High capacity vertical aligned carbon nanotube/sulfur composite cathodes for lithium–sulfur batteries†

Susanne Dörfler,a,b Markus Hagen,c Holger Althues,a Jens Tübbe,c Stefan Kaskel*a and Michael J. Hoffmannd

Received 19th December 2011, Accepted 28th February 2012
DOI: 10.1039/c2cc17925c

Binder free vertical aligned (VA) CNT/sulfur composite electrodes with high sulfur loadings up to 70 wt% were synthesized delivering discharge capacities higher than 800 mAh g⁻¹ of the total composite electrode mass.

Sulfur cathodes potentially offer significant advantages over metal oxide and phosphate based electrodes for future generation lithium ion batteries. While the relatively low capacity of intercalation cathode materials (150–200 mAh g⁻¹) limits the energy density of today’s lithium ion cells to about 200 Wh kg⁻¹, the theoretical estimated capacity of elemental sulfur is 1672 mAh g⁻¹ and thus energy densities larger than 350 Wh kg⁻¹ are in reach for lithium–sulfur (Li–S) cells.1 Other advantages of sulfur are extremely low cost and widespread availability as well as enhanced safety aspects.

However, Li–S batteries still have not been commercialized yet, despite their promising properties. The so-called shuttle mechanism2 lowering the efficiency of Li–S cells by dissolution, migration and parasitic reactions of polysulfides during charging, and the poor electrical conductivity of elemental sulfur (5 × 10⁻³⁰ S cm⁻¹), causing only low utilization of the active material, are still challenges that have to be overcome. Conducting carbon additives were found to significantly enhance the utilization of sulfur in various literature reports.3–9

In a typical approach a slurry is formulated based on elemental sulfur, a polymeric binder and carbon black dispersed in a solvent.10 For electrode production, the slurry is applied on aluminum current collectors via a doctor blade technique or roll coating and finally dried and calendared. While carbon black is an efficient and established conductive additive, many other carbon materials have been recently reported to be advantageous in Li–S cathodes. Ordered mesoporous carbons were found to offer a conducting framework encapsulating and contacting the sulfur.4 In combination with polyethylene glycol coatings the porous carbons were additionally found to inhibit the migration of polysulfides and thereby suppress the shuttle mechanism. Porous hollow carbons are another promising host material and cathodes based on hollow carbon@sulfur composites were reported to deliver stable discharge capacities of over 850 mAh g⁻¹ (in terms of active sulfur mass) upon 100 cycles at 0.5 C.5 While porous carbons offer high utilization and stabilization of the sulfur, the overall energy density of resulting electrodes is reduced by the high carbon content (30 wt%), the addition of binders (typically 5–10 wt%) and the voids resulting from the interparticulate porosity.

Furthermore, one possible degradation mechanism of sulfur/carbon composite cathodes is known to result from structural changes of the interparticle network as observed for carbon black in sulfur electrodes.6 During cycling, the uniform distribution of sulfur in the composite is lost resulting in decreased utilization and consequently in poor cycle life time. Carbon nanotubes (CNTs), when applied as a conductive additive in paste or slurry formulations, were found to be superior due to higher retention of the integrity of the conductive network caused by the high aspect ratio and the high intrinsic conductivity of CNTs.7–12

Similar to the electrodes from porous materials, the reported CNT/S composite electrodes typically have sulfur contents below 70 wt%, as the paste formulations contain binders and other additives. The largest reported specific capacities related to the electrode mass \((m_{\text{sulfur}}, m_{\text{carbon}}, m_{\text{binder}} \text{ and } m_{\text{additives}})\) are below 600 mAh g⁻¹ both for CNT–S composites8 and for porous carbon–S composites4 (estimated from the average capacity values of the first 20 discharge cycles), while in many publications capacities are far below this value. Increasing the sulfur content in the electrode, while retaining its high utilization, is one of the challenging tasks to exploit the full potential of Li–S batteries in terms of energy density.

Here we report a new approach based on binder free VA-CNT/sulfur composite electrodes enabling high sulfur loadings up to
70 wt% and delivering discharge capacities higher than 800 mAh g\(^{-1}\) related to the composite electrode mass. This value significantly exceeds capacity values related especially to the composite mass from previous reports to the best of our knowledge.

In contrast to the common procedure of preparing the sulfur/ carbon composite by mixing and subsequent slurry coating, we start with the production of a VA-CNT electrode using chemical vapor deposition (CVD) on metallic substrates based on a process developed by Hata et al.\(^{13}\) For the VA-CNT growth, a catalyst thin film is deposited on metal foil by wet chemical techniques from organic metal salt solutions (Fig. 1, step 1).

The seeded substrate is transferred to the CVD chamber in a second step and CNT growth is performed at 730 °C using ethene as carbon precursor (Fig. 1, step 2). Details about the synthesis and the influence of catalyst composition and process parameters on the CNT film properties have been reported elsewhere.\(^{14}\) The resulting VA-CNT films are obtained as homogeneous black coatings on the nickel foil. In a typical experiment, the CNT films have a thickness of up to 200–200 μm (Fig. 2a) and a density of about 0.06–0.13 g cm\(^{-3}\). The individual CNTs are multiwalled and exhibit diameters of 7–30 nm. Due to the direct connection between each single CNT and the metal foil, the material offers a highly conductive and stable framework for the direct connection between each single CNT and the metal foil, while the metal foil acts as a current collector. To finally prepare CNT/S composite cathodes using this unique material architecture, elemental sulfur is infiltrated in a subsequent step into the VA-CNT/Ni electrode (Fig. 1, step 3). Infiltration is achieved from sulfur solutions in toluene. SEM investigations indicate a complete and homogeneous infiltration of sulfur in the CNT network after melt infiltration, while capillary forces cause the formation of trenches and walls within the composite layer (Fig. 2b). Sulfur/carbon ratios up to 7:3 are obtained. TEM images (ESI\(^{+}\), Fig. S1) reveal that the CNTs are covered by large amounts of amorphous sulfur. EDX (energy dispersive X-ray spectroscopy) mapping (ESI\(^{+}\), Fig. S2) proves the homogeneous sulfur dispersion, as well.

Electrochemical cells are constructed from the CNT/S composite cathodes, lithium anodes and separators, while a solution of 1 M LiTFSI in DME:DIOX (1:1, v:v) is selected as the electrolyte.

Fig. 3 shows the results of the cycling experiments using a charge and discharge rate of C/13. An electrode with 69 wt% sulfur content, as prepared by solution infiltration, has discharge capacities of over 1300 mAh g\(^{-1}\) (related to sulfur mass) and 800 mAh g\(^{-1}\) (related to composite mass) within the first cycles, representing a very high active mass utilization. In Fig. 2c SEM images of a CNT–S electrode after 20 cycles in the charged state are shown. The electrode was examined after disassembling the test cell in the glove box directly after 24 h of drying in the glove (Fig. 3, the calculated free volume is about 94 vol%. In the VA-CNT system, pores are formed by the tube interspaces.

When pressed to one fifth of the original volume, the free space between the tubes is reduced to 71 vol%. This free space is filled with sulfur to a very large extent, as 69 wt% sulfur in the composite corresponds to 65 vol% (assuming a sulfur density of 2.06 g cm\(^{-3}\)).

After cycling the electrode film is coated by a fine layer that could be identified as elemental sulfur by Raman spectroscopy,\(^{16,17}\) with characteristic modes at 151 cm\(^{-1}\), 217 cm\(^{-1}\) and 472 cm\(^{-1}\). The CNT network could still be identified even through some gaps in the sulfur layer. Gaps are expected to be responsible for the operability of the electrode since they inhibit the complete isolation of the electrode by sulfur or...
This journal is © The Royal Society of Chemistry 2012


Fig. 4 Specific capacities (related to sulfur mass and electrode mass) vs. cycle number with LiNO₃ addition in the electrolyte. The CNT/S composites have a sulfur content of 63%.

Li₂S even at high sulfur loadings in the cell. The thickness of the sulfur layer is estimated to be below 50 nm. SEM observations also indicate the CNT film to be intact, undamaged and still in contact with the current collector.

Large sulfur loadings, the absence of a binder and the high density of the electrode, together with the high sulfur utilization during cycling are clear advantages over existing electrode processing methods. In direct comparison with unoriented CNT/S composites, higher capacity values related to sulfur as well as to composite mass were obtained. However, the large differences between charge and discharge capacities indicate a low coulombic efficiency, and a remarkable degradation is observed over the first 40 cycles. These effects are expected to result from the polysulfide shuttle mechanism and side reactions at the metallic anode.

To enhance the charge/discharge efficiency, LiNO₃ is used as electrolyte additive. LiNO₃ is known to be reduced to Li₂NO₃ on Li and it oxidizes the sulfides to Li₂SO₅ species that passivate the Li electrode and thus prevent the continuous transport of electrons from Li to polysulfides in the electrolyte.18 The addition of LiNO₃ (0.25 M) to the electrolyte leads to a much higher cycling stability and coulombic efficiency for an electrode with 63 wt% sulfur, clearly inhibiting the shuttle mechanism (Fig. 4). After large irreversible capacities in the first cycles, which can be attributed to the formation of the above-mentioned passivation layer, the capacity ranges between 800 and 900 mAh g⁻¹ of sulfur and 500–600 mAh g⁻¹ of sulfur/VA CNT composite (Fig. 3).

In summary, a new binder-free electrode containing VA-CNTs, directly synthesized on a Ni current collector and subsequently infiltrated by sulfur was introduced as a promising choice for Li–S cells.

In cycling experiments discharge capacities higher than 800 mAh g⁻¹ (mass of the composite) were demonstrated over 20 cycles. This value significantly exceeds results obtained from slurry made electrodes, especially with respect to the capacity per composite mass. SEM investigations before and after cycling revealed the CNT film provides a stabilizing, conducting network for the sulfur cathodes. As the soluble polysulfides are not retained actively by the electrode, its benefits are most suitably complemented by other mechanisms to shut off the shuttle mechanism as demonstrated exemplarily by applying the LiNO₃ additive.

Adjusting the CNT-CVD process to deposit on aluminium or carbon substrates and thereby replacing nickel potentially enables even further improvement in terms of weight and costs of resulting electrodes. Preliminary experiments confirm the feasibility of CNT deposition on aluminium and carbon substrates and will be reported in separate publications. Due to improved performance of the electrodes, low cost precursors and scalable continuous processing, the synthesis-on-current collector strategy can be a competitive approach for electrode manufacturing.

This research was supported by the German Bundesministerium fuer Bildung und Forschung (BMBF) through the project Fraunhofer Systemforschung fuer Electromobilitaet (FSEM).

Notes and references


Published on 29 February 2012. Downloaded by SLUB DRESDEN on 26/03/2014 08:42:09.