Insights into the function of silver as an oxidation catalyst by ab initio atomistic thermodynamics

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To help understand the high activity of silver as an oxidation catalyst, e.g., for the oxidation of ethylene to epoxide and the dehydrogenation of methanol to formaldehyde, the interaction and stability of many different oxygen species at the Ag(111) surface has been studied for a wide range of coverages. Through calculation of the free energy, as obtained from density-functional theory and taking into account the temperature and pressure via the oxygen chemical potential, we obtain the phase diagram of O/Ag(111). Our results reveal that a thin surface-oxide structure is most stable for the temperature and pressure range of ethylene epoxidation and we propose it (and possibly other similar structures) contains the species activating the catalysis. For higher temperatures, low coverages of chemisorbed oxygen are most stable, which could also play a role in oxidation reactions. For temperatures greater than about 775 K there are no stable oxygen species, except for the possibility of O atoms adsorbed at undercoordinated surface sites (i.e., imperfections, defects). At low temperatures (≤400 K at atmospheric pressure), provided kinetic limitations can be overcome, thicker oxidoleike structures are predicted. Due to their low thermal stability, however, they can be ruled out as playing an important role in the heterogeneous reactions under technical conditions. Bulk dissolved oxygen and a molecular ozonelike species adsorbed at a surface vacancy, as have been proposed in the literature, are found to be energetically unfavorable. The employed theoretical approach for calculating free energies and predicting the lowest energy structures in contact with species in the environment (“ab initio, atomistic thermodynamics”), affords investigation of a system that seamlessly connects standard (T = 0 K) density-functional theory results, characteristic of “typical” theoretical surface science studies, through to those valid for the conditions of catalysis. Though the error bar of the noted theoretical temperatures is noticeable (± = 55 K), the identified trends and physical descriptions are useful.

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I. INTRODUCTION

The identification of active species for desired chemical reactions is a main quest in the field of heterogeneous catalysis.¹ Whether a species is active or not depends on many factors, including the nature of its interaction with the catalyst material, as well as the temperature and pressure at which the catalytic reactions take place. The main approach for gaining quantitative, microscopic understanding of various properties and processes, has been to perform experiments under standard surface science conditions, i.e., under ultrahigh vacuum (UHV) and relatively low (room or lower) temperatures, with the hope that what is learnt may be relevant and/or extrapolated to the system under the “real” catalytic conditions of high temperatures and pressures. Theoretical studies of adsorption and reactions at surfaces have, at best, been carried out by first-principles approaches, which, while highly accurate, typically do not take into account the temperature or the gas-phase environment with which the surface is in contact. A recent focus, both experimentally and theoretically, is to attempt to study systems which more closely resemble those of true catalytic ones.² Experimentally, this is done by performing in situ real-time measurements carried out under high temperature and pressure conditions, and by the systematic study of, e.g., well-characterized metal clusters deposited on support materials and chemical reactions thereon (see, e.g., Refs. 3–8). Such studies are important in particular for understanding those systems which exhibit a “pressure and/or materials gap,” i.e., where different behavior is exhibited under low and high pressures, or when results for model single-metal catalysts do not represent those of the “real” catalyst, typically consisting of a dispersed metal on a support material. Often the pressure and materials gaps are related and occur simultaneously.

The oxygen-silver system is an example which exhibits a pronounced pressure-dependence: Despite its unique activity as the catalyst for ethylene epoxidation (C₂H₄ + ½O₂: C₂H₂O, performed at 500–600 K and atmospheric pressure) and partial oxidation of methanol to formaldehyde (CH₃OH + ½O₂: CH₂O + H₂O—“oxydehydrogenation” or by direct dehydrogenation, CH₃OH: CH₂O + H₂, performed at 700–900 K and atmospheric pressure), these reactions do not occur with any significant probability under UHV conditions.³⁻¹ This behavior of the oxygen-silver system might be related to a materials gap, because for studies of the partial oxidation of methanol, drastic differences are observed in the performance for fresh silver catalysts and those which have been exposed to high temperature (T > 873 K) and pressure (atmospheric) conditions. The temperature dependence of the conversion of the clean Ag material to the active catalyst exhibits a pronounced hysteresis, i.e., the conversion is different when raising the temperature from low to high, compared to from high to low. This was the case for high temperatures up to 873 K, as well as for lower temperatures up to 560 and 750 K. These hysteresis effects were shown from scanning electron microscopy (SEM) studies to be related to significant morphological...
changes which occur in the catalyst depending on the temperature, and that do not revert to the original structure when lowering the temperature, i.e., the changes are irreversible. Structural changes in silver also occur when in contact with a pure $O_2$ environment at high temperatures ($\sim 900 \text{ K}$) and atmospheric pressure, namely, facets with the Ag(111) orientation result. This is presumably because under these conditions it is unlikely that there will be any oxygen on the surface and the (111) surface of Ag has the lowest surface energy.

It has been suggested that the elusive subsurface or “bulk-dissolved” oxygen species play an essential role in the abovementioned catalytic reactions, which so far have escaped a precise characterization. In addition to this aspect, there are other controversial debates in the literature concerning the nature of the active species, e.g., atomic versus molecular (ozonelike) oxygen. Another puzzling issue is how silver, as a noble metal, can function as a good catalyst when it only binds adparticles relatively weakly on the surface compared to the transition-metal catalysts to the left of it in the Periodic Table. Catalysis clearly involves a number of complex processes, namely, the dissociation of molecules and the creation of chemically active species, and subsequent interaction and reaction between the particles to form the product, which desorbs from the surface. Furthermore, it is possible (maybe likely) that some fragments of the reactant molecules (e.g., H or C) modify the catalyst material and thereby participate in the creation of the active species. In the present paper we tackle only part of the problem, namely, the study of the interaction of silver with oxygen in order to identify and exclude possible active O species for the abovementioned heterogeneous catalytic reactions. This is an essential prerequisite for modeling the full catalytic process.

In our previous publications, the interaction between oxygen and Ag(111) was thoroughly studied by density-functional theory (DFT). In these works we considered almost all possibly relevant oxygen species, namely, pure on-surface chemisorbed oxygen, pure sub-surface and bulk-dissolved oxygen, surface- and bulk-substitutional adsorption, structures involving both oxygen on and under the top Ag layer, as well as surface oxides and a molecular (ozonelike) species adsorbed at a surface vacancy. We studied these oxygen species for a wide range of coverages where the following results were obtained: At low coverages oxygen atoms adsorb on the surface. With increasing coverages, however, strong repulsive interactions develop between the partially negatively charged O atoms and the adatoms are driven to penetrate into the subsurface region, and/or to give rise to a reconstructed, thin $(4 \times 4)$ surface-oxide layer, when the coverage is larger than $0.25 \text{ ML}$ where the latter structure has a stoichiometry of $\text{Ag}_1\text{S}_1\text{O}$ in the top layer. For higher oxygen contents, thicker oxidelike films are predicted to form.

We also found that the presence of subsurface oxygen, when bonded to the same surface Ag atoms as an oxygen atom on the surface, dramatically modifies the electronic properties of the on-surface oxygen and vice versa. Depending on the particular geometry, it can give rise to a notable stabilization or destabilization, but the energetically favorable structures induce a stabilization and a lowering of the density of states at the Fermi level. In contrast to the conclusions concerning the “crucial role” of bulk-dissolved oxygen based on interpretations of experimental results, we found that this species is energetically unfavorable, as is bulk- and surface-substitutional oxygen (in which the O atom takes the site of the Ag atom), as well as the molecularly adsorbed ozonelike species at a vacancy, when the system is in thermal equilibrium. In these structures we assume that the displaced Ag atoms are readorsbed at kink sites at step edges where they gain the cohesive energy. On the other hand, the binding energy of atomic oxygen adsorbed at undercoordinated silver atoms is considerably stronger than on the terraces, but the concentration of such species obviously depends on the number of defects and the morphology of the substrate.

Nevertheless, temperature and pressure can affect the stability of structures and this has not been investigated so far. Clearly this is important for gaining an understanding of the system under catalytic conditions and this is the main topic of the present paper. Some of these results were reported briefly in Ref. 22 and are explained in detail here. Although we study explicitly the (111) surface of silver, our general qualitative findings may be expected to be relevant for silver per se. In particular, for oxygen adsorption on Ag(110), a series of $(n \times 1)$ structures form, which exhibit a similar local O-Ag-O bonding arrangement to the thin $(4 \times 4)$ surface-oxide-like structure that forms on the Ag(111) surface, where the O atoms are bonded to surface Ag atoms which are also bonded to one other O atom. The temperature programmed desorption features of these phases are also similar. In addition, for O/Ag(001), a $(\sqrt{2} \times \sqrt{2})$ structure forms, where again the O atoms bond to surface Ag atoms which are also bonded to one other O atom. Furthermore, similarly to the results we have obtained for O on Ag(111), it has been found from DFT calculations of O/Ag(001) (Refs. 28,29) that for low coverages, the O atoms tend to repel each other and adsorb in the hollow sites, while for higher coverages the $(\sqrt{2} \times \sqrt{2})$ phase forms which could be likened to a thin surface-oxide-like structure. Clearly, on the quantitative level, there are differences between O adsorption on the various Ag crystallographic faces (i.e., detailed atomic structure and adsorption energy), but the general behavior regarding formation of on-surface oxygen, thin surface-oxide, and bulk oxide formation may be similar.

Our results will possibly also have implications for gold, another noble metal oxidation catalyst which, for the (111) surface, exhibits a restructuring and chemisorption of oxygen atoms at elevated temperatures (500–800 K) and atmospheric pressure, similar to Ag(111). They may also relate to copper, which catalyses the oxidation of methanol to formaldehyde.
oxygen chemical potential (see, e.g., Refs. 34–39). From the resulting free energies (obtained using our previously obtained energetics\textsuperscript{20,31}), we derive the phase diagram of oxygen at the silver (111) surface. Since our earlier studies\textsuperscript{20,21} predict that oxidelike structures form for high O concentrations, we also investigate the surface and bulk properties of the most stable bulk oxide, namely, di-silver oxide Ag$_2$O.

The paper is organized as follows. The calculation procedure, as well as the thermodynamical method are explained in Sec. II, and in Sec. III results are presented for the bulk properties of Ag$_2$O. In Sec. IV, we investigate various terminations of the Ag$_2$O(111) surface as a function of the oxygen chemical potential. Section V describes investigations into the transition from oxidelike structures [where the positions of the Ag atoms are commensurate with those of the Ag(111) substrate] to the true oxide film which is laterally expanded in comparison. In Sec. VI, the pressure-temperature phase diagram of the oxygen-silver system is presented and compared to experimental results. Finally, the conclusions are given in Sec. VII.

II. CALCULATION METHOD

The DFT total-energy calculations are performed using the pseudopotential plane wave method\textsuperscript{40} within the generalized gradient approximation (GGA).\textsuperscript{41,42} The pseudopotentials are generated by the scheme of Troullier and Martins with the same functional.\textsuperscript{43,44} The wave functions are expanded in plane waves with an energy cutoff of 50 Ry. In the (1 $\times$ 1) surface unit cell, corresponding to the periodicity of clean Ag(111), 21 special $k$ points are used in the surface irreducible Brillouin zone (IBZ) for the Brillouin-zone integration.\textsuperscript{45} Equivalent $k$ points to these are used for all of the surface structures studied for consistency, i.e., to maximize the accuracy when comparing the energetics of different coverages as calculated in different supercells. We employ a Fermi function with a temperature broadening parameter of $k_B T_{\text{f}} = 0.1$ eV to improve the convergence, and the total energy is extrapolated to zero temperature. ($k_B$ is the Boltzmann constant.) The structures are created on one side of the slab and the resulting induced dipole moment is taken into account by applying a dipole correction.\textsuperscript{46} For the adsorption structures on Ag(111), we use five layers of Ag to model the surface and typically keep the bottom two or three layers fixed at the bulklike positions and fully relax all the other atoms until the forces on the atoms are less than 0.015 eV/Å. Relaxation of two metal layers is completely sufficient for O adsorption on the surface, but it is necessary to relax three layers for the structures involving substitutional or sub-surface O adsorption. The oxide surfaces of Ag$_2$O(111) are simulated by symmetrical slabs (i.e., with inversion symmetry). We tested thicknesses corresponding to five, seven, and nine metal layers, separated by 15 Å of vacuum. Our results show that the systems can be safely described using five metal layers with inversion symmetry.

In \textit{ab initio} theory, the consideration of high temperature and high pressure can be achieved by explicitly taking into account the surrounding gas phase in terms of \textit{“ab initio, atomistic thermodynamics”} (see, e.g., Refs. 34–39). This is also an appropriate (first) approach to steady-state catalysis, which is often run close to thermodynamic equilibrium (or a constrained equilibrium) to prevent catalyst degradation. In the following we outline how the combination of thermodynamics and DFT can be applied to obtain the lowest-energy surface structures with a surrounding gas phase, thus enabling us to construct a ($T, p$) diagram of the stability (or metastability) regions of different surface phases.

We consider a surface in contact with an oxygen atmosphere which is described by an oxygen pressure $p$ and a temperature $T$. The environment then acts as a reservoir, as it can give (or take) oxygen to (or from) the substrate without changing the temperature or pressure. We calculate the surface free energy

$$\gamma(T, p) = (G - N_{\text{Ag}} \mu_{\text{Ag}} - N_{\text{O}} \mu_{\text{O}})/A,$$

(1)

where $N_{\text{O}}$ and $N_{\text{Ag}}$ are the number of oxygen and silver atoms, and $A$ is surface area. The $T$ and $p$ dependence is mainly given by $\mu_{\text{O}}$, the oxygen chemical potential, i.e., by the $O_2$ gas phase atmosphere

$$\mu_{\text{O}}(T, p) = \frac{1}{2} \left[ E_{\text{total}} + \mu_{\text{O}_2}(T, p^0) + k_B T \ln \frac{p}{p^0} \right],$$

(2)

where $p^0$ corresponds to atmospheric pressure and $\mu_{\text{O}_2}(T, p^0)$ includes the contribution from rotations and vibrations of the molecule, as well as the ideal-gas entropy at 1 atmosphere. It can be calculated or taken from experimental values from thermodynamic tables—the difference between theory and experimental values is marginal in the temperature range of interest. The silver chemical potential is

$$\mu_{\text{Ag}} = \begin{cases} g_{\text{Ag-bulk}}, & \text{for adsorption on a silver substrate,} \\ \frac{1}{2} (g_{\text{Ag}_2\text{O-bulk}} - \mu_{\text{O}}), & \text{when bulk Ag}_2\text{O is present.} \end{cases}$$

(3)

We use small symbols ($g_{\text{Ag-bulk}}$, $g_{\text{Ag}_2\text{O-bulk}}$) in Eq. (3) to indicate that these are free energies per Ag atom (top) and per Ag$_2$O unit (bottom). These two quantities are calculated by DFT-GGA. In Eq. (1), we have furthermore

$$G = \begin{cases} G_{\text{slab}}, & \text{to obtain surface energies,} \\ G_{\text{O/Ag}(111)} - G_{\text{Ag}(111)}^{\text{slab}}, & \text{to obtain adsorption energies on Ag(111).} \end{cases}$$

(4)

These quantities are also calculated by DFT-GGA. For the temperature dependence of $\mu_{\text{O}_2}(T, p^0)$, we use thermodynamic tables\textsuperscript{47} (i.e., experimental data), while the $T = 0$ K value of Eq. (2) is

$$\mu_{\text{O}}(T = 0K, p) = \frac{1}{2} E_{\text{O}_2}^{\text{total}} - E_{\text{Ag}_2\text{O-bulk}} - 2E_{\text{Ag-bulk}} + H_{\text{Ag}_2\text{O-bulk}}' .$$

(5)
In the above equation we use the experimental value of the heat of formation $H^{total}_{\text{Ag}_2\text{O-bulk}}$ because of its very small value [0.323 eV at the standard state (room temperature and atmospheric pressure)]\textsuperscript{48} or 0.325 eV extrapolated back to $T = 0$ K, which implies that the error introduced in the later discussion by using the theoretical value (0.092 eV) would be significant. We note that the error in $H^{total}_{\text{Ag}_2\text{O-bulk}}$ is due to the significant overbinding of O$_2$, and the underbinding obtained for bulk Ag and bulk Ag$_2$O.

We choose the zero reference state of $\mu_O(T,p)$ to be the total energy of oxygen in an isolated molecule, i.e., $\mu_O(0 \text{ K}, p) = \frac{1}{2}E^{total}_{O_2} = 0$. This will be called the “oxygen-rich” condition. The so-called “oxygen-poor” value of $\mu_O$ is taken as

$$\mu_O^{poor} = E^{total}_{\text{Ag}_2\text{O-bulk}} - 2E^{total}_{\text{Ag-bulk}}. \tag{6}$$

If the oxygen chemical potential becomes lower (more negative) than $\mu_O^{poor}$, di-silver oxide starts to decompose. For oxides of different transition metals, the “oxygen poor” limit can vary significantly depending on the heat of formation. For example, what may be rather “oxygen rich” for RuO$_2$ (with $H_f = 1.60$ eV) is already too “oxygen poor” for Ag$_2$O to be stable. The typical range of the allowed oxygen chemical potential is determined and limited by the pressures and temperatures used in industry and laboratories, namely, from several to thousand K, and from ultrahigh vacuum ($< 10^{-12}$ atm) to several hundreds of atmospheres.

For our investigations into various terminations of Ag$_2$O(111) surfaces, we will also consider the energy required to remove an oxygen atom from the surface and bring it to the vacuum. We call this the “removal energy,” and it is defined as

$$E^{removal} = E_{O/\text{Ag}_2\text{O}(111)} + (E_{\text{Ag}_2\text{O}(111)}^{ref} + E_O), \tag{7}$$

where $E_{O/\text{Ag}_2\text{O}(111)}$ is the total energy of the surface termination of interest, containing the O atom to be removed and $E_{\text{Ag}_2\text{O}(111)}^{ref}$ is that of the reference system, i.e., the surface structure remaining after the O atom has been removed. $E_O$ is the total energy of a free O atom where the spin-polarization energy ($-1.60$ eV) is taken into account. The removal energy reflects the bond strength of the removed O atom to the surface. It could alternatively be referred to (half) the energy of the free oxygen molecule, where the magnitude would just be reduced by half the O$_2$ binding energy.

To help analyze the electronic structure, we also calculate the density of states $N(\varepsilon)$ as well as the state-resolved DOS, or projected DOS $N_{\sigma}(\varepsilon)$. The definitions are given in Ref. 49.

III. BULK Ag$_2$O

The most stable silver oxide is Ag$_2$O, which decomposes at 460 K under atmospheric pressure.\textsuperscript{48} The crystallographic structure of di-silver oxide Ag$_2$O is shown in Fig. 1(a). It has the cuprite structure with six atoms per cubic unit cell, (two oxygen atoms and four silver atoms) where the oxygen atoms form a body centered cubic lattice, while the metal atoms are located on the vertices of a tetrahedron around each oxygen atom, forming a face-centered-cubic lattice.\textsuperscript{51} A characteristic feature, as seen in Fig. 1(a), is a linear oxygen-metal-oxygen coordination. The calculations for Ag$_2$O are performed with 35 special $k$ points in the irreducible part of the Brillouin zone (IBZ) of the six atom cell, corresponding to 1000 points in the full BZ. We used the same pseudopotentials and energy cutoff (50 Ry) as in our earlier works.\textsuperscript{20,21}

The calculated results, as well as available experimental data, are presented in Table I. The theoretical lattice constant of 4.91 Å is 4% larger than the experimental one which is 4.74 Å (Ref. 51) or 4.72 Å. The nearest-neighbor distance
between Ag atoms in Ag₂O is calculated to be 3.47 Å, which is larger than the (theoretical) value in bulk fcc silver which is 2.97 Å. The calculated O-Ag bondlength is slightly shorter than for oxygen chemisorbed on the Ag(111) surface (compare 2.13 Å to 2.15–2.18 Å, depending on the coverage), and similar to the O-Ag bondlengths of the \((4 \times 4)\) structure [depicted in Fig. 3 (bottom), discussed later and in Ref. 21] which are 2.10–2.11 Å for the upper O-Ag bonds and 2.14–2.16 Å for the lower O-Ag bonds (involving the surface Ag atoms). It is also similar to the O-Ag bond lengths of the favorable on-surface+subsurface commensurate oxide structures \((2.10–2.15)\) Å reported in Ref. 21 [see e.g., the upper O-Ag-O layer of Fig. 3 (top)].

The calculated heat of formation, obtained as the total-energy difference between di-silver oxide \((\text{Ag}_2\text{O})\) and the sum of the total energies of bulk silver (two Ag atoms) and the half the oxygen molecule \((1/2 \text{O}_2)\) is 0.092 eV. The difference to the experimental value of 0.231 eV is due to the significant overbinding of \(\text{O}_2\), which is 0.566 eV per oxygen atom, and the underbinding obtained for bulk Ag and bulk Ag₂O. This leads to some cancellation of errors but it is not complete. In Table I, the calculated bulk modulus refers to the theoretical lattice constant. To the best of our knowledge, there are no theoretical values of the heat of formation with which to compare in the literature, nor lattice constants or bulk moduli. Both experiment and theory, nevertheless, show that formation of di-silver oxide is an exothermic process, and the small value is in-line with the low temperature of decomposition noted above. The heat of formation of di-silver oxide is significantly smaller compared to the transition metal oxides for metals to the left of Ag in the periodic table, e.g., 1.60, 1.19, and 0.97 eV per oxygen atom (experimental data at \(T\to0\) K and atmospheric pressure) for RuO₂, Rh₂O₃, and PdO, respectively.⁴⁸

The total density of states is shown in the right panel of Fig. 1(b). The peak located in the energy range \(-4.0\) to \(-2.0\) eV with respect to Fermi energy arises mainly from the Ag-4d states, where there is only a small overlap with the O-2p band, which has main (but smaller) peaks in the regions of \(-6\) to \(-4\) eV and \(-2\) to 0 eV. The states in the former region are largely bonding and those in the latter are mainly antibonding in character, as determined by inspecting the spatial distribution of the wave-functions in these regions. The occupation of the antibonding states, and small overlap of O and Ag states, explains the low stability of the oxide.

The band structure of Ag₂O is shown in the left panel of Fig. 1(b). It can be seen that at the \(\Gamma\) point, the valence and conduction bands overlap, making Ag₂O metallic, instead of semiconducting with a band gap of 1.3 eV (Ref. 52) as found in experiments. The spatial distribution of the wave functions in the conduction band close to the Fermi energy (investigated for several \(k\) points), shows that the states are of anti-bonding nature, involving a hybridization of Ag-4d–O-2p– and Ag-4d–O-3s-like orbitals. The occupation around \(\Gamma\) of these bands could contribute to the underestimated of the heat of formation of Ag₂O noted above. We note, however, that this down shift of the conduction band has only a small effect on the DOS [see right panel of Fig. 1(b) at energy \(\epsilon =0\)]. We also performed calculations using the slightly smaller experimental lattice constant; this also did not yield a band gap. Using the augmented-spherical wave (ASW) method with an extended basis set and the atomic sphere approximation (ASA) within the local-density approximation (LDA), Czyzyk et al.⁵³ calculated the projected density of states and band structure of Ag₂O, where a band gap 0.40 eV was found. However, no band gap was obtained by Deb and Chatterjee⁵⁴ using the full-potential linearized augmented plane wave (FP-LAPW) method within the local-spin density approximation (LSDA). Their reported density of states and band structure are very similar to those of our present work. In both of the above mentioned works, the lattice constant was not stated, but we believe the experimental one was used because our results calculated at the experimental lattice constant agree more closely. On increasing the energy cutoff and using harder pseudopotentials, namely, \(r_s=r_d\) = 2.20 Bohr and \(r_p=r_d\) = 2.10 Bohr for silver and \(r_s=r_p=r_d\) = 1.2 Bohr for oxygen, and an energy cutoff of 90 Ry, no significant change in the results was found (less than 0.02 eV for the heat of formation, 0.01 Å for the lattice constant, and 0.001 Mbar for the bulk modulus). Thus it appears that Ag₂O is another semiconductor for which the LDA and GGA yield no band gap. Other materials for which this is the case are, for example, Ge,⁵⁵ InN,⁵⁶ and ScN.⁵⁷

Calculations for Ag₂O performed using the screened-exchange local-density approximation,⁵⁸ which affords a more accurate description of the band gap compared to the LDA (or GGA), yield a band gap of 0.73 eV which shows that this material is indeed a direct narrow band-gap semiconductor, in agreement with experiment.

### IV. Ag₂O(111) TERMINATIONS

In this section, we examine various surface terminations of Ag₂O(111). This is interesting for two reasons, first in relation to learning more about oxide surfaces in general through comparison with existing studies, and secondly since our earlier works²⁰,²¹ for oxygen adsorption at the Ag(111) surface predicted that for high oxygen concentrations, thick oxidelike structures form which contain half a monolayer of O between the Ag layers and quarter of a monolayer on the surface (see, e.g., Fig. 3, upper right). On inspection of the geometry, we observe that the structures correspond exactly to that of the Ag₂O(111) surface, but where the positions of the Ag atoms are commensurate with those of the underlying Ag(111) substrate, so the structure is laterally compressed compared to the true oxide [compare Figs. 2 and 3 (upper)]. With respect to different terminations, additional oxygen could adsorb on top of the Ag atoms. Such sites are called coordinatively unsaturated sites (CUS’s), since compared to in the bulk, they are lacking their full O coordination; for the present system, the CUS Ag atoms are missing one O bond. Conversely, it could be favorable that the surface has a metal-atom termination instead. We have therefore studied several different terminations, and calculated the surface free energy as a function of the oxygen chemical potential [see, Eq. (2)] to determine which have the lowest energy.
Atoms the site would be a CUS site! Investigated the surfaces with oxygen in the hollow site, and O atoms and four Ag atoms per unit cell of the layer. We stoichiometric surface can be thought of as a stacking of surface is that where both of these O atoms are missing. The geometry of the Ag\,2\,O\,(111) surface. The silver atoms are represented, as is the “O-Ag-O trilayer” which contains two O atoms and four Ag atoms per surface unit cell.

Figure 2 shows the two surface O atoms: The hollow site corresponds to the bulk stacking and the O atom in the top site makes this termination oxygen rich (without the O\(_{\text{top}}\) atoms the site would be a CUS site). The silver-terminated surface is that where both of these O atoms are missing. The stoichiometric surface can be thought of as a stacking of O-Ag-O trilayers as indicated in Fig. 2, which contains two O atoms and four Ag atoms per surface unit cell of the layer. We investigated the surfaces with oxygen in the hollow site, and

![Image](http://example.com/image1.png)

**FIG. 2.** Top (upper) and perspective (lower) views of the atomic geometry of the Ag\(_2\)O\,(111) surface. The silver atoms are represented by large, pale spheres and the oxygen atoms by small dark spheres. The hollow and top sites of surface oxygen atoms are indicated, as is the “O-Ag-O trilayer” which contains two O atoms and four Ag atoms per surface unit cell.

The silver-terminated surface below the “O-Ag-O” trilayer are represented as the white circles. Note that this surface structure is the same as a layer of Ag\(_2\)O\,(111), except there is one Ag atom missing in the indicated cell, which would sit directly on top of the Ag atom in the (111) surface below (which is energetically unfavorable).

![Image](http://example.com/image2.png)

**FIG. 3.** Illustration of the atomic geometry of an oxidelike structure commensurate with the Ag(111) surface. Top view (upper left) and perspective view (upper right) for an O concentration of 1.25 ML. Ag atoms and O atoms are represented by large pale and small dark spheres, respectively. The structure is very similar to the true oxide (Fig. 2), but laterally compressed in comparison. Lower figure: sketch of the atomic structure proposed in the STM study of Ref. 24 for the (4 \times 4) oxygen phase on Ag\,(111). The surface unit cell is indicated. The oxygen atoms are represented by small dark circles, the uppermost Ag atoms by gray circles, and the intact plane of Ag\,(111) atoms lying below the “O-Ag-O” trilayer are represented as the white circles. Note that this surface structure is the same as a layer of Ag\(_2\)O\,(111), except there is one Ag atom missing in the indicated cell, which would sit directly on top of the Ag atom in the (111) surface below (which is energetically unfavorable).

![Image](http://example.com/image3.png)

**FIG. 4.** Surface free energies \(\gamma\) as a function of the O chemical potential, for the various surface terminations of Ag\(_2\)O\,(111): namely, Ag-terminated (dashed line), oxygen at the hollow site which corresponds to the stoichiometric surface (full line), and oxygen at both the hollow and top sites (dot-dashed line). The left and right vertical lines correspond to the “oxygen-poor-limit” and the “oxygen-rich-limit,” as explained in the text, and \(\Delta \mu_O\) is defined as \(\mu_O - 1/2E_{\text{O}_2}^\text{total}\), using Eq. (5) for the latter.

The results described above are in contrast to those found for other (polar) oxide surfaces, e.g., Ru\(_3\)O\(_5\)(110) (Ref. 39) and \(\alpha-\text{Fe}_3\text{O}_4(0001)\)\(^{,38}\) where “full oxygen occupation” of available sites becomes favorable for high oxygen chemical potentials. The reason for this difference is because here the interaction between oxygen and silver is so weak that even extremely high values of the oxygen chemical potential cannot stabilize additional oxygen at the polar surface.

In Table II various physical properties of the surfaces are listed. In particular, it can be seen that the removal energy of O hollow is greatest when there is no additional oxygen at the neighboring top site (compare 3.97 with 3.72 eV). Thus,
the presence of O in the top site weakens the bond of the O-hollow atom to the surface, showing that there is a repulsive interaction between the O atoms in the hollow and top sites. We note that the value of 3.97 eV is significantly stronger than 3.52 eV as obtained for O bonded to the clean Ag(111) surface in the fcc site at the same on-surface coverage of 0.25 ML as the O-hollow atoms shown in Fig. 2. It is, however, similar to the removal energies we found in our earlier work for the energetically favorable structures involving both on-surface and subsurface oxygen and the compressed oxide-like structures, where the values are in the range 3.90–4.01 eV. These bond strengths can no longer be regarded as weak, but are intermediate and of similar order as those identified for the reactive species involved in certain chemical reactions.

The bondlength between oxygen in the hollow site and the surface silver atom for the energetically favorable Ag₂O(111) structure is 2.11 Å, which is slightly shorter than the corresponding bulk value (2.13 Å) noted in Sec. III. It is similar to that of the energetically favorable compressed oxide-like structures (2.10–2.15 Å) [see Fig. 3 (top)] which have a very similar local geometry, but smaller than that for chemisorbed on-surface oxygen at Ag(111) (2.15–2.18 Å) found in our previous works. The bondlength between oxygen in the top site and the silver atom below 1.96 Å is significantly shorter because of its low coordination. Structurally, except for the Ag-terminated surface, where there is a large contraction of 10.3% of the first metal interlayer spacing compared to that of the bulk truncated surface (calculated using the center of the layers), the relative change in the metal interlayer spacings of the oxygen-terminated structures is modest, i.e., less than 2.0%. For the stoichiometric surface, the rumpling of the silver atoms in surface metal layer is not negligible; the magnitude is of the order 0.10 Å. This is due to an outward movement of the Ag atom in the surface unit cell that is not bonded to the surface O atoms, i.e., the CUS atom. These relaxations are smaller than those of the “true transition-metal” oxides (i.e., with incomplete filling of the d band), e.g., α-Fe₂O₃(0001) and RuO₂(110), which we attribute to the weaker bonding between silver and oxygen.

Similarly, for adsorption of oxygen on the Ag(111) surface, we found only very small relaxations of the substrate metal layers in contrast to O on Ru(0001) (Ref. 60) where the O-metal bond is notably stronger and oxygen adsorption induces a significant expansion of the first metal interlayer spacing. The nature of the bonding of the di-silver oxide surface is similar to that of the bulk oxide: Oxygen and silver hybridize mainly via the Ag-4p orbitals as shown in Fig. 5, where the atom projected density of states (PDOS) in Fig. 5 is similar to that of the clean Ag(111) surface (see Fig. 5 in Ref. 20), the Ag-4d band is significantly narrower, which is due to the increased distance between the silver atoms and the weak hybridization with O-2p states. The PDOS in Fig. 5 is rather similar to that of the oxide-like structures with some bond localization as presented in Fig. 12 of Ref. 20. In Fig. 5 two features, one at −5.53 eV and one at −0.66 eV, with respect to the Fermi energy, can be seen which are well separated from the center of the d band. This is in contrast to the case of chemisorption on the metal substrate, where there is only one peak outside of the Ag-4d band, for example at −1.35 eV for oxygen in the on-surface fcc-hollow site at coverage 0.25 ML.

V. TRANSITION TO THE OXIDE

Having studied the atomic and electronic structure of bulk Ag₂O and determined the thermodynamically most favorable surface termination of the (111) crystallographic plane, we now wish to investigate under what conditions a transition from the oxide-like structures (as reported in Ref. 21), with positions of Ag atoms commensurate with those of Ag(111), to the real oxide structure (which is laterally expanded by 14% in comparison) might occur. Such a commensurate oxide-like structure is depicted in Fig. 3 (top) for an O concentration of 1.25 ML. In order to investigate such a transition, we study the difference of the total energy (per O atom)
between oxidelike films (with commensurate Ag positions to those of the metal substrate) and corresponding “real” (unstrained) oxide layers as a function of thickness. Because the structures can be thought of as the stacking of “O-Ag-O trilayers” where there are two O atoms and four Ag atoms per \((2 \times 2)\) surface unit cell of the trilayer, i.e., stoichiometry \(\text{Ag}_2\text{O}\) (see Fig. 2), we calculate the energy difference with increasing numbers of trilayers. Then the number of layers at which the true oxide structure becomes energetically more favorable, defines a critical thickness for the transition to occur. In this approach, the coupling between the (commensurate) oxidelike film with the underlying metal substrate is not included. We note that this study may be somewhat “academic” since there is so far no evidence for such compressed oxide structures, however, this structure is a plausible intermediate and perhaps can even be stabilized under certain experimental conditions. Furthermore, additional insight into the \(\text{O}/\text{Ag}\) system can be obtained by these calculations, and moreover, when compared to analogous studies performed for other \(\text{O}/\text{transition-metal systems, in particular for O}/\text{Ru(0001)},^61\) various trends may be identified.

The results are shown in Fig. 6. A negative value indicates that the true (unstrained) oxide structures are energetically more favorable. We note that Fig. 6 assumes that the number of O atoms is fixed. However, in many (most) experiments that aim to study very high coverages, the actual thermodynamic variable may be \(\mu_O\).

For one O-Ag-O trilayer, the “strained” [commensurate with \(\text{Ag}(111)\)] oxide film is significantly more favorable than the corresponding true oxide, as indicated by the relatively large positive value of 0.53 eV, but the preference decreases quickly to 0.13 eV at three trilayers. On increasing the thickness further, the difference in the energy between the two structures decreases more slowly, and when the number of trilayers is nine, the true oxide film is energetically more favorable, as indicated by the negative value in Fig. 6. Because each trilayer contributes half a monolayer of oxygen, the critical coverage for the transition from the commensurate oxidelike film to the true oxide film is around 3.5–4.5 ML. As the number of the trilayers approaches infinity, clearly the curve approaches the value corresponding to the energy difference (per O atom) between the bulk oxide and the “strained” [commensurate with \(\text{Ag}(111)\)] bulk material.

FIG. 6. Difference in total energy (per O atom) between a stack of O-Ag-O trilayers, commensurate with Ag(111), and that of a corresponding oxide layer, as a function of the number of trilayers in the stack (denoted by “+”). The commensurate structures are just like the oxide, only laterally compressed by 14% and vertically expanded by ~21%. A negative value indicates that the true oxide structure is energetically more favorable. For comparison, analogous results (from Ref. 61) are shown for Ru(0001) (denoted by crosses) namely, the difference in total energy (per O atom) between a stack of O-Ru-O trilayers [commensurate with Ru(0001)] and that of corresponding RuO\(_2\)(110) layers as a function of the number of layers in the stack. The lines connecting the Ag and Ru results (full and dashed lines) are guides to the eye. Insets illustrate the atomic geometry of the bulk oxide structures where oxygen and metal atoms are represented by small dark and large pale circles, respectively. For RuO\(_2\), the O-Ru-O trilayer structure is also depicted (lower left), as well as its conversion, via an “accordionlike” expansion, into a “true” RuO\(_2\)(110) layer.

The electronic properties of the “strained” [commensurate with \(\text{Ag}(111)\)] oxide film are very similar to those of the true oxide as can be seen from comparison of Fig. 12 in Ref. 21 with Fig. 5. This at first may be somewhat surprising since the lateral dimensions of the commensurate, oxidelike structures are smaller, by 14%. We find, however, that much of the associated strain energy is relieved by expansion of the Ag layers perpendicular to the surface to accommodate the O atoms in between. Through this relaxation, the lattice volume per atom is 92% of that of the bulk oxide.

In Fig. 6, analogous results for the phase transition from O-Ru-O trilayer structures [commensurate with Ru(0001)] to the corresponding real RuO\(_2\)(110) (Ref. 61) oxide layers are also presented for comparison. Here the “O-Ru-O trilayer” contains two O atoms and one Ru atom per \((1 \times 1)\) surface unit cell, i.e., stoichiometry RuO\(_2\). Bulk ruthenium dioxide has the rutile structure. The atomic geometry of RuO\(_2\)(110) is shown as an inset in Fig. 6, as is that of the (commensurate) trilayer and its expansion into a corresponding true RuO\(_2\)(110) layer (bottom, left). The commensurate trilayer structures are, laterally, 15 and 35% smaller than for RuO\(_2\)(110) [i.e., in the \(\hat{a}\) and \(\hat{b}\) directions of the rectangular (110) surface unit cell]. In this case the transformation from commensurate trilayer to corresponding oxide layer, involves not only a lateral expansion similar to the di-silver oxide structures, but also a “rotation” of the O atoms about the Ru atoms, as seen from the inset (explained in more detail in Ref. 61). In particular, every second O-Ru-O component rotates to be horizontal to the surface, while the alternative components rotate so that they are perpendicular to the surface. In contrast to the Ag-O system, the phase transition into RuO\(_2\)(110) oxide happens after only one O-Ru-O trilayer.
Also, for Ru(0001), each trilayer contributes two monolayers of oxygen, so the critical coverage is around 2–4 ML. Because of the considerable energy gain due to formation of the real RuO₂ oxide layers, compared to continuous of stacking of the O-Ru-O trilayers, the formation of the oxide structure will certainly take place once it is kinetically possible.

It is interesting to note that the energy differences between oxygen in the trilayers and the corresponding bulk metal oxide layers, exhibits a rather similar dependence on the number of layers for both O/Ag and O/Ru, except shifted in energy. This makes sense since we consider the energy per O atom, and as the stack becomes larger, there are more and more O atoms in the center of the stack which have atomic environments which are similar (and thus similar energies), i.e., the affect on the energy due to the surfaces of the stack, diminishes. In view of this similarity, together with the fact that the heat of formation of e.g., the Rh (1.19 eV) and Pd (0.97 eV) oxides lie between the extremes of Ru (1.60 eV) and Ag (0.33 eV) oxides, we may expect that investigations of analogous transitions from commensurate structures to the bulk oxide phase for Rh and Pd would exhibit a behavior intermediate to those of Ru and Ag. This would imply the trend that the transition from intermediate, commensurate, two-dimensional oxide films to the true oxide phase occurs sooner (i.e., for thinner films) for the elements more to the left in the periodic table, such as Ru, with larger heats of formation, compared to those towards the right, which have smaller heats of formation like Ag.

From a recent trend study on the oxidation at the basal plane of late 4d transition metals (from middle to right in the Periodic Table, i.e., Ru, Rh, Pd, Ag), it was found that oxygen starts to enter the subsurface region only after a certain critical coverage has been reached on the surface. Namely, at coverages beyond which the heat of oxygen adsorption on the surface becomes unfavorable compared to the heat of formation of the bulk oxide. The reported critical coverages are about 1.0, 1.0, and 0.75, and 0.25 ML for Ru, Rh, Pd, and Ag, respectively. Beyond these critical coverages, onset of bulk oxide formation was proposed to take place. Clearly, however, due to kinetic hindering, bulk oxide formation may not be able to occur “immediately” and then it may proceed via the formation of oxidelike layers as described above.

VI. THERMODYNAMICS OF THE OXYGEN-SILVER(111) SYSTEM

We now turn to investigate the affect of the temperature and pressure on the stability of the various oxygen/silver structures, the energetics and atomic geometries of which were reported in our previous publications. In particular, the energies are summarized in Fig. 14 of Ref. 21. To incorporate the effect of temperature and pressure, we calculate the Gibbs free energy as a function of the oxygen chemical potential [see, Eqs. (1) and (2)] similarly to the oxide surfaces, except here the reference system is the clean metal substrate. As discussed above, the “oxygen poor” limit is that below which di-silver oxide is unstable. For oxygen adsorption at the Ag(111) surface, however, this limitation doesn’t necessarily exist. The results are shown in Fig. 7, where we only show the low energy structures. The dependence of the energy on \( \mu _{O} \), namely, the slope of the line, reflects the O coverage. The higher the coverage, the steeper the gradient. We also include the corresponding temperatures for two given pressures, namely, that which corresponds to UHV and the other to the atmospheric conditions of catalysis.

It can be seen that for low values of the oxygen chemical potential, i.e., \( \mu _{O} < -0.78 \) eV, the clean Ag(111) surface is thermodynamically most stable. For higher values, (i.e., in the range of \( -0.78 \) to \( -0.63 \) eV), a low coverage (1/16 ML) of oxygen on the surface in fcc-hollow sites is thermodynamically most stable, and for values greater than \( -0.63 \) eV and less than \( -0.54 \) eV, slightly higher coverages of adsorbed O (1/9 and 1/4 ML) become most stable. Increasing the oxygen chemical potential to values greater than \( -0.54 \) eV leads to the formation of the reconstructed (4×4) phase (labeled “0.375 ML”), which is stable up to \( \Delta \mu _{O} = -0.325 \) eV, i.e., to the “O-poor” limit (left vertical dashed line), where the bulk oxide becomes stable. The atomic geometry of the (4×4) phase is illustrated in Fig. 3 (bottom). The atomic structure is very similar to a trilayer of Ag₂O(111), except with one Ag in the (4×4) cell missing. The basis for this proposed structure (see Refs. 24,25,63–65) is the observation that the area of the \( (\sqrt{3} \times \sqrt{3})R30^\circ \) Ag₂O(111) surface unit cell is practically four times that of clean (1×1)-Ag(111); the difference in the

![Figure 7](image-url)
FIG. 8. Calculated phase diagram for the oxygen-Ag(111) system. On the horizontal axis is the temperature in K and on the vertical axis is the pressure.

length of the lattice vectors is only 1.2% (0.3% for the experimental lattice constants). Although the mentioned surface unit cells are (practically) commensurate, the positions of the Ag atoms within the \((\sqrt{3} \times \sqrt{3})R30^\circ\) oxide surface cell are, all except one, incommensurate with those of the Ag(111) surface.

It can be seen that for values of \(\Delta \mu_O\) greater than \(-0.54\) eV, the commensurate oxidelike structures (see, e.g., Fig. 3, top) are also quite favorable (labeled by the O coverages 0.75, 1.25, and 2.25 ML). It can be noticed that the thicker the film, the more stable the structure is (i.e., the steeper the curve). Clearly, in the limit of an infinitely thick film, the corresponding line would be very close to the left vertical one. This is so because the average O binding energy in the thick commensurate oxidelike structures is very similar to that of O in the bulk oxide as described above. From Fig. 7 it can be concluded that the bulk oxide is the thermodynamically most favorable structure once the oxygen chemical potential is higher than the “O-poor” condition, which is expected. However, the kinetics of oxide formation can be very slow and the thickness of the oxide film will be limited by O and metal bulk diffusion. Thus, it is unlikely that full thermal equilibrium of O\(_2\) with bulk di-silver oxide will be achieved.

A. The O-Ag(111) phase diagram

The results discussed above can be displayed as a continuous function of temperature and pressure by constructing a two-dimensional phase diagram, if we only consider the lowest energy structures. This has been done in Ref. 22 and the results are shown again in Fig. 8. Before giving a discussion, it is necessary to consider the magnitude of the error bars in the temperature and pressure. An indication can be obtained from knowledge of the stability of the bulk oxide, i.e., that it decomposes at 460 K under atmospheric pressure. From Figs. 7 and 8, the calculations predict that the bulk oxide decomposes at \(\sim 350\) K at atmospheric pressure, i.e., around 110 K lower than experiment. At 460 K, the bulk oxide is unstable when the pressure is lower than \(10^3\) atm., i.e., compared to 1 atm. as observed experimentally. The reason for this disparity may be due to systematic errors of the DFT-GGA approach\(^{66}\) and/or to neglect of the entropy contribution. In particular, we note that the experimental \(-TS\) contribution for bulk silver at 450 K and 1 atm. is \(-0.48\) eV per two silver atoms, and \(-0.70\) for Ag\(_2\)O per formula unit.\(^{66}\) Thus, the entropy contribution from di-silver oxide, compared to silver bulk (the entropy of O\(_2\) has already been included in the chemical potential), is greater by 0.22 eV per two silver atoms, which will shift the left vertical line in Fig. 7 further to the left. Therefore, in Figs. 7 and 8, we generally expect that the temperature is underestimated, but the error less than 110 K, while correspondingly, the pressure is overestimated, by less than three orders of magnitude. For the temperature values quoted below, we give an estimate of the upper limit in brackets, that is, 110 K higher than that obtained from Fig. 8. In spite of these uncertainties, we believe that the relative stability of the various systems and our general understanding and conclusions will not be affected.

Typically, vibrational contributions to differences in the Gibbs free energies of extended systems exhibit some cancellation.\(^{39}\) However, when there are additional atomic or molecular species which are not present in the reference system, the situation may be different as there will be no effective cancellation of the contributions of such species between surface and reference systems [as is the case using the clean Ag(111) as reference]. We investigate this aspect in more detail for the present system; in particular, we consider low coverages of on-surface oxygen and thin oxidelike structures. We calculate the difference in the vibrational and entropic contributions to the Gibbs free energy \(F_{\text{vib}} = F_{\text{vib}} - TS_{\text{vib}}\) as

\[
\Delta F_{\text{vib}}(T, \omega) = \left\{ \begin{align*}
\hbar \omega \left( 1 \frac{1}{2} + \frac{1}{e^{\beta \hbar \omega} - 1} \right) \\
-k_B T \left[ \beta \hbar \omega e^{\beta \hbar \omega} - 1 \right] - \ln(1 - e^{\beta \hbar \omega}) \right\},
\]

where \(\beta = 1/k_B T\) (see, e.g., Ref. 35, 39). The vibrations which enter Eq. (8) are those of the adsorbed O atoms. The calculated perpendicular vibrational frequency for 0.25 ML of oxygen in fcc sites on the surface is 50 meV (400 cm\(^{-1}\)), and we assume that the lateral vibrations are the same (this is a somewhat rough approximation but suffices for the present discussion, see below). Thus, in calculating \(\Delta F_{\text{vib}}(T, \omega)\) we multiply by a factor of 3 corresponding to these three modes. For the thin commensurate oxidelike structure containing one O atom on the surface in fcc sites and one subsurface O in tetrahedral sites (i.e., a total coverage of 0.5 ML), the calculated perpendicular vibrational frequencies are 56 meV (455 cm\(^{-1}\)) and 65 meV (526 cm\(^{-1}\)), respectively, for these two species. Again, assuming each O atom has three modes equal to the perpendicular ones, we obtain the results presented in Fig. 9.
It can be seen from Fig. 9 that for on-surface O, for temperatures up to about 800 K, the contribution is a maximum of about \( \pm 2 \text{ meV/Å}^2 \). For lower coverages of on-surface O such as the 1/16 and 1/9 ML structures, the contribution will be smaller (assuming the frequency is similar). For the oxidelike structure at coverage 0.5 ML, the effect is larger, namely, 5–6 meV/Å\(^2\) for temperatures up to about 400 K, thereafter decreasing. For the \((4 \times 4)\) thin oxide structure with coverage 0.375 ML, we expect the vibrational frequencies of the O species will be similar to those of the structure at 0.5 ML since the local coordination of the O atoms is similar, as well as the electronic structure. In addition, the vibrational frequency of O in bulk \( \text{Ag}_2\text{O} \) is reported to be 66 meV (530 cm\(^{-1}\)).\(^{67}\) which is very close to that of the O atom in the subsurface tetrahedral site of the 0.5 ML coverage structure (65 meV). Because of the lower O coverage of the \((4 \times 4)\) structure, the contribution to the free energy will be smaller than the 0.5 ML structure by about 0.375/0.5 = 0.75, giving a maximum contribution at low temperatures of 4.5 meV/Å\(^2\) (dot-dashed line in Fig. 9).

For the \((4 \times 4)\) structure, in the temperature range 400 to 730 K, it gives rise to a destabilization of less than 3.5 meV/Å\(^2\) (the value at 400 K, which decreases with temperature), which on inspecting the energy scale of Fig. 7 can be seen to be non-negligible. The relative energy difference, however between the on-surface species and \((4 \times 4)\) structure, which determines the sequence of the stable phases, can be seen from Fig. 9 to be affected by less than 1.9 meV/Å\(^2\) (1.9 = 3.5 - 1.6 meV/Å\(^2\)) for temperatures greater than 400 K, which will therefore have a small effect. We note that if the lateral vibrations would be 30% greater than the perpendicular ones for the on-surface and oxidelike structures considered, this would result only in a small increase in the contribution to the free energy, i.e., of 0.5 and 1.2 meV/Å\(^2\), respectively, at \( T \) close to zero.

The above analysis may be regarded as qualitative, but it gives an estimate of the magnitude of the changes in the free energy due to consideration of vibrational and entropic effects of the extended systems for the energetically most important O/Ag structures. Clearly, for other systems with adsorbed species that have high frequencies such as OH (of the order 400 meV),\(^{68}\) the contribution to the free energy from vibrational and entropic effects will be significantly greater and will have to be taken into account in determination of the phase diagram.

Summarizing the results contained in Fig. 8, we obtain the following predictions: Under UHV conditions (\( p = 10^{-12} \text{ atm} \)), chemisorption of oxygen on Ag(111) with low coverage (less than 0.0625 ML) is stable up to about 360 K (including corrections: 470 K). This is in-line with experimental results\(^{64}\) where it has been found that for temperatures less than 490 K, 0.03 ML of oxygen can be adsorbed on the Ag(111) surface. Under UHV conditions, the calculations also predict the stability of the \((4 \times 4)\) phase at the lower temperatures of 160 K (including corrections: 270 K) to 260 K (including corrections: 370 K). This is also consistent with scanning tunneling microscopy (STM) results obtained under UHV conditions at low (4 K) temperatures\(^{24}\) where atomic resolution images of the \((4 \times 4)\) phase have been achieved. The \((4 \times 4)\) phase, however, cannot be directly prepared under these conditions as it is known from experiment that temperatures greater than 400 K are required in order for the necessary atomic rearrangement take place. It is therefore clear that it is the kinetics that prevent the reconstruction and formation of this phase at low pressures and temperatures. From Fig. 8, it can be seen that the \((4 \times 4)\) phase should develop into the oxide if the oxygen pressure is raised or if the temperature is lower than 160 K (including corrections: 270 K). Such transitions, however, may be prevented or hindered by kinetics (e.g., diffusion barriers) since extensive mass transport is required to form the oxide.

At atmospheric pressure (\( p = 1 \text{ atm} \)), typical of that used in industry, the bulk oxide is calculated to be only stable up to around 350 K (including corrections: 460 K). From this point of view, it can be concluded that for industrial reactions which take place at considerably higher temperatures, di-silver oxide cannot be the active phase. The \((4 \times 4)\) surface-oxide phase, however, is stable in the temperature range of 350–530 K (including corrections: 460–640 K) which coincides with that at which epoxidation of ethylene occurs. We therefore propose that such atomic O species are actuating the reaction. With regard to the catalytic reactions that take place at even higher temperatures, it can be seen that only very low coverages of chemisorbed O are stable up to 720 K (including corrections: 830 K); O atoms adsorbed at undercoordinated surface Ag atoms are up to 865 K (including corrections: 975 K) (see below). Therefore these species could possibly play a role in these reactions.

It is interesting to note that regardless of the pressure, Fig. 8 shows that the ordering of the stable phases is the same, so that the same phases should be observable at different pressures if the temperature is adjusted accordingly. This indicates a way to bridge the pressure gap, i.e., by appropriate choice of the pressure and temperatures to stay within the same phase. However, it is clear that sometimes this will not be possible within reasonable times because of kinetic hindering for dissociation and sticking of \( \text{O}_2 \) (i.e., delivery of
sufficient concentrations of atomic oxygen to the surface), as well as to kinetic limitations for adequate atomic rearrangement.

Experimentally, it has been reported that a "strongly bound" oxygen species (named O$_2^*$) forms after high temperature ($\sim$900 K) and atmospheric pressure exposure of polycrystalline silver, as well as Ag(111), to an oxygen environment.\cite{12,17} Temperature programmed desorption spectra show that O$_2^*$ desorbs at around 900 K and it is proposed by the authors to be the active species for the dehydrogenation of methanol\cite{12,13} where it diffuses to the surface from bulk sites. At the high temperature of $\sim$900 K in an oxygen atmosphere, silver reconstructs to form many (111) facets\cite{12,13} at the surface and it is speculated that this means that oxygen diffusion takes place by a so-called "subsidiary diffusion mechanism" which the authors argue has higher barriers than usual interstitial diffusion, and to give rise to the O$_2$ desorption peak at the high temperature of 900 K. From our calculations, this picture is difficult to understand, in particular, how sufficient amounts of bulk-dissolved oxygen can be formed. From our theoretical study, we know that at lower coverage, oxygen will prefer to stay on the surface, instead of in the subsurface or bulk region, and for high O concentrations, oxide-like structures are preferred.\cite{21} Furthermore, our results showed that substitutional oxygen adsorption in the bulk (i.e., O on Ag sites) is energetically unfavorable.\cite{21}

B. O adsorption at undercoordinated metal sites: Ag vacancies

The importance of steps and vacancies as active sites has been well documented, from both experimental and theoretical points of view.\cite{69,70} In our previous studies we investigated the binding of oxygen atoms on surfaces with preexisting vacancies.\cite{20} We found the binding energy is stronger than on the (111) terraces, e.g., for 0.25 ML adsorbed in fcc sites on a (2×2)/Ag(111) vacancy array the binding energy is 3.83 eV while it is 3.52 eV on the perfect terrace, at the same coverage. A similar behavior can be expected for adsorption at undercoordinated sites at step edges, as has been shown theoretically for oxygen at other transition metals, see, e.g., Refs. 59 and 69. To investigate the effect of pressure and temperature on the stability of such O species, we calculated the Gibbs free energy of formation. We find that under atmospheric pressure, the O atoms are stable to around 770 K (including corrections: 880 K) on a (3×3)/Ag(111) vacancy array, and even to 865 K (including corrections: 975 K) on a (2×2)/Ag(111) vacancy array. The existence and concentration of these species, however, clearly depends on the number of defect sites at the surface. In view of the above, it could be speculated that the TPD signal of "O$_2^*$" is due not to substitutional diffusion of oxygen atoms from the bulk to the surface as proposed, but due to O atoms adsorbed at undercoordinated surface silver sites.

C. Adsorption of O$_2$ at an Ag vacancy

As mentioned in the Introduction, in the literature, the existence of surface Ag vacancies has been proposed to stabilize a molecular-like ozone (O$_3$) species, which has been suggested to be active for the partial oxidation of ethylene.\cite{19}

As reported in Ref. 21, we calculated the adsorption energy of such a species on (3×3) and ($\sqrt{3}$×$\sqrt{3}$)R30° vacancy arrays, corresponding to oxygen coverages of 1/3 and 1 ML, respectively. We found that these species were energetically unfavorable compared to the commensurate oxide-like structures described above. If, however, we assume that Ag vacancies exist, i.e., neglecting the energy cost to create the vacancy, then the binding energy is the most favorable of all for coverage 1 ML. We can estimate the concentration of such Ag vacancies using our calculated values of the vacancy formation energy, together with an Arrhenius type equation: For a ($\sqrt{3}$×$\sqrt{3}$)R30° vacancy array with a vacancy concentration of 1/3 ML, the formation energy is 0.53 eV; it is 0.44 eV for a concentration of 0.25 ML, and 0.55 eV for a concentration of 0.11 ML. Considering a temperature of 550 K, corresponding to that around which epoxidation of ethylene takes place, this implies a concentration of thermally induced surface vacancies of less than about 10$^{-5}$−10$^{-4}$ ML, which is very small. Furthermore, calculation of the free energy shows that this species is less stable than the others. Our results thus do not support the molecular oxide-like species as playing an important role, but point to oxygen atoms of the thin (4×4) surface oxide as being the main species uptaking epoxidation of ethylene. This has been recently supported by other DFT calculations.\cite{71}

We would like to mention that several groups have recently reported DFT investigations into the adsorption of ethylene on silver surfaces,\cite{72,73} and also into the reaction of ethylene with chemisorbed on-surface O (Refs. 74, 75) with the aim of gaining a deeper understanding of the epoxidation reaction. As we have shown, however, it is likely that under the conditions of catalysis the thin oxide-like structure is present (with possibly some on-surface chemisorbed atomic oxygen also present), and thus it is of interest to study the interaction and reaction of ethylene with the thin oxide-like surface. Such an investigation has very recently been reported\cite{75} where it was found that the reaction mechanism is similar for both the oxide-like surface and for the surface with chemisorbed O atoms. The most highly activated step of the cycle was, however, found to be smaller (by ~0.18 eV) for the thin oxide-like surface. Although suggestive, this alone does not confirm that this surface is most reactive, and a full kinetic modeling is awaited.

D. Thick oxide formation at Ag(111)

As seen from Fig. 7, our calculations predict that thick oxide-like films can form at Ag(111), but only for low temperatures [e.g., ≤350 K (460 K including corrections) at atmospheric pressure]. The extremely low sticking coefficient of O$_2$ at Ag(111),\cite{14} as well as the found low thermal stability of these structures, may hinder their formation. For example, at high temperatures at which diffusion of O (or Ag) atoms can effectively occur, the material may become unstable. Furthermore, when using NO$_2$ to deliver O atoms to the surface, the substrate has to be held at temperatures higher than the temperature at which di-silver oxide decomposes, in order to desorb the NO molecules. Using, however, low temperatures and atomic oxygen or ozone (which readily disso-
cates at the surface) instead of $\text{O}_2$ (or $\text{NO}_2$), we predict that thick oxide structures could be observed experimentally—providing that diffusion barriers can be overcome.

Interestingly, it was found that di-silver oxide ($\text{Ag}_2\text{O}$) can be prepared using a mixture of ozone and $\text{O}_2$ or microwave generated atomic oxygen, at room temperature on polycrystalline silver foil.76–79 The oxygen generated in these processes is quite active, for example, small amounts of ozone (5 mol. % $\text{O}_3$ in $\text{O}_2$) can generate an O partial pressure equivalent to that of a pure $\text{O}_2$ atmosphere at $10^{-12}$ atm at a temperature of 465 K.77 Actually, in these mixtures of reactants, the oxygen generated in these processes is quite active, for example, small amounts of ozone (5 mol. % $\text{O}_3$ in $\text{O}_2$) can generate an O partial pressure equivalent to that of a pure $\text{O}_2$ atmosphere at $10^{-12}$ atm at a temperature of 465 K.77

The importance of defects. Also diffusion of oxygen and silver atoms is seen significantly improved significantly due to the non-uniformity of the growth. We note that subsequent to completion of our studies, very recent experimental work reported thick oxide formation at Ag(111) by using a flux of atomic oxygen,80 thus supporting our findings.

VII. CONCLUSION

Through density-functional theory calculations, and incorporating the effect of the atmospheric environment, we obtained the pressure-temperature phase diagram for the oxygen/Ag(111) system, which reveals important insights into this system and its function as an oxidation catalyst: The calculations predict that a thin surface-oxide structure is most stable in the temperature and pressure range of ethylene epoxidation and we propose it (and possibly other similar structures) contains the main O species actuating the catalysis. Low coverages of chemisorbed O become more stable for higher temperatures and could also play a role in other oxidation reactions. The only species stable for temperatures in excess of 720 K (including corrections: 830 K), are O atoms adsorbed at undercoordinated surface Ag atoms. Thick bulklike di-silver oxide can safely be ruled out as playing an important role in the technical catalytic reactions due to the low thermal stability, causing decomposition at temperatures lower than those used in the reactors. They are, however, the most stable structures for strongly O-rich conditions and low temperatures (e.g., up to 330–460 K at atmospheric pressure). Interestingly, the electronic structure of thick oxide surfaces are very similar to that of the thin (4×4) surface oxide structure, which raises the question of whether di-silver oxide might be used as a low-temperature oxidation catalyst.

A molecular ozonelike species adsorbed at a surface vacancy, as had been proposed in the literature as being the active species for the epoxidation of ethylene, is energetically unfavorable, as is bulk-dissolved oxygen, which strongly questions the latters hitherto thought crucial role in, e.g., the partial oxidation of methanol to formaldehyde.

We also studied the bulk and (111) surface properties of Ag$_2$O, the most stable silver oxide. The interaction of oxygen and silver is via a weak hybridization between Ag-$4d$-$5s$ and O-$2p$ orbitals, with substantial occupation of antibonding orbitals, which explains its low stability. Various terminations of the Ag$_2$O(111) surface were investigated as a function of the oxygen chemical potential and it is found that the surface with oxygen in the hollow site (i.e., the stoichiometric surface) is most favorable for all of the range of the O chemical potential considered, while the Ag-terminated surface and surfaces with additional oxygen adsorbed on top of exposed CUS surface Ag atoms are unfavorable. On investigating the energetics of a transition from commensurate oxide-like structures to the “real” (unstrained) oxide, we find that the transition becomes favorable only at nine O-Ag-O trilayers, which corresponds to $\sim$4.5 ML oxygen.

The approach of “ab initio” atomistic thermodynamics, i.e., of calculating free energies for many possible atomic geometries and predicting the lowest energy structures in contact with species in a gas-phase environment, is a promising approach to investigate a system under different pressure and temperature conditions, in particular, from those of typical theoretical surface science, right up to those of technical heterogeneous catalysis. The obtained phase diagram also shows that if one stays within the region of one phase or along a phase boundary, then a bridging of the pressure gap should be possible. Furthermore, it may be expected that the regions which lie close to the boundaries in the computed surface phase diagram may exhibit enhanced thermal fluctuations, i.e., here the dynamics of atomistic processes may be particularly strong, and this may be particularly relevant for catalysis under realistic conditions. Finally, we note, however, that an understanding of a full catalytic cycle, requires a kinetic modeling that includes all reactant species and intermediates, which for complex systems such as those discussed in the present paper, has not yet been attempted and is a challenge for the future.

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We compared results obtained using the GGA and the local density approximation (LDA) for two selected structures i.e. on-surface O with coverage 1/4 ML and the 0.50 ML oxidelike structure. Although the absolute value of the free energy is 14.6 and 14.3 meV/Å² larger, respectively, within the LDA, the phase boundary position differs by only 0.021 eV suggesting that the whole phase diagram is very similar within LDA and GGA.


Long Li and Judith C. Yang (unpublished).