Growth, Structure, and Stability of Ag on CeO$_2$(111): Synchrotron Radiation Photoemission Studies

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ABSTRACT: The growth and interfacial electronic properties of Ag on CeO$_2$(111) thin films have been studied by synchrotron radiation photoemission spectroscopy (SRPES), low energy electron diffraction (LEED), and X-ray photoelectron spectroscopy (XPS). Stoichiometric CeO$_2$(111) thin films were grown on a Ru(0001) substrate. Ag grows as three-dimensional (3D) particles on the well-ordered CeO$_2$(111) surface at 300 K with a number density of $\sim 1 \times 10^{12}$ particles/cm$^2$. When the CeO$_2$(111) surface has a high density of defects, Ag initially populates these defect sites, leading to a two-dimensional (2D) island growth at low coverages followed by 3D islanding at high coverages. The binding energy of Ag 3d increases when the Ag particle size decreases, which is mainly attributed to the final-state screening. No strong interaction between Ag and CeO$_2$(111) is found. The CeO$_2$(111) surface is slightly reduced upon Ag deposition, which can be ascribed to the reverse spillover of oxygen atoms from the Ag–CeO$_2$ boundary to the Ag nanoparticles. The Ag particles on CeO$_2$(111) experience significant sintering when the temperature increases before they desorb from the surface.

1. INTRODUCTION

Metal-on-oxide systems have received considerable attention because of their extreme importance in both fundamental research and technological applications, for instance, in microelectronics devices, gas sensors, and heterogeneous catalysts.$^1$–$^3$ Interface properties referring to the interfacial diffusion, charge transfer, hybridization, morphology, and reactivity in the metal–oxide systems play crucial roles in determining their applications.$^4$–$^{11}$In heterogeneous catalysis, ceria is one of the most important supports for metal catalysts, mainly the late transition metals, that have been widely used in a variety of catalytic reactions, such as CO and hydrocarbon oxidation, low-temperature water–gas shift reaction, automotive exhaust emission-control reactions, and low-temperature methanol synthesis.$^{12}$–$^{17}$ To gain fundamental insight into the interaction of metals with the CeO$_2$ support as well as their catalytic properties, substantial fundamental surface science studies in terms of the growth, structure, and reactivity of vapor-deposited metal nanoparticles on ordered CeO$_2$(111) surfaces have been performed.$^{18}$–$^{25}$ It is thought that the superb catalytic properties of the ceria-supported metal catalysts arise from: (1) the excellent redox properties of ceria (ceria is easily reduced and has high oxygen storage capacity)$^{14,24}$ and (2) the stronger interactions of ceria with supported metal nanoparticles compared to other oxides, which can maintain smaller metal nanoparticles and resist the small particles to sinter into large particles under catalytic reaction conditions.$^{11}$ This is related to the easy formation of defects, such as oxygen vacancies, on the ceria surfaces.

Among all the late transition metals, Ag is attractive for industrial applications due to its low cost and easy preparation. Ag supported on CeO$_2$ has been reported for use as electric contact materials$^{29}$ and in solid-oxide fuel cells.$^{30}$ In addition, it can be used as a catalyst in NO reduction, CO and hydrocarbon oxidation,$^{31}$ methane oxidation,$^{32}$ and formaldehyde oxidative decomposition.$^{33}$ Moreover, the use of Ag deposited on ceria was also found to increase the rate of carbon gasification in the oxidation of carbon soot particles compared to other noble metals.$^{34}$–$^{37}$ Despite these tremendous applications of the Ag/ceria system, the nature of the Ag–ceria interfaces in terms of interfacial morphologies, electronic/chemical properties, and interaction of Ag with ceria is still far from well understood. Recently, Farmer at al. employed adsorption microcalorimetry in combination with other techniques such as low-energy ion scattering (LEIS), Auger electron spectroscopy (AES), and low-energy electron diffraction (LEED) to study the adsorption energies and growth of Ag on the reduced CeO$_2$($x$(111) ($x = 0.1$–$0.2$) thin films.$^{38,20}$ A three-dimensional (3D) growth mode with a density of $\sim 4 \times 10^{12}$ particles/cm$^2$ for Ag on reduced CeO$_{1.9}$(111) was reported. The sticking probability of Ag on all reduced ceria surfaces was found to be near unity. In addition, the

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higher heat of adsorption of Ag on the more reduced surfaces was attributed to stronger bonding of Ag particles to oxygen vacancies. Moreover, from the point of adsorption energetics they quantitatively explained why Ag has larger sinter resistance on CeO$_2$-(111) than MgO(100), since it bonds much more strongly to CeO$_2$-(111). However, since they were not able to prepare the stoichiometric CeO$_2$(111) thin films, no results of Ag on CeO$_2$(111) were reported. Furthermore, to our knowledge, no studies of temperature-dependent thermal stability of Ag nanoparticles on CeO$_2$ can be found in the literature.

In this study, we use synchrotron radiation photoemission spectroscopy (SRPES) together with X-ray photoelectron spectroscopy (XPS) and LEED to investigate the growth, nucleation, and electronic properties of Ag on the ordered CeO$_2$(111) thin films which were prepared in situ on a Ru(0001) substrate. The advantage of the tunability of the photon energies enables us to investigate every core level and valence band with high sensitivity. In particular, in the case of a very low concentration of reduced state Ce$^{3+}$, from the Ce 3d spectra it is hard to determine the extent of sample reduction since the changes in the spectra are difficult to quantify without extensive signal averaging. In contrast, valence band and Ce 4d spectra are very sensitive to the reduction of CeO$_2$. From the core-level binding energies and Auger parameters, the electronic properties of the Ag/CeO$_2$ interface with various sizes of Ag nanoparticles are determined. The thermal stability/sintering process of the Ag nanoparticles with different initial sizes at elevated temperature are also studied.

2. EXPERIMENTAL SECTION

All the experiments were performed at the Photoemission Endstation at beamline U20 in the National Synchrotron Radiation Laboratory, Hefei, China. This beamline is connected to a bending magnet and equipped with three gratings that cover photon energies from 60 to 1000 eV with a resolving power ($E$/$\Delta E$) better than 1000. The endstation consists of an analysis chamber and a sample preparation chamber, whose base pressures are $2 \times 10^{-10}$ and $5 \times 10^{-10}$ Torr, respectively, and a sample load-lock system. The analysis chamber is equipped with a VG Scienta R3000 electron energy analyzer, a twin anode X-ray source, rear-view LEED optics, a quadrupole mass spectrometer (Pfeiffer QMS220), and an Ar$^+$ sputter gun. The preparation chamber is connected with a quick load-lock port and houses an e-beam evaporator for Ce and a home-built evaporator for Ag.

The photoemission spectra were recorded at normal emission and have been normalized to the photon flux which was calibrated by measuring the Au 4f spectra from a clean Au foil attached to the sample holder. The following photon energies have been used to probe the different core levels: Ru 3d, 650 eV; Ce 4d, 450 eV; Ag 3d, 650 eV; O 1s, 650 eV. Valence band spectra were recorded with a photon energy of 170 eV. Binding energies (BEs) were calibrated with respect to the Au 4f$_{7/2}$ (BE = 84 eV) feature from the Au foil, which was measured immediately after each spectrum.

The Ru(0001) single crystal (10 mm diameter and 2 mm thickness) sample, purchased from MaTeck GmbH, Germany, was used as the substrate. The sample was mounted on a Mo sample holder which was fixed by Ta wires at the side. A K-type thermocouple was used to monitor the sample temperature. The sample was cleaned by repeated cycles of Ar$^+$ bombardment at room temperature followed by annealing up to 1300 K until no contaminants were detected by XPS and a sharp LEED pattern was observed.

CeO$_2$(111) thin films were prepared following a recipe similar to that reported previously. Briefly, the epitaxial grown ordered CeO$_2$(111) films were produced by evaporating cerium metal in $2 \times 10^{-7}$ Torr oxygen environments onto Ru(0001) with a constant substrate temperature of 700 K. To promote film ordering to form large flat terraces, the films need to be postannealed at 980 K in oxygen. In this study, the CeO$_2$(111) films after growth were postannealed at 980 K in oxygen, unless otherwise specified. Metallic cerium (99.999%, Goodfellow) was evaporated from an electron bombarded W crucible using a water-cooled e-beam evaporator that was equipped with an integral flux monitor, enabling real-time control of the deposition rate. The growth rate of the CeO$_2$ films was estimated, from the attenuation of the Ru 3d signal at a photon energy of 650 eV, to be $0.089 \pm 0.003$ nm/min by assuming the electron mean free path of 1 nm. The thickness of the CeO$_2$(111) films is about 4 nm. With this thickness, the long-range electronic interaction effect with the underlying Ru(0001) substrate can be neglected, as reported by Farmer et al. One CeO$_2$ monolayer (ML) is defined as $7.89 \times 10^{-14}$ CeO$_2$ units per cm$^2$, which is the number of oxygen atoms per unit area in the topmost atomic layer of the CeO$_2$(111) surface (which is terminated in an open oxygen layer with a layer of coordinatively-ununsatured Ce$^{4+}$ ions below). Dividing this by the density of bulk ceria (2.53 $\times 10^{22}$ CeO$_2$ units per cm$^3$) gives the thickness of a CeO$_2$ monolayer to be 0.31 nm. Ag was deposited onto the ceria films by evaporating Ag metal (99.999%, Alfa Aesar) from a tantalum basket, which was thoroughly degassed prior to use. The deposition rate of Ag, estimated from the attenuation of Ru 3d signals at a photon energy of 650 eV after Ag was directly deposited onto Ru(0001), was $0.069 \pm 0.002$ ML/min. One monolayer (ML) of Ag is defined throughout as the number of Ag atoms per area in a closed-packed layer (111) of Ag atoms with lattice parameter of bulk Ag (1 ML = 1.4 $\times 10^{15}$ atoms/cm$^2$), which is about twice the number of oxygen atoms in CeO$_2$(111) exposed to the vacuum per unit area.

3. RESULTS

3.1. Characterization of the CeO$_2$(111) Films. Figure 1 shows the Ce 4d and valence band (VB) spectra for a 4 nm thick fully oxidized CeO$_2$(111) film grown on Ru(0001), along with spectra from a typical partially reduced CeO$_2$$_{-x}$ film for comparison. The reduced CeO$_2$$_{-x}$ film was obtained by annealing the CeO$_2$ film at 1000 K for 10 min. Both Ce 4d and VB spectra from the oxidized CeO$_2$(111) surface are very similar to those reported previously. In the Ce 4d region, the two peaks located at 125.2 and 121.9 eV, respectively, are only related to the presence of Ce$^{4+}$. They vanish when the surface is fully reduced to CeO$_2$. There are no peaks in the CeO$_2$$_{-x}$ region between the Ce metal (99.99%, Goodfellow) and CeO$_2$(111). Therefore, the intensities of these two peaks can be directly used to calculate the extent of reduction assuming a linear relationship of the changes in intensities of these two peaks against the relative amount of Ce$^{4+}$. In the valence band region, only the contribution from O 2p is observable. The absence of a Fermi edge is indicative of neither metallic Ce nor substrate Ru being visible on the surface. For the thermally reduced CeO$_2$$_{-x}$ ($x = 0.15$) film the intensities of the peaks at 125.2 and 121.9 eV in the Ce 4d spectrum are only 70% of those for the CeO$_2$ film, implying that ~30% of the Ce$^{4+}$ has been reduced to Ce$^{3+}$ on the surface. The appearance of the
The (0,0) spot is near the center of the screen. In (b), the unit cells of CeO$_2$(111) and Ru(0001) are indicated.

CeO$_2$(111)

Ce$_3$ of surface with the approximate surface stoichiometry of CeO$_{1.85}$. The Ru(0001). When the CeO$_2$ further intensify at the expense of the p(1$	imes$1/C$_{2}$/C$_{2}$) LEED pattern, as shown in Figure 2(a).

After growth of 1 nm (3 ML) of CeO$_2$, in addition to the original spots from Ru(0001), some new spots with smaller distance to the (0,0) spot compared to the original first-order spots of Ru(0001) are clearly observed as a hexagonal structure. This is the well-known p(1.4$	imes$1.4) LEED pattern of CeO$_2$(111) with respect to the p(1$	imes$1) pattern of clean Ru(0001) (shown in Figure 2(b)), as expected based on prior observations. With increasing thickness of the CeO$_2$ film, the p(1.4$	imes$1.4) spots further intensify at the expense of the p(1$	imes$1) spots of Ru(0001). When the CeO$_2$ film reaches 2 nm thickness, the LEED pattern from the Ru substrate can no longer be observed, and only the pattern from the CeO$_2$(111) film is observed. Shown in Figure 2(c) is a sharp p(1.4$	imes$1.4) LEED pattern which corresponds to a (1$	imes$1) pattern of CeO$_2$(111) obtained from a 4 nm thick CeO$_2$(111).

3.2. Growth of Ag on CeO$_2$(111) at Room Temperature.

The dispersion, nucleation, and wetting properties of metals on oxides are of great interest in heterogeneous catalysis. Study of the growth mode of ultrathin metal films on well-defined oxide surfaces provides important insights into the nature of the bond between metal and oxide surfaces. STM studies revealed that on the ordered CeO$_2$(111) films the deposition of Au, Pd, Rh, and Pt at 300 K follows a three-dimensional (3D) growth mode. At low Au coverages (<0.1 ML), Weststrate et al. observed that Au prefers to grow two-dimensionally on CeO$_2$-(111). In the present study, SRPES was used to investigate the growth mode of Ag on CeO$_2$(111) by monitoring the intensity changes of Ag 3d and O 1s. We have studied the growth of Ag on two different 4 nm thick CeO$_2$(111) films with different roughness. The two CeO$_2$(111) films were prepared with and without postannealing at higher temperature (980 K) in oxygen environments, respectively. According to the STM studies, due to large lattice mismatch (40%) between CeO$_2$(111) and Ru(0001), smooth and well-ordered CeO$_2$ films can be only obtained by further annealing at higher temperatures. The ceria films become better ordered and have much larger terraces after postannealing in oxygen at higher temperatures. The CeO$_2$(111) film without postannealing shows particle-like features and, of course, has higher concentration of defects (step edges and kinks). However, from the conventional LEED observations, the differences in the morphology of these two films can hardly be discriminated, in line with previous literature results. In addition, both CeO$_2$ surfaces should have low density of point defects (oxygen vacancies) on the terraces.

As shown in Figure 3, the solid points represent the data from the CeO$_2$(111) surface without postannealing, while the hollow data points were obtained from the surface with postannealing. On both surfaces, the normalized intensity changes of the Ag 3d$_{5/2}$ and O 1s peaks are plotted as a function of Ag coverage, respectively. Here, the intensities of Ag 3d$_{5/2}$ and O 1s are normalized to those of bulk-like Ag and clean CeO$_2$(111), respectively. As can be seen, on both CeO$_2$(111) surfaces the intensity of the Ag 3d$_{5/2}$ peak increases continuously with increasing Ag coverage, and simultaneously that of the O 1s peak decreases. To clarify the growth mode, we calculated the expected results for both a layer-by-layer growth mechanism and a simple 3D-particle (hemispherical cap) growth model, shown as the dashed lines and the solid curves, respectively. The 3D growth model assumes that the Ag particles have hemispherical shapes and constant island density with uniform radii. The island density is independent of coverage and treated as a fitting parameter (here we use the best fitted island densities of 4 × 10$^{12}$ and 1 × 10$^{13}$ particles/cm$^2$ for the CeO$_2$(111) films without and with postannealing at higher temperature in oxygen, respectively). Therefore, at any given Ag coverage the average particle volume and diameter can be calculated. For example, at a coverage of 1 ML, the average
particle size is calculated to be 3.0 nm in diameter on the CeO$_2$(111) films without postannealing at higher temperature in oxygen.

As seen in Figure 3, on the CeO$_2$(111) surface without postannealing during the early stage of Ag growth (<0.3 ML), the Ag signal changes along a slope very close to that expected for a layer-by-layer growth mode, and the solid curves represent a hemispherical cap model with constant island densities (4 $\times$ 10$^{12}$ and 1 $\times$ 10$^{12}$ islands/cm$^2$ for the CeO$_2$(111) films without and with postannealing at higher temperature in oxygen, respectively). The inelastic mean free path used for calculating the curves is 0.61 nm for Ag and 0.5 nm for O, obtained from ref 52, and the detection angle was 0$^\circ$.

In contrast, on the CeO$_2$(111) surface with postannealing, the O signal decreases with a slope also in accord with that of the expected 2D growth. As the Ag coverage increases to above 0.3 ML, both the substrate and overlayer signals deviate from the layer-by-layer growth mode and rapidly exhibit behavior indicative of 3D island growth with an island density of $\sim$4 $\times$ 10$^{12}$ islands/cm$^2$. Even at the coverage above 6 ML, the O signal can be still clearly seen, implying that a large fraction of the substrate area is still exposed; this is clear evidence for the 3D growth of Ag on the surface. This 2D–3D transition in the growth mode of Ag on CeO$_2$(111) is similar to those reported for Ag/TiO$_2$, Ag/SiO$_2$, and Ag/MgO systems$^{53-56}$ but in contrast to that of Ag on reduced CeO$_2$–x(111) where only 3D growth was found.$^{29}$

In contrast, on the CeO$_2$(111) surface with postannealing both the Ag 3d and O 1s signals indicate that the growth of Ag exhibits 3D behavior during the initial growth stages. In addition, the particles have smaller island density ($\sim$1 $\times$ 10$^{12}$ islands/cm$^2$) in comparison with those on the surface without post-annealing. This means that at a given Ag coverage the mean particle sizes are larger on the better ordered CeO$_2$(111) surface than on the rough CeO$_2$(111) surface. In other words, when the CeO$_2$(111) surfaces have more defects the Ag particles grow smaller and are better dispersed, as we observed. Similar results have been found by Weststrate et al.$^{25}$ for Au on CeO$_2$(111) where Au grows with smaller average diameters on the rough CeO$_2$ surface compared with the smooth surface. It is common that metal nucleates at the defect sites of the oxide surfaces at very low coverages, undergoing a 2D growth.$^{11,22,26,44,51,57}$ Since the CeO$_2$(111) surfaces have low density of point defects, i.e., oxygen vacancies,$^{44}$ it is very likely that Ag preferentially nucleates on the defect sites such as step edges and kink sites initially. After these defect sites are populated, Ag starts to grow three-dimensionally on the surface. On the better ordered surfaces, the density of the defects is much lower, thus the nucleation sites are much less for Ag growth. Therefore, the island density is much lower, and at the same coverage the Ag particle sizes are much larger.

It is of interest to compare the growth mode we derive from experimental data with that predicted from thermodynamic parameters. As we know, if the interfacial energy ($\gamma_{m/o}$) is smaller than the difference between oxide surface energy ($\gamma_{o/o}$) and deposited metal surface energy ($\gamma_{m/m}$), i.e., $\gamma_{m/o} < \gamma_{o/o} - \gamma_{m/m}$, the metal is energetically favored to wet the oxide surface. Otherwise, 3D growth occurs. The criterion can also be expressed in terms of metal–oxide adhesion energy, $E_{a/d}$. If it exceeds the adhesion energy of metal–metal, one would expect that the metal will wet the oxide surface and form a continuous film. Since the adhesion energy of Ag nanoparticles to the reduced CeO$_2$(111) surface is $\sim$2.3 J/m$^2$ and becomes even larger on more reduced surfaces,$^{11}$ it is expected that the adhesion energy of Ag nanoparticles to the oxidized CeO$_2$(111) surface is smaller than 2.3 J/m$^2$. This value is obviously less than the adhesion energy of Ag to itself (2.44 J/m$^2$)$^{11,58}$ therefore, one would expect that the growth of Ag on CeO$_2$(111) follows 3D mode, consistent with our results. However, on the rough CeO$_2$(111) surface where the concentration of defects (step edges and kinks) is higher than that on a better ordered surface, a Ag particle bonds locally to these defect sites with an adhesion energy which is likely much more than the stoichiometric CeO$_2$ terraces. This would result in Ag being more thermodynamically favored to form 2D islands in the initial stages of Ag growth.

Note that although the growth models are strongly supported by both the experimental results and thermodynamic prediction it would be helpful to further confirm these by STM observations which provide more direct structural/morphological information.

3.3. Interfacial Electronic Properties. The electronic, optical and magnetic properties of metal nanoparticles supported on oxide supports can diverge strongly from those of the bulk materials.$^{59}$ The interactions between metal particles and oxide supports have been widely studied using photoelectron spectroscopy. When metal particles are deposited on oxide surfaces, binding energy shifts in core-level spectra are often observed as a function of particle size, which can be interpreted as being due to either initial$^{60-62}$ or final state effects or both.$^{53,64}$ In general, initial state effects can be attributed to reduced metal–metal coordination and/or metal-oxide interaction, while final state effects arise from final state screening, i.e., relaxation effects, which depend on the shape and size of the particles as well as the supports.$^{51,65}$ However, separating the relative contributions of these two effects on the core-level binding energy shifts is an issue.

Here we use the Auger parameter method which was first introduced by Wagner$^{66}$ to shed light onto the origin of the particle-size-dependent Ag 3d level shifts. In Figure 4a and 4b, the Ag 3d$_{5/2}$ core level binding energy and Ag M$_{3,5}$VV Auger transition kinetic energy are plotted as a function of Ag coverage
at 300 K. As can be seen, with decreasing Ag coverage, the Ag 3d5/2 binding energy shifts to a higher BE, while the kinetic energy (KE) of the M5VV Auger transition shifts to a lower KE. At 0.04 ML Ag coverage, with respect to the bulk Ag value, the Ag 3d BE is higher by 0.6 eV and the M5VV Auger peak position is lower by 1.5 eV. Above 10.0 ML Ag coverage, they all approach the bulk Ag value.

It was previously shown by Wagner that the sum of the kinetic energy shift of the Auger peak and the binding energy shift of the core level peak approximately equals twice the final state contributions to the core level shifts.\(^{66-68}\) This can be rationalized only when the energy shifts of different orbitals are the same. For Ag clusters grown on alumina surfaces, Luo et al.\(^{69}\) have demonstrated the validity of the Wagner analysis by the observation of parallel shifts of the Ag 3d peak and the center of the Ag 4d band. The Auger parameter (\(\alpha\)), which is currently defined as the sum of the Ag 3d5/2 core level binding energy (shown in Figure 4a) and the kinetic energy of the M5VV Auger transition (shown in Figure 4b), is plotted as a function of Ag coverage in Figure 4c. The Auger parameter increases with increasing Ag coverage and shows opposite shifts from the Ag 3d core levels. The value of \(\alpha\) increases from 719.6 to 721.6 eV with increasing Ag coverage from 0.04 ML to the bulk value. As mentioned above, the final state contribution (\(\Delta R\)) to the Ag 3d5/2 core level BE shifts can be written as

\[
\Delta R = \Delta \alpha / 2
\]

The shift in the Auger parameter (\(\Delta \alpha\)) can be estimated from the data shown in Figure 4c. The initial state contribution (\(\Delta \epsilon\)) can be calculated by

\[
\Delta \epsilon = \Delta BE + \Delta R
\]

The resulting values for \(\Delta R\) and \(\Delta \epsilon\) are plotted as a function of Ag coverage in Figure 4d. It is obvious that over the entire coverages we investigated the final state contribution (\(\Delta R\)) to the Ag 3d core level binding energy shifts is dominant. At very low coverage (0.04 ML), the final state effects contribution is calculated to be \(\sim 0.5\) eV, while the initial state contribution (\(\Delta \epsilon\)) is about \(\sim 0.1\) eV. When the Ag coverage increases to \(\sim 0.2\) ML, the initial state contribution becomes negligible, and only the final state contribution accounts for the Ag 3d BE shifts.

It is known that the final state effects originate from the screening and relaxation of the core hole created by photoemission, which depends on the shape and size of the metal particles as well as on the substrate. Due to the poor conductivity and coupling of the substrate with the metal particle, the hole created by photoemission cannot be compensated by the transfer of a screening electron from the substrate to the metal particle. As a result, the kinetic energy of escaping photoelectron is lowered due to the Coulomb interaction between the escaping electron and the remaining hole, leading to the observed BE shifts. The initial state effects arise from the electron charge transfer between metal and substrate and/or intrinsic size effects due to reduced metal—metal coordination in the metal particles. Since the direction and the value of the initial state shifts are consistent with the surface core level shifts (SCLS) of Ag (\(\sim 0.15\) eV for Ag(111)\(^{70}\)), this initial state contribution can be attributed to a reduced average coordination number for Ag atoms in small particles. Electron charge transfer from Ag to CeO\(_2\) resulting in the formation of positively charged Ag is unlikely because, although the oxidation of Ag would lead to negative BE shifts of the Ag 3d core level relative to the metallic Ag state, a much larger BE shift is expected if Ag is positively charged (for Ag\(_2\)O and AgO, \(\sim 0.4\) and \(\sim 0.8\) eV shifts relative to metallic Ag 3d were observed, respectively.\(^{71}\)). On the other hand, since Ag bonds more strongly on the defect sites, positively charged Ag would be even more easily observed. However, on the CeO\(_2\)(111) film without postannealing, we found that the initial state effect contribution to the Ag 3d BE shifts during Ag growth is essentially very small and can be negligible. Therefore, it is conclusive that charge transfer between Ag and CeO\(_2\) does not occur to a measurable extent during the Ag growth on CeO\(_2\)-(111). In other words, Ag particles maintain metallic states on the
CeO$_2$(111) surface. Similar results have been reported for Ag on Al$_2$O$_3$/Re(0001).99,72 However, our results are in contradiction to previous results for 1% Ag/CeO$_2$73 and density functional theory (DFT) calculations for Ag on CeO$_2$(111),18 where positive-charged ionic Ag$^+$ was reported. It is worth pointing out that from pure XPS studies it is difficult to distinguish between Ag$^+$ and metallic Ag$^0$ due to the lower binding energy shift of Ag$^+$ compared to Ag$^0$ from the Ag 3d core level only, particularly when the particle size is in the nanometer scale.

Figure 5 shows the valence band spectra for Ag deposited on CeO$_2$(111) at different Ag coverages. The main feature of the spectrum from the clean substrate (bottom curve) is the broad peak ranging from 3 to 8 eV which can be ascribed to O 2p structure.45 With increasing Ag coverage, this broad peak shifts to the higher binding energy side due to the coupling of Ag states. In addition, a peak located at $\approx$1.5 eV below the Fermi level ($E_F$), which is attributed to Ce 4f,45 gradually increases in intensity. As mentioned above, this peak is only related to the reduced state Ce$^{3+}$. The appearance of this state as well as its gradual increase in intensity, as can be clearly seen in the inset of Figure 5, indicates that the CeO$_2$ surface is reduced upon Ag deposition.

Moreover, the density of state (DOS) near $E_F$ gradually increases with increasing Ag coverage. For comparison, the valence band spectrum from a bulk-like Ag film (>40 ML) is shown as the dashed curve in Figure 5, where a prominent feature at 4.8 eV and a second feature centered at 6.3 eV, as well as a sharp Fermi edge (seen in inset of Figure 5), are observed. The features at 4.8 and 6.3 eV can be attributed to the 4d band state of metallic Ag.74,75

It is worth mentioning that in previous studies a new state at $\approx$1.2 eV near $E_F$ was observed with increasing Ag cluster size on amorphous graphite, which was primarily ascribed to the 5s state since Ag has a nearly filled 4d state.75 However, this 5s state was only observable when the Ag coverage was larger than 8.5 $\times$ 10$^{15}$ atoms/cm$^2$ (in our case it corresponds to 6 ML). It primarily appeared at around 2.1 eV for small particles.76 In our experiment, we can rule out the possibility of the increase in Ce 4f intensity induced by the appearance of the Ag 5s state since at very low Ag coverage (<0.4 ML) where the Ag 5s state should have negligible intensity the increase in Ce 4f intensity has already been observed, and moreover, there is no shift of the Ce 4f peak position. Therefore, it is clear that CeO$_2$ has been reduced after Ag deposition. Further evidence can also be provided by the observation of Ce 3d core-level spectra. Figure 6 shows the Ce 3d XPS spectra acquired from a clean CeO$_2$(111) surface as well as the surface covered by 0.5 ML Ag. The spectra exhibit evident changes after Ag deposition. Decomposition of the spectrum for 0.5 ML Ag covered CeO$_2$(111) is presented at the bottom of Figure 5. The appearance of new spectral components at 885.0 and 880.5 eV is clear evidence of the formation of Ce$^{3+}$ ions.40,77

The detailed mechanism for the reduction of CeO$_2$ upon Ag deposition will be discussed later. Here we first rule out the possibility of the reduction being caused by photon-induced desorption of oxygen atoms from the CeO$_2$(111) surface through a control experiment. We performed this experiment on a 0.1 ML Ag covered CeO$_2$(111) by measuring the valence band spectra with a different dosage of synchrotron radiations. The results show no difference in the valence band spectra between the fresh-prepared sample and the sample after exposing it for a long time to synchrotron radiations, indicating that the formation of the Ce$^{3+}$ state does not result from photon-induced desorption of O atoms from CeO$_2$(111).

3.4. Thermal Sintering. For Au, Pd, and Pt on CeO$_2$(111),19,22,78 it has been reported that annealing in UHV leads to significant sintering of metal particles on the surface. Even heating up to 500 K, Au nanoparticles experience an increase in particle size on CeO$_2$(111).10,21 To determine the thermal stability of Ag nanoparticles on CeO$_2$, we have carried out the following experiments.

Two different coverages of Ag deposited on CeO$_2$(111) at 300 K were annealed at elevated temperatures, after which Ag 3d spectra were collected at 300 K. In Figure 7(a), the Ag 3d$\alpha$/$\gamma$ peak intensities are plotted as a function of temperature for Ag/
CeO$_2$(111) samples with 0.05 and 0.5 ML Ag coverage, respectively. As expected, the intensity of the Ag 3d$_{5/2}$ peak for 0.5 ML Ag on CeO$_2$(111) at 300 K is much higher than that of 0.05 ML Ag. Correspondingly, the particle size is also much larger for 0.5 ML Ag. With increasing annealing temperature, the Ag 3d$_{5/2}$ intensity for 0.05 ML Ag/CeO$_2$(111) starts to decrease already at 400 K, while for 0.5 ML the Ag 3d$_{5/2}$ intensity remains nearly unchanged up to 500 K. When the temperature is higher than 500 K, the Ag 3d$_{5/2}$ intensities from both surfaces decrease rapidly. At about 900 K, no Ag signal can be detected.

In principle, there are three possible reasons which account for the decrease of Ag 3d$_{5/2}$ intensity during annealing: (i) small Ag particles agglomerate into large particles, i.e., sintering; (ii) Ag diffuses into the subsurface; and (iii) Ag desorbs from the surface. For metal supported on oxide surfaces, the diffusion of metal atoms into the subsurface of oxide has rarely been reported. Moreover, Ar-ion sputtering depth profile experiments demonstrate that there is no Ag which can be detected after the surface Ag particles are removed. Therefore, the diffusion of Ag into the CeO$_2$(111) subsurface can be ruled out. Desorption of Ag from the CeO$_2$(111) surface before 800 K is also unlikely because even for a weak Ag-interacted substrate such as Al$_2$O$_3$ temperature-programmed desorption (TPD) results demonstrated that Ag desorbs only above 800 K. Since Ag interacts more strongly with CeO$_2$ than Al$_2$O$_3$, the desorption temperature for Ag on CeO$_2$(111) must be 800 K or higher. Therefore, desorption temperature for Ag on CeO$_2$(111) below 800 K is unreasonable.

Thus, the decrease in intensity of Ag 3d$_{5/2}$ spectra during annealing before 800 K can be attributed to particle sintering on the surface. Above 800 K, the peak intensity starts to decrease more steeply due to the Ag desorption. By 900 K, all Ag desorbs from the surface, leading to no detectable Ag signal by XPS. The initial sintering temperature for a low coverage Ag film (0.05 ML) is lower than that for a high coverage film (0.5 ML). This is understandable since the initial particle size is much smaller. Moreover, desorption also completes at lower temperature for a lower coverage. These results are consistent with the fact that small particles desorb more easily than “infinite” particles.

That Ag particles undergo significant sintering on CeO$_2$(111) before desorption is also confirmed by the observation of the shifts of Ag 3d$_{5/2}$ binding energy as well as by the changes of the full width at half-maximum (fwhm) of the Ag 3d$_{5/2}$ peak. Figure 7(b) displays the binding energy of Ag 3d$_{5/2}$ peaks as a function of annealing temperature for 0.05 and 0.5 ML Ag on CeO$_2$(111), respectively. As seen, for both Ag coverages, the binding energy of the Ag 3d$_{5/2}$ peak shifts to the lower BE side during annealing. This shift is more apparent for 0.05 ML Ag than for 0.5 ML since the initial particle size is smaller in the former case. In addition, the fwhm (not shown) of the Ag 3d$_{5/2}$ peak decreases during annealing for both coverages. Along with the previous description, the phenomena associated with the changes of BE and fwhm of Ag 3d$_{5/2}$ during annealing are clearly caused by the size effects. Hence, it provides clear evidence for Ag sintering on the CeO$_2$(111) surface before desorption.

To further prove Ag sintering at elevated temperatures, we performed an experiment of Ag deposition at 500 K. The results show that the increase of Ag signals initially follows the 3D growth mode with a lower number density of particles (<1 × 10$^{12}$ particles/cm$^2$, assuming hemispherical cap growth) than 300 K. This implies that the average particle size at 500 K is larger than at 300 K at a given Ag coverage. Correspondingly, at 500 K a smaller fraction of the substrate area is covered by Ag than at 300 K. However, we cannot rule out the possibility of the lower number of particle density at 500 K caused by a lower sticking probability of Ag on CeO$_2$(111) as compared to that at 300 K. Similar dependence on the substrate temperature has been observed previously for Cu$^{50}$ and Au$^{79–81}$ grown on TiO$_2$. Considering the kinetic effects, as the substrate temperature increases, since Ag atoms become more mobile on the surface and some of the kinetic barriers may be surmounted, the Ag particles grow in size by a combination of Ostwald ripening (growth of larger particles at the expense of smaller ones) and coalescence (particles merge together).

4. DISCUSSION

Our results provide clear evidence that the CeO$_2$(111) surface has been slightly reduced upon Ag deposition. Similar phenomena have also been observed in the growth of Pd, Pt, and Au on ceria.$^{24,28,36}$ However, regarding the mechanism of metal-induced reduction of the CeO$_2$ films, it is still under dispute.$^{24,27,42,77,82}$ Summarized from the literature, there are several explanations for metal-induced reduction of CeO$_2$: (1) charge transfer from the supported metal particles to CeO$_2$; (2) reverse spillover of lattice oxygen to the surface of the supported metal nanoparticles; (3) metal-catalyzed desorption of surface oxygen atoms; and (4) metal-catalyzed reactions of O atoms from CeO$_2$ with background CO gas to make CO$_2$. For the first explanation, for example, Wilson et al.$^{77}$ reported that for Pd/ CeO$_2$ the CeO$_2$ surface was reduced by electron transfer from the Pd adatoms to the CeO$_2$ surface. Recently, Skoza et al.$^{42}$ proposed that the formation of a Au–Ce–O gold–ceria interface caused ceria reduction. This finding is consistent with the theoretical study by Shapovalov et al.$^{83}$ but disagrees with the
experimental results of Baron et al.,26 which strongly suggest that the oxidation state of Au atoms adsorbed on the perfect CeO2-(111) single-crystal surface is essentially zero. The reduction of ceria caused by the reverse spillover of lattice oxygen to the surface of the supported metal nanoparticles can be found in many literatures.18,22,27,38,84–88 For example, Zhdanov and Kasemo88 reported that the energy of oxygen atoms on Pt nanoparticles is essentially very close to that in the surface layer of the CeO2. The activation barrier for reverse spillover from ceria to Pt over the Pt–CeO2 boundary is rather low (10 kcal/mol). Experimental evidence for reverse oxygen spillover on the same system has been reported by Lykhach et al.27 They observed the lattice oxygen migration to the Pt particles while annealing the sample in ultrahigh vacuum (UHV) at temperatures between 400 and 500 K. Other experimental evidence for the oxygen reverse spillover from CeO2 to supported metal nanoparticles can be found in the case of Rh/CeO2,84,85,87 and Pd/CeO2.82 The reduction of CeO2 caused by metal-catalyzed desorption of surface oxygen atoms was reported by Pflau et al.84 for Rh on CeO2. The reaction of O atoms from CeO2 with background CO gas to make CO2 catalyzed by supported metal particles, can also reduce CeO2, which has been found for Pt on CeO2.28,78

In the present case of Ag/CeO2(111), our analysis shows that the final state effects are primarily responsible for the Ag 3d core level binding energy shifts. The small contribution from the initial state effects at very low Ag coverages is attributed to the low average coordination number for Ag atoms of small Ag particles. This means that there is no strong interfacial interaction between Ag and CeO2(111). When the Ag coverage increases to be higher than 0.5 ML, the contribution from initial state effects to the Ag 3d core level shifts becomes negligible, while the reduction of CeO2(111) due to Ag adsorption can still be observed when the Ag coverage is above 0.5 ML. This clearly indicates that the reduction of CeO2(111) does not result from electron transfer from Ag to CeO2(111). Moreover, Ag has a lower heat of oxide formation and therefore a lower oxygen affinity than cerium; therefore, it is unreasonable that Ag clusters are oxidized at the Ag/CeO2 interface. However, it is worth pointing out that our observations do not agree with recent DFT calculations,18 which indicated that the adsorbed Ag metal atoms became oxidized (AgO) by one of the Ce4+ cations in the underlying oxide.

At room temperature CO does not adsorb on the stoichiometric CeO2(111) surface, as previously demonstrated by both experimental and theoretical results.96,99 We also found that the CeO2(111) surface is not reduced after leaving the fresh-prepared CeO2(111) surface in the vacuum chamber for several hours. In other words, the background CO can not be adsorbed on the CeO2(111) surface at 300 K. A similar experiment was also performed for 0.1 ML Ag covered CeO2(111). The electronic structure of the surface, e.g., the unoccupied Ce 4f band, shows little change after the sample stays in the vacuum chamber for a long time. This suggests that Ag-catalyzed reaction of O atoms from CeO2 with background CO to form CO2, which induces the reduction of CeO2, does not occur at 300 K.

Having ruled out the possibilities of charge transfer from the supported Ag particles to CeO2 and Ag-catalyzed reaction of O atoms from CeO2 with background CO to form CO2 at 300 K, we attribute the reduction of CeO2(111) to spillover of lattice oxygen in CeO2 at the Ag–CeO2 boundary to the surface of the supported Ag nanoparticles. This process can happen as long as the CeO2(111) surface is not completely covered by Ag. This explains well why at higher Ag coverages, before a continuous Ag film is formed, the reduction of CeO2 can still be observed. One might argue that since the heat of oxide formation of Ag is significantly lower than that of CeO2, this reverse spillover of oxygen atoms from CeO2 to Ag nanoparticles less likely occurs. However, Pt has heat of oxide formation similar to Ag,2 but as mentioned above, in the nanometer scale the energy of oxygen atoms on Pt nanoparticles does not differ too much from that in the surface layer of the CeO2; the actual barrier for spillover of O from CeO2 to Pt is very low.88 Therefore, it is reasonable that a similar process might also occur for Ag on CeO2. These spillovered oxygen atoms on metal nanoparticles might be very reactive, making the Ag/CeO2 catalysts highly active in the redox reactions. However, we can not rule out the possibility of Ag-catalyzed desorption of surface oxygen atoms from CeO2 since in our SRPES experiments the intensity of O 1s indeed decreases upon Ag adsorption, although this decrease may be induced only by the damping effects of the Ag adsorbate. It should be mentioned that theoretical calculations might clarify the exact reduction mechanism of CeO2 upon Ag deposition.

A similar process is also observed when CeO2 particles are deposited on late transition metals (i.e., so-called inverted model catalysts). There, the CeO2 is always observed to be slightly reduced by XPS.90,91 This effect has also a strong promoting function in CO oxidation on different metal surfaces.

It is interesting that when Ag is deposited onto TiO2(110) in a way similar to that done here on CeO2(111) no evidence was seen in the valence band photoemission spectrum for the reduction of the TiO2 or for Ti3+ states (even though Ti3+ has a feature similar to the 1.5 eV BE peak seen here for Ce3+, when present).57 Similarly, no evidence is seen for the reduction of a SnO2 surface upon Ag deposition.92 Thus, the present results show that the CeO2(111) surface is more easily reduced than both TiO2 and SnO2. That it can even be reduced slightly by deposited Ag is somewhat surprising.

5. CONCLUSION

The growth, nucleation, and electronic properties of Ag on the ordered CeO2(111) thin films grown on Ru(0001) have been systematically studied by SRPES, XPS, and LEED. The results clearly indicate that Ag grows three-dimensionally on the well-ordered CeO2(111) surface at 300 K. When the CeO2(111) surface has a high density of defects, Ag populates these defect sites initially, leading to a 2D island growth at low coverages. The binding energy of Ag 3d increases when the Ag particle size decreases, which is mainly due to the final state screening. The interfacial interaction between Ag and CeO2(111) is weak. Ag particles most likely maintain the metallic state on the CeO2(111) surface. The slight reduction of the CeO2(111) surface upon Ag deposition can be ascribed to the reverse spillover of oxygen atoms on CeO2 at the Ag–CeO2 boundary to the Ag nanoparticles. Heating the Ag/CeO2(111) surface leads to the significant sintering of Ag nanoparticles before they completely desorb from the CeO2(111) surface.

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