A Raman study of ion irradiated icy mixtures

G. Ferini, G. A. Baratta, and M. E. Palumbo

INAF, Osservatorio Astrofisico di Catania, Via S. Sofia 78, 95123 Catania, Italy

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Abstract. In this paper we present a Raman study of pure CH₄, H₂O:CH₄:N₂ and CH₃OH:N₂ frozen films before and after ion irradiation at 12 K, 100 K and 300 K. By means of Raman spectroscopy, we monitor the structural evolution of each film, whose chemical and physical properties are deeply modified by the interaction with the ion beam. For the two methane containing samples, Raman spectra show that the initial ice is partially converted into a refractory residue, which under further irradiation evolves towards an amorphous carbon (AC) with a band near 1560 cm⁻¹ (G line) and a shoulder at about 1360 cm⁻¹ (D line). No evidence of the AC Raman band is seen in the spectra of the methanol-containing mixture. By means of Lorentzian fits, we have determined the specific parameters of the AC band (G and D line peak positions, widths and relative intensities) in our spectra after ion irradiation and we have compared them with the corresponding parameters of the band as observed in the spectra of 11 IDPs (Interplanetary Dust Particles). Here we present the experimental results and discuss their contribution to our knowledge of the origin and evolution of IDPs.

Key words. astrochemistry – molecular processes – methods: laboratory – techniques: spectroscopic – comets: general – interplanetary medium

1. Introduction

Comets are believed to be among the less altered objects in the Solar System (and hence the best reservoir of presolar material) and one of the main sources of interplanetary dust grains that are collected in our atmosphere and analysed in terrestrial laboratories. It is known that the external layers of comets are exposed to background radiation (galactic cosmic rays) that, according to some models, can drive the formation of an organic crust, which can survive when the comet approaches the Sun (e.g., Strazzulla & Johnson 1991). Raman spectroscopy is one of the laboratory techniques used to get information about the constituent materials of these grains. Furthermore it has been used in the laboratory to study the effects of ion irradiation on the formation and further evolution of refractory carbonaceous materials. By means of in situ Raman spectroscopy, it has been shown that amorphous carbon can be formed at low temperatures after ion irradiation of butane and benzene (Strazzulla & Baratta 1992). On the basis of these results it has been suggested that an organic crust could already be formed during ion irradiation at low temperature, i.e. while comets stay in the Oort cloud (T ∼ 10–20 K), without requiring a first passage by the Sun and its consequent heating. Once a comet is formed, the inner layers are somehow shielded, and are altered only by the most energetic but less abundant ions. The frozen gases in these regions can sublimate when the comet approaches the Sun, passing through the porous structure of the crust. In fact, experiments on targets much thicker than the penetration depth of irradiating ions showed that an organic residue is left over after the sublimation of underlying ices, thus giving support to the hypothesis that an ion-produced cometary organic crust can “survive” ejection from deeper layers (i.e. passages near the Sun) and be as old as the comet itself (Strazzulla et al. 1991). Thus, a dark surface can be formed, which could exhibit strong albedo variations corresponding to patches or fissures in the crust through which volatiles are expelled. Albedo variations have indeed been confirmed by recent observations (Soderblom et al. 2002) of the comet 19P/Borrelly, whose nucleus has an average geometric albedo of only 0.03 ± 0.005, with darker spots ranging down to ∼0.01.

Interplanetary dust particles (IDPs) are believed to derive mainly from comets and asteroids (a fraction of them has an interstellar origin), and they are the smallest and most fine-grained extraterrestrial objects available for laboratory investigation. Recent studies (e.g., Joswiak et al. 2000; Thomas et al. 1993) have underlined that morphological and mineralogical differences exist between asteroidal and cometary IDPs. Morphologically, asteroidal IDPs tend toward smoother, compact forms while cometary IDPs exhibit fluffy, porous, aggre-gate textures. Mineralogically, asteroidal IDPs are often composed of phyllosilicate minerals with only minor amounts of anhydrous phases present, while cometary IDPs are dominated by anhydrous phases and may contain significant carbon and GEMS. The average density obtained for asteroidal

Send offprint requests to: M. E. Palumbo, e-mail: mepalumbo@ct.astro.it
IDPs is 2.5 g cm$^{-3}$ whereas cometary IDP densities average 1.1 g cm$^{-3}$. However, it can be difficult to unambiguously classify an individual IDP as either asteroidal or cometary since characteristics of each group can be found within the other. The presence of carbon in IDPs was first detected by Fraundorf et al. (1982) and confirmed by subsequent Raman studies (McKeegan et al. 1984; Fahey et al. 1985; Macklin et al. 1987; Wopenka 1988; Raynal et al. 2001). The amount of carbon can be greatly variable (Thomas et al. 1995); anyway it has been found that the C-rich material is amorphous or poorly crystalline and could present vesicular textures (Keller et al. 1994; Thomas et al. 1993). The IDPs’ amorphous carbon (AC) could be the result of ion irradiation processes of volatile materials (frozen hydrocarbons in the parent bodies) and thus it is interesting to compare the profile of the AC band in such IDPs with that of ion-irradiated laboratory samples.

In this paper we present Raman spectra of three icy samples (pure CH$_4$ and the mixtures H$_2$O:CH$_4$:N$_2$ and CH$_3$OH:N$_2$) irradiated with He$^+$ (30 keV), Ar$^+$ (30 keV) or Ar$^{++}$ (60 keV) ions at low temperature (12 K) and then gradually warmed up. Previous experiments performed on pure frozen hydrocarbons, such as benzene and butane (Strazzulla & Baratta 1992) have shown that ion irradiation at low doses makes the samples fluorescent and, at higher doses, a broad band appears around 1560 cm$^{-1}$, which is attributed to amorphous carbon. A detailed analysis of this feature makes it possible to infer information on the structural properties of a given sample. The new experiments here discussed show that ion irradiation can drive the formation of amorphous carbon also in a pure methane ice or in hydrocarbon-bearing mixtures (e.g. H$_2$O:CH$_4$:N$_2$). Amorphous carbon samples formed after ion irradiation of icy mixtures will be referred to as IPHAC (Ion Produced Hydrogenated Amorphous Carbon). Moreover, we compare the AC band of laboratory samples to that of 11 IDPs. These IDPs are some out of the 20 that Wopenka (1988) analyzed by means of Raman spectroscopy (in particular we considered the ones in whose spectra the AC Raman feature has a higher signal-to-noise ratio). This analysis showed that these particles can be divided in six different groups depending on the profile of the amorphous carbon feature. Particles which exhibit an amorphous carbon feature that indicates a high degree of order have been assigned to group 1 while those with a decreasing degree of order have been classified in groups 2 to 5. Particles which do not show the amorphous carbon feature have been assigned to group 6. Among laboratory samples we have considered icy mixtures after ion irradiation (IPHAC) here discussed and hydrogenated amorphous carbon grains (HAC) produced from non-ice experiments (e.g., Mennella et al. 2001, 2002). We have found that the characteristics of the AC band observed in the spectra of the three icy mixtures discussed in this paper differ from those exhibited by IDPs, implying that the corresponding structural properties are different while the amorphous carbon band profile of HAC is similar to that of IDPs.

In the following section we will briefly describe the experimental set-up, then we will give some general information on the profile of the amorphous carbon band and on what can be inferred from it. The results of the experiments performed and the Raman spectra of the films are presented in Sect. 4. The astrophysical implications are discussed in Sect. 5.

2. Experimental set-up

The in situ analyses have been performed in a stainless steel vacuum chamber facing a Raman spectrometer. Figure 1 schematically shows the arrangement used for in situ Raman spectroscopy. A continuous Ar-ion laser beam ($\lambda = 514$ nm) enters a confocal illuminator perpendicular to its optical axis, into which it is deflected by a microprism. The confocal optical system is arranged in such a way that any parallel beam incident along the optical axis in the opposite direction of the laser is focused onto the entrance slit of the spectrometer. By means of two flat mirrors, the laser beam is reflected towards the vacuum chamber and focused by an objective into a 40 $\mu$m spot on the sample. The back-scattered Raman light is collimated by the same objective into a parallel beam that, according to the setup geometry, goes back along the same path of the laser beam and reaches the confocal illuminator where it is focused onto the entrance slit of the spectrometer.

Inside the vacuum chamber, in which the pressure is kept below $10^{-7}$ mbar, a substrate (crystalline silicon) is placed in thermal contact with a cold finger whose temperature can be varied between 10 K and 300 K. The vacuum chamber is interfaced with an ion implanter (30 kV; Danfysik) from which ions with energy up to 30 keV (60 keV for double ionizations) can be obtained. The ion beam produces a $2 \times 2$ cm$^2$ spot on the target and currents in the range of 100 nA cm$^{-2}$ to a few $\mu$A cm$^{-2}$.

A needle valve is used to admit pre-prepared gases (or mixtures) into the chamber, where they freeze on the substrate. A He-Ne laser can be used to monitor the thickness of the ice film during accretion; this is achieved by looking at the interference pattern (intensity versus time) given by the laser beam reflected at an angle of 45$^\circ$ both by the vacuum-film and film-substrate interfaces (see Baratta & Palumbo 1998 for further details on the technique used to measure the thickness).

The substrate holder is mounted at an angle of 45$^\circ$ both with the ion beam and the laser beam. This configuration offers the advantage that spectra can be easily taken in situ, even during irradiation, without tilting the sample.

The amount of energy released into the sample (dose) is expressed in units of eV per small molecule (16 amu) and is calculated from the knowledge of the ion fluence $F$ (ions cm$^{-2}$), the stopping power $S$ (eV cm$^{-2}$ molecule$^{-1}$) for the chosen projectile or its penetration depth (range) $r$ into the target (molecules cm$^{-2}$). The first is given by a current integrator on the path of the ion beam, which measures the charge that reaches the sample during irradiation; the other two parameters are easily calculated by using available routines (e.g. TRIM, etc.). Thus, if the penetration depth is greater than the sample thickness, the dose is:

$$d = S \times F.$$  

If the range is smaller than the sample thickness, the ions are stopped in the target and only the molecules within a certain depth in the ice, roughly equal to the range, interact with the
ion beam. In this case the average energy absorbed by each molecule is:

\[ d = \frac{1}{r} \times F \times E, \]

where \( E \) is the projectile energy given in eV.

The icy films we report on in the following are all several \( \mu \text{m} \) thick, while the penetration depth of the ions used as projectiles ranges between 0.05 \( \mu \text{m} \) and 0.5 \( \mu \text{m} \). This implies that the incoming ions stop in the target and only the outer layers of the ice are processed.

### 3. Raman spectroscopy of carbonaceous materials

Raman spectroscopy gives valuable information on the vibrational transitions of molecules in the solid, liquid, and gas phase. In particular it is a powerful tool to investigate the structural properties of a given sample and so it has often been used to study the effects of ion-induced lattice damage in carbonaceous solids and organic compounds (Elman et al. 1981; Strazzulla & Baratta 1992; Baratta et al. 1996; Musumeci et al. 2000; Strazzulla et al. 2001 and references therein). Raman spectroscopy makes it possible to perform mineralogical studies (see for example Wang et al. 2000), to identify substances and chemical compounds, and to deduce structural and symmetry properties of molecules. For these abilities Raman analysis may have important applications in planetology, especially in the “in situ” and/or laboratory studies concerned with the evolution of surface rocks and also in the determination of past environments, as recorded by meteorites. Interplanetary dust grains can also be analysed by Raman spectroscopy, inferring information on the composition and processing suffered by their parent bodies.

In this paper we will concentrate on what can be deduced from the study of the 1560 cm\(^{-1}\) AC band. The profile of this band differs from one carbonaceous material to another, reflecting the great variation in chemical composition and structure (“degree of order”), and this makes Raman spectroscopy a powerful technique of classification for samples which exhibit this characteristic feature.

Generally speaking, Raman spectroscopists associate the crystallite size to the crystal order or “perfection”. In HOPG (Highly Ordered Pyrolitic Graphite) or natural graphite with large (>1 \( \mu \text{m} \)) microcrystals a band is present at about 1582 cm\(^{-1}\) (Robertson 1986), which is called the G (or graphitic) line. Because of the momentum conservation selection rule, this is the only active first-order strong Raman band and it is due to the \( E_{2g} \) mode (Tuinstra & Koenig 1970). The spectra of microcrystalline (\( \leq 1 \mu \text{m} \)) graphite show an additional line peaked at 1350 cm\(^{-1}\), called the D (or disorder) line.

It has been found (Tuinstra & Koenig 1970) that a correlation exists between the intensity ratio \( I_D/I_G \) and the coherence length \( L_0 \) of the crystal, which can be taken as an indicator of the degree of order: \( I_D/I_G \propto 1/L_0 \) if \( 2.5 \text{ nm} < L_0 < 1 \mu \text{m} \). In more disordered carbons (\( L_0 < 2.5 \text{ nm} \)) this correlation is no longer valid; for example, in amorphous carbons with a low degree of order an opposite trend is observed. In this case, the \( I_D/I_G \) ratio increases with the cluster size. An indication of the degree of order can be also provided by the G and D bands width: the broader the bands are, the more
disordered the amorphous carbon is. In fact, since the FWHM of both G and D lines depends on the bond angle disorder, and on the relative amounts of crystallites versus the amorphous matrix, a broadening of the bands indicates an increase in the bond angle disorder and a decrease in the mean size and/or number of crystallites. Below a certain size of the basic structural units of the carbonaceous material, the bands become so wide that they can no longer be recognized as individual peaks (Wopenka 1988).

Carbonaceous materials with different degree of order differ in number and/or size of the sp2 clusters. When the sample is ion-irradiated, these parameters change. Several laboratory experiments performed on carbonaceous materials have shown that ion irradiation can increase or decrease the sp2 cluster size depending on the starting material. During ion irradiation two competitive processes are present: graphitization and amorphization. The first one is related to the energy release in the form of heat (“thermal spikes”) inside the collision cascade (sp3 to sp2 conversion with or without hydrogen loss), while the second is caused by displacement collisions and depends on the size of crystallites. Since the relaxation time of the excitation generated by an ion is of the order of one picosecond or less, equilibrium processes (like annealing) are not possible. The saturation effect is due to the competition between these two processes (Compagnini et al. 1992), in which saturation has been reached when they become equal.

The effects of releasing energy to a poorly ordered amorphous carbon by thermal annealing are different. Experiments have been performed on AC samples with a low degree of order (Dillon et al. 1984; Vidano & Fishbach 1981; Vidano 1980), showing that the I_D/I_G ratio increases up to a maximum value at T = 800–900°C and then it goes to zero at higher annealing temperature. In summary, energetic processing greatly affects G and D lines peak positions, widths and relative intensities, because these parameters are directly linked to the lattice structure, which changes after ion irradiation and/or annealing. In order to qualitatively analyse these variations and particularly the relative contribution of the D and G lines, we have done a Lorentzian line shape fitting analysis for the amorphous carbon band in the Raman spectra of the laboratory samples. A similar analysis was performed for the band observed in the spectra of some IDPs.

4. Results

We present the results obtained for three different icy mixtures containing hydrocarbons: pure CH4, H2O:CH4:N2 (1:6:3) and CH3OH:N2 (1:2). Each mixture was first deposited at 12 K, then irradiated (with non-reactive ions such as He+ or Ar++) and finally warmed gradually to room temperature. Raman spectra of the samples were taken for different doses. In Figs. 2, 3 and 7 the corresponding infrared spectra of the mixtures as deposited (top curves) are also shown for comparison. Since a molecule can have infrared active transitions which are not Raman-active (or extremely weak) and vice versa (because of the different selection rules), some features present in the IR spectrum are absent from the Raman spectrum. For example, water bands, very intense in the IR spectrum of thin icy films, cannot be easily observed in the Raman spectrum (because of the small cross-section), while the opposite is the case for the symmetric modes of mono- and poly-nuclear molecules such as N2 and CH4 (which are IR-inactive). These considerations are summarized in Table 1 (where we also provide the observed peak positions of the Raman features in our spectra) and indicate that the two spectroscopic techniques can be used as complementary tools in identifying the solid-phase features of various molecules.
Table 1. IR and Raman features of the molecules studied here.

<table>
<thead>
<tr>
<th>Molecule</th>
<th>Vibrational mode</th>
<th>Position (cm(^{-1}))</th>
<th>IR</th>
<th>Raman</th>
</tr>
</thead>
<tbody>
<tr>
<td>H(_2)O</td>
<td>OH sym &amp; asym stretch</td>
<td>3300</td>
<td>3400 (active)</td>
<td>active</td>
</tr>
<tr>
<td>H(_2)O</td>
<td>OH bend</td>
<td>1660</td>
<td>1640 (active)</td>
<td>active</td>
</tr>
<tr>
<td>H(_2)O</td>
<td>libration</td>
<td>675</td>
<td>800 (active)</td>
<td>active</td>
</tr>
<tr>
<td>CH(_4)</td>
<td>C-H stretch (asym)</td>
<td>3010</td>
<td>3011 (active)</td>
<td>active</td>
</tr>
<tr>
<td>CH(_4)</td>
<td>C-H stretch (sym)</td>
<td>2900</td>
<td>inactive</td>
<td>active</td>
</tr>
<tr>
<td>CH(_4)</td>
<td>C-H deform (sym)</td>
<td>1534</td>
<td>1547 (inactive)</td>
<td>active</td>
</tr>
<tr>
<td>CH(_4)</td>
<td>C-H deformation</td>
<td>1300</td>
<td>active</td>
<td>inactive</td>
</tr>
<tr>
<td>CH(_3)OH</td>
<td>O-H stretch</td>
<td>3260</td>
<td>3360 (active)</td>
<td>active</td>
</tr>
<tr>
<td>CH(_3)OH</td>
<td>C-H stretch (asym)</td>
<td>2961</td>
<td>2950 (active)</td>
<td>active</td>
</tr>
<tr>
<td>CH(_3)OH</td>
<td>C-H stretch (sym)</td>
<td>2830</td>
<td>2829 (active)</td>
<td>active</td>
</tr>
<tr>
<td>CH(_3)OH</td>
<td>bend &amp; def</td>
<td>1460</td>
<td>1470 (active)</td>
<td>active</td>
</tr>
<tr>
<td>CH(_3)OH</td>
<td>bend &amp; def</td>
<td>1126</td>
<td>1107 (active)</td>
<td>active</td>
</tr>
<tr>
<td>CH(_3)OH</td>
<td>C-O stretch</td>
<td>1034</td>
<td>1021 (active)</td>
<td>active</td>
</tr>
<tr>
<td>CH(_3)OH</td>
<td>torsion</td>
<td>694</td>
<td>inactive</td>
<td>active</td>
</tr>
<tr>
<td>N(_2)</td>
<td>N≡N strecht</td>
<td>2328</td>
<td>2322 (inactive)</td>
<td>active</td>
</tr>
</tbody>
</table>

* Liquid phase position.

Table 2. List of studied icy mixtures.

<table>
<thead>
<tr>
<th>Number</th>
<th>Sample</th>
<th>Ratio</th>
<th>Thickness ((\mu)m)</th>
<th>Ion &amp; energy</th>
<th>Fluence (ions cm(^{-2}))</th>
<th>Total dose (eV/16amu)</th>
<th>Temperature (K)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>CH(_4)</td>
<td>3.7</td>
<td>3.7</td>
<td>He(^+), 30 keV</td>
<td>1.8 (\times) 10(^{15})</td>
<td>64</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>He(^+), 30 keV</td>
<td>1.8 (\times) 10(^{15})</td>
<td>130</td>
<td>100</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Ar(^{++}), 60 keV</td>
<td>1.1 (\times) 10(^{15})</td>
<td>1000</td>
<td>100</td>
</tr>
<tr>
<td>2</td>
<td>H(_2)O:CH(_4):N(_2)</td>
<td>1:6:3</td>
<td>5</td>
<td>He(^+), 30 keV</td>
<td>3.2 (\times) 10(^{16})</td>
<td>825</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>He(^+), 30 keV</td>
<td>2.0 (\times) 10(^{15})</td>
<td>300</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>CH(_3)OH:N(_2)</td>
<td>1:2</td>
<td>10</td>
<td>He(^+), 30 keV</td>
<td>1.2 (\times) 10(^{16})</td>
<td>164</td>
<td>12</td>
</tr>
<tr>
<td>4</td>
<td>CH(_3)OH:N(_2)</td>
<td>1:2</td>
<td>10</td>
<td>Ar(^{++}), 60 keV</td>
<td>1.8 (\times) 10(^{15})</td>
<td>654</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Ar(^+), 30 keV</td>
<td>7.1 (\times) 10(^{16})</td>
<td>1900</td>
<td>12</td>
</tr>
</tbody>
</table>

4.1. Irradiation effects

It is well known that when fast ions impinge on an icy sample, many molecular bonds are broken, and fragments are formed along the ion track which recombine giving rise to molecular species not present in the original sample. Beyond the chemical modifications, ion irradiation drives a gradual alteration of the sample structure, causing a rearrangement of the atoms in their lattice sites and a thorough modification of the optical properties of the ice. As we will show, Raman spectroscopy is a powerful tool for monitoring structural changes. In the next sections we will illustrate the effects of ion irradiation on the Raman spectra of the mixtures reported in Table 2. When different ions have been used to irradiate the same sample, we report in the table the maximum fluence of each ion and the total dose suffered after each irradiation step.

4.2. Pure methane ice

Figure 2 shows the evolution of the Raman spectrum of a pure methane frozen film after irradiation with 30 keV He\(^+\) and 60 keV Ar\(^{++}\) ions. The curve labelled with the letter a represents the spectrum of the sample as deposited at 12 K, in which the features due to silicon (substrate) phonon vibrations (∼525 cm\(^{-1}\) and ∼990 cm\(^{-1}\) for the first and the second order, respectively) are clearly distinguishable. In addition, we see the C-H asymmetric and symmetric stretching modes and the C-H symmetric deformation mode of CH\(_4\) molecules, that peak around 3010, 2900 cm\(^{-1}\) and 1550 cm\(^{-1}\) respectively. After irradiation at 12 K with 30 keV He\(^+\) ions (up to a dose of 64 eV/16 amu), the sample was warmed up to 100 K (spectrum not shown), a temperature at which underlying unirradiated CH\(_4\) had already sublimated, and then further
irradiated with different ions (30 keV He⁺, 60 keV Ar⁺) up to 1000 eV/16 amu. From infrared spectroscopy (e.g., Kaiser & Roessler 1998; Baratta et al. 2002; Moore & Hudson 2003) we know that ion irradiation of pure methane produces other hydrocarbons (such as ethane, C₂H₆, and propane, C₃H₈), which sublime during warm up at about 60–80 K, and a refractory residue stable at room temperature (e.g., Strazzulla et al. 1983; Strazzulla & Johnson 1991). The formation of an organic residue is a general ion irradiation effect and occurs for different icy mixtures, as has been discussed elsewhere (Moore et al. 1983; Foti et al. 1984; Strazzulla & Johnson 1991). The nature of the residue formed and its thermal stability depend on several parameters such as the initial composition of the ice and the irradiation dose. The “organic residue” obtained by ion irradiation of frozen icy mixtures containing C, H, N, and O bearing species is mainly constituted by C=C, C=O, C≡N, C-H and O-H bonds, as shown by infrared spectroscopy analysis (Strazzulla et al. 2001).

In Fig. 2 the curves labelled with b, c and d show the spectra of the residue for increasing irradiation doses at 100 K: 130, 290 and 1000 eV/16 amu. In these spectra the silicon features are no longer detectable, implying that the icy film optical properties have been modified (darkening) so that the laser beam is absorbed in the sample and can no longer reach the substrate. Furthermore, after irradiation, an intense fluorescence continuum appears, typical of disordered polymer-like carbons containing a relatively large quantity of hydrogen, and due to radiative recombination of electron-hole pairs produced by laser irradiation. With increasing irradiation fluence, the fluorescence signal becomes weaker and weaker, due to the gradual loss of hydrogen. When higher doses are reached, a broad band appears near 1500 cm⁻¹, which is attributed to C=C stretching vibration of amorphous carbon. These effects were already known for other simple frozen hydrocarbons, both aromatic such as benzene (C₆H₆) and aliphatic such as butane (C₄H₁₀; Strazzulla & Baratta 1992). In summary ion irradiation of frozen methane induces the following effects:

- at low fluences, the sample shows an intense fluorescence signal due to the formation of a damaged molecular solid;
- as irradiation goes on, hydrogen is lost and the fluorescence signal intensity decreases; the Raman feature of amorphous carbon appears and gets more and more intense.

All these effects occur at low temperature and cannot be investigated using infrared spectroscopy, which testifies to the chemical evolution of the sample, but gives little information on its structure.

A fit of the profile of the amorphous carbon feature after the highest dose of irradiation with two Lorentzian components gives νg = 1517 cm⁻¹, FWHMg = 146 cm⁻¹, νD = 1265 cm⁻¹, FWHM_D = 314 cm⁻¹, and I_D/I_G = 0.232.

4.3. The mixture H₂O:CH₄:N₂

Figure 3 shows the spectra of an icy mixture (H₂O:CH₄:N₂ (1:6:3)) before and after irradiation with 30 keV He⁺ ions at 12 K. Nitrogen (~2350 cm⁻¹, for the symmetric stretching mode) and methane features are present, while the Raman features of water ice are undetectable.

The amorphous carbon band is distinguishable at doses of 378 eV/16 amu (the corresponding spectrum is labelled with letter “b” in the figure) and its profile and intensity vary as irradiation goes on (the spectrum labelled with the letter “c” refers to the sample after a dose of 825 eV/16 amu). Figure 4 illustrates the result of the fitting procedure with Lorentzian components; the corresponding fitting parameters are shown on the right hand side of the figure.

We can see, in particular, that the G-line peak position shifts to higher frequencies with increasing dose. These results are consistent with an increasing size or number of the largest sp² clusters (with respect to the sp³ clusters) induced by ion irradiation (Baratta et al. 1996).

After irradiation at low temperature of the mixture H₂O:CH₄:N₂, the icy sample was warmed up to room temperature and left under vacuum overnight and then the spectrum was taken (Fig. 5, top curve). Since the penetration depth of the impinging ions was smaller than the sample thickness, just a thin layer of the target was altered by irradiation and so converted into a refractory material. No features due to the ice constituents are obviously present in the spectrum at room temperature, because the underlying ice has sublimated passing through the porous structure of the residue. Only the C=C 1560 cm⁻¹ band is observed superimposed on the fluorescence continuum background.

The organic residue has been further irradiated at room temperature with He⁺ (30 keV) ions. As we can see from Fig. 5, during irradiation the fluorescence background intensity decreases, as occurs at low temperature.

If we look at the fitting parameters reported in Fig. 6, we note that the I_D/I_G ratio decreases (from 0.74 to 0.57) during irradiation. Moreover, the FWHM of both G and D lines
It is evident that the fluorescence background greatly decreases after irradiation. The characteristic feature of such a disordered structure is the progressive evolution of the sample towards an amorphous carbon. The apparent jumps from a spectral region to another indicate laser induced alteration of the sample.

4.4. The mixture $\text{CH}_3\text{OH}:\text{N}_2$

Finally we describe the spectral modifications induced by ion irradiation on a frozen film made up of methanol and nitrogen. The corresponding Raman spectra at 12 K, both before (curve a) and after (curve b) ion irradiation, are presented in Fig. 7. In this case we can see all the original features even after irradiation with a total fluence of $10^{16}$ He$^+$ ions/cm$^2$. The corresponding Raman spectra at 12 K, both before (a) and after (b) ion irradiation with He$^+$ (30 keV) ions are shown. The characteristic feature of such a disordered structure is the 1560 cm$^{-1}$ band, which forms at low temperature. It was already known that amorphous carbon (IPHAC) can be formed after ion irradiation of icy samples of pure hydrocarbons such as $\text{C}_2\text{H}_6$ and $\text{C}_6\text{H}_6$ (both aliphatic and aromatic). The new experiments here discussed show that the formation of IPHAC occurs not only for pure hydrocarbons ($\text{CH}_4$), but also in the case of a mixture containing methanol as carbon bearing species. In the case of a hydrocarbon, such as methane, benzene, etc., doses of some 10–10$^2$ eV/mol are sufficient to make the 1560 cm$^{-1}$ band observable.

5. Discussion

In this paper we have discussed Raman spectra of three ion-irradiated icy mixtures at low (12 K, 100 K) and at room temperature. Raman spectroscopy is useful to understand the progressive evolution of the sample towards an amorphous carbon. The characteristic feature of such a disordered structure is the 1560 cm$^{-1}$ band, which forms at low temperature. It was already known that amorphous carbon (IPHAC) can be formed after ion irradiation of icy samples of pure hydrocarbons such as $\text{C}_2\text{H}_6$ and $\text{C}_6\text{H}_6$ (both aliphatic and aromatic). The new experiments here discussed show that the formation of IPHAC occurs not only for pure hydrocarbons ($\text{CH}_4$), but also in the case of a mixture containing methanol. This supports the idea that the process of IPHAC formation is general. Anyway, there seem to be great differences in the efficiency of the process itself (Sect. 4.4).

The typical parameters (peak position, width and intensity of both G and D lines) of the AC band depend on the carbonaceous material considered. In Figs. 8 and 9 we compare the FWHM of G and D lines, versus the corresponding peak position, for different carbonaceous material, which are: a) IPHAC (samples 1 and 2 of Table 2) both at low and room temperature, b) different kinds of hydrogenated carbon.
Table 3. Parameters of Lorentzian fit to the Raman spectra of 11 IDPs. Particle names have been assigned by Wopenka (1988).

<table>
<thead>
<tr>
<th>Particle</th>
<th>G line position (cm⁻¹)</th>
<th>G line FWHM (cm⁻¹)</th>
<th>D line position (cm⁻¹)</th>
<th>D line FWHM (cm⁻¹)</th>
<th>I₀/D₀</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tyson</td>
<td>1594</td>
<td>73</td>
<td>1350</td>
<td>141</td>
<td>1.068</td>
</tr>
<tr>
<td>Attila</td>
<td>1594</td>
<td>72</td>
<td>1345</td>
<td>202</td>
<td>0.936</td>
</tr>
<tr>
<td>Essex</td>
<td>1591</td>
<td>82</td>
<td>1361</td>
<td>267</td>
<td>0.896</td>
</tr>
<tr>
<td>Summersville</td>
<td>1589</td>
<td>100</td>
<td>1393</td>
<td>469</td>
<td>1.392</td>
</tr>
<tr>
<td>Elias</td>
<td>1579</td>
<td>101</td>
<td>1347</td>
<td>288</td>
<td>1.326</td>
</tr>
<tr>
<td>Lea</td>
<td>1584</td>
<td>159</td>
<td>1379</td>
<td>269</td>
<td>0.978</td>
</tr>
<tr>
<td>Mosquito</td>
<td>1562</td>
<td>114.3</td>
<td>1349</td>
<td>476</td>
<td>1.044</td>
</tr>
<tr>
<td>Pattonsburg</td>
<td>1577</td>
<td>118</td>
<td>1373</td>
<td>373</td>
<td>1.129</td>
</tr>
<tr>
<td>SkyFB</td>
<td>1591</td>
<td>90</td>
<td>1371</td>
<td>159</td>
<td>0.663</td>
</tr>
<tr>
<td>Spray</td>
<td>1581</td>
<td>126</td>
<td>1364</td>
<td>338</td>
<td>0.872</td>
</tr>
<tr>
<td>Verona</td>
<td>1585</td>
<td>104</td>
<td>1358</td>
<td>284</td>
<td>0.966</td>
</tr>
</tbody>
</table>

Table 4. Parameters of Lorentzian fit to the Raman spectra of amorphous carbon samples from non-ice experiments (Baratta et al. 1996; Mennella et al. 2003).

<table>
<thead>
<tr>
<th>Sample</th>
<th>Fluence (30 keV He⁺ cm⁻²)</th>
<th>G line position (cm⁻¹)</th>
<th>G line FWHM (cm⁻¹)</th>
<th>D line position (cm⁻¹)</th>
<th>D line FWHM (cm⁻¹)</th>
<th>I₀/D₀</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACAR</td>
<td>1589</td>
<td>86</td>
<td>1372</td>
<td>355</td>
<td>1.037</td>
<td></td>
</tr>
<tr>
<td>ACAR_H</td>
<td>1585</td>
<td>102</td>
<td>1360</td>
<td>284</td>
<td>0.953</td>
<td></td>
</tr>
<tr>
<td>ACAR_H</td>
<td>1585</td>
<td>114</td>
<td>1363</td>
<td>307</td>
<td>1.032</td>
<td></td>
</tr>
<tr>
<td>ACAR_H</td>
<td>9.5 × 10¹⁵</td>
<td>1553</td>
<td>159</td>
<td>1356</td>
<td>463</td>
<td>0.679</td>
</tr>
<tr>
<td>ACH2</td>
<td>6.0 × 10¹⁵</td>
<td>1551</td>
<td>147</td>
<td>1356</td>
<td>388</td>
<td>0.667</td>
</tr>
<tr>
<td>ACH2</td>
<td>1.1 × 10¹⁶</td>
<td>1557</td>
<td>162</td>
<td>1363</td>
<td>408</td>
<td>0.716</td>
</tr>
</tbody>
</table>

grains (labelled ACH2, ACAR, and ACAR_H¹) produced in the laboratory from non-ice experiments and c) some IDPs. The mentioned laboratory materials (organic residues obtained after ion irradiation of frozen gases and mixtures, hydrogenated carbon grains) have been considered to test if different kinds of AC samples could be similar, at least from a structural point of view, to the carbon phase of interplanetary grains. Here we have selected 11 different IDPs among the 20 already studied by Wopenka (1988), choosing the ones whose Raman spectra have a higher signal-to-noise ratio. These particles, whose dimensions are between 1 and 50 µm, have been collected by NASA spacecrafts at altitudes between 8 and 20 km and have been studied by means of micro-Raman spectroscopy, infrared transmission spectroscopy, scanning electron microscopy (SEM) and energy-dispersive X-ray (EDX) analyses (e.g., Sandford & Walker 1985). It has been found that the particles are heterogeneous, with different mineralogical and structural properties. They contain silicates, in variable amounts, (whose Raman features have not been observed probably due to the presence of carbonaceous material that masks the signal of silicates), and amorphous carbon with different degree of order, as can be seen from Table 3, in which we report the fitting parameters derived by the Lorentzian fits to the IDPs’ AC band. The fitting parameters derived by the Lorentzian fits of the AC band for the samples obtained from non-ice experiments are reported in Table 4. In the case of ACAR_H samples, two different specimens were analysed and, as expected, the fit parameters are very similar.

Here, we compare the “laboratory” and the “interplanetary” values of these parameters and try to test the hypothesis that the amorphous carbon phase of IDPs could be similar to ion irradiated ices. In fact, according to some models (Strazzulla & Johnson 1991), the surface layers of comets (0.1 – 0.5 m) are exposed to an irradiation dose of up to 600 eV/molecule. This energetic processing drives the formation of a substantial “crust” of non-volatile material, left over when volatile material from the deeper layers sublimes in the passages near the Sun. Part of this crust, in the form of IDPs, could enter the Earth atmosphere and then be analysed by means of laboratory techniques. Looking at Fig. 8 and particularly at Fig. 9, that illustrate D

¹ ACH2 was prepared by condensation of carbon vapor produced by an arc discharge between two carbon rods in a 10 mbar hydrogen atmosphere (Mennella et al. 2001). The label ACAR refers to hydrogen-free carbon grains produced by laser ablation of carbon electrodes in an argon atmosphere. In some cases, these grains were hydrogenated (ACAR_H) by exposure to hydrogen atoms produced by microwave-excited dissociation of molecular hydrogen (further details on the preparation of these grains can be found in Mennella et al. 2002). Some of these samples have been irradiated with 30 keV He⁺ ions (Baratta et al. 1996; Mennella et al. 2003).
...H grains, respectively,

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1250 1300 1350 ... (T ∼400 −1200 ◦C) for a few seconds. This heat pulse
will alter abundant organic matter in IDPs and, for example, it

H are similar to those of IDPs. The latter have a

close to some IDPs, namely Pattonsburg, Mosquito and Verona.

a few data points, relative to unirradiated ACAR samples, are

G line width is reported. From the figure it is evident that only

samples is apparent in Fig. 10, where the

I

position. All the symbols in the graph have the same meaning as in the

previous figure.

Fig. 8. D line full width at half maximum (FWHM) versus peak po-

sition. The stars, all in the middle-bottom part of the figure, refer
to IDPs. The label 1 on the squares indicates the mixture H2 O:CH 4 :N 2
at 12 K at the first irradiation dose, while 2 and 3 refer to the same sam-
ple at increasing irradiation dose. The same criterion is adopted for the

Corresponding residue, which is represented by the up triangles and is

labelled with 4 and 5 respectively before and after irradiation. Labels
ach2 irr and acarh irr refer to ACH2 and ACAR-H grains, respectively,
after ion irradiation.

Fig. 9. G line full width at half maximum (FWHM) versus peak po-

sition. All the symbols in the graph have the same meaning as in the

previous figure.

and G line widths versus the corresponding peak position, we can
say that most of amorphous carbon samples obtained after

ion irradiation of icy mixtures greatly differ from IDPs. On the

other hand the profiles of the amorphous carbon band in ACAR
and ACAR-H are similar to those of IDPs. The latter have a

narrower G line, which peaks at higher wavenumbers than the

AC produced after ion irradiation of icy mixtures, implying that

the structure of IDPs is more ordered than that of the IPHACs

here considered. The most ordered IDPs according to the classification made
by Wopenka, i.e. those with the narrowest G and D lines (viz.

Essex, Attila and Tyson), are far from our IPHACs. The IDP

which is closer to our irradiated ices is Spray, which has a
rather low degree of order. We can conclude that the IDPs here

considered have spectral features which are not reproduced by

IPHAC samples.

The observed results indicate that the AC present in the

studied IDPs differs from that formed by ion irradiation of

frozen gases in the cometary crust. Thus, if the cometary crust
is formed after ion irradiation (Strazzulla & Johnson 1991),
these results question the hypothesis that some IDPs could be

fragments of the cometary crust. Alternatively, we could imagine
that the crust has a different origin or that there is a different
mechanism which leads to the amorphous carbon detected in
interplanetary grains. These may be, for example, “older” than the
crust, in the sense that they could be the sites for the sur-
facial ice to condense when the comet forms. In this case, the
crust should have a different history and composition from the
AC that is embedded in IDPs, suggesting that the crust is very
stable.

Some of the considered IDPs might not have a cometary
origin. Some of them (namely Essex, Attila, Tyson) probably
do have an asteroidal origin, so they could have been processed
at high temperature in the inner Solar System. This hypothe-
sis is confirmed by a terrestrial-like deuterium abundance and
the presence of crystalline olivine. Depending on the IDPs ori-
gin (whether asteroidal or cometary), the interaction with the
atmosphere is different. Dust particles from comets generally
enter the Earth’s atmosphere with more initial energy than that
possessed by asteroidal dust particles (see for example Jackson
& Zook 1992) and hence cometary particles of the same size
and density, on the average, should be heated more in their de-
celeration.

This means that IDPs can be differently annealed by the
terrestrial atmosphere, which heats them by a considerable
amount (T ~ 400–1200 ◦C) for a few seconds. This heat pulse
will alter abundant organic matter in IDPs and, for example, it
may lead in some cases to vesicle formation, and in very extreme cases to a porous structure, made up of interconnected bubbles (Brownlee et al. 1997, 2002). Anyway, though it is known that thermal annealing increases the degree of order in amorphous carbon (since it tends to remove heteroatoms and structural defects in the amorphous carbon, so that the basic structural carbonaceous units re-arrange, raising the “degree of order”) the atmospheric heating alone cannot explain the different spectral characteristic of AC in IDPs. In fact, as Wopenka (1988) points out, Attila and Skywalker, which show nuclear tracks that indicate the same atmospheric heating (below 600 °C), exhibit a different AC Raman feature. This could imply differences in chemical composition and/or structure of the precursor material.

All these considerations indicate that the understanding of the evolution of IDPs needs further experimental work and different kinds of samples are to be considered. So other experiments, such as those here presented, which provide an example of comparison between the AC Raman feature of IDPs and of laboratory samples, are needed for further information.

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