Performance of graphene-zinc oxide nanocomposite coated-glassy carbon electrode in the sensitive determination of para-nitrophenol

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Graphene: zinc oxide nanocomposite (GN:ZnO NC) platform was tried for the sensitive determination of para-nitrophenol (p-NP) through the electrochemical method. ZnO nanoparticles (NPs) were synthesized by the modified wet-chemical method where in potassium hydroxide and zinc nitrate were used as precursors and starch as a stabilizing agent. A green and facile approach was applied to synthesize GN:ZnO NC in which glucose was employed as a reductant to reduce graphene-oxide to graphene in the presence of ZnO NPs. The synthesized NC was characterized using scanning and high-resolution transmission electron microscopy, energy dispersive x-ray analysis, X-ray diffraction and Raman spectroscopic techniques to examine the crystal phase, crystallinity, morphology, chemical composition and phase structure. GN:ZnO NC layer deposited over the glassy carbon electrode (GCE) was initially probed for its electrochemical performance using the standard 1 mM K3[Fe(CN)6] model complex. GN:ZnO NC modified GCE was monitored based on p-NP concentration. An enhanced current response was observed in 0.1 M phosphate buffer of pH 6.8 for the determination of p-NP in a linear working range of 0.09 × 10⁻⁶ to 21.80 × 10⁻⁶ M with a lower detection limit of 8.8 × 10⁻⁹ M employing square wave adsorptive stripping voltammetric technique at a deposition-potential and deposition-time of −1.0 V and 300 s, respectively. This electrochemical sensor displayed very high specificity for p-NP with no observed interference from some other possible interfering substances such as 2,4-di-NP, ortho-NP, and meta-NP. The developed strategy was useful for sensitive detection of p-NP quantity in canals/rivers and ground H2O samples with good recoveries.

Para-nitrophenol (p-NP) is an important phenolic compound in the manufacturing industry. It is a starting material in the synthesis of pesticides, dyes, drugs, explosives and is also used in leather darkening1–3. p-NP is also regarded as one of the most priority poisonous toxins because of its toxicity and persistence4. Headache, drowsiness, nausea, and cyanosis are the common symptoms of p-NP inhalation6. It is also a carcinogen and mutagen7. This harmful compound is also present in fresh and manufacturing unused water where it reaches from farming field run-off produced during the organophosphorus compounds/pesticides degradation4,7,8. Therefore, the development of a precise, cheap, and sensitive method for the determination of p-NP in terms of clinical, environmental, and food safety is important. Many analytical methods, like gas and liquid chromatography9–12, spectrophotometry13, fluorescence14, capillary electrophoresis15 are found in the literature for the determination of p-NP.

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of p-NP. Each process has certain confinements associated with it, such as the heavy instrumentation required, use of costly gases, sample pretreatment, limit of detection, sensitivity and selectivity.

Voltammetric methods have received considerable attention due to their potential benefits like simplicity, cost-effectiveness, portable, exceptional selectivity and sensitivity, cheap instrumentation, ease of miniaturization, and capability in minimum quantity monitoring\(^{16-21}\). The electrochemical technique is a noticeable quick, economical, easy operable, extremely sensitive, and most prominently on-site measurement approach to detect wastes like p-NP that needs mild-reaction conditions and has a very low detection limit. These aspects make it a better substitute technique compared to other non-electrochemical approaches. Because of these factors, several electrochemical sensors were reported for the detection of p-NP. Moreover, the aromatic ring of p-NP contains an easily reducible nitro-group which facilitates determining p-NP easily using some efficient electrochemical method.

Different strategies have been established for the qualitative and quantitative analysis of p-NP employing voltammetric techniques. These include BiO\(_3\) NPs Decorated Carbon Nanotube\(^{25}\), carbon nanotubes\(^{24}\)-silver nanoparticles, nano-gold\(^{23}\), crown ether/silver NPs\(^{26}\), ZnO nanorods\(^{27}\), nanostructured CeO\(_2\)\(^{28,29}\), and graphene oxide\(^{30}\) for p-NP detection with satisfactory results.

Recent literature surveys reports various types of conductive materials on the surface of which different metal NPs are attached and grown-up. These include carbon nanotubes\(^{31-33}\) and conducting polymer polyaniline\(^{34}\). Among these, materials graphene has drawn more attention because of its excellent properties such as first-rate electrical conductivity, huge specific surface-area, extraordinary electron-density, high mechanical strength, and chemical steadiness. All these things make it an ideal supporting material\(^{31,35}\). Nickel-based nanocomposites and bi-metallic-gold also exhibit exceptional electrocatalytic activity towards the reduction of p-NP\(^{31,38,39}\), but their small earth richness and high cost inhibit their useful applications. In order to address these problems, an alternative approach was proposed that replaces these precious expensive metals with noble metal zinc having a good stability and cost-effectiveness.

Recent studies have also shown that nanocomposites based on metal NPs decorated graphene where graphene acts as a support to anchor functionalized nanomaterials, find a variety of applications in sensors/biosensors, catalysis, light energy conversion and in fuel cells\(^{32,33}\). This is because of the excellent electron transfer ability, good stability, and wide potential window of graphene\(^{31,43}\). Planar graphene (GN) and graphene oxide-based nano-materials have been extensively used for sensor applications on account of their outstanding charge transfer characteristics, huge specific surface area, excellent electrocatalytic activity, and low cost\(^{45}\).

Semiconducting zinc oxide (ZnO) NPs have become very popular over the past few years for their applications in optics, optoelectronics, sensors, energy and biomedical sciences because of their abundant availability, low cost, friendly to the environment, and excellent electro-catalytic activity as compared with other metal oxides\(^{46-48}\). ZnO and graphene are good electro-catalysts and the activity could be further enhanced by the coupling of two components using graphene as a support. ZnO NPs directly synthesized on the graphene surface have been reported as a good electro-catalyst material for supercapacitors\(^{41,42,49}\).

Here, we separately prepared ZnO NPs by wet chemical method and then loaded on graphene layers by dispersing it with graphene oxide (GO) for the preparation of GN-ZnO NC employing glucose as a reductant to reduce GO to GN and simultaneously loading ZnO NPs on GN layers. For fabrication of the detection platform, the freshly prepared GN-ZnO NC layer was placed over the glassy carbon electrodes (GCE) through the drop-casting deposition method\(^{34,44}\). The GN-ZnO NC was found to be very effective to electro-chemically catalyze the p-NP concentration. The synergistic effect of GN-ZnO NC improved electrochemical detection of p-NP and enabled its sensitive determination in presence of o-nitrophenol using voltammetric techniques. To the best of our knowledge, no electrochemical method employing GN-ZnO NC was reported for the determination and detection of p-NP in presence of o-NP. In addition, the method is simple and the composite synthesized is clean with no extra stabilizer.

**Experimental work**

**Materials used.** Carbon (graphite powder) was supplied from Sigma-Aldrich. Starch, zinc nitrate, potassium hydroxide, ethanol, 4-NP, meta-NP, and 2, 4-di-NP, 2-aminophenol, 4-aminophenol, and chlorophenol, potassium dihydrogen phosphate, dipotassium monohydrogen phosphate, glucose, ethanol, and nitric acid were purchased from Alfa Aesar. Double-distilled water was used throughout for solution preparations.

**Synthesis of zinc oxide nanoparticle.** The ZnO NPs powder was synthesized by the modified wet-chemical method\(^{45}\). To synthesize ZnO NPs, 2.0 g zinc nitrate was dissolved in 100 cm\(^3\) aqueous media under constant mechanical stirring on a hotplate at 60 °C for 30 min. Later on, separately 1.0% starch solution prepared in 10 cm\(^3\) dist. water was introduced into the ongoing reaction and the solution was constantly stirred to continue on the hotplate for a further 30 min. 0.74 g of KOH was also dissolved separately in 50 cm\(^3\) of double-distilled water and added drop wise touching the walls of the beaker with constant stirring. On complete addition of KOH, the reaction was further carried out for 2 h. After that, the solution was kept overnight to allow the product to get settled down and separate the supernatant. The precipitate formed was separated by centrifugation and washed three times with dist. water to eliminate unreacted reactants. Then it was dried overnight using an electric oven and the temperature was kept at 80 °C\(^{51}\).

**Synthesis of GN:ZnO NC.** Initially, graphene oxide (GO) was synthesized employing the conventional “Hummer’s method”\(^{35,44}\). For the synthesis of GN:ZnO NC, 50 mg of synthesized GO was dispersed in 100 cm\(^3\)
of deionized H2O which was exfoliated in an ultrasonic bath using ultrasonication for 15 min. Then to the exfoliated GO dispersion, 50 mg of as-prepared ZnO NPs were added. The 1:1 mixture (by weight) of GO and ZnO NPs was vigorously magnetically stirred for about 2 h for formation of homogeneous suspension. This 100 ml homogenous suspension, 1 g glucose was added followed by stirring for 30 min to reduce GO to GN escaping the use of extremely toxic hydrazine as a reductant. During this process, GO is reduced to GN, and simultaneously the deposition of ZnO NPs onto the GN sheets is achieved. The same procedure was applied for the synthesis of GN but independently and in absence of ZnO NPs to make a comparison. The blackish dispersion obtained was then filtered using a Millipore filtration system and was given washing three times with each of water and ethanol. The obtained GN and the GN:ZnO NC were dried in an oven. Complete representation for the synthesis of GN:ZnO NC is shown in Scheme 1. Finally, 5 mg of the composite as synthesized was dispersed in 2.5 cm3 of ethanol under ultra-sonication for 5 min to make a suspension for electrode fabrication.

**Fabrication of GN:ZnO NC/GCE.** A bare GCE was washed in a 1:1 nitric acid: water mixture and then polished by alumina powder to form a clean shiny surface. It was then again washed with the nitric acid–water mixture and ethanol ultrasonically and dried in air. 10 µL of the as-prepared suspension of the composite material was deposited onto the conducting part of GCE surface with the help of a micropipette. The solvent was evaporated using an IR lamp. When the solvent was removed, the testing material was adhered on the electrode surface which was again rinsed thoroughly with distilled water and dried again under an IR lamp. This fabricated electrode was designated as GN:ZnO NC/GCE and was used in successive electrochemical studies.

**Characterizations.** X-ray diffraction crystallographic studies were carried out using XRD machine (Shimadzu, Japan). S-4800 field emission SEM system was used to take scanning electron microscopic (SEM) images. It is operating at 20.0 kV and equipped with energy-dispersive X-ray (EDX) spectroscopic technique for elemental analysis. Transmission electron microscopic (TEM) and High-resolution-(HR)-TEM (JEOL JEM-2100F microscope at 200 kV with “Orius SC1000” camera) techniques were used to further analyze the microstructure of the materials. Initially the samples were dispersed in dimethyl formamide and then bath-sonicated for 30 min at room temperature. The samples were drop-casted on a carbon-grid and allowed to dry. When the solvent got removed by evaporation, the sample got adhered to the grid. Raman spectra were obtained using (RE-04) Raman spectrometer having a solid state laser diode-pumping at 514 nm.

All the voltammetric and Chronocoulometric experiments were performed using an Electrochemical-workstation (CH-Inst. Model-CHI1100B) with a three-electrode cell. The bare-glassy carbon electrode (GCE, 3 mm diameter) was modified to act as working electrode. Platinum wire and Ag/AgCl were auxiliary and reference.
electrodes, respectively. Electrochemical-impedance experiments were carried out in 0.1 M KCl soln. along with a mixture of 1.0 mM [Fe(CN)6]−3 and [Fe(CN)6]−4 (1:1) was taken as standard probe employing Electrochemical-workstation (Autolab PGSTAT 30 with Frequency Response Analyzer). The pH readings were taken using ELICO LI-120 digital pH meter.

Results and discussion
The XRD plots of the synthesized ZnO nanopowder, GO and GN:ZnO NC composite are given in Fig. 1. XRD analysis of ZnO NPs depicts diffraction peaks at 31.80°, 34.4°, 36.3°, 47.5°, 56.6°, 62.8° and 69.0° in the (Fig. 1A), confirming the hexagonal wurtzite structure of ZnO NPs41,52,53 (JPCDS number: 36-1451)52. Also, the presence of quite sharp peaks and the absence of any other peak indicate that the synthesized nanopowder does not contain other impurities. The size of ZnO-NPs was also calculated with the help of “Debye–Scherrer formula, $d = \frac{0.89\lambda}{\beta\cos\theta}$”54,55, where, 0.89 is Scherer’s factor, $\lambda$ is wavelength of x-ray used, $\theta$ the “Bragg diffraction angle”, and $\beta$ a constant known as the “full width at half-maximum (FWHM)” of a particular diffraction angle. The mean particle size of the ZnO-NPs corresponding to the FWHM of the most intense peak at 36.3° and corresponding to (101) plane using the “Scherrer formula”56,57 was found to be 57.17 nm. The diffraction peak at a 2θ value of 10.81° characteristics of GO (Fig. 1C) disappears in the XRD pattern of as-prepared GN:ZnO NC composite (Fig. 1D). However, a new diffraction peak characteristic of GN at 26.52° with inter-layer spacing (d-spacing) of 0.35 nm was observed in the XRD outline of GN:ZnO NC composite. This confirms that the exfoliated GO has been reduced to GN. The other XRD peaks of the GN:ZnO NC composite can be ascribed to the crystallize ZnO NPs. Further, the XRD peak in case of GN:ZnO NC composite corresponding to GN shifted slightly as compared to individual GN at 26.20° (Fig. 1B) which again confirms GN-ZnO NPs interaction. The ZnO NPs and GN:ZnO NC composites were further analyzed using SEM and EDX techniques. Figure 2A shows the SEM image of as-synthesized ZnO-NPs. As observed, the approximate spherical-shaped ZnO NPs were formed in the range of 40–90 nm size which agrees with the particle size calculated using the “Debye–Scherrer formula” and the particle size analyzer which agrees with the results as obtained in SEM images. HR-TEM image of ZnO NPs from a single nanoparticle is shown in Fig. 3B. The selected area electron-diffraction (SAED) outline so obtained is given in Fig. 3C and the rotational average profile of ZnO NPs is given in Fig. 3D. The lattice spacing values obtained from the SAED outline and the standard “JCPDS” values are given in Table S1. The lattice spacing calculated from the selected area is 2.48 Å which matches perfectly with the value obtained from the standard JCPDS card (2.47390 Å) corresponding to 101 planes of hexagonal ZnO. The values given in Table S1 agree very well within the error limit of ± 2.5% confirming the hexagonal phase of the ZnONPs. Figure 3D represents the rotational average profile of ZnO-NPs. Figure 4A represents the TEM image and Fig. 4B shows the HRTEM image of as-synthesized GN and Fig. 4C shows its corresponding Fast Fourier Transform (FFT) pattern. The hexa-gonal FFT outline shows that the as-synthesized GN is highly crystalline in

Figure 1. X-ray diffraction pattern depicting peak-indices and 2θ values of (A) ZnO NPs, (B) GN (C) GO and (D) GN:ZnO NC.
nature with minimum defects. TEM image of GN:ZnO NC is shown in Fig. 4D which shows that the ZnO-NPs are well dispersed in the graphene sheets.

These results show that there is a good interfacial contact of GN and ZnO-NPs. The existence of ZnO-NPs within the GN layers will help to prevent the agglomeration as well as restacking of plane GN layers and simultaneously its available surface-area gets also increased to improve its electrochemical performance. The GN:ZnO NC was also characterized by EDAX technique. Figure 5(1) shows the EDAX spectrum of GN:ZnO NC with peaks characteristic of C, O, and Zn elements. The respective weight percentages using EDAX studies of C, O, and Zn are 66.94, 10.85, and 22.21. This reveals a good loading of ZnO-NPs on graphene nanosheets, which agrees with the SEM and TEM interpretations. In this way, a good dispersion of ZnO NPs with GN is achieved.

Raman spectroscopy is a very sensitive technique for electronic structure and is an essential tool to examine well-ordered and disordered crystalline structures and is widely used to characterize carbonaceous materials like graphene48. Raman spectra of the GO, GN, and GN:ZnO-NC are presented in Fig. 5(2). The two characteristic bands in the Raman spectrum of GN and graphene-based materials are low intensity ‘D-band’ and high intensity ‘G-band’ at 1351 and 1602 cm\(^{-1}\), corresponding to the disordered carbon and in-plane stretching vibration of sp\(^2\) C – C bonds, respectively\(^{44,59}\). The \(I_D/I_G\) intensity ratio gives the degree of disorderness and the mean size of the sp\(^2\)-domains in graphitic materials\(^{44,59,60}\). Figures 5(2A,B) shows that \(I_D/I_G\) intensity ratio is increased for GN as compared to GO which indicates that there is a decrease in the in-plane sp\(^2\)-domain size on graphene-oxide reduction\(^{59}\). The \(I_D/I_G\) value gets increased from 0.54 in GO to 1.13 in GN:ZnO NC (Fig. 5(2C). This is because on reduction of GO to GN, oxygen functionalities are removed, which decreases the size of the sp\(^2\)-domains in graphene and interaction between ZnO NPs and graphene nanosheets\(^{44,63,64}\).

During the reduction of GO to GN, most oxygen based functionality are removed and restores the conductivity of graphene, and also increases the stability of GN:ZnO NC\(^{42}\). However, some residual oxygen-containing groups are still left unreduced on GN\(^{41}\). The Zn atoms present in ZnO NPs coordinate with the oxygen atoms of the unreduced functional groups via covalent-interaction which helps in composite formation\(^{42}\). Graphene bearing edges and planes will provide support to spherical ZnO NPs to grow and form GN:ZnO NC.

**Electrochemical characterization of GN:ZnO NC/GCE.** Electrochemical impedance spectroscopic (EIS) and chrono-coulometric (CC) techniques were applied to characterize the GN:ZnO NC modified GCE. EIS is a suitable technique commonly used to characterize a surface modified GCE\(^{55}\). In the Nyquist diagrams, the semicircular-region obtained at high frequency values correspond to the charge transfer limiting phenomena and from the diameter of the semicircular part, charge transfer values (\(R_t\)) are calculated\(^{63}\). EIS analysis shows that the \(R_t\) value for GN:ZnO NC/GCE (Fig. 6(1A)) is 3.6 \(\Omega\) which is lower than that of GN/GCE (6.0
Ω, Fig. 6(1B)). The decrease in Rct value confirms that the ZnO-NPs present in the composite improves the charge transfer characteristics of GN:ZnO NC modified GCE. CC was applied to determine the electrochemically active surface area of the GN and GN:ZnO NC modified electrodes by using Anson charge-time dependence equation \(Q = nFACD^{1/2} \pi^{-1/2} t^{1/2}\), where 'Q' is charge in coulombs, 'n' number of electrons involved in redox reaction is 1 for \(K_3[Fe(CN)_6]\) system, 'F' Faraday Constant (96,485 Cmol\(^{-1}\)), 'A' area of electrode in cm\(^2\), 'C' concentration of \(K_3[Fe(CN)_6]\) (1 mM), 'D' diffusion coefficient is \(7.6 \times 10^{-6}\) cm\(^2\)s\(^{-1}\) for \(K_3[Fe(CN)_6]\) and 't' is time in seconds [30]. The slope obtained from the plot of Q against t\(^{1/2}\) using CC and 1 mM \(K_3[Fe(CN)_6]\) standard complex was used to calculate the effective electrochemical surface areas for GN and GN:ZnO NC modified electrodes (Fig. S1A,B, supplementary material). Based on the linear relationship between Q and t\(^{1/2}\), the slopes obtained for GN:ZnO NC/GCE and GN/GCE were found to be 138.40 Cs\(^{-1/2}\) and 76.44 Cs\(^{-1/2}\), respectively (Fig. S1A,B). The effective electrochemical surface areas for GN:ZnO NC/GCE and GN/GCE were found to be 0.166 cm\(^2\) and 0.092 cm\(^2\), respectively.

Electrochemical performance of GN:ZnO NC modified GCE for p-NP determination. Cyclic voltammetric (CV) study was carried out at a scan-rate of 100 mVs\(^{-1}\) to investigate the electrochemical behavior of 1.96 \times 10^{-6} M p-NP at bare GCE, GN/GCE, and GN:ZnO NC/GCE. Figure 6 (2a) is the CV of the blank containing phosphate buffer at pH 6.8 at bare GCE. However, using bare-GCE at the same scan-rate and concentration of 1.96 \times 10^{-6} M, p-NP gives a weak redox couple at 0.20 V ascribed to p-NP (Fig. 2b), while the peak current progressively increased on GN modified GCE (Fig. 2c), and further enhanced with GN:ZnO NC modified GCE (Fig. 2d). Also, the irreversible reduction peak of p-NP at ~0.75 V grows progressively at both GN and GN:ZnO NC modified GCE. This shows that both ZnO NPs and GN had an electrocatalytic effect where ZnO NPs act as spacer to enhance the effective-surface area of GN and improve the electrochemical activity while GN decreases the resistance by providing a conducting path. Therefore, the synergic effect of ZnO NPs and GN as cocatalysts
is responsible for the electrocatalytic activity of GN:ZnO NC/GCE and its application in the electro-analysis of para-nitrophenol. This is due to the excellent catalytic capability and large effective-surface area of GN:ZnO NC/GCE which facilitates p-NP to accommodate at the electrode-surface and enhances the electron-transfer process. The current intensity increases progressively on GCE modified with GN and GN:ZnO NC. Figure 6(2c) shows that one large reduction peak appears at −0.75 V on the first cathodic sweep for the p-NP and another redox couple with one anodic peak at 0.19 V and another cathodic-peak at 0.11 V is obtained. It is observed from the cyclic voltammograms (Fig. 6) that this redox couple grows in peak current at the expense of first irreversible reduction-peak at 0.75 V. This indicates that the irreversible reduction product of p-NP stays on the modified-electrode surface and gets oxidized in the anodic sweep. This is also in agreement with the literature employing the conventional electrodes. The first irreversible reduction-peak obtained is ascribed to the irreversible reduction of the nitroaromatic moiety to yield the hydroxylamine derivative (NHOH) with the four-electron transfer, according to Scheme 2. The anodic redox couple is due to the redox reaction of hydroxylamine and p-NP involving two electrons in the reaction. Figure 6(3) shows the scan rate effect on peak current using CV for p-nitrophenol (1.96 × 10−6 M) in phosphate-buffer of pH 6.8 at 50, 100, 200, 300, 400, 500, 600, 700, 800 mVs−1 at GN:ZnO NC/GCE. The plot of scan rate against the cathodic and anodic-peak current (Fig. 6(4)) of reduction product of p-NP (hydroxylamine) gives straight lines which indicate that both the electrode processes are adsorption controlled. Also, when reduction peak-current of p-nitrophenol was plotted against scan-rate, a straight line was obtained indicating an adsorption-controlled process.

**Experimental parameters and their optimization for p-nitrophenol determination.** Effect of pH values. Nature as well as the pH of supporting electrolytes play a vital role in the determination of analytes using any chemically modified electrode. Many supporting electrolytes were analyzed. While choosing a specific supporting electrolyte for p-NP determination, peak current as well as peak shape were taken into consideration. 0.1 M each of borate, acetate, phosphate, and Britton Robinson buffer were tested. In citrate buffers of pH 2.1 and KH2PO4–K2HPO4 buffer solution of pH 6.8, high-intensity anodic peak currents were observed. However, in the citrate buffer of pH 2, the anodic peak was unstable. Therefore, for subsequent studies, the optimum buffer solution chosen was phosphate buffer (pH 6.8). Differential pulse voltammetry, square-wave voltammetry, and cyclic voltammetric techniques were applied to optimize the pH values and it was found that square-wave voltammetry responded the best in respect of peak-current and peak-resolution. Square wave adsorptive stripping voltamograms of p-NP (1.66 × 10−6 M) at GN:ZnO NC/GCE and various pH values of 0.1 M potassium phosphate buffer are presented in Fig. 7A. It is observed that the peak current goes on increasing with the increase in pH values up to 6.8 but at 7.4 and 8.0 pH values it decreases. It confirms that the redox reaction of p-NP is favored in the acidic medium. Therefore, for further studies phosphate-buffer of pH 6.8 was selected as the supporting-electrolyte. Also, it is observed from Fig. 7A that the peak-potential is shifting towards a more negative side as...
the pH increase which confirms the proton involvement in the reaction. A plot of pH against shift in peak potential (Fig. 7B) gives a straight line with a slope equal to 61.3 mV/pH which confirms that protons and electrons are equally taking part in the electrode reaction as shown in Scheme 2.

Optimization of deposition potential and deposition time. Deposition potential and deposition time, both affect the degree of adsorption of the electrochemically reduced p-NP at the electrode surface (NH–OH). Therefore, both these parameters need to be optimized. The effect of deposition-potential and deposition-time on peak current was investigated using SW-AdSV. It was found that the deposition-potential largely effects the peak current of para-NP at the modified electrode and the peak current enhanced from − 0.1 to − 1.0 V and then diminished. Hence a deposition-potential of − 1.0 V was optimized for subsequent studies. Figure S2 (Supplementary material) presents the effect of deposition-time on the anodic peak current of NH–OH for 10.70 × 10−7 M p-NP at GN:ZnO NC/GCE under optimized conditions (deposition potential: − 1 V; potassium phosphate buffer of pH 6.8 as supporting electrolyte) for various deposition times (60, 120, 200, 300, 400 and 500 s). The peak current increases with the increase in deposition-time from 60 to 300 s and then remain almost constant. This is because the modified electrode surface gets saturated due to the accumulation of reduction product (NH–OH). In this way, sensitivity can be improved by increasing the deposition time to determine low concentration levels. A deposition time of 300 s was optimized for subsequent experiments.

Calibration curve. Figure 8A shows the calibration curve for the fabricated sensor using the optimized experimental parameters. The calibration curve was used for the quantitative analysis of p-NP solutions using GN:ZnO NC/GCE employing SW-AdSV using the as-optimized experimental parameters. The stripping-peak current gave a linear relation by changing the concentrations of p-NP from 0.09 × 10−6 M to 21.80 × 10−6. Using the linear plot of the peak current (i_{ap}) versus concentration of p-NP (Fig. 8B) the regression equation obtained is: i_{ap} (µA) = 68.14 C(µM) + 10.67 (r² = 0.996). This equation was used to calculate the limit of detection (LOD) and sensitivity of the developed sensor according to IUPAC recommendations:

LOD = 3σ/k

where σ is the standard deviation of intercept for replicate determination values under the similar conditions (n = 5); k is the sensitivity, namely the slope of the calibration graph. The limit of detection (LOD) and sensitivity of the developed sensor were estimated to be 8.8 × 10−9 M (S/N = 3) and 68.14 µA µM−1, respectively. This very low detection limit of the sensor shows that the developed method can potentially be applied for the sensitive determination of p-NP. A comparative analysis for the performance of the developed sensor with different electrochemical sensors based on different nanomaterials for the sensitive determination of p-nitrophenol is summarized in Table 1.

It is obvious from Table 1 that the sensor based on the present nanocomposite material is superior in respect of its wide linear working range and more important its limit of detection is least compared to those reported by the earlier workers. It can be concluded that the GN:ZnO NC-based electrode is comparatively an first-rate platform for the sensitive determination of p-NP. This improved performance of the electrochemical sensor is attributed to the synergistic effect of GN and ZnONPs based composite material due to their high adsorption ability and excellent electrocatalytic activity.
Figure 6. (1) Nyquist plots for (A) GN:ZnO NC modified GCE and (B) GN modified GCE in 0.1 M KCl solution in presence of 1.0 mM each of Fe[(CN)]_6^-3 and Fe[(CN)]_6^-4 (1:1). Insert: Equivalent circuit for fitting the Nyquist plots. (2) CV of (a) phosphate-buffer at pH 6.8 (blank) at bare GCE and 1.96 × 10^-6 M p-NP in phosphate buffer at pH 6.8. (b) Bare GCE, (c) GN:GCE and (d) GN:ZnO NC modified GCE, Scan rate100 mV s^-1. (A) Effect of scan-rate on peak current of p-NP (1.96 × 10^-6 M) using cyclic voltammetry at 50, 100, 200, 300, 400, 500, 600, 700, 800 mV s^-1 using GN:ZnO NC/GCE. (B) Plots of peak current (Ipa, Ipc) vs. potential scan rate (v).

Scheme 2. Electrode reaction mechanism of p-nitrophenol at GN:ZnO NC/GCE.
Effect of potentially interfering substances. Selectivity of the prepared GN:ZnO NC-based sensor for the analysis of p-nitrophenol was evaluated by studying the influence of some important interfering substances of p-NP. The interference study was examined in 0.1 M phosphate-buffer of pH 6.8 in presence of $1.07 \times 10^{-6}$ M p-NP. Most of the phenols like pyrocatechol, o-aminophenol, hydroquinone, hydroxyphenyl, p-aminophenol, and chlorophenol did not affect the signal of p-NP with deviations less than 4.0% up to the 100-fold excess. The common nitrophenols that contain the same nitro group as that of p-nitrophenol are o-nitrophenol, m-nitrophenol, and 2, 4, di-nitrophenol. The interference of these nitrophenols was tested on the GN:ZnO NC modified GCE. It was found that o-NP produced a separate reduction peak at $-0.10$ V at the same concentration and suppressed the signal of p-NP at higher concentrations as shown in Fig. S3 (Suplementary material). Therefore, it can be concluded that both p-nitrophenol and o-nitrophenol can be detected simultaneously at the said modified GCE at a lower concentration. However, the presence of m-NP and 2, 4, DNP did not affect the determination of p-nitrophenol. The results obtained from interference tests indicate that GN:ZnO NC-based sensor will be appropriate for the analytical determination of p-NP in presence of the above-mentioned phenolic compounds and meta and ortho-nitrophenol.

The fabricated electrode was also evaluated for its reproducibility and stability. The reproducibility for the detection of $1.66 \times 10^{-6}$ M p-NP using 0.1 M phosphate-buffer of pH 6.8 employing five electrodes was found to have a relative-standard-deviation (RSD) value of 4.70%, revealing a good reproducibility of the sensor. The stability of the sensor was also examined in 0.1 M phosphate-buffer of pH 6.8 for $1.66 \times 10^{-6}$ M p-nitrophenol at the current response obtained was periodically monitored. An initial peak current response of about 93.40% of the sensor was retained for 15 days which proved good stability of the sensor. The repeatability of the as developed sensor was also carried out for 10 successive measurements with an RSD value of 4.0% indicating good repeatability.

Accuracy and precision of the method. The accuracy of the method as developed was determined by spiking an exactly weighed amount (pre-analyzed amount) of p-NP. The accuracy is represented as a mean relative error (Table 2A). The mean percentage recovery was found to be 98.93% and 99.34% for intra-day and inter-day assays, respectively. The recovery values are useful to provide good confidence in the accuracy of the developed method.

The precision of the developed method was also determined. The concentration of p-nitrophenol was analyzed in pre-analyzed sample solutions five times in intra-day assay and successively for five days in an inter-day assay using the SW-AdSV techniques. The precision and the percentage recovery values of the developed method obtained as the mean of five separate measurements are given in Table 2A. The average values of variation-coefficients for intra-day and inter-day assays based on five measurements were found to be 1.63% and 1.94%, respectively. Thus, the results obtained confirm a good reproducibility of the fabricated electrode and a high precision of the developed method.

Practical application of the developed sensor in the determination of para-NP in real samples. The GN-ZnONPs composite based sensor was used to analyze p-NP in various real water samples. Groundwater sample was collected from Andheri (East) and river water sample from Panchganga River, Mum-
Figure 8. (A) Square wave adsorptive stripping voltammograms at GN:ZnO NC/GCE at different concentrations of p-NP using phosphate buffer (pH 6.8). Deposition-potential: −1.0 V, deposition-time: 300 s. (B) Plot of concentration of p-NP against peak current with error bars. (C) SW-AdS voltamgrams for p-NP determination using standard addition method spiked with (1) $35.61 \times 10^{-8}$ M, (2) $71.22 \times 10^{-8}$ M and $106.83 \times 10^{-8}$ M p-NP in river water sample. (D) SW-AdS voltamgrams for p-NP determination using standard addition method spiked with (1) $35.61 \times 10^{-8}$ M, (2) $71.22 \times 10^{-8}$ M and $106.83 \times 10^{-8}$ M p-NP in ground water sample.

Table 1. Comparison of the performances of different electrochemical sensors for p-nitrophenol. *Gold nanoparticle/reduced graphene oxide, **Multi-wall carbon nanotubes, ***Palladium nano-flowers supported on fullerene.
bai. 0.1 M potassium phosphate-buffer (supporting electrolyte) was directly prepared in 100 cm³ of each sample which was used as an analyte. 25 cm⁻³ of sample volume was taken in an electrochemical cell for detection of p-NP using the standard addition method. The developed method was applied under the optimized experimental parameters (deposition potential: − 1.0 V and deposition time : 300 s) and under the established concentration range. The measurements were performed three times. It was found that neither p-NP nor o-NP was detected in both these water samples. Therefore, the desired amount of standard p-nitrophenol was spiked into the collected water samples for performing the recovery tests. Figure 8C shows the SW-AdSV curves for standard p-nitrophenol spiked in river water sample and Fig. 8D shows the SW-AdSV curves for standard p-nitrophenol spiked in ground water sample under the optimized conditions. The percentage recovery of both the samples is given in Table 2B. A recovery of 97.660% and 97.130% was obtained in river and groundwater samples respectively. These observations validate the suitability of the developed method for the determination of p-nitrophenol in the natural water samples.

**Conclusion**

A novel electrochemical method employing GN:ZnONC was established for the sensitive determination of p-nitrophenol. The good dispersibility of ZnO NPs with GN demonstrated an improved electrocatalytic performance for the determination of p-nitrophenol in contrast with GN-based GCE. This can be attributed to the outstanding electrical conductivity, large adsorptive capacity, and very high effective surface area of GN and ZnO NPs. In addition, it was found that simultaneous detection of ortho and p-NPs can be achieved at the said electrode without the interference of other possible interferents at low concentrations. The developed procedure was applied to determine p-nitrophenol in the river and ground water samples using SW-AdSV with acceptable recoveries of 97.660% and 97.130% respectively. It is evident from these results that the GN:ZnO NC composite is a good candidate for advanced electrode materials for the sensitive determination of p-NP.

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![Table 2.](image)

| Sample                  | Added (10⁻⁸ M) | Found (10⁻⁸ M) | % Recovery |
|-------------------------|---------------|---------------|------------|
| Panchganga river water  | 35.61         | 34.78         | 97.66      |
|                         | 71.22         | 70.01         | 98.30      |
|                         | 106.83        | 103.99        | 97.34      |
| Ground water            | 35.61         | 34.59         | 97.13      |
|                         | 71.22         | 69.20         | 97.16      |
|                         | 106.83        | 104.31        | 97.64      |

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**Table 2.** (A) Precision and accuracy analysis for p-nitrophenol in pre-analyzed samples by the developed method (SW-AdSV), (B) Percentage recovery assay for p-nitrophenol in real water samples by the developed procedure (SW-AdSV). *Average of five measurements.*
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**Competing interests**

The authors declare no competing interests.

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