Enhancing photovoltages at p-type semiconductors through a redox-active metal-organic framework surface coating

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Surface modification of semiconductors can improve photoelectrochemical performance by promoting efficient interfacial charge transfer. We show that metal-organic frameworks (MOFs) are viable surface coatings for enhancing cathodic photovoltages. Under 1-sun illumination, no photovoltage is observed for p-type Si(111) functionalized with a naphthalene diimide derivative until the monolayer is expanded in three dimensions in a MOF. The surface-grown MOF thin film at Si promotes reduction of the molecular linkers at formal potentials >300 mV positive of their thermodynamic potentials. The photocurrent is governed by charge diffusion through the film, and the MOF film is sufficiently conductive to power reductive transformations. When grown on GaP(100), the reductions of the MOF linkers are shifted anodically by >700 mV compared to those of the same MOF on conductive substrates. This photovoltage, among the highest reported for GaP in photoelectrochemical applications, illustrates the power of MOF films to enhance photocathodic operation.
Metal-organic frameworks (MOFs) provide opportunities to build well-ordered three-dimensional structures for the generation of sustainable fuels and chemicals. They combine the advantages of molecular species, with well-defined and precisely tunable active sites, with those of heterogeneous structures, which offer long-term stability and technological readiness. In addition, the porosity and extended architecture of MOFs give access to a high concentration of active sites per surface area. We recently reported a porous interconnectivity of MOFs give access to a high concentration of active sites per surface area. We recently reported a porous interconnectivity of MOFs, a UiO-type MOF (UiO = University of Oslo) built from hexanuclear zirconium organic frameworks (PIZOF), a UiO-type MOF (UiO = University of Oslo) built from hexanuclear zirconium oxide secondary building units (SBUs) and redox-active naphthalene diimide-based (NDI) linkers. Electrochemical studies of Zr(NDI) MOF thin films grown on fluoride-doped tin oxide (FTO) showed that charge propagation through the MOF occurs by a diffusion-like linker-to-linker hopping mechanism, with an apparent diffusion coefficient of $(5.4 \pm 1.1) \times 10^{-11}$ cm$^2$ s$^{-1}$ using KPF$_6$ as the electrolyte. The cyclic voltammograms (CVs) of the Zr(NDI) MOF thin films feature two reversible electrochemical reductions assigned to the NDI$^{0/-}$ and NDI$^{1/-2}$ couples.

In photoelectrochemistry, solar energy is harvested to power redox reactions, capitalizing on absorbed photons to drive those reactions at a lower energetic cost (Fig. 1). The diminished requirement for externally applied voltage is balanced by the photovoltage of the semiconductor, $V_{ph}$, set by the difference between the two quasi-Fermi levels associated with the energy of the holes, $E_{F,n}$ and electrons, $E_{F,p}$, generated under illumination at wavelengths sufficient to excite electrons from the valence to the conduction band. However, the actual photovoltage values often fall short of the maximum obtainable photovoltage due to detrimental recombination events, including surface recombination processes ($I_{tr}$) that compete with productive interfacial charge transfer ($I_{hr}$).

Successful transport of minority charges across the interface is highly sensitive to surface functionalization and attachment strategies. The presence of a redox acceptor in proximity to a photocathode does not guarantee transfer of photogenerated charges, as charge recombinations processes must be overcome to translocate the minority carriers across the interface. For example, in a study of the effects of the linker between a cobaloxime catalyst and TiO$_2$-coated Si(100) wafers, no evidence for the reduction of the molecular species in solution at an illuminated photocathode was found, even though the same species was shown to be reduced when immobilized at the photoelectrode. In another example, immobilization of a nickel–phosphine catalyst at a methylenated Si(111) surface increased the photovoltage of the functionalized semiconductor by 200 mV compared to the Si photocathode when measured in the presence of the catalyst in solution. In both cases, the increased performance upon immobilization was tentatively assigned to improved interfacial charge transfer to the immobilized species.

Atomic layer deposition (ALD) of transparent oxide coatings such as TiO$_2$ has been widely used to slow charge recombination and protect semiconductors against corrosion in photoelectrochemical applications. Formation of a p–n junction at p-GaP using an ALD-deposited layer of n-Nb$_2$O$_5$ and a Pt catalyst resulted in a record photovoltage of 710 mV, an improvement of 360 mV compared to a GaP photocathode with Pt only and no metal oxide layer. Molecular functionalization of semiconductors can also lower barriers for photoelectrochemical reactions. In the context of metal–organic hybrid constructs, Downes and Marinescu demonstrated photoelectrochemical H$_2$ evolution in H$_2$SO$_4$ solution (pH 1.3) by a cobalt dithiolene polymer composing a metal–organic surface (MOS) dropcast onto p-type Si electrodes. In this study, the electrochemical potential required to achieve equivalent rates of H$_2$ evolution was 550 mV less under illumination than that of the MOS at glassy carbon electrodes. More recently, a Cu-based MOF at a Cu$_2$O photodecide operated in acetonitrile was used to enhance the CO$_2$ reduction activity of the Cu$_2$O surface.

While these reports show an increase in performance of the MOF-modified semiconductors, it is somewhat unclear whether the improvement is due to the catalytic activity of the surface species themselves or to changes in surface energetics that modulate the band edges and/or improve charge separation. In addition, in the case of a 3D material being added to the surface, improvements in catalysis could be due to a higher reaction surface area. To date, clear evidence of photoelectrochemical reduction of a discrete molecular species in a MOF framework has not been reported. Herein, we investigate whether p-type semiconductors can be used to reduce the redox-active linkers that comprise the Zr(NDI) MOF introduced above.

Results
Preparation of Zr(NDI) thin films. ALD-deposited TiO$_2$ has been used as an interface to immobilize molecular catalysts at semiconductors, and in our construct, provides a metal oxide surface for anchoring the carboxylate-functionalized NDI linker. TiO$_2$ was deposited by ALD onto freshly etched semiconductor (SC, where SC is Si or GaP) wafers using 100 alternating TiCl$_4$ and H$_2$O cycles at 200°C, which results in TiO$_2$@SC samples amenable to the attachment of carboxylate groups. The oxide layer was 3.7-nm thick as measured by spectroscopic ellipsometry. The MOF was grown on the TiO$_2$-coated substrate as reported previously for Zr(NDI)@FTO. Briefly, the NDI linkers are first attached to the substrate by exposure of the TiO$_2$-coated...
wafer to a solution of NDI in DMF, resulting in NDI[TiO2]@SC. The NDI layer was 2 nm as measured by spectroscopic ellipsometry and using a Cauchy layer model for the NDI film. This is in good agreement with formation of a monolayer of NDI, which has a length of 18 Å. MOF growth on the NDI-coated wafers proceeds in a premixed and sonicated solution of the NDI linker, ZrCl4, and acetic acid in DMF under direct solvothermal synthesis conditions at 120 °C for 3 days, resulting in Zr(NDI)[TiO2]@SC.

Film characterization. Attenuated total reflection Fourier transform-infrared (ATR-FTIR) spectroscopic studies of the Zr(NDI)[TiO2]@SC show that the characteristic peaks of the Zr(NDI) bulk powder sample are retained (Fig. 2a). In particular, and as described previously,5 frequencies associated with the symmetrical and asymmetrical stretches of the imide C = O bond of the NDI linker (∼1650 and 1710 cm⁻¹, respectively) appear in the carbonyl region of the spectrum. In addition, peaks indicative of the C-O-C stretch (∼1250 cm⁻¹), imide C-N bonds (1350 cm⁻¹), and ring deformation of the imide (∼1420 cm⁻¹) are present in the spectra of the SC-grown MOF films as well as those of the bulk MOF and the NDI linker (Supplementary Fig. 1). Functionalization of the SC surface was also examined by X-ray photoelectron spectroscopy (XPS). Spectra of the TiO2-coated SC wafers have peaks corresponding to the binding energies of Ti2p, O1s, and C1s (Fig. 1b and Supplementary Figs. 2 and 3). After exposure to the NDI solution, additional contributions from the nitrogen moieties (Fig. 1b and Supplementary Figs. 2 and 3). After exposure to the NDI solution, additional contributions from the nitrogen moieties of the NDI linker are present in the N1s region. Following solvothermal synthesis conditions, the intensities of these contributions increases and new peaks appear in the region of Zr3d binding energy, indicative of the Zr SBUs of the MOF. The ratio of Zr:N is 1:1.8 ± 0.2 (Supplementary Table 1), consistent with each of the hexanuclear Zr nodes sharing twelve linkers in a repeating crystalline fashion. Extensive efforts to characterize the films by X-ray diffraction (XRD) either by grazing-incidence XRD or with long-scan measurements (70 s/step) were unfortunately unsuccessful, a fact that we ascribe to the thinness of the films and their preferred orientation that precludes observation of XRD. Scanning electron microscopy (SEM), in contrast, provides evidence of the crystallinity of the MOF thin film after directed growth on the titania-coated substrate (Fig. 3 and Supplementary Fig. 5a). Octahedral crystals characteristic of the UiO-type morphology are evident, with a particle size of 100–200 nm. The film thickness is 150 ± 43 nm (where the range arises from sample-to-sample variation) determined by cross sections of the films imaged by SEM (Supplementary Fig. 6). SEM with energy- dispersive X-ray spectroscopy (EDX) data across multiple samples and geometric areas yields a Zr:N ratio of 1:2.0 ± 0.4 (Supplementary Fig. 5b and Supplementary Tables 2 and 3), providing additional evidence for the proposed structure of the film.

Cyclic voltammetry at illuminated silicon photocathodes. The photoelectrochemical response of three modified Si electrodes—TiO2-coated Si (TiO2@Si), those with an NDI monolayer grown on the TiO2-coated Si (NDI[TiO2]@Si), and the complete Zr(NDI) MOF-functionalized electrode (Zr(NDI)[TiO2]@Si)—was tested in a three-electrode configuration using the functionalized Si wafer as the working electrode, a carbon counter electrode, and a Ag wire pseudoreference electrode. In 0.5 M LiClO4 in DMF and under 1-sun illumination, the CV of TiO2@Si has little-to-no photoelectrochemical response at a scan rate of 100 mV s⁻¹ (Supplementary Fig. 7). However, under the same conditions, the photocathode functionalized with a monolayer of the NDI linker, NDI[TiO2]@Si, exhibits two redox events in the CV (Fig. 4a and Table 1). The midpoint potentials are observed at −1.00 and −1.25 V vs. Fe⁴⁺/³⁻, close to those of the NDI linker on FTO-grown MOF (−0.97 and −1.30 V vs. Fe⁴⁺/³⁻). We thus assign these features to two normally ordered one-electron reductions of the NDI linker to NDI⁻ and NDI³⁻ (The Zr nodes solely play a coordinative and structural role in the film, without participating in charge transfer itself, as the Zr(IV)/(III) couple is outside the redox window that we are operating within.). After expanding the monolayer into the third dimension in a MOF thin film (Zr(NDI)[TiO2]@Si), relatively broad reductive and oxidative peaks are observed at potentials more positive than those in the monolayer-functionalized Si sample (E_p = −0.999 and E_a = −0.18 V vs. Fe⁴⁺/³⁻, v = 100 mV s⁻¹, Fig. 4a). At slower scan rates (v = 5 mV s⁻¹), the two reductions separate into distinct features, with midpoint potentials of −0.64 and −0.78 V vs. Fe⁴⁺/³⁻ (Supplementary Fig. 8). The large peak-to-peak separations (0.43 and 0.46 V) are commonly observed in MOF electrochemistry and are attributed to sluggish interfacial electron transfer, ohmic resistance in the film, and/or slow ion transport at the film/electrolyte interface.35–37

Photovoltage of Zr(NDI)[TiO2]@Si. Crucially, the midpoint potentials of the first and second reductions of the NDI linker in Zr(NDI)[TiO2]@Si are shifted positive by 330 mV and 520 mV, respectively, compared to the formal potentials of the Zr(NDI) @FTO construct. In photoelectrocatalytic applications, the open-circuit potential (V_oc) or onset potential (V_on) is typically reported as a measure of the photovoltage, a value that depends on equilibration between the semiconductor and redox couple, as well as the surface recombination rates and overpotential requirements.9,11,38 In the case of NDI[TiO2]@Si, the lack of a photoeffect on the NDI electrochemical potentials suggests poor charge separation at the semiconductor interface, resulting in the need for a large applied bias to increase band bending in order to populate the conduction band with minority carriers. The similar midpoint potentials observed in the dark electrochemistry of the
Electroactive portion of the Zr(NDI) thin film. The electroactive coverage of NDI linkers was calculated from controlled potential electrolysis experiments by stepping the potential from +0.1 to –1.2 V vs. Fe+/0, providing sufficient driving force to fully reduce NDI twice to NDI2– (Supplementary Fig. 10). Analysis of the charge passed using three samples yields an electroactive surface concentration of 68 ± 42 nmol cm−2. For comparison, a tightly packed monolayer of NDI, with the molecule oriented perpendicular to the surface, gives a surface concentration of 1.1 nmol cm−2. Thus, the extended MOF architecture allows for a significantly higher loading of redox-active sites at the electrode. Nonetheless, simply increasing the number of sites may not always result in a current enhancement, as diffusional processes can become limiting as films become thicker. To calculate the diffusion coefficient of the film, the chronoamperometry data were fitted to the Cottrell equation (Eq. (1))

\[
\frac{i}{\sqrt{\pi t}} = \frac{nFSC_{\text{NDI}}D}{\sqrt{\pi t}},
\]

where \(i\) is current (A), \(n\) is the number of electrons, \(F\) is Faraday’s constant (C mol⁻¹), \(S\) is the electrode surface area (cm²), \(C_{\text{NDI}}\) is the concentration of NDI linkers on the surface (mol cm⁻³), \(D\) is the apparent diffusion coefficient (cm² s⁻¹), and \(t\) is time (s). This analysis results in a diffusion coefficient of (1.1 ± 1.2) × 10⁻¹⁰ cm² s⁻¹. For comparison, \(D_{\text{app}}\) for the NDI film at FTO was found to be (5.4 ± 1.1) × 10⁻¹⁰ cm² s⁻¹ using KPF₆ as the electrolyte, and more recent investigations have found that the diffusion coefficient for these constructs can be increased by one order of magnitude using LiClO₄ as the electrolyte (Supplementary Fig. 11). Thus, the apparent diffusion coefficient for the Zr(NDI) MOF film (measured with LiClO₄) is in the same order of magnitude in dark electrochemistry experiments at conducting substrates as on illuminated semiconductors. This provides further evidence for the successful formation of the proposed Zr(NDI) MOF (i.e., not only the molecular identity but also the macroscopic properties of the film are conserved) and indicates that charge transport properties through the film are unchanged, regardless of the substrate or film thickness. The similar apparent diffusion coefficients also suggest that the photocurrents in Zr(NDI)TiO₂@Si are largely determined by the charge transfer characteristics of the MOF.

The amount of the film that is electrochemically probed can be calculated using Eq. (2)\(^{40-42}\)

\[
\delta = \sqrt{\frac{D_{\text{app}}RT}{F\nu}},
\]

where \(\delta\) is the diffusion-layer thickness (cm), \(D_{\text{app}}\) is the apparent diffusion coefficient (cm² s⁻¹), \(R\) is the gas constant (J mol⁻¹ K⁻¹), \(T\) is temperature (K), \(F\) is Faraday’s constant (C mol⁻¹), and \(\nu\) is the scan rate (V s⁻¹). With an apparent diffusion coefficient of 10⁻¹⁰ cm² s⁻¹ and a scan rate of 100 mV s⁻¹, about 50 nm of the film is probed during the timescale of the measurement. At slower scan rates, all of the film can be electrochemically accessed as the diffusion-layer thickness increases, approaching the MOF film boundary to reach a regime of finite diffusion\(^{41}\). In this regime, the current is given by Eq. (3)

\[
i_{pc} = \frac{FSC_{\text{NDI}}d_l}{4RT},
\]

where \(i_{pc}\) is the cathodic peak current (A), \(F\) is Faraday’s constant (C mol⁻¹), \(S\) is the electrode surface area (cm²), \(C_{\text{NDI}}\) is the concentration of NDI linkers on the surface (mol cm⁻³), \(d_l\) is the film thickness, \(\nu\) is the scan rate (V s⁻¹), \(R\) is the gas constant (J mol⁻¹ K⁻¹), and \(T\) is temperature (K). At a scan rate of 5 mV s⁻¹, the diffusion-layer thickness is >200 nm (Eq. (2)), larger than the average film thickness (see above and Supplementary Fig. 6). Under these conditions, the observed current magnitude of −0.19 mA cm⁻² (Supplementary Fig. 8b) is within the theoretical
range of $-0.29 \pm 0.14 \text{ mA cm}^{-2}$, calculated with a finite-diffusion model (Eq. (3)).

Applications of MOF coatings at semiconductors for reductive fuel-forming reactions depend on the ability of the film to not only capture minority carriers but also to transfer those charges to the substrate. Thus, charge transfer through the film was further investigated in the presence of an electron acceptor in the electrolyte solution that is thermodynamically capable of being reduced by electrons from the MOF film. The addition of tris(2,2',bipyridine)cobalt(III) tri(hexafluorophosphate) to the electrolyte solution (1 mM in DMF with 0.5 M LiClO$_4$) results in increased photocurrents under chopped light chronoamperometry conditions (Supplementary Fig. 12a) compared to experiments with no external electron acceptor (The measured photocurrents are due to the underlying SC in the Zr(NDI)|TiO$_2$@SC films, as reference experiments on Zr(NDI)|TiO$_2$ show photocurrents <0.5 $\mu$A/cm$^2$. This illustrates that photogenerated charges diffuse through the entire redox-active film and can be extracted at the film|solution interface. Stirring the solution does not further increase the photocurrent (Supplementary Fig. 12b), indicating that the photocurrent is not limited by a diffusional concentration gradient of the Co(III) acceptor at the film|solution interface. The importance of the Zr (NDI) MOF in the experiments is evident by a >300% increase in photocurrent of the MOF-modified samples compared to those of NDI-only modified photocathodes (Fig. 4b). Spikes in the chronoamperogram upon illumination are characteristic of charge transfer to surface states, followed by a decay to steady-state current that reflects the ratio of interfacial charge transfer ($j_0$) to recombination ($j_\infty$) $^{13}$. The appearance of transient photocurrents evident under chopped light illumination suggests that the photovoltage could be further increased through engineering of future MOFs that focus on slowing recombination processes at the interface.

**Photovoltage of Zr(NDI)|TiO$_2$@GaP.** To demonstrate the broader applicability of MOF thin films as surface coatings to increase photovoltage, we extended this work to p-type GaP, a III–V semiconductor with conduction and valence band positions that fulfill the energetic requirements for both hydrogen and oxygen evolution. The photoelectrochemical responses of TiO$_2$@GaP, NDI|TiO$_2$@GaP, and Zr(NDI)|TiO$_2$@GaP were evaluated. In a 0.5 M LiClO$_4$ DMF solution under 1-sun illumination, TiO$_2$@GaP and NDI|TiO$_2$@GaP samples show similarPEC responses (Supplementary Fig. 11). In the dark, little-to-no current passes, while upon illumination, a slight increase in current is observed at negative potentials ($-0.8$ V vs. Fc$^{+/0}$). In contrast, the MOF-modified samples show two clearly defined redox waves upon illumination, with midpoint potentials at remarkably positive values of $-0.26$ and $-0.50$ V vs. Fc$^{+/0}$ (Fig. 5 and Table 1). The first and second reductions are shifted anodically by 710 mV and 800 mV, respectively, compared to the formal potentials of Zr(NDI)|FTO (Supplementary Figs. 14 and 15). These results illustrate that the Zr(NDI) film effectively captures photogenerated charges at the MOF|GaP interface, with the semiconductor playing an integral role in harnessing solar energy to drive the reduction of the NDI linkers. In addition, the two one-electron reductions confirm that the molecular identity of the MOF-immobilized NDI linkers is preserved. Finally, the Zr (NDI)|TiO$_2$ coating also provides ample protection against hydrolytic degradation, a crucial consideration in solar fuel applications. When tested in aqueous conditions (pH 8.5), the NDI-based reductions are still visible with similar photovoltage, but now as one wave due to potential inversion of the two processes as observed earlier in Zr(NDI)|FTO in aqueous electrolyte (Supplementary Fig. 17)$^9$. Photovoltages of up to 710 mV have previously been reported for GaP in photoelectrochemical applications, and have been shown to depend on a variety of factors, including the nature of back contacts, formation of a p–n junction, surface-preparation techniques, and/or surface modification$^9,44$. In our construct, the reduction of NDI-based linkers within a surface-immobilized MOF occurs at potentials over 700 mV positive of the formal potentials at a conductive substrate. This state-of-the-art photovoltage is achieved by employing a redox-active moiety within a MOF thin film, enabling charges at the interface to be efficiently captured and carried away by molecular components, thereby lessening the probability of charge recombination. An extended spatially ordered network of redox-active linkers aids charge separation through localization of charges away from the interface where detrimental charge recombination processes can occur. This results in the highest photovoltage reported for a GaP photocathode modified with only earth-abundant materials.

**Effect of light intensity on photocurrent of Zr(NDI)|TiO$_2$@SC.** The current response at a film-modified photoelectrode may be limited by the number of photons converted to interfacial charge carriers (the light absorption, charge separation, and charge transfer processes)$^{45}$ or by the diffusion of those charge carriers to redox centers. At both semiconductors (SCs, p-type Si, and GaP), a threshold of light intensity must be reached to produce photocurrent, indicating that light is required to reduce Zr(NDI)|TiO$_2$@SC. Remarkably, decreasing the light intensity from 100 mW cm$^{-2}$ (1 sun) to 60 or 10 mW cm$^{-2}$ (for GaP or Si substrates, respectively) does not decrease the photocurrent of the photoelectrodes (Supplementary Figs. 18 and 19). The lack of linear dependence of the peak current to light intensity indicates that the rate of charge propagation through the film is not controlled by photon flux, but by diffusional electron transport through the MOF, as described above.
Discussion
In this report, we have shown that not only are photogenerated charges effectively transported from a visible-light-absorbing semiconductor to the first layer of molecular linkers in a MOF film, but that the extended framework allows those charges to be extracted at a lower applied potential compared to a conductive electrode. We provide unequivocal electrochemical evidence of the interfacial charge transfer of photogenerated electrons from visible-light-absorbing semiconductors to a MOF–confined molecular species, and highlight the role of the MOF in promoting charge separation and improving the photovoltage of the semiconductor–MOF hybrid material. The improved photo-voltage caused by the MOF coating is likely because of the capture and spatial separation of the minority carriers across the film. We show that the charges captured by the film can be translocated through the MOF by a hopping mechanism between discrete molecular moieties, and finally extracted by an electron acceptor in solution. This highlights the promise of this construct to not only accumulate charges at a mild potential, but also to serve as an electron reservoir for further reductive reactions. The lack of photocurrent dependence on light intensity, along with the lack of current increase upon stirring the solution with an external electrode, indicates that the current is not limited by photon flux or mass transport at the film|solution interface. The calculated diffusion coefficient for the film evaluated at an illuminated semiconductor substrate is similar to that of the film grown on a conductive substrate5, suggesting that the current is limited by the same process, i.e., diffusion of charge through the film. Additional evidence that the photocurrent is under diffusion control is found through modeling the current under finite-diffusion conditions.

P-type Si and GaP have previously been used as substrates for hydrogen evolution and carbon dioxide reduction20–23,44,45. Our findings indicate that the advantages of MOFs, such as their high surface area and porosity combined with their ability to house and stabilize molecular catalysts, may be integrated with SCs for MOF-based photoelectrosynthetic applications. Ion-permeable coatings allow key characteristics such as photovoltage to be optimized by forming adaptive junctions at the interface46. We have shown that the construct can be operated in aqueous conditions, a promising feature in the context of light-driven hydrogen evolution. In particular, MOF thin films can serve as novel semiconductor coatings to enhance photovoltages by promoting interfacial charge separation of photogenerated charges.

Methods
General details. All reagents were purchased from commercial sources and used as received, except for DMF, which was stored over activated 3Å molecular sieves. As received, except for DMF, which was stored over activated 3Å molecular sieves.

B-doped Si (111) substrates (ITME) had a thickness of 265 μM and a resistivity of 0.15 Ω cm, a dopant concentration of 5.0 × 1017 cm−3, and an n+–p+ density of 4.5 × 1019 cm−3. Before ALD deposition, Si wafers were cleaned by three successive 5-min sonications in acetone, dichloromethane, and water. The wafers were then etched in ammonium fluoride for 20 min, followed by rinsing with water and storing in an argon environment. GaP wafers were cleaned with a resistivity of 0.16 Ω cm, a dopant concentration of 5.5 × 1017 cm−3, and an n–p density of 5.0 × 1017 cm−3. Before ALD deposition, GaP wafers were cleaned by three successive 5-min sonications in acetone, dichloromethane, and water. The wafers were then etched in ammonium fluoride for 20 min, followed by rinsing with water and storing in an argon environment. Ta2O5 wafer was grown with alternating cycles of the TiCl4 and H2O precursors. The synthesis was performed in a three-electrode configuration using a custom cell (Pine Instruments) equipped with a quartz window. Working electrodes were prepared by scratching the back of the wafer with GaIn eutectic (Sigma-Aldrich). Electrical contact to the stainless-steel back contact was made with a piece of conductive copper tape (Electron Microscopy Sciences). The counter electrode was a glassy carbon rod, and a Ag wire was used as a pseudoreference that was calibrated against Fe/C13 at the beginning and end of every experimental data set. The illumination source was an Eni Tech Solar Simulator with Xe light source and an AM 1.5 G spectral correction filter. The light intensity was calibrated using a NREL-traceable silicon reference cell.

Instrumental details. ATR-FTIR data were collected on a Bruker ALPHA FTIR spectrometer from 4000 to 450 cm−1 at room temperature. XPS was performed on a PHI Quantera SXM using a monochromatic Al Ka source (hv = 1486.6 eV) operated at 25 W with a beam diameter of 100 μM and a pass energy of 224 eV. Data were analyzed using Multipak or Casa software, and all spectra calibrated by setting adventitious carbon to 284.8 eV. SEM was performed with a Zeiss 1550 SEM with InLens detector. The measurements were performed at 4–5 kV accelerating voltage at working distances of 3–5 mm. ImageJ (NIH) software was used to measure the film thickness. EDX data were collected with an 80-nm2 Silicon Drift Detector using AZtec (INCA energy) software at an acceleration voltage of 5 kV and a working distance of 6.5 nm or 8.5 mm.

Photoelectrochemical measurements. Photoelectrochemical measurements were performed in a three-electrode configuration using a custom cell (Pine Instruments) equipped with a quartz window. Working electrodes were prepared by scratching the back of the wafer with GaIn eutectic (Sigma-Aldrich). Electrical contact to the stainless-steel back contact was made with a piece of conductive copper tape (Electron Microscopy Sciences). The counter electrode was a glassy carbon rod, and a Ag wire was used as a pseudoreference that was calibrated against Fe/C13 at the beginning and end of every experimental data set. The illumination source was an Eni Tech Solar Simulator with Xe light source and an AM 1.5 G spectral correction filter. The light intensity was calibrated using a NREL-traceable silicon reference cell.

Data availability
The data that support the findings of this study are available from the corresponding author upon reasonable request.

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References
1. Downes, C. A. & Marinescu, S. C. Electrocatalytic metal-organic frameworks for energy applications. ChemSusChem 10, 4374–4392 (2017).
2. Sun, L., Campbell, M. G. & Dincă, M. Electrically conductive porous metal-organic frameworks. Angew. Chem. Int. Ed. 55, 3566–3579 (2016).
3. Solomon, M. B., Church, T. L. & D’Alessandro, D. M. Perspectives on metal–organic frameworks with intrinsic electrocatalytic activity. CrystEngComm 19, 4049–4065 (2017).
4. Majewski, M. B., Peters, A. W., Wasilewski, M. R., Hupp, J. T. & Farha, O. K. Metal–organic frameworks as platform materials for solar fuels catalysis. ACS Energy Lett. 3, 611 (2018).
5. Johnson, B. A., Blunia, A., Fei, H., Cohen, S. M. & Ott, S. Development of a UiO-type thin film electrocatalysis platform with redox-active linkers. J. Am. Chem. Soc. 140, 2985–2994 (2018).
6. Walter, M. G. et al. Solar water splitting cells. Chem. Rev. 110, 6446–6473 (2010).
7. Bard, A. & Fox, M. A. Artificial photosynthesis: solar splitting of water to hydrogen and oxygen. Acc. Chem. Res. 28, 141–145 (1995).
8. Boucher, D. G., Speller, J. R., Han, R., Osterloh, F. E. & Rose, M. J. Decoupling effects of surface recombination and barrier height on p-Si(111) photovoltaic in semiconductor|liquid junctions via molecular dopes and metal oxides. ACS Appl. Energy Mater. 2, 66–79 (2019).
9. Malizia, M., Seger, B., Chorkendorff, I. & Veszprémi, P. C. K. Formation of a p–n heterojunction on GaP photocathodes for H2 production providing an open-circuit voltage of 710 mV. J. Mater. Chem. A 2, 6847–6853 (2014).
10. Thorne, J. E., Jiang, J.-W., Liu, E. Y. & Wang, D. Understanding the origin of photoelectrode performance enhancement by probing surface kinetics. Chem. Sci. 00, 1–8 (2016).
11. Lewis, N. S. A quantitative investigation of the open-circuit photovoltage at the semiconductor|liquid interface. J. Electrochem. Soc. 131, 2496 (1984).
12. Guirriero, N. et al. Surface modification of semiconductor photoelectrodes. Phys. Chem. Chem. Phys. 17, 15655–15674 (2015).
13. Gong, L. et al. Influence of anchoring groups on the charge transfer and performance of p–Si/TiO2/cobaloxime hybrid photocathodes for photoelectrochemical H2 production. ACS Appl. Mater. Interfaces 11, 54010–54019 (2019).
14. Seo, J., Pekarek, R. T. & Rose, M. J. Photoelectrochemical operation of A Surface-bound, Nickel-phosphine H2 evolution catalyst on p-Si(111): a molecular semiconductor|catalyst construct. Chem. Commun. 51, 13264–13267 (2015).
15. Hu, S. et al. Amorphous TiO2 coatings stabilize Si, GaAs, and GaP photoanodes for efficient water oxidation. Science 344, 1093–1098 (2014).
16. Seger, B. et al. Using TiO2 as a conductive protective layer for photocathodic H2 evolution. J. Am. Chem. Soc. 135, 1057–1064 (2013).
17. Beiler, A. M., Khuninudnova, D., Wadsworth, B. L. & Moore, G. F. Cobalt porphyrin-polyppyridyl surface coatings for photoelectrosynthetic hydrogen production. Inorg. Chem. 56, 12178–12185 (2017).
18. Beiler, A. M., Khusnutdinova, D., Jacob, S. I. & Moore, G. F. Solar hydrogen production using molecular catalysts immobilized on gallium phosphide (111) A and (111) B polymer-modified photocathodes. ACS Appl. Mater. Interfaces 8, 10038–10047 (2016).

19. Khusnutdinova, D., Beiler, A. M., Wadsworth, B. L., Jacob, S. I. & Moore, G. F. Metalloporphyrin-modified semiconductors for solar fuel production. Chem. Sci. 8, 253–259 (2016).

20. Beiler, A. M., Khusnutdinova, D., Jacob, S. I. & Moore, G. F. Chemistry at the interface: polymer-functionalized GaP semiconductors for solar hydrogen production. Ind. Eng. Chem. Res. 55, 5306–5314 (2016).

21. Cedeno, D. et al. Using molecular design to control the performance of hydrogen-producing polymer-brush-modified photocathodes. J. Phys. Chem. Lett. 5, 3222–3226 (2014).

22. Leung, J. J. et al. Photoelectrocatalytic H2 evolution in water with molecular catalysts immobilised on p-Si via a stabilising mesoporous TiO2 interlayer. Chem. Sci. 8, 5172–5180 (2017).

23. Chandrasekaran, S. et al. A robust ALD-protected silicon-based hybrid photoelectrode for hydrogen evolution under aqueous conditions. Chem. Sci. 10, 4665–4675 (2019).

24. Gu, J. et al. Water reduction by a p-GaInP2 photoelectrode stabilized by an amorphous TiO2 coating and a molecular cobalt catalyst. Nat. Mater. 15, 456–460 (2016).

25. Kumar, B., Beyler, M., Kubiak, C. P. & Ott, S. Photoelectrochemical hydrogen generation by an [FeFe] hydrogenase active site mimic at a p-Type silicon/molecular catalyst incorporated into a metal organic framework coating enhances the performance of Cu2O in photocatalytic CO2 reduction. J. Am. Chem. Soc. 141, 10924–10929 (2019).

26. Downes, C. A. & Marques, C. S. Efficient electrochemical and photoelectrochemical H2 production from water by a cobalt diholine one-dimensional metal–organic surface. J. Am. Chem. Soc. 137, jacs.s077020 (2015).

27. Deng, X. et al. Metal–organic framework coating enhances the performance of Cu2O in photocatalytic CO2 reduction. J. Am. Chem. Soc. 141, 10924–10929 (2019).

28. Nellist, M. R., Laskowski, F. A. L., Lin, F., Mills, T. J. & Boetcker, S. W. Semiconductor–electrocatalyst interfaces: theory, experiment, and applications in photoelectrochemical water splitting. Acc. Chem. Res. 49, 740 (2016).

29. Barroso, M. et al. Dynamics of photogenerated holes in surface-modified Fe2O3 photoanodes for solar water splitting. Proc. Natl Acad. Sci. USA 109, 15640–15645 (2012).

30. Li, W. et al. Comparison of heterogenized molecular and heterogeneous oxide catalysts for photoelectrochemical water oxidation. Energy Environ. Sci. 9, 1794–1802 (2016).

31. Sato, K., Kim, H. J., Pekarek, R. T. & Rose, M. J. Hybrid organic/inorganic band-edge modulation of p–Si(111) photoelectrodes: effects of R, metal oxide, and Pt on H2 generation. J. Am. Chem. Soc. 137, 3173–3176 (2015).

32. Duffey, N. W., Dobson, K. D., Gordon, K. C., Robinson, B. H. & McQuillan, A. J. In situ infrared spectroscopic analysis of the adsorption of ruthenium(II) bipyridyl dicarboxylic acid photosensitizers to TiO2 in aqueous solutions. Chem. Phys. Lett. 266, 451–455 (1997).

33. Johnson, B. A., Bhunia, A. & Ott, S. Electrocatalytic water oxidation by a molecular catalyst incorporated into a metal–organic framework thin film. Dalt. Trans. 46, 1382–1388 (2017).

34. Ahrenholz, S. R., Eley, C. C. & Morris, A. J. Solvothermal preparation of an electrocatalytic metalloporphyrin MOF thin film and its redox hopping charge-transfer mechanism. J. Am. Chem. Soc. 136, 2464–2472 (2014).

35. Andrieux, C. P., Costentin, C., Di Giovanni, C., Savéant, J.-M. & Tard, C. Conductive mesoporous catalytic films. current distortion and performance degradation by dual-phase ohmic drop effects. Analysis and remedies. J. Phys. Chem. C 120, 21263–21271 (2016).

36. Kim, Y. S., Balland, V., Limoges, B. & Costentin, C. Cyclic voltammetry modeling of proton transport effects on redox charge storage in conductive materials: application to a TiO2 mesoporous film. Phys. Chem. Chem. Phys. 19, 17944–17951 (2017).

37. Andrieux, C. P. & Savéant, J. M. Electroneutrality coupling of electron hopping between localized sites with electroinactive counterion displacement. 1. Potential-step plateau currents. J. Phys. Chem. 92, 6761–6767 (1988).

38. Lin, F. & Boetcker, S. W. Adaptive semiconductor/electrocatalyst junctions in water-splitting photoanodes. Nat. Mater. 13, 81–86 (2014).

39. McCarthy, B. D. et al. Analysis of electrocatalytic metal-organic frameworks. Coord. Chem. Rev. 406, 213137 (2020).

40. Andrieux, C. P. & Savéant, J. M. Electron transfer through redox polymer films. J. Electroanal. Chem. Interfacial Electrochem. 111, 377–381 (1980).

41. Aoki, T., Tokuda, K. & Matsuda, H. Theory of linear sweep voltammery with finite diffusion space. J. Electroanal. Chem. Interfacial Electrochem. 146, 417–424 (1983).

42. Johnson, B. A., Beiler, A. M., McCarthy, B. D. & Ott, S. Transport phenomena: challenges and opportunities for molecular catalysis in metal-organic frameworks. J. Am. Chem. Soc. 142, 11941–11956 (2020).

43. Peter, L. M. & Upul Wijayantha, K. G. Photoelectrochemical water splitting at semiconductor electrodes: fundamental problems and new perspectives. ChemPhysChem 15, 1983–1995 (2014).

44. Kaiser, B. et al. Solar hydrogen generation with wide-band-gap semiconductors: GaP(100) photoelectrodes and surface modification. Chem. Phys. Chem. 13, 3053–3060 (2012).

45. Wadsworth, B. L., Beiler, A. M., Khusnutdinova, D., Reyes Cruz, E. A. & Moore, G. F. Interplay between light flux, quantum efficiency, and turnover frequency in molecular-modified photoelectrosynthetic assemblies. J. Am. Chem. Soc. 141, 15932–15941 (2019).

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