Infrared-optical double resonance spectroscopy: A selective and sensitive tool to investigate structures of molecular clusters in the gas phase

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A spectrometer to investigate molecules and molecular clusters in the gas phase has been designed and fabricated. It combines supersonic jet expansion technique with fluorescence and mass detection methods to carry out IR–UV double resonance spectroscopic measurements. The fluorescence-detected infrared spectra of phenylacetylene and phenylacetylene–argon complex reveal that the π-bound argon acts as a perfect messenger to probe the acetylenic C–H stretching vibration of the phenylacetylene moiety in $S_0$ and $S_1$ states. The rare gas tagging method along with the infrared pre-dissociation technique revealed that the spectrum of phenylacetylene cation consists of two transitions, one of which can be assigned to the hot band and the other to acetylenic C–H stretching vibration of phenylacetylene. We have demonstrated that the difference in the internal cooling between argon and neon tagging can be used as a tool to assign hot bands.

**Keywords:** Double resonance spectroscopy, laser induced fluorescence, molecular clusters, resonant two photon ionization.

RARE gas clusters have been a subject of interest because of their role in understanding intermolecular interactions$^{4-3}$. Over the last two decades, the combination of laser spectroscopy and molecular beam techniques has contributed to achieve more insights on intermolecular interactions at an unprecedented microscopic level. The versatile supersonic jets and beams technique can be used to produce clusters which are molecular entities bound by weak intermolecular interactions, such as van der Waals forces or hydrogen bonding, by co-expansion of the required reagents. Molecular clusters produced under cold and isolated condition of supersonic jets allow us to probe structures, intermolecular potentials and dynamics at a microscopic level with the aid of various laser spectroscopic techniques. Much of the earlier work on molecular clusters was carried out using electronic spectroscopy$^4$. Electronic spectra of clusters exhibit characteristic spectral shifts depending on their size and/or structural differences, but do not give any structural information. Infrared (IR) spectroscopy along with various other spectroscopic techniques, such as NMR, has long been realized as an excellent technique to investigate local interactions present in molecules and molecular assemblies. However, IR spectroscopy in the gas phase, especially in supersonic jets and beams, cannot be achieved with conventional absorption measurements due to very low density. Further, selectivity is lost due to the presence of heavy mixture of undesired monomers and clusters along with the cluster/monomer of interest. In this respect, any spectroscopic technique to be used for the investigation of molecular clusters requires merits of having high sensitivity and selectivity. Lee and co-workers first demonstrated the IR spectroscopy of clusters by population labelling of target species using electronic transition, which enables the size/ structure selection$^5$. Brutschy and co-workers applied this for the first time to investigate fluorobenzene–methanol clusters, wherein they observed the C–O stretching vibration of methanol$^6$. Later this technique has been popularized by Mikami’s and Zwier’s groups simultaneously, to investigate hydrogen bonding in phenol–water and benzene–water clusters respectively$^7-8$.

The infrared spectra of argon (Ar) complexes of several aromatic molecules, such as benzene$^9$, aniline$^{10}$, phenol$^{11}$, have been reported in the literature. In comparison with bare molecules those spectra are almost unperturbed. On the other hand, it has been amply demonstrated that Ar-tagging significantly reduces the internal temperatures of the cluster ions, without affecting the structure of the parent cluster$^{12-14}$. These results form the basis for the ‘messenger’ technique$^{15-18}$. However, in a few instances it has been observed that under some conditions Ar binding does affect the vibrational structure. For instance, it has been shown that in phenol and the series of fluorophenols, Ar binding in the cationic state leads to lowering of the O–H stretching vibration, which can be interpreted as formation of charge-assisted ionic O–H...Ar hydrogen bond$^{19,20}$. On the other hand, the ability of neon tagging has not been explored, though it can be expected that neon would not be as effective as argon. In this article, we present the investigations carried out on phenylacetylenes.
lène (PHA) and its argon complex in the neutral ground (S₀), neutral first excited (S₁), and the cationic (D₃o) states probed using double resonance spectroscopic techniques. Further, we also demonstrate that the difference in the ability of internal cooling of neon and argon can be used to assign the hot bands.

Experimental

**Design and working of the spectrometer**

We have recently set up a spectrometer to investigate molecules and molecular clusters in the gas phase using double-resonance spectroscopic technique. The spectrometer combines the supersonic nozzle, which produces a cold jet of molecules, with fluorescence and ion detection systems. It consists of two cylindrical stainless steel (316L) chambers of size 250 mm–400 mm and 225 mm–150 mm, respectively, wherein the first dimension is height and second, the diameter. The two chambers are connected by a 650 mm long and 100 mm diameter stainless steel tube. The bigger chamber is connected to a 3000 l/s oil diffusion pump (OD250; Hid Hivac) backed with a 30 m³/h rotary vane pump (ED30; Hid Hivac), while the smaller chamber is connected to a 700 l/s diff stack diffusion pump (700M; Edwards) backed by the 12 m³/h rotary vane pump (2012; Alcatel). To prevent back-streaming of the diffusion oil pump, the bigger chamber is fitted with a custom-designed cold water buffer. Further, both the chambers are fitted with appropriate gate valves, which can be used to isolate the chambers from the oil diffusion pumps, if necessary. In order to achieve good pumping rates for the oil diffusion pumps and to reduce back-streaming, the highest grade diffusion pump oil (Santovac-5; Varian) has been used. Typical base pressure in the entire vacuum system is about 5 × 10⁻⁷ torr. The bigger chamber consists of four ports at right angles, two of which are used for guiding the lasers into the chamber. The third port is fitted with the fluorescence detection system along with the electrostatic grids for the time-of-flight mass spectrometer (TOFMS). The fourth port is used to connect the long tube which serves as a flight tube for the TOFMS. The pulsed nozzle is guided into the bigger chamber with the help of a stainless steel tube fitted with a sample holder. The sample holder and the pulsed nozzle can be heated up to 200°C, which facilitates investigating compounds which have low vapour pressures under ambient conditions.

The spectrometer has the advantage of detecting fluorescence and mass signals simultaneously in one set-up. A seeded supersonic free-jet is produced by co-expansion of the desired reagent in a buffer gas held at a high pressure (typically 3–5 atm) through a 200 μm diameter pulsed nozzle (Series-9; General Valve Corporation), driven by a pulsed valve driver (Iota One; General Valve Corporation) into a high-vacuum chamber. The pulsed valve driver works at a frequency of 10 Hz, with a typical on time of 120–150 μs, corresponding to a duty cycle of 0.15% and the resulting ambient pressure is about 1 × 10⁻³ torr. The excitation laser intersects the gas pulse at right angles 10 mm downstream from the nozzle. The ensuing fluorescence is collected perpendicular to both gas and laser pulses using 50 mm focal length f/1 lens. Only a single f/1 lens is used in the entire collection set-up to avoid reflection losses. The distance of the f/1 lens from the object (the interaction region of the jet and the laser) is adjusted such that the image of the fluorescence from the jet is focused on the photocathode of the PMT. Total fluorescence is detected using a photomultiplier tube (PMT – 9780SB+ 1252-5F; Electron Tubes Limited)/filter (WG305+BG3, Astro Optical Industries) combination.

Alternatively, mass detection can also be done using the TOFMS, which is based on the Wiley–McLaren design. It consists of three 50 mm grids, the repeller (R), extractor (E) and accelerator (A) and 1.1 m field-free flight tube. The distance between R and E is 2.5 cm and the distance between E and A is 1 cm. The interaction region between the gas and the laser beams is midway between the R and E grids, perpendicular to the direction of the TOFMS. The ions are extracted by high-voltage electric fields. Typically, voltages of +3000 and +1700 V are applied to the R and E grids, respectively, and A is grounded. The ions produced are extracted out of the ionization region by an electric field of 520 V cm⁻¹ and are accelerated by an electric field of strength 1800 V cm⁻¹. The accelerated ions enter the field-free region of length 1.1 m to reach the channel electron multiplier (CEM-KBL-25RS; Sjuts) operated typically around ~1800 V. A Faraday cup is used to collect the electrons, through a filter circuit biased at ~250 V and processed with a broadband preamplifier (EG&G PARC, Model 115). The signal from the PMT/CEM is digitized by a digital storage oscilloscope (TDS-1012; Tektronix), which is interfaced to a PC via the PCI-GPIB card. The waveforms are integrated and averaged utilizing a data acquisition program written using LabView. The integrated signal is plotted against the wavelength to get the fluorescence/R2PI spectrum.

**Spectroscopic techniques**

Three different experimental techniques have been used for complete spectroscopic identification and characterization of bare molecules and molecular clusters in supersonic jets. As mentioned earlier, the much-required selectivity of the neutral clusters is provided by electronic transitions, which can be monitored using either fluorescence and/or resonantly enhanced multiphoton ionization (REMPI) induced by an ultraviolet (UV) laser. The combined technique is called IR–UV double resonance spectroscopy, and has been applied to various neutral
clusters, as long as they show a distinct sharp transition in their S\textsubscript{1} \rightarrow S\textsubscript{0} electronic spectrum. In this technique a pulsed UV laser excites the origin band of the S\textsubscript{1} \rightarrow S\textsubscript{0} transition and the ensuing fluorescence or REMPI signal is monitored as a measure of the ground-state population. Prior to the UV laser an IR laser is introduced and its wavelength is scanned. When the IR wavelength is resonant with a vibrational transition of the species, the IR absorption induces reduction of the population, which is then detected as a decrease in the fluorescence or REMPI signal intensity. The resultant spectrum is called fluorescence detected IR (FDIR) or ion detected IR (IDIR) spectrum depending upon the detection scheme used. In the electronic excited state an alternative method called UV-IR double resonance spectroscopy is used, which is just a time-inverted method of the IR-UV spectroscopic technique. In this method, an UV laser prepares the molecules in the zeroth vibrational level of the S\textsubscript{1} state. An IR pulse, introduced following the UV pulse within the radiative lifetime of the S\textsubscript{1} state, pumps the molecules to higher vibrational levels in the S\textsubscript{1} state. For most of the molecules the fluorescence quantum yield increases with the higher vibrational energy in the S\textsubscript{1} state. When the IR wavelength is resonant with a vibrational transition of the species in the S\textsubscript{1} state, the IR absorption leads to a reduction of the quantum yield, which is then detected as a decrease in the fluorescence intensity. Only the fluorescence detection method is suitable for this technique. To record the IR spectra in the cationic state, infrared pre-dissociation (IRPD) spectroscopic technique is used for molecular clusters. In this technique, the cluster cations are produced via REMPI using and UV pulse. The IR laser pulse is delayed relative to the UV pulse, typically about 10 ns, and scanned. When the IR wavelength is resonant on vibrational transitions of the cluster ion, the vibrational excitation leads to vibrational pre-dissociation of the parent cluster ion, leading to the production of fragment cations and lowering of the parent ion signal. The IR spectrum is thus detected by monitoring the loss of the parent ion signal.

Lasers

In order to perform the double resonance spectroscopic experiments, UV and IR laser sources are required. The UV source is a frequency-doubled output of a tunable dye laser (Radiant Dyes, Narrow Scan GR) pumped with second harmonic of a Nd: YAG laser (Continuum, Surelite I-10). On the other hand, the tunable IR source is an idler output of a LiNbO\textsubscript{3} optical parametric oscillator (Euroscan Instruments, Custom LiNbO\textsubscript{3} OPO) pumped with an injection seeded-single mode Nd: YAG laser (Quantel, Brilliant B). The dye laser was calibrated using the optogalvanic method and the IR OPO was calibrated by measuring the wavelength of red light (~645 nm) generated by the sum frequency of signal and pump beams using a standardized monochromator (Spex 0.3 m). The absolute frequency calibration is within ±3 cm\textsuperscript{-1}. The typical bandwidth of both UV and IR lasers is about 1 cm\textsuperscript{-1}. The timing between the IR and UV lasers was controlled electronically with a digital delay pulse generator (DDG-555; Berkeley Nucleonics Corporation).

Results and discussion

Figure 1 shows the one colour-resonant two photon ionization (1C-R2PI) mass spectrum of the PHA–Ar system recorded using the excitation laser fixed at 35,846 cm\textsuperscript{-1}. Apart from PHA and PHA–Ar, the mass spectrum also consists of peaks due to PHA–water complex and PHA dimer. Figure 2 a shows the LIF spectrum of PHA recorded with argon as buffer gas. Figure 2 b and c also depict the 1C-R2PI spectra of PHA\textsubscript{a} and PHA–Ar complex respectively. The band-origin transitions of PHA and PHA–Ar complex are at 35,876 and 35,846 cm\textsuperscript{-1} respectively, and are in good agreement with those reported in the literature\textsuperscript{23,24}. The origin band of the argon complex is shifted to the red by 30 cm\textsuperscript{-1} relative to the bare PHA and is comparable to that of the benzene–argon complex\textsuperscript{25}. The 1C-R2PI spectrum of PHA–Ar complex also shows two low-frequency transitions at 15 and 41 cm\textsuperscript{-1} respectively relative to the band origin which can be assigned to the bend and stretch along the intermolecular coordinate\textsuperscript{26}.

The infrared spectra in the acetylenic C–H stretching region were recorded using double resonance technique. Figure 3 depicts the FDIR spectra of PHA and its argon complex in the S\textsubscript{0} and S\textsubscript{1} states. The FDIR spectrum of bare PHA (Figure 3 a) shows two intense transitions at

![Figure 1](image-url)
3325 and 3343 cm\(^{-1}\) respectively, along with few weak transitions. The two intense transitions have been assigned to Fermi resonance bands between acetylenic C–H stretching vibration and a combination of one quantum of C≡C stretch and two quanta of C≡C–H out-of-plane bend\(^{25}\). Other weak features observed in the FDIR spectrum have been assigned to transitions arising out of higher order coupling terms\(^{25,27}\). The IR spectrum of the argon complex (Figure 3 b) is almost identical to that of the bare molecule, with the experimental uncertainty of ±1 cm\(^{-1}\). This implies that the attachment of Ar to PHA does not perturb the acetylenic C–H oscillator. The FDIR spectra of PHA (Figure 3 c) and PHA–Ar complex (Figure 3 d) in the S\(_1\) are almost identical. In these two spectra the lone transition at 3325 cm\(^{-1}\) can be assigned to the acetylenic C–H stretching vibration in the S\(_1\) state.

For a set of Fermi resonance bands with a separation of \(\Delta E\) and intensity ratio of \(R\), and assuming that the intrinsic (unperturbed) overtone intensity is negligible, the unperturbed separation \(\Delta E_0\) and the coupling matrix element \(W\) are given by\(^{28,29}\).

\[
\Delta E_0 = \Delta E \cdot \frac{R - 1}{R + 1}, \tag{1}
\]

\[
W = \Delta E \cdot \frac{\sqrt{R}}{R + 1}. \tag{2}
\]

De-perturbation analysis was carried out for the 3325 and 3343 cm\(^{-1}\) peaks using eqs (1) and (2) with \(R \approx 1.00\) and \(\Delta E = 18\) cm\(^{-1}\). We obtained values \(\Delta E_0 \approx 0\) cm\(^{-1}\) and \(W \approx 9\) cm\(^{-1}\). (The average value of \(R\) obtained from ten scans of the IR spectra was 1.02 ± 0.05, hence we have used \(R \approx 1\) for the de-perturbation analysis.) This implies that the energy difference between the zero-order vibrational states corresponding to the acetylenic C–H stretching vibration and the combination band of one quantum C≡C stretch and two quanta of C–H in-plane bend is within the bandwidth of the laser. Further, the stretching frequency of the unperturbed acetylenic C–H group can be estimated as 3334 cm\(^{-1}\). This implies that upon electronic excitation the acetylenic C–H stretching vibration shifts to the red by 9 cm\(^{-1}\).

The high-resolution REMPI spectrum for the S\(_1\) ← S\(_0\) electronic transition of the PHA–Ar complex was simultaneously fitted with the rotational constants for the ground and excited states\(^{30}\). The equilibrium structure consists of Ar atom bound to the \(\pi\) electron density of the benzene ring with the shift in Ar atom towards the acetylene group. On the other hand, Dreizler \textit{et al.}\(^{31}\) measured the microwave spectra of 24 isotopomers of the PHA–Ar complex. Along with high level \textit{ab-initio} calculations carried out at CCSD(T)/CBS, these authors concluded that the Ar atom is placed over the benzene ring. However, there is substantial motion over the benzene ring to predict the exact position of the Ar atom\(^{31}\). The near identical IR spectra of PHA and PHA–Ar complex in the acetylenic C–H stretching region, as depicted in Figure 3, indicates that the binding of Ar to PHA does not perturb the acetylenic C–H oscillator, both in the S\(_0\) and S\(_1\) states. Since the Fermi resonance transitions in the S\(_0\) state involve the coupling of the acetylenic C–H stretching vibration and a combination of one quantum of C≡C stretch and two quanta of C≡C–H out-of-plane bend, the persistence of the Fermi resonance in the PHA–Ar complex clearly indicates that the Ar atom does not interact with the acetylenic moiety of PHA and is therefore expected to bind to

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**Figure 2.** (a) LIF spectrum of phenylacetylene. 1C-R2PI spectrum of (b) phenylacetylene and (c) phenylacetylene–Ar. Spectra a and c were recorded using argon as buffer gas, while (b) was recorded using helium as buffer gas.

**Figure 3.** FDIR spectra of (a) phenylacetylene, (b) phenylacetylene–Ar in the S\(_0\) state, (c) phenylacetylene, and (d) phenylacetylene–Ar in the S\(_1\) state.
the $\pi$ electron density of the benzene ring. The FDIR spectrum is consistent with the structure of the PHA–Ar complex predicted by high-resolution electronic spectroscopy and microwave spectroscopy. The FDIR spectra depicted in Figure 3 indicate that the Ar atom bound to PHA acts as a perfect messenger to probe the vibrations of the acetylenic group of PHA. Extending this analogy to the cationic state, one would expect that the infrared spectrum of [PHA–Ar]$^+$ would mirror the infrared spectrum of [PHA]$^+$. Figure 4a shows the IRPD spectrum of [PHA–Ar]$^+$ in the acetylenic stretching region, which consists of two bands around 3268 and 3276 cm$^{-1}$ respectively. In order to assign the bands it is necessary to understand the origins of these two bands. To begin with, four distinct possibilities exist for the appearance of two bands in the IRPD spectrum of [PHA–Ar]$^+$. (i) Reappearance of Fermi resonance/anharmonic coupling similar to that observed in the S$_0$ state. (ii) Combination bands involving intermolecular modes over the acetylenic C–H stretching vibration. (iii) Hot bands due to the formation of internally hot [PHA–Ar]$^+$ following resonant two-photon ionization. (iv) Appearance of two distinct isomers of [PHA–Ar]$^+$ to two different $\pi$ and $\sigma$ complexes, similar to that observed in the case of [phenol–Ar]$^+$ using electron impact ionization, leading to formation of C–H...Ar hydrogen bond in the cation complex.

To resolve the above issues and to assign the two bands, we recorded the IRPD spectrum of [PHA–Ne]$^+$ complex cation. The rationale behind such an exercise was as follows. (i) In the case of reappearance of Fermi resonance/anharmonic resonance in the D$_0$ state, the IRPD spectrum of [PHA–Ne]$^+$ would be identical to that of [PHA–Ar]$^+$. This is due to the fact that the $\pi$-bound Ar atom does not perturb the acetylenic C–H stretching vibration. Therefore, the $\pi$-bound Ne atom is not expected to make any difference, as the binding of Ne is weaker than that of Ar. (ii) Since the frequencies of the intermolecular vibrations would change upon the binding potential energy, it is expected that the positions of the combination bands of the neon complex would be different compared to the argon complex. (iii) In the instance of observation of a hot band, replacement of Ar with Ne would result in an increase in the intensity of the lower energy band, without change in the band positions. Such an observation can be attributed to the reduced efficiency of internal cooling of Ne relative to Ar. (iv) Finally, if the two peaks are due to $\pi$ and $\sigma$ complexes of [PHA–Ar]$^+$, the low frequency band which can be probably assigned to the $\sigma$ complexes, resulting in C–H...Ar hydrogen bond cation complex would disappear completely, as the ability of neon to form C–H...Ne hydrogen bond would be much less in comparison with argon. Comparison of IRPD spectra of [PHA–Ar]$^+$ and [PHA–Ne]$^+$ should correspond to one of the four possibilities listed above. The IRPD spectrum of [PHA–Ne]$^+$ is shown in Figure 4b. This spectrum clearly shows that the intensity of the lower frequency band at 3268 cm$^{-1}$ is about 25% higher than the corresponding band in the spectrum of the argon complex. Based on the analysis presented above, the 3268 cm$^{-1}$ transition can be unambiguously assigned to the hot band. Therefore, the 3276 cm$^{-1}$ band can now be assigned to the acetylenic C–H stretching vibration in [PHA–Ar]$^+$ and [PHA–Ne]$^+$. Extending the analogy observed for the S$_0$ and S$_1$ states to the D$_0$ state, the acetylenic C–H stretching vibration of the phenylacetylene cation [PHA]$^+$ can be pegged at 3276 cm$^{-1}$.

**Conclusion**

The IR spectra of PHA and its Ar complex were recorded in the acetylenic C–H stretching region using double resonance technique in the S$_0$, S$_1$, and D$_0$ states. In the S$_0$ state, the acetylenic C–H stretching vibration is involved in a Fermi resonance with the combination band of two quanta of out-of-plane C=C–H bend and one quantum of C=C stretch. The perturbation analysis places the C–H stretching vibration at 3334 cm$^{-1}$ in the S$_0$ state. In the S$_1$ state the corresponding frequency is 3325 cm$^{-1}$, a shift of ~9 cm$^{-1}$ relative to the S$_0$ state. The IRPD spectrum of phenylacetylene–argon cation complex shows two transitions at 3268 cm$^{-1}$ and 3276 cm$^{-1}$ respectively. Comparison of the relative intensities of the two transitions with those in the IRPD spectrum of phenylacetylene–neon cation complex reveals that the transition at 3268 cm$^{-1}$ corresponds to a hot band, while the transition at 3276 cm$^{-1}$ is the acetylenic C–H stretching vibration of the PHA$^+$ moiety. Thus the differences in the internal cooling due to argon and neon tagging can be a useful tool to understand and assign hot bands.

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**Figure 4.** IRPD spectrum of (a) phenylacetylene–Ar and (b) phenylacetylene–Ne complex cations.
RESEARCH ARTICLES


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