Novel integrated strategies toward efficient and stable unassisted photoelectrochemical water splitting

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Abstract
© 2020 Unassisted photoelectrochemical (PEC) water splitting for hydrogen evolution has been regarded as a sustainable route for the harvest and utilization of solar energy. To achieve such unassisted PEC water splitting, two novel integrated strategies have been developed: to design tandem structures of photoanodes/photocathodes and to construct hybrid devices of solar/PEC cells. Some key unsolved problems, however, still limit the further development of high-performance unassisted PEC water splitting, such as low efficiency, poor stability, and high cost. Herein, we present a brief summary of the latest development in this area and propose perspectives for further enhancing this state-of-the-art solar-to-hydrogen conversion technology, including all-metal oxide photoelectrodes, nanoarray design, surface modification, device coupling, monolithic configuration, and multi-integration.

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\textbf{Abstract}

Unassisted photoelectrochemical (PEC) water splitting for hydrogen evolution has been regarded as a sustainable route for the harvest and utilization of solar energy. To achieve such unassisted PEC water splitting, two novel integrated strategies have been developed: to design tandem structures of photoanodes/photocathodes and to construct hybrid devices of solar/PEC cells. Some key unsolved problems, however, still limit the further development of high-performance unassisted PEC water splitting, such as low efficiency, poor stability, and high cost. Herein, we present a brief summary of the latest development in this area and propose perspectives for further enhancing this state-of-the-art solar-to-hydrogen conversion technology, including all-metal oxide photoelectrodes, nanoarray design, surface modification, device coupling, monolithic configuration, and multi-integration.

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1. Introduction

Since the first industrial revolution in the mid-eighteenth century, traditional fossil fuels (e.g., coal, oil, and natural gas) have greatly promoted the development of the societies’ productivity. With the rapid industrialization and urbanization in recent decades, nevertheless, fossil fuels have been excessively exploited and consumed. It inevitably
leads to two major crises, including the energy shortage and environmental deterioration [1–3]. This is because fossil fuels have a limited reserve on the earth and will soon be exhausted according to their current annual consumption rates. Besides, the combustion of fossil fuels also releases a huge amount of toxic/green-house gases and harmful particle matters, resulting in severe environmental issues, such as global warming, acid rain, ozone depletion, smog, and so on [4,5].

In order to tackle these problems, a sustainable strategy is to capture and utilize renewable and clean energy sources (i.e., new energy sources), including solar energy [6–8], wind energy [9], nuclear energy [10], geothermal energy [11], wave/tidal energy [12,13], biomass energy [14], natural gas hydrates [15], etc., to replace the aforementioned fossil fuels. Among the various types of new energy sources, solar energy has been regarded as a most promising alternative of the carbon-based fossil fuels because of its unique merits, summarized as following [16–18]:

i) Abundance. The energy carried by sunlight reaching the earth's surface is $5.4 \times 10^{17}$ kWh per year, which is about 5000 times of the global annual energy consumption. That is to say, even if 0.02% of solar energy is effectively used, human energy demand can be satisfied.

ii) Sustainability. Based on the rate of hydrogen fusion in the Sun, its hydrogen reserves can be consumed continuously for more than $6.0 \times 10^{10}$ years, meaning that solar energy is an inexhaustible and renewable energy resource for us.

iii) Clean energy. Unlike traditional fossil fuels, solar energy virtually does not produce any pollution. Thus, solar energy can be considered as green energy, which is very beneficial for protecting the ecological environment.

iv) Convenience. In theory, solar energy can be harvested and further utilized wherever the sunlight can shine, which is especially facile for the remote areas. Besides, miniaturized and flexible solar cells also provide a great possibility for the development of self-powered portable devices.

Although solar energy exhibits the above-mentioned advantages, its inherent shortcomings, including intermittency, instability, and low power density, still impede the large-scale commercial application [19,20]. To achieve the efficient utilization of solar energy, the conversion of solar energy into hydrogen fuel (i.e., hydrogen gas) has become a potential solution in this field [21]. Specifically, as an ideal carrier for solar energy, hydrogen fuel shows a serial of excellent properties, such as carbon-free and non-emission features, high energy density (~140 MJ kg$^{-1}$), storability (e.g., hydrogen storage alloys and liquid hydrogen), and mature technology for electricity generation (e.g., hydrogen fuel cells and hydrogen gas steam boilers) [22–26]. Consequently, the solar-to-hydrogen (STH) conversion can not only overcome the drawbacks of solar energy but also provide a feasible way for the large-scale and long-term development of solar energy.

As one typical artificial photosynthesis process, photocatalytic water splitting has been considered as a highly promising strategy for the STH conversion, in terms of its relatively high energy conversion efficiency, environmentally-friendly process, and abundant hydrogen resource [27–33]. A typical PEC water splitting process can be achieved in a PEC cell, commonly with two solar-driven half-reactions: hydrogen evolution reaction (HER) and oxygen evolution reaction (OER), which are carried out at surfaces of (photo)anodes and (photo)cathodes, respectively (Eqs. (1) and (2)).

\[
\text{HER at (photo)cathode : } 2H^+ + 2e^- \rightarrow H_2, E_{\text{red}}^0 = 0 \text{ V (1)}
\]

\[
\text{OER at (photo)anode : } 2H_2O + 4h^+ \rightarrow 4H^+ + O_2, E_{\text{ox}}^0 = 1.23 \text{ V (2)}
\]

where, \(E_{\text{red}}^0\) and \(E_{\text{ox}}^0\) are H$_2$ evolution potential (0 V vs. normal hydrogen electrode (NHE) at pH = 0) and O$_2$ evolution potential (1.23 V vs. NHE at pH = 0), respectively.

Theoretically, as long as the bandgap of photoelectrode is equal to or larger than the minimum energy for decomposing water molecules (1.23 eV), photogenerated electrons and holes produced on photoelectrodes under illumination can drive the HER and OER, respectively. Unfortunately, undesirable energy loss often exists during the PEC water splitting process, such as over potentials and contact resistances. As a result, an external bias is commonly necessary to realize the PEC water splitting. Obviously, this external bias produced by a power supply will lead to unnecessary consumption of electricity.

Recently, the unassisted PEC water splitting has become a research hotspot due to its more energy-saving and efficient characteristics for the STH conversion [34,35]. The namely “unassisted PEC water splitting” refers to the PEC process free from the external bias, in which all the energy used to split water molecules into hydrogen gas and oxygen gas is only provided by the incident sunlight.

Different from the conventional PEC water splitting with an external bias, the unassisted PEC water splitting is usually based on two novel strategies: to design tandem structures of photoanodes/photocathodes, and to construct hybrid devices of solar/PEC cells [36]. Specifically, for the former, a photoanode is coupled with a photocathode in a PEC cell to form a tandem structure (Fig. 1a). By this strategy, the photogenerated electrodes and holes can be produced simultaneously in both the photoanode and the photocathode under the incident light irradiation. Consequently, the photovoltages generated by the photoanode and the photocathode will be combined to provide sufficient voltage for water splitting. Therefore, this type of unassisted strategy can be described as the non-biased PEC water splitting. As for the latter strategy, a solar cell module is integrated with a PEC cell to construct a hybrid device. When this hybrid device is illuminated by solar light, the solar cell can provide the required bias for PEC cells to carry out the water-splitting reaction (Fig. 1b). On this basis, this second strategy can be regarded as the self-biased PEC water splitting.

Briefly, these two strategies can open up new avenues for the development of high-performance unassisted PEC water splitting technologies. Up to now, a serial of research has been carried out to develop these two types of technologies, including development on novel photoelectrode materials and design on unique device configurations [37–40]. However, there are still several critical aspects remaining unsolved, such as low STH efficiency of <10% (i.e., the standard requirement for practical applications), high cost, and poor stability. Hence, more efforts should be made to further improve the performance of these unassisted PEC systems.

Based on these considerations, we believe that it is necessary to discuss in detail the future development of these two state-of-the-art technologies for PEC hydrogen production, and further propose feasible solutions to achieve the efficient unassisted PEC water splitting. Herein, we focus on the technological trends of the tandem structures and the hybrid devices, respectively.

For the former, three methods could be particularly feasible, including all-metal oxide photoelectrodes, nanoarray design, and surface modification. For the latter, device coupling, monolithic configuration, and multi-integration should be paid more attention to, in order to enhance the STH efficiency and stability of the hybrid systems (Fig. 2). In brief, all aspects mentioned above may attract the focus of research and become potential hot spots. At last, we hope that this perspective is valuable to inspire rational designs and developments for efficient and stable unassisted PEC water splitting.

2. Tandem structure design of photoanodes and photocathodes

The pioneering study of the tandem structure for unassisted PEC water splitting can be tracked back to 1977 [41]. Although it has developed for more than 40 years, the tandem structures are far from satisfying the commercial application, due to its low STH efficiency, poor stability, and high cost. To solve these issues, the following three effective strategies are important, as illustrated in Fig. 2:
2.1. All-metal oxide photoelectrodes

Although some high efficiencies that are close to or even higher than 10% have been obtained in the laboratory, the photoelectrode materials utilized in these unassisted PEC tandem cells commonly involve GaN, InN, InP, or Si-based semiconductors [42,43]. Obviously, some fatal shortcomings of these semiconductors, such as poor stability, high cost, and serious toxicity, make them unsuitable for large-scale applications. Compared with the aforementioned materials, metal oxides are more promising for photoelectrode materials of tandem PEC cells in terms of good stability, low cost, non-toxicity, and facile fabrication.

Moreover, some other excellent features, such as n/p-type characteristics, wide distribution of the bandgap, suitable band edge positions for water electrolysis, and easily tunable electronic properties, show also better potential as photoelectrode materials. Firstly, metal oxides exhibit either intrinsic n-type (e.g., TiO₂, ZnO, and WO₃) or p-type (e.g., Cu₂O and NiO) characteristics. Generally, the former can be used as the photoanodes, while the latter can be used as photocathodes. Thus, this provides a great possibility for simultaneously using both the n-type metal oxide as photoanode and the p-type metal oxide as a photocathode in a tandem PEC cell for unassisted solar-driven water splitting.

Secondly, metal oxides also own a wide-range distribution of bandgaps (~3.4–2.0 eV), which can provide the appropriate band edge positions for water electrolysis and sufficient solar energy absorption. For instance, as a typical wide bandgap semiconductor, TiO₂ has a bandgap of ~3.2 eV, and its edges of the conduction band and valence band locate on the potentials of −0.1 and +3.1 eV vs. NHE, respectively. That is to say, TiO₂ can provide photogenerated electrons and holes with sufficient energy to drive a water-splitting reaction under solar irradiation. On the other hand, as a narrow bandgap metal oxide, α-Fe₂O₃ has a relatively small bandgap of ~2.2 eV, which enables a high visible light absorption. Further, if a wide bandgap metal oxide (i.e., TiO₂) is sensitized by a narrow metal oxide (i.e., α-Fe₂O₃), the all-metal oxide photoelectrode material (i.e., α-Fe₂O₃/TiO₂) can not only provide the appropriate band edge positions for PEC water-splitting reaction but also exhibit the satisfactory visible light response.

Lastly, the electronic properties of metal oxides can be easily adjusted as well, such as by elemental doping or defect engineering, providing sufficient room for further enhancing the photoelectrodes. Consequently, these versatile advantages of metal oxides offer a large number of opportunities to construct the tandem PEC cells based on all-metal oxide photoelectrodes for unassisted PEC water splitting with high practical value.
2. Nanoarray design

For efficient unassisted PEC water splitting, the successful suppression of the recombination of the photogenerated electrons and holes is one of the most critical factors. However, it is very challenging for the conventional mesoporous films, which are typically composed of nanoparticles (e.g., TiO₂ P25 powders), to meet this requirement. This is because the large number of grain boundaries in such materials would act as the charge-trapping sites, leading to a high recombination probability between photogenerated electrons and holes [44]. For this issue, using the highly ordered nanoarrays as photoelectrode materials can be a feasible solution to obtain the low exciton recombination, including nanotube arrays, nanorod arrays, nanowire arrays, and nano-sheet arrays. In these nanostructured photoelectrodes, the highly ordered configurations can significantly shorten the diffusion distance of the photogenerated charge carriers to the surface of the material, resulting in the much-reduced recombination of photogenerated electrons/holes.

Except for the above-mentioned effect, the nanoarray designs also exhibit other unique merits. On the one hand, the highly anisotropic arrays can greatly increase the scattering of incident light in the photoelectrodes, thus enhancing the solar light harvesting. On the other hand, the nanoarrays can provide a large specific surface area, giving them more catalytically active sites and better contact with the electrolyte. Besides, the nanoarrays can also accelerate the release of gas bubbles generated at the surface of photoelectrodes, which is helpful to boost the dynamic process of PEC reactions. As a result, the nanoarray photoelectrodes are very promising potential candidates for high-performance unassisted PEC water splitting.

2.3. Surface modification

Apart from the above two points, the surface modification of photoelectrodes is also essential for achieving a more efficient and stable unassisted PEC water splitting based on the tandem structures. For instance, the OER commonly exhibits sluggish kinetics and a large overpotential, thus significantly limiting the overall STH efficiency of the whole unassisted system [45]. To overcome this problem, depositing cocatalysts onto the photoanode surfaces has been regarded as an effective method [36]. Especially, two emerging types of cocatalysts could exhibit the super-high catalytic activity, including metal oxyhydroxide (e.g., NiOOH, CoOOH, and FeOOH), and single-atom catalysts (SACs) [23,46].

For those photoelectrode materials that are easily photo-corroded, such as Si and CdS, the deposition of a highly stable protective layer can further enhance the durability of the tandem PEC systems. For example, graphitic carbon nitride (g-CN) owns excellent physicochemical performance and can act as such protective layers to enhance the stability of Si or CdS photoelectrodes when it is coated onto them [47,48]. Furthermore, the in-situ deposition of g-CN can also keep the original micro-morphologies of photoelectrodes, which is very beneficial to enhance the PEC performances as well [49].

3. Hybrid devices of PEC/solar cells

Except for the tandem structure design, integrating PEC cells with solar cells to form hybrid devices is another feasible way to achieve unassisted solar-driven water splitting. In such devices, the solar cell provides the bias for the PEC cells to carry out the water photolysis for hydrogen production under sunlight illumination.

Despite of the novel concept of hybrid devices, there are still some stumbling blocks on its road toward the commercialization, such as low STH efficiency, poor stability, and mismatching between PEC cells and solar cells, etc. In order to obtain the practical hybrid devices of PEC/solar cells, we believe that the following three constructive opinions should be paid full attention to (Fig. 2):

3.1. Device coupling of PEC/solar cells

In the early research stage, dye-sensitized solar cells (DSSCs) are frequently integrated with PEC cells for unassisted PEC water splitting due to their similar electrochemical behaviors [50,51]. However, the low open-circuit voltage ($V_{OC} < 0.5 V$) produced by DSSCs is not sufficient for PEC cells, leading to a potential mismatching between solar cells and PEC cells, which greatly influences the STH efficiency of the hybrid devices. The novel perovskite solar cells (PSCs) not only generate a relatively high $V_{OC}$ ($0.9–1.1 V$), but also exhibit an excellent photoelectric conversion efficiency (>20%) [52–54]. Consequently, introducing PSCs into the PEC/solar cell hybrid devices is a promising approach for achieving high-performance unassisted PEC water splitting [34].

Apart from the voltage coupling, the light absorption balance between PEC cells and solar cells should also be carefully considered to more sufficiently harvest solar energy. Specifically, if a PEC cell is placed in front of a solar cell, the photoelectrode will absorb the incident light with a wavelength smaller than its bandgap. Thus, the long-wavelength part of the incident light would transmit through the photoelectrode and be absorbed by the solar cell. In this case, the PEC cell and the solar cell should be designed to be capable of absorbing different portions of the visible light to obtain a broad solar spectrum response. To realize this, two possible types of PEC photoelectrodes could be adopted, either on transparent conducting substrates or on mesh-shaped substrates.

3.2. Monolithic configuration

Two main strategies can be used to connect the PEC cell and the solar cell. One is the wire connection (i.e., external connection), and the other is the monolithic configuration (i.e., inner connection) [35]. In the first mode, the PEC cell and the solar cell are simply connected via external wires. Unfortunately, this technology leads to not only the loss of electric energy caused by the extra resistance but also the difficulty in light path design.

To obtain an efficient hybrid device, the monolithic configuration could be an ultimate solution. In this configuration, the PEC cell and the solar cell are integrated into a stand-alone device for unassisted PEC water splitting. This mode can decrease the device resistance, thus resulting in a high photocurrent and STH efficiency. But it needs to be considered that the solar cell in the monolithic hybrid device has to be immersed in and contact with the electrolyte, thus leading to a possible loss of stability for the whole system. Therefore, a reliable packaging technology, which can isolate the solar cell from the electrolyte and simultaneously provide the full contact between the PEC photoelectrode with the electrolyte, is essential to long-term stable work. This is particularly important for realizing stable unassisted PEC seawater splitting due to the serious corrosion capability of natural seawater.

3.3. Multi-integration

As mentioned, solar energy is greatly affected by day and night alternation, and change of weather and season, thus leading to intermittence and instability [19]. These disadvantages significantly limit the stable and continuous hydrogen production from unassisted PEC water splitting. Recently, a rising technology of converting and storing solar energy in the form of electrochemical energy could provide a solution to the aforesaid issue. Specifically, such devices, which are commonly composed of solar cells and electrochemical energy storage (EES) devices (e.g., lithium-ion batteries, sodium-ion batteries, lithium-sulfur batteries, supercapacitors, etc.), can convert solar energy into electricity, and subsequently store the electricity in EES units [55,56].
For the multi-integration of PEC cells/solar cells/EES units, the solar cells can simultaneously provide the bias for PEC cells and charge for the EES units when the whole system is irradiated by sunlight. On the other hand, the EES units can discharge to drive the water-splitting reaction in PEC cells under the dark condition. Consequently, the multi-integration strategy can achieve an all-weather and 24-h water splitting for hydrogen generation, which is necessary for the large-scale industrial production. Besides, the multi-type energy harvesting, such as introducing nanogenerators into hybrid devices of PEC/solar cells, is also a feasible route to develop advanced unassisted PEC water splitting.

4. Conclusion and outlook

As a sustainable energy strategy, solar-driven water splitting for hydrogen evolution can simultaneously address the energy crisis and the environmental pollution [57–61]. Especially, unassisted PEC water splitting exhibits the greater development potential in comparison with other STH technologies. Specifically, unassisted PEC water splitting can be divided into two categories: the tandem structure of photoanode and photocathode and the hybrid device of solar/PEC cell. Although many efforts have been made, the unsolved problems still limit further development toward a more efficient, stable, and low-cost unassisted PEC water splitting. As for the multi-integration of PEC cells/solar cells/EES units, the solar cells can simultaneously provide the bias for PEC cells and charge for the EES units when the whole system is irradiated by sunlight. On the other hand, the EES units can discharge to drive the water-splitting reaction in PEC cells under the dark condition. Consequently, the multi-integration strategy can achieve an all-weather and 24-h water splitting for hydrogen generation, which is necessary for the large-scale industrial production. Besides, the multi-type energy harvesting, such as introducing nanogenerators into hybrid devices of PEC/solar cells, is also a feasible route to develop advanced unassisted PEC water splitting.

Fig. 3. Further developments on the tandem structure and the hybrid device toward more efficient, stable, and low-cost unassisted PEC water splitting.