Chlorate Electrochemical Removal from Aqueous Media Based on a Possible Autocatalytic Mechanism

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The electrochemical chlorate reduction at the Pt electrode in 0.5 M H2SO4 deaerated solutions has been studied using potentiostatic steady-state voltammetry. The kinetics parameters (Tafel slope, charge transfer coefficient, current density, and reaction order) were evaluated in function of chlorate concentration (1x10^{-2} - 0.2 M KClO3). The process of chlorate reduction is a complex one that implies two charge transfer controlled steps with formation of free radicals and an extent potential region controlled by the concentration polarization. The current density dependence of chlorate concentration tends to an exponential growth at concentration ≥ 0.1 M KClO3 and becomes exponential in the conditions of the Cl\textsuperscript{−}/Cl\textsuperscript{2−} catalyst system presence. In the second charge transfer, a surface reaction between free radical ·Cl\textsuperscript{−} and platinum electrode with formation of complex anions PtCl\textsubscript{4}\textsuperscript{2−} and PtCl\textsubscript{6}\textsuperscript{2−} is responsible for the rapid increase of the reaction rate.

Keywords: electrochemical reduction, platinum electrode, acid solution

Air, soil, and water pollution are very important issues for environmental protection and ecology, also due to their toxic impact on human bodies [1-3]. Different type of pollutants or wastes [4-7] as heavy metals [8-10], pesticides [11-13], pharmaceutical waste [14], other organic or biological [15-18] contaminants could be incriminated and specific methods for their detection are already optimized and developed, as well as efficient methods for their degradation or removal.

The chlorates are produced at a large scale by the chemical industry and used in a wide varieties of applications. In general, oxyanions of chlorine result into environment from different sources, including the manufacture and use, such as disinfection [19,20], bleaching agents [21], fuel cells [22]. Many recent studies involve the remediation of chlorine species [23,24] as is perchlorate in water. A photoelectrochemical mechanism of chlorate ion reduction was also suggested [25].

Chlorite ions appear as undesirable by-products when chlorine dioxide (ClO\textsubscript{2}) is used for water disinfection. Chlorate dioxide is a highly energetic molecule and becomes a free radical even in diluted aqueous solutions. Chlorate ion is produced through the reaction of residual chloride (ClO\textsubscript{2}\textsuperscript{−}) and free chlorine or hypochlorous acid during secondary disinfection. The overall reactions that describe chlorate formation are as follow:

\begin{equation}
\text{ClO}_3^- + Cl_2 + H_2O = ClO_2^- + 2Cl^- + 2H^+
\end{equation}

\begin{equation}
\text{ClO}_3^- + HOCI = ClO_2^- + Cl^- + H^+
\end{equation}

Also, chlorite dioxide disproportionatate under alkaline condition (pH>9) to chlorite and chlorate, according to the following reaction:

\begin{equation}
2\text{ClO}_2^- + 2OH^- = ClO_2^- + ClO_3^- + H_2O
\end{equation}

Chlorate and especially perchlorate ions are considered pollutants for drinking water, being recognized as poisons with a significant risk for health [26-30]. The levels of chlorate may be lowered by reduction to chloride ion, applying biological, chemical, and electrochemical treatment methods.

There are specialized microorganisms that can couple growth to the anaerobic reduction of chlorate or perchlorate to chloride. The processes are based on conversion with microorganisms that use chlorate or perchlorate as electron acceptor and organic compound as electron donor and carbon sources. Numerous investigations on the use of the biological treatment to remove chlorate and perchlorate have been reported in the last twenty years [12, 29-35].

An attractive alternative electron donor may be the hydrogen gas. It has proven its value as a reducing agent in waste water and drinking water treatment [31, 36]. Ferrous salts were reported to remove chlorine disinfection byproducts [37].

It is possible to reduce (per)chlorate to chloride by using the electrochemical treatment, but this method was not studied very much. Researches on perchlorate reduction have used a number of different materials as cathodes, including platinum, tungsten, ruthenium, titanium, aluminum, and carbon doped with chromium (III) oxide or aluminum oxide. However, the method has not been implemented yet in the potable water industry [28].

Different electrodes were used for electrochemical reduction of chlorate ion: a hybrid material, for detection of halogenate ion species in aqueous solutions, modified with: electroactive silver nanoparticles [38], titanium [24,39], carbon [40], boron doped diamond [41], or dimensionally stable anodes for chlorate electrolysis [42]. An autocatalytic chlorate-triiodide reaction was investigated in acid and neutral media [43].

The electrochemical reduction of chlorate becomes a theoretical interest, because its reduction up to chloride ion implies six electrons and therefore more intermediate
steps. The aim of this work is to establish the kinetics of the electrochemical reduction of chlorate at a platinum electrode in acid medium and to suggest a probable reaction mechanism.

**Experimental part**

**Material and methods**

The experimental studies consisting in the recording of the cathodic polarization curves were performed in a conventional three-compartment electrochemical cell separated by fretted glass disk. The working electrodes, having the geometrical area of 0.06 cm², were made of polycrystalline platinum. Before the measurements, the electrode was treated, being boiled in 50% nitric acid solution and washed with double distilled water. A platinum sheet with an area of 1 cm² was used as counter electrode and a saturated calomel electrode (SCE) - as reference electrode versus which are reported the potentials in the work.

The solutions consisting of 0.5 M H₂SO₄ with various concentrations of KClO₃ (1x10⁻⁴, 1x10⁻³, 0.01, 0.1, and 0.2 M), were prepared from reagents of Merck purity and double distilled water. Before the measurements, the solutions were non-aerated with purified nitrogen for one hour. All measurements were carried out at room temperature (23±1°C).

The polarization curves were recorded using potentiostatic steady-state voltammetry, point by point at 60 s intervals, in the range of potential from +440 to -520 mV/SCE, using a PS2 potentiostat.

**Results and discussions**

The cathodic potentiostatic polarization curves of platinum electrode in 0.5 M H₂SO₄, in absence and in presence of various chlorate concentrations are shown in figure 1.

![Figure 1: Cathodic polarization curves at platinum electrode in 0.5 M H₂SO₄ solutions in absence and in presence of various KClO₃ concentrations (1 – Aerated solutions; 2-6 – Deaerated solutions)](http://www.revistadechimie.ro)

According to figure 1, the curves of the chlorate reduction are shifted, showing more positive potential values with increasing chlorate concentration. The polarization curves recorded in presence of chlorate are extended on a potential domain of about 1000 mV and show two charge transfer control regions, separated by an extended limiting current region. The Tafel behavior of the two regions is given in figure 3a and 3b, respectively. The Tafel parameters (Tafel slope, b and charge transfer coefficient, α), corresponding to the two Tafel regions - (a) and (b) - are given in table 1.

![Figure 3: Tafel behavior of the chlorate reduction reaction in 0.5 M H₂SO₄ solutions at Pt electrode and various concentrations of KClO₃: 2-1x10⁻⁴ M; 3-1x10⁻³ M; 4-0.01 M; 5-0.1 M; 6-0.2 M, (a), (b) - the first and the second step controlled by the charge transfer polarization](http://www.revistadechimie.ro)

**Table 1**

<table>
<thead>
<tr>
<th>Step</th>
<th>KClO₃conc→ Parameters</th>
<th>1x10⁻⁴M</th>
<th>1x10⁻³M</th>
<th>0.01 M</th>
<th>0.1 M</th>
<th>0.2 M</th>
</tr>
</thead>
<tbody>
<tr>
<td>(a)</td>
<td>-b, mV.xdecade⁻¹</td>
<td>59</td>
<td>100</td>
<td>116</td>
<td>123</td>
<td>127</td>
</tr>
<tr>
<td></td>
<td>α</td>
<td>0.596</td>
<td>0.590</td>
<td>0.538</td>
<td>0.450</td>
<td>0.437</td>
</tr>
<tr>
<td>(b)</td>
<td>-δ, mV.xdecade⁻¹</td>
<td>49</td>
<td>48</td>
<td>47</td>
<td>46</td>
<td>46</td>
</tr>
<tr>
<td></td>
<td>α</td>
<td>1.20</td>
<td>1.23</td>
<td>1.25</td>
<td>1.28</td>
<td>1.28</td>
</tr>
</tbody>
</table>

http://www.revistadechimie.ro REV.CHIM.(Bucharest) • 70 • No. 1 • 2019
In the potential region (a), the Tafel slope values vary between 99 and 137 mV/decade, which correspond to the theoretical value \((\frac{\partial E}{\partial \ln i}) = \frac{2RT}{nF}\) for the one-electron charge transfer. The charge transfer coefficient \(a\) is close to 0.5 that presumes the absence of the adsorption phenomenon in the reaction kinetics. In the second charge transfer controlled step, the potential region (b), the Tafel slopes have values of 49-46 mV/decade, which corresponds to the catalytic reactions.

The figures 4a and 4b show the relationships between the measured values of \(\log i\) and \(\log \text{ClO}_3^-\) at various constant potentials situated in the two Tafel domains. The reaction order vs. the \(\text{ClO}_3^-\) ion \((p_{\text{ClO}_3^-})\) was calculated according to the relation:

\[
\left(\frac{\partial \log i}{\partial \log \text{ClO}_3^-}\right) = p_{\text{ClO}_3^-} \tag{4}
\]

The reaction order \((p<1)\) indicates the formation of free radicals in the two charge transfer controlled steps of the chlorate reduction.

Figures 4a and 4b illustrate the current dependence of chlorate concentration in the case of the various potentials situated in the region of the limiting current. In the figure 5a the current density dependence tends to exponential growth with chlorate concentration (at concentrations ≥ 01 M \(\text{KClO}_3\)) and becomes exponential in the second part of the limiting current (fig. 5b). This behavior presumes an autocatalytic effect of the chlorate concentration on its reduction.

The redox system \(\text{Cl}_2/\text{Cl}^-\) is a complex one. In the reduction process six electrons are transferred, which presumes several electrochemical steps associated with chemical reactions. In acid media, chlorate may be electrochemically reduced to chlorine or chloride, in accordance with the total reactions [44,45]:

\[
E_0 = +1.5 \text{ V} \tag{5}
\]

\[
E_0 = +1.45 \text{ V} \tag{6}
\]

The chlorate ion is a strong oxidant and can oxidize \(\text{Cl}^-\) to \(\text{Cl}_2\) or \(\text{Cl}_3\), according to the chemical reaction [45]:

\[
\text{ClO}_3^- + 6\text{H}^+ + 5\text{Cl}^- = 3\text{Cl}_2 + 3\text{H}_2\text{O} \tag{7}
\]

Also, in solution the following equilibrium appears:

\[
\text{Cl}_3^+ (aq) + \text{Cl}^- = \text{Cl}_2 + 
\]

In accordance with the potential \(p\text{H}\) equilibrium diagram [46] for the system chlorine-water, at a temperature of 298 K in acid medium \((p\text{H}<3.5)\) the chlorate reduction has as reaction product chloride and as intermediate - chlorine gas or chlorine \((aq)\). Conform to the activity-\(p\text{H}\) diagram [46] for chlorine-water system \((at 298 \text{ K, } p_{\text{Cl}_2}=1 \text{ atm} \text{ and } p\text{H } \leq 3)\), the activities of the species \(\text{Cl}_2(aq)\) and \(\text{Cl}_3\) are relevant.

Chlorate reduction can generate free radicals, such as: \(\bullet \text{ClO}_2\), \(\bullet \text{ClO}_3\), \(\text{ClO}\) or \(\bullet \text{Cl}\) [43].

The experimental results showed an extended domain potential for chlorate reduction with various electrochemical behaviors that indicates different mechanisms of the intermediate reactions. According to them, the first charge transfer controlled step implicates the formation of free radicals (the reaction order is subunit conform to figure 4a). The Tafel slopes values (table 1) correspond to some partial reactions with one-electron transfer. These results could be satisfied by the following mechanism:

\[
\text{ClO}_3^- + 2\text{H}^+ + e^- \rightarrow \text{ClO}_2 + \text{H}_2\text{O} \tag{11}
\]

\[
\bullet \text{ClO}_2 + \text{H}^+ + e^- \rightarrow \text{HClO}_2 \tag{12}
\]

\[
\text{HClO}_2 + \text{H}^+ + e^- \rightarrow \text{OCl} + \text{H}_2\text{O} \tag{13}
\]

\[
\text{OCl} + \text{H}^+ + e^- \rightarrow \text{HOCl} \tag{14}
\]

\[
\text{HOCl} + \text{H}^+ + e^- \rightarrow \text{Cl}^- + \text{H}_2\text{O} \tag{15}
\]

\[
\text{Cl}^- + e^- \rightarrow \text{Cl}^- \tag{16}
\]

\[
\text{ClO}_3^- + 6\text{H}^+ + 5\text{Cl}^- = 3\text{Cl}_2 + 3\text{H}_2\text{O} \tag{7}
\]

\[
\text{ClO}_3^- + 2\text{H}^+ + e^- \rightarrow \text{ClO}_2 + \text{H}_2\text{O} \tag{11}
\]

\[
\bullet \text{ClO}_2 + \text{H}^+ + e^- \rightarrow \text{HClO}_2 \tag{12}
\]

\[
\text{HClO}_2 + \text{H}^+ + e^- \rightarrow \text{OCl} + \text{H}_2\text{O} \tag{13}
\]

\[
\text{OCl} + \text{H}^+ + e^- \rightarrow \text{HOCl} \tag{14}
\]

\[
\text{HOCl} + \text{H}^+ + e^- \rightarrow \text{Cl}^- + \text{H}_2\text{O} \tag{15}
\]

\[
\text{Cl}^- + e^- \rightarrow \text{Cl}^- \tag{16}
\]
The reactions (11) - (16) imply successive reduction reactions with one-electron transfer and formation of intermediate free radicals. A similar mechanism was indicated by other authors [37] with the difference that the electrons were provided by the oxidation of the ferrous ions to ferric ions. Summing the reactions (11) - (16) the overall reaction is obtained (6). Chlorate reduction in the first potential region has the chloride ion as final product.

In accordance with figure 5a the current density increases slowly with chlorate concentration at small concentrations and tends to the exponential growth at higher concentrations. That presumes an autocatalytic effect of chlorate ions. It must be admitted that at higher concentrations, the chlorate ion interacts with active free radicals (•ClO) conform to reaction:

$$\cdot ClO + ClO_3^- + 2H^+ + e^- \rightarrow 2HClO_2$$ \hspace{1cm} (17)

setting up a catalytic loop similar with the free radical BrO₂ for bromate reduction [48].

The exponential growth of the limiting current density with chlorate concentration in the potential region - (100-200) mV/SCE (fig. 5b) presumes other reactions with catalytic effect. Together with the potential shift to negative direction, the chloride ion concentration increases in the presence of the catalyst system. In the second charge transfer controlled step the chlorate reduction is catalyzed by a surface reaction between Pt electrode and free radicals. \(Cl_2^-\). The final product of chlorate reduction is chloride ion in all potential regions.

**Conclusions**

The electrochemical reduction of the chlorate at Pt electrode in acid medium is a complex process that implies two charge transfer controlled steps with the formation of free radicals and an extended potential region controlled by the concentration polarization. The current density dependence of chlorate concentration tends to an exponential growth at concentration ≥ 0.1 M KClO₃, and in the presence of the catalyst system. In the second charge transfer controlled step the chlorate reduction is catalyzed by a surface reaction between Pt electrode and free radicals. •ClO. The final product of chlorate reduction is chloride ion in all potential regions.

**References**


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