Biogas Reforming to Syngas: A Review

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Interest in novel uses of biogas has increased recently due to concerns about climate change and greater emphasis on renewable energy sources. Although biogas is frequently used in low-value applications such as heating and fuel in engines or even just flared, reforming is an emerging strategy for converting biogas to syngas, which could then be used to obtain high-value-added liquid fuels and chemicals. Interest also exists due to the role of dry, bi-, and tri-reforming in the capture and utilization of CO2. New research efforts have explored efficient and effective reforming catalysts, as specifically applied to biogas. In this paper, we review recent developments in dry, bi-, and tri-reforming, where the CO2 in biogas is used as an oxidant/partial oxidant. The synthesis, characterization, lifetime, deactivation, and regeneration of candidate reforming catalysts are discussed in detail. The thermodynamic limitation and techno-economics of biogas conversion are also discussed.

INTRODUCTION

Research on renewable energies has been carried out due to concerns related to global warming and depletion of petroleum-based resources. Renewable power sources have entered the energy markets due to their promising sustainable development, reduced environmental pollution, and enhanced domestic energy supply (NREL, 2013). Biogas, which has been considered as a sustainable and renewable gaseous fuel (EPA, 2014), is suitable for direct conversion into syngas (i.e., a mixture of H2 and CO) via a reforming. The removal of CO2 is not required, which avoids substantial cost related to gas purification.

Biogas dry reforming could make a significant contribution to future hydrogen production, as H2 allows for highly efficient generation of electricity in vehicles and fuel cell plants (Charisiou et al., 2019). However, biogas dry reforming involves a highly endothermic reaction because CH4 and CO2 have high bond-dissociation energies (435 and 526 kJ/mol, respectively). Thermodynamically, an increase in the reaction temperature favors biogas dry reforming. In addition, a number of simultaneous side reactions (e.g., reverse water–gas shift [rWGS] reaction) can influence the catalytic performance and reactant conversions (Charisiou et al., 2019). Another issue for biogas dry reforming is coke formation through the methane decomposition and/or Boudouard reaction. The coke deposited on the outer and/or inner catalyst surfaces can cause catalyst deactivation and plugging of the reforming reactor (Jabbour et al., 2019). Linde (News, 2016) had built a pilot reformer to test dry reforming technology, and there could be others at similar stage. However, there are not any commercial plants using CO2 as a feed in the reformer to the best of our knowledge.

Compared with biogas dry reforming, the steam reforming reaction (without the CO2 separation step) is less energy intensive. The steam reforming of CH4 has been an established technology. The addition of steam to the biogas dry reforming process can lead to significant mitigation of coke deposition on the catalyst. Combined CO2 and steam reforming of methane is referred as bi-reforming or “steam biogas reforming” (Equations 1 and 2) (Roy et al., 2018). However, the O2 (if a high content) has to be separated from the biogas to allow the bi-reforming process to take place.

Tri-reforming is a combination of three methane reforming processes: carbon dioxide reforming (Equation 1), steam reforming (Equation 2), and partial oxidation reforming (Equation 3).

\[
\begin{align*}
\text{CH}_4 + \text{CO}_2 &\leftrightarrow 2\text{CO} + \text{H}_2 \quad \Delta H^\circ = 247 \text{kJ/mol} \\
\text{CH}_4 + \text{H}_2\text{O} &\leftrightarrow \text{CO} + 3\text{H}_2 \quad \Delta H^\circ = 206 \text{kJ/mol} \\
\text{CH}_4 + \text{1/2O}_2 &\leftrightarrow \text{CO} + 2\text{H}_2 \quad \Delta H^\circ = -36 \text{kJ/mol}
\end{align*}
\]

In addition, there are other reactions taking place during the tri-reforming process (Walker et al., 2012).

\[
\begin{align*}
\text{CO} + \text{H}_2\text{O} &\leftrightarrow \text{CO}_2 + \text{H}_2 \quad \Delta H^\circ = -411 \text{kJ/mol} \\
\text{C} + \text{H}_2\text{O} &\leftrightarrow \text{CO} + \text{H}_2 \quad \Delta H^\circ = 131 \text{kJ/mol}
\end{align*}
\]
C + O₂ → CO₂  ∆H° = −394 kJ/mol  (Equation 6)

The tri-reforming technology has numerous advantages: the environmental advantage of carbon dioxide reforming, the high economical advantage of steam reforming due to the utilization of CO₂, no need to remove O₂, coke inhibition due to the presence of H₂O and O₂, high energy efficiency of exothermal partial oxidation reaction, and relatively lower energy consumption compared with dry and bi-reforming (Kozonoe et al., 2019; Lino et al., 2019; Paladino Lino et al., 2019; Yoo et al., 2015). However, there is a risk of oxidation of catalysts due to the oxygen present in the feed (Majewski and Wood, 2014). Despite the fact that tri-reforming is energy efficient, avoiding the oxidation of catalysts during the tri-reforming process is a challenge.

The syngas produced from methane reforming could be used to make methanol, ethanol, hydrogen, gasoline, diesel, jet fuel, dimethyl ether, and other chemicals (Gangadharan et al., 2012; He et al., 2019; Vita et al., 2014). For example, hydrogen is an important energy carrier due to its highly specific energy density, abundance, and eco-friendliness (Yoo et al., 2015). The past 20 years have seen an exponential increase in technical articles and citations on syngas production from biogas reforming, including dry reforming, bi-reforming, and tri-reforming (Figure 1).

Catalysts for reforming must be thermally stable and coke resistant due to the endothermic nature of key reactions (Anchieta et al., 2019; Singha et al., 2016b). Conventional supported nickel catalysts tend to deactivate due to coke formation and metal sintering (Yoo et al., 2015). Noble metal catalysts generally have higher coke resistance than conventional nickel catalysts during the methane reforming process (Jiang et al., 2007). Nickel is less expensive than noble metals. Well-developed mesoporous nickel-based catalysts exhibited excellent activity due to high mass transfer rate of reactants (Yoo et al., 2015). The popular catalysts for biogas tri-reforming use are nickel supported by a wide range of materials including CeO₂, ZrO₂, and Al₂O₃ (Kozonoe et al., 2019).

This review focuses on the reforming of biogas to syngas; the comparison between tri-reforming and other biogas reforming reactions (i.e., dry and bi-reforming); and recent research of biogas reforming at low temperatures. The properties of reforming catalysts, catalyst lifetime, catalyst dimension, and catalyst regeneration are discussed. The composition of biogas, thermodynamic limitations or consideration of biogas reforming, application of syngas produced from biogas, techno-economics of biogas conversion, challenges, and future directions will be discussed as well.

**BIOGAS PRODUCTION, COMPOSITION, AND IMPURITIES**

Biogas can be produced in many ways, and the incidence of its generation is growing fast. The feedstocks for biogas generation include farm-based wood residues, crops, livestock, and manure (Lin et al., 2018; Sarker...
et al., 2018). The most common source is anaerobic digestion of biomass, either in commercial digesters, landfills, or wastewater treatment plants. Biogas compositions (CH4: 25%–75%, CO2: 7%–60%) are affected by the type of waste source utilized (Vita et al., 2015). World biogas production increased by 357% to 59 billion m3 from 2000 to 2014 (Scarlat et al., 2018). Approximately 78 biogas plants were built in Japan by 2012 (Sarker et al., 2018). The potential volume of biogas production from farm animal manure and slaughterhouses in Malaysia was approximately 49.5 million m3/year. The potential volume of biogas production from slaughterhouse waste and livestock manure in Iran was 8,600 million m3/year (Zareei, 2018). The increase of biogas production is encouraged by the sustainable, economic, climate, and environmental benefits. However, the distributed nature of waste/biomass that generates biogas, which leads to small-scale reforming reactors, is a major challenge of biogas application. The supply chain logistics for biogas production faces challenges in feedstock hub location, road network design to transport feedstock to hub, and site selection of biogas reactors (Sarker et al., 2019). The location and allocation of a biogas plant is an issue because transportation cost is affected by distance (Sarker et al., 2018). Optimum biogas plant location selection, efficient management, and appropriate planning are crucial to reduce supply chain cost.

Table 1 shows the typical composition and impurities of biogas from different sources. Biogas contains CH4, CO2, N2, O2, H2O, and trace amounts of other gases (e.g., H2S, NH3, and H2). Biogas composition differs based on the feedstocks used, but CH4 and CO2 are the main components. The ratio of CH4 to CO2 is mainly in the range of 1.0–2.3. The CH4 to H2O ratios and CH4 to O2 ratios in biogas are typically higher than 3 and higher than 20, respectively. The large variance of the CH4, CO2, H2O, and O2 content in biogas from different sources is a major challenge for biogas reforming. The reaction conditions in the anaerobic digestion can be controlled to obtain appropriate ratios of components in biogas. The biogas composition can be influenced by the feed composition (Bierer et al., 2019), temperature, and organic loading rate (Bi et al., 2019). In addition, a large number of experiments (Gao et al., 2020; Hossain et al., 2019; Saebia et al., 2019) examining the reforming of biogas with different component ratios at different reaction conditions can be carried out. Artificial intelligence methods can be used on the obtained data to predict what reaction condition (e.g., temperature, pressure, and gas hourly space velocity [(GHSV)]) is needed for the efficient
reforming of biogas with a specific composition to obtain high reactant conversions and the desired H₂/CO ratio (Alsaffar et al., 2020). The fast development of artificial intelligence in recent years provides an opportunity to assist the optimization of biogas utilization and application.

Biogas without purification contains small amounts of impurities such as siloxanes, NH₃, and H₂S (Elsayed et al., 2017). Volatile methyl siloxanes consist of silicon, oxygen, and methyl groups. Manufacturers have started to set limits on allowable siloxanes in the feed, considering the harmful effect of siloxane decomposition to biogas processing reactors (Elsayed et al., 2017). However, these impurities can limit the biogas application and generate harmful environmental emissions (Abdullah et al., 2018). To utilize the biogas efficiently, undesired components need to be removed (Sun et al., 2015). The H₂S can be adsorbed using porous materials such as activated carbon and ZnO-impregnated zeolite (Abdullah et al., 2018). Other materials such as Sulfatreat and silica gel can be used to remove H₂S and NH₃. Elsayed et al. (Elsayed et al., 2017) studied the effect of silicon poisoning on biogas dry reforming performance, finding that the addition of silicon decreased CH₄ and CO₂ conversions by up to >80% with increased deposited silica amounts. Siloxanes can irreversibly decompose to silica, which can become deposited on the equipment. It is suggested that the biogas should be scrubbed of impurities, including siloxanes and sulfur species, to avoid the frequent replacement of catalysts (Elsayed et al., 2017). Some biogases also include nitrogen, which is an inert gas during the biogas reforming. It is not easy to separate or remove nitrogen from the biogas. The presence of nitrogen in biogas can increase the energy consumption to reach the desired reaction temperature for biogas. Therefore, effective removal of nitrogen can be explored and developed in the future.

**THERMODYNAMIC LIMITATIONS OR CONSIDERATIONS**

Thermodynamic calculation can be conducted to evaluate biogas reforming performing Gibbs-free energy minimization and is common in different software packages such as Aspen Plus and HSC Chemistry. The temperature, pressure, and feed composition are the primary factors investigated.

**Dry Reforming of Biogas to Syngas**

The side reactions occurring during biogas reforming are an issue for the efficient utilization of biogas because they can decrease the reactant conversions. However, selection of the appropriate reaction conditions can lessen the side reactions and promote the conversions of reactants. Han et al. (Han et al., 2020) found that the CO₂ conversion and H₂/CO ratio both increased with the increase of the CH₄ to CO₂ ratio (0.5, 1.0, and 2.0) at 700–950°C. Cao et al. (Cao et al., 2018) studied the effect of the CH₄ to CO₂ molar ratio (3/6, 4/6, 6/6, 9/6, 12/6) and pressure (0.5, 1, 10, and 50 bar) on the biogas dry reforming performance at 800°C. The CH₄ conversion increased but the CO₂ conversion decreased with the decrease in CH₄ to CO₂ molar ratio. The CH₄ conversion and CO₂ conversion both increased with the decrease in pressure. Cui et al. (Cui et al., 2007) found that the dry reforming was accompanied by some side reactions such as CO disproportionation reaction, CO₂/H₂ methanation reaction, CO/H₂ methanation reaction, CO/H₂ reduction reaction, and rWGS reaction, as shown in Equations 7, 8, 9, and 10. When the reaction temperature was higher than 450°C, the CO/H₂ methanation reaction was thermodynamically forbidden. When the reaction temperature was higher than 550°C, the thermodynamic equilibrium conversion of CO₂ for the CO₂/H₂ methanation was lower than 3%. Figure 2A shows the variation of CH₄ conversion with the reaction temperature at the equilibrium state. The CH₄ conversion increased greatly from 58% to 98%, with temperature increasing from 550°C to 750°C (Cui et al., 2007). These results suggest that a high temperature (e.g., 750°C) lessens side reactions.

\[
\begin{align*}
2CO & \rightarrow C + CO₂, \quad \text{CO disproportionation} \quad \text{(Equation 7)} \\
CO + 3H₂ & \rightarrow CH₄ + H₂O, \quad \text{CO/H₂ methanation} \quad \text{(Equation 8)} \\
CO₂ + 4H₂ & \rightarrow CH₄ + 2H₂O, \quad \text{CO₂/H₂ methanation} \quad \text{(Equation 9)} \\
CO + H₂ & \rightarrow C + H₂O, \quad \text{CO/H₂ reduction} \quad \text{(Equation 10)}
\end{align*}
\]

A relatively low reaction temperature without the presence of H₂O and/or O₂ in feed gas generally tends to cause coke formation. The higher conversions at higher temperatures were due to the dry reforming being an endothermic reaction. Coke formation during biogas reforming is a major challenge for the efficient utilization of biogas. Nandini et al. (Nandini et al., 2005) reported that temperatures higher than 870°C are needed to prevent coke formation when CH₄/CO₂ feed ratio is 1 and pressure is 1 bar. Chen et al. (Chen et al., 2017) found with a temperature change from 500°C to 800°C, the generated solid carbon was negligible. The conversions of CH₄ and CO₂ increased with the temperature increase, but the rate of conversion increase was slower at higher.
temperatures (Figure 2B) (Chen et al., 2017). Ayodele and Cheng (Ayodele and Cheng, 2015) found that the yield of H\textsubscript{2} and CO increased with increased temperature from 200\textdegree{}C to 1,000\textdegree{}C. However, the syngas formation was not thermodynamically favored at 200–400\textdegree{}C (Ayodele and Cheng, 2015).

For dry reforming of biogas, some side reactions such as rWGS, CO/H\textsubscript{2} methanation reaction, and CO\textsubscript{2}/H\textsubscript{2} methanation reaction take place. A high temperature (e.g., 870\textdegree{}C) is needed to prevent the coke formation on catalysts. With suitable increased temperature range, the conversions of CH\textsubscript{4} and CO\textsubscript{2} increase, and the yield of H\textsubscript{2} and CO increase due to the endothermic nature of the steam reforming of methane. Almost all CH\textsubscript{4} was consumed above 850\textdegree{}C, except the lowest (CO\textsubscript{2} + H\textsubscript{2}O) to CH\textsubscript{4} ratio (0.9). This indicates that oxidizing agents were limiting reactants so that insufficient amounts of oxidizing agents caused a lower conversion.

**Bi-reforming of Biogas to Syngas**

The reaction conditions have a large effect on the biogas bi-reforming process. Jang et al. (Jang et al., 2016) investigated the effects of the (CO\textsubscript{2} + H\textsubscript{2}O) to CH\textsubscript{4} ratio (0.9–2.9) and temperature (500–1000\textdegree{}C) on the thermodynamic equilibrium. For all (CO\textsubscript{2} + H\textsubscript{2}O) to CH\textsubscript{4} ratios, the CH\textsubscript{4} conversion increased with the increase in temperature due to the endothermic nature of the steam reforming of methane. Almost all CH\textsubscript{4} was consumed above 850\textdegree{}C, except the lowest (CO\textsubscript{2} + H\textsubscript{2}O) to CH\textsubscript{4} ratio (0.9). This indicates that oxidizing agents were limiting reactants so that insufficient amounts of oxidizing agents caused a lower conversion.

Figure 2. The Equilibrium State of Biogas Reforming

(A) Variation of CH\textsubscript{4} conversion with the reaction temperature (the feed molar ratio of CH\textsubscript{4}:CO\textsubscript{2} = 1:1). Reproduced with permission from (Cui et al., 2007). Copyright © 2006 Elsevier B.V.

(B) Conversion efficiencies and selectivity: (I) CH\textsubscript{4} conversion, (II) CO\textsubscript{2} conversion, (III) H\textsubscript{2} to CO ratio, (IV) H\textsubscript{2}O productivity. Reproduced with permission from (Chen et al., 2017). Copyright © 2016 Elsevier Ltd.

(C) Equilibrium composition for biogas bi-reforming at different temperatures and 1 bar. The initial amounts of reactants in kmol are CH\textsubscript{4} = 3, CO\textsubscript{2} = 1, and H\textsubscript{2}O = 2. Allowed products are H\textsubscript{2}, CO, and C(s). The dotted curve shows C(s) formation under the same conditions, but in the absence of steam. Reproduced with permission from (Kumar et al., 2015). Copyright © 2015 Elsevier Ltd.
CH₄ conversion. The CO₂ conversion exhibited an increasing trend above 550°C, suggesting carbon dioxide reforming of methane was dominant at a high temperature. The coke yield decreased with the increase in temperature. The coke yield here is determined based on the mass of feed unless specified. In addition, the coke yield decreased largely with the increase in the (CO₂ + H₂O) to CH₄ ratio (Jang et al., 2016).

Elsayed et al. (Elsayed et al., 2018) found that the methane reforming was not possible thermodynamically below 350°C. The H₂ to CO ratio decreased as the reaction temperature increased from 450 to 600°C. The CO₂ conversion increased with an increase in the reaction temperature as the WGS reaction was favored at lower temperatures (<500°C) (Elsayed et al., 2018). Kumar et al. (Kumar et al., 2015) found that temperatures above approximately 850°C were required to reach basically complete equilibrium to H₂ and CO (Figure 2C). There was a significant thermodynamic driving force for carbon formation at temperatures below ~800°C. In addition, the presence of H₂O decreased carbon formation (Kumar et al., 2015).

Challenges related to coke prevention in biogas reforming include the use of low reaction temperatures due to energy saving and catalyst deactivation prevention, varying content of H₂O due to various biogas sources, and catalysts not being advanced. A high reaction temperature (≥500°C) and high (CO₂ + H₂O) to CH₄ ratio are needed for the biogas reforming to prevent coke formation. The addition of H₂O into biogas reduces coke formation due to their capability of initiating reactions that result in coke destruction.

**Tri-reforming of Biogas to Syngas**

To obtain desired reactant conversions and H₂ to CO ratio, additional components (H₂O and O₂) can be added into the biogas. Maintaining a high reforming performance with a low processing cost is a challenge. Diez-Ramirez et al. (Diez-Ramirez et al., 2016) studied the thermodynamic analysis of the biogas tri-reforming process at different temperatures (850–1000°C) and feed compositions. The optimal molar feed composition was identified as CH₄/CO₂/H₂O/O₂ = 4:1:4:2, where the high thermal efficiency (>70%), high methane conversion (>90%), and desirable H₂ to CO molar ratio (2:1) were achieved. The reaction temperature of 950°C led to higher methane conversion (~98%), a desirable H₂ to CO molar ratio, and superior exergy efficiency value. The exergy destruction mainly occurred in the reactor, mostly due to the high irreversibility of the chemical reactions. Future research should focus on decreasing the exergy destruction within the reactor and heat exchanger (Diez-Ramirez et al., 2016). A high reaction temperature (≥850°C) is helpful for the biogas reforming to improve the conversions of CH₄ and CO₂. However, a high reaction temperature needs a high energy consumption. To reduce the energy consumption, the biogas reforming at low temperatures (<600°C) has been investigated and developed (see Low Temperature Biogas Reforming).

Zhang et al. (Zhang et al., 2014) investigated the effects of process variables such as pressure (1–20 bar), inlet composition (CH₄, CO₂, H₂O, and O₂), and temperature (200–1000°C) on the product distribution of tri-reforming. As the temperature increased from 200 to 850°C, the H₂ molar fraction increased from 0 to ~0.6 (Figure 3). At temperatures <600°C, the CO₂ molar fraction was high, indicating the overall reaction favored the WGS reaction instead of dry reforming. The H₂ composition decreased, and the CH₄ composition increased with increased pressure from 1 to 20 bar. Relatively low pressure allowed the researchers to obtain products rich in H₂ and CO. A high temperature and low pressure favor the high CO₂ conversion and H₂ production rate during tri-reforming. With an increased O₂ to CH₄ ratio, the CH₄ conversion increased greatly, especially at temperatures below 850°C (Zhang et al., 2014). Methane is a limiting reactant during the tri-reforming process. With the increase of H₂O content, the CO₂ conversion decreases because H₂O is more chemically reactive to CH₄ than CO₂. Increased H₂O and O₂ in the feed can result in decreased CO₂ conversion and H₂ yield. A relatively low pressure is helpful to improve the conversion of CO₂. However, the syngas produced from biogas reforming needs to be synthesized to liquid hydrocarbon fuels through Fischer-Tropsch synthesis (FTS) at a high pressure (e.g., 20 bar). When combining the biogas reforming and FTS systems, a relatively high pressure in the biogas reforming section will be helpful for the compressor to easily reach the high pressure needed in the FTS section for industrial-scale application. The gas exiting the reformer needs to be cooled to knock out the water.

In our previous study (Zhao et al., 2018a), the thermodynamic equilibrium analysis of tri-reforming was investigated at conditions of pressure (3 bar), temperatures (830°C and 860°C), and H₂O to CH₄ molar ratios (0.34, 0.69, 1.40, and 2.10). The equilibrium O₂ conversions were nearly 100% at all conditions. The equilibrium CH₄ conversion increased from 95% to 99%, whereas the equilibrium CO₂ conversion decreased from 71% to 8% with the increase of the H₂O to CH₄ molar ratio. In addition, the produced H₂ to CO molar ratio increased as the H₂O/CH₄ molar ratio increased (Zhao et al., 2018a).
The appropriate amount of O₂ in biogas is helpful to improve the CH₄ conversion because the presence of O₂ can compensate for some energy consumption and prevent coke formation. In addition, O₂ is nearly all consumed. With the increase of the H₂O to CH₄ molar ratio in a range, the equilibrium CH₄ conversion increases. However, the presence of O₂ and H₂O can decrease the CO₂ conversion because they are more chemically reactive with CH₄ than CO₂. The produced H₂ to CO molar ratio increased as the H₂O to CH₄ molar ratio increased due to the potential 3:1 H₂ to CO ratio resulting from the steam reforming reaction. It is flexible for biogas reforming to add necessary components into biogas based on the biogas composition to obtain desired reactant conversions and H₂ to CO ratio. However, the addition of other components can increase the biogas reforming cost.

**REFORMING TECHNOLOGIES OF CONVERTING BIOGAS TO SYNGAS**

**Dry Reforming of Biogas to Syngas**

The dry reforming of biogas is a highly endothermic process typically taking place at temperatures of 700–900°C (Table 2). Generally, the pressure of 1 bar and a CH₄ to CO₂ ratio of 1.0–2.0 in a quartz reactor were utilized for the dry reforming of biogas. The typical CH₄ to CO₂ ratio (1.0–2.0) falls in the range of that in biogas according to biogas composition study described above. The catalysts used in biogas dry

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**Figure 3. The Equilibrium State of Biogas Tri-reforming**

Effect of the pressure on (A) H₂, (B) CH₄, (C) CO, (D) CO₂, and (E) H₂O mole fractions and (F) effect of the temperature on product mole fractions. Reproduced with permission from (Zhang et al., 2014). Copyright © 2014 American Chemical Society.
reforming were mainly Ni-based and Ir-based catalysts, which showed high activity. Ir is a noble metal, and its use in large amounts could increase costs. However, noble metals in small amounts can decorate the cheaper metals (e.g., Ni) on the catalysts. Nickel is one of the most popular metals used in dry reforming. Al2O3 and SiO2 were the common supports utilized for dry reforming of biogas. The CH4 and CO2 conversions obtained were generally higher than 42% and 57%, respectively. The produced H2 to CO ratio was 0.7–1.0, which is close to the equilibrium calculation. The coke rate varied depending on the reaction conditions (including time-on-stream, temperature, pressure, and GHSV) and catalyst performance.

| Catalyst                      | Reaction Conditions                                                                 | CH4 Conv. (%) | CO2 Conv. (%) | H2/CO | Coke Rate (g_{cokel}/(g_{cat}*h)) | Ref.                        |
|-------------------------------|-------------------------------------------------------------------------------------|---------------|---------------|-------|-------------------------------|-----------------------------|
| Ni/SrZrO3                     | 700–900°C, CH4/CO2/He = 1/0.5/18.5, 1 bar, quartz reactor                           | 43–55         | 86–99         | −1.0  | 0.018–0.042                   | (Evans et al., 2014)        |
| Ni/Al2O3                      | 750°C, CH4/CO2 = 1/1, 18,000 mL(STP)g⁻¹h⁻¹, Inconel tubular reactor                | 65–97         | −             | −     | [7.5 × 10⁻⁴, 8.1 × 10⁻⁴]     | (Lee et al., 2003a)         |
| 5Ni-10W/Al2O3                  | 750°C, CH4/CO2/Ar = 1/1/1, 36,000 mLg⁻¹h⁻¹, 1 bar, quartz tubular fixed-bed reactor | 60            | −75           | −0.8  | −                             | (Arbag et al., 2015)        |
| Ni@SiO2                       | 750°C, CH4/CO2/N2 = 1/1/2, 48,000 mL(STP)g⁻¹h⁻¹, 1 bar, fixed-bed reactor          | −71           | −58           | −0.7  | 4.9 × 10⁻⁴                    | (Zhang and Li, 2015)        |
| Ir/Al2O3                      | 750°C, CH4/CO2 = 1/1.8, 9,000 h⁻¹, 1 bar, fixed-bed quartz reactor                 | −90           | −68           | −1.0  | −                             | (Yentekakis et al., 2015)   |
| Ir/Zr0.92Y0.08O2–d             | 750°C, CH4/CO2 = 1/1.8, 11,000 h⁻¹, 1 bar, fixed-bed quartz reactor                | −87           | −65           | −1.0  | −                             | (Yentekakis et al., 2015)   |
| Ir/Ce0.9Gd0.1O2–d              | 750°C, CH4/CO2 = 1/1.8, 18,000 h⁻¹, 1 bar, fixed-bed quartz reactor                | −90           | −63           | −0.9  | −                             | (Yentekakis et al., 2015)   |
| Ni-Mg PSNTS (phyllosilicate nanotubes) | 750°C, CH4/CO2/He = 1/1/1, 60,000 mLg⁻¹h⁻¹, 1 bar, quartz tube reactor            | −85           | −89           | −0.7  | 0.022                         | (Bian et al., 2016)         |
| Ni-Mg PSNTS@silica            | 750°C, CH4/CO2/He = 1/1/1, 60,000 mLg⁻¹h⁻¹, 1 bar, quartz tube reactor            | 85            | 89            | 0.8   | Negligible                    | (Bian et al., 2016)         |
| Ni-Y/KIT-6                    | 750°C, CH4/CO2/Ar = 1/1/8, 20,000 h⁻¹, 1 bar, fixed-bed quartz reactor            | −65           | −72           | −0.8  | −                             | (Swirk et al., 2019)        |
| Ru/SiO2                       | 700°C, CH4/CO2/Ar = 1/1/8, 10,000 mLg⁻¹h⁻¹, 1 bar, fixed-bed quartz reactor        | −93           | −93           | −1.0  | 3.8 × 10⁻⁴                    | (Das et al., 2019)          |
| Sr0.92Y0.08Ti1–xRuO3–d         | 800°C, CH4/CO2/N2 = 1/1/2, 12,000 h⁻¹, 1 bar, fixed-bed quartz reactor            | −82           | −90           | −0.9  | −                             | (Kim et al., 2019a)         |
| La(Co0.1Ni0.9)0.5Fe0.5O3       | 750°C, CH4/CO2 = 1/1, 12,000 mLg⁻¹h⁻¹, 1 bar, fixed-bed quartz reactor            | 70            | 80            | −0.9  | Negligible                    | (Wang et al., 2019)         |
| Rh/Al2O3                      | 750°C, CH4/CO2 = 1/1, 120,000 mLg⁻¹h⁻¹, 1 bar, fixed-bed quartz reactor            | −80           | −88           | −1.0  | 8.8 × 10⁻⁴                    | (Yentekakis et al., 2019)   |

Table 2. Recent Studies on Dry Reforming of Biogas to Syngas (the Unit g_{cat} Refers to Grams of Catalyst)
The carbon deposition on the catalyst, low reactant conversion, and high reaction temperature are the main challenges for the dry reforming of biogas.

The high endothermic characteristics of the biogas dry reforming reaction cause a high reaction temperature required, which can cause the deactivation of catalysts. The catalyst deactivation is a drawback of biogas dry reforming applied in the industry. To prevent the catalyst deactivation and maintain the catalyst stability, researchers have investigated and developed different catalysts. Evans et al. (Evans et al., 2014) studied the dry reforming of biogas over Ni/SrZrO3 catalysts, which exhibited a high selectivity toward the reforming reaction (i.e., syngas formation). The CH4 conversion was limited due to CO2 as a limiting reactant (Evans et al., 2014). Zhang and Li (Zhang and Li, 2015) found that the core-shell Ni@SiO2 catalysts exhibited a high stability and activity. Both micropores and mesopores were created in the amorphous SiO2 shell, which suppressed carbon filament growth. The Ni cores (nanoparticles) were accessible to gas molecules through the pores of the SiO2 shell (Zhang and Li, 2015). The catalyst species, catalyst structure, and interaction between metal and support play a significant role in the resistance of carbon deposition on the catalyst. Other factors such as metal particle size, metal particle distribution, metal content, metal oxide interface, promoter, and support can affect the performance of catalysts.

Bian et al. (Bian et al., 2016) studied the dry reforming of biogas over multicore-shell Ni-Mg PSNTS@silica catalysts, which exhibited a stable and high conversion during a run of 72 h. Without the silica coating, the Ni-Mg PSNTS exhibited a high carbon deposition due to the decomposition of nanotubular structure at a high temperature (e.g., 750 °C). After coating, the thermal stability of Ni-Mg PSNTS@silica was significantly improved, as silica was expected to form a strong interaction with the outer and inner surfaces of the PSNTS. The high activity and carbon resistance of the Ni-Mg PSNTS@silica catalyst was ascribed to the confinement effect and interaction between metal and nanotubular support given by the unique multicore-shell structure. The Ni-Mg PSNTS@silica catalyst is believed to be used for other high temperature and carbon-deposition reactions (Bian et al., 2016). Cruz-Flores et al. (de la Cruz-Flores et al., 2020) found that the Ni-SiO2 catalysts exhibited a low carbon deposition at a short time-on-stream (e.g., ~4 h), but a severe sintering of Ni particles at a long time-on-stream (e.g., 50 h). Understanding the reaction mechanism on the catalyst will be helpful to design the catalysts, perform the biogas reforming reactions, and analyze the reaction results. Das et al. (Das et al., 2019) studied the dry reforming of biogas over the Ru-modified catalyst (Ru-Mg-Ce/SiO2). The proposed reaction mechanism (Figure 4A) included adsorption of CO2 preferentially occurred over MgO surface, forming a layer of surface-adsorbed CO2. The adsorbed CO2 transformed over the Ce2O3. A fraction of the CO2 activated over the catalyst surface to oxidize activated metallic Ru. The oxidized Ru metal was reduced to metallic Ru by the generated H2. The selective dissociation of CH4 occurred over the Ru metal surface, which yielded a layer of carbon over the Ru metal surface. The deposited carbon was oxidized by the catalyst surface oxygen to generate CO (Das et al., 2019). Khoja et al. (Khoja et al., 2018) proposed a reaction mechanism of dry reforming of biogas over Ni/Al2O3-MgO catalysts (Figure 4B). The reaction mechanism included activation of CH4 on Ni and gasification of deposited carbon on MgO. The adsorption of elemental and intermediates containing H, O, C, and oxy-carbonates occurred on the surface and active sites of the catalyst support. The dissociation of CH4 and CO2 was initiated by plasma (Khoja et al., 2018).

Bobadilla et al. (Bobadilla et al., 2017) proposed a reaction mechanism of biogas dry reforming over Rh/MgAl2O4 catalysts (Figure 4C). The dissociative adsorption of CO2 initially occurred, and the active oxygen species formed on the Rh metallic sites. The oxygen species promoted the CH4 activation to produce H2. The produced H2 initiated a bi-functional mechanism, where CO2 was activated on the basic sites of the support surface. CO2 can be transformed into CO through the H-assisted dissociation on the metal support interface and the direct dissociation on Rh metallic sites (Bobadilla et al., 2017). The reaction mechanism of biogas reforming has been proposed to some extent. It is proposed that the dissociation of CH4 typically occurs on the active metal surface, and CO2 prefers to be activated on the support surface. Carbon is typically deposited on the active metal surface, and the deposited carbon is oxidized by the catalyst surface oxygen. However, it will be important for researchers to study more on the reaction mechanism over various catalysts, which can promote biogas application.

Hossain et al. (Hossain et al., 2019) found that GHSV (10,000–60,000 h⁻¹) had an influence on the CH4 and CO2 conversions, where the highest CH4 and CO2 conversions were obtained at 35,000 h⁻¹. At a higher temperature during biogas dry reforming, the pyrolysis of methane can become a primary carbon-forming reaction because the Boudouard reaction (2CO → CO2 + C) becomes more thermodynamically
unfavorable (Evans et al., 2014). With the increase of the reaction temperature from 700 to 900°C, the CH₄ conversion increased from 43% to 55% and the CO₂ conversion increased from 86% to 99%, respectively (Evans et al., 2014). In summary, dry reforming of biogas has been frequently studied by academia. A relatively lower GHSV indicates a higher residence time of reactants on the catalyst surface, probably leading to higher reactant conversions. The CH₄ conversion increases with increased reaction temperature. The noble metal–based catalysts (e.g., Ru/SiO₂ and Ir/Al₂O₃) seem to exhibit a higher dry reforming performance than Ni-based catalysts (e.g., Ni-W/Al₂O₃ and Ni@SiO₂) due to the high stability of noble metals. Significant advances on the catalyst development have been made to reduce the carbon deposition. Incorporation of appropriate metals (e.g., La, Fe, Sn, and Zr) (Chen et al., 2020; Li et al., 2020; Zhang et al., 2020a, 2020b) to Ni-based catalysts has proven to improve the coke-resistance property. Some catalysts (e.g., Ni-Mg PSNTS) showed a high coke rate up to 0.022 gcoonk/gcat*h⁻¹. However, the Ni-Mg PSNTS@silica catalysts showed negligible coke rate due to the oxygen vacancy of MgO and the function of the silica support. The addition of H₂O and O₂ into the biogas can help prevent the coke formation, discussed in below bi-reforming and tri-reforming sections. The produced H₂/CO ratio in biogas dry reforming is not suitable to produce many hydrocarbon products. Developments on the reaction condition optimization and catalysis mechanism have been made in recent years, which promote the biogas dry reforming performance. The choice of appropriate metal and support for catalyst synthesis (e.g., Ni-Zr on the CeO₂ support), reaction temperature (≥800°C), pressure (e.g., 1 bar), and composition of reactants, is significant in the stable catalytic performance and low deactivation for the dry reforming of biogas.

**Bi-reforming of Biogas to Syngas**

The bi-reforming of biogas typically takes place at temperatures of 800–850°C and pressure of 1–7 bar (Table 3). The CH₄ to CO₂ ratio and CH₄ to H₂O ratio are typically in the range of 1–3 and 0.5–2.5, respectively.

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**Figure 4. Proposed Mechanism of Biogas Dry Reforming over Different Catalysts**

(A) Ru-Mg-Ce/SiO₂ catalysts. Reproduced with permission from (Das et al., 2019). Copyright © 2018 Elsevier Ltd.

(B) Ni/Al₂O₃-MgO catalysts in a cold plasma dielectric barrier discharge reactor. Reproduced with permission from (Khoja et al., 2018). Copyright © 2018 Elsevier B.V.

(C) Rh/MgAl₂O₃ catalysts. Reproduced with permission from (Bobadilla et al., 2017). Copyright © 2017 Wiley-VCH Verlag GmbH & Co. KGaA, Weinheim.
Ni-based catalysts, with different supports such as Al$_2$O$_3$, SiC, and ZrO$_2$, are mainly utilized. The produced H$_2$ to CO ratio is generally 1.1–2.7, which is higher than that obtained from biogas dry reforming. However, the main challenges for bi-reforming of biogas include low reactant conversions and catalyst deactivation. The CH$_4$ and CO$_2$ conversions are generally higher than 61% and 33%, respectively. The coke rates of catalysts varied from being negligible to $\sim 0.006 \text{ g}_{\text{coke}}/(\text{g}_{\text{cat}} \times \text{h})$.

| Catalyst | Reaction Conditions | CH$_4$ Conv. (%) | CO$_2$ Conv. (%) | H$_2$/CO | Coke Rate | Ref. |
|----------|---------------------|------------------|------------------|----------|-----------|------|
| NiO/MgO  | 830°C, CH$_4$/CO$_2$/H$_2$O = 3/1.2/2.4, 60,000 mL*g$^{-1}$*h$^{-1}$, 7 bar, tubular flow reactor | 71                | −73              | 2.0      | −         | (Olah et al., 2013) |
| LaSrNiAlSiC | 850°C, CH$_4$/CO$_2$/H$_2$O = 1/0.5/1.4, 18,000 mL*g$^{-1}$*h$^{-1}$, 1 bar, fixed-bed Incoloy reactor | 95                | 34               | 2.1      | Negligible | (Kim et al., 2015) |
| Ni/Al$_2$O$_3$ | 850°C, CH$_4$/CO$_2$/H$_2$O = 1/1/2, fixed-bed reactor | −99               | −47              | −1.5     | −         | (Park et al., 2015) |
| Mo$_2$C-Ni/ZrO$_2$ | 850°C, CH$_4$/CO$_2$/H$_2$O/N$_2$ = 1/0.4/0.8/1.6, 60,000 mL*g$^{-1}$*h$^{-1}$, 1 bar, quartz tube fixed-bed reactor | −98               | −79              | −1.9     | $\sim 6 \times 10^{-3}$ | (Li et al., 2015b) |
| NiO/MgO  | 830°C, CH$_4$/CO$_2$/H$_2$O = 3/1.2, 60,000 mL*g$^{-1}$*h$^{-1}$, 7 bar, tubular flow reactor | −70               | −72              | 2.0      | $\sim 7.3 \times 10^{-5}$ | (Olah et al., 2015) |
| NiO/MgO  | 830°C, CH$_4$/CO$_2$/H$_2$O = 3/1/2.25, 60,000 mL*g$^{-1}$*h$^{-1}$, 1 bar, tubular flow reactor | 86                | 94               | −1.9     | $\sim 7.3 \times 10^{-5}$ | (Olah et al., 2015) |
| La$_{0.9}$Ce$_{0.1}$NiO$_3$ | 800°C, CH$_4$/CO$_2$/H$_2$O = 1/1/1, 3,000 h$^{-1}$, 1 bar, fixed-bed down flow reactor | 100               | −61              | −        | Negligible | (Yang et al., 2015) |
| Ni/ZrO$_2$ | 850°C, CH$_4$/CO$_2$/H$_2$O/N$_2$ = 3/0.4/0.8/1.9, 60,000 mL*g$^{-1}$*h$^{-1}$, 1 bar, quartz tube fixed-bed reactor | −90               | −88              | 1.1      | $5 \times 10^{-5}$ | (Li et al., 2015a) |
| LA-Ni/ZrO$_2$ (ligand-assisted) | 850°C, CH$_4$/CO$_2$/H$_2$O/N$_2$ = 3/0.8/0.4/0.2, 48,000 mL*g$^{-1}$*h$^{-1}$, 1 bar, quartz tube fixed-bed reactor | −94               | −92              | 1.1      | $1.7 \times 10^{-4}$ | (Li et al., 2015a) |
| Ni/SBA-15 | 800°C, 36,000 mL*g$^{-1}$*h$^{-1}$, 1 bar, quartz tube fixed-bed reactor | −62               | −59              | 2.1      | −         | (Singh et al., 2018) |
| Ni/La-Si | 800°C, CH$_4$/CO$_2$/H$_2$O = 1/0.4/0.8, 1.584 × 10$^5$ mL*g$^{-1}$*h$^{-1}$, 1 bar, fixed-bed quartz reactor | −90               | −75              | −2.0     | $4.7 \times 10^{-4}$ | (Chen et al., 2019) |
| B-Ni/SBA-15 | 800°C, CH$_4$/CO$_2$/H$_2$O = 1/0.33/0.67, 36,000 mL*g$^{-1}$*h$^{-1}$, 1 bar, packed-bed quartz reactor | −67               | −60              | −2.7     | −         | (Siang et al., 2019) |
| Ni/MgAl$_2$O$_4$ | 700°C, CH$_4$/CO$_2$/H$_2$O = 1/0.52/3.71, fixed-bed reactor | −98               | −60              | −2.6     | −         | (Rahmat et al., 2019) |
| Mo$_2$C-Ni/ZrO$_2$ | 700°C, CH$_4$/CO$_2$/H$_2$O = 1/0.4/0.8, 1 bar, quartz tube fixed-bed reactor, 36,000 mL/(g*h) | −74               | −54              | −        | Negligible | (Ren and Zhao, 2019) |
| Ni/Mg-Al mixed oxide | 775°C, CH$_4$/CO$_2$/H$_2$O = 1/0.4/0.73, 1 bar, fixed-bed reactor, 86,000 h$^{-1}$ | 73                | 64               | 2.0      | −         | (Li and van Veen, 2018) |

Table 3. Recent Studies on Bi-reforming of Biogas to Syngas

Ni-based catalysts, with different supports such as Al$_2$O$_3$, SiC, and ZrO$_2$, are mainly utilized. The produced H$_2$ to CO ratio is generally 1.1–2.7, which is higher than that obtained from biogas dry reforming. However, the main challenges for bi-reforming of biogas include low reactant conversions and catalyst deactivation. The CH$_4$ and CO$_2$ conversions are generally higher than 61% and 33%, respectively. The coke rates of catalysts varied from being negligible to $\sim 0.006 \text{ g}_{\text{coke}}/(\text{g}_{\text{cat}} \times \text{h})$. 
Researchers have developed different catalysts to obtain desired reactant conversions and H2 to CO ratios. Kim et al. (Kim et al., 2015) reported that the conversions of CH4 and CO2 both increased by 14% and 129%, respectively, with the increase of Al2O3 content from 0 to 10 wt% in the LaSrNi/Al/SiC catalysts, and then decreased by 13% and 42%, respectively, with the further increase of Al2O3 content to 20 wt%. The higher catalytic activity of LaSrNi/Al(10)/SiC (with 10 wt% of Al2O3 content) catalysts might be mainly ascribed to the suppressed aggregation of nickel crystallites, which caused a stronger interaction of Al2O3 modified SiC with La2NiO4 crystallites. The LaSrNi/Al(10)/SiC catalysts exhibited a larger surface area and higher dispersion of Al2O3 particles on SiC, compared with other Al2O3 concentrations. The nickel-containing crystallites seemed to be well dispersed on the SiC surface, forming intimately and strongly interacted La3NiO4-Al2O3 particles (Kim et al., 2015). Ren and Zhao (Ren and Zhao, 2019) found the catalytic activity and stability of Mo2C-Ni/ZrO2 catalysts were improved via a new synthesis method (glucose-assisted incipient wetness impregnation), compared with conventional incipient wetness impregnation method. Glucose was used as a metal dispersion promoting agent to improve the Ni dispersion in the catalyst.

Li et al. (Li et al., 2015b) studied the bi-reforming of biogas over Mo2C modified Ni/ZrO2 catalysts (0.2–3.0 wt% of Mo2C loading), which exhibited a superior catalytic activity and stability, compared with the unmodified one. This is attributed to the catalysis of Mo2C, improved Ni dispersion with Mo2C addition, and the different coke morphology caused by the change in Ni-ZrO2 interactions. A proposed reaction mechanism of biogas bi-reforming over Mo2C-Ni/ZrO2 and Ni/ZrO2 catalysts is shown in Figure 5A. The Ni active sites
were accessible to the reactants for the whisker-like coke deposited on the Mo$_2$C-Ni/ZrO$_2$ catalysts. However, the Ni active sites were not accessible to the reactants for the shell-like coke deposited on the Ni/ZrO$_2$ catalysts. An appropriate Mo$_2$C loading (e.g., 0.5 wt%) was required to improve the catalytic activity and stability (Li et al., 2015b). The catalyst support (e.g., Al$_2$O$_3$) and promoter (e.g., Mo$_2$C) affect the metal dispersion on the catalyst. If the catalyst support and promoter content is too high, aggregation of metal particles can occur, which can decrease the catalytic performance. Optimization of the catalyst support and promoter, with a stronger interaction between metal and support, a higher dispersion of metal particles on the support, and a larger surface area, is needed for the efficient utilization of biogas.

Optimization of the reaction conditions to obtain desired reactant conversions and H$_2$ to CO ratio is a main challenge for the biogas reforming. Olah et al. (Olah et al., 2015) found the conversions of both CH$_4$ and CO$_2$ increased when the temperature increased from 830 to 910°C. However, the H$_2$ to CO ratio decreased slightly with increased temperature. The conversion of CH$_4$ decreased and H$_2$ to CO ratio increased slightly with the pressure increase from 7 to 42 bar (Olah et al., 2015). Park et al. (Park et al., 2015) also found the conversions of CH$_4$ and CO$_2$ increased with the temperature increase from 750 to 900°C, whereas the H$_2$ to CO ratio decreased with increased temperature. The H$_2$ was consumed by carbon formation on the catalyst as the temperature increased, leading to the decreased H$_2$ to CO ratio (Park et al., 2015).

Li et al. (Li et al., 2015a) studied the bi-reforming of biogas over LA-Ni/ZrO$_2$ catalysts, which exhibited a superior catalytic activity (e.g., conversions of CH$_4$ and CO$_2$), compared with classical Ni/ZrO$_2$ catalysts. This can be attributed to the intensified Ni-support interaction, higher Ni dispersion, enlarged oxygen vacancy, enhanced reducibility of NiO led by oxygen vacancy, and increased t-ZrO$_2$ content of LA-Ni/ZrO$_2$ catalysts. The conversions of CH$_4$ and CO$_2$ decreased with the increase of GHSV from 24,000 to 72,000 mL·g$^{-1}$·h$^{-1}$, which might be ascribed to the decrease in the residence time on the catalyst surface and limited active sites for the increasing reactant amounts (Li et al., 2015a). Rahmat et al. (Rahmat et al., 2019) found that CH$_4$ and CO$_2$ conversions were achieved only at a suitable GHSV value (not too high or too low).

In the range of 830–900°C, a higher reaction temperature is generally favored to obtain a higher CH$_4$ conversion and CO$_2$ conversion during the biogas bi-reforming process. The high reaction temperature indicates a high processing cost due to the energy consumption. In addition, a high reaction temperature can lead to a low H$_2$ to CO ratio. High pressure can result in low CH$_4$ and CO$_2$ conversions. The optimization of the pressure can depend on the final application of the produced syngas due to specific requirements of the H$_2$ to CO ratio. A low GHSV can aid in equilibrium to be reached for obtaining high CH$_4$ and CO$_2$ conversions. However, for large-scale biogas reforming, a high GHSV may improve the feed speed. Looking for a balance between the high reactant conversions and suitable H$_2$ to CO ratio can be a strategy for biogas reforming.

The H$_2$O was added into the reactor system mainly through a syringe pump. The H$_2$O conversion was not reported in most biogas bi-reforming research, perhaps because the unconsumed H$_2$O is not easy to accurately measure. The low CO$_2$ conversion is a main challenge in biogas bi-reforming due to the competition between CO$_2$ and H$_2$O to react with CH$_4$. In addition, CO$_2$ is thermodynamically stable even at high temperatures. The catalyst development and reaction condition optimization have been made in recent years to look for a balance between the high CO$_2$ conversion and suitable H$_2$ to CO ratio. The produced H$_2$ to CO ratio is in the range of 1–3 because of the reaction of methane with CO$_2$ and H$_2$O. The catalysts exhibited different coke rates because of their characteristics and reaction conditions. Recent developments on the catalyst support have prevented the coke formation on the catalysts. The supports included MgO, ZrO$_2$, and ceria due to their high oxygen storage capability. The high oxygen storage property of ZrO$_2$ can reduce the carbon deposition on catalysts. Some catalysts, such as LaSrNi/Al/SiC and La$_{0.7}$Ce$_{0.3}$NiO$_3$, showed a negligible coke rate, probably due to the stronger metal-support interaction and higher metal particle dispersion. Utilizing noble metals (e.g., Rh) could increase the dispersion of Ni on the catalysts, thus improving catalytic performance.

**Tri-reforming of Biogas to Syngas**

Song and Pan (Song and Pan, 2004) first introduced tri-reforming of methane. Different catalysts with various supports and different biogas compositions have been investigated and developed to improve the catalytic performance for the biogas tri-reforming (Tables 4 and S1). Compared with dry reforming of biogas, both bi-reforming and tri-reforming can produce syngas (with a H$_2$ to CO molar ratio of 2:1).
that can be converted into high-value products (e.g., liquid fuel) without using a WGS reactor (Zhao et al., 2018b). During the process of biogas tri-reforming, additional H2O and O2 are provided to obtain a H2 to CO ratio (e.g., 1.7–2.2) suitable for FTS and to improve the CH4 conversion. The steam reforming and partial oxidation reforming of methane can potentially provide a H2 to CO ratio of 3:1 and 2:1, respectively. Compared with bi-reforming of biogas, the addition of O2 in tri-reforming can help reduce the carbon deposition on the catalyst and lower the endothermic nature. Partial oxidation of methane has been commercialized in recent years. Considering the energy balance of biogas reforming processes, the incorporation of H2O and O2 into biogas increases energy efficiency. Compared with dry reforming of biogas and steam reforming of methane, the tri-reforming uses 45.8% and 19.7% less energy, respectively (Zhao et al., 2019a). The tri-reforming of biogas typically takes place at temperatures of 700–950°C and pressure of 1 bar in a fixed-bed reactor. The CH4 to CO2 ratio, CH4 to H2O ratio, and CH4 to O2 ratio are typically in the range of 1–5, 0.3–5, and 2–15, respectively. Similar to biogas bi-reforming, the catalysts used in biogas tri-reforming are mainly Ni-based catalysts with different supports such as Al2O3, SiO2, ZrO2, and CeO2. The CH4 conversion is mainly higher than 72%, and the CO2 conversion is in the range of 22%–100%, respectively. The produced H2 to CO ratio is typically 1.0–2.3. Catalysts exhibit different coke rates, which could reach 0.028 g coke/(g cat*h).

It is important that catalysts exhibit high stability during the biogas reforming process, which is required for the commercial biogas application. An appropriate catalyst support is helpful for the superior metal dispersion and metal-support interaction. Singha et al. (Singha et al., 2016b) studied the tri-reforming of biogas over Ni/ZrO2 catalysts, which exhibited no deactivation for more than 100 h with almost complete stability.

### Table 4. Recent Studies on Tri-reforming of Biogas to Syngas

See also Table S1.

| Catalyst | Reaction Conditions | CH4 Conv. (%) | CO2 Conv. (%) | H2/CO Coke Rate (g coke/(g cat*h)) | Ref. |
|----------|---------------------|--------------|--------------|-----------------------------------|------|
| Ni/CeO2  | 800°C, CH4/CO2/O2/H2O = 1/0.67/0.1/0.3, 30,000 mL/g*h, 1 bar, continuous flow reactor | 97.4–99.6 | 87.8–90.5 | 1.3–1.4 | (Vita et al., 2014) |
| Ni/ZrO2  | 800°C, CH4/CO2/O2/H2O = 5/1/1/2.1, 80,000 mL/g*h, 1 bar, fixed-bed reactor | 84.4–98.5 | 89.3–98.5 | 1.6–2.2 | (Singha et al., 2016b) |
| Nickel-alumina aerogel | 700°C, 269,000 mL/g*h, 1 bar, fixed-bed reactor | 83.3 | – | 2.0–2.1 | 3.5 × 10⁻³ | (Yoo et al., 2015) |
| Ni-MgO/CeO2-ZrO2 | 800°C, 20,000 mL/g*h, 1 bar, CH4/CO2/O2/H2O = 5/1/1/2.1, fixed-bed reactor | 80.9–97.2 | 4.4–94.8 | – | (Singha et al., 2016a) |
| Ni/MgO/Al2O3 | 850°C, 32,000 mL/h*(g cat)⁻¹, 1 bar, CH4/CO2/O2/H2O = 1/0.21/0.1/0.81, fixed-bed reactor | –94 | –55 | –2.1 | – | (Song and Pan, 2004) |
| Ni/Mo/Ce | 850°C, CH4/CO2/O2/H2O = 1/0.39/0.16/0.30, fixed-bed reactor | –93 | ~100 | – | – | (Zou et al., 2016) |
| Ni/Ce-Zr-Al2O3 | 800°C, 1 bar, 161 g gas*(g cat*h)⁻¹, CH4/CO2/O2/liquid H2O = 1/0.67/0.25/0.0008, fixed-bed reactor | –99 | –42 | –1.9 | – | (Izquierdo et al., 2018) |
| Ni/CeO2-ZrO2 | 800°C, 1 bar, 17,220 mL*(g cat*h)⁻¹, CH4/CO2/O2/H2O = 1/0.23/0.07/0.46, fixed-bed reactor | – | – | 2.1 | Negligible | (Kumar et al., 2019) |
| Ni/TiO2 (calcined at 850°C) | 800°C, CH4/CO2/O2/H2O = 1/0.23/0.07/0.46, 1 bar, tubular reactor, 17,220 mL/g*h | – | – | 2.0 | Negligible | (Kumar et al., 2020) |
| Ni-MgO/Al2O3 | 827°C, CH4/CO2/O2/H2O = 1/1.33/0.47/2.47, 20 bar, multi-tubular reactor | –98 | –12 | –2.0 | – | (Alipour-Dehkordi and Khademi, 2019) |
| NiCe@SiO2 | 750°C, CH4/CO2/O2/H2O = 1/0.5/0.1/0.5, 1 bar, fixed-bed reactor, 60,000 mL/g*h | 79 | 75 | 1.7 | – | (Kim et al., 2019b) |

It is important that catalysts exhibit high stability during the biogas reforming process, which is required for the commercial biogas application. An appropriate catalyst support is helpful for the superior metal dispersion and metal-support interaction. Singha et al. (Singha et al., 2016b) studied the tri-reforming of biogas over Ni/ZrO2 catalysts, which exhibited no deactivation for more than 100 h with almost complete stability.
conversions of CH₄, CO₂, and H₂O. The high nickel dispersion and metal–support interaction of the catalyst increased its surface oxygen species and improved its reactivity. The oxygen atoms at the interface between ZrO₂ and Ni decreased the bond energy of the oxygen species, resulting in the oxygen becoming more easily reducible. These characteristics enhanced the catalytic activity of the Ni/ZrO₂ catalyst (Singha et al., 2016b). Similarly, Vita et al. (Vita et al., 2014) found that Ni/CeO₂ catalysts exhibited constant activity over 150 h of reaction (Vita et al., 2014). The catalysts typically need to be stable for at least 1 year of industry use. The high stability of catalysts can significantly reduce costs related to expensive catalyst precursors and synthesis process. Although some methods have been developed for the recycling of deactivated catalysts, the recycling process is high cost and the recycling times are limited. Therefore, it is important to develop catalysts that exhibit high activity and stability for biogas reforming.

Similar to above dry reforming and bi-reforming of biogas, the tri-reforming performance is affected by the reaction conditions. Damanabi et al. (Damanabi et al., 2019) found that the CH₄ and CO₂ conversions both decreased with the increase of the pressure (1–10 bar). Vita et al. (Vita et al., 2014) found that the conversions of CH₄ and CO₂ increased by 3% and 4%, respectively, but the H₂ to CO molar ratio decreased by 2% as the reaction temperature increased from 800 to 900 °C. A high temperature would favor the dry reforming of methane and steam reforming of methane due to their strong endothermic nature, resulting in higher conversions of CH₄ and CO₂. The conversion of CH₄ increased whereas the conversion of CO₂ decreased with the increased O₂ to CH₄ molar ratio (from 0.05 to 0.10). A high O₂ concentration can promote the conversion of CH₄. The conversion of CH₄ increased gradually whereas the conversion of CO₂ decreased largely with the increased H₂O to CH₄ molar ratio (from 0.3 to 0.7). The H₂ to CO molar ratio increased with the increased H₂O to CH₄ molar ratio (from 0.3 to 0.7). A high H₂O concentration can favor the steam reforming of methane with a lower contribution of the dry reforming of methane. In addition, the high H₂O concentration can facilitate the WGS reaction (Vita et al., 2014). Sadeghi et al. (Sadeghi et al., 2018) found the optimized conditions for maximizing the overall energy efficiency were 1,127°C, CH₄ to CO₂ to air to H₂O molar ratio = 1/0.12/3.80/0.6, and 10.8 bar. Under optimized process conditions, the overall energy efficiency reached 92%. The CH₄ conversion, CO₂ conversion, and produced H₂ to CO ratio were 99.99%, 8.2%, and 2.0, respectively. A higher steam concentration in the reactants can increase the reactor exergy efficiency due to the higher H₂ concentration in the product. A lower CO₂ concentration in the reactants is helpful to improve the reactor energy efficiency and reduce the product cost (Sadeghi et al., 2018). The modification of the reforming reactor system (Fekri Lari et al., 2019) was also found to be able to improve the CO₂ conversion.

The CH₄ conversion and the H₂ to CO ratio can be enhanced by increasing the amount of air or H₂O in the reactants. However, the dry reforming reaction can be suppressed, causing a low or negative CO₂ conversion. The negative CO₂ conversion can result from the WGS reaction (Chein and Hsu, 2018). Maciel et al. (Maciel et al., 2010) proposed a reaction mechanism of the biogas tri-reforming process. Dry and steam reforming reactions took place to produce H₂ and CO. The CH₄ and O₂ could react to yield H₂ and CO. However, the combustion-reforming could happen, where CH₄ and O₂ reacted to yield H₂O and CO₂. The methane cracking took place to yield C and H₂. The Boudouard reaction took place to yield C and CO₂ (Maciel et al., 2010).

A high reaction temperature is favored to obtain a high CH₄ conversion and CO₂ conversion due to the endothermic nature but can reduce the H₂ to CO ratio. A high O₂ concentration (in biogas) within a suitable range can promote the conversion of CH₄. The presence of O₂ is useful for industrial application of biogas reforming because it could cause some exothermic side reactions, which can compensate for the energy consumption during the biogas tri-reforming process. However, how to avoid hot spots in the reactor while adding O₂ is crucial. A high H₂O concentration within a suitable range can increase the conversion of CH₄ and the H₂ to CO ratio. However, the CO₂ conversion will be reduced at a high H₂O concentration.

The development on catalysts and reaction condition optimization (e.g., the addition of more O₂ into the reactor) has been successfully made in recent years to prevent the coke formation. Among many nickel-based catalysts used in biogas reforming, the Ni nanoparticles on the CeO₂–ZrO₂ support seem most promising. The nanoparticle size is helpful to enhance the Ni dispersion. The CeO₂–ZrO₂ support exhibits strong interaction with Ni metal and high oxygen storage capability, which can prevent metal sintering and reduce carbon deposition. In addition, the incorporation of Zr in the support can help disperse Ni particles. For the Ni/Mg₀.₅T₁₀.₂5O, Ni/Mg₀.₇₅T₁₀.₈₅O, and Ni/Mg/Ce₀.₆Zr₀.₄O₂ catalysts, the study of effects of atom
ratio of Mg to Ti and Ce to Zr is lacking. It would be interesting to investigate how the different atom ratios of metals affect the performance of catalysts. A main challenge for biogas tri-reforming is the low CO₂ conversion due to the competition between H₂O and CO₂ to react with methane. Some researchers studied the tri-reforming of biogas at a high pressure such as 10 bar, but the low CO₂ conversion at high pressures can be one issue for following FTS reaction to produce liquid fuel. As bi-reforming of biogas, the reports on the H₂O conversion are lacking due to the component tracking challenge. In addition, how to feed H₂O constantly and accurately during the tri-reforming process can help. The successful incorporation of H₂O and O₂ into biogas for tri-reforming with a high conversion efficiency is a major breakthrough achieved in recent years. The high reaction temperature (≥700°C) used for tri-reforming of biogas indicates that new advanced technologies (e.g., plasma and solar) might be used to lower the reaction temperature to save energy and prevent catalyst deactivation.

Compared with biogas dry reforming, the presence of H₂O in bi-reforming can inhibit the coke formation on catalysts. There is no need to remove the moisture from raw biogas in the biogas bi-reforming. The presence of H₂O can also reduce the energy consumption during the biogas bi-reforming process, as the use of a high reaction temperature in biogas reforming results in a high processing cost. The catalysts utilized in biogas bi-reforming have similar phenomena to biogas dry reforming: Ni as the main active metal and Al₂O₃ as a common support. A higher H₂ to CO ratio (1.1–2.7) is generated in biogas bi-reforming than that in biogas dry reforming (0.7–1.0) due to the methane steam reforming reaction in bi-reforming. Steam reforming of methane has been a commercial technology. However, the methane dry reforming reaction can be suppressed to decrease the CO₂ conversion in biogas bi-reforming. The H₂ to CO ratio generated from biogas dry reforming is suitable for syngas to be used to generate heat. However, the biogas can be directly used to generate heat instead of producing syngas via dry reforming. The H₂ to CO ratio generated from biogas bi-reforming is suitable for syngas to be used to generate heat, methanol, liquid fuel, or hydrogen. When the produced H₂ to CO molar ratio is in the range of <1.5, 1.5–2.0, 1.7–2.2, and >2.2, the syngas is suitable to generate heat, methanol, liquid fuel, and hydrogen, respectively. The control of H₂O content in biogas bi-reforming can adjust the H₂ to CO ratio for a specific target application.

Compared with biogas bi-reforming, the presence of O₂ in tri-reforming can further inhibit coke formation to prevent the catalyst deactivation. The presence of O₂ can further reduce the energy consumption to reduce cost. There is no need to remove the O₂ from raw biogas for tri-reforming use. The O₂ can be nearly fully converted during biogas tri-reforming process due to the low content of O₂ (<6%) in biogas. These advantages of the presence of O₂ may lead to a high CH₄ conversion (typically >72%) in biogas tri-reforming. The catalysts utilized in biogas tri-reforming have similar phenomena to biogas bi-reforming: Ni as a main active metal and common supports include Al₂O₃ and ZrO₂. The H₂ to CO ratio generated from biogas tri-reforming is typically 1.0–2.3, which is suitable for syngas to be used to generate heat, methanol, liquid fuel, or hydrogen. Biogas tri-reforming is a simplified process to produce syngas but an emerging technology. Additional steam and oxygen are typically needed to add into the reactor system to obtain high reactant conversions and suitable H₂ to CO ratio for a specific target application. However, the addition of steam and oxygen can increase the cost. The additional O₂ can increase the risk of oxidation of metal particles (e.g., from metallic state Ni⁰ to Ni²⁺) because the metallic state is considered the active phase for biogas reforming. Adding too much O₂ can cause the methane combustion reaction to generate CO₂. If additional air is added into the tri-reforming reactor instead of O₂, the handling of inert N₂ can be an issue. The H₂O typically is not fully converted so that the handling of unreacted H₂O can be another issue for biogas tri-reforming. The unreacted H₂O can aggregate to form condensed liquid water in the pipe tubing to block the gas flowing. The CH₄ dry reforming reaction can also be suppressed by the competition of H₂O to react with CH₄, which can decrease the CO₂ conversion in biogas tri-reforming. Ensuring the accurate and constant flow of the H₂O feed into the tri-reforming reactor is a challenge. A superior heat and mass management is also needed in biogas tri-reforming.

Low Temperature Biogas Reforming

The typical high temperature (~700–950°C) used in biogas reforming can cause carbon deposition and metal Ni sintering, which can result in the catalysts deactivating. In addition, the high-temperature operation leads to high operation costs (Wang et al., 2018b). Therefore, the development of catalysts and technologies for biogas reforming at low temperatures (<600°C) is needed. Low-temperature biogas reforming typically takes place at a temperature of 25–550°C and pressure of 1 bar, shown in Tables S1 and S2. The fixed-bed reactor and plasma reactor are mainly used for the reaction. The plasma reactor is typically a dielectric barrier discharger reactor. The use of plasma in biogas reforming provides the advantages of
| Catalyst                        | Reaction Conditions                              | CH<sub>4</sub> Conv. (%) | CO<sub>2</sub> Conv. (%) | H<sub>2</sub>/CO | Coke Rate (g<sub>coke</sub>/g<sub>cat</sub>*h<sub>reactor</sub>) | Ref.                                      |
|--------------------------------|--------------------------------------------------|--------------------------|--------------------------|----------------|----------------------------------------------------------------|------------------------------------------|
| Ni-Pt/Co<sub>0.6</sub>Ce<sub>0.6</sub>Zr<sub>0.4</sub>O<sub>2</sub> | 430°C, CH<sub>4</sub>/CO<sub>2</sub> = 1/1, ~60,000 h<sup>-1</sup>, 1 bar, u-tube reactor | 8                        | 14                       | 0.4          | 5.5 × 10<sup>-4</sup>                                                  | (Sokfoun et al., 2019)                  |
| ZrO<sub>2</sub>/Ni-MnO<sub>2</sub>/SiO<sub>2</sub>          | 400°C, CH<sub>4</sub>/CO<sub>2</sub> = 1/1, 1 bar, fixed-bed reactor | 3                        | 5                        | 0.6          | 1 × 10<sup>-3</sup>                                                   | (Yao et al., 2017)                      |
| ZrO<sub>2</sub>/Ni-MnO<sub>2</sub>/SiO<sub>2</sub>          | 500°C, CH<sub>4</sub>/CO<sub>2</sub> = 1/1, 1 bar, fixed-bed reactor | 18                       | 23                       | 0.6          | 1.4 × 10<sup>-3</sup>                                                  | (Yao et al., 2017)                      |
| Ni<sub>22</sub>La<sub>0.025</sub>Mg<sub>2.95</sub>Al<sub>0.225</sub>O<sub>4</sub> | 550°C, CH<sub>4</sub>/CO<sub>2</sub> = 1/1, 20,000 h<sup>-1</sup>, tubular quartz reactor | ~32                      | ~35                      | ~0.9        | ~0.84                                                               | (Liu et al., 2016)                      |
| Ni-Mg-Al hydroxide             | 550°C, CH<sub>4</sub>/CO<sub>2</sub> = 1/1, 20,000 h<sup>-1</sup>, tubular quartz reactor | ~40                      | ~40                      | ~1.0        | –                                                                   | (Débek et al., 2016)                    |
| Rh-Co/SBA-15                   | 550°C, CH<sub>4</sub>/CO<sub>2</sub> = 1/1, 67 L/(g*h)<sup>-1</sup>, 1 bar | ~50                      | ~43                      | ~1.1        | –                                                                   | (El Hassan et al., 2016)                |
| Ni/Al<sub>2</sub>O<sub>3</sub>-MgO          | Room temperature, CH<sub>4</sub>/CO<sub>2</sub> = 1/1, 364 h<sup>-1</sup>, 300 J/mL, 1 bar, dielectric barrier discharge plasma reactor | 75                       | 73                       | 1.0         | ~3.7 × 10<sup>-3</sup>                                                 | (Khoja et al., 2018)                    |
| La<sub>2</sub>O<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub> | 25°C, CH<sub>4</sub>/CO<sub>2</sub> = 1/2, 24 kv, 8 W, 800 Hz, plasma discharge coaxial packed-bed reactor | 33                       | 12                       | 0.67        | –                                                                   | (Yap et al., 2018)                      |
| La<sub>2</sub>O<sub>3</sub>/Al<sub>2</sub>O<sub>3</sub> | 300°C, CH<sub>4</sub>/CO<sub>2</sub> = 1/2, 22 kv, 8 W, 800 Hz, plasma discharge coaxial packed-bed reactor | 48                       | 10                       | 0.63        | –                                                                   | (Yap et al., 2018)                      |
| Ni-K/Al<sub>2</sub>O<sub>3</sub>          | 160°C, CH<sub>4</sub>/CO<sub>2</sub> = 1/0.67, 16 W, coaxial dielectric barrier discharge plasma reactor | 32                       | 23                       | 1.9         | 0.035                                                              | (Zheng et al., 2018)                    |
| NiFe<sub>2</sub>O<sub>3</sub>-SiO<sub>2</sub>  | ~193°C, CH<sub>4</sub>/CO<sub>2</sub> = 1/1, 160 W, coaxial dielectric barrier discharge plasma reactor | 80                       | 70                       | 1.0         | 1.2 × 10<sup>-4</sup>                                                 | (Zheng et al., 2015)                    |
| Ni-La/ZrO<sub>2</sub>           | 311°C, CH<sub>4</sub>/CO<sub>2</sub> = 1/1, 7 mA, 0.8 kV, 5.6 W, reactor with electric field | 34                       | 43                       | 0.8         | –                                                                   | (Yabe et al., 2017)                     |
| No catalyst                    | ~387°C, CH<sub>4</sub>/CO<sub>2</sub> = 1/1, 92 h<sup>-1</sup>, 370 J/mL, dielectric barrier discharge plasma alumina reactor | 74                       | 68                       | ~0.9        | –                                                                   | (Khoja et al., 2017)                    |
| Ag-La loaded protonated carbon nitrides nanotubes (pCNNT) | 100°C, CH<sub>4</sub>/CO<sub>2</sub> = 1/1, fixed-bed reactor, visible light from solar simulator, 1 bar | –                        | –                        | 0.2         | –                                                                   | (Tahir et al., 2019)                    |
| Ni-CeO<sub>2</sub>-Al<sub>2</sub>O<sub>3</sub> | 550°C, CH<sub>4</sub>/CO<sub>2</sub> = 1/1, fixed-bed reactor | ~36                      | ~41                      | ~0.9        | –                                                                   | (Liang et al., 2020)                    |
| Cu<sub>19.8</sub>Re<sub>0.2</sub>          | Room temperature, CH<sub>4</sub>/CO<sub>2</sub> = 1/1, Harrick reactor, white light, 1 bar, 19.2 W/cm<sup>2</sup> | ~58                      | –                        | ~1.0        | –                                                                   | (Zhou et al., 2020)                     |
| Rh/La<sub>2</sub>O<sub>3</sub>-ZrO<sub>2</sub> | 400°C, CH<sub>4</sub>/CO<sub>2</sub>/H<sub>2</sub>O = 1/1/3, 30,000 h<sup>-1</sup>, 1 bar, fixed-bed tubular reactor | ~9                       | –                        | –           | –                                                                   | (Angel et al., 2016)                    |
| Pt-NiMg/Co<sub>0.6</sub>Zr<sub>0.4</sub>O<sub>2</sub> | 500°C, CH<sub>4</sub>/CO<sub>2</sub>/H<sub>2</sub>O = 1/1/1, 136,000 h<sup>-1</sup>, 1 bar | 78                       | 32                       | 1.2         | –                                                                   | (Elsayed et al., 2018)                  |
low energy input, easy operation, and low installation cost. The energy efficiency of the plasma reactor can be improved by using different heterogeneous catalysts (Khoja et al., 2018). Generally, the CH4 to CO2 ratio for biogas reforming is in the range of 0.5–1.5, the CH4 to H2O ratio for bi-reforming and tri-reforming is in the range of 0.3–3.0, and the CH4 to O2 ratio for tri-reforming is in the range of 5–10. The catalysts used in low-temperature biogas reforming are mainly Ni based and precious metal based (e.g., Rh, Ru, and Pd). Different catalyst supports such as Ce0.6Zr0.4O2, ZrO2, SiO2, Al2O3, and SBA-15 have been used. The CH4 and CO2 conversions obtained from biogas dry reforming are both generally lower than 81%, and the produced H2 to CO ratio is in the range of 0.2–2.0. The CH4 conversion obtained from biogas bi-reforming and tri-reforming is generally lower than 79%, and the produced H2 to CO ratio is in the range of 1.2–3.6. The CO2 conversion obtained from biogas bi-reforming and tri-reforming is generally lower than 37%. The coke rate of catalysts varies from being negligible to /C240.84 gcoke/(gcat*h).

The low reactant conversions are the major challenge for low-temperature biogas reforming due to the endothermic nature of biogas reforming. The low reaction temperature regime can induce significant selectivity losses due to some side reactions (e.g., WGS) during biogas reforming. To improve the reactant conversions, researchers have developed various advanced catalysts. Majewski and Wood (Majewski and Wood, 2014) studied the tri-reforming of biogas over Ni@SiO2 core-shell catalysts at 550°C. The core-shell structure of the Ni@SiO2 catalysts limits access to the silica surface to prevent the structure damage of the support and the reduction of the catalysts’ surface area. The core-shell structure provides advantages of high utilization of metal species and enhancement of catalytic activity. At a low reaction temperature (e.g., 550°C), the thermodynamic equilibrium favored the forward direction of the WGS reaction causing conversions of CO and H2Ot oH2 and CO2. Additionally, the CO2 could be produced by the complete oxidation of CH4. The increased CO2 and H2 concentration in generated syngas contributed to the low CO2 conversion (<5%) and high H2 to CO ratio (>3) (Majewski and Wood, 2014).

Elsayed et al. (Elsayed et al., 2018) studied the bi-reforming of biogas over Pt-NiMg/Ce0.6Zr0.4O2 catalysts at 500°C. For low-temperature biogas bi-reforming, a challenge can arise because the WGS reaction is more prevalent, generating less H2, whereas reaction kinetics dominates over thermodynamics for dry reforming. An optimum reactant ratio in the feed can be investigated to still allow the dry reforming reaction to occur while improving the H2 to CO ratio. Different reactant feed compositions, CH4 to CO2 to H2O ratio of 1/1/1 and 1/0.33/0.67, were investigated. When the CO2 to CH4 ratio decreased, the H2 to CO ratio increased from 1.2 to 1.9. The CH4 conversion decreased from 78% to 33% as the feed became stoichiometric. The CO2 conversion increased from 32% to 36% because the CO2 was diluted. The results indicate that reasonable conversions of CH4 and CO2 and H2 to CO ratio near 2 can be achieved at a low reaction temperature, which may enable intensified processes for the conversion of biogas to value-added products (Elsayed et al., 2018). To further improve the reactant conversions and reduce the cost, more research is needed for the low-temperature biogas reforming. The development of novel catalysts, optimization of catalyst synthesis, and introduction of new technologies can be a focus in the future. Developing advanced catalysts without using new technologies to obtain desired reactant conversions and H2 to CO ratio is a challenge.

| Catalyst            | Reaction Conditions                                      | CH4 Conv. (%) | CO2 Conv. (%) | H2/CO | Coke Rate (gcoke/(gcat*h)) | Ref.                  |
|---------------------|---------------------------------------------------------|---------------|---------------|-------|---------------------------|-----------------------|
| Pt-NiMg/Ce0.6Zr0.4O2| 500°C, CH4/CO2/H2O = 1/0.33/0.67, 136,000 h⁻¹, 1 bar   | 33            | 36            | 1.9   | Negligible                | (Elsayed et al., 2018) |
| Ni@SiO2             | 550°C, 1 bar, CH4/CO2/CO/2/H2O = 1/0.5/1/0.5, fixed-bed reactor | ~23           | ~3            | ~3.6  | 0.025                     | (Majewski and Wood, 2014) |
| Ni-Mg/La0.1Zr0.9O2   | 200°C, 3 mA, 60,000 mL*(g*h)⁻¹, CH4/CO2/CO/2/H2O = 1/0.33/0.17/0.33, fixed-bed reactor with electric field | ~12           | –             | 3.2   | –                         | (Yabe et al., 2018)    |
| Ni-Mg/La0.1Zr0.9O2   | 200°C, CH4/CO2/CO2/2/H2O = 1/0.33/0.17/0.33, 2.3 W, fixed-bed reactor with electric field | 30            | –             | 1.9   | –                         | (Oguri et al., 2017)   |

Table 5. Continued
See also Table S2.

The low reactant conversions are the major challenge for low-temperature biogas reforming due to the endothermic nature of biogas reforming. The low reaction temperature regime can induce significant selectivity losses due to some side reactions (e.g., WGS) during biogas reforming. To improve the reactant conversions, researchers have developed various advanced catalysts. Majewski and Wood (Majewski and Wood, 2014) studied the tri-reforming of biogas over Ni@SiO2 core-shell catalysts at 550°C. The core-shell structure of the Ni@SiO2 catalysts limits access to the silica surface to prevent the structure damage of the support and the reduction of the catalysts’ surface area. The core-shell structure provides advantages of high utilization of metal species and enhancement of catalytic activity. At a low reaction temperature (e.g., 550°C), the thermodynamic equilibrium favored the forward direction of the WGS reaction causing conversions of CO and H2Ot oH2 and CO2. Additionally, the CO2 could be produced by the complete oxidation of CH4. The increased CO2 and H2 concentration in generated syngas contributed to the low CO2 conversion (<5%) and high H2 to CO ratio (>3) (Majewski and Wood, 2014).
The reduction of catalysts plays a significant role in biogas reforming. For example, the NiO in the nickel-based catalysts needs to be reduced to metal Ni prior to the reforming reaction. Yao et al. (Yao et al., 2017) studied the dry reforming of biogas over ZrOx/Ni-MnOx/SiO2 catalysts at 500°C. It was reported that Zr improved catalytic activity but the reaction would suffer from coke deposition. The promoter Mn improves the dispersion of Ni species, which enables the catalyst to minimize coke deposition. Catalytic tests showed that the reduction temperature (500°C, 550°C, 600°C, 800°C) had a significant influence on the catalyst activity. The reduced ZrOx/Ni-MnOx/SiO2 catalysts were characterized using techniques including temperature-programmed reduction (TPR), X-ray diffraction (XRD), and transmission electron microscopy (TEM). The TPR results showed that the reduction temperature had an influence on the state of the metal Ni species and Mn species of the catalysts. The XRD results exhibited that the diffraction peak of metal Ni increased, whereas the diffraction peak of NiO decreased with increased reduction temperature. The TEM results exhibited that the reduction temperature affected the particle size of the Ni species on the catalyst. The ZrOx/Ni-MnOx/SiO2 catalysts reduced at 550°C exhibited relatively high content of surface Ni species and small Ni species particles with narrow particle size distribution (mainly 5–6 nm), which resulted in a high catalytic activity (Yao et al., 2017). The catalyst activation process has an influence on the catalyst’s structural features such as metal particle size, metal particle size distribution, and metal species content. Small metal particle size, narrow metal particle size distribution, high metal dispersion due to appropriate promoters (e.g., Mn), and high content of surface metal species will contribute to a high catalytic activity. Beyond the reduction temperature, other catalyst activation parameters such as reduction gas, gas flow rate, and reduction time can be investigated and optimized.

To lower the reaction temperature and obtain reasonable reactant conversions for biogas reforming, new technologies including the plasma and electric field have been developed. Khoja et al. (Khoja et al., 2018) studied the dry reforming of biogas over Ni/Al2O3-MgO catalysts at room temperature in a cold plasma dielectric barrier discharge reactor, shown in Figure 5B. The reactor system mainly consists of gas cylinders, mass flow controllers, alternating current power supply, alumina dielectric tube, condenser, and gas detector. The obtained CH4 and CO2 conversions were 75% and 73%, respectively (Khoja et al., 2018). Yabe et al. (Yabe et al., 2018) studied the tri-reforming of biogas over Ni-Mg/La0.1Zr0.9O2-x catalysts at 200°C in an electrode reactor, shown in Figure 5C. The Ni-Mg/La0.1Zr0.9O2-x catalysts exhibited methane oxidation suppression because NiO-MgO on the catalyst is not reduced at such a low temperature. The steam reforming of methane reaction performed well due to the surface protonics in the electric field. The conversion of O2 was ~52%. The electric field utilized for the tri-reforming of biogas at low temperatures (e.g., 200°C) shows some interest in reducing the energy needed for the process (Yabe et al., 2018).

The feed composition, reaction temperature, voltage, current, catalyst species, and catalyst reduction temperature have an effect on the low-temperature biogas reforming performance. The addition of catalysts in the plasma reactor could promote the conversions of CH4 and CO2. The increase of the reaction temperature could promote the CH4 conversion. However, it is important to reach a balance between the high reactant conversions and low reaction temperature. The low reaction temperature (<600°C) used for the biogas dry reforming seems to help reduce the coke rate of catalysts, compared with that at higher reaction temperatures (700–900°C). The utilization of new technologies and advanced catalysts into low-temperature biogas reforming with a low energy input is a major breakthrough achieved in recent years. However, the carbon deposition on the catalyst is still an issue for the low-temperature biogas reforming application. Achieving high product selectivity at a low temperature regime is also a challenge. The current CH4 and CO2 conversions are still low (≤80% and ≤73%, respectively). The appropriate combination of plasma technology and advanced catalysts could lower the reaction temperature without decreasing the reactant conversions largely.

**REFORMING CATALYST SYNTHESIS, CHARACTERIZATION, AND REGENERATION**

**Catalyst Synthesis**

The catalysts used for biogas reforming exhibit different shapes including powder, pellet, core-shell, monolith, and foam (Figure 6A). Generally, the catalysts during the early stage of development are in the powder form. The number of potential candidate powder catalysts is reduced during the screening process, which includes synthesis and characterization. The powder catalysts need to be shaped into macroscopic forms (e.g., pellet) for industrial use to minimize the pressure drop in the reactor. Catalysts used in the industry should not only have the required mechanical properties (e.g., strength and attrition resistance) and functionality but also be able to reproduce the performance of laboratory-scale preparation (Mitchell et al., 2013). Different synthesis methods have been developed to obtain different catalysts with various shapes for biogas reforming use (Table 6).
In laboratory scale, catalysts in powder form are usually used for the screening process in biogas reforming. For example, Zanganeh et al. (Zanganeh et al., 2013) studied the dry reforming of biogas over Ni-MgO powder catalysts, which were synthesized using a coprecipitation method from solutions of Ni(NO$_3$)$_2$·6H$_2$O and Mg(NO$_3$)$_2$·6H$_2$O using K$_2$CO$_3$ as a precipitant. The excellent anti-coking performance of the Ni-MgO catalyst was attributed to the high dispersion of reduced Ni species, nickel-support interaction, and basicity of the support surface (Zanganeh et al., 2013). After the screening, the powders are shaped into macroscopic forms for future large-scale utilization. Maintaining the same catalytic performance without being brittle is a main challenge for catalysts being transferred from powder to macroscopic form. Wang et al. (Wang et al., 2010) synthesized NiO-MgO solid solution cordierite monolith catalysts using a wet impregnation method that impregnated NiO and MgO on the cordierite monolith support. The catalyst had three main phases: Ni phase, NiO phase, and NiO-MgO solid solution phase. The high stability and activity of the NiO-MgO solid solution cordierite monolith catalyst were attributed to the high nickel dispersion on the catalyst (Wang et al., 2010). During the biogas reforming process, the high metal dispersion on the catalyst provides the high resistance of coke formation.

Catalysts with other macroscopic forms such as core-shell and multicore-shell have been developed for biogas reforming too. Das et al. (Das et al., 2018) studied low-temperature dry reforming of biogas over Ni-SiO$_2$@CeO$_2$ core-shell catalysts. Uniform silica nanospheres were synthesized using a Stöber method using tetra-ethyl orthosilicate. A nickel phyllosilicate layer was formed on the SiO$_2$ spheres using an ammonia evaporation method and the obtained SiO$_2$@Ni-phyllosilicate was coated with a layer of CeO$_2$.
using a precipitation method. CeO$_2$ was chosen as the shell due to its high oxygen storage capacity and redox potential, which can reduce the coke formation under severe dry reforming conditions. The Ni-SiO$_2@$CeO$_2$ core-shell catalyst exhibited an excellent coke inhibition property without coke detected after 72 h of reforming. The high activity of the catalyst was attributed to its high Ni dispersion and reducibility (Das et al., 2018). Bian et al. (Bian et al., 2016) studied the dry reforming of biogas over Ni-Mg PSNTS@silica multicore-shell catalysts (Figure 6B), which were synthesized using a hydrothermal method. With the silica coating, the thermal stability of the Ni-Mg PSNTS@silica multicore-shell catalyst was greatly improved (Bian et al., 2016). The design of the core-shell and multicore-shell can prevent the coke formation on the catalyst, probably due to the high metal dispersion. However, the catalyst synthesis and activation process need to be strictly controlled to obtain highly dispersed and uniform metal particles.

Catalysts in pellet form are usually used in an industrial scale for biogas reforming. Pellets can be shaped from powder by simple extrusion, which is cost-effective. Arbag et al. (Arbag et al., 2015) studied the dry reforming of biogas over W- and Ni-incorporated mesoporous Al$_2$O$_3$ pellet catalysts, which were synthesized using a one-pot sol-gel route method. The W- and Ni-incorporated mesoporous Al$_2$O$_3$ pellet catalysts exhibited a high catalytic activity throughout a 150 h of time-on-stream operation at 750°C. The incorporation of W can significantly enhance and stabilize catalyst performance. Coke formation on the catalyst was minimized due to the redox ability of WO$_x$ (Arbag et al., 2015). The synthesized NiMg/Ce$_{0.6}$Zr$_{0.4}$O$_2$/Al$_2$O$_3$ pellet catalysts (Figure 6C) have been used for biogas tri-reforming to produce syngas with high reactant conversions and desirable selectivity and stability (Zhao et al., 2018a). However, the space velocity was low due to the large volume of Al$_2$O$_3$. To increase the space velocity suitable for large-scale application, the NiMg/Ce$_{0.6}$Zr$_{0.4}$O$_2$ pellets (without the Al$_2$O$_3$ support) have been developed via extrusion, where similar catalyst performance was achieved (Zhao et al., 2018b). However, pellet catalysts exhibit issues due to potential mass transfer limitation and heat transfer limitation in the continuous flow reactors.

Heat and mass transfer of catalysts are important factors in the biogas reforming process due to the combination of endothermic and exothermic reactions. In our previous research (Zhao et al., 2018b), the tri-reforming of biogas was run over cylindrical NiMg/Ce$_{0.6}$Zr$_{0.4}$O$_2$ pellet catalysts at 882°C and 3 bar in a fixed-bed reactor. The NiMg/Ce$_{0.6}$Zr$_{0.4}$O$_2$ pellet catalysts exhibited internal diffusion limitations because the

| Catalyst Shape Synthesis Method Dimension Ref. |
|-----------------------------------------------|---------------------------------|-----------------|-----------------|
| NiO-MgO cordierite Monolithic Wet impregnation 50 × 40 × 1 mm (column height × outer diameter × wall thickness) (Wang et al., 2010) |
| Ni$_x$Mg$_{1-x}$O Powder Co-precipitation 13–21 nm (particle size) (Zanganeh et al., 2013) |
| Ni/La-Si Powder One pot sol-gel 177–250 μm (particle size) (Chen et al., 2019) |
| NiO-Mg/Ce-ZrO$_2$/Al$_2$O$_3$ 10-hole ring – 19 × 16 mm (particle size) (Arbag Aboosadi et al., 2011) |
| NiW/Al$_2$O$_3$ Pellet One pot sol-gel 1–2 mm (particle size) (Arbag et al., 2015) |
| NiMg/Ce$_{0.6}$Zr$_{0.4}$O$_2$ Pellet Wet impregnation and extrusion 1.5 mm (diameter) (Zhao et al., 2018b) |
| NiMg/Ce$_{0.6}$Zr$_{0.4}$O$_2$/Al$_2$O$_3$ Pellet Wet impregnation 4.1 × 3.2 mm (length × diameter) (Zhao et al., 2018a) |
| Ni/Al$_2$O$_3$ Pellet Wet impregnation – (Park et al., 2015) |
| Ni/Al$_2$O$_3$/Ni Foam Sol-gel and impregnation – (Park et al., 2015) |
| Ni@SiO$_2$ Core-shell Water-in-oil microemulsion ~5 × 10 × 30 nm (Ni particle size × SiO$_2$ shell thickness × SiO$_2$ sphere diameter) (Wang et al., 2018a) |
| Ni-SiO$_2@$CeO$_2$ Core-shell Stöber treatment, ammonia evaporation, and precipitation – (Das et al., 2018) |
| Ni-Mg phyllosilicate nanotubes@SiO$_2$ Multicore-shell Hydrothermal treatment and silica coating – (Bian et al., 2016) |

Table 6. Shape, Dimension, and Synthesis Method of Catalysts Used for Biogas Reforming
The catalytic activity of catalysts is a combined effect of reaction parameters (such as transfer limitation, biogas feed composition, and reaction temperature) and catalyst characteristics. Tables 7 and S3 show some typical properties of catalysts used in biogas reforming. The properties of catalysts (e.g., Brunauer-Emmett-Teller [BET] surface area, pore size, pore volume, basic site, metal dispersion) have a relationship to the catalytic performance during the biogas reforming process. The screening of properties can be one strategy for evaluating catalysts. The synthesis-property-function relationships of catalysts are crucial to selecting appropriate formulation and structuring form. One challenge is that the property–function relationships of catalysts are not clear. During the biogas reforming process, carbon formation may fill some pores of the catalysts, which can change their surface characteristics.

The synthesis method and procedure have a significant influence on the catalyst's properties (e.g., surface area and pore volume) as does the selection of metal precursors. For example, the Co/Al2O3 catalysts exhibited BET surface area of 118–144 m²/g due to the different cobalt precursors (CoSO4 and Co(NO3)2) (Ding and Yan, 2002). Wang et al. (Wang et al., 2010) synthesized the NiO-MgO solid solution cordierite monolith catalysts and found the BET surface area of the cordierite was 11 m²/g. With the wet impregnation of Al2O3 superfine powders, the NiO-MgO solid solution cordierite monolith catalyst had a higher BET surface area, which was 40.2 m²/g (Wang et al., 2010). Zanganeh et al. (Zanganeh et al., 2013) studied the dry reforming of biogas over NiₓMg₁₋ₓO solid solution catalysts. The surface area of metal oxide powders at high temperatures depends on their intrinsic properties, particularly phase transformation and melting point. The nickel loading in the range of 3–7 wt% had a
| Catalyst                        | BET Surface Area (m²/g) | Pore Volume (cm³/g) | Pore Size (Å) | Basic Sites (mmol/g) | Metal Dispersion (%) | Ref.                                      |
|--------------------------------|-------------------------|---------------------|--------------|----------------------|----------------------|-------------------------------------------|
| NiO-MgO cordierite             | 40.2                    | –                   | –            | –                    | –                    | (Wang et al., 2010)                       |
| Ni,Mg₁₋ₓO                      | 65–115                  | 0.50–1.08           | 260–494      | –                    | –                    | (Zanganeh et al., 2013)                   |
| Al₂O₃                           | 192                     | 0.30                | 63–75        | –                    | –                    | (Ding and Yan, 2002; Zou et al., 2016)    |
| Co/Al₂O₃                       | 118–144                 | 0.23–0.24           | 66–78        | –                    | –                    | (Ding and Yan, 2002)                      |
| Ni/Al₂O₃                       | 130                     | 0.24                | 75           | 0.03                 | 1.0                  | (Ding and Yan, 2002; García-Vargas et al., 2014) |
| Ni/ZrO₂                         | 128                     | 0.16                | 51           | 0.002                | 0.3–2.0              | (Kumar et al., 2019; Singha et al., 2016a; Song et al., 2010) |
| ZrO₂                            | 130                     | 0.11                | 35           | –                    | –                    | (Singha et al., 2016a; Song et al., 2010) |
| Ce₂₋ₓZrₓO₂                      | 93–232                  | 0.06–0.40           | 32–57        | –                    | –                    | (Sukonket et al., 2011)                   |
| Ni/Ceₓ₋ₓZrₓO₂                   | 59–215                  | 0.06–0.30           | 41–53        | –                    | 4.2–7.4              | (Sukonket et al., 2011)                   |
| Ni/CeO₂                         | 1.8–9.2                 | 0.02–0.08           | 89           | 0.02                 | –1                   | (Pal et al., 2015; Pino et al., 2011; Singha et al., 2016a; Vite et al., 2014) |
| Ni/Mg/β-SiC                     | 21                      | 0.09                | –            | 0.01                 | –                    | (García-Vargas et al., 2015)              |
| Ni-alumina aerogel              | 370                     | 1.18                | 127          | 19                   | –                    | (Yoo et al., 2015)                       |
| Ni-alumina xerogel              | 322                     | 0.58                | 72           | –                    | 12                   | (Yoo et al., 2015)                       |
| Ni/SiC                          | 26                      | 0.18                | –            | 0.003                | 1.6                  | (García-Vargas et al., 2014)              |
| Ni/YSZ (yttria-stabilized zirconia) | 11                   | 0.06                | –            | 0.02                 | 1.5                  | (García-Vargas et al., 2014)              |
| Ni/zeolite L                    | 95                      | 0.45                | 96           | –                    | 4.7                  | (Izquierdo et al., 2014)                  |
| Rh-Ni/zeolite L                 | 64                      | 0.63                | 198          | –                    | 10.5                 | (Izquierdo et al., 2014)                  |
| Mg/CoO₂–ZrO₂                    | 138                     | 0.16                | 45           | –                    | –                    | (Kumar et al., 2019; Singha et al., 2016a) |
| Ni/MgO                          | 241                     | 0.74                | 124          | 0.6                  | –                    | (Kumar et al., 2019; Singha et al., 2016a) |
| Ni/Ceₓ₋ₓZrₓO₂                   | 122                     | 0.18                | 61           | –                    | –                    | (Kumar et al., 2019; Singha et al., 2016a) |
| Ni-Mg/CoO₂–ZrO₂                 | 126–133                 | 0.18–0.20           | 46–47        | –                    | –                    | (Kumar et al., 2019; Singha et al., 2016a) |
| Ce/Al₂O₃                       | 195                     | 0.76                | 150          | –                    | –                    | (Izquierdo et al., 2013)                  |
| Zr/Al₂O₃                       | 180                     | 0.67                | 144          | –                    | –                    | (Izquierdo et al., 2013)                  |
| Ni/Co/Al₂O₃                    | 163                     | 0.59                | 144          | –                    | 0.06                 | (Izquierdo et al., 2013)                  |
| Ni/Co/Al₂O₃                    | 151                     | 0.60                | 153          | –                    | 0.08                 | (Izquierdo et al., 2013)                  |
| Rh-Ni/Co/Al₂O₃                  | 157                     | 0.60                | 150          | –                    | 0.12                 | (Izquierdo et al., 2013)                  |
| NiMo–C                          | 138                     | 0.27                | 73           | –                    | –                    | (Zou et al., 2016)                       |
| NiMoC–La                        | 143                     | 0.27                | 73           | –                    | –                    | (Zou et al., 2016)                       |
| NiMoC–Co                        | 130                     | 0.25                | 74           | –                    | –                    | (Zou et al., 2016)                       |
| NiMoC–K                         | 104                     | 0.18                | 62           | –                    | –                    | (Zou et al., 2016)                       |
| Ni/CeO₂–ZrO₂                    | 4                      | –                   | –            | 0.0008               | 1.5                  | (Kumar et al., 2019)                      |
| Ni/SBA-15                       | 36                      | –                   | –            | 0.004                | 2.2                  | (Kumar et al., 2019)                      |

Table 7. Properties of Catalysts Used in Biogas Reforming

(Continued on next page)
minor effect on the BET surface area of the Ni$_x$Mg$_{1-x}$O solid solution catalysts. However, when the nickel loading increased to 10 wt%, the surface area increased significantly. With the further increase of nickel loading (15–25 wt %), the surface area of the Ni$_x$Mg$_{1-x}$O solid solution catalysts decreased (Zanganeh et al., 2013). García-Vargas et al. (García-Vargas et al., 2014) loaded nickel in four different supports including Al$_2$O$_3$, CeO$_2$, SiC, and YSZ. The support had a significant effect on the surface characteristics of the Ni-based catalysts. The surface area and pore volume were in the range of 7–69 m$^2$/g and 0.06–0.24 cm$^3$/g, respectively (García-Vargas et al., 2014). The synthesis parameters including phase transformation, metal loading, and support species have a large influence on the surface characteristics of catalysts. The synthesis procedure needs a strict control to obtain the desired properties of catalysts.

The surface area of catalysts can be reduced with metal loading. Ding and Yan studied the dry reforming of biogas over Al$_2$O$_3$ catalysts, which had a BET surface area of 192 m$^2$/g, pore volume of 0.30 cm$^3$/g, and pore size of 63–75 Å (Ding and Yan, 2002; Zou et al., 2016). After Co or Ni was loaded on the Al$_2$O$_3$ support, the Co/Al$_2$O$_3$ or Ni/Al$_2$O$_3$ catalysts had a lower BET surface area and pore volume than Al$_2$O$_3$ (Ding and Yan, 2002; García-Vargas et al., 2014). Some of the pores of the Al$_2$O$_3$ support might have been filled with metal oxides after the metal loading, which caused the decrease of the surface area and pore volume. Similarly, with the addition of Ni and Mg on the β-SiC support, the surface area of the Ni-Mg/β-SiC catalysts decreased, reported by García-Vargas et al. (García-Vargas et al., 2015). With the impregnation of 5 wt% of Ni on the Ce$_0.6$Zr$_{0.4}$O$_2$ support, the surface area of the Ni/Ce$_0.6$Zr$_{0.4}$O$_2$ catalysts decreased, as reported by Sukonket et al. (Sukonket et al., 2011). In addition, the surface area and pore volume of the Ni/Ce$_0.6$Zr$_{0.4}$O$_2$ catalysts decreased with the increase of the calcination temperature from 650 to 800°C.

Based on research described above, the BET surface areas of reforming catalysts are mainly in the range of 2–370 m$^2$/g. The BET surface area of catalysts is affected by many parameters including the metal precursor, pore structure of supports, synthesis method, and dispersion of active metal on the support. A suitable metal loading (e.g., 10 wt%) on the support can reach the highest surface area of the catalyst. The pore volumes of reforming catalysts are mainly in the range of 0.004–1.18 cm$^3$/g. Generally, the addition of metals on the support will result in decreased pore volume of catalysts because the metals can lead to partial blockage of the support channels. The pore sizes of reforming catalysts are mainly in the range of 32–494 Å. The pore size of catalysts can affect the product selectivity: small molecules of products can diffuse out of catalyst pores, but large molecules of products can block the catalyst pores. The pore volume and pore size of catalysts can be affected by the particle size, synthesis method, and synthesis procedure. The dispersion of nickel metal on various supports (e.g., Al$_2$O$_3$, ZrO$_2$, CeO$_2$, MgO, zeolite) varies largely from 0.06% to 19%. A high metal dispersion of catalysts is effective to reduce the carbon deposition during biogas reforming process. The metal dispersion of catalysts can be affected by the precursor species, support species, synthesis method, and metal loading level. For example, incorporating a small amount of noble metal (e.g., Rh) can increase the dispersion of nickel metal on the catalyst (Izquierdo et al., 2013, 2014). The basic sites of reforming catalysts are mainly in the range of 0.0008–0.04 mmol/g. The optimal catalyst can exhibit these properties: surface area of >100 m$^2$/g, pore volume of >0.2 cm$^3$/g, pore size of ~50 Å, basic site of >0.02 mmol/g, and metal dispersion of >2%. No clear relationship is found yet between the catalyst’s property and biogas reforming reaction type. The catalyst’s surface area, pore volume, and pore size do not vary largely as a function of the active metal (e.g., Ni and Co), probably due to the same support with only small amounts of active metals. However, the catalyst’s surface area, pore volume, and basic site vary largely as a function of the support (e.g., Al$_2$O$_3$, SiO$_2$, MgO, CeO$_2$, and ZrO$_2$). In the future, more studies on the relationship between catalyst physicochemical properties and catalytic activity can be conducted.

| Catalyst | BET Surface Area (m$^2$/g) | Pore Volume (cm$^3$/g) | Pore Size (Å) | Basic Sites (mmol/g) | Metal Dispersion (%) | Ref. |
|----------|-----------------------------|------------------------|--------------|---------------------|----------------------|------|
| Ni-Y/KIT-6 | 199 | 0.2 | 40 | – | – | (Swirk et al., 2019) |
| Ru/SiO$_2$ | 930 | 1.01 | 48 | – | – | (Das et al., 2019) |
| Sr$_0.92$Y$_{0.08}$Ti$_{1-x}$Ru$_x$O$_{3-d}$ | 37 | 0.27 | 296 | – | – | (Kim et al., 2019a) |

Table 7. Continued
See also Table S3.
Catalyst Regeneration

Lifetime (Table 8) is an important factor for catalyst use in industry. Catalysts could be deactivated by coke deposition, sintering of metallic phase, and sintering of support (Vita et al., 2014). However, the spent catalysts can be regenerated to recover their catalytic activity. The typical regeneration methods for spent reforming catalysts include heating in air, activation in steam, and reduction using H2. A main challenge for the catalyst regeneration is the addition of regeneration cost and decreased catalytic performance.

Carbon deposition on the catalyst can be observed using an apparatus such as a transmission electron microscope. Wang et al. (Wang et al., 2019) studied the dry reforming of biogas over La(Co0.1Ni0.9)0.5Fe0.5O3 catalysts. The nanosheet-like carbon species can be observed on the used La(Co0.1Ni0.9)0.5Fe0.5O3 catalyst, indicating the deposition of carbon on the catalyst during the biogas reforming process. Methane decomposition (Equation 11) and Boudouard reaction (Equation 7) can result in serious carbon deposition issues for the catalyst. The deposited carbon species on the catalyst can be removed by oxygen species adsorbed either on the perovskite matrix or on the metallic particle. The oxygen vacancy can provide active sites for the dissociation of CO2, whereas the active oxygen species can promote the removal of the deposited carbon species. The spin-state features and multiple valences of Co cations may affect the active oxygen species in the perovskite, which affected the removal of deposited carbon species. The overall low carbon formation rate of the La(Co0.1Ni0.9)0.5Fe0.5O3 catalyst might be attributed to the stable perovskite phase that enhanced the metal–support interaction (Wang et al., 2019).

\[
\text{CH}_4 \rightarrow \text{C} + 2\text{H}_2 \quad \Delta H^\circ = 74.9 \text{kJ/mol} \quad \text{(Equation 11)}
\]

Researchers have developed various catalysts to improve the lifetime during the biogas reforming process. Vita et al. (Vita et al., 2014) studied the tri-reforming of simulated biogas over Ni/CeO2 catalysts, which showed stability during 150 h of reaction. The coexistence of different nickel phases (such as Ni0 and NiO) and the characteristics of the CeO2 support can play a role in the stability of the Ni/CeO2 catalyst (Vita et al., 2014). Mortensen and Dybkjær (Mortensen and Dybkjæer, 2015) studied the industrial-scale steam reforming of CO2-rich gas. The nickel-based catalysts revealed insignificant carbon formation (500–1,000 ppm) after 2 years of operation under severe conditions. In general, large nickel particles

| Catalyst Lifetime Regeneration Ref. |
|-------------------------------------|
| Boron nitride defect-confined Ni ≥ 125 h – (Bu et al., 2020) |
| Ni/CeO2 ≥ 150 h – (Vita et al., 2014) |
| Ni-Mo-MgO ≥ 850 h – (Song et al., 2020) |
| Ni/MgAl2O4 insignificant carbon formation after 2 years – (Mortensen and Dybkjær, 2015) |
| 0.15%Ni/1.7%Ln/Al2O3 CH4 conversion decreased from 92% to 62% after 600 h – (Slagtern et al., 1997) |
| Ni/ZrO2 CH4 conversion decreased from 86% to 84% after 600 h – (Wei et al., 2000) |
| Ni-0.5Mo/SBA-15 CH4 conversion remained at ~94% for 600 h – (Huang et al., 2011) |
| 8%Ni/SBA-15/Al2O3/FeCrAl CH4 conversion decreased from 92% to 89% after 1,400 h – (Wang et al., 2008) |
| 9% La0.9Ni0.1O2/Al2O3 – In air at 600°C (Liu and Au, 2003) |
| Ni/SiO2MgO – O2 + CO2 was better than CO2 (O2 accelerates coke removal) (Assabumrungrat et al., 2009) |
| NiCo/MgO-ZrO2 – 1 h regeneration: air was better than N2 or H2 (Fan et al., 2011) |
| Ni/Al2O3 – Flow O2, then reduced with H2 at 650°C for 1 h (Quincoces, 2004) |

Table 8. Lifetime and Regeneration of Biogas Reforming Catalysts

Catalyst Regeneration

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(e.g., >10 nm) are more prone to carbon formation. Noble metals are typically more resistant than Ni toward carbon formation. The reforming of CO₂-rich gas in an industrial setting would require co-feeding of steam to reduce carbon formation (Mortensen and Dybkjær, 2013). Slagtern et al. (Slagtern et al., 1997) studied the dry reforming of biogas over 0.15%Ni/1.7%Ln/Al₂O₃ catalysts. The initial coke formation was 0.13 wt% after 20 h of operation and then became 0.16 wt% after ~100 h. The Ni sintering was initially a major reason for deactivating the 0.15%Ni/1.7%Ln/Al₂O₃ catalyst, whereas coking became increasingly important with longer reaction time. The coking was initially high and then stabilized with a longer reaction time (Slagtern et al., 1997).

Wei et al. (Wei et al., 2000) studied the dry reforming of biogas over Ni/ZrO₂ catalysts, which exhibited a coke rate of 1.8 × 10⁻⁴ g_coke/(g_cat* h) after 600 h. There might be a periodic cycle of carbon deposition and elimination on the nickel surface of the Ni/ZrO₂ catalyst (Wei et al., 2000). Huang et al. (Huang et al., 2011) studied the dry reforming of biogas over Ni-0.5Mo/SBA-15 catalysts. The CH₄ conversion remained at ~94%, and the coke rate was 7.3 × 10⁻⁴ g_coke/(g_cat* h) after 600 h, indicating the superior stability of the Ni-0.5Mo/SBA-15 catalysts. The carbon deposition was a main reason for deactivating the catalyst during biogas dry reforming of biogas; the sintering of nickel particles was another reason. The main factors that prevented the carbon deposition of the Ni-0.5Mo/SBA-15 catalyst were strong basicity, strong metal–support interaction, small metal particles, and the formation of Mo₂C species. The changes in catalyst composition and reactivity of active phases can affect the structure and morphology of the deposited carbon (Huang et al., 2011). The lifetime of catalysts can be affected by the coexistence of different metal phases. The addition of appropriate noble metal can promote the catalyst stability. Understanding the coke formation mechanism during the biogas reforming is a challenge. Whether there is a periodic cycle of carbon deposition and elimination on the catalyst can be further studied and confirmed. Some new advanced technologies (e.g., in situ TEM) can be utilized to observe the catalysts in real time under real experimental conditions.

Coke can be removed through the flowing of O₂, air, steam, and other gases. Liu and Au (Liu and Au, 2003) found that the catalyst activity and carbon deposition depended on parameters such as surface composition, metal-support interaction, metal crystalline structure, support acidity, and support basicity. The used La₃NiO₅/Al₂O₃ catalysts in biogas dry reforming process almost restored the catalytic performance after regeneration at 600 °C in air by burning off the carbon. The La₃NiO₅/Al₂O₃ catalyst exhibited decreasing coke formation with the catalyst calcination temperature increase from 500 to 800 °C due to the formation of a stable NiAl₂O₄ spinel structure (Liu and Au, 2003). Quincoces et al. (Quincoces, 2004) studied the dry reforming of biogas over Ni/Al₂O₃ catalysts. The used Ni/Al₂O₃ catalysts were regenerated with O₂, followed by reduction with H₂. The CH₄ conversion over regenerated Ni/Al₂O₃ catalysts decreased by 28% than fresh Ni/Al₂O₃ catalysts (Quincoces, 2004).

In summary, the lifetime of reforming catalysts can vary between 150 h and 2 years. A periodic cycle of carbon deposition and elimination may take place on the catalysts. The lifetime of catalysts is affected by the catalyst synthesis method such as calcination and activation procedure, pore structure of catalysts, support characteristics of catalysts, and reaction conditions during biogas reforming process. A high metal-support interaction, more oxygen species, strong basicity, small metal particles, and the utilization of noble metals on catalysts can inhibit the carbon formation, thus increasing the lifetime of catalysts. In addition, complete removal of impurities in feedstock gases can prevent catalyst poisoning. One challenge for the catalyst regeneration is to keep the same catalytic activity as exhibited by the fresh catalyst. Flowing air or O₂ seems a promising method to effectively regenerate catalysts, but the temperature and oxygen concentration need to be optimized. The catalysts can be regenerated either outside the reactor or directly inside. However, the regeneration may cause catalyst destruction and agglomeration of metals, which will reduce catalyst lifetime. The regeneration of used catalysts can also increase the cost of the biogas reforming process. Catalysts with a long enough lifetime will be desired for industrial applications.

**INDUSTRIAL APPLICATION**

**Syngas Application**

Syngas derived from biogas reforming is a gas mixture that mainly consists of H₂ and CO. It may contain CO₂, CH₄, N₂, and other gases, depending on the biogas composition, reforming reaction performance, and posttreatment. Syngas can be utilized as raw material for the production of fuels (e.g., hydrogen, synthetic gasoline, dimethyl ether, ethanol, methanol) and other valuable chemicals (Hernandez and Martin,
Researchers have recently developed methods to remove impurities from syngas. Spies et al. (Spies et al., 2017) used a warm (e.g., 250–450°C) cleanup technique to purify the coal-derived syngas for downstream conversions such as methanol synthesis. The coal-derived syngas mainly contained H2, CO, CO2, N2, CH4, and C2H4. A multicontaminant removal method using different materials including ZnO, Na2CO3, CuNi, Ir-Ni/MgAl2O4, and CuZnAl was employed. More than 99% of sulfur in the syngas can be removed using this method. The ZnO sorbents were proven desulfurization materials with fast kinetics and substantial capacity (Spies et al., 2017). Hernandez and Martin (Hernandez and Martin, 2018) purified the syngas (mainly containing H2, CO, CO2, N2, H2O, and hydrocarbons) derived from biogas tri-reforming by using a flash separator to remove water first. The traces of hydrocarbons and N2 were removed using a pressure swing adsorption (PSA) system. Second, the H2 to CO ratio was adjusted through a H2 tuned-up section consisting of a PSA membrane unit for H2 recovery, a bypass, and a WGS reactor. Last, CO2 was removed using zeolite 5A or 13X (Hernandez and Martin, 2018). The effective removal of impurities from syngas can prevent damage to reactors and decreased catalyst activity. The better ways to effectively remove impurities from syngas with low-processing costs needs more study. The efficient separation of H2 from CO in syngas would be an interesting topic to study in the future.

Techno-economics of Biogas Conversion

In some parts of the United States, biogas is flared due to infrastructure challenges, wasting valuable resources that could be converted to energy and/or chemicals. Techno-economics analyses (Tables 9 and S4) summarize the economic feasibility of biogas utilization and application via using different software such as Aspen Plus and Aspen HYSYS v9.0. The analysis can be performed in both small-scale and large-scale systems. A major challenge for biogas application is the low conversion efficiency. The biogas conversion efficiency here is determined based on energy recovery unless specified.

Biogas can be simply used to generate electricity. However, the low conversion efficiency of biogas to electricity is an issue for biogas utilization. For example, White et al. (White et al., 2011) analyzed small-scale biogas conversion systems located on multiple farms in ON, Canada. The study found that beef farms with ≥78 animals and dairy farms with ≥33 animals can operate economically attractive biogas systems (White et al., 2011). Pipatmanomai et al. (Pipatmanomai et al., 2009) conducted an economic assessment of biogas conversion to electricity in small pig farms (170 breeder swines and 255 piglets) in Thailand at 45% of government subsidy. The payback period of this system without H2S removal was approximately 4 years. With H2S removal, the payback period was approximately 8 years. The electricity price and government subsidy have a major effect on the payback period. The H2S removal can add high operational costs to the proposed system, but it is strongly recommended to reduce the emission of pollution gas and prevent reactor engine corrosion (Pipatmanomai et al., 2009). To improve conversion efficiency, biogas can be utilized to generate other products instead of electricity.

Hydrogen has a high calorific value and can be used as a fuel. The use of hydrogen for energy offers the advantages of zero carbon and greenhouse gas emissions. However, hydrogen is highly combustible and explosive. Lachen et al. (Lachen et al., 2018) carried out a techno-economic assessment of biogas conversion to hydrogen. Raw materials were water, electricity, biogas, catalyst, iron oxide, cobalt ferrite, and membrane. The pure hydrogen yield reached 68% at 575°C. The production cost was assumed to be in the range of $4.49–16.85/kg (i.e., $4–15/kg), based on the operating temperature (475–575°C) and the grade of integration (IG, IG = 0, 0.25, 0.5, and 1) (Lachen et al., 2018). Madeira et al. (Madeira et al., 2017) conducted
an exergetic and economic evaluation of cassava wastewater biogas conversion to hydrogen. The biogas and hydrogen production cost was estimated as $0.0518/kWh and $0.13/kWh, respectively (Madeira et al., 2017). Braga et al. (Braga et al., 2013) conducted a techno-economic analysis of biogas conversion to hydrogen for fuel cell application. The analysis was based on the biogas generated by a bovine manure biodigester located in São Paulo, Brazil. The biogas conversion process mainly consisted of steam reforming, dry reforming, and shift reaction. Biogas reforming efficiency was 80%. The hydrogen production cost was estimated as $0.27/kWh with a payback period of 8 years (Braga et al., 2013). Montenegro Camacho et al. (Montenegro Camacho et al., 2017) conducted a techno-economic analysis of hydrogen production from the autothermal reforming of biogas. The cost of generating 100 Nm³/h of hydrogen was $2.78/kg (i.e., $2.5/kg) after 10 years of amortization. The longer the BioRobur plant life is, the more feasible the initial investment is. In addition, the raw material (e.g., municipal solid waste, agroindustry scraps) for the biogas production has a major effect on the hydrogen cost (Montenegro Camacho et al., 2017).

| Product          | Product Price | Conversion Process                                      | Conversion Efficiency (%) | Payback Period (year) | Other                                      | Ref.                                      |
|------------------|---------------|--------------------------------------------------------|---------------------------|-----------------------|--------------------------------------------|-------------------------------------------|
| Electricity      | –             | Use an engine                                          | ~35                       | <10                   | Small scale cattle farms                   | (White et al., 2011)                     |
| Electricity      | 0.067 $/kWh   | Engine with electric generator                         | 20.8                      | ~4–8                  | Small pig farm                             | (Pipatmanomai et al., 2009)              |
| H₂               | ~9.99 $/kg    | Biogas dry reforming and steam-iron process            | >45                       | –                     | Plant model (1,350 kg/h biogas)            | (Lachén et al., 2018)                    |
| H₂               | –             | Steam reforming, shift reaction, and PSA              | 79                        | 7                     | Large-scale wastewater plant (generating ~4019 m³/day biogas) | (Madeira et al., 2017)                  |
| H₂               | –             | Biogas autothermal reforming                           | 65                        | –                     | BioRobur plant (generating 100 Nm³/h H₂)   | (Montenegro Camacho et al., 2017)        |
| Methanol         | 0.9 $/gallon  | Biogas cleaning/reforming and methanol synthesis       | –                         | –                     | Plant model (12,080,000 m³/year biogas)    | (Hernández and Martín, 2016)             |
| Methanol         | ~400 $/metric ton | Biogas cleaning/reforming and methanol synthesis     | –                         | Not economically feasible | Large-scale plant (generating 5,900 Nm³/h biogas) | (Sheets and Shah, 2018)                 |
| Liquid fuel      | ~0.79 $/kg (diesel) | Tri-reforming and FTS (carbon conversion efficiency) | 54                        | Not economically feasible | Plant model (27.22 kg/s CO₂ rich natural gas) | (Graciano et al., 2018)                 |
| Liquid fuel      | –             | Tri-reforming and FTS                                  | 45                        | –                     | Commercial scale plant (2,500 scfm LFG)    | (Zhao et al., 2019a)                    |
| Liquid fuel      | –             | Biogas cleaning/reforming, FTS, hydrocracking, and distillation | 54 (mass basis)          | –                     | Plant model (2,000 Nm³/h biogas)           | (Okeke and Mani, 2017)                  |
| Compressed natural gas | –             | Gas pressurizing and impurity removal                  | –                         | –                     | Medium-sized landfill model                | (Winslow et al., 2019)                  |
| Wax              | ~2.77 $/kg    | Biogas steam reforming, FTS, and product separation   | 56                        | ~7                    | Small-scale plant model (~200 kg/h biogas) | (Herz et al., 2017)                    |

Table 9. Techno-economics Analysis of Biogas Conversion
See also Table S4.
The conversion efficiency of biogas to hydrogen is relatively high, which can reach 80%. This conversion technology can be economically advantageous within ~8 years.

Methanol is the simplest alcohol and is widely used for industrial purposes. Biogas can be converted into methanol for chemical applications. Hernández and Martin (Hernández and Martin, 2016) optimized the operational process of biogas conversion to methanol using a mathematical approach. Part of the biogas was utilized to provide the energy needed for this process. The production cost of converting biogas into methanol was estimated as $1.75/gallon (Hernández and Martin, 2016). Sheets and Shah (Sheets and Shah, 2018) conducted a techno-economic analysis of the biogas conversion to methanol from a large-scale landfill or anaerobic digestion facility. The plant mainly consisted of biogas cleaning via pressurized water scrubbing, biogas reforming, syngas to methanol conversion, methanol purification, and energy recovery from unconverted gas. The methanol production cost was $2.11/kg, and it became $1.99/kg with credits for low-pressure steam (Sheets and Shah, 2018).

Liquid fuel is a value-added product that can be used to replace petroleum fuel to reduce the environmental pollution. Researchers have developed some pathways to convert biogas to liquid fuel. Graciano et al. (Graciano et al., 2018) conducted a model-based analysis of converting biogas (CO2-rich natural gas: mainly containing ~48 mol % CH4 and 30 mol % CO2) to liquid transportation fuels via tri-reforming and FTS at a tax rate of 32%/year. The plant-wide electricity consumption can be nearly fully covered by a Rankine cycle unit. However, the economic viability of the process is highly dependent on the biogas price (Graciano et al., 2018). Okeke and Mani (Okeke and Mani, 2017) carried out a techno-economic assessment of biogas-to-liquid-fuels conversion technology. The unconverted methane, syngas, and syncrude were utilized to generate electricity to meet internal plant demand while the excess power was sold to the grid. The liquid fuel price was $5.67/gal ($5.29/gasoline gallon equivalent) when the plant feed capacity was 2,000 Nm3/h. However, when the plant feed capacity increased to 20,000 Nm3/h, the liquid fuel price reduced to $2.06/gal ($1.92/gasoline gallon equivalent) (Okeke and Mani, 2017). In our previous research (Zhao et al., 2019a), a novel technology of combining tri-reforming and FTS (TriFTS) was created to convert landfill gas to liquid fuel. A preliminary economic analysis of a commercial-scale TriFTS process was conducted based on the bench-scale experimental results with a tax rate of 20%. The simulation results indicated that 45% of the energy contained in the feed LFG can be recovered in the liquid fuel produced, with the rest meeting the energy demand of the TriFTS conversion process. The break-even cost of the diesel fuel produced was estimated as $2.71/gallon if the LFG is assumed to be free of cost (Zhao et al., 2019a). Compared with generating electricity, the conversion efficiency of biogas into liquid fuel is much higher. The economic viability of the biogas conversion to liquid fuel is highly dependent on the feedstock price and current market fuel price.

Biogas can also be converted to other products (e.g., compressed natural gas [CNG] and wax). CNG is simply natural gas, mainly comprising methane, that is stored under high pressure for use as vehicle fuel. Wax is typically used for coating and candles. Winslow et al. (Winslow et al., 2019) conducted an economic assessment on LFG to vehicle fuel conversion for a waste hauling operation with 10 tractor-trailers (each tractor-trailer hauled ~19 Mg of MSW per trip to generate 1,900 m3 CH4). The CNG was produced from LFG at the landfill for vehicle use. The LFG refining process consisted of pressurizing the gas and removing impurities (e.g., siloxane, CO2, and hydrogen sulfide). The net present value per diesel gallon equivalent (DGE) of operating this scenario over a 30-year period was determined to be $2.31 at a renewable identification number (RIN) value of $0/DGE (Winslow et al., 2019). Herz et al. (Herz et al., 2017) evaluated the biogas conversion to waxes (i.e., long-chained hydrocarbons that are solid at ambient conditions) through steam reforming and FTS. The process is profitable without governmental support, compared with the established combined heat and power technology (Herz et al., 2017).

Generally, biogas can be converted to heat, hydrogen, methanol, liquid fuel, CNG, and wax. The overall efficiency of biogas conversion is approximately 35%–80%, based on the product type. Biogas conversion can be economically feasible with a typical payback period of ~4–8 years. However, the plant feed capacity, raw material price, government subsidy, and market price of final product have an effect on the economic viability. Recently, T2C-Energy LLC (Tampa, Florida) successfully tested the pilot scale of converting LFG to liquid fuels using a technology of combining tri-reforming and FTS. However, commercial operations are still progressing. Fulcrum BioEnergy Inc. (Pleasanton, California) and Velocys (Plain City, Ohio) have also developed technologies that can cost-effectively produce low-carbon liquid fuels from municipal solid waste. In general, most biogas is used for electricity production or CNG, but no commercial operations exist yet for the production of liquid fuels or chemicals. A major challenge for the biogas reforming industry is to further reduce costs.
CONCLUSIONS

Summary
Biogas reforming technologies have gained greater attention in recent years. However, no systematic study has been performed to optimize the syngas synthesis from biogas. This paper systematically reviewed the state of the art of biogas reforming, reaction mechanism, catalyst synthesis/characterization/regeneration, and syngas application. Biogas, especially from wastes, has great potential for conversion to syngas for diesel fuel use as improved climate factors. Compared with traditional fossil fuel, the fuel produced from biogas can reduce the cost of waste management and provide environmentally friendly transportation fuel.

Biogas reforming is affected by several factors including temperature, pressure, feed gas composition, space velocity, and catalyst activity. High reaction temperature (e.g., >800°C) is favored to obtain high CH4 and CO2 conversions due to the endothermic nature, but high pressure can suppress reactions to decrease CH4 and CO2 conversions. A relatively lower GHSV (e.g., <60,000 h⁻¹) can lead to higher reactant conversions due to higher residence time of reactants on the catalyst surface. Nickel-based catalysts are usually used for the reforming of biogas, and Ni nanoparticles on the CeO₂-ZrO₂ support seem a promising catalyst. Concerned about reducing energy input and coke formation, novel technologies such as plasma and solar have been developed for low-temperature biogas reforming use. However, the reactant conversions are not high yet and need to be improved further.

Compared with other reforming methods (e.g., dry and bi-reforming), tri-reforming of biogas seems to be an advanced technology toward high conversions of reactants. However, the CO₂ conversion needs to be improved during the tri-reforming process. In the future, advanced and economic catalysts with higher activity need to be developed to improve reactant conversions. In addition, the kinetics of catalyst activity could be investigated. The reforming conditions for reactants with different compositions could be further optimized. Unreacted reactants such as water could be recycled to improve the production economics. Finally, more economic evaluation and scale-up of biogas reforming (especially tri-reforming) should be investigated.

Challenges and Future Directions
The major technical challenge is that converting biogas to syngas needs to be cost-competitive. Energy consumption at high temperatures is one of the most costly aspects, and the cost of catalysts plays a significant role. The catalysts can be deactivated after a certain period. Understanding how to regenerate the used catalysts or develop catalysts with long lifetimes becomes important.

Future directions include:

1. Systematic experimental process development is needed to optimize the syngas synthesis from biogas. The molar ratio of the produced H₂ to CO and reactant conversions during biogas reforming can be affected by biogas composition, reaction temperature, reaction pressure, space velocity, time-on-stream, and catalyst property. There is a need to better control the molar ratio of the produced H₂ to CO, which is critical for the following fuel or methanol or other chemical synthesis. The process conditions need to be optimized for maximizing the reactant conversions and the overall energy efficiency.

2. Research interest in direct conversion of methane to chemicals and fuels is growing as a method to improve productivity and energy efficiency (Holmen, 2009; Karakaya and Kee, 2016; Schwach et al., 2017; Wang et al., 2017). Large reserves of methane indicate that a market exists for the large-scale conversion of methane on-site. Direct conversion of methane to liquid fuels without going through an intermediate syngas generation step can be potentially more environmentally friendly and cost-effective. In addition, one-step reforming of CH₄ and CO₂ into chemicals and fuels is a promising route. Catalysts with a high selectivity toward desired products should be designed and developed.

3. Process integration/intensification can help optimize the fuel produced from biogas, such as combining reforming and Fischer-Tropsch synthesis. Combining different systems together may help to improve the reactant conversions, recover the byproduct, and enhance the energy efficiency. An effective hybrid system may reduce the whole processing cost.
SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.101082.

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DECLARATION OF INTERESTS

Dr. Babu Joseph and Dr. John Kuhn disclose an interest in T2C-Energy, LLC. Dr. Xianhui Zhao, Dr. Babu Joseph, and Dr. John Kuhn disclose an interest in patents on related subjects issued to University of South Florida.

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Supplemental Information

Biogas Reforming to Syngas: A Review

Xianhui Zhao, Babu Joseph, John Kuhn, and Soydan Ozcan
Table S1. Other Recent Studies on Tri-reforming of Biogas to Syngas. (Related to Table 4)

| Catalyst            | Reaction conditions                                                                 | CH₄ conv. (%) | CO₂ conv. (%) | H₂/CO | Coke rate (g coke/(g cat*h)) | Ref.               |
|---------------------|------------------------------------------------------------------------------------|---------------|---------------|-------|-----------------------------|--------------------|
| Nickel-alumina      | 700°C, 269,000 mL/(g*h), 1 bar, fixed-bed reactor                                 | 76.6          | –             | 1.9–2.0| 4.7E-3                      | (Yoo et al., 2015) |
| xerogel             |                                                                                   |               |               |       |                             |                    |
| Ni/zeolite L        | 800°C, 162 h, 1 bar, fixed-bed reactor                                             | ~86           | ~24           | ~1.9  | –                           | (Izquierdo et al., 2014) |
| Rh-Ni/zeolite L     | 800°C, 162 h, 1 bar, fixed-bed reactor                                             | 96.4          | ~33           | ~1.8  | –                           | (Izquierdo et al., 2014) |
| Ni/YSZ-CeO₂         | 800°C, 1,000 h, CH₄/CO₂/O₂/H₂O = 1/1/1/1                                           | 100           | 100           | 1.0–1.1| –                           | (Kang et al., 2007) |
| Ni/MgₓTiₓO          | 850°C, 1.78 g*h/mol, 10 bar, CH₄/CO₂/O₂/H₂O = 1/0.48/0.1/0.54, fixed-bed reactor | ~85           | ~62           | –     | –                           | (Jiang et al., 2007a) |
| Ni/MgₓTi₂O₂         | 850°C, 1.78 g*h/mol, 1 bar, CH₄/CO₂/O₂/H₂O = 1/0.48/0.1/0.54, fixed-bed reactor   | ~95           | ~80           | ~1.6  | –                           | (Jiang et al., 2007a) |
| NiMg/CeₓZrₓAl₂O₃    | 800°C, 61,000 h, 1 bar, CH₄/CO₂/O₂/H₂O = 1/0.7/0.2/0.23, fixed-bed reactor          | 97            | 78            | 2.1   | 5.1E-4                      | (Walker et al., 2012) |
| Ni/Al₂O₃            | 800°C, 161 gₓ/1(g cat*h), 1 bar, CH₄/CO₂ = 4, fixed-bed reactor                    | 97            | 22            | 2.0   | –                           | (Izquierdo et al., 2013) |
| Ni/Ce₃Zr₂Al₂O₇      | 800°C, 161 gₓ/1(g cat*h), 1 bar, CH₄/CO₂ = 4, fixed-bed reactor                    | 99            | 34            | 1.5   | –                           | (Izquierdo et al., 2013) |
| Rh-Ni/Al₂O₃         | 800°C, 161 gₓ/1(g cat*h), 1 bar, CH₄/CO₂ = 4, fixed-bed reactor                    | 99            | 39            | 1.7   | –                           | (Izquierdo et al., 2013) |
| Ni/Al₂O₃            | 800°C, fixed-bed reactor                                                           | 96            | 82            | 1.4   | –                           | (Lee et al., 2003)  |
| Ni@SiO₂             | 750°C, 1 bar, CH₄/CO₂/O₂/H₂O = 1/0.5/0.1/3.0, fixed-bed reactor                    | 73            | 43            | 1.7   | 0.028                       | (Majewski and Wood, 2014) |
| Ni/Al₂O₆            | 950°C, 10,000 h, 1 bar, CH₄/CO₂/O₂/H₂O = 50/12.5/25/12.5, fixed-bed reactor       | ~95           | ~25           | –     | –                           | (Jiang et al., 2007b) |
| Catalyst | Reaction Conditions | CH4/CO2/O2/H2O Ratio | Conversion | Yield | Selectivity | Source |
|----------|---------------------|-----------------------|------------|-------|-------------|--------|
| Ni-MgO   | 850°C, 32,000 mL/(h*g cat), 1 bar, CH4/CO2/O2/H2O = 1/0.48/0.1/0.54, fixed-bed reactor | 98 | 79 | 1.5 | – | (Song and Pan, 2004) |
| Ni-Mg-CeZrO | 850°C, 32,000 mL/(h*g cat), 1 bar, CH4/CO2/O2/H2O = 1/0.48/0.1/0.54, fixed-bed reactor | 98 | 73 | 1.5 | – | (Song and Pan, 2004) |
| Ni-ZrO2  | 850°C, 32,000 mL/(h*g cat), 1 bar, CH4/CO2/O2/H2O = 1/0.48/0.1/0.54, fixed-bed reactor | 98 | 73 | 1.6 | – | (Song and Pan, 2004) |
| Ni/Al2O3 | 850°C, 32,000 mL/(h*g cat), 1 bar, CH4/CO2/O2/H2O = 1/0.48/0.1/0.54, fixed-bed reactor | 98 | 73 | 1.6 | – | (Song and Pan, 2004) |
| Ni-CeO2  | 850°C, 32,000 mL/(h*g cat), 1 bar, CH4/CO2/O2/H2O = 1/0.48/0.1/0.54, fixed-bed reactor | 95 | 73 | 1.6 | – | (Song and Pan, 2004) |
| Ni-CeZrO | 850°C, 32,000 mL/(h*g cat), 1 bar, CH4/CO2/O2/H2O = 1/0.48/0.1/0.54, fixed-bed reactor | 93 | 71 | 1.6 | – | (Song and Pan, 2004) |
| NiMo-C   | 850°C, CH4/CO2/O2/H2O = 1/0.39/0.16/0.30, fixed-bed reactor | 96 | 100 | – | – | (Zou et al., 2016) |
| NiMoC-La | 850°C, CH4/CO2/O2/H2O = 1/0.39/0.16/0.30, fixed-bed reactor | 93 | 88 | – | – | (Zou et al., 2016) |
| NiMoC-Co | 850°C, CH4/CO2/O2/H2O = 1/0.39/0.16/0.30, fixed-bed reactor | 93 | 100 | – | – | (Zou et al., 2016) |
| NiMoC-Mg | 850°C, CH4/CO2/O2/H2O = 1/0.39/0.16/0.30, fixed-bed reactor | 90 | 84 | – | – | (Zou et al., 2016) |
| NiMoC-K  | 850°C, CH4/CO2/O2/H2O = 1/0.39/0.16/0.30, fixed-bed reactor | 17 | 0 | – | – | (Zou et al., 2016) |
| Ni/Ce-Al2O3 | 800°C, 1 bar, 161 g gas*(g cat*h), 1, CH4/CO2/O2/liquid H2O = ~99 | 36 | 1.9 | – | | (Izquierdo et al., 2018) |
| Catalyst Type    | Operating Conditions | Space Velocity | Conversion (%) | Selectivity | Activity (g_mol_C2H4_per_g_cat_per_h) |
|------------------|----------------------|----------------|---------------|-------------|---------------------------------------|
| Ni/Zr-Al₂O₃     | 800°C, 1 bar, 161 g_gas*(g_cat*h), CH₄/CO₂/O₂/liquid H₂O = 1/0.67/0.25/0.0008, fixed-bed reactor | 99             | 34            | ~2          | –                                     |
|                  |                      |                |               |             | (Izquierdo et al., 2018)              |
| Rh-Ni/Ce-Al₂O₃  | 800°C, 1 bar, 161 g_gas*(g_cat*h), CH₄/CO₂/O₂/liquid H₂O = 1/0.67/0.25/0.0008, fixed-bed reactor | ~99            | ~32           | ~2          | –                                     |
|                  |                      |                |               |             | (Izquierdo et al., 2018)              |
| Ni/Al₂O₃        | 800°C, 1 bar, 17,220 mL*(g*h)-¹, CH₄/CO₂/O₂/H₂O = 1/0.23/0.07/0.46, fixed-bed reactor | –              | –             | 2.3         | 0.014                                 |
|                  |                      |                |               |             | (Kumar et al., 2019)                  |
| Ni/ZrO₂         | 800°C, 1 bar, 17,220 mL*(g*h)-¹, CH₄/CO₂/O₂/H₂O = 1/0.23/0.07/0.46, fixed-bed reactor | –              | –             | 2.2         | 3.6E-3                                |
|                  |                      |                |               |             | (Kumar et al., 2019)                  |
| Ni/TiO₂         | 800°C, 1 bar, 17,220 mL*(g*h)-¹, CH₄/CO₂/O₂/H₂O = 1/0.23/0.07/0.46, fixed-bed reactor | –              | –             | 1.9         | 2E-5                                  |
|                  |                      |                |               |             | (Kumar et al., 2019)                  |
| Ni/SBA-15       | 800°C, 1 bar, 17,220 mL*(g*h)-¹, CH₄/CO₂/O₂/H₂O = 1/0.23/0.07/0.46, fixed-bed reactor | –              | –             | 2.2         | 1E-4                                  |
|                  |                      |                |               |             | (Kumar et al., 2019)                  |
| Ni/MgO          | 800°C, 1 bar, 17,220 mL*(g*h)-¹, CH₄/CO₂/O₂/H₂O = 1/0.23/0.07/0.46, fixed-bed reactor | –              | –             | 1.8         | 1.4E-3                                |
|                  |                      |                |               |             | (Kumar et al., 2019)                  |
Table S2. Other Recent Studies on Low-temperature Biogas Reforming to Syngas. (Related to Table 5)

| Catalyst                | Reaction conditions                  | CH\(_4\) conv. (%) | CO\(_2\) conv. (%) | H\(_2\)/CO | Coke rate (g\text{coke}/(g\text{cat}*h)) | Ref.                  |
|-------------------------|--------------------------------------|---------------------|--------------------|-------------|----------------------------------------|------------------------|
| Pd-Ni-Mg/\(\text{Ce}_{0.6}\text{Zr}_{0.4}\)O\(_2\)| 366°C, CH\(_4\)/CO\(_2\) = 1/1, 68,000 h\(_{\text{l}}\), 1-bar, u-tube reactor | –                   | 10                 | –           | –                                     | (Elsayed et al., 2016) |
| Pd-Ni-Mg/\(\text{Ce}_{0.6}\text{Zr}_{0.4}\)O\(_2\)| 383°C, CH\(_4\)/CO\(_2\) = 1/1, 68,000 h\(_{\text{l}}\), 1-bar, u-tube reactor | 10                  | –                  | –           | –                                     | (Elsayed et al., 2016) |
| Ni/La\(_2\)O\(_3\)-ZrO\(_2\)| 400°C, CH\(_4\)/CO\(_2\) = 1/1, fixed-bed reactor | ~7                  | ~11                | ~0.5        | ~                                     | (Sokolov et al., 2013) |
| Ni-CaO/ZrO\(_2\)-La\(_2\)O\(_3\)| 500°C, CH\(_4\)/CO\(_2\) = 1/1, 1 bar, fixed-bed reactor | ~ 63                | ~ 80               | ~ 0.5       | ~                                     | (Bachiller-Baeza et al., 2013) |
| Ni/\(\gamma\)-Al\(_2\)O\(_3\)| 500°C, CH\(_4\)/CO\(_2\) = 1/1, 18,000 cm\(_3\)/(g\text{cat}*h), 1 bar, fixed-bed reactor | 23                  | 17                 | –           | –                                     | (Wang and Lu, 1998)    |
| No catalyst             | 25°C, CH\(_4\)/CO\(_2\) = 1/2, 4 W, 1 bar, cold corona plasma reactor | 28                  | 20                 | 0.7         | ~                                     | (Aziznia et al., 2012) |
| Ni/\(\gamma\)-Al\(_2\)O\(_3\)| 25°C, CH\(_4\)/CO\(_2\) = 1/2, 4 W, 1 bar, cold corona plasma reactor | 36                  | 23                 | 0.6         | ~                                     | (Aziznia et al., 2012) |
| No catalyst             | 70°C, CH\(_4\)/CO\(_2\) = 1/1, 19.5 kv, 4 KHz, 5.7 mJ, 23 W, 1 bar, pulsed plasma reactor | 61                  | 50                 | 1.3         | ~                                     | (Ghorbanzadeh et al., 2009) |
| Ni/Al\(_2\)O\(_3\)| 160°C, CH\(_4\)/CO\(_2\) = 1/0.67, 16 W, coaxial dielectric barrier discharge plasma reactor | 27                  | 20                 | 2.0         | 0.029                                 | (Zeng et al., 2018)    |
| Ni/\(\gamma\)-Al\(_2\)O\(_3\)| ~230°C, CH\(_4\)/CO\(_2\) = 1/1, 50 W, coaxial dielectric barrier discharge plasma reactor | 56                  | 30                 | –           | ~[1.7E-3, 6.4E-3]                      | (Tu and Whitehead, 2012) |
| Pd/Al\(_2\)O\(_3\)| 240°C, CH\(_4\)/CO\(_2\) = 1/2, 5.7–6 kHz, 1.2 bar, | 51                  | 28                 | –           | –                                     | (Sentek et al., 2010)  |
| Catalyst | Reaction Conditions | Activity | Selectivity | Ref. |
|----------|---------------------|----------|-------------|------|
| Ru/ZrO₂-La₂O₅ | 450°C, CH₄/CO₂/H₂O = 1/1/0.5, 1 bar, fixed-bed reactor | ~15 | ~3 | ~2.5 | – | (Soria et al., 2011) |
Table S3. Properties of Some Other Catalysts Used in Biogas Reforming. (Related to Table 7)

| Catalyst                        | BET surface area (m²/g) | Pore volume (cm³/g) | Pore size (Å) | Basic sites (mmol/g) | Metal dispersion (%) | Ref.                                                |
|---------------------------------|-------------------------|---------------------|---------------|----------------------|----------------------|-----------------------------------------------------|
| CeO₂                            | –                       | –                   | –             | 0.01                 | –                    | (García-Vargas et al., 2012; Pal et al., 2015)       |
| Ni-La-CeO₂                      | –                       | –                   | –             | 0.03–0.04            | –                    | (Pino et al., 2011)                                 |
| Ni/YSZ-CeO₂                     | 10                      | 0.004               | –             | –                    | –                    | (Kang et al., 2007)                                 |
| Imperial chemical industries (ICI) | 13                      | 0.08                | –             | –                    | –                    | (Kang et al., 2007)                                 |
| Holder Topsoe (HT)              | 18                      | 0.05                | –             | –                    | –                    | (Kang et al., 2007)                                 |
| Ce-Zr/Al₂O₃                     | 192                     | 0.74                | 151           | –                    | –                    | (Izquierdo et al., 2013)                            |
| Ni-Zr/Al₂O₃                     | 167                     | 0.62                | 146           | –                    | –                    | (Izquierdo et al., 2013)                            |
| NiMoC-Ce                        | 132                     | 0.26                | 73            | –                    | –                    | (Zou et al., 2016)                                  |
| NiMoC-Mg                        | 120                     | 0.22                | 69            | –                    | –                    | (Zou et al., 2016)                                  |
| Ni@SiO₂                         | 68                      | –                   | –             | –                    | –                    | (Majewski and Wood, 2014; Majewski et al., 2013)    |
| Ni/YSZ-O₂                       | 14                      | 0.12                | –             | 0.01                 | 1.6                  | (García-Vargas et al., 2014)                         |
| CeO₂-ZrO₂                       | 178                     | 0.26                | 58            | –                    | –                    | (Singha et al., 2016)                               |
Table S4. Other Information of the Techno-economics Analysis of Biogas Conversion.

(Related to Table 9)

| Product          | Plant life (year) | Cost                                                                 | Ref.                                           |
|------------------|-------------------|----------------------------------------------------------------------|------------------------------------------------|
| Electricity      | –                 | Initial investment cost: $5,462–5,795; Operating and maintenance cost: $755–1,629/year; Annual saving and benefit: $2,024–2,576/year | (Pipatmanomai et al., 2009)                    |
| H2               | –                 | Investment cost in reforming system: $3,142,897                      | (Madeira et al., 2017)                         |
| H2               | 10                | Capital cost: $968,923; Operational cost: $103,438/year             | (Montenegro Camacho et al., 2017)              |
| Methanol         | –                 | Capital cost: $46,000,000                                           | (Hernández and Martín, 2016)                   |
| Methanol         | 20                | Capital cost: $170,000,000; Operational cost: $48,000,000/year      | (Sheets and Shah, 2018)                        |
| Liquid fuel      | 30                | Total permanent investment cost: $42,000 bbl/day                     | (Graciano et al., 2018)                        |
| Liquid fuel      | 15                | Total direct manufacturing cost: $5,720,000; Capital cost: $8,500,000 | (Zhao et al., 2019)                            |
| Liquid fuel      | 20                | Capital cost: $96,520,000; Operational cost: $11,450,000/year       | (Okeke and Mani, 2017)                         |
| Compressed natural gas | 30 | Capital cost: $2,429,000; Operational cost: $192,100/year | (Winslow et al., 2019)                         |
| Wax              | –                 | Capital cost: $3,080,000; Operational cost: $83.21/bbl              | (Herz et al., 2017)                            |
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