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### Introduction

With the rapid development of electronic technology, electromagnetic interference has become the fourth major type of pollution, after water pollution, atmospheric pollution, and noise pollution, and constitutes a serious threat to human health.<sup>1-3</sup> Meanwhile, the leakage of electromagnetic waves also affects the normal operation of electronic equipment, resulting in serious losses.<sup>4</sup> The development of a microwave-absorbing material that is thin while having a low density, as well as having a broad, strong absorption band, would be of great significance to the improvement of the quality of human life and for the industrial production of electromagnetic protection equipment and materials.

In general, magnetic particles such as Fe, $^5$  Co, $^6$  Ni, $^7$  Fe $_3\mathrm{O_4},^8$ and carbonyl iron<sup>9</sup> particles can be used to absorb electromagnetic waves due to their highly complex permeability; however, their high densities and narrow absorption bands limit their applicability.<sup>10,11</sup> Electromagnetic wave loss is such that the overall loss is determined by both the magnetic and electrical losses. Combining a magnetic material with a dielectric can not only reduce the weight but also achieve better microwave absorption through impedance matching. Carbon

# Activated carbon fiber/Fe<sub>3</sub>O<sub>4</sub> composite with enhanced electromagnetic wave absorption properties

Qilong Sun,  $\bullet$ <sup>\*ab</sup> Lei Sun,<sup>ab</sup> YingYing Cai,<sup>ab</sup> Tao Ji<sup>ab</sup> and Guangyu Zhang<sup>ab</sup>

To obtain a low-density material that is capable of absorbing electromagnetic waves over a wide bandwidth, an activated carbon fiber/Fe<sub>3</sub>O<sub>4</sub> composite material (ACF/Fe<sub>3</sub>O<sub>4</sub>) was prepared using an in situ reduction method. Scanning electron microscopy images show that  $Fe<sub>3</sub>O<sub>4</sub>$  nanoparticles, approximately 10–40 nm in size, were spread uniformly over the surface of the ACF. The resulting composite exhibited superparamagnetic behavior at room temperature. The ability of the ACF and ACF/Fe<sub>3</sub>O<sub>4</sub> composite to absorb electromagnetic waves over a frequency range of 8.2–18 GHz was measured using the arch method. The results showed that the maximum reflectivity of an ACF felt was -12.9 dB at 18 GHz, and the effective microwave-absorbing bandwidth ( $R < -10$  dB) was 1.9 GHz (16.10–18 GHz). The absorption performance of the ACF was greatly enhanced by being loaded with  $Fe<sub>3</sub>O<sub>4</sub>$  nanoparticles; the maximum reflectivity of the 2 mm layer of the ACF/Fe<sub>3</sub>O<sub>4</sub> composite was  $-30.07$  dB at 16.45 GHz, and the effective bandwidth ( $R < -10$  dB) increased to 8.62 GHz (9.38-18 GHz). Coating with nano-Fe<sub>3</sub>O<sub>4</sub> magnetic particles can effectively improve the absorption of electromagnetic waves by the ACF, and this technique therefore has great potential for application to the field of electromagnetic shielding. PAPER<br>
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ber is a type of dielectric-loss material with a carbon content of more than 95%. It offers the advantages of low density and a low coefficient of thermal expansion, as well as good absorbency of electromagnetic waves at high frequencies, relative to magnetic metal materials.<sup>12,13</sup> However, not every type of carbon fiber can be used as a microwave-absorbing material; this depends on the conductivity of the carbon fiber. An excessively high electrical conductivity causes the carbon fibers to reflect electromagnetic waves, which strongly affects their ability to absorb microwaves. Therefore, only carbon fibers that have been specially treated can be used to absorb microwaves. Wu et al.<sup>14</sup> suggested that the use of activated carbon fiber (ACF) is an effective approach to increasing electromagnetic wave absorption because of the multiple reflections and longer propagation distance inside the composite.

In the present study, we prepared a lightweight composite material, capable of absorbing electromagnetic waves over a wide bandwidth, by coating ACF with  $Fe<sub>3</sub>O<sub>4</sub>$  magnetic particles. The  $Fe<sub>3</sub>O<sub>4</sub>$  magnetic particles were applied using an in situ reduction method, and the reflectivities of the ACF and ACF/ Fe3O4 felts over a frequency range of 8.2–18 GHz were studied using the arch method.

### **Experimental**

### Materials

Viscose fiber felt (Jiangsu Sutong Carbon Fiber Co., Ltd.), phosphoric acid (Nantong Qixiang Biotechnology Co., Ltd.), anhydrous

a National & Local Joint Engineering Research Center of Technical Fiber Composites for Safety and Protection, Nantong University, Nantong 226019, Jiangsu, P. R. China b College of Textiles and Clothing, Nantong University, Nantong 226019, Jiangsu, P. R. China. E-mail: sunqilong001@ntu.edu.cn

ferric chloride (FeCl<sub>3</sub>; Shanghai Runjie Chemical Reagent Co., Ltd.), and glucose monohydrate  $(C_6H_{12}O_6 \cdot H_2O)$ ; Shanghai Runjie Chemical Reagent Co., Ltd.), all of reagent grade, were purchased and used without any further purification.

#### Preparation of ACF felt

The viscose fiber felt samples were cleaned by ultrasonication in water for 30 min to remove the textile oiling agent. Phosphoric acid was employed as an activator to prepare the viscose-based ACF. The dried viscose fiber felts were soaked in phosphoric acid solution (2.0 mol  $\text{L}^{-1}$ ) for 2 h and then dried at 60 °C. The dried sample was then placed in a vacuum muffle furnace and carbonized under a nitrogen atmosphere. The carbonization temperature was 650 °C; the heating rate was 10 °C  $\mathrm{min}^{-1},$  and the holding time was 1 h. This produced the ACF felt. PSC Advances<br>
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#### Preparation of ACF/Fe<sub>3</sub>O<sub>4</sub> felt

The ACF felts were immersed in saturated  $\text{FeCl}_3$  solution (1000 mL) at 60  $\degree$ C for 2 h, then 15 g of glucose monohydrate was added, followed by ultrasonication for 1 h. The sample was dried at 60 $\degree$ C, after which it was subjected to a further hightemperature treatment in a vacuum muffle furnace; the treatment temperature was 750 °C, the heating rate was 15 °C  $\mathrm{min}^{-1},$ and the holding time was 30 min. This produced  $ACF/Fe<sub>3</sub>O<sub>4</sub>$ felts. The preparation process is illustrated in Fig. 1.

#### Characterization

The surface morphologies of ACF and  $ACF/Fe<sub>3</sub>O<sub>4</sub>$  were characterized using field-emission scanning electron microscopy (FESEM; S-4800, Hitachi High-Technologies) with an acceleration voltage of 3 kV.

The surface chemical compositions of the ACF and ACF/ Fe3O4 were analyzed using X-ray photoelectron spectrometry (XPS; ESCALAB-250Xi, Thermo Fisher Scientific). The X-ray source was Al K $\alpha$ , the vacuum level of the test system was  $10^{-8}$ - $10^{-9}$  Pa, and the angle between the electron beam and sample was 45°.

The magnetic properties of  $ACF/Fe<sub>3</sub>O<sub>4</sub>$  are presented as a hysteresis loop, measured using a superconducting quantum interference device-vibrating sample magnetometer (SQUID-VSM; MPMS3, Quantum Design) at room temperature.

The electromagnetic wave absorption performance was investigated by the arch method. The testing apparatus is schematically illustrated in Fig. 2. The vector network analyzer connected with the receiving and transmitting antennas, one having functions of stimulating and transmitting frequency electromagnetic waves through and the other receiving the reflective signal. According to the feedback signal strength, the size of the samples was  $180 \times 180$  mm with 2 mm thickness.

### Results and discussion

#### Phase composition and morphology

The XRD pattern for the ACF is shown in Fig. 3(a). ACF has a typical amorphous structure,<sup>15</sup> with two strong amorphous diffraction peaks at  $2\theta = 23^\circ$  and  $2\theta = 44^\circ$ , corresponding to the (002) and (101) crystal planes, respectively. Fig. 3(b) shows the XRD pattern for ACF/Fe<sub>3</sub>O<sub>4</sub>. New crystals appear at  $2\theta = 30.42^{\circ}$ , 35.74°, 40.28°, 43.46°, 53.94°, 57.24°, and 62.90°, which



Fig. 1 Preparation of ACF/Fe<sub>3</sub>O<sub>4</sub>.



Fig. 2 Arch-method electromagnetic wave absorption test apparatus.



correspond, respectively, to the (220), (311), (400), (422), (511), and (400) crystal planes of the face-centered cubic  $Fe<sub>3</sub>O<sub>4</sub>$  crystals (JCPDS no. 19-0269).<sup>16</sup> It is noted that glucose plays an important role in  $Fe<sub>3</sub>O<sub>4</sub>$  formation because it can decompose to reductive CO gas, which can fully reduce  $Fe^{3+}$  to  $Fe^{2+}$ . This explains why no  $Fe<sub>2</sub>O<sub>3</sub>$  signal are found in the XRD pattern. The particle size of  $Fe<sub>3</sub>O<sub>4</sub>$  at the (311) crystal surface, calculated using the Scherrer equation, is approximately 16.18 nm, indicating that the  $Fe<sub>3</sub>O<sub>4</sub>$  particles prepared using this method will be nano-sized.

Note that there are some miscellaneous peaks, which may indicate the presence of FeOOH, FeO, Fe<sub>2</sub>C, or Fe $(OH)_2$ . FeOOH and  $Fe(OH)_2$  are likely to be the products of trace amounts of  $Fe<sub>3</sub>O<sub>4</sub>$  undergoing acid–base reactions; FeO and  $Fe<sub>2</sub>C$  may be derived from the carbonization reactions, caused by the reaction of  $Fe<sub>3</sub>O<sub>4</sub>$  with carbon (C) in the raw material or with the cracked gas containing carbon and oxygen.<sup>17</sup>

The microstructures of the ACF and  $ACF/Fe<sub>3</sub>O<sub>4</sub>$  were characterized by FESEM. Fig. 4(a) shows a SEM image of the ACF.

The surface of the ACF is relatively smooth, and there are no particles on the surface that would indicate residual impurities. Fig. 4(b) and (c) shows SEM images of  $ACF/Fe<sub>3</sub>O<sub>4</sub>$  at different magnifications. Most of the  $Fe<sub>3</sub>O<sub>4</sub>$  particles are distributed uniformly over the surface of the ACF. The  $Fe<sub>3</sub>O<sub>4</sub>$  particles are irregular in shape while their sizes are within a range of 15– 60 nm, which corresponds to the particle size as calculated from the XRD pattern via the Scherrer equation. Fig. 4(d) showed that the as-prepared  $ACF/Fe<sub>3</sub>O<sub>4</sub>$  felt was easy to bend, indicative of good flexibility after the treatment.

#### Chemical composition

The elemental compositions and chemical states of the ACF and  $ACF/Fe<sub>3</sub>O<sub>4</sub>$  were characterized by XPS. Fig. 5(a) shows the XPS survey spectra for the ACF and  $ACF/Fe<sub>3</sub>O<sub>4</sub>$ . There are two distinct peaks in these spectra, corresponding to C 1s and O 1s, respectively. An additional distinct peak appears in the XPS spectrum for  $ACF/Fe<sub>3</sub>O<sub>4</sub>$ , which corresponds to Fe 2p. The highresolution Fe 2p spectrum is shown in Fig. 5(b); the two peaks at binding energies of 710.9 eV and 724.5 eV correspond to the Fe  $2p_{3/2}$  and Fe  $2p_{1/2}$  spin–orbit peaks, respectively, indicating the formation of Fe<sub>3</sub>O<sub>4</sub>. Fig. 5(c), (d) shows the O 1s XPS spectra for the ACF and  $ACF/Fe<sub>3</sub>O<sub>4</sub>$  with peak-fitting curves; three peaks are visible for both the ACF and ACF/Fe<sub>3</sub>O<sub>4,</sub> at 531.2 eV, 532.3 eV, and 533.2 eV, corresponding to O–C=O, C=O, and C–O, respectively. Meanwhile, the distinct peak at 530.5 eV in the O 1s spectra of ACF/Fe<sub>3</sub>O<sub>4</sub> is assigned to the Fe–O bonds in Fe<sub>3</sub>O<sub>4</sub>, which further demonstrates the existence of  $Fe<sub>3</sub>O<sub>4</sub>$ .

#### Magnetic and electromagnetic wave absorption properties

Fig. 6 shows the magnetic hysteresis loop for  $ACF/Fe<sub>3</sub>O<sub>4</sub>$  over a range of  $-20 000$  to 20 000 Oe at room temperature; its shape is a typical S-curve, corresponding to the superparamagnetic behavior. The magnetic properties including the saturation magnetization  $M_s$ , magnetic coercivity  $H_c$ , and remanent



Fig. 4 SEM images of (a) ACF and (b), (c) ACF/Fe<sub>3</sub>O<sub>4</sub>, and photographic image of (d) ACF/Fe<sub>3</sub>O<sub>4</sub>.

magnetization  $M_{\rm r}$  are 22 emu  ${\rm g}^{-1}$ , 114 Oe and 3.4 emu  ${\rm g}^{-1}$ , respectively. The  $M_s$  of ACF/Fe<sub>3</sub>O<sub>4</sub> is lower than that of bulk Fe<sub>3</sub>O<sub>4</sub> (*ca.* 84.5 emu g<sup>-1</sup>), which can be explained by the small size of the  $Fe<sub>3</sub>O<sub>4</sub>$  nanoparticles and the addition of ACF, which is in good agreement with the findings of a previous study.<sup>18</sup>

as the reflectivity  $R$ , as measured using the arch method. The arch method is a simple and practical method with which the reflectivity of microwave-absorbing materials can be accurately measured without destroying their structure.10,13 Fig. 7(a) shows the microwave absorption/reflectivity curve for ACF felt. The maximum reflectivity is  $-12.9$  dB at 18 GHz, and the effective absorption bandwidth  $(R < -10$  dB) is 1.9 GHz (16.10-18 GHz).

The microwave-absorbing properties of the  $ACF/Fe<sub>3</sub>O<sub>4</sub>$  felt and ACF felt over a frequency range of 8.2–18 GHz are presented



Fig. 5 XPS spectra: (a) survey spectra of ACF and ACF/Fe<sub>3</sub>O<sub>4</sub>, (b) Fe 2p spectra of ACF/Fe<sub>3</sub>O<sub>4</sub>, and fitted O 1s spectra of (c) ACF and (d) ACF/Fe<sub>3</sub>O<sub>4</sub>.





Since ACF is a nonmetallic conductive fiber material with relative high resistance, it can be regarded as a harmonic oscillator in the electromagnetic field, which resonates with incident electromagnetic waves to form a dissipative current. Such generated photocurrent is finally consumed in the material system. ACFs can also be regarded as a dipole under the action of an electromagnetic field. The surface polarization of ACFs further dissipates and decays the photocurrent, converting the electromagnetic energy to other forms of energy under the action of the surrounding matrix.<sup>19</sup> Fig. 7(b) shows the absorption–reflectivity curve of  $ACF/Fe<sub>3</sub>O<sub>4</sub>$ . The absorption properties of ACF felt are greatly enhanced after loading with  $Fe<sub>3</sub>O<sub>4</sub>$ nanoparticles, with the maximum reflectivity reaching  $-30.7$  dB at 16.4 GHz, and the effective absorption bandwidth  $(R < -10$ dB) is 8.2 GHz (9.8–18 GHz). Such enhanced properties are mainly due to the unique electromagnetic properties of nano Fe<sub>3</sub>O<sub>4</sub>. The attached Fe<sub>3</sub>O<sub>4</sub> have functions of hysteresis loss, natural resonance and exchange resonance toward electromagnetic wave, resulting in significant attenuation of electromagnetic wave. Besides, magnetic nanoparticles increases the effective contact interface to electromagnetic wave, greatly enhancing the interfacial polarization effect. Thus, the enhanced absorbing property of the composite is owing to the synergistic effect of the electric loss ACF and the magnetic loss  $Fe_3O_4.^{20,21}$ Paper Were also controlled the maximal on 15 October 2018. The set of the set



Fig. 7 Reflectivity curves of (a) ACF and (b)  $ACF/Fe<sub>3</sub>O<sub>4</sub>$ .

## Conclusion

In summary, an  $ACF/Fe<sub>3</sub>O<sub>4</sub>$  composite was successfully prepared using an in situ reduction method. The  $Fe<sub>3</sub>O<sub>4</sub>$  was in the form of nanoparticles that are uniformly dispersed over the surface of the ACF and which exhibit superparamagnetism at room temperature. The maximum reflectivity of the ACF reached  $-12.9$  dB at 18 GHz, while the effective absorption bandwidth  $(R < -10$  dB) was 1.9 GHz. The ACF/Fe<sub>3</sub>O<sub>4</sub> felt exhibits greatly improved microwave absorption relative to the ACF felt, with a maximum reflectivity of  $-30.7$  dB at 16.4 GHz and an effective absorption bandwidth  $(R < -10$  dB) of 8.2 GHz, for a thickness of 2 mm. Together, these properties indicate that  $ACF/Fe<sub>3</sub>O<sub>4</sub>$  offers the promise of wide-ranging applicability in the fields of absorbing materials and electromagnetic shielding.

### Conflicts of interest

There are no conflicts to declare.

### Acknowledgements

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