

Ni composite electrodes for hydrogen generation: Activation of Nb-based semiconductors

Esteban A. Franceschini^{a,b,*}, Victoria Benavente Llorente^b, Anabel E. Lanterna^{a,c,*}

^a*Department of Chemistry and Biomolecular Sciences, Centre for Advanced Materials Research (CAMaR), University of Ottawa, Ottawa, Canada*

^b*INFIQC, CONICET, Departamento de Fisico-Quimica, Facultad de Ciencias Quimicas, Universidad Nacional de Córdoba, Córdoba, Argentina*

^c*Current affiliation: School of Chemistry, University of Nottingham, University Park, Nottingham, NG7 2RD, UK*

Abstract

The hydrogen evolution reaction depends on the accumulation of electrons on the catalytic center to enable the two-electron processes involved in water reduction. This work reports on the modification of inexpensive nickel (Ni) composite electrodes with engineered semiconductor heterojunctions based on earth-abundant transition metals that show superior hydrogen generation activity in the presence of a non-toxic electrolyte (K_2CO_3). This is, small amounts of cobalt (Co) or copper (Cu) oxides can improve the reactivity of composite electrodes formed by deposition of titanium (Ti) or niobium (Nb) semiconductors onto Ni surfaces. In general, modified Nb-based semiconductors show better performance and their enhanced activity can be understood in terms of modified surface potentials upon formation of semiconductor-Ni heterojunctions. Photoelectrochemical activity can be detected in the presence of Cu oxides, where hydrogen generation onset potential is reduced under UV-Vis light irradiation. The study demonstrates that small composition changes can greatly affect the activity of Nb-based Ni composite electrodes, showing exciting new applications for Nb-based materials.

Keywords: Hydrogen Generation, Electrochemistry, Semiconductors, Nickel, Niobium Oxide

Highlights

- Small amounts of Co or Cu species greatly affect Nb-based Ni composite electrodes activity.

*Corresponding authors

Email addresses: esteban.franceschini@mi.unc.edu.ar (Esteban A. Franceschini), anabel.lanterna@nottingham.ac.uk (Anabel E. Lanterna)

- Similar Cu loadings suggest electrode enhanced activity depends on the semiconductor.
- Cu oxides-Nb phosphate heterojunction shows the highest catalytic activity.
- Cu species induce slight photoelectrochemical effects.

1. Introduction

The use of hydrogen as an alternative to fossil fuels has gained attention as a way to eliminate the emission of greenhouse gases during combustion. However, current industrial production of hydrogen relies on the use of natural gas generating significant carbon emissions. Since the first report on the photoelectrochemical generation of hydrogen from water[1], many efforts have been focused on the production of “clean” hydrogen using renewable energy sources with zero net carbon emissions. Nevertheless, the price of the green electricity used in the electrolysis process is still high to allow for technology transfer [2, 3, 4]. Therefore, the development of inexpensive electrocatalytic systems with high hydrogen generation efficiency is of paramount importance to develop a sustainable future that can rely on the use of solar energy storage to supply the global energy demands.

The design of electrodes that can improve the hydrogen production whereas using earth-abundant and inexpensive materials can contribute to the advance of H₂ technologies in real applications [5]. In this regard, alkaline electrolyzers using Ni electrodes are an appealing method for large-scale production of H₂ [6, 7, 8, 9, 10, 11]. Ni electrodes usually present low catalytic activity and fast deactivation processes that generate unreactive surface species (hydrides, hydroxides, and other oxygenated species [12, 13]). However, modification of the Ni surface can extend the electrode lifetime, thereby improving the efficiency of the hydrogen evolution reaction (HER). For instance, we have recently shown that the modification of Ni electrodes with n-type semiconductors (SC) such as TiO₂ and Nb₂O₅, using Nafion as a binder, can improve the electron flux into Ni electrocathodes under dark [14, 15] or under photochemical conditions [16]. One of the advantages of using semiconductors to modify the electrochemical behaviour of electrodes is their perceived ability to form rectifying junctions by simple structural modifications [17, 18, 19, 20, 21]. Different semiconductor heterojunctions can be engineered [22, 23] to lower the Schottky barrier formed with the Ni electrode[24]. Based on Schottky’s theory, the inclusion of a third metal such as cobalt (Co) or copper (Cu) could generate an electron flow that improves even further the interaction between nickel and the oxides, and that could improve the catalytic activity of the SC-Ni electrodes [25, 26, 27]. Additionally, the use of multiple band gap electrodes to capture a greater fraction of the solar light has also motivated to engineer multi-electron accumulation systems[28, 29, 30, 31]. Such systems can benefit from the unidirectional electron flow that accumulates charges on the catalytic sites, favouring the two-electron reaction that generates H₂ from water.

Here we show that simple modification of semiconductor heterostructures with earth abundant metals (M@SC) can enhance Ni electrodes efficiency up to 3 orders of magnitude when comparing to electrodes modified with the bare semiconductor (SC-Ni). In particular, we have used copper or cobalt oxides to decorate the surface of transition metal semiconductor —i.e., TiO_2 , Nb_2O_5 , and $\text{Nb}_3(\text{PO}_4)_5$ —which have previously shown enhanced activity towards HER under alkaline conditions using Ni electrodes[14, 15].

2. Experimental

2.1. Materials and Instruments

Ni plates (99.99%) were purchased from Cordes SA Argentina. Nafion and 2-propanol were purchased from Sigma and used as received. DI H_2O was obtained by purification of deionized water using a Thermo Scientific™ Barnstead™ GenPure™ water purification system (conductivity = 18.2 $\text{M}\Omega\text{-cm}$) and TiO_2 P25 was purchased from Univar Canada. Niobium oxide (Nb_2O_5) and niobium phosphate ($\text{Nb}_3(\text{PO}_4)_5$) were generously provided by CBMM (Companhia Brasileira de Metalurgia e Mineração) and used as received. $\text{CuCl}_2\cdot 2\text{H}_2\text{O}$ and $\text{Co}(\text{NO}_3)_2\cdot 6\text{H}_2\text{O}$ were obtained from Aldrich, I-2959 from BASF and I-907 from CIBA-GEIGY.

X-ray powder diffraction (XRD) patterns were recorded on a PANalytical X’Pert PRO diffractometer operating at 40 kV and 40 mA using $\lambda = 1.5406$ Å Cu $K\alpha$ radiation employing in the θ - 2θ Bragg-Brentano geometry. All measurements were carried out at room temperature. The 2θ range used was 10 - 90°, with 0.02° steps and counting time of 5 s per step. The Rietveld method was used to refine the crystal structure employing the FullProf program [32]. A pseudo-Voigt shape function was used to fit the experimental data. The data refined were lattice parameters, atomic positions, isotropic thermal parameters, peak shape, and occupation factors.

Raman spectra of the electrode surface were acquired with a LABRAM-HR 800, HORIBA Jobin-Yvon Raman microscope with a 100x objective lens (NA = 0.9). The laser excitation used was 632.8 nm at 3.5 mW. The instrument allows recording spectra with a resolution of 1.5 cm^{-1} . The illuminated area in all Raman experiments was 1.0 μm^2 . At least five different zones on each sample were analyzed to confirm the homogeneity of the catalysts.

AIST-NT Scanning Probe Microscope (SPM) SmartSPM-1000 was utilized to record topography and non-contact potential measurements using Au-coated tips purchased from MikroMasch with a radius <35 nm, a stiffness of 1.8–13 N m^{-1} , and a resonant frequency of 110 - 220 kHz. All measurements were performed in non-contact AFM mode and single pass Kelvin Probe Force Microscopy (KPFM) in the dark or under visible light irradiation. The 5 x 5 μm area of study was scanned at 1 $\mu\text{m s}^{-1}$. For each scan line, the height profile was recorded in the first pass and then the tip was lifted 10 nm away from the surface and the potential offset was measured.

Scanning electron microscopy (SEM) images were obtained using a JEOL JSM-1600 SE microscope working at an accelerating voltage of 2 kV.

Electrochemical measurements were carried out using the Parstat 2273 advanced electrochemical system in a conventional three-electrode electrochemical cell. A Pt wire was used as counter electrode and saturated calomel electrode (SCE) as reference electrode (0.243 V vs. RHE).

2.2. *Synthesis of semiconductor heterostructures*

The preparation of the SC heterostructures used in this work (denoted as M@SC) was performed using a reported methodology[33, 34]. Briefly, Cu@SC heterostructures were prepared by mixing 21.2 mg of $\text{CuCl}_2 \cdot 2\text{H}_2\text{O}$ with 160 mg of the corresponding SC and 56 mg of Irgacure-2959 (I-2959) in 100 mL of DI Water. The mixture was purged with argon, sonicated for 15 min and irradiated for 5 h under UVA light (Fig. S1). The resulting slurry was filtered through nanopore filter and dried overnight under vacuum. Similarly, Co@SC heterostructures were obtained using 41.6 mg of $\text{Co}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$, 160 mg of SC and 78 mg of I-907 mixed together in 150 mL of dry acetonitrile. The mass of the Cu and Co precursors were selected to keep the number of moles of the metals constant during the synthesis. Metal loading was determined by Inductively Coupled Plasma Optical Emission Spectroscopy (ICP-OES), using an Agilent Vista Pro ICP Emission Spectrometer. Samples (10 mg) were prepared in triplicates by digestion in aqua regia. Further dilutions were prepared and analysed following the Cu 324.754 nm emission line.

2.3. *Electrodes Preparation*

The nickel supports were prepared as previously reported [14]. Initially they were degreased with acetone, and before applying the catalyst the supports were peeled by successively immersing them in 15% HCl and 1M KOH for one minute. The electrodes were prepared by drop casting method [35, 36]. Briefly, 5 mg of catalyst were dispersed in 30 mg of DI H_2O , 300 mg of 2-propanol and 15 mg of 5 wt% Nafion solution; and sonicated for 30 min. Then, 350 μL of the Nafion-based ink were pipetted onto a polished (#600 sandpaper) square-shaped Ni electrode (1 cm^2) and air-dried at 60 $^\circ\text{C}$ temperature.

2.4. *Electrochemical Measurements*

The (photo)electrochemical generation of hydrogen and the electrochemical characterization – cyclic voltammetry (CV), open-circuit potential (OCP), and electrochemical impedance spectroscopy (EIS) – of the electrodes was carried out in a conventional three-electrodes electrochemical cell using a home-made photoreactor previously described [14] (See Fig. S2) with a 0.5 M K_2CO_3 solution (pH = 11). The electrolyte was chosen as a non-toxic alternative to common NaOH or KOH. All solutions were prepared in Milli-Q water and deoxygenated by bubbling high purity nitrogen. Temperature inside the electrochemical cell was kept at 298 K ($\pm 2^\circ\text{C}$).

2.4.1. Open Circuit Potential (OCP)

Open circuit potential measurements were performed using a pre-built method consisting on applying a close-to-zero current density and measuring the transient potential with time. The OCP experiments were performed in the dark and under a combination of UVA and Vis light irradiation (Fig. S3).

2.4.2. Electrochemical Impedance spectroscopy (EIS)

EIS experiments were carried out using a frequency range between 10 mHz and 200 kHz with a 10 mV bias potential at different electrode potentials. The potentials selected corresponds with Ni open-circuit potential (OCP, -0.36 V vs SCE), HER onset potential (OP) over pure Ni (-1.1 V vs SCE) , -1.2 V vs. SCE, and -1.4 V vs. SCE in order to cover the potential range where the HER occurs [37, 38]. The ZView 3.3 software (Scribner Associates, Inc.) was used to fit the measured data with different equivalent circuit models.

3. Results and discussion

3.1. Characterization of M@SC-Ni electrodes

Ni electrodes were modified using different metal-semiconductor heterostructures denoted as M@SC, where M represents the transition metal used (*i.e.*, Cu or Co) and SC the corresponding semiconductors. Noteworthy, the supported M@SCs act as modifying agents enhancing the catalytic activity of the Ni electrodes. For simplicity, the semiconductors TiO_2 , Nb_2O_5 and $\text{Nb}_3(\text{PO}_4)_5$ are denoted as TiO, NbO, and NbP, respectively. M@SC heterostructures were prepared following photochemical deposition of the metal precursors onto the bare SC as previously described (Fig. S1).[33] Metal loading was determined by ICP-OES as described in the experimental section and shown in Table 1. M@TiO heterostructures bear the lowest metal loadings with 0.89 and 1.22 wt% of Cu and Co, respectively, while M@NbP and M@NbO support higher metal content; suggesting a preferential photodeposition of metals onto Nb-based semiconductors.

The chemical nature and surface composition of the M@SC heterostructures was determined by X-ray Photoelectron Spectroscopy (XPS). HR-XPS spectra of the metals loaded onto the three different SCs are shown in Fig. 1 and Fig. S4 - S5. Different Cu species are formed depending on the nature of the SC, in agreement with previous reports [33]. For Cu@NbO and Cu@NbP heterostructures, two Cu $2p_{3/2}$ signals, centered at 932.8 eV and 934.9 eV can be clearly distinguished and are assigned to Cu(I) and Cu(II), respectively. The presence of the Cu(II) satellites further indicate the presence of a mixture of Cu(I)/Cu(II) species. It is worth noting that the amount of Cu(I) species present in Cu@NbO (*ca.* 64%) is slightly higher than in Cu@NbP (*ca.* 54%), which could be related to differences in the surface chemistry of the semiconductor affecting the oxidation state of the metal. In contrast, the Cu 2p HR XPS spectrum for Cu@TiO presents only one Cu $2p_{3/2}$ peak centered at 930.1 eV, which can be attributed to the presence of Cu(I) species —although Cu(0) species cannot be ruled out.

Table 1: Composition of M@SC heterostructures and HR-XPS analysis.

M@SC	^a M (wt%)	^b M 2p _{3/2} (eV)	^b M(X)	^b Nb or Ti 3d _{5/2} (eV)
Cu@TiO	0.89	930.1	Cu(I)	458.4
Cu@NbO	4.56	932.8 & 934.9	Cu(I/II)	207.2
Cu@NbP	4.64	932.8 & 934.9	Cu(I/II)	207.6
Co@TiO	1.22	781.1	Co(II)	458.7
Co@NbO	1.7	781.2	Co(II)	207.0
Co@NbP	3.03	781.3	Co(II)	207.3

^a ICP-OES analysis. ^b Binding energies obtained respectively from Co 2p, Cu 2p, Nb 3d and Ti 3d regions.

Furthermore, no satellite features are found, indicating the absence of Cu (II) species in Cu@TiO. In the case of Co@SC, Co 2p HR XPS spectra show the same features for the three SC, with signals centered at 781.1 eV and 796.9 eV corresponding to Co 2p_{3/2} and Co 2p_{1/2}, respectively. All samples present a spin-orbit coupling ($\Delta_{\text{Co } 2p_{(1/2-3/2)}}$) of 15.7 eV, characteristic of Co (II) species [39]. Furthermore, satellites features of Co(II) are identified. Therefore, CoO species are likely to be found in the surface of all modified SC.

The optical properties of the materials were measured by means of diffuse reflectance (DR) spectroscopy, Fig. 2. All samples exhibit distinct absorption in the UV region, corresponding to the band-gap absorption of TiO, NbO, and NbP components. After the Cu or Co modification of SC, an increased absorption in the visible region is noticed for all samples. As a result, the optical band-gap energy, obtained from Tauc plots (Fig. S7 and Table 2), slightly decreased in relation to the corresponding SC. Therefore, the absorption edge is shifted for M@SC heterostructures to longer wavelengths, which could facilitate the generation of charge carriers upon visible light absorption. Using the theoretical electronegativity of the SC[40], one could roughly estimate the energy of the band edges[41], which suggest the NbO and NbP families are better suited for formation of H₂. This has been recently demonstrated in literature [14] where nickel catalysts modified with NbO and NbP exhibit higher catalytic activity compared to those modified with TiO, and can be seen in Fig. 3 and Fig. S8.

After the synthesis and characterisation of the M@SC, the new electrodes (M@SC-Ni) were prepared by deposition of the M@SC heterostructures onto clean Ni electrodes employing a widely used ink strategy,[44] utilizing Nafion as ionomer binder (See Fig. 3A). Nafion has been the preferred choice to achieve high-performance in fuel cells[45] and electrolyzers[46] due to its stability and great ion transport properties[47]. PXRD analysis (obtained from Rietveld refinement using Fullprof software) suggests TiO is composed by a mixture of anatase (A) and rutile (R, (13 ± 4) %), NbO contains orthorombic (O) and monoclinic (M, (24 ± 5)%) crystalline structures, and NbP is amorphous. [14] Also, crystal sizes obtained from the PXRD patterns suggest TiO particles (A: 20 nm and R: 28 nm) are smaller than the NbO particles (O: 50 nm, M: 48 nm). XRD patterns of the prepared electrodes show characteristic reflections of faced centered cubic Ni corresponding to the substrate (See Fig. S6), and the

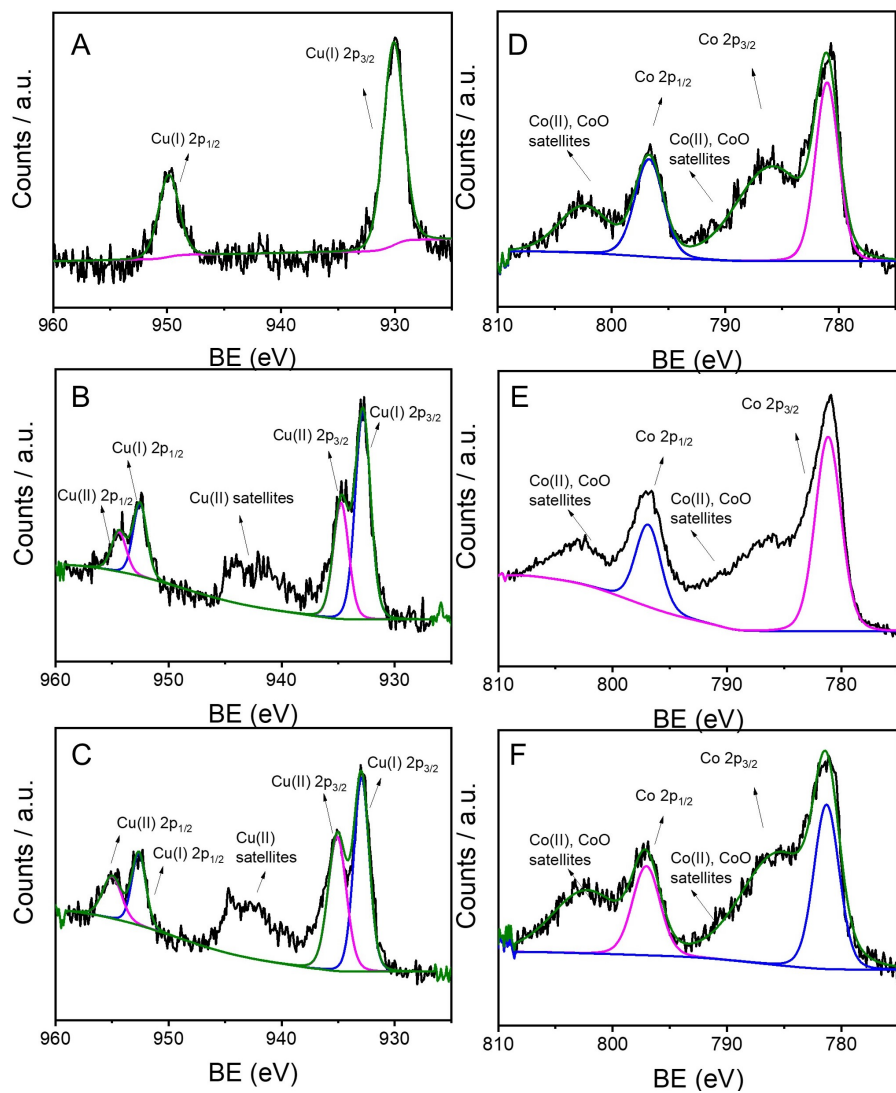


Figure 1: Cu 2p and Co 2p HR XPS spectra of A) Cu@TiO, B) Cu@NbO, C) Cu@NbP, D) Co@TiO, E) Co@NbO, and F) Co@NbP.

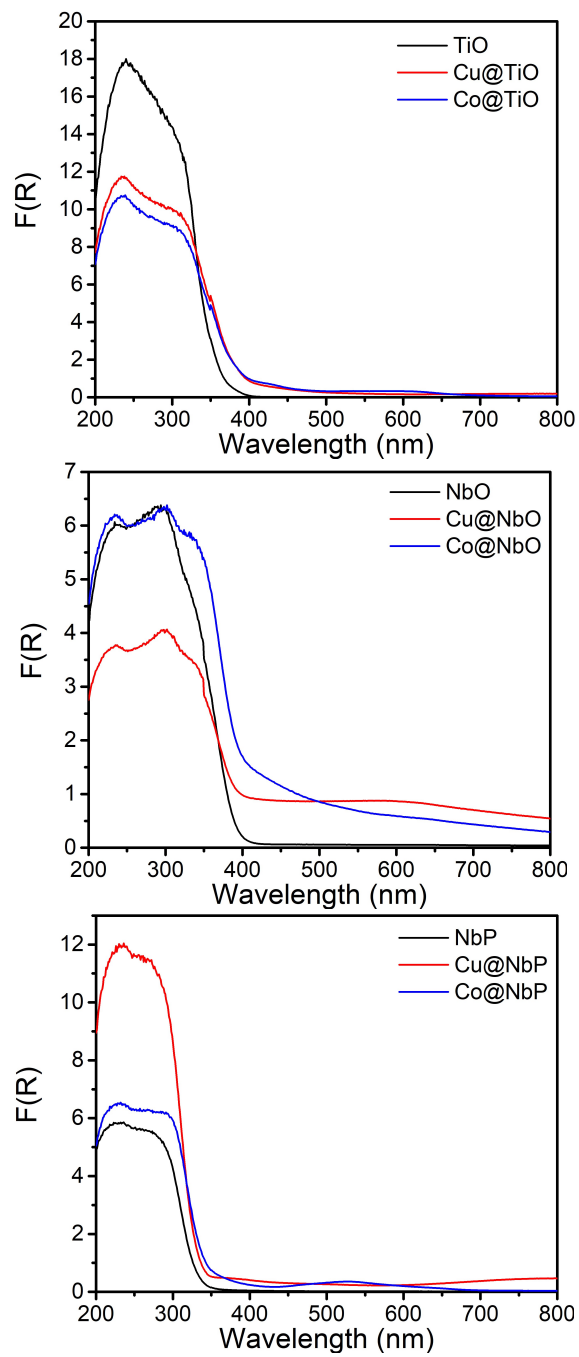


Figure 2: Diffuse reflectance spectra of the SC and their corresponding M@SC composites.

Table 2: Optical band gaps (BG) calculated using Tauc plots (Fig. S7) and valence band (VB) and conduction band (CB) edge energies in eV calculated for each SC and M@SC heterostructures.^a

M@SC	Optical BG	χ_{SC}^b	E_{VB}^c	E_{CB}^c
[14]TiO	3.19	5.81	2.91	-0.28
Cu@TiO	2.87	5.44	2.38	-0.49
Co@TiO	2.83	5.39	2.3	-0.53
[14]NbO	2.99	6.29	3.29	0.30
Cu@NbO	2.73	6.03	2.89	0.16
Co@NbO	2.74	6	2.87	0.13
[14]NbP	3.5	6.68	3.93	0.43
Cu@NbP	3.47	6.59	3.83	0.36
Co@NbP	3.35	6.58	3.76	0.41

^a Electronegativity of the semiconductors (SC) calculated as the geometric average of the absolute electronegativity of the constituent atoms using the electronegativity of each element from literature [40]. ^b Theoretical Ni Fermi level: 0.85 V vs NHE (Work Function of Ni(111): 5.35 eV [42]). ^c Calculated as [41, 43]: $E_{VB} = \chi_{SC} - E_e + 0.5E_g$ and $E_{CB} = E_{VB} - E_g$.

SC preserving their crystalline composition. No reflections of Co or Cu oxides were identified in the diffractograms, potentially due to the low metal loading on each M@SC heterostructure (See Table 1).

3.2. Photoelectrochemical characterization of M@SC-Ni electrodes

3.2.1. Cyclic Voltammetry

Cyclic voltammograms (CVs) were recorded in the range from -1.5 to 0.1 V (vs. SCE) at a scan rate of 5 mVs⁻¹ until stabilization, except for the Co@TiO and Cu@NbP catalyst where a reduced scale (-1.5 to -0.3 V vs. SCE) was used to avoid catalyst electrochemical degradation processes observed at higher potentials. The CVs shown in Fig. 3 were performed in the dark and under UVA-Vis light irradiation (See Fig. S3 and S8) and were used to calculate the corresponding onset potentials (Table 3) for the H₂ generation reaction (see video provided within the supplementary materials). Under dark conditions, it appears as if both NbO and NbP decrease the onset potential when comparing with bare Ni electrodes (OP = -1.1 eV), Table S1; whereas TiO slightly increase the onset potential values. Interestingly, the addition of Cu or Co has different impact on the onset potential values depending on the semiconductor. Overall, these values indicate that the smallest overpotential (η) towards hydrogen evolution is obtained for the Co@NbO-Ni electrode (Table S1). This is, considering the starting onset potential for HER at -0.65 V vs SCE at pH = 11.0 (calculated from Nernst equation), Co@NbO-Ni shows an $\eta = 210$ mV.

As we have recently reported [14], the onset potential for the H₂ generation from SC-Ni electrodes under dark conditions remains unchanged under

Table 3: Exchange current density (j_0), onset potential (OP) and current density (J) obtained at -1.5 V.

	$j_0(\mu\text{Acm}^{-2})$		OP (V)		ΔOP (mV)	J (mAcm ⁻²)		ΔJ (%)
	Dark	Light	Dark	Light		Dark	Light	
TiO-Ni	<1	-1.12	-1.12		0	3.4×10^{-3}	3.7×10^{-3}	8.8
Cu@TiO-Ni	<1	-1.07	-0.95	120		1.4×10^{-3}	1.9×10^{-3}	35.7
Co@TiO-Ni	<1	-1.03	-1.03	0		8.2×10^{-3}	8.2×10^{-3}	0
NbO-Ni	4	-0.96	-0.96	0		5.2×10^{-1}	5.3×10^{-1}	3.3
Cu@NbO-Ni	3	-1.26	-1.19	70		1.6×10^{-1}	1.8×10^{-1}	11.7
Co@NbO-Ni	80	-0.86	-0.86	0		1.6×10^1	1.6×10^1	0
NbP-Ni	<1	-1.05	-1.05	0		7.0×10^{-3}	7.0×10^{-3}	0
Cu@NbP-Ni	140	-1.08	-1.04	40		1.9×10^1	2.0×10^1	4.2
Co@NbP-Ni	12	-1.18	-1.18	0		2.2×10^0	2.2×10^0	0

UVA/Vis light irradiation (Table 3). In contrast, here we show that the presence of Cu in these M@SC heterostructures changes the onset potential towards more positive values upon irradiation, exhibiting some photocurrent (Fig. S8 and Table 3). No changes on the onset potential (ΔOP) and current density (ΔJ close to zero) are observed when Cu is replaced by Co. Therefore, only Cu deposited onto the TiO, NbO and NbP increases the charge carrier generation under irradiation. As shown by XPS analysis, Cu(I) species were detected in all samples, suggesting the presence of Cu_2O in the surface of the semiconductor heterostructures. The values in Table 3 indicate the initial composition of the Cu@SC could be related to the current density changes observed upon irradiation, thus, the greatest changes are obtained for Cu@TiO, which only shows XPS signals corresponding to Cu(I) species. Although NbO and NbP present the same copper loading, Cu@NbO led to higher changes in the current comparing with Cu@NP, in agreement with the slightly higher amount of Cu(I) species found by XPS. Cu_2O is an intrinsic p-type material, presents a suitable conduction band position for water reduction, and a direct band gap of 2.0 eV [48]. Therefore, it is possible that upon UV/Vis irradiation these partly oxidized Cu species can generate photo-excited electrons and holes, driving a shift in the onset potential, and small photocurrents in the composite electrode. Note that no new oxidation peaks were detected for any of the catalysts upon irradiation, although this could be due to the low Cu loading on the M@SC. Further analyses using Bode phase plots suggest these changes are related to HER only in the case of Cu@NbO and Cu@NbP (*vide infra*).

Furthermore, the open circuit potential (OCP) values are indicative of the surface state of the electrode, particularly related to its electronic state and Fermi level [49, 50]. In the case of SC-Ni electrodes, the OCP corresponds to the Ni OCP (-0.36 V vs SCE), modified by the presence of the SC heterostructures [14, 50]. It is expected that the addition of a co-catalyst (*e.g.*, Cu or Co) would modify the OCP of the composite. Interestingly, most of the M@SC heterostructures studied here show differences in the OCP values (Table 4). This is, Cu or Co co-catalysts generate significant changes to the OCP values comparing with the corresponding TiO- and NbO-Ni based electrodes. This

suggests that the presence of these metals affects the Fermi level of the Ni electrode as suggested by Schottky’s theory [27]. The same trend is observed in the case of Cu@NbP-Ni electrodes, whereas the addition of Co does not generate significant changes comparing with the NbP-Ni electrode.

Table 4: Open Circuit Potential (OCP) measured in the dark and under UV/Vis light irradiation.

Catalyst	^a OCP (mV)	
	Dark	Light
TiO	-82	-90
Cu@TiO	-126	-127
Co@TiO	-179	-187
NbO	-109	-123
Cu@NbO	-148	-143
Co@NbO	-176	-172
NbP	-148	-143
Cu@NbP	-143	-141
Co@NbP	-148	-147

^aIllumination conditions as described in Fig. S3B.

The values of the Tafel slope obtained (See Table S1) for each M@SC-Ni electrode are around -120 mVdec^{-1} , indicating that the Volmer step is the rate-determining step during the HER [51]. In alkaline media, the Volmer step involves the transfer of an electron to a water molecule to yield an adsorbed H atom as a reaction intermediate. Fig. S9 shows the different steps of the reactions that constitutes the general mechanism of the hydrogen generation reaction [52]. In order to gain insights into the electrocatalytic properties of the prepared electrodes, the exchange current density (j_0) for each electrode was compared. Typically, Ni electrodes present an j_0 of about $2 \mu\text{Acm}^{-2}$ measured in alkaline media [53]. As shown in Table 3 the addition of Co and Cu to the TiO-Ni electrodes does not generate significant increase in the j_0 , presenting values lower than $1 \mu\text{Acm}^{-2}$. This is in agreement with the low theoretical energy values calculated for the conduction bands in Table 2. Additionally, when analyzing the current density data at -1.5 V (vs. SCE) on the CV, values between 1 and $8 \mu\text{Acm}^{-2}$ are observed for all TiO-Ni electrodes. In contrast, the addition of Cu or Co to NbO-Ni and NbP-Ni electrodes can significantly modify the exchange current density. According to the j_0 and J measured at -1.5 V (vs. SCE), Co@NbO and Cu@NbP present the best performance towards HER in alkaline media. Particularly, the addition of Co to NbO-Ni generates a 20-fold increment in the j_0 values, while Cu addition to NbP-Ni induces a 140-fold increase on j_0 values. The increase catalytic activity might be attributed to several factors, including the modification of the Fermi level. This could be particularly seen in the case of Co@TiO and Co@NbO. In contrast, Co@NbP and Cu@SC do not affect the OCP values in relation to the bare SC. In these cases, other effects could be responsible for their increased catalytic activity;

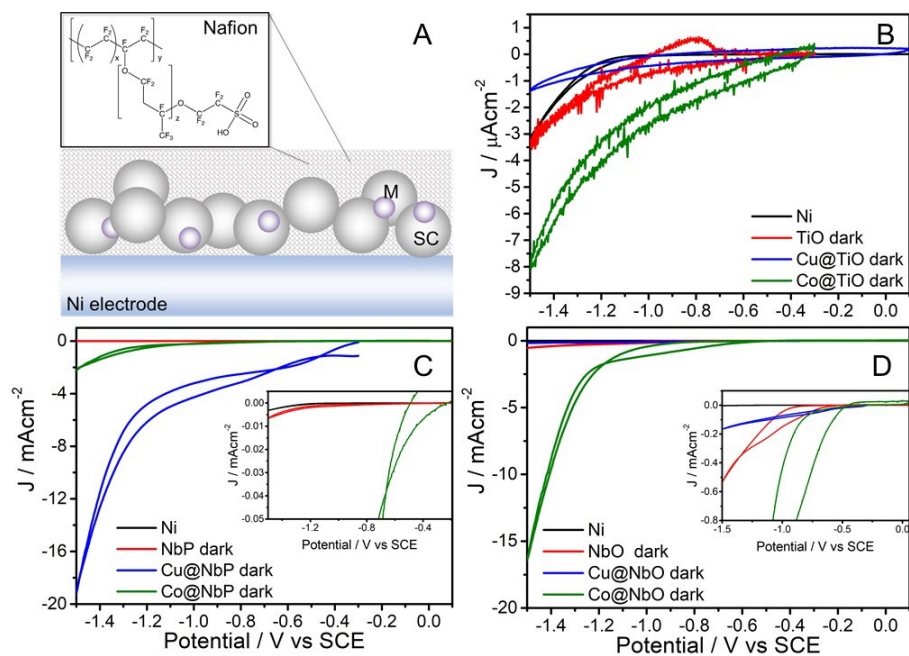


Figure 3: (A) Representative diagram of the working electrodes composed of a Ni substrate and the M@SC heterostructures embedded in a Nafion matrix. (B-D) CV obtained for each M@SC-Ni electrocatalyst under dark conditions in a 0.5 M K₂CO₃ solution (pH = 11). Insets: closer look to the onset potential of H₂ formation. Note panel B lower signal-to-noise ratio is due to low current density values detected (y -axis units in μAcm^{-2}). Panel C: CV corresponding to Cu@NbP (blue line) is out of scale and not shown in the inset.

for instance, the presence of potential active sites onto the oxidized Co and Cu species has been recently suggested in literature. Zhang *et al* have reported that highly disperse Cu_2O nanoparticles in carbon nanotubes can notably increase the catalytic activity towards HER in alkaline media [54]. Additionally, reports on CoO hybrids, supported on carbon[55] or graphene[56], have shown a distinct catalytic activity towards HER.

3.2.2. Electrochemical Impedance Spectroscopy (EIS)

To further characterize the (photo)electrocatalytic properties of the prepared electrodes a series of Electrochemical Impedance Spectroscopy (EIS) measurements were performed in the dark and under irradiation. The general equation for calculating EIS parameters for hydrogen generation predicts the formation of two semicircles in the complex plane of Nyquist plots. A low frequency semicircle, with a radius dependent on the overpotential, is related to the HER process; and a high frequency semicircle, with a constant radius, is related to the electrode surface [57]. The low-frequency semicircle radius decreases with lower charge transfer resistance. The impedance spectra can be fitted using the Armstrong and Henderson equivalent circuit (denoted as AHEC) [58, 52, 57] which allows to consider the roughness of the electrode surface and to analyze the presence of simultaneous processes at different frequencies. AHEC can also account for other processes occurring on the electrode surface, such as photoelectrochemical processes [59, 60]. Pure Ni electrodes are typically evaluated in alkaline electrolysis at the following potentials: i) the pure Ni open circuit potential (OCP) reported as -0.36 V, ii) pure Ni HER onset potential reported as -1.1 V, iii) -1.2 V, and iv) -1.4 V (vs.SCE). Therefore, these four electrode potentials were chosen to evaluate the EIS response of the prepared electrodes. Fig. 4 shows the impedance spectra in the complex plane for the different M@SC deposited onto the Ni substrate, measured at 298 K in the dark and under irradiation. Particularly, the figure shows the behavior of the catalysts at -1.2 V (vs. SCE) in the dark and under irradiation. This potential is ideal to explore the catalyst activity as there is enough hydrogen generation and low electrochemical noise—in comparison with that observed at more negative potentials. It can be seen from the Nyquist plots that the behaviors are varied and depend on both the semiconductor and the metal present in the catalyst, in agreement with what was found in the cyclic voltammetries. This is, smaller semicircles are found for Co@NbO and Cu@NbP in the dark, consistent with their low overpotentials; whereas slight differences can be seen under irradiation, this was not reflected into higher currents during the voltammetry experiments. In the case of M@TiO-Ni catalysts, no major differences are detected despite the marked increase in current density observed in the cyclic voltammetry experiments. This can be attributed to the overall low current density values observed for M@TiO catalysts and to the occurrence of surface processes unrelated to HER, as discussed below. Additionally, in most of the catalysts a process at high frequencies related to the surface structure of the catalyst is observed (Fig. S10—zoom in Fig. 4). The second semicircle observed shows dependence with the applied overpotential and is related to the HER [61]. Fur-

thermore, some catalysts show the appearance of a third semicircle at very low frequencies related to the presence of Nafion in the catalytic layer that limits the diffusion of the generated hydrogen [62, 63].

To gain a better understanding of the catalysts activity we analysed the Bode phase plots (i.e., phase shift vs. frequency plots, Fig. S11). Similar to what is found in the Nyquist plots, Bode phase plots present two phase shift maxima corresponding to the HER process (low frequency) and the surface processes (high frequency). When an electrode is subjected to a gas evolving reaction —e.g., HER —the phase shift angle maximum (at low frequency) is displaced to higher frequencies and becomes less significant at higher overpotentials. This process is associated with an increased electrode surface coverage generated by formation of bubbles that block the exposed surface[64]. Thus, the electrodes with higher HER efficiency exhibit this type of behavior. Fig. S11 shows that M@TiO-Ni catalysts have almost negligible changes as the overpotential increases, suggesting that although electrochemical processes are observed during cyclic voltammetry, the HER might be incipient and other surface processes (such as reduction of metal oxides) are predominant. Thus, it is possible that the increase current observed for Cu@TiO-Ni under irradiation is due to processes other than HER. A similar behaviour can be seen for Cu@NbO-Ni catalyst; although this catalyst shows a higher catalytic activity comparing with M@TiO-Ni catalysts. In contrast, Co@NbO-Ni catalyst shows a marked response to higher potentials, similar to that expected during HER, and in agreement with the response observed by cyclic voltammetry. In the case of the M@NbP-Ni catalysts a correlation between the dependence of the phase shift maximum with the overpotential is also observed. In particular, Cu@NbP-Ni catalyst shows a higher response towards hydrogen generation (low frequency maximum) than in the case of Co@NbP-Ni (see arrows in Fig. S11). Additionally, the phase shift angle maximum at high frequencies (related to the surface response) is more predominant for the Co@NbP-Ni catalyst. On the other hand, small fluctuations in the maximum phase shift are observed when applying light mainly at the onset potential. This fluctuation could be due to the appearance of new electrochemical processes upon irradiation. This is supported by the values obtained from the A & E equivalent circuit fitting model —where increase in the R_{ct} values does not affect the current density. Table 5 shows R_{ct} value is particularly increased for Co@NbP-Ni catalyst while j value decrease is negligible (Figure S8). Thus, the appearance of a new process related to surface irradiation can lead to errors in the fit performed with an equivalent circuit that only considers two processes. The phase shift increase could be due to the irradiation effect, which becomes noticeable at lower external potentials, while at potentials where the hydrogen evolution reaction is being externally forced, small reductions of the charge transfer resistance (R_{ct}) are observed when irradiating with UV-Vis light, as in the case of Cu@SC-Ni catalysts (See Table 5). Considering that the phase shift peak is related to charge transfer, this supports a favourable charge transfer upon irradiation [65]. The complete electrochemical impedance parameters obtained by fitting the experimental data with the Armstrong & Henderson equivalent circuit is presented in Table S2.

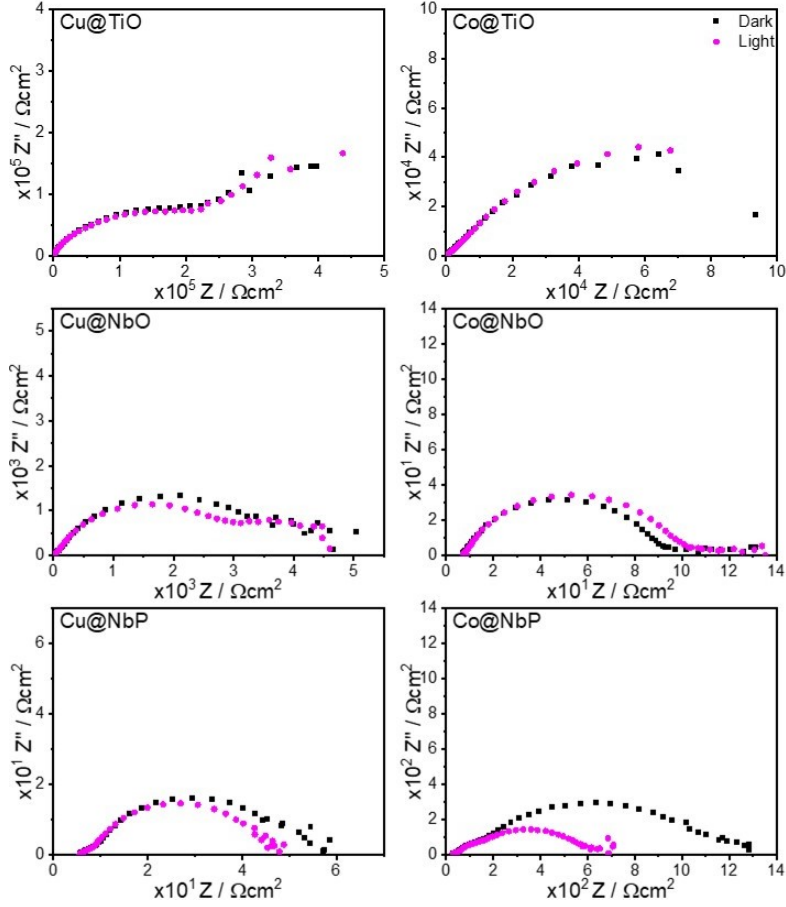


Figure 4: Nyquist plot impedance spectra in the complex plane for the HER for M@SC onto nickel electrodes on 0.5 M K_2CO_3 at 298 K and -1.2 V (vs. SCE). Measurements were performed in the dark and under UV/Vis light irradiation.

Table 5: Charge transfer resistance obtained using the AHEC model for all the catalysts under dark and irradiation

		TiO		NbP		NbO	
		Dark	Light	Dark	Light	Dark	Light
Co	$R_{ct} / \Omega.cm^2$	19422.0	19983.0	353.3	430.7	11.89	12.4
Cu	$R_{ct} / \Omega.cm^2$	132930.0	799.0	6.5	5.0	409.3	367.0

3.2.3. Kelvin probe force microscopy analysis

Kelvin probe force microscopy (KPFM) was used to characterize the impact of the Co and Cu deposition on the surface potential properties of the electrodes in the dark or under visible light irradiation (Fig. S3). In Fig. 5 a mapping of surface potential of representative electrodes under dark conditions can be found. Clearly, the deposition of the M@SC heterostructures onto the Ni substrate modifies the surface potential of the electrodes. Although the absolute work functions values are difficult to obtain under ambient conditions[66], KPFM can measure a contact potential difference (CPD) between the sample surface and the tip (V_{CPD}). The V_{CPD} of the M@SC-Ni electrodes could be used to estimate the work function from different electrodes Fig. S12) considering the Ni electrode V_{CPD} as a reference, corresponding to a work function value equals to 5.35 eV (See table 6) [67]. The analyses suggest that Cu or Co co-catalysts change the work function values of the corresponding M@SC-Ni electrodes. While V_{CPD} values for M@TiO-Ni electrodes are strongly reduced in the presence of Cu or Co, the opposite effect is found on M@NbO- and M@NbP-Ni electrodes —affecting more NbO-based electrodes.

Figs. S13-S15 show the CPD images for all materials in the dark and under irradiation. From the potentials obtained, Table 6 was made. Using the information from Table 6, the work function of each electrode can be estimated. The energy diagram shown in Fig. S12 takes into consideration the Fermi level ($E_f = -\phi$) of the prepared electrodes and bare Ni. The Fermi level of modified electrodes, estimated from KPFM measurements, is shifted towards more negative potentials compared to Ni, which thermodynamically favours the HER. Although the surface energy band diagram may change drastically upon contact with the electrolyte, the estimated values obtained from KPFM and OCP studies indicate that the deposition of Cu and Co onto the SC can alter the surface potential properties of the modified Ni electrocatalysts. On the other hand, under stationary irradiation using visible light (no direct excitation of the SC), Cu@TiO heterostructures presented the highest shift in the V_{CPD} values, indicating a surface potential rearrangement upon irradiation, in agreement with the higher response found in cyclic voltammetry.

4. Conclusions

We have presented a strategy for the preparation of hybrid Nickel electrodes with semiconductor heterostructures using Nafion as a binder. As previously showed, the presence of Nb-based semiconductors (NbO and NbP) can increase the production of hydrogen drastically. Here we showed that deposition of Co or Cu oxides onto these Nb-based semiconductors can increase the current density even more, suggesting semiconductor heterostructures can boost the HER efficiency of Nickel composite electrodes. Remarkably, the Ni composite electrodes composed by Cu and Nb semiconductor heterostructures (Cu@NbSC-Ni) show enhanced activity under photoelectrochemical conditions, even when the corresponding SC-Ni electrode is unaltered under the same irradiation conditions.

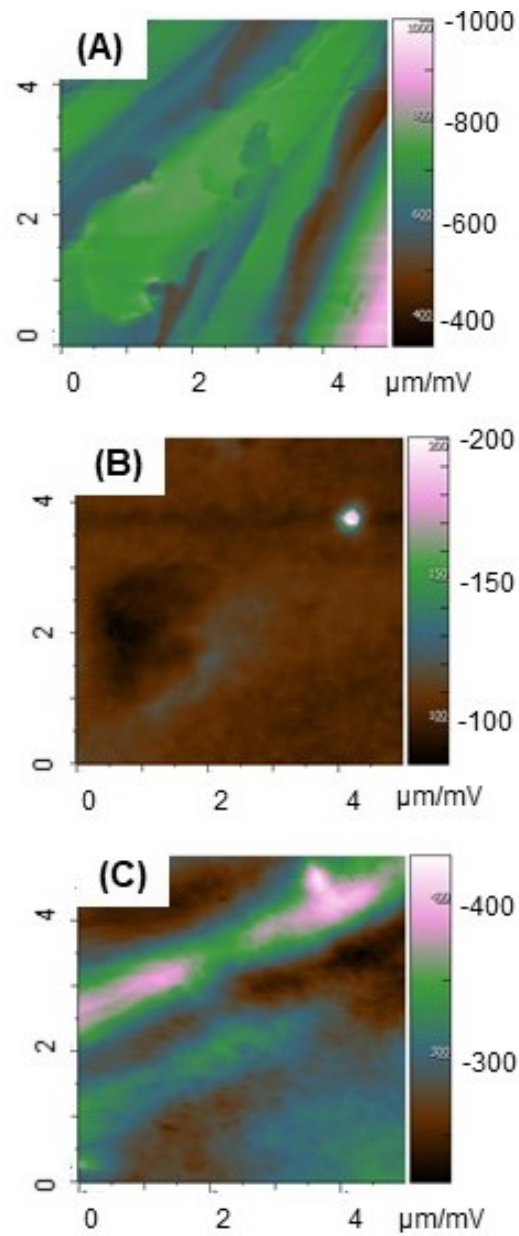


Figure 5: Mapping representing contrast in surface potential of A) Ni, B) Cu@NbO-Ni, C) Co@NbO-Ni electrodes.

Table 6: Contact Potential Difference V_{CPD} and theoretical work function (ϕ).

Catalyst	${}^aV_{CPD}$ (mV)		ϕ (eV)	
	Dark	Light	Dark	Light
TiO	774.1	769.8	5.53	5.53
Cu@TiO	178.7	230.7	4.94	4.99
Co@TiO	-73.9	-73.1	4.68	4.68
NbO	9.9	9.8	4.77	4.77
Cu@NbO	105.3	110	4.86	4.87
Co@NbO	301.3	303.9	5.06	5.06
NbP	133.7	136.4	4.89	4.89
Cu@NbP	185.6	190	4.94	4.95
Co@NbP	242	252.4	5.00	5.01

^aDetermined by KPFM, illumination conditions described in Fig. S3C. V_{CPD} values for Ni plate were 586.9 mV and 583.0 mV in the dark and under illumination respectively.

Nevertheless, EIS studies suggest only the enhanced activity of Cu@NbP-Ni electrodes is related to hydrogen generation, whereas other processes, possibly surface processes, could dominate the increase photocurrent seen for NbO and TiO counterparts. In this sense, it is possible that the amorphous nature of NbP contributes to more favourable Cu-SC surface interactions. Studying differences in the surface and electronic interactions between Cu and the two NbSC could give more insights into this apparent better performance. Based on the Schottky barrier theory, the inclusion of a third metal such as Co or Cu could generate an electron flow that enhances the interaction between the nickel and the oxides improving the response of the catalytic activity of NbSC-Ni heterojunctions. This work shows small amounts of earth-abundant co-catalysts can greatly affect the activity of Nb-based Ni composite electrodes significantly improving hydrogen generation processes under laboratory conditions. This study constitutes the first approach to use this type of materials for hydrogen photoelectrogeneration. Noteworthy, the materials used can be synthesised in large scale —TiO₂ is widely used in different industries and niobium semiconductors have recently attracted attention for potential uses in batteries [68]. Yet, the use of these materials as modifying agents for photocathodes has been little explored in literature. Overall, the materials have interesting properties to be considered for applications in photoelectrochemical generation of hydrogen and open an opportunity for the use of new materials in the production of green hydrogen.

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Supporting information

Additional experimental details and characterization data for all new composites.

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