Carbon Dioxide Activation and Reaction Induced by Electron Transfer at an Oxide–Metal Interface**

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**Abstract:** A model system has been created to shuttle electrons through a metal–insulator–metal (MIM) structure to induce the formation of a CO$_2$ anion radical from adsorbed gas-phase carbon dioxide that subsequently reacts to form an oxalate species. The process is completely reversible, and thus allows the elementary steps involved to be studied at the atomic level. The oxalate species at the MIM interface have been identified locally by scanning tunneling microscopy, chemically by IR spectroscopy, and their formation verified by density functional calculations.

CO$_2$ is known to be one of the crucial greenhouse gases.[1] For decades, there have been efforts to store and utilize CO$_2$ in chemical reactions to transform a stable molecule into a useful chemical.[2] The crucial aspect is the transfer of an electron to the molecule,[3] which costs about 0.6 eV and is associated with a bending of the linear neutral CO$_2$ molecule.[4] This process may be understood on the basis of the so-called Walsh diagram[5] and it has been extensively discussed.[6] The free CO$_2$ radical anion is metastable with respect to electron detachment and has a lifetime of a few milliseconds.[6,7] By attaching a neutral CO$_2$ molecule to the radical anion, forming a (CO$_2$)$_2$ species, a thermodynamically stable entity is formed in the gas-phase of a molecular beam experiment.[8] By transferring a second electron, the dimer anion may be transformed into an oxalate species whereby a carbon–carbon bond is formed. This compound may then be further transformed with water or ammonia to useful chemicals. It is therefore the key issue to transfer electrons. If one looks for a catalytic transformation to drive such a reaction, a source of electrons is needed that shuttles electrons back and forth between the reacting species. The goal herein is to describe and characterize a system that fulfills these requirements.

The concept presented herein is based on a metal–insulator–metal (MIM) system consisting of Au islands as an electron-storage material supported on a substrate that provides electrons to be shuttled back and forth between Au islands and adsorbed CO$_2$ (for conceptual overviews, see Ref. [9]). Specifically, the Au islands are located on an ultrathin MgO film, which covers a metallic substrate. Theoretical and experimental evidence has been provided of the negative charge of such Au islands, which assume a specific flat, raft-like morphology.[10,11] We have demonstrated that molecules such as CO$_2$[12] and isophorone,[13] reside on the rim of such islands where the charge is localized for it to minimize electron–electron repulsion.[14] We will show in this study that the Au islands transfer electrons to carbon dioxide leading to the formation of a CO$_2^-$ radical ion, which may further react to form the oxalate. This model system may provide a playground to build upon a real catalytic system.

Figure 1 shows STM images of two Au islands on a bilayer MgO(001)/Ag(001) film. All experimental details explaining the deposition of the Au particles on ultrathin MgO are described in the Supporting Information. The morphology of the islands is, as proven before, two-dimensional with monolayer (ML) thickness.[11,15] Figure 1 a shows a pristine cluster, which holds an average of 0.2 electrons per interface atoms that originate from the underlying metal substrate.[16] The additional electrons localize preferentially at the cluster edge and thus maximize the local density of the states at the rim (Figure 1 c) resulting in the cluster edge to be the primary site for chemical reactions. This assumption is readily confirmed upon exposing the gold islands to CO$_2$ (Figure 1 b,c). Molecular adsorbates become visible only at scanning values between −0.5 and +0.5 V. The CO$_2$ derivatives are exclusively adsorbed at the rim of the Au islands (Figure 1 b), and

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carbonate, that is, CO$_2$ adsorbed on top of lattice oxygen, as well as with the formation of a carboxylate, where the molecule would reside on an oxygen vacancy or on a metal ion.\cite{18} The carbonate formation may be ruled out by measuring IR spectra after dosing CO$_2$ on MgO films prepared with isotope-labeled oxygen ($^{16}$O$_2$). The observed band does not shift to lower frequency, as would be expected for carbonate formation, because one of the masses of the carbonate would be an $^{16}$O atom. If, on the other hand, the CO$_2$ is either oxygen- or carbon-labeled, the frequency of the carboxylate shifts as expected to 1259 cm$^{-1}$ and to 1275 cm$^{-1}$, respectively (Figure 2a). The frequency as well as an adsorption site possibly on top of a F$^-$ center is also consistent with calculations from various groups for CO$_2$ adsorption on bulk MgO(100) surfaces.\cite{19} A full study of CO$_2$ adsorption on a thin MgO(100) film supported on Ag(001) has not been published to date. The results reported for a bulk MgO(001) may not be directly applicable here owing to the absence of the metal support.

If CO$_2$ is dosed onto Au islands supported on MgO, new vibrational bands are observed. The coverage of Au was 0.06 ML. The IR spectra of this system at CO$_2$ saturation exposure are shown in Figure 2b (top). Three bands are observed; the band at 1294 cm$^{-1}$ has been identified above and may be used as a calibration standard. The other two bands centered at 1220 cm$^{-1}$ and 1398 cm$^{-1}$ are assigned to oxide. There is certain degree of heterogeneity in those bands owing to the irregular rim shape of the Au islands. The amount of adsorbed molecules as calculated from the region of the MgO surface covered by Au islands of 4–5 nm diameter (average nanoparticle size; Supporting Information, Figure S1) agrees well with the observed intensities assuming 15 molecules per island adsorbed. The basis for assigning the molecules to oxide species comes from a comparison with known carbonate transition metal complexes\cite{20} which exhibit two bands with appropriate vibrational frequencies.
Further evidence is again derived from isotopic labeling experiments. First, when labeling the oxide film (Mg\textsuperscript{18}O), we observed no shifts of any of the bands, thus ruling out the possibility that oxygen atoms at the cluster rim are responsible for anchoring the CO\textsubscript{2}. As evident from Figure 2b, the band shift upon CO\textsubscript{2} labeling is consistent with the assignment based on metal oxalate complexes. There are also unavoidable bands that are due to unlabeled CO\textsubscript{2} from the background. In particular, for \textsuperscript{18}O-labeled CO\textsubscript{2}, the carboxylate at MgO sites shifts to 1258 cm\textsuperscript{-1} as before, and the other two bands shift to 1374 cm\textsuperscript{-1} and 1210 cm\textsuperscript{-1}, respectively. Upon \textsuperscript{13}C-labeling, the carboxylate shifts to 1274 cm\textsuperscript{-1} while the band at 1220 cm\textsuperscript{-1} does not shift, and the higher-frequency band moves to 1360 cm\textsuperscript{-1}. For the two oxalate modes, a normal mode analysis is available, performed for a planar oxalate arrangement, and the in-plane vibrational frequencies have been calculated.\textsuperscript{208} Those calculations reveal that the 1220 cm\textsuperscript{-1} band has predominantly C–O stretch and O-C-O angle deformation character, while the 1398 cm\textsuperscript{-1} band has predominantly C=C stretch character with some C-O stretch contributions mixed in. The observed isotopic shifts are fully consistent with those qualitative assignments and support strongly the assignment of the two observed peaks to oxalate species formed at the rim of the Au islands.

Several representative adsorption configurations of CO\textsubscript{2} and C\textsubscript{2}O\textsubscript{4}\textsuperscript{2−} species at an Au ad-atom and an Au\textsubscript{2} ad-dimer attached to a MgO(001)/Ag(001) film were investigated by DFT calculations (Figure 3; Table S1). Two configurations for an AuCO\textsubscript{2} ad-complex were found, which are a surface-planar and surface-normal CO\textsubscript{2} species bound at the side of the Au atom (Figure 3a,b). In the surface-planar configuration, the Au ad-atom transfers −0.7 e\textsuperscript{-} to CO\textsubscript{2}, resulting in an adsorption energy of −0.58 eV. In the surface-normal configuration, the charge transfer is slightly larger but the adsorption is weaker than in the case of the clean film. Similar two configurations are found for CO\textsubscript{2} adsorption on the Au\textsubscript{2} ad-dimer. There, both the CO\textsubscript{2} binding energy and the amount of charge transfer are clearly increased suggesting an even stronger CO\textsubscript{2} binding for larger Au ad-clusters (Table S1).

We turn next to calculations on the oxalate C\textsubscript{2}O\textsubscript{4}\textsuperscript{2−} species. We found two gas-phase configurations ("cross" and planar) with C–C bond lengths of 1.61 and 1.70 Å, respectively (Figure 3c,d). These species were converged to a stable minimum with a total charge of −1.8 e\textsuperscript{-} in the complex, approximating closely the full double anion. When this complex is bound to the Au\textsubscript{2} ad-cluster, again two configurations were found, with adsorption energies of −0.95 eV for the "C–C surface-planar" and −0.72 eV for the "C–C surface-normal" (Figure 3e,f and 3g,h). The C–C bond length is reduced here to 1.56 Å and 1.53 Å, respectively, and the charge on the C\textsubscript{2}O\textsubscript{4}\textsuperscript{2−} part is −1.59 e\textsuperscript{-} and −1.55 e\textsuperscript{-} (Table S1) with the gold atom next to the complex being clearly positively charged (Figure 3e,g). Our calculations suggest that the formation of the oxalate is an activated process, as the formation of the stable gold-oxalate complex seems to depend sensitively on the reaction coordinate.

Vibrational analysis was performed at the Au\textsubscript{2}-oxalate species as well (Table S1). We note that the "C–C surface-normal" configuration (Figure 3h.i) has eigenmodes at 1228 cm\textsuperscript{-1} and 1414 cm\textsuperscript{-1}, which are not far from the observed frequencies at 1220 cm\textsuperscript{-1} and 1398 cm\textsuperscript{-1} in Figure 2b.

We have also performed temperature-programmed desorption experiments to identify desorbing species. Only CO\textsubscript{2} is detected desorbing from the surface and representative TPD spectra are shown in Figure S2. Two desorption states have been identified: The carboxylate species desorbs between 280 and 310 K, while the oxalate species has desorbed at 340 K according to Equation (1):

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**Figure 3.** Computed structures of CO\textsubscript{2} and C\textsubscript{2}O\textsubscript{4}\textsuperscript{2−} species. a,b) two configurations of CO\textsubscript{2} bound at a single Au ad-atom; c,d) "cross" and planar structures of gas-phase C\textsubscript{2}O\textsubscript{4}\textsuperscript{2−}; e,f) side and top views of a "cross" C\textsubscript{2}O\textsubscript{4}\textsuperscript{2−} bound at the Au ad-dimer having a surface-planar C–C bond; g,h) side and top views of the "cross" C\textsubscript{2}O\textsubscript{4}\textsuperscript{2−} bound at the Au ad-dimer having a surface-normal C–C bond. Colors: Mg green, O red, C brown, Au yellow, Ag gray. The blue italics denote atomic Bader charges and black roman numerals interatomic bond lengths and bond angles. The gas-phase structures in (c) and (d) were converged for the total charge of −1.8 e\textsuperscript{-} (see text).
(C₂O₄)²⁻ → 2CO₂ + 2e

(1)

This is by no means self-evident, as in former studies on adsorbed oxalates, an alternate route of decomposition has been observed, as given in Equation (2):[2]

(C₂O₄)²⁻ + CO → 2CO₂ + O²⁻

(2)

In this scenario, CO adsorption should be observable and oxygen would remain on the surface limiting the reversibility of the process. In the present case, CO₂ adsorption-desorption experiments reveal that the process is entirely reversible. One reason for the reversible oxalate-carbon dioxide adsorption-desorption is the low oxophilicity of Au, which hampers the formation of adsorbed carbonate and thus the alternate reaction route. This reversibility renders our system a well-suited model system to study electron shuttling-induced reactions at surfaces.

Our experimental and computational results presented above strongly suggest the following scenario at the supported Au islands on a ultrathin MgO(001) film: The ultrathin MgO(001) film grown on a Ag(001) surface shuttles electrons to nanoscopic Au islands, which assume a flat, raft-like morphology with the transferred electrons located at the rim of the raft. Carbon dioxide adsorbs at the rim of the nano islands, transforms into carboxylate by electron transfer, which then reacts with another CO₂ molecule to form oxalate according to Equation (3):

\[
\begin{align*}
\text{CO}_2 + e^- & \rightarrow \text{CO}_2^- / \text{Au(nano)} \\
\text{CO}_2^- / \text{Au(nano)} + \text{CO}_2 & \rightarrow (\text{CO}_2^-)_2 / \text{Au(nano)} \\
(\text{CO}_2^-)_2 / \text{Au(nano)} + e^- & \rightarrow \text{C}_2\text{O}_4^{2-} / \text{Au(nano)} \\
\text{C}_2\text{O}_4^{2-} / \text{Au(nano)} & \rightarrow 2\text{CO}_2 + 2e^{-}
\end{align*}
\]

(3)

The formed oxalate desorbs from the surface as carbon dioxide and the electrons are shuttled back to the Au islands. The reaction is fully reversible. Evidence for the oxalate C₂O₄²⁻ intermediate is found from the IRAS data and supported by DFT calculations on oxalate adsorption on supported Au, cluster on the MgO film. Further experimental and computational investigations will be conducted to fully characterize the intermediates and to reveal activation barriers of the above-mentioned reaction at the gold island edges.

Keywords: carbon dioxide · electron transfer · metal-insulator-metal structure · oxalate · oxygen


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