Vibrational Analysis of Vacuum Ultraviolet Mass-Analyzed Threshold Ionization Spectra of Phenylacetylene and Benzonitrile

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One-photon MATI spectra of phenylacetylene and benzonitrile were obtained by using vacuum ultraviolet radiation generated by four-wave difference frequency mixing in Kr. Accurate ionization energies of phenylacetylene and benzonitrile were determined from the positions of the 0-0 bands, which are 8.8195 ± 0.0006 and 9.7288 ± 0.0006 eV, respectively. Vibrational frequencies of the molecular ions were determined from the spectra also. Assignments were made by utilizing the symmetry selection rule and frequencies and Franck—Condon factors calculated at the BP86, B3LYP, and B3PW91 density functional theory levels with the 6-311++G(2df,2pd) basis set.

I. Introduction

Structure, thermochemical properties, and dynamics of ions are of fundamental interest in relation to studies of combustion, atmospheric chemistry, cosmochemistry, etc.¹⁻³ Information on ionic vibrational structures is especially useful to probe ions in complex mixtures or follow complicated reaction processes. Nowadays, conventional spectroscopic techniques such as highresolution photoelectron or laser-induced fluorescence spectroscopies have been popular in characterizing polyatomic ions.^{4,5} However, obtaining vibrational spectra of polyatomic ions with these techniques is a formidable job because of their limited capabilities. The resolution of photoelectron spectroscopy (PES), which is typically 10 meV (80 cm⁻¹), is not good enough to obtain vibrational information on polyatomic cations, even though PES is useful to investigate electronic states.⁶ The laserinduced fluorescence spectroscopy, which usually has higher resolution than PES and can resolve vibrational peaks, is not generally applicable because most of the excited electronic states of polyatomic cations do not fluoresce.⁷

Zero kinetic energy (ZEKE) photoelectron spectroscopy has a much better resolution than ordinary PES and hence can obtain even rotational information for simple molecular ions.^{8–10} Massanalyzed threshold ionization (MATI) basically employs the same principle as ZEKE except for detecting ions instead of electrons and hence providing mass-selectivity in the spectra. 11–13 Generally adopted in the ZEKE or MATI spectroscopies is a two-color 1 + 1' scheme. Namely, excitation to a Rydberg state is achieved in two steps via an intermediate state. However, because the first excited states of most of the molecules are located in the region beyond commercial dye laser outputs (>210 nm) and these states in many cases are either unbound or relax rapidly, use of this scheme suffers the major limitation of low transition probability to the Rydberg state. One-photon ZEKE/MATI using tunable vacuum ultraviolet (VUV) radiation can overcome such difficulties because the transition occurs

directly from the ground state to a Rydberg state, not mediated by an excited electronic state of the neutral. 14-17

One-photon MATI spectra of benzene and halobenzenes reported recently by this laboratory showed well-resolved vibrational peaks of the corresponding cations, which consisted mostly of fundamentals with proper symmetries. 18-21 Vibrational assignments were made by referring to the previous results, comparing with calculated frequencies, and invoking the selection rule for one-photon process. Difference in the geometry between the neutral and cation, or geometric change upon ionization, was calculated quantum chemically and used to explain the prominent overtones of some vibrational modes and combinations involving these. As a more rigorous attempt to utilize spectral intensity information for vibrational assignment, Franck-Condon factors were calculated from the quantum chemical results in our recent study of some aliphatic halides. ^{22,23} Theoretical prediction of the intensities for transitions to all of the totally symmetric vibrational states was found to be extremely useful for reliable assignment.

In this paper, we report the vibrational spectra of pheny-lacetylene and benzonitrile cations obtained by one-photon MATI spectroscopy. Successful vibrational assignments made on the basis of the above strategy, namely, by utilizing the calculated frequencies and Franck—Condon factors and symmetry selection rule, will be presented also.

II. Experimental Section

Phenylacetylene and benzonitrile were purchased from Aldrich and Kanto, respectively, and used without further purification. A gaseous sample seeded in He at the stagnation pressure of \sim 2 atm was supersonically expanded through a temperature-controlled pulsed nozzle (diam 500 μ m, General Valve) and introduced to an ionization chamber through a skimmer (diam 2 mm, Beam Dynamics) placed about 3 cm downstream from the nozzle orifice. The background pressure in the ionization chamber was typically \sim 10⁻⁸ Torr.

The experimental setup was described in detail previously. 18,19 Briefly, to excite the Kr $5p[^1/_2]_0-4p^6$ transition for VUV generation by four wave difference frequency mixing, the light at 212.5 nm (\sim 0.5 mJ/pulse) was generated by frequency tripling

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TABLE 1: Vibrational Frequencies (in cm $^{-1}$) of Phenylacetylene Neutral and Cation in the Ground Electronic States Calculated at the B3LYP, B3PW91, and BP86 Levels with the 6-311++G(2df,2pd) Basis Set and Experimental Data for the Neutral

			neutral			cation			
mode	symm	${\sf expt}^a$	B3LYP	B3PW91	BP86	B3LYP	B3PW91	BP86	
1	a_1	763	777	781	756	766	767	746	
2	a_1	3064	3202	3210	3125	3221	3228	3144	
3	b_2	1283	1310	1325	1290	1298	1304	1257	
4	b_1	691	708	707	683	741	738	699	
5	b_1	986	1014	1011	968	1045	1040	999	
6a	a_1	467	474	471	459	470	467	455	
6b	b_2	619	638	648	616	581	578	560	
7a	a_1	3035	3172	3179	3094	3196	3201	3119	
7b	b_2		3198	3206	3121	3219	3226	3142	
8a	a_1	1598	1640	1651	1587	1643	1652	1592	
8b	b_2	1573	1609	1621	1558	1536	1538	1483	
9a	a_1	1178	1202	1198	1163	1210	1206	1172	
9b	b_2	1158	1185	1180	1149	1175	1174	1139	
10a	a_2	842	860	859	825	827	824	789	
10b	b_1	165	142	142	136	120	118	115	
11	b_1	756	782	777	752	817	815	787	
12	a_1	1000	1018	1017	987	998	999	971	
13	a_1	1192	1223	1228	1193	1266	1271	1228	
14	b_2	1331	1358	1357	1325	1387	1393	1348	
15	b_2		161	159	152	153	151	145	
16a	a_2	418	413	409	395	363	358	345	
16b	b_1	531	558	558	536	481	479	457	
17a	a_2	971	998	996	953	1027	1024	982	
17b	b_1	918	946	944	904	991	988	949	
18a	a_1	1028	1050	1053	1020	1009	1011	981	
18b	b_2	1071	1102	1103	1069	1113	1115	1081	
19a	a_1	1489	1526	1525	1474	1486	1483	1435	
19b	b_2	1444	1478	1477	1429	1436	1442	1399	
20a	a_1	3083	3191	3198	3113	3210	3216	3133	
20b	b_2	3058	3180	3188	3103	3208	3214	3130	
β CC	b_2	516	539	538	517	531	528	507	
β CH	b_2	653	689	690	635	694	695	645	
ν CC	a_1	2118	2202	2209	2131	2107	2108	2053	
ν CH	a_1	3291	3468	3472	3393	3409	3412	3334	
γ CC	b_1	352	371	369	354	324	322	311	
γСН	b_1	610	645	633	588	645	642	621	

^a Reference 33.

of 637.6 nm output of a dye laser (Continuum ND6000) pumped by the second harmonic of an Nd:YAG laser (Continuum PL8000). Another dye laser output (430-760 nm) pumped by the second or third harmonic of the second Nd:YAG laser (Continuum Surelite II) was combined with the 212.5 nm light and loosely focused with a fused silica lens (f = 50 cm) in the Kr cell to generate the VUV light tunable in the 123-141 nm region. A MgF₂ lens (f = 25 cm) was placed at the exit of the Kr cell, and the laser beams were aligned off-centered at the lens to separate the residual light beams (UV and visible) from the VUV light, which was focused onto the molecular beam. The optimized Kr pressure in the cell was 5-18 Torr. Small portions of dye laser outputs were used to calibrate their frequencies on the basis of the optogalvanic effect in an Fe/Ne hollow cathode lamp.²⁴ Its precision was ± 0.5 cm⁻¹ in the visible region.

The VUV laser pulse was collinearly overlapped with the molecular beam in a counter-propagation manner and slit-electrodes along the beam path were used to maximize ion collection efficiency. The spoil field of 0–0.2 V/cm was applied in the ionization region to remove directly produced ions. To achieve pulsed-field ionization (PFI) of neutrals in the ZEKE state, an electric field of 15–60 V/cm was applied at a certain delay time after the laser pulse. Ions were then accelerated, flew through a field-free region, and were detected by a dual microchannel plate detector. A short pulse of the scrambling

TABLE 2: Vibrational Frequencies (in cm $^{-1}$) of Benzonitrile Neutral and Cation in the Ground Electronic States Calculated at the B3LYP, B3PW91, and BP86 Levels with the 6-311++G(2df,2pd) Basis Set and Experimental Data for the Neutral

			ne	utral	cation			
mode	symm	expt ^a	B3LYP	B3PW91	BP86	B3LYP	B3PW91	BP86
1	a_1	769	774	774	752	755	756	736
2	a_1	3071	3196	3215	3130	3210	3228	3143
3	b_2	1289	1319	1334	1298	1416	1426	1382
4	b_1	686	706	704	682	631	628	607
5	b_1	987	1021	1016	974	1042	1035	994
6a	a_1	461	467	465	452	459	456	445
6b	b_2	629	641	637	619	504	500	483
7a	a_1	3042	3178	3185	3100	3196	3201	3118
7b	b_2	3027	3204	3212	3127	3219	3225	3141
8a	a_1	1599	1641	1654	1588	1660	1669	1603
8b	b_2	1584	1615	1627	1563	1275	1279	1235
9a	a_1	1178	1203	1200	1165	1210	1207	1172
9b	b_2	1163	1188	1184	1153	1148	1150	1110
10a	a_2	848	863	862	828	814	811	775
10b	b_1	172	147	146	140	118	116	111
11	b_1	758	781	780	751	815	813	784
12	a_1	1001	1019	1019	989	1001	1000	971
13	a_1	1191	1220	1226	1188	1252	1256	1215
14	b_2	1337	1361	1361	1329	1392	1389	1344
15	b_2	162	169	166	160	157	155	148
16a	a_2	401	410	406	392	353	347	335
16b	b_1	548	573	574	552	449	442	413
17a	a_2	978	1002	1000	957	1029	1025	983
17b	b_1	925	954	952	912	990	985	947
18a	a_1	1027	1050	1053	1021	984	988	962
18b	b_2	1071	1105	1106	1073	1088	1093	1054
19a	a_1	1492	1528	1527	1475	1474	1469	1425
19b	b_2	1448	1481	1480	1431	1528	1529	1475
20a	a_1	3080	3207	3203	3118	3221	3216	3132
20b	b_2	3039	3188	3195	3110	3209	3214	3131
β CN	b_2	551	570	570	549	568	570	548
ν CN	a_1	2232	2332	2341	2236	2192	2196	2124
γ CN	b_1	381	392	390	375	317	312	298

^a Reference 33.

field was applied at the laser irradiation time, which significantly lengthened the lifetime of the ZEKE states. Use of a long delay time (\sim 20 μ s) and low spoil field tremendously improved the quality of the MATI spectra. A gold wire was placed in the VUV beam path as a beam monitor.²⁵ Its output was used to normalize the intensity of each vibrational peak in the MATI spectra.

III. Vibrational Analysis

A. Quantum Chemical Calculations. Calculations of equilibrium geometries and vibrational frequencies of phenylacetylene and benzonitrile and their cations in the ground states were performed at the density functional theory (DFT) levels, B3LYP, B3PW91, and BP86, with various basis sets using Gaussian 98 suite of programs. Size of the basis set was systematically increased until the basis set dependence disappeared. Hence, only those obtained with the largest basis set used, the 6-311++G(2df,2pd), will be listed and discussed. For the vibrational frequencies, the results obtained at the BP86/6-311++G(2df,2pd) level showed the best agreement with the experimental data. Frequencies obtained by these calculations are presented without scaling in Tables 1 and 2.

B. Symmetry Selection Rules. In our previous investigation of one-photon MATI spectra of benzene, ²⁰ symmetry selection rules for transitions from the ground-state neutral to Rydberg states were extremely helpful for the vibrational peak assignments. The selection rules pertinent to the present systems are summarized as follows.

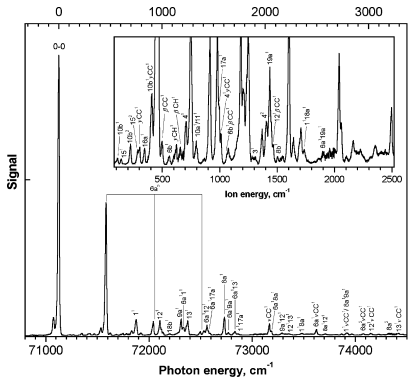


Figure 1. One-photon MATI spectrum of $C_6H_5C \equiv CH$ recorded by monitoring $C_6H_5C \equiv CH^+$ in the ground electronic state. The x-scale at the top of the figure corresponds to the vibrational frequency scale for the cation the origin of which is at the 0-0 band position. Spectrum in the 50-2500 cm⁻¹ region magnified by 15 is shown as an inset to demonstrate the quality of the MATI spectrum obtained in this work.

Under the Born—Oppenheimer approximation, the transition moment for Rydberg (R) ← ground (G) vibronic transition can be expressed as the product of the electronic (e) and vibrational (v) parts. Upon further approximation of the electronic wave function of a Rydberg state as the product of the electronic wave function of the ion core (Ψ_e^C) and that of the Rydberg electron (Ψ_e^R) , these become $\langle \Psi_e^G|\bar{\mu}|\Psi_e^C\Psi_e^R\rangle$ and $\langle \Psi_v^G|\Psi_v^C\rangle$, respectively. For a high Rydberg state, Ψ_e^C and Ψ_v^C can be approximated as those of the corresponding ionic state. Detailed analysis shows that one can always choose a Rydberg orbital that makes the electronic part allowed. Most of the molecules prepared under the supersonic jet condition are in the zero-point vibrational level, which is totally symmetric, a₁. Hence, in an electric dipoleallowed transition, Ψ_{ν}^{C} should also belong to a_{1} . Then, fundamentals and all of the overtones of a₁ modes are allowed, while the $\Delta v = 2, 4, 6, \dots$ selection rule holds for nontotally symmetric modes. Also, all of the transitions to the combination states with a₁ symmetry are allowed. Then, the relative intensity of a transition to an a₁ vibrational state, either fundamental, overtone, or combination, is determined by the Franck-Condon factor. 27,28 The methods to evaluate the Franck-Condon factors from quantum chemical results are well established.²²

Vibrational Analysis of VUV-MATI Spectra

IV. Results and Discussion

MATI spectra of phenylacetylene and benzonitrile recorded by monitoring $C_6H_5C \equiv CH^+$ and $C_6H_5C \equiv N^+$ in the ground electronic states are shown in Figures 1 and 2, respectively. The spectra magnified by 15 are also shown as insets in the figures. The intense peaks appearing at the lowest photon energy, namely, at 71 127 and $78 461 \text{ cm}^{-1}$ in Figures 1 and 2, respectively, correspond to the 0-0 bands. The position of the 0-0 band in a one-photon MATI spectrum is equivalent to the ionization energy of the molecule. However, the ionization energy thus measured is usually a little smaller than the correct value because the molecules in ZEKE states¹⁰ some cm⁻¹ below the threshold can also be ionized when a high PFI field is applied. To correct for this effect, the 0-0 band position was measured using various PFI fields, and the accurate ionization energy was estimated by extrapolation to the zero-field limit. Spoil field was not used in such measurements. The ionization energies to the ground electronic states of phenylacetylene and benzonitrile cations measured in this work are listed in Table 3, together with previous results.²⁹⁻³² There has been no previous report on the accurate ionization energy of phenylacetylene measured by ZEKE or MATI. The ionization energy of phenylacetylene determined in this work, 8.8195 ± 0.0006 eV, is a little different from 8.825 ± 0.001 eV measured by threshold photoelectron spectroscopy (TPES).²⁹ More annoying is that the ionization energy of benzonitrile obtained in this work, 9.7288 ± 0.0006 eV, is smaller than a previous ZEKE result³¹ by ~ 20 cm⁻¹. Our experience is that the present MATI technique, which uses high-voltage electronics, tends to underestimate ionization energies by 0-5 cm⁻¹. ^{19,20} We do not have an explanation for the above discrepancy at the moment even though we would like to point out that quality of the present MATI spectrum is superior to the previous ZEKE spectrum.³¹

Assuming that the shift of a vibrational peak in a MATI spectrum due to the applied electric fields is similar to that of the 0-0 band, the vibrational frequency corresponding to each peak can be determined simply by taking the difference of its position from that of the 0-0 band. Vibrational frequency scales with origins at the 0-0 band positions are also drawn in Figures 1 and 2. Vibrational frequencies of $C_6H_5C \equiv CH^+$ and C₆H₅C≡N⁺ in the ground electronic state calculated at the BP86/6-311++G(2df,2pd) level are compared with the experimental data in Tables 4 and 5, respectively. Also listed in the tables are intensities of each peak in the MATI spectra normalized to that of the 0-0 band. Frequencies of some vibrations measured previously by TPES²⁹ and ZEKE³¹ are also included in the tables.

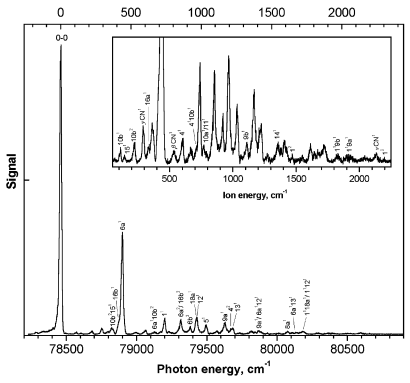


Figure 2. One-photon MATI spectrum of $C_6H_5C\equiv N$ recorded by monitoring $C_6H_5C\equiv N^+$ in the ground electronic state. The *x*-scale at the top of the figure corresponds to the vibrational frequency scale for the cation the origin of which is at the 0–0 band position. Spectrum in the 50–2200 cm⁻¹ region magnified by 15 is shown as an inset to demonstrate the quality of the MATI spectrum obtained in this work.

TABLE 3: Ionization Energies (IE) of Phenylacetylene and Benzonitrile in eV

	IE (X)	ref
phenylacetylene	8.8195 ± 0.0006 8.825 ± 0.001 8.82 ± 0.02	this work TPES ²⁹ PES ³⁰
benzonitrile	9.7288 ± 0.0006 9.7315 ± 0.0002 9.71 ± 0.01	this work ZEKE ³¹ PI ³²

At all of the levels used in the calculation, equilibrium geometries of the phenylacetylene and benzonitrile neutrals and cations belong to the C_{2v} symmetry. If the symmetry selection rule is invoked, prominent peaks in the MATI spectra must be mostly due to transitions to the a1 vibrational states. The Franck-Condon factors for such transitions, either fundamentals, overtones, or combinations, calculated at three density functional theory (DFT) levels, B3LYP, B3PW91, and BP86, using the 6-311++G(2df,2pd) basis set were rather similar. The Franck-Condon factors calculated at the BP86/ 6-311++G(2df,2pd) level and normalized to that of the 0-0transition are listed in Tables 4 and 5 also. Comparing the experimental and calculated frequencies and Franck-Condon factors was helpful to identify a1 peaks. The remaining weak peaks in one-photon MATI spectra must be due to electric dipole-forbidden but vibronically allowed transitions. Only the calculated frequencies, not intensities, can be used to assign these non-totally symmetric transitions.

A. Phenylacetylene. The phenylacetylene cation with $C_{2\nu}$ symmetry has 36 nondegenerate normal modes,³³ 13 of which belong to the a_1 symmetry species, 3 to a_2 , 8 to b_1 , and 12 to b_2 . Among the a_1 -type modes, 2, 7a, 20a, and ν CH in Wilson notation³⁴ are due to CH stretching and have frequencies of ~ 3000 cm⁻¹. According to our previous study on the CH stretching modes,²² they are not expected to appear distinctly in the one-photon MATI spectrum. Then, prominent peaks in

the spectral region $0-1700~{\rm cm}^{-1}$ and near $\sim 2000~{\rm cm}^{-1}$ can mostly be assigned to the fundamentals of the remaining a₁ modes, 1, 6a, 8a, 9a, 12, 13, 18a, 19a, and ν CC, as well as their overtones and combinations. Among these, the calculated Franck-Condon factors are especially significant for the fundamentals of 6a, 1, 12, 9a, 8a, and vCC of which the calculated frequencies at the BP86 level are 455, 746, 971, 1172, 1592, and 2053 cm⁻¹, respectively. Hence, the prominent peaks at 458, 747, 979, 1185, 1604, and 2040 cm⁻¹ in the MATI spectrum of phenylacetylene can readily be assigned to 6a¹, 1¹, 12^1 , $9a^1$, $8a^1$, and νCC^1 , respectively. Even though the fundamental of v_{13} , which is another a_1 -type mode, is expected to be weak according to its calculated Franck-Condon factor, it appears distinctly near its calculated frequency at 1249 cm⁻¹. 19a1 appears rather distinctly at 1435 cm⁻¹, even though its calculated Franck-Condon is only 0.004. We would rather assign this to a composite of 19a1 and 6a1121 on the basis of the calculated frequencies and Franck-Condon factors. A weak shoulder peak at 989 cm⁻¹ is close to 981 cm⁻¹ calculated for 18a¹. We are reluctant to make such an assignment because the calculated Franck-Condon factor for this transition is extremely small, 0.0002. It is to be mentioned that the harmonic frequencies calculated at the BP86 level are usually a little smaller than the experimental ones, while those at the B3LYP and B3PW91 levels are larger. The same trend holds for most of the fundamentals of the phenylacetylene and benzonitrile neutrals as can be seen in Tables 1 and 2. Even though the results must rise from different error cancellations at these levels,35 the correlation can be used advantageously in the peak assignments.

The fact that 6a¹ appeared most prominently in the one-photon MATI spectrum indicates that the geometrical change upon ionization occurs mostly along the 6a eigenvector. This was confirmed by calculation, even though not shown here, by projecting the geometrical change vector on the 6a eigenvector. This also suggests that 6aⁿ overtones and combinations of 6a

TABLE 4: Vibrational Frequencies (in cm⁻¹) and Their Assignments for Phenylacetylene Cation in the Ground Electronic State (\tilde{X}^2B_1)

			cal	culated ^c	thi	s work				cal	culated ^c	thi	s work
$mode^a$	symm	$TPES^b$	freq	$intensity^d$	freq	intensity ^d	$mode^a$	symm	$TPES^b$	freq	intensity ^d	freq	intensity ^d
						Fundame	entals						
1	a_1	759	746	0.131	747	0.076	14	b_2		1348	0		
2	a_1		3144	5×10^{-6}			15	b_2		145	0	143	0.005
3	b_2		1257	0	1287	0.007	16a	a_2		345	0	346	0.011
4	b_1		699	0	706	0.017	16b	b_1		457	0		
5	b_1		999	0	996	0.008	17a	a_2		982	0	989	0.006
6a	a_1	460	455	0.353	458	0.438	17b	b_1		949	0		
6b	b_2		560	0	561	0.008	18a	a_1		981	0.0002		
7a	a_1		3119	3×10^{-6}			18b	b_2		1081	0	1076	0.007
7b	b_2		3142	0			19a	a_1		1435	0.004	1435^{f}	0.044
8a	a_1		1592	0.096	1604	0.073	19b	b_2		1399	0		
8b	b_2		1483	0	1505	0.006	20a	a_1		3133	0.0003		
9a	a_1		1172	0.081	1185	0.102	20b	b_2		3130	0		
9b	b_2		1139	0	1158	0.018	β CC	b_2	504	507	0	499	0.012
10a	a_2		789	0	795^{e}	0.015	β CH	b_2		645	0	658	0.008
10b	b_1		115	0	110	0.008	ν CC	a_1		2053	0.101	2040	0.042
11	b_1		787	0	795^{e}	0.015	ν CH	a_1		3334	0.0001		
12	a_1		971	0.069	979	0.066	γCC	b_1		311	0	303	0.010
13	a_1		1228	0.010	1249	0.087	γСН	b_1		621	0	622	0.010
					O.	vertones and C	Combinations						
$10b^{2}$	a_1		230	0.002	221	0.010	$6a^{1}19a^{1}$	a_1		1890	0.002	1895	0.017
15^{2}	a_1		290	0.0002	286	0.007	$6a^{1}8a^{1}$	a_1		2047	0.035	2058	0.017
$10b^1\gamma CC^1$	a_1		426	0.002	409	0.034	$9a^{1}12^{1}$	a_1		2143	0.005	2164	0.017
6a ¹ 10b ¹ γCC ¹	a_1		881	0.0007	865	0.006	$12^{1}13^{1}$	a_1		2199	0.001	2228	0.013
6a ²	a_1		910	0.065	914	0.063	$1^{1}8a^{1}$	a_1		2338	0.011	2356	0.016
$4^{1}\gamma CC^{1}$	a_1		1010	0.0003	1009	0.012	$6a^1\nu CC^1$	a_1		2508	0.038	2496	0.027
6b ¹ βCC ¹	a_1		1067	0.0004	1064	0.005	$8a^{1}12^{1}$	a_1		2563	0.005	2582	0.012
6a ¹ 1 ¹	a_1		1201	0.045	1205	0.082	$1^1\nu CC^1$	a_1		2764	0.008	2700	0.016
6a ³	a_1		1365	0.008	1370	0.022	$8a^{1}9a^{1}$	a_1		2799	0.012	2789	0.016
42	a_1		1398	0.008	1407	0.028	19a¹19b¹	b_2		2834	0	2837	0.007
6a ¹ 12 ¹	a_1		1426	0.026	1435^{f}	0.044	$6a^2\nu CC^1$	a_1		2963	0.007	2953	0.011
6a ¹ 17a ¹	a_2		1437	0	1448	0.009	$12^1\nu\text{CC}^1$	a_1		3024	0.008	3022	0.010
$12^{1}\beta CC^{1}$	b_2		1478	0	1465	0.007	$6a^{1}8a^{1}12^{1}$	a_1		3018	0.002	3032	0.008
6a ¹ 9a ¹	a_1		1627	0.029	1643	0.017	$8a^2$	a_1		3184	0.006	3206	0.014
6a ¹ 13 ¹	a_1		1683	0.004	1706	0.025	$9a^1\nu CC^1$	a_1		3224	0.010	3227	0.011
1 ¹ 17a ¹	\mathbf{a}_2		1728	0	1739	0.010	$13^{1}\nu CC^{1}$	a_1		3281	0.002	3292	0.010

^a Wilson notation. ^b Reference 29. ^c BP86/6-311++G(2df,2pd) level. ^d Normalized to the intensity of the 0-0 band. ^e A composite of 10a¹ and 11¹. ^f A composite of 19a¹ and 6a¹12¹.

and other a₁ modes would appear prominently in the one-photon MATI spectrum. Accordingly, the prominent peaks at 914 and 1370 cm⁻¹ can be assigned to 6a² and 6a³, respectively. Also, the distinct peaks at 1205, 1435, 1643, 1706, 2058, and 2496 can be assigned to 6a combinations, $6a^11^1$, $6a^112^1$, $6a^19a^1$, $6a^{1}13^{1}$, $6a^{1}8a^{1}$, and $6a^{1}\nu CC^{1}$, respectively, from the calculated frequencies and Franck-Condon factors. The peak at 1435 cm⁻¹ can be alternatively assigned to 19a1 as has been mentioned earlier. Other combinations of the a₁ modes, 1¹18a¹, 9a¹12¹, and 1¹8a¹, appeared distinctly also at 1739, 2164, and 2356 cm⁻¹, respectively. The Franck-Condon factors for $1^1 \nu CC^1$ and $8a^1$ -9a1 calculated at the BP86 level are 0.012 and 0.008, respectively. Hence, the peak at 2789 cm⁻¹, which can be assigned either to $1^{1}\nu$ CC¹ or to $8a^{1}9a^{1}$ on the basis of the frequencies may better be assigned to a composite of the two transitions, 1¹νCC¹/8a¹9a¹. The calculated Franck—Condon factors for the fundamentals of the CH stretching modes 2, 7a, 20a, and ν CH are negligible. This is understandable because the lengths of all of the CH bonds hardly change upon ionization. Extremely weak peaks appeared in the 3000-3200 cm⁻¹ region of the present MATI spectrum (Figure 1). We are reluctant to assign them to 2¹, 7a¹, 20a¹, or ν CH¹ because their calculated Franck— Condon factors are very small. Instead, it is more likely that they are overtones or combinations. Thus, the very weak peaks at 3022, 3206, and 3292 were assigned to $12^{1}\nu CC^{1}$, $8a^{2}$, and $13^{1}\nu$ CC¹, respectively. Some distinct peaks may be assigned to

combinations of nontotally symmetric modes with a₁ overall symmetry. Hence, the distinct peaks at 221, 286, 409, 1009, and 1407 were assigned to $10b^2$, 15^2 , $10b^1\gamma CC^1$, $4^1\gamma CC^1$, and 4², respectively.

The fundamentals of nontotally symmetric modes are forbidden under the Born-Oppenheimer approximation. They still appeared in the actual spectrum, even though very weakly, through vibronic mechanism. Hence, very weak peaks at 303, 346, 499, 561, 622, 706, and 795 cm⁻¹ can be assigned to γ CC¹, $16a^1$, βCC^1 , $6b^1$, γCH^1 , 4^1 , and $10a^1/11^1$ by comparing with the calculated frequencies of 311, 345, 507, 560, 621, 699, and 789/787 cm⁻¹, respectively. A very weak 16b¹ is expected at \sim 457 cm⁻¹ but must have been buried in the strong 6a¹ transition. Similarly, 17a¹ and 17b¹ expected at 950–980 cm⁻¹ may have been buried as shoulders of 12¹. The weak shoulder peak at 989 cm⁻¹ mentioned previously may be 17a¹ rather than 18a¹.

B. Benzonitrile. Kimura and co-workers reported the 1 + 1'ZEKE spectrum of benzonitrile in the $0-1200 \text{ cm}^{-1}$ vibrational energy region.³¹ Tentative assignments were made by comparing with results from PM3 semiempirical calculation. They adopted Mulliken notation for vibration modes, with some minor errors. We reinterpreted their assignments using the DFT frequencies calculated in this work and listed them in Table 5.

A spectrum with much better quality than the above was obtained in the $0-2500 \text{ cm}^{-1}$ region (78 200-81 000 cm⁻¹ in

TABLE 5: Vibrational Frequencies (in cm $^{-1}$) and Their Assignments for Benzonitrile Cation in the Ground Electronic State (\tilde{X}^2B_1)

			cal	culated ^c	this work		
$mode^a$	symm	$ZEKE^b$	freq	intensity ^d	freq	intensity ^d	
		F	undame	ntals			
1	a_1		736	0.213	737	0.098	
2	a_1		3143	0.0001			
3	b_2		1382	0	1386	0.011	
4	b_1		607	0	606	0.022	
5	b_1	1036	994	0	1034	0.047	
6a	a_1	447	445	0.295	438	0.376	
6b	b_2	559	483	0			
7a	a_1		3118	2×10^{-7}			
7b	b_2		3141	0			
8a	a_1		1603	0.143	1612	0.032	
8b	b_2		1235	0			
9a	a_1		1172	0.098	1168	0.072	
9b	b_2		1110	0	1111	0.025	
10a	a_2	854	775	0	771^{e}	0.017	
10b	b_1	110	111	0	111	0.016	
11	b_1		784	0	771^{e}	0.019	
12	a_1		971	0.036	968	0.039	
13	a_1		1215	0.001	1223	0.024	
14	b_2		1344	0	1353	0.020	
15	b_2	144	148	0	143	0.011	
16a	\mathbf{a}_2	332	335	0	336	0.011	
16b	b_1	405	413	0	412	0.022	
17a	a_2	1005	983	0	984	0.011	
17b	b_1	920	947	0	922^f	0.025	
18a	a_1		962	0.081	959	0.047	
18b	b_2		1054	0	4.440:	0.01=	
19a	a_1		1425	0.001	1413g	0.017	
19b	b_2		1475	0	1470^{h}	0.011	
20a	a_1		3132	1×10^{-5}			
20b	b_2	507	3131	0	526	0.015	
βCN	b_2	537	548 2124	0	536 2136	0.015	
νCN	a ₁		2124	0.055	2136	0.037 0.018	
γCN	b_1			0	291	0.018	
				ombinations			
$10b^{2}$	a_1	219	222	0.005	223	0.015	
$10b^215^1$	b_2		370	0	361	0.037	
6a ¹ 10b ²	a_1		667	0.002	662	0.011	
$4^{1}10b^{1}$	a_1		718	0.001	712	0.022	
$16b^{2}$	a_1		826	0.017	854	0.091	
$6a^2$	a_1		890	0.045			
6b ²	a_1		966	0.009	922^{f}	0.023	
42	a_1		1214	0.0004	1212	0.015	
6a ¹ 12 ¹	a_1		1416	0.012	1413g	0.017	
12	a_1		1472	0.020	1470^{h}	0.011	
6a ¹ 13 ¹	a_1		1660	0.0003	1647	0.008	
1 ¹ 18a ¹	a_1		1698	0.017	1724	0.034	
1 ¹ 12 ¹	a ₁		1707	0.006	1041	0.011	
1¹9b¹ 1¹9a¹	b_2		1846	0	1841	0.011	
1 '9a' 13	a_1		1908 2208	0.018 0.001	1903 2205	0.013 0.036	
1	a_1		2200	0.001	2203	0.030	

 a Wilson notation. b Reference 31. c BP86/6-311++G(2df,2pd) level. d Normalized to the intensity of the 0–0 band. Intensity of the peaks above 1700 cm $^{-1}$ is not accurate due to very weak VUV power. c A composite of 10a 1 and 11 1 . f A composite of 17b 1 and 6b 2 . g A composite of 19a 1 and 6a 1 12 1 . h A composite of 19b 1 and 1 2 .

photon energy) by one-photon MATI in this work. The 81 000—81 500 cm⁻¹ region could not be recorded because of a dip in VUV output. According to our experience in MATI of phenylacetylene in this work and halobenzenes in previous studies, ^{18,19} this region is not expected to be important because the CH stretching fundamentals would not be observed anyway. Symmetry selection rule and frequencies and Franck—Condon factors calculated at the DFT levels will be utilized to assign the observed vibrational peaks as has been done for phenylacetylene. VUV power above 1700 cm⁻¹ in vibrational energy was very

weak. Hence, the calibrated relative intensities of peaks above $1700~\rm cm^{-1}$ may be rather inaccurate.

Quantum chemical calculations performed at various levels for this molecule suggest that the cation belongs to the $C_{2\nu}$ point group in the ground electronic state. Twelve normal modes are a₁-type as listed in Table 2. Among these, 2, 7a, and 20a are high-frequency vibrations with CH stretching character and can be neglected in this work. Among the a₁-type vibrations, the fundamental of 6a displays the largest Franck-Condon factor and is readily identified as the peak at 438 cm⁻¹. Franck— Condon factors are significant for 1¹, 18a¹, 12¹, 9a¹, 8a¹, and ν CN¹ with the calculated frequencies of 736, 962, 971, 1172, 1603, and 2124 cm⁻¹, respectively. The prominent peaks at 737, 959, 968, 1168, 1612, and 2136 cm⁻¹ in the MATI spectrum correspond to these transitions. The remaining a₁-type fundamentals are 13¹ and 19a¹ with the calculated frequencies of 1215 and 1425 cm⁻¹, respectively, and the calculated Franck-Condon factors of only 0.001 for each. Even though one might assign the peak at 1413 cm⁻¹ to 19a¹ on the basis of the frequency, 6a1121 seems to be a better assignment when the intensity is considered also. For 131, the peak at 1223 cm⁻¹ is the only candidate even though the calculated and observed intensities show a substantial discrepancy. For the peak at 1223 cm⁻¹, no alternative assignment is possible, to either a fundamental, overtone, or combination. Namely, its assignment to 131 is unavoidable, and its small Franck-Condon factor must be attributed to inaccuracy in the calculation. It is to be noted from Table 4 that the Franck—Condon factor for the same transition in phenylacetylene is also much smaller than the experimental intensity, even though not as dramatically as in benzonitrile. It is to be mentioned that Kimura and co-workers did not identify the a_1 -type fundamentals, except for $6a^1$, in their 1 + 1' ZEKE work.31

Strong fundamental of the 6a mode suggests that its overtones and combinations, especially with other prominent a₁-type modes such as 1, 8a, 9a, 12, 13, 18a, and ν CN, would appear distinctly in the one-photon MATI spectrum. 6a² and 6a³ overtones are expected to appear at somewhat lower than 876 and 1314 cm⁻¹. Their calculated Franck-Condon factors are 0.045 and 0.005, respectively. They are not easy to identify, however, because of the presence of strong spectral features nearby. The strong peak at 854 cm⁻¹ cannot be matched with the calculated frequency of any fundamental. It may be a composite consisting of 6a² and 16b². We do not attempt to identify 6a³ because of its small Franck-Condon factor. The a₁-type combinations involving 6a¹, namely, 6a¹12¹ and 6a¹-13¹, are identified at 1413 and 1647 cm⁻¹, respectively. The mode 1, which has substantial fundamental intensity, also shows an overtone 1² at 1470 cm⁻¹. The a₁-type overtones and combinations involving non-totally symmetric modes are also observed even though with weaker intensities than the a₁ fundamentals. These are 10b², 6a¹10b², 4¹10b¹, and 4² at 223, 662, 712, and 1212 cm⁻¹, respectively.

Fundamentals of some non-totally symmetric modes also appear probably through vibronic mechanism. The most noticeable among these are the fundamental, overtone, and combinations of the 10b mode, namely, $10b^1$, $10b^2$, $10b^215^1$, $6a^110b^2$, and 4^110b^1 . $10b^1$ appeared very prominently in the 1 + 1' ZEKE spectrum reported by Kimura and co-workers. It is not as prominent in the present one-photon MATI spectrum. The fact that the initial electronic states involved in the transition moment integral are different in the two processes must be responsible for the above difference. The remaining distinct feature in the MATI spectrum is the peak at 1034 cm^{-1} . Comparing with the

frequencies calculated at the three DFT levels, it seems to be logical to assign this to 5^1 , in agreement with the assignment by Kimura and co-workers. Even though $6b^1\beta CN^1$ may be an alternative based on the frequency alone, the very small Franck—Condon factor calculated for the latter, 0.001, is incompatible with the observation.

V. Summary and Conclusion

For a molecule with a large number of vibrational degrees of freedom such as $C_6H_5C\equiv CH^+$ and $C_6H_5C\equiv N^+$, assigning its vibrational spectrum can be a formidable job, especially when no additional information is available in the literature. In the case of two-photon ZEKE/MATI, further information can often be obtained through intermediate state selection. This is not the case in the one-photon scheme, even though the fact that excitation via an intermediate is not needed is its clear advantage in experimental terms. In our previous one-photon MATI investigation of C₂H₃Br⁺,²² it was demonstrated that use of the selection rule and calculated frequencies and Franck-Condon factors, especially those at the DFT levels, led to nearly complete assignment for its vibrational spectrum. The same strategy has been used with a striking success for the vibrational assignments for the cations of phenylacetylene and benzonitrile, which are much larger than C₂H₃Br⁺. It is to be emphasized that the frequencies obtained at the DFT levels, especially BP86, provided nearly quantitative fit to the experimental data even though harmonic approximations were adopted for all of the vibrations. Cancellation of various errors must have acted favorably to result in such good fits. It is to be mentioned, however, that agreement between the calculated and measured frequencies is not sufficient for making a definite assignment. Comparing calculated and measured intensities would be helpful in this regard.

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References and Notes

- (1) Glassman, I. Combustion; Academic Press: San Diego, CA, 1996.
- (2) Brasseur, G. P.; Orlando, J. J.; Tyndall, G. S. Atmospheric Chemistry and Global Change; Oxford University Press: New York, 1999.
- (3) Cowley, C. R. An Introduction to Cosmochemistry; Cambridge University Press: Cambridge, U.K., 1995.
- (4) Kimura, K.; Katsumata, S.; Achiba, Y.; Yamazaki, T.; Iwata, S. Handbook of Hel Photoelectron Spectra of Fundamental Organic Molecules; Japan Scientific Societies Press: Tokyo, 1981.
- (5) Miller, T. A.; Bondybey, V. E. *Molecular Ions: Spectroscopy, Structure, and Chemistry*; North-Holland Publishing Company: New York, 1983

- (6) Baltzer, P.; Karlsson, L.; Wannberg, B.; Öhrwall, G.; Holland, D. M. P.; MacDonald, M. A.; Hayes, M. A.; von Niessen, W. Chem. Phys. 1997, 224, 95.
- (7) Maier, J. P. In *Kinetics of Ion-Molecule Reactions*; Ausloos, P., Eds.; Plenum Press: New York, 1979.
- (8) Müller-Dethlefs, K.; Sander, M.; Schlag, E. W. Chem. Phys. Lett. 1984, 112, 291.
 - (9) Hepburn, J. W. Chem. Soc. Rev. 1996, 25, 281.
- (10) Schlag, E. W. ZEKE Spectroscopy; Cambridge University Press: Cambridge, U.K., 1998.
 - (11) Zhu, L.; Johnson, P. M. J. Chem. Phys. 1991, 94, 5769.
 - (12) Krause, H.; Neusser, H. J. J. Chem. Phys. 1992, 97, 5923.
- (13) Park, S. T.; Kim, S. K.; Kim, M. S. J. Chem. Phys. 2001, 114, 5568
 - (14) Park, S. T.; Kim, S. K.; Kim, M. S. Nature 2002, 415, 306.
 - (15) Park, S. T.; Kim, M. S. J. Am. Chem. Soc. 2002, 124, 7614.
 - (16) Ng, C. Y. Annu. Rev. Phys. Chem. 2002, 53, 101.
- (17) Seiler, R.; Hollenstein, U.; Softley, T. P.; Merkt, F. J. Chem. Phys. 2003, 118, 10024.
- (18) Kwon, C. H.; Kim, H. L.; Kim, M. S. J. Chem. Phys. 2002, 116, 10361.
- (19) Kwon, C. H.; Kim, H. L.; Kim, M. S. J. Chem. Phys. 2003, 118, 6327.
- (20) Kwon, C. H.; Kim, H. L.; Kim, M. S. J. Chem. Phys. 2003, 119, 215.
- (21) Kwon, C. H.; Kim, H. L.; Kim, M. S. J. Chem. Phys. 2003, 119, 4305.
 - (22) Lee, M.; Kim, M. S. J. Chem. Phys. 2003, 119, 5085.
 - (23) Lee, M.; Kim, M. S. J. Chem. Phys., in press.
- (24) Becker, U.; Shirley, D. A. VUV and soft X-ray photoionization; Plenum Press: New York, 1996.
- (25) Samson, J. A. R. Techniques of Vacuum Ultraviolet Spectroscopy; Wiley: New York, 1967.
- (26) Frisch, M. J.; Trucks, G. W.; Schlegel, H. B.; Scuseria, G. E.; Robb, M. A.; Cheeseman, J. R.; Zakrzewski, V. G.; Montgomery, J. A., Jr.; Stratmann, R. E.; Burant, J. C.; Dapprich, S.; Millam, J. M.; Daniels, A. D.; Kudin, K. N.; Strain, M. C.; Farkas, O.; Tomasi, J.; Barone, V.; Cossi, M.; Cammi, R.; Mennucci, B.; Pomelli, C.; Adamo, C.; Clifford, S.; Ochterski, J.; Petersson, G. A.; Ayala, P. Y.; Cui, Q.; Morokuma, K.; Malick, D. K.; Rabuck, A. D.; Raghavachari, K.; Foresman, J. B.; Cioslowski, J.; Ortiz, J. V.; Stefanov, B. B.; Liu, G.; Liashenko, A.; Piskorz, P.; Komaromi, I.; Gomperts, R.; Martin, R. L.; Fox, D. J.; Keith, T.; Al-Laham, M. A.; Peng, C. Y.; Nanayakkara, A.; Gonzalez, C.; Challacombe, M.; Gill, P. M. W.; Johnson, B. G.; Chen, W.; Wong, M. W.; Andres, J. L.; Head-Gordon, M.; Replogle, E. S.; Pople, J. A. Gaussian 98, revision A.6; Gaussian, Inc.: Pittsburgh, PA, 1998.
 - (27) Duschinsky, F. Acta Physicochim. URSS 1937, 7, 551.
 - (28) Sharp, T. E.; Rosenstock, H. M. J. Chem. Phys. 1963, 41, 3453.
- (29) Dyke, J. M.; Ozeki, H.; Takahashi, M.; Cockett, M. C. R.; Kimura, K. J. Chem. Phys. **1992**, *97*, 8926.
- (30) Lichtenberger, D. L.; Renshaw, S. K.; Bullock, R. M. J. Am. Chem. Soc. 1993, 115, 3276.
 - (31) Araki, M.; Sato, S.; Kimura, K. J. Phys. Chem. 1996, 100, 10542.
- (32) Watanabe, K.; Nakayama, T.; Mottl, J. J. Quantum Spectrosc. Radiat. Transfer 1962, 2, 369.
- (33) Varsanyi, G. Assignments for vibrational spectra of seven hundred benzene derivatives; Adams Higer: London, 1974.
 - (34) Wilson, E. B., Jr. Phys. Rev. 1934, 41, 706.
 - (35) Neugebauer, J.; Hess, B. A. J. Chem. Phys. 2003, 118, 7215.