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Development of Multimetallic Alloy Catalysts Efficient for CO₂ utilization by Dry Reforming of Hydrocarbons

(炭化水素のドライリフォーミングによる

CO2利用に有効な多元素合金触媒の開発)

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Chapter 1

General Introduction

1. General Introduction

1.1 Recent developments in CO₂ utilization

The extensive emission of CO_2 poses a major threat to humanity, the well-known situation was intensified the greenhouse effect, causing rising seas, melting glacier and ocean acidification. This has aroused human concern about the change in the ecological environment.¹ Besides, the current situation of dwindling fossil fuels has compelled people to seek corresponding alternatives. In this context, the chemical utilization of CO_2 as an abundant and nontoxic C1 block offers an effective solution to the aforementioned two issues, which is the reason why CO_2 utilization has garnered immense global attention. ²⁻³ Many techniques, such as thermal chemical, photochemical, biochemical, electrochemical, have been utilized for CO_2 utilization to convert CO_2 into useful chemicals including syngas, methanol, and formic acid, food and beverages and CO_2 enhanced oil recovery etc..⁴⁻⁶

1.1.1 Chemicals

The use of CO_2 to synthesize chemicals is a promising way to enhance the economic value of CO_2 and relieve energy crisis. The conversion of CO_2 to methanol is of particular importance because methanol could be used not only as a sustainable source of liquid fuel but also as a reactant to produce formaldehyde, olefins, dimethyl ether, and acetic acid. The equation of methanol synthesis from CO_2 as illustrated below (Equation (1)), while the reverse water gas shift reaction also occurs as side reaction (RWGS: Equation (2)), resulting in lower methanol yield. Hence, in order to enhance the methanol yield, a wide number of explorations have recently been carried out.

$$CO_2 + 3H_2 \rightarrow CH_3OH + H_2O \qquad \Delta H_{298K} = -49.5kJ \cdot mol^{-1}$$
 (1)

2

$$CO_2 + H_2 \rightarrow CO + H_2O$$
 $\Delta H_{298K} = -41.2kJ \cdot mol^{-1}$ (2)

Cu based catalyst is the most studied catalyst among many kinds of metal-based catalysts. Some representative Cu-based and other metal catalysts for the synthesis of methanol by CO₂ hydrogenation are summarized in Table 1.1. Industrially, the ternary Cu/ZnO/Al₂O₃ catalyst used in methanol synthesis from syngas (CO/H₂) and also from CO₂ hydrogenation.⁷ However, Cu nanoparticles is prone to aggregation and separate from ZnOx under reaction conditions, which is benefit to the reverse water gas reaction,

Table 1.1 some representative catalysts for the synthesis of methanol by the hydrogenation of $CO_{2.}^{8}$

Catalyst	Preparation method	T/ºC	CO ₂ conversion (%)	Methanol selectivity (%)	Methanol activity (mol kg ⁻¹ cat. h)	Ref.
Cu/Zn/Ga/SiO ₂	co-impregnation	270	5.6	99.5	10.9	9
Cu/Ga/ZnO	co-impregnation	270	6.0	88.0	11.8	10
Cu/ZrO_2	precipitation	240	6.3	48.8	11.2	11
Cu/Ga/ZrO ₂	precipitation	250	13.7	75.5	1.9	12
Cu/B/ZrO ₂	precipitation	250	15.8	67.2	1.8	12
Cu/Zn/Ga/ZrO ₂	coprecipitation	250	n/a	75.0	10.1	13
$Cu/Zn/ZrO_2$	coprecipitation	250	19.4	29.3	n/a	14
Cu/Zn/ZrO ₂	Urea-nitrate combustion	240	17.0	56.2	n/a	15
$Cu/Zn/ZrO_2$	coprecipitation	220	21.0	68.0	5.6	16
Cu/Zn/ZrO ₂	glycine-nitrate combustion	220	12.0	71.1	n/a	17
$Cu/Zn/Al/ZrO_2$	coprecipitation	240	18.7	47.2	n/a	18
Ag/Zn/ZrO ₂	coprecipitation	220	2.0	97.0	0.46	16
$Au/Zn/ZrO_2$	coprecipitation	220	1.5	100	0.40	16
Pd/Zn/ZrO ₂	incipient wetness	250	6.3	99.6	1.1	19
Ga ₂ O ₃ -Pd/ZrO ₂	incipient wetness	250	n/a	70.0	7.9	20
n/a: not available					n/a	

diminishing the methanol yield over time.²¹ Li et al. developed Cu/ZrO₂ catalyst for CO₂ hydrogenation to methanol. It was found that the unique crystal structure of ZrO₂ induced it interact with Cu, which improved the dispersion of Cu, stabilized Cu NPs during catalytic CO₂ hydrogenation, effectively inhibited the RWGS reaction, and promoted the methanol production.²² An et al. introduced MOFs to Cu/ZrO₂ and Cu/ZnO with strong metal-support catalysts for CO₂ hydrogenation. This novel catalyst confines the agglomeration of Cu NPs and phase separation between Cu and ZnOx through utilized the porous structure of MOF, which creating a specific environment for reaction proceed, effectively enhanced methanol selectivity of 100% and high stability over 100 h.²³ Similarly, Qi et al. obtained superior methanol selectivity through introduced a bimetallic MOF template to Cu-ZnO/Al₂O₃ catalyst, in which Cu NPs have higher dispersibility, thereby inhibiting the RWGS reaction.²⁴ In addition to Cu-based catalysts, several other materials exhibit activity in CO₂ hydrogenation. Such as, Pd/ZnO catalyst loaded on multi-walled carbon nanotubes exhibit excellent performance for the formation of methanol due to an increasing concentration of active Pd⁰ species.¹⁹ Studt. *et al.* reported a Ni₅Ga₃ catalyst that reduces CO₂ to methanol at ambient pressure, which showed the same or better methanol synthesis activity, as well as considerably lower production of CO compare to the traditional Cu/ZnO/Al₂O₃ catalyst.²⁵ Besides, Ag- and Au-based catalysts offer superior selectivity of methanol with high content of metal.²⁶

Another important chemical is formic acid, which is widely used in many fields, such as the leather and rubber industries.²⁷ Formic acid has also been considered as hydrogen storage material by combining CO₂ hydrogenation with selective formic acid decomposition. The reaction formula for the CO₂ hydrogenation to formic acid is as shown in Equation (3). Currently, homogeneous catalysts are the most studied catalysts

for this reaction because homogeneous catalytic systems have milder conditions and higher catalytic efficiency. The homogeneous catalytic reaction mechanism of CO_2 hydrogenation to formic acid as shown in Figure 1.1.²⁸

$$CO_2 + H_2 \rightarrow HCOOH$$
 (3)

Wilkinson catalyst, RhCl(PPh₃)₃ was first introduced by Inoue et al. in 1976 for CO₂ hydrogenation.²⁹ Then, Ezhova *et al.* studied this complex in more details, suggesting that formic acid was formed in the presence of rhodium complexes with a phosphine ligand. Reduction of the complex to metallic rhodium is inhibited by excess PPh₃, which increases the yield of formic acid significantly.³⁰ Ru complexes have garnered significant attention, it generally offer favorable activity and selectivity for formic acid. Tai *et al.* found that there is no correlation between the basicity of monophosphines (PR₃) and the activity of the catalysts after comparing the Ru catalysts with total of 44 different phosphines or other ligands. Interestingly, a quite unique interplay of electronic and bite angle effects was observed. Diphosphines with weak basicity exhibit highly active catalysts when their bite angles are small, significantly enhancing the formic acid yield. Conversely, diphosphine with stronger basicity demonstrated the opposite trend.³¹



Figure 1.1 Homogeneous catalytic reaction mechanism of CO_2 hydrogenation to formic acid.²⁸

Syngas, also known as synthesis gas, is another chemical that can be derived from CO₂. Syngas consists of carbon monoxide (CO), CO₂, and H₂, and its applications across various industrial sectors, including fuels, synthetic materials, solvents, and minerals.³² CO₂ reforming represents one method for generating syngas by reacting CO₂ with hydrocarbons. Dry reforming of methane stands out as one of most classic examples, it is a highly endothermic process that requires a high operating temperature. In this context, developing a new hybrid catalyst capable of effectively resisting metal sintering caused by high temperature and enhancing activity is necessary. Ni-based catalysts has been widely studied in this reaction although its susceptibility to sintering at high temperature, virous methods have been conducted to modify the Ni-based catalysts, such as, alloying, controlling the Ni particle size by modifying the synthesis method and changing the type of support etc..³³⁻³⁹ The details of dry reforming of methane will be discussed in the section 1.2 and 1.3.1.

1.1.2 Enhanced oil recovery

Underneath the Earth's surface, oil and gases are enclosed by a substantial insulating layer of shale or caprock, spanning hundreds of feet. This natural barrier functions as a protective shield, preventing the ascent of fluids and gases to the surface. This also complicates the extraction of oil.¹ CO₂ enhanced oil recovery (EOR) is a widely recognized technology that has been successfully applied to extract oil from resevoirs.⁴⁰ The cycle of CO₂-enhanced oil recovery (CO₂-EOR) and the flow of CO₂ within the cycle as illustrated in Figure 1.2. CO₂-EOR involves injecting pressurized CO₂ into oil reservoirs, displacing oil and improving its flow. This mixture is extracted at production wells, separating oil and CO₂. The captured CO₂ is recirculated for reinjection, maintaining pressure and facilitating continuous oil recovery. This process enhances oil

production and has environmental benefits by utilizing and reusing CO₂.⁴⁰ Currently, two significant endeavors employing CO₂-enhanced oil recovery are being executed in offshore oil fields: Sleipner and Snøhvit, both situated in Norway. Similarly, in the United States, prior pilot projects have been undertaken to employ CO₂-enhanced oil recovery within the Gulf of Mexico.⁶ Currently, the industry facing several challenges in implementing EOR, like the non-uniform rock formation, diminishing the efficacy of CO₂ flooding. However, CO₂ EOR remains a promising method for enhanced oil and gas recovery, applicable across diverse reservoirs.⁴¹



Figure 1.2 Overview of the CO₂-EOR (enhanced oil recovery) cycle.⁶

1.1.3 Food and Beverages

The food industry continually seeks advanced extraction technologies to obtain pure natural products and produce healthy, high-quality items applicable in various industries. Regrettably, modern extraction methods employing toxic and unsafe solvents face significant limitations.⁸ Supercritical fluid extraction is a relatively new and promising technology for the "clean" extraction method. CO₂ is a great candidate for food extraction as a supercritical fluid because of its unique properties such as the low critical temperature of 31.06°C, the low critical pressure of 73.83 bar, and its critical density of 0.460 g/cm.⁴² A diverse array of food extraction processes has been effectively achieved using supercritical CO₂, including the extraction of anthocyanins from grape pomace⁴³, and grape seed oil.⁴⁴

Moreover, there has been considerable attention on the synthesis of starch using CO₂. Ma *et al.* reported a chemical-biochemical hybrid pathway for starch synthesis from carbon dioxide and hydrogen in a cell-free system. They designed and constructed a non-natural solid carbon and starch synthesis pathway comprising 11 reaction steps (Figure 1.2). The complete synthesis of starch molecules from carbon dioxide was achieved in the laboratory for the first time. Nuclear magnetic resonance revealed that the structure and composition of artificially synthesized starch molecules matched those of natural starch molecules. Initial laboratory tests indicated that the efficiency of artificial starch synthesis is approximately 8.5 times that of traditional agricultural starch production. This novel pathway makes it feasible to transition starch production from traditional agricultural cultivation to industrial manufacturing, also paving the way for the synthesis of complex molecules from CO_2 .⁴⁵



Figure 1.3 Design and modular assembly of an artificial starch anabolic pathway.⁴⁵

In summary, CO_2 utilization technologies as the current research hotspot, are promising methods to mitigate global warming and chemical compounds derived from CO_2 would be more practical and economical.

1.2 CO₂ reforming of hydrocarbon

Although the commercialization of the CO₂ reforming of hydrocarbons process has not seen significant progress, there still attracted numerous attentions and interests on it. Since their product, syngas (hydrogen and carbon monoxide), is vital in various industries such as chemicals and fuels. Dry reforming of methane (DRM) is the most widely used reaction, the involved reactions in DRM as below. The side reactions depend on the reaction conditions and feed ratio that are applied in the reaction.⁴⁶⁻⁴⁷ DRM has a high risk in carbon deposition, owing to its endothermic property that requires high reaction temperature. In brief, carbon is easier to be formed from reaction (6) at higher reaction temperature, while other three reactions (equation 7-9) tend to be more dominant at lower temperatures as these are the exothermic reaction.⁴⁷

Dry reforming of methane (DRM):

$$CO_2 + CH_4 \rightarrow 2H_2 + 2CO$$
 $\Delta H_{298K} = +247 \ kJ \cdot mol^{-1}$ (4)

Side reactions:

$$CO_2 + H_2 \to H_2O + CO \qquad \Delta H_{298K} = +41 \ kJ \cdot mol^{-1}$$
 (5)

$$CH_4 \rightarrow 2H_2 + C \qquad \qquad \Delta H_{298K} = +75 \ kJ \cdot mol^{-1} \qquad (6)$$

$$2CO \to CO_2 + C$$
 $\Delta H_{298K} = -172 \ kJ \cdot mol^{-1}$ (7)

$$CO_2 + 2H_2 \rightarrow 2H_2O + C$$
 $\Delta H_{298K} = -90 \ kJ \cdot mol^{-1}$ (8)
 $CO + H_2 \rightarrow H_2O + C$ $\Delta H_{298K} = -131 \ kJ \cdot mol^{-1}$ (9)

As alternative to CH₄, ethane,⁴⁸⁻⁵⁰ propane⁵¹⁻⁵² and butane^{49, 53-54} have also attracted many interests for CO₂ reforming due to the high availability of these hydrocarbons in shale gas and liquefied petroleum gas. The generalized equation of dry reforming of hydrocarbons is given as equation (10). Yan et al. conducted a thermodynamic analysis of the DRM, dry reforming of ethane (DRE), and dry reforming of butane (DRB) using the Gibbs free energy minimization algorithm. Their findings revealed that, under equilibrium conditions with a stoichiometric ratio, achieving a 50% conversion of CO₂ in lower temperatures for DRE and DRB compared to DRM. Specifically, for DRE, the conversion could be attained at 488 °C, and for DRB, it could be achieved at 444 °C. These temperatures were notably lower by approximately 100 °C and 150 °C, respectively, in comparison to DRM. This phenomenon can be attributed to the fact that the Gibbs free energies of DRE and DRB reach zero ($\Delta G^\circ = 0$) at lower temperatures.⁴⁹

$$nCO_2 + C_n H_{2n+2} \rightarrow (n+1)H_2 + 2nCO$$
 (10)



Figure 1.4 Typical composition of biomass gasification tars (wt%)⁵⁸

Another important CO₂ reforming reaction is benzene dry reforming. On the one hand, Gasification is a promising technique to convert sustainable biomass into a useful and combustible syngas. However, bio-tar is an unavoidable byproduct originating from the incomplete cracking of organic constituents, which not only reduce the efficiency of gasification but also cause the blockage or corrosion in the downstream devices.⁵⁵⁻⁵⁶ Benzene was selected as a model molecule due to its thermal stability and relatively high weigh percentage in the bio-tar (Figure 1.4). On the other hand, lower the CO₂ content of the produced syngas from biomass could increase its calorific value.⁵⁷ Most importantly, 1-mol of benzene can consume 6-mol of carbon dioxide, greatly enhancing CO₂ utilization. Therefore, dry reforming of benzene provides an interesting solution to reducing CO₂ and simultaneously enhancing biomass gasification efficiency.

1.3 CO₂ reforming over metallic catalysts

For the CO₂ reforming reaction, many works reported that noble metals such as Pt and Pd showed excellent activity, while the expensive price impeded its industrial application.⁵⁹⁻⁶⁰ Ni is considered as a promising candidate due to its high activity for C-H bond cleavage, selectivity to syngas and competitively low-cost. Unfortunately, coke deposition and Ni sintering are two major issues that lead to the deactivation of catalysts.⁶¹⁻⁶²

The coke type mainly including pyrolytic coke, whiskers and gum type.⁶³ Whisker carbon is the most notorious and should be avoided among them, since the formation and gasification of coke would take place at the same time during DRM, catalysts that can either inhibit the formation of coke or enhance the gasification of coke can be applied to attenuate the influence of coke on reaction activity.^{60, 64}

Sintering will occur on the catalysts surface when the affinity energy between metal atoms is higher than the constraint energy of metal-support interaction at high reaction temperature.⁶⁵⁻⁶⁶ Besides, the moisture produced from side reaction of RWGS would promote sintering.⁶⁷ Preventing small active metal particles from sintering to remain a high stable activity is essential especially for the supported catalysts.⁶⁸

1.3.1 CO₂ reforming of methane

DRM catalysts can be classified into two types, namely supported catalysts and reduced solid solution catalysts (Figure 1.5). For a supported catalyst, the active component is anchored onto the surface of a supporting material, typically it's a metal oxide that does not include the metal element in the active species. Over the last decade, there has been significant exploration on bimetallic catalysts based on Ni catalysts, single-atom catalysts and specifically structured metal catalysts for efficient and stable DRM.^{34,69-76}



Figure 1.5 illustrations of (a) a supported catalyst and (b) a solid solution catalyst before (left) and after (right) reduction.⁶⁰

Reduced solid solution catalysts differ from conventional supported catalysts in that their active species originate from the bulk of the solid solution, rather than an additional species. Similarities in crystal structure and lattice parameters of the oxides of the active metal species and the matrix components are the key for the formation of a solid solution.⁶⁰ For example, all MgO, NiO, and CoO possess a face-centered cubic (fcc) structure with closely matched lattice parameters: 4.2112 Å for MgO, 4.1684 Å for NiO, and 4.2667 Å for CoO. Consequently, a composite of MgO and NiO (or CoO) can readily create a stable solid solution (NiO–MgO or CoO–MgO) at a high temperature.⁷⁷ Besides, solid solution catalysts allows generate small-sized active metallic particles that have robust interactions with the solid solution matrix, leading to exceptional activity, remarkable resistance to carbon deposition and high stability during the DRM reaction.⁷⁸ At present, MgO-based reduced solid solution systems have been the most widely studied catalysts in DRM.⁷⁹⁻⁸⁰

1.3.2 CO₂ reforming of benzene

The catalyst of CO₂ reforming of benzene mainly focuses on Ni-based catalyst. Many strategies have been considered to activate both benzene and CO₂ molecules. Caprariis et al. reported a bimetallic NiCo catalyst promoted by Ce for benzene dry reforming, in which Ce improves the resistance to carbon deposition. In addition, the influence of two different support, γ-Al₂O₃ and activated carbon (AC), on the reaction efficiency was evaluated. As a results, the NiCo/Ce-Al₂O₃ was found to be the most active, with an enhancement of 65% and of 85% of CO and H₂ respectively during the reaction.⁵⁷ Moreover, Nam et al. developed NiFe impregnated on silicon carbide (SiC) for benzene dry reforming in a lab-scale fluidized bed reactor. A high benzene conversion rate (>90%) was observed at a higher experimental temperature (above 730°C). However, Ni exerts a significant influence on product selectivity. NiFe/SiC catalyst resulted in higher H₂ production whereas higher CO yield was produced with Fe/SiC catalyst at an elevated temperature.⁸¹ Although many researchers made an effort on CO₂ reforming of benzene, there still remains a room for improvement in the catalysts of this reaction.

1.4 Aim of this thesis

Reforming reactions may have one of greatest commercial potentials among many CO_2 conversion routes. However, it still has many challenges for the catalysts design as we discussed previously, such as carbon deposition and metal sintering. Hence, the aim of thesis focuses on developing a catalyst with high reactant activity, excellent stability and sintering resistance for CO_2 reforming of hydrocarbons. To meet these requirements, I propose the following directions on how to design and develop more efficient catalysts.

1.4.1 Catalyst design concept

Recently, Ni-based intermetallic compounds have been reported to improve catalytic performance and thermal stability. ³⁴⁻³⁵ However, the catalyst design based on binary intermetallic compounds has several limitations in atomic composition and elemental combination. Conversely, a wide range of composition ratios of solid-solution alloys can tune its electronic properties and catalytic performance.⁸² Therefore, hybridizing the advantages of these alloy materials is highly beneficial for designing more functional catalysts. Pseudo-binary alloys, represented as $((A_{1-x}A^*x)_mB_n)$, are the promising candidate for this purpose. A part of the element A in the parent intermetallic A_mB_n was substituted by a third metal A' without changing the crystal structure of A_mB_n .⁸³⁻⁸⁴ Therefore, to reduce carbon deposition while simultaneously enhancing the reactivity of reactants, a suitable combination of metals is crucial. Motivated by this concept, we designed that Ni-based pseudo-binary alloy catalyst (Ni_{0.5}CO_{0.5})₃Ge/SiO₂, in which Ni as the main active metal for C-H scission, the late 3d transition metal Co accelerate CO₂ adsorption and activation, and inert typical element Ge was considered to suppress the direct production of carbon. This combination has a sufficiently high

catalytic activity, coke resistance, and renewability for CO₂ reforming of methane (Scheme 1.1),



Scheme 1.1. The catalyst design concept for the (Ni_{0.5}Co_{0.5})₃Ge/SiO₂.

1.4.2 Fine-tuning metallic catalysts

Moreover, many other factors contributing to carbon deposition and particle agglomeration have also been taken into consideration.⁸⁵ Coking is prone to occur in the large Ni particles, since large particles typically possess lower surface energy, leading to weaker adsorption and reaction capabilities on their surface.⁸⁶ Meanwhile, if the particle agglomeration happened, the unsaturated coordination sites such as edge, corner, and vertex sites with higher catalytic activity on the Ni particles are lost, subsequently affecting their catalytic effectiveness.⁸⁷ More importantly, insufficient activation of CO₂ would break the balance between carbon formation and removal, which finally leading to the carbon deposition.^{71,88} To address these drawbacks, several strategies in this thesis have been explored as follows: (i) introducing other metals to hinder hydrocarbon complete cracking, such as alloying Ni, Co and Ge for CO₂ reforming of methane, in which Ge suppresses directly produce carbon, coking was greatly decreased compared to other corresponding binary catalysts. (ii) utilization of

redox supports to facilitate the activation of CO₂ to gasify the deposited carbon species. CeO₂ support plays an important role in our CO₂ reforming of benzene work, where the basicity and the oxygen-releasing ability of CeO₂ facilitated CO₂ adsorption and coke combustion respectively. No significant deactivation was observed within 20 h in the stability test. (iii) construction of confinement structures to immobilize Ni nanoparticles (NPs) to protect the active species from agglomeration. In the CO₂ reforming of methane, small nanoparticles with average diameter of 5.9 nm were observed on the SiO₂ support. Importantly, the STEM analysis on the spent catalyst showed that the particle size was retained even after a long-time catalytic run.

1.5 Outlines of thesis

In this research, I focused on developing a highly stable and selective alloy catalyst to enhance the CO_2 utilization in CO_2 reforming of hydrocarbons, such as dry reforming of methane, fry reforming of benzene reaction.

Chapter 2 presents a novel catalyst design based on a pseudo-binary alloy structure, $(Ni_{0.5}Co_{0.5})_3Ge/SiO_2$, in which a part of Ni atoms of intermetallic Ni₃Ge was substituted with Co without changing the parent Ni₃Ge structure, as evidenced by the combination of high-angle annular dark-field-scanning transmission electron microscopy-energydispersive system, X-ray diffraction, and X-ray absorption fine-structure analysis. $(Ni_{0.5}Co_{0.5})_3Ge/SiO_2$ exhibited a remarkably high coke resistance, an outstandingly long catalyst life (1000 h) at 700 °C even below the equilibrium conversion. Additionally, the used catalyst could be easily regenerated by a simple and soft oxidation procedure and the initial conversion in the second run was completely recovered in the third run. Alloying Ni with Ge not only inhibits carbon formation, but also promotes CO formation via the CHO intermediate pathway that involves no carbon formation. The dopant Co acts as an efficient site for CO_2 adsorption and activation, which supplies more oxygen atoms to promote carbon combustion. The combination of Ge and Co allows to minimize the coke accumulation, thus achieving outstandingly high stability for long-term operation. Therefore, $(Ni_{0.5}Co_{0.5})_3$ Ge/SiO₂ catalyst showed a sufficiently high catalytic activity, coke resistance, and renewability in CO₂ reforming of methane reaction.

Chapter 3 summarizes a binary intermetallic Ni₃Ga supported on CeO₂ works as an effective catalyst for CO₂-assisted dry reforming of benzene (DRB: $C_6H_6 + 6CO_2 \rightarrow 3H_2 + 12CO$). The combination of high-angle annular dark-field scanning transmission microscopy and X-ray absorption fine structure analysis confirmed the formation of the Ni₃Ga intermetallic structure. The CO₂ conversion of Ni₃Ga/CeO₂ catalyst was two-fold that of the corresponding monometallic catalyst, which might be attributed to the synergetic effect of higher CO₂ adsorption or activation of Ni–Ga alloy and CeO₂ support. Moreover, Ga plays a crucial role in improving catalytic activity and minimizing coke formation. Therefore, the Ni₃Ga/CeO₂ catalyst exhibited significantly enhanced CO₂ utilization ability and product yield.

1.6 Concluding remarks

In conclusion, our research not only demonstrates excellent catalytic performance of multi-metallic Ni-based catalysts in CO_2 conversion, but also offers a new perspective to prolong their lifespan. The findings of this study can promote the carbon-neutralization of industrial processes for CO_2 reforming of hydrocarbons

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Chapter 2

Development of a Highly Stable Ternary Alloy Catalyst for CO₂ Reforming of Methane

2. Development of a Highly Stable Ternary Alloy Catalyst for CO₂ Reforming of Methane

2.1 Introduction

Global warming and climate change have raised increasing attention recently. The development of related research to conduct the obligation for mitigating the emission of greenhouse gases has become a hotspot in academia and industry.¹⁻² Dry reforming of methane (DRM: CH₄ + $CO_2 \rightarrow 2H_2 + 2CO$: $\Delta H^\circ = 274$ kJ mol⁻¹) has been a promising strategy to convert two major greenhouse gases into highly valuable chemical feedstocks to realize a carbon-neutral energy cycle.³ Despite DRM's advantages in both environmental and economic situations, its commercial application still suffers from a bottleneck due to the lack of highly active and cokeresistant catalysts.⁴⁻⁵

Various catalysts have been developed to effectively address this challenge.⁶⁻⁸ The cost of precious metals makes them inefficient for use on a broad scale. Ni-based catalysts are considered as suitable alternatives because of the low cost and high C–H bond activation ability. However, Ni-based catalysts face critical drawbacks of carbon deposition and nickel sintering, which are correlated with the high reaction temperature required by the strong endothermic properties of DRM.⁹⁻¹² The carbon deposition mainly originates from complete dehydrogenation of CH₄ (CH₄ \rightarrow 2H₂ + C: $\Delta H^{\circ} = 75$ kJ mol⁻¹) at high temperatures, which leads to rapid catalyst deactivation¹³ In this context, a unique multifunctional active metal environment that minimizes coke formation while retaining sufficient methane activation ability must be developed for efficient, practical application.

Recently, Ni-based intermetallic compounds have been widely studied as active and stable catalysts for DRM.¹⁴⁻¹⁶ However, the catalyst design based on binary intermetallic compounds has several limitations in atomic composition and elemental combination. However, a wide range of composition ratios of solid-solution alloys can tune its electronic properties and

catalytic performance.¹⁷ Therefore, hybridizing the advantages of these alloy materials is highly beneficial for designing more functional catalysts. Pseudo-binary alloys, represented as $((A_{1-x}A'_x)_m B_n)$, are the promising candidate for this purpose. A part of the element A in the parent intermetallic $A_m B_n$ was substituted by a third metal A' without changing the crystal structure of $A_m B_n$.¹⁸⁻¹⁹ Here, the introduction of the third element A', provides additional functionality to the catalytic system. Additionally, the A' context x can be widely changed to optimize the promotional effect. Therefore, to minimize carbon accumulation for high thermal stability, an appropriate combination of the multimetallic system would optimize the adsorption energy of the CH_x reaction intermediates through the ligand and ensemble effects of alloving.²⁰ We developed a novel type of Ni-based pseudo-binary alloy catalyst using Ni₃Ge intermetallic compound, substituting some of Ni with Co, as directed by the above-mentioned catalyst design concept. Catalytic evaluations at 700°C exhibited an outstandingly high catalytic stability for 1000 h. Co plays an important role in inhibiting coke formation by optimizing the C-H activation ability. Additionally, the used catalyst could be easily regenerated by a simple and soft oxidation procedure and the initial conversion in the second run was completely recovered in the third run. Thus, we developed the (Ni_{0.5}Co_{0.5})₃Ge/SiO₂ catalyst for DRM, which showed a sufficiently high catalytic activity, coke resistance, and renewability.
2.2 Experimental Section

2.2.1 Catalyst preparation

Ni-Co-Ge/SiO2 and Ni-Co/SiO2 (Ni: 3 wt%) were prepared by the pore-filling coimpregnation method using Ni(NO₃)₂·6H₂O (Kanto Chemical, 98.0%), Co(NO₃)₂·6H₂O (Fujifilm Wako, 98.0%) and (NH₄₎₂GeF₆ (Furuya Metal Co. Ltd.) as metal precursors (Ni:Co:Ge = 1.5:1.5:1, Ni:Co = 1:1). A mixture of an aqueous solution of metal precursors was added dropwise to dried SiO₂ (CARiACT G-6, Fuji Silysia, specific surface area; 500 m² g⁻¹, pore volume; 0.7 mL g⁻¹, pore diameter; 6 nm) and then sealed and static overnight at room temperature so that the solutions flowed into the pores of SiO₂. Subsequently, the mixture was transferred to a round-bottom flask and quickly freeze with liquid nitrogen. The frozen sample was dried in a vacuum at -5° C. Next, the resulting powder was dried in an oven at 90°C overnight, calcined in air at 400°C for 1 h, and finally reduced by H₂ (0.1 MPa, 50 mL·min⁻¹) at 700°C for 2 h with a ramping rate of 10°C \cdot min⁻¹. Ni–Ge/SiO₂ (Ni:Ge = 3:1, Ni: 6 wt%), Co-Ge/SiO₂ (Co:Ge = 3:1, Co: 6 wt%) and Ni-Co/SiO₂ (Ni:Co = 1:1, Ni: 3 wt%) were prepared by a similar method, where the amounts of Ni and Co were adjusted to ensure the same active component amount as Ni-Co-Ge/SiO2. Monometallic Ni/SiO2, Co/SiO2 and Ge/SiO₂ were also synthesized by the same method mentioned above, in which the metal amount was comparable to those included in Ni-Co-Ge/SiO₂. Furthermore, Ni-Co-Ge/SiO₂ with different Ni/Co atomic ratios (Ni/Co = 2, Ni: 4%; Ni/Co = 0.5, Ni: 2 wt%) and other silicasupported trimetallic catalysts Ni-M-Ge/SiO₂ (M = Fe and Cu, Ni:M:Ge = 1.5:1.5:1, Ni: 3 wt%) were also prepared in a similar manner. We also tried to prepare Ni-Co-Ge/SiO₂ with high loading amount (Ni: 6 wt%) for greater activity. However, phase separation to Ni, Ni₃Ge and CoGe occurred to some extent (Figure 2.1) probably because of insufficiently homogeneous dispersion of metal precursors. Therefore, choosing an appropriate amount of metal loading is also important for high phase purity of pseudo-binary alloys.



Figure 2.1. XRD patterns of Ni–Co–Ge/SiO₂ with Ni loadings of 3 wt% and 6 wt%.

2.2.2 Characterization

The catalysts' phase formation and facet orientation were determined using XRD, equipped with a Rigaku MiniFlex II/AP diffractometer with Cu Ka radiation. The particle size distribution and the crystal structure of the catalysts were analyzed by high-angle annular darkfield scanning transmission electron microscopy (HAADF-STEM) using a JEOL JEM-ARM200 Microscope EDX analyzer. An accelerating voltage of 200 kV was used for STEM analysis. All samples were ground and sonicated in ethanol before being loaded on the Mo grid. The particle size was estimated based on more than 100 particles in high-resolution STEM images. XAFS spectra of the prepared catalysts were collected at the BL01B1 and BL14B2 beamlines of SPring-8, Japan Synchrotron Radiation Research Institute (JASRI), using Si (111) double-crystals as monochromator at room temperature. The catalyst was pelletized and pretreated with H₂ at 700°C for 0.5 h in a quartz tube to prepare the XAFS specimen, then taken to an Argon glove box ($O_2 < 0.01$ ppm) without exposure to air after cooling to room temperature with N₂ purge. The pellet and an oxygen absorber (ISO A500-HS: Fe powder) were sealed in a plastic film bag (Barrier Nylon) to check for air leakage. The obtained XAFS spectra were fitted by Athena and Artemis software ver.0.9.25 implemented in the Demeter package. FEFF8 was applied for the calculation of the back-scattering amplitude and phase shift functions²¹. Laser Raman spectroscopy experiments were carried out using an XploRA 34 PLUS spectrometer (HORIBA Scientific) at room temperature to investigate the carbonaceous deposits over the spent catalysts, with 1.4 cm⁻¹ resolution of the apparatus and a 638 nm excitation source. The amount of coke on the used catalysts after 20 h of DRM at 700°C was quantified by Temperature-programmed oxidation (TPO) using a BELCAT II (Microtrac BEL) instrument. First, 50 mg spent catalyst placed in a quartz tube was treated at 150°C for 30 min under a flow of He (30 mL·min⁻¹). After cooling to 50°C, 2% O₂/He (50 mL·min⁻¹) was passed through for baseline stabilization. Then, the catalyst was heated from 50°C to 900°C (ramping rate: 5°C·min⁻¹). An online mass spectrometer equipped downstream recorded the CO₂ amount (m/z = 44) in the outlet gas. The formation of carbon from methane and CO₂ is exothermic and downhill: CH₄ + CO₂ \rightarrow 2C + 2H₂O: $\Delta H^{\circ} = -15.2$ kJmol⁻¹, $\Delta G^{\circ} = -20.7$ kJmol⁻¹).

2.2.3 Catalytic test

The catalyst performance was examined in a fixed-bed quartz reactor with 6 mm of internal diameter at atmospheric pressure. Typically, the fresh catalyst of 100 mg was firmly fixed by quartz wool in the reactor tube. Before the DRM test, the sample was pretreated under flowing H₂ (10 mL·min⁻¹, 700°C) for 0.5 h, followed by the residual H₂ gas purged with Ar gas (20 mL·min⁻¹) for 0.5 h. Subsequently, the reactant gas consisting of CH₄:CO₂:Ar = 1:1:2 with a total gas flow rate of 40 mL·min⁻¹ (GHSV = 24,000 mL·h⁻¹·g_{cat}⁻¹) was introduced into the reactor at 700°C. Compositional analysis of the effluent gas via an online gas chromatograph (Shimadzu GC-2014s), equipped with a Shincarbon-ST column and thermal conductivity detector. A long-term stability test was performed under the same conditions. The CH₄ conversion and CO₂ conversion of TOS tests were performed below thermodynamic equilibrium conversions, which reflect the intrinsic activity of catalysts. For all catalysts, the CH₄ and CO₂ conversions were defined as the following equations:

$$CH_{4} \ conversion: X_{CH_{4}} \ (\%) = \frac{F_{CH_{4}}^{in} - F_{CH_{4}}^{out}}{F_{CH_{4}}^{in}} \times 100$$

$$CO_{2} \text{ conversion: } X_{CO_{2}} (\%) = \frac{F_{CO_{2}}^{in} - F_{CO_{2}}^{out}}{F_{CO_{2}}^{in}} \times 100$$
$$\frac{H_{2}}{CO} = \frac{F_{H_{2}}^{out}}{F_{CO}^{out}}$$
$$Carbon \text{ balance: } C_{b} = \frac{F_{CO}^{out} + F_{CH_{4}}^{out} + F_{CO_{2}}^{out}}{F_{CH_{4}}^{in} + F_{CO_{2}}^{in}} \times 100$$

where, $F_{CH_4}^{in}$, $F_{CO_2}^{in}$ and $F_{CH_4}^{out}$, $F_{CO_2}^{out}$, $F_{H_2}^{out}$, F_{CO}^{out} , indicate the inlet flow rate of CH₄, CO₂, and outlet flow rates of CH₄, CO₂, H₂, and CO, respectively.

The deactivation constant and mean catalyst life was defined as follows:

Deactivation constant:
$$k_{d} = \left\{ \ln(\frac{1 - X_{CH_{4}}^{f}}{X_{CH_{4}}^{f}}) - \ln(\frac{1 - X_{CH_{4}}^{i}}{X_{CH_{4}}^{i}}) \right\} (t^{f} - t^{i})^{-1}$$

$$Mean \ catlyst \ life: \tau = \frac{1}{k_{d}}$$

where, $X_{CH_4}^i$ and $X_{CH_4}^f$ indicate the initial (t^i : 0), and final (t^f : 1000 h) X_{CH_4} .

2.2.4 Computational details

Periodic DFT calculations were performed using the CASTEP code²² with Vanderbilt-type ultrasoft pseudopotentials and the revised version of Perdew–Burke–Ernzerhof exchange–correlation functional based on the generalized gradient approximation.²³ The plane-wave basis set was truncated at a kinetic energy of 360 eV. A Fermi smearing of 0.1 eV was used with a spin-polarization condition. Dispersion correlations were considered using the Tkatchenko–Scheffler method with a scaling coefficient of $s_R = 0.94$ and a damping parameter of d = 20.²⁴ The reciprocal space was sampled using a k-point mesh with a typical spacing of 0.04 Å⁻¹, as generated by the Monkhorst–Pack scheme.²⁵ Supercell structures were subjected to geometry optimizations using the periodic boundary conditions. The unit cell size of the bulk

Ni and Ni₃Ge crystal was first optimized. Then, for calculation of paths (A) and (B), slab structures were prepared using Ni(111)–(2×2) and Ni₃Ge(111)–(1×1) supercells (thickness of four atomic layers) with 13 Å as vacuum space on the c-axis, followed by the geometry optimization with the cell size fixed for surface relaxation. For CO₂ adsorption and activation, Ni₃Ge(111)–(2×2) supercell and its Co-doped model were used. For the doped model, a half of Ni atoms in the supercell were randomly chosen using random numbers and substituted with Co atoms to reproduce (Ni_{0.5}Co_{0.5})₃Ge(111) surface, followed by geometry optimization. The convergence criteria for structure optimization and energy calculation were set to (a) an SCF tolerance of 1.0×10^{-6} eV per atom, (b) an energy tolerance of 1.0×10^{-5} eV per atom, (c) a maximum force tolerance of $0.05 \text{ eV} \text{Å}^{-1}$, and (d) a maximum displacement tolerance of 1.0×10^{-3} Å. The net charge was set to zero for all calculations, and spin polarization was considered. A transition state search was performed based on the complete linear synchronous transit/quadratic synchronous transit method²⁶⁻²⁷ with the tolerance for all root-mean-square forces on an atom of 0.10 eV Å⁻¹.

2.3 Results and Discussion

2.3.1 Characterization of the catalysts structure

The silica-supported Ni-Co-Ge catalyst was synthesized using a pore-filling co-impregnation method. The elemental maps obtained by energy-dispersive X-ray (EDX) spectroscopy and the high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) images of Ni–Co–Ge/SiO₂ are shown in Figures 2.2a–d. The nanoparticles consisting of Ni, Co, and Ge were evenly dispersed on the SiO₂ support and constituted ternary alloy nanoparticles. Small nanoparticles with diameters ranging mainly from 3 to 8 nm (average of 5.9 nm) were observed on the SiO₂ support (Figure 2.2e). The high-resolution HAADF–STEM image of a single Ni-Co-Ge nanoparticle shown an atomic arrangement with interplanar distances of 1.83 and 1.85 Å, which were similar but slightly larger than (200) and (020) planes (1.79 Å) of intermetallic Ni₃Ge (Figure 2.2g). A possible interpretation is that the Ni sites of Ni₃Ge lattice were partially substituted with Co, whose atomic size was slightly larger than Ni (2.8% lattice expansion). The X-ray diffraction (XRD) patterns of Ni-Co-Ge/SiO2 showed slight lower-angle shifts of the 111 and 200 diffractions (Figure 2.2g), which supports the lattice expansion caused by the introduction of Co. The inset in Figure 2.2f shows the fast Fourier transform of the HAADF-STEM image, in which superlattice diffraction spots (100 and 110) were clearly observed in addition to the main diffractions of 200 and 220. The corresponding diffraction was also observed as a small peak at ca. 35.2° in XRD (Figure 2.3). These results support the formation of the Ni₃Ge-type L1₂ ordered structure. Thus, he $(Ni_{0.5}Co_{0.5})_3$ Ge pseudo-binary alloy structure was successfully formed on the SiO₂ support.

We also performed an X-ray absorption fine structure (XAFS) analysis to obtain further structural information. The X-ray absorption near-edge spectra (Figures 2.4-2.6) showed that Ni, Co, and Ge were in a metallic state and that the Ni and Ge K-edge features of Ni–Co–

Ge/SiO₂ were similar to those of Ni–Ge/SiO₂, suggesting that Co showed no electronic effect on the electronic states of Ni and Ge. Curve fitting analysis of the EXAFS oscillations (Table 2.1 and Figures 2.7–2.8) revealed that Ni–Ge and Ni–Ni scatterings in intermetallic Ni₃Ge can be distinguished by EXAFS, and the ratio of their coordination numbers (CNs) was 2.0 (CN_{Ni– Ni}: 7.2, CN_{Ni–Ge}: 3.6), which is consistent with the L1₂ Ni₃Ge structure. Similar trends were also observed for Ni–Co–Ge, where the CN_{Ni–Ni(Co)}/CN_{Ni–Ge} and CN_{Co–Co(Ni)}/CN_{Co–Ge} ratios were 2.0 (8.3/4.1 and 6.5/3.2, respectively). Besides, the sum of CN_{Ni–Ge} (4.1) and CN_{Co–Ge} (3.2) agreed with CN_{Ge–Ni(Co)} (7.7) within the error bars. These results strongly support that Co atoms occupy the Ni sites of Ni₃Ge but not the Ge sites, demonstrating the pseudo-binary alloy structure. Considering that CN_{Co–Co(Ni)}(6.5) is lower than CN_{Ni–Ni(Co)} (8.3), the distribution of Co atoms in the Ni sites may be slightly biased to the surface region. Furthermore, the Ni–Ge interatomic distance in the Ni–Co–Ge/SiO₂ catalyst (2.65 Å) was slightly longer than that of Ni₃Ge/SiO₂ (2.57 Å), which is consistent with the lattice expansion due to the Co doping. This excludes the possibility that Ni–Co–Ge/SiO₂ comprises a physical mixture of intermetallic Ni₃Ge and Co₃Ge nanoparticles.



Figure 2.2. Characterization of Ni–Co–Ge/SiO₂ catalyst. **a** HAADF–STEM image of Ni–Co–Ge/SiO₂ (Ni:Co:Ge=1.5:1.5:1), **b-d** corresponding elemental maps for (**b**) Ni, (**c**) Co, and (**d**) Ge, acquired by EDX. **e** Particle size distribution. **f** HAADF–STEM image of a single nanoparticle on Ni–Co–Ge/SiO₂. **g** XRD patterns of Ni–Co–Ge/SiO₂ and Ni–Ge/SiO₂ catalysts. **h** Model of the (Ni_{1-x}Co_x)₃Ge pseudo-binary alloy structure with the crystal structure unit.



Figure 2.3. XRD patterns of Ni–Co–Ge/SiO₂ and Ni–Ge/SiO₂ catalysts.



Figure 2.4. (a) Ni, (b) Co and (c) Ge K-edge X-ray absorption near edge structure (XANES) spectra of the Ni–Co–Ge/SiO₂ and Ni–Ge/SiO₂ catalysts and reference compounds.



Figure 2.5. (a) Ni, (b) Co and (c) Ge K-edge extended X-ray absorption fine structure (EXAFS) spectra of the Ni–Co–Ge/SiO₂ and Ni–Ge/SiO₂ catalysts and reference compound.



Figure 2.6. Fourier-transforms of EXAFS in (a) Ni, (b) Co and (c) Ge K-edge of the Ni–Co–Ge/SiO₂ and Ni–Ge/SiO₂ catalysts and reference compounds.



Figure 2.7. Curve-fitting results of Ni K and Ge K-edge k^3 -weighted EXAFS of Ni–Ge/SiO₂. Solid and dashed lines indicate the results of simulation and experiment, respectively.



Figure 2.8. Curve-fitting results of Ni K, Co K and Ge K-edge k^3 -weighted EXAFS of Ni– Co–Ge/SiO₂. Solid and dashed lines indicate the results of simulation and experiment, respectively. The k^3 -weighted EXAFS oscillation was Fourier-transformed in the *k* range of 3-14 Å⁻¹, 3-11 Å⁻¹ and 3-12 Å⁻¹ for Ni K, Co K and Ge K-edge separately. Curve-fitting was performed using the back Fourier-transforms of the coordination peaks ranging between 1.0–3.1 Å, 1.0–3.0 Å, and 1.3–3.0 Å for Ni K, Co K and Ge K-edge, respectively.

Table 2.1. Results of EXAFS curve-fitting analysis for Ni–Co–Ge/SiO₂ and Ni–Ge/SiO₂ catalysts.

Sample	Edge	Shell	S_o^2	CN	r (Å)	$\Delta E_{\rm o}~({\rm eV})$	$\sigma^{2}(\text{\AA}^{2})$ R-factor
		Co–Co		12 (fix)	2.49 ± 0.0014		0.007
Co foil	Co K	Со–Со	0.84	6 (fix)	$3.52 \hspace{0.2cm} \pm \hspace{0.2cm} 0.013$	$7.4 \hspace{0.2cm} \pm \hspace{0.2cm} 0.8$	0.010 0.016
		Со-Со		24 (fix)	$4.36 \hspace{0.1in} \pm \hspace{0.1in} 0.008$		0.011
		Ni–Ni		12 (fix)	$2.48 \hspace{0.2cm} \pm \hspace{0.2cm} 0.004$		0.007
Ni foil	Ni K	Ni–Ni	0.95	6 (fix)	$3.50 \hspace{0.1 in} \pm \hspace{0.1 in} 0.014$	$7.03~\pm~0.8$	0.010 0.019
		Ni–Ni		24 (fix)	$4.33 \hspace{.1in} \pm \hspace{.1in} 0.008$		0.009
GeO ₂		Ge–O		4 (fix)	1.74 ± 0.005		0.003
	Ge K	Ge–Ge	1.04	4 (fix)	$3.16 \hspace{0.1in} \pm \hspace{0.1in} 0.007$	5.3 ± 1.6	0.004 0.006
		Ge–O		4 (fix)	$3.34 \hspace{.1in} \pm \hspace{.1in} 0.02$		0.001
	Ni K	Ni–Ni	0.05	7.2 ± 0.9	$2.51 \hspace{.1in} \pm \hspace{.1in} 0.006$	10 + 17	0.006
Ni–Ge/SiO		Ni–Ge	0.95	$3.6 \hspace{0.1in} \pm \hspace{0.1in} 0.9$	$2.57 \hspace{0.1in} \pm \hspace{0.1in} 0.116$	1.0 ± 1.7	0.003
	Cak	Ge–Ni	1.04	$8.5 \hspace{0.2cm} \pm \hspace{0.2cm} 0.5$	$2.51 \hspace{.1in} \pm \hspace{.1in} 0.004$	84 + 07	0.009
	UCK	Ge–O	1.04	1.5 ± 1.4	1.78 ± 0.04	8.4 ± 0.7	0.005
		Ni–Ni/Co	.	$8.3 \hspace{0.2cm} \pm \hspace{0.2cm} 0.9$	$2.52 \hspace{0.1 in} \pm \hspace{0.1 in} 0.009$		0.007
	N1 K	Ni-Ge	0.95	4.1 ± 0.9	$2.65 \hspace{0.1 cm} \pm \hspace{0.1 cm} 0.05$	2.5 ± 1.6	0.011 0.014
Ni-Co-Ge/SiO,	C a V	Co–Co/Ni	0.94	$6.5 \hspace{0.2cm} \pm \hspace{0.2cm} 0.6$	$2.56 \hspace{0.1in} \pm \hspace{0.1in} 0.02$	26 ± 14	0.004
2	COK	Co–Ge	0.84	3.2 ± 0.6	$2.43 \hspace{.1in} \pm \hspace{.1in} 0.04$	3.0 ± 1.4	0.002
	Ge K	Ge–Ni (Co)	1.04	7.7 ± 0.7	2.51 ± 0.006	$8.5~\pm~1.0$	0.009 0.008



Figure 2.9. XRD patterns of Ni₃Ge/SiO₂, Co₃Ge/SiO₂, and Ni_{0.5}Co_{0.5}/SiO₂ catalysts.

2.3.2 Catalytic performance

Next, we tested the prepared (Ni_{0.5}Co_{0.5})₃Ge/SiO₂ catalyst in DRM at 700°C and compared it with the control catalysts (Ni₃Ge/SiO₂, Co₃Ge/SiO₂, and Ni_{0.5}Co_{0.5}/SiO₂; see Figure 2.9 for their XRD patterns). Note that the mole of Ni, Co, or Ni + Co was equalized for each catalyst (Table 2.2). The time course of CH₄ and CO₂ conversions is shown in Figures. 2.10a–b (see Figure 2.11 for details of equilibrium conversion). Although Ni_{0.5}Co_{0.5} showed high initial conversions of CH₄ and CO₂ (72%–80%), it was rapidly deactivated within 5 h due to coke accumulation. High conversion and good stability were observed in Ni₃Ge, indicating that alloying with Ge is necessary for high stability. Co₃Ge was much less active than the Ni-based catalysts, which could be ascribed to Co's lower C–H activation ability than Ni.²⁸ For at least 140 h, (Ni_{0.5}Co_{0.5})₃Ge showed high initial conversions and retained them with negligible

deactivation (CH₄: 67%–62%, CO₂ 75%–71%). Thus, the Co doping further promoted the catalyst stability of Ni₃Ge in DRM. CO₂ conversion was always higher than CH₄ conversion, and the H₂/CO ratio was approximately 0.8 (Figure 2.12), which indicates the occurrence of reverse water-gas shift reaction (RWGS: $CO_2 + H_2 \rightarrow CO + H_2O$) to some extent. We also tested other Ni/Co ratios of $(Ni_{1-x}Co_x)_3$ Ge/SiO₂ (see Figure 2.13 for XRD patterns) and Fe- or Cu-doping instead of Co (Figure 2.14 and Figure 2.15, respectively). Deviating from the equimolar ratio of Ni:Co resulted in slightly lower conversions, indicating the optimum Ni/Co ratio of unity. Ni-Cu-Ge and Ni-Fe-Ge showed much lower catalytic activity than Ni-Co-Ge, which is probably due to their lower ability for C-H activation and shows the validity of Co as the third metal. Thereafter, monometallic Ni/SiO₂, Co/SiO₂ and Ge/SiO₂ catalysts were also tested for the DRM reaction as control experiments (Figure 2.16). Although Ni/SiO₂ showed high initial conversion for both CH₄ and CO₂ conversion, it rapidly deactivated within 70 h. Co/SiO₂ retained the low CH₄ conversion (15%), which is probably due to the weak C-H activation ability of Co metal. Ge/SiO₂ was almost inactive for DRM reaction. We conducted further control experiments using physical mixtures of the bimetallic and monometallic catalysts (Ni/SiO₂+Co-Ge/SiO₂, Ni-Ge/SiO₂+Co/SiO₂ and Ni-Co/SiO₂ + Ge/SiO₂). The activity and stability of these mixtures were obviously lower than those of Ni-Co-Ge/SiO₂ (Figure 2.17), demonstrating that the ternary alloy structure is essential for the outstanding catalytic performance.

Then, we investigated the long-term stability of the developed catalyst at 700°C for 1000 h, which is also an important factor for the industrial application of DRM. Although a slight decrease in conversion was observed, approximately 70%–80% of the initial conversions of CH₄ and CO₂ were retained even at 1000 h on stream (Figure 2.10c, see Figure 2.18 for H₂/CO ratio and carbon balance), where the deactivation constant k_d was 0.00077 h⁻¹. The reciprocal deactivation constant ($\tau = k_d^{-1}$), which is often used as a scale of stability, was 1300 h (Figure 4.17).

2.10d). Although some moderately high τ values (400–800 h) have been reported, they were estimated from experiments



Figure 2.10. Catalytic performance of Ni-based catalysts in DRM. Reaction conditions: catalyst amount of 100 mg, CH₄:CO₂:Ar = 10:10:20 mL·min⁻¹, 700°C, 1 bar. **a**, **b** Time course of **(a)** CH₄ Conversions and **(b)** CO₂ Conversions. **c** Long-term stability of CH₄ conversion and CO₂ conversion over (Ni_{0.5}CO_{0.5})₃Ge/SiO₂. **d** Comparison of the stability with reported Nibased DRM catalysts under equilibrium conversion. Numbers correspond to the entries in Table 2.3. **e** Regeneration test using (Ni_{0.5}CO_{0.5})₃Ge/SiO₂ during three catalytic cycles. Regeneration conditions: CO₂:Ar = 10:10 mL·min⁻¹ for 7 h, followed by H₂ of 10 mL·min⁻¹ for 1 h at 700°C. with much shorter periods (<100 h, Figure 2.10d; see Table 2.3 and Figure 2.19 for details; note that the *x*-axis is shown in a log scale for clarity). However, we obtained much higher τ values from an actual long catalytic run. For a fair comparison, Figure 2.10d does not include reported systems in which the CH₄ conversion reached (or exceeded) the thermodynamic equilibrium and/or increased during the reaction. This is because the deactivation trend cannot 46

be accurately evaluated when the conversion is saturated or when some induction period is included, i.e., there is a possibility that catalyst deactivation has not been reflected in the conversion trend due to the use of an excess catalyst or due to some structural change (see Table 2.4 and Table 2.5 and Figure 2.20 for the full comparison with those not included in Figure 2.10d and Table 2.3).

We also examined the reusability of (Ni_{0.5}Co_{0.5})₃Ge/SiO₂ after a simple regeneration process. Generally, deposited carbon on deactivated catalysts is combusted in the presence of oxygen or air in a regeneration process. In this study, however, CO₂ was used as a soft oxidant to combust carbon via the reverse Boudouard reaction (CO₂ + C \rightarrow 2CO). Another merit for easier handling of this is that switching the catalytic run to the regeneration process only requires shutting off the supply of methane in the feed gas at the same temperature. Before the second catalytic run, the catalyst was reduced by flowing H₂ reduction for 1 h at the same temperature after feeding CO₂/Ar for 7 h. The initial conversion of CH₄ and CO₂ were only slightly lower than those in the first runs (Figure 2.10e). They reached a steady-state after 1170 h, probably because CO₂ could combust not all the carbonaceous species at that temperature. The initial conversions in the second run were fully recovered in the third catalytic run after the second regeneration process, demonstrating the continuous reusability of (Ni_{0.5}Co_{0.5})₃Ge/SiO₂. A similar trend was also observed in the H₂/CO ratio change and carbon balance (Figure 2.21). The full recovery of the initial activity after regeneration and the repeated reusability is very rare in reported systems for DRM (Tables 2.3-2.5). Thus, the (Ni_{0.5}Co_{0.5})₃Ge catalyst displayed multifunctional functionalities of high catalytic activity, outstanding long-term stability, and repeated reusability. The STEM analysis on the spent (Ni_{0.5}Co_{0.5})₃Ge catalyst showed that the particle size was retained even after a long time catalytic run (5.9 nm \rightarrow 5.9 nm, Figure 2.22), which evidenced the high thermal stability of the (Ni_{0.5}Co_{0.5})₃Ge pseudo-binary alloy structure. ²⁹ Moreover, we performed a stability test

using $(Ni_{0.5}Co_{0.5})_3$ Ge under a neat condition $(CH_4:CO_2 = 10:10 \text{ mL min}^{-1})$ as a more praction condition. No deactivation was observed up to 200 h (Figure 2.23), demonstrates the outstanding durability for practical application.

Catalyst	metal loa	ding (wt%)	Active metal mole amount / mmol
Cuturyst	Ni	Со	(per 100 mg catalyst)
(Ni _{0.5} Co _{0.5}) ₃ Ge/SiO ₂	3	3	0.1020
Ni ₃ Ge/SiO ₂	6	0	0.1022
Co ₃ Ge/SiO ₂	6	0	0.1019
Ni _{0.5} Co _{0.5} /SiO ₂	3	3	0.1020

Table 2.2. The active metal loading of the catalysts.



Figure 2.11. Calculated equilibrium conversion of CH_4 and CO_2 in DRM ($CH_4:CO_2:Ar = 1:1:2$) as a function of temperature. Calculation was finished by a Gibbs free energy minimization algorithm implemented in HSC Chemistry 8.



Figure 2.12. (a) Product H₂/CO ratio, (b) Carbon balance of (Ni_{0.5}Co_{0.5})₃Ge/SiO₂ and other binary catalysts.



Figure 2.13. (a) XRD patterns of $(Ni_{1-x}Co_x)_3Ge/SiO_2$ (x = 0, 0.33, 0.5, and 0.67) and (b) their magnification of the region designated as a dotted square. Vertical dashed lines indicate the center of these diffractions. $(Ni_{1-x}Co_x)_3Ge/SiO_2$ showed diffraction peaks similar to that of 49

Ni₃Ge/SiO₂ with a small feature of 110 diffraction, which indicates the formation. Moreover, the peak position of the 200 diffraction slightly shifted in the order of x = 0 > 0.33 > 0.5 > 0.67. These results support that pseudo-binary alloys with various Ni:Co ratio were successfully formed with high phase purities.



Figure 2.14. Catalytic activity of different Ni/Co ratio in $(Ni_xCo_y)_3Ge/SiO_2$. (a) CH₄ conversion, (b) CO₂ conversion, (c) H₂/CO ratio, and (d) carbon balance. Reaction conditions: catalyst amount of 100 mg, CH₄:CO₂:Ar = 10:10:20 mL·min⁻¹, 700 °C, 1 bar.



Figure 2.15. Catalytic performance of Ni–M–Ge/SiO₂ (M = Co, Cu and Fe) catalysts. (a) CH₄ conversion, (b) CO₂ conversion, (c) H₂/CO ratio, and (d) carbon balance. Reaction conditions: catalyst amount of 100 mg, CH₄:CO₂:Ar = 10:10:20 mL·min⁻¹, 700 °C, 1 bar.



Figure 2.16. Catalytic activity of monometallic Ni/SiO₂, Co/SiO₂ and Ge/SiO₂ in DRM. (a) CH₄ conversion, (b) CO₂ conversion, (c) H₂/CO ratio, and (d) carbon balance. Reaction conditions: catalyst amount of 100 mg, CH₄:CO₂:Ar = 10:10:20 mL·min⁻¹, 700 °C, 1 bar.



Figure 2.17. Catalytic performance of physical mixtures of Ni/SiO₂ + Co–Ge/SiO₂, Ni–Ge/SiO₂ + Co/SiO₂, and Ni–Co/SiO₂ + Ge/SiO₂ compared with that of Ni–Co–Ge/SiO₂ in DRM. (a) CH₄ conversion, (b) CO₂ conversion (c) H₂/CO ratio and (d) carbon balance.



Figure 2.18. Long-term stability test of (a) H₂/CO ratio and (b) carbon balance over $(Ni_{0.5}Co_{0.5})_3$ Ge/SiO₂ (Ni: 3 wt%). Reaction conditions: catalyst amount of 100 mg, CH₄:CO₂:Ar = 10:10:20 mL·min⁻¹, 700 °C, 1 bar.

No	Cataluat	Tem	GHSV	Gas feed	Equili	ibrium	Stabil	CH ₄	CO ₂ Conv.	$k_d(h^{-1})$:	τ (h):	Regene	Dof
NO.	Catalyst	р. (°С)	L g ⁻¹ h ⁻¹	ratio	CH ₄	CO ₂	ity	(%)	(%)	CH_4	CH_4	rability	Ker.
1	Ni ₃ Ga/MgO	600	54	CH ₄ :CO ₂ :N ₂ =1:1:1	45	58	72	50-46	52-48	0.002	442.8	_	15
2	Ni/MgO-Al ₂ O ₃	600	30	CH ₄ :CO ₂ :N ₂ =1:1:3	53	64	300	30-17	40-21	0.002	405.0	_	30
3	NiCo/Al ₂ O ₃	600	144	CH ₄ :CO ₂ :N ₂ =1:1:1	45	58	24	34-31	-	0.006	175.4	_	31
4	Na-Ni/ZrO ₂	600	120	CH ₄ :CO ₂ :Ar = 68 : 31 :1	26	68	24	17-15	51-43	0.007	147.7	_	32
5	Ni/Ce _{0.9} Eu _{0.1} O _{1.95}	600	60	CH ₄ :CO ₂ :N ₂ =1:1:2	50	62	12	24-22	36-33	0.009	106.2	-	33
6	Ni/La ₂ O ₃	650	240	CH ₄ :CO ₂ :N ₂ =15:15:70	73	81	50	65-62	71-68	0.003	386.1	-	34
7	Ni ₃ Fe ₁ Cu ₁ - Mg _x Al _y O _z	650	120	CH ₄ :CO ₂ :N ₂ =12:12:1	58	70	20	28-26	33-32	0.005	197.0	-	35
8	Ni@SBA-15	650	960	CH ₄ :CO ₂ :A _r =0.5:0.5:9	86	90	12	45-43	45-42	0.007	147.8	-	36
9	NiFe/Al ₂ O ₃	650	600	CH ₄ :CO ₂ :N ₂ =1:1:2	66	76	24	41-38	-	0.005	191.1	+	37
10	(Ni _{0.5} Co _{0.5}) ₃ Ge /SiO ₂	700	24	CH4:CO2 =1:1	80	87	1000	60-41	71-54	0.00077	1299.7	+	This work
11	Ni–Ce–Al oxides	700	36	CH ₄ :CO ₂ =1:1	72	82	80	70-67	-	0.002	546.3	-	38
12	NiCe@SBA-16	700	45	CH ₄ :CO ₂ :He =1:1:1	76	85	100	72-68	77-69	0.002	524.4	-	39
13	Ni/CeO ₂ -SiO ₂	700	48	CH4:CO2:Ar= 1:1:2	80	87	50	78-76	88-87	0.002	442.5	-	40
14	Ni/La ₂ O ₃	700	60	CH ₄ :CO ₂ :N ₂ =15:15:70	84	90	50	74-70	82-75	0.004	251.7	-	41
15	Ni/MgO	700	-	CH ₄ :CO ₂ :A _r = 1:1:8	100	100	500	99-93	95-96	0.004	249.0	-	12
16	NiMnMg/Al ₂ O ₃	700	12	CH4:CO2 =1:1	72	82	20	66-64	70-69	0.004	227.5	-	42
17	Ni/CeO ₂ /Al ₂ O ₃	700	90	CH ₄ :CO ₂ :N ₂ =1:1:1	78	85	60	55-40	62-43	0.010	99.0	_	43
18	Ni/CeO ₂ -ZrO ₂	700	30	CH4:CO2 =1:1	72	82	24	51-38	52	0.022	45.3	-	44
19	Ni-TiN/SBA-15	700	1.558	CH4:CO2 =1:1	72	82	12	72-70	77-73	0.041	38.4	+	45
20	NiSn/Al ₂ O ₃	700	60	CH ₄ :CO ₂ :N ₂ =1:1:6	100	100	20	42-30	75-65	0.035	28.6	_	46

Table 2.3. Summary of the catalytic performance of Ni–Co–Ge/SiO₂ and other reported Nibased catalysts for DRM under equilibrium conversion.

21	NiCo/γ–Al ₂ O ₃ – HY	700	-	CH ₄ :CO ₂ =1:1	72	82	3	61-58	65-65	0.045	22.1	_	47
22	NiLa/ZrO ₂	700	24	CH4:CO2 =1:1	72	82	5	70-59	74-70	0.097	10.3	_	48
23	NiCo/CeO ₂	700	-	CH ₄ :CO ₂ :A _r = 1:1:3	82	88	5	78-65	79-72	0.129	7.7	_	49
24	Ni/Ca-HA1_S	700	15.8	CH ₄ :CO ₂ :N ₂ =1:1:3	82	88	30	70-46	76-57	0.034	29.8	+	50
25	Ni/h-BN	750	60	CH4:CO2:N2 =2:2:1	85	92	40	73-72	81-80	0.001	797.4	_	51
26	2Ni/HAP–Ce	750	60	CH ₄ :CO ₂ :He =1:1:3	90	94	100	90-84	97-94	0.005	183.7	_	6
27	NiMg/Al ₂ O ₃	750	12	CH ₄ :CO ₂ :A _r = 1:1:3	90	94	50	90-87	95-93	0.006	168.8	_	52
28	Ni/Ce-La(Sm) - Cu-O	750	30	CH4:CO2:He =2:2:1	85	92	100	75-61	76-68	0.007	153.5	_	53
29	Ni nanotube $@(T_{2.5}A_{1.5})m$	750	144	CH ₄ :CO ₂ :N ₂ =1:1:2	88	93	60	85-79	89-82	0.007	146.5	_	54
30	Ni@SiO ₂	750	48	CH4:CO2:N2 =1:1:2	88	93	24.5	60-54	73-66	0.011	91.8	_	55
31	Ni/MgFe _x Al _{2-x} O ₄	750	-	CH ₄ :CO ₂ :Ar =1:1:3	90	94	12	48-44	-	0.035	28.4	+	56
32	NiCoCe/Al ₂ O ₃	750	24	CH4:CO2=1:1	83	90	5	68-37	-	0.257	3.9	+	57
33	Ni/ZrO ₂	750	24	CH4:CO2=1:1	83	90	36	83-76	92-90	0.012	80.8	+	58
34	Ni/CeO ₂ -Al ₂ O ₃	800	-	CH ₄ :CO ₂ =1:1	90	95	72	79-69	89-81	0.008	129.6	_	59
35	NiCa/MCM-41	800	143	CH4:CO2:Ar= 1:1:1	92	96	20	62-57	50-43	0.011	91.5	_	60
36	NiGa/MCM-41	800	39	CH ₄ :CO ₂ :N ₂ =30:30:5	91	95	30	88-82	91-86	0.016	63.0	_	16
37	NiW/SiO ₂	800	96	CH ₄ :CO ₂ =1:1	90	95	30	70-59	76-69	0.016	62.1	_	61
38	Ni/Al ₂ O ₃ -La ₂ O ₃	800	180	CH4:CO2:N2 =1:1:1	92	96	8	75-69	86-80	0.037	26.8	_	62
39	Ni/Al ₂ O ₃ -CeO ₂	800	180	CH ₄ :CO ₂ :N ₂ =1:1:1	92	96	8	90-84	93-89	0.067	14.8	_	62
40	Ni/MgAlO _x	800	1440	CH ₄ :CO ₂ :Ar =8:10:7	98	88	20	52-48	-	0.008	118.7	+	63
41	AlSB/MgO@Ni	800	97 000 h ⁻¹	CH ₄ :CO ₂ :N ₂ =3:6:1	99	69	27	35-9	33-7.8	0.068	14.6	Δ	64
42	NiCeLa/MgO- Al ₂ O ₃	800	1700 h ⁻¹	CH4:CO2 =1:1	90	95	76	95-94	92-85	0.003	394.0	Δ	65

- (no data) Δ (not recovered or stable) + (recovered & stable)



Figure 2.19. Catalytic performance of $(Ni_{0.5}Co_{0.5})_3$ Ge/SiO₂ and reported Ni-based systems for DRM under equilibrium conversion. (a) Mean catalyst life ($\tau = k_d^{-1}$), (b) Stability time.

	-			-		•									
		Tem	GHSV		equilit	orium	oper	CH4	CO2	equil					
No.	Catalyst	p.	(F/M)	Feed gas	Conv.	. (%)	ation	Conv	Conv	ibriu	kd (h-1)	τ(h)	Regener	Ref.	
		(°C)	L g ⁻¹ h ⁻¹	ratio	CH ₄	С	time	time . (%)		m	(CH ₄)	(CH ₄)	ability		
			e			O_2	(h)								
1	(Ni _{0.5} Co _{0.5}) ₃ Ge	700	24	CH ₄ :CO ₂ :	80	87	1000	60-	71-	< 60	0.00077	1299.7	+	this	
1	$/SiO_2$	700	24	A _r =1:1:2	00	07	1000	41	54	< cq	0.00077	1277.1	Т	work	
2	NT: /-:1:	(00	(0)	CH4:CO2:	10	50	100	54-	58-	0	0.00280	2565		66	
2	2 INI/SIIICA 60	000	60	He=1:1:1	40	38	100	47	50	w eq	0.00280	550.5	-	00	
				CH ₄ :CO ₂ :				83-	82-	~	0.00.00	a= 1		(7	
3	LaSrNiCuO	650	-	N ₂ =1:1:1	62	74	24	67	57	@ eq	0.03656	27.4	-	07	
				CH ₄ :CO ₂ =				71-	75-						
4	Co/Ni/Al ₂ O ₃	650	12	1:1	58	70	72	63	66	@ eq	0.00504	198.3	-	68	
	NiCu /SBA-		10000 h-	CH ₄ :CO ₂ =				75-	77-						
5	15	650	1	1:1	58	70	6	74	76	@ eq	0.01053	95.0	-	69	
				$CH_4:CO_2=$				72-	82-						
6	6 Ni/MCM-41 700	700	45	1:1	72	82	200	68	78	@ eq	0.00095	1048.8	-	70	
				CH4:CO2:											
NiCoSr/γ-	NiCoSr/γ-	700	3.6	$N_{2}=17.17$	73	83	7.5	85-	82.3-	@ ea	0.02128	47.0	_	71	
,	Al ₂ O ₃	100	010	2	, 0	00	110	83	80.6	ંગ્ય	0102120	1710			
	Ni/Ce-SBA-			- CH ₄ ·CO ₂ =				77-	77-						
8	15	700	36	1.1	72	82	40	73	74	@ eq	0.00594	168.5	-	72	
	10			CHCO				82-	88-						
9	Ni-Ce/W-Zr	700	42	N ₂ -3·3·1	74	83	100	72	83	@ eq	0.00578	173.1	-	73	
				CH.:CO.:				91 ₋	85-						
10	Ni/silica	750	60	$U_{2}=1.1.1$	86	92	100	91- 80	82	@ eq	0.00223	448.6	-	66	
				$CU_{1}CO_{1}$				09	05						
11	NiMgAlCe	750	15	LH4:CO2=	83	90	20	00- 06	95-	@ eq	0.00886	112.9	-	74	
								00	95						
12	NiCo/CeO ₂	800	12	CH4:CO2:	93	96	10	99-	99-	@ eq	0.32088	3.1	-	75	
				$N_2 = 3:3:4$				80	81						
13	NiCe/SiO ₂	800	12	$CH_4:CO_2=$	90	95	30	91.3-	95.6-	@ eq	0.00717	139.4	-	76	
				1:1				89.5	94.6						
14	In0.5Ni@SiO2	800	18	$CH_4:CO_2=$	90	95	430	90-	95-	@ eq	0.00048	2099.7	-	14	
				1:1				88	92						
15	Ni-yolk@Ni	800	40	CH ₄ :CO ₂ :	92	96	90	96-	97-	@ eq	0.01208	82.8	-	77	
asiO ₂	@SiO ₂	800	000	-	N ₂ =1:1:1	12	70	20	89	95	- 1	y 0.01200		-	

Table 2.4. Comparison of catalytic performance in DRM with those of reported Ni-basedcatalysts obtained at thermodynamic equilibrium and showing a positive k_d value.



Figure 2.20 Comparison of the stability with those of reported Ni-based DRM catalysts reach/exceed equilibrium conversion. Numbers correspond to the entries in Table S4.

No.	Catalyst	Temp.	GHSV (F/M) L g ⁻¹ h ⁻¹	Feed gas	equilibrium Conv. (%)		total time	CH4 Conv.	CO ₂ Conv.	equili-	kd (h ⁻¹)	Regene	Ref
		(°C)		ratio	CH ₄	$\rm CO_2$	(h)	(%)	(%)	brium	(CH4)	rability	
1	(Ni _{0.5} Co _{0.5}) ₃ G e /SiO ₂	700	24	CH4:CO2: Ar=1:1:2	80	87	1000	60-41	71-54	< eq	0.00077	+	this work
2	NiCu/Mg(Al) O	600	60	CH4:CO2: N2=1:1:2	50	62	25	50-51	58-57	@ eq	< 0	-	7
3	Ni _{0.2} Co0.3@ S-2	650	100	CH4:CO2: N2=2:2:1	60	82	100	-	-	-	-	-	10
4	Ni/Al ₂ O ₃	700	22000 h-	CH4:CO2 =1:1	72	82	9	91-98	70-50	@ eq	< 0	-	78
5	NiCoCu/Y- Al ₂ O ₃	700	12	CH4:CO2 =1:1	72	82	5	25-30	35-43	< eq	< 0	-	79
6	NiMo/ZSM-5	750	50	CH4:CO2 =1:1	83	90	20	-	-	-	-	-	80
7	Ni/ZrO2@BN	750	25	CH4:CO2 =1:1	83	90	200	68-71	77-80	< eq	< 0	-	81
8	NiCo/SiO ₂	750	60	CH4:CO2: He=1:1:1	86	92	100	84-84	87-87	@ eq	0	-	82
9	Ni–Mg–Al LDH	750	60	CH4:CO ₂ : N ₂ = 1:1:2	88	93	30	86-86	92-92	@ eq	0	-	83
10	NiCo	750	110	CH4:CO2: N2=1:1:1	86	92	2000	90-90	-	@ eq	0	-	84

Table 2.5. Comparison of catalytic performance in DRM with those of reported Ni-based catalysts showing a negative k_d value and/or no deactivation at thermodynamic equilibrium.

	Ni–Co/MgO–			CH4:CO2:		92	40				a <0	4	95
11	ZrO_2	750	126	N ₂ =1:1:1	86	92	40	75-80	80-84	< eq	< 0	+	85
12	NFO/McO	800	20	CH4:CO2	00	05	5	85.02	00.04	< 0.0	< 0		86
12 NIO/MgO	NIO/MgO	800	20	=1:1	90	95	5	85-95	90-94	< eq	< 0	-	
12	Ni/MgO-	800	20	CH4:CO2:	04	97	200	71.80	72.81	< eq	< 0		30
15	Al ₂ O ₃	000	20	N ₂ =1:1:3	24		500	/1-00	72-01	्प्प	< 0	-	
14 NiMo/MgO	NiMo/MgO	800	60	CH4:CO2:	07	97 98	850	100-	100-	@ #0	0		87
	Nilvio/NigO	800	00	He=1:1:8	21		850	100	100	æcq	0	-	
15	NiMo/SBA-	800	4	CH4:CO2	90	95	600	94 94	-	@ #0	0		88
15	15	000	4	=1:1				94-94	-	æcq	0	-	
	NiCoMnCe/Z		243000	CH4:CO2:			24	100-	95 5-		0		
16	rO	800	h-1	O ₂ =1: 0.8:	96	88		100	95.5	@ eq		-	89
	102			0.2				100	75.5				
17	NiCoMgAl	800	32000	CH4/CO2/	94	97	20	96-96	92.5-	@ ea	0	_	90
1,	oxide	800	h-1	Ar=1:1:3	74	,,	20	90 90	92.5	હાવ	0		
	LaNio 24C00 22			CH4:CO2:									
18	Mno 22O2	800	12	N_2	94	94	14	94-94	89-90	@ eq	0	-	91
				= 1:1.05:1									



Figure 2.21. Time course of (a) H_2 /CO ratio and (b) carbon balance over ($Ni_{0.5}Co_{0.5}$)₃Ge/SiO₂ for DRM with catalyst regeneration.



Figure 2.22. (a) STEM image and (b) particle size distribution of spent (Ni_{0.5}Co_{0.5})₃Ge/SiO₂ after 88 h catalytic run.



Figure 2.23. Long-term stability of (a) CH₄ conversion, (b) CO₂ conversion over $(Ni_{0.5}Co_{0.5})_3$ Ge/SiO₂ at neat condition. Reaction condition: catalyst amount of 100 mg, CH₄:CO₂ = 10:10 mL min⁻¹, 700 °C, 1 bar.

2.3.3 The origin of the coke resistance of (Ni_{0.5}Co_{0.5})₃Ge/SiO₂ in DRM

Next, we conducted mechanistic studies to understand the activity trend and the origin of the excellent stability of (Ni_{0.5}Co_{0.5})₃Ge/SiO₂. First, we investigated the behavior of coke on

the spent catalyst. After 20 h of the catalytic run, the spent catalysts were analyzed by temperature-programmed oxidation (O₂-TPO, Figure 2.24a) and Raman spectroscopy. The spent Ni₃Ge showed an intense evolution of CO₂ at 744°C, which is assignable to combustion of graphitic carbon. On the contrary, $(Ni_{0.5}Co_{0.5})_3$ Ge showed little coke combustion, which reflects the significantly improved coke resistance by Co-doping and is consistent with the much greater stability of $(Ni_{0.5}Co_{0.5})_3$ Ge in DRM. The Raman analysis revealed that the spent Ni₃Ge showed the two peaks assignable to G and D band of coke with the intensity ratio of approximately 0.5 (Figure 2.25). The ratio lower than unity suggests the formation of only soft coke;²⁴ therefore, the growth of hard coke has been prevented due to the ensemble effect of Ge. The accumulation of soft coke is also consistent with gradual deactivation. Conversely, no signal was observed for (Ni_{0.5}Co_{0.5})₃Ge, indicating that even the formation of soft coke was inhibited by Co doping. These are also consistent with the results of O₂-TPO and the stability trend. Besides, the XRD patterns (peak position and width) and the XAFS spectra (XANES and EXAFS features) of the two catalysts did not little change after the long-term DRM reaction (Figure 2.26 and Figure 2.27), confirming that the alloy and Co-doped structures were retained even after the reaction and deactivation. These results indicate that the deactivation of Ni₃Ge/SiO₂ was induced simply by coke accumulation on the catalyst, but not by sintering or segregation of the alloy phase. Then, we determined the reaction orders with respect to the partial pressures of CH₄ and CO₂ (P_{CH4} and P_{CO2} , respectively), revealing the first-order dependence on P_{CH4} , while much lower reaction orders were obtained for P_{CO2} (Figure 2.24b) and 3c). This result indicates that the C-H activation of CH₄ is the rate-determining step. For $P_{\rm CO2}$, Ni₃Ge showed a negative reaction order of -0.3, suggesting that CO₂ adsorption on Ni sites competes with CH₄ adsorption. Conversely, (Ni_{0.5}Co_{0.5})₃Ge showed a positive order of 0.4, which implies that the competitive adsorption was eliminated by Co doping. A possible interpretation of this change is that the doped Co acts as an effective CO₂ adsorption site. This

may also have enhanced CO₂ activation, thus serving more oxygen atoms for facile coke combustion. Apparent activation energies (E_a) in DRM were also estimated for each catalyst using Arrhenius-type plots (Figure 2.28). E_a of CH₄ activation on Ni₃Ge was 51 kJ mol⁻¹, whereas that on Co₃Ge was 108 kJ mol⁻¹, which indicates the intrinsic lower ability of Co than Ni for C–H activation and is consistent finely with the experimental activity trend. (Ni_{0.5}Co_{0.5})₃Ge/SiO₂ showed an intermediate E_a of 89 kJ mol⁻¹, suggesting that the C–H activating ability was moderately tuned.

Based on the experimental results, the effects of alloying/doping on the enhanced coke tolerance may be summarized as follows: (1) alloying with Ge significantly suppresses coke formation (Ni₃Ge vs. Ni and Ni_{0.5}Co_{0.5}) and (2) Co-doping enhances coke removal by accelerating CO₂ adsorption/activation and the resulting oxygen supply.^{68, 75} Figure 2.24d shows the reaction pathway of DRM, in which three reaction pathways from CH₄ to CO have been known, i.e., those through the (A) CHO, (B) C, and (C) COH intermediates. Since only the pass (A) does not form carbon atoms among them, facilitating this step should suppress coke formation and accumulation. Besides, enhancing oxygen supply further accelerates path (A) and (B), thus suppressing coke accumulation.



Figure 2.24. Mechanistic study. (a) O_2 -TPD profiles for Ni₃Ge/SiO₂ and (Ni_{0.5}Co_{0.5})₃Ge/SiO₂ used in DRM for 20 h. Reaction orders on the partial pressure of (b) CH₄ and (c) CO₂. (d) Reaction routes from methane and CO₂ to CO in DRM. (e) DFT-calculated E_a of CH decomposition and CH+O reactions over Ni₃Ge(211) step sites and the Co-doped analogs. The inset figures depict the transition states (gray: Ni, pink: Co, green: Ge, light gray: C, red: O, and white: H).

2.3.4 DFT calculation

Finally, we performed density functional theory (DFT) calculations to understand more deeply the roles of Ge and Co on the reaction pathways (A) and (B). The C–H scission of CH to form C and H atoms (CH \rightarrow C + H) and the subsequent oxygenation to form CO (C + O \rightarrow

CO) were considered on Ni(111) and Ni₃Ge(111) surfaces for path (B) (Figure 2.24e, f). Conversely, oxygenation of CH to form CHO (CH + $O \rightarrow$ CHO) followed by the H elimination to CO (CHO \rightarrow CO + H) were considered for path (A). The C-H scission of CH had much higher energy barriers than other steps, indicating that path (A) is minor in this reaction system. This suggests that although carbon formation is a rare event, its growth for coke accumulation eventually deactivates the catalyst. –When comparing Ni(111) and Ni₃Ge(111), the latter showed higher barrier of the C-H scission of CH, whereas lower barriers of path (A), indicating that Ge suppresses path (B), while facilitates path (A). Therefore, carbon formation is further prevented by alloying with Ge. Then, we focused on the role of Co for CO₂ adsorption and activation. We randomly replaced half of Ni atoms in the Ni₃Ge(111) slab to construct (Ni_{0.5}Co_{0.5})₃Ge(111) surface. Since this surface has various metal-metal ensemble sites (i.e., $Ni_x Co_y Ge_z$, x, y, z = 0-2), several possible adsorption/activation configurations were considered (Figure 2.24g, h; 1–6, see Figure 2.29 and Figure 2.30 for details). Bidentate CO_2 with an sp^2 like bended structure was considered as an adsorbate $(CO_2(a))$ because this has been known as a chemisorbed species on transition metal surfaces.^{18,92} The order of adsorption energy was as follows: Ni₃Ge (Ni-C-O-Ni) ~ 1,2 (Co-C-O-Ni) > 5,6 (Ni-C-O-Co) >> 3,4 (Co-C-O-Co). Notably, the adsorption with Co-O bonding resulted in lower adsorption energy, demonstrating that Co acts as an efficient adsorption site of CO₂, probably due to the intrinsic oxophilicity of Co. For the subsequent C–O cleavage, the reactions occurring at the Co sites provided much lower energy barriers and transition state (TS) energies (Figure 2.24g; 3, 4, and 5). Therefore, the doped Co not only stabilizes CO₂ adsorption, but also promotes its activation process to release oxygen atoms. For the optimized catalyst, the EXAFS analysis suggested that the surface was slightly Co-rich. Therefore, the probability of oxygen supply from CO₂ for coke removal seems to be also high. Thus, our DFT calculation well supported the reaction

mechanism derived from the experimental suggestions and clarified the roles of Ge and Co alloying on catalysis.



Figure 2.25. Raman spectra of the spent Ni_3Ge/SiO_2 and $(Ni_{0.5}Co_{0.5})_3Ge/SiO_2$ used in DRM for 20 h.



Figure 2.26. XRD patterns of the fresh and spent (700°C, 75 h) over (a) Ni_3Ge/SiO_2 , (b) $(Ni_{0.5}Co_{0.5})_3Ge/SiO_2$ catalyst and (c) the comparison of spent Ni_3Ge/SiO_2 and $(Ni_{0.5}Co_{0.5})_3Ge/SiO_2$ catalysts. The particles size was calculated by Scherrer formula.



Figure 2.27. (a, e) Ni K- and (b, f) Ge K-edge k^3 -weighted raw EXAFS oscillations for fresh and spent (a, b) (Ni_{0.5}Co_{0.5})₃Ge/SiO₂ and (e, f) Ni₃Ge/SiO₂ catalysts. Fourier transforms of the Ni (c, g) K- and (d, h) Ge K-edge k^3 -weighted EXAFS spectra of fresh and spent (c, d) (Ni_{0.5}Co_{0.5})₃Ge/SiO₂ and (g, h) Ni₃Ge/SiO₂ catalysts.



Figure 2.28. Arrhenius-type plots obtained in DRM over Ni₃Ge/SiO₂, Co₃Ge/SiO₂, and (Ni_{0.5}Co_{0.5})₃Ge/SiO₂.



Figure 2.29. DFT-calculated structures for CO_2 adsorption and activation on $Ni_3Ge(111)$ surface.



Figure 2.30. DFT-calculated structures for CO_2 adsorption and activation on $(Ni_{0.5}Co_{0.5})_3Ge(111)$ surface. Structures of configuration 3 shown in Fig. 3h are shown as representative.

2.4 Conclusion

In this study, we designed and synthesized a (Ni_{0.5}Co_{0.5})₃Ge/SiO₂ ternary catalyst highly efficient for DRM. Substituting a part of Ni in the intermetallic Ni₃Ge with Co formed a pseudo-binary alloy structure of (Ni_{0.5}Co_{0.5})₃Ge, which was confirmed by the combination of XRD, XAFS, and HAADF-STEM-EDX analyses. The (Ni_{0.5}Co_{0.5})₃Ge/SiO₂ catalyst exhibited a remarkably high coke resistance at 700°C and an outstandingly long catalyst life (operation: 1000 h, $\tau = 1300$ h), as well as excellent renewability and reusability with almost full recovery of the initial catalytic performance. Notably, the outstandingly high catalytic performance can be achieved at conversions below the thermodynamic equilibrium for the first time. Alloying Ni with Ge not only inhibits carbon formation, i.e., the origin of coke accumulation, but also promotes CO formation via the CHO intermediate without forming carbon. The dopant Co acts as an efficient site for CO₂ adsorption and activation, which supplies more oxygen atoms to promote carbon combustion. The combination of Ge and Co allows to minimize the coke accumulation, thus achieving outstandingly high stability for long-term operation. The results obtained in this study provide a highly efficient catalyst for converting CO₂ and natural gas and propose an effective catalyst design concept for durable and coke-resistant catalysis for hydrocarbon chemistry.
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Chapter 3

Enhanced CO₂ Utilization in Dry reforming of Benzene over Intermetallic Ni₃Ga Catalyst

3. Enhanced CO₂ Utilization in Dry reforming of Benzene over Intermetallic Ni₃Ga Catalyst

3.1 Introduction

The environmental problems caused by carbon dioxide (CO₂) emissions are a major concern in the scientific and industrial fields. The chemical utilization of CO₂ as an abundant and nontoxic C1 block can contribute to the control of excess CO₂ emissions while generating valuable feedstocks to alleviate the problem of fossil fuel depletion.¹⁻³ A promising method for large-scale CO₂ utilization is dry reforming with hydrocarbons owing to the relatively energyfavorable requirement.⁴ Dry reforming of methane (DRM) has been regarded as a representative route among dry reforming reactions, however, the stoichiometric ratio of CH₄ and CO₂ makes the consumption of CO₂ very limited.

The dry reforming of benzene (DRB; $C_6H_6 + 6CO_2 \rightarrow 3H_2 + 12CO$) has significant potential for reducing CO₂ to the barest minimum. This process uses up to 6 mol of CO₂ per mol of benzene according to the stoichiometric ratio. Aromatic compounds including benzene can be obtained from the degradation of bioderived lignin. Therefore, DRB has a huge potential for carbon neutralization. A reaction temperature above 700°C is required because of the high endothermicity of DRB.⁵⁻⁷ However, there are certain drawbacks at high temperatures, such as side reactions (benzene cracking: e.g., $C_6H_6 \rightarrow 6C + 6H$ and reverse water gas reaction: CO_2 $+ H_2 \rightarrow CO + H_2O$) and poor stability due to coke formation and metal sintering, which hamper the practical application of DRB. In addition, a high-temperature process is energy-intensive.⁸ Therefore, an appropriate catalyst for DRB reactions that functions in a relatively lowtemperature region is highly attractive.

Supported Ni catalysts are promising for the reforming reactions of hydrocarbons owing to their strong ability to break C–C or C–H bonds and their relatively low price.⁹⁻¹¹ Nevertheless,

it is well-known that coke deposition on Ni catalysts severely hinders their catalytic performance. Many strategies have been employed to improve their coke resistance, among which alloying is considered one of the most effective routes.¹²⁻¹⁴ Ni-based intermetallic compounds have been widely utilized for DRM and the steam reforming of benzene reactions, in which Ni–Ni ensembles were normally diluted by catalytically inactive metals, such as early transition and typical metals, to form the intermetallic phase.¹⁵⁻¹⁶ Moreover, the specific ordered atomic arrangement of Ni intermetallic compounds can alter the electronic and geometric structure of Ni and the adsorption behaviors of the reactants, leading to high catalytic activity and selectivity.^{15, 17}

Herein, we report a binary Ni₃Ga intermetallic compound that functions as an effective catalyst for the DRB at 500°C. The Ni₃Ga/CeO₂ catalyst exhibited significantly enhanced CO₂ utilization and long-term stability compared with the corresponding monometallic Ni catalyst (Scheme 3.1). Moreover, alloying with Ga modified the CO₂ activation ability and improved the carbon deposition resistance, resulting in high stability. This study provides insight into Nibased catalysts for CO₂ utilization.



Scheme 3.1 The catalyst design concept for the Ni₃Ga/CeO₂.

3.2 Experimental Methods

3.2.1 Catalyst preparation

 Ni_3Ga/CeO_2 and Ni/CeO_2 (Ni = 3 wt%) were prepared using a conventional impregnation method with Ni(NO₃)₂·6H₂O (Kanto Chemical, 98.0%) and Ga(NO₃)₂·nH₂O (n = 7-9, Wako, 99.9%) as metal precursors (Ni:Ga = 3:1). The CeO₂ support (JRC-CEO-2, SBET = 123.1 m^2g^{-1}) was added to a vigorously stirred aqueous solution (50 mL of H₂O per gram of CeO₂) containing Ni and Ga followed by stirring for 3 h at 30°C. The mixture was dried under a reduced pressure at 40°C using a rotary evaporator, then the sample was calcined under flowing air at 500°C for 1 h and reduced under flowing H₂ (50 mL/min) at 500°C for 1 h at a ramping rate of 10°C min⁻¹. Different Ni/Ga ratios and a monometallic Ga catalyst (a Ga content of 1.19 wt% to ensure the same loading as that in Ni₃Ga) supported on CeO₂ and Ni₃Ga/Al₂O₃ catalysts (the Al₂O₃ was prepared through the calcination of boehmite (γ -AlOOH, supplied by Sasol chemicals) at 900°C for 3 h, y-phase) were prepared using the aforementioned method. In addition, Ni₃Ga/SiO₂ was prepared using a pore-filling co-impregnation method. A mixture of an aqueous solution of metal precursors with a known amount of water was added dropwise to dried SiO₂ (CARiACT G-6, Fuji Silysia, $S_{BET} = 673 \text{ m}^2 \cdot \text{g}^{-1}$). The mixture was sealed and under static overnight at room temperature so that the solutions flowed into the SiO₂ pores. Subsequently, the mixture was transferred to a round-bottom flask and rapidly frozen with liquid nitrogen. The frozen sample was dried in a vacuum at -5° C. The resulting powder was dried overnight in an oven at 90°C, under the same aforementioned calcination and reduction conditions.

3.2.2 Characterization

The crystallite phases of the prepared catalysts were determined by XRD using a Rigaku MiniFlex II/AP diffractometer with Cu Kα radiation. The XAFS spectra of the prepared

catalysts were recorded at the BL01B1 and BL14B2 beamlines of SPring-8, The Japan Synchrotron Radiation Research Institute (JASRI), using Si (111) double crystals as monochromators at room temperature. The catalyst was pelletized and pretreated with H₂ at 700°C for 0.5 h in a quartz tube to prepare the XAFS specimen. Afterward, it was transferred into an Ar glove box ($O_2 < 0.01$ ppm) without exposure to air after cooling to room temperature with N₂ purge. The pellet and an oxygen absorber (ISO A500-HS: Fe powder) were sealed in a plastic film bag (Barrier Nylon) to prevent air leakage. The obtained XAFS spectra were fitted using the Athena and Artemis software (version 0.9.25) implemented in the Demeter package. FEFF8 was applied for the calculation of the back-scattering amplitude and phase shift functions. A JEOL JEM-ARM200 M microscope equipped with an energy-dispersive Xray analyzer was used for the HAADF-STEM analysis. The coke amount on the spent catalysts was analyzed by TPO using a BELCAT II (Microtrac BEL) instrument. First, the spent catalysts (0.5 g of catalyst and quartz sand) in a quartz tube were treated at 150°C for 30 min under a flow of He (30 mL \cdot min⁻¹). After cooling to 50°C, 2% O₂/He (50 mL \cdot min⁻¹) was passed through for baseline stabilization. Thereafter, the catalyst was heated from 50°C to 900°C (ramping rate of 5°C·min⁻¹). An online mass spectrometer equipped downstream recorded the CO₂ amount (m/z = 44) in the outlet gas.

3.2.3 Catalytic test

DRB was conducted in a fixed-bed quartz reactor with an internal diameter of 6 mm under atmospheric pressure. The fresh catalyst (0.1 g) diluted with quartz sand (0.9 g) was firmly fixed in the reactor tube using quartz wool. Prior to the DRB test, the sample was pretreated under flowing H₂ (10 mL·min⁻¹, 500°C) for 0.5 h, followed by residual H₂ gas purged with He gas (20 mL·min⁻¹) for 0.5 h. Subsequently, the reactant gas comprising C₆H₆:CO₂:He = 2.4:14.4:20 with a total gas flow rate of 36.8 mL·min⁻¹ was introduced into the reactor at 500°C. Benzene was supplied using a bubbler at 35°C with He as a carrier gas (10 mL·min⁻¹). The flow rate of benzene was calculated based on the saturated vapor pressure at 35°C. The flow rate of CO₂ was determined to be 6 times higher than that of benzene for stoichiometric conversion. CO₂ was supplied with He carrier gas (10 mL·min⁻¹). The detailed setup of the reactor is shown in Figure 3.1. The compositional analysis of the effluent gas was conducted using an online gas chromatograph (Shimadzu GC-2014s), equipped with a Porapak Q column and thermal conductivity detector. For all the catalysts, the C₆H₆ and CO₂ conversions were defined using the following equations:

$$C_{6}H_{6} \text{ conversion: } X_{C_{6}H_{6}} (\%) = \frac{F_{C_{6}H_{6}}^{in} - F_{C_{6}H_{6}}^{out}}{F_{C_{6}H_{6}}^{in}} \times 100$$

$$CO_{2} \text{ conversion: } X_{CO_{2}} (\%) = \frac{F_{CO_{2}}^{in} - F_{CO_{2}}^{out}}{F_{CO_{2}}^{in}} \times 100$$

$$CO \text{ Yield: } = \frac{F_{CO}^{out}}{6F_{C_{6}H_{6}}^{in} + F_{CO_{2}}^{in}} \times 100$$

Carbon balance:
$$C_b = \frac{F_{CO}^{out} + 6F_{C_6H_6}^{out} + F_{CO_2}^{out}}{6F_{C_6H_6}^{in} + F_{CO_2}^{in}} \times 100$$

where, $F_{C_6H_6}^{in} F_{CO_2}^{in}$ and $F_{C_6H_6}^{out} F_{CO_2}^{out}$, F_{CO}^{out} , indicate the inlet flow rate of C₆H₆, CO₂, and outlet flow rates of C₆H₆, CO₂ and CO₂ respectively.



Figure 3.1. The detailed setup of the reactor for dry reforming of benzene. The thermocouple for the water bath was the actual temperature of benzene. The flow rate of benzene was calculated based on the saturated vapor pressure at 35°C using the bubble system.

3.3 Results and Discussion

3.3.1 Structure Characterization of the catalysts

The Ni₃Ga/CeO₂ catalyst was prepared using a conventional co-impregnation method. CeO₂ was used as a catalyst support suitable for CO₂ adsorption and the subsequent activation.¹⁸⁻¹⁹ The crystal structure of the Ni₃Ga catalyst was examined by X-ray diffraction (XRD). When SiO₂ was used as a support for Ni₃Ga, two intense peaks were observed at 43.7° and 50.9°, corresponding to the 111 and 200 diffractions of intermetallic Ni₃Ga, respectively (Figure 3.2a). However, no corresponding diffraction peaks were observed for Ni₃Ga/CeO₂. A similar phenomenon is frequently observed for CeO2-supported metallic catalysts, mainly due to the strong X-ray scattering by Ce that has a larger atomic weight and scattering coefficient than Si, significantly weakening the X-ray reflection from the Ni₃Ga nanoparticles.¹⁸⁻¹⁹ Therefore, Xray adsorption fine structure (XAFS) analysis was performed to further investigate the structure of Ni₃Ga/CeO₂. The Ni K- and Ga K-edge X-ray absorption near-edge structure (XANES) spectra of Ni₃Ga/CeO₂ were similar to those of the corresponding reference foils, indicating that these metals were reduced to zero-valent states (Figures. 3.2b and 3.2c). Curve-fitting analysis was performed on the extended XAFS (EXAFS; Table 3.1, Figs. 3.3-3.5 and Table 3.2) oscillations and revealed that the Ni-Ga and Ni-Ni scatterings of Ni₃Ga can be distinguished by EXAFS. The ratio of the coordination numbers was approximately 2.0 (CN_{Ni}- $N_i = 7.9$, $CN_{N_i-Ga} = 4.0$), which correlated with the theoretical value of intermetallic Ni₃Ga (2.0, $CN_{Ni-Ni} = 8.0$, $CN_{Ni-Ga} = 4.0$). Therefore, these results supported the formation of intermetallic Ni₃Ga on the CeO₂ support. We also performed high-angle annular dark-field scanning transmission microscopy (HAADF-STEM) analysis on Ni₃Ga/CeO₂. As shown in Figure 3.6, metallic nanoparticles were not visible in the HAADF-STEM image. This is most likely due to insufficient Z-contrast between Ce and the other metals. However, the elemental maps of Ni and Ga clearly indicated the presence of small ($2 \sim 3$ nm) bimetallic nanoparticles on the CeO₂

support. The atomic ratio of Ni:Ga in these nanoparticles were approximately 3:1 (Figure 3.7), which was consistent finely with the formation of nanoparticulate intermetallic Ni₃Ga.



Figure 3.2. (a) XRD patterns of Ni₃Ga supported by SiO₂ and CeO₂. (b), (c) Ni K- and Ga Kedge XANES spectra of the in situ reduced catalysts (solid lines) and reference foils (dashed lines). (d) Model of the Ni₃Ga intermetallic structure with the crystal structure unit.

Shell	CN	r (Å)	$\Delta E_{\rm o}~({\rm eV})$	σ^2 (Å ²)	R-factor	
Ni–Ni	12 (fix)	2.48 ± 0.00	6.5 ± 0.2	0.006	0.000	
Ni–Ni	7.9 ± 0.4	2.48 ± 0.00	75108	0.006	0.001	
Ni–Ga	4.0 ± 0.4	$2.57 {\pm} 0.04$	7.5±0.8	0.015		
Ga—Ni	5.4 ± 1.0	2.51 ± 0.01	6 4+1 0	0.009	0.018	
Ga–O	$1.4{\pm}0.9$	1.86 ± 0.02	0.4±1.7	0.004		

Table 3.1. Results of EXAFS curve fitting analysis of the Ni₃Ga/CeO₂ catalyst.



Figure 3.3. (a) Ni and (b) Ga K-edge extended X-ray absorption fine structure (EXAFS) spectra of the Ni₃Ga/CeO₂ and Ni / CeO₂ catalysts and reference compound.



Figure 3.4. Fourier-transforms of EXAFS in (a) Ni, (b) Ga K-edge of the Ni₃Ga/CeO₂ and Ni/CeO₂ catalysts and reference compounds.



Figure 3.5. Curve-fitting results of Ni K and Ga K-edge k^3 -weighted EXAFS of Ni₃Ga/CeO₂. Solid and dashed lines indicate the results of simulation and experiment, respectively.

Sampl	Edge	Shell	${S_o}^2$	CN	r (Å)	$\Delta E_{\rm o}~({\rm eV})$	$\sigma^2({\rm \AA}^2)$	R-
e								factor
Ni foil	Ni K	Ni–Ni	0.81	12 (fix)	2.48 ± 0.00	6.5 ± 0.2	0.006	0.000
Ga ₂ O ₃		Ga-O	1.13	3 (fix)	1.91 ± 0.02	9.3±2.6	0.002	0.014
	Ga K	Ga-O		1 (fix)	2.10 ± 0.03		0.002	
		Ga-Ga		2 (fix)	3.06 ± 0.01		0.001	
Ni ₃ Ga/ CeO ₂	Ni K	Ni-Ni	0.81	7.9 ± 0.4	2.48 ± 0.00	7.5±0.8	0.006	0.001
		Ni-Ga		4.0 ± 0.4	2.57 ± 0.04		0.015	
	Ga K	Ga-Ni	1.13	5.4 ± 1.0	2.51 ± 0.01	6.4±1.9	0.009	0.018
		Ga-O		1.4 ± 0.9	1.86 ± 0.02		0.004	

Table 3.2. Results of EXAFS curve-fitting analysis for Ni₃Ga /CeO₂ catalyst.



Figure 3.6. (a) HAADF-STEM image and the corresponding elemental maps of Ni₃Ga/CeO₂:(b) Ni+Ga, (c) Ni, and (d) Ga.



Figure 3.7. (a) HAADF-STEM image and the corresponding elemental maps of Ni₃Ga/CeO₂: (b) Ni, (c) Ga, (d) Ce, (e) Ni+Ga. (f) Atomic ratios of Ni and Ga in the regions designated by yellow squares in (e), which agreed roughly with 3:1 (75:25).

3.3.2 Catalytic performance in DRB

Next, the catalytic performance of Ni₃Ga was tested in a DRB reaction at 500°C and compared with that of the monometallic Ni catalyst. Although the Ni₃Ga and Ni catalysts exhibited similar benzene conversions, a large difference was observed in the CO₂ conversion and CO yield, which were markedly enhanced by alloying Ni with Ga (Figure 3.8 and Figure 3.9 for the time course). Moreover, the carbon balance was close to unity (Figure 3.9d), indicating that the selectivity of Ni₃Ga toward undetected products was negligible. Notably, benzene and CO₂ were stoichiometrically converted over intermetallic Ni₃Ga. This reflected the considerably higher selectivity of the intermetallic Ni₃Ga catalyst for dry reforming and CO₂ utilization in contrast with the monometallic Ni catalyst. The stoichiometric conversion of benzene and CO₂ is necessary for effective CO₂ utilization and for preventing coke accumulation due to excess benzene cracking. For Ni/CeO₂, the CO yield was slightly higher than the CO₂ conversion, probably because a part of the accumulated carbon derived from the benzene cracking was oxidized to CO by the lattice oxygen of CeO₂. We also performed control experiments using physical mixtures of Ni/CeO₂ and Ga/CeO₂ and the latter alone (Figure 3.10), which demonstrated a considerably low performance and negligible activity, respectively. This indicates that alloying Ni with Ga is essential for an enhanced catalytic performance. Subsequently, we examined a series of catalysts $Ni_{1-x}Ga_x/CeO_2$ with different Ni:Ga molar ratio (x=0.25, 0.5, 0.6) in DRB reactions in a similar manner as previously conducted (Figure 3.11a and Figure 3.12). The Ga content of the Ni:Ga alloy drastically affected the catalytic performance. The catalyst with a Ni:Ga ratio of 1:1 exhibited a benzene conversion of 10%, whereas its CO_2 conversion was extremely low (<2%). This might have been because excess Ga largely weakened the CO₂ activation ability of Ni. The CO yield was also slightly higher than the CO₂ conversion, probably because the lattice oxygen of CeO₂ may contribute to a part of CO production as we explained earlier.¹⁸ A further increase in the Ga content (Ni:Ga = 2:3)

resulted in a lower benzene conversion, indicating that an appropriate size of Ni–Ni ensembles was necessary for the adsorption/activation of benzene. Next, the support effect was studied using Al_2O_3 and SiO_2 instead of CeO₂ (Figure 3.11b and Figure 3.13 for the time course). The catalytic performance of Ni₃Ga/CeO₂ was significantly higher than that using other supports. Notably, the CO₂ conversion of Ni₃Ga/Al₂O₃ was only half of that of Ni₃Ga/CeO₂, whereas that of Ni₃Ga/SiO₂ was even close to zero. This is probably because the basicity and the oxygen-releasing ability of CeO₂ facilitated CO₂ adsorption (activity) and coke combustion (stability) respectively.¹⁸⁻¹⁹ We also performed a long-term stability test using the Ni₃Ga/CeO₂ catalyst, in which no significant deactivation was observed at least for 20 h (Figure 3.14).



Figure 3.8. Catalytic performance of Ni₃Ga/CeO₂ and Ni/CeO₂ in the DRB. Reaction conditions: catalyst amount, 100 mg; flow rate, C_6H_6 :CO₂:He = 2.4:14.4:20 mLmin⁻¹; temperature, 500°C. Values at a time-on-stream of 125 min are shown.



Figure 3.9. (a) C_6H_6 conversion, (b) CO_2 conversion (c) Product CO yield, (d) Carbon balance of Ni₃Ga/CeO₂ and Ni/CeO₂.



Figure 3.10. Catalytic performance of physical mixtures of Ni/CeO₂ + Ga/CeO₂ and monometallic Ga/CeO₂ in DRB. (a) C_6H_6 conversion, (b) CO₂ conversion (c) CO yield and (d) carbon balance.



Figure 3.11. Catalytic performance of (a) $Ni_{1-x}Ga_x/CeO_2$ with different Ni:Ga ratios and (b) Ni_3Ga on different supports (CeO₂, Al₂O₃, and SiO₂) in the DRB (values at a time-on-stream of 95 min and 125 min are shown, respectively).



Figure 3.12. Catalytic activity of different Ni/Ga molar ratio in Ni_{1-x}Ga_x/CeO₂ (x = 0.25, 0.5, 0.6). (a) C₆H₆ conversion, (b) CO₂ conversion (c) CO yield and (d) carbon balance.



Figure 3.13. (a) C_6H_6 conversion, (b) CO_2 conversion (c) CO yield and (d) carbon balance in DRB on the Ni₃Ga catalysts using various support.



Figure 3.14. Stability test using Ni₃Ga/CeO₂ in benzene dry reforming: (a) benzene and (b) CO₂ conversion. Reaction condition: catalyst amount, 200 mg; flow rate, C_6H_6 :CO₂:He = 2.4:14.4:20 mLmin⁻¹; temperature, 500°C.

To clarify the role of Ga, the coke behavior of spent catalysts was analyzed by temperature-programmed oxidation (O₂-TPO; Fig. 3.15). The spent Ni/CeO₂ catalyst exhibited an intense evolution of CO₂ at approximately 600°C, which was assignable to the combustion of graphitic carbon.²⁰ Contrarily, the Ni₃Ga/CeO₂ catalyst exhibited minimal coke combustion around 350°C, which was ascribed to the facile combustion of amorphous carbon and/or the partial oxidation of Ga species in Ni₃Ga. Overall, these findings indicated that the formation of the intermetallic Ni₃Ga phase significantly inhibited coke formation and accumulation, which was consistent with the high stability of the DRB (Figure 3.14).



Figure 3.15. O₂-TPO profiles of spent Ni₃Ga/CeO₂ and Ni/CeO₂ in the DRB.

3.4 Conclusion

In summary, we prepared a Ni₃Ga/CeO₂ catalyst that is effective for the DRB. The XAFS analysis revealed that the intermetallic Ni₃Ga phase was formed on the CeO₂ support. Alloying Ni with Ga inhibited coke accumulation and significantly enhanced CO₂ utilization. Notably, the CO₂ conversion of Ni₃Ga was two-fold that of the monometallic Ni catalyst, which might be attributed to the synergetic effect of higher CO₂ adsorption or activation of Ni–Ga alloy and CeO₂ support. The study findings provide insight into material design in the CO₂ utilization of hydrocarbon conversion.

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Chapter 4

General Conclusions
4 General Conclusions

In this research, I targeted the effective utilization of CO_2 by dry reforming of hydrocarbons based on multimetallic catalysts. Alloying Ni and other metals remarkably enhanced the reactant conversion and stability in various dry reforming reactions, such as dry reforming of methane and dry reforming of benzene. Moreover, the structure of catalysts and mechanism study were further explored by characterization technique and experimental analysis.

Chapter 2 concludes that an innovative catalyst design concept based on a pseudo-binary alloy structure enables the development of a highly efficient and outstandingly stable catalyst for CO₂ reforming of methane. (Ni_{0.5}Co_{0.5})₃Ge/SiO₂, in which a part of Ni atoms of intermetallic Ni₃Ge was substituted with Co without changing the parent Ni₃Ge structure, exhibited a remarkably high coke resistance, an outstandingly long catalyst life (1000 h) at 700 °C even below the equilibrium conversion (many previous studies were done at equilibrium conversions; therefore, there remains a possibility that deactivation may have not appeared in conversion due to excess activity). Additionally, the used catalyst could be easily regenerated by a simple and soft oxidation procedure and the initial conversion in the second run was completely recovered in the third run. Alloying Ni with Ge not only inhibits carbon formation, but also promotes CO formation via the CHO intermediate pathway that involves no carbon formation. The dopant Co acts as an efficient site for CO₂ adsorption and activation, which supplies more oxygen atoms to promote carbon combustion. The combination of Ge and Co allows to minimize the coke accumulation, thus achieving outstandingly high stability for long-term operation. Therefore, (Ni_{0.5}Co_{0.5})₃Ge/SiO₂ catalyst showed a sufficiently high catalytic activity, coke resistance, and renewability in CO₂ reforming of methane reaction.

Chapter 3 summarizes a binary intermetallic Ni₃Ga supported on CeO₂ works as an effective catalyst for CO₂-assisted dry reforming of benzene (DRB: $C_6H_6 + 6CO_2 \rightarrow 3H_2 +$

12CO), which has great potential for the purpose of reducing as much CO₂ as possible. The CO₂ reforming of methane consumes a limited amount of CO₂ because of its stoichiometric ratio, whereas the CO₂ reforming of benzene could consume up to 6 mol of CO₂ with only 1 mol of benzene. Moreover, aromatic compounds including benzene can be supplied from the degradation of bio-derived lignin. The Ni₃Ga/CeO₂ catalyst exhibited significantly enhanced CO₂ utilization ability and product yield. Notably, the CO₂ conversion was two-fold that of the corresponding monometallic catalyst, which might be attributed to the synergetic effect of higher CO₂ adsorption or activation of Ni–Ga alloy and CeO₂ support. The combination of high-angle annular dark-field scanning transmission microscopy and X-ray absorption fine structure analysis confirmed the formation of the Ni₃Ga intermetallic structure. Moreover, Ga plays a crucial role in improving catalytic activity and minimizing coke formation.

In conclusion, our research not only demonstrates excellent catalytic performance of multimetallic Ni-based catalysts in CO_2 conversion, but also offers a new perspective to prolong their lifespan. The findings of this study contribute to advancing the goal of carbonneutralization in current industrial processes.

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