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Manganese Oxide/Hemin-Functionalized Graphene as a Platform for Peroxynitrite Sensing

Haitham Kalil, Shaimaa Maher, Tiyash Bose, and Mekki Bayacho

Recent clinical research indicates that the cytotoxicity of peroxynitrite (ONOO\(^-\), PON) plays an important role in several pathologies including cardiovascular dysfunctions and other diseases triggered by oxidative stress.\(^1\) Peroxynitrite is a potent oxidative and/or nitrosative agent produced from the diffusion-controlled reaction between nitric oxide radical (NO) and superoxide anion-radical (O\(_2\)\(^-\)). It is a well-known member of the reactive oxygen-nitrogen species family.\(^6\)\(^,\)\(^10\) This non-radical metabolite attacks vital components in cells and initiates deleterious effects via direct and indirect interactions. Peroxynitrite reacts directly with lipids, DNA, and proteins, and indirectly triggers several radical chain reactions.\(^9\)\(^,\)\(^11\)

The elevation of peroxynitrite levels is associated with chronic inflammatory disorders including neurological and vascular diseases, as well as many other pathophysiological conditions.\(^3\)\(^,\)\(^4\)\(^,\)\(^10\)\(^,\)\(^12\)\(^,\)\(^13\) Hence, the precise quantification of this analyte in biological systems is of paramount importance not only to understand the genesis and development of diseases at the tissue/cellular level, but also to assess potential therapies and understand its associated reaction mechanisms. Although some progress has been achieved in terms of peroxynitrite detection and quantification, the techniques of detection are still relatively limited. This deficiency is due to complicated pretreatment, low selectivity, as well as other technical problems.\(^10\)\(^,\)\(^16\)

Fluorescence, chemiluminescence, electron spin resonance, and immunohistochemistry methods are indirect since they rely on the measurement of secondary species derived from oxidation or nitration reactions mediated by peroxynitrite.\(^17\)\(^,\)\(^19\) These indirect methods are of course costly, relatively laborious, and generally suffer from poor sensitivity and low selectivity. Thus, developing sensitive, selective, fast-responding, and direct detection methods is highly desirable.

Graphene nanosheets have become very attractive as an interface material for electrochemical sensing due to their unique physical and chemical properties. Graphene is a single atomic layer of graphite arranged in a two-dimensional structure.\(^20\)\(^-\)\(^22\) Its sp\(^2\) hybridization coupled with its ultra-thin atomic thickness accounts for graphene’s unique properties and wide applications.\(^23\)\(^-\)\(^35\) Graphene can combine with substrates by either π–π stacking or cation–π interaction through catalytic metallic centers.\(^33\)\(^,\)\(^36\) Graphene modification, whether by metal decoration or nonmetal functionality, opens possibilities for its use in catalysis and sensor applications.

Electrochemical detection of analytes such as peroxynitrite at an interface generally offers real-time measurements using simple methods with minimum sample preparation. Metalloporphyrins-modified glassy carbon electrodes (GCE) have shown significant electrocatalytic properties toward peroxynitrite detection, particularly hemin-modified GCE. Manganese phthalocyanine electrodeposited on platinum microelectrodes, also showed a significant increase in electrode sensitivity toward peroxynitrite. Both metalloporphyrins and phthalocyanine compounds have been used as electrochemical interfaces for peroxynitrite detection and were used as such or in a combination with graphene nanostructures.

Several electrochemical sensors based on manganese phthalocyanine, cyanocobalamin, and hemin have been developed to detect peroxynitrite.\(^17\)\(^,\)\(^40\)\(^,\)\(^63\) Amatore and co-workers used a platinized carbon microelectrode under physiological conditions.\(^44\) They were able to detect the signature of one-electron oxidation of peroxynitrite released under oxidative stress at the level of a living cell.\(^44\) Bedioui’s group used a platinum ultramicroelectrode electrochemically modified with manganese tetraamino-phthalocyanine films for the detection of peroxynitrite in alkaline solutions (pH = 10.2).\(^43\)\(^,\)\(^45\) The reported sensitivity of the modified sensor was 14.6 nA/μM with a limit of detection of 5 μM. On the other hand, Wang and Chen showed that electropolymerized cyanocobalamin films on glassy carbon can be used for the electrochemical detection of peroxynitrite at pH of 9.2.\(^17\)

Apparently, the polymerized cyanocobalamin layer catalyzes the electrochemical oxidation of peroxynitrite at the surface of the electrode.\(^17\) Our group used electropolymerized 3,4-ethylenedioxythiophene (PEDOT)-hemin film on carbon fiber electrodes to detect and measure PON.\(^69\) Later, hemin-reduced graphene oxide material was used at an electrode interface for peroxynitrite detection.\(^69\) In this work, we examine the electrocatalytic activity of hemin-reduced graphene oxide material that is further modified with manganese oxide nanoparticles toward PON catalytic detection. Graphene oxide is used as a scaffold and is modified with needle-like MnO\(_2\) nanostructures followed by hemin functionalization as reported before.\(^18\)
The graphene-hemin nanomaterial modified with needle-like manganese oxide nanoparticles was formed and cast on carbon electrodes. We characterized the morphology and composition of this material using scanning electron microscopy (SEM), energy dispersive X-ray analysis (EDX), and X-ray photoelectron spectroscopy (XPS), and absorption spectroscopy. Finally, we investigated the electrochemical catalytic activity of the prepared interfaces using cyclic voltammetry and dose-response amperometry.

Experimental

Chemicals and reagents.—Ferrirhodopsin (hemin) (99%), synthetic graphite powder (<20 micron), sulfuric acid (95-98%), hydrogen peroxide (30%), dichloromethane, hexane, manganese (II) chloride tetrahydrate (99%), manganese (IV) oxide (99%), sodium nitrate (99%), sodium chloride (99%), potassium dihydrogen phosphate, dipotassium hydrogen phosphate, potassium permanganate, isopropanol, methanol, diethylene triamine pentaaetic acid (DTPA), ethanol, 3-morpholinosydnonimine hydrochloride (SIN-1) and isoamyl nitrite were all purchased from Sigma-Aldrich (St. Louis, MO) and used as such. Nano-pure deionized water (specific resistance > 18.2 MΩ cm) was supplied by a Barnstead water purification system model D8961. Indium Tin Oxide (ITO) coated glass slides were from Delta Technologies, Ltd. All other chemicals were reagent grade and used as received.

Instruments.—The electrochemical analyses were performed using CHI-1030B multi-mode potentiostat/galvanostat system. Amperometry and cyclic voltammetry measurements were carried out using the above mentioned electrochemical workstation. Ag/AgCl is used as the reference electrode and all potentiograms reported here are measured versus this reference. A platinum wire was used as the auxiliary electrode, and glassy carbon electrodes (GCEs) with cast graphene-hemin or manganese-modified graphene-hemin material were used as working electrodes. UV-Visible absorbance spectra were recorded on a Perkin Elmer Lambda-45 spectrophotometer. An Accumet AB15 pH-meter was used for pH measurements. SEM and EDX analyses were carried out using a Hitachi S-4500 Field Emission Gun Scanning Electron Microscope. X-Ray Photoelectron Spectroscopy (XPS) was performed using PHI VersaProbe 5000 Scanning spectrometer at a pressure ~ 10⁻⁷ millibar in the vacuum chamber. Monochromatic Al Kα X-ray beam was applied in the survey scans as well as in high-resolution analyses, with an estimated area of 0.35 mm². A pass energy of 93.9 eV was selected for sample surveys (0-800 eV), and 1.175 eV was chosen for high-resolution C1s (278-298 eV), whereas 23.5 eV was used for Fe 2p and Mn 2p (630-740 eV).

Synthesis and preparation.—Graphene oxide preparation.—In this study, a modified method of Hummers and Offeman was used to prepare graphene oxide.7 This method is based on the oxidation of graphite flakes in strong acid. In a typical preparation, 30 mL of concentrated sulfuric acid was added drop-wise to the solid mixture of 0.5 g of graphite powder and 0.3 g of sodium nitrate in a 250 mL round bottom flask at 0°C. The second step consists of slowly adding 2.0 g of solid potassium permanganate to the reaction vessel. The reaction mixture is continuously stirred for 1 hour at 60°C. Then, deionized water (130 mL) is gradually added to the mixture with continuous stirring for 0.5 hour. Finally, a solution of hydrogen peroxide (30%) is slowly added to the reaction flask until the gas evolution stops. The final product was centrifuged for 1 hour to remove water and the excess of oxidizing reagents. The residue is then washed repeatedly with hydrochloric acid solution (5%) followed by washing several times using deionized water and tested for sulfate ions using BaCl₂ reagent. When the washing solution shows no sulfate ions, the final product is dried overnight in an oven at 60°C to obtain the solid graphene oxide (GO).

Graphene Oxide-MnO₂/hemin composite.—The graphene oxide (GO) as prepared was then functionalized with MnO₂ nanoparticles by oxidizing manganese ions in situ as reported by Wang.25 Briefly, GO (0.066 g) and MnCl₂·4H₂O (0.270 g) are dispersed in isopropyl alcohol (50.0 mL), with ultrasonication for 1 hour. The reaction mixture was heated to ~ 80°C under reflux with vigorous stirring. KMnO₄ (0.150 g) is dissolved in 5.0 mL of deionized water and then added to the mixture above and refluxing was allowed to continue for one hour. The mixture was then allowed to cool to room temperature. Finally, the prepared graphene-MnO₂ composite material was centrifuged, washed, and then dried in air at 60°C for at least 12 hours. Finally, hemin was incorporated by adding 5 mM solution of iron protoporphyrin IX in 5.0 mL methanol to 5.0 mg of graphene-MnO₂ dispersed in 5.0 mL of deionized water and put under sonication. The sonication was continued for 5 hours, followed by centrifugation, and washing of the precipitate. The solid was then dried at 60°C to obtain the final composite of GO-MnO₂/hemin. For comparison, we also prepared reduced graphene oxide/hemin (rGO/hemin) using the same method as reported in details in a previous work.18

Peroxynitrite generation.—In this work, we used peroxynitrite released from 3-morpholinosydnonimine hydrochloride (SIN-1) at physiological pH as described earlier.18,49

Electrode modification.—Glassy carbon electrodes (GCEs) were polished on alumina pastes (0.3 and 0.05 μm respectively) using a Buehler microcloth polishing disk, followed by sonication in a mixture of ethanol and acetone for 1 minute to remove any polishing residue, then flushed repeatedly with a deionized water. The electrodes are then modified by the drop-casting method. A suspension of GO-MnO₂/hemin or rGO/hemin composite was sonicated in methanol for 1 hour and then cast on the surface of GCE. The electrode is left for 30 minutes to dry. This process was repeated three times using a 10-μL aliquot each time. Finally, the modified electrodes were washed several times with a deionized water and phosphate buffer solution pH = 7.4 before use in the electrochemical experiments.

Results and Discussion

Preparation and characterization.—A simple and direct method has been developed to prepare the composite GO-MnO₂/hemin matrix. The process uses acid oxidation of graphite flakes using Hummers method to form the GO sheets.47 The GO obtained was mixed with manganese chloride salt and dispersed in isopropyl alcohol and subjected to ultrasonication. Oxidation was allowed by adding KMnO₄ under reflux with stirring to obtain the composite GO-MnO₂ as described by Wang and Zhu.25

Hemin was then introduced to allow for π-π stacking interaction under ultrasonication for few hours to form the final matrix of GO-MnO₂/hemin. Scheme 1 illustrates the various preparation steps to form the first composite (GO-MnO₂), followed by addition of hemin to form the reduced GO/hemin (rGO/hemin) and GO-MnO₂/hemin composite materials. SEM analysis was used to study the change in morphology and surface features of the functionalized graphene-based composite materials. The SEM images of GO, rGO/hemin, GO-MnO₂, and GO-MnO₂/hemin are shown in Figure 1. The characteristic micrograph images illustrate the reaction steps of the final product GO-MnO₂/hemin nano-composite. The solid powder of sample collected at different preparation steps were mounted and subjected for the SEM analyses. The SEM images show a random orientation of GO nanosheets in both GO and rGO/hemin samples. Figures 1A and 1B. The SEM image of GO-MnO₂ shows needle-like structure of this composite nano- material. Figure 1C. This is similar to findings by Wang and Zhu who described the formation of the needle-like manganese oxide crystals on graphene oxide sheets via a mechanism of intercalation and adsorption of manganese ions on graphene oxide sheets. Oxidation then triggers nucleation and oriented growth of needle-like MnO₂ crystals on graphene oxide.25 The length of the nanoneedles in our preparation is in the 100-200 nm range, with ~40-50 nm width. This is consistent with data reported earlier.41,42 Finally, the SEM of GO-MnO₂/hemin
nanocomposite also shows the morphological characteristics needle-like structures GO-MnO2, Figure 1D.

We performed EDX elemental analysis of GO-MnO2/hemin sample and for comparison we also analyzed GO and rGO/hemin under the same conditions. Figure 2. The relative atomic weight ratios for the various samples are listed in Table I. The carbon/oxygen (C/O) atomic ratio is indicative of the degree of graphene oxidation, and assesses the nature of oxygen functionalities. The degree of oxidation of the prepared graphene oxide can be estimated from C/O ratio. Optimally oxidized graphene oxide C/O atomic ratio lies between 2.1 to 2.9. The samples of GO prepared in this work exhibited a C/O atomic ratio of 2.15, consistent with the range reported for well-oxidized graphene.

Another support for the optimal level of oxidation of the prepared GO comes from the yellowish color of the sample when it is dispersed in deionized water. The atomic weight ratio C/O of GO increased from 2.15 to 2.8 after mixing with hemin under ultrasonication. This result is evidence for some degree of reduction of the graphene oxide sheets upon treatment with hemin. This confirms the role of hemin as a reducing agent, as reported earlier using the deconvolution of C1s in high-resolution XPS.

In contrast, the Mn-based material, GO-MnO2/hemin, has a very low value of C/O atomic ratio, indicating more oxygen in the final composite product. The introduction of MnO2 nanoparticles obviously increases the relative amount of oxygen atoms and therefore the C/O atomic weight ratio of the corresponding material is expected to decrease significantly.

Figure 2 also shows the difference between the EDX spectra of rGO/hemin and GO-MnO2/hemin in terms of iron and manganese peaks. As expected, a new peak of Mn appears in the case of GO-MnO2/hemin and is absent in rGO/hemin, consistent with the formation of GO-MnO2/hemin nanocomposite. Also, the atomic weight percentage of iron increased from 1.75 in rGO/hemin to 2.87 atomic weight% in the final GO-MnO2/hemin material. The oxidation of manganese ions in the water-isopropanol solvent system allows for the oriented growth of manganese oxide nanoneedles. This process was shown to cause further exfoliation of graphene oxide sheets. The manganese oxide-driven exfoliation of graphene oxide is expected to allow for more hemin intercalation within graphene sheets in GO-MnO2/hemin, which may explain the increase of the Fe peak in the EDX spectrum of this nanocomposite material.

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<th>Table I. EDX Elemental Analysis of graphene oxide GO as prepared, and of GO after treatment with hemin (rGO/hemin) and of GO-MnO2 after treatment with hemin (GO-MnO2/hemin).</th>
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We then carried out XPS analysis to characterize the modified surfaces, and to identify the chemical changes in the various nanocomposites. As shown in Figure 3, high-resolution spectra have been obtained for C1s, N1s, O1s, Fe2p, and Mn2p of rGO/hemin and GO-MnO2/hemin as prepared. Initially, survey spectra were acquired for the graphene-based composites on the conductive ITO glass coated with the material in question using the drop-casting method. Figure 3A. The conductive ITO glass is also conveniently used, when needed, as a supporting electrode material for electrochemical analysis of the same material subjected to XPS. The XPS spectrum of blank ITO slide was taken under the same conditions and subtracted from the analyzed samples to eliminate the ITO background contribution.

The deconvolution of the C 1s spectrum is often used to measure the degree of oxidation or reduction that occurs on the GO sheets upon chemical changes. Specifically, the three deconvoluted peaks of C 1s at binding energies of 284.8 eV (C-C/C-H), 286.7 eV (C=O/H/C-N), and 289.1 eV (C=O) are used to determine the chemical environment (oxidation state) of the graphene-based structures. The reduction of GO by hemin results in a significant decrease of the level of oxygen-bound carbons (C=O/H/C-N), with concomitant increase in the amount of sp2 carbon hybridization. In the case of rGO/hemin versus GO-MnO2/hemin, we observe an overall decrease in the intensity of sp2/C-C peaks in the presence of MnO2 particles, which seems to indicate that the introduction of MnO2 nanoparticles prevents hemin from reducing GO at the level observed in plain rGO/hemin. The high-resolution XPS spectra of C1s and N1s have almost the same binding energies at 283.8 and 397.6 eV, respectively, for rGO/hemin and GO-MnO2/hemin, indicating similar chemical states for both composites with respect to carbon and nitrogen contributions. Figures 3C and 3D. The intensities of both C1s and N1s peaks of rGO/hemin are higher than the GO-MnO2/hemin nanocomposite due to the predominant graphene and protoporphyrin moieties compared to the manganese-oxide nanoparticles in the final matrix. Interestingly, the O1s peak of the GO-MnO2/hemin matrix shows higher intensity and a slight negative chemical shift in binding energy compared to rGO/hemin (compare 530.0 eV to 531.8 eV). Figure 3B. This negative shift suggests that there is more contribution of oxygen as C=O or HO-C=O moieties in GO-MnO2/hemin compared to rGO/hemin. This conclusion is also supported by the careful analysis of the C 1s spectra of GO-MnO2/hemin, which shows small peaks at 288, 291, and 293.5 eV, assigned to C=O/HO-C=O groups and π-π conjugation, respectively. Together, these observations further support the conclusion that the incorporation of MnO2 nanoparticles seems to prevent hemin molecules from reducing GO sheets to the level observed in rGO/hemin samples.

As expected, GO-MnO2/hemin features characteristic XPS peaks of manganese, which is absent in rGO/hemin samples. Figure 3F. The Mn2p spectrum of GO-MnO2/hemin exhibits two peaks for 2p3/2 and 2p1/2 chemical states at 642 and 653 eV respectively, with binding energy difference of 11 eV as reported in the literature. The incorporation of hemin in the prepared composites is confirmed by the presence of the iron peaks. The two materials exhibit the same binding energy for the iron peak, which confirms similar chemical state of iron in both composites. Figure 3E. The spin-orbital coupling of Fe 2p3/2 and Fe 2p1/2 for both composites appears at binding energies of 712 and 725 eV, which is consistent with published data elsewere. Interestingly, the rGO/hemin and GO-MnO2/hemin composites both have similar iron peak intensities in XPS. However, in EDX analysis shows a relative increase in the iron peak in GO-MnO2/hemin compared to rGO/hemin. The EDX info depth is >500 nm compared to only 1-5 nm surface penetration in XPS analysis. The deeper sampling of EDX explains why this method better reflects the relative Fe % in the GO-MnO2/hemin composite compared to rGO/hemin. The higher Fe content -and thus hemin content- in GO-MnO2/hemin compared to rGO/hemin is consistent with the finding earlier that the oriented growth of needle-like crystals of MnO2 on graphene oxide leads to the exfoliation of GO sheets, and thus a relatively higher surface area for hemin intercalation.

We used UV/Vis analysis to investigate the formation of GO-MnO2 and GO-MnO2/hemin nanocomposites as shown in Figure 4. In addition, UV/Vis absorption analysis of the initial components (starting materials) GO and hemin was conducted for comparison. A dispersed sample of GO and GO-MnO2 nanomaterials in water were analyzed to monitor the composite formation and the observed changes in GO.

As observed in Figure 4A, the UV/Vis spectrum of GO suspension shows a maximum absorption at 228 nm, which is ascribed to the π-π* transitions of the carbon in GO layers. In addition, a small shoulder is observed at 298 nm that is attributed to n-π* transitions of C=O functionalities. Interestingly, the GO-MnO2 nanocomposite exhibits also two predominant absorption peaks at 219 nm and 430 nm corresponding to GO (π-π* transitions) and MnO2 (d-d transitions), respectively. The blue-shift of the graphene peak is attributed to the exfoliation of graphene sheets by MnO2 nanoneedles as reported by Wang et al. They reported blue-shift for the graphene peak and a concomitant red-shift for the d-d of Mn as the reaction of formation of GO-MnO2 proceeds. Increasing the time of reaction causes the peak of MnO2 to gradually shift from 413 to 436 nm, while the GO peak shifts from 250 to 216 nm.

The UV/Vis absorbance spectra of GO, GO-MnO2, hemin, and GO-MnO2/hemin are shown in Figure 4. The absorption
Figure 3. Survey and High-Resolution XPS analyses of rGO/hemin (blue) and GO-MnO₂/hemin (black) samples: (A) XPS survey of the rGO/hemin and GO-MnO₂/hemin samples cast on ITO slide. Background contribution of ITO was subtracted using spectra acquired with non-coated ITO slides as blanks as described in the text (B) O1s High Resolution XPS spectra of rGO/hemin and GO-MnO₂/hemin samples, (C) C1s High Resolution XPS spectra of rGO/hemin and GO-MnO₂/hemin, (D) N1s High Resolution XPS spectra of rGO/hemin and GO-MnO₂/hemin, (E) Fe2p High Resolution XPS spectra of rGO/hemin and GO-MnO₂/hemin samples, (F) Mn2p High Resolution XPS spectra of rGO/hemin and GO-MnO₂/hemin samples under the same conditions.
characteristics of the GO-MnO$_2$/hemin nanocomposite suspended in water shows the Soret absorption band of incorporated hemin, as well as the peak of GO. For comparison, absorption characteristics of hemin in solution confirms that the absorption at approximately the same wavelength (≈400 nm) is that of hemin intercalated within the GO sheets. The incorporation of hemin in the graphene-manganese nanocomposite is also indicated by the presence of weak peaks between 420 nm to 640 nm corresponding to the hemin Q-bands. Together, these spectroscopic data show that the hemin rings are incorporated as monomeric entities within the GO-MnO$_2$ nanomaterial.

The relatively weak d-d transition peak of MnO$_2$ is obscured by the intense hemin Soret-band in the final matrix nanocomposite; however, a careful observation still shows the broad shoulder at 430 nm for the GO-MnO$_2$/hemin final material.

In conclusion, EDX, XPS, and UV/Vis analyses of the GO-MnO$_2$/hemin material show the general features of all matrix components, and together confirm the formation of the target nanocomposite material. We next use this prepared material on electrode surfaces to test its activity in catalytic sensing of peroxynitrite.

**Electrochemical measurements and sensor performance.**— The electrochemical performance of the rGO/hemin and GO-MnO$_2$/hemin-modified GCEs was examined side-by-side using cyclic voltammetry analysis in phosphate buffer. The brown-red suspension of rGO/hemin and the brown-black suspension of GO-MnO$_2$/hemin were drop-cast on freshly polished glassy carbon electrodes (GCEs) and allowed to dry before use as described in detail in the Experimental section. The typical voltammetric response in absence and presence of peroxynitrite (PON) is shown in Figure 5. We used SIN-1 as a peroxynitrite generator in situ at physiological pH. In the presence of PON, the rGO/hemin voltammogram shows an oxidative peak response at ∼+1.1 V attributed to the catalytic oxidation of PON mediated by hemin on graphene. The mechanism of the catalytic activity of hemin-mediated peroxynitrite oxidation has been previously reported. A similar mechanism is proposed for GO-MnO$_2$/hemin and is shown in Scheme 2.

The incorporation of MnO$_2$ in the graphene/hemin composite shifts the catalytic oxidation peak of peroxynitrite to a lower peak potential by about 200 mV. As a result, a well-defined peak for the catalytic oxidation appears at approximately +0.9 V vs. Ag/AgCl as shown in Figure 5B. The shift of the catalytic peak to less positive potentials indicates that the incorporation of MnO$_2$ particles on GO/hemin nanocomposite helps lower the barrier of mediated electrochemical oxidation of PON at the modified interface. Also, as we established earlier, the added exfoliation of GO enabled by the needle-like MnO$_2$ nanoparticles allows for the incorporation of more hemin catalyst in the nanocomposite material. Both effects result in higher current response at less positive potential.

Amperometric measurements have been used to compare and contrast the electrocatalytic activity and the performance of the various composite materials. Again, we used nanomolar-range aliquots of PON released from 1μM stock solution of SIN-1.

The amperometric response of the modified electrodes to added PON aliquots was measured after each injection for sensors with GO-MnO$_2$/hemin as well as with rGO/hemin for comparison. The applied potential in both cases was of +1 V vs. Ag/AgCl.

Figure 5A shows the typical current responses associated with the addition of PON aliquots to the electrochemical cell in phosphate buffer at pH 7.4. As expected based on cyclic voltammetry, we observe a significantly larger response in catalytic oxidation current of PON on electrodes modified with GO-MnO$_2$/hemin as compared to electrodes modified with rGO/hemin. Calibration curves based on amperometry show linear correlation between measured currents and added PON concentration in the range of 2.0 to 10.0 nM for both
systems. Linear regression analyses show a sensitivity of 23.7 nA/nM for GO-MnO$_2$/hemin as opposed to only $\sim$10 nA/nM for rGO/hemin. The relative detection limit of the GO-MnO$_2$/hemin-modified sensors was 2.0 nM with overall good sensor stability.

Conclusions

We have described a preparation method of sensors with a nanomaterial based on hemin- and MnO$_2$-functionalized graphene oxide nanosheets. Glassy carbon electrode sensors with the prepared nanocomposite material showed relatively higher catalytic activity for peroxynitrite detection in phosphate buffer at pH 7.4. This work highlights for the first time the effect of MnO$_2$ incorporation in GO/hemin on carbon-based sensors toward the electrocatalytic detection of PON. The incorporation of MnO$_2$ nanoparticles not only allows for more adsorbed hemin entities into the final composite matrix, but also lowers the barrier of mediated electrochemical oxidation of PON. As a result, amperometric calibration curves of the GO-MnO$_2$/hemin-modified sensors show a significant increase in sensitivity toward peroxynitrite detection in physiological pH compared to rGO/hemin sensors. We are now adapting the same catalytic interface (GO-MnO$_2$/hemin) on a miniaturize platform consisting of carbon fiber electrodes in order to test their use as sensors for peroxynitrite at the level of single cells or collection of cells.

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