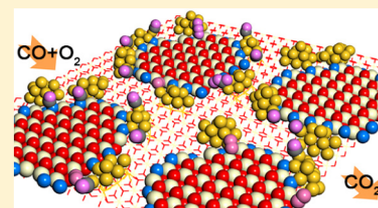


CO Oxidation at the Interface of Au Nanoclusters and the Stepped-CeO₂(111) Surface by the Mars–van Krevelen MechanismHyun You Kim^{*,†} and Graeme Henkelman^{*}

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S Supporting Information

ABSTRACT: DFT+U calculations of CO oxidation by Au₁₂ nanoclusters supported on a stepped-CeO₂(111) surface show that lattice oxygen at the step edge oxidizes CO bound to Au NCs by the Mars–van Krevelen (M-vK) mechanism. We found that CO₂ desorption determines the rate of CO oxidation, and the vacancy formation energy is a reactivity descriptor for CO oxidation. Our results suggest that the M-vK mechanism contributes significantly to CO oxidation activity at Au particles supported on the nano- or meso-structured CeO₂ found in industrial catalysis.



SECTION: Surfaces, Interfaces, Porous Materials, and Catalysis

Cerium oxide is a promising candidate for advanced heterogeneous catalysis as well as a prevalent supporting material for catalysis at supported Au nanoparticles (NPs).^{1,2} The unique properties of CeO₂ have been recognized as a combination of high oxygen storage and release capacity, facile oxygen vacancy formation, and the presence of a sharp Ce *f*-band.^{1,2} Experimental studies have shown that the surface of CeO₂ can be enriched with oxygen vacancies and that metal NPs or nanoclusters (NCs) bind strongly to these vacancies.^{3–5} Moreover, the step edges on CeO₂ surfaces have also been recognized as a binding site of metal NPs/NCs.⁴

The oxidation chemistry of CeO₂ supported Au NPs/NCs (Au/CeO₂) has been extensively studied, and especially the reactive species.^{6–10} The critical role of the interface between the supporting oxide and the supported metal NPs/NCs has been highlighted by experimental and theoretical studies in several systems.^{6–18} Rodriguez and co-workers have shown that the Au supported CeO₂ inverse-catalyst has a high activity for water–gas shift reaction compared to the conventional Au/CeO₂ catalyst.¹¹ Zhou et al. showed a clear relationship between the length of the Au–CeO₂ interface and the CO oxidation reactivity on the Au/CeO₂ multilayers.¹⁹ The Neurock and Yates groups showed that the Au–TiO₂ interface activates the following reactions: CO oxidation,¹² partial oxidation of acetic acid,²⁰ and low-temperature H₂ oxidation.²¹ Very recently, Bruix et al. reported that lattice oxygen of a (Ce–Ti)₂O mixed oxide can directly participate in the water–gas shift reaction by supported Pt NCs.¹⁴

Another interesting feature of CeO₂-supported metal NPs/NCs is the nanosize effect of the CeO₂ support on the catalytic activity. Several recent reports on the nanosize effect of CeO₂ support on the catalytic activity of supported metal NPs/NCs confirm the relevancy of the metal–CeO₂ interface to the oxidation catalysis.^{7,8,13} Guzman et al. reported experimental results showing that the nanosized CeO₂ increases the CO oxidation activity by supported Au and that the CeO₂ support supplies the reactive oxygen species directly to the supported

Au NPs.⁸ The Neyman and Illas groups reported a lower vacancy formation energy (E_{vac}) of nanosized CeO₂, and showed that the reactive oxygen atoms are supplied by oxygen spillover from the CeO₂ to supported Pt NPs.¹³

In our previous studies, we have suggested that O₂ bound to the Au–Ce³⁺ interface oxidizes CO bound to Au NCs.¹⁰ We also found that lattice oxygen from the CeO₂ surface can be directly used as an oxygen source for CO oxidation at the Au–CeO₂ interface by cation doping.⁹ From an industrial perspective, however, a model system of Au NPs/NCs supported on a flat defect-free CeO₂ surface is not particularly relevant. Au NPs/NCs supported on rough polycrystalline surfaces, oxide NPs, or meso-structures such as zeolites would be more appropriate models for industrial-grade catalysts.

In this Letter, we are taking a step toward modeling real catalysts by studying the mechanism of CO oxidation by Au₁₂ NCs supported on stepped-CeO₂, using density functional theory (DFT). We are focusing on the role of the interface between the supporting CeO₂ and the supported Au NCs via previously reported interface-mediated CO oxidation mechanisms: oxygen spillover from the CeO₂ to the Au NPs;⁸ CO oxidation by the O₂ bound to the Au–Ce³⁺ interface;¹⁰ and CO oxidation by the Mars–van Krevelen (M-vK) mechanism.⁹ We find that the low E_{vac} of the stepped-CeO₂ support promotes CO oxidation at the Au–CeO₂ interface by the M-vK mechanism. The experimentally reported CeO₂ NP/Au inverse catalyst¹¹ was also studied to confirm our computational results. Our findings provide insight into the oxidation catalysis by Au/CeO₂ catalysts and the origin of their catalytic activity.

Nilius et al. have resolved the morphology of steps in the CeO₂(111) surface and reported three different structures: two types of $\langle 110 \rangle$ steps and one $\langle 211 \rangle$ step.²² They confirmed that the simulated STM images of these DFT-interpreted steps were

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in close agreement with experimental STM data. We reproduced these three types of steps on the $\text{CeO}_2(111)$ surface (CeO_2 -step) (refer to Figure 1 for the detailed

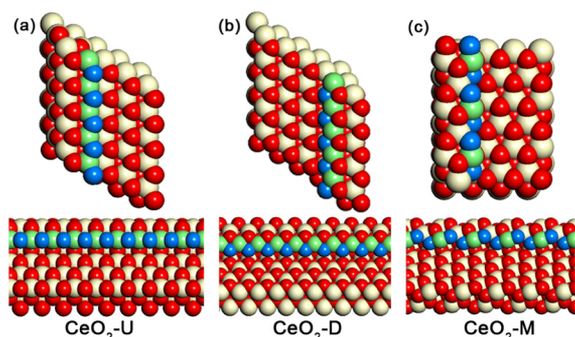


Figure 1. Three kinds of steps on the $\text{CeO}_2(111)$ surface used for the current study. CeO_2 -U (a) and CeO_2 -D (b) describe the $\langle 110 \rangle$ type steps with upper and lower rows of oxygen atoms, respectively. CeO_2 -M (c) describes the $\langle 211 \rangle$ -type step with mixed type of edge oxygen atoms. Ivory and red spheres represent Ce and O atoms, respectively. Ce and O atoms along the step edge are colored in green and blue, respectively.

morphology). The CeO_2 -U and CeO_2 -D ($\langle 110 \rangle$ -type steps) were produced by adding a $2/5$ ML of three atomic layers of CeO_2 (a $\text{Ce}_{10}\text{O}_{20}$ wire) on a 5×5 $\text{CeO}_2(111)$ slab with six atomic layers. The CeO_2 -M ($\langle 211 \rangle$ -type step) was constructed by adding a $3/8$ ML of three atomic layers of CeO_2 (a Ce_9O_{18} wire) on a $4 \times 3\sqrt{3}$ of $\text{CeO}_2(111)$ slab with six atomic layers. The edge atoms along one side of a partial CeO_2 ad-layer was used to model CO oxidation at a CeO_2 -step. Differences in E_{vac} between our systems and thicker models (total 12 atomic layers) are smaller than 30 meV, demonstrating convergence with respect to system size. A Au NC comprising of 12 atoms was optimized at the step edge of each CeO_2 -step to generate the Au/ CeO_2 -step models (see Figure 2a,b,c). Three lowest

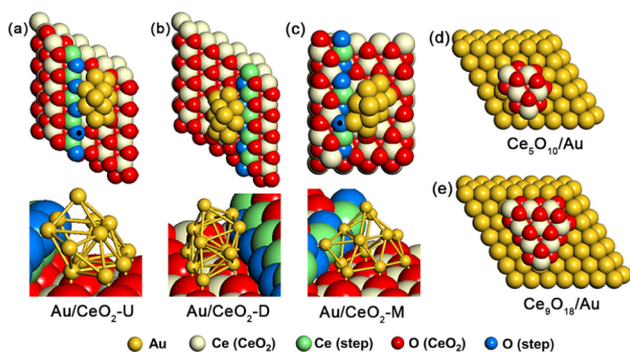


Figure 2. Morphology of studied catalyst models: (a–c) Au/ CeO_2 -step and (d,e) CeO_2 /Au. Black dots represent the location of edge oxygen atoms that were tested for M-vK mechanism of CO oxidation.

atomic layers of CeO_2 were fixed during optimizations. The structure of Au_{12} NC was adapted from our previously reported Au_{13} NC.^{9,10} One corner Au atom was removed from the side of Au_{13} NC to produce a better Au- CeO_2 -step interface structure. Detailed information on the stability of our Au NC and the selection of the CeO_2 support can be found in our previous studies.^{9,10}

Two models of CeO_2 /Au inverse catalysts were constructed optimizing a two-dimensional (three atomic layers) Ce_5O_{10} and Ce_9O_{18} island on four layers of a 6×6 and 7×7 Au(111) slab, respectively (see Figure 2d,e). The two lowest Au layers were held fixed for geometry optimization.

We performed spin-polarized DFT calculations with the VASP code²³ and the PBE²⁴ functional. In order to treat the highly localized Ce 4f-orbital, DFT+ U ²⁵ with $U_{\text{eff}} = 5$ eV was applied.^{9,10} The interaction between the ionic core and the valence electrons was described by the projector augmented wave method,²⁶ and the valence electrons with a plane wave basis up to an energy cutoff was 400 eV. The Brillouin zone was sampled at the Γ -point. The convergence criteria for the electronic structure and the geometry were 10^{-4} eV and 0.01 eV/Å, respectively. We used the Gaussian smearing method with a finite temperature width of 0.1 eV in order to improve convergence of states near the Fermi level. The location and energy of transition states (TSs) were calculated with the climbing-image nudged elastic band method.^{27,28} E_{vac} , the energy required to remove an oxygen atom from the CeO_2 -step, was calculated with reference to the energy of gas phase O_2 .

A comparison of E_{vac} between the reference 5×5 , nine-layer $\text{CeO}_2(111)$ slab and three CeO_2 -step structures (CeO_2 -U, CeO_2 -D, and CeO_2 -M) shows that oxygen atoms at the step edge are weakly bound to the CeO_2 matrix (Table 1). This is natural due to the lower coordination of these oxygen atoms. Considering previously reported low E_{vac} of CeO_2 nanocrystallites²⁹ and their enhancing effect on the catalytic activity of supported Pt NPs,¹³ the CeO_2 -step is also expected to promote CO oxidation by the Au/ CeO_2 -step. The E_{vac} of three CeO_2 -step structures lie between 1.5 and 1.7 eV, which is consistent with literature values.²⁹ These E_{vac} values also fall near the previously suggested maximum E_{vac} for the activation of the M-vK mechanism of CO oxidation by Au NC supported on doped- CeO_2 .⁹ The E_{vac} of the second vacancy at the first nearest neighboring position from the first vacancy (Table 1) shows that the vacancy pairing is a rare event in CeO_2 -U and CeO_2 -D steps, whereas the divacancy can be occasionally found at the step edge of the CeO_2 -M.

Figure 2 shows three different optimized initial Au/ CeO_2 -step models. Low-coordinated surface sites on the CeO_2 surface, such as at step edges or surface vacancies, are regarded as strong binding sites for metal (Au or Ag) NPs.^{4,10} We found that the CeO_2 -step strongly binds Au NCs, making it a preferential binding site of supported Au NPs/NCs (Table S1). The binding energy of a Au NC on the CeO_2 -U and CeO_2 -M

Table 1. E_{vac} Calculated in CeO_2 -step and Au/ CeO_2 -step^a

	CeO_2 -STO ^b	CeO_2 -U	CeO_2 -D	CeO_2 -M	Au/ CeO_2 -U	Au/ CeO_2 -M
E_{vac} (eV)	2.48	1.66 (2.07)	1.62 (2.17)	1.52 (1.62)	1.75	1.02

^aValues in parentheses represent the E_{vac} of the second vacancy. E_{vac} of two Au/ CeO_2 -step systems where the M-vK mechanism of CO oxidation was tested are presented. ^bAdopted from ref 9.

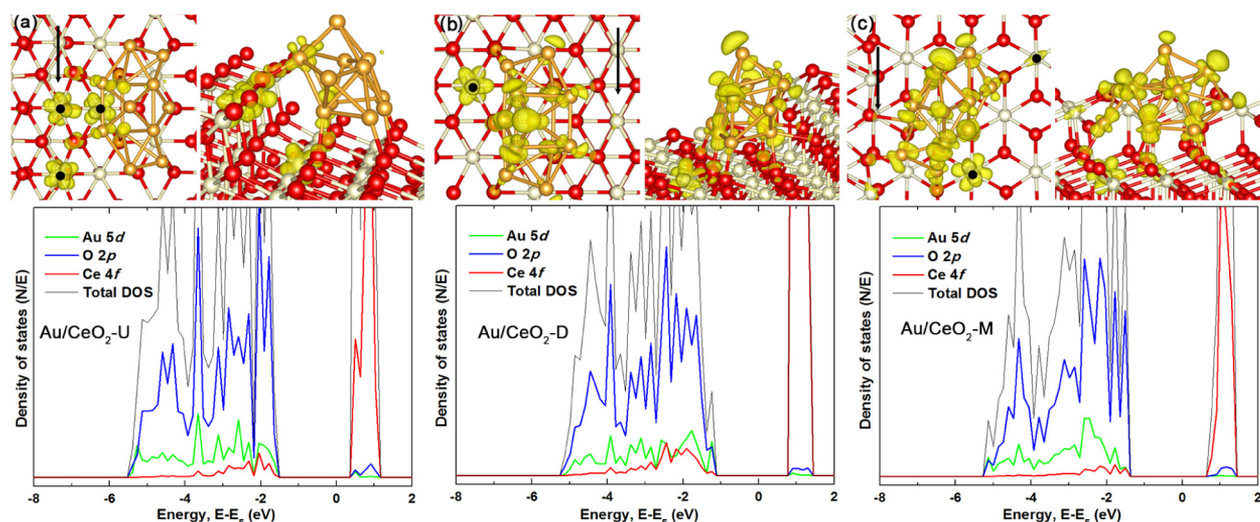


Figure 3. Density plots of the two highest occupied Kohn–Sham molecular orbitals associated with the introduction of Au NP on CeO₂-step: (a) CeO₂-U, (b) CeO₂-D, and (c) CeO₂-M. Corresponding total and partial density of states are presented below. Black dots represent the location of reduced Ce³⁺ ions upon supporting of Au NP. Black arrow represents the step edge. Yellow, ivory, and red spheres represent Au, Ce, and O atoms, respectively. Density of states was calculated with a $4 \times 4 \times 1$ K-points grid (10 irreducible K-points). Hybridization between O-2p and Au-5d orbitals are responsible for the strong Au–CeO₂-step interaction. O-2p, Au-5d, and Ce-4f orbitals are contributing to the frontier orbitals.

steps was even stronger than that on the stoichiometric CeO₂ and partially reduced CeO₂ surfaces with a divacancy (Table S1). Stronger Au binding on CeO₂-step confirms the previously reported binding tendency of Au NPs on polycrystalline CeO₂ substrate; Au NPs prefer the step edge.⁴

The location of the frontier orbitals (the sum of the charge densities of two highest lying Kohn–Sham molecular orbitals) of Au/CeO₂-U, Au/CeO₂-D, and Au/CeO₂-M and their density of states, presented in Figure 3, show that Au binding at the CeO₂-step is through Au–O bonds (hybridization of Au-*d* and O-*p* orbitals). Reduced Ce³⁺ ions are located at the neighboring sites to the Au–O bonds composing the highest occupied molecular orbital (HOMO).

CO oxidation at the Au–CeO₂-step interface was studied considering three mechanisms: the M-vK mechanism of CO oxidation⁹ (oxidation of Au–CO* by an oxygen atom at the step edge), the oxygen spillover mechanism^{8,13} (oxidation of Au–CO* by an oxygen atom spontaneously transferred from the step edge to the Au NC), and CO oxidation by O₂ bound to the Au–Ce³⁺ interface.¹⁰

In the case of Au/CeO₂-step structures, the spillover of lattice oxygen of the CeO₂ step was not found to be energetically favorable. The binding energy of atomic oxygen on the Au NP is -0.5 eV, which is close to the experimental binding energy estimated by Campbell's group for tiny Au particles on TiO₂(110) (-0.85 eV),³⁰ requiring more than 1.0 eV for oxygen transfer. We also have examined the oxygen spillover mechanism on Au NCs supported on stoichiometric CeO₂,¹⁰ partially reduced CeO₂,¹⁰ and doped-CeO₂⁹ and found that oxygen spillover is energetically prohibited in these systems as well. We postulate, therefore, that this mechanism, which requires either a low E_{vac} of CeO₂ support or strong binding of atomic oxygen on Au NPs/NCs, is a special case for Au NPs/NCs supported on small CeO₂ nanocrystallites or NPs, for example, the truncated-octahedral Ce₈₀O₁₆₀ NP reported by Neyman and co-workers²⁹ where the E_{vac} is as low as 0.46 eV (PW91+U calculation). As reported by Neyman and co-workers, CO oxidation by CeO₂-supported strong oxo-phillic

NPs such as Pt NPs would favor the oxygen spillover mechanism.¹³

In the case of Au/CeO₂-U where two Ce⁴⁺ ions at the step edge are reduced (Figure 3a), we examined whether these Au–Ce³⁺ interface binds a gas phase O₂ and activates CO oxidation by O₂ bound to the Au–Ce³⁺ interface. The calculated binding energy of gas phase O₂ at the Au–Ce³⁺ interface is as low as -0.36 eV. Because the entropic contribution to the Gibbs free energy of O₂ desorption at the conventional operating temperature of CO oxidation (-0.64 eV at 298 K and 1 bar; the standard entropy of O₂ at 298 K is 205.14 J mol⁻¹ K⁻¹; see Supporting Information)^{31,32} is greater than the binding energy, O₂ coverage at the Au–Ce³⁺ interface would be lower than that of Au–CO*.¹⁰ Presumably, reduced Ce³⁺ ions and low coordinated oxygen atoms at the step edge stabilize each other in the form of local Ce₂O₃ clusters, thus, reduced Ce³⁺ ions are reluctant to bind additional gas phase O₂.

The next step of CO oxidation by O₂ at the Au–Ce³⁺ interface takes place on the surface of the Au NC.¹⁰ Since the activation energy is 0.08 eV at the interface of the Au NC and the reduced CeO₂(111),¹⁰ we postulate a low barrier in the case of our Au/CeO₂-step as well. In such a case, the relative coverage of CO and O₂ at the interface would account for the preferred reaction mechanism. Because CO adsorption is 3 times stronger than that of O₂, the M-vK mechanism would be favored for CO oxidation by the weakly bound O₂ molecule at the Au–Ce³⁺ interface. In other cases, where the reduced Ce³⁺ ion is located at the terrace of the CeO₂(111) support (Figure 3), we found that the Au–Ce³⁺ interface does not bind O₂ strongly, which is consistent with what we have reported in Au NCs supported on doped-CeO₂.⁹ Likely, the presence of reduced Ce³⁺ is not a sufficient condition for strong O₂ binding at the Au–Ce³⁺ interface. We hypothesize that the binding of O₂ at the Au–Ce³⁺ interface¹⁰ is a result of the systematic attribution of the Ce³⁺ ion adjacent Au^{δ-} or low coordinated Au atom.

O₂ binding at the Au–Ce³⁺ interface was also examined in Au NCs supported on a partially reduced CeO₂-step with one or two vacancies. We found that O₂ prefers to fill the vacancy

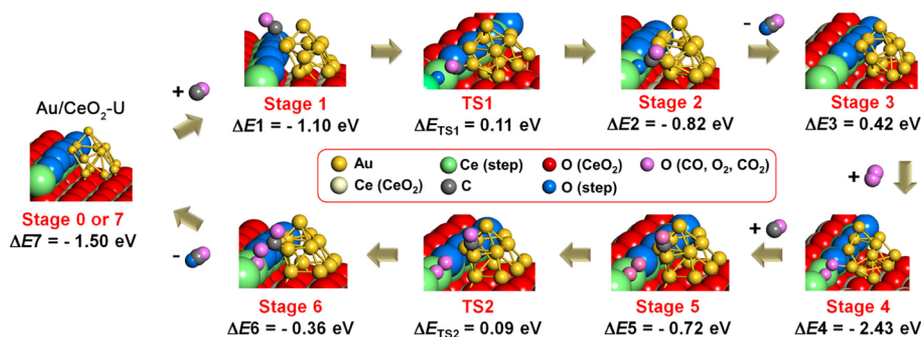


Figure 4. Energetics and the morphology of reaction intermediates of CO oxidation by Au/CeO₂-U. ΔE_x is the energy of the x^{th} state relative to that of the previous stage; for example, ΔE_2 is the energy difference between stage 2 and stage 1. ΔE_{TS_x} is the energy of the x^{th} TS relative to the previous stage (energy barrier of the x^{th} TRI formation).

rather than binding at the Au-Ce³⁺ interface. O₂ bound to the vacancy will oxidize CO by the second-half of the M-vK mechanism.

CO oxidation by the M-vK mechanism was studied in the Au/CeO₂-M and Au/CeO₂-U systems, where CO bound to Au NC (Au-CO*) is in close contact with one of lattice oxygen atoms at the CeO₂-step. The CO oxidation pathway shown in Figure 4 (Au/CeO₂-U) is a typical M-vK mechanism of CO oxidation that we have reported for a Au NC supported on CeO₂ surfaces; CO binding on Au NC (Au-CO*), formation of a triangular reaction intermediate (TRI), CO₂ desorption and vacancy formation, vacancy healing by a gas phase O₂ molecule (V-O₂*), and further oxidation of Au-CO* by a protruding oxygen atom of V-O₂*.⁹ The low barrier for TRI formation confirms that the desorption of the first CO₂, which accompanies vacancy formation at the CeO₂-step, is the rate-determining step of CO oxidation by the M-vK mechanism. In both cases, the energy of CO₂ desorption can be supplied by the entropic contribution to the Gibbs free energy of CO₂ desorption (-0.66 eV at 298 K and 1 bar); the standard entropy of CO₂ at 298 K is 213.79 J mol⁻¹ K⁻¹),^{31,32} suggesting that CO₂ production would not impede the reaction (see Supporting Information).⁹ Experimentally observed apparent activation energies for CO oxidation by Au-CeO₂ catalysts are in the vicinity of 0.3 to 0.5 eV, depending on the size of Au NCs and the dimension of CeO₂.³³⁻³⁵ The energy of CO₂ desorption ($\Delta E_3 = 0.42$ eV, Figure 4) is consistent with these experimental findings.

Previously, we have reported that the energy of CO₂ desorption from the first TRI of Au NCs supported on a doped-CeO₂ surface is a function of the E_{vac} of Au/doped-CeO₂.⁹ Two additional data points from the Au/CeO₂-U and Au/CeO₂-M (see Supporting Information, Table S2, for energetics of CO oxidation by Au/CeO₂-M) were added to Figure 5. These two data points lie on the red line, which is parallel to the previously reported trend line of the Au/CeO₂ surface. The shift in the data points is likely the result of the size effect in CeO₂, and it shows that the CeO₂-step is a more favorable and volatile oxide support than the flat CeO₂ surface for CO oxidation by the supported Au NPs. The maximum E_{vac} for the activation of the M-vK mechanism of CO oxidation at the interface of Au/CeO₂-step is calculated as 2.10 eV, which is higher than that of the Au/CeO₂ surface (1.58 eV) by 0.52 eV; the Au/CeO₂-step whose E_{vac} is lower than 2.10 eV would be eligible for the spontaneous CO₂ production from the first TRI (see Supporting Information for details). It is not clear yet whether data from other CeO₂ nanostructures, for example

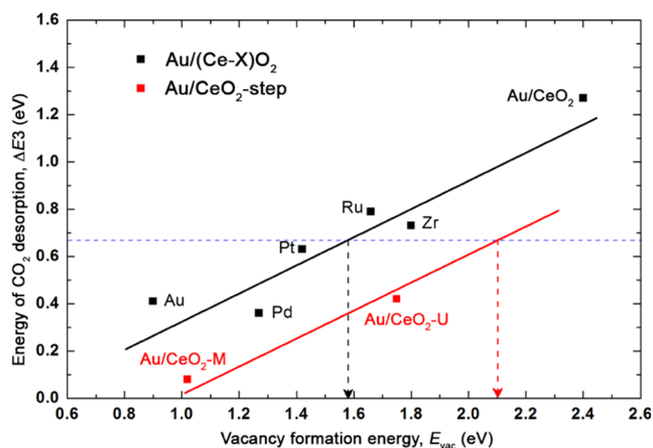


Figure 5. A linear relation is found between E_{vac} and the energy of CO₂ desorption (ΔE_3 as defined in Figure 4) over Au NCs supported on CeO₂-step surfaces, Au/CeO₂-step, (red data), and doped CeO₂ surfaces, Au/(Ce-X)O₂ (X = Au, Pd, Pt, Ru, or Zr) (black data). Data points of Au/(Ce-X)O₂ were adopted from ref 9. The blue dashed line represents the boundary where ΔE_3 can be overcome by the entropic contribution to the Gibbs free energy of CO₂ desorption, which is -0.66 eV at 298 K. The black and red dashed lines represent the maximum E_{vac} for the activation of the M-vK mechanism of CO oxidation by Au/(Ce-X)O₂ (1.58 eV) and Au/CeO₂-step (2.10 eV), respectively.

CeO₂ NPs or curved steps, will fall on the current trend line (red line in Figure 5). Further investigations would provide a comprehensive picture of how the dimension of nanostructured CeO₂ affects their oxidation potential. We speculate that, as shown in Figure 5, a specific morphology of the CeO₂ nanostructure with a lower E_{vac} would be more favorable for oxidation catalysis by the M-vK mechanism.

Rodriguez and co-workers have reported the excellent catalytic activity of CeO₂/metal inverse catalyst, and their results are recognized as a good example of the critical role of the metal-oxide interface on the oxidation catalysis.^{11,36} To examine the oxidation potential of oxygen atoms in a CeO₂ island supported on a metal surface, we constructed CeO₂/Au(111) models (see Figure 2d,e) and calculated the E_{vac} of oxygen atoms of the CeO₂ island at the Au-CeO₂ interface. The E_{vac} of the Ce₅O₉/Au(111) model depends on the location of oxygen atoms varying between 0.95 to 1.32 eV (Table S3). Of course, CeO₂/Au(111) inverse-catalysts could activate CO oxidation by a different mechanism. However, the average E_{vac} of interfacial oxygen atoms (1.19 eV, which is lower than the

threshold E_{vac} (2.10 eV) for the activation of the M-vK mechanism of CO oxidation derived from CeO₂-steps) predicts the strong oxidation potential of these oxygen atoms. The calculated E_{vac} of the larger CeO₂ island, Ce₉O₁₈/Au(111), was increased from that of the Ce₅O₁₀/Au(111) (average E_{vac} = 1.62 eV, see Table S3), predicting that the E_{vac} of the supported CeO₂ of the CeO₂/Au inverse catalyst is highly sensitive to the shape and size, suggesting that the dimension of CeO₂ islands would affect their catalytic activity. We found that Ce atoms in Ce₅O₁₀/Au(111) were reduced as the CeO₂ island acquired extra electron density from Au supports (see Table S4). We will address the size effect of CeO₂ islands on the oxidation catalysis and the detailed mechanism of oxidation catalysis.

Herein, we study various kinds of CO oxidation mechanisms at the interface of the Au NCs supported on the CeO₂-step and find that Au–CO* can be directly oxidized by the protruded oxygen atoms of the CeO₂-step by the M-vK mechanism of oxidation. CO oxidation by the M-vK mechanism at the interface between the CeO₂ support and supported Au NPs/NCs is highly beneficial for the consistency in the catalytic activity because the Au NPs/NCs and CeO₂ are playing different roles without competing for the same binding site; Au and CeO₂ supplies CO and oxygen, respectively. The low activation energy and CO₂ production energy of the Au/CeO₂-step give rise to the high CO oxidation activity of the Au/CeO₂-step, and the E_{vac} is a reactivity descriptor. The maximum E_{vac} that insures spontaneous CO₂ production is higher for the Au/CeO₂-step than that for the Au/CeO₂-surface, suggesting that the CeO₂-step is a better supporting material than the CeO₂-surface for CO oxidation by the Au/CeO₂. Our results also suggest that for CO oxidation by Au NCs supported on nano- or meso-structured CeO₂, which is the case of industrial catalysts, the M-vK mechanism accounts for a large portion of the total activity.

Our previous study and recent report by Bruix et al.¹⁴ shows that doping the CeO₂ surface⁹ or making a mixed (Ce–X)O₂ oxide¹⁴ would query the participation of lattice oxygen atoms of CeO₂ surface in CO oxidation at the interface between CeO₂ surface and supported metal NPs/NCs. Moreover, current results show that the catalytic activity of the CeO₂-metal NPs/NCs interface can be modified by the dimension of the CeO₂ support. We suggest that combining chemical and morphological optimization of the oxide support is a promising way to extensively modify the activity of oxidation catalysis by supported Au NPs/NCs.

■ ASSOCIATED CONTENT

■ Supporting Information

Additional data is presented in Tables S1, S2, S3, and S4, and Figures S1 and S2. 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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