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J. Phys. Chem. C, Just Accepted Manuscript • DOI: 10.1021/acscpcc.5b02667 • Publication Date (Web): 27 May 2015

Downloaded from http://pubs.acs.org on May 31, 2015

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Atomic Structure Control of Silica Thin Films
on Pt(111)

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Abstract

Metal oxide thin films grown on metal single crystals are commonly used to model heterogeneous catalyst supports. The structure and properties of thin silicon dioxide films grown on metal single crystals have only recently been thoroughly characterized and their spectral properties well established. We report the successful growth of a three-dimensional, vitreous silicon dioxide thin film on the Pt(111) surface and reproduce the closed bilayer structure previously reported. The confirmation of the three-dimensional nature of the film is unequivocally shown by the infrared absorption band at 1252 cm\(^{-1}\). Temperature programmed desorption was used to show that this three-dimensional thin film covers the Pt(111) surface to such an extent that its application as a catalyst support for clusters/nanoparticles is possible. The growth of a three-dimensional film was seen to be directly correlated with the amount of oxygen present on the surface after the silicon evaporation process. This excess of oxygen is tentatively attributed to atomic oxygen being generated in the evaporator. The identification of atomic oxygen as a necessary building block for the formation of a three-dimensional thin film opens up new possibilities for thin film growth on metal supports, whereby simply changing the type of oxygen enables thin films with different atomic structures to be synthesized. This is a novel approach to tune the synthesis parameters of thin films to grow a specific structure and expands the options for modeling common amorphous silica supports under ultra high vacuum conditions.

**Keywords:** UPS (He I\(_\alpha\)), MIES, TPD, IRRAS, SiO\(_2\), CVD.
Introduction

Thin metal oxide films have become a standard tool to support model heterogeneous catalysts under ultra high vacuum (UHV) conditions, as well as being studied for an assortment of applications in materials science. Silicon dioxide (SiO$_2$) is one of the most commonly used supports in heterogeneous catalysis, and much progress has been made in growing thin SiO$_2$ films on metal single crystals, as well as revealing the underlying growth mechanisms and structures formed. Mo(110), Mo(112) and Ru(0001) have all been shown to be ideal candidates for the growth of these films and their structural properties have been well documented. From these studies it has been shown that the type of silica film can be solely determined by its infrared vibration band: $\sim$1300 cm$^{-1}$ for a bilayer, $\sim$1250 cm$^{-1}$ for a thick, vitreous film, and $\sim$1065 or $\sim$1135 cm$^{-1}$ for a monolayer film on Mo(112) or Ru(0001), respectively. SiO$_2$ films have also been successfully grown on Ni(111), Pd(100), and Pd(111), but have not been characterized to the extent of the aforementioned surfaces. Pt(111) has only recently emerged as another candidate for growing thin SiO$_2$ films and has been shown to form a closed bilayer structure, similar to the one observed on Ru(0001), and scanning tunneling microscopy (STM) showed that it is vitreous and does not completely cover the surface.

The trend observed for the structure of these thin films was shown to follow two parameters: lattice mismatch and oxygen affinity of the metal support. A greater lattice mismatch results in a non-crystalline thin film, as was shown for the bilayer on Pt(111) compared to Ru(0001). Oxygen affinity influences the ability to form Si-O-metal bonds and this is believed to allow for the formation of a monolayer film seen on Mo(112) and Ru(0001). Both the large lattice mismatch and the low oxygen affinity of Pt(111) mean that silica has only been grown as a closed bilayer with no Si-O-Pt bond (low oxygen affinity) and in a vitreous state (large lattice mismatch). In this paper we reproduce on Pt(111) the bilayer previously reported and successfully synthesize a three-dimensional vitreous film. Using Auger electron spectroscopy (AES), metastable impact electron spectroscopy (MIES), ultraviolet
photoelectron spectroscopy (UPS), and infrared reflection absorption spectroscopy (IRRAS), we show that the atomic structure of the film is determined by the amount of oxygen on the surface after silicon deposition and that the thicker films exhibit similar spectroscopic behavior observed for other thick SiO$_2$ films grown on Mo(112) and Ru(0001) while retaining the vitreous nature observed for the bilayer. Lastly, the applicability of a bilayer or three-dimensional film as a support for model catalyst systems is also investigated using temperature programmed desorption (TPD). The ability to grow a thin silica film on a Pt(111) surface that completely covers the metal enables the direct comparison between a catalytically active single crystal and supported metal particles using the same experimental instrumentation, i.e. random error arising from the use of two single crystals (e.g. Mo(112) for supported clusters and Pt(111) as a model single crystal catalyst) can be ruled out. Furthermore, it has been shown that the metal upon which thin films are grown can influence nanoparticle morphology by a charge transfer mechanism from the thin film/metal interface.$^{21-23}$ The ability to grow thin, closed silica films on Pt(111), allows for the investigation of these structure-property relationships with regards to the underlying metal support. The change in work function between Mo, Ru and Pt, represents a decisive parameter governing surface charging effects, whose influence on nanoparticle morphology, and hence catalytic activity, can now be investigated. This also represents a corollary to dopants in bulk metal oxides, and these effects can now be modeled by simply changing the metal support. This is an important step to expand the variety of parameters of model heterogeneous catalyst systems under UHV and their comparability to more applied systems.

**Experimental**

All experiments were performed in a UHV chamber at a base pressure of $2 \times 10^{-10}$ mbar; details about the setup can be found elsewhere.$^{24}$

Prior to the measurements the Pt(111) crystal (MaTeck, Germany) was cleaned by
repeated cycles of Ar$^+$ sputtering ($T = 900$ K, $p(Ar^+) = 5 \times 10^{-6}$ mbar, $\Delta t = 60$ min, $U = 1$ keV, $I \sim 10 \mu$A) followed by oxidation ($T = 650$ K, $p(O_2) = 5 \times 10^{-7}$ mbar, $\Delta t = 5$ min) and annealing ($T = 1300$ K, $\Delta t = 1$ min); subsequently the single crystal purity was checked by means of AES and MIES/UPS and is in good agreement with the literature.

The SiO$_2$ thin film was prepared by e-beam evaporation (Tectra, Germany) of a silicon rod (99.999% purity, Alfa-Aesar, Germany) in an oxygen (5.5 purity, AirLiquide, Germany) background pressure ($T = 400$ K, $p(O_2) = 2 \times 10^{-6}$ mbar) onto the Pt(111) single crystal surface. The flux of the silicon beam was measured using an electrode on the end of the evaporator, with the normal flux being $\sim 10$ nA and, where specified, half-flux being $\sim 5$ nA. The film was then annealed at 1200 K in a $2 \times 10^{-6}$ mbar oxygen atmosphere before experimental data was acquired, unless otherwise mentioned. A single-shot four minute and ten minute film growth step was performed to synthesize the bilayer and three-dimensional thin film, respectively. The electron emission spectra (EES) were all recorded at 100 K and further details regarding the spectroscopic setup can be found in our previous publications.$^{25,26}$ The IRRAS experiments were performed at 100 K (Thermo-Nicolet FT-6700, Thermo-Nicolet MCT detector) and averaged 256 scans at 4 cm$^{-1}$ resolution. The TPDs (Balzers QMG 430, Lichtenstein) were run with a temperature ramp of 2 K/s and the gas dosage was performed with a calibrated molecular beam doser.$^{27}$

**Results**

IRRAS spectra taken after four and ten minutes of film growth on Pt(111) followed by annealing at 1200 K are shown in figure 1. The peak at 1288 cm$^{-1}$ corresponds to Si-O-Si vibrations within a closed bilayer thin film after four minutes of film growth.$^{16}$ After ten minutes of silicon evaporation a peak at 1252 cm$^{-1}$ with a broad shoulder down to 1100 cm$^{-1}$ appeared and these are the asymmetric longitudinal phonon vibrations, characteristic for a thick vitreous film. Ru(0001) also exhibited this exact trend where a bilayer was first observed.
with a peak at 1300 cm\(^{-1}\) and upon further film growth the bilayer peak disappeared and a peak centered at 1257 cm\(^{-1}\) with a broad shoulder down to 1100 cm\(^{-1}\) appeared.\(^{15}\) The clear formation of the bilayer after 4 minutes indicates that the film deposition rate is on the order of 0.5 monolayer/min. The presence of the thick film after 10 minutes would mean a film thickness of 5 atomic layers, which is similar to the thickness of 3-dimensional films grown on Ru(0001).\(^{15}\) Assuming similar values for the thin films on Ru, the approximate thickness of the 3-dimensional film grown on Pt(111) is 10 Å.

Figure 1: IRRAS spectra for the three-dimensional and bilayer silica film. Each film shows a characteristic peak: bilayer at 1288 cm\(^{-1}\) and the three-dimensional film at 1252 cm\(^{-1}\).

The EES spectra in figure 2 show the SiO\(_2\) valence bands after four and ten minutes of film growth each followed by 1200 K annealing before spectral characterization. The UP spectra have been background corrected by means of a previously introduced method in order to remove secondary electron contributions.\(^{25}\) This procedure was not applied to the MIES spectra as the secondary electron contribution was not as significant. After four minutes of film growth a broad peak appeared, centered at 6.4 and 6.7 eV for UPS and MIES, respectively, belonging to the O(2p) nonbonding state. Additionally, a MIES peak emerged at 10.7 eV belonging to the Si-O bonding state. After ten minutes of film growth the O(2p) peak shifted
to 6.8 and 7.4 eV for UPS and MIES, respectively, while the Si-O peak appeared at 11.0 eV for UPS and MIES.\textsuperscript{12}

Figure 2: MIE (a) and background corrected UP (b) spectra of the silica thin film at different preparation steps. The bottom spectrum shows the clean Pt(111) single crystal surface, then after four minutes of film growth with subsequent annealing at 1200 K and lastly after ten minutes of growth and annealing at 1200 K. The uppermost spectrum can be assigned to the vitreous, three-dimensional SiO\textsubscript{2} and the middle spectrum to the bilayer thin film.
Deuterium TPD spectra were collected from both the three-dimensional and bilayer film can be seen in figure 3. The spectra show that the thick film has no D$_2$ desorption peak, whereas the bilayer shows a small peak centered at 300 K corresponding to desorption from uncovered Pt(111).

Figure 3: Deuterium TPD spectrum of 0.4 D$_2$/surface atom dosed at 100 K on the three-dimensional and bilayer silica film.
Discussion

Figure 4: IRRAS spectra (a) after 20 min film growth using half the silicon flux as previously reported, followed by five minutes of normal flux film growth, both after annealing at 1200 K. AES spectra (b) for a clean Pt(111) surface, the bilayer SiO$_2$ film after 20 minutes of low flux silicon deposition, and the multilayer film after 10 minutes of film growth at normal flux (both without annealing at 1200 K), keeping the total amount of silicon deposited constant. The region above 150 eV has been multiplied by a factor of two.
The results of figure 1 demonstrate that a three-dimensional, vitreous silica film can be grown on Pt(111). This overcomes the limitations which restricted silica films to exclusively a bilayer structure. The reason for the formation of the three-dimensional film in contrast to previous reports, was identified by varying the film growth parameters. A SiO$_2$ film was grown for 20 min using half the silicon flux previously used, followed by acquisition of an IR spectrum seen in figure 4a. Even after 20 min of film growth, only the bilayer formed. On top of this film, five minutes of the normal flux film growth protocol was performed and the three-dimensional film was observed. Auger spectra taken after an identical low flux experiment were compared with those taken after the normal flux experiment (the ten minute standard growth protocol), both seen in figure 4b. Both AES experiments were performed in a way that an equal amount of silicon was evaporated on the Pt(111) surface. As both spectra were taken before high temperature annealing, both are showing the same surface concentration of silicon. No large elemental silicon peak at 92 eV is visible, in contrast to silicon deposited on and segregated from the Pt(111) surface. This is, however, to be expected in the presence of atomic oxygen, where no elemental silicon peak is observed after oxidizing at $\sim$400 K. Instead, when using molecular oxygen as the oxidizing agent, a very large peak at 92 eV is visible. The absence of a well defined peak at 92 eV implies that atomic oxygen, likely being generated in the evaporator, is oxidizing the deposited silicon. We identify the oxygen species as atomic oxygen based on the similarity of our Auger spectra to those in Ref. [29]. A direct identification of atomic oxygen is not possible due to the geometry of the evaporator with respect to the mass spectrometer and therefore a definitive assignment is not possible. It is, however, unlikely that ionized oxygen species are responsible due to the potential of -18 V applied to the electrode at the end of the evaporator, but excited states of molecular and/or atomic oxygen cannot be ruled out.

The normal flux spectrum shows two peaks at 78 and 89 eV, which are known SiO$_x$(LVV) transitions from SiO$_2$ and SiO$_{x<2}$, respectively. The low flux spectrum shows only a very small feature at $\sim$93 eV indicating that the nature of silicon deposited under these conditions
is more varied than with the higher flux, i.e. the degree of oxidation varies from completely unoxidized to fully oxidized which causes the Auger peak to smear out over a wide range of energies, rendering it unresolvable in AES.

The amount of oxygen can be readily quantified from the O(KLL) Auger peak at 508 eV. Comparing the size of the oxygen peak to the Pt transition at 237 eV, the amount of oxygen almost doubles when doubling the flux. As both low and normal flux experiments have deposited the same amount of silicon, the only difference between the two surfaces is the amount of oxygen present. As the same background oxygen pressure was used for both experiments, the increase in oxygen must be originating from the increased evaporation rate (more specifically the increased filament current in the evaporator) and the most probable oxygen source which could be affected is atomic oxygen generated in the evaporator.

As SiO\textsubscript{4} tetrahedra have been proposed as the structural units for three-dimensional films on Ru(0001)\textsuperscript{9}, the concentration of these types of units during deposition and annealing must be a decisive factor discriminating between formation of purely the bilayer, and formation of three-dimensional SiO\textsubscript{2}. The SiO\textsubscript{2}(LVV) Auger peak at 78 eV clearly shows that the three-dimensional growth protocol produces more oxidized silicon which indicates that increasing atomic oxygen has the effect of increasing the surface concentration of fully oxidized silicon species. These species ultimately lead to the formation of a three-dimensional structure that completely covers the metal surface. It is also known that the saturation coverage of atomic oxygen on Pt(111) is approximately double that from molecular oxygen adsorption and dissociation.\textsuperscript{30,31} This evidences a compound effect, whereby increased atomic oxygen leads to a higher oxygen saturation coverage of open platinum surface domains, which facilitates the formation of fully oxidized silicon species and a complete covering of the metal. This shows that limitations on film growth imposed by a surface’s oxygen affinity and adsorption properties can be circumvented by using atomic oxygen while concomitantly changing the atomic structure of the thin film grown.

A second factor governed by the degree of silicon oxidation during deposition is etching
of the film during annealing at 1200 K effectively 'dissolving' the metal oxide. This could also play a factor in the unsuccessful attempts to grow three-dimensional SiO$_2$ on Pt(111) where a combination of a lower concentration of oxidized silicon and a higher concentration of elemental silicon both act to suppress the formation of three-dimensional SiO$_2$. After annealing our film to 1200 K all SiO$_x<2$ features disappeared, which we attribute to the desorption of SiO, in agreement with the known desorption above 1200 K from silicon surfaces.$^{33-35}$

The peak positions from the EES experiments compare well to experiments performed on Mo(112) but do show slight deviations, the most obvious being the O(2p) state, which does not split into two well resolved peaks after annealing at high temperatures ($\sim$1100-1200 K). Instead, this peak remains broad and shifts to higher binding energy for the thicker film. The broad nature of the valence band peaks has been attributed to a higher concentration of defect sites$^{36}$ and a lack of crystallinity within the thin film.$^8$ The deviation of 0.6 eV between the two spectroscopic methods can be explained by the surface sensitivity of MIES compared to UPS. MIES probes only the uppermost surface layer due to impenetrability of helium atoms into the surface. Thus, the UPS spectra originate from a convolution of the pure surface states seen in MIES and the underlying emission from the bulk, leading to the broader UPS peak and its shoulders on the high binding energy side.

The presence of a single peak centered at 11.0 eV, in both the UP spectrum and the MIE spectrum, also corroborates the lack of Si-O-Pt linkages. This is in contrast to EES spectra from SiO$_2$ on Mo(112) where a shoulder at 10.5 eV on the larger Si-O peak at 11.5 eV was observed and attributed to a Si-O-Mo bond.$^{12}$ The presence of only one peak in this binding energy range is to be expected, when Si-O-Pt bonds are absent. As previously mentioned, results on Pt(111) have shown a distinct lack of crystallinity when compared to Mo(112) and Ru(0001)$^9$ and our MIES/UPS results show that silica grown on Pt(111) produces no twin peak O(2p) feature corresponding to a crystalline film and no feature belonging to a Si-O-Pt state is observed.
The D$_2$ TPD experiments showed a desorption peak at 300 K from the bilayer film. This peak originates from an incomplete covering of the Pt(111) single crystal which has been observed using STM.$^{9,16}$ Covering the Pt(111) support entirely by the SiO$_2$ film is essential in order to be able to differentiate between signals originating from the underlying metal and e.g. deposited, sub-nanometer catalyst particles. The three-dimensional SiO$_2$ film showed no such desorption peak, indicating a complete covering of the surface, which makes it an ideal candidate for the use as a catalyst support material which may model industrially relevant supports.

**Conclusion**

In summary, we have reproduced the growth of a bilayer SiO$_2$ thin film on Pt(111) and observed the expected spectral behavior based on previous reports, including an incomplete covering of the surface which supports STM results$^9$. In addition, the growth of a three-dimensional vitreous SiO$_2$ film was achieved on the Pt(111) surface and exhibits the same characteristics as thick SiO$_2$ films produced on Ru(0001), Mo(110) and Mo(112).$^{10,14,15}$ The confirmation that the film belongs to the three-dimensional category was confirmed with IRRAS where a peak at 1252 cm$^{-1}$ with a shoulder down to 1100 cm$^{-1}$ is indicative of a multilayer silica film, in contrast to the sharp bilayer absorption band observed at 1288 cm$^{-1}$.

The absence of O(2p) peak splitting and fine structure within the EES spectra further indicates the lack of crystallinity in both the bilayer and thick silica film. Additionally, no other emission peak was observed that could be attributed to a Si-O-Pt state.

TPD measurements show that the bilayer film is a poor choice as a nanoparticle support for catalytic measurements due to incomplete coverage of the active Pt(111) surface. The thick film, however, shows that it completely covers the Pt(111) surface and can therefore be applied as a catalyst support for UHV experiments which more closely models supports used for ambient pressure experiments. Catalytic measurements on an active metal single
crystal as well as from particles supported on a thin film grown on the same single crystal, eliminates random error and allows for the most accurate comparison of measurements from the pure single crystal and the supported metal particles. The ability to grow the same thin film on three different metals (Mo, Ru and Pt) also allows for a systematic study of their influence on clusters/nanoparticles. The difference in work function between the metals is one interesting parameter which could lead to different particle morphologies by attenuating or enhancing the degree of charge transfer from the thin film/metal interface, while also providing a technique to model dopant effects in bulk metal oxides.

The exclusive growth of a bilayer film was observed when half the silicon flux compared to the normal procedure was used. By increasing the flux it was observed that the amount of oxygen present after deposition doubles even though the deposited amount of silicon, and oxygen background pressure, remain the same. This excess oxygen is tentatively attributed to atomic oxygen being generated by the silicon evaporator. Atomic oxygen would overcome the limitation imposed by the low oxygen affinity of Pt(111) and allows for a higher oxygen coverage on the surface during silicon evaporation compared to molecular oxygen. This leads to the formation of more fully oxidized silicon species needed for three-dimensional film growth and a complete covering of the surface. Excess elemental silicon deposited on the bilayer, and multilayer film is oxidized to volatile SiO when annealing the sample to 1200 K, which etches the film. Since in this case the multilayer SiO$_2$ film is also etched, the ability to grow three-dimensional films is dependent on the more oxidized silicon suppressing the etching along with the concomitant increase in the concentration of the requisite building blocks for a three-dimensional film. The identification of atomic oxygen as the discriminating factor between two atomic structures of silica thin films grown on the same metal support has, to our knowledge, never been demonstrated before. This represents an important parameter that can in principle be investigated for the control of atomic structure of any metal oxide thin film on any metal support.
Acknowledgements

C.J.R. and F.F.S. thank Prof. Hans-Joachim Freund and Dr. Shamil Shaikhutdinov for their hospitality during our stay at the FHI and their insights into the preparation and characterization of silica thin films. We acknowledge financial support from the DFG through the Nanosystems Initiative Munich (NIM) (F.F.S., C.J.R., M.D.R., A.S.C. and U.H.) and the Project He 3454/9-2 (F.F.S. and U.H.).

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