Is High-Density Amorphous Ice Simply a “Derailed” State along the Ice I to Ice IV Pathway?

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ABSTRACT: The structural nature of high-density amorphous ice (HDA), which forms through low-temperature pressure-induced amorphization of the “ordinary” ice I, is heavily debated. Clarifying this question is important for understanding not only the complex condensed states of H2O but also in the wider context of pressure-induced amorphization processes, which are encountered across the entire materials spectrum. We first show that ammonium fluoride (NH4F), which has a similar hydrogen-bonded network to ice I, also undergoes a pressure collapse upon compression at 77 K. However, the product material is not amorphous but NH4F II, a high-pressure phase isostuctural with ice IV. This collapse can be rationalized in terms of a highly effective mechanism. In the case of ice I, the orientational disorder of the water molecules leads to a deviation from this mechanism, and we therefore classify HDA as a “derailed” state along the ice I to ice IV pathway.

A simple compression experiment of the “ordinary” ice Ih at liquid-nitrogen temperature has left a remarkable legacy and tremendous impact across the materials sciences.1 It marked the discovery of high-density amorphous ice (HDA), which provided the first example for the process of pressure-induced amorphization (PIA). Countless other examples of PIA have been reported since for a wide range of inorganic, organic, and even biochemical materials.2 However, questions related to the detailed mechanisms of the PIA processes are still under debate, in particular because some reported cases of PIA were later attributed to either the formation of nanocrystals, chemical decomposition or the nonhydrostatic compression conditions.2,3

The PIA of ice Ih was first rationalized in terms of a thermodynamic melting process implying that HDA is a glassy state of high-pressure water.4,8−12 This, together with the transformation of HDA to low-density amorphous ice upon decompression or heating at ambient pressure,16 provided an experimental background for the two-liquid model of water and the second critical point scenario.13 Yet, later, it was also argued that HDA forms due to a mechanical instability and can therefore be understood as a collapsed “ill-crystalline” material16−23 that lacks a connection with the liquid phase.24,25 A temperature-driven crossover between these two scenarios has also been suggested.9,17,20 Finally, it was even speculated that HDA may consist of strained microdomains of crystalline high-pressure phases of ice.26,27 In summary, it is safe to state that the structural and thermodynamic nature of HDA is still poorly understood. Due to the pivotal position of HDA not only for understanding the condensed states of H2O but also in the wider context of PIA processes, a deeper knowledge of the mechanism of the PIA of ice I is of paramount importance.

Here we first investigate the low-temperature compression behavior of hexagonal and stacking-disordered ammonium fluoride (NH4F), whose hydrogen-bonded networks are similar compared to those of the corresponding members of the ice I family. From this, we then aim to gain new insights into the origin of the pressure collapse of ice upon low-temperature compression and to propose a structural mechanism for the PIA of ice I, which we also test with density functional theory (DFT) calculations.

Ice I and NH4F I are similar materials from a structural point of view. Figure 1a shows the isostructural hydrogen-bonded networks of the hexagonal ice Ih and NH4F Ih materials, which are the stable phases at ambient conditions. The hydrogen bonds in NH4F Ih are less than 2% shorter than those in ice Ih.28 A metastable variant of ice Ih exists that contains interlaced sequences of cubic and hexagonal stacking, generally known as stacking-disordered ice (ice Isd).29−32 The most cubic ice Isd prepared so far, which will be used in the following, was obtained by slowly heating the ice II high-pressure phase at ambient pressure.32 The material obtained by heating the NH4F II high-pressure phase at ambient pressure has been labeled as NH4F V.33 Yet, as we will show, this material is, in
analogy with the situation for ice, best described as stacking-disordered ammonium fluoride I (NH₄F I).

Remarkably, as shown in Figure 1b, NH₄F Ih and NH₄F Isd show similar collapses upon compression at 77 K in terms of the onset pressures and overall volume changes compared to the ice I samples. The X-ray diffraction patterns of the starting materials shown in Figure 1c revealed that they actually consist of crystalline mixtures of ∼90% NH₄F II and ∼10% NH₄F I. Upon comparison of the crystal structure of NH₄F II with the known phases of ice, it was realized that NH₄F II is isostructural with the metastable ice IV. Although NH₄F II is the first high-pressure phase to form upon compression of NH₄F I at room temperature, it is probably not the stable phase at 77 K and ∼1 GPa due to the

Figure 1. Ice I and NH₄F I show remarkably similar pressure collapses upon compression at 77 K. (a) Crystal structures of ice Ih and NH₄F I. (b) Changes in sample volume, as indicated by the piston extension, upon compression and decompression at 77 K of equal amounts of ice I and NH₄F I materials \( n(\text{H}_2\text{O}) = n(\text{NH}_4^+) + n(F^-) \). Powder X-ray diffraction patterns of the materials (c) before and (d) after compression. The percentages of cubic stacking, \( \Phi_c \), of the starting materials are given in (c), and the tickmarks in (d) indicate the positions of the Bragg peaks of NH₄F II and Ih, respectively. Thick gray lines in (c) and (d) are Rietveld or MCDIFFaX fits.

Figure 2. Formal mechanism of the EKC.36,37 (a) Transformation of 2 × 2 × 3 supercells of the ice/NH₄F Ih unit cells viewed along [100] to the ice IV/NH₄F II unit cell in the hexagonal setting. The spheres indicate the nodes of the hydrogen-bonded networks and are colored differently for each layer. The “threading-through” structural feature is shown in the inset. (b) Transformation of a single layer of the Ih/Ic networks to the ice IV network viewed along [001].

On the basis of the similar pressure collapses, we initially assumed that the NH₄F samples had also undergone transitions to high-density amorphous materials in analogy to what has been observed for ice. Surprisingly, the diffraction patterns of the NH₄F materials after compression shown in Figure 1d revealed that they actually consist of crystalline mixtures of ∼90% NH₄F II and ∼10% NH₄F I. Upon comparison of the crystal structure of NH₄F II with the known phases of ice, it was realized that NH₄F II is isostructural with the metastable ice IV. Although NH₄F II is the first high-pressure phase to form upon compression of NH₄F Ih at room temperature, it is probably not the stable phase at 77 K and ∼1 GPa due to the
pronounced slope of the NH$_4$F II/III phase boundary. This suggests that the NH$_4$F II, which results from the low-temperature compression, is a kinetic product that forms as a consequence of a favorable mechanistic pathway.

The transformation of NH$_4$F I to NH$_4$F II can be understood on the basis of a remarkable mechanism that achieves a 37% increase in density while only breaking one in four hydrogen bonds. This collapse was first described by Engelhardt and Kamb for the transition from the hypothetical fully cubic ice I (ice Ic) to ice IV. The corresponding collapse starting from the hexagonal starting material was later described in the NH$_4$F literature. The mechanistic details of this collapse, which we refer to as the Engelhardt–Kamb collapse (EKC), are shown in Figure 2. The ice Ic/Ih networks contain identical layers consisting of puckered six-membered rings in the armchair conformation. These networks are shown in Figure 2a with the network nodes of the individual layers highlighted in different colors.

The transition from the ice Ic to the ice IV network requires breaking of the interlayer hydrogen bonds followed by flattening and ultimately a complete “rebuckling” of the layers. The last step is the formation of hydrogen bonds right through the center of the six-membered rings of the layers above and below toward water molecules of the second layers above and below (see Movie S1).

This threading-through is the hallmark structural feature of the ice IV network, which “pulls” the layers closer together and thereby achieves the increase in density (inset in Figure 2a). Starting off from the ice Ih network, the “rebuckling” of layers is not required. Instead, the individual layers need to shift or rotate with respect to one another to achieve the threading-through of the six-membered rings (see Movie S2). Because both the ice Ic and Ih networks can undergo the EKC, it is of course also possible to start from stacking-disordered starting materials.

The effect of the ice IV network formation on an individual layer is shown in Figure 2b. The six-membered rings labeled with 1 are the ones that experience threading-through with hydrogen bonds during the transition to the ice IV network. This leads to a flattening of these rings, a slight increase in diameter, and rotation. These changes in the 1 rings distort the rings labeled with 2, which either raise or lower the nodes that are not members of the 1 rings so that they can form hydrogen bonds with two layers above or below (see Movie S3).

A remarkable feature of the EKC is that the density increase is highly anisotropic and achieved almost exclusively by contraction along [001]. It is difficult to imagine that there is another mechanism that achieves a similar increase in density while breaking fewer hydrogen bonds. Therefore, the EKC seems to be a particularly important mechanism for densification under kinetically controlled low-temperature conditions.

The similar onset pressures and volume changes of the low-temperature collapses of ice I and NH$_4$F I suggest that both materials suffer from a similar mechanical instability that marks the onset of the EKC. However, in both cases, the EKC is not followed through entirely at 77 K. In the case of NH$_4$F I, the conversion to NH$_4$F II is ~90 wt% leaving ~10 wt% of NH$_4$F I unconverted. The incomplete conversion is attributed to the buildup of macroscopic strain environments, which arise from the highly anisotropic nature of the EKC.

In the case of ice, it is important to recall that the ice I materials are hydrogen-disordered, which means that they display disorder with respect to the orientations of the hydrogen-bonded molecules (Figure 1a). As mentioned earlier, during the EKC, the interlayer hydrogen bonds are broken and re-formed with water molecules one and two layers above or below. The chances of successfully connecting a broken hydrogen bond to another water molecule are 50% because water molecules can be either hydrogen-bond donors or acceptors. This implies that in the case of ice I, not all broken hydrogen bonds will necessarily thread-through the six-membered rings in the final stage of the EKC but form hydrogen bonds with other close water molecules instead that fulfill the donor/acceptor conditions. The resulting material is HDA, which displays a lack of long-range order and a density somewhat lower than that of ice IV.

Accordingly, HDA can be classified as a "derailed" state along the EKC, and we provide further arguments for this in the following.

The hydrogen-bond donor/acceptor mismatches in ice could in principle be resolved with molecular reorientations. These are known to be very slow at 77 K but have been shown to "unfreeze" during the glass transition of HDA at higher temperatures. When HDA is heated at around its “natural” pressure of 1 GPa, the amorphous sample becomes denser at first followed by crystallization. The crystallization to ice IV has been shown to have the lowest activation energy out of all of the possible crystallization pathways around 1 GPa. This means that the derailed state of HDA can be brought "back on track" with respect to the EKC by thermal annealing under pressure as dynamic molecular reorientations resolve the problem of the hydrogen-bond donor/acceptor mismatches. Similar structures of pressure-annealed HDA and ice IV are also evident from their Raman spectra. Furthermore, the first strong diffraction peak of pressure-annealed HDA, which reflects the intermediate range structural order, is in the same position as the strongest Bragg peaks of ice IV. Finally, it has been suggested that HDA still contains “crystalline remnants” of ice I after the PIA. These can now be rationalized in terms of small amounts of ice I-like nanodomains, which arise stochastically wherever the largest numbers of hydrogen-bond donor/acceptor mismatches are present.

An interesting question now arises if the derailment upon low-temperature compression of ice I could be prevented by using hydrogen-ordered starting materials. The hydrogen-ordered ice Ic (ice Xlc) is a promising starting material because the EKC from a cubic starting material does not require the translational movement of layers. All interlayer hydrogen bonds point in the same direction in ice XI, which means that the problem of the hydrogen-bond donor/acceptor mismatches does not apply. Unfortunately, ice Ic has not been prepared so far, and consequently, it is unknown if and how ice Xlc can be prepared. Also, the fact that ice Ih becomes only partially hydrogen-ordered upon doping with potassium hydroxide below 77 K illustrates how much of a challenge the low-temperature compression of ice Xlc would be.

To investigate the feasibility of the ice Xlc to ice IV pathway, we performed a transition-state search using the solid-state nudged elastic band (NEB) method at the dispersion-corrected DFT level. The solid-state NEB method couples the atomic and lattice degrees of freedom of a crystal structure and enables the transition pathway between two defined states to be optimized. A minimum-energy contiguous path was obtained after optimization, and the energy profile and cell volumes of all frames on the transition path are shown in Figure 1.
3. The activation energy for the phase transition is found to be \( \sim 8 \text{ kJ mol}^{-1} \), about three times as large as the energy difference between ice IV and XIc. The initial configurations are associated with the formation of point defects that resemble interstitials. The volume changed slowly during the first stages of the transition, but close to the transition state, there was a marked drop. The largest change in volume at the very end of the transition is clearly associated with the formation of hydrogen bonds passing through the six-membered rings (Figure 2). We note that our estimated activation energy is about four times higher than what is expected for a phase transition induced solely by the thermal motion of the molecules at 77 K (i.e., at 77 K, \( 3RT = 1.9 \text{ kJ mol}^{-1} \)). Several factors could contribute to this overestimation. First, we started from a perfect lattice of ice XIc. In ice, non-negligible concentrations of kinetically trapped point and line defects will be present, which, together with grain-boundary effects, may act as seeds for the transition to ice IV, effectively lowering the transition barrier. Second, our predicted transition mechanism only represents one of the possible transition pathways; lower-energy transition mechanisms may exist for larger simulation cells, where the point defects, which facilitate the transition, are more spatially separated, allowing for greater lattice relaxation. Finally, quantum tunneling effects, which facilitate in particular the movements related to the hydrogen atoms, are not captured in our calculations. Therefore, in summary, it seems very plausible that ice XIc would undergo the EKC upon low-temperature compression and avoid derailment to HDA.

In conclusion, we have shown that the hydrogen-bonded networks of ice I and NH₄F I both suffer from similar mechanical instabilities upon low-temperature compression, which mark the onsets of EKCs in these materials. Yet, neither material follows the EKC through completely. This is most likely due to the buildup of macroscopic strain in NH₄F and the hydrogen-bond donor/acceptor mismatches in ice, as illustrated by the DFT calculations of the ice XIc to ice IV pathway. Put simply, the low-temperature PIA of ice I is a consequence of its hydrogen disorder. This means that HDA does not represent the glassy high-pressure liquid, at least not the material that is obtained from the low-temperature compression of ice I at 77 K. Furthermore, this study also showcases the high efficiency of the EKC in achieving large density increases under kinetically controlled conditions. Future work should focus on low-temperature compressions of other structurally related materials such as diamond, silicon, and the large family of AB materials including silver iodide, zinc sulfide, gallium phosphide, and silicon carbide.

### EXPERIMENTAL AND COMPUTATIONAL METHODS

NH₄F I (99.99%) was purchased from Sigma-Aldrich and stored under dry nitrogen. For the compression experiments at 77 K, 404.0 mg of NH₄F I was placed inside an indium cup, which was quickly transferred into a stainless steel piston cylinder precooled with liquid nitrogen. The ice I samples were prepared by pipetting 393.0 mg of Milli-Q water into an indium cup inside a precooled piston cylinder. The compression to 1.4 GPa at 77 K was performed with a 30 ton hydraulic press. Changes in sample volume were recorded using a GTS000RA-L25 positional transducer from RDP electronics. To obtain NH₄F Iod, NH₄F II was prepared in a first step by compressing 404.0 mg of NH₄F I to 0.25 GPa at room temperature, followed by cooling to 77 K under pressure and heating from 77 to 203 K at ambient pressure. Ice Iod was obtained by heating an ice II sample from 77 to 190 K at ambient pressure. Both stacking-disordered materials were subjected to the same compression experiments at 77 K as previously described for the hexagonal materials. After the compression experiments, the samples were recovered under liquid nitrogen at ambient pressure and characterized with X-ray powder diffraction using a custom-made sample holder with Kapton windows mounted on a Stoe Stadi-P diffractometer (Ge 111 monochromatized Cu Kα, 40 kV, 30 mA). Data were collected using a Mythen 1K linear detector, and the temperature of the sample was controlled with an Oxford Instruments CryojetHT.

The solid-state NEB calculations were performed using the atomic simulation environment and the TSASE code. Thirteen frames of the \( 2 \times 2 \times 3 \) supercells from ice XIc to ice IV were considered on the transition path. The transition-state search was considered converged if the maximum atomic force fell below \( 2.5 \times 10^{-3} \) Hartree/Bohr. The energies, forces, and stress tensors were calculated using the CP2K code, which uses a mixed Gaussian/plane-wave basis set and a 800 Ry polarization quality Gaussian basis set. We employed double-\( \zeta \) polarization quality Gaussian basis sets and a 800 Ry plane-wave cutoff for the auxiliary grid, in conjunction with the Goedecker–Teter–Hutter pseudopotentials. All calculations were performed using the nonlocal van der Waals density functional, vdW-DF2, which has been shown to give excellent agreement with experiment and diffusion Monte Carlo calculations on the sublimation energies of ice I and ice VIII.

### ASSOCIATED CONTENT

#### Web-Enhanced Features

Movie S1 (QuickTime movie) shows the transformation of the ice Ic to the ice IV network viewed along [100]. Movie S2 (QuickTime movie) shows the corresponding transformation of the ice Ih network. Movie S3 (QuickTime movie) shows a view along [001] of the transformation of a single layer of the ice Ic/Ih networks during the transformation to the ice IV network. All movies were prepared with the CrystalMaker software by using linear interpolations between the start and end structures. The movies therefore show the formal mechanism of the EKC in line with Figure 2.
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