Smart Designs of Anti-Coking and Anti-Sintering Ni-Based Catalysts for Dry Reforming of Methane: A Recent Review

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Abstract: Dry reforming of methane (DRM) reaction has drawn much interest due to the reduction of greenhouse gases and production of syngas. Coking and sintering have hindered the large-scale operations of Ni-based catalysts in DRM reactions at high temperatures. Smart designs of Ni-based catalysts are comprehensively summarized in four aspects: surface regulation, oxygen defects, interfacial engineering, and structural optimization. In each part, details of the designs and anti-deactivation mechanisms are elucidated, followed by a summary of the main points and the recommended strategies to improve the catalytic performance, energy efficiency, and utilization rate.

Keywords: Ni-based catalysts; dry reforming of methane; anti-coking; anti-sintering

1. Introduction

Many studies have been focused on the conversion of CO 2 and CH 4 (two greenhouse gases) to value-added chemicals or petrochemicals in various processes, such as reforming of methane [1–8], CO 2 methanation [9–16], membrane-assisted reaction [17–21], chemical looping [22], reversed water-gas-shift [23,24], and catalytic decomposition [25]. Therein, dry reforming of methane (DRM) reaction draws much interest because it contributes to the reduction of greenhouse gases (CO 2 and CH 4) and generates a mixture of CO and H 2 (syngas), which can be used as fuels and further converted to high value-added petrochemicals [26–31]. The reaction equation is shown as below:

\[ \text{CH}_4 + \text{CO}_2 \leftrightarrow 2\text{CO} + 2\text{H}_2 \quad \Delta H_{298K}^\circ = 261 \text{ kJmol}^{-1} \]  

(1)

The low H 2 :CO ratio is favorable to Fischer–Tropsch reaction due to the promotion of chain growth and suppression of methanation. Also, the endothermic forward reaction benefits the Solchem process by converting solar energy to chemical energy. Moreover, DRM reaction is applicable for the storage of seasonal energy in the form of CO and H 2 [32–35].

In order to activate the reactants, noble metal catalysts are applied and proved to be highly reactive. However, the high cost, low abundance, and metal sintering hinder their large-scale applications in industry [30]. Ni-based catalysts have been widely studied due to their low cost and good activities, but carbon deposition, metal sintering, and surface oxidation lead to the deactivation of catalysts and impede their further applications [30]. For example, the in situ XPS technique was applied to monitor the evolution of Ni surface during the reaction. Results showed that during the inactive period, 3 nm
layer of NiO covered the metallic Ni surface, leading to the decrease of reactivity [36]. Solutions have been provided to address this issue, including the size control of active metals, doping second transition element, adding alkali earth or rare earth promoters to the support, tuning metal-support interaction (MSI), structure designs and optimization of reaction parameters [26–31]. Researchers have reviewed the progress of Ni-based catalysts for DRM reaction in terms of the metal doping, porous supports, and kinetics/mechanisms, which shows the high importance of and widespread attention paid towards the DRM reaction [26–31]. However, it is rare to see a summary of the structure designs of supports, architectures of the catalysts, surface species control and interfacial strength adjustment in recent years, which greatly affects the active surface area, activities, and anti-deactivation capabilities. In this review, we summarize the recent advances of Ni-based catalysts’ modifications and corresponding catalytic performances for DRM reaction. The following parts consist of surface regulation, oxygen defects, interfacial engineering, and structural optimization, followed by concluding remarks and prospects.

2. Surface Regulation

For most heterogeneous reactions, reactants are adsorbed and activated on the surface active sites to produce the intermediates and final products. Modifying the surface properties has significant effects on the catalytic process and performances. In DRM reaction, CO₂ and CH₄ are converted to CO and H₂ through adsorption, dissociation, and recombination on the surface of Ni-based catalysts. Surface basicity exerts a great influence on the catalytic conversions and coke resistance [30].

Surface acidity and basicity affect the electron environment of metal active sites and the adsorption of reactants, thus determining the catalytic performances. For example, Lewis acid sites reduce electron density of Ni species and balance the activation of CH₄ and removal of carbon deposits, leading to a high stability in DRM reaction [37]. Also, enhanced basicity of the supports improves the adsorption of CO₂, which subsequently dissociates into CO and oxygen atoms, reacting with the decomposition intermediate CH₄ to produce CO and inhibit carbon formation [30]. To enhance the surface basicity, various compounds have been utilized, including alkali and alkali earth metal oxides and rare earth metal oxides [30,38–41].

La₂O₃ as a rare earth metal oxide, acts as the support to promote the adsorption of CO₂ by reacting with CO₂ to form La oxycarbonate (La₂O₂CO₃) [42]. According to in-situ DRIFT study, carbon species from the activation of methane reacted with the La oxycarbonate to form CO, removing the surface coke and maintaining the active sites [43]. The reaction mechanism was shown as below [43]:

\[
\begin{align*}
\text{CH}_4 + * & \leftrightarrow * - \text{CH}_4 \\
\text{CO}_2 + \text{La}_2\text{O}_3 & \leftrightarrow \text{La}_2\text{O}_2\text{CO}_3 \text{ (RDS)} \\
* - \text{CH}_4 & \rightarrow * - \text{C} + 2\text{H}_2 \\
\text{La}_2\text{O}_2\text{CO}_3 + \text{C} - * & \rightarrow \text{La}_2\text{O}_3 + 2\text{CO} + * \text{ (RDS)}
\end{align*}
\]

To further enhance the CO₂ adsorption, MgO as a basic oxide is promising due to its successful applications [44–47]. When MgO was added into the Ni/La₂O₃, due to its strong basicity, more CO₂ adsorbed on the surface to react with La₂O₃ to form monoclinic La₂O₂CO₃. Interestingly, at a low Mg:La ratio, only hexagonal La₂O₃ and La₂O₂CO₃ mixtures were formed, which was not effective as monoclinic phase in reacting with carbon species, thus deactivating the catalysts much faster (seen in Table 1) [48]. To further elucidate the mechanism of removing the carbon deposits by La₂O₃, a theoretical study was conducted to compare the different catalytic pathways of La₂O₃ and La metal. La₂O₃-Ni followed the reaction pathway below: CH₄ + CO₂ + La₂O₃ →CH + CO₂(La₂O₂-O) + 3H →CHO + CO₂(La₂O₂) + 3H →2CO + 2H₂ + La₂O₃. It showed that the oxygen atom dissociated from La oxycarbonate reacted with CH or C species to form CO. On the other hand, La metal dopant was not effective as the oxide since the faster rate of CH → C + H for La-Ni accelerated the coke formation than La₂O₃-Ni catalysts [49]. As another rare earth metal oxide, CeO₂ may decrease the
total basicity. However, the intermediate and strong basic sites could be enhanced, thus promoting the CO$_2$ adsorption and modifying the activity (shown in Table 1) [50].

Besides La$_2$O$_3$, CeO$_2$ and MgO, other group metal oxides can also enhance the surface basicity. Considering the acidity of SiO$_2$ support, Ga$_2$O$_3$ (Table 1) was added as a promoter to enhance the basicity and facilitate the adsorption of CO$_2$ to form carbonate or bicarbonate species (Figure 1), which reacted with carbon deposits more easily than the physically or linearly adsorbed CO$_2$ on the pristine SiO$_2$ surface [51].

![Figure 1](image-url)  
Figure 1. Activation of CO$_2$ on Ga$_2$O$_3$-promoted SiO$_2$ support. Reproduced with permission from reference [51].

B$_2$O$_3$ as a non-metal oxide has been successfully doped into Ni/Al$_2$O$_3$ to enhance the surface hydroxyl group concentrations. The borated surface was rich in OH group which promoted the oxidation of carbon deposits from CH$_4$ decomposition [52].

In certain situation, if the basicity of the support is too strong, the CO$_2$ adsorption may not inhibit but lead to carbon deposition [53]. As shown in Table 1, to alleviate the basicity of Ni supported on MgO/Al$_2$O$_3$, Y$_2$O$_3$ was added to introduce more weak and medium basic sites on the surface [54,55]. By using 1.5 wt% of Y$_2$O$_3$, the highest conversion of CH$_4$ and CO$_2$ was achieved with the best stability [54]. In addition, La-promoted Ni–Mg–Al hydrotalcite was reported to be anti-coking due to the gasification of carbon deposits where weak and medium basic sites were increased [56].

In a word, adding promoters like MgO, La$_2$O$_3$ and Ga$_2$O$_3$ could enhance the surface basicity, thus facilitating the adsorption of CO$_2$ which dissociated into CO and O and removed carbon deposits. Besides the surface basicity adjustment, surface hydroxyl group concentrations could be increased by adding B$_2$O$_3$ as a promoter, which promoted the carbon oxidation and maintained the stability. But a careful control of the surface basicity may be necessary since a too strong basicity could not inhibit the coke formation.
Table 1. A summary of surface regulation effects on the catalytic performances of Ni-based catalysts for DRM reaction.

| Catalyst | Preparation Method | Reaction Condition | Methane Conversion | CO₂ Conversion | Carbon Formation | Comments | Ref. |
|----------|-------------------|--------------------|-------------------|----------------|-----------------|----------|------|
| Ni/L₂O₃  | Wet impregnation method | 700 °C; CH₄/CO₂ = 1 | 70% for CH₄ after 50 h | 75% for CO₂ after 50 h | 10.3% | Carbon species reacted with the La oxycarbonate to form CO. | [42] |
| Ni/MgO-L₂O₃ | Co-precipitation and impregnation method | 700 °C; CH₄/CO₂ = 1 | 63% for CH₄ after 200 h | 65% for CO₂ after 200 h | 0.031 molC/molCH₄ | When MgO was added into the Ni/L₂O₃, more CO₂ adsorbed on the surface to react with L₂O₃ to form monoclinic L₂O₃CO₃. | [48] |
| Ce-NiMgAl | Co-precipitation method | 550 °C; CH₄/CO₂ = 1 | 54.8% for CH₄ after 5 h | 37.4% for CO₂ after 5 h | Around 90% | CeO₂ enhanced the amount of medium and strong basic sites. | [50] |
| Ni/SiO₂–Gα₂O₃ | Incipient wetness impregnation method | 700 °C; CH₄/CO₂ = 1 | 70% for CH₄ after 10 h | 79% for CO₂ after 10 h | 16.8 mg/gcat | Ga₂O₃ was added as a promoter to enhance the basicity to facilitate the adsorption of CO₂ to form carbonate or bicarbonate. | [51] |
| 5%Ni/B₂O₃–Al₂O₃ | Impregnation method | 700 °C; CH₄/CO₂ = 1 | 75% for CH₄ after 65 h | 67% for CO₂ after 65 h | 13% | The borated surface was rich in OH group which promoted the oxidation of carbon deposits. | [52] |
| HTNi-Y  | Co-precipitation method | 700 °C; CH₄/CO₂ = 1 | 74% for CH₄ after 10 h | 78% for CO₂ after 10 h | 75.8% | Y₂O₃ was added to introduce more weak and medium basic sites on the surface. | [54] |
| YNiMgAl | Co-precipitation method | 700 °C; CH₄/CO₂ = 1 | 85% for CH₄ after 5 h | 90% for CO₂ after 5 h | 35.1% | Basicity increased with the increase loading of Y. | [55] |
| La-NiMgAl | Co-precipitation method | 550 °C; CH₄/CO₂ = 1 | 45% for CH₄ after 8 h | 37% for CO₂ after 8 h | 87.5% | La dopant enhanced the content of medium and weak basic sites. | [56] |

*HT: hydrotalcite.*
3. Oxygen Defects

In addition to the surface basicity and acidity regulation, oxygen defects including lattice and surface oxygen and oxygen vacancies enhance the oxygen mobility and adsorption, which effectively removes the carbon deposits during DRM reaction [30]. Rare earth metal oxides, transition metal oxides, mixed oxides, and perovskites will be discussed as below based on the origin of oxygen defects and corresponding effects on the carbon removal.

3.1. Rare-Earth Metal Oxides

Studies have shown that rare earth metal oxides can generate oxygen defects, promote oxygen mobility, and scavenge carbon species [57–77]. As the representative, CeO$_2$ with a good redox property can release lattice oxygen to generate oxygen vacancies, thus accelerating the oxygen mobility. When CeO$_2$ was used as the support for Ni metals, the catalytic performance benefited from the lattice oxygen defects of CeO$_2$ and the reversible conversion between Ce$^{3+}$ and Ce$^{4+}$. When Ce$^{4+}$ was reduced to Ce$^{3+}$, lattice oxygen was released and oxygen vacancies were formed, which improved the adsorption of oxygen species like O$_2$ or CO$_2$ to generate oxygen radicals, thus reducing the carbon deposition on the surface. However, an excessive amount of CeO$_2$ may cover the Ni active sites and lower the accessibility of reactants to the Ni metal surface, causing a low conversion of CH$_4$ and CO$_2$ [30,65,78,79].

Theoretical study was applied to explore and elucidate the role of CeO$_2$ in Ni catalysts for DRM reaction. Results showed that the activation sites for CH$_4$ and CO$_2$ were different. For CH$_4$ activation, the preferred sites included metal-support interface, oxygen vacancies at the metal-support interface and the metal vacancies. But for CO$_2$, the reactant molecule was mostly adsorbed and dissociated at oxygen vacancies at either the support or the metal-support interface. Moreover, the active sites could be regenerated by the interfacial oxygen of CeO$_2$, maintaining the stability of the catalyst [58]. The detailed mechanistic diagram is shown in Figure 2 below.

Another possible explanation of the catalytic effect of CeO$_2$ was presented as below:

\[ \text{CeO}_2 + n\text{CH}_4 \rightarrow \text{CeO}_{2n} + n\text{CO} + 2n\text{H}_2 \]  
\[ \text{CeO}_{2n} + n\text{CO}_2 \rightarrow \text{CeO}_2 + n\text{CO} \]

The lattice oxygen of CeO$_2$ reacted with CH$_4$ to produce CO and H$_2$, leaving a ceria with oxygen defects, which subsequently grabbed oxygen atoms from CO$_2$ to recover to CeO$_2$ and produce another molecule of CO. The initial high oxidation state of Ce ion is essential to the whole process. However, during the DRM reaction, the existence of hydrogen possibly reduced Ce$^{4+}$ to Ce$^{3+}$, decreasing the conversion of CH$_4$ and CO$_2$. Therefore, the immediate removal of hydrogen product from the system determines the catalytic effect of CeO$_2$ [80].

To stabilize the CeO$_2$ in the Ni catalysts, ZrO$_2$ can be added to substitute the Ce in the lattice to form a cubic phase of Ce$_x$Zr$_{1-x}$O$_2$. This structure contains more oxygen vacancies and higher oxygen storage capacities than tetragonal phase. Because of the enhanced oxygen release capability, carbon deposits around the oxides are easily oxidized to form CO [81]. Likewise, as seen in Table 2, in another catalyst of Ni supported on CeO$_2$-ZrO$_2$ mixed oxides, the bonding degree determined the amount of oxygen vacancies. Also, the different oxidation states of Ce and Zr produced more oxygen vacancies. With the increase of Ni concentration, oxygen vacancies increased accordingly. When Ni reached the optimal molar ratio (9.3 mol%), the highest conversion of CH$_4$ and CO$_2$ was achieved [82].

Besides CeO$_2$, Y$_2$O$_3$ is also proved to be effective in removing the carbon deposits due to its surface oxygen mobility. During the DRM reaction, the movement of surface oxygen was accelerated with the generation of interstitial oxygen ions when lattice defects were produced in Y$_2$O$_3$. Interestingly, Y$_2$O$_3$ with a smaller crystal size exhibited a higher surface oxygen mobility, thus decreasing the carbon formation rate [57]. In another study shown in Table 2, α-oxygen was found on the surface of
Y$_2$O$_3$ to activate CH$_4$ and remove the carbon. Considering the limited amount of $\alpha$-oxygen, it was necessary to feed a small amount of oxygen to maintain the stability of the catalyst by regenerating the $\alpha$-oxygen [83].

Figure 2. Three paths of C* elimination. (a) Relative energy of C* elimination at m, ms, and ovms sites. (b) Relation of d band center with CO* adsorption energy. (c) Corresponding configurations of part a. Gray spheres represent C, red spheres represent O, green spheres represent Ni and cream-colored spheres represent Ce. Reproduced with permission from reference [58].

Another rare earth metal oxide La$_2$O$_3$ also draws attention in recent years as a promoter for Ni catalysts to inhibit coke formation in DRM reaction. As presented in Table 2, besides the effect of
La oxycarbonate formed with adsorbed CO$_2$ mentioned previously, the addition of small amount of Sr could enhance the surface oxygen species mobility, which facilitated the C-H activation and coke removal, leading to a high CO$_2$ conversion and low carbon formation even at 600 °C reaction temperature [84].

In addition to the mixed oxides of CeO$_2$ and ZrO$_2$ above, two rare earth metal oxides can also be mixed to form a support for Ni catalyst, such as Ce$_{0.70}$La$_{0.20}$Ni$_{0.10}$O$_{2-δ}$. During the reaction, the lattice of CeO$_2$ expanded due to the partial dissolution of La$^{3+}$ into the lattice structure to generate oxygen vacancies. Moreover, with the increase of reduction temperature, the change of Ce oxidation state further increased the amount of oxygen vacancies. Under the condition of GHSV = 26,400 h$^{-1}$, no carbon deposition was found, demonstrating the excellent coke resistance (seen in Table 2) [85].

In a word, owing to the redox property and high oxygen storage capacity, rare earth metal oxides can produce oxygen defects (vacancies) with the change of oxidation state and release of oxygen atoms. With the addition of second metals, the synergetic effect may generate more oxygen defects so that the activation of CH$_4$ and CO$_2$ as well as the carbon removal can be enhanced simultaneously, leading to the high activity and stability of Ni catalysts for the DRM reaction.

3.2. Transition Metal Oxides

Due to the incompletely occupied d orbitals, transition metal oxides have been widely studied as a support to enhance the coke resistance by producing lattice oxygens to reduce the carbon depositions during DRM reaction. ZrO$_2$ and TiO$_2$ are two representatives which will be discussed in detail below [86–89]. As for Fe, considering its main existing form as a Ni-Fe alloy instead of metal oxides, the effect of Fe will be illustrated in the interfacial engineering part.

The lattice oxygen species of ZrO$_2$ plays a crucial role in the anti-coking property of Ni catalysts in DRM reaction. As seen in Table 2, the results of XRD and XPS showed that CH$_4$ decomposed into CH$_x$ fragments and H atoms. The latter combined with each other to produce H$_2$; the former adsorbed on the Ni metal to form NiCH$_x$ species. Because of the intimate interaction between Ni and ZrO$_2$, the lattice oxygen of ZrO$_2$ reacted with CH$_x$ to form CO and H$_2$ and regenerated the Ni active sites. In the meanwhile, the coke formation was inhibited due to the release of lattice oxygen of ZrO$_2$ and its affinity for the CO$_2$ adsorption [59].

To enhance the interaction between Ni and ZrO$_2$, a core-shell structure of Ni-ZrO$_2$@SiO$_2$ was prepared by hydrolysis of TEOS to form a mesoporous SiO$_2$ shell and thermal treatment of NiZr(OH)$_x$ to form a Ni-ZrO$_2$ core as shown in Figure 3. More active sites were formed at the interface and perimeter of Ni-ZrO$_2$ with the stabilization effect of ZrO$_2$ on Ni. As presented in Table 2, more abundant active oxygen provided by ZrO$_2$ easily accessed the carbon species at Ni active sites and oxidized the coke to form CO, maintaining the stability of the catalyst [89].

In order to elucidate the effect of different oxygen species in ZrO$_2$ on the carbon removal for Ni catalysts in DRM reaction, three types of calcination atmospheres were applied. Results showed that the optimal ratio of surface adsorbed oxygen to lattice oxygen was achieved under H$_2$ treatment, most favorable for CO$_2$ and CH$_4$ adsorption and activation, thus leading to the highest conversions and stability. Relative speaking, catalysts calcined under N$_2$ and oxidizing atmospheres presented a lower ratio of surface oxygen to lattice oxygen, suffering from the serious carbon formation shown in Table 2 [90].

Besides ZrO$_2$, another transition metal oxide TiO$_2$ is proved to be effective in removing the carbon due to the lattice oxygen. Two oxygen species were found in TiO$_2$, that is, oxygen ions inside TiO$_2$ and lattice oxygen at the interface. The later could react with the coke formed on the surface of Ni to produce CO, thus maintaining the active surface area for the reactants to continue dissociation. Moreover, water vapor provided by the humid environment showed a high affinity for the oxygen vacancies and dissociated on TiO$_2$ to reduce the carbon deposits [86].
In a word, the lattice oxygen of ZrO$_2$ and TiO$_2$ can react with surface carbon species such as CH$_x$ and carbon deposits to form CO. A careful control of the ratio between surface and lattice oxygen may affect the coke resistance. Further strengthening the MSI can lead to the intimate contact between abundant oxygen species and carbon residues, resulting in a high conversion and low carbon deposition.

### 3.3. Perovskites

Represented by the formula ABO$_{3}$, perovskite structures have been widely investigated in heterogeneous catalysis at high temperatures due to its good thermal stability [91–97]. By substitution of A-site ions, the dimension of unit cells changes, thus changing the covalence of B-O bond accordingly, resulting in a lattice defect formation. These defects cause the increased amount of oxygen adsorption and high concentration of oxygen vacancies, leading to a promoted oxygen fluidity in the crystal structure [98–100]. Therefore, perovskites are extensively applied in DRM reaction to enhance the coke resistance of Ni catalysts.

Various elements have been added into LaNiO$_3$ to replace A-site or B-site ions [101–104]. For example, Sr was doped in the perovskite structure to partially substitute A-site La ions to tune the cell dimension and oxygen states. The lattice distortions generated high lattice oxygen mobility, activating C-H bond in CH$_4$ and adsorbing CO$_2$ molecules to form La$_2$O$_2$CO$_3$ [104]. For B-site replacement as presented in Table 2, Cu partially substituted Ni ions to enhance the mobility of lattice oxygen species, leading to a high conversion of CH$_4$ but a low catalytic stability because agglomeration of Ni particle took place with a weak MSI. On the contrary, Fe-substituted LaSrNiFeO$_3$ catalyst exhibited a long-term stability in DRM reaction due to the abundance of lattice oxygen and strong MSI [104]. The modification effect of Fe is elucidated in another work [105]. When Fe was doped into LaNiO$_3$ structure, Fe could be converted to FeO and Fe$_3$O$_4$ by CO$_2$ and Fe$_3$O$_4$ reacted with La$_2$O$_3$ to form LaFeO$_{3.5}$, which distorted the lattice structure and produced oxygen vacancies via compensation of the low valency of Fe ions. To modify the A-site of LaNiFeO$_3$, Ce can be added to partially replace the La to introduce more oxygen vacancies by both activating the B-site ions and reversibly shuttling between Ce$^{3+}$ and Ce$^{4+}$. In detail, unlike Sr$^{2+}$ with a lower valency than La$^{3+}$, Ce$^{3+}$ was not possible to directly provide oxygen vacancies when doped into the perovskite structure. But according to the TPR profiles as Figure 4 below, the reduction peak of Ni(Fe) ions shifted to lower temperatures with the increase of Ce amount, reflecting an improved reducibility and more oxygen vacancies indirectly offered by Ce. On the other hand, due to the local redox fluctuations of Ce ions, the regeneration of transition metals may be facilitated with the promoted migration inside/outside the perovskite structure. Therefore,
more oxygen vacancies brought by Ce enhanced the oxygen mobility and backfilling of lattice oxygen and promoted the adsorption and activation of CO$_2$ to CO and oxygen atoms which in turn replenished the oxygen vacancies (seen in Table 2) [106]. As for the B-site of LaNiFeO$_x$, in another study, Co was doped into LaNiFeO$_3$ to improve the oxygen mobility due to the spin state, multivalent property and oxygen affinity. When the molar ratio of Co equaled to 0.1 and 0.3, the modified perovskite structure presented a structural stability and high conversions [101].

![Figure 4](image_url)  
**Figure 4.** TPR profiles of fresh La$_{1-x}$Ce$_x$Ni$_{0.5}$Fe$_{0.5}$O$_3$ with different $x$ values. Reproduced with permission from reference [106].

Besides LaNiO$_3$ structure, LaAlO$_3$ perovskite catalysts are proved to be more resistant to coke formation than $\alpha$-Al$_2$O$_3$ due to the higher thermal stability and more oxygen vacancies [107]. Two possible mechanisms were illustrated in Figure 5 below, which differed in the location of oxygen vacancies and NiO in the perovskite. When oxygen ions of NiO happened to be in the oxygen vacancies, two electrons of H$_2$ were released to reduce Ni ions to form metallic Ni and produce H$_2$O by reacting with O$^{2-}$, enhancing the dispersion of Ni particles on the surface. When oxygen ions were located adjacent to the oxygen vacancies, H$_2$ would be heterolytically dissociate into H$^+$ and H$.^-$. The later adsorbed on the surface of acidic support while the former combined with O$^{2-}$ to produce water. In this case, the reduction of NiO was not preferred and the metal dispersion became poor. These two mechanisms reflected the promotional effect of oxygen vacancies on the migration of lattice oxygen of NiO and coke removal at the perovskite interface.

![Figure 5](image_url)  
**Figure 5.** Scheme of the reduction/activation mechanism of the Ni catalyst supported on LaAlO$_3$ catalysts. (I)—Terminal oxygen ion of the NiO clusters located on the surface oxygen vacancy. (II)—Terminal oxygen ion of the NiO clusters located adjacent to the surface oxygen vacancy. Reproduced with permission from reference [107].
Table 2. A summary of oxygen defects effects on the catalytic performances of Ni-based catalysts for DRM reaction.

| Catalyst       | Preparation Method                  | Reaction Condition | Methane Conversion | CO₂ Conversion | Carbon Formation | Comments                                                                 | Ref. |
|----------------|-------------------------------------|--------------------|--------------------|----------------|------------------|--------------------------------------------------------------------------|------|
| Ni-CoO–ZrO₂    | Co-precipitation method             | 700 °C; CH₄/CO₂ = 1 | 73% for CH₄ after 35 h | 83% for CO₂ after 35 h | NA               | The lattice oxygen of ZrO₂ reacted with CH₄ to form CO and H₂ and regenerated the Ni active sites. | [59] |
| NiRhCe₂Zr₁.₃₁  | Pseudo sol-gel method               | 700 °C; CH₄/CO₂ = 7:3 | 16% for CH₄ after 25 h | 37.5% for CO₂ after 25 h | 4–10 mg<sub>cat</sub>/100 mg<sub>cat</sub> | ZrO₂ and CeO₂ formed a cubic phase of Ce₂Zr₁.₃₁O₇ generating more oxygen vacancies and higher oxygen storage capacities than tetragonal phase. | [81] |
| NiCeO₂–ZrO₂    | Co-precipitation/molten salt synthesis | 750 °C; CH₄/CO₂ = 1 | 2.56 mmol/(m²·h) | NA             | 3.75 × 10⁻⁴ mg<sub>cat</sub>/h | With the increase of Ni concentration, oxygen vacancies increased accordingly. | [82] |
| PdO–NiO(Y₂O₃) | Wet impregnation method             | 700 °C; CH₄/CO₂O₂ = 2/1/1 | 91.11% for CH₂ after 24 h | 44.98% for CO₂ after 24 h | 16.8 mg<sub>cat</sub>/h | α-oxygen was found on the surface of Y₂O₃ to activate CH₄ and remove the carbon. | [83] |
| Ni–SDL (Sr doped La) | Wet impregnation method       | 600 °C; CH₄/CO₂ = 1 | 78% for CH₂ after 10 h | 60% for CO₂ after 10 h | 50 mg<sub>cat</sub>/h | The addition of small amount of Sr could enhance the surface oxygen species mobility. | [84] |
| Ce₀.₇₅La₀.₂₅Ni₀.₁₀O₂₋₉   | Combustion synthesis             | 750 °C; CH₄/CO₂ = 1 | 71% for CH₂ after 50 h | 83% for CO₂ after 50 h | n.d.           | The lattice of CeO₂ expanded due to the partial dissolution of La⁺⁺ into the lattice structure to generate oxygen vacancies. | [85] |
| Ni–ZrO₂@SiO₂ | Microemulsion method               | 800 °C; CH₄/CO₂ = 1 | 90.5% for CH₂ after 240 h | 93.2% for CO₂ after 240 h | n.d.           | More abundant active oxygen provided by ZrO₂ easily accessed to the carbon species and oxidized the coke to form CO. | [86] |
| Ni/ZrO₂    | Deposition-precipitation method    | 700 °C; CH₄/CO₂ = 1 | 52% for CH₂ after 300 min | 59% for CO₂ after 300 min | 26.7%         | The optimal ratio of surface adsorbed oxygen to lattice oxygen was achieved under H₂ treatment. | [80] |
| La(Co₉Ni₁₋ₓ)₅FeₓO₁₅   | Sol-gel self-combustion method     | 750 °C; CH₄/CO₂ = 1 | 70% for CH₂ after 30 h | 80% for CO₂ after 30 h | <3 mg<sub>cat</sub>/h | Co was doped into LaNi₅FeO₁₅ to improve the oxygen mobility due to the spin state, multivalent property and oxygen affinity of Co. | [102] |
| La₀.₄Sr₀.₆Ni₁₃Co₀.₃O₇ | Sol-gel method                     | 600–800 °C; CH₄/CO₂ = 1 | 80% for CH₂ after 24 h | 80% for CO₂ after 24 h | 70 mg<sub>cat</sub>/h | Partial substitution by Sr and Cu generated higher lattice oxygen mobility. | [105] |
| La₁.₄Ce₀.₆Ni₁₃Fe₀.₅O₇ | Sol-gel self-combustion method | 750 °C; CH₄/CO₂ = 1 | 62% for CH₂ after 25 h | 72% for CO₂ after 25 h | NA             | More oxygen vacancies brought by Ce enhanced the oxygen mobility and backfilling of lattice oxygen. | [107] |
In a word, partial substitution of A-site and B-site can generate oxygen vacancies by lattice distortion or redox property, which promotes the lattice oxygen mobility and adsorption of CO$_2$ to form CO and oxygen atoms. Carbon species derived from the methane activation can be removed by the oxygen atoms, presenting the potential of substituted perovskites as a coke scavenger in the DRM reaction.

For the utilization of oxygen defects, rare earth metal oxides, transition metal oxides and perovskites could be adopted as the supports or promoters which generated oxygen vacancies, surface oxygen or lattice oxygen species, facilitating the mobility of oxygen and oxidizing the coke in time.

4. Interfacial Engineering

Either the metal-metal synergy or metal-support interaction affects the electron density, surface composition and metal dispersion, thus determining the activation of reactants anti-deactivation properties of Ni catalysts [108,109]. In the following, two strategies of interfacial engineering will be discussed in detail, namely Ni-metal alloy formation and Ni-support interaction.

4.1. Ni-Metal Alloy Formation

Bimetallic Ni catalysts enjoy the benefits of synergetic effects between Ni and another metal, such as oxygen affinity, redox cycle, alleviated agglomeration, produced reducibility, dilution effect [109–115]. For example, Cu can modify the Ni environments and prevent carbon formation [116]. Co can inhibit sintering of Ni by formation NiCo alloy and enhance the resistance to oxidation [117]. Noble metals can improve the Ni dispersion on the surface of the support and increase the tendency of CH oxygenation instead of dehydrogenation [118,119]. The following part will cover the effects of Ni alloys with different groups of metals on the physicochemical properties and catalytic performances of Ni-based catalysts in DRM reaction.

4.1.1. Alloy of Ni and Noble Metals

When noble metals are added to Ni catalysts, the reducibility of Ni is enhanced due to the spillover of H$_2$ on the surface of noble metals. In the presence of Pt, the reduction temperatures of Ni-Pt alloy shifted to lower regions in the TPR profiles. This was due to the ready adsorption of H$_2$ molecules on the Pt surface and dissociated into H species which subsequently reduced NiO to form metallic Ni [120]. Besides the promotional effect on the reducibility of Ni ions, noble metals can also alleviate the oxidation of Ni by forming alloys. It is well known that a layer of Ni$^{2+}$ covering the metallic Ni particles reduces the active sites, thus leading to a poor activation of CH$_4$. Due to the resistance to oxidation under atmospheric or pressurized conditions, noble metals prevent Ni metals from being oxidized when forming alloys [121]. Moreover, Ni dispersion can be enhanced due to the dilution effect of noble metals when forming alloy with them. The resulting small particles contributed to the enhanced coke resistance in DRM reaction [113].

A mechanistic study was to explore the surface composition and reconstruction of Ni and Pt under oxidative environment as shown in Figure 6. When heating a layer of Pt atoms covering Ni bulk, Pt atoms would form a Ni-Pt alloy on the surface. Under oxidative treatment, segregation of Ni from the alloy was observed and the Ni-rich surface was exposed to outside to form O/Ni-Pt-Pt structure. Similarly, when a layer of Ni atoms covered Pt bulk, Ni atoms would diffuse into the sublayers to form a stable Pt-Ni-Pt structure upon being heated. When treated with oxidative atmospheres, Ni again segregated to the surface to form the same O/Ni-Pt-Pt structure. By using EXAFS to characterize the Pt coated Ni catalysts and analyze the coordination number of Pt-Ni and Pt-Pt, an increased value of N (Pt-Ni) and decreased value of N (Pt-Pt) reflected the migration of Pt to sublayers and Ni to the surface, transforming the Pt monolayer to a core-shell structure with Ni as the core and Ni-Pt alloy as the shell. This resulting structure facilitated the oxidation of CH and carbon removal (seen in Table 3) [119]. Moreover, recent operando studies suggested that the formation of PtNi alloys could promote the
formation of Ni oxide surface clusters, which related to the strong MSI effect, thus leading to a catalytic performance enhancement [122,123].

![Figure 6. (a) Surface restructuring of Ni/Pt bimetallic systems induced by temperature and adsorbates. (b) Evolution of the surface structure of the Ni/Pt bimetallic NPs upon thermal treatment. Reproduced with permission from reference [119].](image)

Preparation methods may affect the surface composition. When co-impregnation was applied to prepare Ni-Pd/Al₂O₃ catalyst, a uniform Ni-Pd alloy was formed due to the strong interaction between two metals. When sequential impregnation was utilized, segregation of Pd atoms was observed because of the weak interaction. In this case, Pd atoms preferred to locate on the surface instead of diffusing into the sublayers [124].

4.1.2. Alloy of Ni and Transition Metals

Similar with noble metals, transition metals can enhance the anti-sintering and anti-coking properties of Ni catalysts by forming alloys [39,77,125–129]. But the driver force may differ from that of noble metals.

After the addition of Co, Ni-Co alloys exhibited an improved affinity for oxygen but a stronger resistance to oxidation than Co alone [130]. First principle calculations showed that Co was very sensitive to oxygen which was proved by the experimental results that when Co loading was below 9 wt% on Al₂O₃, the Co metals suffered from oxidation, thus losing the active sites and deactivating DRM reaction [131]. However, when Ni-Co alloy was formed, due to the synergy between two metals, both the oxidation of metals and carbon formation were inhibited due to the adsorption of CO₂. At 800 °C, only 1.7 wt% of carbon deposits were produced after 100 h DRM reaction [130,132]. One possible explanation for the inhibition effect of Co on oxidation is demonstrated by Gao et al. [133] in the XPS study of reduced Ni-Co/SiO₂, whereby the shift of the binding energies of Ni and Co reflected the donation of electrons from Co to Ni, increasing the electron density of Ni and prevented Ni from being oxidized. Meanwhile, metal sintering was also impeded. As for the anti-coking ability, DFT calculations showed that the activation of CH₄ on Ni clusters produced mainly CH₃ and H species; while for Ni-Co alloy, due to the high affinity for oxygen, CH₄ reacted with oxygen atoms to form CH₃ and OH species, which removed surface carbon deposits more readily (shown in Table 3) [134].

Compared with Co, the addition of Fe mostly reduces the carbon deposits via redox cycle. In the Ni-Fe/MgAl₂O₃ catalyst, after pulse injection of CO₂, Ni-Fe decomposed to form metallic Ni and Fe. Different from stable Ni phase, Fe was easily to be oxidized to FeOₓ. Following the Mars–van Krevelen (MvK) mechanism, carbon species on the surface of Ni were oxidized by FeOₓ to form CO and Fe [135]. Likewise, in another study, dealloying and realloying process took place during the DRM reaction. In detail as shown in Table 3, CO₂ adsorbed on the Ni-Fe alloy oxidized the Fe to form FeO, which reacted with the carbon derived from the CH₄ decomposition to form CO and regenerate metallic Fe. The redox cycle of Fe/Fe²⁺ was crucial in inhibiting the carbon formation [136].
As another transition metal, Cu has been proved effective in DRM reaction when forming alloy with Ni. On one hand, Cu alleviated the metal sintering after 30 h reaction; on the other hand, with a molar ratio of Cu: Ni equaling to 1:3, a high conversion of reactants with minimal carbon formation was achieved [137]. A possible mechanism was elucidated by DFT calculation that O(OH) species preferred to adsorb on the CuNi (111) phase, which promoted the elimination of carbon deposits. Besides the DFT calculations, experimental evidence is expected [26].

As for the Cu amount, it was found that by increasing the concentration of Cu, methane activation may be inhibited since Ni active sites were covered by Cu which exhibited slower dissociation kinetics than Ni. As shown in Table 3, this competitive adsorption can be minimized by a careful control of the amount of Cu added in the Ni catalysts [137].

4.2. Ni-Support Interaction

Metal-support interaction (MSI) influences the metal size, dispersion, surface area and reducibility of Ni-catalysts, in turn determining the reactivity and stability in DRM reaction [26,30]. MSI can be turned by the support materials and promoters/dopants. For example, though silica is an inert support, by adopting a modified prepared method, a stable Ni phyllosilicate structure can be formed which enhances the MSI and controls the size and dispersion of Ni [29]. Ca as a promoter promotes the interaction between Ni and Al2O3, thus impeding the metal sintering and improving the dispersion of Ni [138]. However, a too strong MSI may exert a negative impact on the metal surface area since a high reduction temperature is needed [30].

When CeO2 was used as the support for Ni, strong MSI was formed, which facilitated the dissociation of methane in DRM reaction [65,67]. By DFT calculations, the activation barrier for Ni alone was 0.9 eV while for Ni-CeO2-x, the value dropped to 0.15 eV only. It is widely known that the rate determining step for DRM reaction is the activation of C-H bond in CH4 molecules [139], therefore, a lower value of activation energy enhanced the reaction rate and energy efficiency. After 80 h reaction at 450 °C, the conversion was 12%, close to the equilibrium value at this condition. As for the reaction pathway, once the first H was released from CH4, the following transformation of CHx species proceeded very quickly. The carbon species was oxidized by the O adatoms of CO2 to form CO, and H desorbed the surface in the form of H2O. Interestingly, the non-defective CeO2 exhibited no signs of CO2 activation. Only when the Ni2+ and Ce4+ were reduced by the decomposition products of CH4 to form metallic Ni and Ce3+, could the CO2 reacted with oxygen vacancies and H adatoms produced by methane activation to form CO. This was proved by the decreased amount of Ce4+ exposed to CH4 and CO2 atmospheres compared with CH4 alone according to Ce 4d XPS results as shown in Figure 7 below. In summary as presented in Table 3, Ni and O atoms cooperatively adsorbed the CH4 molecules due to the Ni-Ce solid solution formation, where a strong MSI tuned the electronic or chemical properties of Ni, thus enhancing the activation of methane by Ni metals and inhibiting the carbon formation proved by the lack of C or NiCx peaks as below [65].

Similarly, in another work, under a small Ni coverage, Ni and O atoms collaboratively activated the C-H bond in CH4, leading to a good conversion at low reaction temperature 377 °C. Due to the strong MSI and electrons transfer from Ni to CeO2, Ni2+ and Ce3+ formed, generating the most stable site Ni2+@O-hollow. Ni2+ was the only Ni species either before or after the adsorption of CH4 at room temperatures. When the coverage of Ni increased to 0.5 ML, NiCx species appeared on the surface; when the Ni coverage was below 0.2 ML, the small Ni atoms or particles contacted closely with the CeO2 support, resulting in a much lower activation energy of CH4 than that with a Ni coverage of 0.5 ML. A small loading of Ni formed small particles, which experienced large electronic perturbations from CeO2. These small Ni particles and O atoms in CeO2 promoted the activation of CH4 in a cooperative way. When Ni2+/CeO2 was transformed to Ni0/ CeO2-x by reacting with C species from CH4 decomposition, CO and oxygen vacancies were generated [67,140,141]; the oxygen vacancies formed on the reduced CeO2-x facilitated the adsorption and activation of CO2, thus leading to an enhanced conversion and stability in DRM reaction [67].
In certain situations, however, strong MSI may not be a benefit. As shown in Table 3, when a layer of CeO$_2$ covered the Ni-Co alloys, the promotional effect of Co on the enhanced MSI was cancelled off by the reduction of active sites exposed to the reactants [63]. Similar findings were shown presented for the Ni/TiO$_2$. Despite the strong MSI, the surface free energy may be reduced due to the coverage of Ni active sites by the migration of TiO$_2$ species over the surface.

In addition to CeO$_2$, owing to the strong basicity of MgO, a strong MSI between Ni and MgO derived from the solid solution probably enhance the anti-coking property [142,143]. However, the surface and under-coordinated Ni still exhibited a sensitivity to coke formation [144]. MgAl$_2$O$_4$ spinel structure may be a promising candidate to produce a highly dispersed Ni particles due to the strong MSI; together with the basicity of the support, CO$_2$ adsorption will be promoted, and carbon deposition will be alleviated (seen in Table 3) [145]. But an extraordinarily strong MSI may bring a high reduction temperature, possibly leading to the metal sintering. To further modify the dispersion of Ni, Co was added to the Ni/MgAl$_2$O$_4$ to slow the reduction of Ni and generate a small and uniform Ni particle, resulting in an enhanced DRM activity [146].

Besides the rare earth metal oxides, alkali earth metal oxides and transition metal oxides, SiO$_2$ as a support can also form a stable structure with Ni species with a modified preparation method [147,148]. With the addition of NH$_3$ to control the pH, Ni phyllosilicate formed between the Ni and SiO$_2$ under thermal treatment of Ni precursors. Upon decomposition of the phyllosilicate, highly dispersed Ni particles interacted strongly with the SiO$_2$, achieving a high and stable conversion of CH$_4$ (74%) and CO$_2$ (82%) over 50 h at 700 °C as seen in Table 3 [149].

In addition to the ammonia-assisted method, thermal treatment also benefited the Ni phyllosilicate formation. Compared with air calcination or H$_2$ treatment, a sequential calcination method was advantageous in terms of the size and MSI of Ni/SiO$_2$ catalysts. In detail, the first step of H$_2$ calcination at 400 °C scavenged oxygen radicals produced from the nitrate decomposition, reducing the Ni particle size and enhance the dispersion. The subsequent air calcination at 700 °C strengthened the MSI by forming the Ni phyllosilicate, anchoring Ni particles and preventing the sintering. As presented in Table 3, the Ni/SiO$_2$ catalyst prepared with the facile modified incipient wetness impregnation method exhibited an outstanding catalytic performance (80% for CH$_4$ and 85% for CO$_2$) with little sintering and no coke formation after 100 h DRM reaction [150].

In a word, by adjusting the Ni-metal interface and Ni-support interaction, the physicochemical properties of Ni catalysts can be improved due to the synergy between Ni and dopant metals and the electronic and chemical perturbation by the support. In turn, catalytic performances and anti-deactivation abilities in DRM reaction can be enhanced by introducing oxygen vacancies, surface basicity, oxygen affinity, redox cycle, low activation barrier, steric hindrance, and electron transfer upon interfacial engineering in Ni-based catalysts.
Table 3. A summary of interfacial engineering effects on the catalytic performances of Ni-based catalysts for DRM reaction.

| Catalyst             | Preparation Method | Reaction Condition | Methane Conversion | CO₂ Conversion | Carbon Formation | Comments                                                                                           | Ref. |
|----------------------|--------------------|---------------------|--------------------|----------------|-----------------|---------------------------------------------------------------------------------------------------|------|
| NiCo/CeO₂            | Incipient wetness impregnation method | 700 °C; CH₄/CO₂ = 1 | 64% for CH₄ after 300 min | 69% for CO₂ after 300 min | 11.5% | When a layer of CeO₂ covered the Ni-Co alloy, the promotional effect of Co on the enhanced MSI was cancelled off. | [63] |
| Ni/Co₂               | NA                 | 450 °C; CH₄/CO₂ = 1 | 10⁻⁷ torr CH₄ after 80 h | 10⁻⁷ torr CO₂ after 80 h | NA | Ni and O atoms cooperatively adsorbed the CH₄ molecules due to the Ni-Co solid solution formation. | [65] |
| Ni/TiO₂              | Impregnation method | 790 °C; CH₄/CO₂ = 1 | 41% for CH₄ initially | 67% for CO₂ initially | 8.0 x 10⁻⁴ g/sec/gcat | The surface free energy may be reduced due to the coverage of Ni active sites by the migration of TiO₂ species. | [108] |
| NiPt/Al₂O₃           | NA                 | 700 °C; CH₄/CO₂ = 1 | 85.8% for CH₄ after 18 h | 91.2% for CO₂ after 18 h | 0.1 molC/mol₁surface-metal/h | Pt monolayer in a core-shell structure with Ni facilitated the oxidation of CH and carbon removal. | [119] |
| NiCo/MgO-ZrO₂        | Co-precipitation and impregnation method | 600 °C; CH₄/CO₂ = 1 | 6.6 mol(g atom NiCoₓsurf-x)⁻¹ | NA | NA | For Ni-Co alloy, due to the high affinity of Co for oxygen, CH₄ reacted with oxygen atoms to form CH₃ and OH species. | [134] |
| Fe-Ni/Mg₆Al₄         | Co-precipitation and incipient wetness impregnation method | 750 °C; CH₄/CO₂ = 1 | 0.13 molClIT s⁻¹ molg⁻¹ after third cycle | NA | 5.02 molCO molNi₄Fe⁻¹ | Carbon species on the surface of Ni were oxidized by FeOₓ, to form CO and Fe. | [135] |
| Fe-Ni/Mg₆Al₄         | Co-precipitation and incipient wetness impregnation method | 750 °C; CH₄/CO₂ = 1 | NA | NA | 5.02 molCO molNi₄Fe⁻¹ | Carbon species on the surface of Ni were oxidized by FeOₓ, to form CO and Fe. | [135] |
| Cu–Ni/SiO₂           | Hydrothermal process | 700 °C; CH₄/CO₂ = 1 | 70% for CH₄ after 30 h | 3% | | When Cu formed alloy with Ni, the metal sintering and carbon formation were alleviated. | [137] |
| Ca-promoted Ni₆Al₄    | Wetness co-impregnation method | 800 °C; CH₄/CO₂ = 1 | 84.8% for CH₄ after 240 min | 85.5% for CO₂ after 240 min | 24.9 mg/gcat/h | Ca promoted the interaction between Ni and Al₂O₃, thus impeding the metal sintering and improving the dispersion of Ni. | [138] |
| AuNi₆Al₄             | Sol preparation method | 650 °C; CH₄/CO₂ = 69:30 | 20% for CH₄ after 1000 min | 48% for CO₂ after 1000 min | 3 mg | MgAl₂O₄ spinel structure increased the basicity of the support, so CO₂ adsorption will be promoted, and carbon deposition will be alleviated. | [145] |
| Ni@Ni embedded SiO₂  | Self-templating method | 700 °C; CH₄/CO₂ = 1 | 74% for CH₄ after 50 h | 82% for CO₂ after 50 h | n.d. | Upon decomposition of the phyllosilicate, highly dispersed Ni particles interacted strongly with the SiO₂. | [149] |
| Ni/SiO₂              | Incipient wetness impregnation method | 700 °C; CH₄/CO₂ = 1 | 80% for CH₄ after 100 h | 85% for CO₂ after 100 h | n.d. | Sequential H₂-air calcination enhanced Ni dispersion and MSI with SiO₂. | [150] |
5. Structural Optimization

A small particle size can enhance the activity of Ni-based catalysts [151]. To alleviate the Ni sintering at high temperatures during DRM reactions, optimizations have been made on the structure of supports, such as porous materials which enhance the surface area and providing the anchoring points for metal active sites [30, 31]. Moreover, hierarchical designs of the catalyst including core-shell, yolk-shell, embedded and hollow structures, sterically hinder the migration of Ni and prevent the agglomeration [27, 30].

5.1. Porous Supports

High porosity of carbon materials inhibits the movement of metal particles and enhances the dispersion on the carbon support. Also, porous carbon materials can be prepared into various physical forms, such as pellets, cloth, and fiber [30]. Li et al. [152] synthesized porous carbon from renewable hydrochar as the precursor and successfully anchored Ni$^{2+}$ ions on the surface, leading to highly dispersed Ni nanoparticles with the diameter of less than 8 nm. Methane conversions reached 52% and 70% for 700 °C and 800 °C pretreated catalyst respectively.

In addition to porous carbon materials, silica with high porosities and ordered structures is widely used to control the particle size and enhance the metal dispersion by confining metal particles in the pores (seen in Table 4) [153–158]. SBA-15 with silanol groups on its surface was able to trap the Ni ions by bonding with amine groups of polyethylenimine (PEI) in the Ni-PEI complexes (Figure 8). Little sintering took place due to the confinement effect of SBA-15 with ordered channels, leading to a high stability over 40 h of reaction [155]. When ethylene glycol (EG) was used as the solvent, Ni-EG complexes were pumped into the pores of SBA-15 by capillary force, hindering the migration of Ni during the drying and calcination process [154].

![Figure 8. Schematic diagram of PEI-aided route. Reproduced with permission from reference [155].](image)

As shown in Table 4, besides Ni monometallic catalysts, bimetallic Ni-Co supported on SBA-15 maintained the activity over 50 h. The small particle size (4–5 nm) and high metal dispersions (17–20%) were achieved by the confinement effect of ordered channel of SBA-15 [159]. Perovskite structures LaNiO$_3$ were also impregnated into the mesoporous SBA-15 supports. With the help of the stable structures, the conversions were kept stable over 60 h without agglomeration of Ni particles [160].

Compared with SBA-15, MCM-41 and KIT-6 supported Ni catalysts exhibited lower conversions and higher carbon depositions in DRM reaction. For KIT-6, the micropores were occupied by Ni particles so that the pathways of reactants were blocked despite the high Ni dispersion [157]. For MCM-41, it suffered poorer structural stability than SBA-15 especially when TiO$_x$ and MnO$_x$ hindered the channels for reactants to pass through and the amorphous form of silica was transformed into crystals [161, 162].
To enhance the diffusion efficiency and thermal stability, as shown in Table 4, hexagonal mesoporous silica (HMS) with thicker pore walls and more interconnected channels was applied as the support of Ni to improve the metal dispersion and inhibit sintering by forming Si-O-Ni bonds, leading to a stable conversion over 100 h at 700 °C [158]. For other mesoporous silica supports, one way to enhance the metal dispersion on porous silica supports is to utilize a novel method (seen in Table 4). Colloidal approach was applied by converting Ni-silicide colloids into highly dispersed Ni nanoensembles supported on porous silica under thermal treatment, achieving a 60% higher syngas yields [163]. Polyols precursors were treated under inert atmospheres and decomposed to produce carbon residues, protecting Ni particles from migrating from internal pores to external surfaces. The near-equilibrium conversions at 750 °C resulted from the well-dispersed Ni particles inside the mesoporous channels exposed completely to the reactants [164].

In addition to the colloidal and polyols approaches, ligand-assisted incipient wetness impregnation (IWI) method is applied to control the size and MSI (presented in Table 4). Similar with PEI and EG forming complexes with Ni ions on SBA-15 supports, a combination of oleic acid (OA) and oleylamine (OAm) were used as co-ligands to protect Ni from agglomeration by ligand bonds between Ni ions and amine groups of OAm and hydrogen bonds between carboxylic groups of OA and silanol groups of mesoporous silica. Together with the steric hindrance of the hydrocarbon chain network, a high dispersion of Ni particles and strong MSI were achieved, leading to a stable activity in DRM reaction [165]. Moreover, this method proved effective for reducing the size bimetallic Ni-Co to 3.1 nm averagely, resulting in a high conversion of methane and CO₂ over 30 h at 700 °C [133]. Furthermore, in a systematic study of the amine effect, the organic amines with a long hydrocarbon chain and high concentration produced a small size of Ni and a high dispersion as well as a strong MSI [166].

Compared with inert materials like carbon and silica, alkali earth metal oxides, rare earth metal oxides and alumina presented both a porous structure and a surface modification effect [167–170]. NiO-MgO-Al₂O₃ mesoporous frameworks provided more active centers for the reactants to be activated and converted to syngas products. Ni particles were anchored within the pores to alleviate the sintering effect and prolong the lifetime in the reaction; with the basic promoters, carbon removal was facilitated due to the adsorption of CO₂ which was dissociated into CO and oxygen radicals (seen in Table 4) [167]. In another example as shown in Figure 9, mesoporous NiO-CeO₂-Al₂O₃ materials was prepared by evaporation-induced self-assembly method with the introduction of CeO₂ into Ni/Al₂O₃. The excellent thermal stability of this mesoporous structures (preserved over 80 h at 700 °C) effectively protected Ni from sintering. Also, the redox property of CeO₂ oxidized carbon deposits to help maintain the exposure of active sites to reactants during the reaction [171].

![Figure 9](image-url) Schematic illustration of the NiCeAl (R) preparation. Reproduced with permission from reference [171].

In a word, various porous materials such as carbon, silica and alumina provide accommodations for Ni nanoparticles to be protected from migrating from internal pores to external surfaces of the supports. Organic ligands, novel methods and promoters are utilized to further enhance the anti-sintering and anti-coking capabilities, leading to a high conversion of methane and CO₂ and long-term stability.
5.2. Hierarchical Designs

Besides the rational design of the porous supports to enhance the surface area and confine the metal nanoparticles inside the pores or channels, architecture optimizations of the catalysts have become an effective method to improve the sintering and coking resistance. Owing to the uniform microporosity, high thermal stability and unique shape selectivity, zeolites draw attentions in stabilizing Ni particles and prevent the sintering during DRM reaction [172,173]. Moreover, other smart designs emerge in recent years, such as core-shell structures. With the first design of “nanoreactor” Pt@CoO core-shell nanostructures, various nanocomposites containing metal and metal oxides protected by silica shells are developed in recent years, including core-shell, yolk-shell, hollow and embedded structures, to partially separate metal particles from each other and enhance the dispersion [30,174–182].

Ni@SiO$_2$ core-shell catalysts with Ni as the core and SiO$_2$ as the shell exhibited a strong resistance towards filamentous carbon deposition (shown in Table 4). However, amorphous carbon still formed within the nanocavities between the core and shell. Minimizing the nanocavities was recommended to modify the structure and improve the coke resistance [183].

To enhance the catalytic performance, a thick SiO$_2$ shell was formed to transform the core-shell structure to yolk-shell structures with Ni as the core surrounded by Ni nanoparticles embedded in the SiO$_2$ shell (Figure 10). With a 11.2 nm of thickness, this unique structure exhibited an enhanced conversion and stability in DRM reaction with the help of strong MSI and small satellite Ni particles [184]. However, the turnover frequency (TOF) can be increased by tuning surface area and Ni exposure. Instead of using micro-emulsion method, a self-templating method was utilized to successfully prepared a yolk-shell Ni@SiO$_2$ catalyst, increasing the TOF by 4.5 times of the previous one (shown in Table 4) [149].

![Schematic illustration of the formation process of Ni–yolk@Ni@SiO$_2$ nanocomposite. Reproduced with permission from reference [184].](image)

To further enhance the active surface area and metal loadings, as presented in Table 4, multi-core shell catalysts with multiple Ni cores and a mesoporous SiO$_2$ shell were synthesized by reversed micelle method. During the reaction, Ni particles migrated into the SiO$_2$ shell, leading to the formation of uni-nuclear Ni core embedded in the shell [185]. To achieve a better anti-sintering property, a layer of silica was coated on the multiple Ni cores supported on the mesoporous silica as shown in Figure 11. With the protective layer of silica, 5.2 nm of Ni particles with a narrow size distribution were prevented from agglomeration due to the strong MSI and steric hindrance, reaching a high stability over 170 h at 800 °C [186].

To improve the MSI between Ni and SiO$_2$, Ni phyllosilicate structures were prepared by ammonia evaporation method, which generated highly dispersed Ni nanoparticles with the size of 6 nm supported on silica. With a layer of SiO$_2$ covering the Ni metals, a good activity was achieved at 600 °C over 24 h because of the enhanced MSI and confinement effect of mesoporous SiO$_2$ [182].
Al layer double hydroxide and MgO, metal sintering could be prevented to a large extent. Together with the protective layer of mesoporous silica, this catalyst maintained active for 72 h at 750 °C [188].

As for the controlled synthesis of Ni phyllosilicate, Ni precursors exerted an influence on the size of Ni particles, thus affecting the diffusion speed through the Ni phyllosilicate layer to form a new phyllosilicate species with silicate ions (seen in Table 4). Ni acetylacetonate, among various precursors, reacted with mesoporous silica spheres to form Ni phyllosilicates by hydrothermal method, exhibiting a unique pore structure with the smallest pore volume and pore size. This strong MSI between Ni and the support retarded carbon nanotube formation by anchoring Ni on the silica [147].

Besides the effect of precursors on the size of Ni, as shown in Table 4, modification of the porosity and basicity can further enhance the adsorption and accessibility of reactants. Therefore, Ni-Mg phyllosilicate were synthesized via facile hydrothermal method as shown in Figure 12. By adjusting the duration of treatment, an ideal porosity and basicity could be realized [187]. On one hand, due to the high affinity of MgO for CO\textsubscript{2}, more reactants were adsorbed on the surface more easily through a modified pore structure, accelerating the conversion of methane and the removal of carbon deposits. On the other hand, because of the strong MSI between Ni and MgO, metal sintering could be prevented to a large extent. Together with the protective layer of mesoporous silica, this catalyst maintained active for 72 h at 750 °C [188].

![Figure 11](image1.png)

**Figure 11.** DRM with highly coke-resistant silica-coated nickel nanoparticle catalysts. Reproduced with permission from reference [186].

![Figure 12](image2.png)

**Figure 12.** Schematic illustration of the formation process of Ni@Ni–Mg phyllosilicate core–shell catalyst. Reproduced with permission from reference [187].

Likewise, as shown in Table 4, NiMgAl layer double hydroxide was firstly synthesized by hydrothermal method, followed by coating a layer of mesoporous silica (Figure 13). After calcination and reduction, Ni nanoparticles were generated inside the cavity, which were confined by both MgO and silica layer. The synergies between the confinement effect of mesoporous silica and basicity of MgO resulted in an 8 h stable conversion at 750 °C with little carbon deposition [189].
The synergies between the confinement effect of mesoporous silica and basicity of MgO resulted in an 8 h stable conversion at 750 °C with little carbon deposition [189].

Besides Ni monometallic catalysts, the core–shell structures can be extended to Ni-based bimetallic catalysts. 12 nm Ni-Cu alloys were successfully synthesized by tuning the precursor concentrations. Being covered by the silica shell, the sintering of alloys could be alleviated, presenting a higher conversion than impregnated samples. Moreover, higher H2 selectivity could be achieved by the suppression effect on RWGS reaction with the addition of Cu (shown in Table 4) [190].

Ni-Co@silica core-shell structures calcined at 500 °C exhibited poor activities and stabilities. However, as presented in Table 4, when calcined at 800 °C, with the increase of crystallinity of SiO2 and enhanced MSI, the activity could be maintained stable over 1000 h, reflecting the high thermal stability of the support [191].

In a word, by designing the hierarchical structures of Ni-based catalysts, Ni particle size can be controlled, metal sintering can be alleviated, and coke resistance can be enhanced due to the confinement effect of silica shell and strong MSI derived from the Ni phyllosilicate. By doping another metal (Co, Cu, and Fe) to form alloys with Ni and adding Mg precursors to prepare Ni-Mg phyllosilicates, the activity and stability of the catalysts for DRM reaction can be further enhanced because of the promotional effects, such as the adsorption of CO2 and suppression of RWGS reaction.
Table 4. A summary of structural optimization effects on the catalytic performances of Ni-based catalysts for DRM reaction.

| Catalyst | Preparation Method | Reaction Condition | Methane Conversion | CO₂ Conversion | Carbon Formation | Comments | Ref. |
|----------|-------------------|--------------------|-------------------|----------------|-----------------|----------|------|
| NiCo/SiO₂ | Incipient wetness | 700 ºC; CH₄/CO₂ = 1 | 73% for CH₄ after 30 h | 77% for CO₂ after 30 h | n.d. | Homogeneous Ni-Co alloy supported on mesoporous silica was synthesized by oleylamine/oleic acid organic pair. | [133] |
| NiPhy (Ni phyllosilicate) hollow sphere (HS) | Hydrothermal method | 700 ºC; CH₄/CO₂ = 1 | 77.5% for CH₄ after 70 h | 86.6% for CO₂ after 70 h | <5% | Ni phyllosilicates exhibited a unique pore structure with the smallest pore volume and pore size. | [147] |
| Ni yolk@Ni@SiO₂ | Self-templating method | 700 ºC; CH₄/CO₂ = 1 | 74% for CH₄ after 50 h | 82% for CO₂ after 50 h | 1.4% | A yolk-shell Ni@SiO₂ catalyst increased the TOF by tuning surface area and Ni exposure | [149] |
| Ni@C | Wet impregnation method | 700 ºC; CH₄/CO₂ = 1 | 68% for CH₄ after 800 min | 80% for CO₂ after 800 min | 79.3% | Porous carbon anchored Ni²⁺ and enhanced dispersion. | [151] |
| Ni/SBA-15 | Modified impregnation method | 750 ºC; CH₄/CO₂ = 1 | 87% for CH₄ after 34 h | 92% for CO₂ after 34 h | 3.8% | Ni-EG complexes were pumped into the pores of SBA-15 by capillary force, hindering the migration of Ni. | [154] |
| Ni/SBA-15 | Modified impregnation method | 750 ºC; CH₄/CO₂ = 1 | 85% for CH₄ after 40 h | 88% for CO₂ after 40 h | 3.63 mg/gcat | Little sintering took place due to the confinement effect of SBA-15 with ordered channels. | [155] |
| Ni/KIT-6 | Impregnation method | 750 ºC; CH₄/CO₂ = 1 | 60% for CH₄ after 20 h | 78% for CO₂ after 20 h | NA | The micropores were occupied by Ni particles so that the pathways of reactants were blocked despite the high Ni dispersion. | [157] |
| Ni/HMS | S⁹ dickopal assembly pathways | 700 ºC; CH₄/CO₂ = 1 | 72% for CH₄ after 100 h | 84% for CO₂ after 100 h | 6.22% | Hexagonal mesoporous silica (HMS) improved the metal dispersion and inhibit sintering by forming Si-O-Ni bonds. | [158] |
| NiCo/SBA-15 | Co-precipitation method | 700 ºC; CH₄/CO₂ = 1 | 70% for CH₄ after 50 h | 78% for CO₂ after 50 h | n.d. | Small particle size (4–5 nm) and high metal dispersions (17–20%) were achieved by the confinement effect of ordered channel of SBA-15. | [159] |
| La₃Ni₂O₆/KIT-6 | Sol-gel method | 800 ºC; CH₄/CO₂ = 1 | 88% for CH₄ after 60 h | 100% for CO₂ after 60 h | 3.41% | Perovskite structures LaNiO₃ were also impregnated into the mesoporous SBA-15 supports. | [160] |
| Ni (Si₄O₉)₃/SiO₂ | Colloidal approach | 500 ºC; CH₄/CO₂ = 1.3 | 465 molCO₂/ h molNi | 450 molCO₂/ h molNi | NA | Colloidal approach was applied by converting Ni-silicide colloids into highly dispersed Ni nanoassemblies supported on porous silica. | [163] |
| Ni/SiO₂ | Incipient wetness impregnation method | 700 ºC; CH₄/CO₂ = 1 | 70% for CH₄ after 1000 min | 72% for CO₂ after 1000 min | n.d. | Co-ligands to protect Ni from agglomeration by ligand and hydrogen bond and steric hindrance. | [165] |
| Ni/SiO₂ | Incipient wetness impregnation method | 700 ºC; CH₄/CO₂ = 1 | 70% for CH₄ after 20 h | 78% for CO₂ after 20 h | 1.8 mg/gcat | With a small size, high dispersion and strong MSI, Ni/SiO₂ prepared with oleylamine showed the most stable catalytic performances. | [166] |
Catalyst & Preparation Method & Reaction Condition & Methane Conversion & CO₂ Conversion & Carbon Formation & Comments & Ref. \\
NiO-MgO-Al₂O₃ & evaporation induced self-assembly strategy & 700 °C; CH₄/CO₂ = 1 & 78% for CH₄ after 100 h & 83% for CO₂ after 100 h & 7.0% & Ni particles were anchored within the pores to alleviate the sintering effect and basicity of support could enhance CO₂ adsorption and remove coke. & [167] \\
NiO-CeO₂-Al₂O₃ & evaporation induced self-assembly strategy & 700 °C; CH₄/CO₂ = 1 & 78% for CH₄ after 80 h & 73% for CO₂ initially & 21.5% & The excellent thermal stability of this mesoporous structures protected Ni from sintering and the redox property of CeO₂ oxidized carbon deposits. & [171] \\
Silica@Ni@Silica & Ammonia evaporation method & 600 °C; CH₄/CO₂ = 1 & 44% for CH₄ after 24 h & 54% for CO₂ after 24 h & 7.6% & With a layer of SiO₂ covering the Ni metals, a good activity was achieved at 600 °C over 24 h & [180] \\
Ni@SiO₂ & Sol-gel process & 750 °C; CH₄/CO₂ = 1 & 54.1% for CH₄ after 25 h & 65.6% for CO₂ after 25 h & 1.2% & Strong resistance towards filamentous carbon deposition was exhibited; however, amorphous carbon still formed within the nanocavities & [183] \\
Ni@SiO₂ & Microemulsion method & 800 °C; CH₄/CO₂ = 1 & 90% for CH₄ after 90 h & 95% for CO₂ after 90 h & 4.8% & With a 11.2 nm of thickness, this unique structure exhibited an enhanced conversion and stability with the strong MSI and small satellite Ni particles. & [184] \\
Ni@SiO₂ & Reverse micelle method & 800 °C; CH₄/CO₂ = 1 & 87% for CH₄ after 100 h & 88% for CO₂ after 100 h & NA & Ni particles migrated into the SiO₂ shell, leading to the formation of uni-nuclear Ni core embedded in the shell. & [185] \\
Silica coated Ni/SiO₂ & Microemulsion method & 800 °C; CH₄/CO₂ = 1 & 42.5% for CH₄ after 170 h & 56.6% for CO₂ after 170 h & n.d. & To achieve a better anti-sintering property, a layer of silica was coated. & [186] \\
Ni@Ni-MgPhy & Hydrothermal method & 700 °C; CH₄/CO₂ = 1 & 78% for CH₄ after 95 h & 81% for CO₂ after 95 h & 4.4% after 20 h & By adjusting the duration of treatment, an ideal porosity and basicity could be realized. & [187] \\
Ni@Ni-MgPhy & Hydrothermal method & 750 °C; CH₄/CO₂ = 1 & 85% for CH₄ after 72 h & 89% for CO₂ after 72 h & n.d. & Strong MSI and the protective layer of mesoporous silica prevented metal sintering. & [188] \\
NiMgAl-(LDH)@mesoporous-SiO₂ & Hydrothermal and sol-gel method & 750 °C; CH₄/CO₂ = 1 & 79% for CH₄ after 8 h & 88% for CH₄ after 8 h & 2.5% & Ni nanoparticles were generated inside the cavity, which were confined by both MgO and silica layer. & [189] \\
Cu–Ni@SiO₂ & Microemulsion method & 700 °C; CH₄/CO₂ = 1 & 75% for CH₄ after 16 h & NA & NA & Being covered by the silica shell, the sintering of alloys could be alleviated. & [190] \\
NiCo@SiO₂ & Microemulsion method & 800 °C; CH₄/CO₂ = 1 & 90% for CH₄ after 1000 h & 90% for CO₂ after 1000 h & n.d. after 150 h & With the increase of crystallinity of SiO₂ and enhanced MSI, the activity could be maintained stable. & [191] \\

[^43]: AC: active carbon; HMS: Hexagonal mesoporous silica; LDH: layered double hydroxides.
6. Conclusions and Prospects

In this review, smart designs of Ni-based catalysts are discussed in four categories, including surface regulation, oxygen defects and interfacial engineering and structural optimization. In surface regulation, enhanced basicity can promote the CO$_2$ adsorption, resulting in active oxygen atoms which react with carbon species to form CO. Various oxides like La$_2$O$_3$, MgO, Ga$_2$O$_3$, and B$_2$O$_3$ can tune the surface basicity but too strong a basicity may inhibit carbon deposition. In oxygen defects, lattice distortion in perovskite structures, redox property of rare earth and lattice oxygen of transition metal oxides can promote the oxygen mobility, thus facilitating the oxidation of CH$_x$ and C deposits to form CO and H$_2$. In interfacial engineering, doping noble metals can enhance the reducibility of Ni, prevent Ni from agglomeration and impede the metal oxidation. Transition metals with oxygen affinity or redox cycle can absorb more CO$_2$ to produce CO and oxygen atoms, accelerating the carbon removal. Besides the Ni-metal synergy, a strong MSI can slow the reduction and retard the migration of Ni, thus leading to a high dispersion on the support, such as MgO and SiO$_2$. Moreover, electron perturbations of Ni from CeO$_2$ generate active Ni sites to adsorb and dissociate CH$_4$. On the other hand, the Ce$^{3+}$ promotes the C-O bond activation in CO$_2$ molecules to produce oxygen atoms, leading to an improved reactivity and stability in DRM reaction. In structural optimization, porous supports and core-shell structures can enhance the surface area and provide steric hindrance, leading to a highly dispersed Ni particle and more exposure to the reactants. In this case, the activation of reactants is facilitated, and metal sintering is inhibited. Despite the progress in designing Ni-based catalysts to resist coke and sintering, there is still room for further enhancement of the catalytic performance and a clear explanation of the mechanism.

First, a facile, economical, and green preparation strategy will be beneficial to the large-scale operations of DRM reaction in industry. Multi-step reaction processes may limit the yields of the catalyst materials. Also, the cost of the raw materials should be carefully controlled to an acceptable level. Furthermore, production and dispose of waste streams are another issue to be considered.

Second, novel techniques such as plasma, light and electricity assisted catalytic process can be a promising way to avoid the high temperature activation and metal sintering. Energy efficiency and stability may be enhanced in this way.

Third, combining with membrane, CO$_2$ can be separated from the raw feed and simultaneously converted into valuable syngas by reforming of methane. The recent advancement of highly CO$_2$-permeable membranes paves the way for developing an integrated system for both CO$_2$ capture and conversions [192–194].

Fourth, single-site or single-atom catalysts have drawn great interest in recent years due to the high efficiency of expensive catalysts and enhanced reactivity and selectivity in various reactions. For DRM reaction, single Ni atom catalysts have been proved to be highly active. However, how to prevent the agglomeration is still waiting to be resolved.

Fifth, by using CeO$_2$ as the support, low temperature or even room temperature activation of the DRM reactions have been reported. This strategy can save the energy input and alleviate the metal sintering. Considering the low equilibrium conversion at these conditions, an integrated system is to be developed for an efficient and immediate recovery of the unreacted molecules to realize the chemical looping process.

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