Dynamic structure of active sites in ceria-supported Pt catalysts for the water gas shift reaction

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Oxide-supported noble metal catalysts have been extensively studied for decades for the water gas shift (WGS) reaction, a catalytic transformation central to a host of large volume processes that variously utilize or produce hydrogen. There remains considerable uncertainty as to how the specific features of the active metal-support interfacial bonding—perhaps most importantly the temporal dynamic changes occurring therein—serve to enable high activity and selectivity. Here we report the dynamic characteristics of a Pt/CeO2 system at the atomic level for the WGS reaction and specifically reveal the synergistic effects of metal-support bonding at the perimeter region. We find that the perimeter Pt0−O vacancy−Ce3+ sites are formed in the active structure, transformed at working temperatures and their appearance regulates the adsorbate behaviors. We find that the dynamic nature of this site is a key mechanistic step for the WGS reaction.
The WGS reaction (CO + H₂O → CO₂ + H₂) is an essential processing step in the industrial production of synthesis gas—one used both to purify and to adjust the H₂ composition in important downstream processes such as the Fischer–Tropsch synthesis of hydrocarbons (CO:H₂ = n:(2n + 1)), methanol synthesis (CO:H₂ = 1:2), and the Haber–Bosch process for the production of ammonia. Among all WGS catalysts, ceria-supported noble metal (Pt and Au) catalysts have been heavily investigated recently because they have excellent low-temperature activity, selectivity, and stability, and they are nonpyrophoric (an important feature for such uses as in polymer electrolyte membrane (PEM) fuel cells). The identity of the active species/sites of ceria-supported Au or Pt catalysts and the roles of the supports in the WGS reaction have been sources of intense debate. For example, several groups reported that the nonmetallic gold or platinum species/sites strongly associated with the support are responsible for the catalytic activity with the metal of the nanoparticles acting as spectator species. Ding et al., however, observed that only metallic Pt nanoparticles show activity in gas that nonmetallic Pt single atoms (Pt⁶⁺) are spectators on a variety of supports. In terms of the support effects, Zhai et al. reported that a support only serves to stabilize the Pt⁶⁺ sites. Mechanistic/kinetic studies, based on advanced transient isotopic techniques, on the other hand, suggest that catalyst support provides active sites within a region (reactive zone) around the Pt nanoparticles. Supports, especially oxygen vacancies of the supports, play an essential role in the activity of the catalyst because water dissociation, involved in two main mechanisms (the regenerative/reodox and the associative mechanisms), is believed to occur at oxygen vacancy sites. Vecchietti et al. introduced Ga³⁺ to ceria supports to increase the concentration of surface oxygen vacancies. The catalytic activity of Pt/C₆Ce₆Ga₂₀, however, was lower than that of Pt/CeO₂. Ricote et al. added Zr⁴⁺ to the ceria support. Although the zirconia doping did not increase oxygen vacancies, it enhanced oxygen surface mobility and increased the reaction rate.

These seemingly conflicting results imply that some important features of the active site that determine the catalytic reactivity are still missing in our current understanding and demonstrate the need for further understanding of all parts of this complex interacting system (metal catalyst, support, adsorbates) at the atomic level—which is difficult to obtain without cumulative, multimodal, in-situ/operando modes of characterization. In this work, we describe the correlative results of such a study. Within this work, combined methods of ex situ and in situ characterization have elucidated the unique electronic structure and bonding environment of the active sites (perimeter Pt⁰ – O vacancy – Ce⁵⁺ sites) of Pt/CeO₂. The data demonstrate that strong dynamical forms of complexity exist in this system, features that convolve strongly mediating perturbations due to adsorbates and metal center (Pt and Ce) valence- and bonding-state speciations.

Results
Size evolution of Pt species under working conditions. An activity test (Fig. 1a) revealed the catalyst first starts to show activity at 180 °C. As the temperature is raised, the production of H₂ and CO₂ continues to increase. The catalyst is not active below 180 °C when the temperature ramp is reversed. To understand the effect of WGS reaction conditions upon the structure of the catalyst, the nature of the as-prepared and reacted (collected after the activity test) Pt/CeO₂ were evaluated using scanning transmission electron microscopy (STEM) and X-ray absorption spectroscopy (XAS). High-angle annular dark field (HAADF)-STEM images of the as-prepared and reacted catalyst are shown in Fig. 1b, c, respectively. The Pt species in the as-prepared catalyst are not easily identified due to their ultra-small size (likely single-atom, vide infra) and highly dispersed nature, as seen by the patches of diffuse contrast in Fig. 1b. We note here that XAS spectra of the as-prepared catalyst show very similar features to those expected for ceria-supported Pt(II) single atoms (Supplementary Fig. 1). Over the course of the reaction, the Pt species aggregate into larger Pt nanoclusters (about 1.7 nm in diameter, Supplementary Fig. 2) which are seen clearly in Fig. 1c. The XAS measurements that are discussed later in the manuscript agree quantitatively with the STEM particle size measurements. These results, together with those obtained from in situ diffuse reflectance infrared Fourier-Transform spectroscopy (DRIFTS) measurements (Supplementary Fig. 3) and XAS measurements (Supplementary Fig. 4), suggest that during the activating temperature ramp, single Pt atom species in the as-prepared sample coarsen to metallic Pt nanoparticles at T ≥ 180 °C. These metallic Pt nanoparticles remain the predominant species upon ramp-down to RT. When correlated with activity results, the data demonstrate that Pt nanoclusters likely exist within the working temperature range of the catalyst (180–300 °C). We hereafter focus on the attributes of atomic structure presented by Pt nanoclusters in situ within this temperature range.

Dynamics and heterogeneity of atoms in the active Pt species. First, we performed in situ TEM measurements to observe the atoms within active Pt nanoparticles. A typical Pt nanocluster is shown first in CO (Fig. 2a1 – a6, Supplementary Movie 1) and then in the WGS reaction environment (Fig. 2b1 – b6, Supplementary Movie 2). Due to the weakening of the Pt–Pt bonds from the strong repulsive interactions occurring between adsorbed CO molecules and as a result of the elevated temperature, the Pt atoms are generally quite dynamic under CO at 200 °C, except notably for those atoms deep in the nanocluster core and at/close to the interface between the nanocluster and the support. In contrast, under the WGS condition wherein water is introduced, almost all the outer Pt atoms become more stabilized/localized except for the ones located at the perimeter sites of the cluster (as shown by the yellow arrows in Fig. 2b3 – b5). At the perimeter sites, the Pt atomic columns are seen to disappear (Fig. 2b3) and reappear (Fig. 2b4) in sequential images, indicating that they remain dynamically mobile. These observations suggest that the atoms at different sites within the nanocluster dynamically and heterogeneously change under in situ conditions. The dynamics of the Pt atoms at the perimeter, in comparison to the relatively static Pt atoms at the terrace/step surface sites, demonstrate their uniqueness as specific metal centers, ones where chemical activities are likely occurring (vide infra).

In the present data, we further observe a distortion in the interplanar spacing (plane bending) that is present at the surface of the nanocluster, here visible in Fig. 2b2 and indicated with dashed yellow lines for clarity. The observed bending is an indication that the surface atoms are not in a bulk terminated configuration and that anisotropic/localized bond strains—ones subject to perturbations due to both adsorbates and specific support bonding interactions—are important features of their structure. We will now focus on the dynamic interactions of the perimeter sites with adsorbates and support atoms in order to better understand the observed phenomena.

In situ variation of adsorbate bonding at perimeter Pt sites. To provide additional insights into the bonding of adsorbates with specific Pt sites, DRIFTS measurements were performed first under CO and then WGS conditions. Under CO (Fig. 3a) at high
temperatures (300 °C, 240 °C, and 180 °C), infrared bands of high frequency (HF) and low frequency (LF) atop/bridge bound CO on Pt atoms are observed; at lower temperatures (120 °C and RT), the LF bands disappear. The LF bands are associated with CO bonded to low coordinated Pt atoms in proximal contact with reduced Ce$^{3+}$ sites along the metal–support interface (Pt$^{0}$–O vacancy–Ce$^{3+}$)\textsuperscript{18–24}. The appearance of the LF bands at high temperatures, therefore, suggests that a correlation exists between Pt atoms proximal to the catalyst perimeter and reduced ceria sites.

As shown in Fig. 3b–f, the introduction of H$_2$O does not affect the line shapes of these modes but does influence their intensities. At low temperatures (26 °C and 120 °C), an increase of the HF peak intensity, together with a redshift, signifies an interaction between water molecules, Pt, and CO via one of several possible mechanisms noted in prior literature: (i) partial charge transfer to the CO 2π* orbital due to the modification of the metal particle potential by the lone pair of a bound water molecule; or (ii) dipole-dipole interactions between co-adsorbed CO and water molecules\textsuperscript{25–30}. Interestingly, this trend is reversed at higher temperatures (at which the dynamic perimeter Pt$^{0}$–O vacancy–Ce$^{3+}$ sites appear). Upon further addition of H$_2$O to CO at 180 °C, 240 °C, and 300 °C, the intensities of the HF peaks decrease while those of the LF peaks increase, an indication of the migration of some CO adsorbates from high to low coordination perimeter Pt sites due to the change of gas condition from CO to WGS. Such changes seem stronger with the decrease of the temperature from 300 °C to 180 °C which could be due to the temperature effect on the CO coverage\textsuperscript{27,31}.

In summary of the TEM and DRIFTS observations, we conclude that the perimeter Pt$^{0}$–O vacancy–Ce$^{3+}$ site responds dynamically to the temperature and that CO (one of the two reactants) migrates to it at high temperature. We, therefore, hypothesize that it is the active site for the WGS reaction. To further investigate this hypothesis, we turn our attention to another reactant, H$_2$O.

Water dissociates into hydroxyl and atomic H species on the surface of ceria supports. The hydroxyl group bound to the
The surface of ceria is easily detected by DRIFTS measurements. Figure 4 shows difference spectra illustrating the changes that occur in the bonding of hydroxyl moieties that are pre-adsorbed on ceria in the presence of CO at 300, 200, 120, and 26 °C. At both 300 °C and 200 °C, the difference spectra show complex negative and positive features, indicating that a redistribution of –OH bonding states on the ceria occurs after the introduction of CO. The increase in the intensity of the type II-B OH band (at 3638 cm$^{-1}$) is an indication that the ceria support has been reduced, likely from surface oxygen elimination, and it suggests the association of OH bonds with surface reduced ceria sites at working temperatures. Previous reports also suggest that the active –OH groups are classified as type-II –OH groups formed on partially reduced ceria and reside within a narrow zone around the periphery of Pt–ceria interface.

Combining these results (Fig. 4) with the insights previously discussed (Fig. 3), we hypothesize that the two reactants of the WGS process (CO and H$_2$O) must associate in part with proximal contacts occurring at the perimeter Pt$^0$ – O vacancy – Ce$^{3+}$ sites at working temperatures. To verify this point, it is necessary to demonstrate that the reduced ceria sites that adsorb the hydroxyl groups are in fact located near the perimeter Pt sites. A precise quantification of their fractional densities can establish the existence of such a correlation between the number of perimeter Pt$^0$ sites and the number of oxygen vacancies/Ce$^{3+}$ on the ceria surface at working temperatures.
Active oxygen vacancies associate with perimeter Pt⁰ sites. To quantify the ratio of perimeter Pt⁰ /Ce³⁺, we carried out AP-XPS measurements of the Pt 4f and Ce 3d regions under WGS conditions (Fig. 5 and Supplementary Fig. 7). Though XPS reveals the effective valence state of Pt averaged over the entire nanocluster, positively charged Pt²⁺ atoms (Supplementary Fig. 8), are most likely located at the interface with the Ce oxide support 35. Therefore, changes in Pt²⁺ concentration indicates the nanocluster-support interface restructures when cooled. Generally, there is a higher fraction of Pt⁰ than Pt²⁺ at all temperatures. Starting from about 65% Pt⁰ and 35% Pt²⁺ at 300 °C, the fraction of Pt⁰ decreases linearly as the temperature decreases to 200 °C, at which point the nanocluster contains approximately 55% Pt⁰ and 45% Pt²⁺. This ratio is retained through further
cooling to RT. The numbers suggest that approximately 22% of interfacial Pt atoms are reduced to Pt$^0$ at 300 °C. Among the interfacial Pt atoms, those reducible atoms are most likely at the perimeter. They are the least coordinated and most affected by the support and adsorbates. Thus, approximately half of the perimeter Pt atoms are reduced to the zerovalent state. These high temperature Pt$^0$ perimeter sites strongly associate with reduced ceria sites as indicated by DRIFTS (the emergence of high-temperature perimeter Pt$^0$–O vacancy–Ce$^{3+}$ sites) and Ce 3d AP-XPS data. Under the WGS reaction, the concentration of Ce$^{3+}$ is 20% at 300 °C, which is 5% higher than that at RT. The rest of the Pt atoms at the interface remain in 2$^+$ state and interact with the ceria support, thereby preventing further small cluster agglomeration.

AP-XPS evaluation of Pt 4f and Ce 3d reveal the relative concentrations of Pt and Ce to be approximately, a 1:9 ratio (Supplementary Table 1). Within the 1.7 nm Pt nanocluster, ~10% of Pt atoms are perimeter Pt$^0$ site and therefore represent about 1% of all metal (Pt and Ce) ions. At 300 °C, the concentration of Ce$^{3+}$ is about 20%. Assuming all Ce$^{3+}$ stay at the surface of the support 36, the total fraction of Ce$^{3+}$ is 18%, corresponding to 9% oxygen vacancies. Analogous estimates for the RT species suggest 7% oxygen vacancies (Fig. 5b and Supplementary Table 1). Assuming they are retained throughout the cooling process, the additional 2% oxygen vacancies correlate with the appearance of perimeter Pt$^0$ sites. Since perimeter Pt$^0$ sites associate with oxygen vacancies, this implies the increased oxygen vacancy sites appear near the perimeter Pt$^0$ sites at working temperatures. This reinforces our hypothesis that perimeter Pt$^0$–O vacancy–Ce$^{3+}$ sites bring the two reactants closer.

**Fluxional bond dynamics of Pt nanoclusters.** To provide a better understanding of the structural dynamics of ceria supported Pt nanoclusters, in situ XAS measurements were performed at Pt L$_3$ edge. As shown in Fig. 6a, b, the XAS spectra collected at working temperatures (200 °C, 250 °C, and 300 °C) show changes compared to those collected at corresponding post-reaction RT, suggesting that the catalyst is dynamic and adopts different structures when it is in its active catalytic state. The perturbations seen in these data illustrate underpinnings related to both atomic and electronic structural origins. At working temperatures, the white line (near 11,566 eV) is consistently shifted to lower energies and becomes narrower compared to RT.

The decreased white line intensity and the redshift suggest a reduction of Pt occurs at elevated temperatures, in agreement with XPS results.

A quantitative analysis was performed to elucidate the structural changes that are evidenced in the R-space EXAFS data (Supplementary Fig. 9). As shown in Fig. 6a–g, the coordination numbers and the distances of Pt-O and Pt-Pt pairs change with temperature under WGS conditions. These changes in bonding are reversible and upon returning to RT revert to the initial resting state of the catalyst. At RT, the coordination number of the Pt-Pt pairs provided by the analysis is 8.3 ± 0.5 (Fig. 6f). This value corresponds closely to the expectations for an idealized, hemispherical cuboctahedral cluster with the size of approximately 1.7 nm$^3$. We also found the bond length of Pt-O to be about 2.6 Å (Fig. 6e), significantly longer than the 2.0 Å to 2.2 Å range expected for direct Pt-O bonding interactions 38. Long metal-oxygen distances (2.5 Å to 2.7 Å) have also been seen for supported Pt treated in H$_2$ at T < 350 °C as well as for clusters bound to partially-hydroxylated supports 39,40. The long bond in each case was ascribed to the presence of an additional atom (i.e. atomic chemisorbed H) residing at the interface between the metal atoms and the support 39,40. The adsorbed H atoms recombine into H$_2$ at working temperatures (Fig. 1a) and such recombination occurs with the appearance of active oxygen vacancies (near perimeter Pt$^0$ sites) which act as specific sites for H$_2$ activation 43.

As shown in Fig. 6e, at high temperatures, the bond distance of Pt-O is within the range from 2.53 Å to 2.57 Å, about 0.1 Å shorter than that at RTs (from 2.61 Å to 2.65 Å). Such change could be related to the reduced ceria support. The missing oxygen atoms in the ceria unit cell elongate the Ce$^{3+}$–O bond (~0.1 Å longer than that of Ce$^{4+}$–O) 43,44, which, via interface interaction, drives the structure modification of Pt clusters. In addition, a distortion in the lattice plane of Pt clusters was observed in electron microscopy results (Fig. 2b).

The catalytic activity at/near the perimeter Pt$^0$ sites. The data described above make it possible to develop a schematic depiction of the atomic structure of the catalyst at low and high temperatures (Fig. 7), which helps to rationalize structure-activity correlations developed for the WGS reaction over Pt/CeO$_2$ catalysts. At low temperatures (< 180 °C), the Pt nanoclusters and the ceria support are inactive. Herein the reactant CO adsorbs on the step/terrace Pt$^0$ sites. The reactant H$_2$O, via dipole-dipole interaction...
with CO, affects the CO-Pt bonding, and further likely dissociates into hydroxyl and atomic H: the hydroxyl groups bond to the surface of the ceria support, leaving atomic H species at the interface of the Pt nanocluster and the support. At elevated temperatures where the catalyst is active, the reduction of the ceria support and Pt nanoclusters produces new Pt\(^0\) – O vacancy – Ce\(^{3+}\) sites localized at the perimeter of the Pt cluster. These changes in valence states and vacancy concentration at the nanocluster-support interface produce significant associated strains in the metal-metal bonds of the Pt clusters. The distorted structure may facilitate the facile transport of CO molecules to the perimeter Pt\(^0\) surface-sites that comprise the active bonding ensemble for the catalytic reaction\(^{3,46}\).

Though the associative mechanism may also play a role (Supplementary Fig. 10), the perimeter Pt\(^0\) – O vacancy – Ce\(^{3+}\) sites are more likely associated with a regenerative redox mechanism. By using steady-state isotopic transient kinetic analysis (coupled with DRIFTS and mass spectrometry), Kalamaras et al. studied Pt/ceria catalysts of different sizes (1.3–8.0 nm). These results also suggest that at 300 °C, the WGS reaction proceeds largely via the redox mechanism (Pt\(^0\) – reactive site located along the periphery of the Pt–CeO\(_2\) interface (the specific rate based on the length of periphery of the Pt–ceria interface increases linearly with respect to particle size)\(^{34}\). In the redox mechanism pathway, the hydroxyl groups donate atomic oxygen to the reduced support, which is subsequently re-reduced by donating the atomic oxygen to CO\(^{34}\). The oxidation of ceria at the perimeter of the nanocluster would cause the corresponding oxidation of the perimeter Pt atoms and strong bonding between the perimeter Pt atoms and the support. Regenerative redox mechanism requires oxidation state changes are dynamically reversible (reversible change of Pt\(^0\) to Pt\(^2+\) and Ce\(^{3+}\) to Ce\(^{4+}\)). This is consistent with the dynamic restructuring of the perimeter atoms observed by TEM. Such repeating transitions in the electronic structure of the perimeter Pt atoms and their interaction with the support would markedly affect CO-Pt binding energy\(^{37}\) and increase the mobility of perimeter Pt atoms observed by TEM. Such repeating transitions in the electronic structure of the perimeter Pt atoms and their interaction with the support would markedly affect CO-Pt binding energy\(^{37}\) and increase the mobility of perimeter Pt atoms observed in situ TEM. The higher mobility of the perimeter atoms has also been observed in ceria supported Au nanoparticles under CO oxidation\(^{45}\), a reaction that most likely proceeds via a metal-assisted Mars–van Krevelen mechanism with oxygen vacancy exchange involved\(^{48,49}\). Here we note the importance of active oxygen vacancies. Oxygen vacancies also exist at low temperature but are not involved in the reaction. At high temperatures, only those newly emerged oxygen vacancies nearby the active Pt sites play important roles including adsorbing active hydroxyl groups, transporting oxygen atoms, and facilitating the recombination of H atoms which remain associated with sites at the nanocluster–support interface. The synergistic effects between O vacancy – Ce\(^{3+}\) and active perimeter Pt\(^0\) sites seem to best explain why only increasing the number of oxygen vacancies is not sufficient for improving the reaction rate\(^{12,14}\), and why not all perimeter Pt sites are able to be involved in the reaction\(^{50}\). Experimentally, to improve the reactivity of Pt/ceria catalysts for the WGS reaction, the dopants should be able to increase the concentration or improve the activity of oxygen vacancies that involved in the perimeter Pt\(^0\) – O vacancy – Ce\(^{3+}\) sites\(^{51}\).

In this work, we studied a system of ceria supported Pt for the WGS reaction by combining multiple in situ techniques including DRIFTS, AP-XPS, XAS, and ETEM. With the multiple sources of information on Pt nanocluster/ceria combined, we revealed the complexity of the catalysts under working conditions. The complexity involves the structural evolution of the catalysts, heterogeneity and dynamic behavior of atoms located at different places of the nanostructure. In this work, the functional species are found to be ceria supported Pt nanoclusters, in which multiple platinum states coexist. The WGS reaction proceeds most likely via the redox mechanistic path at the dynamic perimeter Pt\(^0\)-O vacancy-Ce\(^{3+}\) site at which CO oxidation, water reduction, and H recombination occurs. Accompanying the formation of the new perimeter Pt\(^0\)-O vacancy-Ce\(^{3+}\) sites are a bonding environment change occurring at the interface where H atoms are stored and a localized/substantive intraparticle distortion which facilitates the CO transport to the perimeter Pt\(^0\) sites.

**Methods**

**Sample preparation.** To prepare Pt/ceria sample, 0.5 g of cerium(IV) oxide (CeO\(_2\) nanopowder (<25 nm) was dispersed in a solution consisting of 2.0 g of Urea and 8 mL of water. While stirring, 3.3 g of a solution (approximately 1 wt% Pt) of chloroplatinic acid hydrate in water was added to the dispersion. All chemicals were purchased from Sigma-Aldrich. Ultrapure water (18.2 M\(\Omega\)m) was provided by a Millipore purification system. The mixture was sealed in a vial and stirred for 24h in an oil bath at 95 °C. Afterward, the postdeposition sample was separated and washed by 3 cycles of centrifuging/redispersing in water to remove the residual ions from the precursors. The sample was then dried overnight at 60 °C before being crushed into a powder and calcined at 500 °C (10 °C/min heating ramp). The resulting platinum loading of the sample was measured by Inductively Coupled Plasma (ICP) elemental analysis and determined to be 1.85 wt%. Note, certain commercial equipment, instruments, or materials are identified in this paper in order to specify the experimental procedure adequately. Such identification is not intended to imply recommendation or endorsement by the National Institute of Standards and Technology, nor is it intended to imply that the materials or equipment identified are necessarily the best available for the purpose.

**Activity test.** For the activity test, the as-prepared catalyst was loaded in a quartz tube (<2 mm ID, 2.4 mm O.D.) and mounted in a Clausen cell\(^{12}\). For the reaction, 1% CO (balanced with He; total flow rate of 25 ml/min) traveled through a water bubbler to carry water vapor (2.3%) to the sample. The activity test was executed by a stepwise increase of the reaction temperature from room temperature (RT) to 300 °C and returning to RT. The reactants and products were analyzed by a mass spectrometer (Hiden QGA).

**Electron microscopy.** Ex situ electron microscopy was carried out on a Hitachi 2700D scanning transmission electron microscope equipped with a probe aberration corrector operated at 200 kV. In situ environmental transmission electron microscopy (ETEM) experiments were conducted using an aberration-corrected FEI Titan ETEM operated at 300 kV. The third-order spherical aberration coefficient (C3) of the aberration corrector was tuned to a slightly negative value of approximately −15 μm, yielding a white-column contrast for the atomic columns in the resultant images. TEM samples were prepared by dispersing the Pt/CeO\(_2\) powder onto a windowed micro electro-mechanical system (MEMS)-based SiN\(_4\) chip, which was then loaded into a DESIOnSolutions Wildfire heating holder. After heating the sample into the ETEM, 24 Pt of CO gas was introduced into the environmental cell and the catalyst was reduced in situ at 300 °C for 2 hours. The catalyst was subsequently cooled to 200 °C and imaged under the same pressure of CO gas. To produce a water-gas shift reaction environment, at 200 °C, water vapor was directed to the cell with a H\(_2\)/CO gas ratio 1:1, resulting in a total cell pressure of 73 Pa. Images were acquired using a Gatan K2 IS direct electron detector operated at 20 frames per second with an electron flux of 9.6 × 10\(^{13}\) A\(\cdot\)cm\(^{-2}\) and a total exposure time of 6 s. The electron beam was blanked when images were not being acquired. Static high-resolution images were recorded from many areas of the catalyst under CO and then WGS conditions. The time-resolved images were temporally binned by summing to various time resolutions (e.g., 0.15 s, 1.5 s, and 6.0 s), producing so-called time-averaged images with an increased signal-to-noise ratio.

**Diffuse reflectance infrared Fourier-transform spectroscopy (DRIFTS).** The spectra were collected using a Nicolet 6700 FTIR spectrometer equipped with a rapid-scanning liquid nitrogen-cooled mercury cadmium telluride (MCT) detector and a transient infrared reaction cell (Hiden Analytical Products). Using this reactor, the catalytic behavior of the catalyst was also tested and found to be very similar to that obtained using the Clausen reactor (Supplementary Fig. 11). Two types of temperature-dependent DRIFTS measurements were performed. In Measurement I (Fig. 3), at each temperature, the sample was treated in the following order: He (the background spectrum was collected) → 1% CO/He → 1% CO/2.3% water vapor/He. In Measurement II (Fig. 4), to study the surface OH bonding (3800–3150 cm\(^{-1}\) at each temperature: 2.3% H\(_2\)/O\(_2\) → 1% CO/He mixture. For both measurements, the temperature was increased stepwise from RT to 300 °C and back to RT. The total gas flow rate was 25 ml/min under each condition, the spectra were continuously collected until no further changes were observed. Each spectrum (256 scans) was collected with a resolution of 4 cm\(^{-1}\).
Ambient pressure X-ray photoelectron spectroscopy (AP-XPS). A commercial SPECS AP-XPS instrument equipped with a PHOIBOS 150 EP MCD-9 analyzer at the Chemistry Division of Brookhaven National Laboratory (BNL) was used for XPS analysis (resolution: ~0.4 eV). A Mg K-alpha X-ray anode (1254 eV) was used for all measurements. Incident Mg K-alpha (~1254 eV) incident X-rays have a maximum escape depth of about 6.2 nm, deep enough to probe the entire Pt nanocluster. The Ce 3d photoemission line with the strongest Ce 3d feature (916.9 eV) was used for energy calibration. The powder sample (Pt/CeO2 catalyst) was pressed on to an aluminum plate and loaded into the AP-XPS chamber at ultra-high vacuum. For the reaction, 1.3 Pa of CO + 3.0 Pa H2O (CO:H2O = 1:2.3) was introduced into the reaction chamber through a high precision leak valve and O 1 s, Ce 3d, and Pt 4f regions were collected at different temperatures. The temperature was increased stepwise from RT to 300 °C and back to RT.

X-ray absorption spectroscopy (XAS). XAS measurements were performed at Beamline 20-ID-C of the Advanced Photon Source (APS), Argonne National Laboratory. The sample powders were loaded in a quartz tubing (2 mm I.D., 2.4 mm O.D.) and mounted in a Clausen cell. The Pt L2,3 XAS data were collected in fluorescence mode by using a Vortex (Hitachi) silicon drift detector. For the in situ measurements, 1% CO (balanced with He; the total flowrate of 25 ml/min) carrying water vapor (2.3%) flows through the catalyst. The measurements were performed under WGS reaction conditions through a sequence of heating and subsequent cooling to RT steps (RT → 100 °C → RT → 150 °C → RT → 200 °C → RT → 250 °C → RT → 300 °C → RT). Once the desired temperature was reached, after about 15 min, the XAS data were collected.

Data availability
The data supporting the findings of this study are available within the article and its Supplementary Information file. Additional information and figures are provided in the Supplementary Information file and Supplementary movies. Other data are available from the authors upon request.

Received: 14 September 2020; Accepted: 2 January 2021; Published online: 10 February 2021

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Acknowledgements

Y.L., A.I.F., and R.G.N. acknowledge support for this work by the U.S. Department of Energy, Office of Basic Energy Sciences under Grant No. DE-FG02-03ER15476. Reaction tests and DRIFTS measurements at Brookhaven National Laboratory’s Chemistry Division and XAFS data analysis were made possible due to the Program Development fund 21-017 to A.I.F. M.K. and R.G.N. acknowledge fellowship support from the Department of Chemistry at the University of Illinois. Work by S.D.S. and Z.L. at Brookhaven National Laboratory was supported by the US Department of Energy, Office of Basic Energy Sciences, Division of Chemical Sciences, Geosciences and Biosciences under contract no. DE-SC0012704. S.D.S. is partially supported by a US Department of Energy Early Career Award. This research used Hitachi2700C STEM of the Center for Functional Nanomaterials, which is a U.S. DOE Office of Science Facility, at Brookhaven National Laboratory under Contract No. DE-SC0012704. The XAFS experiments used resources of the Advanced Photon Source Beamline 20ID at Argonne National Laboratory, which is an Office of Science User Facility operated for the U.S. Department of Energy (DOE) Office of Science and was supported by the U.S. DOE under contract no. DE-AC02-06CH11357. I.L.V. and P.A.C. gratefully acknowledge NSF grant CBET-1604971 and the facilities at the National Institute of Standards and Technology at Gaithersburg, M.D. as well as those at the John M. Cowley Center for High Resolution Electron Microscopy at Arizona State University. W.D.Y. acknowledges support under the Cooperative Research Agreement between the University of Maryland and the National Institute of Standards and Technology Physical Measurement Laboratory, award 70NANB14H209, through the University of Maryland. We also gratefully acknowledge stimulating discussions on supported catalyst materials with Dr. Christopher Karwacki that helped sharpen presentations made in the current work.

Author contributions

Y.L., R.G.N., and A.I.F. conceived the project. Y.L., P.A.C., S.D.S., R.G.N., and A.I.F. designed the experiments. M.K. prepared the catalyst. I.L.V., L.Z., W.-C.D.Y., and P.A.C. were responsible for TEM experiments. Y.L. and M.K. tested the catalytic activity of the catalyst and performed DRIFTS measurements. Z.L. conducted AP-XPS experiments and analyzed AP-XPS data. Y.L., M.K., and J.H. conducted XAS experiments. Y.L. analyzed XAS data. Y.L., R.G.N., and A.I.F. co-wrote the manuscript with contributions from all authors. M.I.E. edited the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41467-021-21132-4.

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Peer review information Nature Communications thanks Sebastián E. Collins and the other, anonymous, reviewer(s) for their contribution to the peer review of this work. Peer reviewer reports are available.

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