Self-Powered Ultrasensitive Nanowire Photodetector Driven by a Hybridized Microbial Fuel Cell**

Qing Yang, Ying Liu, Zetang Li, Zongyin Yang, Xue Wang, and Zhong Lin Wang*

Nanostructures are promising materials for photodetection, with advantages of high sensitivity, fast response speed, and low energy consumption.[3] However, until now, highly sensitive nanostructure-based photodetectors have been designed to be driven by external power sources, which not only largely increase the system size but also greatly limit their mobility and independence for applications in areas such as large-area wireless environmental sensing, chemical and biosensing, and in situ medical therapy monitoring. One of the goals of nanotechnology is to build a nanosystem that can function as a living species, with capabilities of sensing, controlling, communicating, and actuating/responding.[2] Such a nanosystem is composed of not only nanodevices but also a power source. For instance, implanted nanoscale biosensors for monitoring diabetes and blood pressure require a nontoxic power source without adding too much weight and size. Although a battery or energy storage unit may be a choice for powering nanodevices, harvesting energy from the environment is an essential solution for building a self-powered nanodevice/nanosystem, which is indispensable for sustainable operation of a large portion of the sensor network.[2] For example, a self-powered nanoscale photodetector network is highly desired for waste-water and air-pollution monitoring systems that feature low energy consumption, low cost, and high sensitivity.[3] Recently, a self-powered UV detector driven by a pn heterojunction photocell was demonstrated,[4] but the limitation of the detectable light intensity is very high (ca. 200 W cm\(^{-2}\)). However, for environmental monitoring and medical therapy treatment, weak light detection is much more desired. Herein we demonstrate an integrated system of a self-powered ultrasensitive single-nanowire-based multicolor photodetector driven by a high-efficiency microscale microbial fuel cell (MFC) that was fabricated using a carbon fiber-ZnO nanowire hybridized structure. One microscale MFC can power an ultrasensitive nanowire photodetector (NWPD) with a responsivity of more than 300 A W\(^{-1}\) and a detection limit as low as nW cm\(^{-2}\). The noise-equivalent power (NEP) of the self-powered NWPD for detection of UV, blue, and green light at 10 Hz is about \(5.0 \times 10^{-14}, 1.5 \times 10^{-13}\), and \(2.8 \times 10^{-17}\) WH z\(^{-1}\), respectively, indicating extremely high sensitivity of the self-powered nanosystem, which is compatible with those of quantum dots and polymer-nanowire-based photodetectors. Thus, it opens exciting opportunities for self-powered detectors with a wide spectral range of high responsivity and high resolution. Our self-powered nanosystem may find a variety of applications in medical therapy, environmental monitoring, defense technology, and personal electronics; for example, light dosemeter measurement in photodynamic therapy and contamination detection in polluted water. Furthermore, parallel or series connection of a hybridized fuel cell can be adopted to improve the output similar to the integration of nanogenerator, as carried out previously.[3]

Our self-powered nanosystem is fabricated by integrating a single-fiber NW hybrid-structured MFC with a single CdS NWPD in series (Figure 1a; Supporting Information, Figure S1). The MFC uses a carbon fiber that is radially surrounded with densely packed ZnO NWs (Figure 1b). The principle of the MFC is shown in Figure 2a. The carbon fiber (5 μm in diameter) acts not only as the substrate onto which the ZnO NWs were grown but also as an electrode (indicated as electrode EII in Figure 1a).[6] Densely packed ZnO NWs with lengths of about 250 nm were grown around the carbon fiber by a physical vapor deposition method.[9] A thin layer of Nafion was spin-coated on the fiber-NW hybrid structure as the anion exchange membrane in the MFC. Then a millimeter-diameter solution micropool was constructed by squeezing poly(methyl methacrylate) (PMMA) polymer solution onto the Nafion film in a ring dam shape to form a micropool (the inner diameter ranges from 1 to 5 mm, the thickness is about 1 mm, and the volume is less than 20 μL). The micropool not only keeps the yeast solution near the surface but also as an electrode (indicated as electrode EII in Figure 1a).[6] Densely packed ZnO NWs with lengths of about 250 nm were grown around the carbon fiber by a physical vapor deposition method.[9] A thin layer of Nafion was spin-coated on the fiber-NW hybrid structure as the anion exchange membrane in the MFC. Then a millimeter-diameter solution micropool was constructed by squeezing poly(methyl methacrylate) (PMMA) polymer solution onto the Nafion film in a ring dam shape to form a micropool (the inner diameter ranges from 1 to 5 mm, the thickness is about 1 mm, and the volume is less than 20 μL). The micropool not only keeps the yeast solution near the hybrid structure, but also improves the stability of the MFC. Yeast and glucose were mixed in solution and dripped into the micropool. The MFC was connected to the outer circuit by two electrodes. The first electrode (E1) was placed on the top surface of the yeast solution, the other electrode (EII) was connected to one end of the carbon fiber where ZnO NWs

** These authors contributed equally to this work.
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The working principle of a MFC is shown in Figure 2a. Glucose was decomposed by yeast to form CO₂ and electron–proton pairs. The electrons produced in the microorganism solution are transferred to the anode and subsequently through an external circuit to the cathode, whilst protons or phosphate anions migrate from anode to the cathode through the separator membrane. At the cathode, H⁺ and electron recombine with an oxygen molecule to form H₂O. A separator is an important part to ensure an efficient and sustainable operation of an MFC. The paradox of ion transfer and oxygen permeation is the major constraint for most separators in MFCs.⁶ Our MFC however is based on a hybridized structure of carbon fiber–ZnO NWs, which has increased ion transfer and avoids simultaneous oxygen permeation. As shown in Figure 2a, a porous membrane rather than a smooth film will form on the ZnO NW-coated carbon fiber when spin coating Nafion film, which allows various charged or neutral species to pass through efficiently and leads to easier ion transfer.⁷ On the other hand, oxygen is adsorbed and immobilized on the ZnO NW surface, as represented by blue spheres on the ZnO nanowire surfaces. Thus, the oxygen permeation through the membrane is reduced. These two advantages may allow the hybrid-structured MFC to give a higher output.

The performance of the MFC is characterized by measuring the short-circuit current I_{sc} and open-circuit voltage V_{oc}. Once a drop of yeast solution was dripped into the micropool, the MFC started to generate a DC output. A typical output of the MFC is shown in Figure 2c and Figure S3; the I_{sc} is about 56 nA, the V_{oc} is about 295 mV, and the corresponding power density is about 30 W m⁻² of electrode geometry area, while the performance reported for µL- and sub-µL-scale MFC is limited by low volumetric power density owing to its high internal resistance.⁸ The output power density of our MFC is among the best of its kind reported to date.⁹

We found that the ZnO NWs around the carbon fiber may play an important role for the enhanced performance of our microsize MFC. For comparison purposes, a MFC was fabricated using a bare carbon fiber without ZnO NWs following the same procedures as for the hybrid structured MFC. The corresponding I_{sc} was about 2 nA, and the V_{oc} was about 80 mV, and the corresponding power density was 0.29 W m⁻² of the electrode geometry area (Figure 2c), which are two orders of magnitude lower than that of the hybrid structured MFC. Thus, the significant enhancement in performance of our hybrid structured MFC may originate from the increased surface area and the absorption of oxygen at ZnO NW surfaces.

By integrating a MFC with a CdS NWPD,⁶⁻⁹ a self-powered photodetector can be achieved. The output voltage of the MFC was about 295 mV, which was high enough to drive a NWPD (Figure 3; Supporting Information, Figure S4). The self-powered nanosystem shows an excellent response to solar light, with a response time of about 30 ms and decaying time of about 40 ms (Supporting Information, Figure S4a,b). The photocurrent arises from the photoconductivity of the CdS NW and the shrinkage of the depletion region of the Schottky contact.¹⁰ The fast response time...
results from the Schottky contact\cite{1} and high crystallinity of the CdS NW.\cite{11} A Schottky contacted device has been proven to have a much higher sensitivity and much faster responsivity than an ohmic contacted device for a gas, chemical, and biosensor.\cite{10} Regarding the intensity dependence of measured current and derived responsivity for solar detection (Supporting Information, Figure S4c,d), the responsivity decreases at high light intensity owing to the elimination of the Schottky barrier and hole-trapping saturation at strong light intensities.\cite{1e,g}

Apart from solar light detection, our self-powered photodetector can be used to detect multicolor light ranging from red light to UV light. For the integration system of the hybrid structured MFC and the CdS NW connected in series (Figure 1a), if light irradiation on the CdS NW is turned off, the system current had a leakage current less than 1 fA. Once the UV light with an intensity of about 1.6 × 10^{-4} W cm^{-2} shed on the CdS NW, the system current increased immediately to 1.4 nA (Figure 3a). The on–off cycles have remarkable reproducibility.

Our self-powered nanosystem can detect light down to a nW/cm^{2} level (fW light illuminating onto the NW). The responsivity for UV, blue, and green light with intensity of 2.3 nW cm^{-2} (equal to 25 fW light illumination onto the NW) of the system were 1180, 344, and 332 AW^{-1}, respectively, which is two to three orders of magnitude higher than that received using a nitride based metal–semiconductor–metal photodetector\cite{12,13} and a silicon NW photodetector.\cite{13} Figure 3d shows the photocurrent versus light intensity for UV, blue, and green light, respectively, indicating that the photocurrent increased linearly with the optical power for sub-μW/cm^{2} light detection. The spectral photoresponse of the self-powered nanosystem (Supporting Information, Figure S5b) showed that the responsivity depended on light wavelength. The responsivity for UV, blue, and green light was much higher than that for red light, as the photon energy of red light is less than the band gap of CdS, and thus the generation of electron–hole pairs was less efficient.

The figure of merit of a photodetector is the NEP, that is, the minimum radiant power that a detector can distinguish from noise. It is difficult to measure the signal when the signal-to-noise ratio is unity. Therefore, it is customary to make the measurement at a high level of incident radiation and calculate the NEP from the following equation:\cite{1c,14}

\[
\text{NEP} = \frac{I_N}{R \sqrt{\Delta f}}
\]

where \(I_N\) is root-mean-square value of the noise current at output of the photodetector, \(R\) is responsivity of the photodetector, and \(\Delta f\) is the electrical bandwidth in Hz. Frequency-dependent responsivity and the noise-equivalent power of the self-powered nanosystem are shown in Figure 4a and 4b, respectively. Surprisingly, we found that the noise current of the NWPD driven by a hybridized MFC is about one order of magnitude lower than that driven by an external voltage provided by a function generator, especially at frequencies that are overtones or combination bands of 60 Hz (Figure 4b, inset image), which is due to the small size, good stability of the microscale MFC, and isolated properties of the nano-

![Figure 3](image1.png)

**Figure 3.** UV–blue–green multicolor detection by a single CdS NWPD driven by a single hybrid-structured MFC. a) UV light (372 nm); b) blue light (486 nm); c) green light (548 nm). Light intensities (light power illuminated onto the CdS wire): black 2.3 × 10^{-4} (25 fW), red 9.1 × 10^{-4} (100 fW), blue 3.6 × 10^{-3} (400 fW), and green 1.6 × 10^{-2} W cm^{-2} (1.7 nW). d) Photocurrent as a function of the excitation intensity on the CdS NW that was illuminated by UV, blue, and green light (curves correspond to the color of the light).

![Figure 4](image2.png)

**Figure 4.** Frequency-dependent performance of the self-powered multicolor NWPD. a) Responsivity versus modulation frequency \(F\) under 2.3 × 10^{-3} W cm^{-2} UV, blue, and green light illumination. Lines: data fitting. b) Noise-equivalent power (NEP) versus modulation frequency \(F\) of the self-powered NWPD for different color light detection. Inset: noise current \(I_N\) of the NWPD driven by a hybridized MFC (red) and by a 0.3 V external voltage source (black) provided through a function generator at a bandwidth of 90 Hz (from 10 to 100 Hz).
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Ein integriertes System aus einem Mehrfarben-Hybridphotodetektor (Kohlenstofffaser und ZnO-Nanodraht (NW)) wird durch eine mikrobielle Brennstoffzelle angetrieben (siehe Bild; PMMA = Polymethylmethacrylat, E = Elektrode).

Dieser Photodetektor ohne externe Stromquelle kann Licht selbst bei einer Intensität im nW cm⁻²-Bereich mit einer Empfindlichkeit von mehr als 300 AW⁻¹ detektieren.