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Transition metal salen complexes (Co, Ni, Cu) are studied as precatalysts for CO₂-to-CO electroreduction in zero-gap electrolyzers. Performance depends strongly on metal choice and electrode characteristics. XPS, UV-Vis, comparisons to simple metal salts, and a RHE-ZGE half-cell setup shed light on precatalyst decomposition and inform methodologies for molecular electrocatalyst heterogenization.

With the current shift towards more sustainable synthetic pathways, CO₂ electrolysis has gained attention as a promising technology. Beyond established Ag-, Au-, and Zn-based catalysts, numerous molecular systems have demonstrated high faradaic efficiencies for CO₂-to-CO conversion.^{1,2} However, the majority of catalysts conceptualized in lab environments fail to make the transition into industrial reactors.³ Bridging this “Valley of Death” for new electrolytic materials, as previously described by our group, requires an application-oriented strategy that couples catalyst development with process optimization in scalable devices, like zero-gap electrolyzers (ZGEs), that enable CO₂ electrolysis at elevated current densities.⁴ For molecular electrocatalysts in particular, the tools available to achieve this transfer to industrially relevant ZGEs remain highly limited.^{2,5} We thus set out to identify key experimental considerations for incorporating molecular electrocatalysts into gas diffusion electrodes (GDEs) for CO₂ reduction in ZGEs. We chose to focus our research on salen complexes, which are affordable, modular, compatible with a wide range of non-precious metals, and have shown good CO₂ electroreduction activity in both homogeneous and heterogeneous reaction environments.^{5–10} Specifically, we aimed to: (1) study how ink

Integrating salen complexes into gas diffusion electrodes for CO₂ electroreduction: considerations for employing molecular precatalysts in heterogeneous electrolyzers

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preparation and operating conditions affect CO₂R activity to find optimized process parameters, (2) determine the influence of different metal centers (Co, Ni, Cu) and salen ligands on CO₂R performance under those optimized process parameters, and (3) use pre- and post-electrolysis surface XPS to identify possible salen complex decomposition occurring after drop-casting or after electrolysis.

The three salen ligands used in this research are illustrated in Fig. 1a. All salen complexes of Co^{II}, Ni^{II}, and Cu^{II} used herein are described by the symbol of the metal and the name of the salen ligand used; for instance, the salen complex with a Co center using ligand L² is described as “CoL².“ All GDEs were prepared by directly adding solid catalyst, powdered carbon

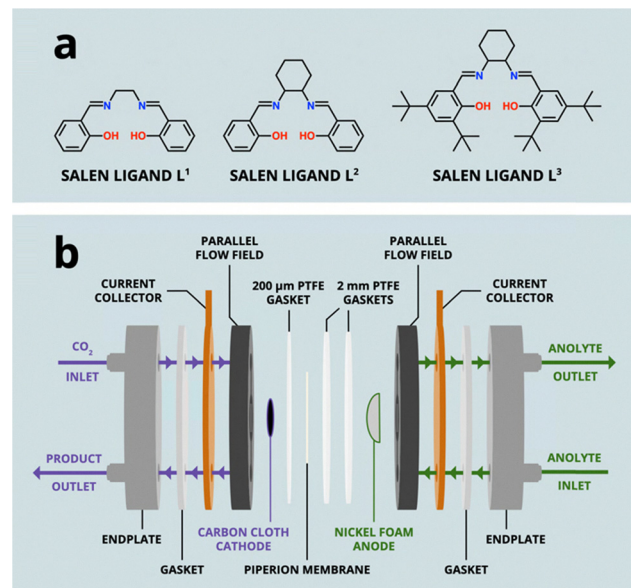


Fig. 1 (a) The salen ligands (L¹, L², and L³) used in this work. (b) Diagram of the internal components of the zero-gap electrolyzer (ZGE) used in this research, including the gas diffusion electrode (carbon cloth cathode).

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black, and PTFE nanoparticles (where noted) to a vial, sonicating in ethanol or THF, adding Sustainion® XA-9 alkaline ionomer, and then dropcasting the resulting ink onto a carbon cloth electrode at 70 °C until a catalyst loading of 0.45 mg cm⁻² was achieved. These electrodes were then used as cathodes inside a ZGE (design shown in Fig. 1b). During operation, an aqueous 1 M KOH anolyte solution was circulated through the anode side of the ZGE, while CO₂ was passed through a 2 cm² parallel flow field on the cathode side employing a PiperION® anion-exchange membrane (AEM) at room temperature, as previously reported.^{11–13} After an operating period of 30 minutes, any gaseous products (CO, CH₄, C₂H₄, and H₂) generated by the cell were analyzed by an in-line GC system. Further details regarding equipment and procedures can be found in the SI.

The commercially available cobalt salen complex CoL² was chosen for process parameter optimization experiments, as we expected that the hydrophobic and electronic properties of ligand L² would represent a reasonable medium of the hydrophobic and electronic properties of ligands L¹ and L³. Preliminary experiments were conducted at a current density of 100 mA cm⁻² with either humidified or non-humidified CO₂ feeding the cathode (Table 1a). Our experiments with non-humidified CO₂ showed a slightly higher average FE_{CO} value

Table 1 Results from process parameter optimization experiments with CoL². Unless otherwise indicated, all experiments were run with dry CO₂ at a current density of 50 mA cm⁻². Each reported FE_{CO} value is the average of two separate trials, and the 95% C.I. from the mean is provided. Values of FE_{H₂} and total FE values for these experiments, as well as further experimental details, are available in the SI (Table S2). All CO₂ humidification experiments were run at 100 mA cm⁻²

(a) CO ₂ Humidification (% relative humidification)	FE _{CO} (%)
Dry CO ₂ (0%)	3.68 ± 0.57
Humidified CO ₂ (100%)	1.339 ± 0.081
(b) Amount of PTFE nanoparticles added (mg cm ⁻²)	FE _{CO} (%)
0.0	10.4 ± 2.5
0.5	10.5 ± 1.2
1.0	8.0 ± 1.2
2.5	8.48 ± 0.85
5.0	10.7 ± 3.6
(c) Type of carbon black added	FE _{CO} (%)
Ensaco® 250G	10.4 ± 2.5
Super P® Li	7.06 ± 0.40
C-Nergy® Super C65	6.8 ± 1.7
Vulcan® XC-72R	4.8 ± 1.6
Multi-walled CNTs (Fe-free)	4.128 ± 0.083
(d) Carbon-to-catalyst weight ratio	FE _{CO} (%)
0.5	10.4 ± 2.5
1.0	9.0 ± 1.3
2.0	7.8 ± 4.0
(e) Current density (mA cm ⁻²)	FE _{CO} (%)
10	41 ± 10
50	10.4 ± 2.5
100	3.68 ± 0.57

of 4% – with the remaining amount of charge being consumed in parasitic hydrogen evolution, as verified by GC analysis (Table S2) – while experiments with humidified CO₂ generated an FE_{CO} of only 1%. This led us to suspect that moisture accumulation at the cathode might hinder CO₂R activity. Even when non-humidified CO₂ is used, the GDE is still wetted by water transport through the AEM, so we sought to directly control moisture content at the cathode through the incorporation of hydrophobic materials. To this end, varying amounts of hydrophobic PTFE nanoparticles (0.5, 1.0, 2.5, and 5.0 mg cm⁻²) were added to the catalyst ink prior to dropcasting (Table 1b).¹⁴ Electrolyses at a lower current density of 50 mA cm⁻² showed that the amount of PTFE added does not affect FE_{CO}. Additionally, seeking to understand whether the carbon support added to the ink plays a role in CO₂R performance, we tested five different varieties of carbon supports (Ensaco® 250G, Super P® Li, C-Nergy® Super C65, Vulcan® XC-72R, and multi-walled carbon nanotubes) (Table 1c). Operating at 50 mA cm⁻², Ensaco® 250G gave the highest average FE_{CO} values, followed by Super P® Li alongside C-Nergy® Super C65. This trend is in line with previous results from our group on molecular Ag catalysts in which the hydrophobic and graphitic character of Ensaco® 250G appears to promote CO₂R.^{9,12} Variations in carbon porosity and particle size may influence the formation of distinct microenvironments with locally altered water activity, potentially facilitating decomposition and hydroxylation of the salen complexes to other active species, as evidenced by our pre- and post-electrolysis XPS analyses (to be discussed later).¹⁵ The fact that CO₂ humidification and the type of carbon black added had a significant influence on observed CO₂R performance – while the addition of PTFE nanoparticles did not – suggests that local water content at the GDE is best controlled either at the carbon-catalyst interface or at the macroscopic interface through operational conditions, such as relative CO₂ humidification or temperature.

Further experiments were carried out with Ensaco® 250G in which the catalyst loading of CoL² was kept constant while the amount of carbon black was varied. 0.5, 1.0, and 2.0 equivalents of carbon black by weight relative to the catalyst were tested (Table 1d). While the obtained FE_{CO} values showed no significant dependence on the carbon-to-catalyst weight ratio, chronopotentiometry data from these trials showed that carbon-to-catalyst weight ratios of 1.0 and 2.0 often led to non-stabilized, elevated cell voltages of up to 4 V (Fig. S13–S17). This can be attributed to the increased amount of non-catalytically active carbon support (inactive towards both CO₂ reduction and water reduction) present in the GDE matrix. Decreasing the current density from 50 to 10 mA cm⁻² causes the FE_{CO} value of GDEs dropcast with CoL² to increase from 10% at 50 mA cm⁻² to 41% at 10 mA cm⁻² (Table 1e), where the change in FE_{CO} is again compensated for by a change in FE_{H₂}, as verified by GC analysis (Table S2). Despite the limited performance of CoL² for CO₂ reduction at current densities above 50 mA cm⁻², our results show how process parameters related to ink preparation and cell operation (CO₂ humidification, addition of PTFE nanoparticles, and the type and amount of carbonaceous support)



can impact faradaic efficiencies for CO production. More specifically, when screening heterogenized molecular electrocatalysts for CO₂ reduction on GDEs, we suggest employing non-humidified CO₂ streams and paying close attention to the variety and amount of carbon support incorporated into the catalyst ink, with more hydrophobic materials appearing the most promising.

Based on these process parameter optimization experiments, we chose to run all subsequent trials with non-humidified CO₂ at successive current densities of 10 mA cm⁻², 25 mA cm⁻², and 50 mA cm⁻² using a catalyst ink prepared with Ensaco[®] 250G in a carbon-to-catalyst weight ratio of 0.5 and without any PTFE nanoparticles added. Experiments were conducted with different salen ligands, providing varying hydrophobic and electronic effects, as well as different metal centers. The resulting FE_{CO} values are presented in Fig. 2a (at 10 mA cm⁻²) and Fig. S22–S28 (all current densities) for Co salen complexes CoL¹, CoL², and CoL³, Ni salen complexes NiL¹ and NiL³, and Cu salen complexes CuL¹ and CuL³. Ni and Cu complexes were not prepared with L², as this ligand (representing an intermediate between L¹ and L³ in terms of hydrophobic and electronic effects) was used only for process parameter optimization. CoL¹ displays a high faradaic efficiency for CO production (FE_{CO}: 91%) at a current density of 10 mA cm⁻², yet this FE_{CO} value drops precipitously with

increasing current density to 2% at 50 mA cm⁻². As described in the prior paragraph, CoL² follows a similar trend, giving a FE_{CO} value of 41% at 10 mA cm⁻² that decreases with rising current density, while CoL³ shows no specific activity for the CO₂R. Moreover, both Ni salen complexes are almost entirely inactive towards CO₂ electroreduction, with GDEs dropcast with NiL¹ only producing small amounts of CO, even at very low current densities (FE_{CO}: 2% at 10 mA cm⁻²) and NiL³ GDEs almost exclusively producing H₂. These findings are in line with the poor activity observed for a Ni porphyrin complex under nearly identical conditions.¹³ Notably, although CuL¹ and CuL³ show lower FE_{CO} values than CoL¹ and CoL² at 10 mA cm⁻², both Cu salen complexes display much smaller drops in their FE_{CO} values as current density increases compared to CoL¹ and CoL²: FE_{CO} values for CoL¹ drop by 97% when current density is increased from 10 to 50 mA cm⁻², whereas FE_{CO} values for CuL¹ drop by only 5% over the same increase in current density. CuL³ is the only complex that produces CH₄, with an FE_{CH₄} of 2% at 50 mA cm⁻² and 4% at 100 mA cm⁻²; no hydrocarbon products are observed for any of the other tested complexes.

Compared to reported systems involving molecular CO₂R electrocatalysts in heterogeneous electrolyzers (see Table S3), the salen complexes tested herein show limited activity for CO₂R on GDEs, despite promising CO₂R performance in homogeneous systems. As we have highlighted in a recent review,¹⁶ the incorporation of molecular electrocatalysts into ZGEs requires *in situ* spectroscopic studies to best identify the catalytically active species (if any) that are formed on the electrode before and after electrolysis.² To assess possible salen complex decomposition, XPS surface scans were collected of pre- and post-electrolysis GDEs dropcast with CoL², CoL³, NiL³, and CuL³ (Fig. S18–S20). Demetallation of the salen complex was often observed directly after preparation of the GDE: pre-electrolysis scans of CoL² and CoL³ (Fig. S18) both show appreciable formation of Co(OH)₂, and a pre-electrolysis scan of CuL³ (Fig. S20) shows formation of both Cu(OH)₂ and CuO.^{13,17–20} Suspecting that interactions with the Sustainion[®] XA-9 binder added to the catalyst ink (in its chloride form) may lead to pre-electrolysis decomposition of our metal salen complexes, we measured UV-Vis of CoL² sonicated in wet ethanol both in the absence and presence of binder (Fig. S21). No significant spectral changes were observed after adding binder, suggesting that pre-electrolysis demetallation of these complexes is a result of the heating and/or desolvation incurred during dropcasting at 70 °C. We also found that demetallation of the salen complexes can occur during electrolysis under alkaline conditions. A pre-electrolysis scan of a GDE dropcast with NiL³ shows that the salen complex remains fully intact after ink preparation and dropcasting, while the markedly different post-electrolysis scan shows Ni(OH)₂ as the majority species on the electrode (Fig. S19), which we attribute to the attack of hydroxide anions that have migrated through the AEM at the Ni^{II} center.^{13,17} These results highlight the general need for routine spectroscopic analyses (both before and after electrolysis) to identify the species actually formed on the cathode during GDE preparation and during electrolysis.²¹ In particular,

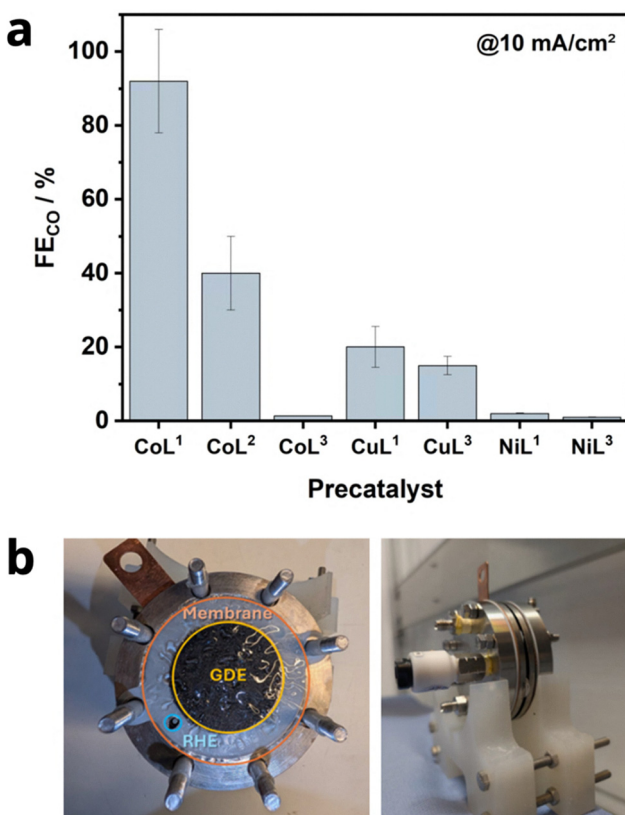


Fig. 2 (a) FE_{CO} values obtained at 10 mA cm⁻² after 30 min of electrolysis in a 2 cm² ZGE. Each reported FE_{CO} value is the average of two separate trials, and the 95% C.I. is provided in the form of error bars. (b) Photographs of the 12 cm² ZGE with an integrated RHE contact used for our half-cell potential experiments.



ensuring that a targeted molecular electrocatalyst is stable at elevated pH is a key consideration when working in alkaline electrolyzers, as metal complexes (like NiL^3) that are otherwise stable to ink preparation and dropcasting steps may react with hydroxide.^{15,22}

To better identify the catalytically active species actually formed on each GDE, we also performed CO_2R experiments with the simple metal salts Co(OAc)_2 , NiCl_2 , and Cu(OAc)_2 in the same metal loadings as their respective salen complexes (Fig. S29). Notably, the Ni and Cu salts outperform their respective salen complexes under identical ZGE conditions: at 10 mA cm^{-2} , we obtain FE_{CO} values of 18% for NiCl_2 (compare to 2% for NiL^1 and 0% for NiL^3) and 27% for Cu(OAc)_2 (compare to 19% for CuL^1 and 14% for CuL^3), suggesting that the metal salen complex is not the principal catalytically active species on GDEs dropcast with salen complexes of these metals. Therefore, we conclude that CuL^1 and CuL^3 primarily act as precatalysts that undergo decomposition to catalytically active Cu(OH)_2 or CuO species. GDEs dropcast with Co(OAc)_2 only evolve H_2 , suggesting that the salen ligand plays a role in either (1) guiding the decomposition of CoL^1 and CoL^2 to catalytically active Co(OH)_2 assemblies of a particular size or shape or (2) stabilizing the Co center such that the non-decomposed complex is catalytically active.

Finally, to expand the toolbox our work offers, we fabricated a ZGE with an integrated RHE (Fig. 2b), allowing for the accurate measurement of half-cell potentials at the cathode across different current densities. The resulting half-cell potentials are plotted alongside FE_{CO} values at 10, 25, 50, and 100 mA cm^{-2} in Fig. S24–S28 for NiL^1 , NiL^3 , CuL^1 , CuL^3 , and the highest performing Co salen complex, CoL^1 . We note that neither CuL^1 , CuL^3 , nor CoL^1 show significant voltage changes with increasing current density, suggesting the presence of conductive metallic particles performing the more kinetically favoured HER. With this approach enabling simultaneous product quantification and half-cell potential measurements in a scalable cell design, we offer the community a valuable tool for molecular electrocatalyst evaluation.

Herein, we evaluated Co, Ni, and Cu salen complexes as CO_2R precatalysts in industrially relevant ZGEs. The salen ligand is believed to play a role in the CO_2R activity of GDEs dropcast with Co salen complexes (either by directing the formation of catalytically active Co(OH)_2 assemblies or by stabilizing a catalytically active, non-decomposed complex). Ni salen complexes are almost entirely inactive for CO_2R , and Cu salen complexes appear to act as precatalysts that decompose to catalytically active Cu(OH)_2 or CuO species. FE_{CO} values were limited above 50 mA cm^{-2} , but we identify key experimental considerations for bridging lab-scale molecular electrocatalysis research and implementation in industrial electrolyzers: (1) control local water content on the GDE either at the carbon-complex interface or through CO_2 humidification, (2) apply *in situ* techniques (like surface XPS) to assess possible decomposition during ink preparation and/or electrolysis, and (3) compare GDEs dropcast with molecular complexes to those dropcast with simple metal

salts to help identify active species. We also present a modified ZGE with an integrated RHE for accurately measuring half-cell potentials during electrolysis, offering molecular electrochemists another tool for probing molecular catalyst behavior in scalable cells.

Conflicts of interest

There are no conflicts to declare.

Data availability

All data supporting the findings of this study are available within the article and its supplementary information (SI). Supplementary information is available. See DOI: <https://doi.org/10.1039/d5cc03236a>.

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