Axial Coordination Site-Turned Surface Confinement, Electron Transfer, and Bio-Electrocatalytic Applications of a Hemin Complex on Graphitic Carbon Nanomaterial-Modified Electrodes

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ABSTRACT: Understanding the relation between the chemical bonding and the electron-transfer (ET) reaction of surface-confined hemin (a five-coordinated Fe-porphyrin-with-chlorine complex) is a special interest in the biomimicking studies of heme proteins. Owing to the difficulty in ET function studies, scanty electrochemical reports of hemin in aqueous solutions were reported. It has been noticed that in most of the reported procedures, the sixth axial coordination position of the hemin complex has been unknowingly turned by attaching with water molecules (potential cycling in alkaline conditions or heating), solvents such as ethanol and dimethyl sulfoxide, and nitrogen-donating compounds that have helped for the heme ET reaction. In this work, a systematic effort has been taken to find out the contribution of hemin and its axial bond coordination with π–π interaction, hydrogen bonding, and hydrophobic binding systems toward the ET reaction. Various graphitic carbons such as graphitized mesoporous carbon (GMC), mesoporous carbon-hydrophilic and hydrophobic units, graphite nanopowder, graphene oxide, single-walled carbon, multiwalled carbon nanotube (MWCNT), and carboxylic acid-functionalized MWCNT (as a source for π–π interaction, hydrogen bonding, and hydrophobic environment) along with the amino functional group of chitosan (Chit; as an axial site coordinating system) have been tested by modifying them as a hemin hybrid on a glassy carbon electrode (GCE). In addition, a gold nanoparticle (Au nanoparticles) system was combined with the above matrix functionalized MWCNT (as a source for π–π interaction, hydrogen bonding, and hydrophobic environment) and Au nanoparticles on GMC@hemin-Chit hybrid system, wherein all interactive features have been utilized. Omitting any of the individual interactions resulted in either decreased (with Au nanoparticles) or nil current response. As applications, efficient bio-electrocatalytic reduction and sensing of dissolved oxygen and hydrogen peroxide have been demonstrated.

1. INTRODUCTION

Hemin, a five-coordinated Fe-porphyrin-with-chlorine complex (Scheme 1), is a derived active site of the heme-containing proteins and enzymes such as hemoglobin (Hb), cytochrome C, horseradish peroxidase, and catalase. It is produced endogenously in the human body during the turnover of red blood cells (RBCs). Indeed, it forms inappropriately as a result of vascular (vessels that carry blood) injury or hemolysis (rupture or destruction of RBCs) in the physiological system. It has been revealed that the endogenously formed hemin helps in intracellular generation of reactive oxygen species (ROS) such as hydrogen peroxide, singlet oxygen, hydroxyl radicals, and superoxide. At the same time, this hemin has the capacity to interact with the peroxide and oxygen molecules and facilitates their bio-electrocatalytic reduction to water by mediated electron-transfer (ET) reaction mechanism (at specific operating potentials). Note that meticulous ROS production in human cells assists in energy production and apoptosis and supports to combat pathogens. Indeed, abandoned ROS leads to permanent damage of the cell organelles and tissues. Biomimicking the ET function of the hemin and probing the mysteries of its biological function are a continued research interest. In this regard, fixing hemin as a surface-confined species similar to the natural system is an important step. Unfortunately, there is no clear and systematic information about ET of surface-confined hemin complex covering π–π and hydrogen bonding and axial ligand coordination, in the literature. The electro-inactive character...
of hemin on solid electrodes is a prime reason for the limitation. Herein, we report a meticulously designed hemin-confined carbon nanomaterial (graphitic base)/chitosan (Chit) chemically modified electrode, designated as glassy carbon electrode (GCE)/carbon@hemin−Chit, for an efficient ET and mediated reduction of dissolved oxygen and hydrogen peroxide in a physiological system.

With the aim of biomimetic studies, since 1928, there has been significant interest on the ET feature of hemin (homogenous condition) in organic and semiaqueous mediums. In 1967, Bednarski and Jordan first reported the adsorption and ET study of hemin on the dropping Hg electrode in 0.1 M KOH solution and stated that hemin forms a soluble dimer, which undergoes a two-electron reversible reduction to yield a ferroheme monomer (wherein two water molecules are attached on the two axial positions). Following this, there were several studies on hemin complex with Hg systems. In 1990, Bianco et al. reported adsorption of hemin on a pyrolytic graphite electrode (PGE) and showed an ET reaction at \( E^{\text{o}} \approx -0.55 \text{ V} \) versus Ag/AgCl with a surface excess of \( \Gamma_{\text{hemin}} = 11 \times 10^{-10} \text{ mol cm}^{-2} \) in pH 7 phosphate buffer solution (PBS). In their preparation procedure, the PGE was simply exposed to hemin solution containing 30% ethanol/borate buffer of pH 11. It has been claimed that the \( \pi-\pi \) interaction between the graphitic sites and hemin is responsible for the adsorption and ET feature. In a similar fashion, hemin was adsorbed on a multiwalled carbon nanotube (MWCNT) via \( \pi-\pi \) interaction and showed \( E^{\text{o}} = -0.34 \text{ V} \) versus Ag/AgCl and \( \Gamma_{\text{hemin}} = 27 \times 10^{-10} \text{ mol cm}^{-2} \). Later on, a covalently immobilized hemin on an amino-functionalized MWCNT-modified electrode via 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide (EDC) \( (E^{\text{o}} \approx -0.5 \text{ V} \text{ vs Ag/AgCl in pH } 1 \text{ buffer}) \) was reported. Recently, hemin-immobilized graphene oxide (GO)-a mixture of ethanolic solution of hemin−GO and gold nanoparticle−GO (a mixture of ethanolic solution of hemin−GO−Au nano was used; \( E^{\text{o}} = -0.4 \text{ V} \text{ vs Ag/AgCl, } \Gamma_{\text{hemin}} \approx 2 \times 10^{-10} \text{ mol cm}^{-2} \))−based chemically modified electrodes were reported in the literature. All of those studies emphasize the \( \pi-\pi \) interaction for the ET process. Meanwhile, there were few reports on hemin surface-confined gold self-assembled monolayer electrodes utilizing nitrogen-containing biomolecules such as histidine and guanine quadruplex (G4) complexes with \( E^{\text{o}} = -0.22 \text{ to } -0.35 \text{ V} \) versus Ag/AgCl and \( \Gamma_{\text{hemin}} = 0.0016\text{ to } 2.2 \times 10^{-10} \text{ mol cm}^{-2} \). These above reports highlight the importance of the planar orientation and axial ligand interactions for the ET activity.

In this report, \( \pi-\pi \) interaction, axial ligand coordination, and multihydrogen bonding were collectively tuned for the enhanced ET reaction of the hemin on a carbon nanomaterial/Chit-modified electrode system. Importance of the axial ligand coordination in the ET of hemin was first reported by Richard et al. by studying a series of 29 pentacoordinate hemin derivatives using the cyclic voltammetry (CV) technique in a noncoordinating solvent \( (\text{CH}_2\text{Cl}_2) \). A linear correlation between the \( E^{\text{o}} \) values of the
Figure 1. Ten continuous CV responses of (A) (a) GCE/hemin, (b) GCE/GMC@hemin, (c) GCE/GMC@hemin–Chit, and (d) GCE/Au nano–GMC@hemin–Chit and various GCE/Au nano–carbon nanomaterial@hemin–Chit systems: (B) mesoporous carbon hydrophilic unit, (C) mesoporous carbon hydrophobic unit, (D) carbon nanofiber, (E) SWCNT, (F) MWCNT, (G) carboxylic-functionalized MWCNT, and (H) GO in N2-purged pH 7 PBS at a scan rate of v = 50 mV s⁻¹.

2. RESULTS AND DISCUSSION

2.1. CV Study of GCE/Au nano–Carbon@Hemin–Chit.

Initial experiments were carried out using GMC as a standard system. At first, the hemin complex is adsorbed on a bare GCE surface and subjected to CV study in N2-purged pH 7 PBS. As can be seen in Figure 1A, curve a, a featureless voltammetric response was noticed. This observation is the same when the hemin is modified on GCE/GMC as GCE/GMC@hemin (Figure 1A, curve b) (Scheme 1A–C). Note that, even after providing a graphitic environment by GMC, which can support π−π interaction, there is no specific redox signal for hemin. This is the reason why, in previous reports, hemin surface-confined graphene and GO/gold nanoparticle systems failed to show any marked characteristic redox peak at an apparent electrode potential of E° = −0.320 V versus Ag/AgCl in neutral pH solutions, affirming that graphitic interaction alone could not be a prime factor for the ET reaction. Interestingly, when a dilute solution of Chit polymer which contains an amino functional group is coated on GCE/GMC@hemin, as GCE/GMC@hemin–Chit, a well-defined redox peak at E° = −380 ± 5 mV versus Ag/AgCl with a surface coverage (Γ_hemin) = 62 × 10⁻¹⁰ mol cm⁻² appeared because of the ET of the surface-confined hemin (Figure 1A, curve c). Meanwhile, a 1% NaFon (Nf) polymer, a characteristic cationic exchange polymer without any amino functional group, is casted on GCE/GMC@hemin, as GCE/GMC@hemin–Nf, and tested (Supporting Information Figure S1). In addition, hemin adsorbed on the Chit-modified electrode, that is, GCE/Chit@hemin, was also tested (data not included). There is no formation of the redox peak in the previous cases. These observations highlight the important role of the amino functional group along with the graphitic environment for the redox tuning of the hemin. In the literature, the amino group in Chit has been referred to as a good coordinating site. Our group utilized Chit for the complexation with the iron site buried in the pristine MWCNT. Similarly, Chit–iron(III)-modified smectites, a [Fe(CN)₆(NH₃)]₃⁻ complex-incorporated Chit hybrid film, and an Fe(III)–Chit complex have
been reported in the literature utilizing complexation of the amino functional group with the iron site. In this work, it is expected that the amino functional group of Chit is bonded on the sixth vacant position of the hemin (axial bond) and helped the ET process.

With an aim to improve the performance and literature reports based on molecular wiring systems, Au nano particles were chosen as a wiring agent and incorporated in the optimal electrode as a GCE/Au nano–GMC@hemin–Chit configuration. The preparation procedure was optimized as casting a dilute solution of Au nano ion on the GCE as the first step followed by other modification procedures. During the electrochemical potential cycling experiment, the Au nano ion was in situ reduced as Au nano particles and molecularly wired in the matrix (Scheme 1H,I). Figure 1A (curve d) shows a typical CV response of GCE/Au nano–GMC@hemin–Chit, showing the redox peak at 10−320 ± 2 mV versus Ag/AgCl with a 2-fold enhancement in the $\Gamma_{\text{hemin}}$ value (120 × 10−10 mol cm−2). Further, $\Delta E_{p}$ noticed here is about 70 mV positive than the value obtained without the Au nano particle-modified electrode (10−390 mV), supporting the molecular wiring within the matrix. The $E^o$ value (10−320 mV) is matching with previous reports on hemin-modified electrodes, for example, hemin-MWCNT ($E^o = 10−0.34$ V vs Ag/AgCl)16 Au nano–GO–hemin ($E^o = 10−0.4$ V vs Ag/AgCl),19 and hemin-self-assembled gold monolayers via histidine and guanine quadruplex (G4) complexes ($E^o = 10−0.22$ to −0.3 V vs Ag/AgCl).20,21 The calculated peak-to-peak separation value ($\Delta E_{p}$) is 10−40 mV at $v = 10$ mV s−1, indicating a fast ET process. It is noteworthy that the $\Gamma_{\text{hemin}}$ value noticed in this work is the highest compared to the previous literature reports on the hemin-surface-confined system such as PGE@hemin (11 × 10−10 mol cm−2),15 MWCNT@hemin (27 × 10−10 mol cm−2),19 and N-doped GO@hemin (13 × 10−10 mol cm−2).29 In continuation, NF polymer-coated GCE/Au nano–GMC@hemin (GCE/Au nano–GMC@hemin–Nf) was also examined for the redox activity (data not enclosed) (Scheme 1D). However, nil redox response was noticed, confirming the specificity of Chit for the axial coordination ET process. On the basis of the results, it has been proposed that π–π interaction, axial coordination, hydrogen bonding (oxygen and hydrogen functional groups between hemin, GMC, and Chit), and internal linking (wiring) factors have been collectively involved in the effective ET process in this work.

From the CV study of the effect of scan rate ($v$), it is confirmed that ET of GCE/Au nano–GMC@hemin–Chit is surface-confined in nature (anodic and cathodic peak currents $i_{pa}$ and $i_{pc}$ vs $v$ are linear starting from the origin) (Figure 2A,B). The ratio of the cathodic and anodic peak currents, $i_{pc}/i_{pa}$, is found to be nearly unity, obeying the reversible nature of the ET process. The heterogeneous rate constant $k_{h}$ was calculated from the Laviron’s model at condition $\Delta E_{p} < 200$ mV using the equation $k_{h} = mFv/RT$, where $m$ is a dimensional-less parameter relating to $\Delta E_{p}$ and transfer coefficient value ($\alpha$), $n$ is the number of ET ($n = 1$), $v$ is the scan rate, and other symbols have their own significance.41 Priorly, the values $E_{pc} = E^o + 1$ and $\Delta E_{p} = 99$ mV at $v = 100$ mV s−1 (Figure 2C) were applied in the Laviron’s working plots,40 and the $\alpha$ value was calculated as 0.5. On the basis of the parameters, the $k_{h}$ value at $v = 100$ mV s−1 is then calculated as 1.3 s−1. This value is comparable with the values reported for HB-based surface-confined systems such as HB–MWCNT–NF (1.25 s−1),16 HB–F–CNT–CTAB surfactant–NF (1.25 s−1), HB–ionic liquid–MWCNT–CPE (0.84 s−1), and HB–IL–MWCNT–carbon paste electrode (0.99 s−1).42 Further, the redox peak is pH-dependent in nature (Figure 2D). The plot of $E_{pc}$ versus pH is linear with a slope of 10−36 mV pH−1, indicating the non-Nernstian behavior (expected slope = 10−58 mV pH−1).43,44

Previously, a similar kind of observation was made with several HB-modified electrodes: HB–polymer-grafted MWCNT (42 mV pH−1)16 and HB–C–CTAB surfactant–NF (42 mV pH−1) systems.45

Partial protonation of ligands is the likely reason for this observation.46 At this stage, it is difficult to predict the exact molecular orientation of the hemin in this new system. Further, because the modified electrode is complex in nature and the concentration of the immobilized species is at the nanogram level, it is not convenient to use conventional characterization techniques such as nuclear magnetic resonance and mass spectroscopy to identify the active site. On the other hand, from the control experiments and physicochemical characterization studies, the possible structure and orientation of the surface-confined hemin system can be predicted.

The concept of immobilizing hemin is extended to other carbon nanomaterials as well. Figure 1B–H shows the typical CV responses of the surface-confined hemin systems on various carbons such as CM-hydrophilic unit, CM-hydrophobic unit, GNPs, SWCNT, MWCNT, f-MWCNT, and GO prepared similar to the GCE/Au nano–GMC@hemin–Chit case. With respect to the $\Gamma_{\text{hemin}}$ value, the carbon materials were sequenced as follows (ascending order): GO (no graphitic structure; no hydrophobic unit (61 × 10−10 mol cm−2) < GNPs (8 × 10−10 mol cm−2) < f-MWCNT (15 × 10−10 mol cm−2) < CM-hydrophilic unit (61 × 10−10 mol cm−2) < SWCNT (90 ×
10^{-10} \text{ mol cm}^{-2} < \text{MWCNT} (101 \times 10^{-10} \text{ mol cm}^{-2}) < \text{GMC} (120 \times 10^{-10} \text{ mol cm}^{-2}). Following important conclusions can be derived from the observations: (i) The graphitic structure is necessary. (ii) Comparing hydrophilic and hydrophobic structures, hydrophobic units have positive assistance for ET process. (iii) Oxygen functional groups such as carboxylic and ether in f-MWCNT and GO have a negative effect with the functional group of hemin. Possibly, there may be electrostatic repulsions between the oxygen functional groups (hemin-COOH, pK_a \approx 3.8)\textsuperscript{22} and hence avoid the communication. Indeed, a fraction of the carbon’s oxygen functional groups helped for the hydrogen bond formation. (iv) The MWCNT structure is better than the single-walled structure, which may be due to the highly porous nature. (v) The graphitic structure along with the porous site provided by the GMC is the best option for the stable entrapment of the hemin system. Overall, it is obvious that tuning of \pi-\pi interaction, hydrophobic interaction, hydrogen bonding, and axial ligand coordination is necessary for the effective ET reaction. Partial or incomplete tuning of the interactions may result in either poor or nil ET process.\textsuperscript{17,19,28,47} Because of the well-defined redox peak response, highest \Gamma_{hemin} and low material cost, GMC has been chosen as a model system in this work.

In order to find out the interaction of Au\textsubscript{nano} on the hemin functional group, control experiments were carried out with other forms of hemin derivative, hematin, modified electrodes, that is, GCE/Au\textsubscript{nano}–GMC@hematin–Chit and GCE/GMC@hematin–Chit (without Au\textsubscript{nano}). The hematin electrodes were prepared similar to GCE/Au\textsubscript{nano}–GMC@hematin–Chit and tested. Figure 3 (curves a and b) shows the typical CV responses of GCE/Au\textsubscript{nano}–GMC@hematin–Chit (curve b) and GCE/GMC@hematin–Chit (curve a) in N\textsubscript{2}-purged pH 7 PBS at \nu = 50 \text{ mV s}^{-1}. Unlike the hemin case with a 50% increment in the \Gamma_{hemin} value (Figure 1A, curve c), there is no significant effect of Au\textsubscript{nano} on the redox feature of the hematin (Figure 3, curves a and b). Both Au\textsubscript{nano}–modified and unmodified hematin electrodes showed almost similar redox peaks signals (Figure 3). Note that, both hemin and hematin have the same molecular structure and varies only in the fifth axial ligands, where in, chloride (Cl) with hemin and hydroxide (OH) with hematin (Scheme 1). From these results, it can be revealed that Au\textsubscript{nano} has a positive interaction with the chloride ligand of hemin (likely chemisorption) as [Fe–Cl–Au\textsubscript{nano}] (molecular wiring) in turn for the enhanced communication.

### 2.2. Physicochemical Characterizations

Transmission electron microscopy (TEM) images of Au\textsubscript{nano}–GMC@hemin–Chit and GMC@Chit systems are shown in Figure 4. There is no marked change in the structural feature, except for black shades on the graphitic structure and fine dots of \textasciitilde 20–40 nm in size with Au\textsubscript{nano}–GMC@hemin–Chit. These preliminary observations may support \pi-\pi interaction between the hemin and the graphitic structure of GMC and the presence of some nanoparticles in the matrix. Extended experiment with TEM–energy-dispersive X-ray spectrometry (EDXS) showed signals corresponding to Cu (because of the copper grid), Au, and Fe atoms (Supporting Information Figure S2). This observation confirms the existence of hemin-Fe and Au systems on the modified electrode. Supporting Information Figure S3 shows comparative Raman spectroscopic responses of Au/GMC@hemin–Chit and unmodified GMC displaying specific D and G bands corresponding to the graphitic (hexagonal sp\textsuperscript{2} carbons) and disordered structure (sp\textsuperscript{3} carbons), respectively.\textsuperscript{46} The calculated intensity ratio between D and G bands, I\textsubscript{D}/I\textsubscript{G}, for Au/GMC@hemin–Chit (0.214) is less than that for the unmodified GMC (0.2707). These data indicate an improvement in the graphitic structure of GMC by the hemin surface confinement through \pi-\pi interaction. Although the porphyrin rings have \pi bonds, because of the improper orientation, they may not be completely aligned on the six-membered \pi structure of the GMC-graphitic units. On the other hand, the alkene tail in hemin is expected to have positive \pi-\pi interaction with the graphitic structure (Scheme 1). UV–vis spectroscopic characterization was also used for the structural identification in this work. For that, respective GCE-modified electrode was sonicated with 500 \mu L of PBS and filtered with a microsyringe. Figure 5A shows a typical UV–vis response of...
Aunano−GMC@hemin−Chit, hemin, and Au\textsuperscript{3+} solution (precursor). Specific absorption signals at $\lambda_{\text{max}} \sim 405$ and 534 nm were noticed. Compared with the control hemin and Au\textsuperscript{3+} solution and the data from the literature related to hemin−GO hybrid ($\lambda_{\text{max}}$ = 413 nm)$^{28}$ and Aunano ($\lambda_{\text{max}}$ = 520−535 nm; 20−70 nm-sized particles),$^{50}$ the UV−vis signals can be identified as hemin (hybrid) and Aunano particles, respectively. Figure 5B shows comparative Fourier transform infrared (FTIR)/KBr responses of various modified electrodes prepared by pealing the layer of the modified electrode and combined with KBr.

The Aunano−GMC@hemin−Chit system showed specific IR signals at 3550 (νOH), 2980 (νC−C), 2356 (νC−H), 1713 (νC=O), 1237−1343 (νC−NH−C and Aunano), 886 (νC5−ring), and 717 (ν−C\textsuperscript{=CH}) cm$^{-1}$ for various characteristic functional groups along with a specific intense signal at 1044 cm$^{-1}$ due to the Fe−N stretching, selectively confirming amino group complexation with the axial position of the hemin-Fe site.$^{51}$ These characteristic IR signals (except the Fe−N stretching) are found to be slightly shifted with respect to the signals of the corresponding control samples: GMC (3450, 3350, 2930, 2355, 1640, 1540, 1370, 1150, 715 cm$^{-1}$).

Figure 5. (A) UV−vis and (B) FTIR spectroscopic analysis results of Aunano−GMC@hemin−Chit with various control samples.

Figure 6. (A) Comparative CV responses of GCE/Aunano−GMC@hemin−Chit without (a) and with (b) dissolved oxygen (10 ppm) in pH 7 PBS. Curve c is GCE/Aunano−GMC@hemin−Chit in dissolved oxygen (10 ppm) in pH 7 PBS. (B) CV responses of GCE/Aunano−GMC@hemin−Chit with increasing concentration of DO at $v = 10$ mV s$^{-1}$ and a photograph of the DO meter. (C) Corresponding calibration plot. (D) Comparative CV responses of GCE/Aunano−GMC@hemin−Chit without (a) with and (b) H$_2$O$_2$ in pH 7 PBS. Curve c is GCE/Aunano−GMC@hemin−Chit in H$_2$O$_2$. (E) CV responses of GCE/Aunano−GMC@hemin−Chit with increasing H$_2$O$_2$ concentrations at $v = 10$ mV s$^{-1}$. (F) Corresponding calibration plot.
The information supports the existence of various functional group interactions such as π–π, hydrophilic, and hydrogen bonding within the new chemically modified electrode.

2.3. Bio-Electrocatalytic Reduction of Dissolved Oxygen and Hydrogen Peroxide. Figure 6A shows the CV response of GCE/Au nano–GMC@hemin–Chit with (10 ppm) and without oxygen gas saturation in pH 7 PBS at v = 10 mV s⁻¹. A N₂ gas-purged pH 7 PBS was used as an electrolyte. A well-defined oxygen reduction signal at an onset potential, 0.25 V versus Ag/AgCl with peak current maximums at 0 and −0.38 V versus Ag/AgCl, where the hemin redox peak exists, was noticed. A control experiment with a hemin unmodified electrode, GCE/Au nano–GMC–Chit showed a reduction signal about 300 mV higher negative potential and about twice lesser in the peak current value than that of the optimal electrode. This observation denotes the hemin redox-mediated reduction of the dissolved oxygen. A pre-reduction peak at about 0 V versus Ag/AgCl noticed with the optimal working electrode is due to immobilization of the hemin at energetically different surfaces such as pores and valleys/asperities of the GMC and its interaction with analytes. Presumably, hemin trapped at the pores of the GMC, which provide significant π–π and hydrogen bonding, is responsible for the low-potential peak observation in this work. Figure 6B shows the effect of dissolved oxygen concentration on the mediated oxygen reduction current signals. The dissolved oxygen concentration was measured discreetly by a DO meter (the inset of Figure 6B). A systematic increment in the peak current against the increase in the dissolved oxygen concentration was noticed. The plot of peak current versus dissolved oxygen concentration was linear in a range of 0–8 ppm with a current sensitivity value of 0.794 µA ppm⁻¹ (Figure 6C). This result highlights the potential use of the new system for dissolved oxygen sensor development.

In addition, electrocatalytic reduction of hydrogen peroxide was also demonstrated with this new hemin-modified electrode system, as in Figure 6D. The H₂O₂ reduction signal was noticed at an onset potential of −0.38 V versus Ag/AgCl with peak maximums at −0.05 and −0.4 V, similar to the case of the oxygen reduction reaction, due to the mediated reduction by the surface-confined hemin sites. The plot of the reduction peak current signal (I_{pc}^{H₂O₂}) versus concentration of H₂O₂ is linear in a window 5 µM−1 mM with a current sensitivity value of 0.044 µA µM⁻¹. With an aim to extend to electroanalytical assays, amperometric i–t analysis of H₂O₂ at an applied potential of 0 V versus Ag/AgCl in N₂-purged pH 7 PBS was also performed by spiking 25 µM H₂O₂ in a stirred electrolyte solution (Supporting Information Figure S4A). A regular increase in the current signals against the H₂O₂ spikes was noticed. The calibration plot of baseline-corrected current signals versus [H₂O₂] was linear in the range 0–250 µM with a relative standard deviation of 3.1% (S/N = 3) (Figure S4B). The calculated detection limit was 6.8 µM. The analytical results are comparable with the previous literature reports on Hb-based chemically modified electrodes, wherein the detection limit value is in a range of 4–45 µM. In addition, the effect of interferences from various biochemicals such as nitrite (NO₃⁻), nitrate (NO₂⁻), glucose (Glu), uric acid (UA), ascorbic acid (AA), xanthine (Xan), hypoxanthine (hyp), cysteine (CysH), and dopamine (DP) on the detection of 50 µM H₂O₂ at an applied potential of 0 V was analyzed (Supporting Information Figure S4C). There is no significant alteration in the reduction current signals upon spiking the aforementioned interfering biochemicals. This observation indicates the selective H₂O₂, electrochemical interaction and reduction similar to the heme-based protein/enzymatic systems. GCE/Au nano–GMC@hemin–Chit is found to be stable at room temperature for 30 days and beyond (Supporting Information Figure S5), unlike the enzymatic systems, which have poor stability.

3. CONCLUSIONS

Highly redox active hemin-complex surface-confined carbon nanomaterial/Chit/Au nano chemically modified electrodes have been prepared by a simple drop-casting technique within 18 ± 2 min. Cyclic voltammetric investigation of the modified electrode showed a well-defined redox peak at E° = −320 ± 2 mV versus Ag/AgCl with peak-to-peak separation and surface excess values of 40 mV (10 mV s⁻¹) and 120 × 10⁻¹⁰ mol cm⁻², respectively, in N₂ gas-purged pH 7 PBS because of the facile ET behavior of the iron site in the hemin. Control experiments with the absence of Chit and graphitic carbon nanomaterial resulted in nil redox peak response. The new hemin-modified electrode was subjected to several physicochemical characterization studies using TEM, EDXS, Raman spectroscopy, UV–vis, and FTIR. The collective results revealed that the axial coordination bonding of amino functional groups of the Chit with sixth vacant site of the hemin is a key step for the ET process of the hemin-surface-confined carbon nanomaterial systems. Incorporating gold nanoparticles in the matrix improved the ET function through molecular wirings. As an electrochemical application, electrocatalytic reductions of dissolved oxygen and hydrogen peroxide in N₂-purged pH 7 PBS have been demonstrated. The new modified electrode showed about 300 mV reduction in the overpotential and two times increment in the reduction peak current signals over the respective unmodified electrode, GCE/Au nano–GMC–Chit. Extension of the new system toward sensor applications of dissolved oxygen and H₂O₂ without interference from common biochemicals was also demonstrated. Unlike the enzymatic systems, the hemin surface-confined carbon nanomaterial system showed redox stability over 30 days at room temperature.

4. EXPERIMENTAL SECTION

4.1. Chemicals and Reagents. Lyophilized hemin from Bovine (purity > 90%), GMC (purity > 99.95%, <50 nm pore size), GO—ethanol dispersed stock solution (GO, 5 mg mL⁻¹, ~80% carbon basis, flake size—0.5–2.0 µm, thickness—0.6–1.2 nm, purity—99%), mesoporous carbon-hydrophilic core (CM, 99.95% pure; particle size 5–50 µm) and mesoporous carbon-hydrophobic core (99.95% pure; particle size 5–50 µm), carboxylic acid-functionalized MWCNT (f-MWCNT; ~90% pure, size 7–15 nm × 0.5–10 nm), SWCNT (~70% pure on carbon basis, 0.7–1.1 nm diameter), and Chit were obtained from Sigma-Aldrich, USA. Standard gold solution (1000 mg L⁻¹) was purchased from SISCO Research Laboratories, India. Other basic chemicals of analytical grade were used without any purification. Nitrogen-purged (15 min) PBS (Na PBS) of ionic strength 0.1 M, pH 7 was used as a supporting electrolyte. Screen-printed carbon electrodes (SPCEs) were purchased from Zensor R&D, Taiwan.
4.2. Apparatus. Electrochemical experiments were carried out using a portable Biologic (USA) instrument with a three-electrode system comprising hemin chemically modified GCE (GCE/GMC@hemin—Chit) as a working electrode (0.0707 cm²), platinum as a counter electrode, and Ag/AgCl with 3M KCl as a reference electrode with a 10 mL working volume of N₂-purged pH 7 PBS. Raman spectroscopy analyses were carried out using AZILTRON, PRO 532, (USA) with a 532 nm laser excitation source. UV—vis analysis was done with a JASCO 4100 spectrophotometer by KBr method. A portable dissolved oxygen meter (HI9142) was purchased from Hanna Instruments, USA, was used to measure the dissolved oxygen concentration.

4.3. Fabrication of a Hemin Chemically Modified Electrode. First, the GCE (3 mm in diameter) was subjected to a series of chemical, mechanical, and electrochemical cleaning procedures, as reported in the literature.26 For the preparation of GCE/Au nano—GMC@hemin—Chit, the clean surface of GCE was first casted with Au³⁺ solution (5 µL; 1000 ppm), followed by drying at room temperature for 2 ±1 min. Then, 5 µL of GMC or other carbon nanomaterial—ethanol suspension (2 mg mL⁻¹) was casted and air-dried (2 ±1 min). Following this, 5 µL of hemin—NaOH (0.1 M) solution (2 mg mL⁻¹) was dropped on the surface of the above modified electrode (GCE/Au nano—GMC) and left to dry in an incubator (40 ± 2 °C) for 5 ± 1 min. As a last step, 5 µL of 0.5% Chit—acetic acid solution was overlayer-coated and air-dried for 2 ± 1 min. In total, within 18 ± 2 min, the modified electrode can be prepared. The modified electrode was potential-cycled in pH 7 PBS at a window −0.8 to 0.4 V versus Ag/AgCl for 20 scans at ν = 50 mV s⁻¹ (Scheme 1). During this process, the Au³⁺ ions were in situ reduced as Au nanoparticles and stabilized by the amino functional group of Chit in the modified electrodes (Scheme 1H,i).21

For Raman spectroscopic characterization, SPCE-coated GMC and Au nano—GMC@hemin—Chit were used. UV—vis spectroscopy measurements were carried out by stripping out the hemin site from the GCE-modified electrode by sonicating the electrode with 500 µL of pH 7 PBS followed by microfiltration. To perform a typical FTIR characterization, respective GCE-modified electrodes were peeled off with a doctor’s syringe needle (1 mm × 4 cm), mixed with KBr as a pellet, and subjected to the analysis. Electrocatalytic reduction of dissolved oxygen and H₂O₂ was carried out by exposing GCE/Au nano—GMC@hemin—Chit to saturated DO (10 ppm, measured by a standard DO meter) taken in N₂-purged pH 7 PBS and 500 µM H₂O₂.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsomega.8b00322.

Control CV response of Ni overlayer-coated GCE/Au nano—GMC@hemin, TEM—EDXS analysis result of GCE/Au nano—GMC@hemin—Chit, comparative Raman spectroscopy responses of GMC and GCE/Au nano—GMC@hemin—Chit, amperometric i−t responses of GCE/Au nano—GMC@hemin—Chit with its control (GCE/Au nano—GMC—Chit) for the detection of continuous spikes of 25 µM H₂O₂, calibration plot and interference effect in N₂-purged pH 7 PBS, and comparative stability CV response of GCE/Au nano—GMC@hemin—Chit (PDF).

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**Notes**

The authors declare no competing financial interest.

**ACKNOWLEDGMENTS**

The authors acknowledge the Department of Science and Technology—Science and Engineering Research Board (DST-SERB-EMR/2016/002818) Scheme. K.A. thanks the Council of Scientific and Industrial Research for the award of her senior research fellowship (09/844(0037)/2016 EMR-1). A.S.K. acknowledges National Taipei University of Technology for the support of distinguished visiting professor program.

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