Adsorption of ochratoxin A (OTA) anodic oxidation product on glassy carbon electrodes in highly acidic reaction media: Its thermodynamic and kinetics characterization

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Abstract

We study the thermodynamics and kinetics of the adsorption of a redox couple having quinone nature on glassy carbon electrodes. This couple is produced by the anodic oxidation of mycotoxin ochratoxin A in 10% acetonitrile + 90% 1 M HClO4 aqueous solution. The quasi-reversible redox couple was studied by both cyclic (CV) and square wave (SWV) voltammetric techniques. The Frumkin adsorption isotherm best described the specific interaction of the redox couple with carbon electrodes. By fitting the experimental data, we obtained values of $-28.4 \text{ kJ mol}^{-1}$ and $0.70 \pm 0.02$ for the Gibbs free energy of adsorption and the interaction parameter, respectively. SWV fully characterized the thermodynamics and kinetics of the adsorbed redox couple, using a combination of the “quasi-reversible maximum” and the “splitting of SW peaks” methods. Average values of $0.609 \pm 0.003 \text{ V}$ and $0.45 \pm 0.06$ were obtained for the formal potential and the anodic transfer coefficient, respectively. Moreover, a formal rate constant of $10.7 \text{ s}^{-1}$ was obtained. SWV was also employed to generate calibration curves. The lowest concentration of mycotoxin was $1.24 \times 10^{-8} \text{ M}$ (5 ppb), measured indirectly with a signal to noise ratio of 3:1.

1. Introduction

Ochratoxin A (OTA), or 7-carboxyl-5-chloro-8-hydroxyl-3,4-dihydro-3R-methylisocoumarin-7R-l-phenylalanine (Fig. 1a), is a colorless crystalline compound that belongs to a group of closely related derivatives of isocoumarin linked to l-phenylalanine and classified as pentaketides [1]. OTA is a secondary metabolite produced by fungi of the Penicillium (e.g. P. verrucosum) and Aspergillus (e.g. A. ochraceus) genera. It is a known teratogen and immunosuppressive agent and has been implicated in Balkan nephropathy in humans. The International Agency for Research on Cancer (IARC) lists OTA as possibly carcinogenic to humans (Group 2B) [2]. Humans can consume OTA in vegetal and animal products [3–5]. For this reason, many countries have restricted OTA levels in food, with upper limits of 1 and 10 ppb depending on the type and quality of the foodstuff [3].

With imports and exports of agricultural commodities increasing in recent years, monitoring of OTA is essential for food safety in the world-wide market. Many research groups have been studying methods for monitoring naturally occurring OTA in a number of agricultural commodities [3,6]. Like many other mycotoxins, OTA is primarily detected and quantified using liquid chromatography in conjunction with fluorimetric detectors and by LC–MS techniques [7–10]. Thin-layer chromatography and enzyme-linked immunosorbent assay (ELISA) are alternative methods. Today, the electroanalytical methods are very valuable alternatives to other methods because no separation procedure is required prior to substrate determination. Furthermore, the required instrumentation is typically less expensive, smaller amounts of solvent are necessary and analysis time is shorter for electroanalytical methods over chromatographic measurements.

To secure these advantages, competitive ELISA methods have been proposed for the quantification of OTA. One such direct method uses polyclonal antibodies [11]. Another uses screen-printed electrodes for quantitative determination of OTA [12], with detection limits in the range 0.9 and 100 µg L$^{-1}$ for the direct and indirect assay format, respectively. Two ELISA strategies were recently investigated for the development of OTA electrochemical immunosensors based on different OTA immobilization procedures. These immunosensors were suitable as screening tools to detect OTA in wine [13]. In addition, a novel electrochemical immunosensor for sensitive detection of OTA based on screen-printed gold electrodes modified with a layer of 4-nitrophenyl diazonium salt was recently reported. A detection limit
of 12 ng ml$^{-1}$ was achieved [14]. Another label-free electrochemical immunosensor developed on modified gold electrodes for sensitive detection of OTA was also recently published [15]. However, fundamental studies on the electrochemical behavior of OTA are necessary to assess the feasibility of employing these electroanalytical methods.

To our knowledge, a few papers presently exist addressing the electrochemical behavior of OTA [16,17]. The electrochemical oxidation of OTA in acetonitrile (ACN) has been reported, as well as that in phosphate aqueous buffer solutions in the pH range from 6 to 8 [16]. A direct correlation with the oxidative behavior of 4-chlorophenol was established. The electrochemical oxidation of OTA in aqueous buffer solutions of different chemical composition in the pH range from 2 to 12 was also recently reported [17]. In this study, we report an adsorbed product of OTA anodic oxidation having quinone nature, which we observed in ACN and in acidic aqueous solutions (OTQ) [16]. Its anodic oxidation voltammetric response depended on pH, but no in-depth investigation of the adsorption characteristics of OTQ was carried out.

In this article, we report the results of our studies on the thermodynamics and kinetics of the adsorption of OTQ on glassy carbon (GC) electrodes in 10% acetonitrile (ACN) + 90% 1 M HClO$_4$ aqueous solution. A surface-based quasi-reversible redox couple was obtained from the adsorption of OTQ, which has been ascribed to an OTA type quinone (OTQ)/hydroquinone (OTHQ) derivative (Fig. 1b and c) [16]. This was characterized by cyclic (CV) and square wave (SWV) voltammetric techniques. Surface charge transfer kinetics and thermodynamic data for the OTQ/OTHQ redox couple were computed by identifying the quasi-reversible maximum and by splitting the square wave voltammetric peaks [18–24]. These results have very promising analytical applications for indirect determination of OTA at trace levels.

2. Experimental

2.1. Reagents

OTA was purchased from SIGMA and used as received. ACN and H$_2$O were HPLC grade from Sintorgan. HClO$_4$ (Merck p.a.) was used without further purification. Stock solutions of OTA were prepared in ACN and kept at 4 °C in the dark. Working solutions were prepared daily by adding aliquots of stock solution to 10% ACN + 90% aqueous 1 M HClO$_4$. The solution used for OTA spontaneous adsorption was 1.2 × 10$^{-4}$ M OTA in 10% ACN + 90% aqueous 1 M HClO$_4$. For safety reasons, plastic gloves were used in all manipulations. Most of the experiments were carried out in blank solutions composed of aqueous 1 M HClO$_4$. The influence of pH on the surface redox couple was studied in buffer solutions prepared by combining 0.2 M Na$_2$HPO$_4$ (Merck p.a.) with 0.2 M KH$_2$PO$_4$ (Merck p.a.). The pH of the solutions ranged from 1 to 12. Solutions were prepared from the above buffer solution and their pH values were adjusted by adding different volumes of 1 M HClO$_4$ or 1 M NaOH.

2.2. Apparatus and experimental measurements

CV and SWV experiments were performed with an AutoLab PGSTAT 12 potentiostat, controlled by GPE 4.9 electrochemical software from Eco-Chemie, Utrecht, The Netherlands. In CV measurements, the scan rate ($\nu$) was varied from 0.005 to 1.000 V s$^{-1}$. For SWV, a square wave amplitude of $\Delta E_{SW} = 0.025$ V and a staircase step height of $\Delta E_s = 0.005$ V were predominantly used. The frequency ($f$) was varied from 9 to 40 Hz. In some experiments, $\Delta E_{SW}$ was varied from 0.025 to 0.125 V.

Electrochemical measurements were performed in a two-compartment Pyrex cell [25]. The working electrode was a GC disk (BAS, 3 mm diameter). It was polished with 0.3 and then 0.05 μm wet alumina powder (from Fischer), copiously rinsed with water and sonicated in a water bath for 2 min. Then, it was electrochemically activated in aqueous 1 M KOH (Merck p.a.) by applying a potential of 1.2 V over 5 min, according to a methodology described by Anjo et al. [26]. Finally, it was transferred to a blank solution (aqueous 1 M HClO$_4$) and cycled 10 times between 0 and 1.4 V. Anodic and cathodic peak currents ($I_{p,a}$ and $I_{p,c}$), peak potentials ($E_{p,a}$ and $E_{p,c}$) and the difference between anodic and cathodic peak potentials ($\Delta E_p$), were highly reproducible when this pre-treatment was applied to the working electrode. The counter electrode was a large-area platinum foil (approximately 2 cm$^2$). The reference electrode was an aqueous saturated calomel electrode (SCE) fitted with a fine glass Luggin capillary containing a bridge solution identical to that containing the sample being measured. While no difference was observed in voltammetric responses in aerated vs. deaerated solutions, to ensure reproducible measurement conditions samples were deaerated by bubbling purified nitrogen through the solutions for at least 10 min prior to measurement. The temperature was 20 ± 0.2 °C.

Experimental data were fitted using a nonlinear least squares procedure in Origin 7.0 to determine the adsorption isotherms which best described the specific interaction of OTQ with GC electrodes. The Chi-square function ($\chi^2$) was employed to choose the best fit between experimental and theoretical data.

3. Results and discussion

3.1. Cyclic voltammetry

Cyclic voltammograms of OTA recorded at different pH on activated GC electrodes resembled those in the literature [16,17], and the OTA oxidation peaks ($E_p \approx 1.1–1.2$ V; pH 1) were shifted to less positive potentials with pH ($E_p \approx 0.8$; pH 8–12). These results agree qualitatively with those of Oliveira et al. [17] who worked within a similar pH range (2–12). On reversing the potential sweep no complementary cathodic peak was observed, but another surface-based peak system appeared at less positive potentials for pH < 6. This has been assigned to the reduction/oxidation of an OTA anodic oxi-
The surface-based nature of the system was confirmed by responding to oxidation and reduction of the OTHQ/OTQ redox couple. The surface-based quasi-reversible peak system was observed, corresponding to oxidation and reduction of the OTHQ/OTQ redox couple. The surface-based nature of the system was confirmed by the linear variation of peak currents with sweep rate. For instance, in aqueous 1 M HClO4 solution, a plot of $I_{p,a}$ vs. sweep rate ($ν$) was linear, with a slope of $(47 ± 3) \times 10^{-6} \text{ A V}^{-1}$ and a correlation coefficient of $r = 0.9963$. For solutions with pH values from 1 to 8, cyclic voltammograms were displaced towards lower potential values as the pH was increased. Plots of $E_{p,a}$ and $E_{p,c}$ vs. pH were linear with slopes of $(0.064 ± 0.002) \text{ V}$ and $(0.064 ± 0.003) \text{ V}$, respectively, indicating clearly that the oxidation/reduction mechanism of OTQ/OTHQ involves the same number of electrons and protons. On the other hand, $E_{p,a}$ and $E_{p,c}$ were practically independent of pH at pH > 8. In addition, the anodic and cathodic currents as well as the anodic and cathodic surface coverage decreased as the pH increased. The surface redox couple response was not observed at pH > 11. Based on these results, 10% ACN + aqueous 1 M HClO4 was chosen as the reaction medium to prepare the adsorbed OTQ, and 1 M HClO4 was chosen for in-depth study on the oxidation/reduction cycle of the adsorbed redox couple.

For these experiments, the working electrode was cycled five times in acidic solution containing the mycotoxin (10% ACN + 90% aqueous 1 M HClO4). Then, the electrode was rinsed with a 1 M HClO4 aqueous solution and transferred to another electrochemical cell which contained only a 1 M HClO4 aqueous solution. Cyclic voltammograms obtained under these experimental conditions at different $ν$ are shown in Fig. 4. A surface-based quasi-reversible redox couple was found, with an anodic peak potential of about 0.64 V. Similar behavior was observed when the electrode was immersed in a solution of OTA, in 10% ACN + 90% aqueous 1 M HClO4, for 60 s under an open circuit and then transferred to another electrochemical cell containing only aqueous 1 M HClO4. During the first anodic scan only one oxidation wave was found in the potential range from 1.2 to 1.4 V. This can be assigned to the oxidation of OTA adsorbed onto the electrode surface. In addition, during the first reverse cathodic scan, only one reduction wave for a surface process was observed, with potential range from 0.75 to 0.55 V, whose complementary anodic peak was found when the second consecutive potential scan was performed (Fig. 5). Successive scans caused an increase in current for this surface quasi-reversible peak system, as well as a decrease of the OTA main oxidation wave. Constant current values were obtained for the surface redox reaction after the third potential scan. Similar
The reference electrode was SCE. A voltammetric cell which contained OTA in 10% ACN + 90% 1 M HClO₄ (at different scan rates after cycling the working electrode five times in a separate solution of 1 M HClO₄, after immersion of the working electrode in a cell containing OTA). Our observation that ∆Eₚ was simultaneously nonzero and independent of sweep rate, suggests that slow charge transfer kinetics are not the origin of the observed response. Slow kinetics would cause ∆Eₚ to increase with faster sweep rate. Omnic effects are also not an explanation for the finite sweep rate independent of ∆Eₚ because a larger ohmic drop would be expected at higher sweep rates due to the passing of a larger total current. Furthermore, diffusion is effectively excluded as an explanation because the peak current increased linearly with ν, rather than with ν¹/₂ as expected for diffusion-controlled reactions. There have been previous reports of finite ∆Eₚ values that are independent of sweep rate for surface-confined redox couples [30]. This unusual quasi-reversibility or apparent non-kinetic hysteresis in cyclic voltammetry has been interpreted in terms of N-shaped free energy curves. In this interpretation, unusual hysteretic quasi-reversibility is a non-equilibrium behavior, in which a finite ∆Eₚ is observed because some rate processes are slow on the time scale of the experiment [31].

The areas under the oxidation and reduction peaks, corrected for any residual charging current, represent charge (Q) associated with the oxidation of OTHQ and the reduction of OTQ, i.e., 
$$Q_{ox} = nFA\Gamma_{OTQ}$$
and
$$Q_{red} = nFA\Gamma_{OTHQ},$$
where n is the number of exchanged electrons per mole of oxidized or reduced substance, F is the Faraday constant, A is the microscopic electrode area and ΓₒTQ and ΓₒTHQ are the surface concentrations of adsorbed oxidized and reduced substance, respectively. Average values of Qox and Qred of (2.7 ± 0.3) × 10⁻⁶ and (1.9 ± 0.2) × 10⁻⁶ C, respectively, were calculated from cyclic voltammograms recorded with scan rates from 0.015 to 0.100 V s⁻¹ when cOTA = 1.2 × 10⁻⁴ M was employed in the accumulation solution. Further, average values of (2.0 ± 0.2) × 10⁻¹⁰ and (1.4 ± 0.1) × 10⁻¹⁰ mol cm⁻² were obtained for ΓₒTHQ and ΓₒTQ, respectively, under these experimental conditions. The value obtained for ΓₒTHQ represents the saturation surface concentration, ΓₒTHQ (Fig. 6a), i.e., that corresponding to the formation of a monolayer of adsorbed substance (see below). In addition, a tentative expression for ΓₒTHQ for a surface redox couple can be expressed as in [27]:

$$\Gamma_{OTHQ} = \left(\frac{4RT}{n^2F^2A}\right) \left(\frac{\ln a}{\nu}\right)$$

An average value of ΓₒTHQ = 1.8 × 10⁻¹⁰ mol cm⁻² was obtained from cyclic voltammograms in the 0.010–0.050 V s⁻¹ range. This value is in a good agreement with the average value previously determined using the charges of the anodic peaks, as well as with expected values for adsorbed redox couples comprised of similarly sized molecules [27,28,30].

We also examined the isotherms and energetics of adsorption in order to better understand the behavior of this adsorbed redox species. A plot of ΓₒTHQ vs. cₒTHQ is shown in Fig. 6a. The surface concentration rose rapidly with cₒTHQ at low cₒTHQ and then asymptotically approached the limiting surface concentration. The surface saturation concentration arose at cₒTHQ ≥ 1.2 × 10⁻⁴ M. Fits of different adsorption isotherms models (Langmuir, Frumkin, Temkin and Freundlich) were tested to describe the adsorption of OTA anodic 

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**Fig. 4.** Cyclic voltammograms of OTHQ obtained in 1 M HClO₄ aqueous solution at different scan rates after cycling the working electrode five times in a separate electrochemical cell which contained OTA in 10% ACN + 90% 1 M HClO₄ (νₒOTA = 1.2 × 10⁻⁴ M). The working electrode was a GC disk of area A = 0.0707 cm², and the reference electrode was SCE. ν = (1) 0.010; (2) 0.015 and (3) 0.020 V s⁻¹.

**Fig. 5.** Consecutive cyclic voltammograms of OTQ/OTHQ recorded in a blank solution of 1 M HClO₄ after immersion of the working electrode in a cell containing OTA in 10% ACN + 90% aqueous 1 M HClO₄ for 60 s under an open circuit potential. (cₒOTA = 1.2 × 10⁻⁴ M), and ν = 0.050 V s⁻¹. (1) and (2) are the first and second scans, respectively.
oxidation product at GC electrodes. The best fits were obtained with the Frumkin adsorption isotherm, which is [27]:

\[ \beta c_{\text{OTA}}^* = \frac{\theta}{1-\theta} \exp(g'\theta) \]  

(2)

where \( \beta = \exp(-\Delta G_{\text{ads}}/RT) \) is the adsorption coefficient, which expresses the strength of adsorption, \( \theta = \Gamma_{\text{OTHQ}}/\Gamma_{\text{OTQ}} \), \( g' = 2g \Gamma_{\text{OTHQ}}/RT \) is a parameter characterizing the interaction between the adsorbed species, and \( \Delta G_{\text{ads}} \) is the Gibbs free energy of adsorption. A plot of \( c_{\text{OTA}}^* \) against \( \theta \) is shown in Fig. 6b. The fit was performed for values of 0.1 < \( \theta \) < 0.8 (solid line in Fig. 6b). For the best fit, the parameters were: \( \beta = (1.144 \pm 0.004) \times 10^5 \), \( g' = 0.70 \pm 0.02 \) and \( \chi^2 = 8.2 \times 10^{-18} \), where \( \chi^2 \) is the chi-square statistical parameter. There is excellent agreement between experimental data and the results of this fit. A value of \( \Delta G_{\text{ads}} = -28.4 \text{ kJ mol}^{-1} \) was obtained for the standard free energy of adsorption. This indicates that the overall adsorption process of OTQ molecules onto the GC surface is energetically favorable. In addition, a positive value for \( g' \) indicates that interactions between adsorbed species on the electrode surface are attractive.

3.2. Square wave voltammetry

SW voltammograms are highly sensitive to the charge transfer kinetics of surface redox processes [18,19]. They may be interpreted using two techniques. In the first technique, often called the “quasi-reversible maximum” method, the ratio between the peak current and the frequency is approximated as a parabolic function of the kinetic parameter, \( k \). In the second technique, the voltammetric peak is split with the SW amplitude at a given frequency [20,21]. According to theory, the maxima of these parabolic functions and the splitting of the voltammetric peak can be used to calculate the formal rate constant, the transfer coefficient and the formal potential of redox couples immobilized on the electrode surface [20–24]. Therefore, a combination of the two techniques is used here to carry out a full characterization of the thermodynamics and kinetics of the OTHQ/OTQ redox couple adsorbed on GC electrodes in 1 M HClO4 aqueous solution.

Forward (\( I_f \)), reverse (\( I_r \)) and net (\( I_n \)) currents were obtained from SW voltammograms of OTHQ adsorbed on GC electrodes after transferring the electrodes to a blank solution (1 M HClO4). Results for the potential region from 0.5 to 0.8 V are shown in Fig. 7 and present strong evidence for the surface-based quasi-reversible nature of this electrochemical signal [29,32].

These findings reveal that the OTA oxidation product selectively interacts with the carbon surface. In principle, this may be explained by the presence of oxygen-containing functional groups, such as carboxylic acid, lactone, o-quinone, p-quinone, carbonyl and phenols, on an activated GC surface [33]. The interaction between these functional groups and the CO and –COOH groups present in the chemical structure of OTQ (Fig. 1b) might be responsible for the adsorption of this electro-oxidation product onto GC electrodes. Such specific interactions have been previously observed in our laboratory for other mycotoxins [29,32].
3.2.1. Determination of thermodynamic and kinetic parameters for OTQ adsorbed at GC electrodes

Plots of $I_{p,n}/\Delta f$ vs. $f$ (where $I_{p,n}$ is the net peak current) obtained as previously described after OTQ accumulation onto the GC electrode, are shown in Fig. 8 for two bulk concentrations of mycotoxin.

The apparent reversibility of the confined redox reaction at the electrode surface depends on the kinetic parameter, $\kappa$, defined as the ratio between the standard (formal) rate constant ($k_s$) and the frequency ($\kappa = k_s/\Delta f$). Net peak currents from these SW voltammograms are linearly proportional to the frequency, but the factor of this proportionality is a function of the reaction reversibility. If the adsorptions of both the reactant and the product of a quasi-reversible redox couple are equally strong, a maximum appears in the plot of $I_{p,n}/\Delta f$ vs. $f$ or $1/\Delta f$, which appears at the SW frequency which is approximately equal to $k_s$ of the redox reaction [19,34]. In the region of the maximum, the curve can be approximated by a parabola, and if $(I_{p,n}/\Delta f)_{\text{max}} = k_s/\kappa_{\text{max}}$ [34]. Thus, the equation:

$$k_s = \kappa_{\text{max}}f_{\text{max}}$$  

is a convenient way to calculate $k_s$.

Also, theory predicts that a useful “quasi-reversible maximum” appears only if $-1.5 < \log \delta < 1.5$ [34], where $\delta$ is the ratio between the adsorption constants of the oxidized and reduced forms of the surface redox couple.

The theoretically calculated critical kinetic parameter, $\kappa_{\text{max}}$, depends on the transfer coefficient $(1 - \alpha)$, on the product of the SW amplitude and on the number of electrons, $n\Delta E_{\text{SW}}$, but it is independent of the normalized potential increment, $n\Delta E$, and of the amount of initially adsorbed reactant [34].

In this experiment, the critical frequency, $f_{\text{max}}$, which corresponds to the maximum of an $I_{p,n}/\Delta f$ vs. $f$ plot is nearly independent of $C_{\text{OTA}}$ in the accumulation solution. This behavior suggests that $\kappa_{\text{max}}$ does not depend on the surface coverage. The ratio between peak current and frequency reaches a maximum at $f_{\text{max}} = (12 \pm 2)$ Hz.

Theory also predicts that if the rate of reaction becomes rapid, the net peak current of a SW voltammogram splits and the peak height approaches zero [20]. The splitting of the net peak is caused by the skew of forward and reverse peaks on the potential scale, as the dimensionless rate constant increases. Experimentally, this behavior may be observed by decreasing the SW frequency or by increasing the SW amplitude [20]. However, changes in the peak shape, which carry kinetics information, are more effectively produced by varying the SW amplitude rather than varying the frequency [20]. Thus, a set of SWV experiments where $C_{\text{OTA}}$ and the SW amplitude were varied at a given frequency. The net, forward and reverse currents recorded at a frequency of 10 Hz and at different $\Delta E_{\text{SW}}$ are shown in Fig. 7. As is theoretically predicted [20,21], large changes in voltammogram shape are obtained when $\Delta E_{\text{SW}}$ increases for an appropriate fixed frequency. For small $\Delta E_{\text{SW}}$ (i.e., $25$ mV) and $f = 10$ Hz, a single net peak is observed. However, at that frequency the peak starts to split at about $\Delta E_{\text{SW}} = 50$ mV and is almost completely split at $\Delta E_{\text{SW}} = 100$ mV. The peak splits due to the relationship between the potential-dependent rate constants for the redox couple and the time scale of the experiment. An average value of $(0.605 \pm 0.003)$ V for the overall formal potential ($E'$) of the adsorbed two-electron redox couple was estimated from the individual potentials. The individual potentials were calculated from the splitting of the anodic and cathodic peaks at frequencies of 10 and 20 Hz (see Table 1), by considering that the two new peaks are symmetrically located about the formal potential of the surface redox couple [20].

It has also been reported that the influence of $(1 - \alpha)$ on $\Delta E_p$ is insignificant [35]. However, the form of the net SW response for an adsorbed reversible redox couple is influenced by the transfer coefficient [20,35]. For $(1 - \alpha) > 0.2$, the ratio between the forward (anodic) and reverse (cathodic) peak currents, $I_{p,a}/I_{p,c}$, can be approximated by a single exponential curve [35]:

$$\frac{I_{p,a}}{I_{p,c}} = 5.6414 \exp[-3.4606(1 - \alpha)]$$  

Experimental values for the anodic ($I_{p,a}$) and cathodic ($I_{p,c}$) peak currents, the ratio between $I_{p,a}$ and $I_{p,c}$ and the overall anodic transfer coefficient for the two-electron OTA oxidation product are shown in Table 1. The corresponding average values for these two parameters are $(I_{p,a}/I_{p,c}) = (1.2 \pm 0.3)$ and $(1 - \alpha) = (0.45 \pm 0.06)$, for frequencies of 10 and 20 Hz.

The overall $k_s$ value was calculated through Eq. (3) by using both methods previously mentioned for SW voltammetric analysis. The value $f_{\text{max}} = (12 \pm 2)$ Hz was taken from the first method, and $\kappa_{\text{max}}$ was extracted from Table 1 as in Ref. [21] by considering values obtained from the second method, i.e., $n\Delta E_{\text{SW}} = 0.050$ mV and $(1 - \alpha) = 0.45 \pm 0.06$. Based on these results, Eq. (3) predicts that $k_{\text{max}} = 0.89$ and that $k_s = 10.7 s^{-1}$ for the overall two-electron redox process. These values indicate slower kinetics for this mycotoxin compared with others studied in our laboratory, such as cercosporin ($k_s = 350 \pm 50 s^{-1}$ and $(1 - \alpha) = 0.50 \pm 0.03$ in aqueous 1 M HClO$_4$) [29] and altetroxin-1 ($k_s = 685 \pm 27 s^{-1}$ and $(1 - \alpha) = 0.48 \pm 0.03$, in 20% ACN + 1 M HClO$_4$) [32] on the same electrode surface. On the other hand, all transfer coefficients are in the 0.44–0.50 range.

3.2.2. Indirect determination of OTA

Some calibration curves of adsorbed OTA oxidation product generated using SWV are discussed here as a simple introduction for a future publication about the analytical applications of SWV in indirect determination of OTA in real infected matrices. The net current-potential curve in SWV is the most useful analytical signal for this purpose [36]. The high sensitivity of an adsorptive accumulation method is obviously its greatest advantage. The combination of adsorptive accumulation with SWV provides a very valuable electroanalytical tool for performing trace analysis of compounds which are both surface-active and also electroactive. Indirect quantification of OTA was carried out on GC electrodes in unstirred 1 M

![Fig. 8. Dependence of the ratio of net peak current and frequency ($I_{p,n}/\Delta f$) and the SW frequency obtained for OTHQ. The reaction medium was 1 M HClO$_4$ aqueous solution. $\Delta E_{\text{SW}} = 0.025$ and $\Delta E = 0.005$ V. The $C_{\text{OTA}}$ in the accumulation solution was: (a) $1.2 \times 10^{-5}$ M ($f_{\text{max}} = 14 s^{-1}$) and (b) $2.2 \times 10^{-4}$ M ($f_{\text{max}} = 11 s^{-1}$); $f_{\text{max}}$ was calculated from the first derivative of the mathematical expression for the parabola resulting from a best fit with experimental $I_{p,n}/\Delta f$ values.]
HClO₄ aqueous solution after cycling the working electrode five times in a solution containing mycotoxin in 10% ACN + 90% 1 M HClO₄ Values of Iₚ,n obtained from SW voltammograms recorded at different c²ₒTA were used to generate calibration curves. A linear relationship between Iₚ,n and c²ₒTA was obtained at f = 10 Hz between 6.2 × 10⁻⁸ and 6.0 × 10⁻⁶ M. One of the linear regressions obtained by a least square procedure is

\[ I_{p,n} = (7.5 \pm 0.3) \times 10^5 c_{\text{OTA}}^2 - (0.03 \pm 0.05), \quad (r = 0.9943) \]  

(5)

\[ I_{p,n} \] is expressed in μA and c²ₒTA has units of mol dm⁻³. Data used in the regression analysis of each calibration curve are the average of three repeated measurements (10 data points were taken). The percent relative standard deviation (RSD %) calculated from three calibration curves was 4%, and day-to-day reproducibility was 4.9%.

The experimentally determined detection limit (dl) measured experimentally was 1.24 × 10⁻⁸ M (5 ppb) with a signal to noise ratio of 3:1. Equivalently, in 10 mL solution, about 50 ng of OTA was indirectly detected. This sensitivity compares favorably with those obtained by other authors using electrochemical techniques. For example, the detection limit was 2.6 × 10⁻⁷ M (104 ppb) using SWV in acetate buffer solution with a pH of 4 [17]. The significant (21-fold) improvement in detection limit shows that the highly acidic medium and/or the electrochemical activation of the GC electrode surface makes the adsorption (pre-concentration) of OTA and its electrochemical discharge much more efficient.

### 4. Conclusions

We have demonstrated that the OTA electrochemical oxidation product (OTQ) obtained in a highly acidic reaction medium strongly adsorbs at the surface of activated glassy carbon electrodes.

The Frumkin adsorption isotherm best described the specific interaction of OTQ with glassy carbon electrodes. A direct 2e⁻/2H⁺ exchange for the redox couple in aqueous solution could be inferred from the dependence of the electrochemical behavior of adsorbed OTQ on solution pH and from the width at half-height of the anodic and cathodic voltammetric peaks. In addition, a multifaceted analysis of SWV results enabled full characterization of the thermodynamics and kinetics of the overall two-electron redox couple of the adsorbed species in the highly acidic reaction medium. These results show that SWV is a powerful technique to study redox phenomena of superficial nature.

Moreover, adsorptive accumulation of OTQ appears to be a very promising analytical tool for indirect determination of OTA in real samples.

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