

Morphology and optoelectronic properties of ZnO rod array/conjugated polymer hybrid films

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Abstract

Vertically aligned zinc oxide (ZnO) nanorods were grown on the ITO glass and then coated with the conjugated polymer poly(2,3-dibutoxy-1,4-phenylene vinylene) (DB-PPV) to make the hybrid films. Nanorods with different diameters were synthesized to study the influences of ZnO nanorod morphology and polymer infiltration on the photocurrent and optical properties of the hybrid films. Increasing the growth time leads to the formation of ZnO rod array with large rod diameter, large surface area and small inter-rod distance. Small inter-rod distance hinders the filling of DB-PPV into the porous ZnO rod microstructure and lowers the PN junction area. It leads to lower photocurrent of the hybrid film. The red shift of the photoluminescence spectra suggests that filling the polymer into the ZnO rod microstructure favors more planar molecular orientations of the conjugated polymers and leads to an increase in the effective conjugation length.

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1. Introduction

Recently, it has been shown that composite materials containing the nanostructured electron accepting component and the electron-donating organic semiconductor in a “bulk heterojunction” structure are promising materials for photovoltaic device [1]. Ultra-fast photoinduced charge transfer occurs between the conjugated polymers and the metal oxide semiconductor such as TiO₂, or ZnO [2].

However, the overall performance of such devices is still disappointing because of limited charge transport and charge separation efficiency. Zinc oxide (ZnO) is of interest because of its availability of low temperature synthesis, the high electron mobility, and the potential for preparing the ordered nanostructure by the solution process. Solution synthesis offers the potential for much lower cost because it eliminates the expenses associated with high temperature and vacuum processing. Besides, solution processing is compatible with roll-to-roll processing of flexible plastic substrates. ZnO nanorods grown

perpendicular to the substrate are particularly interesting. Perpendicularly oriented single-crystalline ZnO nanorod arrays have been recently fabricated on sapphire substrates using VLS [3] and CVD methods [4]. These methods involve complex procedures, sophisticated equipment and high temperature. Recently, O’Brien et al. [5] and Vayssieres [6] reported the low temperature growth of ZnO nanocolumns from aqueous solution. Vertically aligned ZnO nanorods were used to increase the efficiencies of these photovoltaic devices [7,8]. The ordered zinc oxide nanorods acted as electron acceptors and provided the direct path for photogenerated electrons to the collecting electrode. If the electrons diffuse through the film faster than they recombine, the majority of the charge injected into the semiconductor is collected at the transparent conducting electrode. Devices were prepared by infiltrating conjugated polymers into a mesoporous, nanostructured oxide semiconductor network [9].

Increasing the growth time leads to the formation of ZnO rod array with large rod diameter, high surface area and small inter-rod distances. The distances between the rods influenced the difficulty of filling the mesoporous structure with polymer. No effort has been made on the influences of ZnO nanorod array

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structure on the infiltration of conjugated polymer into a mesoporous ZnO rod array and the photocurrent response. In this paper, ZnO nanorods with different average diameters were synthesized by changing the rod growth time. The influences of rod diameter and inter-rod distance on the morphology and the in-situ photocurrent response of vertically aligned ZnO nanorod-DB-PPV polymer hybrid films were studied.

2. Experimental

Perpendicularly oriented single crystalline ZnO nanorods were grown on ITO substrates by the hydrothermal growth method. The procedure consists of two steps: (1) formation of ZnO nanoparticles as seed on ITO glass substrates, and (2) hydrothermal growth of ZnO nanorods in aqueous solution.

At first, 0.01 M zinc acetate [$\text{Zn}(\text{CH}_3\text{COO})_2 \cdot 2\text{H}_2\text{O}$] was dissolved in the ethanol. The solution was cooled to 0 °C. Then, CTAOH (cetyltrimethylammonium hydroxide) was added and the solution was stirred for 30 min. The resulting mixture was then agitated at 60 °C for 2 h to yield a homogeneous and stable colloid solution, which served as the coating solution. After being coated with the colloid solution, the substrates were dried and annealed at 300 °C for 1 h. The ZnO nanorod array was grown at 95 °C in a sealed bottle by immersing the modified substrates in the aqueous solution containing $\text{Zn}(\text{NO}_3)_2$ (0.05 M) and methenamine (0.05 M). ZnO12 and ZnO14 films mean the rod growth time are 2 and 4 h respectively. DB-PPV was dissolved in THF and spin-coated onto the ZnO rod array at 700 rpm for 10 s. DBZ12 and DBZ14 films represent the DB-PPV/ZnO12 and DB-PPV/ZnO14 hybrid films respectively.

XRD studies were carried out with a MAC SCIENCE MXP3 diffractometer. The morphology of the nanorods was characterized using scanning electron microscopy using a HITACHI S-4800 Field Emission Scanning Electron Microscope (FESEM). The photoluminescence spectra were measured by the PL2006 Multifunction Fluorescent Spectrometer (Labguide co.)

3. Results and discussion

The crystal structure of the nanorod was examined by XRD. Fig. 1 shows an XRD pattern of the ZnO12 nanorod array. The sharp shape of the (0 0 2) diffraction peak suggests that ZnO samples are well crystallized. The remarkably enhanced (0 0 2) diffraction peak is much more intensive than the other peaks, implying that these nanorods were perfectly oriented perpendicular to the substrate surface and that they were grown along the *c*-axis.

The morphology of ZnO nanorods was characterized by FESEM. Fig. 2(a) and (b) show top-view images of the ZnO12 and ZnO14 nanorods respectively. All nanorods were grown in a direction perpendicular to the substrate. Vertically aligned arrays of ZnO nanorods were uniformly grown on the ZnO12 sample with the diameters varying from 25 to 50 nm. The diameter of the ZnO rod on the ZnO14 sample altered from 65 to 165 nm. Increasing the rod growth time widened the rod diameter. There are some closely packed rods that the inter-rod

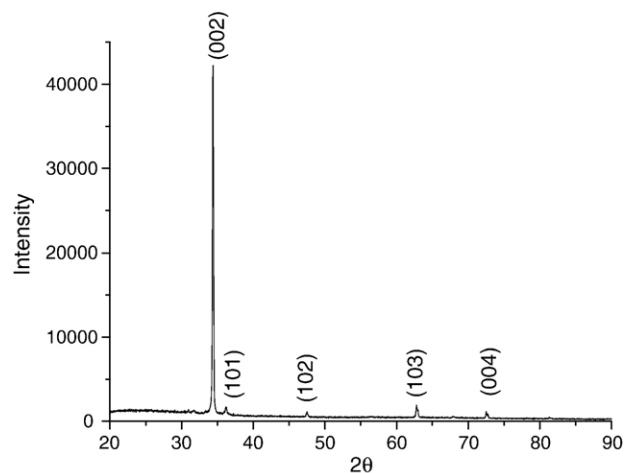


Fig. 1. XRD of ZnO12 nanorods.

distance is close to zero. Although the aspect ratio of the longest nanowires continues to increase with increasing growth time, the number density of nanorods decreases. Some nanorods have stopped growing. Similar phenomena were also observed by Miyashita et al. [10] in the ZnO nanorods prepared by the CVD method. They reported a ‘collision’ between two non-parallel ZnO nanorods during the growing process. When a nanorod collides with the side of another nanorod, the growth of the colliding nanorod is stopped there. On the other hand, the growth of a nanorod being collided on its side continued after the ‘collision’. It is the reason why the number density of nanorods decreases and why the vertical orientation of nanorods is improved after longer growth time.

The DBZ14 film with larger ZnO rod should have larger ZnO surface area than the DBZ12 film did. If the ZnO surface is completely covered by DB-PPV, there will be more donor-acceptor interface area (P–N junction) for the DBZ14 film. More incident photons are absorbed. Because of efficient dissociation of generated excitons and subsequent transport of charge carriers, photocurrent should increase with increasing surface area. To check the coverage of DB-PPV polymer on the ZnO rod, the cross-sectional images of both films were observed by FESEM. Fig. 2(c) showed the cross-sectional FESEM image of DB-PPV polymer infiltrated into the DBZ12 nanorod array structure. The polymer can be effectively intercalated into the ZnO rods film to make a nanostructured oxide/conjugated polymer composite device. The thickness of the hybrid film is about 500 nm. Fig. 2(d) showed the cross-sectional FESEM image of DBZ14 hybrid film. There were some ZnO nanorods that were not covered by the DB-PPV polymer. The closely packed rods with almost zero inter-rod distance (shown in Fig. 2(b)) hindered the infiltration of polymer into the nanorod array. The thickness of the hybrid film is about 1 μm.

For the conjugated polymers, electron transport can be viewed as a series of hops between shallow trap sites, with a fraction of the sites acting as recombination centers. The use of single crystal nanorods may allow electron transport via extended states in the conduction band, rather than by a series of hops between trap states. In this study, the ZnO rod diameter

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