INTRODUCTION

Electrode surface modification is a field of paramount importance in the modern electrochemistry, especially due to the various application possibilities of modified electrodes. In recent years, chemically modified glassy carbon electrodes have received increasing attention due to their potential applications in various analyses and also due to its relative ease of electrode preparation and regeneration [1]. Electrode surfaces coated with electropolymerized conducting polymer films have been paid great attention in the past, due to their unique physical and chemical properties and some possible applications in electrocatalysis, organic batteries, and microelectronic devices. Studies have indicated that surface coated polymer films exhibit enhanced analytical response for the quantification of various biological and clinical analytes. The thickness, permeation and charge transport characteristics of the polymeric films can be controlled by the potential and current applied [2–15]. Research on surface modified electrode has involved studies of the electrochemistry of the attached molecules, the catalysis and inhibition of various electrochemical processes and specific applications to such widely varying areas as photoelectrodes and analytical determinations [16, 17].

In recent years, modification of electrode surfaces has been an important research area in electrochemistry. Compared with other electrode concepts in electrochemistry, the distinguishing feature of a chemically modified electrode is that generally a thin film of a selected chemical is bonded or coated onto the electrode surface to endow the electrode with the chemical, electrochemical, optical, electrical, transport, and other desirable properties of the film in a rational, chemically designed manner [18, 19]. One of the methods used for the modification of electrode surfaces is electropolymerization. Electropolymerization can accelerate transmission of electrons onto the surface of the electrode; it has high selectivity and sensitivity due to the film homogeneity in electrochemical deposition, and it has strong adherence to the electron surface and large surface area [20, 21]. Researchers have employed polymeric film modified electrodes to detect organic and inorganic molecules in recent years. Electrochemical methods, such as differential pulse polargraphy (DPP), stripping voltammetry (SV), differential pulse voltammetry (DPV) and square-wave voltammetry (SWV) have been widely applied for the determination of pharmaceuticals [22–30]. In the present investigation, a simple, effective and sensitive electrochemical method for the determination of ibuprofen on polyaniline nanofiber modified electrodes is explored.

Experimental

Chemicals and apparatus

All reagents were of AR grade purchased commercially. Solutions were prepared using deionized double distilled water. Stock standard solution of ibuprofen was prepared in 50% ethanol. A standard stock solution of ibuprofen (1000 ppm) was prepared. The voltammetric studies were carried out in exploratory and determination mode on a software connected CH Instruments Electrochemical Workstation (model CH 650C). The voltammetric cell consisted of a three electrode assembly with polymer modified glassy carbon electrode as a working electrode, a platinum wire as an auxiliary electrode and Ag/AgCl electrode as a reference electrode. Nitrogen gas was purged through the solution for 5 min. A Hanna instrument pH/ORP meter was used for pH measurements.

Modification of the electrodes

A GCE (3-mm diameter) was polished using 1.0 and 0.05 mm alumina slurry and rinsed thoroughly with Milli-Q water. Ultrasonic agitation for 30 min of 2.0 mg of chemically prepared nanostructured polyaniline in 2 ml of water gave a homogeneous green solution. 20 µl of this solution was placed on the GCE surface. The electrode was then dried at room temperature to obtain a polymer modified GCE.
Pharmaceutical sample preparation

One tablet of containing ibuprofen were weighed, powdered and then placed into a 250 ml of the conical flask; warm water was added into the flask. The sample was swirled to dissolve for 30 min in sonicator and left cool. The sample solution was filtered through a filter paper (Whatman No.42) into 100 ml volumetric flask. An aliquot of the solution was then analyzed according to the proposed voltammetric procedure.

AFM topographic analysis

The structural characterization of polyaniline nanofiber modified GCE and the modified surface adsorbed with ibuprofen was performed by atomic force microscopy (AFM). Nano Surf Easyscan 2AFM microscope operated in tapping mode under ambient conditions was employed. TopAZ08 probes with a spring constant of 20–80 N/m were used.

RESULTS AND DISCUSSION

Characterization of electrode surfaces

The electrode surfaces were characterized by AFM. AFM studies were conducted to give insight into the surface topography of the polymer surface and compound adsorption on the modified electrode surface. Fig. 1 shows the cyclic voltammogram of polymer modified electrode, compound adsorbed surface, particle distribution graphs, and surface roughness values.

Electrochemical behavior of ibuprofen

The cyclic voltammogram of 200 ppm ibuprofen in pH 1.0 on polymer film modified electrode exhibits a single well defined anodic peak in the potential range −0.5 to 1.8V. This anodic peak is assigned to the oxidation of ibuprofen at 1.63 V which is not accompanied by corresponding cathodic reduction. This behavior suggested that the irreversibility of the electrode process. Fig. 2 exhibits the cyclic voltammogram of polymer modified electrode (curve a). 200 ppm ibuprofen on polymer film modified (curve b) in pH 1.0. The polymer film modified surface shows a well defined and a sharp anodic peak is obtained.

The peak current values were plotted against the scan rate in fig. 3. Peak current values increased non-linearly with an increase in scan rate. Logarithmic valid peak currents were correlated well with the logarithmic values of scan rate, and it resulted in a linear line.

These facts reveal that the voltammetric oxidation of ibuprofen is irreversible and controlled by a diffusion process.

Differential pulse voltammetry (DPV)

DPV is one of the most useful and convenient methods for sample identification. By employing this technique parts per billion (ppb), the range of a compound can be determined. The voltammetric behavior of drug allowed employing stripping analysis using polyaniline nanofiber modified GC electrode.

Cyclic voltammetric results revealed the diffusion-controlled of the substrate on polyaniline nanofiber modified GCE at aqueous ethanol medium pH 1.0. It was expected that ibuprofen would be adsorbed on the electrode during the accumulation step and stripped off easily in the stripping step. Thus the stripping voltammetric studies of ibuprofen were done at polyaniline nanofiber modified GCE in medium ethanol pH 1.0 using differential pulse stripping voltammetric method. The drug exhibits a very good stripping signal. A systematic study of various instrumental parameters that affect the stripping response has been carried out with 250 ppb concentration of drug to establish the optimum conditions.

Effect of accumulation potential (Eacc) and effect of accumulation time (Tacc)

Hence optimization of the accumulation potential was done as the first part for this study. This potential was varied from −0.5 to 1.8 V at an accumulation time of 15s. Maximum Peak current was observed at 1.4 V and it was fixed as the optimum accumulation potential.

By varying the deposition time from 15 to 75s, the effect of deposition time was studied after fixing the deposition potential at 1.4 V. This table reveals highest current is observed at 15s deposition time. Hence this is taken for the further consideration.

Effect of initial potential (E0)

The initial scanning potential is another important parameter as it confirms the non-faradaic nature of the pre-concentration step. If also controls both the peak potential and peak current in the stripping voltammogram. The influence of the initial potential on the peak current was studied by varying the initial scan potential from −0.8 to 1.5 V. The peak current is affected by this initial potential in a different way. Better response ie, high peak current with the better resolution was observed at −0.5V. Hence−0.5V was fixed as initial scanning potential and the reproducibility of the method was determined by making successive measurements.

Effect of pulse height (PH) and effect of pulse width (PW)

Effect of pulse height was studied by varying from 25 to 150mV. The sharp peak current was observed at the pulse height of 100mV. Similar to pulse height, pulse width also studied by varying from 25 to 150 ms. The sharp peak nature with higher current was observed at a pulse width of 100 ms. Hence the pulse width of 100 ms was chosen as an optimum value.

Effect of scan increment (SI) and effect of pulse period (PP)

The scan increment was varied between 4 to 16 mV and maximum peak current response was obtained at 8mV scan increment. A study on the effect of pulse period in the range between 2 and 10 s was carried out. The stripping voltammetric signal showed the maximum peak current at 6 s. The optimum conditions that resulted in good peak response were used to study the effect of analyte concentration. The range of study and the optimum values arrive at table 1.

Analytical characteristic

Under optimum experimental conditions, the influence of concentration on the stripping signal was studied. The experimental results showed that the peak current increased with the increase in the concentration of ibuprofen. A representative differential stripping voltammogram is given in fig. 4. A calibration was made, which indicated the linear dependence of peak current with concentration fig. 5. The range of determination was found in between 200 ppb and 400 ppb. The lower limit of detection was 100 ppb. The reproducibility of the stripping signal was understood from relative standard deviation (2.1%) calculated for 7 identical measurements at a concentration level of 250 ppb.

The pharmaceutical samples were collected from medical shops and determined through DPV under optimum experimental conditions. Various tablets having ibuprofen were analyzed to detection of the content of drug. Stripping voltammograms of the drugs at pH 1.0 were recorded under optimized conditions. The concentration of the drug in commercial formulations determined by the proposed method was in good agreement with the reported value of the company (table 2).

Table 1: Optimum experimental conditions in DPVS

| Parameters                        | Range studied | Optimum value |
|-----------------------------------|---------------|---------------|
| pH                                | 1-13          | 1.0           |
| Accumulation potential (V)        | 0.8 to 1.8    | 1.4           |
| Accumulation time (Sec)           | 10-60         | 15            |
| Initial scan potential (V)        | -0.5 to 1.5   | -0.5          |
| Pulse Height (PH) (mV)            | 25 to 150     | 100           |
| Pulse width (PW) mSec             | 25 to 150     | 100           |
| Scan Increment (SI) mV            | 2 to 20       | 8             |
| Scan rate (SR) mV/sec             | 10 to 100     | 50            |
| Stirring rate (rpm)               | 50 to 250     | 250           |
| Rest period (Sec)                 | 2 to 10       | 5             |
Table 2: Amount of IBP presented in tablets determined by DPSV

| Brand name  | Company name            | Tablets (mg) | Experimental value (mg) | % of RSD |
|-------------|-------------------------|--------------|-------------------------|----------|
| Ibuorifen   | Modern laboratories     | 400          | 391                     | 1.7      |
| Brufen      | Abbott India Ltd        | 200          | 197                     | 2.7      |
| Ibuorifen 200 | 200Modern laboratories | 200          | 196                     | 1.7      |
| Ibuorifen   | Synmedic laboratories   | 200          | 195                     | 2.5      |
| Ibugecic    | Cipla                   | 200          | 193                     | 2.6      |

Fig. 1: AFM photographs of (a) polyaniline nanofiber modified GCE (b) Ibuprofen accumulated on electrode surface 2D, 3D, size distribution graph and surface roughness data

Fig. 2: Cyclic voltammetric behavior of (a) modified Glassy carbon electrode and (b) 200 ppm of ibuprofen on polymer modified glassy carbon electrode in aqueous alcoholic pH 1.0 at 100 mV/s

Fig. 3: Plot of peak current vs scan rate

Fig. 4: Differential pulse stripping voltammetric behavior of (a) polymer coated surface (b) 250 ppb ibuprofen on modified GCE under optimum condition

Fig. 5: Calibration plot of peak current vs conc
CONCLUSION

The AFM topography shows the polymer film is uniformly coated on the electrode surface and forms a nano fibrous morphology. The compound adsorbed surface exhibits similar structural topography. The anodic peak was observed at 1.63 V, assigned for the oxidation of ibuprofen, which is not accompanied by corresponding cathodic reduction. This behavior suggested that the irreversibility of the electrode process. The electrochemical response of ibuprofen at the modified surface reveals the irreversible and diffusion electrochemical process. The range of determination was found between 200 ppb and 400 ppb, the lower limit of detection is 100 ppb through DPV.

CONFLICT OF INTERESTS

Declare none

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