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Gold nanoparticle-based supramolecular approach for dye-sensitized H₂-evolving photocathodes†

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Solar conversion of water into the storable energy carrier H₂ can be achieved through photoelectrochemical water splitting using light adsorbing anodes and cathodes bearing O₂ and H₂ evolving catalysts, respectively. Herein a novel photocathode nanohybrid system is reported. This photocathode consists of a dye-sensitized p-type nickel oxide (NiO) with a perylene-based chromophore (**PCA**) and a tetra-adamantane modified cobaloxime reduction catalyst (**Co**) that photo-reduces aqueous protons to H₂. An original supramolecular approach was employed, using β-cyclodextrin functionalized gold nanoparticles (**β-CD-AuNPs**) to link the alkane chain of the PCA dye to the adamantane moieties of the cobaloxime catalyst (**Co**). This new architecture was investigated by photoelectrochemical measurements and via femtosecond-transient absorption spectroscopy. The results show that irradiation of the complete NiO|**PCA|β-CD-AuNPs|Co** electrode leads to ultrafast hole injection into NiO ($\pi = 3$ ps) from the excited dye, followed by rapid reduction of the catalyst, and finally H₂ evolution.

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Introduction

The development of sustainable energy technologies that allows us to decrease our use of fossil fuels, with their negative environmental impact, is one of the greatest challenges of our times. Inspired by natural photosynthesis, the conversion of solar energy into fuels (e.g. H₂, CH₄, CO) from abundant substrates (H₂O, CO₂) by photoelectrochemical cells (PECs) is a promising strategy to address this challenge.^{1–4} The electrodes in dye-sensitized photoelectrochemical cells (DS-PECs) are typically based on a wide band gap semiconductor that is sensitized with molecular dye molecules as light harvesting compounds^{5–8} and catalysts for solar fuel production.^{9,10} In a complete tandem cell, water oxidation occurs at the photoanode, while H₂ is produced at the photocathode.

Significant work has been done to improve the photoanodes, which can now achieve relatively high photocurrents and efficiencies.^{5,7,8} In comparison, the photocathodes have been less explored, and their performances are currently limiting the efficiency of complete DS-PECs.^{6,11–13} In dye-sensitized photocathodes (DS-PCs), light absorption by the surface bound dye is followed by hole injection into the valence band of the semiconductor, creating a reduced dye that can transfer an electron to the co-immobilized catalyst. Two reducing equivalents as well as two protons are needed by the catalyst to form H₂. Rapid charge recombination relative to the rate of proton reduction limits the overall efficiency of DS-PCs. To generate higher photocurrents, the dominant recombination mechanisms between holes in the p-type semiconductor and (1) the reduced dye or (2) the reduced catalyst must be prevented.¹²

The nature of the molecular components, the semiconductor itself, and the way by which photosensitizer and catalyst are arranged on the electrode surface greatly impact the performance of DS-PCs. Several DS-PCs photocathodes have been described with the majority relying on NiO substrates.^{6,14–28} However, problems associated with the high density of traps and low hole mobility have been identified,^{29–32} and other p-type metal oxide semiconductors such as LaFeO₃ and CuCrO₂ have been recently proposed as alternative materials for the construction of DS-PCs for solar-

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driven H_2 generation.^{33–35} As far as the dye is concerned, organic chromophores and ruthenium based complexes are the two families of light-harvesting units that are frequently used. The best photosensitizers have a donor–bridge–acceptor structure, where the HOMO is positioned close to the anchoring group and the LUMO is positioned away from the NiO surface.^{13,36–38} This push–pull structure is thought to promote charge-separation and slow down charge recombination by keeping the electron on the reduced dye physically distant to the holes in the NiO . Among organic chromophores, perylene-based dyes are renowned for their high stability and intense visible light absorption. PMI based photocathodes have been assembled with heterogeneous catalysts,³⁷ molecular catalysts in solution³⁹ and immobilized $\text{Ni}(\text{P}_2\text{N}_2)_2$ – based catalysts,^{26,35} but to our knowledge, never with a cobaloxime type catalysts.

To improve PC performance, many PC designs have focused on optimizing the electrode assembly. The different strategies are DS-PCs combined with a catalyst in solution,^{16,36,39} co-grafted dye/catalyst systems,^{6,19,22,40} layer by layer assemblies of the dye and the catalyst,^{17,18,20} and supramolecular or covalent dye/catalyst assemblies.^{14,15,21,23}

In this study, we report the design and synthesis of a new dye/catalyst supramolecular assembly based on β -cyclodextrin functionalized gold nanoparticles (β -CD-AuNPs) that connect a perylene-based dye (PCA) to a cobaloxime catalyst (Co). A rational and simple procedure of sequential immersion of NiO films into solutions of PCA, β -CD-AuNPs and Co was employed to produce electrodes of the type $\text{NiO}|\text{PCA}|\beta\text{-CD-AuNPs}|\text{Co}$ (Fig. 1A) where the catalyst is spatially separated from the semiconductor electrode to reduce inefficiency from charge recombination. β -CD binds strongly to the alkane chain of the dye and the adamantane moiety of the catalyst to form the supramolecular assembly. Cyclodextrin has been previously used to form host–guest interactions in order to immobilized molecular catalysts on electrode surfaces.^{41,42} In the present architecture, AuNPs are proposed to act as electron transfer bridges by accepting electrons from PCA and transporting them to Co. This function of the AuNPs is motivated by pre-

vious studies in which such constructs were shown to enhance electron transfer rates at protein/electrode interfaces in bioelectrochemical systems.^{43,44} Other studies have shown that small gold nanoparticles (2–3 nm) significantly increase charge separation in chromophore-gold nano-assemblies, where the charge separation is sustained for several microseconds before undergoing recombination.^{45,46} However, to the best of our knowledge, this strategy has never been explored to develop operative photocathodes for H_2 evolution.

Results and discussion

Synthesis and characterization

PCA,⁴⁷ β -CD-AuNPs^{48,49} and $[\text{Co}(\text{dcpG}\text{H})(\text{dcpG}\text{H}_2)]\text{Cl}_2$ ⁵⁰ **1** were synthesized as previously reported. The adamantane moiety was added to **1** via peptide coupling (see ESI†) to afford **Co**. The target complex was characterized by ^1H NMR, Fourier transform infrared (FT-IR) spectroscopy, high-resolution mass spectrometry (HR-MS) and electrochemistry (Fig. S1–S3†). Fig. S11† shows a transmission electron microscopy (TEM) image of the functionalized AuNPs and reveals an average AuNP size of 3.2 ± 0.7 nm, which is in good agreement with previous studies.⁴⁹ This type of AuNPs was selected to form a supramolecular assembly due to the cyclodextrin moieties which are able to interact with a variety of guests.⁵¹ The PCA dye was selected for its ability to interact with both NiO and β -CD. The monoanhydride precursor of PCA opens to form the dicarboxylate when exposed to metal oxide semiconductors for convenient electrode functionalization,⁵² while the hydrophobic alkane chain can interact with the β -CD moiety of the AuNPs, as illustrated in Fig. 1A. For short alkane chain, a host : guest ratio of 1:1 is expected.⁵³ The optical and electrochemical properties of PCA are display in Table 1. The dye anion exhibits sufficient driving force ($E_{\text{PCA}/\text{PCA}^-} = -0.91$ V vs. NHE) to reduce **Co** to a catalytically active state ($E_{\text{Co,inset}} = -0.38$ V vs. NHE).⁵⁴ Hole injection into the valence band of NiO ($E \approx 0.46$ V vs. NHE) is facilitated by the highly anodic

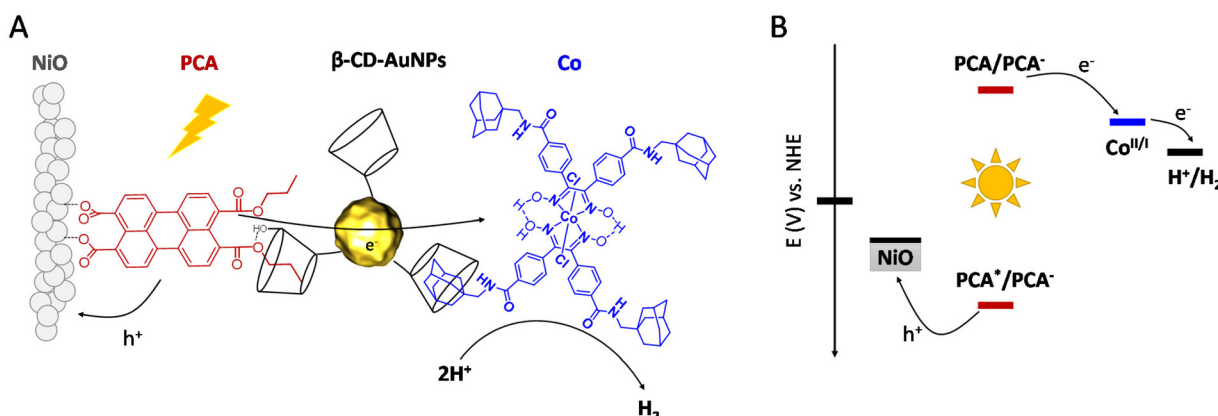


Fig. 1 (A) Representation of the $\text{PCA}|\beta\text{-CD-AuNPs}|\text{Co}$ supramolecular assembly anchored onto a mesoporous NiO thin film, (B) energy diagram of the photocathode.



Table 1 Optical and electrochemical properties

	λ_{max}^a	ϵ^a ($\text{M}^{-1} \text{cm}^{-1}$)	E_{red} (V vs. Fc^+/Fc)	E_{ox} (V vs. Fc^+/Fc)	E_{red} (V vs. NHE) ^e	E_{ox} (V vs. NHE) ^e
PCA ^{c,d}	503	36 000	-1.64	1	-0.91	1.73
	476	29 300				
Co ^b	—	—	-1.13	-0.4	-0.4	0.33

^a Adsorption spectra in CH_2Cl_2 . ^b Redox potentials obtained in a DMF solution. ^c Redox potentials obtained in a CH_3CN solution. ^d Redox potentials obtained for surface bound molecules. ^e Redox potentials vs. NHE were calculated from the values vs. $\text{Fc}^{+/-}$ by addition of 730 mV.⁵⁵

excited state reduction potentials ($E_{\text{PCA}^+/\text{PCA}^-} = +1.73$ V vs. NHE). PCA displays a broad absorption from 450 to 550 nm with a maximum at 503 nm and a shoulder at 476 nm (Fig. S6†). The intense visible light absorption and favourable electrochemical properties of PCA are well suited for the application of the dye in DS-PCs with Co as the catalyst (Fig. 1B). To confirm this ability to reduce the immobilized catalyst, PCs were assembled for photoelectrochemical analysis.

Electrode preparation and pre-catalysis characterization

PCA, β -CD-AuNPs, and Co were used to prepare a photocathode for H_2 evolution based on FTO coated NiO films.^{26,56} NiO electrodes were prepared by immersion into a solution of PCA (0.2 mM) in a mixture of CH_2Cl_2 and methanol (1:1) for 24 hours to give NiO|PCA.

Co was immobilized on the NiO|PCA photocathodes in a layered assembly using β -CD-AuNPs to link the alkane chain of the dye to the adamantane moiety of the catalyst. Sequential loading of molecules was carried out by soaking the NiO|PCA films in an aqueous solution of β -CD-AuNPs (0.5 mg mL^{-1}) for 1 h to give NiO|PCA| β -CD-AuNPs, followed by soaking in a Co

solution (0.2 mM in MeOH-1%DMF) for 30 minutes to afford NiO|PCA| β -CD-AuNPs|Co electrodes.

Control electrodes without β -CD-AuNPs (denoted as NiO|PCA + Co) were obtained by soaking the NiO|PCA films in a Co solution following the same protocol. All immobilization steps were carried out at room temperature in the dark (see experimental part for details). The absorption spectra of the sensitized electrodes are depicted in Fig. 2B. UV-vis spectra of the films show the characteristic visible light absorption profiles of the dye. NiO|PCA electrodes display a blue shift and higher intensity for the peak at 448 nm than at 503 nm as opposed to the solution-based experiments (Fig. S6 and S9†). This is attributed to the anhydride opening and the binding of the carboxylates to NiO.^{57,58} β -CD-AuNPs binding was confirmed by UV-vis spectroscopy. In aqueous solution (Fig. 2A) a relatively weak plasmon resonance absorption maxima at 507 nm is observed.⁴⁹ The difference absorption spectrum of NiO|PCA and NiO|PCA| β -CD-AuNPs electrodes (Fig. 2C) exhibits a peak at 538 nm attributed to the surface plasmon resonance of immobilized β -CD-AuNPs.

The immobilization of the β -CD-AuNPs and the catalyst does not affect the NiO|PCA interface. The amount of dye loading on the photoelectrode was quantified by measuring the UV-vis spectra of PCA solution after sensitization of NiO films. A surface coverage of $20 \pm 2 \text{ nmol cm}^{-2}$ was found, which is in the same range as those determined in previous studies using similar electrodes.⁵⁹ The amount of catalyst loading on the photoelectrodes was quantified by ICP-OES after acid digestion of the electrodes. A coverage of $0.5 \pm 0.3 \text{ nmol cm}^{-2}$ was found for NiO|PCA| β -CD-AuNPs|Co electrodes. No catalyst was observed for the NiO|PCA + Co electrodes confirming the crucial role of β -CD-AuNPs for the immobilization of Co catalysts.

The presence of PCA, β -CD-AuNPs and Co at the surface of NiO electrodes was further investigated by XPS analyses. The

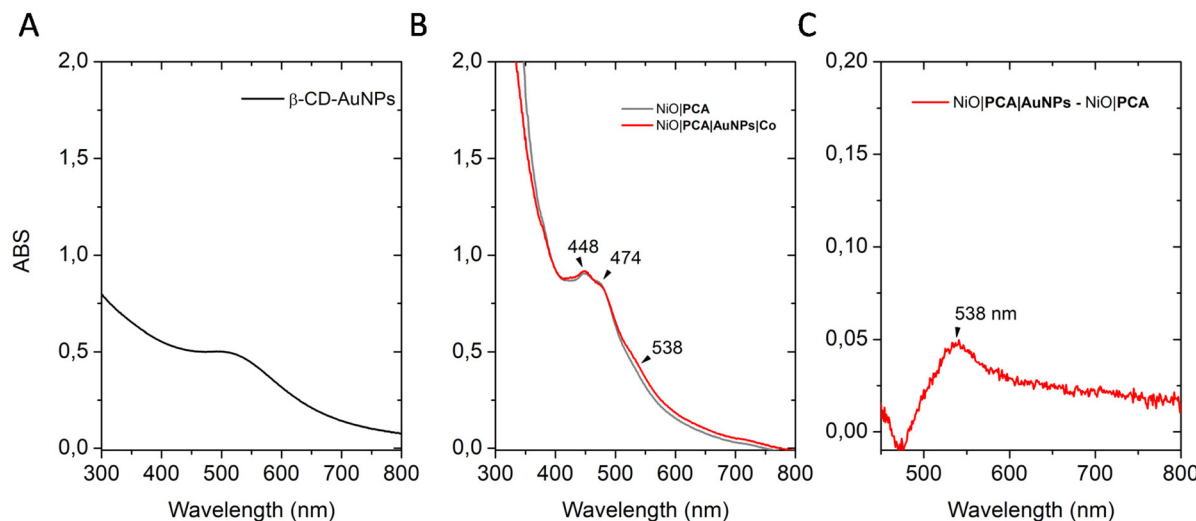


Fig. 2 Absorption UV-vis spectrum of (A) an aqueous solution of β -CD-AuNPs, (B) NiO|PCA (grey) and NiO|PCA| β -CD-AuNPs|Co (red) electrodes (NiO electrodes were used to set the background), and (C) Difference in absorption of NiO|PCA| β -CD-AuNPs and NiO|PCA electrodes.

survey spectrum of the NiO electrodes (Fig. S12†) shows peaks attributed to Ni 2p and NiLMM Auger photoelectrons. After chemisorption of PCA onto NiO, the survey spectrum of the resulting material NiO|PCA shows an increase in the level of C and O compared to the NiO electrodes, consistent with the presence of PCA (Fig. S13†). The survey spectrum of NiO|PCA| β -CD-AuNPs|Co electrodes (Fig. S14†) shows the same features, but also additional information such as the presence of N and Au. A signal in the 398–402 eV range of the N 1s core level region (Fig. S13†) corresponds to N atoms present in grafted Co. The doublets for Au 4f_{7/2} (83.8 eV) and Au 4f_{5/2} (87.5 eV) shown in Fig. S14† are characteristic of Au⁰ of the β -CD-AuNPs.^{60,61} The absence of a signal at 84.9 eV as found for Au¹ in a gold octanethiol complex⁶² indicates that the gold atoms in the clusters are largely present as Au⁰. Further substantiation can be found on Au 4f peaks with slight asymmetry consistent with the presence of small nanoparticles/cluster.⁶³ Au⁰ peaks due to metallic conduction (*i.e.* non-occupied states at the Fermi level) should be asymmetric with a tailing towards higher binding energy. XPS data clearly evidence the immobilization of PCA, β -CD-AuNPs and Co at the surface of NiO electrodes.

Photoelectrochemistry with NiO|PCA and NiO|PCA| β -CD-AuNPs

To ensure that NiO|PCA| β -CD-AuNPs electrodes could function without the kinetic limitations of sluggish catalyst reactions and charge recombination, photoelectrochemical measurements were performed in the presence of a soluble irreversible electron acceptor (IEA) in solution (Fig. 3A). Linear sweep voltammetry (LSV) was carried out in MES buffer electrolyte (0.1 M, pH 5), at room temperature in an Ar-purged one compartment three electrodes electrochemical cell using a Pt counter electrode and a Ag/AgCl reference electrode. Irradiation of NiO|PCA and NiO|PCA| β -CD-AuNPs electrodes resulted in only minor photocurrents without a soluble acceptor (Fig. 3A). Addition of the IEA chloropentamminecobalt(III) chloride (10 mM) in the electrolyte solution allows for estimation of a maximal attainable photocurrent as [Co^{III}(NH₃)₅Cl₂]Cl is known to be irreversibly reduced in solution.⁶⁴ The IEA allows the photo-reduced dye to dispose its photoelectrons and to regenerate the ground state, thereby limiting the effects of reductive dye decomposition and charge recombination, thereby enhancing the photocurrent response for NiO|PCA and NiO|PCA| β -CD-AuNPs electrodes dramatically (Fig. 3A).

With NiO|PCA| β -CD-AuNPs electrodes, an absolute photocurrent response of $|I| \approx 420 \mu\text{A cm}^{-2}$ (0.0 V *vs.* Ag/AgCl) was observed, which indicates efficient light induced hole injection from the dye to the valence band of NiO with reduction of the IEA by PCA⁻. For comparison, NiO|PCA electrodes displayed a lower maximum photocurrent ($|I| \approx 344 \mu\text{A cm}^{-2}$, 0.0 V *vs.* Ag/AgCl), suggesting lower susceptibility to recombination between PCA⁻ and holes in NiO when AuNPs are present. This finding supports that co-anchoring a catalyst could be a viable

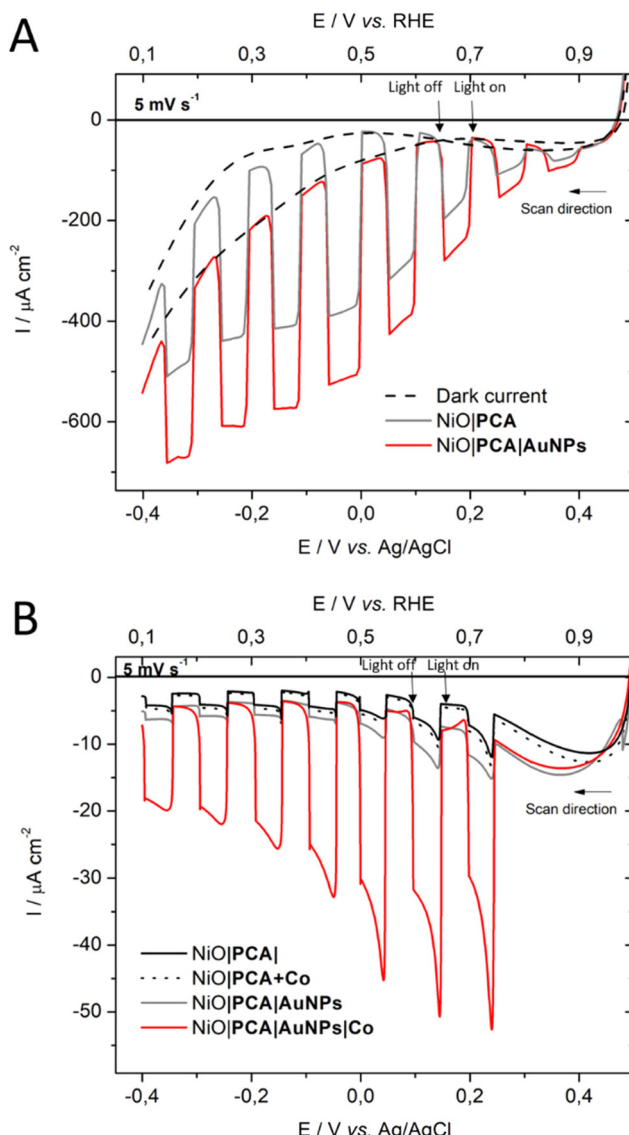


Fig. 3 Linear sweep voltammograms (LSVs) under chopped light illumination of (A) NiO|PCA (grey) electrodes, and NiO|PCA| β -CD-AuNPs (red) electrodes with 10 mM of [Co^{III}(NH₃)₅Cl₂]Cl acceptor in solution. (B) LSV scans of NiO|PCA, NiO|PCA| β -CD-AuNPs, NiO|PCA + Co and NiO|PCA| β -CD-AuNPs|Co. All experiments were performed in MES buffer (0.1 M, pH 5). Illumination with a white LED with 100 mW cm⁻². An active electrode area of 1 cm⁻² was used with a scan rate of 5 mV s⁻¹.

approach to exploit the reductive power of NiO|PCA| β -CD-AuNPs electrodes for H₂ photo-production.

Photoelectrochemistry with NiO|PCA| β -CD-AuNPs|Co and H₂ generation

The photoelectrochemical properties of the NiO|PCA| β -CD-AuNPs|Co photocathodes were evaluated in aqueous solution (0.1 M MES buffer, pH 5). Linear sweep voltammograms were recorded under chopped light irradiation (Fig. 3B). Immobilization of Co catalyst on the electrode resulted in a 10-fold photocurrent increase compared to that of NiO|

PCA| β -CD-AuNPs electrodes ($|j| = 15.8 \mu\text{A cm}^{-2}$, $-0.2 \text{ V vs. Ag/AgCl}$). This enhanced response is attributed to the ability of PCA^- to reduce **Co** and ultimately protons. The absence of current enhancement in the control experiment confirmed the crucial role of β -CD-AuNPs for the immobilization of **Co** catalysts.

Chronoamperometric measurements were performed under chopped light irradiation at $-0.1 \text{ V vs. Ag/AgCl}$ over a period of 15 min showing steady-state cathodic photocurrent (Fig. 4) with a current density of *ca.* $-11 \mu\text{A cm}^{-2}$. Photoelectrochemical H_2 generation was confirmed by long term electrolysis over a period of 1 h (Fig. S16 \dagger), during which the magnitude of the current decreases slowly.

Clark electrode measurements confirmed the evolution of hydrogen in solution during the chronoamperometric measurements with a faradaic efficiency of $10 \pm 2\%$ and a TON of 8. Noteworthy is that these measurements do not take into account H_2 present in the headspace.⁶⁵ Moreover, no H_2 generation was observed in the absence of catalyst or in the absence of light. This relatively low yield is within the current state-of-the-art in NiO-based dye-sensitized H_2 -evolving photocathodes.³⁴ Attempts to increase the yield by increasing the catalyst loading on the electrode resulted in smaller photocurrents. The decrease in the photocurrent (from 9.8 to $4 \mu\text{A cm}^{-2}$) observed during long-term electrolysis can be attributed partly to some leaching or demetalation⁶⁶ of the catalyst since we observed a decrease of catalyst loading by ICP measurements after photoelectrochemical measurements (Table S1 \dagger). Dye aggregation and competing reactions, such as NiO reduction^{15,67,68} may also be considered.

Transient absorption spectroscopy

Femtosecond transient UV/Vis absorption measurements were conducted on **NiO|PCA**, **NiO|PCA| β -CD-AuNPs**, and **NiO|PCA| β -CD-AuNPs|Co** photocathodes to understand the charge dynamics upon PCA excitation. Energetically, hole injection is possible from excited PCA dye to the NiO valence band.⁶⁹ The excited state dynamics of PCA in **NiO|PCA** was compared to that of a **ZrO₂|PCA** reference system, in which neither electrons

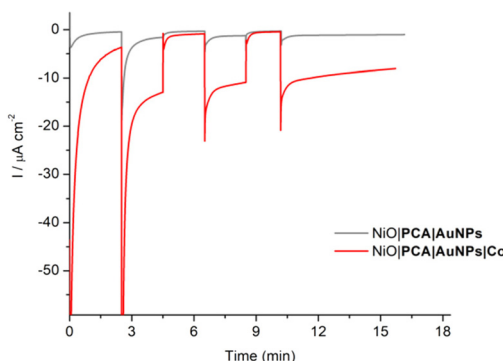


Fig. 4 Chopped-light chronoamperometric measurements of **NiO|PCA| β -CD-AuNPs** and **NiO|PCA| β -CD-AuNPs|Co** photocathodes at an applied bias of $-0.1 \text{ V vs. Ag/AgCl}$ in pH 5 MES buffer.

nor holes can be injected due to the high-energy conduction band and low-energy valence band of **ZrO₂**.⁷⁰ Fig. 5 shows the transient absorption spectra of the PCA dye attached to **ZrO₂** and **NiO** at different time delays after excitation. Both sets of spectra display the ground state bleach (GSB) of the **PCA** dye at around 420 to 550 nm. The spectra of the **ZrO₂|PCA** system shows an excited state absorption (ESA) from 550 to 750 nm with a distinct maximum around 580 nm, as expected.⁵² In **NiO|PCA**, the induced absorption is seen further to the red, without a distinct maximum. This difference agrees with the difference spectrum after spectroelectrochemical reduction of the NiO-bound dye (Fig. S10 \dagger). Fig. S18 \dagger presents the kinetic traces associated with ESA (probed at 580 nm) in **ZrO₂|PCA**, and the reduced dye in **NiO|PCA** (probed at 625 nm), both of which were fitted with a with a double-exponential function.

Clearly, attaching PCA to **NiO** ($\tau_1 = 15 \text{ ps (60\%)}$, $\tau_2 = 300 \text{ ps (40\%)}$) results in faster dynamics than having the dye on **ZrO₂** ($\tau_1 = 30 \text{ ps (60\%)}$, $\tau_2 = 500 \text{ ps (40\%)}$).³⁹ This is attributed to the photo driven hole injection from excited PCA into the **NiO** valence band.³⁹ The thereby produced charge separated state **NiO⁺|PCA⁻** is however very short lived, and recombines to the ground state on the picosecond timescale, giving rise to the observed overall faster kinetics in **NiO|PCA** compared to that

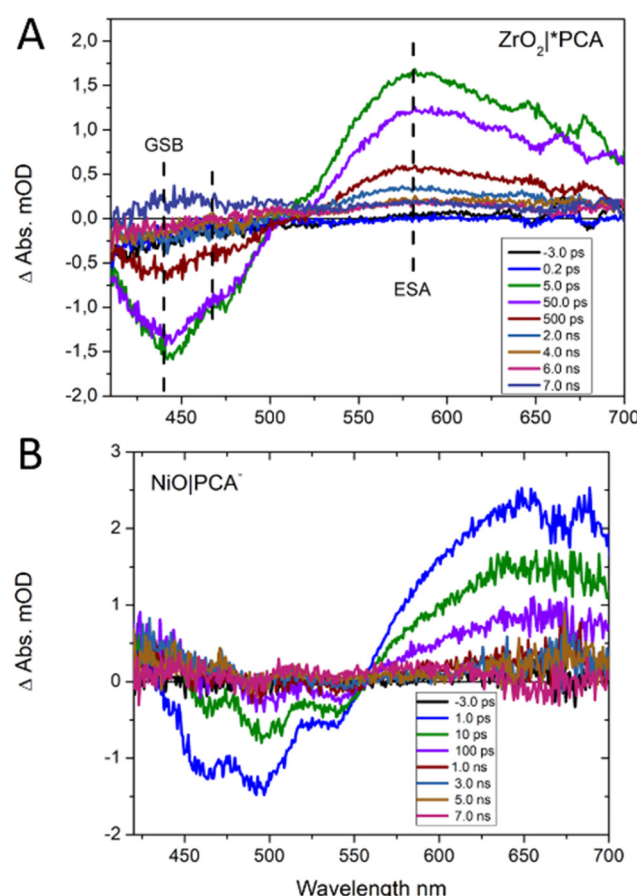


Fig. 5 fs TA spectra of (A) **ZrO₂|PCA** and (B) **NiO|PCA**, after excitation at 440 nm.



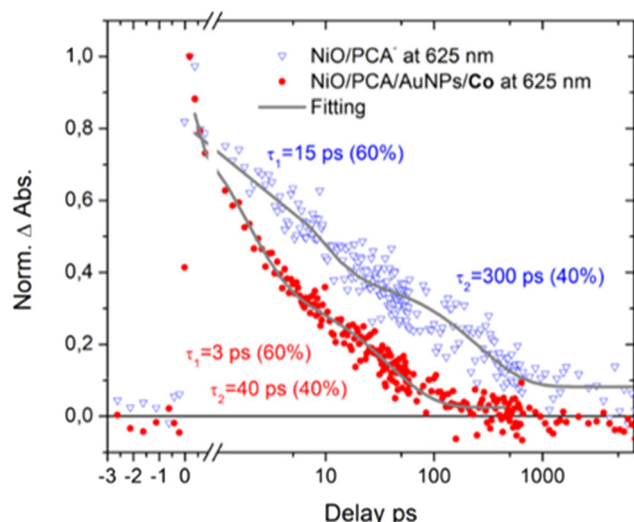


Fig. 6 Time traces of $\text{NiO}|\text{PCA}^-$ (blue triangles) and $\text{NiO}|\text{PCA}|\text{AuNPs}|\text{Co}$ (red dots) at 625 nm (reduced dye).

in $\text{ZrO}_2|\text{PCA}$. The photoelectrochemical properties the $\text{NiO}|\text{PCA}|\beta\text{-CD-AuNPs}|\text{Co}$ photocathodes (Fig. 3B) shows the role of the Au NPs as an important structural component in the supramolecular system. Fig. S17C† presents the TA spectra of the $\text{NiO}|\text{PCA}|\beta\text{-CD-AuNPs}|\text{Co}$ photocathode which qualitatively contains the same features as those of $\text{NiO}|\text{PCA}$, but clearly with even shorter lifetimes of the signal that is attributed to the reduced PCA. Fig. 6 presents the kinetic traces of the decay of the reduced dye in $\text{NiO}|\text{PCA}|\beta\text{-CD-AuNPs}|\text{Co}$ photocathodes, with a fit ($\tau_1 = 3$ ps (60%), $\tau_2 = 40$ ps (40%)), in comparison with the same features in $\text{NiO}|\text{PCA}$. In addition to the fast dynamic that is caused by attaching the dye to NiO (due to the hole injection and recombination), directed electron transfer from the reduced PCA *via* $\beta\text{-CD-AuNPs}$ to the catalyst results in even faster dynamics of PCA^- (Fig. 6). On one hand, inserting the $\beta\text{-CD-AuNPs}$ will improve the loading of the catalyst and at the same time, it will guide the electron transfer from the reduced dye to the catalyst. It's important to note that the transient absorption data were acquired without applied bias, the presence of which can alter charge recombination rates.^{28,32,71} The main focus of the transient absorption study was however to demonstrate hole injection and find support for electron transfer to the AuNPs. Spectroscopically, it was not possible to detect the reduced form of the **Co** catalyst due to its comparably weak spectroscopic features compare to the dye GSB.⁷²

Conclusions

Here, we report the first supramolecular assembly based on cyclodextrin functionalized gold nanoparticles for the construction of a NiO -based photocathode for H_2 evolution from water. $\beta\text{-CD-AuNPs}$ play a crucial role in the immobilization of **Co** catalysts and guide the electron transfer from the reduced dye to the catalyst. It was shown that light irradiation initiates

an electron transfer from the NiO to the dye molecule, and subsequently from the reduced dye to the catalyst. The photocathode produces H_2 with a faradaic efficiency of 10%, which is in the same range as those of previously reported molecular photocathodes based on cobaloxime catalysts.³⁴ Further work to improve this system will target the use of novel p-type transparent materials such as CuCrO_2 , which have already proved promising.^{34,35}

Experimental

Materials

All chemicals were purchased from commercial suppliers (Sigma-Aldrich, VWR and Alfa-Aesar) and used without further purification, unless stated otherwise. Solvents were dried using drying agents and freshly distilled under argon before use. All organic reactions were performed under inert argon atmosphere.

Instrumentation

^1H NMR spectra were measured on a JEOL Eclipse 400 MHz spectrometer. High-resolution mass spectral analyses (HRMS) were performed on a pneumatically-assisted electrospray ionization Fourier transform (FTMS + pNSI) mass spectrometer (orbitrapXL). UV-vis spectroscopic measurements were collected using a Varian Cary 50 Bio UV-vis spectrophotometer. Attenuated total reflectance Fourier transform infrared (ATR-FTIR) spectroscopic measurements were performed using a PerkinElmer Spectrum One FTIR-Spectrometer with a PerkinElmer Universal ATR Sampling Accessory. Inductively Coupled Plasma-Optical Emission Spectrometry (ICP-OES) experiments were performed using a PerkinElmer Optical Emission Spectrometer. X-ray photoelectron spectroscopy (XPS) spectra were recorded on a PHI Quantera II scanning XPS microprobe with monochromated $\text{Al K}\alpha$ radiation (1486.6 eV). Survey spectra were captured at a pass energy of 224 eV, while for high-resolution spectra a pass energy of 55 eV was used. All spectra were referenced against an internal signature ($\text{C } 1s$, 284.8 eV). Transmission electron microscopy (TEM) images were obtained using JEOL JEM 1400 instrument. The particle size distribution was obtained from digitalized amplified micrographs by averaging the larger and smaller axis diameters measured in each particle.

Electrochemical and photoelectrochemical measurements were acquired with an Autolab potentiostat (PGSTAT 302N). Electrochemical measurements were conducted in a traditional three-electrodes system with Ag/AgNO_3 reference electrode, a platinum rod as counter electrode and either glassy carbon or FTO/ NiO as working electrodes. Ferrocene was used as an internal standard and all the potentials converted into the Normal Hydrogen Electrode using $E_{\text{Fe}^{+}/\text{Fe}} = 0.64$ V *vs.* NHE. LED lamp (cool white, 5000K) was used as light source. A Unisense Clark electrode H_2 sensor was used for the detection and the quantification of H_2 . A Dektak150 Surface Profiler was used to measure the NiO film thicknesses.

TA spectroscopy

Femto-second transient absorption spectroscopy (fs-TAS) was performed probing in the UV-vis region using a Newport TAS. A Coherent Libra Ti:sapphire amplifier (1.5 mJ, 3 kHz, 800 nm, fwhm 40 fs) was used and split into pump and probe beams. An excitation wavelength of 440 nm was generated by directing the pump beam into the optical parametric amplifiers (TOPAS-Prime and NIRUVVIS, Light Conversion) and then focused and centered on the dry films with a pump power adjusted to 500 μ W. The 800 nm fundamental of the amplifier was focused on a CaF₂ crystal (Newport TAS), generating the white light supercontinuum probe. A silicon diode array (Newport custom made) was used to record the probe spectrum. A mechanical chopper blocked every other pump pulse, and the transient absorption at each time point was calculated for an average of 1000 ms chopped/un-chopped pulse pairs. To record the transient absorption spectra at different time points, an optical delay line was used to scan the delay of the probe beam relative to the pump beam from -5 ps to 7.5 ns. A total of 10–15 scans were collected and averaged for each sample. Prior to analysis, the data was corrected for the spectral chirp using Surface Xplorer 4.2, where single wavelength fits were also performed.

Synthesis

[Co(dcpgH)(dcpgH₂)]Cl₂⁵⁴ (dcpgH₂ = diphenylglyoxime-4,4'-dicarboxylic acid) **1**, PCA dye⁴⁷ and β -CD-AuNPs^{48,49} were synthesized according to literature method.

Synthesis of **Co** (Benzotriazol-1-yl)tripyrrrolidinophosphonium hexafluorophosphate (PyBOP, 4 equiv., 0.51 mmol) was added to a solution of **1** (1 equiv., 0.13 mmol), diisopropylethylamine (7 equiv., 0.90 mmol) and 1-adamantanemethylamine (4 equiv., 0.51 mmol) dissolved in DMF (7 mL). Upon addition, 1-adamantanemethylamine became completely soluble. The reaction mixture was then stirred at R.T. overnight. DMF was removed under reduce pressure and the resulting oil was dissolved in 10 mL of CH₂Cl₂. The organic phase was washed with brine (2 \times 5 mL), water (2 \times 5 mL), dried (MgSO₄), filtered and reduced to 2 mL. Then the crude orange product was precipitated with Et₂O, filtered and washed with Et₂O. The solid was purified by flash chromatography (0 to 7% methanol in CH₂Cl₂). The pure product was obtained after evaporation of the solvent in combine fractions to give 2 as an orange solid (83 mg, 46%). ¹H NMR (CD₂Cl₂, 400 MHz): δ (ppm) 19.25 (Br, 2H, oxime), 7.51–7.26 (m, 16H, Ar), 7.10 (s, 4H, NH amide), 3.08 (s, 8H, CH₂), 1.94 (s, 12H, CH adamantan), 1.68–1.52 (m, 48H, CH₂ adamantan). HR-MS (ESI⁺): m/z , 1302.6 [M-H-2Cl]⁺.

Electrode preparation

First, FTO-coated glass slides were cleaned *via* sequential sonication steps using detergent, acetone, and isopropanol. Then, NiO was screen printed onto the clean FTO coated glass using a NiO paste as previously reported.^{56,73} After each layer of screen printing, the electrodes were annealed at 120 °C for

9 minutes on a hot plate. Three layers were screen printed and were finally annealed at 450 °C for 30 minutes in an oven, resulting in a thickness of ~1.2 μ m determined by profilometry.

Photocathode assemblies

NiO electrodes were soaked in a 1:1 CH₂Cl₂ : MeOH solution of PCA (0.2 mM) for 24 h at room temperature, then rinsed with CH₂Cl₂; MeOH and dried in air affording NiO|PCA electrodes. For the molecular assemblies PCA| β -CD-AuNP|Co, sequential loading of molecules was carried out by soaking the films in aqueous solution of β -CD-AuNPs (0.5 mg mL⁻¹) for 1 h (giving NiO|PCA| β -CD-AuNPs electrodes), followed by soaking in methanol-1%DMF solution of the catalyst Co (0.2 mM) for 0.5 h. The electrodes were then rinsed with MeOH-1%DMF solution and dried in air. Water and methanol were used to rinse the films between the two loading solutions. NiO|PCA|Co electrodes were obtained following the same procedure without the soaking step in solution of β -CD-AuNPs.

Surface loading quantification

Surface coverage (Γ in mol cm⁻²) of PCA was evaluated by measuring the UV-vis spectra of PCA solution before and after sensitization of NiO films. A surface coverage of 20 \pm 2 nmol cm⁻² was found.

Co was quantified by ICP-OES after digestion of NiO photocathodes in a mixture of conc. H₂SO₄ (1 mL) and HNO₃ (1 mL) by heating in a Biotage (Uppsala Sweden) SPX microwave reactor for 10 min at 100 °C. The resulting clear solution was filtered with a micron filter syringe, then diluted using deionized water and analyzed by ICP-OES (pre-calibrated with known concentrations of each element (sulfur and cobalt) of interest). The geometric surface areas of the dissolved electrodes were measured prior to being digested in order to determine the surface loading.

Photoelectrochemistry measurement

All the PEC experiments were performed in 0.1 M MES buffer at pH 5 using an Ag/AgCl (sat'd NaCl) reference electrode and a Pt wire counter electrode. Linear sweep voltammetric measurements were performed in a one-chamber cell degassed with Ar. A white LED was used to illuminate the photocathode, to generate a power density of approximatively 100 mW cm⁻². Long term chronoamperometric experiments were performed in an H-cell in order to avoid leakage from the Pt counter electrode onto the working electrode. The photocathode and the reference were in one compartment, while the Pt counter electrode was in the other one. All samples were degassed under Ar prior to measurements.

Clark electrode H₂ measurements

H₂ detection using a Clark electrode was performed based on a previously reported method.⁶⁵ Prior to measurements, the Unisense Clark electrode was calibrated using known concentrations of dissolved H₂ in 0.1 M MES buffer at pH 5 and measuring the Clark electrode response signal.



Conflicts of interest

The authors declare no conflict of interest.

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References

- 1 K. E. Dalle, J. Warnan, J. J. Leung, B. Reuillard, I. S. Karmel and E. Reisner, *Chem. Rev.*, 2019, **119**, 2752–2875.
- 2 Y. Tachibana, L. Vayssieres and J. R. Durrant, *Nat. Photonics*, 2012, **6**, 511–518.
- 3 M. G. Walter, E. L. Warren, J. R. McKone, S. W. Boettcher, Q. Mi, E. A. Santori and N. S. Lewis, *Chem. Rev.*, 2010, **110**, 6446–6473.
- 4 B. Zhang and L. Sun, *Chem. Soc. Rev.*, 2019, **48**, 2216–2264.
- 5 M. K. Brennaman, R. J. Dillon, L. Alibabaei, M. K. Gish, C. J. Dares, D. L. Ashford, R. L. House, G. J. Meyer, J. M. Papanikolas and T. J. Meyer, *J. Am. Chem. Soc.*, 2016, **138**, 13085–13102.
- 6 F. Li, K. Fan, B. Xu, E. Gabrielsson, Q. Daniel, L. Li and L. Sun, *J. Am. Chem. Soc.*, 2015, **137**, 9153–9159.
- 7 D. L. Ashford, M. K. Gish, A. K. Vannucci, M. K. Brennaman, J. L. Templeton, J. M. Papanikolas and T. J. Meyer, *Chem. Rev.*, 2015, **115**, 13006–13049.
- 8 Z. Yu, F. Li and L. Sun, *Energy Environ. Sci.*, 2015, **8**, 760–775.
- 9 E. A. Gibson, *Chem. Soc. Rev.*, 2017, **46**, 6194–6209.
- 10 J. Willkomm, K. L. Orchard, A. Reynal, E. Pastor, J. R. Durrant and E. Reisner, *Chem. Soc. Rev.*, 2016, **45**, 9–23.
- 11 E. Benazzi, J. Mallows, G. H. Summers, F. A. Black and E. A. Gibson, *J. Mater. Chem. C*, 2019, **7**, 10409–10445.
- 12 C. J. Wood, G. H. Summers, C. A. Clark, N. Kaeffer, M. Braeutigam, L. R. Carbone, L. D'Amario, K. Fan, Y. Farré, S. Narbey, F. Oswald, L. A. Stevens, C. D. J. Parmenter, M. W. Fay, A. La Torre, C. E. Snape, B. Dietzek, D. Dini, L. Hammarström, Y. Pellegrin, F. Odobel, L. Sun, V. Artero and E. A. Gibson, *Phys. Chem. Chem. Phys.*, 2016, **18**, 10727–10738.
- 13 L. Tong, A. Iwase, A. Nattestad, U. Bach, M. Weidelener, G. Götz, A. Mishra, P. Bäuerle, R. Amal, G. G. Wallace and A. J. Mozer, *Energy Environ. Sci.*, 2012, **5**, 9472–9475.
- 14 P. B. Pati, L. Zhang, B. Philippe, R. Fernández-Terán, S. Ahmadi, L. Tian, H. Rensmo, L. Hammarström and H. Tian, *ChemSusChem*, 2017, **10**, 2480–2495.
- 15 N. Kaeffer, J. Massin, C. Lebrun, O. Renault, M. Chavarot-Kerlidou and V. Artero, *J. Am. Chem. Soc.*, 2016, **138**, 12308–12311.
- 16 L. Li, L. Duan, F. Wen, C. Li, M. Wang, A. Hagfeldt and L. Sun, *Chem. Commun.*, 2012, **48**, 988–990.
- 17 B. Shan, B. D. Sherman, C. M. Klug, A. Nayak, S. L. Marquard, Q. Liu, R. M. Bullock and T. J. Meyer, *J. Phys. Chem. Lett.*, 2017, **8**, 4374–4379.
- 18 B. Shan, A. Nayak, M. K. Brennaman, M. Liu, S. L. Marquard, M. S. Eberhart and T. J. Meyer, *J. Am. Chem. Soc.*, 2018, **140**(20), 6493–6500.
- 19 K. Fan, F. Li, L. Wang, Q. Daniel, E. Gabrielsson and L. Sun, *Phys. Chem. Chem. Phys.*, 2014, **16**, 25234–25240.
- 20 M. A. Gross, C. E. Creissen, K. L. Orchard and E. Reisner, *Chem. Sci.*, 2016, **7**, 5537–5546.
- 21 Z. Ji, M. He, Z. Huang, U. Ozkan and Y. Wu, *J. Am. Chem. Soc.*, 2013, **135**, 11696–11699.
- 22 L. J. Antila, P. Ghamgosar, S. Maji, H. Tian, S. Ott and L. Hammarström, *ACS Energy Lett.*, 2016, **1**, 1106–1111.
- 23 S. Lyu, J. Massin, M. Pavone, A. B. Muñoz-García, C. Labrugère, T. Toupance, M. Chavarot-Kerlidou, V. Artero and C. Olivier, *ACS Appl. Energy Mater.*, 2019, **2**, 4971–4980.
- 24 B. van den Bosch, J. A. Rombouts, R. V. A. Orru, J. N. H. Reek and R. J. Detz, *ChemCatChem*, 2016, **8**, 1392–1398.
- 25 B. Shan, A. K. Das, S. Marquard, B. H. Farnum, D. Wang, R. M. Bullock and T. J. Meyer, *Energy Environ. Sci.*, 2016, **9**, 3693–3697.
- 26 K. L. Materna, N. Lalaoui, J. A. Laureanti, A. P. Walsh, B. P. Rimgard, R. Lomoth, A. Thapper, S. Ott, W. J. Shaw, H. Tian and L. Hammarström, *ACS Appl. Mater. Interfaces*, 2020, **12**, 4501–4509.
- 27 J. Huang, J. Sun, Y. Wu and C. Turro, *J. Am. Chem. Soc.*, 2021, **143**, 1610–1617.
- 28 E. Giannoudis, S. Bold, C. Müller, A. Schwab, J. Bruhnke, N. Queyriaux, C. Gablin, D. Leonard, C. Saint-Pierre, D. Gasparutto, D. Aldakov, S. Kupfer, V. Artero, B. Dietzek and M. Chavarot-Kerlidou, *ACS Appl. Mater. Interfaces*, 2021, **13**, 49802–49815.
- 29 L. D'Amario, J. Fohlinger, G. Boschloo and L. Hammarstrom, *Chem. Sci.*, 2018, **9**, 223–230.
- 30 S. Mori, S. Fukuda, S. Sumikura, Y. Takeda, Y. Tamaki, E. Suzuki and T. Abe, *J. Phys. Chem. C*, 2008, **112**, 16134–16139.
- 31 L. D'Amario, R. Jiang, U. B. Cappel, E. A. Gibson, G. Boschloo, H. Rensmo, L. Sun, L. Hammarström and H. Tian, *ACS Appl. Mater. Interfaces*, 2017, **9**, 33470–33477.
- 32 L. D'Amario, L. J. Antila, B. Pettersson Rimgard, G. Boschloo and L. Hammarström, *J. Phys. Chem. Lett.*, 2015, **6**, 779–783.
- 33 F. Li, R. Xu, C. Nie, X. Wu, P. Zhang, L. Duan and L. Sun, *Chem. Commun.*, 2019, **55**, 12940–12943.
- 34 C. E. Creissen, J. Warnan and E. Reisner, *Chem. Sci.*, 2018, **9**, 1439–1447.
- 35 C. E. Creissen, J. Warnan, D. Antón-García, Y. Farré, F. Odobel and E. Reisner, *ACS Catal.*, 2019, **9**, 9530–9538.
- 36 K. A. Click, D. R. Beauchamp, Z. Huang, W. Chen and Y. Wu, *J. Am. Chem. Soc.*, 2016, **138**, 1174–1179.



37 D. A. Hoogeveen, M. Fournier, S. A. Bonke, X.-Y. Fang, A. J. Mozer, A. Mishra, P. Bäuerle, A. N. Simonov and L. Spiccia, *Electrochim. Acta*, 2016, **219**, 773–780.

38 S. Lyu, Y. Farré, L. Ducasse, Y. Pellegrin, T. Toupance, C. Olivier and F. Odobel, *RSC Adv.*, 2016, **6**, 19928–19936.

39 R. J. Kamire, M. B. Majewski, W. L. Hoffeditz, B. T. Phelan, O. K. Farha, J. T. Hupp and M. R. Wasielewski, *Chem. Sci.*, 2017, **8**, 541–549.

40 N. Kaeffer, C. D. Windle, R. Brisse, C. Gablin, D. Leonard, B. Jousselme, M. Chavarot-Kerlidou and V. Artero, *Chem. Sci.*, 2018, **9**, 6721–6738.

41 H. Li, F. Li, B. Zhang, X. Zhou, F. Yu and L. Sun, *J. Am. Chem. Soc.*, 2015, **137**, 4332–4335.

42 L. Sévery, J. Szczerbiński, M. Taskin, I. Tuncay, F. Bandalise Nunes, C. Cignarella, G. Tocci, O. Blacque, J. Osterwalder, R. Zenobi, M. Iannuzzi and S. D. Tilley, *Nat. Chem.*, 2021, **13**, 523–529.

43 N. Lalaoui, P. Rousselot-Pailley, V. Robert, Y. Mekmouche, R. Villalonga, M. Holzinger, S. Cosnier, T. Tron and A. Le Goff, *ACS Catal.*, 2016, **6**, 1894–1900.

44 C. Gutiérrez-Sánchez, M. Pita, C. Vaz-Domínguez, S. Shleev and A. L. De Lacey, *J. Am. Chem. Soc.*, 2012, **134**, 17212–17220.

45 B. I. Ipe, K. G. Thomas, S. Barazzouk, S. Hotchandani and P. V. Kamat, *J. Phys. Chem. B*, 2002, **106**, 18–21.

46 A. Kotiaho, R. M. Lahtinen, N. V. Tkachenko, A. Efimov, A. Kira, H. Imahori and H. Lemmettyinen, *Langmuir*, 2007, **23**, 13117–13125.

47 R. K. Gupta and A. S. Achalkumar, *J. Org. Chem.*, 2018, **83**, 6290–6300.

48 M. T. Rojas, R. Koeniger, J. F. Stoddart and A. E. Kaifer, *J. Am. Chem. Soc.*, 1995, **117**, 336–343.

49 J. Liu, W. Ong, E. Román, M. J. Lynn and A. E. Kaifer, *Langmuir*, 2000, **16**, 3000–3002.

50 S. Roy and E. Reisner, *Angew. Chem.*, 2019, **131**, 12308–12312.

51 G. Chen and M. Jiang, *Chem. Soc. Rev.*, 2011, **40**, 2254–2266.

52 R. J. Lindquist, B. T. Phelan, A. Reynal, E. A. Margulies, L. E. Shoer, J. R. Durrant and M. R. Wasielewski, *J. Mater. Chem. A*, 2016, **4**, 2880–2893.

53 C. Zhou, J. Huang and Y. Yan, *Soft Matter*, 2016, **12**, 1579–1585.

54 S. Roy, Z. Huang, A. Bhunia, A. Castner, A. K. Gupta, X. Zou and S. Ott, *J. Am. Chem. Soc.*, 2019, **141**, 15942–15950.

55 J. R. Aranzaes, M.-C. Daniel and D. Astruc, *Can. J. Chem.*, 2006, **84**, 288–299.

56 B. Xu, L. Tian, A. S. Etman, J. Sun and H. Tian, *Nano Energy*, 2019, **55**, 59–64.

57 Y. Farré, F. Maschietto, J. Föhlinger, M. Wykes, A. Planchat, Y. Pellegrin, E. Blart, I. Ciofini, L. Hammarström and F. Odobel, *ChemSusChem*, 2020, **13**, 1844–1855.

58 F. Würthner, C. R. Saha-Möller, B. Fimmel, S. Ogi, P. Leowanawat and D. Schmidt, *Chem. Rev.*, 2016, **116**, 962–1052.

59 F. Odobel, Y. Farré, F. Maschietto, J. Föhlinger, M. Wykes, A. Planchat, Y. Pellegrin, E. Blart, I. Ciofini and L. Hammarström, *ChemSusChem*, 2020, **13**, 1844–1855.

60 M. Brust, M. Walker, D. Bethell, D. J. Schiffrin and R. Whyman, *J. Chem. Soc., Chem. Commun.*, 1994, 801–802.

61 O. Swiech, R. Bilewicz and E. Megiel, *RSC Adv.*, 2013, **3**, 5979–5986.

62 A. McNeillie, D. H. Brown, W. E. Smith, M. Gibson and L. Watson, *J. Chem. Soc., Dalton Trans.*, 1980, 767–770.

63 C. Paun, G. Słowiak, E. Lewin and J. Sá, *RSC Adv.*, 2016, **6**, 87564–87568.

64 N. Queyriaux, R. A. Wahyuno, J. Fize, C. Gablin, M. Wächtler, E. Martinez, D. Léonard, B. Dietzek, V. Artero and M. Chavarot-Kerlidou, *J. Phys. Chem. C*, 2017, **121**, 5891–5904.

65 C. D. Windle, J. Massin, M. Chavarot-Kerlidou and V. Artero, *Dalton Trans.*, 2018, **47**, 10509–10516.

66 K. L. Materna, A. M. Beiler, A. Thapper, S. Ott, H. Tian and L. Hammarström, *ACS Appl. Mater. Interfaces*, 2020, **12**, 31372–31381.

67 D. A. Hoogeveen, M. Fournier, S. A. Bonke, A. Nattestad, A. Mishra, P. Bäuerle, L. Spiccia, A. J. Mozer and A. N. Simonov, *J. Phys. Chem. C*, 2017, **121**, 25836–25846.

68 L. Tian, R. Tyburski, C. Wen, R. Sun, M. Abdellah, J. Huang, L. D'Amario, G. Boschloo, L. Hammarström and H. Tian, *J. Am. Chem. Soc.*, 2020, **142**, 18668–18678.

69 Z. Liu, D. Xiong, X. Xu, Q. Arooj, H. Wang, L. Yin, W. Li, H. Wu, Z. Zhao, W. Chen, M. Wang, F. Wang, Y. B. Cheng and H. He, *ACS Appl. Mater. Interfaces*, 2014, **6**, 3448–3454.

70 M. Abdellah, A. M. El-Zohry, L. J. Antila, C. D. Windle, E. Reisner and L. Hammarstrom, *J. Am. Chem. Soc.*, 2017, **139**, 1226–1232.

71 R. J. Dillon, L. Alibabaei, T. J. Meyer and J. M. Papanikolas, *ACS Appl. Mater. Interfaces*, 2017, **9**, 26786–26796.

72 M. Abdellah, S. Zhang, M. Wang and L. Hammarström, *ACS Energy Lett.*, 2017, **2**, 2576–2580.

73 A. Nattestad, A. J. Mozer, M. K. R. Fischer, Y. B. Cheng, A. Mishra, P. Bäuerle and U. Bach, *Nat. Mater.*, 2009, **9**, 31.

