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Advances on methane reforming in solid oxide fuel cells

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ABSTRACT

With the demand for anticipated green hydrogen and power production, novel and upgraded catalytic processes are desired for more effective utilization of precious natural resources. Methane steam reforming is an advanced and matured technology for converting methane to hydrogen and syngas. As a renewable energy resource containing a large amount of methane, biogas is a promising fuel for green hydrogen production. Because of the fuel flexibility and high efficiency relative to alternative technologies, solid oxide fuel cells with internal methane reforming capabilities may become an economically viable technology for hydrogen and power generation. A renewed interest in the flexible application of biogas in solid oxide fuel cells for the co-generation of green hydrogen and power has emerged recently, driven by the spectacular advances in fuel cell technology. However, the methane reforming process suffers from inaccurate or unprecise descriptions. Knowledge of the factors influencing the reforming reaction rate on the novel and improved reforming anode catalysts in solid oxide fuel cells are still required to design and operate such systems. Therefore, a comprehensive review of recent advances in methane steam reforming provides meaningful insight into technological progress. Herein, major descriptors of the methane steam reforming reaction engineering are reviewed to provide a practical perspective for the direct application of biogas in solid oxide fuel cells, which serves as an alternative sustainable, flexible process for green hydrogen and power co-production. Current advances and challenges are evaluated, and perspectives for future work are discussed.

1. Introduction

Climate change has become a major source of concern for the general public in recent years. Policies such as the Kyoto protocol [1] and the Paris agreement [2] have been put in place to limit the impact of global warming to a maximum temperature change of 2 °C above pre-industrial levels by reducing greenhouse gas (GHG) emissions. About 19% of the world's energy is currently supplied by non-fossil resources, such as biomass [3–5], biofuels, nuclear, solar photovoltaic, hydropower, geothermal, or wind energy. Fortunately, the supply of renewable energy resources has steadily increased thanks to the improved energy processing technologies and strict policies for greenhouse gas emissions [6–8]. Predictions indicate that by 2050, non-fossil fuels will make up 46% of the world's energy source.

Furthermore, there is a great concern about energy security because global oil production will not meet the demand within the next 10–20 years [6,9,10]. As a result, the application of renewable and sustainable alternative sources is under development to mitigate global energy and climate concerns. For example, many states in the USA have a long-term goal of achieving 100% renewable electricity-supported grids by 2050. In Australia, New South Wales (NSW) also has a long-term plan to achieve net-zero GHG emissions by 2050 [4,11]. However, the uptake of renewable energy is constrained by its stability and sustainability of supply, adequate distribution, and availability when required.

As a renewable energy carrier, hydrogen (H_2) has the potential to play a significant role in sustainable energy systems [12,13]. H_2 generated from biomass is considered green H_2 as the CO_2 released during the H_2 generation process is mainly compensated by the amount

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of CO₂ absorbed while biomass grows [5,14]. A study conducted by the European Fuel Cells and Hydrogen Joint Undertaking (FCH-JU) on green H₂ production has analyzed the possible pathways to produce H₂ from renewable energy sources. H2 production via biomass gasification and biogas reforming is a promising method [15]. Furthermore, as biogas naturally contains a large amount of CH4, it is a good fuel option for SOFCs where CH4 can be internally reformed on the anode, thus making direct use of the heat and H_2O produced internally [16–20]. Integrating biomass processing technologies, such as gasification, gas cleaning, and upgrading processes, with SOFC technology, can provide more efficient design and simple operation conditions for power and H2 production [5,21-24]. The direct electrochemical oxidation of CH₄ is substantially slower than that of H2 and CO. CH4 is assumed to be reformed to syngas before electrochemical oxidation. Direct internal reforming (DIR) of methane (CH₄) in SOFCs can achieve high electrical efficiencies and even higher efficiencies when applied for combined heat and power (CHP) [25-28], combined cooling, heating, and power (CCHP) [29–32] and combined heat, H_2 , and power (CHHP) [7,33]. This review will not expand on these topics, and the interested readers can refer to the studies mentioned above for more details.

The SOFC is a fuel-flexible electrochemical device that can utilize various fuels to co-produce H_2 and power under proper operation conditions [34,35]. The typical lifetime of state-of-the-art stationary SOFCs is 40,000 h. A single SOFC consists of porous electrodes, a non-porous electrolyte, and interconnects. These can be repeated in a tubular or planar pattern to create a SOFC stack. A schematic illustration of the SOFC's working principle is shown in Fig. 1. The electrochemical reactions are relatively fast if the electrode materials are porous, electronically conductive, and catalytically active. Air/oxygen (O_2) is supplied to the cathode, where O_2 is reduced into oxide ions (O^{2-}) by gaining electrons at the triple phase boundary (TPB) on the cathode side in R1:

$$O_2 + 4e^- = 2O^{2-} \tag{R1}$$

The cathode must reduce O_2 and have sufficient conductivity to provide electrons for this reaction. The readers can refer to the review reported by Sun et al. [36] for more detailed information on the requirement and development of the cathode materials. O^{2-} then travels through the electrolyte towards the anode at the TPB, where they oxide the fuel, as shown in Fig. 1. In case H_2 , the overall electrochemical oxidation reaction can be written as R2:

$$H_2 + O^{2-} \to H_2O + 2e^-$$
 (R2)

SOFCs have been thoroughly researched throughout the literature for all ranges of parameters in experimental studies and computational investigations [22,37,38]. In principle, solid oxide fuel cells (SOFCs) can run on any fuel that can react with oxide ions coming through the electrolyte from the cathode to the anode. Therefore, it is regarded as

one of the most promising candidates for alternative power generation technology, with high energy efficiency and flexibility as a power and H₂ co-producing device. While current SOFCs have demonstrated excellent performance when fuelled with various hydrocarbons, including biogas, significant challenges remain for practical applications [39]. For example, SOFCs produce power and heat by converting the chemical energy of the fuel through MSR, Water Gas Shift (WGS), and electrochemical (half) reactions. Coupling the endothermic reforming and exothermic electrochemical reaction in a single device may lead to local hot and cold spots, inducing unacceptable thermal stresses on the brittle ceramic materials [40]. In addition, several undesirable side reactions may form solid carbon deposits, blocking and damaging the porous electrodes [41]. However, an excessive supply of a reforming agent can suppress the formation of carbon or even remove it but might also oxidize the Ni catalyst in the fuel electrode [42]. Finally, it is difficult to optimize the catalyst for the different internal reactions simultaneously, compromising overall performance [43]. Therefore, a comprehensive understanding of the internal reforming process and its kinetics is indispensable.

Direct CH₄ oxidation in SOFCs is difficult to achieve electrochemically and is more likely to occur as a multi-step process. Therefore, Gur's in-depth review is recommended for a detailed discussion on the kinetics of direct CH₄ oxidation [21]. The effects of material, temperature, current, fuel composition, flow rate, and other experimental parameters on SOFC performance and efficiency have been extensively studied [44–51]. Since the potential utilization of renewable hydrocarbon fuels is a crucial advantage of SOFCs, studies of CH₄ reforming on the existing application of biogas in SOFCs for H2 and power production in various simulations and experiments are very important for the development of such systems. Therefore, the CH₄ reforming in SOFCs and its applications for power and H2 co-production is reviewed comprehensively in this review. The various CH4 reforming kinetic models used in the modelling studies of such systems have been categorized and discussed in detail based on the deviation in complexity and reported assumptions of the models. In summary, this review provides an in-depth examination of the process of CH₄ reformation in SOFCs, providing insight into the potential of using CH₄ derived from a variety of sustainable sources to co-produce power and H2 in SOFCs. Further research in this area is identified to accelerate the implementation of green H2 production using biogas-fed SOFCs and provide significant benefits for climate change and energy security.

2. Modelling studies of methane reforming in SOFCs

CH₄ reforming chemistry is rich and includes various reactions, depending on the available agent, H₂O, O₂, and CO₂ or their mixture. Methane steam reforming (MSR) is the extensively utilised and well-developed of these reforming processes because steam reforming of

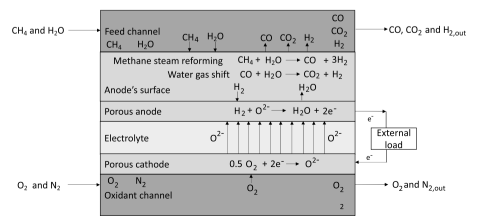


Fig. 1. The methane steam reforming and electrochemical reactions on SOFC anode.

natural gas in centralized plants produces approximately 75% of the H_2 produced globally [16]. As shown in Fig. 1, MSR is a catalytic process expressed in R3 ($H_{298} = +206 \text{ kJ/mol}$) [52]:

$$CH_4 + H_2O \rightarrow CO + 3H_2$$
 (R3)

The CO, besides electrochemical oxidation, is also converted to CO_2 through R4, the water-gas shift (WGS) reaction ($\Delta H_{298} = -41$ kJ/mol) [53], thus maximizing the production of H_2 :

$$CO + H_2O \leftrightarrow CO_2 + H_2$$
 (R4)

In dry methane reforming (DMR), CO_2 is used as an oxidant in place of H_2O ($\Delta H_{298} = +247$ kJ/mol) shown in R5 [54]:

$$CH_4 + CO_2 \rightarrow 2CO + 2H_2 \tag{R5}$$

Interest in this technique is mainly driven by the possibility of capturing and reusing both CH_4 and CO_2 . However, besides the high energy duty, another critical issue is the requirement for a nearly pure source of CO_2 as feedstock.

Partial oxidation of Methane (POM) is an exothermal reaction ($\Delta H_{298} = -38 \text{ kJ/mol}$), which proceeds in R6 [39]:

$$CH_4 + \frac{1}{2}O_2 \rightarrow CO + 2H_2$$
 (R6)

Auto thermal reforming is a hybrid process combining POM with conventional MSR. As the process does not require an external heat source, auto thermal reforming can increase the thermal conversion efficiency of H_2 production and reduce its operational costs. Depending on the purity requirement of the application, syngas production is usually followed by a further purification step. The latter can be achieved using conventional (pressure swing adsorption and cryogenic distillation) or unconventional (membrane separation) technologies [55].

SOFCs are the most promising fuel cell type for CH_4 reforming, opening new avenues for developing green, energy-efficient, and cost-effective H_2 and power co-production technology. One advantage is the fuel flexibility. Since the early 2000s, researchers have been studying the operation of SOFCs on a variety of sustainable fuels, including natural gas, biogas, syngas generated from biomass gasification, and distillate fuel reforming [10,35,42,56–59]. Using different hydrocarbon fuels directly would lower fuel prices and accelerate SOFC adoption, especially for high-value applications [37,39,60–62].

However, fuelling SOFCs with hydrocarbon fuels is inherently complex due to physiochemical, thermal, and transport processes. Given the possibility of competing or co-existing multiple reforming reactions at the SOFC anode, studying the mechanism of CH4 reforming at SOFC operating conditions is a logical first step toward achieving direct CH₄ application. Thus, modelling studies are advantageous for forecasting the operating safety restrictions and designing SOFCs suited for various fuels before actual testing [63-66]. The equilibrium is valuable for anticipating the safety of cells and stacks and the possibility of fuelling them with practical fuels which contain contaminants such as HCl and H₂S [67,68]. System-level modelling can generate the data necessary to develop a holistic view of the system, thus studying its operational regimes, including safety, reliability, and performance [26,57]. The computational fluid dynamic (CFD) simulation is another tool to examine the distribution of mechanical, chemical, and thermal parameters within the system that helps in optimizing the component and system designs [69,70].

There are multiple commercial tools to simulate the SOFC operations, including Aspen Hysys® (Aspen Tech), Aspen Plus® (Aspen Tech), ChemCAD® (Chemstations), COCO/COFE® (AmsterCHEM), Cycle-Tempo (Asimptote), DWSIM® (Open Source CAPE-OPEN process simulator), gPROMS® (Siemens Process Systems Engineering), Pro-SimPlus (ProSim), SimSci PRO/II® (SimSci/Invensys), UniSim® (Honeywell), Comsol ®, and Ansys ®. These software packages can calculate the material and energy balances, the equipment size, and the cost of several chemical processes, including fermentation, distillation,

combustion, gasification, separation, filtration (microfiltration, ultrafiltration, and reverse osmosis), chemical extraction, absorption, adsorption and crystallization. They can also increase productivity and solve chemical process models, modelling, evaluating, and optimizing SOFC systems in a user-friendly environment [63]. These programs have been used for simulations with kinetics implemented in zero-dimensional (0D), one–dimensional (1D), and two–dimensional (2D) models to identify profiles of gas concentration, voltage (V), and feed utilization throughout the SOFC anode [71]. However, models using such software need an accurate description of the reforming process of CH₄.

Yi et al. [72] developed a theoretical framework and thermodynamic simulation capability to investigate an integrated SOFC reformer system working on multiple fuels. The results indicated that feeding various fuels in the same system may present major thermal management issues. Proper operation on different fuels may require significant modifications in operating conditions or system design.

Audasso et al. [73] performed 2D simulations of the direct internal reforming process to study performance degradation and optimize SOFC operating conditions. The developed model may simulate overall cell performance and evaluate the local variables that provide theoretical and feasibility studies insights. Moreover, the model is computationally lean and can be integrated into more complex software for system/plant simulations, such as Aspen Plus.

Another 2D model is developed by Xu et al. [74] to address the carbon deposition problem in SOFCs. The electrolyte can help prevent carbon build-up and lower cell performance by transporting O_2 molecules to the fuel side from the cathode. After being validated by experimental data, the model is expanded to a tubular cell for parametric simulations. All porous SOFCs' carbon resistance and electrochemical performance are investigated under various operating conditions with electrolyte microstructure features. All porous SOFCs have good carbon resistance, as shown quantitatively. The fuel composition and flow rate influence the anode surface's electrochemical performance and oxygen-to-carbon ratio. The findings of this study provide a strong foundation for understanding the principles of operation and promising premises of SOFCs operating with hydrocarbon fuels.

A SOFC stack model comprising 100 anode-supported cells cogenerator with biogas was presented by van Herle et al. [75]. A process flow diagram was defined to vary system operation parameters, including fuel composition, reforming process, stack temperature, current, and fuel utilization. This model can also predict the performance of complex fuel sources such as sewage, landfill sites, and large farms. However, the absence of reforming kinetics and electrode diffusion overpotentials are the shortcomings of this model.

SOFC models for the individual cell or stack are developed using a CFD environment, which may forecast cell potential and the spatial distribution of current density, chemical concentrations, and temperature as time functions for various cell geometries and operating conditions [47,77]. CFD models can also evaluate the effects of different operating variables on SOFC performance. Fan et al. [76] investigated the fuel flexibility of an anode-supported intermediate temperature planar SOFC under co-flow operation using a single channel model containing direct internal reforming (DIR). In Fig. 2, the three reactions considered in this modelling work are the MSR, the WGS, and the electrochemical oxidation of H₂. For the safe operation of biogas-fuelled SOFCs, thermodynamic calculations of Ni oxidation and carbon deposition were performed. The SOFC anodes can be protected against carbon deposition by incorporating enough reforming agents into the fuel stream. While kinetics of carbon deposition and reoxidation is vital in utilizing hydrocarbon fuels, a good understanding of CH4 reforming kinetics is the first necessary step. It will benefit in overcoming the operational safety and stability issues. When fuelling a SOFC with CH₄, the reforming kinetics will significantly affect gas and temperature distributions. These parameters largely determine the fuel cell system performance and occurrence of carbon deposition and thermal stresses

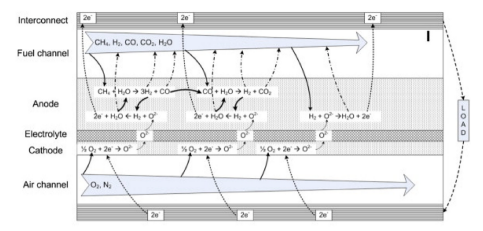


Fig. 2. Schematic representation of reaction and transport processes in the SOFC (Reprinted with the permission from Ref. [76], Copyright (2013), Elsevier).

[44,63]. Because different reforming kinetics have been published in the literature and using them in SOFC simulation may produce varied results. A comparative analysis is required to examine the impact of such kinetic factors on cell performance. The experimental literature on MSR kinetics is thoroughly reviewed and carefully incorporated into this model to determine how internal MSR processes affect cell performance [45,64,78–82]. Based on different reforming kinetics, the modelling results reveal differences in the operational parameters such as current density, temperature, and gas concentration. Calculations of the internal reforming reaction and the possibility of undesirable side reactions are provided, along with recommendations for future theoretical and experimental investigation.

Using the most generally used rate expression described by Achenbach and Riensche [83] was also conducted by Fan et al. [76,77]. However, the Ni concentration (20 wt%) was lower than the 50–70 wt% typically seen in SOFC anodes. In addition, the experimental data collected in this study cannot be directly applicable to other research because of the incompatible operating parameters, such as the anode thickness and feed gas compositions. Another rate expression reported by Xu and Froment [84] was used by Lehnert et al. [85], where first-order dependences of $\mathrm{CH_4}$ and $\mathrm{H_2O}$ concentrations were assumed. However, the latter is rarely observed in laboratory experiments, especially under conditions relevant to direct internal MSR in SOFCs. Therefore, reliable $\mathrm{CH_4}$ reforming kinetic models are needed to describe the reforming process in SOFCs accurately.

Heterogeneous catalysis for MSR over SOFC anode materials must consider the catalyzed reaction and the complexity of the SOFC anode microstructure, contamination of catalytic sites, and adsorption and desorption kinetics. Most research on the MSR reaction kinetics has been conducted during the last decades, particularly on Ni-based catalysts. A considerable variation of MSR kinetics derived from the experimental data was obtained under different testing conditions, including (1) different catalysts; (2) different Ni content/catalytic loadings; (3) various conditions like temperature, steam to carbon ratios, the interaction of electrochemical reactions with the reforming reaction; and (4) different proposed kinetic models and mechanisms. Consequently, there is no single universal model applied in every MSR kinetic study, and the majority of existing data is interpreted through various types of kinetic models. In these SOFC kinetic studies, three types of MSR kinetic models can be distinguished: the multi-step reaction mechanism [78,79,84,86, 87], the surface reaction model assuming a rate-determination step on the surface of a catalyst, and the empirical global gas-phase rate expression [45,78,79,81,83,88-96]. The proposed rate expressions vary among various kinetic models, including First-Order (FO), Power-Law Langmuir-Hinshelwood (LH), muir-Hinshelwood-Hougen-Watson (LH-HW). These are categorized based on the practicality of applying modelling studies to accurately

describe the reforming process and reviewed in the following subsections.

3. Methane steam reforming kinetic models

3.1. Multi-step reaction mechanism

Heterogeneous catalysis is a multistep reaction that involves a sequence of elementary reaction steps [97]. Kinetic studies for MSR were given particular regard, and various kinetic parameters for MSR were reported. Kinetic models can be derived from experimental data, yielding characteristic parameters that describe a reaction. Hecht et al. [98] and Zhu et al. [93] have developed a multistep reaction mechanism for CH₄ reforming on Ni-based SOFC anode. The mechanism comprises $42\,reactions$ involving six gas-phase species and twelve surface adsorbed species. The mechanism encompasses heterogeneous reforming, WGS, and Bouduard reactions [84]. They have also reported a computational model for heterogeneous chemistry and electrochemistry in SOFCs over Ni-based anode materials using the proposed multi-step elementary reactions. A multicomponent transport model based on kinetic theory is used to describe the transport of a gas mixture through a porous matrix of evenly distributed particles. Charge-transfer chemistry is described using a modified Butler-Volmer equation derived from elementary reactions and assuming a single rate-limiting charge transfer process. The underlying theory, on the other hand, is unaffected by geometry. The representation of elementary heterogeneous kinetics in multi-step reaction mechanisms is a key new model feature in SOFCs fed with CH₄-containing fuels. The model considers the effects of channel flow, porous electrode transport, heterogeneous-reforming processes, partial-oxidation chemistry, and electrochemistry. Analyzing oxidative MSR on Ni catalysts helped confirm the heterogeneous chemical reaction process.

Following the investigations by Hetch et al. [98], Janardhanan et al. [99] modelled the internal CH_4 reforming SOFC performance coupled with interactions of transport, heterogeneous catalysis, and electrochemical processes. Experimental data and anticipated cell performance are found in good agreement. At the TPB, the H_2O content significantly impacts the overpotentials and the coverage of surface species. Understanding the underlying chemical processes and optimizing physical operations in any fuel cell can be accomplished using this model.

Wei et al. [96] assessed the reaction mechanisms of CH_4 with CO_2 or H_2O to form syngas and carbon on Nickel–Magnesium Oxide (Ni–MgO) catalysts. The sequence of elementary steps shown in Fig. 3 accounting for CO_2 or H_2O reforming, reforming, CH_4 decomposition, and WGS reactions also provide the steps required to form chemisorbed carbon precursors to carbon filaments. All of the general characteristics of this sequence are in complete agreement with the previously hypothesised

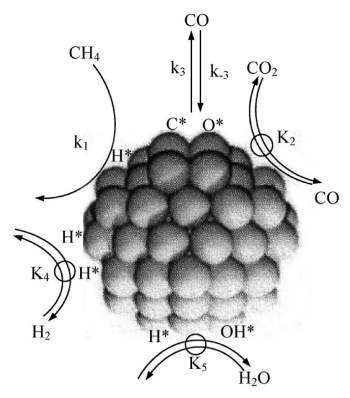


Fig. 3. The sequence of elementary steps for CH₄-reforming and water– gas-shift reactions on Ni-based catalysts (fx1irreversible step, fx1quasi-equilibrated step, fx1 reversible step, k_i is the rate coefficient, and K_i is the equilibrium constant for a given step i) (Reprinted with the permission from Ref. [96], Copyright (2004), Elsevier).

mechanism by Bebelis et al. [92] on supported Ni catalyst. Reaction rates, rate constants and kinetic parameters for the forward CH₄ reactions were identical, indicating that the C–H bond activation is the only kinetically relevant step and reversible. The proposed elementary processes provide a good framework for dealing with the dynamics of carbon synthesis during CH₄ reforming, as evidenced by kinetic and isotopic evidence. The partial pressure ratios of various species in the contacting gas phase are linearly connected to the rate of carbon production and thermodynamic activity of chemisorbed species.

Jones et al. [100] performed first-principles calculations and

experiments on CH₄ reforming over transition metal catalysts. The authors presented a detailed analysis of the reforming process based on experimental observations and proposed nine elementary reactions to represent the MSR reaction. The study considers the adsorption during the reforming process on transition metal surfaces and a series of elementary steps for the MSR reaction. It demonstrates that by integrating scaling principles with thermodynamics and kinetics, one can estimate the catalytic activity of pure metals. A CH₄ dissociation step and a CO formation step are kinetically controlled by the reaction, with the last step dominating at lower temperatures. In a catalyst, CH₄ reforming is a multi-step chemical process that includes gas phase and surface reactions and is highly dependent on operating conditions. The rates of elementary steps are either known with a limited accuracy or are uncertain. As a result, calculating the MSR reaction rate from the rates of elementary steps remains challenging.

3.2. Surface reaction model

Multi-step reaction mechanisms may be reduced to a surface reaction model when a single step is assumed to be rate-determining.

The commonly used LH model shown in Fig. 4(a) represents a catalyst-based mechanism wherein participating species chemisorb onto a heterogeneous catalyst surface in a monolayer and react while adsorbed [79], reflecting the first two scenarios. The Hougen-Watson theory is required to find the rate-controlling step and realize the entire reaction process. The underlying assumption is that a reactant is adsorbed onto the catalyst's surface and subsequently reacts to form the product. The theory is based on the following assumptions: (1) adsorption occurs exclusively on a solid surface's top monolayer; (2) the energy distribution on the surface is uniform; and (3) the adsorbed species do not react with one another.

The surface reaction can be either a single site or dual-site. The site where the reactant is adsorbed in a single-site mechanism participates in the reaction. For example, assuming dissociative adsorption of CH_4 and H_2O on a singular or identical site [101], as shown in R7-18:

$$CH_4 + 2X \leftrightarrow CH_3 - X + H - X$$
 (R7)

$$CH_3 - X + X \leftrightarrow CH_2 - X + H - X \tag{R8}$$

$$CH_2 - X + X \leftrightarrow CH - X + H - X \tag{R9}$$

$$H_2O + 2X \leftrightarrow OH - X + H - X \tag{R10} \label{eq:R10}$$

$$OH - X + X \leftrightarrow O - X + H - X \tag{R11}$$

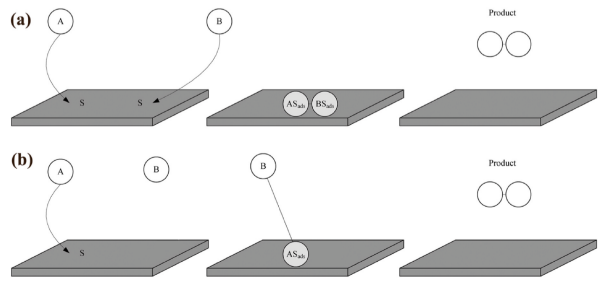


Fig. 4. (a) Langmuir-Hinshelwood kinetic model diagram; (b) Eley-Rideal kinetic model diagram.

$$CH - X + O - X \leftrightarrow CHO - X + H - X$$
 (R12)

$$CHO - X + X \leftrightarrow CO - X + H - X \tag{R13}$$

$$CHO - X + O - X \leftrightarrow CO_2 - X + H - X \tag{R14}$$

$$CO - X \leftrightarrow CO + X$$
 (R15)

$$CO_2 - X \leftrightarrow CO_2 + X$$
 (R16)

$$2H - X \leftrightarrow H_2 - X + X \tag{R17}$$

$$H_2 - X \leftrightarrow H_2 + X$$
 (R18)

Or assuming preferential dissociative CH_4 and H_2O adsorption on two active sites X_1 and X_2 [101], called a "double-site mechanism", as shown in R19-31:

$$CH_4 + 2X_1 \leftrightarrow CH_3 - X_1 + H - X_1$$
 (R19)

$$CH_3 - X_1 + X_1 \leftrightarrow CH_2 - X_1 + H - X_1$$
 (R20)

$$H_2O + 2X_2 \leftrightarrow OH - X_2 + H - X_2 \tag{R21}$$

$$OH - X_2 + X_1 \leftrightarrow O - X_1 + H - X_2$$
 (R22)

$$CH_2 - X_1 + O - X_1 \leftrightarrow CHO - X_1 + H - X_1 \tag{R23}$$

$$CHO - X_1 + X_1 \leftrightarrow CO - X_1 + H - X_1$$
 (R24)

$$CHO - X_1 + O - X_1 \leftrightarrow CO_2 - X_1 + H - X_1$$
 (R25)

$$CO - X_1 \leftrightarrow CO + X_1$$
 (R26)

$$CO_2 - X_1 \leftrightarrow CO_2 + X_1$$
 (R27)

$$2H - X_1 \leftrightarrow H_2 - X_1 + X_1$$
 (R28)

$$2H - X_2 \leftrightarrow H_2 - X_2 + X_2$$
 (R29)

$$H_2 - X_1 \leftrightarrow H_2 + X_1 \tag{R30}$$

$$H_2 - X_2 \leftrightarrow H_2 + X_2 \tag{R31}$$

A reaction between an adsorbed molecule and a gas phase molecule serves as a third mechanism. The Eley–Rideal (ER) mechanism explains a reaction between a chemisorbed reactant and another non-chemisorbed reactant, as shown in Fig. 4(b). Single-layer adsorption, no diffusion limit, and only the reactants affect the reaction rate are some of the approximations. Hofmann et al. [102] analyzed the heterogeneous reaction mechanism published by Hecht et al. [98] into a simplified model. The CH_4 reforming is represented using a global kinetic approach, and the WGS reaction is assumed to be in equilibrium. The catalytic surface coverage can be calculated using the model.

Individual steps for the adsorption or desorption of various species during CH_4 reforming on different Ni surfaces were investigated experimentally and numerically by Delgado et al. [103]. The main pathways for CH_4 conversion processes on the Ni catalyst for H_2 and CO productions are shown in Fig. 5. A model consisting of 52 reactions with six gas-phase species and 14 surface species involving adsorption and desorption steps of all reactants, products and surface reaction steps was proposed. The availability of adsorbed atomic oxygen O(s), produced via dissociative adsorption of oxygen-containing species, plays a vital role in determining the reaction rates supported by the DFT studies [104–107]. A detailed reaction mechanism for CH_4 reforming was developed and evaluated, making it possible to predict the product distribution for methane conversion in the reactor. The developed simulation tools simulate the chemical species profiles and the surface coverage within the reactor.

The heterogeneous reactions have multiple elementary steps that compete with one another. According to this theory, any of the three groups can determine the reaction rate: (1) kinetic group, (2) driving force group, and (3) adsorption group [97]. For a hetero-catalytic reaction, a general form of rate expression could be expressed in Eq (1):

$$r = \frac{(kinetic factor) \cdot (driving force)}{(adsorption isotherm)} \tag{1}$$

In the MSR reaction, assuming a single layer surface hetero-catalytic reaction, the general adsorption isotherm is calculated in Eq (2):

adsorption is other
$$m = (S_0 + K_{CO}p_{CO} + K_{H_2}p_{H_2} + K_{CH_4}p_{CH_4} + K_{H_2O}p_{H_2O})^n$$
(2)

Each term is proportional to the surface coverage by the respective species, scaled such that S_0 is proportional to the vacant surface. n is the

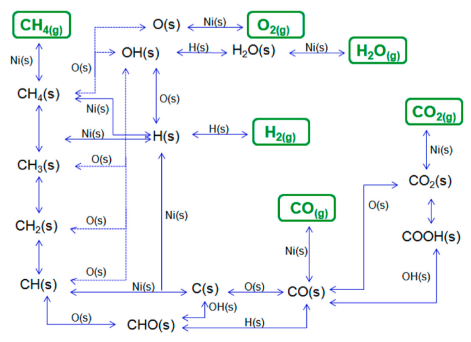


Fig. 5. Reaction pathways for methane oxidation and reforming on the nickel-based catalyst (Reprinted with the permission from Ref. [103], Copyright (2015), MDPI).

number of sites involved in the molecular reaction. The assumption of the bimolecular rate-determining step as the kinetic term can be combined with the Langmuir isotherms to describe the adsorption to the catalyst surface. The temperature dependence of the adsorption coefficient K_j typically follows the van't Hoff equation. The driving force of the reaction is calculated based on the deviation from reaction equilibrium. It can be calculated via the reaction quotient and a temperature-dependent equilibrium constant, where the reaction quotient has been expanded to the MSR reaction. The driving force is calculated in Eq (3) [108]:

driving force =
$$1 - \frac{p_{CO} \cdot p_{H_2}^3}{K_{eq,wgs} \cdot p_{CH_4} \cdot p_{H_2O}}$$
 (3)

The equilibrium constant for the WGS reaction ($K_{eq,wgs}$) is a function of temperature, calculated using empirical equations [84].

The general kinetic factor is calculated in Eq (4):

$$kinetic factor = k \cdot p_{CH_A}^a \cdot p_{H_2O}^b \cdot p_{CO}^c \cdot p_{CO}^d$$
(4)

Experiments were carried out to evaluate the performance of MSR activity over a Nickel-Magnesium Aluminate (Ni-MgAl₂O₄) catalyst, and an effective kinetic model was developed based on the kinetic model proposed by Xu and Froment [84], in which an intrinsic expression for the reforming reaction rate was proposed. The model has been widely accepted as a reliable approximation of MSR kinetics. However, compared to commercial SOFC anodes, which typically have Ni contents ranging from 50 to 70 wt %, this one had a lower Ni concentration (20 wt%). Lehnert et al. [85] applied the MSR kinetic parameters obtained by Xu and Froment [84] to an industrial catalyst. Elnashaie et al. [109] employed the LH reforming rate expression to assess the impact of non-monotonic kinetics on the performance of steam reformers, which explains the discrepancies between the rate expressions found in the literature. Experiments were also carried out by Hou et al. [87] to study the kinetics of the MSR over a commercial Nickel- α Aluminum Oxide (Ni-αAl₂O₃) catalyst. The reforming reaction rate increases in proportion to the CH₄ partial pressure at low product concentrations. The surface reaction between adsorbed species was identified as the rate-controlling step. The intrinsic rate expressions were obtained using the LH-HW method and Freundlich's adsorption theory. The simulation determined a satisfactory model of intrinsic kinetics. Klein et al. [110] modelled the direct internal and gradual internal reforming SOFCs using the kinetic parameters reported by Hou et al. [87]. The kinetic data were obtained from testing a Ni-αAl₂O₃ catalyst, different from the commonly used anode materials.

Dicks et al. [86] studied the MSR reaction on Nickel-Yttria-Stabilized Zirconia (Ni-YSZ) in a tubular plug flow differential reactor. They developed an improved MSR rate expression assuming LH adsorption isotherms. The reaction showed first-order dependence on CH_4 , weak positive dependence on H_2 , and strong negative dependence on H_2O . Because reaction orders in H_2 and H_2O were allowed to vary for different temperatures and gas compositions, the kinetics were convoluted.

Bebelis et al. [92] investigated the MSR catalytic activity and kinetics of a Ni-YSZ anode. The reaction followed an LH kinetic pattern, indicating that CH_4 and H_2O adsorption on the catalytic sites was competitive. The apparent activation energy is greatly influenced by two rate-determining steps: the activated adsorption of CH_4 for the formation of active species and the surface reaction between the adsorbed activated carbon and O_2 for the production of CO.

Additional factors need to be considered due to the complicated nature of the adsorption-desorption mechanism. Different anode manufacturing processes result in varying catalyst availabilities for the adsorbed species. Combined with diffusion and heat/mass transfer limitations, this can result in deviations from the actual results. These problems may have a more significant impact on reforming behaviour than the kinetics of the surface reaction itself. To combat this issue, Wang et al. [111] implemented an 'effective factor' that can be

experimentally calculated and acts as a rate coefficient, allowing greater model adherence to the physical behaviour, although at the cost of computing time. Furthermore, the need for additional kinetic parameters is still not thoroughly demonstrated in the literature.

Thallam Thattai et al. [79] emphasised the need for simply applicable global kinetic models for SOFC systems and a lack of research on MSR global and elementary kinetic mechanisms. They observed the limitations in using previously proposed rate expressions for Ni-based catalyst beds to analyse MSR kinetics of SOFC cermet anodes. Many PL and LH kinetic models for MSR on Ni-GDC anodes were developed and predict significantly different local MSR reaction rates and distributions of species concentrations, highlighting the need for further investigation. The following subsection examines a practical and balanced kinetic treatment based on simplifications and approximations.

3.3. Gas-phase reaction model

The PL kinetic model is the most simplified and commonly used reaction rate expression in modelling studies. This model simplifies the reforming reaction kinetics by assuming a global singular reaction step for a given reaction, determined via a set of experimentally determined variables. A fundamental principle of this model is that the rate of every chemical reaction is proportional to the masses of the reacting substances, as shown in Eq (5):

$$r = k \cdot p_{CH_4}^a \cdot p_{H_2O}^b \tag{5}$$

where p_{CH_4} and p_{H_2O} are the partial pressures of CH₄ and H₂O, and a and b are the reaction orders of CH₄ and H₂O, respectively. According to the previously published data, a typical range for these reaction orders is -2.0 to 2.0, according to the previously published data [44]. Reaction orders commonly resemble first-order for p_{CH_4} , and zero-order for p_{H_2O} .

Achenbach and Riensche [83] report a PL rate expression that has been extensively used in SOFC modelling studies. Ahmed and Foger [91] studied the MSR catalytic activity of basic Ni-YSZ and modified anode materials. A lower $\rm H_2O$ to $\rm CH_4$ ratio was required to avoid carbon deposition. They used a PL kinetic model, and the orders for $\rm CH_4$ and $\rm H_2O$ were higher on the modified anode. In addition, the activation energy was greater on the modified anode, showing that the active catalytic sites in these two anodes were distinct.

MSR on Ni–MgO catalysts at low pressure was investigated by Wei et al. [96]. A PL rate expression was used in this study with a lower temperature range. The reaction rate was proportional to p_{CH_4} , and independent of p_{H_2O} . In this study, a comparison of the reactivity of Ni and noble metal catalysts for CH₄ reforming was made using direct measurements of C–H bond activation rates. The activation energy was similar to that of the CH₄ decomposition, indicating the activation of the C–H bond is the rate-limiting step.

Experimental study of reforming kinetics and the influence of operation conditions on it are essential for an accurate prediction of the performance of a direct internal reforming SOFC. Internal reforming kinetic parameters have been reported by Mogensen et al. [90] on a complete cell with a Ni-YSZ anode under closed-circuit operations. The rate expression was proposed in a PL type with an activation energy of 166.1 kJ/mol. The reaction rate has a dependency on only p_{CH_4} , with a reaction order of 0.7. MSR over a Ni-YSZ anode in stack configuration was also studied by Mogensen et al. [82]. It was discovered that the rates could be well matched to a PL rate expression when the measurements were carried out between 600 $^{\circ}\text{C}$ and 800 $^{\circ}\text{C}$ under different pressures. According to the model, the CH₄ conversion in a stack configuration could be predicted from the inherent dynamics of the anode support materials, and the predictions were consistent with the experimental results in most cases. Because of its simplicity, this model is well suited for simple SOFC stack models, such as those used for flow sheeting applications such as Aspen Plus®.

Sciazko et al. [94] presented a novel approach to the experimental investigation of MSR kinetics utilizing the orthogonal least-squares method. The results were obtained using a plug flow tubular reactor with fine Ni-YSZ catalyst particles as a catalyst. The suggested algorithm is easily adaptable to mathematical modelling of the fuel reforming process on various catalysts and under different experimental conditions. Furthermore, the difference between the published inconsistent data about the reaction kinetic parameters can be clarified using this approach. Fan et al. [44,45] investigated MSR reaction kinetics in complete SOFC with Ni-YSZ and Ni-GDC anodes using this method under varying operating conditions. A PL rate expression was used in both studies. The CH₄ conversion reduces as the temperature decreases, consistent with earlier research. However, the MSR reaction rate increases slightly when a current is drawn. The effect of H₂O concentration on the reaction rate becomes more negative, likely due to the adsorbed O2/H2O molecules/ions blocking reaction sites or the local oxidation of Ni. Thallam Thattai et al. [79] extended the study and reported the modelling results of both PL and LH kinetic models derived from their previous experimental studies [45,78,79]. Both models have successfully predicted the spatial distribution of the reforming reaction.

PL type rate expression requires experimentally determined values of the pre-exponential factor, the activation energy, and the reaction orders to predict the reaction rates for given conditions. These parameters can be readily applied to systems with variant temperatures and inlet gaseous compositions when these parameters are obtained. Several studies, such as those conducted by Achenbach et al. [83] and Belyaev et al. [112] have assumed FO kinetics. This low-order model prioritized the computational time, which was reduced significantly through the first-order assumption, allowing for the development of a real-time feedback control system. However, this simplification sometimes is contrary to multiple experimentally produced data sources, showing reaction orders for $\rm CH_4$ and $\rm H_2O$ vary across numerous experimental conditions.

3.4. Summary and comparison of methane steam reforming kinetic models

MSR kinetics on SOFC anodes have been widely studied. A large variety of reaction rate expressions has been proposed in different studies. The various reaction rate expressions may be selected and used in different modelling studies based on the anode materials and operation conditions. The previously reported PL and LH kinetics for MSR on Ni-based materials are summarized in Table 1. The WGS kinetic models over various catalysts are summarized in Table 2.

A vast majority of these studies used Ni-YSZ cermets. Kinetic data have also been reported on Ni-GDC anode materials. For the MSR kinetic studies, some researchers used Ni-based catalysts, such as Ni-MgAl $_2$ O $_4$ cermet [91] and Ni- α Al $_2$ O $_3$ catalyst [135], for the MSR kinetic studies. As reactions occur on the anode, which is perhaps the most critical factor for the operational performance of SOFCs. SOFC anode materials should ideally be chemically stable, have sufficient mechanical strength, and possess high conductivity and electro-catalytic activity. A series of MSR catalysts with increased activity and stability have been developed. They can be employed as anode catalysts in the SOFC, allowing for relatively inexpensive, safe, and readily available fuel cell development [37,46,128]. Many detailed, comprehensive overviews of those materials can be found in other reviews [128–135].

The reported values for the reforming activation energy differ from as little as 18 kJ/mol up to 240 kJ/mol [44,45,79,113]. Ni contents typically range from 50 to 70 wt%, close to the typical Ni content in SOFC anode materials. Few studies have reported the MSR kinetic parameters on materials with low Ni content [87,96,112,119]. Lee et al. [116] and Minette. et al. [81] reported MSR kinetics on high Ni content materials with a significant activation energy difference. Based on the catalyst materials used, these experimental studies can be broadly grouped into two categories: (1) Ni-based catalytic materials such as

catalyst pellets [83,84,86,92,100,116,136]; (2) active SOFC anodes, such as the porous catalyst layer on the electrolyte support, under conditions with or without producing a current [44,45,78,79,90,91,113, 137–140].

The experimentally derived reforming kinetics mainly were obtained within the same temperature range as the SOFC operating temperature, from 700 $^{\circ}$ C to 900 $^{\circ}$ C as shown in Table 1. Few studies have reported the reforming kinetic parameters below 600 $^{\circ}$ C. Xu et al. [84] have studied the reforming kinetics from 400 $^{\circ}$ C to 800 $^{\circ}$ C, and Hou et al. [87] have tested the catalytic activity from 450 $^{\circ}$ C to 550 $^{\circ}$ C. Most research was carried out on Ni-based catalytic materials operated within a safe range of steam to carbon ratios. The trials carried out in these studies were designed to avoid the formation of carbon deposits. With the elimination of carbon deposition during the reforming reaction, it may be possible to understand better how different operating conditions affect the reforming kinetics in SOFCs.

The overall reaction order in the PL kinetic model also varies significantly from 0.25 to 0.8. The LH exponents mainly were found to be 2. These variations in the reaction orders may be due to the various test materials, different steam to carbon ratios, and temperatures. Even with a similar Ni content in the anode materials, dopants and modifications can significantly impact the kinetic activity [79,95,113]. Mogensen et al. [82,90,141] investigated the kinetics of the internal MSR reaction on Ni-YSZ within a SOFC stack and externally via crushed Ni-YSZ pellets using a packed bed reactor. An expression for the rate of $\rm H_2$ generation was derived via the PL kinetic model. The reaction orders were fixed to 0.9, $\rm -0.2$, and 0.2 for CH₄, H₂, and CO₂, respectively. The activation energy was 185 kJ/mol. However, it was noted that this model on the packed bed had a mean absolute deviation of 43.7%.

Six CH₄ reforming reaction rates: four PL-type rate expressions and two LH-type rate expressions derived on different functional Ni-YSZ anodes versus temperature, the p_{CH_4} and p_{H_2O} are selected for the comparison, and the discussions are as follows.

The influence of temperature on the MSR reaction rates, converted to the standard surface reaction rate over Ni-YSZ anodes, was plotted in Fig. 6. The calculated reaction rates using the selected kinetic parameters increase exponentially along with temperature; however, they vary significantly given the same partial pressures of different species and temperatures. The variations may result from the material properties, such as the surface area, the microstructure of the anode materials, and operational conditions such as the temperature, the flow rate, and the steam to carbon ratio, indicating the challenge of developing a universal reforming rate expression.

The influence of p_{CH_4} on the MSR reaction rate over Ni-YSZ anodes is plotted and shown in Fig. 7 (a). Most of the calculated reaction rates increase linearly with CH₄ because most reaction orders for CH₄ are either one or very close to one. The influence of p_{H_2O} on the MSR reaction rate over various Ni-YSZ anodes were plotted and shown in Fig. 7 (b). Most of the calculated reaction rates slightly decrease with p_{H_2O} because many reaction orders for H₂O are very close to zero or slightly negative. However, the absolute reaction rates predicted by the previously reported kinetic parameters vary significantly for similar temperatures, flow rates, and species partial pressures. There are numerous reasons for these variations. The surface area of the anode materials, for example, and the microstructure of the anode materials are all affected by material properties, as are the temperature and steam to carbon ratio. This indicates the need for a more consistent reforming reaction rate expression.

4. Application of biogas in SOFCs

SOFC is a promising alternative technology that can convert CH_4 to power with potentially high efficiency. In addition, CO_2 can be separated relatively easily from the flue gas as air and fuel are not mixed in the SOFC. Therefore, biogas is a promising fuel for SOFCs. The CH_4 content typically ranges from 45 mol% to 75 mol%, most of the

Table 1MSR kinetics of Ni-based materials.

				Ni	SC	Current (A/ m ²)	E _a kJ/mol	Kinetic parameters
113]	Ni-YSZ-CeO ₂ anode	$r_{CH_4} = \frac{K_1 K_2 K_3 p_{CH_4} p_{H_2O}}{\left(1 + K_2 p_{CH_4} + K_2 p_{H_2O}\right)^2}$	973–1273	50	2–7	6000	≥ 18	-
114]	Ni–ZrO ₂ cermet	$r_{CH_4} = k_0 \exp\left(-\frac{E_a}{RT}\right) \left(1 - \frac{p_{CO}p_{H_2}^3}{p_{CH_4}p_{H_2O}}\right) p_{CH_4}$	973–1313	20	2.6–8	_	82	-
115]	Ni cermet	$r_{CH} = k(p_{CH})^{1.5}$	1223	_	3	_		$k = 2.4e - 3 mol s^{-1} atm^{-1.25}$
116]	Ni-YSZ cermet	$r_{CH_4} = k_0 \exp\left(-\frac{E_a}{RT}\right) p_{CH_4} (p_{H_2O})^a$	1073–1273	50-80	2–7.4	-	17.8, 23.5	$k_0 = 490 \ 4775 \ mol \ g^{-1}h^{-1}, \ \alpha = -1.28, 1.25$
86]	Ni-YSZ anode	$r_{CH_4} = k \frac{p_{CH_4}}{\left(1 + K_H p_{H_2}^{0.5} + K_S \frac{p_{H_2} o}{p_{H_1}}\right)^n}$	973–1273	55	1.2–7	-	135	$k_0 = 21 \; mol \; bar^{-1} cm^{-2} s^{-1}, \; n = 2$
117]	Ni/GDC, Au–Ni/GDC anodes	$r_{CH_4} = \frac{k_a p_{CH_4}}{1 + k \frac{p_{CH_4}}{p_{H_2O}}}$	1073–1173	57	0.25–1	2300	23–28	$k = 0.22 \sim 0.25, k_a = 15 - 45e - 6$
84]	Ni–MgAl ₂ O ₄ cermet	$r_{CH_4} = rac{k_1}{p_{H_2}^{2.5}} \left(p_{CH_4} p_{H_2O} - rac{p_{CO} p_{H_2}^3}{K_1} \right)$	675–1000	15	3–5	-	240.1	-
109]	Ni foil	$\overline{(1 + K_{CO}p_{CO} + K_{H_2}p_{H_2} + K_{CH_4}p_{CH_4} + K_{H_2}op_{H_2}o)^2}}$ $r = \frac{kp_{CH_4}}{1 + a\frac{p_{H_2}o}{p_{H_2}} + bp_{CO}}$	1073–1173	15	1–3	-	-	$a = 0.2 \sim 0.5, b = 0 \sim 2$
91]	Ni–ZrO ₂ anode	$r_{CH_4} = k_0 \exp\left(-\frac{E_a}{RT}\right) (p_{CH_4})^a (p_{H_2O})^{\beta}$	1127–1180	-	1.53–2.5	-	95 ± 2	$\alpha = 0.85 \pm 0.05, \beta = -0.35 \pm 0.04, k_0 = 8542 mol s^{-1} m^{-2} bar^{-0.5}$
71]	Modified Ni–ZrO ₂ anode	$r_{CH_4} = k_0 \exp\left(-\frac{E_a}{RT}\right) (p_{CH_4})^{\alpha} (p_{H_2O})^{\beta}$	1111–1195	-	1.4–3	-	$\textbf{20,8} \pm \textbf{10}$	$\alpha = 1.4 \pm 0.01, \beta = -0.8 \pm 0.02, k_0 = 3.6e8 mol s^{-1} m^{-2} bar^{-0.6}$
92]	Ni-YSZ cermet film	$r_{CH_4} = k_{ad}p_{CH_4} \left(1 - \frac{k_{ad}}{k_r K_{H_2O}} \frac{p_{CH_4} p_{H_2}}{p_{H_2O}}\right)$	1073–1173	70	0.07–16	-	48.2 ± 2.1	$k = 3.44 \sim 23.5e - 6 \text{ mol kPa}^{-1} s^{-1}$
[89]	Ni-CGO anode	$r_{CH_4} = k_0 \exp\left(-\frac{E_a}{RT}\right) (p_{CH_4})^a (p_{H_2O})^{\beta}$	1073–1223	50	0–3	-	26.3	$k_0 = 4.05e - 5 mol / (s * m^2 * Pa^{1.19}), \alpha = 1.19, \beta = 0$
[89]	Ni-YSZ anode	$r_{CH_4} = k_0 \exp\left(-\frac{E_a}{RT}\right) (p_{CH_4})^{\alpha} (p_{H_2O})^{\beta}$	1073–1223	50	0–3	-	52	$k_0 = 4.05e - 5 mol / (s * m^2 * Pa^{1.19}), \alpha = 1.19, \beta = 0$
118]	Ni-YSZ cermet	$r_{CH_4} = k_0 \exp\left(-\frac{E_a}{RT}\right) (p_{CH_4})^a (p_{H_2O})^b$	873–1023	60	2.5–5	-	117	$k_0 = 41.5 mol g^{-1} s^{-1} bar^{-(a+b)}, a = 0.98, b = -0.09$
118]	Ni-SDC cermet	$r_{CH_4} = k_0 \exp\left(-\frac{E_a}{RT}\right) (p_{CH_4})^a (p_{H_2O})^b$	873–1023	60	2.5–5	-	106	$k_0 = 40 \ mol \ g^{-1} s^{-1} bar^{-(a+b)}, \ a = 0.98, \ b = -0.25$
45]	Ni-GDC anode	$r_{CH_4} = k_0 \exp\left(-\frac{E_a}{RT}\right) (p_{CH_4})^{\alpha_{CH_4}} (p_{H_2O})^{\alpha_{H_2O}}$	973–1023	57	1.5–2.45	600, 1000	63–88	$k_0=1.1\sim18.9~mol~g^{-1}s^{-1}bar^{-(a_{CH_4}+a_{H_2O})},~a_{CH_4}=0.64\sim0.67,~a_{H_2O}-0.04\sim-0.08$
79]	Ni-GDC anode	$r_{CH_4} = k_0 \exp\left(-\frac{E_a}{RT}\right) (p_{CH_4})^{\alpha_{CH_4}} (p_{H_2O})^{\alpha_{H_2O}}$	1043–1103	57	1.07–1.35	500–3000	50	$a_{CH_4} = 0.15$, $a_{H_2O} = 0.1$, at open circuit, and both vary with increasing current
79]	Ni-GDC anode	$r_{CH_4} = k \cdot p_{CH_4} \frac{p_{H_2O}^q}{p_{H_2}^p} \frac{\left(1 - \frac{p_{CO}p_{H_2}^3}{K_1 p_{CH_4} p_{H_2O}}\right)}{\left(1 + K_{CH_4} p_{CH_4} + K_{H_2O} \frac{p_{H_2O}}{p_{H_2}}\right)^2}$	1043–1103	57	1.07–1.35	500–3000	160–220	a=0.007,b=1.210 at open circuit, and both vary with increasing current
78]	Ni-GDC anode	$r_{MSR} = k p_{CH_4}^{\alpha} p_{H_2O}^{\beta} p_{H_2}^{\gamma} \left(1 - \frac{Q_{MSR}}{K_{MSR}} \right)$	973–1048	57	1.5–3	_	173.1	$\alpha = 0.8954, \ \beta = -0.0619, \ \gamma = 0.0693,$ $k_0 = 9.472e8 \ mol \ s^{-1} m^{-2} bar^{(\alpha+\beta+\gamma)}$
78]	Ni-GDC anode	$r_{MSR} = k p_{CH_4} \left(1 - rac{Q_{MSR}}{K_{MSR}} ight)$	973–1048	57	1.5–3	_	190.5	$k_0 = 9.472e8 \ mol \ s^{-1}m^{-2}bar^1$
78]	Ni-GDC anode	$r_{MSR} = \frac{kK_{CH_4}K_{H_2O}p_{CH_4}p_{H_2O}^{0.5}}{\left(1 + K_{CH_c}p_{CH_4}^{0.2} + K_{H_2O}p_{H_2O}^{0.5}\right)^2} \left(1 - \frac{Q_{MSR}}{K_{MSR}}\right)$	973–1048	57	1.5–3	-	207.6	$k_0 = 1.467e10 \ mol \ s^{-1} m^{-2}$
78]	Ni-GDC anode	Conditional of the supplemental of the supplem	973–1048	57	1.5–3	-	158.5	$k_0 = 2.787e7 \ mol \ s^{-1}m^{-2}bar$

(continued on next page)

Table 1 (continued)

Ref	Anode Material	Rate Expression	T (K)	Wt% Ni	SC	Current (A/ m ²)	E _a kJ/mol	Kinetic parameters
		$r_{MSR} = \frac{kp_{CH_4}}{\left(1 + K_{CH_4}p_{CH_4} + \frac{K_{H_2O}p_{H_2O}}{p_{H_5}}\right)^2} \left(1 - \frac{Q_{MSR}}{K_{MSR}}\right)$						
82]	Ni-YSZ anode	$r = k_0 \exp\left(-\frac{E_a}{RT}\right) p_{CH_4}^{\alpha_{CH_4}}$	873–1073	51	1.5–6	250–1000	166.1	$k_o = 1.99 \text{e4 mol s}^{-1} \text{m}^{-2} \text{Pa}^{-0.7}, \alpha_{\text{CH}_4} = 0.56 \sim 0.80,$
119]	Ni–ZrO ₂ –CeO ₂ anode	$r_{CH_4} = k(p_{CH_4})$	1073–1123	5	2–4	-	39	$k=2.6e9\exp\left(-\frac{19500}{T}\right)\mu mol min^{-1}$
37]	Ni–α Al ₂ O ₃ catalyst	$r_{CH_4} = r_1 + r_3 (LH - HW type)$	748–823	15–17	3–7	-	209.2, 109.4	5.922e8, 1.093e3
93]	Ni-based anode	Elementary reaction mechanism	1048-1223	-	1.5–4	0-80000	-	$k = AT^n \exp\left(-\frac{E_a}{RT}\right) \exp\left(\frac{\varepsilon_{CO9s0}\theta_{CO9s0}}{RT}\right)$
94]	Ni-YSZ cermet	$r_{CH_4} = k_0 \exp\left(-\frac{E_a}{RT}\right) (p_{CH_4})^a (p_{H_2O})^b$	823–1023	60	2.5–6	-	$121.3 \pm \\2.8$	$a = 0.88 \pm 0.058, b = 0.083 \pm 0.039,$ $k_0 = (6.472 \pm 1.921)e - 3 mol s^{-1} m^{-2} bar^{0.78}$
95]	Ni-YSZ anode	$r_{CH_A} = k(p_{CH_A})$	873-1073	50	3	_	124	$k_0 = 1.82 \text{ mol } g^{-1}h^{-1}$
96]	Ni–MgO catalyst	$r_{CH_4} = k(p_{CH_4})$	823–1023	7, 15	_	_	102	$k_0 = 1.62 \text{ mot g}^{-1}$ $k_0 = 2.5e5 \text{ s}^{-1} Pa^{-1}$
81]	Ni coating on metal	$r = kp_{CH_4} \left(1 - \frac{p_{CO}p_{H_2}^3}{p_{CH_2}p_{H_2O}K_1}\right)$	723–873	75–85	2.87-5.53	-	226.4	$k_0 = 7.48e12 mol bar^{0.5} kg_{cat}^{-1} s^{-1}$
[45]	Ni-GDC anode	$r_{CH_4} = k_0 \exp\left(-\frac{E_a}{RT}\right) (p_{CH_4})^{a_{CH_4}} (p_{H_2O})^{a_{H_2O}}$	923–1023	57	1.5–2.45	0	88 ± 11	$\alpha_{CH_4}=0.64\pm0.15,\ \alpha_{H_2O}=-0.04\pm0.07,\ k_0=18.9\pm17.6\ mol/(\ bar^{0.60}s^{-1})$ 1.7 g catalyst
[45]	Ni-GDC anode	$r_{CH_4} = k_0 \exp\left(-\frac{E_a}{RT}\right) (p_{CH_4})^{\alpha_{CH_4}} (p_{H_2O})^{\alpha_{H_2O}}$	923–1023	57	1.5–2.45	632	63 ± 8	$a_{CH_4} = 0.70 \pm 0.06, \ a_{H_2O} = -0.04 \pm 0.03, \ k_0 = 1.1 \pm 0.9 \ mol/(\ bar^{0.66} s^{-1})$ 1.7 g catalyst
45]	Ni-GDC anode	$r_{CH_4} = k_0 \exp\left(-\frac{E_a}{RT}\right) (p_{CH_4})^{\alpha_{CH_4}} (p_{H_2O})^{\alpha_{H_2O}}$	923–1023	57	1.5–2.45	1052	66 ± 1	$a_{CH_4} = 0.67 \pm 0.05, \ a_{H_2O} = -0.08 \pm 0.05, \ k_0 = 1.5 \pm 0.2 \ mol/(\ bar^{0.59} s^{-1})$ 1.7 g catalyst
[44]	Ni-YSZ anode	$r_{CH_4} = k_0 \exp\left(-\frac{E_a}{RT}\right) (p_{CH_4})^{\alpha_{CH_4}} (p_{H_2O})^{\alpha_{H_2O}}$	923–1048	57	1.83-5.50	0	59.27	$a_{CH_4} = 0.50$, $a_{H_2O} = -1.19$, $k_0 = 5.38 \ mol/(bar^{0.69} g_{-n}^{-1} min^{-1})$
[44]	Ni-YSZ anode	$r_{CH_4} = k_0 \exp\left(-\frac{E_a}{RT}\right) (p_{CH_4})^{a_{CH_4}} (p_{H_2O})^{a_{H_2O}}$	923–1048	57	1.83-5.50	632	50.35	$a_{CH_4} = 0.56, a_{H_2O} = -0.59,$ $k_0 = 7.43 \text{ mol}/(bar^{0.03}g_{ca}^{-1}\text{min}^{-1})$
44]	Ni-YSZ anode	$r_{CH_4} = k_0 \exp\left(-\frac{E_a}{RT}\right) (p_{CH_4})^{a_{CH_4}} (p_{H_2O})^{a_{H_2O}}$	923–1048	57	1.83-5.50	1052	50.35	$a_{CH_4} = 0.59, a_{H_2O} = -0.49,$ $k_0 = 14.29 \ mol/ (bar^{-0.10} g_{cal}^{-1} min^{-1})$
[44]	Ni-YSZ anode	$r_{CH_4} = k_0 \exp\left(-\frac{E_a}{RT}\right) (p_{CH_4})^{a_{CH_4}} (p_{H_2O})^{a_{H_2O}}$	923–1048	57	1.83-5.50	1052	50.35	$k_0 = 14.29 \text{ mol}/(bar \frac{k_{car} mar}{k_{car}})$ $a_{CH_4} = 0.59, a_{H_2O} = -0.49,$ $k_0 = 14.29 \text{ mol}/(bar^{-0.10}g_{car}^{-1} min^{-1})$
[44]	Ni-YSZ anode	$r_{CH_4} = \frac{k \cdot p_{H_2}^c \cdot p_{H_2O}^b \cdot p_{H_2}^c \cdot \left(1 - \frac{p_{CO} p_{H_2}^3}{K_1 p_{CH_4} p_{H_2O}}\right)}{\left(1 + K_{CH_4} p_{CH_4} + K_{H_2O} p_{H_2O} / p_{H_2}\right)^2}$	923–1048	57	1.83–5.50	0	52.43	$k_0 = 14.25 mol/(bat k_{scat}^2 min^2)$ a = 0.37, b = -1.03, c = -0.38 $k_0 = 1.10 mol/(bat^{1.04} g_{cat}^{-1} min^{-1})$
[44]	Ni-YSZ anode	$k \cdot p_{H_2}^c \cdot p_{H_2O}^b \cdot p_{H_2}^c \cdot \left(1 - \frac{p_{CO}p_{H_2}^3}{K_{PDM}}\right)$	923–1048	57	1.83–5.50	632	41.73	a = 0.45, b = -0.49, c = -0.04 $k_0 = 2.37 \text{ mol/}(bar^{0.08}g_{cal}^{-1}min^{-1})$
[44]	Ni-YSZ anode	$egin{align*} r_{CH_4} &= rac{1}{(1+K_{CH_4}p_{CH_4}+K_{H_2O}p_{H_2O}/p_{H_2})^2} \ & k \cdot p_{H_2}^c \cdot p_{H_2O}^b \cdot p_{H_2O}^c \cdot p_{H_2O}^c \cdot \left(1-rac{p_{CO}p_{H_2}^3}{K_1p_{CH_2}p_{H_2O}} ight) \end{split}$	923–1048	57	1.83–5.50	1052	43.55	a = 0.43, b = -0.34, c = -0.02 $k_0 = 4.07 \text{ mol/}(bar^{-0.07}g_{cat}^{-1}min^{-1})$
		$r_{CH_4} = \frac{1}{(1 + K_{CH_4}p_{CH_4} + K_{H_2O}p_{H_2O}/p_{H_2})^2}$						

Table 2
WGS kinetic models over various catalysts.

REF	Catalyst material	Rate Expression	Temperature (K)	Wt% Ni	SC	Current density	Kinetic Parameters
[120]	Iron and Copper-based	$r_{WGS} = \frac{K_0 P_{CO} P_{H_{2O}} \left(1 - \frac{P_{CO_2} P_{H_2}}{P_{CO} P_{H_2O} K_e}\right)}{\left(1 + K_1 P_{CO} + K_2 P_{H_2O} + K_3 P_{CO_2} + K_4 P_{H_2}\right)^2}$	673–973	-	-	-	$\begin{array}{l} k_0 = 0.92 \ mmol \ g^{-1} s^{-1} atm^{-2} \\ E_{ao} = 4080 \ J \ mol^{-1} \\ k_1 = 2.21 mmol \ g^{-1} s^{-1} atm^{-2} \\ E_{a1} = -910 \ J \ mol^{-1} \\ k_2 = 0.4 \ mmol \ g^{-1} s^{-1} atm^{-2} \\ E_{a2} = -1420 \ J \ mol^{-1} \\ k_3 = 0.0047 \ mmol \ g^{-1} s^{-1} atm^{-2} \\ E_{a3} = -24720 \ J \ mol^{-1} \\ k_4 = 0.052 \ mmol \ g^{-1} s^{-1} atm^{-2} \end{array}$
[121]	Ni-YSZ	$r_{WGS} = K_{sf} \left(P_{H_2O} P_{CO} - \frac{P_{H_2} P_{CO_2}}{K_{ps}} \right)$	523–1073	-	0.35–6	7300	$E_{a4} = -14400 J mol^{-1}$ $K_{sf} = 0.0171 \exp\left(-\frac{103191}{RT}\right) mol m^{-3} Pa^{-2} s^{-1}$
[100]	0/70./	F \ / \ P P \	070 670		1–3		$K_{ps} = \exp(-0.2935Z^3 + 0.6351Z^2 + 4.1788Z + 0.3169)$
[122]	Cu/ZnO/ Al ₂ O ₃	$r_{WGS} = k_0 \exp\left(-\frac{E_a}{RT}\right) \left(P_{CO}P_{H_2O} - \frac{P_{CO_2}P_{H_2}}{K_e}\right)$	373–673	-	1–3	_	$E_a = 47.4 kJ mol^{-1}$ $k_0 = 2.96 \cdot 10^5 mol hr^{-1} Pa^{-2}$
[45]	Ni-YSZ and Ni-SDC	$k_{eq,WGS} = \exp\left(-\frac{\Delta G_{wg}}{RT}\right)$	1073–1223	-	0–3	0–6000	_
[123]	Fe ₂ O ₃ /Cr ₂ O ₃ / MgO	Elementary reaction mechanism	473–723	-	-	-	It varies at different steps
[124]	Ni	$r = kp_{CO}^{l}p_{H_{2}O}^{m}p_{CO_{2}}^{n}p_{H_{2}}^{q}\left(1 - \frac{p_{CO_{2}}p_{H_{2}}}{K_{eq}p_{CO}p_{H_{2}O}}\right)$	573–1273	-	-	-	$k_0 = 8e6 s^{-1}, E_a = 85 kJ mol^{-1}$
[124]	Ni-ceria	$r = kp_{CO}^{l}p_{H_{2}O}^{m}p_{CO_{2}}^{n}p_{H_{2}}^{q}\left(1 - \frac{p_{CO_{2}}p_{H_{2}}}{K_{eq}p_{CO}p_{H_{2}O}}\right)$	573–1273	-	-	-	$k_0 = 1.7e8 \ s^{-1}, \ E_a = 85 \ kJ \ mol^{-1}$
[125]	Green NiO	$r = kp_{CO}^{x}p_{H_2O}^{y}p_{CO_2}^{z}$	523–553	-	-	-	$E_a = 15 - 20 \ kcal \ mol^{-1}, x = 0 - 0.7, \ y = 0 - 0.3$
[125]	Black NiO	$r = kp_{\mathrm{CO}}^{\mathrm{x}} p_{\mathrm{H}_2\mathrm{O}}^{\mathrm{y}} p_{\mathrm{CO}_2}^{\mathrm{z}}$	535–553	-	-	-	$E_a = 23 \pm 1 \ kcal \ mol^{-1}, \ x = 0.4 - 0.8, \ y = 0,$ z = -0.4
[105]	Ni–Al ₂ O ₄ / MgO ₂	$m{r} = rac{k_H p_{CO}^a p_{H_2O}^b}{p_{H_2}^c}$	673–873	40	4	-	$k_{H0} = 5.49e6 s^{-1} bar^{-1}, E_a = 83.7 kJ mol^{-1}, c = $ $-\frac{1}{07} - 0.55, a = 1.04, b = -0.03 - 1.01$
[126]	Ni–Al ₂ O ₃	$r = kc_{CO}c_{H_2}^{0.3}$	1069–1112	11	3–5	-	$k_0 = 3100 m^{3.9} kg^{-1} s^{-1} mol^{-0.3}, E_a = 82 kJ mol^{-1}$
[127]	Ni–Re	$r = k_2 p_{CO}^a p_{H_2O}^b p_{H_2}^c p_{CO_2}^d \left(1 - \frac{p_{H_2} p_{CO_2}}{K_{ ho ho} p_{CO} p_{H_2O}} ight)$	573–673	-	0.6–1	-	$k_{20} = 2.55e - 6$, $E_a = 33.48 \text{ kJ mol}^{-1}$, $a = 1.9645$, $b = 1.9645$, $c = 4.2979$, $d = -9.4584$
[53]	Ni–CeO ₂	$r = k_2 p_{CO}^a p_{H_2O}^b p_{f_2}^c p_{CO_2}^c \left(1 - \frac{p_{H_2} p_{CO_2}}{K_{eq} P_{CO} P_{H_2O}} \right)$ $r = k \left(y_{CO} y_{H_2O} - \frac{y_{CO_2} y_{H_2}}{K_{eq}} \right)$	523–873	5–12.5	-	-	$lnk_o = 5.241 \pm 1.6, E_a = 42.51 \pm 7.9$

remainder being CO_2 . This variation means that the energy content of biogas can vary; the lower heating value (LHV) is between 16 MJ/m^3 and 28 MJ/m^3 . In addition, raw biogas often contains considerable quantities of undesirable trace compounds such as hydrogen sulphide and siloxane that can cause SOFC degradation at low concentrations. The amounts of methane and trace contaminants vary significantly on the working condition of the production unit and the feeding biomass composition [3,14].

However, the viability of using biogas in SOFCs has not been systematically and comprehensively investigated. Therefore, recent progress in the effective utilization of biogas in SOFCs is reviewed for $\rm H_2$ and power productions. Perspectives for the current advances, challenges, and the direction of future work are also discussed in this section.

4.1. Power production

The direct utilization of hydrocarbons in a SOFC was first reported by Park et al. [142–144] as early as 1995 but the power density reported was low. Murray et al. [145] reported a direct Methane-Solid Oxide Fuel Cell (CH₄–SOFC) without carbon deposition when a 0.5 μ m thick (Yttrium Oxide)_{0.15}(Cerium (IV) Oxide)_{0.85}(Yttria Doped Ceria) ((Y₂O₃)_{0.15}(CeO₂)_{0.85}(YDC)) porous film was applied between the Ni-YSZ anode and YSZ electrolyte.

Lanzini et al. [146] investigated the behaviour of planar SOFCs fed by two simulated biogas compositions coming from anaerobic digestion, namely bio-hydrogen (bio-H₂) and bio-CH₄. Ni-YSZ anode-supported cells and Ni-GDC anode electrolyte-supported cells were used. The oxidant addition was shown to help prevent carbon deposition and

accelerate CH $_4$ conversion into H $_2$. Furthermore, the cell voltage was stable when loads of $0.5~\text{mA/cm}^2$ and $0.3~\text{mA/cm}^2$ were applied to the anode-supported and electrolyte-supported cells, respectively, for at least 50 h at 800 °C. This study developed an energy model of a complete SOFC system operating on reformed bio-CH $_4$. The steam-reformed CH $_4$ demonstrated the best performance, with more than 41% electrical efficiency.

The performance of three configurations of biogas-fuelled SOFC Micro-Combined Heat and Power (micro-CHP) systems for residential applications has been evaluated by Farhad et al. [147]. Three configurations with anode off-gas recirculation, steam reforming, and partial oxidation were investigated in this study. Depending on the size, location, building type, and design of a residential unit, these systems may be able to provide the domestic hot water and electric power requirements of the unit. A sensitivity analysis of the cell operating voltage, the fuel utilization, the CHP efficiency, the biogas fuel composition, excess aid to control the stack temperature, and the thermal to electric ratio was conducted through a detailed analysis. According to the findings, the optimum cell voltage is higher than the cell voltage at which the SOFC stack's minimal number of cells is obtained. Furthermore, the exergy analysis revealed that the systems under consideration have a significant potential for generating additional electric power, particularly if the partial oxidation system is integrated with other power generation devices and adjusted suitably. However, the sensitivity analysis of the partial oxidation on the system performance was missing in this study for an accurate evaluation of such systems.

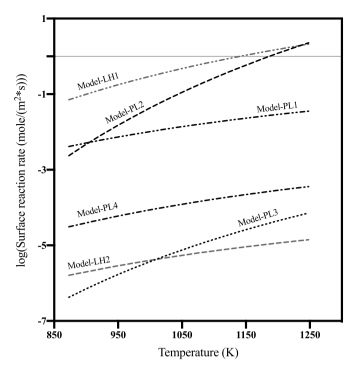
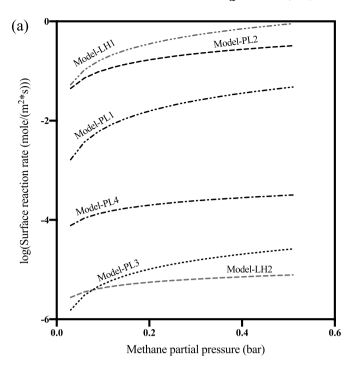


Fig. 6. The methane steam reforming reaction rates versus temperature on Ni-YSZ anodes. The six selected models are: Model PL-1 ([89]), Model PL-2 ([90]), Model PL-3 ([95]), Model PL-4 ([44]), Model LH-1 ([86]), Model LH-2 ([44]).

The direct application of biogas in an anode-supported SOFC can give good MSR performance and assure high conversion of $\mathrm{CH_4}$ even when the temperature is much below the nominal value. Santarelli et al. [66] investigated the direct reforming of biogas on Ni-based SOFC anodes experimentally. A mathematical model comprising the reforming kinetics on Ni catalysts was utilised to forecast the gas composition profile across the fuel channel. The operational maps of a fuel cell based on direct reforming of biogas were developed. Experiment-derived polarization curves were used to validate the model.

A biogas-fed decentralized SOFC combined heat and power (CHP) system model, consisting of a pre-reformer, an SOFC stack, an afterburner and a heat-recovery boiler, was proposed and analyzed by Wang et al. [57]. The system model integrates a multi-scale hierarchical 3D SOFC stack model with 0D balance of power (BoP) component models, enabling simultaneous investigations of the overall system performance and the stack-internal distributed properties down to the electrode scale. The effects of steam to carbon ratio, biogas composition and operation voltage of the SOFC stack on the electrical and CHP efficiencies of the system and the temperature gradient within the SOFC stack were studied. The proposed system model is an insightful and powerful tool for designing hybrid SOFC combined heat and power systems. However, the stack durability may decrease with increasing CH₄ in biogas due to the increase of both the temperature gradient and the carbon coverage on the electrodes. Furthermore, the increase in operating voltage leads to a decrease in the system electrical efficiency and the stack temperature gradient but an increase in the system CHP efficiency.

The power generation using CH_4 in SOFCs has been widely researched and applied. However, power density may improve by modifying the fuel cell geometry or designing better anode materials. The power density of SOFC fuelled with humidified CH_4 has been increased fast from 50 mW/cm² at 700 °C on Ni-YSZ anode and 150 mW/cm² on the Nickel–Cerium (IV) Oxide-Yttria-Stabilized Zirconia (Ni/CeO₂/YSZ) anode [143] in 1999 to 640 mW/cm² at 900 °C on the Ni-Scandia Stabilized Zirconia (ScSZ) zirconia anode in 2004 [148]. A power density of 545 mW/cm² was achieved with 30% CH_4 on micro-channelled Ni-GDC anode support as it provides a rapid gas



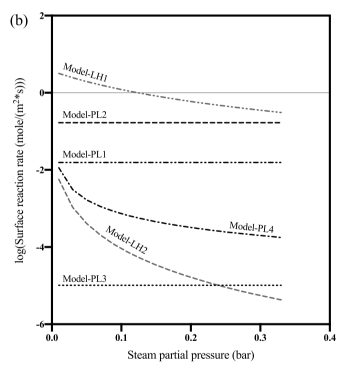


Fig. 7. (a) The methane steam reforming reaction rates versus methane partial pressure on Ni-YSZ anodes; (b). The methane steam reforming reaction rates versus steam partial pressure on Ni-YSZ anodes. The 6 six selected models are: Model PL-1 ([89]), Model PL-2 ([90]), Model PL-3 ([95]), Model PL-4 ([44]), Model LH-1 ([86]), Model LH-2 ([44]).

diffusion pathway [48]. A SOFC with a Ni-YSZ anode and Nickel/Cerium (IV) Oxide-Aluminum Oxide (Ni/CeO $_2$ -Al $_2$ O $_3$) layer achieved a power density of 830 mW/cm 2 with 25% CH $_4$ /25% CO $_2$ /Ar at 800 °C [149]. Recently, cells with samarium modified Ni-YSZ anode operated on humidified CH $_4$ reached a power density of 1540 mW/cm 2 at 800 °C [150]. A complete understanding of the kinetics of various anode material properties helps realize the efficient yet safe application of biogas in

SOFCs. However, this current review will not discuss the anode materials, and the readers can refer to other reviews from different perspectives [16,39,51,128–132].

4.2. Hydrogen/syngas and power co-production

Although many technologies for green $\rm H_2$ production are inefficient and non-competitive with conventional methods, SOFC technology stands out as a potent alternative. Co-generation of electricity and syngas from $\rm CH_4$ via SOFCs to reach high combined efficiency and achieve zero-emission is highly attractive to enhance the cost-effectiveness of biogas utilization. However, achieving high power output and syngas formation rates with sufficient operational stability remains a significant challenge in existing SOFCs.

A prototype SOFC reactor with microchannel reactors integrated into the anode has been designed to directly co-generate power and syngas from CH₄. Fan et al. [151] demonstrated effective co-generation by incorporating a catalytic microchannel reactor into the anode. The integrated anode contains distinct dendritic channels filled with a highly efficient nanofibrous Ni-based composite (Ni/CeO2-Al2O3) that serves as an internal catalytic bed reformer. The novel SOFC anode structure successfully displays thermal and material coupling effects between the exothermal fuel oxidation and endothermal reforming reactions. Compared to the conventional SOFCs, the peak power density was enhanced by 25% and the syngas production and duration of stable operation by more than twice. The new SOFC architecture has a lot of practical uses. The co-generation performance can be increased further by optimizing the electrochemical oxidation of fuels and CH₄ reforming in future experiments, which may be accomplished by adjusting the anode support thickness. The combined SOFC reactor with high energy conversion efficiencies is promising for the application of the alternative solid oxides cell-based electrochemical fuel conversion systems.

Auto thermal reforming of CH4 integrated with exothermal POM, endothermal MSR, and DMR can achieve a high overall thermal efficiency. It can be utilised in conjunction with SOFCs to eliminate the requirement for oxygen while also avoiding the safety risks associated with CH₄/O₂ mixtures. In addition, electrical power is generated together with heat. An investigation on auto thermal reforming of CH₄ over an integrated SOFC reactor for power and syngas co-generation was conducted by Fan et al. [149]. The efficient electrochemical auto thermal reforming of CH₄ over a SOFC with the catalyst beds within an anode channel configuration was demonstrated. The catalyst bed configuration increased the syngas yield significantly by improving the CH₄ conversion and selectivity towards syngas. The increased fuel accessibility shows improved mass transport at the electrode-electrolyte interface, aided by integrated catalyst beds. The catalyst beds are more efficient at catalyzing CH4 reforming than a traditional catalyst layer on the anode surface. Following initial cell microstructure stabilization, the SOFC reactor demonstrated consistent cell performance and syngas output during a 120-h test, with no detectable carbon deposition.

Panagi et al. [152] used SOFCs to produce highly efficient electrical power and syngas from bio-hythane, a gaseous mixture consisting of 60/30/10 vol% CH₄/CO₂/H₂ produced through an improved two-stage anaerobic digestion (AD) process. Fuelling SOFCs with bio-hythane has been experimentally demonstrated to co-produce electricity and valuable chemicals. The increases in SOFC efficiency have shown an increase in energy yield from the two-stage AD process compared to single-stage AD in SOFCs. The presence of H₂ in bio-hythane resulted in up to 77% higher overall electrical energy yields from biomass. The maximum power density of a bio-hythane-fuelled SOFC was around 160 mW/cm² and may be improved further by modifying the bio-hythane composition. The findings show that bio-hythane generation, rather than biogas, can be a more cost-effective way to generate electricity from biomass. Many items have been successfully coproduced. The technologies described in this study could be utilised to dispose of and add value to a wide range of challenging renewable and industrial waste gas streams in

the future.

In the past 20 years, modelling studies on such systems have made the system evaluation quicker, cheaper, and safer, beneficial to the experimental studies. Combined CH $_4$ reforming with CO $_2$ and H $_2$ O in proton-conducting SOFCs (H–SOFCs) for syngas/power co-generation was studied by Chen et al. [50], and the effects of adding H $_2$ O to the CO $_2$ and CH $_4$ fuel mixture on the performance of a tubular proton-conducting SOFC were studied numerically. The results show that the CH $_4$ conversion and current density were improved after adding H $_2$ O to the fuel. Furthermore, sensitivity studies indicate that H $_2$ O addition can effectively control the H $_2$:CO ratio and that H $_2$ starvation can be alleviated, especially at high current density conditions.

Both numerical and experimental analysis of the biogas reforming process on the Ni-YSZ catalyst was carried out by Brus et al. [94,118]. A numerical model containing MSR, DMR, and WGS reactions was proposed to predict the composition of the gas mixture at the outlet of the reformer. Different thermal boundary conditions, steam to carbon ratios, and fuel compositions were examined. The modelling results agreed with the experimental results and successfully predicted the outlet $\rm H_2$ composition under different operating conditions.

Another thermal-electrochemical modelling study on the syngas and power co-generation from H–SOFC assisted by DMR was conducted by Chen et al. [65]. The coupled heat and mass transport with electrochemical and chemical reactions were fully considered, and the co-generation performance was numerically characterized. Advantages have been shown by integrating the H–SOFC process with the internal DMR process for power and syngas co-generation using GHGs.

Under its "Green Village" program, Delft University of Technology has the initiative to build a power plant (car parking lot) with the fuel cells used in vehicles for personal mobility. Exergy analysis of different system designs on the fuel cell electric vehicle as a power plant and SOFC as a natural gas reformer was conducted by Fernandes et al. [153] in 2016. The tri-generation system generates power, heat, and H2. The system is divided into three zones: H2 production, parking, and a pumping station, as shown in Fig. 8. This study evaluates four different system designs in two different facility operation modes: (a) the car as power plant (CaPP) mode, which corresponds to the open period of the facility and uses fuel cell electric vehicles (FCEVs) for electricity and water production while parked; (2) the pump mode, which compresses H₂ and pumps it to the vehicle's fuel tank. The present catalytic reformer (CR) and a SOFC acting as a reformer (SOFCR) differ in terms of reforming technology, as well as the option of combining carbon capture and storage (CCS). The results show that the SOFCR unit greatly minimizes exergy destruction, resulting in a 20% efficiency improvement in SOFCR-based system designs in both operating modes compared to CR-based system designs. It also mitigates the system efficiency reduction to only 2% by integrating a CCS unit, whereas CR-based systems have a 7-8% efficiency reduction. In Pump mode, the SOFCR-based system has a 60% tri-generation efficiency. In addition, the use of bio-CH₄ in these systems can easily be expanded.

An afterburner-powered CH_4 steam reformer for SOFC application was numerically studied by Mozdzierz et al. [154] in 2018. A 0D model of an after burner-powered fuel cell with a reformer has been developed to investigate the effect of fuel composition on SOFC performance. It is proven that SOFC can thermally support the H_2 production system if a heat exchanger-type MSR reactor is adopted. One possible application for an after burner-heated reformer is to feed the syngas produced during the reforming process in a SOFC. The high steam to carbon ratio, which is not always desirable for the MSR, positively affects cell efficiency.

In theory, any combustible fuel can generate electrical power and $\rm H_2$ in SOFCs. Gorte et al. [155] investigated the potential of ethane, 1-butene, $\it n$ -butane, and toluene as fuels for SOFCs. Reformed liquid hydrocarbons like dodecane and diesel have also been reported as good fuels for SOFCs [156,157]. Reformed bio-diesel was also evaluated as a potential fuel for SOFCs by Mehrpooya et al. [158]. However, the fuel

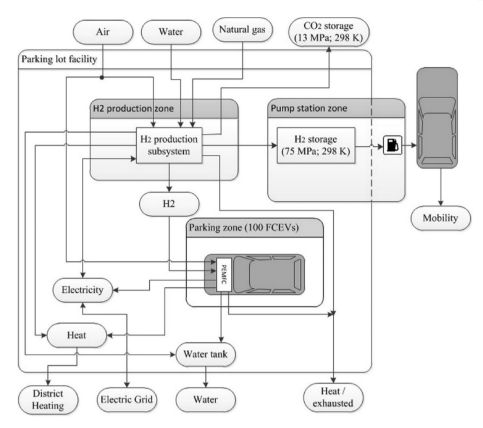


Fig. 8. The layout of the 'car as a power plant facility and flow diagram of inlet and outlet streams ((Reprinted with permission from Ref. [153], Copyright (2016), Elsevier).

cell degradation is higher than H₂-fed SOFC [159].

Leone et al. [160] investigated the performance and the degradation issues of CH₄-free biogas fuelled directly to a Ni-based anode-supported SOFC. The CH₄-free biogas is produced using an innovative process. Biomass is fermented with a pre-treated bacteria inoculum (Clostridia), which can completely inhibit the mechanization step during fermentation and thus produce an H2/CO2 mixture rather than the traditional CH₄/CO₂ anaerobic digester gas (bio-CH₄). Despite the considerable amount of sulphur in the fuel stream, the fuel cell produced an acceptable power output (at 800 °C, 0.35 W/cm² with biogas against 0.55 W/cm² with H₂). A short-term test with as-produced biogas demonstrated that sustainable biomass processing created biogas can be directly fed into SOFCs using typical anode materials with no carbon deposition, eliminating the requirement for a pre-reformer. Teramoto et al. [20] investigated CH₄-ammonia (NH₃) mixed fuel direct reforming on Ni-YSZ anode crushed materials. The reaction kinetics of combined fuel reforming/decomposition was explored experimentally. NH3 decomposition takes priority, while MSR only becomes active after enough NH₃ has been consumed. As a result, the CH₄ to NH₃ ratio in the mixed fuel can be adjusted to control local temperatures. At a specific combined ratio, a H₂ production rate was greater than pure CH₄ or NH₃.

SOFCs have been promoted as an alternative to convert biogas into electricity and heat with high efficiency. However, few studies have considered using the anode exhaust gas to co-produce green $\rm H_2$ together with electricity and heat, which could increase the performance and profitability of these systems. Thus, Nakashima et al. proposed a new approach to model SOFC with direct internal reforming to produce power, $\rm H_2$ and heat [35]. The results indicate that the proposed system can reach exergy efficiencies between 57% and 69% depending on the $\rm CH_4$ content of biogas. Furthermore, $\rm H_2$ separation reduces the amount of fuel to be burned. In addition, significant performance improvements can be achieved by optimizing the $\rm H_2$ recovery of the pressure swing adsorption and the SOFC operating temperature.

A new configuration of an integrated SOFC and gas turbine (GT) combined with a biogas reforming cycle for the cogeneration of power and $\rm H_2$ was proposed by Soleymani et al. [34]. The thermal energy discharged from the SOFC-GT system is used to supply the energy required for the reforming reaction in the biogas reforming cycle for $\rm H_2$ production. A parametric study has been performed to demonstrate the effect of different parameters on the main performance metrics of the devised system. The results revealed that the energy efficiency and exergy efficiency of the proposed combined system have increased compared to the SOFC-GT system by 23% and 28%, respectively. As a result, the net output power and $\rm H_2$ production rate are obtained by 2726 kW and 0.075 kg/s, respectively.

A biogas-based poly-generation system for the combined H2, heat and electricity production was designed and analyzed from energy and economic points of view by Mimutillo et al. [161]. The system consists of four sections: a biogas processing unit consisting of an auto thermal reactor and a water gas shift reactor, an SOFC power unit, a H₂ separation unit and a compression/storage unit. The syngas generated in the autothermal reforming reactor is split into two fluxes. The first one is sent to the SOFC power unit for electricity and heat generation. The second one is sent to the WGS reactor to increase the H2 content. The system behaviour and the energy performances were investigated by the numerical simulation based on thermo-electrochemical models. Four operating conditions related to different SOFC loads (from 30% to 100%) have been analyzed. The evaluated overall efficiencies range from 68.5% to 72.3%, and the energy-saving, calculated for the separate production of H₂, heat and electricity, ranges from about 8% to 26%. Therefore, the biogas polygene ration system can present an alternative solution to operating the existing biogas power plant being more profitable to produce H2 in the near future when the incentives for renewable electric power are not available.

Hydrogen energy systems are essential components of solutions toward reducing the negative consequences of global warming. Hydrogen should be affordable, reliable, safe, clean, and efficient for a sustainable future. Evaluation of the technical, environmental, social, and economic performance of such systems is critical for developing reliable hydrogen production and application [8,10,13,16,35,38,162-165]. By selecting the most sustainable source for hydrogen production and storage systems and enhancing the performance of hydrogen end-user technologies such as fuel cells, the demand for fossil fuels for hydrogen energy production could be reduced. Different hydrogen production sources and systems and some hydrogen storage options are comparatively investigated in detail by Acar et al. [166]. Biomass, geothermal, hydro, nuclear, solar, and wind are the selected hydrogen production sources; biological, thermal, photonic, and electrical are the selected hydrogen production methods; and chemical hydrides, compressed gas, cryogenic liquid, metal hydrides, and nanomaterials are the selected hydrogen storage systems. The economic, environmental, social, and technical performance and reliability of the chosen options are compared, providing a broader sustainability investigation of hydrogen production and storage together. A more in-depth discussion on the effectiveness and difficulties of hydrogen production is not included in this review but can be found elsewhere [5,13,15,55,163,167,168].

4.3. Advances and challenges

SOFCs provide a feasible alternative to the conventional energy conversion processes due to their highly efficient and eco-friendly operation. The high operating temperature (800–1000 °C) of SOFCs, oxide-ion conduction through the electrolyte, and the catalytic activity of the anode material for reforming reaction enable them to operate with a wide range of hydrocarbons. All techniques developed to produce $\rm H_2$ through $\rm CH_4$ reforming can be easily applied for internal reforming in SOFCs to enable carbon-neutral $\rm H_2$ production ultimately. The reforming process can occur internally in the SOFC itself or externally in a catalytic (pre-) reformer. Because SOFCs operate at high temperatures, the electrode polarization losses are often much lower than those observed in their low-temperature equivalents, resulting in significantly lower energy consumption. Furthermore, compared to low-temperature devices based on proton-conducting electrolytes, SOFCs based on $\rm O^{-2}$ conductors offer significantly higher fuel flexibility.

CH₄ reforming occurring directly on the SOFC anodes enables a less complicated system with a high overall thermal efficiency at a lower cost [153]. The current main focus of CH₄-fed SOFC operations is to increase the durability and stability of the SOFC systems and maximise the H₂ and power productions [42,169]. Issues like the fabrication and electrode materials costs [170,171], microstructural changes or degradations of anode materials during operations [172], defect segregation [173–175], insufficient current collection [176,177] and high cost for interconnect materials and fabrication [178–180] and the system design [181–183] are common to all SOFCs, regardless of the fuels. Therefore, these topics are not expanded in this review, and the interested readers can refer to the reviews by Shi et al. [184], Faes et al. [172], Badwal et al. [185,186], Tao et al. [39,134,187] and Gorte et al. [131,132,188, 189].

Biogas is a renewable energy source composed of CH₄, CO₂, and other trace compounds produced from organic matter's anaerobic digestion. Various feedstocks combined with different digestion techniques yield biogas with different compositions. Biogas derived from various biomass would be the perfect fuel for SOFCs if sufficiently clean. However, trace contaminants in the biogas vary on the biomass sources, and the treatment conditions can cause the degradation or failure of such systems. For example, municipal waste biogas contained elevated volatile chemical products such as aromatic hydrocarbons, siloxanes, and halogenated hydrocarbons. Food waste biogas contained high sulphur-containing compounds, including hydrogen sulphide, mercaptans, and sulphur dioxide. Biogas produced from dairy manure generally had lower concentrations of trace chemicals, but the combustion products had slightly higher toxicity responses than the other feedstocks [3,

6,10,14,190,191]. Efficient and cost-effective gas cleaning measures are crucial for the widespread application of such systems, which requires knowledge of the effect of the contaminants on the fuel cell performance and the tolerance limits on the anode. However, the effects of these contaminants on the fuel cell operation are not the focus of the current review and will not be discussed in-depth. Hydrogen production via dry biogas reforming, including the preparation of catalysts, the optimization of operation conditions, and the influence of impurities in biogas, has been reviewed by Gao et al. [192]. In addition, newer reactors, including membrane reactors, microreactors, and solar thermal flow reactors, were reviewed. However, the effects of various impurities in biogas on dry reforming require further study, including the mechanism of H2S and siloxane poisoning and the development of poisoning-resistant catalysts. Almost all research on dry biogas reforming has used simulated biogas as the feed gas. Future research should test dry reforming using actual biogas, an essential step to push this technology toward practical applications. A detailed review of the strategies for carbon and sulphur tolerant SOFC materials, including the conventional CH₄ reforming catalysts and novel SOFC anode materials, has been conducted by Boldrin et al. [193]. Furthermore, the interested readers can refer elsewhere for more detailed information on the advances in biogas cleaning techniques [24,60,68,194,195].

The application of current can mitigate the carbon deposition and sulphur poisoning effect on the anode materials, which is the unique advantage for CH₄ reforming in SOFCs. However, The interaction of the electrochemical reactions and the CH₄ reforming reaction, which is essential for understanding the actual reforming process in biogas-fed SOFCs [8,50,58,60,105,159], is rarely reported and often overlooked.

The effect of the electrochemical reaction on the reforming activity was first experimentally investigated by Nakagawa et al. [113] over the Ni-YSZ-CeO₂ anode. An LH-HW rate equation was proposed for the open-circuit working condition. A catalytic activity deterioration was observed with low p_{H_2} , high p_{H_2O} and high current density, possibly because of the oxidation of the Ni surface by the oxygen ions (O²⁻) or H₂O. However, no extensive investigation into the effects of current on the reforming reaction kinetics has been conducted.

The catalytic behaviour of Nickel–Zirconium Dioxide-Cerium (IV) (Ni–ZrO₂–CeO₂) anode in SOFCs was investigated by Belyaev et al. [112, 119] and how the electrochemical pumping of O₂ to or from the reaction zone influences the MSR reaction was reported. A linear increase in reaction rate with CH₄ concentration was observed. However, the anodic and cathodic polarization of the Ni–ZrO₂–CeO₂ anode did not significantly influence the reforming reaction rate.

Timmermann et al. [89] reported PL rate expressions of MSR reaction over Ni-YSZ and Ni-CGO anodes. The MSR reaction was independent of $\rm H_2O$, and the dependence of $\rm CH_4$ was slightly higher than 1. The reforming reaction has reached equilibrium at the outlet at 950 °C. They have also claimed that less than stoichiometric ratios of steam to carbon could be used under electrical load at the sacrifice of the lack of water.

Fan et al. [44,45] extended this study in which PL and LH kinetic models were used to determine and analyse the MSR reaction rate under various operating conditions. The electrochemical reaction and anode thickness effects were investigated to obtain accurate rate expressions for biogas-fed SOFC modelling studies. The findings of the study suggest that the intricacy and versatility of these models vary, and some are better at simulating real-world behaviour than others. Furthermore, since the experimental data reported were obtained under non-identical operating conditions, such as catalyst loading and feed gas composition, thus cannot be directly applied to other studies.

Additionally, the direct internal reforming of CH₄ may also prompt high-temperature gradients within the SOFC, resulting in the local thermal stresses, which accelerate cell degradation. Recent work has shown that such thermal stress severity depends on the CH₄ concentration in the fuel gas and the kinetics of electrochemical and MSR reactions. Therefore, developing an appropriate kinetic expression for the

MSR reaction over the Ni-GDC anodes within the SOFC is highly beneficial in predicting accurate temperature and concentration profiles within the fuel cell and, ultimately, the safe operation, design, and development of SOFCs.

5. Perspectives and future work

Biogas is an abundant renewable energy source produced by anaerobic treatment of biological waste such as sewage sludge, agroindustrial, and industrial animal waste. Utilizing biogas instead of fossil fuels in a solid oxide fuel cell (SOFC)-based system is an excellent choice to achieve a fossil-free and sustainable energy future. Biomethane is a near-pure source of methane produced either by upgrading biogas or through the gasification of solid biomass followed by methanation. Biomethane has an LHV of around $36~{\rm MJ/m^3}$. It is extremely similar to natural gas; therefore, it can be used without changes in transportation and distribution infrastructure, end-user equipment, and natural gas vehicles.

This review provides insights into the catalytic process of methane reforming in SOFCs. This work helps formulate appropriate kinetic models from experimental studies to aid modelling studies of the direct application of methane in SOFC systems. However, the kinetic models and the kinetic parameters for the methane reforming reaction are still under debate, especially regarding the influence of steam concentration

on the overall reaction rate. Furthermore, the fundamental principles underlying the effects of the electrochemical reaction on steam dependency have yet to be explained, necessitating further research into the effect of the electrochemical reaction on reforming kinetics. To understand the impact of anode (micro-)structure on reforming kinetics, the mass transfer should be considered, and a more extensive current density range should be implemented. Future research on developing the novel anode catalyst should also be done to facilitate the direct application of biogas in the SOFCs for $\rm H_2$ and electricity production.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Nomenclature

Abbreviations

GHG Greenhouse gases

FCH-JU European Fuel Cells and Hydrogen Joint Undertaking

MSR Methane Steam Reforming

WGS Water Gas Shift

DMR Dry methane reforming SOFCs Solid Oxide Fuel Cells DIR Direct Internal Reforming

DIR-SOFCs Direct Internal Reforming-Solid Oxide Fuel Cells

CH₄-SOFC Methane-Solid Oxide Fuel Cells

 $(Y_2O_3)_{0\cdot 15}(CeO_2)_{0.85}(YDC) \quad (Yttrium\ Oxide)_{0.15}(Cerium(IV)\ Oxide)_{0.85}(Yttria\ Doped\ Ceria)$

CHP Combined Heat and Power

Micro-CHP Micro-Combined Heat and Power CCHP Combined Cooling, Heating and Power CHHP Combined Heat, Hydrogen and Power

ER Eley-Rideal
BoP Balance of Power
AD Anaerobic Digestion
GA Gasification Agent
TPB Triple Phase Boundary

Ni-YSZ Nickel-Yttria-Stabilized Zirconia

YSZ Yttria-Stabilized Zirconia

Ni-MgAl₂O₄ Nickel-Magnesium Aluminate

Ni–MgO Nickel–Magnesium Oxide Ni-αAl₂O₃ Nickel-αAluminum Oxide

Ni-YSZ-CeO₂ Nickel-Yttria-Stabilized Zirconia-Cerium (IV) Oxide

Ni-GDC Nickel-Gadolinium Doped Ceria

Au-Ni/GDC Gold-Nickel/Gadolinium Doped Ceria

Ni-ZrO2-CeO2 Nickel-Zirconium Dioxide-Cerium (IV) Oxide

Bio-H₂ Bio-hydrogen Bio-CH₄ Bio-methane

Ni/CeO₂/YSZ Nickel-Cerium(IV) Oxide-Yttria-Stabilized Zirconia

Ni/CeO2-Al2O3 Nickel/Cerium(IV) Oxide-Aluminum Oxide (Ni-based composite)

Ni-ScSZ Nickel-Scandia Stabilized Zirconia POM Partial Oxidation of Methane CFD Computational Fluid Dynamics SC Steam to Carbon (ratio) PL Power Law

PL-1 Power Law model 1
PL-2 Power Law model 2
PL-3 Power Law model 3
PL-4 Power Law model 4
LH Langmuir-Hinshelwood

LH-1 Langmuir-Hinshelwood model 1 LH-2 Langmuir-Hinshelwood model 2

LH-HW Langmuir-Hinshelwood-Hougen-Watson

LHV Lower Heating Value
0D Zero-dimensional
1D One-dimensional
2D Two-dimensional
FO First-Order

Ni-CGO Nickel-Ceria Gadolinium Oxide Ni-SDC Ni-Samarium Doped Ceria Ni-ZrO₂ Nickel-Zirconium Dioxide

H-SOFCs Steam in proton conductive SOFCs

AD Anaerobic digestion
CaPP Car as Power Plant
FCEVs Fuel Cell Electric Vehicles
CR Catalytic Reformer

SOFCR SOFC operating as reformer CCS Carbon Capture and Storage

Cu/ZnO/Al₂O₃ Copper/Zinc Oxide/Aluminium Oxide

 $Fe_2O_3/Cr_2O_3/MgO$ Ferric Oxide/Chromium Oxide/Magnesium Oxide

Ni-Al₂O₄/MgO₂ Nickel-Aluminate/Magnesium Dioxide

Ni-Al₂O₃ Nickel-Aluminum Oxide

Ni-Re Nickel-Rhenium

Ni-CeO₂ Nickel-Cerium(IV) Oxide

Latin symbols

Subscripts

 K_i Temperature dependence of the adsorption coefficient

 $K_{eq,wgs}$ Equilibrium constant for the WGS reaction

k_i Rate coefficient

 $\begin{array}{ll} p_{CH_4}^a & \text{Partial pressures of methane} \\ p_{H_2O}^b & \text{Partial pressures of steam} \\ p_{H_2}^c & \text{Partial pressures of hydrogen} \end{array}$

 p_{CO}^d Partial pressures of carbon monoxide

a Reaction orders of methane
 b Reaction orders of steam
 c Reaction orders of hydrogen

d Reaction orders of carbon monoxide

 $\begin{array}{ll} p_{O_2} & \text{Partial pressure of oxygen} \\ p_{H_2O} & \text{Partial pressure of steam} \\ p_{CH_4} & \text{Partial pressures of methane} \\ p_{H_2} & \text{Partial pressure of hydrogen} \end{array}$

 p_{CO} Partial pressures of carbon monoxide

 X_1 Active site 1 X_2 Active site 2

S₀ Proportional to the vacant surface

n Number of sites involved in the molecular reaction

k₀ Rate Constant

k Reaction Rate Pre-Exponential Factor

 Q_{msr} Quotient of the reaction

- K_{msr} Equilibrium constant for the MSR reaction
- K_i Equilibrium constant for adsorption
- K_{eq} Equilibrium constant
- y_i Concentrations of reagents in mole fraction

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