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Self-adaptive amorphous CoO_xCl_y electrocatalyst for sustainable chlorine evolution in acidic brine

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Electrochemical chlorine evolution reaction is of central importance in the chlor-alkali industry, but the chlorine evolution anode is largely limited by water oxidation side reaction and corrosion-induced performance decay in strong acids. Here we present an amorphous CoO_xCl_y catalyst that has been deposited in situ in an acidic saline electrolyte containing Co^{2+} and Cl⁻ ions to adapt to the given electrochemical condition and exhibits ~100% chlorine evolution selectivity with an overpotential of ~0.1 V at 10 mA cm⁻² and high stability over 500 h. In situ spectroscopic studies and theoretical calculations reveal that the electrochemical introduction of Cl⁻ prevents the Co sites from charging to a higher oxidation state thus suppressing the O-O bond formation for oxygen evolution. Consequently, the chlorine evolution selectivity has been enhanced on the Cl-constrained Co-O⁺ sites via the Volmer-Heyrovsky pathway. This study provides fundamental insights into how the reactant Cl⁻ itself can work as a promoter toward enhancing chlorine evolution in acidic brine.

Chlorine (Cl₂) gas is an essential chemical with an annual production of over 70 million tons for a wide range of applications in water treatment, organic synthesis, and polyvinyl chloride production¹⁻³. It is mainly produced through electrochemical chlorine evolution reaction (CER)⁴ in the chlor-alkali industry⁵⁻⁷, where the oxygen evolution reaction (OER) can compete with CER in aqueous electrolytes owing to their comparable onset potentials and similar types of catalytic active sites⁸⁻¹⁰, thereby it is a great challenge to ensure the highly selective production of Cl₂ gas. The state-of-theart industrial electrolyzers for CER generally use relatively acidresistant mixed metal oxides based on high-cost precious metal (Ru or Ir)^{11,12}, which are highly active for OER as well. To ensure the high efficiency in the CER system and improve the Cl₂ selectivity, a high molar concentration of Cl⁻ ion (5.0 M) and acidic pH conditions are required¹³⁻¹⁵. However, for practical applications, Ru-based CER catalysts are still insufficiently stable under long-term operation due to the formation of soluble ruthenium chloride¹⁶. Hence, some additives or dopants, such as the SnO_x, were used to improve the activity as well as stability of mixed metal oxides-based CER catalysts over the past decades¹⁷⁻¹⁹.

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The CER occurs under harsh reaction conditions according to the Pourbaix diagram of an aqueous saline electrolyte²⁰⁻²², especially at pH <3 with a highly polarized potential range, unavoidably leading to the dissolution of most metal oxide materials. Recently, low-cost nonprecious metal oxides, such as highly crystalline Co₃O₄ and transitionmetal antimonates²³⁻²⁶, have been tested as CER catalysts, which display decent stability since their dense, thick, and crystalline natures somehow prevent their complete dissolution in corrosive and acidic electrolytes. By contrast, an amorphous nonprecious metal oxide film can dissolve in such electrolytes in tens of minutes²⁷. As a result, the intrinsic electrochemical surface amorphization upon anodic polarization is very detrimental to the catalytic stability but technically inevitable. Despite the beneficial effect of enriching the surface-active sites through amorphization, it remains a huge challenge to exploit amorphous non-precious metal oxide catalysts, like the alkaline OER catalysts, toward highly active, selective, and sustainable CER.

In this work, we present a self-adaptive amorphous CoO_xCl_y catalyst, which was in situ deposited on a bare F-doped tin oxide (FTO) electrode at pH \leq 2 in saline solutions containing Co²⁺ and Cl⁻ ions. The final state of the deposited film catalyst depends on a given electrochemical condition, including the applied potential, solution pH, and electrolyte concentration²⁸. We show that CER takes place heterogeneously on the in situ deposited CoO_xCl_y catalyst. We find that an anodic polarization at \geq 1.67 V (versus RHE if not otherwise noted) leads to a moderate deposition rate of CoO_xCl_y and the Cl⁻ in electrolytes aids the co-deposition of Co and Cl into the catalyst film. Experimental results and density functional theory (DFT) calculations show that the introduction of Cl⁻ can share the oxidative equivalent under highly anodic potential and inhibit the oxidation of Co to a higher level for OER, leading to increased selectivity for CER. Online differential electrochemical mass spectrometry (DEMS) and rotating ring-disk electrode (RRDE) verified the efficient CER with ~100% selectivity in 0.5 M Cl⁻ brine solution. A certain time of catalyst deposition at 1.67 V leads to the CER activity increasing to around 10 mA cm⁻² and afterwards a slight increase over 500 h.

Results and Discussion

Screening out metal cations for CER in acidic brine

We first screened several transition metal cations, such as nickel (Ni), manganese (Mn), iron (Fe), copper (Cu), and cobalt (Co) at pH 2 for CER. We demonstrated that adding Cl⁻ ions into Mn, Ni, and Cucontaining brine solutions did not obviously shift the onset potential relative to the Cl-free electrolytes with the same ionic activity. Interestingly, a substantial decrease in the onset potential and an evident increase in current density were observed in the saline solution containing 0.001 M Co²⁺ or Fe³⁺ and 0.5 M Cl⁻ relative to that containing 0.001 M Co²⁺ or Fe³⁺ and 0.5 M ClO₄⁻ (Supplementary Fig. 1). However, although the Fe-catalyst displayed initial CER activity, the subsequent electrodeposition was not favorable, regardless of either increasing Fe³⁺ or Cl⁻ concentration thus being precluded (Supplementary Note 1 and Supplementary Figs. 2-6). In contrast to the large onset difference between CER and OER over deposited Co-catalyst, the dimensionally stable anode (DSA) and RuO2 catalysts loaded on FTO showed close onset potentials in 0.5 M NaCl electrolyte for CER relative to the Clfree electrolyte for OER (Supplementary Fig. 7). Meanwhile, we precluded the anion influence from Co precursors on cyclic voltammograms (Supplementary Fig. 8). In addition, we found that the current densities increase with the concentration of either Co²⁺ or Cl⁻ (Supplementary Fig. 9). The X-ray fluorescence (XRF) Co Ka mapping showed that the amount of deposited Co sites increases with the Cl concentration, suggesting the interplay between Co²⁺ and Cl⁻ (Supplementary Fig. 10).

The linear scan voltammetry (LSV) curve after 2 h of electrodeposition at 1.67 V in Cl-containing electrolyte was used as an example to exhibit a substantial increase in current density and a downshift of onset potential over 400 mV at 10 mA cm⁻² relative to that of Cl⁻free electrolyte (Fig. 1a and Supplementary Fig. 11). The in situ deposition of CoO_xCl_y at 1.67 V was accompanied by Cl₂ production (Supplementary Fig. 12), as well as reduced charge transfer resistance (Supplementary Fig. 13). As expected, a stepwise increase in potentiostatic time brought in an increased mass loading of the Co sites, which was verified by the time-dependent Co K α XRF test (Supplementary Fig. 14). Accordingly, the continuous increase of active Co sites led to a gradual attenuation of overpotential from -0.4 V after the first potential scan to finally -0.1 V (Fig. 1b). During this process, the catalyst film presents an epitaxial-growth-like behavior, of which the applied potential at the electrode/electrolyte interface can oxidize the Co²⁺ for the deposition of outermost catalyst layer until the interfacial potential drop across the catalyst^{29,30} cannot support the subsequent oxidation-deposition of Co²⁺.

Electrochemical CER selectivity and stability

Achieving high selectivity for CER is crucial due to the competing OER³¹⁻³³. Thus, we evaluated the CER selectivity based on the rotating ring-disk electrode (RRDE), where the generated Cl₂ on the disk catalyst electrode was quantified through electrochemical reduction on the ring electrode (Supplementary Fig. 15 and Supplementary Note 2). Through comparing the onset potentials for both O2 and Cl2 reduction on the Pt ring electrode based on the established method³⁴, 0.95 V has been applied for Cl₂ reduction and quantification (Supplementary Fig. 16 and Supplementary Note 3). The CER selectivity was evaluated by the chronoamperometry method at indicated potentials using RRDE³⁵ that was conducted in an Ar-saturated electrolyte containing 0.1 M Co²⁺ with various concentrations of Cl⁻ at pH 2 at 1600 RPM (Supplementary Figs. 17-21). Significantly, in acidic saline containing 0.1 M Co²⁺ and 0.5 M Cl⁻, a CER selectivity of ~100% at 1.67 V was achieved (Fig. 1c). It is worth noting that the Cl₂ selectivity already reached $90 \pm 1.5\%$ at 1.67 V when the Cl⁻ concentration is as low as 0.01 M. Importantly, in 0.5 M Cl⁻, the CER selectivity can easily achieve 100% at a wide range of potentials. These results are in line with the substantially reduced onset potential for CER in comparison to that for OER (Fig. 1a). Compared to the CoO_xCl_y catalyst, the selectivity of DSA (~80%) and RuO₂ (~88%) catalysts were distinctly low even in 0.5 NaCl electrolyte (Supplementary Fig. 22).

To further prove the CER selectivity, online DEMS was applied to discriminate the CER and OER³⁶⁻³⁸. The DMES experiment was implemented in a three-electrode H-cell in acidic saline solutions containing 0.1 M Co²⁺ and 0.5 M Cl (Supplementary Fig. 23). Significantly, as shown in Fig. 1d, the DEMS data clearly exhibit that the ⁷⁰Cl₂ gas was the only product in 0.5 M Cl electrolyte during the CER process, and no other signals were detected, such as ¹H¹⁶O³⁵Cl, ³⁵Cl³²O₂ or ³⁵Cl¹⁶O (Supplementary Fig. 24). In contrast, in the absence of Cl, the ³²O₂ was the exclusive product (Supplementary Fig. 25).

In addition, it is well accepted that, for non-noble metal oxidesbased CER catalysts, the activity decay is also a major issue in acidic electrolytes^{1,7}. A highly crystalline and thick Co₃O₄ catalyst film prepared through calcination was used as a control³⁹, and we found that its current density quickly declined from 5.0 to 0.87 mA cm⁻² (Supplementary Fig. 26). This activity decay resulted from the leaching/ dissolution of Co²⁺ from the surface of Co₃O₄. Similarly, the noble metal oxides-based DSA and RuO2 catalysts exhibited a fast descent in current density. In contrast, the current density of the CoO_xCl_y catalyst deposited at 1.67 V increases to 10 mA cm⁻² and then stabilizes at about 15 mA cm⁻² over 500 h test, which leads to a downshift of potential ~ 300 mV at 10 mA cm⁻² relative to the initial CV activity (Supplementary Fig. 27). Meanwhile, the electrochemical stability of the CoO_xCl_y at 250 mA cm⁻²_{geo} could sustain over 20 h (Supplementary Fig. 28). This catalyst system could even outperform the representative precious and non-precious metal oxide catalysts (Fig. 1e and Supplementary Table 1).



Fig. 1 | **The CER performance of amorphous CoO_xCl_y catalysts. a** The LSV curve of CoO_xCl_y and CoO_x catalysts in 0.5 M NaCl and NaClO₄ at pH 2, respectively. The CoO_xCl_y and CoO_x catalysts were obtained after 2 h electrodeposition at 1.67 V in the electrolyte containing 0.1 M Co²⁺ and 0.5 M Cl or 0.5 M ClO at pH 2, respectively. 1.67 V was selected based on the CER onset potential of the first potential scan. b The overpotential trend at 10 mA cm⁻² with increasing the electrolyte containing 0.1 M Co²⁺ and 0.5 M Cl at pH 2. The overpotential was iR

Cl--induced deposition of amorphous CoO_xCl_y

To understand the catalyst deposition coupled CER process, we characterized the composition and structure of the deposited catalyst film. Compared to the CoO_x film, the Co K α and Cl K α mapping of CoO_xCl_y catalyst electrode by XRF show that the Co and Cl elements were distributed throughout the entire catalyst film, indicating the codeposition of Cl⁻ together with Co²⁺ (Fig. 2a and Supplementary Fig. 29). The energy-dispersive X-ray spectroscopy (EDX) mappings confirmed the presence of the relatively even distribution of Co, Cl, and O for the formation of CoO_xCl_v film even after 500 h stability test (Fig. 2b and Supplementary Fig. 30). The scanning electron microscope (SEM) images demonstrated the comparable dense surfaces of both CoO_xCl_y and CoO_x films (Supplementary Figs. 31, 32). The crosssection of CoO_xCl_y film characterized by scanning transmission electron microscopy (STEM) showed a film thickness of about 600 nm after 40 h of electrodeposition at 1.67 V (Fig. 2c and Supplementary Fig. 33). The EDX mapping of the cross-section further exhibited the relatively homogenous distribution of Co and Cl elements (Supplementary Fig. 34). In addition, we characterized the structure of the electrodeposited catalyst films. X-ray diffraction (XRD) demonstrated that CoO_xCl_v film was absent of characteristic diffraction peaks (Supplementary Fig. 35). The crystal structure of both CoO_x and CoO_xCl_y catalysts was further imaged by the HRTEM. The lattice fringe together with the selected area electron diffraction (SAED) pattern showed that CoO_x film was composed of CoO_x nanocrystals (Supplementary Fig. 36). In contrast, CoO_xCl_y film exhibited an amorphous structure without lattice fringe and SAED rings (Fig. 2d). Even after 500 h of operation, the CoO_xCl_y film still remained the amorphous structure (Supplementary Fig. 37). The amorphous nature was probably ascribed

corrected with 90% ohmic resistance. The electrolyte resistance was 46.0 Ω . **c** The Cl₂ selectivity of CoO_xCl_y catalyst under different Cl⁻ concentrations and anodic potentials during the deposition process at pH 2. The data were recorded based on a rotating disk electrode method at 1600 RPM. The error bars represent the standard deviation for triplicate measurements. **d** The LSV curves and corresponding DEMS signals during CER at pH 2 at 10 mV s⁻¹. **e** The performance comparison between different CER catalysts (the detailed data were summarized in Supplementary Table 1).

to the Cl⁻ co-deposited with Co²⁺ in the catalyst as it may be induced by reconstruction or distortion relative to CoO_x^{40} .

Suppression of Co oxidation state by Cl⁻ in the CoO_xCl_y catalyst Further, compared with the CoO_x film in the absence of C¹ (Supplementary Fig. 38), a peak at 197.6 eV for Cl 2p in CoO_xCl_v film was observed, which was attributed to the Co-Cl bond (Fig. 2e)^{25,41,42}. Analogous to Co (Supplementary Fig. 8 and 39a), the content of Cl increased with the electrodeposition time (Supplementary Fig. 39b and Supplementary Table 2). In addition, the chemical states of the asprepared CoO_xCl_v film were analyzed by XPS to study the role of Cl⁻ inclusion. The Co 2p spectra can be well deconvoluted by the Co²⁺ and Co³⁺ components according to the reported binding energies^{43,44} (Fig. 3a). The Co^{3+}/Co^{2+} ratio in the CoO_x film is higher than that in the CoO_xCl_v film (Fig. 3b). Moreover, the Raman spectrum of CoO_xCl_v film with peaks at 180.7, 456.2, 502.4, 591.5, and 660.1 cm⁻¹ belong to the 3 F_{2g} , 1 E_g , and 1 A_{1g} vibrational modes of cobalt oxides (Fig. 3c)^{45,46}. The blueshift of the vibration bands relative to those of CoO_x film indicated the variation of Co local coordination owing to Cl⁻ introduction, consistent with its relatively lower Co oxidation state (Fig. 3a)⁴⁰.

In situ X-ray absorption fine structure (XAFS) spectroscopy was further applied to elucidate the valence state of Co under CER conditions (Supplementary Fig. 40). The Co K-edge X-ray absorption near edge structure spectra (XANES) of CoO_xCl_y and CoO_x film were collected under anodic polarization in the 0.5 M NaCl and NaClO₄ electrolytes, respectively⁴⁷. As shown in Fig. 3d and Supplementary Fig. 41, the anodic polarization induced a shift of the Co K-edge to a higher energy region for both the CoO_xCl_y and CoO_x film suggesting an increase in the Co oxidation state^{44,48,49}. However, the CoO_xCl_y film



Fig. 2 | **The composition and structure of the CoO_xCl_y catalyst. a** The XRF Co Kα and Cl Kα mapping of the CoO_xCl_y film electrodeposited in the electrolyte containing 0.1 M Co²⁺ and 0.5 M Cl⁻ at 1.67 V for 10 h. The scale bar is 1 mm. **b** The EDX mapping of the CoO_xCl_y film after 40 h electrodeposition at 1.67 V. **c** The cross-section TEM images of the CoO_xCl_y film after 40 h electrodeposition at 1.67 V. **d** The

HRTEM images and corresponding SAED patterns (inset) of the CoO_xCl_y film after 40 h electrodeposition at 1.67 V. **e** The Cl 2p X-ray photoelectron spectroscopy (XPS) spectrum of the CoO_xCl_y film. This peak could be well fitted by 197.6 eV for Cl $2p_{3/2}$ and 199.4 eV for Cl $2p_{1/2}$, respectively^{41,42}.

underwent a larger shift (-0.83 eV) than that of the CoO_x film (-0.41 eV) upon the potential increase from OCP to 2.02 V (Fig. 3e). Clearly, the CoO_xCl_y film showed lower Co K-edge absorption energy relative to that of CoO_x film at all the test potentials. This suggested a relatively lower Co oxidation state in CoO_xCl_v film. Even at OCP, the Co valence state remains lower in the CoO_xCl_y relative to CoO_x, thus upon increasing the potential to 1.67 V, an obvious increase in the Co valence state allows CER to initiate⁴⁰. Further, we analyzed the Fouriertransformed of k³-weighted Co K-edge extended X-ray absorption fine structure (EXAFS) spectra of both in CoO_x and CoO_xCl_v catalysts, which showed two major peaks associated with Co-O (~1.90 Å) and Co-Co (~2.85 Å) coordination (Supplementary Figs. 42, 43). An additional peak located around ~2.35 Å in the CoO_xCl_y catalyst was observed, indicating the presence of Co-Cl coordination (Supplementary Fig. 43). Compared to the CoO_x catalyst, the CoO_xCl_y catalyst exhibited a lower fitted Co-O coordination number during the anodic polarization (Supplementary Tables 3, 4) and displayed a larger increase in Co-O coordination number relative to the CoO_x catalyst, in coincidence with its significant degree of Co oxidation state variation^{50,51}.

To get a deeper understanding of the CER process together with the effect of Cl⁻ introduction, in situ spectroelectrochemical UV-Vis was applied to track the deposition processes of CO_xCl_y and CO_x catalysts (as illustrated in Fig. 4a and Supplementary Fig. 44). Figure 4b demonstrated the variation of spectral absorption of the COO_x film deposited in the absence of Cl⁻ at 1.67 V. The absorption intensity increased rapidly within 30 min of deposition and several characteristic peaks appeared at AI-A4 corresponding to the charge transfer transitions^{52,53}. The absorption centered at A2-A4 was assigned to the higher oxidation state of Co coordinated with highly oxidized oxygen species in the oxygen evolution process⁵⁴⁻⁵⁶. Instead, A2-A4 absorbance bands were absent in CoO_xCl_v film, again suggesting a relatively lower oxidation state of Co than that in CoO_x film (Fig. 4c). Moreover, the absorbance intensity of CoO_xCl_y film is almost two times that of the CoO_x film. Such a phenomenon could be ascribed to the Cl⁻ accelerated electrodeposition and CER process. To clarify this phenomenon, we tracked the absorption of electrodeposited CoO_xCl_y by stepwise increasing the Cl⁻ concentration from 0 to 2.0 M (Supplementary Fig. 45). As expected, the A1 absorption intensity distinctly increased with the concentration of Cl⁻ (Fig. 4d). When Cl⁻ concentration was at 0.01 M, the A1 band intensity was relatively weaker while the A2-A4 appeared, indicating the existence of OER active phase analogous to that in CoO_x. These results together with in situ XANES analyses illuminate that the Cl⁻ introduction promotes the electrodeposition process and improved the Cl₂ selectivity by suppressing the formation of highly oxidized Co sites required for OER.

The role of CI⁻ on the CER process by DFT calculations

Subsequently, we applied the density functional theory (DFT) calculation to further clarify the effect of Cl on Co valence states. Based on the above experiment results, we selected cobalt oxide with both octahedral and tetrahedral Co ($Co_{octahedral} / Co_{tetrahedral} = 3$) (Supplementary Fig. 46 and Supplementary Table 5), and constructed six possible explicit solution models based surfaces without and with Cl in the catalyst (Supplementary Fig. 47). The negative formation energies suggested that Cl was spontaneously introduced into the catalyst (Supplementary Note 4, Supplementary Table 6, and Supplementary





Fig. 3 | **The role of Cl**[•] **on the Co oxidation state for the CoO_xCl_y catalyst. a** The Co 2*p* XPS spectra of the CoO_xCl_y and CoO_x film after 2 h electrodeposition at 1.67 V, respectively. **b** The corresponding Co³⁺/Co²⁺ ratio of the CoO_xCl_y and CoO_x film. **c** The Raman spectra of the CoO_xCl_y and CoO_x films after 10 h electrodeposition at 1.67 V. **d** Co K-edge XANES at different applied potentials from the

open-circuit condition to 2.02 V for $CoO_x Cl_y$ film. **e** Co K-edge energy at different applied potentials for CoO_x and $CoO_x Cl_y$ film. The absorption energy (E₀) is obtained from the first maximum in the first-order derivative as the electronic vacancy. The error bars represent the standard deviation for triplicate measurements.

Fig. 48). We chose two models with lowest formation energy (Cldoped-d and Cl-doped-e) to study the Co valence state in comparison with that of the Cl-free model (Supplementary Fig. 49). Because the surface atoms have lower coordination numbers than bulk atoms, surface Co sites are covered with water ligands. The density of electronic states (DOS) (Supplementary Fig. 50) shows that tetrahedra Co induces the defect states in the bandgap, attributed to the surface oxygens coordinative unsaturation and higher oxidation states of Co (Supplementary Fig. 51 and Supplementary Table 7). The defect states usually lead to significant optical absorption at the infrared and ultraviolet-visible regions⁵⁷, which may correspond to the A2-A4 absorbance of CoO_x film. When the surface oxygen atoms are substituted by Cl⁺, the defect states disappeared, in line with the absence of A2-A4 absorbance bands of CoO_xCl_v film. Moreover, the hole states near the Fermi level appear to suggest Cl⁻ in the catalyst enhanced electrical conductivity with reduced charge transfer resistance (Supplementary Fig. 13). As shown in Supplementary Figs. 52, 53 and Supplementary Tables 8, 9, the calculated Bader charge and bond length suggested that Cl⁻ introduction could increase the Co electronic charge and decrease the valence state, consistent with in situ UV-Vis and XANES results.

To gain an in-depth understanding of the high Cl_2 selectivity of CoO_xCl_y , we further studied the influence of Cl introduction on OER and CER. We first constructed the OER and CER reaction free-energy

diagrams as a function of electrode potential (U) at pH 2 and 0.5 M Cl⁻ based on the thermodynamically stable structures of various adsorbates^{10,58} (i.e. Cl^{*}, ClO^{*}, H₂O^{*}, HO^{*}, O^{*} and HOO^{*}) on Co sites (Supplementary Note 5). Without Cl⁻ in the catalyst, Cl^{*} is favorable at U < 1.4 V, which transferred to Cl-O^{*} at a higher potential (Fig. 5a). On the other hand, in the presence of Cl in the catalyst, OH^{*} became mover favorable at U < 1.4 V (Fig. 5b and Supplementary Fig. 54). Hence, we evaluated the influence of pH and Cl⁻ concentrations on the reaction pathways (Supplementary Fig. 55). The results indicated that even with very low Cl⁻ concentrations, CER is predominant at pH 2 on both Cl-free and Cl-containing catalysts.

To further understand the high selectivity of Cl₂ to O₂, we compared the reaction free-energy diagrams between OER and CER (Fig. 5c, d and Supplementary Note 6). For CER, we consider the mechanism that proceeded through the Volmer (O*/* + Cl \rightarrow ClO*/Cl* + e) and the subsequent Heyrovsky step (ClO*/Cl* + Cl \rightarrow Cl₂ + e), where the Cl directly adsorbed on the active site. On the Cl-free catalyst, the theoretical overpotentials for the OER and CER were 0.63 V and 0.39 V, respectively (Fig. 5c, d, Supplementary Figs. 56–59, and Supplementary Tables 10, 11). On the Cl-containing catalyst, the theoretical overpotential for both the OER and CER were reduced to 0.41 V and 0.2 V, respectively (Fig. 5c, d, Supplementary Figs. 60–63, and Supplementary Tables 12, 13). Even though the theoretical potential of 1.36 V for CER is 0.13 V higher than the theoretical



Fig. 4 | In situ UV-Vis spectra for tracking the electrodeposition process of the CoO_xCl_y and CoO_x film on FTO. a The schematics illustrate the in situ spectroelectrochemical UV-Vis tracking of the catalyst deposition process (left). The schematic diagram for CER on the CoO_xCl_y electrode (right). b The UV-Vis absorption spectra of the deposited CoO_x film in the electrolyte containing 0.1 M

 $\rm Co^{2*}$ and 0.5 M ClO₄ at 1.67 V. c The UV-Vis absorption spectra of the deposited $\rm CoO_x Cl_y$ film in the electrolyte containing 0.1 M $\rm Co^{2*}$ and 0.5 M Cl at 1.67 V. d The UV-Vis absorption spectra of the deposited $\rm CoO_x Cl_y$ film upon increasing the Cl concentration from 0.01 to 2.0 M. Inset shows the absorption spectra around the A2-A4 range for the CoO_x Cl_y electrodeposited in 0.01 M Cl electrolyte.

potential of 1.23 V for OER, such results still suggest that CER is more favorable than OER on the catalysts with and without Cl. However, the presence of Cl⁻ in the catalysts can accelerate the catalysis reaction. In addition, there is a transition of the active site from Co^{*} to the Co-O^{*} for CER with the Cl⁻ introduction into the catalyst (Fig. 5e). The same trend was observed for other Cl-containing models (Supplementary Figs. 64–67, and Supplementary Tables 14, 15).

In this work, we present a highly selective and robust CER system based on an amorphous CoO_xCl_y electrocatalyst electrodeposited in an acidic electrolyte containing both Co²⁺ and Cl⁻ ions. Notably, this work presents the first case in that Cl⁻ ions work as a promoter for catalyst deposition and CER. Experimental results demonstrate that ~100% CER selectivity has been acquired and in situ spectroscopic studies together with DFT theoretical calculations show that the introduction of Cl⁻ into the catalyst suppresses the formation of highly oxidized Co sites required for water oxidation while allowing for the active site transition from Co* to Co-O* for CER with enhanced CER selectivity via the Volmer-Heyrovsky pathway. This work provides a new avenue to stabilize the non-precious metal oxides for anodically electrocatalytic reactions in strong acids and we would suggest in situ characterize the catalyst system with cautions before we categorize a soluble species or molecule as a homogeneous catalyst.

Methods

Chemicals

All chemicals used in this work were commercially available. Cobalt nitrate hexahydrate (Co(NO₃)₂·6H₂O, 99.99%), cobalt chloride hexahydrate (CoCl₂·6H₂O, 99.99%), ferric nitrate nonahydrate (Fe(NO₃)₃·9H₂O, 99.99%), manganese nitrate tetrahydrate 99.99%), potassium ferricyanide trihydrate $(Mn(NO_3)_2 \cdot 4H_2O_1)$ (K₃Fe(CN)₆·3H₂O, 99.99%), urea (CO(NH₂)₂, 99.999%), sodium chloride (NaCl, 99.99%), ammonium fluoride (NH₄F, 99.99%), perchloric acid (HClO₄, 70-72%), and anhydrous ethanol (C₂H₆O, 99.8%) were purchased from Sigma-Aldrich. Cobalt sulfate heptahydrate (CoSO₄·7H₂O, 99.99%), nickel nitrate hexahydrate (Ni(NO₃)₂·6H₂O, 99.99%), copper nitrate trihydrate (Cu(NO₃)₂·3H₂O, 99.99%), sodium perchlorate monohydrate (NaClO₄ · H₂O, 99.99%), and cobalt acetate tetrahydrate (Co(acetate)₂·4H₂O, 99.9%) were purchased from Aladdin. Hydrochloric acid (HCl, 31%), nitric acid (HNO₃, 65-68%), sulfuric acid (H_2SO_4 , 95–98%), and acetone (C_3H_6O , 99.9%) were purchased from KESHI. Commercial ruthenium oxide (RuO_2 , Ru > 75%) and iridium oxide (IrO_2 , Ir > 85%) were purchased from Adamas. In all experiments, ultrapure deionized water (Milli-Q, 18.2 MΩ cm⁻¹) was used for the electrolyte preparation. The clean fluorine-doped tin oxide (FTO) with resistance <15 ohm/sq was used as a substrate for electrodeposition. All chemicals were used without further purification.



Fig. 5 | **DFT calculations to understand the role of Cl⁻ on the reaction mechanism. a** The surface phase diagram of the Cl-free catalyst model. **b** The surface phase diagram of the Cl-containing catalyst model. **c** The reaction free energy diagram of OER on the Co sites of the Cl-free and Cl-containing catalysts,

respectively. **d** The reaction free energy diagrams of CER on the Co sites of the Clfree and Cl-containing catalysts, respectively. Insets: corresponding computational models for each CER step. **e** The schematic illustration of the role of Cl introduction on the CER reaction active sites.

In situ electrodeposition of $\text{CoO}_{x}\text{Cl}_{y}$ catalyst

The electrolyte for the working electrode side was prepared by mixing 0.1 M Co^{2+} salts and 0.5 M NaCl in 40 mL of deionized water. The pH of the electrolyte was adjusted to 2 using HClO₄. Typically, the CoO_xCl_y film was in situ electrodeposited on the FTO in the Co salt-NaCl electrolyte at 1.67 V at 25 °C. The electrochemical tests were conducted in a custom-designed PEEK cell. The Pt counter electrode was separated by an N117 Nafion membrane with a thickness of 183 um and size of 4 × 4 cm and was immersed in 0.5 M NaCl (pH 2) as well, to avoid the corrosion and the cathodic deposition of Co-based film. Before the experiment, the membrane was inserted in 5% hydrogen peroxide soaking for 60 min, deionized water for 30 min, 5% diluted sulfuric acid for 60 min, and deionized water for 30 min at around 80 °C.

Synthesis of Co₃O₄ film

Typically, the Co₃O₄ electrode was prepared by a simple hydrothermal method following a previous report³⁹. Specifically, 0.233 g Co(NO₃)₂· $6H_2O$, 0.07 g of NH₄F, and 0.240 g of CO(NH₂)₂ were dissolved in 100 mL deionized water under stirring for 30 min. Then, the obtained solution was transferred to a 50 mL Teflon-lined autoclave with an FTO inside as substrate. The reaction was conducted at 120 °C for 10 h and then cooled to room temperature. The obtained FTO substrates were washed with deionized water, dried at 60 °C overnight, and annealed in air at 350 °C for 2 h with a heating rate of 2 °C min⁻¹.

Synthesis of RuO₂ and DSA catalysts

To prepare the RuO₂ film electrode, 5 mg commercial RuO₂ was dissolved into 0.95 mL ethanol by adding 0.05 mL 5 wt% nafion117 solution under ultrasonic treatment for 60 min. 10 uL of this suspension was dropped onto the clean FTO (the efficient contact area is 0.283 cm²) for the formation of a catalyst film. The DSA catalyst film was prepared based on the commercial protocol on Ti (the ratio of Ir/Ru is 2).

Electrochemical characterizations

The CER tests were conducted based on a standard three-electrode with a computer-controlled electrochemical workstation (Bio-Logic SP -200). FTO (efficient contact area: 0.283 cm²), platinum mesh, and Hg/ Hg₂SO₄ electrode were used as the working electrode, counter electrode, and reference electrode, respectively. A custom-designed PEEK cell with a quartz window was used as the electrochemical cell. All experiments were carried out in a temperature-controllable box, and the temperature was kept at 25 °C if not specially indicated. Before electrochemical experiments, the reference electrodes were calibrated by a standard reversible hydrogen (RHE) electrode (PHYCHEMI) in the corresponding electrolytes after 60 min Ar bubbling. For the working electrode, magnetic stirring at 1500 RPM was applied to remove chlorine bubbles during the electrochemical tests. All the measured potentials were calibrated with a reversible hydrogen electrode (RHE) as the reference. The pH values of the electrolyte were measured through an Ohaus Starter 2100 pH meter with temperature calibration.

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For the electrochemical tests at different NaCl concentrations, the NaClO₄ \cdot H₂O was added for compensation to maintain a constant ionic strength⁸. The ohmic resistance (R) for iR corrections was determined by electrochemical impedance spectroscopy (EIS), and the recorded ohmic resistance was around 46.0 Ω , as shown in Supplementary Fig. 13. The overpotential in Fig. 1b and potential in Supplementary Fig. 28 were iR corrected with 90% ohmic resistance, while the other data were not iR-corrected. Electrochemical impedance spectroscopy (EIS) was measured at 1.67 V with a frequency scan range from 100 kHz to 10 mHz, and the amplitude of the sinusoidal wave was 10 mV.

In situ X-ray Absorption Fine Structure (XAFS) measurements

The XAFS measurements were recorded at the BL11B beamline of the Shanghai Synchrotron Radiation Facility (SSRF). The beam current of the storage ring was 220 mA in a top-up mode and the Si (111) double-crystal monochromator was applied to monochromatize the incident photons, with an energy resolution $\Delta E/E \sim 2 \times 10^{-4}$. The collected spot size of the sample was $\sim 200 \ \mu m \times 250 \ \mu m$ (H \times V). And the Co foil was used to calibrate the position of the absorption edge (E₀). All XAFS spectra were collected in fluorescence mode. The E₀ was obtained from the largest peak in the 1st derivative XANES according to the previous work^{47,59}. The CoO_xCl_y and CoO_x films were polarized at different anodic potentials in a three-electrode system, respectively. After the corresponding current density reaches a steady state, the XAFS data was collected at the indicated potentials. All XAFS data were processed by the ATHENA and ARTEMIS modules implemented in the IFEFFIT software package.

In situ UV-Vis

The in situ UV-Vis spectroscopy was implemented on a QE Pro UV-Visible spectrometer (Ocean Optics) equipped with an HL-2000 light source and a DH-2000-BAL source. A fiber-optic cable (100 μ m fiber core diameter) was used to direct the light from the light source through the sample to the detector. The UV-Vis spectra were recorded in the different electrolytes as a function of electrolysis times at 1.67 V. The FTO (area: 0.283 cm²), Pt electrode, and Hg/Hg₂SO₄ were used as the working electrode, the counter electrode, and the reference electrode, respectively. The spectrum of the unpolarized cleaned FTO immersed in the same electrolyte was recorded for the background subtraction of the UV-Vis spectra of the deposited CoO_xCl_y and CoO_x catalysts.

Online differential electrochemical mass spectrometry (DEMS)

The online DEMS experiments were implemented with the chamber containing the quadrupole mass spectrometer (MS) with a Faraday-SEM detector array (Hiden Analytical, HPR-40), which allows the isolation of the ion source from the electrochemical cell by forming a small pre-chamber^{37,38}. For the DEMS measurements, a custom gastight electrochemical H-cell with a conventional three-electrode arrangement was employed for the electrochemical experiments. In this experiment, a Nafion membrane was used to separate the cathode and anode. The acquisition probe is vertically attached to the working electrode where the volatile species pass through a porous Teflon membrane into the vacuum inlet of the mass spectrometer. For the DEMS measurements, the multiplier mode was adopted. The LSV and mass spectrometric cyclic voltammograms (MSCVs) were acquired in electrolytes containing 0.1 M Co2+ with 0.5 M NaClO4 or 0.5 M NaCl, respectively. The MSCVs for the concerned species ($^{70}Cl_2$: m/z = 70, ${}^{1}H^{16}O^{35}Cl: m/z = 52$, ${}^{35}Cl^{32}O_{2}: m/z = 67$, ${}^{35}Cl^{16}O: m/z = 51$ and ${}^{32}O_{2}: m/z = 67$, ${}^{35}Cl^{16}O: m/z = 51$ z = 32) were recorded. Before the tests, the electrolyte was purged with Ar (99.999%) over 1 h to remove the dissolved oxygen.

Materials characterizations

The X-ray fluorescence method based on Co and Cl content analyses was conducted on a Bruker M4 Tornado instrument with Rh target for

X-ray generation. The catalyst morphology of the samples was characterized by a SU8010 field emission scanning electron microscope (SEM). The cross-section of the catalyst was characterized by a transmission electron microscope (JEOL JEM2100F) operating at 200 keV in both TEM and STEM modes. The electron beam spot size is 1.0 nm. An EDX detector (Oxford Instruments) was equipped for the element analysis. The TEM sample for cross-section observation was prepared by cutting the electrodeposited FTO electrode into two pieces, of which the cross-section was firstly glued by the epoxy resin and then processed through Ar⁺ ion-milling and polishing (PIPS II, GATAN) at grazing incidence (<5°). The X-Ray Diffraction (XRD) measurement of CoO_xCl_y catalyst was conducted on Bruker D8 ADVANCE A25X. The high-resolution TEM (HRTEM) images of CoO_xCl_y and CoO_x catalysts were taken on an FEI-Tecnai G2 microscope. Moreover, the element mapping and selected area electron diffraction (SAED) were performed on an FEI TECNAI G2 F30 field emission transmission electron microscope. The X-ray photoelectron spectroscopy analyses were carried out using a Thermo Scientific K-Alpha system to evaluate the composition and chemical states of CoO_xCl_y and CoO_x catalysts. Al Ka X-ray source with a power of 250 W was used as the X-ray source and the binding energies were calibrated by referring to the C1s peak (284.8 eV) of the adventitious carbon. The Raman was carried out with an XploRA PLUS Raman spectrometer (Horiba) equipped with a 532 nm He-Ne laser as the excitation source.

Rotating ring-disk electrode detection of selectivity \mbox{Cl}_2 evolution

The Cl₂ selectivity for CER was analyzed based on the RRDE system in which the disk electrode generates Cl₂ while the Pt ring electrode monitors Cl₂ concentration through its electrochemical reduction current. The potential of the Pt ring was fixed at 0.95 V to avoid the current response from O₂ reduction. Before each measurement for Cl₂ quantification, the Pt ring-disk electrode was successively polished with aqueous suspensions of 1.0 µm, 0.3 µm, and 0.05 µm alumina. It was then rinsed with deionized water, ultrasonically cleaned in ethanol for 15 s, and dried. Later on, the Pt ring was further electropolished in 0.5 M H₂SO₄ by 30 CV scans between -0.1 and 1.7 V at a rate of 500 mV s⁻¹. The chronoamperometry method was applied for the Cl₂ selectivity quantification. During the Cl₂ selectivity measurements, the potentiostatic electrolysis was carried out for 5 min at 1600 RPM at different applied disk potentials³⁵. Each experiment was repeated three times with an interval of 5 min in an Ar-saturated electrolyte at pH 2 containing 0.1 M Co(NO₃)₂ and NaCl with different concentrations.

The CER current density (j_{CER}) on the disk electrode can be calculated by the following equation:

$$j_{\text{CER}} = |j_{\text{R}} / N_{\text{I}}| = |i_{\text{R}} / A \times N_{\text{I}}| \tag{1}$$

Where j_{R} is the current density measured on the ring, N₁ is the collection efficiency (0.37), and A is the disk electrode area (0.196 cm²).

The OER current density (j_{OER}) can be calculated by the following equation:

$$j_{\text{OER}} = j_{\text{D}} - j_{\text{CER}} = j_{\text{D}} - |j_{\text{R}} / N_{\text{I}}| = j_{\text{D}} - |i_{\text{R}} / A \times N_{\text{I}}|$$
(2)

Where j_D is the current density measured on the disk.

The Cl₂ selectivity was calculated by the following equation:

$$Cl_2 \text{ selectivity}(\%) = 100 \times 2 \times j_{CER} / (j_D + j_{CER})$$

= 100 \times 2 \times |j_R / N_1| / (j_D + |j_R / N_1|) (3)

Computation details

The first-principles calculations based on density functional theory (DFT) were carried out within the projected augmented wave

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method, as implemented in the Vienna Ab-initio Simulation Package (VASP). The $CoO_{v}Cl_{v}$ film catalyst models were made by cleaving along the (100) facet of Co₃O₄ which contained two tetrahedrons and six octahedrons and were solved by explicit solution models. To generate highly accurate electrochemical stability diagrams, we employed a recently developed approach. which included the use of a Hubbard U term, a van der Waals functional (optPBE), and the use of spin polarization for the calculations. The U-value applied for d-orbitals of Co is taken as 3.50 eV. A cutoff energy of 500 eV was used for the plane wave expansion at all calculations of structure relaxations, vibrational frequencies, and single point energy. Monkhorst-Pack $(4 \times 4 \times 1)$ k-point grids were used for Brillouin zone integration. The equilibrium structures were obtained when the maximum atomic forces were smaller than 0.01 eV/Å and when the total energy convergence of 10⁻⁶ eV was achieved for the electronic selfconsistent field loop. The thermodynamic energy correction used in the free energy calculations was listed in Supplementary Table 6. The detailed descriptions of formation energy, OER, and CER reaction free energies calculations have been discussed in Supplementary Notes 4-6.

Data availability

All relevant data are provided in this article and its Supplementary Information.

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Author contributions

C.C. led the project. M.X. carried out the experiments. Q.W. repeated the experiments. C.L., M.X., J.L. (Jiong Li), and Q.W. finished the XAFS measurements. L.L. and M.X. proceed with the DEMS. A.M. and M.V. characterized STEM. J.L. (Junwu Liang), Z.Z., J.Z., L.Z., and R.K. conducted the density functional theory calculation. M.X. and C.C. wrote the manuscript. All authors commented on the manuscript.

Competing interests

The authors declare no competing interests.

Additional information

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