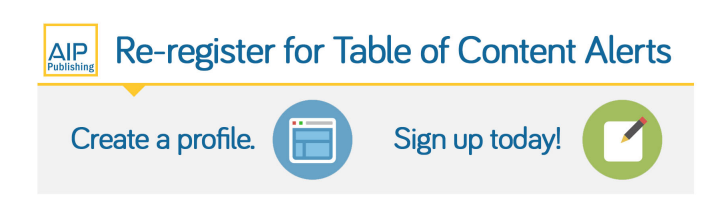




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Quasi-static magnetic measurements to predict specific absorption rates in magnetic fluid hyperthermia experiments

D. F. Coral,¹ P. Mendoza Zélis,¹ M. E. de Sousa,¹ D. Muraca,² V. Lassalle,³ P. Nicolás,³ M. L. Ferreira,³ and M. B. Fernández van Raap^{1,a)}

¹Instituto de Física La Plata (IFLP- CONICET), Departamento de Física, Facultad de Ciencias Exactas, Universidad Nacional de La Plata (UNLP), c.c. 67, 1900 La Plata, Argentina ²Instituto de Física "Gleb Wataghin" (IFGW), Universidade Estadual de Campinas, Brazil ³Instituto de Química del Sur (INQUISUR, UNS-CONICET), Avda. Alem 1253, B8000CPB, Bahía Blanca, Argentina

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In this work, the issue on whether dynamic magnetic properties of polydispersed magnetic colloids modeled using physical magnitudes derived from quasi-static magnetic measurement can be extrapolated to analyze specific absorption rate data acquired at high amplitudes and frequencies of excitation fields is addressed. To this end, we have analyzed two colloids of magnetite nanoparticles coated with oleic acid and chitosan in water displaying, under a radiofrequency field, high and low specific heat power release. Both colloids are alike in terms of liquid carrier, surfactant and magnetic phase composition but differ on the nanoparticle structuring. The colloid displaying low specific dissipation consists of spaced magnetic nanoparticles of mean size around 4.8 nm inside a large chitosan particle of 52.5 nm. The one displaying high specific dissipation consists of clusters of magnetic nanoparticles of mean size around 9.7 nm inside a chitosan particle of 48.6 nm. The experimental evaluation of Néel and Brown relaxation times ($\sim 10^{-10}$ s and 10^{-4} s, respectively) indicate that the nanoparticles in both colloids magnetically relax by Néel mechanism. The isothermal magnetization curves analysis for this mechanism show that the magnetic nanoparticles behave in the interacting superparamagnetic regime. The specific absorption rates were determined calorimetrically at 260 kHz and up to 52 kA/m and were well modeled within linear response theory using the anisotropy density energy retrieved from quasi-static magnetic measurement, validating their use to predict heating ability of a given polydispersed particle suspension. Our findings provide new insight in the validity of quasi-static magnetic characterization to analyze the high frequency behavior of polydispersed colloids within the framework of the linear response and Wohlfarth theories and indicate that dipolar interactions play a key role being their strength larger for the colloid displaying higher dissipation, i.e., improving the heating efficiency of the nanoparticles for magnetic fluid hyperthermia. © 2014 AIP Publishing LLC. [http://dx.doi.org/10.1063/1.4862647]

I. INTRODUCTION

Magnetic hyperthermia is nowadays being settled as a therapeutic for cancer treatment. Abnormally high temperature is induced in a tumor using magnetic nanoparticles (MNPs) and radiofrequency (RF) field to selectively kill tumor cells. After the first pioneer work on Limph Node,¹ latter resumed by Jordan's group,² much effort has been devoted to enhance nanoparticles properties preserving biological requirements (i.e., non-toxicity and stability in suspension in biological medium). MNPs displaying large specific heating, usually characterized by the specific absorption rate (SAR), are desired to lower dose for treatment.³ Although large number of particles and coatings has been studied, iron oxides (magnetite and maghemite) coated with organic molecules dispersed in water are widely accepted as the colloid displaying the best biocompatibility. Most common used surfactants include non saturated monocarboxylic

acids,⁴ such as oleic, lauric, myristic and stearic acids, among others.⁵ Polymeric coating also appears as a suitable alternative, especially Chitosan which is a natural biodegradable polymer with a high density of amine groups in their structure that may be protonated depending on pH to achieve stability in suspension. Colloids formed by MNPs coated with polyethylene glycol,⁶ polyethylene glycol fumarate, polyvinyl alcohol, acrylate-based coatings, dextran-based coatings, synthetic polyesters,⁷ alginate and polyethylenimine, have been reported.⁸ Besides the achieved stability, other main difference between monocarboxylic acids and polymeric coatings, of importance for magnetic hyperthermia application, resides on coated nanoparticles hydrodynamic size. In the former case, a small molecule is bound to the MNP surface then the overall nanoparticle size increase is small. In the latter a large macromolecule is bonded resulting in a large overall nanoparticle size and consequently in a large hydrodynamic size.

Although much knowledge and many assays have been used to synthesize, test and characterize various MNPs type for magnetic hyperthermia, many questions remain still open. First, it is unclear if rather larger dissipation would be

^{a)}Author to whom correspondence should be addressed. Electronic mail: raap@fisica.unlp.edu.ar. Phone: +54 221 4246062 x 257. Fax: +54 221 4236335.

obtained by Néel mechanism using particles of large hydrodynamic size or by Brown mechanism using particles of smaller sizes. Second, the influence of interparticle dipolar interactions on SAR is not clearly understood either. Experimental results indicating that increasing dipolar interactions increase and decrease the SAR have been published. Increases in dipolar interaction strength were attempted in $30-33 \text{ nm MFe}_2O_4$ (M = Mg, Ni, Co) spinel ferrite nanoparticles by increasing the size distribution and it was shown that SAR, and minor loops area increase with increasing distribution standard deviation (s.d).9 However, increasing s.d effects arising from size dependence cannot be discriminated from dipolar effects. On the other hand, experimental studies on monodispersed maghemite nanoparticles of 12 nm magnetic size and 14 nm hydrodynamic size evidence that an increase in concentration leads to a decrease in MNP's heating capacity¹⁰ and this result was later supported by Monte Carlo simulations.¹¹ More recently, high frequency hysteresis loops measurements of various monodispersed Fe_xC_y (core-shell) and FeCo samples displaying same size but different anisotropy has been reported.¹² The authors showed that an increase in dipolar interaction strength (achieved by increasing anisotropy) improve heating efficiency in a colloidal solution when nanoparticles are free to arrange in chains and columns. We have recently showed for aqueous stable suspensions of magnetite MNPs coated with a monocarboxyl acid (citric acid), that interparticle interactions are relevant influencing the magnitude of Neél relaxation time and then dissipation mechanism selection.¹³

For single-domain MNP, coherent magnetization reversal is due to Néel and/or Brown processes. The magnetization lags behind the field when the nanoparticle magnetic moment switches (Neél) or when the particle itself rotates in the medium (Brown). Theoretical treatments^{14,15} of these mechanisms for magnetic hyperthermia application are based on linear response¹⁶ and Stoner Wohlfarth theories.¹⁷ Briefly, when a randomly oriented assembly of uniaxial single-domain particles of volume V and specific saturation magnetization M_s is subjected to a RF field of frequency f and amplitude H_0 , SAR is given by

$$SAR = \mu_0 \pi f H_o^2 \chi''(f, \tau), \tag{1}$$

where μ_0 is the permeability of free space $(4\pi \times 10^{-7} \text{ H/m})$ and $\chi''(f)$ is the out-of-phase susceptibility given by

$$\chi''(f) = \frac{2\pi f \tau}{1 + (2\pi f \tau)^2} \chi_0$$
(2)

and $\chi_0 \cong \frac{\mu_0 \rho M_s^2 V}{3k_B T}$ is the d.c. mass susceptibility. The dynamic of the particle's dipole moment is governed by the effective relaxation time τ given by $\tau^{-1} = \tau_N^{-1} + \tau_B^{-1}$ when independency between modes is assumed. The relaxation times depend on medium viscosity (η) and hydrodynamic volume V_H through Brown component given by $\tau_B = \frac{3\eta V_H}{k_B T}$, and on magnetic volume, effective anisotropy energy density K_e and attempt time $\tau_0 \sim 10^{-10}$ s (Ref. 18) through Néel given by $\tau_N = \tau_0 \exp(E_a/k_B T)$. The energy barrier E_a of a double-well potential is related to K_eV . Equation (1) holds for single-size isolated particles and its applicability has been tested for nearly monodispersed nanoparticles, relative s.d. lower than 0.18, produced by thermal decomposition.¹⁹ However, the more complex case of a polydisperse assembly of nanoparticles, commonly obtained from coprecipitation methods, has not been fully studied. Recently, we show for cubic-like shaped $Zn_xFe_{3-x}O_4$ particles (0.0 < x < 0.5) that normalized SAR data of a large set of colloids merge into a single curve consistent with the frequency factor $\chi''(f)/\chi_0$, but the SAR experimental values were not theoretically reproduced.²⁰ The importance of MNPs size distribution on SAR has been pointed out theoretically^{14,21} and experimentally.^{22,23} Theoretical calculation has indicated that the useful H_0 and f values strongly depend on size distribution parameters²² and predicted a reduction close to 70% in the SAR when size polydispersity increase from monodispersed (lognorm deviation $\sigma = 0$) to $\sigma = 0.4$ at optimum mean size.^{14,21} Salas et al.²³ have measured at 18 nm magnetite coated with DMSA a SAR value three times lower when the size distribution s. d. was duplicated.

Here, aqueous colloids displaying a wide magnetic-core size distribution are used to test the applicability and validity of Eq. (1) to predict heating ability of polydispersed suspensions of particle when their dynamics are probed by isothermal magnetization, i.e., using quasi static measurements. The dispersed phase is magnetite coated with oleic acid and chitosan. We first used several structural techniques like FTIR, TEM, AFM, SAXS, and DLS to completely characterize the nanoparticles and their aqueous suspensions, and DC magnetometry to determine magnetic properties. Once the structural and magnetic parameters are known, we analyze SAR theoretical behavior.

Articles regarding magnetite dispersion containing chitosan synthesis protocols and their physicochemical and magnetic characterization have been reported.^{24,25} However, biomedical applications of these materials were not evaluated. One article deals with the preparation of stable colloidal magnetic dispersions composed of magnetite and chitosan by two steps crosslinking of polymeric networks²⁴ resulting in Fe₃O₄—chitosan magnetic composite nanoparticles with a core—shell structure and a diameter of 30–50 nm. The inductive heating for hyperthermia applications was accomplished but no quantitative evaluation of SAR or any analysis to relate heating efficiency to magnetic and structural properties were attempted.

II. MATERIALS AND METHODS

A. Synthesis of stable aqueous suspension of Fe_3O_4 coated with oleic acid and chitosan

All chemicals used in this work are of analytical grade. Oleic acid (OA) was from Anedra, Chitosan (CS) from Chitoclear, by Primex (Iceland) and solvents from Dorwill (Argentina, SA).

The colloids were prepared by co-precipitation method²⁶ commonly used to prepare magnetite, with some modifications.¹⁰ The same amount of CS (450 mg) was used to hydrophilize 300 mg of oleic-acid coated magnetite core (OA-core)

nanoparticles (3/2 of CS/OA-core w/w) previously synthesized with different mass ratio of oleic acid to magnetite (1/1 and 4/1), In brief, $FeCl_3 \cdot 6H_2O$ (0.0121 mol of Fe^{3+}) and FeSO₄ (6.46×10^{-3} mol Fe²⁺) were dissolved in 100 ml of distilled water. The corresponding proportion of OA was incorporated to the mixture. Then, 25 ml of NaOH 5 M were added to precipitate the oxide. This addition was performed at controlled rate to minimize the magnetite aggregation. The mixture was allowed to complete the magnetite formation during 30 min. The obtained dark solution was decanted and supernatant was extracted. Then, bidistilled water was added to remove the impurities by filtration. These washing cycles were repeated three times to ensure complete elimination of the ionic moieties. The solid was dried in an oven at 45 °C overnight under vacuum. In a second step, the nanoprecipitation method was adopted to coat OA-core with CS.²⁷ In brief about 300 mg of OA-core were dispersed in acetone under ultrasound treatment during 15 min. Then, the corresponding volume of a solution of 9.73 mg/ml of CS in acetic acid (50%) was incorporated. The solid MNPs decanted immediately. The supernatant was removed and the solid re-dispersed in distilled water. At these conditions, a dispersion stable during almost three months was formed. A deeper discussion on synthesis protocol and physicochemical characterization can be found elsewhere.²⁸ From these procedures carried out varying synthesis parameters, we choose two colloids displaying high and low SAR, here in after labeled as HD and LD, respectively.

B. Structural and physical-chemical properties

Fourier Transform Infra Red (FTIR) Spectroscopy was used to ensure polymeric coating accomplishment. A FTIR (DRIFTS) Thermo Scientific Nicolet 6700 spectrometer was used for recording spectra in the range 4000–400 cm⁻¹. A few milligrams of the samples (roughly 10–20 mg) were mixed in a mortar manually with near 50 mg of KBr powder. The mixture was placed in the sample unit of FTIR-DRIFTS.

For AFM analysis the colloids were diluted to 1/10000 from their original concentration [x] to ensure single CS particle observation and were dropped on newly peeled muscovite mica sheet V-1 grade (SPI Supplies) and dried in an oven at 40 °C. The samples were imaged in air and at room temperature using standard semicontact mode in a NT-MDT Solver Pro microscope equipped with a APPNANO-ACTA silicon probe consistent for high resolution imaging. The rectangular cantilever of 40 N/m force constant and 281.8 kHz resonant frequency holds a sharp tip of 6 nm curvature radius. Transmission electron microscopy measurements (TEM) were carried out with a 300 keV JEM 3010 microscope to obtain particle size distribution by drying the aqueous dispersions on carbon coated copper grids.

XRD was carried out for phase identification. The patterns were recorded at room temperature using a PHILIPS PW1710 diffraction spectrometer. Small-angle X-ray scattering (SAXS) was performed to determine nanoparticle structuring in suspension using synchrotron radiation. The data were obtained at a wavelength of $\lambda = 1.822$ Å $(\varepsilon = 6.804 \text{ keV})$, using MAR 165 CCD-detector and corrected for absorption, background due to solvent density fluctuation scattering prior to the fitting procedure. The momentum transfer vector $q = 4\pi \sin(\theta)/\lambda$, being 2θ the scattering angle, range from 0.06 to $2.0 \,\mathrm{nm}^{-1}$ was achieved by measuring with two sample-detector distances of 975 and 1977 mm. Absolute scale to express scattered intensity as differential cross section (cm⁻¹) was determined using water as a secondary standard. For measurement the as prepared colloids were sonicated and injected in between two clear Ruby Mica Discs, $25 \,\mu m$ thick each, of the liquid sample holder.

Nanoparticles hydrodynamic diameter (D_H) was retrieved from Dynamical Light Scattering (DLS). Colloidal system stability is analyzed from Z-potential (ζ) obtained from Laser Doppler Electrophoresis measurements. A Malvern Zetasizer nano ZS90 device was employed. Dispersions of magnetic MNPs in bidistilled water were prepared in a proportion of 0.1 mg MNPs/mL at pH = 5.5 to measure ζ . The same dispersions were ultrasonicated during 1 h to measure the average particle sizes.

Atomic Emission Spectroscopy with Induced Plasma Coupled, (ICP-AES), simultaneous high resolution Shimadzu 9000 was utilized to determine the MNP's composition in terms of total Fe content. Alternatively, a common and standardized chemical volumetric analysis, with $K_2Cr_2O_7$ used as titrant, was used for colloid concentration [x] determination, expressed as magnetite mass per solvent volume with an accuracy of 2%. [x] values were used to calculate SAR values from calorimetric data and to normalize magnetic measurements dividing by magnetite mass concentration [m] = [x]V/M_p. The nanoparticle mass (M_p) was obtained drying V = 100 μ l of colloid at 37 °C during 24 h. Values of [x] and [m] are reported in Table I.

C. Magnetic measurements

MNP's magnetic response was measured by DC. Magnetometry, understood as quasi-static magnetic measurements comparing with the condition of magnetic hyperthermia

TABLE I. Nanoparticles and colloids composition from nominal precursors proportions: OA/core is the mass radio of oleic acid to magnetite core and CS/core-OA the mass radio of chitosan to OA coated magnetite nanoparticle (OA/core) and from ICP and titration: [x] is the concentration expressed as Fe₃O₄ mass per ml of water and [m] the mass of Fe₃O₄ over nanoparticle mass. Magnetic colloids zeta potential (ζ) before chitosan coating (i.e., OA/core) and of final colloid after CS coating.

Label	OA/core	CS/core-OA	[x] mg/mL	[m] mg _{Fe3O4} /mg _{NP}	$\zeta_{OA/core} (mV)$	$\zeta_{\rm CS/OA/core}~({\rm mV})$
HD	1/1	3/2	3.19	0.17	-20.8	+25.7
LD	4/1	3/2	1.73	0.07	-31.3	+39.8

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experiments, on dried powder samples and by calorimetric measurements on colloidal samples under RF fields.

Specific magnetization (*M*) as a function of applied magnetic field (*H*) at room temperature was obtained using vibrating sample magnetometer (VSM) LakeShore 7404 operated with maximum applied fields $\mu_0 H_{max} = 1.5$ T and using a superconducting quantum (SQUID) magnetometer with $\mu_0 H_{max} = 2.5$ T. The later was also used to achieve hysteretic magnetization loops at 10, 47, 100, 170, 240, and 290 K and *M* dependence with T under zero field cool (ZFC) and field cool (FC) protocols with $H_{DC} = 8$ kA/m.

Time-dependent calorimetric experiments for determining SAR were conducted by exposing 1 mL of the aqueous suspensions, hold in a clear glass Dewar, to 260 kHz and field amplitude varying in the range from 16 to 52 kA/m. Field generator consists of a resonant RLC circuit Hüttinger (2.5/300) with a water refrigerated 6 turns coil of 2.5 cm diameter. The temperature was sensed during the experiment with an optical fiber sensor placed in the sample centre. The sensor was connected to a calibrated signal conditioner (Neoptix) of an accuracy of \pm 0.1 °C. Colloid temperature was kept below 50 °C in order to minimize solvent evaporation and prevent its destabilization.

The SAR values were calculated from the initial slope $\partial T/\partial t$ of experimental heating curves as

$$SAR = \frac{C}{[x]} \frac{\partial T}{\partial t},$$
 (3)

where C is the volumetric heat capacity of the solvent $(4.18 \text{ J/cm}^3 \text{ K})$.

III. RESULTS AND DISCUSSIONS

XRD patterns of both colloids (not shown) display welldefined reflection lines belonging to cubic spinel structure (JCPDS #19-629) indicating that MNPs are single phase and that neither CS nor OA incorporations alter the crystalline magnetic core.

Figure 1 displays FTIR-DRIFTS spectra of HD and LD colloids containing different OA and CS proportions and pure polymer. Characteristic peaks of CS could be distinguished in the spectra, such as those located at around 3440 cm^{-1} , ascribed to –OH group, at 1620 cm^{-1} assigned to N–H bending vibration and the peak of 1392 cm^{-1} associated to –C–O stretching of primary alcoholic group in CS.²⁹ Magnetite contribution, observed in both spectra, is evidenced by a band located at almost 570 cm^{-1} assigned to Fe-O bonds. These results confirm integration of stable magnetic colloids.

TEM micrographies of LD and HD nanoparticles are shown in Figure 2, spheroid-like forms are distinguished and atomic planes are resolved indicating good crystallinity. Figures 2(a) and 2(b) display imaging of HD sample, where MNPs seems to be composed of several magnetite cores recovered by a polymeric layer. A large polymeric particle containing several magnetite cores is clearly seen in Figure 2(a) and the polymeric coating can also be observed. As a

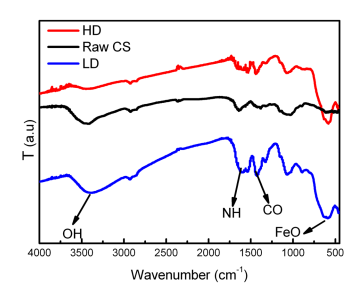


FIG. 1. FTIR-DRIFTS spectra of HD and LD magnetic colloids and raw chitosan. Transmittance (T) as a function of the wavenumber (cm⁻¹). The spectra were shifted for clarity.

difference, LD sample imaging in Figures 2(d) and 2(e) shows almost individual MNPs where the CS layer seems to be acting as spacer between nanoparticles. From the images size histograms were constructed by counting MNPs from several micrographies of each sample. Mean size and s.d. of 9.7 nm ($\sigma_d = 4.5$ nm) and 4.8 nm ($\sigma_d = 2.1$ nm) were obtained for HD and LD samples by fitting the histograms with Gaussian functions (see Figures 2(c) and 2(f)). LD sample present lower sizes and narrower size distribution than HD sample, i.e., HD sample is more polydisperse than LD, consistently with the larger amount of OA used to synthesized LD. OA is used to limit growing after nucleation during synthesis protocol. On a qualitative analysis we observed that, although both samples were synthesized by the same route and dried for TEM measurements under the same conditions, HD sample presents some aggregates which may be due to interparticle interactions acting during dry treatment or to a permanent morphology of polymeric particles containing several particles as can be clearly seen in Figure 2(a). As already mentioned, HD sample displays on mean value larger sizes than LD, i.e., larger magnetic moment, resulting in a stronger dipolar interaction between nanoparticles favoring particles aggregation.

Surface properties were estimated by ζ measurements recorded on colloid precursors (OA-core nanoparticles) and are listed in the Table I for LD and HD suspension. Dispersions of OA-core in bidistilled water (pH around 5.5) render negative ζ with magnitudes between -20 mV and -31 mV. These values correlate with OA double layer and are in agreement with previous work.²⁷ CS incorporation causes positively charge surfaces attributed to surface exposed protonated amine groups on CS moieties. Z potential is considered as a measure of colloid stability. The recorded ζ magnitude between +25 mV and +39 mV correlates with stable suspensions.

According to the mechanism proposed in an earlier work,²⁸ complexes acetate–CS are formed from acetic acid dissolving CS during nanoprecipitation step, and coordinate

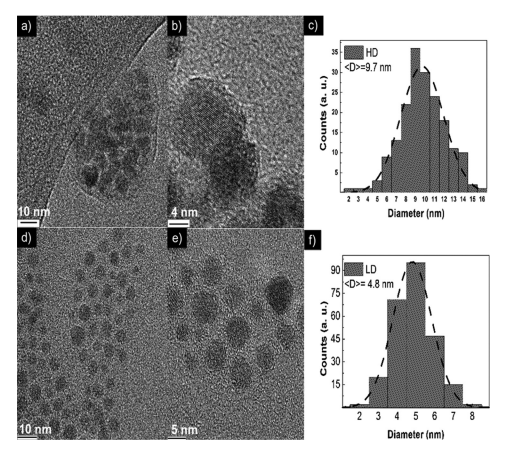


FIG. 2. Representative TEM images of HD sample are in (a) and (b); of LD sample in (d) and (f). Size histograms obtained by sampling 182 nanoparticles are shown in (c) for HD and in (f) for LD considering 250 nanoparticles. The dashed lines stand for a fitted Gaussian function.

with surface Fe atoms of magnetite, leading to surface exposed NH_3^+ groups. Therefore, colloid stabilization takes place by electrostatic repulsions.

A charged particle moving in a solvent drags a layer of fluid molecules and counter ions, resulting in an entity with a quite large hydrodynamic diameter D_H . In fact, measured D_H values are larger than the overall nanoparticle size including coating. Mean values of number-average hydrodynamic sizes (D_H) measured in bidistilled water are reported in Table II. LD colloid sample presents D_H of 134.6 nm ($\sigma_{D_H} = 18.8$ nm) while HD exhibits a lower value of 57 nm ($\sigma_{D_H} = 7.7$ nm).

Scattering techniques like DLS and SAXS can be carried out in the suspension but only SAXS provides information on the internal structure of the scattering objects. Figure 3 shows the one-dimensional (1-D) SAXS pattern for the nanoparticles circularly averaged from the 2-D measured pattern. The 2-D pattern (not shown here) clearly revealed isotropic scattering, which is consistent with spherical objects scattering. The scattering curves shown in Figure 3 for HD and LD colloids are quite different. Two knees are observed for LD (see arrows in Fig. 3), indicating objects scattering at different length scales, while HD displays a

more smooth decreasing q-behavior. HD pattern was well fitted considering spherical form factor weighted with a LogNormal size distribution (median $log(d_s)$ and s.d. σ_s) convoluted with the analytic form of the structure function $S(q)^{30,31}$ derived for a fractal model of aggregation as described in Ref. 32. In this model, the power law form (q^{-d_f}) of the scattering function is limited by finite cluster size ξ and by the primary particle finite size d_s . SAXS analysis is indicating that the morphology seen by TEM for HD sample corresponds to a permanent aggregation with clusters of maximum size $\xi = 28.2$ nm inside larger CS particles of 48.6 nm size and not to a structuring produced during drying process. On the other hand, we have considered LD scattering curve as the superposition of the scattering from a dispersion of individual small magnetite nanoparticles which is observed at $q \ge 0.46 \text{ nm}^{-1}$ and the scattering from a large CS particle which dominates scattering at q values lower than 0.46 nm^{-1} , being the magnetite nanoparticle and CS the two objects scattering at different length scale. The differential cross sections in the large q-range was fitted in the same way as HD sample but setting S(q) = 1, which means no aggregation while in the low q-range with the unified equation of

TABLE II. Structural properties: $\langle d \rangle$ is the mean magnetite nanoparticle size deduced from TEM histograms and σ_d the corresponding standard deviation (s.d); D_H and σ_{D_H} are the mean hydrodynamic size and s.d. from DLS data; SAXS: d_s , σ_s mean magnetite size and s.d, d_f fractal exponent, ξ cluster size and d_{CS} chitosan particle size from SAXS data analysis.

Label	$\langle d \rangle$ (nm)	$\sigma_d (\mathrm{nm})$	$D_H(\mathrm{nm})$	σ_{D_H} (nm)	d_s (nm)	$\sigma_{\rm s}({\rm nm})$	d_f	ξ (nm)	d_{CS} (nm)
HD	9.7	4.5	56.9	7.7	6.0	3.1	2.7	28.2	48.6
LD	4.8	2.1	134.6	18.8	3.4	1.3			52.5

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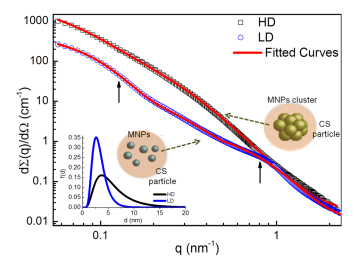


FIG. 3. 1D-SAXS patterns expressed as differential cross section $d\Sigma(q)/d\Omega$ against scattering vector q for HD and LD colloids. Continuous lines stands for fitted curves using the models explained in the text. For each pattern, a schematic representation of MNPs structuring inside chitosan particles is plotted. *Inset*: f(d) is the lognorm magnetite core size distributions obtained from the fits.

Beaucage³³ for one structural level. This fit is consistent with a CS particle of 52.5 nm. SAXS analysis in both cases is in agreement with TEM results and provides reliable values for CS coating and nanoparticles structuring. From this analysis, the overall size of the coated nanoparticles are 48.6 nm and 52.5 nm for HD and LD colloids. D_H size, obtained in the suspension with DLS is calculated from the diffusion coefficient using the Stokes-Einstein and corresponds to the size of the dynamic hydrated particle. This value is close to the overall size for HD and 3 times larger for LD.

AFM images of LD and HD samples are shown in Figure 4, and some individual particles are labeled with their height (size). The topography clearly indicates spherical particles and sizes bellow 50 nm for both samples in agreement with SAXS results. The large shell of CS in LD nanoparticles is probably much less compact then drags a quite large layer of water and counterions.

This complete structural study involving four experimental techniques confidently indicates that in HD colloids each CS particle contain aggregates of several MNPs while in LD colloids the MNPs are spaced inside a large CS particle. A schematic draw is included in Figure 3.

Next, magnetic properties are evaluated using DC magnetometry techniques. The magnetization temperature dependence under ZFC (M_{ZFC}) and FC (M_{FC}) protocols are shown in Figure 5. These curves, displaying magnetization measurements recorded at a low DC field, while temperature is increase at a constant rate, are dominated by magnetic anisotropy and dipolar interactions rather than Zeeman interaction. M_{ZFC} is consistent with a broad energy barrier distribution and peaks at T_m close to 196 K and 71 K for HD and LD nanoparticles, respectively. For HD, the ZFC peak is wider and more shifted to higher temperatures consistently with larger size, broader distribution and stronger interactions between MNPs. For LD sample M_{FC} increase with decreasing temperatures below the temperature of

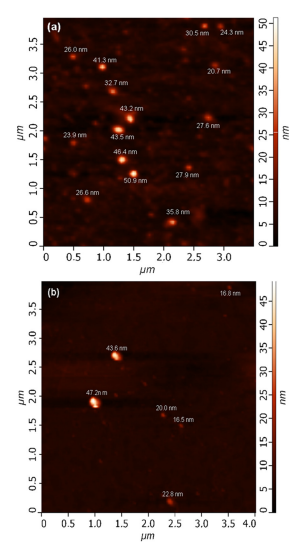


FIG. 4. AFM images (topography) of HD sample (a) and LD sample (b), some nanoparticles are labeled with their height (size).

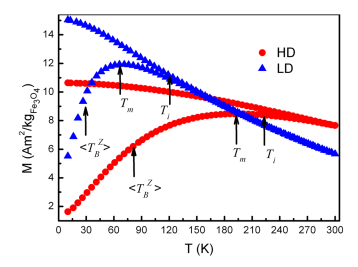


FIG. 5. Specific Magnetization (*M*) against temperature (*T*) measured under zero-field-cooling (ZFC) and field-cooling (FC) protocols with $H_{DC} = 8 \text{ kA/m}$ and heating rate of 5 k/min. $\langle T_B^Z \rangle$ indicates mean blocking temperature obtained from $d(M_{FC} - M_{ZFC})/dT$, T_i the irreversibility temperature and T_m the maximum of $M_{ZFC}(T)$ curve.

TABLE III. Magnetic properties: $\langle \mu \rangle$ is the nanoparticle mean magnetic moment, M_s specific saturation magnetization of magnetite nanoparticle at room temperature, K_e effective anisotropy constant and $\langle T_B^H \rangle$ deduced from coercive field temperature behavior and T_m and $\langle T_B^B \rangle$ are the maximum and mean blocking temperatures deduced from ZFC/FC magnetization data. Specific absorption rate SAR measured at 260 kHz and 52 kA/m and expressed per Fe₃O₄ unit mass. SAR_s are the simulated SAR values.

Label	$\left< \mu \right> \left(\mu_{\rm B} \right)$	M_s (Am ² /kg)	$K_e (\text{kJ/m}^3)$	$T_m(\mathbf{K})$	$\left\langle T_{B}^{H}\right\rangle$ (K)	$\langle T_B^Z \rangle$ (K)	SAR (W/g)	SAR_s (W/g)
HD	8053	46	14.7 ₄	196	83 ₁₃	92.9 _{1.7}	82.7 _{1.0}	79
LD	4009	53	11.6 ₉	71	31 ₁₂	29.6 _{0.2}	8.2 _{3.0}	0.1

irreversibility (T_i), while a flat behavior is observed for HD which is again an indication of HD as a more interacting system. In a previous work carried out on magnetite nanoparticles coated with citric acid a correlation between higher SAR values for larger T_m values has been observed.¹³ Mean blocking temperature $\langle T_B^Z \rangle$ is obtained³⁴ from the energy barrier distribution $f(T) \propto d(M_{FC} - M_{ZFC})/dT$. Values are listed in Table III and indicated with arrows in Figure 5.

Hysteresis loops were recorded in both samples at various temperatures below 300 K. Curve shape and behavior are similar for both samples then results are only shown for HD sample in Figure 6. For the data analysis, each sample is considered as an ensemble of MNPs with magnetocrystalline anisotropy axes randomly distributed. For magnetic data analysis, coating only contribute establishing how close the particles can be to each other, then the particles are considered as composed by a magnetic single domain core of volume V and magnetic moment μ surrounded by a dead magnetic layer of thickness δ . The loops were fitted using a displaced Langevin function weighted with a lognorm distribution of magnetic moments $g(\mu)$ plus a high field susceptibility χ_a linear term as it is described in a previous work.¹³ From this fitting procedure M_s , coercive field H_c , mean magnetic moment $\langle \mu \rangle$, s.d. of $ln(\mu)$ variable σ , and χ_a are obtained. Relevant data is shown in Table III. Nanoparticle sizes determined from $g(\mu)$, using $\mu = M_s \rho V$ (mass density $\rho = 5.18 \text{ g/cm}^3$), results in mean sizes smaller than TEM values and SAXS. We used mean values from histograms in Figures 2(c) and 2(f) to model SAR, because TEM is a direct

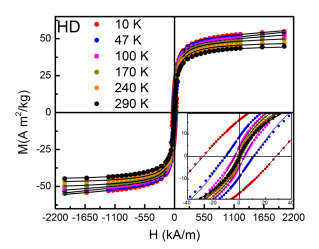


FIG. 6. Hysteresis loops of HD sample recorded at various temperatures. Inset shows a magnification to detail coercive field (H_c) decrease with increasing measurement temperature. Solid lines are the best fits achieved using a displaced Langevin function weighted with a lognorm distribution of magnetic moments plus high field susceptibility.

observation of the magnetite nanoparticle and we were able to count around 200 particles of each sample ensuring reliable statistics.

For both colloids M_s at room temperature is 57% lower that bulk value⁶ (446 kA/m) due to the presence of a dead magnetic layer $\delta \sim 1.8$ nm. This typical M_s lowering is an undesired result because SAR depends on M_s^2 .

Mean magnetic moments $\langle \mu \rangle$ monotonically increases with *T*. This is an unphysical behavior commonly observed in dispersions behaving as interacting superparamagnets (ISP) as explained by Allia *et al.*³⁵ In this regime, the particle moments interact through long-ranged dipolar random forces, whose effect is pictured in terms of a temperature *T** added to measurement temperature in the denominator of the Langevin function argument. Consistently, magnetization data for samples perfectly scale with $H/(T + T^*)$ for $T > T^*$ and ξ/χ increase linearly with T/M_s^2 for both samples, being $\xi = \langle \mu^2 \rangle / \langle \mu \rangle^2$ and χ the low field susceptibility. These plots confirming that the MNPs behave in the ISP regime are shown for HD sample in Figure 7.

The coercive field (H_c) of an assembly of non interacting randomly oriented single-domain MNP exhibiting thermally activated coherent magnetization reversal follows the relation:

$$H_c = 0.48 \frac{2K}{\mu_0 M_s} \left(1 - \left(\frac{T}{\langle T_B^H \rangle} \right)^{1/2} \right) \tag{4}$$

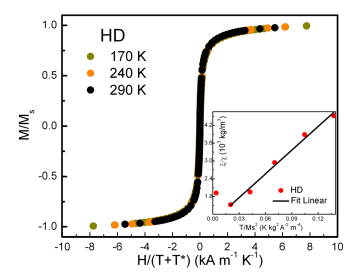


FIG. 7. (a) High temperature scaling regime for hysteresis loops displayed in Figure 6. Inset: plot displaying the linear behavior of $\xi = \langle \mu^2 \rangle / \langle \mu \rangle^2$ (scale second moment of the magnetic moment) over χ (low field DC-susceptibility) against reduce variable T/Ms^2 , with *Ms* the saturation magnetization at the temperature *T*.

below blocking temperature $\langle T_B^H \rangle$,³⁶ where K is the anisotropy constant. For interacting nanoparticles, the anisotropy constant is replaced by the effective anisotropy constant K_e . This constant takes into account the anisotropy energy of each particles and the interparticle interaction energy. H_c values plotted against T are shown in Figure 8 for LD and HD samples. H_c monotonically decrease with T, but the decay is slower for HD sample. The curves were well fitted with $T^{1/2}$ dependence, and $[K_e, \langle T_B^H \rangle]$ were used as fitting parameters, while M_s was fixed at the values measured at 10 K. The values $[14.7 \pm 0.4 \text{ kJ/m}^3, 83 \pm 13 \text{ K}]$ and [11.6] $\pm 0.9 \text{ kJ/m}^3$, $31 \pm 12 \text{ K}$] were obtained for HD and LD, respectively. The fitted $K_{\rm e}$ values are larger than magnetite bulk anisotropy constant³⁷ of 9 kJ/m³ at 300 K. $\langle T_B^H \rangle$ values are in good agreement with those derived from ZFC/FC data analysis. The energy barrier increases by increasing interparticle interactions strength and a shift of blocking temperature towards larger values is observed. Related models have been extensively discussed in Ref. 38. Using a rough estimation for the interparticle interaction strength as $E_d = (K_e - K)V$, HD is 18 times more interacting than LD. This result agrees, considering the dipolar character of the interparticle interaction proportional to the inverse cube of interparticle-distance, with the structural evidence of MNP clusters inside a given CS particle for HD sample where the MNPs are much closer than in the LD case where individuals particles spaced inside a CS particle as happens in LD sample.

Then, SAR values were determined for both colloids using Eq. (3). SAR of HD colloid is 10 times larger than for LD at 260 kHz and 51.76 kA/m. Recently, the largest SAR values, one order of magnitude higher than for isolated single-domain maghemite nanoparticles, have been reported for 50 nm particles composed by several smaller particles.³⁹

The dependence of SAR with field amplitude is shown in Figure 9(a) and the typical heating curves for HD colloid are shown in Figure 9(b). The larger the field amplitude H_0 the faster the achieved heating, as expected from the quadratic dependence of SAR with H_0 (see Eq. (1)). The heating rate of LD colloid was measured at 49.01 and 51.76 kA/m and no significant temperature increase was observed by increasing the field amplitude.

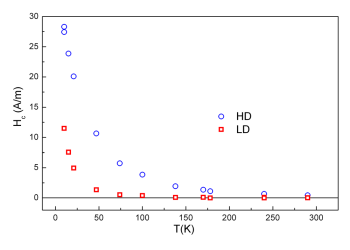


FIG. 8. Coercive field (H_c) dependence on measurement temperature T.

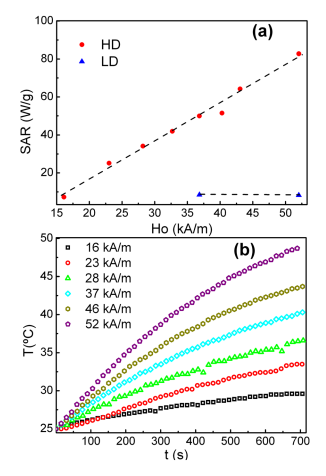


FIG. 9. (a) Specific absorption rate (SAR) derived from calorimetric measurements against excitation field amplitude H_0 obtained under RF frequency of 260 kHz. (b) Heating curves of HD sample.

As already mentioned Eq. (1) holds for single-size isolated particles, i.e., a non-interacting collection of monodispersed nanoparticles. Here, we are analyzing colloids displaying broad size distributions and behaving as interacting superparamagnets, meaning that relaxation times are distributed with a relaxation time distribution function $g(\tau)$ due to size distribution and the energy barrier may be changed by interparticles interaction. When the MNP assembly is monodispersed, the whole set of particle are fluctuating. Then, within a mean field theory, the field produce by dipolar interactions is a fluctuating field which average to zero. For broad size dispersion, the smaller particles are fluctuating, while the larger particles are blocked. The larger particles produce a static mean dipolar-field, increasing the energy-barrier height. In the present analysis, the dipolar interaction is included as a large effective anisotropy constant K_e derived from DC magnetometry data, i.e., this K_e value corresponds to MNPs inside the assembly. The obtained K_e magnitudes are in concordance with the structuring of the MNPs inside the CS particle.

Then SAR is given by $SAR = \mu_0 \pi f H_o^2 \int \chi''(f, \tau) g(\tau) d\tau$ and making explicit τ size dependence is given by

$$SAR = \mu_0 \pi f H_o^2 \frac{\int_0^\infty \chi''(D) V p_N(D) dD}{\int_0^\infty V p_N(D) dD},$$
(5)

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where $p_N(D)dD$ is the number of spherical particles with diameters between D and D + dD. The main MNPs properties listed in Tables II and III were used to address the goodness of Eq. (5) to predict heating efficiency under RF fields of polydispersed MNPs of large hydrodynamic volume. Néel and Brown relaxation times, calculated using data in Tables II and III, are around 10^{-10} s and 10^{-4} s, respectively. This large difference assures that the nanoparticles in both colloids magnetically relax by Néel mechanism and the possibility of a fraction of particles dissipating by Neél and the complement by Brown is excluded. Equation (5) was evaluated, for both colloids at $H_0 = 52 \text{ kA/m}$ and f = 260 kHz, using the size distribution function $p_N(D)$ retrieved from TEM histograms jointly with V_H from DLS and K_e from DC magnetometry (i.e., isothermal magnetization), $\eta = 10^{-3}$ Pa.s and $\tau_0 = 10^{-10}$ s. The attempt time value was choose from the detail analysis of Dormann et al.⁴⁰ who showed that for interacting nanoparticles the dependence of $log(\tau)$ against $1/T_B$ departs from Arrhenius law and τ asymptotically goes to $\tau_0 = 10^{-10}$ s. By this way, Eq. (5) gives SAR values of 78.8 and 0.1 W/g for HD and LD, respectively. The agreement with calorimetric SAR data (also listed in Table III) is very good for HD sample not for LD. However, this calculation results in low values for the low dissipating sample and the correct high values for high dissipating sample, i.e., the right behaviour is achieved.

Our analysis suggest that Eq. (5) combined with a complete set of structural and magnetic parameter $[\langle d \rangle, \sigma_d, \langle D_H \rangle, \sigma_{DH}, K_e, M_s]$ reproduce the SAR behavior of colloids displaying large size polydispersity. This fact validates the use of Eq. (5) to predict the colloids performance under other RF field parameters and to draw tips to improve nanoparticles design for biomedical colloids. It is worth mentioning that our analysis differs from the one carried out in Ref. 41 where the relaxations times τ_N and τ_B were separately averaged using size and hydrodynamic size distribution, respectively, and replaced in Eq. (1), resulting in SAR calculated values one order of magnitude lower than the experimental ones.

Simulations of τ , τ_B and τ_N against D were obtained using parameters listed in Tables II and III and are shown for HD colloid in Figure 10(a). The crossover from Neél to Brown relaxation mechanism appears at D = 20 nm for HD and D = 22 nm for LD. Then, the whole dispersion of MNP composing the colloids studied here dissipates through Neél mechanism consistently with the large hydrodynamic volumes. In Figure 10(b), simulations of the out of phase susceptibility dependence on MNPs size $\gamma''(\tau(D))$ at fixed frequency of 260 kHz and amplitude of 52 kA/m and $p_N(D)$ are plotted together for both samples. It is clearly seen that only when $p_N(D)$ superimposed to a non null value of $\chi''(\tau(D))$ the sample dissipates. $\chi''(\tau(D))$ determines the fraction of dissipating MNP at a given frequency. For LD sample $\chi''(\tau(D))$ is not overlapping with $p_N(D)$, then almost no heating is expected. For HD colloid, there is some overlapping, only a fraction of particles are dissipating and SAR is 10 times larger. The particles of sizes larger than 6 nm represent 34% of the magnetite mass in HD colloid and generate 98% of dissipated energy. One way to improved SAR of HD

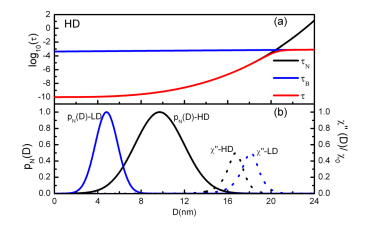


FIG. 10. (a) displays size dependence of τ , τ_N and τ_B simulated for HD colloid using magnetic nanoparticle size from TEM, hydrodynamic volumes from DLS, effective anisotropy constant K_e form DC-magnetometry, $\eta = 10^{-3}$ Pa.s and $\tau_0 = 10^{-10}$ s. (b) displays Gaussian size distribution $(p_N(D))$ against radii (*D*) as derived from TEM data (filled lines) and simulation of the out of phase susceptibility $\chi''(D)$ normalized by DC-susceptibility χ_0 . The simulations were performed using $\tau(D)$ values plotted in (a) and other data listed in Tables II and III.

colloid would be to separate the nanoparticles on the basis of their dissipative properties. For instance, selecting the particles with radii larger than 6 nm, SAR would be 225 W/g. Other way would be raising the field frequency because the maximum of χ'' will shift to lower sizes. For instances, Eq. (5) evaluated at $f = 500 \,\text{kHz}$ predict an increase to 224 W/g for HD colloid. Then, this analysis indicates that for colloids of MNPs with wide size distribution the largest dissipation would be achieved for an assembly of nanoparticles of mean size coincident with the maximum of $\chi''(D)$. Provided this coincidence condition the narrow the size distribution the largest the SAR in agreement with previous work.^{14,21,23} Other way, would be increasing the energy barrier height, i.e., increasing the phase shift between excitation wave and magnetization, which may be achieved in a controlled way by increasing interparticle interactions strength, for example, increasing the amount of particles inside the CS particle. But it should be kept in mind that increasing the energy barrier height the crossover in Fig. 10(a) will shift to lower sizes values and a situation corresponding to part of the dispersion (lower core sizes) dissipating by Néel and the rest (larger core sizes) by Brown mechanism could be reached. This analysis also indicates that for a MNP assembly displaying large polydispersity, the size optimization for improving heating efficiency must take into account both relaxation mechanisms and interparticles interactions.

Finally, we stress that an increase in size distribution width results in a decrease in SAR only in colloids in which mean MNP size $\langle d \rangle$ has been tailored to maximize the heat dissipation at the working conditions, i.e., tailored to fulfill the resonance condition $(2\pi f \tau (\langle d \rangle) = 1)$ at working frequency. When the mean size of the dispersion do not fulfill this condition, for instance, the mean size is located at the left of the maximum of $\chi''(D)$ as in Figure 10(b), increasing the polydispersity would result in SAR increase due to a large fraction of particles satisfying resonance condition. This reasoning can explain results of Ref. 9 were an increase in dipolar interaction strength was attempted by increasing size distribution without verifying that their mean sizes were satisfying resonance condition.

IV. SUMMARY AND CONCLUSIONS

We have performed a complete structural and magnetic study of Fe₃O₄ magnetic nanoparticles coated with Oleic acid and chitosan to analyze their heating efficiency under radio frequency field for magnetic hyperthermia applications. We have studied two stable aqueous suspensions of magnetic nanoparticles displaying broad size distributions and differing markedly in the dispersed phase structure. One colloid consists of magnetic nanoparticles of mean size around 4.8 nm spaced inside chitosan particle of 52.5 nm and hydrodynamic size of 135 nm. The other colloid consists of clusters of magnetic nanoparticles of mean size around 9.7 nm inside a chitosan particle of 48.6 nm and hydrodynamic sizes of 57 nm. Our results indicate that both assemblies of suspended magnetic nanoparticles behave in the interacting superparamagnetic regime and the nanoparticles dissipate through Neél relaxation mechanism. It is shown that combining structural data with quasi-static magnetic measurement, the heating ability of a given colloid can be modeled within the framework of the linear response and Wohlfarth theories. This fact validate the use of the dynamic susceptibility modeled using structural and magnetic parameter derived from quasi-static measurements to predict heating ability of a given polydispersed particle suspension of large hydrodynamic sizes. We also conclude that dipolar interaction plays a key role being larger the dissipation for the more interacting nanoparticles assembly.

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