Nanoporous Gold as a Highly Selective and Active Carbon Dioxide Reduction Catalyst

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Supporting Information

ABSTRACT: Electrochemical conversion of CO2 into useful chemicals is a promising approach for transforming CO2 into sustainably produced fuels and/or chemical feedstocks for industrial synthesis. We report that nanoporous gold (np-Au) films, with pore sizes ranging from 10 to 30 nm, represent promising electrocatalytic architectures for the CO2 reduction reaction (CO2RR) due to their large electrochemically active surface area, relative abundance of grain boundaries, and ability to support pH gradients inside the nanoporous network. Electrochemical studies show that np-Au films support partial current densities for the conversion of CO2 to CO in excess of 6 mA cm−2 at a Faradaic efficiency of ∼99% in aqueous electrolytes (50 mM K2CO3 saturated with CO2). Moreover, np-Au films are able to maintain Faradaic efficiency greater than 80% for CO production over prolonged periods of continuous operation (110 h). Electrocatalytic experiments at different electrolyte concentrations demonstrate that the pore diameter of nanoporous cathodes represents a critical parameter for creating and controlling local pH gradients inside the porous network of metal ligaments. These results demonstrate the merits of nanoporous metal films for the CO2RR and offer an interesting architecture for highly selective electrocatalysis capable of sustaining high catalytic currents over prolonged periods.

KEYWORDS: CO2 reduction, nanoporous cathode, pH gradient, grain boundaries, electrocatalysis

The ability to reduce CO2 into useful chemicals or fuels will not only enable clean technology but also close the carbon cycle by recycling CO2 and preventing its further addition to the atmosphere.1 The CO2 reduction products can either be liquid fuels such as ethanol or gaseous products such as syngas (H2 and CO), which are feedstocks for thermocatalytic transformations via the Fischer–Tropsch process.2–5 To date, CO2 reduction is not a widespread technology because of low energy efficiency associated with high overpotentials, a lack of electrocatalytic stability, and poor selectivity for the CO2 reduction reaction (CO2RR) over the H2 evolution reaction (HER), which results in low partial current densities for the product of interest.6 Various approaches have been explored to improve the activity and selectivity of Au-based electrocatalysts for the CO2RR, from controlling nanocrystal size to tailoring of the exposed crystal facets, and even surface functionalization with molecular coatings.7–12 Recently, nanoporous catalytic architectures have shown promise for electrochemical CO2 reduction due to their large internal surface area and prevalence of stepped sites and grain boundaries inherent in their complex structure of highly curved metal ligaments.13–19 This propensity for under-coordinated atomic sites has been suggested to play a pivotal role in improving the selectivity of CO2 reduction in nanoporous silver (np-Ag) cathodes by stabilizing CO2− intermediates involved in the electrochemical conversion of CO2 to CO.18,19 Similar mechanisms have been invoked to explain the electrocatalytic performance of nanoporous gold (np-Au) films.18,19 While the highly irregular surface atomic structure of np-Au is well-known, relatively less
attention has been devoted to exploring how molecular transport into and out of such a tightly confined catalytic system may also affect the selectivity for CO$_2$ reduction within the porous network. Mesoporous Au films with controlled pore sizes around 200 nm in diameter have previously been shown to exhibit increased selectivity for CO$_2$ reduction with increasing film thickness from 0.5 to 2.7 μm. The improved selectivity is attributed to the formation of a pH gradient within the porous network as protons are consumed during electrolysis faster than they can be replenished by the supporting electrolyte; this effect is increased with increasing thickness of the mesoporous metal cathode. Although this study only adjusted the overall film thickness, these observations strongly suggest that fine-tuning the metal porosity by controlling the pore size could offer a simple route to further improving the selectivity of porous cathodes for electrochemical CO$_2$ reduction in aqueous electrolytes.

Here, we use np-Au films with pore diameters on the order of tens of nanometers to explore the influence of metal porosity on the selectivity for CO$_2$ reduction in aqueous electrolytes. Due to their small pore diameters, the porous network of metal ligaments is able to sustain pH gradients within np-Au films that are half as thick (∼800 nm) as those previously reported in mesoporous Au films (∼2 μm). This effect becomes more prominent upon further decreasing the pore diameter from ∼30 to ∼10 nm, as evidenced by electrochemical studies. We find that np-Au films are highly selective for the conversion of CO$_2$ to CO with high Faradaic efficiency (FE ∼ 99%) at modest overpotentials (η = 0.40 V), while at the same time delivering large current densities for CO (J$_{CO} = 6.2$ mA cm$^{-2}$). Finally, we demonstrate that these np-Au films exhibit excellent electrochemical durability and maintain Faradaic efficiency of ∼80% for CO production over 4.5 days of continuous electrolysis at an applied potential of $E = −0.5$ V vs the reversible hydrogen electrode (RHE).

The np-Au films were fabricated by electron-beam deposition of Ag and Au alloys onto clean silicon (Si) substrates, followed by selectively etching Ag from the Ag/Au alloy with nitric acid (see Experimental Methods in the Supporting Information). Briefly, 2 nm of titanium (Ti) was initially deposited onto Si as an adhesion layer, followed by 50 nm of Au as a planar, nonporous base layer to support the np-Au structure and to ensure that the Ti is not exposed to the electrolyte. The Au/Ag alloy (25/75 (vol %)) was then co-deposited in the electron beam, ranging in thickness from 100 nm to 2 μm. The np-Au morphology was then obtained by etching the Au/Ag alloy films in 70 wt % nitric acid for 10 min at room temperature (∼22 °C; denoted RT np-Au) or at lower temperatures (∼20 °C; denoted LT np-Au) in a freezer. Secondary ion mass spectrometry (SIMS) indicates that approximately 1.3 at. % of residual Ag remains in the structure after etching, consistent with prior reports. Figure 1 shows helium focused ion beam (He FIB) images of np-Au samples that were etched at room temperature (Figure 1a) and at low temperature (Figure 1b), displaying average ligament thicknesses of 28 ± 8 and 10 ± 2 nm, respectively. Chemical etching at low temperatures restricts the surface mobility of the Au atoms during etching and ensures that the ligament diameter is decreased. A scanning electron microscope (SEM) cross-section image of the ∼800 nm thick RT-etched np-Au film shows that the entirety of the film is porous down to the planar Au base layer (Figure 1c). As shown in this cross-sectional image, we routinely observed that the fully etched np-Au films were approximately 20% thinner (∼800 nm) than the initial thickness of the Au/Ag alloy (∼1 μm). To characterize the electrochemical surface area of the np-Au films, we performed Cu underpotential deposition (UPD) experiments (see Experimental Methods in the Supporting Information) and obtained a maximum roughness factor ∼57 for the thickest films. It is also important to note that the surface area increases linearly with film thickness, indicating that the entire surface area of the np-Au film is electrochemically accessible (Figure 1d).

To estimate the average grain size of the np-Au films, we performed X-ray diffraction (XRD) on a RT-etched sample of ∼800 nm thickness (Supporting Information Figure S1a). For comparison, we also examined a 50 nm thick planar Au base layer and a commercial Au foil (Alfa Aesar, 99.9975%; Figure S1b,c). These data show that the average full width at half-maximum of the diffraction peaks from np-Au are larger than those of the Au foil (Figure S1d). According to the Scherrer equation, the np-Au film and Au foil have average grain sizes of 20 ± 4 nm and 77 ± 23 nm, respectively. These calculations assume a shape factor of unity and do not take into account the possibility of microstrain. We also performed transmission electron microscopy (TEM) to directly visualize the distribution of grain boundaries within individual ligaments of the np-Au film. Consistent with prior reports, we observed many grain boundaries along the surface of the curved Au ligaments (Figure S2). Dark-field TEM images were also collected to estimate the average grain size within the np-Au films.
Au film (Figure S3 and Figure S4). The average grain size that we observed in TEM (17 ± 4 nm) is very similar to the estimate obtained through analysis of the XRD data. It is known that grain boundaries and under-coordinated sites in Au electrocatalysts serve as the active sites for CO₂ reduction, suggesting that these np-Au films should exhibit substantial activity for CO₂ reduction.

A two-compartment electrochemical cell was used to evaluate the electrocatalytic properties of the np-Au films for the CO₂RR (see Experimental Methods in the Supporting Information). An anion exchange membrane separated the Pt foil anode from the np-Au cathode in 50 mM K₂CO₃ electrolyte saturated with dissolved CO₂. It is known that higher supporting electrolyte concentrations provide higher current densities during electrolysıs, 30–32 but we found this concentration sufficient to enable reliable evaluation of our electrodes. The reference electrode was a leakless Ag/AgCl electrode; all potentials are reported relative to RHE to aid comparison with literature. The cathode compartment was bubbled with CO₂ at a flow rate of 5 cm³(STP) min⁻¹ through a glass dispersion frit to maximize the delivery of CO₂ into solution during controlled potential electrolysis. The effluent gas was sent directly into a gas chromatograph (SRI instruments) to analyze the chemical products.

Figure 2a shows the FE of a RT np-Au film (661 ± 10 nm thick) for both CO (filled blue bars) or H₂ (white bars) as a function of the applied potential (E) from −0.3 to −1.1 V_RHE (V vs RHE). Each data point shown in Figure 2 represents the average FE for CO or H₂ obtained over 2–3 h of continuous electrolysis at the indicated potential with IR compensation. All data were obtained from the same electrode along the potential sweep. The RT np-Au film exhibits a maximum FE for CO of 90% at E = −0.5 V_RHE with the remainder of the current producing H₂. We note that no liquid products were detected for any of the Au electrodes studied. Notably, the LT np-Au film (664 ± 5 nm thick) obtains a maximum FE for CO (filled gray bars) of 99% at E = −0.5 V_RHE and maintains at least 80% FE for CO from −0.3 to −0.7 V_RHE before the HER (white bars) begins to account for a larger portion of the products at more negative applied potentials (Figure 2b).

To examine the influence of the np-Au morphology on CO₂ reduction selectivity, we tested the activity of a 50 nm thick planar Au film, which is the base Au layer of the np-Au electrodes. As shown in Figure 2c, the planar Au film primarily produces H₂ (white bars) across the entire potential window; the FE for CO production (green bars) only reaches ~40% at −0.5 V_RHE. We also evaluated the activity of a commercial Au foil (Alfa Aesar, 99.9975%) to confirm that our experimental conditions and cell configuration are capable of adequately reproducing commonly observed activity trends for Au films. As shown in Figure 2d, the Au foil obtained a maximum FE for CO of 92% at −0.5 V_RHE (filled red bars), consistent with prior reports. A significant advantage of the np-Au morphology over the planar Au electrodes is illustrated by the high partial current density for CO (J_CO) relative to H₂ (Figure 2e–h). At an applied potential of −0.7 V_RHE the LT np-Au film exhibits a peak J_CO of 8.1 mA cm⁻² (Figure 2f), which is four times higher than the Au foil (Figure 2h) and eight times higher than the planar Au film (Figure 2g). At the optimum applied potential for CO production (~0.5 V_RHE), the LT np-Au film displays J_CO of nearly 6.2 mA cm⁻², while the RT np-Au film J_CO is around 4.5 mA cm⁻². Interestingly, the LT np-Au film shows only a slight increase in J_CO as compared to that of RT np-Au despite the ~3 times increase in surface area between the LT and RT np-Au films (Figure S5a). This lower than expected J_CO from LT np-Au films likely arises due to mass transport limitations, whereby the geometry of the electrochemical cell does not allow for sufficient delivery of CO₂ throughout the porous electrode to keep up with the
Figure 3. Influence of electrolyte concentration on CO₂ reduction selectivity with Au cathodes. (a–c) Faradaic efficiency for CO as a function of applied potential (E) obtained at two different electrolyte concentrations (both saturated with CO₂) for (a) 809 ± 15 nm thick RT-etched np-Au film, (b) 821 ± 22 nm thick LT-etched np-Au film, and (c) 50 nm thick planar Au film. (d–f) Corresponding average current density (J) obtained at E observed at two different electrolyte concentrations for (d) RT np-Au, (e) LT np-Au, and (f) planar Au film. (g–i) Predicted solution pH at the surface of the electrode for (g) RT np-Au, (h) LT np-Au, and (i) planar Au film. A planar electrode geometry is assumed for the simulations.

electrochemically active surface area of the entire film. This hypothesis is confirmed by comparing the FE and Jₖₒ for RT-etched samples that were ~800 and ~150 nm thick (Figure S6). Despite the drastic difference in electrochemical surface area as determined by Cu UPD (Figure 1d), the thicker film only showed a ~30% increase in Jₖₒ under our experimental conditions (Figure S6c). Note that the linear relationship between the surface area of the np-Au film and alloy thickness (Figure 1d) implies that the entire network is accessible to the electrolyte, suggesting that a large fraction of dissolved CO₂ does not penetrate the entire depth of the film at the current densities tested. These results suggest that to better use the full electrocatalytic surface area of np-Au for CO₂ reduction requires that the geometry of the cell be modified to flow the CO₂ directly through the np-Au film so that CO₂ is efficiently delivered to the catalyst, as opposed to simply flowing the CO₂ past the electrode surface. Indeed, it has recently been shown that such a tactic is highly beneficial for improving the rate of electrochemical CO₂ reduction.²⁴,³⁵

Both the RT and LT np-Au films exhibit superior FE for CO (Figure 2a,b) relative to the planar Au film or Au foil (Figure 2c,d) across the entire potential window studied. It has previously been shown that the residual Ag in the np-Au film is not the source of the high FE for CO.¹⁸ We therefore attribute such significant improvements in catalytic selectivity to the prevalence of grain boundaries that exist within the np-Au structure relative to the planar Au film. Another factor that likely contributes to such marked improvements in selectivity is the ability of the np-Au film to support locally alkaline pH conditions within the porous network as protons are consumed during electrolysis. Such an effect has previously been observed in mesoporous Au electrodes, which serves to suppress the rate of HER while the rate of CO₂ reduction is relatively unaffected.²⁰

To examine the influence of local pH gradients within the nanoporous network on the selectivity of np-Au films, we examined the electrochemical activity of a RT and a LT np-Au film (Figure 3d,e) at two different electrolyte concentrations (50 mM K₂CO₃ and 200 mM K₂CO₃—both fully saturated with CO₂). Increasing the electrolyte concentration increases the buffering capacity of the solution,²⁰,³² which reduces any swings in local pH that are anticipated to form within the pores of the np-Au films as protons are consumed during electrolysis. It was therefore anticipated that the np-Au films would exhibit reduced FE for CO in 200 mM K₂CO₃ electrolyte if an increased solution pH within the porous network was responsible for the high selectivity observed on the np-Au films. As shown in Figure 3a,b, the selectivity on both RT np-Au and LT np-Au is essentially unchanged at low applied potentials (~0.3 V₉ₑₙ₉ₑₙₑ) but as the current density increases with increased applied bias (Figure 3d,e), any pH gradient that may form within the np-Au film in the 50 mM K₂CO₃ electrolyte (Figure 3a,b (circles)) is diminished by the improved buffering capacity of the 200 mM K₂CO₃ electrolyte (Figure 3a,b (diamonds)). In contrast, no change in FE for CO is observed on a planar Au film at any applied potential (Figure 3c), confirming that the change in selectivity observed on the np-Au electrodes is not simply a consequence of the increased
electrolyte concentration (Figure 3c,f). These results strongly suggest that a pH gradient is forming within the porous network of the Au electrocatalyst and serves an important role in achieving high selectivity of CO₂ reduction in aqueous electrolytes.

Interestingly, the LT np-Au film shows a larger reduction in FE for CO (Figure 3b) than the RT np-Au film (Figure 3a). Analysis of the pore sizes between these two samples reveals that the LT np-Au film has pores with an average diameter of 10 ± 2 nm, while the RT np-Au sample has pores with an average diameter of 28 ± 8 nm (Figure S5b). We therefore attribute the improved selectivity of the LT np-Au film relative to the RT np-Au film to the smaller pore size of the former (~10 nm) relative to the latter (~30 nm), which more effectively supports a high local pH within the porous network that improves the selectivity for CO₂RR relative to HER (Figure 2a,b). Previous work has shown that increasing the thickness of a mesoporous Au catalyst to ~2 μm helps to achieve a similar effect within ~200 nm pores.²⁰ Our observations suggest that further reducing the pore volume below 10 nm may enable realizing such an effect within even thinner nanoporous metal films than those studied here.

To further explore whether the pH gradient is developed within the porous Au network or occurs largely in the boundary layer, we simulated the pH profile as a function of distance away from the electrode surface using the model previously reported by Gupta et al.²⁰ Briefly, the model assumes a planar electrode geometry, which is a valid assumption for calculating the pH at the outer surface of the electrode because the flux of reactants and products must be the same for either a porous or planar electrocatalyst at this location. The assumption of a planar electrode is clearly incapable of accounting for changes in the transport of reactants and products into and out of the porous film itself, and we therefore interpret any experimental deviations from the model to originate from changes occurring within the porous network of metal ligaments. The inputs into the model are the electrolyte concentration, the total current density, and the Faradaic efficiency for CO and H₂. A 70 μm thick boundary layer was assumed based on the experimental flow rate of CO₂ of 5 cm³(STP) min⁻¹ through the catholyte.³¹ As shown in Figure 3g–i, these calculations predict very little change in local pH at the electrode surface between the two electrolyte concentrations, albeit small deviations from the bulk electrolyte are predicted for the RT np-Au film and the planar Au electrode (Figure 3g,j). While significant reductions in FE for CO were observed on both the RT and LT np-Au films (Figure 3a,b, respectively), no change in FE was observed experimentally on the planar Au film (Figure 3c). This obvious contradiction between the results of experiment with those from the model indicates that the local pH changes must be occurring within the porous network itself. Otherwise, we would have observed a similar reduction in FE for CO with the planar electrode at the higher electrolyte concentration. We note that these experimental observations are consistent with a previous report on mesoporous Au films,²⁰ yet are achieved with much thinner films. Taken together, these results indicate that the pore diameter of porous metal electrocatalysts is a critical parameter for optimizing their selectivity and suggest that control over the pore size on the nanometer length scale may offer further improvements in electrochemical selectivity.

We further evaluated the electrocatalytic stability of these Au films for the CO₂RR at an applied potential of $E = −0.5 \, V_{\text{RHE}}$ (with iR compensation). Significantly, we observed that the np-Au film maintained a high FE for CO (nearly 90%) over the course of 24 h of continuous electrocatalytic testing (Figure 4a). In stark contrast, the Au foil and planar Au films exhibit drastic reductions in FE for CO over just 1 day of testing at the same applied potential (Figure 4b,c). Continued testing of a different RT np-Au film for 4.5 days (110 h) showed continued catalytic stability (Figure S7). Comparison of the SEM images before and after testing show no significant changes in morphology except that the np-Au ligaments appear to coarsen slightly (Figure S8). Analysis of these films by XRD indicates no significant changes in peak width before and after testing, but all films showed a decrease in the overall signal magnitude from diffraction peaks associated with high-index reflections (Figure S9). We note that the activity of the Au foil can be recovered if the flame annealing treatment is repeated, but such a process is undesirable as it hinders long-term continual operation under CO₂RR conditions. These observations serve to highlight the benefit of using the nanoporous metal structure to perform CO₂ reduction: the prevalence of grain boundaries offers numerous active sites on the metal ligaments, while the porous network is able to support a locally alkaline
ph in the film that helps improve electrocatalytic selectivity for the CO₂RR over the HER.

In conclusion, we have demonstrated that np-Au films constitute a promising electrocatalytic architecture for CO₂ reduction to yield CO in aqueous electrolytes. The np-Au films exhibit a maximum Faradaic efficiency for CO of 99% at −0.5 V \text{RHE} while operating at a partial current density for CO in excess of 6 mA cm⁻². We attribute the catalytic performance of np-Au to its high electrochemical surface area possessing a large number of grain boundaries and its ability to support a local depletion of protons within the porous network. Significantly, these np-Au films maintain a Faradaic efficiency of greater than 80% over the course of 110 h of continuous electrolysis at −0.5 V \text{RHE}, while the activity and selectivity of both planar Au films and Au foils diminishes significantly over much shorter periods of operation (~4 h). These studies highlight the benefits of nanoporous metal cathodes for CO₂ reduction and indicate that the pore size is an important parameter to control for improving selectivity in these promising electrocatalytic architectures.

**REFERENCES**


