Studying the cation dependence of CO\textsubscript{2} reduction intermediates at Cu by in-situ VSFG spectroscopy

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Abstract

The nature of the electrolyte cation is known to have a significant impact on electrochemical reduction of CO\textsubscript{2} at catalyst|electrolyte interfaces. An understanding of the underlying mechanism responsible for catalytic enhancement as the alkali metal cation group is descended is key to guide catalyst development. Here, we use in-situ vibrational sum frequency generation (VSFG) spectroscopy to monitor changes in the binding modes of the CO intermediate at the electrochemical interface of a polycrystalline Cu electrode during CO\textsubscript{2} reduction as the electrolyte cation is varied. A CO\textsubscript{bridge} mode is observed only when using Cs\textsuperscript{+}, a cation that is known to facilitate CO\textsubscript{2} reduction on Cu, supporting the proposed involvement of CO\textsubscript{bridge} sites in CO coupling mechanisms during CO\textsubscript{2} reduction. Ex-situ measurements show that the cation dependent CO\textsubscript{bridge} modes correlate with morphological changes of the Cu surface.

Introduction

Anthropogenic climate change has led to the urgent requirement for sustainable carbon cycles to be developed. The use of CO\textsubscript{2} through capture and conversion technologies has been identified as a promising way to generate chemical feedstocks and fuels.\textsuperscript{1} Electrochemical CO\textsubscript{2} reduction (eCO\textsubscript{2}R) has been studied extensively at a range of metal surfaces; however there is still an incomplete mechanistic understanding of the eCO\textsubscript{2}R reaction.\textsuperscript{2} Cu has been identified as the only heterogeneous, monometallic catalyst to be able to reduce CO\textsubscript{2} to products that require more than two electrons in the reaction pathway in appreciable quantities, including many short-chain oxygenates and hydrocarbons (C\textsubscript{2+} products).\textsuperscript{3-5} This is hypothesised to be
due to optimal binding energies of crucial intermediates on Cu surfaces, shown by density functional theory (DFT) studies.\textsuperscript{6}

The nature of the catalyst is only one of the many factors which influence the performance of eCO\textsubscript{2}R; cell architecture and electrochemical environment both also play major roles. The species of the electrolyte has been shown to dramatically alter the product distribution of the reaction at Au,\textsuperscript{7–9} Ag\textsuperscript{10–12} and Cu.\textsuperscript{11–17} Specifically, the alkali metal cation species present in the electrolyte has a profound effect on the Faradaic efficiency (FE) towards target carbon products. When Cu is the electrode, the C\textsubscript{2}/C\textsubscript{1} product ratio is also reported to increase in the order Li\textsuperscript{+}< Na\textsuperscript{+}< K\textsuperscript{+}< Rb\textsuperscript{+}< Cs\textsuperscript{+}. The dependence of product distribution on cations has also been detected in zero-gap cathode structures, where cation cross-over from a liquid analyte through an anion exchange membrane occurred.\textsuperscript{18} The hypotheses for the mechanism of these catalytic enhancements are widely debated within the community; the three predominant arguments can be summarised as (i) the pK\textsubscript{a} of the water molecules in the hydration sphere of the cation at the cathode decreases as the alkali metal group is descended from Li\textsuperscript{+} to Cs\textsuperscript{+} increasing the available CO\textsubscript{2} by minimising conversion to bicarbonate locally.\textsuperscript{12,19} (ii) An increase in the potential drop across the outer Helmholtz plane (OHP), and hence local electric field, due to the smaller hydrated radius going from Li\textsuperscript{+} to Cs\textsuperscript{+}.\textsuperscript{20} The increased electric field can benefit eCO\textsubscript{2}R kinetics via a number of mechanisms, including reducing the activation barrier for charge transfer and through the stabilisation of reaction intermediates.\textsuperscript{20} Finally (iii) stabilisation of intermediates through specific cation/reaction intermediate interactions has been proposed, impacting on both CO\textsubscript{2} and CO binding strengths and the CO coupling reactions\textsuperscript{11,21–24} and supporting this conclusion is the recent demonstration that eCO\textsubscript{2}R does not occur in rigorously prepared solutions where the cation has been excluded.\textsuperscript{9}

Vibrational spectroscopies\textsuperscript{19,25–29} such as surface enhanced infrared absorption spectroscopy (SEIRAS), surface enhanced Raman spectroscopy (SERS) and shell isolated nanoparticle enhanced Raman spectroscopy (SHINERS) have been used extensively to study the electrode–electrolyte interface during eCO\textsubscript{2}R and to explore the role of cations. SEIRAS measurements have studied bicarbonate at an operating electrode providing evidence for the role of the cation in pH buffering during eCO\textsubscript{2}R.\textsuperscript{19} The potential dependent (often labelled vibrational Stark) tuning rates of vibrational probes (such as CO) at Au,\textsuperscript{28,29} Cu\textsuperscript{21} and Pt\textsuperscript{26} have also been reported and these can provide an indication of local field strength. Such studies show that the field strength correlates with an increased interfacial cation concentration and a decreased hydrated cation radius. However, it is important to note that in many cases only a single metal–CO
binding site is discussed. Furthermore a recent study of a Au electrode measured the field strengths arising at the electrode from the applied potential and found these to be relatively small when compared to the solvation induced Onsager reaction field, which is induced by the cation through direct interaction with surface CO.29

Although SEIRAS and SERS have provided important insights into the eCO₂R reaction, both techniques require surface modification of the electrode. This is typically roughening, or in some cases deposition of the catalysts onto a suitably roughened support, in order to gain sufficient enhancement factors to enable identification of surface species. A concern is that samples suitable for SEIRAS/SERS can show significantly different catalytic activities to the polycrystalline electrode surfaces that are routinely employed in eCO₂R and this can be correlated to differences in CO binding sites.25 CO has been identified linearly bound atop (COₐtₜₜₗ) and at bridging sites (COₘᵦᵢₜᵢₜₜ) by attenuated total reflection (ATR) FTIR and also SERS spectroscopies.30,31,40,32–39 Whilst COₘᵦᵢₜᵢₜₜ has been considered to be irreversibly bound and not active on the eCO₂R pathway,33 recent works have proposed that the presence of COₘᵦᵢₜᵢₜₜ leads to an increase in C₂ products,30,34,36,37 with DFT calculations suggesting that CO–CO coupling may be more facile between COₐtₜₜₜ and COₘᵦᵢₜᵢₜₜ.36 Interestingly, a preliminary ATR-FTIR study noted that COₘᵦᵢₜᵢₜₜ sites are present in Cs⁺ containing electrolytes21 which are proposed to promote C₂ formation and a similar observation was made by Raman spectroscopy.35 However, as highlighted in a recent review,41 there is a need for improved understanding of the nature of surface intermediates during eCO₂R at Cu and their dependence on the reaction conditions. Here we use VSFG spectroscopy to study ν(CO) at pc-Cu electrodes to identify relative populations of reaction intermediates and their evolution in the presence of different electrolyte salts. Three alkali metal cations have been chosen for analysis: K⁺, which is the most commonly used electrolyte cation for eCO₂R, Cs⁺, which has been shown to give the greatest enhancement for C₂⁺ products, and Na⁺, which shows poorer eCO₂R performance than K⁺ whilst maintaining appreciable levels of C-based products. The ability of VSFG to study catalyst|electrolyte interfaces without the need for modifications, as required in the aforementioned spectroelectrochemical studies, which can fundamentally alter the electrodes activity, makes it an important tool to assess the mechanisms occurring on the pc-Cu electrodes routinely employed for eCO₂R.
Results and Discussion

The data for the *in-situ* VSFG experiments on a pc-Cu electrode in 0.5 M Na\(^+\), K\(^+\) or Cs\(^+\) bicarbonate electrolytes purged with CO\(_2\) at a range of applied potentials is shown in Figure 1. Full experimental details are provided in the methods section and the experiment design is shown in Figure S1. The potential is held for 150 seconds with the VSFG spectrum collected for 120 s, following a 30 s equilibration period. The potential steps progress in the order \(-0.5 \rightarrow -1.0 \rightarrow -0.1 \rightarrow -0.5\) V vs. Ag\(^+\) in 0.1 V increments. Progressing the potential steps in this order ensures that the Cu surface is fully reduced at the start of the experiment. All potentials from this point onwards will be referenced against the Ag\(^+\) pseudo reference electrode and the choice of reference electrode is discussed in supplementary note 1. The potential window was chosen following linear sweep measurements of the same electrode which showed that the catalytic onset potential under CO\(_2\) was *ca.* \(-0.8\) V for all of the electrolytes (Figure S2). As discussed in supplementary note 2, VSFG spectra are recorded with a short time delay (*ca.* 0.7 ps) between the mIR pulse (centred at approximately 1920 cm\(^{-1}\), 170 fs duration) and a time-asymmetric nIR picosecond laser pulse in order to minimise the non-resonant SFG response that can otherwise dominate the VSFG spectrum.\(^{42}\) Spectra are also normalised to the intensity of the mIR profile as discussed in supplementary note 3.
Figure 1: VSFG spectra showing CO modes on pc-Cu across the studied potential window CO$_2$-purged 0.5 M a) NaHCO$_3$, b) KHCO$_3$ and c) CsHCO$_3$ during sequential potential measurements ($-0.5 \rightarrow -1.0 \rightarrow -0.1 \rightarrow -0.5$ V vs. Ag$^+$ in 0.1 V increments). In each panel the data for the reverse steps ($-1.0 \rightarrow -0.1$ V) is in blue to guide the eye. The red, green and yellow shaded regions highlight the spectral range where bands are assignable to $\nu$(CO$_{atop(defect)}$), $\nu$(CO$_{atop(terrace)}$) and $\nu$(CO$_{bridge}$) sites respectively.

The CO binding site on Cu is dependent on the cation type and applied potential: Figure 1 shows that $\nu$(CO) are detected by in-situ VSFG spectroscopy on the pc-Cu electrode during eCO$_2$R but the nature and presence of $\nu$(CO) is dependent upon the choice of electrolyte cation. In the presence of Na$^+$, a single $\nu$(CO) at approximately 2062 cm$^{-1}$ is present at $-0.5$ V. A second band at 2094 cm$^{-1}$ grows in at $-0.8$ V and follows to become the dominant band by $-0.9$ V. As discussed below the exact positions of the $\nu$(CO) depends on the applied potential. The presence of adsorbed CO at $-0.5$ V, positive of the anticipated onset potential for eCO$_2$R (Figure S2), is due to the production of CO during prior measurements on the cell (supplementary note 2). The 2062 and 2094 cm$^{-1}$ bands can be assigned to CO linearly bound at terraced (CO$_{atop(terrace)}$) and defect (CO$_{atop(defect)}$) sites, respectively.$^{25,27,43,44}$ VSFG band intensities ($I_{VSFG}$) recorded by homodyne detection scale quadratically with number density and are also dependent on the orientation of the vibrational mode and ordering at the surface.$^{45}$ Here we use ($I_{VSFG}$)$^{0.5}$ as a way to estimate the relative populations of the different CO binding modes, Figure 2, with $I_{VSFG}$ determined from the peak area obtained by Lorentzian peak fitting. As shown in Figure 2, when using NaHCO$_3$ CO$_{atop(terrace)}$ accumulates between $-0.5$ and $-0.8$ V, whilst CO accumulates at defect sites between $-0.7$ to $-0.9$ V. It is reported that CO$_2$ reduction to CO occurs preferentially at defect sites, formed following the reduction of surface oxides and the increase in population of CO$_{atop(defect)}$ at potentials around where eCO$_2$R onsets (ca. $-0.8$ V, Figure S2, 2) is in line with this proposal.$^{46}$ However, we note that defect formation on Cu is known to be sensitive to CO coverage and we cannot rule out that the potential dependence of CO$_{atop(defect)}$ is due to the potential dependent formation of the binding site or relative potential dependent stability of the binding sites, and this is addressed in the discussion.
Figure 2: Square root of the area of the different VSFG measured CO modes ($I_{VSFG}^{0.5}$) as an estimate of surface concentration on Cu as a function of applied potential and electrolyte salt (a) NaHCO$_3$, (b) CsHCO$_3$, during eCO$_2$R. Values of 0 counts indicates that the signal:noise for the peak was insufficient to enable convergence for the fit. Populations could not be estimated for the KHCO$_3$ system in the absence of $I_{VSFG}$ for this data set.

When CsHCO$_3$ is used as the electrolyte during eCO$_2$R the two VSFG bands assigned to CO$_{\text{atop(terrace)}}$ and CO$_{\text{atop(defect)}}$ at approximately 2066 and 2094 cm$^{-1}$ respectively are present, alongside an additional band at 1887 cm$^{-1}$ (at $-0.5$ V), Figure 1c. The frequency of the band at 1887 cm$^{-1}$ matches that previously assigned to CO$_{\text{bridge}}$ binding at Cu.$^{30,33,34,38,39,41,47}$ Past studies have proposed that bridge bound CO is electrochemically stable$^{30,33,38,47}$ and we also observe that the band at 1887 cm$^{-1}$ at $-0.5$ V is stable across a wide potential window ($-0.1$ to $-1.0$ V, Figure 2a and Figure S5) and that CO accumulates at this site during the experiment. The observation that CO$_{\text{bridge}}$ is present when Cs$^+$ is used as an electrolyte but not Na$^+$ is notable. When the VSFG experiment in CsHCO$_3$ is repeated under CO instead of CO$_2$, the CO$_{\text{bridge}}$ species is not observed, Figure 3. This demonstrates that the generation of and/or occupancy of the CO$_{\text{bridge}}$ site relies upon the specific local conditions generated during eCO$_2$R. Both the CO coverage and concentration in the electrolyte have also been shown to be important factors...
in the reconstruction of Cu surfaces,\textsuperscript{27,48} which may explain the evolution of different binding motifs when compared to the VSFG data during eCO\textsubscript{2}R.

Figure 1b shows that VSFG experiments in KHCO\textsubscript{3} have very low I\textsubscript{VSFG} bands only. Previously a lack of observation of \(\nu(\text{CO}_{\text{atop}})\) bands by VSFG on pc-Cu has been attributed to the presence of a disordered CO layer on the electrode.\textsuperscript{49} It is well established that CO can be measured at Cu electrodes in KHCO\textsubscript{3} by linear vibrational spectroscopies (e.g. SEIRAS, Raman).\textsuperscript{27,36,37,44,50–55} The lack of detected Cu-CO by VSFG during eCO\textsubscript{2}R using a KHCO\textsubscript{3} is therefore proposed to be due to disorder of the Cu-CO surface. Overall, it is clear that there is a large variation in the manner in which CO binds at the surface in the presence of each of the cations during eCO\textsubscript{2}R.

\textbf{Figure 3:} VSFG spectra showing CO modes on Cu across the studied potential window CO-purged 0.5 M CsHCO\textsubscript{3} at the potentials indicated. The red and green shaded regions highlight the spectral range where bands are assignable to \(\nu(\text{CO}_{\text{atop(terrace)}})\) and \(\nu(\text{CO}_{\text{atop(defect)}})\) sites respectively. A background subtraction using the first \(-0.5\) V spectrum (where no CO modes are observable) has been performed on all spectra to remove interference in the CO\textsubscript{atop} region by supercontinuum generation, shown in Figure S7 and discussed in supplementary note 3.
The CO potential dependent stretching frequency changes as the cation is changed: Variations in local electric field as the cation is changed have been proposed to be a significant factor in controlling eCO_2R activity and these may be a factor behind the potential and cation dependence of the CO binding site observed in Figure 2. Potential dependent ν(CO) frequency changes are often used as an indicator of local electric field strength and these are shown in Figure 4. The deviation from linearity in the potential dependent frequency tuning < −0.8 V (shaded blue in Figure 4) demonstrates that directly equating potential dependent frequency tuning to electric field effects is an oversimplification. Consequently, tuning rates here are determined from linear fits of the data between −0.1 and −0.8 V, where the frequency tuning is relatively linear, and only in cases in which there are 3 or more data points so that an error can be calculated. Tuning rates are annotated on the Figure 4 for the −0.5 to −0.8 V cathodic and the −0.8 to −0.1 anodic scans. A full set of all tuning rates calculated in CO_2 purged electrolytes can be found in Figure S8.
Figure 4: VSFG measured $v$(CO) on Cu as a function of applied potential with CO$_2$ purged (a) 0.5 M NaHCO$_3$ and (b) CsHCO$_3$ in the $-0.5 \rightarrow -1.0$ V (black) and $-1.0 \rightarrow -0.1$ V (red) scan directions. Tuning rates in the potential range $-0.1$ to $-0.8$ V are annotated and the region in which the frequency tuning rates become non-linear is shaded in blue.

The VSFG experiment in CO$_2$ purged CsHCO$_3$ shows a tuning rate of 24.3 ($\pm$ 3.4) cm$^{-1}$ V$^{-1}$ for the $v$(CO$_{\text{atop(terrace)}}$) site when the potential is scanned from $-0.5$ to $-0.8$ V. The $v$(CO$_{\text{bridge}}$) mode tunes at a higher rate of 74.4 ($\pm$ 13.6) cm$^{-1}$ V$^{-1}$. The difference in both absolute frequency and potential dependant frequency between the CO$_{\text{bridge}}$ and CO$_{\text{atop}}$ sites can be explained by the increase in backdonation to $\pi^*$ antibonding orbitals as the CO adsorbate coordinates to more metal atoms.$^{58}$ The tuning rate of the $v$(CO$_{\text{atop(terrace)}}$) site is less than half in NaHCO$_3$ (9.0 ($\pm$ 3.4) cm$^{-1}$ V$^{-1}$, Figures 4a and S8b) compared to CsHCO$_3$. Tuning rates for the $v$(CO$_{\text{atop(defect)}}$) site are more difficult to compare; the number of data points in the linear region are not enough to extract associated errors in the case of the anodic scan in CsHCO$_3$ and the cathodic scan in NaHCO$_3$. Tentative comparison of the tuning rates in opposite scan directions shows a significantly higher rate in the presence of Cs$^+$; however, the uncertainty in the linear regression means conclusions cannot be drawn regarding $v$(CO$_{\text{atop(defect)}}$). The potential dependent tuning rates for CO on Cu surfaces measured here fall within the distribution reported. The VSFG measured decrease in tuning rate for Na$^+$ and Cs$^+$ are in line with past reports, where it has been shown to vary in the order Na$^+$ < K$^+$ < Cs$^+$.$^{20}$

The cation induced change in CO binding mode correlates with ex-situ measured changes in electrode morphology due to eCO$_2$R: VSFG measurements show that the different cations lead to the population of a different set of surface sites by CO during eCO$_2$R on Cu. It is known that eCO$_2$R can lead to both reversible (with applied potential) and irreversible restructuring of the Cu surface, which would modify available sites for CO binding.$^{27,59,60}$ To assess cation dependent surface structuring, atomic force microscopy (AFM) was used to study the Cu surfaces after cycling through the potential step programme used for spectroelectrochemistry in CO$_2$ purged electrolytes, Figure 5a–d. AFM shows the electrochemical experiments lead to changes in the grain size of Cu surface post-experiment. Interestingly, there is a correlation between cation size and post-electrolysis Cu grain size; the mean grain size increases in the order of pre-bulk electrolysis (32.4 ± 7.0 nm) < Na$^+$ (68.7 ± 21.4 nm) < K$^+$ (93.3 ± 23.3 nm) < Cs$^+$ (139.5 ± 44.1 nm). The AFM map post-electrolysis also has the widest distribution of grain size post-electrolysis with Cs$^+$ present, indicating particularly high levels of surface
reconstruction using this cation. As the CO\textsubscript{bridge} site is present in CsHCO\textsubscript{3} when purging with CO\textsubscript{2} but not CO, AFM images have been recorded of the Cu surface after bulk electrolysis in CO purged CsHCO\textsubscript{3}, Figure 5e. Similar reconstruction occurs; however the average particle size is lower for the CO purged system (104.3 ± 29.9 nm), closer to that of the CO\textsubscript{2} purged KHCO\textsubscript{3} system. We note that it is important to take caution when interpreting \textit{ex-situ} characterisation of Cu surfaces as they are not true representations of surfaces under \textit{operando} conditions. \textit{In-situ} TEM shows that metallic Cu structures formed under operating conditions are rapidly changed upon exposure to air and subsequent Cu\textsubscript{2}O formation.\textsuperscript{61} Nonetheless, the clear difference in \textit{ex-situ} morphology post-electrolysis are indicative of differences in structure \textit{in-situ} depending on the cation used offering a rationale for the cation dependence of CO binding site measured \textit{in-situ} by VSFG spectroscopy.
**Figure 5:** AFM images showing surface morphology of the Cu electrode surface a) before bulk electrolysis, after bulk electrolysis in CO$_2$ purged 0.5 M b) NaHCO$_3$, c) KHCO$_3$ and d) CsHCO$_3$ and also in e) CO purged 0.5 M CsHCO$_3$. Image analysis methods are described in the experimental section.

**Discussion:** VSFG spectroscopy allows us to explore the potential dependence (and stability) of the atop binding modes in the presence of Na$^+$ and Cs$^+$, Figure 1. Overall, the potential dependent profiles of the CO$_{\text{atop(terrace)}}$ and CO$_{\text{atop(defect)}}$ $I_{\text{VSFG}}$ response are similar, with the CO$_{\text{atop(defect)}}$ increasing at potentials negative of $-0.7$ V. The slight decreases in $I_{\text{VSFG}}$ for CO$_{\text{atop(defect)}}$ at the most negative potentials studied ($-1.0$ V) in the presence of Na$^+$, but not Cs$^+$, indicates that Cs$^+$ leads to stabilisation of CO binding at the defect site and past studies have also proposed that this may occur.$^{9,11,23}$ The potential dependent tuning rates of the CO stretching modes need to be interpreted with caution. Deviations in the linear applied potential versus CO frequency relationship are observed at catalytic potentials ($<-0.8$ V) which is reasonable given that we anticipate both changes in the local environment (electrode structure), as indicated by the post-experiment AFM studies, and changes in CO concentration which will impact on dipolar coupling. The excellent agreement between the anodic and cathodic tuning rates in each electrolyte (Figure 4), despite clear differences in $I_{\text{VSFG}}$ observed (Figure 2), suggests the effects of dipolar coupling in the $-0.1$ to $-0.8$ V potential range in our experiments is minimal. It is striking that there is a significant increase in tuning rate of the $v$(CO$_{\text{atop(terrace)}}$) mode in the presence of Cs$^+$ compared to Na$^+$ which is indicative of an increase in local field strength at the Cu electrode in the presence of CO which will modify Cu-CO stabilities.

However the most striking result is that the nature of the CO binding site is strongly dependent on the cation used. With Na$^+$ CO is bound atop at both terrace and defect sites. With K$^+$ we propose that a more disordered surface with multiple sites occupied by CO occurs leading to no clear VSFG modes. When Cs$^+$ is used, which is widely reported to give the highest levels of eCO$_2$R activity,$^{11-17}$ there is a build-up of CO bound at bridge sites on the Cu surface which is not seen with other cations. The CO$_{\text{bridge}}$ site is considered electrochemically inactive$^{33}$ making it striking that it is observed using Cs$^+$, but not Na$^+$ which is usually less active for eCO$_2$R. One possible cause of the apparent contradictory results is miss-assignment of the CO$_{\text{bridge}}$ mode. For example, metal impurities can lead to different M-CO modes being present and these have been proposed to be the cause of erroneous assignments in SEIRAS studies.$^{62}$
Common causes of metal ions in the electrochemical cell, such as counter electrode dissolution and impurities from the supporting electrolyte itself, are addressed by the use of carbon counter electrodes and a thorough pre-electrolysis of the electrolyte solution (see experimental section). Furthermore, observation that the ν(CO) at ~1887 cm\(^{-1}\) on Cu is not present during spectroelectrochemical studies of CsHCO\(_3\) purged with CO, but that it is present during studies of the same electrolyte when CO\(_2\) purged, supports assignment to an intermediate formed during eCO\(_2\)R on Cu, namely multiply (bridge) bound CO\(_{x}\).\(^{30,33,34,38,39,41,47}\)

Although electrochemically inactive at the potentials studied here, the presence of bridge bound CO may still enhance eCO\(_2\)R activity for C\(_2\)+ products and a number of studies propose that C-C coupling is facilitated by bridge sites during eCO\(_2\)R, with calculations revealing that energetic barriers are lowest for coupling between CO\(_{atop}\) and CO\(_{bridge}\) sites.\(^{30,34,36,37,63}\) In fact, a recent study reports C\(_2\) FE over 90% at industrially relevant current densities and attributes this catalytic behaviour to the stable ratio of CO\(_{atop}/CO_{bridge}\) sites, observed via Raman spectroscopy.\(^{34}\) The observation by VSFG spectroscopy that CO\(_{bridge}\) is formed with CsHCO\(_3\), an electrolyte with an increased C\(_2\)/C\(_1\) product ratio when compared to K\(^+\) and Na\(^+\),\(^{11-17}\) strongly supports the involvement of the bridging site in CO-CO coupling.

Potential and pH induced surface reconstruction has been reported to trigger formation of the bridge binding site, with a high pH being particularly beneficial.\(^{33}\) Cu surface restructuring is also sensitive to a range of other factors including the initial presence of a surface oxide,\(^{64}\) local CO concentration/surface CO coverage\(^{27,48}\) and the nature of the electrolyte anion.\(^{38,65}\) The presence and nature of alkali metal cations has also been proposed to be a controlling factor\(^{18}\) with one study postulating that increased activity for C-C bond formation using Cs\(^+\) is due to electrolyte induced modification of the Cu surface.\(^{66}\) It is therefore striking that the detection of the CO\(_{bridge}\) mode \textit{in-situ} correlates with increased \textit{ex-situ} measured changes in electrode morphology post-electrolysis when Cs\(^+\) is used as the electrolyte during eCO\(_2\)R.

Conclusions

VSFG spectroscopy enables the study of a pc-Cu electrode without the need for surface modification/nano-structuring of the electrode which is often applied to linear vibrational spectroscopies to obtain suitable levels of surface sensitivity. There is increasing evidence that the morphology of the electrode can have a profound effect on catalytic activity making it important that spectroscopic studies are carried out on the most commonly employed electrode
structures. Past VSFG studies have identified \( \nu(C-H) \) but been unable to identify the important surface bound CO on pc-Cu; here we identify that the detection of CO at the pc-Cu depends on the choice of cation and specific reaction conditions. We measure high levels of CO\text{bridge} by VSFG when a Cs\(^+\) electrolyte is used. We find that there is a correlation between \textit{ex-situ} post electrolysis measurements of electrode morphology, which show that the degree of restructuring is cation dependent, and the presence of the CO\text{bridge} intermediates as measured by VSFG spectroscopy. These results suggest that a high level of bridge site formation is related to, or facilitated by, the Cu restructuring that happens as a result of the use of the Cs\(^+\) cations in the supporting electrolyte. Recent reports have indicated that multiple (bridge) bound CO may be electrochemically inert\(^ {33} \) but this work builds on the emerging evidence that CO\text{bridge} sites are a key intermediate in the CO-CO coupling step that is required for C\(_2\)\(^+\) formation during e\text{CO}2R.\(^ {30,34,36,37,63} \)

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**Data Availability:**

Underpinning data including multicurve fits are freely available at on the University of Liverpool Research Data Catalogue (DOI:10.17638/datacat.liverpool.ac.uk/2557)

**Experimental section**

VSFG cell preparation: All VSFG experiments were performed in a custom spectroelectrochemical “cross-cell”. The cross cell and all components were kept in Milli-Q water between experiments to avoid contamination by organics in the air, with additional thorough rinsing before assembly. The 3 mm diameter Cu (Alvatek) working electrode was
mechanically polished for 10 minutes using 1.0 µm then for a further 10 minutes using 0.05 µm alumina suspension. The electrode was rinsed thoroughly and sonicated in Milli-Q water between each polish. The working electrode was then secured into the cross cell with a Ag wire pseudo reference electrode (sanded, rinsed and sonicated before each experiment) and a glassy-carbon counter electrode (mechanically polished before each experiment). Electrolytes used were 0.5 M KHCO$_3$ (99.5%, Merck), NaHCO$_3$ (99.7%, Fisher) or CsHCO$_3$. Due to issues regarding impurities when using commercial CsHCO$_3$, CsHCO$_3$ was prepared from CsOH (99.95%, Merck) by purging with CO$_2$ (99.995%, BOC) for 1 hour, verified by a pH change from 13.6 to 7.4. All electrolytes were pre-electrolysed at $-0.025$ mA cm$^{-2}$ with Ti plates overnight prior to use to remove any metal impurities. Electrolytes were purged with CO$_2$ for 20 minutes in a sealed vial before being cannula-transferred into the cross cell. For the CO purged controls, the electrolytes were first degassed with N$_2$ for 20 mins before a 1 min purge with CO (99.97%, BOC). Working electrodes were pressed up against the 2 mm thick CaF$_2$ window after electrolyte transfer.

VSFG apparatus: A diagram of the VSFG setup is provided in Figure S1. The VSFG experiments were performed using a newly constructed spectrometer. 90% of the output PHAROS-PH1-SP (Light Conversion, 1030 nm, 10 kHz, 10 W, 170 fs pulse duration) laser system is used to generate both the nIR and mIR laser pulses required for the VSFG experiments. 1 W of the output is used to generate a time asymmetric narrow-band nIR pulse (1030 nm, 10 kHz, ~1.5 ps, ~13 cm$^{-1}$ linewidth) via an etalon (SLS Optics), which is directed to the sample through a half-wave plate (Thorlabs, WPH10M-1030) and polariser (ThorLabs, LPVIS050-MP2) to rotate the light to a horizontal polarisation, and p-polarised w.r.t reflection from the sample, at an angle of incidence of ~45°. A 950 nm long pass filter (Thorlabs, FEL0950) is also used to filter out second harmonic 515 nm light generated within the halfwave plate. This nIR pulse is focused by a 30 cm lens (Thorlabs LB1779-B), with the sample placed ~25 cm away from this lens, resulting in an approximate beam diameter at the sample of ~400 µm (1/e$^2$ diameter) with a power of ~20 mW. 8 W of the laser output is used in an IR OPA (Light Conversion, Orpheus-One-HE) to generate the broadband IR beam which can be tuned across the frequency range of interest (10 kHz, 170 fs pulse duration and ~150 cm$^{-1}$@1900 cm$^{-1}$). The mIR output passes through a twisted periscope to switch the polarisation from vertical to the desired horizontal polarisation. The purity of the polarisation is checked using a polariser (Thorlabs, LPMIR050-MP2) which is removed from the beam path prior to VSFG
measurement. The mIR beam is focussed onto the sample, with an approximate beam diameter of 300 µm using a Au parabolic mirror (Thorlabs, MPD249H-M01) at an angle of incidence of ~50° and $p$-polarisation w.r.t reflection from the sample, with a power of ~20 mW. A gas-purge generator removes H$_2$O, CO$_2$ and other contaminants in the air from the IR beam path. Resulting $p$-polarised VSFG light is directed through 950 (Thorlabs, FES0950) and 900 nm (Thorlabs, FES0900) short pass filters to remove the 1030 nm nIR beam. The beam goes through another polariser (Thorlabs, LPVIS050-MP2) before being focussed using a 15 mm focal length lens (Thorlabs, LA1540-B) through 150 µm slits and into the spectrograph (Andor, Kymera), and is detected on a CCD camera (Andor, iDus16). nIR/mIR delay is introduced using a linear stage (ThorLabs, LTS300C) on the nIR beam path. The spectrograph was calibrated using Ne spectral lines and the rovibrational P, Q and R CO$_2$ branches).

Spectroelectrochemical measurements: Laser beams were always initially aligned off a Au mirror to allow comparison to an IR profile on a clean surface. The potential was stepped from $-0.5 \rightarrow -1.0 \rightarrow -0.1 \rightarrow -0.5$ V on Cu using a potentiostat (PalmSens). Each potential step was held for 150 s, before sweeping at 50 mV s$^{-1}$ to the next step to avoid current spikes when jumping between potentials. After equilibration for 30 s, VSFG spectra were acquired for 120 s. Spectra were collected for nIR/mIR delays of 1000, 670 and 0 fs in that order.

AFM characterisation: A contact-mode cantilever (Pointprobe®, NanoWorld®, Switzerland) was inserted onto an AFM nose cone (Keysight Technologies, United States). The nose cone was loaded into an atomic force microscope (5500 AFM, Keysight Technologies, United States) and then the laser was aligned to situate the beam spot on the cantilever tip. After approaching the cantilever, images were acquired using Keysight NanoNavigator software (version 1.8.2), morphologies were analysed using Gwyddion (version 2.61) and size distributions were obtained using ImageJ (version 1.53t). In ImageJ the scale of the edge of each image was set to 2 µm and the diameters of a randomly selected set of 100 particles were determined using the measure function.

References

1. L. Fu, Z. Ren, W. Si, Q. Ma, W. Huang, K. Liao, Z. Huang, Y. Wang, J. Li and P. Xu,


