InGaN/Cu$_2$O Heterostructure Core-Shell Nanowire Photoanode for Efficient Solar Water Splitting

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INTRODUCTION

The heterostructuring and doping concepts have proved to obtain a novel n-InGaN/p-Cu$_2$O nanowire (NW) photoanode by strong enhancement of the photocurrent compared to a bare InGaN NW photoanode in solar water splitting. The large photocurrent is due to the maximized photocarrier separation and hole transfer to the surface in the depletion zone of the p–n heterojunction established by the p-Cu$_2$O layer, forming a thin, uniform shell-layer around the n-InGaN NW core by electrodeposition. For sufficiently thin Cu$_2$O layers, the upward energy band bending in the depletion zone extends up to the surface for optimized hole transport and surface reaction. Thick Cu$_2$O layers on top of the InGaN NWs act as common photocathodes. The functional InGaN/Cu$_2$O heterostructure core-shell NW photoanode is chemically self-stabilized at positive applied voltage by a thin CuO surface layer. Final deposition of the earth-abundant NiOOH co-catalyst boosts the photocurrent of the InGaN/Cu$_2$O/NiOOH complete NW photoanode into the competitive mA/cm$^2$ range.

Keywords: Cu$_2$O, InGaN nanowires, core-shell, stability, co-catalyst

INTRODUCTION

The key strategies to boost the performance of photoelectrodes are nanostructuring, surface catalyst coupling, light management, heterostructuring, and doping to optimize the crucial processes of light absorption, photocarrier separation and transfer, and surface reaction [1, 2]. To begin with the right material, InGaN is one of the best choices due to the wide tunability of the direct optical bandgap over the whole visible range by varying the In content, the high absorption coefficient, high carrier mobility, and chemical and mechanical resilience [3–5]. In addition, for one of the most important reactions, the photoelectrochemical splitting of water into hydrogen and oxygen [6–8], InGaN possesses the required straddling alignment of the conduction band and valence band with the water H$^+$/H$_2$ reduction potential and O$_2$/H$_2$O oxidation potential for up to $\sim$50% of the In content and shows the best bandgap energy for solar light absorption for 30–40% In [9]. Solar hydrogen produced by this method of photoelectrochemical water splitting is the ideal green fuel to replace fossil fuels in a sustainable hydrogen energy economy$^1$.

Nanostructured n-type InGaN nanowires (NWs) and InGaN layers, activated with InN quantum dots (QDs), employed as photoanodes, showed high efficiencies in the oxygen evolution reaction (OER) in solar water splitting [10–16]. The underlying difference between these two approaches is

$^1$Hydrogen program. Available online at: https://www.hydrogen.energy.gov.
the type of surface states for the m-plane InGaN NW sidewalls and the c-plane surface of the InGaN layers and InN QDs. m-Planes exhibit negatively charged surface states within the energy bandgap, causing near-surface upward energy band bending, which is most commonly encountered in the n-type semiconductors. The c-plane of In-rich InGaN layers exhibits positively charged surface states located within the conduction band [17–21]. These positively charged surface states catalyze the OER by attraction of electrons, which is strongly enhanced by the zero-dimensional electronic properties of InN QDs. For m-planes, in contrast, a low catalytic activity for the OER results from the negatively charged surface states. Therefore, to benefit from the largely increased surface area of NWs, additional measures need to be taken to obtain competitive water splitting efficiencies, such as the decoration of the NW sidewalls with co-catalysts. InGaN NWs have also demonstrated their successful light management and absorption enhancement, acting as efficient anti-reflection structures when grown on pyramid-patterned Si substrates [22].

In this study, we demonstrate the heterostructuring and doping routes for an InGaN/Cu$_2$O core-shell NW photoanode to boost solar water splitting by enhancing the separation and transport of photogenerated carriers. The underlying concept is as follows: A heterostructure with the right staggered band alignment, such that no energy barriers for the transport of electrons and holes are present, can improve the photogenerated carrier separation and transport. However, it is only effective together with the right band bending given by the doping. Based on an n-type photoanode with near-surface upward energy band bending, a p-type layer on top is optimum to increase the upward energy band bending at the p–n junction. Together with a positive band offset, this pushes the holes toward the surface more effectively for the OER and also enhances the electrons to travel toward the cathode for the hydrogen evolution reaction (HER). A p-type layer, however, exhibits the wrong near-surface downward energy band bending, opposing the hole transport toward the surface. Therefore, the p-type layer has to be sufficiently thin, so that it is fully depleted and the upward energy band bending proceeds all the way to the surface. This turns a p-type layer, usually acting as photocathode into an active surface layer, by boosting the efficiency of an n-type photoanode. For the optimized bandgap energy for solar light absorption of the n-type photoanode, the p-type layer on top ideally has a similar, medium bandgap energy to contribute to the absorption. Regarding the OER, however, the surface of the p-type layer is not necessarily reactive, still requiring the deposition of co-catalysts.

Considering the band alignment, doping, and bandgap energy, p-type Cu$_2$O fulfills all the requirements when combined with n-type In-rich InGaN as a heterostructure photoanode. Photogenerated carrier separation and transfer are strongly enhanced in the depletion zone of the p–n heterojunction formed at the n-InGaN/p-Cu$_2$O heterointerface with positive band offset. This is based on a previous study where we showed that Cu$_2$O, a deeply studied p-type metal-oxide, acts as an efficient hole injection and a collection layer on In-rich InGaN NWs, solving the p-type conductivity problem [23]. High-quality p-type InGaN with a high In content is hardly realized. The bandgap energy of Cu$_2$O of 2.1 eV is sufficiently small and the conduction band and valence band of Cu$_2$O are well-aligned with those of the In-rich InGaN with the right band offsets.

Cu$_2$O is electrodeposited on the InGaN NWs. Scanning electron microscopy (SEM), transmission electron microscopy (TEM), and energy dispersive X-ray spectroscopy (EDS) reveal that Cu$_2$O forms a uniform shell-layer on the m-plane InGaN NW sidewalls while nucleating as isolated Cu$_2$O nanocrystals on the c-plane NW tops. p-Type Cu$_2$O is well-known as a photocathode for the HER [24, 25]. This is reproduced for thick Cu$_2$O layers grown on top of the InGaN NWs. Similar to InGaN, Cu$_2$O also exhibits a straddling energy band alignment with the water redox potentials [25–28], making the energy band alignment with InGaN most obvious. For thin Cu$_2$O layer deposition, the Cu$_2$O shell-layers on the InGaN NWs are fully depleted for thicknesses of ~10 nm with upward energy band bending up to the surface. The efficient photocarrier separation and hole transport to the surface strongly enhance the OER reaction for positive applied voltage, as shown by linear sweep voltammetry (LSV) measurements and current density vs. time (I–t) measurements under chopped illumination. For optimized Cu$_2$O deposition amount, an almost 7-fold enhancement of the photocurrent is observed. Electrochemical impedance spectroscopy (EIS), however, shows that the Cu$_2$O surface is catalytically not very active for the OER, such as the m-plane InGaN NW sidewalls, leading to an overall insufficient photocurrent. Therefore, to achieve competitive photocurrents, NiOOH co-catalyst is electrodeposited to complete the photoanode preparation process. NiOOH is chosen because it contains only earth-abundant elements. Functioning as InGaN/Cu$_2$O core-shell NW photoanode at positive applied voltage, the Cu$_2$O layer is chemically self-stabilized by the formation of a thin CuO surface layer due to the self-limiting oxidation of Cu$_2$O, as evidenced by the LSV measurements, cyclic voltammetry (CV) measurements, and X-ray photoelectron spectroscopy (XPS). Although imposing a small energy barrier for the hole transport, the thin CuO surface layer does not significantly influence the photocurrent. In contrast to Cu$_2$O employed as photocathode at negative applied voltage, no other surface passivation is needed. The InGaN/Cu$_2$O heterostructure core-shell NW photoanode, self-passivated with the thin CuO surface layer, shows no degradation for many hours of operation.

**EXPERIMENTAL DETAILS**

**Growth of In-rich InGaN NWs**

The InGaN NWs on planar p-type Si (111) substrates were fabricated by plasma-assisted molecular beam epitaxy (PA-MBE). The In and Ga metal fluxes were provided by standard Knudson effusion cells, and the active nitrogen species were generated with a radio frequency (RF) plasma source. Before loading into the MBE chamber, the Si substrates were etched in 10 wt% HF aqueous solution for 1 min to remove the native surface oxide layer. Then, the cleaned Si substrates were loaded into the MBE buffer chamber and degassed in the MBE middle chamber for 1 h at 300°C. The Si substrates were then transferred to the MBE...
growth chamber, outgassed for 10 min at 900°C (thermocouple reading) to remove any residual native surface oxide, and exposed to active N flux for 5 min to form a thin SiN\textsubscript{x} layer. The RF power for the Si surface nitridation was 350 W, and the molecular N\textsubscript{2} flow rate was 1.7 standard cubic centimeters per minute (sccm). The SiN\textsubscript{x} layer improves the InGaN crystal quality by enhancing strain relaxation and reducing the dislocation density. For not too extended nitridation, the SiN\textsubscript{x} interlayer does not alter the electronic properties/near-ohmic behavior of the Si/InGaN interface [29]. For InGaN growth, the substrate temperature was ramped down to 610°C (thermocouple reading) to remove any residual native surface oxide, and exposed for strain relaxation and reducing the dislocation density. For not too extended nitridation, the SiN\textsubscript{x} interlayer does not alter the electronic properties/near-ohmic behavior of the Si/InGaN interface [29].

Electrode Preparation
The samples were coated with In–Ga eutectic on the back side of the substrate to form an ohmic contact, glued on Cu-conductive adhesive tape, and fixed on a supporting glass plate. The surface was covered with non-transparent silicone rubber, leaving an opening for contact with the electrolyte.

Electrodeposition of Cu\textsubscript{2}O
The Cu\textsubscript{2}O layers were fabricated on the InGaN NWs by electrodeposition in a three-electrode electrochemical cell using an electrochemical work station. The InGaN NWs, a Pt mesh, and a saturated calomel electrode (SCE) were used as working, counter, and reference electrodes, respectively. The aqueous electrolyte comprised 0.4 M CuSO\textsubscript{4}, 3 M lactic acid, and C\textsubscript{3}H\textsubscript{6}O\textsubscript{3}. The pH was adjusted to 12 by adding 5 M NaOH. The deposition voltage was kept at −0.4 V vs. SCE. During deposition with magnetic stirring at 180 rpm, the temperature was set to 27°C by a hot plate with a temperature sensor in the solution.

Electrodeposition of NiOOH
The InGaN/Cu\textsubscript{2}O/NiOOH structures were prepared by electrodeposition in a three-electrode electrochemical cell with the InGaN/Cu\textsubscript{2}O working electrode, a Pt mesh counter electrode, and a saturated KCl Ag/AgCl reference electrode, connected to an electrochemical workstation. The electrolyte was 100 mL of 5 mM Ni(NO\textsubscript{3})\textsubscript{2}·6H\textsubscript{2}O aqueous solution. The potential was kept at −1.0 V vs. Ag/AgCl. The optimized electrodeposition time was 330 s.

Materials Assessment
Structural analysis of the InGaN NWs and Cu\textsubscript{2}O layer was performed using X-ray diffraction (XRD) (PANalytical X’Pert PRO). Field-emission SEM (ZEISS Gemini 500), TEM (JEM-2100HR), and atomic force microscopy (AFM, Bruker MultiMode 8) were used to characterize the surface morphology and cross-section of the structures. EDS element mappings were carried out with energy disperse spectrometers (Aztec X-Max 80) attached to the SEM and TEM. XPS was conducted with XPS spectrometers (EscaLab 250Xi) attached to the SEM and TEM. The XPS spectra were calibrated with respect to the C 1s peak. Raman spectroscopy was performed with the 532 nm line of a Nd:YAG solid-state laser. The diameter of the laser spot was ~1 μm. Photoluminescence (PL) spectra were taken at room temperature with the 100 mW, 532 nm line of a Nd:YAG solid-state laser as excitation source, and a silicon charged coupled device (CCD) attached to a single monochromator for detection.

Electrochemical Measurements
The electrochemical measurements were performed in a three-electrode electrochemical cell with the InGaN, InGaN/Cu\textsubscript{2}O, and InGaN/Cu\textsubscript{2}O/NiOOH working electrodes; a Pt mesh counter electrode; and a saturated KCl Ag/AgCl reference electrode connected to an electrochemical workstation. A scheme of the experimental setup and sample structure is shown in Supplementary Figure 1. The electrolyte was 0.5 M Na\textsubscript{2}SO\textsubscript{4} (pH = 7) aqueous solution. LSV current vs. voltage (I–V) measurements from +1 V to 0 V vs. Ag/AgCl [+1.6 V to +0.6 V vs. the reversible hydrogen electrode (RHE)] with 10 mV s\textsuperscript{−1} scan rate and I-t measurements at 0.6 V vs. Ag/AgCl (1.2 V vs. RHE) were performed under chopped 5 s on and 5 s off simulated sun light (1-sun, AM 1.5, 100 mW cm\textsuperscript{−2}, GLORIA-X500A). EIS measurements were conducted in the frequency range of 10 Hz to 1,000 kHz in the dark and light. CV measurements were performed in the voltage windows of −1 V to +1 V and 0 V to +1 V vs. Ag/AgCl with a 100 mV s\textsuperscript{−1} scan rate.

RESULTS AND DISCUSSION
Structure
Figures 1a,b shows the SEM top-view and cross-sectional images of the as-grown, bare InGaN NWs on a Si (111) substrate. The NWs have an average diameter of 40 nm, 600 nm length, and a flat top. After 40 s of Cu\textsubscript{2}O electrodeposition, which turns out to be the optimum Cu\textsubscript{2}O electrodeposition time, sparse, isolated Cu\textsubscript{2}O nanocrystals are visible in the top-view SEM image in Figure 1b. AFM images are shown in Supplementary Figure 2. The nanocrystals are positioned on the NWs and also in-between NWs with arbitrary orientations. The shape of the nanocrystals reflects the cubic crystal symmetry of Cu\textsubscript{2}O. The more connected appearance of the NWs in the cross-sectional image in Figure 1c indicates the deposition of Cu\textsubscript{2}O on the InGaN NW sidewalls. Formation of the pure Cu\textsubscript{2}O phase for the present electrodeposition condition is confirmed by the omega-two theta XRD spectrum shown in Supplementary Figure 3A and by the Raman spectrum in Supplementary Figure 3B [23]. Only peaks from cubic Cu\textsubscript{2}O, InGaN, and Si, as labeled, are observed. From the XRD spectrum, the in content of the InGaN NWs of 28% is deduced [30]. However, XPS measurements (Figure 1e) that were taken some days after fabrication of the Cu\textsubscript{2}O layer revealed the presence of CuO. As XRD and Raman spectroscopy rather probe the bulk material, XPS is highly surface sensitive, indicating the self-limiting oxidation of Cu\textsubscript{2}O to CuO in air to form a thin surface layer of CuO [31]. This is
Zhao et al. InGaN/Cu$_2$O Heterostructure Core-Shell Nanowire Photoanode

FIGURE 1 | Top-view- and cross-sectional SEM images of the (a,b) bare InGaN nanowires (NWs) and (c,d) InGaN NWs after 40 s Cu$_2$O electrodeposition. (e) X-ray photoelectron spectroscopy taken several days after Cu$_2$O electrodeposition.

Further discussed in the section Stability on the stability of the InGaN/Cu$_2$O heterostructure operated as photoanode.

The element distributions are imaged by EDS mappings shown in Figure 2. The mappings are taken from the NW cross-sectional part shown in the SEM image in Figure 2a. Figure 2b displays the total EDS intensity. The individual mappings of In, Ga, N, Cu, O, and Si are shown in Figures 2c–h, respectively. The mappings of In, Ga, N, and Si show the uniform element distribution in the InGaN NWs on the Si substrate. The Cu and O element distributions reveal the deposition of Cu$_2$O on the NW sidewalls, penetrating from the top to the bottom with decreasing EDS intensities.

The deposition of Cu$_2$O on the InGaN NW sidewalls is directly resolved in the TEM image in Figure 3a, and the corresponding EDS element mappings of In, Ga, N, and Cu are shown in Figures 3b–f. Figure 3b gives the total EDS intensity and Figures 3c–f depict the individual In, Ga, N, and Cu element mappings. The element contrast for O, which is not shown, is very weak. Cu$_2$O forms a rather uniform shell on the InGaN NW sidewalls. The Cu$_2$O shell is composed of ultra-small nanocrystals, of ~5 nm in diameter, with random orientation, seen in the high-resolution TEM image in the inset of Figure 3a. The spaces between the NWs are not completely filled. The element distributions of In, Ga, and N follow the straight geometry of the InGaN NWs. The element distribution of Cu is more blurred and smeared out, as expected for the InGaN/Cu$_2$O core-shell structure. The different Cu$_2$O electrodeposition behavior on the InGaN NW sidewalls and NW tops is addressed in the following section.

Photoelectrochemical Properties

Figure 4A depicts the LSV curves for the InGaN/Cu$_2$O core-shell NWs and for the bare InGaN NWs under chopped 1-sun (AM 1.5, 100 mW/cm$^2$) illumination. The Cu$_2$O electrodeposition time is 40 s. The LSV scan is taken from +1 V to 0 V vs. Ag/AgCl (1.6 V to 0.6 V vs. RHE) with a scan rate of 10 mV/s. In the presence of the Cu$_2$O shell, the photocurrent density at positive voltage is enhanced by up to 700%. At negative voltage, no photoresponse is observed, as seen in the CV scan from +1 V to −1 V in Supplementary Figure 4A. This demonstrates the role of Cu$_2$O in enhancing the OER when the InGaN/Cu$_2$O core-shell NWs operate as photoanodes. The optimum Cu$_2$O electrodeposition time of 40 s is provided from the plot of the photocurrent density vs. electrodeposition time, which is taken for an applied voltage of +0.6 V vs. Ag/AgCl (1.2 V vs. RHE), as shown in Figure 4B. When the Cu$_2$O deposition time is increased much longer, the individual cubic Cu$_2$O nanocrystals on top of the InGaN NWs, as shown in the SEM image in Figure 1c, increase in density and coalesce to form a compact layer. For such thick Cu$_2$O layers, the usual behavior of Cu$_2$O as a photocathode for the HER at negative voltage is recovered, as shown in the LSV and I–t curves in Supplementary Figures 4B–D. In this case, there is no photoresponse for positive voltage. This confirms that the thin Cu$_2$O shell-layer is responsible for the enhanced photoanode performance, whereas the much thicker/larger cubic Cu$_2$O nanocrystals act as photocathodes.

A first indication of the reproducibility and stability of the InGaN/Cu$_2$O core-shell NW photoanode is given by the I–t curve taken at +0.6 V vs. Ag/AgCl (1.2 V vs. RHE) under...
chopped 1-sun illumination in Figure 4C with the curve for the bare InGaN NWs for comparison. The curves reproduce a 700% increase of the photocurrent density in the presence of the Cu$_2$O shell-layer, in agreement with the LSV measurements, and a good photo-switching behavior is observed.

The increased photocurrent density is attributed to the p–n junction with related upward energy band bending, formed at the InGaN/Cu$_2$O interface for the m-plane NW sidewalls. The upward energy band bending enhances the separation of the photogenerated carriers and the transport of photogenerated holes toward the surface. This is maximized for sufficiently thin and fully depleted Cu$_2$O layers on the NW sidewalls when the upward energy band bending continues to the surface. This is guaranteed for up to a Cu$_2$O thickness where the energy band positions at the surface, taking into account only the ionized acceptor density, reach the values given by the downward energy band bending for bulk Cu$_2$O of typically 0.4 V, which are, in turn, given by the surface Fermi-level pinning. This accounts for the increase of the photocurrent at positive voltage with increasing Cu$_2$O shell-layer thickness, passing up to a maximum and finally reverse with photoresponse at negative voltage for thick Cu$_2$O layers on the InGaN NW tops. The sparse cubic Cu$_2$O nanocrystals formed together with the Cu$_2$O shell-layers do not affect the photoanode behavior. Their size rather indicates near-surface downward energy band bending without photoresponse for positive voltage. A scheme of the described InGaN/Cu$_2$O energy band structure and photogenerated carrier transport for increasing Cu$_2$O shell-layer thickness is drawn in Figure 4D.
The thin CuO surface layer [32] formed due to the self-limiting oxidation of Cu$_2$O, which is discussed in more detail below, is also indicated.

The maximum photocurrent density for a Cu$_2$O shell-layer thickness of several 10 nm agrees with typical ionized acceptor densities for Cu$_2$O of $10^{15}$–$10^{16}$ cm$^{-3}$ and the related width of the depletion zone [33, 34]. However, the decrease of the photocurrent density might set in before the maximum width of the Cu$_2$O depletion zone is utilized because of filling up of the spaces between the InGaN NWs with Cu$_2$O, whereas the light absorption in the Cu$_2$O layer might contribute to the increase of the photocurrent density with Cu$_2$O layer thickness. Unfortunately, we have no direct experimental access to the electrical properties of the thin Cu$_2$O shell-layers electrodeposited on the InGaN NW sidewalls.

Apart from that, in contrast to the rectifying p–n junction for Cu$_2$O on the m-plane InGaN NW sidewalls, leading to the upward energy band bending, the electrical contact is more ohmic for Cu$_2$O on the c-plane NW tops [23], not supporting the upward energy band bending. This has been attributed to the varying characteristics of the surface states for the m-plane and the c-plane of In-rich InGaN: Negatively charged surface states in the bandgap for the m-plane vs. positively charged surface states in the conduction band for the c-plane of In-rich InGaN, leading to upward/downward near-surface energy band bending for the m-plane/c-plane for In-rich InGaN and, accordingly, to near-surface electron depletion for the m-plane/near-surface electron accumulation for the c-plane of In-rich InGaN.

The different characteristics of the surface states and resulting InGaN/Cu$_2$O electrical contact for the m-plane and the c-plane of In-rich InGaN may also explain the different electrodeposition behavior of Cu$_2$O on the m-plane NW sidewalls and c-plane NW tops. The formation of the rectifying p–n junction for Cu$_2$O electrodeposition on the NW sidewalls, which is forward biased under the electrodeposition conditions, might lead to a more controlled, charge-transfer limited, quasi-two-dimensional growth with the nucleation and fast coalescence of ultra-small and dense Cu$_2$O nanocrystals. The more ohmic resistance for the NW tops might favor mass-transport limited growth, leading to the larger, randomly oriented three-dimensional Cu$_2$O nanocrystals with lower density, as usually observed on conductive substrates [35].

The PL spectra taken at room temperature from the InGaN/Cu$_2$O core-shell NWs and the bare InGaN NWs are shown in Figure 5A. The PL intensity for the InGaN/Cu$_2$O core-shell NWs is significantly reduced. This is the PL from InGaN. No PL is observed from Cu$_2$O. The PL process is fairly complex, involving light absorption, photogenerated carrier relaxation and
Zhao et al. InGaN/Cu$_2$O Heterostructure Core-Shell Nanowire Photoanode

**FIGURE 4** | (A) Linear sweep voltammetry measurements of the InGaN/Cu$_2$O core-shell NWs and the bare InGaN NWs under chopped illumination. (B) Photocurrent density at +0.6 V vs. Ag/AgCl (1.2 V vs. reversible hydrogen electrode [RHE]) as a function of the Cu$_2$O electrodeposition time. (C) I–t measurements at +0.6 V vs. Ag/AgCl (1.2 V vs. RHE) of the InGaN/Cu$_2$O core-shell NWs and the bare InGaN NWs under chopped illumination. (D) Scheme of the InGaN/Cu$_2$O energy band diagram and photogenerated carrier transport for increasing Cu$_2$O layer thickness. Int. indicates the InGaN/Cu$_2$O interface and Surf. indicates the surface of Cu$_2$O shell-layers with different thicknesses. The thin Cu$_2$O surface layer is also shown.

diffusion, and radiative electron–hole recombination. However, the reduced PL intensity for InGaN and the absence of PL from Cu$_2$O strongly support the enhanced photogenerated carrier separation and transfer in the presence of the Cu$_2$O shell-layer. The radiative electron–hole recombination is suppressed, overcompensating the absorption, which might be enhanced by the Cu$_2$O shell-layer.

With regard to all the steps in photoanode operation, i.e., absorption, photocarrier separation and transfer, and surface reaction, however, the Cu$_2$O surface is not anticipated to exhibit particular surface catalytic activity for the OER reaction. This is seen in the EIS measurements shown in Figure 5B of the InGaN/Cu$_2$O core-shell NWs and the bare InGaN NWs taken at an applied voltage of +0.6 V in the dark and under illumination. The half-circle in the Nyquist plots, whose diameter gives the charge transfer resistance, $R_{ct}$, is large for the InGaN/Cu$_2$O core-shell NWs and for the bare InGaN NWs, indicating an overall large $R_{ct}$. To be precise, $R_{ct}$ includes the charge transfer through the surface in the dark plus the charge transfer toward the surface inside the photoanode under illumination. Therefore, the large $R_{ct}$ for the InGaN/Cu$_2$O core-shell NWs, which is also under illumination, indicates a dominating slow charge transfer through the surface, resulting in a slow surface reaction. Therefore, to demonstrate competitive photocurrent densities, the photoanode fabrication process is completed by the electrodeposition of NiOOH co-catalyst comprising only earth-abundant elements. As shown in Figures 5C,D, the LSV and I–t curves (at +0.8 V, where the dark current continues to be low) of the InGaN/Cu$_2$O/NiOOH complete NW photoanode and the InGaN/Cu$_2$O core-shell NW photoanode demonstrate that photocurrent densities, reaching the mA/cm$^2$ regime, are achieved.

**Stability**

Figure 6 shows the CV curves recorded from −1 V to +1 V vs. Ag/AgCl in the dark and under 1-sun illumination with a scan rate of 100 mV/s for the bare InGaN NWs in
FIGURE 5 | (A) Photoluminescence (PL) spectra taken at room temperature of the InGaN/Cu$_2$O core-shell NWs and the bare InGaN NWs. (B) Electrochemical impedance spectroscopy (EIS) taken at +0.6 V vs. Ag/AgCl (1.2 V vs. RHE) in the dark and under illumination of the InGaN/Cu$_2$O core-shell NWs and the bare InGaN NWs. (C) LSV- and (D) I–t measurements at +0.8 V vs. Ag/AgCl (1.4 V vs. RHE) of the InGaN/Cu$_2$O/NiOOH complete NW photoanode and the InGaN/Cu$_2$O core-shell NWs under chopped illumination.

Figure 6A, the InGaN/Cu$_2$O core-shell NWs in Figure 6B, and the InGaN/Cu$_2$O/NiOOH complete NW photoanode in Figure 6C. The bare InGaN NWs show solely capacitative behavior without reaction peaks, whereas the InGaN/Cu$_2$O core-shell NWs exhibit distinct reduction and oxidation peaks at −0.46 V and +0.3 V, respectively. These are again absent for the InGaN/Cu$_2$O/NiOOH complete NW photoanode. This indicates excellent chemical stability of the bare InGaN NWs, which has been widely reported in previous studies, whereas the InGaN/Cu$_2$O core-shell NWs exhibit reduction and oxidation reactions. This is well-known and has been deeply investigated for the reduction and oxidation of thick Cu$_2$O layers used as photocathodes [25]. The NiOOH co-catalyst stabilizes the Cu$_2$O shell-layer by promoting the water splitting rather than the reduction and oxidation of Cu$_2$O.

Importantly, as we show in this study, the InGaN/Cu$_2$O core-shell NWs without NiOOH are stable when operated solely at positive voltage as photoanode. They are self-stabilized by a thin CuO surface layer. The chain of arguments goes as follows: (i) The oxidation peak at positive voltage occurs only after prior scanning over the reduction peak at negative voltage, as is done for the CV measurements (Figure 6B). When scanning only from 0 V to +1 V and back, while avoiding the reduction peak, no reaction peaks are visible in the CV curves (Figure 6D). Also, when scanning from +1 V to 0 V or to −1 V, no oxidation peaks were visible in the LSV measurements in Figure 4A and Supplementary Figure 4A. (ii) XPS shows that the oxidation peak originates from the oxidation of Cu$_2$O to CuO. In the XPS taken after scanning at positive voltages, as shown in Figure 6E, the CuO-related features are enhanced compared to those in the
XPS taken before, as shown in Figure 1e. We do not exclude that the oxidation of Cu contributes to the oxidation peak after prior reduction of Cu$_2$O [36]. However, this is irrelevant for the photoanode operation as no Cu-related features are observed in the XPS after scanning at positive voltage. (iii) In the I–t measurement taken over an extended duration of time at +0.6 V under chopped illumination, as shown in Figure 6F, an initial dark current is observed. This dark current appears when the measurement is started after prior to reduction at negative voltage. The dark current vanishes within 100 s. The photocurrent appears unaffected over the whole measurement. The total current and the dark current initially decrease with the same rate and finally, the total current equals the photocurrent and becomes very stable. Altogether, from observations (i)–(iii), there is oxidation of Cu$_2$O in air and at positive voltage, forming a thin surface layer of CuO, as already discussed in the context of the XRD, Raman, and XPS measurements. The oxidation quickly saturates such that the Cu$_2$O layer becomes self-stabilized by the thin CuO surface layer. Photo-oxidation is not involved in CuO formation as only the dark current changes but the photocurrent does not. The formation of a nanoscale thin, self-limiting surface oxide is all-too-common for materials which can undergo oxidation, such as those observed for the bare InGaN photoanode and Si. The self-limiting thickness is expected, considering that Cu$_2$O is the most stable Cu oxide. In TEM, the presence of the CuO layer is not seen. Importantly, the thin CuO surface layer does not significantly affect the photocurrent, although it imposes a small energy barrier for hole transport, as indicated in Figure 4D. Hole transport might be easy because of thermionic emission and tunneling or because the thin CuO layer is leaky for the transition of holes. Self-stabilized by the thin CuO surface layer, Supplementary Figure 5 shows that the InGaN/Cu$_2$O photoanode only exhibits a very small photocurrent decrease after many hours of operation.

CONCLUSIONS

Large enhancement of the photocurrent is achieved for a novel n-InGaN/p-Cu$_2$O NW photoanode compared to a bare InGaN NW photoanode for the OER in solar water splitting. The electrodeposited Cu$_2$O layer forms a thin shell around the InGaN NW cores. The enhancement of the photocurrent is due to maximized photogenerated carrier separation and hole transfer to the surface in the fully depleted Cu$_2$O shell-layer of the p–n Cu$_2$O/InGaN heterojunction. Thick Cu$_2$O layers on top of the InGaN NWs act as usual photocathodes for the HER. The functional InGaN/Cu$_2$O heterostructure core-shell NW photoanode is chemically self-stabilized at positive applied voltage by a thin CuO surface layer. The photocurrent reaches the competitive mA/cm$^2$ range after final deposition of the earth-abundant NiOOH co-catalyst, establishing the InGaN/Cu$_2$O/NiOOH complete NW photoanode. Moreover, this is an important proof of the successful formation of a rectifying p–n junction at the InGaN/Cu$_2$O heterointerface with the Cu$_2$O layer acting as an efficient hole collection and transport...
layer on In-rich InGaN, solving the p-type doping problem with far-reaching impact for devices.

**DATA AVAILABILITY STATEMENT**

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author/s.

**AUTHOR CONTRIBUTIONS**

YZ performed the growth and the measurements with the support of LX. GZ and RN discussed and interpreted the results. RN supervised the research and drafted the manuscript together with YZ. All authors discussed and commented on the manuscript and read and agreed to the published version of the manuscript.

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**FUNDING**

This study was supported by: the Program for Chang Jiang Scholars and Innovative Research Teams in Universities (No. IRT_17R41), the Science and Technology Program of Guangzhou (No. 2019050001), the Guangdong Provincial Key Laboratory of Optical Information Materials and Technology (Grant No. 2017B030301007), the Provincial Natural Science Foundation of Guangdong program (Grant No. 2018A030313368), the MOE International Laboratory for Optical Information Technologies, and the 111 Project.

**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fphy.2021.684283/full#supplementary-material
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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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