Study on Distribution of Magnetite (Fe$_{3-x}$Mn$_x$O$_4$) Filler in Fe$_{3-x}$Mn$_x$O$_4$-PEG/PVA/PVP Magnetic Hydrogel by Using Two-lognormal Function Analysis

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Abstract. The trend of nanoparticles technology has been so advanced in several years; for instance, the bio-applications of polymer gel composites that involve micro/nanoparticles to be embedded in polymer gel have been many observed. This complex material composition can express a specific property and low toxicity so it can be used in various domain uses. Besides, the polymers material must possess certain properties to be utilized in the biomedical application (magnetic hyperthermia, cancer therapy, and food industry) such as flexible, biocompatibility, and water swallowability. Magnetite is nanoparticle with a unique property that can be utilized as a filler. The low toxicity of magnetite under the superparamagnetic condition is very useful in biomedical application. This research was succeeded to synthesize PVA/PVP polymer-based hydrogel magnetic with PEG-coated magnetite with Manganese doping filler, which has good biocompatibility and stable particle. An advanced characterization using XRD has shown the crystallite size about 9-11 nm with the magnetite phase confirmed from the sample. The SAXS analysis using two-lognormal functions exhibited primary and secondary particles around 2.40 and 9.74 nm that was well proven by TEM image analysis that showed a close value of the average particles size about 9.78 nm. From the SAXS and TEM analysis, it could be observed that the samples formed clusters from primary and secondary particles.

Keywords: Fe$_{3-x}$Mn$_x$O$_4$-PEG, PVA/PVP, magnetic hydrogel, SAXS analysis, two lognormal function

1. Introduction

Nanoparticles are increasingly becoming the focus of research highly developed by researchers due to their superior nature. Many kinds of nanoparticles being developed by researchers ranging from silica [1–5], cobalt [6–10], ZnO [11–13], and so forth. One of the nanoparticles that have low toxicity [14–16] and biocompatible [17–19] is magnetite so that it can be used in the biomedical field [20–23]. Practically, magnetite can be utilized as a filler material in the magnetic hydrogel to be subsequently used as actuator [24,25] or as artificial [26–28]. Besides, the magnetic hydrogel can be also used as the packaging for environmentally friendly food.
The need for food in the current era is increasing with the increase in the world population [29], where it is in line with the enhancement of the need for high-quality food. Besides, the urgency of the waste produced by the food wrapping is also massively discussed by many scientists [30]. Considering such problems, an environmentally friendly food packaging was developed by the ability to protect the food form environmental contamination that can give the information of the food condition in the real-time and the packaging waste produced is biodegradable [31–33]. One of the developments of this food packaging is the use of hydrogel polymer as the raw material. PVP and PVA polymers have been proven to have a biocompatible characteristic with the low toxicity level and it is biodegradable [34–40]. Meanwhile, the use of PVA and PVP simultaneously becomes the magnetic hydrogel is still rarely reported.

Magnetic hydrogel consists of two important parts namely filler and hydrogel polymer as the place in which the filler materials attach. The filler of hydrogel polymer can be originated from any nanoparticles that one of them is in the form of magnetite nanoparticles [41]. The previous research on the hydrogel magnetic that used magnetic nanoparticles was reported by Safronov et al. [42]. They explained that magnetic nanoparticles and PAAm polymer underwent a strong interaction when they were composed so that the agglomeration appeared. This case is different from the condition when the filler magnetic used came from magnetic nanoparticles that had less agglomeration. This research caused the hydrogel magnetic difficult to develop if it was subjected to an external magnetic field but having the higher shear modulus. Besides, Blyakhman et al. [43] reported that the addition of magnetite in the PAAm hydrogel could increase the mechanic strength of the hydrogel magnetic system even in proportion to the increase in the polymer bonds. This addition also influenced the Young modulus that was in line with the number of magnetites. The magnetization measurement also showed the magnetite percentage which was in line with the magnetization happening in the magnetic hydrogel. The magnetic fabrication has been also successfully conducted by Sunaryono et al. [44] that used the raw material of polyvinyl alcohol (PVA) and magnetite filler. The formed magnetite nanoparticles had the diameter between 13 – 32 nm because of the aggregation between the particles that also correlated to the magnetite filler concentration level in the polymer namely from 1% to 15% of the hydrogel weight. However, the study of the distribution of magnetite filler (Fe$_{3-x}$Mn$_x$O$_4$) in the magnetic hydrogel of Fe$_{3-x}$Mn$_x$O$_4$-PEG/PVA/PVP is rarely reported.

This research had the magnetite raw material as the filler due to its superior nature and superparamagnetic characteristic when its size under 10 nm [45–47]. The synthesis of magnetite particles in this research employed the method of simple degradation namely coprecipitation method. Besides, the use of surfactant could improve the monodispersity of the magnetite nanoparticles [48]. Thereby, this research fabricated the magnetite added with PEG polymer and dopped with Mn. This polymer has the ability to add the biocompatibility of the magnetite [49,50]. Meanwhile, Mn doping was carried out to increase the magnetic characteristic of the magnetite nanoparticles. Subsequently, the study of the distribution of magnetite nanoparticles (Fe$_{3-x}$Mn$_x$O$_4$) in the magnetic hydrogel of Fe$_{3-x}$Mn$_x$O$_4$-PEG/PVA/PVP was undertaken using SAXS instrument and analyzed using global fitting method through a two-lognormal model.

2. Methods

2.1. Synthesis of Fe$_{3-x}$Mn$_x$O$_4$-PEG-coated nanoparticles

Natural iron sand that was used as the source of magnetite was taken from Tulungagung District, Indonesia. Before synthesizing the magnetite nanoparticles, we first mixed the PEG and distilled water to make the surfactant. Magnetite powder as starting material was extracted from the iron sand using a permanent magnet. The certain amount of magnetite powder was then synthesized to nanoparticles through coprecipitation method because of its simplicity. The magnetite powder was initially dissolved in hydrochloric acid (HCl) then mixed with MnCl$_2$ and PEG$_2$H$_2$O using magnetic stirrer followed by the titration process using ammonium hydroxide (NH$_4$OH) to obtain the magnetite nanoparticles as precipitates. The precipitation yielded was then washed using distilled water until pH = 7 was reached.
The variables through the process were under controlled (temperature, reaction time, pH, and mixing speed).

2.2. Fabrication of Fe\textsubscript{3.3}Mn\textsubscript{0.7}O\textsubscript{4}-PEG/PVA/PVP magnetic hydrogels

PVA/PVP hydrogels were prepared by dissolving a particular composition of the polymers powder to distilled water under the controlled temperature to increase the solubility of the hydrogels. The process was succeeded by the indicator of polymers form changing into a paste. The paste was then mixed again with the magnetite nanoparticles and stirred until became homogenous gels. This experiment used the composition of polymer PVA/PVP and distilled water about 40:100 with 15% of Fe\textsubscript{3.3}Mn\textsubscript{0.7}O\textsubscript{4}-PEG filler. The yield of the mixture was then placed in a mold prepared before and continued to F-T process for 3h compared to 1.5h.

2.3. Characterization

To collect the phase and crystal structure data from the powder samples of Fe\textsubscript{3.3}Mn\textsubscript{0.7}O\textsubscript{4}-PEG nanoparticles, a characterization using X-ray diffractometer (Philips X-Pert MD diffractometer) was conducted under 0.02° scanning data step and in the range of CuKα (λ = 1.54) angular position (2θ) from 20° to 90°, and was done at room temperature. The morphology of Fe\textsubscript{3.3}Mn\textsubscript{0.7}O\textsubscript{4}-PEG nanoparticles samples was characterized using a Transmission Electron Microscope (JEM1400 JEOL). In addition, a comparison between the size data results from XRD and SAXS analyses can be made for further investigation.

The characterization using synchrotron SAXS was performed at Siam Photon Laboratory of Synchrotron Light Research Institute (SLRI) of Thailand to obtain the Fe\textsubscript{3.3}Mn\textsubscript{0.7}O\textsubscript{4}-PEG nanoparticles distribution as the filler inside the Fe\textsubscript{3.3}Mn\textsubscript{0.7}O\textsubscript{4}-PEG/PVA/PVP Magnetic Hydrogels. The characterization was also conducted because of the non-destructive measurement and the ability to obtain the primary size of the magnetic hydrogel sample. The energy of X-ray on the range of 6 to 9 keV generated the intensity of the SAXS scattering X-ray showed as the scattering vector function, q, that can cover the momentum transfer from 0.12 nm\textsuperscript{-1} to 2 nm\textsuperscript{-1} then can be noted as \( q = (4π/λ) \sin (θ/2) \), where θ is the angle between the scattered and incident beam, and λ refers to the wavelength of the incident X-ray. The sample-to-detector distances (dd) of CDD detector used in the SAXS measurement for the high and low q ranges were 1200 mm and 4500 mm respectively. The data collected from the SAXS characterization were then normalized using SAXSIT and also calibrated and made the correction between the sample and the background scattering intensities. The final result on using SAXSIT software was the merged data of two different scattering vector ranges. The next step in the SAXS analysis was performed using SASfit software (0.94.10 version); under the two-lognormal distribution function model, the primary and secondary particles size, as well as the fractal dimension and the particles aggregation before dispersing process into the hydrogel could be obtained.

3. Results and Discussion

The X-ray diffraction (XRD) patterns of all powder samples of Fe\textsubscript{3.3}Mn\textsubscript{0.7}O\textsubscript{4}-PEG with various concentrations of dopant in the range of x = 0, 0.2, 0.4, and 0.6 are shown in Figure 1. The patterns produced indicated that the powder samples had a quite similar patterns data with pure magnetite from other reported file before which could be confirmed from the suitable peaks of standard magnetite (JCPDS file, PDF No.19-0629) [51]. The XRD patterns showed some peaks corresponding to the (2 2 0), (3 1 1), (4 0 0), (4 2 2), (5 1 1), (4 4 0), and (5 3 3) planes which became the indication of spinel cubic structure with space group \( Fd\textit{3}m \) [44]. Besides that, a shifting happened to the diffraction peaks to the lower angle as the increasing of x presented by Figure 2, inferred that the increasing of lattice parameter because of the successful insertion of Mn\textsuperscript{2+} ion by replacing Fe\textsuperscript{2+} and Fe\textsuperscript{3+} ions were partially from Fe\textsubscript{3.3}Mn\textsubscript{0.7}O\textsubscript{4}-PEG [52,53]. Another reason from the increase happening was the ionic radius of Mn\textsuperscript{2+} that was larger than Fe\textsuperscript{2+} and Fe\textsuperscript{3+} ionic radius [54]. The lattice parameters of powder sample Fe\textsubscript{3.3}Mn\textsubscript{0.7}O\textsubscript{4}-PEG correspondingly increased from 8.3711 to 8.3939 Å due to the increase of x appropriate with another
report before [55]. The analysis of XRD using Rietica software [56] exhibited an average crystallite size about 9-11 nm.

Figure 1. XRD patterns of Fe$_{3-x}$Mn$_x$O$_4$-PEG sample with various concentrations of dopant in the range of $x = 0, 0.2, 0.4,$ and 0.6

Figure 2. Profile of (311) peak shift of the powders sample of Fe$_{3-x}$Mn$_x$O$_4$-PEG sample with various concentrations of dopant in the range of $x = 0, 0.2, 0.4,$ and 0.6
The complex system of magnetic hydrogel that consists of \( \text{Fe}_{3-x}\text{Mn}_x\text{O}_4\)-PEG nanoparticles as filler and polymer hydrogel PVA/PVP was characterized using SAXS and to investigate the aggregation of the nanoparticle \( \text{Fe}_{3-x}\text{Mn}_x\text{O}_4\)-PEG inside the magnetic hydrogel that included primary and secondary particles, the synchrotron data were analyzed using two-lognormal distribution function model to get the form factor of these particles. Besides that, two-lognormal distribution model also offers good information about the distribution of primary and secondary particles [44]. The function of the scattering vector \( q \) was used to calculate the scattering intensity in SAXS analysis with a certain relation between the scattering intensity \( I(q) \), the form factor \( P(q) \), and the structure factor \( S(q) \) for particular \( N \) of \( \text{Fe}_{3-x}\text{Mn}_x\text{O}_4\)-PEG nanoparticles per unit volume that can be noted as follow.

\[
I(q) = NpP(q)S(q) + bkg \tag{1}
\]

The morphology of the samples yielded from the characterization using TEM in this experiment showed that the nanoparticles formed fractal aggregates between one another that involved polydisperse nanoparticles (in cluster size) in which the cluster size distribution would have a limitation because of the process of aggregation. Moreover, the polydispersity will affect the difference of shape between the observed and the single cluster structure factor. The aggregate of the clustered object model which has a spherical particle formed a fractal-like cluster with \( \xi \) (correlation length) conforming to their whole size and \( D \) as the fractal dimension (self-similarity dimension) [57]. The structure factor of this system can be expressed by [58]

\[
S(q, \xi, D, r) = 1 + \frac{D_1(D-1)\sin([D-1]\tan^{-1}(q\xi))}{(q\xi)^D[1+(q\xi)^2]^{(D-1)/2}} \tag{2}
\]

The \( \text{Fe}_{3-x}\text{Mn}_x\text{O}_4\)-PEG/PVA/PVP magnetic hydrogel SAXS data were well-refined by SASfit software using two-lognormal distribution function shown in a good curve fitting in Figure 3 in the assumption that the system consisted of not only single size distribution. The two-lognormal equation is given in previous work [59]. The two-lognormal analysis result showed that the primary particles distribution of \( \text{Fe}_{3-x}\text{Mn}_x\text{O}_4\)-PEG dispersing inside the PVA/PVP hydrogel had the diameter of \( \sim 2.4 \) nm which had the similar result of the previous experimental report by Sun et al. [60] and Baumgartner et al. [61] that got the cluster aggregates with the diameter \( \sim 2.6 \) nm and \( 1-4 \) nm respectively.

In addition to the presence of primary particles, also there were secondary particles with the bigger size distribution about \( 9.74 \) nm; the diameter of these secondary particles was confirmed by the characterization using the image of transmission electron microscopy (TEM) which had been analyzed using ImageJ and Origin software; the analysis exhibited an average particle of \( 9.78 \) nm that had the similarity with the XRD crystallite size of \( \text{Fe}_{3-x}\text{Mn}_x\text{O}_4\)-PEG in this experiment \((\sim 9-11) \) nm and from another experiment reported before [62,63]. The evidence of the aggregation process in magnetite nanoparticles is shown in Figure 4; it shows that the nanoparticles tended to cluster because of the strong pull force between one particle and another [64]. Another variable from equation (2) and (3) that contributed to the sample was \( D \) which referred to the fractal dimension of \( \text{Fe}_{3-x}\text{Mn}_x\text{O}_4\)-PEG, in the analysis the fractal dimension tends to 3 \((\sim 2.78 \) exactly), corresponding to the sample the structure that grew into three-dimensional building block [52]. The aggregation of \( \text{Fe}_{3-x}\text{Mn}_x\text{O}_4\)-PEG can be illustrated as Figure 5.
Figure 3. SAXS patterns of $\text{Fe}_{3-x}\text{Mn}_x\text{O}_4$-PEG nanoparticles inside magnetic hydrogel with the F-T process of 3 time

Figure 4. The TEM image of magnetite with particle size distribution
Figure 5. Schematic of Fe$_{3-x}$Mn$_x$O$_4$-PEG nanoparticles aggregation inside PVA/PVP hydrogel magnetic

4. Conclusion
The synthesis of Fe$_{3-x}$Mn$_x$O$_4$-PEG with $x = 0$, 0.2, 0.4, and 0.6 nanoparticles has been successfully conducted through coprecipitation method. The magnetite phase was well identified from Fe$_{3-x}$Mn$_x$O$_4$-PEG nanoparticles according to JCPDS file, PDF No.19-0629 with XRD patterns analysis result that showed the crystallite size about 9-11 nm. Fe$_{3-x}$Mn$_x$O$_4$-PEG/PVA/PVP magnetic hydrogel has been characterized by SAXS and analyzed using a two-lognormal function that produced the primary and secondary particle size ~2.4 nm and ~9.7 nm respectively; this result was well confirmed by the TEM image that exhibited the nanoparticles size about 9.8 nm polymeric.

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