Short Communication

Hydrogen from wet air and sunlight in a tandem photoelectrochemical cell

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Abstract

A solid-state photoelectrochemical (SSPEC) cell is an attractive approach for solar water splitting, especially when it comes to monolithic device design. In a SSPEC cell the electrodes distance is minimized, while the use of polymer-based membranes alleviates the need for liquid electrolytes, and at the same time they can separate the anode from the cathode. In this work, we have made and tested, firstly, a SSPEC cell with a Pt/C electrocatalyst as the cathode electrode, under purely gaseous conditions. The anode was supplied with air of 80% relative humidity (RH) and the cathode with argon. Secondly, we replaced the Pt/C cathode with a photocathode consisting of 2D photocatalytic g-C3N4, which was placed in tandem with the photoanode (tandem-SSPEC). The tandem configuration showed a three-fold enhancement in the obtained photovoltage and a steady-state photocurrent density. The mechanism of operation is discussed in view of recent advances in surface proton conduction in absorbed water layers. The presented SSPEC cell is based on earth-abundant materials and provides a way towards systems of artificial photosynthesis, especially for areas where water sources are scarce and electrical grid infrastructure is limited or nonexistent. The only requirements to make hydrogen are humidity and sunlight.

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Introduction

A fundamental societal concern is the impact of climate change on water quality and quantity [1]. It is also expected that the duration, magnitude and frequency of droughts on a worldwide basis will increase due to climate changes [2]. Water electrolysis is already an important and promising method for hydrogen generation and in the near future, clean water supplies will become a vital concern for this technology. Devices that can directly convert the solar energy to fuels, such as hydrogen, are highly desirable because the solar energy is absorbed and stored in the form of fuels in a single step [3–5]. Such “artificial photosynthesis” (AP) devices hold a great
promise by mimicking nature in order to perform environmentally sustainable reactions, such as splitting water into O2 and H2, and CO2 fixation to sugars [6,7]. The device design is also very important and has an impact on the overall efficiency and cost [8]. Monolithic device designs, where the photoabsorbers are integrated together with the electrocatalysts, have certain advantages such as reduced area requirements, less frames and connections, as well as reduced footprint [9,10].

In our previous work [11], we demonstrated a monolithic solid-state photoelectrochemical (SSPEC) water electrolysis cell, in which the photoanode and cathode electrodes were attached to opposite sides of a Nafion-based polymer electrolyte membrane. The role of the membrane is to separate the evolved gaseous products, support and minimize the distance between the electrodes (down to a few μm) and provide protonic conductivity. The monolithic SSPEC cell was operated under asymmetric conditions i.e. 0.1 M Na2SO4 solution at the anode and gaseous Ar or air in the cathode compartment. There are a few more other works in the literature inspired by a polymer exchange membrane, PEM-like approach, where the photoanode is fed with a water stream [12,13]. Iwu et al. were the first to photoelectrolyze water vapor in an all-solid-state photoelectrochemical (PEC) cell [14], while Georgieva et al. have employed an all-solid-state PEC cell for the decomposition of organic vapors under UV and visible illumination [15]. Recently, the possibility of steam electrolysis in a high temperature PEC cell has been demonstrated by Brunauer and co-workers [16].

Inspired by recent advances in understanding of surface proton conduction in the adsorbed water layers in porous ceramic electrolytes [17,18], we further developed our SSPEC cell, which operates now under ambient wet air conditions. The main advantage of this resource-efficient system is that the water source, which is the main reactant, is coming from the abundant and not exploited humidity in air or steam outputs from industry. This SSPEC device is activated by sunlight and catalyzes the water splitting reaction in the gas phase, alleviating the need for clean water resources. Moreover, the noble-metal cathode, which is usually employed in PEC systems, is replaced by an earth-abundant photocathode. We see that such a monolithic system is important for solar fuels generation in rural areas and in areas where grid infrastructure and water supply are limited or non-existent. The milder operating conditions (not requiring strong basic or acidic aqueous solutions) would decrease the degradation rate of the device components (electrodes, connections, gaskets etc.), a fact of great importance for a long-term performance [19].

In this work, we compare the PEC performance of two SSPEC configurations operated under gaseous conditions. The first one utilizes a Pt-based cathode (n-type|Nafion|C-Pt - SSPEC), as traditionally used in the vast majority of the PEC water splitting cells, while in the second, we replace the Pt cathode with a 2D photocathode g-C3N4 to make an n-type|Nafion|p-type tandem configuration (tandem SSPEC) and compare the two configurations. Finally, the mechanism for enabling water splitting is discussed in light of protonic surface diffusion on the photoanode, enabling larger electrochemically active area. It is worth mentioning that in both configurations the anode was supplied with wet air, while the cathode was supplied with dry Ar.

**Experimental**

Experimental procedures and characterization methods are given in the supporting information (SI). Here, we describe the preparation of the SSPEC membranes and mounting cell, as well as the operating conditions.

As shown in Fig. 1 the solid polymer electrolyte was prepared by drop-casting 200 μL Nafion5 solution onto Cyclpr02 (Cyclopore track etched membrane of 0.2 μm pores size, see SI for more information) and left to dry overnight. The Cyclpr5 (Cyclopore track etched membrane of 5 μm pores size) was used in the tandem-SSPEC configuration as this membrane

![Fig. 1 — Schematic representation of the construction of the button-like SSPEC membrane, as well as its mounting on the custom-made SSPEC device. The SSPEC membrane is sandwiched between two silicone O-rings, which have appropriate copper wiring for connection to the potentiostat. The SSPEC device is of continuous flow type.](image-url)
retained approx. 76% of its transparency after treatment with the Nafion5 (see Fig. S1), while the Cyclpr02 became white. In both cases, the pores of the membranes were fully covered by the Nafion® solution (see Fig. S2). The anode was adhered to the membrane by 20 μL Nafion20 solution as the binding agent, while the cathode was adhered to the opposite side by applying 10 μL Nafion5 solution, and having the Pt-C or g-C3N4 side facing towards the membrane. Then, approx. 13 kPa pressure was applied to the stack overnight in order to adhere the three components together and get a button-like PEC membrane. One may note that the Nafion® is at the same time the proton conducting polymer membrane and the binding agent for the fastening of the anode and cathode electrodes. At this point, we did not construct the g-C3N4/Nafion/C-Pt configuration, which could assess the performance of the photocathode against the C-Pt as the anode electrode. For such a configuration, we need a better stabilization method for the photocathode and this will be addressed in our future works.

The nominal geometric area in the SSPEC configuration is considered as the nominal area of the TiO2 nanotubes (TNTs) mesh electrode, which was 1 cm², while in the tandem-SSPEC configuration it is considered the area of the button cell, approx. 2 cm², as in this case both electrodes contribute to the

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**Fig. 2** — Open circuit voltage (OCV) of the two SSPEC configurations under dark (2000 s) and light conditions (1000 s) (a). Linear sweep voltammograms of the SSPEC (b) and tandem-SSPEC (c) cell configurations at a scan rate of 10 mV s⁻¹ under chopped and continuous 1 Sun illumination. The fast response during the chopped light conditions is highlighted in the tandem-SSPEC cell by the much shorter light-ON, light-OFF steps. Cyclic voltammograms under 1 Sun illumination at a scan rate of 50 mV s⁻¹ (d). Chronoamperometry at 3 different applied voltages. Light on period, 180 s and light off, 20 s (e).

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observed photocurrent. The anode compartment was supplied with air of 80% relative humidity (RH), while the cathode was supplied with dry Ar. The flow rates of both gases were set to 4.5 sccm and the system was otherwise at ambient temperature (25 °C) and pressure (1 atm). Gas leakage and crossover were not determined, but assumed insignificant due to relatively small chemical gradients and absence of total pressure differences, and that the pores of the membrane were blocked by the Nafion® (see Fig. 1 and Fig. S2).

**Results**

Scanning electron microscope (SEM) and X-ray diffraction (XRD) data are provided in Figs. S3–S6. Herein, we focus on the photoelectrochemical characterization of the SSPEC cells, as well as the concept of the surface proton transport, which we suggest as the possible mechanism of the observed photoelectrochemical behavior.

The open circuit voltage (OCV) response under dark and light conditions of the two SSPEC configurations is given in Fig. 2a. It can be seen that the tandem-SSPEC cell produced an OCV of approx. 3 times higher than that produced by the SSPEC cell. This difference is attributed to the combined series photovoltages of the n- and p-semiconductors in the tandem-SSPEC cell, which increase the internal voltage of the device by a factor of three. The potential difference under light in both SSPEC cells is approx. 0.6 V (Fig. 2a), which implies that the Fermi level (EF) in g-C₃N₄ reaches the EF of Pt. This suggests that proton reduction by g-C₃N₄ is thermodynamically favorable [20,21]. In addition, the potential differences in dark and light conditions for the SSPEC configuration are quite consistent, as they depend only on the time constants of charge separation and recombination of one photoelectrode, i.e. TiO₂. In the tandem-SSPEC cell, it can be seen that these values differ between the first and the second light-ON, light-OFF cycle, but less in the third cycle. This behavior may be attributed to the combination of two photocatalysts, which have different time constants, as well as different photoelectrode microstructure and morphology.

The better PEC performance of the tandem-SSPEC cell is evident in the current-voltage curves in Fig. 2b, c, and d where the photocurrent density of the tandem-SSPEC cell exceeds the simple SSPEC. It should be noted that the observed photocurrent in the tandem configuration is normalized by a factor of 2 while in the SSPEC configuration by a factor of 1 (see experimental). It should be noted that the dark scan and the dark from the chopped illumination curve do not match very well. A possible explanation might be that the scan rate used in the LSV experiments, was fast and the steady-state conditions were not reached. This may also explain why the current densities in the potentiostatic experiments of Fig. 2e do not agree with the LSV values. A combination of appropriate scan rate and light-ON, light-OFF steps should be considered for such systems, where the water reactant is not in such an abundance as in a liquid PEC cell. Nevertheless, the tandem cell displays an improved steady-state photocurrent density, as depicted in the chronocamperometric curve (Fig. 2e). It is also interesting to observe that the steady state photocurrent density in the case of the SSPEC cell reaches zero after a few minutes of operation at high applied potentials. Conversely, the tandem cell, displays a significant photocurrent density after 50 min of operation, with short light-OFF periods, which highlight the photocurrent generation. In general, we also observe that the photocurrent density is decreasing, but part of it is retrieved when light is turned on, after a short light-OFF period. This may be related to the heating of the surface of the photoanode, so that part of the physisorbed water layers is removed. The physisorbed water layers are compensated during the short light-OFF period. This hypothesis will be tested when the light-Off periods are increased and/or by thermostating the cell at higher temperatures, so that heating from the light is excluded. More details are given in the discussion section.

The superiority of the tandem configuration over the single SSPEC one is also seen by the electrochemical impedance (EIS) measurements of Fig. 3. The Nyquist plot is recorded at 0.5 V in a two-electrode configuration under 1 Sun illumination. Three depressed semicircles are observed in the tandem SSPEC cell, which are related to the charge transfer resistance over the two photoelectrodes, as well as a mass transfer resistance associated with chemical diffusion of reactants and products [17]. The charge transfer resistance of the SSPEC is clearly higher than the tandem-cell, and is attributed to the extra photovoltage provided by the photocathode, giving a higher electrochemical activity than the Pt cathode under illumination. In the dark though, a high to mid frequencies semicircle is seen for the SSPEC cell, but not for the tandem-SSPEC cell (see Fig. S7a). The charge transfer semicircle in the tandem-SSPEC is most probably much higher in the dark and merged with the capacitive behavior in the low frequency domain, indicating a higher resistance for the photocathode in the dark, as compared to C-Pt. The equivalent circuits used to model the raw EIS data are given in Fig. S7b, but details and the physical interpretation are beyond the scope of this communication. Deconvolution of the EIS spectra showed the
same high frequency resistance (approx. 850 Ω cm² in both dark and light conditions), which is related to the proton conductivity of the polymer electrolyte (2.4 10⁻² mS cm⁻¹) at room temperature and 80% RH. The overall resistance was found to be 27 kΩ cm² and 412 kΩ cm² for the tandem-SSPEC and SSPEC cell, respectively. This further supports the larger photocurrent density of the tandem cell.

Discussion

In this work we have demonstrated the generation of photovoltage and photocurrent of a solid-state PEC cell operating with humidified air as the reactant stream. Moreover, the steady state photocurrent density is significantly enhanced by replacing the Pt cathode with an earth-abundant photocathode, g-C₃N₄ (Fig. 2d). The addition of the photocathode – due to its proper alignment with the photoanode (see Fig. S8) – adds to the generated photovoltage due to the series connection and enhances the proton reduction reaction at the cathode. In principle, the tandem-SSPEC resembles a z-scheme photocatalytic system [20], with the advantage that the oxidation and reduction products can be separated.

These SSPEC cells operate with humid gas as the supply of water, and thus resemble that of proton ceramic electrolyzers operated at intermediate to high temperatures. Contrary to traditional PEC cells, the charge transfer reaction (water oxidation) in such systems occur directly at the triple-phase-boundary (TPB) – the interface between electrolyte, electrode and gas, as the protons formed from the water oxidation reaction can directly “enter” the polymer electrolyte [22,23]. The mass transfer reactions – adsorption, dissociation and diffusion – can occur along the entire electrode surface but is then limited by the chemical diffusion to the TPB for the charge transfer reaction to proceed. Thus, at low temperatures where the diffusivity is rather low, the total reaction is commonly limited to a small area close to the TPBs. For our current cell design, this would severely limit the

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The porosity of the cathode electrode, as well as the Ar flow combined with the electrons and hydrogen gas is produced. In the phase, the electrolyte and the cathode and the protons are prevalent in our photoanodes operated at 80% RH, aiding the chemical diffusion of adsorbed species and reactants along the electrode surface and expanding the electrochemically active area beyond that of the triple-phase-boundaries (see Fig. 4).

At the cathode, another TPB is formed between the gas phase, the electrolyte and the cathode and the protons are combined with the electrons and hydrogen gas is produced. The porosity of the cathode electrode, as well as the Ar flow ensures the H₂ gas evolution and removal from the surface of the catalyst (Fig. 4). On the other hand, the dry conditions in the cathode may have an impact on the water content close to the cathode side, resulting in the low proton conductivity observed in the present case (2.4 $10^{-2}$ mS cm$^{-1}$) and limited proton diffusivity along the surface. Adding humid Ar atmosphere could be a possibility, or a composite membrane, which stays hydrated at low RH [29,30]. It should be mentioned that H₂ was not detected due to the low current which stays hydrated at low RH [29,30]. It should be acknowledged.

Thus, it is reasonable to assume surface proton conduction is absent, and sources of clean water are scarce. What is needed is only sun and humid air.

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Conclusions

In this work we introduced the concept of water vapor splitting for hydrogen production in a solid-state PEC cell, after recent developments in the surface proton conduction in porous ceramic-based electrolytes. The traditionally Pt-based cathode is replaced by an earth-abundant photocathode, g-C₃N₄, which increased the intrinsic photo-induced electrical field by a factor of three. Moreover, we replaced the liquid electrolytes by a polymer proton conducting membrane based of Nafion®. We envision that such a monolithic solid-state PEC cell can play an important role for clean energy production in rural or other areas where grid infrastructure is limited or

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References


