Yield—not only Lifetime—of the Photoinduced Charge-Separated State in Iridium Complex–Polyoxometalate Dyads Impact Their Hydrogen Evolution Reactivity

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Abstract: Covalently linked photosensitizer–polyoxometalate (PS-POM) dyads are promising molecular systems for light-induced energy conversion processes, such as “solar” hydrogen generation. To date, very little is known of their fundamental photophysical properties which affect the catalytic reactivity and stability of the systems. PS-POM dyads often feature short-lived photoinduced charge-separated states, and the lifetimes of these states are considered crucial for the function of PS-POM dyads in molecular photocatalysis. Hence, strategies have been developed to extend the lifetimes of the photoinduced charge-separated states, either by tuning the PS photophysics or by tuning the POM redox properties. Recently, some of us reported PS-POM dyads based on cyclometalated IrIII complexes covalently linked to Anderson-type polyoxometalate. Distinct hydrogen evolution reactivity (HER) of the dyads was observed, which was tuned by varying the central metal ion M of the POMMx (M = MnII, CoIII, FeIII). In this manuscript, the photoinduced electron-transfer processes in the three Ir-POM dyads are investigated to rationalize the underlying reasons for the differences in HER activity observed. We report that upon excitation of the IrIII complex, ultrafast (sub-ps) charge separation occurs, leading to different amounts of the charge-separated states (IrII-POM) generated in the different dyads. However, in all dyads studied, the resulting IrII-POM species are short-lived (sub-ns) when compared to reference electron acceptors (e.g. porphyrins or fullerenes) reported in the literature. The reductive quenching of IrII-POM by a sacrificial donor, triethyl amine (1 M), to generate the intermediate Ir-POM, is estimated to be very efficient (70–80%) for all dyads studied. Based on this analyses, we conclude that the yield instead of the lifetime of the IrII-POM charge-separated state determines the catalytic capacity of the dyads investigated. This new feature in the PS-POM photophysics could lead to new design criteria for the development of novel PS-POM dyads.

Introduction

Converting sunlight into chemical fuels is a key goal for green and sustainable energy technologies.[1] The generation of “solar hydrogen” requires the combination of several fundamental steps such as light-harvesting, charge-separation, charge-accumulation and catalysis (i.e. reduction of protons to H2).[2] One strategy in artificial photosynthesis is to integrate all these fundamental processes into one supramolecular assembly, which at least contains a photocatalytic unit combined with an acceptor as catalytic unit for reduction reactions, respectively.[2] As a result of this molecular design, photoexcitation of the supramolecular assembly leads to the formation of a charge-separated state (CSS), in which the photoactive donor is oxidized, while the catalytic center is reduced. Ideally, the CSS is long lived and can thus react in a collision-induced reaction with a sacrificial electron donor or other reactants to give the desired product.[2] To date a number of typically organic electron acceptors such as porphyrin or fullerene have been combined in this fashion with photoactive transition metal...
complex photosensitizers as donors.\textsuperscript{2c-3} More recently, molecular metal oxides, or polyoxometalates (POMs), have emerged as inorganic electron acceptors in the field of artificial photosynthesis.\textsuperscript{4} The beneficial properties of POMs as electron acceptors arise from their strong electron acceptor properties, multi-electron redox capabilities and catalytic activity in their reduced states.\textsuperscript{4} The functionalization of POMs with visible light absorbing PS is critical, as POMs themselves typically only absorb light in the UV region. Thus, the design of covalently linked PS-POM dyads has recently emerged as a new design concept in POMs energy conversion.\textsuperscript{5}

However, due to the synthetic challenges involved,\textsuperscript{6,7} currently there are only a few pioneering studies available. In these initial studies, Re\textsuperscript{I,II}, Ru\textsuperscript{III,II} and Ir\textsuperscript{III} complexes, porphyrins as well as BODIPY dyes have been used as photosensitizers, leading to possible applications in light-driven hydrogen evolution and photocatalysis.\textsuperscript{8} In these studies, the main focus was on Keggin-, Dawson- and Anderson-derivatives as their covalent functionalization is well-established.\textsuperscript{9} However, in the reported PS-POM dyads either no charge-separated states, that is, PS\textsuperscript{−}−POM\textsuperscript{+}, or only relatively short-lived PS\textsuperscript{−}−POM\textsuperscript{+} were detected upon excitation of the photosensitizers.\textsuperscript{10-13} The lifetimes of PS\textsuperscript{−}−POM\textsuperscript{+} range from a few ps to the longest value reported up to now, which is about 500 ns.\textsuperscript{3,14} However, the lifetime of the photoinduced primary CSS, that is, PS\textsuperscript{−}−POM\textsuperscript{+}, is generally considered key to the function of PS-POM dyads in an artificial photosynthetic Scheme, as a long-lived CSS allows for reductive quenching of the oxidized PS by a sacrificial donor and subsequent light-driven or—depending on the specific reaction mechanism—dark processes. As POM-based dyads do not excel in terms of long lifetime of the CSS, this might become one of the limiting factors for developing POM-based artificial systems for solar fuels generation via multi-electron processes. Compared to the more frequently utilized Keggin- and Dawson-POMs, photoinduced dynamics in Anderson-POMs based assemblies are scarcely investigated.\textsuperscript{15} Hasenkopf and co-workers reported the linkage of Zn\textsuperscript{II} porphyrins to Dawson- and Anderson-POM and observed that photoinduced electron transfer takes place from the excited Zn\textsuperscript{II} porphyrin to the Dawson- but not to the Anderson-POM.\textsuperscript{16}

Some of us recently reported three PS-POM dyads based on Anderson-POM as electron acceptors, which were covalently linked to two cyclometalated Ir\textsuperscript{III} complexes as photosensitizer and electron donor (\(\text{[(nBu}_{3}\text{N})\text{[MMO}_{6}\text{O}_{19}\text{]}(\text{OCH}_{3})_{5}\text{CNICH}_2\text{H}_2\text{N}_2\text{]}_2\)) molecular structure see the inset in Figure 1).\textsuperscript{5} In these dyads, the central metal ion M of the POM\textsubscript{6} was varied from \(M = \text{Mn}^{3+}\) to \(\text{Co}^{3+}\) to \(\text{Fe}^{3+}\).\textsuperscript{5} It was found that all Ir-POM\textsuperscript{−} dyads displayed hydrogen evolution reactivity (HER) in the presence of triethylamine as a sacrificial electron donor and acetic acid as the proton source.\textsuperscript{6} The HER activity decreased from Ir-POM\textsubscript{Mn} (TON = 80, measured over seven days) to Ir-POM\textsubscript{Co} (TON = 34) to Ir-POM\textsubscript{Fe} (TON = 20).\textsuperscript{6} Electrochemical data and DFT calculations indicated that the central M alters the redox properties of POM\textsubscript{M}.\textsuperscript{6} For \(M = \text{Mn}^{3+}\) the LUMO energy was the lowest.\textsuperscript{6} The modulation of the LUMO energy (3.95, 4.58 and 4.74 eV for POM\textsubscript{Mn}, POM\textsubscript{Co} and POM\textsubscript{Fe}, respectively)\textsuperscript{6} in turn leads to a variation of the driving force for electron transfer, and hence, distinct rate constants for charge separation (which would significantly affect the yield of Ir\textsuperscript{III}−POM\textsubscript{M} and charge recombination (the stability of Ir\textsuperscript{III}−POM\textsubscript{M}) are expected.

In this work, we explore the charge separation and charge recombination underlying the different HER activity of the aforementioned Ir-POM\textsubscript{M} dyads. In order to do so, we performed femtosecond and nanosecond time-resolved spectroscopy and combine them with steady-state and time-resolved emission spectroscopy. The data presented indicates that the yield of the initial CSS is an important factor influencing the overall catalytic activity of Ir-POM\textsubscript{M}. Thus, not only the lifetime of the CSS, \(\text{Ir}^{3+}−\text{POM}^{\cdot}\), is to be considered as a photophysical metric to relate to the activity of such photocatalytically active dyads. Furthermore, on a more detailed notice, this work provides to the best of our knowledge the first example of a special characterization of a CSS in Anderson-POM based PS-POM dyads.

Results and Discussion

Steady-state emission spectra of Ir-POM\textsubscript{M} dyads are depicted in Figure 1. The emission spectra of each compound were recorded at an optical density of 0.08 upon excitation at 400 nm. Compared to Ir, the emission intensity from the Ir\textsuperscript{III} unit is significantly decreased in Ir-POM\textsubscript{Mn} and Ir-POM\textsubscript{Co} (by 86 and 79\% respectively). For Ir-POM\textsubscript{Co}, the emission intensity is only reduced by 58\%. The partially quenched emission indicates that combining the Ir\textsuperscript{III} complex with the POM\textsubscript{M} provides additional non-radiative decay channels for the excited Ir-POM\textsubscript{M} dyads.
Photoinduced dynamics

The non-radiative decay pathways within the Ir-POM\textsubscript{M} dyads were studied by fs transient absorption (TA) spectroscopy. Figure 2 shows the fs TA spectra of Ir, Ir-POM\textsubscript{Co}, and Ir-POM\textsubscript{Mn} obtained in aerated DMF upon excitation at 400 nm. The data recorded for Ir-POM\textsubscript{Fe} are depicted in the ESI (Figure S1) as all dyads show very similar spectral features but different kinetics. The fs TA spectra of Ir upon excitation of the singlet metal-to-ligand charge transfer (MLCT) states resemble the spectral-temporal evolution as previously reported in literature\cite{7}. A broad excited-state absorption (ESA) spanning from 350 to 750 nm corresponds to the absorption of \(^3\)MLCT states in the photosensitizer and is observed immediately upon photoexcitation.\cite{7b,c} Subsequently, two bands at 375 and at around 500 nm increase on a sub-10 ps timescale (Figure 2a, b). Based on spectro-electrochemical data, the ESA band at 375 nm (the orange dashed line in Figure 2c, for more information see Figure S2b) is assigned to the absorption of the reduced bpy ligand.\cite{7b,c,8} Its increase at early times is attributed to interligand electron transfer (ILET) from an upper-lying \(^3\)MLCT\textsubscript{bpy} state to the lower-lying \(^3\)MLCT\textsubscript{bpy} state, a processes which has been established in literature.\cite{7a,b}

Ir-POM\textsubscript{Co} and Ir-POM\textsubscript{Mn} initially (i.e. at 0.4 ps after photoexcitation) display similar transient absorption features as Ir, that is, a rather broad but red-shifted ESA band in the visible region (Figure 2d, g). However, between 0.4 and 10 ps the dyads reveal spectral changes different to Ir: For one, the evolution of the ESA band in the visible region is faster in the dyads than in Ir. At 1.0 ps the band sharpens compared to Ir, that is, the FWHM (full width at half maximum) of the spectrum is reduced by 5, 16 and 11\% compared to the data at 0.4 ps for Ir, Ir-POM\textsubscript{Co}, and Ir-POM\textsubscript{Mn}, respectively. Meanwhile the spectral intensity already reaches a maximum (Figure 2d, g). Additionally, the ESA band in the UV region decreases instead of increasing as observed for Ir (Figure 2a, d and g). This is seen directly from the kinetic trace at 375 nm (blue lines in Figures 2b, e, h), which reflects the formation of a charge-separated state, that is, \(\text{Ir}^+\text{-POM}^-\). The one-electron-reduced POM\textsubscript{M} (i.e. POM\textsubscript{M}C\textsubscript{2}@) has a negative absorption below 400 nm (Figure S3 a,c, e), which apparently compensates the positive absorption signal of bpy\textsuperscript{2-} (Figure 2c, orange dashed line). At delay times longer than 100 ps, a pronounced decay of the overall TA signal is observed (Figure 2e, h) for both Ir-POM\textsubscript{Co} and Ir-POM\textsubscript{Mn}. However, within the experimentally accessible delay time range of 1.8 ns the decay is not complete, that is, a long-lived component is apparent in the fs data.

The quantitative interpretation of the fs TA data is based on a global fit.\cite{9} For all compounds, two decay components and an offset are sufficient to describe the experimental data. The
decay-associated spectra (DAS, see ESI for a description on the DAS) and the corresponding characteristic time constants are given in Figure 2c, f, i. The kinetic components obtained for Ir are in agreement with previous work by Lochbrunner on the same complex.\(^{[7b,c]}\) Following Lochbrunner’s work, we assign the component associated with \(t_1 = 1.1\) ps (Figure 2c) to vibrational relaxation within the \(^1\text{MLCT}\) manifolds involving both ppy and bpy ligands. This is followed by ILET from a \(^1\text{MLCT}\text{ppy}\) to a \(^3\text{MLCT}\text{ppy}\) state. This process is characterized by \(t_2 = 11\) ps. The long-lived species represents the long-lived thermalized \(^3\text{MLCT}\text{ppy}\) state, the decay of which to the electronic ground state is beyond the experimentally accessible delay-time range.

Considering the DAS-analysis of the fs TA data of the dyads, the fastest kinetic component is associated with a characteristic decay time \(t_1 = 0.5\) ps. The corresponding spectral changes resemble the spectral shape of \(t_1\) in Ir, nonetheless comprising a slight red-shift compared to the spectra of the photosensitizer only (Figure 2f, i and Figure S1c; Figure S5a, c, e). Furthermore, DAS(\(t_1\)) reveals a broader negative feature compared to the DAS(\(t_1\)) of Ir. Specifically, in the dyads an increase of the transient absorption at 550 nm is associated with the \(t_1\)-component (Figure 2f, i, which is not present in Ir. This ESA increase is due to the absorption of the reduced form of POM\(_{\text{m}}\) (i.e. POM\(_{\text{m}^-}\), Figure S4) combined with possible contributions from the oxidized Ir\(^{III}\) center (i.e. Ir\(^{IV}\)), Figure S2a. Note that the Ir photosensitizer in the dyads has additional substituents on the bpy ligand compared to reference Ir which might cause slight spectral difference to the MLCT and charge-separated states). This indicates that in Ir-POM\(_{\text{m}}\) electron transfer from the photo-excited Ir\(^{IV}\) complex to the POM\(_{\text{m}}\) occurs rapid and in concert with vibrational relaxation of \(^1\text{MLCT}\) states. The fast formation of Ir\(^{II}^-\text{POM}_{\text{m}^-}\), which indicates effective electronic coupling between the initially excited MLCT state of the photosensitizer and the POM\(_{\text{m}}\) is corroborated by the difference spectrum of the DAS(\(t_1\)) in Ir and Ir-POM\(_{\text{m}}\) (see Figure S5). Notably, the intensities of the TA signal at 550 nm in DAS (\(t_1\)), which accounts for the charge-separation process, vary within the set of dyads (see Figure 3). Namely, DAbs at 550 nm in Ir-POM\(_{\text{m}^-}\) is roughly 2 times as high as that in Ir-POM\(_{\text{c}o}\) (Figure 3). This indicates different yields of Ir\(^{II}^-\text{POM}_{\text{m}^-}\) being generated in the dyads, that is, the yield of Ir\(^{II}^-\text{POM}_{\text{m}^-}\) is (two times) higher than that of Ir\(^{II}^-\text{POM}_{\text{c}o}\).

The 11-ps-component related to ILET in Ir is absent in the dyads. Instead, a hundreds of ps component is apparent, that is, \(t_3 = 540\) and 290 ps for Ir-POM\(_{\text{c}o}\) and Ir-POM\(_{\text{c}m}\), respectively. This component reflects the decay of the charge-separated state Ir\(^{II}^-\text{POM}_{\text{m}^-}\), whose absorption has been simulated based on UV/Vis spectroelectrochemical data (Figure 2f, i, S2 and S3). In addition to \(t_1\) and \(t_2\), all dyads reveal a long-lived state, whose decay extends beyond the experimentally accessible delay-time window (~1.8 ns). The corresponding infinite component observed for Ir-POM\(_{\text{c}o}\) shows nearly identical features to that in Ir (Figure 2f), that is, a strong band below 400 nm and a moderate band (asymmetric and flat) in the visible region. This may indicate a decay of the \(^1\text{MLCT}\text{ppy}\) state in Ir-POM\(_{\text{c}o}\). In contrast, the infinite component in Ir-POM\(_{\text{c}m}\) (Figure 2f) and Ir-POM\(_{\text{c}e}\) (Figure S1d) displays a rather distinct absorption at 550 nm. This points to a different long-lived state in Ir-POM\(_{\text{c}m}\) and Ir-POM\(_{\text{ce}}\) compared to Ir-POM\(_{\text{c}o}\).

The nature of the long-lived state was assessed by ns TA spectroscopy (Figure 4). The ns TA spectra match the features of the infinite component in fs TA data. For Ir-POM\(_{\text{c}o}\), similar to Ir (Figure S6), the ns TA signal decays mono-exponentially with a characteristic time constant of 120 ns (Figure 4c, d). For Ir-POM\(_{\text{c}m}\) and Ir-POM\(_{\text{c}e}\), the shoulder at 550 nm disappears after 30 ns as revealed by the ns data in Figures 4b and S10a, while the overall decay of the differential absorption is slower. A global fit of the ns TA data yields a comparably short-lived component, \(t_{n1} = 30\) ns, and a relatively long-lived one, \(t_{n2} = 120\) ns (Figure S10b and S12b). The 120 ns-component in Ir-POM\(_{\text{c}c}\) resembles the key spectral features of the 100 ns-component in Ir (Figure 4c and S14). Furthermore, the changes of its spectra and lifetime upon changing the solvent polarity (Figures S16–S20) correlate with the corresponding changes observed for Ir. Thus, we assign the 120 ns-component to the decay of the \(^1\text{MLCT}\text{ppy}\) state. The significant increase of \(t_{n2}\) upon exclusion of oxygen from the solution (120 vs. 640 ns in aerated and deaerated DMF, see Figure 4d, Figures S9, S11, S13 and S15) corroborate this assignment.\(^{[10]}\)

Considering the 30 ns component, the ns TA results in DMF and DMSO indicate that it is a bright state whose lifetime is independent of solvent polarity (\(t_{n1} = 30\) ns in DMF and \(t_{n1} = 40\) ns in DMSO, Figures S10d, S12d, S18d and S19d). These properties exclude the possibility of a CSS being formed. On the contrary, \(t_{n2}\) doubles upon exclusion of oxygen from DMF (inset in Figure 4d; Figure S11d and S13d). Furthermore, its energetic position appears to be drastically shifted when changing the solvent from DMF (emission at 550 nm) to DMSO (emission at 680 nm). Nevertheless, the nature of the 30 ns component is beyond the scope of this study because the different HER activity of Ir-POM\(_{\text{m}^-}\) is not impacted by the presence of this state: Both dyads, which reveal this component, Ir-POM\(_{\text{c}m}\) and Ir-POM\(_{\text{ce}}\) display rather different HER activity (TON = 80 vs. 20, respectively).\(^{[12]}\) Ir-POM\(_{\text{c}c}\), whose ns-decay does not reveal the 30-ns component, displays an intermediate TON of 34, which is 70% higher than that of Ir-POM\(_{\text{c}e}\).\(^{[12]}\)
Relaxation model

Scheme 1 summarizes the photophysical picture of Ir-POM as it emerges from the time-resolved spectroscopy. According to the fs TA results, photoinduced electron transfer in the dyads occurs concerted with vibrational energy dissipation, indicating that electron transfer is most efficient from vibrationally hot 1MLCT states (Scheme 1). Considering electron transfer from the individual 1MLCT states intrinsic to the system, the driving force for electron transfer can be estimated by the Rehm–Weller equation \[ \Delta G_{\text{elas}} = E_{\text{D}} - E_{\text{A}} - \int \left( -e^2 / 4\pi \varepsilon_{\text{D}} \rho_{\text{D}} \right) \, \text{d}x. \]

As the 1MLCTppy state is energetically higher than the 1MLCTppy state (i.e., a larger \( E_{\text{gap}} \)), a more negative driving force will be induced. Hence, we cannot exclude the direct through-space electron transfer from the hot 1MLCTppy to the POM (Scheme 1). The presence of this direct electron-transfer pathway explains the absence of ILET in the dyads Ir-POM as compared to Ir, where ILET dominates the 11-ps process. Besides, ultrafast electron transfer from the initially excited hot 1MLCT states (see resonance Raman data in Figure S21), which would have more favorable driving forces, might be operative as well. However, due to the temporal resolution (~110 fs) of the fs TA setup such ultrafast process cannot be explored.

\( \tau_2 \) in the dyads is attributed to charge recombination yielding the decay of Ir\(^{1-}\)-POM\(^{+}\). The lifetime of Ir\(^{1-}\)-POM\(^{+}\) depends on the central metal ion \( M \), that is, Ir\(^{1-}\)-POM\(^{+}\) \((290 \text{ ps}) < \text{Ir}^{2+}\)-POM\(^{+}\) \((430 \text{ ps}) < \text{Ir}^{3+}\) -POM\(^{+}\) \((540 \text{ ps}) \). In hydrogen evolution experiment investigating the Ir-POM\(_{\text{Co}}\) dyads studied here, the concentration of the sacrificial donor (1 m) was 10,000 higher than that of the dyads (0.1 m). Under these conditions, the diffusion-limited rate constant of the sacrificial donor for quenching is about \( 7.6 \times 10^9 \text{ s}^{-1} \) (calculated by multiplying the diffusion-limited rate constant in DMF, \( 7.6 \times 10^9 \text{ m}^{-1} \text{s}^{-1} \), with the concentration of sacrificial donor, 1 m). This rate corresponds to a characteristic time constant of 126 ps. This indicates that the characteristic lifetimes of the Ir\(^{1-}\)-POM\(^{+}\) (\( \tau_2 = 290 - 540 \text{ ps} \)) provides a sufficiently large temporal window for the interaction with the sacrificial donor, yielding Ir-POM\(^{+}\) as an essential intermediate in the photocatalytic cycle. Based on the time constants for diffusion and charge recombination (\( \tau_2 \)), the quenching efficiency by the sacrificial donor is estimated to be 70 and 80% in Ir-POM\(_{\text{Mn}}\) and Ir-POM\(_{\text{Co}}\), respectively. However, the HER activity of Ir-POM\(_{\text{Mn}}\) (TON = 80) was reported to be 2.4 times as high as that of Ir-POM\(_{\text{Co}}\) (TON = 34). Therefore, we conclude that the HER activity of the dyads as observed under conventionally employed experimental conditions seems not governed by the lifetime of the charge-separated state Ir\(^{1-}\)-POM\(^{+}\). Instead, this study suggests that the yield of Ir\(^{1-}\)-POM\(^{+}\) plays an important role in the catalytic capacity of Ir-POM. As discussed above, the yield of Ir\(^{1-}\)-POM\(^{+}\) is roughly twice the yield of Ir\(^{1-}\) -POM\(^{+}\). This finding is consistent with the ratio of the HER activity of the respective dyads. Furthermore, from the perspective of photostability, more efficient hot 1MLCT-POM electron transfer results in a reduced population of the long-lived thermalized 1MLCT state. Thereby, 1MLCT-POM electron transfer reduces the potential for destructive side-reactions taking place from the high-energy 1MLCT state, which might take place.
modify the redox properties of the POM\textsubscript{\text{\textregistered}} acceptors.\textsuperscript{(52)} As a consequence of this modification, different photophysical properties of the Ir-POM\textsubscript{\textregistered} dyads can be expected. Ir-POM\textsubscript{\textregistered} dyads were investigated spectroscopically. The central metal ion \(M\) of POM\textsubscript{\textregistered} is the lowest state) in frozen conditions. Hence, an increased yield of the CSS might benefit the catalytic HER activity twofold by increasing the yield of a critical charge-transfer intermediate of the catalytic cycle and by diminishing the potential for detrimental side reactions. Our work points out that for a good performance of the catalysts the photoinduced CSS should be sufficiently long-lived for the subsequent reactions (i.e. to react with external reagents according to the specific reaction schemes). When the decay dynamics of the CSS itself are comparable to the rates of subsequent reactions, extending the lifetimes of the CSS would not improve the function of the catalysts. At this point, the yields of CSS have to be considered. In this respect, proper molecular design for fast charge separation (e.g. by inducing a larger driving force or a stronger electronic coupling), which can compete with the decay of the excited state itself (e.g. the cooling process in Scheme 1), could promote the yields of CSS.

Conclusions

The photoinduced electron transfer dynamics underlying the distinct catalytic capacity (Ir-POM\textsubscript{Mo} > Ir-POM\textsubscript{Co} > Ir-POM\textsubscript{Mn}) of three covalently linked Ir-POM\textsubscript{\textregistered} dyads were investigated spectroscopically. The central metal ion \(M\) of POM\textsubscript{\textregistered}, that is, \([\text{MnMo}_2\text{O}_4\text{O}][\text{\textregistered}]\), was changed from Mn\textsuperscript{3+} to Co\textsuperscript{3+} to Fe\textsuperscript{3+} to

### Experimental Section

**General**

Steady-state UV/Vis absorption spectra were recorded in a quartz cell with 1 mm path length (for fs transient absorption experiment, JASCO V-670 spectrophotometer) and with 1 cm path length (for ns transient absorption experiment, Cary 5000 UV/Vis spectrometer, Varian, USA). For all time-resolved experiments, the stability of samples was ensured by recording the steady-state UV/Vis absorption spectra before and after every measurement. The steady-state emission spectra were recorded in a quartz cell with 1 cm path length on a FLS980 spectrofluorimeter (Edinburgh).

**Electrochemistry and spectroelectrochemistry**

Cyclic voltammetry (CV) and spectroelectrochemistry (SEC) measurements were performed in a home-built three-electrode thin-layer cell with a path length of 1 mm. The three-electrode system consists of a glassy carbon working electrode, a platinum wire counter electrode and an Ag/AgCl reference electrode. CV and potential-controlled monitoring were performed using a computer-controlled potentiostat (VersaSTAT 3, Princeton Applied Research). All potentials given in the manuscript refer to the ferrocene/ferro-cenium couple as internal standard. The corresponding UV/Vis spectra were recorded on a single-beam spectrometer (Avantes, Aivalight-DH-S-BAL) at room temperature.
Time-resolved spectroscopy

Femtosecond (fs) transient absorption spectra were collected by using a previously reported home-built pump-probe laser system which is based on an amplified Ti: Sapphire oscillator (Libra, Coherent Inc.). All compounds were excited by pump pulse centered at 400 nm (TOPASwhite, Lightconversion Ltd.) with a duration of 110 fs. The power of the pump beam was kept at 0.4 mW and the beam diameter of the pump was 145 μm at the sample position. This corresponds to 0.95 × 10^{15} photons m^{-2} per pulse. A white light supercontinuum generated by focusing a fraction of the fundamental in a rotating CaF₂ plate is used to probe the samples in a wide spectral range (340 to 750 nm). The probe beam is delayed in time with respect to the pump beam by means of an optical delay line and the polarization between probe and pump is set at the magic angle (54.7°). Each solution (optical density ca. 0.2 at the excitation wavelength) was kept in a 1 mm quartz cuvette. Transient absorption data were displayed after chip correction. The transient absorption data was analyzed by a global multi-exponential fit after exclusion of a temporal window of 200 fs around time-zero in order to avoid contributions of the coherent-artifact region to the data analysis. Furthermore, a spectral band of 20 nm around the pump-wavelength is omitted from the data analysis due to pump-scatter in this spectral range.

A commercial multi-pump head is used to probe the samples in a wide spectral range (340 to 750 nm). The probe beam is delayed in time with respect to the pump beam by means of an optical delay line and the polarization between probe and pump is set at the magic angle (54.7°). Each solution (optical density ca. 0.2 at the excitation wavelength) was kept in a 1 mm quartz cuvette. Transient absorption data were displayed after chip correction. The transient absorption data was analyzed by a global multi-exponential fit after exclusion of a temporal window of 200 fs around time-zero in order to avoid contributions of the coherent-artifact region to the data analysis. Furthermore, a spectral band of 20 nm around the pump-wavelength is omitted from the data analysis due to pump-scatter in this spectral range.

Resonance Raman spectroscopy

Resonance Raman (RR) spectra were recorded through excitation by a 405 nm diode laser (TopMode-405-HP, Toptica, Germany) and detected by an iSPlane 160 spectrometer (Princeton Instruments, USA) with an entrance slit width of 0.05 mm, a focal length of 750 mm, and grating 2400 grooves mm^{-1}. The excitation energy was attenuated to around 8 mW. The Raman signals were recorded with a thermoelectrically cooled PIIXIS eXcelon camera (Princeton Instruments, USA). The Raman spectra were initially baseline corrected and normalized with respect to a solvent band, that is, to the signal at 1404 cm⁻¹ for DMF.

Deutscher Akademischer Austauschdienst DAAD for a Ph.D. fellowship.

Conflict of interest

The authors declare no conflict of interest.

Acknowledgements

This research is supported by the Catalight CRC/TRR 234 (project number 364549901, projects A1 and A4 and Z2) funded by the Deutsche Forschungsgemeinschaft (DFG). S.M. thanks the...
Note: the calculated time constant for intermolecular quenching only considered the rate of encounter between the photo-excited dyad and the sacrificial donor. In reality this process would kinetically compete with backward electron transfer within the solvent cage to reform the original ground-state species. These processes together with the escape of the redox species out of the cage into bulk solution determine the yield of the Ir-POM$_{\text{red}}$ intermediate. But we do not expect that the intermolecular kinetics would depend drastically on the nature of the reduced state of Ir-POM$_{\text{ox}}$.