HIGH-DENSITY AMORPHOUS ICE, THE FROST ON INTERSTELLAR GRAINS

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ABSTRACT

Most water ice in the universe is in a form which does not occur naturally on Earth and of which only minimal amounts have been made in the laboratory. We have encountered this "high-density amorphous ice" in electron diffraction experiments of low-temperature (T < 30 K) vapor-deposited water and have subsequently modeled its structure using molecular dynamics simulations. The characteristic feature of high-density amorphous ice is the presence of "interstitial" oxygen pair distances between 3 and 4 Å. However, we find that the structure is best described as a collapsed lattice of the more familiar low-density amorphous form. These distortions are frozen in at temperatures below 38 K because, we propose, it requires the breaking of one hydrogen bond, on average, per molecule to relieve the strain and to restructure the lattice to that of low-density amorphous ice. Several features of astrophysical ice analogs studied in laboratory experiments are readily explained by the structural transition from high-density amorphous ice into low-density amorphous ice. Changes in the shape of the 3.07 μm water band, trapping efficiency of CO, CO loss, changes in the CO band structure, and the recombination of radicals induced by low-temperature UV photolysis all covary with structural changes that occur in the ice during this amorphous to amorphous transition. While the 3.07 μm ice band in various astronomical environments can be modeled with spectra of simple mixtures of amorphous and crystalline forms, the contribution of the high-density amorphous form nearly always dominates.

Subject headings: dust, extinction — infrared: ISM: lines and bands — ISM: molecules — line: profiles — methods: laboratory — molecular data

1. INTRODUCTION

The frost on interstellar grains in dense molecular clouds is thought to consist mainly of an amorphous water ice with trapped impurities (e.g., Tielens & Allamandola 1987). The observed 3.07 μm absorption band is usually best compared to the OH stretch vibration band of impure water ice, vapor-deposited onto inert surfaces at temperatures less than 30 K (Hagen, Tielens, & Greenberg 1981; Schmitt et al. 1989). The same, or similar, ices are thought to constitute the bulk of comet nuclei (Mumma, Weissman, & Stern 1993; Kouchi et al. 1994). The 3.07 μm ice band has been studied in laboratory experiments of interstellar ice analogs as a function of temperature, deposition conditions, and impurity content (Hagen et al. 1981; Sceats & Rice 1982. Hagen et al. 1983; Tielens, Hagen, & Greenberg 1983; Schmitt, Ocampo, & Klinger 1987). In addition, considerable effort has gone into measuring the changing shapes of infrared absorption bands of trapped impurities, the diffusion and recombination of radicals induced by UV photolysis, and the outgassing properties of ice mixtures (Bar-Nun, Herman, & Laufer 1985; Bar-Nun et al. 1987; Grim & Greenberg 1987; Laufer, Kochavi, & Bar-Nun 1987; Sanford et al. 1988; Schutte 1988; Schutte, Allamandola, & Sanford 1993).

The observed characteristics of vapor-deposited impure water ices are a function of the arrangement of the water molecules in the solid, which can be studied directly by electron or X-ray diffraction. Fourier transformation of the observed diffraction patterns yields a statistical distribution of distances between the oxygen atoms in the water matrix (the oxygen-oxygen radial distribution function), which uniquely characterizes its structure. By applying electron diffraction to dynamic systems, we find that vapor-deposited water ice exists in three amorphous forms, apart from two crystalline states (Jenniskens & Blake 1994).

This paper characterizes the two low-temperature amorphous forms of vapor-deposited water ice, low-density amorphous ice (l, I) and high-density amorphous ice (l, h), and the behavior of these ices in the temperature range 15—120 K. An amorphous form of water ice was first described early in this century. Diffraction studies were performed by Burton & Oliver (1935). This familiar low-density form is made when water vapor is deposited on an inert surface at 77 K. I, l has a density of 0.94 ± 0.03 g cm⁻³ (Narten, Venkatesh, & Rice 1976; Mayer & Pletzler 1986) and has been studied extensively because of its resemblance to liquid water (Olander & Rice 1972; Sceats & Rice 1982; Pohorille et al. 1987; Scintitti, Geiger, & Stanley 1992), and because it can be used to replace liquid water as an embedding material for biological tissues in electron microscopy (Heide 1984; Heide & Zeitler 1985).

High-density amorphous ice was first reported by Narten et al. (1976). X-ray diffraction patterns of water vapor deposited at 10 K on a single crystal copper substrate showed the presence of oxygen atoms at distances between those of the first and second nearest neighbor at the interstitial site of the l, I network, which results in a higher value for the calculated local density of oxygen atoms, i.e., 1.1 ± 0.1 instead of 0.94 g cm⁻³. For this reason, Narten et al. introduced the names of "high density" (I, H—our notation) and "low density" (I, l) amorphous ice. Later experiments failed to reproduce the high-density form on substrates other than single-crystal copper, and it was thought that the high-density amorphous form was an artifact of epitaxial growth on an oriented single crystal (Sivakumar, Rice, & Sceats 1978; Sceats & Rice 1982), depending perhaps on a nucleation step (Sivakumar et al. 1977).

Recently, it was shown that high-density ice also forms during vapor deposition on amorphous carbon films (Heide 1984; Jenniskens & Blake 1994). However, it has not been shown that the resulting ice structure is the same as that of Narten
et al. (1976). This high-density amorphous ice is potentially important in astrophysics, because infrared spectra of low-temperature deposits provide the best fit to the 3.07 \( \mu \)m ice band (Hagen et al. 1981). In the astrophysical literature, the high-density form has only briefly been referred to before by Hagen et al.

Another form of high-density amorphous ice can be made by pressure melting, and is similarly referred to as high-density amorphous ice (hda or HDA). When cooled to 77 \( K \), crystalline hexagonal ice, \( I_h \), undergoes a transition into hda at 1–2 GPa pressure (Mishima, Calvert, & Whalley 1984, 1985). The resulting ice, when recovered at 1 atm, has a density of 1.17 ± 0.03 g cm\(^{-3}\) (determined by weighing in liquid nitrogen). Theoretical estimates of the transition pressure as a function of temperature are given in Tse & Klein (1990) and Poole et al. (1992). HDA can exist metastable for some time at low pressure and 77 \( K \). When this pressure-induced, high-density amorphous ice is warmed at a rate of 0.2–3 \( K \) minutes\(^{-1}\), the ice transforms irreversibly into a low-density 0.94 ± 0.02 g cm\(^{-3}\) amorphous form (lda or LDA) in the range 105–123 \( K \), releasing energy in the process (Mishima et al. 1984, 1985; Handa, Mishima, & Whalley 1986; Bizid et al. 1987; Tse & Klein 1987; Hemley, Chen, & Mao 1989). The radial distribution function of hda (Bosio, Johari, & Teixera 1986; Bizid et al. 1987) differs significantly from that of the \( I_h \) of Narten et al. (1976). It has been argued that pressure-induced hda has more hexagonal order than vapor-deposited \( I_h \) (Tse 1992).

In this paper, we confirm the existence of the vapor-deposited, high-density amorphous form \( I_h \) from electron microscopy studies (§ 2). We measured the radial distribution function (rdf) of \( I_h \) and the gradual change of the diffraction pattern during the \( I_h \rightarrow I_{ha} \) transition (§ 3, § 4). The rdf is compared to that of pressure-induced hda and similar data by Narten et al. (1976). The distinct physical properties of \( I_h \) (§ 5) and the conditions necessary for formation during vapor deposition (§ 6) are determined by reinterpreting experimental data reported in the literature. In § 7, new molecular dynamics calculations are presented that attempt to model the ultrastructure of high-density amorphous ice. The theoretical and experimental results are discussed in § 8. In § 9, it is argued that high-density ice is ubiquitous in space, to an extent that this form is probably the most abundant form of ice in the universe.

2. CHARACTERIZING THE STRUCTURE OF AMORPHOUS ICES

The electron diffraction studies employ a Hitachi H-500H transmission electron microscope (TEM), which was adapted for vapor-depositing water molecules in situ onto a thin amorphous carbon film substrate held at a temperature of 14 K (Blake & Palmer 1991). The structure of the ice film is characterized by selected area electron diffraction (SAED). Morphological features of the ice can be viewed in either bright-field mode (strongly diffracting regions of the film appear dark) or dark-field mode (diffracting regions appear bright).

Degassed distilled water vapor is deposited from a reservoir at 14 \( torr \) and room temperature into the microscope at base pressure of ~3 \( \times \) \( 10^{-7} \) torr, until the partial \( H_2O \) pressure increases to ~4 \( \times \) \( 10^{-6} \) torr. A film of ice is grown by opening a shutter that covers the sample for 5–20 s (rate of growth 6–36 \( \mu \)m hr\(^{-1}\)), while monitoring the diffraction pattern. The resulting film is about 0.05 \( \mu \)m thick, smooth, and homogeneous. Bright-field imaging reveals no sign of the crystallinity reported by Olander & Rice (1972), even at high magnification. There is also no indication of needle- or stalactite-like surfaces as reported by Laufer et al. (1987).

Diffraction patterns from the ice film are obtained from volumes of material of about 0.7 \( \mu \)m\(^3\). A parallel electron beam of 3 \( \mu \)m in diameter is directed through the sample to form the diffraction pattern, which is focused onto an imaging device at the back focal plane of the microscope. The resulting diffraction pattern is a Fourier transform of the statistical distribution of distances between atoms in the ice. Oxygen atoms are the principal electron scatterers (scattering by hydrogen atoms is negligible). The diffraction pattern is electronically recorded with an instrumental resolution of \( \Delta k = 0.08 \) \( Å^{-1} \), sampled at 0.03 \( Å^{-1} \) pixel\(^{-1}\), and the patterns are stored at a rate of 1–3 per minute during a typical experiment. Temperatures can be recorded with an absolute accuracy of ± 2 \( K \) and relative accuracy of about 0.2 \( K \). The warming rate in these experiments varied from 2.3–5.2 \( K \) minutes\(^{-1}\) and was typically 2.7 \( K \) minutes\(^{-1}\).

The two-dimensional circular diffraction patterns are reduced by folding to a one-dimensional radial intensity distribution (Fig. 1). The \( x \)-axis, the momentum transfer coordinate \( k = (4\pi/\lambda) \sin \theta \), is calibrated using crystalline gold (\( \sigma_0 = 4.0786 \) \( Å \)) deposited on a small part of the substrate as an internal standard, which was measured at the beginning and end of each experiment. The relatively small intensity contributed by the amorphous carbon substrate is removed by a background subtraction algorithm.

Low electron dose imaging techniques are employed to minimize sample degradation during viewing. Electron beam damage does not change the diffraction patterns on the timescale of each measurement, and different parts of the ice layer are exposed during each subsequent measurement. The technique offers a powerful means of measuring the structural properties of water ice films in situ in vacuum.

Fourier transformation of the diffraction patterns yields the rdf of distances between oxygen atom pairs (Fujime 1966;
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Moine, Pelton, & Sinclair 1988)

\[ k = \left(\frac{4\pi \sin \theta}{\lambda}\right), \]

\[ I(k) = \frac{[I_{\text{obs}}(k) - I_{\text{bg}}(k)]}{\beta f_0^2}, \]

\[ \text{rdf} = \frac{(2/\pi)}{\int_0^{2\pi}} I(k) \exp(-zk^2) \sin(kr)drdk, \hspace{1cm} (1) \]

where \( I_{\text{obs}} \) is the observed diffraction pattern after subtraction of a substrate contribution, \( I_{\text{bg}} \) is a smooth and continuous function through the inversion points of the pattern (a sum of three power laws; Fig. 1), \( \beta \) is a scaling factor taken such that the intensity is -1 for small \( k \) after subtraction of the background (at onset of first diffraction peak), and \( f_0 \) is the electron scattering function of oxygen. The parameter \( \alpha \), in the damping factor in the Fourier transform, is taken such that \( \exp(-ek^2) = 0.1 \). The latter reflects the rapid decrease of the electron beam intensity away from the center of the beam and the limited range of \( k \) that is measured.

Due to limitations in the dynamic range of our electronic imaging system, we cannot derive an rdf for every recorded pattern. In order to follow the changes in the structural state, we measure the position, height, and width of each peak in the diffraction pattern by fitting Gaussian profiles to individual maxima. The Gaussian profiles usually produce a good fit to the peaks, which is the only and sufficient reason for choosing this particular functional form.

3. THE RADIAL DISTRIBUTION FUNCTION OF \( I_{\text{ah}} \) AND \( I_{\text{al}} \)

When water ice is formed by vapor deposition at 14 K, it yields a distinct diffraction pattern when compared to that of the more familiar low-density amorphous ice at 77 K (Fig. 1). Most notably, the first diffraction peak is shifted to larger values of \( k \).

The resulting radial distribution function of the low-temperature deposit (Fig. 2), which Narten et al. (1976) calls "high-density amorphous ice" (\( I_{\text{ah}} \)—our notation), differs principally from the higher temperature deposit, \( I_{\text{al}} \), in having an additional maximum between 3 and 4 \( \AA \).

It is illustrative to compare the radial distribution functions of \( I_{\text{ah}} \) and \( I_{\text{al}} \) to those of vapor-deposited amorphous ice at 10 K (Narten et al. 1976), and those of pressure amorphosed hexagonal ice (Bizid et al. 1987). Figure 2 shows that there is a striking similarity between our rdf of vapor-deposited \( I_{\text{ah}} \) and that of the pressure-induced D\(_2\)O ice by Bizid et al. (1987). The high-density amorphous form is characterized, in both cases, by a splitting of the second nearest neighbor peak, such that one component lies between the first and second nearest neighbor peaks of the low-density amorphous form. The diffraction maxima in \( I_{\text{ah}} \) are less pronounced, suggesting that \( I_{\text{ah}} \) is less well ordered than Bizid's hda. This apparent reduction in ordering may be due to the thinness of the vapor-deposited ice, differences in annealing times between the two ices, exposure to high fluxes of X-rays, or more microporosity in the vapor-deposited ice (that is, defects in the random hydrogen-bonded network with typical cavity sizes of radius 1 \( \AA \) < \( r \) < 10 \( \AA \)).

The agreement with the pressure-induced hda is better than with the vapor deposit of Narten et al. (1976). The first nearest neighbor distance of both \( I_{\text{ah}} \) and \( I_{\text{al}} \) is 2.81 (+0.05/-0.01) \( \AA \), in good agreement with the hda of Bizid et al. (1987) (2.80 \( \AA \)) and Bosio et al. (1986) (~2.82 \( \AA \)), while Narten et al. (1976) report a slightly smaller value at 2.76 \( \AA \). The second nearest neighbor in our \( I_{\text{ah}} \) is broad and is split into two peaks at 3.70 \( \AA \) and 4.70 \( \AA \), similar to the hda of Bizid et al., 3.70 and 4.64 \( \AA \); and that of Bosio et al., 3.65 and 4.45 \( \AA \), respectively. However, neither our, nor Bizid's, radial distribution functions show a sharp spike at the "interstitial" position as in the Narten et al. (1976) data.

![Graph](image-url)
also be identified indirectly. We have reinterpreted experimental data reported in the literature, in light of the previous results, and suggest that the high-density amorphous form can be identified by the shape of the 3.07 \( \mu \text{m} \) band (Hagen et al. 1981; Mayer & Pletzer 1985), the shape of the dangling OH bond absorption bands (Venkatesh, Rice, & Bates 1975; Sivakumar et al. 1978), the presence of CO impurities (Sandford et al. 1988; Schmitt, Greenberg, & Grim 1989), outgassing behavior (Lauffer et al. 1987; Kouchi 1990), and the diffusion of radicals (Schutte 1988).

Figure 4 summarizes the deposition conditions for the experiments cited above. It is concluded that \( I_{\alpha} h \) forms readily when water is condensed slowly (\(< 100 \mu \text{m hr}^{-1}\)) at low enough temperature (\( T < 40 \text{ K} \)) on any surface with good thermal conductivity (single-crystal copper, amorphous carbon film, aluminum, gold-coated copper, CsI window). Vapor deposits at 40–70 K form mostly \( I_{\alpha} l \) (see, e.g., Sandford et al. 1988).

There is a maximum and a minimum condensation rate at which \( I_{\alpha} h \) can form. Deposition rates close to the maximum of 100 \( \mu \text{m hr}^{-1} \) do result in a structure with more \( I_{\alpha} l \) character. For example, Palumbo & Strazzulla (1993) measured a noticeable change in the position of the C–O stretch in an impure water ice for deposition rates above 1 \( \mu \text{m hr}^{-1} \). For a given temperature, there is, potentially, a minimum deposition rate, below which the water molecules have time to reorient themselves before being incorporated in the \( I_{\alpha} h \) lattice by subsequent condensing molecules. Kouchi et al. (1994) measured this effect for the formation of the cubic crystal form in the low-density amorphous domain at very low deposition rates, and found a temperature dependence, which, to us, suggests a characteristic \( \Delta H = 44 \pm 7 \text{ kJ mol}^{-1} \) activation enthalpy. This value is the same as that for the crystallization of \( I_{\alpha} l \) during warm-up (Jenniskens & Blake 1994). An extrapolation of the measured critical flux at 96–110 K (Kouchi et al. 1994) down to lower temperature suggests that, at realistically low deposition rates, no cubic crystalline ice will be formed in the high-density regime below 40 K. However, a lower activation energy may apply for the reorientation of the water into an open \( I_{\alpha} l \) structure. The horizontal line in Figure 4 shows this deposition rate dependence, if the characteristic activation enthalpy is 12 kJ mol\(^{-1}\) (see § 8.2). The result suggests that this effect plays no significant role in the laboratory experiments, but may be relevant in case of extremely low deposition rates in some astrophysical environments.

The film thickness has an effect as well. Most experiments concern layers of 0.01–10 \( \mu \text{m} \) in thickness. Low-temperature vapor deposits as thick as 1000 \( \mu \text{m} \) sometimes do (Narten et al. 1976), and sometimes do not, show \( I_{\alpha} h \) signature (Venkatesh et al. 1975). Perhaps this is because in thicker films the heat of fusion from impinging molecules is not conducted fast enough to prevent local reordering. There is also some indication that for very thin layers (\(< 0.002 \mu \text{m} \)), substrate surface structure can affect the structural growth during deposition (Moore et al. 1994) and, therefore, the presence of a surface film on the substrate, and the existence of single-crystal orientations of the substrate, may affect the structure of thin deposits through epitaxy or nucleation (Scots & Rice 1982).

Finally, we note that the \( I_{\alpha} h \rightarrow I_{\alpha} l \) transition does not always unfold the same way in vapor-deposited layers. For example, Hagen et al. (1981) studied a 0.5 \( \mu \text{m} \) thick water ice layer, vapor-deposited at 10 K. We assume that the observed shift of the peak frequency and width of the infrared O–H stretching band is due to the \( I_{\alpha} h \rightarrow I_{\alpha} l \) transition, and find that the transition had an onset shortly after the beginning of the warm-up at about 20 K, which is at lower temperature than in our data. The transition was complete at about 70 K, as in our data. On the other hand, the 1000 \( \mu \text{m} \) thick ice layer of Narten et al. (1976) still had a recognizable "interstitial" peak in the rdf after warming to 77 K.

7. MODEL CALCULATIONS

Molecular dynamics (MD) calculations of water structure, that is, solutions of the equations of motion for an ensemble of point masses with the intermolecular potentials of water, go back two decades (for reviews see Angell 1983 and Dore & Teixeira 1991). The resulting structures are best described as an imperfect random network of hydrogen bonds. The models successfully describe many features of the measured structure and physical properties of liquid water and low-density ice \( I_{\alpha} l \).

7.1. Vapor Deposition

MD simulations of water vapor deposition at 10 K on small water clusters (Buch 1992) or on films (Wilson et al. 1995) do not yield \( I_{\alpha} h \). Instead, the resulting network is found to contain a substantial fraction of molecules with hydrogen bond coordination different from 4 (1, 2, 3 and 5) and an abundance of dangling OH bonds. This results in a porous structure with many microvoids and a very large effective surface area. Buch (1992) ascribed this behavior to the preferred attachment of new molecules to dangling atoms on the surface (unbonded H atoms) and to O atoms with only one hydrogen bond, thus propagating the formation of surface protrusions. MD calculations are limited by the amount of relaxation that can be allowed after a molecule has collided with the surface. The ensemble can usually only be followed for tens of picoseconds before allowing the next molecule to hit the surface. In effect, this corresponds to deposition rates of \( 10^8–10^9 \mu \text{m hr}^{-1} \). The
molecular velocities within the cluster are rescaled, after each sticking trajectory, by a constant factor to represent cooling back to the initial temperature of 10 K. However, this quenching may not result in the same final state as a fully relaxed network. In addition, the water molecules are considered rigid and certain energy transfer pathways are not considered in the calculations, including quantum mechanical tunneling and energy transfer to intramolecular O—H stretch vibrations. Note that the energy of binding is of similar magnitude as the energy of an intramolecular O—H stretch vibration or bending mode.

7.2. Pressure Melting

MD simulations have modeled the transition of hexagonal crystalline ice (Ih) into high-density amorphous ice by pressure (Tse & Klein 1987; Tse 1992). The process was found to be driven by a collapse of the open hexagonal ice Ih structure. In the resulting oxygen-oxygen rdf, the second nearest neighbor peak, at 4.5 Å, broadens significantly, and a new band, at 3.2 Å, starts to emerge, reflecting a migration of water molecules into the interstitial sites. A number of hydrogen bonds are broken, creating 3-coordinated and 5-coordinated molecules. Some memory of Ih remains: the network appears to keep a recognizable layering (stacking order) and 6-fold symmetry when viewed along the corresponding crystal axis (Tse 1992).

7.3. Quenching Configurations of Liquid Water

We have attempted to model high-density amorphous ice by quenching configurations, drawn from MD simulations of liquid water in a confined space at 300 K, instead of a crystalline Ih configuration, therefore starting from an amorphous configuration. The system consisted of 343 water molecules placed in a cubic simulation box, assuming periodic boundary conditions and imposing a density of 1.17 g cm$^{-3}$. Instead of a steepest descent minimization of the high-density water configurations, as in our previous model (Wilson et al. 1995), a more gentle quenching was attempted. Each stage of quenching consisted of a 20 ps trajectory during which the temperature was reduced by 50 K, followed by 20 ps of equilibration. Most of the network changes occurred in the higher temperature regime above 150 K. Subsequently, at 77 K, the simulation box was expanded in one dimension, thereby creating a slab of ice with two free surfaces. The structure did not change on a tens of picoseconds timescale when allowed to expand in two dimensions.

To illustrate the disordered nature of these amorphous ices, Figures 5a–5c show a small 10 × 10 × 10 Å cube of the high-density amorphous, low-density amorphous, and crystalline Ih configurations. The planes of molecules, and the cubic and
hexagonal symmetries that are typical of the crystalline polymorphs, cannot be recognized in the amorphous form when viewed from different angles. There are many 4-, 5-, and 7-ringed structures which create a wide range of cavities or "interstitial" sites.

The calculated oxygen-oxygen rdf of the high-density amorphous ice, at 77 K, shows a broad second nearest neighbor oxygen peak, with some indication of splitting into two components (Fig. 6). The peak is not as pronounced as in the collapsed \(I_h\) model of Tse & Klein (1987), but is clearly different from the rdf of the low-density ice model. Also, this peak spans the region between the "interstitial" and second nearest neighbor peak in the observed rdf of \(I_h\) (Fig. 2).

These models of \(I_h\) and \(I_d\) were investigated further by calculating the distribution of coordination numbers and the distribution of cavity sites in the ice, as in Pratt & Pohorille (1992). The coordination number (Table 2) is defined in two ways: either by the number of oxygen atoms within a sphere of 3.2 Å, or by the presence of a hydrogen bond stronger than \(-8\) kJ mol\(^{-1}\). The number distribution of hydrogen bonds shows that (1) there are no interstitial water molecules with zero hydrogen bonds; (2) the most preferred coordination is 4; (3) the distribution of low coordination numbers less than 4 does not differ between the high and low-density amorphous ice models; (4) in high-density amorphous ice, there is a large percentage of molecules with five, six, and even seven oxygen atoms located within 3.2 Å, that are not found in low-density ice; (5) the increased percentage of short oxygen-oxygen distances does not correspond to a large number of 5-coordinated hydrogen bonds, although the percentage of 5-coordinated water is somewhat higher in high-density ice.

Figure 7 shows the probability of placing a rigid sphere of given radius at a random location in the ice structure (allowing for the Van der Waals radius of the water molecules). Large open spaces are conspicuously absent in high-density amorphous ice, relative to low-density amorphous ice. For radii larger than 0.3 Å, there is a fast decline of the relative probability of placing a cavity of given radius at given position. In the model of high-density amorphous ice, cavities larger or equal to the largest cavity in the cubic crystalline lattice are highly improbable.

8. DISCUSSION

8.1. A Model of High-Density Ice

Two models have been proposed for vapor-deposited high-density ice: (1) unbonded water molecules are trapped in interstitial sites in slightly distorted ice I cages, in which 45% of the cavities are filled with interstitial water, or 19% of all water molecules are in cavity positions (Narten et al. 1976); and (2) high-density amorphous ice is a randomized version of ice II or III, high-pressure crystalline polymorphs of water ice which

<table>
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<tr>
<th>TABLE 2</th>
<th>PERCENTILE DISTRIBUTION OF OXYGEN COORDINATION NUMBERS*</th>
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<tr>
<td>coordination number</td>
<td>O-O &lt; 3.2 Å</td>
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<tr>
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<td>hda</td>
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<td>6</td>
<td>7.3</td>
</tr>
<tr>
<td>7</td>
<td>0.06</td>
</tr>
</tbody>
</table>

* Based on the number of oxygen atoms within a radius of 3.2 Å and the number of those oxygen atoms that have a hydrogen bond strength larger than \(-8\) kJ mol\(^{-1}\) (statistical error is \(\pm 2\) in the least significant digit).
have a similar mean density (Narten et al. 1976; Sceats & Rice 1982).

These models are highly unlikely in a distorted network. Some bonding of the interstitial water to other water molecules is expected. The presence of a distorted form of one of the high-pressure crystalline polymorphs of ice is also, in our opinion, not reasonable, because either these polymorphs need a form of proton ordering (preferred orientation of the water molecules) or they are outside their stability regime.

Following Tse & Klein's interpretation of the difference between hda and lda (Tse & Klein 1987; Tse 1992), we propose an alternative model: the high-density amorphous form is a collapsed low-density lattice, characterized by the presence of oxygen-oxygen distances between 3 and 4 Å. Arguments for using the term “collapsed” include that the number density of the large cavities is significantly reduced in high-density amorphous ice compared to the low-density ice. Also, there are more cases of 5 oxygen atoms within 3.2 Å of another oxygen atom, without a similar increase in the number of 5-coordinated hydrogen bonds.

This raises the question of how a collapsed lattice is formed during vapor deposition. Indeed, MD model calculations typically result in the formation of an open I\textunderscore 1 lattice. There is a possibility that ice is deposited in an I\textunderscore 1 form because nature offers a faster redistribution of the energy of binding (some 30 kJ mol\textsuperscript{-1} on average) than the molecular dynamics calculations allow for (e.g., because in the MD model the water molecules are rigid). Alternatively, the ice may deposit itself in an open I\textunderscore 1 form, but a collapse to I\textunderscore h occurs below the binding site upon release of binding energy, analogous to the injection of energy in the ice network, as in charged particle bombardment and photolysis (§4), while the MD calculations, again, do not allow a fast enough dissipation of energy for the distortion to be frozen in.

There is no single structure which characterizes I\textunderscore h. Instead, a wide range of configurations is possible, depending on the number of collapsed cages and the amount of relaxation toward an I\textunderscore 1 structure. Such relaxation apparently causes the less distorted cages to be restructured more easily than the more heavily distorted structures, with water molecules at an “interstitial” position in an ice I\textunderscore 1 lattice. The difference between hda and I\textunderscore h, if any, is that hda may have a more recognizable hexagonal stacking order than I\textunderscore h.

8.2. I\textunderscore h \rightarrow I\textunderscore 1, I\textunderscore 1 \rightarrow I\textunderscore h

The presence of a well-defined transition between one amorphous form and the other suggests that there is an activation energy barrier that prevents the transition at temperatures below 38 K in a vapor deposit, and significantly delays the transition below 70–80 K in electron-irradiated I\textunderscore 1 or I\textunderscore h. The nature of this barrier determines the physical difference between the two amorphous forms.

We speculate that the I\textunderscore h \rightarrow I\textunderscore 1 transformation requires the breaking of one hydrogen bond, on average, per molecule in the I\textunderscore h structure. An argument for this is that the temperature range of 70–90 K that discriminates the two regimes is at half the temperature needed to transform amorphous ice into cubic ice. The latter transition occurs at 143–155 K and has a characteristic activation enthalpy of $\Delta H_a = 44 \pm 2$ kJ mol\textsuperscript{-1} (Jenniskens & Blake 1994), which is equivalent to breaking 2 hydrogen bonds on average per molecule. The onset of the transition will scale with the activation energy, if the kinetic mechanism that drives the transformation is the same. Therefore, the breaking of 1 hydrogen bond on average per molecule, with an activation energy of $\Delta H_a = 22$ kJ mol\textsuperscript{-1}, corresponds to a temperature of about 72–78 K, in agreement with the boundary of the metastability zones of high and low-density amorphous ice (70–90 K).

The reordering from I\textunderscore h to I\textunderscore 1 starts at 38 K, in a gradual warm-up, and proceeds gradually. The onset at 38 K suggests that the activation energy is lowered by some mechanism, and the gradual transition indicates that the effect is of a range of activation energies. This probably reflects the distribution of binding states in the ice. Indeed, MD models of vapor-deposited low-density ice (Buch 1990; Zhang & Buch 1990; Buch 1992) show many defects in this network, causing a range in bond coordination. The model low-density ice has a mean potential energy per molecule of 20.9 kJ mol\textsuperscript{-1}, averaged over time and all molecules in a cluster, but a broad distribution extends from about 7 to 28 kJ mol\textsuperscript{-1} and peaks at 17.5 and 23.5 kJ mol\textsuperscript{-1}. Alternatively, it has been shown that 5-coordination can lower the activation enthalpy by about 10 kJ mol\textsuperscript{-1} (Sciortino et al. 1992). High-density amorphous ice is particularly rich in 5-coordinated water.

Exposure to the electron beam at temperatures below 70 K causes the ice to flow, although the I\textunderscore h structure is preserved. The low viscosity of high-density ice suggests that it can easily restructure its lattice, like liquid water. In liquid water, 5-coordination has been invoked to account for the easy restructuring, because it allows a lowering of the activation enthalpy (Sciortino et al. 1992). Five-coordination is less important in low-density ice, and absent in cubic ice, in agreement with the observed higher viscosity.

The mechanism of electron beam processing of I\textsubscript{c} into I\textsubscript{h} is not yet well understood. Elastic collisions of electrons in the bulk are not important for displacements (Heide 1984) and, hence, for the formation of I\textsubscript{h}. At a given dose, only about $2 \times 10^4$ of all molecules are displaced. Ionization processes, in which a water molecule is dissociated ($\text{H}_2\text{O} \rightarrow \text{H} + \text{OH}$), can be significant. For the given radiation dose, about 30% of all water molecules are dissociated. However, other processes.

![Figure 7](image-url)
may dominate (for example, the electronic or vibrational excitation of the inter- and intramolecular hydrogen bonds). Studies of the influence of photons on this process may reveal new information. Note, for example, that photons that excite a single O-H-O bending or a single O-H stretching vibration (6.27 and 3.07 μm) carry as much energy as a typical hydrogen bond (19 and 39 kJ mol⁻¹, respectively). Visible and near-infrared photons carry more energy than this and may contribute to the structural transformation if absorbed.

8.3. Time-Temperature Diagram

It is reasonable to assume that activation enthalpy is a local phenomenon and activation enthalpies at other sites in the lattice are not much affected by a local restructuring. Given a range of activation enthalpies that determine the onset and the completion of the high- to low-density transition, a time-temperature diagram can be constructed (Fig. 8), according to which the induction time (Δt) of each phase of the transition behaves according to (Schmitt et al. 1989)

$$\Delta t(s) = v_0 \exp \left( \frac{\Delta H_a}{RT} \right).$$

The kinetics of all transitions in water are governed by the intermolecular vibration ($v_0 = 9 \times 10^{-15}$ s) or bending ($v_0 = 2.1 \times 10^{-14}$ s) timescales of water. The uncertainty in the time axis of Figure 8 (the vertical shift) is about a factor of 4, depending on the definition of the onset.

If ΔHₐ is of order 22 kJ mol⁻¹, it follows that high-density amorphous ice can persist on a timescale of 10⁵ yr at temperatures below 22 K, even in the absence of irradiation or charged particle bombardment. Similarly, the high-density ice can persist below temperatures of 19 K on the solar system timescale of 4.5 × 10⁹ yr.

9. ASTROPHYSICAL IMPLICATIONS

9.1. The 3.07 μm Ice Band on Interstellar Dust Grains

Ice in the interstellar medium is usually characterized by the strength and shape of the 3.07 μm absorption band in the infrared (Knacke, Cudaback, & Gaustad 1969; Gillett & Forrest 1973; Merrill, Russell, & Soifer 1976; Hagen et al. 1981; Whitsett et al. 1983; Bult, Greenberg, & Whitsett 1985; Baratta & Stazzulla 1990; Smith, Sellgren, & Brooke 1993). This band is due to the O—H stretch vibration of water. A unique assignment of shoulder and peaks of bands to vibrations is not possible because intra- and intermolecular coupling of O—H oscillators are nearly equal in strength (Haas & Hornig 1960; Bergren et al. 1978; McGraw et al. 1978). Incoherent inelastic neutron scattering gives a density of vibrational states that shows a strong difference between pressure-induced high- and low-density amorphous ices (hda and lda) (Klug et al. 1991; Tse 1992). The strongest differences are in the longitudinal and transverse optic branches of the dispersion relation. The leading edge of the librational band begins at lower frequency in hda ice and rises more slowly. These differences are manifested in the band shapes of infrared absorption bands.

There is a clear difference in the 3.07 μm band shape associated with vapor-deposited high- and low-density amorphous ice, Iₐ, h, and Iₐ, l. If the laboratory spectra of low-temperature deposits by Hagen et al. (1981) are indeed those of Iₐ, h, as we believe they are, it follows that the Iₐ, h band contains additional absorption at high vibrational frequency. When the ice is warmed at a rate of less than 5 K minutes⁻¹, the peak wavelength changes from 3.04 to 3.08 μm, and the feature sharpens from 310 to 230 cm⁻¹ in the temperature range 20–70 K. The 45 μm band is a less sensitive probe of the Iₐ, h → Iₐ, l transition (Klug et al. 1991; Moore & Hudson 1992).

Smith, Sellgren, & Tokunaga (1989) found that some astrophysical objects show ice-band profiles that are best fitted by a combination of ice deposited at low temperature and warmed to 23, 77, and 150 K. We now interpret these deposits as high-density amorphous ice Iₐ, h, low-density amorphous ice Iₐ, l, and cubic crystalline ice Iₐ, c, respectively. For example, spectra from the Becklin-Neugebauer object are best fitted with spectral ratios of Iₐ, h: Iₐ, l: Iₐ, c equal to 3.1:1.2:1, while Mon/IRS 3 is best fitted by ratios of 5.3:2.5:1 and GL 2591 by ratios of 8.6:1.2:1. The 23 K ice band provides a satisfactory fit to the overall shape of most other known spectra. The conclusion must be that ice frozen on grains in the interstellar medium is principally in the high-density amorphous form.

The frost on interstellar grains constitutes the largest reservoir of ice in the universe—in our Galaxy alone about 10²⁷ kg (Jenniskens et al. 1993). Little is known about extrasolar planetary systems, hence the amount of ice in icy planets and comets cannot be estimated with any certainty. However, if all stars have an amount of icy material in the form of icy planets, satellites and Kuiper Belt comets similar to that in our planetary system, then the total adds to perhaps 10¹⁷–10²⁴ kg. We cannot think of other sources of ice. Hence, we conclude that the frost on interstellar grains is the largest reservoir of ice, and most ice in the universe is, therefore, in the high-density amorphous form.

9.2. Formation of Icy Mantles

The physical conditions present in molecular clouds are consistent with the presence of Iₐ, h instead of Iₐ, l. The deposition
The band positions of CO and CO₂ have been studied in the formation of the water mantle at low temperature. Also expected to be distorted into the I, h form during initial the deposition of energy by electrons or protons, then the ice is energy of O--H chemical bonds has an effect similar to that of molecules from setting into a crystalline structure. But does this temperature spike decreases rapidly with increasing distance from surface. Kouchi et al. (1994) argue that the resulting temperature below 30 K. Hence, evaporation and recondensation of icy mantles in dense cloud environments results in I₉h ice mantles, as observed.

Initially, however, the ice mantle is not formed by direct vapor deposition but by surface reactions of atomic oxygen and hydrogen: O + H → OH and OH + H → H₂O. In some environments, this initial ice layer may be preserved. One wonders if that ice would be structurally different from vapor-deposited amorphous ice. The energy released in these surface reactions is on the order of 1 eV, which will locally heat the surface. Kouchi et al. (1994) argue that the resulting temperature spike decreases rapidly with increasing distance from the formation point, sufficiently fast to disallow the water molecules from setting into a crystalline structure. But does this prevent the formation of I₉h instead of I₉h? It probably does. We point to the analogy with electron bombardment, where the input of energy in the ice matrix at T < 70 K results in high-density amorphous ice. Hence, if the release of the binding energy of O-H chemical bonds has an effect similar to that of the deposition of energy by electrons or protons, then the ice is also expected to be distorted into the I₉h form during initial formation of the water mantle at low temperature.

9.3. CO as a Tracer of Structural Changes in the Water Matrix

Impurities in the water ice matrix can be used as probes of the structure of the water mantle ( Grim & Greenberg 1987). The band positions of CO and CO₂ have been studied in the laboratory (Sandford et al. 1988; Schmitt et al. 1989; Sandford & Allamandola 1990; Palumbo & Strazzulla 1993). Both bands are readily observable in astrophysical ices. Notably the 4.7 μm band of CO-O has been studied using ground-based infrared telescopes (Whittet et al. 1983; Latty et al. 1984; Kerr, Adamson, & Whittet 1993; Chiar et al. 1994).

Two absorption maxima were identified in laboratory studies of CO: H₂O 1:20 mixtures at 10 K (Sandford & Allamandola 1988). The main absorption is at 2137 cm⁻¹, which is shifted to lower frequency (i.e., a repulsive bond) with respect to the position of pure CO at 2143 cm⁻¹. In addition, there is a shoulder at 2152 cm⁻¹ (an attractive bond) in fresh deposits at T < 40 K, which accounts for 22%-26% of the total band intensity below 35 K. The latter band progressively disappears (irreversibly) when the ice is warmed above 40 K. The CO loss stops immediately when warm-up is stopped (Sandford & Allamandola 1988), which implies that the loss of CO is controlled by changes in the water structure and that these structural changes are gradual. Indeed, the temperature zone where this band disappears (40–60 K at 2 K minutes⁻¹ warm-up) coincides with our measured transition I₉h → I₉l.

The peak at 2152 cm⁻¹ was assigned to interstitial CO by Sandford & Allamandola (1988), and the peak at 2137 cm⁻¹ was thought to be due to CO that substitutes for a water molecule in the water-ice network. It is unlikely that CO can substitute for water in a water matrix, particularly at temperatures above 60 K, and that upon warming from low temperatures, interstitial CO evolves into a substitutional site. Schmitt et al. (1989) and Grim (1988) assigned the two peaks to two types of water-CO interactions within the incompletely hydrogen-bonded network. While the 2137 cm⁻¹ peak was thought to be due to unbonded CO, the 2152 cm⁻¹ peak was assigned to CO-H₂O hydrogen-bonded complexes in a water ice. This complex has a peak at 2149 cm⁻¹ in an argon matrix (Dubost 1976) and 2146.5 cm⁻¹ in a CO ice (Hagen & Tielens 1981). During warm-up from 10 to 40 K, the hydrogen-bonded complexes were thought to be preferentially replaced by hydrogen bonds of the normal water network. We believe this interpretation to be essentially correct. In light of our observation that the 2152 cm⁻¹ peak is linked to I₉h ice, we propose that this peak is due to CO-H₂O complexes involving dangling OH bonds, while the 2137 cm⁻¹ peak is due to CO trapped in "interstitial" sites in the ice cages, noting that such cages in I₉h have a broader range of shapes and are typically smaller than those in I₉l. This can account for the small shift to lower frequency and the slight narrowing of the C=O stretch vibration during the I₉h → I₉l transition (Sandford & Allamandola 1988).

The CO peak at 2152 cm⁻¹ is absent in interstellar ices. This implies that the CO-containing ice mantles were warmed above 60 K on timescales less than that for water mantle evaporation. Indeed, most of the CO is found in more volatile apolar mantles. However, the water ice itself is not found in the I₉l form. This observation suggests that I₉l is transformed back into I₉h on short timescales. The most likely transformation mechanism is photolysis. The photon flux in the dense cloud medium is high enough to allow a restructuring into the I₉h form on a timescale of less than 10³ yr, when 1%-10% of water molecules absorb a 10 eV photon. Many more lower energy photons are absorbed. This cycling of I₉h → I₉l ice and back can play an important role in the generation of organic compounds of large molecular weight through UV photolysis, as long as traces of CO or CO₂ remain in the ice (Jenniskens et al. 1993; Jenniskens, Tielens, & Blake 1994). The photolysis will also cause the ice to be less viscous and form a smooth mantle around the ice grains. Hence, the effective surface area for H₂ formation is probably not much different from that of a homogeneous surface.

10. CONCLUSIONS

The observed changes of the diffraction pattern of vapor-deposited water ice as a function of temperature, the presence of two stability regimes for electron bombardment, and the similar rdf compared to the pressure-induced high- and low-density amorphous forms, among other arguments, leave no doubt as to the existence of high-density amorphous ice in low-temperature vapor deposits. It is a naturally occurring form which arises when water ice is slowly (< 100 μm hr⁻¹) vapor-deposited on any substrate of good thermal conductivity at temperatures T < 30 K.

The radial distribution function of high-density amorphous ice, obtained by Fourier transformation of the diffraction patterns, has oxygen-oxygen distances between 3 and 4 Å, between the first and second nearest neighbor distance in low-density amorphous ice. The shape of the rdf is similar to that of pressure-induced hda reported by Bizid et al. (1987).

MD calculations can reproduce this 3–4 Å feature, if the water ensemble is allowed to relax in a confined space. The
feature is not due to unbonded interstitial water molecules. Four-coordination is still preferred, although in comparison to the low-density amorphous ice model, there is a somewhat larger number of five-coordinated water molecules. The most striking feature of the high-density amorphous structure is the small probability of finding cavities as large as those found in the two crystalline forms of ice, while low-density amorphous ice has a cavity size distribution more characteristic of an open crystalline structure (Fig. 7). From these results, we conclude that the high-density amorphous form is best described as a (partially) collapsed low-density ice structure.

There is an irreversible first-order transition from high- to low-density amorphous ice. Below 70–80 K, there is a range of activation energy barriers that hinder restructuring. Several processes can transform the low-density amorphous form (or the crystalline cubic polymorph) back into high-density amorphous ice in this regime. These include electron and proton bombardment, pressure melting, and possibly UV photolysis. From similarities between the resulting radial distribution functions, and from results of MD simulations of pressure-induced amorphous ice in the literature, we infer that all of these processes cause some form of collapse of the low-density amorphous ice structure. During vapor deposition, the collapse of the water lattice may be a consequence of a rapid dissipation of the energy of binding before impinging molecules can reorient to form low-density amorphous ice. Alternatively, the ice may form a low-density amorphous ice structure initially, but then be distorted by the release of binding energy (analogous to distortions induced by electron irradiation).

The nature of the energy barrier determines the physical difference between the two amorphous forms. We propose that the restructuring of high-density ice into low-density ice requires the breaking of one hydrogen bond, on average, per molecule. The wide range of activation energies present at different locations in the vapor-deposited high-density amorphous ice may be responsible for the early onset. Alternatively, high-density ice does have a relatively large fraction of five-coordinated water molecules, and there has been a suggestion that five-coordination is responsible for the lowering of the activation enthalpy needed for the rapid restructuring in liquid water.

Given the astrophysical conditions in the dense interstellar medium, we argue that vapor deposition on interstellar grains results in a water frost with high-density amorphous structure. We surmise that water mantles formed through the recombinination of H and OH radicals result also in a high-density ice structure. A brief warming above 40 K, however, will lead to a low-density amorphous form in the absence of sufficient irradiation processes. The structural transition from one amorphous form into another can play an important role in dynamic processes like radical diffusion and outgassing.

Ground-based observations of the O–H stretch vibration at 3.07 μm in a variety of astrophysical environments have been interpreted in terms of a composite of ice deposited at low temperature and warmed to 23, 77, and 150 K. We now interpret these deposits as high-density amorphous ice (Iₜₜ), low-density amorphous ice (Iₜₜ), and cubic crystalline ice (Iₜ), respectively. Most of the observed 3.07 μm absorption bands in dense molecular clouds are best represented by a low-T(<30 K) deposit (Hagen et al. 1981) and, therefore, by the high-density form of amorphous water ice. The frost on interstellar grains constitutes the largest reservoir of ice in the universe—in our Galaxy alone, an estimated 10²⁷ kg—and we therefore conclude that most ice in the universe is in this high-density amorphous form.

In the near future, the Infrared Space Observatory will offer a unique means to measure the structural state of water ice in different astrophysical environments, either directly, from the shape of the infrared absorption and emission bands of the vibration and bending modes of water, or indirectly, from the behavior of trapped impurities in the water matrix.

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