Preparation and characterization of uniform pseudocubic hematite particles by utilizing polyethylene oxide polymers in forced hydrolysis reaction

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The monodispersity of pseudocubic hematite particles, produced from a forced hydrolysis reaction of acidic FeCl₃ solution, were controlled by using five kinds of polyethylene oxides (PEO; AlkoxS, Mesi Chemical Works, Ltd.) (0–2 wt.%). AlkoxS possesses nominal formula of (CH₂–CH₂–O)ₙ. The fairly uniform pseudocubic particles were precipitated for PEO with low Mₗ and narrow molecular weight distribution. However, POEs with large molecular weight and wide molecular weight distribution produced polydispersed pseudocubic hematite particles. The PEO with the largest molecular weight attained nearly spherical with large particle size distribution. From the calculation of crystallite size, all the pseudocubic hematite particles were regarded as a polycrystal as well as the large spherical hematite particles produced without AlkoxS (control system). This polycrystallinity of the particles provided an evidence that the particles are grown by aggregation of polynuclear (PN) primary particles. The specific surface areas of the particles (SSA) were ranged in 4–10 m²/g and no large difference can be recognized among five kinds of POEs. Since the measured SSA values were corresponded to the SSA one calculated by assuming monodispersed cubic particle with 300 nm in edge length, the pseudocubic hematite particles are non-porous. The time resolved XRD, TEM, ICP-AES and TOC measurements employed on the systems produced for pseudocubic particles elucidated that the pseudocubic crystal habit was formed by the specific adsorption of PEO molecules together with chloride ions and/or chloro ferric complexes to the {012} faces, restraining the growth process through stacking of ultrafine PN particles in the direction of normal to the {012} faces but strictly restricting the growth and mutual fusion of PN ones. The uniform pseudocubic particles found to be effective photocatalytic material than spherical and/or polydispersed pseudocubic hematite ones due to their large size with uniform flat crystal faces.

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1. Introduction

Recently design and synthesis of the morphology-controlled colloidal nanoparticles have been intensively investigated for their size and shape-dependent properties. Hematite (α-Fe₂O₃) represents the most important ore of iron and is the most stable iron oxide under ambient conditions. It has been profoundly investigated, therefore, due to its environmental safety and potential applications on gas sensor, catalysis and magnetic storage devices. Nowadays, various well-defined simple shapes of hematite particles, such as rods, wire, belt, tubes, ellipsoids, rings and spindles, has been obtained by vapor-phase processes and solid-based wet method such as a forced hydrolysis method. Our research group has been reported on the formation of spherical hematite particles by the forced hydrolysis reaction under 100°C. The authors revealed that the spherical hematite particles are formed by aggregation of fine polynuclear (PN) primary particles produced after dissolved β-FeOOH particles, which appeared at the initial stage of the reaction. Similar aggregation mechanism was also reported on the formation of micrometer-sized cubic hematite particles.

In the past decade, our research group has extended the investigation on the formation of hematite particles from the forced hydrolysis reaction of acidic FeCl₃ solution in the presence of polyethylene oxide series polymers (polyacrylamide, PAAm and polyacrylic acid, PAAc) with a high complexity to Fe(III) ions and a high adsorption affinity on both β-FeOOH of initial hydrolysis products of FeCl₃ aqueous solution and PN particles, the authors found that these polymers strongly influence on morphology and structure of hematite particles even at extremely low concentrations (less than 4 × 10⁻³ wt.%). The shape of the particles produced with PAAc was ellipsoidal though it was spherical for those formed with PAAm. These findings reveals that the particle shape strongly depends on the adsorption mode and strength of polymers onto β-FeOOH, PN primary and growing hematite particles regardless of the molecular weight of additives. Based on these results, we investigated further the effects of diblock co-polymer with terminal amino groups, [CH₂–(PEO)ₙ–(PPO)ₚ–NH₂], so called as Surfonamine. Surfonamine with the highest total amine content (PEO contents were over 80 mol %) gave spherical but those with lower total amine contents (PEO contents less than 76%) produced pseudocubic hematite particles. However, the uniformity in size of pseudocubic particles was low due to their low molecular weight of Surfonamine employed (600–3000).

It is well known that the growth and/or inhibition of particular crystal faces of the particles usually attain anisotropic particles such as cubic and rod-like shapes. To develop uniform cubic hematite particles, PEO polymers with large molecular weight (larger than 1 × 10⁶) are desired because there are large numbers of
of hydrophilic PEO units of those induce a strong adsorption onto \( \beta \)-FeOOH, PN primary and growing hematite particles. Since the orientation of the uniform cubic particles is ready to take place on the matrix, it is expected to develop new nano-composites. The aim of the present study is to disclose this point. The present study will describe the fundamental information for producing uniform pseudocubic hematite particles and their physicochemical properties. The application of these as-prepared uniform pseudocubic hematite particles was also demonstrated on a photocatalysis reaction.

2. Experimental procedure

2.1 Synthesis of hematite particles

Hematite particles were prepared from the forced hydrolysis of mixed solution of FeCl₃ and HCl dissolving various amounts of PEO (0–2 wt.%). This procedure was almost the same as employed in our previous paper. The total concentrations of FeCl₃ and HCl in a 30 mL solution were fixed at 3.12 × 10⁻² and 9.60 × 10⁻³ mol dm⁻³, respectively. This condition without PEO provides fairy uniform spherical particles of ca. 600 nm in diameter.¹⁶) Pyrex glass vial containing each 30 mL of reacting solution was tightly closed with a Teflon-lined screw cap and stood in a conventional forced-air oven at 100°C for 7 d. Five kinds of PEO [Alkox\(^{3,6}\) (CH₂₇-C₂H₇-O)₉], kindly supplied from Meisei Chemical Works Ltd. (Kyoto, Japan), as listed in Table 1, were used in this study. The number (Mₙ)-, weight (Mₘ)-average molecular weights and their ratio (Mₘ/Mₙ) were listed in Table 1. The molecular weight distribution curves of PEOs are shown in Fig. 1. All the PEOs possess larger Mₘ values, larger than 1 × 10⁴ and its order can be noted as follows, L-11 < R-150 < R-400 < R-1000 < E-60. The L-11 has a narrow and sharp distribution curve, suggesting that the number of PEO units is considerably ordered. On the other hand, molecular weight distribution curves of the rest of L-11 are widely spread out from 1 × 10⁻³ to 10⁻⁵. Since the E-60 has the highest Mₘ, 10 times larger than L-11, the strong effect of E-60 onto particle formation can be presumed. The resulting particles were thoroughly washed by filtration with ultrapure water using 0.2 μm Millipore filter and finally dried in air oven at 40°C for 18 h. The solution pH before and after aging was monitored at room temperature. Guaranteed reagent grade chemicals from Wako Pure Chemical Co. Inc. (Osaka, Japan) were used without further purification. Ultrapure water from Elix (Millipore) were used for the preparation and washing of all particles. Fresh ultrapure water had conductivities ca. 0.06 μS/cm.

2.2 Characterization of the samples

The morphology and crystal structure of particles were investigated by transmission electron microscope (TEM; JEOL JEM-2100), simultaneous thermogravimetry and differential thermal analysis (TG-DTA; SII-7200), Fourier-transform infrared spectrometer (FTIR; Nicolet Protégé 460) embedded in KB pellets (0.2 wt.%) and X-ray powder diffraction (XRD; Rigaku Rad-RC, 40 kV, 120 mA) techniques. The concentrations of PEOs and Fe(III) ions were assayed by total organic carbon analyzer (TOC; Shimadzu TOC-VCSH) and inductively coupled plasma atomic emission spectroscopy (ICP-AES; SII SPS 3520UV-2) measurements, respectively. The specific surface area of the hematite particles was investigated by N₂ adsorption isotherms recorded by a computer-aided volumetric apparatus built-in-house at a boiling point of liquid-N₂. Prior to the adsorption, the samples were treated at 100°C under 1.0 × 10⁻³ Pa for 2 h.

2.3 Photocatalysis measurement

The photocatalytic properties of as-synthesized hematite particles were evaluated by using Rhodamine B (RB) used as a probe molecules as reported by Fang et al.²⁰) A suspension of hematite photocatalysts (3 mL, 5 mg/L) after dispersed by an ultrasonic agitation was added to a quartz cell (1.0 cm × 1.0 cm × 4.5 cm). The RB solution (50 μL, 1.5 mM) was injected into the quartz cell using a micropipet with continuous stirring under UV light irradiation of 365 nm (30 W). The degree of RB degradation was monitored by absorption at a wavelength of 533 nm using a Halogen light source (UV–vis: Hitachi Hi-Tech U-3900H) for 5–20 min intervals. Since this measurement takes only a few second, no variation of solution temperature was encountered. All the measurements were carried out at 25°C.

3. Results and discussion

3.1 Morphology and crystal structure

Figure 2 displays the TEM pictures of typical hematite particles precipitated by aging for 7 d at 0.8 wt.% of PEOs (L-11, R-400 and E-60) together with those produced without PEO (referred to control system). L-11 and R-400 produced pseudocubic particles though E-60 gave nearly spherical with large particle size distribution especially at high E-60 concentration. It is noteworthy that the fairly uniform pseudocubic particles can be seen for L-11 with low Mₘ value and narrow molecular weight distribution.

Since the PEOs are nonionic polymers, there is no large difference can be seen among the solution pHs. All the system exhibit pH drop from ca. 1.8–1.9 to 1.0–1.1 by hydrolysis of FeCl₃ molecules. The mean edge length and/or particle diameter (Lₚ) for the L-11, R-150, -400, -1000 and E-60 are displayed in Fig. 3(a). The Lₚ values of all the systems remarkably dropped

Table 1. Properties of polyethylene oxides (Alkox\(^{3,6}\)) employed

| Alkox\(^{3,6}\) | Number average molecular weight \((Mₙ)\) | Weight average molecular weight \((Mₘ)\) | \(Mₘ/Mₙ\) |
|----------------|------------------------------------------|------------------------------------------|----------|
| L-11           | 7.60 × 10⁷                               | 11.6 × 10⁷                               | 1.53     |
| R-150          | 2.41 × 10⁹                               | 14.4 × 10⁹                               | 5.98     |
| R-400          | 2.98 × 10⁹                               | 20.2 × 10⁹                               | 6.78     |
| R-1000         | 5.70 × 10⁹                               | 29.6 × 10⁹                               | 5.19     |
| E-60           | 1.97 × 10⁷                               | 10.0 × 10⁷                               | 5.10     |

a) Alkox\(^{3,6}\) from Meisei Chemical Works Ltd. (Kyoto, Japan)

Fig. 1. Molecular weight distribution curves of polyethylene oxides employed measured by GPC method.

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from 540 to 300 nm after addition of 0.05% PEOs but their reduction was less up to 2 wt.% PEO (L_p is ca. 260 nm). There is no large difference among five kinds of PEO molecules can be seen in L_p values despite there are large difference in particle size distribution for these systems. To make clear the effect of PEOs onto the particle size distribution of particles produced, particle size distribution curves were made by counting over 300 particles for the systems produced with 0.8 wt.% PEO in Fig. 4. The median, mode, number mean diameter, length mean diameter and standard deviation are listed in a table in Fig. 4 (bottom). It is easy to recognize that particle size distribution curves of L-11 is sharp but they are spread out by increase in M_w of PEOs (R-150 < R-400 < R-1000 << E-60). This result indicated that the uniformity in size of pseudocubic hematite particles strongly depends on the molecular weight distribution of PEO; the nearly monodispersed pseudocubic hematite particles can be precipitated by using L-11 with narrow molecular size distribution. It can be presumed from above results that monodispersed pseudocubic hematite particles with 260 nm in diameter can be produced if we can acquire the PEO molecules with single molecular weight of \(1 \times 10^5\).

**Figure 5** displays the XRD patterns of the particles precipitated with L-11 and R-150 providing pseudocubic particles. Clearly, all the particles exhibit characteristics of hematite (JCPDS33-664). Similar results were obtained for the other PEO systems. However, the normally observed strong peak of the \{012\} face (2\(\theta\) = 24.2°) becomes weak by increase in the concentrations of L-11 and R-150 systems. Hence, the relative area intensity of the \{012\} peak was calculated and plotted as a function of concentrations of PEOs (abbreviated as [PEO]) in Fig. 6. Here the area intensity of the control system was assumed as unity. The relative area intensities are decreased for all the systems. This decrease in the \{012\} peak corresponds to the results of Surfonamine system,19) i.e., the pseudocubic crystal habit is thought to be formed by the specific adsorption of PEO molecules together with chloride ions and/or choloro ferric complexes to the \{012\} faces, restraining the growth in the directions normal to the \{012\} face.21)

The crystallite sizes of the particles (L_{104}) were evaluated using the Scherrer equation from the half height width of XRD peak of \{104\} face (2\(\theta\) = 33.2°). The results are shown in Fig. 3(b). Clearly, the crystallite sizes are decreased with increase in the
PEO concentration in each system and L_{104} are smaller than L_p of the corresponding particles in Fig. 3(a), indicating that all the particles can be regarded as a polycrystal.

The TG–DTA curves of the particles produced with PEOs are shown in Fig. 7. The curves contain two mass loss steps around at 25–600°C and 600–1000°C. Clearly the former mass loss is increased by raising the [PEO]. This mass loss can be ascribed to dehydration and dehydroxylation from the particle surface. The mass loss step at 600–1000°C is derived by elimination of OH\(^-\) ions included in the hydrohematite particle. In hematite, O\(^2-\) may be partly replaced by OH\(^-\) ions accompanying Fe(III) vacancies to provide charge balance of the particles. These anion-substituted hematomles are called as hydrohematite or protohematite by Wolska and Schwertmann.\(^{22}\) The TG–DTA curves of the particles produced with E-60 only exhibit a large weight loss at 200–400°C together with an exothermic peaks, indicating that the E-60 molecules remained in these particles. This result coincides with the result of TOC as will be described later.

### 3.2 Specific surface area

The adsorption isotherms of N\(_2\) for all the samples produced were shown the type II isotherms in the IUPAC classification.\(^{23}\) The specific surface areas of the particles were estimated by fitting the adsorption isotherms of N\(_2\) to the BET equation and plotted as a function of [PEO] in Fig. 8. All the SSA values are
ranged in 4–10 m$^2$/g and no large difference can be recognized. By assuming monodispersed cubic particles with $L_p = 300$ nm with 5.2 g/cm$^3$ density, SSA value can be calculated as 3.8 m$^2$/g. The measured SSA values are almost corresponded to the calculated SSA one, indicating that the pseudocubic hematite particles are non-porous. In the case of Surfonamine, the molecules remained between ultrafine PN particles adsorbing chloride ions and/or chloro ferric complexes on their surfaces and produced highly uniform micropores with diameters of 0.7 nm. This difference between Surfonamine and PEO employed in the present study can be attributed to the difference in adsorption ability of those polymers, i.e., adsorption ability of PEO is lower, especially at high temperature, than Surfonamine with terminated amino groups even though they have large molecular weight.

### 3.3 Formation mechanism of pseudocubic hematite particles in the presence of PEO

The authors reported that pseudocubic shape appers to be characteristic to hematite particles prepared in solution containing high concentration of chloride ions (>$0.1$ M). We revealed that the dramatic shift of the isoelectric point from about 8.5 to 3.0 due to irreversible incorporation of chloride ions in the surface layers of pseudocubic hematite particles by a forced hydrolysis reaction though electrokinetic and XPS measurements after thoroughly washing the particles with alkaline solution. The pseudocubic crystal habit is thought to be formed by the specific adsorption of chloride ions and/or chloro ferric complexes to the Miller faces, retaining the growth in the direction of normal to the Miller faces as described before. Similar mechanism was also reported by a forced hydrolysis reaction of condensed ferric hydroxide gel, so called as gel–sol technique. Hence it can be considered that the pseudocubic hematite particles were formed by the specific adsorption of PEO molecules together with chloride ions and/or chloro ferric complexes to the Miller faces. The preparation of pseudocubic hematite particles from diluted alcoholic solutions of ferric chloride was also reported by Hamada and Matijević. They concluded that alcohol acts as an impurity, destroying the normal interfacial structure and providing an easier transition from solution to crystal thus affecting the growth rate.

To disclose the mechanism of the formation of pseudocubic hematite particles in the present study, we traced on the systems
with 0.8 wt.% of PEOs. In this experiment, we measured the time resolved TEM pictures together with the concentrations of Fe(III) ions (abbreviated as [Fe(III)]) and PEO (abbreviated as [PEO]) in a supernatant at different maturation time by TEM, ICP-AES and TOC, respectively. The hematite fraction in the precipitates was obtained and the results were shown in Fig. 9. Here the area intensity of fundamental vibration band of hematite at 580 cm\(^{-1}\) obtained and the results were shown in Fig. 9. The control system (\(\bullet\)) slowly produced spherical particles and the phase transformation from \(\beta\)-FeOOH to hematite was almost completed after aged for 5 d as well as reported in the previous paper.\(^{18}\) However, the addition of PEOs enhanced the phase transformation, i.e., the phase transformation from tiny rod-like \(\beta\)-FeOOH to pseudocubic hematite particles was completed after aging for 2 d, indicating that these PEO molecules enhance the particle growth. This is due to the formation of fine \(\beta\)-FeOOH particles, i.e., these particles are readily dissolved and transferred to hematite phase though the TEM pictures are not shown.

The changes of [Fe(III)] and [PEO] as a function of aging time were shown in Figs. 10(a) and 10(b), respectively. In the case for the control system in Fig. 10(a), [Fe(III)] decreases very rapidly within 6 h after started aging and slightly increases up to 12 h due to the dissolution of \(\beta\)-FeOOH particles (\(\bullet\)). After aging for 1 d, [Fe(III)] decreases again and reaches a constant value (ca. 5 mmol/dm\(^3\)), but it reduces again remarkably after aging over 5 d. This rapid reduction of [Fe(III)] (secondary step) coincides with the phase transformation of \(\beta\)-FeOOH to hematite as is confirmed in Fig. 9. This agreement between secondary reduction step of [Fe(III)] and completion of phase transformation may indicate that the dissolution of \(\beta\)-FeOOH particle is a rate-determining step in the reaction as the authors reported in a previous paper.\(^{15}\) On the other hand, [Fe(III)] values for the systems with PEOs within 12 h are lower than the control system, indicating that the hydrolysis reaction of Fe(III) ions are accelerated at the initial stage of reaction. This acceleration may be attributed to the transfer of hydrated water from Fe(III) ions to PEO molecules in order to promote PEO dissolution at high temperature. The [Fe(III)] values after 12 h, except for E-60 system (X), decreased steeply rather than the control system. The [Fe(III)] values for the L-11 (\(\square\)) became to zero at 4 d, suggesting a fast phase transformation from \(\beta\)-FeOOH to hematite. This fast phase transformation may induce a monodispersed pseudocubic particles. On the contrary, the E-60 system showed a slow phase transformation exhibiting the high [Fe(III)] value. This extremely slow phase transformation is caused by the adsorption onto \(\beta\)-FeOOH and PN particles to inhibit both their dissolution and migration at 100°C. According to this slow phase transformation, the polydispersed hematite particles could be precipitated. The amounts of [PEO] curves in Fig. 10(b) indicates that PEO molecules, except for E-60, do not adsorbed on \(\beta\)-FeOOH particles and pseudocubic hematite ones because these systems exhibited no change in [PEO]. The PEO molecules adsorbed on \(\beta\)-FeOOH, PN and hematite particles during aging at 100°C by reducing their solubility, but PEO molecules could be readily desorbed by reducing the aging solution to room temperature and completely removed after filtration. On the other hand, only the [PEO] for E-60 was steeply decreased to one-third within 1 d. The desorption of E-60 with largest molecular weight was hardly to take place and remained on the hematite particle surface. The residual E-60 molecules was already confirmed by the TG–DTA curves as was shown in Fig. 7. From above results, it can be regarded the PEOs molecules, especially for the L-11, acted as a morphology control agent without remaining in/on the resulting particles.

### 3.4 Photocatalitic function

Fang et al. fabricated spherical hematite colloidal monocrystal cluster (CNCs).\(^{20}\) The hematite CNCs have 2 \(\mu\)m in diameter of secondary spherical particles and they are composed of numerous nanocrystals of approximately 20 nm in diameter. Since CMCs are aggregates of small nanocrystals, the CMCs have large SSA values of 36 m\(^2\)/g. Fang et al. reported that CMCs exhibit a high photocatalytic activity, i.e., almost 100% of RB was decomposed within 1 h by UV light irradiation. This high photocatalytic
growth and mutual fusion of PN ones. The uniform pseudocubic particles exhibit effective photocatalytic behavior owing to their large size with uniform flat crystal faces.

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