

MALAYSIAN JOURNAL OF ANALYTICAL SCIENCES

Published by The Malaysian Analytical Sciences Society

ISSN 1394 - 2506

THE SYNTHESIS OF ZINC OXIDE/CARBON SPHERES NANOCOMPOSITES AND FIELD ELECTRON EMISSION PROPERTIES

(Sintesis Nanokomposit Zink Oksida/Sfera Karbon dan Sifat Pemancaran Elektron Medan)

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Received: 19 November 2016; Accepted: 6 March 2017

Abstract

Zinc oxide (ZnO)/carbon spheres (CS) nanocomposites were successfully synthesised using waste engine oil as precursor for the CS production. ZnO nanorods were grown using sol-gel immersion method with MgZnO as the seeded catalyst and thermal chemical vapour deposition was used to synthesise CS. Different configurations of ZnO/CS structures were prepared i.e. CS-coated ZnO and ZnO-coated CS. The structures of composite samples were analysed using field emission scanning electron microscopy (FESEM), Energy-Dispersive X-ray (EDX), micro-Raman and X-ray Diffraction Spectroscopy (XRD). FESEM observations revealed the structural changes of pristine ZnO and CS in composite structures. The as-present of ZnO or CS was believed to affect the subsequent growth of another structure. Field electron emission (FEE) properties of both nanocomposites were also investigated. It was found that ZnO-coated CS sample has better FEE properties with lower turn-on (3.48 V μ m⁻¹) and threshold field (6.35 V μ m⁻¹) obtained at current density of 0.1 and 1 μ Acm⁻², respectively. This study highlighted that nanocomposites of ZnO and CS have successfully enhanced the field emission performances of materials compared with pristine ZnO or CS due to the structural changes of material emitter.

Keywords: zinc oxide, carbon spheres, waste engine oil, field electron emission

Abstrak

Nanokomposit zink oksida (ZnO)/sfera karbon (SK) telah berjaya disintesis menggunakan minyak enjin terpakai sebagai pelopor untuk penghasilan SK. Nanorod ZnO telah ditumbuhkan menggunakan kaedah rendaman sol-gel dengan MgZnO sebagai pemangkin berbenih dan pemendapan wap kimia terma telah digunakan untuk mensintesis SK. Konfigurasi yang berlainan bagi struktur ZnO/SK telah disediakan iaitu SK bersalut ZnO dan ZnO bersalut SK. Struktur sampel komposit dianalisis menggunakan *Mikroskop* Imbasan *Elektron* Pancaran *Medan* (FESEM), penyerakan tenaga sinar-X (EDX), spektroskopi mikro-Raman dan pembelauan sinar-X (XRD). Pemerhatian FESEM menunjukkan perubahan struktur ZnO dan CS tulen dalam struktur komposit. ZnO atau CS sedia ada dipercayai memberi kesan kepada pertumbuhan seterusnya struktur lain. Sifat pemancaran elektron medan (PEM) bagi kedua-dua nanokomposit juga turut dikaji. Kajian mendapati bahawa sampel ZnO bersalut SK mempunyai ciri-ciri PEM yang lebih baik dengan nilai medan permulaan (3.48 Vµm⁻¹) dan ambang (6.35 Vµm⁻¹) yang lebih rendah diperolehi pada ketumpatan arus 0.1 dan 1 µAcm⁻². Kajian ini menekankan nanokomposit ZnO dan SK telah berjaya meningkatkan prestasi pemancaran medan bahan berbanding dengan ZnO atau CS tulen disebabkan oleh perubahan struktur bahan pemancar.

Kata kunci: zink oksida, sfera karbon, minyak enjin terpakai, pemancaran elektron medan

Introduction

Field emission is a quantum process of electron emission from a material where the electrons in Fermi level tunnel the energy barrier to vacuum level under high applied electric field [1]. Emission characteristics of a material are based on the Fowler-Nordheim (F-N) theory as followed in Eq. 1 [2]:

$$J = \frac{A(\beta E)^2}{\varphi} \exp\left(\frac{B\varphi^{3/2}}{\beta E}\right)$$
(1)

The current emission density (J) at applied electric field (E) can be calculated from the above equation. A and B are constants of $1.54 \times 10^{-6} \text{ AV}^2 \text{eV}$ and $6.83 \times 10^9 \text{ eV}^{-3/2} \text{Vm}^{-1}$, respectively. Field enhancement factor of emitter (β) is an important FEE characteristic and can be calculated from the slope of ln (J/E²) vs 1/E curve using the equation $\beta = -B\phi^{3/2}/\text{slope}$. A typical benchmark that have been reported in the literature for FEE characterization comparison are the turn-on and threshold field, maximum current obtained (before the destruction of the emitter) as well as emission stability. Turn-on field for FEE is the field required to emit current density which is greater than the electrical noise-floor [3]. There is no agreement on the value of current density to be specified and the definition of turn-on and threshold field, it is determined by the group research themselves. For example, Ghosh et al. [4, 5] and Srivastava et al. [6] defined turn-on and threshold field to be the field required by the film to produce current density of 10 μ Acm⁻² and 1 mAcm⁻² respectively. While Bonard and co-worker [90] specified the current density of 10 μ Acm⁻² and 10 mAcm⁻² respectively. Since there is no specific definition of both turn-on and threshold field and as current density does not increase linearly with applied field, the result cannot be accurately compared. The comparisons between the results were only a rough estimation to elaim that theirs is better than others. In this study, they are defined as the current density at applied field of 0.1 and 1 μ Acm⁻², respectively for turn-on and threshold field.

To increase the FEE performance of the materials, it can be composited to further enhance the electron emission performance of those nanocomposites. Take carbon material such as carbon nanotubes (CNTs) as example, in view of its potential application as good electron emitters, however it can be easily degraded and damaged by the following factors [7-10], therefore the improvement of the FEE can be achieved by fabricating CNTs and zinc-oxide (ZnO) as nanocomposites [11-13]. The order of the configuration of CNTs/ZnO composites is the crucial component that lead to the enhancement of field emission ability, where it frequently occurs that the materials fabricated at the top is more likely to help the FEE performances. ZnO/CNTs nanocomposites structure have been reported to improve the FEE performance of material [1, 11-14]. In our recently published work [11], the investigation was done on the composite configuration to enhance the FEE performance by using two different configurations of ZnO/vertically aligned CNTs (VACNTs) where the order of the nanocomposites plays an important role in improving the FEE properties of pristine VACNTs.

The work revealed that the VACNTs-coated ZnO nanocomposites configuration showed a lower turn-on and threshold field at only 0.49 and 1.41 Vµm⁻¹ as compared to the ZnO-coated VACNTs of 1.60 (turn-on) and 2.03 (threshold field) Vµm⁻¹ respectively. The presence of different ZnO nanostructures including nanorods, nanoflowers and nanorods-nanoflakes for ZnO-coated CNTs configuration were seen to play significant role towards enhancing FEE performances [13]. Among the ZnO nanostructured used the growth of CNTs on ZnO nanoflower demonstrated the highest FEE performance of 0.8 Vµm⁻¹ at 1 uAcm⁻² of turn-on field. ZnO is a semiconducting material with wide direct band gap (3.37 eV), large binding energy of 60 meV and and work function of 5.3 eV [1]. It has been applied in many electronic devices due to its unique and versatile properties [15-17]. ZnO nanostructures also showed promising application in field emission devices due to its high mechanical strength, chemical stability and good optical properties [18, 19]. On the other hand, carbon materials are well known as a conducting material which is good for electron emission application. One type of carbon materials namely carbon spheres (CS) has been reported to have a comparable field emission properties as compared to other materials [20].

Here we presented the composite structures of ZnO/CS and their FEE properties. To the best of our knowledge, this is the first report on the FEE of the CS synthesised from waste engine oil (WEO) with two different configurations of nanocomposite; CS-coated aligned ZnO nanorod and ZnO nanorod-coated CS sample. Recently, we reported the use of WEO for the production of another carbon material namely CNTs [21] and amorphous Al-Cu alloy nanowires decorated with CS [22]. The use of waste [23-27] and natural precursors [28-36] to produce carbon materials may not only be beneficial for nanotechnology research field but also preserve the environmental sustainability at the same time. The CS were synthesised using thermal chemical vapour deposition (TCVD) method while the ZnO were synthesised using the sol-gel method. The structural and field emission properties of ZnO/CS nanocomposites samples were then systematically investigated.

Materials and Methods

The production of ZnO/CS was performed in two different synthesis processes which were (i) the synthesis of ZnO nanorods using sol-gel immersion process and (ii) the synthesis of CS via thermal chemical vapour deposition (TCVD) method. Two different configurations were prepared as illustrated in Figure 1. For the CS-coated ZnO sample, ZnO nanorods were initially grown followed by the synthesis of CS and vice versa for the ZnO-coated CS sample.



Figure 1. Different configurations of composite materials (a) CS-coated ZnO and (b) ZnO-coated CS

The synthesis procedures of CS from WEO precursor was similar to previous report [21, 22]. However, different synthesis parameters were used. An amount 3 ml of WEO mixed with 5.33 wt% ferrocene catalyst was heated at 450 °C in precursor zone and deposited onto Si substrate in 800 °C synthesis furnace. The synthesis process was carried out for 30 min under Ar gas flow at the rate of 0.25 Lmin⁻¹. Meanwhile, the synthesis of ZnO nanorods was adopted from [37]. MgZnO was initially deposited as the seeded catalyst. Collected samples were characterized using field emission scanning electron microscopy (FESEM-Hitachi SU8020), energy-dispersive X-ray (EDX), micro-Raman spectroscopy (Horiba Jobin Yvon-DU420A-OE-325) and X-ray diffraction (XRD) instrument (Bruker AXS D9). Field emission properties of the samples were measured using FEE equipment (Hewlett-Packard 34401A multimeter) at emission area of 0.1 cm² and distance between two electrodes was 100 µm.

Results and Discussion

Figure 2(a) – (f) present the FESEM images of the prepared ZnO nanorods, CS and their nanocomposites with different configurations. It was observed in Figure 2 (a) that the ZnO nanorods were vertically aligned with length up to 1.6 μ m. Wide diameter distribution of the rods in the range of 42.8 – 285.7 nm were measured. This was due to the non-unifom MgZnO particles size as the template for the ZnO nanorods growth. The growth of ZnO nanorods was also believed occurred on the collection of agglomerated MgZnO particles. High density of CS was obtained with diameter of the spheres were ranged from 0.17 to 2.57 μ m as seen in Figure 2(b). The morphology of CS was modified after grown on ZnO nanorods as seen in Figure 2(c) and (d). Most of the spheres were decorated with short tube-like particles and grains which seemed to grow covering the CS. This was believed due to the adsorption of ZnO particles on the surface of CS which further become the catalyst for the secondary growth of short tube-like carbon structure. Somehow, the perfectly round shape of CS was still observed with diameter range from 1.1 to 2.7 μ m.

Figure 2(e) and (f) show the FESEM observation of ZnO-coated CS sample. High density and uniformly grown ZnO nanorods on CS were observed. Slightly different to the pristine aligned ZnO nanorods, flower-like ZnO

nanostructures consisted of many hexagonal rods were also produced on the top of ZnO nanorods. The diameter of the rods was observed to be between 31.3 to 187.5 nm, in the smaller range as compared to pristine ZnO. This indicated the important presence of CS as the nucleation site for the growth of ZnO nanostructures. MgZnO particles were believed to be well deposited with uniform size on the CS, thus produced a better growth of ZnO nanostructures. Only small amount of CS was still observed as seen in the inset of Figure 2(f). The perfect alignment of ZnO nanorods cannot be observed due to the spherical shape of CS template. ZnO nanorods were initially grown on the uneven surface of MgZnO-coated CS template. Once the saturated condition was achieved, nucleation of aligned ZnO nanorods got slower. There was no more space for the growth of nanorods on the MgZnO-coated CS template, hence the ZnO particles started to nuclei on the top of the rods. Then, the aggregated ZnO particles formed the flower-like structure as can be observed in Figure 2(e)-(f).



Figure 2. FESEM images of (a) ZnO nanorods, (b) CS, (c-d) CS-coated ZnO and (e-f) ZnO-coated CS nanocomposites

EDX analysis was performed to investigate the elemental compositions of the composite materials. Figure 3 show the EDX spectrum of both ZnO-coated CS and CS-coated ZnO samples. Carbon, oxygen, and Zn elements were detected from overall structure. The Si peaks was believed to come from the substrate used. The average atomic of zinc and oxygen was found to be 1:1 which indicated a good stoichiometric ratio of ZnO. Moreover, low amount of Mg element can be detected in this sample. The concentration of Zn atoms decreased in the CS-coated ZnO sample. This was in line with FESEM observation where during the synthesis of CS at high temperature of 800°C, the ZnO nanorods were fully etched and turned to be ZnO flakes structure. In the meantime, Zn and oxygen atoms were also suspected to be adsorbed on the surface of CS.



Figure 3. EDX analysis showing elemental compositions of ZnO/CS nanocomposites (a) ZnO-coated CS and (b) CS-coated ZnO

Micro-Raman spectra of the as-prepared samples are presented in Figure 4. The micro-Raman spectrum of pristine CS shows the *D* (disorder) and *G* (graphitic) band at 1357.70 and 1598.2 cm⁻¹, respectively with the intensity ratio of *D* and *G* band (I_D/I_G) ratio was found to be 0.79. Micro-Raman spectrum of ZnO nanorods only shows one typical ZnO characteristics at 438.18 cm⁻¹ attributed to the E₂ (high) modes. It also confirmed wurtzite hexagonal phase of ZnO nanorods. Peak at 303.76 cm⁻¹ was derived from the use of Si as the substrate. Micro-Raman spectra of ZnO/CS nanocomposites present both characteristic peaks of materials. ZnO-coated CS remained show the wurtzite hexagonal phase of ZnO characteristic at 438.18 cm⁻¹ which presented by a clear E₂ (high) mode.

Only CS characteristics can be observed from the Raman spectrum of CS-coated ZnO sample. However, micro-Raman spectrum at lower shift range Figure 4(b) showed a sharp low intense peak at 444.13 cm⁻¹ which corresponded to Raman active mode of wurtzite ZnO structure (E_2 high). This mode was shifted to a higher number. It might due to the increase of the stress or defect in the ZnO crystal [38]. Moreover, I_D/I_G ratio of the CS increased to be 0.91 (ZnO-coated CS) and 0.88 (CS-coated ZnO) after being composited with ZnO. These indicated the lower crystallinity of CS in both composite structures as compared to the pristine CS. The reasonable explanation to these occurrences was due to the changes of CS structure after being combined with ZnO. In case of ZnO-coated CS sample, annealing process of MgZnO thin layer and ZnO nanorods alter the structure of CS where it introduced more defect of C–C bonding in the spheres. The highest ratio of 0.91 was obtained from this structure. In addition, for the growth of CS on ZnO nanostructures, defect-induced from the bonding of carbon from CS and oxygen from the ZnO nanostructures was responsible to the higher defect of CS [13].



Figure 4. (a) Micro-Raman spectra of ZnO/CS nanostructures and (b) Raman profile at lower wavenumber of CScoated ZnO structure shows the shift of E₂ high mode due to the compressive stress

Typical XRD patterns of ZnO/CS nanostructures are presented in Figure 5. Both samples show the characteristic peaks of ZnO and carbon. Similar to the pristine ZnO nanorods, the most intense peak was observed at \sim 34° along the (002) orientation [19]. ZnO-coated CS sample presents a broad and low intense peak at 43.40° which attributed to carbon phase from CS. Another ZnO peak was shown by a low intense peak at 47.49° along the (102) orientation. Diffraction peaks of ZnO in CS-coated ZnO sample are located at 20 of 34.49, 36.15, 47.60 and 56.20° which are in good agreement with the standard of hexagonal ZnO crystal (PDF-01-080-0075). Meanwhile, carbon phase at 44.67 and 52.3° are well matched with the standard of cubic carbon crystal (PDF-01-080-0017) with the lattice parameter of 3.539 Å.



Figure 5. XRD patterns of CS/ZnO nanostructures composites

Figures 6 and 7 present the J-E and F-N plot of different ZnO and ZnO/CS nanocomposites materials. The measurements were taken at fixed electrodes separation of 100 μ m and applied voltage was increased from 0 to 1000 V. Table 1 presents the summary of FEE characteristics of the as-prepared samples. Several repeated measurements have been conducted on the CS sample, but the current emission was still not observed. Due to its spherical structure, CS are less suitable to be applied as electron emitter. ZnO nanorods possessed moderate turn on

and threshold fields at 5.65 and 7.34 V/µm, respectively with maximum current density of 3.46 µA/cm². Composite structures of ZnO/CS successfully enhanced the FEE properties. Turn on and threshold values of CS-coated ZnO sample were founded to be at 6.83 and 7.88 V/µm and maximum current density of 2.51 µA/cm². Modification of CS structure with the presence of short tube-like carbon structure which covered the CS was believed promoted a better emission site as compared to pristine CS. Interestingly, ZnO-coated CS structure where the ZnO nanostructures were grown on CS as the template exhibited an obvious decrement in the turn on value to be as low as 3.48 V/µm and the threshold field was also decreased to be 6.35 V/µm. This structure presented a better FEE performance compared with the pristine ZnO nanorods as well as CS-coated ZnO composite structure. In addition, highest β value up to 5485 and maximum current density of 3.78 µA/cm² were obtained. Higher conductivity of CS compared to ZnO nanorods promoted a better electron transfer from the CS through ZnO nanorods structures to the vacuum level at high applied voltage. Moreover, the high aspect ratio of ZnO nanostructures with a good crystallinity and uniform rods diameter were also responsible for the high β value of ZnO-coated CS composite material [6].



Figure 6. The J-E curves of the ZnO and ZnO/CS nanocomposites materials



Figure 7. F-N plots of ZnO and ZnO/CS nanocomposites materials (a) ZnO, (b) CS-coated ZnO and (c) ZnO-coated CS

Sample	Turn on Field at 0.1 μΑ (V/μm)	Threshold Field at 1 μΑ (V/μm)	J Max (μA/cm ²)	β
ZnO	5.65	7.34	3.46	2452
ZnO-coated CS	3.48	6.35	3.78	5879
CS-coated ZnO	6.83	7.88	2.51	755

Table 1. FEE characteristics of CS, ZnO nanorods and their composite materials

To study the emission current stability, a constant electric field was applied to the ZnO nanorods and ZnO/CS nanocomposites materials for one hour duration time. The current vs time graphs for both samples are presented in Figure 8 and the detail of their current emission fluctuation order is given in Table 2. The current emission of ZnO nanorods was moderately stable since frequent fluctuations to a high current value were observed. This was due to the wide diameter range of ZnO nanorods. Smaller diameter rods proportional with the length and rods spacing is preferable for a better FEE performance. Although ZnO nanorods current emission was less stable, the emission of ZnO nanorods was found to be higher in the order of 10⁻⁶ A than the CNTs which was in the order of 10⁻⁷ A [11]. ZnO nanomaterials have negative electron affinity [18] which can support the current emission due to their ability to emit electrons to the vacuum with little energy loss.



Figure 8. Current stability measurement of (a) ZnO and (b)-(c) ZnO/CS nanostructures composites

Sample	Current Fluctuation Order (Ampere)
ZnO	$10^{-8} - 10^{-6}$
ZnO-coated CS	$10^{-9} - 10^{-6}$
CS-coated ZnO	$10^{-7} - 10^{-6}$

Table 2.	Current emission fluctuation order of carbon material	, ZnO nanorods			
and their composite structures					

The growth of ZnO nanorods on CS sample gave better emission stability as compared to pristine ZnO and CScoated ZnO samples. This was believed due to the presence of as-grown CS and high uniformity with smaller ZnO nanorods, flower-like ZnO nanostructures size which is more favourable for electron emission application. High conductivity of carbon materials promoted better electrons transfer and led to the enhancement of field emission. Moreover, the carbon materials acted as the additional catalyst which improved the uniformity and crystal quality of ZnO nanostructures.

Conclusion

ZnO/CS nanocomposites were successfully synthesised using the combination of sol-gel immersion and TCVD method. Waste material precursor namely WEO was used to produce CS in this study. The structure of ZnO and CS were modified after being composited. In case of CS-coated ZnO, the structural change of CS was believed due to the adsorption of ZnO particles on the surface of CS. Meanwhile, the as-presence of CS in ZnO-coated CS sample has promoted a better growth ZnO nanostructure. This structure also presented the best FEE performances with the lowest turn on (3.48 V/µm) and threshold field (6.35 V/µm) as well as highest maximum current density (3.78 μ A/cm²) and field enhancement factor up to 5879 with better emission current stability. The presence of conductive carbon material which is CS promoted the well electron transfer from the CS through the ZnO nanostructures to vacuum level at high electric field. A better structure of ZnO nanorods with uniform diameter size was also responsible to the better FEE properties of this nanocomposites structure. It can be concluded that the ZnO-coated CS nanocomposites has great future applications in field emission devices such as flat panel displays and flat lamps.

Acknowledgement

The authors are thankful to Fundamental Research Grant Scheme (Grant code: 2015-0154-102-02) and UPSI for financial and technical supports of this work.

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