

Transparent Cuprous Oxide Photocathode Enabling a Stacked Tandem Cell for Unbiased Water Splitting

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Photoelectrochemical water splitting represents an attractive method of capturing and storing the immense energy of sunlight in the form of hydrogen, a clean chemical fuel. Given the large energetic demand of water electrolysis, and the defined spectrum of photons available from incident sunlight, a two absorber tandem device is required to achieve high efficiencies. The two absorbers should be of different and complementary bandgaps, connected in series to achieve the necessary voltage, and arranged in an optical stack configuration to maximize the utilization of sunlight. This latter requirement demands a top device that is responsive to high-energy photons but also transparent to lower-energy photons, which pass through to illuminate the bottom absorber. Here, cuprous oxide (Cu₂O) is employed as a top absorber component, and the factors influencing the balance between transparency and efficiency toward operation in a tandem configuration are studied. Photocathodes based on Cu₂O electrodeposited onto conducting glass substrates treated with thin, discontinuous layers of gold achieve reasonable sub-bandgap transmittance while retaining performances comparable to their opaque counterparts. This new high-performance transparent photo-cathode is demonstrated in tandem with a hybrid perovskite photovoltaic cell, resulting in a full device capable of standalone sunlight-driven water splitting.

1. Introduction

Sunlight is the largest and most widespread source of renewable energy, but although the market for photovoltaics is growing by around 40% per year,^[1] the intermittent and diurnal nature of solar power requires the development of an efficient method of energy storage to accommodate the global energy demand. The use of sunlight toward the direct production

of chemical fuels is a promising route and, in particular, hydrogen generated via photoelectrochemical (PEC) water splitting represents a clean and energy-dense fuel.[2] With the aim of achieving energy conversion on a simple single device, PEC water splitting cells integrate the processes of sunlight collection and water electrolysis to produce hydrogen and oxygen, gaseous molecules which can recombine in fuel cells to efficiently utilize their stored energy.

Under standard conditions, the electrolysis of water has a reversible cell voltage of 1.23 V, but due to reaction overpotentials a voltage of 1.5 V or greater is required to drive water splitting at meaningful rates. Accomplishing this with a single material photoelectrode would require discovery of a semiconductor with demanding energetic requirements, since its conduction and valence bands must properly straddle the water redox potentials plus overpotentials.[2] No ideal single material has been discovered that can drive this reaction with reasonable efficiency, although high-throughput searches are underway.[3] A more promising approach is to employ a two- absorber tandem comprising a wider bandgap transparent top absorber stacked above a smaller bandgap component.[4] By detailed balance calculations, the ideal pair of bandgaps would be approximately 1.7 eV (absorbing wavelengths up to 730 nm) and 1.1 eV (up to 1127 nm).[5] The theoretical maximum water splitting efficiency for such a system exceeds that of a single- absorber system, a result of the more complete solar spectral utilization and the ability to produce additive photovoltages toward the electrolysis demand.

Dual-absorber tandem devices can be accomplished with photoanode–photocathode systems (PEC–PEC) or photoelectrode–photovoltaic coupled devices (PEC–PV) to generate the sufficient driving force for standalone solar water splitting while simultaneously maximizing the fraction of solar energy collected. Among the various studied tandem configurations, the benchmark performances were achieved by devices using efficient III–V materials.[6] Since the balance between materials availability, fabrication cost and device performance must be optimized to realize a competitive device, most recent attention has targeted the discovery and development of Earth-abundant materials for use toward these goals. A tandem cell combining an Earth-abundant solar cell with an oxide-based photoelectrode could be an ideal solution to obtain cost-effective unassisted water splitting.[4]

Studies employing the PEC–PV configuration have mostly focused around the use of n-type semiconductor photoanodes for water oxidation as the PEC component, largely due to the number of promising candidate materials in this class, their suitable bandgaps and their Earth-abundant nature. Among them, Fe₂O₃,^[7] BiVO₄,^[8] and WO₃^[7a,9] have been employed most commonly in approaches succeeding at unassisted water splitting. Comparatively fewer Earth-abundant candidates exist for photocathode PEC devices, with even fewer demonstrated toward complete water splitting in tandem systems.[10] While photocathode stability can prove to be a challenge, Seger et al. have recently suggested that when effective surface passivation strategies are used, tandems based on photocathodes as the larger bandgap component may offer several advantages over photoanode-based approaches.[11]

To this end, p-type photocathode materials for water reduction are a topic of ongoing research.[12] Copper-oxide-based materials, such as cuprous oxide (Cu₂O), have gained significant interest due to their elemental abundance, scalable synthesis techniques, and natural p-type character.[13] Moreover, the Cu₂O bandgap energy around 2.1 eV, and its appropriate

band edge positions, make it appealing for solar hydrogen production from water, although its poor stability in aqueous solutions is a limiting factor for its use. Paracchino et al. developed a highly active and stable multilayer composite photocathode that consists of a p–n junction between electrodeposited p-type Cu₂O and n-type overlayers of Al-doped zinc oxide (AZO) and TiO₂.^[14]

The device was recently demonstrated as part of a PEC– PEC tandem capable of complete water splitting at modest efficiency.^[10a]

Despite the recent success with this device architecture, the photovoltage produced by this photocathode alone is insufficient to drive complete water splitting, so it should ideally be incorporated as the top, wide-bandgap component of a tandem system. However, the fact that the Cu₂O thin films are typically electrodeposited onto an opaque Au layer precludes their use as a top absorber. Therefore the development of an efficient

and stable transparent Cu₂O thin film device is an important goal, wherein the portion of solar radiation not absorbed by the photocathode can be transmitted and utilized by a second photoabsorber. In this work, an innovative tandem device was enabled by the development of a transparent Cu₂O photocathode, which when connected in series with a hybrid perovskite photovoltaic was demonstrated to perform unassisted sunlight-driven water splitting.

2. Results and Discussion

2.1. Transparent Au Substrates for Cu₂O Photocathodes

The photocathode device structure employed herein is depicted in **Figure 1**. Electrodeposited p-type Cu₂O serves as the light absorbing component, producing photogenerated electrons for water reduction. As previously demonstrated, atomic layer deposition (ALD) overlayers of Al:ZnO (AZO) and TiO₂ enable heterojunction formation and corrosion protection, respectively,^[14b] and electrodeposited RuO₂ represents a highly active and stable catalyst for the hydrogen evolution reaction.^[15] In this study, these three overlayers remained unchanged while the substrate and absorber characteristics were varied. In the majority of previous works employing Cu₂O as photoabsorber in PV and PEC devices, a thick and opaque gold film was used as the hole-collecting contact to Cu₂O.^[12d,13,16] This arrangement prevents implementation of such devices in an optical tandem, since the long-wavelength portion of the solar spectrum cannot pass through to the second absorber. Our first task was therefore to adapt the substrate to enable light transmittance. Films of the transparent conducting oxide SnO₂:F (FTO) on glass are routinely used as substrate for transparent electronic devices, so we began by attempting the photocathode synthesis using Au-free, pristine FTO glass as substrate. As shown in **Figure 2**, the current density–potential (*J*–*E*) response of the device formed on bare FTO was considerably poorer than that of a typical device grown on a continuous 150 nm thick Au film. For a photocathode driving water reduction, the goal is to achieve large photocurrents at potentials well positive of the reversible potential of hydrogen evolution, 0 V vs. RHE (reversible hydrogen electrode). The typical device based on thick Au exhibits a photocurrent (*J*_{ph}) onset potential of approximately +0.5 V vs. RHE, and reaches cathodic *J*_{ph} approaching –6 mA

cm^{-2} at 0 V vs. RHE, a performance that is among the best for photoelectrodes based on metal oxide semiconductors. However, the PEC response suffered significantly in the absence of Au, with a very gradual onset and a J_{ph} reaching only -2 mA cm^{-2} at 0 V vs. RHE. This transparent, Au-free device ultimately reached significant photocurrents, but only at large overpotential for water reduction.

This behavior suggests that a significant resistive element is present in this device configuration. A key role of the substrate contact to Cu_2O is to form an ohmic junction for photogenerated hole collection.[13,17] Due to the relatively large work function of p-type Cu_2O , this constrains the contact material to one with a comparable or larger work function. It is likely that the Cu_2O contact with FTO is not an ideal ohmic junction, but rather forms a slight Schottky barrier opposing the collection of holes, and therefore contributes to the worsening of the J - E response. Indeed, electrochemical impedance spectroscopy analysis revealed a resistive element for this bare FTO device, a feature that was nonexistent in devices with Au at the interface with Cu_2O (Figure S1, Supporting Information). Additionally, the electrodeposited Cu_2O exhibited quite different morphology when grown on bare FTO, proceeding by the nucleation and growth of large and dispersed Cu_2O crystals, in contrast to the dense, uniform and continuous growth of Cu_2O films when using a gold-coated substrate. This behavior is depicted in Figure 3 via electron microscopy images of the different substrates before and after device fabrication, and correlates with previous reports on the substrate dependence of Cu_2O electrodeposition.[18] Therefore, the Au substrate seems to affect both the electronics of the junction and the quality of the electrodeposited films, making its presence important for the device performance but a challenge given our goal of transparent devices.

We hypothesized that very thin layers of Au could be used to overcome these limitations while also allowing a high degree of transparency, and therefore explored the effect of using substrates of FTO treated by brief sputter depositions of Au. As shown in Figure 3c, sputtering Au for an equivalent dose (based on calibrated film deposition rates) of 3 nm led to the formation of a discontinuous island coating on the FTO surface. Note that the Au substrates are herein labeled by their nominal thickness based on calibrated sputter deposition rates, although this does not accurately define the discontinuous morphology. Interestingly, Cu_2O electrodeposition onto these Au island substrates resulted in dense, uniform, and crystalline films (Figure 3f), similar to those deposited onto thick Au and in contrast to the large Cu_2O particles which form on bare FTO substrates. Examination of the transmittance of Au-coated FTO glass (Figure 4a) revealed that these substrates exhibit reasonable transparency across the spectrum, but the transparency drops with increasing amounts of gold.

Employing these as substrates for the fabrication of Cu_2O photocathodes, we discovered that the devices were indeed capable of efficient operation. In Figure 4b, the PEC response for devices based on the three different Au-treated substrates reveals that even a small amount of gold can enable photocathodes with onset potentials and J_{ph} values comparable to that obtained on thick Au. The performance is dependent on the amount of Au used, as the shape of the J - E curve improves with increased Au. For the smallest dose of 1 nm Au, the slow, rather linear increase in J_{ph} is likely due to a series resistance effect resulting from the limited interfacial area between Au and Cu_2O . With increased Au, the apparent fill factor of

the curves improves, but the transparency begins to suffer as a result of the coalescence of Au particles and the formation of continuous films. Note that the photocurrent transient behavior near the photocurrent onset, similar across all devices studied here, seems to result from the capacitive charging of the TiO₂ overlayer or the RuO₂ catalyst; these observations are presently under study. We have therefore established FTO-glass substrates with slight Au treatments as suitable substrates for enabling transparent Cu₂O-based devices, while identifying that the balance between transmittance and performance will be important in optimizing the tandem device. In this regard, we selected the 3 nm dose of sputtered Au as substrate for the studies continued below.

2.2. Effects of Cu₂O Absorber Thickness

The next parameter to consider is the Cu₂O absorber layer thickness. Our previous reports employed Cu₂O films of 500 nm or greater, while other groups have used thicknesses of several micrometers for photovoltaics based on electrodeposited Cu₂O.^[16] Meanwhile, the device transparency and the application of Cu₂O in dual-absorber tandems have been little explored.^[19] In this work, transparency was an important factor, and we therefore examined a series of Cu₂O thicknesses by varying the duration of electrodeposition onto substrates of FTO treated with 3 nm doses of Au. Three durations were explored, with 105, 50, and 25 min electrodepositions producing film thicknesses of approximately 500, 260, and 100 nm, respectively (see Figure S2, Supporting Information, for electron images and photographs of the films). As shown in Figure 4c, the devices exhibited varying photocathode performances. The device with the thinnest absorber layer (100 nm) showed diminished plateau photocurrents, whereas the 260 and 500 nm films exhibited nearly identical *J*-*E* responses, both being comparable to the performance on a typical thick Au substrate. Variation in the absorber thickness modifies the light absorption profile and the resulting quantum efficiency spectra. In Figure 4d, the incident photon-to-current conversion efficiencies (IPCE) of the devices under monochromatic illumination reveal the changes in spectral response. Although the bandgap of Cu₂O is often stated as being approximately 2.1 eV, the nature of this transition is direct but forbidden, whereas the first allowed transition occurs at around 2.5 eV.^[13,20] This effect is clearly seen as an inflection in the IPCE spectra, where photons of lower energy (wavelengths longer than 500 nm) are poorly utilized by these thin Cu₂O films. Film thicknesses of several micrometers are required for significant absorption in this range,^[19a] but for a stacked tandem device the transmittance of long-wavelength photons is an important factor, and it can be seen that even a 500 nm Cu₂O film contributes to significantly decreased transmittance as compared to a 260 nm Cu₂O device (Figure 4d). This loss is possibly due to an increase in scattering or reflection due to the visibly larger degree of surface roughness for the thick Cu₂O films (Figure S2a, Supporting Information). Furthermore, the response to short wavelengths decreases with thicker absorber layers, a result of the poor majority carrier (hole) collection in thick films. The 50 min electrodeposition photocathode was therefore selected as a relatively optimized candidate for balancing PEC performance with optical transparency in the complete water splitting device.

2.3. Tandem Device for Complete Water Splitting

A schematic of the assembled tandem device is shown in **Figure 5**, where the two absorbers are placed back-to-back, the photovoltaic electron collector is wired to the photocathode, and its hole collector is wired to a water oxidation anode. In a dual-absorber PEC–PV tandem, the photoelectrode and the photovoltaic utilize photons of different regions of the solar spectrum to enable broad sunlight harvesting. Furthermore, since they will be connected in series and therefore operate at the same current density, it is desirable for their individual photocurrent responses to be well matched. In **Figure 6a**, the IPCE responses of the photocathode and the perovskite photovoltaic (behind the photocathode) are multiplied by the AM1.5G one-sun photon flux to reveal the expected photogenerated electron flux from each device, which when integrated yields their expected photocurrent densities.

In constructing a PEC–PV tandem for complete water splitting, three components are required: a suitably transparent photocathode driving hydrogen evolution, a photovoltaic cell responsive to the transmitted photons, and an anode for oxygen evolution. Among water oxidation catalysts, IrO₂ is known to be a top performer,^[21] and was therefore our anode choice for this proof-of-concept device. Even so, the water oxidation reaction imparts a large energetic demand on the overall water splitting processes, especially in near-neutral solutions, and significant overpotentials beyond the reversible potential for oxygen evolution (1.23 V vs RHE) are required to achieve meaningful current densities. In the tandem configuration employed here, the flat anode electrode is positioned parallel to the path of light, an approach that allows scaling its active area beyond that of the illuminated area as a tactic toward reducing the overpotential required for supporting the tandem photocurrent. In previous reports of PEC–PV tandems for unbiased water splitting, it has been common to either make no mention of the counter electrode dimensions or to employ counter electrode areas that are several times larger than the photoelectrode illuminated area.^[7a,8b,22] This non-trivial parameter plays an important role in the tandem construction and performance, and here we used a catalyst with an active area of approximately 30 times that of the photocathode illuminated area. While the anode overpotential is a key challenge producing an efficient tandem device, this mismatch of electrode areas can actually highlight an advantage of a photocathode-based tandem. Since the oxygen evolution reaction, which exhibits significantly larger overpotentials than the hydrogen evolution reaction, occurs on a non-photoactive component, its relative area may be increased as long as the cell design allows its placement out of the path of illumination, an approach that is even more desirable when abundant catalyst materials are employed.

Figure 6b presents the J – E behavior of all three components tested individually, where the raw current of each was normalized by the photocathode illuminated area. Comparison of the photocathode and anode curves reveals the current-dependent additional voltage needed to enable complete electrolysis. For instance, around 1 V is needed in order to drive a current density of 2 mA cm⁻² between these electrodes. Few single-absorber photovoltaics are capable of photovoltages this large, but the emergent high- V_{OC} hybrid perovskite photovoltaics represent promising candidates for this application.^[23] Here, we employed a PV cell based on the mixed-cation formulation formamidinium methylammonium lead iodide ((MA) _{x} (FA)_{1- x} PbI₃)^[24] that exhibits a V_{OC} under one sun illumination of 1.13 V (see **Figure S3**, Supporting Information, for the photovoltaic cell J – V and IPCE analysis). The PV cell was placed against the back window of the sealed PEC cell and connected by wires to the

electrodes. Since the PV is electrically in series between the anode and photocathode, its J - V response can be plotted between the electrode J - E curves in order to predict the tandem operation current. Under operation, the currents through each component are equal, and the potentials of each contact spontaneously adjust to reach this equilibrium. As shown in Figure 6b, this treatment predicts an operating current density of about 2 mA cm^{-2} for this tandem configuration. We then connected all components in series and illuminated the cell with one-sun intensity simulated sunlight, using a potentiostat to monitor the current flowing between the PV and anode (applying zero bias) while simultaneously performing in-line gas chromatography measurement of the evolved gases. Figure 6c shows the resulting measured current density, where it can be seen that the actual tandem performance corresponded well with that predicted by the separate component analysis. Measurement of both hydrogen and oxygen, important for proving complete water splitting,^[25] led to calculated Faradaic efficiencies around 100% for each gas (Figure 6d). Variations in the gas measurements are a result of the buildup and release of bubbles on the electrode surfaces. A J_{ph} of 2 mA cm^{-2} combined with the near-unity yield of evolved gases corresponds to a solar-to-hydrogen (STH) efficiency of 2.5% by Equation (1):

$$\text{STH} = \frac{J_{\text{ph}} (\text{mAcm}^{-2}) \times 1.23\text{V} \times \eta_{\text{F}}}{P_{\text{in}} (\text{mWcm}^{-2})} \quad (1)$$

where J_{ph} is the photocurrent density during unbiased operation, 1.23 V is the standard-state potential for water electrolysis, η_{F} is the Faradaic efficiency of evolved hydrogen, and P_{in} is the power of the incident illumination, taken here as 100 mW cm^{-2} for the AM1.5G spectrum at one sun intensity. This efficiency, while modest in comparison to more sophisticated systems,^[6] represents an important advance among photocathode-based tandem devices employing Earth-abundant absorbers.

The tandem device was tested under continuous illumination for over 2 h, during which time the J_{ph} slowly decreased to stabilize at around 1.5 mA cm^{-2} . In addition to measurements of J_{ph} and η_{F} , monitoring the device potentials during operation revealed insight into the tandem operation. By periodically measuring the electrode potentials against a reference electrode in the cell as illustrated in Figure 5 and shown in Figure 6e, we observed that the anode potential was quite stable whereas the photocathode potential shifted to more positive values over time. These changes were concurrent with the gradual decrease of device J_{ph} , and the behavior can be interpreted by referring to the analysis in Figure 6b. The measured potentials represent the potentials at the contacts between the PV cell and each electrode, equivalent to the labeled crossover points. Since the anode curve is steep, changes in current are accommodated with relatively little change in potential. For the photocathode, on the other hand, the observed positive shift in potential, combined with the decreasing measured J_{ph} , reveals that the performance decline is likely a result of a decrease in the PV photovoltage. Indeed, when its J - V response was re-tested after tandem operation, the PV cell V_{OC} and fill-factor exhibited a slight decrease (Figure S3, Supporting Information). Despite this observation, we note that several recent works have demonstrated extended stability

for hybrid perovskite photovoltaics,^[26] and this is a topic of ongoing study for the device type employed here. Meanwhile, the photocathode was found to be robust. In an extended test, a representative Cu₂O photocathode of this same transparent configuration was found to be highly stable when tested for 24 h under continuous operation (Figure S4, Supporting Information).

There are several clear paths toward improving upon this tandem efficiency, mostly based on the enhancement of the Cu₂O photocathode performance. As shown above, the J_{ph} is highly dependent on the operating potential. An increase in the photovoltage of the device, manifesting as a positive shift in the $J-E$ curve, would allow tandem operation at higher current density. For instance, a photovoltage increase of 0.2 V could enable a near-doubling of the operating current density to $\approx 4 \text{ mA cm}^{-2}$. Recently it has been shown that improving the nature of the Cu₂O-overlayer junction by using Ga₂O₃ interlayers can lead to significantly enhanced photovoltages,^[16b,27] an approach that may prove useful toward tandem applications. Improvement in the photocathode photocurrent is also needed, although the forbidden electronic transitions for photon energies below 2.5 eV require much thicker Cu₂O films for improved absorption in that range,^[20] posing a challenge when targeting high device transparency.

The tandem performance may also be improved by operating in a highly alkaline or acidic electrolyte more suitable for efficient electrolysis. Although rapid stirring was used here to prevent mass transport limitations and pH gradient build-up, it has been shown that devices in near-neutral solutions can be fundamentally limited in efficiency when operated for extended periods.^[28] Furthermore, catalysts for water oxidation are generally more efficient in alkaline solutions, for which there are several desirable Earth-abundant candidates.^[21] At present, we found this TiO₂ overlayer approach to be insufficiently stable in alkaline conditions to allow a demonstration extended operation, but further study in this direction is ongoing,^[29] since a stable and efficient photocathode in alkaline solution would be highly desirable.

3. Conclusion

In this work, we constructed a PEC–PV water splitting tandem using a Cu₂O photocathode and a hybrid perovskite photovoltaic. In developing a transparent photocathode, we discovered the important role of gold as substrate and explored how different aspects of the device architecture influence the balance between performance and transparency. This transparent photoelectrode enabled the construction of an optically stacked two absorber tandem device capable of performing standalone sunlight-driven water splitting at up to 2.5% solar-to-hydrogen efficiency, a performance that may be significantly enhanced by further development of the Cu₂O photocathode.

4. Experimental Section

Cu₂O Photocathode Preparation: The films of cuprous oxide were electrodeposited onto fluorine-doped tin oxide (FTO, TEC-15, NSG glass) substrates with or without Au treatments. The FTO-glass substrates were cleaned by sequential ultrasonic treatments in soapy water (15 min), acetone (15 min), ethanol (15 min), and deionized water (15 min). The substrates were then coated with various treatments of Au by DC sputtering. The

standard opaque films of 150 nm were deposited at a calibrated rate of 1.1 nm s^{-1} (after depositing a 10 nm Cr adhesion layer). The transparent Au treatments (without Cr layers) were performed at a calibrated rate of 0.2 nm s^{-1} for durations 5, 15, and 25 s to yield the substrates labeled herein by their nominal thicknesses of 1, 3, and 5 nm Au. The electrodeposition of cuprous oxide from a basic solution of lactate-stabilized copper sulfate was performed as described previously.^[30] The electrodeposition was performed in galvanostat mode (constant current density of -0.1 mA cm^{-2}) by using a two-electrode configuration with a platinum mesh as the counter electrode. The time of deposition was varied as described in the main text. Thin n-type oxide overlayers were deposited atop the Cu_2O films by ALD using a thermal ALD system (Savannah 100, Cambridge Nanotech), as described previously.^[15] The ALD protective structure consisted of 20 nm of Al:ZnO (AZO; deposited at $120 \text{ }^\circ\text{C}$ using precursors of diethyl zinc, trimethylaluminum, and water vapor) followed by 100 nm of TiO_2 (deposited at $150 \text{ }^\circ\text{C}$ using precursors of tetrakis(dimethylamino) titanium at $75 \text{ }^\circ\text{C}$ and H_2O_2 50% in water). After ALD, the electrode areas were defined by encapsulation using hot glue or opaque epoxy. Onto the exposed active area, RuO_2 catalyst was galvanostatically deposited under illumination using an aqueous solution of $1.3 \times 10^{-3} \text{ M KRuO}_4$. The deposition was carried out at a current density of $-28.3 \text{ } \mu\text{A cm}^{-2}$ for 15–20 min under simulated one sun illumination, following the procedure described previously.^[15]

Spectroscopic and Microscopic Characterization: The morphology of the substrates and photoelectrodes was characterized using a high-resolution scanning electron microscope (Zeiss Merlin) with an in-lens secondary electron detector. Cross-sectional images were acquired from freshly cleaved surfaces. Total transmittance spectra were measured with a spectrophotometer (Shimadzu UV-3600) equipped with an integrating sphere. The partial devices were tested directly in air, whereas the full devices were wetted with water and sandwiched between quartz slides in order to approximate the optical behavior of the device within the PEC cell. The absence of sample was used as a transmittance blank in order to account for the contribution of every layer in the device.

Electrochemical Characterization: The PEC cell performance was evaluated in a standard three-electrode configuration using the Cu_2O device as photocathode, a Pt wire as counter electrode, and a reference electrode of Ag/AgCl/sat. KCl. The electrolyte solution was Na_2SO_4 (0.5 M) buffered with phosphate (0.1 M) to obtain pH 5.0. An Ivium Potentiostat/Galvanostat was used to acquire the photoresponse under chopped irradiation from a 450 W Xe lamp (Osram, ozone-free) equipped with an IR/UV filter (KG3 filter, 3 mm, Schott). The PEC cell was positioned for illumination at one-sun intensity, the position determined by measuring the short-circuit current on a calibrated silicon diode fitted with a KG3 filter to obtain low spectral mismatch with the AM 1.5 spectrum across the relevant wavelength range of 300–800 nm. The scan rate for all current–potential (J – E) studies was 10 mV s^{-1} in the cathodic direction. The electrolyte was continuously bubbled with nitrogen during the J – E measurements and stability tests to remove oxygen and thus eliminate signals of oxygen reduction.

IPCE measurements were performed under light from a 300 W xenon lamp (Cermax PE 300 BUV) passing through a monochromator (Bausch & Lomb). In three-electrode configuration the photocathode current response was measured while holding the potential constant. This photoresponse was compared against that of a calibrated Si photodiode to determine the IPCE at each wavelength.

Photovoltaic Preparation: A 50 nm compact TiO₂ blocking layer was first deposited onto the surface of a precleaned FTO substrate by spray pyrolysis on a hotplate at 450 °C using Ti-isopropoxide and acetylacetone in ethanol. Then a nanostructured layer of TiO₂ was deposited by spin-coating diluted Dyesol paste (18NR-T), and sintering at 500 °C for 20 min. The desired perovskite solutions of (FAPbI)_{3-x}(MAPbI₃)_x, were prepared by dissolution of CH₃NH₃Br, NH₂CH₂NH₂I, with PbI₂ and PbBr₂ in the mixed solvent of DMSO and DMF. The mixed perovskite film was obtained by spin-coating the precursor solution, followed by antisolvent treatment. The coated films were then placed on a hot plate set at 100 °C to evaporate the solvent. The composition of hole transport material was 2,2',7,7'-tetrakis-(N,N-di- p-methoxyphenyl-amine)-9,9'-spirobifluorene (spiro-OMeTAD, 0.06 M, Lumtec.), bis(trifluoromethane)sulfonimide lithium salt (LiTFSI, 0.030 M, 99.95%, Aldrich), FK209 (Co[t-BuPyPz]₃[PF₆]₃, 0.0024 M), and 4-tert-butylpyridine (0.2 M, 99%, Aldrich) in anhydrous chlorobenzene (99.8%, Aldrich). The perovskite-sensitized TiO₂ films were coated with HTM solution using spin-coating, followed by deposition of Au (80 nm) as electrode by thermal evaporation.

IrO₂ Anode Preparation: A 1.8 cm² square piece of titanium foil (99.7%, 0.25 mm; Sigma Aldrich) was etched for 60 min in boiling oxalic acid (1 M, ≥97%, anhydrous, Fluka). Subsequently, H₂IrCl₆ (30 μL, 0.2 M, 99.9%, hydrate, ABCR) in isopropanol (ACS Reagent, Merck) were drop cast on the foil. This was followed by drying at 70 °C for 10 min and calcination at 500 °C for 10 min in air. The step was repeated three times on each side of the Ti foil, resulting in the deposition of 6.3 mg of IrO₂ onto each side.

Tandem Assembly and Testing: After defining the active area of the transparent photocathode using opaque epoxy (Loctite Hysol 9461), the device was fixed into a custom gas-tight test cell with front and back windows of quartz, gas inlet (submerged) and outlet (headspace) tubes, and feedthroughs for the anode, cathode, and reference electrodes. The IrO₂ anode was positioned to the side, approximately 1 cm away from the photocathode surface. The pH 5 electrolyte solution (15 mL) was filled into the cell, rapid stirring was applied, and a 20 mL min⁻¹ flow of He was continuously bubbled through the cell. Behind the electrochemical cell, against the rear quartz window, was placed the perovskite photovoltaic cell, which was illuminated by light passing through the photocathode aperture. The photovoltaic FTO contact (electron collector) was connected to the photocathode by a short wire, while its gold contact (hole collector) was connected (through a potentiostat in 2-electrode configuration with zero applied bias to measure the short-circuit current) to the IrO₂ anode. The cell was illuminated through the front window by simulated sunlight from a 450 W Xe lamp (Osram, ozone-free) equipped with an AM1.5G filter (LOT-QD), calibrated with a silicon diode to one sun intensity. For in-line characterization of evolved hydrogen and oxygen, the outflow was periodically injected

into a gas chromatograph (TRACE Ultra with PDD detector, Thermo Scientific; ValcoPLOT Molesieve 5Å fused silica column).

Supporting Information

Supporting Information is available from the Wiley Online Library or from the author.

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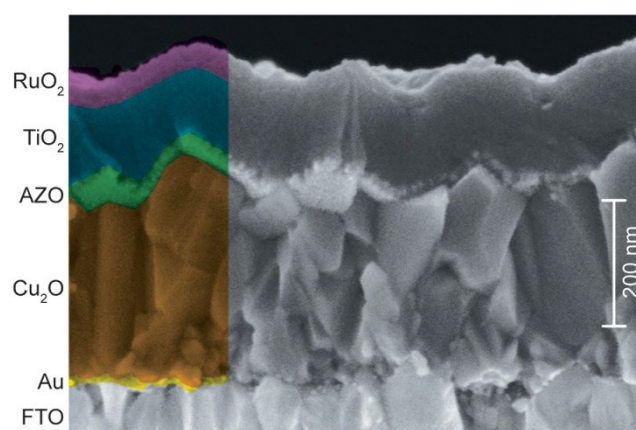


Figure 1. Cross-sectional scanning electron microscopy image of a Cu₂O photocathode device based on an FTO substrate treated with a 3 nm dose of Au. False-color was added to aid visualization of the layers.

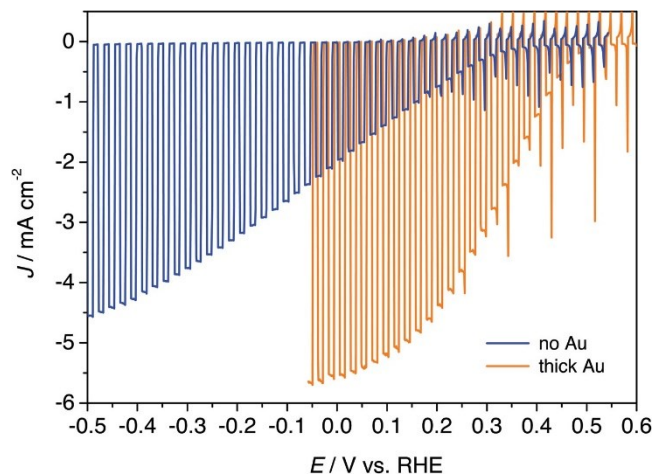


Figure 2. J - E response under chopped illumination for photocathodes synthesized both with and without the presence of a 150 nm thick Au substrate layer. In the absence of the Au layer, the performance of the photocathode is severely worsened. Conditions: pH 5 electrolyte, one-sun intensity chopped illumination, 10 mV s^{-1} scan rate from positive.

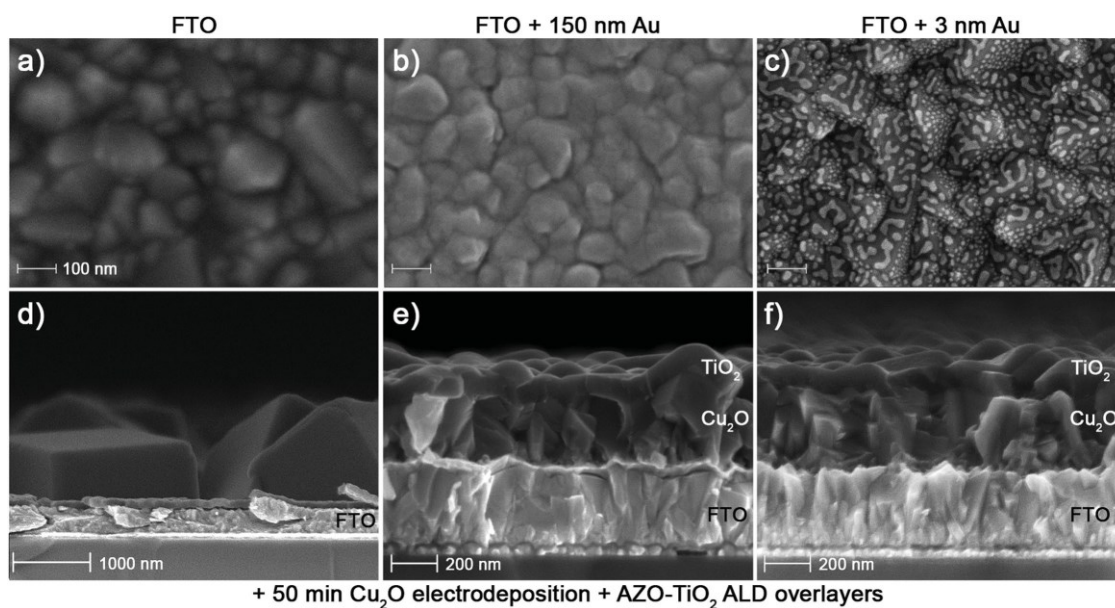


Figure 3. Scanning electron images of different substrates before (top row; top view) and after (bottom row; cross-section view) device fabrication following identical treatments (50 min Cu_2O electrodeposition followed by atomic layer deposition of AZO and TiO_2 overlayers). The substrates examined were a) bare FTO glass, b) FTO with a 150 nm thick Au film, and c) FTO with a 3 nm dose of Au (scale bars: 100 nm). On bare FTO, the Cu_2O nucleates and grows into large, d) distinct crystalline particles, whereas on both e) 150 nm and f) 3 nm Au-treated substrates the Cu_2O growth is uniform, dense, and continuous.

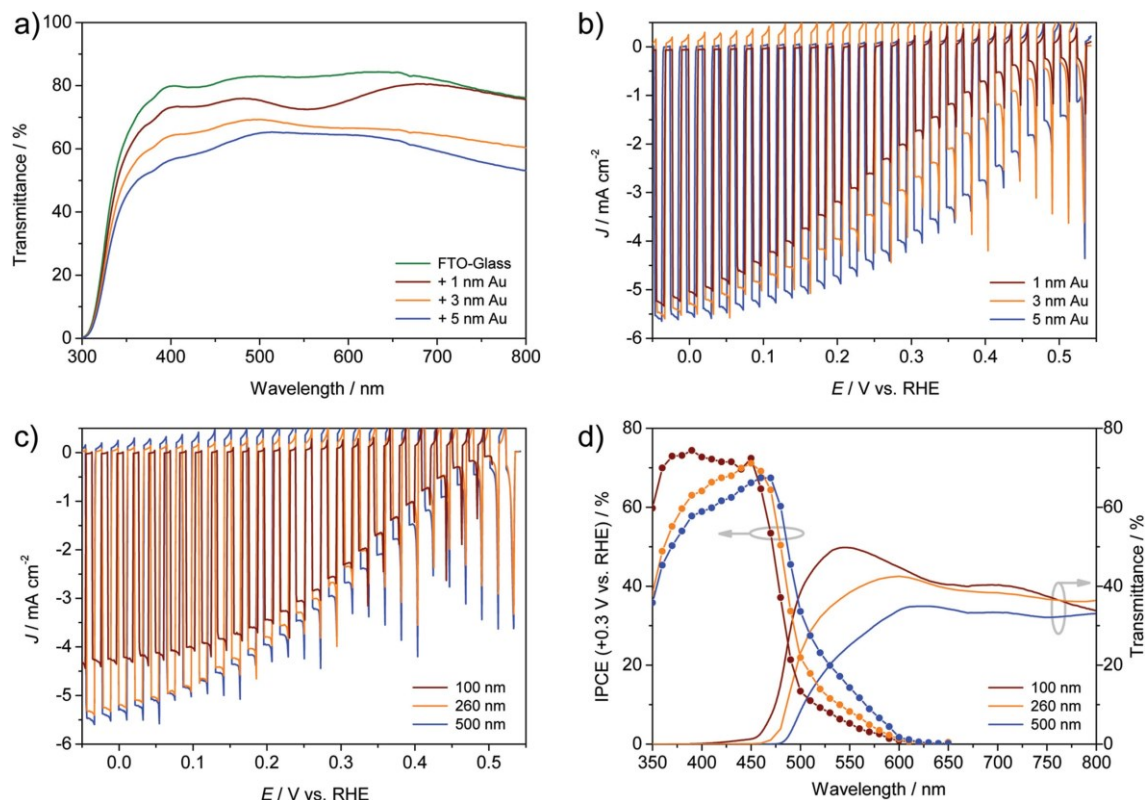


Figure 4. Optical and photoelectrochemical effects of Au and Cu₂O thickness variation. a) Transmittance spectra of FTO-glass substrates with the addition of various doses of sputtered Au. b) J - E responses of photocathodes based on 105 min Cu₂O electrodepositions onto substrates of various Au treatments, tested in pH 5 electrolyte under 1-sun intensity chopped illumination, with a 10 mV s⁻¹ scan rate from positive. c) J - E responses and d) IPCE and transmittance spectra for devices of varied Cu₂O thickness formed onto 3 nm Au-treated substrates. The devices are labeled by approximate Cu₂O thickness. The transmittance spectra were obtained on samples in air. The IPCE responses were measured while biased at +0.3 V vs. RHE. All photocathodes in (b-d) were tested following 15.0 min of RuO₂ catalyst deposition on their surfaces.

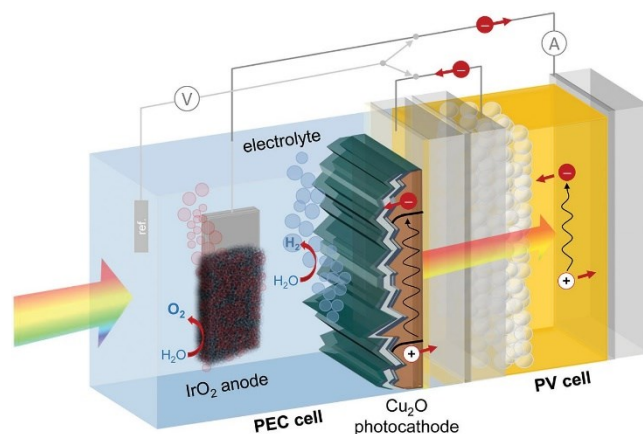


Figure 5. Schematic of the Cu₂O–perovskite–IrO₂ tandem cell during operation. An ammeter (A) is employed to monitor the short-circuit current flowing through the unbiased tandem device, while a voltmeter (V) is used to periodically measure the potentials of the anode and cathode contacts against a reference electrode in the solution.

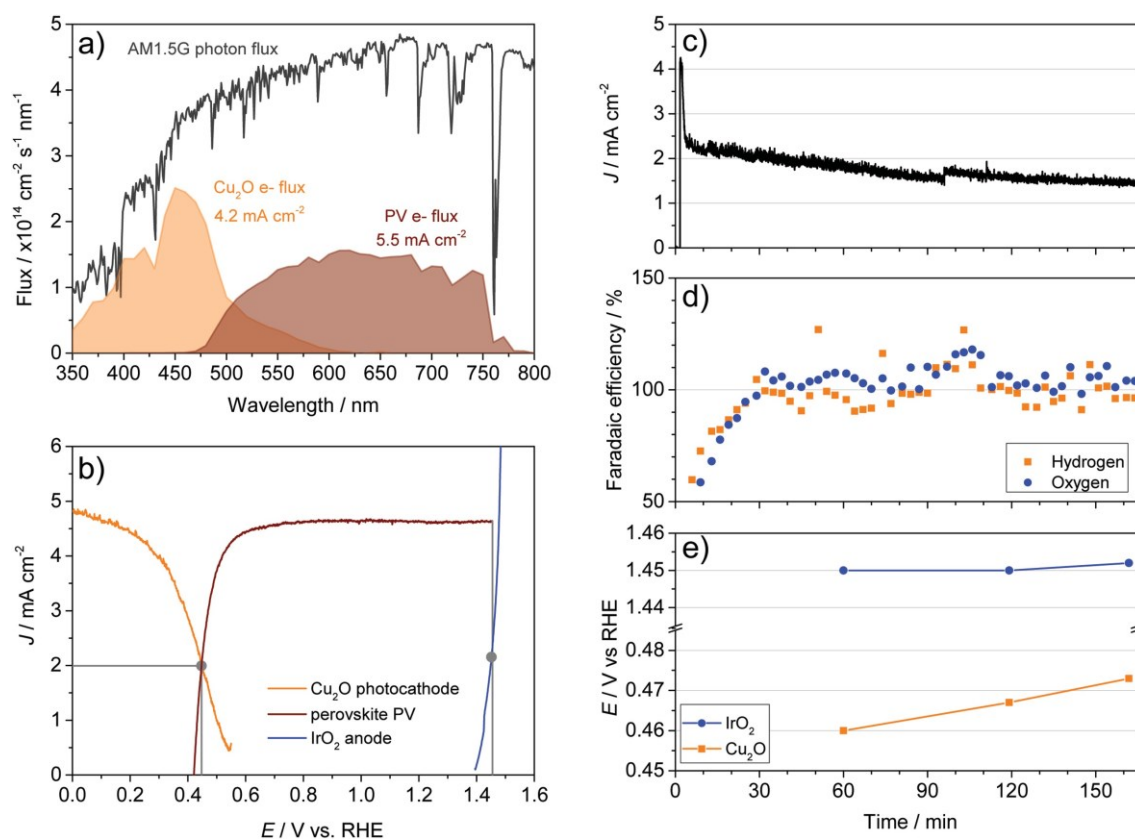


Figure 6. PEC–PV tandem assembly and operation. a) Plot of the spectral flux of photons in the AM1.5G spectrum, and the expected electron current flux of photocathode and PV obtained from multiplication of their respective IPCE responses by the photon flux (note that for IPCE acquisition, the photocathode was biased at +0.3 V vs. RHE and the PV was measured at short circuit). Integration yields the expected current densities labeled for each component. b) J – E plots of the photocathode and anode components with overlaid J –

V response of the photovoltaic cell. The photocathode configuration was 3 nm Au + 260 nm Cu₂O + ALD overlayers + 20 min electrodeposited RuO₂ surface catalyst. The position of the photovoltaic curve was defined by actual potential measurements at the photovoltaic electrode contacts after 60 min of tandem operation, as indicated by gray markers. c) Photocurrent density, d) Faradaic efficiency from in-line gas measurements, and e) potential measurements during operation of the complete assembled tandem in a sealed, stirred cell under continuous flow of He carrier gas, with a photocathode and photovoltaic illuminated area of 0.057 cm². Approximately 30 min were required for the produced gases to reach equilibrium in the cell.