Better Together: Ilmenite/Hematite Junctions for Photoelectrochemical Water Oxidation

Serena Berardi, Jagadesh Kopula Kesavan, Lucia Amidani, Elia Marek Meloni, Marcello Marelli, Federico Boscherini, Stefano Caramori, and Luca Pasquini*

ABSTRACT: Hematite (α-Fe₂O₃) is an earth-abundant indirect n-type semiconductor displaying a band gap of about 2.2 eV, useful for collecting a large fraction of visible photons, with frontier energy levels suitably aligned for carrying out the photoelectrochemical water oxidation reaction under basic conditions. The modification of hematite mesoporous thin-film photoanodes with Ti(IV), as well as their functionalization with an oxygen-evolving catalyst, leads to a 6-fold increase in photocurrent density with respect to the unmodified electrode. In order to provide a detailed understanding of this behavior, we report a study of Ti-containing phases within the mesoporous film structure. Using X-ray absorption fine structure and high-resolution transmission electron microscopy coupled with electron energy loss spectroscopy, we find that Ti(IV) ions are incorporated within ilmenite (FeTiO₃) near-surface layers, thus modifying the semiconductor−electrolyte interface. To the best of our knowledge, this is the first time that an FeTiO₃/α-Fe₂O₃ composite is used in a photoelectrochemical setup for water oxidation. In fact, previous studies of Ti(IV)-modified hematite photoanodes reported the formation of pseudobrookite (Fe₂TiO₅) at the surface. By means of transient absorption spectroscopy, transient photocurrent experiments, and electrochemical impedance spectroscopy, we show that the formation of the Fe₂O₃/FeTiO₃ interface passivates deep traps at the surface and induces a large density of donor levels, resulting in a strong depletion field that separates electron and holes, favoring hole injection in the electrolyte. Our results provide the identification of a phase coexistence with enhanced photoelectrochemical performance, allowing for the rational design of new photoanodes with improved kinetics.

KEYWORDS: photoelectrochemistry, hematite, titanium, EXAFS, electron microscopy, transient absorption spectroscopy, electrochemical impedance spectroscopy, heterointerface, oxygen evolution catalyst

1. INTRODUCTION

The need to find solutions to the global energy crisis has prompted the scientific community to propose various approaches aimed at the exploitation of renewable energy sources. Among them, solar energy conversion in the form of chemical energy (so-called artificial photosynthesis) can be effectively realized in photoelectrochemical cells (PECs). In such devices, the illumination of semiconductor-based materials triggers several complex charge separation processes, leading to the formation of energy-rich molecules, also known as solar fuels. One of the most investigated alternative fuels, hydrogen, can be obtained at the cathodic side of a PEC system for photoinduced water splitting, while at the anodically side, hydrogen evolves. In order to realize optimized composite photoanodes, it is of paramount importance to gain a thorough understanding of the physical properties of commonly used materials as well as to investigate new ones.

In particular, in this work we use advanced spectroscopic and electrochemical techniques in order to shed light on the interfacial composition and the charge-transfer dynamics of Ti(IV)-modified hematite films with enhanced PEC performance.

The photoanodic platform of choice, i.e., hematite (α-Fe₂O₃), is widely used in PEC-based systems, thanks to (i) its intermediate band gap (ca. 2.2 eV), allowing for the absorption of visible photons up to 590 nm, (ii) the valence band maximum favorably located with respect to the potential of the H₂O/O₂ redox couple, (iii) the essentially null toxicity, (iv) the good chemical stability in aqueous alkaline media, and (v) its abundance, leading to low cost. However, the photoelectrochemical performance of hematite is limited by the low mobility of charge carriers and the slow hole transfer to the electrolyte. These detrimental aspects are generally...
addressed using strategies involving doping, nanostructuring, and the functionalization with catalytic layers, or a combination thereof.\(^5\)

In our work, Ti(IV) was chosen as the dopant agent since it is reported to increase the electron density and the overall conductivity of the resulting material,\(^7\) as well as to passivate hematite surface states.\(^7,8\) All these aspects contribute to extend the lifetime of the charge carriers, reducing their recombination and improving the photocurrent. However, the actual origin of the Ti-induced improvement of the PEC performance remains elusive to date. We thus use X-ray absorption fine structure (XAFS) and high-resolution transmission electron microscopy (HRTEM) as powerful tools to study the local atomic and electronic structure of condensed matter. Detailed analysis of the fine structure oscillations in the extended energy range (the so-called EXAFS: extended X-ray absorption fine structure) based on the real-space multiple scattering formalism is able to provide a quantitative determination of the composition of the first few coordination shells around the excited atom, the interatomic distances, and their spread around the average value. Moreover, analysis of the line shape of the spectral region near the absorption edge (so-called XANES: X-ray absorption near-edge structure) can provide important information on the oxidation state, valence, atomic geometry, and site/symmetry selected density of states (DOS) of unoccupied electronic states. Due to its local character, XAFS is particularly useful to study disordered or defective atomic arrangements in condensed matter, including especially thin films. Also very important for the present investigation, XAFS is the premier structural tool to determine the incorporation site of dopants in condensed matter since it is possible in many cases to isolate the XAFS cross section of the excited atom from the background contribution.\(^9,11\) The objectives of the XAFS measurements in the present case were to assess the possible modification of the hematite matrix upon addition of Ti and to determine the incorporation sites. Concerning the second point, we sought to determine whether Ti is found in a local structure reminiscent of one of the titanium oxides, as an isolated Ti impurity substitutional to Fe, or in ilmenite, an Fe-Ti mixed oxide, or in any other form. The spatial distribution of Ti-containing phases within the mesoporous thin films was determined by TEM equipped with electron energy loss spectroscopy (EELS), which provides information complementary to XAFS together with the high spatial resolution typical of TEM.

At the same time, we employ transient absorption spectroscopy (TAS), transient photocurrent (TPC) experiments, and electrochemical impedance spectroscopy (EIS), in order to evaluate the effect of the interfacial modifications on the collection and recombination dynamics of the photo-generated charges, involving the participation of different kinds of surface states. It has recently been pointed out that this is a critical (and often overlooked) issue for the operation of hematite-based photoanodes for PEC water oxidation, which can be usefully engineered and exploited.\(^1,2\)

Our work thus provides spectroscopic and photoelectrochemical insights enabling us to unravel the role of Ti(IV) and catalytic layers on the performance of composite hematite electrodes, as well as to rationally design new photoanodes with optimized kinetics.

2. EXPERIMENTAL METHODS

The photoanodes were prepared by depositing mesoporous hematite thin films (300–400 nm thick, indicated as MPH) on fluoride-doped tin oxide (FTO) slides following an easy-to-scale electrophotoretic method.\(^13,14\) When the preparation is made in the presence of a given amount of dissolved titanium (IV) butoxide (nominal Ti content of 5 and 10 wt %), Ti-modified MPH_5Ti and MPH_10Ti electrodes are obtained, respectively. Detailed information on the procedure is provided in the Supporting Information. Some of the MPH_5Ti photoanodes were further functionalized with an amorphous iron oxyhydroxide oxygen-evolving catalyst (FeOEC) deposited by successive ionic layer adsorption and reaction (SILAR), as previously reported in conjunction with different absorbing materials.\(^13,15,17\) These electrodes are named MPH_5Ti-FeOEC throughout the manuscript.

Photoelectrochemical measurements were carried out with a PGSTAT 30 electrochemical workstation in a three-electrode configuration, using a saturated calomel electrode (SCE) and a Pt wire as reference and counter electrodes, respectively. A LOT-Oriel solar simulator, equipped with an AM1.5G filter and set to 0.1 W/cm\(^2\) incident irradiance power, was used as the illumination source. J–V curves were recorded at 20 mV/s scan rate.

For the EIS measurements under illumination, the photoanodes were sampled in the selected potential range at 50 mV intervals, employing an FRA2.v10 frequency response analyzer controlled by software Nova 1.10. A 10 mV amplitude sinusoidal perturbation, the frequency of which ranged between 100 kHz and 0.1 Hz, was used. The EIS data were fitted by means of the equivalent circuit reported in Figure S13D using the ZView software, with typical relative errors lower than 15%.

A nanosecond TAS was carried out using pump pulses with 355 nm wavelength and ca. 500 µJ/cm\(^2\)/pump fluence. The spectra were collected both under open-circuit potential \(V_{OC}\) and under anodic bias in a two-electrode configuration using a Pt wire as a counter electrode. TPC experiments were performed under the same excitation conditions, with the photoanodes placed in a three-compartment cell. TPC measurements were also conducted under a white light bias from a solar simulator set to 0.4 W/cm\(^2\). Further details on the experimental setup for transient measurements are provided in the Supporting Information.

All experiments were performed in a 0.1 KOH aqueous electrolyte (pH 13.3). Unless otherwise stated, the potential values are given versus the reversible hydrogen electrode (RHE), applying the following conversion:

\[
V_{V,SCE} = V_{V,RHE} + 0.24 + 0.059(pH)
\]

XAFS measurements were carried out at the Ti and Fe K edges at BM23 of the European Synchrotron Radiation Facility (ESRF), Grenoble. Measurements on Ti-modified MPH were performed in fluorescence mode using a Vortex Si drift diode detector placed in the horizontal plane at right angles to the impinging beam, at room temperature. Reference powder samples of hematite, magnetite, rutile, anatase, brookite, ilmenite, and pseudobrookite were also measured in transmission mode. The pretreatment process for all XAFS spectra was performed using the Athena program of IFEFFIT software package.\(^18\) The pre-edge features were fitted with Gaussian profiles using the Fityk package.\(^19\) The EXAFS spectra were analyzed using the Artemis code\(^19\) using simulated scattering paths calculated using FEFF 6.0.

TEM analyses were conducted on specimens prepared by scratching the photoanodes surface with a scalpel and collecting the powder by adherence onto a copper TEM grid coated by a holey carbon film.\(^20\) The employed instrument was a ZEISS LIBRA200 FE electron microscope equipped with an in-column Ω-filter analyzer. ESI (electron spectroscopy imaging) elemental maps for Ti and Fe were collected at the L\(_{3,3}\) edge (at energy loss 462 and 708 eV, respectively) by a three-windows methods. EELS spectra were collected at the Ti L\(_{3,3}\) edge in the energy loss range of 420–500
3. RESULTS AND DISCUSSION

The photoelectrochemical performances of the photoanodes, collected under 1 sun illumination (0.1 W/cm², AM1.5G) in 0.1 M KOH, are reported in Figure S1A. The results confirm a 4-fold improvement of the maximum photocurrent density for the MPH_5Ti electrodes (up to 1 mA/cm² at 1.67 V vs RHE) with respect to the bare MPH. A decrease in the donor density for MPH_5Ti (and MPH_5Ti-FeOEC) versus the bare MPH, likely due to the formation of a junction between hematite and a Ti-containing phase. However, the exact identification of the local structure of this phase could not be determined either by X-ray photoemission spectroscopy (XPS) or by X-ray diffraction (XRD). XPS shows that the oxidation states of Fe and Ti are 3+ and 4+, respectively. However, this does not allow us to distinguish between different Ti-containing phases like TiO₂ (rutile, anatase, brookite), FeTiO₃, and Fe₂TiO₅. In order to provide an in-depth understanding of the origin of the enhanced PEC performance of these modified photoanodes, the local structure and oxidation state of Fe and Ti have been studied by XAFS and TEM–EELS.

3.1. XAFS and TEM Measurements on the Photoanodes. The background-subtracted Fe K edge EXAFS spectra of MPH_5Ti and MPH_10Ti and reference hematite are reported in Figure S2. It is clear that the spectra are all very similar to each other. Hematite has trigonal crystal structure with space group 167 (i.e., R3c:H) in which each Fe atom is found in a distorted octahedron formed by six O ones (three atoms at 1.93 Å and three at 2.08 Å). Figure 1 shows as continuous lines the Fourier transforms of the EXAFS spectra performed in the range of 3–13.9 Å⁻¹. These spectra were fitted in R-space in the range of 1–4.8 Å using the known structure of hematite to calculate scattering paths.

The many-body amplitude reduction factor was fixed to the value obtained from a fit of a hematite reference spectrum (S₀² = 0.90); the fitting parameters were a common value of the energy origin shift (∆E₀) and, for each contribution listed below, the interatomic distances (R) and Debye–Waller factors (σ²). R goodness-of-fit factors were ~0.01 for all samples, and the fits are reported as empty circles in Figure 1.

We found that the first peak at ~1.5 Å and a small shoulder at ~1.90 Å are due to the contribution from single scattering of the six first-shell O atoms, the second peak is due to one Fe atom at 2.86 Å and three Fe atoms at 2.97 Å, and the third peak is due to three Fe atoms at 3.38 Å and six Fe atoms at 3.68 Å. No multiple scattering contributions were included because of their very low amplitude. The quantitative results of the fit are listed in Table S1. The fitting parameters of Ti-modified MPH and those of reference hematite exhibit negligible differences, being both compatible with literature data for hematite; this indicates that the presence of Ti(IV) does not alter significantly the local structure of Fe, which remains very similar to that of hematite.

The Ti K edge XANES of Ti-modified MPH photoanodes and of pseudobrookite (Fe₂TiO₅), anatase, brookite, rutile, and ilmenite reference samples are shown in Figure 2. The pre-edge features arise from dipole and quadrupole transitions to bound electronic states in the bottom of the conduction band originating from Ti 3d/4p and O 2p hybridization, and their line shape is strongly affected by the oxidation state and local symmetry around Ti. The pre-edge features are highlighted in the left panel of Figure 3. It is known that the A1, A2, and A3 features are due to transitions to t₂g states and feature B to transitions to e_g states. The rising and main edge region is attributed to the electronic transition of 1s electrons to continuum states. It can be noted that all spectral features of Ti-modified MPH are similar to those of ilmenite (with some extra broadening of the main edge) and are significantly different from those of all titanium oxides and pseudobrookite. This indicates clearly that Ti is found in a local structure very similar to that of ilmenite.

![Figure 1. Fourier-transformed Fe K edge EXAFS (k²-weighted) spectra of Ti-modified MPH and reference hematite with their best fits (data reported as continuous line and fits as empty circles). Spectra have been vertically offset for clarity.](https://dx.doi.org/10.1021/acsami.0c12275)  

![Figure 2. Ti K edge normalized XANES spectra of Ti-modified MPH compared to pseudobrookite, anatase, brookite, rutile, and ilmenite reference spectra.](https://dx.doi.org/10.1021/acsami.0c12275)
The pre-edge spectral features of the Ti-modified MPH and ilmenite were fitted by Gaussian components, as shown in the left panel of Figure 3. The numerical results are reported in Table S2. Since it is known that the intensity and the energy position of each peak are related to oxidation state and local coordination, we report these quantities in the right panel of Figure 3 as an intensity versus energy position plot. It is clear that, within the uncertainties, the energies of all components are equal, while there are some differences in the intensities and widths. Our conclusion is that the local electronic structure around Ti near the bottom of the conduction band are similar to that in ilmenite. This result points to a similar atomic structure, which was assessed by Ti K edge EXAFS as described below.

The background-subtracted Ti K edge EXAFS spectra are shown in Figure S3. The spectra of Ti-modified MPH have a line shape similar to the ilmenite spectrum. This indicates that the local coordination around Ti is very similar to the ilmenite local structure, as expected. Ilmenite has trigonal crystal structure with space group 167 (i.e., R3c:H) as hematite, and this structure was used to fit the spectra.

The Fourier transforms of Ti K edge EXAFS spectra, obtained in the range of 2.5–9.5 Å⁻¹, are shown in Figure 4. The only notable differences between the samples and ilmenite are a small shift to higher distances of the first peak and a smaller intensity of the second peak with respect to the first.

In the quantitative EXAFS analysis, we sought to distinguish between two options: (i) substitutional Ti atoms within the hematite matrix or (ii) local structure around Ti similar to ilmenite. The key difference between the two is the contributions giving rise to the second peak in the Fourier transforms: in the first case only Ti–Fe contributions would be present, while in the second both Ti–Ti and Ti–Fe are expected. Any attempt to fit the samples’ spectra with only Ti–Fe contributions yielded unphysical structural parameters; we thus conclude that the local structure is in fact similar to ilmenite. Therefore, all fits were performed on the basis of the known local structure of ilmenite. The many-body amplitude reduction factor was obtained from the best fit of the ilmenite spectrum \(S_0^2 = 0.84\), and this was fixed in the fits of Ti-modified MPH. The other fitting parameters were a common energy origin shift \(\Delta E_0\) and interatomic distances \(R\) and Debye–Waller factors \(\sigma^2\) for each contribution listed below. The first peak in the ilmenite spectrum was fitted with contributions originating from three O atoms at \(\sim 1.84\) Å and three more at \(\sim 2.02\) Å, the second peak with three Ti atoms at \(\sim 2.99\) Å, three Fe atoms at \(\sim 3.41\) Å, and six O atoms at \(\sim 3.29\) Å, and the third by six Fe atoms at \(\sim 3.76\) Å. No multiple scattering contributions were considered due to their low amplitude. The photoanodes spectra were fitted in a similar fashion, and slight variations in the local structural parameters were obtained, as summarized in Table S3. Debye–Waller factors are nearly always higher in Ti-modified MPH, indicating a more structurally distorted environment. A significant difference is that the Ti–O distances in the photoanodes are significantly higher than in ilmenite and tend to the values of the corresponding Fe–O distances in hematite. This might reflect the influence of the hematite matrix on the ilmenite-like regions.

This view is reinforced by TEM analyses that additionally provide local information on the phase distribution at the nanoscale. Structural analyses by selected area electron diffraction (SAED) and HRTEM confirm hematite as the main phase but also reveal the presence of ilmenite. SAED analysis (Figure S4) highlights a polycrystalline structure; the
diffraction rings characteristic of hematite are clearly visible. Ilmenite shares the trigonal crystal structure with hematite, and despite the fact that part of the reflections partially overlap, some planes specific to ilmenite, such as (101) and (003), are clearly detected in the diffraction micrograph. Moreover, HRTEM structural analysis in proximity of the surface (Figure 5, parts A and B) reveals again the hematite main phase (Figure 5B, zone axis [2 −3 2]) together with reflections indexed to ilmenite (104) planes (marked by arrows Figure 5B). From the morphological point of view, the MPH_STi sample appears to be constituted by well-shaped crystalline nanoparticles of about 40–70 nm diameter enveloped by a conformal layer with a thickness in the range of 2–4 nm (pointed by arrows in Figure 5, parts C and D). The ESI analysis shown in Figure 6, performed by mapping the Ti and Fe L_{2,3} energy loss feature, clearly shows Ti enrichment of this layer.

The similarity between the Ti L_{2,3} edge EELS spectra for sample MPH_STi and for an ilmenite reference sample (Figure 7) clearly indicates that the Ti-rich outer layers around the nanoparticle are ilmenite, in agreement with the outcome of EXAFS analysis. In fact, the energies of the edge onsets for both the L_{2} edge at 464.3 eV and the L_{3} edge at 458.9 match well the reference ones at 464.7 and 459.3 eV, respectively; moreover, the overall line shape (including the corresponding branching ratios) of the MPH_STi spectrum corresponds quite well with that of the ilmenite reference. Moreover, the EELS spectrum of sample MPH_STi differs significantly from those of rutile, anatase, and amorphous TiO₂ (Figure S5).

Although ilmenite is reported to have a valence band edge which is not properly aligned to accept holes from the valence band (VB) of hematite, in our case the very thin (2–4 nm) FeTiO₃ layer does not hinder the hole transfer to the electrolyte, as clearly demonstrated by the significant improvement of the PEC performance of the MPH_STi electrodes compared to the MPH counterpart (Figure S1). A favorable band alignment is instead obtained in other kind of composites, such as those constituted by hematite and pseudobrookite Fe₂TiO₅,7,30,31 In this case, the Fe₂TiO₅ phase allows for the formation of a type II heterojunction with α-Fe₂O₃, leading to maximum photocurrent values up to 3 mA/cm² when a nickel/iron oxide OEC is introduced.31

Furthermore, to the best of our knowledge, our report is the first in which a thoroughly characterized Fe₂O₃/FeTiO₃ composite is used in a photoanodic setup for water oxidation. Indeed, this combination is generally used for its magnetic properties, while photoelectrochemical or photocatalytic applications often exploit Fe₂TiO₅/TiO₂ junctions.34–36

### 3.2. TAS and TPC Measurements on the Photoanodes

In order to evaluate the effect of the ilmenite phase on the interfacial dynamics of the photogenerated charges, we have performed TAS and TPC measurements on unmodified MPH and on MPH_STi, i.e., the best-performing photoanode in terms of Ti(IV) content. The latter was also functionalized with FeOEC with the aim to investigate its role in photoelectrochemical water oxidation.

The TA spectra for MPH and MPH_STi photoanodes at V_{OC} are reported in Figure 8, parts A and B. The laser excitation power was kept as low as possible in order to observe dynamics that could be related to the behavior of the photoanode under solar illumination, resulting in a small optical density difference (ΔOD) in the order of 10⁻⁴. TA spectra at V_{OC} are dominated by a broad absorption band at wavelengths between 540 and 640 nm peaking around 580–590 nm, which decays in the millisecond time scale. This feature, common to both MPH and MPH_STi, can be attributed to photohole trapping in intragap states located a few hundreds of millielectronvolts below the conduction band, as demonstrated by the Durrant group for other nanostructured hematite photoanodes.37 In the absence of an applied bias, these intragap states (below the Fermi level) are occupied by electrons and therefore do not contribute to ground-state absorption. After laser pulse excitation, valence

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**Figure 5.** (A) HRTEM micrographs of MPH_STi and (B) related fast Fourier transform (FFT) of the selected region of interest; arrows indicate the reflections indexed as ilmenite (104) planes. (C) Low-magnification TEM micrograph and (D) HRTEM micrograph that shows the 2–4 nm thin conformal layer around nanoparticles.

**Figure 6.** TEM micrographs of MPH_STi, related ESI Fe and Ti maps, and the combined map.

**Figure 7.** EELS spectra for MPH_STi specimen (above) and reference ilmenite (below) recorded under the same experimental conditions at the Ti L_{2,3} edge.
Band photoholes are rapidly (≪milliseconds) trapped at these states, making them available for optical transitions of valence band electrons that lead to the observed transient absorption band. Its decay is due to the filling of the intragap states by photoexcited electrons in the conduction band. In other words, the intragap states act as recombination centers. The fact that the absorption band is more intense (almost double) in the case of MPH_5Ti points to a higher number of available states for the holes to be trapped in, due to the formation of the hematite−ilmenite heterointerface.

When the photoanodes are biased at 0.8 V versus Pt (the potential at which the limiting photocurrent is observed in a two-electrode configuration; see Figure S1B for the corresponding J−V curves), the spectral characteristics change. An intense bleach appears at a similar wavelength of $V_{\text{OC}}$ transient absorption (centered at 580−590 nm for MPH and MPH_STi, respectively; see Figure 8, parts C and D). Moreover, a broad absorption band at $\lambda > 630$ nm extending into the red is visible. This spectral behavior is consistent with previous observations.37

The origin of the bleaching can be understood considering again the intragap states, as follows. Under applied anodic bias, optical absorption at 580−590 nm takes place in the ground state because the intragap states are empty, at least within the space charge region. However, after laser excitation, the intragap states are rapidly (≪milliseconds) filled by photoexcited electrons in the conduction band. This reduces absorption at 580−590 nm, generating a negative $\Delta\text{OD}$ signal, approximately 90% of which is recovered within 10 ms (Figure S6A). Moreover, as reported in Figure S6B, for both MPH and MPH_STi electrodes, the bleach intensity increases (in absolute value) with the square root of the applied bias, further confirming that this dynamics originates from the extension of the space charge layer in the photoanode upon positive polarization.

On the other hand, the broad feature at $\lambda > 630$ nm has been attributed to photoholes swept by the electric field at the semiconductor/electrolyte interface (SEI).37 It is interesting to note that it is at least 2 times higher in the Ti-modified electrode, consistent with the improved photoelectrochemical performance.
In order to investigate the role of the different components of the photoanodic interfaces in more detail, we have performed TPC measurements. The photocurrent decay was recorded when striking the photoanode from the “front” (i.e., the electrolyte side, as opposed to the FTO collector “back” side) under variable bias (Figure S7). These experiments are schematized in Figure 9, considering the different behavior of the three interfaces under 1.62 V (vs RHE) applied bias and highlighting the possible pathways for the fate of the charge carriers generated by the laser pulse.

Figure 10A reports a typical transient chronoamperometric trace in the 1–100 ms time domain, showing the following distinctive features: (i) a positive photocurrent peak, which is originated by collection of laser-generated electrons at the back contact and usually decays on a 10 ms time scale, in agreement with the transient lifetime observed by TAS. The corresponding photocurrent peak rise; in the following we call $\tau_1$ the time at half-amplitude; (ii) a negative transient (with half-amplitude time as $\tau_2$), corresponding to a photocharge that extends up to 100 ms, originated by the recombination of the photogenerated electrons with holes trapped in surface states, and thus not injected into the electrolyte. The integral of these transient features yields the temporal evolution of the photogenerated charge (inset of Figure 10A), thus allowing for the identification and quantification of the different processes framed by the two different time regimes (10 and 100 ms). In particular, we define (i) $Q_{gen}$, as the area subtended by the positive photocurrent peak, which is the result of the kinetic competition between electron transport at the collector and fast recombination processes involving reaction with mobile holes (VB holes) or holes contained in shallow traps, (ii) $Q_{rec}$, as the charge lost due to slower recombination processes involving long-lived surface-trapped holes, and (iii) $Q_{coll}$, as the actual collected charge, given by the difference between the other two contributions, according to

$$Q_{coll} = Q_{gen} - |Q_{rec}|$$

We calculate these charges for all the TPC; traces reported in Figure S7. Figure 10B shows the resulting bias dependence of $Q_{coll}$ and $Q_{rec}$ (inset) for all the investigated photoanodes, while the curves corresponding to $Q_{gen}$ are reported in Figure S8A. In general, the magnitude of the photocurrent transients increases in the order MPH < MPH_{STi} < MPH_{STi-FeOEC} (Figure S7), and the resulting trend in $Q_{coll}$ is consistent with the $J−V$ curves obtained under 1 sun illumination (see Figure S1A), confirming that the presence of the ilmenite overlayer improves the collection of photogenerated charge. We note that the $Q_{coll}$ versus $V$ trend reported in Figure 10B is exacerbated compared with the conventional $J−V$ curves (Figure S1), due to the front-side illumination geometry herein adopted, which is the most unfavorable for the collection of charge carriers in hematite.

Furthermore, the time at which the $Q_{gen}$ reaches half of its peak value ($\tau_1$; see also inset of Figure 10A) is essentially constant within the explored potential range, being ca. 0.5 ms for MPH, ca. 1.5 ms for MPH_{STi}, and ca. 4 ms for MPH_{STi-FeOEC} (see Figure S8B). This suggests that fast recombination processes are partially inhibited by the presence of the ilmenite layer, allowing it to extend the useful time frame for the collection of photogenerated electrons.

The applied bias dependence of $Q_{rec}$ (inset of Figure 10B) gives further insight on the differences in the photoanodic performance. Indeed, we obtain a distribution of low (ca. 0.05 $\mu$C) and almost constant values for MPH, meaning that the fraction of holes that are able to survive fast recombination and undergo slow recombination with electrons is smaller in such electrodes. On the contrary, a well-defined bell-shaped behavior is observed for both MPH_{STi} and MPH_{STi-FeOEC} within the interval 0.6 $\leq$ V $\leq$ 1.4 V, indicating that a more significant fraction of holes is now trapped in long-lived surface states, as a result of the interfacial modification. The distribution of $Q_{rec}$ peaks between 1.0 and 1.2 V, in agreement with the surface chemical capacitance distribution found by EIS experiments in these (vide infra) and other types of bare and Ti-modified hematite electrodes.7,13,17,38,39 We note that the $Q_{rec}/Q_{gen}$ ratio, shown in the inset of Figure 10C, is higher in the Ti-modified electrodes and that, contrary to MPH, it drops to 0 for an applied potential $\geq$1.4 V. This behavior can be explained by the build-up of a stronger electric field inside the Ti-modified electrodes compared to unmodified ones as the positive potential increases. The enhanced field...
sweeps electrons away from the surface to the collector, leaving holes free to react with water. We can argue that the space charge inside unmodified hematite does not develop enough to guarantee efficient collection, while the structural changes induced by Ti incorporation at the interface allow it to sustain a stronger electric field and bring about a redistribution of surface states. These effects, pointed out by the Andreu group for FeO$_x$/Fe$_2$TiO$_4$, heterojunctions,' may be also related to Fermi-level pinning that was observed in the presence of a thin buffer layer of hematite underneath the MPH and greatly improved electron collection. Strikingly, under white light illumination the magnitude of the photocurrent transients in the best-performing Ti-modified MPH electrodes is strongly reduced, resulting in $Q_{gen,L}$ values that are ca. 5 and 10 times lower than $Q_{gen}$ for MPH$_{5Ti}$ and MPH$_{5Ti}$-FeOEC, respectively. By contrast, $Q_{gen,L}$ remains substantially unvaried with respect to the dark counterpart in the case of MPH. Figure 11A shows that $Q_{gen,L}$ values are higher (up to 1.4 V) in unmodified MPH compared to Ti-modified MPH, in spite of its poorer photoanodic performance. At the same time, $Q_{gen,L}$ maintains the expected increase when the bias becomes more anodic. This counter-intuitive behavior can be rationalized in terms of charge recombination at the surface of the electrodes between electrons produced by the laser pulse and holes generated by white light. It should be recalled that the laser pulse strikes the electrolyte side and that the penetration depth of 355 nm radiation in MPH films is estimated to be in the 25–50 nm range [note: the optical penetration depth ($\lambda$) of 355 nm photons in our hematite samples was directly calculated from the film thickness ($l$) and absorbance ($A$), using the formula $\alpha$ (cm$^{-1}$) = ln(10)$A/l$ (cm), in good agreement with the known extinction coefficient of hematite].

The constant white light illumination generates electrons and holes, which are swept in opposite directions by the electric field created inside the photocatalyst. Electrons travel to the collector, while holes accumulate at the SEI. The stronger the electric field at a given potential, the higher the hole density at the interface with the electrolyte. Thus, in the electrodes displaying a stronger depletion layer, i.e., MPH$_{5Ti}$ and MPH$_{5Ti}$-FeOEC, the laser pulse generates electrons in a spatial region densely populated by holes, causing fast recombination and decreasing the intensity of $Q_{gen}$ and Qcoll transients. In agreement with this interpretation, the time at half-amplitude measured for $Q_{gen,L}$ (named $\tau_{1,L}$ and reported in Figure S11) is ca. half of the dark $\tau_1$ value for MPH$_{5Ti}$ and up to 4 times lower than $\tau_1$ for MPH$_{5Ti}$-FeOEC (compare Figures S8B and S11). On the contrary, $\tau_{1,L}$ and $\tau_1$ are very similar in the case of MPH photoanodes. The stronger decrease in the $\tau_{1,L}$ values in the presence of the catalyst can be well-explained by the occupation of the surface FeOEC sites by the photogenerated holes at $V \geq 1.0$ V.

Due to fast recombination within the penetration depth of 355 nm laser pulses, $Q_{rec,L}$ also decreases by at least a factor of 5 with respect to $Q_{rec}$ for both MPH$_{5Ti}$ and MPH$_{5Ti}$-FeOEC (compare Figures 11B and 10B). In any case, $Q_{rec,L}$ maintains the bell-shaped distribution observed in the absence of white light bias and still drops to 0 for $V \geq 1.4$ V. After this

![Figure 11](https://dx.doi.org/10.1021/acsami.0c12275)
threshold, a relative increase in both $Q_{\text{gen,L}}$ and, as a consequence, $Q_{\text{coll,L}}$ can be also observed (Figure S12). Furthermore, the MPH_STi-FeOEC photoanode displays higher $Q_{\text{rec,L}}$ values with respect to those of MPH_STi. This finding suggests that, in the presence of the amorphous and electrolyte-permeable catalyst, an increased number of electronic states are available on the FeOEC for the localization of photogenerated holes.\textsuperscript{16,38}

The transient photocurrent dynamics of MPH in the presence of white light illumination are consistent with the presence of a thinner space charge layer, which can be ascribed to Fermi-level pinning by deep traps, as previously discussed. Indeed, for MPH the $Q_{\text{rec,L}}$ values are almost constant for $V \geq 0.8$ V and approximately doubled with respect to dark $Q_{\text{rec}}$ values. A plausible interpretation is that holes generated under white light and trapped in the MPH surface states do not experience a significant surface accumulation upon application of anodic bias. The larger number of photoproduced holes, with respect to the dark case, enhances their probability of slow recombination with the laser-produced electrons, explaining the increase in $Q_{\text{rec,L}}$.

The overall behavior of the analyzed photoanodes is summarized in Figure 13, in which we report the bias dependence of the ratio between the transiently collected charge in the presence ($Q_{\text{coll,L}}$) and in the absence ($Q_{\text{coll}}$) of the white light source. Once again, the observed trends clearly illustrate the differences between MPH and the ilmenite-modified MPH photoelectrodes in terms of charge separation. When the FeTiO$_3$ phase at the interface with the electrolyte is present, the $Q_{\text{coll,L}}/Q_{\text{coll}}$ drops as soon as the voltage reaches 1.0 V due to accumulation of photoholes at the SEI where laser-induced electron generation occurs. These surface hole localization effects are strongest in the best-performing electrode (MPH_STi-FeOEC), in which the steepest decrease to the lowest $Q_{\text{coll,L}}/Q_{\text{coll}}$ is related to the highest photocurrent and clearly tied to photocurrent generation, as can be appreciated from the onset of the $J$–$V$ characteristics between 0.92 and 1.02 V (Figure S1A). Thus, although the band edges of ilmenite are not properly aligned with those of hematite to promote a downhill electron-transfer cascade capable to enhance charge separation, they contribute to heterointerfacial effects, which equally result in the suppression of carrier recombination. Hence, by combining the information gained from both TPC and CV experiments, we can conclude that the formation of the FeTiO$_3$ overlayer passivates deep traps, which otherwise cause recombination, and induce a greater concentration of donor levels close to the conduction band edge, building up a stronger depletion layer.

The insights gained by TPC measurements were corroborated by means of EIS under illumination. The resulting Nyquist plots are reported in Figure S13A–C and fitted using the equivalent circuit model indicated in Figure S13D. The elements of this particular circuit, often used to model hematite-based photoanodes\textsuperscript{38,47,48} include the space charge capacitance of hematite ($C_{\text{SC}}$), the charge-transport resistance through the space charge ($R_{\text{CT,SC}}$), the surface-state capacitance ($C_{\text{SS}}$), and the resistance of the charge transfer from the surface states to the electrolyte ($R_{\text{CT,SS}}$). While $C_{\text{SC}}$ and $R_{\text{SC}}$ are related to high-frequency (i.e., faster) processes, $R_{\text{CT,SS}}$ and $C_{\text{SS}}$ are associated with the low-frequency (slower) processes taking place at the interface with the electrolyte. This results in Nyquist plots composed by two different arcs, which in our specific case collapse into a single one at high applied bias ($V \geq 1.4$ V).

The adequacy of the circuit model is confirmed by the good correlation observed when the plots of the differential resistances ($dI/dV$, calculated from the $I$–$V$ curves) versus the applied potential are compared to the corresponding $R_{\text{TOT}}$ values (Figure S14). In particular, the total resistance ($R_{\text{TOT}}$) is calculated by adding $R_{\text{SC}}$, $R_{\text{CT,SS}}$, and the serial resistance ($R_e$).

The EIS data also confirmed the expected bell-shaped distribution of $C_{\text{SS}}$ for the Ti-modified electrodes (Figure S15A), exhibiting at least 1 order of magnitude higher values with respect to those of MPH. This observation is consistent with the better charge separation observed for MPH_STi and MPH_STi-FeOEC electrodes, yielding a higher surface concentration of high-valence iron–oxo species promoting water oxidation.\textsuperscript{49} As a result, the corresponding $R_{\text{CT,SS}}$ values are considerably lower with respect to those of MPH (Figure S15B). A further contribution to the higher $C_{\text{SS}}$ values can be ascribed to the presence of a larger surface concentration of Fe(III)–OH upon Ti incorporation, as reported by the Hamann group for atomic layer deposited hematite electrodes.\textsuperscript{49}
Moreover, when MPH_5Ti is functionalized with FeOEC, the $C_{SS}$ values further increase (up to a factor 4 with respect to MPH_5Ti) and the differential capacitance peak shifts slightly in the cathodic direction. This confirms the picture of hole trapping into catalyst sites during MPH_5Ti-FeOEC operation. At the same time, it is worth noting that the maximum $C_{SS}$ nicely correlates with both the inflection point of the $J-V$ curve and the minimum value of $R_{CT,SS}$ (observed at 1.22 V for MPH_5Ti-FeOEC, see Figure S16).

Finally, we used the EIS data to calculate the efficiency for the water oxidation process ($\eta_{WO}$), as reported by the groups of Peter$^{50}$ and Mendes$^{51}$ and briefly illustrated in the Supporting Information.

Figure 14 reports the $\eta_{WO}$ values obtained for the different photoanodes, which, as expected, mirror the trend of the $J-V$ curves (see also Figure S1A). Indeed, the $\eta_{WO}$ accounts for the fact that only the photogenerated charges able to survive recombination can reach the interface and perform water oxidation. The best results were obtained for MPH_5Ti-FeOEC electrodes that are able to operate with 80% efficiency already at applied potentials close to the thermodynamic value for water oxidation. These results are in line with those obtained for other hematite-based interfaces functionalized with different metal oxides catalysts, e.g., iridium/ruthenium-based OECs$^{52}$ and nickel/iron-based OECs.$^{53}$

4. CONCLUSIONS

State-of-the-art XANES and EXAFS measurements have been used to gain advanced structural/chemical knowledge about the origin of the enhanced PEC performance in Ti(IV)-modified mesoporous hematite photoanodes. In particular, we demonstrate that the inclusion of Ti(IV) does not influence the local structure of Fe, which remains that of hematite, and that the local electronic and atomic structures around Ti are similar to those of ilmenite (FeTiO$_3$).

As illustrated by TAS, TPC, and EIS analyses, the formation of this mixed phase results in a main beneficial mechanistic pathway for the modified interfaces, related to the passivation of deep electron traps, which act as recombination centers and cause Fermi-level pinning in unmodified hematite. At the same time, it induces a higher concentration of oxygen vacancies acting as donor states close to the conduction band edge. Therefore, better charge separation is possible thanks to the buildup of a stronger electric field inside the semiconductor, which directs holes at the interface with the electrolyte.

Despite the misalignment between hematite and ilmenite band edges, the thin and discontinuous nature of the FeTiO$_3$ layer allows for a sufficient coupling between the surface-trapped holes and water for the charge transfer to occur.

We also confirmed that the functionalization of the Fe$_2$O$_3$/FeTiO$_3$ interface with an electrolyte-permeable and amorphous catalyst (FeOEC) yields a ternary composite electrode with enhanced hole injection capabilities toward the electrolyte, resulting in photocurrents up to 1.45 mA/cm$^2$ at 1.7 V versus RHE.

The present understanding of charge-transfer dynamics in low-cost and highly stable materials, made of earth-abundant and nontoxic elements, will guide the development of tailored composite photoanodes for solar water splitting with enhanced photoelectrochemical outputs.

Author Information

Corresponding Author
Luca Pasquini — Department of Physics and Astronomy, Alma Mater Studiorum—Università di Bologna, 40127 Bologna, Italy; orcid.org/0000-0001-8939-2204; Email: luca.pasquini@unibo.it

Authors
Serena Berardi — Department of Chemical and Pharmaceutical Sciences, University of Ferrara, 44121 Ferrara, Italy; orcid.org/0000-0002-0275-6501
Jagadesh Kopula Kesavan — Department of Physics and Astronomy, Alma Mater Studiorum—Università di Bologna, 40127 Bologna, Italy; orcid.org/0000-0002-0591-2493
Lucia Amidani — Helmholtz-Zentrum Dresden-Rossendorf, European Synchrotron Radiation Facility, 38000 Grenoble, France; orcid.org/0000-0003-2234-4173
Elia Marek Meloni — Department of Chemical and Pharmaceutical Sciences, University of Ferrara, 44121 Ferrara, Italy
Marcello Marelli — CNR-SCITEC, Istituto di Scienze e Tecnologie Chimiche ”Giulio Natta”, 20138 Milano, Italy
Federico Boscherini — Department of Physics and Astronomy, Alma Mater Studiorum—Università di Bologna, 40127 Bologna, Italy; orcid.org/0000-0002-9703-3903
Stefano Caramori — Department of Chemical and Pharmaceutical Sciences, University of Ferrara, 44121 Ferrara, Italy

Complete contact information is available at: https://pubs.acs.org/10.1021/acsami.0c12275

Notes
The authors declare no competing financial interest.
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