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Diclofenac determination using CeO$_2$ nanoparticle modified screen-printed electrodes – A study of background correction

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ABSTRACT

The detection of low levels of drugs including Non-Steroidal Anti-Inflammatory Drugs (NSAIDs) in natural waters and wastewaters is becoming increasingly important. Electrochemical methods offer an effective means of detection, as they overcome many disadvantages associated with the currently available analytical methods. Cerium dioxide nanoparticles were synthesised and then incorporated onto the working electrodes of commercial graphite-based Screen Printed Carbon Electrodes (SPCE) then used to determine diclofenac levels. Following initial Cyclic Voltammetry studies, Square Wave Voltammetry (SWV) investigations were conducted over a range of conditions to optimise the peak potential separation and sensitivity of the method. The use of background correction as signal processing is highlighted since it constitutes a mandatory pre-treatment of data before the analysis of results. SWV study was carried out on diclofenac over a concentration range from 0.4 μM up to 26 μM which indicated that the response was linear with a limit of detection of 0.4 μM and a sensitivity of 0.058 μA/μM. The SWV method provides a rapid means of diclofenac detection where cerium dioxide nanoparticles combined with electrode vacuum heat treatment and use of background correction all play an important role.

1. Introduction

In the last decade, the investigation and use of electrochemical techniques in the analysis of pharmaceuticals drugs and other compounds have shown promising results. Such compounds and their degradation products may accumulate in an ecosystem causing harmful environmental damage. There is a wide range of analytical techniques available for quantification of pharmaceutical compounds, notably spectrophotometric and chromatographic techniques such as UV–Visible, Raman spectroscopy, Mass spectrometry, High-Performance Liquid Chromatography, Liquid Chromatography-Mass Spectrometry and Gas Chromatography among others [1,2]. When compared with other analytical techniques, electrochemical methods provide simple, rapid, selective detection using relatively low-cost instrumentation.

The efficacy of the electrochemical methods is largely determined by the electron transfer capability of the working electrode surface. Therefore, to improve the electrochemical performance of this electrode, a typical strategy is to design composites combining electrocatalytically active materials with conductive additives to modify the working electrode surface. Numerous materials have been developed with carbon-based materials showing particular promise. These include boron-doped carbon diamond electrodes (BDDE) [4], glassy carbon electrodes [5], carbon paste electrodes [3], graphene electrodes [6], modified glassy carbon electrodes and screen-printed carbon/graphite electrodes (SPCE) [7,8]. Currently, such SPCE find applications in diverse areas of electrochemistry; however, it has been used most often in the bioelectrochemical field for sensing applications [8,9].

The use of catalyst materials to improve the stability and conductivity of the WE has become an area of increasing interest for electroanalysis [5]. Diclofenac 2-[(2,6-dichlorophenyl)amino] benzene acetic acid sodium salt or DCF is well known as a Non-Steroidal Anti-Inflammatory Drug (commonly abbreviated as an NSAIDs) and is often used as an analgesic [10,11]. Due to its widespread use, diclofenac residues can be found in seawater, rivers and lakes [12–14]. Diclofenac and its residues have been identified as compounds of concern in aquatic systems particularly when accumulated and combined with by-products of DCF and other pharmaceuticals drugs [15]. Thus, the development of an electrochemical method for the detection of diclofenac and its degradation products along with many other pharmaceutical substances in-situ is extremely important for environmental monitoring. Electrochemical techniques using modified electrodes have

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been investigated for the estimation of levels of easily oxidised organic molecules such as diclofenac (DCF), acetaminophen (paracetamol ACOP), aspirin (acetylsalicylic acid, ASA), cocaine (COD), codeine (COD) and caffeine (CAF) when presented individually, as well as their simultaneous determination in certain pharmaceutical formulations [3,10,16,17]. For example, a research group from Brazil developed a working electrode using BDD electrode and Square Wave Voltammetry (SWV) for the determination of COC and screening of the most common adulterants (benzoic acid, caffeine, licorice, phenacetin, paracetamol, and procaine) in seized COC samples [4]. Khairy et al. developed a screen-printed electrode coated with cerium dioxide (CeO2) nanoparticles (NPs) which were utilised for the simultaneous determination in certain pharmaceutical formulations [3,10,16,17]. All measurements were conducted using a three-electrode one compartment configuration, in which the working electrode was a graphite-based screen-printed electrode with a geometric area of 4 mm diameter, and 15 ± 4 µm thickness. These screen-printed electrochemical sensors (SPCE) were purchased from Methom-DropSens, model DRP-110 with dimensions: 3.4 × 1.0 × 0.05 cm (Length × Width × Height). The counter electrode consisted of a graphite film, and a silver pseudo-reference electrode was also part of the three-electrode system. However, to avoid any interference due to the instability of the reference electrode (Ag-serves as a pseudo-reference electrode in this SPCE), an external reference electrode was utilised instead (Hg/Hg2Cl2, KCl (sat’d) – saturated calomel electrode (SCE) (from Radiometer model REF421). A Labquest2 digital pH meter was utilised together with a calibrated pH electrode to measure the pH of solutions. A Gallenkamp (from Fisstrem International Ltd) vacuum oven was used at 7.6 torr for further treatment of the sensors.

2.2. Instrumentation

The voltammetric measurements (both CV and SWV) were performed on an Eco Chemie B.V, Electrochemical Work Station, model Autolab PGSTAT 12 (The Netherlands) using GPES software (version 4.9). For analytical purposes, SWV is usually the ideal method for analysis as it suppresses the background (non-faradaic or capacitive) current effectively and also has higher sensitivity compared to other electrochemical techniques [24,25]. This method often leads to increases in anolyte peak height and enables much faster scan rates into the Vs⁻¹ range to be employed while diminishing the non-faradaic charging contribution [26]. In this work the use of CeO2 NPs modified SPCEs in conjunction with SWV for the detection of DCF is shown to be beneficial. Towards this goal, a novel and simple precipitation synthesis of CeO2 NPs was initially developed and then investigated using both Cyclic Voltammetry (CV) and SWV techniques in aqueous conditions pertinent to the analysis of DCF in environmental water samples, but not in pharmaceutical products or biological fluids. The role of background current correction which is especially important at higher potential values is also explored.

2.3. Preparation of the modified electrode (SPCE-CeO2)

In order to overcome some inherent variability in responses between different SPCEs, all SPCEs were subjected to an electrochemical pre-treatment (EP). This was done in an attempt to remove impurities and thus improve their electroactivity, and also to alter the SPCE WE surface wettability. A potential of +1.4 V (vs. SCE) was initially applied for 300 s in 0.05 M phosphate buffer solution (pH 7.2) to each SPCE WE, after which it was rinsed with deionised water. Such a pre-treatment has been reported previously [27–29]. For example, Moreno et al. reported that the EP can also prove beneficial through the introduction of new electrochemically active sites by removing impurities on the SPCE active working electrode surface [29].

A suspension of the synthesised CeO2 NP annealed at 900 °C (see supplementary information, Fig SI-2) was prepared by adding 0.5 mg of NPs in 1 mL of ultrapure water. The temperature of 900 °C was chosen based on the data provided in the supplementary information. Following the SPCE pre-treatment, 5 µL of this suspension was then drop cast directly on the WE surface of the SPCE by a micropipette and the sensor was then subjected to a heat treatment in a vacuum oven operating at 7.6 torr for 1 h at 100 °C, thus completing the preparation of the modified SPCE-CeO2. A summary of the procedure is shown in the flowchart in Fig. 1.

This vacuum heat treatment (VHT) was necessary to allow the NPs to strongly adhere to the WE surface causing the binder present in the SPE to melt and flow on the WE, thus avoiding dislodgement during the subsequent electrochemical analysis. Scanning electron microscopic images of the sensor surfaces were used to compare the morphological features of the WE surface of the screen-printed electrode, before and after adding the CeO2 NP. The SEM image of a bare SPCE, Fig. 2(a) indicated a porous, highly faceted structure. In contrast, the relatively smooth surface observed for the drop cast WE surface presented in

![Fig. 1.](image-url)
Fig. 2(b) was possibly due to the CeO$_2$ NP filling the graphite gaps and thus causing the smoother surface. Similar results have also been reported by Kadara et al. whereby a comparison between different electrochemical platforms was carried out [27]. All experiments were conducted at room temperature (22–25 °C) using a fresh electrode (SPCE) for each analysis. Moreover, all solutions were stirred for 60 s with an equilibration time of 30 s between each consecutive analysis.

2.4. Acquisition and presentation of voltammetric data

All results obtained by SWV are presented after a baseline-correction using the “moving average” algorithm (peak width = 0.03 V) included in the Autolab GPES software (version 4.9.007). This version can be found on the metrhom website (https://www.ecochemie.nl/) and is very effective when peaks appear as shoulders on steep flanks. After baseline correction, individual peaks can often be seen far more clearly. The curve fitting approaches are usually created using the simulation of a complex signal and the sum of single peak models, utilising least-squares minimization as reported by Jakubowska et al. [30]. This mathematical treatment improves the imaging and identification of the peaks above the baseline without the introduction of artefacts, although the peak current is in some cases decreased slightly (<10%) compared to that of the untreated curve [30]. In the quantitative voltammetric analysis, the peak height (the maximum current with respect to the baseline) is the most widely used parameter to determine concentrations. This parameter will, therefore, be utilised for further calculation [24].

3. Results and discussion

3.1. Cyclic voltammetry characterisation of the modified electrode

In order to determine and compare the electrochemically active surface area (EASA) of the SPCE-CeO$_2$, and the SPCE, a CV was conducted at 0.05 V s$^{-1}$ in a solution of 5 mM K$_4$Fe(CN)$_6$ in 0.1 M KCl. The electrochemical reaction of K$_4$Fe(CN)$_6$, is described according to the following equation for a quasi-reversible electron transfer process; the modified-quasi-reversible Randles–Ševčik (RS) equation [31]:

$$I_p = \frac{2.99 \times 10^5 n^{3/2} A^{1/2} D^{1/2} C^{1/2} \nu^{1/2}}{\alpha \cdot \langle n \rangle^{1/2}}$$

(1)

where $I_p$, n, α, A, D, C, ν are peak current in amps, n is the number of electrons transferred (n = 1 for [Fe(CN)$_6$]$_{3-}$/[Fe(CN)$_6$]$_{4-}$), α is the transfer coefficient (usually taken to be close to 0.5), electrochemically active surface area (cm$^2$), diffusion coefficient (cm$^2$ s$^{-1}$), concentration of ferrocyanide (mol/cm$^3$) and scan rate (V s$^{-1}$), respectively. The diffusion coefficient of potassium ferrocyanide is 6.3 × 10$^{-6}$ cm$^2$ s$^{-1}$ at 25 °C [31]. A typical CV is shown in Fig. 3, obtained at a scan rate of 0.05 V s$^{-1}$ and the resultant anodic peak current was used to calculate the EASA using the Randles–Ševčik (Eq. (1)). An alternative approach using the potential step (chronoamperometric) technique in conjunction with the Cottrell equation is often used to assess the EASA [32]. For an inner sphere reactant however such as potassium ferrocyanide or ascorbic acid, they showed that both approaches were comparable. Other recent investigations utilising the CV/Randles–Ševčik approach described herein have also been reported [19–22,32–35].

The original carbon surface of VHT SPCE (bare) is characterised by a large number of edges of graphite particles and a rather porous, rough surface as can be seen in Fig. 2. However, the modified SPCE-CeO$_2$ is far smoother in appearance and perhaps the nanoceria along with the heat treatment, made the WE surface more conductive with enhanced electron transfer rates. Such a statement can be confirmed with a better current response recorded in the voltammogram for the SPCE-CeO$_2$ electrodes.

The effective area of a bare SPCE and SPCE-CeO$_2$ was evaluated in order to compare the EASA for both electrodes. It has been reported that the WE active areas of SPCE from these DropSens SPCE are exposed edge plane-like sites/defects formed during its fabrication, and such changes can affect the roughness of the electrode surface [31,36]. Table 1 indicates the electrochemically active surface area of SPCE-CeO$_2$ and SPCE. It can be observed that the geometrical area of the
SPCE is 3.5 times greater than the EASA. This strongly suggests that SPCE contain only a small fraction of their surface which is electrochemically active, which is most likely to be edge plane graphite flakes protruding above an inert (non-conductive) polymeric binder matrix.

For reversible and quasi-reversible redox reactions such as the potassium ferrocyanide/ferricyanide (hexacyanoferrate II/III) couple, the heterogeneous rate constant $k^0$ can be estimated using Nicholson’s method \[37\]. For a single electron transfer, this is described by Eq. (2) below. In this method, the dimensionless rate parameter $\Psi$ is expressed as:

$$
\Psi = \frac{k^0}{\sqrt{\pi D v (F/RT)}} \gamma^{\nu/2}
$$

where $D$ is the diffusion coefficient of the ferrocyanide species, $\alpha$ is the transfer coefficient, $v$ is the scan rate and $F$, $R$, $T$ has the normal meanings. In addition, $\gamma$ is given by:

$$
\gamma = \frac{D^*}{D}
$$

where $D^*$ is the ferricyanide ion’s diffusion coefficient (7.6 × 10$^{-6}$ cm$^2$ s$^{-1}$) \[31,38\]. For this redox couple then $\gamma$ is equal to 0.83.

Using the cyclic voltammogram of 5 mM K$_4$[Fe(CN)$_6$] in 0.1 M KCl at the SPCE and SPCE-CeO$_2$ NP electrode obtained at a scan rate of 0.05 V s$^{-1}$ displayed in Fig. 3, the measured value of $\Delta E_p$ was 360 mV for both VHT electrodes.

Swaddle developed an empirical equation for $\Psi$ estimation from $\Delta E_p$ \[39\]

$$
\ln \Psi = 3.69 - 1.16 \ln(\Delta E_p - 59)
$$

For a $\Delta E_p$ of 360 mV this generates a value of $\Psi$ of 0.0534. Hence a value of $k^0$ of 3.4 × 10$^{-4}$ cm s$^{-1}$ is determined for the potassium ferrocyanide/ferricyanide couple at a sweep rate $v$ of 0.05 V s$^{-1}$.

This value of $k^0$ is comparable to those reported by Morrín et al. who quoted values of less than 10$^{-4}$ cm s$^{-1}$ for two commercial SPCEs displaying very large $\Delta E_p$ values (over 400 mV) \[38\]. They attributed their large SPCE $\Delta E_p$ values to poor charge transfer characteristics and hence measured low $k^0$ values. In the present work, the peak separation value ($\Delta E_p$ of 360 mV) is also very large, beyond the range of values of (61–212 mV) tabulated by Nicholson, including those used for quasi-equilibrium conditions.

Curiously the consistent value of $k^0$ found in our work for both SPCE and SPCE-CeO$_2$ NP seems to suggest that the CeO$_2$ NP does not affect the rate of electron transfer at the WE. This cannot be the case however, as the higher currents obtained in the SPCE-CeO$_2$ NP electrode displayed in Fig. 3 attest. There are a number of possible reasons for the existence of large $\Delta E_p$ values in SPCEs leading to low $k^0$ values. Aside from the poor internal charge transfer characteristics of the VHT SPCE, the existence of an ohmic drop (uncompensated resistance) may be the reason, a fact recognised by Nicholson where an additional correcting factor is suggested in an amended version of Eq. (1) \[37\]. Stevens et al. investigated this in a range of conditions including those without added supporting electrolyte recommending use of high concentrations (50 mM) of potassium ferrocyanide/ferricyanide \[31\].

Another explanation for such high peak separations is the conduction of the CV sweeps in aerated (oxygenated) conditions. For example significantly smaller $\Delta E_p$ values of less than 100 mV yielding $k^0$ values of 5.7 × 10$^{-3}$ cm s$^{-1}$ and 5.76 × 10$^{-3}$ cm s$^{-1}$ in a potassium ferrocyanide/ferricyanide redox couple on similar commercial SPCEs to those studied in this work were reported using deaerated electrolytes \[40,41\]. In contrast high $\Delta E_p$ were obtained in this work and others conducted in aerated conditions \[38\]. Under such oxidising conditions it may be that formation of Prussian Blue Fe$_4$[Fe(CN)$_6$]$_3$ (iron(III) hexacyanoferrate(II)) occurs, passivating the electrode surface \[42,43\].

Finally, it is worth noting that both the SPCE and SPCE-CeO$_2$/NP electrodes were subjected to a vacuum heat treatment before each CV sweep. Melting of the binder during the VHT in the electrodes leading to flow of the polymer matrix may have caused different surface

### Table 1

| Electrochemical Area A (cm$^2$) | Geometrical surface area WE (cm$^2$) |
|---------------------------------|------------------------------------|
| SPCE 0.0361                     | 0.1256                             |
| SPCE-CeO$_2$ 0.0833             | 0.1256                             |

\[Fig. 4.\] Square Wave Voltammogram showing the background current on an SPCE-CeO$_2$ WE in BRB (pH = 7) as a function of potential. SWV conditions: $f = 10$ Hz, $E_{\text{step}} = 10$ mV and $E_{\text{pulse}} = 25$ mV. Scan conducted from 0 to 1.5 V at a rate of 0.05 Vs$^{-1}$ (scan rate determined by the $E_{\text{step}}$ and the frequency). The CE was carbon ink and an external SCE served as a RE.
adsorption characteristics and re-orientation of the conductive graphite flakes within the electrodes.

An SWV plot of background current of the BRB pH 7 (blank) after the EP treatment of an SPCE-CeO2 is shown in Fig. 4. The parameters used in the SWV were a frequency of 10 Hz, potential step height of 10 mV and pulse height of 25 mV [26]. In this work, these conditions were optimised to improve the sensitivity and quality of the signal of the compounds to be analysed. The high overpotential for oxygen evolution (above +1.0 V) and possibly the instability of carbon electrodes under oxidising conditions invariably leads to a high background current, notably at higher potential values which could adversely affect the baseline correction at high potentials [44,45]. Such high background currents have also been observed at high potentials for other SPCE [3,18].

3.2. Electrooxidation of diclofenac

The electrochemical behaviour of DCF at the surface of the modified screen-printed electrode was investigated initially utilising the CV technique. CV scans were obtained in 100 µM DCF solution in 0.1 M BRB at pH 7 in the potential range from −0.4 to 0.8 V (vs SCE) at a scan rate of 0.05 V s⁻¹ and is shown in Fig. 5. A comparison was made between bare-SPCE (a) and the modified screen-printed electrode (a), with the CVs displayed in Fig. 5. The formation of an anodic peak (Ia) with the highest current at about 0.60 V (vs. SCE), is most probably due to the oxidation of the anionic form of DCF as the pKa is 4.18 and sodium salt was utilised in the CV resulting in the initial formation of a radical nitrogen species [46,47]. As the potential first scan was reversed, a cathodic peak (IIc) was observed at 0.26 V (vs. SCE). Aguilar-Lira et al. proposed that this peak corresponds to the reduction of 2-(2-hydroxyphenyl) acetic acid to 1-hydroxy-2-(hydroxyphenyl) ethanoate [48], although at pH 7 it is likely to be the acetate rather than the acetic acid form that is reduced. During the second and subsequent sweeps, a new anodic peak (IIa) appears at 0.34 V (vs. SCE). This indicates that peaks IIb and IIc are related to the same redox process [48]. Thus in the case of multiple scans peak IIa appears which upon the next forward scan oxidises, producing peak IIb. This process is described in the supplementary information, scheme SI-1 showing the initial formation of a nitrogen radical DCF species.

The voltammogram in Fig. 5 confirms the existence of a higher peak current for the modified electrode, which is likely due to the incorporation of CeO2 nanoparticles increasing the electrochemically active surface area of the working electrode and also enhancing the electrocatalytic activity of the CeO2 sensor for electrooxidation of the DCF molecule. DCF is irreversibly oxidised and thus there is no reverse peak associated with peak Ia. Such behaviour has also been reported elsewhere [6,11–13]. Since the peak (Ia) displays the highest current intensity (6.7 µA) and possesses better definition, it was chosen as an analytical signal for DCF quantification. It is worth noting that Aguilar-Lira et al. [48] selected peak IIa in their analysis which was closely associated with another peak IIc as part of a redox couple. In contrast peak Ia is an irreversible peak which is not associated with a comparable cathodic peak. It is also worth noting that in the presence of the CeO2 nanoparticles another anodic peak (IIIa) is evident at approximately 0.46 V (vs SCE) whilst a peak IIIc occurs at about −0.05 V (vs SCE). This peak has not been reported before and is obviously associated with the CeO2 NPs, as it is not present in the CV of the unmodified (bare) SPCE. Although the reactions responsible for the well-separated peaks are uncertain, one possibility is the formation of a polyaniline-like conducting polymer [49]. According to the reaction scheme proposed by both Cid-Cerón et al. [47] and Aguilar-Lira et al. [48], one of the products following the initial formation of the dehydrogenated nitrogen radical in the initial single electron transfer oxidation step is 2,6-dichloroaniline. Possibly catalysed by the CeO2 NPs on the modified SPCE, this may then react to form a poly-dichloroaniline species, although further work is required to confirm this. Alternatively, it could be a reaction involving one of the DCF degradation species reported by Brillas et al. in their study of the anodic treatment of DCF solutions [50].

Enhanced reactivity of SPCE-CeO2 after VHT could also possibly be explained by an increase of oxygen vacancies on the CeO2 surface. Such behaviour can change the concentration and recombination ratio of...
Ce⁴⁺ and Ce³⁺ species (see supplementary information, Fig S1-3) and in addition may introduce further levels of point defects due to the highly defective nature of the nanosized CeO₂ [51]. Choudhury et al. reported that annealing (at 100 °C) of CeO₂ NPs under vacuum introduces Ce³⁺ and oxygen vacancies in the CeO₂ lattice sites and on the crystal surfaces as well as on the grain boundaries.[52]. Such properties can enhance the catalytic performance of CeO₂, especially on the nanoscale. Thus the incorporated CeO₂ NPs stabilised the SPCE surface and promoted higher catalytic activity and enhanced sensor response.

3.2.1. Effect of scan rate

In order to better understand the nature of the reaction and the role of the potential sweep rate on the oxidation peak current (Ip of peak Ia) of DCF, CVs were performed at various selected scan rates in the range 0.05–1.5 V s⁻¹. As shown in Fig. 6(a), the anodic peak current of peak Ia increases with increasing scan rate and a slight shift in the oxidation peak potentials (Ep) occurs towards more positive values.

A linear relationship was observed between the anodic peak current (Ip with background correction) and the scan rate (ν) in Fig. 6(b). This indicates that the DCF oxidation process on the CeO₂ modified screen-printed electrode is adsorption-controlled [24,25]. A plot of Ip vs. ν¹/² shown in Fig. 6(c) is clearly curved (ie. non-linear). Finally, a plot of ln (Ip) vs ln (ν) in Fig. 6(d) is also linear, with a slope of 0.72, for sweep rates above 0.05 V/s. A slope of 0.5 in such double logarithm plots is thought to indicate a semi-infinite diffusion process, whilst a finite (thin layer) diffusion process possesses a slope of 1.0[13]. Thus a slope of 0.72 probably indicates the occurrence of a mixed process [40,53]

This suggests that an adsorbed species is involved and therefore peak Ia is not solely due to a diffusion-controlled reaction involving a solution-based molecule. Costa-Rama et al. reported that there are regions exposed to the bulk solution in which planar diffusion operates and a region in which “thin layer” diffusion effects occur for a carbon-based WE[54]. As indicated in Fig. 6(a), a reverse peak for peak Ia is often absent, apart from some scan rates (0.4 V/s, 0.5 V/s and 1.0 V/s) in which a peak appears at about −0.02 V (vs SCE) to 0.05 V (vs SCE) which is far apart from the peak Ia at about 0.6 V(vs SCE) to 0.7 V (vs SCE). This indicates that the electron transfer reaction associated with peak Ia is essentially irreversible [55].

With the knowledge that the oxidation reaction of DCF probably involves an adsorbed surface species process involving a single electron transfer, resulting in the formation of an adsorbed nitrogen radical, it is possible to estimate the heterogeneous rate constant k⁰. Originally proposed for an irreversible reduction of an adsorbed species by Laviron [56], this was later summarised in a review by Honeychurch and Rechnitz [57] who provided three diagnostic criteria for the

Fig. 6. (a) Cyclic voltammograms of DCF at SPCE-CeO₂ at different scan rates from 0.05 to 1.5 V s⁻¹ in BRB (pH = 7) and [DCF] = 100 µM. The SPCE-CeO₂ is the WE. CE was carbon ink and the external SCE was the RE. (b) Plot of Ip (from peak Ia) vs. ν with background correction for the oxidation of DCF at SPCE-CeO₂. (c) Plot of Ip vs. ν¹/² with background correction for the oxidation of DCF at SPCE-CeO₂. (d) Plot of ln (Ip) vs. ln (ν) with background correction for the oxidation of DCF at SPCE-CeO₂.
voltammetry of an adsorbed redox species undergoing irreversible adsorption kinetics.

(i) Linear plot of Ip vs v (as displayed in Fig. 6(b))
(ii) Plot of Ep vs ln v should also be linear with a slope of $(RT/\alpha_a F)$ for a single electron transfer oxidation reaction 
(iii) The peak potential separation at half height ($\Delta E_{1/2}$ in mV) is $62.5/\alpha_a$.

Compton et al. derived an expression for such a process allowing both the transfer coefficient $\alpha_a$ and the value of $k_0$ to be determined [58]. Considering the single electron transfer involving the oxidation of an adsorbed species in an irreversible manner, it is possible to derive the following expressions;

$$k^h = \left(\frac{\alpha_a F v}{RT}\right) \exp\left(\frac{RT}{\alpha_a F} \left(E_p - E_F^0\right)\right)$$ (5)

$$E_p = E_F^0 + \left(\frac{RT}{\alpha_a F}\right) \ln\left(\frac{\alpha_a F v}{RTk_0}\right)$$ (6)

Thus for a surface-bound adsorbed species undergoing an irreversible oxidation reaction, as $v$ (or ln $v$) increases, then so too does the peak potential $E_p$ according to the relationship described by Eq. (6). This equation predicts that a plot of $E_p$ versus ln $v$ will yield a straight line of gradient $\left(\frac{RT}{\alpha_a F}\right)$ enabling $\alpha_a$ to be determined from the slope. If however $E_p$ is plotted against $v$, then the intercept indicates the formal potential $E_F^0$ determined from Eq. (5).

From the results presented in Fig. 6, it is possible to estimate $\alpha_a$ from Eq. (6), $E_F^0$ from the linear extrapolation of $E_p$ versus $v$ to 0 V/s and then finally $k_0$. Initially, the slope of the plot of $E_p$ versus ln $v$ (See supplementary information, , Fig SI-4) was evaluated giving an $\alpha_a$ of 0.46 (estimated for sweep rates of > 0.05 V/s). Sweep rates above 0.05 V/s were used for this purpose, as this was the most linear section of the $E_p$ versus ln $v$ plot. Fotouhi et al. used a similar approach in their investigation of a modified MWCNT carbon paste electrode involving a surface-confined species for the detection of NADH [62]. Sweep rates above 100 mV/s were used by them to ascertain a $k_0$ of 2.239 s$^{-1}$. In the next step, extrapolation of a plot of $E_p$ versus $v$, to 0.0 V/s yielded an intercept ($E_F^0$) of 0.577 V (vs SCE). Then, using these values the heterogeneous rate constant $k_0$ was estimated to be 4.1 s$^{-1}$ based on an $E_p$ of 0.6 V (vs SCE) at a sweep rate of 1 V/s from Eq. (5).

There are many mechanisms proposed for the oxidation reaction of DCF in the literature. It is often described that the mechanism of this reaction is a reversible process of 2e$^-$, 2H$^+$ as reported by Goyal et al., whereby 5-OH diclofenac is formed [55]. In these studies, the DCF oxidation process begins with the protonated molecule. However, at pH 7 the predominant DCF species would be the deprotonated molecule (see supplementary information, , Fig SI-5). An alternative description is that given by Cid-Ceron et al. [47] and Aguilar-Lira et al. [48] involving an EC process at pH 7. Initially, the DCF reacts with water and after undergoing an oxidation reaction losing one electron and one proton. The nitrogen resonates between the creation of a double bond and formation of a radical (See supplementary information, , Scheme SI 1). This may then react with the medium (water), thus producing the irreversible DCF rupture into two products, 2,6-dichloroaniline and 2-hydroxyphenyl acetate.

3.2.2. Analytical performance

The SWV method was utilised for the determination of different concentrations of DCF solutions, using the modified SPCE-CeO$_2$ under the optimised experimental conditions. Following preliminary work conducted in BRB over a range of pH values, a pH of 7 was selected as optimal giving the best current response. The current values are obtained by automatically subtracting the background current. Fig. 7 shows that the peak current increases with successive additions of DCF. The determination of the limit of detection (LOD) and limit of quantification (LOQ) of the proposed modified electrode was also performed. The calibration curve is linear in the low concentration range of 0.4–1.6 µM with the regression equation: $I_p = 0.05787 C + 0.01323$. 

![Fig. 7. Square wave voltammograms of DCF at different concentrations in the range 0.1–100 µM. The SPCE-CeO$_2$ is the WE. A carbon ink CE and an external SCE was the RE. SWV was performed under the optimum parameters. Analysis conditions: f = 40 Hz, $E_{\text{step}} = 20$ mV and $E_{\text{pulse}} = 25$ mV. SWV performed in BRB (pH = 7).](image-url)
This has a correlation coefficient ($R^2$) of 0.9964. In the higher concentration range of 3.6–25.6 µM, another very good linear relationship was obtained with the regression equation:

$$I_p = 0.05226 \, C + 0.15738,$$

and a correlation coefficient ($R^2$) of 0.9969. The occurrence of two separate linear ranges for the detection of DCF was also reported by Shalauddin et al. [14]. Using differential pulse voltammetry (DPV) in conjunction with nanocellulose/f-MWCNT on glassy carbon electrodes, they studied DCF levels in pharmaceutical products and biological fluids reporting two linear calibration ranges, low (0.05–1 µM) and high (2–250 µM). These are similar ranges to those found in this work. Guzmán et al. also reported the existence of two linear ranges (low ~0–5 µM and high, 5–100 µM) in their investigation of DCF undertaken using DPV in pH 7 phosphate buffer solution [13].

No reasons were suggested however for the occurrence of the two distinct linear regimes. One possibility is that at low concentrations the adsorption of the reactant DCF species is only sufficient for a sub-monolayer level of reactant to adsorb, whereas at higher concentrations the reactivity is hindered as layers build up on the top or outermost surfaces, thereby shielding some reactive sites and effectively hindering electron transfer processes.

In the present investigation the LOD and LOQ were based on 3 s/m and 10 s/m respectively, where s is the standard deviation of the peak currents (n = 3, three runs) and m is the slope of the calibration curve [64]. The LOD was estimated to be 0.4 µM and 0.12 µM was found to be the LOQ. The sensitivity was determined as 0.058 µA/µM. This indicated that DCF can be estimated with a high level of confidence in the given concentration range (0.4–25.6 µM) using this SWV/SPCE-CeO$_2$ NP system.

### 3.2.3. Signal processing effects

The analytical conditions of a range of different modified electrodes for the determination of DCF using a variety of materials as the WE and a variety of nanoparticles for enhanced sensitivity are compared in Table 2. As can be observed in this Table, the LOD value that was obtained for the sensor fabricated in this work is either lower than or similar to those reported for other electrochemical sensors. However, this comparison strictly should only be carried out taking into account other factors such as sensor sensitivity, simplicity and its low cost of fabrication. Also, the difference in background correction procedure used and the number of data points utilised varies widely, as reported in Table 2. It is well known that the background correction can have a huge effect on the current response and peaks detected. However, it is rarely dealt with in the literature and consequently, comparison of LOD should not be directly undertaken without taking this into account.

Furthermore, the number of data points (not the range of concentration) selected to obtain the regression equation (and hence determine the LOD) is not the same within existing reports. However, in the majority of studies of electrochemical sensors typically the number of data points varies from 3 to 10 [1–5,15–16]. In the table above the number of data points selected for all studies was 6 or 7, and in this work, it was 7 as can be seen in Fig. 8.

It can be seen in Fig. 9, that it is necessary to carry out background correction in order to reveal the true magnitude of the peaks. It should be pointed out that in the reports cited in Table 2, no mention is made of how the background correction was carried out or it is often mentioned that it was carried out simply by subtracting the background current. Error in peak height determination is more evident at low peak sensitivities, thus generating larger inaccuracies at lower concentrations.

### 3.2.4. Stability and reproducibility of the modified electrode

The long term stability of the CeO$_2$-SPCE was evaluated by measuring the voltammetric current response of a fixed concentration of DCF (100 µM) after the modified electrode was stored for 2 weeks in a refrigerator at 4 °C and the peak current was measured every week. A relative standard deviation (R.S.D) of about 15% was observed. The reproducibility of the modified sensor was evaluated by using five independent electrodes and checking the current response of a 100 µM DCF solution. Here the R.S.D value obtained was 2.7%, indicating excellent reproducibility of the method. The repeatability of the modified electrode was investigated. The precision of the method was evaluated by repeating five sequential measurements in the same solution containing 100 µM of DCF using the same SPCE-CeO$_2$/NP. The R.S.D was found to be 8.5%, indicating that SPCE-CeO$_2$/NP is more suitable as a disposable electrode.

### 3.2.5. Selectivity and interfering species

The work reported in this paper focussed on the detection of DCF in aqueous conditions using a modified SPCE-CeO$_2$/NP sensor together with SWV in a BRB of pH 7. In order to assess the selectivity of the method, comparison can be made to the voltammetry of DCF reported in the literature. For example, Shalauddin et al. [14] reported that diclofenac had an $E_p$ of about 0.65 V (vs Ag/AgCl (3 M)), very close to the values found in this work, whereas dopamine displayed an $E_p$ of about 0.42 V (vs Ag/AgCl (3 M)). Thus the peaks were separated by over 0.2 V. Ascorbic acid, displaying a well-separated peak at approximately 0.17 V (vs Ag/AgCl (3 M)) was also reported. This peak is more than 0.45 V distant from the diclofenac peak. Besides, they also reported that uric acid and various carbohydrates (fructose, glucose, lactose and sucrose), even when in 100 fold excess did not interfere with the diclofenac oxidation peak.

In the case of xanthine in biological fluids, most investigations appear to have been conducted at slightly lower pH 5 PBS solution with $E_p$ values of near 0.8 V (vs Ag/AgCl) which is more than 0.15 V more positive than the diclofenac $E_p$ value found in this work [20,21]. Another study using Co-doped CeO$_2$ nanoparticles, in an electrochemical sensor for the simultaneous determination of hypoxanthine, xanthine, and uric acid reported similar $E_p$ values to those given in these references [23]. Guzmán et al. [13] found that ibuprofen did not interfere with DCF detection at any level. Ascorbic acid, citric acid, oxalic acid and sodium dodecyl sulfate only interfered at concentrations greater than or equal to that of DCF. Paracetamol (acetaminophen, ACOP) however was found to cause the most interference and the authors recommended the use of the standard addition method to minimize potential interference from paracetamol when determining DCF in an electrochemical sensor. For the SPCE-CeO$_2$/NP electrodes however it was found that it was possible to detect paracetamol, aspirin and

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### Table 2

Comparison of the analytical conditions of different modified electrodes for the determination of Diclofenac.

| Method                        | Technique | Linear range (µmolL$^{-1}$) | LOD (µmolL$^{-1}$) | Sensitivity (µA/µM) | Reference |
|-------------------------------|-----------|----------------------------|-------------------|--------------------|-----------|
| Reduced graphene oxide (rGO)/Co(OH)$_2$-nano-flakes rGO/CHNF/CPE | C.V      | 3.7–140                    | 0.05              | 1.370              | [6]       |
| Au-PtNPs/f-MWCNTs/AuE         | DPV       | 0.5–1000                   | 0.30              | 0.012              | [11]      |
| MWCNTs/Co(OH)$_2$/NPs/IL /GCE | DPV       | 0.18–119                   | 0.04              | 0.014              | [12]      |
| MWCNT/CPE                    | DPV       | 0.1–100                    | 0.001             | 0.290              | [13]      |
| F-MWCNTs/nanocellulose/GCE   | DPV       | 0.05–250                   | 0.012             | 2.619              | [14]      |
| CeO$_2$-SPCE                 | SWV       | 0.1–25.6                   | 0.40              | 0.058              | This work |
Fig. 8. Plots showing linear dependence of the anodic peak current as a function of DCF concentration in the range of (a) 0.4–1.6 µM and (b) 3.3–25.6 µM (number of replicates = 3).

Fig. 9. Square wave voltammograms of DCF 10 mg/L (a) without background correction and (b) with background correction. Using the SPCE-CeO_2 as the WE, carbon ink CE and an external SCE as a RE. SWV analysis conditions: $f = 40 \text{ Hz}$, $E_{\text{step}} = 20 \text{ mV}$ and $E_{\text{pulse}} = 25 \text{ mV}$. SWV performed in BRB (pH = 7).
caffeine in a common pharmaceutical product using SWV under the same conditions [63]. None of these compounds showed a current response near the DCF value of 0.6 V (Vs SCE) (as is evident in the supplementary information, Fig S1-6). Further exploration of the selectivity of the modified SPCE-CeO2/NP sensor and its response to a range of other compounds that may be found in natural or wastewaters would be worthwhile.

4. Conclusions

The SPCE-CeO2/SWV electrochemical method outlined in this paper shows many advantages over other sensors reported in the literature; notably a simple fabrication process, and hence the possibility of large scale-production, use of relatively low-cost materials, and the potential for use over wide concentration ranges of DCF. The main advantage of this electrochemical sensor is that it can be used portably, possibly in the field with a small portable potentiostat instrument utilising a nanoparticle sized catalyst (CeO2) that is relatively inexpensive and abundant, in conjunction with a commercially available SPCE. From a practical perspective the main advantage of SWV over many other analytical techniques is that accurate analyses can be carried out in a short time (of the order of seconds as fast scan rates can be used) without the need for bulky expensive equipment, solvents and special gases. When the baseline correction (the moving average) in a high potential region is performed, there is an inherent interference with the peak height of the analyte, resulting from the presence of high background currents due to the onset of oxygen evolution and possibly carbon oxidation reactions, thereby making the determination of compounds above +0.8 V unreliable. In the case of DCF however which undergoes an oxidation reaction at around 0.2 V less than this figure, this problem is greatly diminished using SWV combined with background correction. This allows an accurate assessment of DCF levels to be undertaken with good sensitivity and reproducibility. This approach shows considerable promise as a means of carrying out a quick and accurate assessment of DCF levels in environmental water samples, although it may also possibly be extended to the determination of DCF levels in pharmaceutical and/or samples of biological interest. Finally, an investigation of the selectivity of the modified SPCE-CeO2 NP sensor is worth pursuing to accurately determine the role of possible interfering species and EIS analysis as another method to determine kth values for charge transfer reactions.

CRediT authorship contribution statement

Rafaela C. de Carvalho: Writing - original draft, Conceptualization, Methodology, Software, Investigation, Validation, Visualization, Data analysis. Anthony J. Betts: Writing - review & editing, Supervision, Conceptualization, Data analysis. John F. Cassidy: Writing - review & editing, Supervision, Conceptualization, Data analysis.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

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