Effect of a titania covering on CNTS as support for the Ru catalysed selective $\tau$ CO methanation


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ABSTRACT

One of the major set-backs in the selective CO methanation process, as the final clean-up step in removing residual CO from reformate gas feed, is the reverse water gas shift (RWGS) reaction. This reaction is an undesired reaction because, it runs parallel with the selective CO methanation reaction. This increases the CO outlet concentration. The catalytic performance of ruthenium supported on carbon nanotubes (CNTs), nitrogen doped carbon nanotubes (NCNTs), titania coated carbon nanotubes (NCNT-TiO$_2$ and CNTs-TiO$_2$) and TiO$_2$ anatase (TiO$_2$-A) for selective CO methanation was investigated. The feed composition relevant to reformate gas was used but in the absence of steam. The experiments were conducted within a temperature range of 100 °C and 360 °C. It was observed that carbon dioxide methanation was suppressed until CO methanation attained a maximum conversion for all the catalysts studied. The Ru/NCNT showed higher activity than Ru/CNT at all temperatures examined due to the nitrogen incorporation in the carbon domains. Both Ru/CNT and Ru/NCNT however promoted the RWGS reaction at temperatures above 250 °C. The Ru/CNT-TiO$_2$ catalyst recorded the highest activity for both the CO and selective CO methanation followed by Ru/TiO$_2$-A. The presence of titania on the carbon nanotubes significantly retarded the RWGS reaction from about ~120% CO conversion to about 80% CO conversion, while selectivity towards methane increased in all catalysts with increasing temperature.

1. Introduction

Numerous efforts to fine tune the proton exchange membrane fuel cell (PEMFC) technology have been reasonably successful, especially as relates to the transportation industry which uses electrical power sources. This would apply to both stationary fuel cell applications and portable fuel cells. One of the major setbacks hindering possible com-mercialisation of a low temperature PEMFC is the generation of a “nearly-CO-free” gas feed but the current processes use may leave residual quantities of CO in the H$_2$ rich feed [1]. Hydrogen rich fuels for PEMFCs are obtained by reforming hydrocarbons, e.g. liquefied petro-leum and natural gas [1–3]. Trace amounts of CO impurities can de-activate the anode electrode of a PEMFC. For example, the anode electrocatalyst composed of Pt-Ru only tolerates a <50 ppm CO con-centration [4–6], while a Pt anode electrocatalyst can only tolerate concentrations of <10 ppm [7–9].

Usually, the water gas shift (WGSR) reaction is employed in initial process for clean-up when the CO concentration is in the range of I–8%.

Thus secondary processes are needed to reduce the CO level to ppm levels after the initial process.

There are several techniques that have been used to reduce CO in a reformate gas feed to lower concentrations, for use in a PEMFC [10]. These include using pressure swing adsorption (PSA), a method that makes use of special adsorptive materials (e.g. zeolites), which are employed as molecular sieves. The preferential CO oxidation (PROX) reaction Eq. (3) has also been widely studied over the years but today much focus has shifted from PROX to selective CO methanation called SMET (Eq. (1)) as a technique for removal of the final traces of CO in reformate gas feed. This is because SMET can avoid the limitations experienced by PROX such as addition of air from an external source. The CO and CO$_2$ methanation reactions (as indicated in Eqs. (1) and (2)) are less exothermic than the CO and H$_2$ oxidation reactions (Eqs. (3) and (4)), which makes it inherently easier to control SMET than PROX.

\[
\text{CO} + 3\text{H}_2 \leftrightarrow \text{CH}_4 + \text{H}_2\text{O} \quad \Delta H^\circ = -206 \text{ kJ/mol} \quad (1)
\]
CO + 4H₂ → CH₄ + 2H₂O \[\Delta H_{298} = -165 \text{ kJ/mol}\] (2)

CO + (1/2)O₂ → CO₂ \[\Delta H_{298} = -284 \text{ kJ/mol}\] (3)

H₂ + (1/2)O₂ → H₂O \[\Delta H_{298} = -244 \text{ kJ/mol}\] (4)

CO₂ + H₂ → CO + H₂O \[\Delta H_{298} = 41 \text{ kJ/mol}\] (5)

The SMET process is considered as a most promising process to remove CO in a reformate gas stream for PEMFC applications. The key to development in SMET is the design of an efficient catalyst, capable of removing CO, retarding CO₂ and inhibiting the RWGS reaction (Eq. (5)) at a sufficiently low temperature. Otherwise the methanation of carbon dioxide would occur which would then be a major hydrogen-consuming reaction.

A number of catalysts have been tested for this methanation of CO. They include Rh [11–14], Ni [12,15,16] and Ru [14,17,18]. In the selective CO methanation reaction Ru and Rh are still considered to yield high activities and unlike Pt and Pd do not promote the undesired reverse water gas shift reaction [19]. The nature of the support plays a role in the overall performance of a catalyst and this suggests a strategy to modify the behaviour of SMET catalysts [13,20,21].

Recently the use of carbon nanotubes (CNTs) as model catalyst supports for hydrogenation reactions such as the Fischer-Tropsch (FT) reaction has increased. CNTs have become attractive due to a number of advantages they possess such as facile modification of their surface properties, good hydrothermal stability and above all, their moderate interaction with the active metal [22]. The impressive mechanical nature of carbon supports and the absence of micro-porosity, which reduces intraparticle mass transfer in reaction media, as well as the easy accessibility of the active metal phases makes the CNT an excellent catalyst support.

Carbon supports do however have a very low density which is a problem in terms of using the CNTs in commercial reactors. Other types of supports, for example the metal oxides such as TiO₂, SiO₂ and Al₂O₃ can be used but they suffer from strong metal support interactions and even the formation of mixed metal compounds in some cases or spinel. Carbon does not experience spinel or possibly no strong metal support interactions.

Combining carbon and titania, could improve or merge the properties of both the carbon and oxide supports. The photocatalytic abilities of TiO₂ are known to be enhanced by doping or coating titania with carbon. The position that carbon occupies in a TiO₂ matrix will determine its effect on the carbon [23]. Since the density of carbon materials is enhanced by coating carbon with TiO₂ (or vice versa) [24] these methodologies suggest a process for increasing the catalyst/support density when C is used in the support.

In this paper we report on the enhanced catalytic activity and suppression of the RWGS reaction using Ru as the active metal phase supported on TiO₂ coated CNTs. A simple hydrothermal route was employed to coat the CNTs with TiO₂.

2. Experimental

2.1. The synthesis of CNTs

CNTs and nitrogen doped CNTs (N-CNTs) were synthesized by using the chemical vapour deposition (CVD) technique [25]. The CNTs and N-CNTs were grown using a 5%Fe-5%Co/CaCO₃ catalyst [25,26]. About 1 g catalyst was placed in a quartz boat and placed in the middle of a quartz tube in a furnace. The furnace was heated to 700 °C at a rate of 10 °C per minute under a nitrogen flow rate of 240 ml/min. Once the reaction temperature was attained, acetylene gas was introduced into the reaction chamber at a flow rate of 90 ml/min. The reaction was then allowed to continue for 1 h. After the reaction, the acetylene gas flow was stopped and the reactor was allowed to cool to room temperature under a flow of nitrogen gas. For the N-CNT synthesis, acetylene was bubbled through acetonitrile (acetonitrile the source of nitrogen) [26], and the CH₃CN/C₂H₄ introduced into the CVD reactor.

The CNTs were washed with 55% HNO₃ to remove both residues of the catalyst and to functionalise the CNTs at the same time. To do this, a concentrated solution of 55% HNO₃ was added to the as-synthesised CNTs or the N-CNTs in a round bottom flask. The mixture was then refluxed at 110 °C for a period of 12 h. The residue was washed with a large amount of deionised water till the solution had a neutral pH. The CNTs were collected by filtration and then dried overnight in an oven at 100 °C.

2.2. Synthesis of CNT-TiO₂ composites and TiO₂ anatase

CNT-TiO₂ was synthesized by modifying the procedure used by Yan et al. [27]. In a typical synthesis, 1.0 g of CNTs were dispersed in 30 ml of butanol and sonicated for 30 min. Titanium butoxide was dissolved separately in 10 ml of butanol, sonicated for 20 min and then added dropwise to the functionalised CNTs while stirring at a temperature of 30 °C. Diethyamine (0.6 ml) was added dropwise to the above mixture which was stirred for 10 min at room temperature to catalyst Ti-O-Ti formation reactions. The mixture was then transferred into a Teflonlined autoclave and the reaction was carried out at 150 °C for 14 h. The product was centrifuged, washed several times with ethanol followed by water and then calcined at 250 °C for 4 h under nitrogen. The same procedure was repeated without any CNTs to synthesize titania with an anatase polymorph. This support was denoted as TiO₂-A.

2.3. Preparation of the catalyst by wet impregnation

The catalysts were prepared by using RuCl₃ as metal precursor. Approximately 1.74 mL of RuCl₃ solution was dissolved in 10 mL of deionised water and then sonicated for 15 min. The RuCl₃ solution was then added drop-wise into a beaker containing 1 g of the support (CNT, N-CNT, TiO₂ or CNT-TiO₂). The thick paste formed was dried in an oven overnight and then calcined at 250 °C for 3 h to remove unwanted residues.

2.4. Catalyst activity studies

(FBTR) of height 250 mm and an internal diameter of 16 mm. The reaction was carried out at atmospheric pressure. The temperature of the reaction was measured in the catalyst bed using a K-type thermocouple. For each run, 1 g of the catalyst was weighed and placed in a reactor. Prior to the reactions, the catalyst was reduced at 250 °C for 4 h at 1 bar pressure under 99.995% H₂ gas. The various reactions (CO methanation, CO₂ methanation and selective CO methanation in the presence of CO₂) were investigated in the temperature range 160–360 °C using a gas feed with composition of 1% CO, 20% CO₂, 10% N₂ and H₂ balance. A flow rate of 90 ml/min was used to feed the gas feed to the reactor. Online GCs were connected to the FBTR to monitor the reaction. Both a flame ionization detector (FID) and a thermal conductivity detector (TCD) were used.

2.5. Characterization

The surface areas of all the catalysts and supports were analysed by BET using N₂ physisorption (Micromeritics ASAP-2000 Tri-star analyser). Before the surface area analysis, 0.2 g of the sample was degassed for 6 h at 150 °C under a N₂ flow using a Micromeritics flow Prep 060. The weight losses of the carbon samples were monitored on a Perkin-Elmer STA6000 TG/DTA Thermogravimetric analyser. The analyses were performed in air at a heating rate of 10 °C/min. Powder X-ray diffraction (XRD) studies were done on a Siemens D2 diffractometer using Co Kβ radiation equipped with a Ni filter. Steps of 0.02° with a scan range of 10–90° were used. The morphologies of the
samples were identified using SEM (FEI Nova Nanolab 600) and TEM (Tecnai T12) images. TEM samples were prepared by dissolving the catalyst in ethanol followed by sonication for 2 min. A drop of the ethanol suspension was transferred to a copper grid and dried. The disorder in the carbon based supports was measured with a micro-Raman attachment of a Jobin-Yvon T64000 Raman spectrometer. The excitation wavelength used was 514.5 nm from an Ar ion laser and the spot size on the sample was ca. 1.5 μm in diameter.

3. Results and discussion

3.1. Support characterization

3.1.1. TEM and SEM analysis

TEM analysis was used to elucidate the structural and morphological features of the different supports. From the images (both SEM and TEM) shown in Fig. 1, the expected tubular structure for the CNTs and N-CNTs was observed. Incorporation of nitrogen in the CNT’s is usually characterized by the presence of bamboo like structures, which are typically seen as a series of compartments inside the tube. The TEM insert of Fig. 1a shows this structure. Both CNTs and N-CNTs were identified to be multi-walled. Their diameters were found to be in the range of 40–70 nm and 30–80 nm for CNTs and N-CNTs respectively. Analysis carried out to quantify the nitrogen content present in the N-CNTs was found to be 3.7% by mass using CHNS analysis. This was confirmed with the presence of the bamboo structure seen in the TEM image. Fig. 1c shows the SEM and TEM images of the CNTs coated with TiO₂. A closer view of the TEM image (insert in Fig. 1c; higher resolution image) showed that the coverage of the TiO₂ was not complete on the CNTs. There were some regions with little coating and other regions where the titania nanoparticles formed aggregates along the length of a carbon nanotube. The anatase coating on N-CNT (NCNT-TiO₂) produced smaller TiO₂ particles and these covered the N-CNTs (Fig. S1) more than observed for the CNT-TiO₂ sample. This could be due to the presence of surface nitrogen.

3.1.2. Raman spectral analysis

Raman spectroscopy was used to determine the crystallinity and information on the defects in the synthesised CNTs and N-CNTs. Two main characteristic peaks are associated with carbonaceous materials. These peaks are the D-band (approximately 1350–1360 cm⁻¹) which identifies with the defects, while the G-band (1590–1600 cm⁻¹) corresponds to the C=O bond stretching mode in the graphite plane. The area ratio of the G to D bands (I₆/I₉) gives an idea of the extent of defects in the carbon material. A high ratio suggests high levels of defects [28]. The Raman spectrum (Fig. 2) showed that nitrogen
incorporation into the CNTs increased the defect concentration as seen by the increased \( I_D/I_G \) ratio (Table 1). The broadening of the D and G bands coupled with a slight shift to higher wave numbers again confirmed an increased defect concentration in the N-CNTs [29]. Usually, defects result in the creation of anchorage sites for nano metal particles which serve as the active species in catalysis.

3.1.3. PXRD analysis

The crystalline phases in the support materials were identified by PXRD. As shown in Fig. 3, graphite carbon peaks at \( 2\theta \) angles of ca. 31\(^\circ\) were present in all the carbon containing supports (N-CNTs not shown). The diffraction patterns for the hydrothermally produced TiO\(_2\) and the TiO\(_2\) coated CNTs (N-CNT-TiO\(_2\) similar to the CNT-TiO\(_2\)) corresponded to the anatase polymorph of TiO\(_2\). The peaks obtained were similarly sharper as the pure anatase which indicates that the titania and carbon nanotubes were crystalline.

3.1.4. Thermogravimetric analysis (TGA)

The TGA plots in Fig. 4 allows for the determination of the per-centage purity and the thermal stability of the CNTs, N-CNTs and the coated CNTs. The TGA plots of the unpurified samples (as-synthesised CNTs and N-CNTs) are shown in Fig. S2. It is noted that the HNO\(_3\) acid treatment was effective in removing the bulk of the residue of the Fe and Co catalyst. The functionalization process significantly removed impurities (residue from catalyst) from about 30\% < 1.5\% and from 45\% to < 2\% for CNTs and N-CNTs respectively (Fig. S2). There were no weight loss registered below 400 \(^\circ\)C which suggested that there was no amorphous carbon present [30,31]. The N-CNTs decomposed below 500 \(^\circ\)C i.e. about 100 \(^\circ\)C less than the CNTs. This is also an indication that presence of defects present in the carbon framework reducing the thermal stability of the carbon. The TGA results also showed a small weight loss in the low temperature region of Fig. 4; this is due to the decomposition of butanol, diethylamine or other residual organic materials that were adsorbed onto/into the CNTs. The maximum decomposition temperature for the CNT-TiO\(_2\) was around 63 \(^\circ\)C, relatively higher than that for the CNTs. The TGA data indicated that about 25\% of the CNT-TiO\(_2\) was composed of TiO\(_2\) (Fig. S2). The improvement in the thermal stability was as a
Fig. 5. Representative images showing RuO2 particles (and their size distribution) dispersed on the different supports: (a) TEM image of 5%Ru/CNT, (b) particle size distribution diagram for 5%Ru/CNT (c) TEM image of 5%Ru/N-CNT, (d) particle size distribution diagram for 5%Ru/N-CNT (e) TEM image of 5%Ru/CNT-TiO2 and (f) particle size distribution diagram for 5%Ru/CNT-TiO2 (g) TEM image of 5%Ru/TiO2 (h) particle size distribution diagram for 5%Ru/TiO2.
result of the TiO$_2$ covering which shielded the carbon from oxidation.

3.1.5. N$_2$ adsorption studies
The surface area and porous structure of materials were determined using the nitrogen adsorption technique [32]. Introduction of nitrogen into the carbon matrix increased the surface area and increased the pore volume of the supports. This was attributed to both the functionalization process and the doping [25,29]. The hydrothermally produced anatase TiO$_2$ had the lowest surface area (52 m$^2$/g) which correlated with literature data. Combining CNT with TiO$_2$ to form a CNT-TiO$_2$ composite significantly improved the surface area (from 75 to 52 to 119 m$^2$/g).

3.2. Catalyst characterization
3.2.1. TEM analysis on the catalyst
TEM images of the various catalysts prepared and studied are presented in Fig. 5. The darker spots observed in all the TEM images are due to RuO$_2$ nano particles. The average RuO$_2$ particle size distribution calculated from the TEM images are shown in the size distribution plots. The RuO$_2$ particles on the CNT were observed to have an average size of 6.4 nm and to have a wider size distribution range compared to the other catalysts (Fig. 5b). The RuO$_2$ particle sizes on the N-CNTs were smaller with an average size of 3.8 nm (Table 1) with a relatively narrower size distribution (Fig. 5d). The presence of the nitrogen enhanced anchoring of the RuO$_2$ particles [33]. The Ru supported CNT-TiO$_2$ particles were smaller (2.3 nm) than those found on the pure anatase TiO$_2$ support which also had fairly small RuO$_2$ particles (3.6 nm). This was probably due to the synergetic effect offered by the CNT-TiO$_2$ composite and the larger surface area. The Ru particles were distributed on both TiO$_2$ and the CNT surfaces as show in the TEM images.

3.2.2. BET analysis on the catalyst
The surface areas of the catalysts decreased compared to the supports (Table 1). This was due to the occupation of the pores of the supports by Ru, particles. The pore volume increased slightly due to occupation of smaller pores after loading of the Ru except for the 5% Ru/CNT-TiO$_2$ sample [34].

3.2.3. PXRD analysis on the catalyst
The phases present in the Ru catalysts on the support were investigated by XRD and are presented in Fig. 6. The peaks observed at 32.5 and 40.8° were indexed to RuO$_2$ (110) and (101) planes respectively. No peaks for Ru were observed on 5%Ru/CNT-TiO$_2$ and 5% Ru/N-CNT-TiO$_2$ (Fig. S2). This was due to the small undetected RuO$_2$ particles which were well dispersed as observed on the TEM analysis and also the difficulty of observing Ru on TiO$_2$. The RuO$_2$ peaks observed on the CNT sample indicated larger particle sizes (7.9 nm from XRD, Table 1). The RuO$_2$ peaks for 5%Ru/N-CNTs and 5%Ru/TiO$_2$-A were almost similar (broad and less intense) which confirmed the relatively smaller particles observed by TEM analysis.

3.2.4. TPR studies
The reduction profiles of the different catalysts are shown in the H$_2$-TPR plots in Fig. 7. Reduction of RuO$_2$ to Ru for all the catalyst occurred below 200 °C. The reduction profile for the carbon and titania composites revealed maximum broad reduction peaks at elevated temperatures 163 °C and 174 °C Ru/CNT-TiO$_2$ and Ru/N-CNT-TiO$_2$ respectively. This suggested Ru particles were dispersed on both the CNTs and TiO$_2$ components of the composites as observed in the TEM images. The broadness was attributed to RuO$_2$ reduction on the different support (carbon and titania supports). The RuO$_2$ particles were believed to have interacted slightly more strongly with the functional groups present on the carbon supports. This was revealed by the reduction RuO$_2$ to Ru occurring at relatively higher temperature compared to TiO$_2$-A supported catalyst. Reduction on the N-CNTs (RuO$_2$ to Ru) occurred at slightly higher temperature than the RuO$_2$ on CNTs. This could be due to the extra lone pair of electron made available by the surface nitrogen (pyridinic nitrogen) which formed a stronger bond with the RuO$_2$ i.e. the nitrogen on the surface provided strong anchorage to Ru particles. Surprisingly, the RuO$_2$ supported on the TiO$_2$-A reduced at an unexpectedly lower temperature (113 °C) than both CNT and N-CNTs supports. To confirm this reduction observed on the anatase TiO$_2$, a series of anatase supports were synthesized using a sol gel method and a resin gel method and used as a Ru catalyst support. Their reduction profiles were compared to Degussa P25 TiO$_2$ Ru supported catalyst as shown in the Fig. S3. This may suggest that the CNTs and N-CNTs binds the Ru particles firmly through the functional groups or alternatively,
The density (catalyst bed height) of the Ru/CNT and Ru/N-CNT contributed in a longer interaction of the hydrogen (reducing gas) which resulted in the shift in RuO₂ reduction peaks. This possibly affected the composite graphs. Further studies should be conducted (as stated in the recommendations) on TPR to clarify this.

The methanation or gasification of the carbon support was the reason assigned to the peaks occurring at higher temperatures. This peak was absent on the Ru/TiO₂-A profile.

4. CO hydrogenation

The activity and selectivity performance of the different supports (CNTs, NCNT, CNT-TiO₂ and TiO₂-A) for CO hydrogenation over Ru catalyst with performed with the same Ru loading (5 wt.%). The reactions were performed using CO or CO/CO₂ to study methanation and selective CO methanation conversion in the presence of CO₂. Fig. 8 summarises the conversion data when only CO was used. Above ca.140 °C, all the catalysts investigated showed a steady increase in CO conversion (Fig. 8) and all attained 100% conversion at temperatures beyond 280 °C. The N-CNT Ru supported catalyst showed a higher activity than observed when CNT was used as a support. This was attributed to the presence of pyridinic nitrogen with an extra lone pair of electron which contributed extra electrons making the surface of the tubes more electonegative. This resulted in increased nucleation sites which served as anchorage sites thereby enhancing dispersion of the active phase while restricting the Ru particle sizes [33]. Again, the extra electrons due to the incorporation of nitrogen atom increased the electron density around the Ru and the electrons were then back-donated from Ru d-orbitals to a π* antibonding molecular orbital of CO. This strengthened the Ru-C bond (back bonding) weakening the CeO bond, promoting the dissociation of CO and increasing the methanation of CO with Ru/N-CNT over Ru/CNT [35]. It should be noted that, titania coating on the carbon nanotube as support exhibited the highest activity, performing better than Ru/TiO₂-A. This was due to the higher specific surface area of the coated support and the smaller par-ticle sizes of the Ru.

Anatase on its own has a high surface energy which tends to facilitate agglomeration of metal particles supported on it [36–38]. Defects on titania surfaces (Ti³⁺ are related to anionic oxygen) which are known to stabilize metallic species from agglomerating [37]. These Ti³⁺ surface defects are mostly associated with anatase-rutile transformation but Kongkiat et al. reported they could modify the surface structure (Ti³⁺ defects) of anatase by using varying oxygen content in their calcination process under temperatures below the anatase-rutile transformation [39]. These defects could have been inroduced on the surface of our coated CNTs due to the presence of oxygen (made available through functionalizing the CNTs) present during the synthesis or calcination process. Alternatively, the small Ru particles found on the coated support may have resulted from synergetic effects between the CNT-TiO₂ which is believed to have improved the specific surface area. The NCNT-TiO₂ performed slightly better than the CNT-TiO₂ composite. It can be concluded that the activity of CO hydrogenation is dependent on the type of support used in our experiments.

In spite of the small surface area recorded for the TiO₂-A support, the Ru catalyst showed much higher activity than the Ru/N-CNT and Ru/CNT catalysts. Besides the presence of anionic vacancies that are present at the edges of the oxide-metal boundaries on the TiO₂, a complex is formed between the oxygen groups on the TiO₂ and the adsorbed CO [40]. Sachler and Ichikawa and co-authors suggested that the Lewis acid/base bonding interaction that occurs between the adsorbed CO and oxygen end and a promoter influence at the oxide/active metal interface [41,42]. These effects are said to enhance the cleavage of the C=O bond which results in higher activity for TiO₂.

4.1. CO₂ Hydrogenation

The hydrogenation of CO₂ was initiated above ca. 150 °C as shown in Fig. 9. Conversion remained below 20% until 325 °C for all catalyst except Ru/CNT. It should be noted that no significant conversion occurred within the temperature window in which our catalysts were tested. This means significant methanation of CO₂ (above 50% conversion) may only occur at higher temperatures. The supports 5%Ru/CNT and 5%Ru/N-CNT had higher activity than the other catalysts, and suggested the supports had effects on the CO₂ conversion. Several studies acknowledge that CO₂ mostly adsorbs on the supports while hydrogen is adsorbed on the metal to produce methane. The presence of OH groups could have contributed to the higher activity of the functionalized carbon supports [43]. The 5%Ru/CNT was slightly more active than 5%Ru/N-CNT, which could mean particle size played a role in the reaction.

5. Selective CO methanation

Fig. 10 summarises the performances of all the catalyst investigated for the selective methanation of CO in the presence of CO₂. The Figure
The effect of the support on the catalytic performance of Ru in the selective methanation of CO in the presence of CO$_2$ (XCOXCO$_2$) as a function of reaction temperature using 5% wt.% Ru. Solid symbols: CO conversion; open symbols: CO$_2$ conversion. Experimental conditions: mass of catalyst: 1 g; reactant composition: 1% CO, 20% CO$_2$, 10% N$_2$ (balance H$_2$); total flow rate: 90 cm$^3$/min.

showed a similar activity sequence for all the catalysts performance as seen when only CO was hydrogenated. The same explanations can be given to the activity trend noted for the selective CO methanation reaction. CO conversion was initiated at about ca. 140 °C for all catalysts. The conversion of CO increased with increasing temperature for all the catalyst; at 280 °C, both Ru/CNT and Ru/N-CNT attained maximum conversions of 64% and 78% respectively. Increasing the temperature further above (>280 °C) indicated a drop in CO conversion revealing “negative” percentage values for CNT and N-CNT at about 320 °C. This is due to the CO hydrogenation reaction (Eq. (1)) running parallel to the RWGS reaction (Eq. (5)) under the experimental conditions. The “negative” CO conversion showed that the effluent CO concentration drastically increased due a slower rate of CO hydrogenation compared to the rate at which CO was produced via the RWGS reaction. This behaviour implies that both CNT and N-CNT promoted the RWGS reaction. Although, a drop in CO conversion was observed for NCNT-TiO$_2$ and CNT-TiO$_2$ at ca. the same temperature, the effect is small. Clearly, the presence of anatase TiO$_2$ suppressed the RWGS reaction relative to the Ru/N-CNTs and Ru/CNTs. This result is in agreement with the study by Jiménez et al. who used different carbon nanofibers; they observed that the carbon material promoted a high RWGS.

Hydrogenation of CO$_2$ in the presence CO was minimal at temperatures before CO attained maximum conversion for all the catalysts investigated. This confirms that CO interacted more strongly on the surface of the catalyst than CO$_2$, in agreement with reports by several authors [44–46]. It is widely accepted that CO$_2$ hydrogenation proceeds by the adsorption of CO$_2$ on the support with H$_2$ on the metal [47–50]. The work of Shohei et al. suggested that the presence of hydroxyl groups on TiO$_2$ at elevated temperatures facilitated the hydrogenation of CO$_2$ [43]. The CNT and N-CNT supports are known to have hydroxyl groups on the surface from the acid treatment. This could possibly explain the relatively higher conversion of CO$_2$ on Ru/CNT.
6. Methane selectivity

Several studies that have focused on selective CO methanation have shown that besides methane other higher hydrocarbons are formed, especially at low temperatures when CO conversions are low. In our studies, plots revealed that besides methane, the hydrogenation of CO/CO₂ resulted not only in methane but other higher hydrocarbons (Fig. 11). The catalyst supported on pure anatase (5%Ru/TiO₂-A) showed the highest selectivity towards methane. Fig. 10d shows that other than methane only C₂ and C₃ molecules were formed at low temperatures. Above 260 °C only methane was observed. The CNT-TiO₂ composite showed 100% methane selectivity at temperatures above 300 °C. Both 5%Ru/N-CNT and 5%Ru/CNT did not attain total methane selectivity in the temperature range of this study. The data show that larger Ru particle sizes support higher carbon growth. Similar results were observed for CNT-TiO₂ and NCNT-TiO₂ (not shown).

7. Conclusions

The coating of CNTs and N-CNTs using titania was successfully done. However, the coating did not result in total coverage. The coating of CNTs and N-CNTs with titania produced a catalyst support with better thermal stability and a larger surface area compared to the functionalised CNTs. The catalytic activity as well as selectivity in this study depended on the support used and experimental conditions used (e.g. only CO methanation, the mixture of CO and CO₂ reaction temperature etc.). Total CO conversion was attained for all catalysts at a temperature ca. 280 °C when only CO was methanated. In the presence of CO₂ a slight variation of conversions was recorded for all catalysts. The titania coated CNT showed significantly higher activity compared to the uncoated CNTs and N-CNTs. The presence of TiO₂ on the CNTs suppressed the RWGS reaction and the CO methanation rate. Higher temperatures increased methane selectivity at the expense of higher hydrocarbon formation.

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Appendix A. Supplementary data

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