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## Coincidence study of core-ionized adamantane: site-sensitivity within a carbon cage?

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We investigate the fragmentation dynamics of adamantane dications produced after core-ionization at the carbon edge followed by Auger decay. The combination of high-resolution electron spectroscopy, energy-resolved electron-ion multi-coincidence spectroscopy and different theoretical models allows us to give a complete characterization of the processes involved after ionization. We show that energy- and site-sensitivity is observed even for a highly-symmetric molecule that lacks any unique atomic site.

## 1 Introduction

Diamondoids have been of great interest in recent years due to their role in many applications<sup>1</sup> such as in nanotechnology,<sup>2</sup> in medicine,<sup>3,4</sup> and in astrochemistry.<sup>5,6</sup> In space, diamondoids are expected to be abundant due to their high stability but after comparison with laboratory measurements based on infrared spectroscopy,<sup>7</sup> astronomical observations show a deficiency of diamondoids in the interstellar medium which to date is not completely understood.<sup>8</sup> Adamantane ( $C_{10}H_{16}$ ) is the smallest diamondoid molecule with a carbon cage formed by  $C(sp^3)-C(sp^3)$ -hybridized bonds and fully terminated by hydrogen atoms. The molecule comprises two different carbon sites  $CH$

and  $CH_2$ , so-called methanetriyl and methylene groups, respectively. Previously, valence dissociative ionization of the adamantane has been studied using VUV radiation,<sup>9</sup> XUV femtosecond pulses,<sup>10,11</sup> electron ionization<sup>12,13</sup> as well as other spectroscopic works.<sup>14,15</sup> These studies reveal dissociation *via* a number of parallel channels which all start with an opening of the carbon cage and hydrogen migration indicating that the low stability of adamantane after excitation/ionization could explain its deficiency in astronomical observations. However, little is known about X-ray dissociative ionization of adamantane and diamondoids in general. It is known that X-rays travelling through the interstellar medium ionize the atoms, molecules and grains in the medium.<sup>16,17</sup> The interaction of diamondoids with X-rays is therefore relevant to the chemistry of interstellar medium<sup>18–21</sup> and could partially explain the lack of diamondoids in space.

The only studies of carbon K-edge spectra we are aware of, for adamantane and other diamondoids, are the absorption spectra of Willey *et al.*<sup>22,23</sup> In addition, the Auger electron spectra of diamondoids (bulk) have been measured and rationalized by calculations on adamantane derivatives by Endo *et al.*<sup>24</sup> The theoretical results could resolve different contributions both regarding the Auger process ( $1s-2p2p$ ,  $1s-2s2p$ ,  $1s-2s2s$ ) and the chemical environment ( $C-C$  or different  $C-H_n$  atoms), raising the possibility of finding site-sensitive fragmentation upon core-hole excitation. Thus we have studied the dissociation of adamantane molecules in the gas phase after C-1s core-ionization using synchrotron X-ray radiation and explore site-sensitive dissociation. In fact, K-shell ionization by synchrotron radiation has been extensively used to probe the chemical environment of many molecular and cluster systems<sup>25–33</sup> even though site-specific dissociation is often considered as weak<sup>34</sup> and that fragmentation dynamics is mostly driven by internal energy.<sup>35,36</sup> The fact that adamantane

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is a saturated hydrocarbon implies that the detection of site-selective dissociation (using the direct photoelectron) is highly challenging due to small energy difference between the two different carbon sites (similar chemical environment) and the effect is expected to be rather small. Nevertheless, multi-coincidence tools using Auger electron and ion spectroscopy can reveal such processes as we will show in the following.

In this work, we present the results of electron-ion spectroscopy of the adamantane dication after core-ionization at the carbon edge using X-ray photons. The first part consists of high-resolution spectroscopic measurements such as C-edge X-ray photoelectron spectrum (XPS), near edge X-ray absorption fine structure (NEXAFS), normal and resonant Auger electron spectroscopy (AES and RAES). These studies are used to determine the conditions and interpret the results of correlated photoion and Auger electron spectroscopy (AEPIPICO). This enables the characterization of the charged products of interaction (identification and energetics). Combination of state-of-the-art computational techniques is required to treat such complex system and the variety of processes occurring. We demonstrate that thanks to coincidence measurements and good understanding of the electronic structure of ionized adamantane we can reveal both energy- and site-sensitivity of photodissociation, even for such symmetric molecules containing carbon as the only heavy element.

## 2 Methods

### 2.1 Experimental

The experiments were carried out at the soft X-ray beamline PLEIADES at Synchrotron SOLEIL.<sup>37</sup> The soft X-rays were generated by an Apple II HU 80 permanent magnet undulator and monochromatized using the modified Petersen plane grating monochromator in combination with a high-flux 600 lines per mm grating. For the present experiment, the photon energy was calibrated according to the maximum position of the CO<sub>2</sub> C-1s → π\* absorption peak.<sup>38</sup>

The electron spectroscopy measurements were performed using a newly developed heated gas cell, in combination with a wide-angle lens VG-Scientia R4000 electron energy analyzer.<sup>39</sup> High purity commercial adamantane powder (99%, Sigma-Aldrich) was used without further purification except vacuum pumping. The sample was placed in an external container and introduced into the gas cell through a heated injection gas line. The gas cell was heated at 54 degrees Celsius. During the measurements, the pressure in the Scientia chamber (outside the gas cell) was  $7.1 \times 10^{-7}$  mbar (the base pressure was  $4.3 \times 10^{-8}$  mbar). The binding energy scale of the adamantane XPS spectrum was calibrated according to the XPS spectrum of CO<sub>2</sub> and the reference C-1s IP value reported by Myrseth *et al.*<sup>40</sup> The CO<sub>2</sub> was introduced into the gas cell under the same experimental conditions, through the same injection gas line. The kinetic energy scale of the adamantane Auger spectra was calibrated at the same time according to the measured CO<sub>2</sub> Auger spectrum and the position of reference lines reported by

Modeman *et al.*<sup>41</sup> The overall uncertainty of binding and kinetic energy scales due to the calibration is estimated to be up to 0.1 eV. The adamantane electron spectra were recorded at the photon energy of 350 eV for the XPS and AES, 287 and 287.6 eV for the RAES, with an overall resolution of 55 meV.

The AEPIPICO spectra were recorded using the EPICEA setup,<sup>32,37</sup> which consists of a double toroidal electron analyzer (DTA)<sup>42</sup> and a 3D focusing ion TOF spectrometer. Electrons emitted at an angle of 54.7° enter the DTA, where they are retarded to a predetermined pass energy (E<sub>p</sub>). The detection electron kinetic energy window is about 12% of the defined E<sub>p</sub>, whereas the energy resolution is about 1% of the E<sub>p</sub>.<sup>43</sup> In the present experiments, a pass energy of 250 eV was used, which leads to an analyzer resolution of about 2.5 eV and a detection range of ±15 eV. The retardation voltage and DTA lenses were then adjusted to detect Auger electrons with a kinetic energy corresponding to a binding energy (BE) centered at 20 eV in the case of resonant Auger electron spectra and 40 eV in the case of normal Auger electron spectra. After exiting the DTA, the electrons are recorded by a delay-line position sensitive detector (DLD40, Roentdek GmbH). The distance of the electron position from the center is inversely proportional to its kinetic energy. The calibration of the kinetic energy scale was performed by recording radius-positions of Xe 5p and Xe 5s IPs as a function of the photon energy, and by using a previously proposed empirical formula to transform the detector radius-position (mm) to the electron kinetic energy (eV).<sup>43</sup>

For the EPICEA experiment, the adamantane is introduced into the vacuum chamber through a heated injection gas line and a heated needle, in a crossed-beam experimental arrangement. The base pressure was  $4.2 \times 10^{-8}$  mbar. The interaction region is placed in a zero-electric-field space between two extraction grids. The detection of an electron triggers a pulsed field that accelerates all ions toward a hexagonal delay-line detector (HEX75, Roentdek GmbH) placed behind the TOF drift tube. The positions on the detector and time of flight of all extracted ions are recorded and make part of the same coincident “event” marked by the triggering electron. Additionally, a pulse generator (Stanford DG645) was used to produce random triggers, which extract ions present in the interaction without a coincident electron event, thus allowing to analyze the background ion signal recorded due to false coincidences. False coincidences were taken into account using the statistical procedure reported by Prümper and Ueda.<sup>44</sup> The NEXAFS spectra recorded on the EPICEA setup were obtained as a total photoion yield measured in a continuous ion extraction field.

### 2.2 Theoretical

**2.2.1 XPS spectra.** In a series of papers<sup>45–48</sup> we have proposed a quantum approach to calculate total and differential cross sections for the electron impact ionization of complex molecules (formic acid, tetrahydrofuran, pyrimidine...). By integrating the most differential cross sections (here the triply differential cross sections), we can easily retrieve the doubly differential (DDCS), simply differential and total cross sections. In order to obtain the XPS intensities one computes the zero momentum transfer limit ( $q \simeq 0$ ) of the DDCS. The calculation is performed

for a given energy deposit provided by the electron projectile that corresponds, under these special kinematical conditions, to the XPS photon energy (e.g.  $h\nu = 350$  eV). Finally, the modeling of an experiment using molecules in gas phase necessitates that the DDCS results must be averaged out over the molecular orientations.

The theoretical framework is based on the First Born Approximation (FBA) and the ejected electron is described by a distorted wave (a complete theoretical description of the model is given in ref. 48). The molecular wavefunction was computed at the HF/6-31G(d) level over the optimized geometry at the B3LYP/6-31G(d) level, and using Gaussian16 code.<sup>49</sup> Such wave function is then converted to a single center wavefunction, thereby reducing the complexity of this multi-atom system.

**2.2.2 Auger spectra.** Auger electron spectra of adamantane were obtained with a protocol that uses configuration interaction (CI) wavefunctions and the one-centre approximation.<sup>50–52</sup> A convolution with Gaussian functions with a full width at half maximum (FWHM) of 1 eV provided the resulting theoretical spectra. The protocol is very similar to the one described in more detail in ref. 53 and further details of the approach as well as references to them can be found in ref. 54.

Hartree–Fock (HF) orbitals of the adamantane molecule obtained with the SV(P) basis set<sup>55</sup> were extended by the space of atomic valence orbitals according to ref. 56. Localized C-1s core orbitals were obtained with a Mulliken localization of the ten HF-orbitals with lowest orbital energies. The core-ionized states were described with CI calculations with one hole in the core orbital of interest and additionally one excitation of an electron from the occupied valence orbitals to one atomic virtual orbital while the final state energies and wavefunctions were obtained with a CI calculation with all configurations that result from two holes in the occupied valence orbitals of the ground state adamantane molecule.

**2.2.3 Dication photodissociation.** Exploration of the potential energy surface in doubly ionized adamantane was carried out at the B3LYP/6-31G(d) level of theory. We have located relevant minima and the corresponding transition states connecting them. The nature of the stationary points (both minima and first order transition states) was verified by computing the harmonic frequencies, and the connection of the transition states by running the intrinsic reaction coordinate. Relative energies were corrected with the zero-point energy, ZPE. These calculations were also carried out using Gaussian16 code.<sup>49</sup>

### 3 Results and discussion

#### 3.1 C-edge X-ray photoelectron spectroscopy

We used X-ray photoelectron spectroscopy (XPS) to probe the chemical environment of the adamantane carbon cage. Fig. 1 shows the experimental C-1s edge XPS spectrum recorded at  $h\nu = 350$  eV and where several contributions can be observed. The two main features at 289.9 eV and 290.1 eV come from the two different carbon sites (CH and CH<sub>2</sub>) and are spaced by about 200 meV. The visible tail at higher binding energies can be associated with the population of some vibrational states.

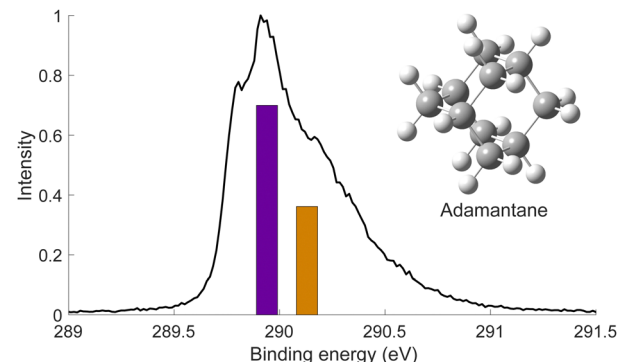


Fig. 1 C-1s XPS spectrum of adamantane recorded at  $h\nu = 350$  eV. The two bars represent the theoretical contributions of the 2 carbon sites (purple for CH<sub>2</sub> sites and orange for CH sites) and are normalized to 0.7. The structure of adamantane is depicted in inset.

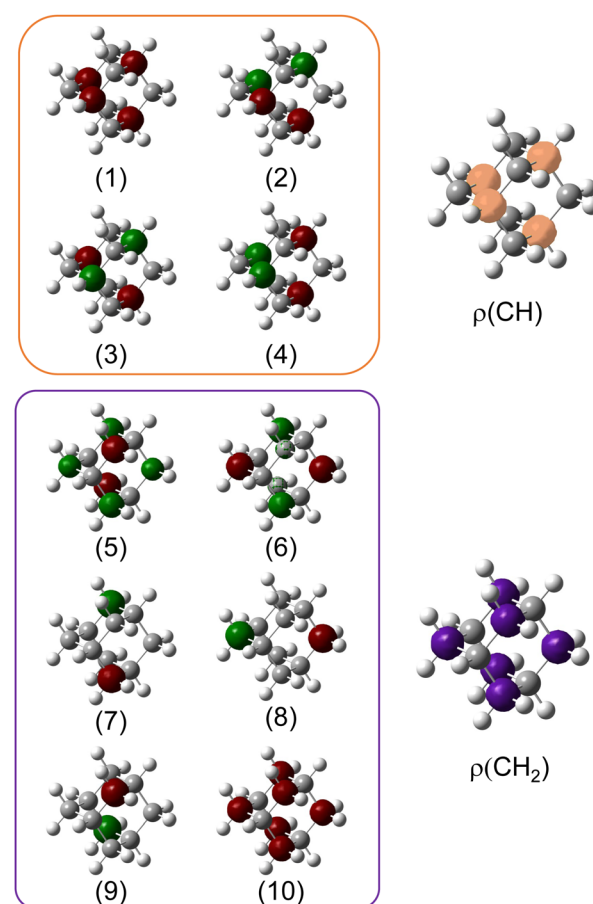


Fig. 2 Molecular orbitals (MO)  $\phi_i$  of neutral adamantane, computed at the HF/6-31G(d) level, are depicted, in red and green the two phases of the MO, and with a surface isovalue of 0.01 a.u. (1–4) correspond to 1s molecular orbitals of CH sites and (5–10) to CH<sub>2</sub> sites. The electronic density associated with the two types of carbon atoms has been computed as  $\rho(\text{CH}) = \sum_{i=1}^4 |\phi_i|^2$  and  $\rho(\text{CH}_2) = \sum_{i=5}^{10} |\phi_i|^2$  and have been depicted in orange and purple, respectively and with an isovalue of 0.001 a.u.

To compare with the experimental measurement, theoretical calculations (see Section 2.2.1) have been performed. First, they



give access to the 10 different molecular orbitals that are displayed in Fig. 2. The first four (1–4) correspond to methanetriyl groups and  $\rho(\text{CH})$  is the corresponding electronic density associated to the core orbitals in this site (see caption of Fig. 2). This shows that the electronic density is mainly located on the carbon of the initial site. Similar behaviour is observed for the ones of the methylene groups (5–10) and  $\rho(\text{CH}_2)$ . This indicates that orbitals seem to be very well localized on the carbon groups involved in the ionization and should give a well defined site-sensitivity. The Koopmans energy values of these molecular orbitals have an energy difference of  $\sim 200$  meV between the two carbon sites matching the experimental observation. The intensities of the different orbitals are calculated with a method based on the First Born Approximation (see Section 2.2.1) and are reported as bars normalized to 0.7 in Fig. 1 (purple for  $\text{CH}_2$  sites and orange for  $\text{CH}$  sites) in good agreement with the experimental measurement.

### 3.2 Auger-yield near edge X-ray absorption fine structure

Fig. 3(a) shows the photon energy dependence of Auger electron energy of adamantane near the C-1s ionization edge. We observe that below the ionization edge ( $\sim 289.7$  eV) there are two resonances at 287 eV and 287.6 eV attributed to  $\text{CH}$  and  $\text{CH}_2$  valence orbitals, respectively, and one resonance above threshold at 292 eV coming from  $\sigma^*$  states of C-C bonds (respective attributions from ref. 22 and 23). Strong post-collision interactions (PCI) are visible between 288 and 292 eV (curved features). Fig. 3(b) shows the NEXAFS spectrum where the two narrow resonances at 287 eV and 287.6 eV and a broad one at 292 eV are clearly observed same as in Fig. 3(a). For comparison, a total ion yield X-ray absorption spectrum was recorded including higher resolution (30 meV) around the narrow resonances where the different vibration modes can be observed as in the works of Willey *et al.*<sup>22,23</sup>

### 3.3 Auger electron spectra

The normal Auger electron spectrum was recorded at  $h\nu = 350$  eV photon energy well above threshold ( $\sim 289.7$  eV) in order to avoid PCI effects and is shown as a thick black line in Fig. 4. We also estimate the internal energy axis (top axis of Fig. 4) according to the equation proposed by Kukk *et al.*<sup>35</sup>  $E_{\text{int}}(\text{Ada}^{2+}) =$

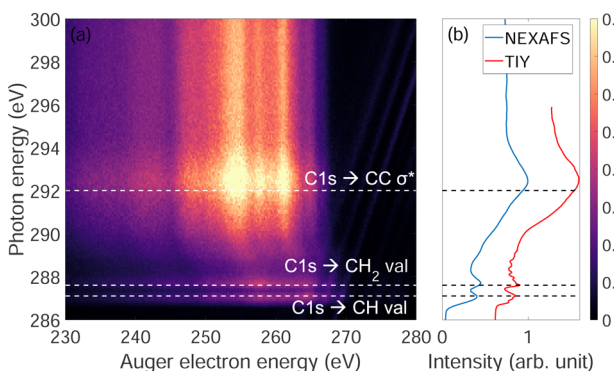


Fig. 3 (a) Two dimensional map of Auger electron kinetic energy as a function of incident photon energy near the C-1s ionization edge of adamantane. (b) Near edge X-ray absorption fine structure (NEXAFS) spectrum and the total ion yield X-ray absorption (TIY) spectrum of adamantane.

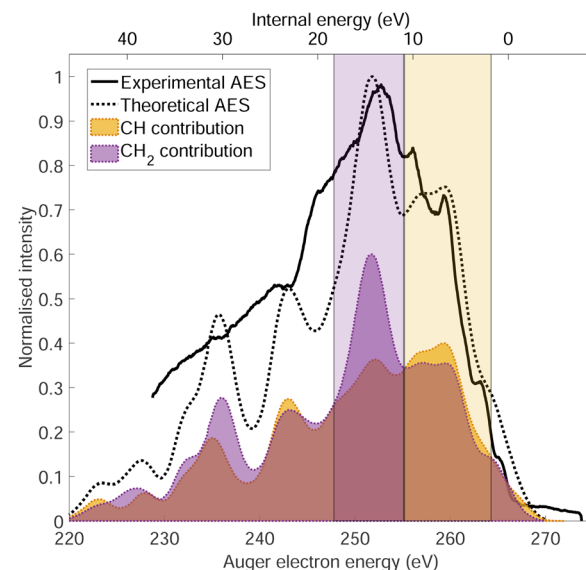


Fig. 4 Normal Auger electron spectrum of adamantane. The thick black line corresponds to the experimental measurement recorded at  $h\nu = 350$  eV, the dotted black line to the theoretical calculations and the orange and purple shaded curves to the calculated contributions from  $\text{CH}$  and  $\text{CH}_2$  sites, respectively. The two shaded rectangles (purple for  $\text{CH}_2$  and orange for  $\text{CH}$  sites) represent the two bands used later for the ion-electron data analysis.

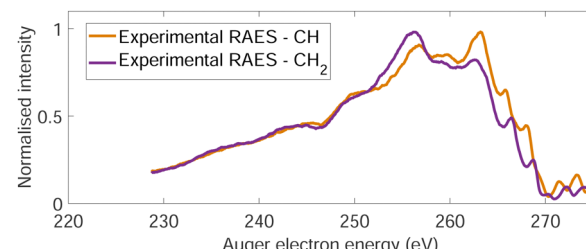


Fig. 5 Two resonant Auger electron spectra (RAES) at the  $\text{CH}$  (orange) and  $\text{CH}_2$  (purple) valence orbital resonances after photoionization of adamantane at  $h\nu = 287$  eV and  $h\nu = 287.6$  eV, respectively.

$E_b(\text{Ada}^{2+}) - V_{\text{DIT}} - \text{KE}_{\text{AES}}$  where  $E_b(\text{Ada}^{2+})$  is the binding energy of the C-1s core hole state approximated at 290 eV in our case (see Fig. 1),  $V_{\text{DIT}}$  is the double ionization threshold known at 23.9 eV according to our previous study<sup>10</sup> and  $\text{KE}_{\text{AES}}$  the kinetic energy of the Auger electrons. We can see that the internal energy starts nicely from 0 eV with a maximum around 15 eV. The theoretical Auger electron spectrum was obtained using CI energies and intensities according to the one-centre approximation (see Section 2.2.2) and is reported as a dotted black line in Fig. 4 showing a good agreement with the general features of the experimental spectrum and a previous work from Endo *et al.*<sup>24</sup> The theoretical spectrum can be decomposed in two contributions coming from the two different carbon groups ( $\text{CH}$  and  $\text{CH}_2$ ) and are shown in plain color (orange and purple, respectively) in Fig. 4. The strongest contributions are observed around 252 eV in the case of the  $\text{CH}_2$  sites (purple) and 259 eV in the case of  $\text{CH}$  sites (orange). The same predominance can be

seen in the two resonant Auger electron spectra (RAES) at the CH and CH<sub>2</sub> sites (Fig. 5) demonstrating the site-sensitivity of this two peaks. Therefore, we label two bands in the AES in purple for CH<sub>2</sub> (between 247.8 and 255.1 eV) sites and orange for CH sites (between 255.2 and 264.3 eV) visible in Fig. 4 even though these bands are not purely coming from one or another site. These bands will be used afterwards as filters in the core-ionization ion-electron spectroscopy analysis recorded at  $h\nu = 350$  eV.

### 3.4 Dication photodissociation dynamics

When the adamantane molecule is ionized with a photon energy above the C-1s edge, it undergoes normal Auger decay and produces an unstable C<sub>10</sub>H<sub>16</sub><sup>2+</sup> state. Our previous study<sup>10</sup> showed that the adamantane dication goes through barrier-free structural changes like cage-opening or hydrogen migration(s) before dissociating into different ion pairs labelled as C<sub>n</sub>H<sub>x</sub><sup>+</sup>/C<sub>m</sub>H<sub>y</sub><sup>+</sup>. In this section, we present the dissociation dynamics of the adamantane dication after C-1s ionization at a photon energy of 350 eV.

**3.4.1 AEPIPICO.** Multi-particle coincidence data of Auger electrons and photoions (AEPIPICO) of core-ionized adamantane were recorded using the EPICEA setup (see Section 2.1). Fig. 6(a) shows the different correlated ion pairs measured in coincidence with Auger electrons in the energy range 230 to 270 eV. The AEPIPICO map shows that the core-ionized

adamantane dication mostly dissociates *via* multi-body break-ups leading to the production of small fragments (C<sub>n</sub>H<sub>x</sub> with  $n = 2, 3$ ), unlike valence ionization where the dication mainly breaks *via* 2-body dissociations.<sup>10</sup>

The ion pair islands in the AEPIPICO were identified and for a quantitative analysis we calculated their branching ratios. Fig. 6(b) shows the branching ratios using a representation similar to the AEPIPICO map shown in Fig. 6(a) for direct comparison. We observe that C<sub>2</sub>H<sub>3</sub><sup>+</sup>/C<sub>3</sub>H<sub>3</sub><sup>+</sup> is the most intense channel with 15.2% of the total counts measured. The second most intense ion pair (8.4%) accounts for C<sub>3</sub>H<sub>3</sub><sup>+</sup>/C<sub>3</sub>H<sub>3</sub><sup>+</sup>. These two ions are observed in space<sup>57</sup> and can be reactive species since they are radical cations which could contribute to the formation of large hydrocarbon species in certain interstellar regions.

Interestingly, we observe quite some ion pairs containing the ion CH<sub>3</sub><sup>+</sup> (BR = 21.6%) while the neutral structure of adamantane does not contain any carbon group with 3 hydrogen atoms, therefore the abundance of methyl cations in our data is an evidence for hydrogen migration<sup>10</sup> before the dication dissociates. Methyl cations are observed in coincidence with ions containing two to seven carbon atoms. Methyl radical cations are reactive species also detected in the interstellar medium.<sup>58</sup>

It is also worth mentioning that the combined branching fractions of ion pairs that include CH<sub>3</sub><sup>+</sup>, C<sub>2</sub>H<sub>3</sub><sup>+</sup>, and C<sub>3</sub>H<sub>3</sub><sup>+</sup>,

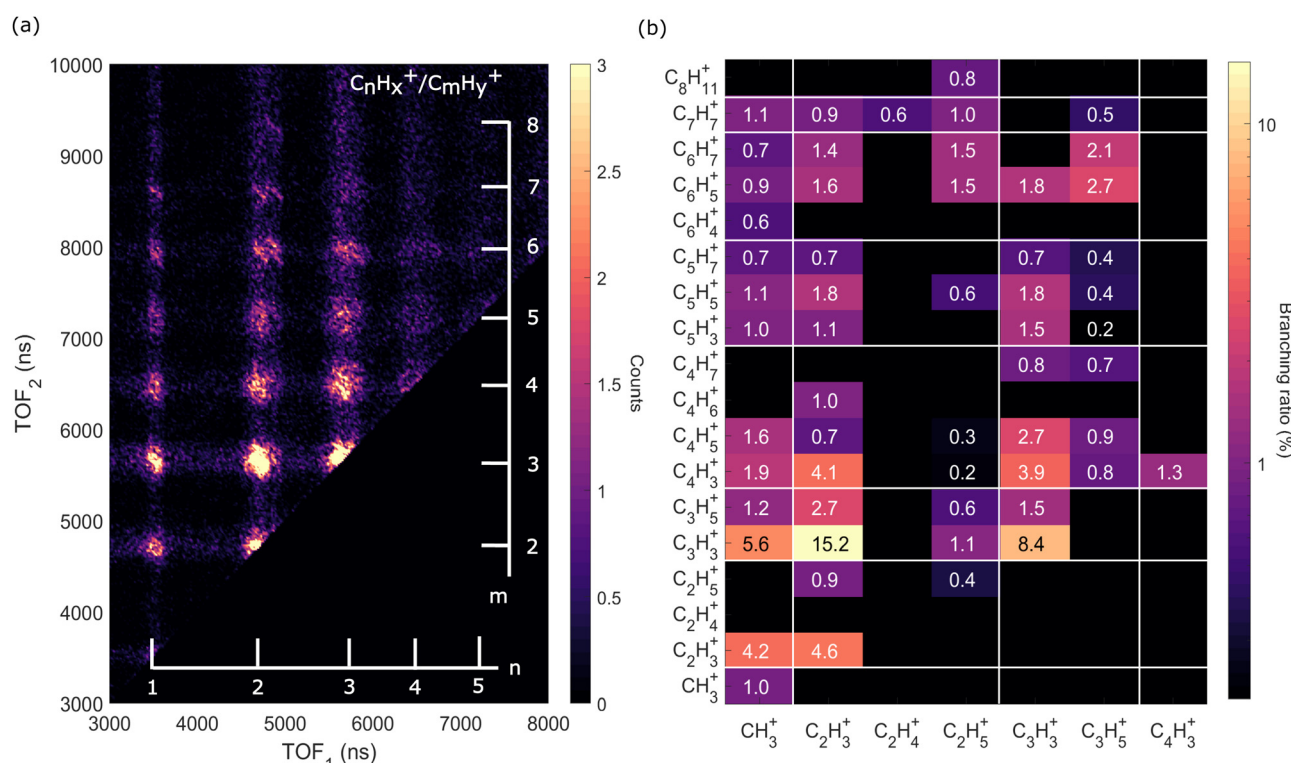


Fig. 6 Two-dimensional time-of-flight (TOF) coincidence map of ion pairs in correlation with Auger electrons with energy between 230 and 270 eV following ionization of adamantane at  $h\nu = 350$  eV, commonly referred to as AEPIPICO. The data are filtered to remove false coincidences using the statistical technique described by Prümper *et al.*<sup>44</sup> (a) All the ion pairs recorded in coincidence. (b) Heat map showing the branching ratios of ion pairs produced after photoionization of adamantane at  $h\nu = 350$  eV. White lines delimit the different carbon groups. The errors are estimated to be around 10% of the value.



account for almost 40% thus could contribute to the astrochemistry of certain regions,<sup>59</sup> for instance planetary atmospheres such as Titan's one where formation of large polycyclic aromatic hydrocarbons (PAHs) are observed.<sup>60</sup>

Another fragment,  $\text{C}_2\text{H}_5^+$ , involving at least two hydrogen migrations is observed in several channels. For instance, the correlation island of the 2-body breakup channel  $\text{C}_2\text{H}_5^+/\text{C}_8\text{H}_{11}^+$  is observed with an intensity of around 0.8% which is much smaller than the one observed for valence ionization from our previous study<sup>10</sup> since it was the main dissociation channel (BR = 23.3%). As expected, the correlation island has a slope of  $-1$  indicating a 2-body breakup with two ions ejected in opposite directions with equal momenta.<sup>61</sup> In general, the slopes of the other islands of the map are well defined and equal to  $-1$  implying multi-step dissociation process that involves neutral fragment evaporation(s) prior to Coulomb explosion into two singly charged fragments with negative momentum correlation.

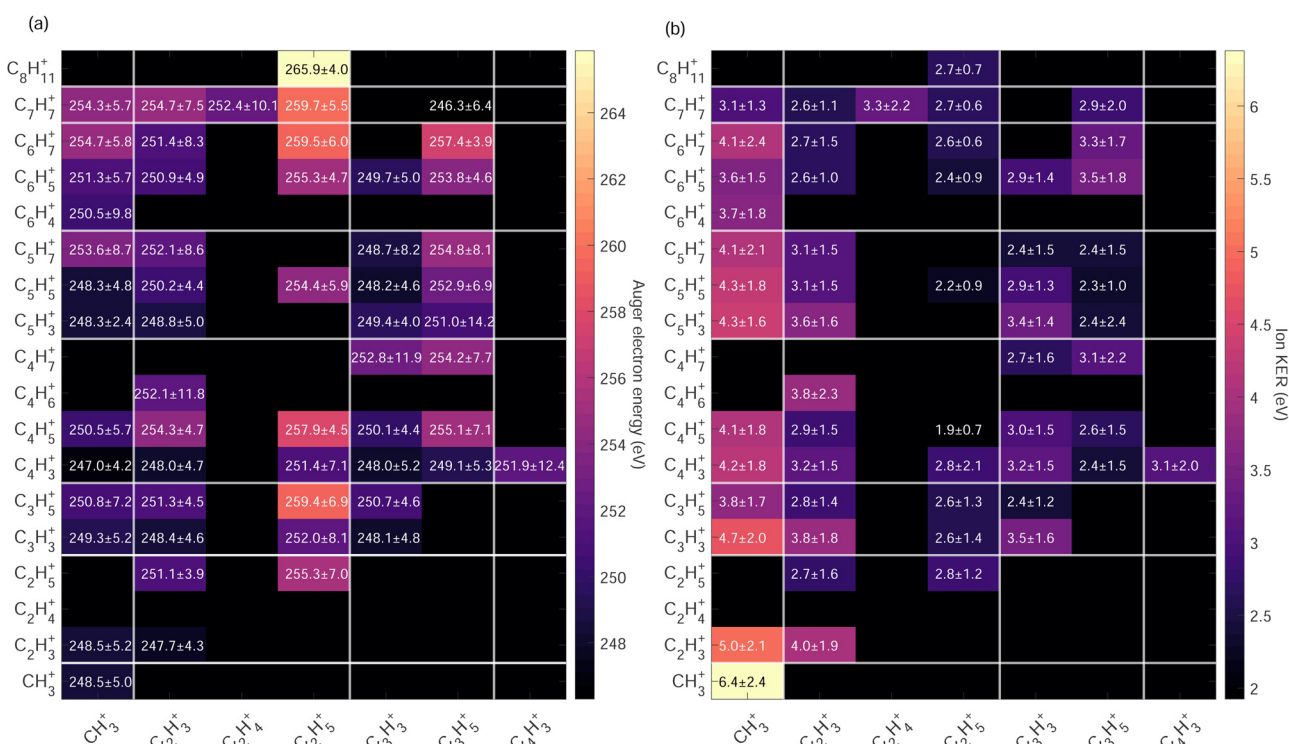
**3.4.2 Kinematics.** In order to gain an insight into the fragmentation dynamics of the dication of adamantane AEPI-PICO measurements were used to retrieve the kinetic energy release for each ion pair as well as the kinetic energy of the correlated Auger electron. Fig. 7(a) shows the weighted-average value and the standard deviation of the Auger electron kinetic energy distributions recorded in coincidence with the different ion pairs while Fig. 7(b) shows the average value and the standard deviation of the total ion kinetic energy released

(KER) distributions of the ion pairs. For the most part, ion pairs that have high energy Auger electrons have lower KER values.

High Auger electron kinetic energies indicate low binding energies of the states *i.e.* the first HOMOs – valence shells were involved in the Auger cascade. After such a Auger decay, the dication is left with low internal energy. The fragmentation dynamics of the dication depends on the internal energy and energy barrier of the potential energy surfaces. In Fig. 7(a) we observe that within the same  $\text{C}_x/\text{C}_y$  groups ion pairs with more hydrogen atoms have higher Auger electron kinetic energies indicating a strong correlation between the internal energy of the dication upon Auger decay and hydrogen migration/dissociation before fragmentation.

The highest Auger electron kinetic energies are observed for ion pairs containing the fragment  $\text{C}_2\text{H}_5^+$ . The maximum is observed for the channel  $\text{C}_2\text{H}_5^+/\text{C}_8\text{H}_{11}^+$  with a weighted average value of 265.9 eV and a standard deviation of 4.0 eV, denoted as  $265.9 \pm 4.0$  eV. The associated ion kinetic energy release for this channel is  $2.7 \pm 0.7$  eV. The dominant ion pair  $\text{C}_2\text{H}_3^+/\text{C}_3\text{H}_3^+$  has a low Auger electron kinetic energy of  $248.4 \pm 4.6$  eV and ion kinetic energy release of  $3.8 \pm 1.8$  eV.

We compare the calculated potential energy surface (PES) (Fig. 8) to the experimental results (Fig. 9). We have selected the  $\text{C}_2\text{H}_5^+/\text{C}_8\text{H}_{11}^+$  (blue) and  $\text{C}_2\text{H}_3^+/\text{C}_3\text{H}_3^+$  (green) ion pairs to exemplify 2-body and multi-body breakup dynamics in adamantane dication respectively.



**Fig. 7** (a) Heat map showing the weighted-average and standard deviation of Auger electron kinetic energy (eV) measured in coincidence with the ion pairs produced after photoionization of adamantane molecules at  $h\nu = 350$  eV. White lines delimit the different carbon groups. (b) Heat map showing the weighted-average and standard deviation of kinetic energy released (eV) of ion pairs produced after photoionization of adamantane molecules at  $h\nu = 350$  eV. White lines delimit the different carbon groups.



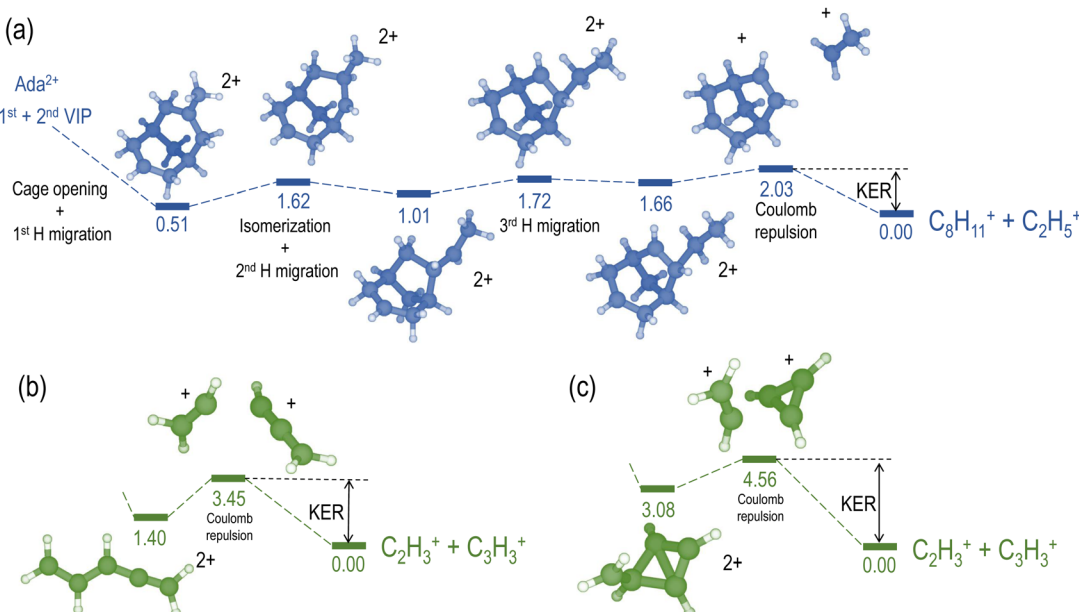


Fig. 8 Potential energy surfaces (DFT-B3LYP/6-31G(d) level) for the channels  $\text{C}_2\text{H}_5^+/\text{C}_8\text{H}_{11}^+$  (a) and  $\text{C}_2\text{H}_3^+/\text{C}_3\text{H}_3^+$  (b) and (c). Relative energies in eV corrected with ZPE and referred to the corresponding exit channel.

Fig. 8(a) shows the PES of the channel  $\text{C}_2\text{H}_5^+/\text{C}_8\text{H}_{11}^+$ . This ion pair is formed by a barrier-less cage-opening, followed by three hydrogen migrations and then finally a 2-body breakup. In Fig. 9(a), we present the coincidence map of the ion KER vs. AES for the two selected ion pairs. For  $\text{C}_2\text{H}_5^+/\text{C}_8\text{H}_{11}^+$ , we see a sharp island (in blue) indicating narrow distributions for AES

and ion KER with a well defined single peak. The theoretical ion KER can be estimated as the difference between the energy levels of the exit channel and the last transition state in the PES, because it corresponds to the Coulomb repulsion.<sup>62,63</sup> In this case, the theoretical ion KER is 2.03 eV (vertical dashed blue line) which lies within the experimental distribution ( $2.7 \pm 0.7$  eV) shown in Fig. 9(b). Fig. 9(c) shows that AES of  $\text{C}_2\text{H}_5^+/\text{C}_8\text{H}_{11}^+$  (in blue) is in the range 264–272 eV, which corresponds to a low internal energy distribution spreading from 0 to  $\sim 3$  eV. The negative part of the internal energy can be due to approximation of the binding energy  $E_b(\text{Ada}^{2+})$  (vibrational states) and calibration uncertainties at the edges of the DTA detector.

Fig. 8(b) and (c) show the PES of two different fragmentation pathways producing the  $\text{C}_2\text{H}_3^+/\text{C}_3\text{H}_3^+$  ion pair. Pathway (b) assumes the fragmentation of a linear  $\text{C}_5\text{H}_6^{2+}$  and pathway (c) a more complex cyclic structure leading to the formation of the cyclopropenyl cation. Both kind of structures can be produced assuming a high internal excitation energy after Auger decay. The coincidence map of ion KER vs. AES in Fig. 9(a) shows a broad distribution (in green) for  $\text{C}_2\text{H}_3^+/\text{C}_3\text{H}_3^+$ . Unlike  $\text{C}_2\text{H}_5^+/\text{C}_8\text{H}_{11}^+$ , for this ion pair there are several contributions in the AES and ion KER distributions indicating the presence of multiple fragmentation pathways. The maximum intensity in Fig. 9(b) is observed for KER around 3 eV but widely spread from 0 to almost 9 eV. The theoretical ion KER from the PES exploration gives two values of 3.45 and 4.56 eV (Fig. 8(b) and (c)) which are reported as vertical dashed green lines in Fig. 9(b) and shows that the broad distribution come from the fact that various pathways lead to the same final exit channel. Then, in Fig. 9(c) the AES distribution ranges from 240 to 260 eV and corresponds to a rather high and broad internal energy distribution spanning from 6 to 26 eV which also partly explain the broad feature for the KER distribution (up to 15 eV of vibrational energy left in the fragments).

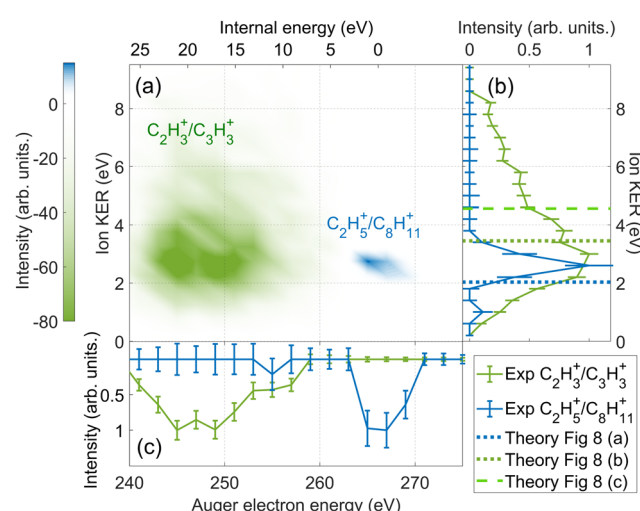


Fig. 9 The difference in energy-resolved AEPIPICO data for the  $\text{C}_2\text{H}_3^+/\text{C}_3\text{H}_3^+$  (green) and the  $\text{C}_2\text{H}_5^+/\text{C}_8\text{H}_{11}^+$  (blue) ion pairs. (a) 2D map showing the energy correlation of the Auger electron energy with the total ion kinetic energy released (KER) for the selected ion pairs. The intensities of the  $\text{C}_2\text{H}_3^+/\text{C}_3\text{H}_3^+$  events is set to be negative and the  $\text{C}_2\text{H}_5^+/\text{C}_8\text{H}_{11}^+$  events is set to be positive for better contrast in the map. (b) The projection of (a) showing the experimental total ion KER of the  $\text{C}_2\text{H}_3^+/\text{C}_3\text{H}_3^+$  (green) and the  $\text{C}_2\text{H}_5^+/\text{C}_8\text{H}_{11}^+$  (blue) ion pairs. The theoretical ion KER values obtained from PES (Fig. 8) are shown as vertical dashed lines. (c) The projection of (a) showing the experimental Auger electron energy for the  $\text{C}_2\text{H}_3^+/\text{C}_3\text{H}_3^+$  (green) and the  $\text{C}_2\text{H}_5^+/\text{C}_8\text{H}_{11}^+$  (blue) ion pairs.



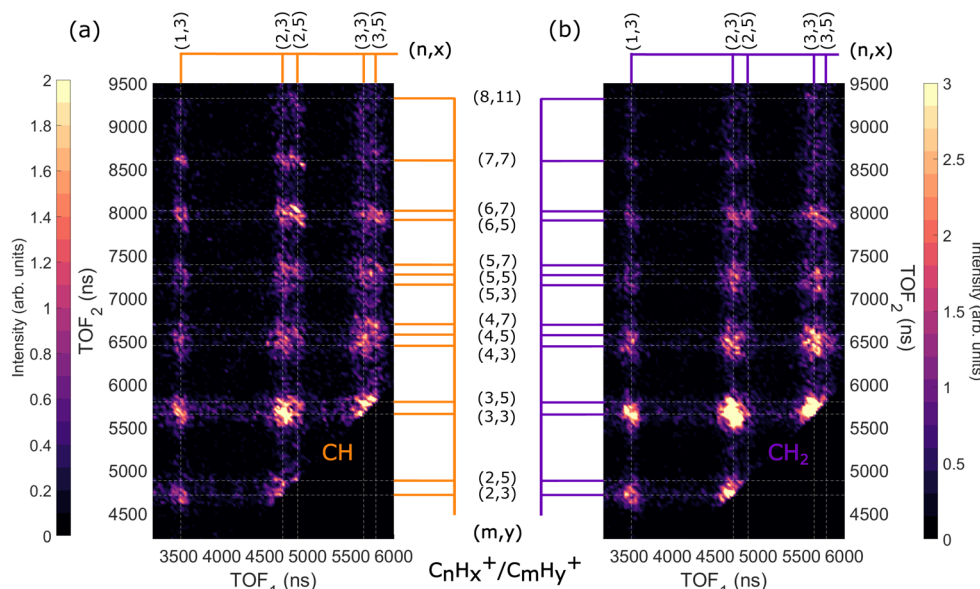


Fig. 10 Zoomed in AEPICO of ion pairs recorded in coincidence with CH AES band (a) and CH<sub>2</sub> AES band (b) following ionization of adamantane at  $h\nu = 350$  eV. The maps have been normalized to maintain the 4:6 (4CH:6CH<sub>2</sub>) ratio that we expect in the adamantane molecule.

From these two examples, we illustrate that the large amount of internal energy deposited by core-ionization compare to valence ionization privileges multi-body breakups and the production of small fragments.

**3.4.3 Site-sensitive dissociation.** Unlike heteronuclear site-specific studies, it is rather difficult to selectively ionize the different carbon sites (CH and CH<sub>2</sub>) in adamantane. Nevertheless, the use of the AEPICO technique enables the observation of such site-sensitivity by filtering the Auger electron kinetic energy.

Fig. 10(a) and (b) shows the filtered AEPICO maps recorded in coincidence with the CH and CH<sub>2</sub> AES bands (Fig. 4), respectively. The intensities of the maps have been normalized to maintain the 4:6 (4 CH:6 CH<sub>2</sub>) ratio that we expect in the adamantane molecule. We observe that the map in Fig. 10(a) has more ion pairs containing C<sub>2</sub>H<sub>5</sub><sup>+</sup> and C<sub>3</sub>H<sub>5</sub><sup>+</sup> than the map in Fig. 10(b). Generally, ions with more hydrogen atoms attached are more likely to be created when we ionize the CH site than the CH<sub>2</sub> site. There is a significant difference in the intensities of the dominant ion pair C<sub>2</sub>H<sub>3</sub><sup>+</sup>/C<sub>3</sub>H<sub>3</sub><sup>+</sup> in Fig. 10(a) and (b). Also, the C<sub>2</sub>H<sub>5</sub><sup>+</sup>/C<sub>6</sub>H<sub>5,7</sub><sup>+</sup> and C<sub>2</sub>H<sub>5</sub><sup>+</sup>/C<sub>7</sub>H<sub>7</sub><sup>+</sup> ion pairs are enhanced for CH site whereas the C<sub>3</sub>H<sub>3</sub><sup>+</sup>/C<sub>4</sub>H<sub>3,5</sub><sup>+</sup> ion pairs are enhanced for CH<sub>2</sub> site.

Therefore, we find that ions with more hydrogen atoms attached are more likely to be created when we ionize the CH site than the CH<sub>2</sub> site. It seems that CH<sub>2</sub> site-ionization creates a final dicationic state that prefers to undergo hydrogen evaporation and produces ions with fewer hydrogen atoms attached.

## 4 Conclusions

In this work, we investigated the dissociation of adamantane dicationics produced after core-ionization at the C-1s edge and

Auger decay using energy-resolved electron-ion multi-coincidence spectroscopy and theoretical calculations. We first determined the electronic structure of the ionized adamantane molecule using high-resolution spectroscopic techniques XPS, NEXAFS, AES and RAES measurements. The XPS hinted to the contributions of the two carbon sites (CH and CH<sub>2</sub>) of the molecule supported by theoretical calculations based on FBA methods. RAES and theoretical simulations showed that two energy bands in the AES can be identified as main contributors from these two carbon sites. We then performed AEPICO spectroscopy to assess the fragmentation dynamics of the dication of adamantane. We succeeded to identify and access the kinematics of the different fragmentation pathways and determined their branching ratios. The energetics of two cases C<sub>2</sub>H<sub>3</sub><sup>+</sup>/C<sub>3</sub>H<sub>3</sub><sup>+</sup> and C<sub>2</sub>H<sub>5</sub><sup>+</sup>/C<sub>8</sub>H<sub>11</sub><sup>+</sup> have been presented to illustrate the differences in the mechanisms of dissociation involved. Higher internal energies are associated with stronger dissociation. Finally, we used the filtering of the PIPICO measurement with both CH and CH<sub>2</sub> Auger electron bands. Our results suggest the presence of site-sensitive dissociation even though the effect can be blurred by the non-single state contributions in the Auger spectrum. Thus, this study, which combines both experimental and theoretical results, pushes the boundaries of understanding site sensitivity in a carbon cage. Further work is needed with a higher degree of theory to reveal the role of electronic excitation, and its interplay with nuclear dynamics, in the dissociation of the adamantane dication.

Our results show that the adamantane molecule fragments into small hydrocarbons after exposure to soft X-rays and could partially play a part in the deficiency of diamondoids in space. In addition, diamondoids could contribute as a source of reactive species for carbonaceous cyclic compounds formation.<sup>60</sup> Therefore, this study could be used to model the X-ray driven photodissociation of adamantane and perhaps



other diamondoids in accompaniment to the X-ray driven photodissociation of PAHs.

## Author contributions

ARM, PR and SM performed the experiments. SG and SM analyzed the data. RFF made the theoretical calculations for the Auger spectra. SDT computed the HF orbitals and PES. PAH and ZA performed theoretical calculations for XPS C-1s spectrum. All authors participated to the scientific discussions concerning the interpretation of the results. SG and SM wrote this manuscript and all authors helped to improve it.

## Conflicts of interest

There are no conflicts to declare.

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