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Preparation of artificial graphite coated with sodium alginate as a negative electrode material for lithium-ion battery study and its lithium storage properties[†]

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In this paper, artificial graphite is used as a raw material for the first time because of problems such as low coulomb efficiency, erosion by electrolysis solution in the long cycle process, lamellar structure instability, powder and collapse caused by long-term embedment and release of lithium ions when it is used as a cathode material. The strategy of modifying a graphite anode with a sodium alginate (SA) coating was proposed and a 5.0% SA-1000 composite anode material was obtained. As a result of using sodium alginate (SA) to coat graphite particles, the SEI film formed was stable and compatible with electrolytes. One benefit of the coating is that it limits the amount of irreversible lithium consumed during film formation. Compared with artificial graphite, the first specific capacities of the initial charge and discharge (412.809 mA h g^{-1} and 472.067 mA h g^{-1}) of the 5.0% SA-1000 composite anode material are significantly increased. Furthermore, the organic coating reduces the interface impedance between the electrolyte and the negative electrode. After 600 cycles, the reversible capacity of the composite remains at 48.298 mA h g⁻¹, significantly improving the cyclic stability of graphite. On the other hand, the introduction of sodium alginate improved the migration of lithium ions inside and outside of the graphite, and the reversible capacity of the 5.0% SA-1000 composite sample is 461.197 mA h g⁻¹, 444.457 mA h g⁻¹, 386.633 mA h g⁻¹, 304.794 mA h g⁻¹, 140.846 mA h g⁻¹ and 72.196 mA h g^{-1} at 100 mA g^{-1} , 200 mA g^{-1} , 300 mA g^{-1} , 500 mA g^{-1} , 1000 mA g^{-1} and 2000 mA g^{-1} , respectively. When compared with graphite samples at the same magnification, 5.0% SA-1000 significantly improved its reversible capacity and stability, especially at 300 mA g^{-1} and 500 mA g^{-1} . Its reversible capacities were increased by 196.911 mA h q^{-1} and 184.818 mA h q^{-1} . Therefore, 5.0% SA-1000 composites make it possible to improve the lithium storage capacity of lithium-ion batteries.

1. Introduction

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Recently, the production and storage of energy has become the most important issue in the world.^{1,2} In the field of energy storage, lithium-ion batteries are developing rapidly as a new



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Fig. 1 The crystal structure of graphite.

doping.²⁰ Although the spheroidization process is relatively advanced, it is difficult to significantly improve graphite's rate properties. The doping process is difficult to control because of its elements and amounts. Therefore, spheroidization and doping have to be explored extensively in commercial production, causing difficulties for industrial applications. However, the coating process has been developed and it can improve commercial graphite's rate performance and cycle stability.²¹

Sodium alginate (SA for short), which is made up of β -D-mannuronic acid and α -l-guluronic acid linked by a $(1 \rightarrow 4)$ bond (Fig. 2), has good chemical stability and antioxidant performance. Sodium alginate has been widely used in the fields of national defense, civil industry and medicine and it is used as a high-quality raw material for the production of high power and ultra-high power graphite electrodes, special graphite, lithium anode materials and high-end carbon products.^{22,23} The cyclic stability and rate properties of sodium alginate (SA) can be improved by coating with a modified anode material. However, SA has rarely been reported to have been used as an anode modification material.²⁴

This study used commercial graphite as the raw material and sodium alginate was used as a coating material. To disperse the samples, magnetic stirring was used to stir them in the liquid phase under heat and then they were synthesized by firing them in a tubular furnace in a nitrogen environment.²⁵ The composition and structure of the materials was analyzed by Brunauer–Emmett–Teller (BET) measurements, X-ray diffraction (XRD), Raman spectroscopy, scanning electron microscopy (SEM) and transmission electron microscopy (TEM) characterization. A constant current charge–discharge test, CV test, and EIS test were performed to study the effect of a sodium alginate coating on commercial graphite's performance.

2. Experimental section

2.1 Preparation of the sodium alginate coated graphite composite SA/G-C

• An electronic balance (BS224S, Xiamen Jing vi Xing ve Technology Co., Ltd) was used to accurately weigh 0.50 g of sodium alginate powder (Analytical Reagent, Shanghai Zhan yun Chemical Co., Ltd, molecular weight (198.11)n, CAS No. 9005-38-3). Then 9.50 g graphite powder (Analytical Reagent, Shenzhen Beitrui New Energy Materials Co., Ltd) was added to 100 mL deionized water (Analytical Reagent, made by lithium Laboratory of Jiangxi University of Science and Technology), which was placed on a magnetic stirrer (85-2 digital display constant temperature magnetic stirrer, Changzhou Yue xin Instrument Manufacturing Co., Ltd) and stirred mechanically under heat until the graphite was completely dispersed. It was then dried at room temperature to obtain the precursor powder and then the precursor was placed in a high-purity nitrogen atmosphere. The initial temperature was 25 °C and the heating rate was increased to 300 $^\circ$ C at 10 $^\circ$ C min $^{-1}$ and then to 1000 $^\circ$ C at 30 $^{\circ}$ C min⁻¹. The sample was kept at 1000 $^{\circ}$ C for 2 hours, cooled to room temperature naturally, then sieved to obtain the SA/G-C composite anode material samples. This process is illustrated in Fig. 3.

3. Results and discussion

3.1 Structure and morphology analysis of the 5.0% SA-1000 composite and graphite samples

Through surface adsorption experiments of different electrode materials, a series of physical properties such as specific surface area (Fig. 4), total pore volume and average pore diameter of the composite material 5.0% SA-1000 and graphite samples were tested (Table 1).

The 5.0% SA-1000 sample has a porous structure, resulting in a total pore volume of 0.018 cm³ g⁻¹, which is much larger than the total pore volume of traditional commercial graphite samples of 0.0023 cm³ g⁻¹. Its smaller average pore diameter (3.068 nm) is the key to forming a fluffy structure, which results



Fig. 2 The crystal structure of sodium alginate.



Fig. 3 The preparation process of the composite anode material



Fig. 4 BET for the best coating of the 5.0% SA-1000 and graphite samples.

 Table 1
 BET surface adsorption parameters of the 5.0%
 SA-1000 composite sample and graphite sample

Sample	Surface area $(m^2 g^{-1})$	Pore volume $(cm^3 g^{-1})$	Pore diameter $D_{\rm v}$ (d) (nm)
5.0% SA-1000	3.686	0.018	3.068
Graphite	0.50	0.0023	18.54

in a larger specific surface area and more exposed active sites. The specific surface area of the 5.0% SA-1000 sample is $3.686 \text{ m}^2 \text{ g}^{-1}$, while that of the traditional commercial graphite sample is only 0.50 m² g⁻¹. The 5.0% SA-1000 composite sample with a larger surface volume will be more favorable for improving the lithium storage performance of lithium-ion batteries.

We further confirmed this result by examining the XRD diffraction patterns of the 5.0% SA-1000 composite and commercial graphite samples. The XRD results (Fig. 5) show that the crystalline structure of 5% SA-1000 is obviously weaker than the graphite and the Raman results (Fig. 6) also show the weakening of the crystalline structure and the increase in disorder. In contrast, the (002) crystal plane diffraction peaks of the 5.0% SA-1000 composite are very sharp, indicating that the preparation of the composite anode material has a high graphitization degree so that it will have a high initial charge-discharge capacity.

Raman spectroscopy analysis of the order degree of the 5.0% SA-1000 composite material and graphite sample was conducted to further explore the existing forms of various elements



Fig. 5 XRD of the 5.0% SA-1000 composite and graphite sample.

after SA-coated graphite and to analyze the modification mechanism (Fig. 6). For the graphite samples, a typical G-band at 1568.63 cm⁻¹ and a 2D band at 2703.66 cm⁻¹, a typical G-band was observed, corresponding to the sp² hybrid structure of ordered graphite, which further indicated that the graphite is composed of layered ordered materials. However, for the 5.0% SA-1000 composite sample, the two typical D-band peaks at 1325.41 cm⁻¹ correspond to the sp³ hybridization of the disordered state and the G-band peak at 1568.63 cm⁻¹ corresponds to the sp² hybridization of ordered graphite structure. There is a 1.13 intensity ratio between the D-band peak and the G-band peak and the G-band peak and the S.0% SA-1000 contains disordered sp³ hybrid carbon and graphite-like



 sp^2 hybrid carbon, in which the disordered amorphous state is dominant. Consequently, the hybrid structure of the 5.0%

SA-1000 composite material is more effective at accelerating faster diffusion and improving the rate performance of the lithium-ion battery.

SEM characterization was conducted to investigate further the structural difference between the 5.0% SA-1000 composite and graphite sample (Fig. 7). As shown in (Fig. 7a–c), the graphite samples are regular spherical particles with smooth surfaces and no adhesion between each other. Fig. 7(d–f) shows that the surface of the 5.0% SA-1000 composite sample is relatively fluffy and the grains are closer together, which is caused by the viscous sodium alginate wrapped around the graphite surface. We speculate that this is also the primary reason why the specific surface area of the 5.0% SA-1000 composite sample increased slightly after coating.

Further exploration of the material's crystal structure was performed with transmission electron microscopy (TEM) of the samples with a concentration of 5.0% SA-1000 and raw graphite (Fig. 7g–n). A TEM image of graphite specimens reveals a



Fig. 7 Electron microscopy images of the 5.0% SA-1000 composite and graphite samples (a-f) SEM and (g-n) TEM.

high degree of crystallinity and an orderly and regular lamellar structure, as shown in (Fig. 7g–i), showing regular lattice stripes of 0.335 nm. However, for the 5.0% SA-1000 sample, it was observed that the fringes are curved and disordered, as shown in Fig. 7j–l, indicating that they are stacked together and disordered. A material with an isotropic microstructure is more conducive to the rapid transport of lithium ions since it can facilitate the smooth embedding and expulsion of lithium ions. In addition, Fig. 7m and n shows that the layer spacing of the 5.0% SA-1000 sample is 0.362 nm, which is larger than the layer spacing of graphite (0.335 nm), indicating that the 5.0% SA-1000 electrode is more conducive to the rapid insertion and removal of lithium ions, so it will have better rapid charging and discharging performance than the graphite electrode.

 Table 2
 The initial charge-discharge capacity and Coulomb efficiency of the 5.0% SA-1000 composite and graphite samples

Sample	First specific charge capacity (mA h g ⁻¹)	First specific discharge capacity (mA h g ⁻¹)	Charge–discharge efficiency (%)
5% SA-1000	412.809	472.067	87.447
Graphite	342.306	392.159	87.287

3.2 Electrochemical performance analysis of the 5.0% SA-1000 composite material and graphite sample

Using the charge-discharge specific capacity of the 5.0% SA-1000 composite sample and graphite (Table 2) and the charge-discharge curve distribution (Fig. 8a and b), the charge-discharge curve of the first circle (Fig. 8c) is further compared. After coating, the initial discharge specific capacity of 5.0% SA-1000 increases from 392.159 mA h g⁻¹ to 472.067 mA h g⁻¹ and the specific capacity of 5.0% SA-1000 increases from 342.306 mA h g⁻¹ to 412.809 mA h g⁻¹. The first Coulomb efficiency increased by 0.16% (87.287% \rightarrow 87.447%). Similar to the SA-1000 composite material, 5.0% SA-1000 has a greater capacity and Coulomb efficiency in subsequent cycles of 2–3 curves. Additionally, the 5.0% SA-1000 composite material has improved electrochemical activity because of its low impedance.^{26,27} Due to its specific capacity and Coulomb efficiency, sodium alginate effectively improves the graphite electrodes.

Subsequently, we tested the rate performance of the composite sample 5.0% SA-1000 and graphite sample under different current densities (Fig. 8d). It can be seen from the curve that the reversible capacity of the 5.0% SA-1000 composite sample is 461.197 mA h g^{-1} , 444.457 mA h g^{-1} , 386.633 mA h g^{-1} , 304.794 mA h g^{-1} , 140.846 mA h g^{-1} and 72.196 mA h g^{-1} at 100 mA g^{-1} ,



Fig. 8 Charge-discharge curve for (a) 5.0% SA-1000 and (b) graphite samples. (c) First-ring charge-discharge comparison curve. (d) Ratio performance curve of the composite sample 5.0% SA-1000 and graphite sample.

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Fig. 9



200 mA g⁻¹, 300 mA g⁻¹, 500 mA g⁻¹, 1000 mA g⁻¹ and 2000 mA g⁻¹. Compared with the graphite sample at the same magnification, its stability was significantly improved, especially at 300 mA g⁻¹ and 500 mA g⁻¹ magnification. The reversible capacities were increased by 196.911 mA h g⁻¹ and 184.818 mA h g⁻¹. The increased rate performance is attributed to the graphite coating with sodium alginate reducing the electrochemical impedance of the composite.²⁸ Additionally, the Li⁺ rapid inlay is made possible by combining the SA-1000 composite material with electrolyte at 5.0% SA.^{29,30} Therefore, the 5.0% SA-1000 composite material has better magnification performance at higher magnification.

In order to study the cyclic stability of the 5.0% SA-1000 composite samples, we tested the cyclic performance curves (Fig. 9) at a low magnification of 500 mA g^{-1} (Table 3) and a high magnification of 4000 mA g^{-1} (Table 4). The results show that the initial specific capacity of the 5.0% SA-1000 composite sample is 575.439 mA h g^{-1} . The capacity retention rate is as high as 97.90% after 100 cycles at a low rate of 50 mA g^{-1} , which is much higher than the initial specific capacity of 276.561 mA h g^{-1} and capacity retention rate of 54.14% of graphite. Especially at the high rate of 4000 mA g^{-1} current density, the initial specific capacity of graphite is very low, only

Table 3 $\,$ Cyclic data of 5.0% SA-1000 composite and graphite samples at 500 mA g^{-1}

Sample	First specific charge capacity (mA h g ⁻¹)	100th specific charge capacity (mA h g^{-1})	Capacity retention (%)
5% SA-1000	575.439	563.359	97.90
Graphite	276.561	149.731	54.14

Table 4 Cyclic data of the 5.0% SA-1000 composite and graphite samples at 4000 mA g^{-1}

Sample	First specific charge capacity (mA h g^{-1})	600th specific charge capacity (mA h g^{-1})	Capacity retention (%)
5% SA-1000	45.552	48.298	106.03
Graphite	35.362	32.891	90.49

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35.362 mA h g⁻¹, and its function basically fails. However, the initial specific capacity of the 5.0% SA-1000 composite samples at 4000 mA g⁻¹ was still maintained at 45.552 mA h g⁻¹. Even after 600 cycles, the initial specific capacity remained at 48.298 mA h g⁻¹. It can be concluded that the 5.0% SA-1000 composite sample can show better cyclic stability and a higher capacity retention rate, mainly because the coating layer of the composite sample contributes to the stable SEI film formed in the process of lithium embedding for the first time,^{31,32} which keeps the structure stable and improves the cyclic stability.^{33,34} Therefore, 5.0% SA-1000 has a significant advantage in the long cycle as the anode material of lithium-ion batteries.

The cyclic voltammetry curves of graphite and 5.0% SA-1000 samples were compared before and after coating to study the modification mechanism (Fig. 10). For the 5.0% SA-1000 composite samples, the reduction peaks at 0.01, 0.07 and 0.18 V correspond to Li⁺ embedding in the graphite layer. In comparison, the SEI of the 5.0% SA-1000 composite sample formed a lower electrical potential than that of the graphite sample (Fig. 10a), indicating that the CV curve was consistent with the microstructure of the material, which was confirmed by the XRD emission pattern and Raman spectroscopy confirmed that the 5.0% SA-1000 composite sample is composed of an internal ordered lamellar structure and an external amorphous structure. In contrast, the graphite sample has two sharp redox peaks at 0.26 V and 0.11 V, representing the insertion and extraction of the ordered graphite lamellar structure, respectively (Fig. 10b). Additionally, it is apparent from the CV curves of the front and rear circles (Fig. 10c and d) that the cyclic voltammetry curves of the 5.0% SA-1000 composite samples are larger than those of the graphite samples, indicating that the 5.0% SA-1000 composite samples will generate a larger battery capacity.

An EIS graph test was conducted to determine whether coating modification reduced the composite's electrochemical impedance (Fig. 11). The graphite sample and composite sample with 5.0% SA-1000 exhibit a similar shape composed of two semicircles.³⁵ At the 50% charged state, the charge transfer impedance³⁶ and diffusion transfer impedance³⁷ are both lower than those of the graphite samples as determined by the fitting calculation, indicating that the coating process reduces the charge

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Fig. 10 Cyclic voltammetry curve of (a) the composite material 5.0% SA-1000(b) graphite sample. (c) A comparison of the first ring of the 5.0% SA-1000 and graphite samples. (d) A comparison of the tail ring of the 5.0% SA-1000 and graphite samples.



Fig. 11 Ac impedance curves of the 5.0% SA-1000 composite and graphite samples.

transfer resistance of materials³⁸ and promotes the migration and diffusion of lithium ions.³⁹ Therefore, the 5.0% SA-1000 composite sample has a better rate performance than the graphite sample.

4. Conclusions

This paper reports the preparation of a 5.0% SA-1000 composite material by magnetic stirring, heating and mixing with deionized water, air drying and sintering at 1000 °C. After magnetic stirring and heating, drying to get a dispersed powder, and sintering under a nitrogen environment at 1000 °C to remove the crystal water in the composite, the 5.0% SA-1000 composite material was obtained. From the analysis of the whole paper, the following conclusions can be drawn:

(1) Compared with SA-800 and SA-1100, SA-1000 has a higher specific charge and discharge capacity. After 200 cycles, the capacity retention rate of SA-1000 is 84.73%.

(2) The reversibility of 2.5% SA-1000 and 7.5% SA-1000 during the first charge and discharge process is poor and below 80%, while 5.0% SA-1000 and 10% SA-1000 are above 87% when sodium alginate is coating the graphite in different proportions. In the process of converting chemical energy into electrical energy, the conversion rate is better.

(3) Compared with graphite, 5.0% SA-1000 has a larger specific surface area and is more favorable for the lithium storage performance of lithium-ion batteries. From the perspective of multiplier performance, at 300 mA g⁻¹ and 500 mA g⁻¹, the reversible capacities were increased by 196.911 mA h g⁻¹ and 184.818 mA h g⁻¹, 5.0% SA-1000 has advantages over graphite, which can be used in high-rate fast charging battery systems.

(4) In addition, 5.0% SA-1000 has more advantages in initial capacity and cycle performance, and the specific capacity of the first discharge is increased by 79.908 mA h g^{-1} after 600 cycles.

The capacity remains 48.298 mA h g^{-1} . From the perspective of battery life, it has more application value.

In the later stage of this study, further research will be conducted to explore more advanced production technology for industrial applications to serve society and to obtain broad application prospects in the field of batteries.

Author contributions

Xianfa Rao: Conceptualization, resources, methodology, project administration. Lixia Zhang: validation, form analysis, writingoriginal draft, writing-review & editing. Baobao Li: data curation, curation, investigation, validation. Xinxiong Zeng: form analysis, data curation. Wenlong Xiao: curation, visualization. YitaoLou: investigation, supervision. Huanmeng Xie: software, visualization. Huchen Yan: resources, visualization. Zixuan Yi: validation, supervision. Shengwen Zhong: conceptualization, funding acquisition.

Conflicts of interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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