Solar-Driven Water Splitting at 13.8% Solar-to-Hydrogen Efficiency by an Earth-Abundant Electrolyzer

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ABSTRACT: We present the synthesis and characterization of an efficient and low cost solar-driven electrolyzer consisting of Earth-abundant materials. The trimetallic NiFeMo electrocatalyst takes the shape of nanometer-sized flakes anchored to a fully carbon-based current collector comprising a nitrogen-doped carbon nanotube network, which in turn is grown on a carbon fiber paper support. This catalyst electrode contains solely Earth-abundant materials, and the carbon fiber support renders it effective despite a low metal content. Notably, a bifunctional catalyst-electrode pair exhibits a low total overpotential of 450 mV to drive a full water-splitting reaction at a current density of 10 mA cm\(^{-2}\) and a measured hydrogen Faradaic efficiency of ∼100%. We combine the catalyst-electrode pair with solution-processed perovskite solar cells to form a lightweight solar-driven water-splitting device with a high peak solar-to-fuel conversion efficiency of 13.8%.

KEYWORDS: Solar-driven electrolysis, Earth-abundant materials, Nanostructured catalyst, Perovskite solar cells, Cost analysis

INTRODUCTION

The ever-increasing energy demand, spurred by an overall global increase in living standards and a larger world population estimated to reach 10 billion people in 2060, in combination with global warming, mandates a fast transition to renewable energy sources.\(^1\) The Sun offers a continuous, albeit intermittent, source of energy, but the direct conversion of solar energy to electricity will not alone solve the energy demand, due to poorly developed electrical grids in large parts of the world and due to the fact that less than 20% of the energy consumption today is in the form of direct electricity.\(^2\) Converting solar energy to transportable hydrogen fuel by solar-driven water electrolysis or photocatalysis are promising routes to lower fossil-fuel dependence, since the stored hydrogen can be used in fuel cells that convert the chemically stored energy back to electricity, with water as the only byproduct.\(^3,4\) Since H\(_2\) has a very high gravimetric energy density, fuel cells are preferable for long distances and heavy transportation over battery-driven vehicles, and several countries are now rapidly developing an infrastructure for fueling vehicles by hydrogen.\(^9\) Besides usage as a high density energy carrier, H\(_2\) can also be used in many industrial applications, such as producing CO\(_2\)-free steel and NH\(_3\). However, around 95% of all hydrogen is today produced by reforming natural gas and coal gasification, releasing almost 1 Gt CO\(_2\) to the atmosphere every year, with the consequence being that the current hydrogen economy is far from CO\(_2\) neutral.\(^1,10\) Solar-driven water electrolysis thus represents a potentially efficient way for CO\(_2\)-free hydrogen production.

Sustainable water electrolysis requires that the anode and cathode catalysts are noble metal free and contain small amounts of other metals in order to lower system costs and facilitate recycling. Using bifunctional catalysts that work for both hydrogen evolution and oxygen evolution is also advantageous since it facilitates upscaling. In alkaline conditions, transition metal oxides, based on Ni, Fe, and Co, have been shown to exhibit low overpotential for the oxygen evolution reaction (OER), in line with the best noble metal oxides RuO\(_2\) and IrO\(_2\), but few examples of such catalysts are reported for the hydrogen evolution reaction (HER),\(^11\) although approaches such as vacancy engineering,\(^12\) bimetallic alloying, and addition of phosphides\(^13,18\) have shown to be promising strategies to lower the overpotential for metals toward HER in alkaline conditions.\(^17,18\) In recent years, the best catalyst electrodes are usually supported on
rather expensive nickel foams for achieving a high surface-area 3D network that concurrently is stable and relatively active as a catalyst itself. An alternative support is nanostructured, high surface-area carbon, as carbon is abundant, has a high conductivity, and a good ability to anchor nanocatalysts, especially if properly functionalized. However, at harsh alkaline conditions, the carbon at the anode is at risk of being oxidized, albeit this risk is lowered for crystalline carbon phases such as graphite or carbon nanotubes. Catalysts such as FeNi-layered double hydroxides feature low overpotential for OER and offer better stability than nanoparticles, since a layered structure could assist in protecting the carbon support against direct exposure to the alkaline electrolyte. Fe–Ni alloys, however, exhibit rather high overpotential for HER.

In this study, we investigate the effects of adding a third transition metal, molybdenum, to the Fe–Ni alloy. Previously, bimetallic transition metal alloys doped with a third high melting-point transition metal have shown good performance for OER, and molybdenum-based materials have in addition shown excellent performance as HER catalysts. Therefore, for further optimizing the performance of NiFe toward OER but also promote HER, molybdenum was added in the NiFe complex. Here, we report on the design of a trimetallic hybrid catalyst, which drives the water-splitting reaction at 10 mA cm⁻² at an overpotential as low as 450 mV. The low overall overpotential allows us to operate our complete water-splitting device, comprising our hybrid catalyst electrodes and two serially connected solution-processed perovskite solar cells, with a high solar-to-hydrogen conversion efficiency (STH) of 13.8%.

**RESULTS AND DISCUSSION**

To achieve our goal of an abundant electrode, we opted to use carbon paper (CP) comprising carbon fibers (with a typical diameter of 10–20 μm) as the starting material. The CP fibers allow for good anchor points for an extremely fine network of nitrogen-doped carbon nanotubes (NCNTs, typical diameter of 20–120 nm). The NCNTs were grown directly on the carbon fibers using chemical vapor deposition for achieving high adhesion and reduced contact resistance. An essential aspect for a well-coated catalyst structure is the high number of adhesion sites on the nitrogen-functionalized carbon nanotubes, which previously has been proven to be beneficial for catalysts anchoring. Furthermore, it has been shown that the introduced nitrogen defects in the NCNTs result in an increased electrical conductivity compared to pristine CNTs. This combined support material, consisting of a carbon-based NCNT/CP network, provides a relatively large surface area (70 m²/g) with good conductivity, stability, and high density of anchoring sites for the metal catalysts.

To finalize the catalyst electrode synthesis, a trimetallic NiFeMo catalyst was decorated onto the NCNT/CP support through hydrothermal synthesis (see Experimental Section for details). While NiFe hydroxides have been used as very efficient catalysts for OER previously, Mo oxides have shown near optimal hydrogen adsorption energies and are seen as promising candidates for HER in alkaline electrolytes and also as an OER catalyst. Therefore, we have added Mo as a third metal in the catalysts to improve the HER activity and still retain or improve the high OER activity of NiFe. The XPS data in Figure S1 show that the Ni:Fe:Mo ratio is 1.1:1.0:0.4 with an oxygen:metal ratio of 2.05:1.00 in agreement with a metal oxide/hydroxide phase for the catalyst.

![Figure 1](image-url) Figure 1. SEM images of the (a–c) NiFeMo-NF and (d) NiFeMo-NP catalysts. The (a) low magnification image reveals the overall catalyst electrode structure with CP fibers coated with NCNTs which in turn are decorated with the trimetallic NiFeMo catalyst. The close-up on a single CP fiber in (b) reveals the dense coating of NCNTs and a NiFeMo catalyst on the CP. The high magnification image in (c) of a single decorated NCNT on a NiFeMo-NF OER electrode reveals that the NCNT is densely covered with a nanoflake-structured NiFeMo catalyst. (d) The NiFeMo catalyst on the HER electrode has been subjected to a reduction process leading to a structural change from nanoflakes to nanoparticles, NiFeMo-NP.
and higher electrical conductivity.\textsuperscript{17,36,37} The nanoflake structures of the NiFeMo-NF catalysts are concurrently transformed into nanoparticle (NP) structures, as highlighted by the SEM images in Figure 1d. The SEM images also reveal that the catalyst particles remain firmly attached to the NCNT network with a homogeneous coverage of the NCNT/CP support, as indicated by the SEM-EDS images in Figure S2. The reduced catalyst electrode is denoted NiFeMo-NP.

To evaluate the electrochemical performance of each catalyst separately, 0.2 cm\(^2\) pieces of the electrodes were punched out and connected in three-electrode cell setups with Ag/AgCl and Pt wire as the reference and the counter electrode, respectively. The measurements were performed in 1.0 M KOH with a 2 mV s\(^{-1}\) scan speed. The performances of the individual NiFeMo-NP electrode in the HER and that of the NiFeMo-NF electrode in the OER are shown in Figure 2a and b, respectively. For comparison, the Mo-free NiFe catalysts anchored on a similar NCNT/CP support, blank NCNT/CP support as well as Pt wire (HER), and IrO\(_2\) on CP (OER) were also measured. We note that for the NCNTs/CP support, there is no background currents at the gas production potentials, which indicates that the electrode is stable and inert at these conditions. Also, here the NiFe catalyst was reduced before HER tests.

Alloying NiFe with Mo leads to a significant increase in HER activity as manifested by a lowering of the overpotential needed to reach 10 mA cm\(^{-2}\) from 250 mV for NiFe/NCNT/CP to 170 mV for NiFeMo-NP. The overpotentials for the noble metal-free hybrid catalyst are in fact only 105 mV higher than that for Pt wire at the same current density. We also note in agreement with our earlier study that the "blank" NCNT/CP support electrodes possess very poor catalytic activity for HER.\textsuperscript{17} A Tafel analysis of the voltammetry data (Figure S5a) indicates that the HER for both NiFeMo catalysts proceeds via a Volmer-Heyrovsky pathway, likely with strong hydrogen adsorption manifested by a lack of a linear Tafel region at low current densities in the Tafel plot, and therefore, the Heyrovsky reaction step would be rate determining.\textsuperscript{38,39} A low Tafel slope of around 45 mV/dec is identified at low current densities for the Pt reference electrode. At higher current densities, the Tafel slope for all materials increases to 120 mV/dec, which is a general characteristic for metallic catalyst since the hydrogen coverage is then constant.\textsuperscript{19}

The OER activity of NiFeMo-NF shows a remarkable low potential at 10 mA cm\(^{-2}\) of only 1.49 V (Figure 2b), which is superior to any of the tested reference materials including IrO\(_2\) and comparable to the state-of-the-art catalysts.\textsuperscript{11,40} The small oxidation peak near 1.4 V likely corresponds to an oxidation reaction, similar to \(\alpha\)-Ni(OH)\(_2\) to \(\gamma\)-NiOOH transformation in pure nickel samples, although the exact origin is currently unclear due to the complexity of the trimetallic alloy.\textsuperscript{41–43} It is clear that the addition of NCNTs to the CP increases the electrode surface area, which in turn renders the normalized current density larger than for both Pt and IrO\(_2\) on CP, manifesting the advantage of NCNT to boost the catalytic activity. The Tafel plot for NiFeMo-NF (Figure S5b) is harder to analyze since the oxidation peak is observed at relatively high current densities. Even though, if we extrapolate the current density, an initial Tafel slope of \(\sim30\) mV/dec at low current densities that increases to 60 mV/dec after the oxidation peak and then to 120 mV/dec can be observed, which is consistent with a metal–OOH to metal–OO\(^-\) reaction as the rate-determining step in the OER process.\textsuperscript{38}

However, the reaction mechanism for OER is normally difficult to confirm due to the high complexity. Similar behavior is seen for all tested materials, and the Tafel slopes eventually increase to greater than 300 mV/dec at larger current densities (\(\sim100\) mA cm\(^{-2}\)). In solar-driven electrolysis, these limitations are not necessarily of importance since the devices usually operate at the lower current densities produced by the solar cells, typically around 10 mA cm\(^{-2}\).

To measure the performance for complete water splitting, the two catalysts were positioned parallel to each other at 1 cm separation. In this two-electrode setup (denoted NiFeMo-NF/NiFeMo-NP), the NiFeMo-NF was the anode and the NiFeMo-NP electrode the cathode. The scanning was done in the negative direction to eliminate any oxidation peaks that might overlap with the Faradaic current at low current densities. In Figure 2c, the polarization curve shows that for the full water-splitting reaction to occur only 1.68 V is required to reach 10 mA cm\(^{-2}\), which represents a total overpotential of 450 mV for a current density of 10 mA cm\(^{-2}\).

To certify that the measured current corresponds to gaseous production, the Faradaic efficiency (\(\eta_{\text{Far}}\)) of the system was quantified by membrane-inlet mass spectrometry. A constant current density of 10 mA cm\(^{-2}\) was applied to each of the catalysts, and the produced gases were measured at certain time intervals (detailed information in Experimental Section). The resulting efficiencies are shown in Figure 2d as solid green triangles (H\(_2\)) and solid green squares (O\(_2\)), while the corresponding black markers represent the quantified amount of produced gases. The Faradaic efficiencies were near 100% for both gases within experimental error. We find that the average molar ratio of hydrogen to oxygen produced in this experiment to be 2.06:1.00, which is close to the expected 2:1 ratio. Using the 100% Faradaic efficiency and the measured total overpotential of 450 mV at 10 mA cm\(^{-2}\), we can conclude that the NiFeMo-NF/NiFeMo-NP electrode setup is able to...
produce H₂ with an electric-to-hydrogen conversion efficiency of 73.2% at a production rate corresponding to a current density of 10 mA cm⁻².

The stability of the catalysts is important for applicability in real devices, and this was investigated by applying a steady current density of 10 mA cm⁻² while monitoring the evolution of the applied potential of each catalyst electrode over a 10 h test period. We find that the catalyst electrodes are fairly stable as the combined overpotential for the NiFeMo-NF NiFeMo-NP electrode pair increases from 1.68 to 1.78 V during the first 10 h of operation (Figure S5c), which arises from an overpotential increase of 40 mV for NiFeMo-NF and 60 mV for the NiFeMo-NP, corresponding to an overall performance decrease of 6%. From other studies on similar catalysts, such a decrease in performance, at relatively low current densities and potentials, is usually related to a lowering of the active catalyst surface area due to catalyst migration and catalyst dissolution.⁴⁴

We now turn our attention to the solar-driven hydrogen production device. To target the requirements of a low-cost, lightweight, and stand-alone system, we opted to use perovskite solar cells (PSCs) for the power generation. PSCs have recently emerged as potential low-cost competitors to traditional Si-based solar cells, thanks to their solution-based fabrication using only Earth-abundant materials for their active layers and their high solar-to-electric conversion efficiency.⁴⁵⁻⁴⁷ In addition, the high output voltage of PSCs renders them simpler (i.e., fewer devices connected in series) device configurations for water-splitting operations where a working potential of at least 1.23 V (not including the catalyst overpotentials) is required.⁴⁸,⁴⁹

A schematic of the PSC device structure is presented in Figure 3a. The device consists of a mixed halide FAI:PbI₂:MABr: PbBr₃ (1:1:1:0.2:0.2 molar ratio) photoactive layer positioned in between a doped spiro-OMeTAD hole transport layer and a compact/mesoporous TiO₂ bilayer for the electron transport. The top Au cathode is in contact with the spiro-OMeTAD hole transport layer, whereas the bottom FTO anode is in contact with the TiO₂ bilayer. The FTO anode is positioned in between a doped spiro-OMeTAD hole transport layer, whereas the bottom FTO anode is in contact with the TiO₂ bilayer. The FTO anode is deposited on a glass substrate, which provides mechanical stability. All layers were deposited by solution-based processes, with the exception of the vacuum-deposited Au cathode and FTO anode. The PSC fabrication protocol is detailed in the Experimental Section.

A total of 116 different PSC devices were fabricated and characterized under AM1.5 illumination, using a shadow mask with an opening of A_PSC = 0.118 cm² to accurately define the illuminated area. Their performance is summarized in the histogram plots in Figure S6, and the narrow distributions of all of the performance metrics indicate a robust fabrication protocol. The current density—voltage (J−V) characteristic for our best performing PSC is shown in Figure 3b, and it features a short circuit current density of J_{SC} = 23.4 mA cm⁻² (average J_{SC} = 22.6 mA cm⁻²), an open circuit voltage of V_{OC} = 1.09 V (average V_{OC} = 1.05 V), and a solar-to-electric conversion efficiency of PCE = 19.4% (average PCE = 17.5%). A high fill factor FF = 76% (average FF = 73.5%) is the result of the sharp knee-like J−V trace, ensuring that a high operational voltage can be maintained when the illuminated solar cell is driving a load.

The J−V data for the PSC (Figure 3b) were matched with the J−V data of the two-terminal NiFeMo-NF NiFeMo-NP complete pair (Figure 2c) in a load line analysis (Figure 4a) in order to predict the performance of a number of connected complete solar-driven water-splitting devices. Specifically, the calculated solar-generated currents of two-, three-, and four-series-connected PSCs, with a constant total illuminated area of A_{PSC} were compared with the calculated catalyst-consuming current of one NiFeMo-NF NiFeMo-NP catalyst pair, with a varying active electrode area (A_{CAT}). The operation point for each configuration is found at the intersection between the PSC and the NiFeMo-NF NiFeMo-NP catalyst pair load lines. This analysis reveals that it is preferable to utilize a two-series connected PSC configuration for the complete solar-driven water-splitting device, because of the high V_{OC} and FF of the PSC in combination with the low overpotential of the NiFeMo-NF NiFeMo-NP catalyst pair.

From the operating points derived by our load line analysis, we can extract the operating current (I_{OP}) which in turn (through the operating current density, J_{OP}/A_{PSC}) directly reveals the expected solar-to-hydrogen conversion efficiency (STH) of our device via eq 1, as described in the Experimental Section. The calculated I_{OP}/A_{PSC} (left y-axis) and the corresponding STH (right y-axis) are summarized in Figure 4c as a function of the A_{CAT}/A_{PSC} ratio (i.e., an increase of catalyst area in comparison to the total PSC area). The data are plotted for the two-, three-, and four-series-connected PSCs. We again find that the two-series connected PSC results in the highest performance; specifically, that an I_{OP}/A_{PSC} > 11 mA cm⁻², corresponding to an STH > 13.5%, is obtained for A_{CAT}/A_{PSC} > 1. Increasing A_{CAT}/A_{PSC} to much larger values than 1 result in a very small increase of STH, but with the drawback that the material cost of the catalyst increases. With these predictions at hand, we designed and fabricated a solar-driven water-splitting device comprising a two-series PSC, with a combined A_{PSC} = 0.236 cm², and a NiFeMo-NF NiFeMo-NP catalyst—electrode pair with A_{CAT} = 2 cm², so that A_{CAT}/A_{PSC} = 8.5. This combination is marked with an open X in Figure 4c and predicts an I_{OP}/A_{PSC} = 11.2 mA cm⁻², corresponding to an STH = 13.8%.

In order to experimentally test this prediction, two high-performance PSCs were electrically connected to the NiFeMo-NF NiFeMo-NP catalyst—electrode pair, facing each other at a 4−5 mm interelectrode spacing. The catalyst electrodes were submerged into a 1 M KOH electrolyte, after which a constant AM1.5 illumination was directed at the PSCs. This activation of the two PSCs immediately switched on the generation of O₂ and H₂ gases at the catalyst electrodes by water splitting, as made visible by the bubble generation on the NiFeMo-NF...
NiFeMo-NP catalyst electrodes (a photograph of the H₂-producing water-splitting device is shown in Figure S7). The measured initial value of $I_{OP}/A_{PSC}$ for the water-splitting device is 11.2 mA cm$^{-2}$, which corresponds to an STH of 13.8%. These measured numbers are in excellent agreement with the prediction from the load line analysis and demonstrate the validity of our analysis and, most importantly, the functionality of our developed solar-driven water-splitting device. The measured STH efficiency is improved compared to all previous electrolyzers driven by low-cost PSCs that used solely Earth-abundant electrocatalysts but lower compared to the highest performing perovskite/Si tandem devices that uses noble metal electrocatalysts, which reached an initial STH efficiency of 17% STH.3,50 We note that in a longer perspective, using Earth-abundant electrocatalysts is necessary for sustainable large-scale H₂ production, both for cost-wise and environmental perspectives.55

The stability of the water-splitting device is displayed in Figure 4d. A continuous decrease of $I_{OP}/A_{PSC}$ and STH during the 10 h measurement is observed, after a very short onset period originating from the shutter of the solar test system. Since the NiFeMo-NF$|$NiFeMo-NP catalyst electrodes alone demonstrate relatively stable operation during such time period (Figure S5c and d), we turn to the PSCs for an explanation. Figure S8 compares the $I_{OP}/A_{PSC}$ of the solar-driven water-splitting device to the measured $J_{SC}$ of a pristine two-series connected PSC during a continuous AM1.5 illumination stress test at short-circuit conditions. The data clearly reveal that the measured reduction in $I_{OP}/A_{PSC}$ matches the decline of $J_{SC}$, implying that the lowered water-splitting performance with time is primarily due to the degradation of the PSCs. We note that the PSCs were operated in ambient atmosphere without encapsulation and that various efforts into stabilizing nonencapsulated PSC devices are ongoing56−66 but are outside the scope of this study.

In addition to lab-scale demonstrations, there is also value in performing relevant cost analysis on the key components early on in development, simply to point out potential limitations and to motivate topics and set targets for future development. We therefore performed a rudimentary cost analysis focusing on the material costs for fabrication of the NiFeMo-NF$|$NiFeMo-NP catalyst-electrode pair and the PSC in two different scenarios: (i) a lab-scale scenario, with actual high-waste fabrication and actual lab-scale material costs and (ii) a no-waste scenario, with an estimated no-waste fabrication and actual material costs for similar quality materials. The cost analysis for the NiFeMo-NF$|$NiFeMo-NP catalyst-electrode pair and the PSC in these two scenarios is detailed in Table S2.

For the lab-scale fabrication scenario, we use actual material volumes and costs to calculate an enormous cost of $19,100.
m² of coated area for the catalyst—electrode pair. The more interesting cost-per-active-area metric for the catalyst (CPACAT) is obtained when the cost per coated m² is multiplied with the geometric fill factor (GFF), defined as the ratio between active and coated area. With a GFF = 98% for the lab-scale catalyst, we obtain a lab-scale CPACAT = $21,500 m². Surprisingly, this high cost is mainly due to the high volume of DMF solvent used during the catalyst decoration step. For the PSC, we found an even more extreme cost of $43,400 m². Considering the low GFF of 18.7%, this results in a cost-per-active-area for the PSC (CPAPSC) of $233,000 m². This cost is mainly driven by the high cost of the glass/FTO substrates and the material-inefficient deposition of the expensive spiro-OMeTAD and Au materials.

In the no-waste scenario, the material usage is calculated as the minimum amount of material needed to produce the lab-scale device structure by assuming 100% material utilization as well as reuse of solvents in the catalyst decoration step. This aims to investigate the effects of wasteful fabrication steps on the total cost. We find that no-waste fabrication of the catalyst—electrode pair results in a cost reduction of a factor of 35 to a cost of $540 m² of coated area. The main cost driver is now the carbon paper as well as the catalyst precursors (mainly the Ni and Mo precursors). For the PSC, we find a cost reduction of a factor of 130 with a cost of $325 m² of coated area. Here, the glass substrates, the spiro-OMeTAD, and Au are still the main culprits for the PSC.

On the basis of these results, we conclude that further development is needed in order to (i) significantly improve the stability of the PSCs and the catalysts, (ii) replace the expensive Au electrodes and glass substrates and develop low-cost alternatives to the spiro-OMeTAD for the PSCs, and (iii) develop solvent-efficient, low-cost, and large-scale fabrication of the PSC and catalysts. We finally note that such development is already ongoing.6,67–72

## CONCLUSIONS

We present the synthesis, characterization, and application of a solar-driven electrolyzer. The trimetallic NiFeMo catalyst takes the shape of nanometer-sized flakes anchored to a fully carbon-based current collector comprising a nitrogen-doped carbon nanotube network, which in turn is grown on a carbon fiber paper support. This catalyst electrode exhibits a low total overpotential for water splitting of 0.45 V at 10 mA cm⁻² and combined with high-performance solution-processed PSCs forms a solar-driven water-splitting device with an impressive initial STH of 13.8%. We highlight that this performance is forms a solar-driven water-splitting device with an impressive ratio between active and coated area. With a GFF = 89% for the lab-scale catalyst, we obtain a lab-scale CPACAT = $21,500 m². Surprisingly, this high cost is mainly due to the high volume of DMF solvent used during the catalyst decoration step. For the PSC, we found an even more extreme cost of $43,400 m². Considering the low GFF of 18.7%, this results in a cost-per-active-area for the PSC (CPAPSC) of $233,000 m². This cost is mainly driven by the high cost of the glass/FTO substrates and the material-inefficient deposition of the expensive spiro-OMeTAD and Au materials.

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## EXPERIMENTAL SECTION

### Catalyst Synthesis.

Thin films of Ti (10 nm) and Fe (5 nm) were thermally evaporated (Kurt J. Lesker PVD 75) on carbon paper (Sigraet GDL 34 AA). The carbon paper was then placed into a quartz tube (3 cm diameter) and inserted in a CVD chamber for NCNT growth. The thin metal films were reduced into small particles by heating the chamber to 800 °C in an Ar:H₂ (95:5) flow and then further reduced for 15 min with ammonia (25 mL min⁻¹). The NCNTs were grown at 800 °C with a reactor pressure of 13.3 kPa and a 10 μL min⁻¹ flow of Ar:H₂ (95:5) by injecting pyridine and a syringe (NEMSYS) at 10 μL min⁻¹ for 45 min and then being cooled in Ar. To synthesize NiFeMo-NP, a small piece (1.5 cm x 2 cm) of the coated carbon paper was mixed with a solution containing 6 mL of DMF and a total concentration of 3 mg mL⁻¹ of Ni(acac)₂, Fe(acac)₃, and MoO₂(acac)₃ with a molar ratio of 5:1:4. The solution mixed together with the carbon paper was then inserted into a Teflon-lined autoclave and heated at 180 °C for 20 h before being cooled at room temperature and washed carefully with milli-Q water. To create NiFeMo-NP, the final material was further reduced by annealing in an Ar:H₂ (95:5) flow at 450 °C for 2 h and being cooled in the same reducing atmosphere before being taken out. The reference sample NiFe-NP and BioMoNPs were synthesized following the same recipe without adding molybdenum.

### Membrane Characterization.

SEM was performed on a ZEISS Merlin FESEM microscope, and EDS was measured with an Oxford Instruments X-MAX 80 mm² X-ray detector at 5 kV. The XRD patterns were recorded with a Panalytical Xpert X-ray diffractometer with Cu Kα radiation (λ = 1.5418 Å). XPS analyses were done on a Kratos axis ultradelay line detector electron spectrometer using a monochromatic Al Kα source operated at 150 W.

### Electrochemical Measurements.

The polarization curves were recorded with linear sweep voltammetry at a scan rate of 2 mV s⁻¹ in 1.0 M KOH with a potentiometer (Metrohm-AutoLab PGSTAT302N, FRA32 M module). For analyzing individual electrodes, a three-electrode cell setup was used with Ag/AgCl (CH1111-CH instruments) as a reference electrode and a coiled Pt wire (99.999%) as the counter electrode. The voltage was converted into RHE by adding 0.222 V + 0.059 pH (pH 14). All polarization curves were postcorrected from the iR resistance (2 Ω) measured from electrochemical impedance spectroscopy. The scanning direction was positive for all polarization curves except for the full device setup, where any interfering oxidation is omitted by instead scanning in the negative direction.

### Faradaic Efficiency.

Membrane-inlet mass spectrometry (MIMS) was used to quantify the gas product of OER and HER, and its sensitivity toward O₂ and H₂ was calibrated using known amounts of air and a gas mixture of Ar:H₂ (95:5), respectively. The reference and counter electrodes were the Ag/AgCl/1 M KCl and Pt coil, respectively. A gas-tight electrochemical cell was used to house the WE and RE in the main compartment, which was separated by a glass frit from another compartment that housed the CE. Both compartments contained 1 M KOH as the electrolyte and were purged with N₂ for 1 h prior to measurement. Chronopotentiometry at −10 mA cm⁻² (for HER) or 10 mA cm⁻² (for OER) was performed for 4 h, during which the gaseous aliquots in the headspace of the main compartment were analyzed using MIMS at 30 min intervals. The sum of the gas in the headspace and electrolyte (calculated using Henry’s law)53 of the main compartment yielded the amount of gas product, which was compared with the value calculated from the assumption of perfect charge-to-O₂ (4-electron oxidation) or charge-to-H₂ (2-electron reduction) conversion to obtain the Faradaic efficiency.

### Perovskite Solar Cell Fabrication.

Fluorine-doped tin oxide (FTO)-coated glass substrates (7 Ω sq⁻¹, Thin Film Devices, USA) were cleaned by 10 min subsequent steps of ultrasonication in detergent (Extran MA01); DI water (volume ratio 1:10), DI water, acetone, and isopropanol. A compact TiO₂ layer was deposited through spray pyrolysis by placing the clean and dry substrates on a hot plate (PZ28-3TD, Harry Gestigkeit) and raising the temperature stepwise to 500 °C. A precursor solution comprising a mixture of 0.2 M titanium isopropoxide (>97.0%, Sigma-Aldrich) and 2 M acetylacetone (puriss >99.5%, Sigma-Aldrich) in isopropanol (IPA, anhydrous 99.5%, Sigma-Aldrich) with a volume ratio of 9.2:20:5:4:1000 was sprayed at a rate of 14 μL s⁻¹ using a handheld airbrush (Art. 17–372, Biltema), operated at p = 20 psi, with a sweeping motion over the substrates at a 15 cm substrate-to-nozzle distance. After spray pyrolysis, the substrates were kept on the hot plate at 500 °C for 30 min for the TiO₂ to crystallize to the rutile phase (final thickness ~30 nm) before slowly cooling to room temperature. A mesoporous TiO₂ was subsequently deposited by first spin-coating (spin speed = 4000 rpm, acceleration = 4000 rpm s⁻¹, time = 30 s) a 150 mg mL⁻¹ precursor solution consisting of TiO₂ nanoparticles (DSL 18NR-T, Dyesol) dispersed in isopropanol.
followed by annealing at 500 °C for 30 min (final thickness ~200 nm).

The perovskite solution consists of formamidinium iodide (FAI, Lumtec), methylammonium bromide (MABr, Lumtec), lead iodide (PbI2, TCI chemicals), and lead bromide (PbBr2, TCI chemicals) in a 1:1:0.2:2.0 M concentration of FAI:PbI2:MABr:PbBr2. The FAI:PbI2 and MABr:PbBr2 were dissolved separately in a mixture of N,N-dimethylformamide (DMF, anhydrous 99.8%, Sigma-Aldrich) and dimethyl sulfoxide (DMSO, anhydrous >99.9%, Sigma-Aldrich) with a volume ratio of 4:1, before they were stirred together at room temperature to form the perovskite solution. The perovskite solution was spin-coated in a N2-filled glovebox ([O2], [H2O] < 1 ppm) using a two-step spin-coating process (step 1: spin speed = 1000 rpm, acceleration = 1000 rpm s⁻¹, time = 10 s; step 2: spin speed = 4000 rpm, acceleration = 1000 rpm s⁻¹, time = 30 s), and 65 μL of chlorobenzene (anhydrous 99.8%, Sigma-Aldrich) was quickly dropped onto the spinning substrate when there was 15 s left of step 2. The coated substrates were annealed at 100 °C for 1 h resulting in a uniform dark brown perovskite active layer film with a thickness of ~600–700 nm.

The hole transport layer comprised 2,2’,7,7’-Tetrakis(N,N-di-p-methoxyphenylamino)-9,9’-spirofluorene (spiro-OMeTAD, Lumtec) doped by bis(trifluoromethylsulfonyl)imide lithium salt (Li-TFSI, Sigma-Aldrich), tris(2-(1H-pyrazol-1-yl)-4-tert-butylyphridine)-cobalt(III) tris(bis(trifluoromethylsulfonyl)imide) (Co-TFSI, FKO29, Dyenamo), and 4-tert-butylpyridine (TBP, Sigma-Aldrich). Li-TFSI and Co-TFSI were separately dissolved in acetonitrile (anhydrous 99.8%, Sigma-Aldrich) to a molar concentration of 2 and 0.5 M, respectively, before being added to form a 70 mM spiro-OMeTAD solution in chlorobenzene. The hole transport layer was deposited by spin-coating the solution (spin speed = 4000 rpm, acceleration = 1000 rpm s⁻¹, time = 30 s) on top of the perovskite layer resulting in a film thickness of ~200 nm. The substrates were then stored in a desiccator with a dry ambient atmosphere for 24 h before the device was finalized with a 70 nm gold electrode, deposited by thermal evaporation (p < 5 × 10⁻⁶ mbar). An FTO/Au electrode overlap of 3 mm × 7 mm defined the active device area.

Solar Simulator Measurements. The unencapsulated perovskite solar cells were illuminated using a solar simulator (Oriel sol 3A, Newport, model 940233 A) calibrated with a reference silicon solar cell (Newsport, model 91150 V) to represent AM1.5 irradiance at 100 mW cm⁻². The operating current (Iop) and the solar cells during illumination, and a 100% Faradaic efficiency (ηH₂) was used when calculating the solar-to-hydrogen conversion efficiency (STH) through

\[
\text{STH} = \frac{I_{\text{op}} \times E_{\text{H}_2} \times \eta_{\text{H}_2}}{P_{\text{in}}}
\]

where \(E_{\text{H}_2} = 1.23 \text{ V}\) is the thermodynamic potential for water splitting.

ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acssuschemeng.1c03565.

XPS, SEM, XRD, electrochemical characterization, solar cell performance measurements, PV-driven electrolysis performance measurements, and cost analysis (PDF)

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Notes
The authors declare no competing financial interest.

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