Retraction Notice

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Heterojunction Photoelectrode of Polyaniline/ZnS Film/ZnO Nanorod on FTO Glass

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Abstract
To enhance the absorption of visible light for wide-band-gap semiconductors, methods such as sensitizing with nanoparticles or quantum dots, and band gap engineering using dopants have been reported. However, these can cause lattice mismatch, inherent disorders, or imperfect charge balance, which serve as recombination sites and significantly reduce the photocatalytic efficiency. Herein, photoelectrodes of PANI/ZnS/ZnO on F:SnO2 (FTO) were fabricated to analyze these issues and examine their interface microstructural and photocatalytic properties. ZnO nanorods (NRs) were grown on FTO glass by potentiostatic electrodeposition, and ZnS and PANI films were coated by liquid processes. These PANI films were applied as a visible light sensitizer and photocorrosion prevention layer for ZnS/ZnO photoelectrodes. Subsequently, the prevention effect of photocurrent loss due to the photocorrosion of semiconductors was systematically investigated. The photocurrent of the PANI/ZnS/ZnO photoelectrode measured at 0.5V under white light illumination was five times higher than that of the ZnO NR photoelectrode. This was attributed to photocorrosion prevention and visible light absorption of PANI layers due to proper energy band alignment of the hybrid heterojunction semiconductors.

Keywords
Polyaniline, Visible Light Sensitizer, Photocorrosion, Heterojunction, ZnS Shell/ZnO Core

1. Introduction
The solar energy received by the earth consists of ultraviolet (~5%) and visible light (~43%). Nanostructures are important not only for visible-light absorption in the solar spectrum, but also to obtain a high photocatalytic efficiency for semiconductor photoelectrodes. There has been extensive research on various nanostructured photoelectrodes such as nanowires, nanorods, and nanotubes...
Many multilayered heterojunction nanostructures with great penetration depth have been also reported for the absorption of long wavelengths in the visible light spectrum, as well as the efficient charge separation and transfer of excited electrons and holes [7] [8] [9] [10].

One of the methods to enhance visible-light absorption is to adhere inorganic semiconductors with narrow-band-gap materials such as nanoparticles or quantum dots (QDs) on the surface of wide-band-gap semiconductors, as a sensitizer [11] [12] [13]. Another way is to manipulate the band gap sufficiently to obtain a visible light spectrum response of 1.7 - 2.9 eV by doping impurities such as carbon, nitrogen, or hydrogen [14] [15].

Regarding visible-light sensitizers, studies on chalcogenide compound QDs such as CdS (Eg = 2.4 eV), CdSe (1.7 eV), and CdSSe (1.9 eV) have been reported [16], as well as those on plasmonic Ag and Au nanoparticles [17], transition metal dichalcogenide sheets such as MoS2 and WS2 [18], and metal halide perovskites such as CH3NH3PbI3 and CsPbBr3 [19]. However, it is important to develop alternative sensitizer materials, owing to the significant disadvantages of the abovementioned materials, such as the harmful nature of Cd, high cost of noble metals, and unstable moisture process.

In band gap engineering through doping of single or several impurities, the recombination probability of excited electrons and holes is very high near doping sites, which include deteriorative crystal defects such as lattice mismatch at the interface between the sensitizers and wide-band-gap semiconductors, unstable inherent disorders [20], and charge imbalance among dopants. Hence, even though the absorption of visible light is improved by doping, these recombination sites can cause significant losses of photocatalytic current during the separation and transfer of visible-light-excited electrons and holes.

Meanwhile, there are few reports on the photocorrosion of sensitizers such as QDs and nanoparticles attached on wide-band-gap semiconductors or those of semiconductors themselves. As alternatives, polymeric materials have recently been proposed as a prevention layer for photocorrosion. However, systematic studies on the photocatalytic performance using these films have not been reported thus far.

In this study, conductive organic polyaniline (PANI, ΔE_{LUMO-HOMO} = 2.4 eV) was applied for the core/shell composites of ZnS film/ZnO nanorods (NRs) on F:SnO2 (FTO) substrates. Subsequently, the performance of PANI as a sensitizer as well as a prevention layer to improve the photocatalytic properties, was evaluated.

### 2. Methods

In this study, to fabricate the hybrid core/shell heterojunction photoelectrodes, after ZnO NRs were grown on FTO substrates by electrodeposition, ZnS films were formed on the surface of ZnO NRs by chemical dipping of ZnO NRs into Na2S solution. Finally, the PANI films were coated by chemical dipping techniques. The detail processes are described below.
2.1. Preparation of ZnO NRs/FTO Glass Photoelectrode

ZnO NRs were grown on FTO (resistance = 8 Ω·m) glass by potentiostatic electrodeposition [21] [22]. The FTO glass was placed in an aqueous solution of 0.5 mM ZnCl₂ and 0.1 M KCl under oxygen bubbling in a three-electrode electrochemical cell, comprising a counter electrode of Pt mesh and reference electrode of Ag/AgCl/sat. KCl. Electrodeposition was carried out at −1.0 V for ~3 h at room temperature.

2.2. Particulate ZnS Film Coating on ZnO NRs

A particulate ZnS film was coated on the surface of the ZnO NRs by immersing the ZnO NR electrode in an aqueous 0.32 M Na₂S·9H₂O solution for ~12 h. Sulphurization by anion-exchange reaction was carried out in a deionized water bath at 60°C. Na₂S·9H₂O was used as an S²⁻ anion source to form ZnS on the surface of the ZnO NRs, according to the following reaction:

\[
\text{ZnO(s)} + \text{S}^{2-}_{(aq)} + \text{H}_2\text{O(l)} \rightarrow \text{ZnS(s)} + 2\text{OH}^-_{(aq)}
\]

The electrode was then washed with deionized water and absolute ethanol several times and dried at 80°C for 2 h in an oven [23] [24].

2.3. PANI Sensitizer Coating on Particulate ZnS Film/ZnO NRs

The PANI sensitizer was coated by successive ionic layer adsorption and reaction using ammonium persulfate as an oxidizing agent. First, 0.4 M aniline was dissolved in 1 M sulfuric acid, which served as a cationic precursor in the first beaker. The second beaker contained an oxidant solution of 0.2 M ammonium persulfate in 1 M sulfuric acid, which served as an anionic precursor. The electrodes were then immersed into the aniline solution for 60 s for the surface polymerization of PANI on the particulate ZnS film/ZnO NR. Subsequently, the electrodes were immersed in the ammonium persulfate solution for 30 s [25] [26].

2.4. Microstructural Characterization

The microstructures of the photoelectrodes were characterized by field-emission scanning electron microscopy (FE-SEM; JSM-6500 F, JEOL), field-emission transmission electron microscopy (FE-TEM; 200 kV/JEM-2100F HR, JEOL), and X-ray diffraction (XRD, SWXD, Rigaku). Ultraviolet-visible (UV-vis) absorbance spectra of the electrodes were obtained using an ultraviolet-visible-near infrared (UV-VIS-NIR) spectrophotometer (UV 2600, Shimadzu). XPS was performed using a Thermo VG ESCALAB 250 instrument equipped with a microfocused, monochromatic Al Kα X-ray source (1486.6 eV) and magnetic lens. The X-ray spot size was 500 µm (15 kV, 150 W). The spectra were acquired in the constant analyzer energy mode with a pass energy of 150 eV and 40 eV for overview scans and high-resolution scans, respectively.

2.5. Photoelectrochemical Characterization
The photocurrent densities of the photoelectrodes were measured using a potentiostat (AMT VERSASTAT 3, Princeton Applied Research) with a three-electrode cell comprising Pt mesh as the counter electrode and Ag/AgCl/sat. KCl as the reference electrode separated by a proton exchange membrane in aqueous 0.5 M Na$_2$SO$_4$ (pH = 7.15) electrolyte. Using a 1 kW xenon lamp (Newport) with the infrared wavelengths filtered out by water, 1 cm$^2$ of the working electrode was exposed. The light irradiance, measured using a thermopile detector, was 100 mW/cm$^2$.

3. Results and Discussion

3.1. Microstructural Characterization

In this study, photocatalytic properties of ZnS film/ZnO NR heterojunctions with PANI layers were investigated. For heterojunction composite fabrication, after the ZnO NRs were grown on FTO substrates by electropinning, ZnS layers were formed via sulfurization followed by PANI coating via cationic surface polymerization. The cross-sectional and surface FE-SEM images of ZnO NRs (Figure 1) show that the hexagonal ZnO NRs, with diameters of 300 - 500 nm and lengths of 0.5 - 1.5 μm, were grown vertically.

![Figure 1](image_url)

**Figure 1.** FE-SEM images of coated PANI on sulfurized ZnO NRs core. Cross sectional and surface views for ZnO NRs/FTO (a) & (b), particulate ZnS film/ZnO NRs/FTO (c) & (d), and PANI/particulate ZnS film/ZnO NRs/FTO (e) & (f), respectively.

The top surface of the prepared ZnO NRs was dominantly composed of slanted side faces of \{10\bar{1}\} crystal planes rather than flat hexagonal \{0001\} crystal planes. Compared to the hexagonal \{0001\} planes, \{10\bar{1}\} crystal planes have lower stacking density, and thus, higher surface energy. The hydrothermal growth
rate of ZnO NRs on FTO is determined primarily by the diffusion rate of Zn\(^{2+}\) ions during the process, and ion diffusion into side \{10\(\bar{1}\)\} planes is much more than that into top \{0001\} surfaces, such that the hydrothermally grown ZnO NRs show wide \{0001\} crystal planes on their top surfaces. On the other hand, electroplated ZnO NRs have narrow and sharp top surfaces composed of hexagonal \{0001\} planes, because Zn\(^{2+}\) concentration is much higher in the top surfaces due to higher electroplating potential. However, many voids are observed on the surface of ZnO NRs (Figure 1), probably formed by O\(_2\) gas bubbles in the electroplating bath, which disturbed the migration of Zn\(^{2+}\) ions on the surface.

Meanwhile, for fabrication of heterojunction composites with efficient charge separation, the ZnS films were coated on the surface of ZnO NRs via ion exchange sulfurization. When the ZnO NRs were dipped in the Na\(_2\)S solution, the O\(^{2-}\) ions on the ZnO surface and the S\(^{2-}\) ions in the Na\(_2\)S solution were exchanged to form ZnS particles on the surface of ZnO NRs. The cross-section and surface images of ZnS films/ZnO NRs fabricated by these processes are shown in Figure 1(c) and Figure 1(d). As shown in Figure 1(d), the ZnS films consist of stacked aggregates of \(~25\) nm nanoparticles. The cross-section and top surface images of the conductive organic sensitizer, PANI, coated on the ZnS film/ZnO NRs are shown in Figure 1(e) and Figure 1(f), respectively. As shown in Figure 1(f), the smooth surface of a ZnO NR is surrounded by both ZnS film composed of stacked agglomerated nanoparticles and PANI film.

TEM images of PANI/ZnS/ZnO NRs (Figure 2(a) and Figure 2(b)) show that a core electrodeposited ZnO NR is surrounded by both \(~5\)-nm-thick PANI film and \(~30\)-nm-thick particulate ZnS film. Thus, this core/shell structure should prevent direct contact of ZnO NRs with the electrolyte in the photocatalytic cells, thus preventing deterioration of photocatalytic current due to photocorrosion.

ZnS films on ZnO NRs can be typically formed by several methods like chemical absorption of ZnS nanoparticles and vapor-liquid-solid (VLS) methods.
Zn and S vapors on the surfaces of ZnO NRs, which have different surface states depending on crystal orientations, results in anisotropic growth of ZnS. Thus, it is difficult to obtain dense ZnS films using this technique, but thick films can be obtained. However, in the surface sulfurization processes, uniform ZnS films can be feasibly formed, because ZnS nanoparticles are formed by ion exchange of O\textsuperscript{2−} ions on the ZnO surface and S\textsuperscript{2−} ions in the solution. Furthermore, the crystal orientations of ZnO NRs have negligible effect on the growth of ZnS because the ZnO surface is surrounded by water molecules in the solution.

The PANI film synthesized by polymerization was deposited relatively uniformly on the ZnS film. Selected area electron diffraction patterns for the interface between particulate ZnS film and ZnO NRs are shown in Figure 2(c). Polycrystalline ring patterns and single-crystalline dot patterns were observed for the particulate ZnS film and the ZnO NR, respectively. Electron diffraction patterns from (100), (101), and (102) ZnO (zincite space group P6\textsuperscript{3}mc; JCPDS card no. 36-1451) for the [0\textit{0}0\textit{0}] zone axis were observed, confirming the hexagonal Wurtzite structure of ZnO NRs.

However, the electron diffraction patterns of the ZnS film corresponded to both the (111) sphalerite cubic phase (space group F43m; JCPDS card no. 05-0566) and the hexagonal (008) hexagonal phase (space group P6\textsuperscript{3}mc; JCPDS card no. 39-1363), which have interplanar distances of 3.123 Å and 3.120 Å, respectively. However, other groups using similar ZnS synthesis processes reported cubic zinc blende (111) peaks at 2\textdegree = 28.62˚ in XRD patterns.

The XRD patterns for PANI/particulate ZnS film/ZnO NRs/FTO samples (Figure 3) indicate that the peaks (2\textdegree = 31.8˚, 34.5˚, 36.3˚, 47.6˚, 56.5˚, and 62.9˚) of ZnO NRs correspond to the (100), (002), (101), (102), (110), and (103) crystal planes, respectively, of the hexagonal crystal lattice of zincite (space group P6\textsuperscript{3}mc; JCPDS card no. 36-1451). The peak at 2\textdegree = 28.61˚ corresponds to the cubic ZnS (111) crystal plane. However, the XRD peaks of PANI layer were not observed. This resulted from the amorphous phase of the PANI layer after the synthesis and dip-coating, whereas other group has reported for the polycrystalline PANI synthesis. This discrepancy was attributed to different processes including sources for the PANI synthesis [27].

The PANI layer and compositional surface analysis of the ZnS film/ZnO NRs were characterized by using X-ray photoelectron spectroscopy (XPS), and the analysis results are shown in Figure 4. Zn 2p\textsubscript{1/2} (1.045 eV), Zn 2p\textsubscript{3/2} (1.020 eV) and O 1s (531 eV) peaks were observed for ZnO NRs, and S 2s (226 eV) and S 2p
3.2. Photoelectrochemical Characterization

The UV-Vis absorption spectra of PANI/ZnS film/ZnO NRs, ZnS film/ZnO NRs, and ZnO NR electrodes were measured as shown in Figure 5(a). All the spectra present a strong absorption in the UV region between 300 and 400 nm. The band gaps of each layers were estimated from the Tauc plots [29] as shown in Figure 5(b). The values from the line intersections were measured as 3.12, 3.16, and 2.7 eV for ZnO NRs, ZnS film/ZnO NRs, and PANI/ZnS film/ZnO
Figure 5. (a) UV-Vis absorption spectra and (b) Tauc plot for band gap calculation of ZnO NRs, ZnS film/ZnO NRs, and PANI/ZnS film/ZnO NRs electrode, respectively.

The ZnO NRs coated with the ZnS film of the wide band gap showed slightly increased band gap, that is blue-shifting absorption. However, the wide and weak absorption in the visible region between 400 and 500 nm was observed for the PANI coated ZnS film/ZnO NRs sample. This resulted in the reduced band gap of 2.7 eV in the Tauc plot. Thus this is due to the narrow band gap of the PANI layer in the range from ~2.4 eV to ~2.8 eV [30].

Photocurrent densities of the samples were measured in a photoelectrochemical (PEC) cell under exposure of white light consisting of UV and visible light (see Figure 6(a) ZnO, Figure 6(b) ZnS/ZnO, and Figure 6(c) PANI/ZnS/ZnO, respectively). Photocurrent density corresponding to the pure photoresponse by using a chopping light source (on-off) was measured. The measured photocurrent density was the photoresponse difference between the on- and off-states of the light source.

Under white light illumination, the representative photocurrent densities of ZnO NRs, ZnS film/ZnO NRs, and PANI/ZnS film/ZnO NRs were 0.173 mA/cm², 0.461 mA/cm², and 0.865 mA/cm², respectively, at 0.5 V. The PANI/ZnS film/ZnO NR electrode showed about 5- and 1.8-times larger photocurrent density than the ZnO NRs and ZnS film/ZnO NRs, respectively. The ZnS film/ZnO NR electrode showed about 2.7-times larger photocurrent density than the ZnO NRs.

These measurement results confirm the light absorption spectral analysis results (Figure 5). The photocurrent densities of PANI/ZnS film/ZnO NRs were significantly improved due to superior visible light absorption properties of the
Figure 6. Photocurrent density of (a) ZnO NRs, (b) ZnS film/ZnO NRs, and (c) PANI/ZnS film/ZnO NRs electrodes under white light (UV and visible light) irradiation.

PANI layers. The PANI band gap was 2.7 eV as shown in Figure 5(b). Furthermore, both PANI/ZnS film/ZnO NRs and ZnS film/ZnO heterojunction have good enough combination of potential levels (energy band alignment) to transfer and efficiently separate excited electrons and holes [31] [32] [33] [34].

In summary, PANI/ZnS film/ZnO NR photoelectrodes were fabricated and evaluated for their photocatalytic performance. Single-crystalline hexagonal ZnO NRs of lengths 0.5 - 1.5 μm were grown on FTO glass. Cubic ZnS films consisting of 5 - 10-nm nanoparticles were coated with thickness of ~30 nm on the ZnO NRs and organic PANI layers with thickness of ~5 nm were coated on the ZnS films. XRD and XPS analyses showed strong peaks for ZnS (111) and ZnO (002) crystal planes, and C/N elements and molecular bonding structure were observed for PANI.

4. Conclusion

In this study, PANI/ZnS/ZnO/FTO electrodes were studied on their interface and photocatalysis. Under white light illumination, the PANI/ZnS film/ZnO NR electrode showed five times higher photocurrent density than ZnO NRs. This was due to the anti-photocorrosion and visible light absorption of PANI layer, and also provided proper energy band alignment of the hybrid heterojunction (PANI/ZnS film/ZnO NRs) to efficiently transfer and separate excited electrons and holes. It is expected that this hybrid heterojunction photoelectrode with proper band alignment using organic materials could be utilized for other applications such as photovoltaics, solar CO₂ fuel conversion, and solar pollutant decomposition.

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Conflicts of Interest

The authors declare no conflicts of interest regarding the publication of this paper.

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