Dry reforming of glycerol over Rh-based ceria and zirconia catalysts: New insights on catalyst activity and stability

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ABSTRACT

Effects of reaction temperature and feed composition on reactant conversion, product distribution and catalytic stability were investigated on syngas production by reforming of glycerol, a renewable waste, with CO\textsubscript{2} on Rh/ZrO\textsubscript{2} and Rh/CeO\textsubscript{2} catalysts. For the first time in the literature, fresh and spent catalysts were characterized by \textit{in-situ} FTIR, Raman spectroscopy, transmission electron microscopy and energy dispersive X-ray analysis techniques in order to unravel novel insights regarding the molecular-level origins of catalytic deactivation and aging under the conditions of glycerol dry reforming. Both catalysts revealed increased glycerol conversions with increasing temperature, where the magnitude of response became particularly notable above 650 and 700 °C on Rh/ZrO\textsubscript{2} and Rh/CeO\textsubscript{2}, respectively. In accordance with thermodynamic predictions, CO\textsubscript{2} transformation occurred only above 700 °C. Syngas was obtained at H\textsubscript{2}/CO \textasciitilde 0.8, very close to the ideal composition for Fischer-Tropsch synthesis, and carbon formation was minimized with increasing temperature. Glycerol conversion decreased monotonically, whereas, after an initial increase, CO\textsubscript{2} conversion remained constant upon increasing CO\textsubscript{2}/glycerol ratio (CO\textsubscript{2}/G) from 1 to 4. In alignment with the slightly higher specific surface area of and smaller average Rh-particle size on ZrO\textsubscript{2}, Rh/ZrO\textsubscript{2} exhibited higher conversions and syngas yields than that of Rh/CeO\textsubscript{2}. Current characterization studies indicated that Rh/CeO\textsubscript{2} revealed strong metal-support interaction, through which CeO\textsubscript{2} seemed to encapsulate Rh nanoparticles and partially suppressed the catalytic activity of Rh sites. However, such interactions also seemed to improve the stability of Rh/CeO\textsubscript{2}, rendering its activity loss to stay below that of Rh/ZrO\textsubscript{2} after 72 h time-on-stream testing at 750 °C and for CO\textsubscript{2}/G = 4. Enhanced stability in the presence of CeO\textsubscript{2} was associated with the inhibition of coking of the catalyst surface by the mobile oxygen species and creation of oxygen vacancies on ceria domains. Deactivation of Rh/ZrO\textsubscript{2} was attributed to the sintering of Rh nanoparticles and carbon formation.

1. Introduction

A majority of the world’s existing energy demand is met by fossil fuels such as crude oil, coal and natural gas, all of which accounts for more than 80% of total energy consumption \cite{1}. However, increasing costs of exploration and production of fossil fuels together with the environmental and societal impacts of global warming caused by accumulation of CO\textsubscript{2}, accelerated efforts towards research, development and commercial use of renewable fuels and energy conversion technologies. Among a number of renewable fuels, biodiesel is receiving increased attention as it can be blended with the crude-oil based diesel without losing its compatibility with the existing diesel engines \cite{2}. Since 2005, biodiesel market grew by \textasciitilde 23% per year, which corresponds to a seven-fold market expansion in the last decade \cite{3}.

Biodiesel production is carried out by transesterification of animal-based or vegetable oils with methanol or ethanol. In this process, one mole of glycerol is produced as a side product for every three moles of biodiesel \cite{4,5}. This stoichiometry, however, leads to a significant surplus of glycerol. It is predicted that cumulative glycerol supply will be \textasciitilde 3 \times 10\textsuperscript{6} ton by 2020, whereas the demand will remain below \textasciitilde 5 \times 10\textsuperscript{5} ton/year \cite{4}. Along these lines, production costs of biodiesel can be lowered by catalytic valorization of excess glycerol into value-added products such as syngas (i.e. synthesis gas), which is the raw material of key commodities such as synthetic fuels, methanol and...
dimethyl ether [6].

Steam reforming is the main route to transform glycerol into a hydrogen-rich gas mixture. As summarized in various comprehensive reviews [4,6-8], glycerol steam reforming (GSR) has been investigated extensively in various aspects ranging from catalyst development at fundamental levels to the testing of different reactor configurations. Steam reforming generates syngas with low CO content (i.e. H2/CO > 2), rendering it disadvantageous in the production of long chain hydrocarbons via Fischer-Tropsch (FT) synthesis [9-11]. Reforming of glycerol with CO2 on the other hand, is capable of not only delivering syngas with molar H2/CO ratios close to 1, but also satisfying the feed conditions for FT synthesis to obtain long chain hydrocarbons [9,12]. Moreover, dry reforming offers the advantage of obtaining syngas from molecules which can cause serious economic and environmental penalties. These benefits make the dry reforming route a promising option for glycerol valorization.

Glycerol dry reforming (GDR) is an endothermic process where one mole of glycerol reacts with one mole of CO2 to produce H2, CO and H2O through the following overall reaction:

\[
\text{C}_3\text{H}_6\text{O}_3 + \text{CO}_2 \rightarrow 4\text{CO} + 3\text{H}_2 + \text{H}_2\text{O} \quad \Delta H^\circ = 292 \text{ kJ/mol}
\] (1)

In dry reforming conditions that involve temperatures in excess of 500 °C and presence of CO2 in the feed, reverse water gas shift (RWGS - Reaction 2) affects the product distribution [13]. Therefore, Reaction 1 can be envisioned as the combination of RWGS and glycerol decomposition (Reaction 3):

\[
\text{CO}_2 + \text{H}_2 \rightarrow \text{CO} + \text{H}_2\text{O} \quad \Delta H^\circ = 41 \text{ kJ/mol}
\] (2)

\[
\text{C}_3\text{H}_6\text{O}_3 \rightarrow 3\text{CO} + 4\text{H}_2 \quad \Delta H^\circ = 251 \text{ kJ/mol}
\] (3)

In addition to Reaction 3, decomposition of glycerol involves a series of dehydration and dehydrogenation reactions which lead to the production of various species such as methane, ethane, ethylene, acetaldehyde, acrolein, acetone, methanol, ethanol and acetic acid [14]. These species can eventually be converted into coke via homogeneous/heterogeneous thermal cracking reactions. Other possible side reactions are steam and dry reforming of methane (Reactions 4 and 5) and of higher hydrocarbons, and coke gasification (Reactions 6-8):

\[
\text{CH}_4 + \text{H}_2 \rightarrow \text{CO} + 3\text{H}_2 \quad \Delta H^\circ = 206 \text{ kJ/mol}
\] (4)

\[
\text{CH}_4 + \text{CO}_2 \rightarrow 2\text{CO} + 2\text{H}_2 \quad \Delta H^\circ = 247 \text{ kJ/mol}
\] (5)

\[
\text{C(s)} + \text{H}_2 \rightarrow \text{CO} + \text{H}_2 \quad \Delta H^\circ = 131 \text{ kJ/mol}
\] (6)

\[
\text{C(s)} + \text{CO}_2 \rightarrow 2\text{CO} \quad \Delta H^\circ = 172 \text{ kJ/mol}
\] (7)

Endothermic nature of GDR requires temperatures above ~500 °C, where the reaction is thermodynamically favored [12,15,16]. Wang et al. [15] reported that 727 °C and molar inlet CO2/G of 1 were optimum thermodynamic conditions for maximizing H2 yield, and showed that molar H2/CO ratios produced by GDR changed between 1 and 2.15 by varying the temperature between 500–700 °C and CO2/G ratios within 1–5. They also mentioned that coke formation decreased with increasing temperature. Thermodynamically, coke formation became insignificant above 677 °C at CO2/G = 1 [15].

GDR has become the focus of experimental studies only recently and the available information in the literature regarding its catalysis is rather scarce. Siew et al. [17-20] studied GDR over La-promoted Ni/Al2O3 catalysts at temperatures between 650–850 °C and CO2/G = 0–5. The authors concluded that La promotion (i) provided better metal dispersion (i.e. a finer crystallite size and higher BET specific surface area), (ii) significantly reduced carbon deposition, and (iii) reduced deactivation rate as quantified by the 72 h time-on-stream stability tests carried out at 750 °C to give an average glycerol conversion of 90%. They also reported that presence of CO2 was essential in reducing carbon deposition through the gasification reactions. Lee et al. [21,22] investigated GDR over Ni-based catalysts supported on cement clinker (CC), a material composed mainly of CaO and MgO, with the intention of utilizing CO2 emitted during cement production. They concluded that use of CC facilitated the suppression of carbon formation. Furthermore, increasing Ni loading from 5 wt.% to 20 wt.% improved BET specific surface area from 0.6 to 18 m2/g. The authors also reported H2/CO ratios below 2 and glycerol conversions up to ~80% upon reaction at 750 °C and CO2/G of 1.67, which were claimed to be optimum for the 20 wt.% Ni loaded catalyst. Arif et al. [23] compared the activities of CaO and ZrO2 supported Ni catalysts with different metal loadings in GDR. It was reported that at 700 °C and for CO2/G ratio of 1, Ni/CaO catalyst gave a H2 yield and a glycerol conversion higher than those obtained over Ni/ZrO2. Superiority of CaO supported samples over ZrO2 supported ones was attributed to the higher metal dispersion and smaller NiO crystallite size over CaO as revealed by XRD analysis. Harun et al. [24] studied 0–5% Ag-promoted 15% Ni/SiO2 catalysts for GDR. It was revealed by XRD analysis that addition of Ag did not change the metal crystallite size significantly, but improved H2 yield and glycerol conversion. Carbonaceous deposits were found to exist on the catalysts upon SEM analysis of the spent samples.

As summarized above, a comprehensive molecular-level fundamental understanding of GDR catalysis is clearly missing. Moreover, detailed information regarding the effect of reaction parameters on CO2 conversion, yield and syngas composition is not available. Thus, in the current work, we follow a systematic experimental approach in order to address some of these issues and combine catalytic activity/selectivity studies with detailed molecular-level in-situ/ex-situ spectroscopic/imaging investigations in an attempt to obtain new structure-functionality relationships about GDR. We focus on Rh-based catalysts supported on ZrO2 and CeO2. Although Rh-based catalysts were reported to exhibit superior activity and stability features in dry reforming of various hydrocarbons [25-29], no such catalysts exist in the literature for GDR process. As for the supports, both ZrO2 and CeO2 are known as promising materials for dry reforming reactions due to their advantageous oxygen transfer properties. Oxygen vacancies on ZrO2 are reported to help dissociation of CO2 into CO and O, which in turn, facilitates the oxidation of the surface carbon species [30-32]. CeO2 is known to have a high oxygen storage capacity, which can create an oxygen reservoir enabling gasification of coke [33,34]. In the light of these points, Rh/ZrO2 and Rh/CeO2 can be envisioned as promising novel catalysts that can be investigated in GDR process. Along these lines, in the current study, we report Rh-based GDR catalysts which outperform the existing Ni-based ones [17–24] in terms of activity and stability. Furthermore, through in-situ/ex-situ spectroscopic/imaging experiments, for the first time in the literature, we provide valuable insights regarding the origins of catalytic activity, stability, aging and sintering in GDR reaction.

2. Experimental

2.1. Catalyst synthesis and pretreatment

Catalysts, 1 wt.% Rh/CeO2 (RhCe) and 1 wt.% Rh/ZrO2 (RhZr), were prepared by conventional incipient wetness impregnation method. The first step in synthesizing RhCe was the preparation of the support. For this purpose, Ce(NO3)3·6H2O (purity: 99.99%, Sigma-Aldrich) was first calcined in air at 600 °C for 4 h for achieving thermal decomposition to CeO2 [35]. The resulting material was calcined at 800 °C in a muffle furnace under air atmosphere for 4 h in order to enhance thermal stability (i.e. for preventing its sintering during reaction conditions). A necessary amount (7 × 10−2 mol/g catalyst) of liquid Rh-precursor (Rh(NO3)3, purity: 10% (w/w) Rh in > 5 wt.% HNO3 solution, Sigma-Aldrich) was dissolved in deionized water and this aqueous precursor solution was impregnated onto the CeO2 support with a particle size range of 2.5–3.5 × 10−4 m by means of a peristaltic pump under vacuum. The resulting slurry was dried overnight in an oven at 158
110 °C and then calcined in a muffle furnace at 800 °C for 4 h. An identical protocol was used for the synthesis of RhZr. Prior to its use in impregnation, the zirconia support (ZrO₂, purity: 99%, Alfa Aesar) was brought to a particle size range of 2.5–3.5 × 10⁻⁴ m and calcined at 800 °C for 4 h. Before reaction tests, the catalysts were reduced in-situ at 800 °C for 2 h under 40 Nml/min H₂ (purity > 99.99%, Linde GmbH) flow measured by a Brooks 5850E Series Mass Flow Controller.

2.2. Catalyst characterization

2.2.1. N₂ physisorption

BET isotherms were obtained by using a Quantachrome Nova 2200e automated gas adsorption system with liquid nitrogen at a temperature of −196 °C. Specific surface areas of the pure support materials (i.e. ZrO₂ and CeO₂) calcined at 800 °C for 4 h were determined via multi-point BET analysis. Pore sizes and pore diameters were calculated using the BJH method.

2.2.2. Transmission electron microscopy (TEM) and energy dispersive X-Ray (EDX) analysis

TEM imaging and EDX analysis of the fresh (i.e. reduced) and spent catalysts (after 5 h reaction at 750 °C, CO₂/G = 4, residence time = 3.75 mg.min/Nml) were performed via a FEI, Tecnai G2 F30 microscope using an electron beam voltage of 300 kV. Before TEM-EDX analysis, each sample was dispersed in ethanol and sonicated for 5 min. Then, the sample suspension was transferred on a copper TEM grid by using a micropipette. The excess solution was removed and the copper grid was dried in the fume hood at room temperature overnight. While bright field imaging mode was used for the high resolution TEM (HR-TEM) measurements, high angle annular dark field scanning transmission electron microscopy (HAADF-STEM) was utilized for the EDX analysis. Average particle size values were determined using the ImageJ digital image processing software.

2.2.3. Ex-situ Raman spectroscopic analysis

Ex-situ Raman spectroscopic experiments were performed on fresh (i.e. reduced) and spent catalysts (after 5 h reaction at 750 °C, CO₂/G = 4, residence time = 3.75 mg.min/Nml) using a Renishaw inVia-Reflex confocal Raman microscope/spectrometer by utilizing a 532 nm laser with an adjustable power where the maximum output laser power at the end of the fiber was 50 mW.

2.2.4. in-situ FTIR spectroscopic analysis

in-situ FTIR measurements were carried out in a custom-designed batch-type spectroscopic reactor coupled to an FTIR spectrometer (Bruker Tensor 27) in transmission mode. FTIR spectra were recorded using a Hg-Cd-Te (MCT) detector, where each spectrum was acquired by averaging 32 scans with a spectral resolution of 4 cm⁻¹. Finely-ground powder catalyst samples of ~20 mg were pressed onto a lithographically etched W-grid sample holder (i.e. in the absence of KBr support/diluent material) materials. All of the FTIR spectra were acquired at 50 °C. Other details regarding experimental setup can be
found elsewhere [37]. In-situ CO adsorption experiments were carried out using FTIR technique by exposing the fresh (i.e., reduced) or spent (obtained after either 5 h or 72 h time on stream experiments carried out at 750 °C, CO2/G = 4 and residence time = 3.75 mg.min/Nml) catalyst surfaces to 10.0 Torr of CO(g) for 10 min at 50 °C. Next, spectroscopic chamber was evacuated to ~10−2 Torr at 50 °C, where FTIR spectra were acquired. Prior to each analysis, background spectrum was collected for each sample after reactor was evacuated to ~10−2 Torr at 50 °C. Otherwise mentioned, prior to in-situ FTIR analysis, no pretreatment protocols were applied to the fresh and 5 h spent catalysts.

On the other hand, since the surfaces of the 72 h-spent catalysts were significantly covered with carbonaceous species due to coke deposition during GDR reaction, 72 h-spent catalysts were treated with 10.0 Torr of H2 (purity > 99.999%, Linde GmbH) for 10 min at 300 °C prior to CO adsorption experiments via in-situ FTIR, in an attempt to remove some of the coke deposition and unravel Rh nanoparticles to make them available to CO adsorption. The background spectrum for 72 h-spent catalyst was collected after pre-treatment as mentioned above.

Nature and relative abundance of the carbonaceous species (which were generated during the GDR process) on the spent catalyst surfaces were also investigated using in-situ FTIR spectroscopy. For this purpose, FTIR spectra of the spent catalysts (without further treatment) were used as the background spectra and the sample in-situ FTIR spectra of the spent catalysts were obtained after exposing them to 10.0 Torr of H2 for 10 min at 300 °C, followed by evacuation and cooling to 25 °C.

2.3. Catalytic performance experiments

The catalytic performance tests were carried out in a down flow, quartz tubular packed bed reactor positioned in a three-zone furnace (Protherm PZF 12/50/500) as presented in Fig. 1. The quartz tube was 8 × 10−3 m long with an internal diameter (ID) of 2 × 10−2 m and narrowed down to an ID of 1 × 10−2 m along the central 1.5 × 10−1 m length of the tube. The vertically positioned furnace had a heated length of 5 × 10−1 m which involved three equidistant zones with dedicated PID controllers capable of regulating the pertinent zone temperature to ±1.0 °C via K-type thermocouples positioned at the midpoints of the zones. The catalyst bed, composed of a physical mixture of 20 mg of catalyst (RhZr or RhCe) and 700 mg of diluent (α-Al2O3, Alfa-Aesar), both at the particle size range of 2.5–3.5 × 10−4 m, was positioned at the middle of the second zone via a quartz wool plug that was supported by a specially designed inset existing within the narrower section of the quartz tube (with ID of 1 × 10−2 m) to prevent any physical movement of the bed. The resulting configuration gave a packed bed height of ~1 × 10−2 m to obtain bed height-to-particle diameter and tube diameter-to-particle diameter ratios of ~35, which were acceptable for ignoring diffusive transport terms and assuming plug flow behavior, respectively [38,39]. In all conditions pressure drop remained negligible (<1%) as simulated in CHEMCAD 7.1.4 chemical process simulation software. Absence of pressure drop was also confirmed experimentally by the steady, pulse-free flow of reactor effluent that was periodically monitored via directing it to a soap-bubble meter.

Liquid glycerol (Sigma-Aldrich, purity: 99.5%) was precisely metered by a Shimadzu LC-20AD HPLC pump and transported via a 1/16-inch outer diameter (OD) stainless steel tubing to the inlet of the quartz tube, where it contacted with CO2 and N2, both supplied by Linde GmbH with purities above 99.995% and metered by Brooks 5850E Series Mass Flow Controllers (Fig. 1). The gas-liquid mixture was then introduced into the quartz tube through a 1/16-inch OD stainless steel tubing acting as an injector. The tubing extended until 1 × 10−1 m into the first zone of the furnace, where the temperature was kept above 400 °C. This scheme enabled complete evaporation and continuous feeding of glycerol and ensured that the desired CO2/G feed ratio was obtained. After evaporation and mixing within the injector, the reactive mixture reached the catalyst bed located at the middle of the second zone of the furnace, where the temperature was set to the specified reaction temperature. The third zone of the furnace was kept above 350 °C to prevent any condensation within the downstream of the catalyst bed. After having left the quartz tube, the effluent stream was passed through two consecutive cold traps to knock out any condensable components such as water, unreacted glycerol or any other possible liquid-phase products of glycerol decomposition (Fig. 1). The remaining gaseous mixture was transferred to two on-line gas chromatographs (GCs) for qualitative and quantitative analysis. The first GC, Shimadzu GC-2014, was equipped with a thermal conductivity detector (TCD, detector temperature and current of 150 °C and 50 μA), respectively and a 60–80 mesh Molecular Sieve 5 A packed column operated under 25 Nml/min Ar (purity > 99.99%, Linde GmbH) carrier gas flow at 50 °C to detect H2, N2, CH4 and CO. The second GC, Agilent 6890 N, involved an 80–100 mesh Porapak Q packed column and 20 Nml/min He (purity > 99.99%, Linde GmbH) carrier gas flow at 40 °C to quantify the amounts of N2, CO2, CH4, C2H4 and C3H6 in the product mixture via TCD (detector temperature and current of 150 °C and 120 μA, respectively). Settings of the GC units ensured reproducible detection of the specified molecules at ppm levels. Sample injection to both GC units was realized by six-way sampling valves, each of which involved sample loops of 1 mL volume.

Catalytic activity measurements were carried out by varying reaction temperature (T), CO2/G and residence time within 600–750 °C, 0–4 and 0.5–5.5 mg.min/Nml, respectively. Residence time was defined as the ratio of mass of catalyst (mgcatalyst) to the inlet volumetric flow rate of the reactive mixture (Nml/min) and was varied by regulating the catalyst quantity. Parametric study was carried out by changing a single parameter value in its pertinent range, while keeping other parameters constant at their default values (i.e. 750 °C, 1 and 0.5 mg.min/Nml). Catalyst samples tested in characterization studies as well as in 72 h time-on-stream tests, however, were obtained from experiments carried out at residence times above 0.5 mg.min/Nml for magnifying the impacts of phenomena such as sintering, deposition of carbonaceous species, etc. on the catalyst surface via its elongated interaction with the reactive mixture. In all experiments, inlet flow rate of glycerol vapor was fixed at 4 Nml/min. CO2 flow rate was determined according to the value of CO2/G ratio and N2 was used as an inert balance gas to fix the total inlet flow rate at 40 Nml/min. Except the 72 h time-on-stream runs, duration of all experiments was set as 5 h. Product sampling and analysis were first carried out for 30 min after the initiation of the experiments, and continued periodically at every 45 min. The results reported in this study were based on the arithmetic average of the outcomes of the product analysis done between 2nd and 5th h of the experiments where the reaction was found to exhibit a steady state pattern.

Catalytic performance was evaluated in terms of CO2 conversion (XCO2), glycerol conversion to gaseous products (Xglyc) and product yields (Y):

$$X_{CO_2} (%) = \frac{F_{Glyc, in} - F_{CO_2, out}}{F_{Glyc, in}} \times 100$$

$$X_{glyc} (%) = \frac{2F_{CO_2} + 4F_{CH4} + 4F_{C_2H_6} + 6F_{C_3H_8}}{8F_{Glyc, in}} \times 100$$

$$Y_i = \frac{\text{moles of species } i \text{ in gaseous products}}{\text{moles of glycerol fed}}$$

In Eqs. (9)–(11) Fi and Fout refer to the molar flow rate of species i in the product and feed streams (both in mol/min), respectively. Since product analysis was done on the basis of gaseous species, it was not possible to calculate glycerol conversion from the molar flow rate of glycerol in the product stream. In this case, as commonly done in the literature [17–24] an atomic balance over hydrogen was conducted to calculate the amount of converted glycerol. Molar flow rates of all gaseous products that contain H atoms were used in the calculation, as
shown in Eq. 10. It should be noted, however, that condensable species such as steam and oxygenated C₂+ hydrocarbons that involve H-atoms and potentially exist in the product mixture were not included in the calculation. Possible existence of hydrocarbons other than CH₄, C₂H₆ and C₂H₄ within the reaction mixture were controlled by thermodynamic analysis calculations explained in Section 3.2.1. The outcomes did not predict presence of any oxygenated C₂+ species within the field of parametric study. Moreover, owing to the very high activity of Rh-based catalysts for steam reforming of hydrocarbons [40], complete removal of the in-situ generated steam (via RWGS, Reaction 2) by its interaction with CH₄ (via Reaction 4) and with C₂H₆ and C₂H₄ was very likely to occur as also shown in Section 3.2.2. These statements could then serve for the validation of the assumptions made in the formulation of Eq. (10) which would eventually give the conversion of glycerol to gaseous products, namely H₂, CH₄, C₂H₆ and C₂H₄.

Reliability of the catalyst testing and product analysis systems were found to be reproducible in all cases within < 1% of the measurements and was verified by the outcomes (XCO₂, XH₂, and YH₂). In addition to the catalytic experiments, blank tests involving only quartz wool and diluent (α-Al₂O₃) were also conducted within the entire parameter range (T = 600–750 °C, CO₂/G = 0–4) in order to detect any activity associated with α-Al₂O₃ and to quantify the effect of temperature and CO₂/G ratio on homogeneous decomposition of glycerol. As expected, outcomes of the blank tests did not change with the existence of α-Al₂O₃ due to its low surface area (< 5 m²/g) and well-known inert nature. As additional blank experiments, pure support materials (i.e. ZrO₂ and CeO₂) were also tested at 750 °C and CO₂/G = 4 (i.e. conditions leading to the highest CO₂ conversions) to provide insight into their possible contributions to the breakdown of CO₂.

3. Results and discussion

3.1. Structural and functional characterization studies

3.1.1. Surface area and porosity measurements

Results of specific surface area (SSA), average pore volume and average pore size measurements obtained via N₂ physisorption on the support materials calcined at 800 °C for 4 h are presented in Table 1. It is observed that both ZrO₂ and CeO₂ had limited SSA, as expected from the high calcination temperatures used in the synthesis (i.e. 800 °C). These findings are in alignment with those reported in the literature. For instance, Zhao et al. [41] stated a strong correlation between ZrO₂ and calcination temperature, and reported a BET specific surface area of 15 m²/g for ZrO₂ calcined at 800 °C. Moreover, da Silva et al. [27] used the same method for preparing CeO₂ that is used in the current study and reported a SSA of 14 m²/g. A 60% greater SSA observed for ZrO₂ as compared to that of CeO₂ is in line with the better gasification ability of ZrO₂ calculated in Section 3.2.1. The outcomes associated with linear (atop) CO(ads) and CO(ads) species. Vibrational features at 2090 and 2011 cm⁻¹ can be assigned to the symmetric and antisymmetric stretchings of gem-dicarboxyl species on Rh²⁺ sites (i.e. Rh²⁺(CO₃)₄ [42–44]. Intense IR band at 2048 cm⁻¹ is associated with linear (atop) CO(ads) on metallic Rh sites. On the other hand, vibrational features appearing at lower frequencies correspond to CO adsorbed on high-coordination metallic Rh sites. Namely, 1912 cm⁻¹ signal can be attributed to CO adsorbed on two-fold (bridging) metallic Rh sites; while the IR feature at 1835 cm⁻¹ can be assigned to CO adsorption on three-fold (hollow) metallic Rh sites [45–47]. It has been demonstrated in the literature that CO adsorbed on Rh³⁺ sites in a linear (atop) fashion led to a weak vibrational signal located at 2137 cm⁻¹. As Rh³⁺ sites could readily be reduced to Rh⁺ in the presence of CO(g), detection of this feature was reported to be rather elusive [48]. Although a very weak signal is probably present in Fig. 4a at 2137 cm⁻¹, due to the extremely low intensity of this feature, it is not possible to conclusively suggest the presence of Rh³⁺-CO species on the fresh RhZr catalyst surface.

In-situ FTIR spectra given in Fig. 4a provide valuable insights regarding the electronic nature of the Rh sites on ZrO₂ support surface. Co-existence of Rh and Rh⁺ features indicates that both metallic and oxidic Rh sites may be simultaneously present on the RhZr fresh
catalyst surface. This can be explained by the exchange of oxygen between ZrO₂ lattice (generating oxygen vacancies) and the Rh metal (creating oxidic Rh⁺ species). Rh⁺ species could be also generated with the assistance of gas phase oxygen during the calcination step of the catalyst synthesis protocol.

Currently mentioned observations/explanations about the oxidation state of the precious metal sites on the metal oxide support after calcination is nothing but classical and have been reported in the literature frequently. As we clearly mentioned in the current text, rhodium can be present on the ZrO₂ surface in the forms of Rh₀, Rh⁺, or Rh³⁺. Rh³⁺ can exist on the surface due to the preservation of the original oxidation state of the Rh(III) nitrate precursor with or (more likely) without nitrate. After calcination, replacement/decomposition of the nitrate precursor to NO + O₂ and/or NO₂+1/2O₂ may lead to the replacement of nitrates with oxide anions. Moreover, detection of Rh³⁺ becomes difficult during the in-situ FTIR experiments as, in the presence of CO, oxide ions coordinated to Rh³⁺ can be consumed in CO oxidation (CO + O₂ → CO₂) leading to reduction of Rh³⁺ to Rh⁺. Finally, formation of metallic sites after calcination occurs either due to the total thermal decomposition of the Rh(III) nitrate (i.e. electron transfer from nitrate anion to Rh³⁺ and/or Rh⁺ cations during nitrate decomposition to NO + O₂ and/or NO₂+1/2O₂ or due to the reduction of Rh³⁺ and Rh⁺ with CO during the in-situ FTIR experiments. As also mentioned in the text, an alternative way of Rh⁺ generation is the oxygen vacancy formation on ZrO₂ and oxygen transfer to metallic Rh.

In-situ FTIR data also offer information about the morphology of the Rh nanoparticles. Presence of an intense IR band at 2046 cm⁻¹ corresponding to the linear (atop) CO(ads) on metallic Rh sites; co-existing with weak IR bands for CO adsorbed on two-fold and three-fold metallic Rh sites suggests that Rh particles on ZrO₂ do not expose extremely...
large (i.e. extended) facets and possess a high concentration of point defects or extended defects (e.g. corners, kinks, step edges etc.) which favor linear CO adsorption. These findings are consistent with the presence of small Rh particles on the fresh RhZr catalyst with a relatively fine dispersion. This is in line with the TEM results given in Fig. 2 revealing an average Rh particle size of 2.1 nm.

Upon aging of the RhZr catalyst for 5 h under GDR reaction conditions, striking changes appeared in the corresponding in-situ FTIR spectrum as shown in Fig. 4b. It is apparent that the Rh-CO feature at 2046 cm$^{-1}$ drastically attenuated, while the IR bands associated with CO adsorbed on high-coordination metallic Rh sites grew in intensity. One possible cause for the attenuated Rh-CO signal for the 5h-spent catalyst could be the deposition of carbonaceous species on the Rh sites and/or burial of the Rh sites by the ZrO$_2$ support material. This argument will be further justified by the current ex-situ Raman analysis of the 5 h-spent RhZr catalyst (Fig. 5). In addition, attenuation of the linear-bound CO species at the expense of the growing two-fold and three-fold adsorbed CO signals may also point to the fact that upon catalyst aging and deactivation, average particle size of Rh increased and Rh nanoparticles started to expose wider extended facets revealing a greater number of two-fold and three-fold adsorption sites. Such an argument is in very good harmony with the average Rh particle size value of the 5 h-spent RhZr catalyst obtained from TEM analysis given in Fig. 2(4.1 nm) which was significantly greater than that of the fresh RhZr catalyst (2.1 nm).

Interestingly, CO(g) adsorption on 72 h-spent RhZr revealed no vibrational features (data not shown) due to relatively severe aging, deactivation and coke deposition. In order to unravel the underlying Rh sites on the coke-covered 72 h-spent RhZr catalyst, we treated the catalyst with H$_2$(g) at 300 °C in the in-situ FTIR spectroscopic cell; in an attempt to gasify the coke and create available Rh sites for CO adsorption. These results are shown in Fig. 4c. General characteristics of the in-situ FTIR spectrum corresponding to the 72 h-spent RhZr catalyst were comparable to that of the 5 h-spent RhZr catalyst although IR intensities were typically lower in the former case. This is possibly due the fact that even after H$_2$ treatment and removal of some of the coke deposit, there exist still considerable amount of carbonaceous species on the 72 h-spent RhZr catalyst.

In the current results, decreasing intensities of the CO(ads) IR signals for the 72 h-spent RhZr catalyst may also indicate the loss of exposed Rh sites due to Rh particle size growth (i.e. decreasing dispersion) and/or migrating of Rh into the ZrO$_2$ matrix by strong metal support interaction. Evolution of the in-situ FTIR data for CO/RhZrO$_2$ upon exposure to reaction conditions may correspond to Rh diffusion into the ZrO$_2$ matrix because, increasing durations of reaction time leads to a monotonic and significant decrease in the FTIR peak intensities consistent with the loss of exposed Rh sites. On the other hand, particular loss of the intense 2046 cm$^{-1}$ and the less prominent 2080 cm$^{-1}$ bands of the fresh catalyst originating from linearly bound (on-top/atop) CO species adsorbed on kinks/edges/corners of the small clusters upon aging under reaction conditions suggest that these defect sites are replaced with non-defective terraces on larger particles where CO can bind with higher coordination (bridging & three-fold) yielding much lower vibrational frequencies in FTIR.

Fig. 4d–f show the analogous in-situ FTIR spectra obtained after CO (g) adsorption on fresh, 5 h-spent and 72 h-spent RhCe catalysts. As described earlier, for the 72 h-spent catalyst, a coke removal procedure
was also employed with H2(g) at 300 °C. Interestingly, none of these RhCe catalysts revealed any significant CO(adss) vibrational signals. It is interesting that even though the fresh RhCe catalyst did not possess any coke deposit, it still did not reveal any signs of CO adsorption. This indicates that there were no exposed Rh sites on the surface of the fresh RhCe catalyst and the Rh sites were covered with the ceria support material. It was previously highlighted that reducible oxides (i.e. CeO2) can go through the formation of strong chemical bonds to precious metals and embedding metal particles inside the lattice which can even further enhance reaction rate by up to 100 fold [49,50]. Moreover, a similar interaction between platinum and ceria was shown to lead up to 20-fold enhancement in the catalytic activity for the water gas shift reaction. This increase in activity was further confirmed by the density functional theory calculations which revealed that the electronic interaction between metal particles smaller than ca. 7 nm and ceria support can significantly decrease the activation barrier for water dissociation [51,52]. Therefore, although Rh sites seem to be covered with ceria for the fresh and spent RhCe, higher activity of RhCe catalyst could be attributed to strong electronic interaction between small Rh particles and CeO2 lattice which may possibly lead to formation of Rh-O-Ce type of surface species as well as Rh/Rh+ particles on support-metal interface which are not feasible to probe the properties of Rh sites further with CO(g) adsorption.

In addition, in the current study, it should be noted that, all catalysts were treated under identical conditions including pressure, time, temperature, mass, etc. and all measurements were repeated at least twice in order to assure reproducibility of the results. Utilized, CO pressures, exposures and durations are within typical/standard values which have been used on similar catalysts in hundreds of former studies in the literature since 1970’s. Here it is obvious that differences in the CO adsorption of Rh/Me and Rh/Zr are associated with the differences in the intrinsic nature Rh species, Rh dispersion, particle size and extent of the interaction of Rh with different support oxides, CeO2 and ZrO2. Moreover, it was pointed out that ceria can adsorb CO in the form of carbonates, carboxylates and formates [53]. These species have vibrational features below the carbonyl region of interest (i.e. < 1800 cm⁻¹) and were also observed in our current experiments (data not shown). Thus here, we merely report the absence of carbonyls on Rh sites of Rh/ Ce but do not mean to entirely exclude CO adsorption on ceria in the form of carbonates, carboxylates and formates. However, valuable information regarding the aging and coking of the spent catalysts can also be inferred via in-situ FTIR spectroscopy. This can be accomplished by investigating the differences between the vibrational features of the carbonaceous species on the 5 h-spent RhZr and 5 h-spent RhCe catalysts before and after their treatment with 10 Torr H2(g) at 300 °C for 10 min. In these set of experiments, IR and 5 h-spent Rh catalysts before and after their treatment with vibrational features of the carbonaceous species on the 5 h-spent RhZr support, respectively [57]. Furthermore, negative vibrational features appearing at 2987 and 2876 cm⁻¹ can be readily assigned to the loss of –CH2 and –CH functionalities from the RhZr catalyst surface [58,59]. Thus, it is clear that 5 h-spent RhZr catalyst was subject to severe coke deposition where a large variety of CxHyOz surface species covered both Rh/Rh+ active sites as well as the ZrO2 support. Lack of formation of significant amount of additional –OH species (except the weak signal at 3680 cm⁻¹) on 5 h-spent RhZr upon hydrogen treatment also points to the fact that most of the ZrO2 domains of the 5 h-spent RhZr catalyst was initially covered with coke and atomic hydrogen species were not able to access surface O2⁻ sites to form –OH functionalities even after gasification.

3.1.4. Ex-situ Raman spectroscopic experiments
In an attempt to further elucidate the structural properties of the fresh (i.e. reduced) and spent RhZr and RhCe catalysts (after 5 h reaction at 750 °C, CO2/G = 4, residence time = 3.75 mg.min/Nml), we performed ex-situ Raman spectroscopic analysis of these catalysts as shown in Fig. 6a and b, respectively. Topmost (black) spectrum in Fig. 6a belongs to the fresh RhZr catalyst. The feature at 173 cm⁻¹ can be attributed to RhO3 species [60–62]. This observation is in line with the in-situ FTIR data given in Fig. 4 indicating the presence of oxidative Rh species (i.e. Rh+). Six other Raman signals were also discernible in this spectrum at 211, 325, 367, 468, 548 and 611 cm⁻¹. Among these features, while 325, 468, 611 cm⁻¹ can be ascribed to tetragonal-ZrO2 structures, 211, 367 and 548 cm⁻¹ can be attributed to the characteristic features of monoclinic-ZrO2 [63–67].

The middle (red) Raman spectrum in Fig. 6a was obtained for the 5 h-spent RhZr catalyst. Observation of an extremely intense oblique baseline in this spectrum is a clear indication of fluorescence due to the presence of coke deposition on the 5 h-spent RhZr catalyst. This strong fluorescence signal overwhelms most of the Raman signatures, rendering them poorly discernible. In order to remove coke deposition (at least by a certain extent), we performed photoeleaching by increasing the laser power and irradiating the 5 h-spent RhZr catalyst using the excitation laser of the Raman spectrometer. Bottommost (blue) spectrum in Fig. 6a obtained after this process possesses almost all of the characteristic RhO3 and ZrO2 Raman signals present for the fresh RhZr catalyst suggesting that most of the coke deposit could be removed by photoeleaching. These ex-situ Raman experiments (along with the supporting in-situ FTIR data in Fig. 4) clearly indicate that RhZr catalyst suffered from severe coking during the GDR reaction.

Fig. 6b shows ex-situ Raman spectra of the fresh and 5 h-spent RhCe catalyst. The most characteristic Raman signal for the stoichiometric CeO2 lattice is located at ca. 463 cm⁻¹ associated with the F2g mode of the cubic fluorite-structure (i.e. symmetric breathing mode of the oxide ions around cerium(IV) ions) [55,68,69]. The most prominent Raman signal for the fresh RhCe catalyst appeared at 454 cm⁻¹ which was red-shifted by 9 cm⁻¹ with respect to the stoichiometric ceria. Such a shift was also observed in numerous former reports and was ascribed to the ceria unit cell expansion due to reduction of some of the Ce4+ ions in the ceria lattice to Ce3+. In addition to this intense feature, fresh RhCe catalyst also presented weaker and broader features at 240 and 571 cm⁻¹ which can be assigned to oxygen vacancies and defects in the CeO2 lattice leading to a sub-stoichiometric structure. These defects can either originate from Rh incorporation into the ceria lattice due to strong metal support interaction between Rh particles and the reducible CeO2 lattice [70–74] or from the reduction of ceria with hydrogen
which was used during the routine activation protocol of the fresh catalysts after synthesis. Moreover, another weak feature appearing at 167 cm$^{-1}$ for the Raman spectrum of the fresh RhCe catalyst can be due to the defects in the ceria lattice or possibly to Rh-O-Ce and/or RhO$_x$ species [60–62,72,73].

It can be readily seen in Fig. 6b that 5 h-spent RhCe catalyst did not reveal any fluorescence due to lack of coke deposit on this surface under GDR reaction conditions. On the other hand, it should be pointed out that there were significant changes in the spectral line shape of the Raman signals for RhCe after GDR reaction. Three features located at 167, 240 and 571 cm$^{-1}$ were significantly suppressed after the GDR reaction. It is likely that this was caused by the healing of the oxygen vacancies and defects in the substoichiometric CeO$_{2-x}$ structure and oxidation to CeO$_2$. This oxidation phenomenon is consistent with the frequency shift of the $F_{2u}$ mode to 462 cm$^{-1}$ after the GDR reaction, presumably through a mechanism resembling to the Mars-van Krevelen type. Under GDR conditions, carbon deposits on the RhCe surface could be oxidized by the utilization of mobile/active lattice oxide ions whose formation was also facilitated by Rh incorporation weakening the Ce–O bond [75,76].

3.2. Catalytic activity and stability studies

3.2.1. Effect of reaction temperature

Glycerol is a thermally unstable molecule, which tends to decompose at elevated temperatures [14]. It is worth mentioning that non-catalytic/homogenous decomposition of glycerol was often ignored in various former GDR studies in the literature, which resulted in inaccurate determination of catalytic conversion values [17–22]. Thus, in order to emphasize this important aspect and to distinguish the relative extents of catalytic and non-catalytic glycerol conversion routes, we performed detailed blank experiments. Effect of reaction temperature on glycerol conversion to gaseous products over RhZr and RhCe catalysts as well as in the blank experiments (i.e. without any catalysts) is presented in Fig. 7a. These results showed that homogeneous breakdown of glycerol was promoted with increasing temperature. At 600 °C, glycerol conversion to gaseous product in the blank tests was found to be 7%, which increased up to 38% at 750 °C. It is also apparent that at
T ≤ 650 °C and in the presence of RhZr or RhCe catalysts; no significant change in the glycerol conversion to gaseous products was observed. At 600 °C, the highest glycerol conversion was observed over RhZr, which differed from the blank test only by 6%. Similarly, at 650 °C, catalytic and non-catalytic glycerol conversions were found to be between 21–24% and ~17%, respectively. At higher temperatures, however, influence of the catalyst became notable, evident by 46% and 38% increase in glycerol conversion over RhZr (between 650–700 °C) and RhCe (between 700–750 °C), respectively. The difference between conversions obtained at 700 °C in favor of RhZr was associated with its higher catalytic activity due to the finer dispersion of Rh nanoparticles that was confirmed both by TEM-EDX analysis and in-situ FTIR spectroscopy studies in Sections 3.1.2 and 3.1.3, respectively. Fig. 7b shows the effect of temperature on CO2 conversion. Unlike glycerol, CO2 remained intact in the blank tests within the entire operating range. CO2 was also relatively stable in the presence of RhZr and RhCe catalysts where no CO2 conversions were observed in the 600–700 °C range. Breakdown of CO2 became noticeable at 750 °C where conversions of 12.6% and 7.1% were observed over RhZr and RhCe, respectively. Owing to the fact that at 700 °C glycerol conversion obtained on RhZr differed notably from that of the blank experiment (Fig. 7a), whereas no CO2 conversion was obtained under the same conditions (Fig. 7b), it could be stated that RhZr favored Reaction 3 only at 700 °C.

In order to interpret these findings, thermodynamic analysis was carried out to determine the natural limits of reactant conversions and product distributions under investigated reaction conditions. The analysis was carried out using the Gibbs Free Energy Reactor Unit-Operation (GIBS) of CHEMCAD 7.1.4 chemical process simulation software. GIBS utilizes the Gibbs free energy minimization method provided that the inlet stream is identified. During the thermodynamic analysis, glycerol, CO2, H2, CO, H2O, C(s) as well as hydrocarbons such as methane (CH4), ethylene (C2H4), ethane (C2H6), ethanol (C2H6O), acetaldehyde (C2H4O), propionaldehyde (C3H6O), allyl alcohol (C3H8O), acrolein (C3H6O) and acetyl (C2H5O2) were considered as the components that could exist in the product mixture, and the operating conditions existing during the activity tests were mimicked. The results clearly showed that neither of the conditions favored formation of C2H4 and C2H6 both of which existed only in trace quantities and presence of C(s) was suppressed with increasing temperature. Moreover, thermodynamic glycerol conversions, calculated based on the equilibrium amounts of the species specified in Eq. (10), showed that both catalysts were able to deliver 82% of the equilibrium glycerol conversion of 88.6% at 750 °C with RhZr being capable of delivering similar performance also at 700 °C (Fig. 7a). Thermodynamic limits of CO2 conversion at the studied reaction conditions are presented in Fig. 8. The results showed that below 700 °C, CO2 conversion could not be achieved thermodynamically and CO2 conversion was favored. This is in accordance with the results of Wang et al. [15], who pointed out that CO2 conversion was thermodynamically possible only above 677 °C. Thus, it is apparent that CO2 production routes such as carbon gasification with steam (e.g. Reaction 7) was dominant at lower temperatures, whereas endothermic CO2 consumption routes (e.g. Reactions (2), (5), and (8)) were thermodynamically favored at higher temperatures.

Product distributions obtained over RhZr and RhCe catalysts as well as in the blank tests at different temperatures are presented in Fig. 9a–c, respectively. It is clearly observed that yields of H2 and CO were promoted with increasing temperature in all experiments. Monotonically increasing syngas yield with temperature can be explained by the facilitated decomposition of glycerol into CO and H2 via Reaction 3, and by steam reforming and dry reforming of CH4 (Reactions (4) and (5), respectively). Fig. 9d also provides the syngas ratios (H2/CO) obtained as a function of temperature and catalyst type. H2/CO ratio seemed to have a weak dependence on temperature and remained below 0.5 for the blank tests. However, in the presence of RhZr and RhCe catalysts, they converged to ~1.1 (i.e. very close to the ideal syngas composition of 1) upon increasing the temperature to 750 °C. Increase in the H2/CO ratios in the presence of RhZr and RhCe catalysts as a function of temperature (Fig. 9d) was found to be concomitant to the corresponding changes in the H2 yields (Fig. 9a and b).

CH4 is an unwanted by-product that necessitates the post-purification of syngas. CH4 yield increased with increasing temperature both in the presence and absence of a catalyst. Increase in CH4 yield, however, was more notable at temperatures below 700 °C, above which the rate of change decreased significantly on both RhZr and RhCe. For example, rate of increase in CH4 yield between 650–700 °C and 700–750 °C was 65% and 11%, respectively on RhZr, and 78% and 13%, respectively, on RhCe (Fig. 9a and b). Conversion of CO and H2, both of which already existed in the product mixture as a result of glycerol decomposition (Reaction 3), into CH4 via reverse of Reaction 4 seemed to be the main route of CH4 production. Faster increase in CH4 yield below 700 °C was in alignment with thermodynamics, which promoted exothermic CH4 formation at lower temperatures, and with the lack of steam due to the limited impact of endothermic RWGS. The suppressed rate of increase in CH4 yields above 700 °C was likely to be caused primarily by a shift in the direction of Reaction 4 (i.e. in favor of steam reforming of methane) and by the onset of Reaction 5. These arguments were supported by the facts that CO2 (dry) reforming and steam reforming of CH4, both of which are endothermic, started to become thermodynamically significant above 650 and 620 °C, respectively [77], and steam needed to drive Reaction 4 in forward direction was provided at higher temperatures under operando conditions via RWGS, which was also endothermic and promoted at elevated temperatures. The suggested pathway seemed to hold for explaining CH4 and C2H6 yields, both of which increased with temperature and decreased significantly above 700 °C on both RhZr and RhCe. In contrast with the catalytic experiments, dampening effect of temperature between 700 and 750 °C on hydrocarbon yields was much less in blank runs (Fig. 9a–c). This finding would confirm that RWGS, which supplied steam needed for hydrocarbon consumption via reforming, occurred only in the presence of RhZr or RhCe, and the extent of homogeneous CO2 reforming of hydrocarbons was significantly smaller than that obtained heterogeneously.

Reaction temperature dictated the extent of coking in GDR [78]. Coke formation was inhibited at elevated temperatures due to the endothermic carbon gasification routes (Reactions 6–8) and became thermodynamically unfavorable above 700 °C for CO2/G = 1. A visual proof of coking hindrance at elevated temperatures was presented in Fig. 10 revealing images of the catalyst bed (RhZr + α-Al2O3) taken after 5 h testing at different temperatures. It can be observed that the extent of coke deposition decreased with temperature, as verified by the color of the bed becoming progressively lighter from 600 to 750 °C. Apart from temperature, both the extent and nature of coke formation depended also strongly on the catalyst type, as illustrated by the in-situ
FTIR spectroscopic results presented in Figs. 4–6.

Results discussed above suggest that elevated temperatures (e.g., 750 °C) could be favorable in GDR on RhZr and RhCe catalysts as they lead to the suppression of coke formation. However, it should be noted that extremely high temperatures also may result in catalyst aging via sintering of the Rh active sites. The likelihood of this phenomenon can be assessed by checking the Hüttig and Tamman temperatures of the currently utilized support materials corresponding to 0.3 and 0.5 times their melting temperatures, respectively. It is reported that at Hüttig temperature, atoms in the lattice defects become mobile, while at Tamman temperature atoms at the bulk start to demonstrate mobility causing rearrangement and sintering of the active metals [79]. By considering bulk melting points, Hüttig and Tamman temperatures of CeO₂ and ZrO₂ can be estimated to be 720 and 812.7 °C and 1200 and 1354.5 °C, respectively [79,80]. Based on these findings, default value of the GDR reaction temperature was chosen to be 750 °C, which secured the thermal stability of the support materials and was used in the rest of the performance tests where the effects of CO₂/G ratio and the residence time were explored. Even though it slightly exceeded Hüttig temperature of CeO₂, 750 °C was significantly below the related Tamman temperature (1200 °C). Moreover, as the supports materials were already calcined at 800 °C for 4 h prior to their use in catalyst preparation, no changes in their structure were expected during the reactions carried out at 750 °C.

3.2.2. Effect of CO₂/G ratio

In order to observe the effect of CO₂ concentration in the feed, CO₂/G ratios between 1 and 4 were tested at 750 °C. Experiments in the absence of CO₂ in the feed (CO₂/G = 0) were also conducted to investigate the extent of glycerol decomposition into gaseous products. The results, presented in Fig. 11a, give a clear trend of decreasing glycerol conversion to gaseous products with increasing CO₂ in the feed. The same trend characterized the relation between CO₂/G ratio and thermodynamic glycerol conversions predicted by the Gibbs free energy minimization method (Section 3.2.1) and calculated by inserting equilibrium quantities of H₂, C₂H₄, C₂H₆ and C₂H₅ into Eq. (10). The findings can be explained by the negative correlation of H₂ production with amount of inlet CO₂ due to the occurrence of RWGS (Reaction 2). As glycerol conversion to gaseous products was dictated by H₂ in the product stream (Eq. 10), its consumption by RWGS caused progressive decline of conversion with increasing CO₂/G, as commonly noted for both catalysts. This qualitative finding was observed also by theoretical predictions [15] and experimental studies [17,19] reported in the literature. It is worth noting that Eq. (10) was based on elemental
hydrogen balance over the gaseous species and did not include H2O, which was condensed in the cold traps and removed from the reactor outlet stream before GC analysis. Since RWGS produces H2O while consuming H2, the lack of H2O in the definition of Eq. (10) leads to the prediction of lower glycerol conversions together with their notable decline at higher amounts of inlet CO2. In order to test the impact of the absence of H2O on the results, it was assumed that all of the converted CO2 was spent in RWGS to produce H2O which was included in the hydrogen balance to calculate so called “modified” glycerol conversions. In other words, H2O was assumed to remain within the product mixture without being consumed. Fig. 11a presents the “modified” glycerol conversion with respect to CO2/G ratio over RhCe. It is seen in Fig. 11a that there is a less steep, but still decreasing trend for the CO2/G ratio, indicating that the absence of H2O on the results, it was assumed that all of the converted CO2 is spent in RWGS to produce H2O.

The results presented in Fig. 11b also include the evolution of CO2 conversion (T = 750 °C, residence time = 0.5 mg.min/Nml). Note that data labelled as ‘RhCe Modified’ in Fig. 11a was calculated by assuming that all of the converted CO2 is spent in RWGS to produce H2O.
residence time (0.5 mg.min/Nml) involved in the current experiments. In order to test this hypothesis, additional experiments were conducted at longer residence times which was achieved by increasing the amount of catalyst packed in the reactor while keeping the total flow rate constant at 40 Nml/min. These results, presented in Fig. 13, showed that, upon changing the residence time from 0.5 to 3.75 mg.min/Nml, CO2 conversion on RhZr increased from 22.9% to 29.2%. A further increase in residence time to 5.5 mg.min/Nml, however, led to a limited change in conversion to 29.5%. In other words, CO2 conversion converged to the thermodynamic limit of 32.6%. A similar trend was also observed for RhCe, though the relative conversion values were less than that of RhZr.

Owing to the fact that it was reported in neither of studies in the literature on Ni-driven GDR, CO2 conversion could not be used as a metric for comparison of RhZr and RhCe catalysts with the Ni-based counterparts. In this respect, the results presented in Figs. 7b and Figure 11b were unique in the literature in terms of reporting CO2 conversions under GDR conditions. Comparisons made on the basis of glycerol conversions calculated by the methodology followed in Eq. (10) showed that, even though they were tested at much shorter residence times, both RhZr and RhCe outperformed Ni-based catalysts. For example, glycerol conversion of 80% was reported on 20% Ni/cement clinker (CC: CaO + MgO) catalyst at 750 °C, CO2/G = 1.67 and residence time of ~1.5 mg.min/Nml [21], whereas more than 65% of glycerol was converted on RhZr and RhCe under the same temperature and feed composition, but with a shorter residence time of 0.5 mg.min/Nml (Fig. 11a). Moreover, glycerol conversions remained ~30% on 15% Ni/CaO, 10% Ni/ZrO2 and 5% Ag-15% Ni/SiO2 catalysts at 700 °C, CO2/G = 1, and residence times ~10 times higher than involved in the present work [23,24]. At the same temperature and CO2/G, however, RhZr and RhCe catalysts gave glycerol conversions of 70% and 35%, respectively (Fig. 7a).

3.2.3. Catalyst stability

Stability of the catalysts were examined through 72 h time-on-stream (TOS) tests carried out at 750 °C, CO2/G = 4 and residence time of 3.75 mg.min/Nml. Outcomes presented in Fig. 14a showed that even though RhCe delivered lower CO2 conversions, it exhibited superior stability as compared to RhZr. At the end of 72 h, CO2 conversion of RhZr decreased from 29% to 17.5%, which corresponded to a 40% conversion loss. In contrast, corresponding loss in conversion was only 23% for RhCe. Due to the dissimilar rates of deactivation, conversion gap between RhZr and RhCe catalysts monotonically diminished.

Comparison of the CH4 production rates, shown in Fig. 14b, also provides insight regarding differences in catalytic deactivation. While the fresh RhZr did not produce CH4, its throughput reached to ~0.8 Nml/min at the end of 72 h. In the same time span, however, CH4 production rate increased by only ~0.6 Nml/min on RhCe. As CH4 consumption was primarily due to the catalytic reforming routes, existence of CH4 in the product stream can be linked to reduced catalytic activity. Along these lines, it is apparent that RhZr deactivated faster than RhCe. Deactivation also suppressed the syngas production rate over both catalysts (Fig. 14c). However, H2/CO ratios remained almost invariant (i.e. stayed within H2/CO ~0.66 and 0.8) on both catalysts.
The absence of catalyst stability in terms of CO2 conversion in the GDR literature prevents benchmarking of the stability of the RhZr and RhCe catalysts provided in Fig. 14. Limited number of studies in the literature involved 72 h TOS testing of 3% La-20%Ni/Al2O3 and 20% Ni/CC catalysts only in terms of glycerol conversion, which decreased by ~25% and > ~50% respectively [17,21]. Interpretation of catalyst stability using glycerol conversion, however, is somewhat questionable as it does not reflect the inherent stability of the catalysts due to the presence of homogeneous reactions. The lack of contribution of non-catalytic reactions to the breakdown of glycerol in the literature also makes the reported outcomes of the TOS experiments debatable. Despite these uncertainties, RhZr and RhCe catalysts delivered truly stable glycerol conversions, which changed only by <4% on both catalysts at the end of 72 h (data not shown) and clearly outperformed the Ni-based counterparts.

4. Conclusions

Syngas formation by dry reforming of glycerol was investigated on ceria and zirconia supported Rh-based catalysts in a series of activity and stability tests, and characterization studies. Conversion of glycerol started at 600 °C, increased with temperature and reached up to 76% and 72% at 750 °C on Rh/ZrO2 and Rh/CeO2, respectively. Higher temperatures were also beneficial for suppressing carbon formation and for increasing H2/CO up to ~1, the syngas composition for improving long chain hydrocarbon selectivity in Fischer-Tropsch synthesis. In accordance with the thermodynamic calculations, CO2 conversions, reported for the first time in the literature for catalytic glycerol dry reforming, were observed only above 700 °C. Increasing the inlet molar CO2/G ratio from 1 to 4 led to a steady decline in glycerol conversions from 76% to 59% on Rh/ZrO2 and from 72% to 52% on Rh/CeO2. The same change, however, resulted in an opposite trend in CO2 conversions which first increased up to and then remained constant at 23% and 16% on Rh/ZrO2 and Rh/CeO2, respectively. These values were improved further up to 30% on Rh/ZrO2 and 20% on Rh/CeO2 by increasing the residence time. Despite its higher activity, which was related to the slightly higher specific surface area and smaller average Rh-particle size, Rh/ZrO2 deactivated faster than Rh/CeO2 in a time span of 72 h at 750 °C and CO2/G = 4. For the first time in the literature, molecular-level root causes of deactivation characteristics of the glycerol dry reforming catalysts were investigated by in-situ FTIR, Raman spectroscopy, transmission electron microscopy and energy dispersive X-ray analysis of the fresh and spent samples. These techniques consistently confirmed that the loss in activity in Rh/ZrO2 was associated with sintering of Rh nanoparticles and coke formation. These phenomena, however, occurred to a remarkably lesser extent on Rh/CeO2 and improved its stability due to (i) encapsulation of Rh nanoparticles by CeO2 as a result of strong metal-support interaction and (ii) impact of the mobile oxygen species and creation of oxygen vacancies on ceria domains that inhibited coking on the catalyst surface. Strong metal-support interaction on Rh/CeO2, however, partially suppressed the catalytic activity of the metal sites.

Declaration of interest

None.

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