

Electro- and Photoinduced Interfacial Charge Transfers in Nanocrystalline Mesoporous TiO₂ and TiO₂/Iron Porphyrin Sensitized Films under CO₂ Reduction Catalysis

Beatriu Domingo-Tafalla, Tamal Chatterjee,* Federico Franco, Javier Perez Hernandez, Eugenia Martinez-Ferrero, Pablo Ballester,* and Emilio Palomares*



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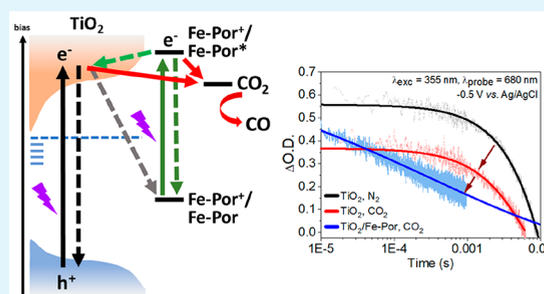
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ABSTRACT: Electro- and photochemical CO₂ reduction (CO₂R) is the quintessence of modern-day sustainable research. We report our studies on the electro- and photoinduced interfacial charge transfer occurring in a nanocrystalline mesoporous TiO₂ film and two TiO₂/iron porphyrin hybrid films (*meso*-aryl- and *β*-pyrrole-substituted porphyrins, respectively) under CO₂R conditions. We used transient absorption spectroscopy (TAS) to demonstrate that, under 355 nm laser excitation and an applied voltage bias (0 to −0.8 V vs Ag/AgCl), the TiO₂ film exhibited a diminution in the transient absorption (at −0.5 V by 35%), as well as a reduction of the lifetime of the photogenerated electrons (at −0.5 V by 50%) when the experiments were conducted under a CO₂ atmosphere changing from inert



N₂. The TiO₂/iron porphyrin films showed faster charge recombination kinetics, featuring 100-fold faster transient signal decays than that of the TiO₂ film. The electro-, photo-, and photoelectrochemical CO₂R performance of the TiO₂ and TiO₂/iron porphyrin films are evaluated within the bias range of −0.5 to −1.8 V vs Ag/AgCl. The bare TiO₂ film produced CO and CH₄ as well as H₂, depending on the applied voltage bias. In contrast, the TiO₂/iron porphyrin films showed the exclusive formation of CO (100% selectivity) under identical conditions. During the CO₂R, a gain in the overpotential values is obtained under light irradiation conditions. This finding was indicative of a direct transfer of the photogenerated electrons from the film to adsorbed CO₂ molecules and an observed decrease in the decay of the TAS signals. In the TiO₂/iron porphyrin films, we identified the interfacial charge recombination processes between the oxidized iron porphyrin and the electrons of the TiO₂ conduction band. These competitive processes are considered to be responsible for the diminution of direct charge transfer between the film and the adsorbed CO₂ molecules, explaining the moderate performances of the hybrid films for the CO₂R.

KEYWORDS: electro- and photochemical CO₂ reduction, mesoporous TiO₂ film, mesoporous TiO₂/iron porphyrin hybrid films, interfacial charge transfers, laser transient absorption spectroscopy

1. INTRODUCTION

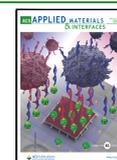
In the coming decades, enabling an infrastructure to produce commodity chemicals and fuels using renewable energy to fulfill global demand is a necessary challenge. Worldwide active research is going on to make such a novel industry in hard reality.^{1,2} Solar-driven water splitting and CO₂ reduction (CO₂R) are long-standing routes for the sustainable production of carbon-based fuels and feedstocks, as they can be easily utilized in existing fossil-fuel-assisted energy infrastructure.^{3–5} The ultimate inspiration to perform artificial water splitting and CO₂ reduction started with emulating natural photosynthesis. Nature performs the machinery in several concerted redox-favored processes, from light-triggered single-electron-transfer events to multielectron catalysis processes. Similarly, the artificial version of photosynthesis is envisaged by designing a hybrid assembly that integrates light absorption, electron transfer, and catalysis concurrently.^{6,7} For the CO₂R,

such an assembly can be incorporated as a photocathode, in which nanocrystalline semiconductor materials can be used to generate and store multiple charges for a considerable time with the accessible band gap across the solar spectrum and a molecular catalyst counterpart that can perform the multiredox electro- or photoelectrochemical CO₂ reduction with optimum activity and product selectivity.⁸ The quest for developing CO₂R photocathodes has been well documented in recent reviews covering the various combination of light-absorbing semiconductors (p-type Si, GaP, GaAs, n-type TiO₂, ZnO,

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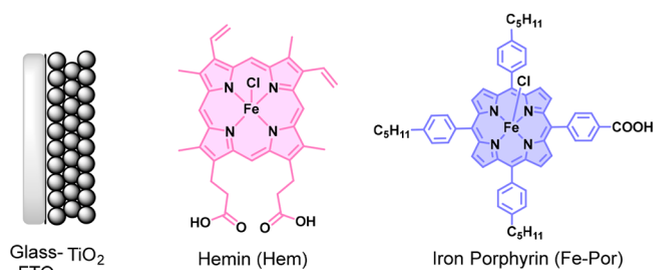
NiO, Cu₂O, etc.) and transition-metal-based molecular catalysts such as Re(vinyl-bpy)(CO)₃Cl, Ru(bpy-R₂)(CO)₂Cl₂ (R = PO₃H₂, COOH), Ni(cyclam)²⁺, Re(bpy)(CO)₃Cl, Fe-porphyrin, Co(II) bis(terpyridine), etc.^{8–10} For hybrid electrode fabrication, the desired covalent immobilization of the molecular catalyst over the semiconductor surface is obtained either by electropolymerization or via phosphonic, carboxylic, and hydroxamate anchoring groups.⁸ Different types of hybrid systems showed photocurrents in the range of 100–150 mA cm⁻² to 0.5–3 mA cm⁻² with moderate stability, having CO as the main CO₂R product along with CH₃COOH, HCOO⁻, and CH₃OH as minor products, although the formation of C₂₂ reduced products remains elusive.^{8,9} Many reports have explored different types of catalyst immobilizations, dissimilar anchoring methods, CO₂R product selectivity, and the overall stability of the photocathodes. However, the number of reports focusing on investigating the interfacial charge-transfer events occurring in the hybrid electrodes is low. This point is especially relevant due to the fact that poor charge accumulation and fast recombination of charges have a major impact on the multiredox reactions occurring at the molecular catalysts and thereby limit the overall photocathode CO₂R efficiency.¹¹ CO₂R involves multielectron and multi-proton transfer processes; therefore, by tuning the band gap and the conduction band energy of the semiconductor pertinent to the energy levels of the molecular catalysts provide scope to improve the interfacial electron transfers from the semiconductor to the molecular catalyst. The optimization of the energy levels of the photocathode components enhances the photoelectrochemical CO₂R efficiency and feasibility of seldom C₂₂ product formation.¹² Hence, the study of the interfacial charge transfer kinetics of such hybrid photocathodes has become of special interest. Durrant et al. have conducted important studies in this area. For instance, they explored the correlation between the photoelectrochemical hydrogen evolution and CO₂R activity with the lifetime of the charge-separated state formed in the hybrid systems made up of TiO₂-Co(II) terpyridine/cobaloxamine complexes via phosphonic acid linkages.^{13–15} Under light irradiation, charge accumulation, Co(II)/Co(I) redox shuttling, and transfer kinetics of the TiO₂ excited electron to the Co(II) catalysts showed a dependence on light intensity, applied external bias, catalyst loading, and the length of the linkers spanning the cobalt ion from the TiO₂ surface.

For our current study, we constructed a model photocathode system made of mesoporous TiO₂ and iron porphyrins as a molecular catalyst. We aimed at studying the interfacial charge transfer processes taking place during the CO₂R reaction. The choice of TiO₂ was guided by its advantageous properties, including transparency, stability, low cost, and its wide use in studies of dye-sensitized solar cells (DSSCs).^{10,16,17} For the CO₂R, bare nanocrystalline mesoporous TiO₂ is known to undergo multielectron-transfer processes to adsorb CO₂, resulting in the formation of CH₃OH and CO in acetonitrile. These electron-transfer processes were further examined by using spectroelectrochemical (ATR-IR and UV-vis) methods.^{18,19} On the other hand, iron porphyrin families with modified *meso*-aryl groups are well-known catalysts for the electrochemical transformation of CO₂ to CO with high Faradaic efficiency and overpotential values between 220 and 800 mV, depending on the solvent systems.²⁰ Furthermore, the catalytic activity of iron porphyrins was also explored in flow cell electrolyzers. In this case, the iron porphyrins were

employed as catalytic films obtained from their heterogenization onto nanocarbon support materials.²¹ Heterogenization of iron porphyrins on semiconductor supports has been previously reported; Naruta et al. studied CO₂ to CO conversion with an Fe *meso*-arylporphyrin immobilized on FTO-SnO₂ and FTO-TiO₂ surfaces by using phosphonic acid as an anchor group.²² Moore et al. studied the hydrogen evolution reaction (HER) after covalent grafting of Co and Fe *meso*-arylporphyrins on a GaP semiconductor.²³ More recently, the Aukauloo group investigated charge accumulation during the photocatalytic CO₂R mediated by an Fe porphyrin, [Ru(bpy)₃]²⁺ as photosensitizer, and DMF as solvent. These studies suggested the possibility of an alternate mechanism for the CO₂R involving a Fe(I) intermediate instead of the Fe(0) counterpart invoked in the classical pathway.^{24,25} Meyer et al. reported a non-Nernstian type multielectron transfer using mesoporous TiO₂-Co(II) *meso*-aryl porphyrin heterogeneous films. They performed nano- to microsecond transient absorption spectroscopy experiments upon excitation of the Co porphyrin at 532 nm. Nevertheless, they did not undertake any study of the catalysis.²⁶ To the best of our knowledge, studies of the intriguing interfacial charge transfer and charge recombination processes taking place on mesoporous TiO₂ and TiO₂/iron porphyrin films for the CO₂R under operando conditions have not been reported and we decided to focus on them in this work.

Herein, we report two model hybrid material-based catalysts for the CO₂R. The catalysts consist of a mesoporous nanocrystalline anatase polymorph TiO₂ functionalized with two different iron porphyrin-based catalysts. Specifically, from the iron porphyrin family, we choose (1) the commercially available hemin (Hem), a protoporphyrin IX ligand containing Fe(III) as a central metal atom having an axial chloride ligand, and (2) a A₃B type Fe(III) *meso*-tetraphenylporphyrin (Fe-Por) bearing three pentyl chains and one carboxylic acid group in the *para* positions of its *meso*-phenyl groups (Chart 1). We

Chart 1. (Left) Cartoon Representation of the Glass-FTO-TiO₂ Surface Used for the Covalent Anchoring of the Porphyrin Derivatives and (Right) Molecular Structures of the Iron Porphyrins (Hem and Fe-Por) Employed in the Preparation of the Porphyrin Hybrid Assemblies



report the synthesis, characterization, interfacial charge transfer kinetics, and catalytic performances in the CO₂R reaction of these two TiO₂/iron porphyrin hybrid assemblies. We also compare their performances with that of a mesoporous TiO₂ film used as a reference.

2. RESULTS AND DISCUSSION

2.1. Preparation and Characterization of the TiO₂/Iron Porphyrin (Hem and Fe-Por) Films. The free base 5-(4-carboxyphenyl)-10,15,20-tris(4-pentylphenyl)porphyrin has

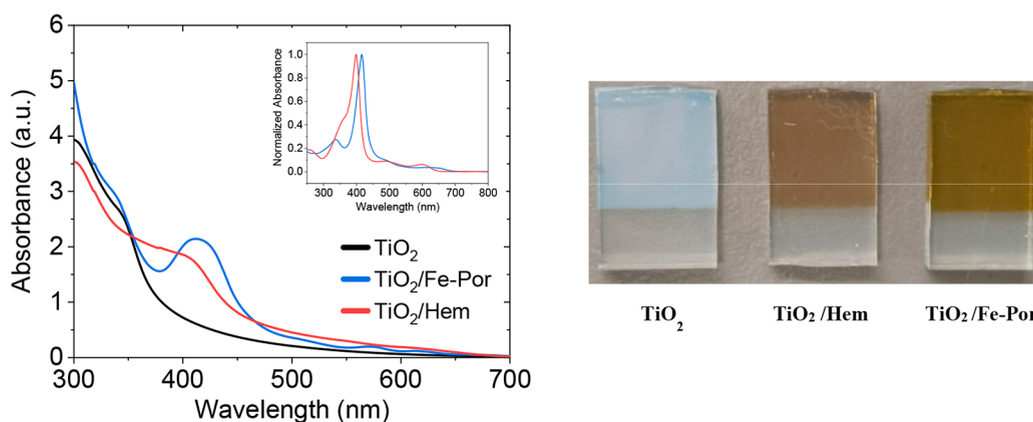


Figure 1. (left) UV–visible spectra of Hemin (**Hem**) and Fe-porphyrin (**Fe-Por**) on a transparent 5 μm thick mesoporous TiO_2 film measured in air with the normalized absorption spectra of only **Hem** and **Fe-Por** porphyrins in CH_3OH solution being shown in the inset. (right) Visual appearance of the TiO_2 , TiO_2/Hem , and $\text{TiO}_2/\text{Fe-Por}$ films on FTO glass.

been previously explored as sensitizer in TiO_2 solar cells,²⁷ in which the role of the *para*-pentyl chains in the *meso*-phenyl substituents was proved effective to deter porphyrin aggregation on the TiO_2 layer. The Fe metalation of the porphyrin core yielded **Fe-Por** (the detailed synthesis process is described in the [Supporting Information](#)), and its absorption characteristics and ^1H NMR shifts are in good agreement with those of similar iron porphyrins reported in the literature.^{28–30} We prepared separate solutions of **Hem** and **Fe-Por** in methanol by prolonged sonication of the initial suspensions. Subsequently, the Fe porphyrin derivatives were readily immobilized on the mesoporous TiO_2 surface using a dip-coating methodology. Most likely, the porphyrins were covalently anchored on the TiO_2 surface via the typical bidentate chelating mode of their carboxylic groups.³¹ The resulting films were stable at room temperature and kept in the dark until use ([Figure 1](#)).

The absorption maxima of **Hem** and **Fe-Por** in MeOH solution were observed at $\lambda = 400$ and $\lambda = 416$ nm, respectively, with broad Q-bands in the region of $\lambda = 500$ – 650 nm ([Figure 1](#), inset). However, upon anchoring the porphyrins to the TiO_2 surface, their absorption spectra experienced significant broadening, which can be attributed to the light-scattering effect of the mesoporous TiO_2 .^{32,33} The TiO_2/Hem and $\text{TiO}_2/\text{Fe-Por}$ hybrid assemblies showed broad Soret bands with maxima at $\lambda = 404$ and $\lambda = 414$ nm, respectively, and weakly absorbing Q-bands ([Figure 1](#)). Adsorption kinetics studies showed that the adsorption of both iron porphyrins on the TiO_2 surface reached a plateau within 20 min ([Figure S4](#) in the [Supporting Information](#)). Approximately, this should correspond to a monolayer of covalently anchored porphyrins on the TiO_2 surface. Thus, we conclude that despite the **Hem** and **Fe-Por** having structural differences (*meso*-aryl and β -substituents) and possessing different numbers of anchoring $-\text{COOH}$ groups, they experience similar adsorption kinetics onto the TiO_2 surface. The average loading of the iron porphyrins on the TiO_2 was estimated from the UV–visible measurements of solutions obtained after leaching treatments of the $\text{TiO}_2/\text{Fe-Por}$ and TiO_2/Hem under a high reductive and mild oxidative potential (cycles between -1.8 and 0.8 V vs Ag/AgCl). This result also evidenced the low stability of the linkages between the porphyrin carboxylic group and the TiO_2 surface under the above electrochemical conditions. We estimated that the average loading of the iron porphyrins on

the TiO_2/Hem and $\text{TiO}_2/\text{Fe-Por}$ films are 11 ± 3 and 22 ± 3 nmol cm^{-2} , respectively ([Figure S4](#)). The variation in loading arises from the different structures of the Fe porphyrins, but both values are comparable with those previously reported for cobalt porphyrin loadings on mesoporous TiO_2 .²⁶ However, they are smaller than those of cobalt complexes bearing phosphonic acid anchors.¹⁴ Attenuated total reflection infrared (ATR-IR) spectroscopy was also used for the characterization of TiO_2/Hem and $\text{TiO}_2/\text{Fe-Por}$ hybrid assemblies (spectra are presented in [Figures S5](#) and [S6](#)). In **Fe-Por**, the Fe–N bond stretching frequency was centered at ~ 1000 cm^{-1} , those of the C–H bonds of benzene and pyrrole appeared at ~ 2923 – 2852 cm^{-1} , the C=C stretching mode at ~ 1494 – 1687 cm^{-1} , and that of the C=N at ~ 1333 cm^{-1} . Also, bands at 1733 and 1272 cm^{-1} are assigned to the characteristic C=O and C–O stretching of the carboxylic acid group, respectively. The values follow the same trend as reported for other iron(III) chloride porphyrin complexes.^{34,35} The broad peaks at ~ 3400 and ~ 1650 cm^{-1} for the FTO- TiO_2 film correspond to the stretching vibration of hydroxyl functional groups and bending vibration of free water, respectively.³⁶ The spectrum of $\text{TiO}_2/\text{Fe-Por}$ showed the near disappearance of C=O and C–O stretching modes of the free carboxylic acid group, confirming the anchoring of **Fe-Por** on the TiO_2 surface. However, in the $\text{TiO}_2/\text{Fe-Por}$ spectrum, broadening of the **Fe-Por** signals is attributed to the presence of a nanomolar amount of porphyrin loading compared to a high excess of TiO_2 molecules (3.91 $\mu\text{mol cm}^{-2}$ of TiO_2). The IR spectrum of TiO_2/Hem exhibited similar characteristics feature ([Figure S6](#)). XPS analysis was further carried out with the TiO_2 , TiO_2/Hem , and $\text{TiO}_2/\text{Fe-Por}$ films ([Figures S7](#) and [S8](#)). In TiO_2 , the Ti $2p_{1/2}$ and Ti $2p_{3/2}$ peaks appeared at 464.4 and 458.7 eV, respectively, which matches the literature report.³⁷ There is no marked shift and no shape change in the Ti $2p_{1/2}$ and Ti $2p_{3/2}$ signals when comparing bare TiO_2 to $\text{TiO}_2/\text{iron porphyrins}$ films, which confirms that there is no change in the coordination and electronic environment of Ti in TiO_2 after the porphyrin loading. Furthermore, [Figure S8](#) reveals the Fe 2p signals at ~ 709 – 711 eV (Fe $2p_{3/2}$) and ~ 723 eV (Fe $2p_{1/2}$) in TiO_2/Hem and $\text{TiO}_2/\text{Fe-Por}$ films, respectively, which is in accordance with the +III oxidation state of iron.³⁸ Thus, an XPS analysis confirms the presence and anchoring of the iron(III) porphyrin structures in the hybrid catalyst films.

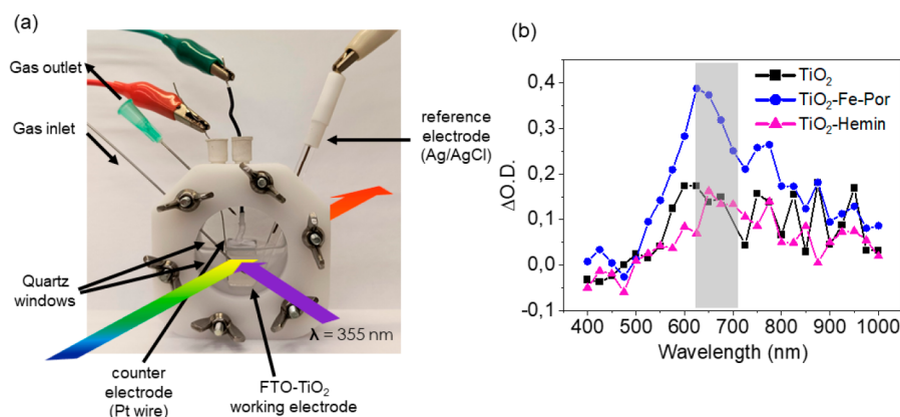


Figure 2. (a) Picture of the Teflon L-TAS cell setup with double quartz windows, equipped for measurements under a controlled atmosphere and applied external bias. (b) Survey L-TAS spectra of TiO_2 , TiO_2/Hem , and $\text{TiO}_2/\text{Fe-Por}$ films measured in acetonitrile solution containing 0.1 M TBAPF₆ after $\lambda = 355$ nm excitation under an N_2 atmosphere. The probe wavelength was fixed at $\lambda = 680$ nm.

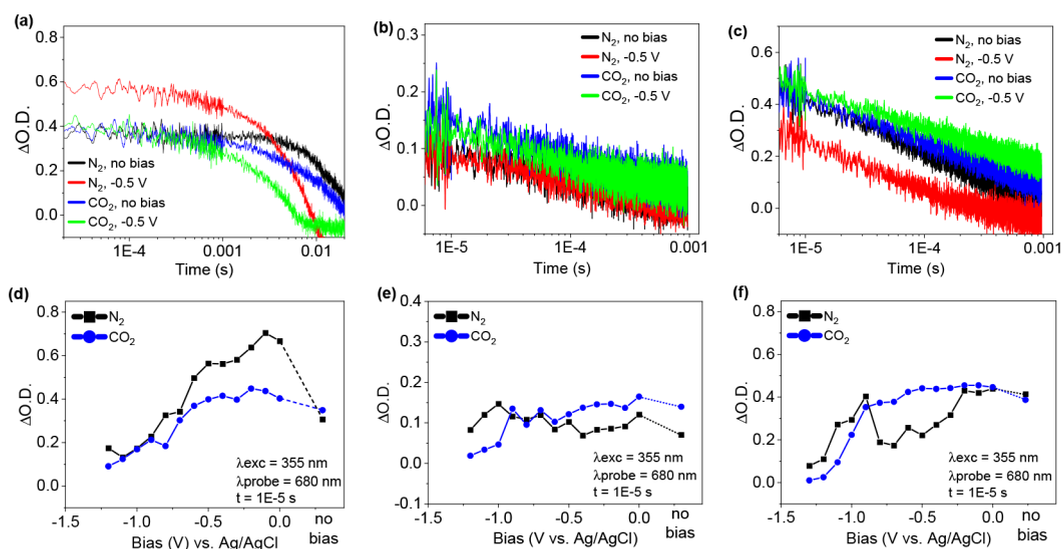


Figure 3. TAS spectra of (a) TiO_2 , (b) TiO_2/Hem , and (c) $\text{TiO}_2/\text{Fe-Por}$ recorded in the presence of acetonitrile containing 0.1 M TBAPF₆ after excitation at $\lambda = 355$ nm and under N_2 and CO_2 atmospheres using applied bias (-0.5 V vs Ag/AgCl) conditions. Only the L-TAS of TiO_2 was measured in the presence of 0.01 M TEOA as a hole scavenger. (d–f) Comparison of the transient absorption for TiO_2 , TiO_2/Hem , and $\text{TiO}_2/\text{Fe-Por}$, respectively, acquired under N_2 and CO_2 atmospheres. The TAS spectra were recorded by applying a bias of 0 to -1.2 V vs Ag/AgCl. The transient absorption values were taken at the time 1×10^{-5} s. The probe wavelength was fixed at $\lambda = 680$ nm.

2.2. Interfacial Charge Transfer Kinetics. Laser transient absorption spectroscopy (L-TAS) was employed to investigate the charge transfer and recombination kinetics between the components of the TiO_2 /iron porphyrin hybrid assemblies by monitoring the charge carrier lifetimes (Figure 2a). For our current study, we focused on micro- to millisecond charge carrier lifetimes of TiO_2 and their corresponding values pertinent to the CO_2R reaction. Transient decays of the anatase TiO_2 photoelectron in an acetonitrile-TBAPF₆ (supporting electrolyte) solution under an N_2/CO_2 atmosphere were measured following a band gap excitation at $\lambda = 355$ nm. After laser excitation at $\lambda = 355$ nm in the dark of the TiO_2 film, the survey spectrum (black trace in Figure 2b) showed transient absorption throughout the $\lambda = 600$ – 900 nm range, where the electrons in the conduction band of TiO_2 absorb. We chose to study and compare all our results with the $\lambda = 680$ nm transient absorption signals and obscure the $\lambda = 690$ – 710 nm region signals to avoid any interference in the signals arising from the second harmonics of the excitation wavelength ($\lambda_{\text{exc}} = 355$ nm; $\lambda_{\text{har}} = 710$ nm).

Previously, it has been also shown that the use of a chemical scavenger during the L-TAS measurement can specifically quench either the electrons (AgNO_3) or the holes (methanol or amine) depending on the choice.^{14,39} In the present case, we were particularly interested in the photogenerated electrons and their recombination kinetics during the CO_2R reaction. Hence, we used a sacrificial hole scavenger (triethanolamine, TEOA) to quench the TiO_2 photogenerated holes during the measurement. The use of TEOA as a hole scavenger increased the lifetime of the photoexcited electrons and induced an increment in the amplitude of the transient absorption signal (Figure S9a). Thus, in the presence of 0.01 M TEOA (black trace in Figure 3a), we corroborated that the photoexcited electrons in TiO_2 exhibit decay kinetics on the millisecond time scale following typical stretched exponential kinetics.^{40–42}

Next, to mimic a typical photoelectrochemical CO_2 reduction experiment, we purged CO_2 into the electrolyte and/or we applied a bias varying from 0 to -1.3 V vs Ag/AgCl to the TiO_2 film. In doing so, we fixed the Fermi level at the TiO_2 electrode. Ideally, this bias range is less than the TiO_2

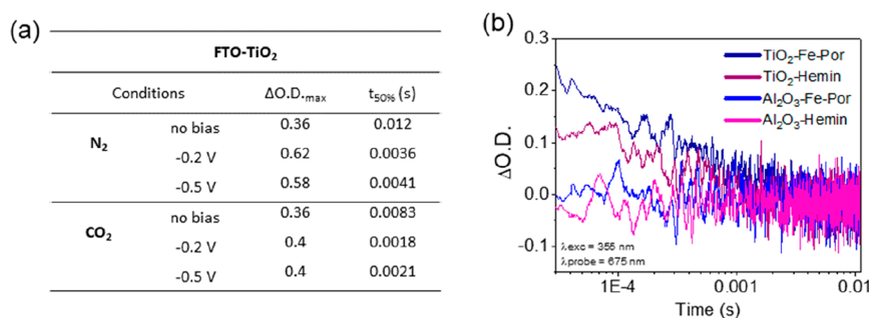


Figure 4. (a) Table of the $t_{50\%}$ values obtained with the TAS measurements on TiO₂ films. (b) Comparison of TAS spectra of Al₂O₃/Hem and Al₂O₃/Fe-Por with those of TiO₂/Hem and TiO₂/Fe-Por in acetonitrile containing 0.1 M TBAPF₆ after excitation at $\lambda = 355$ nm under an N₂ atmosphere and without an applied bias voltage. The probe wavelength was fixed at $\lambda = 675$ nm.

conduction band potential, so that we restrict the continuum filling of the valence band to the conduction band level of TiO₂. However, we observed differences in the decay kinetics of the photogenerated TiO₂ electron when CO₂ was purged in the electrolyte even in the absence of applied potential. In fact, under a CO₂ atmosphere, the transient signal intensity remains at the same ΔOD initial value but its lifetime is shortened in comparison to an N₂ atmosphere (blue trace in Figure 3a). The corresponding " $t_{50\%}$ " values (time taken to reach half of the transient absorption amplitude values, ΔOD) and maximum ΔOD at $\lambda = 680$ nm are compared in Figure 4a; $t_{50\%}$ decreases from 12 ms under an N₂ atmosphere to 8 ms under a CO₂ atmosphere. Afterward, we chose in particular a bias of -0.5 V (vs Ag/AgCl) to compare the transient decay with the nonbiased case under N₂ and CO₂ atmospheres (red and green traces in Figure 3a). The overlay of all the transient decays recorded in the range of 0 to -1.3 V vs Ag/AgCl applied bias under N₂ and CO₂ atmospheres for the TiO₂ film is shown in Figure S10a,b. Under N₂ and -0.5 V applied bias (vs Ag/AgCl) the transient signal intensity of the TiO₂ film increases while the $t_{50\%}$ value decreases to 4 ms in comparison to a no-bias experiment (red trace in Figure 3a). Most likely, this is due to the fact that the tail states below the TiO₂ conduction, which are likely involved in the trapping of the photogenerated electrons, are filled up electrochemically.

Under CO₂, an application of the external bias produces a marked difference from the transient decay for the N₂ atmospheres. With CO₂, the maximum transient absorption decreases by 0.2 ΔOD , and the lifetime of photogenerated electrons features a reduced $t_{50\%}$ value of 2 ms (green trace in Figure 3a). When bias is applied, in the 0 to -1 V (vs Ag/AgCl) region, the maximum absorption values, measured at time 1×10^{-5} s, also show noticeable differences between N₂ and CO₂ atmospheres (Figure 3d). The absorption value under a CO₂ atmosphere always remains lower than that of the experiment under N₂. For both cases, after applying -1 V vs Ag/AgCl, the transient signal starts to disappear due to electrochemical filling up of the conduction band of TiO₂ with electrons.

We conducted the same measurements for TiO₂/Hem and TiO₂/Fe-Por films, and we observed some differences in the transient absorption signals compared to the bare TiO₂ film (Figure 3b,c). First, the use of TEOA in the case of the TiO₂/Fe-Por and TiO₂/Hem films did not have any improvement effects in the transient absorption or in the decay kinetics (Figure S9b,c). This observation is remarkably different from that observed previously for the bare TiO₂, suggesting that TEOA does not stabilize the oxidized porphyrin catalyst. For

this reason, we skipped the use of TEOA in the TAS measurements of TiO₂/Hem and TiO₂/Fe-Por films. Second, the presence of Fe-Por on the TiO₂ caused an increase of the transient signal in the region of $\lambda = 550$ – 800 nm, with a maximum at $\lambda = 650$ nm (blue trace in Figure 2b). In the case of Hem a related but less defined transient absorption spectrum was observed (pink trace in Figure 2b). The recorded signal in both spectra correspond to the oxidized porphyrin catalysts as a result of the photoinduced electron injection to the TiO₂ conduction band. Third, under identical conditions, we observed that for the TiO₂/Hem and TiO₂/Fe-Por films their transient decays are 100 times faster than that of the bare TiO₂ film. The transient signal of the porphyrin-functionalized films disappeared by 1 ms (Figure 3b,c). Last but not least, the initial steady state (time scale faster than 1 μ s) is not clearly recordable within our time resolution range. This is particularly important for TiO₂/Hem, with ΔOD as low as 0.1 ΔOD . This limitation does not allow us to determine accurate initial transient absorption and $t_{50\%}$ values for the TiO₂/Fe-Por and TiO₂/Hem hybrid systems. Moreover, these findings indicate the existence of faster charge recombination processes in the hybrid systems compared to the bare TiO₂ film.^{26,43,44}

By looking in more detail at the effect of the presence of CO₂ and the application of bias to the TiO₂/Hem and TiO₂/Fe-Por films, we observed a different behavior of the hybrid films in comparison to bare TiO₂. Under an N₂ atmosphere, the application of -0.5 V caused a reduction in the transient signal (red trace in Figure 3b,c). The reduction of the intensity was more significant for the TiO₂/Fe-Por film than for the TiO₂/Hem counterpart. The presence of CO₂ does not pronouncedly affect the L-TAS signal in the nonbiased case but, interestingly, it prevents the decrease in the transient signal when a -0.5 V bias is applied (blue and green traces in Figure 3b,c). Overlays of the transient absorption decays recorded over the range of 0 to -1.3 V vs Ag/AgCl applied bias under N₂ and CO₂ atmospheres are shown in Figure S10c–f. In the 0 to -1 V vs Ag/AgCl range, the maximum transient absorption values obtained at time 1×10^{-5} s showed that TiO₂/Fe-Por and TiO₂/Hem follow similar trends (Figure 3e,f) with always smaller ΔOD amplitude transient signals for TiO₂/Hem. For the TiO₂/Fe-Por film under an N₂ atmosphere, the maximum of the transient signal decreases with negative bias until -0.8 V vs Ag/AgCl is applied. In contrast, under a CO₂ atmosphere, the application of a negative bias does not have a significant effect on the maximum of the transient signal until a bias of -1 V vs Ag/AgCl is applied. Under N₂ and CO₂ atmospheres, the

Table 1. CO₂R Performances of the TiO₂ Film and TiO₂/Hem and TiO₂/Fe-Por Hybrid Films under Electro-, Photo-, and Photoelectrochemical Conditions^a

| electrode | potential (vs Ag/AgCl) | without light irradiation | | | with 365 nm LED irradiation | | |
|------------------------------|------------------------|---------------------------|----------------------------------------------------------|------------------------------|-----------------------------|----------------------------------------------------------|------------------------------|
| | | product | production rate (nmol h ⁻¹ cm ⁻²) | selectivity (%) ^b | product | production rate (nmol h ⁻¹ cm ⁻²) | selectivity (%) ^b |
| FTO-TiO ₂ | no bias | | | | CO | 22 ± 2 | 100 |
| | | | | | CH ₄ | traces | |
| | -0.5 | | | | CO | 19 ± 3 | 56 |
| | | | | | CH ₄ | 15 ± 2 | 44 |
| | -1.25 | | | | CO | 51 ± 10 | 51 |
| | | | | | CH ₄ | 48 ± 10 | 49 |
| | -1.5 | H ₂ | 140 | 100 | H ₂ | 160 ± 9 | 70 |
| | | | | CO | 67 ± 2 | 30 | |
| | -1.8 | H ₂ | 496 ± 118 | 85 | | | |
| | | CO | 90 ± 51 | 15 | | | |
| FTO-TiO ₂ /Hem | no bias | | | | CO | 17 ± 2 | 100 |
| | -0.5 | | | | CO | 19 ± 8 | 100 |
| | -1.25 | | | | CO | 29 ± 3 | 100 |
| | -1.5 | CO | 280 ± 60 | 100 | CO | 211 ± 1 | 100 |
| | -1.8 | | | | leaching of porphyrin | | |
| FTO-TiO ₂ /Fe-Por | no bias | | | | CO | 25 | 100 |
| | -0.5 | | | | CO | 40 ± 10 | 100 |
| | -1 | | | | leaching of porphyrin | | |

^aMeasurements were done in a single-compartment cell using Pt wire as the counter electrode, Ag/AgCl as the reference electrode, and acetonitrile solution containing 0.1 M TBAPF₆ as the supporting electrolyte. ^bSelectivity = mol of one product/mol of total product formed.

application of potentials more negative than -1 V induced the disappearance of the transient signals. This is due to the existence of very fast charge recombination kinetics between the electrons at the TiO₂ conduction band and the oxidized catalyst.

As a control system, we prepared an iron porphyrin supported on Al₂O₃ nanocrystalline mesoporous films. We chose Al₂O₃ since the energy values of the conduction band (~-1.5 eV vs vacuum) of this metal oxide (high compared to TiO₂ ~-4.5 eV vs vacuum) prevents efficient photoinduced electron transfer from the porphyrin's excited state upon λ = 355 nm excitation.⁴⁵ Figure 4b presents a comparison of the TAS signals of the hybrid assemblies with immobilized iron porphyrin on TiO₂ and of Al₂O₃ analogues. As expected, no transient absorption signals are observed in case of Al₂O₃/Hem and Al₂O₃/Fe-Por hybrid assemblies. Furthermore, we also performed L-TAS by exciting the TiO₂/iron porphyrin assemblies at the Soret and first Q-band of the Fe porphyrin (excited with λ = 420 nm and λ = 500 nm for TiO₂/Hem or λ = 530 nm for TiO₂/Fe-Por, laser). We monitored the transient absorption signals associated with the excited porphyrins. Notably, we did not observe any transient signals using TiO₂/Hem at the millisecond time scale (Figure S11b) at both excitation wavelengths. Conversely, for the TiO₂/Fe-Por hybrid, the excitation at 420 and 530 nm produced a weak transient signal at 650 nm (Figure S11a) in the millisecond time scale. The nanocrystalline mesoporous TiO₂ film does not have a significant absorption at 420 and 530 nm. Thus, the transient decay must be due to the photoinjection of electrons in the TiO₂ conduction band from the photoexcited iron porphyrin. This result is in good agreement with our previous measurements depicted in Figure 2b for the TiO₂/Fe-Por film. In short, the results of the control studies support that, for the TiO₂/Hem and TiO₂/Fe-Por sensitized films, the transient

decays correspond to the photoinduced charge recombination kinetics between the electrons at the TiO₂ conduction band and the oxidized porphyrins.

We became interested in studying the electron transfer kinetics of these hybrid systems for the CO₂R under operando conditions. Thereby, we studied the fate and efficiency of the hybrid systems during the CO₂R using similar L-TAS conditions.

2.3. CO₂ Reduction Studies using TiO₂, TiO₂/Hem, and TiO₂/Fe-Por Films. To understand the CO₂R performances using the TiO₂/Hem and TiO₂/Fe-Por films, we investigated the process in a one-compartment glass cell under both electro- and photoelectrochemical conditions. The cyclic voltammograms (CV) of Hem and Fe-Por in homogeneous DMF solution showed three typical iron-center-based reduction peaks (Fe^{III/II}, Fe^{II/I}, Fe^{I/0}) (Hem, -0.35, -1.21, -1.81 V; Fe-Por, -0.24, -1.06, -1.67 vs Ag/AgCl). The CV experiments were carried out under an argon atmosphere and using ferrocene as an internal standard (Figure S12a). Further, in the presence of CO₂, we observe a catalytic current response at the Fe^{I/0} wave in both cases (Figure S12b,c). In a DMF solution, the controlled-potential electrolysis (at -1.6 V vs Ag/AgCl for Fe-Por or -1.7 V vs Ag/AgCl for Hem) produced exclusively CO as the CO₂R product for both iron porphyrins. The CV of the bare TiO₂ film in acetonitrile solution under an argon atmosphere exhibited the appearance of a signal at -1.3 V vs Ag/AgCl. This value matched well with the reported reduction of Ti^{IV} to Ti^{III} at the TiO₂ surface (Figure S13a).¹⁸ In the presence of CO₂, an increase in the reduction current is noticed to have an onset at the same -1.3 V potential. It is important to note that the -1.3 V vs Ag/AgCl potential is greater than the TiO₂ conduction band edge and at this potential accumulation of electron-rich Ti³⁺ species at the surface of the film is initiated. These species are key for the

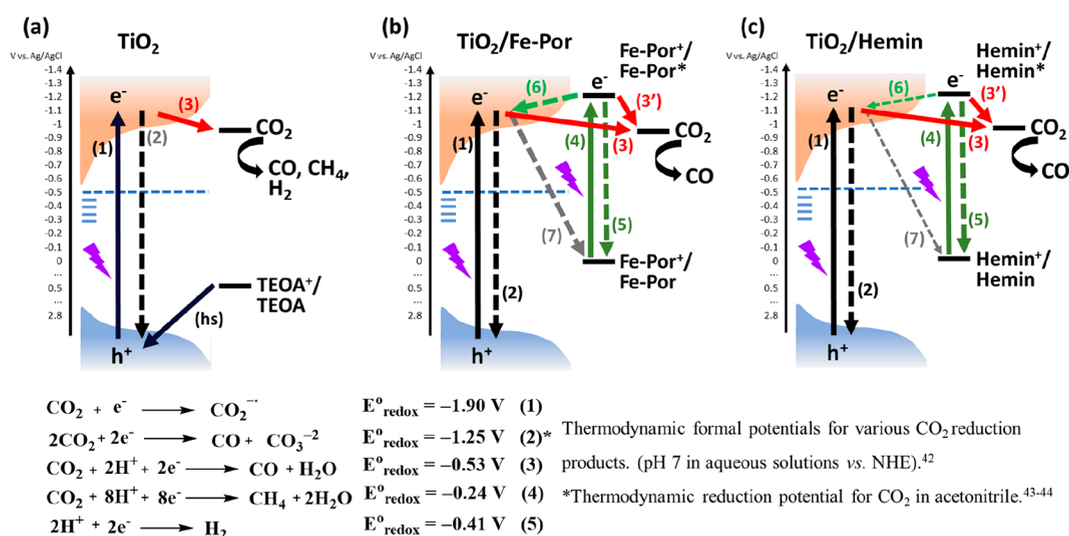


Figure 5. Suggested pathways for the different electron transfer processes occurring in (a) TiO₂, (b) TiO₂/Fe-Por, and (c) TiO₂/Hem after laser excitation at $\lambda = 355 \text{ nm}$ and application of an external bias. “hs” denotes scavenging of holes by TEOA as a hole scavenger. (1) and (4) denote light excitation of TiO₂ and iron porphyrins, (2) and (5) denote the back-electron-transfer processes between the TiO₂ (CB) to TiO₂ (VB) and Fe-Por* to Fe-Por, (3) denotes the charge/electron transfer from TiO₂ (CB) to an adsorbed CO₂ molecule, (3') denotes the charge transfer from Fe-Por* to an adsorbed CO₂ molecule, and (6) and (7) represent the back-electron-transfer processes from the reduced Fe-Por* to TiO₂ (CB) and TiO₂ (CB) to the Fe-Por ground state or to the electrolyte. At the bottom, CO₂ reduction potentials are given for different product formations in aqueous and acetonitrile solvents.

commencement of the CO₂ reduction catalysis. In the oxidative scan, the peak observed at -1.5 V vs Ag/AgCl is ascribed to the oxidation of some Ti³⁺ to Ti⁴⁺ state (Figure S13b,c). The CVs of TiO₂/Hem and TiO₂/Fe-Por film electrodes under an argon atmosphere did not display the diagnostic redox couples for three iron-centered reversible reactions. In TiO₂/Hem, a reduced signal at -1.47 V and in the oxidative scan a peak at -1.40 V were noted (Figure S13a). For TiO₂/Fe-Por a reversible wave at -1.12 V vs Ag/AgCl was observed. However, in the presence of CO₂, both TiO₂/Hem and TiO₂/Fe-Por films showed a catalytic reduction current starting at an onset potential of $\sim -1.3 \text{ V}$. The onset potential of the catalytic currents of the hybrid films is also similar to that of the bare TiO₂ film (Figure S13b,c).

The electro-, photo-, and photoelectrochemical CO₂R product selectivities of the hybrid assemblies were further evaluated using controlled-potential electrolysis, at potentials ranging from -0.5 to -1.8 V vs Ag/AgCl of the CO₂-saturated acetonitrile solutions. The evolved gases were analyzed using gas chromatography, and no liquid products were detected in solution. CO and CH₄ were obtained as CO₂ reduction products and H₂ was also formed in some cases (Figures S14 and S15). The obtained results are summarized in Table 1. Under electrochemical conditions, the TiO₂ film produced a small amount of CO only when the applied potential reached a value of -1.8 V vs Ag/AgCl, although under those conditions, H₂ was the main product (15% CO selectivity). In contrast, the simple irradiation of the TiO₂ film with a 365 nm LED light produced the formation of CO and a trace of CH₄. The application of a bias potential of -0.5 V to the light-irradiated electrode did not modify the production of CO but initiated significant formation of CH₄ ($\sim 50\%$ selectivity). At -1.25 V , the production rates of the two gases increased, maintaining a similar product selectivity. However, at -1.5 V , H₂ evolved as the major product. CO was also produced ($\sim 30\%$ selectivity), but no CH₄ was detected. Taken together, these observations support that the photoelectrons injected in the conduction

band of TiO₂ are responsible for the photoelectrochemically catalyzed CO₂R occurring at lower overpotentials (-0.5 to -1.5 V) than that required under pure electrochemical conditions (-1.8 V). Also, under light irradiation, it is possible to obtain CH₄, the 8-electron-reduction product of CO₂, along with suppression of the undesired HER. Previously, it was described that Ti³⁺ sites could enhance the chemisorption of CO₂ and entail its activation to undergo C–O bond cleavage.⁴⁶ Also, the photogenerated electrons accumulated in the TiO₂ intraband states were claimed to facilitate multielectron-transfer processes, generating C₂ and C₃ hydrocarbons.⁴⁷ Nevertheless, we have not observed the presence of C₂ and C₃ products in the present study.

Both TiO₂/Hem and TiO₂/Fe-Por hybrid films exclusively formed CO as the CO₂R product using either electro- or photoelectrochemical conditions. Under electrochemical conditions, the TiO₂/Hem film started to furnish CO starting at -1.5 V , but under light irradiation the TiO₂/Hem film started formation of CO starting at -0.5 V . However, at -0.5 and -1.25 V , the rate of formation of CO became similar to that of bare TiO₂ (Table 1). Formation of CO under light irradiation increased drastically at -1.5 V . For the TiO₂/Fe-Por film, only light irradiation furnishes CO formation similar to that observed in previous TiO₂ and TiO₂/Hem cases. A slight increase in the CO formation catalyzed by the TiO₂/Fe-Por film was observed under the photoelectrochemical conditions at -0.5 V applied bias (Table 1). As indicated, the TiO₂/Fe-Por hybrid system was unstable in controlled-potential electrolysis at further potentials more negative than -0.8 V ; thus, we were restricted to further explore its activity for the CO₂R.

It is worth noting that we did not detect the formation of CO₂R products under pure electrochemical conditions (without light irradiation) using the TiO₂-sensitized/hybrid films in the range of applied bias (0 to -1 V vs Ag/AgCl) used for the L-TAS measurements. The studied TiO₂ hybrid films show a reduced performance in the CO₂R reaction compared to

reported analogous hybrid assemblies of FTO and FTO-SnO₂-supported Fe porphyrin dimer.²² Nevertheless, it is remarkable that the TiO₂-sensitized films reported here (TiO₂/Hem and TiO₂/Fe-Por) can be used as highly selective (100% CO selectivity) photocathodes. Also, the superior performance of the materials for the formation of CO₂R products, under the photoelectrochemical conditions at much more positive reduction potentials, indicates that the photogenerated electrons within the sensitized TiO₂ hybrid films play a pivotal role in the catalysis.

3. DISCUSSION

In this work, we report our results of CO₂ reduction catalysis using nanocrystalline mesoporous TiO₂ and TiO₂/iron porphyrin hybrid films. The reported findings are supported by the characterization of the decay dynamics of the photoinduced charge transfer processes occurring in the films using micro- to millisecond L-TAS spectroscopy. Previously, it was reported that the conduction band electrons of TiO₂ can be trapped in the intraband defect states of the surface and the bulk. The electron entrapment strongly influences the charge transport properties of the TiO₂ film.⁴⁸ In addition, the application of a negative potential results in the accumulation of electrons in the conduction and intraband trap states. The accumulation of the electrons produced a modulation of the Fermi level of the TiO₂ electrode. The rate of charge recombination increases as the applied negative voltage does.⁴⁹ In fact, we observed that in the range of 0–0.8 V (vs Ag/AgCl) applied voltage, the nanocrystalline mesoporous TiO₂ displayed a maximum ΔOD transient absorption in comparison to an analogous experiment without any applied voltage bias (Figure 3a,d). This is due to the continuum filling of the electrons in the valence band to the Fermi level of the TiO₂ electrode. The $t_{50\%}$ value (0.0041 s) at –0.5 V applied bias is smaller than that measured without applying a bias voltage (0.012 s). This result demonstrated that the application of a bias voltage caused a significant increase in the rate of charge recombination (65% reduction in $t_{50\%}$ value) (Figure 4a). In Figure 5a, we denoted the charge recombination process (2), occurring in the millisecond to second time scale, as that observed in the transient absorption decay process of the FTO-TiO₂ film. In the presence of CO₂ and under a –0.5 V bias, a decrease in both the transient absorption (~35%) and $t_{50\%}$ values (50%) are noted (Figure 4a). This indicates that the photogenerated electrons at the TiO₂ surface are rapidly transferred (faster than 10 ms) to the adsorbed CO₂ molecule at the surface and further reduction (process (3) in Figure 5a and Figure 3a) to C₁ products. Also, the presence of the adsorbed CO₂ molecules causes (process (3)) faster charge recombination of the TiO₂ excited electrons. Subsequently, we tested our hypothesis by performing controlled-potential electrolysis using FTO-TiO₂ films as electrodes under similar conditions employed for the TAS experiments. Our aim was to detect the formation of any multielectron-assisted CO₂R product triggered by the accumulated photogenerated TiO₂ electrons. Table 1 shows that the photogenerated TiO₂ electrons induced the formation of both CO (2e[–]) and CH₄ (8e[–]) products in either the presence or absence of a voltage bias. The use of the TiO₂ electrode under typical electrochemical conditions required an applied voltage of at least –1.8 V to induce the formation of products and furnish CO and H₂.

It worth noting that, based on the standard potential values for the reduction reactions of CO₂ to CO and CH₄ in H₂O and acetonitrile solution,^{50–53} there is a gain in the overpotential value when the FTO-TiO₂ electrode is used under photoelectrochemical conditions. This result correlates directly with the faster charge recombination kinetics measured in the corresponding TAS experiments. To the best of our knowledge, this is the first report studying the charge transfer kinetics of mesoporous TiO₂ films during CO₂ reduction, monitoring the micro-/millisecond lifetimes of the photoexcited electrons of TiO₂. In the L-TAS measurement, TEOA was used in excess as a hole scavenger. However, TEOA can also bind at the TiO₂ surface, as well as induce the formation of a zwitterionic alkylcarbonate adduct⁵⁴ with CO₂ that also interacts with the surface via unreacted terminal –OH groups. The two surface binding processes involving TEOA may produce a faster decay in the L-TAS experiments, as observed for the TiO₂ electrodes. For our electrolysis studies, we did not use TEOA. Nevertheless, multiple reports have described the use of TEOA as a sacrificial electron donor for the catalytic CO₂ reduction in organic media.^{54–56} The presence of TEOA can alter the reaction rate and the product selectivity of the CO₂ reduction, but it does not inhibit the reaction.

For the TiO₂/Hem and TiO₂/Fe-Por hybrid films the situation changed. In an ideal photocathode, electron transfer from the TiO₂ conduction band to the iron porphyrins is desired. This process should cause the reduction of the iron metal center and subsequent CO₂ activation/reaction. However, both TiO₂ and iron porphyrins absorb light in the visible region. The light absorption initiates intra- and interfacial electron communications within the TiO₂/Iron porphyrin hybrid systems. The standard potential of the TiO₂ conduction band (–0.85 V vs NHE) is well below the reduction potential of the Fe^{II/I} and Fe^{I/0} processes of iron porphyrins (Hem, –1.03, –1.61 V; Fe-Por, –0.86, –1.47 V vs NHE). Fe^I and Fe⁰ are potent reductants to activate the reduction of CO₂. Considering that in the hybrids film the energy levels of the different redox states are similar to those in acetonitrile solution, the electron transfer from the photoexcited Hem and Fe-Por units to the conduction band of TiO₂ is feasible and thermodynamically favored.⁵⁷ In fact, the micro- to millisecond L-TAS spectra of TiO₂/Hem and TiO₂/Fe-Por imply the existence of such an interfacial charge transfer process. The excitation with 355 nm light initiates ultrafast electron injection in the excited state of the iron porphyrins from the ground singlet states, which can be recombined back (processes (4) and (5) in Figure 5b,c), producing a transient signal. However, the rate of this recombination step (5) for iron porphyrins is known to occur in the pico- to femtosecond time scale.^{43,44,58–60} Consequently, it is not possible to detect this process in the presented data set due to the time resolution limitation of the used equipment. Two additional interfacial charge transfer processes can be foreseen: process (6), corresponding to the photoinjection of electrons in the TiO₂ conduction band/intraband states from the excited state of the iron porphyrins, and process (7) corresponding to the back electron transfer from the TiO₂ conduction band to the oxidized porphyrin ground state (Figure 5b,c). From the spectra shown in Figure 3b,c, it is evident that from 10 μ s to 1 ms similar decay signals are observed for the both iron porphyrins, which are quite different from that measured for the reference TiO₂ film, recorded at the same time. Thereby, considering Figure 5b,c, we suggest that the decay of the

TiO₂/iron porphyrin film signals does not correspond to the recombination process (2). The decay of the signal is fast (decay in 1 ms) compared to that of electrons in the conduction band of TiO₂ (up to 0.1 s) (Figures 3a and 5a). Comparing the electron transfer processes (6) and (7), it is more likely that process (6) (nano- to microsecond) exhibits faster relaxation dynamics than process (7) (micro- to millisecond). Hence, the transient signals in Figure 3b,c featuring decays in the 10 ms scale are attributed to the electron-transfer process (7). In this process, the oxidized iron porphyrin cation is being reduced by the back-electron transfer from electrons accumulated or photoinjected in the TiO₂ conduction band/intraband states. The relative position of the oxidized porphyrin cation makes process (7) energetically more favorable than process (2), leading to ~100 times faster decay of the L-TAS signals in the hybrid materials. In the presence of CO₂, the photoexcited electrons can be transferred to the adsorbed CO₂ molecule on the surface. This is depicted via processes (3) (from the conduction band of TiO₂) and (3') (from excited state of the iron porphyrin). Electron-transfer processes (4), (6), and (7) take place efficiently in a closed-loop manner within the covalently attached iron porphyrins and the TiO₂ interface. This is detrimental to the direct transfer of photogenerated electrons to adsorbed CO₂ molecules (processes (3) and (3')). This hypothesis could serve to explain our observations related to not having significant changes in the decays of the transient absorption signals for TiO₂/Hem and TiO₂/Fe-Por hybrid films in the presence or absence of CO₂ (Figure 3b,c). The presence of TEOA as a hole scavenger has no effect on the stabilization of the oxidized iron porphyrin cation state (Figure 5b,c and Figure S9b,c). Therefore, and in agreement with experiment, the decay of the signals corresponding to process (7) are not modified. In support of these L-TAS observations, photoelectrocatalysis results confirm that both TiO₂/iron porphyrin systems form CO as the reduced product at -0.5 V with 100% selectivity. As shown in Table 1, under photoelectrocatalysis conditions with an applied voltage bias of -0.5 V, the rates of product formation for bare TiO₂ and for TiO₂/iron porphyrin films are similar; however, the TiO₂ film also produces CH₄. We conclude that the presence of the Fe porphyrins causes a change in product selectivity in favor of CO formation. Applying a more negative potential bias (≥ -1 V) favors the reduction of the iron center in the porphyrin layer, resulting in a better CO₂ reduction performance. As explained above, we do not observe any L-TAS signals in this range due to faster back-electron transfer between the electrons at the TiO₂ and the oxidized porphyrin catalysts.

4. CONCLUSION

In summary, we determined the charge transfer kinetics of mesoporous TiO₂ and two TiO₂/iron porphyrin hybrid films by employing L-TAS. The results were corroborated by those obtained in the evaluation of their performance as photocathodes in the CO₂R. Our findings demonstrate that the kinetics and the nature of the photoinduced electron transfer, as well as charge recombination processes occurring in the films, have a noticeable impact on the efficient electron transfer to the adsorbed CO₂ molecules and their CO₂R reactivity. Under excitation with $\lambda = 355$ nm light and applied external bias, the photogenerated electrons in bare TiO₂ showed a large decay of the transient signal intensity of the charge-separated state. In contrast, we observed faster recombination kinetics in

the presence of CO₂ and TEOA, used as a hole scavenger. For the TiO₂/iron porphyrin films, the recombination kinetics of the charge-separated states were fast and not modified in the presence of CO₂. The efficiencies of the materials in the CO₂R were comparable. Although the film functionalized with Hemin units is quite stable, the analogue based on the iron porphyrin had leaching issues at moderate negative bias potentials.

The interfacial charge recombination processes, occurring between the oxidized iron porphyrin and the electrons in the conduction band of TiO₂, caused a diminution of the direct charge transfer process to the CO₂ molecules. This phenomenon is translated into a depletion of the CO₂ reduction efficiencies of the hybrid film materials. Under photo- and photoelectrochemical conditions, the TiO₂ film produced CO and CH₄ as CO₂ reduction products at potentials ranging from 0 to -1.25 V. In contrast, TiO₂/Hem and TiO₂/Fe-Por films exclusively produced CO under comparable conditions. All films were better cathodes under light irradiation, in terms of efficiency and overpotential requirement for the CO₂R. This finding made us investigate in detail the electron-/charge-transfer processes occurring in the materials. We emphasize that the use of L-TAS we applied for the investigation of the photochemical CO₂ reduction could be extended to a wide variety of catalytic systems. Using this methodology, it is possible to gain further insight into the charge-transfer kinetics occurring at the catalyst surface. We anticipate that a better understanding of the detrimental steps involved in the interfacial charge-transfer processes and other limiting factors regarding the stability of the TiO₂/iron porphyrin assemblies will help significantly in designing improved and unprecedented metalloporphyrin-semiconductor-based photocathodes for the CO₂R.

5. EXPERIMENTAL SECTION

5.1. Materials and Reagents. All reagents, including pyrrole, pentylbenzene, methyl 4-formylbenzoate, acetic acid 99%, Hemin $\geq 96\%$, acetonitrile and *N,N*-dimethylformamide anhydrous 99.8% (ACN and DMF), triethanolamine 99% (TEOA), tetrabutylammonium hexafluorophosphate $\geq 99.0\%$ (TBAPF₆), 2,6-lutidine, and iron(II) bromide (FeBr₂) were purchased from Sigma-Aldrich and were used as received without further purification. TBAPF₆ was dried under vacuum before use. FTO conducting glass substrates (TEC 15 Ω/cm^2 resistance) were purchased from Xop Glass, Castellón, Spain. Nanocrystalline titania paste was purchased from Dyesol (30NR-D titania paste, 20 nm particle size). Aluminum oxide, 20% in H₂O, colloidal dispersion was purchased from Alfa Aesar.

5.2. Alumina Paste Synthesis. The Al₂O₃ paste was prepared using a commercial water-based colloidal dispersion of Al₂O₃ nanoparticles with a uniform diameter of 50 nm (Alfa-Aesar). Briefly, 37.5 mg of Al₂O₃ solution and 3 mg of acetic acid 99% were mixed and stirred for 1.5 days, until the mixture became thick.

5.3. TiO₂ and Al₂O₃ Porphyrin Film Fabrication. A nanocrystalline TiO₂/Al₂O₃ paste was deposited onto FTO conducting glass substrate by the well-known doctor blade technique. The glass-FTO substrate was previously cleaned by washing with soap, distilled water, and ethanol and dried by heating at 500 °C for 15 min. After the film fabrication, the resulting films were dried for ~30 min at room temperature and gradually heated in a muffle furnace at 325 °C for 5 min, 375 °C for 5 min, 450 °C for 15 min, and 500 °C for 15 min for sintering. The glass films, with a resulting thickness of 4–5 μm , were cut into pieces with an active area of 1 cm². TiO₂/Al₂O₃ electrodes were functionalized by dip-coating into a solution of 0.5 mM Hemin (Hem) or 0.5 mM Porphyrin (Fe-Por) in MeOH for 30 min at room temperature in the dark. Afterward, they were rinsed with MeOH, dried with air, and placed in an oven at 70 °C for 1 h.

5.4. TiO₂ and Al₂O₃ Porphyrin Film Electrode Preparation.

An electrical wire connection to the small glass film (electrode with 1 cm² area) was made by using ultrasonic shouldering (USS-9210, MBR electronics) systems and Cerasolzer as an active solder alloy. Except for the TiO₂-coated film area, other parts of the electrode were well-wrapped with Teflon tape before placing it inside the electrochemical/TAS measurement cell to protect it from any types of solvent-assisted contamination during the measurements.

5.5. Transient Spectroscopic Characterization Techniques.

The charge-transfer dynamics of TiO₂/Al₂O₃ iron porphyrin films were measured using laser transient absorption spectroscopy (L-TAS) from the microseconds to milliseconds time scale at room temperature. Measurements were carried out employing a NdYAG laser as the excitation source (355 nm, 7 ns pulse width, 1 mJ of power) and a 150 W tungsten lamp as the light probe. All experiments were performed in a homemade Teflon cell with two quartz windows. The cell was purged with N₂ for 15 min filled with anhydrous TBAPF₆/acetonitrile (0.1 M) electrolyte, containing 0.01 M triethanolamine (TEOA) when stated. For measurements under CO₂, the electrolyte was purged with CO₂ for 15 min. Bias was applied using a Keithley 2400 instrument in a 4-wire configuration as a potentiostat. Ag/AgCl (1 M KCl) was used as the reference electrode, and Pt wire was used as the counter electrode.

5.6. Electrochemical and Photoelectrochemical Measurements.

All the electrochemical experiments were carried out with a Bio-Logic potentiostat equipped with EC-Lab software in a homemade single-compartment glass cell. Ag/AgCl (1 M KCl) was used as the reference electrode, Pt wire was used as the counter electrode, and a TiO₂ or TiO₂/iron porphyrin film was used as the working electrode. The cell was purged with Ar for 15 min, and it was filled with anhydrous acetonitrile/TBAPF₆ (0.1 M) electrolyte prior to the measurements. For measurements under CO₂, the electrolyte was purged with CO₂ for 15 min. For controlled-potential electrolysis (CPE), the same single-compartment glass cell was used. The cell was purged with CO₂ for 15 min and filled with anhydrous acetonitrile/TBAPF₆ (0.1 M) electrolyte, followed by purging for 15 min more. The counter electrode bridge was filled with 0.05 M Et₄COONH₄ salt along with acetonitrile/TBAPF₆ (0.1 M) electrolyte. Then, the CO₂ flow was removed, ensuring the perfect sealing of the cell. The electrolyte was maintained under constant stirring in the course of the experiment. During the photoelectrochemical experiments, a homemade single-compartment glass cell with a one-sided quartz window was used for the electrolysis experiment, keeping the same electrolyte and reference, counter, and working electrodes as mentioned for the electrochemical experiments. The electrode was irradiated with a high-power light source purchased from Sahlmann Photochemical Solutions consisting of 3 LED diodes from Nichia (365 nm, 241.5 mW cm⁻²). The sample was located 1.5 cm from the power source (LED). The CO₂ reduction gaseous product was characterized with an Agilent GC instrument by manual injection of the sample taken from the head space of the glass cell by gastight Hamilton micro syringe. The GC is configured with HP-PLOT-Q and a Molsieve column, argon as the carrier gas, and flame ionization (CH₄, CO) and thermal conductivity (H₂) detectors (FID and TCD). The calibration of each gas (CH₄, CO, H₂) were done prior to the CPE measurements using a standard mixture of the analyte gases obtained from Messer. The product selectivity (SE) and turnover numbers (TON) were calculated following a standard literature practice.^{61,62}

■ ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at <https://pubs.acs.org/doi/10.1021/acsami.2c22458>.

Synthesis of Fe-Por, UV-vis spectra of Fe-Por, ¹H NMR spectra of Fe-Por, MALDI mass of Fe-Por, adsorption kinetics of Hem and Fe-por on TiO₂, IR spectra of FTO-TiO₂/Fe-Por and FTO-TiO₂/Hem, XPS of FTO-TiO₂, FTO-TiO₂/Fe-Por, and FTO-

TiO₂/Hem, TAS of TiO₂, TiO₂/Fe-Por, and TiO₂/Hem under bias, no external bias, and different conditions, CV of Hem and Fe-Por in DMF, CV of TiO₂, TiO₂/Hem, and TiO₂/Fe-Por films in DMF, GC chromatogram for the CO and CH₄ gases measured after an electrolysis study (PDF)

■ AUTHOR INFORMATION

Corresponding Authors

Tamal Chatterjee – Institute of Chemical Research of Catalonia (ICIQ), The Barcelona Institute of Science and Technology, 43007 Tarragona, Spain; Email: tchatterjee@iciq.es

Pablo Ballester – Institute of Chemical Research of Catalonia (ICIQ), The Barcelona Institute of Science and Technology, 43007 Tarragona, Spain; Catalan Institution for Research and Advanced Studies (ICREA), 08018 Barcelona, Spain; orcid.org/0000-0001-8377-6610; Email: pballester@iciq.es

Emilio Palomares – Institute of Chemical Research of Catalonia (ICIQ), The Barcelona Institute of Science and Technology, 43007 Tarragona, Spain; Catalan Institution for Research and Advanced Studies (ICREA), 08018 Barcelona, Spain; orcid.org/0000-0002-5092-9227; Email: epalomares@iciq.es

Authors

Beatriu Domingo-Tafalla – Institute of Chemical Research of Catalonia (ICIQ), The Barcelona Institute of Science and Technology, 43007 Tarragona, Spain; Universitat Rovira i Virgili (URV), Departament D'enginyeria electrònica Elèctrica i Automàtica, Avinguda Paisos Catalans, 43007 Tarragona, Spain; orcid.org/0000-0001-8579-7677

Federico Franco – Institute of Chemical Research of Catalonia (ICIQ), The Barcelona Institute of Science and Technology, 43007 Tarragona, Spain

Javier Perez Hernandez – Institute of Chemical Research of Catalonia (ICIQ), The Barcelona Institute of Science and Technology, 43007 Tarragona, Spain; orcid.org/0000-0002-8503-5717

Eugenia Martinez-Ferrero – Institute of Chemical Research of Catalonia (ICIQ), The Barcelona Institute of Science and Technology, 43007 Tarragona, Spain; orcid.org/0000-0001-7558-0271

Complete contact information is available at: <https://pubs.acs.org/doi/10.1021/acsami.2c22458>

Notes

The authors declare no competing financial interest.

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