

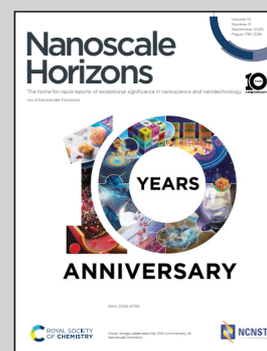
Showcasing research from Professor Nguyen Thi Kim Thanh's laboratory, Department of Physics and Astronomy, University College London, London, UK.

$\text{Co}_{0.6}\text{Fe}_{0.4}\text{O}-\text{Co}_{1.4}\text{Fe}_{1.6}\text{O}_4$  core-shell nanoparticles with colossal exchange bias

The work explored the interplay between antiferromagnetic and ferrimagnetic phases in a colloidally stable core-shell nanoparticle system, which exhibits a record exchange bias of 10.34 kOe. The nanoparticles were synthesised *via* a simple, highly reproducible one-step thermal decomposition method. Atomic-resolution STEM confirms the epitaxial relationship between the core and the shell. Time-of-flight neutron diffraction and magnetometry measurements reveal a Néel temperature of 397 K in the antiferromagnetic core. These findings highlight the system's potential for future applications in nanospintronics and nanomedicine.

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# Co<sub>0.6</sub>Fe<sub>0.4</sub>O–Co<sub>1.4</sub>Fe<sub>1.6</sub>O<sub>4</sub> core–shell nanoparticles with colossal exchange bias†

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The exchange bias (EB) effect is widely utilized in spintronics with 2D materials like thin films. Exploring the EB effect in nanoparticles opens up tremendous opportunities, such as miniaturization of devices, enhanced efficiency, and tunable properties, all of which are size-dependent. Due to the increased surface area to volume ratio, magnetic nanoparticles display unique characteristics, allowing for the manipulation of their magnetic properties, such as the EB effect commonly observed between antiferromagnetic (AFM) and ferro-/ferrimagnetic (FM/FiM) materials. This work employs a simple and highly reproducible one-step thermal decomposition method to fabricate colloiddally stable Co<sub>0.6</sub>Fe<sub>0.4</sub>O–Co<sub>1.4</sub>Fe<sub>1.6</sub>O<sub>4</sub> core–shell (CS) nanoparticles with a lattice-matched interface and strong exchange coupling. We investigate their temperature and field-dependent magnetic properties using time-of-flight neutron diffraction and magnetometry. These nanoparticles exhibit the highest reported EB values among core–shell nanoparticles, reaching a maximum of 10.34 kOe. Additionally, the core exhibits antiferromagnetism above room temperature, with a Néel temperature of approximately 397 K, making it more suitable for high-temperature applications. This study paves the way for designing core–shell biphasic nanoparticles to enhance the EB effect and tune the effective magnetic anisotropy, offering potential future applications in nanospintronics and nanomedicine.

## New concepts

The exchange bias effect is vital in modern magnetic storage, sensors, and information processing technologies. While state-of-the-art devices utilize this effect using layered antiferro-ferro/ferrimagnet thin films, synthesizing colloiddally stable biphasic nanoparticles remains challenging due to complex multi-step processes. We have achieved a major advancement in obtaining record colossal exchange bias by synthesizing high-quality core–shell Co<sub>0.6</sub>Fe<sub>0.4</sub>O–Co<sub>1.4</sub>Fe<sub>1.6</sub>O<sub>4</sub> colloiddal nanoparticles using thermal decomposition. Using double aberration-corrected atomic resolution transmission electron microscopy, we confirm the epitaxial relation between the antiferromagnet and ferrimagnet layer. Additionally, the core exhibits a Néel temperature of 397 K, as determined using time-of-flight neutron diffraction and magnetometry. This provides insightful knowledge on the magnetic phase transition of antiferromagnetic system at the nanoscale. The conceptual novelty lies in the straightforward and highly reproducible fabrication process, record colossal exchange bias, and a redefined phase stability limit for cobalt-iron oxide antiferromagnetic nanoparticles. Exchange bias in nanoparticle systems opens tremendous technological opportunities, such as the miniaturization of devices, enhanced efficiency and reactivity, better dispersibility for flexible applications, and tunable magnetic, electronic, and optical properties, all of which are size-dependent. Furthermore, the observed Néel temperature in the antiferromagnet core enhances its potential for high-temperature applications.

## Introduction

Magnetic nanoparticles have garnered significant attention in recent years due to their potential applications across various fields, such as biomedicine, catalysis, data storage, high-performance permanent magnets, and spintronic devices.<sup>1–5</sup> Due to a high surface-area-to-volume ratio, these nanoparticles exhibit unique size-dependent properties. One such property is the exchange bias (EB) effect, an interfacial phenomenon where an antiferromagnetic (AFM) material couples with a ferro-/ferrimagnetic (FM/FiM) material (Fig. 1), resulting in unidirectional anisotropy. The anisotropy is due to the “pinning” of the FM/FiM spins by the spins of the AFM material at the interface, which results in the shift of the magnetic hysteresis loop.<sup>6</sup> The EB effect is widely utilized in the spintronic sector for data storage and information processing applications using thin

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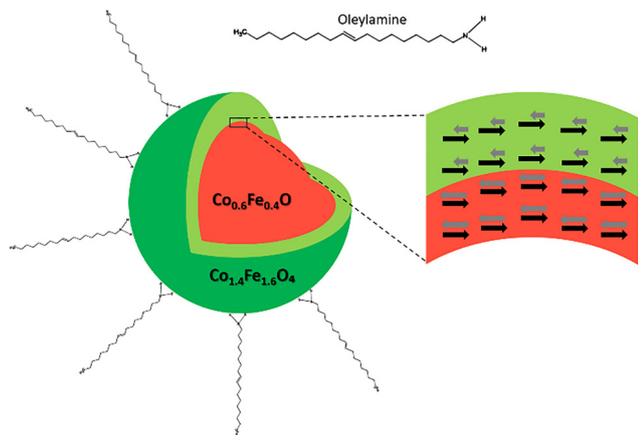


Fig. 1 Schematic illustration of the colloiddally stable AFM-FiM core-shell nanoparticle.

films and single-phase materials.<sup>7–17</sup> This can be attributed to well-established fabrication protocols and a deep understanding of the interfaces between the two phases. However, practical applications of the EB effect using nanoparticles are virtually non-existent. As a result, the advantages of nanoparticles, such as device miniaturization, enhanced efficiency, and tunability of properties due to their size dependence, have not been fully realized. A significant challenge in nanoparticle-based systems is the superparamagnetic limit, which results in low or insignificant anisotropy.<sup>18</sup> Engineering a core-shell system with lattice-matched interfaces offers a potential solution, as it can help enhance and tune the magnetic anisotropy of nanoparticles.

In general, the EB effect is influenced by several factors, such as the formation of defects, the thickness of the AFM and FM/FiM layers, the interface's roughness, the material's crystallinity, and the interface's crystallographic orientation.<sup>19–25</sup> In the case of nanoparticles, it is more complex as additional factors such as disordered surface spins and interparticle (dipolar) interactions can also affect the observed shift in the hysteresis loop. In ultrasmall nanoparticles (typically <3 nm), disordered surface spins can couple with the ordered core spins, and the differences in the magnetic anisotropies can lead to the EB effect.<sup>26–29</sup> The choice of core and shell materials has been shown to affect dipolar interactions, which can influence the EB effect. For example, in nanoparticles with magnetically soft core and shell materials, the EB effect is

significantly influenced by dipolar interactions. In contrast, in systems containing magnetically hard and soft materials, EB primarily originates from interfacial exchange interactions.<sup>30</sup> Studies reporting giant EB generally employ a two-step synthesis process. First, core materials like Fe are synthesized using a wet chemical or gas-phase method, followed by the growth of a shell material like Fe<sub>2</sub>O<sub>3</sub> through an oxidation process.<sup>31</sup> A wide variety of core-shell systems, such as FeO-Fe<sub>3</sub>O<sub>4</sub>, Fe-Fe<sub>3</sub>O<sub>4</sub>, CoO-CoFe<sub>2</sub>O<sub>4</sub>, Co-CoO, MnO-Mn<sub>3</sub>O<sub>4</sub>, have shown varying EB due to differences in composition, size, interface quality, and material selection.<sup>32–36</sup> Recent work demonstrated a 9 kOe EB in Co<sub>0.3</sub>Fe<sub>0.7</sub>O-Co<sub>0.6</sub>Fe<sub>2.4</sub>O<sub>4</sub> core-shell nanoparticles at 10 K under a 50 kOe cooling field, though lower than our system's value.<sup>37</sup> Core-shell morphologies using transition metal oxides offer a promising approach for tuning nanoparticles' effective magnetic anisotropy ( $K_{\text{eff}}$ ). A comparison between the current work and other core-shell systems is presented in Table 1.

Here, we employ a one-step thermal decomposition technique to synthesize Co<sub>0.6</sub>Fe<sub>0.4</sub>O-Co<sub>1.4</sub>Fe<sub>1.6</sub>O<sub>4</sub> core-shell (CS) biphasic nanoparticles, which exhibit a remarkably high EB of 10.34 kOe. For comparison, we have also synthesized pure cobalt monoxide (CO) and cobalt ferrite (CFO) nanoparticles in similar conditions. Here, the spinel phase has a general formula AB<sub>2</sub>O<sub>4</sub>, where A and B represent the cations in the tetrahedral and octahedral sites, respectively. Ligands attach to the nanoparticles' surface during synthesis, enhancing their colloidal stability and achieving a narrow size distribution. This method not only improves the nanoparticles' homogeneity but also enhances the reproducibility of the synthesis process.<sup>40</sup> These factors are crucial in nanoparticle synthesis, as even slight variations in size or interface area can lead to significant differences in the nanoparticles' properties and behavior. To our knowledge, 10.34 kOe is the highest reported EB in core-shell nanoparticles. The AFM core in the CS system and the pure CO nanoparticles also display Néel temperature ( $T_N$ ) at around 397 K, which is higher than the reported bulk value of CoO at around 293 K.<sup>41</sup> This is attributed to Fe substitution in the CoO lattice and/or nanoscale effects, supported by neutron diffraction refinement and magnetometry results. The atomic-resolution transmission electron microscope (TEM) image shows a smooth lattice-matched interface between the core and the shell. The CS system has a significantly higher  $K_{\text{eff}}$  value than the pure CO nanoparticles. This is attributed to (1) the strong exchange coupling between the AFM and FiM spins, (2) the high crystallinity of the Co<sub>0.6</sub>Fe<sub>0.4</sub>O core, and (3) its higher phase fraction in the CS system.<sup>16</sup>

Table 1 A comparison of EB core-shell nanoparticles presented in this work with other systems in the literature

Composition (core-shell)	EB (kOe)	Size (nm)	Cooling field (kOe)	Synthesis method	Ref.
Co <sub>0.6</sub> Fe <sub>0.4</sub> O-Co <sub>1.4</sub> Fe <sub>1.6</sub> O <sub>4</sub>	10.3	8.6	40	TD <sup>a</sup>	This work
Fe <sub>5</sub> C <sub>2</sub> -Fe <sub>3</sub> O <sub>4</sub>	0.5	18.0	50	TD	38
Co-CoO	0.8	17.5	10	Sol-gel	39
Co-CoO	3.6	12.0	50	TD + oxidation	35
CoFe <sub>2</sub> O <sub>4</sub> -CoO	3.7	12.4	70	Seed-mediated polyol	34
CoO-CoFe <sub>2</sub> O <sub>4</sub>	5.6	14.9	70	Seed mediated TD	18
Co <sub>0.3</sub> Fe <sub>0.7</sub> O-Co <sub>0.6</sub> Fe <sub>2.4</sub> O <sub>4</sub>	9.0	9.0	50	TD	37

<sup>a</sup> TD = Thermal decomposition.

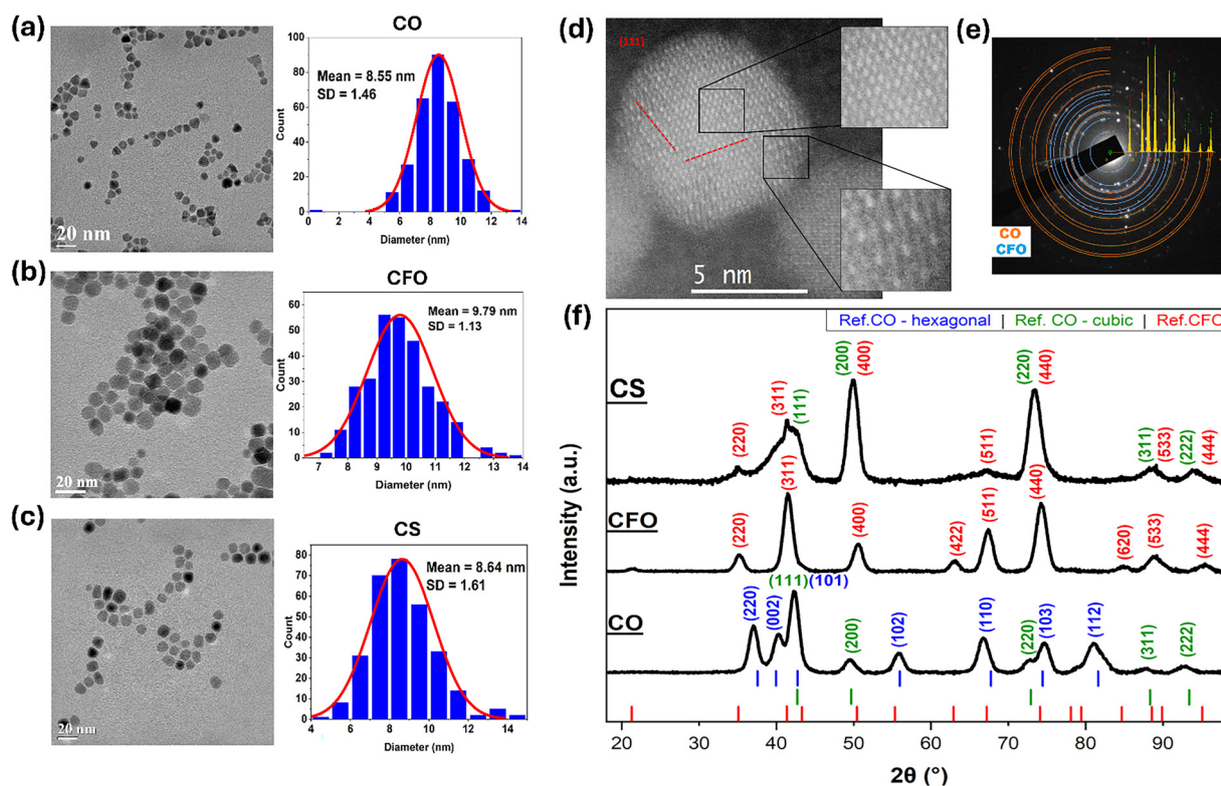


## Results and discussion

The X-ray diffraction (XRD) pattern of synthesized samples shows no impurities. The pure CO peaks match with hexagonal close-packed ( $P63mc$ ,  $a = 3.21 \text{ \AA}$ , and  $c = 5.24 \text{ \AA}$ ) and face-centered cubic ( $Fm\bar{3}m$ ,  $a = 4.26 \text{ \AA}$ ) JCPSD reference pattern (Fig. 2f) where the most intense lines are that of hexagonal crystal plane (101). The CFO matches with the cubic ( $Fd\bar{3}m$ ,  $a = 8.40 \text{ \AA}$ ) crystal structure as expected, whereas the CS nanoparticle is a mixture of cubic CO and CFO. The strongest peak in the CS nanoparticles corresponds to the cubic CO (200) and CFO (400) planes. Hence, we assume that the AFM core in CS nanoparticles has a face-centered cubic structure ( $Fm\bar{3}m$ ). This is confirmed by atomic resolution TEM (Fig. 2d).

Pure CO and CFO nanoparticles exhibit well-resolved peaks that were fitted in OriginPro software. A rough estimate of the crystallite size was performed using the Scherrer equation for comparison with TEM and dynamic light scattering (DLS) measurements. The crystallite size of pure CFO obtained using the (311) peak is 7.08 nm. The crystallite size of cubic CO obtained using (200) is 6.87 nm, whereas the hexagonal CO obtained using (220) is 6.85 nm in the pure CO sample. The phase fraction is estimated using the third most intense (220) cubic and (002) hexagonal peaks (eqn (S1), ESI<sup>†</sup>). Cubic CO accounts for 24.11% of the sample, whereas the rest contain the hexagonal crystal structure. For the CS sample, only the  $32^\circ$ – $42^\circ$

peaks could be deconvoluted due to significant peak overlap in others (Fig. S2, ESI<sup>†</sup>). The crystallite size of the AFM and FiM phases in the CS sample is 3.95 nm and 2.57 nm, respectively (Table 2). All fittings achieved an adjusted  $R^2$  value greater than 0.99. The TEM images display the spherical morphology of all synthesized nanoparticles with a narrow particle size distribution (Fig. 2a–c). The mean sizes of CO, CFO, and CS are 8.55 nm, 9.79 nm, and 8.64 nm, respectively, which are similar to the sizes obtained using the Scherrer equation (Table 2). Since nanoparticles smaller than  $\sim 80 \text{ nm}$  are typically single-domain,<sup>42</sup> the synthesized nanoparticles are expected to be single-domain. Low magnification TEM images of all three nanoparticles are shown in Fig. S1a–c in the ESI.<sup>†</sup> Further investigation of the CS nanoparticles using atomic resolution Scanning TEM (STEM) reveals a well-matched lattice interface between the core AFM and the shell FiM phase, with the CO(111) crystal plane separating the spinel (outer) and rock-salt (inner) structures. The differences in the core and shell lattice can be observed in the magnified image (Fig. 2d). The epitaxial relation between the AFM and the FiM phases is further supported by the overlapping XRD peaks in the CS sample, such as the cubic CO (200) and CFO (400) planes, as well as the cubic CO (220) and CFO (440) planes (Fig. 2f). Selected area electron diffraction (SAED) clearly shows the presence of two structural phases: a rock salt structure associated with the AFM core and a spinel structure corresponding



**Fig. 2** TEM image of (a) pure CO, (b) pure CFO, and (c) biphasic CS nanoparticles with their particle size distribution ( $n = 300$ ). (d) Double aberration corrected atomic-resolution STEM image of a CS nanoparticle with magnified core and shell section. CO(111) plane separates the core and the shell phases. (e) SAED pattern showing diffraction rings corresponding to the CO (orange) and CFO (light blue) phases. (f) XRD of all samples with hexagonal CO (reference code: 01-089-2803), cubic CO (reference code: 01-071-1178), and cubic CFO (reference code: 01-088-2152) reference patterns.



Table 2 Comparison of XRD crystallite size of all phases with their TEM and DLS diameters

Sample	Crystal phase	Magnetic behavior	$D_{\text{XRD}}$ (nm)	$D_{\text{TEM}}$ (nm)	$D_{\text{DLS}}$ (nm)
Pure CO	Cubic	AFM	$6.87 \pm 0.12$	$8.55 \pm 1.46$	$11.74 \pm 4.89$
	Hexagonal	AFM	$6.85 \pm 0.03$		
Pure CFO	Cubic	FiM	$7.08 \pm 0.02$	$9.79 \pm 1.13$	$13.53 \pm 5.54$
Biphasic CS	Cubic	AFM	$3.95 \pm 0.15$	$8.64 \pm 1.61$	$10.16 \pm 5.22$
	Cubic	FiM	$2.57 \pm 0.18$		

to the FiM shell (Fig. 2e). DLS is used to measure the solvodynamic size, *i.e.*, the size of colloidal nanoparticles in a solvent. The DLS size-distribution of the synthesized nanoparticles functionalized with ligands and dispersed in *n*-hexane is shown in Fig. S1d, ESI†. CO, CFO, and CS are 11.74 nm, 13.52 nm, and 10.16 nm, respectively, which agrees with the size determined by TEM and XRD (Table 2).

Neutron diffraction measurements were conducted on the biphasic CS powder sample over a temperature range of 2 K to 813 K using the WISH diffractometer at ISIS, the UK Neutron and Muon source.<sup>43</sup> Multi-phase Rietveld refinement was performed against the variable temperature data using FullProf,<sup>44</sup> with representative data and fits shown in Fig. 3a–c. The temperature dependences of other key parameters extracted

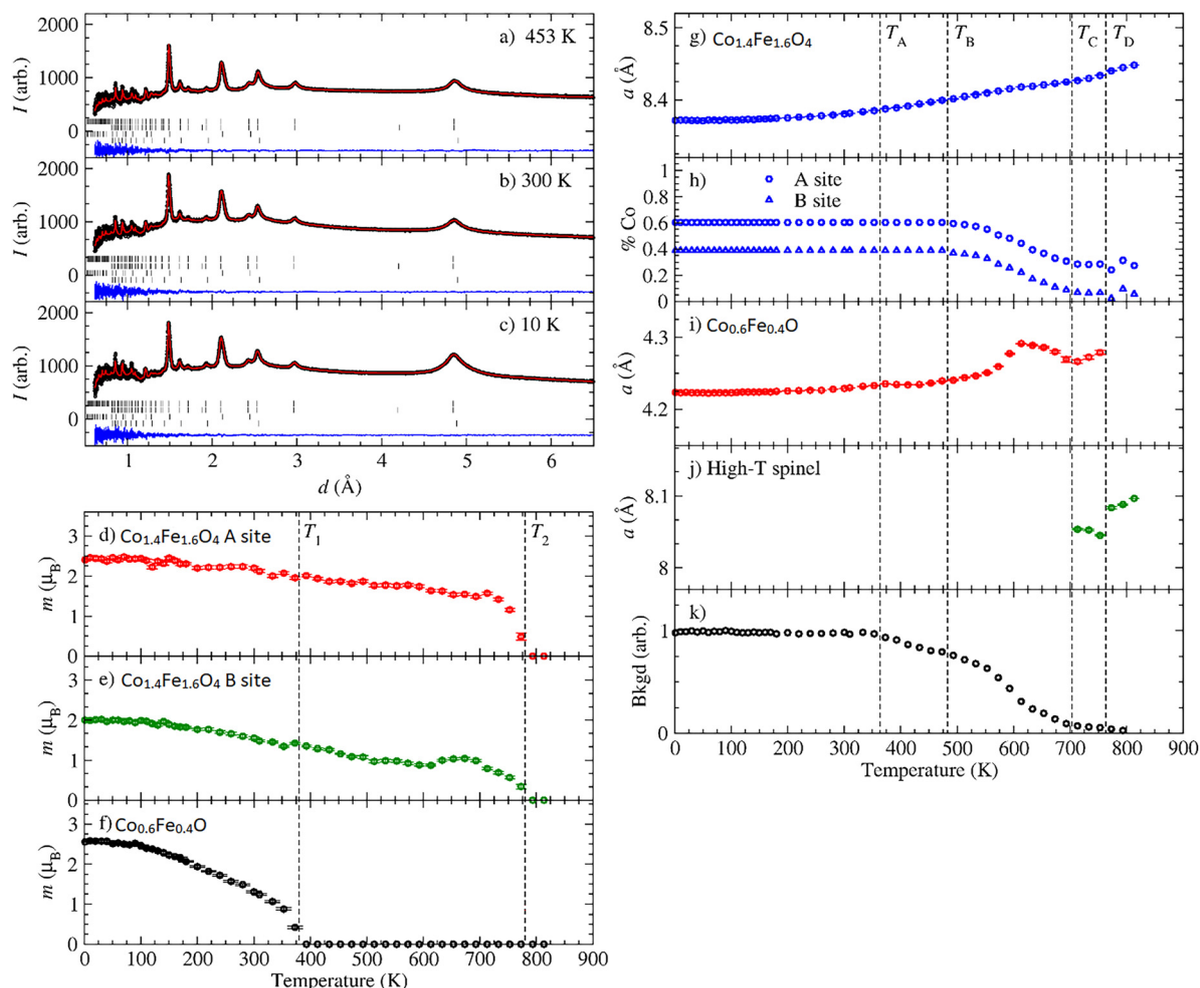


Fig. 3 (a)–(c) Representative Rietveld refinements of neutron powder diffraction data of the AFM–FiM  $\text{Co}_{0.6}\text{Fe}_{0.4}\text{O}$ – $\text{Co}_{1.4}\text{Fe}_{1.6}\text{O}_4$  core–shell measured at 453, 300, and 10 K. The black points and the red line represent the observed and calculated patterns, respectively. The black vertical tick marks indicate the positions of the nuclear CFO (first/top), magnetic CFO (second), nuclear CO (third), and magnetic CO (fourth/bottom) diffraction peaks. The blue line represents the difference between the observed and calculated plots. (d)–(f) Temperature dependence of the FiM–A site, FiM–B site, and AFM’s magnetic moments, respectively. (g) and (h) The temperature dependence of the FiM lattice parameter, and composition plotted in terms of % cobalt on the A and B sites, respectively. Note that the difference in A and B site % cobalt values was constrained to be constant. (i) The temperature dependence of the lattice parameter of the AFM core. This phase decomposes above  $\sim 760$  K, at which point the whole sample undergoes recrystallization. (j) The lattice parameter of a new spinel phase that appears at high temperature prior to recrystallization. (k) The relative change in background is due to hydrogen-containing material being burnt off on heating.



from Rietveld refinement are summarized in Fig. 3d–k. We note that on modelling the diffraction data, it was necessary to include a degree of preferential orientation of the nanoparticles along the [110] axis. While pure CO exhibits distinct peaks characterized mostly by the hexagonal crystal structure according to XRD results (Fig. 3f), the neutron diffraction and XRD data for the core AFM phase within the CS system are consistent with the cubic crystal structure (Fig. 1f and 2a). Furthermore, the magnetic propagation vector of the core's long-range antiferromagnetic order was found to be  $k = (1/2, 1/2, 1/2)$ , consistent with previous reports of the bulk cubic CoO magnetic structure.<sup>45</sup> The lattice parameter and composition of the FiM shell, expressed in terms of the percentage of Co on the A and B sites, are shown in Fig. 2g and h. The lattice parameter varies smoothly with temperature, exhibiting a gradual expansion as the temperature increases. The composition is relatively constant, up to around 480 K, beyond which the iron-rich spinel phase transforms towards Fe<sub>3</sub>O<sub>4</sub> (we label this temperature point  $T_B$ ). We hypothesize that the cobalt ions start diffusing out of the nanoparticle beyond this point; however, the fate of diffused Co remains uncertain, as neutron diffraction only provides information from well-ordered structures. The lattice parameter of the core AFM component is shown in Fig. 3i and displays some anomalous behavior at high temperatures prior to its decomposition above  $\sim 760$  K ( $T_D$ ), at which point the CS nanoparticles recrystallize. Pre-empting recrystallization, a new crystalline phase appears above  $\sim 700$  K ( $T_C$ ), which we tentatively assign to a high-temperature spinel phase (Fig. 3j). The figure also illustrates the lattice parameter of this high-temperature spinel phase, which likely corresponds to

Co<sub>3</sub>O<sub>4</sub>. The relative change in the background due to the hydrogen-containing material (ligands) being burnt off is shown in Fig. 3k. This occurs at a temperature above  $T_A$ , which closely agrees with the thermogravimetric analysis (TGA) data (Fig. S5a, ESI†). The reduction of hydrogen-containing material from  $T_A$  to  $\sim T_B$  is attributed to the desorption of moisture, while the reduction observed beyond  $\sim T_B$  is likely due to the burning of surface ligands from the nanoparticles. The refinement data reveal a mixed cobalt/iron composition in both the AFM core and the FiM shell. At room temperature, the composition of the cations in AFM core is (Co<sub>0.6</sub>Fe<sub>0.4</sub>)O, while the FiM shell has a composition of (Co<sub>0.6</sub>Fe<sub>0.4</sub>) for the A sites and (Fe<sub>1.2</sub>Co<sub>0.8</sub>) for the B sites.

Bulk elemental analyses of the CS sample indicate excess Fe compared to the ordered Fe stoichiometry derived from neutron diffraction (Section S6, ESI†). This discrepancy arises because bulk methods detect both ordered and disordered Fe-rich phases, while neutron diffraction selectively probes the crystalline AFM-FiM CS structure. Complementary, the Fe<sub>2p</sub> spectrum (Fig. S7, ESI†) obtained using X-ray photoelectron spectroscopy (XPS) indicates the presence of Fe<sub>2</sub>O<sub>3</sub>/Fe(OH)<sub>3</sub>, which likely exists as the disordered phase, since we do not observe it in our neutron diffraction experiments.

Fig. 3d shows the refined magnetic moment at the A-site of the FiM shell, which was found to decrease gradually with temperature, reaching zero at  $T_2$  ( $\sim 770$  K). This indicates the Curie transition temperature, which agrees with the TGA results (Fig. S5b, ESI†). The FiM B-site moment also tends to zero at  $T_2$ , having undergone a slight increase observed at around 600 K (Fig. 3e), consistent with the thermally induced change in composition towards Fe<sub>3</sub>O<sub>4</sub>.

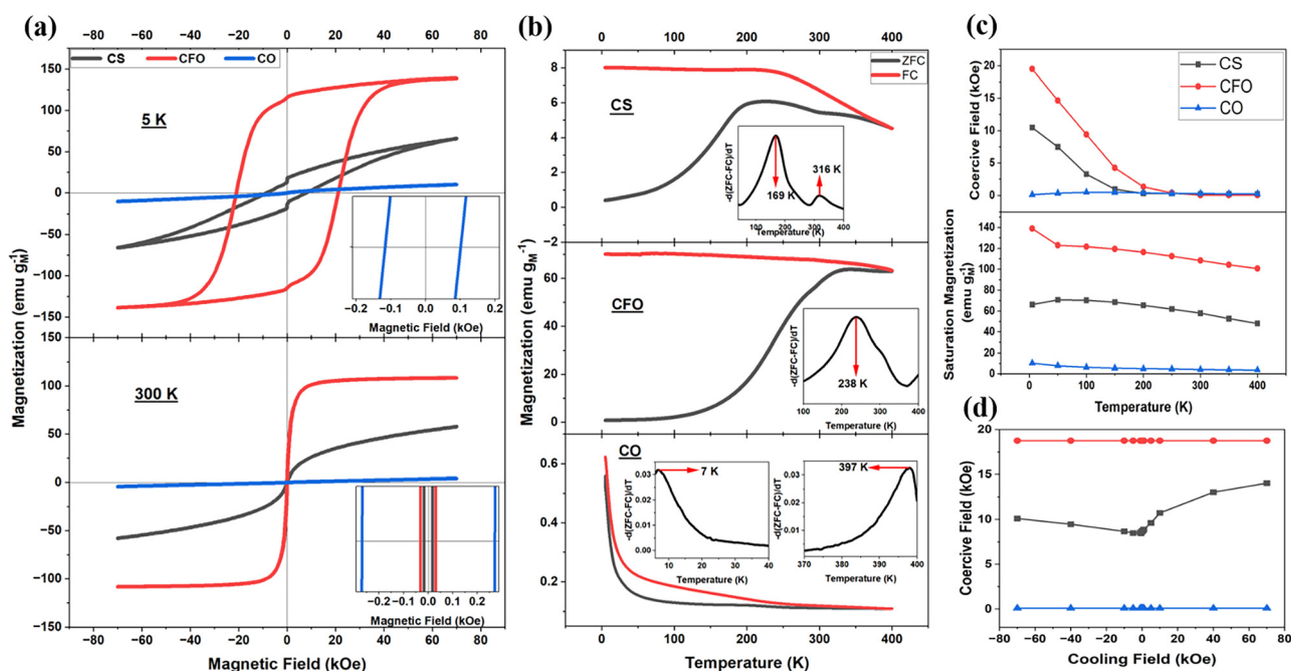


Fig. 4 (a) Magnetic hysteresis loop of CO, CFO, and CS nanoparticles at 5 K and 300 K. (b) ZFC–FC curves and their derivative plots (inset) depicting points of divergence between ZFC and FC curves. (c) The coercive field (top pane) and maximum magnetization (bottom pane) obtained at 70 kOe as a function of temperature. (d) Coercive field as a function of cooling field strength measured at 5 K.



The temperature-dependent magnetic moment of the AFM  $\text{Co}_{0.6}\text{Fe}_{0.4}\text{O}$  core is presented in Fig. 3f. The moment vanishes at  $T_1$  ( $\sim 397$  K), representing its  $T_N$ . Additionally, the zero-field cooled (ZFC–FC) graph of the pure CO nanoparticles shown in Fig. 4b shows two distinct points of divergence: the first at 7 K, supposedly the blocking temperature ( $T_b$ ), and the second around 397 K, the  $T_N$  of the material, which aligns well with the neutron diffraction data. The observed  $T_N$  in the AFM nanoparticles may originate from nanoscale effects or Fe substitution. To our knowledge, such a high  $T_N$  in similar AFM nanoparticles has not been previously reported.

The magnetic hysteresis loops of all three samples were measured at different temperatures from 5 K to 400 K (step size 50 K) to the magnetic field of  $\pm 70$  kOe (Fig. S4a, d, e, ESI†). CFO and CS display FM/FiM characteristics at 5 K and superparamagnetic behavior at 300 K (Fig. 4a). Pure CO, in general, displays a linear hysteresis at all temperatures, characteristic of an AFM material. However, the presence of weak ferromagnetism is seen as a non-linear S-shaped curve at a lower field, particularly at 5 K (Fig. S4d, ESI†). This is attributed to the disordered spins on the surface of the CO nanoparticles, which can act as a weak ferromagnet and couple to the primary AFM phase. Consequently, the pure CO nanoparticles display a weak EB effect (Fig. S4e, ESI†). Repeated hysteresis loop measurements on this sample reflect the random nature of disordered surface spins, where the magnetization and coercive field differ slightly. Interestingly, when a cooling field of 10 kOe was applied, the overall magnetization of the sample decreased (Fig. S4b, ESI†), suggesting that the cooling field aligns the surface spins antiparallel to the neighboring spins, thereby reducing the total magnetization.

The magnetic hysteresis loops for all samples also exhibit kinks at zero applied magnetic field, as shown in Fig. 4a (top pane), characteristic of so-called exchange spring behavior. In both pure CFO and the CS, the non-uniform distribution of cobalt ions (with large anisotropy energy) and iron ions (with relatively small anisotropy energy) within the lattice can lead to such behavior. These loops generally result from the superposition of individual hard and soft hysteresis loops.<sup>46</sup> For the CS nanoparticles, the interdiffusion of ions at the interface further contributes to variations in anisotropy energy across the system, resulting in pronounced exchange spring behavior.<sup>47–49</sup> The relatively weaker kink observed in the pure CO sample (Fig. S4e, ESI†) is attributed to anisotropy energy differences between disordered surface and ordered core spins.

Temperature-dependent magnetization is studied by measuring ZFC–FC curves (Fig. 4b), where the point of divergence of the ZFC and FC curves is estimated using a derivative method, as outlined in the literature.<sup>50</sup> For nanoparticles, this divergence typically represents the  $T_b$ , below which the magnetic moments of a given nanoparticle are ‘frozen’. Above the  $T_b$ , thermal energy is sufficient to cause the magnetic moments of individual nanoparticles to flip randomly, resulting in superparamagnetism. Pure CFO nanoparticles exhibit a  $T_b$  at 238 K, while biphasic CS exhibits a major peak at 169 K and a minor

peak at 316 K (Fig. 4b). Such a bimodal distribution is observed in exchange-coupled biphasic nanoparticles and can be attributed to the competing magnetic anisotropies between the distinct phases within the CS structure.<sup>51,52</sup> We also observe two divergence points for pure CO nanoparticles (Fig. 4b, bottom panel). The first is at 7 K, which most likely represents the  $T_b$ , as AFM metal oxides are known to exhibit relatively weak magnetic anisotropy energy.<sup>53,54</sup> The divergence at 397 K and the neutron diffraction data (Fig. 3f) suggest elevated  $T_N$  of CO nanoparticles, higher than the bulk value of  $\sim 293$  K. Using  $T_b$  values of 7 K, 238 K, and 169 K for CO, CFO, and CS nanoparticles, respectively, the  $K_{\text{eff}}$  is calculated (eqn (S2) and Table S1, ESI†). The  $K_{\text{eff}}$  value represents the energy required per unit volume to reorient the magnetization of a nanoparticle from the easy axis. It is affected by a combination of crystalline, shape, surface, and exchange anisotropy.<sup>55,56</sup> The CS nanoparticles exhibit the highest  $K_{\text{eff}}$  of  $17.83 \times 10^4 \text{ Jm}^{-3}$ , while pure CO shows the lowest at  $0.71 \times 10^4 \text{ Jm}^{-3}$ , and pure CFO has a similar value to the CS system ( $16.72 \times 10^4 \text{ Jm}^{-3}$ ). The comparable  $K_{\text{eff}}$  values of the CS and pure CFO nanoparticles suggests that the CS architectures retains the intrinsic high anisotropy of the FiM shell while enabling additional stabilization mechanisms, such as the exchange coupling with the AFM core. This synergy could help overcome the superparamagnetic limit in FiM nanoparticles by leveraging both enhanced anisotropy and interfacial effects.

At lower temperatures, the CS's coercivity is smaller than pure CFO's, consistent with their respective blocking temperatures. The differences are attributed to the CO phase, which has a weaker  $K_{\text{eff}}$  (Table S1, ESI†). The coercivity of CFO and CS approaches zero near their respective blocking temperatures, indicating the switch to superparamagnetism (Fig. 4c, top pane). Additionally, the coercivity of CS is influenced by both the type (positive or negative) and the strength of the applied cooling field, as shown in the figure (Fig. 4d). The maximum magnetization of all samples generally decreases with increasing temperature as the thermal fluctuations destabilize the alignment of the dipoles (Fig. 4c, bottom pane).

Fig. 5a shows the EB effect in the CS nanoparticles as positive and negative fields are applied while cooling to 5 K. The cooling field aligns the FiM spins relative to the AFM spins. When a positive magnetic field is applied, the hysteresis loop shifts to the negative ( $x$ -axis) direction and *vice-versa*. This is because, in the core–shell system, the AFM spins exert a unidirectional force on the FiM spins due to exchange interaction. This leads the FiM spins to experience an extra energy barrier when trying to flip its magnetization, causing a shift in the hysteresis loop.<sup>55</sup> The maximum EB observed is  $-10.34$  kOe at 5 K when applying a 40 kOe cooling field (Fig. 5b). The EB in the CS is highest at 5 K, gradually decreasing with increased temperature and disappearing at temperatures above 150 K (Fig. 5c). This is close to the primary derivative ZFC–FC peak observed at 169 K for the CS sample (Fig. 4b, inset of top pane graph). We attribute this peak to the CS system's primary  $T_b$ . As the temperature reaches  $T_b$ , thermal energy is large enough to flip the magnetization of the FiM spins randomly, and the sample as a whole no longer



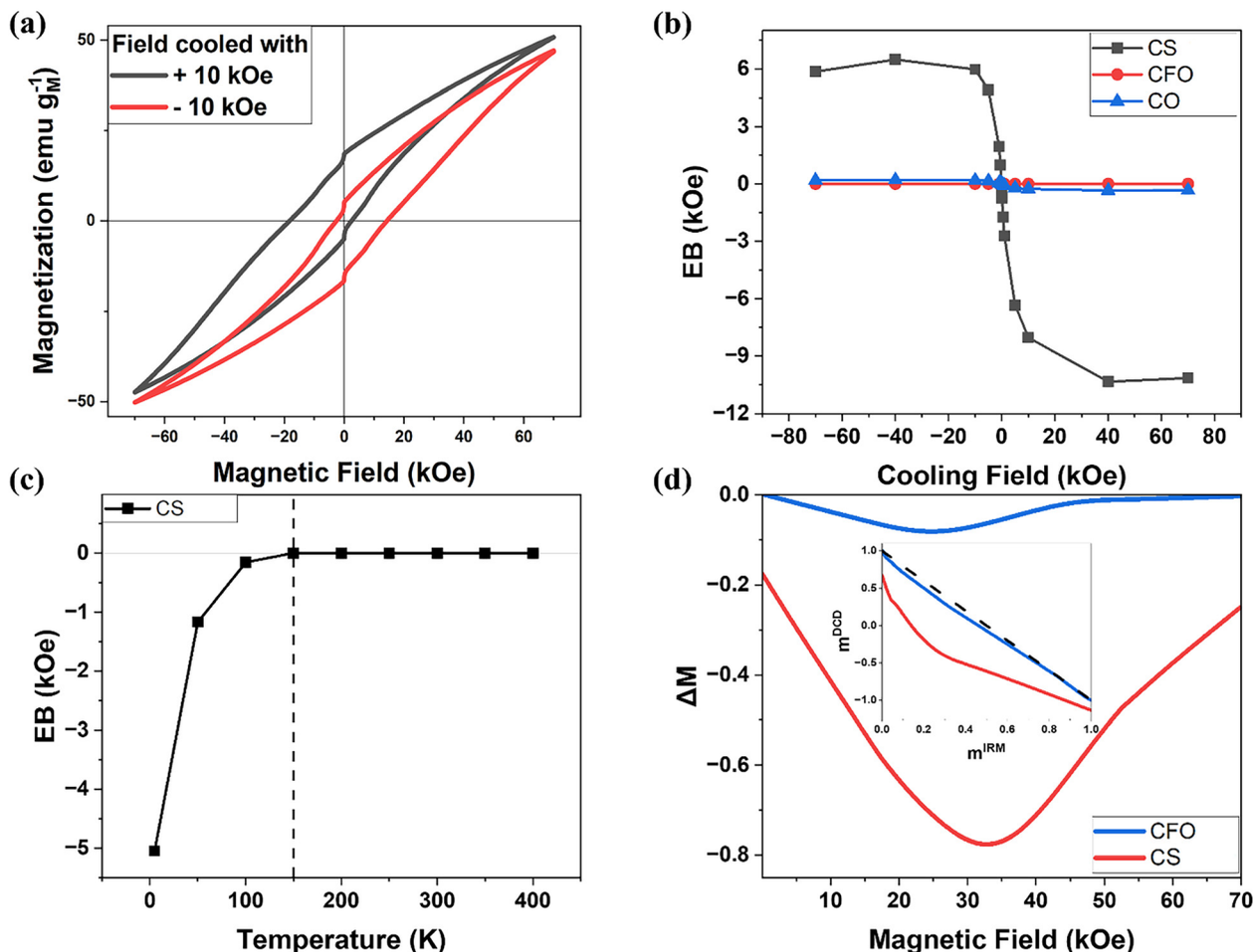


Fig. 5 (a) Shift in magnetic hysteresis loop at 5 K observed upon cooling biphasic CS sample with +10 kOe and -10 kOe. EB effect as a function of (b) cooling field strength at 5 K and (c) temperature for the CS nanoparticles. (d) Henkel plots of pure CFO and biphasic CS measured at 5 K.

displays EB. Nevertheless, the EB may still be present above  $T_b$ , but magnetometry might not capture its subtle effects due to increased thermal fluctuations at high temperatures. Since EB allows precise control over the magnetic switching behavior of nanoparticles, it can enhance their responsiveness to external magnetic fields, making it particularly useful for applications like magnetic hyperthermia in nanomedicine.

Interparticle (dipolar) interactions in pure CFO and the CS nanoparticles are studied at 5 K using Henkel plots, which are generated from a series of isothermal remanent magnetization (IRM) and direct current demagnetization (DCD) measurements (Fig. 5d). These measurements are based on the Stoner-Wohlfarth model, which describes a system of non-interacting, single-domain particles.<sup>57</sup> Henkel plots are obtained by comparing the IRM curve  $M_r(H)$  with the DCD curve  $M_d(H)$ , as outlined in the literature.<sup>58,59</sup> For a system of non-interacting magnetic particles, the reduced IRM and DCD curve is a straight line with a slope of -0.5, following the Wohlfarth relationship:

$$m_d(H) = 1 - 2m_r(H)$$

$$\text{where, } m_d(H) = \frac{M_d(H)}{M_d(H_{\max})} \text{ and } m_r(H) = \frac{M_r(H)}{M_r(H_{\max})}.$$

In systems with interacting magnetic nanoparticles, the Henkel plot departs from the linear graph in  $\Delta M$  vs.  $H$  plot. The  $\Delta M$  of the system is calculated using the following formula:<sup>57-59</sup>

$$\Delta M(H) = m_d(H) - [1 - 2m_r(H)]$$

Here,  $\Delta M < 0$  indicates demagnetizing or inhibiting dipolar interactions, while  $\Delta M > 0$  indicates magnetizing or enhancing interactions. The magnitude of  $\Delta M$  depends on the degree of the magnetic interactions.<sup>57-59</sup> Pure CFO and CS exhibit demagnetizing interactions, as indicated by the deviation from the linear graph (dashed line) in the inset of Fig. 4d. The magnitude of the dipolar interaction is greater in the CS sample and is observed at all magnetic field strengths. For pure CFO nanoparticles, the effect of dipolar interaction is only visible up to  $\sim 45$  kOe, above which it disappears. The stabilizing ligands may act as spacers and influence dipolar interaction during the measurement.<sup>60</sup> TGA measurements show that the amount of stabilizing ligands attached to CFO nanoparticles is higher (12.6%) than the CS (8.8%), which may lead to a higher degree of aggregation in the CS (Fig. S5a and c, ESI<sup>†</sup>). Despite efforts to synthesize similar nanoparticles for comparison, slight



differences in the final products could explain the varied Henkel results. Additionally, the disappearance of dipolar interaction effects in CFO at 45 kOe aligns with the magnetic field at which the hysteresis loop saturates. In contrast, the dipolar interaction effects in the CS do not disappear even at the maximum applied field of 70 kOe. Our results indicate a stronger dipolar interaction in biphasic CS nanoparticles than pure CFO. Complex spin interactions between the two phases in core-shell nanoparticles and dipolar interactions can impact the magnetic properties, including the anisotropy energy barriers of individual nanoparticles.<sup>61</sup> The evaluation of such interactions and their impact on the observed EB is outside the scope of this study. The small size of the nanoparticles also challenges the analysis of the experimental results in this study, introducing various uncertainties.

## Conclusion

We have synthesized colloidally stable CS  $\text{Co}_{0.6}\text{Fe}_{0.4}\text{O}-\text{Co}_{1.4}\text{Fe}_{1.6}\text{O}_4$  nanoparticles with a remarkable EB of 10.34 kOe. The synthesis was performed using a one-step thermal decomposition of metal complexes. We also synthesized pure CFO and CO nanoparticles for comparison. The colossal EB observed in the CS nanoparticles is attributed to the highly crystalline AFM core and a lattice-matched interface observed using atomic resolution STEM. Neutron diffraction reveals structural changes of the AFM core in the CS at high temperatures, leading to its recrystallization into a new phase, most likely  $\text{Co}_3\text{O}_4$ . Additionally, because of its core-shell structure, we observe the interdiffusion of iron ions into the AFM lattice. The  $K_{\text{eff}}$  of the CS system is significantly larger than pure CO nanoparticles. As a result, the EB is observed from 5–150 K, close to the blocking temperature of the CS nanoparticles. Above 150 K, superparamagnetic behavior dominates, with vanishing hysteresis. ZFC-FC measurement of the pure CO nanoparticles reveals a divergence at  $\sim 397$  K, which we attribute to its  $T_{\text{N}}$ . This is supported by neutron diffraction measurements conducted on the CS system, where its AFM core ( $\text{Co}_{0.6}\text{Fe}_{0.4}\text{O}$ ) also exhibits  $T_{\text{N}}$  at  $\sim 397$  K. The robust AFM stability at such elevated temperatures could hold significant promise for high-temperature applications. We also observed stronger demagnetizing (inhibiting) dipolar interactions in the CS than in pure CFO nanoparticles, which may affect the observed EB. We envision that this study will pave the way for engineering biphasic CS nanoparticles to overcome the superparamagnetic limit for smaller particle sizes and obtain room-temperature EB by tuning the magnetic anisotropy energy and controlling the magnetic switching behavior. Such systems could advance applications in nanospintronics (e.g., high-density storage) and nanomedicine (e.g., magnetic hyperthermia), while motivating further exploration of doped AFM nanoparticles and their interplay with magnetic properties.

## Experimental

### Materials

Cobalt(II) acetylacetonate,  $\geq 99\%$ ; iron(III) acetylacetonate,  $\geq 99\%$ ; 1,2-hexadecandiol, 90%; dioctyl ether, 99%; oleylamine,

70%; oleic acid, 90%; octadecene, 99%; *n*-hexane,  $\geq 99\%$  and ethanol absolute,  $\geq 99\%$ , were purchased from Sigma-Aldrich and VWR. All chemicals were used without further purification.

### Synthesis of pure CO nanoparticles

The synthesis of all nanoparticles is conducted in a Schlenk line under a nitrogen atmosphere. In short, cobalt(II) acetylacetonate (0.324 g, 1.26 mmol), oleylamine (1.714 ml, 5.20 mmol), 1,2-hexadecandiol (0.388 g, 1.5 mmol), and octadecene (10 ml) are added to a 100 ml three-neck round bottom flask. The flask is then placed in the Schlenk setup and degassed for 30 min at room temperature while stirring with a magnetic stir bar. Nitrogen is purged at a flow rate of  $300 \text{ ml min}^{-1}$ , and the speed of the magnet bar is maintained at 1000 rpm.

Following the initial degassing step, the temperature is raised from room temperature with a ramping rate of  $6 \text{ }^\circ\text{C min}^{-1}$  to  $100 \text{ }^\circ\text{C}$  and then held for 30 min under continued degassing. The temperature is further increased to  $200 \text{ }^\circ\text{C}$  with the same ramping rate and then held for 30 min. Finally, the temperature is ramped to  $295 \text{ }^\circ\text{C}$ , and the reaction was carried out for another 60 min.

At the end of the reaction, the heating mantle is removed, and the solution is cooled naturally to room temperature, which takes approximately 15 min. The resulting product is a black solution divided equally into two 50 ml centrifuge tubes. To each tube, 10 ml of absolute ethanol is added, and the mixture is centrifuged at 9500 rpm for 20 min. The supernatant is discarded, and 5 ml of *n*-hexane is added to each centrifuge tube. The precipitate is quickly redispersed by sonication for 1 min. Subsequently, 10 mL of absolute ethanol is added, and the mixture is vortexed and centrifuged under the same conditions. This washing procedure is repeated once more, after which the precipitate is dried overnight at room temperature for further characterization.

### Synthesis of pure CFO nanoparticles

For the synthesis of pure CFO nanoparticles, cobalt(II) acetylacetonate (0.108 g, 0.42 mmol), iron(III) acetylacetonate (0.297 g, 0.84 mmol), 1,2-hexadecandiol (0.388 g, 1.5 mmol), oleic acid (0.866 ml, 2.6 mmol), oleylamine (0.816 ml, 2.6 mmol) and octadecene (10 ml) is added to a 100 ml three-neck round bottom flask. The reaction and washing processes are carried out following the same protocol as described in the previous section.

### Synthesis of biphasic CS nanoparticles

For the synthesis of core-shell (CS) nanoparticles cobalt(II) acetylacetonate (0.108 g, 0.42 mmol), iron(III) acetylacetonate (0.297 g, 0.84 mmol), 1,2-hexadecandiol (0.388 g, 1.50 mmol), oleylamine (1.714 ml, 5.20 mmol) and dioctyl ether (10 ml) are added to a 100 ml three-neck round bottom flask. The reaction and washing steps follow the protocol described in the previous section.

### Characterization

Characterizations are performed in powder or liquid form, where the nanoparticles are dispersed in *n*-hexane. For



characterizations requiring liquid samples, the solution was mixed vigorously with a vortexer for 1 min and sonicated in an ultrasonicator for 5 min.

The sample's morphology is characterized using a TEM (JOEL EM- BEI10). For this, the liquid is drop-cast onto a carbon-coated copper grid and dried for 3 h to allow the evaporation of *n*-hexane. Double aberration corrected atomic resolution TEM/STEM (JEOL NeoArm 200F) is used for atomic columns imaging and SAED of the core-shell structure. JEOL 2100F was used for energy-dispersive X-ray Spectroscopy (EDX). The solvodynamic size of the nanoparticles is measured using DLS (Malvern Instruments Zetasizer Nano). The optical properties are characterized using UV-VIS (Molecular Devices Max Pro 5). The liquid sample was placed in a quartz cuvette for DLS and UV-VIS measurements. Powder samples for inductively coupled plasma optical emission spectroscopy (ICP-OES) were dissolved in 70% nitric acid and diluted to 5% for measurement (Varian 720).

Structural characterization is performed on the powder sample using XRD (PANalytical X'Pert Pro), with cobalt radiation at a wavelength of 1.79 Å. XPS measurements were collected using a thermo Fischer scientific K-Alpha+ photoelectron spectrometer. Thermal analysis is conducted using TGA (TA Instruments Discovery Series). Surface functional groups are characterized using FTIR (Perkin Elmer Spectrum 65). The atomic and magnetic structure of the nanoparticles is studied using time-of-flight neutron diffraction (ISIS Neutron and Muon Source). Finally, DC magnetic measurements are performed with a superconducting quantum interference device-vibrating sample magnetometer, SQUID-VSM (Quantum Design MPMS 3).

## Author contributions

S. A., R. D. J., and N. T. K. T. conceived and designed the study. S. A. conducted the experiments, including nanoparticle synthesis, XRD, TEM, EDX, UV-Vis, DLS, FTIR, ICP-OES, SQUID measurements, and data analysis. L. D. T. and S. A. performed TGA. P. M., R. D. J., and S. A. conducted neutron powder diffraction. L. L. and V. K. L. carried out SAED and atomic-resolution STEM. R. D. J. analyzed the neutron powder diffraction data. S. A. wrote the manuscript with input from R. D. J., V. K. L. and N. T. K. T. All authors reviewed the manuscript, provided critical feedback, and approved the final version.

## Conflicts of interest

There are no conflicts to declare.

## Data availability

The data supporting the findings of this study are available within the article and ESI.†

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