

Interfacial Engineering of Gallium Indium Phosphide Photoelectrodes for
Hydrogen Evolution with Precious Metal and Non-Precious Metal Based
Catalysts

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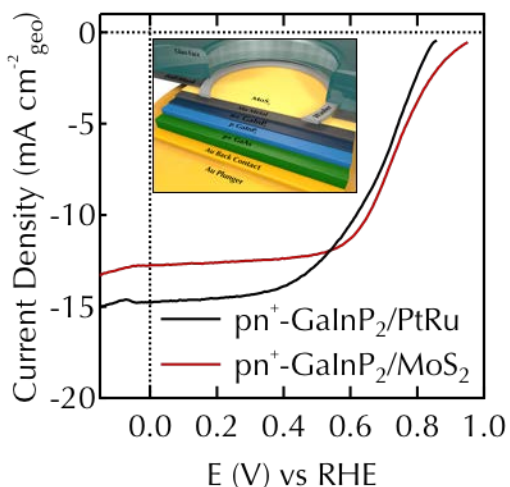
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ABSTRACT

Gallium indium phosphide (GaInP_2) is a semiconductor with promising optical and electronic properties to serve as the large bandgap, top junction in a dual absorber tandem solar water splitting device. Poor intrinsic catalytic ability and surface corrosion in aqueous electrolyte remain key obstacles. Significant progress has been made developing thin-film protection layers and active catalysts for photoelectrochemical devices, but combining these into a catalytic protection layer that can provide long-term stability without sacrificing performance has proven difficult due, in large part, to challenges in developing active and stable interfaces. In this work, we demonstrate that a nanoscale molybdenum disulfide (MoS_2) film functions both as an effective protection layer and excellent hydrogen evolution catalyst for GaInP_2 photocathodes, with only a $\sim 10\%$ loss in initial light-limited current density after 100 h, and a photocurrent onset potential better than that of the same state-of-the-art device with a platinum-ruthenium catalyst. Using transient photoreflectance spectroscopy, we probed the carrier dynamics of these photocathodes and show that the MoS_2 coated device exhibits improved electron transfer at the surface interface compared to the PtRu catalyzed device. These MoS_2 protected devices are among the most active and stable single-absorber photocathodes for solar water splitting to date and offer a promising pathway towards generating hydrogen with high efficiency and significant longevity.

TOC GRAPHICS



KEYWORDS

photoelectrochemistry, solar water splitting, hydrogen production, electrocatalysis, stability, III-V semiconductors.

TEXT

Hydrogen is a staple chemical in today's society. Millions of tonnes are utilized in processes ranging from oil refining, which produces 30% of the world's fuel, to ammonia production, which feeds 48% of humanity^{1,2,3}. Currently, hydrogen is primarily produced from fossil fuels, but one pathway toward more sustainable production is to split water into hydrogen and oxygen using the energy from sunlight. Designing a device that can accomplish this efficiently and with long-term stability could have a major impact on today's industry by producing this commodity in a clean, economical fashion, with the possibility of growing its market as a primary energy carrier in place of fossil fuels⁴.

Stability is a critical concern for many solar water splitting devices as semiconductors oftentimes undergo significant surface corrosion in the aqueous conditions required for efficient operation⁵. Substantial progress has been made extending the lifetime of semiconductors such as Si and GaAs with thin-film protection layers, such as titanium dioxide. However, to achieve high efficiency, these protection layers, which may be inert for the hydrogen or oxygen evolution reaction, often require the addition of a cocatalyst^{5, 6}. Furthermore, while the energetics of the interfaces of these protection layers + cocatalysts have been studied experimentally and theoretically^{7, 8, 9}, further insight into the complex carrier dynamics of these systems is needed to understand and effectively engineer high performing devices.

In a dual-junction tandem absorber device, the protection + cocatalyst layers for the top absorber must be engineered to avoid significant light reflection and parasitic absorption. As a result, the devices with the greatest efficiencies use highly active precious metal cocatalysts, such as Pt and Rh, since they provide high turnover per active site and favorable HER onset potentials, limiting the loading needed and minimizing deleterious light blocking^{10, 11, 12}. An earth-abundant thin-film material that can provide both long-term protection and efficient catalysis can eliminate the need for a separate protection layer and precious metal cocatalyst, reducing light blocking and cost.

Earth-abundant molybdenum sulfide nanomaterials possess excellent stability in acidic electrolyte and have edge sites with exceptional activity for the HER making them a superb candidate for a surface protection layer that has intrinsic catalytic ability^{13, 14, 15}.

Furthermore, recent studies have also demonstrated the effectiveness of MoS₂ as a protection layer on multiple semiconductors^{15, 16, 17, 18}. Herein, we demonstrate that a thin coating of molybdenum disulfide can impart both long-term stability as well as effective catalytic ability to a pn⁺-GaInP₂ photocathode. We further probe the GaInP₂ semiconductor interfacial carrier dynamics using transient photoreflectance spectroscopy (TPR) and highlight an important difference between conformal vs island catalysts in this system. GaInP₂ is a semiconductor with promising optical and electronic properties as the top absorber in a dual absorber tandem device but degrades within hours in the acidic conditions optimal for the hydrogen evolution reaction^{17, 18, 19, 20, 21, 22}. In this work we show that after 100 h of testing, the MoS₂ protected device showed only a ~10% loss in light-limited current density, and the surface of the photocathode remained in pristine condition with no exposed GaInP₂ as measured using x-ray photoelectron spectroscopy (XPS). Furthermore, the MoS₂ protected device showed an improved onset potential of 0.91 V vs the reversible hydrogen electrode (RHE) compared to a similarly prepared pn⁺-GaInP₂ photocathode coupled with a PtRu catalyst that exhibited an onset potential of 0.83 V vs RHE and which failed within only 6 hours of testing. Transient photoreflectance spectroscopy explains this improved onset by revealing improved electron transfer at the MoS₂-GaInP₂ interface compared to the PtRu-GaInP₂ interface.

This study, in concert with previous studies on Si and GaInP₂ photocathodes, demonstrates the promise of thin-films of MoS₂ as a non-precious protection layer whose activity is competitive with precious metal cocatalyzed devices in addition to exhibiting improved stability. These MoS₂ protected photocathodes are among the most active and

longest-lasting single junction devices to date and show promise to generate hydrogen with high efficiency and significant longevity. This will help enable the transition from lab-scale solar water splitting devices to prototype-scale devices that can generate meaningful quantities of hydrogen.

Photocathodic structure

Two different photocathode architectures were examined in this study and are illustrated in figures 1a and 1b. Both architectures use the same single crystal pn^+ -GaInP₂ semiconductor epitaxially grown on a degenerately doped p-type GaAs substrate. The addition of ~25-nm thick n^+ -GaInP₂ layer to a p-GaInP₂ photocathode provides ~0.5 V more positive photocurrent onset potential by improving band edge energy alignment at the electrolyte²³. For the pn^+ -GaInP₂/MoS₂ photocathode, the MoS₂ coating was synthesized by a partial thermal sulfidization of a sputtered thin film of Mo metal (see the Methods section). This method creates a thin, conformal surface coating of MoS₂ that provides an electronically conductive and protective barrier from the electrolyte, catalyzes the HER, and can be applied while minimizing parasitic light absorption^{16, 17}. For the pn^+ -GaInP₂/PtRu photocathode, PtRu nanoparticles were deposited onto a different part of the same pn^+ -GaInP₂ wafer using flash sputtering in a manner identical to that of a 16.4%-efficient tandem III-V cell, one of the highest STH efficiencies for an integrated PEC device reported to date^{10, 23}. This method was chosen to create a device that would be among the most active single junction photocathodes and represent a competitive comparison point for the pn^+ -GaInP₂/MoS₂ photocathode.

To confirm successful fabrication of the desired architectures, cross-sectional scanning electron microscopy (SEM) was performed, as shown in figures 1c and 1d. The SEM images reveal the degenerately-doped GaAs epitaxial substrate and 1 μm thick pn^+ -GaInP₂ layer common to both architectures. The n^+ layer is expected to be about 25 nm thick and have a doping density of about $7 \times 10^{18} \text{ cm}^{-3}$ (see Methods). For the pn^+ -GaInP₂/MoS₂ photocathode, the synthesis of the Mo/MoS₂ layer is identical, except for a small temperature reduction in the sulfidization step, to one performed previously on a crystalline silicon substrate that was characterized with cross-sectional transmission electron microscopy (TEM)¹⁶. On the basis of these studies, the MoS₂ layer is expected to consist of several conformal stacked basal planes that reside on top of a thin layer of Mo metal. The total thickness of all the layers above the pn^+ -GaInP₂ substrate is ~ 5 nm. For the pn^+ -GaInP₂/PtRu photocathode, the deposition of the PtRu catalyst is identical to one performed previously on a tandem III-V cell that was characterized with STEM²³. On the basis of this study, the PtRu is expected to consist of 2-5 nm particles with a surface coverage of about $\sim 30\%$ ²³.

The composition and chemical state of the surface of both architectures were studied using XPS and are shown in figures 2 and 3. The pn^+ -GaInP₂/MoS₂ photocathode shows peaks in the Mo 3d region and S 2p region. The peaks in the Mo 3d can be deconvoluted into 3 doublets indicating that Mo exists in 3 different states: a reduced metallic state consistent with Mo metal, a 4+ state consistent with MoS₂, and a 6+ state consistent with MoO₃^{13, 24}. The S 2p region shows a peak corresponding to sulfur in the 2- state consistent with MoS₂^{13, 24}. These XPS peaks are consistent with the Mo/MoS₂ layer

expected from TEM characterization of the MoS₂-Si system¹⁶. The absence of detectable peaks in the Ga 2p, In 3d, or P 2p regions further verifies the existence of a conformal Mo/MoS₂ layer on the surface of the pn⁺-GaInP₂.

The pn⁺-GaInP₂/PtRu photocathode shows peaks in Pt 4f, Ru 3p, Ga 2p, In 3d, and P 2p regions. The Pt 4f doublet is consistent with a reduced metallic Pt²⁵. While the Ru 3p 1/2 peak is obfuscated by a Ga Auger peak, the Ru 3p 3/2 peak confirms that Ru also exists in a reduced metallic state^{26, 27}. The Ga 2p and In 3d regions each show doublets of Ga and In in the 3+ oxidation state consistent with GaInP₂.^{17, 27, 28} The P 2p region shows two doublets, one for P in the 3- oxidation state consistent with GaInP₂, and one for P in the 5+ oxidation state consistent with a metal phosphate^{29, 30}. The presence of peaks in the Ga 2p, In 3d, and P 2p regions indicates that the PtRu catalyst does not form a conformal layer. This is consistent with the nanoparticulate conformation observed in the STEM characterization of the PtRu morphology on GaInP₂²³.

Electrochemical Performance

Following an established methodology³¹, the PEC performance of the pn⁺-GaInP₂/MoS₂ and pn⁺-GaInP₂/PtRu photocathodes was evaluated in a three-electrode compression cell using 3M sulfuric acid electrolyte with 1 mM Triton X-100 (OmniTrace® EMD Millipore) surfactant added to expedite H₂ bubble evolution (additional details in Methods). The working electrodes were illuminated by a tungsten-halogen lamp with a water-filled IR filter and light shaping diffuser, and illumination intensity was set to “one sun” using a GaInP₂ reference cell calibrated to the AM1.5G standard³². The performance

of the photocathodes was first measured with a linear sweep voltammogram (LSV) scanning, under reverse bias, towards more positive potentials to determine the onset potential and light-limited photocurrent density. The voltammogram was halted when the current density reached about -0.5 mA cm^{-2} to prevent any surface oxidation from positive oxidative current. The LSV was immediately followed by a chronoamperometry (CA) measurement conducted at a constant potential of -0.3 vs. MSE (0.334 V vs. RHE). At this potential, the photocurrent is light limited, but the devices are still operating near the maximum power point. Such an operating point mimics the ideal operating point of a tandem device at short circuit, without excessive reverse bias that can inflate durability by increasing charge separation and cathodic protection while suppressing anodic corrosion mechanisms²³.

As shown in figure 4a, the $\text{pn}^+\text{-GaInP}_2/\text{MoS}_2$ photocathode demonstrated an onset potential of 0.91 V vs RHE at 1 mA cm^{-2} , while the $\text{pn}^+\text{-GaInP}_2/\text{PtRu}$ demonstrated an onset potential of 0.83 V vs RHE . The light-limited current density attained by the $\text{pn}^+\text{-GaInP}_2/\text{MoS}_2$ photocathode was $\sim 13 \text{ mA cm}^{-2}$ while the $\text{pn}^+\text{-GaInP}_2/\text{PtRu}$ photocathode achieved $\sim 15 \text{ mA cm}^{-2}$. The $\text{pn}^+\text{-GaInP}_2/\text{MoS}_2$ photocathode was able to maintain its light-limited current density for 100 h without noticeable degradation, at which the point the experiment was terminated, and the sample was immediately transferred to a glove box and packaged air-free for post-analysis. The $\text{pn}^+\text{-GaInP}_2/\text{PtRu}$ photocathode was able to maintain its light-limited current density for $\sim 6 \text{ h}$ at which point the device exhibited a sharp drop in current density indicating failure. (Figure 4b).

Catalytic onset

The onset potential is defined here as the potential at which the photocathode is able to begin catalyzing hydrogen production at an appreciable rate (i.e. 1 mA/cm²). The pn⁺-GaInP₂/MoS₂ and pn⁺-GaInP₂/PtRu photocathodes demonstrated onsets that are among the best for a single absorber PEC device to date⁵. (Figure 4a, 6) This performance was anticipated for the pn⁺-GaInP₂/PtRu photocathode given that it matches the paradigm common to the highest efficiency devices of a high quality III-V semiconductor coupled to a Pt-based catalyst^{5, 10}. The more counterintuitive result is the ~80-mV improved onset potential for the pn⁺-GaInP₂/MoS₂ photocathode since the intrinsic dark catalytic onset of MoS₂ for HER is known to be worse than that of PtRu^{33, 34, 35, 36}. (Figure S1)

To help explain this result, transient photoreflectance (TPR) spectroscopy was used to probe the carrier dynamics at the surface of these photocathodes⁹. The prototypical TPR spectra for pn⁺-GaInP₂, pn⁺-GaInP₂/PtRu and pn⁺-GaInP₂/MoS₂ are recorded at delay time of 5 ps and compared in Figure 5. All three spectra show sharp oscillatory peaks near the bandgap (1.8 eV), highlighted by the red shaded region. These spectral features suggest the presence of a strong surface electric field that causes oscillations in reflection spectra near the bandgap due to the Franz-Keldysh effect, and thus these oscillations are known as the Franz-Keldysh oscillations (FKO)⁹. After optical excitation, the separation of photocarriers driven by the surface field can partially screen the surface field, resulting in the oscillatory TPR spectra. In addition to these oscillatory features, a broad positive peak above the bandgap, near 1.96 eV, is present in the TPR spectra of pn⁺-GaInP₂ and pn⁺-GaInP₂/PtRu, indicated within the blue shaded regions. This broad band feature has

been attributed to the presence of electrons within the surface region of GaInP₂ through the band filling effect and bandgap renormalization⁹. This feature is also present in the TPR spectra for a related system, p-GaInP₂ (Figure S2), however this system exhibits a shorter lifetime (<1ns) due to recombination and diffusion processes leading to depopulation of surface carriers^{37, 38}. The longer-lived photo-generated electrons at the surface of pn⁺-GaInP₂ and pn⁺-GaInP₂/PtRu suggest lower surface recombination rates and slower carrier diffusion into the bulk for GaInP systems with a pn⁺ junction. Combined with the surface electric field evidenced by the FKO⁹, we can conclude that the electrons in pn⁺-GaInP₂ and pn⁺-GaInP₂/PtRu must be localized in a potential-well-like energy band induced by band bending near the surface while the holes are spatially separated from electrons (Figure 6). Interestingly, the same broad band feature near 1.96 eV is barely visible in the TPR spectra of pn⁺-GaInP₂/MoS₂, even at short delay times, implying the absence of the well-like band bending (Figure 6) which correlates to improved electron transfer to MoS₂ at the interface.

As the TPR results indicate more efficient charge separation in pn⁺-GaInP₂/MoS₂ relative to pn⁺-GaInP₂/PtRu, greater photovoltage would be expected in the pn⁺-GaInP₂/MoS₂ system, which could explain the better photocurrent onset potential despite MoS₂ having lower intrinsic catalytic activity than PtRu. We see that for this pn⁺-GaInP₂ semiconductor system, a conformal catalyst layer is important, demonstrating a potential advantage of continuous, 2D catalysts such as MoS₂ over metal catalysts, such as PtRu, that do not readily wet surfaces as necessary to achieve continuous coverage. Poor wetting characteristics require greater thickness for complete coverage, leading to

parasitic absorption/reflection losses by a metal catalyst. This finding may be relevant to other photoelectrodes with catalytic overlayers as there appears to be a fundamental and important dependence of performance on the catalyst conformality. This study only begins to reveal the complexity of this catalyst-semiconductor interface; we encourage additional investigations of various conformal and non-conformal catalysts fabricated with different semiconductors and characterized with photoelectrochemical methods (aqueous and non-aqueous) combined with advanced interface-focused techniques such as TPR.

Light-limited current density

The light-limited photocurrent density represents the yield of incident photons that are converted into current. Sufficiently thick GaInP₂ that has no reflection or recombination losses could achieve a maximum light-limited photocurrent density of ~19 mA cm⁻² under AM1.5G illumination^{17, 39}. Previous work done on a GaInP₂ PEC system has also established that Fresnel normal incidence reflectance can contribute to a 20-30% loss in maximum achievable external quantum efficiency³⁹. The ~15 mA cm⁻² (~18% STH) attained by the pn⁺-GaInP₂/PtRu photocathode indicates that this device is subject to some reflective and parasitic absorptive losses^{17, 40} (Figure 4a) which is expected given that no antireflective or light-trapping strategies were employed. Despite these losses, if paired with the proper photoanode, this photocathode could be the top absorber in an ~18% STH device. If an antireflective coating or other light trapping mechanism is used, an STH efficiency around ~20% could feasibly be achieved making this photocathode

attractive for inclusion in an unassisted tandem PEC device. The pn^+ -GaInP₂/MoS₂ photocathode demonstrated a slightly smaller light-limited photocurrent of $\sim 13 \text{ mA cm}^{-2}$, indicating that the Mo/MoS₂ layer may contribute to a greater amount of parasitic absorption (Figure 4a). This increased absorption is expected given the thicker and more conformal nature of the Mo/MoS₂ film. Previous absorption experiments on Mo/MoS₂ protected photocathodes have shown that the Mo/MoS₂ film can absorb as much as 40% of the incoming light across the spectrum^{16, 17}. Even with this high parasitic absorption, pairing this photocathode with the right photoanode could create a device a $\sim 16\%$ STH efficiency. Future work will focus on improving the light-limited current density through strategies such as anti-reflective coatings, light-trapping mechanisms, and thinning of the protection layers.

Photocathode Stability

The pn^+ -GaInP₂/MoS₂ photocathode demonstrated over 10x better stability than the pn^+ -GaInP₂/PtRu photocathode (Figure 4b). The XPS spectra collected after the CA measurements, shown in figures 2 and 3, can explain why the pn^+ -GaInP₂/PtRu photocathode failed catastrophically after 6 hours while the pn^+ -GaInP₂/MoS₂ photocathode retained its activity after 100 hours. The pn^+ -GaInP₂/PtRu photocathode, after testing, shows peaks in Pt4f, Ru 3p, Ga 2p, In 3d, P 2p and As 3p regions. The Pt 4f and Ru 3p doublets are consistent with a metallic PtRu catalyst indicating that the catalyst remains reduced on the surface^{25, 26, 27}. The peaks in the Ga 2p region have gained shoulders and can be deconvoluted to show the formation of an oxide that likely formed from corrosive contact with the acidic electrolyte²⁷. The In 3d and P 2p regions show

peaks relatively unchanged from spectra taken before testing, except for the appearance of As 3p peaks⁴¹. These peaks indicate that the GaInP₂ semiconductor layer corroded away so extensively as to expose the GaAs substrate underneath. The exposed GaAs substrate most likely created significant shunt and recombination pathways that contributed to the abrupt loss of light-limited photocurrent density⁴².

The pn⁺-GaInP₂/MoS₂ photocathode after testing only shows peaks in the Mo 3d and S 2p regions with no evidence of Ga, In, P, or As peaks. (Figure 3) In the Mo 3d region, the peaks can be deconvoluted into 2 doublets indicating that Mo exists in 2 different states: a reduced metallic state consistent with Mo metal and a 4+ state consistent with MoS₂^{13, 24}. The peaks corresponding to Mo in the 6+ state consistent with MoO₃ are no longer present which is expected given that MoO₃ is unstable in sulfuric acid and dissolves¹³. The S 2p region shows a peak corresponding to sulfur in the 2- state consistent with MoS₂ and a new peak corresponding to S in the 6+ state consistent with SO₄²⁻. This new peak is most likely due to residual sulfuric acid from the electrolyte²⁴. The absence of peaks in the Ga 2p, In 3d, P 2p, and As 3p regions indicates that the substrate underneath the MoS₂ protection layer has not been exposed over the course of this test. The pn⁺-GaInP₂/MoS₂ photocathode, after 100 hours of continuous operation in 3M sulfuric acid, appears to be in nearly pristine condition.

Comparing performance

When studying a photocathode for a dual-junction tandem absorber device, several important metrics for evaluating performance include the onset potential for the HER, the

light-limited current density, and the lifetime of the device. A better onset potential for a photocathode is a proxy for efficiency, and is critical because it relaxes the onset potential requirements for the photoanode, which involves the more sluggish kinetics of water oxidation, and/or allows the use of lower bandgaps for higher photocurrents and thus higher STH efficiency²⁰. Light-limited current density and lifetime each provide useful information about a photocathode, with a key metric resulting from these two properties: the amount of hydrogen the photocathode can produce before failure. A high light-limited current density is not useful if the photocathode fails rapidly, and, conversely, a long lifetime is not valuable if the light-limited current density is too small—in both cases, very little hydrogen is produced. Multiplying these two properties together results in the combined metric of charge passed, a measurable value which directly correlates to total hydrogen produced.

Figure 7 compares the onset potential at 1 mA cm⁻² vs RHE and the demonstrated charge passed for high-performing single absorber photocathodes with various thin-film protection schemes. This figure puts the different photocathodes that have been studied by researchers in this field in within context to help elucidate trends and useful insights for future device improvement. We see that crystalline silicon photocathodes are able to pass the largest amount of charge before failure but are unable to achieve high onset potentials even with precious metal catalysts. Conversely, III-V photocathodes tend to achieve high onset potentials but are unable to pass as much charge before failure. This trend is the basis for the strategy employed in this work: the charge passed by a pn⁺-III-V photocathode could be improved by leveraging an MoS₂ protection scheme previously

shown to work on other systems^{15, 16, 17}. Figure 7 also shows that most of the high performing devices, in terms of both onset potential and demonstrated charge passed, contain precious metal cocatalysts. The exception to this trend is the pn^+ -GaInP₂/MoS₂ photocathode presented in this work. This photocathode used an earth-abundant cocatalyst, MoS₂, to achieve the highest reported onset potential for any single absorber III-V photocathode with or without precious metal catalysts. The MoS₂ thin-film also functioned as a very effective protection layer enabling the photocathode to also pass the most charge of any single absorber III-V photocathode reported to date.

As this MoS₂ protection scheme has been used successfully on semiconductors such as silicon and GaInP₂, this work further highlights the potential for MoS₂ nanomaterials to be used as catalytic thin-film protection schemes for many different water splitting semiconductors that are unstable in acid^{15, 16, 17}. Successful stabilization of other III-V semiconductors could enable a multitude of highly efficient dual-absorber unassisted water splitting devices that can split water with high durability.

CONCLUSION

The pn^+ -GaInP₂/MoS₂ device presented in this work is among the most active and stable single-absorber photocathodes for solar hydrogen production from water splitting. TPR spectral studies provided fundamental insights into the device's high performance, revealing that a conformal MoS₂ catalyst interface imparts improved electron transfer in addition to catalytic activity compared to a nonconformal PtRu catalyst. This insight on semiconductor-catalyst interfaces is potentially relevant to catalytically protected

photoelectrodes across the field. As the $\text{pn}^+\text{-GaInP}_2/\text{MoS}_2$ photocathode presented in this work has a better onset and is significantly more stable than a sister $\text{pn}^+\text{-GaInP}_2/\text{PtRu}$ photocathode, it demonstrates that a precious metal catalyst may not be a requirement for achieving high efficiencies in a photocathode. In fact, due to the substantial stability and interfacial properties it imparts, an MoS_2 catalytic protection layer may even be preferred. Future efforts to apply these MoS_2 protection layers to dual-junction tandem absorber devices offer the opportunity for highly efficient unassisted water splitting over long periods of time. These devices will enable sizeable quantities of hydrogen to be produced and help transition solar-water splitting from the lab-scale to the prototype-scale.

METHODS

Device Synthesis

The epitaxial GaInP_2 films were grown by ambient pressure organometallic vapor-phase epitaxy on degenerately Zn-doped $\text{GaAs}(100)$ substrate pieces with 4° offcut toward (111)B. The substrates were precut to 20 mm x 30 mm and etched for 60 seconds in a dilute $\text{NH}_4\text{OH}:\text{H}_2\text{O}_2:\text{H}_2\text{O}$ solution (1:2:10 by volume) before growth. A 1- μm thick Zn-doped GaInP_2 layer was grown first, with a nominal p-type carrier concentration of 10^{17} cm^{-3} . Then, the dopant was switched to Se to grow a 25-nm thick layer of n-type GaInP_2 with a nominal carrier concentration of $\sim 7 \times 10^{18} \text{ cm}^{-3}$.²³ Following this, a 200 nm GaAs layer was grown and selectively etched off with an ammonium hydroxide (NH_4OH), hydrogen peroxide (H_2O_2), and water mixture (H_2O) in a 2:1:10 ratio. The GaAs contact layer was included to replicate the layer structures and post processing of inverted metamorphic multijunction devices (IMMs), the devices we plan to work on in the future²³. After growth, Au ohmic contacts were electroplated to the back of the substrate.

MoS₂ deposition

The part of the wafer to be coated with MoS₂ was diced and transferred to a DC magnetron sputter coater. A thin layer of Mo metal was deposited onto the wafer at a rate of 7.2 nm min⁻¹ for 30 seconds to achieve a nominal thickness of 3.6 nm. Immediately after Mo deposition, the wafer was transferred to a tube furnace and sulfidized in 90%H₂/10%H₂S held at 150 °C for 1 h to create the MoS₂ layer.

PtRu deposition

The PtRu was deposited via flash sputtering²³, in which the sample was passed briefly (< 1 s duration) underneath the sputter head to achieve very low PtRu loading (~500 ng/cm²).

Physical and Chemical Characterization

X-ray photoelectron spectra of the pn⁺-GaInP₂/PtRu and pn⁺-GaInP₂/MoS₂ photocathodes before and after testing were collected using a Phi VersaProbe Spectrometer with an Al Kalpha source. Binding energies were calibrated to the adventitious C 1s peak at 284.6 eV. SEM cross-sectional images were obtained using an FEI Magellan XHR microscope operated with the assistance of Pong Chakthranont. Cross-sectional images were taken instead of surface images because both samples appeared as blank gray surfaces since the Mo/MoS₂ layer is flat and conformal and the PtRu nanoparticles were too small to readily resolve by SEM.

Electrochemical Characterization

To make photoelectrodes, the sample was cleaved into about 7 mm x 7 mm pieces, with the MoS_x- and PtRu-modified sample edges trimmed off to prevent the possibility of shorting. The PEC characterization and durability testing were performed in a compression cell described elsewhere³¹ that facilitated the post-durability analyses. The electrolyte was 3M H₂SO₄ with 1 mM Triton X-100 (both OmniTrace® EMD Millipore) surfactant added. The Pt foil (Premion® 99.997%) counter electrode (1 cm x 2.5 cm) was housed in a glass tube with medium-porosity glass frit end that was filled with the same electrolyte but without surfactant to mitigate counter electrode fouling and solution yellowing (same reference). A Koslow Scientific Co. mercury/mercurous sulfate (Hg/Hg₂SO₄) reference electrode (MSE) with 3M H₂SO₄ filling solution was used (reference potential of 0.634 V vs. normal hydrogen electrode). Illumination was provided by a 250 W quartz tungsten halogen lamp with water-filled IR filter and light-shaping diffuser (Newport). The illumination intensity was set to match the AM1.5G calibration current of a 1.81 eV bandgap GaInP₂ reference cell. Current-voltage measurements were taken at a 20 mV/s scan rate from negative to positive potential while blocking/unblocking the light source at 0.1 V intervals twice to simultaneously measure the dark current. The scan would begin in the light limited current density region and the electrolyte in the vicinity of the working electrode would rapidly saturate with H₂ while scanning towards the onset potential. Durability testing was performed at constant potential of -0.3 V vs MSE (0.334 vs RHE) while monitoring photocurrent density. For post-analysis, the

compression cell was transferred to an Ar-filled glove box immediately after stopping the durability test, where the sample was removed from the cell, rinsed with DI water, dried with Ar, and vacuum packaged in Ar. After characterization with XPS, no additional chronoamperometry or voltammetry was performed since the characterization process (rinsing, pumping cycles, etc.) can accelerate failure.

Transient Photoreflectance Characterization

Femtosecond TPR spectroscopy is based on a regeneratively amplified Ti:sapphire laser system. The 800nm output beam is ~ 4 mJ per pulse with 1 kHz repetition rate. The pump for TPR is generated by doubling the frequency of the fundamental pulses (~ 1.5 mJ/pulse) in a BBO crystal, and it is then chopped at a frequency of 500 Hz. The pump power is attenuated by neutral density filter wheels. The broadband probe pulses (420-830 nm) are generated by focusing 800nm beam into a sapphire crystal. The probe pulses are delayed in time with respect to the pump pulses using a motorized translation stage. The pump and probe are spatially overlapped on the surface of the sample, and the reflected probe pulses are directed to the multichannel CMOS sensors by optical cables. The beam spot diameters on the sample for probe and pump are 250 μm and 1.4 mm, respectively. The total pump photon flux is determined by measuring the pump power after a pinhole with radius of 200 μm at the sample position. The input photon flux is obtained by subtracting the reflected photon flux from the total photon flux. The carrier density is calculated as the ratio of input photon flux to the penetration depth that is $1/\alpha$ over absorption coefficient at the excitation photon energy.

ASSOCIATED CONTENT

Supporting Information.

Additional details on the dark catalytic performance and the transient photorefectance measurement.

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Notes

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FIGURES

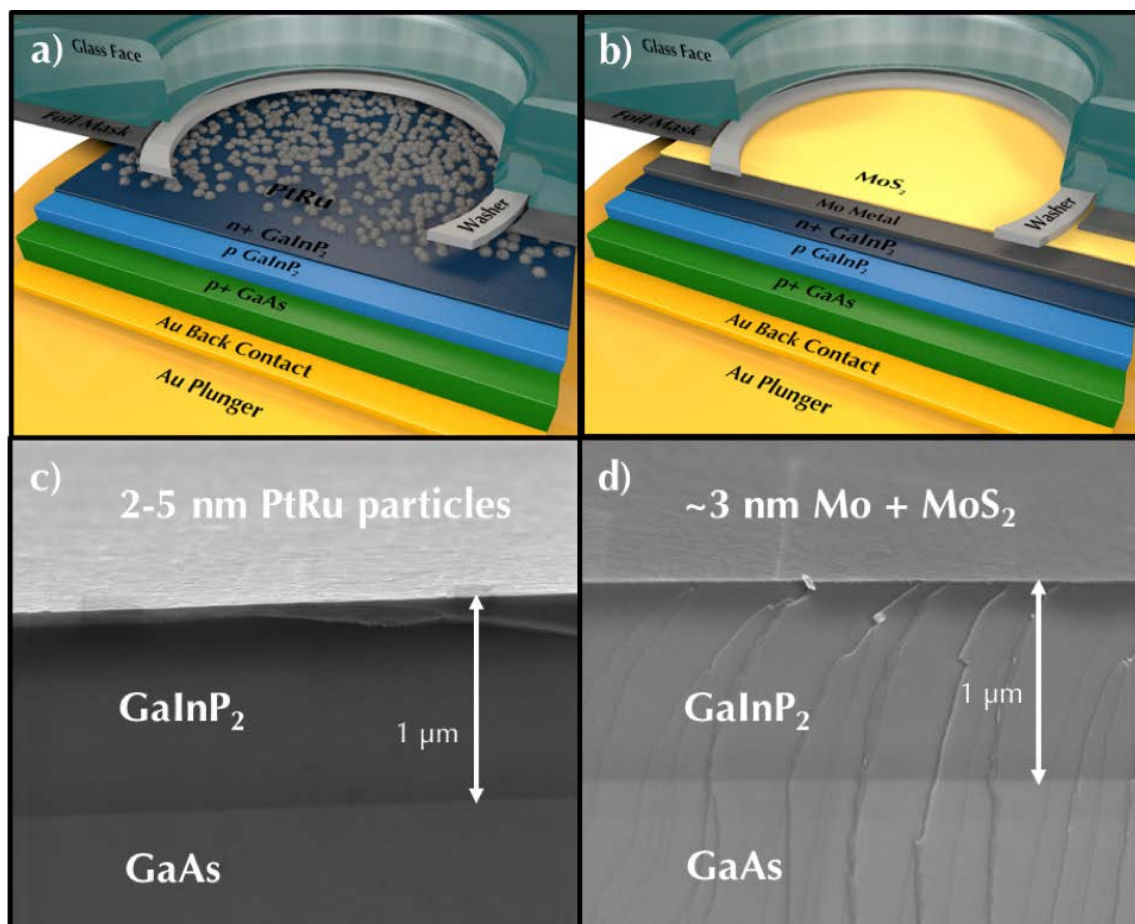


Figure 1. Photocathode device structures. Schematic of a fully synthesized a) pn^+ -GaInP₂/PtRu and b) pn^+ -GaInP₂/MoS₂ photocathode being tested in a compression cell. Cross-sectional SEM images of the surface region of c) pn^+ -GaInP₂/PtRu and d) pn^+ -GaInP₂/MoS₂ photocathode.

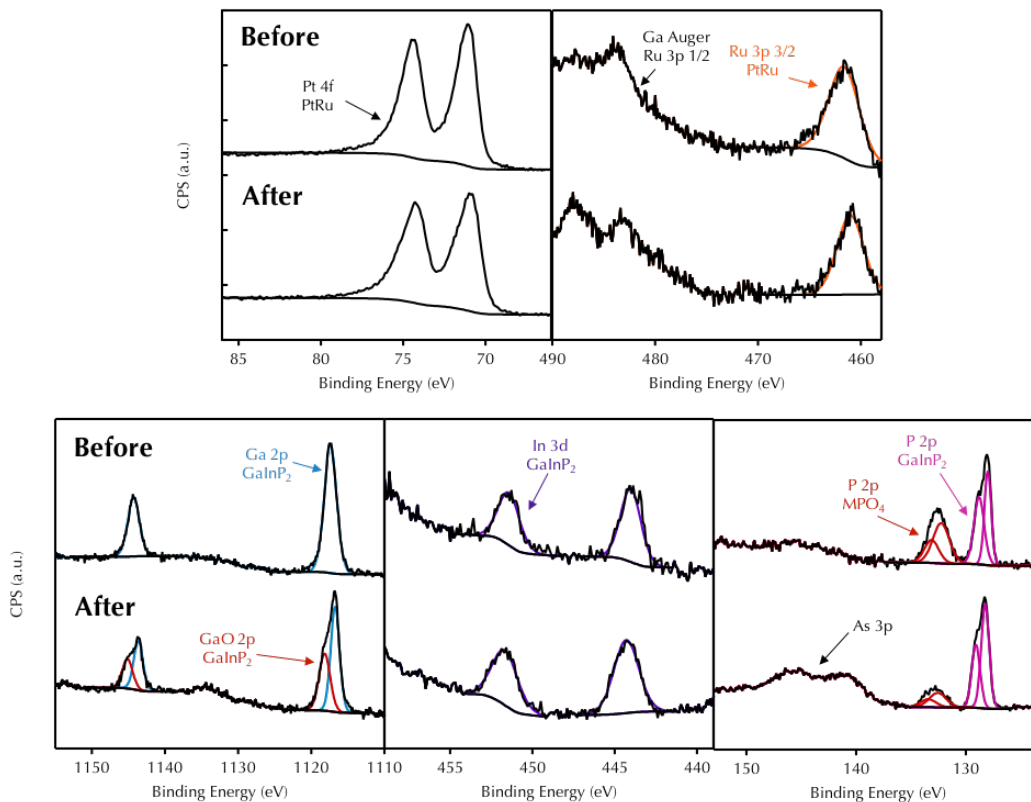


Figure 2. XPS measurements of a $\text{pn}^+\text{-GaInP}_2/\text{PtRu}$ photocathode before and after electrochemical LSV and CA testing. Before testing, the surface of the device consists of Pt and Ru in a reduced metallic state, Ga and In in the 3+ oxidation state consistent with GaInP_2 , and P in a 3- oxidation state corresponding to GaInP_2 , and a 5+ oxidation state corresponding to a metal phosphate. After testing, a Ga surface oxide has grown, and the underlying GaAs substrate has been exposed due to the appearance of an As 3p peak.

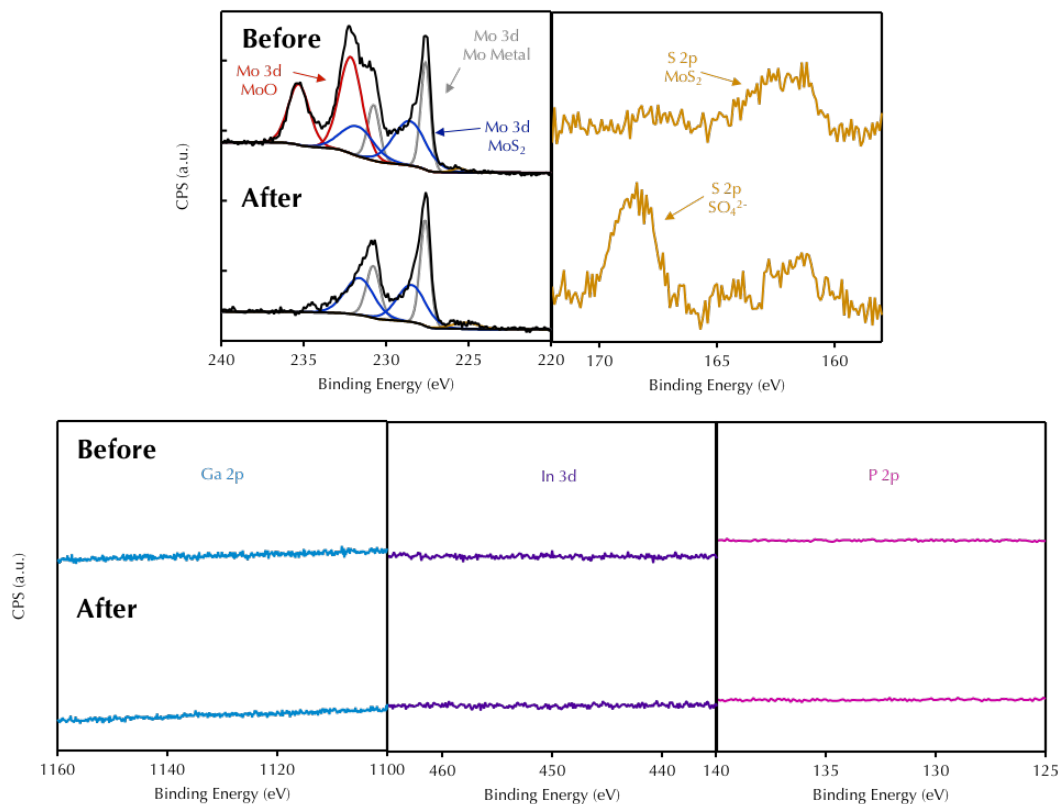


Figure 3. XPS measurements of a $\text{pn}^+\text{-GaInP}_2/\text{MoS}_2$ photocathode before and after electrochemical LSV and CA testing. Before testing, the surface of the device consists of Mo in a reduced metallic state consistent with Mo metal, a 4+ state consistent with MoS₂, and a 6+ state consistent with MoO₃ and S in a 2- state consistent with MoS₂. After testing, the only change is the absence of peaks corresponding to Mo in a 6+ state corresponding to MoO₃, indicating that the surface remains relatively pristine.

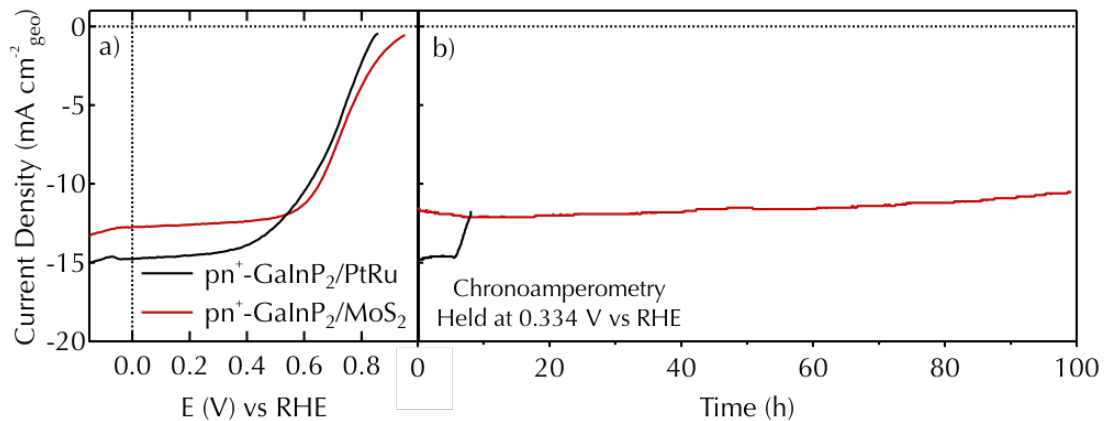


Figure 4. Electrochemical characterization of the $\text{pn}^+\text{-GaInP}_2/\text{PtRu}$ and $\text{pn}^+\text{-GaInP}_2/\text{MoS}_2$ photocathodes in 3M sulfuric acid. a) LSV collected prior to stability testing b) CA measurement taken at a constant potential of 0.334 V vs RHE.

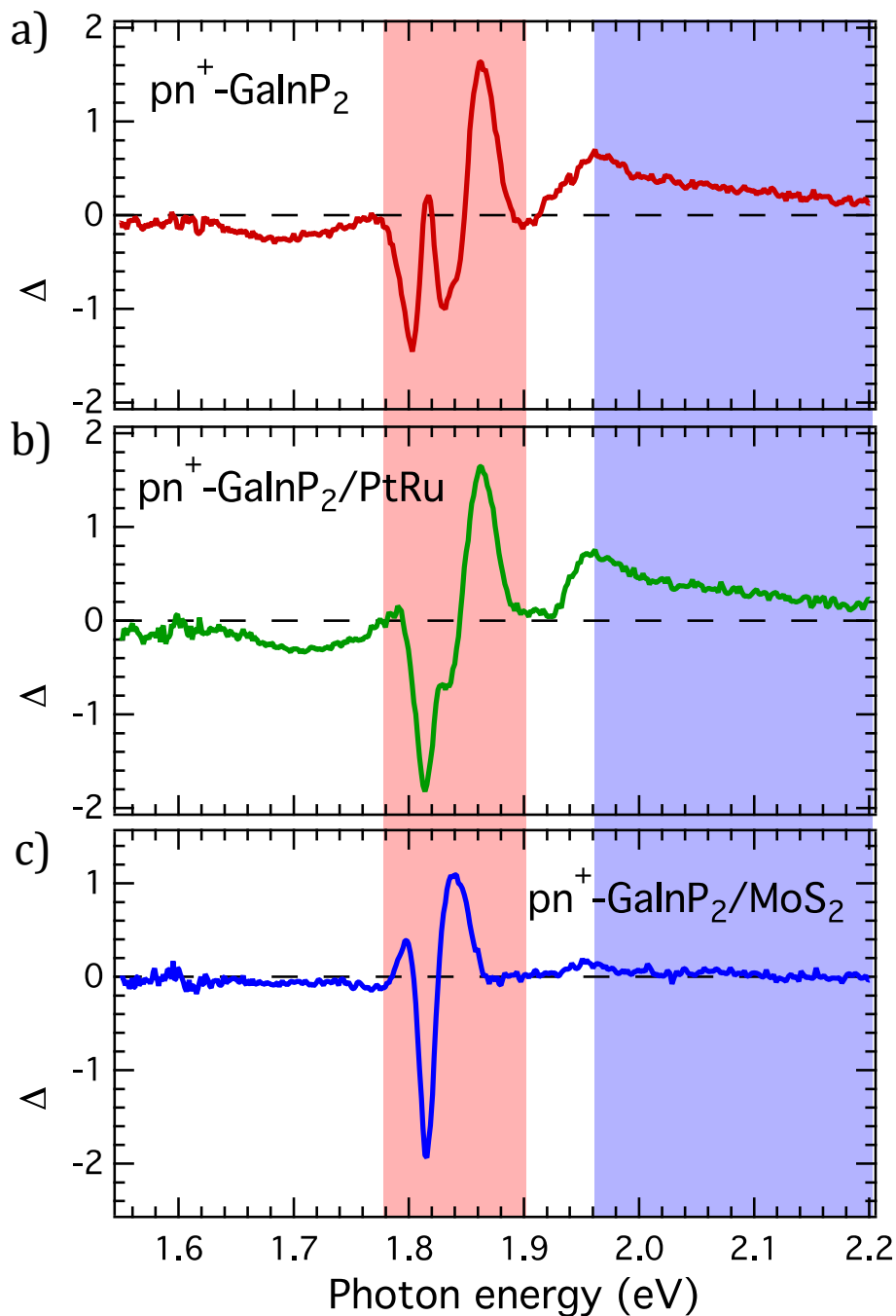


Figure 5. TPR spectra of the samples. These TPR spectra are recorded at 5 ps delay time. The y-axis and x-axis represent the photo-induced reflection change and probe photon energy. The red and blue shade areas indicate the FKO and surface electron induced reflection change, respectively.

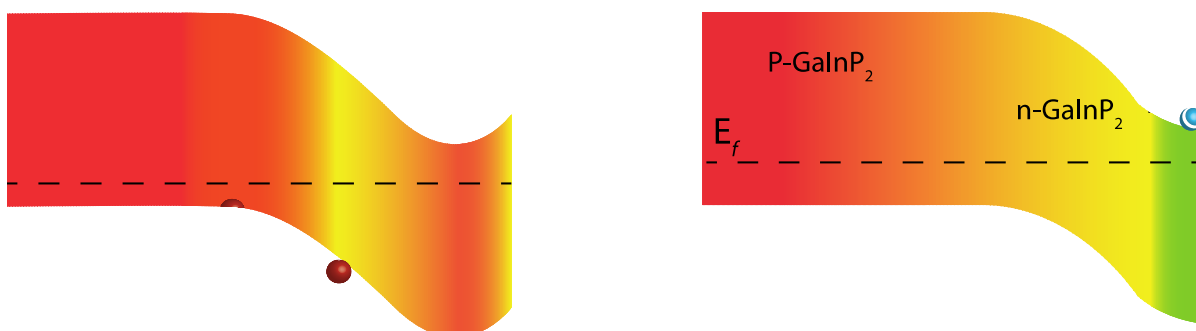


Figure 6. Illustrations of the proposed energy bands for pn^+ -GaInP₂ and pn^+ -GaInP₂/MoS₂. For pn^+ -GaInP₂, the surface electric field bends the band up near the surface, leading to an energy valley that traps electrons. For pn^+ -GaInP₂/MoS₂, the n-MoS₂ contact removes the surface states of n-GaInP₂ and avoids forming the energy valley, and thus the photo-generated electrons in GaInP₂ are quickly injected into the MoS₂ layer.

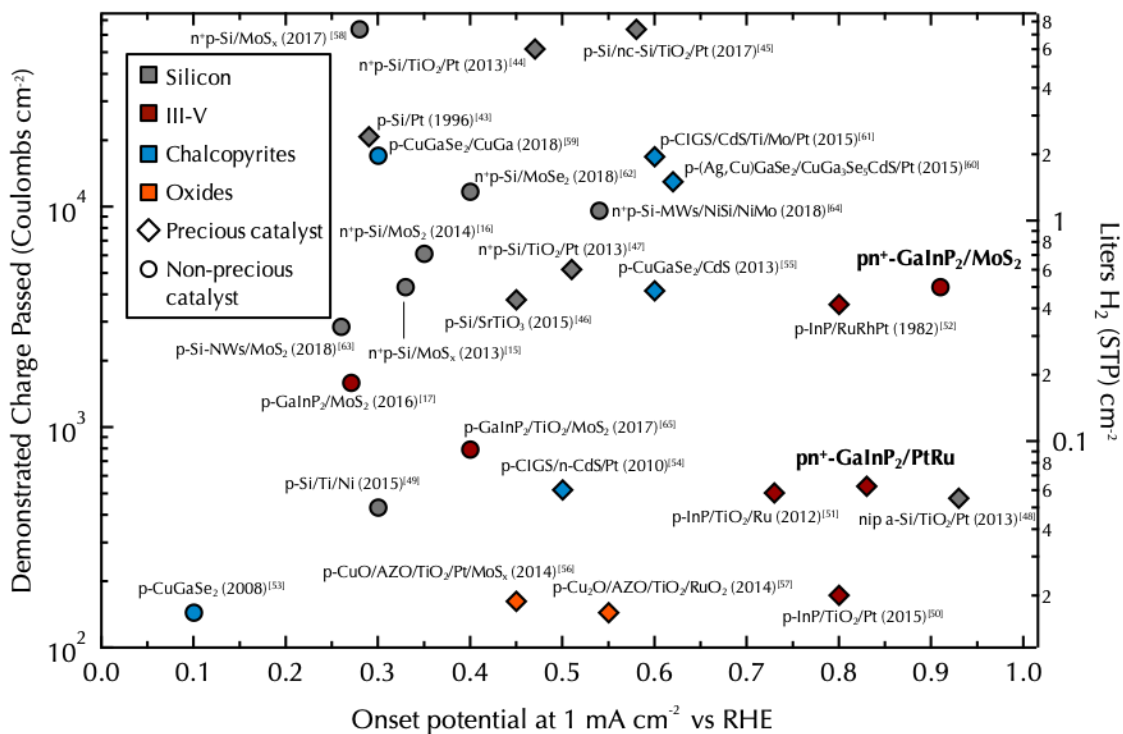


Figure 7. Comparison of the onset potential at 1 mA cm^{-2} vs RHE and the demonstrated charge passed for high-performing single absorber photocathodes with various thin-film protection schemes. The photocathodes with the highest reported activities and stabilities using silicon, III-V, chalcopyrite, and oxide light absorbers were included and delineated by the use of precious and non-precious metal cocatalysts. The pn^+ -GaInP₂/PtRu and pn^+ -GaInP₂/MoS₂ photocathodes (labeled in bold) are from this work. The demonstrated charge passed was calculated by combining the reported light-limited current density and the reported stability. The liters of H₂ at STP was calculated assuming 100% faradaic yield for hydrogen production from the light limited current density. The tabulated data

from this figure can be found in the supplementary information in Table S1.^{15,16, 17, 43, 44,}
45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58,59,60,61,62,63,64,65

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