Electrochemical Dopamine-Imprinted Sensor Based on Tio₂ Nanoparticles and Polypyrrole-Chitosan Composites Modified Glassy Carbon Electrode

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Received 19/06/2023; accepted 01/09/2023 https://doi.org/10.4152/pea.2024420605

Abstract

In this study, a composite material containing PY and CTS, as functional material, and dopamine, as template, was electrodeposited onto a GCE surface modified with NP from TiO₂. The prepared MIP sensor was characterized by CV, SEM, FTIR and UV-vis techniques. The developed MIP matrix was used for dopamine detection, using DPV method. Diverse analytical parameters were optimized, such as: monomer, template, SE concentrations, electropolymerization cycles, pH medium, incubation time and scan rate. In the best conditions, the sensor response to dopamine was linear, in the concentration range from 1 x 10⁻⁶ to 1 x 10⁻⁵ mol/L⁻¹. LOD was about 2.81.10⁻⁷ mol/L⁻¹, and sensitivity was 3.202 μ A/L/mol⁻¹/cm⁻². Furthermore, the proposed MIP sensor showed good selectivity, repeatability, reproducibility and stability. Also, it was successfully applied for dopamine determination in urine samples.

Keywords: CV; dopamine; DVP; FTIR; MIP; PPY-CTS composite; SEM; TiO₂; urine samples; UV-vis.

Introduction•

Dopamine was classified as the most abundant catecholamine and special neurotransmitter, and plays a critical role in human health related to the central nervous, cardiovascular, renal and hormonal systems [1]. In addition, dopamine extends its role to control stress responses, consciousness, information flow and attention span, learning, sleep-wake cycle, motivation, motions and memory formation, i.e., it rules brain-body integration [2-4]. A deficiency or rise in dopamine concentration leads to many mental diseases [5]. For these reasons, it is necessary to develop a sufficiently sensitive method for dopamine detection in real samples. Since the 1950s, much effort has been made on developing new methods for dopamine

[•] The abbreviations and symbols definition lists are in pages 469-470.

determination. Several electrochemical biosensors were tested for dopamine detection using NP and conducting polymers, as recognition elements [6, 7]. During recent years, a wide range of metal NP has been widely used for the fabrication of sensors. They have many superb properties like large surface-to-volume ratio, good electrical features, strong adsorption ability, high surface reaction activity, smaller size and good surface characteristics [8]. These wonderful properties are helpful for biomolecules immobilization. TiO₂ has been used as a model transition metal, due to its simple electronic configuration, with empty conduction and filled valance band [9]. Additional application of these molecules as a surface modifier, in electrochemistry [10-12] and sensor technology, [13-16] has increased due to their suitable and useful surface properties for electrodes surface modification. The successful use of NP in sensor technology area is stimulated by the addition of conductor polymers. Among existent conducting polymers, PPY and its derivatives play a leading role, because of their versatile applicability and wide variety of molecular species that are covalently linked to PY groups [17].

As biocompatible polymer, CTS has many properties such as excellent filmforming ability, remarkable biocompatibility, nontoxicity and high mechanical strength [18, 19]. Besides, since it has many amino and hydroxyl groups, it has been widely used as a modifier.

Recently, due to obvious advantages such as low cost, physical robustness, thermal stability and easy preparation over biological receptors and other functionalized materials, MIP have become a competitive tool in the field of molecular recognition. The mechanism of molecular recognition between target molecule and MIP material comes from the field of immunology, and it was inspired by the recognition between antigen and antibody. The idea originated with Breinl and Haurowitz, and dates back to 1930 [20]. The first successful work using imprinted polymeric material was performed in 1973, employing silica particles imprinted with D-glyceric acid and D-mannitol [21].

The principle of this imprinted technique consists in creating a pre-polymerization complex between the monomer function and the target molecule, through Van der Waals or H bonds. A cross linker is used early in the polymerization stage, to form a three-dimensional cross-linked network, in which the target molecule is trapped by interfacial interactions established during the pre-polymerization complex step. After this process, the target molecule is extracted from the matrix, leaving a cavity with spatially oriented functionalities in the cross-linked polymer network [22].

In fact, electrochemistry has often received attention as a means of preparing MIP for developing high performance electrochemical sensors [23-26].

The present study aimed to complete the benefits given by biomolecules immobilization strategy used for biosensors elaboration. In addition, a major advantage was established by analytical and electroanalytical methods applied to the determination of biological and environmental analyses [27]. Adopted electrochemical sensitive methods included voltammetric and impedimetric measurements to evaluate the biosensor electrochemical response [25]. Additionally, this work endeavored to incorporate the benefits of MIP technique and electrochemical sensors technology. A new MIP electrochemical sensor was developed based on GCE modified with NP from TiO₂ and PPY-CTS composite

for dopamine detection. Electrodeposition parameters optimization and the MIP sensor characterization were investigated.

Experimental

Chemicals

Commercial nanopowder of TiO₂ was obtained from Fluka. PY was purchased from Merck. CTS, H₂SO₄, CH₃COOH, Glu, dopamine, AA and UA were supplied by Sigma Aldrich, and used as-received. PBS with various pH values was prepared using K₂HPO₄ and KH₂PO₄ obtained from Merck. K₄(Fe(CN)₆ and K₃(Fe(CN)₆) were purchased from Fluka chemika. Ultra-pure water was used for the solutions preparation and modified electrode rinsing.

Apparatus and instruments

Electrochemical experiments were carried out with a three-electrode system consisting of: GCE (3 mm diameter), Pt wire and Ag/AgCl (3 M KCl) as working, auxiliary and reference electrodes, respectively. CV and DPV were carried out using an Autolab PGSTAT 320 N potentiostat equipped by Nova version 1.5 software. All electrochemistry experiments were performed at room temperature. The solutions pH were measured with a pH meter (3505 JENWAY).

The morphology of the different matrices deposited on the carbon screen printed electrode surface was characterized using JEOL JSM 7100F SEM. Spectrum data were collected with FTIR spectroscopy, using a Perkin Elmer 1600 FTIR spectrometer. UV-vis measurements were recorded by 67 spectrophotometer models 6705 UV-vis JENWAY, after the deposited layers redissolution in a DMF/water mixture.

Preparation of the modified working electrode

Prior to use, GCE was polished with 0.30 and 0.05 mm alumina powder, employing a polishing cloth, rinsed with water, and then ultrasonicated in ethanol and distilled water, for five min, to remove adsorbed alumina particles on its surface. Afterwards, the polished GCE was electrochemically cleaned by cycling the potential scan from -0.2 to 1 V, in 0.5 mol/L⁻¹ H₂SO₄, until a stable CV was obtained.

Finally, GCE surface was dried under a stream of high purity nitrogen, for further use. The modification process consisted in the electrodeposition of a fresh TiO_2 aqueous solution (5.10⁻³ mol/L⁻¹) on the GCE, using CV from -1 to 1 V, for 5 cycles, at a scan rate of 50 mV/s. Then, the electrode was rinsed with distilled water and dried in air.

After the solution was deoxygenated by bubbling nitrogen gas, for 20 min, the MIP was built by electropolymerization of 5.10^{-3} mol/L⁻¹ PY and 1 mg/mL CTS in 0.1 mol/L⁻¹ CH₃COOH as SE, with 10^{-3} mol/L⁻¹ dopamine as template on the modified GCE/TiO₂ electrode, using CV, in the potential range from -1 to 1 V, for 10 cycles, at a scan rate of 50 mV/s.

The recognition sites in MIP film were formed after the template removal from the membrane. Dopamine extraction was conducted by CV in 0.1 mol/L⁻¹ PBS (pH 7).

NIP-modified electrode was also prepared under the same protocol, but without adding dopamine. Every imprinting and elution procedure are illustrated in Fig. 1.



Figure 1: Schematic representation of the fabrication procedure for GCE/TiO₂/(PPY-CTS) MIP.

Results and discussion

Preparation of the GCE/TiO₂ modified electrode

Fig. 2 illustrates electrodeposition of TiO_2 layer on the GCE surface by CV, in the potential range from -1 to 1 V, for 5 cycles, at a scan rate of 50 mV/s. As a result, the initial oxidation-reduction cycle shows a large and irreversible anode wave with a potential peak at 0.35 V, revealing the onset of TiO_2 oxidation on GCE surface. This peak increased until stabilization, which explains the fixation of the TiO_2 deposit layer on the working electrode surface.



Figure 2: CV of the TiO_2 electrodeposition layer on the GCE surface at a scan rate of 50 mV/s.

Electrodeposition of MIP PPY-CTS

During the polymerization process, it was seen that the oxidation peak of PPY-CTS was at 0.5 V. Hence, the reduction peak was at -0.1 V. The decrease in the oxidation peak was due to the deposition of the conducting PPY-CTS layer on the GCE/TiO₂ surface. The reduction peak is assigned to the polymer degradation [28]. For the MIP structure, dopamine template molecules had a major effect on the electroactivity of the deposit sensitive layer. The absence of an oxidation peak corresponding to the dopamine template during the formation of the MIP from PPY-CTS demonstrates the efficient blocking of dopamine electrooxidation by this barrier film. The polymer film development and growth is shown in Fig. 3.



Figure 3: Electrodeposition of PPY-CTS at GCE/TiO₂ by CV, at scan rate of 50 mV/s, using CH₃COOH as SE, containing: (black) 5.10^{-3} M PY + 1 mg/mL CTS; and (red) 5.10^{-3} M PY + 1 mg/mL CTS + 10^{-3} M dopamine.

The film oxidation revealed that dopamine was hosted by the polymer [29]. Dopamine molecules, which are in the vicinity of the electrode surface, get trapped in the polymer matrix, during the electropolymerization process, due to their ability to interact favorably with the PPY units. This behaviour was confirmed by the increase in oxidation and reduction peaks intensity, when adding dopamine, which made the PPY-CTS film with dopamine grow faster than the one without it.

Factors influencing MIP electrode performance

Different influencing factors, such as monomer, SE and template concentrations, pH medium, scan cycles of electropolymerization process and incubation time were optimized to make up an efficient sensor. The effect of the cited factors on the MIP dopamine sensor electrochemical performance was investigated by DPV.

Influence of monomer concentration

The influence of monomer concentration during electropolymerization is very important to sensors properties for dopamine detection. To determine the influence of PY concentration on the response of MIP modified electrodes, the films were prepared using a constant dopamine concentration (1 mM), and various PY amounts in the range from 1 to 100 mM.

As illustrated in Fig. 4A, the sensor responses increased with higher PY concentrations up to 5 mM in the solution mixture, and subsequently decreased. High monomer concentrations may result in a rapid polymerization process, improving the sensor sensitivity. However, too elevated monomer concentrations might lead to the formation of a thick membrane that makes some of the recognition sites not to be easily accessible [30]. So, the present work chose 5 mM as optimum monomer concentration.



Figure 4: Influence of different parameters on the electrocatalytic oxidation of dopamine in 0.1 mol/L PBS (pH = 6) solution. (A) PY concentration, (B) CH₃COOH concentration, (C) dopamine concentration, (D) electropolymerization cycle, (E) pH value, (F) incubation time.

Influence of the SE concentration

One of the important factors that affect MIP efficiency is SE concentration. This was studied under constant concentrations of CTS (1 mg/mL), dopamine (1 mM), PY (5 mM), and various CH₃COOH amounts, via CV measurements. MIP sensor response increased with SE concentrations up to 100 Mm, and then decreased (Fig. 4B). According to this result, 0.1 M was chosen as optimum SE concentration during all measurements.

Influence of template concentration

The quantity and ability of MIP recognition sites depend primarily on the mechanism and degree of interactions between monomer concentration and template amount in prepolymerization mixture [31]. Fig. 4C displays the influence of template concentration on electrodeposition of MIP layer. It is clearly seen that maximum peak current was observed when template concentration was about 1 mM. Therefore, this template concentration was selected to achieve polymerization step.

Influence of scan cycles during electropolymerization process

Electropolymerization scan cycles are one of the important factors for the formation of a successful MIP layer [32, 33], which can influence the thickness and compactness of the film. Fig. 4D shows the optimum number of scan cycles required to deposit a MIP onto the GCE, which possessed maximum current response during polymerization process. The modified electrode performances were evaluated in a 1 mM dopamine solution in PBS (pH = 6), at several MIP modified electrodes prepared by applying different numbers of voltammetric cycles in the pre-polymerization. The suitable number of electropolymerization cycles was estimated to be 10.

Influence of pH detection solution

The pH of the working solution can influence electrochemical performances of MIP-GCE/TiO₂/(PPY-CTS) sensor towards dopamine molecules detection. MIP response was tested in pH range from 4 to 7 (Fig. 4E). DPV responses of the sensor in contact with 1 mM dopamine, after 5 min, in PBS, revealed that the highest electrode response was at pH 6.

Influence of incubation time

After removing the template in the PBS solution (pH 7), MIP electrode was immersed in 1 mM of dopamine dissolved in 1 mM PBS (pH 6), for different incubation times. After each immersion, current response was measured by DPV method. As a result, 5 min was estimated as the optimum incubation time that led to maximum current. In fact, during this optimum time, the highest number of binding interactions can be established between MIP sites and dopamine molecules (Fig. 4F).

Effect of scan rate

Scan rate effect on voltammetric response current was studied for 1 mM dopamine in 0.1 mol/L⁻¹ PBS (pH 6). Fig. 5A presents the background subtracted voltammograms recorded under different scan rate values from 10 to 250 mV/s. Fig. 5B shows the linear relation between I_p and $v^{1/2}$.



Figure 5: (A) CV obtained for MIP at different scan rates from 10 to 250 mV/s, in 1 mM dopamine; **(B)** Plot of peak current versus square root of scan rate, in 1 mM dopamine.

Peak response increased linearly with an increase in square root of scan rate ($v^{1/2}$), in the range from 10 to 250 mV/s, according to the equation:

$$I_{pa} = 2.67 v^{1/2} + 7.321, (R^2 = 0.998)$$
 (1)

where it occurs a diffusion controlled process [34]. D diffusion coefficient was calculated using Randles–Sevcik equation.

Ip = (2.69.10⁵) A
$$n^{3/2} D^{1/2} v^{1/2} C$$
 (2)

where I_p is peak current, n is electron transfer number (2), D (cm²/s⁻¹) is diffusion coefficient, C is dopamine bulk concentration (10⁻⁶ mol/cm³), A is electrochemical active electrode area (A = 0.070 cm²) and v is scan rate (mV/s⁻¹). With the slope and other required values, 2.10⁻⁸ cm²/s⁻¹ D was obtained, in good agreement with literature data [35].

Surface morphology of the prepared materials

The morphology of the modified electrodes was examined by SEM, after verifying the homogeneity and adhesion of the deposit membrane. Fig. 6A shows the surface morphology of the TiO_2 deposit layer, characterized by spherical spongy module shaped particles, homogeneously distributed with aggregation [36].



Figure 6: SEM images of: **(A)** GCE/TiO₂; **(B)** GCE/TiO₂/PPY-CTS; **(C)** NIP-GCE/TiO₂/PPY-CTS; **(D)** GCE/TiO₂/PPY-CTS/dopamine; **(E)** MIP-GCE/TiO₂/PPY-CTS. Resolution of 100 nM, magnification of 80.000 and accelerating potential of 15.0 kV were used.

After modification of GCE/TiO₂ surface with PPY-CTS nanocomposite, a compact, uniform and homogeneous structure was observed (Fig. 6B). With SEM, it is hard to see the difference in morphology between GCE/TiO₂/PPY-CTS (B)

and GCE/TiO₂/PPY-CTS/dopamine (Fig. 6D) sensors, because the surface of the film formed is in the order of few nanometers.

Compared with NIP-GCE/TiO2/PPY-CTS (Fig. 6C) sensor, MIP-GCE/TiO2/PPY-CTS (Fig. 6E) one has a rougher surface, with several large pores. This behavior may be due to the template removal process, which only the MIP surface went through, resulting in considerable recognition sites for target dopamine interaction. In addition, the rough surface structure supports mass transfer and the formation of three-dimensional recognition sites.

FTIR and UV-visible studies

Fig. 7A presents FTIR spectra of dopamine powder, NIP-GCE/TiO₂/(PPY-CTS) and MIP-GCE/TiO₂/(PPY-CTS) deposit layers. The bands observed in FTIR spectra of dopamine powder were at 3342 cm⁻¹ (amine N–H stretching), 3207 cm⁻¹ (phenol O–H stretching), 2972 cm⁻¹ (aromatic C–H stretching), 2895 cm⁻¹ (alkyl C–H stretching), 1610 cm⁻¹ (amine N–H bending), 1496 cm⁻¹ (aromatic C=C stretching), 1248 cm⁻¹ (amine C–N stretching) and 1177 cm⁻¹ (phenol C–O stretching) [37-39]. FTIR spectra of the NIP-GCE/TiO₂/PPY-CTS and MIP-GCE/TiO₂/PPY-CTS modified surfaces were similar to each other in terms of patterns, and they both had some new peaks when compared with dopamine only. The main difference between the FTIR spectra from MIP and NIP sensors is the increase in bands, which can be attributed to the template extraction.

FTIR results for NIP-GCE/TiO₂/PPY-CTS and MIP-GCE/TiO₂/PPY-CTS sensors showed peaks at 3396 cm⁻¹, which may be due to N-H stretching of PPY and CTS. The peak at 2961 cm⁻¹ may correspond to C-H stretching of PPY and CTS. The small peak at 2120 cm⁻¹ stretching of PPY-CTS may be assigned to O-H vibration stretching of CTS.

Furthermore, CTS incorporation into PPY backbone is evident by the new band formed at 1655 cm⁻¹. This band could be due to C=O stretching vibrations of CTS. This band was clearly absent in dopamine FTIR spectrum. Also, the band at 1438 cm⁻¹ in both spectra may be attributed to C=C vibration stretching of PPY and CTS. The band at 1385 cm⁻¹ is due to N-H angular deformation couplings, while bands at 1249 cm⁻¹ and 1100 cm⁻¹ were ascribed to stretching vibrations of glycosidic bond C-O-C linkage [40-43]. The band centered at 656 cm⁻¹ is assigned to metal-O bonds Ti-O of TiO₂ [44, 45]. FTIR spectra of the deposit showed characteristic matrix bands that indicated the formation of a composite material.

TiO₂/(PPY-CTS), UV-vis spectra of TiO₂, NIP/TiO₂/(PPY-CTS), MIP/TiO₂/(PPY-CTS) and TiO₂/(PPY-CTS-dopamine) were displayed in Fig. 7B. A mixture of DMF and ultrapure water was used as reference solution. UV-visible absorption spectra of GCE/TiO₂/(PPY-CTS) show three absorption bands at 472, 513 and 704 nm. The first absorption band was assigned to π - π * transition, and the second band to the formation of composite PPY-CTS. The third band was attributed to the film oxidation. After dopamine addition to the composite matrix, the second band intensity, at 513 nm, decreased, which was due to the template adsorption on the electrode surface. After dopamine extraction, the band at 472 nm disappeared, and the absorption peak of the composite PPY-CTS has moved, which confirmed the formation of artificial cavities to recognize dopamine molecules.



Figure 7: (A) FTIR spectra of dopamine powder, NIP-GCE/TiO₂/(PPY-CTS) and MIP-GCE/TiO₂/(PPY-CTS); **(B)** UV-vis absorption spectra of TiO₂, TiO₂/(PPY-CTS), NIP/TiO₂/(PPY-CTS) and TiO₂/(PPY-CTS-dopamine).

Electrochemical characterization by using CV measurements

CV is a suitable and effective way for visualizing the current peak of the analytes in electrochemical detection [46]. In this work, it was also used to characterize electrodes modification process. CV results of the electrochemical sensor were recorded in a 1 mmol/L⁻¹/K₃[Fe(CN)₆] aqueous solution containing 0.1 mol/L⁻¹ KCl, as shown in Fig. 8.



Figure 8: CV obtained for: (a) bare GCE; (b) GCE/TiO₂; (c) GCE/TiO₂/(PPY-CTS); (d) GCE/TiO₂/(PPY-CTS-dopamine); and (e) MIP-GCE/TiO₂/(PPY-CTS), in 1 mM K_3 [Fe(CN)₆]/ K_4 [Fe(CN)₆] containing 0.1 M KCl, at a SR of 50 mV/s.

 $[Fe(CN)_6]^{3-}/[Fe(CN)_6]^{4-}$ redox couple was chosen as an electro-active probe to investigate its rate of electron mobility [47, 48]. From the CV, Fig. 8 shows that bare GCE gave well-resolved redox peaks, at 0.234 and 0.141 V, respectively, which are typical of a quasi-reversible CV. As seen in curve b, the current signal increased greatly. This result was due to the NP, which increased the active surface area and improved current response of the modified electrode.

After the electrodeposition of PPY-CTS-dopamine, anodic peak and cathodic peak currents were higher (Curve c) than those of the bare GCE and GCE/TiO₂, which implies that the composite can greatly enhance the effective electrode surface area. This increase in current may be due to the affinity of positively charged CTS with the negative charge of $[Fe(CN)_6]^{3-}/[Fe(CN)_6]^{4-}$ [49].

 Δ Ep of GCE/TiO₂/(PPY-CTS-dopamine) modified electrode was 0.110 V, as against 0.148 V, for bare GCE/TiO₂. This low Δ Ep indicates a faster and better electron transfer process in PPY-CTS-dopamine modified GCE [50].

However, after dopamine template extraction, electroactive probe redox peaks at the MIP-GCE/TiO₂/PPY-CTS surface were absent. This phenomenon shows that dopamine MIP film was successfully prepared.

Electrochemical sensing of dopamine at the MIP modified electrode

Electrochemical responses of bare GCE, GCE/TiO₂, NIP-GCE/TiO₂/(PPY-CTS), and MIP-GCE/TiO₂/(PPY-CTS) modified electrodes towards dopamine recognition were investigated using CV in 0.1 M PBS with pH 6, after 5 min incubation time (Fig. 9A).



Figure 9: (A) CV and (B) DPV spectra performed on GCE electrodes modified with 1 mM dopamine in a PBS (0.1 mol/L, pH 6) solution.

The bare GCE shows a significantly low oxidation peak at 0.25 V, compared to NIP-GCE/TiO₂/PPY-CTS and MIP-GCE/TiO₂/PPY-CTS, which gave a well-resolved oxidation peak, at 0.33 and 0.41 V, respectively, in PBS containing dopamine. DPV responses also showed clearly that MIP-GCE/TiO₂/PPY-CTS modified electrode presents a high recognition performance in dopamine presence (Fig. 9B).

The high affinity of MIP-GCE/TiO₂/PPY-CTS film with dopamine was due to: hydrophobic interaction between MIP-GCE/TiO₂/PPY-CTS film and dopamine (the aromatic part of dopamine molecules is hydrophobic in nature) [51]; acceleration of dopamine diffusion through the porous structure of MIP-GCE/TiO₂/PPY-CTS film; and MIP mechanism operating within the film [52].

The imprinted cavities in the film and the functional groups on the cavities produced by dopamine molecule template on the modified electrode displayed much higher binding properties with dopamine than with the NIP electrode.

Extraction of dopamine molecules

In this work, the extraction of dopamine from the composite matrix was obtained by CV from -0.2 to 1 V, in 0.1 mol/L PBS, at pH 7. This process was repeated for several cycles, until all dopamine molecules were extracted from MIP material [53]. Recorded CV are shown in Fig. 10, indicating that peak current decreased with more scans, and then it gradually tended nearly zero, which demonstrated the template molecules complete removal.



Figure 10: Extraction of dopamine template (10 scans) in 0.1 mol/L PBS (pH 7) solution, at a scan rate of 20 mV/s.

Performance of the modified electrode for dopamine detection

Typical DPV was employed to detect dopamine. Fig. 11A shows DPV curves of dopamine at MIP modified electrode recorded for various dopamine concentrations in a PBS (pH 6), with an incubation time of 5 min.

As it can be seen, dopamine anodic current increased with its higher concentrations. A well-resolved peak at 0.21 V was assigned to dopamine oxidation. When an adequate potential is applied to MIP electrode, it appears that dopamine electroanalytical behavior follows by exchanging two electrons and two protons, leading to dopamine-o-quinone.



Figure 11: (A) DPV obtained for MIP film with dopamine various concentrations in a PBS (pH 6). DPV parameters were: step E: 0.004V; modulation amplitude: 0.05 V; modulation time: 0.05 s; interval time: 0.2 s. (B) Plot of peak oxidation current vs. dopamine concentration of MIP and NIP. Average of three measurements (mean \pm SD).

Dopamine electrochemical oxidation mechanism on the modified MIP electrode is given by the following reaction (Scheme 2):



Scheme 2: Dopamine electroanalytical oxidation reaction on the MIP electrode.

Fig. 11B illustrates that calibration curve has a linear relation between I_p and dopamine concentration in the range from 10^{-6} to 10^{-5} M, with a correlation coefficient of 0.991. Additionally, the electrode sensitivity for dopamine detection is $3.202.10^{-6} \,\mu\text{A}/\mu\text{M}^{-1}/\text{cm}^{-2}$.

Based on IUPAC definition [54], LOD was calculated as:

$$LOD = 3 Sb/q \tag{3}$$

where Sb is standard deviation of the blank analyte and q is the slope of calibration plot. Hence, LOD of GCE modified in the optimum fabrication condition was estimated to be 0.281μ M.

Analytical parameters, such as linear response range, sensitivity and LOD, were compared with literature data, and are given in Table 1. Thus, results showed higher sensitivity and lower LOD than those from other electrochemical sensors used for dopamine determination.

Tuble 1. Comparison of various mounted electrodes for department detection.						
Modified electrode	Method	Linearity (mol/L ⁻¹)	LOD (mol/L ⁻¹)	References		
3DOM-MIPs/GCE	DPV	2×10 ⁻⁶ -2.3×10 ⁻⁴	9×10^{-7}	[57]		
Au/OPPy-DA	SWV	5×10 ⁻⁶ -5×10 ⁻⁵	1.04×10^{-6}	[58]		
PPy/DA-MIP	DPV	$1 \times 10^{-5} - 1 \times 10^{-4}$	7.97×10^{-6}	[59]		
OPPy/ERGO/GCE	DPV	2×10 ⁻⁶ –1.6×10 ⁻⁴	5×10^{-7}	[60]		
Nanostructured conducting polymer	VC	$1.0 \times 10^{-2} - 2.0 \times 10^{-2}$	1.5×10^{-6}	[61]		
MIP-GCE/TiO ₂ /(PPY-CTS)	DPV	$1 \times 10^{-6} - 1 \times 10^{-5}$	2.81×10 ⁻⁷	This work		

 Table 1: Comparison of various modified electrodes for dopamine detection.

Selectivity study of MIP-dopamine sensor

MIP-GCE/TiO₂/(PPY-CTS) and NIP-GCE/TiO₂/PPY-CTS modified electrodes DPV responses were recorded in PBS solutions with 10⁻⁵ mol L⁻¹ dopamine and 10⁻⁴ mol L⁻¹ of each interferent, separately. Fig. 12A shows the effect of interfering species Glu, UA and AA on the current response of MIP sensor in presence of 10⁻⁵ M dopamine. Fig. 12B shows electrochemical results of MIP and NIP electrodes given as a histogram graph. It can be clearly seen that MIP electrode interacted with dopamine more selectively than other analogue interferent species. This further confirmed the outstanding specificity of MIP towards dopamine template molecule, which was mainly due to the specific binding between it and the tailor made imprinted cavities.



Figure 12: (A) Effect of interfering species Glu, UA and AA on the current response of MIP sensor in presence of 10^{-5} M dopamine. (B) Anodic current peak value offered by MIP sensor in purple and NIP sensor in black for the detection of dopamine 10^{-5} M with different interfering substances in 10^{-4} M prepared in a PBS (0.1 M, pH 6) solution. Error bars indicate SD from three measurements.

Repeatability, reproducibility and stability of MIP-dopamine sensor

The fabricated MIP dopamine sensor was assessed for stability, in terms of repeatability, reproducibility and response time. To study the repeatability, 1 mM dopamine solution was analyzed for four times using MIP-GCE/TiO₂/PPY-CTS sensor under optimized conditions, employing DPV technique. RSD of the peak current intensities was about 2.4%, which indicates good repeatability.

To study reproducibility, four MIP-GCE/TiO₂/PPY-CTS sensors were fabricated under identical conditions, and their peak current intensity showed a RSD of 4%, indicating excellent reproducibility of the fabrication method.

MIP-GCE/TiO₂/PPY-CTS sensors were stored at room temperature, and the storage stability was evaluated for 1 month. During the first week, the sensor response to dopamine remained stable without any significant disruption. After 30 days, the sensor maintained about 89% of its initial current response, indicating that the prepared electrode had excellent long-term stability (Fig. 13).



Figure 13: MIP response stability with 1 mM dopamine.

Dopamine detection in real urine sample

In order to evaluate the applicability and feasibility of the prepared electrochemical sensor, a standard addition method was used to determine dopamine concentration in real urine samples from healthy individuals. The urine samples were diluted by 100-folds with a 0.1 M PBS (Ph 6), to overcome the effect of urine matrix interferences [56]. Initially, no dopamine was detected. Thereafter, successive dopamine additions were performed, and recovery percentages were calculated based on its determined concentrations. Table 2 shows very good recovery (between 93.3 and 116%) and reasonable RSD (less than 2%), indicating a hopeful applicability, with a good reliability of the prepared sensor for the measurements of trace dopamine in real samples analyses.

Sample	Added dopamine (M)	Found dopamine (M)	Recovery(%)	RSD*(%)
	5. 10-6	4.81. 10-6	96.2	1.023
Urine	10-5	9.71. 10 ⁻⁶	97.1	1.121
	1.5. 10-5	1.42. 10-5	94.6	0.571
	5. 10-5	5.15. 10 ⁻⁵	100.3	1.412

Table 2: MIP performances during dopamine determination in human urine samples (n = 3).

*Measurement values taken from three experiments.

Conclusion

The developed MIP-GCE/TiO₂/PPY-CTS sensor was successfully applied for detecting dopamine. The electrochemical MIP electrode was elaborated by electropolymerization of PPY-CTS composite onto the GCE/TiO₂ matrix, in the presence of dopamine as template molecule. Under optimum conditions, this sensor showed a rapid current response, wide linear range from 1×10^{-6} to 1×10^{-5} mol L⁻¹, low LOD of 2.81×10^{-7} mol/L⁻¹, high sensitivity and selectivity, good reproducibility and excellent stability for dopamine determination. It was shown that incorporating NP from TiO₂ in the sensitive matrix enhanced electrode was successful applied for monitoring dopamine in a real medium. The MIP sensor showed good dopamine recovery in urine samples. Hence, the simple and efficient strategy reported in this research can be further employed to fabricate MIP based electrochemical sensors for other target neurotransmitters recognition.

Acknowledgements

Authors would like to acknowledge the PRF-2017-D4P1/Tunisia project (No. 12Mag088) and the Ministry of Higher Education and Scientific Research of Tunisia (LR11ES55). The authors thank Dr. Florence GENESTE for his assistance for the SEM measurements.

Authors' contributions

Chama Mabrouk: writing. All authors have read and agreed to the published version of the manuscript **Houcine Barhoumi**: conceptualization and validation, review and editing. **Nicole Jaffrezic Renault**: conceptualization and validation; review and editing.

Abbreviations

AA: ascorbic acid CH₃COOH: acetic acid CTS: chitosan CV: cyclic voltammetry DMF: N,N-dimethylformamide DPV: differential pulse voltammetry FTIR: Fourier transform infrared spectroscopy GCE: glassy carbon electrode Glu: glucose H₂SO₄: sulfuric acid **IUPAC:** International Union of Pure and Applied Chemistry K₂HPO₄: dipotassium hydrogen phosphate K₃(Fe(CN)₆: ferrocyanide K₄(Fe(CN)₆: ferricyanide KH₂PO₄: potassium dihydrogen phosphate LOD: limit of detection **MIP**: molecular imprinted **NIP**: non-imprinted polymer **NP**: nanoparticle **PBS**: phosphat buffer solution **PPY**: polypyrrole **PY**: pyrrole **Redox**: reduction/oxidation reactions **RSD**: relative standard deviation **SD**: standard deviation **SE**: supporting electrolyte **SEM**: scanning electron microscopy UA: uric acid UV-vis: visible ultraviolet

Symbols definition

 ΔEp : peak-to-peak separation potential

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