Structure and Morphology Properties of Nanoparticle Hematite Pigment from Lathe Waste

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Abstract

Hematite coated PEG was synthesized by the sonication-calcination method. A variation of Na₂CO₃ is investigated to know the effect on structure and morphology. Characterization of samples are using X-ray fluorescence, X-ray diffraction, Scanning electron microscopy, and color reader techniques. XRF data showed that iron is the highest element in the precursor. The X-ray diffraction data confirm that Fe(OH)₃, α-FeOOH, and Fe₂O₃ established at the sonication stage are then transformed into the α-Fe₂O₃ phase after calcination. The X-ray diffraction data also was found that α-Fe₂O₃ at 0.5 M formed with the highest crystallinity degree. The scanning electron microscopy showed that the particle's shape is spherical, bar-shaped, and aggregate. However, the distribution of particles is not uniform and still displays agglomeration. The Color reader shows the highest degree of lightness obtained is at 1 M variation.

Keywords: nanoparticle, α-Fe₂O₃, lathe waste, Na₂CO₃, sonication

1. Introduction

1.1 Introduce the Problem

Nanoparticle hematite has attracted more attention and more developed in recent years for functional color properties. R. M. Cornell and U. Schwertmann (2003) explained that structure, size, and shape of particles could affect the color of hematite. Bilalodin et al. (2015) on the research explained that iron oxide of 926.4 nm particle size colored black, iron oxide of 72.2 nm particle size colored red, and iron oxide of 349 nm particle size colored yellow. Each iron oxide compound has different morphology distinguishable from one another. Based on Wei Wang et al's research (2008), the morphology of Iron oxide β-FeOOH is spherical with an amorphous structure and stacked as cubic hematite.

The synthesis of nanoparticle hematite pigment is influenced by several factors, including synthesis method, precursor type, and concentration of the precipitating agent. These factors will affect crystallinity, particle size, shape, and distribution of particles characterized by the differences in hematite color (Pramanik et al., 2005). Synthesis of iron oxide nanoparticle hematite has been carried out using several methods, including the precipitation-calcination route (Aji et al., 2007; Khoiroh, 2019), hydrothermal (He et al., 2016; Khalil et al., 2014), sol-gel (Dar et al. 2007), and ultrasonic sonochemistry (Hassanjani-Roshan et al., 2011). The sonochemistry is the easiest and most effective method for large-scale production with precise size control, high morphology, and crystallinity (Khalil et al., 2017). According to Firnando (2015), it was proven that ultrasonic waves in the sonochemistry method can separate aggregation and that the effect of sonication (ultrasonic) can produce a product of nanocrystal size. According to Khalil et al. (2017), using ultrasonic sonochemical methods has successfully produced monodispersed hematite nanoparticles with a uniform shape and particle size of about 14 nm.

Lathe waste contains high iron, which can potentially be used to produce hematite. Hematite can be synthesized from pure compounds or high-iron containing waste. Based on Sobirin et al., (2015) research, hematite synthesized from waste steel (mill scale) yielding product of orthorhombic structures. Therefore, it is counted to reduce pollution by processing lathe waste into hematite pigment (Khoiroh et al., 2011) (Dražić et al., 2017). Based on Blanco-Andujar et al., (2012) research, sodium carbonate is a suitable precipitating agent to be used at the synthesis of nanoparticle hematite. The average particle size obtained was about 10 nm, and the shape of the particles was cubic and uniform.

In this study, hematite was synthesized from lathe waste through the sonication-calcination method and coated with polymer just as polyethylene glycol (PEG) as a stabilizer. An attempt has been made to understand the effect of the concentration of precipitating agent Na₂CO₃ on nanoparticle formation. The precursor was characterized by XRF to
know the content of the metal. The products were well characterized by various characterization techniques such as X-Ray Diffraction (XRD), Scanning Electron Microscopy (SEM), and color reader.

2. Materials and Methods

2.1 Ferric (Fe³⁺) Precursors Preparation

The ferric precursor (Fe³⁺) was prepared by mixing 30 gr iron samples with 500 mL diluted nitric acid for 24 hours, then heated until forming a slurry. The presence of ferric ion was determined by qualitative test; the sample was dripped with NaOH 0.1 M, which carried out the red sediment that shown ferric ion.

2.2 Synthesis of Hematite Pigment

Ferric precursor was dissolved into 250 mL distilled water and then added with precipitating agent Na₂CO₃ of various concentrations; 0.5; 1; 2 M. The pH of the mixture was regulated to remain at 6 then the sample was heated at 70° C for 1 hour. The solution was cooled at room temperature then decanted. After that, it was filtered and washed with distilled water to pH 7 and dried at room temperature. Brownness solid was formed from this step, then calcined at 750°C as Figure 1. The iron oxide α-Fe₂O₃ nanoparticles are produced by the following solid-state reaction:

Figure 1. Synthesis of hematite through sonication method
2Fe(NO$_3$)$_3$ (aq) + 4H$_2$O (l) + 3Na$_2$CO$_3$ (aq) $\rightarrow$ 2Fe(OH)$_3$ (s) + 6NaNO$_3$ (aq) + H$_2$O (l) + 3CO$_2$ (g) (Eq. 1)

Fe(OH)$_3$ $\rightarrow$ FeOOH (s) + H$_2$O (g) (Eq. 2)

2FeOOH (s) $\rightarrow$ $\alpha$-Fe$_2$O$_3$ (s) + H$_2$O (g) (Eq. 3)

2.3 Characterizations

The composition of the precursor was characterized using X-R Fluorosense analysis. The crystalline structure and crystallite size were examined with X-ray diffraction (XRD) analysis using a Cu K$_\alpha$ (1,5418 Å) source (40 kV, 40 mA) from Philip X-Pert. The crystallite size was determined with the Debye Scherer formula. The synthesized hematite morphology was evaluated through a scanning electron microscope (SEM, FEI type Inspect-S50). Color characteristics were analyzed using a white Xenon lamp with D65 illuminate from Minolta CR-10. The value of C* and H$^o$ was calculated with the following formula (Cornell and Schwertmann. 2000):

\[ C^* = [(a^*)^2 + (b^*)^2]^{1/2} \]  \hspace{1cm} (Eq. 4)

\[ H^o = \tan^{-1}(b^*/a^*), \text{degree} \quad 0^\circ \leq H^o \leq 360^\circ \]  \hspace{1cm} (Eq. 5)

3. Results and Discussion

3.1 Characterization with XRF

The chemical composition of the precursor of the manufactured from lathe waste is shown in Table 1, which shows that iron element was found in form Fe$_2$O$_3$ as the highest content by 96.3 %, which means that Fe$_2$O$_3$ is the primary oxide in the precursor. The presence of oxide and another element because of the precursor is made from lathe waste, which contains another element in its preparation, although not exceed 0.9 %. The high content of iron shown that the lathe waste is the potential to synthesize to be hematite.

| Component | % Content |
|-----------|-----------|
| Fe$_2$O$_3$ | 96.3 |
| Eu$_2$O$_3$ | 0.81 |
| MnO | 0.47 |
| Br | 0.35 |
| SiO$_2$ | 0.81 |
| OsO$_4$ | 0.24 |
| Cr$_2$O$_3$ | 0.27 |
| CuO | 0.19 |
| P$_2$O$_5$ | 0.38 |
| CaO | 0.13 |
| La$_2$O$_3$ | 0.06 |
3.2 Characterization with X-ray Diffraction

The X-ray diffraction pattern of the sample (0.5 M) after sonication showed in Figure 2 displayed many phases in the sample, which is bernalite, goethite, and magnetite phase is formed. The characteristic of the bernalite phase is indicated by the presence of peaks at $2\theta$ (º) 23.2573 according to ICSD No. 73411, goethite ($\alpha$-FeOOH) at $2\theta$ (º) 14.6080; 26.8934; dan 36.2991 according to ICSD No. 159957 and magnetite ($\text{Fe}_3\text{O}_4$) at $2\theta$ (º) 35.2904 according to ICSD No. 158505 whereas the sharp peak at $2\theta$ (º) 19.1420 revealed the presence of polyethylene glycol (PEG) (Tu et al., 2019). PEG content on the surface of iron oxide is due to the addition of PEG during sonication, which aims to stabilize it so that the particles formed are uniform in size even though they are given ultrasonic energy.
The transformation of bernalite, gothite, and magnetite to hematite phase is displayed in the X-ray diffraction pattern in Figure 3. The peaks observed at $2\theta$ (24.16, 33.18, 35.64, 41.00, 49.48, 54.08)Å correspond to those of hexagonal of hematite with hkl (012) (104) (110) (113) (024) (116) (214) (300) (1010) (220) according to ICSD 66756, with a space group of $R\overline{3}c$. During calcination, high temperature and energy can transform bernalite, gothite, and magnetite into pure hematite phases, shown in the absence of another peak beside the hematite peak and high crystallinity indicated by sharp peaks. The highest crystallinity degree of hematite is achieved at hematite 0.5 M of Na$_2$CO$_3$, which is indicated all of the peaks of diffraction are sharper than others. The Na$_2$CO$_3$ precipitating agent's effect on the diffraction pattern of hematite can be possible due to the presence of anions such as carbonate ions during the synthesis process. The carbonate ion in Na$_2$CO$_3$ tends to precipitate in the form of molecules such as amorphous of Fe$_2$(OH)$_2$CO$_3$ or Fe(OH)$_3$ phase (Tangale et al., 2013).

The diffraction pattern was obtained by varying the concentration of precipitation agent at the synthesis after calcination and concluded that the peak intensity between one concentration and the other is different, as displayed in Figure 4. Based on Figure 4, the $2\theta$ positions for 1M and 2 M hematite have shifted toward the smaller $2\theta$; this indicates that the sample has a larger crystal size. A position shift of $2\theta$ allows changes in lattice parameters (Kumar et al., 2015). Data obtained also indicates that peak intensity gets higher as the Na$_2$CO$_3$ concentration smaller.
Table 2. Crystallite size of hematite

| Hematite | 2θ     | The crystallite size (nm) |
|----------|--------|---------------------------|
| 0.5 M    | 33.1936| 46.29                     |
|          | 35.6575| 59.89                     |
|          | 54.1232| 32.02                     |
| 1 M      | 33.1522| 40.85                     |
|          | 35.6193| 30.84                     |
|          | 54.0656| 65.92                     |
| 2 M      | 33.0886| 54.42                     |
|          | 35.5127| 35.22                     |
|          | 54.1115| 52.73                     |

The crystallite size of various samples is listed in Table 2. Crystallite size is calculated with The Debye Scherrer equation using broadening of the most intense peaks from the X-ray diffraction pattern of hematite. The smallest crystallite size of hematite synthesized is achieved at 1 M. Crystallite size can affect sample color. The smallest crystallite size exudes the lightest pigment color, as displayed in Figure 5. The visual color of hematite pigment shows that at 1 M variation, the pigment exhibits the lightest color. The pigment's color grows brighter as the crystal size becomes smaller (Sobirin et al., 2015).

3.3 Characterization Through Scanning Electron Microscope

SEM analysis result of hematite 0.5 M variation with 50,000 times magnification shows that particle shape distribution is relatively uniform, but agglomeration still occurs. The particle size of hematite 0.5 M variation is around 38.33 nm. The particle's shape at a magnification of 50,000 times is spherical, as seen in the SEM results in Figure 6. At 1 M variation, the synthesized hematite is bar-shaped with surface morphology seems uniform; however, aggregation still occurs with the particle size obtained is around 37.99 nm. Meanwhile, the 2 M variation's product after 50,000 magnification showed particles taking the shape of aggregate and not uniformly distributed with a particle size around 43.03 nm.
3.4 Characterization with Color Reader

Based on Table 3, the lightness degree of all concentration variation's product matches that of the standard with the highest lightness degree obtained at 1 M Na$_2$CO$_3$, which is possible as 1 M variation's product has the smallest crystal size. The values of a* and b* of the three concentration variation are positive; this indicates that the sample is dominantly red and yellow. The red color appeared is due to the presence of red iron oxide or hematite - the presence of peaks following those of hematite in the XRD results. Hematite with 1 M Na$_2$CO$_3$ exhibits the highest a* value; it is attributed to the degree of lightness and color purity of the product, which is higher than those of 0.5 M and 2 M variation.
Table 3. Optical data of hematite synthesized

| Color system | Hematite 0.5 M | Hematite 1 M | Hematite 2 M |
|--------------|---------------|--------------|--------------|
| $L^*$        | 33.8          | 35.1         | 33.7         |
| $C^*$        | 21.439        | 26.792       | 23.284       |
| $H^*$        | 33.385        | 34.059       | 35.716       |
| $a^*$        | 17.9          | 22.2         | 18.9         |
| $b^*$        | 11.8          | 15.0         | 13.6         |

The value of $H^*$ (color pattern) increases with the increasing of precipitating agent concentration. The hue value of the synthesized pigment is closer to the red spectrum due to hematite presence. It is evident from the peaks of the diffraction pattern appears at 20 (33.1). The chroma of the synthesized pigment is also directly proportional to its particle size, indicating that as the particle size gets smaller, it exudes a higher level of lightness and purity (Buxbaum, 2005). The higher the crystallinity level, the brighter the hematite color and the higher the color values have the smaller particle size. Based on the data of $L^*$, $C^*$, $H^*$, $a^*$, and $b^*$, the best color value is obtained at 1 M variation.

4. Conclusion

Nanoparticle hematite coated PEG was synthesized from lathe waste that contains 97% of iron using the sonication method has a hexagonal structure with space group R-3C. The highest crystallinity of hematite was obtained at 0.5 M but the highest lightness of color and degree of redness obtained at 1 M. The morphology of hematite was like spherical, bar-shaped uniform and aggregates. The highest lightness of color and degree of redness obtained at 1 M.

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1. Lightness of color
2. Chroma
3. Hue
4. degree of redness
5. degree of yellowness
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