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Article Optimization of Magnetic Cobalt Ferrite Nanoparticles for Magnetic Heating Applications in Biomedical Technology

Diana Zahn ^{1,†}^(b), Joachim Landers ^{2,†}, Marco Diegel ³, Soma Salamon ², Andreas Stihl ^{4,5}, Felix H. Schacher ^{4,5}^(b), Heiko Wende ², Jan Dellith ³ and Silvio Dutz ^{1,3,6,*}^(b)

- ¹ Institute of Biomedical Engineering and Informatics (BMTI), Technische Universität Ilmenau, D-98693 Ilmenau, Germany
- ² Faculty of Physics and Center for Nanointegration Duisburg-Essen (CENIDE), University of Duisburg-Essen, D-47057 Duisburg, Germany
- ³ Leibniz Institute of Photonic Technology (IPHT), D-07745 Jena, Germany
- ⁴ Institute for Organic Chemistry and Macromolecular Chemistry, Friedrich-Schiller-University Jena, D-07743 Jena, Germany
- ⁵ Jena Center for Soft Matter (JSCM), Friedrich-Schiller-University Jena, D-07745 Jena, Germany
- ⁶ Leupold Institute for Applied Natural Sciences (LIAN), Westsächsische Hochschule Zwickau, D-08056 Zwickau, Germany
- Correspondence: silvio.dutz@tu-ilmenau.de
- + These authors contributed equally to this work.

Abstract: Using magnetic nanoparticles for extracorporeal magnetic heating applications in biomedical technology allows higher external field amplitudes and thereby the utilization of particles with higher coercivities (H_C). In this study, we report the synthesis and characterization of high coercivity cobalt ferrite nanoparticles following a wet co-precipitation method. Particles are characterized with magnetometry, X-ray diffraction, Mössbauer spectroscopy, transmission electron microscopy (TEM) and calorimetric measurements for the determination of their specific absorption rate (SAR). In the first series, $Co_x Fe_{3-x}O_4$ particles were synthesized with x = 1 and a structured variation of synthesis conditions, including those of the used atmosphere (O_2 or N_2). In the second series, particles with x = 0 to 1 were synthesized to study the influence of the cobalt fraction on the resulting magnetic and structural properties. Crystallite sizes of the resulting particles ranged between 10 and 18 nm, while maximum coercivities at room temperatures of 60 kA/m for synthesis with O_2 and 37 kA/m for N₂ were reached. Magnetization values at room temperature and 2 T (M_{RT,2T}) up to $60 \text{ Am}^2/\text{kg}$ under N₂ for x = 1 can be achieved. Synthesis parameters that lead to the formation of an additional phase when they exceed specific thresholds have been identified. Based on XRD findings, the direct correlation between high-field magnetization, the fraction of this antiferromagnetic byphase and the estimated transition temperature of this byphase, extracted from the Mössbauer spectroscopy series, we were able to attribute this contribution to akageneite. When varying the cobalt fraction x, a non-monotonous correlation of H_C and x was found, with a linear increase of H_C up to x = 0.8 and a decrease for x > 0.8, while magnetometry and in-field Mössbauer experiments demonstrated a moderate degree of spin canting for all x, yielding high magnetization. SAR values up to 480 W/g (@290 kHz, 69 mT) were measured for immobilized particles with x = 0.3, whit the external field amplitude being the limiting factor due to the high coercivities of our particles.

Keywords: magnetic nanoparticles; cobalt ferrite; magnetic heating

1. Introduction

Magnetic nanoparticles (MNP) have been extensively studied for medical applications, including therapeutic hyperthermia [1–3]; drug targeting where MNP are used as vehicles for the transport of therapeutic agents [4,5]; or imaging that utilizes MNPs as contrast agents [6,7]. Each of the approaches mentioned above requires the particles to be applied



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Copyright: © 2023 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). intracorporeally, leading to restrictions for the particles concerning their biocompatibility, but also for the external magnetic fields used to navigate or heat the particles. Safety restrictions limit an external alternating magnetic field to $H \times f = 5 \cdot 10^9 \text{ Am}^{-1} \text{s}^{-1}$ [8], to prevent unspecific heating of tissue by eddy currents and to ensure patients' comfort.

In addition to these intracorporal medical approaches, MNP are also promising candidates for medically related extracorporeal approaches: their ability to be moved by a magnetic gradient can be used to extract pathogens from specimens and use them as magnetic markers for immunomagnetic separation [9,10]. Furthermore, their mechanism of generating heat can be used extracorporeally, for example as thermal markers in place of the common colorimetric markers used in immunoassays. In this context, external fields with remarkably higher field amplitudes can be used to heat the particles, enabling the use of hard magnetic MNP with higher anisotropy. Such particles are not of interest for intracorporal heating applications, since the required field amplitude to efficiently heat these particles needs to be two or three times the particles' coercivity [11], exceeding the limit for patient safety.

Commonly used iron oxide nanoparticles can be tuned towards a higher anisotropy by changing their shape from spherical to cubic, disc-like or elongated needles [12–15]. Increasing their size while remaining in a single domain state also increases their coercivity [16–18]. Another approach to the synthesis of particles with large coercivities is the introduction of metal atoms such as Zn, Mg, Ba or Co into the crystal lattice to synthesize ferrites. Starting with the inverse spinel magnetite (Fe₃O₄ or [Fe³⁺]_{tetra}[Fe²⁺Fe³⁺]_{octa}O₄), where Fe³⁺ ions are distributed equally between octahedral and tetrahedral sites and Fe²⁺ ions are located on the octahedral site [19], Co²⁺ ions can replace the Fe²⁺ ions. Particles of pure cobalt ferrite, where Fe²⁺ is completely replaced by Co²⁺, are well known to exhibit exceptionally high coercivities of up to several hundred kA/m at room temperature [20,21]. As stated above, an external magnetic field used to heat such particles would need an impractically high field amplitude. Therefore, a partial replacement of Fe²⁺ by Co²⁺ can be beneficial for obtaining materials with the formula Co_xFe_{3-x}O₄, resulting in coercivities and magnetization values between magnetite and cobalt ferrite.

When examining the literature that focuses on this kind of partially substituted magnetite particles, controversial results can be found regarding the correlation between cobalt content x and resulting particle properties. The coercivity of H_C , for example, is found to increase monotonously with cobalt content [22] or to reach a maximum and then decrease for higher cobalt contents towards pure cobalt ferrite (x = 1) [23,24]. In addition, no clear trend for the saturation magnetization (M_S) can be found— M_S can increase with more cobalt [25], decrease [22,26], or show a maximum for intermediate cobalt contents [24,27]. Crystal size and lattice parameters show linear trends with rising cobalt content, but whether this is an increasing or decreasing trend is not consistent, and most likely depends on the degree of oxidation from magnetite to maghemite for the particles with no or small amounts of cobalt [22,23,27,28].

To develop efficient cobalt-substituted magnetite particles that match the application specific requirements by tuning the particle properties, the above-mentioned correlations need to be understood. Therefore, in the first series of experiments we used a co-precipitation method to synthesize a broad range of cobalt ferrite particles (x = 1) with varied synthesis parameters, to evaluate the influence of reaction temperatures, durations, and atmosphere (O₂ or N₂) on the resulting particles. In the second series, we used fixed optimized synthesis parameters to produce partially substituted magnetite particles of $Co_xFe_{3-x}O_4$ with x = 0, 0.3, 0.5, 0.7, 0.8, 0.9, 0.95 and 1, to study the correlations between cobalt content x and the resulting particle properties. The particles were characterized using vibrating sample magnetometry (VSM), X-ray diffraction, transmission electron microscopy, Mössbauer spectroscopy as well as calorimetric measurements for determination of the heating capability.

As the main conclusions, we found thresholds for the reaction temperatures, which, if exceeded, would lead to the formation of crystalline impurity phases—apart from

CoFe₂O₄—and the significant decrease of magnetization. Oxygen-free synthesis under N₂ atmosphere leads to higher magnetization values and decreasing H_C compared with synthesis under air. With increasing amounts of cobalt, H_C at 300 K reaches a maximum at x = 0.8, while M_{RT,2T} ranges from 71 to 77 Am²/kg for 0 < x < 0.9 and slightly decreases. In-field Mössbauer experiments were employed to analyze the particles' magnetic structures, check for the presence of Fe²⁺, determine the Co and Fe ion site occupation and its influence on the particles' magnetization, and to evaluate the correlation between Co content x and the samples' degree of spin frustration.

2. Materials and Methods

2.1. Synthesis of Cobalt Ferrite Nanoparticles

Cobalt ferrite nanoparticles were prepared using a wet co-precipitation method. For this method, iron(III) chloride hexahydrate and cobalt(II) chloride tetrahydrate are dissolved in 120 mL deionized water in a two-neck glass flask under stirring in a water bath, heating the salt solution to the addition temperature (T_{add}). An amount of 12 mL of a 3M sodium hydroxide solution is added with a syringe pump over a specified duration, hereafter called duration of NaOH addition or d_{add}. The flask is then transferred to an oil bath, heating the solution to a higher temperature named end temperature or T_{end} and kept in this bath under stirring for a defined time, named duration of reaction or dreact. For a structured analysis of the influence of the named synthesis parameters on the resulting particle characteristics, for the first part of this study (hereafter named series 1) we varied the parameters over a broad range. Each set of parameters was used for the synthesis of three particle batches to enable a basic statistical analysis of the results. Additionally, all parameter sets were reproduced under the exclusion of oxygen by washing the used solutions with nitrogen prior to the synthesis and keeping the flask under nitrogen atmosphere throughout the whole reaction. Table 1 shows the used settings for the four parameters T_{add}, d_{add}, T_{end}, d_{react}. When one of the parameters is varied, the remaining parameters are kept constant at the value written in bold in the table. For the parameter "duration of addition", 0 min means an addition of the total volume of NaOH solution as a single bolus over a period of seconds. For evaluation of the synthesis parameters, a stochiometric cobalt-to-iron ratio was used in this series of particles to synthesize completely substituted cobalt ferrite nanoparticles with x = 1 and a ratio of Co^{2+} ions to Fe^{3+} ions of 1:2. When needed, samples are named in the following results as O₂/N₂_d_{add}_T_{add}_T_{end}_d_{react}.

Table 1. Varied synthesis parameters and used values. Bold type values are declared as standard and used during the variation of other parameters.

Parameter	Abbreviation	Unit		Used	Values	
Duration of NaOH addition	d _{add}	min	0	2	4	8
Addition temperature	T _{add}	°C	25	40	65	88
End temperature	T _{end}	°C	70	85	97	
Duration of reaction	d _{react}	min	30	60	90	

As a second part of the study (hereafter named series 2), the amount of cobalt was altered and iron(II) chloride tetrahydrate was introduced to the synthesis. By varying the ratio of Co^{2+} ions to Fe^{2+} ions and while keeping the amount of Fe^{3+} ions constant, a series of $Co_xFe_{3-x}O_4$ particles with varying cobalt content was synthesized. The percentage of Fe^{2+} ions replaced by Co^{2+} ions is named cobalt fraction x in the following and was evaluated in the range from 0 (no Co^{2+} ions) to 1 (Fe^{2+} is replaced completely by Co^{2+}) via the intermediate steps x = 0.3, 0.5, 0.7, 0.8, 0.9, 0.95. Three batches of particles were synthesized for each x. In those experiments, the synthesis parameters were set to $T_{add} = 40$ °C, $d_{add} = 2 \text{ min}$, $T_{end} = 97$ °C and $d_{react} = 90 \text{ min}$ and the synthesis was carried out under nitrogen atmosphere, to minimize nanoparticle oxidation. These values were chosen considering the results of series 1 of the study, since they enable a high H_C while providing a high $M_{RT,2T}$ at the same time.

After synthesis, particles were washed magnetically with deionized water and stored in deionized water. About 200 mg of each sample were dried under air at room temperature to produce powders for following characterization measurements.

2.2. Characterization of Cobalt Ferrite Nanoparticles

2.2.1. Transmission Electron Microscopy (TEM)

For the evaluation of shape and size of the resulting particles, TEM images were acquired for the samples of series 2. One sample for each cobalt substitution rate x was used as a dry powder and suspended in micropure water at a concentration of 5 mg/mL. The sample was deposited on a carbon support film on a 400-mesh copper grid manufactured by Quantifoil Micro Tools (Großlöbichau, Germany). The films were hydrophilized in an Argon plasma produced by a Diener Electronics (Ebhausen, Germany) plasma oven for 120 s prior to sample deposition. An amount of 10 μ L of the suspension was then placed on the film, the excess blotted off using filter paper, and allowed to air dry. TEM images were acquired with a 200 kV FEI Tecnai G² 20 (Hillboro, OR, USA) using a 1 k × 1 k Olympus MegaView camera (Münster, Gemany) with the acceleration voltage set to 120 kV.

2.2.2. Magnetometry

For a comparative evaluation of the magnetic parameters of the particle batches synthesized in series 1 of the study, VSM measurements at room temperature were conducted using an MSE-EZ9, Microsense, Lowell, MA, USA. A weighed amount of the particle powders, around 3 to 5 mg, was transferred to 3D-printed polymeric measurement vials and hysteresis curves were recorded within a field amplitude of ± 2 T, using a sampling interval of 0.2 T for the saturated branches of the curve and 0.005 T between ± 0.1 T to determine coercivity more precisely. Magnetization at room temperature and a field of 2 T (M_{RT,2T}) as well as coercivity (H_C) were extracted from each measurement and the mean values were calculated for the three particle batches synthesized with the same parameter set.

For a more detailed evaluation of the magnetic behavior of the particles synthesized in series 2 with varied cobalt substitution ratios, magnetometry measurements at 5 and 300 K up to a field of \pm 9 T were carried out using the VSM option of a Quantum Design PPMS DynaCool (San Diego, CA, USA) for one representative sample out of each set of three particle batches with the same x.

2.2.3. Mössbauer Spectroscopy

Mössbauer spectroscopy was utilized for selected samples of series 1 and all samples of series 2, where again one representative sample for each x was characterized. Mössbauer spectra were measured in transmission geometry using a constant acceleration driving unit and ca. 20 mg/cm² of oxide nanoparticle powder. To study sample composition based on magnetic phase transitions, spectra between 4 and 300 K were recorded in a closed-cycle cryostat (SHI-850-5, Lake Shore Cryotronics, Westerville, OH, USA), while in-field spectroscopy was performed using a l-He bath cryostat with a superconducting split-coil magnet allowing maximum magnetic fields of 10 T (Spectromag-4000-10, Oxford Instruments, Abingdon, UK) to analyze magnetic alignment behavior in more detail. Upper estimations of the hydroxide fraction are based on the relative spectral intensity of the corresponding subspectrum observed in spectra at 300 K.

2.2.4. X-Ray Diffraction (XRD)

Crystal structure and size of the particles were evaluated using X-ray diffraction. Diffractograms were measured using a Panalytical X'Pert Pro MPD theta-theta diffractometer (Malvern Panalytical Ltd., Malvern, UK) with parallel beam setup and 1D line detector in scanning mode. For fluorescence reduction, the lower discrimination threshold of the PIXCEL detector was set to 45.5%. The divergence and anti-scatter slits were 0.5° and beam mask was 10 mm. Measurements were taken from 15° to 85° 20 with 2 h measurement time. Phase identification and crystallite sizing was undertaken with Malvern Panalytical

Highscore Plus V. 4.9 [29] and the databases ICDD pdf2 from 2001 and COD 2021. Crystallite sizing was completed with the "size-strain-analysis-R"-method (SSA-R) for the phase peaks, which provides more accurate values compared with the commonly used Scherrer method, which only takes one peak into account. Previously, the instrumental broadening was determined for the X-ray optics used with a 100% anatase powder sample (TiO₂). The powder samples were ground by hand with a mortar prior to measurement. The sample holder was a "zero background holder" (ZBH) with a recess of 15 mm diameter and 0.2 mm depth, into which the powder was filled flush. The ZBH rim was adjusted to the target height of 1 mm (\pm 1 µm) to avoid peak shift due to height error. From the highest peak ((311) for CoFe₂O₄) the interplanar spacing d was used to calculate the lattice parameter a using Equation (1).

$$a = \sqrt{d^2 * (h^2 + k^2 + l^2)} \tag{1}$$

2.2.5. Calorimetric Measurements

The heating power of the particles was determined using a calorimetric hyperthermia measurement setup. As a field generator a SINAC 12 SH, from EFD Induction GmbH, Freiburg im Braisgau, Germany, was used together with a FOTEMP 2 fiber optical temperature probe from Weidmann Technologies Deutschland GmbH, Dresden, Germany. The resolution of the FOTEMP 2 was 0.1 K with a standard resolution of ± 0.3 K. Measurements were made for all synthesized samples and averaged for the three samples with the same x. Specific absorption rate (SAR) as a measure for the heating power of nanoparticles was determined in fluid and agarose samples. Fluid samples consisted of 0.5 mL deionized water with 1 wt% of particles. Agarose samples for immobilized particles contained 1 wt% particles in 0.5 mL of a 1 wt% agarose gel. The external alternating magnetic field was set to a frequency of 290 kHz and amplitudes of 40, 54 and 69 mT. For each particle sample, three fluid and three gel measurement samples were prepared, and the temperature increase was measured at each field amplitude with a sampling interval of 0.5 s. The temperature probe was placed in the center of the sample, and no sedimentation of the particles was observed within the measurement time. Second, SAR values were calculated using Equation (2) for each measurement. Last, the three SAR values for each particle and field amplitude were averaged. This procedure was used because SAR measurements are subject to a rather large statistical error that can be minimized by averaging.

$$SAR = \frac{\Delta T}{\Delta t} * c * \frac{m_S}{m_{MNP}}$$
(2)

The *SAR* is given by Equation (2), where *c* is the heat capacity of water, m_S is the mass of the sample and m_{MNP} is the mass of particles within. The same heat capacity was used for the agarose samples, as the concentration of agarose in water of 1 wt% did not change the heat capacity significantly. Δt was set to 10 s and ΔT as an absolute value over this time span was extracted from the temperature curves (see Figure 1) by utilizing a MATLAB script for automatization. For further details on *SAR* measurements, refer to [30].

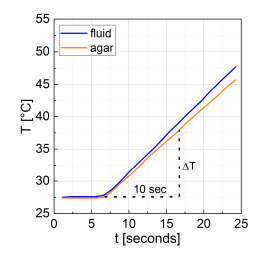


Figure 1. Representative temperature curve of a particle sample with cobalt content x = 0.3 measured as a fluid and in agar gel at 290 kHz and 69 mT. ΔT was extracted as an absolute value after 10 s of heating.

3. Results and Discussion

3.1. Series 1—Variation of Synthesis Parameters

3.1.1. XRD

For each parameter set of series 1, a sample was characterized using XRD. The results can be seen in Table 2, where the crystallite size, the crystallinity and the lattice parameter a are given. Some samples did show additional crystalline phases in XRD and are marked red in Table 2.

Table 2. Size, crystallinity and lattice parameter a for all parameter sets. The varied parameters are written in bold. Red values represent samples with additional crystalline phases, apart from CoFe₂O₄, determined by XRD.

					O ₂ Atmosphere	N ₂ Atmosphere			
d _{add}	T _{add}	T _{end}	d _{react}	Size	Crystallinity	а	Size	Crystallinity	а
min	°C	°C	min	nm	%	Å	nm	%	Å
0				12.9	42.7	8.391	11.4	43.6	8.396
2	10		<i>(</i> 2	12.6	43.6	8.383	15.7	41.4	8.395
4	40	97	60	12.0	35.6	8.364	17.0	37.8	8.392
8				13.6	40.0	8.384	17.4	36.3	8.395
	25			13.3	42.3	8.389	11.0	36.0	8.392
	40	07		12.9	42.7	8.391	11.4	43.6	8.396
0	65	97	60	13.8	26.9	8.415	12.1	44.4	8.398
	85			13.8	38.9	8.381	14.1	38.4	8.395
	25			11.3	33.0	8.375	14.4	38.5	8.401
2	40		60	12.6	43.6	8.383	15.7	41.4	8.395
	65	97		13.0	33.3	8.372	18.0	36.9	8.393
	85			14.2	32.7	8.426	12.5	28.5	8.417
		70		12.8	30.7	8.342	10.3	28.2	8.364
2	40	85	60	11.9	35.4	8.355	15.4	38.4	8.402
		97		12.6	43.6	8.383	15.7	41.4	8.395
	40		30	12.0	34.8	8.376	15.5	33.1	8.400
2		97	60	12.6	43.6	8.383	15.7	41.4	8.395
			90	11.9	37.4	8.361	16.4	34.3	8.405

Most samples of series 1 were single phase $CoFe_2O_4$ and 7 samples were multi-phase $CoFe_2O_4$ plus NaCl and/or FeO(OH) with $CoFe_2O_4$ fractions between 35% and 89%. In one sample ($O_2_85 \ ^\circC_2 \ min_60 \ min_97 \ ^\circC$) $Co(OH)_2$ and magnetite (Fe₃O₄) were indicated together with FeO(OH) and NaCl to form the most likely phase composition. Determining phase composition with XRD for our samples needs to be handled with care, as the diffractograms of magnetite, maghemite and cobalt ferrite in nanoparticulate form are difficult to distinguish. Quantification was carried out using the "direct derivation method" [31]. In Tables 2 and 3, the samples with additional phases are marked red.

Table 3. Mean H_C and $M_{RT,2T}$ with standard deviation for all parameter sets. The varied parameters are written in bold. Red values represent samples with additional crystalline phases, apart from CoFe₂O₄, determined by XRD.

				O ₂ Atm	osphere	N ₂ Atmosphere	
d _{add}	T _{add}	T _{end}	d _{react}	M _{RT,2T}	H _C	M _{RT,2T}	H _C
[min]	[°C]	[°C]	[min]	[Am ² /kg]	[kA/m]	[Am ² /kg]	[kA/m]
0				51.67 ± 1.1	28.30 ± 4.1	59.44 ± 0.8	10.16 ± 0.2
2	40	07	(0	39.40 ± 0.5	38.72 ± 2.2	52.21 ± 0.8	26.87 ± 1.4
4	40	97	60	36.44 ± 1.1	44.91 ± 3.0	49.90 ± 2.4	27.81 ± 3.4
8				36.79 ± 0.5	41.65 ± 2.1	49.54 ± 0.8	24.05 ± 2.0
	25			47.53 ± 0.5	39.84 ± 2.6	60.27 ± 0.3	9.22 ± 0.1
0	40	07	60	51.67 ± 1.1	28.30 ± 4.1	59.44 ± 0.8	10.16 ± 0.2
0	65	97		52.13 ± 0.8	29.41 ± 1.8	61.47 ± 0.5	11.63 ± 0.5
	85			48.51 ± 2.5	12.33 ± 1.5	54.08 ± 8.0	10.86 ± 2.3
	25			34.70 ± 0.8	24.98 ± 2.3	55.03 ± 0.5	26.60 ± 0.9
2	40	07	(0	39.40 ± 0.5	38.72 ± 2.2	52.21 ± 0.8	26.87 ± 1.4
2	65	97	60	28.49 ± 1.6	61.28 ± 11.6	51.74 ± 1.8	30.45 ± 2.8
	85			14.06 ± 4.0	18.11 ± 2.6	7.55 ± 0.4	1.01 ± 0.4
		70		13.27 ± 0.6	108.31 ± 2.3	19.52 ± 2.1	4.65 ± 0.7
2	40	85	60	24.70 ± 1.8	60.14 ± 11.4	51.89 ± 1.9	25.50 ± 1.9
		97		39.40 ± 0.5	38.72 ± 2.2	52.21 ± 0.8	26.87 ± 1.4
			30	29.90 ± 1.0	40.00 ± 5.3	43.68 ± 1.4	23.05 ± 1.4
2	40 97	97	60	39.40 ± 0.5	38.72 ± 2.2	52.21 ± 0.8	26.87 ± 1.4
			90	32.51 ± 2.8	39.19 ± 7.6	55.33 ± 1.5	32.22 ± 2.3

Figure 2 shows typical diffraction patterns of our samples. The peaks of the nanocrystalline powders are rather low because of the large amounts of amorphous components, resulting in crystallinity values from 26.9 to 44.4%. The rise of the background between 15° and 30° 20 is caused by air scattering. For the determination of the crystallinity, the diffraction pattern of a ZBH measurement must be subtracted beforehand, which reflects the influence of the air scattering. The amorphous components and thereby low reflection peaks complicate the phase identification: for reliable detection of a peak, an SNR > 3 is needed, a threshold that is not reached for some of the peaks in our XRD measurements. Phase identification for our samples therefore needs to be considered with care.

The crystallite sizes are between 10.3 and 18.0 nm determined with the "size-strainanalysis-R"-method (Rietveld) for the $CoFe_2O_4$ phase. The microstrain values are near zero ranging from 0% and 0.6% for single phase samples and between 0.6% and 1% for multi-phase samples.

Lattice parameters are derived from the interplanar spacing of the (311) plane, with former ranging between 8.342 Å and 8.426 Å, and most likely depend on the formation of a hydroxide phase. This may result in an iron deficit of the spinel phase, which is described in more detail below.

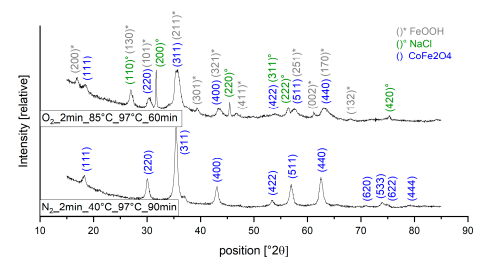


Figure 2. XRD spectra for sample $O_2_2 \min_{85} C_{97} C_{60} \min$ as an example for particles containing multiple phases and sample $N_2_2 \min_{40} C_{97} C_{90} \min$ as an example for pure cobalt ferrite particles.

Only a partial correlation between magnetic properties (described in detail in the next chapter) or varied synthesis parameters and crystallite size can be found. For single varied parameters under N₂ atmosphere, a correlation between synthesis parameter and crystal size can be seen—particles become larger with increasing d_{react} (15.5 to 16.4 nm), as well as with increasing d_{add} (11.4 to 17.4 nm) and increasing T_{add} (11.0 to 14.1 nm). Under O₂ atmosphere, no correlation of size and synthesis parameters can be found. Therefore, other factors beyond the macroscopic particle size need to be considered when discussing the change in magnetic parameters with variations of the synthesis process. Crystallinity, determined from XRD measurements, also seems to correlate with few synthesis parameters. For both O₂ and N₂, increasing the end temperature of the reaction increases crystallinity (for N₂ from 28.2 to 41.4%, for O₂ from 30.7 to 43.6%), indicating a dependency of the formation of a well-structured crystal on the temperature to which the precipitate is heated.

3.1.2. Magnetic Properties

All samples of series 1 were characterized magnetically using VSM measurements at room temperature (RT). $M_{RT,2T}$ and H_C were averaged for the three samples synthesized with the same parameter set. The results are listed in Table 3 and shown graphically in Figures 3 and 4.

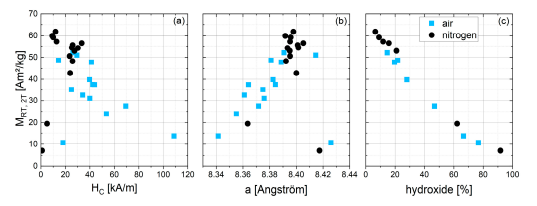


Figure 3. Correlation of $M_{RT,2T}$ of series 1 samples with (**a**) coercivity H_C , (**b**) lattice parameter a, and (**c**) estimated hydroxide content from Mössbauer spectra for selected samples of series 1.

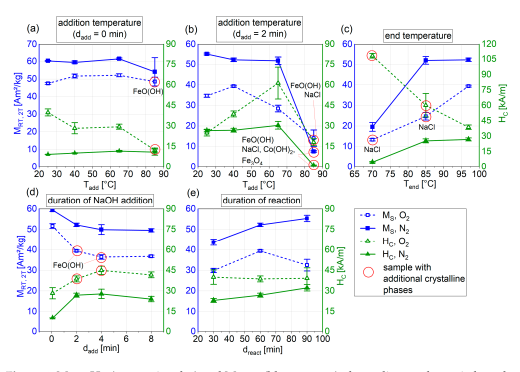


Figure 4. Mean H_C (green triangles) and $M_{RT,2T}$ (blue squares), depending on the varied synthesis parameter (**a**) addition temperature with $d_{add} 0 = \min$ (**b**) addition temperature with $d_{add} = 2 \min$ (**c**) end temperature (**d**) duration of NaOH addition and (**e**) duration of reaction. Full symbols represent samples synthesized under the N_2 atmosphere, while empty symbols represent the O_2 atmosphere. Red circles mark samples with crystalline phases, apart from CoFe₂O₄, determined with XRD. Possible phases are indicated by the circles.

Please note that the magnetization values at 2 T and 9 T for series 2 of this study, described below, were extracted from the measurement as high field magnetization values, while the M(H) curves indicate that the particles were not yet saturated at 9 T. Therefore, the measured values are named $M_{RT,2T}$ and M_{9T} and not saturation magnetization M_S .

All of the varied synthesis parameters show an influence on the resulting magnetic characteristics of the particles. Figure 3 gives an overview of the magnetization of all particles from series 1 at room temperature and a field of 2 T in dependence of H_C , lattice parameter a and hydroxide content.

Concerning the used atmosphere, oxygen or nitrogen, an overall trend can be seen (Figure 3a): with values ranging from 12 to 108 kA/m and an average of 40.7 kA/m, the particles synthesized with O_2 show significantly higher H_C , compared with the reaction under N₂, where H_C ranges from 1 to 32 kA/m with an average of 19.8 kA/m. $M_{RT,2T}$, in contrast, is higher for N₂ samples ranging from 7.6 to 61.5 Am^2/kg with an average of 49.9 Am^2/kg , and 13.3 to 52.1 Am^2/kg with an average of 36.7 Am^2/kg for O₂ samples. Figure 3b shows the lattice parameter a, derived from the position of the (311) peak in the XRD data. A linear correlation can be seen between the lattice parameter and $M_{RT,2T}$, whereby an increase in lattice parameter a correlates with a higher $M_{RT,2T}$. A shift in lattice parameter of a crystal lattice can originate from a rearrangement of the ion distribution on the octahedral and tetrahedral sites [32]. However, since the effect on the magnetization is rather large, ion redistribution as the only reason for the increasing lattice parameter and decreasing magnetization seems unlikely. A more likely explanation is the formation of an additional phase of low magnetization-described in more detail below and as indicated by the Mössbauer spectra of some samples-pointing towards akageneite (FeO(OH)). Depending on pH during synthesis, akageneite can precipitate in an amorphous structure [33], explaining why XRD did find crystalline FeO(OH) only in some of these samples. The estimated amount of hydroxide material in those samples

can be seen in Figure 3c and correlates well with the magnetization. The formation of such an iron-containing hydroxide phase would result in a slight excess of cobalt in the remaining particles, shifting the stoichiometry of CoFe₂O₄ towards more cobalt-rich phases such as FeCo₂O₄. Lattice parameters for CoFe₂O₄ are reported in the range of 8.371 A [34] to 8.40 Å [35], while smaller lattice parameters have been observed for cobalt-rich ferrites. Ateia et al. [36] found the lattice parameter of $Co_{1.5}Fe_{1.5}O_4$ to be 8.366 A, while $CoFe_2O_4$ resulted in a = 8.382 Å in their study. Increasing Co content further to $FeCo_2O_4$, results in a lattice parameter a as small as 8.242 Å [37]. Discussing the formation of cobalt-rich phases, one needs to note that FeCo₂O₄ is stable only between 850 and 950 °C and decomposes into two spinel phases below 850 °C [38]. Therefore, we expect the cobalt-rich phase in our particles to have a stoichiometry close to $Co_xFe_{3-x}O_4$ with $x \ge 1$. Cedeno-Mattei et al. [20] and Whaba et al. [39] synthesized cobalt ferrites with varying x up to values larger than 1 and observed a slight decrease in lattice parameter a for x > 1 as well, supporting our assumption of decreasing lattice parameter with increasing cobalt content. Higher M_{RT.2T} and larger lattice parameters for particles synthesized under N2 indicate the suppression of the hydroxide phase under an inert atmosphere due to reduced oxidization. Looking at Figure 3a, an overall negative correlation between H_C and M_{RT2T} can be seen—particles with a high H_C show low M_{RT.2T} and vice versa. This can be explained by the slightly higher H_{C} values found for x > 1 [40] and the lower net magnetization due to the antiferromagnetic character of the mainly amorphous hydroxide phase.

In the following section, the influence of the varied synthesis parameters will be discussed for each parameter.

Looking at the varied temperatures during synthesis (addition temperature T_{add} and the end temperature T_{end} of the reaction, Figure 4a–c), thresholds seem to exist, which, if exceeded, lead to impurity phases beyond the CoFe₂O₄. Samples, where additional phases were observed via XRD, are marked red in Tables 2 and 3 and Figure 4.

The temperature T_{add} of the metal salt solution when NaOH is added for the precipitation process was evaluated for fast ($d_{add} = 0 \text{ min}$) and slow ($d_{add} = 2 \text{ min}$) addition of the alkaline medium. For a slow addition, particle M_{RT,2T} and H_C decrease drastically for $T_{add} = 85 \,^{\circ}C$ for both the O₂ and N₂ atmospheres (Figure 4b). This drop in the magnetic behavior could be related to the occurrence of crystalline nonmagnetic phases in the particles as detected by XRD measurements. Under N_2 atmosphere, FeO(OH) and NaCl were detected in addition to CoFe₂O₄, while the sample synthesized under O₂ is assumed not to contain any crystalline CoFe₂O₄. Instead, a small amount of Fe₃O₄, together with FeO(OH), NaCl and Co(OH), was indicated by XRD. Akageneite can precipitate in solutions containing Fe^{3+} ions, if pH < 4 or pH > 8 and a high concentration of chloride is present, which is the case for our synthesis method using iron- and cobalt chloride. Without chloride, goethite (α -FeO(OH)) would form under the same conditions [41]. This process starts around 55 °C and is enforced with rising temperature, [42] explaining the formation of akageneite that results from increased addition temperatures in our experiments. The precipitation of akageneite was also observed during the synthesis, as the metal salt solution turned opaque above a temperature of approximately 70 °C.

For the slow addition of the alkaline medium and a T_{add} between 25 and 65 °C only O_2 samples show an influence of T_{add} -H_C increases with increasing T_{add} and $M_{RT,2T}$ decreases for T_{add} = 65 °C. This indicates an influence of O_2 on the hydroxylated ferric ions in the salt solution. Hydroxylation of ferric ions takes place at 1 < pH < 4, as does the rapid condensation of the hydroxylated ion–aquo complexes, to form ferrihydrite [41]. Co^{2+} ions, in contrast, need pH > 8 to hydroxylate and condensate [43]. A temperature increase of the metal salt solution before the alkaline medium is added, and while the metal salt solution is still strongly acidic, therefore most likely has an influence on the condensation of the ferrihydrites, matching the assumed formation of FeO(OH), either crystalline or amorphous.

Compared with the slower addition of the alkaline medium over 2 min discussed above, increasing T_{add} when NaOH is added rapidly to the metal salt solution ($d_{add} = 0$ min,

Figure 4a) results in different trends. For the N₂ atmosphere, only a slight increase in H_C with increasing T_{add} and a slight drop in M_{RT,2T} for T_{add} = 85 °C can be seen. For the O₂ atmosphere, T_{add} = 85 °C again leads to the detection of crystalline akageneite FeO(OH) in XRD measurements; however, most likely only very small amounts of akageneite are formed, as M_{RT,2T} does not decrease significantly. For T_{add} = 25–65 °C, H_C decreases with increasing T_{add}.

Looking at T_{end} (Figure 4c), which represents the maximum temperature that the reactants are heated up to after the precipitation, the O₂ samples show remaining amounts of NaCl for T_{end} = 70 and 85 °C. For T_{end} = 70 °C, a very high H_C (108 kA/m) and a low M_{RT,2T} (20 Am²/kg) can be seen, indicating the incomplete formation of a CoFe₂O₄ crystal lattice. In XRD measurements of this sample, no other crystalline phase, apart from NaCl and CoFe₂O₄, was found; however, Mössbauer spectroscopy, as shown below does display a doublet subspectrum, indicating the presence of an antiferromagnetic phase, most likely an amorphous iron hydroxide, as discussed above. With increasing T_{end}, H_C decreases and M_{RT,2T} increases, leading to particles with a high magnetization of almost 39.4 Am²/kg and a H_C of 38.7 kA/m for T_{end} = 97 °C. For synthesis under N₂ atmosphere, a T_{end} of 70 °C seems also to be insufficient, leading to particles with a low M_{RT,2T} of 19.5 Am²/kg and no significant H_C and indicating that the formation of ferrimagnetic CoFe₂O₄ was hindered. T_{end} > 85 °C enables the formation of ferrimagnetic particles with M_{RT,2T} above 48 Am²/kg and H_C of 25 to 29 kA/m.

Apart from synthesis temperatures, the duration of NaOH addition and of the overall reaction were evaluated. Looking at the duration of NaOH addition (Figure 4d), all samples synthesized under N₂ were pure CoFe₂O₄ derived from XRD and no crystalline impurities were found. For O₂ atmosphere, samples with d_{add} = 2 and 4 min showed some crystalline FeO(OH) in XRD measurements. Interestingly, the sample synthesized with d_{add} = 8 min in O₂ atmosphere did not show a second phase in XRD. Because all of the particles studied are only partially crystalline, and amorphous components contribute significantly, there are only few relatively low and broad reflections, making phase identification difficult. Therefore, it must be considered that the contribution of additional phases may be too small to allow reliable detection. Concerning magnetic properties, increasing d_{add} results in similar trends for O₂ and N₂ samples (Figure 4d). With increasing d_{add}, leading to a slower increase in pH and thereby a slower precipitation process, H_C increases up to d_{add} = 4 min and slightly decreases again for d_{add} = 8 min. M_{RT,2T} simultaneously decreases with increasing d_{add}, reaching a plateau around 50 Am²/kg for N₂ samples and 36 Am²/kg for O₂ samples.

The duration of the reaction (d_{react}), starting after NaOH addition and ending with the removal of the reactants from the water bath and the washing of the particles, has a minor influence on the resulting parameters (Figure 4e). For synthesis under N₂ atmosphere, $M_{RT,2T}$ can be increased by prolonging the reaction time with a maximum value of $M_{RT,2T} = 55.3 \text{ Am}^2/\text{kg}$ for d_{react} = 90 min, while slightly increasing H_C at the same time. For O₂ samples, H_C is not affected by the reaction time while $M_{RT,2T}$ has its maximum at a medium duration of reaction of 60 min.

Summarizing the results of the first series of synthesis with a variation of synthesis parameters, we can conclude that, for high $M_{RT,2T}$, the synthesis under nitrogen atmosphere should be favored, while synthesis under oxygen leads to higher H_C . The increase in temperature of the salt solution when adding NaOH should not exceed 65 °C in any case, for synthesis under oxygen 40 °C may not be exceeded to prevent a decrease in $M_{RT,2T}$. The end temperature of the reactants should be 97 °C, to prevent evaporation of the solvent by boiling while enabling crystallization and high $M_{RT,2T}$. Duration of NaOH addition can be used to tune H_C , but a trade-off between H_C and $M_{RT,2T}$ needs to be made, wherefore 2 min is suggested as d_{add} for further experiments. The overall duration of the reaction should be 90 min for synthesis under N_2 and 60 min for O_2 . The parameters for the synthesis of the samples in series 2 with varied amounts of cobalt were therefore chosen to be $T_{add} = 40$ °C, $d_{add} = 2 \min$, $T_{end} = 97$ °C and $d_{react} = 90 \min$.

3.1.3. Mössbauer Spectroscopy

To better understand the magnetic structure as well as the composition of the particle powders prepared under different conditions, we performed an in-depth Mössbauer spectroscopy analysis. Spectra recorded at 4 K in an external magnetic field of 8 T parallel to the γ -ray propagation direction are shown in Figure 5, illustrating samples of very low (a), moderate (b), and high (c) hydroxide fraction appearing in addition to the expected spinel phase. For synthesis parameters as used in (a), the spectrum consists primarily of well-resolved contributions of Fe³⁺ in tetrahedral (A-site, blue) and octahedral (B-site, green) surroundings, as expected for CoFe₂O₄. Ideal synthesis parameters described above and used for the preparation of particle powders in series 2 with varying stoichiometry, as presented in the following section, resulted in hydroxide fractions far below 10%, wherefore this component will not be considered separately in the Mössbauer data analysis of series 2.

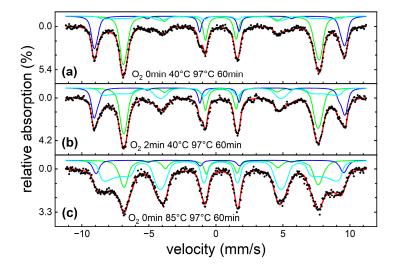


Figure 5. Mössbauer spectra of exemplary samples showing low (**a**), moderate (**b**), and high (**c**) hydroxide fraction recorded at 4 K in an external magnetic field of 8 T along the γ -ray propagation direction. Black dots represent experimental data points, red lines the overall theoretical fit function. Spinel subspectra corresponding to iron on A-sites (blue) and B-sites (green) were reproduced using narrow hyperfine field distributions, while the additional contribution assigned to an akageneite-like Fe³⁺ antiferromagnetic phase (cyan) was represented using a simulation of an ensemble of randomly oriented antiferromagnetic particles in an external magnetic field.

For higher hydroxide fractions, the antiferromagnetic character of this phase became more evident, resulting in the high intensity of lines 2 and 5, representing non-aligned spins [44] and characteristic lineshapes, respectively, as reported before for antiferromagnetic nanoparticles exposed to high magnetic fields [45]. This component was therefore reproduced by simulating an ensemble of randomly oriented antiferromagnetic particles in an external magnetic field, using literature values of exchange and anisotropy fields of bulk akageneite [46], closely matching the shape of the experimental spectra. Independent of XRD analysis, the identification of the byphase as akageneite can also be inferred from the temperature-dependent Mössbauer spectroscopy below. In general, the nanoparticles' degree of spin frustration does not seem to exhibit a strong correlation to the amount of hydroxide, with the spinel phase showing moderate spin canting angles, based on observed subspectral line intensity ratios. As it is reasonable to assume that ferri- to antiferromagnetic interfaces of mixed spinel–hydroxide particles would result in strong interface spin frustration, this would indicate that spinel and hydroxide species grow as individual particles.

All nanoparticle powders were analyzed via Mössbauer spectroscopy between 4 and 300 K, to identify their composition and to examine them with regards to superparamagnetic relaxation effects. Figure 6a shows a sample composed almost entirely of spinel

material. At elevated temperatures, a minor asymmetry of the absorption lines is visible, originating from beginning Néel-type superspin relaxation. A small doublet is also visible at this temperature, containing ca. 14% of spectral area. This could represent a fraction of the smallest spinel particles, already reaching fast superparamagnetic relaxation, but likely also contains small fractions of paramagnetic byphase, as becomes evident based on Figure 6b. At low temperatures, we see that the hydroxide subspectrum partially overlaps with spinel components due to similar hyperfine parameters: an isomer shift of ca. 0.48 mm/s (relative to α -Fe at RT), which is characteristic for Fe³⁺; a hyperfine magnetic field B_{hf} of ca. 48.2 T; and a nuclear quadrupole level shift 2 ϵ of ca. -0.1 to -0.15 mm/s. These parameters match those reported previously for akageneite [47,48], in agreement with results from XRD analysis. This identification is further supported by studying the temperature-dependent decrease in B_{hf}, yielding a Néel temperature of ca. 290 K, again in agreement with literature values of akageneite, before reaching the paramagnetic doublet state (orange) at room temperature.

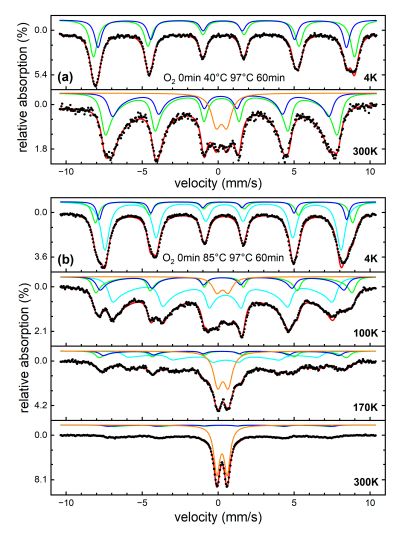


Figure 6. Mössbauer spectra recorded between 4 and 300 K for exemplary samples showing low (**a**) and high hydroxide contents (**b**), respectively. Black dots represent experimental data points, red lines the overall theoretical fit function. Asymmetric deformation was reproduced using a multilevel relaxation model within the slow relaxation regime [49]; spectra consist of A- and B-site spinel subspectra (blue and green), as well as akageneite in the antiferromagnetic low-temperature state (cyan). The high temperature doublet state (orange) is assigned to a superposition of superparamagnetic spinel particles and paramagnetic akageneite.

3.2. Series 2—Variation of Cobalt Fraction x

In a second series of experiments, $Co_x Fe_{3-x}O_4$ samples with varying cobalt fraction x were synthesized with the same set of synthesis parameters under a nitrogen atmosphere.

Table 4 gives an overview of the characterization of series 2 particles, including magnetometry and XRD-derived results. They will be described and discussed in detail in the following sections.

Table 4. Results of magnetometry and XRD for samples of series 2 with varied cobalt fraction x. H_C and M_{9T} were measured at 5 and 300 K with a maximum field of 9 T.

x	Co ²⁺ / Fe ²⁺ +Fe ³⁺	H _C @ 5 K [kA/m]	H _C @ 300 K [kA/m]	M _{9T} @ 5 K [Am ² /kg]	M _{9T} @ 300 K [Am ² /kg]	a [Å]	Size [nm]	Crystallinity [%]
0	0.00	18.31	0.80	81.57	70.77	8.380	12.4	50.6
0.3	0.11	464.07	11.54	91.62	76.52	8.391	14.4	50.1
0.5	0.20	431.43	20.70	93.65	77.43	8.395	16.5	45.8
0.7	0.30	548.44	32.64	90.01	72.48	8.407	16.6	45.0
0.8	0.36	612.92	37.41	90.97	72.81	8.397	15.2	40.4
0.9	0.43	757.79	37.41	92.08	72.38	8.406	14.5	40.5
0.95	0.46	717.99	34.23	88.41	68.57	8.410	14.5	41.3
1	0.50	751.42	29.45	81.98	61.92	8.405	16.4	34.3

3.2.1. TEM Analysis

TEM micrographs (Figure 7) show spherical to slightly irregular shaped particles that confirm the size range determined from XRD measurements between 12.4 and 16.6 nm. An influence of cobalt fraction x on the particle shape and size can be assumed: with increasing x, size distributions seem to become broader, with an increasing amount of very small particles, resulting in a rather inhomogeneous distribution for x = 1. Mean particle size determined by XRD did not show a clear trend with increasing x. One has to keep in mind that XRD only takes into account crystalline components, while the investigated particles most likely contain amorphous phases to some extent, leading to uncertainties regarding the size determination.

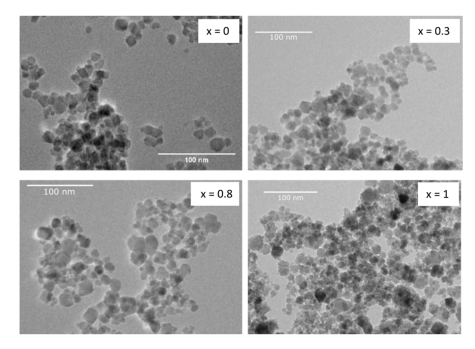


Figure 7. TEM micrographs of samples x = 0, 0.3, 0.8, 1—deposited from solutions of 5 mg/mL concentration.

3.2.2. XRD

Diffractograms of all series 2 samples can be seen in Figure 8. All samples were single-phase, identified with either a Fe₃O₄ (magnetite) reference card for x = 0 or CoFe₂O₄ reference card for x > 0. Co doping from magnetite to CoFe₂O₄ has an effect on the lattice spacing, but not on the crystal lattice structure, which remains cubic. The lattice spacing d and thus the lattice parameter a become about 0.4% larger, which is shown in Figure 9. The crystallite size for the series 2 samples varies from 12.4 nm (x = 0) up to 16.6 nm (x = 0.7), the micro strain from 0% (x = 0) up to 0.13% (x = 0.7), determined with the "size-strain-analysis-R"-method (Rietveld) for the CoFe₂O₄ phase.

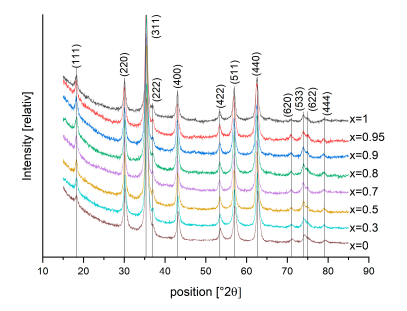


Figure 8. XRD diffractograms of all samples with varying Co fraction x. For better visibility, the curves are shifted on the *y* axis.

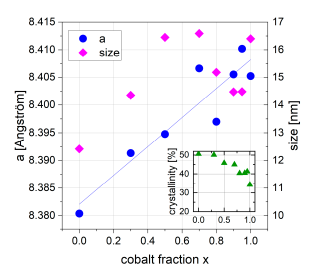


Figure 9. Lattice parameter a as blue dots (**left**) and crystallite size as pink rhombuses (**right**) plotted against cobalt fraction x. Inset shows crystallinity (green triangles) depending on cobalt fraction x.

The Co doping of magnetite up to cobalt ferrite increases the calculated lattice parameter from 8.38 Å to 8.41 Å (Figure 9). Thus, our $CoFe_2O_4$ sample has a slightly larger lattice parameter compared with reference card values from the ICDD, ICSD or COD database, where the lattice parameter for $CoFe_2O_4$ varies from 8.350 Å up to 8.400 Å. An increase in lattice parameter a with increasing cobalt fraction can be found in the literature; Kumar

et al. 2021 [23] found a to increase from 8.351 to 8.395 Å with x increasing from 0 to 0.8. In contrast, the opposite trend is also described, with a decreasing lattice parameter with increasing x [27]. The mechanisms behind these contrasting phenomena seem unclear and to be influenced by several factors. Oxidation of magnetite to maghemite for particles with low cobalt content could be one factor, leading to smaller lattice parameters for lower x. The crystallinity varies from 40.4% up to 50.6%, with a decreasing trend for increasing x (see Figure 9, inset), indicating higher amounts of amorphous components for higher cobalt doping. Mössbauer spectroscopy did not find significant amounts of additional iron containing amorphous phases in series 2 samples, which indicates the presence of some cobalt ferrite in an amorphous structure.

3.2.3. Magnetic Properties

To evaluate the magnetic parameters strongly varying upon the change in Co-content x, M(H) hysteresis loops were recorded at 4.3 K up to 9 T as displayed in Figure 10, as well as at 300 K. All samples, except for the x = 0.0 sample, show open hysteresis loops with relatively high coercive fields in the 1 T (ca. 800 kA/m) range, typical for cobalt ferrite nanoparticles, and high-field magnetization values close to 90 Am²/kg. A clearly visible decrease in magnetization when going through zero magnetic field indicates the presence of a soft magnetic fraction, showing a relevant contribution for low Co-content and representing a small fraction of particles of lower magnetic anisotropy. Studying the cobalt fraction x-dependent trend in the 9 T magnetization M_{9T} at 4.3 K and 300 K (see Figure 11), as extracted from the data in Figure 10, we observe a maximum at ca. x = 0.5. Lower values of M_{9T} for higher x could be explained by slightly higher degrees of spin frustration due to higher magnetic anisotropy in Co-rich spinel material, as discussed in more detail in the Mössbauer spectroscopy section. At the same time, the decrease in magnetization when going towards x = 0.0 can be assigned to oxidation, reducing the Fe²⁺ fraction in the spinel particles and thereby their saturation magnetization when vacancies replace B-site Fe²⁺, reducing the B-site sublattice magnetization. One has to keep in mind that a magnetite/maghemite transformation does not necessarily require aerial oxidation and can also happen under inert conditions by desorption of Fe²⁺ ions and hydroxylation in the solvent from the surface of particles, leaving behind a cationic vacancy and only trivalent Fe ions. As a result, oxide layers of ca. 2–3 nm may form within days to weeks, composed chiefly of maghemite, which explains the decrease of the overall magnetization of these oxidized samples [41,50]. At the same time, the difference between the M_{9T} values at 300 K and 4.3 K becomes smaller when considering lower Co-fractions, which could be attributed to the higher Curie temperature of magnetite and maghemite, as compared to cobalt ferrite with $T_C \approx 790$ K [51].

A monotonous increase in coercivity H_C can be observed at 4.3 K when moving from low to high Co-fractions, with a maximum at ca. 750 kA/m for x = 0.9–1.0, while a maximum of ca. 35–40 kA/m is visible at x = 0.8–0.9 at 300 K, followed by a slight decrease when reaching even higher cobalt fractions. A maximum for H_C with medium cobalt ratios and a decrease of H_C when cobalt ratio x approaches 1 can be found in the literature: Yu et al. [21] found a maximum of H_C for x = 0.6 and stated that the co-existence of Fe²⁺ and Co²⁺ ions seems to enhance anisotropy. Kumar et al. [23] found H_C to be highest for x = 0.4–0.6 and suggested that with higher x the particle structure changed from single to multidomain, indicated by the increasing particle size with increasing x. A slightly increasing trend for particle size with increasing cobalt content can also be assumed for our particles (see Figure 9).

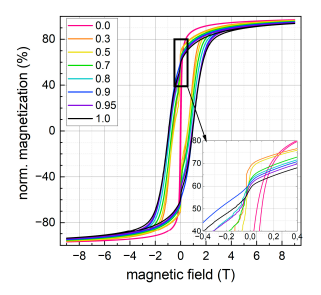


Figure 10. M(H) loops recorded at 4.3 K up to maximum fields of 9 T for x = 0.0–1.0. M(H) data are normalized to the saturation magnetization Ms, estimated by extrapolation of the high-field region.

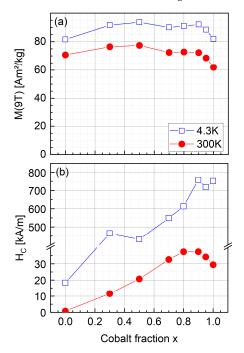


Figure 11. High-field magnetization M_{9T} (**a**) and coercivity H_C (**b**) observed at 4.3 K (open symbols) and 300 K (full symbols), respectively.

3.2.4. Mössbauer Spectroscopy

Mössbauer spectra of spinel particles with x = 0.0 to 1.0 recorded at 4 K in a magnetic field of 8 T (Figure 12) do not exhibit the characteristic features assigned to hydroxides, as discussed above for series 1. As a result, this fraction was disregarded in further analysis, showing subspectra of octahedral (green) and tetrahedral (blue) Fe sites. However, a slight inner shoulder is visible for the B-site sextet subspectrum for x = 0.0 and x = 0.3, which could originate from a minor Fe²⁺ fraction. It seems reasonable to assume that for x = 0.0 the particles consist mainly of maghemite with a small magnetite core, limited to less than 20% of the particle volume based on the fine structure of the B-site sextet. Comparing relative spectral areas of A- and B-site subspectra, we find a ratio R_{AB} close to 0.6, as expected for maghemite, while pure magnetite would translate to R_{AB} = 0.5. Upon rising cobalt content, R_{AB} decreases monotonously until reaching a value of ca. 0.52 for x = 1.0, corresponding to

an inversion parameter of ca. 0.68, which represents a mainly random ion placement with only a very minor B-site preference of Co^{2+} ions.

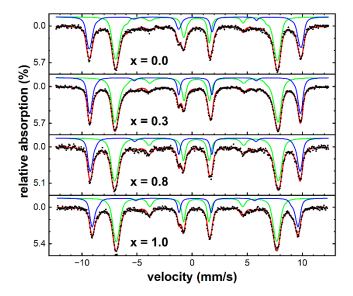


Figure 12. Mössbauer spectra of exemplary samples with cobalt content x = 0.0, 0.3, 0.8 and 1.0 recorded at 4 K in an external magnetic field of 8 T along the γ -ray propagation direction. Black dots represent experimental data points, red lines the overall theoretical fit function. Spinel subspectra corresponding to iron on A-sites (blue) and B-sites (green) were reproduced using narrow hyperfine field distributions.

Although the degree of spin frustration is moderate in all samples, as indicated by the low intensity of lines 2 and 5, average spin canting angles show a slight increase from ca. 19° (x = 0.0) up to ca. 24° (x = 1.0). This could be explained by the higher magnetocrystalline anisotropy of cobalt ferrite as compared with iron oxides, which could add to the slight decrease in M_{9T} from x = 0.5 to x = 1.0 observed in magnetometry experiments. Considering the average spin misalignment angle $\theta_{(8T)}$ of ca. 20° for x = 0.5, determined in an external field of 8 T at 4 K, a saturation magnetization M_S of ca. 99.3 Am²/kg can be estimated via $M_{(8T)} \approx M_S \cdot \cos(\theta_{(8T)})$ from the corresponding magnetization $M_{(8T)} \approx 93.3$ Am²/kg. This value is far higher than expected when assuming an inverse spinel structure, as often discussed for bulk cobalt ferrite; however, it can be explained in terms of the mainly random placement of Co²⁺ ions [52], further supporting findings on inversion parameters discussed above.

3.2.5. SAR

SAR was determined using a calorimetric measurement setup for particles in fluid and immobilized in agar gel at three different field amplitudes. The results can be seen in Figure 13. In general, *SAR* values measured in fluids are slightly higher than those measured in agar gel for the same sample. This indicates that the mechanism of heat generation can be attributed partially to a physical rotation of the particle that is prohibited in agar gel. A physical rotation of the particle can be expected for our particles, since their anisotropy constant is rather high, as can be seen from the high coercivity values. High magnetic anisotropy leads to a blocking of the intrinsic rotation of the magnetic moment within a particle and thus a physical rotation of the particle to follow the external magnetic field. However, the differences in *SAR* for fluid and agar samples are small, indicating that the particles may not have been completely immobilized in the agar gel. The influence of the used excitation field amplitude is as expected: with higher external field amplitudes, *SAR* values increase, leading to highest *SAR* values for the highest field of 69 mT. *SAR* also depends on the cobalt fraction x, with a maximum of 480 W/g (fluid sample, 69 mT) for x = 0.3 and a decrease in *SAR* for higher cobalt fractions. For x > 0.8, a slight increase in *SAR* can be seen, which might be related to the decreasing H_C for those samples (see Figure 11). Looking at Figure 13b, one can see that *SAR* monotonously decreases with increasing H_C, after a maximum for H_C = 11.54 kA/m. For higher H_C, the particles seem to be too magnetically hard to be heated efficiently by the used external field. The external field amplitude needs to be two or three times the coercivity as a rough guideline; however, depending on the particle size distribution and the anisotropy field distribution, even larger field amplitudes may be required to switch the particles' magnetization [53]. In our case, the magnetization of particles with H_C higher than 12 kA/m, about a fourth of the external field amplitude, no longer switch efficiently, leading to the conclusion that the limiting factor for the heating power of the particles is the available external field amplitude. With the development of field generators producing higher field amplitudes for extracorporeal magnetic heating applications, this limitation can be overcome and the full potential of hard magnetic particles can be exploited.

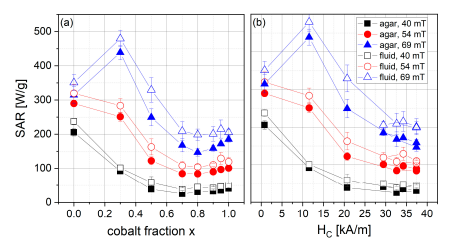


Figure 13. Mean *SAR* values and standard deviation for samples of series 2 in agar gel or fluid samples at varying field amplitude and a frequency of 290 kHz depending on (**a**) cobalt fraction x or (**b**) coercivity.

4. Conclusions

We synthesized cobalt ferrite nanoparticles in the size range of 10 to 18 nm using a wet co-precipitation method. By varying the synthesis parameters in the first series of experiments over a broad range, we were able to identify thresholds that need to be met for the synthesis of crystalline and of pure cobalt ferrite nanoparticles and to prevent the formation of additional phases. Akageneite was found in some samples, indicated by XRD and Mössbauer spectroscopy, leading to a decrease in magnetization. Depending on the synthesis parameters, coercivities of up to 60 kA/m at room temperature and magnetizations (room temperature, 2 T) of 60 Am^2/kg can be achieved. Second, the substitution of Fe²⁺ with Co²⁺ in Co_xFe_{3-x}O₄ with 0 < x < 1 leads to a linear increase in H_C up to x = 0.8, enabling the application-specific tuning of the particles' magnetic behavior from a soft to a hard magnetism. For x > 0.8, H_C slightly decreases. High magnetization values for all $Co_x Fe_{3-x}O_4$ particles, regardless of their cobalt fraction x, between 62 to 77 Am^2/kg (room temperature, 9 T), were measured as a result of the optimized synthesis parameters found in the first part of this study. In-field Mössbauer spectroscopy experiments revealed moderate spin canting angles and a mainly random ion placement, both attributing to the observed high magnetization values. SAR values of up to 480 W/g (290 kHz, 69 mT) were measured for particles with x = 0.3, leading to the conclusion that the available external magnetic field amplitude is the limiting factor for the efficient heating of the hard magnetic high coercivity samples with higher x. The

herein presented particles are promising candidates for extracorporeal magnetic heating applications with high magnetic field amplitudes.

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