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Structural perspective on revealing heat dissipation behavior of CoFe\textsubscript{2}O\textsubscript{4−Pd nanohybrids: great promise for magnetic fluid hyperthermia

This interdisciplinary work with contributions from physics, chemistry, and medicine introduces combinations of superparamagnetic CoFe\textsubscript{2}O\textsubscript{4} (CFO) and palladium nanoparticles as thermo-seeds for magnetic fluid hyperthermia treatment. In comparison with bare CFO, the palladium decorated nanohybrids exhibit higher magnetization, larger magnetocrystalline anisotropy, and an improved specific absorption rate under applied alternating magnetic fields due to their magnetic moment interaction. The present piece of application-driven basic research gives an important insight into utilization of tailored nanohybrids.

As featured in:

Structural perspective on revealing heat dissipation behavior of \( \text{CoFe}_2\text{O}_4–\text{Pd} \) nanohybrids: great promise for magnetic fluid hyperthermia†

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Loss mechanisms in fluid heating of cobalt ferrite (CFO) nanoparticles and CFO–Pd heterodimer colloidal suspensions are investigated as a function of particle size, fluid concentration and magnetic field amplitude. The specific absorption rate (SAR) is found to vary with increasing particle size due to a change in dominant heating mechanism from susceptibility to hysteresis and frictional loss. The maximum SAR is obtained for particle diameters of 11–15 nm as a result of synergistic contributions of susceptibility loss, including Néel and Brownian relaxation and especially hysteresis loss, thereby validating the applicability of linear response theory to superparamagnetic CFO nanoparticles. Our results show that the ferrofluid concentration and magnetic field amplitude alter interparticle interactions and associated heating efficiency. The SAR of the CFO nanoparticles could be maximized by adjusting the synthesis parameters. Despite the paramagnetic properties of individual palladium nanoparticles, CFO–Pd heterodimer suspensions were observed to have surprisingly improved magnetization as well as SAR values, when compared with CFO ferrofluids. This difference is attributed to interfacial interactions between the magnetic moments of paramagnetic Pd and superparamagnetic/ferrimagnetic CFO. SAR values measured from CFO–Pd heterodimer suspensions were found to be 47–52 W g\(_{\text{ferrite}}^{-1}\), which is up to a factor of two higher than the SAR values of commercially available ferrofluids, demonstrating their potential as efficient heat mediators. Our results provide insight into the utilization of CFO–Pd heterodimer suspensions as potential nanoplatforms for diagnostic and therapeutic biomedical applications, e.g., in cancer hyperthermia, cryopreserved tissue warming, thermoablative therapy, drug delivery and bioimaging.

1. Introduction

Magnetic fluid hyperthermia (MFH) is a promising new therapeutic approach to synergistically improve efficacy in a wide range of biomedical applications, including controlled thermal therapy of cancer, cellular magnetic activation (magnetogenetics), magnetic targeted drug delivery and the adjustment of glucose metabolism and the neuronal activity of cells.\(^1–3\) Magnetic ferrite nanoparticles have attracted the most attention because of their biocompatibility, superparamagnetic behavior, strong magnetization (\(M\)) and effective anisotropy (\(K\)). Cobalt ferrite (CFO) nanoparticles, in particular, exhibit significant heat-generation ability as a result of their higher magnetocrystalline anisotropy\(^4\) when compared with other ferrites.

CFO nanoparticles can be delivered to targeted tissues/organisms by direct infusion into cancerous tissues or initial intravenous infusion and subsequent magnetic guidance to a tumor using DC magnetic field gradients. Local magnetic
heating of nanoparticles (up to 43 °C) using an AC magnetic field is used to selectively damage cancerous cells in their direct vicinity. In addition, there may be beneficial effects for blood flow and for conventional cancer therapies. For example, an improvement in blood supply by heating increases the efficiency of chemotherapy, while for temperatures above 40 °C the ability of cancerous cells to repair radiation damage is severely decreased, significantly increasing tumor regression in radiation therapy. The heat generation ability of magnetic nanoparticles can also be used for targeted drug delivery applications. For example, during liposomal drug delivery by magnetic nanoparticles local heat generation can be used to trigger the release of an encapsulated drug by the activation of a thermal release mechanism, which improves the drug release rate.

Although the generation of heat can be regulated by the amplitude ($H$) and frequency ($f$) of the applied magnetic field, a practical safety limit should be observed to avoid possible adverse phenomena, such as additional non-localized heating in the body by eddy currents during clinical treatment. The heat-generation efficiency should therefore be improved by tailoring the properties of nanoparticles, such as their size, chemical composition, structural and magnetic properties, as well as their biocompatibility. The size of a nanoparticle directly influences its crystallinity, stability, biocompatibility and magnetic properties, e.g., its superparamagnetic limit. In addition, the ferrofluid concentration affects heat generation, agglomeration and the cell-cytotoxicity of nanoparticles.

Since the magnetic responsivity of nanoparticles decisively influences possible applications in magnetic hyperthermia, there were several intense research efforts during the past years to develop new approaches including the design of magnetic nanoparticles with non-spherical shapes such as cubic, flower-shape, and elongated particles. For non-spherical particles, the total magnetic anisotropy is enhanced due to the additional shape anisotropy. Furthermore, two-phase magnetic nanoparticles consisting of a magnetic core and a non-magnetic shell could provide a higher heat generation, although in many cases the total mass magnetization is significantly decreased. In some other cases, agglomeration of nanoparticles to larger magnetic clusters could increase the heat generation ability, but limits their use for in vivo applications.

Colloidal hybrid nanoparticles with diverse structures, such as mosaic, core–shell, multi-core–shell and heterodimer, act as multipurpose agents for biomedical applications. Among these structure models, heterodimers of CFO nanoparticles and noble metal particles of palladium (Pd) have been recognized as efficient multifunctional agents for potential biomedical purposes, including photothermal therapy, cell separation and optical tracking of therapeutic agents. However, the relatively poor heat transfer efficiency of CFO necessitates the use of a highly concentrated ferrofluid for effective hyperthermia treatment, which inhibits its practical utilization in clinical treatments.

In addition, there are some concerns about the biocompatibility of CFO nanoparticles but several studies introduce CFO nanoparticles as a promising candidate for bioapplications. Additionally, it is experimentally confirmed that the CFO nanoparticles could effectively inhibit the proliferation of carcinoma cells.

There are two general approaches to further improve the biocompatibility of CFO: the first is based on covering the magnetic cores through biocompatible ceramic/polymeric shells such as silica, PVA, PEG, and citrate. The second is based on the synthesis of CFO substituted with biocompatible metals like Ag or Cu. However, in both cases, the total magnetization might be negatively affected.

In contrast to most diamagnetic noble metals, such as Cu, Ag and Au, Pd is a paramagnet with a large magnetic susceptibility. Ultrasmall paramagnetic Pd nanoparticles may exhibit even enhanced magnetic properties compared to the corresponding bulk material and can maintain the basic net magnetic features of CFO, accompanied by emerging new optical properties. Moreover, the biocompatibility and physicochemical stability of Pd nanoparticles result in the formation of biocompatible CFO–Pd heterodimers. The fabrication and magnetic characterization of nano-hybrids consisting of magnetic nanoparticles and noble metals with different morphological structures, such as core–shell, phase segregated and heterodimers, have been discussed. However, a study of the magnetic heat generation ability of CFO–Pd heterodimers has not yet been reported.

In the present work, the dominant mechanisms of magnetic heat generation of CFO nanoparticles upon application of an alternating magnetic field are studied by the experimental measurement of the specific absorption rate (SAR). The effects of particles size, ferrofluid concentration magnetic field amplitude on the heat-generating ability of CFO nanoparticles are investigated. Furthermore, magnetic features and the associated heat generation properties of CFO–Pd heterodimers in two different ranges of particles size are characterized and discussed. Correlations between SAR value and structural factors, i.e. degree of crystalinity, crystallite size and cation distribution in CFO, are evaluated and optimized. To the best of our knowledge, there have been no previous studies of the magnetic and heat-generating properties of CFO–Pd heterodimers. Accordingly, this is the first investigation of CFO–Pd hybrid architectures for use in magnetic hyperthermia biomedical applications.

2. Experimental section

2.1. Chemicals

In order to synthesize CFO nanoparticles, precursors consisting of FeCl$_3$·6H$_2$O and CoCl$_2$·6H$_2$O (> 99%, Merck) as magnetic cation resources, NaOH (1 M, Merck) as an adjusting agent of the solution pH, and citric and nitric acids (> 99.5%, > 65%, Merck) as chelating agents, were employed. Materials for preparing CFO–Pd heterodimers structures included PdCl$_2$ (> 99%, Merck), KCl (> 99.5%), HCl (37–38%) and citric acid. All materials, solutions and surfactants were used without any further purification.
2.2. Synthesis of CFO nanoparticles and CFO–Pd nano-heterodimers

Superparamagnetic CFO nanoparticles were synthesized using a modified coprecipitation method. The pH of a mixture solution of Fe and Co chloride salts was adjusted using NaOH solution (1 M) to prepare the required condition for magnetic ferrite phase reduction. The size of the synthesized CFO particles was controlled by the reaction temperature, which was selected according to the design of experiments approach. Citric and nitric acids were used as chelating agents to control the size, size distribution and physical stability of the nanoparticles, which were dispersed in a colloidal ferrofluid after magnetic separation and washing progressions. Further information about the CFO nanoparticle synthesis process is provided in our previous works.\textsuperscript{39,40} In order to randomly attach metallic Pd nanoparticles to CFO nanoparticles and create CFO–Pd heterodimers, 2 vol% of HCl solution was added and sonicated for 20 min to disperse the agglomerated CFO clusters and activate their surfaces to connect with the ultra-small Pd particles. Mixing of the PdCl\textsubscript{2} and KCl salts in aqueous medium for 30 min led to the formation of a water-soluble intermediate phase of K\textsubscript{2}PdCl\textsubscript{4}. After adding CFO nanoparticles and homogenizing for 60 min at room temperature, the pH of the medium was regulated to be in the range 12–13 using NaOH solution. The resulted colloidal solution was incubated for 24 h with controlled nitrogen flow. The resulting samples were centrifuged, magnetically separated, washed with ethanol and deionized water, pH-naturalized and dried at ambient temperature using a vacuum dryer.

2.3. Characterization

The sizes and structures of the synthesized CFO nanoparticles were characterized using dynamic light scattering (DLS, MALVERN, Zetasizer Nano ZSP) and X-ray diffraction (XRD, Bruker Advance 2), respectively. Characterization of the microstructural and elemental features of the CFO nanoparticles and CFO–Pd heterodimers was carried out using high-resolution scanning transmission electron microscopy in high-angle annular dark-field mode (HAADF-STEM, FEI Titan G2 80-200 ChemiSTEM),\textsuperscript{41} and energy dispersive X-ray spectroscopy (EDS, Titan G2 80-200 ChemiSTEM) at a primary energy of 200 keV. Magnetic properties were measured using a vibrating sample magnetometer (VSM, VSM-PPMS-14T) at up to 13 T and ambient temperature (300 K).

2.4. Magnetic fluid heating and SAR measurement

In order to study magnetic fluid heating, ferrofluid temperature variations as a function of time were measured at constant volume (1 ml for all samples) at different concentrations in the presence of an AC magnetic field with $H = 19$ mT ($I = 33$ A) and $f = 318$ kHz. In order to assess the effect of the magnetic field amplitude, some experiments were also conducted at $H = 12$ mT ($I = 22$ A). All of the measurements were performed at ambient temperature and atmosphere using a TruHeat HF 3010 system equipped with a fiber-optic temperature probe. The four-turn induction coil had a total height of approximately 6 cm and an inner diameter of 10 cm. In order to eliminate the effect of a different starting temperature, curves were initially plotted (Fig. 3) as a function of temperature difference ($\Delta T$) and the magnetic field was applied after fixing the indicated probe temperature. The SAR factors of the samples were calculated based on the well-established equation\textsuperscript{42}

$$\text{SAR} = C \times \left( \frac{\Delta T}{\Delta t} \right) \times \left( \frac{m_s}{m_f} \right).$$

where $C$ is the specific heat of water (4.184 J K\textsuperscript{-1} g\textsuperscript{-1}), ($\Delta T/\Delta t$) is the initial slope of the temperature vs time curve, $m_s$ is the mass of the sample and $m_f$ is the mass of ferrite, which is equal to the total mass of the magnetic cations and oxygen anions in the bare CFO samples and the total mass of the magnetic cations, oxygen anions and Pd nanoparticles in the CFO–Pd samples. The total mass of CFO (or CFO + Pd) is only preferred over the total mass of magnetic cations in SAR calculations to achieve more realistic values for applications.

The change in temperature with time is related to the dissipated power $P$ according to the well-known relation

$$P = m_s C \left( \frac{\Delta T}{\Delta t} \right).$$

In this regard, the SAR can be understood as the dissipated power per mass of magnetic material (here, CFO). For magnetic heating by superparamagnetic particles in an AC magnetic field, the dissipated power depends on external factors, such as magnetic field amplitude $H$ and frequency $f$, as well as internal factors, such as the imaginary part of the AC magnetic susceptibility $\chi''$ and the effective relaxation time $\tau$, according to the equations\textsuperscript{43}

$$\chi'' = \frac{\omega \tau}{1 + (\omega \tau)^2} z_0,$$

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

where $\omega$ is the angular frequency $f = \frac{\omega}{2\pi}$, $\chi_0$ is the equilibrium susceptibility, $H_0$ is the small applied field and $\mu_0 = 4\pi \times 10^{-7}$ T m A\textsuperscript{-1} is the vacuum permeability.

In order to explore the effective influence of size and concentration of the CFO nanoparticles, as well as the amplitude of the magnetic field, on heat generation, several experimental runs were performed. The characteristics of the sample series that were used to evaluate the size, concentration, and field amplitude dependent SAR factors of bare CFO and CFO–Pd architectures are presented in the ESI\textsuperscript{†} (Tables S1–S3). In all of the MFH measurements, the approximate uncertainty is 2%.

2.5. Optimization

Optimization studies were carried out based on the central composite design (CCD) approach of response surface methodology (RSM) using the DESIGN EXPERT software program to understand the correlation between SAR values and structural factors, such as crystallite size, degree of crystallinity and cation
disorder. Predictor models were used to determine optimal values of different factors to achieve the targeted responses. More details about the samples, as well as the optimization process, are presented in the ESL.†

3. Results and discussion

3.1. Structural characterization and magnetometry

The first step in the preparation of CFO–Pd heterodimers and studies of their magnetic fluid heating was the synthesis of CFO nanoparticles with controlled sizes, size distributions and crystal structures. Results of X-ray diffraction (XRD) analyses (Fig. S1 in the ESL†) confirmed the formation of nanoparticles with an inverse spinel structure of the cobalt ferrite phase CoFe$_2$O$_4$, without the detectable formation of undesirable hydroxide or oxide phases such as CoO. Quantitative analyses of the XRD results using Rietveld structure refinement confirmed the crystallization of nanoparticles in the target phase with a significant degree of crystallinity (>$85\%$). Crystallite sizes calculated from pseudo-Voigt profiles showed acceptable correspondence with particles sizes measured using dynamic light scattering (DLS) and microscopic approaches. The hydrodynamic size $d_{	ext{H}}$, size distribution $D_V$ and polydispersity index (PDI) of the synthesized nanoparticles were obtained using DLS. Based on the results (Table S1, ESL†), the mean hydrodynamic diameters of particles in the different samples were determined to be between 7.2 and 40 nm with a narrow distribution, while the corresponding crystallite diameters $d_{	ext{Cryst}}$ varied between 5.8 and 32.2 nm. The hydrodynamic PDI values of all of the samples were below 0.1, indicating the suitability of the nanoparticles for use in biomedical and especially in vivo applications.44

In the next step, the formation of CFO–Pd heterodimers as novel hybrid nano-architectures was evaluated using microscopic analysis. Fig. 1 shows high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) micrographs of CFO–Pd nanostructures obtained from suspension. The mean diameters of the CFO and Pd nanoparticles are approximately 7 and 2 nm, respectively, with a PDI value of 0.07. Morphologically, the nanoparticles are uniform and close-to-spherical. Crystalline CFO nanoparticles with attached ultra-small Pd particles are clearly visible in the STEM images. Element-specific mapping of the CFO–Pd nanostructures was performed using energy-dispersive X-ray spectroscopy (EDS), as shown in Fig. 2. The CFO nanoparticles contain Fe, Co, and O, while Pd is visible in the attached particles.

Magnetometry measurements recorded in applied magnetic fields of up to 13 T at a temperature of 300 K are shown in the ESL† (Fig. S2) for bare CFO nanoparticles with two different hydrodynamic diameters (7.2 and 24.1 nm) and for corresponding CFO–Pd heterodimers. Vanishing coercivity ($\leq 5\text{ mT}$) and remanence ($\leq 4–6\%$ of the saturation magnetization ($M_s$)) are indicative of a primarily superparamagnetic state. Even in high external magnetic fields of 10 T and above, magnetic saturation was not reached. The high-field data, i.e. magnetometry data in fields of above 3 T, were fitted to the law of approach, as shown in the ESL† (Fig. S2 and Table S4) and the effective magnetic anisotropy was extracted from the fitting parameters. In addition, the magnetic DC susceptibility was analyzed. The results are summarized in Table 1. For both nanoparticle sizes, the high-field magnetization, effective

![Fig. 1] HAADF-STEM micrographs of 7.20 nm CFO–Pd heterodimers. The mean sizes of the CFO and Pd nanoparticles are approximately 7 and 2 nm, respectively, with a PDI value of 0.07.
anisotropy and low-field susceptibility were significantly enhanced upon Pd decoration. These effects were more prominent for smaller CFO nanoparticles. The magnetometry results further motivate the investigation of magnetic heating properties for bare and Pd-decorated CFO nanoparticles of different sizes.

3.2. Specific absorption rates

**AC magnetic field amplitude-dependent fluid heating.** In order to obtain a better understanding of the effect of magnetic field amplitude, SAR values were measured from samples with a constant concentration of 15 mg ml⁻¹ for three different ranges of particle hydrodynamic diameters (10.24, 13.70 and 17.77 nm) in applied magnetic fields with amplitudes of 12 and 19 mT (and currents of 22 and 33 A, respectively). The frequency \( f = 318 \) kHz was constant. The product \( H f \) was below the safety limit for both amplitudes (\( 3.0 \times 10^9 \) and \( 4.8 \times 10^9 \) A m⁻¹ s⁻¹, respectively) and appropriate for in vivo biomedical applications.⁴⁄ The resulting field-amplitude-dependent SAR values are given in the ESI† (Fig. S3). The dissipated power and, as a consequence, the SAR values of all of the samples decreased considerably for smaller magnetic field amplitudes, in agreement with previous studies.⁴⁶,⁴⁷

Since the magnetic susceptibility is field-dependent (Fig. S4, ESI†) and a ferromagnetically blocked fraction of nanoparticles is present, in particular for the larger particles, there are deviations from a simple \( H^2 \) dependence. As a minor additional effect, the effective relaxation time may be changed as a result of magnetic dipolar interactions between the particles, which depend on the field-dependent magnetic moments. Further experiments were carried out with AC magnetic field amplitudes of 19 mT, for which the larger SAR values were obtained.

**Particle-size-dependent fluid heating.** All particle-size-dependent measurements were carried out on samples with the same concentration (15 mg ml⁻¹) in the presence of an AC magnetic field with \( H = 19 \) mT and \( f = 318 \) kHz. The measured temperature variation as a function of time \( (\Delta T/\Delta t) \) is shown in Fig. 3a. The temperatures of all of the samples were raised to \( 43 \) °C, the desired temperature for hyperthermia, in reasonable times (less than 15 minutes), confirming the potential effectiveness of CFO nanoparticles for thermal therapy applications. Fig. 3b shows the initial slopes of the \( \Delta T/\Delta t \) curves and corresponding fitted linear equations to calculate SAR values. The difference in heat-generation ability of the different samples within the first two minutes of the measurements is clearly visible.

The extracted SAR values as a function of mean hydrodynamic particle diameter are shown in Fig. 4a. The variation in particle size leads to irregular changes in SAR factor, meaning that two SAR peaks are observed for particle hydrodynamic diameters of 12.05 and 14.77 nm. The activation of different heat loss mechanisms in different size ranges can result in such features in the dependence of SAR value on nanoparticle size. Increasing the particle hydrodynamic diameter from 7.2 to 40 nm activated different loss mechanisms sequentially, including susceptibility (Néel and Brownian relaxation), hysteresis and frictional losses as the dominant mechanisms. In order to estimate the size dependence of the different relaxation channels, Néel and Brownian relaxation times were calculated based on the equations⁹,⁴⁸

\[
\tau_N = \frac{\text{K}_N V}{4 \pi \mu_0} \quad \text{(5)}
\]

\[
\tau_B = \frac{4 \pi \eta_0 \gamma_i^3}{k_B T} \quad \text{(6)}
\]

where \( \tau_N \) is the Néel relaxation time, \( \tau_B \) is the Brownian relaxation time, \( \tau_0 = 10^{-9} \) s,⁴⁶,⁴⁷,⁴⁹ \( V \) is the geometric volume of the
nanoparticle, $K$ is the effective anisotropy, $k_B$ is the Boltzmann constant, $T$ is the measurement temperature, $\eta$ is the viscosity of the ferrofluid and $r_H$ is the hydrodynamic radius of the nanoparticle. The effective relaxation time was calculated from the Neél and Brownian relaxations using the expression

$$t_{\text{eff}} = \frac{t_N \times t_B}{t_N + t_B}. \quad (7)$$

Fig. 4b shows the Neél, Brownian and effective relaxation times plotted as a function of particle hydrodynamic diameter. Initially, in the range of small particle diameters up to 11 nm, the dominant mechanism of heat generation is Neél relaxation loss. Further augmentation of particle size changes the dominant mechanism into Brownian relaxation loss, reaching a maximum SAR value at a particle diameter of approximately 12.05 nm. For particle sizes above the superparamagnetic dimension $D_S$, the dominant mechanism is hysteresis loss. The size-dependent heating trends and associated dominant mechanisms are consistent with linear response theory (LRT) for superparamagnetic nanoparticles and the Stoner–Wohlfarth model of hysteresis for ferrimagnetic structures. The Neél relaxation loss in smaller nanoparticles is associated with a rotation of the magnetic moments inside them, while Brownian relaxation depends on the ferrofluid viscosity, as it is associated with particle rotation in a fluid medium. The dominant mechanism of magnetic heat generation changed from Neél to Brownian relaxation loss with increasing particle diameter, followed by a subsequent increase in fluid viscosity. Ferrimagnetic states and corresponding hysteresis loops were formed in particles with sizes larger than $D_S$, with hysteresis loss identified as the dominant mechanism with increasing particle size, in accordance with the dynamic hysteresis model (DHM), as a result of an enhancement in their magnetization and magnetic anisotropy. The area of influence of each mechanism could be distinguished in the particle-size-dependent SAR variation curve. In order to determine the dominant loss mechanism, the critical diameter of susceptibility loss limit, $D_{\text{Sus}}$, in zero/small magnetic field is given by the equation

$$D_{\text{Sus}} = \left[\frac{6k_B T \ln(f_m \tau_0)}{\pi K}\right]^{\frac{1}{3}}, \quad (8)$$

where $f_m$ is the maximum frequency of the magnetic field and $\tau_0$ is the characteristic spin-lattice relaxation time.
attributed to the maximum efficiency of Neél relaxation loss. An increase in particle size leads to a gradual decrease in the effectiveness of Neél loss and to the development of Brownian relaxation as the dominant loss mechanism. The Brownian mechanism efficiency is maximized at the utmost point of the SAR first peak at approximately 12.05 nm. Although the areas of influence of these mechanisms overlap each other, only one of them is dominant in each size range. An increase in particle size up to the ferrimagnetic range leads to a dramatic weakening of the relaxation loss mechanisms and associated SAR values. Above a particle size of 13.67 nm (from eqn (8)), the dominant mechanism changed to hysteresis loss and the SAR value increased with particle diameter. Accordingly, two pronounced SAR values were detected in the size ranges 14–15 nm and 11–12 nm, in order of intensity. The SAR value was gradually reduced with increasing particle diameter, especially above 17 nm, due to the insufficient field amplitude (19 mT) to switch the magnetic moments of larger particles. In this particle size range, frictional loss is likely to be the dominant heating mechanism because of fluid viscosity enhancement.

**CFO concentration-dependent fluid heating.** In order to achieve a better understanding of the influence of ferrofluid concentration on the magnetic heating of CFO nanoparticles, SAR values were measured for samples with different particle diameters for both low and high concentrations (approximately 1 and 15 mg ml\(^{-1}\), respectively) in applied magnetic fields with \(H = 19\) mT and \(f = 318\) kHz. Fig. 5 shows measured SAR values of CFO samples with five different hydrodynamic particle diameters for the two different concentrations. In all of the analyzed samples, the SAR values decreased dramatically (by more than 50%) at high concentrations, probably because of particle agglomeration and inter-particle dipolar interactions (IPDI), such that the superparamagnetic properties of particles almost vanished, the Neél and in particular the Brownian relaxation mechanisms were weakened and the SAR values were reduced. The inter-particle distance could be decreased in highly-concentrated fluids, leading to a bulk-like state performance and to the activation of hysteresis and frictional losses as the dominant mechanisms of fluid heating. Whereas the susceptibility loss vanished for the highly-concentrated samples, the hysteresis loss had a small contribution at \(H = 19\) mT to fluid heating for highly-concentrated particles in agglomerated assemblages. The inverse proportionality of IPDI strength and remanence ratio \(M_r/M_s\) had been confirmed both theoretically and experimentally, while the hysteresis loss was affected strongly by the remanence ratio. Although the results about the effect of concentration were obtained from samples with specific nanoparticle sizes and cannot be generalized to all particle size intervals in size-dependent SAR values, including the maximum peaks, the analogous SAR variations of measured low- and high-concentrated samples (in five ranges of particle size) increase the probability of a similar trend over all size ranges, including the maximum SAR values.

With reference to Fig. 5, although the time required for the fluid to reach the temperature needed for successful hyperthermia treatment was shortened for the highly-concentrated suspensions (corresponding to a sharper \(AT/\Delta t\) slope), their heating efficiency was lower. As a result of the unpleasant effects of using a high concentration on the biocompatibility and superparamagnetic behavior of the fluids, the use of low-concentration samples has an important role in biomedical applications.

In order to obtain an understanding of the reproducibility of the measurements and its influence on fluid heating, the SAR values of CFO ferrofluids with four different ranges of particle diameters (11.04, 12.17, 14.77 and 15.70 nm) were obtained from repeated measurements performed for the same concentrations of 15 mg ml\(^{-1}\) and magnetic field amplitudes of 19 mT. SAR values obtained from the first and second runs of measurements varied slightly for all of the samples, as a result of the enhancement of viscous and isothermal remanent magnetizations (VRM and IRM, respectively) during repeated measurements. The results are presented in the ESI† (Fig. S5). Whereas the SAR values decreased in the second run for the two smallest diameters, they increased for the larger diameters. A critical diameter \(D_0\) may be introduced, with \(D_0 = 12.5–14.5\) nm, which is consistent with the value calculated from eqn (8). For single-domain ferrimagnetic nanoparticles with \(d > D_0\), the primary loss mechanism leading to magnetic heating is hysteresis loss, which is increased by contribution from IRM and especially VRM in repeated runs of measurements, leading to improved SAR values. For smaller particles with \(d < D_0\), an increase in remanent magnetization attenuates relaxation losses, particularly those related to the Brownian mechanism, leading to negative SAR differences in samples with particle sizes of 11.04 and 12.17 nm.
Magnetic heating of CFO–Pd ferrofluids. In order to obtain a detailed understanding of the influence of Pd decoration on the magnetic heating of CFO ferrofluids, SAR values of heterodimer CFO–Pd hybrid nanoparticles were monitored for two different ranges of CFO particle size and a constant Pd concentration of \((2 \pm 0.5)\) wt\%. All of the measurements were performed for the same ferrofluid concentration (1 mg mL\(^{-1}\)) and magnetic field \((H = 19\) mT, \(f = 318\) kHz). Fig. 6 shows measurements of SAR values for bare and Pd-decorated CFO nanoparticles with mean hydrodynamic diameters of 7.20 and 24.10 nm. In both sample series, the measured SAR values increased after Pd decoration. This effect was most significant for the larger particles. Although the initial SAR value of the larger bare CFO nanoparticles (24.10 nm) was lower than that of the smaller nanoparticles (7.20 nm), after Pd decoration the larger particles showed higher SAR values.

Interactions between the magnetic moments of the ultra-small paramagnetic Pd and CFO nanoparticles can create a spin-glass-like state.\(^{65–67}\) The Pd structures can then pin or freeze the ordered moments of the magnetic core, ameliorating the SAR values, particularly for larger particles. This mechanism does not affect relaxation losses, which originate at the interfaces of the Pd and CFO structures. Magnetic coupling between the Pd moments and the canted surface spins of the CFO nanoparticles improves the effective anisotropy and increments the SAR values,\(^{68,69}\) in agreement with anisotropy values determined from magnetometry (Table 1). In order to further explain these observations, the magnetometry results are discussed in more detail. We follow the approach of introducing a spin disorder layer \(t_d\) according to the expression\(^{70}\)

\[
t_d = d \cdot \frac{1 - \left( \frac{M_S}{M_{bulk}} \right)}{6}.
\]

where \(d\) is the mean nanoparticle diameter, \(M_S\) is the saturation magnetization of the nanoparticles and \(M_{bulk}\) is the saturation magnetization of cobalt ferrite in its bulk state \((90\) emu \(g^{-1}).dis.\) Based on this relation, the thickness of the spin disorder layer decreases from 3.3 to 2.7 nm for the larger CFO particles and from 1.1 to 0.8 nm for the smaller CFO particles upon decoration with Pd. These results suggest that the relative thickness of the spin disorder layer \(t_d/d\) exhibits a non-linear inverse proportionality relationship to the SAR value of the samples.

Fig. 7 shows the magnetic anisotropies of bare and Pd-decorated CFO samples. After Pd decoration, the anisotropy values of both the large and in particular the small nanoparticles increased significantly. This behavior is associated with the enhanced SAR values. This connection between magnetic anisotropy and SAR is now discussed in more detail for the two different particle sizes and their corresponding loss mechanisms. For the 7.20 nm nanoparticle suspension, combined Néel and Brownian relaxation is the dominant mechanism of fluid heating. As Néel relaxation depends directly on magnetic anisotropy (eqn (5)), the enhanced magnetic anisotropy that results from Pd decoration (Table 1) slows down Néel relaxation. The fluid viscosity of the Pd-decorated samples was also slightly augmented. As this is a critical factor for the Brownian loss mechanism (eqn (6)), it also slows down Brownian relaxation. The Néel and Brownian relaxation times – and therefore also the effective relaxation time – are larger for the CFO–Pd heterodimers, shifting the curves in Fig. 4 (lower panel) upwards. This behavior is qualitatively connected to a shift in SAR towards smaller diameters (Fig. 4, upper panel), such that the small diameter of 7.20 nm is closer to the (first) maximum SAR. The SAR value for the 7.20 nm-sized CFO ferrofluid was subsequently increased after Pd decoration.

The most effective heating mechanisms for the 24.10 nm-sized nanoparticle suspension were hysteresis loss and primarily frictional loss. By increasing the magnetic anisotropy, hysteresis loss was augmented. This effect is enhanced further by the increased remanent (and saturation) magnetization after Pd decoration and
can explain the larger SAR for the 24.10 nm particles. To investigate the hysteresis loss mechanism in detail, in addition to the calorimetry under AC magnetic field, the AC magnetometry and related hysteresis loop could be helpful as has been discussed for the example of the iron ferrite structure in the literature.74–76 For all ranges of particle size, the shape uniformity of the particles decreases as a result of Pd decoration. Frictional loss, especially in larger sized nanoparticles, is then activated as the dominant heating mechanism. Frictional loss results from translational and rotational motions of the nanoparticles in the ferrofluid and magnetic forces according to the expressions47.

![Fig. 8](image-url)  
Fig. 8  (a–c) SAR value variation plotted as a function of synthesis parameters: (a) reaction temperature and pH at a Co²⁺/Fe³⁺ cation ratio of 0.50; (b) reaction temperature and cation ratio at a pH of 11; (c) pH and cation ratio at a temperature of 95 °C. (d–f) SAR value variation plotted as a function of structural characteristics: (d) crystallite size and crystallinity at a cation distribution factor (X_B–X_A) of 0.42; (e) crystallite size and cation distribution at a crystallinity of 76%; (f) crystallinity and cation distribution at a crystallite size of 13.5 nm. The SAR values were calculated from measurement results. The fluids concentration, magnetic field amplitude and frequency were 15 mg ml⁻¹, 19 mT, and 318 kHz, respectively. The error bar was estimated to be approximately 4% in (a–c) and 3% in (d–f) using a predictive model.
values. The SAR values area also augmented by an elevated higher probability of CFO phase formation at higher pH results in significantly enhanced SAR values, as a result of a increase in initial pH value in the range 8–12 parameters. An increment in initial pH value in the range 8–12

\[ F_t = m \nabla H \quad (10) \]

\[ F_r = m \times H \quad (11) \]

where \( F_t \) is the transitional force, \( F_r \) is the rotational force, \( m \) is the magnetic dipole moment and \( H \) is the magnetic field. These magnetic forces are much more effective for larger nanoparticles, especially for single-domain ferrimagnetic nanoparticles, which are characterized by \( d > D_s \) and whose magnetic moments are blocked.

Our results show that the SAR values of CFO–Pd heterodimers improved significantly, when compared to those of bare CFO nanoparticles, due to differences in active heating mechanisms, including Néel and Brownian relaxations, hysteresis and frictional losses. The CFO–Pd samples that we studied have significantly higher SAR values than commercial hyperthermia agents. For example, MagForce AG, which is a pioneer commercial company in hyperthermia treatment, uses agents with SAR values of 10–40 \( \text{W g}^{-1} \text{magnetic element} \) whereas the CFO–Pd ferrofluid studied here has SAR values of 66–68 \( \text{W g}^{-1} \text{magnetic element} \).

### 3.3. Optimization

Given the importance of the heating ability of CFO nanoparticles for magnetic fluid hyperthermia applications, we now address the optimization of synthesis factors, including reaction temperature, initial pH and Co\(^{2+}\)/Fe\(^{3+}\) cation ratio, to achieve a desired SAR value. We also make use of a correlation between SAR value and structural properties such as degree of crystallinity, cation distribution and crystallite size for optimization. More details about the influence of synthesis parameters and structural properties on the magnetic properties of nanoparticles are available in our previous works.\(^{35,36}\) In order to consider the impact of cation distribution, a factor \((X_B - X_A)\) was defined based on the following chemical formula for CFO nanoparticles:

\[ \text{(CoX}_A\text{Fe}_{1-X_A})\text{(CoX}_B\text{Fe}_{2-X_B})\text{O}_4. \]

Fig. 8a–c shows SAR values plotted as a function of synthesis parameters. An increment in initial pH value in the range 8–12 results in significantly enhanced SAR values, as a result of a higher probability of CFO phase formation at higher pH values.\(^{39}\) The SAR values area also augmented by an elevated Co\(^{2+}\)/Fe\(^{3+}\) cation ratio and in particular reaction temperature, due to an increase in both the degree of crystallinity and the crystallite size of the CFO phase.\(^{39}\) Among these synthesis parameters, the reaction temperature and pH have the greatest influence on the SAR value of the CFO nanoparticles, whereas the cation ratio has a minor effect.

Fig. 8d–f shows SAR values plotted as a function of structural properties. Higher degrees of crystallinity of the CFO phase can be seen to improve the SAR values, whereas the influence of crystallite size depends on the degree of crystallinity. At the maximum level of crystallinity, the SAR values increase significantly on enlarging the crystallite size. An increment in cation distribution factor \((X_B - X_A)\) leads to a gradual increase in SAR value, as a result of a change in the magnetic structures of the nanoparticles.\(^{40}\) Experimental measurements can be used to predict optimized correlations for a broader range of factors. Table 2 shows optimized synthesis parameters and structural properties for achieving maximal SAR values. Upon applying an AC magnetic field with \( H = 19 \text{ mT and } f = 318 \text{ kHz} \), the SAR value is maximized at 78.205 \( \text{W g}^{-1} \text{CFO-Pd} \) by adjusting the temperature, pH and cation ratio to 105 °C, 10.61 and 0.52, respectively. The maximal SAR value is predicted for structural properties of the CFO nanoparticles of crystallite size, crystallinity and \((X_B - X_A)\) factor equal to 12.67 nm, 76.32% and 0.45, respectively.

### 4. Summary and conclusions

A systematic study of magnetic fluid heating of CFO nanoparticles and CFO–Pd heterodimers suspensions has been presented, resulting in the introduction of new hyperthermic agents with significantly increased SAR values (by up to two times) when compared to established commercial treatment agents. Studies of the dominant loss mechanism provided an improved interpretation based on a combination of the Rosensweig and dynamic hysteresis models. The measured SAR values were found to depend on the sizes of the magnetic CFO nanoparticles, which resulted in the activation and/or strengthening of different heat generation mechanisms, including susceptibility, hysteresis and frictional losses. The maximal SAR values of bare CFO nanoparticles, which were achieved for particle diameter ranges of 14–15 nm and 11–12 nm in order of intensity, were attributed to hysteresis and susceptibility mechanisms, respectively. The calculated critical limit for susceptibility loss \( D_{\text{Sus}} = 13.66 \text{ nm} \) is consistent with this interpretation and with data reported in the literature.\(^{78,79}\) Our experimental size-dependent SAR measurements validate the LRT model for CFO nanoparticles, which is not sufficient to describe the properties of magnetic iron oxide nanoparticles.\(^{4,70}\) In accordance with the Rosensweig model, fluid heating of CFO nanoparticles is
found to depend significantly on magnetic dipole–dipole interactions, which are influenced by the ferrofluid concentration, magnetic field amplitude and $H_f$ factor. Although hysteresis loss showed the highest SAR intensity in a size-dependent manner and represents the most effective fluid heating mechanism, it was found to be valid for ferrimagnetic nanoparticles characterized by $d > D_s$, which are not ideal candidates for biomedical in vivo applications because of the probability of magnetic agglomeration and complications such as thrombosis. For superparamagnetic nanoparticles, susceptibility loss, including Néel and Brownian relaxations, was the primary contributor to fluid heating, demonstrating optimal magnetic, hyperthermic and biological properties. The SAR values of CFO–Pd heterodimers were substantially improved, when compared to those of bare CFO nanoparticles, because of interfacial interactions between the magnetic moments of Pd and CFO nanoparticles, resulting in reduced spin canting, stronger internal fields and enhanced anisotropy without any substantial effect on the relaxation mechanisms. The calculated SAR values of CFO–Pd heterodimer suspensions were 47–52 W g_{Ferrite}^{-1}, which are higher than the best-reported values in commercially available ferrofluids. These findings present an important insight into the utilization of hybrid nanostructures of CFO–Pd heterodimers with excellent time-dependent temperature enhancement for biomedical applications.

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**Conflicts of interest**

The authors declare no conflict of interest.

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