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Graphene-directed two-dimensional porous carbon frameworks for high-performance lithium–sulfur battery cathodes†

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Graphene-directed two-dimensional (2D) nitrogen-doped porous carbon frameworks (GPF) as the hosts for sulfur were constructed via the ionothermal polymerization of 1,4-dicyanobenzene directed by the polyacrylonitrile functionalized graphene nanosheets. As cathodes for lithium–sulfur (Li–S) batteries, the prepared GPF/sulfur nanocomposites exhibited a high capacity up to 962 mA h g⁻¹ after 120 cycles at 2 A g⁻¹. A high reversible capacity of 591 mA h g⁻¹ was still retained even at an extremely large current density of 20 A g⁻¹. Such impressive electrochemical performance of GPF should benefit from the 2D hierarchical porous architecture with an extremely high specific surface area, which could facilitate the efficient entrapment of sulfur and polysulfides and afford rapid charge transfer, fast electronic conduction as well as intimate contact between active materials and the electrolyte during cycling.

Introduction

Motivated by the ever-growing requirement for advanced batteries, lithium–sulfur (Li–S) batteries with high theoretical energy density, cost effectiveness and environmental benignity have been lately regarded as a promising successor to lithium–ion batteries (LIBs). 1–5 Although the concept of Li–S batteries emerged in the 1960s, their large-scale commercial applications have long been precluded by the following factors: (i) fast capacity fade and unsatisfactory cycle life due to the diffusion of polysulfide (Li₂Sₙ) species in the electrolyte; (ii) the low utilization rate of active materials for the insulating nature of both sulfur and lithium sulfides (Li₂S and Li₂S₂). 6–5

To address the above-mentioned problems, numerous novel sulfur cathodes including nanoporous carbon–sulfur composites, 6–8 graphene–sulfur composites, 9–12 one dimensional (1D) carbon–sulfur composites, 13–16 polymer–sulfur composites, 17,18 and inorganic–sulfur composites (metal oxides, 19,20 metal–organic frameworks, 21 and mesoporous molecular sieves 22) have been extensively explored. Among these materials, nano-structured porous carbons are confirmed to be the most promising hosts for sulfur in Li–S batteries. Given that nano-structured porous carbons can improve the utilization of an active material by keeping sulfur particles nanometer-sized and electrically connected in their well-defined porous structure with intrinsic high electronic conductivity, 23 meanwhile, they are able to adsorb sulfur and polysulfides on their large internal surfaces and mitigate polysulfide diffusion. 24 Moreover, compared to zero-dimensional (0D) or one-dimensional (1D) counterparts, two-dimensional (2D) porous carbon materials provide large surface-to-volume ratios for better contact between active materials and the electrolyte, continuous pathways for electron conduction, short distance for charge transfer, and enormous potential in tuning the pore and channel configuration with a large specific surface area. 25,26 The rational design and construction of 2D porous carbons for energy-storage devices have attracted extensive interest and aroused favorable results. 27–30 However, developing effective templates to control the formation of 2D porous carbon frameworks remains a challenge for researchers. In this respect, functionalized graphene nanosheets such as graphene oxide (GO), reduced graphene oxide (RGO) and other graphene derivatives containing diversified functional groups are believed to be qualified templates for the construction of 2D porous carbon materials since they can direct the synthesis of porous polymers in a 2D manner, which can be further converted to porous carbons via appropriate thermal treatments. 31–33

Recent research studies of graphene-based sulfur cathodes mostly focused on graphene-coated sulfur nanoparticle composites 31 and sandwich-like graphene (oxide)–sulfur
composites. Nevertheless, further applications might be restricted due to the hindering of lithium ion transportation by the graphene lattice. Moreover, the conductivity of graphene oxide greatly depends on its oxidation degree. Therefore, it is constructive to fabricate graphene-based 2D porous carbons as the sulfur host, which will combine the merits of 2D materials with the advantages of hierarchical porous carbons. The strategy aims at realizing efficient entrapment of sulfur and polysulfides as well as facilitating electron and charge transportation to improve the cell performance.

Herein, we present the graphene-directed 2D nitrogen-doped porous carbon frameworks (denoted as GPF-n, n = 1, 2, and 3) as sulfur hosts in Li–S batteries. The 2D porous structure with a large specific surface area of up to 1683 m² g⁻¹ benefits the intimate contact between active materials and the electrolyte and provides short pathways for charge transfer. The hierarchically micro-/meso-porous carbon doped by nitrogen facilitates the chemical adsorption of polysulfide by enhancement of the surface electronegativity. As the cathode material in Li–S batteries, the nanocomposite of GPF-3 and sulfur (GPF-S-3) manifested an initial specific capacity of 1461 mA h g⁻¹ and a stabilized capacity of 962 mA h g⁻¹ after 120 discharge/charge cycles at a large current density of 2 A g⁻¹. It also exhibited an excellent rate performance by retaining a high reversible capacity of 591 mA h g⁻¹ even at an extremely large current density of 20 A g⁻¹.

**Experimental section**

**Materials**

Flake graphite was purchased from Aldrich. Acrylonitrile was of industrial polymerization grade and purchased from Xiya Reagent Co. (Chengdu, China) and used without further purification. Organic solvents were purified, dried, and distilled under dry nitrogen. All the other chemicals were purchased from Aladdin Reagent (Shanghai) and used without further purification.

**Preparation of GPF-n and PF**

Graphene oxide (GO) was synthesized from natural graphite flakes by a modified Hummers method and then reduced by hydrazine hydrate to obtain RGO. The preparation of polyaniline-functionalized graphene nanosheet (RGO-PAN) templates was based on our previous report. GPF-n was synthesized through a classic ionothermal polymerization in molten ZnCl₂. The mixtures of as-prepared RGO-PAN (30 mg), 1,4-dicyanobenzene (pDCB, 300 mg) and ZnCl₂ (3.2 g) in quartz ampoules were heated to 400 °C for 40 h, 600 °C for 20 h, or 400 °C for 20 h and then 600 °C for 20 h. The resulting products were denoted as GPF-1, GPF-2 and GPF-3, respectively. A nitrogen-doped porous carbon framework (PF) was prepared as a control sample by heating to 400 °C for 40 h without using RGO-PAN.

**Preparation of GPF-S-n and PF-S nanocomposites**

Based on a facile melt-diffusion method, GPF-n (n = 1, 2, and 3) and PF were mixed and ground separately with a certain amount of sublimed sulfur and loaded into a closed container before being heated in a muffle furnace at 153 °C for 12 h (denoted as GPF-S-n and PF-S). The theoretical maximal amounts of sulfur that GPF-n and PF can load were calculated according to the results from N₂ adsorption/desorption isothermal analysis and by the following method: the loading amount of sulfur = weight of the sample × its total pore volume × density of lithium sulfide (1.66 g cm⁻³) × the weight ratio of sulfur in lithium sulfide (69.78%). Next, GPF-1, GPF-2, GPF-3, and PF were hybridized with sulfur to obtain GPF-S-1, GPF-S-2, GPF-S-3 and PF-S with sulfur proportions of 50%, 75%, 65% and 43%, respectively.

**Characterization**

The morphology and elemental mapping information of the samples were characterized by scanning electron microscopy (SEM, Sirion 200, 25 kV) and transmission electron microscopy (TEM, JEOL JEM-2010, 200 kV). X-ray diffraction (XRD) measurements were executed on a D/max-2500/PC (Rigaku Corporation, Japan) using Cu (40 kV, 30 mA) radiation. Raman spectra were recorded on a SENTERRA with a 532 nm excitation of an Ar-ion laser with a power of about 5 mW. N₂ adsorption was measured with a Micromeritics ASAP 2010 analyzer at 77 K. The Brunauer–Emmett–Teller (BET) method and density functional theory (DFT) pore model were utilized to calculate the specific surface area and pore size distribution. Thermogravimetric analysis (TGA) of the samples was performed with a Q5000IR (TA Instruments, USA) thermogravimetric analyzer at a heating rate of 20 °C min⁻¹ under nitrogen. X-ray photoelectron spectroscopy (XPS) measurements were carried out on an AXIS Ultra DLD system from Kratos with Al Kα radiation as the X-ray source.

**Electrochemical measurements**

Each sample was mixed with carbon black (Super-P), and a polyvinylidene fluoride (PVDF) binder in 7 : 2 : 1 ratio in 1-methyl-2-pyrrolidone (NMP) solvent (Aldrich, 99.5%) then cast onto Al foil before drying in a vacuum oven at 60 °C for 12 h. All electrochemical performances were measured by using CR2016-type coin cells assembled in an argon-filled glovebox (M. BarunGlovebox) with the test electrodes by using Celgard 2400 as the separator. The electrolyte was lithium bis-(tri-fluoromethanesulfonyl)imide (LiTFSI, 1 M) with 0.1 M LiNO₃ in a 1 : 1 (v/v) mixture of 1,3-dioxolane (DOL) and 1,2-dimethyl carbonate (DMC). Electrochemical charge/discharge experiments were performed using a battery cycler (LAND-CT2001A) with current rates from 0.5 to 20 A g⁻¹ in the voltage range of 1.7–2.8 V. The cyclic voltammograms were obtained over the potential range of 1.7–2.8 V at a scanning rate of 0.1 mV s⁻¹. Electrochemical impedance spectroscopy (EIS) measurements of the electrodes were carried out on an electrochemical workstation (PARSTAT 2273). The impedance spectra were recorded by applying a sine wave with an amplitude of 5.0 mV over the frequency range from 100 kHz to 0.01 Hz. Fitting of the impedance spectra to the proposed equivalent circuit was performed by the code Z view.
Results and discussion

To ensure efficient electrolyte penetration, fast ion transfer and intimate contact between active materials and the electrolyte, a hierarchically porous carbon was constructed on the reduced graphene oxide (RGO) substrates as schematically illustrated in Fig. 1. A series of GPF-\(n\) (\(n = 1, 2,\) and 3) samples were prepared by employing the polyacrylonitrile functionalized graphene nanosheets\(^{33}\) (RGO-PAN) as the template and 1,4-dicyano-benzene (pDCB) as the precursor via ionothermal polymerization.\(^{32}\) This strategy relies on the construction of a 2D architecture induced by functionalized graphene nanosheets through the chemical bonding between templates and monomers with the introduction of nitrogen atoms throughout the porous structure. The control of porosity of the carbon matrix was achieved by adjusting the temperature and time in polymerization procedures varying from 400 °C for 20 h, 600 °C for 40 h, 1000 °C for 20 h then 600 °C for 20 h. The synthesized products were denoted as GPF-1, GPF-2, and GPF-3, respectively. For comparison, the corresponding porous carbon framework was synthesized at 400 °C for 40 h without RGO-PAN, which is denoted as PF. The sulfur impregnation of the cathode material was operated based on a facile melt-diffusion strategy\(^{39}\) with the resulting products denominated as GPF-S-1, GPF-S-2, GPF-S-3, and PF-S, respectively.

The 2D architecture of samples was confirmed by the FE-SEM and TEM analyses. The SEM image of GPF-S-3 displayed in Fig. 2a reveals that the 2D structure was well maintained after sulfur impregnation and the surface of the nanosheet was smooth without agglomerated sulfur (the morphology of GPF-3 before sulfur loading is shown in Fig. S1\(^\dagger\)). All GPF-S-\(n\) nanocomposites showed similar 2D structures while the PF without the template displayed a bulk morphology (Fig. S1\(^\dagger\)). The TEM image in Fig. 2b indicates the existence of wrinkles and folds on the GPF-S-3 nanosheet, no agglomerated sulfur can be observed either. Furthermore, the carbon, nitrogen and sulfur mapping images of GPF-S-3 suggest that nitrogen was evenly distributed throughout the carbon framework and that sulfur was homogeneously impregnated into the micro- and meso-pores of the carbon frameworks (Fig. 2c–f).\(^{40}\)

The construction of porous carbon/sulfur nanocomposites is based on the porous nature of GPF-\(n\) (\(n = 1, 2,\) and 3) and PF, which was investigated by \(N_2\) adsorption/desorption measurements at 77 K. The \(N_2\) adsorption/desorption curve of PF in Fig. 3a shows a type-I isotherm, which declares a micropore-dominated porous structure in PF.\(^{41,42}\) Interestingly, the conversion from the micropore-dominated structure in PF to the mesopore-rich textures in GPF-\(n\) (\(n = 1, 2,\) and 3) occurred with the addition of the RGO-PAN template and the changing of the polymerization time and temperature. The isotherms of GPF-1, 2 and 3 displayed more evident hysteresis loops at higher relative pressure, which indicates the formation and propagation of mesopores.\(^{41}\) This result implies that the RGO-PAN template played a fundamental guiding role in tuning the hierarchically porous configuration in GPF-\(n\) (\(n = 1, 2,\) and 3) samples. In addition, the Brunauer–Emmett–Teller specific

![Fig. 1](image1.png)

Fig. 1 Schematic illustration of the fabrication processes of GPF-S-\(n\) (\(n = 1, 2,\) and 3) nanocomposites. (a) The preparation of RGO: SDS, \(\text{N}_2\text{H}_4\cdot\text{H}_2\text{O},\) 100 °C, 10 h; (b) anionic polymerization of acrylonitrile on the surface of RGO: sodium, THF, sonication, 10 h, RT, then acrylonitrile, sonication, 10 h, RT; (c) the ionothermal polymerization of pDCB in the presence of RGO-PAN at the designed temperature and time periods in \(\text{ZnCl}_2\) (vacuum) to form GPF-\(n\); (d) melt-diffusion of sulfur into the pores of GPF-\(n\) at 155 °C for 12 h.

![Fig. 2](image2.png)

Fig. 2 (a) SEM image, (b) TEM image and (c–f) the corresponding elemental mapping images of GPF-S-3.

![Fig. 3](image3.png)

Fig. 3 (a) Nitrogen adsorption/desorption isotherms and (b) the pore size distribution of GPF-\(n\) and PF based on the density functional theory (DFT) model; (c) the pore volumes and average pore sizes of GPF-\(n\) and PF; (d) TGA curves of GPF-S-\(n\) and PF-S under nitrogen atmosphere (\(n = 1, 2,\) and 3).
The evaluation of pore size distribution of GPF-n and PF samples was investigated by the density functional theory (DFT). The curves in Fig. 3b verify that a large quantity of mesopores with the size from 2 to 4 nm existed in GPF-3, while an even larger proportion of mesopores with the size from 4 to 10 nm was exhibited in GPF-2, besides micropores in both. The relatively lower specific surface areas of GPF-2 might be ascribed to the existence of more larger-sized mesopores. The fact signifies that the adjusting of polymerization conditions exerted a significant effect on the regulation of the pore size in GPF-n. After impregnation of sulfur, each nanocomposite exhibited a dramatic fall in the specific surface area and total pore volume (Table S1†). Among them, GPF-S-3 demonstrated the most remarkable reduction (specific surface area dropped from 1683 to 14 m² g⁻¹ and the total pore volume fell from 1.57 to 0.07 cm³ g⁻¹). The results indicate that these carbon hosts with a hierarchically porous structure and pretty high specific surface areas possessed an efficient entrapment of elemental sulfur. In addition, the total pore volumes of GPF-S-n and PF-S are smaller than their theoretical values, which could be attributed to the possibility of water entrapment during the sulfur loading. Furthermore, the micro-pores in GPF-n and PF are likely to be blocked and form small enclosed spaces, which could not be reached in the N₂ adsorption/desorption tests and inevitably lead to lower detected results. The contents of sulfur in the nanocomposites were confirmed by thermogravimetric analysis (TGA) under a nitrogen atmosphere (Fig. 3d). The obvious weight loss of PF-S and GPF-S-n (n = 1, 2, and 3) caused by sulfur vaporization happened between 250 and 350 °C, much higher than the temperature range when pure sulfur volatilizes (160–270 °C), indicating the strong adsorption of sulfur in the abundant micro-/meso-pores of carbon matrices. Furthermore, the TGA results demonstrated an increase in sulfur loading of sulfur, the Raman spectroscopy was conducted to further characterize the structural rearrangement in porous frameworks during the polymerization process with optimized temperature and reaction time.\(^{38,41–46}\)

![Image](310x69 to 546x155)

The XRD patterns of GPF-S-1, 2, and 3) and PF in Fig. S2b† displays two peaks at 1350 and 1580 cm⁻¹, which are assigned to D (disordered, defect-activated) and G (graphitic, in-plane stretching of sp² bonds) bands of carbon,\(^{13}\) respectively. After the loading of sulfur, the Raman spectra of GPF-S-n (n = 1, 2 and 3) and PF-S (Fig. 4b) exhibited the exactly similar characteristic peaks with their original samples, GPF-n (n = 1, 2 and 3) and PF. They even showed very similar integral intensity ratios, \(I_D/I_G\), as labeled above each spectrum in Fig. 4b and S2b.† Hence the fact signifies that the sulfur loading did not change the structural characteristics and the graphite layer defectiveness of carbon matrices. Furthermore, no characteristic peaks of elemental sulfur were observed in the region between 100 and 500 cm⁻¹ in Raman spectra of PF-S and GPF-S-n (n = 1, 2, and 3), which are related to vibration of the S–S bond in S₈ species (Fig. S2d†). This observation suggests that sulfur particles were impregnated into pores and adsorbed on the internal surface of pores in GPF-n and PF hosts,\(^{44}\) which was in good accordance with the results from XRD spectra and the sulfur mapping image.

X-ray photoelectron spectroscopy (XPS) analyses (Fig. 5 and S3†) revealed that the nitrogen atom proportion in GPF-S-1, GPF-S-2, and GPF-S-3 were 6.62, 4.27 and 4.20 at%, respectively. The N 1s spectra of GPF-S-3 (Fig. 5a) and GPF-S-1, 2 (Fig. S3 in ESI†) depicted that four distinct nitrogen configurations, pyridine \(N\) (398.3 eV, N1), pyrrole \(N\) (400.1 eV, N2), quaternary \(N\) (401.1 eV, N3), and pyridine-N-oxide (403.1 eV, N4), existed in these nitrogen doped carbon matrices,\(^{32,44,55}\) suggesting the formation of graphitized fragments. Among them, the N 1s signal with binding energy (BE) at about 401.1 eV (N3) is assigned to quaternary nitrogen, namely graphite-like nitrogen, which is believed to enhance the conductivity of N-doped carbons by providing additional electrons to the delocalized \(\pi\)-system.\(^{56}\) Quantitative analysis based on XPS suggests that the ratios of N3 gradually increases while the ratio of N1 decreases.

![Image](317x642 to 539x729)

![Image](513x912 to 635x1087)

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in GPF-S-1, GPF-S-2 and GPF-S-3, which confirms the occurrence of rearrangement reactions and a gradual increase of graphitized fragments. In addition, XPS has been successfully used to identify the chemical state of sulfur molecules and distinguish the sulfur types in Li–S batteries. All three S 2p spectra in Fig. 5b and S3 contain the characteristic sulfur splitting of the S 2p signal into two components S 2p3/2 and S 2p1/2 as a result of spin–orbital coupling effect at 163.6 and 164.9 eV, which suggests the existence of S8 molecules in GPF-S-n samples. The two more peaks at 168.7 and 169.9 eV indicate the existence of sulfur atoms located at the chain end of the small S2–4 molecules, further proving the strengthened interaction between sulfur molecules and 2D porous carbon hosts. This finding implies that the effective adjustment of pore architecture can implement molecule level regulation of sulfur molecules in micro-/meso-pores.

To evaluate the potential of GPF-S-n and PF-S as Li–S cathodes, coin cells were fabricated by the method described in the experimental section. Their electrochemical performance for S electrodes at various current densities from 0.5 to 5 A g\(^{-1}\) is illustrated in its galvanostatic discharge–charge profiles (Fig. 6b). Consistent with the peak voltages in the CV curves, these discharge voltage profiles demonstrated the two-plateau characteristic of a typical Li–S cell at different current densities from 0.5 A g\(^{-1}\) to 5 A g\(^{-1}\), which further proves the excellent capacity reversibility of the GPF-S-3 cathode. Furthermore, the GPF-S-n cathodes exhibited favourable cycling performance at a constant current rate of 2 A g\(^{-1}\) (Fig. 6c). Among the samples, the GPF-S-3 cathode achieved an initial discharge and charge capacity of 1461 and 1424 mA h g\(^{-1}\) with the first cycle coulombic efficiency (CE) up to 97.5%. The PF-S cathode, by contrast, obtained initial discharge and charge capacities of 1435 and 1227 mA h g\(^{-1}\) with the first cycle CE of only 85.5%. After 120 cycles, GPF-S-1, GPF-S-2 and GPF-S-3 cathodes still sustained stabilized specific capacities of 791, 706 and 962 mA h g\(^{-1}\), respectively, greatly surpassing a capacity of 386 mA h g\(^{-1}\) for the PF-S cathode. GPF-S-1, GPF-S-2 and GPF-S-3 cathodes also exhibited better rate performance than PF-S cathode as depicted in Fig. 6d. A high reversible capacity of 810 mA h g\(^{-1}\) was still preserved even at a high rate of 5.0 A g\(^{-1}\) for the GPF-S-3 cathode, which was higher than 710 mA h g\(^{-1}\) for GPF-S-2, 696 mA h g\(^{-1}\) for GPF-S-1 and much higher than 317 mA h g\(^{-1}\) for PF-S. The recovery of a reversible capacity of 1040 mA h g\(^{-1}\) for GPF-S-3 was also achieved at 0.5 A g\(^{-1}\) following a series of high rate charge–discharge processes. The promising cycling and rate performance of GPF-S-n could be ascribed to their 2D architecture with RGO as the template, which guaranteed a rapid ion transfer, fast electronic conduction and intimate contact between the active materials and electrolyte. In addition, the functionalized RGO templates could provide stable frameworks to sustain the strain from the volumetric change of active sulfur and help GPF-S-n to preserve the conductive matrices well during the charge/discharge cycles. The superior electrochemical performance of GPF-S-3 among three GPF-S-n cathodes might be attributed to the abundant hierarchical micro-/mesopores with an optimal pore size in its carbon matrix and nitrogen-doped inner surface, which can increase the carbon surface affinity to active sulfur. This feature could contribute to the effective entrapment of active sulfur and polysulfides in carbon hosts. Interestingly, the GPF-S-2 cathode with the...
highest sulfur loading of 72 wt% presented inferior specific capacities to those of GPF-S-3 with a sulfur loading of 63 wt% under the same testing conditions, which might be attributed to the weaker entrapment of sulfur and polysulfides inside the relatively larger-sized (over 4 nm) mesopores in the GPF-2 framework (Fig. 3b) So, both 2D architecture and optimized micro-/meso-pore structure are indispensable to guarantee a fast electronic/ionic transport and enhanced reaction kinetics of sulfur in a nanocomposite skeleton. The improved rate charge–discharge capability and cycling performance of GPF-S-3 were further confirmed via a Nyquist plot and its fitted results of an equivalent circuit depicted in Fig. S4 † The cell with the GPF-S-3 cathode demonstrated smaller impedance of a cell component ($R_1 = 18.52 \Omega$) than that with PF-S ($R_1 = 56.89 \Omega$) as shown in Table S3 † which might be due to the stronger confinement of active materials and less dissolution of polysulfides. Moreover, the cell with the GPF-S-3 cathode exhibited a pretty low charge transfer impedance ($R_{ct}$) value of 24.33 $\Omega$ compared with 146.5 $\Omega$ for the cell with the PF-S cathode, suggesting that the former one preserved highly rapid charge transportation kinetics benefited from its 2D architecture as well as the uniform distribution of sulfur in a hierarchically well-tailored porous carbon matrix. To simulate the realistic situation where a faster discharge/charge process is adopted, the high-rate performance tests for GPF-S-3 and PF-S cathodes were consequently carried out at current densities from 3 to 20 A g$^{-1}$. As illustrated in Fig. 6e, GPF-S-3 displayed extremely high specific capacities of 753, 652, and 591 mA h g$^{-1}$ even at very high current densities of 10, 15, and 20 A g$^{-1}$. After cycling at an ultrahigh current density of 20 A g$^{-1}$ and being reverted to 3 A g$^{-1}$ again, GPF-S-3 still achieved both an outstanding reversible capacity of 960 mA h g$^{-1}$ and excellent CE of 99.9%. The PF-S electrode, by contrast, exhibited reversible capacities of only 95, 21, and 16 mA h g$^{-1}$ at 10, 15, and 20 A g$^{-1}$ respectively. Such an impressive performance of the GPF-3 host for sulfur preceded most of the state-of-the-art porous carbon host materials for Li–S battery cathodes (see Table S4 †), including hollow carbon spheres, layered graphene-based porous carbon, nanoarchitected graphene/CNT@porous carbon, and the like.

Conclusions

In this work, 2D GPF-n ($n = 1$, 2 and 3) with a well-tailored porous structure and high specific surface area were synthesized via in situ ionothermal polymerization of 1,4-dicyanobenzene directed by the functionalized graphene nanosheets as sulfur hosts for the cathodes in Li–S batteries. The as-prepared GPF/sulfur cathodes exhibited a high capacity up to 962 mA h g$^{-1}$ after 120 cycles at 2 A g$^{-1}$. A high reversible capacity of 591 mA h g$^{-1}$ was still retained even at an extremely large current density of 20 A g$^{-1}$. Such an outstanding rate and cycling performance demonstrate that the construction of 2D hierarchically micro-/meso-porous carbon frameworks as sulfur hosts is an instructive and efficient strategy to fabricate Li–S batteries for the practical application of high power devices. It is expected that the synthesis strategy can provide a platform for the preparation of unprecedented hierarchical porous carbons by utilizing various monomers and diversified functionalized templates for sulfur hosts in high performance Li–S batteries.

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Notes and references
