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Deactivation of a Single-Site Gold-on-Carbon Acetylene Hydrochlorination Catalyst: An X-ray Absorption and Inelastic Neutron Scattering Study


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Supporting Information

ABSTRACT: Single-site Au species supported on carbon have been shown to be the active sites for acetylene hydrochlorination. The evolution of these single-site species has been monitored by Au L₃ X-ray absorption spectroscopy (XAS). Alternating between a standard reaction mixture of HCl/C₂H₂ and the single reactants has provided insights into the reaction mechanism and catalyst deactivation processes. We demonstrate that oxidative addition of HCl across an Au(I) chloride species requires concerted addition with C₂H₂, in accordance with both the XAS measurements of Au oxidation state and the reaction kinetics being first order with respect to each reactant. Excess C₂H₂ changes the Au speciation and results in the formation of oligomeric acetylene species which were detected by inelastic neutron scattering. Catalyst deactivation at extended reaction times can be correlated with the formation of metallic Au particles. These Au(0) species generated during the sequential gas experiments, or after prolonged reaction times, results in the analysis of the normalized near-edge white line intensity becoming an unsuitable guide for identifying the active Au species, affecting the strong correlation between normalized white line height and VCM productivity usually observed in the active catalyst. Thus, a combination of scanning transmission electron...
INTRODUCTION

Vinyl chloride monomer (VCM) is a large-scale chemical intermediate used almost exclusively as a precursor to polyvinyl chloride (PVC). Annually over 13 million tons of VCM are produced by acetylene hydrochlorination. Traditionally, acetylene hydrochlorination has been catalyzed using mercuric chloride supported on high surface area activated carbon. While this catalyst is highly active and has been industrially employed for several decades, it is extremely volatile and highly toxic.\textsuperscript{1–6} Following the prediction of Hutchings in 1985 that Au would be an effective catalyst for the reaction based on a correlation with electron affinity,\textsuperscript{7} Au supported on carbon has recently been validated as a replacement catalyst for this large-scale industrial process.\textsuperscript{8}

The preparation method for making the Au/C catalysts has been demonstrated to strongly influence the corresponding structure and activity of the catalyst.\textsuperscript{7–9} Active catalysts were prepared by the impregnation of HAuCl\textsubscript{4} onto C in strongly oxidizing solutions of nitric acid or aqua regia, while it was found that using an aqueous HAuCl\textsubscript{4} solution produced catalysts with low activity. Characterization of these catalysts by a range of techniques, including X-ray photoelectron spectroscopy (XPS) and scanning transmission electron microscopy (STEM), suggested that the active catalysts had high Au dispersion and contained both oxidized Au(III) and Au(I) species. While these catalyst structure–activity studies revealed the importance of cationic Au species, the exact distribution of oxidation states and their chemical nature proved difficult to determine. First, XPS and STEM analyses can easily alter the relative distributions of Au(0), Au(I), and Au(III) species and can induce the formation of nanoparticle in the samples through beam modification.\textsuperscript{10–15} Second, the reaction occurs under an aggressive chemical environment which can cause dynamic changes to the catalyst structure during reaction.

Recently, we reported the first detailed in situ X-ray absorption spectroscopy characterization study following the behavior of these Au/C catalysts during the acetylene hydrochlorination reaction.\textsuperscript{9} Analysis of the Au L\textsubscript{3}-edge X-ray absorption near-edge structure (XANES) and the extended X-ray absorption fine structure (EXAFS) showed the operating catalyst comprised of atomically dispersed cationic species with no Au nanoparticles being present in the most active Au/C catalysts. During the reaction, distinct changes in the XANES were observed, which clearly correlated to the catalyst productivity, showing that the active catalysts were comprised of atomically dispersed cationic gold in both Au(III) and Au(I) oxidation states. While this provided new details of the nature of the active catalyst under operating conditions, we reason that studying the deactivation of the catalyst will provide a more profound understanding of the catalytic behavior and the reaction mechanism to generate the information needed to design catalysts that exhibit extended lifetimes under industrial operating conditions.

Previously, from post-reaction ex situ XRD and Mössbauer spectroscopy studies, deactivation of such Au/C catalysts operating above 120 °C has been attributed to the reduction of gold from its cationic form to generate metallic Au nanoparticles.\textsuperscript{16,17} Also, the formation of polymeric organic species on the catalyst surfaces, as observed after long reaction periods under industrial conditions, resulting in carbon fiber formation\textsuperscript{7} has been considered as a vector for the deactivation of this class of catalyst. Strategies to inhibit both deactivation pathways include the addition of other metal chlorides or dopants such as nitrogen and phosphorus.\textsuperscript{18,19}

This study investigates if the formation of Au(0), which has previously been stated as a deactivation mechanism from post-reaction studies, can be directly correlated with a decrease in VCM productivity under reaction conditions.\textsuperscript{16,20–23} In addition, the role of the individual reactants, in particular, acetylene, has been studied and correlated with the observed Au speciation, particularly the formation of polymeric organic species on the catalyst surface.\textsuperscript{1} In addition, observations of changes in Au L\textsubscript{3}-edge XANES spectra during gas-switching experiments have been used to further elucidate the reaction mechanism.

RESULTS AND DISCUSSION

Catalyst Structure during Thermal Treatment. Highly active Au/C catalysts prepared by the impregnation of Au from a solution of aqua regia have previously been shown to consist of atomically dispersed cationic Au and be highly active for acetylene hydrochlorination.\textsuperscript{9} To test the thermal stability of highly dispersed Au species, the catalyst was heated to 350 °C (at a rate of 5 °C/min) under an inert N\textsubscript{2} atmosphere (to prevent the combustion of the carbon support) while being characterized using in situ XRD (Figure 1a). Given that the single-site cationic Au species, shown by XANES analysis\textsuperscript{7} to be present in catalyst, are undetectable by XRD, no diffraction peaks from Au were observed at room temperature. The dispersed Au(III)/Au(I) species in the 1 wt% Au/C (1% Au/C-AR) catalyst showed high thermal stability from ambient temperature up to 250 °C, with no characteristic reflections of fcc Au being observed over this temperature range. Upon reaching 300 °C, reflections corresponding to the (111), (200), (220), and (310) Au lattice planes became evident, demonstrating the formation of discrete Au nanoparticles and implying that the Au species have now become mobile on the support material. Scherrer analysis of the (111) reflection at 300 °C gave an average crystallite size of 29 nm. As expected, the mean nanoparticle size increased even further as the temperature was increased to 350 °C as did the relative intensity of the Au diffraction pattern, demonstrating that a greater amount of Au is in a crystalline state.

Clearly 1% Au/C-AR is thermally stable from the perspective of XRD analysis up to 300 °C under an inert atmosphere. However, it should be noted that even if single-site cationic species are thermally stable under an inert atmosphere, there is significant scope for different AuCl\textsubscript{4} species, which are not differentiable by XRD, to exist and evolve during heating. X-ray absorption fine structure (XAFS) can be used to show potential changes in Au oxidation state and speciation. Monitoring the changes in white line height, a feature detectable in the XANES region which corresponds to the Au 2p\textsubscript{3/2} → 5d primary transition (see experimental section for more details), showed that while heating the catalyst to the reaction temperature (200 °C) under Ar, the Au(III)/Au(I) ratio significantly changed from Au(III) chloride to Au(I) chloride (Figure 1b).\textsuperscript{9} Therefore, while it is true to state single-site cationic species are retained on
heating, the speciation and oxidation state changes even in the absence of reactant gases.

**XANES Analysis of Au/C-AR Catalyst under Different Reactant Gases.** It has been postulated, based on combined in situ XAFS and DFT studies, that the reaction mechanism for Au-catalyzed acetylene hydrochlorination proceeds via an oxidative addition and reductive elimination process to generate VCM, which makes the redox properties of the cationic Au species an essential requirement for activity.\(^7,9\) Given the postulated redox mechanism and likely deactivation mechanism, namely, the agglomeration and reduction of cationic species to metallic Au, it can be expected that exposure to the reaction mixture or indeed each individual reactant gas has the potential to affect Au speciation. In previous studies, ex situ XPS of Au/C catalysts collected after exposure to different combinations of reactants showed that HCl increased the concentrations of Au(III) and Au(I) at the expense of Au(0), while C\(_2\)H\(_2\) resulted in a reduction of Au(III) to Au(I).\(^7\) To reiterate, these experiments were carried out ex situ, potentially resulting in significant changes to the Au speciation during removal from the reactor and exposure to air and moisture before insertion into the XPS instrument. Another important point to note is that relative changes in the concentrations Au(III) and Au(I) species appear minor in comparison to the dominant signal associated with Au(0). A key point to consider during XPS experiments is the significant photoreduction of gold chloride compounds that can occur, which results in overestimation of the Au(0) concentration. While appearing counterintuitive, the higher incident photon energy associated with Au L\(_3\)-edge XAFS (11.919 keV for Au L\(_3\)-edge XAFS versus 1.487 keV for Al source XPS) results in lower rates of photoreduction for XAFS experiments compared to Al source XPS due to the significantly lower absorption cross-section in the former case.\(^8\) For this reason XAFS can be considered a less artifact prone and more representative characterization technique for Au valence state determination than XPS.

To clarify the influence of both combinations and the individual reactants on Au speciation, under reaction conditions, a sequential flow in situ Au L\(_3\)-edge XAFS experiment (as detailed in the Experimental Section) was performed to study the 1% Au/C-AR catalyst. A plot of VCM productivity with respect to reaction time and the normalized white line intensity is shown in Figure 2 during exposure to a well-defined sequence of different reaction gas mixtures.

The catalytic behavior of the 1% Au/C-AR catalyst for the first 240 min under the reaction mixture shows the same trend as in our previous study in which an induction period of ca. 180 min was noted before VCM productivity reached steady state.\(^9\) During this induction period, the normalized Au L\(_3\)-edge white line intensity dramatically increased from 0.67 to 0.94 before slowly decreasing back to a value of 0.70 at steady state. As previously reported,\(^7\) this initial change in white line height is associated with the rapid formation of Au(III) chloride-like species followed by its gradual reduction to one principally containing Au(I) chloride.

After 240 min of reaction the catalyst was then exposed to HCl only diluted in argon [HCl/Ar] for ca. 70 min (i.e., no C\(_2\)H\(_2\)) at a flow rate matching the total flow of the reaction mixture. As expected, while flowing HCl/Ar only, no vinyl chloride monomer formation was observed. Interestingly, in the

![Figure 1](image1.png)  
**Figure 1.** In situ characterization of the 1% Au/C-AR. (a) Powder X-ray diffraction (pXRD) patterns acquired when increasing temperature under inert nitrogen atmosphere. (b) Changes in white line height during heating to reaction temperature (200 °C) under Ar at 5 °C min\(^{-1}\). (Dotted lines represent the white line intensities of the Au(I) [AuCl\(_4\)]\(^-\) standard (value of 0.6) and the Au(III) KAuCl\(_4\) standard (value of 1.1)).

![Figure 2](image2.png)  
**Figure 2.** VCM productivity during the sequential gas exposure experiment combined with in situ Au L\(_3\)-edge XANES data of 1 wt % Au/C-AR. VCM production (black triangles) as a function of time online and the corresponding normalized white line intensity values (blue squares) derived from XANES measurements. Reaction sequences: (i) HCl + C\(_2\)H\(_2\)/Ar, (ii) HCl/Ar, (iii) second HCl + C\(_2\)H\(_2\)/Ar, (iv) C\(_2\)H\(_2\)/Ar, and (v) third HCl + C\(_2\)H\(_2\)/Ar exposure. (*) No white line data is shown for step iv in the sequence.
induction period was seen. As the 1%Au/C-AR catalyst was not deactivated by the HCl/Ar gas treatment but neither underwent oxidation, it is reasonable to conclude that C₂H₂ must somehow assist in the oxidative addition of HCl, facilitating the reaction, based on the Au(I)–Au(III) redox couple. We postulate that C₂H₂ assists in the oxidative addition of HCl in the first step of the reaction mechanism with a concerted HCl and C₂H₂ addition to the Au(I)Cl species. This differs from our previously proposed mechanism in which the oxidative addition of HCl to form AuCl₂H was then followed by C₂H₂ addition and the reductive elimination of VCM. Further supporting evidence for this revised mechanism has been obtained by studying the order of reaction with respect to HCl and C₂H₂ (as discussed later).

Once the steady state in step iii of the sequence had been reached, the catalyst was exposed to dilute acetylene only [i.e., C₂H₂/Ar] (step iv). In common with the HCl/Ar environment, no VCM production was observed with exposure to acetylene alone. On introduction of C₂H₂/Ar the XANES spectra (Figure 3b) were significantly altered, with the gradual loss of features at 11 921 (white line) and 11 931 eV and the simultaneous appearance of a new feature at 11 923 eV along with the absorption edge shifting to higher energy. The new XANES spectra were not clearly attributable to either Au(III) or Au(I) chloride-like speciation. Figure 4 shows a comparison between C₂H₂/Ar XANES spectra and Au(I) chloride, and a metal standard: (black solid line) Au (III) standard of [AuCl₅]⁻, (green dashed line) Au(I) standard of [AuCl₃]⁻, and (red dashed line) Au (0) metal foil.

The new XANES spectra were not clearly attributable to either Au(III) or Au(I) chloride-like speciation. Figure 4 shows a comparison between C₂H₂/Ar XANES spectra and Au(I) chloride, and a metal standard: (black solid line) Au (III) standard of [AuCl₅]⁻, (green dashed line) Au(I) standard of [AuCl₃]⁻, and (red dashed line) Au (0) metal foil.

Reintroduction of the full reaction mixture (i.e., HCl/C₂H₂/Ar), after HCl/Ar treatment, resulted in an almost immediate return of VCM production (Figure 2), although initially at a marginally lower rate of 2.28 mol kg⁻¹ h⁻¹ compared to the previous steady-state productivity rate of 2.74 mol kg⁻¹ h⁻¹. The VCM productivity then steadily increased over the next 115 min before reaching a new steady-state productivity value of 2.70 mol kg⁻¹ h⁻¹. During this period of increasing activity the normalized white line intensity also gradually increased. Clearly, the catalyst was not significantly deactivated under the HCl/Ar atmosphere, although the system was perturbed and a second, albeit smaller,
interaction that is potentially detrimental to the catalytic performance. Further analysis of this species from EXAFS is presented in the following section.

Reintroduction of reactant gases to the 1% Au/C-AR catalyst after C₂H₂/Ar treatment (step v) resulted in VCM productivity returning. However, unlike the original induction period (step i) or following the HCl/Ar treatment (step iii), the productivity did not increase with reaction time. Rather it decreased from 2.24 to 1.76 mol kg⁻¹ h⁻¹ over 90 min, providing evidence that the Au acetylene interaction occurring in step iv had partially poisoned the catalyst. Interestingly, the XANES of the catalyst on reintroduction of HCl/C₂H₂/Ar reactant gases in step v show an almost instantaneous return to the more recognizable spectrum with the characteristic Au(I)/Au(III) features seen in the previous reaction steps i–iii. The re-emergence of an identifiable white line feature could therefore be monitored, whose intensity was found to remain relatively constant at ~0.69, with only a marginal increase in height over this final reaction period (Figure 2). Notably, the relationship between the normalized white line and VCM productivity was clearly altered relative to that observed initially and after HCl/Ar treatment.

A linear correlation between the normalized white line height and VCM productivity for the initial induction period (step i in Figure 2), as previously reported⁹ and shown in Figure 5, is strong with a Pearson’s correlation coefficient value r of −0.994 and observed gradient of 0.104 (±0.002). A similar correlation between the normalized white line height and VCM production for the reaction period after HCl treatment (step iii in Figure 2) is also evident, except that the gradient of the trend line changed to 0.057 (±0.007). A similar plot for the reaction period (step v) after C₂H₂ treatment gave a gradient of only 0.009 (±0.005), indicating that the relationship between productivity and normalized white line height was now almost invariant.

The simplest explanation for the loss of correlation after step v is that one or more additional spectator species, which contribute to the XANES in the region of the Au(I)/Au(III) white line, are present after treatment with C₂H₂/Ar and to a lesser extent after exposure to HCl/Ar. Such an explanation is supported by comparison of the XANES spectra of the catalyst at steady state for each reaction period (Figure 6), i.e., at the end of the initial induction period (step i in Figure 2), at steady state after HCl treatment (step iii in Figure 2) and at the very end of the reaction after C₂H₂ (step v in Figure 2). A 1 eV shift in the position of the white line, between the initial and the final steady-state spectra, indicates a notable change in Au speciation that is not apparent from white line height measurements alone.

Linear combination fitting (LCF) of these three spectra and those of the catalyst under HCl/Ar and C₂H₂/Ar using Au(III), Au(I) and Au metal standards was performed to gain greater understanding of the changes in the XANES spectra and explain the loss in correlation between white line and VCM productivity in step v (Table 1). First, fitting of the catalyst under C₂H₂/Ar was not successful in that no acceptable fit was found with the three standards chosen, further highlighting a strong interaction between C₂H₂ and the gold species present on the catalyst.

Figure 5. Comparison of white line height versus VCM productivity for the 1 wt% Au/C-AR catalyst under HCl/C₂H₂/Ar reaction gas after different single-reactant gas treatments: (solid black circles) catalyst during the first induction period (productivity and white line data from step i, Figure 2), (open triangles) catalyst after HCl/Ar treatment (productivity and white line data from step iii, Figure 2), and (half-filled diamonds) catalyst after C₂H₂/Ar treatment (productivity and white line data from step v, Figure 2).

Table 1. Linear Combination Fitting of the 1 wt% Au/C-AR Catalyst at the End of Each Sequence of Gas Switchingᵃ

<table>
<thead>
<tr>
<th>reaction step</th>
<th>quantification (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Au(III)</td>
</tr>
<tr>
<td>first reaction mixture (step i)</td>
<td>28(4)</td>
</tr>
<tr>
<td>HCl/Ar (step ii)</td>
<td>25(5)</td>
</tr>
<tr>
<td>second reaction mixture (step iii)</td>
<td>24(5)</td>
</tr>
<tr>
<td>third reaction mixture (step iv)</td>
<td>20(3)</td>
</tr>
</tbody>
</table>

“Percentage of Au associated with Au(I) chloride (standard [AuCl₃⁻]); Au(III) chloride (standard KAuCl₄) and Au (0) metal (standard Au foil).” Reaction sequence; step i = HCl/C₂H₂/Ar; step ii = HCl/Ar; step iii = HCl/C₂H₂/Ar; step iv = C₂H₂/Ar, and step v = HCl/C₂H₂/Ar. “NB. Analysis of step iv (C₂H₂/Ar) is excluded due to a poor fit.

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The speciation of the of catalyst at the end of the first induction period under HCl/Ar and during the second reaction period was comparable in terms of their Au(III) and Au(I) ratios; however, a small amount of metallic Au was possibly present from step ii. Given that the calculated Au(0) concentration was comparable to the estimated error, this result taken in isolation appears of limited value. However, clear evidence of metallic Au (ca. 21 wt %) was found from LCF of the catalyst after C2H2/Ar treatment (step iv in Figure 2). As metallic Au is known to be catalytically inactive for acetylene hydrochlorination, this observation correlates well with the loss in activity seen after C2H2 treatment. Furthermore, given that Au(0) has a normalized absorption value of 0.6 at the energy of the Au(III) white line (11 920 eV) the presence of this species will reduce the white line intensity in a XANES spectrum of a sample containing Au(III), Au(I), and Au(0). In doing so the presence of Au(0) will disrupt the correlation between normalized white line height and VCM productivity. An example of how 0, 25, or 50 mol % Au(0) effects the gradient of the white line height versus Au(III):Au(I) is shown in Figure S1 where increasing Au(0) content within the sample decreases the size of the gradient of the plot.

Given that LCF using reference KAuCl4 [AuCl4]− and Au foil materials cannot comprehensively model single-site AuCl4 species supported on a carbon surface, previous studies used the simpler analysis of white line height to give an average Au oxidation state. While white line height was satisfactory for analysis of the initial analysis (induction period and initial steady-state testing) of a two-component system (Au(I) chloride and Au(III) chloride), LCF despite its flaws becomes required with this more complex multispeciation.

EXAFS Analysis of Au/C-AR Catalyst under Different Reactant Gases. While XANES analysis provides relevant information concerning Au speciation, the complex nature of the spectra requires additional EXAFS analysis to clarify interpretation. EXAFS of the catalyst at steady-state reactivity, before and after HCl/Ar treatment and after C2H2 treatment, can provide further evidence of an increased contribution from metallic Au after sequential gas treatments. Figure 7 shows the magnitude of the Fourier transform data of the catalyst at the end of steps i, iii, and v in the reaction profile. The predominant feature in all samples is that of first shell Au−Cl interactions with multiple Au−Cl scattering events beyond 3 Å. However, features associated with Au−Au interactions between 2.5 and 3 Å also become noticeable after treatment with C2H2 (although the FT magnitude is small), confirming the deduction form linear combination fitting of the XANES that metallic Au nanoparticles are generated during exposure to C2H2/Ar.

EXAFS analysis (Figures S2 and S3, Table 2) can also provide further information on the nature of the Au speciation, especially while the catalyst was under a C2H2/Ar-only environment. While the Au L3 edge XANES of this species was clearly different to Au (I) or Au(III) chloride, EXAFS shows that Au−Cl interactions are still present in the C2H2/Ar-treated sample. The magnitude of this scattering in the Fourier transform data (Figure S3) is significantly lower under C2H2/Ar than the catalyst in the presence of both reactants (C2H2 + HCl/Ar). No additional features associated with Au−C or Au−Ar interactions were observed in the C2H2-treated material, suggesting no significant Au−C2H2 complex or Au(0) nanoparticle formation. Taking this into account, along with the dampened oscillations at higher wavenumbers in the χ-space data (Figure S2), a model of AuCl4 with two Au−Cl bond lengths that were partially out of phase was considered. The results of fitting two first-shell Au−Cl lengths to the data are given in Table 2, which yields an Au−Cl combined coordination number of 1.8(4) compared to that of 2.5(1) for the catalyst under normal steady-state conditions (fitted with a single Au−Cl distance). It should be noted that no reasonable fitting of 2 Au−Cl paths was obtainable for any other sample investigated in steps ii, iii, and v. It was observed that a small component of the Au−Cl interactions in the catalyst under C2H2/Ar were notably shorter, at 2.110(57) Å compared to 2.273(6) Å for the catalyst at steady-state conditions. Therefore, we conclude that the presence of excess acetylene results in the formation of some shortened Au−Cl interactions but with no direct evidence of Au−C2H2 bond formation. It should also be noted that evidence of multiple Au−Cl distances was also previously observed during the induction period of the Au/C-AR catalyst in step i and illustrates that complex interactions between AuCl4 are present even under standard reaction conditions.

EXAFS analysis (Table 2) of the catalyst at steady state under HCl/Ar (step ii) and the second reaction mixture HCl/C2H2/Ar (step iii) showed little difference from that at steady state in the first reaction period. A slight reduction in Au−Cl CN number under HCl/Ar was observed, in line with the moderate reduction in white line height also noted during this period. For the sample at steady state after the C2H2/Ar cycle (i.e., at the end of step v), a model with both a Au−Cl and a Au−Au interaction was found to provide good fitting parameters, further supporting the presence of metal in the deactivated catalyst.

XAFS sequential flow experiments highlighted several interesting observations that have not been observed in previous ex situ studies, namely, (i) the oxidation of the Au(I)−Cl species appears far more facile in the presence of C2H2 and (ii) that C2H2 directly interacts with AuCl4 species, significantly altering the XANES spectra and changes Au−Cl lengths when no HCl is present. The first point regarding Au(I)−Cl oxidation can be further clarified by performing kinetic studies of the reaction to determine orders of reaction with respect to the reactant gases. If the apparent requirement for Au(I)−Cl oxidation is the concerted addition of HCl and C2H2 (as suggested from XANES data) and if this step is rate limiting then the reaction should be first-order dependent with respect to each reactant. To prevent apparent zero-order responses, each reactant investigated was
Figure 9

Table 2. In Situ EXAFS Fitting for Au L₃-Edge Data of the 1 wt % Au/C-AR Catalyst during the Sequential Gas Experiment

<table>
<thead>
<tr>
<th>reaction step</th>
<th>scattering path</th>
<th>coordination number (Å)</th>
<th>σ (Å)</th>
<th>θ (°)</th>
<th>E_f (eV)</th>
<th>R factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>first reaction mixture (step i)</td>
<td>Au–Cl</td>
<td>2.5(1)</td>
<td>2.27(1)</td>
<td>1.2(9)</td>
<td>0.0012</td>
<td></td>
</tr>
<tr>
<td>HCl/Ar (step ii)</td>
<td>Au–Cl</td>
<td>2.4(1)</td>
<td>2.27(1)</td>
<td>3.3(0.99)</td>
<td>0.007</td>
<td></td>
</tr>
<tr>
<td>second reaction mixture (step iii)</td>
<td>Au–Cl</td>
<td>2.5(1)</td>
<td>2.27(1)</td>
<td>2.55(66)</td>
<td>0.004</td>
<td></td>
</tr>
<tr>
<td>C₂H₂/Ar (step iv)</td>
<td>Au–Cl (1)</td>
<td>1.4(2)</td>
<td>2.29(3)</td>
<td>1.04(2.84)</td>
<td>0.026</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Au–Cl (2)</td>
<td>0.5(2)</td>
<td>2.11(6)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>third reaction mixture (step v)</td>
<td>Au–Cl</td>
<td>1.7(3)</td>
<td>2.26(1)</td>
<td>2.24(1.89)</td>
<td>0.013</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Au–Au</td>
<td>1² ²</td>
<td>2.84(5)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Fixed parameters:

S² = 0.75 and 2σ² (Å²) = 0.0037

EXAFS data was taken under steady-state conditions during each sequential gas sequence step (i.e., when no change in three consecutive spectra was observed). Fitting with multiple Au–Cl paths was attempted for all spectra. However, apart from C₂H₂/Ar (step iv), unrealistic fits such as high R factors and/or negative amplitudes for second paths were obtained when using multiple paths. Debye–Waller and amplitude reduction factors were calculated from the fitting of a KAuCl₄ standard with a fixed CN of 4. Fittings of the Au–Cl and Au–Au for step v were performed fixing the Au–Au coordination number to 1.

studied over a range of highly dilute concentrations. The linear dependence of both HCl and C₂H₂ reaction rate, as shown in Figure 8, demonstrates first-order dependence with respect to each reactant and second-order reaction kinetics overall. The combined evidence from in situ XANES experiments and the measured first-order dependence for each reactant provides strong evidence for a concerted HCl and C₂H₂ addition to the Au(I)–Cl active site.

**Inelastic Neutron Scattering (INS): Effect of C₂H₂ on Au Complex Speciation.** Vibrational spectroscopy provides a viable method to clarify the effect of C₂H₂ on Au complex speciation, as seen during the C₂H₂/Ar treatment in step iv of the reaction sequence. As the carbon support in the catalyst would be highly absorbent of infrared radiation, we attempted inelastic neutron scattering (INS) spectroscopy. While INS is suited to observing hydrogen speciation in the C₂H₂/Ar-treated sample and therefore the nature of the Au complex, it suffers from low signal-to-noise (due to the required interaction of the neutron with the small nucleus of the absorber). To mitigate this problem a more heavily loaded 2 wt % Au/C-AR sample was prepared to increase the concentration of Au species available for interaction with acetylene in the sample. XAFS, XRD, and STEM analysis of the 2 wt % Au/C-AR material demonstrated that this more highly loaded catalyst was still comprised of single-site cationic Au species with a negligible concentration of metallic gold nanoparticles (Figure S4, Figure 9a–d).

The 2 wt % Au/C-AR catalyst was sequentially treated under helium and a dilute acetylene gas mixture (5% C₂H₂/Ar or concentrated acetylene (100%) in a reaction cell for ca. 30 min at 200 °C. Between each step the sample was cooled (to minimize Debye–Waller contributions) and the cell flushed with He prior to acquisition of the INS spectrum. The resulting spectrum provides detail of acetylene species while the catalyst is exposed to an excess of acetylene under reaction conditions. The spectra of 2 wt % Au/C-AR sample prior to acetylene treatment (used as a background spectrum), 2 wt % Au/C-AR after treatment in 5% acetylene, and 100% acetylene are shown in Figure S5. Reaction with the C₂H₂(5%)/He mixture resulted in no observable changes in the spectrum, whereas in contrast reaction with 100% C₂H₂ results in bands at 2990, 1295, 961, 700, and 640 (and possibly 400) cm⁻¹. The difference spectra C₂H₂ (5%) dosed minus the background, C₂H₂ (100%) dosed minus the background, and C₂H₂ (100%) dosed minus C₂H₂ (5%) dosed are shown in Figure 10.

INS spectra of metal–acetylde–silver, silver/copper, copper, silver/copper, and gold complexes have been widely reported. The bands seen in the difference spectra of (100%) C₂H₂ – background were compared to that of Au–C≡C–H modeled as an isolated molecule, an adsorbed molecule on a single carbon atom of a graphene sheet (to model the carbon
support), and an adsorbed molecule above the center of a graphene hexagon (see Figure S6). All three models exhibit $\equiv$ $C$$-H$ stretches at $\sim 3300$ cm$^{-1}$, a $\equiv C$$-C$ stretch at $\sim 2000$ cm$^{-1}$, a $C$$-C$$-H$ bending mode at $\sim 600$ cm$^{-1}$, and an Au$-C$ stretch at $\sim 250$ cm$^{-1}$. As shown in Figure 11, none of the calculated spectra match the observed spectrum of the catalyst after acetylene treatment. Given the possibility of Au(0) formation after acetylene treatment as suggested from LCF of XANES and EXAFS analysis, models for acetylene bound to Au nanoparticles were also considered. However, in common to the metal–acytlyde spectra no correlation could be found between the models of acetylene bound to an Au$_4$ tetrahedron (representing a small Au cluster) and the experimental spectrum.

An alternative assignment to the observed bands on the 2 wt % Au/C-AR catalyst after C$_2$H$_2$ treatment is that of oligomerized acetylene species. Previously, carbon fibers have been observed by TEM analysis to have grown on spent Au/C-AR catalysts after 24 days time-online. The INS spectrum of polyacetylene shown in Figure 12 exhibits bands at similar energies to those seen on the 2 wt % Au/C-AR catalyst after C$_2$H$_2$ treatment. However, there are several modes absent in the 2 wt % Au/C-AR catalyst after C$_2$H$_2$ treatment, as compared to polyacetylene. Potentially these absent modes are associated with shorter oligomeric species on the catalyst surface. To test this possibility, all-trans-1,3,5,7-octatetraene (inset in Figure 12) was used as a model for a short-chain acetylene oligomer. Figure 12 compares the experimental and calculated spectra and indicates good agreement, particularly regarding the 1295 cm$^{-1}$ mode that is clearly present.

The used catalyst after the INS experiment was then analyzed ex situ by XAFS and STEM (Figure S7). The XANES spectrum recorded is almost identical to the one obtained for the 1% Au/C-AR catalyst during step iv of the sequential flow experiment (C$_2$H$_2$/Ar). Moreover, also in this case, the Au$-Cl$ intensity in the EXAFS FT decreased, compared to the spectrum of the freshly prepared catalyst shown in Figure S7, while the STEM-HAADF images (Figure 13) show that the gold is still predominantly dispersed as atomic on the carbon support.

The INS data shows for the first time the presence of oligomeric acetylene species, while the Au XANES shows a clearly altered Au species after C$_2$H$_2$ treatment. Unfortunately, there is no clear evidence that these two species are connected, as no evidence of Au–acytlyne bonding was found from either XANES or INS. However, we provide clear evidence of the significant surface acetylene species that become evident on exposure to a C$_2$H$_2$ excess in a reactant gas mixture, which is known to be detrimental to catalytic activity.

In Situ Deactivation of 2% Au/C-AR during the Acetylene Hydrochlorination Reaction. Catalyst deactivation, associated with Au(0) formation and a loss of the correlation between white line height and VCM productivity, was observed in the 1% Au/C-AR catalyst over multiple different sequential gas experiments. To clarify if a similar deactivation process occurs, without various sequential gas treatments, a continuous extended time-online experiment with a constant gas composition coupled with in situ XAFS monitoring could in
principle be performed. Such an experiment is not easily carried out with a 1 wt % Au/C-AR catalyst as the deactivation period is too long to be monitored over a single synchrotron beam time experiment. However, the 2 wt % Au/C-AR catalyst used for INS studies, with its comparable initial single-site dispersion of AuCl₅⁺ species, would be expected to undergo a more rapid deactivation associated with Au(0) particle formation, due to the higher concentration of Au available for agglomeration.

The structural behavior of the 2 wt % Au/C-AR catalyst was found to be initially comparable with that of the 1 wt % Au/C-AR material with a decrease in normalized white line height upon heating under Ar from 1.07 at 25 °C to 0.68 at 200 °C (Figure S8). Linear combination fitting of the 2 wt % Au/C-AR catalyst at 200 °C confirmed the absence of Au(0) before introduction of the reaction mixture. As the reaction mixture was introduced the normalized white line increased from 0.68 to 0.85 (Figure 14a), as observed in the 1 wt % Au/C-AR catalyst. The white line height then began to decrease over a 20 min period, commensurate with an increase in VCM productivity, as seen for the induction period of the 1 wt % catalyst. From this point in the reaction the evolution of VCM productivity and association with normalized white line height, diverges for the 2 wt % Au/C-AR catalyst. After ca. 40 min reaction, the higher loading catalyst undergoes a gradual and prolonged loss in activity, with VCM productivity dropping by 40% between 40 and 200 min time-online. As predicted from the sequential gas experiments, on the 1 wt % Au/C-AR catalyst, the observed normalized white line height remained relatively consistent over the deactivation period. As the reaction proceeded for another 150 min, EXAFS and LCF showed that the catalyst developed considerable amounts of metallic Au (ca. 22.5%) in addition to AuCl₅⁺ species (Figures 14a, 14b, and 15).

HAADF-STEM images of the used 2 wt % Au/C-AR catalyst confirmed the presence of single-site species in addition to larger particles of Au (Figures 16a–c) in the 20–100 nm size range. This observation suggests that once the AuCl₅⁺ single-site species reduced to Au(0), sintering occurred very efficiently under reaction conditions. Counting the numbers of Au atoms over certain areas of the carbon support from several HAADF-STEM images gave estimated Au atom densities of 0.470 and 0.154 atoms/nm² for the unused and used 2 wt % Au/C-AR catalysts, respectively (see Table 3). This suggests that ~2/3 of atomic Au transformed into large Au particles during the reaction, leading to a significantly decreased Au dispersion.

**CONCLUSIONS**

Single-site Au/C catalysts, prepared by impregnation and using aqua regia Au solution, were studied by XAFS and INS under...
various reactant gas compositions to gain a greater understanding of the reaction mechanism and mode of catalyst deactivation. Sequential gas experiments showed that HCl in the absence of C\textsubscript{2}H\textsubscript{2} cannot reoxidize Au(I)−chloride species under the reaction conditions studied and that HCl addition requires concerted addition with C\textsubscript{2}H\textsubscript{2}. Exposure of the Au/C catalyst to C\textsubscript{2}H\textsubscript{2} (in the absence of HCl) resulted in a XANES spectrum that did not resemble Au(III) chloride, Au(I) chloride, or Au(0), while EXAFS analysis suggested that AuCl\textsubscript{x} species were retained but with distorted Au−Cl distances. INS studies of a Au/C catalyst exposed to C\textsubscript{2}H\textsubscript{2} showed that oligomeric acetylene species formed on the catalyst surface, which could act as potential precursors to carbon fiber formation and thus become a significant deactivation process.

Catalyst deactivation, after C\textsubscript{2}H\textsubscript{2} treatment or prolonged time-online studies with higher Au loadings, was found to be associated with the formation of metallic Au particles. The presence of Au(0) significantly affected the strong correlation between normalized white line height and VCM productivity seen in a completely single-site catalyst (i.e., in the absence of Au(0) particles). Care must be taken to consider other spectator species that may be present from catalyst preparation or evolve during the reaction. Simple descriptors of catalyst structure, such as white line height, while useful, must be carefully used in systems of increased complexity.

**EXPERIMENTAL SECTION**

**Catalyst Preparation.** One and 2 wt % gold supported on activated carbon catalysts were prepared by wet impregnation of the HAuCl\textsubscript{4} precursor dissolved in aqua regia (denoted, respectively, as 1 wt % Au/C-AR and 2 wt % Au/C-AR materials). Activated carbon (Norit ROX 0.8) was initially ground to obtain a 100–140 mesh powder. The gold precursor, HAuCl\textsubscript{4}·xH\textsubscript{2}O (Alfa Aesar, 99.9% (metals basis), Au 49%), was dissolved in aqua regia (3 parts by volume HCl [(Fisher, 32 wt %):1 part by volume HNO\textsubscript{3} [(Fisher, 70 wt %)])]. The gold precursor...
solution was then added dropwise with stirring to the activated carbon. Stirring was continued at ambient temperature until NO₃ production had subsided. The product was then dried for 16 h at 140 °C under an inert flow of nitrogen.

In Situ Powder X-ray Diffraction (XRD). Powder XRD data were acquired using an X’Pert Pro PAN Analytical powder diffractometer employing a Cu Kα radiation source operating at 40 kV and 40 mA. Analyses of the spectra were carried out using X’Pert High Score Plus software. The mean crystallite size of the metallic gold nanoparticles was determined using the Scherrer equation assuming a spherical particle shape and a K factor of 0.9 on the reflection arising from the {111} Au planes (at 2θ = 38°).

In Situ X-ray Absorption Fine Structure (XAFS). XAFS spectra were recorded in transmission mode at the Au L₁ absorption edge, at the B18 beamline of the Diamond Light Source, at Harwell, UK. The measurements were performed using a QEXAFS setup with a fast-scanning Si (111) double-crystal monochromator. For the in situ measurements, the time resolution of the data acquisition was 20 s per spectrum.

X-ray absorption near-edge structure (XANES) analysis was mainly focused on understanding the details of the white line feature detectable at an absorption energy of ~11 920 eV, which corresponds to the Au 2p₃/2 → 5d primary transition. Three different Au standards were used to perform a linear combination fitting (LCF) analysis of the Au L₁-edge XANES, namely, KAuCl₄/[AuCl₅]⁻ for Au(III), [AuCl₄]⁻ for Au(I), and a bulk Au-foil for Au(0). The normalized white line intensity values observed for the [AuCl₄]⁻ and [AuCl₅]⁻ standards were 1.1 and 0.6, respectively.

Analysis of the extended X-ray absorption fine structure (EXAFS) data was performed using IFEFFIT with the Demeter software package (Athena and Artemis). Firstly, Au paths were fitted at all k-weighted χ data using a k space window of 3–11 and an R window of 1.25–2.60. The Debye–Waller (2σ²) and the amplitude reduction (S²)2 factors were fixed during fitting of data at 0.0037 Å² and 0.75, respectively. These values were determined from fitting of KAuCl₄ at a known temperature and with a fixed coordination number (CN) of 4.

Inelastic Neutron Scattering (INS). For INS analysis, the required amount of 2% Au/C-AR catalyst was loaded into a Conflat-sealed stainless steel cell and then dried at 200 °C under vacuum. INS spectra of the dried catalyst were recorded with the MERLIN spectrometer at the ISIS Facility using incident energies of 600, 250, and 100 meV to observe the C–H/O–H stretch, in-plane C–C stretches/C–H bends, and the out-of-plane deformations, respectively. These spectra were used as backgrounds for all subsequent spectra. The catalyst was then dosed with C₆H₆ (5%)/He at 200 °C. After measurement of the spectra, the catalyst was then dosed with 100% C₂H₂ at 200 °C and the spectra measured. All of the spectra were recorded at <20 K to minimize the Debye–Waller factor.

Computational studies of model systems were carried out using the plane-wave pseudopotential code CASTEP. Isolated molecule calculations were carried out with the molecule in the center of either a 10 × 10 × 10 Å or a 15 × 15 × 15 Å box. To investigate the effect of the surface, the same molecules were placed above a single graphene layer with a 10 Å vacuum gap. The Perdew–Burke–Ernzerhof (PBE) functional was used; the plane-wave cutoff was 750–1000 eV. For the isolated molecules only the Γ point was included; for the surface calculations a Monkhorst–Pack grid of 8 × 8 × 1 (12 k points) or 9 × 9 × 1 (20 k points) was used. The calculations were converged to better than ±0.003 eV Å⁻¹. After geometry optimization, the vibrational modes were calculated using density functional perturbation theory as implemented in CASTEP. The CASTEP output includes the vibrational transition energies and the atomic displacements of the atoms in each mode. The latter enables visualization of the modes using Materials Studio and is also the input to the program AbINS that creates simulated INS spectra. AbINS can generate the full 2D S(Q,ω) map for comparison with the output of direct geometry INS spectrometers such as MERLIN. Cuts through the data assume constant resolution and have the correct Q dependence and treatment of the Debye–Waller factor so that comparison with experimental data is both straightforward and reliable.

Scanning Transmission Electron Microscopy (STEM). Materials for examination by STEM were dry dispersed onto a holey carbon TEM grid. These supported fragments were examined using BF- and HAADF-STEM imaging mode in an aberration-corrected JEOL ARM-200CF scanning transmission electron microscope operating at 200 kV. This microscope was also equipped with a Centurio silicon drift detector (SDD) system for X-ray energy-dispersive spectroscopy (XEDS) analysis.

Catalytic Testing Experiments: Activity and Deactivation Measurements. The 1 wt % and the 2 wt % Au/C-AR catalysts were tested using a completely automated reactor system with the same setup as previously described. All of the predilute gases 5% C₂H₂/Ar (BOC), 5% HCl/Ar BOC, and Ar (99.99% BIP, Air Products) were dried using moisture traps before being introduced into the reactor. In all cases the reactor was heated to 200 °C at a ramp rate of 5 °C/min and held at this temperature for 30 min under a flow of argon prior to admitting the hydrochlorination reaction mixture. The tests were performed using a fixed-bed polyimide (Kapton) microreactor containing the catalysts, keeping the total flow of 50 mL min⁻¹ and a total gas hourly space velocity (GHSV) of ~14 000 h⁻¹. When both reactants were present, the C₂H₂:HCl ratio was kept at a constant value of 1.1:0.2.
The sequential flow experiment (reaction sequence 1) was performed using the 1% Au/C-AR catalyst while simultaneously monitoring the Au L3-edge XAFS and catalytic activity.

Reaction sequence 1 employed the following gas compositions: step i = HCl/C2H2/Ar, step ii = HCl/Ar, step iii = HCl/C2H2/Ar, and step iv = C2H2/Ar, and step v = HCl/C2H2/Ar.

The duration of each step in the sequence was not the same. The gas composition during the experiment was altered only when no further change in the XAFS spectra was observed. For this reason, the duration of each step in the reaction sequence was as follows: step i = 240 min, step ii = 75 min, step iii = 120 min, step iv = 70 min, and step v = 90 min.

The reaction mixture was analyzed online by mass spectrometry (Hiden QGA), and Professional Edition software was used for both qualitative and quantitative analyses. The catalyst activity presented is shown in terms of productivity toward VCM. The results are presented in terms of acetylene conversion (%).

The reaction mixture was analyzed online by mass spectrometry and identification of the products was carried out using a Varian 450 gas chromatograph equipped with a flame ionization detector (FID). Chromatographic separation and identification of the products was carried out using a Porapak N packed column (6 ft x 1/8” stainless steel).

The reaction mixture was analyzed online by mass spectrometry (Hiden QGA), and Professional Edition software was used for both qualitative and quantitative analyses. The catalyst activity presented is shown in terms of productivity toward VCM.

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**REFERENCES**


(12) Ozkaraoglu, E.; Tunc, I.; Suzer, S. Preparation of Au and Au-Pt Nanoparticles within PMMA Matrix Using UV and X-Ray Irradiation. Polymer 2009, 50 (2), 462−466.


(29) Bertus, P.; Fécourt, F.; Bauder, C.; Pale, P. Evidence for the in Situ Formation of Copper Acetylides during Pd/Cu Catalyzed