Enhanced CO Oxidation Rates at the Interface of Mesoporous Oxides and Pt Nanoparticles

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Supporting Information

ABSTRACT: The interaction of the metal and support in oxide-supported transition-metal catalysts has been proven to have extremely favorable effects on catalytic performance. Herein, mesoporous Co₃O₄, NiO, MnO₂, Fe₂O₃, and CeO₂ were synthesized and utilized in CO oxidation reactions to compare the catalytic activities before and after loading of 2.5 nm Pt nanoparticles. Turnover frequencies (TOFs) of pure mesoporous oxides were 0.0002−0.015 s⁻¹, while mesoporous silica was catalytically inactive in CO oxidation. When Pt nanoparticles were loaded onto the oxides, the TOFs of the Pt/metal oxide systems (0.1−500 s⁻¹) were orders of magnitude greater than those of the pure oxides or the silica-supported Pt nanoparticles. The catalytic activities of various Pt/oxide systems were further influenced by varying the ratio of CO and O₂ in the reactant gas feed, which provided insight into the mechanism of the observed support effect. In situ characterization using near-edge X-ray absorption fine structure (NEXAFS) and ambient-pressure X-ray photoelectron spectroscopy (APXPS) under catalytically relevant reaction conditions demonstrated a strong correlation between the oxidation state of the oxide support and the catalytic activity at the oxide−metal interface. Through catalytic activity measurements and in situ X-ray spectroscopic probes, CoO, Mn₃O₄, and CeO₂ have been identified as the active surface phases of the oxide at the interface with Pt nanoparticles.

1. INTRODUCTION

The chemistry of heterogeneous catalysis on transition-metal surfaces involves the covalent bonding of neutral molecules, which leads to dissociation and rearrangements to produce other neutral atoms and molecules.¹⁻² Many previous studies have demonstrated large enhancements of catalytic behavior due to the role of the oxide−metal interface, wherein charge plays a crucial role in the catalytic chemistry.³⁻¹³ Examples include the hydrogenation of CO₂ on rhodium by many different transition-metal oxides, the use of titanium oxide as a catalyst support for CO hydrogenation (Fischer−Tropsch) reactions, and even tandem catalysis.⁵⁻¹³ Despite the large amounts of research on oxide−metal interfaces, the mechanism that causes the enhanced catalytic properties is not fully understood. This is mainly due to a limitation of insight of the interfaces under catalytic reaction conditions, as opposed to pre- and postcatalysis characterizations. Through the use of synchrotron-based characterization techniques it is possible to study the surface chemistry of many catalytic systems on the molecular level in order to find vital mechanistic insights under the catalytic working conditions.¹⁴

Herein, numerous novel mesoporous oxides (Co₃O₄, NiO, MnO₂, Fe₂O₃, and CeO₂) were synthesized and loaded with size-controlled Pt nanoparticles to investigate the oxide−metal interface effects on catalytic CO oxidation reactions in both excess O₂ and excess CO. While pure mesoporous oxides are poor catalysts compared with pure Pt nanoparticles, all of the mesoporous oxide/Pt catalysts exhibit large enhancements of the CO oxidation rate beyond the turnover rates of pure Pt nanoparticles and pure mesoporous oxides. These systems were extensively characterized, both ex situ and in situ, in order to provide insight into the working conditions of the catalyst systems. In situ studies showed that the redox behavior of the oxides provides the charge that changes the mechanism of CO oxidation from covalent-bond chemistry to so-called acid−base or charge-transfer chemistry. This change of mechanism was found to be a general phenomenon for all of the oxide−metal interfaces studied. Through the utilization of near-edge X-ray absorption fine structure (NEXAFS) and ambient-pressure X-ray photoelectron spectroscopy (APXPS) under catalytically relevant reaction conditions, we were able to demonstrate a strong correlation between the oxidation state of the oxide support and the catalytic activity at the oxide−metal interface.

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2. EXPERIMENTAL SECTION

2.1. Synthesis of Mesoporous Co₃O₄, NiO, MnO₂, Fe₂O₃, and CeO₂. Mesoporous silica KIT-6 with a bicontinuous pore structure was used as a hard template. For the synthesis of KIT-6, 27 g of P123 and 43.5 mL of concentrated HCl were dissolved with 980 mL of water in a polypropylene bottle, and 33.3 mL of n-butanol was added to the solution at 308 K with vigorous stirring. After 1 h, 58 g of TEOS was added to the solution followed by stirring at this temperature for 24 h. The capped bottle was stored at 313 K for another 24 h in an oven. The solid was filtered, dried at 363 K overnight, and calcined at 823 K for 6 h.

Mesoporous oxides were prepared through the hard-templating approach using KIT-6, which was developed by Schüth and co-workers. Co(NO₃)₂·6H₂O, Ni(NO₃)₂·6H₂O, Mn(NO₃)₂·xH₂O, Fe(NO₃)₃·9H₂O, and Ce(NO₃)₃·6H₂O (Sigma-Aldrich) were used to synthesize mesoporous Co₃O₄, NiO, MnO₂, Fe₂O₃, and CeO₂, respectively. In a typical synthesis, 16 mmol of metal nitrate dissolved in 8 mL of water was added to 4 g of KIT-6 in 50 mL of toluene at 338 K with stirring. After evaporation of toluene, the precipitated solid was collected and dried at 333 K overnight, followed by calcination at 573 K for 6 h. The silica template was removed completely using 2 M aqueous NaOH solution heated to 333 K overnight. The solid was diluted in 8 mL of water, followed by several washing steps with water and a final drying step at 333 K.

2.2. Preparation of Oxide-Supported Pt Nanoparticle Catalysts. Poly(vinylpyrrolidone) (PVP)-capped Pt nanoparticles with an average diameter of 2.5 nm were synthesized by following the reported methods. For the preparation of oxide-supported nanoparticle catalysts, colloidal solutions of Pt nanoparticles (1 mg mL⁻¹) were diluted in ethanol. The desired amounts of solution (0.3–0.5 wt % Pt) were added to the mesoporous oxides, and the mixtures were sonicated for 3 h at room temperature using a commercial ultrasonic cleaner (Branson, 1510R-MT, 70 W, 42 kHz). The brown precipitates were separated by centrifugation (3000 rpm, 20 min), thoroughly washed with acetone and ethanol four times, and dried in an oven at 353 K overnight.

2.3. Characterization. Structural characterizations of Pt nanoparticles were performed using a Hitachi H-7650 transmission electron microscope operated at 120 kV. X-ray diffraction (XRD) patterns were measured on a Bruker D8 GADDS diffractometer using Co Kα radiation (1.79 Å). Nitrogen physisorption data were obtained on a Quantachrome Autosorb-1 analyzer. Elemental analyses by inductively coupled plasma atomic emission spectroscopy (ICP-OES) were conducted using a PerkinElmer optical emission spectrometer (Optima 7000 DV). Before an ICP-OES measurement, calibration was conducted using platinum standard (Fluka, TraceCERT, 1000 mg L⁻¹) as a certified reference material (CRM). Catalysts dissolved in aqua regia were diluted with DI water and filtered for the measurement.

2.4. Catalytic CO Oxidation. CO oxidation was performed in a laboratory-scale flow reactor operated at atmospheric pressure over the temperature range between 373 and 573 K. Gas flows (Praxair, UHP) were regulated using calibrated mass-flow controllers. Temperature was controlled using a type-K thermocouple and a PID controller (Watlow 96E). Catalyst (0.01–0.1 g) was diluted with low-surface-area quartz sand (washed with acid, rinsed with DI water, and calcined before use) and loaded into quartz reactors. Two different CO oxidation rates were acquired, in 10 Torr CO and 40 Torr O₂ with a balance of He for net reducing conditions or in 40 Torr CO and 100 Torr O₂ with a balance of He for net oxidizing conditions. Gas compositions were analyzed with a thermal conductivity detector (TCD) on an HP 5890 Series II gas chromatograph (GC). For comparison, mesoporous silica SBA-15 was used with and without loading of Pt nanoparticles for this reaction.

Figure 1. Preparation of Pt-nanoparticle-loaded Co₃O₄ catalysts. (a, b) TEM images of (a) the mesoporous silica template and (b) the resulting Co₃O₄ replica. (c) TEM image of Pt/Co₃O₄ catalysts and (d) their corresponding energy-dispersive spectroscopy (EDS) phase mapping, showing the merged image of the Co K (red) and Pt L (green) lines. (e) High-resolution TEM image of Pt/Co₃O₄ catalysts. (f) Illustration of the hard-templating (nanocasting) approach for the preparation of mesoporous-oxide-supported Pt nanoparticle catalysts.

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molecular production rates measured per unit surface site of the oxides based on their BET surface areas (m² g⁻¹) and crystalline structures. The TOFs of the Pt–oxide interfaces (i.e., after subtraction of the contributions from pure Pt and metal oxide) were calculated from the number of surface Pt sites.

2.5. In Situ Characterizations. APXPS and NEXAFS experiments were performed at beamlines 9.3.2 and 7.0.1 of the Advanced Light Source (ALS) at Lawrence Berkeley National Laboratory. NEXAFS studies were carried out in our purpose-built flow cell. Supported catalysts were exposed to pure CO or O₂ or a CO/O₂ mixture (either 39 Torr O₂ and 15 Torr CO or 39 Torr CO and 15 Torr O₂) at 473 or 523 K. The NEXAFS total electron yield (TEY) spectra with 0.3 eV resolution at the metal L (Co and Mn) or M (Ce) edge were monitored via a current amplifier by detecting compensating electrons from ground to the sample. The oxidation states of the metal oxides in the near-surface regions were determined by a linear-combination fitting of the reference compounds. APXPS at beamline 9.3.2 of the ALS was conducted because it overcomes the limitations of short mean free paths of emitted photoelectrons by bringing a small aperture cone, which is differentially pumped, very close to the sample surface, which is in the reaction gas mixture.

3. RESULTS AND DISCUSSION

3.1. Preparation of Mesoporous Oxides of Co₃O₄, β-MnO₂, NiO, α-Fe₂O₃, and CeO₂ and Pt-Loaded Oxide Catalysts. Mesoporous Co₃O₄, NiO, MnO₂, Fe₂O₃, and CeO₂ were prepared through the hard-templating (nanocasting) approach using mesoporous silica templates. The hard template determines the final structure of the oxide and provides stability during high-temperature crystallization.

In the present study, KIT-6 mesoporous silica was used as a hard template with an ordered bicontinuous mesostructure with cubic Ia₃d symmetry. When metal nitrates as metal oxide precursors were combined with KIT-6 in solution, they were completely impregnated into the silica templates and readily converted to the desired crystalline oxides after calcination at 573 K. The Kit-6 was removed completely by washing with aqueous NaOH solution, leaving mesoporous Co₃O₄, β-MnO₂, NiO, α-Fe₂O₃, and CeO₂ replicas. Figure 1 shows a general nanocasting approach for the preparation of ordered mesoporous oxides and representative transmission electron microscopy (TEM) images of the KIT-6 silica template, mesoporous Co₃O₄, and Pt-nanoparticle-loaded Co₃O₄ catalysts. TEM images of the mesoporous oxides demonstrated that the well-ordered structures and the wall thicknesses (ca. 3 nm) of the oxides were in accordance with the replicated KIT-6 (see Figures S1–S3 and Table S1 in the Supporting Information). PVP-capped Pt nanoparticles with an average diameter of 2.5 nm were incorporated into the mesoporous oxides for the preparation of Pt-nanoparticle-loaded mesoporous oxide catalysts. Sonication induced efficient dispersion of the Pt nanoparticles over the inner pores of the mesoporous oxides.

The TEM image in Figure 1c clearly shows that the Pt nanoparticles were deposited and well-dispersed on mesoporous Co₃O₄ homogenously. Elemental analyses by energy-dispersive spectroscopy (EDS) phase mappings on Pt/Co₃O₄ (Figure 1d) were in agreement with the values obtained by ICP-OES (i.e., 0.38 wt % Pt).
3.2. CO Oxidation on Pure Mesoporous Oxides.

Mesoporous oxides have shown notable catalytic activity as heterogeneous catalysts even in the absence of noble metals. Ren et al. studied ordered mesoporous oxides of Co₃O₄, Cr₂O₃, CuO, Fe₂O₃, MnO₂, Mn₂O₃, NiO, and CeO₂ as catalysts for CO oxidation. In their study, Co₃O₄, β-MnO₂, and NiO exhibited high CO oxidation activities, which were determined by measuring the temperatures required for 50% CO conversion \( T_{50} \). In order to study the catalytic activity of the metal-support interaction and the effect of gas feed composition, CO oxidation was carried out under two different sets of conditions, with either reducing (100 Torr CO and 40 Torr O₂ with a balance of He) or oxidizing (40 Torr CO and 100 Torr O₂ with a balance of He) gas feeds. The reaction was performed in a laboratory-scale flow reactor operated at atmospheric pressure over the temperature range between 373 and 573 K. It has been reported that Co₃O₄ shows extraordinarily high low-temperature activity, and Co³⁺ is believed to be an active site in CO oxidation. Jansson et al. proposed that CO oxidation over Co₃O₄ proceeds through a redox cycle in which gas-phase CO adsorbs on a cobalt site and subsequently reacts with a lattice oxygen atom. This then forms CO₂(g) and an oxygen vacancy, thus reducing the oxidation state of the cobalt site to Co²⁺. Reoxidation of the cobalt site occurs with gas-phase oxygen. As shown in Figure 2a, pure Co₃O₄ and NiO showed higher TOFs (normalized to surface area) than MnO₂, Fe₂O₃, and CeO₂ under both reducing and oxidizing conditions. The TOFs of MnO₂, Fe₂O₃, and CeO₂ under the net reducing reaction conditions were much higher than those under the net oxidizing reaction conditions. On the contrary, the catalytic activity of NiO was found to be higher under O₂-rich conditions than under CO-rich conditions. Intrinsic properties of the oxides seem to determine the catalytic behavior under these oxidizing or reducing reaction conditions.

3.3. CO Oxidation Studies on a Pt-Nanoparticle-Loaded Silica Support.

In order to find the contribution to the catalytic activity of only Pt nanoparticles, mesoporous silica was used as an inert support. It has been shown that mesoporous silica, such as SBA-15 or MCF-17, does not contribute to the catalytic activity of Pt for many reactions. When pure SBA-15 was solely used for CO oxidation in the current study, no conversion was found up to 673 K. In Figure 2a, the TOF for the Pt/SiO₂ catalyst was substantially higher under the O₂-rich reaction conditions than under the CO-rich reaction conditions across the temperature range studied. The TOFs for CO oxidation over Pt/SiO₂ were 0.014 and 0.11 s⁻¹ at 473 K under CO- and O₂-rich reaction conditions, respectively. Goodman and co-workers reported TOF values of 0.01−0.1 s⁻¹ at 473 K and an activation energy of ca. 26 kcal/mol over Pt/SiO₂ catalysts and Pt(100) single crystals under CO-rich reaction conditions, which are in agreement with the values for the Pt/SiO₂ catalyst from this work (0.014−0.11 s⁻¹ and 23 kcal/mol). CO oxidation on Pt-group metal surfaces has been shown to be structure-insensitive under CO-dominant conditions, under which the reaction rate is limited by CO adsorption because CO blocks the active sites for O₂ adsorption and dissociation. This agrees with our findings on CO oxidation over Pt nanoparticles supported on SiO₂, which exhibited no size dependence of the TOFs over the size range from 1.5 to 5 nm under both CO- and O₂-rich reaction conditions.

3.4. CO Oxidation Studies of Pt-Nanoparticle-Loaded Mesoporous Oxide Systems.

When Pt nanoparticles were...
loaded into the mesoporous oxides, the observed catalytic TOFs were significantly higher than those expected simply by summing the contributions from the oxide support and the pure Pt nanoparticles. This enhancement can be attributed to activity at the oxide–metal interface coupled with the activities of the oxide and Pt (Figure 2b). The TOFs over the Pt-loaded oxide catalysts were 495.27, 1.12, 0.57, 0.53, and 0.11 Pt $^{-1}$ s $^{-1}$ at 473 K under CO-rich reaction conditions and 443.31, 1.14, 0.60, 0.48, and 0.15 Pt $^{-1}$ s $^{-1}$ at 473 K under O$_2$-rich reaction conditions for Pt/Co$_3$O$_4$, Pt/NiO, Pt/CeO$_2$, Pt/MnO$_2$, and Pt/Fe$_3$O$_4$, respectively. The contributions of the oxide–metal interfaces were found to be orders of magnitude greater than those of Pt and the oxides regardless of the gas composition (Figure 2c,d). When Pt nanoparticles were loaded on the mesoporous oxides, the reaction rates were further altered by the redox behaviors of the oxide supports. Since the 3d-block metal oxides are subject to alternation of their oxidation states under redox gas atmospheres, synergistic acid–base and/or redox sites are regarded as the key for the catalytic oxidation reaction.36–38

3.5. In Situ Characterization of Oxide Catalysts during CO Oxidation. In order to understand the observed behaviors of activity enhancement, in situ characterizations using NEXAFS and APXPS were conducted under catalytically relevant reaction conditions. Briefly, supported Pt/Co$_3$O$_4$, Pt/MnO$_2$, and Pt/CeO$_2$ catalysts were exposed to pure CO or O$_2$ or a CO/O$_2$ mixture [~39 Torr O$_2$ and 15 Torr CO (denoted as O$_2$-rich) or 39 Torr CO and 15 Torr O$_2$ (denoted as CO-rich)] at 473 or 523 K. Figure 3a shows NEXAFS TEY spectra of the Pt/MnO$_2$ catalyst at 523 K under the O$_2$-rich and CO-rich reaction conditions. A linear-combination fitting of the reference compounds indicated that an excess of the MnO$_2$ phase (a fraction of 0.8) was present in equilibrium with the spinel Mn$_3$O$_4$ phase under the O$_2$-rich conditions. Under the CO-rich reaction conditions, however, the surface regions were further reduced, resulting in a 50:50 mixture of the MnO$_2$ and Mn$_3$O$_4$ phases (Figure 3). Figure 3b shows the proportions of MnO$_2$ on the surface and TOFs that have been decomposed into the contributions of pure Pt (in gray) and the Pt–CeO$_2$ interface (in orange). The contribution of pure mesoporous CeO$_2$ was too small to be represented. It should be noted that both the line plots and bar graphs are given to the same scale.

Figure 4. (a) NEXAFS TEY spectra at the Ce M edges for the Pt/CeO$_2$ catalyst and (b) graphs correlating the catalytic activity (TOF) and oxidation state of Ce in CO oxidation. Shown in (a) are representative linear-combination fittings obtained at 523 K under (left) 15 Torr CO and 39 Torr O$_2$ (O$_2$-rich) and (right) 39 Torr CO and 15 Torr O$_2$ (CO-rich) conditions. The top panels in (b) show the proportions of Ce$^{4+}$ obtained by linear-combination fitting of the reference compounds to the NEXAFS spectra and analysis of Ce 4d XPS spectra, and the corresponding TOF plots at both 473 and 523 K under different reaction conditions are given in the bottom panels. The bar graphs showing the total TOFs of the Pt/CeO$_2$ catalyst have been decomposed into the contributions of pure Pt (in gray) and the Pt–CeO$_2$ interface (in orange). The contribution of pure mesoporous CeO$_2$ was too small to be represented. It should be noted that both the line plots and bar graphs are given to the same scale. 

3.5. In Situ Characterization of Oxide Catalysts during CO Oxidation. In order to understand the observed behaviors of activity enhancement, in situ characterizations using NEXAFS and APXPS were conducted under catalytically relevant reaction conditions. Briefly, supported Pt/Co$_3$O$_4$, Pt/MnO$_2$, and Pt/CeO$_2$ catalysts were exposed to pure CO or O$_2$ or a CO/O$_2$ mixture [~39 Torr O$_2$ and 15 Torr CO (denoted as O$_2$-rich) or 39 Torr CO and 15 Torr O$_2$ (denoted as CO-rich)] at 473 or 523 K. Figure 3a shows NEXAFS TEY spectra of the Pt/MnO$_2$ catalyst at 523 K under the O$_2$-rich and CO-rich reaction conditions. A linear-combination fitting of the reference compounds indicated that an excess of the MnO$_2$ phase (a fraction of 0.8) was present in equilibrium with the spinel Mn$_3$O$_4$ phase under the O$_2$-rich conditions. Under the CO-rich reaction conditions, however, the surface regions were further reduced, resulting in a 50:50 mixture of the MnO$_2$ and Mn$_3$O$_4$ phases (Figure 3). Figure 3b shows the proportions of MnO$_2$ on the surface and TOFs that have been decomposed into metal–oxide interface and Pt metal contributions during the CO/O$_2$ reactions at 473 and 523 K. We found a strong correlation between the oxidation state of Mn and the normalized catalytic activity at the metal–oxide interface. In the O$_2$-rich reaction, the fraction of MnO$_2$ was changed very
little (from 0.81 at 473 K to 0.79 at 523 K). In parallel with this, the turnover rate at the metal−oxide interface (color-coded in orange) was slightly increased from 0.37 to 0.77 s\(^{-1}\), exhibiting a marginal change in CO oxidation rate at the interface compared with pure Pt (color-coded in gray) (Figure 3b, left). However, in the CO-rich reaction, both the extent of the reduction of Mn and the relative change in turnover rates at the metal−oxide interface were substantial. The fraction of MnO\(_2\) phase decreased from 0.72 to 0.49, while the turnover rate at the interface jumped from 0.51 to 2.16 s\(^{-1}\), which was an order of magnitude change with respect to pure Pt. On the basis of the observed trend, spinel Mn\(_3\)O\(_4\) was identified as the dominant phase at the metal−support interface and thus the species responsible for the orders of magnitude enhancement in the CO oxidation rate.

Both a bifunctional mechanism\(^{44}\) and a Mars−van Kravelen mechanism\(^{36−38}\) could potentially explain the catalytic activity enhancement by the support in this case; however, neither could give insight into the detailed reaction scheme of the CO-rich or CO-deficit atmospheres. The O 1s APXPS spectra of the Pt/MnO\(_2\) catalyst were obtained under CO-rich and O2-rich CO/O2 reaction conditions at 473 K. In these spectra, the peak at 529 eV was assigned to the lattice oxygen of MnO\(_2\)\(^{45,46}\) and the higher-binding-energy peak at 531 eV is due to OH species or adsorbed H\(_2\)O\(^{46}\) (Figure S8 in the Supporting Information). The normalized XPS intensities indicate that the lattice oxygen was more abundant under the O2-rich reaction conditions than under the CO-rich conditions, although the CO/O2 reaction was more favorable under the CO-rich conditions. From these results, we concluded that the interfacial reaction is governed by the oxidation state of MnO\(_2\).

This phenomenon was not limited to the case of the Pt/MnO\(_2\) catalyst but could be generalized to the other Pt/metal oxide systems. The Pt/CeO\(_2\) catalyst exhibited greater overall interfacial enhancement under the O\(_2\)-rich conditions, while it was more dramatic relative to pure Pt under the CO-rich reaction condition. APXPS and NEXAFS independently revealed that the near-surface region (ca. 2 nm) of the Pt/CeO\(_2\) catalyst mainly was composed of Ce\(^{4+}\) during the CO/O2 reaction at 523 K. On the other hand, the Ce\(^{3+}\) concentration was appreciable (ca. 15%) under CO-rich conditions but was negligibly small (less than 5%) under O\(_2\)-rich conditions at 523 K, suggesting that CO reduces CeO\(_2\) without apparent reaction turnovers (Figure 4). The O 1s APXPS spectra also indicated no correlation between lattice oxygen (or OH) and the reaction rate (Figure S9 in the Supporting Information). The overlapping conclusion was that Ce\(^{4+}\) sites give rise to the interfacial enhancement over the Pt/CeO\(_2\) catalyst during the CO/O2 reaction: Ce\(^{3+}\) sites formed upon reduction of CeO\(_2\) under the CO-rich conditions, rendering the overall enhancement (i.e., higher turnover rates under O\(_2\)-rich vs CO-rich conditions) without significantly impacting the enhancement factor (i.e., similar orders of magnitude enhancement at the interface relative to pure Pt).

For the Pt/Co\(_3\)O\(_4\) catalyst, the characteristic NEXAFS spectra of spinel Co\(_3\)O\(_4\) at the Co L edge were mainly maintained under CO-rich reaction conditions at and below 473 K, whereas CoO formed as the dominant phase at the expense of the spinel Co\(_3\)O\(_4\) phase above 523 K under the CO-rich conditions (Figure S5). Our linear-combination fitting indicated the simultaneous formation of two CoO phases: one with Co(II) in octahedral sites and another with Co(II) in tetrahedral sites.\(^{20}\) For the reaction in the O\(_2\)-rich environment,
the spinel Co$_3$O$_4$ phase remained dominant over the whole temperature range studied, in agreement with the findings of Oku and Sato.\textsuperscript{47} From the viewpoint of catalysis, the CO oxidation rates were enhanced by 4 orders of magnitude at the metal–oxide interface under both the CO- and O$_2$-rich conditions; however, the enhancement factor was greater in the case of the CO-rich reactions, where CoO phases were dominant at the near-surface regions (Figure 5). While Co(III) sites were usually attributed to the CO/O$_2$ reaction turnovers on pure Co$_3$O$_4$\textsuperscript{48} we found that Co(II) sites are responsible for the orders of magnitude enhancement in the reaction kinetics at the metal–oxide interface.

Through catalytic activity measurements and in situ surface probe experiments, we identified that Mn$^{2+}$ and Mn$^{3+}$ in spinel Mn$_3$O$_4$, Ce$^{4+}$ in CeO$_2$, and Co$^{3+}$ in Co$_3$O$_4$ are the active surface phases of the oxides in contact with Pt nanoparticles. The mesoporous oxides of 3d transition metals (Co and Mn) appeared to be more active in CO oxidation at a reduced form of the oxide in contact with Pt, while CeO$_2$ exhibited the opposite behavior, as Ce$^{4+}$ was catalytically more active than Ce$^{3+}$ when interfaced with Pt. This reflects on the fact that the catalytic TOF on the Pt/CeO$_2$ catalyst under the O$_2$-rich reaction conditions was significantly larger than that under the CO-rich reaction conditions.

On the basis of our measurements of oxidation state and catalytic activities, we propose the existence of a charge transfer during the chemical process. For example, the rate of oxygen desorption isotherms of metal oxides (S1 and S2) and mesoporous oxides (S1 and S5) in CO and H$_2$O$_2$ reactions using catalytic nanodiodes constructed from ca. 4 nm thick Pt films deposited on titanium oxide to form Schottky barriers.\textsuperscript{49} The current flow across the rectifying barrier was linearly correlated with the reaction turnover rate, indicating the major role of electron flow at the Pt–TiO$_2$ interface in the catalytic reactions. However, the turnover rates were 10$^4$ times greater than the electron flow rates measured at steady state, indicating that the electrons have several competing reaction paths (such as absorption at defect sites, heating by the thermoelectric effect, etc.) in addition to becoming part of the reaction path that enhances the catalytic turnover for some reason that is yet to be determined. It should be mentioned that cobalt oxide is an active material in studies of water splitting using solar energy, probably for similar reasons of efficient charge transfer during the chemical process.

4. CONCLUSIONS

At the interfaces of Pt nanoparticles and mesoporous oxides, we have observed great enhancements of catalytic activity in CO oxidation, which has rendered the “structure-insensitive” reaction on pure Pt surfaces very much “structure-sensitive” at oxide metal interfaces. Compared with pure Pt nanoparticles and pure mesoporous oxides, the turnover rates were increased by orders of magnitude with catalysis by the Pt–oxide interface. The reaction rates over Pt-nanoparticle-loaded oxide catalysts were further controlled by the redox properties of the oxides at the oxide–metal interface under reducing reaction conditions. Even under O$_2$-deficient reaction conditions, active oxygen could be provided from the oxide lattice. The redox chemistry in the near-surface regions of the oxide catalysts was dynamic, depending on the temperature and whether CO- or O$_2$-rich atmospheres were utilized during the reaction. It appears that the redox behaviors of the oxides provide the charges that participate at the oxide–metal interface, which act to amplify the CO oxidation. Through in situ characterizations using NEXAFS and APXPS under alternating redox conditions combined with catalytic activity measurements, we found that CoO, Mn$_3$O$_4$ and CeO$_2$ are responsible for the orders of magnitude enhancement in CO oxidation rate as the active surface phases of the oxide at the interface with Pt nanoparticles. This discovery of the relationship between the surface redox chemistry and catalytic activity offers great advancement in the knowledge of how the oxide–metal interface functions during catalytic reactions. The in-depth understanding of these phenomena will have wide implications in understanding catalytic selectivity on charged and polar oxide surfaces for a number of heterogeneous transformations.

ASSOCIATED CONTENT

S Supporting Information

TEM, XRD, and nitrogen adsorption—desorption isotherms of mesoporous oxides (Figures S1–S3 and Table S1); Pt size distribution histograms of the supported catalysts (Figures S4 and S5); conversions, Arrhenius plots, TOFs, and activation energies for CO oxidation (Figures S6 and S7 and Table S2); and APXPS spectra of Pt/oxide catalysts (Figures S8 and S9). This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes

The authors declare no competing financial interest.

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