The moment of initial crystallization captured on functionalized nanoparticles

Hironobu Machida, Takeshi Sugahara & Izumi Hirasawa

Even if a liquid is cooled below its melting point, the liquid state can be maintained under certain conditions. This state is called supercooling. Spraying fine particles of dry ice or silver iodide induces a phase change from supercooled droplets to ice grains. However, the mechanism by which crystallization seeds diminish supercooling is not well understood. Here, we captured the moment when a cluster, which is the smallest structural unit of a crystal, envelops a silver nanoparticle. As a result of observing the structure of a supercooled aqueous solution of a clathrate hydrate, we found that silver nanoparticles accelerate the formation of clusters, whereas the noble metals palladium, gold and iridium likewise form nanoparticles but do not promote crystallization. Our discoveries elucidate the mechanism of heterogeneous nucleation during a phase change in clathrate hydrates. We anticipate our discovery to be the starting point for the control of supercooling, a technique that can be applied to enhance the production efficiency and quality of manufactured products.
Heat storage technology is anticipated to have the potential for the effective use of unused thermal energy currently vented into the environment, such as automobile exhaust heat, factory exhaust heat, and household waste heat. There are various technologies for heat storage, of which latent heat stored by phase change at the desired temperature is of particular interest due to its ease of use and low cost. Latent heat storage materials have been actively studied since around 1980, but practical examples are currently limited to cold storage boxes for outdoor use and pocket heaters. One major impediment to its practical use is supercooling, a phenomenon in which the liquid phase is maintained even when it is cooled below the freezing point. Supercooling phenomena lead to problems such as unreliable heat storage and increased cooling costs. To minimize supercooling, it is important to understand the crystallization mechanism of supercooled aqueous solutions.

Here, we focus on clathrate hydrate crystallization from an aqueous solution. A clathrate hydrate is composed of water and guest molecules. The hydrogen-bonded water molecules surround the guest molecule to form a cage structure. Among clathrate hydrates, the hydrates formed from aqueous solutions of quaternary ammonium salts have been classified as semiclathrate hydrates. Tetra-n-butylammonium bromide (TBA) semiclathrate hydrate has been investigated in the fields of gas separation, as well as thermal storage.

We have observed the nanometer-scaled solution structure in supercooled TBAB aqueous solutions by applying the freeze-fracture replica method and have elucidated the relationship between the hysteresis phenomenon in recrystallization (called the memory effect) and the residual solution structure after solid dissociation. In both the supercooled aqueous solution and the crystal, granular structures (called clusters) measuring 10–30 nm in diameter are present as a basic structural unit. During crystallization, the number density of the clusters significantly increases. Immediately prior to crystallization, the clusters are densely packed in the supercooled aqueous solution. To diminish supercooling, therefore, it is necessary to promote cluster formation and densification. The memory effect contributes considerably to raising the formation temperature of TBAB semiclathrate hydrate and reducing the degree of supercooling. However, it is not sufficient, because few residual solution structures (clusters) can survive long enough, even at a temperature 2 K above the decomposition temperature of TBAB semiclathrate hydrate. The development of chemically and thermodynamically stable materials that promote cluster formation (hereinafter called “crystallization seeds”) is therefore necessary. One example of the prevention of supercooling is seen with repetition fields of gas fields and household waste heat. There are two kinds of rainy clouds: a cold cloud below 273 K and a warm cloud above 273 K. To make it rain artificially in cold clouds, it is necessary to form ice crystallites in a supercooled cloud. The fine silver iodide particles nucleate ice crystallization. It is thought that the lattice constants of the hexagonal form of silver iodide are very similar to those of ice, which is why silver iodide works as a nucleus. However, the mechanism of how crystallization seeds diminish supercooling is not fully known because the crystal structure of seeds is not always similar to that of the crystal nucleated by crystallization seeds. Many researchers have investigated the role of crystallization seeds and the secondary nucleation mechanism induced by crystallization seeds.

Here, we reveal the moment of initial crystallization, which is the formation of 10–30 nm clusters encircling silver (Ag) nanoparticles, in the tetra-n-butylammonium 3-methylpentanoate and tetra-n-butylammonium 2-ethylbutyrate (TBA-3MP) and tetra-n-butylammonium 2-ethylbutyrate (TBA-2EB) semiclathrate hydrate systems. The synergetic effect of Ag nanoparticles and tetra-n-butylammonium fluoride (TBAF) diminishes supercooling to a practicable level during the formation of these semiclathrate hydrates.

**Results and discussion**

Effect of metallic carboxylates with or without TBAF on the degree of supercooling. Figure 1 shows the degree of supercooling (ΔTsup.) in the formation of TBA-3MP and TBA-2EB semiclathrate hydrates with various metallic carboxylates and or TBAF added. The decomposition temperature of TBA-3MP semiclathrate hydrate coexisting with TBAF and silver pentanoate (Ag-PA) is 278.1 ± 0.2 K at a mass fraction, wTBA-3MP, of 0.42 (Supplementary Fig. 1), which is the same as the reported decomposition temperature of TBA-3MP semiclathrate hydrate without any additives. Without any additives present (Sample #1), a degree of supercooling greater than 10 K is necessary to...
form a TBA-3MP semiclathrate hydrate. Even if only TBAF (Sample #2) and a transition-metallic acetate, as well as TBAF (Samples #3-9 and #14-16), were added, the degree of supercooling was little changed. When either silver acetate or Ag-PA was added, the degree of supercooling was brought down to approximately 6 K. Further improvement (approximately 4 K) of the degree of supercooling results from the synergetic effect of silver acetate (or Ag-PA) with TBAF. The synergetic effect of silver carboxylate with TBAF is also similar to that in TBA-2EB aqueous solution. The time required by the end of crystallization of TBA-3MP semiclathrate hydrate also depends on the amounts of Ag-PA and TBAF present (Fig. 2b), whereas the degree of supercooling is almost independent of their amounts (Fig. 2a).

Why did only Ag, but not Pd, Ir, or Au nanoparticles diminish supercooling? We have already reported the formation of corresponding metal nanoparticles of Pd, Ir, and Au32. In either system, carboxylate ion is thought to contribute to preventing metal nanoparticles from aggregating due to steric hindrance. To understand the unique nature of Ag and the chemical forms of Ag on an atomic or molecular level, we measured the XAS of Ag, Pd, Ir, and Au. XAS is a tool for gaining information on the type of neighboring atoms, which can be usually deduced by comparison of the XAS spectrum of the sample with those of suitable reference compounds. Shown in Figs. 4a and b are the Ag K-edge X-ray absorption near-edge structure (XANES) spectra of the sample solutions (#13 measured at 273 K and 293 K) along with the reference compound silver fluoride (AgF), Ag-PA, silver nitrate (AgNO3), Ag foil, silver monoxide (AgO), and silver(I) oxide (Ag2O). On comparing them at the rising-edge region of 25.50–25.51 keV, the rising-edge behavior of the sample solutions measured at 273 K and 293 K was somewhat similar to that of AgF, AgNO3, and Ag-PA, whereas it was different from that of Ag foil, AgO, and Ag2O. At the energy side higher than the absorption edge, the Ag K-edge XANES spectrum (orange) of sample solution #13 measured at 293 K was somewhat flat. It appears that Ag has no interaction with any specific atoms in the sample solution. On decreasing the temperature to 273 K, the XANES spectrum (red) changed and showed a peak at 25.51 keV. The spectral shape is similar to that of AgF in an aqueous solution. These XANES spectra suggest that Ag exists without any preferential binding partner at 293 K, but, at 273 K, the
Interaction between Ag and fluorine, in addition to Ag and pentanoate, strengthens. We attempted to analyze the Fourier-transformed-extended X-ray absorption fine structure (FT-EXAFS) region of Ag. However, the S/N ratio was so low that, due to insufficient Ag concentration, we could not obtain any meaningful data. The Pd K-edge, Ir LIII-edge, and Au LIII-edge XANES, and FT-EXAFS spectra of the sample solutions #14, #15, and #16 are shown in Fig. 4c, d, and e along with possible reference compounds. Both the XANES and FT-EXAFS spectra reveal that the environment of a Pd, Ir, or Au atom is similar to that of their respective metal acetates: in other words, Pd, Ir, and Au are present as acetate-coordinated complexes in the sample solutions. These XAS results indicate an answer to the question as to why Ag effectively diminishes supercooling, but Pd, Ir, and Au do not. Since Pd, Ir, and Au ions are divalent, tetravalent, and trivalent, respectively, they are coordinated with two, four, and three acetate ions. These multiple acetate ions prevent other ions from approaching the Pd, Ir, and Au atoms. Since the Ag ion is monovalent, however, there is just enough space for other ions to approach Ag, even if it is coordinated with an acetate ion. In Sample solution #13, fluoride ion is present as a counter anion to the TBA cation. Therefore, with a decrease in temperature, the spectral element of AgF increases in addition to that of Ag-PA in the XANES spectrum of Sample solution #13. The approach of fluoride ions to Ag nanoparticles yields an enriched area of fluoride ions around the Ag nanoparticles. Local enrichment of

![Fig. 3 Cluster formation of semicladrate hydrate around silver nanoparticles. a] HAADF-STEM image of replica films prepared from samples #1, #2, #11, and #13 of TBA-3MP aqueous solutions listed in Fig. 1a. A number of silver nanoparticles smaller than 10 nm were observed in samples #11 and #13. b) SE-STEM image (different area) of the same replica film as “13-a” in a. EDX measurements were performed at points 1-9. c) Typical EDX charts at points 1, 5, and 6 in b. EDX charts of points 1-4 reveal the existence of silver nanoparticles incorporated in replica film. Like enclosing silver nanoparticles, clusters of 10-30 nm were formed.
fluoride ions increases the probability of formation of 10–30 nm TBAF (or TBAF + TBA-3MP mixed) semiclathrate hydrate clusters around Ag nanoparticles at 281 K (as shown in Fig. 3b) since Ag nanoparticles also effectively diminish supercooling in the TBAF semiclathrate hydrate formation (Fig. 5). The 10–30 nm clusters of the semiclathrate hydrate act as crystalization seeds for TBA-3MP semiclathrate hydrate formation due to the similarity (tetragonal lattice) of the two crystal structures (Supplementary Table 1 and Supplementary Fig. 2). It follows that the TBA-3MP semiclathrate hydrate can be formed at a relatively small degree of supercooling.

To summarize this study: we have reported that Ag nanoparticles act to diminish supercooling under the coexistence of TBAF during the formation of TBA-3MP and TBA-2EB semiclathrate hydrates and have elucidated the mechanism, which is the promotion of the formation of 10–30 nm clusters around Ag nanoparticles, through STEM observation using the freeze-fracture replica method and XAS. Furthermore, neither Ag nanoparticles nor TBAF has an effect on diminishing supercooling alone. The present results suggest a useful strategy for diminishing supercooling of TBA salt semiclathrate hydrate formation when used as heat storage materials. A role for Ag compounds in diminishing supercooling and a possible mechanism can also be proposed that does not depend on the similarity of the crystal structure to Ag compounds.

Methods

Samples. Tetra-n-butylammonium-3-methylpentanoate (TBA-3MP) and Tetra-n-butylammonium-2-ethylbutyrate (TBA-2EB) were synthesized by aqueous neutralization of tetra-n-butylammonium hydroxide (TBAOH, Tokyo Chemical Industry) with 3-methylpentanoic acid (Tokyo Chemical Industry) and 2-ethylbutyric acid (Tokyo Chemical Industry), respectively. The resultant aqueous solutions were completely dried under vacuum at 40 °C to obtain TBA-3MP and...
TBA-2EB. The obtained products were confirmed by $^1$H NMR. The purity of the synthesized TBA-3MP or TBA-2EB was virtually the same as that of TBA-OH (on a water-free basis) and 3-methylpentanoic acid or 2-ethylbutyric acid. The majority of the impurities were tri-n-butylamine, 1-bromobutane, and KBr that were originally present in the TBA-OH aqueous solution. In the NMR and ion chromatographic (Thermo Fisher Scientific, Integrion HPIC) analyses, the concentrations of these expected impurities, as well as the others, were below the detection limits. TBA-3MP (mass fraction $\omega_{\text{TBA-3MP}} = 0.42$) and TBA-2EB (mass fraction $\omega_{\text{TBA-2EB}} = 0.36$) aqueous solutions were prepared. Close to equimolar amounts of silver nitrate (Fujifilm Wako Pure Chemical Corporation) and pentanoic acid (Tokyo Chemical Industry) were added to water under dark conditions. The addition of sodium hydroxide (Tokyo Chemical Industry) aqueous solution for neutralization led to the precipitation of silver pentanoate (Ag-PA). This precipitate was filtered, washed, and dried. Ag-PA and tetra-n-butyrammonium fluoride (TBAF, Tokyo Chemical Industry) solution was added to TBA-3MP aqueous solution ($\omega_{\text{TBA-3MP}} = 0.42$) at a molar ratio of Ag: F: TBA-3MP (or TBA-2EB) = 1: 10: 2. The addition of TBAF supplementarily improves the solubility of Ag-PA. Other sample solutions of the metallic acids listed in Fig. 1a were also prepared at the same molar ratio. They were then stirred ultrasonically and placed under diffuse sunlight and a fluorescent room lamp (Panasonic, FHF32EX-N H) for one day. The wavelength and energy irradiated to the samples were measured with an illuminance spectrophotometer (Konica Minolta, CL500A). The result (not shown) indicated low optical energy in the ultraviolet region below 400 nm.

**Structural observation in aqueous solution.** A freeze-fracture replica method was used for structural observation in aqueous solution. First, a droplet (1 μL) of the aqueous solution at the desired temperature was injected into a platinum cell with an internal diameter of 1 mm. Next, the platinum cell containing the droplet was dipped into liquid 2-methyl butane at its melting point (113 K). This caused the droplet to vitrify. The vitrified droplet was then cut in a vacuum chamber (JEOL, EM-19501 JFD-V) maintained at approximately 80 K to expose the cut surface. To replicate these irregularities, carbon was vapor-deposited on the cut surface immediately after cutting the vitrified droplet. The vitrified droplet was then dissolved in pure water to obtain an approximately 4 nm-thick carbon replica film. The obtained replica film was picked up on a 200-mesh copper grid (VECO, square type). We prepared at least two replica films under a condition. For each, more than 10 fields were observed using an SEM (Carl Zeiss, Ultra plus, acceleration voltage: 2 kV, detector: InLens) or STEM (Hitachi, HD-2700, acceleration voltage: 200 kV, detector: secondary electron detector (SE) or High angle annular dark-field (HAADF)). The metal nanoparticles remaining in the replica film were analyzed via EDX (EDAX, TEAM, STEM attached, acceleration voltage: 200 kV). In Fig. 3c, the copper originates in the mesh used for SEM observation. The EDX peak of the Si component was also detected as an analysis point outside the Ag nanoparticles. There was no sulfur in the chemicals used in this study. Sulfur derived from rubber washers might have reacted with Ag nanoparticles being kept under dry conditions before EDX measurement.

**Measurement of the degree of supercooling.** Aliquots of approximately 1 g of the sample solutions (Fig. 1a) were introduced into glass vials. A T-type thermocouple was placed on the outside of the bottom of each glass vial. All the glass vials were placed in a thermostatic bath (Espec, SU-241). The bath was programmed to hold the temperature at 283 K for 60 min and then cool down at 0.05 K min$^{-1}$ from 283 K to 263 K. The onset temperature and offset time of the exothermic peak derived from crystallization was determined as the crystallization temperature and the time for crystallization to be completed, respectively. The difference between the crystallization and equilibrium temperatures was taken as the degree of supercooling. These measurements were repeated 14 times and the average degree of supercooling was calculated.

**Data availability**

The data that support the findings of this study are available from the corresponding authors upon reasonable request.

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**Author contributions**

H.M., T.S. and I.H. conceived the research idea. H.M. performed the measurements of the degree of supercooling and analyzed the XAS data. H.M. and T.S. summarized the research results and wrote the manuscript after discussion with and obtaining input from all the coauthors.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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Correspondence and requests for materials should be addressed to H.M. or T.S.

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