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1	A Novel High Conductive Ferroferric Oxide/Porous Carbon Nanofibers		
2	Composites Prepared by Electrospinning as Anode Materials for High		
3	Performance Li-ion Batteries		
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10	Abstract: In this paper, ferroferric oxide (Fe <sub>3</sub> O <sub>4</sub> ) nanoparticles/porous carbon nanofibers		
11	(Fe <sub>3</sub> O <sub>4</sub> /PCNFs) composites were successfully fabricated by elctrospinning and subsequent		
12	calcinations. The composites were characterized by X-ray diffraction, thermogravimetric		
13	analysis, scanning electron microscopy and transmission electron microscopy to analyze the		
14	structure, composition and morphology. The electrochemical performance was evaluated by		
15	coin-type cells versus metallic lithium. The results indicated that Fe <sub>3</sub> O <sub>4</sub> /PCNFs composites		
16	exhibited high reversible capacity and good capacity retention. The discharge capacity		
17	maintained 717.2 mA h g <sup>-1</sup> at 0.5 A g <sup>-1</sup> after 100 cycles. The excellent performances of		
18	Fe <sub>3</sub> O <sub>4</sub> /PCNFs composites are attributed to good crystallinity and uniformly dispersive Fe <sub>3</sub> O <sub>4</sub>		
19	nanoparticles, and porous carbon shell with high conductivity. The carbon coating buffered		
20	the tremendous volumetric changes between $Fe_3O_4$ nanoparticles and Fe atoms in the		
21	charge/discharge processes and kept the structure integrity of Fe <sub>3</sub> O <sub>4</sub> nanoparticles. Porous		
22	carbon nanofibers prepared by unique calcination process improved the conductivity of		
23	composites and provided free space for migration of lithium ions. The preparation of strategy		
24	is expected to be applied to the preparation of other transition metal oxides materials as		
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superior anode materials for lithium-ion batteries.

26 Keywords: Porous nanofiber; Fe<sub>3</sub>O<sub>4</sub> nanoparticles; Electrospinning; Hybrid anode materials;

27 Li-ion battery

28

# 29 1 INTRODUCTION

Haze and greenhouse effect have become severe challenges to environment, which are 30 closely connected to excessive use of fossil fuels <sup>[1-3]</sup>. Rechargeable lithium ion batteries are 31 tools of energy exchange between chemistry energy and electric energy <sup>[3-4]</sup>, which can store 32 energy from clean energy <sup>[5]</sup>. As rechargeable lithium ion batteries have the characteristics of 33 high energy density, long lifespan, environment friendly and fast charge-discharge rates <sup>[6-7]</sup>, 34 they have been widely applied to mobile phones, digital cameras and laptops, etc <sup>[7]</sup>. But 35 hybrid electric vehicles and electric vehicles require Li-ion batteries to have higher energy 36 density and rate capability <sup>[3,8]</sup> to match the performance of internal combustion vehicles <sup>[3]</sup>. 37

Natural graphite seems to be the most promising candidate for the anode material in 38 lithium-ion batteries because of its numerous advantages <sup>[9,10]</sup>. Graphite and graphitized 39 carbon as anode materials in lithium-ion batteries have been used in many commercial 40 products on the market <sup>[9]</sup>. Modification of carbonaceous anode materials which has been a 41 research focus <sup>[11]</sup> can greatly improve materials' electrochemical performance. Composites 42 of carbon nanomaterials and sulfides [12-16]/metal oxides [17-21] have been intensively studied 43 for efficient energy storage. Transition metal oxides (MO) (Fe<sub>2</sub>O<sub>3</sub>, Fe<sub>3</sub>O<sub>4</sub>, NiO, CoO, Co<sub>3</sub>O<sub>4</sub>, 44 Cu<sub>2</sub>O, CuO, RuO<sub>2</sub> and Cr<sub>2</sub>O<sub>3</sub> *etc.*) have much higher theoretical capacities (~1000 mA h g<sup>-1</sup>) 45 than graphite based on the conversion between MO and M<sup>[17]</sup>. But pure transition metal 46 oxides as anodes in lithium ion batteries often have poor cycling performance owing to the 47 collapse of lattice structure of the original crystal over several discharge/charge cycles as the 48 tremendous volume changes <sup>[18-19]</sup>. A number of research methods about transition metal 49

50 oxides as anodes in lithium ion batteries have been carried out to improve the deficiency and obtained excellent electrochemical performance by constructing nanostructured materials, 51 hollow nanostructures, hybrid nanostructures, etc. Carbon coating is the most widely used 52 modification techniques for transition metal oxides. On the one hand, carbon coating on the 53 surfaces of metal oxide nanoparticles can reduce the side reactions of solid electrolyte 54 interface (SEI) at the interface between metal oxide and electrolyte <sup>[19]</sup>. On the other hand, 55 good electrical conductivity of carbon can make up for the poor conductivity of metal oxides 56 and promote the electron transport. What's more, carbon coating as elastic buffer 57 layers/supports <sup>[20]</sup> can confine the position of metal oxides and prevent the agglomeration 58 and cracking of crystal structures which can enhance the cycle stability of electrode. In 59 addition, nanostructured electrode materials have some special characteristics, such as large 60 61 proportion of surface atoms, small size, etc. which lead to higher electrode/electrolyte contact area, shorter path lengths for Li<sup>+</sup> transport and higher charge/discharge rates<sup>[21]</sup>. So carbon 62 coating nanostructured metal oxides as anodes in lithium ion batteries would strengthen the 63 cycle stability and improve the high-rate charge-discharge performance. Iron oxides have the 64 characteristics of low price, environment-friendliness and abundant resources, which make 65 them charming anodes for high performance lithium-ion batteries <sup>[20]</sup>. 66

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A lot of Fe<sub>3</sub>O<sub>4</sub>/carbon nanocomposites  $^{[22-25]}$  and Fe<sub>2</sub>O<sub>3</sub>/carbon nanocomposites  $^{[26-28]}$ 67 have been fabricated and detected as electrode materials for lithium ion batteries. Carbon 68 coating iron oxides nanoparticles can form 0 D nanospheres <sup>[29-30]</sup>, 1 D nanowires <sup>[31-32]</sup>, 2 D 69 nanoflakes <sup>[33-34]</sup> and 3 D structures of porous carbon foam loading iron oxide <sup>[35-36]</sup>. 70 Electrospinning is one of the carbon coating methods and has been used to fabricate1 D 71 hybrid carbon coating iron oxides nanofibers composites <sup>[37-41]</sup>, 1 D iron oxides nanofibers 72 <sup>[42-43]</sup> and 1 D carbon nanofibers for Li ion batteries <sup>[44-45]</sup>. The method can embed the iron 73 oxides into a conductive carbon by subsequent heat treatment and enhance the 74

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electrochemical properties effectively. Zhang, X. [37] fabricated Fe<sub>2</sub>O<sub>3</sub>-carbon composite 75 nanofibers as durable anode materials for lithium ion batteries. The cell exhibits a reversible 76 capacity of 820 mA h g<sup>-1</sup> at a current rate of 0.2 C even after 100 cycles. Chaudhari, S.<sup>[42]</sup> 77 synthesized hollow-structured  $\alpha$ -Fe<sub>2</sub>O<sub>3</sub> nanofibers by a simple electrospinning technique and 78 subsequent calcination at 500 °C for 4 h in air. The hollow fibers anodes obtained a high 79 reversible capacity of 1293 mA h g<sup>-1</sup> at a current density of 60 mA g<sup>-1</sup> (0.06 C). Kim, C. <sup>[44]</sup> 80 fabricated high-purity carbon nanofiber webs by combination of electrospinning 10 wt. % 81 polyacrylonitrile (PAN) polymer - dimethyl formamide (DMF) solution and thermal 82 treatments. Nanofiber webs thermally treated at 1000 °C in Ar have the highest reversible 83 capacity of 450 mA h g<sup>-1</sup> than that of 700 °C or 2800 °C at a discharge current density of 30 84 85  $mAg^{-1}$ .

Here, 1D Fe<sub>3</sub>O<sub>4</sub> nanoparticles/porous carbon nanofibers (Fe<sub>3</sub>O<sub>4</sub>/PCNFs) composites are 86 fabricated by electrospinning a 10 wt. % PAN and 3 wt. % Fe(acac)<sub>3</sub> dissolving in DMF 87 solution and subsequent thermal treatments (Fig. 1). The innovative thermal treatments are 88 89 made up of four calcination processes for the first time. Firstly, the electrospun nanofibers were pre-oxidized at 250 °C in air to maintain their morphologies. Secondly, carbonization 90 was proceeded at 1000 °C in argon to obtain high conductive graphitized carbon. Thirdly, 91 oxidation was continued at 400 °C in blowed air to change Fe atoms and Fe C compounds 92 into iron oxides and remove some graphitized carbon of the nanofibers to form pore 93 structures. Carbothermic reduction was finally proceeded at 500 °C in Ar to change Fe<sub>2</sub>O<sub>3</sub> 94 nanoparticles into high conductive Fe<sub>3</sub>O<sub>4</sub> nanoparticles with better crystal structures and 95 further more increase porosity of the nanofibers. What should be noted is that the flow rate of 96 air and calcinating equipment have very significant effects on content of carbon and types of 97 iron oxides in the third calcination process. The excellent electrochemical properties of the 98 novel high conductive Fe<sub>3</sub>O<sub>4</sub>/PCNFs composites anode is evident from the high capacity of 99

100 717.2 mA h  $g^{-1}$  after 100 cycles at 0.5 A  $g^{-1}$ .

# 101 2 EXPERIMENTAL SECTION

## 102 **2.1 Fabrication of Fe<sub>3</sub>O<sub>4</sub>/PCNFs Composites**

PAN with 150,000 average molecular weight was purchased from Aladdin. Analytic
grade ferric acetylacetonate (Fe(acac)<sub>3</sub>) and DMF were obtained from Sinopharm Chemical
Reagent Co.,Ltd. All the reagents were not further purified before using.

In a typical process, 1 g PAN and 0.3 g Fe(acac)<sub>3</sub> were added into 8.70 g DMF and 106 formed a 10 g mixture, followed by vigorous stirring at room temperature for at least 24 h. 107 The obtained homogeneous dispersion was used as the Fe<sub>3</sub>O<sub>4</sub>/PCNFs composites precursor 108 109 solution for electrospinning. The precursor solution was loaded into a 10 mL plastic syringe connected to a blunt-tip needle with a inner diameter of 0.33 mm. The distance between the 110 needle tip and collector is 15 cm. The injection flow rate of the solution was set to 0.1 mm 111 min<sup>-1</sup> (about 1 mL h<sup>-1</sup>) driven by a pumb. A total voltage power of 15 kV was applied 112 between the needle (12 kV) and a plate (-3 kV) covered with aluminum foil. Electrospinning 113 114 was proceeded with a commercial setup (Beijing Ucalery Technology Development Co., Ltd, 115 SS-2535DC). The environmental humidity of electrospinning was  $45\pm10$  %, and temperature 25±5 °C. The electrospun nanofibers were firstly preoxidized at 250 °C for 120 min with a 116 heating rate of 1 °C min<sup>-1</sup> in air atmosphere in a tube furnace (Hefei Ke Jing Materials 117 Technology Co., Ltd., OTF-1200x $\Phi$ 50) to keep fibrous morphology, and then carbonized at 118 1000 °C for 120 min with a heating rate of 5 °C min<sup>-1</sup> in argon atmosphere to obtain high 119 conductive graphitized carbon. Then the composite nanofibers were calcined at 400  $\,^\circ C$  for 3 120 h with a heating rate of 5  $^{\circ}$ C min<sup>-1</sup> in air atmosphere to obtain the Fe<sub>2</sub>O<sub>3</sub> and Fe<sub>3</sub>O<sub>4</sub> 121 122 nanoparticle-loaded porous carbon nanofibers (Fe<sub>x</sub>O<sub>y</sub>/PCNFs) composites. It should be noted that the flow of air and the calcination time both have a profound influence on the 123 124 compositions and morphology of the Fe<sub>x</sub>O<sub>y</sub>/PCNFs composites because it is likely that

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graphitized carbon is completely oxidized to  $CO_2$  and  $Fe_xO_y$  transforms into pure  $Fe_2O_3$  by much longer time calcination and bigger air flow rate at 400 °C in air. At last,  $Fe_3O_4$ /PCNFs composites were prepared by further calcination at 500 °C for 2 h with a heating rate of 5 °C min<sup>-1</sup> in argon atmosphere.

For comparison, homogeneous dispersion containing 1.00 g PAN and 9.00 g DMF was also prepared, and then proceeded electrospinning and calcination with the same processes. In addition, the preoxidized nanofibers were calcined at 700 °C or 900 °C in Ar to observe electrical conductivity of calcinated nanofibers at different calcination temperature. For ease of description, the above products were renamed in this article as follows.



temperature.

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136 137  $Fe(acac)_3$ PAN DMF Temp (atm) Temp (atm) Temp (atm) Temp (atm) 138 Samples First Second Third Last g g g 139 140 PAN0-250 0.00 1.00 9.00 250 °C (air) PAN0-700 0.00 1.00 9.00 250 °C (air) 700 °C (Ar) 141 142 PAN0-900 0.00 1.00 9.00 250 °C (air) 900 °C (Ar) 250 °C (air) 1000 °C (Ar) 143 PAN0-1000 0.00 1.00 9.00 144 PAN0-400 0.00 1.00 9.00 250 °C (air) 1000 °C (Ar) 400 °C (air) PAN0-500 250 °C (air) 1000 °C (Ar) 400 °C (air) 500 °C (Ar) 145 0.00 1.00 9.00 146 PAN3 0.30 1.00 8.70 \_\_\_\_ 250 °C (air) 147 PAN3-250 0.30 8.70 1.00 8.70 250 °C (air) 1000 °C (Ar) 148 PAN3-1000 0.30 1.00 149 PAN3-400 0.30 250 °C (air) 1000 °C (Ar) 400 °C (air) 1.00 8.70 500 °C (Ar) PAN3-500 0.30 250 °C (air) 1000 °C (Ar) 400 °C (air) 150 1.00 8.70

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Comment: Temp = Temprature, atm = atmosphere.

# 152 **2.2 Materials characterizations**

153 The crystal structures of as-prepared materials were characterized using powder X-ray 154 diffraction(XRD) on a D8 Advanced with lynxEye and SolX (Bruker AXS, WI, USA) with a 155 Cu-K $\alpha$  radiation source between 10 ° and 80 °. The morphologies were characterized using a 156 field-emission scanning electron microscopy (SEM) (S-4800 II, Hitachi, Japan) and a field

157 emission transmission electron microscopy (TEM) (Tecnai 12, Philips, Holland) in the Test Center of Yangzhou University. A SEM (Zeiss Supra<sup>™</sup> 55, Sapphire Carl Zeiss Group, 158 Germany) and a TEM (JEM-2100, JEOL, Japan) in College of Chemistry and Chemical 159 160 Engineering of Yangzhou University were also used to characterized the morphologies. High-resolution TEM (HRTEM) and high-angle annular dark-field scanning transmission 161 electron microscopy (HAADF-STEM) were conducted using a FEI Tecnai G2 F30 STWIN 162 (USA) operating at 200 kV. Thermogravimetric analysis (TGA) (Pyris 1 TGA, PerkinElmer, 163 USA) was performed in air atmosphere up to 800 °C at a heating rate of 10 °C min<sup>-1</sup>. Raman 164 165 spectra was conducted with a laser Raman spectrometer (In Via, Renishaw, UK) at the 532 nm wavelength. The magnetic measurements of PAN3-400 and PAN3-500 were performed 166 on a vibrating sample magnetometer (VSM) (EV7, ADE, USA). The electrical conductivities 167 168 of different pure carbon nanofibers were measured by a direct volt-ampere method (SZT-2A, Suzhou Tong-Chang electronics company Ltd, CHN). Nitrogen physisorption-desorption 169 measurements at 100 °C were performed by a surface area and porosity analyzer (ASAP 2020 170 171 HD88, Micromeritics, USA). Brunauer-Emmett-Teller (BET) analyses were done by software to characterize surface properties of porous carbon nanofibers. X-Ray photoelectron 172 spectroscopy (XPS) measurements were conducted with an Al Ka (1486.8 eV) X-ray source 173 (ESCALAB 250Xi, Thermo Fisher Scientific, USA). 174

**2.3 Electrochemical measurements** 

The calcined flexible PAN0-500, PAN3-400 and PAN3-500 mats were cut into 176 electrodes with size  $\Phi 16$  mm, which were assembled into lithium ion batteries by attaching 177 onto a current collector copper foil with 10 wt. % PVDF which dissolved in 178 1-methyl-2-pyrrolidone (NMP) as binder. The electrodes were first dried in a 179 vacuum drying oven at 80 °C for 12 h. The PAN0-500, PAN3-400 or PAN3-500 electrode 180 was about 2 mg. Then coin cells were assembled with metallic lithium as the 181

182 counter/reference electrode, 1 M LiPF<sub>6</sub> in ethylene carbonate (EC), diethyl carbonate (DMC) and ethylmethyl carbonate (EMC) (1:1:1 by volume) as electrolyte, and Celgard 2400 183 polypropylene as separator in an high-purity argon-filled glovebox (VAC-Omni, OMNI-LAB, 184 185 Vacuum atmospheres company, CA). Cyclic voltammetry (CV) measurements were performed using an electrochemical workstation (CHI660 E, Chenghua, CHN) at a scan rate 186 of 0.0001 V s<sup>-1</sup> between 0.01 and 3.0 V. Galvanostatic charge (lithium extraction) and 187 discharge (lithium insertion) cycling of the cells were carried out using a battery test system 188 (CT-3008W, Xinwei, CHN) at the different current densities of 0.05, 0.1, 0.2, 0.5 and 1.0 A 189  $g^{-1}$  between 0.01 and 3 V (vs. Li<sup>+</sup>/Li) to observe ratio performance, and at the current density 190 of 0.5 A  $g^{-1}$  to study cycle stability. Electrochemical impedance spectroscopic 191 192 (EIS) measurements were carried out on an electrochemical analyzer (Autolab, Ecochemie, 193 NL). All the current densities and capacities in this study were calculated on the basis of the 194 weight of integral maps.

Finally, the morphology of PAN3-500 anode was observed by S-4800 II SEM and Tecnai 12 field emission TEM after 100 charge/discharge cycles at  $0.5 \text{A g}^{-1}$ .

# 197 **3 RESULTS AND DISCUSSION**

### 198 **3.1 Preparation process**

The overall synthesis procedures of Fe<sub>3</sub>O<sub>4</sub>/PCNFs composites are schematically 199 200 illustrated in Fig. 1. PAN and Fe(acac)<sub>3</sub> were dissolved in DMF forming uniform viscous 201 reddish-brown electrospinning solution by stirring 24 h. And electrospinning was carried out using above conditions in experimental section. The original electrospinning nanofibers 202 (PAN3) films are yellow. Wide brown slices (PAN3-250) are obtained by pre-oxidation of 203 204 PAN3 films at 250 °C in air. Black graphitized nanofibers (PAN3-1000) composites are prepared by calcination of PAN3-250 films at 1000 °C in Ar. Black porous graphitized carbon 205 coating Fe<sub>x</sub>O<sub>y</sub> nanofibers (PAN3-400) sheets are gotten by calcinating PAN3-1000 at 400 °C 206

207 in air. At last, black Fe<sub>3</sub>O<sub>4</sub>/PCNFs (PAN3-500) compounds are gained by calcinating PAN3-400 in Ar at 500 °C. The pieces have continuous macroscopic area shrinkages by 208 pre-oxidation and graphitization, which can be demonstrated from the microscopic 209 characterizations of SEM and TEM images below. PAN3-400 flakes have obvious weight 210 loss and become more fluffy than PAN3-1000 pieces. A portion of carbon should be oxidized 211 to CO<sub>2</sub> at 400 °C and form a large amount of pores in air atmosphere, which can be 212 demonstrated by the BET results of PAN0-1000 and PAN0-400 as shown in Fig. 2. 213



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Fig. 1 Schematic illustration of the process of electrospinning and products of different

calcination conditions.

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### **3.2 Materials characterization** 217

It is well known that high temperature is advantageous to the graphitization of 218 nanofibers. The pure PAN0-250 was calcinated at the temperature of 700, 900 and 1000 °C 219 under Ar. Table 2 shows the conductivities of carbon nanofibers with the various calcination 220 temperatures and BET results of PAN0-1000 and PAN0-400. PAN0-1000 has the highest 221 conductivity with the value of 476.2  $\mu$ S cm<sup>-1</sup>, which indicates that the graphitization degree 222 increases with the carbonization temperature from 700 to 1000 °C. However, the GCNFs 223 have small specific surface area. In order to increase the specific surface area of graphitized 224 carbon nanofibers, the carbon nanofibers are continually calcinated at 400 °C under air 225 atmosphere. Fig. 2(A) shows the nitrogen adsorption-desorption isotherms of PAN0-1000 226

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227 and PAN0-400. The BET specific surface areas of PAN0-1000 and PAN0-400 are calculated to be 40.6 and 542.6 m<sup>2</sup> g<sup>-1</sup>, respectively. After further calcination at 400 °C in air, the 228 specific surface area of PAN0-400 increased 12.4 times than that of PAN0-1000. Fig. 2(B) 229 displays the distributions of the pore size of PAN0-1000 and PAN0-400. Most pore diameter 230 of PAN0-1000 ranges from 15 to 50 nm and a main peak centers at ~22 nm. The pore 231 diameter of PAN0-400 has relatively large variation range of 3.5 - 50 nm. Most pore 232 diameter of PAN0-400 ranges from 3.5 to 23 nm and a main peak centers at ~6.4 nm. As 233 shown in Table 2, it is worth noting that the conductivity of PAN0-400 is also much higher 234 than PAN0-250 calcinating at 700 °C (PAN0-700). The method will be applied in the 235 preparation of the follow-up composite of nanofibers in this paper. 236



Fig. 2 Nitrogen adsorption-desorption isotherms of PAN0-1000 and PAN0-400 (A) and and pore size distributions of PAN0-1000 and PAN0-400 (B).

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The morphology and the diameter of the electrospun nanofibers of PAN0 and PAN3 are compared. It is found that there is no distinct difference between pure PAN nanofibers and

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247 composite nanofibers as shown in Fig. S1, indicating that the electrospinning solution together with operation conditions of electrospinning in this experiment are appropriate to 248 obtain uniform and perfect nanofibers. Fig. 3 shows SEM images of PAN3-250, PAN3-1000, 249 PAN3-400 and PAN3-500. All nanofibers exhibit homogeneously distributed diameters. Fig. 250 S2 shows the diameter distributions of PAN3-400 and PAN3-500. The mean sizes of 251 PAN3-400 and PAN3-500 are 250 and 160 nm, respectively. More interesting, when the 252 carbon nanofibers was successively calcinated at 500 °C under Ar atmosphere, the average 253 diameter of PAN3-500 decreased obviously comparing with that of PAN3-400, which was 254 255 due to the consumption of carbon in ferric iron reduction reaction. In the control experiment, the average diameter of PAN0-500 has no distinct decrease relative to that of PAN0-400 as 256 shown in Fig. S3. 257



**Fig. 3** SEM images of PAN3-250 (a), PAN3-1000 (b), PAN3-400 (c) and PAN3-500 (d).

As shown in Fig. 4(a) and (c), PAN3-400 and PAN3-500 have homogeneous

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261 morphologies. But the porosities are distinctly increasing along with continuous calcination. As can be seen from Fig. 4(b) and (d), there are many tiny pores and particles in PAN3-400 262 and PAN3-500. The portion of darker colour of PAN3-400 and PAN3-500 would be iron 263 oxides nanoparticles. The lighter colour would be carbon and tiny pores of them. The little 264 white spots should be pores. The above-mentioned iron oxides nanoparticles, carbon and 265 pores can be certified by following XRD patterns in Fig. 8(A), Raman spectra in Fig. 8(E) 266 267 and BET values in Table.2, respectively. What should be noted is that in the third calcination process, both the flow of air and the placed apparatuses of PAN3-1000 have great influence 268 269 to integral morphology of nanofiber and respective content of carbon and iron oxides. Fig. S4 shows two differrent equipments in the third calcination process. Fig. 5 shows the SEM and 270 TEM images of product calcinated using apparatus in Fig. S4(b) in the third calcinating 271 272 process. Fig. 5(a) and Fig. 5(b) have obvious different morphologies comparing with Fig. 3(c) and Fig. 4(b), respectively. The degree contacting with air of PAN3-1000 in the third 273 calcination process is responsible for the results. What's more, large percentage shrinkage in 274 area would occur or red brown product would generate at much higher temperature or larger 275 flow rate of air. In this paper, the relevant experiments of a series of air flow and different 276 equipments in the third calcinating process will not be further discussed. And all of the other 277 products (PAN3-400 and PAN0-400) were calcinated using the apparatus in Fig. S4(a). 278



Fig. 4 Typical TEM image of PAN3-400 (a) and its partial magnification (b), representative

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TEM image of PAN3-500 (c) and its partial magnification (d).





Fig. 6 shows HRTEM images of PAN3-500. The clear shell lattice fringes with

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d-spacing of 0.30 nm in HRTEM picture (Fig. 6(c)) shows are in good agreement with that of 286 (220) plane of cubic Fe<sub>3</sub>O<sub>4</sub>. Fig. 6(d) demonstrates the clear shell lattice fringes with average 287 d-spacing of 0.34 nm, corresponding to the (002) plane of hexagonal graphite. But the 288 lattice structures are seldom observed. As shown in table 2, the conductivity of PAN0-400 is 289 reduced significantly comparing with PAN0-1000. The lattice structures of graphitized 290 291 carbon nanofibers would have been destroyed to a great extent in the calcination processes at 400 °C in air for 3 h and carbon thermal reduction at 500 °C for 2 h. 292







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295	PAN3-500 (b), the boxed region of part (b) - the observed crystallites in the HRTEM image
296	corresponding to $Fe_3O_4$ (c), and seldom observed HRTEM image of lattice structure porous
297	carbon of PAN3-500 (d).
298	The PAN3-500 composites are expressed by the element maps from EDX spectroscopy.
299	The distributions of C, O and Fe are shown in Fig. 7(c), (d) and (e), respectively. The Fe and
300	O elements prove the existence of iron oxide. All elements distribute uniformly in the
301	nanofiber. Therefore, the element maps of EDX confirm the composition of PAN3-500
302	composites.



Fig. 7 HAADF-STEM images of PAN3-500 (a) and partial enlargement (b), element 304 305 mapping of C (c), O (d) and Fe (e).

Fig. 8(A) shows the XRD patterns of PAN0-1000, PAN3-400 and PAN3-500. The 306 307 identified diffraction peaks of PAN0-1000 nanofibers can be clearly assigned to the hexagonal graphite (JCPDS, card 41-1487). The peak at  $2\theta = 26.38$  can be indexed to (002) 308 lattice plane of hexagonal graphite, which manifests the precursor-PAN has changed into 309 high conductive graphitic carbon. The peaks at  $2\theta = 24.14$ , 33.15, 35.61, 40.85, 49.48, 310

311 54.09, 62.45 and 63.99 can be indexed to (012), (104), (110), (113), (024), (116), (214) and (300) lattice planes of hexagonal Fe<sub>2</sub>O<sub>3</sub> (JCPDS, card 33-0664), respectively. The peaks at 312  $2\theta = 18.27, 30.10, 35.42, 43.05, 56.94$  and 62.52 can be indexed to (220), (311), (400), 313 (511) and (440) lattice planes of cubic Fe<sub>3</sub>O<sub>4</sub> (JCPDS, card 19-0629), respectively. In order 314 315 to obtain the pure Fe<sub>3</sub>O<sub>4</sub> in the nanofibers, PAN3-400 composites were calcinated at 500 °C in Ar. Under inert atmosphere, carbon is used as reductive agent and Fe<sub>2</sub>O<sub>3</sub> transforms into 316  $Fe_3O_4$ . No impurity peaks from other iron oxide are observed, implying the high purity of 317 Fe<sub>3</sub>O<sub>4</sub> in the nanofibers. The intense diffraction peaks of Fe<sub>3</sub>O<sub>4</sub> in PAN3-500 demonstrate 318 319 the better crystal forms of Fe<sub>3</sub>O<sub>4</sub> than that in PAN3-400. But the strong diffraction peak of graphitized carbon is disappeared in XRD patterns of PAN3-400 and PAN3-500, which 320 indicates the structures of graphitized carbon have been destroyed to a great degree. The 321 322 hysteresis curves of as-prepared PAN3-400 and PAN3-500 are shown in Fig. 8(B). The saturated magnetizations of PAN3-400 and PAN3-500 are about 15.0 and 29.2 emu g<sup>-1</sup>, 323 respectively. The results of hysteresis curves are consisted with their compositions shown 324 325 by XRD results. The surface electronic state and composition of PAN3-500 were further investigated by XPS analysis as presented in Fig. 8(C) and (D). Fig. 8(C) reveals the 326 existence of C, N, O and Fe elements in the PAN3-500. The existence of N of PAN3-500 in 327 Fig. 8(C) should be attributable to nitrogen-containing groups of PCNFs originated PAN of 328 the raw materials. Fig. 8(D) shows the high resolution of Fe 2p spectrum. The two main 329 peaks located at 711.0 and 725.7 eV correspond to Fe 2p3/2 and Fe 2p1/2, respectively. The 330 results shown the  $Fe^{3+}$  and  $Fe^{2+}$  states were coexisted in PAN3-500, which 331 further confirmed the iron oxide in PAN3-500 is Fe<sub>3</sub>O<sub>4</sub>. 332

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PAN3-400 and PAN3-500 (B), XPS survey spectrum of PAN3-500 (C) and high resolution of Fe 2p spectrum (D), Raman spectra of PAN0-1000, PAN0-500 and PAN3-500 (E), and TG-DSC curves of PAN0-500 and PAN3-500 (F).

Fig. 8(E) shows the Raman spectra of PAN0-1000, PAN0-500 and PAN3-500. All samples show strong D (disorder) peak at about 1360 cm<sup>-1</sup> and G (graphite) peak at about 1590 cm<sup>-1</sup>. The intensity ratios R ( $I_D/I_G$ ) of PAN0-1000, PAN0-500 and PAN3-500 are 0.96,

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0.99 and 0.99, respectively. The large values of R indicate the degrees of disordered and 341 defected carbon. From Fig. 8(F), the differential scanning calorimetry (DSC) and 342 thermogravimetry (TG) curves of PAN0-500 indicate that the reaction is exothermic and the 343 loss weight is mainly between 550 °C and 600 °C. But the TG curve of PAN3-500 reveals that 344 the weight loss is mainly at the temperature range of 400 °C - 490 °C. It can be speculated 345 that the graphitized carbon structures of PAN3-500 are destroyed more seriously 346 than PAN0-500 due to the reaction of Fe<sub>2</sub>O<sub>3</sub> and carbon etc. and the exothermic reaction of 347 Fe<sub>3</sub>O<sub>4</sub> oxidizing into Fe<sub>2</sub>O<sub>3</sub> promotes the oxidation reaction of porous carbon at the lower 348 temperature. The residual proportion of PAN3-500 is 36.9 wt. %. According to the results of 349 TGA tests, the original percentages in weight of carbon and Fe<sub>3</sub>O<sub>4</sub> in PAN3-500 are 350 calculated to be 66.0 wt. % and 34.0 wt. %, respectively. 351

# 352 Electrochemical Performance

Porous carbon in the final mat PAN3-500 is sufficient to form a stable framework. So 353 the PAN3-500 mat has very good flexibility, as shown in Fig. S5. The CV measurements 354 were carried out to investigate the electrochemical reactions of PAN3-500 in the range of 3.0 355 to 0.01 V at a scan rate of 0.1 mV s<sup>-1</sup> at room temperature. It is seen in Fig. 9(A) the 356 PAN0-500 exhibits a cathodic peak at about 0.63 V which represents the formation of SEI 357 film and the decomposition of the electrolyte <sup>[46]</sup> and two broad anodic peaks at about 1.16 V 358 and 2.11 V attributing to irreversible reactions with electrolyte. In the subsequent cycles, the 359 peak P2 at about 0.18 V and the peak P3 at about 0.25 V should be the insertion and 360 extraction of lithium ion from the graphitized structures of porous carbon nanofibers, 361 respectively. And the peak P1 at about 0.63 V disappears, indicating the stable and complete 362

363	SEI film has formed on the surface of carbon. As shown in Fig. 9(B), at the first cycle,
364	PAN3-500 electrode exhibits a clear cathodic peak. The peak P1 at about 0.54 V might be the
365	initial lithium insertion into the $Fe_3O_4$ to form $Li_2Fe_3O_4$ , given by Eqn (1) <sup>[33,47]</sup> and the
366	reduction reaction of $Li_2Fe_3O_4$ to $Fe^0$ , along with the formation of amorphous $Li_2O$ , given by
367	Eqn (2), as well as the formation of SEI film <sup>[33,41,47]</sup> and the decomposition of the electrolyte.
368	The peak P2 should be the extraction of lithium ion from the graphitized structures of porous
369	carbon nanofibers and the peak at about 1.20 V should be irreversible reactions with
370	electrolyte like as the peak at about 1.16 V of PAN0-500 in Fig. 9(A). The continuous broad
371	peaks P3 and P4 at about 1.64 V and 1.82 V are the oxidation reactions of $Fe^0$ to $Fe^{2+}$ and
372	$Fe^{2+}$ to $Fe^{3+}$ , respectively <sup>[33,36,47]</sup> . The two electrochemical reactions of $Fe^{0}$ to $Fe^{2+}$ and $Fe^{2+}$ to
373	Fe <sup>3+</sup> owning nearby peak potentials and the irreversible reactions could be responsible for the
374	phenomenon that the peaks P3 and P4 merged into a more broaden peak in the subsequent
375	cycles. The total reaction is given by Eqn (3) $^{[33,41,47]}$ .
376	$\operatorname{Fer}\Omega_{4} + 2\operatorname{Li}^{+} + 2\operatorname{e} \longrightarrow \operatorname{Lir}\operatorname{Fer}\Omega_{4}(1)$

377 
$$\text{Li}_2\text{Fe}_3\text{O}_4 + 6\text{Li}^+ + 6\text{e} \rightarrow 3\text{Fe}^0 + 4\text{Li}_2\text{O}(2)$$

378 3 
$$\operatorname{Fe}^{0} + 4 \operatorname{Li}_{2}O \rightarrow \operatorname{Fe}_{3}O_{4} + 8 \operatorname{Li}^{+} + 8 \operatorname{e}(3)$$

EIS measurements were carried out at open circuit potential with an AC voltage 379 amplitude of 5.0 mV in a frequency range from 100 kHz to 0.01 Hz to understand the 380 electrochemical performance of PAN3-500 comparison with PAN3-400, PAN0-500 and 381 PAN0-1000. Fig. 9(C) and (D) display the Nyquist plots of PAN3-500 and PAN3-400 382 electrodes, PAN0-1000 and PAN0-500 electrodes after 1 cycle at 0.5 A g<sup>-1</sup>. From Fig. 9(C), 383 384 the Nyquist plots of PAN3-500 and PAN3-400 have similar profiles, which are consisted of two semicircles, as well as a line, respectively. The two semicircles and the line from high 385 frequency to low frequency were related to the resistance of SEI, charge-transfer resistance 386

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387 on the electrolyte/electrode interface and the solid-state diffusion resistance of Li ion in the electrode, respectively<sup>[13,15]</sup>. From Fig. 9(D), it clearly shows that the radius of the semicircle 388 for PAN0-500 in the medium frequency region is much smaller than that of PAN0-1000, 389 390 indicating the porous structured PAN0-500 has much lower electron-transfer resistance than graphitized PAN0-1000. The PAN3-500, PAN3-400 and PAN0-500 electrodes have 391 remarkably small electron-transfer resistance compared with that of PN0-1000. The results 392 indicates the greatly diminutive charge-transfer resistance at the electrode/electrolyte 393 interface should be due to the porous structure in PCNFs of PAN3-500, PAN3-400 and 394 395 PAN0-500, which can greatly improve the diffusion of lithium ions as well as the transfer of electrons for better elctrochemical performance<sup>[12-16]</sup>. 396



Fig. 9 CV measurements of PAN0-500 (A), PAN3-500 (B) cycled between 0.01 and 3 V at a scan rate of 0.1 mV s<sup>-1</sup>, Nyquist plots of PAN3-500 and PAN3-400 (C), PAN0-500 and

PAN0-1000 (D).

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401 capacitance of the composites is calculated as approximate 560 mA h g<sup>-1</sup> based on the 402 theoretical capacities of Fe<sub>3</sub>O<sub>4</sub> 926 mA h  $g^{-1}$  and graphite 372 mA h  $g^{-1}$ . Fig. 10(a) shows the 403 charge-discharge cycle performance of PAN0-1000, PAN0-500, PAN3-400 and PAN3-500 at 404 0.5 A g<sup>-1</sup> in the range of 3.0 to 0.01 V. In the 100th cycle, the discharge capacity of 405 PAN0-1000, PAN0-500, PAN3-400 and PAN3-500 were 173.1, 261.1, 579.3 and 717.2 mA 406 h  $g^{-1}$ , respectively. The test results show that the capacities of composite nanofibers are 407 obviously higher than that of carbon nanofibers. The high capacity of PAN3-500 is attributed 408 409 to superior porosity of porous carbon and carbon coating good crystallinity of Fe<sub>3</sub>O<sub>4</sub>. The discharge capacities of PAN3-500 in the 1st, 2nd, 10th, 30th, 50th, 70th and 90th cycles were 410 1422.1, 811.7, 672.9, 665.0, 674.6, 692.7 and 700.6 mA h g<sup>-1</sup>, respectively, indicating that 411 PAN3-500 had high capacity and remarkable capacity retention. The SEM images of 412 PAN3-500 after 100 charge-discharge cycles and one discharge cycle are shown in Fig. 11. 413 And the diameter distribution of PAN3-500 after 100 charge-discharge cycles and one 414 discharge cycle is shown in Fig. S6. The mean diameter of PAN3-500 after 100 415 charge-discharge cycles and one discharge cycle is 180 nm. The results shows that the mean 416 diameter of PAN3-500 after 100 charge-discharge cycles and one discharge cycle is the same 417 as the original PAN3-500, which can demonstrate that PAN3-500 basically has no volume 418 change in the charging-discharging process at the density of 0.5 A g<sup>-1</sup>. The amounts of small 419 pores which can afford space for inserted lithium should be responsible for this phenomenon. 420 Fig. 10(b) shows the detailed charge and discharge curves of PAN3-500 at different cycle 421 numbers. As shown in Fig. 10(c), the rate capabilities of PAN3-500 were investigated at 422 various rates from 0.05 to 1.0 A  $g^{-1}$  to further evaluate the electrochemical performance. 423 When the densities were 0.05, 0.1, 0.2, 0.5 and 1.0 A  $g^{-1}$ , the PAN3-500 retained high 424 specific capacities of 1050.6, 931.1, 857.0, 709.0 and 501.6 mA h  $g^{-1}$  in their own 5th cycle, 425

426 respectively. And then back to 0.1 A  $g^{-1}$ , the capacity still achieved 919.3 mA h  $g^{-1}$ . Fig. 10(d) 427 shows the charge/discharge curves of PAN3-500 at different current densities, which display 428 approximate symmetrical shape, indicating fine reversibility of Li<sup>+</sup> insertion/extraction.





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Fig. 11 SEM image of PAN3-500 after 100 charge-discharge cycles and one discharge cycle
(a) and relevant magnifation (b).

437 It is worth noting that the electrochemical performance of PAN3-500 in this work are 438 comparable to most of the electrospinning and calcined products, such as  $Fe_2O_3$  nanofibers 439 <sup>[42-43]</sup>, carbon nanofibers <sup>[44-40]</sup> and iron oxides/CNFs composites <sup>[37-41]</sup> reported in previous 440 literatures, as shown in Table 3.

441

Table 3 Comparison of the retaining capacity of  $Fe_xO_y$  nanofibers, carbon nanofibers or

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$Fe_xO_y/CNFs$ composites	s preparing	by el	lectrospinnin	g.
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443	Nanocomposites	Precursors	Calcination temperature	Retaining capacity
444	Fe <sub>2</sub> O <sub>3</sub> -carbon	PAN-Fe(acac) <sub>3</sub>	At 500 °C for 3h in air	820 mA h g <sup>-1</sup> at 0.2 C [37]
445	a-Fe <sub>2</sub> O <sub>3</sub> -CNFs	PAN-FeCl <sub>3</sub>	At 600 $^{\rm o}{\rm C}$ for 8 h in Ar	$600 \text{ mA h g}^{-1} \text{ at } 50 \text{ mA g}^{-1} [38]$
446	$\alpha$ -Fe <sub>2</sub> O <sub>3</sub> nanorods	PVP-Fe(acac) <sub>3</sub>	At 500 $^{\circ}$ C for 5 h in air	1095 mA h g <sup>-1</sup> at 0.05 C [39]
447	C/Fe <sub>3</sub> O <sub>4</sub>	PAN-Fe(acac) <sub>2</sub>	At 600 °C for 10 h in Ar	1096 mA h $g^{-1}$ at 0.2 A $g^{-1}$ [40]
448			At 700 $^{\rm o}{\rm C}$ for 10 h in Ar	$300 \text{ mA h g}^{-1}$ at 0.2 A g $^{-1}$ [40]
449	Fe <sub>3</sub> O <sub>4</sub> @PCFs	PAN-PS-Fe <sub>3</sub> O <sub>4</sub>	At 600 °C for 2 h in Ar	541 mA h $g^{-1}$ at 2.0 A $g^{-1}$ [41]
450	Hollow Fe <sub>2</sub> O <sub>3</sub> nanofibers	B PVP-Fe(acac) <sub>3</sub>	At 500 $^{\circ}$ C for 4 h in air	1293 mA h g <sup>-1</sup> at 0.06 A g <sup>-1</sup> [42]
451	Porous $Fe_2O_3$ nanotubes	PVP-Fe(acac) <sub>3</sub>	At 500 °C for 3 h in air	987.7 mA h $g^{-1}$ at 0.2 A $g^{-1}$ [43]
452	Carbon nanofibers	PAN	At 700 °C in Ar	275 mA h g <sup>-1</sup> at 0.03 A g <sup>-1</sup> [44]
453			At 1000 °C in Ar	450 mA h g <sup>-1</sup> at 0.03 A g <sup>-1</sup> [44]
454			At 2800 °C in Ar	140 mA h g <sup>-1</sup> at 0.03 A g <sup>-1</sup> [44]
455	Carbon nanofiber	PAN	At 800 °C for 1 h in Ar	407 mA h g <sup>-1</sup> at 0.15 A g <sup>-1</sup> [45]
456			At 1300 °C for 1 h in Ar	239 mA h $g^{-1}$ at 0.15 A $g^{-1}$ [45]
457	Fe <sub>3</sub> O <sub>4</sub> /PCNFs	PAN-Fe(acac) <sub>3</sub>	At 1000 °C for 2 h in Ar,	
458			at 400 °C for 3 h in Air,	579.3 mA h $g^{-1}$ at 0.5 A $g^{-1}$ [This]
459			at 500 $^{\rm o}{\rm C}$ for 2 h in Ar	717.2 mA h $g^{-1}$ at 0.5 A $g^{-1}$ [This]

460 Comment: PS = polystyrene, PVP = polyvinylpyrrolidone, PCFs = porous carbon fibers.

461 **CONCLUSIONS** 

In summary, we synthesized  $Fe_3O_4$ /porous carbon nanofibers by electrospinning and subsequent unique calcination processes. Graphitized carbon was obtained by calcination at 1000 °C in Ar. Porous structured carbon was obtained by further calcination at 400 °C in air, which had much higher porosity than graphitized carbon.  $Fe_3O_4$ /porous carbon nanofibers

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were prepared via the carbon-thermal reduction process by the final calcination at 500 °C in 466 Ar. Porous carbon coating nano-sized  $Fe_3O_4$  nanofibers composites had excellent 467 performance as anodes for lithium ion batteries. The reversible capacity was higher than 468 Fe<sub>x</sub>O<sub>y</sub>/porous carbon nanofibers composites and carbon nanofibers. Fe<sub>3</sub>O<sub>4</sub>/porous carbon 469 nanofibers kept a reversible capacity of 717.2 mA h  $g^{-1}$  at 0.5 A  $g^{-1}$  after 100 cycles. At 0.05 A 470  $g^{-1}$ , the composites delivered a reversible capacity as high as 1050.6 mA h  $g^{-1}$ . The porous 471 carbon nanofibers of coating Fe<sub>3</sub>O<sub>4</sub> nanoparticles have good conductivity, which is useful for 472 electronic transmission and fast diffusion of lithium in the charge-discharge process. What's 473 474 more, carbon can buffer the volume changes between nano-sized Fe<sub>3</sub>O<sub>4</sub> nanoparticles and Fe atoms in charging and discharging process and make the transition reactions in situ to 475 maintain the crystal structures. So the combinations of conductive porous carbon nanofibers 476 477 and nano-sized Fe<sub>3</sub>O<sub>4</sub> are responsible for the higher capability and good cycling stability. We believe much higher capacity Fe<sub>3</sub>O<sub>4</sub>/porous carbon nanofibers can be fabricated by more 478 appropriate calcination conditions containing calcinating apparatuse, temperature and flow 479 480 rate of air. More importantly, the method of preparing Fe<sub>3</sub>O<sub>4</sub>/porous carbon nanofibers is novel. 481

# 482 ASSOCIATED CONTENT

### 483 Supporting Information

Additional experimental data including SEM images of PAN0 and PAN3, SEM images of PAN3-400 (a), PAN3-500 (c) and relevant diameter distributions of PAN3-400 (b), PAN3-500 (d), SEM images of PAN0-400, PAN0-500 and relevant diameter distributions, two differrent equipments in the third calcination process, picture of flexible PAN3-500 electrodes, SEM image of PAN3-500 after 100 charge-discharge cycles and one discharge cycle and relevant diameter distribution.

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- 493 **Notes**

494 The authors declare no competing financial interest.

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Ferroferric oxide/porous carbon nanofibers composites were synthesized by electrospinning and

subsequent innovative thermal treatments, exhibiting enhanced electrochemical performance.