Sandwich-Stacked SnO$_2$/Cu Hybrid Nanosheets as Multichannel Anodes for Lithium Ion Batteries

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Lithium-ion batteries (LIBs) have revolutionized portable electronics and become the dominant power sources for potential applications in electric vehicles (EV) and plug-in hybrid electric vehicles (HEV). Further improvements in energy and power density put high demands on either new electrode materials or novel structure design. As a potential substitute for commercially available graphite anodes (theoretical capacity 372 mAh g$^{-1}$), SnO$_2$ has a high theoretical capacity (781 mAh g$^{-1}$) and low potential of Li$^+$ intercalation. Generally, SnO$_2$-based anodes suffer from poor electronic conductivity due to intrinsic nonstoichiometry arising from oxygen vacancies. In addition, large volume changes (~300%) during the lithiation/delithiation process inevitably generate mechanical strain, which leads to electrode disintegration and induces rapid capacity fading, making it a significant challenge to achieve superior electrochemical performance. In order to improve the lithium storage capability for SnO$_2$-based electrodes, considerable efforts have been made toward tailoring material structures to enhance the electronic conductivity, accommodate volume changes and facilitate ion diffusion.

Two-dimensional (2D) nanoarchitectures are particularly attractive for Li$^+$ storage owing to their unique structural properties and surface characteristics. However, most 2D nanostructures have smooth facets, easily leading to tightly overlap and form agglomerates. The tight overlapping of nanosheets reduces the active surface area and elongates ion diffusion paths, therefore, heavily hampering their application potential for energy storage. One promising solution to prevent such agglomeration is to create loosely stacked sandwich structures by employing additional additives such as nanotubes, ultrafine nanocarbons and graphene. Recently, progress has been reported for SnO$_2$-graphene sandwich-like 2D architectures, which...
exhibited superior electrochemical behaviors due to their fast ion diffusion and electron transport. In spite of that, most of these fabrication methods require complex processing, and the 2D framework of the structures is mainly limited to graphene nanosheets, upon which active materials are weakly anchored. Therefore, alternative facile and reliable synthetic routes are highly desirable to fabricate sandwich-like 2D nanostructures for high performance LIBs.

Rolled-up nanotechnology is a method employing strain engineering to change thin film in-plane dimensions. Because of stresses generated during film deposition, the stress (strain) gradient through the film thickness causes bending of the film upon release from the substrate. The thin film rolls up into a tubular structure if the film length is longer than one circumference along the rolling direction. This approach has been successfully employed to prepare strain-released microtubes for energy storage as reported previously.21 The main drawback of the rolled-up tubes is the large hollow center, which limits their packing density and therefore the potential for further practical applications. In an attempt to increase the volume efficiency of self-wound nanomembranes, here, we apply rolled-up nanotechnology and a subsequent compression step to create compact sandwich-like nanosheet stacks. SnO2/Cu bilayers are carefully strain engineered and, upon release from the substrate, automatically roll up into tubular structures with precoated carbon black (CB) periodically clamped between each winding. After compressing the rolled up microtubes with a pressure around 2.9 kPa, compact 2D nanosheet stacks with well-defined channels are readily created. With CB acting as the intersheet spacer, restacking and agglomeration of the SnO2/Cu nanosheets are prevented during repeatable charge/discharge process, and volume changes can be accommodated by free channel space. In addition, these 2D nanosheets have large exposed surfaces for electrochemical reactions, from which ion kinetics as well as the interaction between active materials and the electrolyte is enhanced. Furthermore, the hybridized Cu thin layers not only function as a conductive layer to provide good electronic conductivity of the overall electrode, but also act as a structural buffer to preserve the integrity of the SnO2 active layers. The multichannel anodes based on sandwich-stacked 2D SnO2/Cu hybrid nanosheets thus exhibit a high reversible capacity of 764 mAh g−1 at 0.1 A g−1 and a stable cycling performance of ∼75% capacity retention at 0.2 A g−1 after 150 cycles, as well as superior rate capability.

**RESULTS AND DISCUSSION**

The processing of sandwich-stacked SnO2/Cu nanosheets with multiple channels is schematically illustrated in Figure 1a. A 50 nm-thick SnO2 layer is first deposited on top of a precoated sacrificial layer (photoresist AR-P 3510), followed by deposition of a 3 nm-thick Cu layer by electron beam evaporation. Then, homogeneously dispersed CB/ethanol suspension is sprayed on the SnO2/Cu bilayer films forming a cross-linked conductive framework. Metallic Cu, which does not react with the electrolyte and lithium ions during the electrochemical process, is employed to serve as the supporting layer because of its high stability and ideal conductivity. The physically deposited bilayers are hence uniformly contacted with strong affinity, and the structural integrity is well-maintained due to electrochemical inactivity of Cu. After drying at ambient conditions, the CB-coated SnO2/Cu nanomembranes are released from the substrate by dissolving the sacrificial layer with acetone. Meanwhile, the SnO2/Cu composite membranes together with CB automatically peel off from the substrate. As a result of the strain gradient induced by thermal expansion during deposition, the nanomembranes break up into micrometer-sized pieces because of strain pulling, during which these pieces roll up into tubular structures and clamp the
CB framework in between the neighboring SnO2/Cu windings (Figure S1). Under ~2.91 kPa external compression pressure, the tubular structures are subsequently flattened and partially slit at the edges26 to form sandwich-stacked nanostructures (Figure S2). With CB embedded between alternating SnO2/Cu hybrid nanosheets, these multichannel stacks are featured with enhanced mechanical property and effective utilization of interior space. Figure 1b shows schematic configuration of a cell model including stacked SnO2/Cu nanosheets with CB spacer, which could establish lithium ion and electron “superhighways” for lithium ion delivery and storage.

The sample morphology was first characterized by scanning electron microscopy (SEM). Figure 2a shows a top-view SEM image of the as-prepared SnO2/Cu sandwich-stacking, which reveals that the size of inplane sheets is in the micrometer scale according to the pristine tube diameter (φ ~ 20 μm). The cross-sectional profile prepared by focused ion beam (FIB) cutting shows that CB as sandwich stuffing is tightly clamped between adjacent nanosheets (Figure 2b).

The zoomed-in cross-sectional image in figure 2c displays smooth SnO2/Cu hybrid layers with a total thickness of around 53 nm. A statistical study relating the CB cluster sizes to the heights of the multichannels is shown in Figure S3. The gaps, created by CB between adjacent layers, will not only provide open channels for Li+ ion diffusion and electrolyte mass flow, but also offer free space for the volume expansion of SnO2 during lithiation/delithiation processes, which opens up great opportunities to improve battery performance.

To further determine the element composition and uniformity, energy-dispersive X-ray (EDX) elemental mapping (Figure 3a) was carried out with a primary energy of 8 keV, which corresponds to a spatial resolution of about 300 nm. The sum spectrum confirms the composition of the as-prepared stacked SnO2/Cu/CB nanosheets (Figure S4). The results suggest as-prepared samples are indeed composed of Sn, Cu and O elements as well as uniformly dispersed CB. X-ray photoelectron spectroscopy (XPS) analysis was performed to reveal the element stoichiometry of SnO2. E-beam deposited SnO2 was compared with a bulk SnO2 reference sample. Figure 3b presents the Sn 3d core level binding energy spectra of the composite membranes at the surface, where two peaks at ~486.9 and ~495.0 eV, which are assigned to Sn 3d5/2 and 3d3/2, respectively, are detected.27 At the sample surface, the Sn 3d spectra are in good agreement with the characteristic band of SnO2, indicating that surface Sn is fully oxidized to Sn4+. With 4 min presputtering (Ar+, 3.5 keV), both Sn 3d5/2 and Sn 3d3/2 peaks are found to be slightly asymmetric, which indicates a nonstoichiometric oxygen deficiency (SnO2−x) and a small amount of Sn2+ existed in SnO2 active layer. This is not found for the bulk reference sample and therefore not an effect resulting from ion beam damaging during sputtering. The phase structural of SnO2/Cu was found to be amorphous, which was confirmed by X-ray Diffraction (XRD) (Figure S5).

To investigate the electrochemical behavior of the sandwich-stacked SnO2/Cu hybrid nanosheets, control experiments are performed on pure SnO2 nanosheets and SnO2/Cu hybrid nanosheets, which are also prepared by self-rolling and further compressing. All the samples are tested at the same condition, with a current density of 100 mA g−1 and a voltage cutoff in the range from 0.05 to 1.5 V. Figure 4a presents the first 10-cycle profiles of galvanostatic discharge/charge. It is clear that both cycling performance and Coulombic efficiency are improved when combining SnO2 film with Cu thin layer (Figure S6), which attributes to the enhanced conductivity and structural stability.28,29 The produced Li2O during lithiation has been reported to be electrochemically inactive, which substantially decreases the Coulombic efficiency. But the incorporated ultrathin Cu can lead to partial electrochemical decomposition of Li2O particles during the charge process and improve the electrochemical performance of SnO2 nanosheets.30 Moreover, when CB is embedded into the hybrid nanosheets to create multichannels and prevent overlapping, a much better performance is achieved under the same measurement condition.

The detailed electrochemical processes concerning lithiation/delithiation of the sandwich-stacked SnO2/Cu nanosheets were further investigated by cyclic voltammetry (CV). Figure 4b presents the CV curve at 0.1 mV/s scan rate in the potential window of 0–2 V. The reaction mechanism of SnO2-based active materials is described by the following equations:31

Figure 2. (a) Top-view SEM image of the sandwich-stacked SnO2/Cu nanosheets formed by rolled-up nanotech; (b–d) cross-sectional images of the electrodes composed of stacked SnO2/Cu nanosheets with multichannel design using CB as the spacer. The CB maintains the electrical connectivity between the SnO2/Cu nanosheets as a soft medium to buffer the stress of volume expansion and as a barrier against the reaggregation of SnO2/Cu nanosheets (Figure S2). Under ~2.91 kPa external compression pressure, the tubular structures are subsequently flattened and partially slit at the edges26 to form sandwich-stacked nanostructures (Figure S2). With CB embedded between alternating SnO2/Cu hybrid nanosheets, these multichannel stacks are featured with enhanced mechanical property and effective utilization of interior space. Figure 1b shows schematic configuration of a cell model including stacked SnO2/Cu nanosheets with CB spacer, which could establish lithium ion and electron “superhighways” for lithium ion delivery and storage.

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SnO₂ + 4Li⁺ + 4e⁻ → Sn + 2Li₂O (1)

Sn + xLi⁺ + xe⁻ ↔ LiₓSn (0 ≤ x ≤ 4.4) (2)

In the initial cycle, a broad reduction peak is observed at 1.25 V, which relates to eq 1 and describes the reduction of SnO₂ into element Sn and the formation of inactive Li₂O. The cathodic peak located around 0.9 V is attributed to the formation of a solid electrolyte interface (SEI) layer, which consumes Li⁺ ions but can buffer and constrain the volume change thereafter. The two discussed peaks disappear in the following cycles, indicating the irreversibility of these two processes. In the reduction reaction, there are another two main peaks below 0.6 V, which are assigned to the alloying process of LiₓSn shown in eq 2. During the reverse scanning, four anodic peaks in the voltage range of 0.4–0.9 V are observed, which relates to the LiₓSn dealloying and Li⁺ release process. The coincidence of cycle 2 and cycle 3 indicates good repetition of the cycling performance, which is very consistent with the results of the galvanostatic discharge/charge curves shown in Figure 4a.

Figure 5a shows cycling performance of the sandwich-stacked SnO₂/Cu hybrid nanosheets as anode materials at a current density of 100 mA g⁻¹ for 100 cycles. The material delivers a high reversible capacity of 572 mA h g⁻¹ after 100 cycles, which retains 75% of its initial capacity. Such a stable cycling performance is superior to that of SnO₂ nanosheets and many other previously reported SnO₂-based nanostructures. With higher current rates (500 and 1000 mA g⁻¹) applied, highly stable cycling performance is also obtained at each current density for 100 cycles. Similar to the electrode behavior at a current density of 100 mA g⁻¹, there is slight capacity decay over initial 10 cycles, which is plausibly explained by consumption of Li⁺ for the formation of Li₂O inactive phase and the SEI layer. Beyond the 10th cycle, the cycling stability becomes extremely good and the capacity retention remains at around 90% after 100 cycles.

In addition to the excellent cycling performance, these sandwich-like stacks also exhibit convincing rate capabilities. Figure 5b shows the rate capability of the anode, where a gradually increased current rate is applied to an assembled cell. Starting from 100 mA g⁻¹, a reversible capacity of 470 mA h g⁻¹ is obtained after 40 cycles at a current density as high as 1000 mA g⁻¹. When the discharge/charge current density is reduced back to 100 mA g⁻¹, the reversible capacity increases to 590 mA h g⁻¹ and stays above 580 mA h g⁻¹ in the following cycles. To evaluate the cycle life of the stacked SnO₂/Cu nanosheet electrodes, a current density of 100 mA g⁻¹ was applied for 100 cycles, and the results showed a stable cycling performance of 90% after 100 cycles.

Figure 3. (a) SEM image of the stacked SnO₂/Cu nanosheets and the corresponding C, O, Cu, Sn elemental mapping images using EDX at a primary energy of 8 keV; (b) XPS spectra of Sn 3d₅/₂ and Sn 3d₃/₂ for the stacked SnO₂/Cu nanosheets.

Figure 4. (a) Charge-discharge voltage profiles for pure SnO₂ nanosheets, SnO₂/Cu nanosheets, and stacked SnO₂/Cu nanosheets cycled at a current density of 100 mA g⁻¹; (b) cyclic voltammogram of the stacked SnO₂/Cu nanosheets from 2 to 0 V versus Li/Li⁺ at 0.1 mV/s scan rate; the first three cycles are shown.
200 mA g$^{-1}$ is applied for 150 cycles. As shown in Figure 5c, the first reversible capacity is 713 mAh g$^{-1}$, and a charge capacity of 535 mAh g$^{-1}$ can be retained after 150 cycles with capacity retention of 75%. The Coulombic efficiency is well above 98% from and subsequent to the second cycle.

The outstanding electrochemical performance is mainly ascribed to the new structure design of the SnO$_2$/Cu nanosheets. Using CB plays a key role in forming such sandwich-stacked structure. With CB as conductive intersheet spacer, restacking and agglomeration of SnO$_2$/Cu nanosheets are prevented, and the free channel space effectively accommodates volume changes during repeated charge/discharge processes; hence, stable cycling performance could be ensured by the structural stability. As the electrolyte seeps through the nanogaps between adjacent SnO$_2$/Cu nanosheets, there is an additional pathway created for Li$^+$ to reach the interior of the SnO$_2$/Cu nanosheets. Therefore, the sandwich-stacked SnO$_2$/Cu nanosheets help to enhance the rate capabilities of Li-ion batteries by providing higher electrode/electrolyte contact areas, increased number of Li-ion insertion sites, and reduced diffusion length in the intercalation SnO$_2$ nanosheets. Taking into account that the characteristic time constant $\tau$ for diffusion is proportional to the square of the diffusion length $L$ ($\tau \approx L^2/D$, where $L$ is the diffusion distance and $D$ is the coupled diffusion coefficient for Li$^+$ and e$^-$), the diffusion time $\tau$ for Li$^+$ will decrease by decreasing the diffusion length $L$, which finally leads to an excellent rate performance. In addition, the hybridization of Cu thin layer effectively enhances the electronic conductivity of the overall electrode, and the integrity of SnO$_2$ active layer is well preserved due to its structure restriction. Moreover, the nanosheets are formed by self-winding during film strain release. This could effectively ease the intrinsic strain and offer a minimization of the system energy, and thereby enhance the tolerance to stress cracking caused by lithiation/delithiation processes.

To further investigate the lithium-driven structural and morphological changes of the sandwich-stacked SnO$_2$/Cu nanosheet systems, we examined the cross-sectional profile of the stacked SnO$_2$/Cu nanosheets at the 100th fully charged state by FIB cutting. As depicted in Figure 5d, the stacked SnO$_2$/Cu nanostructures are still maintained and the whole structural integrity of the active electrode materials is preserved. The electron backscattering contrast in the image relates to different elements detected by the detector, and continuous white lines indicate the integrity of the SnO$_2$/Cu film. With repeated expansion and shrinking during lithiation/delithiation, the flat sheets twist and curl to form interconnected porous compounds because of the embedded CB, which in turn strengthens the stability of the electrode material. This is in accordance with the results in Figure 5c that there is no obvious capacity decay even after 150 cycles. Therefore, we conclude that the outstanding cycling performance relies on the distinctive stacked 2D nanosheet structures with its excellent structural stability.

CONCLUSION

In summary, we employed compression of self-rolled nanostructures to fabricate novel sandwich-stacked two-dimensional nanosheets with CB uniformly...
METHODS

Sample Preparation. A photoresist (ARP 3510, Allresist) as the sacrificial layer was first spin coated onto Si wafer substrates (3 in.) at 3500 rpm for 35 s prior to baking at 90 °C in an oven. A 50 nm SnO2 film and a 3 nm Cu film were then deposited sequentially by electron beam evaporator (Edwards AUTO500). After sprinkling carbon black (CB) (Timcal) on the hybrid film (100 μL of 0.5 mg/mL CB-Ethanol suspension on each wafer), we thoroughly dried the CB-coated nanomembranes in air at room temperature. Then the samples were directly immersed into acetone to remove the sacrificial layer. In the process of photoresist dissolution, the composite membranes self-rolled into tubular structures clamping with CB. The tubes were then uniformly dispersed on a 3 in. Si wafer surface with a film thickness of roughly single tube height. By means of another Si wafer, 2.91 kPa pressure was applied to compress and flatten the tubes. During this process the edges of the tubes split to form sandwich-like 2D nanosheets.

Material Characterization. A FIB/SEM equipment (NVision 40 CrossBeam, Carl Zeiss) was used to investigate the morphology and cross-sectional profiles of the samples. The microstructure and chemical composition of the samples were investigated using a field-emission gun SEM scanning electron microscope (Zeiss Ultra 55 Plus, 8 keV) equipped with an energy-dispersive X-ray (EDX) spectroscopy instrument. The XPS measurements were carried out using a PHI 5600 CI (Physical Electronics) spectrometer. Raman analysis (from Renishaw) was performed with 442 nm wavelength to check the compositions.

Electrochemical Characterization. Swagelok cells were assembled in an Ar-filled glovebox (H2O, O2 < 0.1 ppm, Mbraun, Germany). Working electrodes were prepared by mixing the SnO2/Cu nanosheets with conductive additive CB (Super P, Timcal) and polyvinylidene difluoride (PVDF, Aldrich) binder at a weight ratio of 70:20:10 in N-methyl-2-pyrrolidinone (NMP, Aldrich) solvent. The slurry was pasted onto a current collector (Cu foil, Goodfellow) and then dried in a vacuum oven at 80 °C for 10 h. The dried electrode was punched into φ = 10 mm discs for cell assembly. Metallic Li foil (Aldrich) was used as both the counter and reference electrodes, and a glass fiber membrane (Whatman) was used as the separator. The electrolyte consisted of a solution of 1 M LiPF6 in ethylene carbonate (EC)/ dimethylcarbonate (DMC) (1:1, in wt %) obtained from Merck, including 2 vol % vinylene carbonate (VC) electrolyte additive (Merck). All the electrochemical tests are conducted on these cells, and the control samples are prepared at the same conditions. The cyclic voltammetry test was performed between 0 and 2 V versus Li/Li+ at a scanning rate of 0.1 mV s\(^{-1}\) via a Zahner electrochemical workstation (IM6ex). Gelvanoastatic cycling was carried out with an Arbin instrument BT2000 at a voltage range of 0.05–1.5 V versus Li/Li\(^+\) at different current densities. The pristine samples are composed of 90.9% SnO2, 6.8% Cu, and 2.3% CB by weight; therefore, SnO2 accounts for 64% of each electrode. The capacity in our work was calculated based on the total mass of the SnO2/Cu nanosheets with CB.

sandwich-stacked SnO2/Cu hybrid nanosheets exhibit significant improvement in cyclability compared to SnO2 nanosheets and SnO2/Cu hybrid nanosheets. A high capacity of 535 mAh g\(^{-1}\) can be retained after 150 cycles at a current density of 200 mA g\(^{-1}\) with 75% capacity retention. The proposed strategy is facile and general, which can be applied to the fabrication of other electrode materials with multichannel design, and enhanced electrochemical performance for lithium ion batteries should be expected.

REFERENCES AND NOTES


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