Catalytic effects of molybdate and chromate–molybdate films deposited on platinum for efficient hydrogen evolution

Oscar Diaz-Morales, Aleksandra Lindberg, Vera Smulders, Athira Anil, Nina Simic, Mats Wildlock, Germán Salazar Alvarez, Guido Mul, Bastian Mei and Ann Cornell

Abstract

BACKGROUND: Sodium chlorate (NaClO₃) is extensively used in the paper industry, but its production uses strictly regulated highly toxic Na₂Cr₂O₇ to reach high hydrogen evolution reaction (HER) Faradaic efficiencies. It is therefore important to find alternatives either to replace Na₂Cr₂O₇ or reduce its concentration.

RESULTS: The Na₂Cr₂O₇ concentration can be significantly reduced by using Na₂MoO₄ as an electrolyte co-additive. Na₂MoO₄ in the millimolar range shifts the platinum cathode potential to less negative values due to an activating effect of cathodically deposited Mo species. It also acts as a stabilizer of the electrodeposited chromium hydroxide but has a minor effect on the HER Faradaic efficiency. X-ray photoelectron spectroscopy (XPS) results show cathodic deposition of molybdenum of different oxidation states, depending on deposition conditions. Once Na₂Cr₂O₇ was present, molybdenum was not detected by XPS, as it is likely that only trace levels were deposited. Using electrochemical measurements and mass spectrometry we quantitatively monitored H₂ and O₂ production rates. The results indicate that 3 mmol L⁻¹ Na₂Cr₂O₇ (contrary to current industrial 10–30 mmol L⁻¹) is sufficient to enhance the HER Faradaic efficiency on platinum by 15%, and by co-adding 10 mmol L⁻¹ Na₂MoO₄ the cathode is activated while avoiding detrimental O₂ generation from chemical and electrochemical reactions. Higher concentrations of Na₂MoO₄ led to increased oxygen production.

CONCLUSION: Careful tuning of the molybdate concentration can enhance performance of the chlorate process using chromate in the micromolar range. These insights could be also exploited in the efficient hydrogen generation by photocatalytic water splitting and in the remediation of industrial wastewater.

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Keywords: hydrogen evolution reaction; chlorate process; molybdenum; chromate

INTRODUCTION

Sodium chlorate (NaClO₃) is one of the largest (by volume) electrochemically produced materials, with a yearly production capacity of approximately 4 million tons.¹ It is the precursor of chlorine dioxide (ClO₂), which is extensively used in the paper industry to bleach pulp feedstock. This bleaching technology, i.e. elemental chlorine-free (ECF) bleaching, is regarded as the best available technique. No other bleaching technology provides the same brightness while retaining the strength of the pulp, good economy and very low environmental impact.²

The chlorate process is an electrochemical process carried out in undivided electrochemical cells. It consists of a complex series of electrochemical and chemical reactions that convert chloride ions into chlorate, with hypochlorous acid and hypochlorite anions as intermediates. Concomitantly, hydrogen evolution (2H₂O + 2e⁻ → H₂ + 2OH⁻) takes place on the cathodes.³

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Since the process is performed in undivided cells, electroreduction of hypochlorite and chlorate anions can occur. These unwanted reactions reduce the energy efficiency of the process and are conventionally minimized by adding 3–6 g dm\(^{-3}\) \(\text{Na}_2\text{Cr}_2\text{O}_7\) to the electrolyte.\(^6\) The Cr\(^{VI}\) additive gets reduced during electrolysis and forms a poorly electrically conductive film of hydrous chromium(III) oxide/hydroxide (here denoted as \(\text{Cr(OH)}_3 \times n\text{H}_2\text{O}\)) on the cathode, which hinders the transport of hypochlorite and chlorate anions to the electrode surface, thus preventing their electroreduction. \(\text{Cr(OH)}_3 \times n\text{H}_2\text{O}\) still allows water to reach the cathode surface so the hydrogen evolution reaction (HER) can proceed in the presence of the film.\(^4,5\) An alternative mechanism explaining the selectivity of the \(\text{Cr(OH)}_3 \times n\text{H}_2\text{O}\) film has been suggested to be electrocatalytic, where Density Functional Theory (DFT) calculations show that the active site for hypochlorite reduction becomes blocked in the course of reduction by irreversible hydroxylation of the transition metal site during the reaction.\(^6\)

Na\(_2\)Cr\(_2\)O\(_7\) is pivotal in producing NaClO\(_3\) efficiently; without this additive it is not possible to produce chlorate with the currently available technologies. Unfortunately, all Cr\(^{VI}\) compounds are environmentally harmful and have proven to be carcinogenic, mutagenic and reprotoxic to humans, and therefore included in environmentally harmful and have proven to be carcinogenic, mutagenic and reprotoxic to humans, and therefore included in the REACH annex XIV list of the European Union (EU).\(^7,9\) Thus, since 2017, a special authorization is required to use Cr\(^{VI}\) compounds within the EU. Despite many research efforts devoted to finding less harmful alternatives to the Na\(_2\)Cr\(_2\)O\(_7\) additive for the chlorate process,\(^10–16\) finding a suitable replacement has proven to be very challenging due to the complex role of this additive\(^4\) as well as the harsh operating conditions (i.e. highly oxidizing electrolyte and high temperature).

Molybdate (Mo\(_2\)O\(_7^{2−}\)) has been investigated as a full or partial replacement to Na\(_2\)Cr\(_2\)O\(_7\) addition.\(^11,17–20\) Molybdate addition influences the chlorate process in several ways: (i) it activates the HER on cathodes of titanium,\(^18,19\) molybdenum\(^19\) and steel;\(^17\) (ii) it enhances the current efficiency towards HER on steel/iron and titanium;\(^11,12,15,16,20\) and (iii) it increases the rate of unwanted oxygen formation, in particular at high molybdate concentrations.\(^17,18\) Note that all these studies were made on the industrially relevant electrode materials iron/steel and titanium, which undergo reactions such as oxidation and hydride formation in the chlorate electrolyte. Steel corrodes when not under cathodic protection, and the different types of corrosion products influence the chemical and electrochemical reactions.\(^12\) Even under cathodic polarization, corrosion products are not completely reduced under conditions for hydrogen evolution at non-noble materials such as steel and titanium.\(^5\) On a hydrogen-evolving titanium cathode, a layer of titanium hydride is formed, changing the electrochemical properties of the electrode. This continuous change of properties of steel and titanium cathodes is also seen in full-scale operations.\(^21\) Thus, to pinpoint and further understand the role of the molybdate additive on HER electrocatalysis, we have conducted a study using platinum cathodes. Concentration limits of Na\(_2\)MoO\(_4\) were explored, aiming to run the chlorate process at low Na\(_2\)Cr\(_2\)O\(_7\) concentrations. Formed films were characterized by scanning electron microscopy (SEM) and X-ray photoelectron spectroscopy (XPS), and their influence on activity and selectivity towards HER, oxygen formation and hypochlorite reduction was investigated. Furthermore, it was found that in situ formed Cr–Mo mixed metal oxide films are chemically stable, providing essential benefits for cathode corrosion protection during process shutdown.

### EXPERIMENTAL

#### Surface characterization of the formed films

The Pt used as working electrode (rotating disc or foil) was mechanically polished to a mirror finish with alumina of decreasing grain size of 5, 0.3 and 0.05 μm L\(^{-1}\), prior to each experiment. Afterwards, it was thoroughly rinsed and sonicated for 10 min with ultrapure water (resistivity > 18.2 MΩ cm) to remove polishing residues. After mechanical polishing, the Pt electrode was electrochemically cleaned/activated by performing cyclic voltammetry between 0 and 0.75 V versus Ag/AgCl at 100 mV s\(^{-1}\) in a 0.1 mol L\(^{-1}\) NaOH (>99% Sigma Aldrich, Stockholm, Sweden).

For surface characterization experiments, cathodic depositions were formed on Pt foil polarized in 0.1 mol L\(^{-1}\) NaOH, at −0.8 V versus Ag/AgCl for 5 min, using a Pt wire as a counter electrode and a Hg/HgO as a reference, while purging the electrolyte by N\(_2\). After 5 min the samples were removed from the electrolyte without interrupting polarization. Different concentrations of additive in the form of Na\(_2\)MoO\(_4\) × 2H\(_2\)O (used as received from Sigma Aldrich, Stockholm, Sweden) were added and samples are named accordingly; for example, the sample with 10 mmol L\(^{-1}\) Mo salt formed at 22 °C (which was the room temperature when preparing these samples, noted as RT) is 10Mo-22 (see also Table 1). Finally, the samples were rinsed with water, ethanol and acetone.

Scanning electron microscopy (SEM) images were acquired using a Hitachi S-4800 instrument at 1 kV accelerating voltage, using platinum cathodes. Concentration limits of Na\(_2\)MoO\(_4\) were explored, aiming to run the chlorate process at low Na\(_2\)Cr\(_2\)O\(_7\) concentrations. Formed films were characterized by scanning electron microscopy (SEM) and X-ray photoelectron spectroscopy (XPS), and their influence on activity and selectivity towards HER, oxygen formation and hypochlorite reduction was investigated. Furthermore, it was found that in situ formed Cr–Mo mixed metal oxide films are chemically stable, providing essential benefits for cathode corrosion protection during process shutdown.

#### Table 1. Sample naming for characterization by SEM and XPS

<table>
<thead>
<tr>
<th>Sample</th>
<th>Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>(A) 10 mmol L(^{-1}) Mo, 22 °C</td>
<td>10Mo-22</td>
</tr>
<tr>
<td>(B) 80 mmol L(^{-1}) Mo, 22 °C</td>
<td>80Mo-22</td>
</tr>
<tr>
<td>(C) 10 mmol L(^{-1}) Mo, 80 °C</td>
<td>10Mo-80</td>
</tr>
<tr>
<td>(D) 80 mmol L(^{-1}) Mo, 80 °C</td>
<td>80Mo-80</td>
</tr>
<tr>
<td>(E) 10 mmol L(^{-1}) Mo, 3 μmol L(^{-1}) Cr, 22 °C</td>
<td>10Mo-3Cr-22</td>
</tr>
<tr>
<td>(F) 80 mmol L(^{-1}) Mo, 3 μmol L(^{-1}) Cr, 22 °C</td>
<td>80Mo-3Cr-22</td>
</tr>
</tbody>
</table>
to avoid damaging samples. A Quantera II scanning XPS microprobe instrument having monochromatic Al Kα radiation with a total power of 50 W was used for the XPS measurements. The beam spot size was 100 μm², and the base pressure of the main chamber was 7 × 10⁻⁷ bar. Survey scans were taken using five different sample regions to check the repeatability and uniformity of the samples. Survey scans were recorded from 1040 to 0 eV with a pass energy of 224 eV. In addition, high-resolution core shell element electronic spectra were taken using 55 eV pass energy and 0.05 eV resolution. CASA XPS® was used for further XPS analysis and deconvolution of peaks. The Mo individual peak area was determined using integration, having Shirley as the background function.

**Catalytic effect of Mo on HER**

The experiments were performed with a platinum rotating disc electrode (RDE) of 5 mm diameter from PINE Research (model WaveVortex 10 RDE). Polarization curves were recorded galvanostatically from high to low current density using a BioLogic model VSP (Figs 3 and 4) or a PAR 273A potentiostat (Figs 2 and 5). To correct for the IR drop a built-in high-frequency impedance method (85% correction) was used with BioLogic, whereas a current interrupt technique (100% correction) was used with the PAR potentiostat. In the latter, for each current density level the electrode was polarized for 15 s and then the current was interrupted repeatedly. The potential decay was measured for 500 μs at 1 MHz data sampling and recorded potential data were fitted using a third-order polynomial expression to establish the IR-corrected potential values.

A three-electrode system was employed, in which a Pt RDE was the working electrode, a Pt wire was the counter electrode and Ag/AgCl (3 mol L⁻¹ KCl) or Hg/HgO (1 mol L⁻¹ NaOH) was the reference electrode. When the electrolyte did not contain chloride ions a single compartment cell was used. For the polarization curve experiments using chloride in the electrolyte, the CE was placed in a compartment separated from the WE by a frit to inhibit the reaction products at the CE (hypochlorite) to reach the WE. The potentials reported in this work are all referred to an Ag/AgCl reference electrode, which were calculated according to Eqn (1):

\[ E_{\text{Ag/AgCl}} = E_{\text{Hg/HgO}} - 0.065 \text{V} \]

Electrode transfer for experiments with *ex situ* formed films were performed by first pre-polarizing the electrodes in hypochlorite-free solutions containing the desired additives for 5 min at −0.8 V versus Ag/AgCl and removed without interrupting polarization. The electrodes were then rinsed with water and transferred to a clean electrolyte containing 0.1 mol L⁻¹ NaOH, 2 mol L⁻¹ NaCl and 80 mmol L⁻¹ NaClO or only 0.1 mol L⁻¹ NaOH. Polarization curves were recorded from high to low current density, with constant N₂ flow.

**Effect of Mo and Cr in situ electrolyte additions on the Faradaic efficiency for hydrogen and oxygen evolution**

These measurements were done with a residual gas analyzer (RGA) operating with a triple-filter quadrupole mass spectrometer from Hiden (model HPR-20). This instrument is equipped with a heated quartz capillary that allows connecting the headspace of the electrochemical cell (at room pressure, ~760 torr) with the mass spectrometer chamber (~10⁻⁶ torr).

A gas-tight, two-electrode electrochemical cell was used for the mass spectrometry (MS) experiments, as previously reported by our group. A Pt foil was used as working electrode and a dimensionally stable anode (TiO₂RuO₂) as a counter electrode. Both working and counter electrodes had ~0.5 cm² of geometric surface area. Measurements were performed during galvanostatic electrolysis at 300 mA cm⁻² with flow of Ar as a carrier gas of 75 mL min⁻¹. In addition, to study oxygen formation from homogeneous reactions in the bulk electrolyte, MS measurements were made without the presence of any electrodes in the cell. The electrolyte contained 2 mol L⁻¹ NaCl and an initially 80 mmol L⁻¹ NaClO (both chemicals were purchased from VWR International, Spånga, Sweden and were used as received). Experiments were performed at 80 °C with the electrolyte adjusted and kept at pH 6.5 with an automatic titrator Metrohm 907 Titran titrator equipped with a Unitrode and Pt 1000 combined pH and temperature sensor. The electrolyte pH was kept constant by adding either 6 mol L⁻¹ HCl or 1 mol L⁻¹ NaOH.

NaClO concentration was monitored as a function of time by analyzing liquid samples by *ex situ* ultraviolet–visible spectroscopy, following a previously reported procedure.

**RESULTS AND DISCUSSION**

**Surface characterization of formed films**

The formed films indicated in Table 1 were analyzed by SEM and energy-dispersive X-ray analysis (EDX). No clear depositions could be seen on the platinum surface (Supporting Information Fig. S1). Probably, the formed films were of the order of nanometer thickness and therefore too thin for EDX measurements, which only recorded Pt peaks (Supporting Information Fig. S2). In addition, X-ray emission lines of Pt and Mo are very close in energy, and the Pt peak responses possibly overlap the Mo peaks.

However, the more surface-sensitive XPS identified elements present in the samples as well as oxidation states of Mo (Supporting Information Fig. S3). Molybdenum has oxidation states ranging from +2 to +6. Considering the importance of Mo oxidation state for different applications such as gas sensors, energy storage (desired Mo⁶⁺) or as anodic material for the solid oxide fuel cells (desired Mo⁴⁺), different techniques to obtain certain Mo oxidation states have been employed. Here, we deposited Mo layers from aqueous solutions and used XPS to evaluate the oxidation state of the electrodeposited Mo films as a function of the plating conditions (i.e. temperature and molybdate concentration). The deconvoluted core-level electronic spectra of Mo and Pt are shown in Fig. 1 and Supporting Information Fig. S4, respectively. The Pt core-level spectra show two well-resolved peaks corresponding to the Pt 4f7/2 and Pt 4f5/2 asymmetric representing metallic Pt. In the case of Mo, the prominent Mo peak and its intensity vary in all four samples, suggesting that each sample is unique regarding the different oxidation states. Compared to the electrolyte solution of MoO₃, where the Mo oxidation state is +6, partial reduction to lower oxidation states is observed in all the four samples. Peak positions and oxidation state percentages are shown in Table 2, and the peak positions go well with earlier reported values. Films deposited at higher temperatures show higher percentages of higher oxidation states (+6 and +5) compared to the films deposited at room temperature.

In all samples Ag was recorded, and by recording pure Pt foil used for samples deposition the presence of Ag was confirmed, indicating that the Ag response comes from the substrate itself.
The film deposited in an electrolyte containing 3 μmol L\(^{-1}\) Cr and 10 (or 80) mmol L\(^{-1}\) Mo displayed response characteristics only for Cr, with no signs of any Mo deposited (Supporting Information Fig. S5). An earlier study employing XPS found that Mo did not deposit from solutions containing Mo and phosphate ions.\(^{19}\) It was discussed that there could be a few different reasons for that, such as the formation of some interfering metal phosphate or competitive adsorption of molybdate and phosphate. In the case of Mo–Cr films, a similar phenomenon could be operating. Another possibility is that a Cr-hydroxide film is formed, which hinders molybdate reduction and, therefore, subsequent Mo deposition.\(^{18}\)

Interestingly, it was found that more Mo is electrodeposited from pH-neutral solutions compared to deposition at higher pH.\(^{19}\) Deposits formed in alkaline solutions showed two main peaks, located at 232.6 eV and 230.4 eV, assigned to MoO\(_3\) and MoO\(_2\). Reports indicate that formed films were either very thin or formed locally, as the signal from substrate (in the reported case, Ti) was recorded on these samples. Here, we agree with previous reported values for positions of the peaks for Mo +4 and +6 (Table 3).\(^{19}\) We also record Mo in +5 state, mainly in the samples 80Mo-22 and 10Mo-80.

### Catalytic effect of Mo on HER

IR-corrected polarization curves were recorded to analyze the effect of Mo addition on HER on a Pt electrode (Fig. 2). The study was performed under alkaline conditions as it represents the electrolyte close to a cathode where hydroxide ions formed during HER cause a locally increased pH. The cathode potential at the industrially relevant current density 300 mA cm\(^{-2}\) shifted 126 mV to a less negative potential when Mo was present in the electrolyte at 10 mmol L\(^{-1}\), compared to Mo-free electrolyte (Fig. 2(a)). Further increase in the molybdate concentration to 80 mmol L\(^{-1}\) deactivated the electrode; compared to the 10 mmol L\(^{-1}\) case, it shifted 50 mV to a more negative potential. This was likely due to a thick layer being formed by the higher Mo concentration and is in line with previous observations,\(^{19}\) where it was realized that the concentration of

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**Table 2.** Mo oxidation states determined by XPS and its percentage in each sample

<table>
<thead>
<tr>
<th>Sample/Mo oxidation state</th>
<th>+4</th>
<th>+5</th>
<th>+6</th>
</tr>
</thead>
<tbody>
<tr>
<td>10Mo-22</td>
<td>39.5</td>
<td>15</td>
<td>45.5</td>
</tr>
<tr>
<td>80Mo-22</td>
<td>—</td>
<td>85.5</td>
<td>14.5</td>
</tr>
<tr>
<td>10Mo-80</td>
<td>7</td>
<td>63.6</td>
<td>29.4</td>
</tr>
<tr>
<td>80Mo-80</td>
<td>15.7</td>
<td>18.5</td>
<td>65.8</td>
</tr>
<tr>
<td>10Mo-3Cr-22</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>80Mo-3Cr-22</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

**Table 3.** Peak positions (binding energy) for different Mo oxidation states

<table>
<thead>
<tr>
<th>Sample ID</th>
<th>Mo(^{+4}) BE (eV)</th>
<th>Mo(^{+5}) BE (eV)</th>
<th>Mo(^{+6}) BE (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>10Mo-22</td>
<td>230.3</td>
<td>231.2</td>
<td>232.4</td>
</tr>
<tr>
<td>80Mo-22</td>
<td>—</td>
<td>231.2</td>
<td>232.4</td>
</tr>
<tr>
<td>10Mo-80</td>
<td>230.1</td>
<td>231.3</td>
<td>232.5</td>
</tr>
<tr>
<td>80Mo-80</td>
<td>230.3</td>
<td>231.5</td>
<td>232.6</td>
</tr>
</tbody>
</table>
4 mmol L$^{-1}$ Mo addition had a stronger activating effect than 100 mmol L$^{-1}$ Mo, regardless of the substrate material. The same trend was observed also at higher temperature (80 °C; Fig. 2(b)).

Polarization curves recorded at RT reveal two regions. Without any Mo additive the Tafel slope was around $-190$ mV dec$^{-1}$ in the higher current density region. Molybdate addition lowered the Tafel slopes: with addition of 80 mmol L$^{-1}$ Mo it was $-175$ mV dec$^{-1}$ and with 10 mmol L$^{-1}$ Mo addition it amounted to $-150$ mV dec$^{-1}$. A similar trend was recorded in the lower current density region: without Mo addition the Tafel slope was $-73$ mV dec$^{-1}$; with 80 mmol L$^{-1}$ Mo it was $-63$ mV dec$^{-1}$; and with 10 mmol L$^{-1}$ Mo addition it was $-57$ mV dec$^{-1}$. A Tafel slope of 125 mV dec$^{-1}$ has been reported for HER on Pt in 0.5 mol L$^{-1}$ NaOH. However, values in the range 100–130 mV dec$^{-1}$, or even lower, were found with Pt-based and modified catalysts.

The differences can be assigned to different pH values of the electrolytes in the studies, and to the potential dependence of the Tafel slope.

Another difference among the samples is the difference in oxidation state of Mo, directly after deposition (Table 2). The highest activity towards HER was recorded on the sample where the +6 oxidation state dominates, i.e. with addition of 10 mmol L$^{-1}$ Mo.

Pt catalysts typically have lower activity for HER in alkaline media compared to acidic, due to the sluggish water dissociation step. Therefore, some elements such as Mo may be added to Pt to provide active sites for the sluggish step and significantly improve Pt activity in alkaline media. When combining molybdenum oxide sheets with Pt nanoparticles, the activity towards HER compared to pure Pt was improved.

The polarization curves measured on the Pt RDE at 2500 rpm in 2 mol L$^{-1}$ NaCl (Fig. 3) illustrate the system with three different concentrations of Na$_2$CrO$_4$. The black open squares curve, acquired in the absence of NaClO, shows a monotonic increase in potential at high current densities, as expected for the HER. The grey dashed line indicates the current density typically used industrially (300 mA cm$^{-2}$).

The black filled squares curve was acquired in the presence of NaClO (without chromate and molybdate additives). This curve shows a limiting current between 0 and $-1$ V versus Ag/AgCl, which indicates that chromate reduction is suppressed, whereas HER is the dominating reaction at potentials negative to $-1$ V versus Ag/AgCl.

Upon adding 0.5 μmol L$^{-1}$ Na$_2$CrO$_4$ to the electrolyte (red curve in Fig. 3(a)), the limiting current corresponding to hypochlorite electroreduction shifts to lower current densities, indicating partial inhibition of the reaction. At chromate concentrations higher than 3 μmol L$^{-1}$ in the electrolyte (see red curves in Figs. 3(b, c)) full inhibition of the hypochlorite electroreduction occurred, in agreement with previous reports. With an increase in chromate concentration, the thickness of the chromate film formed increases, leading to an increase in the performance. However, here we see that there is a critical concentration of Cr needed for suppression of hypochlorite reduction. It is also important to note that the polarization curves acquired with Cr in solution overlap with the curve obtained without hypochlorite at potentials negative to $-1$ V versus Ag/AgCl, indicating that HER is not much affected by the presence of the Cr film.

The molybdate additive affects both the hypochlorite electroreduction and the HER, as seen in the purple curve in Fig. 3. The limiting current corresponding to hypochlorite electroreduction shifts to lower current densities when 10 mmol L$^{-1}$ Na$_2$MoO$_4$ is added to the electrolyte, depicting a small retarding effect of the MoO$_4^{2-}$ film on the hypochlorite electroreduction. An increase in cathodic current efficiency at small chromate concentrations in the presence of molybdate was recently reported. The cathode potential shifts to less negative potentials in the presence of the molybdate additive: when the Cr$^{3+}$ concentration amounts to 13.5 μmol L$^{-1}$, the cathode potential shifts 260 mV to less negative values at 300 mA cm$^{-2}$ compared to the values obtained in molybdate-free electrolytes. Also, the inhibiting effect of molybdate on the hypochlorite reduction could be further enhanced by adding low amounts of chromate to the electrolyte (see polarization curves in green in Fig. 3), while the activation towards hydrogen evolution remains as in the case of only molybdate present. The inhibiting and activating effects of the molybdate additive have been previously reported. However, those studies used higher molybdate concentrations ($>$40 mmol L$^{-1}$ Na$_2$MoO$_4$), leading to undesired enhancement in O$_2$ generation. Here, we show hypochlorite electroreduction hindrance with chromate (3 μmol L$^{-1}$) and molybdate (10 mmol L$^{-1}$).
in solution, having less negative cathode potential compared to the reaction performed in additive-free electrolyte. Note that although the XPS results indicate that Mo deposition is hindered by the presence of Cr, there is still an activation of HER in Fig. 3 in a Cr-containing electrolyte. To better understand this, the HER was investigated on ex situ formed films.

Figure 3. IR-corrected polarization curve for hydrogen evolution reaction (HER) in NaClO\(^{-}\)-containing electrolyte, as a function of the concentration of Na\(_2\)Cr\(_2\)O\(_7\) additive \((x)\) in the absence and presence of molybdate co-additive: (a) \(x = 0.5\) \(\mu\)mol L\(^{-1}\); (b) \(x = 3\) \(\mu\)mol L\(^{-1}\); (c) \(x = 13.5\) \(\mu\)mol L\(^{-1}\). Measurements were made at 2500 rpm at RT. Pre-polarization at \(-0.8\) V versus Ag/AgCl for 5 min before measuring the polarization curve from high to low current density, with constant purging by N\(_2\). The grey dashed line indicates the current density typically used to run the industrial chlorate process (300 mA cm\(^{-2}\)).
Ex situ formed catalytic cathodic films

The apparent activation of HER by Mo may be explained by either a catalytically active Mo oxide species or by continuous reduction of dissolved molybdate.\textsuperscript{17,19} To determine the source of the activating effect, polarization curves for hydrogen evolution on \textit{ex situ} coated cathodes were obtained in hypochlorite-containing electrolyte (Fig. 4). The electrodes with \textit{ex situ} formed films were removed from the depositing electrolyte while still polarized. Afterwards, the electrode was rinsed and transferred to the measuring electrolyte (NaOH 0.1 mol L\textsuperscript{-1} + NaCl 2 mol L\textsuperscript{-1} + NaClO 80 mmol L\textsuperscript{-1}). Insertion in the measuring electrolyte was done under open circuit conditions. As revealed by the pink curve in Fig. 4, the \textit{ex situ} formed films, deposited in chromate-containing electrolyte (13.5 μmol L\textsuperscript{-1} Na\textsubscript{2}Cr\textsubscript{2}O\textsubscript{7} only), are not stable. In fact the inhibiting effect on hypochlorite reduction disappears upon transfer (compare red curves in Figs 3(c) and 4). Conversely, the film deposited in a mixture of chromate and molybdate maintains the inhibiting effect on hypochlorite reduction after transfer. This clearly shows that molybdenum enhances the stability of the film in an alkaline environment under open circuit conditions, enhancing the corrosion protection greatly. Furthermore, for \textit{ex situ} coatings formed by Mo addition, the polarization curves resemble the trend observed for \textit{in situ} formed films (Fig. 2); that is, 10 mmol L\textsuperscript{-1} Mo improves the activity for HER most efficiently, indicating a true activation of HER by the formed films rather than a simultaneous continuous Mo reduction shifting the potential (Fig. 5).

Effect of Mo and Cr electrolyte in situ additions on the Faradaic efficiency for hydrogen and oxygen evolution

Faradaic efficiencies of hydrogen and oxygen evolution were measured under experimental conditions approaching the industrial ones, by increasing the temperature up to 80 °C and maintaining the bulk electrolyte at pH 6.5. To assess the role of additives, galvanostatic electrolysis with simultaneous MS measurements were performed. The effect of chromate and molybdate additives on the Faradaic efficiency towards hydrogen evolution, measured after 30 min of galvanostatic electrolysis, is depicted in Fig. 6 (the hypochlorite concentration in the electrolyte reached a constant value of ∼30 mmol L\textsuperscript{-1}, as shown in Supporting Information Fig. S6). The Faradaic efficiency towards hydrogen evolution is calculated as the ratio of the volumetric flow of hydrogen after 30 min of electrolysis with respect to the expected flow of hydrogen for the applied current (22.65 μL s\textsuperscript{-1}). The volumetric flow of H\textsubscript{2} as a function of time is shown in Supporting Information Fig. S7.

The results show that the Faradaic efficiency to hydrogen is only ∼65% in the absence of chromate and molybdate additives. This observation agrees well with the polarization curve in Fig. 3, where it is clearly shown that, in the absence of chromate or molybdate, hypochlorite anions can be electrochemically reduced on the Pt working electrode. The MS results in Fig. 6 also indicate that 0.5 μmol L\textsuperscript{-1} of chromate additive does not significantly affect the efficiency of hydrogen production on Pt in NaClO\textsuperscript{-containing electrolyte. With these low concentrations of chromate in solution, adding 80 mmol L\textsuperscript{-1} of molybdate induces a small enhancement, in line with previous reports.\textsuperscript{11}

By increasing the concentration of chromate in solution (i.e. 3 and 13.5 μmol L\textsuperscript{-1}), the efficiency towards hydrogen generation increases to >75%, as one would expect due to the inhibiting effect of chromate.\textsuperscript{10,12,20,46,48,51} Note that this is far below the Faradaic efficiency obtained industrially of close to 100%, then with electrolyte levels of ∼3–6 g L\textsuperscript{-1} Na\textsubscript{2}Cr\textsubscript{2}O\textsubscript{7}, corresponding to ∼10–20 mmol L\textsuperscript{-1} Na\textsubscript{2}Cr\textsubscript{2}O\textsubscript{7}.

Figure 6 shows that molybdate does not significantly affect the HER Faradaic efficiency on platinum in the NaClO-containing electrolyte. There is a small enhancement in the efficiency when 80 mmol L\textsuperscript{-1} molybdate is in the electrolyte, as shown by the blue bars in Fig. 6. However, > 40 mmol L\textsuperscript{-1} molybdate in solution has been shown to enhance O\textsubscript{2} formation,\textsuperscript{17} detrimentally affecting the anode efficiency and raising safety concerns (due to the risk of reaching an explosive mixture in the hydrogen-containing cell gas). These results allow us to unravel the oxygen formation and to evaluate the effect of molybdate on it.

The volumetric flow of oxygen generated during galvanostatic electrolysis as a function of the additives (Supporting Information Fig. S8) shows a sudden fast increase of the oxygen signal at the beginning of the experiment (t < 5 min) and a monotonic signal decrease after ~5 min. The behavior at about t < 5 min has been previously observed for the chemical decomposition of hypochlorite.\textsuperscript{15,52} There are several possible sources of oxygen in the chlorate electrolysis cell: coming from (i) anodic reactions so called electrochemically produced oxygen, or (ii) chemical heterogeneous decomposition of hypochlorite forming oxygen where the electrode surface catalyzes the oxygen formation, and
The MS results (Supporting Information Fig. S8) further show that the concentration of Na$_2$Cr$_2$O$_7$ does not significantly affect either the chemical decomposition of hypochlorite (i.e. no significant differences in the oxygen maximum at the beginning of the experiment) or the electrochemical oxygen production (depicted as a sharp decrease in the O$_2$ signal upon turning the current off). On the contrary, 80 mmol L$^{-1}$ of molybdate both promotes the chemical decomposition of NaClO (see Fig. 7) and enhances the electrochemically produced oxygen, seen as the drop in Supporting Information Fig. S7 once the current was switched off. Still, our MS results show that 10 mmol L$^{-1}$ molybdate in the electrolyte does not promote the formation of oxygen compared to the measurement without additives.

These results are in line with previous reports from our group, concluding that molybdate at low concentration (10 mmol L$^{-1}$) does not significantly affect the chemical decomposition of hypochlorite into oxygen.$^{32}$ Given the presented experimental data, chemical decomposition is avoided at a concentration of 10 mmol L$^{-1}$ molybdate (Fig. 7 and Supporting Information Fig. S9). Following hypochlorite concentration (Supporting Information Table S1) indicated that addition of 10 mmol L$^{-1}$ molybdate does not significantly affect the rate constant for the hypochlorite decomposition compared to the case without additives. However, in the case of 80 mmol L$^{-1}$ molybdate the rate constant for the hypochlorite decomposition was increased by $\sim$30%.

Both the overpotential for HER and the rate of unwanted oxygen evolution increase at high concentrations of molybdate, while the hydrogen Faradaic efficiency does not depend much on the concentration of molybdate (range 10–80 mmol L$^{-1}$). We show that by fine-tuning the molybdate concentration in the range 10–30 mmol L$^{-1}$ a high Faradaic efficiency and activity towards the HER can be obtained without promoting O$_2$ generation.

Coatings of Cr and Mo may be formed in situ or ex situ. The former has the advantage of a continuously refreshed electrode surface, where cracks etc. in the coating are mended during operation, as is the case with the chromate addition today. The latter allows a better control of the composition and morphology of the protective layer, and has shown promising results.$^{20}$ We show that Mo is important for the stability of the selective Cr layer.

CONCLUSIONS

This work has quantitatively addressed the effect of additives, namely Na$_2$Cr$_2$O$_7$ (in the micromolar range) and Na$_2$MoO$_4$ (in the millimolar range), on the production of H$_2$ and O$_2$ from NaClO-containing electrolytes using inert Pt as cathode material. Films formed from these additives were examined by SEM and EDX, where only the platinum surface could be recorded, as most likely the deposited films were of nanometer thickness. XPS studies of the as-deposited films revealed a partial reduction in Mo oxidation states compared to the electrolytic solution. Their oxidation states and oxidation state percentages were highly dependent on the deposition conditions (concentration of additives and temperature). In the presence of Cr it is likely that only trace levels of Mo are deposited as XPS could not detect any traces of Mo in such films.
Catalytic effects of molybdate and chromate

Still molybdate, particularly at electrolyte concentrations of 10 mmol L\(^{-1}\) Mo, has a strong activating effect towards HER on Pt and additionally enhances the stability of the chromium hydroxide film at open circuit. The electrocatalytic effect is also preserved when Cr and Mo are combined, leading to a shift of 260 mV to less negative potentials at current densities of 300 mA cm\(^{-2}\) compared to measurements in molybdate-free electrolytes.

The Faradaic efficiency towards hydrogen evolution can be enhanced by more than 15% in the presence of 3 μmol L\(^{-1}\) Na\(_2\)Cr\(_2\)O\(_7\)·Na\(_2\)MoO\(_4\) as a co-additive has a minor effect on the hydrogen Faradaic efficiency at a high concentration (80 mmol L\(^{-1}\)). However, such high concentration enhances the chemical and electrochemical reactions leading to O\(_2\), which is detrimental for the chloride process, whereas 10 mmol L\(^{-1}\) Mo does not affect homogeneous hypochlorite decomposition or oxygen generation in the cell.

Our finding allows us to optimize molybdate concentration further to obtain the highest hydrogen evolution efficiency and film stability while minimizing O\(_2\) generation even at micromolar chromate levels. The insights on the synergy of Cr–Mo could also be capitalized for photocatalytic hydrogen production and the degradation of industrial wastewater. However, the influence of the investigated additives on the reduction of chloride ions still remains to be investigated. Also, as it is unclear how the films containing Cr and Mo are formed, more investigations to enlighten this synergy are needed. Related to this is how to produce such films in an industrial process – in situ or ex situ. The findings presented in this paper may be applicable to the next generation of sodium chlorate process technology where a more active and stable cathode material is used which also requires a much higher electrolyte purity compared to the industrial standard today. Further elucidate the commercial viability of these results, further research should focus on applying more industrial conditions by, for instance, using electrolytes containing high ionic strength and disturbing agents such as calcium or sulfates.

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SUPPORTING INFORMATION

Supporting information may be found in the online version of this article.

REFERENCES


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