Visible-Light-Driven Water Oxidation on Self-Assembled Metal-Free Organic@Carbon Junctions at Neutral pH

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ABSTRACT: Sustainable water oxidation requires low-cost, stable, and efficient redox couples, photosensitizers, and catalysts. Here, we introduce the in situ self-assembly of metal-atom-free organic-based semiconductive structures on the surface of carbon supports. The resulting TTF/TTF⁺@carbon junction (TTF = tetrathiafulvalene) acts as an all-in-one highly stable redox-shuttle/photosensitizer/molecular-catalyst triad for the visible-light-driven water oxidation reaction (WOR) at neutral pH, eliminating the need for metallic or organometallic catalysts and sacrificial electron acceptors. A water/butyronitrile emulsion was used to physically separate the photoproducts of the WOR, H⁺ and TTF, allowing the extraction and subsequent reduction of protons in water, and the in situ electrochemical oxidation of TTF to TTF⁺ on carbon in butyronitrile by constant anode potential electrolysis. During 100 h, no decomposition of TTF was observed and O₂ was generated from the emulsion while H₂ was constantly produced in the aqueous phase. This work opens new perspectives for a new generation of metal-atom-free, low-cost, redox-driven water-splitting strategies.

KEYWORDS: water oxidation, hydrogen evolution, photochemistry, sustainable, redox shuttles

INTRODUCTION

The increase of the global energy demand and the negative environmental impact triggered by the overexploitation of fossil fuels accentuate the need for the development of clean, efficient, and sustainable energy technologies.¹⁻¹³ Hydrogen is rapidly becoming the energy vector of choice, and advanced water electrolysis is one of the most efficient approaches to produce high-quality green H₂ from renewable energy sources despite their intermittencies.¹⁻¹³ To carry out what is often referred as artificial photosynthesis to produce hydrogen and oxygen directly from water, one usually combines an antenna such as a dye to absorb the light, a semiconductor to carry out the charge separation, and some electrocatalysts to drive both water reduction and the water oxidation reaction (WOR).¹⁻¹⁸ As of today, the photosensitized oxidation of water to oxygen under visible-light irradiation is still the bottleneck of artificial photosynthesis hindering large-scale developments. Light-driven water oxidation on semiconductor photoelectrodes has been extensively studied for nearly five decades with slow progress due to the difficulty of finding electrode materials exhibiting high chemical stability, suitable optoelectronic properties, and high catalytic efficiencies. Materials, such as titanium dioxide, absorbs only a very small part of the solar spectrum.¹⁻¹⁸ The sensitization of semiconductors has been significantly improved by using a wide range of metal complexes, organic dyes, and porphyrins.⁴⁻¹²,¹⁴,¹⁵,¹⁹⁻²⁶ Photocatalytic junctions based on BiVO₄, Fe₃O₄, WO₃, and Ag₃PO₄ among others, have been widely studied for water oxidation in both suspensions and photoelectrochemical devices.⁸,¹⁰,²⁷⁻³³ Metal–molecule interactions play a key role in the water oxidation reaction.³⁴⁻³⁶ However, the need for sacrificial electron acceptors such as Ag⁺, Fe³⁺, and peroxodisulfate (S₂O₈²⁻) restricts its scalability.³⁷ Silicon and III–V compounds (GaAs, GaP, etc.) exhibit excellent optoelectronic properties but due to their instability under illumination in aqueous solutions, they require high-quality protective layers to prevent photocorrosion.²

On the molecular side, many organometallic compounds have been proposed as antennae and redox catalysts, but none have yet found industrial applications.¹⁷ Nonetheless, it can also be noted that some fully organic compounds have been found to be electrocatalytically active for water oxidation on glassy-carbon or Pt working electrodes in the dark.³⁸⁻⁴⁰

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Due to their strong self-assembly properties, the radical cation of the organic electron donor tetraphiafulvalene (TTF\(^+\)) and its derivatives are frequently used as building blocks for the synthesis of highly electrically conducting structures.\(^{34,41-43}\) Such properties have been extensively exploited to develop organic conductors, superconductors, photovoltaic cells, and solar cells, among others.\(^{41-45}\) Metal–ligand coordination strategies have been widely used to create self-assembled three-dimensional TTF-based architectures where the TTF\(^+\) sites enhance conductivity. Indeed, TTF-based metal–organic frameworks (MOFs) show an extended charge transport pathway within the rigid MOF structure,\(^{48,49}\) exhibiting semiconductive properties comparable to that of common organic semiconductors.\(^{48,49}\)

Some examples include Zn\(_2\)(TTFTB) (10\(^{-6}\) S cm\(^{-1}\)), Co\(_3\)(TTFTB) (10\(^{-6}\) S cm\(^{-1}\)), Mn\(_3\)(TTFTB) (10\(^{-3}\) S cm\(^{-1}\)), and Cd\(_3\)(TTFTB) (10\(^{-4}\) S cm\(^{-1}\)).

To overcome the efficiency, scale-up, and cost-related roadblocks of the conversion of solar energy into H\(_2\) through photosensitized water splitting, the development of low-cost and highly stable photosensitizers/redox shuttles and photocatalysts is crucial. Furthermore, a system operating with water at neutral pH and without ion-exchange membranes is highly desirable.

Here, we introduce an efficient and highly stable visible-light-driven WOR by in situ self-assembled semiconductive tetraphiafulvalene-based structures (TTF/TTF\(^+\)) deposited on a solid carbon support. A waterbutyronitrile emulsion was used as a platform for the spontaneous separation of the generated lipophilic TTF from the generated hydrophilic protons, with O\(_2\) evolving as a gas. Such a separation allows the subsequent effective reduction of the protons to H\(_2\) and the electrochemical recycling of the electron acceptor TTF\(^+\).

## EXPERIMENTAL SECTION

### Synthesis of TTF/TTF\(^+\)BF\(_4\)^{−}\)

TTF/TTF\(^+\)BF\(_4\)^{−} was synthesized entirely inside a glovebox with nitrogen by the stoichiometric oxidation of TTF (99%, Acros, structure in Scheme 1) with dry nitrosonium tetrafluoroborate (NOBF\(_4\), 98%, Alfa Aesar) in dry butyronitrile (butyronitrile, 99.9%, Acros). Prior to the reaction, butyronitrile was degassed in a Schlenk line and dried with molecular sieves (3 Å, Fisher Chemical) during 3 days inside of a glovebox. For the photochemical experiments, an excess of 5% of TTF was used in the synthesis of TTF\(^+\) in order to consume any excess of oxidant, therefore avoiding oxidation of TTF\(^+\) to TTF\(^{++}\). The resulting product in solution (Figure SI1) is further referred to as TTF/TTF\(^+\)BF\(_4\)^{−}. After the completion of the reaction, the solution was thoroughly degassed inside of a glovebox in order to evacuate the NO released during the reaction. The TTF/TTF\(^+\)BF\(_4\)^{−} solution was always kept inside the glovebox and protected from the light.

### Cyclic Voltammetry (CV)

A three-electrode configuration composed of a commercial 3 mm diameter glassy-carbon electrode as the working electrode, Pt wire as the counter electrode, and an Ag/AgCl/3 M KCl double-junction reference electrode was used to analyze by CV 1 mM of the synthesized TTF/TTF\(^+\)BF\(_4\)^{−} in 50 mM tetrabutylammonium tetrafluoroborate (TBABF\(_4\), 99%, Acros) in butyronitrile. The scan rate was 10 mV s\(^{-1}\). The CV (Figure SI1) was recorded inside a glovebox with a Metrohm Autolab PGSTAT 302N (Herisau, Switzerland) potentiostat.

For the electrochemical recycling of TTF\(^+\) depicted in Scheme 2, CVs were recorded before and after the recycling to determine and verify the potential to be applied at the anode (WE). A glassy-carbon electrode (D = 3 mm) was used as the WE, and the scan rate was 25 mV s\(^{-1}\).

### Visible-Light-Driven Water Oxidation by TTF\(^+\)@GF and TTF\(^+\)@RVC Junctions upon In Situ Electrochemical Recycling of TTF\(^+\) and Continuous Reduction of Protons by V\(^{2+}\)/Mo\(_2\)C

The visible-light-driven WOR by TTF/TTF\(^+\)@carbon and the in situ electrochemical regeneration of TTF\(^+\) were performed in the anodic compartment of the H-cell depicted in Scheme 2 (left side). The anode was illuminated with a 455 nm LED, and stirred inside of a glovebox filled with N\(_2\). The protons resulting upon the WOR were spontaneously transferred to the aqueous phase and then to the cathodic compartment where they were reduced by V\(^{2+}\)/Mo\(_2\)C (Scheme 2, right side) by redox electrocatalysis.\(^{36}\) The two septum-sealed compartments of the H-cell were separated by a glass wool plug, to avoid the mixing of TTF\(^+\) and VCl\(_2\). The glass wool plug ensured that the cathodic compartment was purely aqueous and that of the anodic compartment was a biphasic system kept as an emulsion during vigorous stirring.

The anodic compartment (left): organic phase (butyronitrile), 1 mM TTF/TTF\(^+\)BF\(_4\)^{−} in 100 mM TBABF\(_4\); aqueous phase, pure water. Anode, RVC or GF. Cathodic compartment (right): 100 mM VCl\(_2\) in water, neutral pH, 40 mg of Mo\(_2\)C microparticles. Cathode: Pt mesh. The contents of the septum-sealed compartments were vigorously stirred, and the anodic compartment was illuminated with a 455 nm LED. Constant anode potential electrolysis of TTF: 0.6 V vs an Ag/AgCl/3 M KCl double-junction reference electrode. Bridge: glass wool plug to avoid butyronitrile from entering the cathodic compartment, while the two aqueous phases were kept in contact.
the emulsion was initially neutral. Two different carbon anodes (working electrode (WE)) were studied independently: reticulated vitreous carbon (RVC, Duocel) and graphite foil (GF, high-purity flexible graphite foil, SIGRACLAD).

The cathodic compartment (right) contained 16 mL of 100 mM VCl2 in water at neutral pH and 40 mg of Mo2C microparticles (<10 μm). A Pt mesh (Sigma-Aldrich) was used as the cathode (counter electrode (CE)).

The reference electrode was an Ag/AgCl/3 M KCl double-junction electrode connected to the anodic compartment. The recycling potential of TTF+ was set at 0.6 V vs the Ag/AgCl/3 M KCl double-junction reference electrode.

The gas products of the visible-light-driven WOR on the anodic side and the HER on the cathodic side were analyzed every 15 min by gas chromatography (GC-TCD-MS). Electrochemical impedance spectroscopy (EIS) under illumination was also performed every 15 min. To avoid overpressure of the system upon gas evolution, which in fact has a detrimental effect in the progress of the WOR and the HER, the recycling and illumination were paused after each GC and EIS analysis, and the cell was opened and degassed with N2. Once the anode and the cathode did not show remnant O2 or H2, respectively, the cell was septum-sealed again and the WOR/electrochemical recycling/proton reduction was resumed.

Three control experiments were performed following the same methodology:

Control 1. without TTF/TTF**BF4− in the anodic compartment but with V2+/Mo2C in the cathodic compartment; GF as the anode and Pt mesh as the cathode.

Control 2. with TTF/TTF**BF4− in the anodic compartment but without V2+/Mo2C in the cathodic compartment; GF as the anode and Pt mesh as the cathode.

Control 3. carbon electrode replaced by platinum mesh with TTF/TTF**BF4− in the anodic compartment and with V2+/Mo2C in the cathodic compartment.

For all three controls, the reference electrodes and all the other conditions were the same as for the system in Scheme 2. After every experiment the glass H-cell was washed with water, acid, and isopropanol. The carbon electrodes were new for each experiment.

Gas Chromatography (GC)
The headspaces of both septum-sealed compartments of the H-cell shown in Scheme 2 were sampled every 15 min by using a lock-in syringe with a push—pull valve (SGE Analytical Sciences). The gas was then injected into a TRACE 1300 GC equipped with a thermal conductivity detector (TCD) and an ISQ single quadrupole (mass spectrometry detector, MSD) from Thermo Fisher Scientific (GC-TCD-MSD). The chromatographic column was 5 Å molecular sieves, 80/100 mesh. Oxygen was detected with the MSD, using helium as the carrier gas, and H2 was detected with the TCD, using argon as the carrier gas. The instrument was calibrated for O2 and H2 by using 8.0% carrier gas. The instrument was calibrated for O2 and H2 by using 8.0% O2 and 8.0% H2 in N2 or H2 in N2 standards (99%, Carbages). Prior to injection, the injection line was purged with N2 and the exit of the line was connected to a water trap in order to avoid suction of air during the injections.

Electrochemical Impedance Spectroscopy (EIS)
Each EIS spectrum was recorded after every 15 min of electrolysis of TTF to TTF**. Prior to impedance spectra recording, stirring was turned off such that the electrode was immersed only in the organic phase. Stirring was then resumed to continue the electrolysis. Conditions: 10 mV amplitude perturbation, 100 kHz to 100 mHz, frequency step 10 points per decade. All experiments were recorded in the dark and under visible-light illumination at 0.6 V vs an aqueous Ag/AgCl/3 M KCl reference electrode.

More details of the methods are given in the Supporting Information.

## RESULTS AND DISCUSSION

In 2018, we demonstrated that the in situ self-assembly of tetrathiafulvalene (TTF), TTF**, and BF4− or PF6− on the surface of Pt microparticles in wet acetonitrile is active for the WOR upon visible-light irradiation. In short, we had demonstrated that TTF**+ is highly oxidizing and that the molecular assembly TTF/TTF**BF4−, once photoexcited, can provide the four electrons required to oxidize water when it is dissolved in an organic solvent. However, this system is short-lived because it suffers from a gradual increase in the proton concentration in the wet organic phase and subsequent protonation of TTF in the molecular assemblies, triggering competing reactions such as proton and oxygen reduction and preventing the electrochemical recycling of TTF back to TTF**.

In this work, we eliminate the use of metallic catalysts for the WOR, replacing them with a low-cost, and highly stable, TTF/TTF**BF4−/carbon junction, which works as an all-in-one redox-shuttle/photosensitizer/molecular-catalyst triad for the visible-light-driven WOR at neutral pH. Thus, we take advantage of the possibility of adjusting the Gibbs energy transfer across a polarized waterbutyronitrile interface, to separate in situ and in a spontaneous manner the hydrophilic protons from the lipophilic TTF formed upon the visible-light-driven WOR by TTF/TTF**BF4− assemblies. The goal of such a separation is to prevent TTF from being protonated and therefore allow its electrochemical (in situ) reoxidation to perform continuous photosensitized WOR. Subsequently, the protons generated are chemically consumed in the cathodic compartment by redox electrocatalysis. Indeed, the standard redox potential of the V3+/V2+ couple is −0.26 V and is therefore reducing enough to generate hydrogen on an electrocatalyst such as molybdenum carbide. Here, the strategy is to set up a “proton pump” to extract the photogenerated protons in the anodic compartment toward the cathodic compartment, where they are directly reduced to produce hydrogen. The reason to choose the V3+/V2+ couple lies in the long experience that our group has in producing hydrogen on a very large scale using a vanadium redox flow battery (200 kW–400 kWh). However, on consideration that our long-term objective is to develop a fully environmentally friendly WOR-HER system, the next step is to improve the HER part of the cell, replacing vanadium by a nonmetallic hydrogen evolution system, which is a work in progress in our laboratory.

Because the primary objective of this work is to demonstrate long-lived water photo-oxidation by TTF/TTF**BF4− assemblies on carbon electrodes without any metal catalyst, we chose to operate the electrochemical regeneration of TTF** from TTF at the carbon electrode by applying a constant anode potential using a potentiostat where the carbon electrode is the working electrode (see Scheme 2). The electrode in the aqueous compartment acts as a counter electrode, and its potential cannot be directly controlled. It is, of course, the locus of a reduction reaction such as the reduction of V3+ to V2+.

Scheme 2 schematically shows the H-cell used to test the visible-light-driven WOR by TTF/TTF**@RVC and TTF/TTF**@GF junctions (anodes), coupled to the in situ electrochemical regeneration of TTF** in the anodic compartment, and the reduction of protons by V2+/Mo2C in the cathodic compartment, using Pt mesh as a cathode.

Once the visible-light-driven WOR is triggered by illumination of the anodic compartment, the protons that formed are transferred to the aqueous phase of the emulsion, while TTF is transferred to the organic phase, as it is poorly soluble in water, the salt TTF**BF4− being soluble in both phases. As the two septum-sealed compartments are physically connected by an aqueous glass wool plug, the protons are extracted to the
cathodic compartment, where they are reduced by V²⁺/Mo₂C to produce H₂. The glass wool plug avoids a convective mixture of TTF** and VCl₂. Indeed, the transfer of VCl₂ to the anodic compartment results in the quenching of the visible-light-driven WOR and, therefore, in the inhibition of oxygen evolution and proton production, as the oxidation of V²⁺ by TTF** is thermodynamically much more feasible than the oxidation of water.

In accordance with the CV recorded prior to the experiment (Figure 1), the onset current of the electrochemical WOR on the carbon electrode is observed above 1.15 V, and the electrochemical oxidation of TTF is observed above 0.4 V vs carbon electrode is observed beyond 1.15 V, and the accumulated in the headspace of the anodic compartment (3 M KCl) basically no aqueous species, including water, butyronitrile phase and that at this potential (0.6 V vs Ag/AgCl/3 M KCl which is about 600 mV less positive than the Ag/AgCl/3 M KCl, which is about 600 mV less positive than the

Figure 1. The black plot shows the cyclic voltammogram of the organic phase of the anodic compartment shown in Scheme 2 (1 mM TTF**) prior to the experiment. The green plot corresponds to the CV obtained after 9 h of experiment for control 2: with TTF** in the anodic compartment but without V²⁺/Mo₂C in the cathodic compartment and GF as the anode and Pt mesh as the cathode. The reference electrode and all the other conditions were the same as for the system in Scheme 2. A glassy-carbon electrode (D = 3 mm) was used as the WE, with 50 mM TBABF₄ as the supporting electrolyte. Scan rate: 25 mV s⁻¹.

The anodic compartment and only H₂ was produced in the cathodic compartment and they do not cross to the adjacent compartment during the experiment (Figure S12).

Figure 3a compares the activity of TTF/TTF**@RVC and TTF/TTF**@GF junctions. The TTF/TTF**@RVC and the TTF/TTF**@GF junctions achieved an impressive Y₁₈,WOR = 0.5 × 10⁻²% and 2.2 × 10⁻²% efficiencies, respectively, accounting for a recycling of TTF** of 530 and 2200 times, respectively. These results prove that the TTF/TTF**@carbon systems alone act as a redox shuttle/photosensitizer/catalyst triad for the visible-light-driven WOR, without the need for any metallic catalyst.

As a control, a Pt mesh that yielded the same initial current as the GF electrode (Figure 3b) was also tested as an anode. This TTF/TTF**@Pt junction reached half of the efficiency of the TTF/TTF**@GF junction (Figure 3a). In addition, as shown in Figure 3b, the TTF/TTF**@GF junction is electrochemically stable for as long as 5 h, while the stability of the TTF/TTF**@Pt junction starts to decrease after 1 h of reaction.

The SEM-EDX analyses of the GF (Figure 4a–i) after constant visible-light-driven WOR and anode potential electrolysis of TTF show the self-assembly of micrometer-sized (0.3–2 µm) cauliflower-like arrays, composed by sulfur and fluorine (Figure 4b–i), indicating the self-assembly of TTF/TTF**BF₄⁻ on the surface of the GF. These arrays are likely to be the nuclei of the microtubes (0.1–0.5 µm) observed in Figure 4c–i, which are similar to the microtubes observed in our previous work on Pt microparticles in wet acetonitrile. The size of the microtubes is consistent with the smaller diameter of the seminal graphite filaments (ca. 0.3 µm, Figure 4a). Indeed, Favier et al. showed the slow electrocrystallization of TTF(Br)_x conductive microtubes on Pt nanoparticles deposited on a GF electrode, whose diameters were directly proportional to the diameter of the Pt particles.

The growth of conductive TTF/TTF** microtubes on the surface of the GF observed by SEM explains the decrease in resistance of the electron transfer through the GF observed by EIS as the visible-light-driven WOR and electrochemical recycling of TTF** advanced (Figure 5).

In contrast, for the control where a Pt mesh was used as the anode instead of carbon, the SEM-EDX analyses showed the growth of a TTF-based film on the surface of Pt (Figure 4j,k), which correlates with the increase in capacitance of the TTF/TTF**@Pt junction as the WOR advances (Figure 5). The EIS and SEM analyses show that the filamentary structure of the GF (electric conductivity 9.1 × 10⁵ S m⁻¹) offers countless micrometer-sized seeds for the growth of TTF-based arrays, explaining the better performance of the TTF/TTF**@GF junction in comparison with the flat Pt mesh that, although it exhibits a much higher conductivity (9 × 10⁵ S m⁻¹), does not offer the micrometer-sized seeds required to grow the TTF-based conductive arrays.

After 5 h of reaction, both the current recorded during the electrochemical recycling of TTF to TTF** (Figure 3b) and the efficiency of the visible-light-driven WOR by the TTF/TTF**@GF junction (Figure 3c) started to decrease due to passivation of the GF. At this point, we switched the light off and stopped the electrochemical process to replace the anode by a clean GF.
Once the light and the constant anode potential were turned back on, the current and efficiency recovered to their initial values (Figure 3b,c). The time depicted in Figures 3b,c corresponds to the time during which the light and the recycling were on. The used GF was then cleaned up with butyronitrile and used again, showing a total recovery of the activity. This process was repeated three times up to 16 h (Figure 3b), exhibiting a stable current and a decrease of 15% in the efficiency of the WOR ($Y_{H_2,WOR} = 1.1 \times 10^6$%), which is mainly due to the depletion of the $\gamma^2$-solution, which was also replaced twice during the 16 h reaction. The fact that the whole system can be turned off and on without evident loss of efficiency and stability is a competitive advantage if the system is to be connected to an intermittent supply of energy. The stability of the TTF/TTF$^{\ddagger}$@GF junction upon continuous visible-light WOR and in situ constant anode potential electrolysis of TTF was further tested up to 100 h. The results of this experiment showed the constant production of gases, a constant current, and a constant charge increase (Figure SI3) proving the high stability of the photocatalytic system, which could indeed run further, as it did not show signs of decomposition.
The main advantage of the electrochemical recycling of TTF+ over the use of sacrificial electron acceptors is that no waste products which could decompose and eventually quench and slow down the WOR are produced. Attempts of electrochemical recycling of other electron donors/acceptors have been unsuccessful due to their long-term instability. The UV absorption spectrum (Figure SI4) and MS (Figure SI5) show no evidence of decomposition of the TTF species, showing that the cell can operate on a long-term basis.

Figure 4, SEM analysis. (a) New GF cleaned with water and butyronitrile (ETD SE). (b–f) GF as the anode after 150 min of experiment (Scheme 2); T1D: bright area, higher atomic number (TTF); gray area, lower atomic number (carbon). (g) ETD SE: bright area, self-assembly; dark area, support (GF). (h) EDX composition: sulfur in red (TTF), (i) EDX: composition, fluorine in blue (BF4). (j) Pt mesh as the anode after 150 min of constant anode potential electrolysis (Scheme 2). T1D: bright area, higher atomic number (Pt); gray area, lower atomic number (TTF-based film). (k) ETD SE: bright area, covering film; dark area, Pt mesh.

Control 1 was made without TTF+ in the anodic compartment, with V2+/Mo2C in the cathodic compartment, and using GF as the anode and Pt mesh as the cathode (all other conditions were the same as for Scheme 2). This experiment produced neither O2 in the anodic compartment nor H2 in the cathodic compartment, proving that, without the TTF/TTF+ assemblies, the visible-light-driven WOR is not triggered in the system described in Scheme 2 and therefore no protons were produced to be reduced by V2+/Mo2C in the cathodic compartment, explaining the absence of H2.

Control 2 was made with TTF+ in the anodic compartment, without V2+/Mo2C in the cathodic compartment, and using GF as the anode and Pt mesh as the cathode (all other conditions were the same as for Scheme 2). This blank produced a small amount of O2 and H2 in the anodic and cathodic compartments, respectively, accounting for a recycling of TTF+ of only 35 times, which is far below the recycling achieved when V2+/Mo2C is used for the proton reduction (2200 times). In this blank, the protons obtained upon visible-light-driven WOR by the TTF/TTF+@GF junction in the anodic compartment were reduced by the Pt mesh alone in the cathodic compartment; however, the pH was not sufficiently low to trigger efficient electrochemical reduction of protons on the Pt mesh at the potential applied during the recycling (0.6 V). The CV recorded after control 2 was performed (green plot in Figure 1) shows the total protonation of TTF upon accumulation of the protons formed during the visible-light-driven WOR by TTF+ in the anodic compartment.

This explains why the recycling ceases. The oxidation potential of HTTF+ is close to 1.05 V vs the Ag/AgCl/3 M KCl reference electrode, which is too high in comparison to the potential set for the oxidation of the
photogenerated TTF to TTF$^\bullet\bullet$ (0.6 V). Therefore, the protonation of TTF also stops the anode potential electrolysis, the current dropping to almost zero after only 30 min of the experiment. This control proves that, under the conditions of the experiment, the chemical reduction of protons by Ve/MoC is much more efficient than that of Pt alone.

The mechanism of the WOR by TTF$^\bullet\bullet$ assembles at neutral pH (eqs 1 and 2) is the reverse mechanism of that previously proposed for the slow four-electron reduction of oxygen to water by TTF assembles in acidic pH (see ref 57). Indeed, in ref 57 it was computationally established that the self-assembly of TTF-TTFH$^+$ dimers could provide the four electrons required to reduce oxygen to water, which is in fact the opposite reaction to the four-electron oxidation of water to O$_2$ studied in this paper. $^{57}$ However, in ref 57 the self-assembly of TTF was not proved experimentally; therefore, this manuscript offers experimental proof of the computational mechanism proposed in 2011. $^{57}$ The mechanism proposed in ref 57 involves the helical tetramer [TTF$_4$H$_2$]$^{2+}$ formed by the interaction of two TTF-TTFH$^+$ dimers [TTF$_2$H$_2$]$^{2+}$ (eq 1), which delivers the four electrons needed for the reduction of oxygen to water: $^{57}$

$$4\text{TTF}^{\bullet\bullet} + 2\text{H}_2\text{O} \rightleftharpoons [\text{TTF}_4\text{H}...\text{HO}_2]^{2+} + 4\text{H}^+$$

(eq 1)

$$\rightleftharpoons [\text{TTF}_4\text{H}_2...\text{O}_2]^{2+} + 2\text{H}^+ \rightleftharpoons [\text{TTF}_4\text{H}_2]^{2+} + \text{O}_2 + 2\text{H}^+$$

$$+ 2\text{H}^+ \rightleftharpoons 4\text{TTF} + \text{O}_2 + 4\text{H}^+$$

(1)

Electrochemical recycling of TTF$^\bullet\bullet$ on GF:

$$4\text{TTF} \rightleftharpoons 4\text{TTF}^{\bullet\bullet} + 4e^-$$

(2)

Indeed, [TTF$_4$H$_2$]$^{2+}$ binds O$_2$ to form a [TTF$_4$H$_2...\text{O}_2$]$^{2+}$ intermediate that is converted into the more stable [TTF$_4$H...HO$_2$]$^{2+}$ complex (eq 1). $^{57}$ The highly reactive HO$_2^\bullet$ and TTF$^\bullet\bullet$ moieties of the [TTF$_4$H...HO$_2$]$^{2+}$ complex combine and dissociate from the tetramer and get the needed protons from the solution to form two molecules of water. $^{57}$ Counteranion--cation complexes such as (TTFH$^+$)$_2$(BF$_4^-$)$_2$ exhibit cation--anion electrostatic interactions rather than direct interactions between the two TTFH$^+$ units. $^{57}$ This paper offers evidence of the formation of such a TTF-based assembly.

# CONCLUSIONS

A waterbutyronitrile emulsion was successfully used as a platform for the spontaneous separation of the products of the visible-light-driven WOR by self-assembled TTF/TTF$^\bullet\bullet$@GF junctions at neutral pH. Such a separation allowed the successful and efficient electrochemical recycling of TTF back to TTF$^\bullet\bullet$ at 0.6 V vs. Ag/AgCl/3 M KCl in the organic phase, while simultaneously performing the molecular photosensitized WOR. Moreover, the protons partitioned to the aqueous phase were reduced to H$_2$ by Ve acting as electron donors on Mo$_x$C microparticles acting as redox electrocatalysts. This work shows that, as long as the TTF$^\bullet\bullet$/TTF assembles have an appropriate substrate with a small grain size to grow on, they do not need any metal or organometallic catalyst to perform the photosensitized WOR efficiently. In fact, the TTF/TTF$^\bullet\bullet$@GF junction showed 2 times the efficiency and 5 times the stability of a TTF/TTF$^\bullet\bullet$@Pt analogue. During 16 h of recycling, no decomposition of TTF was observed and H$_2$ was consistently produced, achieving a yield of H$_2$ production of 1.1 $\times$ 10$^{-5}$% in relation to the initial concentration of TTF$^\bullet\bullet$. Constant gas production, constant current, and constant increase of the charge were recorded during 100 h of visible-light-induced WOR and in situ constant anode potential electrolysis of TTF, and no signs of decomposition were observed, indicating that the recycling could be continued further. In conclusion, we have demonstrated a sustainable photosensitized WOR system operating in an emulsified system, where a TTF/TTF$^\bullet\bullet$@GF junction performs as a photosensitizer/molecular-catalyst/redox-shuttle triad with the total absence of metallic or organometallic catalysts and sacrificial electron acceptors. This work opens new perspectives for sustainable metal-atom-free visible-light-driven water splitting in biphasic systems at neutral pH.

## ASSOCIATED CONTENT

### Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacsau.1c00408.

Methods, mechanism of separation of the products of the visible-light-driven WOR by TTF$^\bullet\bullet$, details on the role of vanadium in the proton reduction reaction, an analysis of the headspace of the two compartments of the H-cell depicted in Scheme 2 after 15 min of experiment, a 100 h study of the stability of the TTF/TTF$^\bullet\bullet$@GF junction upon continuous visible-light WOR and in situ constant anode potential electrolysis of TTF, and the study of the stability of TTF by UV-vis absorption and mass spectrometry (PDF).

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