Imaging, understanding, and control of nanoscale materials transformations

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The development of liquid cells for transmission electron microscopy has enabled breakthroughs in our ability to follow nanoscale structural, morphological, or chemical changes during materials growth and applications. Time-resolved high-resolution imaging and chemical analysis through liquids opened the opportunity to capture nanoscale dynamic processes of materials, including reaction intermediates and the transformation pathways. In this article, a series of work is highlighted with topics ranging from liquid cell developments to in situ studies of nanocrystal growth and transformations, dendrite formation, and suppression of lithium dendrites through in situ characterization of the solid–electrolyte interphase chemistry. The understanding garnered is expected to accelerate the discovery of novel materials for applications in energy storage, catalysis, sensors, and other functional devices.
self-assembly of nanoparticles, electrode–electrolyte interfaces, catalysis, and beyond. Other areas of study include nanoscale chemical reactions, solid–liquid–gas interfaces, catalysis, and biological samples.

This article highlights the liquid cell development and its associated work with a focus on nonspherical nanoscale materials, including the growth and transformations of PtFe nanowires,\(^2\) facet development of Pt nanoparticles,\(^1\) reversible giant deformation of PbSe nanocrystals during superlattice transformations,\(^1\) and Li dendrites formation.\(^22\) Not only do these material systems have unique anisotropic properties,\(^23,24\) they are also ideal systems for exploring nonequilibrium behaviors at the nanoscale, in which a small perturbation can drive them out of equilibrium. To this end, direct observations with high spatial and temporal resolution are necessary for yielding critical insights to a range of related questions. These include how a system responds to perturbations, and what are the governing rules for a system to evolve away from its initial thermodynamic equilibrium. Coupled with theoretical calculations, these fundamental studies foster the discovery of novel materials and promote their real-world applications in functional devices.

This article is adapted and extended from the presentation I gave at the MRS Medal Award session of the 2019 MRS Fall Meeting. It is my hope that researchers from materials science to physics, chemistry, and engineering, find the materials presented informative and useful.

**Development of liquid cells allowing high-resolution imaging and chemical analysis**

The challenges of liquid phase TEM imaging arise largely from the liquid cell being too thick to achieve high resolution. Previously, nitrocellulose thin films,\(^25\) aluminum thin films,\(^26\) and thick SiN\(_x\) membranes more than 50 nm in thickness were typically used; liquid cell fabrication features the stacking or gluing two chips together. Making ultrathin freestanding membranes and controlling the liquid thickness are some of the key attributes to recent developments. These were achieved by using low-stress silicon nitride, SiN\(_x\) (\(x < 4/3\)), 25 nm in thickness on a thinner silicon wafer (100-μm-thick, as compared to the standard 500-μm-thick standard wafer). Even thinner SiN\(_x\) membranes have been developed and used more recently. A unique problem associated with the use of thin wafers is that they are too fragile to be processed using standard nanofabrication processes. To overcome this problem, one solution is to first bond a thin wafer to a standard 500-μm wafer with photoresist, thus batch fabrication of liquid cells with automated lithography patterning can be accomplished (Figure 1a). To control the liquid thickness inside the cell, a thin In spacer (70–100 nm) may be deposited by thermal evaporation on the bottom chip of the cell and bottom chip can be bonded with the top chip. The thin wafer design minimizes the attenuation of x-ray signals in EDS (Figure 2). The self-contained liquid cells (Figure 1c) with dimensions typically of 2.6 mm × 2.6 mm and 200 μm in thickness fit well into a standard TEM holder.\(^6,7\)

Electrochemical liquid cells may be fabricated by incorporating patterned electrodes (Figure 1e–f). Various metals (e.g., Au, Ti, and Pt) may be used as the electrode materials,\(^3,16,22,27\) and they may be electrically connected to the electric pads on the TEM holder with Au or Al wires. Here, the TEM holder must be specially designed to enable electrochemical biasing. EDS chemical analysis is possible with these thin electrochemical liquid cells. For most of the studies on deposition of alkali metals on the electrode, high-resolution imaging is possible, but low-dose imaging is often necessary to avoid electron-beam damage, as will be elaborated later.

Superior high-resolution imaging has been accomplished with a range of newer membranes and 2D materials ranging from carbon films\(^10,11\) to MoS\(_2\) combined with graphene\(^12\) (Figure 1g–h); lipid nanoreactors\(^28\) (Figure 1i), on the other hand, offer unique opportunities for study of controlled nanochemistry and nanoscale chemical reactions.

**In situ high-resolution tracking of colloidal nanoparticle growth and transformations**

The ability to directly monitor colloidal nanoparticle transformations with high resolution opens the opportunities to uncover their growth and transformation mechanisms. Many nanoparticles show no distinct facets (Figure 2a–c). A group of nanoparticles with certain shape and facets, such as nanowires, nanosheets, and nanocubes, has attracted significant interests due to their shape-dependent properties. Synthesis of nanoparticles with controlled shapes has been achieved recently.\(^29\) However, our understanding of the shape-control mechanisms remains limited in many nanoparticle systems.

The conventional way of investigating nanocrystal growth is by *ex situ* measurements (e.g., by periodically taking out nanoparticles from the solution during synthesis, drying them and examining them under TEM). The drawbacks of *ex situ* measurements are the possible structural changes during sample preparation and the need to infer crystal growth between the examining points. Direct imaging during synthesis resolves this problem, as we can track the trajectories of individual nanoparticle growth and capture the intermediate states on the fly.

Liquid cell imaging can provide valuable insights into the growth mechanisms of 1D nanowires from quantum-dots as the building blocks. Figure 3a shows the sequential images of the growth of a single PtFe nanorod from a molecular precursor solution. Nanoparticles were formed initially. They subsequently attached together to form a nanowire, followed by atomic-level structural rearrangement between the nanoparticles eventually leading to the formation of a single-crystal PtFe nanowire.\(^2\) The role of ligands in the nanoparticle shape evolution is revealed through theoretical simulation coupled
with a series of control experiments.\textsuperscript{2,21,30} With advanced image analysis, it is possible to capture the interaction energetics during nanoparticle self-assembly.\textsuperscript{20}

Pt nanocube formation has been studied using liquid cell TEM (Figure 3d–f).\textsuperscript{13} For the shape evolution of nanocrystals, Wulff construction\textsuperscript{31,32} is often used to explain their growth. For instance, high-energy facets grow faster than the low-energy facets. In other words, the fast growing facets are eventually replaced by low-energy facets.\textsuperscript{29,33} Surfactant ligands, however, modify the facet energy, thus influencing the relative growth rate of different facets and altering the final shape of a nanocrystal.\textsuperscript{34,35} An \textit{in situ} study of Pt nanocube growth demonstrated that the facet development can deviate from the Wulff construction. As shown in Figure 3e, the growth rates of all low-energy index facets, \{100\}, \{110\}, and \{111\}, are similar until the \{100\} facets stop growing at about 2.5 nm \((d_c)\). The \{110\} facets continue to grow until an edge is formed, whereas the growth of \{111\} facets fills the corners of the nanocube. Density functional theory (DFT) calculation revealed that the different mobilities of the ligands on the different facets play an important role here. Specifically, the mobility of ligands is several orders of magnitude lower on the \{100\} than the \{111\} facets, thus retarding the growth of the \{100\} facets. A selective facet-arrested shape-control mechanism, mediated by the ligand mobility on different facets, is thus proposed from the complementary theoretical calculations and experiments. Interestingly, the growth of nanocrystals was found to be largely stochastic when they are smaller than a critical size \((d_c = 2.5 \text{ nm})\), below which no obvious differences in the growth rate could be observed.

Drastic shape evolution was also observed in the growth of cobalt oxide or cobalt nickel oxide nanoparticles (Figure 3g–i). Nanoparticles are formed first from a precursor solution and they transform into nanosheets after they reach a critical size. Different from the previously reported 2D nanosheet formation through oriented attachment of primary 3D nanocrystals\textsuperscript{36,37} or the “soft template”-assisted growth of 2D nanostructures,\textsuperscript{38–40} the 3D-to-2D transformation involves thinning down at the edges of the nanoparticles first followed by the formation of a uniform single-crystal nanosheet.
Individual PbSe nanocrystals were observed to experience shape changes during superlattice transformations. When oleate ligands were removed from the nanoparticle surface by ethylene glycol (EG), negligible changes of individual nanocrystals were observed. In contrast, a small amount of ethylenediamine (EDA) with EDA:EG = 1:1000 in the ligand exchange solution induced large shape changes (up to 40%) as the PbSe nanoparticle interacting with neighboring particles. The deformation of individual PbSe nanocrystals introduces defects in the final superlattice, for example, lower degree of connectivity and higher degree of misorientation in the superlattice formed by ligand exchange with EDA than that formed with pure EG (Figure 4c–d). Further in situ study resolved the necking formation mechanism at the atomic level and revealed the origin of defects in PbSe superlattice transformations.

In situ study of dendrite formation at the nanoscale

Understanding nanodendrite pattern formation

Dendritic materials are widely found both in nature and through laboratory synthesis. Their formation mechanisms have been explored theoretically. Using liquid cell TEM with fine control of the growth conditions and advanced data collection and image analysis, the growth mechanism of iron oxide nanodendrites has been studied with a focus on validating the applicability of available theories for nanoscale systems (Figure 5a–b). Tracking the trajectories of their morphology development reveals the relationship between the tip curvature and growth rate, tip splitting relative to instabilities, and the effects of precursor diffusion and depletion on the morphology evolution. Interestingly, the dendritic morphology evolution observed during the growth of iron oxide nanodendrites was remarkably consistent with existing theoretical predictions, despite occurring at the nanoscale.

Interests in metal dendrites stem from the fact that their growth is a primary cause for short-circuit failures in rechargeable batteries and other electronic devices. In a study of the crystallization and morphology evolution of lead dendrites, we observed dendrite growth through tip splitting and dissolution of dendrites with mass loss when electric biasing was reversed (Figure 5c). The electrochemically reduced lead atoms form nanoparticles on the electrode. They subsequently aggregate into clusters and eventually form a dendritic structure. In situ study of Li dendritic growth with liquid phase TEM was challenging because of the use of air/moisture-sensitive electrolyte, and alkali metals being electron-beam sensitive. Nevertheless, the deposition and dissolution of Li dendrites were observed in situ for the first time using a commercial electrolyte for Li-ion batteries (Figure 5d–f). More discussion is provided in this article.

Understanding and controlling of Li dendrite growth through in situ chemical analysis

Alkali metal (e.g., Li and Na) plating and stripping have been studied using liquid cell TEM with an emphasis on analysis of the role of solid–electrolyte interphase (SEI). As a passivation interfacial film formed from the reduction of the electrolyte, SEI plays an important role in the cycle stability of Li-ion batteries. Previous efforts on the study of SEI were primarily through ex situ FTIR, Raman spectroscopy, atomic force microscopy, x-ray photoelectron spectroscopy, and scanning tunneling microscopy. In situ characterization has been attempted using x-ray scattering and neutron reflectometry avoiding complications due to sample exposure to air and moisture. In situ electron microscopy imaging with nanoscale chemical analysis complements these studies. Liquid phase TEM observations and nanobeam diffraction of SEI on the electrode revealed that the SEI layer contained LiF nanocrystals distributed in an amorphous matrix in contrast to the previous understanding.
of a denser layer of inorganic components adhering to the electrode with a porous organic outer layer exposing to electrolyte.

Since the Ti electrode is not reactive with the LiPF6/EC/DEC electrolyte, the formation of SEI should be stepwise from preferential reduction of certain electrolyte components. In the contrary, the Li metal electrode reacts with the electrolyte directly, leading to instantaneous SEI formation. For this reason, the reduction of electrolyte components should be indiscriminate to all species.

Further advances of in situ study of electrochemical liquid cells allowed the mapping of spatially resolved SEI chemistry on individual lithium nanograins. The observation revealed that Li dendrites can be suppressed by a cationic polymer coating on the electrode (Figure 5e–h), thus uncovering a potential path to Li dendrite suppression.

Electron-beam effects

The high-energy electron beam is known to interact with liquids, causing molecular dissociation, ionization, and other chemical reactions. Concerns about sample damage due to electron-beam exposure have been discussed throughout the history of electron microscopy development. Recently, there have been an increased number of studies on this problem especially for beam-sensitive materials in liquids. There have also been reports about simulation of radiolysis-induced damages of water and organic molecules. Real-time time-dependent density functional theory (rt-TDDFT) calculations revealed the strong competition among different dissociation paths in C2H6O2 radiolysis. Obviously, blocking the dissociation paths during electron microscopy imaging may reduce the electron-beam damage, even though radiolysis damage to a liquid solution can be rather complex. Many factors
must be considered, including the electron-beam energy, molecules being studied, and their competing chemical reactions.

While no universal solution exists, lowering the electron-beam current density reduces the electron-beam effects. For example, low-dose imaging has been applied to electrochemical studies\textsuperscript{3,4,16,17,22} and no discernible reactions should be observed before applying an electric bias. In the study of nanocrystal growth and transformations, strong electron-beam intensity was often needed, in which case, control experiments would become critical to observing and isolating the beam damage effects.\textsuperscript{64}

Understanding of materials transformations accelerates materials discovery

With the fundamental understanding of nanoscale materials growth and transformations garnered from \textit{in situ} studies, materials may be designed for achieving special functions. For example, a series of functional materials was prepared through laser synthesis involving nonequilibrium phonon-induced reactions.\textsuperscript{65–67} Nanoporous metal–organic composites were designed and prepared recently, showing nearly 100\% selective of CO production over H\textsubscript{2} generation from CO\textsubscript{2} reduction reaction. Tandem catalysts prepared by incorporating Rh or Ag nanoparticles into the metal–organic composites, on the other hand, produced formic and acetic acids. Remarkably, laser synthesis significantly shortens the growth processes, from 24+ h with the conventional solvothermal method to 30 min–2 h.

Conclusions

With the development of \textit{in situ} liquid cell TEM, unprecedented advances in high-resolution imaging and chemical analysis have been made for a wide range of

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\caption{Liquid phase transmission electron microscope (TEM) study of PbSe nanocrystal deformation during superlattice transitions. (a) Reversible large deformation of PbSe nanocrystals (No. 11). Scale bar = 5 nm. (b) Quantification of the deformation. (c) Low- and high-magnification TEM images showing that nearly perfect PbSe superlattice was achieved with ethylene glycol (EG) as the ligand exchange solution. Scale bars = 20 nm (left) and 5 nm (right). (d) Comparing to (c), defects were observed with ethylenediamine (EDA) in the ligand exchange solution. Scale bars = 20 nm (left) and 5 nm (right). Reprinted with permission from Reference 11. © 2019 AAAS.}
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\caption{In situ study of dendrite growth allowing fundamental understanding of the dendrite formation at the nanoscale and controlling of dendrite growth in functional devices (i.e., Li-ion batteries). (a) Iron oxide dendrite formed in a liquid cell overlapping with the contour map from computer tracing of the dendrite evolution. (b) Large data analysis revealed tip split arising from instabilities. Adapted with permission from Reference 50. © 2018 American Chemical Society. (c) Electrodeposition and dissolution of Pb dendrites in an electrochemical liquid cell. Reprinted with permission from Reference 16. © 2013 Nature Publishing Group. (d) Electrodeposition and dissolution of Li dendrites in an electrochemical liquid cell. Adapted with permission from Reference 22. © 2018 American Chemical Society. (e–h) Suppressing Li dendrite formation with a cationic polymer film (PDDA) coated on the electrodes inside the liquid cell. (e) Li nanogranular growth was achieved on the electrode coated with PDDA. (f) Li dendrite growth was achieved on a bare electrode without polymer coating. (g) Energy-dispersive x-ray spectroscopy (EDS) chemical mapping of a Li nanograin with SEI on the surface. (h) EDS line scan profile showing the solid–electrolyte interphase (SEI) chemistry. (e–h) Reprinted with permission from Reference 4. © 2020 Royal Society of Chemistry.}
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problems involving nanoscale materials growth and transformations. These advances opened many opportunities to resolve dynamic processes of nanoscale materials that were unreachable previously. This article highlights some examples of liquid phase TEM studies of nanoscale materials, from one-dimensional nanocrystals to faceted nanocrystals, 2D nanosheets, and superlattice transformations. Other applications discussed include nanodendrite formation and Li dendrite suppression by resolving nanoscale structure and chemistry of SEI. Clearly, the advances in liquid cell electron microscopy accelerate materials discovery.

There are tremendous future opportunities in liquid phase TEM as the liquid cell development is tuned for specific experiments with better control of reactions under various stimuli. And, as advanced imaging with fast data collection has become widely available, large data processing and data interpretation are highly integral components of the studies. Liquid phase TEM will continue to attract great interest in various research fields from materials science to chemistry, physics, and biology.

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Data availability

Data are available in the online version of this paper. Data that support the findings of this study are available from the author upon reasonable request.

Conflict of interest

The author declares no competing financial interests.

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450 • MRS BULLETIN • VOLUME 46 • MAY 2021 • mrs.org/bulletin

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