Magnetic enhancement of carbon-encapsulated magnetite nanoparticles

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Abstract
In this paper, we report the effect of carbon encapsulation on the magnetic behavior of magnetite (Fe$_3$O$_4$) nanoparticles obtained through chemical coprecipitation. Using starch, Fe$_3$O$_4$/carbon core–shell nanoparticles were produced through a hydrothermal method. Magnetic studies revealed that both the Fe$_3$O$_4$ and Fe$_3$O$_4$/carbon core–shell nanoparticles were nearly superparamagnetic at room temperature; more importantly, carbon encapsulation significantly enhanced the room-temperature ferrimagnetism. Transmitting optical magnetic circular dichroism analysis indicated the existence of interfacial charge transfer in the Fe$_3$O$_4$/carbon heterosystem. The ferrimagnetic enhancement was attributed to the charge transfer to polarized states of the A-site in Fe$_3$O$_4$, which is equivalent to chemical reduction of the A-site iron of Fe$_3$O$_4$. We concluded that Fe$_3$O$_4$/carbon core–shell nanoparticles formed a heterosystem so that the amorphous carbon passivated the magnetic semiconducting Fe$_3$O$_4$ nanoparticles, to which the sp$^2$ electron was transferred.

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1. Introduction

Nanocomposite materials with extended functionality have drawn research attention due to their potential applications in chemistry and nanotechnology. Carbon encapsulation is a promising method for forming core–shell nanostructures because carbon is light weight, biocompatible, and thermally and chemically stable. Carbon-encapsulated magnetic nanoparticles (NPs) are interesting because of their potential biomedical applications [1–3]. Carbon encapsulation can mitigate chemical corrosion–induced instability and can modify the magnetic properties of core materials.

Magnetite (Fe$_3$O$_4$) has been extensively studied as a core material because of its ease of preparation and functionalization in the laboratory. Its superparamagnetism and low toxicity may enable Fe$_3$O$_4$ NPs to meet the growing demand for biomedical applications, such as diagnostic magnetic resonance imaging, thermal therapy, and drug delivery [4–8]. Fe$_3$O$_4$ NPs benefit biomedical applications because they are superparamagnetic [i.e., high saturation magnetization ($M_s$) and nearly zero coercivity] [9,10] at room temperature and are usually smaller than 20 nm. Nevertheless, Fe$_3$O$_4$ NPs often exhibit the chemical corrosion–induced instability, which may cause partial oxidation into γ-Fe$_2$O$_3$ [9,11]. To alleviate this, they can be coated with protective layers of various materials [12]. Such nanocomposite materials are primarily composed of Fe$_3$O$_4$ cores coated with organic or inorganic shells.

Fe$_3$O$_4$ is a soft ferrimagnetic material with low magnetic anisotropy at room temperature. Among iron oxides, it possesses the highest $M_s$ (92 emu/g) at room temperature [13]. Magnetite crystallizes into a structure with a space group of Fd$ar{3}$m at room temperature. All iron cations are in the trivalent state in maghemite (γ-Fe$_2$O$_3$), whereas magnetite contains both Fe$^{2+}$ and Fe$^{3+}$ cations. The cell content of ideal magnetite is [Fe$^{3+}$]$_8$[Fe$^{3+}$]$_8$[Fe$^{2+}$]$_8$[O$_{32}$], in which one-third of Fe ions are tetragonally coordinated with four oxygen atoms, and two-third of Fe ions are octahedrally coordinated with six oxygen atoms; which are called the A- and B-site, respectively. Magnetic moments of Fe ions in each site are
oppositely oriented, forming a ferrimagnetic configuration [14,15]. Unlike \( \gamma \)-FeO\(_2\), which is an insulator, Fe\(_3\)O\(_4\) is reported to be a semiconductor at room temperature [16].

Enhanced magnetization was reported for Fe\(_3\)O\(_4\) capped with alginic acid [17]. This enhancement was attributed to encapsulation-improved Fe\(_3\)O\(_4\) surface crystallinity. Intrinsic electronic origins accounting for magnetization enhancement in the Fe\(_3\)O\(_4\)/graphene heterostructure were also predicted from first-principle calculations [18]. A-site ions in Fe\(_3\)O\(_4\) were concluded to play a crucial role in determining the magnetism of hetero-systems. This conclusion explained the intrinsic electronic effect engendering ferrimagnetic enhancement and prompt our studies on magnetic behavior of carbon-encapsulated magnetite core–shell NPs. Fe\(_3\)O\(_4\)/carbon core–shell NPs (Fe\(_3\)O\(_4\)/C NPs) can be synthesized through various approaches, including solvothermal, mechanochemical, and thermal decomposition methods as well as hydrothermal synthesis [12,19–22]. Nevertheless, to the best our knowledge, magnetic enhancement in a carbon-encapsulated magnetite system has not been reported. A carbon shell is usually formed through a high-temperature annealing process, which carbonizes hydrocarbon precursors but also reduces iron oxide [23]. To avoid this reduction, a low-temperature carbon encapsulation procedure was employed.

In this paper, we report that room-temperature ferrimagnetic enhancement can be achieved by controlling the amount of starch precursor used when forming Fe\(_3\)O\(_4\)/C NPs hydrothermally. The preparation temperature during the hydrothermal process was found to be crucial for the carbonization of the starch precursor and therefore for magnetic enhancement. Transmitting optical magnetic circular dichroism (OMCD) confirmed the existence of interfacial charge transfer in Fe\(_3\)O\(_4\)/C NPs which engendered the enhancement of ferrimagnetism; that is equivalent to the chemical reduction of the Fe\(^{3+}\) in the \( A\)-site caused by carbon encapsulation. From our experimental observations, the band alignment of a Fe\(_3\)O\(_4\)/C heterosystem was deduced. The rest of this paper is organized as follows. Section 2 describes the sample preparation and the experimental setups. Section 3 presents the structural [X-ray diffractometry (XRD), Raman spectroscopy, and transmission electron microscopy (TEM)] and magnetic [using a vibrating-sample magnetometer (VSM)] characterization results. In Section 4, to explain the observed ferrimagnetic enhancement, we discuss the channels of charge transfer that possibly occurred in the Fe\(_3\)O\(_4\) core based on electronic configuration description. Main optical transitions in Fe\(_3\)O\(_4\) based on band descriptions are delivered. The results of OMCD experiments for determining the existence of interfacial charge transfer in the Fe\(_3\)O\(_4\)/C NPs are presented. Band alignment of the carbon-encapsulated Fe\(_3\)O\(_4\) heterosystem suggested by our VSM and OMCD experiments is also proposed. Section 5 gives the conclusions.

2. Experimental procedure

2.1. Sample preparations

Magnetite NPs were synthesized in an aqueous solution through the coprecipitation method. Magnetite NPs were prepared by slowly adding NH\(_4\)OH to an aqueous mixture of FeCl\(_3\)·6H\(_2\)O and FeSO\(_4\)·7H\(_2\)O at a 1:1 M ratio until pH was equal to 10 at room temperature. The black iron oxide product was found to be strongly magnetic. To isolate NPs, the black particles were separated from the liquid by using a bar magnet and were washed with absolute ethanol. This separation and rinsing process was repeated three times. The final product was dried under vacuum at room temperature.

Fe\(_3\)O\(_4\)/C NPs were prepared through a hydrothermal method, in which pristine 0.5-g Fe\(_3\)O\(_4\) NPs were dispersed in deionized water containing soluble starch, as previously described [9]. The resulting suspensions were transferred to the autoclave and maintained at 160 and 200°C for 12 h. After completion of the reaction, the autoclave was cooled, and the resulting Fe\(_3\)O\(_4\)/C NPs were collected using a magnet. The final product was washed with deionized water and subsequently dried under vacuum at room temperature.

XRD was performed using a BL01C beamline at the National Synchrotron Radiation Research Center in Taiwan. A monochromatic X-ray with a wavelength of 0.6888 Å was used as the probing source. Raman measurements were performed using a microscopic Raman system (RAMaker, Protrudech Co., Ltd., Tainan, Taiwan). An exciting line of 532 nm was supplied by a diode laser (CNI) with a power of 1 mW. The exposure time was 60 s with two accumulations. Calibration was conducted using a silicon standard where the band is generally observed at 520 cm\(^{-1}\). The corresponding spectral resolution was in the range of 1 cm\(^{-1}\). Saturation magnetization and \( M-H \) loop measurements were performed using a VSM (Lake Shore Model 7400, USA) under magnetic fields \( H \) up to 15,000 G. Particle sizes and shapes were studied through TEM on a JEOL JEM-2100F CS STEM. TEM samples were prepared by placing one drop of a dilute particle suspension in anhydrous alcohol on a carbon-coated copper grid and evaporating the solvent at room temperature. For OMCD measurement, a photoelastic modulator in the system (Jasco J-815) produced left-circularly polarized and right-circularly polarized light waves for the measurement. The light source was a 450-mW Xenon lamp; wavelength accuracy was ±0.3 at 250–500 nm and ±0.8 at 500–800 nm. All measurements were performed at room temperature.

3. Structural and magnetic characterization

3.1. XRD phase analysis

Fig. 1 shows the XRD results for the Fe\(_3\)O\(_4\)/C and pristine Fe\(_3\)O\(_4\) samples. The Fe\(_3\)O\(_4\)/C samples exhibited similar patterns as as-prepared Fe\(_3\)O\(_4\), suggesting the crystalline form was nearly unchanged during the hydrothermal process. The diffraction peaks could be indexed to a spinel structure of either magnetite (JCPDS no. 75-0449) or maghemite (JCPDS no. 89-5892). Phase identification of magnetite and maghemite through conventional XRD is not simple, because both have the same cubic structure with a minute distinction in lattice parameters. The (511) peak could be deconvoluted to identify and quantify mixed magnetite/maghemite [24]. Accordingly, most of the crystalline phases in the pristine Fe\(_3\)O\(_4\) NPs were magnetite (approximately 86%), and the Fe\(_3\)O\(_4\)/C samples in the study were nearly pure magnetite. However, a small amount of the hematite phase (\( \alpha \)-Fe\(_2\)O\(_3\)) was detected both in the
Fe₃O₄@C-160 and Fe₃O₄@C-200 samples. No additional crystalline phase apart from α-Fe₂O₃ after hydrothermal treatment was concluded. The average crystallite sizes ($D_{311}$) of the Fe₃O₄, Fe₃O₄@C-160, and Fe₃O₄@C-200 samples were estimated to be 18, 24, and 25 nm, respectively, through Scherrer’s formula [25].

3.2. Raman analysis

Fig. 2 shows the Raman spectrum for the phase identifications of the samples prepared from various amounts of the starch precursor. For the magnetite NPs, which were more prone to oxidation by laser heating, the output power of the Raman excitation source was reduced to 1 mW. The spectrum of the as-prepared Fe₃O₄ powder had a main peak at 660–680 nm, consistent with the peaks described in the related literature [26]. All Fe₃O₄@C in this region were in accordance with the characteristic bands of magnetite. In the high wavenumber region (1100–1800 cm⁻¹), the Fe₃O₄@C-200 samples exhibited a typical spectrum of diamond-like carbon [27]. This band region could be deconvoluted into a broad band at 1540–1580 cm⁻¹ (G band) and another band at 1330–1350 cm⁻¹ (D band). The intensity ratio ($I_D/I_G$) was found to correlate linearly with the $sp^3/sp^2$ ratio [27]. The $I_D/I_G$ ratios remained nearly the same (approximately 4) for the samples prepared from various amounts of starch. The $sp^3/sp^2$ ratio was estimated to be approximately 2.5 based on the data reported by Ferrari and Robertson [27,28]. Moreover, $I_D$ and $I_G$ became more prominent when more starch was used as a carbon source when forming the Fe₃O₄@C-200, whereas those in Fe₃O₄@C-160 were severely suppressed, suggesting that the starch was not fully carbonized through hydrothermal treatment at 160 °C. According to our Raman analysis, the hydrothermal carbonization process converted organic starch into a carbon-like product at 200 °C.

3.3. Saturation magnetization and M–H loop measurements

Fig. 3 shows the saturation magnetization ($M_s$) of pristine Fe₃O₄ NPs (approximately 89 emu/g) and Fe₃O₄@C-200 samples; the mean crystallite sizes estimated from XRD and TEM are also indicated. $M_s$ increased initially until reaching a maximum value of 134 emu/g when approximately 0.19 g of starch precursor before gradually decreasing when additional starch content was used. Both TEM and Scherrer’s analyses showed a consistent diameter growth of ~1.5 times with carbon encapsulation, but saturate and then slightly decrease with the increasing amount of starch precursor. Possible explanation for the decreasing crystallite size of Fe₃O₄@C-200 was that a large amount of starch facilitates the heterogeneous polymerization on the surface of magnetite, thus hindering the further growth of magnetite NPs.

The $M_s$ of Fe₃O₄@C-200 was usually higher than that of the pristine samples. The significantly decreased $M_s$ of samples that were prepared using as much starch as the carbon precursor can be explained by the relatively low Fe₃O₄ quantity because the difference in the mean crystallite size revealed by XRD was only in the range of 2 nm. These observations suggest that the room-temperature ferrimagnetic enhancement can be achieved through precise control of the amount of starch precursor. The effect of the preparation temperature on $M_s$ was also examined using samples produced from the same amount of starch (0.19 g) precursor added...
in the hydrothermal process. Fig. 4 shows the $M-H$ loops of the as-prepared Fe$_3$O$_4$ NPs, Fe$_3$O$_4$@C-160, and Fe$_3$O$_4$@C-200 samples. All displayed nearly superparamagnetic behavior. The $M_s$ value of Fe$_3$O$_4$@C-200 was significantly higher than those of the Fe$_3$O$_4$@C-160 and Fe$_3$O$_4$ powders. The crystalline phase and the mean crystallite size shown on XRD were almost the same for Fe$_3$O$_4$@C-160 and Fe$_3$O$_4$@C-200. Therefore, the observed 35% magnetic enhancement of Fe$_3$O$_4$@C-200 was primarily related to the formation of the core–shell interface in Fe$_3$O$_4$@C.

3.4. TEM measurements

Fig. 5 shows the TEM images of as-prepared Fe$_3$O$_4$ and Fe$_3$O$_4$@C-200 produced from 0.19 g of the starch precursor. The dark core of the Fe$_3$O$_4$ NPs and the grey carbon shell are clearly observable in the TEM image of the Fe$_3$O$_4$@C-200 sample. Although the Fe$_3$O$_4$@C-200 sample was enlarged after the hydrothermal process, the $M_s$ have shown excessively higher than the bulk Fe$_3$O$_4$ (92 emu/g) at room temperature [13], suggesting the intrinsic electronic origins rather than NP size effect were accounted for the observed ferrimagnetic enhancement. To the best of our knowledge, the observation of room-temperature magnetic enhancement in amorphous carbon-encapsulated magnetite NPs has not been reported elsewhere. The typical carbon shell thickness was estimated to be 2 nm, and that may suggest the appropriate diameter-to-thickness ratio of Fe$_3$O$_4$@C NPs is a prerequisite for observation of this enhancement.

4. Origins of magnetic enhancement

4.1. Electronic configuration description

In this section, we discuss the charge transfer that possibly occurs to explain the observed ferrimagnetic enhancement based on the electronic configuration of Fe$_3$O$_4$. The cubic phase of Fe$_3$O$_4$ comprises a mixture of Fe$^{3+}$ and Fe$^{2+}$ ions in the B-site, whereas only Fe$^{3+}$ ions exist in the A-site. The spins of Fe ions in each site are in opposite directions, forming a ferrimagnetic configuration. Following Hund’s rule, the high-spin configurations for each subcell are as follows: the Fe$^{3+}$ ($t^6$) in the A-site can be represented by $|t^6_t^3_e^0_e^0_t^3>\text{or} |t^6_t^3_e^0_e^0_t^3+$; the Fe$^{2+}$ ($d^5$) and Fe$^{3+}$ in the B-site can be represented by $|t^5_3_e^2_2>$ and $|t^5_3_e^2_2>$, respectively. Here, the notations 1 and 2 denote spin direction parallel (majority) and antiparallel (minority) to total magnetization ($\uparrow$). We suggest that two channels possibly account for the experimentally observed magnetic enhancement. First, a $1$ electron may be transferred to the empty $e$ state of the A-site Fe$^{3+}$ [$A(e)$], which is equivalent to the chemical reduction of Fe$^{3+}$ in the A-site; second, a $2$ electron may be removed from the $t_{2g}$ state of the B-site Fe$^{2+}$ [$B(t_{2g})$], which is equivalent to the oxidation of Fe$^{2+}$ in the B-site. Here, we use the formulae: $e^0_1 \leftrightarrow t^7_1$, $e^0_1 \leftrightarrow t^7_1$ and $e^0_2 \leftrightarrow t^7_1 \leftrightarrow t^7_1 \leftrightarrow e^0_2$, with $t^6_2$ and $t^6_1$ (denotes the empty state to/from which a polarized electron is transferred) to represent the two possibilities accounting for the observed saturation moment enhancement of Fe$_3$O$_4$@C-200. Each represents electron addition to the $e$ state of the A-site and electron removal from the $t_{2g}$ state of the B-site, respectively. Both cases explain the increase in total magnetic moment but with opposite flow directions and sign of spin that may occur in the Fe$_3$O$_4$@C heterosystem.

Amorphous carbon (a-C) contains both a three-fold coordinated sp$^3$ configuration, as in graphite, and four-fold sp$^3$ sites, as in diamond. For sp$^3$ sites, each of the four valence electrons lies in a sp$^3$ hybrid, which then forms a $\sigma$ bond with a neighbor. At a sp$^3$ site, only three of these electrons are used in the $\sigma$ bond; the fourth enters a $\pi$ orbital that lies normal to the $\sigma$-bond plane. The $\pi$ states are only weakly bonded and therefore are the most likely electron donors at the heterointerface, which infers that the transfer of the $\pi$ electron to the A-site of Fe$_3$O$_4$ engenders the ferrimagnetic magnetic enhancement.
4.2. Electronic structure and spin-dependent optical transitions in magnetite

Density functional theory (DFT) employing Hubbard-\(U\) correction (DFT + \(U\)) [29] has proven to be essential for describing \(\text{Fe}_3\text{O}_4\) in terms of optical and magnetic properties [30–35]. On-site Coulomb interactions (\(U\)) rectify the overestimated \(p-d\) hybridization and provide accurate energy positions for both occupied and unoccupied states. Our recent studies [30] have concluded that the transmitting OMCD spectra can describe the valence electronic structures after consideration of electron correlation effects, and hence the dominant spin-dependent optical transition in magnetite. Fig. 6(a) shows the spin-polarized projected density of states of \(\text{Fe}_3\text{O}_4\). The valence structure comprises mainly crystal field splitting of \(\text{Fe}(3d)\) into \(t_2g\) and \(e_g\) group hybridized with \(O(2p)\) bands. The notable half-metallicity is indicated by the \(B\)-site \(t_2g\) spin-minority band, denoted by \(B(t_{2g})\)\(\uparrow\), across the Fermi level \((E_F)\). A valence gap of spin-majority (\(-2.2\) eV) is formed by the unoccupied electronic states of \(A\)-site, denoted by \(A(e_{t_2})\)\(\uparrow\), and the occupied \(B(e_g)\)\(\uparrow\); a relatively smaller valence gap of spin-minority (\(-2.0\) eV) is formed by \(B(t_{2g})\)\(\downarrow\) and \(O(2p)\)\(\downarrow\). The most relevant to optical transition is the band interval between binding energy \(2\) and \(-6\) eV, of which the spin-majority and -minority are dominated by polarized Fe(3d) and O(2p) bands, respectively. The occupied electronic states of \(A\)-site \(A(e_{t_2})\)\(\downarrow\) are located deeper beyond this region, and \(A(e_{t_2})\)\(\uparrow\) immediately above the \(E_F\) is fully empty.

Fig. 6(b) shows the schematic site-projected density of states of \(\text{Fe}_3\text{O}_4\), in which the main optical transitions are depicted by arrows. Main transitions are categorized into three types, namely of intra-site, inter-site, and ligand-to-metal optical transition. They are also known as intervalence charge transfer (IVCT), intersublattice charge transfer (ISCT), and ligand-to-metal charge transfer (LMCT), respectively. The IVCT relaxing from Laporte selection is mainly minority-spin transfer of \(B(t_{2g})\)\(\downarrow\) \(\rightarrow\) \(B(e_g)\)\(\uparrow\); the ISCT, mediated by \(2p\) oxygen, is the majority-spin transfer of \(B(e_g)\)\(\uparrow\) \(\rightarrow\) \(A(e_{t_2})\)\(\uparrow\). The LMCT is the minority-spin transfer of \(O(2p)\)\(\downarrow\) \(\rightarrow\) \(B(t_{2g})\)\(\downarrow\).

OMCD in Faraday geometry manifesting the absorptive difference of incident photons with spins parallel (\(\sigma_+\)) and antiparallel (\(\sigma_-\)) to the applied magnetic field was recorded simultaneously as a function of wavelength. With the resolution of both energy and dominant absorptive spin-channel, the OMCD signal (\(\sigma_+ - \sigma_-\)) can resolve the dominant channels of spin-dependent optical transition in a spin-split valence system. This technique enables a simple laboratory probe for the interfacial charge transfer in the nanocomposite through observations of the relative shift of spectrum between the pristine \(\text{Fe}_3\text{O}_4\) NPs and the \(\text{Fe}_3\text{O}_4@\text{C}\) heterosystem. Fig. 7 shows the OMCD spectra of pristine \(\text{Fe}_3\text{O}_4\) NPs and \(\text{Fe}_3\text{O}_4@\text{C}-200\) at the zero field and the saturation field of \(\pm 0.8\) T. The OMCD profile of the pristine \(\text{Fe}_3\text{O}_4\) NPs is consistent with that described in the literature [37–39]. It mainly comprises regions according to the sign of the OMCD signal. The interval of negative OMCD from pristine \(\text{Fe}_3\text{O}_4\) NPs \(1.50–2.30\) eV, as depicted in Fig. 7(c), is attributed to the IVCT of \(B(t_{2g})\)\(\downarrow\) \(\rightarrow\) \(B(e_g)\)\(\uparrow\) [dotted line in Fig. 6(b)] i.e., the crystal-field transition [40] of the \(B\)-site. The broad band with positive OMCD (\(2.30–4.00\) eV) mainly reflects the competition between the ISCT of \(B(e_g)\)\(\downarrow\) \(\rightarrow\) \(A(e_{t_2})\)\(\uparrow\) [dashed line in Fig. 7(b)] and the LMCT of \(O(2p)\)\(\downarrow\) \(\rightarrow\) \(B(t_{2g})\)\(\downarrow\) [solid line in Fig. 7(b)], which are characterized by the paramagnetic response of the main raising edge (\(2.30–2.86\) eV) and the diamagnetic response of the falling edge (\(2.86–4.00\) eV) in the broad band, respectively. Note that the onset of ISCT define the optical gap of spin majority, which was enlarged for \(\text{Fe}_3\text{O}_4@\text{C}-200\) as suggested from the OMCD blue shift (\(0.25\)-eV) of the main raising edge. This blue shift can also be considered from the work function difference of the core-shell material. The work function (\(\phi\)) of \(\text{C}-\text{Fe}\) determined by Keivin probe can vary in the range 4–5 eV [41]; that of bulk \(\text{Fe}_3\text{O}_4\) was reported to be 5.78 eV, truncated to 5.32 eV at the surface [42,43]. Hence, the \(\text{Fe}_3\text{O}_4@\text{C}\) heterosystem can be represented by a metal to magnetic-semiconducting contact of the case of \(\phi_m<\phi_m\), which also indicates the charge flow direction is from the metal (\(a\)-C) to the semiconducting (\(\text{Fe}_3\text{O}_4\)) side. The shift value of 0.25-eV is close to the work function difference (\(\phi_m - \phi_m\)), suggesting the channel of

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**Fig. 6.** (a) Spin-polarized projected density of states (PDOS) of \(\text{Fe}_3\text{O}_4\) (Fd\(\overline{3}\)m) calculated through DFT + \(U\) as implemented in Vienna Ab initio Simulation Package (VASP) [36]. (Left panel) Minority-spin DOS; (Right) majority-spin DOS. The simulation is \(\text{Fe}_3\text{O}_4@\text{C}-160\) with 56 total atoms in a cell of ferrimagnetic configuration. The Fermi level (\(E_F\)) is indicated by the dotted horizontal line. For calculation details refer to Ref. [30]. (b) Schematic of valence band electronic structure and associated optical transitions in \(\text{Fe}_3\text{O}_4\). Dotted, dashed, and solid lines represent the inter-valence charge transfer (IVCT), inter-sublattice charge transfer (ISCT), and ligand-to-metal charge transfer (LMCT) which corresponding to minority-spin transfer of \(B(t_{2g})\)\(\downarrow\) \(\rightarrow\) \(B(e_g)\)\(\uparrow\), majority-spin transfer of \(B(e_g)\)\(\uparrow\) \(\rightarrow\) \(A(e_{t_2})\)\(\uparrow\), and majority-spin transfer of \(O(2p)\)\(\downarrow\) \(\rightarrow\) \(B(t_{2g})\)\(\downarrow\), respectively.
electron addition in the A-site $\text{Fe}^{3+}$ ($e^2 \rightarrow t_2^1 e^1$) of Fe$_3$O$_4@C$-200 due to carbon encapsulation. Therefore, the OMCD blue shift have implied the charge transfer into the polarized $e$ band of the A-site.

4.4. Band alignment

The observations of the ferrimagnetic enhancement and the OMCD blue shift of Fe$_3$O$_4@C$-200 sample have indicated the existence of spin-polarized interfacial charge transfer in a core–shell heterosystem. We propose the band alignment of the Fe$_3$O$_4@C$ heterosystem, as shown in Fig. 8. Two isolated systems of $\varphi_m < \varphi_{ms}$ were referred to a common vacuum level. The band descriptions of $\alpha$-C generally suggest the $\pi$ states lie on the $E_F$ [43]. We focused on the bands of majority spin, which essentially account for the ferrimagnetic enhancement. The conduction band maximum (CBM) and valence band minimum (VBM) of Fe$_3$O$_4$ are represented by $A(e^2; t_2^1)$ and $B(e^1_g)$, respectively. The optical gap of the majority spin joining the CBM and VBM is approximately 2.0 eV. When forming the contact, the $\pi$ electrons in the vicinity of the $E_F$ flow into the polarized $A(e^2; t_2^1)$ band until the induced dipole field brings the $E_F$ on the two sides together, and that leads to the magnetic moment enhancement in Fe$_3$O$_4@C$-200 as well as the increasing optical gap of majority spin. The later engendering the OMCD blue shift (approximately 0.25 eV) as well as the ferrimagnetic enhancement was demonstrated by a valence band offset ($\Delta_{\text{VBO}}$) of the Fe$_3$O$_4@C$ heterosystem.

Fig. 7. OMCD spectra of (a) pristine Fe$_3$O$_4$ NPs and (b) Fe$_3$O$_4@C$-200 samples. (c) Overlapping of these two curves (eV) under the applied voltage of 0.8 T. Blue shift of the main raising edge representing ISCT of $B(e^1_g) \rightarrow A(e^2; t_2^1)$ [dashed line in Fig. 6(c)] is indicated by the arrow. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

![Fig. 7](image-url)

![Fig. 8](image-url)

Fig. 8. (Top) Schematic diagram of valence electron states of isolated amorphous carbon ($\alpha$-C) and Fe$_3$O$_4$. Vacuum level and Fermi energy are indicated by VAC and $E_F$ (thin dashed line), respectively. CBM and VBM are represented by the unoccupied state of the A-site [$A(e^2; t_2^1)$] and the occupied state of B-site [$B(e^1_g)$]; $\varphi_m$ and $\varphi_{ms}$, as indicated by the arrow lines, are work functions of $\alpha$-C (metal) and Fe$_3$O$_4$ (magnetic-semiconductor). The optical gap of majority spin ($E_g$) is indicated by the thick arrow joining VBM and CBM. (Bottom) Band alignment in representing Fe$_3$O$_4@C$ as a metal-to-magnetic semiconducting contact. The heterointerface is indicated by the thin vertical solid line. The valence band offset ($\Delta_{\text{VBO}} = 0.25$ eV) of the heterosystem obtained through OMCD measurement is also indicated.
5. Conclusions

The magnetization properties of carbon-encapsulated Fe₃O₄ NPs synthesized through chemical coprecipitation were examined in this study. Fe₃O₄/C particles were formed through the hydrothermal method, with the Fe₃O₄ NPs as a magnetic core and soluble starch as a carbon source. The Mₛ of Fe₃O₄/C samples was generally higher than that of pristine Fe₃O₄ after hydrothermal treatment at 200 °C. The magnetite–carbon interface charge transfer played a major role in the increase in ferrimagnetic magnetization. We proposed the observed 35% magnetic enhancement was primarily attributed to electron addition of the heterojunction so that the higher than that of pristine Fe₃O₄ after hydrothermal treatment at equivalent to the chemical reduction of synthesized through chemical coprecipitation were examined in and stability than those of pure Fe₃O₄ can benefit medicinal applications.

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