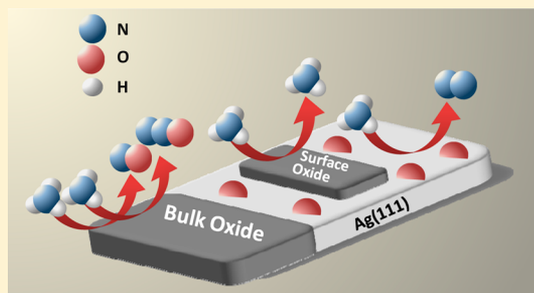


Selective Catalytic Ammonia Oxidation to Nitrogen by Atomic Oxygen Species on Ag(111)

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Supporting Information

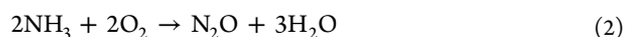
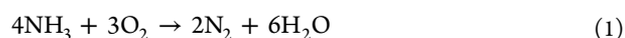
ABSTRACT: Ammonia-selective catalytic oxidation was studied on the planar Ag(111) single-crystal model catalyst surface under ultra-high-vacuum (UHV) conditions. A variety of oxygen species were prepared via ozone decomposition on pristine Ag(111). Surface coverages of oxygen species were quantified by temperature-programmed desorption (TPD) and X-ray photoemission spectroscopy techniques. Exposure of ozone on Ag(111) at 140 K led to a surface atomic oxygen (O_a) overlayer. Low-energy electron diffraction experiments revealed that annealing of this atomic oxygen-covered Ag(111) surface at 473 K in UHV resulted in the formation of ordered oxide surfaces (O_{ox}) with $p(5\times 1)$ or $c(4\times 8)$ surface structures. Ammonia interactions with O/Ag(111) surfaces monitored by temperature-programmed reaction spectroscopy indicated that disordered surface atomic oxygen selectively catalyzed N–H bond cleavage, yielding mostly N_2 along with minor amounts of NO and N_2O . Higher coverage O/Ag(111) surfaces, whose structure was tentatively assigned to a bulklike amorphous silver oxide (O_{bulk}), showed high selectivity toward N_2O formation (rather than N_2) due to its augmented oxygen density. In contrast, ordered surface oxide overlayers on Ag(111) (where the order was achieved by annealing the oxygen adlayer to 473 K) showed only very limited reactivity toward ammonia. The nature of the adsorbed NH_3 species on a clean Ag(111) surface and its desorption characteristics were also investigated via infrared reflection absorption spectroscopy and TPD techniques. Current findings demonstrate that the Ag(111) surface can selectively oxidize NH_3 to N_2 under well-defined experimental conditions without generating significant quantities of environmentally toxic species such as NO_2 , NO, or N_2O .



INTRODUCTION

Ammonia is an environmental pollutant whose exposure above 300 ppm is known to cause significant health risks such as lung diseases.^{1,2} European Environment Agency (EEA) reports indicate that the biggest contribution to ammonia emissions originates from agricultural activities³ while power plants and transportation are among other major sources of environmentally toxic ammonia in the atmosphere.⁴ The effects of ammonia on the ecosystem as an environmental pollutant were also reviewed comprehensively by Fangmeier et al.⁴ Various technologies have been developed for gas-phase ammonia abatement.⁵ Selective catalytic oxidation (SCO) of ammonia is a technology which can be used as an environmentally friendly method of ammonia abatement where N_2 and H_2O are produced as the major products. Both transition metals^{6–10} and noble metals^{11–16} were utilized as catalysts in the SCO of ammonia. Various catalytic systems in SCO of ammonia, including noble metals, transition metal oxides, and modified zeolites, were reviewed by Chmielarz et al.¹⁷ Among these metals, silver is considered as a promising low-temperature catalyst due to its high activity and high N_2 selectivity.^{14,18} The catalytic effect of silver nanoparticles on SCO reactions as well as their interactions with other metal systems were also investigated in the literature.^{6,12,19,20} However, a detailed

fundamental understanding of the ammonia SCO reaction mechanism on silver single crystals at the atomic/molecular level is still missing. As can be seen in the principal ammonia oxidation reactions given in eqs 1–3, the product distribution in ammonia SCO reaction depends on the active oxygen quantity, and thus, understanding the nature of oxygen is critical in terms of the selectivity of the catalyst. Some of the proposed mechanisms of the ammonia SCO process can be found in the literature.¹⁷ Technical ammonia SCO catalysts including silver nanoparticles as active sites reveal silver facets dominated by the (111) orientation.²¹ Thus, it is essential to elucidate the reactivity of the Ag(111) single crystal model catalyst surfaces in the ammonia SCO reaction.



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The O/Ag(111) system is an important catalytic system which has been thoroughly investigated within the context of different catalytic reactions in the past few decades both experimentally^{22–32} and theoretically.^{33–38} However, recent studies indicate that the O/Ag(111) structure proves to be an elusive system to understand in a comprehensive manner at the atomic scale.^{39–46} Oxygen-containing Ag(111) surfaces are known to undergo surface reconstruction readily. The most commonly observed reconstruction of this system is the p(4×4) structure, which was first observed by Rovida et al.²² The atomic structure of the p(4×4) reconstruction was revised in 2000,⁴⁷ and an even more recent model was proposed by Schnadt et al. in 2006.³⁷ In addition, other reconstructions of the O/Ag(111) system were also observed, such as ($\sqrt{3}\times\sqrt{3}$)R30°,⁴⁸ c(3×5 $\sqrt{3}$)rect,³⁷ (7×7),³² c(4×8),³⁸ and p(4×5 $\sqrt{3}$)rect structures.³⁸ The reconstruction process of the Ag(111) surface was reviewed by Michaelides et al.⁴⁹ As will be described below, none of these reconstructions besides c(4×8) were observed in the current study, presumably due to the use of a particularly different atomic oxygen dosing method comprised of ozone as the active atomic oxygen delivery agent. To the best of our knowledge, the ozone decomposition method was not used on Ag(111) single-crystal surfaces under UHV conditions in any of the former studies in the literature.

O₂ dissociation is not observed on the Ag(111) surface under UHV conditions due to the extremely low sticking coefficient of the oxygen molecule (ca. $\sim 10^{-6}$) on the Ag(111) surface at low pressures.²⁶ The minimum pressure for the observation of molecular oxygen dissociation on Ag(111) was reported to be 10^{-3} Torr in the literature.^{24,44} Thus, in most of the former studies, high-pressure O₂ exposures were used to investigate the nature of oxygen on a Ag(111) model catalyst surface.^{25–29,32} Other methods were also utilized for atomic oxygen delivery onto the Ag(111) surface under UHV conditions, such as NO₂ decomposition^{31,36,38,40,47,50} and thermal gas cracking.^{41,42,45} Ozone decomposition is an extremely efficient method for dosing high concentrations of oxygen atoms on metal surfaces under UHV conditions (i.e., in the absence of elevated-pressure exposures). Efficient atomic oxygen delivery onto single-crystal model catalyst surfaces with ozone was successfully carried out for Al(111),⁵¹ Pd(111),⁵² Pt(111),^{52,53} and Au(111).^{52,54} With this method, atomic oxygen can be accumulated on single-crystal surfaces even at pressures on the order of 10^{-10} Torr, while significantly minimizing the introduction of unwanted contaminations onto the catalyst surface. High-pressure oxygen accumulation leads to surface reconstruction and oxide formation rather than the generation of a mobile surface atomic oxygen overlayer on Ag(111). Heine et al. reported that oxide formation was observed to be more facile on the surface regions with high step density⁴⁴ whose generation can be triggered by high-pressure O₂(g). Thus, atomic oxygen accumulation under UHV conditions may limit oxide formation and allow exclusive investigation of high-coverage surface oxygen overlayers on Ag(111). Along these lines, an important advantage of ozone utilization over other atomic oxygen delivery vectors such as NO₂ decomposition is the fact that the ozone decomposition method enables the use of a greater variety of surface coverages of atomic oxygen species.

In the current work, we focus on the reactivity of oxygen atoms and oxide overlayers in ammonia SCO reactions on Ag(111) and report new catalytic structure–functionality relationships at the molecular level. We believe that the current findings may also provide valuable insights regarding the nature

of oxygen on Ag(111) which can also possibly be extrapolated to other crucial heterogeneous catalytic processes such as partial oxidation reactions of ethylene and methanol.^{27,57,71}

■ EXPERIMENTAL SECTION

Experiments were performed in a custom-made UHV chamber with a base pressure of 2×10^{-10} Torr which is equipped with X-ray photoemission spectroscopy (XPS; Riber Mg/Al dual anode with a Riber EA150 electron energy analyzer), low-energy electron diffraction (LEED; custom-made), infrared reflection absorption spectroscopy (IRAS; Bruker Tensor 37 with custom-design external IR optics and a tailor-made N₂ purge box, 4 cm⁻¹ resolution, 100 scans, 10 kHz scanner velocity, mercury cadmium telluride (MCT) detector), temperature-programmed desorption (TPD), and temperature-programmed reaction spectroscopy (TPRS) capabilities. A quadrupole mass spectrometer (QMS; Ametek Dycor Dymaxion DM200) and a proportional–integral–derivative (PID)-controlled linear sample heater (Heatwave, model 101303) were used for the TPD/TPRS experiments. All of the TPD/TPRS experiments were performed with a heating rate of 1 K/s, a 70 eV QMS beam energy, and a 30 ms dwell time for each desorption channel. The Ag(111) single-crystal sample (10 mm diameter × 1 mm thick disk, both sides polished, purity >99.999%, MaTeck GmbH) was affixed on Ta wires, through which the sample could be resistively heated to 1073 K. The cooling of the sample was achieved via a liquid nitrogen reservoir located inside the sample manipulator probe holding the Ag(111) single crystal. The temperature of the sample was measured using a K-type thermocouple spot-welded on the lateral facet of the Ag(111) disk. The Ag(111) sample surface was cleaned by cycles of Ar⁺ sputtering (Ar(g), Linde AG, purity $\geq 99.999\%$) with a sputtering gun (LK Technologies, NGI3000, 1.5 kV × 15 mA) at room temperature followed by annealing at 773 K in UHV for 20 min. After Ar⁺ sputtering, the Ag(111) sample was exposed to ozone (2.0×10^{-9} Torr × 5 min) at room temperature for further cleaning followed by annealing to 773 K for adsorbed oxygen removal. The cleanliness of the surface prior to experiments was confirmed by XPS and LEED. An Al K α X-ray source (1486.6 eV, 10 kV × 20 mA) was used in the XPS measurements. O 1s and Ag 3d regions were analyzed with an energy resolution of 0.1 eV for 10 scans and 5 scans, respectively. All of the XPS O 1s signals were positioned and normalized using the Ag 3d signals that were measured along with the O 1s signal.

Atomic oxygen was accumulated on the Ag(111) model catalyst surface by ozone exposure using a methodology whose details can be found in one of our former papers.⁵⁵ Ozone was dosed to the UHV system via Teflon tubings and a high-precision leak valve. NH₃ gas (Linde AG, purity $\geq 99.995\%$) was used without further purification and introduced onto the Ag(111) surface via a separate leak valve. In the rest of the text, exposures of adsorbate species (ϵ) are given in langmuirs (1 langmuir = 1×10^{-6} Torr·s) and the estimated surface coverages (θ) of the corresponding adsorbates are reported in monolayer equivalents (MLE).

■ RESULTS AND DISCUSSION

Oxygen on Ag(111). Before the investigation of SCO of NH₃ with oxygen, the behavior of oxygen species generated upon ozone decomposition on a clean Ag(111) single-crystal model catalyst surface was studied via TPD. For this purpose,

increasing exposures of ozone were introduced on Ag(111) at 140 K, and the TPD spectra of molecular oxygen ($m/z = 32$) desorbing from the Ag(111) single-crystal surface as a function of temperature were acquired as shown in Figure 1a. As can be

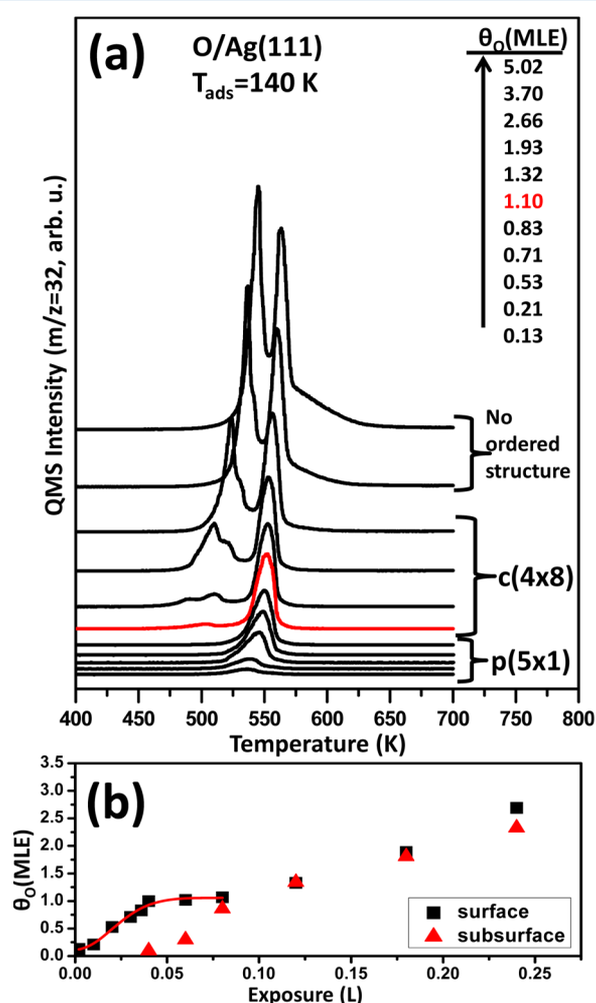


Figure 1. (a) TPD profiles for oxygen desorption ($m/z = 32$) obtained by increasing ozone exposures on clean Ag(111) at 140 K. (b) Relative oxygen surface coverages (obtained via integrated TPD desorption signals in part a) for surface and subsurface oxygen species on Ag(111) as a function of the ozone exposure.

seen in Figure 1a, oxygen desorption starts at 535 K for an oxygen surface coverage of $\theta_{\text{O}} = 0.13$ MLE, and the desorption temperature slowly increases with increasing oxygen surface coverage. Furthermore, various additional low-temperature ($T < 525$ K) desorption features (e.g., see the red spectrum in Figure 1a) start to appear and grow in intensity with increasing θ_{O} . These features can be associated with subsurface atomic oxygen (O_{sub}) as previously reported in the literature.^{26,42} These low-temperature features do not converge to a saturation, in agreement with a former paper,⁴² presumably due to the diffusion of atomic oxygen into the subsurface. Campbell et al. argued that oxygen atoms could be more stable on the silver surface rather than in bulk silver oxide on the basis of the heat of adsorption of oxygen atoms on Ag(111) as compared to the heat of formation of bulk silver oxide.²⁶ Therefore, bulk oxide formation due to oxygen diffusion to the silver subsurface seems to be rather unlikely at low oxygen coverages. However, formation of bulk silver oxide at high

oxygen coverages is feasible and will be discussed in more detail below. Along these lines, by exploiting isotopically labeled oxygen molecules, Backx et al. showed that subsurface oxygen in the Ag(110) surface desorbs at lower temperatures than the surface oxygen atoms.⁵⁶ TPD data for O/Ag(111) (Figure 1a) obtained via ozone decomposition are in good agreement with a former study, where NO_2 decomposition was utilized for atomic oxygen delivery on Ag(111).⁵⁰ Integrated TPD signals corresponding to surface (O_{a}) and subsurface (O_{sub}) oxygen species in Figure 1a were also calculated individually and plotted as a function of the ozone exposure (Figure 1b). In this latter plot, monolayer coverage reaches a plateau around 0.04 langmuir, which may be attributed to the saturation of the first overlayer on the surface. The absolute surface saturation coverage of the first oxygen monolayer on Ag(111) was previously calculated by Campbell et al.²⁶ for a (4 \times 4)-O/Ag(111) surface structure at this coverage. It is worth mentioning that, in the current work, the (4 \times 4)-O/Ag(111) structure was not observed in any of the investigated coverages, because this structure is not stable when oxygen is accumulated under UHV conditions.^{44,45} Nevertheless, in the current work, the saturation exposure of 0.04 langmuir was attributed to 1 MLE and was used to calibrate and quantify other desorption signals. As can be seen in Figure 1b, for the ozone exposures above 0.08 langmuir, surface oxygen coverage starts to exceed the convergence coverage of 1 MLE. This can be explained by the transformation of the surface oxygen overlayer to form a new structure with a higher oxygen density. We tentatively assign this latter structure to a bulklike silver oxide.

Analysis of the Ag(111) single-crystal surface after various ozone exposures at 140 K by LEED revealed no ordered structures, suggesting randomly located oxygen species at 140 K for all oxygen coverages. However, annealing of the O/Ag(111) surfaces originally prepared at 140 K indicated that thermal activation led to the ordered oxide formation with a p(5 \times 1) structure for $\theta_{\text{O}} < 1.10$ MLE (Figure 2a,b), as well as an additional oxide structure which can be tentatively assigned to a

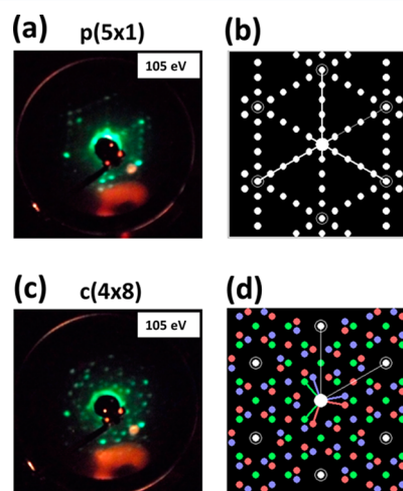


Figure 2. LEED images of O/Ag(111) overlayers obtained by introducing different ozone exposures at 140 K on a clean Ag(111) surface followed by pumping and annealing to 473 K in UHV. (a) LEED image of the p(5 \times 1)-O/Ag(111) overlayer ($\theta_{\text{O}} = 0.83$ MLE) and (b) its simulated (theoretical) LEED pattern. (c) LEED image of the c(4 \times 8)-O/Ag(111) overlayer ($\theta_{\text{O}} = 1.10$ MLE) and (d) its simulated (theoretical) LEED pattern.

c(4×8) structure appearing at $1.10 \text{ MLE} < \theta_{\text{O}} < 1.93 \text{ MLE}$ (Figure 2c,d). Oxide formation due to the annealing of oxygen overlayers on Ag(111) was also previously observed by Bukhtiyarov et al.,²⁹ where it was demonstrated via XPS that atomic oxygen adsorbed on Ag(111) at 300 K can form silver oxide when the sample is heated to 420 K. LEED images in Figure 2 were obtained by dosing different exposures of ozone on Ag(111) at 473 K (i.e., a temperature which is slightly below the beginning of the oxygen desorption) followed by pumping and cooling the sample to 140 K for LEED analysis. It is worth mentioning that ozone accumulation using relevant exposures at 140 K and heating to 473 K yielded LEED images identical to that of direct ozone dosing at 473 K. Starting from the lowest oxygen coverage (i.e., $\theta_{\text{O}} = 0.13 \text{ MLE}$) up to $\theta_{\text{O}} = 1.10 \text{ MLE}$, sharp spots of the p(5×1) structure were detectable after annealing at 473 K. For $\theta_{\text{O}} \geq 1.10 \text{ MLE}$, another ordered LEED structure started to appear after annealing at 473 K along with the emergence of the subsurface oxygen signals in TPD (red spectrum in Figure 1a). We tentatively assigned this structure to a c(4×8) overlayer consisting of three separate domains rotated by 120° with respect to each other. These domains are color-coded in the theoretical LEED pattern given in Figure 2d. c(4×8) reconstruction on O/Ag(111) was also observed by Schnadt and co-workers via scanning tunneling microscopy (STM) and was associated with a significantly oxidized silver structure.³⁸ In the current study, the c(4×8) structure was only observed at relatively high oxygen coverages (i.e., $1.10 \text{ MLE} < \theta_{\text{O}} < 1.93 \text{ MLE}$) where the surface oxygen species converged to a constant value (black curve in Figure 1b) while subsurface oxygen monotonically increased (red curve in Figure 1b). For $\theta_{\text{O}} > 1.93 \text{ MLE}$, LEED images became extremely fuzzy, suggesting the lack of an ordered structure at high coverages. This fuzzy structure may be associated with an amorphous bulklike oxide structure. Current results are in good agreement with the density functional theory (DFT) calculations of Li et al., who proposed that increasing oxygen coverages on the Ag(111) surface by ozone exposure might lead to ordered atomic oxygen overlayers followed by oxide-like thick films at higher θ_{O} .³³ In a related study, Martin et al. deposited atomic oxygen on Ag(111) by thermal cracking of $\text{O}_2(\text{g})$ at 500 K with $P_{\text{O}_2} = 10^{-7} \text{ mbar}$.⁴¹ They reported bulklike silver oxide formation at high θ_{O} and argued that the c(4×8) overlayer is the precursor state for bulklike silver oxide formation.

Oxygen accumulation on the Ag(111) surface at 140 K was also analyzed by XPS. O 1s spectra of the O/Ag(111) surface as a function of increasing oxygen coverage are given in Figure 3. Due to the detection limit of our photoelectron detector, we were able to observe a reliable O 1s signal at $\theta_{\text{O}} \geq 0.53 \text{ MLE}$. Signals in the region between 530 and 531 eV were assigned to atomic oxygen either on the surface (O_{a}) or in the subsurface (O_{sub}), whereas surface oxide (O_{ox}) signals were found between 528.1 and 528.5 eV.^{26–30,38–40,44,50,57} As mentioned above, ordered surface oxide (O_{ox}) species were not detectable at 140 K according to LEED experiments. Schlögl and co-workers previously reported that the O 1s signal of bulk silver oxide appeared at 529.0 eV.⁵⁸ In another study, Heine et al. assigned an O 1s signal at 528.9 eV to silver oxide species at the step edges of the Ag(111) surface⁴⁴ and suggested that subsurface oxygen increases the step density of the Ag(111) surface. In Figure 3, a noticeable increase in the O 1s signal at ca. 529 eV is visible at $\theta_{\text{O}} \geq 1.10 \text{ MLE}$ (i.e., in correlation with the

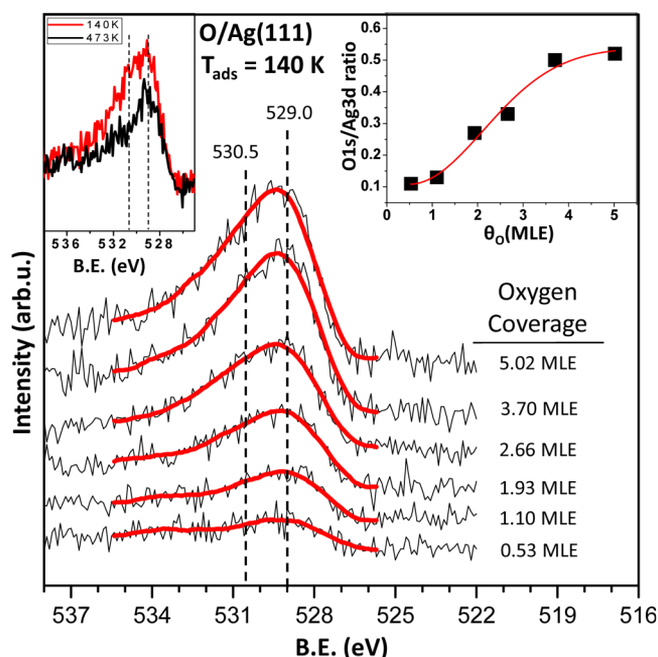


Figure 3. O 1s XPS spectra of O/Ag(111) as a function of increasing oxygen coverage obtained by increasing exposures of ozone at 140 K. The inset on the right shows the integrated O 1s/Ag 3d signal ratio as a function of the oxygen coverage. The inset on the left shows the O 1s XPS spectra for O/Ag(111) with $\theta_{\text{O}} = 2.66 \text{ MLE}$ dosed at 140 K (red) and dosed at 473 K (black).

appearance of the subsurface oxygen features in TPD data given in Figure 1a). This O 1s feature at ca. 529 eV may be associated with disordered oxide and/or amorphous bulklike oxide (O_{bulk}) species. Indications of the formation of bulklike silver oxide species were also present in the Ag 3d XPS spectra (Figure S1, Supporting Information), revealing a broadening in the full width at half-maximum (fwhm) of the Ag 3d peak with increasing oxygen coverages $\theta_{\text{O}} > 1.90 \text{ MLE}$. Broadening of the Ag 3d XPS signal due to the formation of silver oxide was also previously reported by Bukhtiyarov et al.²⁹

The inset on the right-hand side of Figure 3 shows the variation of the O 1s/Ag 3d integrated signal ratios in XPS as a function of oxygen coverage calculated from the ozone exposure values using the TPD data in Figure 1. It is apparent that the O 1s/Ag 3d integrated signal ratio reaches a convergence for $\theta_{\text{O}} \geq 3.70 \text{ MLE}$ (Figure 3) while the total $m/z = 32$ TPD desorption signal intensity still continues to increase within this coverage window (Figure 1b). These observations may indicate continuous transport of oxygen species toward the subsurface region (i.e., possibly below 10 nm of the surface) of the O/Ag(111) system at high oxygen coverages which cannot be readily detectable by the surface-sensitive XPS technique. It should be noted that X-ray-induced oxygen desorption during XPS analysis could also contribute to the high-coverage convergence of the O 1s/Ag 3d signal ratio.

The O 1s signal of the 2.66 MLE O/Ag(111) surface obtained at 140 K was also compared to the analogous surface prepared at 473 K using an identical ozone exposure. These results are shown in the left-hand-side inset of Figure 3 (red and black spectra, respectively). It is apparent that the O 1s spectrum corresponding to 473 K reveals attenuated features at ca. 529 and ca. 530.5 eV, which may be attributed to the partial loss of bulklike oxide (O_{bulk}) and surface/subsurface ($\text{O}_{\text{a}}/\text{O}_{\text{sub}}$)

atomic oxygen species, respectively. The O 1s feature at 529 eV corresponding to bulklike oxide species shows relatively less significant attenuation. These observations indicate transport of oxygen from the surface and subsurface regions of silver toward the bulk upon thermal activation provided by annealing at 473 K.

Other than the O 1s signals due to surface atomic oxygen (O_a) and bulklike oxide (O_{bulk}) species, a less prominent shoulder at 532 eV is also detectable in the O 1s spectra presented in Figure 3 which can be associated with hydroxide (minority) species due to background water adsorption. Note that the presence of physisorbed oxygen species can be readily excluded due to the extremely low desorption temperature of such species (i.e., ~ 50 K).²⁶

Ammonia on Ag(111). Ammonia adsorption was carried out on the clean Ag(111) surface at 90 K, and the $m/z = 17$ signal was monitored in TPD experiments along with the $m/z = 28$, 18, and 2 signals to check the presence of potential decomposition products and water desorption from background adsorption (Figure 4). In all of the TPD experiments,

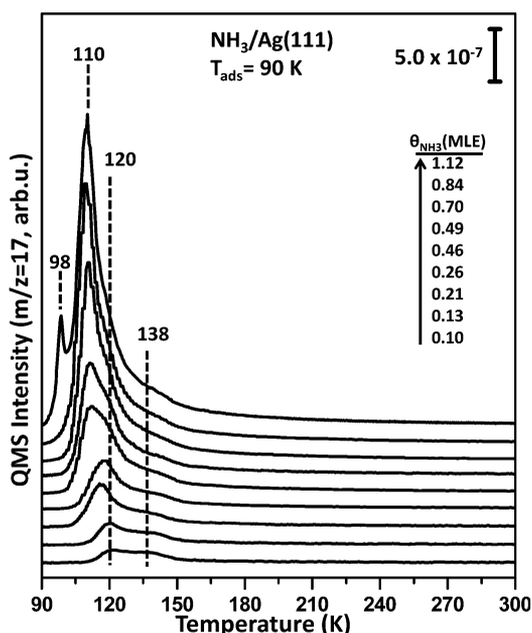


Figure 4. TPD profiles ($m/z = 17$) obtained after ammonia adsorption on a clean Ag(111) surface at 90 K as a function of increasing ammonia coverage.

ammonia was found to desorb molecularly in a reversible manner from the clean Ag(111) single-crystal surface, where no ammonia decomposition products, namely, N_2 and H_2 desorption, were detectable at any ammonia coverage. Limited background water desorption was observed, and TPD spectra for the $m/z = 18$ channel are given in the Supporting Information (Figure S2). In Figure 4, two ammonia desorption maxima located at 138 and 120 K were observed for the lowest ammonia coverage of $\theta_{\text{NH}_3} = 0.10$ MLE. With increasing ammonia coverage, the desorption signal at 120 K became broader and its desorption maximum shifted to lower temperatures. Broadening and the low-temperature shift with increasing ammonia coverage in the submonolayer range are common behaviors of ammonia TPD profiles obtained from numerous metal surfaces such as Pt(111),⁵⁹ Au(111),⁶⁰ Rh(111),⁶¹ and Ir(100)⁶² which can be attributed to the

heterogeneity of the adsorbed ammonia species, as well as repulsive interactions (or decreasing attractive interactions) between the adsorbates in the first monolayer.⁶³ Figure 4 shows that the major ammonia desorption feature at 110 K reached saturation, which was immediately followed by the appearance of an additional desorption maximum at 98 K that can be assigned to the formation of multilayer ammonia overlayers. TPD data given in Figure 4 can be used to estimate the relative ammonia surface coverages. For this purpose, the multilayer signal at 98 K was deconvoluted and subtracted from the whole desorption spectrum and the integrated desorption signal between 90 and 480 K associated with the desorption maximum at 110 K was used to estimate $\theta_{\text{NH}_3} = 1$ MLE. It is worth mentioning that TPD experiments were also performed for ammonia coverages greater than 1.12 MLE, revealing a typical zeroth-order desorption kinetics (data not shown). In the literature, analogous monolayer and multilayer ammonia desorption signals were also observed on other metal surfaces^{61,62} as well as on Ag(110).⁶⁴ The less prominent ammonia desorption signal at 138 K in Figure 4 can be attributed to stronger ammonia adsorption on surface defects and/or adsorption of ammonia with dissimilar geometries/orientations on different surface adsorption sites. Overall, TPD results for NH_3 adsorption on clean Ag(111) given in Figure 4 clearly point to the fact that, in the absence of oxygen species, Ag(111) is incapable of activating NH_3 . On the other hand, as will be shown below, ammonia activation and its selective catalytic oxidation are achievable by fine-tuning the nature and surface coverage of oxygen species on Ag(111).

IRAS spectra corresponding to the adsorption of ammonia on clean Ag(111) for various coverages of ammonia at 90 K are given in Figure 5. For the lowest coverage of ammonia ($\theta_{\text{NH}_3} =$

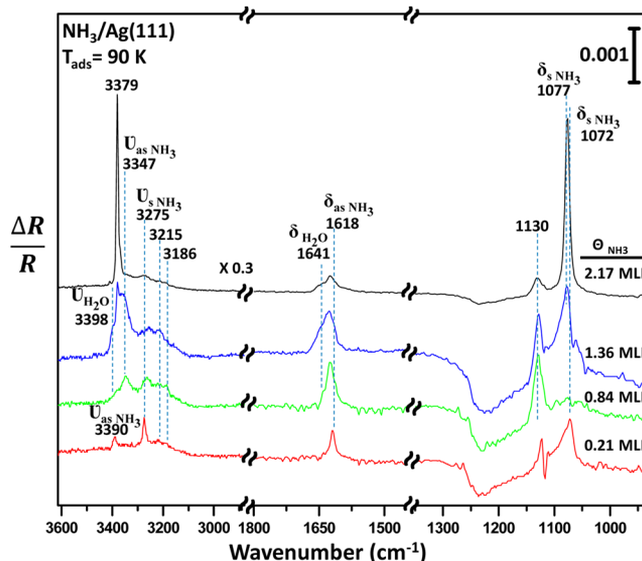


Figure 5. IRAS spectra of ammonia adsorption on a clean Ag(111) surface at 90 K as a function of the ammonia surface coverage.

0.21 MLE), symmetric (δ_s) and antisymmetric (δ_{as}) NH_3 deformation signals appeared at 1072 and 1618 cm^{-1} , respectively; while symmetric (ν_s) and antisymmetric (ν_{as}) NH_3 stretching modes can be seen at 3275 and 3390 cm^{-1} , respectively.⁶⁴ For submonolayer coverages, it is commonly observed that ammonia molecules are typically adsorbed on transition-metal surfaces through unpaired electrons of nitrogen

and the molecular axis stays perpendicular to the surface.⁶⁵ In such cases, δ_{as} and ν_{as} modes cannot be observed as reported in the literature for ammonia adsorption on Ni(111),⁶⁶ Ru(0001),⁶⁷ Pt(111),⁶⁸ and Cu(110).⁶⁹ In contrast, IRAS data in Figure 5 suggest that, on a clean Ag(111) surface, the molecular axis of ammonia is tilted with respect to the Ag(111) surface for submonolayer coverages. A similar behavior was also observed for ammonia adsorption on clean Ag(110) via electron energy loss spectroscopy (EELS).⁶⁴ For the IRAS spectrum corresponding to $\theta_{\text{NH}_3} = 0.84$ MLE, the δ_{s} signal was found to vanish and the NH_3 stretching modes were observed to broaden and shift to lower frequencies, namely, to 3347 and 3266 cm^{-1} . These changes in the IRAS spectra are probably due to increasing intermolecular interactions between adsorbed ammonia molecules on Ag(111) with increasing surface coverages as well as due to the formation of hydrogen-bonding interactions. Loss of the δ_{s} feature was also reported on $\text{NH}_3/\text{Ru}(0001)$ by Rodriguez et al. for the second layer (i.e., multilayer) of ammonia on Ru(0001) and was attributed to the formation of intermolecular hydrogen bonding and/or a smaller IR absorption cross section of the ammonia molecules in the second layer.⁶⁷

The IRAS spectrum corresponding to $\theta_{\text{NH}_3} = 0.84$ MLE also revealed water stretching and bending signals due to background water adsorption located at 3398 and 1641 cm^{-1} , respectively.⁷⁰ Furthermore, the IRAS signals at 3186 and 3215 cm^{-1} might be related to hydronium ions and hydrogen-bonded water molecules located on the surface or on the ammonia overlayer with different orientations.⁷⁰ When the ammonia coverage was increased to 1.36 MLE, the symmetric deformation mode (δ_{s}) at 1072–1077 cm^{-1} reappeared and the antisymmetric (ν_{as}) NH_3 stretching signal became broader. Furthermore, at $\theta_{\text{NH}_3} = 2.17$ MLE, an additional sharp signal appeared at 3379 cm^{-1} which can be assigned to the formation of solid multilayer ammonia.⁶⁸ These results are in very good agreement with the former reports in the literature for multilayer ammonia adsorption on various single crystals.^{67,71} Vibrational frequencies of NH_3 on Ag(111) for different NH_3 coverages obtained by the currently presented IRAS spectra are summarized in Table 1. IRAS data given in Figure 5 for $\text{NH}_3/\text{Ag}(111)$ are in agreement with the corresponding TPD data presented in Figure 4, suggesting molecular adsorption (i.e.,

Table 1. Vibrational Frequency Assignments for the $\text{NH}_3/\text{Ag}(111)$ IRAS Data Presented in Figure 5

	$\delta_{\text{s}}(\text{HNNH})$ (cm^{-1})	$\delta_{\text{as}}(\text{HNNH})$ (cm^{-1})	$\nu_{\text{s}}(\text{NH})$ (cm^{-1})	$\nu_{\text{as}}(\text{NH})$ (cm^{-1})
NH_3 on Ag(111) ($\theta_{\text{NH}_3} = 0.21$ MLE)	1072	1618	3275	3390
NH_3 on Ag(111) ($\theta_{\text{NH}_3} = 0.84$ MLE)		1640	3266	3347
NH_3 on Ag(111) ($\theta_{\text{NH}_3} = 1.36$ MLE)	1077	1625	3257	3379
NH_3 on Ag(111) ($\theta_{\text{NH}_3} = 2.17$ MLE)	1077	1651	3275	3379
NH_3 on Ag(110) (submonolayer) ⁶⁴		1651		
NH_3 on Ag(110) (submonolayer) ⁶⁴	1050	1640	3320	3400
solid NH_3 ⁶⁴	1060	1646	3223	3378

lack of dissociation) of ammonia on Ag(111) in the absence of additional oxygen species.

Ammonia SCO on O/Ag(111). The ammonia SCO reaction was carried out on O/Ag(111), and the resulting products were followed in TPRS experiments. To prevent poisoning of the O/Ag(111) surface with excess ammonia, NH_3 was dosed on the O/Ag(111) at 250 K (i.e., at a temperature above the desorption maximum of the first monolayer of ammonia on clean Ag(111)). During TPRS experiments, $m/z = 17$ (NH_3), 28 (N_2), 30 (NO), 44 (N_2O), 46 (NO_2), 32 (O_2), 18 (H_2O), and 2 (H_2) desorption channels were simultaneously monitored.

Two different sets of TPRS spectra for the oxygen–ammonia reaction are presented in Figure 6, where the main difference

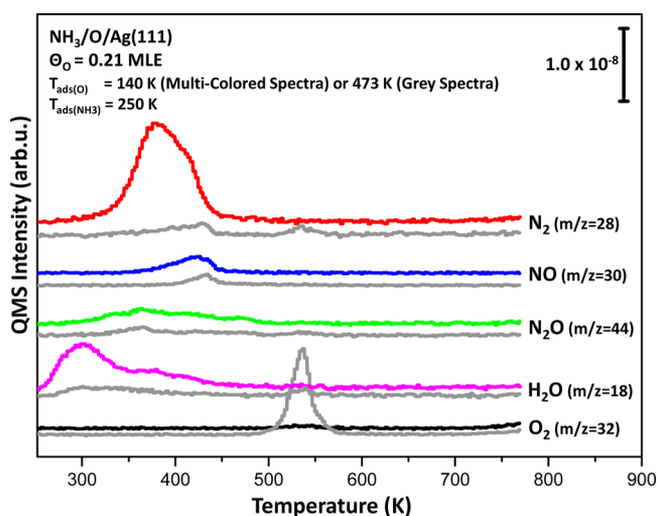


Figure 6. TPRS profiles for the reaction of ammonia with oxygen on Ag(111). In all of the experiments, an identical exposure of NH_3 ($P_{\text{NH}_3} = 2.0 \times 10^{-9}$ Torr \times 60 s) was dosed at 250 K on an oxygen-covered ($\theta_{\text{O}} = 0.21$ MLE) Ag(111) surface, where oxygen was delivered by ozone exposure either at 140 K (multicolored spectra) or at 473 K (gray-colored spectra).

between these different sets of spectra is the temperature of oxygen delivery. While the oxygen delivery at 140 K (multicolored spectra) led to surface atomic oxygen (O_{a}) species on Ag(111), oxygen delivery at 473 K (gray spectra) resulted in the formation of a $p(5 \times 1)$ surface oxide.

Multicolored TPRS spectra in Figure 6 corresponding to the presence of surface atomic oxygen (O_{a}) species on Ag(111) obtained via oxygen delivery at 140 K revealed the obvious reaction between surface oxygen species and ammonia yielding SCO products such as N_2 (red), NO (blue), and N_2O (green) in a wide temperature range between 300 and 450 K. In addition, water desorption within the same temperature range was also in line with the N–H bond activation and hydrogen abstraction from ammonia by surface oxygen atoms on Ag(111). Neither unreacted oxygen (black spectrum in Figure 6) nor NO_2 formation (see Figure S3, Supporting Information) were detectable for the multicolored TPRS data. Molecular ammonia desorption was not also observed as a result of the NH_3 dosage temperature (250 K), which is above the desorption temperature of the first monolayer of ammonia on clean Ag(111) (see Figure S3).

On the other hand, for the gray-colored TPRS data given in Figure 6 corresponding to the interaction of ammonia with the

p(5×1) oxide overlayer (confirmed by LEED), a very limited extent of reaction between ammonia and oxygen was observable. The strikingly intense desorption signal of the unreacted oxygen species at 537 K was consistent with the limited ammonia oxidation by the oxide overlayer. A similar behavior was also reported by Madix and co-workers for CO oxidation on Ag(111), suggesting that the silver oxide overlayer was not the most reactive phase for the CO oxidation reaction.⁷² Thus, comparative analysis of the TPRS data shown in Figure 6 pointed to the fact that surface atomic oxygen (O_a) on Ag(111) could readily activate N–H bond cleavage in NH_3 , while ordered silver oxide (O_{ox}) overlayers had limited activity for ammonia oxidation.

NH_3 SCO reactions on the Ag(111) single-crystal surface containing different quantities of oxygen (delivered by ozone exposure at 140 K) were performed, and the corresponding TPRS spectra of the selected products (i.e., N_2 , NO, and N_2O) are given in Figure 7. Alternative depictions of the same set of data are also provided in Figure S4 and S5 (Supporting

$NH_3/O/Ag(111)$ $T_{ads(O)} = 140$ K
 $T_{ads(NH_3)} = 250$ K

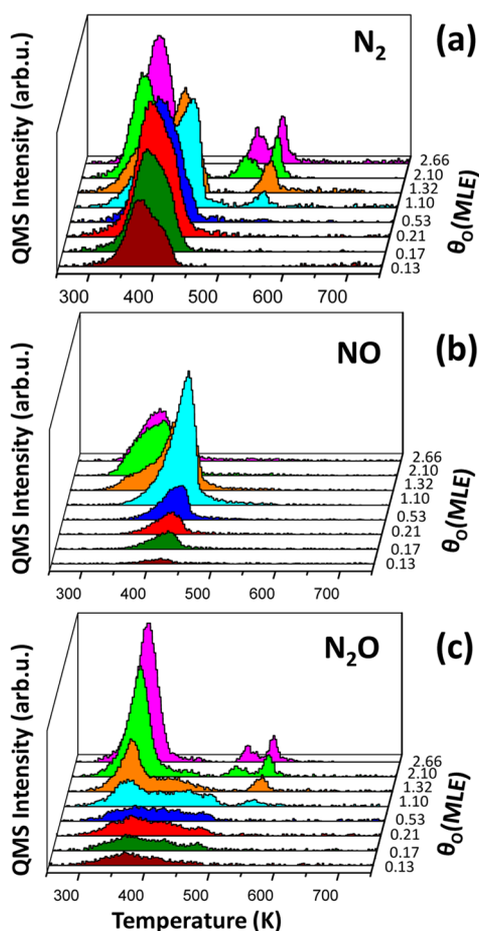


Figure 7. TPRS profiles of various ammonia SCO reaction products (i.e., N_2 , NO, and N_2O) for the reaction between ammonia and oxygen-precovered Ag(111) surfaces as a function of the oxygen coverage. For all of the data, oxygen was delivered on Ag(111) by ozone exposure at 140 K, while NH_3 was delivered using an identical ammonia exposure at 250 K ($P_{NH_3} = 2.0 \times 10^{-9}$ Torr \times 60 s).

Information). Integrated desorption signals of the relevant products in Figure 7 were also calculated (after compensating for the interferences between various desorption signals due to mass spectroscopic fragmentation). These results are presented as histograms in Figure 8, revealing relative ammonia SCO

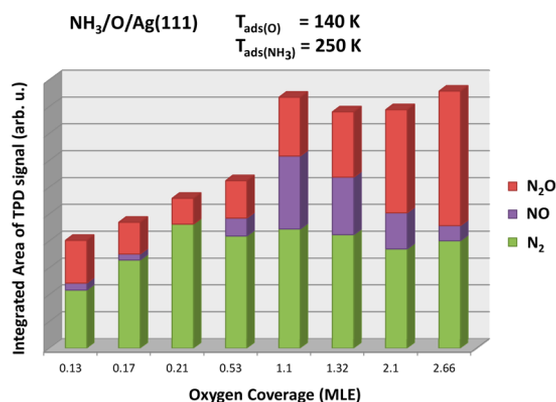


Figure 8. Integrated TPD desorption signals for the main ammonia SCO products (i.e., N_2 , NO, and N_2O) on an O/Ag(111) surface as a function of the oxygen coverage extracted from the TPRS data given in Figure 7. See the text for details.

selectivity for dissimilar oxygen coverages. To calculate total integrated TPRS signals of N_2O in Figure 8, mass spectroscopic fragments of N_2O other than $m/z = 44$ were also considered. By referring to the National Institute of Standards and Technology (NIST) mass spectroscopy database,⁷³ relative intensities of the main fragments of N_2O were obtained as 100%, 32%, and 11% for $m/z = 44$, 30, and 28, respectively. Accordingly, the integrated $m/z = 44$ desorption signal was multiplied by a factor of 1.43. On the other hand, the contributions of N_2O to the NO and N_2 integrated signals were subtracted (removed) from the integrated $m/z = 30$ and 28 signals. As can be seen in Figures 7 and 8, the main ammonia SCO product for low oxygen coverages (i.e., $0.13 \text{ MLE} < \theta_O < 1.10 \text{ MLE}$) was clearly N_2 and the maximum N_2 formation was obtained for $\theta_O = 0.21 \text{ MLE}$. It is also apparent that NO selectivity increased drastically for $\theta_O = 1.10 \text{ MLE}$ and became comparable to that of N_2 .

It is worth mentioning that the N_2 desorption maxima shifted to higher temperatures for $\theta_O \geq 1.10 \text{ MLE}$ (Figure 7a and Figure S4a, Supporting Information) and additional N_2 desorption states also became visible. This observation was concomitant with the appearance of excess oxygen desorption associated with surface and subsurface/bulk oxide states (Figure S5, Supporting Information). Thus, it can be argued that formation of oxide species abundant in oxygen led to an attractive interaction between oxygen species and atomic nitrogen, resulting in the stabilization of the atomic nitrogen species generated by N–H bond activation of NH_3 and a shift in the N_2 desorption maxima to higher temperatures. Furthermore, generation of additional high-temperature N_2 desorption states at elevated oxygen coverages can also be attributed to the increasing heterogeneity of the adsorption sites of the substrate surface due to the formation of bulklike silver oxide.

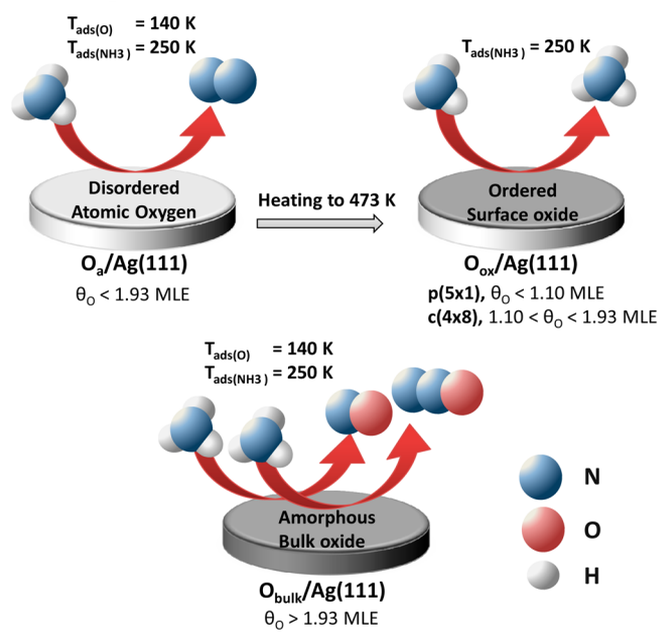
Furthermore, unreacted NH_3 was also monitored (Figure S6, Supporting Information) for the set of experiments presented in Figure 7. These results revealed limited amounts of ammonia desorption (i.e., a lack of the presence of excess ammonia),

suggesting a high conversion of ammonia for the particular set of experimental conditions chosen for Figure 7.

It is worth mentioning that IRAS measurements were also performed at 250 K for SCO of ammonia on the atomic oxygen (O_a) covered Ag(111) surface using experimental conditions (coverages) similar to the ones chosen for the TPRS experiments given in Figure 7. However, no IRAS signals were detected, possibly due to the unfavorable adsorption geometry or small IR absorption cross sections of the reactants and products lying possibly close to parallel to the surface.

The overall reactivity and selectivity trends of ammonia SCO on O/Ag(111) as a function of the adsorption temperature, oxygen surface coverage, and nature of the oxygen species deduced from the currently presented experimental findings are summarized in Scheme 1.

Scheme 1. Reactivity and Selectivity Trends of Ammonia SCO on O/Ag(111) as a Function of the Oxygen Coverage and Temperature



CONCLUSIONS

In the current study, SCO of ammonia on O/Ag(111) was investigated by utilizing a multitude of surface-sensitive spectroscopic and diffraction techniques where a unique oxygen delivery agent (i.e., ozone) was utilized under UHV conditions, eliminating the need for high-pressure O_2 exposures. TPD/TPRS, XPS, and LEED data suggest that a variety of oxygen species can be generated on the Ag(111) surface upon ozone adsorption, such as atomic (chemisorbed) oxygen, surface oxide, and bulklike oxide species. Indications of oxygen transport to the subsurface region of silver at elevated ozone exposures were also detected. Oxygen overlayers prepared via ozone exposure at 140 K in UHV led to disordered overlayers (in the form of either atomic oxygen or bulklike silver oxide) for all oxygen coverages, while annealing of these overlayers for intermediate oxygen coverages at 473 K resulted in the formation of ordered surface oxides with $p(5 \times 1)$ and $c(4 \times 8)$ structures. The clean Ag(111) surface was found to be incapable of activating N–H bonds of ammonia under UHV conditions in the absence of ozone exposure. Disordered

atomic oxygen overlayers prepared on Ag(111) at 140 K for $\theta_{\text{O}} < 1.10 \text{ MLE}$ were found to be highly selective in SCO of ammonia, producing N_2 as the dominant product. On the other hand, ordered $p(5 \times 1)$ and $c(4 \times 8)$ surface oxide overlayers on Ag(111) prepared via ozone adsorption at 473 K were found to be almost entirely inactive in SCO where NH_3 and O_2 desorbed in an intact manner in TPRS experiments. It was observed that extreme oxygen exposures on Ag(111) ($\theta_{\text{O}} > 1.93 \text{ MLE}$) at 140 K led to bulklike silver oxide species with poor N_2 selectivity in ammonia SCO and increasing extent of formation of toxic pollutants such as NO and N_2O . Currently presented experimental findings on the Ag(111) single-crystal model catalyst surface reveal valuable insights as well as new structure–functionality relationships at the molecular level regarding the reactivity of oxygen species in ammonia SCO. These findings may also be relevant to other crucial heterogeneous catalytic oxidation/partial oxidation/oxidative coupling processes of CO, alkenes, and alcohols.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcc.7b08291.

Ag 3d XPS spectra for clean Ag(111), $O_a/\text{Ag}(111)$, and $O_{\text{bulk}}/\text{Ag}(111)$, TPD profiles for the H_2O ($m/z = 18$) desorption channel for the data given in Figure 4, TPRS profiles for the NO_2 ($m/z = 46$) and NH_3 ($m/z = 17$) desorption channels for the data given in Figure 6, alternative depiction of the TPRS profiles of various ammonia SCO reaction products given in Figure 7, and TPRS profiles for the O_2 ($m/z = 32$), NH_3 and OH ($m/z = 17$), and H_2O ($m/z = 18$) desorption channels for the data given in Figure 7 (PDF)

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Notes

The authors declare no competing financial interest.

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