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Design of screen-printed potentiometric platform for sensitive determination of mirabegron in spiked human plasma; molecular docking and transducer optimization†

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The integration of molecular modelling simulation and electrochemical sensors is of high interest. Herein, for the first time, a portable solid-contact potentiometric electrode was designed for the sensitive determination of mirabegron (MIR) in human plasma and pharmaceutical formulation. A two-step optimization protocol was investigated for the fabrication of an ion on sensing polymeric membrane. First, molecular docking was used for optimum ionophore selection. Calix[6]arene showed the highest affinity towards MIR with a better docking score (-4.35) and potential energy (-65.23) compared to other calixarene derivatives. Second, carbon nanotubes and gold nanoparticles were investigated as ion-electron transducers using a drop-casting procedure. Gold nanoparticle-based sensors showed better slope, potential stability, and rapid response compared to carbon nanotubes. The proposed solid contact sensors (V–VII) showed comparable sensitivity and ease of handling compared to liquid contact sensors (I–IV). The optimized gold nanoparticles sensor VII produced a Nernstian response over the range of 9.77×10^{-7} to 1×10^{-3} M with LOD of 2.4×10^{-7} M. It has also been used to determine MIR in its pharmaceutical formulation in the presence of a co-formulated antioxidant butylated hydroxytoluene and spiked human plasma. This would offer a feasible and economic platform for monitoring MIR in pharmaceutical preparation and biological fluids.

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Introduction

Overactive bladder (OAB) is a disorder characterized by urine urgency, with or without urinary incontinence, and nocturia. MIR is a beta-3 adrenergic receptor agonist that provides a better safety profile for OAB patients with advanced age and cognitive deficits. Recently, MIR has been approved for the treatment of neurogenic detrusor over-activity in children. It is chemically known as 2-(2-amino-1,3-thiazol-4-yl)-*N*-[4-[2[[(2*R*)-2-hydroxy-2-phenyl-ethyl]amino]ethyl]phenyl]acetamide (Fig. S1†). Few analytical methods were developed for the analysis of MIR such as chromatographic, 5 spectrophotometric, 6,7 and voltammetric methods. To the best of our knowledge, no potentiometric sensor has been reported for the determination of MIR.

Ion-selective electrodes (ISEs) have been widely used in pharmaceutical, biological fluids, and environmental analysis. 9,10 The ionophore is a key ingredient in membrane fabrication. It is responsible for the desirable selectivity in ISEs. Host-guest chemistry, represented by the ionophore-analyte interaction, accounts for better ionophore selection. It results in low limits of detection (LOD) and enhancement of sensitivities of the fabricated sensors.11 Calixarene derivatives are macrocyclic compounds with endless potential. They have large cavities which allow efficient binding with large ions. They are used as ionophores in sensors that detect nitrogen-containing ions like protonated amines and amino acid esters.11 Due to the lack of a potentiometric technique suitable for the assay of MIR in the examined matrices (formulation, and spiked human plasma), the selection of the optimum ionophore is of interest. Molecular modelling is a powerful alternative to predict and evaluate the binding between the host ionophores and the studied guest molecule. The binding energy could be assessed through docking score functions. Such calculations would help in selecting the optimum target molecule with minimum experimental runs and thus save time and effort.12

Great challenges were exerted to eliminate the inner filling solution to get a stable and reproducible response with long-

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term storage resulting in the development of solid-contact ISEs. Later, the application of screen-printed electrodes (SPEs)¹³⁻¹⁵ is considered a breakthrough. SPEs have extra advantages over other conventional solid-contact electrodes (coated wire electrodes and glassy carbon electrodes), as being cheap, disposable, easily handled, and stored with reproducible and reliable response in addition to their versatility and commercial availability. However, SPEs are susceptible to instability as other classical solid contact electrodes because of internal water layer formation and ion-to-electron transduction hindrance. Hence, to reduce the formation of the water layer, a hydrophobic conducting polymer layer was added on the surface of the solid contact and acted as an ion-to-electron transducer. Even though these conducting polymers decreased detection limits and signal drifts. 17

Another alternative to reduce water layer and increase signal stability is the use of nanoparticles. Although nanoparticles are not part of the direct electrode's sensing mechanism, they have been incorporated into the SPEs to offer higher signal, increase electrode stability, and eradicate the undesired water layer. Single-walled carbon nanotubes,18 multiwall carbon nanotubes (MWCNT),19 graphene,20 and most recently gold nanoparticles (GNP)17 are the most commonly used candidates. In the production of electrochemical sensors, GNP are known to be a suitable modifier for electrode surfaces. Due to their great quality such as tunable physiochemical properties, outstanding electrical conductivity, oxidation resistance, high stability, simple production, limited size distribution, excellent biocompatibility, capacity for surface modification, vast surface area, and excellent catalytic activities.21 Also, MWCNT have one of the simplest chemical compositions and atomic bonding configurations among nanomaterials. Their distinct structural, electrical, and mechanical characteristics make them a particularly appealing material for a wide range of applications.²²

The aim of the work was to optimize a sensitive potentiometric screen-printed electrode for MIR assay in its formulation matrix and spiked human plasma. A computer-aided molecular design platform was performed to study the binding characteristics of each ionophore with the drug. Then, the electrochemical performance of gold nanoparticles and carbon nanotubes as transducers was compared for better potentiometric stability, selectivity, and sensitivity.

2. Experimental

2.1 Chemicals and reagents

A pure sample of MIR was supplied by the Egyptian Drug Authority (EDA), Egypt. Its purity was found to be $99.99\pm0.612.$ A pure sample of butylated hydroxytoluene (BHT) was supplied by Oxiris Chemicals SA, Spain. Bladogra® tablets, batch number (AT02880322), were manufactured by APEX Pharma, Egypt, and labelled to contain 50 mg of MIR per tablet. High molecular weight polyvinyl chloride (PVC), 2-nitrophenyl octyl ether (oNPOE), potassium tetrakis(4-chlorophenyl)borate (K-TCPB), calix[4]arene (CX-4), calix[6]arene (CX-6), calix[8]arene (CX-8). Carbon screen-printed electrodes (SPE) 3 mm diameter were purchased from CH Instruments, Inc. (USA). Gold nanoparticles

(5 nm diameter), multi-wall carbon nanotube powder (MWCNT), tetra-butyl ammonium bromide, sodium hydroxide, xylene, orthophosphoric acid solution, and potassium dihydrogen orthophosphate were purchased from Sigma Aldrich (USA). Tetrahydrofuran (THF) was obtained from BDH (England). Sodium chloride, magnesium sulfate, ammonium chloride, potassium chloride, and calcium chloride were obtained from El-Nasr Pharmaceutical Chemicals Co. (Egypt). Human plasma was obtained from Holding Company for Biological Products and Vaccines, VACSERA (Egypt). Phosphate buffer (pH 7.4 ± 0.1) was prepared by adding 2.8 g of potassium dihydrogen orthophosphate in 700 mL of distilled water then pH was adjusted to 7.4 using 0.1 M sodium hydroxide solution. Volume was completed to 1000 mL with water.

2.2 Instrument and software

Jenway digital ion analyzer model 3330 (UK) with Ag/AgCl double-junction as reference electrode (Aldrich Chemical Co., Germany) was used to measure the potential difference. pH glass electrode (Jenway, UK) was used for pH adjustment. The Molecular Operating Environment software (MOE) version 2014 was used to study docking.

2.3 Standard solutions

MIR stock standard solution (1 \times 10⁻² M) was prepared in phosphate buffer pH 7.4. Working solutions of varying concentrations (1 \times 10⁻⁷ to 1 \times 10⁻³ M) were freshly prepared by serial dilutions from the stock solution using phosphate buffer pH 7.4 as a solvent.

2.4 Procedure

2.4.1 Fabrication of sensors

For the liquid contact membrane (LC) sensor I. 33.17 wt% PVC, 66.60 wt% NPOE, and 0.23 wt% K-TCPB (total 600 mg) were dissolved in about 6.00 mL THF. The solution was then poured into a Petri dish (5 cm diameter), the dish was covered with filter paper, and the solvent was allowed to evaporate overnight at room temperature, resulting in a master membrane with a thickness of \approx 0.1 mm.

For ionophore-doped membranes. The CX-4 membrane sensor II contained 32.79 wt% PVC, 66.60 wt% NPOE, 0.25 wt% K-TCPB, and 0.36 wt% CX-4. The CX-6 membrane sensor III contained 32.54 wt% PVC, 66.60 wt% NPOE, 0.25 wt% K-TCPB, and 0.64 wt% CX-6. The CX-8 membrane sensor IV contained 32.21 wt% PVC, 66.60 wt% NPOE, 0.25 wt% K-TCPB, and 0.94 wt% CX-8. The ion exchanger and the ionophore were doped in a molar ratio of 1:2 in all membranes.

For solid contact membranes. Preparation of multiwall carbon nanotubes: the solution dispersion method was used¹⁶ to prepare carbon nanotubes. 10.0 mg of carbon nanotubes powder were dispersed in 1 mL xylene by ultra-sonication for 5 min. 95.0 mg PVC was dissolved into 3.00 mL of THF using 0.2 mL NPOE as a plasticizer. The THF solution was mixed with carbon nanotubes dispersion and sonicated at least for 20 min to obtain a uniform suspension.

For screen-printed sensor V (without transducer): 10.0 μ L of the CX-6 membrane before dryness was applied on a carbon SPE and allowed to evaporate overnight at room temperature.

For multi-wall carbon nanotubes screen printed (MWCNT SP) sensor VI: MWCNT solid layers were prepared by drop-casting 10.00 μ L of MWCNT dispersion on a carbon SPE and allowed to evaporate overnight at room temperature. Then, 10.0 μ L of the CX-6 membrane before drying was applied to a carbon SPE and allowed to evaporate overnight at room temperature.

For gold nanoparticles screen printed, (GNP SP) sensor VII: GNP solid layers were prepared by drop casting 10.00 μL of GNP dispersion on a carbon SPE and allowed to evaporate overnight at room temperature. Then, 10.0 μL of the CX-6 membrane before drying was applied to a carbon SPE and allowed to evaporate overnight at room temperature.

SPEs sensors V, VI, VII were kept in MIR 1×10^{-4} M stock standard solution for 24 hours and stored in the same solution when not in use.

2.4.2 Liquid contact (LC) ion selective electrode assembly. A disk of about 8 mm in diameter was cut from the master membrane using a cork borer and then pasted using THF to a transposable PVC tip. Equal volumes of 1×10^{-4} M MIR and 1×10^{-4} M KCl (prepared in phosphate buffer, pH 7.4) were mixed and used as an internal reference solution. An Ag/AgCl wire was immersed in the internal reference solution as an internal reference electrode. The sensor was conditioned by soaking in a 1×10^{-4} M MIR stock standard solution for 24 hours and stored in the same solution when not in use. The same procedure was applied to the ionophore-doped

2.4.3 Screen-printed ion selective electrodes (SPEs) assembly. SP sensors V, VI, VII were fabricated as mentioned under 2.4.1, and then soaked in MIR 1 \times 10⁻⁴ M stock standard solution for 24 hours.

2.5 Docking study

membranes.

The docking study was performed using the Molecular Operating Environment software (MOE) version 2014. The hosting ionophores (CX-8, CX-6, and CX-4) and the guest molecule (MIR) were prepared for docking using 3D protonation of the structure at pH 7.4. The conformational analysis using a systematic search was scanned then the lowest energetic conformer was chosen.

2.6 Sensors potential stability and water layer test

The water layer test, 20 was performed by measuring the response of the screen-printed electrodes (SP sensors V, VI, VII) for three hours. First hour in 1 \times 10 $^{-3}$ M MIR. Second hour in 1 \times 10 $^{-2}$ M tert-butyl ammonium bromide and again in 1 \times 10 $^{-3}$ M MIR for the third hour. The signal drift was monitored by recording the emf values every 2 min during the first hour.

2.7 Sensors calibration

Each sensor was separately conjugated with a double-junction Ag/AgCl reference electrode, calibrated by being immersed in MIR drug solutions (1 \times 10⁻⁷ to 1 \times 10⁻³ M). Solutions were

allowed to equilibrate while stirring by a magnetic stirrer until achieving a constant reading of the potentiometer. The electromotive forces (emf) were recorded to within ± 1 mV for LC sensor I, ± 0.5 mV for SP sensor II, ± 0.3 mV for MWCNT SP sensor III, and ± 0.3 mV for GNP SP sensor IV. Calibration graphs were plotted that related the recorded emf from the seven proposed sensors *versus* the $-\log$ molar concentrations of MIR. Regression equations were calculated for the seven proposed sensors. The electrochemical performance of the proposed sensors was evaluated according to the IUPAC recommendations.²³

2.8 Effect of pH on gold nanoparticles screen printed sensor VII

The influence of pH on the response of the most stable GNP SP sensor VII was studied over a pH range of 2–10 using the 1 \times 10⁻⁴ M and 1 \times 10⁻³ M MIR solutions. At each pH value, the potential was recorded.

2.9 Selectivity of gold nanoparticles screen printed sensor VII

The degree of foreign substance interference with MIR using GNP SP sensor VII was studied using the separate solutions method (SSM). The potential response of the sensor was recorded in the presence of inorganic-related compounds and in the presence of the main excipient BHT and co-administered drug solifenacin. The potentiometric selectivity coefficient (K^{Pot}) was calculated.

2.10 Direct potentiometric determination of mirabegron

2.10.1 In the pharmaceutical formulation. Ten tablets of Bladogra® were accurately weighed, their average weight was calculated, and then finely powdered. An amount equivalent to 3.96 mg of MIR was accurately weighed, transferred into a 100 mL volumetric flask, and diluted in a 70 mL phosphate buffer (pH 7.4). Sonication was done for 15 minutes till complete dissolution, then the volume was completed to the mark with buffer. The concentration of the resulting solution is claimed to be 1×10^{-4} M MIR. GNP SP sensor VII in conjunction with the double-junction Ag/AgCl reference electrode was immersed in the prepared solution. The resulting potentials were recorded, and the relevant concentrations were determined using the sensor's regression equation.

2.10.2 Spiked human plasma. One mL of MIR-free human plasma and aliquots of MIR working standard solutions (1 \times 10⁻⁶ to 1 \times 10⁻² M) were added into a series of 10 mL volumetric flasks. The volumes were completed to the mark with phosphate buffer pH 7.4 to yield final respective concentrations of 9.77 \times 10⁻⁷ M, 1 \times 10⁻⁵ M, 6.25 \times 10⁻⁵ M, 1 \times 10⁻⁴ M, and 1 \times 10⁻³ M of MIR. Quality control (QC) samples (low-quality control (QCL); 2.9 \times 10⁻⁶ M, medium quality control (QCM); 4 \times 10⁻⁴ M and high-quality control (QCH); 8 \times 10⁻⁴ M) were prepared using the same procedure. GNP SP sensor IV in conjunction with the double-junction Ag/AgCl reference electrode was immersed in the prepared solutions without matrix pre-treatment. The resulting potentials were recorded.

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3. Results and discussion

SPEs provide the choice to construct eco-sustainable, portable, and quick analytical procedures for detecting analytes in different matrices.²⁵ The proper selection of the ionophore and the transducer have a synergic effect on the selective and sensitive determination of the analyte. As a result, the current study introduces a novel solid-contact sensor for measuring MIR in pharmaceutical formulations and spiked human plasma. The study investigated the impact of various ionophores on the designed sensors' response stability, sensitivity, and selectivity.

3.1 Membrane fabrication

A polymeric PVC membrane was used to provide consistent support. NPOE was incorporated as a plasticizer to provide membrane malleability. The strongest basic pK_a value of MIR is 9.62, thus it behaves as a cation in an acidic medium. Therefore, an LC-ISE was fabricated for MIR determination with a cation exchanger. It offered anionic sites inside the membrane matrix to increase conductivity and reduce the interference caused by lipophilic anions coextraction (Donnan's effect).26 A preliminary

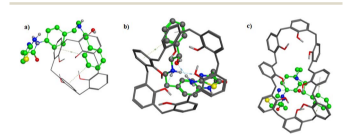


Fig. 1 Three-dimensional docking of mirabegron in (a) calix[4] arene, (b) calix[6]arene, and (c) calix[8]arene.

study was conducted using LC sensors to investigate the effect of the cation exchanger type on the potentiometric performance. Using sodium tetra phenyl borate (NaTPB) resulted in a weak Nernstian slope (43.202 \pm 0.23). Upon replacing NaTPB with a more lipophilic and stable ion exchanger, potassium tetrakis(chlorophenyl)borate (K-TCPB), enhancement of the Nernstian slope was observed.

3.2 Molecular docking

All ionophores have an easily well-defined cavity for molecular recognition. The structure of these ionophores is flexible allowing the binding sites to have an optimized binding conformation with minimal energy.26 The selectivity of ionophores depends on the cavity size and the conformation of the molecule binding strength of these receptors to the target ion via cation interactions.27 Our study studied calixarene derivatives, such as CX-4, CX-6, and CX-8, to recognize MIR selectively. They can develop separate interactions with guest molecules, such as hydrogen bonding, π - π , and cation-electrostatic interactions. The additional lipophilic moieties are compatible with the solvent mediator, and their polar functional groups generate a basket shape suitable for molecular recognition.28

Molecular docking was conducted to study the stability of the inclusion complex of MIR with CX-4, CX-6, and CX-8. A docking study was performed using MOE (2014) software package, and the 3D-crystal structures of different calixarenes were obtained from their corresponding files on pdb. The hosting ionophores and the guest molecule (MIR) were prepared for docking using 3D protonation of the structure scanning the conformational search analysis to select the global minimum energy conformers. A molecular docking study illustrated a well-fitted MIR conformer within the cavity of CX-6 than CX-4 or CX-8. MIR demonstrated a hydrogen bonding along with π - π and π -alkyl interactions with CX-6, unlike CX-4

Table 1 Response characteristics of the investigated sensors for mirabegron determination

Parameter	LC sensor I	LC sensor II	LC sensor III	LC sensor IV	SP sensor V	MWCNT SP sensor VI	GNP SP sensor VII
Slope (mV per decade)	50.398 ± 306.15	55.761 ± 328.71	58.886 ± 367.55	556.235 ± 339.85	50.207 ± 1.49	52.233 ± 0.09	57.513 ± 0.06
Intercept (mV)	306.15	328.71	367.55	339.85	319.73	350.54	414.02
Correlation	0.9997	0.9994	0.9999	0.9997	0.9996	0.9996	0.9998
coefficient (r)							
Concentration	1.56×10^{-5}	3.91×10^{-6}	1.95×10^{-6}	3.91×10^{-6}	3.91×10^{-6} to 1 \times	9.77×10^{-7}	9.77×10^{-7}
range (M)	to 1×10^{-3}	10^{-3}	to 1×10^{-3}	to 1×10^{-3}			
Working pH range	6-8	6-8	6-8	6-8	6-8	6-8	6-8
Response time (s)	25	25	25	25	7	5	2
Stability (weeks)	4	4	4	4	4	4	4
Average accuracy ^{a} \pm S.D.	100.11 ± 0.603	99.21 ± 0.782	100.25 ± 0.182	100.33 ± 0.381	$\textbf{100.09} \pm \textbf{1.668}$	$\textbf{100.74} \pm \textbf{1.164}$	101.96 \pm
							0.205
Repeatability b (RSD%)	0.217	0.251	0.625	0.812	0.533	0.532	0.274
Intermediate precision ^c	0.752	0.812	0.823	0.798	1.014	1.159	0.804
(RSD%)							
$LOD^{d}(M)$	1×10^{-5}	5.3×10^{-6}	8.9×10^{-7}	5.3×10^{-6}	6.1×10^{-7}	4.0×10^{-7}	2.4×10^{-7}

^a Average of three determinations (n = 9). ^b Repeatability: the intraday precision (n = 9), average of three concentrations repeated three times within the day. c Intermediate precision: the inter-day precision (n = 9), average of three concentrations repeated three times on three consecutive days. ^d LOD (limit of detection) was measured by interception of the extrapolated arms of Fig. 5 and S2.

or CX-8 which forms only weak π - π interaction with MIR (Fig. 1). Additionally, the docking score (s) and potential energy of the molecular system (E) values confirmed that the CX-6 forms a more stable inclusion complex with mirabegron than CX-4 or CX-8. The (s, E) values were (-4.35, -65.23), (-4.19, -62.30), and (-4.15, -64.19) for CX-6, CX-4, and CX-8, respectively. The highest docking score value was recorded for CX-6 MIR complex showing the best stability. Results from the computational model agreed with the experimental findings, where sensor III (CX-6) showed the best slope and sensitivity among the examined sensors (Table 1). Therefore, CX-6 was chosen for further experiments.

3.3 Screen printed electrodes using calix[6]arene membrane

Three disposable solid contact screen-printed ISEs were fabricated for on-spot monitoring (Fig. 2). They were used for determination of MIR utilizing the CX-6 doped membrane as the sensing membrane. SP sensor V contained only the CX-6 membrane. MWCNT SP sensor VI was first doped with a thin interlayer of carbon nanotube then CX-6 membrane was applied. SP GNP sensor VII was first doped with a thin interlayer of gold nanoparticles then CX-6 membrane was applied. The signal drifting experiment was carried out via chronopotentiometric measurements performed using open circuit potential. It was tested by taking emf data every 2 minutes for an hour for an average of three electrodes for each SP sensor. Both MWCNT SP sensor VI and GNP SP sensor VII showed improvement in stability and minimal signal drifting of approximately 0.7 mV h⁻¹ and 0.5 mV h⁻¹ only when compared to SP sensor V which showed signal drifting of about 8.3 mV h⁻¹ (Fig. 3). These findings demonstrated that adding nanoparticles to an electrode improves both electrode stability and signal drifting significantly.

3.4 Sensors potential stability and water layer test

An increase in potential was observed upon carrying the measurements in a 1×10^{-2} M solution of a cationic structurally relative compound (tetra butyl ammonium bromide). This increase is due to the substitution of MIR in the water film

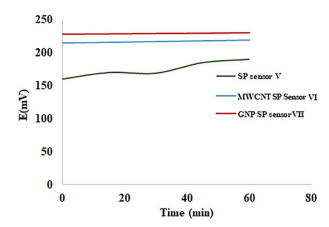


Fig. 3 Signal drift for screen printed sensor V, multi-wall carbon nanotubes screen printed sensor VI, and gold nanoparticles screen printed sensor VII. Measurements were recorded in 1×10^{-3} M MIR.

formed between the PVC sensing membrane and the carbon layer of the screen-printed substrate (SP sensor V), the MWCNT layer (MWCNT SP sensor VI), and the GNP layer (GNP SP sensor VII) by tetra butyl ammonium bromide. The final replacement of the test solution $(1 \times 10^{-2} \text{ M})$ tetra butyl ammonium bromide) by the original 1×10^{-3} M MIR solution exhibited an almost null potential drift GNP SP sensor VII compared to MWCNT SP sensor VI and SP sensor V. The data presented in Fig. 4 clearly demonstrated that GNP showed comparable results with MWCNT. Thus, it proved that GNP offered the required hydrophobicity for the water layer test. Accordingly, a gold nanoparticle screen-printed electrode (GNP SP sensor VII) was employed for all the measurements. Hence, it was used to determine the pure drug; MIR, in Bladogra® tablets and spiked human plasma.

3.5 Sensors calibration and response time

The response properties of the proposed sensors were measured over a four-week time frame and the results are shown in Table 1. Calibration curves were plotted relating the recorded electrode potential from the proposed sensors *versus* the —log molar

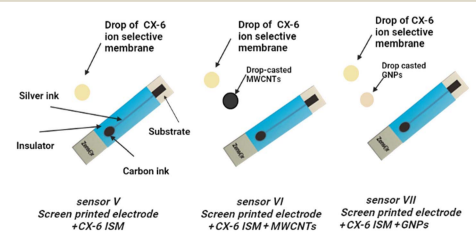


Fig. 2 Schematic representation of screen printed electrodes (V–VII)

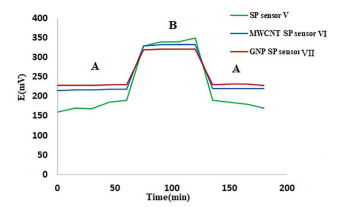


Fig. 4 Water layer test for screen printed sensor V, multi-wall carbon nanotubes screen printed sensor VI, and gold nanoparticles screen printed sensor VII. Measurements were recorded in 1 \times 10 $^{-3}$ M MIR (A) and 1 \times 10 $^{-2}$ M tetra butyl ammonium bromide (B).

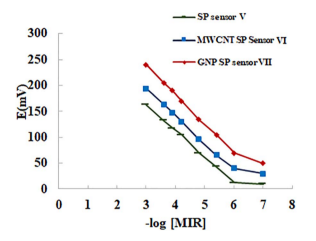


Fig. 5 Profile of the potential in mV/–log MIR molar concentration using sensor V, multi-wall carbon nanotubes screen printed sensor VI, and gold nanoparticles screen printed sensor VII.

concentrations of MIR as shown in Fig. 5 and S2.† The slope of the curves using the calibration graph's linear portion was calculated as presented in Table 1. Over the course of four weeks, the sensors were tested and found to have reliable and accurate results. The quick response time of ISEs is still a key characteristic that helped the use of these electrodes for quality control since it reduces the amount of required analysis period. The recommended time for the sensors to obtain stable potential readings after raising the MIR level 10-fold was found to be 25 seconds for the LC sensors, 5 seconds for the SP sensor V, 4 seconds for the MWCNT SP sensor VI, and approximately 2 seconds for GNP SP sensor VII.

3.6. Electrochemical impedance spectrometry (EIS) measurements

Electrochemical impedance spectroscopy (EIS) was conducted to characterize the GNPs transducer layer. The Nyquist plot $(-Z'' \nu s. Z')$ for GNP SP sensor VII and SP sensor V was obtained using the frequency range (100 kHz to 0.1 Hz) as shown in Fig. 6. The

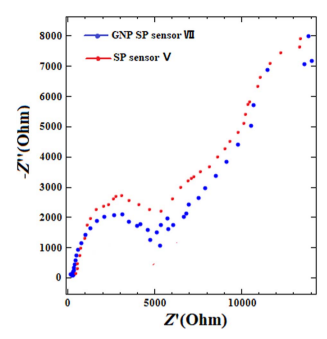


Fig. 6 Nyquist plots of GNP SP sensor VII and SP sensor V.

EIS results were fitted to equivalent Randles circuit and the double layer capacitance ($C_{\rm dl}$) in GNP SP sensor VII and SP sensor V was calculated and found to be 1.14 μF and 173 nF, respectively. While the charge transfer resistance measured for the GNP SP sensor VII electrode was 4.19 $k\Omega$ and for the SP sensor V electrode was 8.13 $k\Omega$. These results confirmed that GNPs incorporation increased the electrode capacitance and reduced the transfer resistance which can explain the stability of the interfacial potential and improved charge transfer at the interface.

It should be noted the overall electrochemical performance of the GNP SP sensor was achieved through (i) the use of potassium tetrakis(chlorophenyl)borate as a lipophilic cationic exchanger which offered the required ion exchange sites for the cationic drug (MIR). (ii) The optimized selection of CX6 as an ionophore that enabled molecular recognition for optimal binding with minimal energy. (iii) The incorporation of GNP as a modifier for electrode's surface. Such efforts resulted in optimum sensor's selectivity and sensitivity as well as signal stability.

3.7 Effect of pH on gold nanoparticles screen printed sensor VII

The pH of the working solutions is a very critical factor to achieve complete drug ionization. Fig. 7 showed the effect of pH on the response of the most stable electrode (GNP SP sensor VII) over a pH range of 2–10 to achieve the optimum experimental conditions. In solutions with pH values of 6–8, where MIR was entirely ionized and detected, the sensor response was clearly stable. This finding, together with the fact that the MIR pK_a value is 9.62, justifies our decision to employ phosphate buffer pH 7.4 for MIR determination.

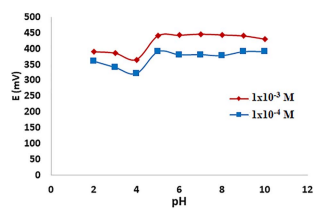


Fig. 7 Effect of pH on the response of gold nanoparticles screen printed sensor VII using the 1×10^{-4} M and 1×10^{-3} M mirabegron solutions [working pH range: 6-8].

3.8 Gold nanoparticles screen printed sensor VII selectivity

The potentiometric selectivity coefficient was calculated using the following equation:

$$\log(K^{\text{Pot}}) = (E_{\text{MIR}} - E_{\text{I}}/S)$$

where $K^{\rm Pot}$ is the potentiometric coefficient, $E_{\rm I}$ is the potential measured in a 10^{-3} M solution of interferent ion and $E_{\rm MIR}$ is the potential measured in a 10^{-3} M solution of MIR solution. S is the slope of the calibration plot in MIR. The estimated selectivity coefficients revealed that the proposed sensor is highly selective to MIR without interference from the interfering species (Table 2).

3.9 Potentiometric determination of MIR

3.9.1 In pharmaceutical formulation. GNP SP sensor VII was used to determine MIR concentration in Bladogra® tablets. The average of three determinations showed accurate and precise percentage recoveries, Table 3 indicating the selectivity of the sensor to detect MIR in pharmaceutical formulations without any interference from the excipients present in the formulation.

3.9.2 In spiked human plasma. GNP SP sensor VII was dipped in conjunction with the double-junction Ag/AgCl

Table 2 Potentiometric selectivity coefficients of GNP SP sensor VII using separate solution method (SSM)

Interfering compound (10 ⁻³ M)	$\log \mathit{K}^{^{\mathrm{Pot}a}}$
внт	1.87×10^{-3}
Solifenacin	1.34×10^{-3}
MgCl_2	2.24×10^{-3}
NaCl	1.48×10^{-3}
KCl	2.35×10^{-3}
CaCl ₂	2.14×10^{-3}
NH ₄ Cl	2.24×10^{-3}

^a Average of three determinations.

Table 3 Determination of mirabegron in Bladogra® tablets by the gold nanoparticles screen printed sensor VII

	Gold nanoparticles screen printed sensor VII		
Pharmaceutical formulation	Found ^a (%)	R.S.D. (%)	
Bladogra® tablets, batch no. AT02880322	99.33	1.01	
a Average of three determinations			

^a Average of three determinations.

reference electrode in MIR solutions (9.77 \times 10⁻⁷ to 1 \times 10⁻³ M) spiked with 1.0 mL human plasma. The potential was recorded without prior extraction or matrix pre-treatment procedures. The electrode succeeded to detect MIR in plasma down to a concentration level of 9.77×10^{-7} M which is below multiple dose MIR C_{max} (461 ng mL⁻¹ equivalent to 1.16 × 10⁻⁶ M).30 MIR is extensively metabolized in the liver through the CYP450 system and has a broad safety profile. However, plasma monitoring and dose adjustment are required in patients who are poor metabolizers of CYP2D6 enzyme or taking other medications that are well-known potent inhibitors (such as amiodarone, fluoxetine, and diphenhydramine) or substrates (such as metoprolol, carvedilol, and amitriptyline) of the CYP2D6 enzyme.31 To assess the accuracy and reproducibility of the proposed sensor in quantifying MIR in plasma, quality control (QC) samples (QCL, QCM, and QCH) were analyzed three times within the day and on three consecutive days (Table 4).

Table 4 Precision and accuracy for determination of mirabegron in spiked human plasma by gold nanoparticles screen printed sensor VII

Nominal concentration	Interday (n	= 9)	Intraday $(n = 9)$	
(M)	Accuracy	% RSD	Accuracy	% RSD
QCL (3×10^{-6})	97.79	0.79	97.66	0.38
QCM (4×10^{-4})	100.58	1.07	99.77	0.40
QCH (8×10^{-4})	101.24	1.02	101.39	1.02

Table 5 Statistical comparison of the results obtained using the proposed gold nanoparticles screen printed sensor VII as compared with the reported method for mirabegron determination in its pure powdered form

Item	GNP SP sensor VII	Reported method ^a
Mean	100.00	99.99
S.D.	0.633	0.612
Variance	0.401	0.375
n Student's <i>t</i> -test ^b F test ^b	7 0.027 (2.228) 1.069 (6.16)	5

^a HPLC method⁴ using C₁₈ column (150 mm \times 3.9 mm; 5 μm) and phosphate buffer pH 5: acetonitrile (50:50 v/v) as a mobile phase at flow rate 1 mL min⁻¹ and UV detection at 249.0 nm. ^b Corresponding tabulated values for t and F (at p = 0.05).

Table 6 Penalty points for greenness assessment of the proposed potentiometric method using gold nanoparticles screen printed sensor VII compared to the reported method

	Penalty points		
Hazard	Proposed method	Reported method ^{4a}	
Reagents			
Tetrahydrofuran	6	_	
Acetonitrile	_	15	
Potassium dihydrogen phosphate	0	0	
Instruments			
Energy	0	1	
Occupational hazard	0	0	
Waste	6	8	
Total penalty points	12	24	
Analytical eco-scale total score	88	76	

^a HPLC method using C_{18} column (150 mm \times 3.9 mm; 5 μ m) and buffer pH 5: acetonitrile (50: 50 v/v) as a mobile phase at flow rate 1 mL min⁻¹ and UV detection at 249.0 nm.

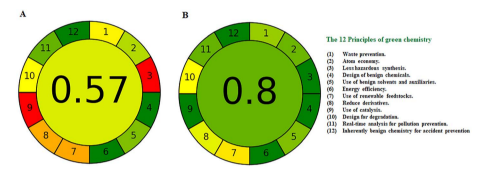


Fig. 8 Agree green profile assessment of the reported method (A) [14] and (B) the developed potentiometric method for determination of mirabegron.

3.10 Statistical data analysis

The results obtained by the GNP SP sensor VII were compared with those obtained by applying the reported method for the determination of MIR pure sample. The *t* and *F* test values were used to compare the mean and variance between the two compared methods where no significant difference in the results was found (Table 5).

3.11 Greenness assessment of the proposed method

The green index of the proposed method was assessed using two metrics: analytical eco-scale and analytical greenness metric procedures. Analytical eco-scale is a semi-qualitative tool based on giving penalty points to each element of analytical methodology subtracted from a base of 100. Penalty points were calculated for the proposed electrochemical method and the reported method for the determination of MIR.⁴ The proposed method showed a better score (eco-score = 88) than the reported method (eco-score = 76) due to the use of a large amount of acetonitrile organic solvent (Table 6). AGREE metric³² was also used to evaluate the method greenness. The full AGREE concept circulates around the twelve principles of green analytical chemistry (GAC) to allow thoroughness of input and clarity of output.³² Each input principle is converted into a score

between 0 and 1. The final score is calculated from the assessment of all the principles. Our proposed method showed a greater score than the reported method⁴ as shown in Fig. 8.

4. Conclusion

The scope of this research was to develop an excellent green ionselective electrode methodology, with good selectivity and low limit of detection, for the determination of MIR. Ionophore selection was the key measure of this work. Three calixarenes (4), (6), and (8) were consequently tested via molecular docking. Calixarene (6) formed the most stable complex with MIR compared to (4) or (8). Consequently, seven different sensors (four liquid contact and three solid contact SPEs electrodes) were designed. The plain solid contact SPE suffered from the formation of a water layer and then substantial drift. Hence, carbon nanotubes and gold nanoparticles were trialed as transducers. They showed negligible signal drifting and steady potential using the water layer test. The best detection limit $(2.4 \times 10^{-7} \text{ M})$ was obtained with the GNP SP sensor VII. Our proposed method using GNP SP sensor VII revealed to be an excellent green method for MIR detection in its formulation and in spiked human plasma without the need of using risky and hazardous solvents, sample pretreatment, or sophisticated equipment.

Author contributions

Aya A. Mouhamed: methodology, investigation, formal analysis, visualization, writing – original draft. Basma M. Eltanany: supervision, validation, writing – review & editing. Nadia M. Mostafa: conceptualization, project administration, writing – review & editing, supervision. Tamer A. Elwaie: resources, data curation, formal analysis. Ahmed H. Nadim: methodology, data curation, formal analysis, validation, resources, writing – review & editing.

Conflicts of interest

The authors state that they have no known competing financial interests or personal ties that could have influenced the research presented in this study. There is no conflict of interest to declare.

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