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Tailored oxide interface creates dense Pt single-atom catalysts with high catalytic activity

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Highly reactive dense Pt single-atoms stabilized on an oxide support can resolve a grand challenge in the economic use of Pt in catalysis. Maximized number density of the reaction sites provided by dense Pt single-atoms guarantees the improved catalytic performance of Pt combined with the high efficiency. By manipulating the chemical nature of multi-component interfaces, we synthesized CO-tolerant dense Pt single-atoms highly reactive for the CO oxidation reaction, which governs the key steps for chemical energy conversion and emission control. The addition of 1 wt.% of Ce to TiO₂ support particles creates a CeO_x-TiO₂ interface that stabilizes Pt single-atoms by strong electronic interaction. Dense Pt single-atoms formed on CeO_x/TiO₂ oxides exhibit 15.1 times greater specific mass activity toward CO oxidation at 140 °C compared with a bare Pt/TiO₂ catalyst. We elaborate how the CeO_x-TiO₂ interfaces activate the interface-mediated Mars-van Krevelen mechanism of CO oxidation and protect Pt single-atoms from CO-poisoning. Through a comprehensive interpretation of formation and activation of dense Pt single-atoms using operando X-ray absorption spectroscopy, density functional theory calculation, and experimental catalyst performance test, we provide a key that enables the catalytic performance of noble metal single-atom catalysts to be optimized by atomic-scale tuning of the metal-support interface.

Broader context

Oxide supported Pt nanoparticles (NP)-based catalysts accelerate many critical chemical reactions for energy conversion and emission control. Achieving high catalytic performance of Pt catalysts, while using less amount of Pt, is key for sustainable use of earth unabundant Pt. Relevant catalysis communities are responding to such needs by reducing the size of Pt NPs to subnanometer or even pulverizing Pt NPs into single-atoms (SAs). Dense Pt-SAs homogeneously distributed over the surface of oxide supports guarantee the optimum materials' efficiency by using all individual Pt atoms as reaction sites. However, even for simple CO oxidation reaction, achieving the high catalytic performance of Pt-SAs is challenging because Pt-SAs are structurally vulnerable and Pt-SAs are easily CO-poisonable. In this study, we accomplished the catalytic supremacy of Pt-SAs using CeO_x/TiO₂ hybridoxide support and manipulating the chemical nature of multi-component interfaces. Highly reactive and CO-tolerant dense Pt-SAs formed at the CeO_x-TiO₂ interfaces exhibit outstanding catalytic performance toward CO oxidation compared with conventional Pt NPs-based catalysts. Our atomic-scale interface tuning strategy provides a promising approach toward rational design of SA catalysts with excellent catalytic performance and high materials' efficiency.

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Introduction

The cutting edge of rational design of heterogeneous catalysts drives the efficient use of expensive noble metals. Intriguing studies on the catalytic properties of isolated metal ions embedded within oxide supports^{1, 2} or sub-nanometer-sized cluster catalysts³⁻⁵ have provided innovative ideas on the use of low-dimensional motifs of noble metals for enhancing catalytic activity. Recent studies have explored the catalytic functionality of single-atom (SA) catalysts⁶⁻¹². The greater surface area of SAs spontaneously increases the number density of reaction sites, which in turn enhances the specific mass activity (MA) of the catalyst^{8, 11-13}. Moreover, given the improved chemical and

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physical homogeneity of the reaction sites, SA catalysts often comprise outstanding selectivity along with the enhanced activity^{14, 15}.

Although the design of SA-based heterogeneous catalysts is newly emerging, several fundamental issues have not yet been resolved¹⁶. Achieving high catalytic activity¹⁶ in SA catalysts requires the design of catalysts with well-distributed, dense metal monomers. Recent efforts toward stabilization of SAs showed that the chemical potential of SAs can be tuned by utilizing the SA-support¹⁷⁻¹⁹ and SA-reactant interactions^{10, 18-20}. Particularly, CeO₂ and TiO₂ are known as the best supporting oxides for stabilization of Pt-SAs^{16, 21}. The strong interaction between Pt-SAs and supporting oxides leads to preferred formation of ionic Pt species²¹. However, synthesis of dense SAs on such supports for further improved conversion performance is still challenging¹⁶.

In addition to the structural issues, it is still unclear whether the inherent catalytic nature of SAs is different from nanoparticle (NP) counterparts, and SAs are superior to NPs for a given reaction. The interfaces between Pt NPs and oxide supporting materials are generally regarded as a reaction site for oxidation²²⁻²⁴. Naturally, as Pt NPs are pulverized into Pt-SAs, the total length of the Pt-oxide interface increases. However, whether the interface between a Pt-SA and an oxide support exhibits the same catalytic activity with the interface between a Pt NP and an oxide is under veil. Even in the case of CO oxidation, the catalytic behaviour of Pt-SAs is controversial^{25, 26}. Stair and coworkers identified Pt NPs rather than Pt-SAs as the more efficient catalyst for CO oxidation since Pt-SAs are readily deactivated by CO-poisoning²⁵. Meanwhile, Pt-SAs supported on 5 nm-sized TiO₂ NPs are reported to reveal the Mars-van Krevelen (MvK) type of CO oxidation (CO_{ad} + O_{lattice} \rightarrow CO₂ (g)) are inherently immune to CO-poisoning²⁷. Moreover, Nie et al. demonstrated that the CO oxidation activity of Pt-SAs supported on CeO₂ NPs can be improved by steam treatment, corroborating the importance of the chemical nature of the oxygen species at the Pt-CeO₂ interface for activation of the MvK type CO oxidation by Pt-SAs⁷. These results suggest that the catalytic reaction mechanism of oxide supported SAs is strongly correlated with the chemical nature of the metal-oxide interfaces, whose areal fraction is maximized in SAs-based catalysts. Designing an appropriate supporting material is essential to achieve an ideal performance of SA-based catalysts.

CO oxidation has been applied as a prototypical reaction for catalysis studies due to its relatively simple chemistry²⁸. The US Department of Energy 150 °C challenge for emissions (converting 90% of pollutants in automotive exhaust gas at $150 \,^{\circ}\text{C}^{29}$ is driving the design of highly reactive and efficient CO oxidation catalysts for environmental applications¹⁶. Moreover, CO oxidation is a key for chemical energy conversion such as water-gas shift reaction.^{30, 31}

Pt catalysts are prone to CO-poisoning, therefore removal of CO from hydrogen feedstock (below 10 ppm^{32, 33}) is critical when Pt catalysts are applied in polymer electrolyte fuel cell (PEMFC)^{32, 33}. In this aspect, hydrogen purification through preferential oxidation (PROX) of CO under the high H₂ partial pressure conditions³³⁻³⁷ is being more important. A highly CO-



---- TiO2

2.94 eV

CeO_-TiO_

-- ---

Fig. 1 DFT-calculated binding trend of Pt-SAs on $TiO_2(101)$ and at the CeO_x- TiO_2 interface with fully oxidized or reduced Ce ions. The E_{bind} of *n*Pt represents the average binding energy of total number of Pt atoms.

tolerant Pt catalyst would guarantee the improved reliability of the fuel cell-based energy generating systems. Moreover, a Ptbased catalyst with both CO-immunity and high catalytic activity can significantly increase the reliability and efficiency of the catalytic reaction activated by Pt under the CO-rich conditions.

In parallel, the formation of ionic Pt-SAs on oxide supports is accompanied with the oxidation state change of the supporting oxide, especially at the interface region³⁸. Thus, observing the oxidation state of the supporting oxide plays a crucial role in uncovering the mechanism of CO oxidation. For example, reduction of Ce ions at the Pt-CeO₂ interface during CO oxidation accounts for the operation of the MvK type CO oxidation by Pt-SAs. Recently, site-specific infrared (IR) spectroscopy²⁵ and near-ambient pressure X-ray photoelectron spectroscopy³⁸ provided insights for identifying the catalytic active sites but these methods lack spatial information. Additionally, a combination of extended X-ray absorption fine structure (EXAFS) and environmental transmission electron microscopy traced the in-situ structural rearrangement of Pt NPs³⁹, however, quantitative chemical information remains elusive. Thus, qualifying the spatio-temporal evolution of the oxidation state of supporting materials can elucidate both the preferred sites of Pt-SA formation during synthesis and the underlying catalytic mechanism. Particularly, such observation under the real synthesis/working conditions in liquid/gas have not been achieved owing to a lack of spatio-temporal resolution and information about beam-induced damage⁴⁰.

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---- CeO-TiO,



Fig. 2 Morphology of CeO_x -TiO₂ support and *n*PCT catalysts ($n \le 1$). (a) High-angle annular dark-field (HAADF)-STEM images of CeO_x -TiO₂ supporting oxide with 1 wt.% initial Ce-loading. (b) HAADF-STEM and Ce EDS scan images of the CeO_x -TiO₂ presented in (a). HAADF-STEM image and Ce and Pt EDS scan image of 0.25PCT [(c) and (d)], 0.5PCT [(e) and (f)], and 1PCT [(g) and (h)]. Atomic-sized bright spots in (c), (e), and (g) denote Pt-SAs and CeO_x clusters. Some of the CeO_x -clusters and Pt-SAs are highlighted in white circles. Ce and Pt EDS scan images [(d), (f), and (h)] confirm the locational correlation of CeO_x and Pt-SAs on TiO₂ powder surfaces. (i), (j) Intensity profile of the selected highlighted area of 0.5PCT, which clearly show the intimate contact between Pt, Ce, and Ti ions. Arrows in (e) and (g) indicate Pt NPs formed under the higher Pt-loading.

Here, we demonstrate a high-performance dense Pt-SAs catalyst by designing the chemical nature of the multicomponent interface in atomic precision. A CeO_x -TiO₂ hybridoxide support was synthesized to provide strong binding sites and reactive oxygen species for Pt-SAs, therefore, activate the interface-mediated MvK CO oxidation mechanism. We adopt synchrotron-based *operando* liquid/gas scanning transmission X-ray microscopy (STXM) to probe the spatiotemporal evolution of the Ce oxidation state, by which the preferred growth of Pt-SAs at the CeO_x-TiO₂ interfaces and the interface-mediated reaction mechanism can be elucidated.

Results and discussion

 CeO_2 and TiO_2 are the most widely applied supporting oxide materials which can exceptionally stabilize Pt-SAs by formation

of Pt ionic species¹⁶. We designed out Pt/CeO_x -TiO₂ (PCT) catalyst system with reference to theoretical modelling: We constructed a CeO_x -TiO₂ interface model with a single CeO_2 cluster on the (101) surface of anatase-phase TiO_2 (Fig. 1). The formation of ceria clusters on TiO₂(101) surface are thermodynamically preferred under low Ce-loading conditions^{41, 42}. Density functional theory (DFT)-calculated energies of Pt deposition on TiO₂ and at the CeO_x-TiO₂ interface confirmed that the affinity of the CeO_x-TiO₂ interface toward Pt stabilizes the Pt₂ cluster at the interface (Fig. 1). Upon deposition, Pt donates electrons to the Ce and O ions at the CeO_x -TiO₂ interface (Fig. S1 in the ESI⁺). Based on these promising theoretical findings, CeO_x-TiO₂ hybrid-oxides were prepared by the chemical synthesis of CeO_x nanoarchitectures on the surface of ~25 nm-sized TiO_2 particles via thermal reduction of cerium nitrate $(Ce(NO_3)\cdot_3 6H_2O)^{42, 43}$. Small CeO_x

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Fig. 3 Electronic interaction between Pt and Ce upon formation of Pt-SAs at the CeO_x-TiO₂ interfaces, which is directly observed by *operando* X-ray absorption spectroscopy. (a) Averaged optical density map across Ce M_5 -edge. (b) Spatio-temporal chemical evolutions with respect to reaction time. Chemical phase maps obtained by linear combination fits of XAS data at each pixel (see Methods in the ESI⁺). The fitting results (chemical information) and averaged optical density (morphological information) are presented by color legend and transparency, respectively. The presence of CeO₂ and CeO_{2-x} assigned colors blue and red, respectively. (c) *Operando* XAS from entire particle region in (a). Energy positions of M_5 -edge absorption features for Ce³⁺ and Ce⁴⁺ are indicated as black arrows. (d) Ce⁴⁺ fraction versus reaction time. Scatters and yellow shaded area indicate average and standard deviation of Ce⁴⁺ fraction from entire particle region, respectively. The error range of the reaction time is estimated by the liquid injection time (see Methods in the ESI⁺). (e) Distribution plot for initial Ce⁴⁺ fraction versus fraction difference between the first and second measurements (solid red scatters). Solid green line is a guide to eye.

clusters were predominantly observed in the CeO_x-TiO₂ samples with 1 wt.% of Ce addition (Fig. 2a, b), which assures the maximum concentration of CeO_x-TiO₂ interfaces and closely resembles our model structure: a single CeO₂ cluster on TiO₂(101) that can stabilize Pt-SAs by electron donation from Pt. While Ce-loading in the CeO_x-TiO₂ hybrid-oxide was kept at 1 wt.%, various amounts of Pt were impregnated on the CeO_x-TiO₂ support (hereafter, *n*PCT, where *n* = 0.25, 0.5, and 1.0 wt.%; see Table S1 in the ESI⁺ for the composition analysis result). Corresponding control group samples, Pt catalysts on the TiO₂ support without any Ce addition (*n*PT, where *n* = 0.25, 0.5, and 1.0 wt.%), were also synthesized.

Scanning transmission electron microscopy (STEM) images of the supporting CeO_x-TiO₂ (1CT) and, and *n*PCT catalysts are presented in Fig. 2. STEM images of *n*PCT show that tiny, bright, and highly distinguishable Pt-SAs are distributed over the surface of *n*PCT catalysts (Fig. 2c, e, and g). Energy-dispersed Xray spectroscopy scan images (Fig. 2d, f, and h) and the intensity profiles of the selected highlighted areas of 0.5PCT (Fig. 2i, j) corroborate that the Pt-SAs are strongly correlated with the preformed CeO_x clusters on TiO₂, thus affirming the strong Pt affinity of CeO_x clusters. Consistent with the recent findings^{10, 17, ⁴⁴, Pt-SAs could be preferentially synthesized owing to the SAsupport interaction. On the other hand, the population of the sub-nanometer-sized Pt NPs were relatively greater in *n*PT samples than *n*PCT samples (Figs. S2 and S3 in the ESI⁺).}

Spatio-temporal observation was performed using operando synchrotron-based STXM combined with a liquid flow nanoreactor (see Methods and Fig. S4 in the ESI⁺) to probe the preferred in-situ Pt-SAs formation onto CeO_x-TiO₂. In this case, a CeO_x -TiO₂ sample with 6 wt.%, instead of 1 wt.%, was used to obtain distinct absorption features (both Ce³⁺ and Ce⁴⁺, Figs. S5 and S7 in the ESI⁺) and measurable absorption contrast. Using a microfluidic heating cell, we tracked the change in Ce oxidation state that accompanies the Pt-SAs growing onto CeO_x-TiO₂ (see Supplementary discussion, Fig. 3b and Figs. S6 and S7 in the ESI⁺). Fig. 3b shows operando image spectra of a typical particle aggregate, which contains a statistically significant number of ~ 25 nm-sized CeO_x -TiO₂ particles. Although, the spatial resolution of the STXM image spectra (~ 55 nm with the 45 nm outer-zone-width Fresnel zone plate) cannot directly visualize the individual CeOx-TiO2 particle or Pt-SA, the heterogeneous distributions of the Ce oxidation states were clearly observed by analyzing X-ray absorption near-edge structure (XANES) on a pixel-by-pixel basis with reference spectra (Fig. 3b). Of particular note is that the maximum dose for individual pixel was carefully limited to eliminate partial photoreduction of CeO₂ (see Methods and Fig. S6 in the ESI⁺), therefore highly reliable quantitative analysis about Ce oxidation state was available. During the initial stage of the Pt-SAs formation, the Ce oxidation states averaged over each particle were continuously reduced (Fig. 3c, d). Interactions

er from information (Fig. S9 and Table S2 in the ESI⁺). The relative amounts of Pt and PtO_2 in the *n*PT and *n*PCT catalysts also



Fig. 4 DFT-predicted MvK-type CO oxidation pathway catalyzed by a Pt₂ cluster pinned at the CeO_x-TiO₂ interface. ΔE_n denotes the relative energy of the n^{th} stage with respect to the $(n-1)^{\text{th}}$ stage. For example, ΔE_3 was calculated as follows: ΔE_3 = Energy of stage 3.

A surprising observation for the heterogeneous Ce oxidation states during the Pt-SAs formation indicates the strong correlation between the initial $\mbox{Ce}^{\mbox{\tiny 3^+}}$ concentration and the relative Ce oxidation state change, ΔCe^{4+} (pixel-by-pixel Ce⁴⁺ concentration differences between the first and the second image spectra, Fig. 3e). In general, the pixels in which the initial $\mbox{Ce}^{\mbox{\tiny 3+}}$ fraction is higher were more reduced during the Pt-SAs formation. Since, the Ce³⁺ phase is highly localized at the CeO_x-TiO₂ interface (see Supplementary discussion in the ESI⁺), the strong spatial correlation in Fig. 3e suggests that the CeO_x-TiO₂ interface and not the CeO₂ surface is the preferred site for Pt-SAs growth. Therefore, along with our DFT calculation results and STEM observations, the operando STXM observations confirm the effectiveness of the $\text{CeO}_{x}\text{-}\text{TiO}_{2}$ interface in disintegrating Pt NPs. Interestingly, the concentration of the reduced Ce³⁺ upon Pt-SAs formation recovers to the original value after further Pt deposition (after 250 min, Fig. 3d), upon emergence of Pt NPs that do not excessively donate electrons to Ce ions.

The XANES and EXAFS across the Pt *L*-edge also indicate that CeO_x -TiO₂ stabilizes part of the Pt-SAs even under high Ptloading conditions (n = 1, Fig. S8 in the ESI⁺) by forming ionic Pt phases. This is shown by the rightward shift of the peak in the XANES, indicating an increased oxidation state for Pt, and the appearance of a peak in the EXAFS Fourier magnitude at the same position (1.7Å indicated) as in PtO₂. Because the XANES spectra show that the Pt in the lower *n* in *n*PCT and *n*PT catalysts is the more ionic (PtO₂ like, Fig. S9 in the ESI⁺), all the XANES spectra were fit by a linear combination using reference standard spectra of Pt and PtO₂ to provide quantitative confirm the increased content of ionic Pt in nPCT, compared to nPT.

Based on our findings of the preferential formation of Pt-SAs at the CeO_x-TiO₂ interface, DFT calculations were geared to scrutinizing the mechanism of CO oxidation that could be catalyzed by Pt-SAs at the CeO_x-TiO₂ interface. Fig. 4 suggests the CO oxidation pathway by the Pt_2/CeO_x -TiO₂ model catalyst. We found that the Pt₂ cluster at the CeO_x-TiO₂ interface binds two CO molecules together (stages 1 and 2, hereafter S1 and S2, respectively). The interfacial oxygen ion bridging the Ce and the Ti ions and binding the Pt₂ cluster oxidizes a Pt-bound CO molecule (Pt-CO*) through the interface-mediated MvK mechanism. The formation of an O-C-O-type intermediate requires an activation energy barrier, Eact, of 0.75 eV (marked as TS1, Fig 4). After the relatively easy production of CO_2 (S3), the Ce ion of Pt₂/CeO_x-TiO₂ binds and activates an oxygen molecule (S4). One oxygen atom of the Ce-bound O₂ molecule oxidizes another Pt-CO* and the other O atom heals the vacancy, thereby completing the overall MvK type reaction pathway. An E_{act} of 0.43 eV was required for dissociation of the Ce-bound O₂ (TS2). Our results show that the CO oxidation pathway involves the lattice oxygen at the tri-phase interface of Pt-CeO_x-TiO₂ oxidizing the first Pt-CO^{*} via the MvK mechanism. The relatively higher E_{act} of the first CO oxidation indicates that activation of the lattice oxygen at the Pt₂-CeO_x-TiO₂ interface is the ratedetermining step.

The catalytic properties of both *n*PCT and *n*PT were also investigated for CO oxidation in the temperature range of 50-300 °C. Fig. 5a show the light-off performance of the *n*PT and *n*PCT catalysts, respectively. Clearly, the *n*PCT catalysts oxidize

CO more actively than the *n*PT for all Pt-loadings. T_{50} and T_{100} (the temperature of 50% and 100% conversion, respectively) of *n*PT are higher than those of *n*PCT with the same *n* (Table S3 in the ESI⁺). As shown in Fig. 5b and Table S3 in the ESI⁺, the specific MA of *n*PCT samples at 140 °C is always higher than that of *n*PT with the same amount of Pt-loading.

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role of the MvK mechanism with the higher E_{act} . Once the MvK mechanism becomes active above 100 °C, the MA of 0.25PCT rapidly increases at the elevated temperature. In contrast, the MA of 0.25PT does not significantly increase by the temperature increase despite its lower E_{act} . In addition, the MA of 0.25PT is less than that of 0.5PT, which is consistently observed in the



Fig. 5 CO oxidation performance of *n*PT and *n*PCT catalysts. (a) Temperature-programmed CO conversion profile (light-off curve) of *n*PT and *n*PCT catalysts. (b) The specific mass activity (MA) of *n*PT and *n*PCT catalysts at 140 °C as a function of Pt-loading and the ECSA of *n*PT and *n*PCT catalysts. The MA of *n*PCT catalysts is an inverse function of Pt-loading, whereas the MA of 0.25PT is lower than the others. (c) The specific MA values of *n*PT and *n*PCT catalysts plotted as a function of T_{s_0} . Reference values were limited to Pt-SAs or NPs supported on conventionally synthesized or purchased CeO₂ or TiO₂. (refer to ESI⁺ for the full reference list). (d) CO-TPR analysis results of 0.25PT and 0.25PCT catalysts.

Here, we emphasize that the E_{act} of *n*PCT is close to the DFTcalculated E_{act} (0.75 eV) *via* the MvK-type CO oxidation at the Pt₂-CeO_x-TiO₂ interface (Fig. S10 in the ESI⁺). Therefore, the MvK type mechanism plays a major role in *n*PCT catalysts. Despite the relatively higher E_{act} of the *n*PCT catalysts, they exhibited outstanding catalytic performance, higher MA, compared with *n*PT catalysts with even lower E_{act} . This enhancement can be understood by the presence of Pt-SAs at the CeO_x-TiO₂ interfaces in *n*PCT, as already confirmed by experimental observation (STEM and XAS). It is further supported by additional CO adsorption measurement, which reveals the electrochemical surface area (ECSA) in 0.25PCT is 55% higher than that in 0.25PT owing to the preferential formation of Pt-SAs (Fig. 5b).

Notably, despite high ECSA values, the MA of 0.25PT was significantly lower than that of 0.5PT (Fig. 5b and Table S3 in the ESI⁺). The MA of *n*PT and *n*PCT catalysts plotted as a function of temperature (Fig. S11 in the ESI⁺) shows that the MA of 0.25PCT increases upon temperature increase, assuring the essential

whole temperature range. These findings are indicative of CO poisoning of the Pt species in 0.25PT.

Our DFT calculation results confirm that the Pt-SAs formed at the CeO_x -TiO₂ interface are immune to surface poisoning by CO and thus activate the MvK type CO oxidation (Fig. S12 in the ESI⁺). In contrast to 0.25PCT, the high CO₂ production energy of 1.28 eV calculated from the CO-saturated TiO₂(101) supported Pt₉ NP, which stands for small Pt NPs in 0.25PT, predicts that the MvK type CO oxidation pathway is energetically forbidden in 0.25PT (Fig. S12 in the ESI⁺). As a result, our 0.25PCT exhibits the best MA-T₅₀ balance among the previously reported TiO_2 or CeO₂ supported Pt-SA and NP catalysts (Fig. 5c). Moreover, our 0.25PCT catalyst was synthesized by simple and hierarchical chemical route. The consistent light-off performance of 0.25PCT catalyst for repeating six CO oxidation cycles suggests that the catalytic nature of the Pt-SAs at the CeO_x-TiO₂ interfaces is wellpreserved during long-term operation (Fig. S13 in the ESI⁺). This result convinces that stabilizing potential based interface design is successful to improve the MA as well as stability of Pt-SAs. The

long-term temperature- and time-resolved stability test results of Pt-SAs supported on TiO_2 and CeO_x - TiO_2 will be studied for future work.

To directly unravel the operation of the MvK mechanism in *n*PCT catalysts, we performed complementary CO-temperature programmed reduction (TPR) analysis with 0.25PT and 0.25PCT catalysts (Fig. 5d). The CO-TPR analysis was performed under the stream of 1% of CO balanced with 99% of Ar without any oxygen supply, therefore the oxygen to participate in the CO oxidation should be supplied through the Pt-CeO_x-TiO₂ or the Pt-TiO₂ interfaces. Fig. 5d shows that CO₂ was produced in 0.25PCT at above 135 °C whereas no CO₂ was detected in



Fig. 6 *Operando* gas phase XAS analysis. (a) Schematic diagram of Ce reduction upon CO oxidation by the MvK mechanism at the Pt-CeO_x-TiO₂ interface. (b) *Operando* XAS with respect to reaction temperatures. Owing to low signal-to-noise ratio, XAS was obtained from entire particle region. De-convoluted spectral features for Ce³⁺ (red and magenta shaded areas, energy positions are indicated as red arrows) and Ce⁴⁺ (blue shaded area, energy positions are indicated as blue arrow) are superimposed on averaged XAS. (c) Area ratio between de-convoluted spectral features of Ce³⁺ and Ce⁴⁺ in (b).

0.25PT. This result provides convincing evidence about the presence of the CO-tolerant Pt-SAs and the CO-poisoned Pt species in 0.25PCT and 0.25PT, respectively.

Synchrotron-based operando gas STXM was utilized to validate the operation of the MvK mechanism (Fig. 6a). The MvK mechanism can be completely identified by probing XAS across the Ce *M*-edge under the same condition with CO-TPR. The results presented in Fig. 6b and 6c reveal that the production of CO₂ in 0.25PCT accompanies the reduction of Ce⁴⁺ to Ce³⁺, which results from the loss of oxygen at the Pt-CeO_x-TiO₂ interfaces as shown schematically in Fig. 6a. The reaction temperature at which the Ce⁴⁺/Ce³⁺ ratio is significant is consistent with that of CO₂ production from CO-TPR. Our CO-TPR and operando XAS results strongly indicate the presence of the CO-tolerant Pt-SAs in 0.25PCT catalyst and the operation of the interface-mediated MvK mechanism of CO oxidation.

We present the catalytic supremacy of dense Pt-SAs achieved by tailoring the chemical nature of the multi-component interface with atomic precision. Addition of 1 wt.% Ce to TiO₂ supporting oxide powders creates CeO_x-TiO₂ interfaces that physically disintegrate Pt into Pt-SAs and activate the interfacemediated MvK mechanism of CO oxidation, while preserving its catalytic activity against CO-poisoning. As a result, the Pt-SAs stabilized at the CeO_x-TiO₂ hybrid interfaces increase the MA of Pt toward CO oxidation by 15.1 times at 140 °C, showing the highest mass activity among similar systems reported so far. We adopt synchrotron-based operando liquid/gas scanning transmission X-ray microscopy (STXM) to probe the spatiotemporal evolution of the Ce oxidation state which indicates the preferred growth of Pt-SAs at the CeO_x-TiO₂ interfaces and the interface-mediated reaction mechanism in CO oxidation for the first time. Our comprehensive study of DFT calculations, operando STXM analyses, and experimental catalyst performance test results present a novel design strategy of Pt-SA catalysts by atomically precise interface engineering toward efficient use of Pt catalysts.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

Conclusions

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TOC entry

Catalytic Supremacy of Pt-Single Atoms Achieved by ${\rm CeO}_x{\rm -TiO_2}$ Interfaces

