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Vacuum Ultraviolet Photoionization of Complex Chemical Systems Oleg Kostko, Biswajit Bandyopadhyay, Musahid Ahmed

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Keywords

Mass spectrometry, Synchrotron, Hydrogen Bonds, Combustion, Proton Transfer, Laser Ablation,

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Abstract

Tunable VUV radiation coupled to mass spectrometry is applied to the study of complex

chemical systems. The identification of novel reactive intermediates and radicals is revealed in

flame, pulsed photolysis and pyrolysis reactors leading to the elucidation of spectroscopy, reaction

mechanisms and kinetics. Mass resolved threshold photoionization-photoelectron coincidence

measurements provides unprecedented access to vibrationally resolved spectra of free radicals

present in high temperature reactors. Photoionization measurements in water clusters, nucleic acid

base dimers and their complexes with water provide signatures of proton transfer in hydrogen

bonded and π -stacked systems. Experimental and theoretical methods to track ion-molecule

reactions and fragmentation pathways in intermolecular and intramolecular hydrogen bonded

systems in sugars and alcohols are described. Photoionization of laser ablated molecules, clusters,

and their reaction products inform thermodynamics and spectroscopy of relevance to

astrochemistry and catalysis. New directions in coupling VUV radiation to interrogate complex

chemical systems are discussed.

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Introduction

The valence shell, the outermost shell of electrons in an atom or molecule, contributes most to the physical properties of chemical systems. It is these electrons that are shared in covalent bonding, transferred in ionic systems, coupled to form bands in bulk material, and interact most strongly with the environment. Excitation of valence electrons in molecules to higher lying electronic states typically require vacuum ultraviolet (VUV) radiation, spanning 6-13 eV, which contributes to three processes: photodissociation, photon emission and photoionization (1). In this review, we will focus on application of VUV photoionization to decipher kinetics, dynamics and transformations in complex chemical systems.

Traditionally, VUV photoionization, has contributed to the understanding of the details of ionization energies (IEs) and thermodynamic properties in molecules (2-4). In 1958, a discharge lamp coupled to a mononchromator led to the first publication in VUV photoionization based mass spectrometry (5). Photoionization efficiency curves (PIEs) for the parent and fragments were recorded, and already as stated in the abstract – "the results can be interpreted in terms of potential surfaces and electronic transition probabilities" sets the tone for this review. The advantageous nature of tunable VUV radiation to study fundamental properties in gases (those could be obtained in "lecture bottles" and cylinders) was seen at the weekend runs at a French synchrotron in the early 70's (6). The coupling of a molecular beam to a synchrotron was first reported in 1973, measuring PIE curves for acetylene and carbon dioxide (7). In the following decades, as synchrotrons became more accessible to the world of physical chemists, in addition to fundamental photoionization studies on atomic and molecular systems, cluster beams coupled to quadrupole mass spectrometers gave rise to a series of studies, mostly on H-bonded and van der Waals clusters.(8) Ionization and appearance energies (IEs and AEs), proton affinities, solvation energies and intermolecular bond energies were determined for a series of small clusters, while

intermolecular interactions in ionic clusters were tracked via proton transfer (PT) upon photoionization. With the advent of tunable VUV radiation from 3rd generation synchrotron based sources (9) the scope of work shifted to the realm of chemical dynamics (2; 10). Based on the success of fundamental studies on small molecules, it became evident that VUV photoionization could play a unique role in providing insight into complex systems.

Single photon ionization with VUV photons is an efficient (and "soft") method of ionizing fragile molecules, whereby the tunable radiation provides precise threshold ionization imparting little excess internal energy into the molecular ion. This greatly reduces the extensive fragmentation usually observed with other multiphoton or electron impact ionization techniques (11). Furthermore, tunable VUV ionization has been shown to be a selective, yet universal technique in elucidating molecule-specific information since it can resolve isomers via IE, absorption cross-sections follow rules which are quantifiable, and all molecules will ionize beyond its IE and hence can be detected in a mass spectrometer. In the last few years, there has been an explosion of work around various synchrotrons worldwide where physical chemists are using tunable VUV photons coupled to mass and electron spectrometers to study elementary reactions pertinent to combustion, heterogeneous chemistry on aerosols, PT in H-bonded clusters, as well as spectroscopic properties of molecules relevant to astrochemistry and catalysis. As experimental techniques mature, and new ones emerge, processes those are more complicated can be studied, providing significant gains in our understanding of real world systems (11).

1. Photoionization in Combustion Relevant Processes and Identification of Reactive
Intermediates in Gas Phase Chemical Kinetics.

One example of a real world process is combustion. With the advent of the internal combustion engine over 150 years ago, it would appear that all is understood in a fundamentally simple system that comprised fuel (hydrocarbons) and an oxidizer. However, the chemistry that occurs upon ignition coupled to the fluid dynamics over a wide range of temperatures and pressures, is complicated. The introduction of new fuels, demands for higher efficiency, reduced emissions for pollution and environmental mitigation requires detailed understanding of combustion at the fundamental level. VUV ionization coupled to a number of reactors simulating combustion environments have been deployed at various synchrotron radiation facilities worldwide since the first low-pressure, laminar premixed flame was coupled to a molecular beam photoionization mass spectrometer at the Advanced Light Source (ALS) in Berkeley (12). A comprehensive and detailed view of the very exciting work that has been accomplished over the last decade is described by Qi (13), and below we highlight two examples showcasing the unique application of tunable VUV to combustion chemistry.

The ALS-based photoionization technique played a significant role to identify a class of molecules, the enols, which used to be absent from flame chemistry modeling (14). Enols have an OH group adjacent to a C=C double-bond, and are isomers of aldehydes/ketones. The characteristic PIE curves of enols can easily be differentiated from those of aldehydes/ketones as shown in Figure 1. It was observed that enols were important mainly in alcohol flames where they can be as abundant as the isomeric aldehydes/ketones, and with bio-based fuels which have higher alcohol content (15). VUV photoionization was also crucial in the identification of C₄H₃ and C₄H₅ radical species in hydrocarbon flames. A long-lasting debate about the importance of the so-called *n*-C₄H₃ (CHCHCCH) and *n*-C₄H₅ (CH₂CHCHCH) isomers compared to their resonantly stabilized

i-isomers (CH₂CCCH and CH₂CHCCH₂) came to an end when Hansen *et al.* showed that predominantly, the resonance-stabilized radicals (i-C₄H₃ and i-C₄H₅) are present (16).

Since these pioneering studies with low pressure flames, a set of reactors designed to simulate realistic combustion conditions under various temperature and pressure regimes to probe detailed chemical mechanisms have been deployed at various synchrotron facilities. Pressures up to few hundred torr, timescales of microseconds, and temperatures up to 1500 C became accessible in a flow reactor (hot nozzle) coupled to molecular beams (17; 18). Photoionization mass spectrometry (PIMS) studies of the unimolecular decomposition pathways of acetaldehyde (17), furan (19), the benzyl radical (20), and cyclopentadienone (21) provide glimpses to processes relevant in renewable biofuel processing while the reactions of the phenyl radical with propyne (18), allene, propylene (22), 1,3-butadiene (23), acetylene (24), oxygen (25) and the reactions of benzyl (26), and napthyl (27) radicals with acetylene, shed light on molecular growth mechanisms relevant to soot formation. For instance, the phenyl radical with acetylene study (24) provided first experimental evidence of the HACA (hydrogen abstraction followed by acetylene addition) mechanism, postulated over three decades ago, to be operative leading to PAH formation (naphthalene) under combustion relevant conditions. However, subsequently it was shown, formation of the third aromatic ring (anthracene or phenanthrene) was not so facile when the napthyl radical was reacted with acetylene (27).

The power of tunable VUV coupled to electronic structure calculations for spectroscopic characterization of transients in combustion is nicely illustrated in the case of cyclopendienone, a cyclic ketone ($C_5H_4=O$) that formally has 4π electrons and is therefore "antiaromatic" according to the Hückel 4n+2 electron rule. A photoionization and photoelectron photoion coincidence (PEPICO) study performed on cyclopentadienone (28) formed via pyrolysis of o-phenylene sulfite

showed that the nominal highest occupied molecular orbital has π -bonding character within the five-membered ring. The study supports the notion that the lowest-lying excited electronic state results from the associated $\pi\pi^*$ excitation, in contrast to most aldehydes and ketones those do not have the antiaromatic character of C₅H₄O. A closely related species, the hydroxycyclopentadienyl radical formed in the unimolecular decomposition of ortho-methoxyphenol (guaiacol) has been identified via a mass selected threshold photoelectron spectroscopy (ms-TPES) study complemented by Franck Condon (FC) simulations (29).

Hemberger et al. employed imaging photoelectron photoion coincidence (iPEPICO) to a similar heated tubular reactor to identify isomers in the decomposition of radicals and transients (30). Pyrolysis studies of m-xylyl bromide studied with ms-TPES and FC simulations provided unambiguous evidence for the formation of m-xylyl radical with the exclusion of o- and p-xylyl, and also methyl tropyl radicals (Figure 2). The m-xylyl radical subsequently isomerizes and decomposes with the elimination of a hydrogen atom to form para-xylene. Vibrational features were identified corresponding to the activity of two ring-deformation modes associated with C-C-C bending vibrations (m-xylyl radical) and for para xylene, it was C-C-C bending and C-C stretching vibrations which provides signatures for structural elucidation. Lang et al. used the ms-TPES to identify a series of C₄H₅ isomers – 2 butyn-1-yl and 1-butyn-3-yl, and C₄H₇ isomers, 1methylallyl and 2-methylally radicals formed via the pyrolysis of C₄H₅ and C₄H₇ nitrite and bromide radical precursors (31). The quality of the fits of the simulated FC envelope to experimental spectra for the C₄H₅ isomers was diminished due to the internal rotation of the methyl groups which would cause broadening of the spectra and the harmonic approximation often fails for these internal rotations. The 2-methylallyl provided for a much better fit with a series of vibrational progressions observed quite clearly in the spectrum arising from transitions to the X⁺

 1 A₁ and A⁺ 3 A states from the X 2 A₂ ground state. However, one complication in these comparisons between FC simulations and experiments are that typically hot band contributions are not included as clearly observable in figure 2 (b), where at 7.05 eV, the peak in the m-xylyl photoelectron spectrum is absent in the theoretical simulation.

The latest addition to the combustion relevant experiments is the jet-stirred reactor with molecular beam sampling capabilities, where IEs and fragmentation patterns were used to identify a series of alkyl peroxides formed in the oxidation of n-butane (32; 33) and probe the low-temperature oxidation of dimethyl ether (32; 34). For example, the keto-hydroperoxide (HOOCH₂OCHO) was identified to the dominant species based on calculated conformer-dependent relative ground state populations accessible in the experiment, calculating their adiabatic IEs (AIE) and comparing these to experiment (34). The lowest energy conformer featured a H-bonded seven membered ring while a range of other structures were within 8 kcal/mol relative to the most stable structure.

In the future, it would be interesting to see what new chemistry will be identified when even higher pressures and temperatures are accessed via the introduction of miniature shock tubes coupled to VUV photoionization (35). Another promising trend is the coupling of flame sampling instruments to iPEPICO instruments demonstrated at the Swiss Light Source (36) and synchrotron Soleil (37). It is encouraging to see that a ms-TPES of the hydroxyl radical have been recorded in flame (36) and flow reactors (36; 38). To date, the combustion intermediates detected have been fairly low molecular weight species and a detailed comparison of the various ionization techniques does point to the superiority of VUV photoionization in this regard (39). However, combustion chemistry is increasingly moving towards complexity and nowhere is this more relevant than in observing molecular growth processes. A coupling of a flame sampling apparatus to aerosol based

mass spectrometry and VUV ionization (40) led to the intriguing suggestion that aliphatic bridged or oxygen bonded aromatic associations might be formed in soot processes as opposed to the more thermodynamically stable polycyclic aromatic hydrocarbons (PAH).

The need for isomer resolved mechanistic information and chemical kinetics for complex molecules, led Osborn and Taatjes to develop a laser photolysis based flow reactor, which they have put to use to study transients, radicals and reactive intermediates (41). VUV-PIMS has been used almost a decade ago to identify the methyl peroxy radical directly by the measurement of a PIE curve, while the ethylperoxy radical was identified via dissociative ionization products, since the C₂H₅OO cation was unstable even at threshold energies (42). In both cases, calculated IEs, and an FC simulated PIE curves were used to identify the transient species. In 1949, Rudolf Criegee proposed a mechanism for the oxidation of alkenes by ozone. This mechanism has gained wide acceptance despite the lack of any direct observation of the so-called Criegee intermediates, of which the simplest is carbonyl oxide (CH₂OO). In 2008, Osborn, Taatjes and co-workers succeeded in observing CH₂OO in the Cl-initiated oxidation of dimethyl sulfoxide in the presence of excess O₂ (43) and the Criegee intermediate was identified by comparing PIE curves with FC simulations. This was followed by measuring CH₂OO formed in the reaction of CH₂I₂ with O₂ (44). In this particular case, a calculated FC envelope fit the observed PIE curve extremely well, and other structural isomers, dioxirane and formic acid could be eliminated. The ability to separate conformers via threshold allowed for the measurement of the reactivity of syn- and anticonformers of a larger Criegee intermediate (CH₃CHOO) (45). A similar strategy has been applied to detect directly the hydroperoxyalkyl radical (QOOH) intermediate in the oxidation of 1,3cycloheptadiene (46). Typically peroxy radicals isomerize to QOOH which then leads to unimolecular decomposition, but in this particular time resolved experiment, the QOOH

intermediate could be captured in the 0-60 ms time scales accessible in the time-of-flight mass spectrometer. The identification of these intermediates allowed for novel elementary kinetic and mechanistic details which have important ramifications in combustion (46) and environmental (47) chemistry.

2. Water Clusters, Proton & Excitonic Charge Transfer studied with photoionization

The use of clusters as a vehicle for studying molecular properties allows for systematic investigations of how bulk properties of a substance arise from the isolated molecule as size is increased. Water is fundamentally very interesting and its importance in biological and atmospheric processes on Earth is unquestioned, and photoionization of its clusters provides insight into its thermodynamic and bonding properties. Water clusters were ionized using tunable VUV energy (10 - 14 eV) and analyzed by reflectron time-of-flight mass spectrometry (48). Photoionization of neutral clusters creates unstable cations, $[(H_2O)_n]^+$, which undergo very fast intracluster charge redistribution on the subpicosecond time scale to produce protonated clusters. The AEs of a series of protonated water clusters up to 79 molecules were determined from the PIE curves. These AEs represent an upper limit of the adiabatic IE of the corresponding parent neutral water cluster in the supersonic beam. The experimental results show a sharp drop in the AE for the small neutral water clusters (from 12.62 ± 0.05 eV to 10.94 ± 0.06 eV, for (H₂O) and (H₂O)₄, respectively), followed by a gradual decrease for clusters up to (H₂O)₂₃ converging to a value of 10.6 eV (± 0.2eV). The decrease of the protonated water cluster ion AE values as a function of cluster size for n < 20 is probably a manifestation of the stabilization of the overall cluster geometry and intracluster H-bonding network. A comparative study where the valence photoelectron spectra of water clusters were measured and compared to theory explicitly showed the size dependence of the vertical IEs (VIE) of the outermost orbitals (49). This is an interesting experiment where by changing source conditions, different distributions of water clusters were produced and photoelectron spectra recorded. The authors then compared calculated VIE's to the peak maxima in the photoelectron spectrum. A shift toward lower values is observed in agreement with mass spectrometry based approach discussed earlier, and this lowering for the smaller clusters sizes was attributed to charge delocalization. The ionization energies of the larger clusters decreased linearly with inverse cluster radius and at the asymptote approached that of liquid water. A quantum mechanical sampling of the distribution of different initial state geometries of the clusters provided a qualitative explanation of the photoionization process reflected in the measured photoelectron spectrum.

The water dimer has been studied over decades as a prototypical H-bonded system predominantly in the neutral state. Ionization allows it to become a test-bed for theory and also to glean spectroscopic signatures for PT dynamics. An added advantage is that, the dimer does not dissociate till 11.72 eV, when ionization occurs by removing an electron from the lone pair of the H-bond donor. Experimentally this allowed for a PIE to be measured for the unfragmented dimer, and upon differentiation, produces a spectrum similar to a photoelectron spectrum which was then compared to theory (50). This study revealed that the dynamics of the PT process can be highly dependent, both qualitatively and quantitatively, on the nature of the ionized state. In the water dimer, PT is driven by the localization of the hole on one individual monomer. The shared proton stretch and the oxygen-oxygen stretch are strongly coupled yielding large anharmonic effects. The calculations showed that the FC factors can entirely hide the adiabatic onset of ionization as well as shift the peak to the red of the vertical value (50). Complementary to these spectroscopic studies, the thermodynamic properties derived by measuring OH⁺ and H₃O⁺ AEs using iPEPICO led to the determination of the proton affinity of water with unprecedented accuracy (51).

Water clusters fragment extensively and lead to very efficient PT even upon threshold ionization. However, with aggressive jet cooling, unfragmented water (cationic) clusters beyond the dimer have been observed (52; 53). Infrared spectroscopy coupled to VUV photoionization have been performed on size-selected unfragmentated water clusters (54) and the discussion has stemmed around evaporative cooling by the carrier gas (Ar) leading to the stabilization on these clusters. To test this hypothesis, PIMS measurements with high pressure Ar (7 bar) in a supersonic expansion of water through a small (50 µm) nozzle led to the observation of unfragmented water clusters up to n=9 (55). Efficient charge transfer was shown to occur via an Ar exciton process which occurred below the IE of water, shown clearly in Figure 3. It appears that solvation of the ionized water cluster by argon atoms stabilizes the (H₃O)⁺OH core, and the excess energy formed upon ionization leads to the evaporation of the Ar atoms and/or water molecules. Infrared studies have shown that ionized water clusters maintain a structure of H⁺(H₂O)_{n-1}(OH), very similar to that observed for protonated water clusters (54). Figure 3 shows that following the water cluster ionization onset, the unfragmented water cluster signal appears over the first Ar cluster exciton region and then dramatically decreases. Protonated water clusters, on the other hand, follow the Ar fluorescence signal (measured from pure Ar clusters, (56) and showing the signature of electronic excitation) for the full measured region. This difference in behavior suggests that evaporative cooling of the Ar cluster absorbs the excess electronic energy up to a limit of ~1.2 eV above ionization. In this regard, water-doped argon clusters behave in a similar way to an intermolecular columbic decay (ICD) process, where electronic energy excited at one site is channeled to a neighboring location where an electron is ejected (57). These kinds of processes are not limited only to VUV photoionization since a similar scheme of ICD was demonstrated for electron impact ionization of mixed Ar-water clusters (58).

3. Photoionization and Proton Transfer in Gas Phase Biomolecules & Their Complexes

Photoionization of gas phase biomolecules, their complexes and solvation provides a fertile ground to learn about intra- and intermolecular properties of these systems. Studying these species under well-defined conditions can distinguish between intrinsic and externally imposed properties (i.e. solvation) (59). Timely reviews by Qi and coworkers(60), and Schwell and Hochlaf (61) have extensively discussed studies of biomolecules and nonvolatile molecules with tunable VUV radiation paying particular attention to electronic structure and analytical chemistry applications and below the focus will be on DNA base dimers and their microhydration.

PIMS coupled with electronic structure calculations were performed for a series of nucleic acid base monomers and dimers to track ionization, proton and charge transfer in H-bonded, π -stacked and solvated systems (62-64). These studies showed that both the vertical and adiabatic IEs in these dimers are strongly affected by non-covalent interactions. For the thymine dimer (TT), asymmetric H-bonded and symmetric π -stacked isomers showed the largest VIE shift between monomer and dimer (0.4 eV) because of a charge stabilization of the ionized fragment by the dipole moment of the neutral fragments. On the other hand, the shift was much smaller for symmetric H-bonded isomers (0.1 eV) since the hole is delocalized and change in IEs is directly related to the extent of overlap between the MO. While H-bonded isomers undergo barrierless (or almost barrierless) PT producing protonated monomers, symmetric π -stacked structures form covalent bonds between fragments. The AE for the TT dimer suggests an origin from a π -stacked isomer. There was experimental evidence for a number of additional H-bonded isomers, as the AE for AH⁺ agrees well with the calculated dissociation threshold energies for an asymmetric H-bonded dimer. For the adenine dimer (AA), a more delocalized MO leads to larger geometrical relaxation and

poor FC factors for the lowest band. The onset of the PIE curve for the dimer and the AE of the protonated monomer was in reasonable agreement with the computed threshold for the lowest energy AA isomer (symmetric H-bonded). The analysis of vertical IEs revealed that the states derived from thymine ionization exhibit larger shifts that those derived from ionization of A in these dimers. The IE of T is affected more by non-covalent interactions (both stacking and H-bonding) than those of A, which has a lower IE. In the case of cytosine (63), evidence was presented for several isomers. In contrast to the dimers of AA, AT and TT, the IEs of cytosine were strongly affected by the inter-fragment interactions in the dimer, i.e., the lowest IE of the most stable dimer is red shifted by almost 1 eV relative to the monomer. However, here too, extensive barrierless PT occurred upon photoionization leading to the formation of CH⁺. In these proton-transferred structures, the positive charge is localized on the closed-shell protonated fragment, whereas the unpaired electron resides on the deprotonated moiety.

Theoretical calculations suggest that the photoionization relaxation patterns are different for π -stacked and H-bonded dimers (65; 66) and that PT requires a H-bonded pathway. To test this hypothesis, PIMS was performed with uracil (U) and 1,3-dimethyluracil (dmU) dimers along with theoretical calculations, since dmU dimers can only be formed in the stacked arrangement as the methyl substituents restrict H-bonding (67). As expected, extensive PT from the uracil dimer was observed when it becomes thermodynamically accessible. Following ionization of the methylated sample, the π -stacked dimer also showed a significant signal for protonated dmU cation demonstrating that PT is efficient in the methylated species despite the absence of H-bonds in the neutral dimer. The observed AEs agree with theoretically calculated values. Deuterated dimethyluracil (where only the two methyl moieties are deuterated) studies confirmed that the PT originates from the methyl group and not from the aromatic C-H sites. The nature of motions

promoting PT in the ionized state was very different for hydrogen-bonded and π -stacked systems and was explored theoretically guided by the experimental results. In the former, the PT reaction coordinate corresponds to a simple proton motion between the two uracil moieties due to an almost linear H-bond (N–H–O = 177°) and a relatively short distance between the proton-accepting and proton-donating atoms (2.794 Å at the FC geometry). In contrast, the structure of the stacked dmU dimer was not optimal for PT; that is, the C–O separation is 3.136 Å and (C–H–O) = 107° at the FC relaxed geometry. Therefore, a considerable rearrangement of the bases along the PT pathway was required to make the process energetically possible. The lowest-energy PT path involves a concerted change in the distances between the proton and the donating (C–H) and accepting (O–H) atoms. The C–O distance decreases by 0.6 Å as the system moves from the FC relaxed structure to the transition state. Near the transition state, the dimer assumes a T-shaped structure and as the system passes over the transition state, it relaxes back to a stacked geometry.

The occurrence of PT through a π -stacked dimer is not unique. Similar PT processes have been shown to occur across CH...O type H-bonds in the dimethyl ether dimer, using iPEPICO (68). Theory predicted that the neutral dimer would have multiple low energy isomers with variable CH..O interactions and all of these isomers undergo barrierless PT upon photoionization to the ground electronic state of the cation. The calculated AEs matched the experimental onsets for the protonated monomers confirming that the dissociation is dependent only on the product energetics.

Using coincidence imaging coupled to threshold ionization, Hochlaf and co-workers performed photoionization studies on valerolactam (piperidin-one) and its dimer (69; 70). A number of bands (combinational and pure) were recorded with vibrational resolution and their origins were captured via high level calculations. While the slow photoelectron spectra identified the cation in the ground state, S/N, competition from autoionization and predissociation, and congestion of features leading

to fairly structureless spectra precluded assignment to the high lying monomer B and C states. Four low lying doublet states of a cyclic dimer (2 H-bonds) gave rise to a featureless spectrum; however, protonated monomers could be clearly identified from AE measurements. Upon ionization, one of the H-bonds breaks resulting in unfurling of the ring and isomerization to a more stable structure followed by PT to form the protonated monomer.

Water plays a central role in chemistry and biology by mediating the interactions between molecules, altering energy levels of solvated species, modifying potential energy profiles along reaction coordinates, and facilitating efficient proton transport through ion channels and interfaces. A PIMS study on dmU clusters however revealed, that upon microhydration, PT is effectively blocked (71). Instead, a new pathway opens up in which the protonated dmU is generated by PT from the ionized water molecule to dmU. To gain further insight into the electronic structure of hydrated species and to validate theory, a PIE and its derivative of the smallest hydrated cation, dmU(D₂O), were measured. The differentiation of the PIE curve allows identification of multiple ionized states (the peaks on the differentiated PIE curve correspond to the VIEs). The PIE (black) and the differentiated (red) curves are shown in Figure 4, along with the computed VIEs. The curve features the ionization onset at ~8.6 eV and a series of peaks between 8.5 and 11.5 eV. The character of these states are illustrated by the respective MOs, and suggests that low-lying electronic ionized states correspond to the ionization from dmU, whereas the 3A" state at 11.2 eV is localized on water. The AE for protonated dmU is at 10.6 eV followed by a sharp rise at 11.2 eV suggesting efficient PT occurs when the 3A" state is accessed upon photoionization. Electronic structure calculations revealed that PT from water to dmU is barrierless when an ionized state localized on water is accessed. Possible adiabatic passage on the ground electronic state of the ionized system, though energetically accessible at lower energies, is not efficient. Thus, PT is controlled electronically, by the character of the ionized state, rather than statistically, by simple energy considerations.

4. Photoionization of Alcohols, Ethers & Sugars

Photoionization and dissociative photoionization studies were performed on a series of alcohols, ethers and sugars to observe and quantify how multiple sites for H-bonding – both inter and intramolecular, effect fragmentation dynamics. The ethanol dimer, due to the possibility of multiple dissociation pathways upon photoionization provides a rich space to test statistical theories of fragmentation. Li et al. (72) demonstrated with PIMS, that the dimer undergoes a rapid barrierless PT followed by two dissociation channels, one leads to protonated ethanol, and the other to a protonated formaldehyde-ethanol complex. The latter arises from a carboncarbon bond cleavage in the donor. Bodi (73) performed an elegant coincidence study and modelled the dissociation pathways using statistical theory, and ab initio calculations. He determined the 0 K AE for the C-C bond breaking process, and combining this with thermochemical cycles, quantified the energetics for the formation of the protonated formaldehyde-ethanol complex. Excited state calculations coupled with experimental data suggested that the dissociative photoionization process below 12 eV is statistical in nature, i.e. even if the initial ionization process may be site or state specific, internal degrees of freedom are completely scrambled within the lifetime of the parent ion dissociation. Upon photoionization beyond 11.55 eV, where vertical excitation occurs to a higher ionic state, Xiao et al. (74) provided evidence for a methyl migration in the ethanol dimer, giving rise to a methyl-ethanol ion complex. They came to this conclusion by calculating stationary points along the reaction coordinate and comparing AEs of various fragments to theoretical predictions.

The photoionization and dissociative photoionization of glycerol and its dimer were studied with both PIMS and theory, using isotopic labelling to investigate the nature of the major fragments and their corresponding AEs (75). It was discovered that the primary fragmentation of the glycerol radical cation occurs only via two routes. The first channel proceeds via a six-membered hydrogentransfer transition state leading to a common stable ternary intermediate - comprised of neutral water, neutral formaldehyde, and a vinyl alcohol radical cation, which exhibits a binding energy of about 42 kcal/mol and a very short (1.4 Å) H-bond. Fragmentation of this intermediate gives rise to an experimentally observed m/z 74, 62, 44 and 45. Fragments m/z 74 and 62 both consist of hydrogen-bridged ion-molecule complex with binding energy above 25 kcal/mol, whereas the m/z 44 species lacks such stabilization. This explained why water- or formaldehyde-loss products are observed first. The second primary fragmentation route arises from cleaving the elongated C-C bond. Also for this channel, intermediates comprised of hydrogen-bridged ion-molecule complexes exhibiting binding energies above 24 kcal/mol are observed. Energy decomposition analysis revealed that electrostatic and charge-transfer interactions are equally important in hydrogen-bridged ion-molecule complexes. The fragmentation patterns for the glycerol dimer radical cation can be described as a monomeric glycerol radical cation in the presence of a spectator glycerol, thus giving rise to a dissociation pattern similar to that of the monomer. However, the spectator glycerol lowers the barrier affiliated with separating the resulting products (water, formaldehyde, and a vinyl alcohol cation) as it aids the delocalization of positive charge and spin density and no water or formaldehyde loss is observed.

The photoionization of deoxyribose, a cyclic sugar, provides for an interesting comparison to glycerol, which starts out with an essentially linear configuration but upon ionization takes on a cyclic configuration held together by strong ionic H-bonds (75). A recurring theme of ionization

of sugars and alcohols is the elimination of neutral water, and ab initio molecular dynamics calculations, MO and FC analysis coupled with experimental AEs shed light into this channel. Most of the fragmentation events occur with a significantly weakened C4-C5 bond breaking (bond length of 1.84 Å), causing the cyclic structure to unfurl. This allows the OH groups on C3 and C5 to come close to one another, followed by water elimination suggesting the transient ionic structures are quite similar to the strong H-bonded ion complexes formed in glycerol photoionization.

Nahon and coworkers have studied dissociative photoionization of 2, 3- and 1, 3-butanediol using threshold PEPICO (76) and observed very different fragmentation patterns. This was due to the stability of the corresponding radical cation and breakdown diagrams were prepared from the internal-energy selected data and showed significant isomer specific behavior. For the 2, 3-butanediol, the neutral H-bridged structure was lower in energy. On the other hand, for 1, 3-butanediol, H-bridged structure had more energy than that of the C-C one, which results in different fragmentation pathways.

A new method where ionization is performed at various distances from the nozzle to sample different regimes of a supersonic expansion was implemented for studying ion-molecule and neutral reactions (77). Using methanol as a test case, since its VUV ionization properties have been well documented (78), it was shown that the intensity patterns of the cluster distributions vary significantly as a function of distance. The series of protonated methanol, methanol-dimethyl ether, and methanol-water clusters provide signatures of the ionization and clustering processes that occur in the expansion. At closer ionization distances, ion-molecule reactions dominate the cluster formation process resulting in formation of protonated species. The intensity distributions at various distances are qualitatively explained by Thomson's liquid drop model (79). At closer

distances, both supersaturation and ion-induced growth reduces the free energy barrier of formation. On the other hand, low neutral density at farther distances increases the free energy barrier, reducing the ion-induced growth. This experimental approach with a control on growth and decay processes can be used to study ion-molecule association reactions (80) and site selective ion reaction dynamics could be followed in mixed systems in which constituents have different IEs.

5. Photoionization of Laser Ablated Molecules and Clusters

5.1. Carbonaceous Clusters

Laser ablation (81) to generate metal atoms and clusters, oxides and carbides, have had tremendous impact in the fields of astrochemistry and catalysis (82). Much of the reactivity and spectroscopy of these systems have been probed with visible and ultraviolet laser based methods which provides information mostly on the ground state neutral molecules. However both synchrotron and laboratory based methods have also been coupled to laser ablation to provide energetic and spectroscopic information for the corresponding ions. Ng describes high resolution VUV studies on a number of small metal containing species, of relevance to astrochemistry, probed via VUV pulsed field ionization-photoelectron measurements (83). Yin and Bernstein (82) have reviewed oxidation-reduction and bond activation reactions of a series of neutral metal clusters which were all investigated by fixed wavelength (10.5 eV) PIMS. Below we describe synchrotron based photoionization studies on a series of carbon and metal clusters, and their products on oxidation and reaction with hydrocarbons.

Small carbon-containing species are ubiquitous in interstellar space, around massive stars, and in our own environment as critical intermediates in flame chemistry and soot formation. They have

also been invoked as precursors of large carbon molecules including aromatic species and fullerenes, such as C₆₀. Photoionization studies of clusters provide accurate input parameters of astrochemical models simulating the formation of carbon grains and PAH's. Small carbon (C_n, n=2-15) clusters, (84) carbon nitrogen-terminated (C_nN, n=4-12) clusters (85), produced by pulsed laser vaporization of graphite in atmospheres of rare gas and N₂O respectively, and polyacetylenes $(H(C \equiv C)_n H, n=1-9)$ (86) formed via ablation of graphite in the presence of argon-seeded benzene or by ablation of silicon in neat acetylene were studied by PIMS. The comparison of computed and experimental IEs allowed for determination of the isomeric structures of the neutral clusters (linear vs cyclic) produced in the experiment. In the case of pure carbon clusters (84), for the sizes n=4-6, unquenched excited electronic states complicated the comparison between theory and experiment, however the prominence of linear structures for n= 7, 9, 11, and 13 species and cyclic C₁₀ is revealed. For the C_nN series of clusters (85), a comparison of AEs to calculated IEs suggests that similar to the bare carbon clusters, linear isomers dominate the spectra. In the case of polyacetylenes (H(C≡C)_nH) (86), clusters containing odd numbers of carbon atoms have polyacetylenic character with alternating single and triple bonds, while the even number of carbon based clusters tend to have more cumulenic type structures and ionization from these states were reflected in the measured PIE curves allowing for structural elucidation.

5.2. Silicon Systems

The thermodynamic properties of organo-silicon compounds are of paramount importance in understanding the formation of silicon-bearing nanostructures together with their precursors from the 'bottom up' in the interstellar medium, in our Solar System, and in chemical vapor deposition processes. Towards this end, structure determination of bare silicon clusters as well as reactivity studies of Si clusters generated by laser ablation with a variety of hydrocarbons and oxidizing

molecules have been performed. Generation of silicon clusters in an atmosphere of acetylene led to formation of small organo-silicon molecules of the formula SiC_2H_x (x=0, 1, 2) (87; 88). Theoretical IEs and FC factor calculations for SiC_2 , SiC_2H and SiC_2H_2 allowed for isolation of the isomers prevalent in the experiment. This is nicely illustrated in the case of SiC_2H_2 where Figure 5(a) shows theoretical PIE curves for five different isomers superimposed with the experimental data and it is clear that only cyclic SiC_2H_2 (2B_2) isomer is present. Even though three isomers have IEs overlapping with the experimental data, the FC analysis of the experimental PIE curves points to the right isomer.

SiO₂ in the solid state is ubiquitous on earth as silica in its various amorphous and crystalline forms and in space in the form of interstellar dust, while there is a paucity of its spectroscopy in the gas phase. Gas phase SiO₂ was observed in the laser ablation of silicon in a CO₂ molecular beam and studied with VUV-PIMS (89). PIE curves were recorded for SiO and SiO₂ and IE estimates were revealed from such measurements. A state-to-state IE of 12.60 ± 0.05 eV was recorded by fitting two prominent peaks in the PIE curve for the following process: $^{1}\Sigma$ O-Si-O \rightarrow $^{2}\Pi_{g}$ [O-Si-O]⁺. Electronic structure calculations aided in the interpretation of the photoionization process and allowed for identification of the symmetric stretch of $^{2}\Pi_{g}$ [O-Si-O]⁺ which was observed in the PIE spectrum to be 0.11 eV (890 cm⁻¹) above the ground state of the cation and agreed with the 892 cm⁻¹ symmetric stretch frequency calculated at the CCSD(T)/aug-cc-pVTZ level.

Since there has been controversy in the measurements of the IE of the silicon dimer and no direct measurements of the higher clusters, the IEs of Si_n (n=1-7) using VUV-PIMS coupled to laser ablation of silicon in a supersonic molecular beam were recorded (90). The extracted AIEs obtained by fitting a FC simulation to the experimental PIE curves represented a significant

improvement in the literature values with error bars ranging from ± 0.05 to 0.1 eV depending on cluster size. In the case of the silicon dimer, a feature in the PIE spectra, coupled to a FC simulation allowed for the observation of the ionization of the X $^3\Sigma_g^-$ neutral state to the $a^2\Pi_u$ excited state of the dimer cation. Time-dependent density functional theory calculations coupled to observed trends in the measured PIE spectra revealed the energies of electronically excited cationic states in some of these clusters.

5.3. Transition Metal Oxides and Carbides

Transition metal oxides and carbides have been the subject of numerous experimental and theoretical studies due to their relevance to active species in catalytic processes such as the oxidation of CH₄ and CO (91). Platinum is of particular interest as it is extensively used in the petroleum industry as a catalyst for hydrocarbon dehydrogenation, cracking, isomerization and aromatization. These molecules are challenging to describe with electronic structure theory due to the number of unpaired electrons and many low-lying electronic states (often having different electron spin states), as well as the difficulty of accurately treating electron correlation and relativistic effects. A major goal of photoionization experiments in this case is to determine accurate bond strengths for neutral metal oxides and carbides. As the IE's of the metal atoms are precisely known, measuring the IE of the metal oxides determines the metal-oxide bond strength in the neutral. Metz et al. previously determined IEs of FeO and CuO (92), systems which are relevant in corrosion processes, and extended them to platinum compounds because PtO+ can directly convert methane to methanol (93), while Pt⁺ reacts with methane to produce PtCH₂⁺, also at thermal energies (94). Citir et al. determined IE(PtC)=9.45±0.05 eV, IE(PtO)=10.0±0.1 eV and IE(PtO₂)=11.35±0.05 eV (95). The IE values for the oxides are ~0.5 eV higher than those derived from the existing neutral and ion bond strengths (96; 97). This independent measurement provides

an excellent check on the accuracy of the measured bond strengths. A range of metal oxidation states and variety of oxygen binding motifs were explored via the photoionization of TaO_x (x=3-6) clusters (98) and comparing their PIE's to FC simulations. TaO_3 and TaO_3^+ have pseudo- C_{3v} symmetry, with Jahn-Teller distortions along two degenerate antisymmetric Ta-O stretches. TaO_4 and TaO_4^+ maintains a superoxide form where O_2 is electrostatically bound to TaO_2^+ . In contrast, a $(\eta^1$ - $O_2)$ TaO_3 dioxygen structure was recovered for TaO_5 while for TaO_6 , $(\eta^2$ - $O_2)(\eta^1$ - $O_2)$ TaO_2 structure is favored.

Although several third row transition metal cations M⁺, including platinum, react with methane at thermal energies to produce MCH₂⁺ + H₂, this reaction has not been observed for neutral metal atoms. The photoionization studies observe the H-Pt-CH₃ reaction intermediate formed by Pt insertion into the C-H bond in methane. The intermediate has been predicted by calculations, and postulated to be responsible for the high termolecular rate for Pt + CH₄ clustering (99-101). Combination of the photoionization results with guided ion beam measurements of the Pt+ + CH4 potential energy surface (102) allowed for a derivation of a potential energy surface for the neutral reaction (Figure 5(b)). $IE(H-Pt-CH_3)=8.89\pm0.03 \text{ eV}$ and $IE(PtCH_2)=8.78\pm0.03 \text{ eV}$ were determined for species formed by reacting laser-ablated Pt with methane (103). This reveals that formation of H-Pt-CH₃ is 164 kJ/mol exoergic, and the overall reaction is only 9 kJ/mol endoergic. In addition, DFT calculations show that spin-orbit coupling is so strong that there is no barrier to C-H insertion. These studies show that the PES for reactions of Pt and Pt⁺ with CH₄ are very similar, with the cation intermediates and products only ~20 kJ/mol more stable, relative to reactants. So, in activation of methane by Pt, covalent forces dominate and charge matters very little.

6. Future Directions

New directions in enhancing sensitivity in detecting isomers, and complex molecules beyond just IE and fragmentation mechanisms are being taken by coupling VUV ionization to multiplexing detection of ions and electrons with coincident imaging and integration of separation techniques such as gas chromatography. The synchrotron groups at Hefei (104), SLS (105) (106), and Soleil (107) have developed new versions of ion-electron coincidence detection. As documented throughout this review, the ms-TPES method shows tremendous potential for quantitative, timeor space-resolved, isomer specific analysis of complex systems. The photoelectron spectrum contains vibrational signatures of the final quantum state and therefore provides higher selectivity compared to VUV-PIMS. It remains to be seen whether the reduced resolution in mass spectra and lower S/N observed in these coincidence measurements are outweighed by the superior ability of mass selected threshold photoelectron spectra to identify isomers. Notwithstanding, these groups have demonstrated that these methods are very useful to study complex chemical systems. Hemberger et al. have studied unimolecular reaction mechanisms of an Arduengo-type (stable Nbased heterocyclic) carbene (108; 109). Nahon, Garcia and co-workers at synchrotron Soleil have used ms-TPES, to study the spectroscopy of gas phase molecules generated in molecular, radical or aerosol beams. One elegant example is that of a model PAH (coronene) and its clusters (dimer and trimer), whose ms-TPES revealed a number of autoionization resonances near the first adiabatic ionization threshold relaxing to both vibrationally and electronically excited cationic states (110). Another direction that Nahon and co-workers are pushing is their combination of VUV photoionization with photoelectron circular dichroism (PECD) to investigate chirality in biomolecules such as alanine (111). The PECD measurements at different photon energies allowed the detection of various conformers present in the beam, promising both isomer and conformer selectivity.

While Mass Analyzed Threshold Ionization (112) and two color laser + VUV synchrotron photoionization (113; 114) experiments have been demonstrated at a synchrotron, the enhanced sensitivity of lab based laser techniques as reviewed by Ng (83) will continue to dominate the field of state to state ionization dynamics and spectroscopy. Notwithstanding, the coupling of high repetition rate (kHz) lasers to synchrotron based PIMS (115) did allow for identification of molecular details in complex systems such as mineral organic associations in soil (116) and melanin in bird feathers (117). While synchrotron experiments have superior S/N when sampling occurs from continuous sources such as flames and flow reactors, tunable VUV pulsed lasers with more photons accessible per pulse, ties well with pulsed sources. New laser sources at higher repetition rates have been deployed for high resolution VUV spectroscopy in the gas phase (118) while fiber based frequency comb lasers (119) to generate synchrotron like tunable VUV light should find increased application in the future. Free electron lasers (120) are burgeoning all over the world, heralding novel applications in spectroscopy, scattering and imaging of chemical systems. A dedicated VUV free electron laser (121) for chemical applications particularly for combustion and energy science is being commissioned in Dalian China, and should produce first results within this decade. Synchrotrons are not lagging behind, and new diffraction limited light sources (122) with transverse coherence will allow interrogation of complex chemical systems using interferometry and correlation spectroscopy.

An exciting new direction is the coupling of gas chromatography to VUV-PIMS (123) to interrogate complex aerosol chemistry as demonstrated by Wilson and co-workers (124). A chromatography column separates complex molecules both by volatility and polarity and are subsequently ionized by VUV. This combination of separation and threshold ionization leads to elucidation of very complex chemical processes such as heterogeneous oxidation chemistry by

radicals (OH) on atmospheric aerosols (125) and identification of emissions from gasoline and diesel engines (126; 127). The coupling of ion trap mass spectrometers to synchrotrons to perform VUV and X-ray spectroscopy of mass selected ions will provide valuable information on the electronic structure of complex chemical systems (128). A glimpse of future possibilities in spectroscopy and dynamics are revealed in the photoelectron spectroscopy of size-selected (MoO₃)_n+ clusters, (129), VUV multiphoton ionization of size-selected silicon cluster ions (130), VUV photoionization of peptides (131) and proteins (132), VUV photofragmentation (133) and electron detachment spectroscopy (134; 135) of small peptides. While the bias has been towards gas phase ions, the possibilities of selecting ions out of solution via electrospray and then interrogating them for spectroscopy bodes well for solution phase chemistry.

Finally, theory will continue to play a critical role in advancing new directions in VUV photoionization studies of complex systems. A theoretically determined absolute photoionization cross section will be very valuable for quantification of complex molecules and transient species, which are prevalent in combustion and environmental chemistry mechanisms. Another direction would be to develop electronic structure methods for larger molecular systems, complexes and clusters.

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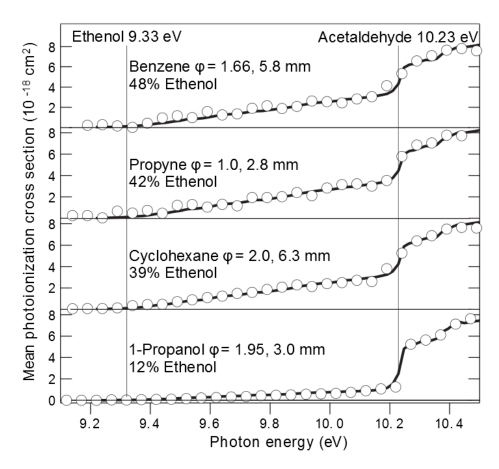


Figure 1. Photoionization efficiency curves taken for m/z = 44 ion sampled from four representative flames. The fuel/oxygen equivalence ratio for each flame is denoted as ϕ in the figure. Photoion signals (circles) taken at the stated distance from the burner (indicated in mm in the figure) have been background-corrected and normalized by the measured photon flux. Signals are then scaled to compute mean photoionization cross sections (lines) for hypothetical mixtures of ethanol and acetaldehyde that give best fits to the data. Ionization energies for ethanol (CH₂=CHOH) and acetaldehyde (CH₃CHO) are indicated by the thin vertical lines. From (14). Reprinted with permission from AAAS.

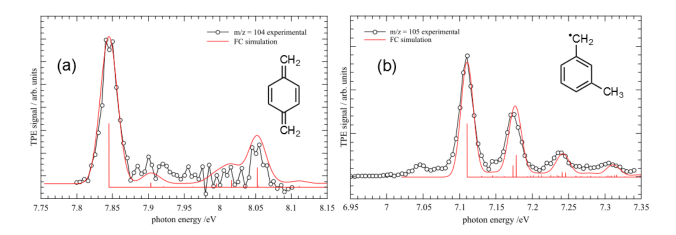


Figure 2. Threshold photoelectron spectra for (a) the m/z=105 species formed in the pyrolysis of m-xylyl bromide compared with a Franck-Condon simulation for the m-xylyl radical and (b) the m/z=104 product from m-xylyl bromide pyrolysis, with comparison to a FC simulation for p-xylylene. Reprinted with permission from (30). Copyright 2013 American Chemical Society.

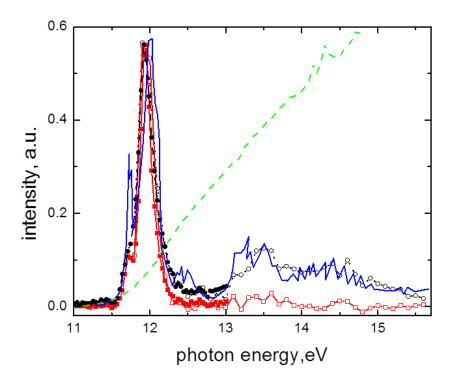


Figure 3. Photoionization efficiency curves of the water tetramer $(H_2O)_4^+$ (red filled (30 meV step size) and unfilled squares (100 meV step size)) and the protonated tetramer $(H_2O)_4H^+$ (black filled (30 meV step size) and unfilled circles (100 meV step size)) formed in a high pressure (7 bar Ar) expansion. The fluorescence spectrum of Ar_n clusters $\langle n=20 \rangle$, formed via VUV excitation, taken from ref (56) is shown as a blue solid line. The green dashed line is the photoionization efficiency curve of protonated tetramer, $(H_2O)_4H^+$ formed in a low pressure (1.3 bar Ar) expansion. Reprinted with permission from (55). Copyright 2012 American Chemical Society.

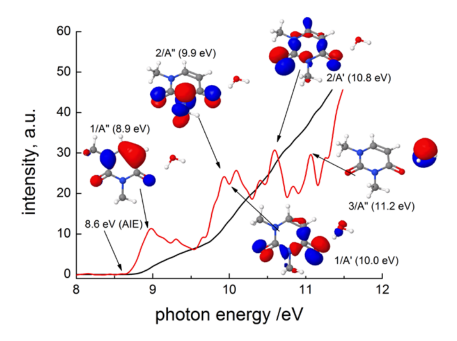


Figure 4. Photoionization efficiency curve (black) of dimethyluracil-D₂O complex and its derivative (red), observed using 8–12 eV photons. The derivative plot reveals multiple ionized states derived by removing the electron from different molecular orbitals (theoretically calculated). Black arrows point toward the calculated ionization energies. Reprinted with permission from (71). Copyright 2013 American Chemical Society.

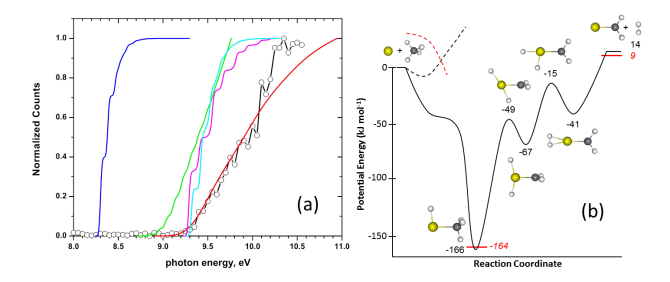


Figure 5. (a) Experimental (open symbols and black line) and computed (solid colored lines) PIE curves of the low lying individual isomers for $SiC_2H_2^+$, blue line H_2CCSi , green line HSiCCH, gray line SiC_2H_2 cyclic (2A_1), magenta line H_2SiCC linear, red line SiC_2H_2 cyclic (2B_2), (b) Potential energy surface for Pt + CH₄ \rightarrow PtCH₂ + H₂ reaction. DFT results dashed, spin-orbit DFT/aug-cc-pVTZ results in solid black, experimental energies in red. Reprinted with permission from (103) Copyright 2012 John Wiley and Sons, Inc.

Acronyms and definitions:

VUV: Vacuum Ultraviolet.

PT: Proton Transfer.

PIMS: Photoionization Mass Spectrometry.

H-bond: Hydrogen bond. IE: Ionization Energy.

PIE: (Photoionization Efficiency) Integrated ion intensity as a function of photon energy, normalized to incident photon intensity.

AE (Appearance Energy): Minimum photon energy required to detect a fragment ion.

AIE: (Adiabatic Ionization Energy) Minimum energy required to remove an electron from a neutral molecule in its vibrational ground state.

VIE: (Vertical Ionization Energy) Ionization energy corresponds to vertical transition when maximum overlap of vibrational ground and excited state wavefunctions is achieved.

PEPICO: (Photoelectron Photoion Coincidence Spectroscopy) Mass spectrometric technique to investigate energy selected ions which provides dissociation rates and onsets.

iPEPICO: (Imaging Photoelectron Photoion Coincidence Spectroscopy) Simultaneous mass spectrometry and velocity map imaging to investigate energy selected ions and electrons typically produced by VUV ionization.

Franck Condon (FC): Assuming the fixed nuclear approximation, square of overlap integral between the vibrational wavefunctions involved in a transition.

ms-TPES: (mass-selected Threshold Photoelectron Spectroscopy) Using tunable VUV, photoions from a specific species are collected in coincidence with near zero kinetic energy electrons.