that order. The CC2 vertical transition energies reported in Table 1 are always larger than the corresponding experimental energies since the excited state potential is not optimized.

<table>
<thead>
<tr>
<th></th>
<th>$E_0$/eV ($\lambda_0$/nm)</th>
<th>$E_{\exp}$/eV ($\lambda_{\exp}$/nm)</th>
<th>$f$</th>
<th>transition</th>
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<tr>
<td>$S_1$</td>
<td>2.94 (421.7)</td>
<td>2.58* (480.3)</td>
<td>0.91</td>
<td>LUMO$\rightarrow$HOMO</td>
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<tr>
<td>$S_2$</td>
<td>3.33 (372.3)</td>
<td>3.08 (407.6)</td>
<td>0.03</td>
<td>LUMO$\rightarrow$HOMO$-1$</td>
</tr>
<tr>
<td>$S_3$</td>
<td>3.34 (371.2)</td>
<td>3.21 (389.9)</td>
<td>0.01</td>
<td>LUMO$\rightarrow$HOMO$-2$</td>
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Table 1. Left panel: CC2 vertical excitation energies ($E_0$) for trans-ABH$^+$ in the $S_0$ geometry, transition wavelengths ($\lambda_0$), and oscillator strengths ($f$). Experimental wavelengths ($\lambda_{\exp}$) are taken from band origins; (*) weak band, the onset may not correspond to the 0–0 transition. Ground state geometry was optimized, in the Cs symmetry, at MP2 level of theory, using the cc-pVDZ basis set. Right panel: isodensity of the involved frontier orbitals. The ring closer to the protonation site is indicated as 1.

The band assignment presented in Table 1 allowed us to propose a plausible photofragmentation mechanism for the (ABH$^+$→ BD$^+$ + B) reaction which, as shown in Fig. 1b, only takes place when the excitation wavelength is restricted to the 300-410 nm spectral range. Under these conditions, both $S_2 \leftrightarrow S_0$ and $S_3 \leftrightarrow S_0$ excitations are able to neutralize the positive charge in the protonated azo group by CT from one of the rings, and this triggers the concurrent homolytic cleavage of the NH bond and an H radical migration to the other ring (see fragmentation pathways in $S_2$ and $S_3$ excited states in Scheme 3).
Unlike what happens with AB, where the molecule has two equivalent protonation sites in the azo group, dmaAB can be protonated in three different positions corresponding to each one of the lone electron pairs on the nitrogen atoms (see Scheme 1b). Due to the asymmetry of the molecule, the two nitrogen atoms of the azo group are slightly dissimilar, leading to the azonium structures indicated as (I) and (II). The $S_0$ geometry optimization of both azonium ions of trans-dmaABH$^+$ evidenced that the structure (I) is 0.06 eV lower in energy than (II) at the MP2 level, and 0.20 eV lower when using the DFT method. The small computed energy difference between isomers (I) and (II) points out that both azonium ions might in principle coexist in the trap under our experimental conditions. However, recent Raman studies$^{19}$ performed on dmaAB/acetonitrile acidic solutions excited on the band maximum of Fig. 2c evidenced selective enhancement of quinoid modes, implying that the preferred protonation site corresponds to the structure (I) (see Scheme 1). The third conceivable protonation site on the amino group leads to a significantly less stable tautomeric structure in the ground state, being 0.29 eV higher in energy than the azonium structure (I) at the MP2 level, and 0.75 eV higher from DFT calculations. The preference of structure (I) is
consistent with previous $^{13}$C- and $^{15}$N-NMR measurements combined with MINDO calculations$^{35}$ done on 4-aminoazobenzene, and is in accordance with the higher proton affinity computed for the dmaABazo nitrogen using semiempirical AM1 and ab initioSTO-3G studies.$^{36}$ By considering the low temperatures achieved in our experiment and taking into account the larger stability of the azonium structures derived from the calculations, it is possible to assume that the ammonium tautomer will likely not be present in the trap. Consequently, we have assigned the entire spectrum shown in Fig. 2b to the azonium form of $trans$-dmaABH$^+$, without the possibility to distinguish between isomers (I) and (II).

The vertical transition energies (ground state geometry) calculated for the first three excited states of isomers (I) and (II) of $trans$-dmaABH$^+$ are given in Table 2. The resulting energy values are close to those calculated previously$^{19}$ for the azonium ion of 4-aminoazobenzene. The excitation to the $S_1$ state (LUMO$\rightarrow$HOMO) can be characterized as a $\pi\pi^*\leftarrow\pi\pi$ transition with a significant oscillator strength, which can be associated to the first band of the dmaABH$^+$ spectrum starting at $\lambda = 558.3$ nm.

<table>
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<tr>
<th>isomer</th>
<th>$E_0$/eV ($\lambda_0$/nm)</th>
<th>$E_{exp}$/eV ($\lambda_{exp}$/nm)</th>
<th>$f$</th>
<th>transition</th>
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<tbody>
<tr>
<td>$S_1$</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(I)</td>
<td>2.62 (473.2)</td>
<td>2.22 (558.3)</td>
<td>1.35</td>
<td>LUMO$\rightarrow$HOMO</td>
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<tr>
<td>(II)</td>
<td>2.38 (520.9)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$S_2$</td>
<td></td>
<td></td>
<td>0.01</td>
<td>LUMO$\rightarrow$HOMO$\rightarrow$1</td>
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<tr>
<td>(I)</td>
<td>3.76 (329.7)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(II)</td>
<td>3.60 (344.4)</td>
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</tr>
<tr>
<td>$S_3$</td>
<td></td>
<td></td>
<td>0.01</td>
<td>LUMO$\rightarrow$HOMO$\rightarrow$2</td>
</tr>
<tr>
<td>(I)</td>
<td>4.00 (310)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(II)</td>
<td>3.77 (328.8)</td>
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**Table 2.** Left Panel: CC2 vertical excitation energies ($E_0$) of the azonium tautomer of $trans$-dmaABH$^+$ in the $S_0$ geometry, transition wavelengths ($\lambda_0$), and oscillator strengths ($f$). Experimental wavelength ($\lambda_{exp}$) is taken from the band origin. Ground state geometry was optimized in the Cs symmetry, at MP2 level of theory, using the cc-pVDZ basis set. Right panel: isodensity of the involved frontier orbitals. The ring closer to the protonation site is indicated as 1.

According to our calculations, the second band system observed in the experiment starting at $\lambda = 376.7$ nm (3.29 eV) could originate from the $S_2$$\leftarrow$$S_0$ or
$S_3 \leftarrow S_0$ transitions although, unlike what happens in unsubstituted ABH$^+$, these two transitions cannot be easily assigned in the spectrum of dmaABH$^+$. The excited state energies of isomers (I) and (II) in Table 2 are very close and, since the experimental bands are very broad, it is not possible to exclude the contribution of isomer (II) to the spectrum.

Finally, both vertical ($E_0$: $S_0$ geometry) and adiabatic ($E_1$: $S_1$ geometry, and $E_2$: $S_2$ geometry) low-lying excitation energies were computed for the benzenediazonium fragment and summarized in Table 3, relative to the ground state energy. The two band systems observed in the spectrum of BD$^+$ were re-assigned to the first two $\pi\pi \leftarrow \pi\pi^*$ transitions ($S_1 \leftarrow S_0$ and $S_2 \leftarrow S_0$) shown in the table. As usual, calculated vertical excitations $E_0$ are about 0.5 eV higher than the experimental values, and the adiabatic transition energies $E_1$ ($S_1$ geometry) and $E_2$ ($S_2$ geometry) become closer to the experimental values. The experimental band onsets included in the table may not correspond to 0–0 transitions, considering the absence of sharp absorption peaks in the spectrum.

<table>
<thead>
<tr>
<th></th>
<th>$E_0$/eV ($\lambda_0$/nm)</th>
<th>$E_1$/eV ($\lambda_1$/nm)</th>
<th>$E_2$/eV ($\lambda_2$/nm)</th>
<th>$E_{exp}$/eV ($\lambda_{exp}$/nm)</th>
<th>$f$</th>
</tr>
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<tr>
<td>$S_1$</td>
<td>4.40 (281.8)</td>
<td>3.98 (311.6)</td>
<td>3.79* (327.1)</td>
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<td>0.04</td>
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</tr>
<tr>
<td>$S_2$</td>
<td>4.97 (249.5)</td>
<td>4.82 (257.3)</td>
<td>4.49* (276.0)</td>
<td></td>
<td>0.36</td>
</tr>
</tbody>
</table>

Table 3. Left Panel: CC2 vertical ($E_0$: $S_0$ geometry) and adiabatic ($E_1$: $S_1$ optimized Cs geometry, $E_2$: $S_2$ optimized Cs geometry) excitation energies of BD$^+$, transition wavelengths ($\lambda_{\text{vis}}$), and oscillator strengths ($f$). Experimental wavelengths ($\lambda_{\text{exp}}$) are taken from band origins; (*) weak band, the onset may not correspond to the 0–0 transition. Ground and excited state geometries were optimized in the Cs symmetry, respectively at MP2 and CC2 level of theory, using the cc-pVDZ basis set. Right panel: BD$^+$ geometries corresponding to the ground state ($S_0$) and the conical intersections ($S_0 = S_1$) and ($S_1 = S_2$), determined by performing out-of-plane optimizations of the first and second excited states.

4. Discussion:
The slow rising of the first trans-ABH\textsuperscript{+} band system (Fig. 1b) implies a substantial change in the potential upon photoexcitation and hence a low Franck Condon factor for this transition. The large geometrical change adopted by the molecule upon excitation can be visualized by comparing the optimized structures of trans-ABH\textsuperscript{+} in the S\textsubscript{0} and S\textsubscript{1} states that are displayed in Scheme 2. It results that half of the molecule flips to a perpendicular position leading to a chair-like structure, with an CNNC dihedral angle of about 96 degrees. This implies that in the S\textsubscript{1} excited state the observed planar-to-chair isomerization reaction of trans-ABH\textsuperscript{+} proceeds via a mechanism that resembles the rotation pathway described for the trans\textrightarrow{cis} isomerization in neutral ABs. Rotation about the NN bond will be certainly favored by protonation on one of the N atoms because it decreases the double bond character.

The loss of planarity of ABH\textsuperscript{+} in S\textsubscript{1} could also be responsible for the absence of BD\textsuperscript{+} fragments (m/q = 105) in the photodissociation spectrum above \( \lambda = 410 \) nm. According to the fragmentation mechanism proposed in Scheme 3, the (ABH\textsuperscript{+} \rightarrow BD\textsuperscript{+} + B) reaction would be favored by an in-plane ring\textrightarrow{azo} electron transfer, in which one \( \pi \) electron of the ring migrates to the empty p orbital on the positively-charged N atom in order to extend the conjugation. In addition, the twisting process occurring in the S\textsubscript{1} state would convert very quickly the electronic energy into S\textsubscript{0} vibrational energy through the S\textsubscript{1}\textrightarrow{S\textsubscript{0}} conical intersection found in the calculations, and this will favor the release of m/q = 77 fragments like in the case of CID fragmentation in the ground state. The 41 \text{cm}^{-1} active vibrational progression observed in the ABH\textsuperscript{+} spectrum (inset of Fig. 1b) may also be associated with the twisting of the azo bond if one considers that the calculated NN torsional mode in the ground state is very close in energy (v = 35 \text{cm}^{-1}, in Scheme 2).

The next two quasi-degenerate bands in the spectrum of ABH\textsuperscript{+} exhibit sharper onsets than that of the first band. This observation is compatible with having sufficiently large Franck Condon factors in order to compensate the low oscillator strength values calculated for S\textsubscript{2}\textleftarrow{S\textsubscript{0}} and S\textsubscript{3}\textleftarrow{S\textsubscript{0}} excitations. As a result, it can be assumed that the geometry of S\textsubscript{2} and S\textsubscript{3} states will be akin to that of the S\textsubscript{0} state.

As shown in Fig. 3, the first band in the spectrum of dmaABH\textsuperscript{+} exhibits a different behavior than that of ABH\textsuperscript{+}. The onset is markedly steeper in the case of dmaABH\textsuperscript{+} evidencing a smaller distortion of the S\textsubscript{1} equilibrium geometry. One can tentatively assign this difference to the resonance donating effect of the terminal dimethylamino group, which contributes to
preserve a quasi-planar geometry in the excited state. The similar geometries of the ground and first excited states of dmaABH⁺ may also account for the production of \( m/q = 105 \) fragment ions.

As it was already pointed out, in the conditions of the trap, protonation of AB or dmaAB leads to the azonium form of the molecular ions. We have neglected the presence of the ammonium mesomer of dmaABH⁺ because of the large energy difference found in the \( S_0 \) states. Moreover, supplementary excited state calculations carried out for the ammonium tautomer of dmaABH⁺ have shown that the first transition having a non-negligible oscillator strength value appears at 3.79 eV (\( \lambda = 327 \) nm), and this does not match with the observed spectrum or in any case the ammonium band would be obscured by the \( S_2 \leftrightarrow S_0 \) transition of the azonium form.

The spectrum of cold BD⁺ given in Fig. 4c does not present well resolved vibrational structures as those observed for similar systems in comparable conditions.\(^{30,37}\) This is an indication that a substantial change in geometry occurs in the excited state or that its lifetime is very short. In order to study this possibility we have optimized \( S_1 \) and \( S_2 \) excited states of BD⁺ by releasing the constraint of maintaining the ground state Cs geometry at cc-pVDZ/CC2 level. The computed \( S_1 \) and \( S_2 \) geometries show that the diazonium group (\( -^iN\equiv N: \)) abandons the plane defined by the aromatic ring and bends. Two conical intersections associated with the crossings \( S_1=S_0 \) and \( S_2=S_1 \) emerge from the calculations at the geometries depicted in the right panel of Table 3, and no barrier was found in these optimization processes. A fast internal conversion through the two aforementioned conical intersections could account for the absence of a well-defined vibrational progression in the BD⁺ absorption bands. Accordingly, a fast deactivation of BD⁺ is expected to take place by relaxation processes linked to the torsion of the diazonium group.

5. Conclusions:

UV-VIS electronic spectra of ABH⁺, dmaABH⁺ and BD⁺ ions confined in a cryogenically-cooled trap were measured by photodissociation spectroscopy and the results were interpreted with the aid of ab initio calculations that include vertical excitation energies, normal modes in the \( S_0 \) state, and geometry optimizations in the ground and low lying excited states.
According to our ab initio computations, cisomers do not exist at the temperature of the trap, and the protonated forms of trans-AB and trans-dmaAB in the trap correspond to azonium structures, with no preference between isomers (I) and (II) in the case of dmaABH⁺. We have visualized that the S₁←S₀ transition of ABH⁺ and dmaABH⁺ mainly correspond to a LUMO←HOMO excitation with ππ←ππ⁺ character, the band of the substituted azobenzene being strongly red-shifted due to the donating effect of the dma group. At slightly higher energies, two intense CT transitions (S₂←S₀ and S₃←S₀) are clearly distinguished in the ABH⁺ spectrum. In contrast to S₁, S₂ and S₃ states preserve a quasi-planar structure in which an electron transfer takes place via the π orbitals system, along with the production of BD⁺ photofragments; this fragmentation channel is not active for S₁ and nor for S₄ excited states. Despite the size of the molecular ions involved in this study, we conclude that fragmentation is not exclusively due to internal conversion but some electronic properties of the excited states are revealed in the measured fragmentation patterns.

An efficient photoisomerization process connected to the torsion of the azo bond was shown in this study for ABH⁺ ions. It consists in a planar-to-chair dynamics that is related to the trans→cis rotation mechanism described in photoexcited neutral ABs. However, the yield of cis-isomers might be very low in the case of protonated ABs, since the chair-to-planar back reaction is expected to be very fast. Photoisomerization was found very sensitive to substitution by dma groups. The more pronounced rising of the first absorption band measured in the dmaABH⁺ electronic spectrum, and the good yield of BD⁺ fragments upon S₁ excitation, allowed us to speculate that the planar-to-chair dynamics is hindered in this case. A reason for that is the larger resonance delocalization induced by the dma terminal group, which may contribute to preserve a quasi-planar geometry in the S₁ excited state.

In addition, we are presenting the photofragmentation UV spectrum of BD⁺, a very stable and fundamental ion which results from photofragmentation of ABH⁺. Although the ion is cooled down in the trap, the observed spectrum is not well resolved, which is indicative of a strong change in the excited state geometry by torsion of the diazonium group or/and fast relaxation processes.

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6. Bibliography:


