

Dual role of CO in the stability of subnano Pt clusters at the Fe₃O₄(001) surface

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Interactions between catalytically active metal particles and reactant gases depend strongly on the particle size, particularly in the subnanometer regime where the addition of just one atom can induce substantial changes in stability, morphology, and reactivity. Here, time-lapse scanning tunneling microscopy (STM) and density functional theory (DFT)-based calculations are used to study how CO exposure affects the stability of Pt adatoms and subnano clusters at the Fe₃O₄(001) surface, a model CO oxidation catalyst. The results reveal that CO plays a dual role: first, it induces mobility among otherwise stable Pt adatoms through the formation of Pt carbonyls (Pt₁–CO), leading to agglomeration into subnano clusters. Second, the presence of the CO stabilizes the smallest clusters against decay at room temperature, significantly modifying the growth kinetics. At elevated temperatures, CO desorption results in a partial redispersion and recovery of the Pt adatom phase.

single-atom catalysis | adsorbate induced sintering | redispersion | Fe_3O_4 | nucleation and growth

Subnanometer metal particles exhibit a range of interesting electronic or catalytic properties that can vary substantially with the removal or addition of a single atom (1-6). Understanding the mechanistic details underlying the rearrangement of the active phase is important because changes in cluster size and shape are known to be commonplace under the conditions used in heterogeneous catalysis (7, 8), and because such processes are associated with deactivation phenomena such as sintering. Although sintering is usually regarded as a thermally activated process, there is increasing evidence that adsorbates influence sintering rates in a reactive environment by formation of mobile metal-molecule intermediates (2, 8–30). Indeed, in a previous study we demonstrated that the formation of highly mobile Pd1-CO species led to enhanced sintering in the Pd/Fe₃O₄(001) system (31). Here, we turn our attention to Pt. Mobility is induced in the form of Pt₁-CO. In addition, CO stabilizes the smallest clusters. When it desorbs, Pt dimers break up into single atoms; thus, the CO is necessary for preserving nuclei that act as seeds for further growth. Using roomtemperature scanning tunneling microscopy (STM), complemented by X-ray photoelectron spectroscopy (XPS) and density functional theory with an on-site Hubbard U (DFT+U), we follow the COinduced diffusion and coalescence of Pt atom-by-atom, creating catalytically active (32) subnano clusters with a well-defined size distribution. On heating, desorption of CO leads to significant redispersion of Pt into the adatom phase.

Results

Fe₃O₄(001) is one of the most stable facets of magnetite (Fe₃O₄), a spinel-type ferrimagnetic conductive oxide (33) frequently used as a catalyst support. The surface exhibits a ($\sqrt{2} \times \sqrt{2}$)R45° reconstruction based on an ordered array of subsurface Fe vacancies and interstitials (34), which stabilizes metal adatoms [e.g., Au, Ag, and Pd (31, 35, 36)] in one specific site per unit cell against thermal sintering to temperatures as high as 700 K. Fig. 1A shows an STM image of the Fe₃O₄(001) surface following deposition of 0.2 ML Pt (1 ML = 1 adatom per reconstructed unit cell = 1.42×10^{14} atoms per cm²). The characteristic undulating rows of the reconstructed $Fe_3O_4(001)$ substrate are indicated by dark-blue lines (34). There are no O-related states in the vicinity of the Fermi energy $(E_{\rm F})$; thus oxygen atoms, although present in the surface layer, are not imaged with the scanning conditions used here. Surface hydroxyl groups (O_sH), a common adsorbate at this surface after ultrahigh vacuum (UHV) preparation, are observed as a pair of extra-bright Fe atoms (cyan arrow) (37, 38). The Pt adatoms adsorb in two distinct configurations. First, protrusions located midway between the surface Fe rows indicate Pt twofold-coordinated to oxygen (Pt₁, Fig. 1B), the configuration commonly observed for other metal adatoms at this surface (31, 35, 36, 39, 40). DFT+U calculations find an adsorption energy $\Delta E_{ads}(Pt_1)$ of 3.89 eV compared with free Pt atoms in vacuum and little charge transfer to the surface (<0.5 e⁻). A second configuration, labeled Pt1*, not previously observed for other metals, appears offset to one side in STM images. Our DFT+ U calculations find a stable adsorption site $[\Delta E_{ads}(Pt_1^*) = 3.01 \text{ eV},$ charge transfer $<0.3 \text{ e}^{-}$] bridging two oxygen atoms parallel to the rows of surface Fe (Fig. 1C). The Pt atom is observed to switch between these two configurations in STM image sequences collected in UHV conditions at room temperature (Fig. S1). The transition from Pt₁* to the Pt₁ state is strongly preferred, consistent with the higher binding energy of Pt1 obtained in DFT+U calculations. Considering the large energy difference between the two adsorption sites, the high density of Pt1* can only be explained by a large diffusion barrier between the sites. In DFT+U calculations, the energy barrier along an optimized diffusion path is found to be of the order of $\Delta E_{\text{barr}}(\text{Pt}_1^* \rightarrow \text{Pt}_1) \ge 0.9 \text{ eV}$. Assuming an Arrheniustype behavior, this energy results in a slow, thermally activated

Significance

The catalytic activity of metal particles is highly size-dependent in the subnanometer regime, which makes understanding how and why particle sizes change in reactive atmospheres particularly important. Here, we show that carbon monoxide plays a dual role in the coarsening of otherwise highly stable Pt atoms on an Fe₃O₄(001) support: CO adsorption weakens the adatomsupport interaction inducing mobility, and stabilizes the Pt dimer against decay into two adatoms. Our results illustrate how molecules modify the clustering dynamics on surfaces, provide much-needed insight into how deactivation and redispersion can occur in single-atom catalyst systems, and demonstrate an approach to prepare size-distinguished clusters for studies of the size effect.

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Fig. 1. (*A*) STM image of 0.2 ML Pt/Fe₃O₄ (001). The surface Fe_{oct} atoms are imaged as rows along the [110] direction (illustrated by the parallel blue lines). The bright features on the rows of Fe (example highlighted by cyan arrow) are due to surface OH groups (37, 38). The red and yellow frames highlight Pt adatoms in two different geometries: bridging surface O atoms in between (Pt₁) and parallel to (Pt₁*) the Fe_{oct} rows. (*B* and C) DFT+U-based force-relaxed structural models of the two adsorption configurations.

process at room temperature (of the order of $10^{-3} \cdot s^{-1}$ for an excitation frequency of $10^{13} s^{-1}$). Annealing to 280 °C in UHV reduces the coverage of Pt₁* drastically, allowing the conclusion that the transition is thermally induced and the remaining Pt₁* features are likely stabilized by surface defects.

In Fig. 2, we show that CO adsorption on Pt₁ adatoms presents a clear signature in STM that can be interpreted with the aid of DFT+U calculations. Fig. 2 *A*–*D* shows a sequence of STM images acquired while exposing 0.15 ML Pt adatoms to 2×10^{-10} mbar CO. Fig. 2*A*, acquired before CO exposure ($P < 10^{-10}$ mbar), contains several Pt₁ and Pt₁* adatoms and a bright, double-lobed feature in the center of the scanning area. Increasing the CO pressure to 2×10^{-10} mbar (Fig. 2*B*) results in a second double-lobed feature at the former position of a Pt₁ adatom (labeled Pt₁–CO), and a new, single-lobed feature at the former position of a Pt₁* adatom (labeled Pt₁*–CO). Between Fig. 2 *B* and *D*, transitions between the Pt_1^* -CO and Pt_1 -CO state occur. Likely, both entities are chemically identical, i.e., they consist of one Pt atom and one CO molecule. The only difference between these species would be their adsorption geometry. To check whether adsorption of another molecule from the residual gas could be responsible for the observed features, we also exposed the as-deposited Pt adatoms to the most common components of the residual gas as measured by a quadrupole mass spectrometer: H₂, CO₂, CO, CH₄, H₂O, and O₂. (The presence of trace amounts of O₂ in our vacuum system is linked to the use of this gas during sample preparation.) No bond formation or other interactions with the Pt adatoms were observed for any gas molecule other than CO at room temperature.

DFT+U calculations for CO adsorption on Pt₁ adatoms are consistent with strong Pt–CO binding. The configuration of Pt₁– CO shown in Fig. 2*E* is favored by 5.53 eV compared with a free



Fig. 2. CO adsorption and Pt site interchange. (*A*–*D*) STM image sequence (same place) acquired during exposure to 2×10^{-10} mbar CO. (*A*) Pt adatoms in two configurations (Pt₁, Pt₁*) and bright, double-lobed features are observed. (*B*) CO adsorption on Pt₁ and Pt₁* adatoms results in bright, double-lobed and oval features, respectively. (*C*) The Pt₁*–CO labeled in red in *B* changes site, transforming into a Pt₁–CO feature. (*D*) The Pt₁–CO formed and labeled in cyan in *B* transforms into a Pt₁ *–CO feature. (*E*) Structural model of Pt₁–CO/Fe₃O₄(001). Upon adsorption of CO, the Pt adatom is lifted up and shifts perpendicular to the Fe rows to an off-centered position. (*F*) Simulated STM images at +2-V sample bias: Pt adatom, Pt* adatom, Pt₁–CO, mirrored Pt₁–CO. The latter is an overlay image of Pt₁–CO in the two equivalent off-centered positions to simulate switching during the scan, induced thermally or by the STM tip.

Pt atom and a free CO molecule in vacuum. The Pt₁ adatom is lifted up by ~ 1.3 Å away from the surface plane, and shifted perpendicular to the surface Fe-row direction. In this asymmetric geometry, one Pt-O bond is stretched to ~2.98 Å, which compares to ~ 2.03 Å for the second Pt–O bond and ~ 1.98 Å for both Pt–O bonds before CO adsorption. In STM images at 78 K (Fig. S2) and in simulated STM images (Fig. 2F) the Pt₁ carbonyl is imaged as a single, bright, oval protrusion, offset from the center of the Pt adsorption site. This resembles one of the two lobes observed in experiment. When this feature is scanned with a sample bias of +2 V (Fig. S2C), the protrusion switches repeatedly from one side of the unit cell to the other, suggesting that the STM tip can move the Pt-CO between two nearby sites that are energetically and symmetrically equivalent. Similar tipinduced switching behavior has been observed for dangling bond pairs on hydrogenated semiconductor surfaces (41, 42). To estimate the switching barrier we calculated the DFT energy of the likely transition state, a Pt1-CO species centered between the Fe_{oct} rows. We found this energy to be only ~ 0.25 eV higher than the minima, suggesting that the double-lobed feature observed at room temperature is a time average of thermally activated switching. The Pt1* adatom is also lifted up from the surface (by ~ 0.7 Å) by CO adsorption, but remains in the same location laterally. We considered the possibility that the double-lobed feature could be a Pt_1 -(CO)₂ species. However, we never observed a different type of CO adsorption event, which would lead to Pt₁-CO in this case, although it should be clearly visible in STM. Moreover, the observed switching between the Pt_1^* -CO and Pt1-CO states agrees well with the energies derived from the DFT+U calculations.

Fig. 3 shows STM images of Pt/Fe₃O₄(001) in a constant CO pressure of 2×10^{-10} mbar, selected from a 55-frame movie acquired over approximately 3 h. The CO induces mobility, and Pt clusters are formed over time. The sizes of the individual clusters are determined by tracking each individual Pt atom. Fig. 3A shows a large-area image of the surface after 10 min of CO exposure. The yellow frame highlights a smaller region selected for detailed analysis in Fig. 3 B-G. In Fig. 3B, four Pt adatoms are observed, two in Pt1 configuration and two in Pt1* configuration. Going through the panels from left to right we find that CO adsorbs initially on one Pt_1 and one Pt_1^* (Fig. 3C), and that the Pt₁-CO feature then coalesces with the remaining Pt₁* adatom to form a Pt dimer in Fig. 3D. The Pt₁*-CO subsequently joins the dimer to form a trimer in Fig. 3E. Finally, the remaining Pt₁ adatom adsorbs a CO molecule in Fig. 3F, and coalesces with the trimer to form a Pt tetramer in Fig. 3G. It should be emphasized that, because of tip convolution effects, the identification of cluster sizes by their appearance is very hard if not impossible with STM (43). With our approach, the STM signature of the different cluster sizes can be ascertained, which allows determining the exact number of Pt atoms contained within each cluster in the larger scanned area (Fig. 3H). In Fig. 3 the largest clusters contain four atoms. The cluster size distribution in areas scanned with the STM during the CO exposure is comparable to those in areas located further away. Thus, the Pt₁-CO mobility and sintering is not induced by the STM tip.

It is important to note that the Pt carbonyl is the only diffusing species observed in the STM movies. Dimers, trimers, and larger clusters are immobile, even in CO pressures up to 10^{-6} mbar. Fig. 4A shows an STM image of 0.15 ML Pt/Fe₃O₄(001) acquired as part of a different STM movie starting from adatoms where the CO pressure was increased up to 10^{-6} mbar in a stepwise manner. As in Fig. 3, we determine the number of Pt atoms in each cluster by watching its assembly. The resulting size distribution is narrow, ranging from one to five atoms. The majority of clusters contain two or three atoms. The bar graph illustrates the number density of Pt clusters (blue, 71.5 ± 4.7% of all features, 5.69 × 10^{16} clusters per m²) versus the adatom phase (yellow,



Fig. 3. CO-induced Pt diffusion and coalescence. STM image sequence acquired during exposure to 2×10^{-10} mbar CO. (A) After 5 h in UHV and 10 min CO exposure: Pt adatoms coexist with a few double-lobed, bright features, identified as Pt₁-CO created by CO adsorption in Fig. 2. (*B*-*G*) Formation of a Pt tetramer: CO adsorption on Pt₁ and Pt₁*, mobility, and coalescence are observed atom by atom. (*H*) Partially sintered state: Only a few adatoms remain; the biggest clusters in this image contain four atoms.

 2.27×10^{16} adatoms per m²). The graph represents the average from two experiments in which 0.15 ML Pt adatoms were sintered in different amounts of CO. In both cases the remaining adatom density was between 0.01 and 0.02 ML. The shaded area corresponds to the respective standard deviation.

To investigate the stability of the subnano clusters formed by COinduced sintering at reaction temperatures, we heated the surface to 520 K in UHV. The average coverage of Pt adatoms increased by almost 130%, and the number of clusters decreased by more than 60%. The remaining clusters were larger, with an average size of ~7.6 atoms (~3.3 atoms before annealing). This trend is again illustrated by the bar graph beneath the STM image in Fig. 4*B*, which clearly shows that adatoms are the dominant Pt-related feature after annealing (71.2 ± 8.9%). The complete absence of small clusters suggests that these species are unstable without CO, and that CO desorption drives the breakup. In a control experiment, the



Fig. 4. Sintering at high CO pressure and redispersion of small clusters by annealing. The bar graphs at the bottom show the average adatom and cluster density in each case. (A) $Pt/Fe_3O_4(001)$ after exposure to CO pressures up to 10^{-6} mbar. The main features are small clusters (Pt_2 , Pt_3); the largest observed clusters consist of five atoms. (B) A different region of the sample after annealing to 520 K in UHV. Interestingly, more adatoms are now present at the surface. The number of clusters has decreased and the average cluster size is considerably larger.

adatoms were heated in UHV without prior exposure to CO. This treatment leads to a small decrease in the adatom density, but no discernible change in the cluster density. This suggests that the adatoms become mobile at 520 K and can incorporate within existing clusters. Crucially, the constant cluster density highlights the importance of CO for new cluster nucleation. The crucial role of CO for the stability of the clusters is supported by the XPS measurements in Fig. 5, which show that CO was adsorbed on the clusters following room-temperature exposure, but desorbs after heating to 520 K. Before CO exposure (blue curves) the Pt $4f_{7/2}$ peak at 71.4 eV resembles that of metallic Pt, consistent with the very small electron transfer predicted by DFT+U. Following 120 L CO exposure (red) a peak appears in the C 1s region at 287.7 eV, which is higher than CO adsorbed in an atop site on Pt(111) (286.8 eV). The adsorption of the CO causes the Pt 4f to shift up in binding energy to 72.4 eV, which is also higher than when CO is adsorbed atop on Pt(111). Finally, heating above the desorption temperature of CO on Pt (~500 K), the peak in C1s is gone and the Pt 4f peak shifts back to the metallic position. Note that CO does not adsorb on the pristine $Fe_3O_4(001)$ surface at room temperature, where all experiments were performed.

The assertion that Pt cluster stability is linked to adsorbed CO is supported by DFT+U calculations. In the absence of CO, two Pt₁ isolated adatoms are preferred to a Pt dimer (two adatoms: 7.78 eV, one Pt dimer: 6.35 eV with respect to Pt in the gas phase). A Pt₂(CO)₁ species is also unfavorable with respect to one Pt₁ carbonyl and one adatom ($E[Pt_2(CO)] = 8.44 \text{ eV}$ and $E[Pt_1 + Pt_1CO] = 3.89 \text{ eV} + 5.53 \text{ eV} = 9.42 \text{ eV}$). Crucially, a Pt dimer with two adsorbed CO molecules is slightly more stable than two Pt₁-CO species (11.16 eV compared with 11.06 eV). Thus, the formation of dimers can only proceed in the presence of (at least) two adsorbed CO molecules. When CO is desorbed, the strong Pt₁/Fe₃O₄ interaction leads to a redispersion of Pt, and a partial recovery of the adatom phase.

Discussion

The agglomeration phenomena described here are a clear case of gas-induced coarsening, a well-known process that causes



Fig. 5. XPS spectra of the Pt 4*f* and C 1*s* regions acquired after the deposition of 0.7 ML Pt (blue), after CO exposure (20 min, 10^{-7} torr, black), and after annealing to 520 K (red). After deposition Pt adatoms are the main species, coexisting with a small number of clusters and Pt carbonyls formed by adsorption from the background gas. After CO exposure all Pt adatoms have adsorbed CO and coalesced to small clusters. Annealing to 520 K causes the CO to desorb. The Pt peak shifts back to an energy slightly lower than its initial position, indicating the complete absence of CO and the coexistence of adatoms and clusters.

catalyst deactivation. Nevertheless, unequivocal identification of the mobile species is rare, and through our combined use of STM, XPS, and DFT, we clearly demonstrate that a Pt_1 -CO species drives the coalescence of single Pt atoms into clusters. Moreover, we demonstrate this occurs because the adsorption of the CO molecule weakens the Pt–O bonds, facilitating diffusion. Adsorption of one CO (which, as argued above, appears as a double-lobed feature in STM) is sufficient to allow the Pt to move across the surface.

In recent years knowledge of the size effect in subnano catalysis has been revolutionized by the ability to deposit size-selected clusters on model support surfaces (6, 44–47). Our approach to monitor cluster formation in an atom-by-atom fashion produces a model system where the cluster size and distribution is accurately known. Following the behavior of such "size-distinguished" ensembles in reactive atmospheres could provide valuable insights into catalytic reactions in the nonscalable cluster regime. Indeed, we recently demonstrated that subnano Pt clusters grown by the CO-induced sintering approach are active for CO and H_2 oxidation via a Mars–van Krevelen-type mechanism (32).

Surprisingly, we find that the CO also stabilizes dimers against decay. Thus, in addition to providing mobility, CO plays a crucial role in the growth kinetics by providing stability to the smallest clusters that act as further nucleation sites. In the UHV environment, the binding of the metal adatoms to the substrate is sufficiently strong that spontaneous formation of a Pt dimer is energetically unfavorable. Similar behavior was observed previously for Ag on the same surface, and extremely high adatom coverages were achieved because the nucleation of stable clusters was suppressed (35, 36). Here, this limitation is bypassed when two Pt₁–CO species meet, because their merger produces a $Pt_2(CO)_2$ dimer that is stable and can function as a seed for further growth. When the sample is heated, the CO desorbs, and the small clusters break up into single Pt atoms.

The results described here contain several important lessons for the rapidly emerging, yet controversial field of single-atom catalysis. Here, the goal is to use single atoms as active sites for catalytic reactions to maximize the efficient use of noble metals such as Pt. Sintering clearly runs counter to this objective, and the ability to rapidly recover the adatom phase is considered vital to economic viability. In our work, the strong CO-Pt1 interaction is clearly detrimental as it accelerates the sintering process. The problem of the strong CO-Pt₁ bond has been recognized before, and has been proposed to poison low-temperature oxidation reactions on a variety of substrates (48). Based on our results, it seems plausible that the catalytic activity reported for single-atom catalysts based on Pt could result from subnano particles formed under reaction conditions, which subsequently redisperse after the fact. In this light, although Pt is clearly an excellent catalyst as a nanoparticle, a less reactive (cheaper) metal would almost certainly yield a better and more stable single-atom catalyst. It would

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be fascinating indeed if, in an attempt to reduce the amount of Pt, it becomes possible to circumvent its use entirely.

In summary, we have analyzed the interaction of CO with Pt adatoms and clusters in detail. CO-induced mobility leads to the agglomeration of Pt into subnano clusters, and the presence of the CO stabilizes the smallest clusters against decay. Our results provide insights into coarsening processes at surfaces, which should be applicable to established catalysts based on oxide-supported metal clusters. CO-induced adatom sintering provides a welldefined initial state for experiments into the size-dependent properties of small Pt clusters.

Experimental Methods

The STM experiments were performed in a two-vessel UHV system. It consists of a preparation chamber and an analysis chamber. The latter is equipped with an Omicron µ-STM, which was operated in constant current mode. The substrate material was a synthetic Fe₃O₄(001) crystal grown using the floating zone method (49). The crystal was prepared by cycles of sputtering with Ar^+ ions (E_{Ar} = 1 keV, $j_{\text{sample}} = 6.5 \,\mu\text{A/cm}^2$, 10 min) followed by annealing at 870 K in 6.6 \times 10⁻⁷ mbar O₂ (15 min). Temperatures were measured with a K-type thermocouple attached to the sample holder. The systematic error of the measurement is estimated as ± 20 K (at 520 K). At high temperatures, the uncertainty of the temperature readout increases (\pm 50 K at 900 K) because the thermocouple is not attached directly to the sample. The XPS data were acquired in a second UHV chamber equipped with a He-flow cryostat, a twin-anode X-ray source (Al, Ag), a Focus 500 monochromator, and a SPECS Phoibos 150 analyzer. In this system, the sample is prepared by cycles of sputtering with Ne⁺ ions (E_{Ne} = 1 keV, $j_{sample} \sim 2.3 \,\mu\text{A/cm}^2$) and annealing in O₂ using a directional doser creating a local pressure of $P = 10^{-6}$ mbar at the sample. In both chambers Pt was deposited using a modified Omicron single-pocket electron-beam evaporator. The deposition rate (~0.3 ML/min, 1 ML = 1 atom per reconstructed unit cell = 1.42 \times 10¹⁸ m⁻²) was calibrated using a water-cooled quartz-crystal microbalance.

The DFT calculations are based on the augmented plane wave + local orbital method as implemented in Wien2K (50). We used the generalized gradient approximation with a Hubbard U (U_{eff} = 3.8 eV) to treat the strongly correlated Fe 3d electrons. The adsorption of Pt adatoms was modeled on a 17-layer 2 × 2 supercell of the $\sqrt{2}$ × $\sqrt{2}$ -reconstructed Fe₃O₄(001) surface with inversion symmetry and a vacuum layer of \sim 13 Å. In total, the slab contains 244 atoms plus 2 or 4 Pt atoms and 2 or 4 C and O atoms in the calculations including CO molecules. The atomic sphere sizes used for Fe, O, and Pt were 0.98, 0.79, and 1.11 Å, respectively. A plane-wave cutoff of $RK_{max} = 7.0$ corresponding to 296.6 eV was used, and the Brillouin zones of the surface models were sampled with a $3 - \times 3 - \times 1 - k$ mesh. The surface models were relaxed until all forces were below 1 mRy/bohr (26 meV/Å). A Fermi broadening of 0.08 eV was used. The charge transfer from Pt adatoms to the slab was estimated from the change in the partial charge distribution in the Pt atom and the change in the magnetic moment of the total slab, the Pt adatom, and the adjacent O atoms.

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