Voltammetric study of formic acid oxidation via active chlorine on IrO₂/Ti and RuO₂/Ti electrodes

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Abstract: This work aimed to contribute to the mechanism electrochemical oxidation study of organic compounds on DSA electrodes. To do this, IrO₂ and RuO₂ electrodes were prepared thermally at 40°C on Titanium substrate. The prepared electrodes were characterized using voltammetric and SEM techniques. The electrochemical measurements in acid media made it possible to show the presence of IrO₂ and RuO₂ on the surface of the electrode. These electrodes have identical electrocatalytic behaviors both for oxygen evolution and chlorine evolution. The prepared electrodes make it possible to oxidize the organic compounds in an acid media in the absence or in the presence of Cl⁻. In acidic electrolytes, water molecules produce hydroxyl radicals that contribute to the higher oxides (RuO₃ or IrO₃) formation. The higher oxides obtained produce O₂ and regenerate the active sites of our electrodes. In the electrolytes containing chlorides, higher oxides fix them (IrO₃(Cl) or RuO₂(Cl)) in competition with the production of O₂. Then IrO₃(Cl) or RuO₂(Cl) reacts with Cl⁻ to produce Cl₂ and regenerate the adsorbed hydroxyl radicals. The higher oxides also react as a mediator in HCOOH oxidation in competition with O₂ evolution. In the electrolytes containing HCOOH and Cl⁻, the organic pollutant's oxidation occurs indirectly via the hypochlorite ions produced in the solution and on the electrodes. This study showed that the produced OH⁻ and Cl₂ in situ are involved in the oxidation of HCOOH.

Keywords: electrode; metallic oxides; oxygen; chlorine; formic acid.

1. Introduction

The application of electrochemical technologies to environmental pollution abatement has been the topic of several books 1–6. The main advantage of these technologies is the ability to mineralize harmful pollutants into simple inorganic compounds such as H₂O or CO₂, which are natural for the environment. Generally, the organic pollutants mineralization occurs on the anode surface via the production of hydroxyl radicals 6–8 or/and other oxidants. Thus, anode material plays a crucial part in the electrochemical technologies 9–12.

In recent years, with the introduction of dimensionally stable anodes (DSA) by H. Beer 13, metal oxide electrodes (such as RuO₂ and IrO₂) have become increasingly technologically important. Major characteristics of these anodes are longer operating lifetime 14 and comparatively low cost 15,16. Thus, the DSA anodes constitute an important class of stable electrocatalysts 17 mainly used in the chloro-alkali industry 18, in the production of oxygen and hydrogen in water electrolysis, and have received attention wastewater treatment 19. The main drawback of using DSA is the generation of highly reactive oxidants limited (like hydroxyl radicals), reducing their wastewater treatment efficiencies. Therefore, it is necessary to overcome that limit.

According to the literature 20,21, the generation of highly reactive oxidants might be overcome if DSA is irradiated with UV light. Besides, these anodes have the facility to generate chloro oxidant species (Cl₂, HOCl, and OCI) when electrolysis is carried out at chloride-containing medium 15. In several studies, NaCl has been employed as the salt to increase electrolyte 1,3,6. However, few works have reported organic compounds' mechanism degradation via chloro oxidant species on DSA anodes.

Different voltammetry techniques can be used to study an electrochemical system’s response to polarization: linear, cyclic, and square wave voltammetry. Voltammetry is one of the most important electroanalysis techniques 22,23, also used...
for the reaction mechanisms determination. Its strength lies in the simplicity of implementation and the wealth of information collected for the redox processes study.

For this reason, this work aims to study the oxidation mechanism of synthetic effluent in the presence of different Cl− concentrations, using DSA anodes. Owing to their high electrocatalytic properties, we used RuO2 and IrO2 as electrodes to study the oxidation of the formic acid in the presence of oxygen (O2) and chlorine (Cl2) evolution by voltammetric measurements. Formic acid was chosen because it is the smallest molecule with the four most common chemical bonds in organic compounds (C−H, C=O, C−O, O−H), making formic acid an ideal model molecule for studying electrooxidation reactions.

2. Experimental

2.1. Preparation of RuO2 (ruthenium oxide) and IrO2 (Iridium oxide) anodes
RuO2 and IrO2 electrodes were prepared by the thermal decomposition method on titanium substrates. The precursor solution is made by the dissolution of RuCl3·xH2O (Fluka, 99.98 %) and H2IrCl6·6H2O (Fluka, 99 %) in pure isopropanol (Fluka, 99.5 %). Before the deposition and to ensure maximum adhesion of the coating, the substrate (Titanium) was sandblasted. The precursor was applied by a painting procedure on the titanium (Ti) substrate then put in an oven for 10 minutes at 80°C to allow the solvent evaporation. It is then placed in a furnace at 400°C for 15 minutes to allow the precursor's decomposition. These steps were repeated until the desired mass of the coating is reached. A final decomposition of 1 h was done at 400°C. The deposits' loading is around (5, 0 ± 0, 3) g/m².

2.2. Electrochemical measurements

The electrochemical measurements were performed in a three-electrode cell. The working electrode was IrO2 or RuO2 electrode, the counter electrode was a platinum wire, and the reference electrode was a saturated calomel electrode (SCE). The electrochemical study (potentiodynamic polarization) is carried out using an experimental device composed of an Autolab Potentiostat de ECHOCHEMIE (PGSTAT 20), interfaced with a computer. To overcome the potential ohmic drop, the reference electrode was mounted in a luggin capillary and placed close to the working electrode by a distance of 1 mm. The apparent exposed area of the working electrode was 1 cm².

The chemicals used for the electrochemical study are HClO4 (Suprapur Merck, 70 %), NaCl (Fluka, 99 %), and HCOOH (Fluka, 98 %). All solutions were prepared with distilled water, and the experiments were carried out at room temperature at 25°C.

To decrease the potential ohmic drop's contribution, the reference electrode was mounted in a luggin capillary and placed close to the working electrode by a distance of 1 mm. The ohmic drop is known to increase the Tafel slope value, and careful attention was paid to it during the experiment. To reduce the residual ohmic drop, the polarization curve should be corrected according to the method described elsewhere. A brief description of the correction is given as followed.

The overpotential η / V observed in the experiment can be given by

\[ \eta / V = a + b \ln j + j R \]  

(1)

where \( a / V \) is the Tafel constant, \( b / V \) dec⁻¹ is the Tafel slope, \( j / A \) cm⁻² is the current density and \( R / \Omega \) cm⁻² is the total area-specific uncompensated resistance of the system, which is assumed to be constant.

The derivative of equation (1) for current density gives equation (2) from which b and R can be easily obtained by plotting \( d\eta / dj \) as a function of \( 1/j \)

\[ \frac{d\eta}{dj} = \frac{b}{j} + R \]  

(2)

The estimation of R allows correcting the experimental overpotential by subtracting the ohmic drop jR according to equation (3).

\[ \eta_{corr} = \eta - j R \]  

(3)

where \( \eta_{corr} \) stands for the corrected overpotential.

During the calculations, the derivatives \( d\eta / dj \) was replaced by their finite elements \( \Delta\eta / \Delta j \) estimated from each pair of consecutive experimental points.

3. Results and Discussion

3.1. Physical characterization of the electrodes

In Figure 1, the micrographs of the prepared electrodes were presented. It appears that the deposit has covered the surface. The surface of the IrO2/Ti and RuO2/Ti electrodes is rough with a mud cracked structure. That observation is similar to that is generally observed for thermally prepared RuO2/Ti and IrO2/Ti electrode. Pores are also present on the deposit. The cracks observed on the electrodes surfaces occur during the cooling of the deposit because of the thermal shock to which these deposits are subjected during their retreat from the furnace.
3.2. Electrochemical characterization of the electrodes in acid media and acid media containing chloride

In Figure 2a, the voltammogram of the IrO$_2$ electrode recorded in 1 M HClO$_4$ at 100 mV/s was presented. A high voltammetric charge is observed with localized peaks at 0.7 and 1 V vs. SCE. These reversible broad peak characteristics of the redox transition of Ir(III)/Ir(IV) and Ir(IV)/Ir(VI) was calculated from the measurements carried out in 1 M HClO$_4$ media for different values of the scan rate presented in Figure 2b. This report shows that the cathode charge is greater than the anode charge at low potential scan rates. This result would indicate that at low potential scan rates, probably irreversible (slow) processes at the electrode have not been fully recovered during anodic scanning. On the other hand, for the high scan rate, whatever the potential scan rate, the cathodic charge is practically equal to the anode charge. Therefore, the curves are symmetrical throughout the potential domain regardless of the high scan rates' potential scan rate. This result shows that the observed anodic and cathodic peaks are related and can be attributed to the formation and reduction of the higher oxide (IrO$_3$).

The ratio of voltammetric charges $|q(+) / q(-)|$ was calculated from the measurements carried out in 1 M HClO$_4$ media for different values of the scan rate. This report shows that the cathode charge is greater than the anode charge at low potential scan rates. This result would indicate that at low potential scan rates, probably irreversible (slow) processes at the electrode have not been fully recovered during anodic scanning. On the other hand, for the high scan rate, whatever the potential scan rate, the cathodic charge is practically equal to the anode charge. Therefore, the curves are symmetrical throughout the potential domain regardless of the high scan rates' potential scan rate. This result shows that the observed anodic and cathodic peaks are related and can be attributed to the formation and reduction of the higher oxide (IrO$_3$).

Voltammetric measurements were performed in perchloric acid containing various concentrations of NaCl on IrO$_2$ electrode. Figure 3 shows the different linear voltammograms obtained. This figure shows a
rapid increase in current intensity between 1.13 and 1.35 V vs. SCE, followed by another rapid growth in current from 1.35 V vs. SCE. The first increase in intensity would correspond to chloride oxidation to chlorine because the current intensity increases with chloride concentration. The chlorine evolution reaction starts at a lower potential than the oxygen evolution reaction obtained in the presence of HClO₄. This result shows the chlorine evolution reaction before the oxygen evolution reaction on this electrode.

Figure 3. Linear voltammograms recorded on IrO₂/Ti electrode in 1 M HClO₄ containing NaCl at 100 mV/s

In Figure 4a, the voltammogram of the ruthenium dioxide electrode recorded in 1 M HClO₄ at 100 mV/s was presented. A rectangular shape voltammogram is observed with waves at 0.6 and 0.9 V vs. SCE. The presence of these waves could be due respectively to the redox transitions of Ru (III) / Ru (IV) and Ru (IV) / Ru (VI) on RuO₂ electrode surface. This voltammogram is identical to that obtained in our previous work with the RuO₂ electrode. This result confirms the presence of RuO₂ on our electrode. On this electrode, the oxygen evolution reaction starts at 1.2 V vs. SCE.

Figure 4b shows the RuO₂ voltammograms in 1 M HClO₄ containing chloride. A rapid increase of the current intensity is observed after 1.15 V vs. SCE for the concentrations investigated. This potential is lower than that of oxygen evolution. Then this rapid increase of the current would be related to the Cl⁻ oxidation in Cl₂. Figure 5 shows the characteristics of RuO₂ electrode in the presence of Cl⁻ only with the correction of the baseline linked to the supporting electrolyte. Oxidation peaks are observed. These peaks are Cl⁻ to Cl₂ oxidation peaks because they increase with the Cl⁻ concentrations. This result confirms that the rapid increase in current intensity observed from 1.15 V vs. SCE is related to the Cl₂ evolution reaction.

Figure 4. (a) Cyclic voltammogram recorded on RuO₂/Ti electrode in 1 M HClO₄ at 100 mV/s; (b) linear voltammograms recorded on RuO₂/Ti electrode in 1 M HClO₄ containing NaCl at 100 mV/s
From the obtained voltammograms, the Tafel slopes for the oxygen evolution reaction were determined. 39 and 40 mV/dec were obtained respectively for IrO$_2$ and RuO$_2$. This result shows that the prepared electrodes have the same oxygen evolution reaction kinetics in an acidic electrolyte. Taking into account these values of the Tafel slopes, obtained voltammograms and our previous work $^{24,26}$ and the literature $^{32,49}$ the following mechanism is proposed for the oxygen evolution reaction in acid media on our electrodes:

$$\beta O_2 + H_2O \rightarrow \beta O_2(HO) + H^+ + e^- \quad (4)$$
$$\beta O_2(HO) \rightarrow \beta O_3 + H^+ + e^- \quad (5)$$
$$2 \beta O_3 \rightarrow 2 \beta O_2 + O_2 \quad (6)$$

Where \(\beta\) is Ir or Ru

The mechanism described by eqs. (4-6) predicts the Tafel slope of 120 mV/dec if the first step is represented by eq. (4) the rate-determining step (rds), 40 mV/dec if the second step is described by eq. (5) rds and 30 mV/dec if the third step is represented by eq. (6) is rds. For OER on IrO$_2$ or RuO$_2$ in HClO$_4$ medium, the rate-determining step is the step described by eq. (5), because the Tafel slope is 40 mV/dec.

The Tafel slopes were also determined for the chlorine evolution reaction in acidic media. 40 mV/dec was obtained with IrO$_2$ and RuO$_2$ electrodes. This result shows that these electrodes have the same kinetic of the chlorine evolution reaction in acidic media. According to our previous work $^{25,28}$, and the Tafel slopes value, a global chlorine evolution reaction mechanism is described as followed in acidic media. After the formation of higher oxides (\(\beta O_3\)) in the eq.5, chloride ions attach themselves to the higher degree oxide (eq. 7), then chlorine is produced (eq. 8).

$$\beta O_3 + Cl^- \rightarrow \beta O_3(Cl^-) + e^- \quad (7)$$
$$\beta O_3(Cl^-) + H^+ + Cl^- \rightarrow \beta O_2(HO^-) + Cl_2 \quad (8)$$

Where \(\beta\) is Ir or Ru

This mechanism predicts the following Tafel slopes: 120 mV/dec if the first step (eq. 5) is the rate-determining step (rds), 40 mV/dec for the second step (eq. 7), and 30 mV/dec for the third step (eq. 8). For chlorine evolution reaction on IrO$_2$ or RuO$_2$ in HClO$_4$ medium, the rate-determining step is described by eq. (7), because the Tafel slope is 40 mV/dec.

### 3.3. Formic acid oxidation without chloride

Figure 6a shows the voltammograms recorded on IrO$_2$ in formic acid media. No peak characteristic of the formic acid oxidation for the potential domain between -0.25 and 1 V vs. SCE is observed when an organic compound is added to these voltammograms. This result indicates that IrO$_2$ electrode is not electrocatalytic for HCOOH oxidation on this domain. After 1 V vs. SCE, a modification is observed on the voltammograms. The current intensity increases as the organic compound concentration increase for a given potential. The oxygen evolution potential decreases with the rise of the formic acid concentration. In the absence of formic acid, the oxygen evolution starts at 1.22 V vs. SCE. In the presence of formic acid, the oxygen evolution starts at 1.1 V vs. SCE for a formic acid concentration of 0.5 M and 1.04 V vs. SCE for 1 M.

In order to determine the behavior of IrO$_2$ in a formic acid media with correction of the baseline linked to the support electrolyte, the curves of Figure 6b were carried out. To get it, the current intensities obtained in the absence of formic acid were subtracted from the current intensities obtained in the presence of formic acid. This subtraction makes it possible to get the current linked to the organic compound oxidation. From -0.25 to 1 V vs. SCE, no oxidation peak of formic acid is observed. There is a rapid increase in current after 1 V vs. SCE. We observe a shift of the curve to the left when the concentration of formic acid increases. The current intensity increases with the rise.
of formic acid concentration. The rapid increase after 1 V vs. SCE would be related to the formic acid oxidation in which the species from the electrochemical decomposition of water are likely to intervene.

HCOOH oxidation on ruthenium oxide is shown in Figure 7a. This figure shows that the voltammetric curves change recorded in the absence and then in formic acid are observed at potentials higher than 1 V vs. SCE. From -0.25 to 1 V vs. SCE, these voltammograms are almost superimposable. No peak characteristic of formic acid oxidation is observed in this potential domain. On the other hand, there is a shift in the oxygen evolution reaction potential to the left as the amount of formic acid increases. For 0; 0.5 and 1 mM of formic acid, the oxygen evolution reaction start respectively at 1.17; 1.05 and 1.01 V vs. SCE. The current intensity increases with the increase of formic acid concentration. For a potential of 1.17 V vs. SCE, the current intensity is 0.24 mA in the absence of formic acid. In the presence of 0.5 and 1 mM formic acid, the current intensities are 0.99 mA and 1.68 mA, respectively.

Indeed, it follows from the previous observations that the organic compound oxidation occurs after 1 V vs. SCE in a domain where the voltammetric measurements had revealed the redox transitions Ir (IV) / Ir (VI) IrO2 / IrO3 with IrO2 electrode. With RuO2, the Ru (IV) / Ru (VI) redox transitions are observed in the same potential domain. These observations made it possible to propose the following mechanism for the organic compound oxidation.
oxidation and the simultaneous oxygen evolution reaction in an acid media without chloride.

During the formic acid oxidation, the first reaction is the electrochemical decomposition of water molecules, leading to hydroxyl radicals $^{46}$. These hydroxyl radicals resulting from the electrochemical decomposition of water are instantaneously adsorbed on the electrode surface. Since iridium oxide and ruthenium oxide are active electrodes, there is a strong interaction between the electrode and the hydroxyl radicals. In this case, the adsorbed hydroxyl radicals can interact with the anode followed by a redox transition at the electrode surface, forming a higher oxide ($\text{IrO}_3$ or $\text{RuO}_3$). $\text{IrO}_3$ or $\text{RuO}_3$ can act as a mediator in the organic compounds (eq. 9). This organic compound oxidation reaction competes with the oxygen evolution reaction eq. 6.

$$\beta\text{O}_3 + \text{HCOOH} \rightarrow \beta\text{O}_2 + \text{CO}_2 + \text{H}_2\text{O} \quad (9)$$

### 3.4. Formic acid oxidation in acid media containing chloride

The formic acid oxidation in an acid media containing chloride on the $\text{IrO}_2$ electrode is shown in Figure 8. This figure's voltammograms show that no significant change is observed at -0.25 to 1 V vs. SCE when the organic compound is added. By exploring high potential domains from 1.1 to 1.4 V vs. SCE, it is noted that the current increases when $\text{HCOOH}$ is added to the electrolytic solution. The anode wave observed at 1.27 V vs. SCE in the absence of $\text{HCOOH}$ is almost invisible in the presence of $\text{HCOOH}$. In the presence of formic acid, the current increases rapidly after the wave at 1.27 V vs. SCE. Chloride would therefore intervene indirectly in the $\text{HCOOH}$ oxidation.

In the presence of chloride, Figure 9 shows the voltammograms recorded on ruthenium oxide. The voltammograms show an oxidation peak at 1.29 V vs. SCE. These peaks could be due to the chloride oxidation in the electrolyte. The intensity of the oxidation peak increases with the increase of formic acid concentration while the reduction peak remains constant. The increase in the oxidation intensity wave would be characteristic of the organic compound oxidation. This oxidation probably occurs consecutively to that of the chloride $^{47}$. Then in the presence of Cl-, the HCOOH oxidation on these electrodes would proceed according to the following mechanism:

$$\text{Cl}_2 + \text{H}_2\text{O} \rightarrow \text{HOCl} + \text{H}^+ + \text{Cl}^- \quad (10)$$

$$\text{HOCl} \rightarrow \text{ClO}^- + \text{H}^+ \quad (11)$$

$$\text{HCOOH} + \text{ClO}^- \rightarrow \text{CO}_2 + \text{H}_2\text{O} + \text{Cl}^- \quad (12)$$

![Figure 8](image1.png) **Figure 8.** (a) Formic acid oxidation in acid media containing 0.1 M chloride on IrO$_2$/Ti electrode; (b) IrO$_2$/Ti electrode in a formic acid with the baseline linked to the electrolyte support at 20 mV/s

![Figure 9](image2.png) **Figure 9.** (a) Formic acid oxidation in acid media containing 0.1 M chloride on RuO$_2$/Ti electrode; (b) RuO$_2$/Ti electrode in a formic acid with the baseline linked to the electrolyte support at 20 mV/s
There is, therefore, a first step, which consists of chloride oxidation. This leads to the chlorine evolvement on the different electrodes. The formed chlorine is then hydrolyzed to hypochlorous acid (HOCI), which is a powerful oxidant. In acidic media, hypochlorous acid releases its proton (H\(^+\)) and forms the hypochlorite ion (ClO\(^-\)). It is these ions (ClO\(^-\)) that oxidize the organic compound.

By relying on the mechanisms of oxygen evolution reaction, chlorine evolution reaction, and formic acid oxidation above, Figure 10 makes it possible to propose a mechanism for the formic acid oxidation in acid media contains chloride. Concerning the oxygen evolution in acid media, we have steps 1, 2, and 3. In acid media containing Cl\(^-\), Cl\(^-\) is fixed on the higher oxide, then the chlorine occurs, and the adsorbed radical is regenerated (steps 4 and 5). In the presence of HCOOH, the higher oxide (IrO\(_2\) or RuO\(_2\)) allows the organic compound oxidation to CO\(_2\) and H\(_2\)O (step 6). This organic compound oxidation competes with the oxygen evolution reaction (steps 3 and 6).

In acid media containing chloride, the oxidation mechanism of organic compounds is complex. The chlorine produced indirectly intervenes in the organic compound oxidation according to steps 7, 8, and 9 of the mechanism. The chlorine makes it possible to oxidize HCOOH and regenerates Cl\(^-\). In the presence of chloride, the HCOOH oxidation on the IrO\(_2\) and RuO\(_2\) electrodes takes place at high potentials. At high potentials, the Cl\(^-\) oxidation leads to the production of Cl\(_2\) or ClO\(^-\). Thus, the formic acid oxidation occurs indirectly via hypochlorite ions and directly on the electrodes' active sites in competition with the oxygen evolution reaction.

4. Conclusion
The physical and electrochemical characterizations performed on these electrodes revealed the presence of IrO\(_2\) or RuO\(_2\). This work has shown that RuO\(_2\) and IrO\(_2\) have the same kinetics of oxygen evolution and chlorine evolution. The voltammograms obtained made it possible to show that the prepared metal oxide electrodes could oxidize the organic compounds. However, the oxidation current of the organic compound increases in the presence of chloride. In HCOOH, the higher oxide (IrO\(_2\) or RuO\(_2\)) acts as a mediator in the oxidation of the organic compounds in competition with the oxygen evolution reaction. In acid media containing chloride, the oxidation mechanism of organic compounds is complex. At high potentials, the oxidation of Cl\(^-\) leads to the production of Cl\(_2\). This reaction competes with the oxygen evolution reaction and the reaction's formic acid oxidation reaction via hydroxyl radicals. The chlorine produced is transformed into a hypochlorite ion, which is involved in the formic acid oxidation.

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