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Characterization of γ - and α -Fe₂O₃ nano powders synthesized by emulsion precipitation-calcination route and rheological behaviour of α -Fe₂O₃

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Abstract

Nano crystals of γ -Fe₂O₃ (maghemite) were synthesized by emulsion precipitation method using kerosene as oil phase, SPAN-80 (sorbitane monooleate) as the surfactant and sodium hydroxide as the precipitating agent. The characterization of the samples by FTIR (Fourier transform infra-red) and XRD (X-ray diffraction) techniques confirmed the formation of γ -Fe₂O₃ (maghemite). Analysis by SEM (scanning electron microscope) and TEM (transmission electron microscope) was carried out to study the morphology and particle size. The as prepared samples contained inverse spinel cubic phase maghemite. Effect of initial iron concentration on crystallite size of maghemite showed that it decreased with the decrease in initial iron concentration. Transformation of γ -Fe₂O₃ to α -Fe₂O₃ (hematite) was studied by calcining the precursor in the temperature range of 500 to 850°C. Formation/transformation of phases at different temperatures was confirmed by FTIR and XRD studies. Images, obtained by SEM and TEM showed the morphology and nanocrystal formation of hematite. Room temperature rheological behaviour of the synthesized α -Fe₂O₃ nano powder has been studied.

Keywords: Iron oxide, emulsion, nanocrstals, crystal growth, rheological property

1. Introduction

Among the various forms of iron oxides, maghemite (γ -Fe₂O₃) and hematite (α -Fe₂O₃) are of great importance in technological and industrial applications (Huo *et al.*, 2000). Maghemite has numerous applications like recording, memory devices, magnetic resonance imaging, drug delivery or cell targeting (Laurent *et al.*, 2008; Daou *et al.*, 2010). Hematite exhibits high resistance to corrosion, therefore, it has been extensively used in many fields which include photo-anode for photo assisted electrolysis of water. It is an active component of gas sensors, catalyst, lithium ion battery, pigments and oxidizer in thermite composition (Kodama *et al.*, 1997; Fischer and Grubelich, 1998; Dong and Zhu, 2002; Prakash *et al.*, 2004; Gupta and Gupta, 2005; Figuerola *et al.*, 2010). It is also used in magnetic fluids, also called ferrofluids, for damping in inertial motors, shock absorbers, heat transfer fluids, etc (Wang and Meng, 2001; Espi'n *et al.*, 2004). All these engineering applications require study of rheological behavior. Because of extensive applications, nano maghemite and hematite have attracted much attention for their synthesis and applications.

Various methods have been developed to produce maghemite and hematite with controlled shape, size, and size dispersion (Daou *et al.*, 2010). Micronic acicular maghemite was prepared either directly from FeOOH or through a topotatic oxidation of magnetite (Bate, 1975). Maghemite nanoparticles have been obtained by co-precipitation of FeCl₃ and FeCl₂ in an alkali medium, (Massart, 1981), microemulsion method with ionic surfactants, (Tueros *et al.*, 2003) or physical methods such as high-energy ball-milling (Randrianantoandro *et al.*, 2001). Hematite nanoparticles with different structures have also been synthesized by various methods, such as the sol–gel process (Woo *et al.*, 2003), thermal oxidation of iron metal (Wen *et al.*, 2005), forced hydrolysis (Musi'c *et al.*, 2003), hydrothermal method (Cao *et al.*, 2005; Giria *et al.*, 2005; Baruwati *et al.*, 2006; Jia *et al.*, 2007), gas-phase flame synthesis (Kumfer *et al.*, 2010), ultrasonic spray pyrolysis (Gurmen and Ebin, 2010). Jia *et al.* (2005)) prepared monodispersed α-Fe₂O₃ single-crystal nanotubes by a facile hydrothermal method (Jia *et al.*, Chen *et al.* (2005) synthesized α-Fe₂O₃ nanotubes by a templating method and studied their gas sensor and lithium-ion battery applications. Gupta and Gupta (2005)

prepared various nano iron oxides for biomedical applications. Vayssieres and co-workers investigated the one-dimensional quantum-confinement effect in α -Fe₂O₃ ultrafine nanorod arrays (Vayssieres *et al.*, 2005). The present study is aimed at synthesizing nano maghemite and hematite using simple and facile emulsion-precipitation followed by calcination method.

The inverse spinel crystal structure is the most general case for maghemite (Pecharroman *et al.*, 1995; Glotch and Rossman, 2009). General formation of hematite is found to be rhombohedrally hexagonal. Phase transformation of α -Fe₂O₃ from other forms of iron oxides, hydroxide and oxy-hydroxides has been reported by many researchers. The formation of Fe(OH)₃ by precipitation of Fe³⁺ salt and its transformation to β -FeOOH (akaganeite) and then to α -Fe₂O₃ was proposed by Cao *et al.* (2006). Phase transformation from β -FeOOH to α -FeOOH (goethite) to α -Fe₂O₃ in alkaline medium was reported by Cornel and Schwertmann (1996). It is generally accepted that the phase transformation from β -FeOOH to α -FeOOH and then to α -Fe₂O₃ occurs via the dissolution precipitation mechanism (Music *et al.*, 1997) in aqueous medium. Dar *et al.* (2005) investigated the formation of α -FeOOH to α -Fe₂O₃. Evaluation and investigation of various phases of iron oxides as a function of heat treatment time and temperature has been reported by Deb and Basumallick (2004). Increase in holding time at 250°C caused transformation of Fe₃O₄ (magnetite) to γ -Fe₂O₃ and from γ -Fe₂O₃ to α -Fe₂O₃. Zhao *et al.* (2007) investigated the phase transformation of γ -Fe₂O₃ to α -Fe₂O₃ and found that such a transformation occurred at 770 °C with an exothermic peak in DSC curve. Yen *et al.* (2002) studied the crystallite size variation on formation of α -Fe₂O₃ from γ -FeOOH (lepidocrocite), Fe₃O₄ and amorphous Fe(OH)₃ to α -Fe₂O₃. Effect of calcination temperature on crystallite size and crystal growth of iron oxide nano-crystallites was studied by Praserthdam *et al.* (2003).

In view of the importance of nano maghemite and hematite, the present investigations were undertaken to synthesize maghemite nano-crystallites by a facile emulsion method and their transformation to nano-crystals of hematite. Surfactant mediated precursors were prepared at two initial iron concentrations and effect of calcination temperature on phase transformation was followed. The synthesized α -Fe₂O₃ has been used to prepare its colloidal solution (ferrofluid) of different concentrations with a view to study its rheological behavior.

2. Experimental

Anhydrous FeCl₃, NaOH, SPAN-80 (all of Analytical grade) and kerosene were used without further purification. 200 mL of FeCl₃ solution of 0.5M was taken in a 1000 mL beaker. 2g of SPAN-80 was added with vigorous stirring and the contents were stirred for 15 minutes. 4 mL kerosene was added and stirring was continued for another 30 minutes. The system transformed to turbid emulsion of milky brown colour. NaOH solution (5M) was added drop wise till the pH reached up to 10-10.5. Reddish brown precipitate of iron hydroxide started forming from pH 3 onwards. Precipitate became thick as pH increased. After reaching the desired pH, addition of NaOH was stopped and the stirring was continued for 1h. The contents in the beaker were then left undisturbed for 30 min to allow the precipitate to settle. The precipitate was filtered and washed with distilled water till free of ions and excess alkali. Then it was washed with acetone to remove kerosene and surfactant and kept in an oven at 100 °C overnight. The dried precursor was subjected to calcination at various temperatures i.e. 250, 500 and 850 °C for two hours. After cooling and drying, final product was ground with agate mortar to get fine powder. The sample details and calcination temperature are given in Table 1. Similarly for 0.1M FeCl₃ solution (lower concentration), 1M NaOH was used following the same method.

Table 1. Conditions of sample preparation and their coding

Serial No.	Sample	Calcination temp.	Initial FeCl ₃	Surfactant %	Oil %
		(°C)	Conc (M)		
1	A	100 (oven dried)			
2	A250	250	0.5	1	2
3	A500	500			
4	A850	850			
5	В	100 (oven dried)			
6	B250	250			
7	B500	500	0.1	1	2
8	B850	850			

Infrared spectra of samples were recorded on a Thermonicolet FTIR spectrophotometer (Model Nicolet 5700, make USA) using KBr matrix. XRD patterns were obtained using (Model X'Pert PRO make PANalytical, Netherland) with $CuK\alpha$ source. Particle morphology was studied by obtaining micrographs using Carl Zeiss, EVO50 low vacuum SEM, Germany. TEM and and HRTEM were obtained using FEI make Technai F-30, the Netherland, with 300kV field emission gun,.

Three types of ferrofluids were prepared by dissolving 50 mg, 100 mg and 300 mg each of α -Fe₂O₃ (B850) in 100 ml of ethylene glycol (EG). These solutions were sonicated for 1 h and subsequently 0.5, 1, and 3 ml of acetylacetone (acac) was added and sonicated again for 10 min. The rheological behavior of the suspensions was investigated using Rheologica Nova Rheometer with 40 mm plate-plate geometry at 25 °C temperature.

3. Results and discussion

3.1 FTIR Analysis

Figure 1 shows the FTIR spectra of iron oxide samples of series A and sample B850. From the spectra it can be seen that no strong bands were identified for sample A. Two broad peaks at 555 and 463 cm⁻¹ were identified for sample A250. These peaks may correspond to Fe-O stretching and bending vibration mode, respectively of γ -Fe₂O₃ (Kim *et al.*, 2010). These two peaks became slightly stronger for sample A500, which may correspond to a partial vacancy ordering in the octahedral positions in the maghemite inverse spinel crystal structure (White and Deangelis, 1967; Faria *et al*, 1997; Millan *et al*, 2007). The characteristic absorption bands of sample A850 at 535 and 462 cm⁻¹ and for sample B850 at 538 and 466 cm⁻¹ are assigned to Fe-O stretching and bending vibration mode of α-Fe₂O₃ respectively (Zhao *et al.*, 2007).

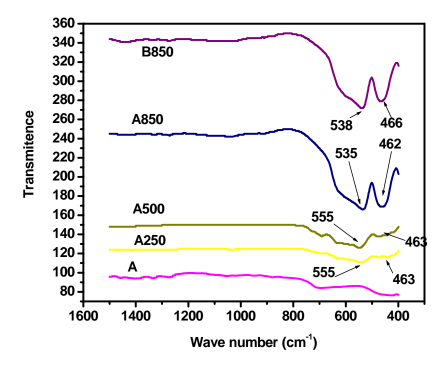


Figure 1. FTIR spectra of iron oxide samples series A and B850.

3.2 XRD analysis

XRD spectra of the A series samples are presented in Figure 2 and those for B series samples are given in Figure 3. The peaks observed at d values of 2.95, 2.51, 2.08, 1.60, 1.47 Å in sample A correspond to cubic phase of maghemite, γ -Fe₂O₃ (matched with PDF No- 01-089-5892).) XRD peaks of sample A250 are almost similar to those of precursor A showing better crystallinity. Sample A500 contains more intense peaks of cubic γ -Fe₂O₃ along with weak α -Fe₂O₃ (peaks marked as 'a'). Sample A850 contains peaks of rhombohedral α -Fe₂O₃ at d-values of 3.62, 2.67, 2.49, 2.19, 1.83, 1.68, 1.59, 1.48, 1.44, 1.30 Å (matched with PDF No-01-079-007). The results show that the crystal formation of hematite phase started at 500 °C. Thus, A500 is the state of both γ-(major) and α -Fe₂O₃ (minor as indicated in figure). γ -Fe₂O₃ particles have cubic unit cells with both octahedrally and tetrahedrally coordinated Fe³⁺ sites (defect spinel structure) whereas the unit cell of α -Fe₂O₃ is hexagonal and contains only octahedrally coordinated Fe³⁺ atoms (corundum structure). Complete phase transformation occurs at 850°C. Crystallographic phases obtained for sample B series are also similar to those of sample A. The difference is that in case of B500, α -Fe₂O₃ peaks were not found. Thus B500 contains peaks of mainly γ -Fe₂O₃. This data helps to synthesize pure γ -Fe₂O₃ for further application and confirms that phase transformation at lower concentration occurs after 500°C.

The values of lattice constants, phase present and crystallite size of various samples are listed in Table 2. Average crystallite size is calculated using broadening of most intense peaks from XRD pattern and Scherrer equation. The line broadening is due to grain surface relaxation of nanocrystalline powders (Scardi, 1999; Leoni *et al.*, 2004; Maensiri *et al.*, 2007). Lattice parameters 'a' and 'b' show marginal decrease with increase in calcination temperature from 250 to 500°C but abrupt decrease is observed at 850 °C with the formation of α -Fe₂O₃ Sudden increase in lattice parameter 'c' is observed with the formation of pure α -Fe₂O₃ phase. In

case of crystallite size of nanoparticles, with increase in calcination temperature, marginal increase in crystallite size was observed upto 500 °C and with the formation of α -Fe₂O₃ at 850°C, the crystallite size increased from 25.8 to 42.3 nm. Similar results were obtained for B series of samples for lattice parameters.

There is a lot of difference in the particle size of precursor sample A and B with their MCD values as 2.1 and 21.2 nm respectively. It may be explained as follows: an emulsion is composed of two phases, namely water, oil and a mixing agent i.e., surfactant. In oil-in-water emulsion, volume of water phase is more than oil phase. The materials were synthesized with very low concentration of oil and surfactant. Due to vigorous stirring, oil phase is transformed into tiny micro-droplets stabilized by surfactant within the water phase and move very fast. These moving droplets act as collider. At the time of precipitation, the precipitated iron hydroxide is collided by these fast moving oil droplets. Continuous and very fast collision leads to break the hydroxide into nanoparticles. By increasing the number of collisions, smaller particle size may be obtained. As the concentration of iron hydroxide decreases, probability of number of collision per molecule/particle increases and particle size becomes smaller. Thus the precursor sample B is much smaller (2.1nm) when compared to the precursor sample A (21.2nm).

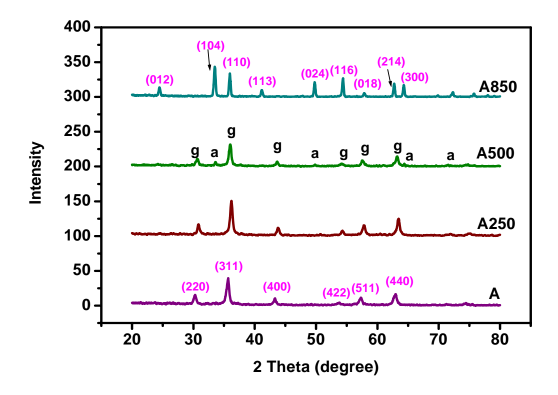


Figure 2. XRD patterns of sample A, A250, A500 and A850. (g: γ -Fe₂O₃ and a: α -Fe₂O₃).

Table 2. Crystallographic data obtained from XRD

Sample	Phases present	Lattice parameters (Å)			$d_{ m XRD}$ (nm)
		а	b	С	,
A	γ-Fe ₂ O ₃	8.3363	8.3363	8.3363	21.2
A250	γ -Fe ₂ O ₃	8.2680	8.2680	8.2680	24.5
A500	γ -Fe ₂ O ₃ , α -Fe ₂ O ₃	8.2331	8.2331	8.2331	25.8
A850	α -Fe ₂ O ₃	4.9876	4.9876	13.6114	42.3
В	γ -Fe ₂ O ₃	8.5220	8.5220	8.5220	2.1
B250	γ -Fe ₂ O ₃	8.2915	8.2915	8.2915	8.6
B500	γ -Fe ₂ O ₃	8.2341	8.2341	8.2341	35.4
B850	α -Fe ₂ O ₃	4.9584	4.9584	13.5070	58.5

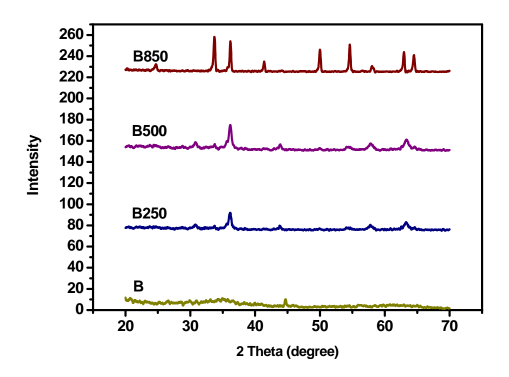
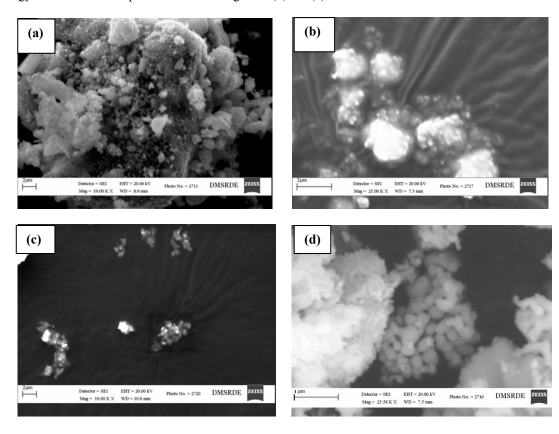


Figure 3. XRD patterns of sample B, B250, B500 and B850.

3.3 SEM and TEM Analysis

The morphology of iron oxide samples are shown in figures 4(a) to 4(e) as obtained from SEM and TEM.



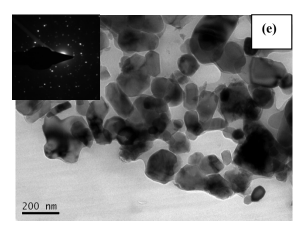


Figure 4. SEM micrograph of (a) Sample A, (b) Sample A500, (c) Sample A850, (d) Sample B850 and (e) TEM image of Sample B850.

Sample A is irregular, bulkier and agglomerated. Increasing calcination temperature resulted in particles becoming more fine and uniform as observed from figures 4(b) and 4(c)]. Fig 4(d) and 4(e) show the SEM and TEM micrographs, respectively for sample B850. The particles are well dispersed, very fine and hexagonal single crystals with particle size of ~70-100 nm. The selected area electron diffraction (SAED) pattern of the sample shows clear diffraction spots (inset in figure 4e), which indicates the high crystallinity of hematite nanoparticles (Kim *et al.*, 2010). The average crystallite size of B850 was found 58.5 nm from XRD, which is in good agreement with TEM result. The well dispersed and single crystal particles of nano hematite (B850) are because of growth of bigger crystals with less strain among the particles at the time of synthesis. Surface energy and lattice strain play important role for synthesis of nanoparticles and growing of crystal (Yen *et al.*, 2002).

3.4 Rheological behavior

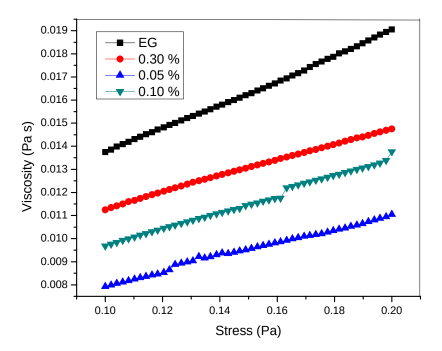


Figure 5. Variation of viscosity with stress for α -Fe₂o₃ nanoparticles in ethylene glycol (EG) for different concentrations.

The rheological tests were carried out for B850 ferrofluids with stress varying from 0.2 to 0.1 Pa to avoid the effects due to inertia yielding shear rate of 1 to 50 s⁻¹ and secondary flows due to the electro rheological ferrofluids (ERF) having very low concentration. All the prepared ferrofluids show linearity in nature, which indicate Newtonian behavior of the ferrofluids (Espi'n et al., 2004).

4. Conclusions

The following conclusions are drawn from the above study:

- 1. Nano crystalline α and γ -iron oxides were synthesized by emulsion assisted precipitation-calcination method.
- 2. The as synthesized precursor samples were calcined at 250, 500 and 850 °C. The inverse spinel maghemite phase transformed completely into rhombohedral hematite at a calcination temperature of 850 °C. With increase in calcination temperature, average crystallite size increased and lattice parameters 'a' and 'b' decreased. The lattice parameter while 'c' decreased upto a calcination temperature of 500 °C followed by increase for samples calcined at 850 °C. Also for lower initial iron concentration, pure maghemite phase was formed at a calcination temperature of 500 °C.
- 3. Scanning electron micrographs clearly showed that with the increase in calcination temperature, agglomeration decreased. A typical TEM image confirmed the well dispersed particles of hematite obtained when lower initial concentration (0.1M) of iron salt was used during synthesis.
- 4. From rheological study, it was observed that the prepared α -Fe₂O₃ based ferrofluids show Newtonian behavior.

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