Innovative Methane Conversion Technology Using Atmospheric Pressure Non-thermal Plasma

Tomohiro Nozaki* and Ken Okazaki

Dept. of Mechanical and Control Engineering, Tokyo Institute of Technology, 2-12-1 O-okayama, Meguro-ku, Tokyo 152-8552, JAPAN

(Received November 9, 2010)

Non-thermal plasma assisted methane reforming is reviewed. Plasma catalysis is one of the innovative next generation green technologies that may meet the needs for energy and materials conservation as well as environmental protection. Non-thermal plasma uniquely generates reactive species independently of reaction temperature, and these species are used to initiate chemical reactions at unexpectedly lower temperatures than normal thermochemical reactions. Non-thermal plasma thus broadens the operation window of existing chemical conversion processes, and ultimately allows modification of the process parameters to minimize energy and material consumption. The general aspects of plasma assisted fuel reforming including arc plasma to non-thermal plasma are described. We specifically focus on dielectric barrier discharge (DBD) as one of the viable non-thermal plasma sources for practical fuel reforming. Two contrasting approaches of DBD-oriented plasma catalysis of methane are introduced: (1) Low temperature (300-500°C) methane steam reforming using a plasma-catalyst hybrid reactor, and (2) Room temperature direct methane conversion to methanol using a microplasma reactor. The practical background and unique characteristics of each application such as the plasma-catalyst synergistic effect and highly non-equilibrium product distribution are explained.

Keywords
Plasma, Microreactor, Synergy, Non-equilibrium, Methane, Methanol

1. Introduction

Current industrial material and energy conversion technology is based on thermochemical processes including various catalytic reactions. Present industry scale technology and related science is well established, but further improvements in energy efficiency and material saving are required. Drastic reductions in CO₂ emissions are also important with the growing concern for energy and environmental issues. Green chemistry is a rapidly growing field of science and technology, and often emphasizes renewable bioenergy, bioprocesses, and solar photocatalysis of water splitting and CO₂ regeneration to provide synthetic fuels. Plasma catalysis of hydrocarbon feedstocks is also an important component of innovative next generation green technologies that are expected to satisfy the needs for energy saving, environment protection, and resource conservation. Non-thermal plasma has the specific characteristic of generating reactive species almost independently of reaction temperature. Such plasma-generated reactive species are used to initiate chemical reactions at much lower temperatures than the corresponding normal thermochemical reactions. Therefore, non-thermal plasma extends the operation window of existing chemical conversion processes, which will ultimately enable better control over the process parameters to save energy and resources, and to protect the environment.

This review briefly describes the general aspects of plasma fuel reforming from high temperature arc plasma to cold non-thermal plasma. Various plasma sources are available, but this review specifically focuses on dielectric barrier discharge (DBD) as one of the most viable non-thermal plasma sources for practical fuel reforming. First, elementary electron collision processes in non-thermal plasma are briefly introduced for those who may not be familiar with plasma chemistry. Two contrasting approaches of DBD-oriented plasma catalysis of methane, based on our work, are then introduced: (i) Low temperature (300-500°C) methane steam reforming using a DBD-catalyst hybrid reactor, and (ii) Direct conversion of methane to methanol at room temperature using a microplasma reactor, providing flexible solutions to future C₁ chemistry. Both the practical background and the mechanistic aspects of plasma catalysis such as plasma-catalyst synergistic effect and highly non-equilibrium product distribution are described. Finally, future trends are discussed.
2. Plasma Assisted Fuel Reforming

2.1. Overview of Plasma Sources for Fuel Conversion

Plasma sources can be divided into three types based on the purpose of fuel reforming: First example is low gas temperature non-thermal plasma such as dielectric barrier discharge (DBD), corona discharge, and spark discharge\(^5\). Low gas temperature non-thermal plasma consists of partially ionized gas, in which the gas temperature remains near room temperature, is readily created using a simple electrode configuration and inexpensive power source. Reactive species produced by electron impact are extremely important in fuel conversion processes. Gaseous feedstocks or light liquid fuels, after vaporization, are used as the initial feedstocks. The second example is the arc plasma where high temperature thermal energy is essential to decompose carbon containing complex materials into basic elements such as hydrogen and carbon monoxide. Arc plasma has been historically used for acetylene synthesis\(^6\) and hydrogen and carbon black co-production\(^7\). Recent applications of arc plasma have focused on gasification of biomass, coal, and municipal waste to produce syngas due to the increasing concern for energy and environmental issues. The third example is the arc plasma where high temperature thermal energy is essential to decompose carbon containing complex materials into basic elements such as hydrogen and carbon monoxide. Arc plasma has been historically used for acetylene synthesis\(^6\) and hydrogen and carbon black co-production\(^7\). Recent applications of arc plasma have focused on gasification of biomass, coal, and municipal waste to produce syngas due to the increasing concern for energy and environmental issues. The third example is relatively new, and known as “warm discharge”\(^1\). The definition of warm discharge is not clear, but may be characterized as non-thermal (cold) plasma generated in a relatively high temperature gas media. Warm discharge focuses on the combination of radical species created by non-thermal plasma and thermal energy to improve chemical conversion efficiency. Gliding arc and related technology is frequently used for this purpose\(^3,\(^9\)). The thermal energy is supplied either by the plasma, heat released by exothermic reactions such as partial oxidation, or external waste heat via an appropriate heat exchanger. Thermal energy is partly recovered through endothermic reactions such as steam reforming to enhance overall energy and material conversion efficiency. We have been focusing on DBD exclusively as one of the viable plasma sources for fuel reforming due to its flexibility and better controllability compared with other plasma sources.

2.2. Dielectric Barrier Discharge (DBD)

DBD, corona discharge, and spark discharge are commonly used for fuel reforming\(^5\), but we exclusively used DBD due to several reasons. First, DBD has a long history originating in ozone synthesis, so the fundamental physical and chemical properties, and industrial installation requirements are well understood\(^10\). Second, non-thermal plasma is readily generated: DBD rarely transits to unstable spark or arc plasma due to the presence of the dielectric barrier in the gap. Finally, electrode and reactor configurations allow great flexibility: Combination of DBD and functionalized porous media such as reforming catalysts has been rapidly growing since 2000\(^2\). The reactor can be also constructed with inexpensive materials such as glass and polymers. A peculiarity of DBD is the presence of dielectric insulator on one or both metallic electrodes, which leads to the formation of numerous filamentary microdischarges of nanosecond duration. **Figure 1** shows the typical voltage and current waveforms observed in methane-fed DBD\(^11\) as well as an image of the microdischarge taken with a high-speed intensified CCD camera with 10 ns exposure time (i-Star DH712; Andor Technology). A number of nanosecond current pulses with 1-10 ns duration are observed during every half cycle of applied voltage. The immediate termination of the developing discharge as a result of charge built up on the dielectric barrier creates highly reactive non-equilibrium conditions at atmospheric pressure. Because of the charge accumulation on the barrier, microdischarges are dispersed randomly between the electrodes, providing a wide range of ionizing discharge gaps in time and space average. Although DBD in air has been studied extensively for ozone synthesis\(^10\), limited information is available for methane-fed DBD. Detailed gas phase plasma chemistry of methane-fed...
DBD has been described previously\textsuperscript{22,(13)}.

2.3. Elementary Collision Process between Electron and Methane

The reaction rate for electron impact excitation of methane is expressed as follows:

\[
d\langle \text{CH}_4 \rangle \over dt = k \cdot Ne \cdot \langle \text{CH}_4 \rangle
\]  

Eq. (1)

Here, \( k \) is the rate constant (cm\(^3\)/s/molec), and \( Ne \) is the electron density (molec/cm\(^3\)). Electron density in a single filamentary discharge formed in air, for example, is approximately \( 10^{14} \) cm\(^{-3}\). \( k \) is estimated using the Eq. (2):

\[
k = \int \sigma(E) \cdot v \cdot f(E)dE
\]  

Eq. (2)

Here, \( \sigma(E) \) is the electron-methane collision cross section, \( E \) is the electron energy, \( v \) is the electron drift velocity, and \( f(E) \) is the electron energy distribution function (EEDF). EEDF is a function of temperature, pressure, type of gas, and electric field, and incorporates a large displacement from the Boltzmann distribution, so the Boltzmann equation for practical EEDF must be solved under given conditions to evaluate the appropriate rate constant. Assuming the standard conditions of methane (298 K and 101 kPa), EEDF and other swarm parameters were calculated using ELENDIF software that solves the time evolution of the electron energy distribution function using two-term spherical harmonic expansion of the Boltzmann transport equation\textsuperscript{30}. Assuming Boltzmann and non-Boltzmann electron energy distribution functions, rate constants for vibrational (\( k1 \)), dissociation (\( k3 \)) and ionization (\( k7 \)) collisions\textsuperscript{15} were calculated as shown in Fig. 2 (see also Table 1). There is a small deviation for vibrational excitation because the excitation threshold (0.16 and 0.36 eV) is small enough that electron energy distribution and vibrational cross section overlap considerably; therefore, vibrational excitation rate constants can be estimated by assuming the Boltzmann EEDF. However, the Boltzmann EEDF overestimates the non-Boltzmann rate constants for dissociation and ionization, although the electron mean energy is identical for both cases. Electrons at the high energy tail of EEDF can cause dissociation and ionization of methane, and lose energy through those collisions. As a result, the probability of the presence of high energy electrons decreases, so the rate constants with experimental EEDF are smaller than those estimated with the Boltzmann EEDF.

| \( k1 \) | \( \text{CH}_4(\nu_{2a}) + e \) | 0.162 | bending |
| \( k2 \) | \( \text{CH}_4(\nu_{3c}) + e \) | 0.361 | stretching |
| \( k3 \) | \( \text{CH}_4 + H + e \) | 9.0 | dissociation |
| \( k4 \) | \( \text{CH}_2 + \text{H}_2 + e \) | 10.0 | dissociation |
| \( k5 \) | \( \text{CH} + \text{H}_2 + H + e \) | 11.0 | dissociation |
| \( k6 \) | \( \text{C} + 2\text{H}_2 + e \) | 12.0 | dissociation |
| \( k7 \) | \( \text{CH}_3 + e + e \) | 12.6 | ionization |
| \( k8 \) | \( \text{CH}_3^+ + e + e \) | 14.3 | ionization |

Table 1  Eight Elementary Processes Considered in Fig. 2

The rate-determining step of methane conversion is dissociation of the strong C-H bond. Here, three different approaches are compared: R1, homogenous gas phase reaction; R2, heterogeneous surface reaction; and R3, plasma-enhanced reaction.

\[
\text{CH}_4(\text{+thermal energy}) = \text{CH}_3 + \text{H} \quad \text{El} = 434 \text{ kJ/mol (R1)}
\]

\[
\text{CH}_4(\text{+Ni}) = \text{CH}_3^+ + \text{H}^+ \quad \text{E2} = 100 \text{ kJ/mol (R2)}
\]

\[
\text{CH}_4 + e = \text{CH}_3 + H + e \quad \text{E3} = 868 \text{ kJ/mol (R3)}
\]

E1-E3 express activation energy and superscript “*” denotes chemisorbed fragments. The activation energy for methane pyrolysis (R1) is 434 kJ/mL\textsuperscript{17} so requires high temperature thermal energy (>1000°C). The activation energy and the reaction temperature are greatly reduced by the use of transition metal catalysts (R2)\textsuperscript{18}. A peculiarity of non-thermal plasma is that chemically stable methane is dissociated by electron impact almost independently of reaction temperature, but inelastic electron collision requires a large activation energy\textsuperscript{15}. Compared to the endothermic enthalpy of methane dis-
Biogas is obtainable from landfills, coal mines, and other organic resources. Methane content decreases below 40% if air enters and oxidizes methane during fermentation, resulting in much poorer biogas. Although a huge amount of poor biogas is obtainable from landfills, coal mines, and other sources, most is simply flared and wasted because of the high global warming potential (GWP) of biogas, which is 5-15 times as potent as carbon dioxide, depending on the biogas's composition. The poor ignition stability and low heating value of such biogas make its use in a conventional energy utility system extremely difficult. We developed a low-temperature (300-500°C) method for upgrading simulated biogas using DBD generated in a reforming catalyst bed(12,13,19-21). The simulated biogas is partially converted into hydrogen via low-temperature steam reforming; then the hydrogen-enriched biogas is fed into an internal combustion engine. Ignition stability and combustibility of hydrogen enriched biogas are remarkable, and the internal combustion engine was powered efficiently. Low-temperature steam reforming is essential so that part of the biogas, with low heating value (20-60% of CH₄), or additional fuel does not have to be used to provide a high-temperature heat source. Low-temperature exhaust gas from an internal combustion engine (300-500°C) is useful to activate the catalyst bed and heat is partly recovered as chemical energy of hydrogen.

3. Endothermic Reaction System: Methane Steam Reforming Using DBD-catalyst Hybrid Reaction

3.1. Introduction

The primary role of a non-thermal plasma is to generate reactive species at the lowest reasonable temperature possible. The reactive species are used to initiate chemical reactions at low temperature or to create unusual reaction pathways which result in non-equilibrium product distributions at a given temperature and pressure. Therefore, non-thermal plasma can enable better process control over operating parameters. However, radical injection is not satisfactory to enhance energy and material conversion efficiency if non-thermal plasma is combined with endothermic reactions such as steam reforming. Therefore, for conservation of energy, thermal energy must be supplied simultaneously. Without appropriate combination with thermal energy, non-thermal plasma cannot promote the overall chemical conversion process. In this section, the DBD-catalyst hybrid reaction for the low temperature (300-500°C) methane steam reforming is discussed. Appropriate combination of DBD and low temperature thermal energy is the key to establish an economically viable plasma fuel conversion process.

3.2. Background of the Project

This project was initiated to develop profitable recovery of methane from poor bio-resources using non-thermal plasma technology. Biogas generally consists of 60% methane and 40% CO₂ if methane fermentation of organic matter is the main digestion process. Methane content decreases below 40% if air enters and oxidizes methane during fermentation, resulting in much poorer biogas. Although a huge amount of poor biogas is obtainable from landfills, coal mines, and agricultural residues, most is simply flared and wasted because of the high global warming potential (GWP) of biogas, which is 5-15 times as potent as carbon dioxide, depending on the biogas's composition. The poor ignition stability and low heating value of such biogas make its use in a conventional energy utility system extremely difficult.
between the electrodes. Gas breakdown occurred at the pellet contacts where the electric field is concentrated and a number of filamentary discharge channels propagated along the pellet surface. Detailed characteristics of this type of discharge were described previously. The total gas flow of simulated biogas was adjusted to a gaseous hourly space velocity (GHSV) of 10,000-18,000 h⁻¹.

3. 4. Synergistic Effect between Non-thermal Plasma and Reforming Catalysts

Figure 4 (a) shows methane conversion in (1) a catalyst bed reactor (without DBD), (2) a hybrid reactor, and (3) a DBD reactor with α-Al₂O₃ pellets (without catalysts). The results are plotted with respect to catalyst bed temperature measured using an infrared camera. The dotted line in Fig. 4 (a) represents the equilibrium methane conversion. The relationship between discharge power and endothermic reaction enthalpy is also important parameter because the bed temperature is determined as a result of energy balance. The results for hybrid reactor operation are presented in Fig. 4 (b). Briefly, methane was decomposed only slightly by DBD because the discharge power was minimized, which is possible to avoid excess heating. The main product was C₂H₆ and selectivity for hydrogen was much lower than the chemical equilibrium. The catalyst reaction was more efficient than DBD, but the methane conversion did not reach equilibrium due to low temperature catalysis. On the other hand, methane conversion in the hybrid reactor was clearly much greater than the simple sum of the DBD and catalyst reactions at given bed temperatures.

Methane conversion increased from points A₄₀₀, B₄₀₀, to C₄₀₀ depending on the discharge power of DBD, where the subscript denotes the temperature of the ceramic radiant heater, used to externally heat the catalyst bed. The ceramic heater functioned as a constant temperature bath. The bed temperature slightly decreased from the initial condition (400°C) because of the small amount of heat absorbed by the methane steam reforming when the discharge power was zero (at A₄₀₀) (see Fig. 5 (a)). Methane conversion increased steeply with increasing discharge power, whereas the bed temperature increased slightly (A₄₀₀ → B₄₀₀). The average bed temperature was 420°C at B₄₀₀ (Fig. 5 (b)) and methane conversion reached the chemical equilibrium. A further increase in discharge power created a localized high temperature zone in the downstream portion of the bed (Fig. 5 (c)). After chemical equilibrium was established at B₄₀₀, further increase in methane conversion beyond the equilibrium was not anticipated because the reverse reaction (CO₂ + 2H₂ → CH₄ + 2H₂O) is much faster than the forward reaction. In the equilibrium-limiting regime, the synergistic effect was no longer anticipated and heat produced by DBD becomes the main driving force for the methane steam reforming. In fact, the bed temperature increased up to 570°C and the equilibrium methane conversion was established at C₄₀₀. With the initial bed temperature was 500°C (at A₅₀₀), the forward methane reaction was also accelerated with input power until the chemical equilibrium was established (A₅₀₀ → B₅₀₀). Excited species, produced by high-energy electron impact, presumably accelerated the low temperature catalysis of methane. Simultaneously, heat produced by DBD promoted the catalytic conversion of methane. Both mechanisms were important in the hybrid reaction. The effects of radical injection and heat generation on the reaction enhancement mechanism was analyzed based on Arrhenius plots.

3. 5. Kinetic Analysis by Arrhenius Plot Method

The forward methane reaction rate based on the overall steam reforming using power-law kinetics is expressed as Eq. (3).
Here, \( r \) is the forward methane reaction rate and \( k \) is the forward methane reaction rate constant for the overall steam reforming. The reaction order for overall methane steam reforming on nickel catalyst is \( \alpha = 0.85-1.4 \) in \( \text{CH}_4 \) and \( \beta = -0.8-0 \) in \( \text{H}_2\text{O}^{26,29} \). The phenomenological understanding is that the rate-determining step is the activation of methane on the metallic catalyst, whereas the overall forward reaction rate is less dependent on the \( \text{H}_2\text{O} \) concentration\(^{18} \). The overall reaction order for \( \text{H}_2\text{O} \) (\( \beta \)) is negative if the metallic catalysts lose activity due to oxidization by \( \text{H}_2\text{O} \). In the hybrid reactor, both \( \text{CH}_4 \) and \( \text{H}_2\text{O} \) are excited primarily by electron impact, which is the driving force for modifying the catalytic methane conversion. For this reason, we determined the reaction orders (\( \alpha \) and \( \beta \)) independently with and without DBD.

Assuming that the overall rate constant is expressed in Arrhenius form, the results are shown in Fig. 6.

\[
k = A \exp \left( -\frac{E}{RT} \right)
\]

The activation energy \( E \) and the pre-exponential factor \( A \) for the overall steam reforming reaction are listed in Table 2 along with the respective reaction orders. During the experiment, the discharge power was controlled appropriately so that methane conversion did not reach chemical equilibrium. The detailed numerical procedure to evaluate the reaction orders (\( \alpha \) and \( \beta \)), pre-exponential factor (\( A \)) and overall activation energy (\( E \)) was described previously\(^{20} \).

The convex nature of the resultant Arrhenius plot implies that the overall steam reforming was in the reaction-limited regime when the bed temperature was

\begin{table}[h]
\centering
\begin{tabular}{|c|c|c|c|c|c|}
\hline
 & \( \alpha \) & \( \beta \) & \( T \text{[°C]} \) & \( E \text{[kJ·mol}^{-1}\] } & \( A \) \\
\hline
Hybrid & 0.54 & 0.20 & \( T<420 \) & 102 & \( 3.90 \times 10^7 \) \\
 & & & \( T>420 \) & 53 & \( 6.15 \times 10^7 \) \\
Catalyst & 0.45 & 0.08 & \( T<460 \) & 90 & \( 8.01 \times 10^8 \) \\
 & & & \( T>460 \) & 48 & \( 8.20 \times 10^2 \) \\
\hline
\end{tabular}
\caption{Reaction Orders, Activation Energy, and Pre-exponential Factor of the Overall Methane Steam Reforming Reaction}
\end{table}
lower than 460°C, but transited to the diffusion-limited regime when the bed temperature is higher than 460°C. Although the overall rate constant in the hybrid reaction was larger than that of the normal catalyst reaction, the overall activation energy was essentially unchanged independently of DBD. The overall activation energy in the reaction-limited regime was approximately 100 kJ/mol, which agrees well with the reported value, implying that methane dehydrogenation is the rate-determining step regardless of DBD application. On the other hand, the pre-exponential factor was enhanced by a factor of 50 by DBD. The same trend was observed in the diffusion-limited regime, where only the pre-exponential factor was enhanced by a factor of 7, whereas the overall activation energy was not influenced by DBD.

3.6. Role of Excited H₂O in Methane Steam Reforming

To enhance overall methane conversion, both methane dehydrogenation over the solid catalyst and water chemisorption should be promoted simultaneously so that unreacted chemisorbed carbon intermediates are oxidized. Otherwise, chemisorbed carbon ultimately builds up carbon filaments, which block active sites and catalyst pores. In this respect, excited H₂O is expected to increase the concentration of chemisorbed oxygen-related intermediates such as O and OH, and efficiently oxidize chemisorbed solid carbon, leaving sufficient numbers of active sites available for successive methane dehydrogenation. In fact, the reaction order for H₂O was more than doubled by applying DBD, which eventually increased the pre-exponential factor of the overall forward rate constant \( k \). The removal of chemisorbed carbon is a vital reaction pathway together with the methane dehydrogenation reaction. Nevertheless, DBD-induced carbon precipitation was uniquely identified on catalyst pellets. Therefore, the carbon to steam (S/C) ratio must be increased to prevent carbon formation as reaction temperature increases.

3.7. Operational Regime of Hybrid Reaction

The operational regime of the DBD-catalyst hybrid reaction was clearly established, as schematically shown in Fig. 7. In the presence of DBD, methane conversion increases from 0 to 1, then further increases beyond chemical equilibrium at given temperature (\( \text{I} \rightarrow \text{II} \) at \( T_{\text{initial}} \)). However, the excess amount of CO₂ and H₂ present in the reactor drives the backward reaction (\( \text{II} \rightarrow \text{I} \)). This process results in net-zero methane conversion. The energy absorbed by reforming during the \( \text{I} \rightarrow \text{II} \) process, which could be either electrical energy or heat, is released as thermal energy through the reverse reaction. This thermal energy simply increases the bed temperature from 1 to 3. The increase in methane conversion, \( \Delta \text{CH}_4 \) shown in Fig. 7, originates from the net temperature increase of the bed. Here again, DBD enhances methane conversion from 3 to 4; but reverts to equilibrium through the reverse reaction (\( \text{IV} \rightarrow \text{III} \)). Even if DBD produces reactive species, the final effect is as a heat source once chemical equilibrium is established and a synergistic effect is no longer anticipated.

4. Exothermic Reaction System: Direct Methane to Methanol via Partial Oxidation

4.1. Introduction

In this section, direct methane conversion via partial oxidation using a non-thermal plasma combined micro-
reactor, known as a microplasma reactor, is introduced. Microplasmas are an emerging new field of plasma science and technology for materials processing. Many projects are related to chemical vapor deposition, micro fabrication, medical treatment of tissue, de-contamination of hazardous biomaterials, lighting, elemental analysis, and so on. In addition, chemical synthesis is an important part of electron-driven micro-chemistry. Non-thermal plasma “stimulates” the methane-oxygen mixture, leading to spontaneous oxidation of methane into various oxygenates at room temperature, while the heat generated by methane partial oxidation must be removed efficiently so that full combustion of methane is avoided. Unlike methane steam reforming (endothermic reaction) as discussed in the previous section, generation of reactive species at low temperature is the key to achieve direct methane conversion into the desired products at high yields.

4.2. Background of the Project

Direct conversion of natural gas into synthetic fuels such as methanol is important because the direct process can reduce the capital and operating costs of the high temperature, energy intensive multi-step processes via syngas production. Tremendous efforts have been invested in direct methane conversion to oxygenates in homogeneous gas phase reaction and over solid catalysts, but the yields of desired products have been below economic values. More recently, atmospheric pressure non-thermal plasma has been suggested as a viable synthesis method because high energy electron impact initiates methane partial oxidation, enabling single step methane conversion into various oxygenates. Nevertheless, the one-pass yield for useful oxygenates was unsatisfactory. We proposed DBD generated in a microreactor which enabled moderate methane oxidation at room temperature, and selectively produced methanol (CH$_3$OH), formaldehyde (HCHO), and formic acid (HCOOH) with one-pass yield of more than 10%. Use of non-thermal plasma in a microreactor allows application of unconventional thermochemical conditions to materials processing, enabling better control over process parameters for the selective synthesis of desirable products.

4.3. Thermodynamic Analysis

Equilibrium calculations were performed using STANJAN software that solves the adiabatic equilibrium state of gas mixtures based on the JANAF thermochemical database. Figure 8 shows the equilibrium composition of CH$_4$, O$_2$, and oxidative products at 101 kPa with O$_2$/CH$_4$ = 0.5 at atmospheric pressure. These components were considered based on the experimental results as discussed later. Oxygen (O$_2$) is fully consumed even at room temperature. Correspondingly, 50% of methane is converted, producing full combustion products such as H$_2$O and CO$_2$ when the temperature is well below 300°C. As the temperature increases, the fraction of syngas (i.e. CO and H$_2$) increases and the remaining methane is gradually consumed. Eventually, the H$_2$/CO ratio approaches 2 as temperature approaches 800°C. This reaction scheme expresses the typical two-step methane reforming for syngas production. Liquid components were negligibly small over the temperature range. Thermodynamic analysis emphasizes several important facts: First, to achieve highest selectivity for desired products, abrupt termination of methane oxidation must be achieved at the desired moment before chemical equilibrium is established. Catalysts can accelerate specific reaction pathways preferentially for better selectivity of the process; but previous results have not been satisfactory. Non-thermal plasmas can provide a totally different methane...
activation route with different kinetics in an unusual
temperature range, thus providing better feasibility and
controllability of partial chemistry instead of the equi-
librium state.

4. Experimental

A detailed description of the microplasma reactor
was presented previously\(^{49,50}\). A schematic diagram
of the experimental setup is presented in Fig. 9. The
microplasma reactor consists of a quartz tube (I.D:
1.5 mm, O.D. 2.7 mm, length: 50 mm) equipped with
internal wire electrodes (0.5 mm stainless steel) and an
externally grounded electrode. The CH\(_4\) and O\(_2\) mix-
ture is introduced into the microplasma reactor. The
total flow rate is varied so that the reaction time of gas
flow is between 100 ms and 500 ms. A bipolar pulsed
high voltage with peak intensity of 10 kV and fixed f
re-
quency at 10 kHz is applied between the metallic elec-
trodes. The microplasma reactor is immersed into the
water bath maintained near room temperature (10
\(^{\circ}\)). The microreactor configuration efficiently removes heat
generated by methane partial oxidation. Low temper-
ature synthesis is essential because oxygenates are con-
densed on the microreactor wall which results in product
separation from the O\(_2\)-rich reactive plasma. The idea
of product separation is essential because organic oxy-
genates are much more reactive than methane, and are
readily decomposed during plasma activation of meth-
ane. Liquid water is also injected intermittently into the
microplasma reactor (1.2 cm\(^2\)・min\(^{-1}\) for 10 s and
2 min interval) to wash out condensed oxygenates,
which suppresses successive destruction of reactive
oxygenates, enabling selective synthesis of oxygenates
with high methane conversion in a single reactor.

4.5. Methane Conversion and Selectivity for

Liquid Components

Figure 10 shows methane conversion with reaction
time. The initial O\(_2\)/CH\(_4\) ratio was also varied. Methane conversion increased linearly with reaction
time. Methane steam reforming was also anticipated
because water vapor was present as a result of liquid
water injection. D\(_2\)O was injected to trace the D\(_2\)O
(i.e. H\(_2\)O) dissociation pathway, but neither HD nor D\(_2\)
were identified by quadrupole mass spectrometry
(Prima-100; Pfeiffer Vacuum Technology AG).
Electronic H\(_2\)O dissociation to produce OH, which is a
viable radical chain propagator, was anticipated, but did
not participate in methane oxidation, as liquid water
mechanically washed out condensed oxygenates before
further destruction of the desired products occurred.
Liquid components were quantitatively analyzed by
GC-MS (Shimadzu Corp., QP2010Plus), showing
methanol (CH\(_3\)OH), formaldehyde (HCHO), and formic
acid (HCOOH) as the main components.

Selectivity for individual liquid components is de-
scribed in Fig. 11. Total liquid selectivity was 40-
50% and almost independent of initial O\(_2\) content, which
implies that product separation due to liquid condensa-
tion effectively suppressed excessive oxidation of
organic oxygenates. Selectivities for oxygenates and
syngas were 5-6 orders of magnitude greater than the
chemical equilibrium as shown in Fig. 8, demonstrating that methane oxidation was successfully terminated
before the equilibrium was established. The tempera-
ture of the microplasma reactor was increased to 300-
400\(^{\circ}\) without external cooling. As a result, selectivity
for formaldehyde and formic acid decreased dramati-
cally. Selectivity for methanol was surprisingly high
in this temperature range.

4.6. Syngas Production and DME Synthesis as

Post Plasma Reaction

In addition to oxygenates, relatively large amounts of
syngas were produced. Figure 12 shows the CO and
CO\(_2\) selectivities and Fig. 13 shows the H\(_2\)/CO ratio.
CO selectivity increased with methane conversion. In
contrast, H\(_2\)/CO ratio was independent of methane con-

\[\text{O}_2/\text{CH}_4 = 0.2 (\text{△}), 0.5 (\square), 1.0 (\bullet).\]

Fig. 10 Methane Conversion vs. Reaction Time
version, and was determined by the initial O2/CH4 ratio. Small amounts of oxygen, such as O2/CH4 \( \approx 0.2 \), produced hydrogen rich syngas (H2/CO \( \approx 1.8 \)), but formation of a tar-like product was observed on the reactor wall. Note that solid carbon formation was fully eliminated because of low temperature synthesis and the absence of metallic catalysts. Syngas is the fundamental building block for C1 chemistry, and is readily converted into various chemicals under moderate conditions using well-established catalysts. We assume that one-pass dimethyl ether synthesis (DME; CH3OCH3) from syngas with H2/CO \( \approx 1.3 \) occurred as a post plasma reaction, given by,

\[
3\text{CO} + 3\text{H}_2 = \text{CH}_3\text{OCH}_3 + \text{CO}_2 \quad \text{(R4)}
\]

When initial O2/CH4 = 0.5, syngas with H2/CO \( \approx 1.2 \) was obtained. Eventually, overall liquid selectivity, \( S_{\text{total}} = S_{\text{liq}} + S_{\text{DME}} \), would be feasible:

\[
S_{\text{liq}} = [\text{CH}_3\text{OH}]_{\text{rel}} + [\text{CHOH}]_{\text{rel}} + [\text{HCOOH}]_{\text{rel}} \quad (5)
\]

\[
S_{\text{DME}} = \frac{2}{3} \times [\text{CO}]_{\text{rel}} (n > 1) \quad (6)
\]

\[
S_{\text{total}} = S_{\text{liq}} + S_{\text{DME}} \quad (7)
\]

4.7. Comparison with Conventional Thermochemical Methods

Figure 14 shows the selectivities for liquid components (\( S_{\text{liq}} \) and \( S_{\text{total}} \)) with respect to methane conversion obtained for a wide spectrum of operating conditions and compared with three conventional thermochemical methods. The conventional methods required addition
of relatively high temperature thermal energy to initiate methane oxidation, whereas small amounts of oxygen, typically 10% at most, were introduced so that successive oxidation of methanol was suppressed. Effective cooling methods, such as the water bath used in this study, could not be used because low temperature conditions do not initiate methane oxidation. As a result, methane conversion beyond 10% was hardly achieved. Methane conversion was increased by increasing either oxygen content or reaction temperature, but the selectivity for useful oxygenates dropped sharply. In our method, liquid product selectivity is slightly decreased with methane conversion, implying that liquid product separation is effective in the microplasma reactor. Although one-pass liquid yield reaching 20% is obtainable, liquid selectivity was below 60% (●). Assuming post DME synthesis with the given syngas (H₂/CO ~ 1), the overall liquid selectivity can be enhanced to 80% (▲).

5. Conclusions and Future Trends

This review article introduced the principle of plasma fuel reforming, followed by two contrasting examples of non-thermal plasma catalysis of methane. Two major conclusions were derived: First, only non-thermal plasmas might be an inefficient way of converting methane due to the large activation energy required for the electron impact process. Second, appropriate combinations of thermal energy (including heat removal) and radical species produced by non-thermal plasma can establish efficient fuel reforming. Both radical species and thermal energy are required for efficient methane conversion in methane steam reforming. Combination of reforming catalysts minimizes the discharge power required for low temperature methane steam reforming, but methane conversion as well as product distribution are dominated by the chemical equilibrium under given conditions. In contrast, methane partial oxidation by radical injection at room temperature is the key to achieve high methanol yields where the product distribution is highly non-equilibrium. In addition to radical injection, efficient heat removal using a microreactor configuration is crucial.

Hydrogen energy and related technology such as the PEFC (Polymer Electrolyte Fuel Cell) and FCV (Fuel Cell Vehicle) have motivated research activity in on-board plasma fuel reforming. However, mass production of high purity hydrogen (CO < 10 ppm) is hard to achieve in an economical way. On the other hand, fuel modification by plasma catalysts for the SOFC (Solid Oxide Fuel Cell) would be feasible because the initial feedstock does not need to be modified entirely. Hydrogen enrichment of poor bio-resources by non-thermal plasma, introduced in section 3, is also feasible because initial feedstocks are partially converted into hydrogen that greatly reduce electrical energy requirements, whereas the combustion capability of poor biogas is sufficiently improved as a result of hydrogen enrichment. Note that in principle, biogas is carbon neutral. Plasma-assisted combustion of landfill gas, which emits methane under extremely fuel-lean conditions, is interesting because the global warming potential (GWP) of landfill gas is 10 times as potent as that of carbon dioxide. Development of a portable plasma reactor may be beneficial to burn trace amounts of residual hydrogen in a fuel cell exhaust for safety purposes. Direct methane to methanol conversion using DBD has a relatively long research history, but adequate yield has not been achieved yet. The combination of DBD and a microreactor enables excellent control over heat and mass transport in the discharge field, enabling direct methanol synthesis at high yield. Much research regarding plasma fuel conversion is ongoing on the laboratory scale worldwide, but we have to be aware that plasma fuel conversion is a part of the energy utility system. Even if the energy efficiency of a plasma reactor is not as excellent as existing technology, the overall energy and material conversion efficiency of the system could be enhanced because the plasma can broaden the operation window of the system, and enables chemical reaction under unconventional situations. Further development of plasma fuel reforming has excellent potential to eventually provide viable solutions for future energy and material use.

Acknowledgments

The authors would like to thank former postdoctoral researcher Dr. Shigeru Kado (Chiyoda Corporation), Dr. Sreejit Nair (Dow Chemicals International) and Dr. Anıl Ağıral (Lawrence Berkeley National Laboratory), and Dr. Valentin Goujard (currently JSPS PD fellow) for supporting the projects. Projects were partly supported by KAKENHI, Japan.

References

要  旨

大気圧非平衡プラズマを用いた新規なメタン転換技術の開発

野崎 智洋・岡崎 健

東京工業大学大学院理工学研究科機械制御システム専攻．152-8552 東京都目黒区大岡山2-12-1

化石燃料利用体系の省エネルギー、省資源、CO₂排出抑制が求められている。現在は、熱化学的手法を基盤としたエネルギー・物質変換が重要な役割を果たしているが、長期的なエネルギー資源利用体系のあり方を考えたとき、既成技術の延長線上にない革新的な反応プロセスの開発が不可欠である。このような背景から、我々は大気圧非平衡プラズマを利用して、天然ガスやバイオガスの主成分であるメタンを高効率に転換するための技術を開発している。本稿では一般的なプラズマ燃料改質の概要を述べた後、これまで我々が取り組んできた研究を二つ紹介する。まず、プラズマ・触媒ハイブリッド反応を用いた低温メタン水蒸気改質について概要を述べた後、マイクロプラズマによるメタンからメタノールの常温直接合成を紹介する。それぞれ、具体的な応用を目指したプラズマ燃料改質の位置づけを明確にした後、プラズマ固有の反応特性、たとえばプラズマと触媒のシナジー効果、マイクロプラズマによる高速に非平衡化された化学変換の可能性などを示唆する新しい技術の概要を紹介する。最後にプラズマ燃料改質の展望を述べて本稿のくくりとする。