Real-Time Observation of Water-Soluble Mineral Precipitation in Aqueous Solution by In Situ High-Resolution Electron Microscopy

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Supporting Information

ABSTRACT: The precipitation and dissolution of water-soluble minerals in aqueous systems is a familiar process occurring commonly in nature. Understanding mineral nucleation and growth during its precipitation is highly desirable, but past in situ techniques have suffered from limited spatial and temporal resolution. Here, by using in situ graphene liquid cell electron microscopy, mineral nucleation and growth processes are demonstrated in high spatial and temporal resolution. We precipitate the mineral thenardite (Na$_2$SO$_4$) from aqueous solution with electron-beam-induced radiolysis of water. We demonstrate that minerals nucleate with a two-dimensional island structure on the graphene surfaces. We further reveal that mineral grains grow by grain boundary migration and grain rotation. Our findings provide a direct observation of the dynamics of crystal growth from ionic solutions.

KEYWORDS: in situ graphene liquid cell electron microscopy, water-soluble mineral, nucleation and growth, grain boundary migration, grain rotation

The crystallization of dissolved salts in aqueous solution is involved in bio-, geo-, and soil mineralization and is the most common way of producing crystals in nature.$^{1-3}$ A fundamental understanding of crystallization from solution is necessary to control and optimize crystal structure, morphology, and the size of the products. The key steps affecting the final structures and properties are the crystal nucleation and subsequent growth.$^{4-6}$ Despite great interest in this pathway, the pathways of crystallization are still ambiguous because processes that occur in liquid are difficult to access with current experimental methods. While an aqueous environment presents many challenges to experimental investigation, grazing-incidence small-angle X-ray scattering (GISAXS), atomic force microscopy (AFM), and optical microscopy have previously been employed to obtain dynamical information on mineral nucleation and growth, such as calcium carbonate, iron oxide, sodium chloride, and silica.$^{7-10}$ Using these in situ methods, it has been demonstrated that, depending on the supersaturation, precipitation occurs predominantly by attaching molecules at screw dislocations or through two-dimensional (2D) surface nucleation.$^{4-6}$ However, these experimental methods have their limitations. For example, in situ spectroscopy methods cannot provide morphological information, and in situ AFM can be applied only for fixed samples on a substrate with limited temporal resolution, constraining the utility in colloidal synthesis investigation.

Liquid cell transmission electron microscopy (TEM) provides unique advantages, by allowing direct observation of the morphological and structural changes of particles in solution during chemical and physical reactions. This method has been employed for the observation of nucleation and growth of colloidal nanocrystals.$^{10-16}$ Since mineral precipitation by water evaporation is technically difficult to achieve in a TEM, application of in situ liquid cell TEM has been limited to the experiments initiated by electron-beam-induced radiolysis of water instead of water evaporation occurring in nature but also show the precipitation processes of the mineral.

Received: July 2, 2015
Accepted: December 3, 2015

DOI: 10.1021/acsnano.5b04064
ACS Nano XXXX, XXX, XXX--XXX
thenardite (Na₂SO₄), which is used in the glass, paper, and medical industries. 

RESULTS AND DISCUSSION

For observation of mineral precipitation in high spatial and temporal resolution, we employ graphene liquid cells (GLCs) in a TEM to achieve high contrast imaging of the target materials in solution. For observation of mineral precipitation in high spatial and temporal resolution, we employ graphene liquid cells (GLCs) in a TEM to achieve high contrast imaging of the target materials in solution. 

Figure 1. Time-lapse TEM imaging of mineral nucleation from aqueous solution. (A) Schematic of the GLC with sodium persulfate aqueous solution trapped between FLG sheets. Sodium and persulfate ions are solvated by water molecules. (B) Time-lapse, color-contrasted, dark-field TEM images of mineral precipitation by electron beam irradiation. (C) Total projected area (A) of all precipitates versus electron beam irradiation time (t). (D) A versus the number of precipitates (N). Inset, mean radius (r) of precipitates versus t. Line in (d) indicates the linear fitting.

Figure 2. HR-TEM imaging of the sodium sulfate nucleus. (A) HR-TEM image of hexagonal sodium sulfate mineral with [12̅13̅] zone axis. Inset is the FFT image of (A). (B) Structure of hexagonal sodium sulfate crystal (a = b = 0.54 nm and c = 0.72 nm). (C) HR-TEM image of the sodium sulfate nucleus with thickness less than 5 nm deposited on the FLG with ~12 layers.

Figure 3. HR-TEM imaging of mineral grain growth by the grain boundary migration and the grain rotation. (A) Time-serial HR-TEM images (top row of images) and the corresponding false-colored schematics identifying grains and grain boundaries (bottom row of images). At 0 s, the precipitate is composed of four crystalline grains of G1, G2, G3, and G4 and five grain boundaries of GB₁₂, GB₁₃, GB₂₃, GB₂₄, and GB₃₄. α₁₂, α₁₃, α₁₄, and α₁₅ of the angles between grain boundaries are 122, 128, and 110°, respectively. Au particles are loaded as an indicator of liquid environment. Due to the migration of GB₁₂ and GB₂₄ toward center of curvature, the G2 changes shape, which assumes a planar rectangular shape at 7.0 s and a triangular shape at 30.1 s, and eventually disappears at 54.6 s. In the schematics at 0, 7.0, and 30.1 s, arrows indicate the directions of grain boundary migration. (B) (0001) diffraction spot of G1 rotates relative to vertical lines by 8° and 1° at 34.3 and 54.6 s, respectively. (11̅01) diffraction spot of G4 rotates relative to vertical lines by 17° and 16° at 34.3 and 54.6 s, respectively.
biological samples. The addition of a few graphene sheets has minimal impact on TEM resolution compared to the impact of the liquid itself.

Since sodium sulfate nanocrystals are indiscernible in bright-field TEM images due to their low mass–thickness contrast, to observe the mineral nucleation, we acquire time-lapse dark-field TEM images by choosing the precipitates’ diffraction spots with an objective aperture (Figures 1B and S4 and Movies S1 and S2). Initially, sodium persulfate salts are totally dissolved in water, which is indicated by the halo feature in the diffraction pattern (Figure S5 and Movie S3). Once the electron beam irradiation begins, minerals precipitate and nucleate. The diffraction spots of the precipitates correspond to sodium sulfate crystals (PDF #27-0791) (Figure S5). Figure 1C shows the total projected areas of the mineral precipitates as a function of the electron beam irradiation time. The amount of precipitates increases with the electron beam irradiation time because the water is depleted by the electron beam radiolysis. Although the mean radius \( r = (S/n)^{1/2} \), where \( S \) is the projected area of each precipitate) of the precipitates is constant (7.5 and 13.2 nm at 101 and 208 e⁻/nm² s, respectively, in the inset of Figure 1D), the total projected areas of all precipitates increase linearly with the number of precipitates (Figure 1D). This indicates that mineral nuclei spread during electron beam irradiation. Mineral precipitation by the spread of nuclei is known to occur in a highly supersaturated solution using conventional synthesis methods.

To investigate the structure of the mineral nuclei, we observe a single mineral nucleus at high resolution. Although sodium, sulfur, and oxygen atoms are relatively light atoms, Figure 2A shows a clear high-resolution TEM (HR-TEM) image, and the corresponding fast Fourier transformation (FFT) image (Figure S6) shows the hexagonal structure of the sodium sulfate crystal (Figure 2B). Figure 2C shows a HR-TEM image of the precipitates on the FLG edge. It indicates that sodium sulfate nanocrystals nucleate on graphene surfaces to lower the interfacial energy, which also occurs in the conventional SiN liquid cell. The 2D shape of the nuclei has a small thickness of less than 5 nm (Figure S4), which is formed in a highly supersaturated solution using conventional synthesis methods.

We monitor real-time mineral grain growth in the aqueous solution. Figure 3A shows time-serial HR-TEM images of grain boundary migration (Movie S4). The motion of the gold nanoparticles indicates that there is a liquid environment. Initially, the precipitate consists of four grains G1, G2, G3, and G4 with different crystallographic orientations (Figure S7) and five grain boundaries (GB12, GB23, GB123, GB13, and GBG23). In the aqueous solution, water molecules can be structurally incorporated in mineral grain boundaries with a highly disordered lattice. In the grain boundaries incorporating water molecules, unstable adatoms in convex surfaces dissolve and reprecipitate on concave surfaces (Figure S8), and hence the grain boundaries migrate, which does not occur in a dried environment (Figure S9). When the grain boundaries are mobile, at a triple junction where G1, G2, and G3 meet, a local mechanical equilibrium is established and the requirement for equilibrium is \( \gamma_{12}/\sin \alpha_{12} = \gamma_{13}/\sin \alpha_{13} = \gamma_{23}/\sin \alpha_{23} \), where \( \gamma_{12}, \gamma_{13}, \) and \( \gamma_{23} \) are the relative boundary energies of the GB12, GB13, and GB23, respectively, and \( \alpha_{12}, \alpha_{13}, \) and \( \alpha_{23} \) are the angles between GB12, GB13, GB123, and GB23 and GB13 and GB23, respectively. Since \( \sin \alpha_{12} \) is larger than \( \sin \alpha_{13} \) and \( \sin \alpha_{23} \), \( \gamma_{12} \) should be relatively higher than \( \gamma_{13} \) and \( \gamma_{23} \). Thus, the GB12 migrates toward the center of curvature to reduce the \( \gamma_{12} \). The GB23 also migrates for the same reason. At 30.1 s, the migration of GB12 and GB23 leads to the formation of a three-sided G2, and GB12 stops migration when GB13 is mechanically stabilized by GB14. At 54.6 s, the G2 eventually disappears due to the migration of curved GB23. It is noted that the migration speed of grain boundaries is independent of the presence or absence of gold particles, indicating that the gold particles do not affect the grain growth of minerals.

Figure 3B shows the FFTs corresponding to HR-TEM images of G1 and G4 during grain growth. For 10.3 s, G1 and G4 rotate 7 and 1°, respectively, which indicates that grains rotate separately during grain growth. In nature, grain boundary migration and grain rotation occur in aqueous solution to reduce the energy of dislocations, point defects, and grain boundaries.

Figure 4A–D shows the schematics representing a proposed mineral precipitation mechanism by electron beam irradiation. The sodium and persulfate ions are initially solvated by water molecules (Figure 4A), and electron beam irradiation yields spurs, spaced according to the electrons’ mean free path (Figure 4B). In the spurs, water molecules are excited or ionized, producing radicals, ions, and molecular species such as \( e^− \) (the hydrated electron), \( \text{H}_3\text{O}^+, \text{H}^+, \text{OH}^−, \text{H}_2\text{O}_2, \text{H}_2\text{O}_2^−, \text{O}^−, \text{H}_2\text{O}_2, \text{H}_2\text{O}_2^− \). Radiolytic \( \text{H}_2 \) and \( \text{O}_2 \) contribute to the formation of bubbles (Figure S10). By electron beam radiolysis in the spurs, sodium and persulfate ions are dehydrated and persulfate ions decompose to sulfate ions. In the undersaturated solution, dehydrated sodium and sulfate ions are solvated again by water molecules. As the electron beam dose accumulates, the amount of water decreases and hence mineral ions become saturated in solution. When mineral ions are supersaturated, clusters of dehydrated mineral ions are generated in the spurs and form stable nuclei (Figure 4C). Under continuous electron beam irradiation, the nuclei spread out and grains grow by grain boundary migration and grain rotation (Figure 4D).

CONCLUSION

In conclusion, we present a new in situ approach for high-resolution imaging of mineral precipitation using the GLC-TEM method. We show nucleation and growth processes of sodium sulfate, which is difficult to achieve with traditional thick SiN liquid cells because sodium sulfate crystals are thin and composed of light atoms. Since graphene is a relatively chemically inert semimetal, the graphene windows of our cell are not expected to appreciably influence the charge distribution and spur formation in solution other than to confine it to a small volume. Our findings faithfully mimic mineral precipitation processes occurring in nature, such as mineral nucleation and mineral grain growth in highly supersaturated solution. This work goes beyond the specific material system studied here or methodology applied here and hence will likely benefit precipitation studies of many different naturally occurring minerals, such as calcium carbonate and sodium chloride.

METHODS

GLC Fabrication. FLG sheets were synthesized on Cu foil (99.8% Alfa Aesar, Ward Hill, MA) with flowing 200 sccm \( \text{H}_2 \) and 18 sccm \( \text{CH}_4 \) gas at 1050 °C for 1 h using atmospheric pressure chemical vapor deposition. After the supporting Cu foil was etched away with a \( \text{Na}_2\text{S}_2\text{O}_8 \) solution (concentration of 0.1 g of \( \text{Na}_2\text{S}_2\text{O}_8 \)/1 mL of water)
overnight (Figure S2A), the FLG sheet was rinsed with deionized water (Figure S2B). The free-floating FLG sheet was intact without any supporting substrates and clearly visible optically under normal illumination condition (see optical image in Figure S2B), while a single graphene sheet was visible optically by using evanescent waves due to its negligible refractive index.20 The FLG sheet was moved to target solutions via a van der Waals interaction where the solution was floating FLG sheet was intact without fraction patterns were acquired using a JEM-2010 LaB6 instrument (JEOL Ltd.) at 80 kV. Dark-field TEM imaging was performed by selecting diffracted spots from precipitates in diffraction mode with a 0.5 nm−1 objective aperture. HR-TEM imaging was conducted with the TEAM 1 operated at 80 kV at the Molecular Foundry. Energy-disperse X-ray spectroscopy was performed in the diffraction mode using an Oxford INCA energy-dispersive X-ray detector, installed in the Philips CM200/FEF operated at 200 kV at the Molecular Foundry.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsnano.5b04064.

Additional figures and movie descriptions (PDF)

Movie 1 (AVI)
Movie 2 (AVI)
Movie 3 (AVI)
Movie 4 (AVI)

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Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

We thank C. Song at the Molecular Foundry for experimental assistance with TEM, and S. Nguyen at U.C. Berkeley for helpful discussions. J.M.Y., Q.Z., J.C., and A.Z. acknowledge support from the Director, Office of Energy Research, Basic Energy Sciences, Materials Sciences and Engineering Division, of the U.S. Department of Energy under Contract DE-AC02-05CH11231 within the SP2-Bonded Materials Program and The Molecular Foundry, which provided for construction of the GLC and TEM characterization, respectively; the Office of Naval Research under Grant N00014-12-1, which provided for graphene growth; the NSF under Grant DMR-1206512, which provided for development of graphene transfer methods; and DTRA Grant HDTRA1-13-1-0035, which provided postdoctoral support.

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