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# A comparative study of different electrodeposited NiCo<sub>2</sub>O<sub>4</sub> microspheres anchored on a reduced graphene oxide platform: electrochemical sensor for anti-depressant drug venlafaxine

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Two different fabrication methods were performed and compared for preparation of binary metallic oxide microstructures supported on a reduced graphene oxide (rGO) modified graphite electrode. Nickel–Cobalt oxide microspheres (NiCo<sub>2</sub>O<sub>4</sub> MSs) were prepared by two different deposition methods: wet chemical and *in situ*-electrical deposited methods. Different characterization methods were conducted, including cyclic voltammetry (CV), scanning emission microscopy (SEM), electrochemical impedance spectroscopy (EIS) and Raman spectroscopy. The deposition methods of NiCo<sub>2</sub>O<sub>4</sub> MSs were found to affect the electrochemical behavior of the modified electrodes towards the oxidation of venlafaxine (VEN), an anti-depressant drug. The fabricated electrode showed linearity over the range 5–500 nmol L<sup>-1</sup> and an excellent sensitivity with a limit of detection (LOD) and limit of quantitation (LOQ) of 3.4 and 10.3 nmol L<sup>-1</sup>, respectively. It was revealed that the wet-NiCo<sub>2</sub>O<sub>4</sub>@rGO modified electrode prepared by the wet chemical method showed an improved electrochemical behavior for determination of VEN in pharmaceuticals and human plasma with high recovery results in the range of 96.7–98.6% and 96.0–100.7%, respectively without any interference from the co-existing components.

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#### Introduction

Venlafaxine (VEN), an anti-depressant drug, is a selective serotonin-norepinephrine reuptake inhibitor (SNRI) used in the treatment of depression, social phobia, generalized anxiety and panic disorder. It increases the concentration of the neurotransmitters serotonin and norepinephrine in the brain.<sup>1,2</sup> However, an overdose of VEN leads to serious adverse effects; it might cause symptoms of depression, serotonin toxicity, seizure, or cardiac conduction abnormalities.<sup>3</sup> To avoid the adverse effects and toxicity, monitoring its therapeutic efficiency, as well as the drug level in body fluids should be carefully followed. Hence, the main aim of this research is to develop a simple, highly sensitive and cost effective analytical method to quantify VEN in biological fluids.

After a careful literature survey, it was revealed that most of the analytical methods for determination of VEN are HPLC methods. However, these methods face some drawbacks including the cost of equipment, consumption of large volumes of solvents and requirement of several pretreatment procedures for biological samples. Also, thin layer chromatography (TLC) was described for VEN analysis in bulk and capsules. Many

Pharmaceutical Analytical Chemistry Department, Faculty of Pharmacy, Assiut University, Assiut, 71526, Egypt. E-mail: marwazahry@aun.edu.eg; Tel: +20-88-2411520 spectrophotometric methods<sup>14-22</sup> and spectrofluorimetric methods<sup>23,24</sup> have been reported for quantitation of VEN as well. Capillary electrophoresis was used for stereoselective analysis of VEN and its metabolites<sup>25</sup> and for determination of the toxic levels of VEN in human serum.<sup>26</sup>

Only few research articles were reported for the utility of electrochemical techniques in the determination of VEN<sup>27-31</sup> especially in biological fluids.<sup>27-30</sup> Two of them used hanging mercury drop electrode (HMDE) as working electrode<sup>27,28</sup> which suffers from several drawbacks such as toxicity, fluctuation of mercury drop surface area, and its limited applications in analysis of more positive potential range.

Pencil graphite electrode (PGE) has several attractive advantages compared with HMDE or other carbon-based electrodes. It has high electrochemical reactivity, good mechanical rigidity, low background current, low cost, commercial availability, offer simple polishing procedure easy in modification and it is considered user and environmental-friendly electrode.<sup>32–34</sup>

Due to its remarkable electrochemical properties, bimetallic oxides such NiCo<sub>2</sub>O<sub>4</sub> have been used in different fields as electrocatalyst<sup>35,36</sup> and in drug delivery system.<sup>37</sup> They have tunable chemical and physical properties, low cost, low toxicity and natural availability. Additionally, they show remarkable magnetic and electrochemical behaviors differing from monometallic oxide.<sup>35</sup>

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Recently, a lot of efforts have been made to improve the properties of the prepared NiCo2O4 nanostructures which related to its aggregation behavior that reduces its applicability. In order to reduce their aggregation and increase their potentialities, many nanoplatforms have been developed as backbone substrates to support these nanostructures. Graphene is a monolayer of carbon atoms used for two decades in modification of electrode. Graphene oxide (GO) is considered one of the most promising substrates due to its ability to bind with nanoparticles via its extra-ordinary chemical and physical properties. Comparing with other carbon-based platforms, such as multi-walled carbon nanotubes (MWCNTs), GO is considered the best because of its availability, cheapness and the ease of preparation. Removal oxygen containing functional groups from GO surface producing the reduced form of graphene oxide (rGO); having a lot of advantages; high surface area and excellent electrocatalytic performance compared with GO.38-40 Different reduction methods have been reported for preparation of rGO; such as chemical or thermal, or electrochemical reduction.39 Among carbon based electrode, rGO is considered an excellent material owing to its outstanding properties; such as high conducting activity, large surface to volume ratio, and chemical stability and inertness.32,40

The presented article describes a comparative study of utilizing two different fabrication approaches to prepare integrated microstructures of binary metallic nickel/cobalt oxide anchored on reduced graphene platform. Different characterization methods such as SEM, EIS and Raman spectroscopy were carried out to evaluate the morphology, spectral properties and the electrochemical behavior of the resultant hybrid microstructures modified electrode. Square wave voltammetric (SWV) curves were recorded using the fabricated wet-NiCo2O4@rGO modified sensor for analysis of VEN in pharmaceutical formulation and in human plasma without any interference from the complicated matrix. The proposed SWV method showed high sensitivity, selectivity, and accurate results when compared with previously reported electrochemical methods<sup>27-31</sup> for VEN determination.

## **Experimental**

#### Chemicals and reagents

Venlafaxine was kindly gifted from Mash premiere company, Ni  $(NO_3)_2 \cdot 6H_2O$  and  $Co(NO_3)_2 \cdot 6H_2O$  were obtained from Sigma Aldrich. Venlaxin® tablet (containing 75 mg VEN per tablet) and Venllamash® capsule (containing 75 mg VEN per capsule) were purchased from local market.

Graphene oxide (GO) was synthesized following the modified Hummer's method.38 Potassium ferrocyanide with purity of 98% was purchased from Alpha Chemika, India.

All chemicals were of analytical reagent grade and were used as received. Deionized water was used for the preparation of standard solutions and electrolytes.

#### **Apparatus**

Using a Princeton VersaSTAT MC (VersaSTAT 3, Model RE-1, Princeton Applied Research, AMETEK, USA) controlled by 394 software in conjunction with a PAR Model polarographic analyzer, the electrochemical measurements were carried out. A three-electrodes cell containing bare or modified pencil graphite electrode PGE as a working electrode, Ag/AgCl (3 mol L<sup>-1</sup> KCl) as a reference electrode and a platinum wire as an auxiliary electrode was employed. Z-view software (Z-view version 3.5d) was applied for electrochemical impedance spectroscopy calculations.

The electrical contact with the pencil was conducted using a metallic wire wrapped to the metallic part of the lead inside the pencil. The pencil electrode (with a diameter of 0.5 mm and a length of 60 mm) was then fitted vertically and immersed into the electrolyte solution inside the electrochemical cell. The electrochemical measurements were carried out at room temperature (25  $\pm$  5 °C).

Surface morphological studies of bare and modified electrodes were investigated using scanning electron microscope (SEM), JEOL JSM-5400 LV instrument (Oxford, USA). Raman spectra were monitored for GO and the fabricated in situ and wet-NiCo2O4@rGO using a Bruker Senterra Raman microscope (Bruker, Optics Inc., Germany) equipped with 785 nm excitation using a Nikon objective ( $\times 20$ ) and a holographic grating with 1200 lines per mm along with a charge coupled device detector (CCD). The spectra were recorded over the range 400 to 2000 cm<sup>-1</sup> using 8 s integration time.

All pH-metric measurements were made on Hanna pH meter (Hanna Instruments, São Paulo, Brazil) digital pH-meter with glass combination electrode. OriginPro 8.5 software was used for graphing and analyzing the experimental data.

#### Fabrication of NiCo2O4 microspheres anchored on rGO modified PGE

Using wet chemical method. A weight of 2 mg of GO powder was transferred to a mixture solution containing 5 mL of  $0.01 \text{ mol } L^{-1} \text{ Ni(NO}_3)_2 \cdot 6H_2O \text{ and } 5 \text{ mL of } 0.02 \text{ mol } L^{-1}$  $Co(NO_3)_2 \cdot 6H_2O$ , the contents were mixed well. The pH of the solution was adjusted to 11 using ammonia solution (25%), and kept in agitation for 30 min. The solution was further heated at 90 °C and kept at this temperature for 10 min. After cooling to room temperature, the precipitate was filtered, washed with deionized water three times and dried in desiccator overnight.

The resultant NiCo2O4@rGO were electrodeposited on the surface of PGE by applying a potential ranged from -0.9 to 1.5 V for 30 s. The modified PGE electrode is denoted with wet-NiCo2O4@rGO.

Using in situ-electrical deposition method. Initially, in order to produce rGO modified PGE; a suitable concentration of GO was transferred into the electrochemical cell and an electrical deposition was carried out over the range from -1.5 to 1.5 V using  $100 \text{ mV s}^{-1}$  scan rate. The deposition cycles were repeated 7 times and the electrode was denoted with rGO/PGE. The content of the electrochemical cell was replaced with a suitable volume of Ni/Co solution mixture (0.02 mol  $L^{-1}$ ) in 0.1 mol  $L^{-1}$ 

KCl and a reduction potential ranged from -0.1 to -1.3 V (for 5

cycles using 50 mV s<sup>-1</sup> scan rate) was given for in situ electrodeposition of NiCo2O4 microspheres over the surface of rGO/ PGE, the electrode was further denoted as in situ-NiCo2O4@rGO.

Electro-analytical procedure for quantitation of VEN. The quantitation of VEN was carried out using two modified pencil electrodes following the previous procedures. The electrochemical cell was filled with  $0.04~\mathrm{mol~L}^{-1}~\mathrm{Britton}$ Robinson (BR) buffer at pH 10.5 and supplemented with suitable amount of VEN. Further; the voltammograms were recorded using wet-NiCo2O4@rGO or in situ-NiCo2O4@rGO modified electrodes. The developed SWV plots recorded at wet-NiCo2O4@rGO, were monitored using a potential range -0.2 to 1.5 V with different experimental parameters after optimization procedures.

#### **Applications**

Paper

Determination of VEN in pharmaceutical formulations. The contents of 20 tablets of Venlaxin® or 20 capsules of Venllamash® (containing 75 mg VEN per formulation) were combined and weighed. An accurately weighed amount of the finely powdered equivalent to 75 mg of VEN of each formulation was transferred to 25 mL volumetric flask separately and made up to volume with methanol. The solution was sonicated for 20 min, filtrated and the filtrate was diluted with methanol to obtain the required concentration that lies within the calibration range. The analytical procedure was conducted as mentioned under analytical procedure.

Determination of VEN in human plasma. Drug-free human plasma samples were collected from four healthy donors and stored at −4 °C until analysis. Each 0.5 mL of the plasma was transferred to 2 mL Eppendorf tube, spiked with 0.1 mL of VEN standard solution at different concentration levels. Then, 1.0 mL of acetonitrile was added for plasma protein precipitation. After vortexing for 20 s, the contents were separated by centrifugation at 4000 rpm at room temperature for 20 min. The clear supernatant was filtered through 0.45 µm membrane filter (Millipore-Whatman, Kent, UK), transferred into a 10 mL volumetric flask and diluted with 0.04 mol L<sup>-1</sup> BR buffer at pH 10.5. Further, appropriate amount of the extracted samples were transferred to the electrochemical cell for determination of VEN. The application on human plasma samples was performed in accordance with the Declaration of Helsinki and approved by the Egyptian Network of Research Ethics Committees (ENREC). Informed consent was obtained for any experimentation with human subjects.

Electrochemical impedance spectroscopy (EIS) measurements. EIS measurements for bare PGE, wet-NiCo2O4@rGO and in situ-NiCo<sub>2</sub>O<sub>4</sub>@rGO modified electrodes were monitored using Princeton VersaSTAT MC. The EIS were recorded with the frequency range 0.1 Hz to 10 kHz using 1.0 mmol L<sup>-1</sup> potassium ferrocyanide/ferricyanide system dissolved in 0.5 mol L<sup>-1</sup> KCl. The amplitude of the applied sine wave potential was 10 mV. EIS results were fitted to Randles equivalent circuit and obtained as Nyquist plots using Z-view software for analyzing EIS results.

#### Results and discussion

#### The electrocatalytic behavior of VEN on different carbon electrodes

The effect of various types of carbon based electrodes, such as glassy carbon electrode (GCE), bare PGE and modified PGE, was investigated towards the electro-oxidation of VEN. Additionally, wet chemical or in situ-electrically deposited modified supported on PGE electrode were utilized to examine the electrooxidation behavior of VEN.

According to Fig. 1, VEN has a weak oxidation peak current at GCE (0.57 V), while a slightly intense and sharp oxidation peak current was provided at PGE bare (0.59 V). Using the wet-NiCo<sub>2</sub>O<sub>4</sub>@rGO modified electrode, the peak was shifted to 0.68 V and the intensity of the peak was increased by 4-fold compared with bare PGE. It can be noticed that wet-NiCo2-O4@rGO modified electrode showed the highest sensitivity in terms of VEN oxidation peak current intensity. This shift could be attributed to the electrodeposited layer of composites on PGE, and the change in the charge transfer rate between the analyte and the modified NiCo<sub>2</sub>O<sub>4</sub> electrode. These results confirm the significance of the modification step, suitability, and sensitivity of fabricated sensor for trace analysis of VEN in different matrices.

The prepared wet-NiCo2O4@rGO which was selected for subsequent work was electrically deposited on PGE surface, where different variables such as initial potential and deposition time were studied and optimized. Initial potential values ranged from -1.0 to +1.5 V and deposition time in the range from 10 to 90 s were investigated and it was concluded that the optimum results were obtained using initial potential of -0.9 Vfor 15 s as deposition time.

#### Characterization of the modified PGE

Upon synthesis of wet-NiCo2O4@rGO, hydroxyl ions will be released from ammonia on heating and consumed in complexation with metal salts (Ni2+ and Co2+). The concentration of the free metals is decreased in the formation of metal hydroxides. When it is mixed with GO solution, the metal ions attached to the negative surface of GO platform which is reduced to rGO due to the effect of ammonia.41

#### Morphological characterization

The morphology of cross sections of bare PGE, in situ-NiCo2-O4@rGO and wet-NiCo2O4@rGO modified electrodes were carried out by SEM technique. As shown in Fig. 2A, thin fragile layers of graphite with smooth surface morphology could be seen for bare electrode. In case of in situ-NiCo2O4@rGO modified electrode (Fig. 2B), the surface is covered with few microsphere like structures. Fig. 2C represents the wet-NiCo<sub>2</sub>O<sub>4</sub>@rGO modified one, the hybrid composite represented as microsphere like structures which are anchored allover PGE surface in the form of cavities and protrusions owing to the deposition of NiCo<sub>2</sub>O<sub>4</sub> and rGO structures. Such modified structures may provide a high surface area of the interfacial surface between the analyte and the electrode; leading to increase the capability

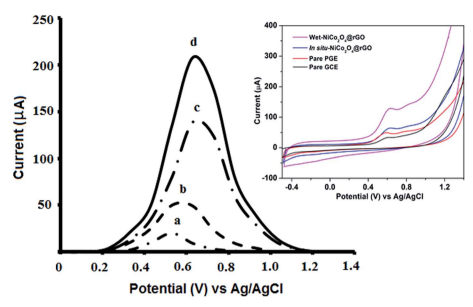


Fig. 1 Square wave voltammograms of 0.1  $\mu$ mol L<sup>-1</sup> VEN solution at (a) bare GCE, (b) bare PGE, (c) in situ-NiCo<sub>2</sub>O<sub>4</sub>@rGO and (d) wet-NiCo<sub>2</sub>O<sub>4</sub>@rGO in 0.04 mol L<sup>-1</sup> BR buffer (pH = 10.5) under optimum conditions. The upper inset: cyclic voltammograms of 0.1  $\mu$ mol L<sup>-1</sup> VEN recorded at different electrodes.

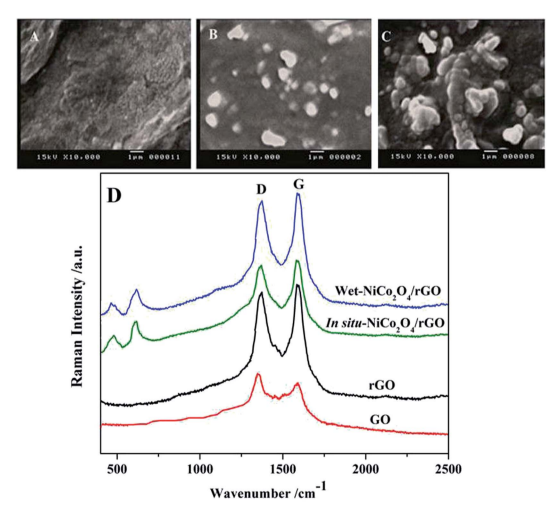


Fig. 2 SEM images of (A) bare PGE, (B) in situ-NiCo<sub>2</sub>O<sub>4</sub>@rGO, (C) wet-NiCo<sub>2</sub>O<sub>4</sub>@rGO surfaces and (D) Raman spectra of GO, rGO, in situ and wet-NiCo<sub>2</sub>O<sub>4</sub>/rGO composite.

of hosting larger amount of drug; resulting in an increase in the oxidation peak current of VEN.

In Fig. 2D, Raman spectra were recorded for GO, rGO, and the composites of *in situ* and wet-NiCo<sub>2</sub>O<sub>4</sub>/rGO. Two significant bands at 1350 (D band) and 1580 cm<sup> $^{-1}$ </sup> (G band) assigning to the carbon bands were observed in the GO and rGO spectra. In case of both types of NiCo<sub>2</sub>O<sub>4</sub>/rGO, it clearly shown new bands appeared around 506 and 653 cm<sup> $^{-1}$ </sup> which assigned to the vibration of Ni–O and Co–O bonds.<sup>35,42</sup>

# Electrochemical characterization of the modified electrode using ferrocyanide-ferricyanide system

Ferrocyanide–ferricyanide redox couple (1.0 mmol  $L^{-1}$  Fe  $(CN)_6^{3-/4-}$  in 0.5 mol  $L^{-1}$  KCl) was utilized to examine the electrochemical performance of bare PGE, *in situ* and wet-NiCo<sub>2</sub>O<sub>4</sub>@rGO electrodes.

Electrochemical impedance spectroscopy (EIS) experiments as meaningful tool for studying the interface properties of the modified electrodes were conducted. The EIS results were fitted to a Randles indented as equivalent circuit. The circuit composed of solution resistance  $(R_s)$ , the charge transfer resistance  $(R_{ct})$ , and Warburg element  $(W_s)$ . Fig. 3 exhibits the Nyquist plots of bare PGE and modified electrodes. Nyquist plot describes charge transfer resistance  $(R_{ct})$  across the electrode surface. The  $(R_{ct})$  is equal to the diameter of the semicircle portion which corresponds to electron-transfer limited process across PGE electrode. Rct values were measured for bare, in situ-NiCo<sub>2</sub>O<sub>4</sub>@rGO and wet-NiCo<sub>2</sub>O<sub>4</sub>@rGO. Comparing with in situ-NiCo2O4@rGO electrode, the Nyquist diameter of the semicircle portion  $(R_{ct})$  of the bare PGE electrode (curve a,  $R_{ct}$  $\sim$ 3600  $\Omega$ ) is much larger than that deposited with in situ-NiCo<sub>2</sub>O<sub>4</sub>@rGO (curve b,  $\sim$ 2600  $\Omega$ ) indicating an acceleration of electron transfer. While deposition with wet-NiCo<sub>2</sub>O<sub>4</sub>@rGO, the  $R_{\rm ct}$  is markedly decreased to about  $\sim$ 1400  $\Omega$  (curve c) which is agreed with the fact that the deposited layer of wet-NiCo<sub>2</sub>O<sub>4</sub>@rGO provided an increase in the electron transfer owing to their excellent conductivity.

It is well known that the electrochemical performance of any electrode is mainly proportional to the real contact surface area with the solution. So the main aim of the following section is to calculate the surface active area of the studied electrodes using Randles–Sevcik equation.<sup>43</sup>

$$I_{\rm P} = (2.69 \times 10^5) n^{3/2} A_{\rm eff} D_{\rm R}^{1/2} C_0 \nu^{1/2}$$

where n is the number of electron transfer,  $A_{\rm eff}$  is the surface area of the electrode (cm²),  $D_{\rm R}$  is the diffusion coefficient (cm² s<sup>-1</sup>),  $\nu$  is the scan rate (V s<sup>-1</sup>) and  $C_0$  is the concentration of Fe(CN)<sub>6</sub><sup>3-/4-</sup> (mol cm<sup>-2</sup>). According to Randles-Sevcik equation, the surface active area of bare PGE, in situ-NiCo<sub>2</sub>O<sub>4</sub>@rGO and wet-NiCo<sub>2</sub>O<sub>4</sub>@rGO has been calculated to be: 46, 100.3 and 163.2 mm². The results revealed that the prepared hybrid composite was successfully deposited (as confirmed by SEM images Fig. 2) on the PGE surface giving high surface area; hence hosting more drug for oxidation on the modified surface, leading to enhancement of the current peak on fabricated wet-NiCo<sub>2</sub>O<sub>4</sub>@rGO electrode towards VEN oxidation.

#### Effect of scan rate

The effect of scan rate on the electro-oxidation of VEN at the wet-NiCo<sub>2</sub>O<sub>4</sub>@rGO was studied by CV technique. The cyclic voltammograms of 0.1  $\mu mol~L^{-1}$  VEN in BR buffer (pH 10.5) using modified electrode at different scan rates varying from 0.05 to 1.0 V s $^{-1}$  were recorded and presented in Fig. 4. It can be seen that the anodic oxidation peak potential is shifted to more

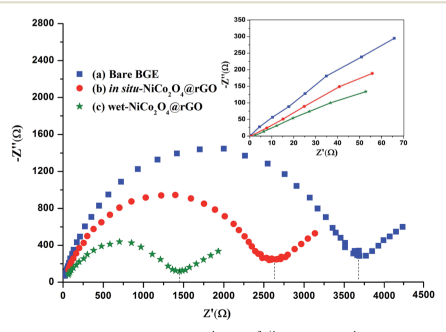


Fig. 3 Nyquist plots of electrochemical responses of 1.0 mmol  $L^{-1}$  Fe(CN)<sub>6</sub><sup>3-/4-</sup> in 0.5 mol  $L^{-1}$  KCl recorded at (a) bare PGE, (b) in situ-NiCo<sub>2</sub>O<sub>4</sub>@rGO and (c) wet-NiCo<sub>2</sub>O<sub>4</sub>@rGO at scan rates 100 mV s<sup>-1</sup>. The upper inset: enlarged view at low impedance range.

**RSC Advances** Paper

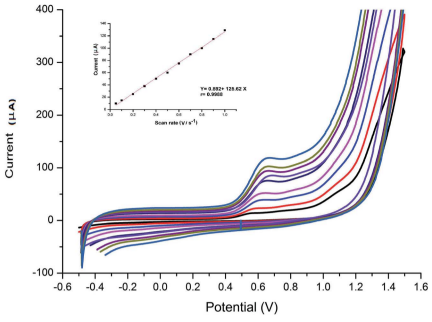


Fig. 4 Cyclic voltammograms of  $0.1 \,\mu\text{mol} \, L^{-1} \, \text{VEN}$  solution at different scan rates (0.05 to 1.0 V s<sup>-1</sup>) recorded at wet-NiCo<sub>2</sub>O<sub>4</sub>@rGO electrode. Inset: relation between scan rate and oxidation peak current of VEN.

positive values by increasing the scan rate along with an increase in current signal indicating the irreversibility of the electro-chemical oxidation of VEN. According to the following equation, the peak current is directly proportional to scan rate. Good linearity was obtained between the oxidation peak current  $(I_{\rm P})$  and the scan rate  $(\nu)$ , with high correlation coefficient equal to 0.9988, suggesting that the electrode reaction is controlled by adsorption rather that diffusion.

$$I_{\rm P}(\mu A) = 0.892 + 125.62\nu (r = 0.9988)$$

Further, the effect of logarithm scan rate ( $\log \nu$ ) on the logarithm peak current ( $\log I_P$ ) was examined giving a straight line as shown in Fig. 5, with a slope of 0.953, very close to the theoretical value of 1,

$$\log I_{\rm P} (\mu A) = 0.953 \log \nu + 2.09 ({\rm V s}^{-1}) (r = 0.9977)$$

This result implied that the electrode process is adsorption controlled which is matched with the previously reported studies on VEN oxidation.29,30

Moreover, a linear relationship between  $E_P$  and  $\log \nu$  was observed (Fig. 5) following the next equation:

$$E_{\rm P}({\rm V}) = 0.06 \log \nu ({\rm V s}^{-1}) + 0.682 (r = 0.9929)$$

According to Laviron equation;44 a relationship between the potential and the scan rate; the number of electron (n) transferred in the rate limiting step can be deduced from the following equation:

Slope = 
$$2.303RT/\alpha nF$$

where T is the absolute temperature (298 K),  $\alpha$  is the electron transfer coefficient, n the number of electrons transferred in the rate determining step, F is the Faraday constant (96.480 C  $\text{mol}^{-1}$ ) and R is the universal gas constant (8.314 J  $\text{mol}^{-1}$  K<sup>-1</sup>). The charge transfer coefficient ( $\alpha$ ) identifies the potential effect on the activation energy of an electrochemical reaction, and its acceptable value ranged between 0.3 and 0.7 in most electrochemical systems. In case of our system, the value of  $\alpha n$  can be easily calculated from the slope of  $E_P$  versus  $\log \nu$  plot, which was found to be 0.06 and  $\alpha n$  value was calculated as 1.05. In case of totally irreversible reactions, it has been reported<sup>29-31</sup> that ( $\alpha$ ) equals 0.5; hence the number of electrons transferred in the oxidation process will be  $\sim$ 2. These calculations agreed well with the reported values in the oxidation reaction of VEN.

Since the electrochemical oxidation of VEN is known to occur by transferring two electrons, thus the number of protons involved was also predicted to be two. These results suggested that two electrons and two protons participated in the irreversible electrochemical oxidation of VEN NiCo2O4@rGO.29-31

#### Optimization of experimental and electrochemical parameters

Effect of supporting electrolyte and pH. Various buffer solutions, such as borate, phosphate and Britton-Robinson (BR) buffer solutions were studied. From all the investigated buffer solutions, BR buffer gave the best response in terms of higher oxidation peak current intensity and better peak shape. The influence of pH on the oxidation peak of VEN was investigated in the range from 8.0 to 11.0 using BR buffer on wet-NiCo<sub>2</sub>O<sub>4</sub>@rGO. As can be seen from Fig. 6A, by increasing pH, the oxidation peak potential of VEN was shifted toward negative

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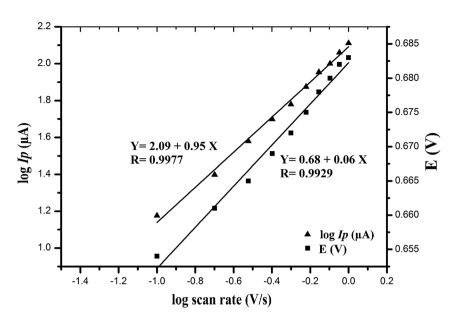


Fig. 5 Dependence of the logarithm of peak current (log  $I_P/\mu$ A) and the oxidation peak potential (E/V) of VEN on logarithm of scan rate (V s<sup>-1</sup>).

potentials which can be explained by the change in the protonation state of the acid-base function of VEN molecule during the oxidation process, indicating that the number of electron transferred was equal to the proton number involved in the electrode reaction process. The linear plot between pH values and the potential is represented in Fig. 6B. Thus, the probable electro-oxidation mechanism of VEN is shown in Scheme 1. The loan pair of electrons in the quaternary nitrogen of VEN is more susceptible to be oxidized than that in oxygen and in aromatic ring. However, oxidation of aromatic  $\pi$ -electrons in VEN is more favorable to happen as the nitrogen lie in the side chain has no other group for stabilization after the loss of electrons whereas the loss of electron from aromatic ring produces cationic radical which can be easily stabilized by the oxygen attachment to the ring.  $^{29,30}$ 

Further optimization of buffer concentration was carried out using different concentrations ranged from 0.02 to 0.10 mol  $\rm L^{-1}$ . The best oxidation peak response was observed in the range from 0.04 to 0.08 mol  $\rm L^{-1}$  BR buffer. Hence, 0.04 mol  $\rm L^{-1}$  was selected as the best buffer concentration.

**Square wave voltammetric parameters.** Various SWV variables such as pulse height over the range of 3–25 mV, frequency within the range of 50–250 Hz, and step height over the range from 3 to 25 mV were studied. The obtained results showed that the pulse height of 5 mV, frequency of 225 Hz and step height of 15 mV were the optimum parameters. These optimum parameters showed higher current peak intensity and consequently provided sensitive quantitation of VEN. Also, the effect of the accumulation potential and accumulation time on VEN oxidation peak was examined. The accumulation potential,  $E_{\rm acc}$  was investigated between -0.8 V and 1.0 V. The oxidation peak current was increased from -0.8 V to -0.6 V and then remained stable till -0.2 V followed by a gradual decrease. Therefore,  $E_{\rm acc}$  of -0.5 V was selected for further subsequent analysis. The

effect of accumulation time;  $t_{\rm acc.}$  had also a great influence on VEN oxidation peak current and it was studied from 10 to 60 s. The peak current of VEN recorded using wet-NiCo<sub>2</sub>O<sub>4</sub>@rGO modified electrode was increased with  $t_{\rm acc}$  up to 30 s after that the peak current intensity was decreased. This behavior may be attributed to the blocking effect of the electrode surface leading to low amount of VEN get oxidized with increasing the accumulation time, so 30 s was selected as an optimum accumulation time.

**Validation study.** Under the optimum experimental and electrochemical conditions, the proposed SWV method was validated according to ICH guidelines<sup>45</sup> for the following parameters:

**Linearity.** Under the optimized conditions, the modified electrode showed a linear response in the concentration range from 0.5 to  $50 \times 10^{-8}$  mol L<sup>-1</sup> with good correlation coefficient of 0.9989. Detection and quantification limits were statistically estimated as  $3.3\sigma/b$  and  $10\sigma/b$ , respectively, where b is the slope and  $\sigma$  is the standard deviation of intercept. The proposed method showed detection and quantification limits of 3.4 and 10.3 nmol L<sup>-1</sup>, respectively. The proposed sensor showed higher sensitivity for determination of VEN when compared with previously reported electrochemical methods.<sup>27–31</sup> Fig. 7 shows the typical SWV of different concentrations of VEN recorded at wet-NiCo<sub>2</sub>O<sub>4</sub>@rGO modified electrode. All statistical data were calculated and summarized in Table 1, the low values of LOD and LOQ reflect high sensitivity of the proposed modified electrode for electro-oxidation of VEN.

**Precision.** The precision of the proposed SWV method was determined through inter- and intra-day repeatability measurements. Intra-day precision was studied by repeating the analysis of three concentration levels of the working solutions of VEN. While inter-day precision was examined by analyzing three concentration levels of the working solutions of VEN over

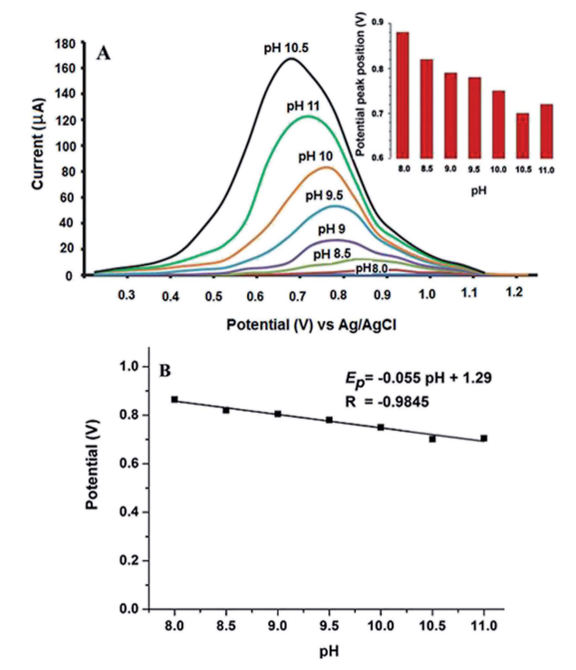


Fig. 6 (A) Effect of different pH values on square wave voltammograms of  $0.1 \,\mu$ mol L $^{-1}$  VEN solution in  $0.04 \,$ mol L $^{-1}$  BR buffer. The upper inset: histogram of potential peak position against pH values. (B) Linear plot between peak potential (V) and pH values of supporting electrolyte.

a period of three working days. The intra- and inter-day precision results were expressed as percent relative standard deviation (% RSD) which was found to be less than 2.97%, indicating good repeatability and reproducibility of the developed method as shown in Table 1.

Interferences study. In order to examine the selectivity of the proposed method for determination of VEN, the influence of some potentially interfering substances found in pharmaceuticals or biological samples was investigated. The tolerance limit for these interfering species was considered as the maximum concentration that gave a relative error less than  $\pm$ 

5.0% at a concentration of  $0.1~\mu mol~L^{-1}$  of VEN under optimum conditions. As shown in Table 2, the results inferred that 100-fold excess of ascorbic acid, citric acid, starch, glucose, sucrose, uric acid and magnesium stearate had no significant effect on the oxidation peak current of VEN (signal change less than 4%). The results obtained showed recoveries over the range 95.5-103.2%, indicating that VEN can be determined in pharmaceuticals and biological samples with high selectivity without any interference from most common interfering substances using the modified electrode.

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Scheme 1 The probable mechanism of VEN electro-oxidation.

#### Application of the proposed method

Determination of VEN in pharmaceutical formulations. The fabricated wet-NiCo2O4@rGO electrode was successfully utilized to determine VEN in pharmaceutical tablets. Table 3 shows the results and it can be seen that good recoveries were

obtained in the range of 96.7-98.6% with good agreement with the claimed amount without any interference from pharmaceutical excipients. The obtained results were statistically compared to those obtained by reported method.23 The calculated t and F values at 95% confidence level were less than the tabulated ones indicating that there is no significant difference

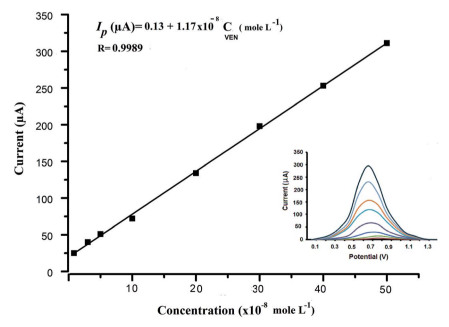


Fig. 7 Square wave voltammograms of different concentrations of VEN (0.5 to  $50 \times 10^{-8}$  mol L $^{-1}$ ) recorded at wet-NiCo<sub>2</sub>O<sub>4</sub>@rGO. Optimum parameters:  $0.04 \text{ mol L}^{-1}$  BR buffer (pH = 10.5),  $E_{\text{acc}} = -0.5 \text{ V}$ , frequency = 225 Hz, pulse height = 5 mV, step height = 15 mV,  $t_{\text{acc}} = 30 \text{ s}$ . Inset: calibration curve between VEN concentration and peak current.

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Table 1 Quantitative statistical parameters of the proposed SWV method for determination of VEN

Parameters	The calculated value
Linearity range ( $\times 10^{-8} \text{ mol L}^{-1}$ )	0.5-50
Intercept $(\mu A) \pm SD^a$	$0.13\pm0.121$
Slope $(\mu A) \pm SD^b$	$1.17\pm0.312$
Correlation coefficient $(R)$	0.9989
$LOD^c$ (×10 <sup>-9</sup> mol L <sup>-1</sup> )	3.4
$LOQ^{d} (\times 10^{-9} \text{ mol } L^{-1})$	10.3
Intra-day precision <sup>e</sup> (%RSD) $n = 3^f$	≤2.55
Inter-day precision <sup>e</sup> (%RSD) $n = 3^f$	≤2.97

<sup>&</sup>lt;sup>a</sup> Standard deviation of intercept. <sup>b</sup> Standard deviation of slope. <sup>c</sup> Limit of detection. d Limit of quantitation. Relative standard deviation. f Mean values of three measurements at three different concentration levels.

Table 2 Interference study using different concomitant substances found in the pharmaceutical formulations of VEN

	Ratio		
Concomitant substances	VEN : concomitant	Recovery $\% \pm SD^a$	
Ascorbic acid	1:10	$96.6 \pm 0.16$	
	1:100	$98.5 \pm 0.76$	
Citric acid	1:10	$99.0\pm0.72$	
	1:100	$103.2\pm0.38$	
Starch	1:10	$98.6 \pm 1.24$	
	1:100	$99.4 \pm 0.92$	
Magnesium stearate	1:10	$101.5\pm0.46$	
8	1:100	$97.5 \pm 0.97$	
Uric acid	1:10	$95.5\pm1.22$	
	1:100	$96.4 \pm 1.05$	
Glucose	1:10	$97.5 \pm 0.37$	
	1:100	$97.8 \pm 0.88$	
Sucrose	1:10	$97.9 \pm 1.12$	
	1:100	$98.1\pm0.53$	
<sup>a</sup> Average of three deterr	ninations.		

with respect to accuracy between the proposed method and the reported one.

**Determination of VEN in human plasma.** The applicability of the proposed method for determination of VEN in human plasma was attempted. The results presented in Table 4 confirm the efficiency of wet-NiCo<sub>2</sub>O<sub>4</sub>@rGO for determination of VEN in plasma sample. The obtained results were ranged from 96.0 to 100.7%, and RSD percentages were excellent, indicating the

Table 4 Determination of VEN in spiked human plasma by the proposed SWV method

Spiked concentration $(\mu \text{mol L}^{-1})$	Found concentration $(\mu \text{mol L}^{-1})$	% Recovery <sup>a</sup>	RSD%
0.03	0.029	96.7	2.43
0.10	0.098	98.0	1.74
0.20	0.192	96.0	1.56
0.30	0.296	98.7	1.82
0.40	0.394	98.5	2.53

suitability of the developed wet-NiCo2O4@rGO modified electrode for VEN determination in biological samples without any interference from co-existing substances in plasma samples. The achieved results ensure the suitability, high sensitivity and cost effective of the proposed SWV method for the analysis of VEN in biological fluid when compared with previously reported electrochemical methods.27-31

Stability and reproducibility. Stability of the developed wet-NiCo<sub>2</sub>O<sub>4</sub>(a)rGO was investigated by immersing the electrode in BR buffer (pH 10.5) for 14 days and SWVs were monitored and compared with the voltammogram of VEN ( $20 \times 10^{-8}$  M) before immersion in buffer. The results show that the current peak remained stable, which indicating excellent stability of wet-NiCo2O4@rGO electrode.

In order to study the repeatability and reproducibility, the RSD of six repetitive measurements of  $20 \times 10^{-8}$  M VEN using the same modified electrode was found to be 1.57%. Additionally, six modified electrodes were prepared and used for determination of  $20 \times 10^{-8}$  M VEN. The RSD was calculated and found to be 2.38%. These results indicate that the developed electrode has high repeatability and reproducibility in synthesis procedure and voltammetric quantitation.

#### Comparison of the proposed with previously reported methods

Table 5 shows a comparison between the analytical behavior of the proposed method and the previously reported methods for quantitation of VEN either in pharmaceutical dosage forms or in biological fluids. As shown, the proposed method has a large advantage over the reported methods considering linearity range, and LOD values. Additionally, using PGE provides another advantage of the proposed method, in terms of its

Table 3 Determination of VEN in pharmaceutical formulations using the proposed and reported methods

Drug	Labeled content (mg)	Recovery $^a$ (%) $\pm$ SD			
		Proposed method	Reported <sup>b</sup> method	<i>F</i> -Test <sup>c</sup> value	<i>t</i> -Test <sup>c</sup> value
Venlaxin®	75	$96.7\pm1.2$	$97.4 \pm 1.6$	2.346	0.634
Venllamash®	75	$97.4 \pm 1.5$	$98.6 \pm 1.3$	1.126	0.279

<sup>&</sup>lt;sup>a</sup> Average of 5 determinations. <sup>b</sup> Ref. 23. <sup>c</sup> Theoretical values at 95% confidence level, t-value = 3.182 and F-value = 2.36, respectively.

Table 5 Comparison between different electroanalytical methods for VEN determination with the present method

Electrode	Linearity working range	LOD (mol L $^{-1}$ )	Ref.
Mercury film microelectrode Hanging mercury drop electrode (HMDE) Multi-walled carbon nanotubes NAF-CNT-GCE Multiwall carbon nanotubes-ionic liquid gel modified GCE	$1.27 \times 10^{-6}$ to $24.3 \times 10^{-6}$ $3.58 \times 10^{-7}$ to $6.48 \times 10^{-6}$ $10 \times 10^{-6}$ to $500 \times 10^{-6}$ $3.81 \times 10^{-8}$ to $6.22 \times 10^{-5}$ $2.0 \times 10^{-6}$ to $2.0 \times 10^{-3}$	$6.93 \times 10^{-7}$ $3.58 \times 10^{-7}$ $4.70 \times 10^{-7}$ $1.24 \times 10^{-8}$ $1.69 \times 10^{-6}$	27 28 29 30 31
NiCo <sub>2</sub> O <sub>4</sub> @rGO modified PGE electrode	$5 \times 10^{-9}$ to $500 \times 10^{-9}$	$3.40 \times 10^{-9}$	The present work

simplicity, high electrochemical reactivity, cost effective and more environmentally friendly electrode.

#### Conclusion

In the present work, a comparative study using two different deposition approaches for synthesis of NiCo2O4@rGO modified graphite electrode was developed. The differences between the two modified electrodes, wet-NiCo<sub>2</sub>O<sub>4</sub>@rGO and in situ-NiCo<sub>2</sub>-O4@rGO modified electrodes were characterized by various techniques; cyclic voltammetry (CV), electrochemical impedance spectroscopy (EIS), scanning electron microscopy (SEM) techniques and Raman spectroscopy. The wet-NiCo2O4@rGO modified electrode is considered an easy and excellent sensing tool to be utilized for voltammetric determination of VEN, an antidepressant drug. In addition, different electrochemical parameters were studied and optimized. The SWV method was validated according to ICH guidelines and showed higher sensitivity (LOD =  $3.4 \text{ nmol L}^{-1}$ ) with high precision (% RSD < 2.97). The proposed method was successfully applied to determine VEN in pharmaceutical formulation and in human plasma. No interference from matrix of real samples was observed. The developed electrode presented excellent stability and reproducibility. Bimetallic microspheres hybrid with a highly conductive reduced graphene oxide (wet-NiCo<sub>2</sub>O<sub>4</sub>@rGO) probe is considered a selective and sensitive tool for trace analysis of VEN in different matrices.

#### Conflicts of interest

The authors declare that there is no conflict of interest.

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