Core/Shell Nanoparticles of Non-Stoichiometric Zn–Mn and Zn–Co Ferrites as Thermosensitive Heat Sources for Magnetic Fluid Hyperthermia

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Supporting Information

1. INTRODUCTION

Hyperthermia is a powerful and well-known oncological therapy, aiming to heat cancer cells up to 41–45 °C to trigger cytotoxic effects with minimum harm on the surrounding tissues. Local heating of tumorous cells can be achieved by using magnetic nanoparticles (NPs) as heat sources in magnetic fluid hyperthermia (MFH) given their ability to convert magnetic energy into heat when submitted to a time-varying magnetic field. In addition, since magnetic NPs provide contrast enhancement in magnetic resonance imaging (MRI), they are being explored as dual-purpose systems for therapy (MFH) and diagnostics (MRI) in a theranostic approach.

The heat produced by ferrimagnetic single-domain NPs under a time-varying magnetic field is related to dynamic hysteresis losses produced by the relaxation of the magnetic moment. The phenomenology of MFH arises from two mechanisms associated with the relaxation of the NPs’ magnetic moment. When dispersed in a liquid carrier, an isolated magnetic NP can rotate, and the associated Brownian relaxation time is \( \tau_B = 3\eta V_H/k_B T \), \( \eta \) being the viscosity of the medium, \( V_H \) the NP hydrodynamic volume, and \( k_B T \) the thermal energy, with \( k_B \) being the Boltzmann constant. The magnetic moment would also relax due to thermally induced fluctuations by overcoming an energy barrier of magnetic anisotropy (\( E_A \)), a phenomenon well-characterized by a Néel relaxation time \( \tau_N = \tau_0 \exp(E_A/k_B T) \), \( \tau_0 \) being the spin relaxation time on the order of \( 10^{-9} \) s (for a detailed review of Néel relaxation time expression, see, for example, ref 8). Assuming these two mechanisms are independent, the effective relaxation time is given by \( \tau^{-1} = \tau_N^{-1} + \tau_B^{-1} \), and the dominant mechanism for the heating generation will be the faster one.

The power absorbed by the NPs is commonly described by the linear response theory (LRT) when the applied field \( H_0 \) is much smaller than the anisotropy field \( H_A \) of the NPs and the dynamic magnetization of the system varies linearly with \( H_0 \). Within this approximation, the heating rates of the NPs present an optimum particle size, the values of which depend strongly on the magnetic anisotropy.

ABSTRACT: We report on the suitability of core/shell nanoparticles (NPs) for magnetic fluid hyperthermia in a self-regulated and theranostic approach. Aqueous magnetic colloids based on core/shell \( n\text{Zn,Mn,Fe}_2\text{O}_4@\gamma\text{-Fe}_2\text{O}_3 \) and \( \text{Zn,Co}_2\text{Fe}_2\text{O}_4@\gamma\text{-Fe}_2\text{O}_3 \) NPs were produced by a three-step chemical synthesis. Systematic deviations from stoichiometry were observed with increasing Zn substitution for both series of samples. We investigated how the chemical composition affects the saturation magnetization, magnetic anisotropy, and thermomagnetic properties of these core/shell NPs. The heating efficiency through specific power absorption (SPA) was analyzed in the framework of the linear response theory. SPA values obtained for NPs present a different contrast of anisotropy between the core and shell materials, indicating no evidence of the contribution of enhanced exchange coupling to the heating efficiency.

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efficiency has been experimentally investigated as a function of several parameters, such as the particle size, saturation magnetization, morphology, and magnetic anisotropy. Moreover, the efficiency of NPs with varying anisotropies and diameters in media of different viscosities has been explored. More recently, the role of magnetic dipolar interaction and NPs aggregation in power loss efficiency has emerged as a controversial issue, and further studies are needed to better understand it.

Since the heating efficiency in MFH is strongly dependent on individual properties of NPs like size, magnetization, and anisotropy, a combination of different materials in a unique core/shell NP would be an interesting strategy to improve the MFH performance. In the last years, few studies pointed out that the design of core/shell nanoparticles with a high contrast of anisotropy between shell and core provides a remarkable enhancement of the power loss efficiency. It has been proposed that this observed increase of the specific power absorption (SPA) would be related to the interfacial exchange coupling between magnetically soft and hard phases. Very recently, the power absorption of such systems has been calculated by using a theoretical approach based on the LRT and considering the influence of both core and shell contributions, as well as the exchange coupling at the interface. It is found that increasing the interface exchange constant would be an interesting strategy to achieve better efficiency with small core/shell soft/hard NPs.

We propose here to investigate the heating performance of core/shell NPs based on hard/soft and soft/soft magnetic ferrite phases. Indeed, in a previous work, an exchange bias field in ultrasmall (~3 nm) core/shell MnFe₃O₄@γ-Fe₂O₃ and CoFe₂O₄@γ-Fe₂O₃ NPs has been measured. It is found that the magnitude of this exchange coupling field is larger for core/shell NPs based on a harder ferrite core. Following ref 44, this result would indicate that they could be more efficient for the conversion of electromagnetic energy into heat. In this paper, we also choose to modulate the thermomagnetic response by doping the NPs core with diamagnetic Zn ions, since larger zinc content provides the weakening of the superexchange interactions. It therefore leads to an increasing thermomagnetic coefficient $f_{TM} = -\Delta M/\Delta T$, where $\Delta M$ is the magnetization per unit mass and decreasing Curie temperature, hence limiting the upper temperature reached by MFH to the safety range (41−45 °C). This characteristic would be very promising because it could enable a local self-regulated temperature control, which would avoid damage to healthy tissues due to overheating.

In such a context, this paper reports on the MFH performances of two series of electric double-layered magnetic fluids (EDL-MF) based on core/shell Zn,Mn,Fe₃O₄@γ-Fe₂O₃ and Zn,Co,Fe₂O₄@γ-Fe₂O₃ NPs, with different Zn/Mn and Zn/Co proportions. These colloidal NPs are dispersed in aqueous medium by using a well-controlled synthesis route and present a long-term thermodynamic stability thanks to the introduction of adjustable Coulomb’s repulsive forces between them. Intrinsic magnetic properties like magnetization and coercive field are explored in detail and allow one to correlate them with the heating efficiency. The strong anisotropic character of Co cations when compared to Zn and Mn ones allows one to tune the magnetic anisotropy constant and the contrast of anisotropy between the core and shell materials. Finally, we discuss these results in light of interfacial exchange coupling between core and shell phases.

2. MATERIALS AND METHODS

2.1. Synthesis of Nanoparticles and Magnetic Colloids. Magnetic colloids are synthesized in three steps, as described in more detail elsewhere. In brief, Zn–Mn and Zn–Co ferrite nanoparticles are prepared by hydrothermal coprecipitation of aqueous solutions of Fe³⁺, Zn²⁺, and Mn²⁺ or Co²⁺ (0.5 mol/L) in strong alkaline medium (2 mol/L NaOH) at 100 °C under vigorous stirring. Aiming for the obtention of a stoichiometric mixed ferrite core (Zn,Mn₁₋ₓFe₂O₄ M being Mn or Co), the metallic salt mixtures Zn:M:Fe are tuned as $x = 1 + x$ (with $x$ varying between 0.1 and 0.9). Next, the as-prepared precipitate is cleaned and acidified with 2 mol/L HNO₃ to reverse the surface charge of the NPs and eliminate undesirable byproduct. At this stage, the release of the metallic cations from the surface is more significant for the divalent metals when compared to Fe³⁺, indicating the beginning of the iron-rich shell formation onto the NPs surface. As the chemical stability of the NPs in acidic medium is achieved by incorporating more iron atoms onto the surface, the precipitate is hydrothermally treated with a 0.5 mol/L Fe(NO₃)₃ solution at 100 °C for 15 min. Finally, the particles are peptized by pH and ionic strength adjustment in the aqueous medium, to control the surface charge density and the screening of the electrostatic repulsion between particles to ensure colloidal stability. The NPs of varying composition are indexed as ZC1−ZC9 for core/shell NPs based on Zn–Co ferrite cores and ZM1−ZM9 for those obtained with Zn–Mn ferrite ones.

2.2. Chemical, Structural, and Morphological Characterization. The chemical composition of NPs is checked by determination of Zn, Mn, Co, and Fe concentrations with energy-dispersive X-ray spectroscopy (EDX) and atomic absorption spectroscopy (AAS) techniques. Both measurements are performed at Universidade de Brasília (UnB) with a Shimadzu EDX 720HS and with a Thermo Scientific Spectrometer model S series AA, respectively. For AAS measurements, specific lines are chosen for each metal—Fe (248.3 nm/372.0 nm), Zn (213.9 nm), Mn (279.5 nm), and Co (240.7 nm)—to avoid interference effects.

The crystalline structure of NPs is investigated by X-ray powder diffraction (XRPD) after evaporation of the liquid phase from the samples. For samples based on Zn–Mn ferrites, measurements are carried out at Laboratório Nacional de Luz Síncrotron (LNLS) using the D12A-XRD1 beamline and a X-ray wavelength $\lambda = 1.771 55$ Å in the 2θ range 15°−120°. For Zn–Co ferrite-based samples, diffractograms are collected using a D8 Focus (Bruker) diffractometer at UnB, using Cu Kα radiation with $\lambda = 1.5406$ Å in the 2θ range 20°−80°. The diffracted lines are related to the characteristic interplanar spacings of the structure and compared with bulk standard data of the International Centre for Diffraction Data (ICDD) for MnFe₂O₄ (ICDD no. 00-073-1964), ZnFe₂O₄ (ICDD no. 98-007-2033), CoFe₂O₄ (ICDD no. 00-022-1086), Zn,Mn₁₋ₓFe₂O₄ (ICDD no. 98-002-8512 to -8516), Zn,Co₁₋ₓFe₂O₄ (ICDD no. 98-016-6201 to -6204), γ-Fe₂O₃ (ICDD no. 98-008-7119), and Fe₂O₄ (ICDD no. 98-002-6410). Crystalline sizes are obtained using Scherrer’s formula $D_{XRD} = 0.9\lambda/\beta \cos \theta$ applied to the most intense peak [311], $\beta$ being its width at half-maximum (the intrinsic width is discounted using a Si standard).

For all samples, morphology, crystallinity, and size distributions of NPs are investigated by transmission electron microscopy (TEM; JEOL JEM100 CX II) and high-resolution
TEM (HRTEM; JEOL JEM2010), both with a LaB6 filament operated at 200 kV, carried out at Université Pierre et Marie Curie (UPMC). Size distribution histograms are obtained by counting approximately 600 particles and are fitted by a log-normal distribution $P(D) = \exp\left(-\ln^2(D/D_0)/2s^2\right)/\sqrt{2\pi} D_0 s$ and $s$ being the characteristic diameter and the polydispersity index, respectively. Moreover, the interplanar spacings associated with the observed reticular planes of the spinel cubic structure can be determined by fast Fourier transform (FFT) analysis and compared to ICDD data. In order to go further in detail on the NPs’ core/shell structure, sample ZM5 is probed by HRTEM using a TECNAI F30 (FEI Co.) microscope equipped with a field emission gun (FEG) operated at 300 kV at Laboratorio de Microscopias Avanzadas (LMA), University of Zaragoza (UZ). Z-Contrast imaging is obtained using scanning transmission electron microscopy (STEM) mode with a high-angle annular dark field (HAADF) and with an Oxford ultrathin window EDX detector to test the chemical composition profile.

2.3. Magnetic Measurements. Magnetization measurements are performed at UnB (Instituto de Física) and at UPMC (MONARIS Laboratory) using the same model of vibrating sample magnetometer (VSM; PPMS, Quantum Design model 6000). Magnetization curves are obtained as a function of the applied field (up to 7160 kA/m), at 5 K after a zero-field-cooling (ZFC) procedure and at 300 K for dilute magnetic fluids (MF) samples (~10 mg/mL). In this regime of negligible magnetic dipolar interactions and at room temperature, the reversible curves are analyzed using Langevin contributions of spheroidal particles weighted by a log-normal distribution of NPs sizes. Magnetic colloids can be considered as a gas of noninteracting particles that presents characteristic giant paramagnetic behavior, so that the magnetization can be expressed by

$$\sigma / \sigma_s = \frac{M}{M_s} = \frac{\int_0^\infty dV \xi V \mu V H / k_B T}{\int_0^\infty dP(D) dD}$$

(1)

where $\sigma$ is the magnetization per unit mass [$\sigma = M/\rho_{NP}$, $M$ being the magnetization per unit of volume and $\rho_{NP}$ the NPs’ density (see Supporting Information (SI))], $\sigma_s$ is the saturation magnetization per unit mass [$\sigma_s = M_s/\rho_{NP}$], $M_s$ is the saturation magnetization of the NP of moment $\mu = M_s V$, $\xi = \mu_0 u H / k_B T$ is the Langevin parameter (with $\mu_0$ being the permeability constant), and $L_q = \coth \xi - 1/\xi$ is the first Langevin function. In the high-field limit $\mu_0 u H / k_B T \gg 1$ the magnetization follows a linear dependence on $H^{-1}$, thus allowing the determination of $\sigma_s$ by performing a linear extrapolation of $H^{-1}$ in the high magnetic field region.

From hysteresis loops of ZFC MFs, obtained at 5 K, we determine the high-field magnetization at this temperature and the coercive field $H_C$ (5 K). The effective magnetic anisotropy constants ($K_{eff} = \mu_0 M H / 2$) are associated with $E_s = K_{eff} V$, $H_C$ being the anisotropy field, here deduced by assimilating $H_C$ (5 K) to $H_{C1}$, its maximum value (to be reached at $E_s/k_B T \gg 1$ with this ensemble of randomly oriented NPs). It is then possible to compare altogether the respective magnetic hardnesses of the various NPs.

Finally, for both Zn−Mn− and Zn−Co-based ferrites samples, we measure the thermal dependence of the magnetization of pressed powder samples (~2 ton) obtained at 80 kA/m, above room temperature up to 600/700 K using an oven coupled to the VSM at UnB. The Curie temperature is obtained by the extrapolation in temperature of the magnetization $\sigma \to 0$. The thermomagnetic coefficient ($k_T = -\Delta \sigma / \Delta T$) is determined at 80 kA/m in the temperature range of interest for hyperthermia applications (300−350 K).

2.4. Specific Power Absorption Measurements. Power absorption experiments are taken at INA-UZ using a commercial device (DM100 from nB nanoScale Biomagnetics) to determine the SPA of ferrofluid samples. In adiabatic systems, the amount of heat released by the NPs (power losses) is equal to the power absorbed and is given by the area enclosed by the ac dynamic hysteresis loop. Thus, the SPA (W g$^{-1}$) index is defined as $SPA = P / c_{NP}$ with $P$ (W m$^{-3}$) being the volumetric power loss and $c_{NP}$ the concentration of magnetic NPs within the solution in mg/mL. It is experimentally determined by

$$SPA = c_{solvent} m_{solvent} \frac{\Delta T}{\Delta t}$$

(2)

where $c_{solvent}$ is the specific heat of the solvent, $m_{solvent}$ and $m_{NP}$ are respectively the mass of the solvent and the NPs, and $\Delta T / \Delta t$ is the initial slope of the experimental heating curve. For all the samples, measurements are carried out at a NP concentration of around 10 mg/mL (aiming to minimize changes in collective behavior due to dipolar interactions between NPs) using an ac magnetic field amplitude of 12 kA/m at a frequency of 265 kHz. Data are also collected for frequencies between 265 and 831 kHz at 12 kA/m for MF based on ZC1 NPs and by varying the field amplitudes in the range of 4−24 kA/m for the MF samples based on ZM2, ZM3, ZM6, and ZM9 NPs.

3. RESULTS

3.1. Core/Shell Nanoparticles. We synthesized mixed ferrite NPs with a core/shell structure portrayed as Zn$_x$M$_{1-x}$Fe$_2$O$_4$@γ-Fe$_2$O$_3$ ($M$ being Mn or Co), to disperse them in acidic aqueous medium ($pH \sim 2$). The NPs’ surface modifications induced by the surface treatment and the formation of the core/shell structure are well followed by EDX and AAS measurements of metal concentrations at each step of the synthesis (refs 58, 59). Table SI (SI) collects the chemical characteristics of both bare NPs and surface-treated ones. The results show that, after the coprecipitation step, the stoichiometry of bare NPs does not correspond to that of an ideal ferrite. For each type of sample, the fractions of $(Zn + Mn)$ and $(Zn + Co)$ decrease with increasing Zn substitution, from about 0.33, the exact value of an ideal ferrite, obtained at low Zn content, to much smaller values. Thus, we write the chemical formula as $Zn_xM_{1-x}Fe_{2}O_{4+\delta}$ to obtain the stoichiometry of bare NPs. An excess of positive charges coming from two mechanisms exactly balances the extraelectronnegative charge appearing per formula unit. The first one occurs in both kinds of samples and is related to the lower zinc content and to the larger iron content when compared to the expected nominal values. Thus, the substitution of a divalent ion by a trivalent one leads to an excess of positive charges. Moreover, in Mn-based NPs, mixed valence states of Mn ions also have to be considered, as oxidation of Mn ions would occur during the synthesis in alkaline medium. Very recently, we studied in detail the local structure of core/shell NPs based on pure Mn-ferrite core elaborated by the same synthesis procedure. The oxidation state of Mn ions has been
investigated by using X-ray absorption spectroscopy. It is found to be 2.9 in bare NPs, a value that we will use here. More details on the stoichiometry calculations are given in the SI. Table S1 lists the values of the oxidation degree \( \delta \) and the chemical formula obtained as described and renormalized in order to correspond to a close-packed arrangement of 32 oxygen anions. It therefore enhances the existence of metal ion vacancies randomly distributed in both tetrahedral and octahedral interstitial sites. A detailed analysis of structural studies of our mixed ferrite-based NPs is in progress and will be published elsewhere.

After the surface treatment, the fractions of (Zn + Mn) and (Zn + Co) also decrease. This is a direct consequence of both the removal of divalent metal cations from the surface of NPs during the acid cleaning and the incorporation of more iron ions into the surface layer during the surface treatment with ferric nitrate.\(^{59}\) This effect is more pronounced for Zn–Mn ferrite NPs, mainly due to their higher surface/volume ratio as the NP size becomes smaller for larger Zn content. The fractions of maghemite shell deduced as shown in the SI from the metal concentrations determined after the surface treatment are collected in Table S1 (SI).

XRPD patterns of representative powder samples, all exhibiting characteristic peaks of cubic spinel structure, are seen in Figure S1a,b. Samples’ characteristics, such as size, lattice parameter, and density, obtained from these diffractograms, as well as information about the procedure, are also found in the SI. The cubic cell size varies with Zn content for both sets of samples. They are compared to the lattice parameters calculated using Vegard’s law. The existence of a maghemite surface layer as well as the iron enrichment at the NP’s core induce an overall decrease of the lattice parameters (see Figure S2, SI). Both Zn–Co and Zn–Mn ferrite nanoparticles present a decrease of the crystalline size as the Zn fraction increases, and this effect is more pronounced for Zn–Mn ferrites (see Table S2, SI), in good agreement with previous reports.\(^{49,50,53,65,66}\) This indicates that the nucleation process would be favored in preference to the crystalline growth during the coprecipitation.\(^{57,66}\) Moreover, the local structure must be considered, more particularly in Zn–Mn-based samples. As an example, interatomic distances Mn–O are larger than their Zn–O counterparts and should lead to a reduction of the cubic cell,\(^{69}\) therefore inducing smaller NPs sizes. The oxidation of Mn ions and the location of the metallic cations at interstitial sites may also contribute to a decrease of both cell size and NPs diameter.\(^{64,70}\)

Conventional TEM images (see Figure 1a,b) show well-crystalline, “rocklike”-shaped NPs (see the HRTEM in Figure 1a1,b1). The corresponding size histograms are shown in Figure 1a2,b2. Results are listed in Table S2 (SI). As already observed for crystalline XRPD sizes, both Zn–Co ferrites and Zn–Mn ferrites present a downward change of characteristic diameter with increasing Zn substitution, although it is more intense for Zn–Mn ferrites.

Moreover, the core/shell nature of our NPs is clearly verified in sample ZM5 by using FEG-HRTEM measurements (Figure 1c), for which the larger contrast allowed us to observe regions of the same nanocrystal with different intensities, pointing toward the existence of a thin and uneven shell. We further investigate this issue using a STEM-HAADF image of the same NP (Figure 1c1). The contrast between the regions of the sample provides information about the morpho-chemical composition. Then, the chemical composition profile is obtained for this representative NP and it is depicted in Figure 1c2,c3. We delimit the surface layer in the presented graphs in order to clarify the analysis. Figure 1c2 show high concentrations of Fe and almost zero for Zn and Mn in the shaded area. It also indicates that the transition at the interface with the core is rather smooth. In Figure 1c3, we present the ratio between the sum of the intensities relative to Zn and Mn and that relative to the total metals. It is expected to be 0.33 for a stoichiometric Zn–Mn ferrite. In the demarked core region, this ratio is constant around 0.3, in agreement with the value 0.28 predicted by the chemical analysis presented in Table S1 (SI). In both shell regions, we observe a decrease of this value, clearly indicating the iron-enriched surface. We have also verified the core/shell chemical structure for ZC3 sample, as depicted in Figure S3 (SI). Even though the surface layer is not homogeneous and presents some remaining content of core divalent metals in the composition, the obtention of the core–shell structure by hydrothermal soft chemistry is remarkable. Indeed, our chemical core/shell approach well accounts for several structural and magnetic properties, such as coordination of core and shell metal ions and intrinsic NPs’ magnetization.\(^{63,72,73}\)

### 3.2. Magnetic Characterization of NPs at 300 and 5 K

Parts a and b of Figure 2 show typical room temperature magnetization curves obtained for dilute magnetic fluids based on ZM9 and ZC8 NPs. At room temperature, because of the rotational degrees of freedom of their NPs in the fluid carrier, the magnetic fluid samples present a giant paramagnetism behavior. Their \( M(H) \) curves thus show no coercive field. In main Figure 2a (respectively Figure 2b), the solid line represents the best magnetization fit calculated using eq 1.

![Figure 1](image-url)
From the linear fit at high field (insets of Figure 2a,b), we deduce the saturation magnetization values $\sigma_s$ (300 K) given in Table S3 (SI). They are presented as a function of the zinc content in Figure 3a,b.

Figure 3. (a and b) Saturation magnetization of NPs in dilute regime at 300 K (open symbols) and 5 K (filled symbols) for Zn–Mn and Zn–Co samples, respectively. (c) Coercive field $H_c$ obtained at 5 K as a function of Zn substitution, for Zn–Co and Zn–Mn series, both in dilute regime. (Error bars in $H_c$ are not represented, since they are about 1.6 kA/m.)

At room temperature, these same NPs, mechanically blocked in disorder in pressed powder samples, all present a small coercive field ($H_c < 9$ kA/m); see Figure S5 (SI).

At very low temperature, in ZFC MF, the magnetic NPs are also mechanically blocked and the magnetization process leads to a hysteresis loop well-described by a Stoner–Wohlfarth formalism for ferromagnetic NPs.74 Figure 2c,d shows typical magnetization hysteresis loops recorded at 5 K with ZFC-MF samples based on ZM2, ZM9, ZC1, and ZC9 NPs. The characteristic saturation magnetization $\sigma_s$ (5 K) and the coercive field $H_c$ (5 K) are plotted as a function of the zinc content $x$ in Figure 3. Both series of samples based on Zn–Mn and Zn–Co ferrites have the same qualitative behavior of $\sigma_s$ (5 K) with the zinc content. It well compares with the well-known variations obtained for Zn mixed-ferrite bulk materials.75 Although the NPs cores are here surrounded by a surface shell of maghemite, the magnetization still presents a typical maximum with Zn content around $x = 0.5$.

The magnetization decrease observed for larger $x$ values can be understood as in bulk materials. The preference of the diamagnetic Zn ions for tetrahedral sites is responsible for the weakening of the A–B superexchange interaction with increasing Zn concentration.75,76 In the present core/shell NPs, this decrease is smoother and several reasons could account for this behavior. First, the chemical composition of the NPs is not homogeneous, with a maghemite surface shell surrounding a nonstoichiometric core, this effect being more pronounced for large zinc content. Second, there is a supplementary contribution to the NP magnetization due to the disordered surface spins freezing at very low temperatures,72,73 mostly observed for smaller NPs (those with the larger zinc content). At 300 K, the magnetization is smaller, as expected, and for Zn–Mn samples slightly decreases with zinc percentage. The difference between $\sigma_s$ values at 5 and 300 K increases with Zn content. This could be due to finite-size effects which would modify the thermal dependence of the NPs magnetization.

From hysteresis loops of Figure 2c,d we deduce the coercive field $H_c$ at 5 K, and their values are presented against Zn substitution in Figure 3c for both Zn–Mn and Zn–Co series of samples. For the Zn–Co series, the $K_{ef}$ values (see Table S3, SI) well-compare with that of cobalt ferrite77 around 3.53 $\times 10^5$ J/m$^3$. They also decrease by almost 1 order of magnitude from $2.5 \times 10^5$ to $5.5 \times 10^4$ J/m$^3$ with increasing Zn content. It reflects the incorporation of less anisotropic ions in the structure of the NPs’ core. On the other hand, for Zn–Mn series the deduced $K_{ef}$ values also can be compared to those of Mn ferrite bulk materials78 (2.50 $\times 10^5$ J/m$^3$). They are larger and increase with the Zn content, from 1.0 to $1.5 \times 10^4$ J/m$^3$. This is most probably due to a surface contribution (as in maghemite79) associated with the NPs size,80 which is smaller than those of Zn–Mn ferrite-based NPs. As magnetic anisotropy affects Néel relaxation and as we deal here with a wide range of anisotropy values, we expect to probe the relation between both magnetic hardness and exchange coupling between hard/soft and soft/soft core/shell NPs on the heat generation performance.

3.3. Curie Temperature and Thermomagnetic Coefficient. The thermal dependence of the magnetization $\sigma$ at 80 kA/m of pressed powders of NPs is illustrated with ZM9 and ZC9 NPs in Figure S4 (SI). For both kinds of samples, the magnetization decreases with increasing temperatures, as expected. This thermal dependence allows one to determine the Curie temperature ($T_C$) and the thermomagnetic coefficient ($k_{TAM}$). Both characteristics are obtained by linear extrapolation of $\sigma$ as a function of temperature. Their respective values are plotted as a function of the Zn content in Figure 4. In Zn–Mn samples, $T_C$ decreases from 645 to 480 K when the amount of Zn in the NPs’ core increases from $x = 0.11$ to 0.65. In the investigated range of Zn composition, the values found here are larger than those extracted from the literature for NPs of stoichiometric Mn–Zn ferrite.85,50,68,89,90 It should be related first to the chemical composition of the core, which is nonstoichiometric with Zn loss and Fe enrichment. Second, the presence of the surface shell of maghemite should also lead to an increase of the $T_C$ of the core/shell NPs since for
maghemite bulk materials $T_C$ lies in the range 820–986 K, a value much larger than that of Mn–Zn ferrites. In Zn–Co ferrite-based samples, $T_C$ decreases from 687 to 480 K for increasing Zn substitution from $x = 0.11$ to 0.59. Such values are also larger than those commonly observed in pure Zn–Co ferrite NPs and reflect once more the combined effects of nonstoichiometry of the core composition and surrounding maghemite shell.

Figure 4 also depicts, as a function of Zn substitution, the values of the thermomagnetic coefficient $k_{TM} (-\Delta r/\Delta T)$ obtained in the presence of an external field of 80 kA/m and averaged over the $T$-range 300–350 K. For the Zn–Mn series, $k_{TM}$ is approximately constant, independent of the Zn content (except for ZM9 powder sample that showed a slight decrease). These measured values are smaller than those of pure Zn–Mn ferrite NPs. For Zn–Co ferrite NPs, $k_{TM}$ increases with Zn content; nonetheless, these values are approximately 3 times lower than those reported in the literature. The smaller values observed in the two series of samples are closely related to iron enrichment in both core and shell phases. Indeed, the Curie temperature of maghemite is larger than that of Zn-substituted ferrite materials, leading to a smoother decrease of the magnetization with temperature. Moreover, since the applied field is too low to saturate the core of the NPs, the different behavior between both series might be related to the magnetic anisotropy, thus increasing the thermomagnetic coefficient with Zn content for Zn–Co samples. As seen in Figure 5 (SI), powder samples ZC1–ZC4 and ZM9 show the largest coercive fields at 300 K and also the smallest values of thermomagnetic coefficient. For softer magnetic NPs, the thermal energy would disorient the magnetic moments more efficiently, decreasing the magnetization more abruptly.

3.4. Specific Power Absorption. The SPA results for some representative NPs dispersed in aqueous medium are presented in Figure 5 as a function of both magnetic field amplitudes and frequencies. They are analyzed in the framework of the linear response theory (LRT), which calculates the dynamic hysteresis area by assuming that the magnetic response of an assembly of magnetic NPs is linear with the external field.$^7$ The validity of LRT takes place when $\xi \ll 1$, which is satisfied when the amplitude of the applied magnetic field is much smaller than the anisotropy field of the NPs ($H_0 \ll H_k$)$^6$ (it is true here as $E/k_B T \geq 1$ for all the magnetic fluid samples at 300 K). Then, under such conditions and for the randomly orientated magnetic NPs, the density of absorbed power is given by

$$P = \frac{1}{2} \mu_0 \chi_0 H_0^2 \frac{(2\pi f)^2 \tau}{1 + (2\pi f\tau)^2}$$

where $\chi_0 = \mu_0 M_s^2 V/3k_B T$ is the initial static susceptibility. Figure 5a shows the SPA values as a function of the square of the magnetic field amplitude for ZM2, ZM6, and ZM9 NPs, both dispersed in aqueous medium. Gray zones indicate the range of validity of LRT. (b) Frequency dependence of SPA (12 kA/m). The fit represents the frequency dependence of SPA in the framework of LRT.

Let us be precise about the domain of validity of the LRT for the different NPs with different hardnesses. If the condition $H_0 \ll H_k$ is valid for $H_0 = 12$ kA/m with ZM2 NPs, which are the less anisotropic among the two series of NPs, it will also stand for the other ones, which are all magnetically harder. Figure 5b depicts the frequency dependence of SPA at $H_0 = 12$ kA/m for ZC1, ZM2, ZM3, and ZM9 NPs. Data are fitted using eq 3 with two free parameters: $a = -\mu_0 \chi_0 H_0^2$ and $\tau$. The best fit (dashed lines) values shown in the figure well-compare to those reported in the literature.$^{11,12}$

It is worth noting that, considering only Néel relaxation, the $\tau$ values could be translated into magnetic anisotropy constants at 300 K. They are in agreement with the $K$ values calculated at 300 K from the data obtained in the low-temperature
experiments using the methodology applied by Garaio et al.84 Future studies intend to investigate more precisely, in these core/shell NPs, the combined effect of size variations and anisotropy on the relaxation times

To compare all the samples investigated here, assuring LRT to be valid in all samples, the measurements have been performed at \( f = 265 \text{ kHz} \) and \( H_0 = 12 \text{ kA/m} \), since in this regime all the samples follow the LRT. The results are collected as a function of Zn content for both series in Figure 6.

Figure 6. SPA values obtained at 12 kA/m and 265 kHz for the entire Zn−Mn and Zn−Co series as a function of Zn content (x). (a) The inset depicts the variation of the NPs crystalline diameter with x.

In the case of NPs based on Zn−Mn core, those with the lower Zn-content exhibit the best efficiency for heat generation (SPA = 130 W/g for \( x = 0.18 \)) and the SPA almost vanishes for larger values of Zn substitution. In this series with negligible variations of magnetic anisotropy, this feature seems to be most likely related to the decrease of both the diameter and saturation magnetization of NPs at 300 K with Zn substitution (see Figure 6a and the inset of Figure 3a). For the Zn−Co series, SPA values are at the same level (∼20 W/g) as those obtained for Zn−Mn series with \( x \geq 0.25 \), except for ZC3 and ZC4 samples, which reach a maximum around 60 W/g. The presence of this maximum is most likely related to the combination of several characteristics, which will be discussed in detail in the following section.

### 4. DISCUSSION

We discuss here the performances of the studied core/shell NPs based on Zn−Mn and Zn−Co ferrite cores, focusing mainly on heat generation phenomena when dispersed in an aqueous medium. As described above, these NPs present SPA values comparable to those previously reported with similar ferrite NPs dispersed in water.11,12,23,85 The increase of Zn content induces the weakening of the superexchange interactions and affects not only the saturation magnetization but also the Curie temperature and power absorption. These NPs could therefore be good candidates for magneto-hyperthermia treatments in a more-safe strategy, since it would help one to avoid overheating and the necrosis of healthy tissues. It is worthwhile to remark that the efficiency of NPs’ heat generation is affected by the diameter of the NPs, the magnetic anisotropy constant, and the saturation magnetization, all parameters that were carefully determined in previous sections.

Figure 7a depicts SPA as a function of the NPs’ crystalline diameter for samples with similar Mn or Co content (y index ranging between 0.67 and 0.48) and \( \sigma_\sigma \) in each Zn−Mn and Zn−Co series. We observe that the SPA values, which are approximately the same for both kind of samples, are obtained in different ranges of diameters. It illustrates the role of the magnetic anisotropy constant on the heat generation efficiency, which shifts the value of the NP diameter that corresponds to the maximum of the SPA5,6 defined by \( 2\pi f \tau_\text{N} = 1 \). However, an increment of anisotropy constant leads to an increase of \( \tau_\text{N} \) and consequently \( \tau_\text{t} \), but it does not necessarily maximizes the heating efficiency. Indeed, for highly anisotropic NPs, the magnetic moment is strongly coupled with the anisotropy field and preferentially relaxes by mechanical rotation. In this case, larger SPA will be obtained for smaller NPs when compared to lower anisotropic ones. In such a scenario, an increment of media viscosity, such as often occurs in biological media, will hamper the Brownian motion, decreasing the heat generation efficiency of the NPs.23 Moreover, the larger SPA values observed for ZC3 and ZC4 samples probably have to be attributed to the large values of saturation magnetization (see Table S3, SI) conjugated with the lowering of the anisotropy barrier induced by increasing zinc content.

Figure 7. (a) Comparison between SPA values (at 12 kA/m and 265 kHz) of some Zn−Mn- and Zn−Co-based ferrite NPs with similar Mn or Co fractions (y) and (b) SPA results obtained at different frequencies and magnetic field amplitudes.
Figure 7b illustrates the SPA values at different frequencies (265–831 kHz) and field amplitudes (4–24 kA/m) for NPs of ZM2 and ZM3 samples, both dispersed in water. At 24 kA/m and 831 kHz, we obtain a SPA value of 799 W/g for the ZM2 sample decreasing to 304 W/g for ZM3. Even if the $H_f$ product obtained with the field amplitude and frequency of our experiments is well above the human tolerable limitation of $H_f < 4.85 \times 10^4$ A m$^{-1}$ s$^{-1}$, these SPA values are comparable with others reported for similar or even more severe conditions. In such magnetically soft ferrites, increasing both NPs’ diameter and saturation magnetization (by slightly modifying the synthesis conditions) might be the key to increase the SPA. However, the size should not overcome the limit where Brownian relaxation becomes the dominant process, since the heating efficiency of the NPs must be maintained in medium with larger viscosities. Nonetheless, considering that the NP diameters of some approved MRI contrast agents lies in the range of 20–150 nm and that the size reduction leads to larger longitudinal relaxivities, our rather small NPs could be also suitable as MRI contrast agents in a theranostic approach.

The SPAs obtained for the Zn–Co series are very similar when compared to those of the Zn–Mn one, except for ZM2 NPs, which exhibits the highest value, most likely related to its higher diameter and saturation magnetization. In fact, considering that the Zn–Co series of samples present the highest anisotropy contrast between core and shell materials, a larger exchange coupling at the interface would be expected, which could increase the heating efficiency.

Here, one would wonder about the nature of this exchange coupling, an issue which is not very clear in the literature. Indeed, an exchange bias field has already been observed in ultrasmall core/shell ferrite NPs (∼3 nm), synthesized by the same chemical route as used here. The exchange bias field originates at an internal interface between the well-ordered ferrimagnetic core and a surface layer of disordered spins. This exchange bias is found to be much larger for NPs based on a harder ferrite core. Nevertheless, this interfacial exchange coupling typically vanishes at temperatures above 50 K and has not been observed with the present NPs at room temperature. Thus, in our case, it seems highly improbable that this exchange bias field can be involved in the mechanism responsible for increasing the SPA. However, the thickness of the chemical shell of the here probed NPs is ∼0.5 nm, which is smaller than their magnetically disordered surface layer. Then, the elaboration and study of core/shell NPs with different proportions of core and shell ferrite phases may enlighten the nature of the exchange coupling and its contribution to the heating efficiency in hyperthermia.

5. CONCLUSIONS

Zn,M,F,Fe$_2$O$_4$ (M = Mn, Co) NPs were synthesized by hydrothermal coprecipitation. Zn substitution yielded deviations from nominal stoichiometry in both sets of samples and a significant change in diameter that was more pronounced in the Zn–Mn series. We succeeded in producing a core/shell structure, consisting of a mixed ferrite core coated with a thin layer of maghemite. This strategy allowed one to disperse the Zn,M,F,Fe$_2$O$_4$@γ-Fe$_2$O$_3$ NPs in acidic medium to obtain long-term-stable ferrofluids.

Regarding the magnetic measurements properties of the NPs, we found that Zn substitution modulates the NPs’ magnetization and anisotropy in different ways. The magnetic anisotropy of Zn–Co ferrite-based NPs changes drastically from $K_f = 2.5 \times 10^5$ J/m$^3$ (low zinc content) to $3.5 \times 10^4$ J/m$^3$ (high zinc content), while the anisotropy of Zn–Mn ferrite-based NPs changed less than 40% ([0.9–1.4] ×10$^4$ J/m$^3$) for any Zn contents.

Consistently, these differences of magnetic anisotropies were reflected in the behavior of the NPs’ thermomagnetic coefficient as a function of their Zn content (investigated in pressed powders). $T_c$ values decreased with the Zn content for both series of samples, resulting in NPs with lower $T_c$ values than other iron oxides, specially magnetite and maghemite. Decreasing $T_c$ down to hyperthermic temperatures would be very interesting, in that it would avoid overheating in biological media. In addition to the magnetization decrease, augmenting the thermal energy allied to the thermally induced changes to both viscosity and magnetic anisotropy is reflect in a decrease of $\tau$, thus shifting the peak of absorption to higher NP diameters, possibly hindering even further the heating efficiency.

The two series of NPs, having either hard or soft ferrite cores and soft ferrite shells, did not present evidence of any interfacial exchange coupling contribution to their power absorption efficiency, as previous reports suggested. Thus, in the absence of exchange coupling effects, synthesizing NPs based on Zn–Mn ferrites with both larger sizes and Zn content would be the key to increase power absorption efficiency while achieving a self-regulated hyperthermia.

In the very near future, we intend to improve the synthesis procedure in order to reach larger nanoparticle diameters and low polydispersity. Using different strategies of synthesis routes and magnetic size sorting (using an ionic-strength-induced process as in ref 79) would be the primary directions to be investigated.

## ASSOCIATED CONTENT

### Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcc.7b11014.

Figures showing X-ray diffractograms of core/shell NPs, lattice parameters of Zn–Mn and Zn–Co sets of samples, core/shell chemical structure of ZC3 sample, Curie temperature determination, and hysteresis cycles at 300 K for NPs samples in powder form and tables summarizing the chemical characterization of core/shell NPs, the structural characteristics of core/shell NPs, and the magnetic properties of the samples (PDF)

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Notes

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