The Electrophilicity of Surface Carbon Species in the Redox Reactions of CuO-CeO₂ Catalysts

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Electronic metal–support interactions (EMSI) describe the electron flow between metal sites and a metal oxide support. It is generally used to follow the mechanism of redox reactions. In this study of CuO–CeO$_2$ redox, an additional flow of electrons from metallic Cu to surface carbon species is observed via a combination of operando X-ray absorption spectroscopy, synchrotron X-ray powder diffraction, near ambient pressure near edge X-ray absorption fine structure spectroscopy, and diffuse reflectance infrared Fourier transform spectroscopy. An electronic metal–support–carbon interaction (EMSCI) is proposed to explain the reaction pathway of CO oxidation. The EMSCI provides a complete picture of the mass and electron flow, which will help predict and improve the catalytic performance in the selective activation of CO$_2$, carbonate, or carbonyl species in C1 chemistry.

Introduction

The interaction between metals and supports plays a vital role in modulating the catalytic performance of active sites. Concepts such as SMSI$^{[3]}$ and EMSI$^{[2]}$ are well-established to describe the geometric$^{[3]}$ electronic$^{[26-4]}$ and bifunctional$^{[3]}$ modification of active sites from the support. With respect to the electronic interaction, there is a flow of electrons either from a metallic centre to an oxidative support or from a reductive support to an oxidative metal centre. The net electron flow leads to modified electronic structures of the active centre and its surroundings, and thus to different behaviours in the adsorption and activation of reaction molecules compared to its unperturbed state. In addition to the oxide support, the destination of the electron flow can also be the surface carbon species resulting from the decomposition and deposition from carbonaceous reactants. Carbon atoms, with nine oxidation states and an electronegativity of 2.55, can accept electrons from metals. Carbon materials are widely used as electron acceptors in lithium-ion batteries$^{[6]}$ and organic photovoltaics.$^{[7]}$ There are some representative studies in catalysis find electrons are transferred from metal to carbon$^{[9]}$, especially in electrocatalysis.$^{[9]}$ Reduction of carbon can form metal-carbon species in Fischer–Tropsch synthesis$^{[10]}$ and alkenes hydrogenation.$^{[11]}$ A CO$_2$ species is formed when the 2π, orbital of surface CO accepts electrons from metallic Cu.$^{[12]}$ Theoretical calculations also predict the modification of the Cu electronic structure via surface carbons.$^{[13]}$

Cu has moderate adsorption strength for carbonaceous intermediates,$^{[14]}$ and is widely used for C1 chemistry, including CO oxidation,$^{[15]}$ water-gas shift,$^{[16]}$ steam reforming,$^{[17]}$ methanol synthesis,$^{[18]}$ and electrochemical reduction of CO$_2$.$^{[14,19]}$ The well-documented Ce$^{4+}$/Ce$^{3+}$ redox pair enables the transfer of oxygen atoms and electrons between Cu$^{2+}$/Cu$^{+}$ active sites and the CeO$_2$ support.$^{[20]}$ Validation of this catalytic cycle still requires precise quantification of their oxidation states to match the balance of electron transfer.$^{[21]}$

Here we report the full picture of electron transfer in this system by considering the electrophilicity of surface carbon species, which are in situ deposited from CO molecules. At 453 K, electrons are initially enriched on metallic Cu via CO reduction (CO stage) to built up the chemical potential for an electron flow within the catalyst. The electrophilicity of surface carbon species is then studied in an inert atmosphere (He stage) to exclude electron transfer from or to the gaseous molecules. Finally, O$_2$ is used to extract electrons that are originally injected into the catalyst from CO (O$_2$ stage). The CO oxidation activity of the catalyst at individual stage is compared below 353 K to elucidate the impact of initial oxidation states of Cu and Ce as well as the surface carbon species.

In a full cycle of CO oxidation, electrons are transferred from CO to Cu, then to Ce and carbon (Figure 1) and finally to O$_2$. We therefore extend the concept of EMSI to “electronic metal-support-carbon interactions” (EMSCI) in...
order to address this interplay among Cu, Ce and carbon species.

**Results and Discussion**

**Highly Dispersed CuO Clusters on CeO$_2$**

Fine powder of highly dispersed 20 wt% CuO clusters on CeO$_2$ with 71 m$^2$g$^{-1}$ specific surface area were prepared via flame spray pyrolysis.[20b,22] The high-resolution aberration-corrected high angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) images show crystalline CeO$_2$ particles with an average size of around 6 nm (Figure S1f,g). The corresponding energy-dispersive X-ray spectroscopy (EDS) maps identify the presence of Cu in the form of ca. 5 nm particles surrounded by CeO$_2$ (Figure S1a-e). The X-ray photoelectron spectroscopy (XPS) shows that ca. 18% of the Ce is Ce$^{3+}$, indicating a considerable number of oxygen vacancies in the lattice (Figure S2 and Table S1).[23] The surface Cu/(Cu + Ce) ratio determined by XPS is 0.36, which is close to the theoretical bulk Cu/(Cu + Ce) ratio (0.35) for 20 wt% CuO-CeO$_2$. This indicates good dispersion of the Cu species over CeO$_2$, leading to large Cu/Ce interface. In comparison, Cu species are usually enriched on CeO$_2$ surface using conventional synthesis methods, showing much higher surface Cu content compared to the bulk composition (Table S2).[24] Laboratory X-ray diffraction (XRD) pattern shows broadened CeO$_2$ diffraction peaks (Figure S3a), whereas weak CuO diffraction peaks can be recognised by synchrotron X-ray powder diffraction (SXPD; Figure S3b). Therefore, the initial catalyst structure of 5 nm CuO and 6 nm CeO$_2$ is confirmed with 18% Ce$^{3+}$ present.

This 20 wt% CuO-CeO$_2$ catalyst has been applied for the preferential CO oxidation in the presence of 50% H$_2$, achieving a wide temperature window from 377 to 388 K with 99% conversion and selectivity of CO oxidation.[19b] Our previous in situ study used a sequence of CO/N$_2$/O$_2$ flows at 453 K to probe the change of Cu oxidation state and the corresponding gas profile.[21b] Two interesting phenomena were observed: 1) under inert N$_2$, Cu$^0$ was slowly reoxidised; 2) when N$_2$ was changed to O$_2$, CO$_2$ was released without an external CO feed. In addition, the near ambient pressure-near-edge X-ray absorption fine structure (NAP-NEXAFS) spectra of a similar system reveals the conversion from oxidised carbon species (288.3 eV) to reduced carbon species (284.9 eV) in ultra-high vacuum (Figure S4). We hypothesise that carbon species will deposit on the surface of CuO-CeO$_2$ from gas-phase CO. Such carbon species are reduced under inert conditions, whereas Cu$^0$ is oxidised simultaneously. When O$_2$ is introduced, the carbon species are oxidised to CO$_2$. This hypothesis of EMSCI is verified via a combination of operando X-ray absorption fine structure (XAFS), SXPD, NAP-NEXAFS, diffuse reflectance infrared Fourier transform spectroscopy (DRIFTS) and gas component quantification at the CO, He and O$_2$ stages, as presented below.

**CO Stage: Electron Transfer from CO to Cu**

Introducing CO at 453 K to the catalyst reduces Cu$^{2+}$ and Ce$^{4+}$ (Figure 2). Most of the Cu reduction follows the sequence of Cu$^{2+}$—Cu$^{3+}$—Cu$^0$, as indicated in the Cu K-edge X-ray absorption near edge structure (XANES) and SXPD (Figure 2a,b). The Ce$^{3+}$ content increases from 20% to 24% according to the peak fitting results of the Ce L$_3$-edge XANES (Figure 2c, note that the initial 20% Ce$^{3+}$ content is slightly different from the 18% Ce$^{3+}$ obtained from XPS study).

DRIFTS shows the absorption peaks of Cu’-carbonyl species (Cu’-CO) in the range of 2160-2080 cm$^{-1}$ suggesting the presence of Cu’ on the surface of metallic Cu (Figure 2d). NAP-NEXAFS at the Cu L$_3$-edge validates that Cu’ dominates the surface layer after the reduction of CuO (Figure S10). Such surface Cu’ species on Cu$^0$ clusters over CeO$_2$ have also been reported in the water gas shift reaction.[19] In general, the surface CO absorption band undergoes a slight blue shift from 2098 to 2111 cm$^{-1}$, and then a red shift to 2100 cm$^{-1}$ with a significant intensity decrease. The initial blue shift indicates the increase of CO coverage[25] on Cu’ along with the release of CO$_2$ detected at 2364 and 2337 cm$^{-1}$. A new shoulder peak at 2138 cm$^{-1}$ appears when the CO coverage is maximised. The peak is similar to the reported band at 2135 cm$^{-1}$ of CO on partially reduced CuO.[25a] The increase of a broad feature between 1700-1200 cm$^{-1}$ (Figure S11 from blue to dark red) that corresponds to surface carbonate species is observed.[18b,26] The formation of the carbonate stems from the oxidation of surface carbonyl species, a process that is reported in the literature.[18b] These carbonates can increase the electrophilicity of neighbour Cu’,[25a] leading to a broad absorption band of Cu’-CO at 2173 cm$^{-1}$ when the reduction of Cu is complete as reflected by the disappearance of CO$_2$ peaks (Figure 2d). The second stage red-shift to the steady and broadened band centred at 2100 cm$^{-1}$ corresponds to a combination of Cu’-CO and Cu$^2$-CO bands at 2160-2080 cm$^{-1}$ and 2090-
2060 cm\(^{-1}\), respectively.\(^\text{[25]}\) A slight decline in surface carbonate species is observed (Figure S11 from dark red to black).

The CO reduction of 20 wt% CuO-CeO\(_2\) releases 2.99 mmol g\(^{-1}\) of CO\(_2\) at 453 K (Figure 2e). Theoretically, the complete reduction from Cu\(^{2+}\) to Cu\(^0\) will generate 2.52 mmol g\(^{-1}\) of CO\(_2\). Furthermore, there is a 4\% reduction of Ce\(^{4+}\) to Ce\(^{3+}\) (according to XANES fitting) results in 0.09 mmol g\(^{-1}\) of CO\(_2\) formation. These results indicate an additional 0.38 mmol g\(^{-1}\) of CO\(_2\) generation that is not related to the Cu and Ce redox. According to the literature, this additional CO\(_2\) may come from the disproportionation of CO\(_2\).\(^\text{[27]}\) A very recent work on CO\(_2\) methanation also finds that the presence of Ce\(^{3+}\) can also help dissociate CO and CO\(_2\), leading to diverse carbon species.\(^\text{[28]}\) Carbonyls and carbonates are detected by DRIFTS, suggesting the high complexity of those surface carbon species. (Figure 2d and Figure S11).

The combination of operando XAFS of the Cu and Ce oxidation states, SXPD of small crystalline clusters and DRIFTS of surface carbon species reveals the change of Cu, Ce and C during the CO stage (Figure 2f). The majority of CuO is fast reduced to metallic Cu with a trace amount of surface Cu\(^+\), whereas 4\% Ce\(^{4+}\) is reduced to Ce\(^{3+}\). Surface carbon species (i.e., carbonyls and carbonates) are deposited from gaseous CO.

**He Stage: Electron Transfer from Cu\(^+\) to Ce\(^{4+}\) and Surface Carbon**

The inert He flow enables observation of the internal reaction among Cu\(^\circ\), Ce\(^{3+}\)/Ce\(^{4+}\) and surface carbon species. Cu\(^\circ\) clusters are immediately oxidised to Cu\(^{+}\) once CO is replaced by He (Figure 3a,b). Simultaneously, 9\% Ce\(^{4+}\) is reduced to Ce\(^{3+}\) (Figure 3c). These results suggest a fast redox reaction [Eq. (1)].

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\text{Cu}^\circ + \text{Ce}^{4+} \rightarrow \text{Cu}^+ + \text{Ce}^{3+} 
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To the best of our knowledge, the observation of Cu oxidation and Ce reduction here is the first direct evidence for this reaction, proving the EMSI between Cu and Ce. The Cu\(^+\) state is stable in the He flow (Figure 3a), whereas the crystalline Cu\(_2\)O feature gradually decreases with time at 453 K, suggesting the amorphization of Cu\(_2\)O (Figure 3b).
The oxidation of all Cu⁰ to Cu⁺ would require 54 % Ce⁴⁺ to be reduced to Ce³⁺, which is much higher than the 9% formation of Ce³⁺. Within the system, the only destination to receive the additional electrons from Cu⁰ are surface carbon species with their abundant valence states, given that both carbonyl species and carbonate species are present on Cu⁺-CeO₂ at the He stage (Figure 3d, Figure S15). The blue shift from 2100 cm⁻¹ to 2127 cm⁻¹ suggests the increased electrophilicity of Cu due to its oxidation. The surface carbon species as electron acceptors have also been reported for CO₂ activation on metallic Cu, where CO₂⁺ species were formed via electron transfer from Cu⁰ into the 2π orbital of surface CO₂.[12] Such CO₂⁺ species were identified by surface enhanced Raman spectroscopy[29] and XPS.[30] In addition, the decomposition of these carbon species can generate surface O atoms to form Cu–O bond.[26,31]

The flow of electrons from Cu⁰ to Ce³⁺ and surface carbon species under inert atmosphere proves the EMSCI concept (Figure 3f). Temperature programmed desorption (TPD) is applied to support the presence of those surface carbon species at 453 K in addition to the DRIFTS evidence. To obtain a similar metallic Cu and CeO₂ surface, the catalyst is reduced first with CO at 453 K and then cooled to 298 K under the CO atmosphere. Subsequent heating in He leads to CO₂ desorption. The released CO₂ may come from the decomposition of surface carbonates[26] or the desorption of CO₂ formed during CO reduction.[31] 0.20 mmol g⁻¹ and 0.39 mmol g⁻¹ of CO₂ release are observed below and above 453 K, respectively (Figure 3e). The latter is stable under He in the in situ study and is partially responsible for the oxidation of Cu (Figure 3a,b). The remaining surface carbon species that cannot be desorbed by heating in He may also accept electrons and oxidise Cu. Density functional theory (DFT) calculations with Bader charge analysis are performed to evaluate the electron transfer between Cu⁰ and adsorbed carbon species. Each Cu atom that adsorbs CO has extra +0.11e and +0.12e at low and high CO coverages, respectively whereas each CO molecule gains −0.13e from Cu (Figure S16, S17 and Table S4). When the coverage of carbon species is further increased, the adsorbed carbon atoms, the number of which equals to half of the Cu atoms, gain −16.58e from Cu, resulting in +0.20e to +0.70e on individual Cu atom
(Figure S18). The theoretical calculations results validate the electrophilicity of surface carbon species and its impact on reducing the electron density of Cu. An O2 stage is then carried out to study those residual carbon and the further oxidation of Cu+//Ce4+.

O2 Stage: Electron Transfer from Ce4+, Surface Carbon, and Cu+ to O2

When switching to O2 at 453 K, the Ce3+ content recovers to the initial level, whereas only a slight increase of Cu2+ is found (Figure 4a,c). No significant change is found in SXPD, suggesting Cu+ remains amorphous (Figure 4b). While the majority of Cu remains in Cu+ state, a full reoxidation to Cu2+ is found on the surface as indicated in the NAP-NEXAFS (Figure S22). A possible explanation is that a dense layer of CuO is formed over the Cu2O surface, preventing further oxidation of Cu+. We hypothesise that when the size of the Cu2O decreases, the curvature of the surface CuO layer increases and more Cu can be oxidised into Cu2+. An extreme case is the atomic Cu sites, which can be completely oxidised back to Cu2+ even under He flow.[22]

The residual surface carbonyl species with a band at 2137 cm−1 are removed via oxidation with simultaneously released CO2 (Figure 4d,f), whereas the carbonates only slightly decrease under O2 at 453 K (Figure S23). In addition to the 0.59 mmol g−1 CO2 released during TPD in He up to 673 K (Figure 3e), temperature programmed oxidation (TPO) of the residual surface carbon can further generate 0.28 mmol g−1 CO2 (Figure 4e). Therefore, at least 0.87 mmol g−1 of carbon species can be deposited and 0.67 mmol g−1 of them can contribute to the oxidation of Cu0 via EMSCI. The amount of carbon deposited on the surface may be directly proportional to the CO pressure (P CO). The different P CO in each experiment may lead to small inconsistencies in the carbon deposition as well as the time required for achieving steady states (Table S5). Nevertheless, the general picture on the direction of the electron flow via EMSCI is valid.

Figure 4. Operando characterisation of Cu, Ce and surface carbon during the oxidation of Cu+//CeO2 at 453 K with O2. a) Contour map of the Cu K-edge first derivative XANES spectra, showing the incomplete oxidation of Cu+ to Cu2+. XANES spectra are shown in Figure S19. b) Contour map of the SXPD patterns, showing the crystalline Cu2O preserved during oxidation. SXPD patterns are shown in Figure S20. c) Contour map of the Ce L3 edge first derivative XANES spectra, showing the recovery of Ce3+ content after oxidation. XANES spectra are shown in Figure S21 and their fitting results are given in Table S3. d) DRIFTS spectra of the surface carbonyl species as they are oxidised and released as CO2. The initial spectrum obtained in He at 453 K is labelled in purple and the progress in O2 at 453 K is colour coded from light blue to blue. e) Exhaust gas profile shows the released CO2 during TPO from 300 K to 673 K, then holding at 673 K for 30 min indicated by the pink-shaded region. f) Simplified models of the structural evolution from Cu+//CeO2 to Cu2+/Cu4+//CeO2.
EMSCI describes the flow of electrons in the sequence of 1) CO; 2) Cu²⁺; 3) carbon + Ce³⁺; 4) O₂. At these four stages, the Cu oxidation states are Cu²⁺, Cu³⁺, Cu⁴⁺, and Cu⁵⁺/Cu²⁺, respectively. The CO oxidation kinetics is then studied at these four stages (Figure S24) to understand the influence of surface carbon and Cu species in catalysis.

The reaction temperature is controlled below 353 K to preserve the initial states of the catalysts. The distinct CO conversion profiles (Figure 5a) indicate the different catalytic behaviours at each stage. In general, the CO oxidation activity of the initial Cu species on CeO₂ follows the order Cu²⁺ > Cu³⁺ > Cu⁴⁺. The fully oxidised CuO-CeO₂ shows the highest turnover frequency (TOF) at 323 K and lowest apparent activation energy (Eₐ) (Figure 5b,c blue) whereas the Cu³⁺-CeO₂ shows the lowest activity and increased Eₐ (Figure 5b,c red). The Cu⁴⁺-CeO₂ catalysts separately reduced by CO and H₂ show similar TOF and Eₐ in the kinetic region (Figure 5b,c red and wine). However, the CO-reduced catalyst with considerable amount of surface carbon shows much higher activity in the high conversion region compared to the Cu₂⁺-reduced catalyst which is free of carbon (Figure 5a red and wine). It suggests that surface carbon on Cu⁶⁺ species is irrelevant to the activity in the kinetic region but can significantly promote the reaction in the high conversion region. After the CO-reduced Cu⁴⁺-CeO₂ is annealed in He, the obtained Cu²⁺ with reduced surface carbon shows increased TOF (Figure 5). The obtained Cu²⁺ cannot be fully oxidised to the initial Cu³⁺ in 20% O₂/He at 473 K, thus the activity cannot fully recover although its Eₐ significantly decreases (Figure 5a,c, red and wine). The Ce³⁺ content is generally accepted to be inversely proportional to thermodynamic oxygen vacancy formation energy (Eᵥ[32]) which is regarded as a descriptor for CO oxidation activity.[32] Unfortunately, there is no clear relationship between the Ce³⁺ content and CO oxidation performance can be correlated in this work. It suggests that influence of Ce³⁺ content towards CO oxidation is not as obvious as that of the Cu oxidation state.

In conclusion, the lower oxidation state of Cu species on CeO₂ leads to inferior CO oxidation activity. Surface carbon is irrelevant to the activity in the kinetic region but can significantly promote the reaction in the high conversion region.

**Conclusion**

The concept of EMSCI is established to describe the electron flow for the CuO-CeO₂ system during CO oxidation. More specifically, the electrons flow from CO to Cu, then to Ce and surface carbon species, and finally to O₂. The EMSCI concept sheds light on the catalytic cycle of other CO and CO₂ involved reactions promoted by polyvalent metal oxides. The probability of transition metals transferring the d electrons to the π* orbitals of surface carbon species will govern how CO₂ or CO can be activated over the catalysts.

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**Figure 5.** The catalytic performance of 20 wt% CuO-CeO₂ in CO oxidation after different pretreatments. CO oxidation conditions: Weight hourly space velocity (WHSV) per gram of catalyst: 1500 mL·g⁻¹·h⁻¹. 80 mg 20 wt% CuO-CeO₂, 200 mL·min⁻¹ flow, 1% CO, 10% O₂. He balanced.

a) Conversion of CO as a function of temperature and catalysed by the catalysts with different Cu oxidation states: Cu²⁺ (Stage 1, blue solid squares); Cu³⁺ (Stage 2, red circles); Cu⁴⁺ (Stage 3, dark yellow stars); Cu⁵⁺/Cu²⁺ (Stage 4, navy open squares); H₂-reduced Cu²⁺ (dark red triangles). b) CO TOF at 323 K. c) Arrhenius plots in the range of 313–328 K. The calculated Eₐ values (kJ mol⁻¹) are listed in parentheses.