Abstract: Highly efficient photoelectrochemical (PEC) water oxidation under solar visible light is crucial for water splitting to produce hydrogen as a source of sustainable energy. Particularly, silver-based nanomaterials are important for PEC performance due to their surface plasmon resonance which can enhance the photoelectrochemical efficiency. However, the PEC of ZnO/Ag₂WO₄/AgBr with enhanced visible-light water oxidation has not been studied so far. Herein, we present a novel photoelectrodes based on ZnO/Ag₂WO₄/AgBr nanorods (NRs) for PEC application, which is prepared by the low-temperature chemical growth method and then by successive ionic layer adsorption and reaction (SILAR) method. The synthesized photoelectrodes were investigated by several characterization techniques, emphasizing a successful synthesis of the ZnO/Ag₂WO₄/AgBr heterostructure NRs with excellent photocatalysis performance compared to pure ZnO NRs photoelectrode. The significantly enhanced PEC was due to improved photogeneration and transportation of electrons in the heterojunction due to the synergistic effect of the heterostructure. This study is significant for basic understanding of the photocatalytic mechanism of the heterojunction which can prompt further development of novel efficient photoelectrochemical-catalytic materials.

Keywords: ZnO nanorods, Silver tungsten, Silver bromide, Heterojunction, Photoelectrodes, Water oxidation
photoelectrode [24]. The result showed high PEC performance in comparison with the ZnO which is attributed to the suppression of the high recombination rate and the shift of the absorption toward the visible-wavelength region due to surface plasmonic resonance (SPR) [24]. In spite of the high performance of the ZnO/Ag/Ag2WO4 due to the deposition of the Ag2WO4 into the ZnO nanorods (NRs), still the band gap did not utilize enough visible light to yield an efficient PEC electrode. Thus, smaller-band-gap semiconductor, e.g., AgBr, can be deposited onto the ZnO/Ag2WO4 heterostructure to further increase the absorption of the visible-light wavelength of the solar spectrum, which further improves the PEC efficiency. AgBr can be a good sensitizer because the same Ag-based material of the Ag2WO4 could give better light absorption and photoelectron transportation with ZnO/Ag2WO4/AgBr heterostructure.

In this work, hydrothermal growth route followed by successive ionic layer adsorption and reaction (SILAR) method was used to prepare ZnO/Ag2WO4/AgBr NRs photoelectrode for PEC water splitting analysis. This implies electrode preparation, characterization, PEC experiments and presenting the proposed electron path during the photocatalytic reaction. Ag2WO4 was deposited into the sample in order to develop plasmonic sensitizer to improve the utilization of the solar power and accelerating the charge carrier transfer. The Ag in the sample with SPR effect can enhances the photocatalytic performance due to the enhancement of the absorption of the solar light. The loading of the AgBr could further enhances the absorption of light in the visible range due to the lower band gap of the AgBr. As far as we know, it is the first time to prepare ZnO/Ag2WO4/AgBr NRs photoelectrode by deposition of Ag2WO4 and AgBr on the ZnO NRs using SILAR method for PEC performance. The ZnO/Ag2WO4/AgBr NRs photoelectrodes were prepared with 5, 10, 15 and 20 SILAR cycles for AgBr amount optimization.

**Experimental Part**

**ZnO/Ag2WO4/AgBr Photoelectrodes Preparation**

The photoelectrodes were prepared using three steps: growth of the ZnO NRs, deposition of Ag2WO4 and AgBr as illustrated in Fig. 1.

The pristine ZnO NRs were prepared on the FTO covered by a seed layer of ZnO nanoparticles (NPs) prepared by the hydrothermal growth similar to our previous work [24]. The precursor solutions were prepared using equal molecular (0.05 M) of zinc nitrate hexahydrate (Zn(NO3)2·6H2O) and hexamethylenetetramine (HMT). Then, the FTO substrates with the seed layer were fixed upside down in a Teflon sample holder and then placed in a beaker containing the growth solution and were kept in a preheated oven at 90 °C for 5 h. After the completion of the growth duration, the samples were taken out and left to cool down to room temperature. Finally, the samples were washed with DI water to avoid any unwanted particles or residuals and then dried with flowing nitrogen gun.

The ZnO/Ag2WO4 NRs photoelectrode was prepared using SILAR method in the same way as our previous work [24, 25]. An anionic and cationic aqueous precursor solutions were prepared separately using 0.1 M of silver nitrate Ag(NO3) and 0.1 M of sodium tungstate (Na2WO4·2H2O). The deposition took place by immersion of the prepared ZnO NRs samples into Ag(NO3) solution for 2 min to absorb the silver ions (Ag+), and then, they were washed with DI water to remove excess ions or any other unwanted particles. Then, the samples were immersed into the Na2WO4·2H2O solution for 2 min and again washed with DI water. This cycle was repeated several times to obtain enough Ag2WO4 particles on the ZnO NRs. Then, the samples were dried in an oven at 60 °C for 3 h to obtain good adhesion of the Ag2WO4 on the ZnO NRs.

The ZnO/Ag2WO4/AgBr NRs photoelectrode was prepared also using the SILAR method. Anionic and cationic aqueous precursor solutions were prepared separately using 0.1 M of silver nitrate Ag(NO3) and 0.1 M of sodium bromide (NaBr). The deposition took place by immersion of the prepared ZnO NRs samples into Ag(NO3) solution for 2 min and again washed with DI water. This cycle was repeated several times to obtain enough Ag2WO4 particles on the ZnO NRs. Then, the samples were immersed into the Na2WO4·2H2O solution for 2 min and again washed with DI water. This cycle was repeated several times to obtain enough Ag2WO4 particles on the ZnO NRs. Then, the samples were dried in an oven at 60 °C for 3 h to obtain good adhesion of the Ag2WO4 on the ZnO NRs.

The ZnO/Ag2WO4/AgBr NRs photoelectrode was prepared also using the SILAR method. Anionic and cationic aqueous precursor solutions were prepared separately using 0.1 M of silver nitrate Ag(NO3) and 0.1 M of sodium bromide (NaBr). The deposition took place by immersion of the prepared ZnO NRs samples into Ag(NO3) solution for 2 min to absorb the silver ions (Ag+), and then, they were washed with DI water to remove excess ions or any other particles. Then, the sample is
immersed into the NaBr solution for 2 min and again washed with DI water. This cycle was repeated for several times to obtain enough AgBr particles on the ZnO/Ag2WO4 photoelectrode followed by drying in the oven at 60 °C for 3 h for better adhesion of the nanoparticles. Also, the ZnO/Ag2WO4/AgBr NRs photoelectrodes were prepared with 5, 10, 15 and 20 SILAR cycles for PEC analysis to optimize the AgBr content.

Characterization
The structural properties of our photoelectrodes were studied using powder X-ray diffraction (XRD) with a Philips powder diffractometer (1729 PW) connected to a Cu Kα radiation source at the generator voltage of 40 kV and the current of 40 mA. A field-emission scanning electron microscope (FE-SEM) equipped with a Sigma 500 Gemini field emission gun operating at 10 kV was used to investigate the sample morphology. The chemical composition of the samples was investigated by X-ray photoelectron spectroscopy (XPS) using a Kratos AXIS Ultra DLD equipped with a monochromatic Al Kα X-ray source. CasaXPS software was used to analyze the data. In order to analyze the optical properties, the UV–visible spectroscopy (Perkin-Elmer Lambda 900 system) was used.

Photoelectrochemical Measurements
The photoelectrochemical performance was studied by three electrode photoelectrochemical measurements using SP-200 potentiostat (Bio-Logic, Clai, France). A platinum (Pt) mesh (as the counter electrode) and a standard silver/silver chloride (Ag/AgCl) in 3 M KCl (as a reference electrode) were used with (0.1 M) of sodium sulfate (Na2SO4) electrolyte. The total immersed area of the electrode in the electrolyte was 1 cm x 1 cm. The visible-light radiation was obtained by a solar simulator that uses a 100-W ozone free xenon lamp with an output power of 1 sun (AM 1.5).

Result and Discussion
Characterization Analysis
Figure 2 shows the XRD spectra of ZnO, ZnO/Ag2WO4 and ZnO/Ag2WO4/AgBr samples. It is clear that all the obtained XRD diffraction peaks in the ZnO sample correspond to the hexagonal wurtzite of ZnO (JCPDS no. 36-1451) suggesting no other phases of ZnO were observed. In the XRD pattern of the ZnO/Ag2WO4 sample, additional peaks were identified, which belonged to Ag2WO4 (JCPDS no 33-1195), indicating the successful deposition of the Ag2WO4 into the ZnO NRs. In the XRD pattern of the ZnO/Ag2WO4/AgBr heterostructure, the planes (200) and (220) labeled in the figure were assigned to the AgBr (JCPDS no 06-438), confirming the formation of AgBr on the ZnO/Ag2WO4 heterostructure (Additional file 1).

The FE-SEM images of the ZnO/Ag2WO4 and ZnO/ Ag2WO4/AgBr NRs photoelectrodes are presented in Fig. 3a, b, respectively. In Fig. 3a, we could clearly see that Ag2WO4 particles were deposited onto the ZnO NRs. In Fig. 3b, we can notice that a larger particle size is shown for the ZnO/Ag2WO4/AgBr sample compared to the ZnO/Ag2WO4 NRs photoelectrode, with semi-crystal structure suggesting the successful deposition of the AgBr onto the sample. Also, tiny particles can be observed, which could be due to different counterpart deposited onto the ZnO NRs. Figure 3c shows EDX spectrum of the ZnO/Ag2WO4/AgBr NRs photoelectrode which display the relative amount of Zn, O, Ag, W and Br. Surprisingly, unexpected Cl element was detected which is attributed to contamination. In addition to that, the EDX mapping is displayed in Fig. 3d to show the elemental distribution over the sample. Figure 3e shows cross-sectional SEM image and mapping of the ZnO/Ag2WO4/AgBr photoelectrode. It is worth to note that good elemental distribution on the sample could result in high-quality heterostructure yielding an enhanced PEC performance. Moreover, the effect of the AgBr loading on the morphology was investigated for the ZnO/Ag2WO4/ AgBr samples with different cycles of AgBr loading as shown in Fig. 4. As the loading cycles increases, more AgBr particles are deposited onto the sample, reaching full coverage of the surface by the AgBr with 20 cycles of loading cycles.

To further evaluate the ZnO/Ag2WO4/AgBr NRs heterostructure, XPS measurement was taken on all
elements present in the structure. Different high-resolution XPS (HR-XPS) spectra are shown in Fig. 5. The Zn 2p spectrum shows two peaks which matches to Zn 2p\textsubscript{3/2} and Zn 2p\textsubscript{1/2} with a peak position at 1022.5 and 1045.7 eV, respectively \cite{26, 27}, as shown in Fig. 5a. The Ag 3d spectrum is also divided into two peaks with a peak position at 365.1 and 371.1 eV which is attributed to Ag 3d\textsubscript{5/2} and Ag 3d\textsubscript{3/2}, respectively, as shown in Fig. 5b. The Ag 3d\textsubscript{5/2} is then divided into two different peaks at 365.0 and 365.8 eV, and the Ag 3d\textsubscript{3/2} peak is also divided into two different peaks at 371.0 and 371.6 eV.

The peaks at low energies 365.0 and 371.0 eV are assigned to the Ag\textsuperscript{2+} in AgBr, whereas the peaks at higher energies 365.8 and 371.6 are assigned to metallic Ag\textsuperscript{0} \cite{28, 29}. Figure 5c shows the spectrum for W 4f which has two peaks positioned at 35.0 and 37.1 eV corresponding to W 4f\textsubscript{5/2} and W 4f\textsubscript{7/2}, respectively \cite{29}. The Br 3d also exhibits two separate peaks positioned at 68.0 and 69.0 eV assigned to Br 3d\textsubscript{5/2} and Br 3d\textsubscript{3/2}, respectively, as shown in Fig. 5d \cite{30}. The high-resolution XPS analysis agrees with the EDX measurement and confirms that the intended ZnO/Ag\textsubscript{2}WO\textsubscript{4}/AgBr NRs composite heterostructure has been successfully achieved.

From the UV–Vis absorption spectra, a redshift in the optical absorption in the visible-light region was observed for the ZnO/Ag\textsubscript{2}WO\textsubscript{4}/AgBr NRs sample compared to the other samples. Also, the optical band gap energy was reduced to 2.94 eV, which was found from the plots of (αhν\textsuperscript{2}) versus photon energy (hν) from the UV–Vis absorption spectra as shown in Fig. 6a. The enhancement in the visible-light absorption ability of the ZnO/Ag\textsubscript{2}WO\textsubscript{4}/AgBr NRs heterostructure can be assigned
Fig. 3 continued
to the deposition of the AgBr due to its lower band gap energy compared to the ZnO. Higher ability to absorb visible light in the heterostructure will be valuable for increasing the efficiency of the PEC reaction.

**Photoelectrochemical Analysis**

The PEC properties of the different photoelectrodes investigated by the photoresponse over time of the NRs-based photoelectrodes using chronoamperometry measurements which recorded the photocurrent density versus time in the dark under on/off solar irradiation with an applied potential of +0.5 V are shown in Fig. 6b. The photocurrent densities were found to be 1.6 mA/cm² for the ZnO photoelectrode which increases up to 2.7 mA/cm² for ZnO/Ag₂WO₄ photoelectrode. Further enhancement of the photocurrent density was obtained after the deposition of the AgBr. The enhancement of the photocurrent density is observed when increasing the amount of the AgBr loading with different SILAR cycles. At the lower cycle of 5 times, small amount of photocurrent density (2.7 mA/cm²) was detected due to small number of particles that distributed into the surface as it can be seen in the SEM of the 5 time. Increasing the number of cycles to 10 resulted in the highest photocurrent density of 3.3 mA/cm² with more particles distribution. Further increase of AgBr loading result in a reduction of the photocurrent density to 2 and 1.8 mA/cm² for ZnO/Ag₂WO₄/AgBr (15 SILAR cycles) and ZnO/Ag₂WO₄/AgBr NRs (20 SILAR cycles) photoelectrodes, respectively, but the interface structure became more uniform size distribution with more crystallization. The possible reason for this reduction is that increasing the AgBr amount leads to form larger aggregates around the ZnO NRs, which might destroy the junctions and reduces the separation of the charge carriers at the interfaces of the heterojunction. Therefore, it is important to optimize the photoelectrodes for higher PEC activity. Due to the best PEC result observed from the ZnO/Ag₂WO₄/AgBr NRs (10 SILAR cycles) photoelectrode, this sample is used for all the measurement as optimum photoelectrode.

Linear sweep voltammetry (LSV) measurement was taken under the illumination of solar light and dark conditions at a potential of 1.23 V (vs. Ag/AgCl) electrode. Under the dark condition, a negligible photocurrent...
density of the ZnO NRs photoelectrode was noticed, which indicates good surface quality of the ZnO NRs as shown in Fig. 6c. In comparison with the ZnO NRs photoelectrode, the ZnO/Ag\textsubscript{2}WO\textsubscript{4} and the ZnO/Ag\textsubscript{2}WO\textsubscript{4}/AgBr NRs photoelectrodes showed a small photocurrent density under dark condition 0.009 and 0.015 mA/cm\textsuperscript{2}, respectively, measured at a potential of 1.23 V indicating an enhancement in the electrical conductivity. Under the illumination of solar light, a lower photocurrent density of 0.7 mA/cm\textsuperscript{2} was observed for ZnO NRs, whereas the photocurrent density was highly increased to 2.3 and 3.8 mA/cm\textsuperscript{2} for the ZnO/Ag\textsubscript{2}WO\textsubscript{4} and ZnO/Ag\textsubscript{2}WO\textsubscript{4}/AgBr NRs photoelectrodes, respectively. The remarkably improved photocurrent density of the ZnO/Ag\textsubscript{2}WO\textsubscript{4}/AgBr NRs photoelectrode could be ascribed to the heterojunction effect upon the deposition of Ag\textsubscript{2}WO\textsubscript{4} and the AgBr NPs onto the surface of the ZnO NRs. Which indicates that a high density of the photogenerated electrons can be transferred from the ZnO/Ag\textsubscript{2}WO\textsubscript{4}/AgBr NRs photoelectrode to the counter electrode through the external circuit providing higher photocurrent density. This is attributed to the enhanced charge carrier separation, transportation efficiency, the high absorption of solar light and the shift of the absorption into the visible-light range.

Mott–Schottky (M–S) analysis is commonly used in PEC for photoelectrodes characterization to understand the electronic properties and the change in the carrier density, which also gives valuable information about the flat band potentials (E\textsubscript{FB}) of the samples [24,
Mott–Schottky plot was obtained by electrochemical impedance measurement at room temperature with selected frequency ~ 1 kHz which is based on the capacitance versus applied potential measurements and is shown in Fig. 7. Figure 7 shows the corresponding M–S plots of pristine ZnO NRs, ZnO/Ag2WO4, and ZnO/Ag2WO4/AgBr photoelectrodes. It can be seen that all of the prepared samples exhibited positive slopes, showing their n-type nature as expected.

The position of the EFB is estimated by linear extrapolation of the linear region of the curve; the x-axis intercept gives the values of the EFB, which were approximately equal to 0.41, 0.70 and 0.74 V for pristine ZnO NRs, ZnO/Ag2WO4 and ZnO/Ag2WO4/AgBr NRs photoelectrodes, respectively. Thus, the shift in EFB to higher values could be attributed to the change in charge carrier

Fig. 6 a The plots of (αhν)² versus hν. b Chronoamperometry I-t curves with solar irradiation on/off cycles, and c LSV curves under dark and visible-light conditions of the ZnO NRs, ZnO/Ag2WO4 and ZnO/Ag2WO4/AgBr photoelectrodes

Fig. 7 M–S plots of pristine ZnO NRs, ZnO/Ag2WO4, and ZnO/Ag2WO4/AgBr photoelectrodes
concentration in the heterojunctions and the change in Helmholtz layer potential [25]. The presence of more surface states can lead to a significant change in the band position that might shift the Fermi level to higher value [24, 31].

Using the dielectric permittivity of the ZnO and the permittivity of the vacuum, the charge carrier density \( N_d \) could be estimated using the following formula [32]:

\[
N_d = \frac{2}{q \varepsilon \varepsilon_0} \left[ \frac{1}{d(1/C^2)/dv} \right]
\]

The charge carrier densities were estimated to be \( 7.5 \times 10^{18} \) and \( 1.1 \times 10^{20} \) cm\(^{-3} \) for the pristine ZnO NRs and ZnO/Ag\(_2\)WO\(_4\)/AgBr NRs photoelectrodes, respectively. Obviously, \( N_d \) of the ZnO/Ag\(_2\)WO\(_4\)/AgBr NRs photoelectrodes is enhanced, which explains the improved PEC activities of this photoelectrode under solar light compared to pristine ZnO NRs.

**Proposed Mechanism**

The energy band position of the ZnO/Ag\(_2\)WO\(_4\)/AgBr heterojunction with possible electron transfer path is illustrated in Fig. 8. Light with lower-energy photons will excite electrons from the VB to the CB of the AgBr due to its suitable band gap (~2.6 eV), whereas higher-energy photons can excite electrons in the ZnO and Ag\(_2\)WO\(_4\) semiconductors. Rapid electrons transfers will take place quickly where electrons are transferred from the CB of the AgBr to the CB of the ZnO and Ag\(_2\)WO\(_4\) consequently. Then, electrons will then be captured and transferred to the photoelectrode contact for reduction reaction in the Pt electrode site where H\(_2\) should be released. Holes that are left in the VB will perform oxidation reaction and O\(_2\) should be released. The efficient electrons transfer between the heterojunction can reduce the recombination rate and enhances the photocatalytic reaction.

**Conclusion**

In summary, ZnO/Ag\(_2\)WO\(_4\)/AgBr NRs heterostructure photoelectrode was prepared successfully using the hydrothermal growth route followed by the SILAR method. The characterization analysis revealed that the Ag\(_2\)WO\(_4\)/AgBr was successfully deposited onto the ZnO NRs. The photocurrent density of the ZnO/Ag\(_2\)WO\(_4\)/AgBr NRs (10 SILAR cycles) photoelectrode was increased 5 times compared to the ZnO NRs photoelectrode under visible sun radiation. Also, the photoresponse over time showed an improvement in the photocurrent density for the ZnO/Ag\(_2\)WO\(_4\)/AgBr NRs (10 SILAR cycles) photoelectrode (3.3 mA/cm\(^2\)) in comparison with that of the ZnO NRs photoelectrode (1.6 mA/cm\(^2\)). The enhancement in the PEC response is attributed to the synergistic effect due to the deposition of the Ag\(_2\)WO\(_4\) and the AgBr NPs onto the surface of the ZnO NRs. This deposition increased the
absorption of the visible light and the accompanied lowered recombination rate. Also, it was found that higher amount of AgBr can lead to larger aggregates into the heterostructure which might destroy the heterojunction and reduces the PEC performance. The high potential of the ZnO/Ag2WO4/AgBr NRs photoelectrode for the PEC water splitting makes this photoelectrode a promising candidate for hydrogen production.

Supplementary Information

The online version contains supplementary material available at https://doi.org/10.1186/s11671-021-03586-z.

Acknowledgements

The authors acknowledge partial financial support from the department of Science and Technology (ITN), at Campus Norrköping, Linköping University, Sweden.

Authors’ contributions

EM, RA, MW and ON conceived the research idea and analyzed the data. EM and RA prepared, measured and characterized all samples and wrote the first draft of the manuscript. PR conducted the XPS measurements and wrote the XPS part. All authors read and revised the manuscript and discussed the research results. All authors read and approved the final manuscript.

Funding

Open access funding was provided by Linköping University, Sweden.

Availability of data and materials

All data relevant for the reproduction of the results presented in this work are included in this published article or in its supplementary information (SI) file.

Declarations

Competing interests

The authors declare that they have no competing interests.

Author details

1Department of Sciences and Technology, Linköping University, Campus Norrköping, 601 74 Norrköping, Sweden. 2Department of Physics, Chemistry and Biology (IFM), Linköping University, 58183 Linköping, Sweden.

References

1. Fujishima A, Honda K (1972) Electrochemical photolysis of water at a semiconductor electrode. Nature 238:37–38
2. Ahmad H, Kamarudin SK, Minggu LJ, Kassim M (2015) Enhancing methanol oxidation with a TiO2-modified semiconductor as a photo-catalyst. Renew Sust Energ Rev 43:599
3. Ros C, Andreu T, Monarte JR (2020) Photoelectrochemical water splitting: a road from stable metal oxides to protected thin film solar cells. J Mater Chem B 8:10625–10669
4. Barreca D, Carraro G, Gasparotto A, Maccato C, Altantset T, Sada C, Kaunisto K, Ruoko T-P, Bals S (2017) Vapor phase fabrication of nanoheterostructures based on ZnO for photoelectrochemical water splitting. Adv Mater Interfaces 4:1700161
5. Hernández S, Hidalgo D, Sacco A, Chiiodoni A, Lamberti A, Cauda V, Tresso E, Saracco G (2015) Comparison of photocatalytic and transport properties of TiO2 and ZnO nanostructures for solar-driven water splitting. Phys Chem Chem Phys 17:7775–7786
6. Zamiri R, Abbastabar Ahangar H, Tolbadi DM, Rebelo a, Seabra MP, Shaban M, Ferreira JMF (2014) Fabricating and characterising ZnO–ZnS–Ag2 S ternary nanostructures with efficient solar-light photocatalytic activity. Phys Chem Chem Phys 16:22418–22425
7. Zhang H, Chen G, Bahnmann DW (2009) Photoelectrocatalytic materials for environmental applications. J Mater Chem 19:5089
8. Ma L, Fan H, Fu K, Lei S, Hu Q, Huang H, He G (2017) Protonation of graphitic carbon nitride (g-C3N4) for an electrostatically self-assembling carbon@g-C3N4 core–shell nanostructure toward high hydrogen evolution. ACS Sustain Chem Eng 5:7093–7103
9. Valenti M, Jonsson MP, Biskos G, Schmidt-Ott A, Smith WA (2016) Plasma- monic nanoparticle-semiconductor composites for efficient solar water splitting. J Mater Chem A 4:17891–17912
10. Tian H, Fan H, Ma J, Liu Z, Ma L, Lei S, Fang J, Long C (2018) Pt-decorated zinc oxide nanorod arrays with graphitic carbon nitride nanosheets for highly efficient dual-functional gas sensing. J Hazard Mater 341:102–111
11. Li X, Chen W, Zhang S, Wu Z, Wang P, Xu Z, Chen H, Yin W, Zhong H, Lin S (2015) 18.5% efficient graphene/GaAs van der Waals heterostructure solar cell. Nano Energy 16:310–319
12. Zhang J, Wang T, Pohl D, Lellinghaus B, Dong R, Liu S, Zhuang X, Feng X (2016) Interface engineering of MoS2/Ni3S2 heterostructures for highly enhanced electrochemical overall-water-splitting activity. Angew Chem Int Ed 55:8114–8119
13. Ishihara H, Kannarpady GK, Khedir KR, Woo J, Trigwell S, Brins AS (2011) A novel tungsten trioxide (WO3)/ITO porous nanocomposite for enhanced photo-catalytic water splitting. Phys Chem Chem Phys 13:19553–19560
14. Zhou R, Lin S, Zong H, Huang T, Li F, Pan J, Cui J (2017) Functional surface coating on cellulose flexible substrates with improved water-resistant and antimicrobial properties by use of ZnO nanoparticles. J Nanomater 2017:1–9
15. Liu Q, Gao F, Wu F, Tian W, Li L (2015) Interface reacted ZnFe2O4 on α-Fe2O3 nanorays for largely improved photoelectrochemical activity. RSC Adv 5:79440–79446
16. Bora DK, Braun A (2014) Solution processed transparent nanoparticulate ZnO thin film electrode for photoelectrochemical water oxidation. RSC Adv 4:23562–23570
17. Nga YH, Iwase A, Kudo A, Amal R (2010) Reducing graphene oxide on a visible-light BiVO4 photocatalyst for an enhanced photoelectrochemical water splitting. J Phys Chem Lett 1:2607–2612
18. Yang JL, An SJ, Park WJ, Yi GC, Choi W (2004) Photocatalysis using ZnO thin films and nanoneedles grown by metal–organic chemical vapor deposition. Adv Mater 16:1661–1664
19. Kamat PV (2007) Meeting the clean energy demand: nanostructure architectures for solar energy conversion. J Phys Chem C 111:2834–2860
20. Willander M, Nur O, Zhao QX, Yang LI, Lorenz M, Cao BQ, Pérez JZ, Czekalla C, Zimmermann G, Grundmann M, Bakin A, Behrends A, Al-Suleiman M, El-Shaer A, Moffat AC, Postels B, Waag A, Boukos N, Travlos A, Kwock HS, Guinard J, Dang DLS (2009) Zinc oxide nanorod based photonic devices: recent progress in growth, light emitting diodes and lasers. Nanotechnology 20:332001
21. Moussa H, Girot E, Mozet K, Alem H, Medjahdi G, Schneider R (2016) ZnO rods/reduced graphene oxide composites prepared via a solvothermal reaction for efficient sunlight-driven photocatalysis. Appl Catal B 185:11–21
22. Paula CHR, Andrade Neto NF, Garcia LMF, Nascimento RM, Paskocimas CA, Bomio MRD, Motta FV (2019) Increased degradation capacity of methylene blue dye using Mg-doped ZnO nanoparticles decorated by Ag 0 nanoparticles. J Electron Mater 48:3017–3025
23. Kumar SG, Rao KSRK (2014) Zinc oxide based photocatalysis: tailoring surface-bulk structure and related interfacial charge carrier dynamics for better environmental applications. RSC Adv 5:3306–3351
24. Adam RE, Pirhashemi M, Elhag S, Liu X, Habibi-Yangjeh A, Willander M, Nur O (2019) ZnO/Ag/Ag2WO4 photoelectrodes with plasmonic behavior for enhanced photoelectrochemical water oxidation. RSC Adv 9:8271–8279
25. Pirhashemi M, Elhag S, Adam RE, Habibi-Yangjeh A, Liu X, Willander M, Nur O (2019) n–n ZnO–Ag2CrO4 heterojunction photoelectrodes with enhanced visible-light photoelectrochemical properties. RSC Adv 9:7992–8001
26. Etacheri V, Roshan R, Kumar V (2012) Mg-doped ZnO nanoparticles for efficient sunlight-driven photocatalysis. ACS Appl Mater Interfaces 4:2717–2725
27. Alnoor H, Pozna G, Khranovskyy V, Liu X, Iandolo D, Willander M, Nur O (2016) Influence of ZnO seed layer precursor molar ratio on the density of interface defects in low temperature aqueous chemically synthesized ZnO nanorods/GaN light-emitting diodes. J Appl Phys 119:165702
28. Pirhashemi M, Habibi-Yangjeh A (2017) Ultrasonic-assisted preparation of plasmonic ZnO/Ag/Ag2WO4 nanocomposites with high visible-light photocatalytic performance for degradation of organic pollutants. J Colloid Interface Sci 491:216–229
29. Dai K, Lv J, Lu L, Liang C, Geng L, Zhu G (2016) A facile fabrication of plasmonic g-C3N4/Ag2WO4 ternary heterojunction visible-light photocatalyst. Mater Chem Phys 177:529–537
30. Smykalla L, Shukryna P, Korb M, Lang H, Hietschold M (2015) Surface-confined 2D polymerization of a brominated copper-tetraphenylporphyrin on Au(111). Nanoscale 7:4234–4241
31. Zhang H, Wang G, Chen D, Lv X, Li J (2008) Tuning photoelectrochemical performances of Ag–TiO2 nanocomposites via reduction/oxidation of Ag. Chem Mater 20:6543–6549
32. Gelderman K, Lee L, Donne SW (2007) Flat-band potential of a semiconductor: using the Mott-Schottky equation. J Chem Educ 84:685–688

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.