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ABSTRACT

A novel method of measuring the core level binding energies of multiple sized nanoparticles on the same substrate is demonstrated using the early stage of Au nanoparticle growth on reduced r-TiO$_2$(110). This method employed in situ scanning tunneling microscopy (STM) and microfocused X-ray photoemission spectroscopy. An STM tip-shadowing method was used to synthesize patterned areas of Au nanoparticles on the substrate with different coverages and sizes. Patterns were identified and imaged using a UV photoelectron emission microscope. The Au 4f core level binding energies of the nanoparticles were investigated as a function of Au nanoparticle coverage and size. A combination of initial and final state effects modifies the binding energies of the Au 4f core levels as the nanoparticle size changes. When single Au atoms and Au$_3$ clusters are present, the Au 4f$_{7/2}$ binding energy, 84.42 eV, is similar to that observed at a high coverage (1.8 monolayer equivalent), resulting from a cancellation of initial and final state effects. As the coverage is increased, there is a decrease in binding energy, which then increases at a higher coverage to 84.39 eV. These results are consistent with a Volmer-Weber nucleation-growth model of Au nanoparticles at oxygen vacancies, resulting in electron transfer to the nanoparticles.

INTRODUCTION

Photoelectron spectroscopy (PES) has been widely employed to explore the electronic characteristics of nanoparticles and their interactions with single crystalline substrates. Here, we refer to earlier studies of nanoparticles on TiO$_2$(110) since this is the model system studied in the present work. Au nanoparticles are chosen because of their significance in several industrial processes, including oxidation and water gas shift catalysis. The choice of support material has been shown to have a critical effect on the catalytic activity of nanoparticles, with TiO$_2$ being particularly effective. Moreover, the question of charge transfer between the nanoparticle and the substrate is a much-debated aspect of Au based catalysis. The surface chemistry of Au supported on TiO$_2$ has been recently reviewed. PES core level electron binding energy shifts of nanoparticles can be explained by changes in electron distributions and, in some cases, the charge transfer between a nanoparticle and the substrate. In the case of Au nanoparticles, the binding energy of core level electrons has been shown to vary with the nanoparticle size on polycrystalline TiO$_2$. In addition to the Au charge state and particle size, the extent to which the substrate has been reduced prior...
to Au deposition has been shown to influence the catalytic activity of the supported nanoparticles and the degree of charge transfer.

Therefore, describing the interactions between the centers of reduction (e.g., oxygen vacancies) and the Au nanoparticles is key to understanding the relationship between the catalytic activity and the reduction state of the substrate. There are many independent studies that have examined the nature of electron transfer occurring between Au nanoparticles and reduced TiO$_2$. The consensus has been that anionic gold is formed at a low coverage in UHV, although recent calculations suggest that neutral or cationic gold is obtained.

Reduction of TiO$_2$(110) gives rise to an electronic state within the bandgap of the material. The presence of this band-gap-state (BGS) is associated with the generation of defects such as interstitial Ti atoms and bridging oxygen vacancies (Ov) in the bulk and at the surface, respectively. Oxygen vacancies on rutile TiO$_2$(110) act as electron trap sites and have been shown to play a role in surface-mediated molecular reactions. They are associated with an area of excess electron density, and low temperature scanning tunneling microscopy (STM) has been used to show that this has polaronic character. Previous experimental and theoretical work has pointed to the initial adsorption of Au atoms, Au$_1$, at Ov sites on reduced r-TiO$_2$(110). This has recently been directly visualized using STM. Au$_1$ and Au$_3$ are bound at Ov, but larger nanoparticles are mobile at room temperature and agglomerate to form larger particles with an increased coverage.

Interpretation of core level electron binding energy shifts of nanoparticles as a function of size is made difficult by the presence of both initial and final state effects. Initial state effects arise from intrinsic properties of the nanoparticles, i.e., the initial electronic state of the system. For metal nanoparticles on a metal oxide, this would be evident if there was charge transfer. For example, if the nanoparticle becomes positively charged, then the binding energy will increase. Final state effects arise from the photoemission process, i.e., its electronic state immediately after photoemission. Final state effects diminish as the particle size increases due to the screening of the newly generated core hole by its local electronic environment. Hence, the final state binding energy shift becomes less positive as the particle size increases. For Au on stoichiometric TiO$_2$(110), earlier work observed a decrease in Au 4f binding energies with an increase in Au coverage. In contrast, for reduced TiO$_2$(110), the binding energy initially decreased but then increased at a higher coverage. The latter behavior is ascribed to a reduction in the averaged charge transfer to the nanoparticles at higher coverages.

A potential drawback in studies of reducible oxides such as TiO$_2$ is that different levels of reduction can be inadvertently introduced during a series of measurements, which can modify the degree of charge transfer to/from supported nanoparticles. In this work, we demonstrate a novel experimental approach employing micro-focused X-ray photoemission spectroscopy ($\mu$-XPS) and STM that can be employed to study multiple sized nanoparticles on the same substrate. Au nanoparticles were deposited onto TiO$_2$(110) in discrete regions using a scanning tunneling microscope tip shadowing effect, with each region containing Au nanoparticles of a different size distribution. This minimizes any discrepancies arising from the variable reduction state of the substrate, such as defect density. Moreover, imaging the nanoparticles in situ removes any uncertainty regarding the size of the nanoparticles and their binding sites.

**EXPERIMENTAL AND INSTRUMENTAL DETAILS**

The experiment was carried out on the XPEEM end-station of the I06 Nanoscience beamline at Diamond Light Source. This end station has a preparation chamber equipped with STM (Omicron VT STM), low energy electron diffraction (LEED)/retarding field Auger electron spectroscopy (AES), and an analysis chamber equipped with an X-ray photoelectron microscopy (XPEEM) instrument (Elmitec SPELEEM-III), which can also be used to image with low energy electron microscopy (LEEM) and UV-PEEM with a Hg lamp (Quantum Design Europe). Both chambers operate at a base pressure in the $10^{-10}$ mbar region. $\mu$-XPS spectra were acquired in dispersive mode at a photon energy of 200 eV with a field of view of 8 $\mu$m. The photoelectron energy was calibrated using the Fermi level and the bandgap state. This was systematically cross-checked using the Ti 3p binding energy.

Two TiO$_2$(110) (MaTecK GmbH) samples were introduced into the UHV system. Both were prepared by five cycles of 1 keV Ar+ sputtering and annealing at 1100 K for 10 min. After preparation, a LEED pattern was observed and the absence of surface contamination was confirmed using AES. The clean samples were then introduced into the STM, where images of the clean surface were recorded to check the presence of large terraces. Au deposition was carried out at room temperature using an e-beam evaporator (Omicron EFM3) oriented toward the STM tip in its scanning configuration. The evaporator built-in flux monitor was stabilized to a given flux, which was constant throughout the experiment. The height histogram of a large assembly of nanoparticles, typically over 100, was obtained using STM. For Au$_1$ and Au$_3$, the nanoparticles were simply counted. The monolayer equivalent (MLE) coverage was then determined by summing all the heights with the assumption that the nanoparticles for Au$_x$ and larger are hemispherical and that the lattice parameter of Au in nanoparticles is that of the bulk.

To produce a stepped gradient of the Au nanoparticle size over the surface, a STM tip shadowing effect was employed. In this methodology, the STM tip acts as a shutter, blocking the pathway between an area of the sample and the incoming metal atoms emitted by a line of sight evaporator. By moving the STM tip over the surface in the precise direction of the evaporator, the previously shadowed area of the surface is progressively exposed to the incoming flux of metal atoms. This gives rise to a continuously varying gradient of nanoparticle coverage on the surface of the substrate. By halting metal deposition during each incremental movement of the STM tip (i.e., closing the shutter on the evaporator; see Fig. 1), well-defined discrete areas of monotonic nanoparticle coverage could be deposited, rather than a continuously varying gradient.

The sample was then transferred to the XPEEM analysis chamber, where a few-micrometers wide patterns created using STM were identified using UV-PEEM. This imaging method offers the best contrast between areas of Au nanoparticles that have different but similar diameters. As the tip shadow area measures only 100 nm
FIG. 1. Schematic diagram of the STM-tip shadow method used to deposit discrete regions of different Au coverages onto a TiO$_2$ substrate. The motion of the tip between deposition stages is indicated in the diagram. Two tip-shadows are shown: the first (outer) exhibits a lower Au coverage than the nonshadowed area of the sample while the second (inner) shows the bare substrate.

at its apex (see below), it was not feasible to locate it by exploring the entire sample with UV-PEEM. To circumvent this problem, we employed an area retrieval method detailed in the next paragraph. This was also employed to optimize the tilt of the sample, which is an important parameter during the PEEM alignment. It effectively defines an area of the sample where lateral movement using the in-plane piezomotor translations does not compromise the alignment of the sample and the analyzer. This allows for lateral movement, within a range of approximately 500 μm, from one area of interest to another with no impact on the reliability of the μ-XPS signal.

The definition of a set of coordinates that can be located and recalled in both the STM and XPEEM represents a critical step in this experiment. To do so, a special sample (Siemens Star™, shown in Fig. 2) was used. This has a diameter of 500 μm with a 5 μm ring that marks its exact centre, which is visible both optically, via the STM CCD camera, and with LEEM. The coordinates of the tilt free zone were then indexed using LEEM, and the optimal area for depositing the nanoparticle shadow pattern was determined. The star sample was then transferred to the STM, which was equipped with a CCD camera capable of resolving movements of the STM tip as small as 30 μm over a sample’s surface. LEEM relative coordinates of the STM tip were determined and recorded with respect to the optically visible central ring. For future samples, the optimum position of the STM tip for nanoparticle shadowing (i.e., within the tilt free zone) could be recalled. This position was then located using the STM coordinate system before recording an image. After creating the tip-shadowed Au/TiO$_2$ sample and having defined the position of the shadow’s apex in both the STM and XPEEM apparatus, the sample’s surface was imaged using UV-PEEM. Figure 3 shows a series of stitched UV-PEEM images taken from the Au/TiO$_2$ sample. In the images, multiple well-defined impressions of the STM tip are seen on the surface, which become incrementally darker toward the sample’s edge. The change in image contrast can be explained by an increased local work function of the surface for areas with a higher Au coverage. The inset of Fig. 3 shows a magnified view of the apex of an STM tip impression (nanoparticle shadow). This is also evident in an STM image of an area of the shadowed surface (see Fig. 4).
RESULTS AND DISCUSSION

A feature of μ-XPS when used in the study of reducible metal oxides is that the high brightness of the incident X-rays can induce beam damage in the form of reduction. This has been observed for TiO$_2$ and CeO$_2$. This effect is likely caused by ionization, excitation, and de-excitation events that reduce the surface. For CeO$_2$(111), this effect was removed by measuring in a partial pressure of O$_2$. We adopt this methodology in the present work, its efficacy being demonstrated by the μ-XPS results shown in Fig. 5. In the presence of 10$^{-6}$ mbar O$_2$, there is no X-ray induced reduction of Au-nanoparticle decorated TiO$_2$(110), as evidenced by the absence of an increase in the bandgap state peak at 0.8 eV BE and Ti 3p features in the range 34–36 eV BE. Reducing the pressure of O$_2$ results in the appearance of these features within a few seconds, indicating reduction to Ti$^{2+}$ and Ti$^{3+}$. There is a corresponding increase in the intensity of bandgap states that arise from Ti 3d electrons. Subsequently, an attempt to reoxidize the sample was made by reintroducing 1 × 10$^{-6}$ mbar of O$_2$ into the chamber, although this did not result in a complete reoxidation.

To exclude the possibility that Au nanoparticles on TiO$_2$(110) are modified by the X-ray beam, the Au 4f signal from a 0.4 MLE covered surface was measured as a function of O$_2$ pressure. The evolution of the Au 4f$_{7/2,5/2}$ spectra shown in Fig. 6 was recorded immediately after the acquisition of the corresponding Ti 3p and bandgap spectra presented in Fig. 5. The spectra in Fig. 6 show that the peak intensities are not affected by the decrease in oxygen partial pressure. The slight shift in energy observed at 3 × 10$^{-6}$ mbar of O$_2$ can tentatively be attributed to the interaction between Au NPs and highly reduced Ti species. This last point means that the interpretation of spectrum acquired at low O$_2$ partial pressure has to consider beam-induced reduction of the substrate. Here, we focus on Au 4f data collected at 10$^{-6}$ mbar of O$_2$, where the chemical shift is independent of beam exposure.

We now turn to the variation of the electronic structure of Au nanoparticles as a function of size. This was monitored through the Au 4f binding energy shift for six different coverages obtained on two TiO$_2$(110) single crystal samples, one of them reprepared once. The first sample supported a uniform coverage of 1.8 MLE of Au. The second and third samples were prepared using STM tip-shadowing to support coverages of 0.4, 0.1, and 0.05 MLE, and 0.2 and 0.3 MLE, respectively. As noted above, this method of nanoparticle growth is employed to minimize effects arising from a variation in the level of substrate reduction.

The coverages at which Au 4f spectra were obtained are calculated from the analysis of the STM images shown in Fig. 7, which cover the Au coverage range 0.007–1.8 MLE. For 1.8 MLE coverage, the Au 4f spectrum (see Fig. 8) corresponds to the surface imaged by STM [Fig. 7(f)]. To obtain the remaining Au 4f coverages, we employed the STM images to calibrate the Au doser. At the lowest coverage, the image contains only Au; and Au nanoparticles, identified by their height of 1.5 Å and 3 Å, respectively. At higher coverages, these species are still present, but larger nanoparticles increasingly dominate. As an example of the height distribution observed, the histograms measured from the images in Figs. 7(c) and 7(f) are shown in Fig. 8.

The Au 4f spectra were recorded for six coverages (0.05, 0.1, 0.2, 0.3, 0.4, 0.5, and 1.8 MLE) at four different partial pressures of oxygen (1 × 10$^{-6}$, 3 × 10$^{-7}$, 1 × 10$^{-7}$, and 3 × 10$^{-8}$ mbar). For a given coverage, the binding energy and intensity of the Au 4f$_{7/2,5/2}$ components are independent of the oxygen partial pressure. This indicates that any beam damage occurring on the substrate in the vicinity of the nanoparticles does not affect the Au 4f binding energies. The data recorded at an oxygen partial pressure of 10$^{-8}$ mbar are shown in Fig. 9 with the Au 4f$_{7/2}$ binding energy variation with the coverage.
plotted in Fig. 10. They display a clear binding energy shift that can be separated into two regions. In the first region, corresponding to 0.05 to 0.3 MLE Au, the Au 4f_{7/2} peak shifts by $-109$ meV. In the second region, corresponding to 0.3 to 1.8 MLE Au, the Au 4f_{7/2} peak shifts by $+80$ meV (from 84.31 to 84.39 eV).

As noted above, studies of stoichiometric surfaces report only a negative binding energy shift of the Au 4f signal as Au coverage increases. This was ascribed to a final state effect. In contrast, for a reduced surface, an initial decrease in binding energy is followed by an increase in binding energy at a higher coverage, arising from a reduction in the charge transfer per Au atom. The same trend is seen in the present work, although details of the binding energies differ, presumably because of differences in the deposition conditions that give rise to different particle sizes in the earlier work.

In addition to measuring the binding energy shifts on substrates with the same level of surface reduction, we have access to the nanoparticle size distribution through our in situ STM measurements. This allows additional qualitative comments on the
binding energy shifts to be made. The predominant species observed below 0.05 MLE are $\text{Au}_4$ and $\text{Au}_5$. The reason is that these species grow rather than larger particles pointing to the reduced nature of the substrate, which contains oxygen vacancies. Hydroxylated surfaces are known to form larger nanoparticles.11 Both $\text{Au}_4$ and $\text{Au}_5$ are bound to oxygen vacancies on the basis of earlier work,12 which is expected to give rise to charge transfer to the nanoparticles to form negatively charged Au.13 This should shift the Au 4f/2 peak to lower binding energy, although the lack of screening of the core hole in such small, well-dispersed nanoparticles will provide a shift in the opposite direction. These effects essentially balance each other, resulting in an Au 4f/2 binding energy of 84.42 eV similar to that at 1.8 MLE of 84.39 eV. At higher coverages, the size distribution increases substantially (see Fig. 8) with $\text{Au}_4$ and $\text{Au}_5$ still visible in the 1.8 MLE-coverage STM image [Fig. 7(f)]. This mid-nucleation phase arises when a significant proportion of the surface $\text{O}^-$ has been filled, and a subsequent addition of Au results in diffusion and attachment of atoms to an existing nanoparticle.

An earlier room temperature STM study found the transition between nucleation and growth of Au on $\text{TiO}_2$(110) to occur around 0.15–0.2 MLE,10 consistent with the results presented here. The corresponding shift of Au 4f/2 to lower binding energy with an increase in the mean nanoparticle diameter (Figs. 9 and 10) indicates that a reduction in the final state effect dominates in this regime. In other words, the increase in core hole screening produces a larger shift than that arising from a decrease in the charge transfer per Au atom. During the late growth phase, as the NPs grow bigger and begin to merge together due to coalescence and/or Ostwald ripening,12 the charge transfer per gold atom approaches zero and, hence, the Au 4f binding energy increases. Although this interpretation is in line with previous calculations of the Au-$\text{TiO}_2$(110), at first sight, it is not consistent with recent calculations that predict the formation of neutral or cationic nanoparticles.13

**SUMMARY**

A novel combination of STM, UV-PEEM, and μ-XPS has been employed to study the interaction between the surface of $\text{TiO}_2$(110) and Au nanoparticles in the coverage range 0.05–1.8 MLE. The Au 4f photoemission binding energies of size-selected nanoparticles were measured as a monitor of the charge transfer to the nanoparticles and the screening of the core hole. Nanoparticles were fabricated using an in situ STM, using a tip shadowing method to generate a surface pattern of different coverages of Au nanoparticles. The pattern was aligned using UV-PEEM to realize μ-XPS measurements of discrete region successively. This has the advantage of allowing the measurement of different sized nanoparticles on the same substrate. μ-XPS spectra were recorded in a background pressure of 10⁻⁶ mbar of $\text{O}_2$ to minimize the effect of beam reduction of the substrate. The μ-XPS data show Au 4f binding energies at the ultradilute limit (0.05 MLE) are similar to those at a higher coverage (1.8 MLE), arising from a cancellation of initial and final state effects for $\text{Au}_4$ and $\text{Au}_5$. As the coverage increases, the binding energy initially decreases and subsequently increases. This is consistent with charge transfer to $\text{Au}_4$ and $\text{Au}_5$ bound to oxygen vacancies in the ultradilute limit.

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