New Insights into Ethene Epoxidation on Two Oxidized Ag\{111\} Surfaces

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Ethene epoxidation is an important heterogeneous catalysis process that takes place selectively on silver catalysts. The mechanism of this partial oxidation has been debated for many years, and to date a detailed atomic level understanding of this process has not been obtained. Much of the debate has centered on the nature of the active O species.\(^1,2\) Recently, two stable O phases on Ag\{111\} have been characterized with the scanning tunneling microscope (STM) and density functional theory (DFT).\(^3,4\) One is a low coverage (0.05 ± 0.02 ML) O adatom phase, and the other is an Ag\(_{1.8}\)O oxide overlayer. A DFT-derived phase diagram predicted that the Ag\(_{1.8}\)O oxide overlayer would be stable under typical industrial conditions for epoxidation.\(^5\) However, the temperature and pressure boundaries with the low coverage O adatom phase were close, implying that both phases could easily coexist in a reactor. Additionally, Barteau and co-workers made a significant step forward with the identification and characterization of a surface oxametallacycle on Ag\{111\} (labeled OME or OMME if they include one or more than one surface atoms, respectively).\(^5\) With the aid of high-resolution electron energy loss spectroscopy (HREELS) and DFT-based cluster calculations, they showed that this was a precursor to ethene epoxide formation.\(^6\) However, that study focused on a single elementary step for the reverse reaction starting with epoxide.

The partial oxidation reaction, C\(_2\)H\(_4\) + 1/2O\(_2\) → C\(_2\)H\(_4\)O, is exothermic by 117 kJ/mol at room temperature, and so the conversion is merely inhibited by kinetics. In this work, we use periodic DFT to determine accurate values for the kinetic barriers to the formation of ethene epoxide. Reaction mechanisms and their associated barriers for the complete catalytic cycle of the conversion of ethene to epoxide have been determined on the recently characterized low and high coverage oxygen phases. We find that on both catalysts epoxidation proceeds similarly via a two-step nonconcerted process. Concerted mechanisms in which O adds bonding directly to the surface. This OMME is the most stable of the eight oxametallacycles investigated, and its structure is similar\(^11\) to that predicted by previous cluster calculations.\(^6,12\) A barrier of 0.92 eV was then identified for OME ring closure to produce a weakly adsorbed ethene epoxide (0.09 eV). To determine the

Figure 1. Relative energy diagram for the conversion of gas-phase ethene into ethene epoxide (C\(_2\)H\(_4\)O) via an oxametallacycle (OMME) intermediate on Ag\{111\} + O. Energies shown in eV refer to the differences between adjacent equilibrium states, or between adjacent equilibrium and transition states. States to the right-hand side of the vertical dashed line do not contain epoxide molecules and are simply related to O\(_2\) dissociative adsorption on clean Ag\{111\} (O, red; C, gray; *, transition state).

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energetics of the entire catalytic cycle, the dissociation of O₂ on clean Ag[111] was also examined as the last regeneration step. A barrier of 0.64 eV, relative to a weakly adsorbed O₂ precursor state (E_ads = 0.17 eV), has been determined for this process.

The energetic profile for epoxidation on the high coverage Ag₁₁O oxide surface is displayed in Figure 2. A weak ethene adsorption precursor is first seen. This state (not shown) has recently been characterized in a combined STM and DFT study and shows ethene symmetrically adsorbed above an Ag of the oxide ring. The barrier to produce the OMME from this state is 0.74 eV (0.46 eV relative to the initial C₂H₄ gas-phase state). At the transition state, a C–O bond has been created through a lateral shift of ethene toward an adjacent oxygen in the underlying oxide. The structure of the OMME product of this step is 0.15 eV more stable than the initial C₂H₄ gas-phase state. It is worth noting that the extraction of an O from the oxide has resulted in a significant reconstruction of the oxide ring. In the next step, the second C–O bond is formed to produce an epoxide with a barrier of 0.74 eV. The epoxide binds weakly to the oxygen-deficient oxide (0.09 eV), above a triangle of Ag atoms. To mimic an entire catalytic cycle, a second O was removed from the oxide overlay (equivalent to performing a second epoxidation cycle), and O₂ dissociation was examined on this doubly reduced oxide overlay. The most favorable dissociation route identified, with a barrier of 0.40 eV, involves O₂ initially adsorbed parallel to the surface.

Overall, we see that on the two surfaces examined the epoxidation mechanisms are reasonably similar, in terms of both structures and energetics. On both catalysts, epoxidation proceeds via an OMME intermediate, and it is the ring closure of this OMME intermediate that is the most highly activated step of the cycle (with a barrier of 0.74 eV on the oxide and 0.92 eV on the O adatom phase). This lower barrier to OMME ring closure on the oxide surface could be explained by the reduced stability of the OMME intermediate on this surface. If we compare the barriers determined here to those from experiment, we find that satisfyingly they fall within the broad range reported (0.6–1.1 eV).

Other differences between the two substrates arise in the OMME formation step (first step) and the catalyst regeneration step (O₂ dissociation). Ethene is first trapped into a chemisorption state before reacting on the oxide, whereas on the O adatom phase it reacts directly from the gas phase. However, we anticipate that this state, with moderate binding of ethene to the oxide, will not be populated at reaction temperatures. This would render the first step similar on both surfaces. Finally, O₂ dissociation is favored on the reduced oxide surface (Ag₁₁O₄ in Figure 2), with a barrier of 0.40 eV as opposed to 0.64 eV on clean Ag[111]. Given that the reduced Ag₁₁O₄ oxide surface is unstable as compared to the equilibrium Ag₁₁O₃ (Ag₁₁O₃) oxide overlayer, a lower barrier to O₂ dissociation on this surface is to be anticipated.

In conclusion, mechanisms and barriers for the conversion of ethene to epoxide on two oxidized Ag surfaces have been determined. On both catalysts, epoxidation proceeds via an oxametallacycle intermediate, and it is the ring closure of this intermediate that is the most highly activated step. The particular question of which catalyst is a priori the most reactive goes beyond the present study and will be the subject of future kinetic modeling. Here, we conclude that both oxidized surfaces are efficacious selective oxidation catalysts and that Ag may act as a dual phase catalyst inside the reactor.

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**Supporting Information Available:** Geometries of the most important intermediates and transition states (PDF). This material is available free of charge via the Internet at http://pubs.acs.org.

**References**

11. Some discrepancy is found with the relative stability of the OMME and epoxide states as compared to the DFT results of ref 6. Possibly this is due to the different surface models used here and in ref 6.

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