Characterization of Sulfur and Nanostructured Sulfur Battery Cathodes in Electron Microscopy Without Sublimation Artifacts

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Abstract: Lithium sulfur (Li–S) batteries have the potential to provide higher energy storage density at lower cost than conventional lithium ion batteries. A key challenge for Li–S batteries is the loss of sulfur to the electrolyte during cycling. This loss can be mitigated by sequestering the sulfur in nanostructured carbon–sulfur composites. The nanoscale characterization of the sulfur distribution within these complex nanostructured electrodes is normally performed by electron microscopy, but sulfur sublimes and redistributes in the high-vacuum conditions of conventional electron microscopes. The resulting sublimation artifacts render characterization of sulfur in conventional electron microscopes problematic and unreliable. Here, we demonstrate two techniques, cryogenic transmission electron microscopy (cryo-TEM) and scanning electron microscopy in air (airSEM), that enable the reliable characterization of sulfur across multiple length scales by suppressing sulfur sublimation. We use cryo-TEM and airSEM to examine carbon–sulfur composites synthesized for use as Li–S battery cathodes, noting several cases where the commonly employed sulfur melt infusion method is highly inefficient at infiltrating sulfur into porous carbon hosts.

Key words: Cryo-TEM, airSEM, lithium sulfur, battery, energy

INTRODUCTION

Lithium sulfur (Li–S) batteries have the potential to provide greater energy storage density at lower cost than current lithium ion batteries. One of the main challenges to improving the performance of elemental sulfur cathodes for Li–S batteries is dissolution and loss of sulfur, in the form of polysulfides, to the electrolyte during battery operation (Bruce et al., 2011). Recent research has focused on electrodes which attempt to sequester sulfur in nanostructured host materials, most prominently porous carbons, to prevent the loss of sulfur and associated capacity reduction as the battery is cycled (Ji et al., 2009; Jayaprakash et al., 2011; Wang et al. 2011; Xiao et al., 2012; Seh et al. 2013; Zheng et al., 2013; Song et al., 2014; Zhao et al., 2014; Sahore et al. 2015; Werner et al., 2015). These electrode materials are typically referred to as carbon–sulfur composites. Accurate characterization of the distribution of sulfur in these composites, over 100 nm to sub-nanometer length scales, is critical for developing an understanding of how the level of infiltration of sulfur into the host material relates to battery performance, which will aid in the design of more durable, high energy density Li–S batteries (Ma et al. 2015).

Electron microscopy offers both direct imaging and spectroscopic techniques for characterization of battery electrodes. For reliable characterization of a material using electron microscopy, it is essential that the sample is not altered by conditions in the microscope. However, elemental sulfur readily sublimes under high-vacuum conditions similar to those of an electron microscope sample chamber. Sulfur sublimation under high vacuum has been observed in vacuum chamber experiments on bulk sulfur (Nash, 1987), and can be predicted from sulfur’s measured vapor pressure (Ferreira & Lobo, 2011), but the sulfur battery community has only recently begun to become aware of the problem of sulfur sublimation in electron microscopy (Raij et al., 2014). As an illustration of this challenge, Figure 1a shows the vapor pressure curve of sulfur, with the conditions of an FEI Tecnai F20 (FEI Company, Hillsboro, OR, USA) transmission electron microscopy (TEM) sample chamber (8.8 × 10⁻⁸ Torr at ~18°C room temperature) indicated. Figures 1b–1g show the result of placing a sample of ball-milled sulfur particles into the microscope under these conditions. The sulfur is observed to sublimate at a rate of ~1 monolayer of sulfur atoms per second, leaving behind only a small residue of “super-sublimated” polymeric sulfur, which remains relatively stable under vacuum. The measured sublimation rate and residual product are fully consistent with previous macroscopic experiments (Nash, 1987).

In nanostructured sulfur composites, sublimation effects in high vacuum may be very severe. In TEM, for example, exposed nanoscale features of the sulfur distribution in a sulfur...
battery cathode composite would disappear within minutes at most. Only super-sublimated polymeric sulfur residue (Fig. 1e), and sulfur encapsulated by another material lacking large enough pores for sulfur to escape (Zhou et al., 2014; Kim et al., 2015), will remain present under vacuum.

In scanning electron microscopy (SEM), the sample is typically held at a higher pressure than in TEM. However, recent studies have observed that even in the vacuum of an SEM sample chamber (~10^{-6} Torr), sulfur sublimation artifacts occur, including the redistribution of sublimated sulfur into the pores of nearby carbon particles. The study concluded that characterization of sulfur/carbon composite materials by vacuum-based methods, including SEM, is very challenging, and results might be misleading (Raiβ et al., 2014). Sulfur sublimation and the resulting artifacts, including sulfur redistribution, present a serious problem for accurate characterization of the inherent sulfur distribution in battery cathodes in electron microscopy. This may be impeding the scientific community’s efforts to gain a systematic understanding of how sulfur distribution in different carbon–sulfur composites relates to their performance in batteries. More reliable alternative techniques for sulfur characterization that suppress sublimation are clearly needed.

The sulfur vapor pressure curve (Fig. 1a) indicates that sulfur has an equilibrium vapor pressure of ~6 × 10^{-7} Torr at 18°C. This means that at 18°C, a sulfur particle will sublimate until the partial pressure of sulfur surrounding the particle reaches 6 × 10^{-7} Torr. If a sample chamber is pumped to too low a pressure, sulfur will not be able to reach equilibrium, and will sublimate continually. To avoid sublimation artifacts, either the ambient pressure during imaging must be increased to a level much greater than sulfur’s equilibrium vapor pressure, or the ambient temperature must be reduced to much less than 18°C. In this paper, we demonstrate characterization of sulfur and nanostructured carbon–sulfur composite materials in vacuum at low temperature using cryogenic transmission electron microscopy (cryo-TEM), and at atmospheric pressure at room temperature using scanning electron microscopy in air (air-SEM). Sulfur sublimation artifacts are not observed using either technique. Our results demonstrate that sulfur infiltration by melt infusion is significantly more efficient in activated porous carbons than in nonactivated porous carbons, carbon nanotubes (CNT), and hollow carbon spheres, all of which have recently been investigated as Li–S battery electrode materials. Adoption of cryo-TEM, airSEM, and other similar techniques, for more reliable characterization of the sulfur distribution in different carbon–sulfur composites will enable scientists to observe the inherent sulfur distribution in their composite materials, providing important insight and feedback to guide the design of improved Li–S batteries.

**EXPERIMENTAL**

Cryo-TEM is a well-established method for imaging hydrated biological samples that cannot be exposed to vacuum at room temperature (Dubochet & McDowall, 1981; Adrian et al., 1984; Kourkoutis et al. 2012). Cryogenic cooling of sulfur cathode samples inside the microscope was achieved using a Gatan model 626 cryo-holder (Gatan Inc., Pleasanton, CA, USA), with liquid nitrogen as the cryogen. The cryo-TEM loading method for sulfur cathode samples is simpler than for biological samples, since the sulfur need only be cooled before loading into the microscope column, whereas biological samples must be vitrified. Carbon–sulfur composite particles were dispersed from an ethanol solution onto TEM grids, which were then allowed to dry in air.
The cryo-holder was cooled so that the tip temperature was approximately −173°C. The sample TEM grids were loaded into the cryo-holder under nitrogen gas near liquid nitrogen temperature. An FEI Tecnai F20 STEM/TEM (FEI Company, Hillsboro, OR, USA) equipped with cryogenically cooled beryllium blades, and operated in scanning TEM (STEM) mode at 200 kV was used to image the samples. An Oxford Instruments XMAX detector (Oxford Instruments PLC, Tubney Woods, Oxfordshire, UK) was used for X-ray energy dispersive spectroscopy (XEDS), and a Gatan 865 HR-GIF spectrometer (Gatan Inc., Pleasanton, CA, USA) was used for electron energy loss spectroscopy (EELS) acquisition.

Experiments in air were performed using a B-nano airSEM (B-nano Ltd., Rehovot, Israel), operated at 30 kV. The airSEM is a relatively new design of electron microscope (a schematic diagram of an airSEM is shown in Supplementary Fig. S1), which enables characterization of samples in air, with no sample vacuum chamber (Nguyen et al., 2013, 2016; Solomonov et al., 2014; Vidavsky et al. 2014). Carbon–sulfur composite particles for analysis in airSEM were dispersed onto TEM grids from an ethanol solution and allowed to dry in air, in the same manner as for cryo-TEM. Sample TEM grids were placed directly on top of an airSTEM detector (Nguyen et al., 2014, 2016; Han et al., 2015), which was mounted on an optical slide. A white light optical reflectance microscope (Olympus, Centre Valley, PA, USA) was used both for optical imaging, and to set the height of the sample to ~50 μm below the electron window, the optimal working distance for airSEM imaging. XEDS maps were acquired using a Bruker XFlash 6160 detector (Bruker Corporation, Billerica, MA, USA).

Pure sulfur particles were prepared for analysis by ball-milling sulfur flakes. Carbon–sulfur composites synthesized with ordered mesoporous carbon structures GDMC-15-1600°C and aGDMC-15-10h, were obtained from block copolymer co-assembly and heated to 1,600°C to remove any intrinsic microporosity. The high-temperature treatment yielded low oxygen content and surface functionalization (hydrophobic surface). Carbon aGDMC-15-10h underwent activation by heating in CO2 at 950°C for 10 h. Sulfur infiltration by melt infusion was attempted at a sulfur:carbon ratio of 1:1 by weight. Mesoporous hollow carbon sphere–sulfur composites, MHCS-S, were prepared by an organic hard template method. In a typical synthesis, size tunable mono-dispersed melamine-formaldehyde (MF) resin sphere templates (0.5 g) were synthesized by the in situ polymerization method reported by Xie et al. (2008). These were then suspended in a 50 mL water/ethanol (volume ratio 3/1) solvent containing 0.5 g of Triton X-100 surfactant (Dow Chemical Co. Midland, MI, USA). Formaldehyde solution (1.0 mL), and resorcinol (0.75 g) were then added to the reaction solution, and stirred for 30 min at 60°C. The solution was subsequently heated for 2 h at 75°C. The solid product was recovered by filtration and air-dried at 90°C for 8 h. MHCS were formed by carbonization of the as-made resorcinol–formaldehyde encapsulated MF spheres at 950°C for 2 h under flowing nitrogen gas with a heating rate of 3°C/min. Sulfur infiltration was attempted by melt infusion at a sulfur:carbon ratio of 7:3 by weight.

**RESULTS AND DISCUSSION**

**Sulfur Characterization in Cryo-TEM**

In order to investigate whether sublimation effects were still apparent at cryogenic temperatures, ball-milled sulfur particles were prepared in an identical manner to those in Figure 1, and imaged by cryo-TEM at a temperature of approximately −173°C. A time series of cryo-TEM images from a sulfur particle is shown in Figure 2. No change in the morphology of the particle was observed over a 5-h period, in stark contrast to the sulfur particle imaged by standard room temperature TEM shown in Figure 1, which sublimated and disappeared in under 40 min. The cryogenically cooled sulfur remained stable enough under the electron beam to allow imaging in annular dark field (ADF) cryo-scanning TEM (cryo-STEM, Fig. 2d) and XEDS mapping (Fig. 2e). The suppression of sulfur sublimation by cryogenic sample cooling demonstrates that cryo-TEM is a viable method for both imaging and spectroscopic characterization of composites containing elemental sulfur such as carbon–sulfur nanocomposites for advanced lithium batteries.

Having established that sulfur sublimation is suppressed at cryogenic temperatures, we used cryo-TEM to analyze the sulfur distribution within three different carbon–sulfur nanocomposite materials recently investigated for Li–S battery electrodes. The first two samples, GDMC-15-1600°C and aGDMC-15-10h, are examples of ordered carbon nanonetworks (Werner et al., 2015). These have attracted interest as sulfur battery cathode materials due to the potential of sequestering sulfur in the pores of a tortuous carbon network. GDMC-15-1600°C and aGDMC-15-10h were prepared in an identical manner except that, before sulfur melt infusion, aGDMC-15-10h underwent an additional activation procedure to substantially increase its microporosity and associated surface area, whereas the nonactivated carbon, GDMC-15-1600°C, exhibits very little microporosity.
section). The third sample, CNT-S used a network of carbon nanotubes (CNT) as the host material for sulfur. CNT have attracted interest as Li–S battery electrodes due to the potential to sequester sulfur inside the body of the nanotubes (Fujimori et al., 2013). In each of our three samples, successful melt infusion would ensure that electrically insulating sulfur is in contact with electrically conducting carbon at the nanoscale. Cryo-STEM imaging and XEDS maps of the activated C–S composite aGDMC-15-10h (Fig. 3a), showed a high degree of infiltration of sulfur into the carbon following melt infusion. However, in the nonactivated C–S composite GDMC-15-1600°C we observed that a majority of the sulfur had not infiltrated the carbon, and remained external to the carbon particles (Fig. 3b). External sulfur particles typically had a diameter of several micrometers or larger. In reported battery performance tests, the activated carbon aGDMC-15-10h outperformed its nonactivated counterpart GDMC-15-1600°C in terms of initial capacity, capacity retention over cycling, and rate capability (Werner et al., 2015). This may be partly explained by our observation that sulfur infiltration was much more successful in the activated carbon aGDMC-15-10h than the nonactivated carbon GDMC-15-1600°C, because the external sulfur particles observed in GDMC-15-1600°C would be in poorer electrical contact with the carbon, and more exposed to dissolution and loss to the electrolyte in the form of polysulfides during battery cycling.

High-magnification STEM imaging and EELS spectra of sample CNT-S indicated that little or no sulfur had infiltrated into the CNT, which had remained hollow (Fig. 3c). XEDS mapping of sample CNT-S at low magnification showed that much of the sulfur in the composite remained external to the nanotube network after melt infusion, forming particles several microns in diameter (Fig. 3d), in a similar way to the sulfur in sample GDMC-15-1600°C. X-ray sum spectra from the datasets used to generate element distribution maps contain information about the relative quantities of carbon and sulfur in the field of view that are not evident from X-ray maps alone. The X-ray spectra used to generate the maps in Figure 3 are shown in Supplementary Figure 2. Each of these spectra show a strong sulfur peak relative to the carbon peak, indicating significant quantities of sulfur in the field of view.

Our results show that the degree to which sulfur infiltrates into carbon host particles can vary significantly depending on the porosity and structural characteristics of the carbon host, and how the carbon was prepared. In this case, sulfur infiltrated far more efficiently into the activated, high surface area carbon aGDMC-15-10h exhibiting both 15 nm mesopores and a large fraction of small micropores (<4 nm), than into the nonactivated carbon GDMC-15-1600°C, exhibiting only 15 nm mesopores, or into a network of CNT. In a related cryo-TEM study of sulfur infiltration into highly porous carbons synthesized by ice templation, sulfur was observed to have infiltrated far more efficiently into activated carbon 20-2-1.5-80S exhibiting both mesopores and micropores, than into nonactivated carbon 20-2-0-80S, exhibiting only mesopores (Sahore et al., 2016), a very similar trend to the results described above.

Supplementary Figure 2
Supplementary Figure 2 can be found online. Please visit journals.cambridge.org/jid_MAM.

Sulfur Characterization in airSEM.
An alternative approach to cryogenic cooling to avoid sulfur sublimation is to keep the sample at room temperature, but increase the pressure around the sample, for example, by imaging the sulfur at atmospheric pressure. The recently developed airSEM by B-nano enables correlative optical and electron microscopy of samples in air, with no vacuum specimen chamber, by using an electron transparent silicon-nitride window to separate the sample in air from the electron optics in vacuum (Solomonov et al., 2014; Vidavsky et al. 2014; Nguyen et al., 2016). Our cryo-TEM observations of external sulfur particles several micrometers in diameter in carbon–sulfur composite samples suggested that correlative optical–electron microscopy would be a useful method of high-throughput characterization of these materials. Optical microscopy can screen samples for large external sulfur particles as external sulfur (yellow) is optically distinct from...
carbon (black). Electron microscopy and XEDS can then be used to image carbon particles at higher magnification to analyze the degree of sulfur infiltration into the carbon, and investigate nanoscale features of the sulfur distribution.

As a demonstration of this technique, we used airSEM to investigate the distribution of sulfur in sample MHCS-S, a composite of sulfur and 3 µm diameter hollow carbon spheres with porous shells. These have attracted interest as candidate materials for Li–S batteries because of the potential for large quantities of sulfur to be sequestered in both the porous shell, and in the hollow interior of the sphere, though some recent studies have questioned whether melt infusion is actually successful at infiltrating sulfur into the hollow interior (Jayaprakash et al., 2011; He et al., 2013).

In our sample, sulfur infiltration was attempted by melt infusion at a ratio of 7:3 sulfur to carbon by weight before imaging, sufficient to ensure that both the pores in the shell wall, and the interior cavity could be filled if melt infusion was successful.

Particles of external sulfur were identified by optical microscopy (Fig. 4a). Closer inspection of a ~30 µm wide external sulfur particle (Fig. 4b) with airSEM, using a bright-field STEM detector positioned directly below the sample (Nguyen et al., 2014, 2016; Han et al., 2015), shows a morphology that suggests this particle was formed by freezing from a liquid state. XEDS mapping confirmed the identity of the particle as external sulfur (Figs. 4c, 4d, spectrum in Supplementary Fig. 4a). Clusters of carbon spheres were identified in the optical microscope (Fig. 4e) and imaged at high resolution in the airSEM (Fig. 4f). The spheres appeared to be unfilled in the airSEM images, suggesting that sulfur had not infiltrated into the sphere interior. This was confirmed by XEDS mapping (Figs. 4g, 4h, spectrum in Supplementary Fig. 4b). However, XEDS maps did show that sulfur had infiltrated the walls of the spheres, despite not infiltrating the hollow interior as had been intended.

Supplementary Figures 4a and 4b

Supplementary Figure 4a and 4b can be found online. Please visit journals.cambridge.org/jid_MAM.

Comparison of airSEM with Cryo-TEM

The spatial resolution set by electron optics in the airSEM will be more limited than for cryo-TEM. However, an advantage of the airSEM is that samples do not need to be cooled and placed in a vacuum chamber, meaning that airSEM can achieve greater sample throughput. AirSEM will be most suitable for use in conjunction with optical

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Figure 3. Cryo-scanning transmission electron microscope annular dark field (STEM ADF) images and X-ray energy dispersive spectroscopy maps of (a) Composite of sulfur and activated gyroidal mesoporous and microporous carbon aG3MC-15-10h, showing a high degree and homogeneity of sulfur infiltration (overlap in carbon and sulfur signals). b: Composite of sulfur and the same gyroidal mesoporous carbon G3MC-15-1600°C without prior activation and very low microporosity, showing pure sulfur external to the carbon host, indicating poor infiltration efficiency. c: Cryo-STEM image of carbon nanotubes after attempted sulfur melt infusion. Tubes remain hollow. Electron energy loss spectroscopy (EELS) spectrum from body of tube (red) on carbon support film shows little or no sulfur L-edge signal. EELS spectrum from an elemental sulfur particle shown for comparison in blue. d: Carbon nanotube sulfur composite (CNT-S), also showing most sulfur external to nanotubes.
microscopy for high-throughput characterization of samples using a broader field of view where image resolution of <5 nm is not required, whereas cryo-TEM will be most suitable for characterization of samples that require resolution of finer, nanometer scale features.

Limitations Due to Radiation Damage
Even after sublimation has been suppressed, radiation damage will ultimately limit the resolution at which sulfur can be characterized in electron microscopy. Electrical insulators, such as sulfur, are vulnerable to ionization damage (Egerton et al., 2004). Furthermore, sulfur molecules are weakly bound, making sulfur vulnerable to knock-on displacement damage (Chisney et al., 1988). When sulfur is imaged at relatively low magnification, as in the results presented in this paper, electron dose is spread over a wide area, and beam damage effects to sulfur will be minimal. However, at high magnification, electron dose becomes more concentrated and radiation damage will cause mass loss from sulfur particles. Radiation damage effects have been observed in Supplementary Figure 5, which is accompanied by further discussion of damage.

Supplementary Figure 5
Supplementary Figure S5 can be found online. Please visit journals.cambridge.org/jid_MAM.

Conclusions
The high vapor pressure of sulfur creates a serious risk that sublimation artifacts will lead to a mischaracterization of samples containing sulfur in standard high-vacuum electron microscopy. As a solution to this problem, we have demonstrated that sulfur sublimation is suppressed by cooling sulfur samples below the sulfur sublimation point in vacuum.
using cryo-TEM, or by using airSEM to image sulfur in air. Both techniques are able to detect features of the sulfur distribution across C–S nanocomposite materials that are extremely challenging to observe reliably in standard electron microscopy due to sulfur sublimation in vacuum and resulting artifacts such as sulfur redistribution. Most notably, in three of the four carbon–sulfur composite samples investigated in this paper, we have directly observed that sulfur infiltration by melt infusion is unsuccessful or inefficient, leaving much of the sulfur external to the carbon host material. It may be of interest for future studies to investigate whether inefficient sulfur infiltration by melt infusion occurs in other composite materials, and to what extent this affects battery performance.

In conclusion, we strongly recommend adoption of cryo-TEM, airSEM for more reliable characterization of sulfur and sulfur battery cathodes. Other techniques that may also be able to suppress sulfur sublimation artifacts could include environmental SEM, cryo-SEM, and environmental-cell TEM. Applying these techniques to Li–S batteries will allow researchers to reliably characterize the inherent distribution of sulfur in their composite electrodes, helping to guide the design of improved Li–S batteries.

ACKNOWLEDGMENTS

This project was supported by the Energy Materials Centre at Cornell, an Energy Frontier Research Centre funded by the US Department of Energy, Office of Science, BES Award DE-SC0001086; and the New York State Centre for Future Energy Systems (CFES), a joint Center for Advanced Technology between Cornell University and Rensselaer Polytechnic Institute, supported by the New York State, Empire State Development Division of Science, Technology and Innovation (NYSTAR), under contract number C100126. This work made use of the Cornell Centre for Materials Research Shared Facilities which are supported through the NSF MRSEC program (DMR-1120926). The authors thank John Grazul for assistance in the TEM facilities.

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