Electrochemical Investigation of the Hydrogen Evolution Reaction on Electrodeposited Films of Cr(OH)3 and Cr2O3 in Mild Alkaline Solutions

Downloaded from: https://research.chalmers.se, 2020-04-11 07:08 UTC

Citation for the original published paper (version of record):
Gomes, A., Simic, N., Wildlock, M. et al (2018)
Electrochemical Investigation of the Hydrogen Evolution Reaction on Electrodeposited Films of Cr(OH)3 and Cr2O3 in Mild Alkaline Solutions
Electrocatalysis, 9(3): 333-342
http://dx.doi.org/10.1007/s12678-017-0435-1

N.B. When citing this work, cite the original published paper.
Electrochemical Investigation of the Hydrogen Evolution Reaction on Electrodeposited Films of Cr(OH)$_3$ and Cr$_2$O$_3$ in Mild Alkaline Solutions

Adriano S. O. Gomes$^{1,2}$ · Nina Simic$^2$ · Mats Wildlock$^2$ · Anna Martinelli$^3$ · Elisabet Ahlberg$^1$

Abstract The hydrogen evolution reaction (HER) from water reduction is the main cathodic reaction in the sodium chlorate process. The reaction typically takes place on electrodes covered with a Cr(III) oxide-like film formed in situ by reduction of sodium dichromate in order to avoid reduction of hypochlorite and thereby increase the selectivity for the HER. However, the chemical structure of the Cr(III) oxide-like film is still under debate. In the present work, the kinetics of the HER were studied using titanium electrodes covered with electrodeposited Cr(OH)$_3$ or Cr$_2$O$_3$, which were characterized by means of scanning electron microscopy (SEM), energy-dispersive x-ray spectroscopy (EDX), x-ray photoelectron spectroscopy (XPS), and Raman spectroscopy. A clear difference in the morphology of the deposited surfaces was obtained, and the structure could be revealed with Raman spectroscopy. The kinetics for the HER were investigated using potentiodynamic and potentiostatic techniques. The results show that the first electron transfer is rate limiting and that the activity decreases in the order Cr$_2$O$_3$@Ti > bare Ti > Cr(OH)$_3$@Ti. The low activity obtained for Cr(OH)$_3$@Ti is discussed in terms of the involvement of structural water in the HER and the slow ligand exchange rate for water in Cr(III) complexes, while the high activity obtained for Cr$_2$O$_3$@Ti is rationalized by a surface area effect in combination with reduction of surface water and water in solution.

Keywords Electrocatalysis · Raman spectroscopy · Sodium chlorate · Electrodeposition

Introduction

The electrosynthesis of sodium chlorate is one of the most important industrial processes, especially for the pulp and paper industry [1]. The production of chlorate occurs via the electrolysis of sodium chloride and water (Eq. 1). Electricity is the main driving force of electrolysis and accounts for up to 70% of the total production cost [2]. The main by-products are heat and hydrogen gas, which can be re-used in the process or surroundings [2]. Chlorate is further reduced to form chlorine dioxide (Eq. 2), the state-of-the-art bleaching chemical for kraft pulp [1, 3].

\[
\begin{align*}
NaCl + 3H_2O & \overset{6e^-}{\rightarrow} NaClO_3 + 3H_2 \\
2NaClO_3 + H_2O_2 + H_2SO_4 & \rightarrow 2ClO_2 + O_2 + Na_2SO_4 + 2H_2O
\end{align*}
\]  

(1)
\(2\)  

The electrochemical and chemical reactions yielding chlorate have been well described in the literature [4–11]. The main half reactions are the cathodic reduction of water (Eq. 3) and the anodic chloride oxidation (Eq. 4). The formation of chlorate occurs in the bulk of the electrolyte as follows: Cl$_2$ immediately undergoes hydrolysis.
chlorate formation (Eq. 7); hypochlorous acid is in equilibrium with hypochlorite (Eq. 6) while chlorate is produced from a disproportionation reaction between hypochlorous acid and hypochlorite (Eq. 7). The optimum pH for the process is empirically known to be between 6 and 7 [12]. At this pH, the equilibrium in Eq. 6 is shifted to the left, keeping the hypochlorous acid in equilibrium with hypochlorite (Eq. 7). The optimum pH for the process is empirically known to be between 6 and 7 [12]. At this pH, the equilibrium in Eq. 6 is shifted to the left, keeping the hypochlorous acid in equilibrium with hypochlorite (Eq. 7).

2H2O + 2e−→H2 + 2OH− E° = −0.828 V vs.NHE (3)
2Cl−→Cl2 + 2e− E° = +1.360 V vs.NHE (4)
Cl2 + H2O→HClO + H+ + Cl− (bulk) (5)
HClO→H+ + ClO− (bulk, pKa = 7.5) (6)
2HClO + ClO−→ClO3− + Cl− + 2H+ (bulk) (7)

Chlorate is produced in undivided cells, using mild steel cathodes and dimensionally stable anodes (DSA) [2]. The electrolyte is rich in ions such as chloride, chlorate, and hypochlorite (HClO + ClO−) [2]. In this setup, water reduction competes with the cathodic reductions of chlorate and hypochlorite (Eqs. 8 and 9), which are thermodynamically favorable. Therefore, in chlorate electrolytes, the addition of sodium dichromate is necessary as it provides selectivity towards water reduction (note that the mild steel electrodes are not selective). Dichromate is cathodically reduced forming a chromium(III) oxide-like film, which inhibits the reactions given in Eqs. 8 and 9 [13, 14]. The composition and effects of such a film concerning chlorate electrochemistry has been investigated [4, 8, 15–17]. Other works, not directly related to the chlorate process, can be found in [18–20].

ClO− + H2O + 2e−→Cl− + 2OH− E° = +0.89 V vs NHE (8)
ClO3− + 3H2O + 6e−→6OH− + 6H+ E° = +1.45 V vs NHE (9)

The use of sodium dichromate, even though optimum from the process point of view, is undesirable owing to the presence of Cr in the hexavalent form (Cr(VI)), which is poisonous, carcinogenic, reprotoxic, and mutagenic. Comprehensive reviews on the toxicity of Cr(VI) can be found in [21, 22]. The removal of chromium(VI) from chlorate electrolytes has long been pursued, and is even more motivated now, since Cr(VI) compounds have been immobilized onto Ti and Fe electrodes in a Cr2O3 matrix, yielding a surface very selective towards water reduction in the presence of hypochlorite [28]. However, it was shown elsewhere that electrodeposited Cr2O3 completely prevents hypochlorite reduction on Ti electrodes [11] and, hence, the effects of MoO2 particles as presented in [28] cannot be precisely evaluated.

It turns out that to find a suitable replacement for sodium dichromate, a deeper knowledge of the selective mechanism is necessary. In this context, a recent paper focusing on Cr(OH)3 and Cr2O3 surfaces has shown that both are capable of blocking hypochlorite reduction, probably due to the p-type semi-conducting properties of Cr(III) oxide-like surfaces [11]. However, studies regarding water reduction on such films were not deeply investigated. With the aim to fill this gap, the present paper provides a comprehensive investigation of the mechanism and kinetics of water reduction on Cr(OH)3- and Cr2O3-electrodeposited films. Notably, the idea of depositing Cr(III) oxide-like films offers the opportunity to characterize them prior to electrochemical analysis, and thus interpret the results based on the films’ structure and composition.

Experimental

Electrode Preparation

Working rotating disk electrodes were home-made by molding Ti disks, with a 1-cm2 geometrical surface area, in epoxy, referred in the text as Ti. The electrodes were used as the substrate for electrodeposition of Cr(OH)3 and Cr2O3 films, henceforth referred to as Cr(OH)3@Ti and Cr2O3@Ti, respectively.

Prior to electrodeposition, Ti substrates were polished with SiC paper (Struers) and later with alumina slurries (OP-AN Struers) until mirror-like surfaces were obtained. To remove any contamination from the polishing steps, the electrodes were ultrasonicated in Milli-Q water for at least 10 min. The Ti electrodes were etched in a HF solution (3.5 mM in H2O, BASF) for 10 s, rinsed, and then directly immersed into the deposition solution.

Cr(OH)3 and Cr2O3 were electrodeposited from two different Cr(VI) solutions as previously described by Aguilar et al. [29] (Table 1). The electrolyte for Cr(OH)3 deposition consists of CrO3 (Sigma-Aldrich) and BaCO3 (Sigma-Aldrich). For Cr2O3 electrodeposition, the electrolyte consists of the same...
Electrocatalysis (2018) 9:333–342

Table 1 Composition of the electrolyte used for the deposition of Cr(III) oxide and hydroxide films [29]

| Electrolyte  | Cr(OH)₃ | Cr₂O₃ |
|--------------|---------|-------|
| CrO₃ (g/L)   | 364     | 364   |
| BaCO₃ (g/L)  | 9.1     | 9.1   |
| H₂SiF₆ (g/L) | –       | 1.8   |
| KNO₃ (g/L)   | –       | 2.3   |

CrO₃ and BaCO₃ with the addition of H₂SiF₆ (20–23.5 mM in H₂O; Sigma-Aldrich) and KNO₃ (Merck). The deposition potential was maintained at −1.60 V (vs Ag/AgCl) during a period of 30 min using a PGSTAT12 Potentiostat (Autolab). The excess of solution was rinsed several times with Milli-Q water, and the electrodes were stored immersed in Milli-Q water.

Electrochemical Experiments

All electrochemical measurements were performed using a Gamry600™ potentiostat. The three-electrode glassy cell configuration was used, where a high-surface area platinum mesh was used as counter-electrode and the reference electrode was a double-junction Ag/AgCl (KCl 3 M, Metrohm, E = +0.210 V vs nhe). Potential sweeps were recorded between −0.0 and −2.2 V (vs Ag/AgCl) at 50 mV s⁻¹ to investigate surface stability. Capacitive currents were measured between 0.0 and −0.8 V (vs Ag/AgCl) to estimate the capacitance of Cr(OH)₃− and Cr₂O₃-electrodeposited films.

Potentiostatic sweeps from −0.2 to −2.2 V (vs Ag/AgCl) in potential steps of −0.1 V were also performed. At the same time, electrochemical impedance spectroscopy (EIS) measurements were carried out at every potential. EIS frequencies varied between 100 kHz and 10 mHz. For impedance experiments, the working electrode was placed in the centre of a round platinum mesh to improve the electrochemical response.

Cyclic voltammetry (CV), potentiostatic sweeps, and EIS were performed in de-aerated 0.2 M Na₂SO₄ at pH 11. The pH adjustment was made with addition of NaOH; N₂ gas flow was maintained in the top of the cell to prevent dissolution of O₂ in the electrolyte during experiments.

Surface Characterization

The surface morphology and composition were investigated with scanning electron microscopy (SEM) and energy-dispersive X-ray spectroscopy (EDX) in a LEO Ultra 55 FEG SEM equipped with Oxford Inca EDX system. The molecular structure of the films was analyzed with Raman spectroscopy using an InVia Reflex spectrometer from Renishaw. The excitation source was the 532-nm line of a diode laser, which together with a grating of 2400 lines per millimeter gives a spectral resolution better than 1 cm⁻¹. The presented Raman spectra are the result of 10 scans with a duration of 10 s each, if not otherwise stated. X-ray diffraction (XRD) was performed using a Siemens D5000 Diffractometer with a Bragg-Brentano setup equipped with a Cu kα X-ray source and a scintillation detector from 20° to 63° at 0.100° intervals of 2θ, and cumulated count time of 4 s for each step.

Results and Discussion

Surface Characterization

In electrochemistry, the determination of surface composition and surface area is a prerequisite for a complete investigation of mechanisms and kinetic parameters. However, the investigation of Cr(III) films formed during the chlorate electrolysis was always limited by the uncertainty of the surface composition. For instance, for a long time it was believed that the reduction of chromate yields a hydrated Cr(OH)₃ film on platinum and gold [15] electrodes, but recently Hatch and Gewirth [20] determined with in situ Raman spectroscopy and atomic force microscopy (AFM) that the structure of chromate electrochemically reduced on gold is in fact a Cr₂O₃ thin film. In addition, a dual composition of the film had also been reported, with an inner Cr₂O₃-like structure covered by an outer Cr(OH)₃ film [9, 30].

As an alternative to in situ deposited films, ex situ electrodepositions of Cr(OH)₃ and Cr₂O₃ onto Ti RDE were performed in the present work. The electrodeposition method includes two electrolytes rich in CrO₃. The difference is the use of KNO₃ and H₂SiF₆ for obtaining Cr₂O₃. KNO₃ acts as an oxidizing agent [31] while fluorosilicates stabilizes the oxide phase [32]. It has been reported that NO₃⁻ and fluoride compounds catalyze the formation of the oxide phase [32]. Differences between Ti and Ti covered with either Cr(OH)₃@Ti or Cr₂O₃@Ti were first analyzed with an optical microscope (Fig. 1). In Fig. 1a, the pure Ti surface shows a smooth surface with scratches from the polishing, which are seen already with a ×50 magnification. The Cr(OH)₃@Ti film is shown in Fig. 1b and presents a light-gray smooth surface. On the other hand, Cr₂O₃@Ti consists of a dark, rough, and brittle layer (Fig. 1c). After drying, the Cr₂O₃ layer cracks and easily detaches from the electrode surface, revealing a black thin film bound to the substrate (Fig. 1d).

The morphology and chemical composition of the Cr(OH)₃@Ti and Cr₂O₃@Ti surfaces were further analyzed with SEM and EDX. The Cr(OH)₃ film displays a rather rough surface in the micrometer scale (Fig. 2a), while EDX analyses revealed that Cr and O are present in a ratio close to 1:3 as expected, and that the signal of the Ti substrate accounts for 15 at% of the total atomic composition (Table 2). By contrast, the Cr₂O₃ film presents plate-shaped crystals less than
1 μm long (Fig. 2b), and EDX measurements show larger amounts of Cr, compared with Cr(OH)₃ (Table 2). Still, a smaller amount of Ti, only 2 at%, was recorded during the Cr₂O₃@Ti analyses. EDX is not suitable for obtaining the absolute film thickness due to the interaction volume. However, lower signals from the substrate (under the same acceleration voltage) indicate that the Cr₂O₃ layer is thicker than the Cr(OH)₃ layer.

Raman Spectroscopy

Even though EDX analyses have shown the expected Cr/O ratio for both the Cr(OH)₃@Ti and Cr₂O₃@Ti phases (Table 2), this is not sufficient to characterize the molecular structure of the films, which we have investigated by Raman spectroscopy. The experiments were carried out by starting with low laser power as suggested in a recent study [33], and the results are summarized in Figs. 3 and 4.

Figure 3 presents a collection of spectra recorded at increasing laser power from an electrode covered with Cr(OH)₃. With the lowest laser power (i.e., 0.35 mW), a weak signature is observed at ≈850 cm⁻¹, which is characteristic for amorphous Cr(OH)₃ [33, 34]. This peak is still visible when a higher laser power is used, although other peaks also appear whose presence is attributed to the in situ formation of Cr₂O₃ as a result of local heating [33]. Compared to the Raman spectra previously recorded for nanoparticles of amorphous Cr(OH)₃ (see, e.g., Fig. 6 in [33]), the spectra recorded from Cr(OH)₃@Ti appear more noisy (at comparable experimental conditions), which we attribute to the deposited film being very thin. In addition, the spectra recorded at low laser power (e.g., 0.35 and 0.70 mW) display a weak or inexistent peak at 530 cm⁻¹ as compared to the intensity at 850 cm⁻¹, a feature that, based on the assignment proposed in [33], suggests films of Cr(OH)₃ with partially condensed –OH groups. Moreover, Raman spectra recorded at different spots but using the same laser power (3.50 mW) show some spatial inhomogeneities with a variable proportion of Cr₂O₃ versus C(OH)₃ (see Fig. S1 in Supporting Information). We believe that these differences can arise from a variable thickness of the film, and thus different responses to the local heating induced by the laser.

Figure 4 shows the Raman spectra for Ti covered with Cr₂O₃ recorded at the same experimental conditions used in Fig. 3. For increasing laser power, the peaks characteristic of Cr₂O₃ [34–38] gain intensity, as already observed for nanoparticles of Cr₂O₃ [33]. However, differently from the case of nanoparticles [33] the Raman spectrum of Cr₂O₃@Ti recorded with a laser power of 0.7 mW looks featureless, which could be attributed to the film here investigated being amorphous (see x-ray diffractogram in Supporting Information, Fig. S2) and consisting in elongated crystals with a length of ~600 nm and a width of less than 100 nm (different from the case of ~50-nm-sized nanoparticles grouped together into large particles studied in [33]).

It is known that the electrodeposition of so-called black chromium yields amorphous Cr(III) oxide films/deposits [29, 32, 39], whose actual composition is difficult to determine [32]. XPS has thrown some light towards characterizing this type of films [29, 32, 39, 40], showing either a pure Cr₂O₃ phase [29, 39], a mixture of Cr₂O₃ and Cr(OH)₃ [40], or a much more complex matrix of different Cr(III) oxyhydroxides [32]. The XPS investigation made in the present work (Fig. S3) has revealed that the surfaces correspond to...
Cr(III) oxide-like materials, with some percentage of Cr metal. The complexity of the C1s reference peaks, however, hinders a precise peak positioning and a distinction between Cr(OH)₃ and Cr₂O₃ was not possible due to the proximity of the corresponding binding energies [41, 42]. Thus, we assume that the temperature increase induced by light absorption, in this case, promotes amorphous-Cr₂O₃ (or a mixture of oxides) to transform into the more thermodynamically stable crystalline-Cr₂O₃.

To confirm that the transition from amorphous to crystalline Cr₂O₃ was caused by light adsorption at laser intensities higher than 0.7 mW, experiments using 0.7 mW of laser power were carried out at scan times of 50 s (data not shown).

**Table 2** Chemical composition of Ti electrodes covered with electrodeposited Cr(OH)₃ and Cr₂O₃

| Surface   | Cr (at%) | O (at%) | Ti (at%) |
|-----------|----------|---------|----------|
| Cr₂O₃     | 37 ± 5   | 47 ± 10 | 2 ± 1    |
| Cr(OH)₃   | 20 ± 1   | 55 ± 8  | 17 ± 7   |

**Fig. 3** Raman spectra of Cr(OH)₃ electrodeposited onto Ti recorded at a different laser power from the same spot. The line at 855 cm⁻¹ marks the characteristic peak for amorphous Cr(OH)₃.

**Fig. 4** Raman spectra of electrodeposited Cr₂O₃ onto Ti recorded at four different laser powers.
However, the Raman spectra showed no peaks related to Cr$_2$O$_3$, indicating that the temperature increase by Raman laser absorption at 0.7 mW is not enough to crystallize amorphous Cr$_2$O$_3$. Also, a pure temperature conversion was studied by electrodepositing Cr$_2$O$_3$ onto a Ti disk, which was further inserted in an oven and heated up to 900 °C for 1 h. The results are summarized in Fig. S4 (Supporting Information), and the Raman spectra obtained for the heat-treated sample show the Cr$_2$O$_3$ features already with 0.7 mW laser energy.

**Electrode Surface Area Estimation**

The capacitance of the Cr(OH)$_3$@Ti and Cr$_2$O$_3$@Ti electrodes was measured by means of cyclic voltammetry in the non-faradaic region of the voltammogram, between −0.8 and 0.0 V. Capacitance measurements were used to estimate differences in the active surface area of the electrodes. In Fig. 5, the capacitive currents ($I_{cap}$) are plotted as a function of sweep rate (v). Observe that the currents obtained for Cr$_2$O$_3$ (left y-axis) are a thousand times higher than for Cr(OH)$_3$ (right y-axis). The slopes of the curves give the capacitances associated with the Cr(OH)$_3$ (squares) and Cr$_2$O$_3$ (diamonds) surfaces. Note that the capacitive currents for the oxide are three orders of magnitude higher than for hydroxide; see also the inset in Fig. 5. The Cr$_2$O$_3$ layer displays a capacitance of $1.84 \times 10^{-3}$ F, while the capacitance of Cr(OH)$_3$ is $3.83 \times 10^{-6}$ F. It will be seen later that the reduction currents found using Cr$_2$O$_3$@Ti electrodes are larger than for Cr(OH)$_3$@Ti electrodes, and that the surface area effect cannot be ignored in further discussions.

**Potential sweeps**

Potential sweeps performed between 0.0 and −2.2 V (vs Ag/AgCl) at 50 mV s$^{-1}$ were used to evaluate any possible changes of the Cr(OH)$_3$ and Cr$_2$O$_3$ films. Water reduction was the only reaction observed in this potential window, with no further reduction of the Cr(III) or re-oxidation back to chromium(VI) (see Supporting Information, Fig. S5). Cyclic voltammograms at 5 mV s$^{-1}$ were performed to investigate the kinetics of water reduction. The capacitive response was calculated and subtracted from the original data, to allow for a comparison of the onset potential for water reduction. The capacitive currents were calculated using Eq. 10, where j is the current density, $\nu$ the sweep rate, $C_d$ the capacitance, and $R_0$ the solution resistance. The background was fitted to a straight line and subtracted. The corrected currents for water reduction at Ti, Cr(OH)$_3$@Ti, and Cr$_2$O$_3$@Ti are summarized in Fig. 6.

$$j = \nu C_d \left[1 - \exp \left( -\frac{t}{R_0 C_d} \right) \right]$$ (10)

From Fig. 6, it is noticeable that the overpotential for water reduction increases in the order Cr$_2$O$_3$@Ti < Ti < Cr(OH)$_3$@Ti. At -2 mA, the difference between Cr(OH)$_3$@Ti and Cr$_2$O$_3$@Ti is approximately 0.36 V. Ti sits in the middle, indicating that the chromium oxide layer activates the electrode towards water reduction, while the chromium hydroxide deactivates it.

**Steady-State Experiments**

Tafel slopes and kinetic parameters could not be precisely evaluated based on the potential sweeps, due to the high capacitance. Therefore, potentiostatic measurements were used instead. For these experiments, freshly prepared Cr(OH)$_3$@Ti and Cr$_2$O$_3$@Ti electrodes were used. The steady-state current densities ($j_{ss}$) were measured together with impedance (EIS), at potentials for which water reduction takes place.

The $j_{ss}$ versus potential measurements using Ti, Cr(OH)$_3$@Ti, and Cr$_2$O$_3$@Ti are presented in Fig. 7 (symbols). The dotted lines show the data fitting with the kinetic current equation (Eq. 9). It is immediately seen that Fig. 7 reproduces results presented in Fig. 6, i.e., Cr$_2$O$_3$ is more active towards water reduction than Cr(OH)$_3$. To ensure a measurable difference between Cr(OH)$_3$ and Cr$_2$O$_3$, the experiments were repeated several times (Table 3). By using Eq. 11, the Tafel slopes, charge transfer coefficients (\alpha), and rate constant (k) were calculated and summarized in Table 3, where the number of measurements for each electrode and the uncertainty of the data are also shown.

$$I = -nFADe^{-\frac{\alpha F \eta}{RT}}$$ (11)
Electrochemical Impedance Spectroscopy

Complex plane plots from 100 kHz to 10 mHz were obtained for every applied potential during the j_{ss} measurements. Complex plane plots obtained for Cr(OH)$_3$@Ti and Cr$_2$O$_3$@Ti at each potential are summarized in Fig. 8a, b, respectively. The impedances observed at the highest frequencies were attributed to phase shifts in the instrument and will not be further discussed. The equivalent circuit used to fit the data is given in Fig. 8c. In Fig. 8c, $R_0$ is the solution resistance, CPE are constant phase elements, and $R_1$ and $R_{ct}$ are resistances. For the oxide, a phase shift independent of potential is observed prior to the semi-circle. This phase shift is probably due to the porosity of the material. In both cases, Cr(OH)$_3$ and Cr$_2$O$_3$, CPE1 and $R_1$ denote mass transport within the films. While CPE2 and $R_{ct}$ arise from electron transfer, with $R_{ct}$ being the charge transfer resistance.

The faradaic impedance for a kinetically controlled process with only one adsorbed intermediate can be described as in Eq. 12 [43]. Assuming that the reaction mechanism has no coverage dependence with potential, the last term of the equation can be eliminated and the impedance is directly proportional to the $R_{ct}$.

\[
\frac{1}{Z_F} = \frac{1}{R_{ct}} + \frac{\partial i}{\partial \theta} \frac{\Delta \theta}{\Delta E}
\]  

(12)

By plotting log $1/R_{ct}$ versus E, as presented in Fig. 9, a Tafel-like curve is obtained. Linear fits revealed slopes of 266 and 170 mV dec$^{-1}$ for Cr$_2$O$_3$ and Cr(OH)$_3$, respectively. These results are in agreement with Tafel slopes found using the steady-state current measurements in Table 3.

Reaction Order

The electrochemical reaction orders with respect to OH$^-$ ions were investigated. The plot presented in Fig. 10 was constructed by varying the pH from 9.5 to 12.5, by NaOH addition, and further acidifying back to 9.5 with H$_2$SO$_4$, in 0.5 pH steps. The currents were measured at $-1.5$ V (vs Ag/AgCl). Cr(OH)$_3$ presents a clear change in slope, indicating a change in the reaction order from zero to $-0.3$. Opposite to that, results found for Cr$_2$O$_3$ show no change in the reaction order in the pH region investigated. The result from acidifying the solution did not show any dramatic discrepancies and was omitted.

The HER Mechanism

Tafel Slopes

The rate-determining step (rds) for the HER on Cr(OH)$_3$ and Cr$_2$O$_3$ films was determined by the Tafel slopes. The Tafel slopes obtained using Cr$_2$O$_3$@Ti and Cr(OH)$_3$@Ti were, respectively, $-239 \pm 27$ and $-145 \pm 28$ mV dec$^{-1}$ (Table 3), indicating that the first electron transfer is the rds at both surfaces. The low $\alpha$ values indicate that the transition state is shifted to the right on the reaction coordinate axis, and the structure of the activated complex is closer to H ads than H$_2$O.

The $\alpha$ value for the Cr(OH)$_3$ film is higher than for the Cr$_2$O$_3$ film, $\sim 0.4$ compared with $\sim 0.25$. A possible explanation for the difference in the transfer coefficient is that water is integrated as part of the Cr(OH)$_3$ film and the film itself is probably involved in the reduction reaction. In this way, the
activated complex has an intermediate state between water and H\textsubscript{ads}. For the Cr\textsubscript{2}O\textsubscript{3} film, on the other hand, the reaction occurs at the outmost layer and surface water or water from the solution is involved in the reaction. Therefore, the activated complex becomes closer in structure to the fully reduced form, H\textsubscript{ads}.

Acid-Base Properties

The mechanism for the HER on oxides/hydroxides can be discussed based on the acid-base properties of the surface. The acid-base equilibrium for oxides in solution is described by the 2pK model \[44\] as given in Eq. 13. The surface charge of an oxide in solution is governed by the solution pH, and it can be positive, neutral (zero), or negative. The charge is controlled by either protonation or deprotonation of surface oxygen atoms. For Cr\textsubscript{2}O\textsubscript{3} and Cr(OH)\textsubscript{3}, surface charges are not well defined and different numbers are obtained depending on conditions according to Kosmulski \[45\]. The value for Cr(hydr)oxides is between 7.9 and 10.3. It is difficult to say that there should be a difference between the oxide and the hydroxide, but commonly hydroxides have higher pzc compared to the oxide. If this is the case, the oxide is negative in the entire pH range studied 9.5–12.5, while the hydroxide may have a change in the surface charge. The reaction order shows no dependence for the oxide and for the hydroxide in the low pH range. Hence, a correlation between different α values observed for the hydroxide and for the oxide cannot be made.

\begin{equation}
\begin{align*}
\equiv\text{Cr}^{-}+\text{OH}^{2-} & \leftrightarrow \equiv\text{Cr}^{-}\text{OH} + \text{H}_{2}\text{O}^{+} \quad K_{a1} \\
\equiv\text{Cr}^{-}\text{OH} + \text{H}_{2}\text{O}^{+} & \leftrightarrow \equiv\text{Cr}^{-}\text{O}^{-} + \text{H}_{3}\text{O}^{+} \quad K_{a2}
\end{align*}
\end{equation}

Hydrogen evolution proceeds as shown in Eqs. 14–16. First, adsorption of a proton accompanied with an electron transfer yields a $^*$ – OH\textsubscript{2} group (Eq. 14), and hydrogen evolution occurs either by recombination of 2 $^*$ – OH\textsubscript{2} groups (Eq. 15) or by the transfer of a second electron as in Eq. 16. This mechanism was proposed by Hedenstedt et al. \[46, 47\] on

![Complex plane plots of Cr(OH)\textsubscript{3}@Ti (a) and Cr\textsubscript{2}O\textsubscript{3}@Ti (b). Color lines show experimental data fitted (thin black lines) with the equivalent circuit in (c)](image)

### Table 3 Kinetic parameters for water reduction on both Cr (III) films electrodeposited

| Number of repetitions | Tafel slope\(\text{(mV dec}^{-1}\) | Charge transfer coef. (\(\alpha\)) | Reaction rate (\(k\)) (mol cm\(^{-2}\) s\(^{-1}\)) |
|-----------------------|-----------------|-----------------|-----------------|
| Cr\textsubscript{2}O\textsubscript{3} | 5               | - 239 (± 27)    | 0.25 (± 0.03)   | 2.3 (± 1.5) \times 10\(^{-10}\) |
| Cr(OH)\textsubscript{3} | 6               | - 145 (± 28)    | 0.41 (± 0.06)   | 9.9 (± 12) \times 10\(^{-13}\) |
| Ti                    | 1               | - 180           | 0.31            | 8.0 \times 10\(^{-12}\) |

\(a\) Experimentally obtained

\(b\) Simulated values
Fe oxides, and used by Larses et al. [48] to investigate the water reduction mechanism on Fe\(_{1-x}\)Cr\(_x\)OOH and Cr(OH)\(_3\) electrodes. In [48], phase diagrams made using density function theory (DFT) demonstrated that M-OH is the most likely structure under HER conditions. Interestingly, they found that the OH coverage increases with increasing Cr content. Taking that into account, a clear distinction between Cr(OH)\(_3\) and Cr\(_2\)O\(_3\) cannot be made in the acid-base or reaction order point of view. We then assume that the mechanism for HER is the same and follows what is presented in Eqs. 13–16.

\[
\equiv \text{Cr}^-\text{OH}^+ + e^- \leftrightarrow \equiv \text{Cr}^-\text{OH}_2^+ \tag{14}
\]

\[
2\equiv \text{Cr}^-\text{OH}_2 \leftrightarrow \equiv \text{Cr}^-\text{OH} + \text{H}_2 \tag{15}
\]

\[
\equiv \text{Cr}^-\text{OH}_2 + \text{H}_2\text{O} + e^- \rightarrow \equiv \text{Cr}^-\text{OH} + \text{H}_2 + \text{OH}^- \tag{16}
\]

**Reaction Kinetics**

The kinetics of the HER were investigated by potential sweeps and steady-state polarization (Figs. 6 and 7). Although the reaction mechanism follows the same path on oxide and hydroxide, the overpotential for the HER is the lowest in the presence of Cr\(_2\)O\(_3\), and the highest when Ti is covered with Cr(OH)\(_3\). The lower overpotential and low \(\alpha\) value observed on Cr\(_2\)O\(_3@\)Ti can be rationalized to be due to the larger surface area and the lack of water in the Cr\(_2\)O\(_3\) crystallites. On the other hand, as water in the Cr(OH)\(_3\) structure is involved in the HER, water has to be replaced by water molecules from the solution, otherwise the film will be destroyed. However, this is hampered by the slow ligand exchange of Cr(OH)\(_3\) as described by Tang et al. [49] and Rustad and Casey [50]. This is corroborated by the results presented by Larses et al. [48], who showed increased overpotentials for the HER with increasing Cr content in the Fe\(_{1-x}\)Cr\(_x\)OOH structure.

**Conclusion**

Hydrogen evolution was investigated on Ti electrodes covered with either Cr(OH)\(_3\) or Cr\(_2\)O\(_3\). The main outcomes of this work are as follows:

- The composition of the electrodeposited films was revealed by Raman spectroscopy.
- Cr\(_2\)O\(_3\) displays lower overpotentials towards water reduction compared to both Cr(OH)\(_3\) and bare Ti electrodes.
- For Cr(OH)\(_3\), on the other hand, the HER kinetics are slower resulting in the largest overpotentials.
- Structural water is involved in the HER reaction on Cr(OH)\(_3\), and the slow kinetics are explained by the slow water exchange rate for Cr(III) complexes.
- On Cr\(_2\)O\(_3\), only surface water or water from solution is involved in the reaction, and the low overpotential was rationalized to be due to the high surface area.

This study has shown that in the search for a replacement for Cr(VI) in the chlorate process, both the composition and the morphology of the deposited layer need to be considered to avoid increasing overpotentials for the HER and subsequent environmental issues related to the resulting increase in energy consumption.

**Acknowledgements**

Financial support from the Swedish research council, 621-2010-4035, is gratefully acknowledged.
Open Access This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made.

References

1. H. Vogt, J. Balej, J.E. Bennett, P. Wintzer, S.A. Sheikh, P. Gallone, S. Vasudevan, K. Pelin, Chlorine oxides and chlorine oxygen acids, in: Ullman’s Encyclopedia of Industrial Chemistry (Wiley-VCH Verlag GmbH & Co. KGaA, 2010)
2. A. Cornell, Encyclopedia of applied electrochemistry (Springer-Verlag, New York, 2014)
3. P. Axegård, E. Bergnor, Environmental performance of modern ECP bleaching (TAPPI IPBC, Portland, 2011)
4. G. Lindbergh, D. Simonsson, J. Electrochem. Soc. 3094, 137 (1990)
5. A. Cornell, D. Simonsson, J. Electrochem. Soc. 3123, 140 (1993)
6. A. Cornell, B. Hakansson, G. Lindbergh, J. Electrochem. Soc. D6, 150 (2003)
7. A. Cornell, B. Hakansson, G. Lindbergh, Electrochim. Acta 473, 48 (2003)
8. G. Lindbergh, D. Simonsson, Electrochim. Acta 1985, 36 (1991)
9. A. Cornell, G. Lindbergh, D. Simonsson, Electrochim. Acta 1873, 37 (1992)
10. J. Wullf, A. Cornell, J. Appl. Electrochem. 181, 37 (2007)
11. K. Hedenstedt, A.S.O. Gomes, M. Busch, E. Ahlberg, Electrocatalysis 326, 7 (2016)
12. B.V. Tilak, K. Viswanathan, C.G. Rader, J. Electrochem. Soc. 1228, 128 (1981)
13. J. Landin, Förfaringsätt vid framställning af klorater och perklorater, K.p.-o. Registreringsverket, Editor (Sweden, 1892)
14. C. Wagner, J. Electrochem. Soc. 181, 101 (1954)
15. A.A. Tidblad, G. Lindbergh, Electrochim. Acta 1605, 36 (1991)
16. N. Krstajić, V. Nakić, M. Spasojević, J. Appl. Electrochem. 77, 17 (1987)
17. A. Ahlberg Tidblad, J. Märtensson, Electrochem. Acta 389, 42 (1997)
18. W.J. Clark, R.L. McCreery, J. Electrochem. Soc. B379, 149 (2002)
19. I.M. Kolthoff, A.M. Shams El Din, J. Phys. Chem. 1564, 60 (1956)
20. J.J. Hatch, A.A. Gewirth, J. Electrochem. Soc. D497, 156 (2009)
21. A. Leonárd, R.R. Lauwerys, Mutat. Res. 227, 76 (1980)
22. R. Saha, R. Nandi, B. Saha, J. Coord. Chem. 1782, 64 (2011)
23. REACH, No 348/2013 (2013)
24. J. Gustavsson, L. Nylen, A. Cornell, J. Appl. Electrochem. 1529, 40 (2010)
25. L. Nylen, J. Gustavsson, A. Cornell, J. Electrochem. Soc. E136, 155 (2008)
26. J. Gustavsson, G. Lindbergh, A. Cornell, Int. J. Hydrog. Energy 9496, 37 (2012)
27. M. Li, Z. Twardowski, F. Mok, N. Tam, J. Appl. Electrochem. 499, 37 (2007)
28. U.C. Lačnjevec, B.M. Jović, L.M. Gajić-Krstajić, J. Kovač, V.D. Jović, N.V. Krstajić, Electrochim. Acta 34, 96 (2013)
29. M. Aguilar, E. Barrera, M. Palomar-Pardavé, L. Huerta, S. Muhl, J. Non-Cryst. Solids 31, 329 (2003)
30. H. Tsuchiya, S. Fujimoto, O. Chihara, T. Shibata, Electrochim. Acta 4357, 47 (2002)
31. E. Barrera, I. Gonzalez, T. Viveros, Sol. Energy Mater. Sol. Cells 69, 51 (1998)
32. N. Vasudevan, V.K.W. Grips, I. Rajagopalan, Surf. Technol. 119, 14 (1979)
33. A.S.O. Gomes, N. Yaghini, A. Martinelli, E. Ahlberg, J. Raman Spectrosc. 48, 1256 (2017)
34. F.D. Hardcastle, I.E. Wachs, J. Mol. Catal. 173, 46 (1988)
35. P. Li, Z. Zhou, H. Xu, Y. Zhang, Thermochim. Acta 71, 544 (2012)
36. S. Khamilč, E. Manikandan, B.D. Ngom, J. Electroanal. Chem. 350, 37 (2011)
37. J. Yang, H.F. Cheng, W.N. Martens, R.L. Frost, J. Raman Spectrosc. 1069, 42 (2011)
38. J. Yang, W.N. Martens, R.L. Frost, J. Raman Spectrosc. 1142, 42 (2011)
39. M. Aguilar-Sánchez, M. Palomar-Pardavé, M. Romero-Romo, M.T. Ramirez-Silva, E. Barrera, B.R. Scharifker, J. Electroanal. Chem. 128, 647 (2010)
40. S. Surviliene, A. Cesuniene, V. Jasulaitiene, I. Jureviciute, Appl. Surf. Sci. 837, 324 (2015)
41. E. Desimoni, C. Malitesta, P.G. Zambonin, J.C. Riviere, Surf. Interface Anal. 173, 13 (1988)
42. Z.A. Hamid, Surf. Coat. Technol. 3442, 203 (2009)
43. E. Ahlberg, H. Anderson, Acta Chem. Scand. 1063, 47 (1993)
44. D.A. Dzombak, F.M.M. Morel, Surface complexation modeling: hydrous ferric oxide (Wiley, New York, 1990)
45. M. Kosmulski, Adv. Colloid Interf. Sci. 69, 51 (1998)
46. J. Gustavsson, L. Nylen, A. Cornell, J. Appl. Electrochem. 1529, 40 (2010)
47. L. Nylen, J. Gustavsson, A. Cornell, J. Electrochem. Soc. E136, 155 (2008)
48. J. Gustavsson, G. Lindbergh, A. Cornell, Int. J. Hydrog. Energy 9496, 37 (2012)
49. M. Li, Z. Twardowski, F. Mok, N. Tam, J. Appl. Electrochem. 499, 37 (2007)
50. Y.Z. Tang, F.M. Michel, L.H. Zhang, R. Harrington, J.B. Parise, R.J. Reeder, Chem. Mater. 3589, 22 (2010)
51. J.R. Rustad, W.H. Casey, J. Phys. Chem. B 7107, 110 (2006)