

# The ever-evolving active site: transformation of single atoms to extended structures during the Rh-catalyzed reverse water–gas shift reaction

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At low temperatures (<400 °C), single atoms of Rh supported on rutile TiO<sub>2</sub> (rTiO<sub>2</sub>) are responsible for the formation of CO during the reverse water gas shift (RWGS), while methane production is associated with the Rh–TiO<sub>2</sub> interface due to the observed correlation between methane formation rates and the volume-averaged Rh nanoparticle diameter. As the temperature is increased to >540 °C, there is a notable increase in CO selectivity as the methane production rates tend towards zero. The time to reach zero depends on the temperature but is independent of the initial Rh structure (single atoms and/or nanoparticles), which is controlled by the catalyst preparation method (wetness impregnation *versus* colloidal nanoparticles). At 600 °C and >4 h time on stream, the catalytic behaviour becomes completely agnostic to the initial Rh structure as well as weight loading, and the catalysts are highly selective for the RWGS reaction. Post-reaction HR-TEM image analysis confirms Rh nanoparticles crystallize/order during the reaction; at 400 °C, most of the Rh particles are disordered, while at 600 °C, they are more ordered (*i.e.*, there is the development of defined faceting). Infrared spectroscopy of CO adsorption on Rh nanoparticles confirms the appearance of defined facets after annealing in nitrogen at high temperatures. Annealing the Rh/rTiO<sub>2</sub> catalysts prior to the RWGS reaction demonstrates the structural transformation

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of Rh depends only on time and temperature and not on reactant or product fugacity. Sites responsible for stabilizing Rh single atoms are no longer competent at higher temperatures, enabling single atom integration into existent nanoparticles. As the reaction temperature is increased to temperatures  $>540$  °C, the dominant Rh structure for CO production evolves from single atoms to extended surfaces.

## 1. Introduction

The reverse water gas shift reaction, (RWGS,  $\text{CO}_2 + \text{H}_2 \rightarrow \text{CO} + \text{H}_2\text{O}$ ) is an industrially relevant reaction to produce carbon monoxide (CO) from carbon dioxide ( $\text{CO}_2$ ) and hydrogen ( $\text{H}_2$ ). This reaction has been discussed as a central component in the so-called power to liquids scenario where water electrolysis is used to make  $\text{H}_2$ , RWGS converts this  $\text{H}_2$  and  $\text{CO}_2$  to CO, and then the resulting syngas is converted into liquid fuels using Fischer–Tropsch catalysis.<sup>1–3</sup> One of the most significant challenges with the RWGS reaction is selectivity to CO due to the competing Sabatier reaction,  $\text{CO}_2 + 4\text{H}_2 \rightarrow \text{CH}_4 + 2\text{H}_2\text{O}$ . This reaction is heavily exothermic ( $-165$  kJ mol<sup>-1</sup>), while RWGS is endothermic (41.5 kJ mol<sup>-1</sup>).

Therefore, the RWGS presents fundamental selectivity challenges. If a catalyst is unselective for RWGS *versus* methanation, then the product distribution is dictated solely by thermodynamics and high temperatures ( $\geq 600$  °C) are required to suppress  $\text{CH}_4$  formation and achieve high CO yield. Such high temperatures typically promote sintering of the dispersed active phase of the catalyst. If a catalyst kinetically controls  $\text{CH}_4$  formation, then substantial CO selectivity is possible at lower temperatures. Most heterogeneous catalysts for RWGS utilize transition metals or oxides as the active species, which are deposited on high surface area oxide supports.<sup>4</sup> Interestingly, it has been found that when Ni, Ru, and Rh supported catalysts are synthesized to promote the formation of atomically dispersed or single atom metal species, as opposed to metal nanoparticles (NPs), they are highly selective for RWGS.<sup>5–9</sup> These studies support the notion that provided the active metal can be stabilized in the form of single atoms, there are opportunities to perform RWGS at lower temperatures due to the high selectivity of the single atoms at lower temperatures. However, because of RWGS thermodynamics, operation at high temperatures is mandatory. For example, at 400 °C, for 3 : 1  $\text{H}_2/\text{CO}_2$  mixture, the equilibrium conversion is only 38.1%, it increases to 51.9% at 500 °C and is 63.1% at 600 °C (the highest temperature explored in this manuscript). Moreover, the literature teaches that regardless of whether thermodynamics are used to control RWGS selectivity by operating at very high temperatures, or kinetics are used to control selectivity by operating at low temperature using atomically dispersed metal catalysts, sintering of the active metal domains must be suppressed to promote long term catalyst stability.

As such we have explored the use of Rh/rTiO<sub>2</sub> for the RWGS reaction. Earlier work using atomically dispersed Rh/TiO<sub>2</sub> (P25) demonstrated isolated atoms and nanoparticles of the same metal on the same support can exhibit uniquely different catalytic selectivity in competing reaction pathways at a reaction temperature of 200 °C and low conversion conditions.<sup>5</sup> Moreover, the dispersion of nanoparticles under reaction conditions can play a significant role in controlling stability.<sup>5</sup> More recently, under high conversion conditions with Rh/rTiO<sub>2</sub>, it was demonstrated the product selectivity is governed by the evolving



distribution of Rh moieties under reaction conditions: the balance between Rh single atoms and Rh nanoparticle–TiO<sub>2</sub> interfacial sites determines CO *versus* CH<sub>4</sub> formation. Moreover, it appeared Rh single atoms are unexpectedly robust over at least 90 h on stream at 500 °C, even at high Rh density, whereas Rh NPs sinter.<sup>10</sup> Additionally, we reported the results of a round robin study of Rh/rTiO<sub>2</sub> catalysts for RWGS across four laboratories,<sup>11</sup> where the time-dependent production rates of CO *via* RWGS and CH<sub>4</sub> *via* methanation were measured at an appreciable approach to equilibrium. In this study the CH<sub>4</sub> formation rate rapidly decreased between 500 °C and 600 °C for higher Rh weight loadings. These studies indicate the reaction temperature plays an important role in the type and stability of the Rh structures present on the titania, motivating the current study.

In this work we report on the evolution of the reactivity and structure of a series of Rh/rTiO<sub>2</sub> catalysts under RWGS conditions at high conversions (400–600 °C, 1 atm, 75% H<sub>2</sub>, 25% CO<sub>2</sub>) as a function of time on stream (16 h). A series of rutile-supported catalysts were prepared by both wetness impregnation (WI) and colloidal synthesis (CS/NP) with nominal weight loadings ranging from 0.1 to 2 wt % Rh. The catalysts were characterized using transmission electron microscopy (TEM), CO diffuse reflectance infrared spectroscopy (CO DRIFTS), and X-ray photoelectron spectroscopy (XPS). We show that irrespective of both the catalyst preparation method and the Rh weight loading, all catalysts operated at 600 °C exhibit similar selectivity to CO (>98%) after prolonged (16 h) time on stream. TEM analysis after reaction for 16 h shows a similar mean Rh nanoparticle size among all catalysts. We show pre-treatment in an inert gas (annealing) prior to the RWGS reaction significantly improves the initial CO selectivity. We attribute these observations to a change in the ordering/crystallinity of the Rh NPs, which enables the establishment of new active sites for CO production. This work clearly demonstrates that at temperatures >500 °C, evolution of the Rh structure requires only time and temperature; the details of the CO<sub>2</sub> hydrogenation atmosphere is not a prerequisite for Rh restructuring.

## 2. Methods

### 2.1 Catalyst synthesis

**2.1.1 Synthesis of wet impregnation catalysts.** Rhodium-supported rTiO<sub>2</sub> (Rh/rTiO<sub>2</sub>) catalysts with nominal Rh loadings of 0.1, 0.3, and 2 wt% were prepared *via* wet impregnation (WI), following a previously reported procedure.<sup>11</sup> Briefly, Rh(III) nitrate hydrate (Sigma-Aldrich, ~36% Rh) was dissolved in high performance liquid chromatography-grade water and added to a suspension of rTiO<sub>2</sub> (US-Nano, 99.9%, 30 nm particle size) prepared at 12 g TiO<sub>2</sub> per liter. The Rh precursor solution was fed at 0.5 mL min<sup>-1</sup> with a syringe pump, after which the water was evaporated at 100 °C. The resulting solids were dried overnight at 100 °C, ground, and calcined under dry air at 400 °C for 4 h with a ramp rate of 10 °C min<sup>-1</sup>. These catalysts are referred to as wetness impregnated (WI) catalysts, with the designation X wt% Rh/rTiO<sub>2</sub> (WI).

**2.1.2 Synthesis of colloidal Rh nanoparticles (NPs).** Rhodium nanoparticles (Rh NPs) were synthesized based on previous work.<sup>12</sup> Briefly, RhCl<sub>3</sub> (Sigma-Aldrich, 41.8 mg), polyvinylpyrrolidone (PVP) (Sigma-Aldrich, 222 mg, MW = 55 000), glucose (Sigma-Aldrich, 720 mg), and KBr (Sigma-Aldrich, 71.4 mg) were dissolved



in 10 mL of water and added to a pressure flask (Ace Glass Inc.). The flask was sealed and heated to 140 °C under magnetic stirring for 3 h. The nanoparticles were then washed in water and separated *via* precipitation with acetone three times before being dispersed and stored in water. Thermogravimetric analysis (TGA) was performed to determine the Rh NP concentration in the solution.

**2.1.3 Deposition of the Rh NPs.** The Rh NPs were supported on a rTiO<sub>2</sub> support following an antisolvent-induced adsorption method. Typically, rTiO<sub>2</sub> (500 mg) was dispersed in water (5 mL) and sonicated for 5 min. The Rh NP solution (volume calculated to achieve desired weight loading) was then added under magnetic stirring for 5 min. Acetone was then added dropwise until precipitation occurred (final acetone : water ~ 3 : 1, v/v). The precipitate was then collected and dried in an oven overnight at 80 °C. The sample was then sieved below 180 μm grain size and calcined at 500 °C for 3 h to remove the organic ligands. Removal of the PVP ligands was ensured by conducting temperature programmed oxidation (Fig. S1) and XPS (Fig. S2). All chemicals were purchased from Sigma-Aldrich and the rTiO<sub>2</sub> was purchased from US-Nano (US-Nano, 99.9%, 30 nm particle size). These catalysts are referred to as colloidal synthesis nanoparticle (CS/NP) catalysts, with the designation *X* wt% Rh/rTiO<sub>2</sub> (CS/NP).

**2.1.4 Catalyst testing.** A full description of the reactor used in this study is presented elsewhere.<sup>11</sup> Briefly, two identical packed-bed reactors share a common gas supply (H<sub>2</sub> (99.9999%), CO<sub>2</sub> (99.999%), O<sub>2</sub> (20% in N<sub>2</sub>, >99.995%), and N<sub>2</sub> (>99.995%), Linde Praxair) which is directed through Silcolloy 2000-coated stainless lines and split into individual manifolds of thermal mass flow controllers (MFCs). The gases from each MFC were mixed downstream and passed through a 50 mL Silcolloy-2000 coated stainless preheater packed with 30-mesh SiC at 200 °C before entering a Silcolloy-coated stainless steel reactor tube (9.5 mm OD, 7 mm ID, 300 mm length). The temperature of the reactor was controlled with a clamshell furnace (Applied Test Systems, Butler, PA) using a K-type thermocouple (DwyerOmega, Michigan City, IN) inserted directly in the bed and a second K-type contacted the reactor tube wall. Reactor operations were controlled *via* LabView. The exhaust was maintained at ambient temperature and pressure to condense water and fix the water vapor pressure prior to gas chromatography analysis. Products (CO<sub>2</sub>, CH<sub>4</sub>, CO) were quantified by gas chromatography (Shimadzu 2030) with an Rt-Q-Bond PLOT column (Restek Corporation, Bellefonte, PA, 30 m, 0.53 mm ID × 20 μm). Nitrogen was used as a carrier gas and an FID with a methanizer was used for gas quantification. The initial catalytic testing of the catalysts followed a previous procedure.<sup>11</sup> In this study, we used Rh/rTiO<sub>2</sub> catalysts with variable weight loading (0.1–2 wt% nominal). All reactions were conducted with a total of 20 μg of Rh; the total catalyst amount varied for each catalyst (20 mg for 0.1 wt%, 6.7 mg for 0.3 wt%, 1.0 mg for 2.0 wt% total Rh loading) unless otherwise noted. This mass of catalyst was then Vortex mixed with nominally 500 mg of SiO<sub>2</sub> diluent (Product 84878, MilliporeSigma) for 15 s before being formed into a packed bed on top of a quartz wool support bed inside the reactor tube. The catalyst bed height was standardized within the furnace's uniform temperature zone by compressing 88 ± 3 mg quartz wool into the tube bottom and using a fixed length mounting jig to set the distance of the quartz wool support bed from the bottom of the reactor tube.

The reaction environment and thermal profile as a function of time for a representative sample is illustrated in Fig. 1a. The overall protocol for reaction



testing was (i) purge reactor tube with 20 sccm of nitrogen (>99.995%) and 100 sccm of 20% O<sub>2</sub> in N<sub>2</sub> (balance) (>99.995%), (ii) oxidation at 400 °C for 30 min (10 °C min<sup>-1</sup> ramp rate) under 20% O<sub>2</sub> in N<sub>2</sub> (balance) at 32 sccm, (iii) cool catalyst down to <40 °C under same gas flow as in (iii), (iv) purge the system of oxygen with N<sub>2</sub> (>99.995%) at 20 sccm for 25 min, (v) switch to 3 : 1 ratio H<sub>2</sub> : CO<sub>2</sub> at a total flow rate of 32 sccm for 30 min with the sample temperature <40 °C, and (vi) ramp the catalyst to reaction temperature at 10 °C min<sup>-1</sup> and hold for 16 h under same gas flow as in (v). This procedure is referred to throughout the manuscript as the normal/standard RWGS protocol (Fig. 1a).<sup>11</sup>

For annealed catalyst samples, the preparation and the reaction protocol was identical through the post oxidation purge (step iv). After this step, the catalyst was reduced and annealed prior to exposure to the 3 : 1 H<sub>2</sub>/CO<sub>2</sub> mixture. In these experiments, the Rh catalyst was reduced by exposing the sample at <40 °C to 20% H<sub>2</sub> in N<sub>2</sub> (Linde, >99.995% grade), ramping at 10 °C min<sup>-1</sup> to 200 °C and holding for 5 min under at a total flow rate of 25 sccm before purging the reactor with 20 sccm of N<sub>2</sub> (Linde, >99.995%) for 25 min. After the post reduction purge, the sample was heated to the annealing temperature at 10 °C min<sup>-1</sup> in 20 sccm of N<sub>2</sub> for a predetermined amount of time (from 1–2 h) prior to exposure to a 3 : 1 H<sub>2</sub> : CO<sub>2</sub> mixture. A representative reaction environment and thermal profile for a sample annealed at 500 °C for 2 h is shown in Fig. 1b. In Fig. 1a and b, the main reaction processes of oxidation, reduction, annealing, and being under a RWGS environment at reaction temperature are summarized. The relevant definition of elapsed time and time at reaction temperature are indicated in Fig. 1. Elapsed time is only shown to help guide the reader on the total time for the pre-treatments before collection of the catalytic data using ‘time at reaction temperature.’

The CO<sub>2</sub> conversion ( $X_{\text{CO}_2}$ ) and CO selectivity ( $S_{\text{CO}}$ ) are defined as follows, where  $n_i$  denotes the moles of species  $i$  leaving the reactor:

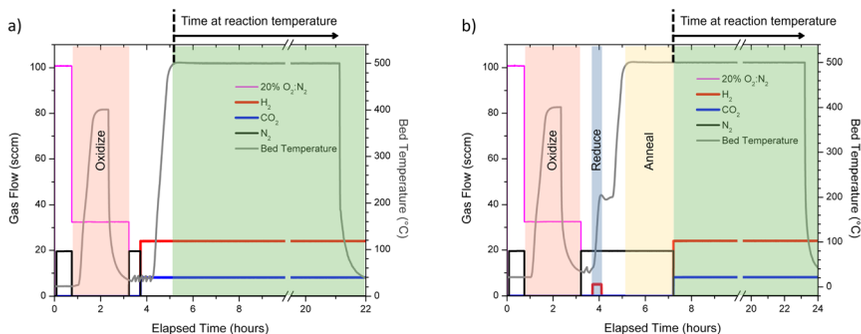


Fig. 1 Representative reaction environments and thermal profiles for non-annealed samples with the standard protocol (a) and samples that undergo annealing (b) before being exposed to the same 3 : 1 H<sub>2</sub> : CO<sub>2</sub> environment. In (b), the reduction step occurs at 200 °C under 20% H<sub>2</sub> in N<sub>2</sub> after the post oxidation purge and prior to annealing under nitrogen and introduction of the RWGS gases. The main processes of purge, oxidation (light orange), reduction (light blue), annealing (light yellow), and time at which the sample is exposed to 3 : 1 H<sub>2</sub> : CO<sub>2</sub> RWGS environment at reaction temperature (light green) are labelled.



$$X_{\text{CO}_2} = \frac{n_{\text{CO}} + n_{\text{CH}_4}}{n_{\text{CO}} + n_{\text{CH}_4} + n_{\text{CO}_2}}$$

$$S_{\text{CO}} = \frac{n_{\text{CO}}}{n_{\text{CO}} + n_{\text{CH}_4}}$$

The net production rates are defined as: “CO production rate” (and analogously “CH<sub>4</sub> production rate”) denotes CO (or CH<sub>4</sub>) production rates ( $r_{\text{CO}}/r_{\text{CH}_4}$ ) normalized per mole of total Rh. These are integral rates, rather than commonly used differential rates for turnover frequency (TOF) or space-time yield (STY) calculations and rate, and  $r_i$  is defined as  $r_i = \frac{F_i}{\text{mol}_{\text{Rh}}}$ , where  $F_i$  is the molar flow rate of CO or CH<sub>4</sub> exiting the reactor and  $\text{mol}_{\text{Rh}}$  is the total moles of Rh in the rTiO<sub>2</sub> catalyst.

## 2.2 Catalyst characterization

**2.2.1 Inductively coupled plasma-optical emission spectroscopy (ICP-OES).** The ICP-determined wt% Rh amounts (ICP-OES; Galbraith Laboratories) of the catalysts were 0.1 wt% Rh/rTiO<sub>2</sub> (WI): 0.0993 wt%, 0.3 wt% Rh/rTiO<sub>2</sub> (WI): 0.277 wt%, 2 wt% Rh/rTiO<sub>2</sub> (WI): 1.6 wt%, and 0.3 wt% Rh/rTiO<sub>2</sub> (CS/NP): 0.230 wt%. Throughout the manuscript we refer to each catalyst by their nominal loadings.

**2.2.2 CO diffuse reflectance infrared Fourier transform spectroscopy (CO DRIFTS).** Catalyst samples were characterized by CO-DRIFTS using a high-temperature Harrick Reactor cell. This reactor cell was mounted inside a Thermo Scientific Praying Mantis diffuse reflectance adapter for a Nicolet iS50 FTIR Spectrometer. The reactor cell was heated by resistive cartridge heaters controlled by a PID temperature controller with a thermocouple positioned at the bottom of the catalyst bed for accurate temperature readings. Most of the sample cup was filled with Al<sub>2</sub>O<sub>3</sub> powder (50 nm particle size), topped with a thin, flat layer of Rh/rTiO<sub>2</sub> catalyst.

All samples were pretreated *in situ* in 20% O<sub>2</sub>/He at 400 °C for 30 min (ramp 10 °C min<sup>-1</sup>) and then cooled to room temperature in the same O<sub>2</sub> flow. After purging with He, the samples were reduced in 20% H<sub>2</sub>/He at 200 °C for 5 min (ramp 10 °C min<sup>-1</sup>), followed by a 20 min He purge to remove residual hydrogen. Subsequently, the samples were heated in He to the desired annealing temperature (400, 500, or 575 °C) at a rate of 10 °C min<sup>-1</sup> and held for the specified dwell time (30 min, 1 h, or 2 h). The temperature (575 °C) represents the highest temperature the Harrick reactor cell was able to reach. After cooling to room temperature, a background spectrum was collected. The samples were then exposed to 10% CO/Ar (>99.995%, Airgas) at room temperature until the CO adsorption reached saturation coverage, after which gas-phase CO was purged with He for 45 min until all gas phase CO signatures disappeared and the final CO-DRIFTS spectrum was then recorded.

Peak deconvolution was done using OMNIC software with Voigt functions. Peak heights were allowed to be in the range of 0–10 absorbance units, Gaussian full width at half heights were allowed to be in the range of 0–32 cm<sup>-1</sup>, and



Lorentzian widths were allowed to be in the range of 0–450  $\text{cm}^{-1}$ . The noise was set to 0.05 and a baseline was not used. The initial number of peaks was determined using the peak finder with low sensitivity to reduce overestimation of the number of peaks. The number of peaks was increased until the fit was constant and the fit parameter no longer decreased.

**2.2.3 Transmission electron microscopy (TEM).** TEM characterization was performed using an objective-lens aberration-corrected transmission electron microscope (FEI Titan 80–300) at the Center for Functional Nanomaterials, Brookhaven National Laboratory (Upton, NY). The microscope was operated at an accelerating voltage of 300 kV, with a spatial resolution of 0.8 Å in the high-resolution TEM (HRTEM) mode. Post-reaction samples were deposited onto TEM grids after gently shaking the container containing both the sample and TEM Cu grid, enabling the particles to disperse onto the grid. All TEM images were acquired with a 0.5 s exposure at a “safe illumination” dose rate, as defined by continuous exposure experiments (Fig. S3) and consistent with the imaging protocol in our previous work<sup>13</sup> ImageJ was used to measure the diameter of the Rh NPs, with each size distribution derived from more than 400 particles. Potential Rh single atoms and ultra-small clusters were not included in the HRTEM-based size distribution analysis because their weak atomic contrast is overwhelmed by the strong scattering from the  $\text{TiO}_2$  support. The evaluation of ordered *versus* disordered structure was carried out using Fast Fourier Transform (FFT) analysis of the HRTEM images of Rh NPs. Only edge-on Rh particles acquired under appropriate defocus conditions were included to avoid artifacts arising from support-induced reduction in the signal-to-noise ratio. We classified NPs as ordered when sharp diffraction spots appeared in their FFTs; in the absence of such spots, they were categorized as disordered. The absence of lattice fringes and sharp diffraction spots in small crystalline Rh NPs may result from defocus, off-axis orientation, or electron-beam-irradiation-induced damage, and these factors were carefully considered to avoid misinterpretation.

**2.2.4 Ambient pressure X-ray photoemission spectroscopy (APXPS).** Ambient-pressure XPS (APXPS) measurements were performed at the Center for Functional Nanomaterials (CFN) at Brookhaven National Laboratory (Upton, NY).<sup>14</sup> The system consists of a SPECS PHI/OBOS 150 NAP hemispherical analyzer in combination with a monochromatic Al  $K\alpha$  X-ray source (1486.6 eV,  $\sim 0.25$  eV line width), focused to a spot size below 300  $\mu\text{m}$ . To reduce charging, the catalyst powder was lightly pressed onto a tantalum (Ta) foil to enhance electrical conductivity. The Ta foil with the catalyst was then mounted directly onto the sample holder *via* spotwelding. The Rh 3d spectra were energy calibrated by aligning the Ti  $2p_{3/2}$  peak of the  $\text{TiO}_2$  support to 459.3 eV to account for minor support charging effects. The spectra were fitted using CasaXPS. A Shirley background was subtracted from the spectra.

## 3. Results and discussion

### 3.1 Methane selectivity decreases with reaction temperatures $>500$ °C

We explored the impact of reaction temperature ( $>500$  °C) on CO selectivity for Rh/ $r\text{TiO}_2$  catalysts produced by two distinct methods (WI and CS/NP) containing both Rh single atoms and nanoparticles ( $d < 4.5$  nm). Fig. 2 summarizes CO and  $\text{CH}_4$  net production rates at reaction temperatures between 500 and 580 °C for 0.3 wt% Rh/



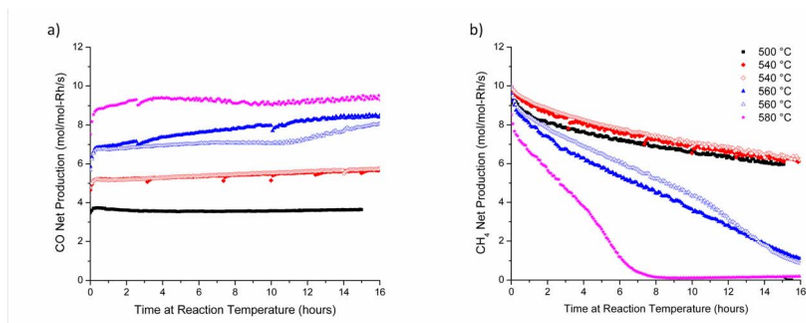


Fig. 2 CO (a) and  $\text{CH}_4$  (b) net production rates as a function of temperature for 0.3 wt% Rh/rTiO<sub>2</sub> (WI). CO net production rates increase with increasing temperature and are stable with time-on-stream. Initial  $\text{CH}_4$  production rates do not follow the same trend with temperature and decrease with increasing time on stream. The methane production rate decreases significantly for temperatures greater than 540 °C. The RWGS reaction at 540 and 560 °C were conducted in duplicate (open symbols) to demonstrate reactor reproducibility.

rTiO<sub>2</sub> (WI) catalysts over 16 h. In a recent round robin study,<sup>11</sup> the 0.3 wt% Rh/rTiO<sub>2</sub> had the highest  $\text{CH}_4$  production yield at 500 °C of all examined Rh/rTiO<sub>2</sub>; therefore, it was chosen as the initial catalyst to examine the impact of temperature above 500 °C on the production of  $\text{CH}_4$ . Fig. 2a clearly demonstrates CO net production rates increase with temperature and remain relatively constant over 16 h. Duplicate measurements at 540 °C and 560 °C demonstrate reproducible behavior between the experiments conducted on two separate diluted catalyst beds. The fate of  $\text{CH}_4$  production rates with time on stream follows very different behavior (Fig. 2b). At 500 °C (black data), the  $\text{CH}_4$  production rate continually decreases over the 16 h experimental window at a rate of  $\text{CH}_4$  loss of 0.1–0.2 mol mol<sub>Rh</sub><sup>-1</sup> s<sup>-1</sup> per hour. Similar temporal behavior in the decline of  $\text{CH}_4$  production rates is observed at 540 °C, and a duplicate catalyst bed demonstrates a similar temporal trend. At 560 °C, the rate of methane production decreases at an average rate of  $\text{CH}_4$  loss of 0.56 mol mol<sub>Rh</sub><sup>-1</sup> s<sup>-1</sup> per hour and is <1 mol mol<sub>Rh</sub><sup>-1</sup> s<sup>-1</sup> at 16 h. At 580 °C, the rate of methane production decreases at ~1.3 mol mol<sub>Rh</sub><sup>-1</sup> s<sup>-1</sup> per hour and reaches 1 mol mol<sub>Rh</sub><sup>-1</sup> s<sup>-1</sup> after 6 h, which corresponds a 60% decrease in time compared with a reaction temperature of 560 °C. The disparate behavior with temperature and time-on-stream between CO and  $\text{CH}_4$  production rates implies a loss of active sites associated with  $\text{CH}_4$  formation while the active sites associated with CO formation appear resilient, ultimately suggesting these products are produced on unique sites. These CO-resilient sites could be associated with our prior study where Rh single atoms exhibited stability for the duration of 90 h reactivity measurements. This prior study was conducted at a reaction temperature of 500 °C, which is substantially below the onset of the behavior discussed here. Utilizing the time-on-stream production rates, the CO selectivity is greater than 97% at 580 °C for times at reaction temperature greater than 8 h.

### 3.2 Methane selectivity at 600 °C is agnostic to the initial Rh structure

To determine if these observations of decreasing  $\text{CH}_4$  net production rate with temperature in excess of 560 °C were not limited solely to the 0.3 wt% Rh/rTiO<sub>2</sub>



(WI) catalyst, additional rutile-supported Rh catalysts (0.1 wt% Rh/rTiO<sub>2</sub> (WI), 2 wt% Rh/rTiO<sub>2</sub> (WI), and 0.3 wt% Rh/rTiO<sub>2</sub> (CS/NP)) were tested under the standard RWGS protocol (Fig. 1a) at a temperature of 600 °C for 16 h. Fig. 3 summarizes the CO net production rate (Fig. 3a) and CH<sub>4</sub> net production rate (Fig. 3b) for all catalysts tested. The CO net production rate (Fig. 3a) shows a general trend of constant CO production with time greater than 6–8 h on stream after a 0–30% decrease in initial CO production rates over the first several hours. The CH<sub>4</sub> production rates in Fig. 3b demonstrate all rutile-supported Rh catalysts exhibit 20–100× lower CH<sub>4</sub> production than CO production at 600 °C regardless of Rh weight loading and preparation method. During the first 4 hours on stream at 600 °C, the rapid 20–100× decrease in CH<sub>4</sub> production coupled with the more constant CO production yields a substantial increase CO selectivity such that all samples exhibit a maximum in CO selectivity during this time. There is a slow decrease in CO selectivity after the maximum, but this is not due to a decrease in CO production rate but is due to the slow increase in the CH<sub>4</sub> production rate (Fig. 3b). This behavior in CH<sub>4</sub> net production rates is also seen in Fig. 2b for the 0.3 wt% Rh/rTiO<sub>2</sub> (WI) sample at 580 °C suggesting this phenomenon is not limited to a temperature of 600 °C. Several weight loadings and catalyst preparation methods were run in duplicate to demonstrate the reproducibility with time on stream trends across different catalyst beds; all duplicate runs are noted with open symbols in Fig. 3a and b.

### 3.3 *Ex situ* TEM characterization of catalysts after reaction displays final particle size distribution

*Ex situ* TEM characterization was conducted on the post-reaction Rh/rTiO<sub>2</sub> catalysts with Rh loadings of 0.1, 0.3, and 2.0 wt% following the standard RWGS reaction conditions at 400 °C, 500 °C, and 600 °C for 16 h. Catalytic performance data for the samples tested at 400 °C and 500 °C are the same as reported previously.<sup>11</sup>

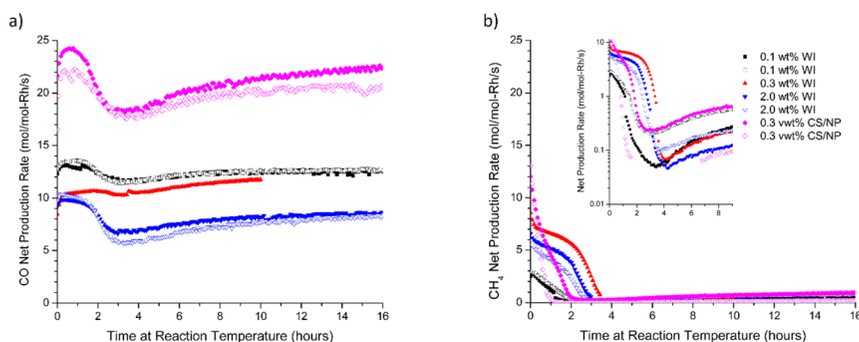


Fig. 3 (a) CO net production rates and (b) CH<sub>4</sub> net production rates for all Rh/rTiO<sub>2</sub> catalysts exposed to the standard RWGS procedure and a reaction temperature of 600 °C. A general trend of constant CO production rate with time-on-stream regardless of Rh weight loadings and preparation method is observable in (a) while (b) shows all Rh/rTiO<sub>2</sub> catalysts have very low CH<sub>4</sub> production rates (typically 0.1–0.5 mol mol<sub>Rh</sub><sup>-1</sup> s<sup>-1</sup>) at 600 °C regardless of Rh weight loading and preparation method. Duplicated reaction conditions are represented by open symbols.



For the 0.1 wt% Rh/rTiO<sub>2</sub> (WI) catalyst, a predominantly single atom catalyst in the fresh state (see Fig. S4),<sup>10</sup> the TEM image (Fig. 4a) after RWGS reaction at 400 °C reveals the formation of discrete Rh NPs, indicating reaction-induced sintering. The particle-size distribution (Fig. 4b) is relatively uniform, with an average diameter of  $\sim 1.8$  nm. The 2.0 wt% Rh/rTiO<sub>2</sub> (WI) (Fig. 4c and d) exhibits a similar mean particle size as the 0.1 wt% catalyst even though it has an order of magnitude higher loading of Rh. The comparable size across catalysts with different Rh loadings after RWGS at 400 °C regardless of weight loading suggests thermodynamic factors dominate the reaction-induced restructuring.<sup>15,16</sup>

The 0.1 wt% catalyst (Fig. 4e and f) post reaction at 500 °C exhibits a size distribution comparable to the same catalyst post reaction at 400 °C, whereas the 2.0 wt% catalyst (Fig. 4g and h) shows a broader size distribution but a similar mean size. Furthermore, the 0.1 wt% catalyst post reaction at 600 °C maintains a nearly identical mean particle size ( $\sim 1.8$  nm) and narrow size distribution

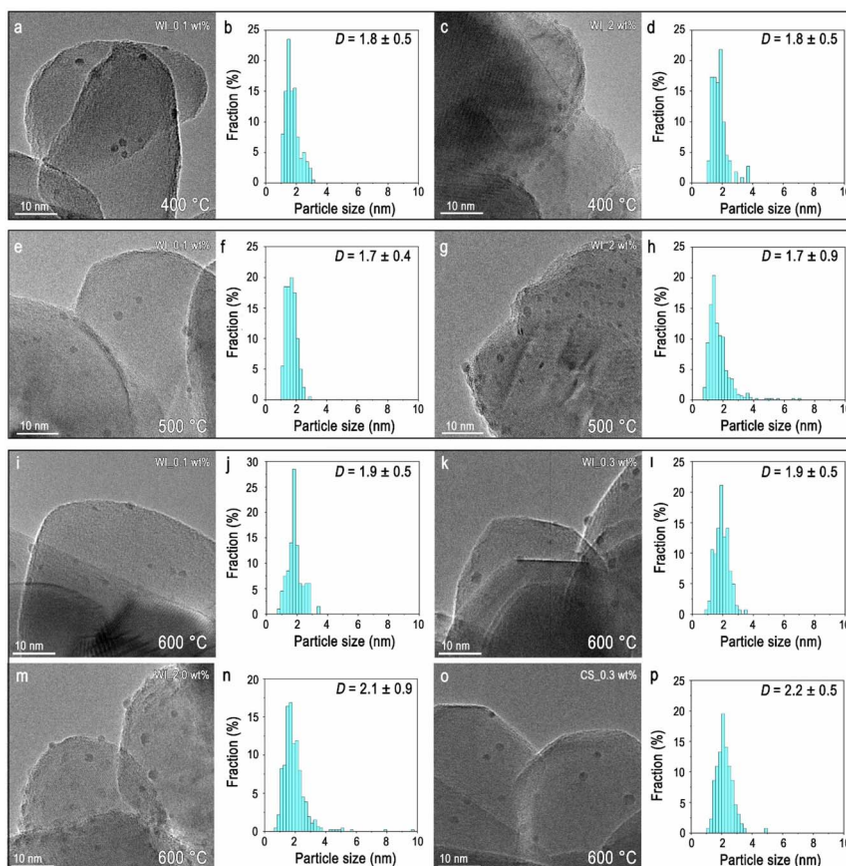


Fig. 4 TEM characterization of the post-reaction samples subjected to the standard RWGS procedure at different temperatures. Low-magnification TEM images and corresponding particle size distributions are shown for the 0.1 and 2 wt% Rh/rTiO<sub>2</sub> (WI) catalysts after 16 h of RWGS at 400 °C (a)–(d), the 0.1 and 2 wt% Rh catalysts after 16 h at 500 °C (e)–(h), and for the 0.1, 0.3, 2 wt% and 0.3 wt% Rh (CS/NP) catalysts after 16 h at 600 °C (i)–(p).



(Fig. 4i and j), consistent with large interparticle spacing suppressing particle coalescence at low loading.<sup>17</sup> For the higher-loading catalysts (0.3 and 2.0 wt%) post reaction at 600 °C shown in Fig. 4k–n, the mean particle size increases from 1.9 nm to 2.1 nm, respectively, along with a broader size distribution. To further investigate the sintering dynamics of post-reaction samples at different temperatures, the size distributions revealed low-loading samples (0.1 wt% and 0.3 wt%) maintain a narrow particle size distribution without the formation of large particles. In contrast, the 2 wt% post-reaction samples exhibit an obvious increase in the fraction of sintering-induced larger particles (6–10 nm) due to the shorter average distance between Rh species.<sup>10</sup> Importantly, the population of large particles (6–10 nm) remains a minor contribution, less than 2% as shown in Fig. 4. For the 600 °C post-reaction, 0.3 wt% Rh/rTiO<sub>2</sub> (CS/NP) catalyst, TEM imaging (Fig. 4o and p) reveals unexpected particle shrinkage after reaction, relative to the fresh sample (see Fig. S4), possibly arising from fragmentation during the catalytic reaction.<sup>18,19</sup> Surprisingly, the post-reaction particle size distributions after RWGS at 600 °C remain relatively consistent across samples with different loadings (0.1–2.0 wt% Rh) and initial Rh structures (WI or CS/NP), suggesting the final particle size is controlled by a thermodynamic equilibrium between sintering and fragmentation.

### 3.4 Differences in internal particle order post RWGS reaction

While the number of particles sampled by TEM imaging is small compared to the total number of particles present in each catalyst, the changes in particle size at the different reaction temperatures do not explain the time-dependent trends in CH<sub>4</sub> production rate and selectivity. A prior study of the RWGS conducted at 500 °C, concluded that Rh single atoms are responsible for CO formation, and Rh NPs harbour sites (assigned to Rh nanoparticle–TiO<sub>2</sub> interfacial sites) responsible for CH<sub>4</sub> production.<sup>10</sup> Fig. 3b clearly demonstrates the 0.1 wt% Rh/rTiO<sub>2</sub> (WI) catalyst produces the least amount of methane on ramp to 600 °C but ultimately has the lowest CO selectivity (albeit still >95%) at 600 °C and 16 h of reaction time. While we previously postulated a population of Rh single atoms on rTiO<sub>2</sub> are stable at 500 °C;<sup>10</sup> at 600 °C, the appearance of Rh NPs with a mean diameter of ~1.8 nm suggests these Rh single atoms may no longer be stable. Fig. 3b further demonstrates CO selectivity is >95% for all catalysts, regardless of the Rh weight loading and preparation method at 600 °C, which differs from our recent round robin observations that both a 2 wt% Rh/rTiO<sub>2</sub> (WI) or 0.3 wt% Rh/rTiO<sub>2</sub> (CS/NP) at 500 °C demonstrated a CH<sub>4</sub> selectivity of 15–20%.<sup>11</sup> Given our analysis demonstrated similar mean particle sizes after 16 h time-on-stream regardless of temperature (>400 °C), we examined the particles further utilizing HRTEM imaging. Specifically, we sought to determine if structural differences in the supported particles could provide an explanation of the temperature dependent CH<sub>4</sub> selectivity.

HRTEM images and the corresponding fast Fourier transform (FFT) patterns in Fig. 5 illustrate structural differences between Rh/rTiO<sub>2</sub> after reaction at different temperatures. We chose to examine the 2.0 wt% Rh/rTiO<sub>2</sub> (WI) since this sample had the highest particle density. HRTEM images of the catalyst post-reaction at 400 °C (Fig. 5a and S5) shows disordered atomic arrangements and irregular nanoparticle morphologies, as further evidenced by the corresponding FFT pattern lacking sharp diffraction spots. HRTEM images of the post-reaction



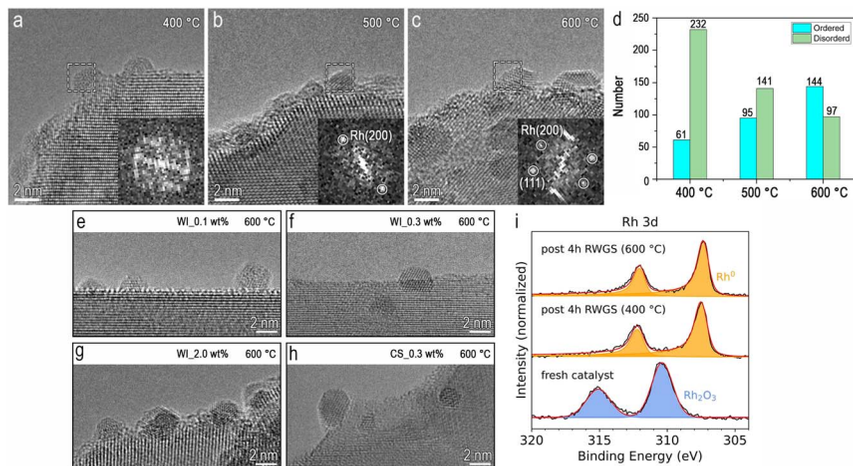


Fig. 5 HRTEM imaging and structural analysis of the post-reaction samples exposed to the standard RWGS procedure at different temperatures. (a)–(c) HRTEM images of the 2 wt % Rh/rTiO<sub>2</sub> (WI) catalyst after 16 h of RWGS at 400, 500, and 600 °C, respectively. Insets are the corresponding FFT images of the selected NPs highlighted by the dashed white rectangles. (d) Number of ordered *versus* disordered structures in the post-reaction samples as a function of reaction temperature. HRTEM imaging of post-reaction Rh/rTiO<sub>2</sub> catalysts at 600 °C with different loading: (e) 0.1 wt% Rh (WI), (f) 0.3 wt% Rh (WI), (g) 2 wt% Rh (WI), and (h) 0.3 wt% Rh (CS/NP), showing the well-crystallized and faceted Rh NPs. (i) *In situ* XPS measurements of the Rh 3d core level of the 2.0 wt% WI catalysts show the fresh catalyst has oxidized Rh and the post-reaction Rh oxidation state is metallic.

samples (Fig. 5b and S6) at 500 °C contain well-arranged atomic columns, indicating increased structural ordering. In contrast the post-reaction samples at 600 °C exhibit well-defined lattice fringes, faceted morphologies, and enhanced structural ordering, suggesting thermally promoted Rh crystallization under RWGS conditions (Fig. 5c). The crystallized Rh nanoparticles in Fig. 5b and c are the face-centered cubic (FCC) structure of metallic Rh. Fig. 5d quantifies the distribution of ordered and disordered Rh NPs at different temperatures. The results show an increase in the number of ordered Rh NPs with increasing reaction temperature. Most of the Rh NPs are well-ordered after reaction at 600 °C, whereas a vast majority possess defective or disordered structures at 400 °C. It should be noted that the disappearance of lattice fringes in some Rh NPs classified as disordered may arise from slight defocus or off-axis orientation. In this study, we maintain such variations in slight defocus and orientation affect all samples similarly and therefore do not alter the observed temperature-dependent trend. However, the majority of Rh nanoparticles have characteristic sizes of approximately ~2 nm and therefore primarily govern the observed structural trends. Moreover, the *in situ* HRTEM images (Fig. S7) reveal dynamic structural evolution between disordered and ordered states during the RWGS reaction at 500 °C. Overall, these temperature-dependent structural variations under RWGS conditions suggest dynamic restructuring processes that impact catalytic performance.

Moreover, the Rh NPs are crystallized/faceted with a similar particle size. These are depicted in Fig. 5e–h. Additional HRTEM images are depicted in Fig. S8. This



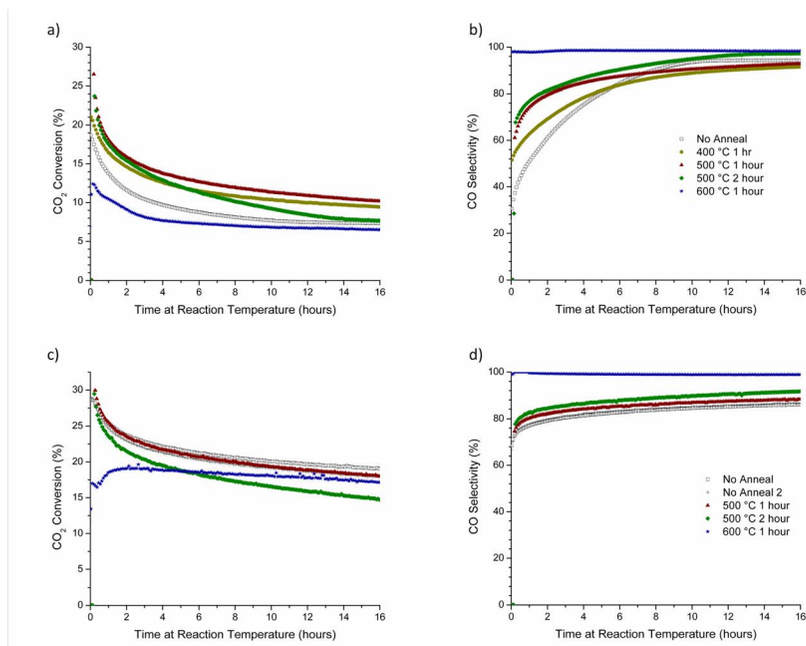
structural consistency across sample preparation methods aligns with the temperature-dependent structural variations after RWGS, providing a potential explanation for the similar catalytic behaviour at 600 °C and >4 h time on stream. These results highlight the critical role of nanoparticle crystallinity in catalytic performance of Rh/rTiO<sub>2</sub> (WI) catalysts at 600 °C. Fig. 5i shows Rh 3d XPS data that are consistent with the Rh being metallic post RWGS. The 600 °C condition results in narrower peaks, supporting the more ordered Rh seen in the TEM results (additional discussion in SI). Notably, no strong metal–support interaction (SMSI) overlayer was observed in TEM on any of the samples (400, 500, 600 °C) studied, while Rh–TiO<sub>2</sub> interactions may contribute to nanoparticle stabilization to avoid long-distance particle migration.<sup>10</sup> The lack of SMSI is further supported by XPS measurements, where the intensity ratio of Rh 3d to Ti 2p peaks is constant for the 400 and 600 °C samples, suggesting no overgrowth and Rh signal attenuation (Fig. S9 and Section S3).

### 3.5 Pre-reaction annealing of Rh/rTiO<sub>2</sub> in an inert atmosphere reduces transient formation of CH<sub>4</sub>

Given the apparent lack of SMSI, we examined whether the ripening and ordering/crystallization of the Rh is induced by the reactive gases (CO<sub>2</sub>, H<sub>2</sub>, and RWGS products – CO and H<sub>2</sub>O), temperature, time, or a combination of these. First, to gain an understanding of the effect of temperature and time on the observed CO selectivity changes during the first several hours, we examined the impact of an annealing step (*i.e.*, heating the metal catalyst in N<sub>2</sub> at a certain temperature and time) prior to the introduction of the 3 : 1 H<sub>2</sub>/CO<sub>2</sub> mixture to the catalyst. For this investigation we chose the 0.3 wt% Rh/rTiO<sub>2</sub> (CS/NP) catalyst, due to the expected structural changes it must undergo to wet the support, and the 2.0 wt% Rh/rTiO<sub>2</sub> (WI) Rh, due to the propensity to produce nanoparticles and CH<sub>4</sub> in the earlier study.<sup>10</sup>

In these annealing experiments, after the oxidation pre-treatment, Rh/rTiO<sub>2</sub> catalysts were ramped at 10 °C min<sup>-1</sup> in 20% H<sub>2</sub> in N<sub>2</sub> to 200 °C and held for 5 min. Following this reduction step the catalysts were directly heated at 10 °C min<sup>-1</sup> in N<sub>2</sub> to the stated annealing temperature and held for the different times. After this step the catalyst was heated to the appropriate reaction temperature and the gas composition was changed to the 3 : 1 H<sub>2</sub>/CO<sub>2</sub> mixture, as depicted in Fig. 1b. It is evident from Fig. 2 and 3 that all major changes are in CH<sub>4</sub> production rate while CO production rate is rather constant with time. As a result, changes in CO<sub>2</sub> conversion and CO selectivity can be used as indicators of the trends in CH<sub>4</sub> production. Fig. 6a and b show the CO<sub>2</sub> conversion and CO selectivity for 0.3 wt% Rh/rTiO<sub>2</sub> (CS/NP) and Fig. 6c and d are the CO<sub>2</sub> conversion and CO selectivity for 2.0 wt% Rh/rTiO<sub>2</sub> (WI). Regardless of whether a catalyst was annealed or not, there are no large differences in the final CO<sub>2</sub> conversion and CO selectivity after 16 h. However, there are noticeable differences in the initial CO selectivity and the rate at which CO selectivity becomes constant with annealing time (Fig. 6b and d) at temperatures from 400–500 °C. In particular, the 0.3 wt% Rh/rTiO<sub>2</sub> (CS/NP) exhibits a rapid increase in initial CO selectivity as the annealing temperature is increased from 400 °C to 500 °C with a further dramatic increase in the initial CO selectivity to >98% for an annealing treatment of 600 °C for 1 h. There is also an increase in CO selectivity with increasing annealing





**Fig. 6** (a) and (c) CO<sub>2</sub> conversion and (b) and (d) CO selectivity for 0.3 wt% Rh/rTiO<sub>2</sub> (CS/NP) (top) and 2 wt% Rh/rTiO<sub>2</sub> (WI) (bottom) after annealing in nitrogen for variable times and temperatures. For each catalyst, the total CO<sub>2</sub> conversion appears independent of annealing conditions prior to the start of the RWGS reaction at 500 °C under a 3 : 1 H<sub>2</sub> : CO<sub>2</sub> environment. Annealing the catalyst prior to exposure to RWGS conditions improves the initial CO selectivity with more dramatic improvements observed for the 0.3 wt% CS/NP catalyst than the 2.0 wt% WI catalyst. CO selectivity increases with increasing annealing temperature and time on stream for both catalyst types. For improved clarity, the repeat of the non-annealing reaction condition for the 2.0 wt% WI catalyst is represented by the '+' sign.

temperature and time for the 2.0 wt% WI catalyst, with the largest increase in CO selectivity being observed as the annealing temperature is again increased from 500 °C to 600 °C for the same 1 h duration. We attribute the different transient behavior in CO selectivity for  $T < 600$  °C for the CS/NP *versus* WI catalysts to their initial morphology. Particles present on fresh higher weight loading WI catalysts are raft-like (Fig. S4), indicative of favorable wetting behavior to the support, while the colloidal Rh NPs on the Rh/rTiO<sub>2</sub> (CS/NP) catalysts are nominally spherical and non-wetting on rTiO<sub>2</sub>. Fig. 4 and 5 demonstrate after 16 h time on stream, the NPs formed in both catalysts are similar in size and shape. The restructuring of spherical Rh NPs ( $d \sim 4.5$  nm) requires the movement of many more atoms to form faceted  $\sim 2$  nm particles compared to the raft-like particles found in WI catalysts. This temporal difference in catalyst restructuring leads to the dramatic improvement in CO selectivity for CS/NP catalysts.

### 3.6 Infrared spectroscopy of CO adsorption confirms defined Rh faceting after annealing

Fig. 7 summarizes the observed CO DRIFTS spectra for three 0.3 wt% Rh/rTiO<sub>2</sub> (CS/NP) catalysts after oxidation, reduction, and then annealing under He at



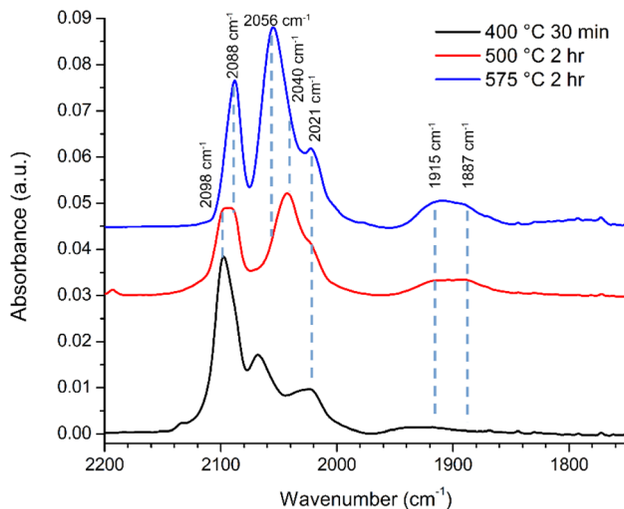


Fig. 7 Infrared spectra of CO adsorbed at room temperature under a He purge on a 0.3 wt % Rh/rTiO<sub>2</sub> (CS/NP) catalyst after annealing at 400 °C for 30 min (black), 500 °C for 2 h (red), and 575 °C for 2 h (blue). The spectra were offset vertically. Vertical dashed lines are added to aid the eye and come from peak positions found through spectra deconvolution (see Section S3). All deconvoluted spectra were assigned to specific absorption coverages on Rh crystal facets or for single atoms based on prior literature assignments for CO adsorption. Default fit parameters were used in the software with no forcing of peaks to be at any specific frequency or intensity.

various temperature/time combinations prior to exposure to CO at room temperature. The samples were annealed at 400 °C for 30 min (black), 500 °C for 2 h (red), and 575 °C for 2 h (blue). Since catalyst amounts varied between the CO DRIFTS experiments, only relative peak absorption intensities are used to compare results between different annealing conditions. However, peak intensities within each CO DRIFTS experiment can be used to define and compare relative contributions of different adsorbed CO configurations in each spectrum.

These CO DRIFTS absorption spectra are generally described by assigning the peak in the 1850–1970 cm<sup>-1</sup> region to bridge bound CO on a Rh NP, the peaks at 2020 cm<sup>-1</sup> and 2090–2100 cm<sup>-1</sup> to the asymmetric and symmetric stretches of gem-dicarbonyl on rhodium single atoms and the peak in the 2040–2070 cm<sup>-1</sup> region due to linear bound carbonyls on Rh NPs.<sup>20–22</sup> We use infrared absorption studies of CO adsorption on Rh single crystal surfaces to provide insight regarding the evolution of the different Rh crystal facets as a function of annealing condition. We emphasize that while Rh single crystal studies are free of interference from gem-dicarbonyl absorption due to their absence on extended low-energy Rh surfaces, the exact CO absorption frequencies and intensities depend on the surface coverage of CO ( $\theta_{CO}$ ) on each crystal facet.

The FTIR spectra in Fig. 7 clearly show changes in CO binding configuration as a function of annealing condition. To better understand the influence of annealing temperature each spectrum was deconvoluted. The results of this peak deconvolution for the annealed catalysts in Fig. 7 are shown in Table 1. Full



**Table 1** List of deconvoluted peaks from OMNIC using default peak fit parameters and Voigt functions. The vibrational frequencies for the bridged and corresponding linear configurations that are observable at known surface coverages of CO on different Rh surfaces as well as vibrational frequencies for the gem-dicarbonyls on single atoms are color coded for clearer matching

575 °C for 2 h	500 °C for 2 h	400 °C for 0.5 h	Peak assignment
	2097.9	2098.8	Rh(CO) <sub>2</sub> symm
2088.9	2087.9	2088.4	Rh(111)–O linear
	2069.6	2067.8	0.9–1.5 nm Rh NP linear
2056.7		2054.8	Rh(111) linear Rh(100) linear $\theta = 0.50+ \text{ML}$
2041.5	2043.5	2039.7	Rh(100) linear $\theta = 0.35\text{--}0.40 \text{ML}$
2021.1	2021.5	2022.4	Rh(100) linear $\theta = 0.20 \text{ML}$ Rh(CO) <sub>2</sub> asymm
2001.3			Rh(100) linear
		1940.1	Rh(100) bridge $\theta > 0.50 \text{ML}$
1912.6	1918.7	1912.9	Rh(100) bridge $\theta = 0.35\text{--}0.40 \text{ML}$
1885.3	1888.9		Rh(100) bridge $\theta = 0.20 \text{ML}$
		1869.7	0.9–1.5 nm Rh NP bridge

detailed peak deconvolution and fitting parameters are listed in Tables S1–S3 while the fits are shown in SI Fig. S10–S12. The deconvoluted peaks in Table 1 show that overall, the frequencies of the deconvoluted peaks were invariant in frequency and only changed in intensity with annealing conditions further supporting the idea that NPs and their final morphology evolve with time and temperature.

The deconvoluted data (Table 1) reveal a total of 11 different absorption peaks can be observed over the 3 different annealing conditions with 4 peaks assignable to CO in a bridging configuration on Rh surfaces (1870–1940  $\text{cm}^{-1}$ ), 6 peaks assignable to linear CO on Rh surfaces and/or the asymmetric stretch of Rh single atoms (2001–2088  $\text{cm}^{-1}$ ), and 1 peak associated with the symmetric stretch of Rh single atoms ( $>2090 \text{cm}^{-1}$ ).<sup>20–34</sup> (All deconvoluted peaks contained shifts of  $<\pm 1 \text{cm}^{-1}$  except the low frequency bridging and the peak at 2041  $\text{cm}^{-1}$  which shifted by  $\pm 3 \text{cm}^{-1}$  and  $\pm 2 \text{cm}^{-1}$  respectively). Of these peaks, only 3 (2021, 2056, and 2088  $\text{cm}^{-1}$ ) can be interpreted as multiple possible CO configurations belonging to either vibrations on multiple Rh crystal surfaces (2056  $\text{cm}^{-1}$ ) or Rh surfaces and single atoms (2021 and 2088  $\text{cm}^{-1}$ ). The bridge and linear frequencies for CO on Rh{100} appear for the same CO surface coverages from the surface science literature without forcing the number of peaks or the peak properties (*e.g.*, center frequency or full width at half maximum) suggesting the sizes of the facets developed during the annealing are energetically equivalent to extended Rh surfaces that can hold up to 0.75 monolayers of CO.<sup>23–25</sup>

As noted, detailed descriptions of the deconvoluted peaks and their frequencies can be found in Tables S1–S3 and Fig. S10–S12. The most important results of the deconvolutions of the FTIR spectra are the physical changes in Rh–CO binding configurations as a function of annealing temperature. These observed



changes and their physical meaning regarding Rh NP development are detailed below.

First, there is a noticeable shift in frequency and an increase in intensity for the infrared bands associated with bridging CO ( $1850\text{--}1970\text{ cm}^{-1}$ ) on the Rh{100} and Rh{111} surfaces<sup>23–26</sup> as the annealing temperature is increased. Table 1 shows these bridging carbonyls can be assigned to Rh{100} surfaces that accommodate surface coverages of  $\theta_{\text{CO}} = 0.20$  and  $\theta_{\text{CO}} = 0.35\text{--}0.40$  ML. Our FTIR results suggest Rh{100} surfaces develop as the annealing temperature is increased, which agrees with the Rh{100} surfaces observable in the post-reaction HRTEM images (Fig. 5, S5, S6 and S8).

Second, all deconvoluted spectra exhibit absorption peaks at  $2021\text{ cm}^{-1}$  and  $2041\text{ cm}^{-1}$ . The absorption peak at  $2041\text{ cm}^{-1}$  has been observed on Rh{100} also with a surface coverage of  $\theta_{\text{CO}} = 0.35\text{--}0.40$  ML further suggesting the existence of Rh{100} facets. The absorption peak around  $2020\text{ cm}^{-1}$  can be associated with CO bound in a linear configuration on Rh{100} with  $\theta_{\text{CO}} = 0.20$  ML thus matching the same surface coverage for the bridging CO at nominally  $1887\text{ cm}^{-1}$ . It should be noted the peak at  $2020\text{ cm}^{-1}$  could also be associated with the asymmetric stretch of CO bound to Rh single atoms implying the Rh CS NPs are fragmented by CO with increasing annealing temperature.

Third, the peak at  $2056\text{ cm}^{-1}$  grows from near zero intensity for the catalyst annealed at  $400\text{ }^{\circ}\text{C}$  to a dominant peak for the catalyst annealed at  $575\text{ }^{\circ}\text{C}$  while the readily observable  $2068\text{ cm}^{-1}$  peak observable at  $400\text{ }^{\circ}\text{C}$  decreases to zero with increasing annealing temperature suggesting these peaks are linked to Rh crystallite development. The  $2056\text{ cm}^{-1}$  peak is most appropriately assigned to linear bound CO on Rh{111}<sup>25,26</sup> suggesting Rh{111} facets develop with increasing annealing temperature and support for the existence of these Rh{111} facets can be observed in Fig. 5. The broadening of the  $2056\text{ cm}^{-1}$  peak on the low frequency side is ascribed to linear CO absorption peaks on Rh{100} facets in the  $2041\text{ cm}^{-1}$  region. However, the peak at  $2068\text{ cm}^{-1}$  cannot be assigned to CO bound to low index Rh crystal faces. Instead, a peak at  $2068\text{ cm}^{-1}$  has been reported for CO in a linear configuration on small Rh NPs ( $0.9\text{--}1.5\text{ nm}$ ).<sup>32</sup> These small Rh NPs also exhibited an absorption peak for bridge bound CO at  $1868\text{ cm}^{-1}$  and a discernible absorption peak is observable at  $1869\text{ cm}^{-1}$  in the spectrum for the catalyst annealed at  $400\text{ }^{\circ}\text{C}$  (Fig. S10 and Table S1). Typically, small NPs are highly stepped and the adsorption position of CO on (111) stepped surfaces such as (331) and (755) have not been widely reported. The closest report is for CO in a linear configuration on Rh(331) at  $2060\text{ cm}^{-1}$ .<sup>33</sup> Therefore, it is reasonable to associate this peak at  $2069\text{ cm}^{-1}$  with highly stepped surfaces of Rh{111}, such as {211} and {311} that would be present in irregularly shaped Rh NPs and observable in the HRTEM images in Fig. 5 of Rh/rTiO<sub>2</sub> exposed to the RWGS reaction at temperatures less than  $575\text{ }^{\circ}\text{C}$ . As noted above, this peak at  $2068\text{ cm}^{-1}$  associated with Rh NPs ( $0.9\text{--}1.5\text{ nm}$ ) and stepped Rh{111} surfaces is noteworthy because of its prominence in Fig. 7 for the catalyst annealed at  $400\text{ }^{\circ}\text{C}$  while it is absent in the IR spectra of catalysts annealed at  $575\text{ }^{\circ}\text{C}$ . Its presence at low temperatures but disappearance at higher annealing temperatures suggests the development of extended Rh structures with increasing annealing temperature. Rh nanoparticles are  $>1.5\text{ nm}$  in diameter and/or Rh{111} facets are no longer stepped when the  $2068\text{ cm}^{-1}$  peak disappears. The  $<10\text{ cm}^{-1}$  shift in frequency can easily be due to oxygen also bound to the Rh surfaces since CO is known to dissociate on {211},<sup>34</sup>



{331} and {755} steps<sup>35</sup> and B5 and B6 defect sites associated with {111} crystal facets.<sup>36</sup> The peak at 2068 cm<sup>-1</sup> agrees well with a stepped or 'roughened' Rh surface as an asymmetrical shaped CO vibrational peak at 2070 cm<sup>-1</sup> was observed on roughened Rh.<sup>37</sup>

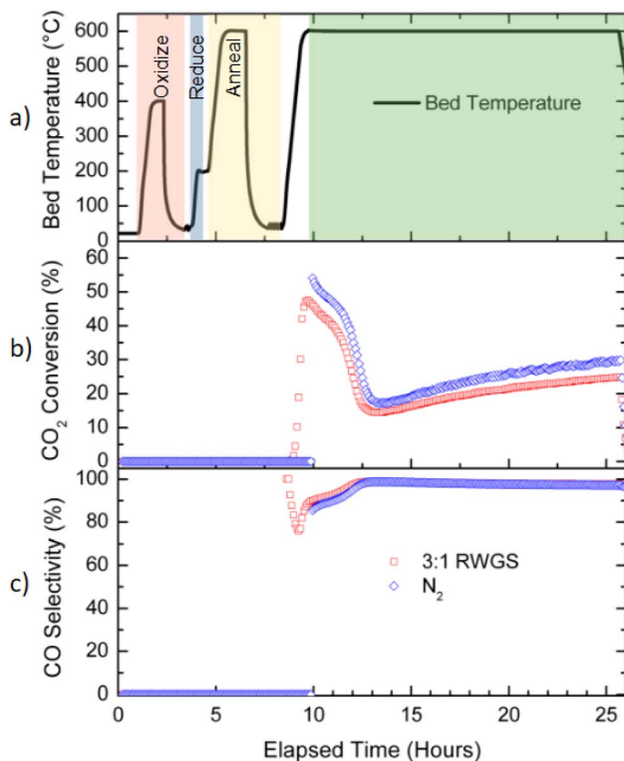
Finally, the peak between 2080 and 2100 cm<sup>-1</sup> is comprised of two distinct peaks whose intensities shift depending on the annealing temperature. For the sample annealed at 400 °C, the asymmetry of the peak suggests the higher frequency peak is larger in intensity than the lower frequency peak while for the sample annealed at 575 °C, the opposite is true. Peak deconvolution yields peaks at 2088 cm<sup>-1</sup> and 2098 cm<sup>-1</sup> for catalysts annealed at 400 and 500 °C with only the 2088 cm<sup>-1</sup> peak being present with a slightly larger full-width at half maximum (Tables S1–S3) for the sample annealed at 575 °C. The peak at 2098 cm<sup>-1</sup> can be assigned to the symmetric absorption vibration of the rhodium gem-dicarbonyl since CO bound to Rh{111} has not been observed at frequencies higher than 2088.4 cm<sup>-1</sup>.<sup>38</sup> Since Rh{111} facets can be observed from the truncated and non-regularly shaped octahedral Rh NPs in the HRTEM images in Fig. 5 and S8, and since it is well known CO can dissociate at stepped {111} and undercoordinated sites in fcc metals,<sup>31–33</sup> it is highly plausible the 2088 cm<sup>-1</sup> peak can be assigned to CO in a linear configuration on Rh{111} with O also present on the Rh{111} surface. Beyond the TEM results, support for this can be inferred from the 2098 cm<sup>-1</sup> : 2088 cm<sup>-1</sup> peak area ratio which decreased from ~3 : 1 for the 0.3 wt % Rh/rTiO<sub>2</sub> (CS/NP) catalyst annealed at 400 °C to 0 for the catalyst annealed at 575 °C. The loss of the Rh–CO vibrational signature is most definitively associated with the disappearance of Rh single atoms with increasing annealing temperature. These general observations and their changes with increased annealing temperature suggest Rh is dynamic during the annealing process and evolving with increasing annealing temperature.

### 3.7 Pre-annealing of Rh/rTiO<sub>2</sub> (CS/NP) demonstrates time and temperature determine the agnostic behavior of CO selectivity to initial Rh structure

When heating to the reaction temperature (600 °C) under RWGS reaction conditions (Fig. 3) or with the inclusion of a pre-anneal step prior to exposure to RWGS reactions conditions (Fig. 6), the initial CO selectivity is significantly higher because of the annealing step. There are slight differences in the non-annealing and annealing pre-reaction protocols that we eliminate in a revised reaction protocol. In the revised procedure, two 0.3 wt% Rh/rTiO<sub>2</sub> (CS/NP) catalysts were oxidized at 400 °C, reduced at 200 °C, and then annealed at 600 °C for 1 h under N<sub>2</sub>. However, instead of exposing the catalysts to the RWGS environment as was done for annealed catalysts (in Section 3.5), both catalysts were cooled back down to < 40 °C under N<sub>2</sub>. One of the catalyst samples was exposed to a 3 : 1 H<sub>2</sub> : CO<sub>2</sub> gas mixture while the other remained under an inert nitrogen environment before both were simultaneously ramped to 600 °C. Once at a reaction temperature of 600 °C, the sample under N<sub>2</sub> was exposed to the 3 : 1 H<sub>2</sub> : CO<sub>2</sub> gas mixture. The CO<sub>2</sub> conversion and CO selectivity as function of elapsed time can be seen in Fig. 8b and c respectively. In Fig. 8a, the reactor temperature, oxidation, reduction, and annealing processing steps are shown using the same color scheme as in Fig. 1.



The catalysts show very similar  $\text{CO}_2$  conversion (Fig. 8a) and CO selectivity as a function of time once both were under RWGS reaction conditions at 600 °C. The catalyst heated from <40 °C to 600 °C under  $\text{N}_2$  prior to the start of the RWGS reaction had a noticeable relative improvement of 10+% in  $\text{CO}_2$  conversion throughout the entire time-on-stream period. Both samples have the same CO selectivity (Fig. 8b) as a function of time despite one catalyst being ramped to the reaction temperature under  $\text{N}_2$  and the other under RWGS; the similarity in CO selectivity suggests the Rh catalyst is structurally modified during the annealing step and prior to exposure to RWGS reaction conditions. It is not clear why these catalysts started with an initial CO selectivity of >85% rather than >95% as was previously observed for the sample in Fig. 6b that did not undergo cooling back to



**Fig. 8** Influence of heat to reaction temperature environment on RWGS performance. (a) Pretreatment and reaction environment with corresponding thermal profile. (b)  $\text{CO}_2$  conversion and (c) CO selectivity for two 0.3 wt% Rh/rTiO<sub>2</sub> (CS/NP) as a function of reaction time at 600 °C. Both catalysts were exposed to the same pre-reaction conditions of oxidation, reduction, annealing at 600 °C for 1 h, and cooled back to <40 °C before being ramped to reaction temperature. One catalyst was ramped to a reaction temperature of 600 °C under a 3 : 1  $\text{H}_2$  :  $\text{CO}_2$  environment while the other catalyst was ramped to a reaction temperature of 600 °C under  $\text{N}_2$  and only exposed to the 3 : 1  $\text{H}_2$  :  $\text{CO}_2$  environment after the catalyst bed reached 600 °C. Both samples exhibit very similar  $\text{CO}_2$  conversion as a function of time and almost identical CO selectivity as a function of time suggesting the changes in the Rh catalyst occurred prior to being exposed to the RWGS on the ramp to the reaction temperature and the RWGS environment does not affect the Rh catalytic performance on the ramp to the reaction.



<40 °C before being ramped back to 600 °C. Both samples started with a substantially higher CO selectivity than the sample shown in Fig. 6b that underwent a 2 h anneal at 500 °C which only had a starting CO selectivity of about 65%. It is evident the annealing of Rh at higher temperatures (600 °C) improves CO selectivity. Annealing of Rh prior to RWGS reaction at 600 °C for as little as 1 h and/or a RWGS reaction temperature more than 560 °C yields a RWGS catalyst that is highly selective to CO and stable with respect to time.

### 3.8 Invariance of the apparent activation energy for CO formation with the initial Rh structure demonstrates a common active site

After 16 h time-on-stream, temporal changes in CO<sub>2</sub> conversion and CO selectivity are rather small, suggesting the reaction has reached steady-state and/or changes in the catalyst structure are minimal. At this point, the samples were cooled *via* thermal loss under the 3 : 1 H<sub>2</sub> : CO<sub>2</sub> mixture. During this cool down, CO and CH<sub>4</sub> production rates were calculated as a function of temperature to generate an Arrhenius plot. Fig. 9a and c shows the procedure and the Arrhenius plot for a non-annealed 2.0 wt% Rh/rTiO<sub>2</sub> (WI) for both CO and CH<sub>4</sub> over a temperature range of 390 °C to 240 °C. The apparent activation energy ( $E_a$ ) for CO production for the 2.0 wt% Rh/rTiO<sub>2</sub> (WI) is 57 kJ mol<sup>-1</sup>. Duplicate measurements of the “cool down”  $E_a$  over the non-annealed 2.0 wt% Rh/rTiO<sub>2</sub> (WI) showed good reproducibility.

The 0.3 wt% Rh/rTiO<sub>2</sub> (CS/NP) catalyst annealed at 600 °C for 1 h prior to the RWGS reaction in Fig. 8 yields a “heat-up”  $E_a$  for CO production of 60.1 kJ mol<sup>-1</sup> and a “cool down”  $E_a$  for CO production of 60.8 kJ mol<sup>-1</sup> both of which agree with the post reaction CS/NP average CO activation energy of 60.5 kJ mol<sup>-1</sup> for the non-annealed CS/NP samples that were exposed to RWGS at 500 °C for 16 h. This suggests that catalysts which undergo a high temperature (600 °C) pre-annealing step prior to exposure to RWGS reaction conditions obtain a Rh morphology (size and degree of ordering) similar to catalysts exposed to RWGS reaction conditions at 500 °C for 16 h.

The fresh 0.3 wt% Rh/rTiO<sub>2</sub> (CS/NP) contains NPs with a mean spherical particle size of 4.5 nm with little to no single atoms (Fig. S4), while the fresh 2.0 wt% Rh/rTiO<sub>2</sub> (WI) contains a mixture of single atoms and raft-like Rh NPs (Fig. S4). Upon reaction under 3 : 1 H<sub>2</sub> : CO<sub>2</sub> at 600 °C for 16 h, both catalysts contain faceted ~2 nm Rh NPs. Since these catalysts contained a large fraction of NPs in the fresh state (albeit with different structure), we conducted a 600 °C anneal in N<sub>2</sub> for 1 h prior to the RWGS reaction on a single atom catalyst, 0.1 wt% Rh/TiO<sub>2</sub> (WI). TEM analysis (Fig. 4, S5, S6 and S8) demonstrates that regardless of reaction temperature, Rh NPs of ~1.8 nm in diameter are observed. However, the HRTEM analysis was unable to observe Rh single atoms, and we anticipate the population of stable Rh single atoms decreases with reaction temperature. The apparent activation energy on Rh single atoms supported on rutile TiO<sub>2</sub> is 101 kJ mol<sup>-1</sup>,<sup>28</sup> while the apparent activation energy on the Rh/rTiO<sub>2</sub> (CS/NP) catalyst is 60 kJ mol<sup>-1</sup>. If the 0.1 wt% Rh/rTiO<sub>2</sub> (WI) catalyst still contains single atoms after reaction or pre-annealing, the measured apparent activation energy should reflect a contribution from both the single atoms and ~1.8 nm NPs. We annealed the 0.1 wt% Rh/TiO<sub>2</sub> (WI) catalyst at 600 °C in N<sub>2</sub> for 1 h, cooled to <40 °C and heated (10 °C min<sup>-1</sup>) to 600 °C under RWGS reaction conditions, followed by 16 h time-



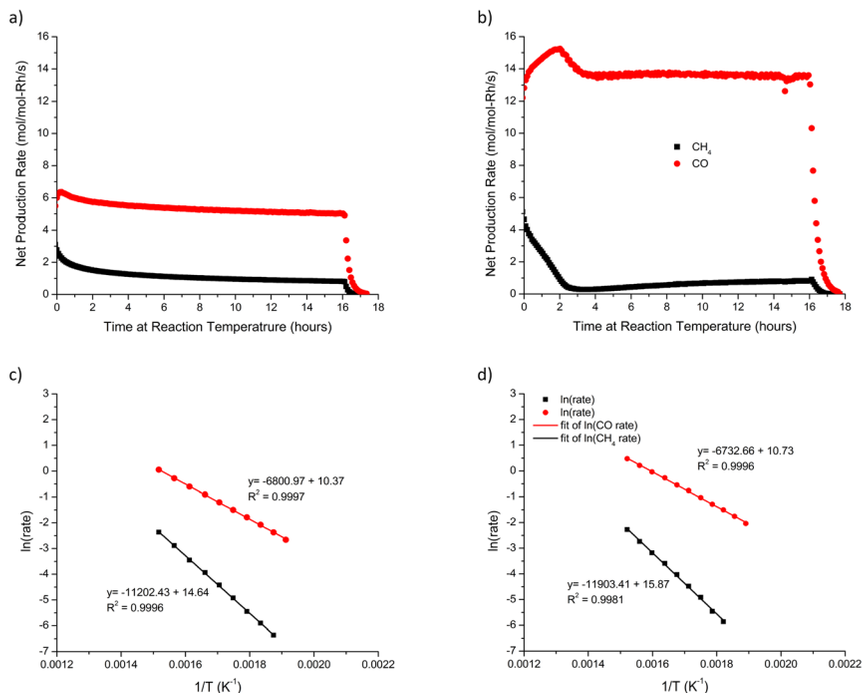


Fig. 9 CO and CH<sub>4</sub> rates for (a) non-annealed 2.0 wt% Rh/rTiO<sub>2</sub> (WI) and (b) pre-annealed (600 °C, 1 h) 0.1 wt% Rh/rTiO<sub>2</sub> (WI) exposed to 3 : 1 H<sub>2</sub> : CO<sub>2</sub> mixture for 16 h and their associated cool down activation energy plot data, (c) and (d) respectively, as function of time and temperature. The 2.0 wt% Rh/rTiO<sub>2</sub> (WI) sample was run at 500 °C and the 0.1 wt % Rh/rTiO<sub>2</sub> (WI) sample was run at 600 °C. The data used for the cool down dynamic activation energy plot can be seen in both image (a) and (b) for times greater than 16 h. The activation energy data plot was fit over a temperature range of nominally 240–390 °C where conversion was 0.1–6%.

on-stream, and subsequent measurement of the  $E_a$  upon “cool-down”. Fig. 9b represents the time on stream CO and CH<sub>4</sub> production rates of a pre-annealed (600 °C, 1 h) 0.1 wt% Rh/rTiO<sub>2</sub> (WI) catalyst. Upon completion of 16 h RWGS reaction, the catalyst sample was cooled down *via* thermal losses from the furnace. Fig. 9d is the Arrhenius plot for the annealed 0.1 wt% Rh/rTiO<sub>2</sub> (WI). The value for the  $E_a$  for CO production is 57.3 kJ mol<sup>-1</sup>. A second measurement on a separate bed of the annealed 0.1 wt% Rh/rTiO<sub>2</sub> (WI) yielded an apparent  $E_a$  of 56 kJ mol<sup>-1</sup>. The measured  $E_a$  during the cool down after 16 h time-on-stream suggests Rh NPs are responsible for CO formation even though the initial Rh structure was dominated by single atoms. The measured apparent activation energy for the RWGS reaction over Rh NP/TiO<sub>2</sub> (P25) has been reported to be 55 kJ mol<sup>-1</sup> for a 4 : 1 H<sub>2</sub> : CO<sub>2</sub> ratio.<sup>39</sup> The measured apparent activation energy in this work at a 3 : 1 H<sub>2</sub> : CO<sub>2</sub> ratio is similar, but significantly lower than the value (101 kJ mol<sup>-1</sup>) reported on rutile-supported single atoms of Rh.<sup>28</sup> A value of 55–60 kJ mol<sup>-1</sup> for the apparent activation energy appears to be independent of the TiO<sub>2</sub> polymorph (rutile *versus* anatase).<sup>39</sup>



The apparent  $E_a$  for  $\text{CH}_4$  formation for the 2.0 wt% Rh/rTiO<sub>2</sub> (WI) catalyst in a 3 : 1 H<sub>2</sub> : CO<sub>2</sub> mixture upon cool down is  $\sim 102.8$  kJ mol<sup>-1</sup> (Fig. 9c), while the  $E_a$  upon cool down on the pre-annealed 0.1 wt% Rh/rTiO<sub>2</sub> (WI) is 99.7 kJ mol<sup>-1</sup> (Fig. 9d). Both experiments were run in duplicate, and the reproducibility between individual catalyst beds was good. These values for the apparent  $E_a$  for methane formation agree well with the reported value of Verykios and co-workers of 103 kJ mol<sup>-1</sup>.<sup>39</sup> Reported  $E_a$  values for CO<sub>2</sub> hydrogenation to CH<sub>4</sub> in a 3 : 1 H<sub>2</sub> : CO<sub>2</sub> mixture over a Rh foil<sup>40,41</sup> or an inverse TiO<sub>2</sub>/Rh foil catalyst<sup>42</sup> are in the range of 130–140 kJ mol<sup>-1</sup>, while a value of 100 kJ mol<sup>-1</sup> was reported over an oxidized Rh foil.<sup>41</sup> The latter value agrees well with the apparent activation energy for CH<sub>4</sub> formation reported here and by Verykios *et al.*<sup>39</sup> The difference in the activation energy reported here compared with those from a Rh foil suggest that undercoordinated sites or tightly bound atomic oxygen at those undercoordinated sites on the Rh nanoparticle surface may be critical for the kinetically relevant CO hydrogenation step.

## 4. Conclusions

Rh single atoms supported on rutile TiO<sub>2</sub> are responsible for CO formation during the reverse water gas shift reaction at low temperatures (<400 °C). As the reaction temperature is increased, the formation of CH<sub>4</sub> is observed which is associated with the growth of Rh nanoparticles and the formation of a Rh–TiO<sub>2</sub> interface, the presumed location of the active site for CH<sub>4</sub> formation. The Rh nanoparticles formed at low temperature are primarily disordered, and CO IR spectroscopy on these catalysts confirms significant Rh speciation (single atoms and nanoparticles). Upon heating to higher temperatures (550–600 °C), the formation of methane decreases significantly even though nanoparticles and their interface with TiO<sub>2</sub> persist. At higher temperatures, the particles become more ordered regardless of their initial structure (single atoms or nanoparticles), and IR spectroscopy of adsorbed CO suggests the formation of CO–Rh terrace atom configurations. At a reaction temperature of 600 °C under RWGS conditions (3 : 1 H<sub>2</sub> : CO<sub>2</sub>), the selectivity to CO for all Rh/rTiO<sub>2</sub> catalysts is >95%, and the apparent activation energy for multiple Rh/rTiO<sub>2</sub> with different initial Rh structures, either pre-annealed at 600 °C or post RWGS (600 °C for 16 h) was 55–60 kJ mol<sup>-1</sup>, in good agreement with reported activation energies measured on nanoparticle Rh/rTiO<sub>2</sub> catalysts. The development of faceted Rh nanoparticles leads to the predominant formation of CO at higher reaction temperatures. Single atoms may persist at these higher temperatures, but the predominance of adsorbed CO associated with extended Rh surfaces and the agreement in measured apparent activation energy on nanoparticle-based Rh/rTiO<sub>2</sub> catalysts suggest single atoms are kinetically irrelevant at reaction temperatures of  $\sim 600$  °C. This study demonstrates detailed examination of catalyst activation is critical for the prediction of catalyst performance with time on stream.

## Author contributions

GB: conceived the initial research idea, catalyst testing, data analysis, original draft writing. XC, AK, JH, SG, LM, DZ: catalyst synthesis, catalyst characterization, discussions, writing and editing. JY, AH, MC, and RR: research supervision,



writing and editing. SRB: funding acquisition, project administration, resources, supervision, writing and editing.

## Conflicts of interest

There are no conflicts of interest to declare.

## Data availability

All data supporting the findings of this investigation are incorporated within the publication and the supplementary information (SI). Supplementary information is available. See DOI: <https://doi.org/10.1039/d5fd00172b>.

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