Formaldehyde Selectivity in Methanol Partial Oxidation on Silver: Effect of Reactive Oxygen Species, Surface Reconstruction, and Stability of Intermediates

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ABSTRACT: Selective oxidation reactions on heterogeneous silver catalysts are essential for the mass production of numerous industrial commodity chemicals. However, the nature of active oxygen species in such reactions is still debated. To shed light on the role of different oxygen species, we studied the methanol oxidation reaction on Ag(111) single-crystal model catalyst surfaces containing two dissimilar types of oxygen (electrophilic, Oe and nucleophilic, On). X-ray photoelectron spectroscopy and low energy electron diffraction experiments suggested that the atomic structure of the Ag(111) surface remained mostly unchanged after accumulating low Oe coverage at 140 K. Temperature-programmed reaction spectroscopic investigation of low coverages of Oe on Ag(111) revealed that Oe was active for methanol oxidation on Ag(111) with a high selectivity toward formaldehyde (CH₂O) production. High surface oxygen coverages, on the other hand, triggered a reconstruction of the Ag(111) surface, yielding Ag oxide domains, which catalyzes methanol total oxidation to CO₂ and decreases the formaldehyde selectivity. This important finding indicates a trade-off between CH₂O selectivity and methanol conversion, where 93% CH₂O selectivity can be achieved for an oxygen surface coverage of θO = 0.08 ML (ML = monolayer) with moderate methanol conversion, while methanol conversion could be boosted by a factor of ~4 for θO = 0.26 ML with a suppression of CH₂O selectivity to 50%. Infrared reflection absorption spectroscopy results and density functional theory calculations indicated that Ag oxide contains dissimilar adsorption sites for methoxy intermediates, which are also energetically less stable than that of the unreconstructed Ag(111). The current findings provide important molecular-level insights regarding the surface structure of the oxidized Ag(111) model catalyst directly governing the competition between different reaction pathways in methanol oxidation reaction, ultimately dictating the reactant conversion and product selectivity.

KEYWORDS: methanol oxidation, selective catalytic oxidation, silver catalyst, formaldehyde, atomic oxygen

INTRODUCTION

Silver is one of the essential metals used in various selective catalytic oxidation (SCO) reactions carried out in the chemical industry. Ethylene oxidation yielding ethylene oxide (CH₂OCH₂) and methanol oxidation producing formaldehyde (CH₂O) are among the most prominent SCO reactions occurring on heterogeneous silver catalysts.¹,² These products are used in the synthesis of numerous commodity chemicals and are also utilized directly in other chemical processes. For example, ethylene oxide is used to produce pharmaceuticals, detergents, and plastics, while formaldehyde is heavily consumed to make a variety of chemicals for the construction, automotive, and furniture sectors.³⁻⁴ Recent reports indicate that the global annual consumption of ethylene oxide and formaldehyde is over 35 million tons, each.³⁻⁵ The annual global production of these chemicals is estimated to grow monotonically due to the increasing demand by the chemical industry. Thus, the SCO continues to be an active field of not only applied but also fundamental research.⁶⁻⁷

A critical challenge in heterogeneous SCO is to understand the nature of the oxygen species and their catalytic role under reaction conditions. Particularly, distinguishing various oxygen species responsible for the partial and complete oxidation pathways is essential in order to design novel heterogeneous catalysts with high reactivity and selectivity. Along these lines,
there exist immense efforts in the literature focusing on the identification of different oxygen species that can be generated on model catalytic Ag surfaces,\textsuperscript{8–10} as well as their chemical interactions with reactants such as ethylene\textsuperscript{25–27} and methanol.\textsuperscript{28–31} In former studies, two different types of oxygen species were identified via X-ray photoelectron spectroscopy (XPS) revealing distinct electronic properties.\textsuperscript{16,17,27} The O 1s XPS signals for these species were determined to be at 530.3 and 528.3 (±0.2) eV and were attributed to the so-called "electrophilic (O\text{\textsuperscript{\text{-}E})}" and "nucleophilic (O\text{\textsuperscript{\text{-}N})" oxygen species, respectively. While nucleophilic oxygen is commonly considered to be a strongly bonded oxygen atom in Ag oxide structures, the precise nature of the electrophilic oxygen is still being debated.\textsuperscript{32–34,44} Moreover, the roles of these dissimilar oxygen species in the SCO are yet to be understood at the molecular level.

Apart from the electronic properties, the thermal stability of the oxygen species is also a key factor in catalytic oxidation selectivity. Former ultra-high vacuum (UHV) studies showed that the atomic oxygen is stable on the Ag(111) surface only up to ca. 600 K and cannot be present on the model catalyst surfaces during the methanol oxidation reaction conditions involving elevated temperatures such as ca. 900 K.\textsuperscript{35} Instead, surface-embedded atomic oxygen (O\text{\textsuperscript{\text{-}A})}, which is stable up to 900 K was suggested to be the reactive oxygen species in methanol oxidation.\textsuperscript{18,30,36} In contrast, a computational microkinetic modeling study argued that surface atomic oxygen species could be present at 900 K under oxygen pressures above 1 kPa.\textsuperscript{37} Furthermore, recent studies revealed evidence that "kinetically trapped" or metastable oxygen species can form under reaction conditions and become decisive in product selectivity.\textsuperscript{38,39} Such discrepancies in the literature clearly indicate that there are still unresolved fundamental questions regarding the nature of oxygen species on catalytic silver surfaces and their role in the partial oxidation reactions.

Former investigations on the nature of oxygen species in the SCO over heterogeneous silver catalysts were mostly performed under oxygen-rich conditions.\textsuperscript{16,30,41} Such reactivity studies typically assumed that a Ag oxide surface with a p (4 × 4) structure was the dominant phase under reaction conditions because this was the dominant phase at oxygen pressures close to the ambient conditions.\textsuperscript{30} In contrast, formaldehyde production in the industry is typically carried out under methanol-rich conditions (e.g., CH\textsubscript{3}OH/O\textsubscript{2} = 2.0–2.5).\textsuperscript{32,42} A former in situ Raman study conducted at atmospheric pressure and elevated temperatures showed that the polycrystalline silver existed in a reduced (i.e., metallic) state during methanol oxidation.\textsuperscript{43} A recent computational study also showed that a low oxygen surface coverage (<0.2 ML, ML = monolayer) was needed for high formaldehyde selectivity in methanol oxidation.\textsuperscript{9} These studies suggest that the precise control of the low surface coverage oxygen species on the catalyst surface is essential in the SCO of methanol.

While methanol oxidation reaction on p (4 × 4) reconstructed Ag oxide surfaces on the Ag(111) substrate was investigated in a former report,\textsuperscript{44} the literature lacks a comparative study examining methanol oxidation on atomic oxygen-containing Ag(111) model catalyst surfaces [O/Ag(111)] and Ag oxide surfaces. In the current work, we focus on these two dissimilar surface states of the silver model catalyst surfaces. Along these lines, we present experimental findings indicating that various types of adsorbed methoxy intermediates having different stabilities can be generated on O/Ag(111) and Ag oxide domains during methanol oxidation reaction. We used ozone decomposition as an effective oxygen delivery method to prepare O/Ag(111) and Ag oxide model catalyst surfaces, enabling the precise control of the quantity of the oxygen species introduced on the model catalyst surface particularly at low ozone partial pressures (e.g., <5 × 10\textsuperscript{−10} Torr). Chemisorbed oxygen atom formation with this method was previously reported on various model catalyst surfaces such as Au(111)\textsuperscript{45} and Pt(111)\textsuperscript{46} under UHV conditions. This method also precludes the use of high-pressure adsorbate exposures which may lead to excessive surface oxygen coverages and/or deposition of unwanted contaminants on the model catalyst surface. In contrast to the elevated/ambient pressure O\textsubscript{2} experiments, the currently employed oxygen delivery protocol allows the preparation of unreconstructed O/Ag(111) surfaces possessing very low (e.g., 0.04 ML) oxygen adsorbate coverages. Because the surface reconstruction of Ag(111) and formation of silver oxide domains typically start at oxygen coverages above 0.05 ML,\textsuperscript{47} it is challenging to study unreconstructed O/Ag(111) surfaces under elevated pressures, rendering the currently utilized oxygen delivery protocol preferable.

\section*{METHODS}

\subsection*{Experimental Details.} All experiments were performed in a UHV chamber with a background pressure of <5.0 × 10\textsuperscript{−10} Torr which was equipped with low-energy electron diffraction (LEED; custom-made), infrared reflection absorption spectroscopy (IRAS; Bruker Tensor37 with custom design external IR optics and a tailor-made N\textsubscript{2} purge box, 4 cm\textsuperscript{−1} resolution, 100 scans, 10 kHz scanner velocity, and mercury cadmium telluride IR detector), XPS (Riber nonmonochromatized Al K\textalpha\ X-ray excitation, 300 W, Riber EA150 electron energy analyzer), and temperature-programmed reaction spectroscopy (TPRS) capabilities.

The Ag(111) single crystal (Matech GmbH, 10 mm diameter × 1 mm thick disk, >99.9999% purity) 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 sample was cleaned by cycles of Ar\textsuperscript{+} sputtering [Ar(g), Linde AG, 99.999% purity] with a sputter gun (LK Technologies, NGI3000, 1.5 kV × 15 mA) and repetitive cycles of O\textsubscript{3} treatment at room temperature, followed by annealing to 800 K for 5 min.

TPRS experiments were performed by using a quadrupole mass spectrometer (QMS; Ametek Dycor Dymaxion DM200) and a proportional–integral–derivative controlled linear sample heater (Heatwave, model 101303). All of the TPRS experiments were performed with a heating rate of 1 K/s, a 70 eV QMS electron ionization energy, and a 30 ms dwell time for each desorption channel. Methanol (Sigma-Aldrich, 99.9% purity) was purified by several freeze–pump–thaw cycles in a glass bulb connected to a gas manifold and dosed onto the sample via a leak valve. In order to eliminate the molecular (undissociated) methanol adsorption and generate predominantly methoxy species with a high surface coverage on O/Ag(111), in each of the TPRS experiments, 0.06 Langmuir (1 L = 10\textsuperscript{−6} Torr × s\textsuperscript{−1}) of methanol [i.e., corresponding to 2
monolayer equivalent, monolayer equivalent (MLE) of methanol dose at 100 K) was introduced at 200 K on O/Ag(111). In the TPRS experiments, \( m/z = 2, 18, 28, 29, 30, 31, 32, 44, \) and 46 desorption channels were simultaneously monitored (see Supporting Information, SI Section 1, Figure S1).

The ozone decomposition method was used for the delivery of atomic oxygen on the Ag(111) model catalyst surface under UHV conditions. Oxygen was produced by a commercial ozone generator (Genozon, GN-G1001S, Turkey) and directly dosed on the sample through a leak valve. Additional details regarding the ozone delivery system can be found elsewhere. In that report, the oxygen desorption signal converged to saturation at 550 K, and the saturation coverage was designated as 1 MLE. In former studies, oxygen coverage for a single layer of silver oxide with saturation coverage obtained at the top layer of Ag(111) in the oxygen TPD experiments was assumed to be 0.375 ML and used as a reference.

**Results and Discussion**

**Methanol Reactivity on Ag(111) and O/Ag(111).** As an initial control experiment, the reactivity of the clean Ag(111) surface toward methanol was investigated using TPD, where various coverages of methanol were dosed on clean Ag(111) at 100 K (Supporting Information, SI Section 2, Figure S2). In accordance with the literature, in the absence of oxygen,

\[
E_{\text{ads}} = E_{\text{tot}} - E_{\text{surf}} - E_{\text{mol}}
\]

where \( E_{\text{tot}} \), \( E_{\text{surf}} \), and \( E_{\text{mol}} \) denote the total energy of the system, the energy of the surface, and the energy of the molecule in the gas phase, respectively. According to eq 1, negative adsorption energies correspond to exothermic adsorption.
methanol was found to adsorb and desorb reversibly from the clean Ag(111) surface in a molecular fashion without dissociation at T ≤ 170 K. Clearly, the Ag(111) surface was incapable of initiating methanol bond activation in the absence of the oxygen species. Molecular methanol adsorption on clean Ag(111) was also verified with IRAS and is discussed in detail below.

However, the interaction of methanol with oxygen-containing Ag(111) surfaces [O/Ag(111)] generated formaldehyde and carbon dioxide in TPRS experiments as the SCO and total oxidation products, respectively (Figures 1 and S1). In the current TPRS experiments, methanol exposure at 200 K on O/Ag(111) surfaces containing 0.04 or 0.08 ML of oxygen mainly yielded formaldehyde (Figure 2a) which desorbed at 350 K. In addition to formaldehyde production, a weak CO2 desorption signal was also observed at 365 K (Figure 2b) for oxygen coverages (θO) of 0.04 and 0.08 ML, indicating only a limited extent of total oxidation. However, the CO2 signal intensity increased with increasing oxygen coverage (i.e., θO > 0.08 ML).

Additional quantitative information was extracted from the current TPRS results, revealing the variations in catalytic selectivity (Figure 1c) and catalytic conversion (Figure 1d) as a function of oxygen coverage. The TPRS results shown in Figure 1c,d clearly indicate that catalytic selectivity and activity of the Ag(111) surface change drastically as a function of oxygen coverage for identical methanol exposures and identical dosing temperatures. While θO ≤ 0.08 ML favors the SCO of methanol to formaldehyde with an enhanced formaldehyde selectivity that can reach up to 93%; θO ≥ 0.26 ML promotes the total oxidation pathway by boosting the CO2 selectivity to 74% at θO = 0.49 ML. As shown in Figure 1d, the combined total amount of formaldehyde and CO2 production also increases by ca. 375% from θO = 0.04 ML to θO = 0.26 ML, followed by some decrease at θO = 0.49 ML. Thus, it is apparent that a trade-off exists between the selectivity and total catalytic conversion of methanol. Accordingly, while formaldehyde selectivity can be maximized at low surface oxygen coverages, maximum methanol conversion is obtained at higher oxygen coverages.

One of the important aspects of the reactivity results shown in Figure 1a is the fact that increasing oxygen coverage not only alters the formaldehyde yield but also modifies its desorption kinetics noticeably. Another formaldehyde desorption feature at 282 K was obtained for θO = 0.26 in addition to the existing formaldehyde signal at 350 K; while the 350 K peak disappears for θO = 0.49. Based on these experimental observations illustrating different reactivity trends of methanol on oxygen-containing silver surfaces, we hypothesize that the observed reactivity trends originate from the following causes: (a) differences in the electronic nature of the oxygen species generated on Ag(111), (b) variations in surface morphology of the Ag(111) model catalyst, and (c) relative adsorption energies and adsorption geometries of the generated intermediates or products during the methanol oxidation reaction at various oxygen coverages. We will address and corroborate each of these points in the following sections.

**Electronic Nature of Oxygen Species on Ag(111).** In an attempt to shed light on the chemical and electronic nature of the oxygen species formed on Ag(111) as a function of oxygen coverage, the XPS spectra of clean and oxygen covered Ag(111) surfaces were obtained by exposing the Ag(111) crystal to different exposures of ozone at 140 K. Based on the corresponding O 1s binding energy (B.E.) values, the electrophilic oxygen (Oe) species (530.3 eV) were observed to form on the Ag(111) surface at low oxygen coverages (i.e., θO = 0.20 ML), while the nucleophilic oxygen (On) species (528.4 eV) were dominant at higher coverages as a result of silver oxide formation (Figure 2a). These assignments are in good accordance with the former studies which reported the O 1s B.E. values for O on within 530.0–535.0 eV on silver8,13,15,18 and 528.1–528.5 eV for On of silver oxide films with p(3 × 3)rect, p(3 × 5)rect, or p(4 × 5)rect structures.8,13,15,18

Figure 2a shows that upon increasing the oxygen surface coverage from 0.20 to 1.00 ML, the integrated XPS signals for Oe and On species increase by a factor of 4 and 8, respectively. Two times greater boost in the On signal as opposed to that of Oe provides evidence that high oxygen coverages induce silver oxide formation (along with surface reconstruction as discussed below) even at temperatures as low as 140 K. Based on the former studies in the literature, the growing signal at 530.3 eV for θO > 0.4 ML can also be attributed to the silver oxide formation because the O 1s signal for the multilayer (bulk-like) silver oxide species (accompanied by the formation of subsurface oxygen) appears at a similar B.E. to...
the O<sub>e</sub> species. Therefore, it is likely that three-dimensional silver oxide growth also contributes to the signal at 530.3 eV. Silver oxide formation is also consistent with the currently observed \(-1.1\) eV core-level shift of the Ag 3d signal with respect to that of the metallic silver signal (Figure 2b).

It is worth mentioning that another type of atomic oxygen on unreconstructed Ag(111) surfaces with a characteristic O 1s signal at 527.9 eV was also reported in the literature at very low oxygen coverages (\(\theta_O < 0.04\) ML). Due to the detection limit of the currently utilized photoelectron detector, in the current work, the lowest oxygen coverage yielding a reliable O 1s signal was 0.20 ML. Thus, the presence of such minority species in the currently presented XPS data at low coverages (e.g., \(<0.20\) ML) cannot be ruled out.

**Morphology of the Oxygen-Containing Ag(111) Surfaces.** The current LEED results (Figure 2c–f) verified that silver oxide formation and surface reconstruction events were relatively insignificant for low oxygen coverages at 140 K. A sharp (1 \(\times\) 1) LEED pattern that is characteristic of a clean Ag(111) surface was obtained for the O(0.08 ML)/Ag(111) surface at 140 K (Figure 2d), indicating the lack of (or negligible) surface reconstruction at this oxygen coverage. XPS results shown in Figure 2a also provides additional evidence for negligible Ag oxide formation at low oxygen coverages. In contrast, higher oxygen coverages (i.e., \(\theta_O > 0.08\) ML) on Ag(111)-induced surface reconstruction due to silver oxide formation is evident by the diffuse (1 \(\times\) 1) LEED pattern obtained for \(\theta_O = 0.26\) ML on Ag(111) at 140 K (Figure 2e). A further increase in the oxygen coverage to 0.49 ML boosted the surface corrugation and reconstruction, resulting in the complete disappearance of the LEED pattern (Figure 2f). It is worth mentioning that the 0.49 ML oxygen coverage was sufficient to generate a silver oxide surface which was also verified by annealing the O(0.49 ML)/Ag(111) surface originally prepared at 140–450 K in UHV (to improve ordering). Such a protocol revealed the formation of ordered silver oxide domains (i.e., \(c(4 \times 8)\) structure) with characteristic and well-defined LEED patterns which were discussed in one of our former reports.

![Figure 3](https://doi.org/10.1021/acscatal.1c00344)

**Figure 3.** Optimized adsorption sites and their corresponding adsorption energies calculated via DFT computational models for different adsorbates on (a) clean Ag(111) and unreconstructed O/Ag(111) and (b) reconstructed p(4×4)–O(0.375 ML)/Ag(111) surface.

Our findings demonstrate that oxygen accumulation on Ag(111) by ozone exposure resulted in surface structures that are comparable to the existing studies in the literature utilizing different oxygen delivery methods. Former scanning tunneling microscopy studies showed that for low oxygen coverages (\(\theta_O < 0.05\) ML), oxygen atoms were randomly located on Ag(111) without a surface reconstruction, while at higher oxygen coverages (\(\theta_O > 0.05\) ML), silver oxide domains coexisted with unreconstructed Ag(111) regions. Very high oxygen coverages (\(\theta_O > 0.8\) ML) on Ag(111) led to bulk-oxide formation.

In light of the XPS and LEED data shown in Figure 2, TPRS reactivity trends shown in Figure 1a can be discussed in more depth. It can be argued that for \(\theta_O \leq 0.08\) ML, O<sub>e</sub> species are the prevalent species on the unreconstructed Ag(111) surface catalyzing high-temperature formaldehyde production at 350 K, while O<sub>s</sub> species exist as minority species at these coverages promoting the formation of only minuscule amounts of CO<sub>2</sub> at 370 K. On the other hand, increasing oxygen coverage (i.e., for \(\theta_O = 0.26\) ML) and the emergence of O<sub>e</sub> species (initiating the growth of the Ag oxide overlayer) result in the observation of both the low-temperature formaldehyde production at 285 K as well as the formation of surging amounts of CO<sub>2</sub> at 370 K. At the highest oxygen coverage studied in the current TPRS experiments (\(\theta_O = 0.49\) ML), the Ag oxide layer which predominantly comprised O<sub>e</sub> species, almost totally suppresses the high-temperature formaldehyde production at 350 K, facilitating low-temperature formaldehyde production at 285 K, while also enhancing CO<sub>2</sub> production. Note that this particular oxygen coverage (\(\theta_O = 0.49\) ML) not only significantly boosts the CO<sub>2</sub> production but also lowers the CO<sub>2</sub> formation temperatures to 350–310 K. A former study reported that formaldehyde desorption occurred at 290 K for methanol reaction on the commonly observed reconstructed p(4 \(\times\) 4) Ag oxide overlayer on Ag(111). Thus, the new formaldehyde desorption feature shown in Figure 1a at 282 K can be readily attributed to the methanol activation on the reconstructed Ag oxide overlayer.

**Relative Stabilities and Adsorption Geometries of the Intermediates and Products.** Optimized adsorption
geometries and the adsorption energies of selected species (i.e., methanol, methoxy, formaldehyde, and formate) acting as reactants, intermediates, or products in the methanol oxidation were investigated using DFT computational modeling on the oxygen-containing unreconstructed Ag(111) surface as a function of oxygen coverage (Figure 3a) and on the reconstructed p(4 × 4)–O(0.375 ML)/Ag(111) surface (Figure 3b). Due to the repulsive interaction between oxygen adatoms that destabilize the surface at high coverages, we only report results on the O/Ag(111) surface as shown in Figure 3a for oxygen coverages up to 0.33 ML. Methanol and formaldehyde bind to the clean surface only through weak van der Waals forces, as evidenced by the low adsorption energies. However, as the concentration of O atoms increases, there is a marked increase in the corresponding adsorption energies of both species due to the attractive interaction between the O atoms on the surface and the H atoms of the molecules via hydrogen bonding. Methoxy and formate, on the other hand, are already adsorbed strongly onto the clean Ag(111) surface. While methoxy stability is appreciably enhanced by the presence of surface oxygen species, the adsorption energy of formate species is relatively weakly altered by the oxygen coverage variations.

In contrast to the O/Ag(111) surfaces, methoxy species display widely varying adsorption energies on the reconstructed p(4 × 4)–O/Ag(111) surface, indicating a much broader assortment of adsorption geometries (Figure 3b). While the most favorable methoxy geometry yields similar adsorption energy on the O/Ag(111) and p(4 × 4)–O/Ag(111) surfaces at a comparable oxygen coverage (i.e., 2.04 eV at θO = 0.33 ML and 2.13 eV at θO = 0.375 ML, respectively), the reconstructed Ag oxide surface provides multiple adsorption sites for methoxy species, yielding an adsorption energy range between 1.22 and 2.13 eV. Formate adsorption, on the other hand, proves to be relatively independent of the silver surface structure and the adsorbate environment. Formate, the species with the highest adsorption energy on both surfaces, is found to be exclusively in the bidentate adsorption geometry. These findings suggest that adsorption energy and thermal stability of both methoxy and formate intermediates are lower for the Ag oxide overlayers on the Ag(111) surfaces. Overall, the results presented in Figure 3 reveal that the investigated adsorbates can be ranked in the following increasing order of stability: formaldehyde on Ag(111) or unreconstructed O/Ag(111) < methanol on Ag(111) or unreconstructed O/Ag(111) < methoxy on reconstructed Ag oxide < methoxy on Ag(111) or on unreconstructed O/Ag(111) < formate on reconstructed Ag oxide < formate on Ag(111) or unreconstructed O/Ag(111).

The surface functional groups generated during the interaction of methanol with the clean or oxygen-containing Ag(111) model catalyst surface and their corresponding adsorption geometries were studied via IRAS (Figure 4). On the clean Ag(111) surface lacking any oxygen, methanol adsorbs in a molecular fashion (Figure 4a) and desorbs from the surface at <170 K (Figure S2) without being activated to methoxy species. Accordingly, characteristic vibrational signals of molecular methanol detected upon saturation methanol exposure (i.e., 0.03 L) on the clean Ag(111) surface at 100 K (Figure 4a and Table 1) are in very good agreement with former TPD and IRAS results for methanol on pristine Ag(111).\(^{44,55}\)

![Figure 4](https://doi.org/10.1021/acscatal.1c00344)

Table 1. IR Vibrational Frequencies and Vibrational Mode Assignments for Methanol, Methoxy, and Formate Species on Ag(111), O/Ag(111), and Ag Oxide Overlayers on Ag(111)*

<table>
<thead>
<tr>
<th>vibrational modes</th>
<th>methanol</th>
<th>methoxy</th>
<th>formate</th>
<th>ref</th>
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<tr>
<td>(\nu(\text{CO}))</td>
<td>1035 a</td>
<td>1041 b</td>
<td>1036 c</td>
<td>44</td>
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<tr>
<td>(\delta_{\text{CH}})</td>
<td>1454 a</td>
<td>1430 b</td>
<td>1470 c</td>
<td>44</td>
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<tr>
<td>(\nu_{\text{HC}})</td>
<td>2820 a</td>
<td>2790 b</td>
<td>2788 c</td>
<td>44</td>
</tr>
<tr>
<td>(2\delta_{\text{CH}})</td>
<td>2873 b,c</td>
<td></td>
<td></td>
<td>44</td>
</tr>
<tr>
<td>(2\nu_{\text{CH}})</td>
<td>2917 b,c</td>
<td>2909 b,c</td>
<td></td>
<td>44</td>
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<tr>
<td>(\nu_{\text{OCO}})</td>
<td>2939 a</td>
<td></td>
<td></td>
<td>44</td>
</tr>
<tr>
<td>fermi resonance doublet of (\nu(\text{CH}))</td>
<td>1320 a</td>
<td></td>
<td></td>
<td>58</td>
</tr>
<tr>
<td>with 2(\delta(\text{CH})) or (\nu_{\text{OCO}} + \delta(\text{CH}))</td>
<td>2807 b &amp; c</td>
<td>2883 b &amp; c</td>
<td>58</td>
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On the other hand, on oxygen-containing Ag surfaces (Figure 4b,c), methanol can be readily activated to methoxy species via O–H bond cleavage even at temperatures as low as 100 K. While vibrational signatures of the generated methoxy species on the reconstructed Ag oxide overlayer (Figure 4c and Table 1) resemble that of the unreconstructed O/Ag(111) surface (Figure 4b and Table 1), the former surface reveals weaker and broader IRAS signals. The broadening of the IRAS signals, as shown in Figure 4c, can be explained by considering the increasing heterogeneity of the surface and the presence of nonuniform adsorption sites on the Ag oxide overlayer. Relatively weaker IRAS signal intensities, as shown in Figure 4c, can be attributed to multiple phenomena such as (i) decreasing IR absorption cross section of the adsorbed methoxy species due to the presence of the Ag oxide overlayer attenuating the image dipole strength in the underlying Ag(111) substrate\(^{57}\) and (ii) tilting of the surface methoxy species. The latter is likely to occur because varying adsorption geometries of methoxy species were predicted by DFT calculations on the reconstructed Ag oxide surface (Figure 3b). Methoxy species adsorbed on the unreconstructed O(0.04 ML)/Ag(111) surface in an upright configuration (Figure 4b) yields a sharp and intense \(\nu(\text{CO})\) signal at 1041 cm\(^{-1}\). Weakening and broadening of this signal for the reconstructed Ag oxide (\(\theta_O = 0.40 \text{ ML}\)) surface (Figure 4c) suggests that the
C–O bond axis of methoxy species was oriented in a tilted fashion on the reconstructed Ag oxide surface. It is important to note that the upright versus tilted methoxy geometry may lead to significant changes in bond-breaking processes and alter the reaction pathways on reconstructed/unreconstructed silver model catalyst surfaces.

Marked differences in the relative thermal stabilities of the reaction intermediates can be considered as an indication of the presence of distinct reaction pathways. The current IRAS and TPRS results demonstrate that the stability of methoxy intermediates is significantly different on the reconstructed Ag oxide compared to that of the unreconstructed O/Ag(111). For sufficiently low oxygen coverages (e.g., \( \theta_O = 0.04 \) ML) precluding the formation of Ag oxide overlayers, methoxy species generated at 100 K are extremely stable and manage to survive on the surface in an intact manner even at 300 K (Figure S3). In contrast, methoxy decomposition and subsequent formaldehyde desorption occur below 300 K on the reconstructed Ag oxide surface (Figure 1a). This experimental observation is in good accordance with the relative adsorption energy trends estimated by the current DFT calculations (Figure 3), suggesting higher methoxy stability on unreconstructed O/Ag(111).

In methanol SCO, formaldehyde production occurs via C–H bond rupture in the methoxy intermediate. Because adsorbed formaldehyde is extremely weakly bound to all of the currently investigated silver model catalyst surfaces (Figure 3), it can be safely argued that formaldehyde desorption occurs immediately after the C–H bond activation in the adsorbed methoxy species and production of formaldehyde on the catalyst surface. Based on the desorption signals of formaldehyde as shown in Figure 1a for O(0.26 ML)/Ag(111), the activation energy for methoxy decomposition on the reconstructed Ag oxide and unreconstructed Ag(111) can be estimated (assuming first-order kinetics\(^58\) and a pre-exponential factor of \(10^{13} \) s\(^{-1}\)) to be 75 and 94 kJ mol\(^{-1}\), respectively. The estimated activation energy difference of 19 kJ mol\(^{-1}\) indicates that C–H bond rupture in methoxy species is kinetically favored on the reconstructed Ag oxide overlayers as opposed to the unreconstructed O/Ag(111) surfaces. This is also consistent with the higher total methanol conversion on the reconstructed Ag oxide overlayers (Figure 1d). However, it is important to note that easier C–H activation of the methoxy species and the associated lowered formaldehyde desorption temperature on the reconstructed Ag oxide overlayers does not lead to an enhanced formaldehyde selectivity (Figure 1a,c). This is probably because methoxy activation also leads to boosted formate production on the reconstructed Ag oxide overlayers, where these formate species can be further oxidized and generate the total oxidation product, CO\(_2\), as indicated in a prior study.\(^44\) Relatively high adsorption energies of formate species on the reconstructed Ag oxide overlayers (Figure 3b) also suggest a long surface residence time for formate species, allowing them to be totally oxidized to CO\(_2\) rather than desorbing as other partial oxidation products (e.g., esters).

Along these lines, it can be argued that while methoxy species are more stable on the unreconstructed O/Ag(111) surface, this surface is inefficient in formate production, which is an essential step that should occur before CO\(_2\) production. Despite the high CO\(_2\) yield detected in the current TPRS experiments on the strongly oxidized O(0.49 ML)/Ag(111) (Figure 1), the lack of formate intermediates at 100 K in the IRAS data presented in Figure 4c indicated that formaldehyde production is kinetically hindered at 100 K, and further oxidation of methoxy species to formate occurred at a higher temperature (i.e., >100 K). Similarly, IRAS data show that the formate production rate was also low on unreconstructed O(0.04 ML)/Ag(111) at 100 K (Figure 4b), leading to the predominance of methoxy species on this surface. Thus, we infer that C–H bond rupture in methoxy yielding formaldehyde/formate is a kinetically hindered process, and formate intermediates could be generated via methanol exposure only at a temperature high enough to overcome the associated kinetic barrier.

Along these lines, formate production was readily achieved on O(0.10 ML)/Ag(111) at 300 K (Figure 4d). Figure 4d indicates that the IRAS spectrum acquired after 0.03 L of methanol exposure at 300 K on the O(0.10 ML)/Ag(111) surface, followed by cooling to 140 K, led to bridging bidentate formate species which adsorbed symmetrically on the surface via its oxygen atoms to two adjacent Ag atoms, where the C–H bond stayed orthogonal to the surface plane.\(^59\) These formate species facilitated the CO\(_2\) formation evidenced by the corresponding TPRS control experiment (Supporting Information Section 4, Figure S4) obtained after acquisition of the IRAS spectrum, as shown in Figure 4d. Thus, it can be argued that at sufficiently high surface oxygen coverages, the reconstructed silver oxide surface tends to react with formaldehyde at 300 K to produce formate, followed by complete oxidation to CO\(_2\).

The overall reactivity and selectivity trends of catalytic methanol oxidation on unreconstructed O/Ag(111) and reconstructed Ag oxide overlay on Ag(111) deduced from the current experimental and theoretical findings are summarized in Scheme 1.

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Scheme 1. Reactivity and Selectivity Trends of Methanol SCO on Unreconstructed O/Ag(111) and Reconstructed Ag Oxide Overlayer on Ag(111)

**Unreconstructed O/Ag(111) Surface**

**Reconstructed Ag Oxide Overlayer on Ag(111)**

<table>
<thead>
<tr>
<th>Temperature</th>
<th>Product</th>
<th>Reaction Pathway</th>
</tr>
</thead>
<tbody>
<tr>
<td>(50 K)</td>
<td>Methanol</td>
<td>Adsorption</td>
</tr>
<tr>
<td>(350 K)</td>
<td>Formaldehyde</td>
<td>C–H Bond Activation</td>
</tr>
<tr>
<td>(282-365 K)</td>
<td>Carbon dioxide</td>
<td>Complete Oxidation to CO(_2)</td>
</tr>
</tbody>
</table>
CONCLUSIONS
In the current work, we focused on the fundamental molecular origins of the catalytic selectivity in methanol oxidation over silver model catalyst surfaces using ozone as a uniquely efficient oxygen delivery vector. Particularly, we addressed three critical aspects governing catalytic activity and selectivity: (a) electronic nature of the oxygen species, (b) morphology of the silver model catalyst surfaces, and (c) stability of the intermediates and products formed during the methanol oxidation reaction on various silver model catalyst surfaces. Our results indicated that at least two different oxygen species existed on silver model catalyst surfaces, namely, electrophilic (Oe) and nucleophilic (On) oxygen. Surface oxygen coverages within 0.04 ML < θ ≤ 0.08 ML favored Oe species and mostly unreconstructed O/Ag(111) surfaces, yielding boosted formic acid selectivity but low total methanol conversion. In contrast, elevated surface oxygen coverages such as θ ≥ 0.26 ML predominantly generated On species, resulting in the reconstruction of the Ag(111) surface and formation of a Ag-O bond overlayer, enhancing methanol conversion to CO2 while suppressing selectivity toward formaldehyde. Surface stability of the methoxy and formate intermediates was found to be lower for the reconstructed Ag-O overlayers as compared to the unreconstructed O/Ag(111) surfaces, where the former surfaces promoted lowering of the activation energies for C–H bond scission in the methoxy intermediate and boosted CO2 production.

ASSOCIATED CONTENT
 Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acscatal.1c00344.
TPRS for methanol on O/Ag(111); methanol adsorption on clean Ag(111); IR spectra of methoxy on O(0.04 ML)/Ag(111) at 100 and 300 K; and temperature-programmed reaction of formate on O(0.10 ML)/Ag(111) (PDF)

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Notes
The authors declare no competing financial interest.

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REFERENCES


