The composition of the protosolar disk and the formation conditions for comets

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Abstract Conditions in the protosolar nebula have left their mark in the composition of cometary volatiles, thought to be some of the most pristine material in the solar system. Cometary compositions represent the end point of processing that began in the parent molecular cloud core and continued through the collapse of that core to form the protosun and the solar nebula, and finally during the evolution of the solar nebula itself as the cometary bodies were accreting. Disentangling the effects of the various epochs on the final composition of a comet is complicated. But comets are not the only source of information about the solar nebula. Protostellar disks around young stars similar to the protosun provide a way of investigating the evolution of disks similar to the solar nebula while they are in the process of evolving to form their own solar systems. In this way we can learn about the physical and chemical conditions under which comets formed, and about the types of dynamical processing that shaped the solar system we see today.

This paper summarizes some recent contributions to our understanding of both cometary volatiles and the composition, structure and evolution of protostellar disks.

Keywords protostellar disks · solar nebula · comets · chemistry

Abbreviations

AGB asymptotic giant branch
ALMA Atacama Large Millimeter Array
CAI calcium-rich aluminium-rich inclusion
FUN Fractionated and unknown nuclear isotopic effects
ISM Interstellar medium
ISRF interstellar radiation field
IRAM Institut de Radioastronomie Millimétrique telescope
MRI magneto-rotational instability
MHD magnetohydrodynamical
OPR ortho-to-para ratio
PSN protosolar nebula
SLR short-lived radionuclide
Gasous inner disk Inside the iceline
Inner disk Inside 35 AU
Outer disk Outside 35 AU
1 Introduction

Cometary volatiles are some of the most pristine of solar system materials, having remained relatively unprocessed since the comets formed in the inner solar system some 4.2 billion years ago. Consequently they retain in their composition signatures of the chemical and physical conditions under which they formed. Observations of cometary comae provide the mixing ratios or relative abundances of various species and have demonstrated that these can vary greatly between comets, e.g., A'Hearn et al. (2012) and Mumma and Charnley (2011) and references therein, with little correlation with dynamical family. The origin of these compositional variations is not well understood but must lie in the physical and chemical processes that were active at the time and location at which the cometary volatiles were formed. To understand these processes requires not only a study of the comets themselves but also of protostellar disks similar to the one from which our solar system formed. Observations of protostellar disks allow for the important disk processes to be investigated while they are occurring and can test models of the evolution of the early solar system.

Some key questions that can potentially be answered by interdisciplinary investigations involving both protostellar disk modeling and observations, and the observations of comets are:

1. Can synergy between protoplanetary disk modeling and compositional studies of comets inform us about the chemical history and processes in the early solar disk?
2. What is the origin of the diversity of observed cometary compositions?
3. Where did comets form in the solar nebula?
4. Are D/H ratios in volatile molecules and spin temperatures cosmogonic indicators for comets? If so, what can they tell us about the formation conditions in the early solar system?
5. What is the range of D/H ratios in cometary water and can we infer the degree to which comets contributed volatiles to the early terrestrial planets?
6. Can cometary volatile abundances be used to constrain the extent of radial and vertical mixing in the early protoplanetary disk?

This paper summarizes some recent developments in our understanding of cometary compositions and their relationship to the conditions under which the comets formed. The paper is arranged as follows. Section 2 covers some observational and modeling studies of protostellar disks. Section 3 discusses fractionation processes in protostellar disks including using deuterium fractionation as a tracer of the links between comets, interstellar chemistry and nebular chemistry. Section 4 considers the origin of the nitrogen deficiency in comets and Section 5 discusses spin temperature as a cosmogenic indicator in comets. Section 6 presents a possible explanation for the observations of neutral sodium cometary tails. Finally, Section 7 summarizes this paper and briefly discusses the potential impact of Rosetta on our understanding of the formation conditions of comets.

2 Overview of the structure and composition of protostellar disks

Protostellar disks form during the star formation process. A molecular cloud core collapses under gravity forming a protostar in the center. As material continues to
fall inwards conservation of angular momentum results in the formation of a disk around the central protostar. In the early stages, the disks are gas-dominated with masses $\sim 1\%$ of the mass of the protostar. They rotate about the protostar with a Keplerian velocity profile. Disks contain a wide range of physical conditions, ranging from cold ($\sim 10 - 20$ K in the outer disk midplane) to hot (with gas temperatures up to several thousand K in the surface layers). Irradiation conditions also vary, with the surface layers experiencing the full force of the stellar UV field, but with little radiation reaching the midplane. Consequently there are a wide variety of chemical conditions depending on location within the disk. The star and disk formation process, and the chemistry and dynamics of disks are discussed in more detail in recent reviews by Dutrey et al. (2014); Pontoppidan et al. (2014) and Turner et al. (2014).

Disks can be very turbulent places. Turbulence is difficult to detect directly but evidence for its effects can be seen in the presence of small dust grains high above the disk midplane. Without turbulence these grains would quickly be removed from these high altitudes by settling (Dullemond and Dominik 2004). The dynamical motions also drive collisions between grains, which can lead to both grain growth, eventually forming planetesimals, and to creation of new small grains or dust. With grain growth, the grains will eventually reach a size at which their motions decouple from those of the gas and they begin to settle under the influence of gravity to the midplane, where they can undergo further collisions and growth to form planetesimals. The removal of grains from the surface layers allows the stellar UV to penetrate deeper into the disk, changing its composition. Turbulence can also drive chemical changes by bringing together material processed in different parts of the disk and conversely chemistry can affect the dynamics by controlling the ionization levels. Consequently dynamics and chemistry strongly influence each other (see Section 2.3.2).

In recent years protostellar disks have been an area of active research, with their molecular content, dust properties, and physical conditions being studied in detail. The observational work has been coupled with simulations that can interpret the observations and make predictions about those regions of the disks which are not amenable to direct observational study. This synergy of the observations and modeling work has revealed a wealth of information about the chemistry and physics of disks, and in turn has informed our understanding of the early history of our own solar system. This section reviews some of the recent observational and modeling work related to protostellar disks, concentrating on the early stages of evolution when the disk is still gas-dominated (the T Tauri phase).

2.1 The structure of a protostellar disk

Protostellar disks are characterized by strong radial and vertical gradients in temperature and density (Figure 1). The stellar UV field controls the temperature and hence the chemistry throughout much of the disk. The penetration of the UV into the disk is governed by the opacity which is mainly provided by dust absorption. In the surface layers, where the UV field is unattenuated, the photoelectric effect can efficiently heat the gas to much higher temperatures than the dust. Close to the midplane, where the UV field is reduced, the dust and grain temperatures are coupled.
These gradients in physical conditions lead to a wide variety of chemical environments (Figure 2). Models have found that outside of the water ice line the disk can be divided vertically into three chemically distinct layers. At the surface, the disk is dominated by the effects of UV. Ices are desorbed (either thermally or by photodesorption) and efficient photodissociation means that the gas is mainly composed of atoms and their ions. In the midplane most molecules are frozen out onto the surfaces of dust grains. Between these two layers is a region where the grains are warm enough to allow at least some molecules to thermally desorb and which is shielded enough that photodissociation is inefficient, allowing the molecules to survive once in the gas. It is this warm molecular layer that is detected in most gas phase observations of the outer regions of protostellar disks (see Section 2.2).

As the distance to the protostar decreases the disk warms up. In the midplane, the ice mantles will begin to desorb, with different molecules being released at different radii, depending on their binding energies – the most volatile species being released at larger radii than the less volatile ones. This results in a series of ice lines, which mark the transition from ice to vapor for each molecule (see Section 2.4). The icelines are curved, and in the outer disk are almost horizontal with the molecule existing as ice below this line and as gas above it (see Pontoppidan et al. 2014). Once inside the water iceline (within a few AU of the protostar) the molecules are all in the gas phase.

Figure 2 also illustrates some of the main physical processes that are active in disks. In particular, turbulence is a crucial process, enabling dust coagulation and driving chemical changes by mixing material processed under different physical conditions within the disk. What drives this turbulence is still a matter of debate but one of the main candidate processes is the magneto-rotational instability (MRI; e.g., Balbus and Hawley 1991; Hawley and Balbus 1991; Hawley et al. 1996) which can efficiently transport angular momentum in ionized accretion disks. A minimum level of ionization is required (in the minimum mass solar nebula the ionization fraction, x(e), must be $> 10^{-12}$ at 1 AU; Inutsuka and Sano 2005). This can be difficult to achieve in the shielded midplane regions of a protostellar disk, where the high column densities prevent UV and cosmic rays from penetrating. Hence
there are regions in the disk where the MRI is not active. These are known as “dead zones” and they are predicted to occur throughout much of the midplane in the planet-forming region. The approximate location of the dead zone in the disk shown in Figure 1 is indicated by the black contour line in the density plot.

The dependence of MRI-driven turbulence on the ionization means that there is a close link between the chemistry and the dynamics. The chemistry determines the ionization level and therefore the strength of the turbulence, but the dynamical motions themselves can drive chemical changes through mixing. This in turn affects the ionization and consequently the dynamics (e.g., Inutsuka and Sano 2005; Ilgner and Nelsen 2008; Ilgner and Nelson 2006; Turner et al. 2007). Hence the evolution of chemistry and dynamics are inextricably linked.

2.2 Observations of protostellar disks

Protostellar disks have been observed at wavelengths from the UV to the millimeter. Each wavelength can probe a different region of the disk. The majority of information about the molecular composition of disks comes from the infrared and the (sub)-millimeter. The disk is mostly optically thick in the infrared, therefore these observations are restricted to the warm gaseous region inside of the iceline or to the surface layers. At these wavelengths gaseous molecules can be observed in both absorption and emission. Ices can also be observed in absorption if illuminated by the central star in an edge-on disk, e.g., Pontoppidan et al. (2005), but the spectra can be difficult to interpret due to contributions from material outside of the disk. Near- and mid-infrared observations are most sensitive to the inner regions of disks, while the far-infrared can probe the surface layers at larger radii (see Henning and Semenov (2013).
Spitzer, operating in the mid-infrared, has shown that inside of the iceline the disk is chemically rich, with the terrestrial planet-forming region containing organics such as C$_2$H$_2$, HCN, as well as CO, CO$_2$, water and OH (e.g., Carr and Najita 2008; Salyk et al. 2008; Pontoppidan et al. 2010). The detected molecules are the result of a rich gas phase molecular chemistry, in a region where temperatures range from a few hundred kelvin up to 1000 K, and densities are high (> 10$^8$ cm$^{-3}$).

Spitzer observations of the gaseous inner disk have also been used to make inferences about the conditions beyond the iceline. For example, Najita et al. (2013) found the HCN/H$_2$O ratio inside of the iceline to be correlated with disk mass, suggesting that this ratio may reflect changes to the C/O ratio induced by the formation of icy planetesimals outside of the iceline. Outside of the iceline much of the oxygen is tied up in water ice, whereas the carbon is mainly in the gas phase in the form of more volatile molecules such as CO. Planetesimals outside of the iceline will form from grains coated with water ice. Since the planetesimals do not move about the disk as easily the small grains and the gas, this will trap the oxygen outside of the iceline and the material that moves inwards to the inside of the iceline will be enriched in carbon. Consequently the gaseous inner disk will be enriched in organics compared to water. Since more massive disks are likely to form planetesimals more efficiently than their lower mass counterparts they will have a higher C/O ratio inside of the iceline.

In the far-infrared, Herschel has detected a number of molecules including cold water in the outer disks of TW Hya (Hogerheijde et al. 2011) and DG Tau (Podio et al. 2013). In both cases the water emission comes from a region of the disk where grain temperatures are low enough for this molecule to be expected to be frozen onto dust grains. Its presence in the gas can be explained by photodesorption driven by the stellar UV field (Willacy and Langer 2000; Dominik et al. 2005). Hogerheijde et al. (2011) found that the predicted flux from photodesorbed water to be higher than observed, and suggested that the answer to this discrepancy could be grain coagulation. As the grains grow they decouple from the gas motions and sink towards the midplane, taking their ices with them. This removes water ice from the molecular layer, leaving less to be photodesorbed, and thus reducing the gas phase abundance.

Emission lines from some molecules in the disk can be optically thin in the (sub)-millimeter and so in theory these wavelengths can probe the whole of the disk. In practice, however, resolution constraints restrict these observations to $\gtrsim 30 – 100$ AU from the protostar, although ALMA allows observations to within a few AU of the closest protostars e.g. (ALMA Partnership et al. 2015). Observations by both interferometers and single dish facilities have detected the emission from molecules in the warm molecular region and their distributions can be mapped using interferometers. Relatively simple molecules have been observed including CO (and its isotopologues), H$_2$CO, CS, C$_2$H, HCN, CN, HNC, DCN, HC$_2$N, c-C$_3$H$_2$, N$_2$H$^+$, HCO$^+$, DCO$^+$, H$_2$D$^+$ (for a recent review see Dutrey et al. 2014). The fractional abundances relative to H$_2$ are generally lower than observed in molecular clouds because of the combined effect of depletion in the cold dense midplane and photodissociation in the surface layers. Both processes reduce the gas phase molecular abundances.

In addition to the composition of the disk, the detection of molecules can also provide information about the physical conditions under which they exist and the
chemical processes that are active (Table 1). Observations of molecular deuteration are especially important since they trace the thermal history of the disk (see also Section 3.1). Deuterated molecules, with their lower column densities, also provide a tracer of the gas in the midplane, e.g., ALMA observations of DCO$^+$ have been used to trace the location of the CO iceline in a Herbig AeBe disk (Mathews et al. 2013) and H$_2$D$^+$ is a potential tracer of the ionization level in the midplane (Ceccarelli et al. 2004).

| Process              | Molecules                                      |
|----------------------|------------------------------------------------|
| Temperature          | CO, $^{13}$CO, $^{17}$O, $^{18}$O, H$_2$CO, H$_2$O, OH, H$_2$ |
| Density              | CO, H$_2$CO, HCO$^+$, HC$_3$N                   |
| Ionization           | HCO$^+$, N$_2$H$^+$, CH$^+$                    |
| Velocity             | CO, CS, HC$_3$N                                |
| Photodissociation    | CN, HCN, HNC, OH, H$_2$O, C$_2$H               |
| Photodesorption      | H$_2$O, OH                                    |
| Grain surface chemistry | H$_2$CO, complex organics, CO$_2$, C$_2$H$_2$   |
| High temperature chemistry | Complex organics                               |
| Deuteration          | HD, DCO$^+$, DCN, H$_2$D$^+$                  |

Table 1 Molecules observed in protostellar disks and the processes they trace.

2.3 Protostellar disk modeling

Chemical models of protostellar disks vary in their complexity, but they all predict the three-layer structure beyond the iceline as shown in Figure 2. The chemical simulations require as their basis, a background disk model to provide the density, temperature and UV field as a function of location in the disk. These can be provided by hydrodynamical models of a 1+1D steady-state $\alpha$-disk model in vertical hydrostatic equilibrium such as those of D’Alessio et al. (1998) (see also Figure 1). These physical disk models are computationally intensive and are generally run independently of the chemical simulations. Current chemical models are based on detailed chemical kinetics and can contain hundreds of species (in both the gas and the ices) and thousands of reactions. These models include the ionizing effects of UV photons, cosmic rays, X-rays and short-lived radionuclides such as $^{26}$Al. Some models have also focused on isotopic chemistry such as deuterium (e.g., Aikawa and Herbst 1999; Woods and Willacy 2007; Willacy and Woods 2009; Kavelaars et al. 2011; Furuya et al. 2013; Albertsson et al. 2014) or carbon and oxygen isotopes (Woods and Willacy 2009; see also Section 3.2). The simplest models assume that the conditions in the disk did not change with time and ignore dynamical motions, although a few also consider advection (e.g., Willacy et al. 1998; Aikawa et al. 1999; Woods and Willacy 2007, 2009; Willacy and Woods 2009). More recent models have included processes such as grain growth.

1 http://www.cfa.harvard.edu/youngstars/dalessio/
and settling, e.g., Aikawa and Nomura (2006); Fogel et al. (2011); Vasyunin et al. (2011); Akimkin et al. (2013), dynamical motions, (e.g., Ilgner et al. 2004; Willacy et al. 2006; Semenov et al. 2006; Semenov and Wiebe 2011; Ilgner and Nelson 2006; Turner et al. 2006; Aikawa 2007; Hersant et al. 2009; Heinzeller et al. 2011) and hydrodynamical evolution of the disk structure itself (e.g., Dodson-Robinson et al. 2009; Yu et al. 2014). Here we highlight a few recent results.

2.3.1 Formation of complex organics in protostellar disks

Of particular interest to the chemistry of comets is the work of Walsh et al. (2014) who modeled the possible formation of complex organics through ice chemistry in protostellar disks. Figure 3 shows the range of abundances they find in ices located outside of 20 AU in a solar-type disk. The large molecules form by sequential addition of atoms to existing mantle molecules, e.g., ethylene glycol can form by adding hydrogen, oxygen and carbon atoms to CO:

\[
\text{CO} \xrightarrow{H} \text{HCO} \xrightarrow{C} \text{HC}_2\text{O} \xrightarrow{O} \text{OCCHO}
\]

[Charnley and Rodgers 2008, see Figure 6]. The models of Walsh et al. suggest that chemistry in disks will greatly increase the abundance of these large molecules relative to what can be created in the parent cloud. The predicted abundances relative to H\(_2\)O are consistent in many cases with observations of comets (Bockelée-Morvan et al. 2004; Crovisier 2006; Crovisier et al. 2004). If these molecules can be desorbed they will also be present in the disk gas, and should be detectable by ALMA providing a test of the model.

Fig. 3 Complex organics can be formed by chemistry in the ice mantles of dust grains in a protostellar disk. This figure shows the predicted range of abundances of some of these molecules relative to water ice (red lines) compared to those derived from cometary comae (blue lines; Bockelée-Morvan et al. 2004; Crovisier 2006; Crovisier et al. 2004). The green asterisks are the results from the initial molecular cloud model used as the input for the disk model. From Walsh et al. (2014).
2.3.2 The effects of dynamics on the chemistry

There are strong links between the disk chemistry and its dynamics. The temperature, density and radiation field in a disk control the chemistry, but the chemistry (and grain evolution) in turn controls the dynamics of the disk through the ionization structure. Models have found that better agreement with observations of the molecular layer in the outer disk can be obtained if vertical mixing is included, e.g., Willacy et al. (2006); Semenov et al. (2006), and more sophisticated models have now considered 2-D mixing (e.g., Heinzeller et al. 2011; Semenov and Wiebe 2011; Albertsson et al. 2014). Mixing smooths out chemical abundance gradients but also brings together species that might not otherwise exist in the same location, e.g., in the inner disk, vertical mixing combines OH and H$_2$O in a region where the temperature is high enough for the neutral-neutral reaction, OH + H$_2$O = H$_2$O + H to occur (Heinzeller et al. 2011). As a result the abundance of H$_2$O increases.

In addition to affecting molecular abundances, mixing also changes isotopic ratios. One problem with the non-mixing models is the high deuteration levels predicted in the comet formation zone, e.g., Willacy and Woods (2009). These reflect the high levels set in the parent molecular cloud, whereas those detected in comets tend to be somewhat lower. As D/H ratios track the thermal history of the material this suggests that at least some of the cometary ices formed at warmer temperatures than are found in the interstellar medium (ISM), e.g., Meier and Owen (1999). Figure 4 compares the observed molecular deuteration in cometary water with the predictions of several models. The non-mixing models uniformly over-predict HDO/H$_2$O, but the addition of mixing (either radial or vertical) brings the ratio into closer agreement with the observations (Bonev et al. 2014).

With vertical mixing this reduction in the deuteration of water is achieved by the destruction and reformation of ice mantles. When dust grains are mixed vertically they can reach the surface layers where high temperatures and/or UV fields desorb their ices. The water molecules are dissociated and the resulting oxygen, hydrogen and deuterium atoms travel back down towards the midplane into more shielded regions where they reform molecules. The molecular deuteration depends on the formation temperature. Since the disk is warmer than the 10 K parent molecular cloud core, the reformed water ice has a lower D/H ratio. Hence vertical mixing can reduce the initial interstellar D/H ratio in water ice ($\sim 10^{-2}$) to the level detected in comets ($\sim 10^{-4}$) (Figure 4 and see also Furuya et al. 2013). Willacy and Turner (2015) and Furuya et al. (2013) also predict radial variations in the deuteration of water. A radial gradient in HDO/H$_2$O is also found in the (radial) mixing models of Kavelaars et al. (2011).

2.3.3 Grain evolution

Grain-grain collisions in protostellar disks lead to efficient grain growth and ultimately result in the formation of planetesimals. As the grains grow they will decouple from the gas motions and sink towards the midplane. This removes grains from the upper layers of the disk, reducing the opacity and allowing UV to penetrate deeper, which increases photodissociation and heating of the newly irradiated layers, and moves the molecular layer closer to the midplane. An additional effect of grain growth is a reduction in the freezeout rate, which depends on the surface area of the grains. Figure 5 shows the effect on the chemistry of CO in the models
The composition of the protosolar disk and the formation conditions for comets

Fig. 4 Current disk models can only account for the observed D/H ratios in comets if they include mixing. The non-mixing models of Willacy and Woods (2009) and T. Millar (private communication) predict D/H ratios that are consistently higher than the observations. Albertsson et al. (2011) treat the evolution of the molecular cloud to the hot core stage only and predict (D/H)$_{\text{water}} > 10^{-3}$. Turbulent mixing is included in the disk models of Kavelaars et al. (2011) (radial mixing) and Willacy and Turner (2015) (vertical mixing at 5 AU) and these produce D/H$_{\text{water}}$ in agreement with the observations. Furuya et al. (2013) predict similar D/H ratios to Willacy and Turner (2015). References: (1) Eberhardt et al. (1995), (2) Hartogh et al. (2011), (3) Bockelé-Morvan et al. (2012), (4) Liu et al. (2013), (5) Bockelé-Morvan et al. (2013) (6a-d) Meier et al. (1998) and Crovisier et al. (2004), (7) Hutsemékers et al. (2008), (8) Biver et al. (2006), (9) Villanueva et al. (2009). Adapted from Bonev et al. (2014).

Fig. 5 The CO abundance distribution in a protostellar disk, showing the effects of grain growth. The highest abundances are shown in red. Grain growth reduces the opacity in the surface layers, allowing UV to penetrate further and pushing the molecular layer closer to the midplane. With 1 micron dust (left) the molecular layer is much thicker, and reaches closer to the midplane than in the model with 0.1 micron dust (center). On the right are the results from an evolving dust model where the grains are allowed to both grow and sediment. In this case the molecular layer is thicker than in the small grain model, and it moves closer to the midplane. From Vasyunin et al. (2011). ©AAS. Reproduced with permission.
of Vasyunin et al. (2011) (see also Aikawa and Nomura 2006; Walsh et al. 2014). Vasyunin et al. found that CO, CN and SO, as well as the ratio of C$_2$H$_2$/HCN, should be good tracers of the grain growth.

2.4 Gas-dust chemistry and icelines in protoplanetary disks

Icelines in the early nebula are defined as radii at which specific molecules undergo a shift in abundance from the gas phase to the condensed phase. There are three of specific interest related to H$_2$O, CO$_2$ and CO and these occur at radii where the temperature is approximately 150 K, 47 K and 20 K respectively ("Oberg et al. 2011). Recent observations by ALMA have allowed icelines to be identified in protoplanetary disks. For example, in HD 163926 (Qi et al. 2011; de Gregorio-Monsalvo et al. 2013; Mathews et al. 2013). The icelines associated with H$_2$O, CO$_2$, and CO are regions of dramatic chemical change and these could be reflected in the molecular composition of the gas and ice that is transported and mixed, both vertically and radially, in the evolving disk.

The chemical signature of such an environment may be imprinted in cometary and meteoritic composition if the Sun formed in a stellar cluster containing at least one massive star (Adams 2010; Mumma and Charnley 2011). Nearby OB stars strongly affect the positions of the CO$_2$ and CO icelines and the chemical structure of the disk (Walsh et al. 2013). The CO iceline occurs at tens of AU from the central protostar and is most accessible to observations at millimeter and sub-millimeter wavelengths. The physical conditions here most closely resemble those in the densest molecular cloud cores and so the related gas-dust chemistry could also be similar (Henning and Semenov 2013). This chemistry can lead to formation of many complex organic molecules on cold dust grains, as well as potentially large isotopic fractionation in both gas phase and grain-surface reactions.

For example, outside the CO iceline, CO-rich ices on dust grains can undergo (tunneling) addition reactions with H (and D) atoms accreted from the gas (Tielens 1983; Charnley 1997). Atoms of O, C and N will also accrete and these processes can lead to rich organic chemistry (Figure 6; Charnley 1997; Charnley and Rodgers 2008; Herbst and van Dishoeck 2009). Many of the proposed hydrogenation sequences have now been demonstrated in surface chemistry experiments (Watanabe and Kouchi 2008; Theulé et al. 2013) and such an ice chemistry explains the presence of the distinctive and inter-related suite of organic molecules detected around massive and low-mass protostars (in hot cores and hot corinos, respectively), where the ice mantles have been evaporated into the hot gas (Herbst and van Dishoeck 2009). Figure 6 indicates that many of the relevant organic molecules could also be present in comets and that, if correct, ketene, ethanol, propynal, acrolein, propionaldehyde and propanol should be searched for in suitably bright apparitions. Several of the organic compounds formed in the scheme of Figure 6 may be detectable in protoplanetary disks with ALMA. Vertical transport of ice-covered dust to the less well-shielded layers of the disk, as well as (inward) radial transport, can lead to these organic compounds being released by evaporation and/or photodesorption. This is probably the origin of the H$_2$CO detected in HD 163296, which lies outside of the CO iceline at about 160 AU (Qi et al. 2013b).
2.5 Two distinct reservoirs of water ice in the protosolar nebula

Formation scenarios of the protosolar nebula (PSN) invoke two main reservoirs of ices present in the disk midplane and that took part in the production of icy planetesimals (see Figure 7). The first reservoir, located in the inner region of the
PSN, contains ices (dominated by H$_2$O, CO, CO$_2$, CH$_4$, N$_2$ and NH$_3$) originating from the ISM, which, due to their near solar vicinity, were initially vaporized. The ice vaporization distance never exceeds $\sim 30$ AU from the Sun, depending on the total source luminosity and characteristics of the collapsing cloud (Chick and Cassen 1997). With time, the decrease of temperature and pressure conditions allowed the water in this reservoir to condense at $\sim 150$ K (at typical nebula pressure conditions) in the form of microscopic crystalline ice (Kouchi et al. 1994), leaving negligible water in the vapor phase to condense at lower temperatures where amorphous ice would be expected. It is postulated that a substantial fraction of the volatile species were trapped as clathrates during this condensation phase as long as free water ice was available within 30 AU in the outer solar nebula (Mousis et al. 2000) and there was enough time to overcome the kinetics of clathration. On the other hand, the remaining volatiles that were not enclathrated (due to the lack of available water ice or a low kinetics of clathration) probably formed pure condensates at lower temperatures in this part of the nebula (Mousis et al. 2012a,b). The other reservoir, located at larger heliocentric distances, is composed of ices originating from the ISM that did not vaporize when entering into the disk. In this reservoir, water ice was essentially in the amorphous form and the other volatiles remained trapped in the amorphous matrix (Owen et al. 1999; Notesco and Bar-Nun 2005).

Interestingly, it has been shown recently that amorphous ice could form from photodesorption and freeze-out of water molecules near the surface layers of the PSN (Ciesla 2014). In these conditions, the transport of the icy grains throughout the outer solar nebula would lead crystalline ice to be lost and reformed as amorphous ice. In turn, amorphous ice generated on the disk’s surface could be transformed to crystalline ice as it migrates through the protoplanetary disk (Ciesla 2014). This cycling scenario needs to be further investigated if one wants to assess its role in the formation or destruction of the two original icy reservoirs. Also, the different transport mechanisms must be simultaneously taken into account at the different stages of the disk’s evolution to depict the abundance of water ice in the two reservoirs as a function of the heliocentric distance. For example, as described in Sec. 2.3.2, it has been shown that vertical mixing can decrease the abundance of water inside the $\sim 15$ AU of the disk. In contrast, it has also been proposed that the outward vapor diffusion induced by the presence of the iceline with its subsequent condensation in a narrow location (with a width around 0.5 AU) would deplete the region inside the iceline of vapor down to subsolar values, and increase substantially the ice abundance at the iceline location by factors larger than 10 times the protosolar value (Ali-Dib et al. 2014). In addition, recent works depicting the chemical evolution of both solids and gases from the pre-stellar core to the protostar and circumstellar disk phases (Visser et al. 2009, 2011; Harsono et al. 2013) should be taken into account to update the representation of the two water ice reservoirs present in the PSN. In these cases, midplane ices may have experienced multiple desorption events during the evolution of the disk. Nevertheless, if one follows this classical picture, icy solids that formed at heliocentric distances less than $\sim 30$ AU mainly agglomerated from a mixture of clathrates and pure condensates, whose ratio depends on the amount of available crystalline water and its clathration efficiency (Mousis et al. 2009b). In contrast, solids produced at higher heliocentric distances (i.e., in the cold outer part of the solar nebula) were formed from primordial amorphous ice originating from the ISM. Therefore,
The composition of the protosolar disk and the formation conditions for comets

Infalling material at 30 AU

Fig. 7 Two reservoirs of ices in the PSN. A first reservoir (cold zone), located at distances higher than ∼30 AU, is constituted from amorphous ices coming from ISM. The second reservoir (hot zone), located within the ∼30 AU, is made from volatiles initially in the form of amorphous ices that were transported from ISM towards the inner and hot part of the disk. When reaching regions with temperatures higher than ∼150 K, these ices vaporized. During the cooling of the disk, volatiles located in the inner 30 AU condensed again but in crystalline forms, including both pure crystalline ices and clathrates (see text).

depending where the outer solar system bodies formed, clathrates may have been agglomerated in comets (Marboeuf et al. 2010, 2011, 2012), and in the building blocks of the giant planets (Gautier et al. 2001; Alibert et al. 2005a,b; Mouiss et al. 2009a, 2012a) and in their surrounding satellite systems (Lunine and Stevenson 1985; Mouiss 2004; Mouiss and Gautier 2004; Mouiss et al. 2009a). Regardless the possible presence of clathrates, there is today indirect evidence that comets formed from crystalline water ice because their deuterium-to-hydrogen ratios measured in H$_2$O are substantially lower (∼6–12 times the protosolar value) than the most deuterium-rich primitive meteorite (∼35 times the protosolar value), whose level of deuteration is expected to be close to the one found in the ice infalling from the presolar cloud (Kavelaars et al. 2011; Hartogh et al. 2011; Ceccarelli et al. 2014). This difference can only been explained by the isotopic exchange between the deuterium-rich water and molecular hydrogen in the gas phase (Kavelaars et al. 2011; Hartogh et al. 2011; Ceccarelli et al. 2014). Hence, in order to enable this reaction, amorphous ice entering the disk from ISM must have been vaporized prior recondensing again in crystalline form, otherwise the isotopic exchange did not occur.

3 Fractionation in protostellar disks

3.1 The deuterium fractionation: The link between comets and the earliest phases of the solar system and interstellar chemistry

The origin of the volatile ices found in comets is a central issue for understanding the formation and early evolution of the solar system. There are three possibilities. First, one view is that all the ices are comprised of pristine interstellar molecules (e.g. Irvine et al. 1996). There is indeed a strong similarity with the many known interstellar molecules believed to form on interstellar grains, although there are differences in the relative abundances of the volatile ices (Mumma and Charnley 2011). Until recently this idea was widely considered to be untenable (see Section 3.1.2 and Cleeves et al. 2014a). Second, at the other extreme, it has been advocated that any interstellar molecules experienced total obliteration and "chem-
ical reset’ in the disk (Pontoppidan & Blevins 2014). In this case all the volatile molecules observed in comets must have been formed by nebular chemistry.

Comets and meteorites contain both high-temperature processed materials, such as crystalline silicates, and matter that apparently formed at very low temperatures, as indicated by the isotopic enrichments measured in D and $^{15}$N (Mumma & Charnley 2011). These characteristics are most naturally explained by the large-scale turbulent transport of dust and gas from different nebular chemical environments into the comet-formation zones (Ciesla and Charnley 2006). The recognition that non-equilibrium models of nebular chemistry are essential, particularly in the comet-forming regions, has motivated many detailed studies of protoplanetary disk chemistry, as recently reviewed by Henning and Semenov (2013) (see also Section 2.3.2). Turbulence leads to an outward radial diffusion of material from the hot inner nebula and an inwards advection of material from the cooler outer nebula. It also leads to vertical mixing and thus to midplane material experiencing the enhanced UV and X-ray fluxes of the upper disk layers (Glassgold et al. 1997). Hence, in this third partially-mixed scenario some specific volatile molecules and isotopic ratios could be remnant interstellar matter, or derive from warm and cold regions of the nebula (as reviewed by Pontoppidan et al. 2014). Deciding between such molecular origins has been problematic.

3.1.1 The molecular deuteration process

Molecular deuteration provides a potential clue to the history of cometary volatiles. Although deuterium in the Universe is a tiny fraction with respect to the hydrogen ($D/H = [2.535 \pm 0.05] \times 10^{-5}$; Pettini & Cooke 2012), its abundance in interstellar molecules has an enormous diagnostic power, because it may be much larger that the elemental D/H ratio. This generally occurs in cold ($\leq 30-50$ K) environments and in trace species, namely molecules with abundances smaller than $\sim 10^{-5}$.

The first key point to keep in mind is that, in cold gas, the major deuterium reservoir is HD and only a small fraction of deuterium is locked in other species. An exception to this rule is represented by the water, whose abundance can reach $\sim 10^{-4}$ and, consequently, water might trap a substantial fraction of deuterium.

The second key point is that the deuterium atoms are “extracted” from the HD molecules because the cosmic rays that hit H and $H_2$ create $H_3^+$ ions, which in turn react with HD and form $H_2D^+$ ions.

\begin{equation}
H_3^+ + HD = H_2D^+ + H_2 + 220K
\end{equation}

(see also Figure 3). The reverse reaction has a small energy barrier, which cannot be overcome at low ($\leq 20-30$ K) temperatures, so that the $H_2D^+/H_3^+$ abundance ratio becomes larger than the D/H elemental ratio. The kinetics of Reaction 1 at low temperatures is also strongly influenced by the spin state of the reacting hydrogen molecules, i.e., whether they are predominantly in the ortho or para states. The higher internal energy of ortho-$H_2$ means that the reverse process can be driven efficiently if the $H_2$ ortho/para ratio is large. Thus, ortho-$H_2$ acts as a “poison” for interstellar deuteration which requires that most of the $H_2$ be converted to para-$H_2$ (see Pagani et al. 2011). Reaction 1 is the first step towards the deuteration of the trace species, as $H_2D^+$ reacts with all neutral molecules and passes, once every three times, the D atom to the product species. This is
The composition of the protosolar disk and the formation conditions for comets

Fig. 8 Schema of the molecular deuteration process in cold gas (Ceccarelli et al. 2014). Molecular deuteration occurs through three basic steps: 1) formation of $\text{H}_3^+$ ions from the interaction of cosmic rays with H and $\text{H}_2$; 2) formation of $\text{H}_2\text{D}^+$ ($\text{HD}_2^+$ and $\text{D}_2^+$) from the reaction of $\text{H}_3^+$ ($\text{H}_2\text{D}^+$ and $\text{HD}_2^+$) with HD; 3) formation of other D-bearing molecules from reactions with $\text{H}_2\text{D}^+$ ($\text{HD}_3^+$ and $\text{D}_3^+$) in the gas phase (Step 3a) and on the grain mantles (Step 3b).

schematically shown in Figure 8. A similar process can also occur through the CH$_3^+$ ions at slightly higher temperatures.

In practice, therefore, a molecular deuteration enhanced with respect to the D/H elemental abundance is a thermometer of the gas temperature.

The third and final key point to keep in mind is that neutral molecules can freeze-out into the grain mantles and, in this way, they can be transmitted through the different stages of the star and planet formation. A sketch of this overall process is shown in Figure 8. Therefore, ice species formed during the very first phases might be conserved and “reappear” in the gas at much later stages: for example when comets and planets are formed. Conversely, the molecular deuteration in comets and planets can help us to understand in what epoch the relevant species was formed, at least in principle. In this sense, molecular deuteration can be thought of as an Ariadne’s thread which links the various phases of solar system evolution. For a more detailed review see Ceccarelli et al. (2014).

3.1.2 Deuteration in the solar nebula

Deuteration in disks is driven mainly via the reaction of $\text{H}_2\text{D}^+$ ions (Reaction 1). At low temperatures the reverse process is inhibited and this leads to enhanced deuteration in gaseous molecules through ion-molecule isotope exchange reactions. Elevated atomic D/H ratios also pertain and lead to significant deuteration in
Fig. 9  Schematic summary of the different phases of the formation of a solar-type star and planets, highlighting the formation and fate of deuterated molecules (adapted from Ceccarelli et al. (2014). Likely, the same process gave birth to the solar system.

Grain-surface reactions, such as those depicted in Figure 10 and in the surface reduction of accreted O atoms to form HDO, condensation of CO leads to enhanced abundances of gaseous H$_2$D$^+$ and to significant deuterium fractionation beyond the CO iceline, as traced by DCO$^+$. Enhanced D/H ratios in simple molecules, such as water, ammonia and methane, can be produced both in ion-molecule reactions and grain-surface additions of H and D to accreted atoms (Ceccarelli et al. 2014). At the CO iceline the grain-surface chemistry depicted in Figure 10 can lead to very high D/H ratios and multiply deuterated molecules (Charnley et al. 1997; Ceccarelli et al. 2014). Observations show that simple deuterated species like DCN and by DCO$^+$ are present in disks (Qi et al. 2008) and we would expect that more complex fractionated organics, e.g., HDCO, D$_2$CO, CH$_3$OD and CH$_2$DOH, to also be present. Between the CO and CO$_2$ icelines (see Section 2.4), large deuterium enrichments can persist in organic molecules since ion-molecule deuteration involving C$_2$HD$^+$ and CH$_2$D$^+$ can persist up to higher temperatures (∼ 70 K and ∼ 50 K) than H$_2$D$^+$ (∼ 30K) (Millar et al. 1989). It is possible that the D/H ratios of several organic molecules can become re-fractionated even in the warm inner nebula, prior to the condensation of acetylene and methane, both of which will occur outside the CO$_2$ iceline. This means that deuteration can still occur in cold gas that is mixed radially inwards, and be initiated in warmer gas coming from the inner nebula (Willacy 2007; Albertsson et al. 2014).

However, the assumption of interstellar rates of cosmic-ray driven ionization (ζ) in the cold regions of the outer midplane has recently been called into question. Cleeves et al. (2013) showed that the effects of either deflection by stellar winds or magnetic field topology (through the magnetic mirror effect) could significantly
Figure 10 Grain surface deuteration pathways associated with methanol formation from hydrogenation of CO ice. The probabilities for each reaction pathway, \( P_i \), assume a statistical steady state and are related to the atomic D/H ratio of the accreting gas and the energy barrier for addition of an H or D atom (from Charnley et al. 1997).

Cleeves et al. (2014a) developed this scenario further by pointing out that effectively extinguishing cosmic ray ionization from the nebular midplane would greatly reduce the efficacy of Reaction 1 and consequently the generation of enhanced molecular D/H ratios by nebular ion-molecule chemistry. They demonstrated that if the enhanced interstellar HDO/H\(_2\)O ratios initially available were lost, then nebular chemistry could not regenerate those currently measured in the solar system, concluding that as much as 50% of the Earth’s oceans and perhaps all cometary water is of interstellar origin. Low values of \( \zeta \) will also reduce the atomic D/H ratio in the disk midplane and so reduce the D/H ratios and the level of multiple deuteration in the molecules formed in the scheme of Figure 10. This may account for the upper limits on HDCO, CH\(_2\)DOH and CH\(_3\)DOD in comet Hale-Bopp (Crovisier et al. 2004) relative to the large observed interstellar D/H ratios measured in these molecules, which could have been lost or modified upon incorporation into the nebula. Although these molecules could be partially reformed in the cold outer nebula, this process, and any related deuteration, may be less efficient than in...
molecular clouds. Cometary CH$_3$OH/ H$_2$O ratios are typically $\sim 1 - 4\%$, whereas $\sim 5 - 15\%$ are typical for the ISM, and can be as high as 40% [Mumma and Charnley 2011]. Cleeves et al. (2014b) also modeled the general protoplanetary disk chemistry for a variety of assumed $\zeta$ values and identified molecular ions that may eventually allow the ionization structure to be determined. If these spatial distributions are observable with ALMA they could provide valuable information on the actual role of cosmic rays in nebular chemistry.

### 3.1.3 The message from molecular deuteration

Figure [11] shows the molecular deuteration in the different objects of the solar system and in objects which are believed to eventually form a solar-like system. It gives us the following major messages:

- All reported solar system objects, including Earth, possess a molecular deuteration larger, by more than a factor 10, than the D/H elemental ratio of the environment where the Sun was born, and that is represented by the line marked with “Proto Solar Nebula”. This implies that the process of molecular deuteration was at work and it was at least partially inherited from the earlier phases of the solar system formation era.
There is a general trend, with the objects that formed earlier being also the ones with the largest molecular deuteration, in agreement with the scenario that the process started in very cold conditions. This implies that there was only a partial transmission between the subsequent steps of Figure 9. Besides, earlier objects in the plot also correspond to material whose distance from the central heating source is larger. Therefore, the trend also tells us that there was not a general and substantial remixing of the different regions (outer and inner, where molecular deuteration is different because of the different temperatures). Exactly how much material was passed through each step or remixed remains, however, unanswered.

The organic matter is systematically more enriched in deuterium atoms than the water. This is true in protostellar objects, comets and meteorites. However, while this result is robust for the protostellar objects and meteorites, where several measurements of the deuteration have been made, it is less certain for the comets, where so far the only organic species for which molecular deuteration has been measured is HCN. The reason for this dichotomy in the molecular deuteration is completely unclear. In protostellar objects, it is believed that this reflects the different phases of formation of water and organics, with water being formed very early when the gas is not very cold, while the organics were synthesized later from frozen CO. It is possible that a similar process is at the origin of the different deuteration in meteorites and comets. Alternatively, they may conserve a “core” of the protostellar molecular deuteration.

Molecular deuteration in a given molecule has a strong temperature dependence at low temperatures. This is true for both gas phase and icy grain surface formation. In ice mantles the higher binding energy of deuteration atoms at < 20 K leads to significant enhancements of deuterated species that form via hydrogenation reactions (see also Figure 10). These low temperatures correspond to what modelers predict for the cold midplane in the giant planet formation zone of the protoplanetary disk. Measurements of the molecular deuteration in comets provide a potential tracer of the thermal conditions under which they formed. For further discussion see Bockelée-Morvan et al. (this volume).

3.1.4 A model of the deuteration of water in the early stages of the evolution of the solar nebula

Molecular deuteration potentially provides a means of tracing the history of solar system material and its links to the parent molecular cloud. In this section we present the results of a specific model of the evolution of the deuteration of water during the collapse of the parent molecular cloud, since the composition of the protosolar nebula depends in part on the chemical history of this core (Figure 12).
The evolution of the core has been modeled recently by Aikawa et al. (2012); Wakelam et al. (2014); Taquet et al. (2014) and Tornow et al. (2014a,b).

In these models the cloud core is initially described as a Bonner-Ebert sphere (Aikawa et al. 2008) which is embedded in a constant density gas-shell with a visual extinction of up to 4 magnitudes (Wakelam et al. 2014), and irradiated by an external UV field. It is allowed to evolve for between 1 and 6 Myrs (Tornow et al. 2014a). Both the gas phase and grain chemistry is followed. Since the core is only partially shielded, photo-chemistry has a clear effect on the radial profile of the relative water abundance in the outer core region (Figure 13). At the end of the quasi-stationary core stage the temperature of the gas and dust decreases to 8 – 9 K in the center for boundary values between 13 – 15 K. Consequently, water ice builds up on the cold grains, reaching a fairly constant relative abundance of $\sim 10^{-4}$ within $10^5 – 10^6$ years. In contrast, Figure 13 shows a clear radial gradient for the gas phase water abundance. Relatively low ratios (about 0.01 – 10 ppm) of gas to dust phase abundance were obtained for the outer core range with up to 25 ppm in the inner region. Caselli et al. (2012) determined higher ratios due to a higher efficiency of water photodesorption; a consequence of the larger grain size and smaller photodesorption yield used in the Tornow et al. models. If the deuterium chemistry is included according to the method of Albertsson et al. (2013) the temporal and radial distribution of the gas phase HDO/H$_2$O ratio can be derived. It reaches a maximum of about 0.02 in the medium shielded core region, and a minimum of $10^{-4}$ in the low shielded outer boundary region (Figure 14). In contrast, the dust phase ratio of HDO/H$_2$O is clearly larger and only varies between 0.01 and 0.1. Its maximum is located in the well shielded inner core region at very early times ($\sim (2 –3) \times 10^5$ years).

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2 Nahoon_public_april.tar.gz at http://kida.obs.u-bordeaux1.fr/models/
3 disk_chemistry_OSU08ggs_UV.zip at http://www.mpia.de/homes/semenov
Fig. 13 Relative gas phase abundance of ater versus time and radius (1 kAU = 10^3 AU). The intensity of the standard interstellar radiation field (IRSF) is scaled by Rad = 1, 2, 3. Two grain distributions are used. MRN refers to Mathis et al. (1977) and Zubko refers to Zubko et al. (2004). The mean grain size is 5.5 \times 10^{-5} \text{ cm} and 3.5 \times 10^{-5} \text{ cm} for MRN and Zubko respectively. The cosmic ray ionization rate is taken to be 10^{-17} \text{ s}^{-1}. The related gas and grain chemistry is modeled using a combined code (Tornow et al. 2014a).

The collapse of the cloud core forms a protostar surrounded by an optically thick disk and an outer envelope (Saigo et al. 2008; Schönke and Tscharnuter 2011). During this stage the chemical evolution can be followed along a Lagrangian path from the outer core boundary to the center of the core (Tornow et al. 2014b).
Figure 15 shows the HDO/H$_2$O ratio along each member of the Lagrangian path set. Water formed in the quasi-stationary core stage is transported into the region of terrestrial planet formation. Signatures of large-scale, supersonic water inflows have been seen with Herschel (Mottram et al. 2013) who determined that these flows occur on the core scale ($\leq 10^4$ AU), e.g., IRAS 15398 and L1527 or at least on the envelope scale ($\geq 3 \times 10^4$ AU), e.g., NGC1333-IRAS4A. Based on their data Mottram et al. suggest an outside-in collapse, in agreement with the model shown in Figure 15.

The gas phase deuteration of water varies widely, depending on location and evolutionary time. At 1 AU it ranges from 0.001 for 0.1 – 0.2 Myrs, to 0.1 for 0.3 Myrs and to 0.03 for times later than 0.4 Myrs. A minimum HDO/H$_2$O ratio near 0.001 extends from the radial region around 0.1 AU at early collapse times to nearly 9 AU at later times. The predicted HDO/H$_2$O ratios of Wakelam et al. (2014) and Tornow et al. (2014b) are slightly lower than those of Aikawa et al. (2012) but larger than the ratios observed by Persson et al. (2014). An overestimation of the water deuteration may be caused by a low collapse temperature which would suggest perfect cooling. Including the ortho-to-para ratio of H$_2$ can also lead to lower HDO/H$_2$O ratios (Taquet et al. 2013, 2014). In addition, water formed during the quasi-stationary core stage must be destroyed by the UV radiation of the interstellar radiation field, nearby stellar sources or X-rays (producing secondary UV photons in a similar way as cosmic rays) and UV photons from the protostar (France et al. 2014). As a result, photodissociation of deuterium-rich water molecules (HDO + h$\nu$ $\rightarrow$ D + OH, or H + OD), together with 1-D radial or vertical mixing enable a subsequent reformation of water by hot neutral reactions in the inner disk plane (Furuya et al. 2013).
3.2 Fractionation of carbon and nitrogen in the solar nebula

Isotopic enrichment of $^{15}\text{N}$ in cometary organics (the nitriles CN and HCN) and ammonia is commonly observed (Jehin et al. 2009, Bockelée-Morvan et al., this issues). Chemical models have shown that fractionation in low-temperature ion-molecule reactions can produce large $^{15}\text{N}$ enrichments in interstellar clouds (Rodgers and Charnley 2008; Wirström et al. 2012). However, a recent study by Roueff et al. (2015) has demonstrated that several key reactions may in fact possess energy barriers, leading to doubts as to the efficacy of ion-molecule processes in N fractionation. Of particular interest for comets is that $^{15}\text{N}$ fractionation in the outer nebula should be similarly impaired by low values of the cosmic ray ionization rate, $\zeta$. Thus, as found by Cleeves et al. (2014a) for water deuteration, the low $^{14}\text{N}/^{15}\text{N}$ ratios commonly measured in both Oort Cloud and Jupiter Family comets is strongly suggestive of an interstellar origin.

The $^{12}\text{C}/^{13}\text{C}$ ratios measured thus far in simple cometary molecules (CO, HCN, CN) are solar (Jehin et al. 2009) and Bockelee-Morvan (these proceedings). In regions of the nebula where water has condensed, the gaseous C/O elemental ratio can increase to order unity (Oberg et al. 2011). Because water, OH and O$_2$ recycle carbon nuclei back into CO (Langer et al. 1984), ion-molecule fractionation in regions with C/O $\sim$ 1 will act instead to channel carbon nuclei into simple organic molecules. Thus, as the sole carbon fractionation process involves the $^{13}\text{C}^+$ exchange reaction with $^{12}\text{CO}$ (Langer et al. 1984), CO condensation may be the reason for the almost uniformity of cometary $^{12}\text{C}/^{13}\text{C}$ ratios measured to date. On the other hand, the condensed CO is predicted to produce distinctive $^{12}\text{C}/^{13}\text{C}$ fractionation patterns in the organic molecules formed in the grain-surface reaction scheme of Figure 6 (Charnley et al. 2004). These involve

Fig. 15 HDO/H$_2$O ratio in the gas phase on Lagrangian paths. At 1 AU the HDO/H$_2$O ratio varies between $10^{-3}$ and 0.1, while in the dust phase it is much smoother, and only varies between 0.03 and 0.05 (Tornow et al. 2014a,b).
molecules whose $^{13}$C isotopologues have not yet been detected in comets (e.g., H$_2$CO, CH$_3$OH) but for which the predicted fractionation has been confirmed by observations of protostellar envelopes (Wiström et al. 2011). In this case, once again, ion-molecule pathways to $^{13}$C fractionation in the nebula are predicated on the presence of significant flux of cosmic rays.

If ion-molecule reactions indeed play a greatly reduced role in the cooler parts of the disk then the major sources of nebular fractionation will involve UV photochemistry. Isotope selective photodissociation of CO has been proposed as an explanation of the oxygen isotope ratios measured in meteorites (Lyons and Young 2005). Isotopic enrichment of $^{13}$C, $^{15}$N and D though isotope-selective photodissociation of isotopologues of CO, N$_2$ and HD has also been considered (Woods and Willacy 2009; Heays et al. 2014; Cleeves et al. 2014a). However, these processes will only occur in thin upper layers of the disk and so will be negligible at the midplane. Thus, to be viable, these processes require efficient downward vertical mixing in the disk. On the other hand, ISP occurring in the outer layers of the presolar core, where the interstellar UV field is attenuated, could in principle produce isotopic anomalies in the oxygen and carbon isotopes (Yurimoto and Kuramoto 2004; Lee et al. 2008; Charnley and Rodgers 2009). This environment could also ultimately produce significant fractionation in nitrogen-bearing species but this has not yet been demonstrated quantitatively.

3.3 Solar nebula chemistry and the origin of oxygen isotopic anomalies in the solar system

Oxygen is a very common element that is a major constituent of many minerals and rocks. It exists in three stable isotopes: $^{16}$O, $^{17}$O and $^{18}$O. The fractionation between these isotopes is not consistent across all solar system samples, as might be expected if the oxygen isotopes were homogeneously distributed across the solar nebula, and if only mass-dependent fractionation processes were active. Figure 16 is a three-isotope plot of the oxygen isotopic compositions of primitive solar system materials such as chondrites, chondrules, shows that their relative isotopic compositions are different compared to those of terrestrial samples (McKeegan et al. 2011; Yurimoto et al. 2007; Clayton 1993). The terrestrial line is characteristic of mass-dependent fractionation, where $^{18}$O is twice as fractionated as $^{17}$O relative to $^{16}$O, whereas the steeper, slope -1, line, of the primitive materials suggests it is the result of mass-independent fractionation.

There are three possible mechanisms that could achieve this mass-independent effect (Yurimoto et al. 2007). Firstly the isotope anomalies could represent an inherited heterogeneity in solar system materials, where the solids are rich in $^{16}$O and the gas in the rarer isotopes. This could occur either from nucleosynthesis in other stars (Clayton et al. 1973) or from nuclear reactions triggered by energetic particles produced by the Sun or by galactic cosmic rays (Lee 1978). These nuclear processes could produce fluctuations in the $^{16}$O abundance. The lack of observed isotopic anomalies in other elements suggests this mechanism is unlikely (Clayton 1993). Additionally, $^{16}$O presolar grains are rare in meteorites (Nagashima et al. 2004; Nittler 2003).

A second suggestion is chemical fractionation within the solar nebula. Differences in the reaction rates of symmetric molecules, e.g. $^{16}$O$^{16}$O, compared to
Fig. 16 A three-isotope plot for oxygen in the solar system. The Earth and Moon lie along a fractionation line with slope 0.52. This line is consistent with fractionation of a single homogeneous source, by processes that depend on the mass of the isotopes, so that $^{18}$O is nearly twice as fractionated as $^{17}$O. Data from meteorites and the Genesis mission falls along a slope-1 line indicating they contain material that has undergone some kind of mass-independent fractionation. The Genesis results suggest that much of the rocky material in the inner solar system was enriched in $^{17}$O and $^{18}$O relative to $^{16}$O before the accretion of the first planetesimals. Hence rocky planets are not made of the average nebular material that formed the Sun. From McKeegan et al. (2011) (reprinted with permission).

asymmetric ones, e.g. $^{16}$O$^{17}$O, can lead to the observed isotopic distributions, since the most abundant isotope is almost entirely contained in the symmetric molecules, while the rarer isotopes are in the asymmetric ones (Heidenreich and Thiemens 1983). However, the exact reactions by which this effect might be achieved are uncertain (Yurimoto et al. 2007).

The final possibility is isotope-selective photodissociation of CO, either in the parent cloud (Yurimoto and Kuramoto 2004) or at the surface of the protosolar disk (Young 2007; Lyons and Young 2005) could also result in enhancements of the rare isotopes of oxygen. CO dissociates when irradiated by far-ultraviolet photons with wavelengths between 91.2 nm and 110 nm. The dissociation process occurs in two steps, with CO first entering a bound excited state before dissociating. As a result the CO absorption spectrum consists of a large number of narrow lines at specific wavelengths determined by the vibrational and rotational levels involved. The change in mass between C$^{16}$O and its isotopologues C$^{17}$O and C$^{18}$O slightly
shifts the lines in the absorption spectra, and consequently the spectra for the different isotopologues do not overlap. Hence when CO is dissociated, either in the protostellar disk or in the molecular cloud core, the different isotopologues do not shield each other (see, for example, Visser et al. 2009). C\textsuperscript{16}O will become optically thick closest to the surface, in a region where C\textsuperscript{17}O and C\textsuperscript{18}O can still dissociate. When coupled with turbulent mixing that can transport the \textsuperscript{17}O and \textsuperscript{18}O towards the colder, shielded midplane of the disk this can result in the formation of water molecules that are enhanced in the rarer isotopes and which could later be incorporated into CAIs and other meteoritic material.

The in situ measurements that Rosetta will make will will directly probe the isotopic composition of the volatile component of a comet. This will test the viability of the potential causes of the oxygen isotopic anomalies and provide new clues as to the origin of the cometary material and the conditions under which it formed.

4 The origin of the nitrogen deficiency in comets

Nitrogen is one of the most abundant elements in the PSN (Asplund et al. 2009). Its abundance was measured in the giant planets atmospheres in which its stable thermodynamic form is NH\textsubscript{3}. The nitrogen abundance is found moderately to substantially enriched, compared to its solar value, in Saturn and Jupiter (Mousis et al. 2014), but is loosely constrained in Uranus and Neptune (de Pater et al. 1989; de Pater and Richmond 1989; Ali-Dib et al. 2014). A puzzling revelation was the apparent severe depletion of nitrogen in comets. Mass spectrometry measurements of 1P/Halley’s dust composition allowed to find that the nitrogen-to-oxygen ratio is depleted by a factor $\sim 3$ with respect to the solar value (Jessberger and Kissel 1991). Subsequent analysis showed that the ice part of the comet is depleted in nitrogen by a factor up to 75 (Wyckoff et al. 1991). This apparent depletion appears controversial because the N\textsubscript{2} molecule, thought to be the main nitrogen-bearing volatile in the whole PSN (Lewis and Prinn 1980; Mousis et al. 2002), has a molecular mass close to that of the very abundant CO molecule, implying that the mass spectrometer aboard the Giotto spacecraft did not have enough resolution to separate the two molecules (Eberhardt et al. 1987).

More recently, Cochran et al. (2000) used the 2DCoude spectrograph on the 2.7-m Harlan J. Smith telescope of McDonald Observatory to observe the N\textsuperscript{2+}/CO\textsuperscript{+} in comets 22P/1995 S1 (deVico) and C/1995 O1 (Hale-Bopp), and found values around $\sim 1 \times 10^{-4}$. These values hint at a severe depletion in N\textsubscript{2} since the N\textsubscript{2+}/CO\textsuperscript{+} ratio is usually equal to N\textsubscript{2}/CO. This is surprising specially that N\textsubscript{2} and CO have very close condensation temperatures (Fray and Schmitt 2009). Moreover, efforts were made to find N\textsubscript{2} using the Far Ultraviolet Spectroscopic Explorer, but with no success (Bockelée-Morvan et al. 2004). All these observations, and many more, motivated theorists to try to explain this depletion.

In Kro et al. (2003), the authors assumed that volatiles are accreted by comets as clathrates (Lunine and Stevenson 1985), and explain the nitrogen depletion by the low efficiency of N\textsubscript{2} trapping in water ices. One of this work’s conclusions is that N\textsubscript{2} can be trapped only if water ice is present in abundance higher than 2.8 $\times$ solar value. This is due to the relative ease by which CO can be trapped in comparison with N\textsubscript{2}, so this last can be clathrated only if there is enough water left after all the present CO was captured. Since N\textsubscript{2} is trapped in crystalline
water ice at temperatures lower than CO (Mousis et al. 2010), their work implied that comets might have formed at temperatures higher than 45 K, allowing the clathration of CO but not N$_2$. This temperature lower limit is one of shortcomings of this model since in many protoplanetary disks models, temperature is thought to decrease below this value (Hueso and Guillot 2005). The recent observation of the CO iceline in TW Hya also supports these models (Qi et al. 2013a). Another caveat is the reliability of the Iro et al. (2003)'s determination of the N$_2$/CO ratio in clathrates since the potential parameters describing the guest–clathrate interaction in their statistical model correspond to guest–guest interactions (case of pure solutions) and not to the guest–water interactions usually used in such models (Mousis et al. 2012b). Also, the model is unable to explain the thick N$_2$ ice cover on Triton and Pluto, both bodies thought to be formed in the comets region (Lellouch et al. 2011).

Motivated by the shortcomings of Iro et al. (2003), Mousis et al. (2012b) introduced another statistical model based on an appropriate set of potential parameters and that seems to address these issues. In this model, volatiles are also accreted as clathrate-hydrates, except N$_2$ which, due to its poor trapping propensity, is assumed to condense at $\sim$20 K in the PSN and subsequently trapped in pure crystalline form in comets. Later radiogenic heating by the decay of short-lived nuclides will heat up these bodies to temperatures enough to liberate the trapped N$_2$, but not CO. In Triton and Pluto, the gravity is sufficiently important to prevent the gaseous N$_2$ from escaping, and let it condense on the surface and form the observed cover. In the case of comets however, the gravity is negligible and the gaseous N$_2$ will escape, leaving comets depleted in nitrogen. A shortcoming of this model is that no radiogenic particles were found by stardust (Flynn 2008).

A recent model, applied to the description of Uranus and Neptune formations, also provides an alternative solution. Ali-Dib et al. (2014) proposed that the diffusive redistribution of vapor will remove the gaseous N$_2$ from the region below its iceline, possibly explaining the probable nitrogen depletion in Uranus and & Neptune if they formed in this region. One can invoke a similar argument for the nitrogen depletion in comets. In this case, the CO–rich comets will need to form between the CO and N$_2$ icelines, in order to accrete CO in solid form (since its vapor will also be depleted below its iceline) but not N$_2$. The relatively small distance between the two icelines ($\sim$4–5 AU) is however a caveat for this model since it imposes the formation of all comets in this restricted area of the outer PSN.

Finally, all these models took the N$_2$ depletion at face value. As discussed above, the N$_2$ molecule’s mass is very close to CO, and the two molecules can be distinguished only using an instrument with a very high spectral resolution, such as ROSINA on board the Rosetta spacecraft (Gulkis and Alexander 2008). If this (or another) instrument is able to detect important amounts of N$_2$ in a comet, this might resolve at least part of the problem. Since important chemical differences are known to exist between Oort Cloud comets and Jupiter Family comets (Mumma and Charnley 2011), positive detections or definitive exclusion of N$_2$ in both types of populations are needed before drawing final conclusions.
5 Spin temperature as a cosmogonic indicator in comets

The volatile composition, D/H ratio in molecular volatiles and the spin temperature, as given by the ortho-to-para ratio in water or other molecules, are considered to be possible cosmogonic indicators for comets and as such may be interpreted as providing information about the formation history of the molecules in question. In this section, we discuss spin temperature as a possible cosmogonic indicator for comets.

Molecules that contain identical hydrogen nuclei (like H$_2$O and CH$_4$) display nuclear spin isomers. Water, for example, has two spin species referred to as ortho-H$_2$O if the nuclear spins of the hydrogen atoms are parallel and para-H$_2$O if they are anti-parallel. The lowest energy level of para-H$_2$O lies 23.8 cm$^{-1}$ ($\sim$34 K) below the lowest ortho level, so when water is formed at temperatures below about 50 K, the formation of para states is preferred and the ortho-para ratio (OPR) is less than the statistical equilibrium value of 3. Likewise for methane, statistical equilibrium corresponds to abundance ratios A:E:F of 5:2:9 and statistical equilibrium is reached for temperatures above about 60 K.

OPRs have been routinely measured in water in comets since the apparition of comet 1P/Halley (Mumma et al. 1987). This is currently done by fitting synthetic fluorescent emission models independently for each spin state as shown in Figures 17 and 18. Corresponding spin temperatures are determined by placing the OPR on a theoretical curve such as that for water shown in Figure 19. OPR is also commonly inferred for ammonia by measuring the OPR for NH$_2$, thought to be formed primarily via photodissociation of ammonia (Kawakita et al. 2004; Shin-naka et al. 2011). More recently, spin temperature has been measured in cometary methane for comets C/2007 N3 Lulin (Gibb et al. 2012), C/2007 W1 (Villanueva et al. 2011), C/2004 Q2 (Bonev et al. 2009; Kawakita and Kobayashi 2009), and C/2001 Q4 (Kawakita et al. 2005).

The meaning of spin temperatures has been the subject of much debate. For an isolated molecule of water or methane in the gas phase, for example, nuclear spin conversion is strongly forbidden. For this reason, spin temperature has been considered a cosmogonic indicator in comets that can provide information on the formation temperature (Crovisier 1984; Mumma et al. 1987). Observational evidence supports the suggestion that OPRs do not change once volatiles are released into the gas phase. For example, recent results mapping the spin and rotational temperatures in cometary comae are consistent with a scenario in which the OPR is unchanged by coma processes (Bonev et al. 2008). Also, the OPR for NH$_2$ does not correlate with the heliocentric distance (and hence rotational temperature) of the comet, nor does it correlate significantly with abundances of other volatiles (Shinnaka et al. 2011).

However, it has also been suggested that the spin temperature of water depends on the evaporation history of the ice (Buntkowski et al. 2008; Hama et al. 2011) found experimentally that the OPR of water desorbed from ice after being vapor deposited at 8 K or formed via irradiation of a CH$_4$/O$_2$ mixture at 8 K was consistent with statistical equilibrium, suggesting that OPR may equilibrate during the thermal desorption process or that energetically produced water does not undergo nuclear spin conversion (to a lower OPR) in water ice. Both formation processes for water in the Hama et al. (2011) experiment were energetic, and the authors note that the spin temperature of water molecules produced by non-energetic processes
Fig. 17 Spectrum of comet C/2007 N3 (Lulin) on 31 Jan 2009 with the sum of the telluric and best-fit synthetic fluorescent emission model (red) overplotted. The best fit water ortho (orange) and para (purple) models are shown. Also shown is OH prompt emission (olive). To illustrate the quality of the fit, the residual (scaled by a factor of two) with the best-fit models subtracted is shown. The 1-sigma error envelope is overplotted in green. The best fit is for statistical equilibrium (T_{spin} > 60 K).

Fig. 18 Residual spectrum of comet C/2007 N3 (Lulin) on 31 Jan 2009 (C) with the sum of the best-fit synthetic fluorescent emission model (red) overplotted (A) and offset vertically for clarity. The OH prompt emission is shown in the inset (B). To illustrate the quality of the fit, the residual (scaled by a factor of two) with the best-fit models subtracted is shown in (D) with the 1-sigma error envelope overplotted in green. Beneath the residuals, individual fluorescent emission models are shown for the three spin components of CH$_4$ (E) and CH$_3$D (F) expanded vertically for clarity. The best fit is for statistical equilibrium (T_{spin} > 60 K). Adapted from Gibb et al. [2012].
Fig. 19 OPR vs Spin Temperature in water with measurements for individual comets. Individual OPR measurements are from Mumma et al. (1988: 1a, 1b, 2); Crovisier et al. (1997: 3, 1999: 4a, b); Dello Russo et al. (2005: 5, 6, 7a, 7b, 2007: 11, 12, 2008: 16); Kawakita et al. (2006: 8); Bonev et al. (2009: 9, 2008: 10); Woodward et al. (2007: 13); Villanueva et al. (2011: 14); Paganini et al. (2014: 15, 2015: 17, 2012: 18); DiSanti et al. (2013: 19); Radeva et al. (2010: 20); Gibb et al. (2012: 21).

Can the measurements of OPR in comets shine any light on this question? We show the measurements of cometary OPR in water to date in Figure 19. Water was chosen due to the greater number of OPR measurements than for other molecules in comets. Early measurements suggested a clustering near $T_{\text{spin}} \sim 30$ K, leading many authors to conclude a formation temperature of $\sim 30$ K for a large population of comets. Similar conclusions have been drawn from studies based on CH$_4$ [Kawakita et al. 2006] and NH$_2$ [Shinnaka et al. 2011], particularly given the lack of correlation between spin temperature and other properties of the observed comets [Shinnaka et al. 2011]. However, it must be noted that among the 20 comets measured to date, nearly all of them are within 1- or 2-sigma of the statistical equilibrium value, excepting the measurements reported for 1P/Halley, one of the two measurements for C/2001 A2, and C/1995 O1 (Hale-Bopp). Likewise, half of all measurements are at or above the statistical equilibrium value. It is also notable that no measurements exist below a spin temperature of 20 K, and this observation holds for methane and ammonia spin temperatures as well.

Possible interpretations of spin temperatures measured in comets are therefore (1) that it is a cosmogonic indicator that gives the formation temperature of the molecule in question and therefore perhaps the formation conditions of the comet, (2) that it is a measurement that, while unchanged by coma processes, is set by
processes occurring in the solid ice during or prior to sublimation. Current data are not sufficient to draw firm conclusions at this point. Even if further observations result in most comets consistent with statistical equilibrium, we may still be sampling cosmogonic material since it is known that comets contain material processed at high temperatures, such as crystalline silicates, which are interpreted to originate from turbulent radial mixing in the solar nebula (Bockelée-Morvan et al. 2002). Mixing of volatile material that formed at high temperatures in the inner disk to the comet forming region would have resulted in a higher OPR than for uncontaminated material that formed in the cold midplane. Clearly a great deal more work needs to be done to understand the significance of molecular spin states in cometary material.

6 On the origin of neutral sodium in comet tails

The presence of sodium D line emission has been confirmed in a large number of comets close to perihelion since it was first reported more than a century ago (Bredichin 1882). Observations of comet C/1995O1 Hale-Bopp during the spring of 1997 led to the discovery of a new tail connected with the sodium D line emission. This neutral sodium gas tail is entirely different from the previously known ion and dust tails, and its associated source is unclear. Up to now, only physical reasons have been advanced to rationalize its origin. It has been proposed that this third type of tail is shaped by radiation pressure due to resonance fluorescence of sodium atoms (Cremonese et al. 1997). Further possibilities were considered and then rejected, as photo-sputtering and/or ion sputtering of nonvolatile dust grains (Ip and Jorda 1998). In the same way, collisions between the cometary dust and very small grains were also questioned (Ip and Jorda 1998). The scenario presented in this review is completely different since entirely based upon chemical grounds. It is shown that the Na$^+$ ions, washed out of the refractory material during the hydration phase, loose their positive charge to evolve progressively into neutral species with their migration towards the surface of cometary ices.

Ellinger et al. (2015) follow the chemical path of sodium, starting from the incorporation into the ice until the final transformation into a neutral atom when released from the sublimating crystalline (see Section 2.5) cometary ice. The modeling of bulk, surface structures and reactivity of ices has been essentially developed for environmental applications (Casassa and Pisani 2002; Casassa et al. 2005; Catalayud et al. 2003) Levering on these results, high-level numerical simulations based on first principle periodic density functional theory (DFT) were used to describe the solid structure of the ice in the form of apolar hexagonal ice (Bussolin et al. 1998) composed of bi-layers of water molecules.

A code, specifically designed for the study of periodic systems, namely, the Vienna ab initio simulation package (VASP) (Kresse and Hafner 1993; Kresse and Hafner 1993b) was used to carry out all the calculations. The generalized gradient approximation (GGA) was employed in the form of the Perdew and Wang exchange-correlation functional (Perdew et al. 1992) coupled to the Grimme correction (Grimme et al. 2010) to take care of the long-range van der Waals interactions. The evolution of the ionic character of the sodium atom as it reaches the ice surface was obtained by a topological analysis (Silvi and Savin 1994) performed by means of the TOPMOD
package (Llusar et al. 1999) and its recent extension to periodic systems (Kozlowski and Pilme 2011).

A first calculation on isolated Na(H$_2$O) showed that the interaction of Na with one H$_2$O is similar to that between two H$_2$O molecules in the H$_2$O dimer ($\approx 0.3$ eV), which gives a hint in view of a possible replacement of a molecule of water by a sodium atom in organized structures. In liquid water, microsolvation of sodium atoms has been studied on small clusters of Na(H$_2$O)$_n$ ($n=1$-6) to mimic the water environment. Both the structures in which Na is positioned on the surface and the encapsulated structure in which Na is surrounded by a solvation shell are minima on the potential surface. They are very close in energy (Hashimoto and Morokuma 1994).

![Fig. 20 Typical structures of Na-H$_2$O$_{ice}$ arrangements. Top: Na in the cavity inside hexagonal ice. Middle: Na replacing H$_2$O in the crystal. Bottom: Na adsorbed on the surface of the ice. Color code Yellow (Na); red (O); white (H).](image)

In the ice, the periodic structure of the crystal imposes strong geometrical constraints. The most stable structure, not found in the liquid, is the one where Na takes the place of a water molecule in the lattice (middle structure in Figure 20) when the ice is formed ($\Delta E \approx 1$ eV). Contrary to microsolvation, the stability of Na encapsulated in the bulk of the crystal ($\Delta E \approx 0.5$ eV), shown in the top structure in Figure 20 differs from that of the adsorbed Na according to the position on the surface. For example, the adsorption energies vary from ($\Delta E \approx 0.6$ to $\approx 0.5$ and $\approx 0.1$ eV) when the adsorption takes place over the center of a surface hexagon, over a surface oxygen (bottom structure in Figure 20) and over an OH dangling bond, respectively.

The most relevant point to this study is that the charge $q_{Na}$ on the sodium atom depends strongly on the position with respect to the ice surface, as does the adsorption energy. When constrained to stay in the cavity inside the hexagonal
lattice, Na remains largely ionized ($q_{\text{Na}} \approx +0.9$) which is close to the value of $\approx +0.8$ when Na takes the place of an H$_2$O molecule in the bulk. When Na is adsorbed on top of an oxygen of the surface layer, $q_{\text{Na}} \approx +0.2$, a value similar to that found for the isolated Na(H$_2$O) complex. Then, as the ice is progressively eroded by the radiation field, Na eventually reaches the surface where it can desorb as a neutral atom when the upper layer of the ice vaporizes.

7 Summary

The observed compositions of comets are the end product of a chemical history that began in the cold parent molecular cloud core. The change in physical conditions experienced by the core material as the core collapsed and the protosun and protosolar disk formed and evolved led to chemical changes that are reflected in the cometary volatiles that we observe today. Each stage of evolution left its imprint in the mixing ratios of molecules observed in comets and disentangling their effects requires interaction between comet and disk researchers. The combination of the two fields of study provide a powerful tool in understanding the drivers that determine comet composition and conversely what the variations in comet composition can tell us about the relative formation conditions of these comets.

Protostellar disks provide complementary information to the comet observations. Their chemistry is complex, covering ice chemistry in the cold, shielded midplane, warm gas phase chemistry and hot, highly ionized chemistry. Comets form from the ices in the midplane, yet dynamical processes ensure that this region does not evolve in isolation but rather is the end product of material that has been processed in a variety of different environments and then brought together in the cometary ices. Establishing the origin of cometary ices therefore requires not only studying the midplane but also an understanding the chemical and physical evolution of the disk as a whole. By studying disks we can observe the chemistry and physics that shaped the solar system as they shape other protoplanetary systems.

Molecular deuteration is a link that might tie together the disparate objects of the solar system and their formation history and give us the picture of the whole solar system formation history. However, at present this link has still weak connections that need to be reinforced. The advent of the powerful new facilities, as for example ALMA and IRAM-NOEMA, will certainly allow this field to progress rapidly in the next few years.

Rosetta will provide detailed ground-truths for the organic and isotopic makeup of cometary volatiles. The new insights obtained into the chemical processes which likely produced these molecules – interstellar or nebular – will allow us to constrain models of the formation and evolution of the early solar system. Its instruments will characterize the composition of the comet 67P/Churyumov-Gerasimenko and distinguish between the molecules native to the comet nucleus (parent species) and those generated by chemistry or by fragmentation in the comae (daughter species). The observations it will make will provide new insights into the formation conditions of the comet. They will also elucidate the degree to which interstellar material can survive its journey into the solar nebula and from there into comets.

Several measurements will characterize the its thermal history, e.g., measurements of molecular spin temperatures (Gulkis and Alexander 2008) and deuteration and of the abundance of argon (Hassig et al. 2014). The relative abundances
of elements can tell us something about the dominant molecular form of each element at the time of formation – if an element is depleted relative to others then the major carrier of that element must have been too volatile to be condensed at the time of formation, or must have desorbed in the intervening time. In particular Rosetta measurements may help to elucidate the mystery of the lack of nitrogen in comets (Altwegg et al. 2015).

A key measurement will be the ratio of oxygen isotopes in water. Meteoritic water has a very different ratio of the oxygen isotopes compared to oxygen isotopes in the Sun. Rosetta provides our best opportunity to directly detect the isotope ratio from the early solar system, and in so doing will distinguish between the various theoretical models of how the oxygen isotope fractionation occurred.

In summary, Rosetta’s observations will provide new insights into the chemical processes that resulted in the formation of cometary volatiles and consequently into the conditions that were present in the solar nebula at their time of formation.

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