Shaping Magnetite with Poly-L-arginine and pH: From Small Single Crystals to Large Mesocrystals

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ABSTRACT: Control over particle size, size distribution, and colloidal stability are central aims in producing functional nanomaterials. Recently, biomimetic approaches have been successfully used to enhance control over properties in the synthesis of those materials. Magnetotactic bacteria produce protein-stabilized magnetite away from its thermodynamic equilibrium structure. Mimicking the bacteria’s proteins using poly-L-arginine we show that by simply increasing the pH, the dimensions of magnetite increase and a single- to mesocrystal transformation is induced. Using synchrotron X-ray diffraction and transmission electron microscopy, we show that magnetite nanoparticles with narrow size distributions and average diameters of 10 ± 2 nm for pH 9, 20 ± 2 nm for pH 10, and up to 40 ± 4 nm for pH 11 can be synthesized. We thus selectively produce superparamagnetic and stable single-domain particles merely by controlling the pH. Remarkably, while an increase in pH brings about a thermodynamically driven decrease in size for magnetite without additives, this dependency on pH is inverted when poly-L-arginine is present.

Supporting Information

Properties of materials typically change when confined to nanoscopic dimensions, opening the door to tailoring nanomaterials toward desired applications. Nanoparticles made from ferrimagnetic magnetite (Fe₃O₄) exhibit superparamagnetic (SP), stable single-domain (SSD), or multidomain behavior depending on their size.¹,² SP nanoparticles are found below a size threshold of 20–30 nm and are widely employed in applications ranging from magnetic ink, data recording, drug delivery, magnetic resonance therapy (MRI), to hyperthermal cancer treatment.³,⁴ Above the SP size threshold and below 80–100 nm, magnetite nanoparticles show SSD character with an intrinsic remanent magnetization that is advantageous for certain biomedical and bioengineering applications.⁵ Controlling the size, and especially size distribution, of magnetite nanoparticles is thus essential to fine-tune their properties toward the desired application.⁶–¹⁰ For most applications, it is not only size that matters, but also colloidal stability, which prevents aggregation, and sedimentation is of equal importance.¹¹,¹² Colloidal suspension stability results when attractive and repulsive forces are in equilibrium. Attractive dipolar forces, which cause aggregation, especially in SSD magnetic nanoparticles, seriously compromise the usefulness of these nanoparticles in applications, such as medical ones where aggregation and sedimentation could clog blood vessels. Thus, to make these particles more operational, a continuous steric or electrostatic stabilization during the growth of these nanoparticles is necessary to counteract attractive magnetic forces, preventing aggregation.

Synthesizing SSD magnetite nanoparticle under mild conditions by co-precipitation of ferrous and ferric iron has so far been a challenge as particle sizes typically range between 2 and 12 nm with a rather broad size distribution when prepared at different pHs and ionic strengths, failing in addition to being colloidal stable.¹³,¹⁴ We recently showed that magnetite particles can be grown beyond the SSD sizes at mild temperatures¹⁵ and/or by increasing the reaction times,¹⁶ but they still form colloidal unstable precipitates.

One encouraging way to overcome these challenges has been adopting biomimetic approaches.¹⁷ Baumgartner et al. emulated the proteome architecture of magnetotactic bacteria and used the polyamino acid poly-L-arginine (polyR) in the in vitro precipitation of magnetite, which resulted in colloidal stable, mesocrystalline nanoparticles of ~40 nm diameter. The magnetic behavior of the mesocrystals surprisingly reflected that of the superstructure¹⁸ rather than that of the building blocks, as reported earlier.¹⁹ Encouraged by these results, we engaged in further developments of these biomimetic functional materials controlling the size and the inherent magnetic properties while exploring polyR properties to alter magnetite’s nucleation and growth characteristics. Here we report the co-
precipitation of ferrous and ferric iron in the presence of polyR at pH values between 9 and 11, resulting in colloidal stable magnetite nanoparticles with a tunable diameter between 10 and 40 nm, showing a morphological transition from spherical single crystals to substructured mesocrystals for high pH. This approach enabled us to selectively produce SP as well as SSD particles with an exceptionally narrow size distribution, as evidenced by (high-resolution) transmission electron microscopy (HR)TEM, synchrotron-based X-ray powder diffraction (XRPD), and vibrating sample magnetometry (VSM).

The polyR/magnetite nanoparticles were synthesized as in Baumgartner et al.20 Briefly, a computer-controlled titration device was connected to a pH electrode, and a dosing device added the iron solution into an alkaline solution of polyR. Particles were grown at pH values of 9, 10, and 11 for 2 h and analyzed using synchrotron X-ray diffraction. The analysis showed an increase in particle size as the pH increased. The azimuthally integrated full XRPD diffraction patterns are shown in Figure 1, with the inset showing a magnified view of the (311) reflection. A broadening of this, and any other reflection, can be directly attributed to a decrease in grain size. This decrease is visible at lower-precipitation pH. Using the Scherrer equation,21 we calculated nanoparticle sizes from the full width at half-maximum of the reflection, which shows particle diameters ranging from 10 ± 0.5 nm for pH 9 to 14 ± 1 nm for pH 10 and 24 ± 2 nm for pH 11. Standard deviations were calculated from the mean particle size of triplicates, showing a high reproducibility of the synthesis. Lattice constants for the cubic system (a=b=c) were obtained from the (311) peak position and calculated to be 8.3734, 8.3741, and 8.3768 Å for pH 9, 10, and 11, respectively, indicating that the obtained particles are indeed magnetite that has undergone slight surface oxidation,22 resulting in a decrease of the lattice constant compared to compositionally pure magnetite (8.3965 Å). The effect of surface oxidation decreases with increasing particle size, which is also observed for magnetite prepared without any additives.16

Complementary TEM measurements, seen in Figure 2, confirm the increase in particle size as the pH is increased. In determining individual sizes for at least 200 particles, we determined intrinsic particle size distributions of 10 ± 2 nm for pH 9, 20 ± 1.8 nm for pH 10, and 38 ± 4.8 nm for pH 11. For comparison, TEM micrographs of corresponding particles prepared in the absence of polyR are shown in the SI. Fast Fourier transformations (FFTs) of HRTEM images of single particles prepared in the presence of polyR were indexed according to an inverse spinel structure with crystal lattice spacing of magnetite. Note the discrepancies in particle size found in Table 1, which are determined by TEM or XRPD. As XRPD is only sensitive to an effective volume measured in the direction of the [311] vector along which the diffraction is coherent, results may differ from TEM, where the radius is calculated from a spatially averaged area projected from a three-dimensional structure. Further, thresholding for size determination in TEM may exclude small particles, which in XRPD contribute to a significant peak broadening, lowering calculated mean sizes. Both methods, however, demonstrate an increase in particles size as the pH increases.

In addition to the observed change in particle size, a clear pH-dependent change in morphology is visually evident: while
maximum magnetization could not be normalized to the mass of the sample, we focus only on the coercivity \((H_C)\) to discriminate between SP and SSD particles. While SP particles show no hysteresis \((H_C = 0)\), SSD particles are characterized by an open hysteresis loop \((H_C > 0)\) when a reversed magnetic field is applied. A change from SP to SSD behavior can be seen between pH 9 and 10, illustrated in the inset of Figure 3, in the opening of the hysteresis loop. Moreover, the increase in coercivity from 6 to 12 mT for particles prepared at pH 10 and 11 were determined from the field strength to completely demagnetize the particles after full magnetization.

To determine the magnetic properties of the nanoparticles, we performed magnetic hysteresis measurements on a VSM device on bulk samples at room temperature (Figure 3). As the maximum magnetization could not be normalized to the mass of the sample, we focus only on the coercivity \((H_C)\) to discriminate between SP and SSD particles. While SP particles show no hysteresis \((H_C = 0)\), SSD particles are characterized by an open hysteresis loop \((H_C > 0)\) when a reversed magnetic field is applied. A change from SP to SSD behavior can be seen between pH 9 and 10, illustrated in the inset of Figure 3, in the opening of the hysteresis loop. Moreover, the increase in coercivity from 6 to 12 mT for particles prepared at pH 10 and 11 were determined from the field strength to completely demagnetize the particles after full magnetization.

"Particle size distributions from TEM were calculated from single particle size in an ensemble of at least 200 particles, while for XRPD mean particle sizes were calculated from mean sizes of triplicates. VSM shows a change from SP to SSD behavior with the appearance of a coercive force \((H_C)\) for particles prepared at pH 10 and 11. The lattice parameter \((a)\) increases with increasing pH.

Table 1. Mean Particle Sizes at Different pHs Analyzed by TEM and XRPD

| pH | \(D_{XRPD}\) [nm] | \(D_{TEM}\) [nm] | \(H_C\) [mT] | \(a\) [Å] |
|----|-----------------|-----------------|-------------|---------|
| 9  | 10 ± 0.5        | 10 ± 2          | 0           | 8.3734(1) |
| 10 | 14 ± 1.7        | 20 ± 2          | 6           | 8.3740(6) |
| 11 | 24 ± 2.0        | 38 ± 5          | 12          | 8.3767(3) |

"Particulate morphology is even more visible in the HAADF STEM image shown in the middle inset of Figure 2c. The morphology, size, and size distribution of the particles, as seen in the TEM images, are found to remain unaltered in storage up to 7 days, when kept in solution under anaerobic conditions (see Figure 2 in the SI). Colloidal stability could also be visually confirmed for more than 3 months (see the SI). To determine the magnetic properties of the nanoparticles, we performed magnetic hysteresis measurements on a VSM device on bulk samples at room temperature (Figure 3). As the maximum magnetization could not be normalized to the mass of the sample, we focus only on the coercivity \((H_C)\) to discriminate between SP and SSD particles. While SP particles show no hysteresis \((H_C = 0)\), SSD particles are characterized by an open hysteresis loop \((H_C > 0)\) when a reversed magnetic field is applied. A change from SP to SSD behavior can be seen between pH 9 and 10, illustrated in the inset of Figure 3, in the opening of the hysteresis loop. Moreover, the increase in coercivity from 6 to 12 mT for particles prepared at pH 10 and 11 were determined from the field strength to completely demagnetize the particles after full magnetization.

Figure 3. Magnetic hysteresis loops of particles prepared at different pHs. The gray region is shown in higher magnification in the inset to show the opening of the hysteresis loop for particles prepared at pH 10 and 11. Particles prepared at pH 9 show no opening of the loop and thus have no coercivity at the given temperature typical for SP particles. Coercivities of 6 and 12 mT for particles prepared at pH 10 and 11 were determined from the field strength to completely demagnetize the particles after full magnetization.

Figure 4. Particle sizes calculated using the Scherrer equation from the (311) magnetite reflection at \(\sim 25\) nm\(^{-1}\) of particles prepared in the presence of polyR compared to literature values \(*\) of magnetite prepared with the same protocol without additives. Error bars were calculated from the mean size of triplicates.

The arrest of particle sizes and morphologies, as obtained from TEM measurements of freshly synthesized and 7 days aged samples (see Figure 1 in the SI), may suggest that the system is at thermodynamic equilibrium. However, polymer induced electrostatic and steric repulsion, which inhibits coagulation of nanoparticles, as well as the ultralow solubility of iron at strongly alkaline conditions, preventing Ostwald ripening, are considerable energetic barriers that may kinetically trap the nanoparticles in a thermodynamically unfavorable state. Still, this stabilization—preventing coagulation and molecular ripening processes—are most favorable for prospective applications, especially for SSD particles, which normally aggregate and precipitate due to magnetic interaction.

While we find an increase in particle size as the pH increases, magnetite nanoparticles in the absence of additives formed by co-precipitation in alkaline solution decrease in size with increasing alkalinity. We use particle sizes of magnetite prepared under the same conditions but without polyR, as described in the literature, and compared them to pH-dependent particle sizes obtained in the presence of polyR (Figure 4). An inversion of the pH dependence of the nanoparticle size is evident. The decrease in particle size with increasing pH for bare particles was modeled on the basis of classical nucleation theory, following a semiquantitative approach by Jolivet et al. They proposed that an increase in surface charge density caused by a stronger deprotonation at high pH is directly related to a decrease in oxide/water surface tension, thereby decreasing particle size to minimize the system’s free energy. Using this model and potentiometric determination of the surface charge density of freshly prepared magnetite nanoparticles makes it possible to predict magnetite particle sizes as a function of pH. Further, the degree of condensation of counternions at the charged magnetite/water
interface has a significant influence on the sizes of magnetic nanoparticles in solution. An increasing counterion concentration will decrease the interfacial tension because of stronger adsorption, as predicted by the Gibb's adsorption isotherm, and will thus lead to a decrease in particle size. Vayssieres et al.,

**demonstrated that the diameter of magnetite nanoparticles varies, from between 4 nm at low NaN₃ concentrations at a given pH to 0.8 nm for high concentrations. As for ions, we may infer that adsorption of polyR at the magnetite/water interface induces a lower surface tension that would lead to a decrease in particle size. We expect this effect to be even more pronounced in our case as the adsorption of a polyelectrolyte with n charges will come along with a smaller entropic penalty compared to the adsorption of n monovalent counterions, as predicted purely based on statistical placement of the n charges. Unexpectedly, for the polyR magnetite hybrids, we observed the opposite effect as they increase in size with increasing pH. This unusual pH effect rules out that the formation of magnetite in the presence of polyR can be easily explained by notions of classical nucleation based on thermodynamic energy minimization, where particles are thought to grow from stable nuclei by the accretion of ions, resulting in dense, homogeneous particles.

There are many examples of biological and synthetic systems following nonclassical nucleation pathways. These systems often exhibit an excess surface charge, where oppositely charged additives are thought to kinetically trap transient precursor phases, altering the crystallization pathway and resulting in a wealth of sophisticated structures and morphologies of nanoparticles. For magnetotactic bacteria, it was possible to identify transient ferric hydroxide phases stabilized by charged proteins that crystallize to form magnetite via nanometric (oxyhydro)oxide intermediates. Polymorph selection in precipitation of CaCO₃ forming either vaterite or amorphous calcium carbonate in the presence of polyaspartic acid has been demonstrated in in vitro experiments. Moreover, prior research has already demonstrated that adding negatively charged polysaccharides inhibits magnetite nucleation at pH values up to 13.

The unifying concept of the given examples, of both natural and synthetic origin, is the existence of thermodynamically less stable polymorphs such as iron oxyhydroxide or amorphous calcium carbonate from which the thermodynamically more stable phases magnetite and calcite are formed. Past research has shown that interaction with charged macromolecules results in kinetic stabilization, preventing crystallization into the thermodynamically more stable (and also more dense) phase. Proving (or disproving) the existence of these structures will thus be the next logical step toward understanding the influence of polyR on the stabilization of magnetite.

In conclusion, we show that SP as well as SSD magnetite nanoparticles can be selectively grown in the presence of polyR simply by changing the pH of the precipitation solution. Even though the effect of pH, and of biological additives on the precipitation of magnetite has been extensively studied within recent years, we believe that the presented control over size and size distribution of the nanoparticles is exceptional. We thus combine many of the previously reported advantages in one system. These properties in combination with their long-term colloidal stability render these biomimetic functional materials ideal for applications. Unexpectedly, the size of the nanoparticles increases with increasing pH, in contrast to naked magnetite particles produced under the same conditions, which decrease in particle size when the precipitation pH is increased.

Classical notions of nucleation fail to explain this unexpected pH dependence as well as the complex, substructured morphology of the mesocrystals formed at pH 11. Thus, reference to nonclassical nucleation theories suggests that the magnetite/polyR nanoparticles may crystallize from polymer-stabilized transient, low-density iron phases. These noncrystalline iron phases then crystallize only when they attach to the growing nanoparticle's surface, adopting the nanoparticle’s crystallography. This mechanism may explain the single-crystalline behavior of our substructured particles. Magnetite formation via these transient iron phases was already considered by others, a polymer induced stabilization of them, however, has so far not been reported. If this mechanism will be proved, we may have found a way to alter the nucleation pathway of magnetite that, in the presence of poly-arginine, would not follow a pathway of thermodynamic energy minimization, consistent with magnetite biomineralization in the presence of charged proteins. Further measurements will provide necessary data for an extensive understanding of the formation pathway of the polyR/magnetite particles, an understanding that promises to deepen our knowledge of crystal morphology control by charged macromolecules in natural as well as synthetic systems.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcl.9b01771.

Full description of the materials and methods, histograms of particle size distributions of fresh and aged nanoparticles, and photographs of solutions stored for 1 week and 3 months (PDF)

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Notes

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