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31 32 FOCUS ISSUE OF SELECTED PAPERS FROM IMLB 2016 WITH INVITED PAPERS CELEBRATING 25 YEARS OF LITHIUM ION BATTERIES

Effective Suppression of Polysulfide Dissolution by Uniformly Transfer-Printed Conducting Polymer on Sulfur Cathode for Li-S **Batteries**

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The lithium-sulfur battery has received much attention in recent years owing to its high gravimetric capacity, far beyond that of current Li-ion batteries. Overcoming the shuttling effect caused by the dissolution of polysulfides during the charge-discharge process is a major challenge for the realization of Li-S cells. Here we report transfer-printing of a conductive polymer to cover and securely passivate the surface of sulfur electrodes to prevent the dissolution of polysulfides and, simultaneously, to provide high electronic conductivity of sulfur cathodes. Highly uniform polyaniline film can be controllably formed on sulfur cathodes via the transfer printing method, and a sulfur cathode with the printed polyaniline layer showed improved cycle performance (capacity retention of 96.4% and an average Coulombic efficiency of 99.6% for 200 cycles) compared to conventional sulfur cathodes. In situ measurement of transmittance during discharge demonstrated that the dissolution of sulfur in electrolytes is considerably suppressed by the printed polyaniline layer, substantiating that transfer-printed polyaniline film can provide robust protection as well as supplement electrical conductivity to the sulfur cathode. This strategy could be extensively applied to sulfur cathodes of diverse morphology and further extended to large-scale production.

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The development of high-energy density rechargeable batteries 33 is of great significance, given the continuously increasing demand 34 for portable electronic devices, and for a variety of electric vehicles 35 (EVs).¹⁻⁶ However, the energy density of commercial lithium-ion bat-36 teries, which are based on lithium intercalation oxides and graphite, 37 is limited to about 150 Wh kg⁻¹, still far from what is needed for 38 39 practical EVs. This is chiefly due to the heavy weight and insufficient storage capacity of transition-metal insertion compounds (e.g., LiCoO₂, LiMn₂O₄, and LiFePO₄).^{7,8} As an alternative material for 41 light-weight cathodes, sulfur can react with metallic lithium to form 42 Li₂S via a two-electron reaction, leading to a high theoretical capacity 43 44 of 1675 mAh g^{-1} and a high theoretical energy density of 2500 Wh 45 kg⁻¹ in lithium-sulfur battery systems. This is almost an order-ofmagnitude higher than that of conventional Li-ion batteries.^{9,10} More-46 over, the natural abundance, low cost, and environmental friendliness 47 of sulfur could make lithium-sulfur materials highly practical for next 48 generation, high-energy-density batteries. 49

However, the practical application of Li-S cells is still challenging, 50 mainly due to their high volumetric expansion and shrinkage (up to 51 80%), to the dissolution of polysulfides resulting in the so-called 52 shuttling mechanism during the charge-discharge process, and due 53 to the low electronic conductivity of sulfur ($\sim 10^{-30}$ S cm⁻¹).¹¹⁻¹³ 54 In particular, the polysulfide shuttling mechanism is caused by the 55 phenomenon that, during a charge cycle, dissolved polysulfides in 56 electrolytes diffuse to the Li anode, are reduced; then diffuse back 57 to the sulfur cathode during discharge cycles as they consume the 58 active materials.^{14–17} These challenges must be met to enable more 59 extensive, practical utilization of lithium-sulfur batteries. 60

Considerable efforts have been made to enhance the electrochem-61 ical performance of sulfur cathodes for more advanced lithium-sulfur 62

cells. One of the most effective approaches is the encapsulation of sul-63 fur particles in a conducting matrix to improve electrical conductivity, 64 and to prevent the dissolution of high-order polysulfides.^{18,19} For ex-65 ample, mesoporous carbon was used to trap high-order polysulfides 66 in their small, numerous pores.²⁰ However, despite the successful im-67 provement of electrochemical performance, the weight fraction of the 68 sulfur in these composite electrodes is usually low (<50%), result-69 ing in a loss of total energy density in the cell.^{21,22} Other approaches 70 that achieved relatively higher sulfur content inevitably involved a 71 complicated process for scaling up lithium-sulfur cells.² 72

To address these important issues, here we report transfer-printing 73 of conductive polymer (polyaniline) to cover and securely passivate 74 the surface of sulfur electrodes to prevent the dissolution of poly-75 sulfides and, simultaneously, to provide high electronic conductivity 76 of sulfur cathodes. As a result, the trapping effect of polysulfides 77 by the uniformly printed thin polyaniline, enhanced cycling stabil-78 ity substantially. The capacity retention was 96.4% with an average 79 Coulombic efficiency of 99.6% during 200 cycles, which is markedly 80 improved compared to a bare counterpart. Moreover, unlike previous 81 approaches, our printing method could be widely applied to the diverse 82 morphology of sulfur cathodes and could be extended to large-scale 83 production. 84

Experimental

155°C for 12 h in a closed vessel for sulfur infiltration.²⁴ The sulfur

Preparation of the graphene oxide-sulfur composite (GO-S).-All chemicals used in the synthesis of graphene oxide were purchased from Sigma Aldrich. Graphene oxide (GO) was prepared using a 88 modified Hummer's method. After washing and exfoliating the GO 89 samples, the centrifuged GO was freeze-dried under vacuum for two 90 days. The graphene oxide (GO) and sulfur (Sigma Aldrich) (weight 91 ratio of 1:3) were mixed by ball milling. The mixture was heated at

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Polymer coating on metal foil Stainless steel foil Sulfur cathode A foil - Crosslinked PANI Hore annealing Release Sulfur cathode Sulfur cathode A foil - Crosslinked PANI Thermal annealing Thermal annealing Release Sulfur cathode Sulfur cathode Sulfur cathode Crosslinked PANI Hore Annealing Ho

Figure 1. Schematic illustration of the PANI-layer coated by a printing method.

⁹⁴ content of GO-S composite is about 74%, which is about the same as ⁹⁵ the initial GO and sulfur ratio.

96 Preparation of polyaniline (PANI).—The aniline monomer solution (Sigma Aldrich) was mixed with 1 M hydrochloric acid (HCl) 97 (Sigma Aldrich) in a volume ratio of 100:1 under stirring. After 30 98 min, the equivalent number of moles (with respect to aniline) of 22.8 99 wt% aqueous ammonium persulfate solution (APS) (Sigma Aldrich) 100 was slowly added, followed by oxidative polymerization at 0°C for 101 102 24 h. The resultant green solid was obtained by centrifugation and washed thoroughly with water and ethanol, to remove excess ions and 103 monomers. 104

Preparation of the PANI-printed electrode.- To prepare working 105 electrodes, the as-prepared GO-S was mixed with the conducting car-106 107 bon Super P (Timcal) and the binder sodium alginate (Sigma Aldrich) 108 in a weight ratio of 75:15:10 using an agate mortar to the DI-water. The slurry thus prepared was cast on an aluminum foil (UACJ) and 109 dried at 60°C under vacuum for 24 h. The PANI solution in n-methyl-110 2-pyrrolidone (NMP) (20 wt%) was spin-coated and tape-casted onto 111 stainless steel foil. The polymer-coated substrate was annealed at 112 180°C for 1 h. The PANI layer was printed on top of the electrode 113 surface using a roll press. 114

Preparation of the air-tight cell for UV/Vis measurement.—To 115 measure the in situ UV/Vis transmittance spectra for the lithium-116 117 sulfur cell during discharge, the electrode was fixed on one side of a rectangular-tube cell using paraffin film, followed by placement of the 118 Li metal (HONJO) on the opposite side. Then, the cell was completely 119 filled with electrolyte and capped with polydimethylsiloxane (PDMS). 120 All procedures were carried out in a glove box filled with high-purity 121 Ar gas. 122

Characterization .- The weight fractions of carbon and sulfur 123 were determined using an Elemental Analyzer (Flash 2000, Thermo 124 Scientific). The morphology of the materials was observed using field-125 emission scanning electron microscopy (FESEM, Hitachi S-4800). 126 The transmittance of the lithium-sulfur cell was measured by UV/Vis 127 spectrometer (UV-3101PC, Shimadzu Co., Japan). The reflectance 128 spectra were recorded using a spectral incident photon-to-current ef-129 ficiency (IPCE) measurement system, equipped with an integrating 130 sphere assembly (K3100, McScience, Korea). 131

Cell assembly and battery testing.-Electrochemical measure-132 ments of all electrodes were performed by preparing 2032-type coin 133 cells that were assembled in an Ar-filled glove box. The PANI-134 printed electrode and lithium foil were used as the working and 135 counter/reference electrodes, respectively. The electrolyte was pre-136 pared by dissolving 1 M lithium bis(trifluoromethanesulfonyl)imide 137 (LiTFSI) (Sigma Aldrich) in a solvents of 1,3-dioxolane (anhydrous, 138 contains \sim 75 ppm BHT as an inhibitor, 99.8%, Sigma-Aldrich) and 139 1,2-dimethoxyethane (anhydrous, 99.5%, Sigma-Aldrich) (volume ra-140 tio = 1:1). Before the electrolyte preparation, these solvents were 141 stored over molecular sieves for 24 h to remove moisture. Polypropy-142 lene membranes (Celgard, Inc.) were used as separators. Galvanos-143 tatic measurements were performed in the potential range of 1.0 to 144 3.0 V vs. Li⁺/Li using a battery cycler (WBCS3000, WonAtech). The 145 C-rates for all coin cells were calculated based on the theoretical ca-146 pacity of sulfur in the composite, for which 1 C is 1675 mA g^{-1} . The 147 mass of sulfur loaded as the active material is about 5 mg cm⁻². All 148 of the electrochemical measurements were performed at 25°C. For 149 the in-situ UV/Vis measurement, the quartz cuvette cell was com-150 pletely covered with the lid to avoid interference by ambient light. 151 After taking the initial spectrum of the as-prepared cell, the Li-S cell 152 was discharged at a rate of C/20 using a potentiostat (VSP-200, Bio 153 Logic). Meanwhile, UV/Vis spectra were recorded every 0.1 V from 154 the open-circuit-voltage (~ 2.8 V) to 1.5 V. 155

Results and Discussion

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One of the key advantages of the transfer-printing method is that 157 the polymer layer thickness and heat-treatment conditions are control-158 lable without affecting the underlying electrodes. As schematically 159 summarized in Figure 1, the PANI solution in NMP was spin-coated 160 onto a stainless steel foil to obtain a thin, uniform film the thickness of 161 which could be controlled by adjusting the solution concentration and 162 spin-casting speed. The PANI-coated substrate was then annealed at 163 180°C for crosslinking.²¹ It was expected that the cross-linked PANI 164 layer would improve the trapping of polysulfides and play a role 165 as a buffer layer against the volume changes of the sulfur electrode 166 during repeated charge-discharge cycles. Then, the PANI layer was 167 transfer-printed on top of the sulfur electrode via simple contact of 168 the PANI-coated side of the stainless foil. Detailed procedures were 169 described in the Experimental Section. 170

The morphologies of the electrode with and without PANI coating are shown in Figure 2a. Figure 2a shows the rough surface of the pristine GO-S electrode, which is composed of GO-S, active 173

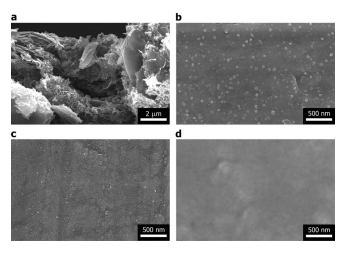


Figure 2. SEM images of (a) Unprinted electrode, (b) PANI-coated stainlesssteel foil, (c) PANI-printed sulfur electrode, and (d) PANI-printed electrode; after 100 cycles.

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avoid not only the detachment of the active materials from the current 180 collector, but also has the advantage of preventing the formation of 181 high-order polysulfide and its diffusion to lithium metal, thus sup-182 pressing the shuttling mechanism to a great extent. Therefore, it is 183 expected that both the cyclability and Coulombic efficiency of the 184 Li-S cell could be improved. In order to examine the morphology 185 changes after charge/discharge, a repeatedly cycled cell was disas-186 sembled in a glove box and washed thoroughly with propylene car-187 bonate (PC) solvent. It can be seen from Figure 2d that, except for the 188 formation of slight cracks, most of the PANI layer retained integrity 189 after electrochemical cycling, implying the stability of the PANI layer 190 during charge-discharge in the electrolyte. 191

The electrochemical performance of the PANI-printed electrode 192 was evaluated using coin-type half-cells. Figure 3a shows the thick-193 ness of the PANI layers measured using a surface profiler. The thick-194 ness of spin-coated PANI layer was controlled (from 1 to 5 μ m) by 195 adjusting the spin casting speed from 1000-3000 r/min. For compar-196 ison, a tape-casted PANI layer with a thickness of 30 µm was also 197 prepared. The mass of PANI layer with a thickness of 1 µm was about 198 0.1 mg. Then, we compared the galvanostatic capacity-voltage profiles 199 of the sulfur electrodes containing different types of PANI protection 200 layer. Figure 3b presents the second discharge-charge profiles dur-201 ing 0.1 and 1C cycling. It was found that there exist two plateaus, 202 at approximately 2.3 and 2.0 V, for bare sulfur electrodes due to the 203 two-step reduction of sulfur in the presence of Li ions.^{24,25} On the 204 other hand, the PANI-printed samples presented a large overpotential 205 and decrease in capacity, due to the PANI capping layer. 206

However, the cycle performance test results in galvanostatic mode 207 showed that the PANI-covered electrode presented both significantly 208

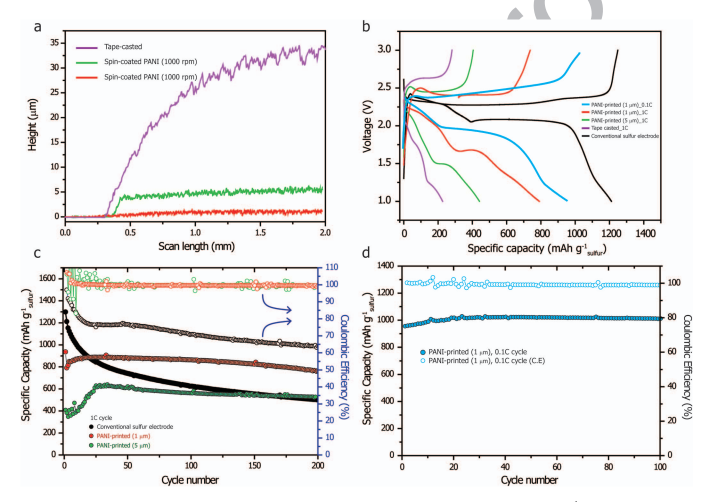


Figure 3. (a) Thickness of the PANI layer, (b) Galvanostatic capacity-voltage profiles of the sulfur electrodes (1 C = 1675 mA g⁻¹), (c) Capacity retention and Coulombic efficiency of the sulfur electrodes at 1 C-rate, (d) Cycle performance of the PANI-printed electrode with a 1 μ m PANI layer at the rate of 0.1C.

improved cyclability and high Coulombic efficiency. Whereas the 209 conventional sulfur electrode delivered an initial discharge capacity 210 of 1298 mAh g^{-1} , the retained capacity was only 500 mAh g^{-1} (41.2%) 211 retention with respect to the capacity in the second cycle) after 200 212 cycles at a rate of 1C (Figure 3c). The average Coulombic efficiency 213 of the conventional sulfur electrode in the cycle range 2-200 was low 214 (\sim 71.6%). The PANI-printed cell with a 5-µm PANI layer initially 215 delivered 400 mAh g^{-1} (discharge capacity), but this capacity increased to 600 mAh g^{-1} after 50 cycles. Interestingly, the Coulombic 216 217 efficiency of the cell with the PANI-printed layer showed significant 218 improvement (close to 100%) compared with the conventional sulfur 219 electrode. In the case of printed cells with a 1 µm PANI protection 220 layer, despite a relatively lower initial discharge capacity (935 mAh 221 g^{-1}), the capacity retention (96.4%) was markedly higher, with an 222 average Coulombic efficiency of 99.6%. 223

To confirm the cyclability with a low C-rate, the cycle performance 224 of the PANI-printed cell with a 1-µm PANI layer was further eval-225 uated at 0.1C for 100 cycles (Figure 3d). Even when the operation 226 time for a cycle was increased 10-fold from 1C to 0.1C, 105% of the 227 original capacity was still retained, with 99.3% of average Coulombic 228 efficiency. The galvanostatic capacity-voltage profile of the PANI-229 printed cell with a 1-µm PANI layer was also shown in Fig. S1. This 230 can be attributed to the fact that the PANI layer printed onto the elec-231 trode surface effectively prevented the dissolution of the polysulfides. 232 However, it was also found that the printing of thicker PANI layers 233 (1 and 5 μ m) caused much lower capacity values (Figure 3b). This 234 phenomenon implies that there exists an optimal range of PANI-layer 235 thickness, because a too thick layer would hinder the diffusion of Li 236 ions, while a too thin layer could not function as a robust blocking 237 layer. Electrochemical impedance spectroscopy (EIS) was measured 238 between 100 mHz and 1 MHz at the open circuit potential of the 239 cell with a signal peak-to-peak amplitude of 5 mV for the samples 240 with conventional sulfur electrode and PANI-printed (1 and 5 μ m) 241 as shown in Fig. S2. As the PANI layer becomes thicker, the second 242 semicircle in a low frequency region become apparent and all the 243 charge transfer resistance (R_{ct}) values increase. 244

245 In order to verify the hypothesis that the outstanding capacity retention and Coulombic efficiency of the PANI-printed electrode is 246 due to the suppression of sulfur dissolution, in situ UV-Vis tests were 247 performed with beaker-type cells. The cells were designed to allow 248 the penetration of the beam from the UV/Vis spectrometer, and the 249 250 sulfur cathode and the Li-metal anode were fixed on opposite sides 251 of the cell to avoid interference from the ambient light. In Figure 4a the UV spectrum of the conventional bare electrode showed almost 252 100% of transmittance at the open circuit voltage. During the dis-253 charge cycle, however, a significant change of transmittance spectra 254 $(\Delta T \sim 21.1\%)$ at the wavelength of 605 nm at 2.1 V) was observed, 255 indicating serious dissolution of the long-chain polysulfides in the 256 electrolytes. However, the PANI-printed sample showed much less 257 change in the transmittance ($\Delta T < 10\%$) during the entire discharge 258 process, as confirmed in Figure 4b. These results are consistent with 259 the excellent cyclability provided by the PANI protection layer, which 260 prevents the polysulfides from shuttling during the electrochemical 261 reactions in the cell. 262

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Conclusions

To summarize, we demonstrated that the formation of a conduct-264 ing polymer protection layer could resolve the chronic polysulfide 265 shuttling issue occurring in conventional Li-S batteries. We showed 266 that a very uniform PANI film could be controllably formed on sulfur 267 cathode via the transfer printing method, with outstanding manufac-268 turability, scalability, and cost-effectiveness. The effectiveness of the 269 printed conducting polymer layer was confirmed by improved cycle 270 performances (capacity retention of 96.4% and an average Coulom-271 bic efficiency of 99.6% for 200 cycles) compared to conventional 272 sulfur electrodes (poor retention of $\sim 41\%$). In situ transmittance 273 measurement during discharge demonstrated that the dissolution of 274 sulfur in electrolytes could be substantially suppressed by the printed 275

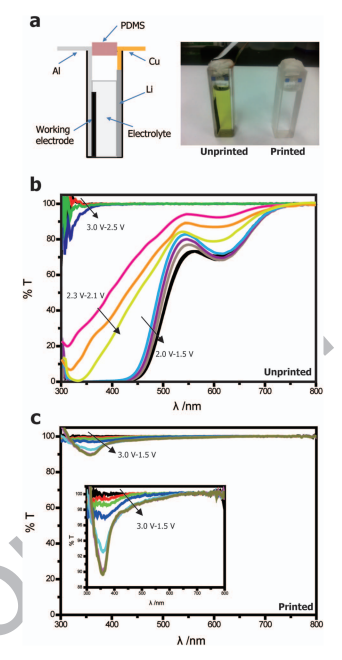


Figure 4. UV/Vis spectra measured in operando mode during first discharge of (a) Unprinted and (b) Printed electrode.

PANI layer, substantiating that transfer-printed PANI film can provide robust protection as well as supplement electrical conductivity to the sulfur cathode. We expect that the strategy proposed in this work could be extensively applied for a variety of other energy-storage electrodes that require both secure protection and electrical conductivity during repeated charge-discharge cycles. 281

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