Assessment of the Interstellar Processes Leading to Deuterium Enrichment in Meteoritic Organics

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Abstract - The presence of isotopic anomalies is the most unequivocal demonstration that meteoritic material contains circumstellar or interstellar components. In the case of organic compounds in meteorites and interplanetary dust particles (IDPs), the most useful isotopic tracer has been deuterium (D). We discuss four processes that are expected to lead to D enrichment in interstellar materials and describe how their unique characteristics can be used to assess their relative importance for the organics in meteorites. These enrichment processes are low temperature gas phase ion-molecule reactions, low temperature gas-grain reactions, gas phase unimolecular photodissociation, and ultraviolet photolysis in D-enriched ice mantles. Each of these processes is expected to be associated with distinct regiochemical signatures (D placement on the product molecules, correlation with specific chemical functionalities, etc.), especially in the molecular population of polycyclic aromatic hydrocarbons (PAHs). We describe these differences and discuss how they may be used to delineate the various interstellar processes that may have contributed to meteoritic D enrichments. We also briefly discuss how these processes may affect the isotopic distributions in C, O, and N in the same compounds.

INTRODUCTION

Isotopic measurements provide definitive evidence for presolar materials within meteoritic samples, including those from both circumstellar and interstellar environments. The presence of circumstellar materials in meteorites is demonstrated by isotopic anomalies measured in a variety of different types of presolar materials (e.g. nanodiamonds, Al₂O₃, graphite, SiC, etc.) (cf. Zinner 1988, 1997). The carriers of these isotopic anomalies are generally believed to have formed in circumstellar ejecta and the anomalies are thought to be of nucleosynthetic origin (Bernatowicz and Zinner, 1997). Indeed, the measurement of isotopic ratios and their correlations in multiple elements is leading to unprecedented advances in our understanding the details of nucleosynthesis in a number of different stellar environments (cf. Gallino et al., 1997).
The existence of interstellar materials in meteoritic samples is also indicated by isotopic evidence, although its nature is significantly different. In this case, the isotopic anomalies are primarily in the form of deuterium (D) enrichments (Robert and Epstein, 1982; Yang and Epstein, 1983; Kerridge and Chang, 1985; Kerridge et al., 1987), although smaller effects have also been seen in nitrogen (Robert and Epstein, 1982; Franchi et al., 1986; Sugiura and Zashu, 1995). In some cases these enrichments are seen in bulk meteoritic materials (Zinner, 1988), but D enrichments have also been observed in meteoritic subfractions and even within specific classes of molecular species, such as amino and carboxylic acids (Epstein et al., 1987; Pizzarello et al., 1991; Krishnamurthy et al., 1992). These anomalies are not thought to be the result of nucleosynthetic processes, but are instead ascribed to chemical processes occurring in the interstellar medium (ISM).

The traditional explanation of these D excesses has been to invoke the presence of materials made in the ISM by low temperature gas phase ion-molecule reactions (Geiss and Reeves, 1981; Dalgarno and Lepp, 1984). However, ion-molecule reactions represent only one of several processes that can produce strong D-H fractionation in the ISM. We will discuss four chemical mechanisms that are expected to generate D-H fractionation in the ISM: (i) low temperature gas phase ion-molecule reactions, (ii) low temperature gas-grain reactions, (iii) gas phase unimolecular photodissociation, and (iv) ultraviolet photolysis in D-enriched ice mantles. Each process is discussed in two parts: first, we describe the basic chemical process and how it leads to 'generic' molecular D fractionation, and second, we describe how the fractionation process manifests itself in the specific case of polycyclic aromatic hydrocarbons (PAHs).

We chose to highlight PAHs for several reasons. First, PAHs are probably the most abundant and widespread class of organic compounds in the universe (Puget and Leger, 1989; Allamandola et al., 1989, 1999). They are observed in the gas phase in a wide variety of interstellar environments, including protoplanetary and planetary nebulae, reflection nebulae, HII regions, and the diffuse ISM, in the ices in dense molecular clouds, and in carbonaceous grains in the diffuse ISM (Allamandola et al., 1989, 1999; Sandford et al., 1991; Sellgren et al., 1995; Brooke et al., 1996, 1999; Onaka et al., 1996; Roelfsema et al., 1996; Chiar et al., 2000; Bregman
et al., 2000). Second, PAHs are extremely stable molecules and represent one of the few classes of materials that can exist and survive in all the interstellar environments associated with the four chemical processes listed above. Third, PAHs and related aromatic materials are common in carbonaceous chondrites (Cronin et al., 1988; Hahn et al., 1988; Cronin and Chang, 1993; Gardinier et al., 2000; and references therein) and IDPs (Allamandola et al., 1987; Clemett et al., 1993), and have been reported in the comae of comet P/Halley (Moreels et al., 1994). In addition, the aromatic fractions of meteorites are known to be significant carriers of D (Kerridge and Chang, 1985). Thus, PAHs represent a known link between the ISM and solar system samples and could potentially serve as probes of the chemical processes associated with interstellar D fractionation.

INTERSTELLAR CHEMICAL PROCESSES AND DEUTERIUM ENRICHMENT

We will consider the processes by which interstellar molecules, including PAHs, can become D-enriched, including gas phase ion-molecule reactions at low temperatures (Dalgarno and Lepp 1984; Tielens, 1997), gas-grain reactions (Tielens, 1983, 1992, 1997), unimolecular photodissociation (Allamandola et al., 1987, 1989), and UV photolysis in D-enriched ice grains (Sandford et al., 2000). Each of these processes is expected to leave a distinct signature in the distribution of D enrichment in the PAH population.

Low Temperature Gas Phase Ion-Molecule Reactions

The Chemical Process

Deuterium fractionation by ion-molecule reactions is perhaps the longest and best known process for the enrichment of interstellar species, and it is not surprising that it is often invoked to account for the D enrichments seen in meteoritic materials (cf. Geiss and Reeves, 1981). It is generally assumed that the fractionation occurred in the ISM rather than in the solar nebula, because this effect falls off rapidly with increasing temperature. The temperatures and time scales associated with inner protosolar disks are not expected to allow significant fractionation to occur, although fractionation could potentially occur in their outer regions (Aikawa and Herbst, 1999).
Deuterium fractionations via gas phase ion-molecule reactions require very low temperatures. Chemical reactions between neutral gas phase species that are endothermic or possess activation barriers will not proceed at any appreciable rate at the low temperatures characteristic of dense clouds ($T < 50$ K). However, exothermic ion-molecule reactions can proceed even at very low temperatures (cf. Herbst, 1987; Tielens, 1997). Thus, most of the gas phase chemistry occurring in dense clouds is the result of ion-molecule reactions and exoergic reactions involving radicals.

Such reactions are expected to produce mass fractionation at low temperatures due to the different zero point energies of H and D, deuterium having a lower energy and hence the potential to form slightly stronger chemical bonds temperatures (Qalgaro and Lepp, 1984; Tielens, 1997; Millar et al., 2000). This results in the two having exchange rates that differ by a factor of $e^{-(\Delta E/T)}$, where $\Delta E$ is the energy difference and $T$ is the temperature. At higher temperatures the difference in energies is proportionally less important and fractionation is a minor effect, but as the gas temperature drops below 70 K, fractionation becomes increasingly more significant (e.g. Millar, Bennett, and Herbst, 1989).

There is good observational evidence that such fractionation occurs in the ISM. Radio telescopic observations of the rotational lines of several simple molecules and their deuterated counterparts in dense clouds demonstrate that these species are often highly D-enriched; D enrichments exceeding a factor of 5000 above the cosmic value of $\text{HD/H}_2 = 3 \times 10^{-5}$ have been reported (Jefferts et al., 1973; Hollis et al., 1976; Mauersberger et al., 1988). However, this process only operates on molecules in the gas phase, and the majority of species are condensed onto grains at the low temperatures where fractionation is significant. Thus, while ion-molecule reactions clearly contribute to D fractionation in the ISM, they may not be the main contributor to the bulk fractionation of D in dense clouds.

**PAH Regiochemistry**

Like smaller molecules, gas phase PAHs should become enriched in D by ion-molecule reactions in dense molecular clouds. The principle reaction path involves a two step process
First, a PAH reacts with an \( \text{H}_2\text{D}^+ \) ion to pick up an excess deuterium atom and charge.

\[
\text{PAH} + \text{H}_2\text{D}^+ \rightarrow \text{PAHD}^+ + \text{H}_2
\]

Subsequent reaction with an electron neutralizes the charge and the PAH eliminates an excess H or D atom. The fraction in which an H is lost, as opposed to a D, can be described by a branching ratio, \( \alpha \).

\[
\text{PAH-D}^+ + e^- \rightarrow \alpha_h(\text{PAH-D}) + \text{H} \quad \text{and} \quad (1-\alpha_h)\text{PAH} + \text{D}
\]

The branching ratio, \( \alpha_h \), is largely determined by statistics (Millar et al., 1989; S. Charnley, private communication) modified by the difference in the the H and D zero point energies. A lower limit to the D-enrichment can be derived easily by ignoring the effect of the zero point energy difference and simply assuming that every H or D on the molecule has about the same chance of being ejected [i.e., \( \alpha_h \sim N_h/(N_h+N_D) \)]. Thus, this process would be expected to initially preferentially deuterate the larger PAHs in dense clouds since they have more H atoms and will lose a higher fraction of H relative to D. Ultimately, all PAHs in the gas phase would be expected to show increasing D/H ratios as they approach steady-state with the gas phase. In this manner, over the lifetime of a dense cloud core (~3\times10^6 years, \( n_H = 3\times10^4 \text{ cm}^3 \)), individual PAHs would be expected to acquire a few peripheral D atoms. Since PAHs are considerably more stable and more abundant in the ISM than many of the smaller species that show D enrichments (PAH/\( \text{H}_2 \sim 10^{-6} \) compared to HCN/\( \text{H}_2 \sim 10^{-8} \)), PAHs are likely to represent the largest reservoir of materials D-enriched by ion-molecule reactions.

**Low Temperature Gas-Grain Reactions**

*The Chemical Process*

At the temperatures characteristic of dense clouds (T < 50 K) most of the volatile species in dense clouds are condensed out of the gas phase into icy grain mantles (Sandford and Allamandola 1993). Evidence for this condensation is found in the large quantities of mixed-molecular ices...
(Sandford, 1995) and gas phase depletions (Mauersberger et al., 1992) seen in dense clouds. The composition of the ice mantles does not simply reflect gas phase abundances, however. Instead, additional species are formed when condensing species participate in gas-grain reactions on the grain surfaces, reactions that cannot occur solely in the gas phase (Tielens and Hagen, 1982; d'Hendecourt et al., 1985; Brown and Charnley, 1990; Hasegawa et al., 1992). The compositions of these new species are strongly dependent on the local H/H₂ ratio. In environments where the H/H₂ ratio is large, surface reactions with H atoms dominate and atoms like C, N, and O are converted to simple hydride species like CH₄, NH₃, and H₂O. In contrast, if H/H₂ is substantially less than one, reactive species such as O and N could, in principle, react with one another to form O₂ and N₂. Thus, two qualitatively different types of ice mantle can produced by grain surface reactions, one dominated by polar, H-bonded molecules and the other dominated by less polar, highly unsaturated molecules. Infrared spectral observations of dense clouds confirm this basic dichotomy (Sandford et al., 1988; Tielens et al., 1991; Chiar et al., 1995).

Gas-grain chemistry should produce large D enrichments in the grain mantles as the result of mass fractionation that become important at low temperatures (Tielens, 1992, 1997). In this case, except for H-abstraction reactions, the role of the zero-point energy difference between H and D counterparts in creating D fractionation is limited (Tielens, 1983). Instead, D enrichment results primarily from the high atomic D fractionation of the accreting gas. Both the atomic H and D accreted onto grain surfaces are light enough to tunnel through activation barriers on interstellar time scales and thereby hydrogenate species that are not fully reduced. Models indicate that such a process could result in mantle D/H ratios as high as 0.1 for simple molecules like H₂O and CH₃OH (Tielens, 1983, 1992, 1997; Charnley et al., 1997). Values in this range have been reported for HDO/H₂O in interstellar ices (Teixeira et al., 1999). Such values meet or exceed the enrichments produced by ion-molecule reactions in gas phase species like C₃HD/C₃H₂ and DCO⁺/HCO⁺.

Since the grain mantles in dense clouds represent a much larger fraction of the total reservoir of material than does material in the gas phase (H₂ excepted), grain surface processes are likely to be far more important for the total D fractionation in dense clouds than ion-molecule reactions.
PAH Regiochemistry

D fractionation during gas-grain reactions should only proceed efficiently for molecules that can be hydrogenated. Fully reduced species will not accept additional D in this fashion and simple exchange is unlikely under these low temperature conditions. Whether PAHs can become D-enriched by this process is unclear. There is currently no laboratory evidence suggesting simple H-atom exposure on grain surfaces leads to the excess hydrogenation of PAHs.

It is possible, with energetic processing, for PAHs with fully aromatic bonding to accommodate additional peripheral H atoms (Bernstein et al., 1999). The effect of these excess H atoms in these molecules, which are collectively termed \( \text{H}_n\)-PAHs, is to convert the affected C atoms from aromatic sp\(^2\) bonding to aliphatic sp\(^3\) bonding (see Figure 2). This results in molecules that contain both aromatic and aliphatic rings. In the extreme case when the maximum possible number of H atoms have been added, the result is a polycyclic aliphatic hydrocarbon. There is some evidence for the existence of \( \text{H}_n\)-PAHs in space (Bernstein et al., 1996). In particular, their presence may explain a number of enigmatic weak infrared emission features seen in the outflows of aging stars (protoplanetary and planetary nebulae), but the environmental conditions in these objects are very different from the conditions pertinent for low temperature gas-grain reactions.

Since there is currently no good theoretical rationale or experimental evidence that suggests PAHs can be hydrogenated to form \( \text{H}_n\)-PAHs by simple gas-grain interactions, it is likely that PAHs do not accumulate significant H atom excesses, and thus D enrichments, by gas-grain reactions. Thus, while gas-grain reactions may dominate the mass distribution of D in dense clouds, it may leave PAHs largely unaffected. To the extent that this process occurs to PAHs, any D enrichments generated would be expected to correlate with the presence of \( \text{H}_n\)-PAHs, increasing H/C ratio, or aliphatic structures.

Unimolecular Photodissociation Reactions

In contrast to ion-molecule and gas-grain reaction processes, D-enrichment by unimolecular photodissociation does not require low environmental temperatures and is expected to be
essentially restricted solely to PAHs (Allamandola et al., 1987, 1989; Tielens, 1997). Because of their multiple C bonding and π electrons, PAHs are much more stable against photolytic disruption than most other molecules. This allows PAHs to survive in environments where other species would quickly be reduced to fragments. This is one of the chief reasons why PAHs are one of the most ubiquitously and abundantly distributed classes of molecules in the ISM (cf. Allamandola et al., 1989; Roelfsema et al., 1996).

While PAHs are relatively immune to complete photodestruction by interstellar UV photons, individual PAHs may still absorb UV photons that contain more energy than they can accommodate vibrationally. In this case, a bond or bonds will break. The most likely bond to break is that of a peripheral C-H, and because of the zero point energy difference of the C-D and C-H bonds, the rate of D loss from interstellar PAHs should be lower than that of H (Figure 3). Since both H and D will return to this molecular site from the gas phase, repeated processing should gradually enrich interstellar PAHs in D relative to the ambient gas. In steady state, which is expected to be reached in photodissociation regions and, at least for small PAHs, in dense clouds, the fraction of peripheral D relative to H will be 3 times the local gas phase ratio of D to H which can range from $10^{-5}$ to $10^{-3}$ (Allamandola et al., 1989; Tielens, 1997). Thus, in these interstellar environments, small PAHs (≤ 25 C atoms) are expected to have one or two peripheral D atoms.

This process differs from enrichment by ion-molecule and gas-grain reactions in a number of ways. First, low ambient temperatures are not required to produce fractionations by this process. Second, the extent of the D-enrichment depends critically on the size of the PAH. The D photoenrichment of PAHs is expected to be most significant for PAHs in the range of sizes spanned by naphthalene ($C_{10}H_8$) and hexabenzocoronene ($C_{42}H_{32}$). Deuterium enrichment is not expected in benzene because this molecule is not stable in the interstellar radiation field. Enrichment in PAHs having more than ~40 C atoms is not expected because these larger PAHs have large numbers of vibrational modes, can accommodate the maximum energy of typical UV photons, and are stable against photolytic bond rupture (Figure 3).
Ultraviolet Photolysis Reactions in D-enriched Ice Mantles

The Chemical Process

Most of the volatile species in dense clouds reside in icy grain mantles. The dominant species in these ices is usually H$_2$O, but they also contain CO, CO$_2$, CH$_3$OH, NH$_3$, H$_2$CO, CH$_4$, and a number of other simple species in abundances greater than 1% that of the H$_2$O (see Sandford, 1996). Many of these species should be significantly enriched in D, either by enrichment in the gas phase by ion-molecule reactions followed by condensation onto the grain mantle, or via gas-grain reactions occurring on the mantle surface. Once in the ice, these molecules can serve as a D-enriched reservoir from which other enriched species can be made.

One would normally expect little in the way of chemical reactions to occur within these ice mantles because of their very low temperatures. However, interstellar ices in dense clouds can be further modified by energetic processing by ambient UV radiation and cosmic rays (Norman and Silk, 1980; Prasad and Tarafdar, 1983). When these ices are irradiated, bonds are broken, molecules are destroyed, and new molecules, radicals, and ions are formed. Most of these new, reactive species remain trapped in the ice matrix. Ultimately their accumulation is interrupted by a thermal event (grain-grain collision, passage of a cosmic ray, exposure to a stellar system, etc.) that causes the ice to warm to the point at which they can become mobile, react, and form new and more complex molecules. Thus, a host of species formed by the irradiation and thermal processing of ices made of simpler (D-enriched) molecules is expected to be present in dense interstellar clouds (Moore et al., 1983; Agarwal et al., 1985; Allamandola et al., 1988; Bernstein et al., 1995; Dworkin et al., 2001) and infrared spectra of the ices in dense clouds provide evidence for this process (Tegler et al., 1993; Pendleton et al., 1999; Bernstein et al., 2000).

Laboratory studies of mixed-molecular ices in which some of the original species are D-enriched have shown that the D ends up in the reaction products (Bernstein et al., 1994, 1995, 2000; Sandford et al., 2000). Unlike the processes described above, this process does not produce a fractionation of H from D, but instead propagates previous fractionations into new molecular
species. Thus, the irradiation of D-enriched interstellar ices produced by other processes will lead to the production of a host of more complex organic species that are also D-enriched.

**PAH Regiochemistry**

At the low temperatures of dense molecular clouds, PAHs will condense onto refractory dust grains along with most other gas-phase species. The presence of PAHs in these environments has been demonstrated by detection of their infrared vibrational bands in absorption in the spectra of stars embedded within dense clouds (Sellgren *et al.*, 1995; Brooke *et al.*, 1996, 1999; Chiar *et al.*, 2000; Bregman *et al.*, 2000). Once condensed into interstellar ice mantles, PAHs will be subjected to the same radiation processing as the other ice components. Laboratory irradiation simulations of PAH-bearing interstellar ice analogs have shown that under these conditions PAHs undergo both oxidation and reduction reactions that affect their edge structures (Bernstein *et al.*, 1999, 2001).

Oxidation results in the production of aromatic alcohols, ketones, and, in PAHs with 'bay' regions in their edge structures, ethers, while reduction leads to the formation of $H_n$-PAHs, i.e., mixed aromatic and aliphatic rings (Figure 4). Many of these aromatic species are very similar to some of those seen in carbonaceous chondrites and IDPs (see Bernstein *et al.*, 1999, 2001).

Laboratory studies of PAHs in D-enriched $H_2$O ices have shown that UV photolysis results in rapid D enrichment of the PAHs through aromatic D$\rightarrow$H exchange, D-atom addition, and exchange through keto-enol tautomerism. Each of these processes produce different compounds with characteristic deuteration patterns and D labilities (Sandford *et al.*, 2000) (Figure 5). For example, D$\rightarrow$H exchange reactions result in D-enriched PAHs in which the D is sited in relatively non-labile locations (Figure 5a). D-atom addition reactions result in D-enriched $H_n$-PAHs in which the D is sited in relatively non-labile aliphatic rings attached to aromatic rings (Figure 5b). In contrast, enrichment via keto-enol tautomeric exchange results in the D being located exclusively on rings that contain oxygen atoms in sites that are relatively labile (Figure 5c). This latter enrichment would likely not be preserved during aqueous alteration.
Deuterium enrichments produced in aromatic species in this manner are expected to occur largely independent of PAH molecular size, but should show specific regiochemical behaviors. In addition, since this same irradiation process produces both reduced PAHs (H₂-PAHs) and oxidized PAHs (aromatic alcohols, ketones, and ethers) (Bernstein et al., 1999), D enrichments produced by this process would be expected to correlate with the presence of these other species.

**Possible Correlated Effects in Carbon, Nitrogen, and Oxygen**

Some of the interstellar processes described above can also drive chemistry involving C, N, and O, so in principle they could also result in enrichments in these elements. However, since the zero-point energy differences for the isotopes of C, N, and O are significantly smaller than for H (10-50 K vs. a few 100 K), fractionations in these elements should be considerably less pronounced (see Tielens, 1997). Nonetheless, D enrichments could potentially correlate with isotopic fractionations in these other elements.

Gas phase ion-molecule reaction models predict \(^{12}\text{C}/^{13}\text{C}\) fractionations can be produced in organic species from gas phase \(\text{C}^+\) generated by isotopic switching reactions with CO. Such fractionations could favor \(^{12}\text{C}\) in the more complex species by as much as a factor of 2, but generally much smaller fractionations are expected (Langer et al., 1984; Langer and Graedel, 1989). Radio observations provide some general support for this scheme (Henkel et al., 1982; Langer and Penzias, 1993). Isotopes of carbon can also be exchanged in proton transfer reactions involving species like \(\text{HCO}^+\), but these effects are expected to be quite small (Langer et al., 1984). Finally, C can be fractionated by isotope-selective photodissociation of CO. In dense clouds regions where \(A_v < 6\), UV photodissociation of CO is mediated by excited \(\text{H}_2\) and is expected to lead to enhanced \(^{13}\text{C}/^{12}\text{C}\) in the gas phase, i.e., the opposite fractionation expected from ion-molecule reactions (Tielens, 1997). As a result, the overall C isotope fractionation in the gas phase depends on the relative importance of atomic C and \(\text{C}^+\) in the generation of larger molecules, an issue that has yet to be fully understood.

Differences in zero-point energies are relatively unimportant for gas-grain reactions, so carbon-bearing species derived from accreting gas phase CO should not be fractionated. Only
species that have a direct heritage to gas phase C, for example CH$_4$, should be fractionated (Tielens, 1997). Thus, as with D, the primary enrichment from this process results from prior fractionation of the accreting gas, which we noted earlier could potentially be fractionated in either a positive or a negative sense.

Oxygen in the gas phase resides primarily in CO, O, and/or O$_2$. Theoretical calculations suggest that very little fractionation in the $^{16}$O/$^{18}$O ratio should occur in these species or their products (Langer et al., 1984). Lacking a significant fractionation in the gas phase, gas-grain reactions are not expected to generate fractionated species either and, consequently, ice irradiation processes should also not produce new species with fractionated oxygen. Thus, one would not expect D-enrichments to correlate with any significant O isotopic fractionation.

The molecular form of most of the nitrogen in dense clouds has yet to be determined. Some of it resides in NH$_3$ (Lacy et al., 1998), but the majority is probably in the form of N$_2$ (Sandford et al., 2001). Some N fractionation is likely driven by gas phase ion-molecule exchange reactions between N$_2$H$^+$ and N$_2$. Most of the $^{15}$N-enhanced N$_2$H$^+$ is simply cycled back to N$_2$, but some can be propagated through the exchange of both a N and H to species like NH, NH$_2$, HCN, etc. The ultimate fractionations expected for such simple hydrides lie between $\delta^{15}$N values of 10 and 1000‰, depending on the molecule and evolutionary time scale (Terzieva and Herbst, 2000). For the most abundant molecules, such as NH$_3$, $^{15}$N enrichments as high as $\delta^{15}$N = 250‰ might be expected. Such values are similar to the enrichments seen in many meteoritic materials, but they cannot explain some of the more extreme values seen in meteorites and IDPs (Messenger, 2000). However, very recent calculations of $^{15}$N gas-grain fractionation suggest that much higher $^{15}$N/$^{14}$N ratios might be attained in N$_2$ and NH$_3$ at high gas phase densities (Charnley and Rodgers, 2001). Thus, moderate fractionations of N can occur in the gas phase and interstellar ices should be $^{15}$N-enriched by both the condensation of enriched gas phase species and via gas-grain reactions. Subsequent ice irradiation will propagate this fractionation into new and more complex species.
Currently, no theoretical or experimental work has been done addressing how gas phase and
gas-grain processes involving PAHs might fractionate C, N, or O. However, limited laboratory
work has been done on how PAHs are modified when they are UV irradiated in ice mantles
containing C, N, and O. Photolysis of PAHs in H₂O ices adds −OH and =O groups and excess
hydrogen atoms to PAH edges (Bernstein et al., 1999, 2001). Preliminary experiments involving
other interstellar ice species (CO, CO₂, CH₃OH, and NH₃) indicate that the formation of aromatic
amino and alkyl moieties is also possible (Elsila et al., 2000). As with D, such photochemistry
could result in the addition of peripheral groups containing N, O, and possibly C, that propogate
any isotopic fractionations found in the simpler molecules present in the ice. Again, these
exchanges should produce fractionations having specific regiochemical systematics characteristic of
the ice photolysis process.

Since none of the processes described above involve changes to the carbon in the PAH
skeletal structure, their C isotopic ratios should be primarily defined by the C reservoir from which
they were first made. Since interstellar PAHs are probably produced in the outflows of aging stars
(Allamandola et al., 1989), their C isotopic ratios should primarily be of nucleosynthetic origin.
Any C isotopic fractionation produced by interstellar chemistry at the edges of PAHs would be
superimposed on this original nucleosynthetic ratio.

In summary, all the processes described above may produce isotopic fractionations in C and
N, but these should be considerably smaller than those seen for D. Fractionation of O is expected
to be smaller still. In the case of carbon, the existence of multiple reaction schemes that can cause
fractionation in opposite directions preclude expectation that C isotopes will correlate in any simple
way with D fractionations. Correlations will be further muddled by the fact that the C, N, and O
fractionations are much more sensitive to temperature than D. Nonetheless, it is possible that
significant ¹⁵N excesses could be produced in the ISM (up to the order of δ¹⁵N ~ 1000‰) and that
these might correlate, at least qualitatively, with D excesses. How these fractionations might
manifest themselves in aromatic species is currently poorly understood.
Comparison of the Signatures of the Various Interstellar D-enrichment Processes

The four different processes described above can all result in the D enrichment of polycyclic aromatic hydrocarbons and related species in the interstellar medium. However, each of these processes produce enrichments that display distinctly different 'signatures' within their associated product populations. These differences are summarized in Table 1.

IMPLICATIONS FOR THE AROMATIC FRACTION OF METEORITIC ORGANICS

Aromatic Materials in Meteorites and IDPs

The organic fraction of primitive meteorites contains a major aromatic component (Studier et al., 1972; Hayatsu et al., 1977; De Vries et al., 1993; Gardinier et al., 2000), and a host of specific aromatic molecules have been identified (Pering and Ponnamperuma, 1971; Basile et al., 1984; Tingle et al., 1991; Kovalenko et al., 1992; Zenobi et al., 1992; Naraoka et al., 2000). Many of the PAHs reported in meteorites have peripheral methyl groups (Basile et al., 1984; Kovalenko et al., 1992; Zenobi et al., 1992), and other functionalities, including ketones, have been reported (Hayatsu et al., 1977; Krishnamurthy et al., 1992). Aromatics have also been detected in IDPs (Allamandola et al., 1987; Clemett et al., 1993).

To date, relatively little attention has been paid to oxidized and reduced aromatics in meteorites. Several groups have used chromatography to detect various 3-ringed ketones in aromatic fractions from the Murchison meteorite (Basile et al., 1984; Krishnamurthy et al., 1992). Cronin and Pizzarello (1990) have shown that the principle aliphatic components of the Murchison meteorite appear to be a structurally diverse set of C_{15}-C_{30} branched and alkyl-substituted mono-, di-, and tricyclic alkanes. In addition, infrared spectra of their extracts suggest the presence of carbonyl groups and some double bonds. Similar structures (partially reduced aromatics with peripheral carbonyl groups) are formed when aromatic hydrocarbons are irradiated in H_2O-rich ices (Bernstein et al., 1996, 1999, 2001). Aromatic alcohols have not been reported in meteorites. Whether the lack of detection of aromatic alcohols is due to their absence or whether it is the result
of selection effects associated with their extraction from meteorites is not yet clear. Targeted searches for aromatics with -OH edge groups are needed to address this issue.

**Fractionations of D, C, N, and O in the Aromatics in Meteorites and IDPs**

The aromatic fraction of meteorites is known to contain carriers of excess D. Early evidence came from the observations that D enrichments as high as $\delta D \geq 1000\%$ can be found in acid-soluble hydrocarbon phases in meteorites and that the enrichment increases as C/H increases from 0.2 to 1.5, i.e., as the material became more aromatic (Yang and Epstein, 1983). In addition, acid insoluble phases with C/H ratios of 2.3 to 3.0 also showed significant D enrichment (Robert and Epstein, 1982). If we assume the fractions with C/H > 1 are dominated by aromatics, these C/H ratios correspond to aromatic moieties containing from 20 to several hundred C atoms.

Using combined carbon and hydrogen isotopic data, Kerridge et al. (1987) demonstrated that the organic fraction of Murchison contains at least three distinct components that carry D-enrichment signatures indicative of an interstellar relationship. Of these, PAHs were a major carrier of the D [Kerridge et al.'s component C(II)H(II), $\delta D \geq +1300\%$]. In addition, Krishnamurthy et al. (1992) examined meteoritic organics and found $\delta D$ values that ranged from about +100 to +1000, with aromatics having higher values than aliphatics. Unfortunately, there is currently little information about the distribution of the D within the aromatic fraction, either in terms of its regiochemistry or its distribution among the various aromatic species present.

The C isotopic character of the D-enriched aromatics is also not fully understood. Studies have demonstrated that the larger deuterium enrichments ($\delta D$ in excess of $+1000\%$) in meteorites do not correlate with C isotopic ratios (e.g., Robert and Epstein, 1982; Yang and Epstein, 1983, 1984; Halbout et al., 1990), which tended to be normal ($-40 \leq \delta^{13}C \leq +20\%$). [The relative importance of aromatics as carriers in the samples measured in these studies is not well established, although the high C/H ratios of some of the samples imply the presence of large amounts of aromatic]
materials (Robert and Epstein, 1982; Yang and Epstein, 1983).] In a study that specifically targeted the isotopic ratios of C and H in meteoritic aromatics, Kerridge et al. (1987) found the aromatics to have $\delta^{13}$C values of about -20‰ that did not correlate with D content. Thus, there does not appear to be any clear correlation between D enrichments and C isotopic systematics in meteorites.

Considerably less work has been done in the area of the O isotopic compositions of these materials. Halbout et al. (1990) studied the H and O isotopic compositions of kerogen extracted from the Orgueil meteorite by acid demineralization. While they did not focus on the aromatic materials in this sample per se, these kerogens are known to be dominated by aromatic structures (cf. Cronin and Chang, 1993). Halbout et al. measured large D-excesses in their sample ($\delta D = +1360‰$) but found relatively normal oxygen ($\delta^{17}$O = +3.3‰) and carbon ($\delta^{13}$C ≤ -48‰). Thus, there is currently no evidence for anomalous O in these types of materials and, hence, no correlation of D enrichments with O isotopic ratios.

Nitrogen is potentially more interesting. Acid residues from carbonaceous chondrites measured by Robert and Epstein (1982) showed $\delta D$ values up to +3500‰ that did not correlate with $\delta^{15}$N. In 1986, Franchi et al. reported $^{15}$N excesses in an acid resistant, presumably carbonaceous, carrier from the Bencubbin meteorite. The carrier yielded $\delta^{15}$N values as high as +1033 ‰ but showed normal C with a $\delta^{13}$C value of about +25‰. Kerridge et al. (1987) identified several isotopically distinct reservoirs of N ranging from light ($\delta^{15}$N < -6‰) to heavy ($\delta^{15}$N > +60‰) which they characterized as being associated with aromatics, but they were unable to closely define the reservoirs. Grady and Pillinger (1990) noted that the primitive chondrite ALH 85085 was highly enriched in $^{15}$N, a portion of which resides in a carrier that is probably carbonaceous material with C/N = 65-75, $\delta^{15}$N = +860‰, and $\delta^{13}$C = 1‰. In a recent comprehensive study, Alexander et al. (1998) examined the C and N isotopic behaviors of the insoluble macromolecular
organic matter in thirteen primitive chondrites. They found that the range of C isotopic compositions varied by only about $\delta^{13}C = 30\%o$ within their samples, while the N isotopes ranged from $\delta^{15}N = -40$ to $+260\%o$. They conclude that the most $^{15}N$-rich material probably formed in the ISM and that the fraction of organic N in meteorites having an interstellar origin is probably very large. For example, in the Renazzo meteorite, Alexander et al. estimate 40-70% of the organic nitrogen may have a direct interstellar heritage. Most recently, Pearson et al. (2000) have examined the highly aromatic, macromolecular kerogen in Murchison using hydrous pyrolysis techniques and demonstrated that it contains a significant portion of organic N that is somewhat isotopically heavy ($\delta^{15}N < +30\%o$). Thus, at least a fraction of the aromatics in meteorites appear to be a carrier of isotopically anomalous N, but the nature of the relationship is not understood.

IDPs are also known to contain D-enriched components, and these components are often distributed heterogeneously within the particles on very small scales (McKeegan et al., 1984, 1985; Messenger, 2000). Ion microprobe studies of the D distribution and correlation with other elements suggest tentative links between the D and a carbonaceous carrier (McKeegan et al., 1985; Messenger et al., 1995; Aléon et al., 2000; Messenger, 2000). IDPs are known to contain abundant PAHs (Allamandola et al., 1987; Clemett et al., 1993), but the relationship between the PAHs and D-enrichments is not clear. So far, combined ion probe and two-step laser mass spectrometry studies of the same IDP fragments have only detected PAHs in D-rich IDPs, but D-enrichments have been seen in IDPs in which PAHs were not clearly detected (Messenger et al., 1995). Aléon et al. (2000) recently reported that $\delta D$ increases in IDPs with increased C/H, with most enrichments occurring when C/H > 1. They suggested their measurements could be explained by the presence of three phases: (i) a material similar to the kerogen in carbonaceous chondrites with C/H = 1.0-1.5 and a relatively small $\delta D$, (ii) a component dominated by small molecules with C/H = 1 and larger D enrichments, and (iii) a refractory organic material with C/H
= 3 and the largest D enrichments. The latter phase was observed to be the major carrier of the D and its C/H ratio well in excess of 1 suggests the carrier is aromatic.

Stadermann et al. (1989) used ion microprobe techniques to examine the C and N isotopes in 24 IDPs. They found no $^{13}$C enrichments and confirmed earlier observations that $\delta^{13}$C values showed little variation between different fragments. In contrast, they found that 10 of the 24 IDPs contained heavy N with values of $\delta^{15}$N up to +442%o and, as with the D, the $^{15}$N showed large spatial variations within an individual particle. The normal $^{13}$C ratios and high degree of spatial variability of D and $^{15}$N enhancements within IDPs was confirmed by Messenger (2000), who also noted that the D and $^{15}$N appeared to be qualitatively correlated, at least in the broad sense that fragile "cluster" IDPs seem to be more generally D- and $^{15}$N-rich than individual IDPs. One of these IDPs, Dragonfly, was found to contain some of the most extreme $\delta$D and $\delta^{15}$N values ($\delta$D > 24,800%o and $\delta^{15}$N > 340%o) ever seen in a natural material. Keller et al. (2000) used a variety of analytical techniques to show that $\delta$D and $\delta^{15}$N correlate with each other in this particle. EELS measurements indicated that the N was associated with the carbonaceous fraction, although the concentration was too low to specifically identify the N-containing component. The XANES measurements of the carbonaceous material showed prominent C-C, C-H, and C-O absorptions and suggested that the D is carried in C-rich grains that contain aldehydes.

In summary, studies of the D-enrichments in meteorites and IDPs suggest that at least a portion of the PAHs in meteoritic material have an interstellar origin. The D enrichments appear to be decoupled from carbon isotopic ratios, while there is a possible relationship between D excesses and excess $^{15}$N. The processes by which the D enrichments were created and the true relationships (if any) of the D enrichments to other isotopic systems remain to be clarified.
Placing Constraints on the Interstellar Processes Responsible for Meteoritic D-enrichments

It is clear that the regiochemistry of D-enrichments in the PAHs in meteoritic materials and correlation (or lack thereof) with the presence of specific aromatic compounds or the isotopic systematics of C, O, and N could place constraints on the interstellar enrichment processes being sampled (see Table 1). We now discuss the meteoritic evidence for or against contributions from the various enrichment processes and point out potential measurements that might help elucidating their relative importance for the creation of meteoritic organics.

Low Temperature Gas Phase Ion-Molecule Reactions

There are several tests that could constrain the relative importance of ion-molecule reactions for the D fractionation in meteoritic PAHs. First, it would be useful if the presence of PAHD$^+$ ions could be verified in the gas phase in dense interstellar clouds. In principle, it might be possible to determine PAD/PAH ratios at infrared frequencies using the relative strengths of the aromatic C-H and C-D stretching bands near 3050 and 2270 cm$^{-1}$, respectively (Hudgins et al., 1994). However, the low expected gas phase abundances of PAHs in such cold environments will make this very difficult. In addition, PAHs are most easily observed in the gas phase when they are excited by UV photons and radiatively cool by emitting a cascade of infrared photons. At the low UV fluxes present in typical dense clouds, this excitation mechanism is strongly curtailed. There may be more hope of measuring the relative strengths of the C-H and C-D aromatic stretching bands in absorption, but in this case the bands would presumably be dominated by PAHs frozen onto grains and it would be difficult to assign any observed fractionation specifically to ion-molecule reactions, as opposed to other processes.

Theoretical models for D enrichments in PAHs by gas phase ion-molecule reactions predict that this process will initially preferentially deuterate the larger PAHs in dense clouds, although if sufficient time is available the D/H ratio of all the PAHs in the gas phase should ultimately equilibrate with the gas phase value. Thus, a second, meteoritic test for this process would be to examine meteoritic organics for correlations between the D/H ratio and the size of the aromatic...
moieties of the carriers. In this respect, the report by Aléon et al. (2000) that in IDPs the value of δD increases with increased C/H, with most enrichments occurring when C/H > 1 is suggestive; such a distribution is qualitatively consistent with a fractionation of the aromatic via ion-molecule reactions. If ion-molecule reactions were the sole source of the observed fractionations seen by Aléon et al., the generally lower δD at lower C/H (smaller aromatics) would then be explained by short interstellar evolutionary times (<3x10⁵ years).

Low Temperature Gas-Grain Reactions

Theoretical models and telescopic observations suggest that gas-grain reactions probably result in the single largest reservoir of D fractionated materials in dense clouds (Tielens, 1983; Teixeira et al., 1999). However, the D enrichments produced by gas-grain reactions are expected to reside predominantly in simple hydrides like CH₄, NH₃, and H₂O, not PAHs. These simple, volatile molecules are not significant carriers of the D found in meteorites, implying that interstellar gas-grain reactions did not play a direct role in the creation of the D enrichment seen in meteorites.

As mentioned earlier, there are no strong theoretical or experimental reasons to expect that PAHs can be directly enriched in D by gas-grain reactions. If PAHs condensed in interstellar grain mantles can participate in gas-grain reduction reactions, the resulting products would be partially reduced PAHs (Hₐ-PAHs). Such structures are seen in meteoritic organics (Cronin and Pizzarello 1990), but so far no attempt has been made to determine the δD systematics of these materials. A better understanding of the δD values in reduced meteoritic PAHs (Hₐ-PAHs and cyclic aliphatics) would clearly be of interest.

Independent of its ability to directly produce D-enriched PAHs or Hₐ-PAHs, this process may still play a key secondary role in the production of D-enriched meteoritic organics. Since gas-grain reactions probably do produce large reservoirs of D-enriched, simple hydrides in ices in dense clouds, this process may be largely responsible for providing the "feedstock" that subsequent chemistry converts into the more complex and refractory species that carry the D in meteorites.
Unimolecular Photodissociation Reactions

Unimolecular photodissociation reactions involving interstellar PAHs are expected to generate a characteristic D-enrichment signature in which the largest D enrichment occur in small PAHs ($C_{10}$ to $C_{50}$). This is in contrast to the signature expected from ion-molecule reactions, which should tend to preferentially enrich larger PAHs. There are good theoretical and experimental reasons to expect this process occurs in the interstellar medium, but there is currently no confirming observational evidence. The challenge is to use the infrared C-D and C-H stretching emission bands near 2270 and 3050 cm$^{-1}$, respectively, to measure the PAD/PAH ratio in a variety of interstellar environments that sample different UV radiation fields and astrochemical evolutionary histories. If D-enrichment by unimolecular photodissociation is occurring, PAD/PAH ratios should be very low in the outflows from carbon stars, where the PAHs have undergone relatively little UV processing and the outflows have D/H = 0 (due to the burning of much of the D in the star). PAD/PAH ratios should however, be much larger in diffuse interstellar 'cirrus' clouds, HII regions, and reflection nebulae, since these PAHs will have had longer UV exposures in environments having higher gas phase D/H ratios.

While it will still be difficult to attain the high signal-to-noise emission data required to detect PADs at the expected PAD/PAH ratios of $< 10^2$, it should be easier than detecting deuterated PAHs in absorption in dense clouds. First, a larger fraction of the PAH population will be in the gas phase in the warmer environments where most unimolecular photodissociation reactions will occur. In addition, the same UV photons driving the unimolecular photodissociation will excite the PAHs and cause them to emit in the infrared, thereby making them easier to detect. The quality of current telescopic infrared spectra is constantly improving and with appropriate spaceborne instrumentation it should be possible to place meaningful constraints on the efficiency of PAH D-enrichment by unimolecular photodissociation reactions in a variety interstellar environments.

There is some inconclusive evidence that suggests that unimolecular photodissociation reactions could have played role in the production of the D-enrichments seen in meteorites. The D/H ratio of acid-soluble hydrocarbon phases in meteorites increases as C/H increases from 0.2 to
1.5 (Yang and Epstein, 1983), while the D/H ratio in acid insoluble phases drops slightly as the C/H ratio increases from 2.3 to 3.0 (Robert and Epstein, 1982). Allamandola et al. (1987) noted that if it is assumed that the soluble and insoluble phases represent different degrees of molecular complexity of related materials (potentially a gross oversimplification), then the peak in D enrichment at C/H ~ 2 would imply the dominance of PAHs containing 20-50 C atoms, consistent with unimolecular photodissociation. As is the case for investigating the role of ion-molecule reactions, the most useful tests of meteoritic organics would be to measure the correlation (if any) between D/H ratios and the size of the aromatic moieties of the carriers.

**Ultraviolet Photolysis Reactions in D-enriched Ice Mantles**

The UV irradiation of PAHs in ices in dense molecular clouds should result in the production of a variety of aromatic ketones, alcohols, ethers, and H$_x$PAHs (Bernstein et al., 1999, 2001). If the ices are themselves D-enriched, the same irradiation results in the equilibration of D in the ice with the PAHs (Sandford et al., 2000). Thus, this process may play an important role in 'fixing' isotopic fractionations produced by other processes into molecular forms that can survive incorporation into the solar nebula and subsequent parent body accretion and alteration.

In principle, one could examine meteoritic organics for evidence of interstellar UV ice irradiation by searching for isotopic fractionations in putative or probable photolysis products. Such an approach is complicated by several issues, however. First, the photoproducts will exhibit H, C, N, and O isotopic ratios that reflect the relative mixing, in the ice, of components produced by several other fractionation processes. Second, each of the fractionation processes represented in the ice will produce a different characteristic pattern of fractionation in different molecules which will, in turn, be converted with different efficiencies into different complex organics. Thus, the ultimate isotopic signature of the irradiation products of interstellar ices probably represents a complex convolution of many effects.

Fortunately, aromatics, which carry most of the D in meteorites, also offer the best hope of providing a distinctive signature of ice irradiation. The deuteration of PAHs by photolysis in D-
enrich ices proceeds along several different paths: H→D exchange reactions on aromatic rings, D-atom addition reactions that convert aromatic rings to aliphatic rings, and keto-enol tautomeric exchange (see Figure 5). Each of these processes results in the placement of acquired D into characteristic regiochemical sites on the PAHs. Ideally, one would attempt to search for correlations of D-enrichments associated with these different kinds of regiochemical sites on meteoritic PAHs. We would note that since these sites have different labilities, they would be expected to suffer differing degrees of exchange with their surroundings during parent body processing, delivery, recovery, and analysis.

It would also be extremely useful to search for correlations of D-enrichment with PAH oxidation and reduction state in meteoritic organics. In this regard, the recent report by Keller et al. (2000) that the degree of D-enrichment in IDPs may correlate with the presence of C-O bonds is of particular interest. Reports that the enrichments of D and 15N may be correlated in IDPs (Keller et al., 2000; Messenger, 2000) is worthy of additional comment. Preliminary photolysis experiments of PAHs in NH3-containing ices suggest that nitrogen can also be affixed to the edges of PAHs in the form of -NH2. Since N in NH3 is expected to become 15N-enriched via ion-molecule reactions (Terzieva and Herbst 2000; Charnley and Rodgers, 2001) and NH3 is abundant in interstellar ices (Lacy et al., 1998), it would clearly be interesting to see if the observed meteoritic 15N-enrichments correlate in any way with the aromatics, particularly in the form of -NH2 sidegroups.

CONCLUSIONS

There are a number of interstellar processes that can produce significant D fractionation in their products, the four main ones being gas phase ion-molecule reactions, low temperature gas-grain reactions, gas phase unimolecular photodissociation, and ultraviolet photolysis in D-enriched ice mantles. Some of these processes may also produce isotopic fractionations in the elements C, N, and O, with the fractionations in N most likely to be significant. In the case of polycyclic aromatic hydrocarbons (PAHs), each of these processes is associated with distinct regiochemical signatures (D placement on the molecule, correlation with specific chemical functionalities, etc.).
Gas phase ion-molecule reactions initially preferentially deuterate the larger PAHs in dense clouds, but if sufficient time is available, all PAHs in the gas phase in dense clouds should achieve steady-state with the gas phase. Thus, the characteristic signature of this process would be fully aromatic PAHs whose D/H ratios are uniform across the PAH population or that increase with increasing PAH size. Gas-grain reactions produce D fractionations in dense clouds that are manifested primarily in simple hydride molecules. PAHs probably cannot be enriched in D in this manner, but if they are, D enrichments would be manifested in $H_n$-PAHs, i.e., hydrogenated PAHs that contain cyclic aromatic and aliphatic rings. Deuterium enrichment via gas phase unimolecular photodissociation reactions differs from other processes in that low ambient temperatures are not required to produce fractionations. This process should produce a fractionation pattern in which PAHs in the C$_{10}$-C$_{20}$ size range show the largest enrichments. The principle effect of energetic irradiation of interstellar ices is to transfer pre-existing fractionations from simple, volatile compounds to more complex, refractory compounds. This process may play an important role in 'fixing' isotopic fractionations produced by other processes into molecular forms that can survive incorporation into the solar nebula and subsequent parent body accretion and alteration. Deuterium enrichments produced in aromatic species by this process will be largely independent of PAH molecular size and should show very specific regiochemical behaviors. PAHs enriched in this manner should show correlations between $\delta^D$ and the presence of both reduced PAHs ($H_n$-PAHs) and oxidized PAHs (aromatic alcohols, ketones, and ethers).

Our poor current understanding of the molecular siting of the excess D and $^{15}$N in meteorites makes it difficult to evaluate the relative contributions of these interstellar processes to the organic inventory in meteorites. The present data on the relationship of $\delta^D$ with C/H ratio in macromolecular meteoritic organics does not preclude significant contributions from ion-molecule reactions, unimolecular photodissociation reactions, or UV driven exchange in interstellar ices. Reported correlations of D enrichment with the presence of C-O bonds in IDPs is consistent with PAHs enriched by photolysis in ices enriched in D by gas-grain reactions.
The isotopic fractionations in N, and the lack of strong fractionation effects in C and O, are also qualitatively consistent with these processes. The $^{15}$N enrichments, and their possible correlation with $\delta$D, may be explainable in terms of low-temperature ion-molecule reactions, possibly followed by UV driven exchange reactions in ices. That D and $^{15}$N enrichments do not correlate tightly could be due to several effects - (i) the fractionations of these two isotopes have different temperature dependences, (ii) meteoritic organics may contain differing contributions from processes like gas-grain reactions that produce D enrichments without corresponding $^{15}$N enrichments, (iii) the efficiency with which UV photolysis will 'fix' anomalies into more complex species depends on the molecular composition of the ices, which can vary within dense clouds, and (iv) differences in the labilities of D and $^{15}$N may alter the enrichment ratios during parent body processing of the meteoritic organics. The lack of correlated isotopic effects in O are not surprising since current theory and experimental data suggest that these processes should not produce strong fractionations in O in the ISM. The lack of C isotopic effects is a little more surprising, but the fact that C fractionations can proceed along two main paths that fractionate their products in opposite directions may moderate the size of any final effect in the ISM.

Thus, there is currently no reason to reject the possibility that all the processes described in this paper may have played a part in the production of the D enrichments seen in meteorites. Determining which processes dominated the overall inventory in meteorites must await additional measurements. Studies that search for the characteristic enrichment patterns expected for these four process (summarized in Table 1) would be particularly useful.

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REFERENCES


Table 1: PAH Interstellar D-Enrichment Processes and their Signatures

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<tbody>
<tr>
<td>Gas Phase Ion-Molecule Reactions</td>
<td>YES</td>
<td>NO</td>
<td>NO</td>
<td>YES</td>
<td>Favors PAHs with larger numbers of peripheral H atoms (larger PAHs). Probably independent of regiochemistry.</td>
</tr>
<tr>
<td>Unimolecular Photodissociation Reactions</td>
<td>NO</td>
<td>YES</td>
<td>NO</td>
<td>Probably NO</td>
<td>Favors smaller PAHs (Number of C atoms ≤ 50). Probably independent of regiochemistry.</td>
</tr>
<tr>
<td>Gas-Grain Reactions</td>
<td>YES</td>
<td>NO</td>
<td>NO</td>
<td>YES?</td>
<td>May not occur. If it does, the enrichment will reside on the reduced rings of H₈-PAHs.</td>
</tr>
<tr>
<td>Ice-PAH Photolysis Reactions</td>
<td>YES</td>
<td>YES</td>
<td>Variable*</td>
<td>Possibly</td>
<td>Size independent. Enrichment will correlate with oxidation, reduction, and regiochemistry.</td>
</tr>
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</table>

(a) The deuterium associated with oxidized rings on aromatic alcohols and ketones will be labile. Deuterium residing on the reduced rings of H₈-PAHs and on the fully aromatic rings of PAHs will be non-labile.
FIGURE CAPTIONS

Figure 1 - PAHs become enriched in D via gas phase ion-molecule reactions in a two step process (shown here for the PAH pyrene). First, a PAH reacts with an H₂D⁺ ion to pick up an excess deuterium atom and charge. Subsequently, reaction with an electron neutralizes the charge and the PAH eliminates an excess H or D atom. This process should initially preferentially deuterate larger PAHs in dense clouds, but ultimately, all PAHs in the gas phase would be expected to D enrichment as they approach steady-state with the gas phase.

Figure 2 - In dense clouds where H/H₂ is large, gas-grain reactions primarily serve to hydrogenate, and D-enrich, species in the ice. It is not clear if PAHs can be hydrogenated in this manner, but to the extent that this process occurs, any D enrichments generated would be expected to correlate with the presence of Hₙ-PAHs and cyclic aliphatics.

Figure 3 - Gas phase PAHs can become enriched in D by repeated unimolecular photodissociation events that favor retention of D over H because of its lower zero point energy. The extent of the D-enrichment depends critically on the size of the PAH. The D photoenrichment of PAHs should be most significant for PAHs in the range of sizes spanned by naphthalene (C₁₀H₈) and hexabenzocoronene (C₄₂H₁₆). Enrichment in larger PAHs is not expected because they have large numbers of vibrational modes and are stable against photolytic bond rupture. Unlike other D fractionation processes, enrichment by unimolecular photodissociation does not require low ambient temperatures.

Figure 4 - PAHs undergo both oxidation and reduction reactions that affect their edge structures when they are irradiated in H₂O-rich ices. Oxidation results in the production of aromatic alcohols, ketones, and, in PAHs with 'bay' regions in their edge structures, ethers. Reduction leads to the formation of Hₙ-PAHs, molecules having mixed aromatic and aliphatic rings.

Figure 5 - The UV photolysis of PAHs in D-enriched ices results in rapid D enrichment of the PAHs through aromatic D→H exchange, D-atom addition, and exchange through keto-enol tautomerism. Deuterium enrichments produced in this manner are largely independent of PAH molecular size, but should show specific regiochemical patterns and would be expected to correlate with the presence of both reduced PAHs (Hₙ-PAHs) and oxidized PAHs (aromatic alcohols, ketones, and ethers).
The Deuteration of PAHs by Ion Molecule Reactions

Figure 1
Deuterium Enrichment of PAHs by Gas-Grain Reactions

Figure 2
Deuterium Enrichment of Gas Phase PAHs by Unimolecular Photodissociation

Small PAHs become D-enriched

UV Photon
Cold Pyrene → Hot Pyrene → Cold Pyrene

Larger PAHs Don't

UV Photon
Cold PAH → Hot PAH

IR Photon
Cold PAH
The Altered Edge Structures Produced when PAHs are UV Irradiated in H$_2$O Ice
Reaction Paths for the Deuteration of Coronene

(a) UV radiation needed
\[ \text{D}_2\text{O ice at 10 K} \]

(b) UV radiation needed
\[ \text{D}_2\text{O ice at 10 K} \]

(c) \[ \text{radiation not required} \]