Synthesis, characterisation, and the electrochemical sensing behaviour of nicotinic acid hydrazide anchored on graphene oxide (NAHGO)

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Abstract

A simple modified sensor was developed with nicotinic acid hydrazide anchored on graphene oxide (NAHGO), synthesised by ultrasonic assisted chemical route, using hydroxy benzotriazole. Structural and morphologies of NAHGO samples were characterised by Fourier-Transform Infrared spectroscopy (FT-IR), Powder X-ray diffraction (P-XRD), Raman spectroscopy, Scanning electron microscopy (SEM), X-ray photoelectron spectroscopy (XPS) and Thermogravimetric analysis (TGA). Morphology modification of the material provided the route for more electroactive surface, that influenced the electro oxidation of caffeine with increased current. We have demonstrated the electrochemical behaviour of NAHGO on glassy carbon electrode (GCE) for caffeine detection, by employing voltammetric techniques. The influence of scan rate, pH, and concentration on the peak current of caffeine was also studied. The NAHGO sensor was employed for the determination of caffeine in imol plus and energy drink. The detection limit determined was 8.7 nM, and the best value reported so far. The results show that NAHGO modified electrodes is one of the best preferences to establish new, efficient, and reliable analytical tools.

1. Introduction

Caffeine (1,3,7-trimethyl xanthine) is one of the most consumed purine alkaloid ingredients, as it is a central nervous system stimulant, fatigue alleviator, sharpener of mental function and concentration, taste refresher with unique taste, antioxidant, booster of metabolism and associated health benefits. It is commonly found in coffee beans, tea leaves, guarana berries, pharmaceutical drugs, processed foods and energy drinks. The sensible amount of caffeine in medications reduces constipation, depression, headache etc. Like all other drugs, the overdose of caffeine produces anxiety, heartburns, increase of blood pressure, seizures, bone mass loss and cardiovascular diseases. The global average consumption of caffeine is 70 mg/person/day. On the other hand, the extracts of caffeine act as a pesticide for plants and it has been considered as an indicator for water contamination in sewers. Hence, there is a need for detecting caffeine reliably at lower concentrations, especially, in common beverages, caffeinated food and drugs.

Currently, there are different analytical techniques that are available for the determination of caffeine, such as: high performance liquid chromatography (HPLC), UV spectrometry, capillary electrophoresis, spectrophotometry. Though these methods are good to detect caffeine, their complicated, lengthy, time consuming system procedures increase the complexity to monitor the analysis. It makes the requirement of highly sensitive, consistent, simple, and fast methods. In correlation with carbon materials, electrochemical method has proved its quality and efficiency for the fabrication of sensing devices successfully. Caffeine has already been electrochemically detected with boron doped diamond electrode, DNA-functionalised single walled carbon nanotubes (SWCNT), attapulgite/nafion, gold nanoparticle, gold-chitosan, multiwalled carbon nanotubes (MWCNT)/diamond like carbon films, glutathione-rGO etc. These literature demonstrate that carbon materials has pivotal role for the
detection of harmful organic molecules like caffeine due to their highly porous structure with large surface area and excellent electronic properties.

Graphene oxide (GO) has got much attention among the carbon nanomaterials owing to its high mechanical strength, thermal and electrical conductivity\textsuperscript{20}. Besides, the carboxyl, hydroxy, and epoxy groups create GO as a desirable platform for further functionalization of molecules\textsuperscript{21,22}. Other than functionalization approaches of GO, heteroatom doping also important since it modify the properties of GO extensively. Among various heteroatom doping, nitrogen doping is widely utilized to employ doping of GO. This is because of the easiness for manipulating GO to improve more active sites for fuel cell, biosensing, adsorption, supercapacitor and solar cell applications\textsuperscript{23,24}. Nitrogen doping of GO create different kinds of nitrogen such as, pyrrolic-N, pyridinic-N, and graphitic-N in the GO lattice which enrich the electronic properties of GO\textsuperscript{25,26}. It has reported that the nitrogen doping changes the spin density and charge distribution on the surface of GO, with the strong interaction between the π-system of GO and lone pair of electrons of nitrogen\textsuperscript{27}. Hence, we propose a new kind of electrochemical sensor supported on GO and nicotinic acid hydrazide (NAH) for the detection of caffeine. In this study, nicotinic acid hydrazide anchored graphene oxide (NAHGO) was synthesised via ultrasonication-assisted chemical route. The as synthesised NAHGO was analysed by different spectroscopic morphological methods and then studied the electrochemical detection of caffeine. We found that NAHGO showed catalytic activity for the electrochemical oxidation of caffeine.

### 2. Results And Discussion

#### 2.1 Characterizations of NAHGO

X-ray photoelectron spectrum of the hybrid NAHGO is done to assess more about the surface composition and chemical states of the elements present in NAHGO. The wide scan spectrum (survey) of GO and NAHGO are displayed in Fig. 1a. GO shows only the peaks of C1s and O1s, where as NAHGO exhibits the presence of carbon, nitrogen and oxygen, with major prominent peaks at 285.12, 401.23 and 533.41 eV corresponds to C1s, N1s and O1s respectively. The C1s high-resolution spectrum is de-convoluted into four components, and is shown in Fig. 1b. The peaks at 284.5, 285.4, 286.1 and 287.6 eV attributed to C-C/C=C of graphene oxide framework, C-N, C-O and HN-C=O bonds. The deconvoluted peaks of O1s at 531.5 and 532.9 eV, as indicated in Fig. 1c, and are due to the characteristic C=O, C-O of hydroxyl and epoxy bonds of graphene oxide\textsuperscript{28,29}. The individual peaks of N1s at 399.6 and 400.8 eV, which are exhibited in Fig. 1d, attribute to the presence of pyridinic N and pyrrolic N, respectively. The XRD patterns of GO and NAHGO is presented in Fig. 1e. The intense 2θ peak at 10.7° (001) diffraction plane of as prepared GO shows an inter-planar distance 0.83 nm, calculated by the Bragg’s equation. In NAHGO, 10.7° of GO is disappeared and shifted to high 2θ value, shows the reduction of oxygen containing functional groups and the incorporation of nitrogen atoms onto graphene surface. The Raman spectra of GO and NAHGO display two major bands as shown in Fig. 1f. The C=C stretching vibration of sp\textsuperscript{2} hybridised carbon systems illustrate the G band at about 1592 cm\textsuperscript{-1} and D band at 1352 cm\textsuperscript{-1}.
corresponds to the distortions in the edge-centred band structure of carbon framework of GO\textsuperscript{29}. In NAHGO, the positions of these two peaks shift to 1581 cm\textsuperscript{-1} and 1341 cm\textsuperscript{-1}, confirming the reduction in hybrid GO\textsuperscript{30}. The intensity ratio $I_D/I_G$ of GO and NAHGO are 0.96 and 1.03 respectively. The increased intensity ratio of $I_D/I_G$ in NAHGO, which attributed to defects present after the functionalization of GO\textsuperscript{31}.

The IR spectra of GO and NAHGO are shown in Fig. 2a. The broad peak at 3375 cm\textsuperscript{-1} is attributed to the stretching vibration of carboxyl -OH for GO. The C=O, C-OH, and C-O of carboxylic and epoxy groups represent peaks at 1721 cm\textsuperscript{-1}, 1215 cm\textsuperscript{-1}, and 1030 cm\textsuperscript{-1}, respectively\textsuperscript{32,33}. Three new peaks were observed in the spectra of NAHGO. They are at 1625 cm\textsuperscript{-1}, correspond to the amide carbonyl stretch; 1561 cm\textsuperscript{-1}, is due to the N-H of amide band; 1445 cm\textsuperscript{-1}, the C-N stretching of amide bond\textsuperscript{34,35}. This is in accordance with XPS, support the nitrogen doping of GO by nicotinic acid hydrazide. Thermal stability of the GO and NAHGO were investigated by TGA, and are shown in Fig. 2b. The slow degradation of NAHGO with GO indicates the high stability which acquire by the reduction of GO and the presence of nitrogen atoms in NAHGO. The XPS data of GO and NAHGO also supports the same. The GO and NAHGO were examined by SEM to understand more about the morphology and microstructure of these compounds. As seen in Fig. 2c-d, GO demonstrated the extended sheet-like structures and NAHGO results the deformation of sheet GO owing to the functionalization of NAH on GO. In NAHGO, the small tube like structures are distributed among the layered sheets of GO. The incorporation of nitrogen moieties on GO delaminate the sheet structure leads to more voids with crumpled structure. This justifies the significant disorder in the graphitic structure on NAHGO.

2.2 Electrochemical activity of NAHGO

The surface area of GO and NAHGO modified working electrodes were probed using 0.001Molar $K_4Fe(CN)_6$ in 0.1 Molar KCl at scan rate of 50 mV s\textsuperscript{-1} in potential window of -0.2 to 0.8 V, and are shown in Fig. 3a. The poor electrochemical response of $K_4Fe(CN)_6$ in 0.1 M KCl at bare GCE implies slow transfer of electrons at the surface. The slight increase of redox peak currents at GCE modified NAHGO than GO possibly owing to fast electron transfer and the large electroactive surface area at NAHGO. This is effective to offer more active sites for caffeine oxidation. The scan rates of NAHGO in a mixture of 0.001Molar $K_4Fe(CN)_6$ and 0.1 Molar KCl and plots of anodic current ($I_{pa}$) and cathodic current ($I_{pc}$) vs square root of the scan rate ($u^{1/2}(mVs^{-1})^{1/2}$ (Fig S1 and S2a and b) were observed. The electrochemically active surface areas of all the GCE modified electrodes were calculated with the individual slopes of the plot using Randles Sevick equation

$$I_p = (2.69 \times 10^5) n^{3/2}D^{1/2}u^{1/2}A^* (1)$$

which relates the peak current of the active species with surface area for a reversible process\textsuperscript{36}. $I_p$ refers to the anodic peak current, $n$ is the total number of electrons transferred ($n = 1$), $A$ is the effective surface area of the electrode, $D$ is the diffusion coefficient of $K_4Fe(CN)_6$, $C^*$ is the concentration of $K_4Fe(CN)_6$ and $u$ is the scan rate. The calculated electroactive surface areas for bare GCE, GO, and NAHGO modified
electrodes are 0.078 cm$^2$, 0.126 cm$^2$ and 0.29 cm$^2$, respectively. NAHGO seemed to provide an enhanced active surface area presumably owing to the defects in GO with electron rich nitrogen moieties$^{37}$.

**2.3 Electrocatalytic characteristics of NAHGO for oxidation of caffeine**

The electrochemical activities of bare GCE, GO, and NAHGO were further investigated in the presence of caffeine. Fig. 3b shows CVs of bare GCE, GO, and NAHGO modified GCE electrodes, at pH 2 with 100 μM of caffeine, at a scan rate of 50 mV s$^{-1}$. The electrocatalytic behaviour of the NAHGO was also examined in the absence and presence of caffeine and is shown in Fig. 3c The occurrence of the caffeine oxidation peak in 1.5 V indicates the oxidation of caffeine on NAHGO modified GCE which is in accordance with the previous reported results$^{19,38}$. Caffeine has tertiary amine group with three alkyl groups and an amide group in which a carboxyl group is connected to a nitrogen atom. In addition to these groups, an imine, carboxyl, methyl, and alkene groups are also present in caffeine. The carbon-oxygen and carbon-nitrogen covalent bonds make caffeine more polar due to its molecular geometry. The distinct positive and negative parts that are created by the overall molecular dipole of caffeine have strong affinity to the ends of other oppositely charged ends of the NAHGO by intermolecular attractions. Though the three nitrogen atoms of caffeine are methylated (N1, N3, and N7) as seen in Fig. S3, caffeine serves as a hydrogen bond acceptor with the ring nitrogen atom (N9) and carbonyl oxygen atoms (O2 and O6)$^{39}$. Therefore the interaction of hydrogen is highly dominated on these positions of caffeine. The effective probes of interactions in NAHGO are epoxide, alcohol, carboxylic acid, pyrrolic, and pyridinic nitrogen moieties$^{40}$. The alcohol and carboxylic acid oxygen containing functional groups and the NH of pyrrolic nitrogen can act as both hydrogen bond donor and acceptor. In this case, they play the role of hydrogen donor and make electrostatic interactions with caffeine. Hence, the increased electrochemical response of NAHGO arise from the efficient interactions of highly polar functional groups of caffeine and NAHGO, which generate charge accumulations on highly polar functional groups turned out to create electrostatic interactions. These interactions are less in GO due to the less number of heteroatoms which lead to not as much current response of NAHGO.

**2.4 Effect of supporting electrolytes**

To understand the effect of supporting electrolyte on the cyclic voltammetric behaviour of caffeine, different electrolyte solutions of pH were used. The effect of pH on the peak currents and peak potentials of caffeine electro-oxidation (100 μM) by NAHGO electrode was studied, and are represented in Fig. 3d. As the pH changes from 3 to 8 the oxidation peak current weakens, and strongest current was obtained at pH 2. In pH 2, the electrostatic interactions are more likely to occur with caffeine as mentioned earlier. So, it was considered as the sensible pH for further experiments. It is also clear that the peak potential shifted negatively as the increase of pH value of the supporting electrolyte. The linear dependence of anodic current and anodic peak potential with pH for caffeine are shown in Fig. 3e and f respectively. The corresponding regression equations are: $I_{pa} = 0.214pH + 0.307$ (R=0.9963) and $E_{pa}=2.56-0.0491pH$ (R=0.9955). This implies that the electrochemical oxidation occurs with equal number of proton and electron transfer process$^{41}$. In the presence of acidic pH, the exchange of electrons between the π bonds
of graphene sheet and the lone pair of electrons bring the polarization effect at the electrode surface. The interactions of NAHGO with acidic protons are shown in Fig. 4.

Cyclic voltammograms of NAHGO (c) with and without caffeine, (d) at pH values of: 2, 3, 5, 6, 7, 8, in 100 μM of caffeine, at a scan rate of 50 mVs⁻¹, Calibration plot of (e) pH vs anodic current (Ipa), and (f) pH vs anodic peak potential (Epa).

2.5 Effect of scan rate

In order to obtain more knowledge about the adsorption properties of NAHGO the CVs of 100 μM caffeine on NAHGO at various scan rates (Fig. S4) were studied for the electro-oxidation process of caffeine. As displayed in Fig. 5a the anodic peak current of caffeine increases linearly with the increase of scan rate from 10 to 60 mV s⁻¹. The linear relationship of the oxidation current and the square root of the scan rates present that the electrochemical reactions occur on the proposed electrode is diffusion controlled. The oxidation peak potential (Epa) shifts to more positive potential as the gradual increase of scan rates. Laviron model was used to describe the kinetic parameters of the electrochemical oxidation of caffeine. The linear relationship of anodic oxidation potential with logarithms of scan rate is illustrated by this method. According to Laviron, the correlation between the scan rate and the oxidation peak potential is explained as follows:

\[ E_{pa} = E^0 + \frac{2.3RT}{[(1-\infty)nF]} \log \nu \]  

(2)

As shown in Fig. 5b, Epa exhibits the linear dependence with logarithms of scan rate (log \( \nu \)) and the equation for linearity is \( E_{pa} = 1.4326 + 0.0507 \log \nu \) with \( R^2 = 0.9965 \). The electrochemical transmission coefficient of an irreversible process is about 0.4-0.6 and for those processes the value is considered to be 0.5. Accordingly, the electronic transmission number (n) of electrochemical oxidation of caffeine is calculated as 3.85, that is in good agreement with the previous results. The mechanism of electrochemical oxidation of caffeine is shown in Fig. 6 as reported earlier. Substituted uric acid is formed in the first step, by the oxidation of two electrons and two protons at C-8-N-9 position, followed by formation of 4,5 diol of uric acid analogue through the immediate two electron and two proton oxidation.

2.6 Analytical performance

In the present work, the differential pulse voltammetric method was found to be a more sensitive method, as it produced better electrochemical performance indicators, than the CV technique. Therefore, in order to achieve the favourable analytical performance using differential pulse voltammetric (DPV) method, the parameters such as, pulse amplitude (from 0.05 V to 0.15 V), pulse width (from 0.005 to 0.5 s) and pulse period (0.1 to 3 s) is optimised. The best oxidation peak of caffeine was recorded as follows: pulse amplitude 0.05 V, pulse width 0.06 s and pulse period of 0.5 s with NAHGO modified electrode in 0.1 M \( \text{H}_2\text{SO}_4 \) at pH 2 in caffeine. Different concentrations from 0.01 μM to 40 μM of caffeine solutions were prepared to examine the relationship between the peak current and the concentration of caffeine. Fig. 5c
displays the linear increase of the anodic peak current (Ipa) with concentrations of caffeine. The calibration graph (Fig. 5d) for caffeine at NAHGO obeys the linear regression equation: 

\[ I_{pa}(\text{mA}) = 6.883 + 0.0989C; \quad R^2 = 0.9906. \]

The limit of detection (LOD) of caffeine is calculated to estimate this method for the determination of caffeine using the formula 3.3s/S, where s represents residual standard deviation and S is the slope of the calibration curve\(^{38}\) and the LOD for NAHGO modified electrode is found to be 8.7 nM, the best reported so far. Moreover, it is compared with other reported values for the determination of caffeine using electrochemical techniques. They are presented in Table 1 and which shows the effectiveness of the proposed NAHGO sensor.

Table 1 An overview of electrochemical methods for the determination of caffeine

| Modified electrodes       | Technique | Linear range (µM) | Detection limit (µM) | References |
|---------------------------|-----------|-------------------|----------------------|------------|
| BQ/CPE                    | SWV       | 500-8000          | 51.00                | 46         |
| Bi-CNT/CPE                | SWV       | 51.03-1026        | 0.182                | 47         |
| DNA/CNT/CPE               | SWV       | 0.5120-61.70      | 0.350                | 48         |
| MIP/CPE                   | DPV       | 0.0600-25.00      | 0.015                | 49         |
| CTAB/GR/GCE               | DPV       | 0.300-100.0       | 0.091                | 50         |
| SWCNT/CPE                 | DPV       | 0.250-100.0       | 0.120                | 51         |
| PT/TiO₂-GR/GCE            | DPV       | 25.00-200.0       | 0.500                | 52         |
| Nitrogen doped graphene   | SWV       | 0.060-50.0.0      | 0.020                | 53         |
| Poly(Alizarin Red S)/GCE  | SWV       | 0.500-250.0       | 0.060                | 54         |
| Attapulgite/Nafion/GCE    | DPV       | 1.000-4.000       | 0.040                | 16         |
| NAHGO/GCE                 | DPV       | 0.010-40.00       | 0.008                | This work  |

2.7 Determination of caffeine in energy drink and pharmaceutical sample

Finally, in order to estimate the practical application of this method, NAHGO/GCE was used to determine caffeine in imol plus and O'cean one8 energy drink. Energy drink was degasified by ultrasonication for 10 minutes before the analysis and diluted with supporting electrolyte (100 µM). The tablet was grounded with mortar to a fine powder, dissolved in supporting electrolyte (100 µM) and sonicated for 10 minutes. Subsequently, the appropriate amount of the resulting dispersion was selected, and then was centrifuged. 9 cm\(^3\) of 0.1 mol L\(^{-1}\) pH 2 solution was added to 1 cm\(^3\) of the sample solution and recorded the DPVs (Fig.5e and f). In the mean time to calculate the recovery standard solutions of caffeine is added and it is
represented in Table S1. The recovery range of 94.9-101.9 % suggests that NAHGO is adequate for practical sensing applications.

3. Experimental

3.1 Materials

Graphite, potassium permanganate, sulphuric acid (36 Molar), hydrochloric acid (5%), sodium hydroxide, hydrogen peroxide (30%), hydroxy benzotriazole, dicyclohexyl carbodiimide, nicotinic acid hydrazide, potassium chloride, potassium ferrocyanide, citric acid, sodium citrate, caffeine powder, potassium dihydrogen phosphate, dipotassium hydrogen phosphate were purchased from Sigma Aldrich. All the reactions were carried out under nitrogen atmosphere with the use of standard Schlenk technique. All the solvents and reagents were of analytical grade, and used without further purification. The O’cean one8 energy drink and imol plus were purchased from a local market and pharmaceuticals respectively. The stock solutions of caffeine (100 μM) were prepared fresh each time of experiments. Aqueous solutions of different pH of buffers were prepared, diluted using millipore water.

3.2 Synthesis of nicotinic acid hydrazide anchored graphene oxide (NAHGO)

Graphene oxide (GO) was synthesised from graphite, by modified Hummers method. The nicotinic acid hydrazide anchored graphene oxide (NAHGO) was synthesised by the dispersion of GO (0.2 g) in 20 cm³ dimethyl formamide (DMF), by ultrasonication method. To the solution, sodium hydroxide pellets (0.2 g, 5.0 mmol) were added and the resulting solution was stirred for 60 minutes at room temperature. Consequently, nicotinic acid hydrazide (0.2 g, 1.45 mmol), hydroxybenzotriazole (0.2 g, 2.81 mmol) followed by dicyclohexylcarbodiimide (0.2 g, 2.66 mmol) were added to the above reaction mixture, and stirred for 24 hours in nitrogen atmosphere. The resultant NAHGO powder collected by centrifugation was added to pure DMF and the resulting suspension was again centrifuged to remove the side products. This process was repeated twice with DMF and then with water to remove DMF to give pure NAHGO. The NAHGO prepared was dried overnight at 60 °C. The amide, amine, and pyridinic groups are present in NAHGO. The reaction of amino group of NAH with hydroxyl of carboxylic acid generates another amide bond in NAHGO. The imine can form with the amine group of NAH and the –C=O or –HC=O group of GO. The nearby carboxylic acid and hydroxyl groups can undergo esterification reaction followed by the generation of pyrrole ring. Hence we believe that the treatment of NAH in GO create more defects as a result of the reduction of GO with nitrogen doping. The suggested reactions which occur during the synthesis of NAHGO are shown in Fig. 7.

3.3 Characterisation techniques

The characteristic functional groups of different samples were recorded using an Agilent technologies FTIR spectrometer in the range 400-4000 cm⁻¹. Powder X-ray Diffraction (P-XRD) patterns were carried out
in a Brucker AXSD8 Advance using Ni-filtered Cu-Kα X-ray source (λ=1.5406 Å), with a scan speed of 2° min⁻¹. Thermogravimetric analysis (TGA) was recorded in Perkin Elmer STA 6000. The composition and disorders of anchored compounds were analyzed by X-ray photoelectron spectroscopy (Thermoscientific K-alpha surface analyser). Raman spectra of GO and NAHGO were recorded in HORIBA Jobin Yvon Lab RAM HR 800, equipped with a thermoelectrically cooled CCD detector. The morphology and size of GO and NAHGO were recorded on a JEOL Model JSM-6390LV scanning electron microscope. Cyclic voltammetric (CV) and differential pulse voltammetric (DPV) measurements were carried out on a CHI608E electrochemical work station with a three electrode system, using saturated Ag/AgCl acted as the reference electrode, Pt wire as a counter electrode and the modified glassy carbon electrode as the working electrode. The preparation of the NAHGO modified glassy carbon electrode and the different buffers used are given in supporting information.

4. Conclusion

In summary, the synthesis and properties of NAHGO for electrochemical detection of caffeine was discussed. Structure and morphologies of NAHGO were studied by IR, P-XRD, Raman, SEM, and TGA. The electrochemical oxidation of NAHGO towards the alkaloid, caffeine was analyzed. NAHGO displayed enhanced electrochemical activity towards the determination of caffeine, with improved detection limit of 8 nM. The NAHGO sensor was also employed for the determination of caffeine in Imol plus and O’cean one8 energy drink. The results of electrochemical studies exhibited that the NAHGO can be used as an efficient and reliable analytical tool for caffeine detection.

Declarations

Declaration of competing interest

The authors declare that they have no competing interest.

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**Figures**

![Chemical reactions diagram](image)

**Figure 1**

Reactions occur during the synthesis of NAHGO
Figure 1

Mechanism of the electrochemical oxidation of caffeine at NAHGO
Figure 1

(a) XPS wide scan spectra of GO and NAHGO, deconvoluted peaks of (b) C1s, (c) O1s, and (d) N1s (e) Powder XRD patterns, and (f) Raman spectra of GO and NAHGO
Figure 1
Cyclic voltammograms of the bare GCE, GO and NAHGO (a) in the presence of 0.001 Molar K4Fe(CN)6 in 0.1 Molar KCl, (b) at pH 2 with 100 μM of caffeine at a scan rate of 50 mV s-1, Cyclic voltammograms of NAHGO (c) with and without caffeine, (d) at pH values of: 2, 3, 5, 6, 7, 8, in 100 μM of caffeine, at a scan rate of 50 mVs-1, Calibration plot of (e) pH vs anodic current (Ipa), and (f) pH vs anodic peak potential (Epa).
Figure 1

(a) FTIR spectra, (b) Thermogravimetric analysis, (c) and (d) SEM images of GO and NAHGO
Figure 1

Calibration plot of (a) the square root of the scan rate ($\sqrt{\nu}$) vs anodic peak current ($I_{pa}$), and (b) logarithm of scan rate ($\log \nu$) vs anodic peak potential ($E_{pa}$), (c) Differential pulse voltammogram (DPV) of the NAHGO modified GCE electrodes at a pH value of 2 with standard additions of caffeine in the order 0.01 $\mu$M, 1.0 $\mu$M, 10 $\mu$M, 20 $\mu$M, 30 $\mu$M, 40 $\mu$M (d) Calibration plot of caffeine concentration ($\mu$M) vs anodic
peak current ($I_{pa}$), Differential pulse voltammogram of the NAHGO in (e) O’cean one8 energy drink and (f) imol plus.

Figure 1

The interactions of NAHGO at pH 2

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