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A Low-Surface-Area Hard Carbon Anode for Na-Ion Batteries via Graphene Oxide as a Dehydration Agent

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ABSTRACT: Na-ion batteries are emerging as one of the most promising energy storage technologies, particularly for grid-level applications. Among anode candidate materials, hard carbon is very attractive due to its high capacity and low cost. However, hard carbon anodes often suffer a low first-cycle Coulombic efficiency and fast capacity fading. In this study, we discover that doping graphene oxide into sucrose, the precursor for hard carbon, can effectively reduce the specific surface area of hard carbon to as low as 5.4 m²/g. We further reveal that such doping can effectively prevent foaming during caramelization of sucrose and extend the pyrolysis burn-off of sucrose caramel over a wider temperature range. The obtained low-surface-area hard carbon greatly improves the first-cycle Coulombic efficiency from 74% to 83% and delivers a very stable cyclic life with 95% of capacity retention after

200 cycles.

KEYWORDS: hard carbon, Na-ion batteries, anode, low surface area, high first-cycle Coulombic efficiency

Introduction

Advanced energy storage devices are the key enabler for renewable energy sources, including wind and solar power, as well as for the realization of electric transportation. Ideally, they should exhibit high energy/power density and are of low cost and great safety properties. Among different devices, Li-ion batteries (LIBs) have been intensively pursued as power sources for portable electronic devices and electric vehicles (EVs) due to their high energy/power density and long cycle life.^{1–3} However, the rarity and being geographically concentrated of lithium resources may severely restrict LIBs' perspective applications in grid-level energy storage.⁴ Currently, attention starts to shift to Na-ion batteries (NIBs)^{5–7} that exhibit slightly inferior performance metrics compared to LIBs due to the lower cell voltages and less capacities. However, NIBs represent a great fit for grid-level storage because they are potentially very cost-effective due to the abundance and wide distribution of sodium resources. Furthermore, NIBs can utilize inexpensive aluminum current collectors for anodes while LIBs have to use copper.^{8–10}

Over the last few years, significant progresses have been made for NIB cathodes by updating the knowledge learnt from LIBs.^{11–15} However, with respect to anodes, graphitic carbon, the commercial LIB anode, does not function well for NIBs due to its extremely low capacity, ~31 mAh/g, according to the formation of NaC₇₀ upon sodiation.^{16,17} It is the large

ionic size of Na⁺, ~98–102 pm (depending on the coordination number), that causes such a low capacity.^{18,19} Alternatives, including metal alloys,^{20–24} metal oxides^{25–27} or sulfides^{28–30} can provide much higher capacities than carbon-based anodes. However, they often exhibit less favorable high redox potentials and, sometimes, low first-cycle Coulombic efficiency. Among the candidates for NIB anodes, carbon-based materials still holds the promise considering the cost for the potential scaling up, and the most promising one is expected to be hard carbon.

Hard carbon, also known as non-graphitizable carbon, is very difficult to be graphitized even at a temperature higher than 2000 °C.^{31,32} Its structure typically comprises small turbostratic nano-domains with nanosized voids sitting in between these nano-domains.³² Pioneering studies have demonstrated promising performance of hard carbon as anodes for both LIBs and NIBs with low operating potentials and relatively high specific capacities.^{33–36} Unfortunately, hard carbon usually suffers a low first-cycle Coulombic efficiency, lower than 75%, and fast capacity fading over cycling. To date, much attention focuses on fabricating functional carbon nanostructures with high surface areas, which can enhance the rate capability by facilitating better contacts between the electrolyte and the electrode.^{37–39} However, the first-cycle Coulombic efficiency is typically very low for these carbon anodes.⁴⁰⁻⁴² The poor efficiency may be caused by the more prominent formation of solid electrolyte interphase (SEI) layer on the large surface area.^{43,44} This can be a serious issue because the low first-cycle Coulombic efficiency of anodes causes the irreversible loss of active Na⁺ ions, provided by the cathode, which decreases the full-cell capacity. It is well-known that a lower surface area increases the first-cycle Coulombic efficiency for

carbon anodes in LIBs.^{45–47} However, an effective strategy to minimize the surface area of hard carbon remains a significant challenge.⁴⁸

Sucrose is a typical precursor for producing hard carbon due to its abundance, low cost and renewability.^{48,49} The conventional synthesis of hard carbon from sucrose consists of two steps: dehydration and pyrolysis. During the dehydration, water from sucrose caramelization is eliminated to a large degree at a modest temperature, e.g., 180 °C. The following pyrolysis is normally carried out under an inert gas at temperatures, ranging from 900 to 1600 °C. It has been reported that the specific surface area of hard carbon decreases upon increasing high pyrolysis temperature, e.g., 1400 to 1600 °C.^{50,51} However, a high-temperature treatment not only raises the cost but sometimes reduces the reversible capacity.^{52,53} Therefore, it is critical to find an alternative method that is effective to reduce the surface area of hard carbon at a relatively low pyrolysis temperature.

Herein, we report an effective strategy to minimize the surface area of sucrose-derived hard carbon simply by introducing graphene oxide (GO) suspension into the sucrose solution. We discover that a hard carbon of surprisingly low specific surface area, e.g., $5.4 \text{ m}^2/\text{g}$, can be prepared by pyrolysis of such a mixture of sucrose/GO. In sharp contrast, hard carbon derived from sucrose powder shows a much higher specific surface area of $137.2 \text{ m}^2/\text{g}$. When evaluated as an anode for NIBs, sucrose/GO derived hard carbon, referred to as G-HC, exhibits a much enhanced first-cycle Coulombic efficiency of 83% compared to that of sucrose derived hard carbon (designated as HC), 74%. Furthermore, G-HC shows a high desodiation capacity of 280 mAh/g and very stable cycling performance over ~ 200 cycles with capacity retention of ~95%, while HC exhibits a fast capacity fading.

Experimental Section

Materials synthesis. For synthesis of G-HC, GO aqueous suspension is first prepared via a modified Hummer's method.^{54,55} Then, sucrose is dissolved in the GO aqueous suspension (mass ratio of sucrose/GO is 80:1) under sonication. The resulting sucrose/GO solution is dried at 80 °C, dehydrated at 180 °C for 24 h and then pyrolysed at 1100 °C for 6 h under Ar. In a control experiment, sucrose powder is directly dehydrated at 180 °C for 24 h and pyrolysis at 1100 °C for 6 h under Ar.

Materials Characterization. X-ray diffraction (XRD) patterns are collected using a Rigaku Ultima IV Diffractometer with Cu K α irradiation ($\lambda = 1.5406$ Å). The morphology is examined by a scanning electron microscope (SEM) (FEI, Quanta 600F). N₂ sorption measurements are performed on a TriStar II 3020 analyzer. Thermogravimetric analysis (TGA) is performed under Ar from ambient temperature to 1000 °C (ramp rate: 10 °C/min).

Electrochemical measurements. The electrodes consist of hard carbon active mass and polyvinylidene fluoride (PVDF) binder with or without carbon black additive. For preparation of the electrodes, hard carbon was ground in a mortar for 10 minutes, and then a PVDF/NMP solution was added for further 30 minutes grinding. The as-obtained hard carbon/PVDF slurry was coated on Al foil by a doctor blade and dried at 120 °C under vacuum for 12 hours. The mass loading of hard carbon is about 2.5 mg/cm² with a thickness of ~40 μ m. Coin cells, 2032-type, are assembled with sodium pellets as the counter/reference electrodes and 1.0 mol/L NaPF₆ solution in ethylene carbon (EC) / diethyl carbonate (DEC) (1:1 in volume) as the electrolyte. Galvanostatic cycling in a potential window of 0.01–2 V versus Na⁺/Na is conducted on an Arbin BT2000 system at room temperature.

Results and Discussion

Both G-HC and HC exhibit a typical amorphous carbon structure, revealed by the XRD patterns with two broad peaks, indexed to (002) and (101) planes, respectively (Figure 1a). Close observation of the XRD patterns indicates that the (002) diffraction in HC locates at 2θ of 22.8°, which corresponds to an interlayer spacing of 0.39 nm. Interestingly, the (002) peak of G-HC shifts to 24.0°, suggestive of a smaller inter-planar distance of 0.37 nm. N₂ adsorption/desorption measurements are carried out to determine the specific surface area of G-HC and HC. As shown in Figure 1b, HC exhibit a Type I sorption isotherm, suggestive of some microporous porosity with a Brunauer–Emmett–Teller (BET) surface area of 137.2 m²/g. In sharp contrast, G-HC exhibits a surprisingly low BET surface area of 5.4 m²/g. The SEM images reveal that both G-HC and HC consist of bulk particles (Figure 2a and 2c). Moreover, despite the much higher surface area, HC exhibits a much smoother surface, compared to that of G-HC, as the zoomed-in SEM images show (Figure 2b and 2d).

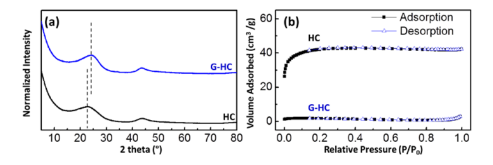


Figure 1. (a, b) XRD patterns and nitrogen adsorption/desorption isotherms of HC and G-HC.

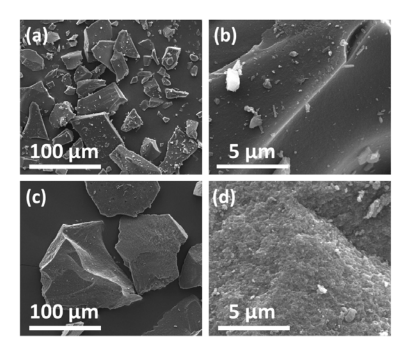


Figure 2. SEM images of HC (a, b) and G-HC (c, d) at different magnifications.

It has been reported that the introduction of graphene normally increases the surface area of the composites due to the high theoretical surface area of single-layer graphene, 2630 m²/g.⁵⁶⁻⁵⁸ It is quite interesting that adding GO into sucrose dramatically decreases the specific surface area of the hard carbon product. In order to investigate this interesting phenomenon, we look back to the synthesis procedure. For preparing hard carbon directly from sucrose powder, dehydration step is critical because it can improve the yield and simplify the sample handling process. The dehydration step is very similar to the caramelization process, where volume expansion of table sugar occurs during melting and foaming. As expected, sucrose is turned into foamed caramel after dehydration (Figure 3a). However, such a foaming phenomenon is not observed when dehydrating the mixture of sucrose/GO (Figure 3a). The mixture of sucrose/GO is obtained by drying its aqueous suspension/solution at 80 °C, which begs a question whether such a solution-to-solid drying

process is why foaming does not take place. In a control experiment, a sucrose aqueous solution is also dried at 80 °C, which results in viscous syrup rather than a solid in contrast to the case of sucrose/GO mixture. When the drying temperature is increased to 110 °C, solid forms out of viscous syrup but foams dramatically (Figure S1). The dried sucrose obtained from its aqueous solution at 110 °C is further dehydrated and pyrolysed, yielding a carbon, referred to as HC-S that exhibits a very high specific surface area of 510.2 m²/g (Figure S2).

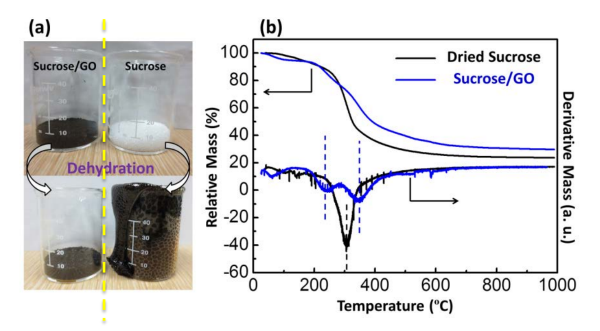


Figure 3. (a) Digital picture of the dried fine mixture of sucrose/GO from its solution (left) and sucrose powder (right) before and after dehydration at 180 °C for 24 h. (b) TGA and differential TGA results comparing the pyrolysis behavior of sucrose/GO mixture and dried sucrose from its solution at a ramp rate of 10 °C/min under Ar.

From the observations above, it is evident that GO plays a major role in preventing the foaming process during drying and dehydration. In order to further investigate this phenomenon, we compared different mass ratios between sucrose and GO during mixing (Table 1). When the mass ratio of sucrose/GO is N, the suspension/solution is referred to as

SG-N. Drying SG-1000 at 80 °C results in viscous syrup, similar to what happens when drying a pure sucrose solution. At 110 °C, SG-1000 foams as well and its final carbon product exhibits a specific surface area of 257.6 m²/g. Interestingly, SG-200 can be dried into firm solid at 80 °C without foaming, and as expected, its carbon product exhibits a specific surface area as low as 4.8 m²/g, even lower than that of G-HC (the product of SG-80), 5.4 m²/g. SG-10 can be dried at 80 °C without foaming as well. The specific surface area of this carbon sample bounces back to 37.2 m²/g. However, it is still much lower than that of the hard carbon obtained without GO addition.

Table 1. Experimental conditions and BET surface areas of the hard carbons derived from sucrose powder, pure sucrose solution and sucrose/GO suspension/solution.

Sample No.	Precursor	Mass ratio of sucrose/GO	Drying temperature of the solution (°C)	BET surface area of the carbon product (m ² /g)
НС	Sucrose powder	No GO	N/A	137.2
HC-S	Pure sucrose solution	No GO	110	510.2
SG-10	Sucrose/GO solution	10	80	37.2
SG-80 (G-HC)	Sucrose/GO solution	80	80	5.4
SG-200	Sucrose/GO solution	200	80	4.8
SG-1000	Sucrose/GO solution	1000	110	257.6

In order to gain further understanding on why the sucrose/GO mixture gives a low surface area, TGA is carried out under Ar for dried sucrose with and without GO. As shown in Figure 3b, the derivative of the TGA results reveal two peaks at ~230 and 350 °C for the dried sucrose/GO mixture while the dried sucrose only exhibits one sharp peak at ~305 °C. It

appears that GO helps spread the burn-off of sucrose during pyrolysis over a much wider temperature range. In summary, without foaming during dehydration, a burn-off over a wider temperature range facilitates the much lower specific surface area of G-HC. We hypothesize that volatile molecules, including water, generated during dehydration and pyrolysis migrate out along GO nanosheets without creating pores. Further research is ongoing in our lab to investigate this phenomenon.

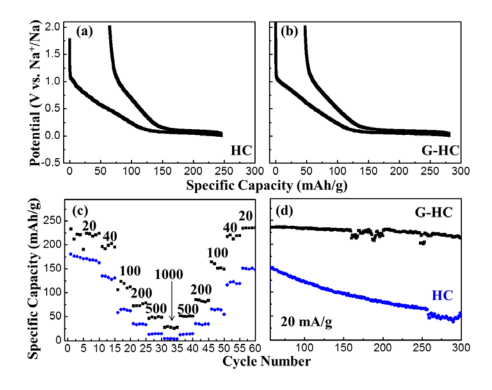


Figure 4. Galvanostatic sodiation/desodiation profiles of the first cycles of HC (a) and G-HC (b) tested in the potential range of 0.01-2 V at a current density of 20 mA/g. (c) Rate and (d) cycling performance of HC (blue) and G-HC (black) tested at current densities from 20 to 1000 mA/g and then at 20 mA/g for long-term cycling in the potential range of 0.01-2 V.

We evaluate the Na-ion storage properties of the as-obtained hard carbons. Working electrodes are only composed of G-HC or HC (93 wt%) active mass and PVDF binder (7

wt%), which is to avoid the impact from carbon black additive on the first-cycle Coulombic efficiency. Galvanostatic sodiation/desodiation cycling is first performed at a current density of 20 mA/g in a potential window of 0.01-2 V versus Na⁺/Na. As previously reported for typical hard carbon anodes in NIBs, HC exhibits two distinct potential regions in the first sodiation profile (Figure 4a). As proposed by Dahn et al., the sloping region from 2.0 to ~0.15 V corresponds to the insertion of Na^+ ions into the inter-layers inside the small turbostratic nano-domains along with the SEI formation, whereas the plateau region near 0.1 V is attributed to the process when Na⁺ ions 'adsorbs' into the nano-voids in between the turbostratic nano-domains.³⁵ The overall first sodiation capacity is 245 mAh/g, where the contributions from the sloping and plateau regions are 130 and 115 mAh/g, respectively. The following desodiation delivers a capacity of 181 mAh/g, yielding a first-cycle Coulombic efficiency of 74%. Moreover, the capacity contribution from the sloping region in the sodiation profile (from ~0.15 to 2.0 V) is 87 mAh/g. Subtracting 87 mAh/g from 130 mAh/g, the capacity from the sloping region during first sodiation, there is an irreversible capacity loss of 43 mAh/g. As shown in Figure 4b, G-HC also exhibits similar characteristic potential profiles of hard carbon. The first sodiation/desodiation capacities are 281 and 233 mAh/g, respectively, leading to a high first-cycle Coulombic efficiency of 83%. This is one of the highest first-cycle Coulombic efficiency values reported for carbon-based anodes in NIBs.⁴⁸ We also compare the capacity contributions from the sloping regions in the sodiation and desodiation profiles, 128 and 96 mAh/g, respectively. The irreversible capacity loss from the sloping region is 32 mAh/g for G-HC, lower than 43 mAh/g of HC. It is known that SEI typically forms during the sloping region of the first sodiation. The less capacity loss from the sloping region of G-HC indicates that the effect of a smaller surface area that does help mitigate the SEI formation, thus enhancing the first-cycle Coulombic efficiency.

Also importantly, G-HC exhibits a much more stable cycling performance than that of HC. Over the following cycles at 20 mA/g, the desodiation capacity of G-HC is stabilized at ~220 mAh/g while HC shows a lower desodiation capacity of ~170 mAh/g (Figure 4c). After cycling at 20 mA/g for 10 cycles, the cycling is conducted with various current densities, ranging from 40 to 1000 mA/g. Evidently, G-HC exhibits a higher capacity than HC at all current densities (Figure 4c). When the current density is switched back to 20 mA/g, G-HC exhibits a desodiation capacity of 235 mAh/g, compared to 151 mAh/g for HC. Then, both G-HC and HC are cycled at 20 mA/g for a long-term cycling. After the total 300 cycles, G-HC still retains a desodiation capacity of 213 mAh/g, compared to 54 mAh/g from HC (Figure 4d).

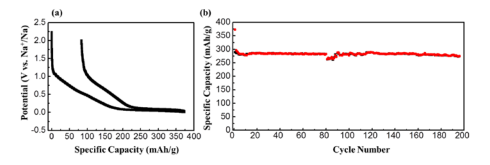


Figure 5. Electrochemical performance of G-HC electrodes with 10 wt% carbon black addition: (a) galvanostatic sodiation/desodiation profiles of the first cycles at 20 mA/g; (b) long-term cycling performance at 20 mA/g in the potential range of 0.01-2 V.

We are also interested in determining the impact from carbon black additive on the performance of hard carbon electrodes. Carbon black (TIMCAL Super C45), 10 wt%, is

added into the G-HC electrodes (mass ratio of G-HC/carbon black/PVDF is 8:1:1). As shown in Figure 5a, the first sodiation and desodiation capacity reaches 373 and 289 mAh/g at a current density of 20 mA/g, respectively, compared to 281 and 233 mAh/g without carbon black. Obviously, higher electronic conductivity in the electrode helps enhance the capacity. However, the first-cycle Coulombic efficiency declines from 83% to 78% with the addition of 10 wt% carbon black, which should be attributed to the greater surface area of carbon black (45 m²/g) and the resulted additional SEI formation. After the first cycle, the desodiation capacity stabilizes at ~280 mAh/g when cycled at 20 mA/g (Figure 5b). Most importantly, G-HC still delivers a desodiation capacity of 274 mAh/g after 195 cycles. It corresponds to capacity retention of 94.8% and the capacity decay is as small as 0.026% per cycle. A few unstable sodiation/desodiation cycles in the cycling curves is due to the temperature change in our laboratories (Figure 4d and Figure 5b).

Conclusion

In summary, we have developed a GO-doping strategy for the synthesis of hard carbon with a very low specific surface area. We discovered that GO effectively prevents foaming of sucrose during caramelization or dehydration and also helps spread the burn-off over a wider range of temperatures during pyrolysis. Both factors facilitate the extremely small specific surface area of the final hard carbon product. When evaluated as an anode for NIBs, with a high active mass loading of 2.5 mg/cm², we report one of the highest first-cycle Coulombic efficiency of 83% for NIB carbon anodes, compared to 74% from HC. Over 300 cycles, almost no capacity fading is observed for G-HC while the conventional HC fades to only 30%

of original capacity. When adding 10 wt% carbon black in the G-HC electrode, the desodiation capacity is increased to 280 mAh/g, and quite stable cycling is still achieved.

ASSOCIATED CONTENT

Conflict of Interest: The authors declare no competing financial interest.

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Supporting Information Available: Additional figures. This material is available free of charge via the internet at http://pubs.acs.org.

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Table of Contents

