Anomalous Behavior of the Homogeneous Ice Nucleation Rate in “No-Man’s Land”

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Supporting Information

ABSTRACT: We present an analysis of ice nucleation kinetics from near-ambient pressure water as temperature decreases below the homogeneous limit \( T_H \) by cooling micrometer-sized droplets (microdroplets) evaporatively at \( 10^3 \)−\( 10^4 \) K/s and probing the structure ultrafast using femtosecond pulses from the Linac Coherent Light Source (LCLS) free-electron X-ray laser. Below 232 K, we observed a slower nucleation rate increase with decreasing temperature than anticipated from previous measurements, which we suggest is due to the rapid decrease in water’s diffusivity. This is consistent with earlier findings that microdroplets do not crystallize at <227 K, but vitrify at cooling rates of \( 10^6 \)–\( 10^7 \) K/s. We also hypothesize that the slower increase in the nucleation rate is connected with the proposed “fragile-to-strong” transition anomaly in water.

Understanding the phase transition from supercooled water or amorphous ice to crystalline ice is key to various fields ranging from cryobiology to atmospheric and astrophysical sciences.\(^1\)−\(^5\) Accurate determination of ice nucleation kinetics and the involved structural transformation is essential for modeling atmospheric cloud formation and the corresponding thermostating effects on the world climate. Most experimental studies of homogeneous ice nucleation (where ice here refers to crystalline ice unless noted otherwise) have been limited to temperatures above the homogeneous nucleation temperature \( (T_H) \) of \( \sim 232 \) K\(^6\)−\(^11\) and below the homogeneous crystallization temperature \( (T_X) \) of \( \sim 160 \) K.\(^12\) This limitation is due to rapid crystallization upon deep supercooling which also prevents measurements of bulk water properties using standard techniques. In the context of ice nucleation, \( T_H \) and \( T_X \) correspond, respectively, to the median temperature limit of metastability of micrometer-sized liquid droplets at moderate

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cooling rates (from ~1 K/min to ~100 K/s) and to where amorphous ice crystallizes upon heating. Thus, the temperature region below $T_{h}$ and above $T_{x}$ is referred to as "no-man’s land". As water is supercooled toward "no-man’s land", the decrease in diffusivity ($D$) becomes more pronounced which may have a significant effect on the ice nucleation kinetics. There have been several simulations that observe in detail the formation of ice nuclei. However, the lack of direct measurements of the nucleation rate, the diffusivity and the solid—liquid (where solid refers to crystalline ice) water interfacial free energy ($\sigma_{li}$) limits our knowledge on ice nucleation kinetics within this important region of the water phase diagram.

There have been several attempts to circumvent the limitation set by $T_{h}$ to achieve deeper supercooling and observe homogeneous ice nucleation events in "no-man’s land". This is often done by increasing the sample cooling rate and reducing the sample dimensions to nanometer sizes that effectively decreases the nucleation probability. However, reducing the sample volume to such small dimensions also dramatically increases the surface-to-volume ratio and the sample internal pressure due to the reduced radius of curvature (described by the Young—Laplace equation). This may alter $D$, and thereby the bulk homogeneous ice nucleation rate ($f$), significantly such that it is no longer representative of bulk water at ambient pressure. For microdroplets, attempts to scale the nucleation rate with the surface area instead of volume improves the general agreement between different data sets, but surface nucleation has not been observed directly in such experiments. When applied to nanodroplets, the 2.5 nm subsurface layer used to analyze surface nucleation effects is about the size of the nanodroplets themselves, which makes the definition of subsurface volume ambiguous in nanodroplets. Therefore, using nanodroplets in ice nucleation experiments may represent water at significantly different conditions than in microdroplets that better represent bulk water at ambient pressure.

Within the framework of classical nucleation theory (CNT), the formation of an ice nucleus in metastable supercooled water requires overcoming a thermodynamic energy barrier ($\Delta F^*$) that includes the Gibbs free energy change to form bulk ice and the Gibbs free energy increase associated with making an interface. This $\Delta F^*$ is strongly dependent on $\sigma_{li}$ and to some extent on the increase in pressure via the change in chemical potential difference between solid and liquid phases ($\Delta \mu_J$) (see Supporting Information). In addition, formation of a nucleus also requires overcoming the kinetic barrier ($\Delta f^*$) that accounts for the movement of water molecules across the solid—liquid water interface, which can be estimated from $D$ in liquid water. As mentioned previously, pressure increase may also significantly alter liquid water $D$, and thus change the $\Delta f^*$ considerably (see Supporting Information). Also interestingly, $D$ of water at ambient pressure has a peculiar behavior as the temperature approaches $T_{h}$. Below the melting point, $D$ decreases more rapidly with decreasing temperature than what is expected from the Arrhenius equation, and this behavior is categorized as that of a "fragile" liquid. Based on the "fragile" liquid concept, $D$ is expected to decrease even more strongly as the temperature decreases toward and below $T_{h}$, but this has not been experimentally verified. Near the glass transition temperature ($T_g$) of ~136 K, liquid water has been proposed to behave like a "strong" or even "superstrong" liquid, which would mean that the temperature dependence of $D$ follows (in contrast to a "fragile" liquid) the Arrhenius equation with an energy barrier higher than that of a higher-temperature "fragile" liquid. Within "no-man’s land", $D$ has been proposed to approach an Arrhenius-like temperature dependence as the temperature is decreased, thus water becomes a "strong" liquid, with a crossover temperature $T_C$ at ~228 K. Overall, the ice nucleation rate is expected to increase as temperature decreases and approaches $T_{h}$, reach a maximum below $T_{h}$, and then decrease with further temperature decrease toward $T_{x}$. However, in light of the possible large variation of $D$ at ambient pressure and the corresponding effect on $\Delta f^*$ over a wide temperature range, particularly near $T_{h}$, it is less certain how fast the nucleation rate changes as it approaches and crosses $T_{h}$.

We recently developed a new method, based on evaporative cooling and ultrafast probing, that allows studies of supercooled water and ice formation at temperatures where ice is formed very rapidly. Here, we present an analysis of the ice nucleation kinetics of near-ambient pressure water as it approaches and enters into "no-man’s land". We are also adding additional information obtained from hyperquenching experiments of micrometer-sized water droplets in order to place upper and lower limits on the maximum nucleation rate at temperatures further into "no-man’s land". These limits can be defined based on the observation that essentially all droplets crystallize in huge ensembles of droplets of 3 μm in diameter, when they are cooled at 10^4 K/s, whereas crystallization was not detected when cooled at 10^7 K/s through the 70 K broad "no-man’s land".

The experiments were conducted using a gas dynamic virtual nozzle (GDVN) to generate, in vacuum, a train of microdroplets with uniform diameter of 9 or 12 μm that are subsequently probed by X-rays. The droplets cool evaporatively, decreasing their temperature as they travel farther from the nozzle and spend longer time in vacuum. This results in cooling rates of 10^3–10^4 K/s around $T_{h}$ prior to crystallization. Hard X-ray laser pulses, each ~50 fs in duration, from LCLS were used to probe the structure of individual droplets at various distances from the nozzle exit, hence varying the droplet temperature. The scattering patterns from individual droplets hit by the X-ray pulse were recorded over an estimated temperature range of 227–252 K (Figure 1) and sorted according to whether they consisted of only diffuse rings indicating scattering from pure liquid water or contained intense and discrete Bragg reflections that indicate diffraction from ice crystals. We collected at least 1800 individual scattering patterns at each distance or temperature. As the droplets spend more time in vacuum and evaporatively cool, the fraction of ice-containing shots ($f_{isc}$) remains near zero for about 1.2 and 3 ms travel time for the 9 and 12 μm droplets, respectively (Figure 1). This fraction increases rapidly to 0.2 in the next 1 ms for both droplet sizes and to 0.97 in the next 2 ms for 12 μm droplets. The error bars in $f_{isc}$ represent the standard deviation of individual recordings collected at each distance and include variations in the hit rate and droplet trajectory jitters. The droplet trajectory jitter is expected to arise from jet breakup to form droplets and from the freezing process of the droplets. The onset temperatures where ice is first detected on these time scales are 232 and 230 K for the 9 and 12 μm droplets, respectively. We use $f_{isc}$ to estimate $J$ assuming, as in other nucleation rate studies, that the nucleation rate follows Poisson statistics and the observed Bragg reflections expected from ice in each shot originate from...
The minimal nucleation rate reached in the “no-man’s land”, \( f_{\text{max}} = 1.1 \times 10^{13} \text{ cm}^{-3} \text{ s}^{-1} \), is calculated analogously using \( q \approx 10^{7} \text{ K/s} \). This is very close to the nucleation rates measured in the present work on evaporatively cooled microdroplets and suggests that indeed the nucleation rate does not need to increase very much in the “no-man’s land” compared to its value at 227 K.

The crystallization kinetics of hyperquenched glassy water to cubic ice was measured by Hage et al.\(^{52}\) on layers of many micrometers in thickness, who report that it takes about 10^4 seconds to reach full crystallization of a volume of about 10^{-2} \text{ cm}^3 at 140 K and about 10^3 seconds at 146 K. Both nucleation and growth are diffusion-controlled under these conditions, and so it is not straightforward to extract nucleation rates from the data. Hage et al.\(^{52,53}\) also report that not only crystallization rates, but even the mechanism of crystallization as inferred from the Avrami-exponent is significantly altered when comparing annealed and unannealed films. Assuming only a single nucleation event, i.e., growth being much faster than nucleation, the nucleation rate calculates to \( J \sim 10^{-12} \text{ to } 10^{-15} \text{ cm}^{-2} \text{ s}^{-1} \) at 155–138 K.\(^{53}\) Nucleation rates around \( T_{\text{fl}} \) have also been measured on much thinner annealed amorphous thin films.\(^{1,31}\) Although these rates are much lower than those of nanodroplets as expected, the measured rates (\( J \sim 10^{-10} \text{ cm}^{-2} \text{ s}^{-1} \)) by Safarik and Mullins\(^{51}\) are higher by 10 orders of magnitude lower rate and show weaker temperature dependence than the data of Hagen et al.\(^{19–21}\) The observation that microdroplets vitrify entirely at cooling rates of \( 10^5 \text{ K/s} \) sets an upper limit to the maximum nucleation rate within the “no-man’s land” of \( \sim 10^{16} \text{ cm}^{-3} \text{ s}^{-1} \) (see \( f_{\text{max}}\)-limit in Figure 2b). At a rate of \( q \approx 10^{7} \text{ K/s} \), it takes \( \Delta t \approx 7 \text{ ms} \) to cool through the \( \sim 230 \) to \( \sim 160 \text{ K} \) interval. Assuming that a single nucleation event (\( N = 1 \)) already suffices to crystallize a droplet of radius \( \langle r \rangle \), one obtains \( f_{\text{max}} = N/(4/3)\pi\langle r \rangle\Delta t \approx 1.1 \times 10^{16} \text{ cm}^{-3} \text{ s}^{-1} \) for \( \langle r \rangle = 1.5 \mu\text{m} \). This estimate represents a hard limit for the maximum nucleation rate of microdroplets within the “no-man’s land” for bulk water at \( p \leq 1 \text{ bar} \). That is, the nucleation rate of microdroplets of 3–12 \( \mu\text{m} \) diameter is at least 7 orders of magnitude lower than the one in nanodroplets of 6–12 nm diameter. This discrepancy may be attributed to the much higher surface-to-volume ratio that affects the internal pressure of the nanodroplets and potentially increases surface nucleation (see Supporting Information on nucleation rate sensitivity to pressure). Given these differences between nano- and microdroplets, we only compare our data against those from previous microdroplet experiments where we expect to have similar conditions.\(^{40–48}\)

The obtained nucleation rates are consistent with earlier hyperquenching experiments on micrometer-sized droplets,\(^{10–4}\) indeed, practically all evaporatively cooled droplets crystallize at cooling rates of up to \( 10^7 \text{ K/s} \). Going beyond this earlier work,\(^{40–41}\) this suggests that the crystallization of droplets observed in these experiments takes place only slightly below \( T_{\text{fl}} \sim 232 \text{ K} \). Since droplets cooled faster by only 3 orders of magnitude vitrify completely, this suggests that the maximum of the homogeneous ice nucleation rate is located below 227 K and most likely not too far below. This suggestion finds confirmation in the nucleation rate data from the current measurements (red and brown filled circles) in Figure 2a, which turn out to be quite insensitive to temperature in the 227–232 K interval. Connecting all data-points measured on microdroplets together with the hyperquenching experiments it would be consistent to assume that the nucleation rate further bends down and eventually flattens out as the temperature is lowered beyond 227 K as indicated by the red line in Figure 2a.

Figure 2b shows ice nucleation rates plotted as a function of temperature for previous studies (black, blue, and green symbols). Data above \( T_{\text{fl}} \)\(^{46–50}\) (Figure 2a) represent the current state of rate measurements (see ref 40 for an exhaustive list of recent measurements) and are, within the uncertainties due to both temperature and cooling rate, consistent with each other and with the qualitative picture of how the nucleation rate should increase with decreasing temperature as described earlier. As temperature decreases below \( T_{\text{fl}} \), data from both Hagen et al.'s\(^{30}\) and our experiments result in higher rates than those above \( T_{\text{fl}} \). However, our measurements yield a 3–7 orders of magnitude lower rate and show weaker temperature dependence than the data of Hagen et al. Notably, Hagen et al. relied on using a droplet growth model\(^{50}\) that may introduce large uncertainties in the estimation of the temperature and droplet size.\(^{51}\) In addition, Riechers et al.\(^{47}\) have reported a slow-down in the rate increase similar to the present study, albeit to lesser extent, as temperature decreases toward \( T_{\text{fl}} \) which they attributed to a rapid decrease in \( D \).

Ice nucleation experiments from nanodroplets of 6–12 nm diameter allow rate measurements at temperatures between 170 and 215 K, which is well below \( T_{\text{fl}} \) (Figure 2b). These rates (\( \sim 10^{53} \text{ cm}^{-3} \text{ s}^{-1} \)) are much higher than those from microdroplets and appear less dependent on temperature.\(^{19–21}\) The observation that microdroplets vitrify entirely at cooling rates of \( 10^5 \text{ K/s} \) sets an upper limit to the maximum nucleation rate within the “no-man’s land” of \( \sim 10^{16} \text{ cm}^{-3} \text{ s}^{-1} \) (see \( f_{\text{max}}\)-limit in Figure 2b). At a rate of \( q \approx 10^{7} \text{ K/s} \), it takes \( \Delta t \approx 7 \text{ ms} \) to cool through the \( \sim 230 \) to \( \sim 160 \text{ K} \) interval. Assuming that a single nucleation event (\( N = 1 \)) already suffices to crystallize a droplet of radius \( \langle r \rangle \), one obtains \( f_{\text{max}} = N/(4/3)\pi\langle r \rangle\Delta t \approx 1.1 \times 10^{16} \text{ cm}^{-3} \text{ s}^{-1} \) for \( \langle r \rangle = 1.5 \mu\text{m} \). This estimate represents a hard limit for the maximum nucleation rate of microdroplets within the “no-man’s land” for bulk water at \( p \leq 1 \text{ bar} \). That is, the nucleation rate of microdroplets of 3–12 \( \mu\text{m} \) diameter is at least 7 orders of magnitude lower than the one in nanodroplets of 6–12 nm diameter. This discrepancy may be attributed to the much higher surface-to-volume ratio that affects the internal pressure of the nanodroplets and potentially increases surface nucleation (see Supporting Information on nucleation rate sensitivity to pressure). Given these differences between nano- and microdroplets, we only compare our data against those from previous microdroplet experiments where we expect to have similar conditions.\(^{40–48}\)

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magnitude than the nucleation rates deduced here from the data of Hage et al.\textsuperscript{52,53} and even comparable to those of microdroplets above $T_H$ ($\lesssim 10^{10}$ cm$^{-3}$ s$^{-1}$). Jenniskens and Blake\textsuperscript{1} measured rates ($J \sim 10^{12}$–$10^{15}$ cm$^{-3}$ s$^{-1}$) that are even higher than our rates below $T_H$ which is problematic considering that amorphous ice can be observed up to 160 K under similar time scales.\textsuperscript{12} The huge discrepancy between the different data sets suggests that different mechanisms of crystallization were probed in these studies. Thin film nucleation rates that are greater than or equal to $\sim 10^{11}$ cm$^{-3}$ s$^{-1}$ seem unlikely considering that in these experiments the sample is heated at much lower rates ($\sim 0.1$–1 K/s) than the cooling rates ($\sim 10^5$ K/s) required to measure nucleation rates $\gtrsim 10^{11}$ cm$^{-3}$ s$^{-1}$ from microdroplets within “no-man’s land”. It might be possible that their films were preseeded with ice contamination, and so their rates might be limited by growth rather than by nucleation. However, the submicrometer sample thickness used in these thin film experiments may reduce sample volume significantly and thus potentially allow such high nucleation rate measurement. In addition, the rates measured in these experiments on submicrometer samples\textsuperscript{1,51} differ by 4–7 orders of magnitude and show very different temperature dependence. Although it is unclear why there are such large differences in the crystallization studies on submicrometer films\textsuperscript{1,51} and much thicker films,\textsuperscript{52,53} different sample preparation and measurement techniques in these experiments can affect the resulting rates. More nucleation rate studies are necessary to understand the differences between these results and, more importantly, to improve our understanding of water and crystallization kinetics near $T_g$.

To account for the results of previous rate measurements, Jenniskens and Blake have considered liquid water as either “fragile” or “strong”, which is associated with how the diffusivity $D$ changes with temperature. By assuming the temperature-dependent $D$ of a “fragile” liquid within the CNT framework, one can connect most nucleation rate data within “no-man’s land” to those at higher temperatures,\textsuperscript{9–21,46–49,51} with the exception of the data from Hagen et al.\textsuperscript{50} for reasons discussed above and also our present data. At low temperatures near $T_g$ liquid water has been shown to behave like a “strong” liquid by recent dielectric relaxation and calorimetric measurements.\textsuperscript{36} This “strong” description of water predicts nucleation rates that show a temperature dependence similar to the one given by Jenniskens and Blake.\textsuperscript{1} In contrast, the rates reported by Safarik and Mullins\textsuperscript{51} exhibit a temperature dependence that is closer to that of a “fragile” than a “strong” liquid.

The current nucleation rate measurements in conjunction with previous microdroplet measurements\textsuperscript{46–49} and the nucleation rate limits within “no-man’s land” deduced from the hyperquenching experiments pioneered by Mayer,\textsuperscript{40–44} however, show a slower than expected nucleation rate increase as temperature decreases toward and crosses $T_H$ giving a hint of an anomalous behavior. This slower increase makes the nucleation rate deviate away from the CNT-predicted rate of a “fragile” liquid and move toward that of a “strong” liquid just below $T_H$ (Figure 2b). We suggest that an expected rapid
decrease in liquid water diffusivity $D_{\text{w}}^{22,29}$ may lead to a significant increase in $\Delta f^4$, which can explain the slow nucleation rate increase just below $T_{\text{nl}}$. In addition, recent ultrafast X-ray scattering measurements have shown a continuous but accelerated change toward a low-density liquid at similar temperatures.\(^{39}\) We thus hypothesize that these observations might be related to the possible “fragile-to-strong” liquid transition\(^{27,38}\) and the findings from molecular dynamics (MD) simulations using the mW water model\(^{34}\) that suggest that the structural transformation controls the ice crystallization rate in “no-man’s land”.\(^{10}\) By contrast, the nucleation rate measurements from nanodroplets appear to follow the CNT-predicted nucleation rate of a “fragile” liquid at temperatures below $T_{\text{nl}}$. Thus, we also hypothesize that the pressure increase within the nanodroplets may have kept the liquid as a “fragile” liquid rather than transitioning into a “strong” liquid within the temperature range of these measurements in “no-man’s land”. However, further investigations at a wider range of temperatures and pressures are needed to definitively understand the anomalous water properties and their influence on ice nucleation kinetics, particularly within “no-man’s land”.

In conclusion, we extend ambient pressure nucleation rate measurements to $\sim 227$ K, well below $T_{\text{nl}}$ and within “no-man’s land”, and observe a slower increase in nucleation rate compared to that expected from prior measurements. Overall, the microdroplet ice nucleation rate data in “no-man’s land” appear consistent with the hypothesis of supercooled water at ambient pressure crossing over from being a “fragile” liquid to a “strong” liquid just below $T_{\text{nl}}$.\(^{31,37,38,52}\) However, more work is needed in the future to clearly determine if this indeed is the case.

# EXPERIMENTAL SECTION

X-ray scattering from deionized water (PURELAB Ultra Genetic, resistivity 18.2 M$\Omega$cm at 298 K) droplets of 9 and 12 $\mu$m diameters upon deep supercooling (<255 K) was measured at the LCLS Coherent X-ray Imaging instrument (CXI).\(^{56}\) A 9.4 keV photon energy and 50 fs nominal pulse duration X-ray laser with spot size of $\sim 1$ $\mu$m$^2$ (9 $\mu$m droplets) and $\sim 10$ $\mu$m$^2$ (12 $\mu$m droplets) at the interaction point were used. The Cornell-SLAC pixel array detector (CSPAD) located 132 mm (9 $\mu$m droplets) and 139 mm (12 $\mu$m droplets) downstream from the interaction point was used to record single-pulse scattering patterns from individual droplets at 120 Hz repetition rate. A 3-axis motorized sample stage was used to control the distance between the nozzle and the interaction point, and thus to determine the droplet temperature. The temperature estimate was based on the Knudsen theory of evaporative cooling with a calibration made by comparing the liquid water wide-angle scattering signal in this experiment against larger microdroplets evaporating in vacuum with diameters of 34 and 37 $\mu$m as well as against synchrotron measurements using a cooling cell down to 251 K.\(^{39}\) The evaporative cooling model was further verified through MD simulations of droplet cooling.\(^{39}\) The temperature difference between core and surface within 9 and 12 $\mu$m droplets is less than 1 K after 2 ms of travel time. In addition, we estimate the droplet internal pressure to be $\sim 0.03$ MPa, primarily due to surface tension exerting Laplace pressure on the droplet. We assumed this to be a good approximation to ambient pressure ($\sim 0.1$ MPa). Finally, data at each distance were sorted according to water-only or ice-containing scattering patterns, and was then used to estimate the fraction of ice-containing shots ($f_{\text{ice}}$). Further details are given in the Supporting Information of ref 39. The final data with their respective accuracy are tabulated in Supporting Information.

We used $f_{\text{ice}}$ to calculate the ice nucleation rate ($J$) at time $t_{n+1}$ using

$$J(t_{n+1/2}) = -\frac{\ln[(1 - f_{\text{ice}}(t_{n+1}))/ (1 - f_{\text{ice}}(t_n))]}{V_{\text{drop}} \times (t_{n+1} - t_n)}$$

(refs 21 and 46) where $V_{\text{drop}}$ is the droplet volume, and subscripts $n$ and $n+1$ represent successive measurements. The corresponding temperature, $t_{n+1/2}$, is obtained from the droplet temperature estimate. We assume, as in other nucleation rate studies,\(^{39–41,46–49}\) that nucleation is a stochastic process that follows Poisson statistics, that diffraction collected in each shot comes from a single crystal, that the growth of a nucleus to a detectable size is much faster than formation of a nucleus, and that the difference in ice nucleation rate is negligible between adjacent data points (i.e., that the temperature difference between adjacent data points is small). These are reasonable assumptions because the cooling rate is slow when ice nucleation occurs (Figure 1) and our measurements generally are limited by thermodynamics (although kinetics becomes increasingly important upon decreasing temperature). In addition, we can detect ice crystals as small as $\sim 50$ nm in diameter and detect ice in the droplet above 0.05% by mass.\(^{39}\) However, our detection limit is much larger than the estimated critical ice nucleus diameter of $\sim 2.5$ nm. Although the structure of a critical ice nucleus has not been determined experimentally, simulations suggest that the critical ice nucleus is rich in cubic ice.\(^{14}\) The average of the ice-containing diffraction patterns, however, shows a hexagonal ice pattern,\(^{55}\) but we cannot exclude that the heat released during growth effectively anneals the ice nuclei into hexagonal ice crystals. Moreover, we observed $\sim 30$ Bragg reflections on average in diffraction patterns classified as hexagonal ice compared to $\sim 2$ Bragg reflections expected on average from a randomly oriented perfect hexagonal ice crystal. These extra Bragg reflections might be due to dendritic growth with defects where the heat release during rapid growth could lead to splitting of the original nucleus into several crystallites. Alternatively, the observed Bragg reflections might originate from $\sim 15$ individual nuclei. We thus account for the possibility of observing $\sim 15$ nuclei in the upper nucleation rate error bar in addition to the $f_{\text{ice}}$ uncertainty. Finally, we note that the crystallization times may represent a combination of nucleation and growth at the lowest temperature of $\sim 227$ K (see Supporting Information).

# ASSOCIATED CONTENT

Supporting Information

CNT that accounts for increased pressure, discussion on crystallization time that may represent nucleation and growth at the lowest temperature of $\sim 227$ K, analysis of nucleation rate sensitivity to increased pressure according to CNT, table of experimental data and calculated ice nucleation rates, and table of thermophysical and transport properties of water. The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcl.5b01164.

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