Bipolar nitrogen-doped graphene frameworks as high-performance cathodes for lithium ion batteries†

Yanshan Huang,ab Dongqing Wu,*a Arezoo Dianat,c Manferd Bobeth,c Tao Huang,a Yiyong Mai,a Fan Zhang,*a Gianaurelio Cuniberti and Xinliang Fengad

Hierarchically porous nitrogen-doped graphene frameworks (N-GFs) are fabricated through the ice-templating of GO with polyethylenimine and the thermal treatment of the resultant hybrids. As cathode materials in lithium ion batteries (LIBs), the obtained N-GFs exhibit an outstanding specific capacity of 379 mA h g⁻¹ at 0.5 A g⁻¹ for 2500 cycles. Even at an ultrahigh current density of 5 A g⁻¹, the N-GFs maintain a capacity of 94 mA h g⁻¹, superior to that of most reported LIB cathode materials. The experimental results and quantum mechanics calculations suggest that pyridinic-like N and pyridinic N-oxide in graphene are responsible for the excellent cathodic performance of the bipolar N-GFs by providing fast surface faradaic reactions with both p- and n-doped states.

Introduction

As a crucial factor in determining the electrochemical performance of lithium ion batteries (LIBs), cathode materials with high specific capacity and rate capability are essential for satisfying the increasing demand for miniaturized LIBs with improved energy density.1,2 The cathode materials in present commercial LIBs are mainly based on inorganic intercalation compounds such as lithium manganese oxide (LiMn₂O₄) and lithium iron phosphate (LiFePO₄). In addition to their high costs, these inorganic LIB cathodes have disadvantages of limited mineral resources, low theoretical capacities, and complicated manufacturing processes. As an alternative to inorganic intercalation materials, organic LIB cathodes including n-type (such as organo-disulphide and conjugated carbonyl compounds), p-type (conjugated amine and conjugated thiourea compounds), and bipolar organics (conjugated hydrocarbon and nitroxy radical) have recently received considerable attention because of their relatively low cost, easy processability and the molecular-level adjustable electrochemical properties.3–14 Compared to n- and p-type organic electrodes, the neutralize state of bipolar organic electrodes can be continuously oxidized by anions (such as ClO₄⁻, PF₆⁻, BF₄⁻ or N(CF₃SO₂)₂⁻ in the electrolyte) with high redox potential and reduced by cations (Li⁺) with low redox potential, respectively.15–17 Such a bipolar redox reaction with a linear transition from each doped state in the energy storing process can result in a remarkably high specific capacity. However, the major problems associated with organic cathode materials are their low electrical conductivity and high solubility in electrolytes, which hinder their utilization in high rate operation and long-cycle process of LIBs.

Profiting from its excellent conductivity and high specific surface area, graphene has been extensively applied as a matrix to improve the conductivity of inorganic or organic cathode materials, thereby boosting their electrochemical performances.15–17 However, the structural defects of graphene in such hybrid electrodes inevitably retard Li diffusion during intercalation and the capacity of such hybrids is still almost lower than 200 mA h g⁻¹.18–21 Several recent publications demonstrated that graphene oxide (GO) alone can serve as the cathode materials in LIBs and its Li storage properties are highly sensitive to the oxygen-functional groups.22,23 Nevertheless, the presence of oxygenated groups in the plane or at the edge of graphene also reduces the electrical conductivity, thus influencing the charge and ion transport behavior. Therefore, the tradeoff between the capacity and the conductivity must be addressed to enhance the effectiveness of graphene as a LIB cathode. By contrast, nitrogen (N)-doped graphene can both boost the conductivity of graphene and improve the wettability of graphene with an electrolyte. Previous studies have focused only on applications of N-doped graphene as LIB anodes,24,25 supercapacitors26 and electrocatalysts.27–29 In this study, for the
first time, we demonstrate that N-doped graphene can function as a LIB cathode with excellent capacity and cycle stability. Hierarchically porous N-doped graphene frameworks (N-GFs) were fabricated through the ice-templating of GO with poly-ethylenimine (PEI) and the following thermal treatment of the resulting composites. With a N doping content of ~5 at% and in the presence of pyridinic nitrogen and the pyridinic-oxide nitrogen, the resulting bipolar type N-GFs exhibit an outstanding specific capacity of 379 mA h g\(^{-1}\) at 0.5 A g\(^{-1}\) for 2500 cycles. Even at an ultrahigh current density of 5 A g\(^{-1}\), the N-GFs maintain a capacity of 94 mA h g\(^{-1}\), superior to that of most reported organic and inorganic LIB cathode materials (Table S1†).\(^{11,29-34}\) The strong affinity of lithium ions to active nitrogen species in the hybrids is verified by quantum mechanics calculations. All the results suggest that pyridinic-like nitrogen and pyridinic N-oxide in graphene provide fast surface faradaic reactions with both p- and n-doped states, which are responsible for the excellent cathodic performance of the N-GFs.

**Experimental section**

**Preparation of N-GFs**

Graphene oxide (GO) was synthesized from natural graphite flakes by a modified Hummers method. PEI (5 mg) was dissolved in the suspension of GO (100 mL, 1 mg mL\(^{-1}\)). The mixture was stirred for 5 h. After that, the as-prepared suspension was washed and centrifuged with water 3 times. The resulting mixture was then placed in a 1 mL insulin syringe, which was then immersed in liquid nitrogen. The unidirectional frozen cryogels were then freeze-dried for 48 h, and removed from the syringe as intact monoliths. Finally, the as-prepared suspensions were thermally treated at 300, 450, and 600 °C in N\(_2\) for 2 h, respectively. And with the different thermal treatment temperatures, the samples were denoted as N-GF-300, N-GF-450, and N-GF-600, respectively. Instead of PEI, aqueous ammonia (NH\(_3\)) and chitosan were also utilized as the nitrogen source to prepare N-GFs, and the resulting hybrids were denoted as N-GF-450-A (from NH\(_3\)) and N-GF-450-C (from chitosan), respectively. The weight ratios of nitrogen sources and GO were both 1 : 20. The fabrication processes of N-GF-450-A and N-GF-450-C were similar to that of N-GF-450. As a control experiment, GF-450 without N-doping was also synthesized by the following process. The suspension of 100 mg, 5 mg mL\(^{-1}\) GO was placed in a vial and then immersed in liquid nitrogen. The unidirectional frozen cryogels were then freeze-dried for 48 h. Finally, the GF was thermally treated at 450 °C in N\(_2\) for 2 h.

**Characterization**

Scanning electron microscopy (SEM) measurements were performed on an FEI Sirion-200 field emission scanning electron microscope. Transition electron microscopy (TEM) studies were conducted on a JEOL-2100 electron microscope at an operating voltage of 200 kV. The samples were dispersed in water and the suspension was dropped onto a copper grid covered with a carbon film. The X-ray diffraction (XRD) analysis was performed on a Rigaku D/Max 2500 X-ray diffractometer with Cu K\(_\alpha\) radiation (\(\lambda = 1.54 \text{ Å}\)) at a generator voltage of 40 kV and a generator current of 50 mA with a scanning speed of 5° min\(^{-1}\) from 10 to 80°. Nitrogen adsorption/desorption isotherms at 77 K were determined by using a Micromeritics ASAP 2010. X-ray photoelectron spectroscopy (XPS) experiments were carried out on an AXIS Ultra DLD system from Kratos with Al K\(_\alpha\) radiation as the X-ray source for radiation. Elemental analysis was applied for the determination of C/N on a Vario EL Cube. Fourier transform infrared (FTIR) spectra were recorded on a NEXUS 670 spectrometer. Raman spectra were recorded on a SENTERRA with excitation from the 532 nm line of an Ar-ion laser with a power of about 5 mW.

**Electrochemical measurements**

The cathode electrodes were prepared by mixing the hybrids, carbon black (Super-P), and poly(vinyl difluoride) (PVDF) with a weight ratio of 80 : 10 : 10 and pasting the mixture on aluminum foil with a thickness of 100 μm. Pure lithium foil was used as the counter-electrode. The electrolyte consisting of a solution of 1 M LiPF\(_6\) in ethylene carbonate (EC)/dimethyl carbonate (DMC) (1 : 1 by volume) was obtained from Ube Industries Ltd. The cells were assembled in an argon-filled glovebox with the concentrations of moisture and oxygen below 1 ppm. The electrochemical performance was tested at various rates in the voltage range of 1.50–4.50 V. The active material loading in these electrodes is about 0.6 mg cm\(^{-2}\).

**Computational details**

DFT calculations were performed with the Gaussian 09 program. All geometry optimizations were carried out at the RB3LYP level of density functional theory with the 6-31G(d) basis set. To compute the electronic charge density, a mixed basis-set approach was used here, where the Kohn–Sham orbitals are expanded into linear combinations of contracted Gaussian type orbitals (GTO) and complemented by a plane-wave basis set. In particular, for geometry optimization, we applied a DZVP (double zeta for valence electrons plus polarization functions) basis set complemented with a plane-wave basis with energy cut-off at 350 Ry. In all calculations, we employed the BLYP exchange–correlation functional and its corresponding norm-conserving pseudo-potential GTH (Goedecker, Teter and Hutter). Dispersion corrections were included through the standard D3 Grimme parameterization. Afterwards, all energies were computed by single point calculations with the B3LYP exchange–correlation functional. The convergence criteria for both geometry and energy calculations were set to 1 × 10\(^{-7}\) Hartree for the SCF energy and 9 × 10\(^{-4}\) Hartree per Å, respectively.

**Results and discussion**

Scheme 1 illustrates the overall procedure for the fabrication of N-GFs. PEI is first added to the aqueous suspension of GO, and the rich amine groups on PEI enable it to anchor to the hydroxyl...
and carboxyl groups on GO through hydrogen bonding, forming PEI–GO.\textsuperscript{35} When the dispersion of PEI–GO is plunged into liquid nitrogen, the water in the suspension rapidly solidifies into ice crystals, which expel the PEI–GO nanosheets to form a three-dimensional (3D) framework along the solidification front. After the freeze-drying process, a macroscopic free-standing monolith (PEI-GOM) with a continuous and interconnected network architecture is formed. Finally, the thermal treatment of PEI-GO in a nitrogen atmosphere converts to N-doped graphene frameworks (N-GFs). During the process, PEI does not only serve as a cross-linker for binding GO sheets into a highly interconnected monolith in the ice-templating step. It also provides a nitrogen source for the N-doping in the graphene frameworks.\textsuperscript{36} The resulting N-GFs are denoted as N-GF-300, N-GF-450, and N-GF-600 according to the thermal treatment temperature at 300, 450, and 600 °C, respectively. In addition to PEI, NH\textsubscript{3} and chitosan are also selected as nitrogen sources for the fabrication of N-GFs (ESI†). The resulting samples are denoted as N-GF-450-A (from NH\textsubscript{3}) and N-GF-450-C (from chitosan).

Cross-sectional scanning electron microscopy (SEM) images of the as-prepared N-GFs (Fig. 1a, b and S1†) indicate that all of the samples have highly interconnected 3D macroporous structures with diameters ranging from 5 to 20 \( \mu \text{m} \). Moreover, alternative dark and light domains with different contrast levels are evident on the N-GF surfaces through transmission electron microscopy (TEM) and high-resolution TEM (HR-TEM), which are attributable to the defects of the N-doped graphene framework (Fig. 1c and d). Furthermore, elemental mapping images of N-GF-450 show that C, N, and O atoms are homogeneously distributed in the N-GF (Fig. 1e–h). In addition, nitrogen physisorption measurements were conducted to gain insight into the porosities of N-GFs (Fig. S2a†). All samples exhibit type IV adsorption branches associated with a well-defined capillary condensation step. The Brunauer–Emmett–Teller (BET) surface areas of N-GF-300, N-GF-450, and N-GF-600 are 77, 111, and 121 m\(^2\) g\(^{-1}\), respectively. Furthermore, the existence of a high number of mesopores and micropores in N-GF-450 and N-GF-600 was verified according to the corresponding pore size distribution based on density functional theory (DFT, Fig. S2b†). Compared to N-GF-300, the considerably enhanced surface areas of N-GF-450 and N-GF-600 might be attributable to the micro- and mesopores generated through the conversion of PEI to N-doped carbon in the graphene frameworks.

According to the X-ray photoelectron spectroscopy (XPS) results, the N contents of N-GF-300, N-GF-450, and N-GF-600 are calculated to be 4.16, 5.14, and 4.31 at\%, respectively, which are in agreement with the elemental analysis results (Table S1†). Moreover, the high resolution N1s and C1s and O1s spectra of the samples are shown in Fig. S3.† The N 1s spectra of N-GFs can be fitted to four peaks at approximately 398.9, 400.2, 401.0, and 402.6 eV, corresponding to the pyridinic nitrogen (N1), pyrrole nitrogen (N2), graphitic nitrogen (N3), and pyridinic N-oxide (N4), respectively (Fig. S3†).\textsuperscript{27} As summarized in Fig. S3c† the amounts of different N atoms in the N-GFs change with the thermal treatment temperature. The N4 in N-GF-300 and N-GF-450 is attributed to N1 coupled with the remaining O atoms on the graphene surface after thermal reduction at 300 or 450 °C. However, the amount of O atoms in N-GF-600 is considerably reduced and not sufficient to form N4. The C1s peak is dominated by a C–C bond at 284.8 eV. A small amount of carbonyl carbon (C=O, 290 eV) can also be detected from the XPS data for the O1s (531.5 eV). This type of the functional group on the graphene surface is well known to increase the capacity of the engineered graphene for its fast reaction with lithium.\textsuperscript{38}
Cyclic voltammetry measurement (CV) was conducted to investigate the fundamental electrochemical behavior of the N-GFs ranging from 1.5 to 4.5 V versus Li/Li⁺ by using 1 M LiPF₆ in ethylene carbonate/dimethyl carbonate (1 : 1) as the electrolyte at room temperature (Fig. 2a–c and S6†). Clearly, N-GFs exhibit the considerably improved electrochemical reactions of association and disassociation with Li⁺ compared with undoped graphene.⁶⁹ Furthermore, the electrochemical reaction of N-GF-450 is quite more active than those of N-GF-300 and N-GF-600. According to the aforementioned composition analysis, the doped N and O atoms are responsible for the electrochemical reaction. Among three samples, N-GF-600 with the least of the total of N and O contents shows the lowest electrochemical activity. With the most O content, the N-GF-300 does not show much higher gravimetric current and capacitance than N-GF-600. However, for N-GF-450, with the highest N contents, both the gravimetric current and capacitance are improved. The results indicate that the capacity of the sample is influenced by heteroatoms (mostly by N atoms) in their graphene framework. In particular, N1 and N4 on the N-GFs provide greater contribution than those of the other N species. To evaluate the reversible association and disassociation of Li⁺ in N-GFs, the CV of N-GF-450 at a slower scan rate of 0.1 mV s⁻¹ is shown in Fig. 2b. In the initial cycle, three couples of redox peaks are observed at about 1.7/3.3, 3.8/4.1 and 4.2/4.3 V, resulting from the association and disassociation of Li⁺ with the heteroatoms (N and O) on the electrode materials. In the second cycle, the two cathodic peaks at 1.7 and 3.8 V shift to the higher potential at 1.8 and 3.9 V, respectively, hence reducing hysteresis between the anodic and cathodic peaks. This result indicates the good reversibility of the electrode.⁶⁹ The redox reactions happening below and above the open-circuit voltage of 3 V suggest that the electrode has reversible p-doped (an oxidized state above 3 V vs. Li/Li⁺) and n-doped (a reduced state below 3 V vs. Li/Li⁺) redox activity. On the basis of the previously proposed lithium storage mechanism for N-rich organic electrodes,⁴⁴ energy storage by N-GFs occurs through the reversible redox reaction accompanied by the association and disassociation of Li⁺ or electrolyte anions (PF₆⁻); this mechanism can be defined as reaction (1) and (2):⁴¹–⁴³

\[
(N4)^{x+}(PF_6^{-})_y + xLi^+ + xe^- \rightarrow (N1)^{x+}(PF_6^{-})_y \quad (1)
\]

\[
(N1) + yLi^+ + ye^- \rightarrow [(N1)^{y+}(Li^+)]_y \quad (2)
\]

The PF₆⁻ electrochemically doped in N-GF-450 at a full charge state (4.5 V, p-doped region) can be detected via FTIR spectra (Fig. S7a†). The coordination interaction between Li and N lone-pair electrons (Li–N bonding) is confirmed through XPS after full discharge to 1.5 V (Fig. S7b†), which indicates that the N1 and N4 atoms in the N-GFs are responsible for bipolar faradaic reactions to accommodate Li ions, and thus hold potential for the LIB cathode materials. Certainly, the C=O also provides the capacity for lithium storage by faradaic reactions with lithium ions (3), similar to previously reported composites of carbon nanotubes.⁴⁸,⁴⁴,⁴⁵ Thus a large working potential window (1.5–4.5 V versus Li/Li⁺) was achieved, which can thereby increase the energy density by yielding a high specific capacity, as shown in the typical profiles of voltage versus specific capacity at a current density of 0.5 A g⁻¹ (Fig. S6a†). Furthermore, the CV profiles of N-GF-450 show a gradual deviation from the ideal rectangular shape with increased scan rate from 10 to 100 mV s⁻¹ (Fig. 2a, S6b† and 2b), indicating the existence of both the redox reaction and double layer capacitance in the sample. In addition, the gravimetric current and the capacity increased considerably in the voltage window of 1.5–4.5 V vs. Li (blue solid curve), compared with a restricted voltage window of 1.5–3.0 V vs. Li (black dashed dotted curve) or 3.0–4.5 V vs. Li (red dotted curve) (Fig. 2c). The results indicate that the electrode cycling over the full voltage range yields much higher gravimetric currents from the faradaic reactions than double-layer capacitance.⁴¹,⁴⁴

The galvanostatic discharge-charge measurements were performed on half cells at a current density of 0.5 A g⁻¹. The capacity of N-GF-450 gradually increases from 163 to 370 mA h g⁻¹ with almost 100% coulombic efficiency after 2500 cycles (Fig. 2d). Nevertheless, the capacities of N-GF-300 and N-GF-600 under the same conditions decrease to 60 and 40 mA h g⁻¹, respectively. Notably, the capacity of the N-GF-450-based cathode remains at 250 mA h g⁻¹ at a current density of 1 A g⁻¹.
The electrochemical impedance spectroscopy spectra of the samples resistance within the electrode, as verified according to the electrochemical impedance spectroscopy spectra of the samples (Fig. S9†). According to the equivalent circuit (inset Fig. S9†), N-GF-450 has lower contact resistance ($R_L = 4.0 \Omega$) and charge-transfer resistance ($R_{ct} = 429.9 \Omega$) than other samples (Table S2†). By contrast, N-GF-600 without N4 species exhibits the highest charge-transfer resistance among the three N-GFs, which explains its low electrochemical activity as the LIB cathode.

The redox properties of N1, N2 and N3 species were estimated using density functional theory (DFT) calculation (Fig. S10†). The lowest-unoccupied molecular orbital (LUMO) energy level of the N1 (−1.742 eV) is lower than those of N2 (−1.611 eV) and N3 (1.646 eV). The net energy change of N1 (0.62 eV) is the highest compared to those of N2 and N3 (−1.37 and 0.14 eV, respectively), leading to the highest activity for association of Li atoms. The deeper insight into the association of Li ions with the different species of N atoms in the N-GFs is further supported through quantum mechanics and calculations. For our calculations, the association of Li ions with N-doped graphene was investigated within the framework of DFT by implementing the standard implementation in the CP2K software package.57,58 An N-doped graphene sheet with a total nitrogen doping concentration of 5 at%, of which N1, N2, and N3 atoms constituted approximately 57.4%, 35%, and 7.6%, respectively, was used in the simulation (Fig. 3a). As a first step, the interaction of Li atoms with N-doped graphene at different adsorption positions was investigated. The lithium adsorption energy was defined as:

$$E_{ads} = E_{Li@gr} - E_{gr} - E_{Li(gas phase)}$$  \(\text{(4)}\)

The Li adsorption energies and the corresponding adsorption positions are illustrated in Fig. 3b. The most stable adsorption position is that of the Li atom between the N1 and N2 atoms, followed by the hollow sites of benzene and pyridinic rings. The Li atom on top of the N3 atom has the lowest adsorption energy. To evaluate the electrochemical activity of the Li/N-doped-graphene system, the change in the partial charge of the Li ions was calculated through Bader charge analysis. For Li atoms adsorbed between the N1 and N2 atoms, a Li partial charge of approximately +0.44e was used as an example.

Battery voltages or potential differences between the bulk lithium metal as the anode and N-doped graphene with different Li concentrations as the cathode were calculated. The voltage between the two electrodes was derived from the difference in the chemical potential of Li in the anode and cathode. At zero temperature, the average voltage $V$ was obtained from the total energy through the formula:

$$V = \frac{E(Li_{x1}/N-Gr) - E(Li_{x2}/N-Gr)}{e(x_2 - x_1)}$$  \(\text{(5)}\)

where $E(Li_{x1}/N-Gr)$ and $E(Li_{x2}/N-Gr)$ are the total energies of the systems with Li concentrations $x_1$ and $x_2$, respectively. $E_{bulk}(Li)$ is the total energy of the Li metal in the bcc structure. The concentrations of the Li atoms with adsorption sites between the N1 and N2 atoms on the upper and lower sides of the graphene sheet gradually increase, followed by those of atoms with the second, third, and fourth most favorable adsorption positions (inset of Fig. 3b). Except for extremely low Li concentrations, the average voltages are between 2 and 3 V. This value is in close agreement with the measurements of the open circuit voltage between the two electrodes.
Conclusions

Herein, 3D macroporous GFs coupled with N atoms were successfully developed as the cathodes for rechargeable LIBs with high capacity and a long cycle life. And their energy storage principle was further investigated by experiment and calculation, which implies that the excellent cathodic performance of the hybrids is mainly due to pyridinic N and pyridinic N-oxide in graphene. More importantly, the effect of nitrogen doping demonstrated in our study will inspire the design and development of various heteroatom doped carbon-based cathode materials for LIBs in future.

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