1. Introduction

Limited reserves of fossil fuels and their negative environmental effects impose significant problems in our energy security and sustainability. Consequently, researchers are looking for renewable energy sources, for instance solar energy, to meet the energy demands of a growing world population. However, terrestrial solar energy is a dilute resource per footprint area and is intermittent showing substantial variability depending on the season, time of the day, and location.

One strategy to overcome these drawbacks of solar energy is to concentrate and use it for cleaning and upgrading dirty fuels such as coal and other hydrocarbons or converting renewable feedstocks such as biomass into carbon-neutral solar fuels. In this way, the intermittent and dilute solar energy can be concentrated and stored as a chemical fuel which can be easily integrated to our existing energy infrastructure. These advantages of solar fuels produced with concentrated solar radiation make them an attractive solution in our quest for renewable and clean fuels. Figure 1 shows the energy potential and carbon emissions by most commonly used fuels along with solar hydrogen.

Most common and available methods for solar fuel production are thermolysis, cracking, reforming, gasification and through thermochemical cycles. All these methods require high temperatures to produce solar fuel. Therefore, in these methods, there are some qualities of the feedstock or the reactor that need to be satisfied to attain high temperatures and efficient solar fuel production. For instance, the physical size and porosity of the feedstock play an important role. As the surface area-to-volume ratio of the feedstock increases, more reaction sites will be available for the reaction to occur, which increases the process efficiency. The feedstock should also have a narrow bad gap to lower the energy requirement for chemical process. Additionally, the material on the reactor walls should have high optical absorption to increase the temperature of the reactor and withstand high temperatures, and the win-
dow material should have high transmissivity to let the solar energy in to the reactor. More detailed property requirements are given by Nowotny et al. [1].

This review chapter consists of four sections. Following the introduction, the second section “Concentrated Solar Fuel Production Methods” reviews the different routes of producing solar fuels according to the feedstock material used in the processes. These include (i) thermolysis of water, (ii) thermochemical cycles, (iii) cracking of gaseous hydrocarbons, and (iv) gasification and reforming of coal and biomass. These methods are compared with each other based on their temperature, pressure, thermodynamic efficiencies, and by-products. The third section “Concentrated Solar Reactors” provides a comprehensive review of different concentrated solar reactor designs reported in the literature. This section first reviews the current solar concentration methods and describes in detail the effects of concentrating factors on the heat flux and temperatures that can be achieved. Then, the section describes the design and basic principles of operation of different solar reactors, their applicability for the different methods described in the preceding section, and their temperature and pressure capabilities. Moreover, the section summarizes the reported solar to fuel conversion efficiencies of each design. Finally, the chapter ends with the conclusions and outlook of fuel production with concentrated solar energy outlining the challenges, new research directions and novel applications.
2. Concentrated solar fuel production methods

This section describes different methods of producing solar fuels according to the feedstock material used in the respective processes.

2.1. Thermolysis of water

The term “thermolysis of water” refers to the thermal decomposition of water molecules into hydrogen and oxygen gases. Historically, due to high availability and simple molecular form of water, researches on solar fuel production started with direct hydrogen production by thermolysis of water using solar energy as,

\[
\text{H}_2\text{O} \rightarrow \text{H}_2 + \frac{1}{2}\text{O}_2
\]

\[
\Delta H_{300K} = 286 \text{ kJ/mol}
\] (Equation 1)

The reaction given in Equation (1) is an endothermic process, i.e., it requires energy to break the bonds. However, breaking all the bonds in water molecules requires temperatures as high as 2500 K [2]. At lower temperatures, partial decomposition occurs. Although it is possible to reach 2500 K with concentrated solar energy, the reactor where this process takes place shows material issues related to high temperatures. Additionally, after the dissociation of water molecules, hydrogen and oxygen gases require separation at high temperatures in order to prevent back-bonding, i.e., reproduction of water molecules with an exothermic process. Some solutions include cooling the reactor down by injecting a gas or expanding these gases through nozzle at the end of the reactor [2, 3]. Other solutions include using double or tubular membranes or using multi-stage steam ejectors to lower the exit pressure [4]. However, these solutions further reduce the efficiency of the process, and thus no commercial plant using this technology exists.

2.2. Thermochemical cycles

Some metal oxides are reduced using solar energy since metals provide good storage and transport of energy, such as solar energy. Such metal oxides include, but not limited to ZnO, MgO, SnO₂, CaO, Al₂O₃, and Ce₂O₃. The reduction step of these metal oxides is generally followed by an oxidation step at lower temperatures than reduction step in order to produce solar fuel, mainly hydrogen. The reduced metal oxides generally react with CO₂ or steam. If steam is used in oxidation that step is called hydrolysis. The thermochemical cycles of different metal oxides are generally compared based on their temperature requirements for the reduction step, the reaction or dissociation rates and reaction kinetics.

ZnO is one of the most popular oxides mainly due to its abundance and relatively low temperature requirement for complete dissociation when compared to other metal oxides. Additionally, since ZnO is a simple metal oxide, it does not undergo multiple reactions before its full dissociation. The dissociation of ZnO occurs as according to,
The complete dissociation of ZnO to Zn requires temperatures higher than about 2300 K whereas, for instance, the dissociation of MgO as another simple metal oxide requires about 3700 K at atmospheric pressures [3, 5]. As in water thermolysis, partial dissociations can occur at lower temperatures. Although hydrolysis of zinc is exothermic as given by Equation (3), only 24% of Zn could be oxidized to produce H₂ at a reactor temperature of 800 K and an atmospheric pressure [6].

\[
\begin{align*}
\text{Zn} + \text{H}_2\text{O} &\rightarrow \text{ZnO} + \text{H}_2 \\
\Delta H_{300K} &\approx -62 \text{ kJ/mol}
\end{align*}
\] (3)

Figure 2 shows the overall process of hydrogen production from zinc-oxide.

Figure 2. Flowchart for thermochemical hydrogen production from zinc-oxide using concentrated solar energy [5].
As an alternative to ZnO reduction, Abanades et al. [7] proposed SnO$_2$ reduction using solar energy. Once the SnO$_2$ is reduced to SnO in gaseous form using solar energy at temperatures nearly 1600°C, hydrolysis of SnO with steam at about 550°C and ambient pressure takes place in another step to form hydrogen gas as,

$$\text{SnO}_2(g) \rightarrow \text{SnO}(g) + \frac{1}{2} \text{O}_2$$

$$\Delta H_{1873K} = 557 \text{ kJ/mol}$$

(4)

$$\text{SnO}(g) + \text{H}_2 \text{O}(g) \rightarrow \text{SnO}_2(g) + \text{H}_2$$

$$\Delta H_{773K} = -49 \text{ kJ/mol}$$

(5)

The advantages of SnO$_2$/SnO reduction when compared to ZnO/Zn reduction are that (i) the SnO$_2$-to-SnO conversion can be increased in Equation (4) by decreasing the pressure of the solar reactor which increases the overall conversion efficiency [7] (ii) SnO has higher melting and boiling points when compared to those of Zn, so that quenching rate of SnO is not as important as of Zn [7] (iii) in ZnO/Zn dissociation, Zn needs to be quenched rapidly below its condensation temperature to prevent recombination, while this is not the case with SnO$_2$/SnO system.

There are some other metals that can be reduced with faster reaction kinetics such as Ce$_2$O$_3$. However, the reduction of Ce$_2$O$_3$ to CeO$_2$ starts at temperatures higher than 2300 K [8, 9]. Full dissociation requires higher temperatures. This requirement of high temperatures creates some material limitations on the material of the reactor and increases the cost of the reactor significantly. Although there are some lab-scale prototypes of Ce$_2$O$_3$/CeO$_2$ solar reactor, it is not preferred due to these limitations and high cost.

Another research was also started with producing hydrogen gas from hydrogen sulfide, H$_2$S, as,

$$\text{H}_2\text{S} \rightarrow \text{H}_2 + \frac{1}{2} \text{S}_2$$

$$\Delta H_{300K} = -91.6 \text{ kJ/mol}$$

(6)

Hydrogen sulfide is a toxic by-product gas of sulfur removing process from natural gas, petroleum and coal. Thermal decomposition of hydrogen sulfide requires about 1800 K [10]. It is advantageous over the other metal oxide reduction processes discussed above since this thermochemical process is only a one-step process that does not require additional oxidation step to produce hydrogen. Additionally, the temperature requirement for dissociation is lower than that for the direct water thermolysis. However, the product gases need to be cooled down after the dissociation as in the water thermolysis or other metal oxide reduction processes [11]. Some studies showed that the temperature of reduction could be reduced to about 1500 K, and they showed that the reproduction of hydrogen sulfide is unimportant below 1500 K [3, 12, 13].
In general, the solar chemical process is a clean way to produce hydrogen without any carbon prints. Therefore, the hydrogen as a product of the solar chemical process can be used in fuel cells directly as it is pure. The solar chemical reduction step of the process produce nanoparticles with high surface area to volume ratio, e.g., Zn, SnO which also create additional reaction centers for the hydrolysis to occur [7]. Therefore, the oxidation or hydrolysis occurs fast due to high mass transport of gases in the solid phase [7]. As in the other dissociation processes, the products of the dissociation also need to be cooled in order to prevent re-oxidation. Sandia National Laboratories of US released a comprehensive report on the thermochemical cycle selection with initial selection for further research [14], and Table 1 summarizes the studied thermochemical cycles [15].

2.3. Cracking of gaseous hydrocarbons

The term “solar thermal cracking” or “solar cracking” is used for thermal decarbonization of natural gas or other hydrocarbons. As a result of cracking, hydrogen, carbon and other possible products are formed without CO₂ emissions. Therefore, this process is another method for clean fuel production. Cracking requires high temperatures of about 1500 K [16] that can be reached using concentrating solar collectors. For example, Maag et al. [17] tested a concentrated solar collector with a concentrating factor of 1720, and obtained a maximum temperature of 1600 K within the solar cavity reactor. In general, the advantages of solar cracking are the increase in value of feedstock using solar energy, pure and uncontaminated products and no CO₂ emission [16].

As being the simplest hydrocarbon and the main constituent of natural gas as given in Table 2, methane has been mainly considered for solar cracking. Chemical reaction of evolution of carbon black and methane is given in equation (7) [18, 19]. The kinetic mechanism of methane cracking at 1500 K and atmospheric pressure was proposed as [20, 21],

\[
2\text{CH}_4 \rightarrow \text{C}_2\text{H}_6 + \text{H}_2 \rightarrow \text{C}_2\text{H}_4 + 2\text{H}_2 \rightarrow \text{C}_2\text{H}_2 + 3\text{H}_2 \rightarrow 2\text{C}_{\text{(solid)}} + 4\text{H}_2
\]

(7)

Another important aspect of producing hydrogen and carbon black (solid carbon) is their market values. Hydrogen and carbon black have a market value of about $135 billion per year and between $7 and $11 billion per year depending on the grade of the carbon black in the world, respectively [22].

Under an EU project named SOLHYCARB, a 50-kW_{th} indirectly heated, cavity type solar reactor was developed for methane cracking [8]. Its 10-kW_{th} prototype was built and tested using natural gas, and 97% conversion was obtained with a maximum temperature above 2000 K under concentrated solar irradiation of 4 MW/m² [23]. The difficulties that prevent this technology to become commercial are mainly the cost of the reactor and the complicated flow pattern inside the reactors. For example, in order to prevent particle accumulation on the window, some inert gas is introduced to the reactor with high flow rates and pressures, or indirectly heated solar reactors are used which decreases the solar-to-fuel conversion efficiency and further increase the cost.
### High Temperature Cycles

<table>
<thead>
<tr>
<th>Cycle</th>
<th>Reaction Steps</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zn/ZnO</td>
<td>[ \text{Fe}_2\text{O}_4 \rightarrow 3\text{FeO} + \frac{1}{2}\text{O}_2 ]</td>
</tr>
<tr>
<td></td>
<td>[ 3\text{FeO} + \text{H}_2\text{O} \rightarrow \text{Fe}_2\text{O}_4 + \text{H}_2 ]</td>
</tr>
<tr>
<td>FeO/Fe$_3$O$_4$</td>
<td>[ \text{CdO} \rightarrow \text{Cd} + \frac{1}{2}\text{O}_2 ]</td>
</tr>
<tr>
<td></td>
<td>[ \text{Cd} + \text{H}_2\text{O} + \text{CO}_2 \rightarrow \text{CdCO}_3 + \text{H}_2 ]</td>
</tr>
<tr>
<td></td>
<td>[ \text{CdO}_3 \rightarrow \text{CO}_2 + \text{CdO} ]</td>
</tr>
<tr>
<td>Cadmium carbonate</td>
<td>[ \text{CdO} \rightarrow \text{Cd} + \frac{1}{2}\text{O}_2 ]</td>
</tr>
<tr>
<td></td>
<td>[ \text{Cd} + 2\text{H}_2\text{O} \rightarrow \text{Cd(OH)}_2 + \text{H}_2 ]</td>
</tr>
<tr>
<td></td>
<td>[ \text{Cd(OH)}_2 \rightarrow \text{CdO} + \text{H}_2\text{O} ]</td>
</tr>
<tr>
<td>Hybrid cadmium</td>
<td>[ \text{Mn}_2\text{O}_3 \rightarrow 2\text{MnO} + \frac{1}{2}\text{O}_2 ]</td>
</tr>
<tr>
<td></td>
<td>[ 2\text{MnO} + 2\text{NaOH} \rightarrow 2\text{NaMnO}_2 + \text{H}_2 ]</td>
</tr>
<tr>
<td></td>
<td>[ 2\text{NaMnO}_2 + \text{H}_2\text{O} \rightarrow \text{Mn}_2\text{O}_3 + 2\text{NaOH} ]</td>
</tr>
<tr>
<td>Sodium manganese</td>
<td>[ \text{Fe}_{3-x}\text{M}_x\text{O}<em>4 \rightarrow \text{Fe}</em>{3-x}\text{M}<em>x\text{O}</em>{4-y} + y\frac{1}{2}\text{O}_2 ]</td>
</tr>
<tr>
<td></td>
<td>[ \text{Fe}_{3-x}\text{M}<em>x\text{O}</em>{4-y} + y\text{H}<em>2\text{O} \rightarrow \text{Fe}</em>{3-x}\text{M}_x\text{O}_4 + y\text{H}_2 ]</td>
</tr>
<tr>
<td>M-Ferrite ($M = \text{Co}, \text{Ni}, \text{Zn}$)</td>
<td>[ \text{H}_2\text{SO}_4 \rightarrow \text{SO}_3 + \text{H}_2\text{O} + \frac{1}{2}\text{O}_2 ]</td>
</tr>
<tr>
<td></td>
<td>[ \text{I}_2 + \text{SO}_3 + 2\text{H}_2\text{O} \rightarrow 2\text{HI} + \text{H}_2\text{SO}_4 ]</td>
</tr>
<tr>
<td></td>
<td>[ 2\text{HI} \rightarrow \text{I}_2 + \text{H}_2 ]</td>
</tr>
</tbody>
</table>

### Low Temperature Cycles

<table>
<thead>
<tr>
<th>Cycle</th>
<th>Reaction Steps</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sulfur-iodine</td>
<td>[ \text{H}_2\text{SO}_4 \rightarrow \text{SO}_3 + \text{H}_2\text{O} + \frac{1}{2}\text{O}_2 ]</td>
</tr>
<tr>
<td></td>
<td>[ \text{SO}_3 + 2\text{H}_2\text{O} \rightarrow \text{H}_2\text{SO}_4 + \text{H}_2 ]</td>
</tr>
<tr>
<td>Hybrid sulfur</td>
<td>[ \text{Cu}_2\text{OCl}_2 \rightarrow 2\text{CuCl} + \frac{1}{2}\text{O}_2 ]</td>
</tr>
<tr>
<td></td>
<td>[ 2\text{Cu} + \text{H}_2\text{O} \rightarrow \text{H}_2 + 2\text{CuCl} ]</td>
</tr>
<tr>
<td></td>
<td>[ 2\text{CuCl}_2 \rightarrow 2\text{Cu} + 2\text{CuCl}_2 ]</td>
</tr>
<tr>
<td></td>
<td>[ 2\text{CuCl}_2 + \text{H}_2\text{O} \rightarrow \text{Cu}_2\text{OCl}_2 + 2\text{HCl} ]</td>
</tr>
<tr>
<td>Hybrid copper chloride</td>
<td>[ 2\text{CH}_4 \rightarrow \text{C}_2\text{H}_6 + \text{H}_2 \rightarrow \text{C}_2\text{H}_4 + 2\text{H}_2 \rightarrow \text{C}_2\text{H}_2 + 3\text{H}_2 \rightarrow 2\text{C}_2\text{H}_4 + \text{4H}_2 ]</td>
</tr>
</tbody>
</table>

Table 1. Summary of Thermochemical Cycles [15].
<table>
<thead>
<tr>
<th></th>
<th>CH$_4$</th>
<th>C$_2$H$_6$</th>
<th>C$_3$H$_8$</th>
<th>C$<em>4$H$</em>{10}$</th>
<th>CO$_2$</th>
<th>N$_2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methane</td>
<td>100</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Modified Algeria Gas</td>
<td>91.2</td>
<td>6.5</td>
<td>2.1</td>
<td>0.2</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Modified Groningen Gas</td>
<td>83.5</td>
<td>4.7</td>
<td>0.7</td>
<td>0.2</td>
<td>-</td>
<td>10.8</td>
</tr>
<tr>
<td>North Sea Gas</td>
<td>88.2</td>
<td>5.4</td>
<td>1.2</td>
<td>0.4</td>
<td>1.4</td>
<td>3.2</td>
</tr>
</tbody>
</table>

Table 2. Compositions of Natural Gas from Different Sources [20].

### 2.4. Gasification and reforming of coal and biomass

Gasification is a chemical process that converts carbonaceous feedstock into gaseous fuels under a controlled amount of oxygen and/or steam [24]. Main difference between gasification and combustion is that products in gasification have useful heating value. In gasification, pressure inside the gasifier is generally in the range from 20 to 40 bar, whereas methanol or ammonia synthesis requires 50 to 200 bar [25]. In addition, temperatures inside the gasifier is generally in the range from 1400 to 1700°C [25].

Pyrolysis is a thermochemical process that occurs before gasification, and it decomposes the complex hydrocarbons into smaller and less complex molecules in the absence of oxidizers. In pyrolysis, the yield of solar char can be maximized by slowing the heating rate, lowering the temperature or allowing a longer residence time [26]. On the contrary, a higher heating rate, a higher temperature, and a shorter residence time maximize the gas yield. Additionally, liquid yield at an intermediate temperature can be maximized by increasing the heating rate or minimizing the residence time. Tar is an undesired by-product of gasification and pyrolysis. It can cause condensation and consequent plugging, formation of aerosols and polymerization into more complex structures [26].

Gasification is an endothermic process and requires energy to occur. In case of conventional gasification, this energy is supplied from the partial combustion or gasification of feedstock which emits CO$_2$ to the atmosphere. Use of concentrated solar energy eliminates or reduces the CO$_2$ emission and utilizes the clean high-temperature gasification process. Additionally, fuel value of the feedstock is increased with solar gasification. For example, fuel value of coal can be increased by about 45% using solar coal gasification [27], and CO$_2$ emission can be reduced by about 30% when compared to conventional coal gasification [28].

Solar gasification of coal and other carbonaceous products is the process of converting these feedstock materials into some synthesis gas (syngas) which includes H$_2$, CO, CO$_2$ and water vapor using solar energy [29]. The gasification products can be further processed. For example, syngas can be processed to form methanol or ammonia or used in cement production, and lean gas can be combusted for heating or used in power stations to generate electricity.
Solar gasification can be performed using CO₂ or steam. In general, steam gasification of coal can be written as,

\[ \text{Coal} + a\text{H}_2\text{O} \rightarrow b\text{H}_2 + c\text{CO} + d\text{CO}_2 \]  

This process is endothermic and requires temperatures above 1000°C. Solar gasification of petcoke, coal and other carbonaceous feedstock started with directly irradiated solar reactors [31]. These designs have high reaction rates and kinetic and high fuel-to-product conversion. However, these reactors have problems with their aperture cover. As a cover, quartz window is commonly used to allow the concentrated solar power into the reactor. In directly irradiated reactors, quartz window has to withstand the high pressures inside the reactor and should not be covered with particles as the gasification occurs. As in the solar cracking, additional flows of inert gases are introduced into the reactor to prevent particle accumulation on the quartz window, but these additional flows introduce additional complexity and cost to the reactor [30]. In EU project SOLSYN [31], a 5-kW reactor prototype was built for solar coal gasification. The temperatures in this reactor could go up to 1700°C with the solar concentrating ratio of nearly 3000, but the general operation temperature was kept at 1220°C. The peak conversion efficiency was found to be 29% [31].

Similar to coal and other carbonaceous feedstock, biomass can also be gasified in solar reactors. Conventionally, gasification of biomass has been done using the exhaust gas of combustion of fossil fuels or biomass itself. Biomass includes demol wood, wood chips, sewage sludge, almond shells, straw, etc. If the biomass is used, nearly 30% of the initial biomass has to be combusted with oxygen to drive the gasification process due to the temperature requirement [15]. This temperature requirement varies between 600 – 1000°C [32]. Additionally, one of the other disadvantages of conventional biomass gasification is the formation of tar which blocks and clogs the equipment. There have been some efforts to eliminate the tar formation with proper selection of materials, operating conditions and the design of the gasifier [32].

Solar-assisted gasification of biomass has advantages over the conventional process. The main advantages are the elimination of tar formation, even at temperatures as high as 1200°C, and high and rapid conversion of biomass to syngas. At the National Renewable Energy Laboratory (NREL) of US, bluegrass was gasified with a maximum conversion of 95% and about 5% of the products were hydrocarbons, ash and char [33]. The resident times can be less than 5 seconds [33]. There is also a solar reactor design to combine solar biomass gasification and steam reformation [33].

### 3. Concentrated solar reactors

This section defines and compares different solar concentrators and gives examples of directly irradiated and indirectly heated solar reactors for the solar fuel production processes defined in the previous section.
3.1. Solar concentrators

There are two main types of concentrated solar collectors, categorized depending on their optical configurations. First type is parabolic trough systems in which there is an absorber tube in the focal line of parabolic reflectors. Linear Fresnel reflectors can also be included in this type of concentrated solar collectors. Second type is point focus solar collectors which include dish systems and heliostats. Dish systems have a solar receiver located in the focal point of the paraboloidal concentrator, and heliostats direct sun light to a solar receiver located at the top of a solar tower. Figure 3 shows the schematic of each solar collector type. Before going into details of each collector type, some terms need to be defined.

Figure 3. Solar concentrators, (a) parabolic trough, (b) linear Fresnel, (c) dish collectors, and (d) heliostats with solar tower [34].
Concentrating ratio is the ratio of the radiation intensity at the receiver by the radiation intensity received by the concentrator. Concentrating ratio plays an important role on the maximum achievable temperature at the receiver. Solar collector efficiency, $\eta$, is a product of Carnot efficiency, $\eta_{\text{Carnot}}$, and the receiver efficiency, $\eta_{\text{receiver}}$, as,

$$\eta = \eta_{\text{Carnot}} \cdot \eta_{\text{receiver}} = \left( 1 - \frac{T_o}{T_{\text{rec}}} \right) \left( \frac{\alpha \cdot G \cdot C - \varepsilon \cdot \sigma \cdot T_{\text{rec}}^4}{G \cdot C} \right)$$

(9)

where $T_o$ and $T_{\text{rec}}$ are the surroundings and receiver temperatures, respectively, $\alpha$ and $\varepsilon$ are the absorptivity and emissivity of the receiver, $G$ is the solar irradiation, $C$ is the concentrating factor and $\sigma$ is the Stefan-Boltzmann constant ($5.67 \times 10^{-8}$ W/m$^2$.K$^4$).

Figure 4 shows the solar collector efficiency as a function of receiver temperature when ambient temperature is 300 K, absorptivity and emissivity are both 1, and the solar irradiation is 1000 W/m$^2$. The figure shows that higher thermal efficiencies and higher receiver temperatures can be obtained with increasing concentrating ratios. Therefore, the selection of solar concentrators mainly depends on the temperature requirement of the application.

![Figure 4](http://dx.doi.org/10.5772/54057)

**Figure 4.** Theoretical solar reactor efficiency, $\eta$, as a function of receiver temperature, $T_{\text{rec}}$, for different concentrating ratios, $C$.

### 3.1.1. Parabolic-trough systems

Parabolic-trough collectors consist of several tubes interconnected in the focal line of highly reflective sheet material. These linearly connected tubes are generally referred as evacuated tubes since they consist of two concentric tubes whose annulus is vacuumed. The working fluid is circulated in the inner tube. The inner tubes are generally made of metals, and out-
side of the inner tube has selective coating to increase the absorption and decrease the heat loss. The outer tubes are generally made of glass, and they can also have selective coatings.

Applications of parabolic-trough collectors can be divided into two sections depending on the temperature of application. The low temperature applications, varies from 100°C to 250°C, include domestic hot water, space heating and heat-driven refrigeration [35]. Their concentrating ratios are between 15 and 20. The high temperature applications have temperatures up to 400°C, concentrating ratios of 20 to 30 [35]. Theoretical limit for concentrating ratio can go up to 100 [2]. These collectors are mainly used in power plants that are driven by steam. In the parabolic trough collectors, the pressure within the inner tube can reach 10 MPa.

There are other configurations of cylindrical absorber tubes which are not commonly used [36-38]. One type has a circulation tube inside the inner cylinder which carries the working fluid. This circulation tube is attached to the inner tube with a cylindrical fin. In this type, the absorbing surface is the fin itself, not the inner tube. Another type cylindrical absorber has a delivery tube inside the inner tube, and the working fluid delivered by this delivery tube fills the space in the inner tube. In another tube type, working fluid fills the annulus [37].

These parabolic-trough collectors and other cylindrical absorber tubes may have diffuse or specular reflectors at their back. Diffuse reflectors are generally flat surfaces that cover the entire back of arrays of tubular collectors. However, specular reflectors have parabolic surfaces, and they cover the back of only one cylindrical absorber. If specular reflectors are used, the absorber tubes have to be in the reflector’s focal line.

Line focus collectors are mounted with axes either in north-south or east-west direction. Hence, single axis tracking for this kind of collectors is sufficient to track the sun throughout the year. Collectors with axes in north-south direction track the sun from sunrise to sunset each day. Alternatively, collectors with axes in east-west direction track the sun seasonally. The spacing between each line of collectors in a solar farm with parabolic trough collectors is determined considering sun shadow line in the winter when the solar radiation comes with a larger azimuthal angle [38].

As an example, Nevada Solar One is in operation in Boulder City, Nevada, USA since 2007, covers 400 acres and has a capacity of 64 MW [39]. Figure 5 shows a picture of this power plant. Another and largest power plant with parabolic trough collectors is Solar Energy Generating Systems (SEGS) VIII-IX, built on approximately 240 acres and operating at 80 MW each near Harper Lake, California, USA. SEGS are also integrated with conventional natural gas turbines to operate at nights. It was reported that solar energy covers about 90% of the power production [40].

Another design is the absorber tubes with Fresnel reflectors. Fresnel reflectors that are mounted close to the ground direct and concentrate solar irradiation to the absorber tubes that are elevated at a higher level than reflectors. A secondary reflector on the back side of the absorber tube is also use to direct all the irradiation to the absorber tube. The main advantages of using Fresnel reflectors are that they have less wind load than the reflectors of
parabolic trough collectors since these reflectors are located at a lower position, and no requirement for vacuum in the absorber tubes and for rotating joints [42]. However, the main disadvantage of Fresnel reflectors is that they have lower concentrating ratios than parabolic trough collectors [43].

Figure 5. Parabolic-trough collectors in Nevada Solar One power plant [41].

In Calasparra, Spain, Novatec Biosol built a power plant with 28 rows of linear Fresnel reflectors that produces 30 MW of electrical power in an area of nearly 200 acres. Figure 6 shows a picture of this power plant. The power plant uses steam, and the temperature and pressure of the steam produced reach to 270°C and 55 bars, respectively [44, 45].

Figure 6. Linear Fresnel reflectors in the power plant Thermosolar Power Plant (PE2) in Spain [46].
3.1.2. Point focus collectors

Parabolic dish collectors concentrate sunlight to the focal point of the parabolic reflectors. These collectors have two-axis sun tracking system. In the focal point of the parabolic reflector, a working fluid is heated directly to a maximum temperature of about 1000 K [47]. This working fluid is generally used to drive a Stirling engine or a gas turbine to produce electricity. The typical parabolic dish collectors have a diameter of 5 to 10 m, and each can produce up to 0.4 MW [47, 48]. The concentrating ratios parabolic dish collectors vary between 1,000 and 10,000 [2]. The reflector is usually made of silver or aluminum coated glass. This kind of collectors can be used in applications with relatively low power requirement in remote areas.

One of the first examples of power plant using parabolic dish collectors was supposed to be Maricopa Solar Plant in Arizona, USA before its contractor company was announced bankruptcy in 2011. Figure 7 shows a picture of this power plant. This power plant consists of 60 parabolic dish collectors that heat the hydrogen to drive Stirling engines. The power plant has a capacity of 1.5 MW. This technology is not commercially operational and available in large scale power production.

Some solar thermal power plants use arrays of heliostats which are sun-tracking flat mirrors. These mirrors or heliostats with two-axis tracking system direct solar irradiation to the receiver located at the top of a tower on a concrete support. In order to direct the sunlight to the receiver with sufficient accuracy all the times, a motor drive system with a large gear reduction is necessary [50]. Due to the presence of the tower, these power plants are sometimes referred as power

Figure 7. Parabolic dish collectors in Arizona, USA [49].
tower systems. Concentrating ratios for these systems vary between 500 and 5000 [2], and the temperature at the receiver can exceed 2000 K depending on the concentrating ratio. These power plants can be used for converting solar energy to chemical energy, such applications include reduction of zinc oxide and coal gasification [51].

Throughout the technological development of heliostats, their sizes become larger and larger in order to decrease the production cost since their cost is a strong function of production rate. Although the initial development of heliostats started in 1975, one of the first prototypes of heliostats in 1980s by Sandia Labs, USA had an area of 37 m$^2$ [52]. Currently, Planta Solar (PS) 20 solar power plants use heliostats with each of their area as 120 m$^2$ [53]. Another improvement in the development of heliostats is the material choice. Glass mirrors with steel support structure are being replaced by silver polymer mirrors with silver-steel alloy structure in order to increase the structural durability and reduce the weight of heliostats [52]. Some designs also have circular mirrors instead of rectangular ones to reduce stress on the support structure.

The Crescent Dunes Solar Energy Project in Tonopah, Nevada, USA will be an example to the central tower power plants once it is completed late 2013. It will be built on approximately 1600 acres, and it will produce 110 MW of electrical power using molten salt as a phase changing storage medium [54]. As another example, PS 20 which is operational since 2009 has a cavity receiver at the top of a 165-m tower. Figure 8 shows the picture of this power plant. It heats up water in the cavity, and steam reaches an outlet temperature of maximum 550 K. This solar power plant consists of 1255 heliostats with a total area of 30 acres. The solar power plant is backed up with natural gas burnt conventional turbine, and the total power production is 20 MW. PS 20 power tower is cooled with water which is generally replaced by air cooling if the power plant is built on deserts due to lack of water resources. The cooling is necessary for the materials used in the power towers.

![Figure 8. PS 10 (back) and PS 20 (front) solar thermal power plants with heliostats with solar towers [55].](http://dx.doi.org/10.5772/54057)

To reach the necessary temperatures for the solar fuel production methods given in the previous section, tower or dish type collectors should be used. For further reference, a review paper on the volumetric receivers for the concentrating solar thermal power plants discussed different designs from the projects of the last 3 decades [56]. Another good review was done on comparison of parabolic trough, dish systems, solar towers and tubular systems with Fresnel reflectors by Pavlovic et al. [57].
3.2. Solar reactors

In this section, different solar reactors that were designed for fuel production using concentrated solar energy are discussed and compared in terms of their operating conditions and design parameters. Depending on the reactor design, these reactors were fed feedstock with or without solid particles to perform hydrolysis, cracking, gasification, etc. These particles not only allowed a more uniform temperature distribution inside the reactor, but also helped the reactor to reach higher temperatures faster. These particles also acted as additional reaction sites due to their high surface to are to volume ratios. In some designs, feedstock was diluted with some inert gas, such as Argon, to increase the produced fuel yield. An auxiliary gas was also fed to prevent particle deposition on the window surface. In some designs, reactors were supplemented with a cooling system for products to prevent them recombine.

In general, the solar energy is transformed into thermal energy in the structure of volumetric receivers. In some designs, solar energy directly heats the feedstock in the reactor which is referred as directly irradiated solar reactors. Additionally, some of these receivers have a porous metal or ceramic absorber to be heated by solar energy. Metal absorbers can be heated up to 1000°C whereas SiC absorbers can reach 1500°C as maximum temperatures. Then, this thermal energy is transferred to a working fluid that passes through the porous absorber. This kind of reactors is referred as indirectly heated solar reactors.

3.2.1. Directly irradiated solar reactors

In this section, examples of directly irradiated solar reactors are presented with their design parameters, temperature and pressure allowances, their power outputs and their solar fuel production rates. These examples are selected to give a wide range of applications and designs.

Maag et al. [17] tested a 5 kW-prototype of a solar reactor seeded with particles for thermal cracking of methane. The cylindrical reactor was 200 mm in length and 100 mm in diameter. It had a 60 mm aperture area. The concentrator was covered with a 240 mm-diameter quartz window as shown in Figure 9. In their experiments, they used a sun-tracking parabolic concentrator that has a diameter of 8.5 m and could reach a concentrating factor up to 5000 suns. They tested the reactor in the temperature range from 1300 to 1600 K with a concentrating factor of 1720 suns. They varied the volume fraction of carbon in the range of 0 to 7.2x10⁻⁵ and gas inlet flow rate in the range of 8.6 to 15.6 l/min. They reported maximum methane-to-hydrogen conversion of 95% at a residence time less than 2 seconds, and an experimental solar-to-chemical energy conversion efficiency of 16%, whereas their theoretical prediction of the same conversion efficiency was 31%.

Yeheskela and Epstein [58] developed and tested 10-kW particle-seeded solar chemical reactor for producing hydrogen and carbon nanotubes from methane. They used iron pentacarbonyl and ferrocene as catalysts to produce multi-walled carbon nanotubes. The reactor was 300 mm in length, and the quartz window which covered the reactor as shown in Figure 10 was 200 mm in diameter. Additionally, He was used as a screen protector gas to eliminate the particle deposition on and near the window, and N₂ was used as a tornado generator gas. The average reported temperature within the reactor core was 1450°C.
Figure 9. Schematic of design of Maag et al. [17].

Figure 10. Schematic of design of Yeheskela and Epstein [58].
Abanades and Flamant [59] designed nozzle-type directly-irradiated solar reactor for methane cracking. They used a graphite nozzle with an inner diameter of 10 mm and a length of 65 mm. The schematic of the proposed design is shown in Figure 11. The reactor walls were made of stainless steel, and they were water cooled for their durability. Additionally, the products were also cooled to eliminate recombination of products [60]. The obtained conversion of methane to hydrogen exceeded 95% in molar basis, while the rest of the by-products were \( \text{C}_2\text{H}_2 \), \( \text{C}_2\text{H}_4 \) and \( \text{C}_2\text{H}_6 \). With a direct normal irradiation of 980 W/m\(^2\), the temperature of graphite nozzle had a maximum temperature of 1385°C, while their model estimated the maximum wall temperature as 1890 K.

Klein et al. [61] investigated the performance of directly irradiated solar methane cracking process with and without \( \text{CO}_2 \) present in the reactor. The reactor, as shown in Figure 12, had a diameter of 160 mm, a length of 266 mm and an aperture diameter of 60 mm. The gas exit temperatures with \( \text{CO}_2 \) only (no methane) in the reactor were in the range from 1000 to 1250°C. Additionally, the exit temperatures were in the range from 1100 to 1450°C with \( \text{CO}_2 \) and methane, when the \( \text{CO}_2 \) and methane molar ratio were varied from 1:1 to 1:6. Overall, the experimental results were similar to the studies where no \( \text{CO}_2 \) was introduced into the reactor. Moreover, when the reactor was fed with \( \text{CO}_2 \) and carbon black and the gas exit...
temperature reached 1000°C, 20% of carbon particles reacted with CO$_2$. When the exit gas temperature was increased to 1350°C, about 70% of the carbon particles reacted with CO$_2$.

Z’Graggen et al. [62] designed a 5-kW prototype reactor for steam-gasification of petroleum coke using concentrated solar energy. The reactor had a 5-cm diameter aperture which was covered by 3-mm-thick quartz window. The window was cooled by oil, and swept by an inert gas to prevent particle accumulation on the window. The solar concentrating ratio was about 5000, and the maximum temperature in the reactor was about 1800 K. The walls of the cavity were covered with Al$_2$O$_3$ and insulated from the backside with Al$_2$O$_3$-ZrO$_2$. Figure 13 shows the design of Z’Graggen et al. [62]. As a result of the steam-gasification of petroleum coke, H$_2$, CO, CO$_2$ and CH$_4$ were produced with a chemical conversion ratio of 87%. The overall solar-to-chemical conversion efficiency was about 9%.
3.2.2. Indirectly heated solar reactors

Directly irradiated solar reactors work with high conversion efficiencies. However, they have problems, such as accumulation of particles on the window. In order to overcome this problem and the need for inert gas feeding, indirectly heated solar reactors are alternatively introduced. This section gives some design examples of indirectly heated solar reactors for different solar fuel production methods.

Gordillo and Belghit [63] modeled the reaction kinetics without pyrolysis using finite volume analysis in a two-phase biochar solar gasification reactor with a bubbling fluidized bed as shown in Figure 14. Bubbling was used to introduce fluidizing gases to the reactor. They found that concentrating solar energy and high gas flow rates affected the temperature distribution within the packed bed. Therefore, a uniform temperature distribution could not be obtained which adversely affected the reaction kinetics. Additionally, they showed that although energy conversion efficiency ($\eta$), defined as in Equation (10), could be as high as 55%, it decreased with increasing the steam velocity or the bed temperature [64].

$$\eta = \frac{m_{\text{product}} \cdot LHV_{\text{product}}}{Q_{\text{solar}} + m_{\text{feedstock}} \cdot LHV_{\text{feedstock}}}$$

where $m$ and $LHV$ refer to the mass flow rate and lower heating value, $Q_{\text{solar}}$ is the solar irradiation, subscripts product and feedstock denote gaseous products and fed feedstock, respectively.

According to Hathaway et al. [65], problems with the preceding reactor designs had poor heat transfer characteristics [63], formation of ash and tar which block the radiative heat transfer and insulate the reaction zone, and intermittency of solar energy. Hathaway et al. [65] investigated the effects of using molten salt on the reaction kinetics in solar gasification of biomass. For the analysis of pyrolysis which occurs before gasification, they prepared tablets using microcrystalline cellulose, and for the analysis of steam gasification, they used tablets of wood charcoal powder. They carried out a series of experiments in the temperature range from 1100 to 1250 K to show the effects of molten alkali carbonate salts (lithium, sodium and potassium carbonate) on reaction rates using the experimental setup shown in Figure 15. They showed that introducing molten salts increased the rate of pyrolysis by 74% and increased the rate of gasification by more than an order of magnitude since molten salts acted as a heat transfer medium for gasification which ended up with more uniform temperature distribution within the solar reactor. On the contrary to the other studies, the catalytic effect of molten salt on pyrolysis was not observed for the reason that pyrolysis happened rapidly, and then gasification occurred. However, the catalytic effect of molten salt on gasification was observed. Introducing the molten salt increased the pre-exponential factor (i.e., rate of reaction in steady state process) by 24.4 times and increased the activation energy by about 4%. Additionally, using molten salt avoids the tar production (as a by-product of uncatalyzed gasification, occurs especially on startup), and molten salts can act as an energy storage unit to overcome the intermittency effect of solar energy.
Figure 14. Model of Gordillo and Belghit [63].
Rodat et al. [66] developed a 10 kW tubular reactor prototype, which was indirectly heated, for methane cracking as shown in Figure 16. They used a graphite cubic cavity as a receiver and a quartz window. The quartz window was swept by nitrogen which prevented O\textsubscript{2} to enter the cavity. The reactor reached 2070 K, and the products included C\textsubscript{2}H\textsubscript{2} with maximum mole fraction as 7%. As given in Equation (7), C\textsubscript{2}H\textsubscript{2} is the last step of H\textsubscript{2} and carbon black evolution. The graphite cavity was purged by N\textsubscript{2}. For this configuration, the reactor required about 4000 seconds to reach the required temperature of 1800 K when the experiment started at 300 K under the direct normal irradiance of 1000 W/m\textsuperscript{2} [67].

**Figure 15.** Experimental setup of Hathaway et al. [65].

**Figure 16.** Schematic of design of Rodat et al. [66].
Lichty et al. [69] designed and analyzed the thermal characteristics of a cavity reactor prototype for solar-thermal biomass gasification as shown in Figure 17. The maximum recorded temperature was 1660 K on the central tube under 7.5 kW power input. They quantified the reacted biomass based on CO and CO$_2$ as these gases showed the ratio of reactants underwent a complete reaction, and the authors reported an average biomass-to-CO and CO$_2$ conversion as 58.4%. The residence time was about 4 seconds. They also compared the syngas production of grass and lignin pyrolysis and cellulose gasification using mass spectrophotometer.

![Figure 17](Image)

*Figure 17. Reactor design of Lichty et al. [69].*

In the design of the German Aerospace Center for directly irradiated solar reactor to reform natural gas is given in Figure 18, porous ceramic absorber coated with Rh catalyst was used [12]. A concave quartz window was also mounted on the concentrating solar collector [12]. The operating conditions were chosen as 1400°C and 3.5 bars, and a volumetric flow rate of 3.8 l/min with 5% methane in argon [70, 71].
Maag et al. [72] simulated the performance of a 10 MW commercial-size reactor. The reactor consisted of four graphite absorber tubes with an outer diameter of 24 mm placed in a 0.2 m-cubic graphite cavity as shown in Figure 19. The graphite cavity had an aperture of diameter 9 mm which was covered by quartz window. They predicted 100% methane-to-hydrogen conversion when flow rate of methane was 0.7 kg/s at a reactor exit temperature of 1870 K. Spectral properties of quartz window were estimated using a band model, and view factors were calculated using Monte Carlo ray-tracing method. The energy balance for the overall system was solved with finite volume method. The results showed that it was possible to increase solar-to-chemical energy conversion efficiency from 42% to 60% when the outlet temperature was lowered to 1600 K and, subsequently, the methane flow rate was doubled, but then quality of carbon black as a product would be poorer.

Figure 18. Design of the German Aerospace Center for natural gas reforming [12].

Figure 19. Schematic of design of Maag et al. [72].
Osinga et al. [73, 74] designed 5-kW indirectly heated solar reactor for the reduction of ZnO. There were two different versions of the reactor. First one had the inner cavity made of graphite, and the second one had the inner cavity made of SiC. Figure 20 shows the second type, reactor with SiC absorber. Both reactors could reach temperature of 1700 K in about 80 minutes after the solar energy was input to the reactor. The reactor with graphite absorber had a vacuum pressure of 10 mbar whereas the pressure inside the reactor with SiC absorber was kept at 1 bar. ZnO and C mixture was reduced to Zn, CO and CO$_2$ from which Zn can be reacted with water to produce ZnO and H$_2$ as in Equation (3) [75].

Figure 20. Reactor design of Osinga et al. [73].
The reactor with SiC absorber was scaled up to a capacity of 300 kW and operated by Wieckert et al. [76] under the EU project, called SOLZINC. They used solar tower with heliostats to deliver concentrated solar power of 300 kW to the volumetric reactor. The second cavity where the mixture of ZnO and C was present reached to maximum temperature of nearly 1500 K, and overall ZnO-to-Zn conversion reached 95% with a Zn production of 50 kg/h.

Piatkowski et al. [31, 77] designed a 5-kW indirectly heated solar reactor with packed bed for solar steam-gasification. Figure 21 shows the reactor design of Piatkowski et al. The authors used different carbonaceous feedstock, such as African coal, sludge and charcoal. Beech charcoal gave the maximum solar-to-chemical conversion efficiency of 29% at a temperature of nearly 1500 K. The solar reactor has two cavities separated an emitter plate made of SiC-coated graphite. The aperture diameter was 6.5 cm which was followed by a 3-mm-thick quartz window. The walls of the lower cavity was also covered by SiC, and insulated by Al₂O₃-SiO₂. The reactor received the concentrated solar energy with concentrating ratios up to 3000, and the maximum temperature measured in the upper cavity was about 1700°C.

![Figure 21. Reactor design of Piatkowski et al. [31, 77].](image-url)
Summary of the operating conditions of the discussed designs for methane cracking is given in Table 3.

<table>
<thead>
<tr>
<th>Reference</th>
<th>Maximum Temperature (°C)</th>
<th>Inlet CH&lt;sub&gt;4&lt;/sub&gt; Dilution (%vol.)</th>
<th>Reactor Dimensions (mm)</th>
<th>Aperture Diameter (mm)</th>
<th>Inlet Flow Rate (l/min)</th>
<th>Catalytic or Fed Conversion</th>
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<td>Maag et al. [17]</td>
<td>1327</td>
<td>6-30 (in Argon)</td>
<td>100 (diameter) 200 (length)</td>
<td>60</td>
<td>8.6-15.6</td>
<td>Carbon black seeded</td>
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<tr>
<td>Yeheskela and Epstein [58]</td>
<td>1450</td>
<td>98 (in catalysts)</td>
<td>200 (diameter) 300 (length)</td>
<td>200</td>
<td>5-9.7</td>
<td>Flow with Fe(CO)&lt;sub&gt;5&lt;/sub&gt;, Fe(C&lt;sub&gt;5&lt;/sub&gt;H&lt;sub&gt;5&lt;/sub&gt;)&lt;sub&gt;2&lt;/sub&gt;</td>
</tr>
<tr>
<td>Abanades and Flamant [59, 60]</td>
<td>1110</td>
<td>11-20 (in Argon)</td>
<td>10 (diameter) 65 (length)</td>
<td>10</td>
<td>0.9</td>
<td>No particle feeding</td>
</tr>
<tr>
<td>Klein et al. [61]</td>
<td>1471</td>
<td>10-24 (in Argon or CO&lt;sub&gt;2&lt;/sub&gt;)</td>
<td>160 (diameter) 266 (length)</td>
<td>60</td>
<td>37-60</td>
<td>Carbon black seeded</td>
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<tr>
<td><strong>Indirectly Heated Solar Reactors</strong></td>
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<tr>
<td>Rodat et al. [66]</td>
<td>1800</td>
<td>10-20 (in Argon)</td>
<td>18 (tube diameter) 200 (cube side)</td>
<td>90</td>
<td>-</td>
<td>No particle feeding</td>
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<tr>
<td>German Aerospace Center [12]</td>
<td>1400</td>
<td>5 (in Argon)</td>
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<td>24 (tube diameter) 200 (cube side)</td>
<td>9</td>
<td>10-48</td>
<td>No particle feeding</td>
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</tbody>
</table>

Table 3. Operating Conditions of Different Reactor Designs for Methane Cracking.

4. Conclusions and outlook

The research to find an alternative fuel to fossil fuels is led by how the new technologies are economically competitive with the fossil fuel technologies, rather than their efficiencies. However, the economical aspect of fossil fuels should also include the cost for CO<sub>2</sub> emissions or sequestration of CO<sub>2</sub> when fossil fuels are compared to solar fuels since solar fuels have no CO<sub>2</sub> emission. Furthermore, as the fossil fuels deplete and the demand for fossil fuels will exceed their production, their prices will be subjected to significant increase. In this way, the investors in fuel or electricity production sector would see solar fuels as an alternative to fossil fuels with the current level of both technologies. As an outcome, the world
would become a more sustainable environment with reduced atmospheric CO\textsubscript{2} level and less pronounced risk for global warming.

It should also be noted that the solar fuel production methods introduced in this chapter are at different levels of maturity. For instance, most of the thermochemical cycles are in laboratory and research scale, whereas gasification and reforming processes are in fully operational or pilot stage. To give some examples on solar fuel production, the targets and predictions of the US Department of Energy (DOE) [78] for both cost and process efficiency are summarized in Table 4 for the ZnO/Zn thermochemical cycle. The predictions show that it is possible and feasible to meet the DOE efficiency and 2015 cost targets. However, the cost target of DOE in 2025 is a challenging objective. The main constituent of cost in thermochemical solar production is the plant capital cost, i.e., cost for heliostats and solar tower, rather than the direct cost for the process. Although process efficiencies are reported and predicted as given in Table 4, the overall solar-to-fuel conversions are still low, less than 10% [1].

<table>
<thead>
<tr>
<th></th>
<th>by 2015</th>
<th></th>
<th>by 2025</th>
<th></th>
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</thead>
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<td></td>
<td>DOE Target</td>
<td>Prediction\textsuperscript{1}</td>
<td>DOE Target</td>
<td>Prediction\textsuperscript{1}</td>
</tr>
<tr>
<td>Cost ($/gge\textsuperscript{1})</td>
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<td>6.07</td>
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<td>4.18</td>
</tr>
<tr>
<td>Process Efficiency (%)</td>
<td>30</td>
<td>35</td>
<td>35</td>
<td>42</td>
</tr>
</tbody>
</table>

\textsuperscript{1}gge refers to gallon-of-gas-equivalent.

\textsuperscript{1}Based on predicted ZnO-to-Zn conversions of 70% in 2015 and 85% in 2025.

| Table 4. Targets of Department of Energy of US [14] and predictions [78] for cost and efficiency for ZnO/Zn thermochemical cycle. |

In summary, the advantages of solar fuels include:

- Energy content or heating value of feedstock is increased by converting it to another form, solar fuel.

- Producing storable and transportable fuel which is not possible if solar energy is directly used. Thus, eliminates the intermittency problem of solar energy.

- Solar fuels are clean and sustainable. The thermochemical cycles and thermolysis of water that are used to produce solar fuels have no CO or CO\textsubscript{2} emissions. However, carbon emission occurs for the gasification or reforming of carbonaceous feedstocks. If these feedstocks are biomass cultivated with CO\textsubscript{2} from the atmosphere, they are carbon neutral.

On the other hand, these are not mature technologies and still suffer from technical challenges which form the basis for future research including:

- High temperatures needed for solar fuel production processes. High temperatures can be reached with high concentrating ratios. However, high concentrating ratios bring high cost to the system, and high temperatures restrict the material choice.
Recombination of product gases, especially in thermochemical cycles, is a significant problem. This recombination significantly decreases both the process and overall solar-to-fuel efficiency.

Quenching is introduced to products in order to reduce the recombination. However, quenching adds additional cost and complexity to the reactor and the process management. For some solar thermochemical processes, membranes are also required to separate product gases.

Particle accumulation on the window of the reactor is a problem in directly irradiated solar reactors. This problem can be eliminated by introducing an inert gas with high flow rates to the reactor which further complicates the management of reaction in the reactor. Another solution is to heat the reactor indirectly which reduces the solar-to-fuel efficiency.

Multiple-step chemical reactions are needed to produce hydrogen in most of the thermochemical cycles. More reactions add further components to the system which increase the cost and the management of the overall fuel production process.

CO and CO₂ formation can be noteworthy in case of solar gasification and reforming of carboneous feedstock, although solar fuels are accepted as clean fuels.

These drawbacks of the solar fuel production prevent the technology to be converted to large scale commercially available power plants. However, solar fuel production processes are thermodynamically efficient, favorable developments to increase the feedstock’s heating values with the unlimited free solar energy. Therefore, in a long-term prospect, solar fuel production is a promising technology that needs significant research efforts for efficiently producing clean and sustainable fuels.

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References


