1. Introduction

Energy, which can exist in a variety of forms, describes the capacity or power of a system to do work. There are many energy resources, including nonrenewable and renewable ones. Except for nuclear, tidal and geothermal energies, all resources of energy ultimately come from the sun. Energy conversion is the process of transforming one form of energy to another, which complies by the laws of thermodynamics. Correspondingly, there exist many different energy conversion devices that can convert energy from different energy sources into useful forms (in particular, electricity and heat) to meet our needs. Among those energy conversion devices are solar photovoltaic (PV), thermoelectric, electrochemical and photoelectrochemical (PEC) cells that have been main research areas particularly in recent years when people realized that renewable energy is the key to creating a clean energy future for the world. In the past, these cells have been developed independently with each other. For example, fuel and solar cells have been traditionally regarded to belong to different disciplines of electrochemistry and semiconductor, respectively, in spite of the facts that both the first demonstration of a fuel cell done by William Grove and the first PV effect observed by Edmund Becquerel are in the same year of 1839, and both of them are electricity-producing devices. In this review article, however, the authors showed that there are close connections between the working principles of solar PV and other energy conversion cells, which may provide more insights in designing better energy conversion devices.
into electrical energy. According to the Carnot’s theorem that is a result of the second law of thermodynamics, the energy conversion efficiency of heat engines is low. An electrochemical cell (fuel cell or battery cell), however, can directly convert the chemical energy of a fuel into electricity. Therefore, electrochemical cells create electricity chemically, rather than by combustion, at a higher efficiency at least in principle, because they are not subjected to thermodynamic laws that limit a conventional power plant. In addition, waste heat from some electrochemical cells can also be harnessed, boosting the system efficiency still further. Alternatively, heat can be produced from solar energy with solar collectors, such as parabolic troughs/dishes and linear Fresnel reflectors for electricity production, that is, concentrating solar power. In addition, heat can be directly converted into electrical energy with thermoelectric cells. Because of the heating (or uneven heating for the case of wind energy, to be more accurate) of solar radiation on the fluid (air or water), air or water flows from a higher energy level to a lower energy level so that wind or hydro energy is obtained that can be harnessed to produce electrical energy with wind or water turbines. With even fewer steps, a solar cell converts solar energy directly into electricity using the PV effect.

Before discussing in detail on each kind of these cells, it may be helpful to get some perspective on them with a simple picture of energy conversion (Figure 2). There are a few key things for a working energy conversion device, explained as follows:

1. Key components: At least two different energy levels with available energy states for mobile carriers and a path loop for carriers to flow through.

2. Carrier pumping: Carriers must be pumped from a lower energy level to a higher energy level with the energy input, which can be done either in situ or ex situ.

3. Carrier separation: Carriers at the higher energy level have to be physically separated from their corresponding empty states at the lower energy level inside the cell to reduce recombination. Otherwise, the energy leakage is high, and significant amount of energy is wasted inside the cell.

4. Carrier transport: The carrier transport must be as smooth as possible with minimum resistance except at the load to achieve high energy conversion efficiency.

5. Energy conversion efficiency (η) = (Energy output) / (energy input). Power output (P) = current × potential = I × V. The energy not converted into work done on the load as energy output is often dissipated as waste heat.

In fact, the above picture can be put in the more basic nonequilibrium theory, which states that under nonequilibrium condition, there is always a current of something trying to reduce a gradient in something else that is the actual cause of the nonequilibrium such that equilibrium can be re-established. As far as the charged
particles such as electrons are concerned, a gradient in the electrical potential causes drift currents and a gradient in a chemical potential (which is a function of concentration and temperature) causes diffusion currents. The total driving force for electrons is then the gradient of the so-called electrochemical potential (the sum of electrical potential and chemical potential).

In the following discussion, the authors focused on solar cells, thermoelectric cells, batteries, fuel cells and PEC cells, because these cells have been the active and main research areas in recent years as aforementioned. It is also important to note that all these cells produce direct current (DC) due to the flow of electrons. So it is important to know the energy levels of electrons in different mediums in these cells.

3. Solar cells
PV is a technique of directly converting solar radiation into electricity using semiconductors that exhibit the PV effect. As solar energy is the most abundant energy resource, it is almost guaranteed to be an indispensable part of the energy system. Compared with other energy conversion paths from solar energy to electricity, PV has a higher overall efficiency. In addition, PV is the only renewable energy source that can be used everywhere. So far, there have been many kinds of solar cells. Traditionally, solar cells have been classified according to the materials used in absorber layers, that is, essentially charge generation. Here, the authors made a slightly different approach, which may provide more insights on the working principles of solar cells. On the basis of junctions applied, these cells can be classified into three major groups, that is, inorganic p-n junction–based solar cells, organic e-donor/e-acceptor junction–based solar cells and hybrid junction–based solar cells (Figure 3). In addition, hybrid junction-based solar cells can be further divided into two subgroups: dye-sensitized solar cells and other hybrids. New concept solar cells, in particular those based on nanostructures and quantum confinement, have been investigated also in recent years.

Figure 4 illustrates the basic working principles of a solar cell under sunlight illumination. There are three key steps for a working solar cell. First of all, there is charge carrier generation (or light pumping): Electrons in the semiconductor (with right band gap $E_g$) absorber are pumped from the valence band (VB) to the conduction band (CB) in the presence of sunlight. Second, there is charge carrier separation: Electrons in the CB and holes in the VB have to move in different directions so that they are physically separated from each other to minimize direct recombination inside the device, which is often realized with selective contacts. Finally, there is charge carrier transport/collection: Carrier transport through the absorber, selective contacts, metal wire and interfaces between them has to be smooth, that is, the resistance should be minimized, which requires that suitable materials be selected with the right energy band alignment between them and less defect states. Overall, electrons flow along the path loop, absorbing solar energy when pumped from the lower level VB to the higher level CB and releasing their energy by doing work on the load, in a similar fashion as the hydrologic cycle in hydro power.

4. Thermoelectric cells
In this section, the discussion is limited to the thermoelectric cells for power generation only; other thermoelectric devices based on different thermoelectric effects will not be considered here. This kind of cell is applied to convert heat energy directly into electrical energy based on the Seebeck effect. A typical thermoelectric device essentially consists of metal–semiconductor junctions as the one shown in Figure 5(a). The device is configured as a power generator, and it creates a voltage when there is a temperature difference $\Delta T$ between the two sides of semiconductors.

![Figure 3. Classification of solar cells based on junctions applied.](image-url)

- **Inorganic semiconductor solar cells:** Si, GaAs, CdTc, CIGS, CZTS...
- **Organic semiconductor solar cells:** P3HT/PCBM
- **Dye sensitized solar cells**
- **Other hybrid solar cells:** inorganic nanostructure + organic matrix...

ClIGS, copper indium gallium (di)selenide; CZTS, copper zinc tin sulfide; P3HT, poly(3-hexylthiophene); PCBM, [6,6]-phenyl C61-butyric acid methylester.
If there exists no temperature gradient, the Fermi energy must be equal everywhere once the materials are brought into contact, and the equilibrium condition is reached. In thermoelectric cells, however, the temperature is not constant throughout the semiconductors. However, the Fermi energy can still be regarded to be constant throughout the material because the effect of the temperature on the Fermi energy are small in particular when the carrier density is high, for example, in heavily doped semiconductors. Therefore, it is reasonable to treat approximately the thermoelectric effect with equilibrium thermodynamics by simply assuming that locally we are not very far from equilibrium and thus can still use band structure models with a constant Fermi energy \( E_F \) (Figure 5(b)). Because of the presence of a thermal gradient across the p- and n-type semiconductors, carriers (free electrons or free holes) have a higher average energy at the hot side than at the cold side. The diffusion of carriers from the hot side to the cold side then occurs because carriers intend to move to a lower energy level so that more free electrons (or holes) are accumulated at the cold side than at the hot side of the n-type (or p-type) semiconductor. However, there exists a backward force that prevents the diffusion – the charged donors/acceptors left behind at the hot side result in an electric field causing the backflow of current that will eventually cause the system to reach steady-state equilibrium under open-circuit condition. The electric field in each semiconductor corresponds to a thermovoltage. So semiconductor legs in a thermoelectric device are connected electrically in series but thermally in parallel. The thermovoltage can be calculated according to Nernst equation, which describes the chemical potential difference \( \Delta U \) between two locations solely due to the activity (concentration related) difference of mobile carriers.

\[
\Delta U = -\frac{kT}{ze} \ln \frac{n_a}{n_c} = -\frac{kT}{ze} \ln \frac{\gamma_c}{\gamma_a}
\]

where \( k \) is the Boltzmann constant, \( T \) the temperature, \( e \) the elementary charge, \( z \) the number of elementary charges per carrier, \( a \) the activity, \( c \) the carrier concentration and \( \gamma \) the activity coefficient. It is obvious then that there exists a driving force that will push electrons to flow from the anode to the cathode once the two electrodes are connected via an external circuit. Unlike the case of solar cells where only the high-frequency part of the radiation is used to pump electrons across the band gap, the low-frequency heat energy is applied in thermoelectric power cells, which makes them complementary to solar cells to increase the overall efficiency of converting solar radiation into useful electricity.

## 5. Electrochemical cells: batteries and fuel cells

Here, the simple Daniel cell has been taken as an example to show the general working principles of batteries from energy (rather than traditional electrochemical) point of view. In the Daniel cell, copper and zinc electrodes are immersed in a solution of copper...
(II) sulfate and zinc sulfate, respectively, and a salt bridge is often used to connect the two half cells to help maintain charge neutrality (Figure 6(a)). Electron-transfer reactions are oxidation and reduction at the anode and cathode, respectively. Taking the standard hydrogen electrode (SHE) as the energy reference point (0 V), the standard electrode reduction potentials can be considered to be −0.76 and 0.34 V, respectively, for the following two reduction reactions:

2. \[ \text{Zn}^{2+} + 2e^- \rightarrow \text{Zn} \]

3. \[ \text{Cu}^{2+} + 2e^- \rightarrow \text{Cu} \]

Therefore, the difference between the two reduction potentials as the standard electromotive force (EMF) for the cell \( \Delta E_{\text{cell}}^0 \) can be considered as follows:

4. \[ \Delta E_{\text{cell}}^0 = -E^0_{\text{cathode}} - E^0_{\text{anode}} = 0.34 - (-0.76) = 1.10 \text{ V} \]

Considering the above situation in energy space, the working principles of the Daniel cell in a straightforward way are shown in Figure 6(b). Compared with Figure 4, the net results are quite similar in both a solar cell and a battery cell, that is, electrons are at a higher energy level at the anode side than their corresponding empty states (i.e., holes) at the cathode side, and they intend to flow to the lower energy level via the external circuit. Of course, there are some differences between a battery cell and a solar cell. For example, there are no chemical reaction and ion transport involved in the latter (except for dye-sensitized solar cells where a redox electrolyte has been applied for charge transport), and a solar cell is essentially a current source while a battery cell is a voltage source as pointed out by Nelson. In addition, the authors considered the standard condition for a working battery cell. The real situation to calculate the EMF is more complicated due to the possible variations in temperature, pressure, chemical activity and so forth. However, the authors think that the complication will not blur the clear connection between the working principles of these two kinds of energy conversion cells.

A fuel cell essentially works in a very similar way as a battery in which both convert chemical energy into electrical energy, produce DC and use electrolytes to conduct ions (Figure 6 and Figure 7). The similarity can be further verified by comparing the chemical reactions in the Daniel cell with those in the five major types of fuel cells shown in Table 1. However, there are differences between them mainly in fuel supply. A battery is a closed system with a fixed amount of chemical energy stored in the reactants while a fuel cell will keep producing electricity as long as fuel is supplied externally, that is, fuel cell reactions do not degrade over time.

6. PEC cells
Hydrogen (H\(_2\)) is a clean noncarbon fuel, and as aforementioned, the sun is the most abundant energy resource. The energy issue would be greatly alleviated if H\(_2\) could be produced by directly splitting water using solar energy with affordable cells. Water splitting to produce H\(_2\) can be traditionally realized with electrolysis, where the decomposition of water (H\(_2\)O) into oxygen (O\(_2\)) and hydrogen (H\(_2\)) occur when DC is connected to the two inert electrodes immersed in the water (Figure 8(a)). Half reactions at the anode and cathode, respectively, are as follows:

5. \[ \text{H}_2\text{O} \rightarrow \frac{1}{2} \text{O}_2 + 2\text{H}^+ + 2e^- \]
The corresponding standard reduction potentials with respect to the SHE are 1.23 and 0.00 V, respectively. Therefore, the standard EMF for the total reaction (2H₂O → O₂ + 2H₂) to proceed can be calculated to be −1.23 V in a similar way as in Equation (4). The negative value means that the reaction is not thermodynamically spontaneous, and the driving force must be provided externally.

The above redox reactions, from semiconductor point of view, have been rearranged as follows:

6. \[ 2H^- + 2e^- \rightarrow H_2 \]

7. \[ \frac{1}{2}H_2O + h^+ \rightarrow \frac{1}{4}O_2 + H^+ \]

8. \[ H^+ + e^- \rightarrow \frac{1}{2}H_2 \]
Electron–hole pairs are necessary for the redox reactions to proceed. It is well known that electron–hole pairs can be generated in semiconductors illuminated with suitable light as seen in solar cells. It is natural then to think of producing hydrogen from water splitting with direct solar energy by combining suitable semiconductors with redox reactions.

Figure 8(b) shows an n-type semiconductor-based PEC cell (other configurations such as those based on p-type semiconductor or both are also possible).26–28 The Fermi levels of the n-type semiconductor and the counter electrode are aligned. The reaction energy levels have to stay somewhere between the Fermi level and the VB edge of the semiconductor. Note that the absolute electrode potential of SHE is estimated to be around −4.44 V at 25°C.29 The semiconductor band gap must be at least 1.6–1.7 eV (>1.23 eV) to drive the redox reactions after accounting for thermodynamic losses, but not greater than 2.2 eV for efficient solar light absorption and charge carrier generation.30 In addition, electron–hole pairs must be separated efficiently to prevent carrier recombination, and these free electrons and holes have to be transported smoothly to the cathode and anode to participate chemical reactions for producing hydrogen and oxygen, respectively. Some similarities between a PEC cell and a solar cell in terms of charge generation, separation and transport are witnessed.

7. Summary

In summary, there exist close connections between solar PV cells and other energy conversion cells, including thermoelectric, electrochemical and PEC cells in terms of their working principles. In these energy conversion cells, the working principles basically follow a simple picture. To be more specific, the following fundamental requirements are necessary: 1) there must exist mobile energy carriers, 2) a path loop with low resistance must be built up for carriers to move through, 3)
there must be available energy states/reservoirs for carriers at different energy levels along the loop, 4) there must be some energy input source(s) such as radiation and heat to pump carriers in situ from lower energy states to higher energy states or energy carriers occupy already higher energy states beforehand (for example, electrons in fuels) and 5) the load must be connected to the external circuit of the path loop so that work can be done on it as energy output (the rest of the energy is often wasted as heat). The picture is essentially a reflection of thermodynamics laws at large scale and quantum mechanics at small scale.

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REFERENCES


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