Activating [FeFe] hydrogenase mimic for hydrogen evolution under visible light**

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Dedication ((optional))

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Abstract

Inspired by the active center of the natural [FeFe] hydrogenases, we designed a compact and precious metal-free photosensitizer-catalyst dyad (**PS-CAT**) for photocatalytic hydrogen evolution under visible light irradiation. **PS-CAT** represents a prototype dyad comprising π -conjugated oligothiophenes as light absorbers. **PS-CAT** and its interaction with the sacrificial donor 1,3-dimethyl-2-phenylbenzimidazoline were studied by steady-state and time-resolved spectroscopy coupled with electrochemical techniques and visible light-driven photocatalytic investigations. *Operando* EPR spectroscopy revealed the formation of an active [Fe^IFe⁰] species – in accordance with theoretical calculations – presumably driving photocatalysis effectively (TON \approx 210).



Nature provides paradigms for photocatalysis to generate solar fuels in a renewable and climate-neutral fashion. In this context, scientists have created various architectures to perform catalytic key processes, *e.g.* the hydrogen evolution reaction (HER), under light excitation. Commonly, 4d and 5d transition metals (*e.g.* Ru, Cd, Re, Ir, Pt) play a key role in molecular artificial HER photocatalysis – both as light-harvesters and catalysts – due to their favourable (photo)redox chemistry and stability.^[1] However, rare abundance or undesired side effects (*e.g.* toxicity) hamper a larger scale application.

Nature uses earth-abundant, inexpensive reaction centers in the [FeFe] hydrogenase enzyme for HER.^[2] Inspired by the natural paradigm [FeFe] hydrogenase mimics have been reported.^[3] Such catalysts, however, often rely on photosensitizers based on precious metals. Few exceptions are given by *e.g.* xanthene dyes, (zinc) porphyrins^[1d, 4] or carbon dots^[5] as light harvesters. Recently, we reported another fully precious metal-free example, a dibenzosilole photosensitizer directly attached to the catalyst, yield a TON of 539 for HER under UV light (254 nm).^[6] *π*-Conjugated oligothiophenes present metal-free, readily accessible and strong light absorbers in the visible spectral region. The optical and redox properties of such dyes, which are used *e.g.* in organic^[7] or dye-sensitized^[8] solar cells. Recently, Hammarström and colleagues combined a *π*-conjugated oligothiophene photosensitizer and a [FeFe] hydrogenase mimic in a cosensitized NiO photocathode.^[9]

We report a fully precious metal-free, visible-light absorbing, bioinspired and compact photocatalyst dyad **PS-CAT** as a prototype hybrid comprising π -conjugated oligothiophenes as photosensitizer for the light-driven HER. We present **PS-CAT**, its photocatalytic activity in the presence of the sacrificial donor 1,3-dimethyl-2-phenylbenzimidazoline (BIH) and a detailed mechanistic characterization. The photosensitizer unit represents a major improvement over the previous reported UV-active dibenzosilole light harvester, which showed promising electron relay properties to the covalently bound catalyst, leading to fundamental changes in the photophysical and photocatalytic features.

PS-CAT combines the light-harvesting properties of π -conjugated oligothiophenes (PS) with a bioinspired [FeFe] hydrogenase mimic (CAT). It was synthesized according to Scheme 1 (also detailed in Scheme S1) from the dithioacetato compound **PS**. Both, **PS** and **PS-CAT** were characterized by NMR, IR spectroscopy and mass spectrometry. The molecular structure of **PS-CAT** (Figure 1; Figure S2 and Table S1/S2; Supporting Information for crystallographic details) reveals a similar binding mode compared to that of a previous reported, structurally related complex.^[6] A view on the ab plane shows stacking of planar **PS-CAT** molecules (Figure S2b). The cyclic voltammogram of **PS-CAT** in CH₂Cl₂ at a scan rate of 0.2 Vs⁻¹ shows a quasi-reversible reduction event at $E_{1/2} = -1.61$ V vs. Fc⁺/Fc (Figure S8a).



Scheme 1: Synthetic pathway to **PS-CAT** *via* **PS**. i) BnNEt₃Cl, $[Pd(PPh_3)_4]$, K_2CO_3 , toluene; 60 °C, 12 h ii) *n*-BuLi, THF; -78 °C to r. t., 12 h iii) KSCOCH₃, THF; r. t., 20 h iv) LiAlH₄, Et₂O; 0 °C to r. t., 12 h v) Fe₃(CO)₁₂, toluene/NMP (20:1); r. t., 12 h.



Figure 1: Molecular structure and atom labeling scheme of **PS-CAT**. The ellipsoids represent a probability of 30 %. Hydrogen atoms are omitted for clarity. For selected bond lengths and angles see Table S2.

The photocatalytic proton reduction behaviour of **PS-CAT** was studied under irradiation with visible light in a 5:1 mixture of CH₃CN and *N*-Methyl-2-pyrrolidone (NMP) and with the two electron, one proton sacrificial donor 1,3-dimethyl-2-phenylbenzimidazoline (BIH)^[10] (1000 equivalents). The solvent mixture ensures an overall high dielectric constant for good solubility supporting the catalytic performance. Moreover, the weak base NMP can potentially react with the oxidized donor (BIH⁺⁺) to capture the proton, thus inhibiting potential back

electron transfer processes.^[11] The photocatalytic reactions were performed in a custom-made, modular 3D printed photoreactor setup (Figure S1). Figure 2 shows the hydrogen evolution time profile. Within the initial 23 hours the catalytic turn-over increases slightly sigmoidally, before the hydrogen evolution reaches a plateau at TON \approx 210 (TOF \approx 6.4 h⁻¹). This visiblelight activity is the highest reported for molecular dyads employing [FeFe] hydrogenase mimics.^[1d, 4, 12] No hydrogen was detected in corresponding experiments without **PS-CAT**, BIH or both or in the dark. Further photocatalytic results are given in Table S3.



Figure 2: Photocatalytic hydrogen production by **PS-CAT** (10 μ M) in CH₃CN/NMP (5:1) with BIH (1000 equivalents) upon irradiation at 455 nm in a 3D printed photoreactor platform (see inset). Hydrogen was quantified by GC-TCD with samples drawn from the head space. Each point was determined in duplicate and the error bars give the range of variation.

Figure 3a depicts the UV-vis spectra of **PS** and **PS-CAT**, which show a strong absorption in the visible spectral region ($\lambda_{max} = 409 \text{ nm}$, $\varepsilon \approx 8.1 \cdot 10^4 \text{ L} \text{ mol}^{-1} \text{ cm}^{-1}$ for **PS** and $\lambda_{max} = 410 \text{ nm}$, $\varepsilon \approx 6.1 \cdot 10^4 \text{ L} \text{ mol}^{-1} \text{ cm}^{-1}$ for **PS-CAT**) resulting from a $\pi\pi^*$ transition localized within the oligothiophene moiety (see S₁ in Figure 3d). For **PS-CAT** TDDFT reveals a dipole allowed charge-transfer transition in the visible region (see S₂ in Figure 3d) shifting electron density from a photosensitizer-localized π orbital to the [FeFe] hydrogenase mimic unit, *i.e.* into the σ^* orbital of the Fe-Fe bond of **PS-CAT**, contributing to the absorption in the visible range (Figure S12). Emission spectra ($\lambda_{exc} = 410$ nm) of both **PS** and **PS-CAT** display three nearly identical vibronic peaks at 455, 484 and 515 nm (shoulder) spaced by 1250 cm⁻¹, corresponding to a ground state vibrational mode coupled to the electronic transition (Table S6). The emission lifetimes were determined to 1 ns and the quantum yields are $\Phi_F = 0.54$ in **PS** and $\Phi_F = 0.28$ in **PS-CAT**. The reduced emission quantum yield in **PS-CAT** are associated with the population of a charge-separated state (S₂) by quantum chemical simulations. S₂, which is not available in **PS**, is in close energetic proximity to the $\pi\pi^*$ S₁ state.



Figure 3: a) Steady-state UV-vis absorption (solid lines) of **PS** (red) and **PS-CAT** (blue) and emission (dashed lines) in deaerated THF excited at 410 nm. b) Transient absorption spectra at indicated delay times of **PS** (top, blue) and **PS-CAT** (bottom, red) upon excitation at 420 nm in deaerated THF. c) Experimental (black, dashed) and simulated UV-vis absorption spectrum of **PS-CAT** in THF; electronic transitions of interest are labelled. d) Potential excited state relaxation cascades associated to electron and energy transfer competing with radiative recombination; relative energies of involved singlet (black) and triplet states (red) are given within their respective equilibrium structures. Electron transfer and energy transfer pathways both leading to singly reduced **PS-CAT** (D₀, ²MC, blue) upon BIH reduction are highlighted. Electronic characters are indicated by charge-density differences (CDDs, electronic transitions) and spin densities (opened-shell ground state); charge transfer takes place from red to blue (Table S7).

Nanosecond-transient absorption spectroscopy was performed to detect the impact of the [FeFe] hydrogenase mimic on the relaxation of the triplet state populated via intersystem crossing (ISC). The data (Figure 3b, Figure S6) for **PS** and **PS-CAT** exhibit similar spectral features. Both, **PS** and **PS-CAT** show a negative signal between 350 and 550 nm, which stems ground-state bleach and emission contributions between 460 and 550 nm.^[13] Excited-state absorption (ESA) is detected at wavelengths longer than 550 nm for both PS and PS-CAT which is characteristic for the ${}^{3}\pi\pi^{*}$ state of the sensitizer unit.^[14] TDDFT simulations on the dipole-allowed triplet-triplet excitations of **PS-CAT** within the equilibrium structure of the ${}^{3}\pi\pi^{*}$ ground state, see $T_2({}^3\pi\pi^*)$ in Figure 3d, associate these ESA features to ${}^3\pi\pi^*$ transitions of the excited oligothiophene at 600 and 529 nm, respectively (see T_{11} and T_{16} in Figure S14) confirming this assignment.^[15] At wavelengths above \approx 750 nm, a second negative signal can be detected which can be assigned to the triplet phosphorescence of the thiophenes.^[15] Both transient signals of **PS** and **PS-CAT** decay biexponentially with lifetimes of $\tau_1 = 20$ ns (reflecting the time-resolution of the experimental setup) and $\tau_2 = 530$ ns for **PS** and $\tau_1 = 20$ ns and $\tau_2 = 210$ ns for **PS-CAT**. The fast component indicates the decay of the residual emission. The long-lived component is associated with the decay of the ${}^{3}\pi\pi^{*}$ state of the sensitizer unit. The shortening in τ_2 reflects changes in the ${}^3\pi\pi^*$ state lifetime of the excited sensitizer in **PS**-**CAT** indicating the presence of additional relaxation channels occurring from the ${}^{3}\pi\pi^{*}$ state with an estimated time constant of 348 ns. Nevertheless, besides the changes in lifetime of the triplet state we do not observe any absorption features at 400, 580 and 700 nm, which could be ascribed to the reduced [FeFe] hydrogenase according to literature and UV-vis SEC data (Figure S10/S11). Hence, no direct indication for a charge-separated state (³CS, Figure 3d) is observed.^[13a, 16] The absence of these features indicates either that no charge transfer takes place to reduce the Fe-Fe unit or that the CS state is too short lived, recombines quickly and escapes experimental detection in absence of electron donor.

Quantum chemical simulations indicate two relaxation pathways for **PS-CAT**: i) electron transfer causing charge separation and which is accessible from the initially excited singlet states. ii) Energy transfer deactivating the ${}^{3}\pi\pi^{*}$ state (Figure 3d). The electron transfer pathway leads from the initially excited ${}^{1}\pi\pi^{*}$ state *via* ISC to the adjacent charge-separated T_{10/3}(FC/³CS equilibria) state (Figure 3d), resulting in a pronounced elongation of the Fe-Fe bond (Table S4) of the formally [Fe^IFe⁰] active site. Upon population of the ${}^{3}\pi\pi^{*}$ state, either *via* ${}^{1}\pi\pi^{*} \rightarrow {}^{3}\pi\pi^{*}$ ISC or *via* relaxation from the ${}^{3}CS$ state, an energy transfer pathway is available. In this case, hole transfer from the photooxidized photosensitizer to the photoreduced iron cluster yields an excited metal centered (${}^{3}MC$) state. During this process, the ESA decays as the initial configuration of the oligothiophene-based chromophore is reformed (Figure S14) in agreement with the experimental observations. In this ${}^{3}MC$ state, cleavage of the Fe-Fe bond (Table S4)

takes place as both the σ_{Fe-Fe} as well as the σ_{Fe-Fe}^* orbitals are singly populated (bond order: 0). The population of the ³MC state can be an explanation for the missing signatures of a CS state in the transient absorption experiment. These findings indicate that the presence of sacrificial agent under real catalytic conditions is crucial for the photoinduced reduction of the [FeFe] unit, a key step of the light-induced catalytic process. In this case, according to TDDFT simulations both relaxation channels, *i.e.* electron transfer and energy transfer, yield the same singly reduced doublet species of [Fe^IFe⁰] character (see D₀ spin density in Figure 3d), blue) – either upon reduction by BIH of the charge-separated species (³CS) or upon reduction of the ³MC state. Alternatively, the sacrificial agent could also interfere with the described relaxation processes already in an initial stage of the relaxation cascade reducing the sensitizer directly after excitation. The quenching behaviour of the fluorescence of **PS** by BIH gives first indication for such a fast process (Figure S4).

The role of the sacrificial agent is elucidated by *operando* UV-vis spectroscopy in the presence of sacrificial donor BIH (1000 equivalents) upon irradiation at 455 nm. After 30 minutes of irradiation, the absorption band of **PS-CAT** at 403 nm in CH₃CN/NMP (5:1) disappeared in favour of a characteristic band at 399 nm, which continues to build up within 17 hours (Figure S3a). Upon irradiation in absence of the donor the absorption at 403 nm decays only slowly, indicating reaction of **PS-CAT** with BIH (Figure S3c). The newly formed species decomposed slowly in the dark and quite rapidly in the presence of oxygen (Figure S3b). It is tentatively attributed to a sensitive follow-up product of the catalytic species, not to the catalytic species itself.

Figure 4: EPR spectrum of **PS-CAT** after four hours of illumination under catalytic conditions (30 µM **PS-CAT**). Illumination was carried out in a MD5 resonator and the EPR measurement was done at 4 K. The negative signal at 330 mT stems from resonator background. Inset: Spin density localized at the [FeFe] moiety of the singly reduced **PS-CAT** (doublet).

Operando electron paramagnetic resonance (EPR) spectroscopy reveals the formation of a reduced [FeFe] moiety under photocatalytic conditions. The catalytic mixture (30 μ M **PS-CAT**, 1000 equivalents BIH) was illuminated in the resonator at 270 K and measured at 4 K. The data show the formation of a paramagnetic [Fe^IFe⁰] intermediate (Figure 4), which has been postulated as key species in the light-induced HER mechanism.^[6c] No signal was detected in the dark or at room temperature (Figure S7a). The *g* values (*g*₁ = 2.003, *g*₂ = 1.99557, *g*₃ = 1.86197) derived from the EPR data are similar to those reported on [Fe^IFe⁰] species with the third *g* tensor being shifted further low field.^[17] This could be due to a low concentration of the [Fe^IFe⁰] intermediate in photochemical reduction consequently leading to a lower EPR spectral resolution compared to (electro)chemical reduction processes.^[17-18]

In summary, we synthesized a precious metal-free photocatalyst dyad **PS-CAT** for photocatalytic hydrogen generation under visible light. The dyad as prototype comprising π -conjugated oligothiophenes as light absorbers shows a remarkable, long-term photocatalytic activity, the best reported for comparable complexes in the visible spectral range. In accordance with theory, *operando* EPR spectroscopy using BIH as sacrificial donor reveals the generation of an active [Fe^IFe⁰] species, which presumably drives the light-induced hydrogen generation.

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Keywords

Photocatalytic hydrogen evolution • Quantum chemistry • [FeFe] hydrogenase • Cyclic voltammetry • Operando electron paramagnetic resonance (EPR) spectroscopy • Conjugated oligothiophenes • Photosensitizer-catalyst dyad • Time-resolved spectroscopy

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