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Red-Shifted Absorptions of Cation-Defective and Surface-Functionalized Anatase with Enhanced Photoelectrochemical Properties

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1. INTRODUCTION

Owing to its semiconductor character enabling light-induced productions of charge carriers, titanium dioxide has been largely investigated.1 In particular, the pioneering work of
Fujishima and Honda on water splitting\textsuperscript{2} led to intensive research on photoelectrochemical systems using TiO\textsubscript{2}, including photoelectrochemical water splitting, photocatalysis, and dye-sensitized solar cells.\textsuperscript{3} However, current limitations of (anatase) TiO\textsubscript{2} are a limited light absorption due to a band gap energy of 3.2 eV yielding an absorption of less than 5\% of the incident solar radiation and a rapid recombination of electrons and holes produced upon photoexcitation.

Enabling the practical uses of TiO\textsubscript{2} as photoelectrodes requires a concomitant red-shift light adsorption into the visible light range and an improved charge separation upon photoexcitation. This can be achieved by using different strategies including surface engineering and/or doping/substitution with heteroatoms.\textsuperscript{1,6,7} In the latter strategy, metallic and nonmetallic dopants can produce donor/acceptor states in the band gap of TiO\textsubscript{2} to effectively extend the light adsorption into the visible light range.\textsuperscript{6,8} Cationic metal dopants, however, produce localized d states deep in the band gap of TiO\textsubscript{2} and create new recombination centers for electrons and holes.\textsuperscript{8}

Among all anionic dopants, fluoride is unique because it can act as both a morphology director and electron scavenger to reduce the recombination rate of charge carriers.\textsuperscript{9−11} There is, however, no clear understanding on how fluorine doping affects the band gap energy and the photoactivity of anatase TiO\textsubscript{2}.\textsuperscript{12,13} Czoska et al.\textsuperscript{15} reported that fluorine doping of TiO\textsubscript{2} did not show a red shift in the visible light region nor reacted with adsorbed O\textsubscript{2} to form superoxide radicals. Li et al.\textsuperscript{16} demonstrated that fluorine doping did not produce any effect on the optical absorption of TiO\textsubscript{2}, although a high visible photocatalytic activity was achieved because of the creation of surface oxygen vacancies. On the contrary, Calatayud et al.\textsuperscript{14} reported that lattice fluoride doping enabled a 0.2 eV decrease of the band gap energy. Yu et al.\textsuperscript{12} previously reported that the F-doped TiO\textsubscript{2} showed stronger absorption in the UV–visible range and a red shift. Overall, such discrepancies might come from the nature of defects induced by the partial substitution of O\textsuperscript{2−} by F\textsuperscript{−}. For instance, the stabilization of fluorine through a mechanism of valence induction,\textsuperscript{14} that is, $\text{Ti}^{IV}(1-y)\text{Ti}^{III}y\text{O}_2(2-x)\text{F}_x$, yielded a 0.2 eV decrease of the band gap energy. In this substitutional mechanism, the presence of Ti\textsuperscript{III} centers is suggested to be the physical origin of the reduced band gap.\textsuperscript{17−19}

Another substitutional mechanism associates the formation of titanium vacancies (□) through the partial substitution of divalent O\textsuperscript{2−} by monovalent F\textsuperscript{−} and OH\textsuperscript{−}. This gives the opportunity to better understand the impact of particular structural features related to the presence of F\textsuperscript{−}, OH\textsuperscript{−}, and vacancies on the band gap energy and the photogenerated carriers.

Experimentally, nanoparticles of anatase featuring the general chemical formula $\text{Ti}_{1−x}□_x\text{O}_2(2-x)\text{F}_x(\text{OH})_y$ can be synthetized by a sol–gel process activated under mild solvothermal conditions.\textsuperscript{20−23} This synthesis method allows to stabilize anisotropic particles with a large percentage of reactive (001) facets along with a high concentration of titanium vacancies arising from a large substitution rate of O\textsuperscript{2−} by F\textsuperscript{−}/OH\textsuperscript{−}.\textsuperscript{2,3,22}

In the present paper, we report the study of the impact of titanium vacancies on the band gap energy and the trapping of photoexcited electrons and holes. The reactivity of the charge carriers diffusing to the surface was monitored by photoelectrochemical measurements. An anatase $\text{Ti}_{1−x}□_x\text{O}_2(2-x)\text{F}_x(\text{OH})_y$ shows a 2-fold enhancement of quantum efficiency with a slight band gap reduction of 0.08 eV as compared to anatase TiO\textsubscript{2}. Furthermore, we took advantages of the high concentration of under-coordinated Ti to functionalize the surface by a donor molecule, that is, N-methyl-2-pyrrydolone (NMP), which further allows reducing the band gap and reaching a 5 times higher quantum efficiency as compared to anatase TiO\textsubscript{2}.

2. RESULTS AND DISCUSSION

2.1. Cation-Defective Anatase TiO\textsubscript{2} ($\text{Ti}_{1−x}□_x\text{O}_2(2-x)\text{F}_x(\text{OH})_y$). The cation-defective anatase was prepared by a previously reported method employing sol–gel chemistry performed in the presence of Hydrofluoric acid (HF). This method allows to stabilize the nanoparticles of anatase having the general chemical formula $\text{Ti}_{1−x}□_x\text{O}_2(2-x)\text{F}_x(\text{OH})_y$, where □ represents a titanium vacancy formed by the negative charge deficiency induced by the partial substitution of oxide by fluoride and hydroxide anions (Figure S1).\textsuperscript{24} In the present study, a compound featuring a high concentration ($x+y$) of titanium vacancy was prepared (see the Experimental Section). The accurate chemical composition was assessed by determining the vacancy concentration through structural refinement of the pair distribution function (PDF) data, the fluorine content was determined by $^{19}$F solid-state nuclear magnetic resonance (NMR), and the OH content was deduced by electroneutrality leading overall to $\text{Ti}_{0.69−x}□_{0.31}\text{O}_{0.76}\text{F}_{0.48}(\text{OH})_{0.76}$ (Figure S1). For comparison purpose, the cation-defective free anatase TiO\textsubscript{2} was prepared. Powder X-ray diffraction (XRD) confirms the phase purities of both anatase samples (Figure S2). The optical absorption properties of anatase TiO\textsubscript{2} and $\text{Ti}_{0.69−x}□_{0.31}\text{O}_{0.76}\text{F}_{0.48}(\text{OH})_{0.76}$ were investigated by UV–vis absorption spectroscopy (Figure 1). Doped and undoped anatase showed an absorption edge within the 350–400 nm range, which corresponds to the typical charge transfer region of O(2p)–Ti(3d).\textsuperscript{25} Although the deduced band gap energy is 3.2 eV for anatase TiO\textsubscript{2}, the band gap energy of $\text{Ti}_{0.69−x}□_{0.31}\text{O}_{0.76}\text{F}_{0.48}(\text{OH})_{0.76}$ is red-shifted to 3.12 eV. For TiO\textsubscript{2}, the conduction band is characterized by the 3d states of Ti and the valence band is composed of the 2p orbitals of O\textsuperscript{2−}. In the case of $\text{Ti}_{0.69−x}□_{0.31}\text{O}_{0.76}\text{F}_{0.48}(\text{OH})_{0.76}$, the valence band is composed of the 2p orbitals of O\textsuperscript{2−}, F\textsuperscript{−}, and OH\textsuperscript{−}. It was shown that the 2p states of F\textsuperscript{−} ions are localized rather at the bottom of the valence band, hence leading to a higher band gap energy.\textsuperscript{25} On the other hand, OH groups were proposed to decrease the band gap energy value of a titanium...
hydrosyfluoride compound featuring titanium vacancies.\textsuperscript{26} We propose that the observed reduction of the band gap energy observed is due to the presence of OH groups in larger quantity than $F^-$ ions.

Electron paramagnetic resonance (EPR) spectra were recorded (Figure 2) under dark and light irradiation (in the spectral range of 250–1000 nm) to investigate the ability of the anatase samples to generate paramagnetic centers (PCs), that is, radicals and reduced Ti\textsuperscript{IV}.\textsuperscript{27–30} The anatase TiO\textsubscript{2} EPR signal features a slightly asymmetrical shape and a g-factor value of 2.000 which can be assigned to electrons trapped at oxygen vacancies (V\textsubscript{O} + e = VO\textsuperscript{−}).\textsuperscript{31} The EPR signal of such oxygen vacancies in TiO\textsubscript{2} does not change with illumination. The most likely reason for this phenomenon may be because the defects act as recombination centers of the photogenerated electrons and holes. Therefore, the amplitude of the EPR signal does not change with irradiation. Such species can contribute to the absorption of light in the visible range because they can create levels in the forbidden band (impurity absorption). However, they are inactive for photoconversion, such as photocatalysis. Under dark condition, no PCs were detected for Ti\textsubscript{0.69}O\textsubscript{0.31}F\textsubscript{0.48}OH\textsubscript{0.76} (Figure 2), in agreement with the sole presence of Ti\textsuperscript{IV}.\textsuperscript{32} Under illumination, a weak EPR signal at around 3300–3340 G, with $g_1 = 2024$, $g_2 = 2010$, and $g_3 = 2003$, is observed to be characteristic of O\textsubscript{2−} radicals. Furthermore, an intensive signal at 3350–3370 G with $g_1 = 1.9845$ and $g_2 = 1.958$ assigned to Ti\textsuperscript{III}/oxygen vacancy PCs.\textsuperscript{27,29} Moreover, the line width of the Ti\textsuperscript{III} centers signal is small (approximately 15 G), suggesting that these PCs are isolated (no overlapping of the wave functions) and mainly localized in the bulk. Indeed, the dipole–dipole interaction between PCs localized on sample surface and surrounding paramagnetic oxygen molecules strongly increases the line width of the EPR signal.\textsuperscript{32}

The photoelectrochemical properties of anatase TiO\textsubscript{2} and Ti\textsubscript{0.69}O\textsubscript{0.31}F\textsubscript{0.48}OH\textsubscript{0.76} were evaluated using linear sweep voltammetry recorded under dark and UV–vis light conditions. The photocurrent response (Figure 3a) was calculated by subtracting the current measured under dark from the current measured under UV–vis light. Overall, the measured activity for water oxidation was low, which is probably due to the porosity of the deposited film that might affect electron–hole recombination, their electronic conductivity, and in fine the photocurrent density. However, the photocurrent response of Ti\textsubscript{0.69}O\textsubscript{0.31}F\textsubscript{0.48}OH\textsubscript{0.76} is ca. 2 times higher than that of TiO\textsubscript{2} at 0.70 V versus NHE. More interestingly, a shift of the onset potential is observed going from 0.10 V versus NHE for anatase TiO\textsubscript{2} to −0.10 V versus NHE for Ti\textsubscript{0.69}O\textsubscript{0.31}F\textsubscript{0.48}OH\textsubscript{0.76} (Figure 3a).

The higher photocurrent and the negative shift of the photocurrent onset potential clearly indicate that the anatase Ti\textsubscript{0.69}O\textsubscript{0.31}F\textsubscript{0.48}OH\textsubscript{0.76} exhibits lower electron–hole recombination rate compared to anatase TiO\textsubscript{2}. Because of the similar specific surface area for TiO\textsubscript{2} (~180 m\textsuperscript{2} g\textsuperscript{−1}, data not shown) and Ti\textsubscript{0.69}O\textsubscript{0.31}F\textsubscript{0.48}OH\textsubscript{0.76} (~190 m\textsuperscript{2} g\textsuperscript{−1}, data not shown), this phenomena can be related to the presence of surface and bulk defects in Ti\textsubscript{0.69}O\textsubscript{0.31}F\textsubscript{0.48}OH\textsubscript{0.76}.\textsuperscript{33} It has been shown that O\textsubscript{2−} radicals are very active species for photocatalytic processes, for example, to decompose toxic substances on oxide surface. Therefore, the presence of these radicals at the surface of Ti\textsubscript{0.69}O\textsubscript{0.31}F\textsubscript{0.48}OH\textsubscript{0.76} upon light irradiation, might play

![Figure 2. EPR spectra of anatase TiO2 and Ti\textsubscript{0.69}O\textsubscript{0.31}F\textsubscript{0.48}OH\textsubscript{0.76} at T = 77 K under dark and illumination.](image)

![Figure 3. (a) j\textsubscript{ph}–E plots under UV–vis irradiation and (b) photocurrent response with chopped light (frequency of 0.1 s\textsuperscript{−1}) at 1 V/NHE for TiO2 and Ti\textsubscript{0.69}O\textsubscript{0.31}F\textsubscript{0.48}OH\textsubscript{0.76} in a N\textsubscript{2}-purged electrolyte (1 M Na\textsubscript{2}SO\textsubscript{4}, pH = 5). Scan rate is 5 mV s\textsuperscript{−1}. (c) IPCE spectra of TiO\textsubscript{2} and Ti\textsubscript{0.69}O\textsubscript{0.31}F\textsubscript{0.48}OH\textsubscript{0.76} photoanodes at 1 V/NHE. (Inset: Tauc plots for the direct optical transitions of TiO\textsubscript{2} and Ti\textsubscript{0.69}O\textsubscript{0.31}F\textsubscript{0.48}OH\textsubscript{0.76} photoanodes.)](image)
a role in the modification of the charge separation process observed between these two anatases.

To further illustrate this point, photocurrent transient measurements were performed to measure the charge recombination at the semiconductor/electrolyte interface. In Figure 3b, the evolution of the photocurrents is reported when light is turned on and off, with a frequency of 0.062 Hz. Anatase TiO2 and Ti0.69□0.33O0.76F0.48(OH)0.76 show distinct behaviors. On the one hand, a sharp anodic transient photocurrent occurs upon illumination of TiO2, showing that the photogenerated charge carriers move to the semiconductor/electrolyte junction and accumulate (the carriers oxidize trap states in the bulk and on the surface).24 This is followed by photocurrent stabilization when equilibrium is reached between water oxidation and charge recombination.25 Conversely, when the light is turned off, the photocurrent decay is assigned to electrons recombining with the accumulated holes. Regarding anatase Ti0.69□0.33O0.76F0.48(OH)0.76 a gradually increase of the transient photocurrent is observed and a stabilization (stable photocurrent) occurs upon illumination of TiO2, showing that donor molecules are adsorbed on its surface, as revealed by the structural deviation occurring at the short-range order. Figure 7 shows the redefined PDF patterns of Ti0.69□0.33O0.76F0.48(OH)0.76 and NMP-Ti1-x□xO2-4(xy)F4x(OH)4y samples. The extracted structural parameters are gathered in Table S1. Both anatase samples showed similar unit cell parameters and a close coherence length of ca. 4 nm, suggesting that the long-range order was not modified upon functionalization. The refinement of the Ti site occupancy, however, shows a slight decrease of the band gap energy from 3.12 to 3.03 eV and a second feature occurring at 2.25 eV.

NMP molecules (Figure S4). These reactions are specific to Ti0.69□0.33O0.76F0.48(OH)0.76 because no color change was noticed for pure anatase TiO2 (Figure S5). Figure 4c shows the UV−vis absorption spectra of anatase Ti0.69□0.33O0.76F0.48(OH)0.76 before and after the surface functionalization with NMP molecules. The sample treated with NMP shows an additional absorption feature within the visible range, that is, ~500 nm, in agreement with its yellow color. Moreover, a plot of (ahv)1/2 versus hv (see inset of Figure 4c) using the modified Kubelka−Munk method reveals a slight decrease of the band gap energy from 3.12 to 3.03 eV and a second feature occurring at 2.25 eV.

Elemental analysis was performed at the level of an aggregate of nanoparticles of NMP-Ti1−x□xO2-4(xy)F4x(OH)4y (Figure S3a). Elemental mapping (Figure Sb−e) obtained by energy-dispersive X-ray spectroscopy (EDX) in a scanning transmission electron microscope revealed the presence of nitrogen atoms present in the whole area of the aggregate, suggesting a homogeneous grafting of NMP molecules at the surface of the nanoparticles.

The effective grafting of NMP molecules was also confirmed by X-ray photoelectron spectroscopy (XPS) analysis. The XPS spectrum of NMP-Ti1−x□xO2−4(xy)F4x(OH)4y (Figure 6) shows the appearance of a N 1s peak centered at 400.5 eV, which is characteristic of pyrrole nitrogen atom in NMP.41,42 2.2.1. Structure and Spectroscopic Analysis of NMP-Functionalized Anatase Ti1−x□xO2−4(xy)F4x(OH)4y. To probe the structural impact of the NMP functionalization, we first performed XRD analysis which confirms that the long-range order remains intact after the NMP treatment (Figure S2). Subsequently, we used the PDF obtained by Fourier transformation of synchrotron-based XRD data to analyze any structural deviation occurring at the short-range order. Figure 7 shows the redefined PDF patterns of Ti0.69□0.33O0.76F0.48(OH)0.76 and NMP-Ti1−x□xO2−4(xy)F4x(OH)4y samples. The extracted structural parameters are gathered in Table S1. Both anatase samples showed similar unit cell parameters and a close coherence length of ca. 4 nm, suggesting that the long-range order was not modified upon functionalization. The refinement of the Ti site occupancy, however, shows a slight decrease of the titanium vacancy with site occupancies of ca. 69(3)% for pristine Ti0.69□0.33O0.76F0.48(OH)0.76 sample and 75(2)% for NMP-Ti1−x□xO2−4(xy)F4x(OH)4y sample, indicating a
modification of the chemical composition, that is, a decrease of F (x) and/or OH (y) contents.

The treatment with NMP results in a substantial decrease of the fluorine wt % (from ~13.3 to ~9.3 wt %) measured by $^{19}$F solid-state NMR. According to the general chemical formula $\text{Ti}_{1-x-y} \square_{x+y} \text{O}_{2-4(x+y)} \text{F}_{4x} \text{(OH)}_{4y}$, the pristine $\text{Ti}_{0.69} \square_{0.31} \text{O}_{0.76} \text{F}_{0.48} \text{(OH)}_{0.76}$ becomes $\text{Ti}_{0.75} \square_{0.25} \text{O}_{1.00} \text{F}_{0.34} \text{(OH)}_{0.66}$ after the NMP treatment. This may be explained by the hydrolysis of water, yielding a partial defluorination of the network according to $\text{Ti}^- + \text{H}_2\text{O} \rightarrow \text{Ti}^0 + \text{H}_2\text{O}$, thereby decreasing the OH content.

The change in the chemical composition was confirmed by XPS analysis. Atomic percentage analysis of $\text{Ti} 2p_3/2$, $\text{O} 1s$ (that includes $\text{Ti}^0$ and $\text{Ti}^1$-OH contributions) and $\text{F} 1s$ peak was performed for $\text{Ti}_{0.69} \square_{0.31} \text{O}_{0.76} \text{F}_{0.48} \text{(OH)}_{0.76}$ and NMP-$\text{Ti}_{0.75} \square_{0.25} \text{O}_{1.00} \text{F}_{0.34} \text{(OH)}_{0.66}$. The atomic percentages of $\text{Ti}$/$\text{O}$/$\text{OH}$/$\text{F}$ change from 30/40/5/25 for $\text{Ti}_{0.69} \square_{0.31} \text{O}_{0.76} \text{F}_{0.48} \text{(OH)}_{0.76}$ to 30/48/3/19 for NMP-$\text{Ti}_{0.75} \square_{0.25} \text{O}_{1.00} \text{F}_{0.34} \text{(OH)}_{0.66}$, confirming changes in the sample stoichiometry induced by the treatment.

Figure 5. (a) STEM image at low magnification coupled with elemental mapping EDX (shown in (b) green for Ti, (c) blue for O, (d) red for F, and (e) violet for N element) of the NMP-$\text{Ti}_{1-x-y} \square_{x+y} \text{O}_{2-4(x+y)} \text{F}_{4x} \text{(OH)}_{4y}$ sample.

Figure 6. High-resolution N 1s XPS spectra of anatase $\text{Ti}_{0.69} \square_{0.31} \text{O}_{0.76} \text{F}_{0.48} \text{(OH)}_{0.76}$ before and after NMP treatment.

Figure 7. PDF refinements of anatase $\text{Ti}_{0.69} \square_{0.31} \text{O}_{0.76} \text{F}_{0.48} \text{(OH)}_{0.76}$ and NMP-$\text{Ti}_{0.75} \square_{0.25} \text{O}_{1.00} \text{F}_{0.34} \text{(OH)}_{0.66}$.

Figure 8. $^{19}$F solid-state MAS (64 kHz) experimental (black) and calculated (red) NMR spectra of $\text{Ti}_{0.69} \square_{0.31} \text{O}_{0.76} \text{F}_{0.48} \text{(OH)}_{0.76}$ and NMP-$\text{Ti}_{0.75} \square_{0.25} \text{O}_{1.00} \text{F}_{0.34} \text{(OH)}_{0.66}$. The asterisks indicate the main spinning side bands. The individual contributions used for the fits are shown below the fits, and parameters used for these fits are given as the Supporting Information (Tables S2 and S3).
T1−F at −87 ppm, Ti2−F at −4 ppm, and Ti3−F at 76 ppm.\textsuperscript{20} The \textsuperscript{19}F MAS NMR spectrum of NMP-Ti0.75\textsubscript{2−}O\textsubscript{1.00}F\textsubscript{0.34}(OH)\textsubscript{0.76} differs from those of the pristine sample. The three main lines are broader and those assigned to Ti2−F and Ti3−F are asymmetric (and are then fitted with two contributions). The fluorine environments are then even more disordered in NMP-Ti0.75\textsubscript{2−}O\textsubscript{1.00}F\textsubscript{0.34}(OH)\textsubscript{0.76} than in Ti0.69\textsubscript{2−}O\textsubscript{0.76}F\textsubscript{0.46}(OH)\textsubscript{0.76}. Because of the decrease of the titanium vacancy rate, the proportion of Ti3−F decreases and those of Ti2−F remains constant. An increase of the proportion of Ti1−F species is expected but with a lower extent than observed. Indeed, despite the partial defluorination, the sample contains a larger quantity of Ti1−F species after the treatment. This further highlights that the treatment is accompanied by a local structural rearrangement. Finally, a resonance appears at about 2.3 ppm. This new contribution is assigned to the H atoms of NMP in accordance with their \textsuperscript{1}H δH values in solution ranging from 2.0 to 3.4 ppm (Table S5). Second, the NMR signal of the Ti2−OH species vanishes or, more likely, is substantially reduced and hidden by the broad contribution of NMP. The decrease of the quantity of Ti2−OH species cannot be explained solely by the decrease of the titanium vacancy rate. Then, it can be assumed that they are preferentially involved in the oxolation reactions.

To probe the presence of PCs in the NMP-Ti0.75\textsubscript{2−}O\textsubscript{1.00}F\textsubscript{0.34}(OH)\textsubscript{0.76} sample, we performed EPR spectroscopy (Figure 10). Similarly, to the pristine compound, under dark condition, the EPR spectrum indicates the absence of Ti\textsuperscript{III}. However, weak EPR signals appear in the range of 3300–3340 G, characteristic of O\textsuperscript{−} radicals,\textsuperscript{47} indicating that the NMP treatment induces changes in reactivity even under dark.

Under illumination, we observed significant changes on the EPR spectrum features. The previous signal at around 3300–3340 G, characteristic of O\textsuperscript{−} radicals, significantly increases. Moreover, similarly to the pristine compound, an intense signal at 3350–3370 G is assigned to Ti\textsuperscript{III} (g\textsubscript{∥} = 1.9845 and g\textsubscript{⊥} = 1.958). The line width of this signal reflects that these PCs are isolated and mainly localized in the bulk.\textsuperscript{32} A quantitative assessment of the Ti\textsuperscript{III} and O\textsuperscript{−} radicals shows that the NMP treatment significantly modifies the reactivity of anatase. Indeed, the content of Ti\textsuperscript{III} centers is 6 × 10\textsuperscript{16} spin/g for Ti0.69\textsubscript{2−}O\textsubscript{0.76}F\textsubscript{0.46}(OH)\textsubscript{0.76} and 5 × 10\textsuperscript{17} spin/g for NMP-Ti0.75\textsubscript{2−}O\textsubscript{1.00}F\textsubscript{0.34}(OH)\textsubscript{0.76}.

The O\textsuperscript{−} radical content further grows under illumination from ~10\textsuperscript{14} spin/g for Ti0.69\textsubscript{2−}O\textsubscript{0.76}F\textsubscript{0.46}(OH)\textsubscript{0.76} to 3 × 10\textsuperscript{17} spin/g for NMP-Ti0.75\textsubscript{2−}O\textsubscript{1.00}F\textsubscript{0.34}(OH)\textsubscript{0.76}. Most strikingly, the Ti\textsuperscript{III}/O\textsuperscript{−} ratio strongly decreases after the NMP treatment, going from 600 (=6 × 10\textsuperscript{10}/3 × 10\textsuperscript{17}) to 1.67 (=5 × 10\textsuperscript{7}/3 × 10\textsuperscript{10}). This indicates that the treatment not only increases the concentration of Ti\textsuperscript{III} but also most importantly seems to limit the electron–hole recombination. Indeed, the photoexcited electrons are captured by oxygen molecules and Ti\textsuperscript{III} centers, leading to a space separation of the charge carriers and prevents their recombination with photoexcited holes. We do not detect photoexcited holes by the EPR technique, which is probably due to their very short relaxation times and, correspondingly, to the large width of the EPR signal line.

2.2.2. Photoelectrochemistry on NMP-Functionalized Photoanodes. Photoelectrochemical measurements were
As it was observed for Ti 0.69 measurement, under light, indicated the formation of Ti III hole recombination rate is low compared to anatase TiO2. Ti0.75 photocurrent onset potential of NMP-photocurrent. Additionally, one should notice that the shows a weak contribution in the visible range up to ca. 475 of a band gap of 2.97 eV with a second threshold featured at Ti 0.69 intermittent light (Figure 11b) have the same behavior for Δ of these values with anatase TiO2 shows a total potential shift negative potential, that is, −0.16 V versus NHE. A comparison with the results obtained by UV alization and a chemical composition change. These two e effects on the band gap energy. The ionocovalent character of Ti–X (X = F, OH) bonds is dictated by the speciation of these species with Ti–X bonds being more covalent in Ti0.31O0.76F0.48(OH)0.76 than in Ti0.31X−X. This point should be more carefully investigated. The second effect observed after the NMP treatment is the appearance of a broad absorption band within the visible range and responsible of the yellow color, leading to a reduced band gap energy at 2.25–2.50 eV. We assigned this broad absorption band to the effective grafting of the electron-donating NMP molecules expected to reduce the band gap energy. 48 Hence, NMP-Ti0.75□0.25O□1.00□F□0.34□(OH)□0.66 can be better described by a core–shell structure that synergistically favors the formation and separation of electron–hole carriers, similar to the previous report activating anatase TiO2 with a coated carbon shell, producing the same color features and increased photoelectrochemical activities. 49 It should be noted that after several photoelectrochemical tests, the coloration of the powder did not evolve, suggesting that NMP molecules are strongly bound to the surface.

3. CONCLUSIONS

In this work, we report on the optical and photoelectrochemical properties of anatase featuring an oxy-hydroxyfluorinated network with titanium vacancies (Ti0.69□0.31O□0.76F□0.48(OH)□0.76). Compared to pure TiO2, this compound showed a red shift of 0.08 eV assigned to changes in the chemical composition, particularly to the partial defluorination of the network and possibly to the modification of the proton environments. The speciation, that is, the number of neighboring cations, of OH groups and fluoride can also induce different effects on the band gap energy. The ionocovalent character of Ti–X (X = F, OH) bonds is dictated by the speciation of these species with Ti–X bonds being more covalent in Ti0.31X−X. This point should be more carefully investigated. The second effect observed after the NMP treatment is the appearance of a broad absorption band within the visible range and responsible of the yellow color, leading to a reduced band gap energy at 2.25–2.50 eV. We assigned this broad absorption band to the effective grafting of the electron-donating NMP molecules expected to reduce the band gap energy. 48 Hence, NMP-Ti0.75□0.25O□1.00□F□0.34□(OH)□0.66 can be better described by a core–shell structure that synergistically favors the formation and separation of electron–hole carriers, similar to the previous report activating anatase TiO2 with a coated carbon shell, producing the same color features and increased photoelectrochemical activities. 49 It should be noted that after several photoelectrochemical tests, the coloration of the powder did not evolve, suggesting that NMP molecules are strongly bound to the surface.

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gifting of the electron-donating NMP molecules, which induces the appearance of a new absorption band in the visible range corresponding to an estimated band gap of 2.25−2.50 eV. Hence, the as-prepared composite material showed enhanced photoelectrochemical properties. Such an increasing performance was ascribed to an increasing ability to generate and separate photogenerated charge carriers. This work provides new opportunities in the development of high-performance photocatalysts.

4. EXPERIMENTAL SECTION

4.1. Synthesis of Ti$_{1-x}$O$_{2-4x}$F$_{4x}$(OH)$_{8y}$ and NMP - Ti$_1$ - x - y O$_{2-4x}$F$_{4x}$(OH)$_{8y}$ and NMP - Ti

In short, 1.2 mL of HF was added to a mixture solution containing 24.8 mL of isopropanol and 4 mL of titanium isopropoxide in a 45 mL Teflon line container. The Teflon line container was loaded into a stainless steel autoclave. After sealing the autoclave, the solution was heated at 90 °C for 12 h, and the white powder was recovered from the solution. The recovered powder was dried at 80 °C overnight. Such a low temperature was used to avoid the thermal dehydroxylation of the compound, enabling to preserve a high vacancy concentration.

Anatase functionalized with NMP was obtained by stirring 500 mg of the as-prepared Ti$_{1-x}$O$_{2-4x}$F$_{4x}$(OH)$_{8y}$ in 20 mL of NMP at room temperature for 7 days. A yellow powder was recovered from the solution using centrifugation and washed several times with ethanol. The solid was dried at 80 °C overnight.

For the sake of comparison, anatase TiO$_2$ was synthesized without HF followed by a post-heating treatment at 300 °C under air for 2 h to obtain crystallized anatase TiO$_2$ (Figure S2). The heating temperature was adjusted to obtain pure TiO$_2$ with particle sizes close to that of anatase TiO$_2$ (Table S1).

4.2. Physical Characterizations. XRD measurements were carried out on a Rigaku diffractometer using Cu Kα radiation in a Bragg–Brentano geometry.

TGA was performed on a METTLER TOLEDO thermogravimetric analyzer under argon atmosphere at a heating rate of 5 °C min$^{-1}$ from room temperature to 600 °C.

UV–visible adsorption spectra were recorded using an Agilent Technologies Cary Series spectrophotometer.

Scanning transmission electron microscopy (STEM)−EDX measurement was performed using FEI-TITAN 80−300 kV with ChemiSTEM capability (Cs probe corrector and extreme field emission gun (X-FEG)).

X-ray scattering measurements were performed for PDF analysis at the 11-ID-B beam line at the Advanced Photon Source of Argonne National Laboratory. Similar to previous reports, high-energy X-rays ($\lambda = 0.2128$ Å) were used to collect data yielding high values of momentum transfer $Q \approx 22$ Å$^{-1}$. By the integration of the diffraction images, one-dimensional diffraction data were achieved. After the correction of the background and the Compton scattering, the $G(r)$ function was extracted from the data using PDFgetX2. The PDF data were refined with the PDFgui software, and the refined parameters were the scale factor, $s$ ratio, sp diameter (i.e., particle diameter), lattice parameters, oxygen atomic position, titanium (4a Wyckoff site) rate occupancy, and isotropic atomic displacement parameters.

$^1$H and $^{19}$F solid-state magic angle spinning (MAS) NMR experiments were performed in the same way (spectrometer, probe head, pulse sequences, and references) as previously employed for Ti$_{1-x}$O$_{2-4x}$F$_{4x}$(OH)$_{8y}$. $^1$H and $^{19}$F spectra were fitted by using the DMFit software. $^{19}$F solid-state NMR was also used to quantify the fluorine content on the studied samples by using reference samples.

XPS characterization was performed with a VG ESCALAB 250 spectrometer at the operating pressure of 3 × 10$^{-9}$ mbar using a monochromatic Al Kα radiation ($h\nu = 1486.6$ eV) as the X-ray source, as described in the previous report. The energy of spectrometer was calibrated with reference to the binding energy (BE) of clean Au sample (Au 4f$_{7/2}$ at 84 eV). The pass energy of the analyzer was 100 eV for survey and 20 eV for the high-resolution spectra (Ti 2p, C 1s, O 1s, N 1s, and F 1s). The BE of peaks was calibrated by setting the C 1s signal corresponding to −CH$_3$−CH$_2$− bonds at 285.0 eV. The Advantage software (version 5.954) was employed to perform a peak fitting using a Shirley background and Gaussian/Lorentzian line shape with a ratio of 70/30.

EPR spectra were recorded by the standard Bruker EPR spectrometer ELEXYSYS-500 (X-band, sensitivity is around ~10$^{10}$ spin/G). To illuminate the samples directly in the spectrometer, the optical cavity was used. The samples were positioned in flat ampoules. Etalon CuCl$_2$-2H$_2$O was used for the calculation of PC concentration. Mn$^{2+}$ in MgO was employed as a reference for g-values. The samples were illuminated in situ (directly in the EPR cavity) with a high-pressure tungsten lamp in the spectral range 250−1000 nm. The intensity of the illumination was 40 mW/cm$^2$. The measured temperature was 77 K. Experiments at liquid nitrogen temperature were carried out using a finger cryostat.

4.3. Photoelectrochemical Characterizations. 4.3.1. Photoanodes. A suspension containing 10 mg of active powder and 2 mL of ethanol was prepared. The mixture was ultrasonically stirred for 2 h to ensure a homogeneous dispersion; after that, 300 μL of suspension was sprayed via an airbrush over the conductive SnO$_2$:F (YSUB/ASASHI120/1010 spin/G). To illuminate the samples directly in the spectrometer, the optical cavity was used. The samples were positioned in flat ampoules. Etalon CuCl$_2$-2H$_2$O was used for the calculation of PC concentration. Mn$^{2+}$ in MgO was employed as a reference for g-values. The samples were illuminated in situ (directly in the EPR cavity) with a high-pressure tungsten lamp in the spectral range 250−1000 nm. The intensity of the illumination was 40 mW/cm$^2$. The measured temperature was 77 K. Experiments at liquid nitrogen temperature were carried out using a finger cryostat.

4.3.2. Photoelectrochemical Measurements. Current–potential curves were measured in a three-electrode photoelectrochemical cell with a working Ti-based porous layer photoanodes, a Pt wire counter electrode, and an Ag/AgCl reference electrode. The photoelectrochemical measurements were conducted in an argon-purged 1 M Na$_2$SO$_4$ electrolyte under UV–vis radiation. A shutter was used during these experiments. The measured potentials versus Ag/AgCl were converted to the normal hydrogen electrode (NHE) according to the following equation: $E$ versus NHE $= E$ versus Ag/AgCl + 0.197 V. IPCE action spectra were recorded at 1.0 V/NHE under UV–vis illumination at 300−550 nm with a 150 W xenon lamp (Spectral Products, ASB-XE-175SX) with an incident irradiation power at the electrode of 34 mW/cm$^2$. Note that the lower light intensity was used in the present study, generating lower photocurrent, compared to the commonly used light intensity (100 mW/cm$^2$) in the literature. Generally speaking, the photocurrent increases with the light intensity. Indeed, solar light between 400 and 700 nm with 2−4% UV is called AM 1.5, with ca. 100 mW/cm$^2$, which is essential for solar cells. In this study, the incident
power density is only essential to calculate the IPCE by applying the light source photoemission correction. All aqueous solutions were prepared using 18.2 MΩ-cm Millipore Milli-Q water, and all measurements were conducted with illumination of the film–electrolyte interface through a 0.19 cm² mask at the working electrode in the UV–vis light range.

## ASSOCIATED CONTENT

### Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsomega.9b01219.

XRD patterns, visual appearances, TGA curves, and individual contributions used for the fits of 19F and 1H MAS NMR spectra (PDF)

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### Notes

The authors declare no competing financial interest.

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