An Atomic-Scale View of CO and H₂ Oxidation on a Pt/Fe₃O₄ Model Catalyst

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Abstract: Metal-support interactions are frequently invoked to explain the enhanced catalytic activity of metal nanoparticles dispersed over reducible metal oxide supports, yet the atomic-scale mechanisms are rarely known. In this report, scanning tunneling microscopy was used to study a Ptₓ/Fe₃O₄ model catalyst exposed to CO, H₂, O₂, and mixtures thereof at 550 K. CO extracts lattice oxygen atoms at the cluster perimeter to form CO₂, creating large holes in the metal oxide surface. H₂ and O₂ dissociate on the metal clusters and spill over onto the support. The former creates surface hydroxy groups, which react with the support, ultimately leading to the desorption of water, while oxygen atoms react with Fe from the bulk to create new Fe₃O₄(001) islands. The presence of the Pt is crucial because it catalyzes reactions that already occur on the bare iron oxide surface, but only at higher temperatures.

Variations in the CO oxidation activity of nominally similar metal nanoparticles dispersed over different metal oxides[1i] are a clear indicator of metal-support interactions, but the mechanisms of how the metal oxide gets involved are difficult to ascertain. The support is known to transfer electrons to (or from) the clusters,[2] modifying their shape and adsorption properties, and adsorb reactants and promoters (e.g., water) that can speed up the reaction.[3] At elevated temperature, the so-called Mars–van Krevelen (MvK) mechanism[4i] can operate, where CO molecules extract lattice oxygen atoms (O_{lattice}) at the cluster perimeter forming CO₂, and gas-phase O₂ repairs the surface to complete the catalytic cycle. Although there is mounting evidence that the MvK mechanism plays a role when reducible metal oxides are utilized as the support,[5] there is little atomic-scale information available to better understand how these processes occur.

Herein, we used scanning tunneling microscopy (STM) to follow the evolution of a Ptₓ/Fe₃O₄(001) model catalyst exposed to CO, O₂, and H₂ at 550 K. Holes and islands in the vicinity of Pt clusters provide direct evidence of the reduction and oxidation of the metal oxide in CO- and O₂-rich atmospheres, respectively, while dissociation and spillover of H₂ lead to hydroxylation of the support lattice. We interpret our results as the metal catalyzing reactions that otherwise occur only between the reactants and the support at higher temperatures.

An STM image of the Ptₓ/Fe₃O₄(001) model catalyst utilized as the basis for this work is shown in Figure 1. The Fe₃O₄(001) support[4i] exhibits large, flat terraces (Figure 1, inset) characterized by rows of protrusions in STM images related to surface Fe atoms. The rows rotate by 90° from terrace to terrace, which is a consequence of the spinel structure. Surface O atoms are not imaged because they have no density of states in the vicinity of the Fermi energy[5] Surface OH groups, formed through the reaction of water and oxygen vacancies during sample preparation,[6i] modify the density of states of the neighboring Fe atoms, making them

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**Figure 1.** The Ptₓ/Fe₃O₄(001) model catalyst. The Fe₃O₄(001) support exhibits rows of protrusions related to surface iron atoms. Elongated protrusions on the Fe rows are surface OH groups. Pt₁₄ clusters (large, oval protrusions) are formed by exposing 2.7 × 10⁻⁷ m⁻² Pt₁₄ adatoms to CO (1 × 10⁻⁶ mbar) for 10 min at room temperature. Upon CO adsorption, otherwise stationary Pt, adatoms become mobile and agglomerate. The Pt cluster sizes were assigned on the basis of STM movies in which the agglomeration was tracked atom-by-atom. Inset: On a large scale, the support exhibits large, flat terraces on which the metal particles are randomly distributed.
appear brighter in STM images.\[7\] The surface exhibits a $\sqrt{2} \times \sqrt{2}$R45° reconstruction,\[11\] which binds metal adatoms in one particular site per unit cell at temperatures as high as 700 K.\[8,9\] In this work, randomly distributed Pt clusters with sizes in the range of one to six atoms (Figure 1a) were prepared by CO-induced sintering of $2.7 \times 10^{17}$ Pt adatoms per m$^2$, as demonstrated previously for Pd.\[8,9\] X-ray photoelectron spectroscopy (XPS) measurements (see the Supporting Information, Figure S3) revealed that the CO remained on the clusters, but desorbed when the sample was heated to 550 K.

Exposing the Pt$_{1.6}$/Fe$_2$O$_4$ system to CO, O$_2$, and H$_2$ at room temperature resulted in no discernible difference in STM (Figure S2). However, exposure to the same gases at 550 K induced significant modification of the support morphology in the vicinity of the clusters. Following CO exposure, (Figure 2a, $p_{CO} = 1 \times 10^{-7}$ mbar, 60 min) large holes covering 3.7% of the surface were observed in the Fe$_2$O$_4$(001) terrace. The step height (2 Å) is consistent with the distance between the Fe$_{oct}$–O planes, and the rows (rotated by 90°) of the underlying Fe$_{oct}$–O layer are clearly visible inside the larger holes (inset). Each hole is associated with (at least) one Pt cluster, which typically sits at the step edge, but only half of the clusters are associated with holes. Holes were also created when the model catalyst was exposed to H$_2$ (Figure 2b, $p_{H_2} = 1 \times 10^{-7}$ mbar, 20 min), but now most clusters appeared inside the holes that cover 8.7% of the surface. As this occurred in just 20 min, the process is clearly more efficient with H$_2$. Some redispersion results in the observation of isolated Pt adatoms on the surface.

None of the morphological changes described above occur in the absence of Pt (see Figure S2), but CO and H$_2$ reduce iron oxides at higher temperatures, producing CO$_2$ and water, respectively.\[5\] In the case of CO, the Pt cluster efficiently adsorbs the molecule and delivers it to the cluster–support interface where it extracts O$_{lattice}$ atoms to create CO$_2$. As some clusters are associated with the removal of many tens of O$_{lattice}$ atoms, while others remain on the pristine substrate, extraction of the first O$_{lattice}$ atom most likely presents a significant energetic barrier. This is in line with recent DFT + U calculations for CeO$_2$(111), where undercoordinated O atoms at step edges were found to be more easily removed by CO than terrace oxygen atoms.\[10\] As the clusters are mostly found on the upper step edge of large holes, they most likely diffuse along the step once a hole is nucleated, extracting more and more lattice oxygen atoms as they go. An interesting question remains: Why are holes observed instead of isolated point defects? The answer is that undercoordinated Fe atoms created by the surface reduction diffuse into the bulk, forming interstitials at elevated temperature, as observed previously.\[6,11\] Oxygen vacancies are not common defects in iron oxides, which handle stoichiometry variation via the cation sublattice.\[11\]

With H$_2$ dissociation on the clusters followed by spillover creates surface OH groups, which are known to react with O$_{lattice}$ atoms, leading to the desorption of water above 500 K.\[6,12\] As OH groups can diffuse over the metal oxide surface at 550 K,\[6,12\] O$_{lattice}$ extraction can occur away from the cluster. In this way, once a hole has nucleated, the step edge recedes from the cluster with time, leaving the cluster in

\[\text{Figure 2. STM images acquired following exposure of the Pt/Fe}_2\text{O}_4(001) \text{ model catalyst to } 1 \times 10^{-7} \text{ mbar of CO (a), H}_2 \text{ (b), or O}_2 \text{ (c) at 550 K. a) Following exposure to CO (60 min), approximately 50% of the Pt clusters sit at the edge of or inside monolayer holes in the Fe}_2\text{O}_4(001) \text{ terrace. The rows (rotated by 90°) of the underlying Fe}_\text{oct}–\text{O layer can be seen inside the holes (inset). The illustration shows how CO extracts O}_{lattice} \text{ atoms from the cluster perimeter and how CO}_2 \text{ desorbs from the surface. Undercoordinated Fe atoms diffuse into the Fe}_2\text{O}_4 \text{ bulk. b) Following exposure to H}_2 \text{ (20 min), the clusters reside inside monolayer holes in the Fe}_2\text{O}_4(001) \text{ terrace. Pt adatoms are observed at regular terrace sites, and the coverage of surface OH groups is increased. The illustration shows how H}_2 \text{ dissociates on the Pt cluster, spils over onto the support, and reacts with O}_{lattice} \text{ atoms, finally leading to the desorption of molecular water. c) Following exposure to O}_2 \text{ (20 min), all Pt clusters reside on top of islands with a step height of 2 Å. On the larger islands, the rows (rotated by 90°) of the next Fe}_2\text{O}_4(001) \text{ layer are clearly visible. The islands result from the spillover of oxygen atoms onto the Fe}_2\text{O}_4 \text{ support, which react with Fe that diffuses out from the bulk.} ]

the bottom of the hole. Such an autocatalytic reduction of iron oxide by hydroxy groups has been reported previously.\[16\]

Interestingly, as the clusters appear to be trapped inside the holes that they create, pretreating the Pt/Fe$_3$O$_4$(001) system in H$_2$ at 550 K could be an effective strategy to mitigate thermal sintering during CO oxidation.

When the Pt$_{15}$/Fe$_3$O$_4$ model catalyst was exposed to O$_2$ at 550 K (Figure 2c; p$_{O_2}$ = 1 × 10$^{-7}$ mbar, 20 min), the Pt clusters appeared atop small islands with a step height of 2 Å. The characteristic rows of Fe$_3$O$_4$(001) are clearly visible on the larger islands, suggesting that the clusters catalyze the growth of a new stoichiometric Fe$_3$O$_4$(001) surface. A significant reduction in the density of clusters suggests that gas-assisted sintering occurs in the O$_2$ atmosphere. The growth of new Fe$_3$O$_4$(001) layers has been observed on the clean surface above 900 K.\[14\] In the present study, however, the Pt clusters catalyze the rate-limiting step, O$_2$ dissociation, and highly reactive atomic O species spill over onto the support where they react with Fe supplied from the bulk at 550 K. Thus, as well as providing a sink for excess Fe during reduction, the Fe$_3$O$_4$ bulk also supplies Fe for the growth of new Fe$_3$O$_4$(001) surfaces during reoxidation.\[14\] Similar metal-activated regrowth leads to encapsulation of the metal clusters and a loss of the catalytic activity of Pt and Pd clusters on TiO$_2$ surfaces.\[15\] But our low-energy ion scattering (LEIS) measurements (Figure 5) suggest that the Pt clusters are not encapsulated by the Fe$_3$O$_4$(001) islands. Zhang et al. recently observed the encapsulation of larger Pt clusters to occur above 650 K in ultrahigh vacuum.\[16\]

The data in Figure 2 clearly show that the etching and regrowth of the Fe$_3$O$_4$(001) support are catalyzed by Pt clusters. To test whether similar processes still occur when multiple gases are present, we heated the Pt/Fe$_3$O$_4$ model catalyst in mixtures of CO, O$_2$, and H$_2$ at 550 K. In a 1:1 mixture of CO and O$_2$, the changes to the support morphology are significantly smaller than with the individual reactants (Figure S5). This could be due to two effects: Either both O$_{\text{ads}}$ extraction and island growth occur at the same time, and their effects on the morphology cancel out, or the presence of one gas blocks the activation of the other (poisoning). To investigate this, we exposed the sample to CO/O$_2$ mixtures of various compositions, and found that small holes began to appear for CO/O$_2$ ratios of 50:1, whereas for the formation of small islands, an O$_2$ excess of 1.5 was required. These findings suggest that the etching and regrowth processes still occur when both CO and O$_2$ are present in the gas phase, although we cannot exclude that direct reaction processes also take place under such conditions. Finally, exposure of the system to a 1:1 mixture of CO and H$_2$ for 20 min resulted in holes that extended several layers deep into the support (Figure S5). Such a roughening of the surface suggests that the addition of H$_2$ to the gas mixture should help activate Pt clusters for the CO oxidation reaction.

In conclusion, the holes/islands that form in CO/O$_2$ mixtures provide clear evidence of the reduction and oxidation of the Fe$_3$O$_4$ support being catalyzed by Pt clusters. The legacy of these reactions is strikingly clear in STM images because the Fe$_3$O$_4$ bulk provides a sink for Fe atoms, which readily diffuse to and from the surface at the reaction temperature. In general, our data provide an atomic-scale view into metal–support interactions of importance in heterogeneous catalysis.

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