

## CONTROLLED GROWTH OF IRON OXIDE MAGNETIC NANOPARTICLES VIA CO-PRECIPIATION METHOD AND NaNO<sub>3</sub> ADDITION

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Received: 13 January 2017

Revised: 13 March 2017

Accepted: 4 April 2017

### ABSTRACT

**CONTROLLED GROWTH OF IRON OXIDE MAGNETIC NANOPARTICLES VIA COPRECIPIATION METHOD AND NaNO<sub>3</sub> ADDITION.** Size controlled magnetic nanoparticle (MNPs) of iron oxide were prepared in the presence of NaNO<sub>3</sub> via co-precipitation method followed by HNO<sub>3</sub> peptization according to Massart's method. The MNPs size were reduced by addition of NaNO<sub>3</sub> in varied molarity and at different stage of process. As an end product, stable water-base colloids were formed. XRD pattern analysis using Rietveld method confirmed Fe<sub>3</sub>O<sub>4</sub>/γ-Fe<sub>2</sub>O<sub>3</sub> phase formation with nanoscale crystallite size. This crystallite size significantly decrease with NaNO<sub>3</sub> addition from 12 nm to smaller than 8 nm, and give end-result of decreasing magnetization as measured by VSM. Langevin fitting of magnetic hysteresis curve also revealed the magnetic core size of nearly the same behaviour. TEM results show bigger value for single magnetic nanoparticle of > 10 nm and < 10 nm for MNPs without and with NaNO<sub>3</sub> addition, respectively. However, PSA measurement still trace a low nanoparticle agglomeration of ~ 20 nm, even after surface peptization using HNO<sub>3</sub>. A possible mechanism is proposed to explain these characteristics formation especially of the MNP's size.

**Keywords:** Magnetic nanoparticle, Fe<sub>3</sub>O<sub>4</sub>, γ-Fe<sub>2</sub>O<sub>3</sub>, NaNO<sub>3</sub>, Size

### ABSTRAK

**PENGENDALIAN PENUMBUHAN NANOPARTIKEL MAGNETIK OKSIDA BESI DENGAN METODA KO-PRESIPITASI DAN PENAMBAHAN NaNO<sub>3</sub>.** Nanopartikel magnetik oksida besi dengan ukuran terkendali telah dipreparasi dengan metoda ko-presipitasi dan penambahan NaNO<sub>3</sub> serta peptisasi HNO<sub>3</sub> mengikuti metoda Massart. Pengendalian ukuran nanopartikel magnetik dilakukan dengan memvariasikan nilai molaritas serta tahapan proses penambahan NaNO<sub>3</sub>. Hasil analisis pola difraksi sinar-X dengan metoda Rietveld mengkonfirmasi terbentuknya fasa Fe<sub>3</sub>O<sub>4</sub>/γ-Fe<sub>2</sub>O<sub>3</sub> dengan penurunan ukuran kristalit dari 12 nm untuk sampel tanpa penambahan NaNO<sub>3</sub> menjadi 8 nm setelah penambahan NaNO<sub>3</sub>. Pengcilan ukuran ini juga teranalisis pada hasil *fitting* kurva magnetisasi dengan persamaan Langevin serta pengamatan dengan TEM. Namun hasil pengukuran PSA tetap mendeteksi adanya aglomerasi nanopartikel dengan ukuran aglomerat terkecil ~20 nm meski telah dilakukan upaya peptisasi dengan HNO<sub>3</sub>. Mengacu pada berbagai data karakteristik dan hasil analisis ini, mekanisme pengendalian pertumbuhan ukuran partikel ini dicoba dijelaskan dalam artikel ini.

**Kata kunci:** Nanopartikel magnetik, Fe<sub>3</sub>O<sub>4</sub>, γ-Fe<sub>2</sub>O<sub>3</sub>, NaNO<sub>3</sub>, Ukuran

## INTRODUCTION

Synthesis of iron oxide magnetic nanoparticles with small and uniform size is still a challenge for researchers to date, mainly motivated by the growing extent of the potential application of these nanoparticles [1]. Each application has a specific requirement for a magnetic nanoparticle to be used, either size, surface properties or its other physico-chemical properties [2].

Iron oxide nanoparticles can be synthesized by various methods including co-precipitation, microemulsion, thermal decomposition, and mechanical synthesis [3]. The co-precipitation method is more common method compare to the others due to its simple, save and economic procedure. The process of co-precipitation involves initial nucleation followed by slow growth as the solutes diffuse to the surface of the crystal. The size and other properties of the growth nanoparticles will be altered by some parameters including pH, temperature, reaction time, and iron ion ratios [4]. Normal co-precipitation method usually produce nanoparticles in the size range of 10 to 40 nm and the maximum magnetic saturation could reach almost 90 emu/gram. However, some application prerequest the nanoparticles with smaller size while maintaining its high magnetic saturation value, in the form of water-base stable colloid.

Stable colloid usually synthesizes following the Massart's method in which nanoparticle surface are peptized by acidic or alkaline surfactant such as  $\text{HNO}_3$ ,  $\text{HClO}_3$  and  $\text{TMAOH}$  [5]. In some research, for reducing nanoparticles size, counter ion was added to increase its ionic strength within precipitation process. Some type of counter ion usually used are  $\text{NaNO}_3$ ,  $\text{NH}_4\text{NO}_3$  or  $\text{N}(\text{CH}_3)_4\text{NO}_3$  which result in nanoparticle size smaller than 10 nm, even on the size as small as  $\sim 2\text{nm}$  [6]. But there is no clear explanation about its magnetic value and the mechanism of size reduction.

In this article, the mechanism of this size controlled was discussed and a model was proposed to get more complete understanding about the role of  $\text{NaNO}_3$  counter ion on the size reduction of magnetic nanoparticle. The data was collected from serial magnetic nanoparticles synthesized by addition of  $\text{NaNO}_3$  in varied molarity and at different stage of coprecipitation process, followed by peptizing with  $\text{HNO}_3$ .

## EXPERIMENTAL METHOD

### Materials

The chemical used in the synthesis of magnetic nanoparticles are  $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$ ,  $\text{FeCl}_3 \cdot 6\text{H}_2\text{O}$ ,  $\text{NaOH}$ ,  $\text{NaNO}_3$ ,  $\text{HNO}_3$  from Merck, Germany with pro analysis grade (pa) and deionized water of  $18.2\text{ M}\Omega\text{cm}^{-1}$ .

## Synthesis of Magnetic Nanoparticles

Magnetic nanoparticles were synthesized by co-precipitation method. Fe (II) and Fe (III) salt in a mole ratio of 1: 1 were dissolved in deionized water. Mixed solution of Fe-salt was added at once into  $\text{NaOH}$  solution of 3.2 M to nucleate the precipitation, continued with ultrasonic mixing for 5 minutes to ensure formation of nanoscale iron-oxide particles. For original sample, the process was continued with aging the solution overnight for enhancing crystallization process of iron oxide phase. Magnetic precipitates were separated from its supernatant by using permanent magnet. Peptizing of nanoparticles were done following the Massart's method by adding  $\text{HNO}_3$  to get acidic sol [5]. The process was done ultrasonically for 2 minutes, followed by decantation of supernatant by centrifugation at 9000 rpm for 10 minutes. To get end result of stable water-base magnetic colloid, the precipitates were dissolved in deionized water and ultrasonically stirred for 5 minutes. 2 mL of samples were evaporated to get dry powder sample for characterization.

Modified samples with  $\text{NaNO}_3$  were done in varied manner as listed in Table 1. Two different stages of  $\text{NaNO}_3$  addition were choosen, either mixed together with Fe(II)/Fe(III) salt before precipitation or added after precipitation of Fe(II)/Fe(III) with  $\text{NaOH}$ .

**Table 1.** List of the prepared magnetic nanoparticles

No.	Sample Code	Magnetic nanoparticle modification
<i>1<sup>st</sup>-series</i> , peptizing with varied $\text{HNO}_3$ molarity, without $\text{NaNO}_3$ addition		
1.	Na0H0	M ( $\text{HNO}_3$ ) = 0
2.	Na0H0.2	M ( $\text{HNO}_3$ ) = 0.2
3.	Na0H0.4	M ( $\text{HNO}_3$ ) = 0.4
4.	Na0H0.6	M ( $\text{HNO}_3$ ) = 0.6
<i>2<sup>nd</sup>-series</i> , fix $\text{HNO}_3$ molarity of 1M and varied molarity of $\text{NaNO}_3$ added before precipitation process		
1.	Na2H1B	M ( $\text{NaNO}_3$ ) = 2M
2.	Na4H1B	M ( $\text{NaNO}_3$ ) = 4M
3.	Na6H1B	M ( $\text{NaNO}_3$ ) = 6M
4.	Na8H1B	M ( $\text{NaNO}_3$ ) = 8M
5.	Na10H1B	M ( $\text{NaNO}_3$ ) = 10M
<i>3<sup>rd</sup>-series</i> , fix $\text{HNO}_3$ molarity of 1M and varied molarity of $\text{NaNO}_3$ added after precipitation process		
1.	Na2H1A	M ( $\text{NaNO}_3$ ) = 2M
2.	Na4H1A	M ( $\text{NaNO}_3$ ) = 4M
3.	Na6H1A	M ( $\text{NaNO}_3$ ) = 6M
4.	Na8H1A	M ( $\text{NaNO}_3$ ) = 8M
5.	Na10H1A	M ( $\text{NaNO}_3$ ) = 10M
<i>4<sup>th</sup>-series</i> , fix $\text{NaNO}_3$ of 6M molarity added before precipitation and varied $\text{HNO}_3$ molarity		
1.	Na6H0	M ( $\text{HNO}_3$ ) = 0
2.	Na6H0.2	M ( $\text{HNO}_3$ ) = 0.2
3.	Na6H0.4	M ( $\text{HNO}_3$ ) = 0.4
4.	Na6H0.6	M ( $\text{HNO}_3$ ) = 0.6

## Characterization of Magnetic Nanoparticles

Dry powder was characterized by X-ray diffractometer (XRD) with Cu-K $\alpha$  targets to identify the phases and nanoparticles crystallites size. Data were collected with XRD PANAnalytical model Empyrean at diffraction angle of 20° to 80°. Phases were identified using Rietveld analysis method implemented in RIETAN-2000 software package [7] using 2-phases assumption of Fe<sub>3</sub>O<sub>4</sub> (JCPDS No.82-1533, *space group Fd3m*) and  $\alpha$ -Fe<sub>2</sub>O<sub>3</sub> (JCPDS No. 39-1346, *space group P4<sub>1</sub>32*), while crystallite size of the nanoparticles were calculated using Debye-Scherrer equation. Morphology of static size of magnetic nanoparticles was observed using Transmission Electron Microscope (TEM) JEOL JEM 1400 for colloid samples diluted 5 folds before measurement. The size distribution of NPs in colloidal system were analyzed using Particle Size Analyzer Malvern NanoZetasizer also using 5-fold diluted samples. Magnetic hysteresis curve was measured using a Vibrating Sample Magnetometer (VSM) of OXFORD 1.2H with a measurement speed of 0.25T / min. The hysteresis curves were further fitted using Langevin function of superparamagnetic system for determining magnetic size of nanoparticles.

## RESULT AND DISCUSSION

### XRD Pattern Analysis

Generally, all samples displayed typical Fe<sub>3</sub>O<sub>4</sub>/ $\gamma$ -Fe<sub>2</sub>O<sub>3</sub> XRD pattern as shown in Figure 1, ensuring no other oxide and non-magnetic phase formation. The difference between original and after modified samples are only noticed in its peak broadening behaviour showing different degree of crystallinity of each sample. Figure 2 shows some of Rietveld refinement result using RIETAN2000 program for sample before and after modification. From this refinement, informations including type of phases, their % weight fraction and FWHM (full width half maximum) data could be obtained. This FWHM value contains information about crystallinity of the samples in which bigger FWHM value, more broader peaks and more amorphous the samples. Crystallite size of the sample could be calculated by Debye-Scherrer equation of:

$$D = \frac{k\lambda}{\beta \cos\theta} \dots\dots\dots (1)$$

- $k$  = A constant related to instrumental arrangement has a value of 0.916
- $\lambda$  = Wavelength of Cu-K $\alpha$  X-ray source equal to 1.5406 Å
- $\beta$  = Denoted for FWHM
- $\theta$  = Diffraction angle of the peak respectively

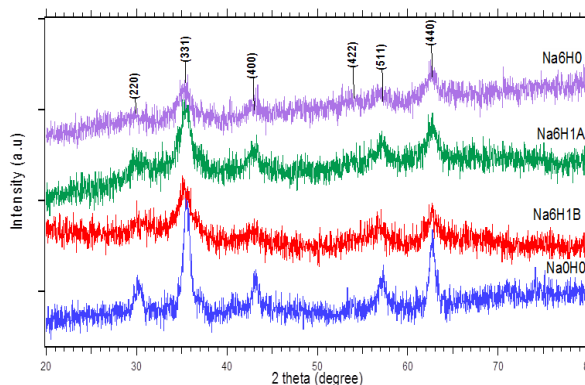


Figure 1. X-ray diffraction pattern of (from the bottom to the top): NaOH0, Na6H1B, Na6H1A, Na6H0 samples

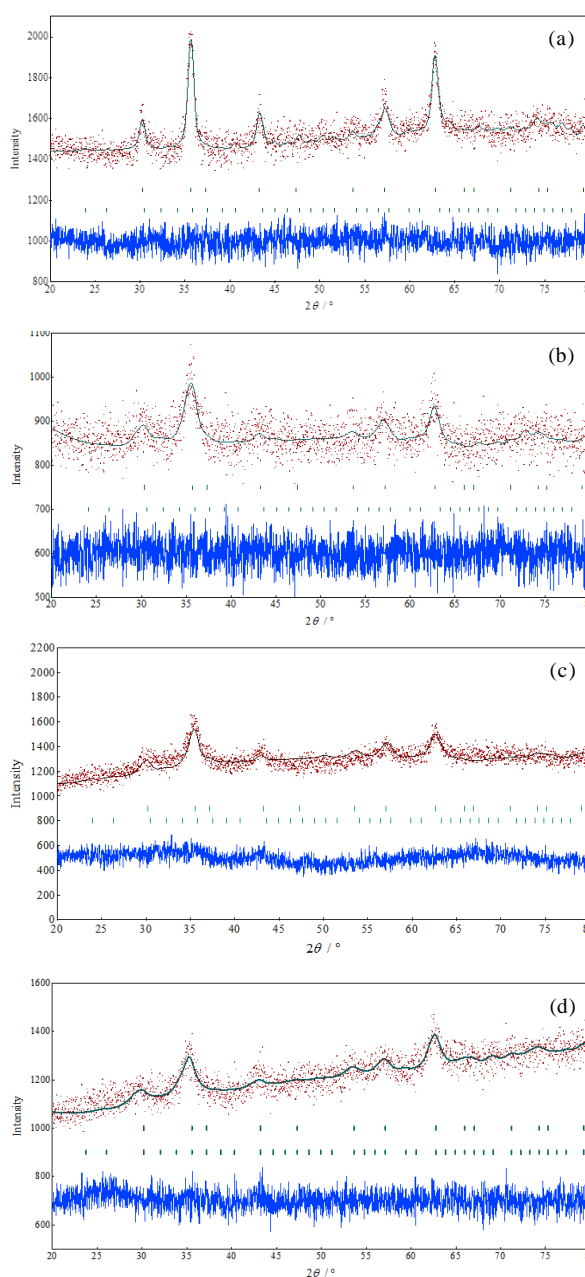


Figure 2. Rietan profile for NaOH0, Na6H1B, Na6H1A and Na6H0 samples (from top to the bottom).

**Table 2.** Characteristic of Magnetic Nanoparticles Fe<sub>3</sub>O<sub>4</sub>/γ-Fe<sub>2</sub>O<sub>3</sub> after some modification with NaNO<sub>3</sub> and HNO<sub>3</sub>

No.	Sample Code	Phase fraction of Fe <sub>3</sub> O <sub>4</sub> (%)	Ms (emu/gram)	Particle Size (nm)			
				PSA	TEM	XRD	VSM
<i>1<sup>st</sup>-series</i>							
1.	Na0H0	0.2445	77.97	268.08	> 10	11.53683	8.43555
2.	Na0H0.2	0.2693	65.58	127.58		12.73387	9.09387
3.	Na0H0.4	0.6154	49.61	84.78		13.07568	10.2102
4.	Na0H0.6	0.566	46.33	85.58		10.86988	10.282
<i>2<sup>nd</sup>-series</i>							
1.	Na2H1B	0.6909	41.89	62.94		6.391233	8.80253
2.	Na4H1B	0.5711	38.04	46.04		5.064378	8.90071
3.	Na6H1B	0.4751	37.25	31.60		6.767192	8.4052
4.	Na8H1B	0.4990	40.76	118.3		5.701094	8.0476
5.	Na10H1B	0.4136	32.00	49.38		4.902013	8.54175
<i>3<sup>rd</sup>-series</i>							
1.	Na2H1A	0.5004	41.44	60.85		6.185035	8.84123
2.	Na4H1A	0.3906	36.97	71.08		3.864834	9.09461
3.	Na6H1A	0.4355	43.97	20.02		8.225166	8.91152
4.	Na8H1A	0.4639	36.49	181.20		4.487137	8.94456
5.	Na10H1A	0.4865	35.10	109.10		7.372910	8.56649
<i>4<sup>th</sup>-series</i>							
1.	Na6H0	0.5172	38.50	65.10	< 10	5.129831	8.38461
2.	Na6H0.2	0.4378	41.02	25.75		7.160345	8.87663
3.	Na6H0.4	0.5415	30.36	27.30		7.811543	9.35783
4.	Na6H0.6	0.4679	34.01	27.10		7.732329	8.953

Complete phase identification and crystallite size calculation result of all the samples are presented at Table 2.

Data on Table 2 show that HNO<sub>3</sub> peptizing have the role on controlling the growth of Fe<sub>3</sub>O<sub>4</sub> phase in which higher HNO<sub>3</sub> molarity will result in higher Fe<sub>3</sub>O<sub>4</sub> weight fraction. The crystallite size tends to be unaffected by HNO<sub>3</sub> modification. NaNO<sub>3</sub> addition give more pronounce effect on nanoparticle crystallite size. The stage of NaNO<sub>3</sub> addition is also affecting the growth of magnetic nanoparticle crystalline. The crystallite size of the 2<sup>nd</sup> series are more stable related to NaNO<sub>3</sub> molarity changing compared to the 3<sup>rd</sup> series samples. However, crystallite sizes of the 3<sup>rd</sup> series samples are only randomly related to the NaNO<sub>3</sub> molarity and magnetic nanoparticle of as low as 3 to 4 nm crystallite size could be achieved.

### TEM Observation

Figure 3 displayed the TEM photograph of typical morphology of magnetic nanoparticle without and after having some modification with HNO<sub>3</sub> and NaNO<sub>3</sub>. Generally, the photograph clearly displayed spherical nanoparticle with different pattern of agglomeration. For samples without modification, Na0H0 at Figure 3(a), magnetic nanoparticle tend to agglomerate with the size of more than 10 nm.

Peptizing with HNO<sub>3</sub> made the magnetic nanoparticle are more separated to each other, leaving

only a slight agglomeration, but with the nanoparticle size of still around 10 nm. Nanoparticle size of NaNO<sub>3</sub> modified samples are more uniform in the range smaller than 10 nm but with no significant difference between 2<sup>nd</sup> series and 3<sup>rd</sup> series samples. It could be assumed that HNO<sub>3</sub> have a role on dispersing the magnetic nanoparticle within the medium and will be responsible for stabilizing the magnetic nanoparticle colloid. These TEM data were also picturing the role of NaNO<sub>3</sub> on reducing magnetic nanoparticles size, which fit with magnetic nanoparticle crystallite size behaviour analyzed from XRD pattern.

### PSA Data

Agglomeration tendency of any colloidal samples could be analyzed using dynamic laser scattering (DLS) technique. This technique usually implemented on Particle Size Analyzer (PSA) instrument which measures the “real” particle size within the colloid samples and usually describe as distribution curve of groups of magnetic nanoparticle with specified size range. Such distribution curve of magnetic nanoparticle synthesis in this experiment are shown in Figure 4.

The curve show the maximum particle distribution at the size of more than 260 nm for the sample without any modification (Figure 4(a)) and at the size of around 20 nm for after NaNO<sub>3</sub> modification. Complete data of particle size measured by PSA are listed at Table 2.

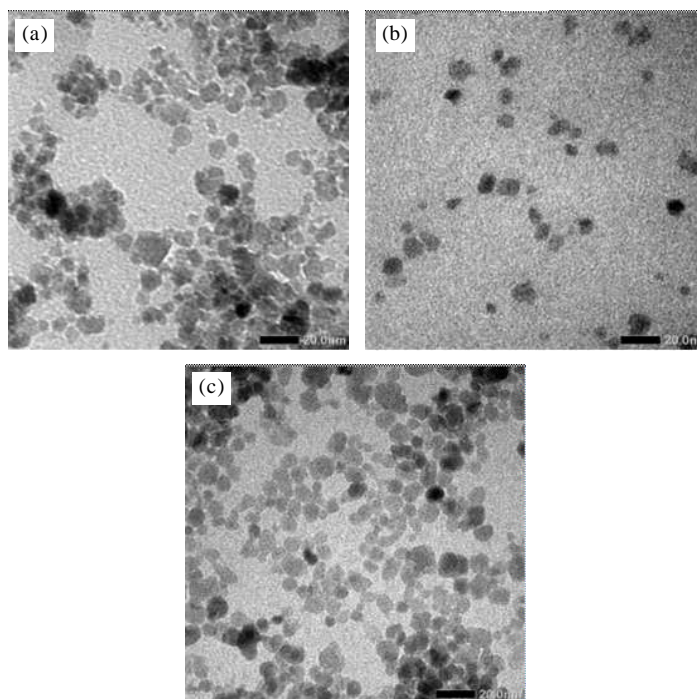


Figure 3. TEM photograph of (a). NaOH0, (b). NaOH1 and (c). Na6H1B samples (the scale bar is equal to 20 nm).

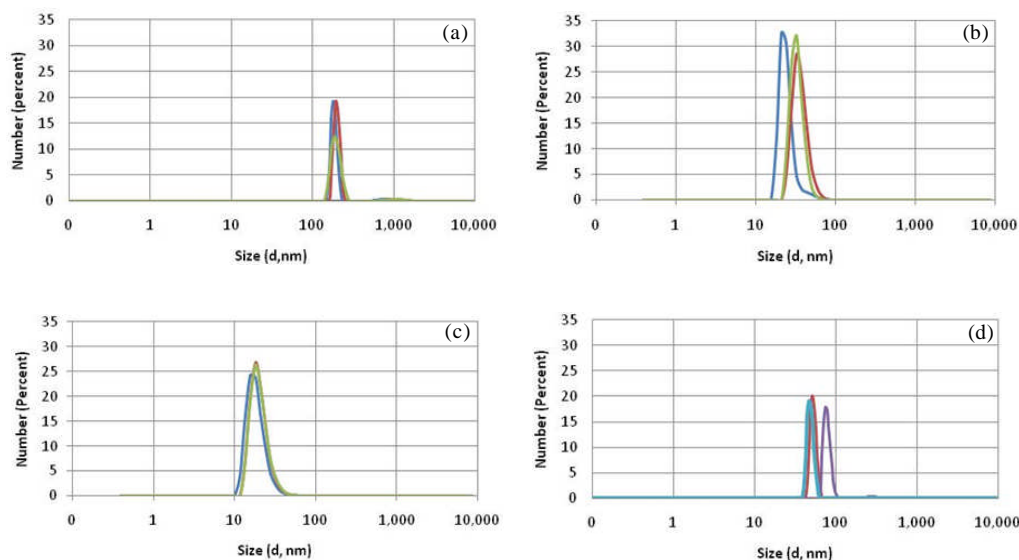


Figure 4. Size distribution curve of (a). NaOH0; (b). Na6H1B; (c). Na6H1A and (d). Na6H0 samples.

Referring to the single particle size measured by TEM, these PSA data could be considered as the size of a group of particles which forming agglomeration within the colloid. A significant decrease of the size with increasing  $\text{HNO}_3$  molarity for 1<sup>st</sup> series samples indicates that  $\text{HNO}_3$  will modify nanoparticles surface and inhibited agglomeration possibility between the particles. At the contrary,  $\text{NaNO}_3$  modification will give contra-productive result on nanoparticle agglomeration tendency. There is an optimum  $\text{NaNO}_3$  molarity that give smallest agglomerate in the size of around 20 nm for samples modify with 6M  $\text{NaNO}_3$ . PSA data of the

4<sup>th</sup>-samples revealed the same size which affected no more with increasing of  $\text{HNO}_3$  molarity. For higher  $\text{HNO}_3$  molarity, the agglomerates tend to increase again to hundreds size.

### Magnetic Hysteresis Curve Analysis

Magnetic properties of nanoparticles system usually analysed from its magnetic hysteresis curve. The curve is generated by measuring the magnetic flux or magnetization (M) of a ferromagnetic material while the magnetizing force (H) is changed. Figure 5

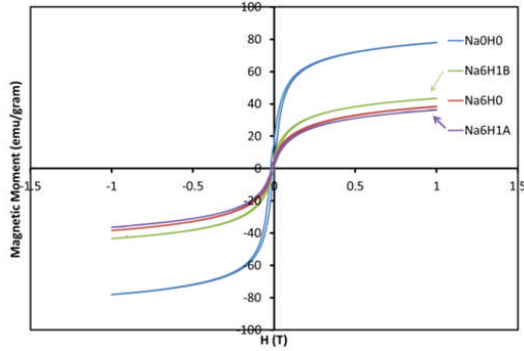


Figure 5. Magnetic hysteresis curve of NaOH0, Na6H1B, Na6H1A and Na6H0 samples.

showed the curve for some samples with different synthesis parameters, which show typical curve for superparamagnetic materials. For sample without any modification, NaOH0, magnetization was saturated at  $M_s$  value of 78.14 emu/gram and decreased to a range of 30 to 40 emu/gram after HNO<sub>3</sub> and NaNO<sub>3</sub> modification with HNO<sub>3</sub> modification being more responsible than NaNO<sub>3</sub>.

For superparamagnetic systems, magnetization values as a function of the external field,  $M(H)$  at  $T > T_B$  could be modeled using Langevin function approach,  $L(\alpha)$  and Equation [8]:

$$M(H) = M_s L(\alpha) \quad \dots\dots\dots (2)$$

In which  $M_s$  is spontaneous magnetization,  $k_B$  is Boltzmann constant equal to  $1.38064852 \times 10^{-23}$  J/K,  $m$  is magnetic moment for 1 particle unit

$$L(\alpha) = \coth \alpha - \frac{1}{\alpha} \quad \dots\dots\dots (3)$$

$$\alpha = \frac{mH}{k_B T} = aH \quad \dots\dots\dots (4)$$

For spherical magnetic nanoparticle,

$$m = M_s w = M_s \rho V = \frac{\pi}{6} d^3 M_s \rho \quad \dots\dots\dots (5)$$

with  $\rho$  being density of mass of Fe<sub>3</sub>O<sub>4</sub>. From Equation (4) and Equation (5),

$$d = \sqrt[3]{\frac{6k_B T}{\pi \rho} \left[ \frac{\alpha}{M_s} \right]} \quad \dots\dots\dots (6)$$

Typical fitting result of the samples are presented at Figure 6. From this fitting, it was found that the  $d$ -value of magnetic cores for samples without NaNO<sub>3</sub> modifications were around 10 nm, while for NaNO<sub>3</sub>-modified samples were 8 to 9 nm. Complete magnetic core value of all the sample are listed at Table 2.

### Controlled Growth Mechanism of Magnetic Nanoparticles

Data on Table 2 shows some characteristic behaviour of magnetic nanoparticles related to the

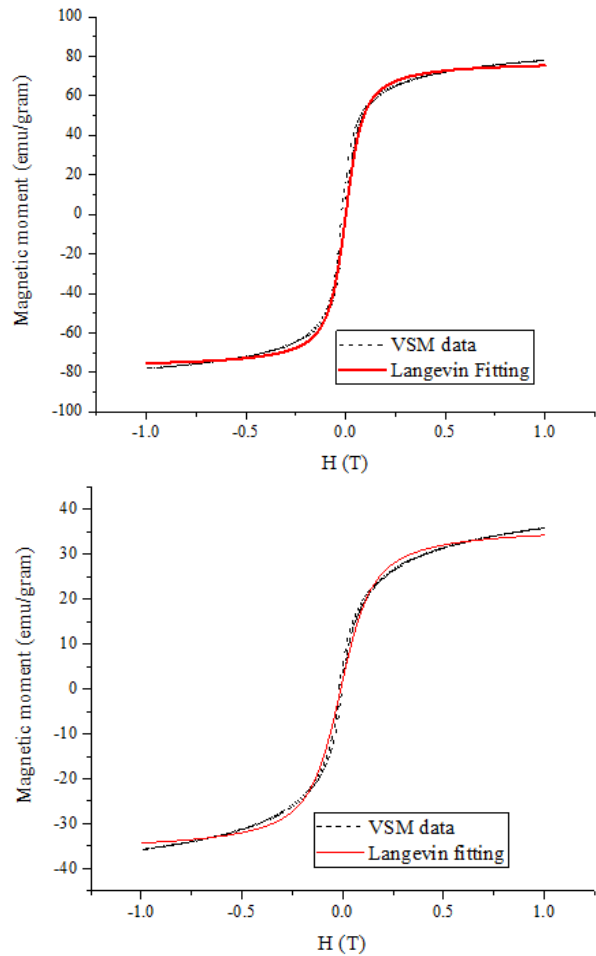
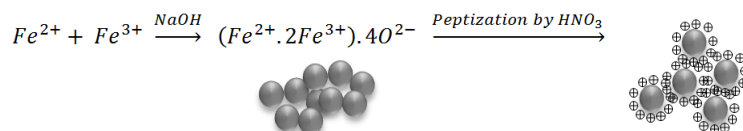


Figure 6. Langevin Fitting of NaOH0 and Na6H1B samples.

modification with HNO<sub>3</sub> and NaNO<sub>3</sub>. It can be resumed that three characteristics including magnetization, crystallization and particle size were all depend on some sort to the modification. HNO<sub>3</sub> modification shows more pronounce effect on magnetization and agglomeration, while addition of NaNO<sub>3</sub> affected more on magnetic nanoparticle crystallinity which also depend on the stage of NaNO<sub>3</sub> addition.

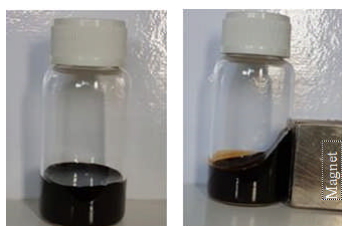
Principally, co-precipitation process on magnetic nanoparticles synthesis usually involves several steps including nucleation of precipitate, growth of precipitate and completion of magnetic nanoparticles phases [4]. At nucleation stage, Fe<sup>2+</sup> and Fe<sup>3+</sup> ions will react with O<sup>2-</sup> from NaOH base immediately after mixing which represent as the direct change of solution colour from yellowish to black solution and forming magnetic nanoparticles nuclei. These nuclei then growth with more diffusion of the ions afterward to the nuclei and arrange them to build crystal structure and phases favour for the system, which take time during overnight aging. At the end of the reaction, system will have a high alkaline environment at high pH ~ 12 and chlorine residue within medium.



**Gambar 7.** Schematic illustration of precipitation and peptization on iron oxide magnetic nanoparticles.

Without further peptization, the precipitate, earned after several washing steps until neutral pH, will be agglomerated due to strong magnetic interaction between nanoparticles and will result in un-stable magnetic liquid. Massart's method solve this problem by peptization of nanoparticles surfaces with acidic or alkaline surfactant such as  $HClO_3$  and TMAOH respectively [5]. The magnetic sol formed after peptization will be easily dispersed in water and give stable colloid system. Such processing step is schematically present at Figure 7.

From this schema, it could be figure out the function of  $HNO_3$  on stabilizing magnetic nanoparticle within water medium. magnetic nanoparticle will be separated by positive polar repulsion and bound to medium via hydroxyl group [9]. As a result, the magnetic



**Figure 8.** Photo of liquid magnetic sample before (left) and after (right) attracted by permanent magnet.

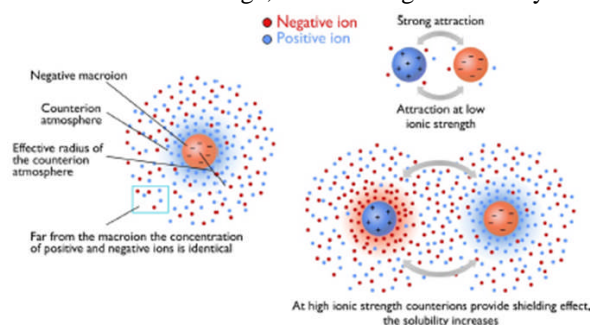
nanoparticle will be attracted by magnet as a magnetic liquid, instead of as separate magnetic particle which can be seen in Figure 8.

For samples with low molarity of  $HNO_3$ , the magnetic interaction between the nanoparticles remained strong and led to the agglomeration. Increasing  $HNO_3$  molarity, will give thicker layer of surfactant. As the results, the particles are more separated, agglomerate size will be decreased and more stable colloid will be obtained. However magnetic interaction between nanoparticles will also be weakened and there is a dilution effect to the magnetic fraction which consequently decreasing the magnetization value. The size of crystalline show no significant relation to  $HNO_3$  addition, which confirm that the growth of nanoparticle size are already completed within overnight-aging time or before peptization [10].

On the other hand, addition of  $NaNO_3$  will alter ionic strength on the reaction system. Illustration of counter ion role on giving shielding effect to the interaction between positive and negative ions is presented at Figure 9 [11].

$NaNO_3$  is a kind of inert activity salt and can be used to make an increase of ionic strength of solution,

which will slow the growth and nucleation rate at the same time based on the decrease of the activity of precursor ions. Addition of  $NaNO_3$  will give shielding effect to positive Fe ions and decreasing the possibility to be reacted and precipitated by negative  $OH^-$  ions. When  $NaNO_3$  added before precipitation or being mixed with Fe-salt at first stage, the shielding was already take



**Figure 9.** Schematic illustration of counter ion role on altering ionic strength between interacting macroion [11].

place even before nucleation which result in amorphous nature of nanoparticles nuclei.

On the other hand, when  $NaNO_3$  added after precipitation,  $NaNO_3$  shielding around iron oxide nuclei will hindered diffusion of the ions afterward to the nuclei and inhibited further growth of nuclei. These analyses could explain the more stable particles size for 2<sup>nd</sup> series comparing to 3<sup>rd</sup> series. After all,  $NaNO_3$  addition will slow down and inhibited the growth of nanoparticles to certain extent which depend on  $NaNO_3$  molarity. Decreasing of  $M_s$  on  $NaNO_3$  addition but rather insensitive to its molarity was also found as a result. Regarding all the parameters used within the synthesis and their characteristic, it can be proposed that modification with  $NaNO_3$  of 6M molarity will give an optimum characteristic.

## CONCLUSION

Size controlled magnetic nanoparticle (MNPs) of iron oxide have been prepared in the presence of  $NaNO_3$  via co-precipitation method followed by peptizing with  $HNO_3$ . The magnetic nanoparticle properties including phase, crystallite size and magnetic properties were optimized by addition of  $HNO_3$  and  $NaNO_3$  in varied molarity and at different stage of process. As an end product, stable water-base colloids were formed, containing magnetic nanoparticle of  $Fe_3O_4/\gamma-Fe_2O_3$  with nanoscale crystallite size. From data analyses result, it can be concluded that  $NaNO_3$  addition will significantly

affect crystallite formation due to its capability to increase ionic strength within the medium which slowed down the growth of magnetic nanoparticles and decreasing crystallite size. However magnetic hysteresis curve, measured using VSM, still displayed a typical superparamagnetic behaviour of magnetic nanoparticles with high enough magnetization value,  $M_s$ . At average, uniform magnetic nanoparticle with nanoparticles size of smaller than 10 nm, crystallite size and magnetic domain smaller than 8 nm and  $M_s$  value of 40 emu/gram is obtained for samples modified by NaNO<sub>3</sub> of 6M molarity and HNO<sub>3</sub> of 1M molarity.

## ACKNOWLEDGEMENT

This work was supported by DIPA of PSTBM at fiscal year of 2016/2017. The authors would like to thank PSTBM-BATAN management for providing this fund. Acknowledgement would also be delivered to Drs. Sudirman, M.Sc for valuable discussion and technical support during the experiment.

## REFERENCES

- [1]. J. D. Robertson, L. Rizello, M.A. Olias, J. Gaitzsch, C. Contini, M.S Magon, S.A. Renshaw and G Battaglia. "Purification of Nanoparticles by Size and Shape." *Sci. Rep.*, vol. 6, pp. 27494, 2016.
- [2]. A. Yadollahpour. "Magnetic Nanoparticles in Medicine: A Review Of Synthesis Methods and Important Characteristics." *Orient. J. Chem.*, vol. 31, 2015.
- [3]. W. Wu, Z. Wu, T. Yu, C. Jiang, and W.-S. Kim. "Recent Progress on Magnetic Iron Oxide Nanoparticles: Synthesis, Surface Functional Strategies And Biomedical Applications." *Sci. Technol. Adv. Mater.*, vol. 16, no. 2, pp. 23501, 2015.
- [4]. N. T. K. Thanh, N. Maclean, and S. Mahiddine. "Mechanisms of Nucleation and Growth of Nanoparticles in Solution." *Chem. Rev.*, vol. 114, no. 15, pp. 7610-7630, 2014.
- [5]. R. Massart, J. Roger, and V. Cavuil. "New Trends in Chemistry of Magnetic Colloids: Polar and Non Polar Magnetic Fluids, Emulsions, Capsules and Vesicles." *Brazilian J. Phys.*, vol. 25, no. 2, pp. 135, 1995.
- [6]. L. Vayssières, C. Chanéac, E. Tronc, and J. P. Jolivet. "Size Tailoring of Magnetite Particles Formed by Aqueous Precipitation: An Example of Thermodynamic Stability Of Nanometric Oxide Particles." *J. Colloid Interface Sci.*, vol. 205, no. 2, pp. 205-212, 1998.
- [7]. F. Izumi and T. Ikeda. "A Rietveld-Analysis Programm RIETAN-98 and its Applications to Zeolites." *Mater. Sci. Forum*, vol. 321-324, pp. 198-205, 2000.
- [8]. P. Tuček, M. Tučková, E. Fišerová, J. Tuček, and L. Kubáček, "Design of Experiment for Measurement of Langevin Function." *Meas. Sci. Rev.*, vol. 12, no. 4, pp. 121-127, 2012.
- [9]. A. K. Hauser, R. Mathias, K. W. Anderson, and J. Z. Hilt. "The Effects of Synthesis Method on The Physical and Chemical Properties of Dextran Coated Iron Oxide Nanoparticles." *Mater Chem Phys*, vol. 160, pp. 177-186, 2015.
- [10]. J. Baumgartner, A. Dey, P. H. Bomans. C. L. Coadou, N.A. Sommerdijk, D. Faivre. "Nucleation and Growth of Magnetite from Solution." *Nat. Mater.*, vol. 12, no. 4, pp. 310-4, 2013.
- [11]. G. Hegyi, J. Kardos, M. Kovacs, A.M Csizmadia, L. Nyitray, G. Pal, L. Radnai, A. Remenyi, I. Venekei. *Introduction to Practical Biochemistry*. Eötvös Loránd University, 2013.