Role of Mixed Oxides in Hydrogen Production through the Dry Reforming of Methane over Nickel Catalysts Supported on Modified γ-Al₂O₃

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Abstract: H₂ production through dry reforming of methane (DRM) is a hot topic amidst growing environmental and atom-economy concerns. Loading Ni-based reducible mixed oxide systems onto a thermally stable support is a reliable approach for obtaining catalysts of good dispersion and high stability. Herein, NiO was dispersed over MOₓ-modified-γ-Al₂O₃ (M = Ti, Mo, Si, or W; x = 2 or 3) through incipient wetness impregnation followed by calcination. The obtained catalyst systems were characterized by infrared, ultraviolet–visible, and X-ray photoelectron spectroscopies, and H₂ temperature-programmed reduction. The mentioned synthetic procedure afforded the proper nucleation of different NiO-containing mixed oxides and/or interacting-NiO species. With different modifiers, the interaction of NiO with the γ-Al₂O₃ support was found to change, the Ni²⁺ environment was reformed exclusively, and the tendency of NiO species to undergo reduction was modified greatly. Catalyst systems 5Ni₃MAl (M = Si, W) comprised a variety of species, whereby NiO interacted with the modifier and the support (e.g., NiSiO₃, NiAl₂O₄, and NiWO₃). These two catalyst systems displayed equal efficiency, >70% H₂ yield at 800 °C, and were thermally stable for up to 420 min on stream. 5Ni₃SiAl catalyst regained nearly all its activity during regeneration for up to two cycles.

Keywords: H₂ production; mixed oxide; NiAl₂O₄; NiWO₃; NiSiO₃; NiO-interacting species

1. Introduction

Hydrogen (H₂) is a green energy source. In fact, the energy content of this molecule is three times higher than that of gasoline [1]. H₂ is a critical feedstock for the chemical industry in oil refining, fertilizer production, and fuel cell manufacture [2]. Notably, biomass pyrolysis and biomass gasification have always been environmentally questionable approaches to hydrogen production; moreover, the implementation of these methods renders H₂ isolation and purification quite difficult, as the desired product associates with many side-products [3]. A more convenient approach is hydrogen production from compounds such as ethanol [4], glycerol [5], glucose [6], starch, and catechol [7], by steam or thermal reforming over a heterogeneous catalyst. Environmental concerns and the high demand for atom-economies have driven the development of hydrogen production methods, relying...
on clean sources, such as water splitting, thermal reforming of methane, stream reforming of methane, and CO\textsubscript{2} (dry) reforming of methane (DRM) \cite{8,9}. DRM in particular, has drawn great deal of attention globally because it fulfils the goal of hydrogen production while raising hope for the reduction of the atmospheric concentrations of carbon dioxide and methane, the two gases whose emissions are most responsible for global warming.

From a catalytic standpoint, metal oxide and mixed metal oxide systems have attracted the attention of the industry, due to their ease of formation, development of mixed oxide potential, tuneable surface topology, high temperature sustainability, and ease of handling in drastic industrial conditions \cite{10}. Ni-based catalysts supported on thermally-stable neutral SiO\textsubscript{2} \cite{11}, acidic Al\textsubscript{2}O\textsubscript{3} \cite{12}, and reducible TiO\textsubscript{2} \cite{13,14} have been investigated thoroughly in recent years. Due to the need of acid-base bi-functional catalyst for DRM, TiO\textsubscript{2} and Al\textsubscript{2}O\textsubscript{3} supports were investigated more than SiO\textsubscript{2}. Reducibility characteristics of TiO\textsubscript{2} controls the coke deposition. However, the coverage of Ni species by the TiO\textsubscript{2} and partial transformation of amorphous titania phase into the anatase phase limit its DRM potential, tuneable surface topology, high temperature sustainability, and ease of handling.

Incorporation of Cu, Mo, Si, Sr, B, Ca, Mg, Ba, Ga, Gd, Zn, Ti, W, Mn, Co, Zr, Ce, La, and Yb, in the alumina supported Ni catalyst was examined in the search of high catalytic activity due to its abundant confined spaces and steady chemisorption behavior. Under other synthetic methodology, such as evaporation-induced self-assembly method \cite{20} and microwave-assisted combustion \cite{21} and atomic layer deposition \cite{22} method have drawn attentions. As a sandwiched catalyst Al\textsubscript{2}O\textsubscript{3}/Ni/Al\textsubscript{2}O\textsubscript{3} (prepared by atomic layer deposition) prevented Ni gathering over alumina by coating of porous Al\textsubscript{2}O\textsubscript{3} thin-film over Ni/Al\textsubscript{2}O\textsubscript{3}. Moreover, Ni had double interaction with Al\textsubscript{2}O\textsubscript{3} in this catalyst system and so it was claimed nearly 100% CO\textsubscript{2} and CH\textsubscript{4} conversion with excellent H\textsubscript{2} and CO selectivity \cite{19}. However, Ni-based catalysts supported on nanofiber exhibited superior catalytic stability due to its abundant confined spaces and steady chemisorption behavior.
CH$_4$ and CO$_2$ conversion up to 300 min) due to formation of MgAl$_2$O$_4$ mixed oxide [30]. Promotional addition of Ga, Gd and Zn claimed high CH$_4$ and CO$_2$ conversions (close to 80% during 420 min time on stream) with H$_2$/CO = 1.0 due to content of NiO interacted moderately and strongly with the support [24]. Co-addition to Ni/Al$_2$O$_3$ nano catalyst showed high Ni/Al ratio than Ni/Al$_2$O$_3$ which led to better Ni dispersion over the alumina surface (by reducing the ensemble size of Ni) [23]. It was claimed good catalytic activity (CO$_2$ and CH$_4$ conversion >80% at 750 °C) with H$_2$/CO ratio was close to one and less coke deposition. Mn-addition caused partial blockage of large Ni particle which was assumed for rapid coke formation. Further addition of potassium caused stability in activity [31]. ZrO$_2$ addition enhanced the dissociation of CO$_2$ near the contact between ZrO$_2$ and nickel where the deposited coke was gasified afterwards [32]. Optimum ceria addition caused formation of CeAlO$_3$ mixed phase. Ceria added mobile lattice oxygen in the surface reaction mechanism of DRM which endorsed minimum coking [33,34]. The addition of lanthana enhanced Ni stability and alkalinity of catalyst [35]. Alkalinity caused profound carbon dioxide adsorption, activation and oxidation of carbon deposit. The addition of 1–2 wt% Yb in NiO-containing Al$_2$O$_3$ (prepared by sol-gel process) matrix controlled the size/mobility of Ni nanoparticles. Moreover, the high ease of reducibility and low carbon deposition led to high catalytic activity and catalyst stability (CH$_4$ conversions 75–81.5% and CO$_2$ conversions 83.5–89% for 22 h time on stream), and H$_2$/CO close to unity (~0.95) [36]. The active elements in doped/promoted Al$_2$O$_3$-supported Ni catalyst systems are presented as the color blocks in periodic table in Figure 1.

Figure 1. Active elements in a doped/promoted Al$_2$O$_3$-supported Ni catalyst system for dry reforming of methane.

In these systems, apart from the dispersed, stable NiO, other Ni-containing phases, like Ni$_2$SiO$_4$, NiAl$_2$O$_4$, MgAl$_2$O$_4$, CeAlO$_3$, NiTiO$_3$, and Ni$_2$TiO$_3$, have also been detected. The presence of NiWO$_4$ [37] and NiMoO$_4$ species [38] has also been reported in Ni-based catalysts supported on WC or MoC, respectively, for DRM. In this context, understanding the role played by different metal oxides or mixed metal oxides matrices in the production of hydrogen through DRM is imperative. Recently, we prepared a modified γ-Al$_2$O$_3$ support by incipient wetness impregnation followed by calcination; notably, the modifier was loaded as 3.0 wt% MO$_x$ (M = W, Mo, Si, or Ti; x = 2 or 3) [39]. The active Ni catalyst was then loaded onto the modified γ-Al$_2$O$_3$ support by incipient wetness impregnation, followed by calcination to reach 5.0 wt% NiO loading. The catalysts, thus obtained, are abbreviated as 5Ni3MAI (M = W, Mo, Si, or Ti). This synthetic strategy gave the chance of interaction of all types of oxides and, hence, the formation of mixed metal oxides. The study of this type of catalytic systems may provide information that would help understand the role of metal oxides and mixed metal oxides in DRM. Herein, we report the preparation of the 5Ni3MAI (M = W, Mo, Si, or Ti) catalyst systems, their characterization by infrared (IR), ultraviolet–visible (UV-Vis), X-ray photoelectron (XPS) spectroscopies,
and by H₂ temperature-programmed reduction (H₂-TPR). We also report the results of tests run on these systems for H₂ production. We attempted to identify the metal, metal oxides, and mixed metal oxides responsible for H₂ production through DRM. Crucially, this information would be useful in the design of metal oxide matrix catalyst systems suitable for the industrial production of H₂ through DRM.

2. Results

2.1. Catalytic Activity

Data reflecting the catalytic activity of the prepared catalysts, in terms of H₂ yield over 420 min on stream at 700 °C are reported in Figure 2A. On the other hand, Figure 2B shows values for the H₂ yield at different reaction temperatures. Among the investigated catalysts, 5Ni3SiAl and 5Ni3WAI exhibited the highest catalytic performance, and the activities and stabilities of these two species were found to be identical. The H₂ yield recorded for both catalysts over 420 min on stream was about 62% at 700 °C. The H₂ yield over 5Ni3MoAl catalyst was 47% after 20 min on stream, which decreased to 39% after 420 min on stream. The catalytic performance of 5Ni3TiAl was found to be the worst: ~30% H₂ yield after 420 min on stream. Notably, the H₂ yield in the presence of all catalyst systems increased as the reaction temperature increased. At 800 °C, the use of 5Ni3SiAl and 5Ni3WAI was associated with more than 70% H₂ yield, whereas in the case of 5Ni3MoAl and 5Ni3TiAl, H₂ yield was measured to be around 60%. Higher conversion trend at higher temperature indicated the endothermic nature of DRM reaction. For silicon- or tungsten-modified alumina-supported nickel catalyst system, the rise of H₂ yield with temperature were similar. For titanium- and molybdenum-modified catalyst systems, the rise of H₂ yield with temperature were similar up to 650 °C, and after this temperature, molybdenum-modified catalyst system showed better H₂ yield.

![Figure 2.](image)

**Figure 2.** (A) H₂ yield over different catalyst systems up to 420 min TOS at 700 °C; (B) H₂ yield over different catalyst systems in the 500–800 °C reaction temperature range.

2.2. Result

SEM images and EDX patterns of 5Ni3MAI (M = W, Si, Mo, Ti) are shown in Figures S1 and S2, respectively. EDX patterns showed the presence of all claimed elements in the catalyst systems (Figure S2). In all IR spectra, the sharp peak at 1644 cm⁻¹ and the broad intense peak at 3450 cm⁻¹ could be due to the bending and stretching vibrations, respectively, of adsorbed water molecules or surface hydroxyl groups [40]. The IR peaks in the 600–750 cm⁻¹ wavenumber range could be attributed to the vibrations of AlO₆ octahedral units (Al-O bonds in an octahedral environment), whereas those in the 750–850 cm⁻¹ range could be attributed to AlO₄ tetrahedral units (Al-O bonds in a tetrahedral environment) [41]. The three peaks in the 1640–1350 cm⁻¹ wavenumber range
might be assigned to the bending vibration of the water molecules coordinated to different unsaturated surface sites. Furthermore, the peak at 1635 cm$^{-1}$ could be due to the vibration of physically adsorbed water molecules, whereas the peak at 1380 cm$^{-1}$ could be due to the bending vibration of water molecules coordinating tetrahedral aluminum [40]. The IR spectrum of γ-Al$_2$O$_3$ was characterized by the presence of three peaks at 1635 cm$^{-1}$, 1517 cm$^{-1}$, and 1382 cm$^{-1}$ [42]. IR peaks in the 400–470 cm$^{-1}$ wavenumber range could be due to the vibration of the Ni-O bond [43] in free NiO or to the said bond in mixed oxides such as NiSiO$_4$, NiAl$_2$O$_4$, NiTiO$_3$, NiWO$_4$, and NiMoO$_4$. The IR spectrum of the titanium-modified catalyst sample, 5Ni3TiAl, (Figure 3A,B and Figure S3A,B), comprised an absorption peak at 433 cm$^{-1}$, which could be due to the Ni-O vibration of free NiO species in a cubic lattice, and peaks at 455 cm$^{-1}$ and 547 cm$^{-1}$, which may be due to the Ni-O bond vibration of NiTiO$_3$ [44,45]. After NiO loading, the peak due to TiO$_2$ in the low-frequency region of the spectrum (below 500 cm$^{-1}$) was replaced by peaks due to Ni-O vibrations, which was an indication of the extent of interaction of NiO species with the modifier over the support.

The IR spectrum of the molybdenum-containing catalyst, 5Ni3MoAl, (Figure 3C,D and Figure S3A,B), comprised a band at 424 cm$^{-1}$ due to Ni-O bond vibration, which appeared at lower wavenumber values than that attributed to free NiO (433 cm$^{-1}$) in the spectrum of the titanium-modified catalyst. No other prominent peaks due to Ni-O vibrations were noticed in the spectrum of the molybdenum-containing catalyst, indicating the deformation of cubic NiO species and the formation of species, whereby NiO interacted with the support, the modifier, or both [46,47]. After NiO loading, the peak due to MoO$_3$ in the low-frequency region was completely replaced by the vibrational peak of Ni-O, indicating the extent of the interaction of NiO species with the modifier over the support.

In the IR spectrum of 5Ni3SiAl (Figure 3E,F and Figure S3A,B), Ni-O vibrational peaks at 430 cm$^{-1}$, due to free NiO, and 403 cm$^{-1}$, due to strongly interacting-NiO (as a result of the weakening of the Ni-O bond), were noticed, alongside a peak at 453 cm$^{-1}$, due to the Si-O-Si bending vibration [48], and one at 481 cm$^{-1}$, due to the asymmetric vibration of Si-O [49]. However, the S-O asymmetric vibration peak was not noticed in the spectrum of SiO$_2$-Al$_2$O$_3$, indicating that this vibration was related to an interaction of NiO with silica species or to the presence of amorphous nickel silicate species. This finding is in agreement with our previous claim of an interaction of NiO species, as deduced from H$_2$-TPR data, and the absence of nickel silicate phases, as deduced from X-ray diffraction data. The IR peak and its shoulder appearing in the 549–560 cm$^{-1}$ wavenumber range could be attributed to the presence of Al-O-Ni bonding interactions in NiAl$_2$O$_4$ [50].

In both 5Ni3SiAl and 5Ni3WAl, following NiO loading, the infrared spectra of the modified catalysts became populated with intense peaks, indicating the presence and dispersion of a wide variety of species, whereby NiO interacted with catalyst components on the catalyst surface. In the IR spectrum of 5NiWAl (Figure 3G,H) the low-intensity absorption bands below 500 cm$^{-1}$ could be attributed to the stretching vibrations of NiO$_6$ polyhedral in NiWO$_4$ species [51].

The infrared spectra of the catalyst samples showed a clear shift of Ni–O peak from 433 cm$^{-1}$ (for 5Ni3TiAl) to 424 cm$^{-1}$ (for 5Ni3MoAl) and 403 cm$^{-1}$ (for 5Ni3SiAl). That reflects the presence of “free NiO” species in 5Ni3TiAl; NiO species interacted with the modifier in 5Ni3MoAl and NiO species strongly interacted with the support/modifier in 5Ni3SiAl. 5Ni3WAl catalyst samples had a wide interaction NiO species as it showed stretching vibration of NiO$_6$ polyhedral.
Figure 3. (A–H) Infrared spectra of different catalyst systems. In (D), Ni-O$^a$ indicates NiO species that interacted with the support. In (F) Ni-O$^b$ indicates NiO species that strongly interacted with the support. In (H) inset indicates stretching vibration peaks of NiO$_6$ polyhedra in NiWO$_4$ species.
UV–Vis spectroscopy is the most suitable characterization technique to understand the electronic environment of Ni^{2+} or d^{8}-configuration systems. The UV–Vis absorption spectra of all catalyst samples are reported in Figure 4. The peak observed at 250–350 nm is associated with the charge transfer transition from O^{2−} to Ni^{2+} in an octahedral coordination environment (O^{2−} → Ni^{2+}) in the NiO lattice [29]. The peak centered at around 410 nm is associated with the d-d transition from the $^3A_{2g}(F)$ state to the $^3T_{1g}(P)$ state of Ni^{2+} in an octahedral environment, whereas the doublet peak with maxima at 593 nm and 634 nm were attributed to the d-d transition from the $^3T_1(F)$ state to the $^3T_1(P)$ state of Ni^{2+} in a tetrahedral environment [52]. Among all four catalysts, only the silicon- and tungsten-containing catalysts (5Ni3SiAl and 5Ni3WAl) included Ni^{2+} ions in both tetrahedral and octahedral environments. The NiAl_{2}O_{4} phase is known to comprise Ni^{2+} ions in both tetrahedral and octahedral coordination environments. Therefore, 5Ni3SiAl and 5Ni3WAl may contain NiAl_{2}O_{4} phases. In the case of the other catalysts, the mentioned bands either displayed low intensity or had shifted toward higher wavelengths, indicating a weak interaction between the modifier and Ni^{2+} and the support. In the case of the titanium-containing catalyst (5Ni3TiAl), no peaks attributable to Ni^{2+} ions in tetrahedral environments were observed, indicating the absence of an NiAl_{2}O_{4} phase. By contrast, the UV-Vis spectrum of 5Ni3TiAl displayed a peak at 740 nm attributable to Ni^{2+} in an octahedral coordination environment, as segregated or crystallized NiO which was as near as bulk NiO [53], indicating that octahedral coordinated Ni^{2+} was present in segregated form in 5Ni3TiAl. An additional hump in the 260–330 nm wavelength range for 5Ni3TiAl could be assigned to a charge transfer from O^{2−} to octahedral/tetrahedral Ti^{4+} (O^{2−} → Ti^{4+}) in a possible NiTiO_{3} phase [44]. The hump in the 280–330 nm wavelength range observed in the UV–Vis spectrum of 5NiMoAl could be attributed to a charge transfer from O^{2−} to octahedral/tetrahedral Mo^{6+} (O^{2−} → Mo^{6+}) in a possible NiMoO_{4} phase/Al_{2}(MoO_{4})_{3} phase [54].

**Figure 4.** Ultraviolet–visible spectra of different catalyst systems. Peak at 410 nm is associated with d–d transition $^3A_{2g}(F) → ^3T_{1g}(P)$ in Ni^{2+} in an octahedral environment. Peaks at 593 and 634 nm are associated with the d–d transition $^3T_1(F) → ^3T_1(P)$ in Ni^{2+} in a tetrahedral environment.

In order to clarify further the interaction of Ti and Mo with the support and with catalytic active sites (NiO), Al (2p, 2s) XPS analysis was conducted on 5Ni3TiAl and 5Ni3MoAl (Figure 5A). The Al (2p) XPS spectrum of 5Ni3TiAl was characterized by peaks
at 74.30 eV and at 119.17 eV. These peaks were quite similar to those observed in the Al (2p) XPS spectrum of pure γ-Al₂O₃ [55]. This evidence indicated that the γ-Al₂O₃ support doesn’t interact with TiO₂ and NiO. However, in the presence of molybdenum as a modifier, a substantial shift to lower energy values (to 71.5 eV and 116.34 eV) was observed for the mentioned peaks in the Al (2p) spectra. This shift was indicative of changes in the electronic environment around Al ions, resulting from an increase in negative charge in the vicinity of Al³⁺ due to an excess of oxygen atoms around it. It may indicate an oxide enrichment resulting from the formation of Al₂(MoO₄)₃ [56]. However, the absence of peak shifts to higher energy in the case of the Al (2p) and Al (2s) XPS spectra of 5Ni3MoAl indicates a poor interaction between NiO and the γ-Al₂O₃ support (i.e., the formation of a NiAl₂O₄ phase was not observed) [57,58]. Notably, Mo (3d) XPS analysis was also carried out on 5Ni3MoAl (Figure 3B). The Mo (3d) XPS spectrum of this catalyst showed a doublet peak with maxima at 234.75 eV and 235.8 eV, which is attributable to the interaction of NiO with MoO₃ species [59,60], and a peak at 233.52 eV, which is attributable to a strong interaction between MoO₃ and γ-Al₂O₃ or the formation of Al₂(MoO₄)₃ [61]. Overall, 5Ni3TiAl can be said to comprise negligibly interacting species, whereas 5Ni3MoAl may include species whereby NiO interacted with MoOₓ, and MoO₃ interacted with γ-Al₂O₃, although little interaction appeared to exist between NiO and the γ-Al₂O₃ support [62].

Figure 5. (A) Al (2p, 2s) X-ray photoelectron spectra of catalysts 5Ni3TiAl and 5Ni3MoAl and (B) Mo (3d) X-ray photoelectron spectrum of 5Ni3MoAl.

Till now, catalysts 5Ni3SiAl and 5Ni3WAl were thus found to be enriched with NiO species interacting with the support and the modifier. The tendency of these species to undergo reduction needs, however, to be determined because only reduced nickel species (i.e., metallic nickel) can initiate CH₄ decomposition for hydrogen production through DRM [63,64]. In order to determine their reducibility, H₂-TPR experiments were carried out for 5Ni3SiAl and 5Ni3WAl catalysts (Figure 6). The deconvoluted peak profiles are shown in Figure S4. These reduction peaks can be categorized into four regions; region I 200–400 °C, region II 400–700 °C, region III 700–900 °C, and region IV 900–1000 °C [65]. The H₂-TPR profile of 5Ni3SiAl was characterized by a low prominent reduction peak in region I and largely overlapped peaks in regions II and III. The less prominent peak in region I was due to reduction of free NiO species. Region II showed reduction peak for NiO interacting with the modifier or NiSiO₃, whereas region III showed reduction peaks for NiO interacting with the support or NiAl₂O₄ [55,59]. Formation of NiAl₂O₄ was explained by Scheffer et al. [66]. Nickel ions diffused into the Al₂O₃ support, where it was coordinated tetrahedrally and octahedrally to form NiAl₂O₄. UV of 5Ni3SiAl sample also detected Ni²⁺ in tetrahedral and octahedral coordination. On the other hand, the H₂-TPR profile of 5Ni3WAl catalyst was characterized by absence of region I peak (for free NiO)
and an additional peak in region IV [67]. The absence of free NiO species suggested that some interaction was occurred between NiO and WO₃ or Al₂O₃ species. Scheffer et al. discussed that from NiAl₂O₄ species some of aluminium ions may be replaced and form new NiO species (or NiWOAl) and the reduction peak of this new NiO species would be at higher temperature beside reduction peak of NiAl₂O₄ species. Thus, additional peak in region IV can be claimed to reduction peaks of NiWOAl species. Further, the reduction peaks in regions II and III were largely overlapped in which region III was more prominent. Region II showed reduction peaks for NiWO₄ species, whereas region III showed reduction peaks for NiAl₂O₄ [56,60]. Overall, it can be said that 5Ni3SiAl catalyst system had NiSiO₃ and NiAl₂O₄ species, whereas 5Ni3WAl catalyst system had NiWO₄ and NiAl₂O₄ species majorly.

3. Discussion

The chief route of H₂ production via DRM comprises the dissociation of CH₄ over the surface of Ni (or supported Ni) to produce carbon deposits and H₂, followed by the oxidation of the carbon deposits by CO₂ to produce CO (CH₄ + CO₂ → CO + H₂). Nevertheless, the reverse water gas shift (RWGS) reaction due to a spill over effect of molecular hydrogen, produced over the surface of Ni, (CO₂ + H₂ → CO + H₂O) and the gasification of the carbon deposits by hydrogen gas (C + 2H₂ → CH₄) [68] are not reaction routes that can be ignored. In fact, these parallel reactions can affect H₂ yield. Notably, not all parallel reactions, taking place over the catalyst surface, are detrimental to the mean yield of H₂. For instance, the presence of water at the catalyst surface (possibly due to RWGS) may contribute to the gasification of carbon deposits, leading to the production of molecular hydrogen and carbon monoxide (C + 2H₂O → CO + H₂), and thus increasing H₂ yield.

The dispersion and stability of “Ni-related species” over a support determine the H₂ yield of a DRM reaction, taking place over the surface of a catalyst. The 5Ni3TiAl catalyst was characterized by infrared-active Ni-O vibration frequencies close to those of free NiO; in fact, NiO interacted with the modifier to produce the characteristic peaks of NiTiO₃. UV–Vis absorption evidence also confirmed the presence of Ni²⁺ ions in an octahedral environment, in the form of segregated, crystallized, or bulk NiO, and the presence of Ti⁴⁺ in both tetrahedral and octahedral environments in NiTiO₃. Overall, 5Ni3TiAl comprised free NiO and NiTiO₃. XPS Al (2p) evidence on 5Ni3TiAl also confirmed the intact nature of TiO₂ and NiO over the Al₂O₃ support. The inferior catalytic performance of 5Ni3TiAl with respect to those of the other systems could be due to the free NiO species present in it. Use of 5Ni3TiAl as DRM catalyst was observed to be associated with a 30% H₂ yield at 700 °C and more than 60% at 800 °C. The IR spectrum of 5Ni3MoAl did not include any peaks attributable to the vibration of the Ni-O bond of free NiO. The UV–Vis spectrum of 5Ni3MoAl was indicative of the presence of Mo⁶⁺ in octahedral/tetrahedral environments, as environment of Mo⁶⁺ in NiMoO₄. XPS Mo (3d) evidence on 5Ni3MoAl
indicated the presence of a species, whereby NiO interacted with MoO\textsubscript{x}, which was the dominant one, with a smaller prevalence of a species whereby NiO interacted with the support (\gamma-Al\textsubscript{2}O\textsubscript{3}). The absence of free NiO in 5Ni3MoAl caused a larger fraction of Ni-interacted species with the modifier (as NiMoO\textsubscript{4}) at the surface than that at 5Ni3TiAl. An increased fraction of species whereby Ni interacted with MoO\textsubscript{x} (i.e., NiMoO\textsubscript{4}) caused the catalyst to increase in stability and resulted in >45% H\textsubscript{2} yield of the DRM reaction at 700 °C. At high reaction temperature, about 800 °C, an endothermic feature of DRM reaction promoted enhanced CH\textsubscript{4} decomposition at “Ni-interacted species surfaces” as NiMoO\textsubscript{4} and NiTiO\textsubscript{3} \cite{69}. Therefore, NiMoO\textsubscript{4} and NiTiO\textsubscript{3} species (in 5Ni3MoAl and 5Ni3TiAl catalyst systems, respectively) displayed high catalytic efficiency and thermal stability at high temperatures, with an over 60% H\textsubscript{2} yield of the DRM reaction at 800 °C.

The IR spectra of 5Ni3SiAl and 5Ni3WAl were characterized by vibrational frequencies due to a variety species, whereby NiO interacted with the modifier or the support. 5Ni3SiAl comprised species, whereby NiO interacted with the modifier to form NiSiO\textsubscript{3} and with the support to produce NiAl\textsubscript{2}O\textsubscript{4}, much like 5Ni3WAl comprised species, whereby NiO interacted with the modifier to form NiWO\textsubscript{4} and with the support to form NiAl\textsubscript{2}O\textsubscript{4}. UV-Vis absorbance data confirmed the presence of Ni\textsuperscript{2+} in tetrahedral and octahedral environments in NiAl\textsubscript{2}O\textsubscript{4} in both catalysts. H\textsubscript{2}-TPR evidence validated the ability of these “interacting-Ni species” in both 5Ni3SiAl and 5Ni3WAI catalyst systems to undergo reduction. As both catalysts comprised a variety of stable interacting-NiO species, they displayed equally high catalytic efficiency. Use of both catalysts in the DRM reaction resulted in >60% H\textsubscript{2} yield at 700 °C and >70% H\textsubscript{2} yield at 800 °C over 420 min on stream. Evidence suggested that the NiSiO\textsubscript{3}, NiWO\textsubscript{4}, and NiAl\textsubscript{2}O\textsubscript{4} mixed oxides present in 5Ni3SiAl and 5Ni3WAI were more thermally stable than the NiTiO\textsubscript{3} and NiMoO\textsubscript{4} mixed oxides present in 5Ni3TiAl and 5Ni3MoAl, respectively. The catalyst system, composed of 12.5 wt% Ni and 12.5 wt% Co and supported on La\textsubscript{2}O\textsubscript{3}, showed 31% and 60% H\textsubscript{2} yield for the catalyst calcined at 700 °C and 900 °C, respectively \cite{70}. The 5 wt% MgO-promoted Ni-Co/Al\textsubscript{2}O\textsubscript{3}-ZrO\textsubscript{2} nanocatalyst showed about 20% H\textsubscript{2} yield at 650 °C and 40% hydrogen yield at 750 °C \cite{71}. Ni catalysts supported on Gd-doped ceria (prepared by conventional impregnation method) showed 64% H\textsubscript{2} yield \cite{72}. The Sr-promoted Al\textsubscript{2}O\textsubscript{3} supported Nickel catalyst showed 70% H\textsubscript{2} yield \cite{73}. In our catalyst system, only 3 wt% promoter of either W or Si in Al\textsubscript{2}O\textsubscript{3}-supported nickel catalyst resulted into more than 60% H\textsubscript{2} yield at 700 °C and more than 70% H\textsubscript{2} yield at 800 °C. In the regeneration study with 5Ni3SiAl catalyst, the H\textsubscript{2} yields after the first and second recycling were 58 and 59%, respectively at 700 °C. It can be inferred that the catalyst regained nearly all its performance after the removal of carbon deposits by using oxygen (Figure 7).

![Figure 7. H\textsubscript{2} yield over fresh 5Ni3SiAl catalyst, H\textsubscript{2} yield after first cycle (R1), and H\textsubscript{2} yield after second cycle (R2).](image)
4. Materials and Methods

4.1. Catalyst Preparation

The 5Ni3MAl (M = Si, Ti, Mo, or W) catalyst systems were prepared by incipient wetness impregnation, followed by calcination. The detailed procedure for catalyst preparation is described as follows. The required amounts of well-grounded γ-Al$_2$O$_3$ and metal oxides precursors ((NH$_4$)$_6$Mo$_7$O$_{24}$·4H$_2$O (Sigma-Aldrich, St. Louis, MO, USA) or (NH$_4$)$_{10}$H$_2$(W$_2$O$_7$)$_6$ (Aldrich)) were mixed together and then crushed mechanically by mortar and pestle. Droplets of deionized H$_2$O was added to form a paste. The mechanical mixing was carried out until the dried compact mixture was accomplished. Wetting and drying processes of the solid mixture were done three times. The dried mixture was calcined in a programmable muffle furnace at 600 °C at the rate 3 °C/min for three hours. γ-Alumina with titania (3.0 wt% TiO$_2$/γ-Al$_2$O$_3$) and silica (2.0 wt% SiO$_2$/γ-Al$_2$O$_3$) in the form of pellets, were gifts from the Inorganic Chemistry Laboratory, Oxford University, Oxford, UK. Same as the above discussed procedure, the essential amount of Ni(NO$_3$)$_2$.6H$_2$O along, with doped γ-Al$_2$O$_3$ support were grounded, were made in paste form, and then were dried and calcined. The calcined 5 wt% NiO /3 wt% MOx-γ-Al$_2$O$_3$(M = Ti, Mo, W, Si) catalysts were abbreviated as 5Ni3MAl (M = Ti, Mo, W, Si).

4.2. Catalyst Characterization

Catalysts were characterized by SEM, EDXS, IR, UV–Vis, and X-ray Photoelectron spectroscopy techniques and H$_2$-TPR. The morphology of the catalyst samples was investigated by using a field emission scanning electron microscope (FE-SEM, model: JEOl JSM-7100 F, St. Lucia, Queensland, Australia), furnished with energy dispersive X-ray spectroscopy (EDXS) for surface elemental analysis. The Fourier transform infrared (FTIR) measurements were carried out by using IR Prestige-21 SHMADZU, spectrophotometer, Kyoto, Japan. The spectra were documented in the range 400-4000 cm$^{-1}$ with 4 cm$^{-1}$ energy resolution, using KBr pellet. Ultraviolet-visible measurement was carried out by using V-570 JASCO, Eston, PA, USA) spectrophotometer. The spectra were registered in the range of 200–800 nm wavelength with resolution of 1 nm at a scanning speed of 200 nm/min. The XPS was carried out using Thermo Scientific X-ray Photoelectron Spectrometer, Manchester, UK. Monochromatic Al Kα (1486.6 eV) radiation source operating at a power of 72 W with a pass energy of 50 eV for high resolution area scans and 200 eV for inspection scans was used. For charge adjustment, a one-point scale with the C1s peak moved to 285.0 eV was used. Temperature-programmed reduction (TPR) was carried out with the Micromeritics AutoChem II, Atlanta, GA, USA. 0.07 g of the sample (in sample holder) was degassed using argon at 150 °C for 60 min and then cooled to 25 °C. Further, it was heated to 800 °C at the rate of 10 °C/minute under 10% H$_2$ in argon flow. The flow rate of 10% H$_2$ in argon was set at 40 mL/minute. At the reactor outlet, thermal conductivity detector (TCD) monitors the gas mixture and peaks corresponding to the consumption of H$_2$ as a function of temperature were obtained.

4.3. Catalytic Reaction

The reaction was performed at atmospheric pressure in a tubular, stainless steel, fixed-bed micro-reactor (ID = 9 mm) (supplied by PID Eng and Tech, Madrid, Spain). 0.10 g catalyst was reduced under 20 mL/min flow of H$_2$ at 800 °C for 60 min. Now, nitrogen gas was passed through the reactor for 20 min to eliminate adsorbed H$_2$ at 700 °C. Afterwards, dry reforming of methane was carried out by passing CH$_4$, CO$_2$, and N$_2$ gas mixture feed at flow rates of 30, 30, and 10 mL/min respectively through the catalyst bed. The temperature, pressure and reaction variables were monitored through the reactor panel. GC-2014 (SHIMADZU, Kyoto, Japan) gas chromatograph equipped with Porapak Q and Molecular Sieve 5A columns and thermal conductivity detector was joined in series/bypass connections to have a whole examination of the reaction products and the feed. For regeneration process, fresh 5Ni-3SiO$_2$ + Al$_2$O$_3$ catalyst is packed in the reactor. The sample was activated with H$_2$ flow of 20 mL/min for an hour and then followed by a
reaction for 820 min at 700 °C. The spent catalyst is then treated with 10 mL/min O2 for 30 min to remove the deposited carbon. Subsequently, the recycled catalyst was activated with H2 to take care of any oxidized Ni metal during the recycling and thereafter, reaction was performed. Again, the recycling operation was repeated and the reaction carried out the third time.

5. Conclusions

The titanium-modified catalyst (5Ni3TiAl) had free NiO and NiTiO3 species. Free NiO species were completely unsuitable for H2 production through DRM. The presence of the free NiO fraction caused 5Ni3TiAl to display about 30% H2 yield at 700 °C. The molybdenum-modified catalyst (5Ni3MoAl) had no free NiO species, but had NiO interacting with MoO3 species or NiMoO4 species, which displayed a moderate catalytic performance of about 45% H2 yield at 700 °C. Notably, at a higher reaction temperature (800 °C), the endothermic feature of the DRM reaction enhanced CH4 decomposition, and both catalysts displayed a higher but equal efficiency of about 60% H2 yield. On the other hand, the silicon-modified catalyst (5Ni3SiAl) and tungsten-modified catalyst (5Ni3WAl) were comprised of a variety of thermally stable species, whereby NiO interacted with the modifier and the support. The 5Ni3SiAl catalyst system had NiSiO3 and NiAl2O4 species, whereas the 5Ni3WAI catalyst system had NiWO4 and NiAl2O4 species. The sets of mixed oxides present in 5Ni3SiAl and 5Ni3WAI were responsible for displaying higher thermal stability and catalytic performance than 5Ni3TiAl and 5Ni3MoAl. In fact, the 5Ni3SiAl and 5Ni3WAI catalyst systems displayed equally high catalytic efficiency of about 60% H2 yield at 700 °C and about 70% H2 yield at 800 °C, over 420 min on stream. The 5Ni3SiAl catalyst regained nearly all its activity after the removal of carbon deposits by using oxygen.

Supplementary Materials: The following are available online at https://www.mdpi.com/2227-9717/9/1/157/s1, Figure S1: SEM image of different catalyst system; Figure S2: EDX spectra of different catalyst system, Figure S3: Infrared spectroscopy of different catalyst system, Figure S4: Deconvoluted H2 temperature-programmed surface reduction profiles of (A) 5Ni3SiAl catalyst, and (B) 5Ni3WAl catalyst.

Author Contributions: A.S.A.-F., A.A.I., and S.O.K. synthesized the catalysts, carried out all the experiments and characterization tests, and wrote the manuscript. R.P., F.A.-M., and R.K. prepared the catalyst and contributed to proofreading of the manuscript. F.A.-M., M.L.C., Y.A.A., and A.A.B., writing—review and editing; A.H.F. contributed to the analysis of the data and proofread the manuscript. All authors have read and agreed to the published version of the manuscript.

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