Synthesis and Characterization of ZnO-decorated GO Nanocomposite Material with Different ZnO Loading through Sol-gel Method

(Penghasilan dan Perincian Komposit Nano GO-ZnO dengan Muatan ZnO yang Berbeza melalui Kaedah Sol-gel)

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ABSTRACT

This work aimed is to synthesis a well dispersed zinc oxide (ZnO) nanoparticles (NPs) decorated on graphene oxide (GO) nanosheet with a practical way by using sol-gel technique. Zinc acetate dehydrate (Zn(CH$_3$COO)$_2$·2H$_2$O) was used as precursor of ZnO and absolute ethanol as solvent. 1 weight percent (wt%), 5 wt%, 10 wt%, and 20 wt% of ZnO was decorated on GO nanosheet. A series of analysis was carried out to characterize the synthesized ZnO-decorated GO nanocomposite material. The results of XRD analysis show some long area of peak at 25° to 80° allocate for ZnO in the ZnO-decorated GO nanocomposite material. By performing zeta potential analysis, the findings show that there was increment of negative surface charge on ZnO-decorated GO nanocomposite material. The experiment result also found that the hydrodynamic particle size of ZnO-decorated GO nanocomposite material become larger when high ZnO loaded. FESEM micrographs demonstrated that spherical-shaped of ZnO NPs appeared on the GO nanosheet with further proved by EDX where the content of ZnO-decorated GO nanocomposite material was composed by 71.3 wt% of C, 17 wt% of O, and additional element of 11.7 wt% of Zn. Thus, it can summarize that the synthesized ZnO-decorated GO nanocomposite material was high in purity. The findings in this study proved that ZnO NPs loading in ZnO-decorated GO nanocomposite material were successfully synthesized by sol-gel method. A ZnO-decorated GO nanocomposite material with layering ZnO NPs on GO nanosheet was produced.

Keywords: Sol-gel; Zinc oxide; Graphene oxide; Nanoparticles

ABSTRAK

Kajian ini bertujuan untuk sintesis serakan zink oksida (ZnO) zarah nano (NPs) yang baik pada permukaan kepingan nano grafen oksida (GO) dengan cara praktikal iaitu melalui kaedah sol-gel. Zink asetat diyahhidratkan (Zn(CH$_3$COO)$_2$·2H$_2$O) digunakan sebagai pelopor ZnO dan etanol mutlak sebagai pelarut. Sebanyak 1 berat peratus (wt%), 5 wt%, 10 wt%, dan 20 wt% ZnO dimuatan ke atas kepingan nano GO. Beberapa analisis telah dilaksanakan untuk mencirikan komposit nano GO-ZnO yang dihasilkan. Keputusan XRD memperahkan beberapa puncak luas dalam lingkungan 25° hingga 80° di mana ia menunjukkan kepada ZnO dalam komposit nano GO-ZnO. Dengan melakukan analisis potensi zeta, keputusan menunjukkan bahawa terdapat peningkatan caj permukaan negatif pada komposit nano GO-ZnO. Keputusan uji zeta juga mendapatkan bahawa saiz zarah hidrodinamik komposit nano GO-ZnO menjadi besar apabila banyak ZnO dimuatan. FESEM mikrograf menunjukkan bahawa ZnO NPs berbentuk sfera muncul pada permukaan kepingan nano GO dihักukat selanjutnya oleh EDX di mana kandungan komposit nano GO-ZnO terdiri daripada 71.3% berat C, 17% berat O, dan elemen tambahan 11.7% berat Zn. Oleh itu, dapat disimpulkan bahawa komposit nano GO-ZnO yang disintesis adalah tinggi ketulenannya. Penemuan yang terdapat dalam kajian ini membuktikan bahawa pemukan ZnO NPs pada komposit nano GO-ZnO berjaya disintesis oleh kaedah sol-gel. Komposit nano GO-ZnO dengan salutan seragam dan ikatan yang kuat antara ZnO NPs dan permukaan kepingan nano GO telah dihasilkan.

Kata kunci: Sol-gel; Zink oksida; Grafen oksida; Zarah nano
INTRODUCTION

There are ZnO NPs has emerged as one of the highly promising metal NPs used in many applications, including gas sensor, protective coatings, electro photography, solar cells (Hasnidawati et al. 2016), healthcare, optics, ultraviolet, and blue light emitting diodes, and in the evolution of membrane area, due to their excellent properties of antimicrobial, anti-corrosive, thermal, and mechanical stability (Fan & Lu 2005; Vaseem et al. 2010; Wang et al. 2002), high catalytic activity, strong adsorption capacity, low toxicity, and environmental friendly feature (Vaseem et al. 2010).

In membrane technology area, incorporation of ZnO NPs into polymer matrix has proven could improve the implementation of membrane such as water flux, rejection capability, and anti fouling property (Liang et al. 2012; Shen et al. 2012; Zhao et al. 2015). These positive improvements attributed by ZnO NPs make it an excellent NPs filler for the improvement of membrane quality and lead to high efficiency of protein concentrated and fractionated without change the properties of food components.

However, high surface energy of ZnO NPs often resulted an agglomeration in membrane matrix, leading to low functional surface area and apparently gives drawback for its applications (Amat et al. 2012). GO nanosheet have recently introduced as a reliable platform for better nanomaterial dispersion (Bao et al. 2011). GO formed from oxidation of graphite were fascinating carbon material that carried hydroxyl and epoxy functional groups on their basal planes, in addition carboxyl and carboxyl groups located at the sheet edges (Qin et al. 2014). These functional groups of GO nanosheet are essential in forming hybrid nanostructures with various kinds of NPs such as silver (Ag), silicon dioxide (SiO₂), and titanium dioxide (TiO₂) (Ng et al. 2013).

These functional groups makes GO nanosheet become more hydrophilic, enable the GO to readily exfoliate in water to yield stable dispersion (Paredes et al. 2008). The synergistic effects between the hydrophilic-layered GO nanosheet and NPs make it adaptable. By having these interactions, the combination of both ZnO and GO would form nanohybrids with special properties. Recently, many researchers have explored the synthesize of ZnO-decorated GO nanocomposite material and it exhibit potential application in photocatalysis and supercapacitor (Gayathri et al. 2014; Li et al. 2012). In addition, Chen et al. (2013) has successfully proved that ZnO-decorated GO nanocomposite material has been shown to have better photo catalytic activity.

Researchers have developed variety of technique to synthesize these nanomaterials. The principle methods that have been put forward in the literature to synthesize these materials, namely spray pyrolysis, hydrothermal, chemical vapor condensation, precipitation, sol-gel method (Tavakoli et al. 2007; Van Werde et al. 2002), electrochemical method, and electrochemical depositions (Chu & Li 2012; Stypuła et al. 2014). The crystallite size and morphology of nanomaterials strongly depend on synthesis method. It is particularly interesting to synthesize ZnO by the sol-gel process because of its simplicity, reproducibility, low cost of the chemicals (Lučko Golič et al. 2011), and sol-gel chemistry is effective for controlling morphology and reactivity of solids (Benhebal et al. 2013). Instead, in recent years it has shown improvement for dispersing materials, promoting good homogeneity and purity (Bodson et al. 2010; Lambert et al. 2004).

Herein, we synthesize the ZnO-decorated GO nanocomposite material using sol-gel method. A uniformly distributed of ZnO NPs onto GO nanosheet to produce ZnO-decorated GO nanocomposite material was investigated. In this work, the effects of varying the ZnO loadings onto ZnO-decorated GO nanocomposite material were characterized and reported in this study for better understanding the properties of nanohybrid material.

METHODOLOGY

MATERIALS

Graphite powder (particle size < 50 µm), zinc acetate dehydrates (Zn(CH₃COO)₂·2H₂O), oxalic acid (C₂H₂O₄·2H₂O), and sodium nitrate (NaNO₃) were bought from Merck Co., Germany. 98% sulfuric acid (H₂SO₄), 38% hydrochloric acid (HCl), 30% hydrogen peroxide (H₂O₂), absolute ethanol, and potassium permanganate (KMnO₄) were bought from Sigma Aldrich, USA. Analytical grade of chemicals was used.

SYNTHESIS OF GO

Following the modify Hummers method, graphite powder was chemically oxidized at low temperature, resulting a brown colloidal suspension known as GO (Hummers JR & Offeman 1958). Typically, 5 g of graphite powder and 2.5 g of NaNO₃ were added into 115 mL of 98% H₂SO₄, and stirred at 300 rpm for 1 h under room temperature. The mixture was further stirred for 30 min in ice bath until it hit 10°C before KMnO₄ was gradually added under continuous stirring. The mixture was left t for 2 h while keeping the temperature less than 10°C. It was then stirred at 35°C for another h. Following, 230 mL deionized water was added to dilute the mixture and stirred at temperature below 100°C. Further dilution was done with 300 mL deionized water and stirred for another h to remove metal ion. The reaction was terminated by adding 10 mL of 30% H₂O₂ into the mixture. The dispersion was washed, rinsed with 5% HCl aqueous solution, followed by second rinse with deionized water. The final fine brown powder product was freeze dried under vacuum at -40°C for further study.

SYNTHESIS OF ZNO-DECORATED GO NANOCOMPOSITE MATERIAL

Initially, 0.3 g of GO was dispersed in 10 mL of absolute ethanol via sonication to produce GO suspension. Then, Zn(CH₃COO)₂·2H₂O mixed into 90 mL of absolute ethanol under vigorous stirring at 65°C for 30 min. Next, Zn(CH₃COO)₂·2H₂O/absolute ethanol mixture was added slowly into GO suspension with stirred for 20 min at 45°C.
Later, 1 g of \(\text{C}_2\text{H}_5\text{O}_2\text{H}_2\text{O}\) dissolved in 50 mL absolute alcohol was poured into the mixture solution and let it stir for another 1 h and 30 min. Collect the precipitate by centrifugation and dried at 65°C overnight in an oven, followed by calcination at 400°C for 3 h to get final product. ZnO-decorated GO nanocomposite material with different weight percent of ZnO NPs (1%, 5%, 10%, and 20%) were earned by manipulating the ration of \(\text{Zn(CH}_2\text{COO)}_2\cdot\text{H}_2\text{O}\) in the synthesis process.

SYNTHESIS OF ZNO-DECORATED GO NANOCOMPOSITE MATERIAL

X-RAY DIFFRACTION (XRD)

The crystal phase composition of synthesized ZnO-decorated GO nanocomposite material was interpret using D8 Advance X-ray diffractometer (Bruker AXS, Germany) with CuKα radiation source (\(1.5406\) Å) at 20 scan range of 5° to 85°. The average crystallite size (D) of the nanomaterial was calculated from its XRD pattern based on Scherrer’s equation.

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

Where \(k\) is the Scherrer constant (0.89), is the X-ray wavelength (0.1540 nm), \(\beta\) is the peak width at half maximum (°), and \(\theta\) is the Bragg diffraction angle (°).

FOURIER TRANSFORM INFRARED SPECTROSCOPY (FTIR)

Nicolet 6700 FTIR spectrometer (Thermo Fisher Scientific Inc., USA) recorded the FTIR spectrum coupled with a diamond crystal at incidence angle at 45° to point out possible functional groups of GO nanosheet. 32 scans with 4 cm\(^{-1}\) resolutions to obtained and spectrum and has operated at wavelength from 4000 cm\(^{-1}\) to 500 cm\(^{-1}\).

FIELD EMISSION SCANNING ELECTRON MICROSCOPY (FESEM) AND ENERGY DISPERSIVE X-RAY SPECTROSCOPY (EDX)

High resolution FESEM Gemini model SUPRA 55VP (Carl Zeiss, Germany) was used to investigate the morphology and distribution of the GO nanosheet and ZnO NPs. All nanomaterials were covered with a thin layer of platinum under vacuum to reduce the charge on the surface so that contrastive images formed. FESEM imaging was observed at the magnification of 100 k for finding out the shape of ZnO NPs deposition on GO. In a meantime, the element composition of the same sample was analyzed using EDX, Oxford INCA PentaFETX3 (Oxford, UK).

HYDRODYNAMIC PARTICLE SIZE AND ZETA POTENTIAL

The zeta potential and hydrodynamic particle size of the synthesized ZnO-decorated GO nanocomposite material were examined using ZetaSizer Nano-ZS (Malvern instruments Inc., UK) which works on the principle of dynamic light scattering (DLS) where it measures the intensity of the laser light scattered from suspended particles. Prior hydrodynamic particle size and zeta potential measurement, ZnO-decorated GO nanocomposite material was dispersed in water at the concentration of 0.1 w/v% (weight of ZnO-decorated GO nanocomposite material/volume of water, solvent), and sonicated using a bath sonicator for 30 min to ensure a uniform suspension.

RESULTS AND DISCUSSIONS

CHARACTERIZATION OF GO AND ZNO-DECORATED GO NANOCOMPOSITE MATERIAL

The XRD pattern of graphite, GO, and 20 wt% of ZnO loaded in ZnO-decorated GO nanocomposite material are set out in Figure 1. Pristine graphite depicted a tenacious peak at 26.5° and relocated to 10.8°, as graphite powder was chemically oxidized to form GO through modified Hummers method. Exfoliation of graphite into GO (Ahmed et al. 2016; Chao et al. 2008) disrupted the basal plane of graphite by adding carbonyl (C=O) and carboxyl (C-OH) groups located at the edge of graphene nanosheet (Wang et al. 2014; Williams & Kamat 2009).

![XRD pattern of graphite, GO, and 20 wt% of ZnO loaded in ZnO-decorated GO nanocomposite material](image)

FIGURE 1. XRD pattern of graphite, GO, and 20 wt% of ZnO loaded in ZnO-decorated GO nanocomposite material

Followed by decoration of ZnO NPs on GO nanosheet, the GO nanosheet peak at 10.8° was gone. It was reported that the exfoliation of regular stacks of GO or graphite could diminish or even disappeared the diffraction peak (Chao et al. 2008). However, ZnO-decorated GO nanocomposite material demonstrate some broad peaks in the range 25° to 80° which assigned by ZnO NPs decorated on GO nanosheet (Xu et al. 2014). The average crystallite size of ZnO-decorated GO nanocomposite material was calculated from its XRD pattern based on Scherrer’s equation. The calculated result showed that the average crystallite size of ZnO-decorated GO nanomaterial was 9.92 nm which was in the range 5-20 nm as reported by Chung et al. (2017).
The presence of functional groups on GO nanosheet was put forward by FTIR spectra as shown in Figure 2. Peak at 1627.40 cm\(^{-1}\) was pointed to C=O stretching vibration whereas the broad peak around 3586.10 cm\(^{-1}\) resemble O-H stretching vibration (Chung et al. 2017). In addition, two bands at 1070.40 cm\(^{-1}\) and 1417.13 cm\(^{-1}\) are referred to by C-OOH and alkoxy (C-O) groups, respectively. The presence of the aforementioned functional groups confirmed the successful of graphite oxidized to GO, coincident with other researchers (Anitha & Jayakumari 2015; Kavitha et al. 2013). However, black color of ZnO-decorated GO nanocomposite material limited the analysis using FTIR as poor infrared transmittance was produced due to the strong absorption characteristic from the dark color of sample. Therefore, ZnO-decorated GO nanocomposite material was characterized using FESEM and EDX analysis.

The morphology and distribution of GO and ZnO NPs were characterized using FESEM, while the elemental compositions that has been existent on both nanomaterials were inspect using EDX. Figure 3(a) and Figure 3(b) showed the FESEM micrographs and EDX spectra of GO and ZnO-decorated GO nanocomposite material, respectively. EDX analysis confirmed the deposition of ZnO NPs (white spots in FESEM micrograph) on ZnO-decorated GO nanocomposite material. As depicted in Figure 3(b), the element compositions attributed by GO consist of carbon and oxygen. In which their weight percent were 65 wt% and 35 wt% for carbon and oxygen, respectively. On the other hand, ZnO-decorated GO nanocomposite material was having additional element, zinc (Zn) as compared to GO. The weight percent for carbon, Zn, and oxygen element existed in ZnO-decorated GO nanocomposite were 71 wt%, 11.7 wt%, and 17 wt% respectively. The EDX analysis has confirmed the success deposition of ZnO NPs on GO in synthesizing ZnO-decorated GO nanocomposite material with high purity.

Table 1 showed the hydrodynamic particle size of ZnO-decorated GO nanocomposite material with different weight percent of ZnO loading. The hydrodynamic particle size was first decreased with the increasing of ZnO loading up to 5%. Larger hydrodynamic particle size at 1 wt% of ZnO loading was noted from the result may be attributed to the rapid combination of reagents during particles synthesized as reported by (Duffy et al. 2007). Zhang et al. (2009) reported that appropriate amount of mixture between precursor and solvent causing higher viscosity and faster increase in evaporative gelation of solution will complete the solution-gel transition in a short time and prevent aggregation. Thus, other reason of larger hydrodynamic particle size at 1 wt% of ZnO loading may due to the amount of Zn (CH\(_3\)COO)\(_2\)·2H\(_2\)O was not appropriate with oxalic acid causing agglomeration occurred.

The hydrodynamic particle size was increased as the weight percent of ZnO was loaded further at higher amount. When the weight percent of ZnO was increased, it leads to the increase higher viscosity, where an increase in hydrodynamic particle size is expected. Since the frequency of particle collision is a strong function of particle concentration (Marsalek 2014; Friedlander 2000), ZnO-decorated GO nanocomposite material at high ZnO loading has high risk to collide, and agglomerate in suspension, hence resulted an increase in hydrodynamic particle size (Domingos et al. 2009; Suttiponparnit et al. 2011).

Figure 4 showed the zeta potential of ZnO-decorated GO nanocomposite material with different ZnO loading. Zeta potential is denoted for surface charge of the particle (Mandzy et al. 2005). In general, zeta potential below -30 mV or beyond +30 mV considered as the criterion for a
stable dispersion, high electrostatic repulsion is expected to manage the particles in suspension to repel each other (Ho et al. 2017; Zhang et al. 2013).

As depicted by Figure 4, ZnO-decorated GO nanocomposite material with desired ZnO loading were having zeta potential value of -17.4 mV, -16.90 mV, -20.30 mV, and -28.23 mV, respectively, suggesting ZnO-decorated GO nanocomposite material was approaching stability to produce uniform NPs distribution at higher ZnO loading, which expected a uniform nanomaterials distribution when incorporation with variety applications for enhancing the process efficiency.

CONCLUSION

In this study, ZnO-decorated GO nanocomposite material was successfully synthesized via sol-gel method. The inclusion effect of difference weight percent of ZnO loading in ZnO-decorated GO nanocomposite material on crystallite phase, hydrodynamic particle size, and zeta potential were also investigated. Besides, the morphological and structure properties of ZnO-decorated GO nanocomposite material were investigated. It was found that the ZnO-decorated GO nanocomposite material was composed of spherical-like ZnO NPs anchored on GO sheet. The XRD patterns of ZnO-decorated GO nanocomposite material indicate the formation of wurtzite ZnO NPs. The increasing weight percent of ZnO loading results in increasing hydrodynamic particle size, which the range size generated approximately 28 nm to 57 nm, and negatively zeta potential value from -16.90 mV to -28.23 mV. Therefore, the results demonstrated a stable suspension of ZnO-decorated GO nanocomposite material and thus these features make the ZnO-decorated GO nanocomposite material an excellent candidate for various applications relating to the enhancing process efficiency.

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