RSC Advances



PAPER

View Article Online
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Cite this: RSC Adv., 2024, 14, 7836

Bimetallic metal organic framework/Ni doped ZnO nanomaterials modified carbon paste electrode for selective electrochemical determination of ciprofloxacin†

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In this work, an efficient and sensitive electrochemical sensor for the determination of ciprofloxacin (CIP) is reported. The sensor was prepared by using a carbon paste electrode (CPE) modified with a combination of bimetallic copper/cerium-based metal organic framework (Cu/Ce-MOF) and nickel doped zinc oxide nanoparticles (NZP). The modifiers were characterized by Brunauer–Emmett–Teller (BET) analysis, Fourier-transform infrared spectroscopy (FT-IR), X-ray diffraction (XRD), scanning electron microscopy (SEM), energy dispersive X-ray spectroscopy (EDX), and elemental mapping analysis (EDS). The electrochemical behavior of the modified electrode was studied by electrochemical impedance spectroscopy (EIS) and cyclic voltammetry (CV). The developed electrode was employed for the detection of CIP by differential pulse voltammetry (DPV). Under optimal conditions, the anodic peak current response of the electrode was linearly correlated with CIP concentration in the range of 0.75–100 μ mol L⁻¹ with a sensitivity of 1.29 μ A μ mol⁻¹ L⁻¹. The limit of detection and reproducibility of the method were 0.142 μ mol L⁻¹ and 2.7%, respectively. The developed sensor showed good selectivity to CIP against possible interferents. The method was applied to determine CIP in water, milk and urine samples. The results indicated that this method has potential to be applied in the analysis of residue CIP in complex matrices with high selectivity, and good reproducibility.

Received 29th November 2023 Accepted 23rd February 2024

DOI: 10.1039/d3ra08174e

rsc.li/rsc-advances

1. Introduction

Along with the rapid growth of the world's population, the demand for the use of antibiotics has increased significantly. As a result, soil, surface water, and underground water are severely affected by residues of antibiotics in human waste disposal, hospital and domestic wastewater effluents.1 Even trace amounts of antibiotics in the environment harm the ecosystem and the health of animals and humans, including interrupting photosynthetic cycles and biological functions of plants and increasing resistance to antibiotics. This global threatening problem led the United Nations to declare in 2017 that the discharge of antibiotics into environmental waters may host wild microorganisms that are completely resistant to the drugs.1 Among broad-spectrum antibiotics, fluoroquinolones with a quinolone ring structure show more activity against Grampositive and Gram-negative aerobic pathogens, due to the substitution of the fluorine atom at the C-6 position.² Ciprofloxacin (CIP) is a second-generation fluoroquinolone and the

most effective antimicrobial and anti-inflammatory antibiotic widely used in veterinary medicine and the treatment of various bacterial infections in humans. CIP affects bacterial growth and reproduction by inhibiting the function of an essential enzyme that maintains superhelical turns in bacterial DNA.3 Due to the poor metabolizing process and widespread use of CIP, its high concentration in municipal and hospital wastewater has been reported between 2.5 and 245 μg L⁻¹.4 It has also been detected at higher concentrations of 4.9 to 50 mg L⁻¹ in freshwater near pharmaceutical factories. Also, CIP has been found in cow's milk at concentrations of 0.1 to 100 ng mL⁻¹.3,5 CIP shows low biodegradability and high solubility in water at different pH values.6 Considering the residue of CIP in various media such as water and food samples, it is necessary to develop new strategies or methods that are simple, accurate, highly selective, and sensitive for its monitoring. So far, several analytical techniques have been employed to determine CIP in different matrices including liquid chromatography-mass spectrometry,7 high-performance liquid spectrophotometry,9 chromatography,8 spectrofluorometry,10 electrophoresis,11 immunoassay,12 iluminescence,13 photoelectrochemical detection,14 and electrochemical techniques. 15-17 Although these techniques are reliable and sensitive, most of them require several time-consuming treatment steps for sample preparation, expensive instruments,

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† Electronic supplementary information (ESI) available. See DOI: https://doi.org/10.1039/d3ra08174e

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or an expert operator. Among the above techniques, electrochemical techniques are characterized by the inherent advantages of high sensitivity and selectivity, ease of use, fast response, easy miniaturization, cost-effective analysis, and enable real-time monitoring of the target analyte.18 Various working electrodes are based on boron-doped diamond, 4,19 carbon paste electrode, 20 glassy carbon electrode,21 pencil graphite electrode,22 screen printed electrodes, 23,24 and etc. have been used to determine different classes of antibiotics. In recent years, carbon paste electrodes have been widely used for the monitoring of fluoroquinolones due to their low cost, wide potential window, rapid fabrication and regeneration of the electrode surface. Modification of the electrode surface with different electroactive materials increases electroactive sites for accumulation of analyte on the electrode surface and facilitates electron transfer. Up to now, various materials including surfactants, polymer layers, inorganic, organic and metal-organic framework materials, biomaterials, metal and semiconductor nanoparticles, and nanocomposites have been used to improve the electrochemical response of electrodes to the target analyte. 25,26 An ideal electrocatalyst should generally possess characteristics such as reasonable cost, high conductivity for charge transfer, and stability at high potentials.²⁷ Recently, organic non-precious metals such as transition metal nitrides, carbides, and phosphides have been considered as electrocatalysts, because of their variable oxidation states, presence of 3d electrons, stability, and morphological properties. However, low conductivity, aggregation and instability during electrocatalytic process are drawbacks of these materials.28 Graphitic carbon nitrides, carbon nanotubes, or carbon-based nanocomposites have been considered as metal-free catalysts as alternatives to conventional nobel metal catalysts owing to their high surface area, low cost, robustness, and tunable structures.29 Nonetheless, hybrid systems of inorganic and organic nanomaterials through covalent or noncovalent bonding have opened a new perspective to electrocatalysts. These combinations modify the morphology, improve the efficiency of electrocatalysts, and enhance the accessible catalytic active sites.30,31

Metal-organic frameworks (MOFs) consisting of a metal ion that connected with an organic linker represent a very important class of hybrid materials. Compared with other inorganic porous materials, MOFs can provide a sensing platform for electrocatalysis owing to their properties such as intrinsic porosity, tunable pore sizes, and high surface area.³² Howbeit, most MOFs have relatively low stability in aqueous solutions and poor electrical conductivity due to the reversible nature of the coordination bonds.³³ Several strategies have been developed to overcome these drawbacks. One common approach is to combine MOFs with other highly conductive functional materials such as carbon or metal oxide nanomaterials.³⁴ On the other hand, to improve high stability, metals with high oxidation states are used to form strong coordination bonds with the linker.

Recently, bimetallic MOFs have received much attention in various fields.³⁵ The incorporation of a second metal with the similar electronic property of the metal in the framework may affect the electronic property of the bimetallic MOF.³⁶ Nonetheless, most of the metal sites in the bulk bimetallic MOF are not

accessible to the electrolyte and buried within the framework. To overcome this issue, several protocols have been suggested, including increasing the surface area with increasing porous structure, selection of adequate bimetallic centers, and appropriate MOF substrate.37 Among the synthesized MOFs, ceriumbased metal-organic frameworks (Ce-MOFs) have attracted attention due to their unique optical and catalytic properties including low energy 4f orbitals, low cost, remarkable thermal and chemical stability, redox activity, and permanent porosity.38 Although Ce-MOF materials are promising electrocatalysts, but due to poor conductivity, the catalytic activity of Ce-MOF needs further improvement.39 Copper benzene-1,3,5-tricarboxylate MOF has been one of the most studied MOFs because of its large surface area, high pore volume, high chemical stability, and also the ability to bind small molecules. 40 It has proven that MOFs with different metal centers exhibit higher conductivity compared to the analogous structures and provide excellent electrochemical activity due to the synergistic electronic effect of the redox sites.41 In addition, metal oxide/bimetal-MOF composites can synergistically enhance activation with higher catalytic performance.42

Herein, we prepared a bimetallic MOF (Cu/Ce MOF) with benzene-1,3,5-tricarboxylic acid (H₃BTC) as an organic linker through a facile precipitation approach. Nickel doped zinc oxide nanoparticles (NZP) are non-toxic and facilitate electron transfer. 43 MOF is a promising candidate for supporting material in the fabrication of the electrochemical sensors. Then, a combination of Cu/Ce-MOF and nickel doped zinc oxide nanoparticles (NZP) was incorporated into carbon paste to fabricate modified carbon paste electrode for the detection of ciprofloxacin. The synergistic effect of Cu/Ce-MOF and NZP helped improve the ability of the sensor for the selective determination of ciprofloxacin under optimal conditions. The resulting material showed a large specific area, and enhanced electrocatalytic activity. The prepared electrode was characterized and the parameters affecting the response of the developed sensor were investigated and discussed. The electrochemical behavior of the sensor for CIP detection is reported in terms of detection limit and dynamic range. The fabricated electrode was applied for the detection of CIP in water, milk and urine samples with satisfactory results.

2. Materials and methods

2.1 Reagents and materials

Ciprofloxacin was purchased from Darupaksh pharmaceutical company (Karaj, Iran). Graphite powder was obtained from Fluka. Nitrate salts of zinc, nickel, cerium and copper metal ions, as well as 1,3,5-benzenetricarboxylic acid (H₃BTC), citric acid, NH₃, sodium phosphate mono and dibasic salts, and ethanol were all prepared from Merck company (Darmstadt, Germany). All solutions were prepared with ultrapure water provided by AquaMaX water production system (Young Lin, South Korea).

2.2 Apparatus

All electrochemical experiments were performed on an AutoLab potenstiostate PGS302N (Metrohm, Utrecht, Netherlands)

equipped with a three-electrode configuration cell in which a Cu/ Ce-MOF modified carbon paste electrode, Ag/AgCl (3 M), and Pt rod were used as the working, reference, and counter electrodes, respectively. EIS measurements were carried out in the AutoLab potentiostate. The powder X-ray diffraction (XRD) pattern was obtained by Phillips Xpert X-ray diffractometer with copper X-ray source. Morphology, mapping, and elemental analysis of the synthesized nanomaterials were studied by field emissionscanning electron microscopy (FE-SEM, MRA3 TESCAN, Czech) with an accelerating voltage of 15 kV. Functional groups of the synthesized nanomaterials were studied by recording the absorption spectra in wavenumber range of 400-4000 cm⁻¹ using a AVATAR 370 Fourier transform infrared spectrophotometer (FTIR, Thermo Nicolet, USA). The surface area and pore size of the synthesized MOF modifier were calculated from N2 gas adsorption/desorption isotherm by BET and BJH analysis at 77 K (Micromeritics ASAP 2020, USA). A professional Benchtop pH meter BP3001 (Trans Instruments, Singapore) was used to adjust the pH of solutions. For comparison, determination of CIP in real samples was performed by fluorometry with excitation/emission wavelength at 275/452 nm. The fluorescence spectra were recorded on a RF-5301PC spectrofluorometer (Shimadzu, Japan) equipped with a 10 mm path-length quartz cell and a xenon discharge lamp with an excitation/emission slit width of 5.0 nm.

2.3 Synthesis of the modifier

The bimetallic organic framework was prepared according to a previous report. ⁴² Briefly, solution A was prepared by dissolving 1 mmole of each $Ce(NO_3)_3 \cdot 6H_2O$ and $Cu(NO_3)_2 \cdot 3H_2O$ in 30 mL of 1:1 (v/v) ethanol/water solvent. Solution B contained 3.0 mmol linker H3BTC in 70 mL of 1:1 (v:v) ethanol/water mixture. Next, solution A was slowly added to solution B under stirring for 30 min until a white precipitate appeared. Then, it was aged for 24 h at room temperature. The precipitate was washed with ethanol/water mixture several times, and then dried in an oven at 60 °C. The resultant is named as Cu/Ce-MOF.

Nickel doped zinc oxide nanoparticles was prepared by sol gel method using citrate precursor according to our previous report. First, $Zn(NO_3)_2 \cdot 6H_2O$ and $Ni(NO_3)_2 \cdot 6H_2O$ salts were mixed thoroughly and dissolved in 100 mL of deionized water with a stoichiometric ratio of 0.98 : 0.02. After addition of citric acid as gelling agent to the mixture solution, ammonium hydroxide solution was slowly added. The pH of the mixture was adjusted to 7 and stirred for 3 h to form a homogeneous gel. The mixture was dried at room temperature to form a xerogel. Then, it was ground and sintered at 600 °C with a heating rate of 10 °C min⁻¹ for 2 h. The greenish-gray powder (Ni_{0.02}Zn_{0.98}O) abbreviated as NZP was used in subsequent experiments.

2.4 Fabrication of Cu/Ce-MOF/NZP/CPE

The carbon paste electrodes were prepared using graphite powder and paraffin oil (2 drops) as binder. The components of 0.15 g in total weight in the absence and presence of modifiers (1.3% NZP, and 2.7% Cu/Ce-MOF) were mixed thoroughly in a mortar for at least 25 min to obtain homogeneous paste. Then, the paste was packed into one end of a polyethylene insulin

syringe in 0.4 cm deep, and a copper wire passing through the center of the electrode body for electrical contact. The surface of each electrode was polished using a suitable paper to produce a smooth, shiny and reproducible working surface. Then, the prepared electrode was washed with deionized water to remove impurities. The Cu/Ce-MOF/NZP/CPE electrodes were fabricated by employing different amounts of Cu/Ce-MOF and NZP modifiers in paste composition.

2.5 Analytical procedure

The selective detection of CIP by Cu/Ce-MOF/NZP/CPE was conducted by the following procedure: the electrode was first activated in phosphate buffer at pH 3.0. Then, cyclic voltammetry (CV) was carried out in the potential range of 0–1.6 V (νs . SCE) at a scan rate of 50 mV s⁻¹. Quantification of CIP was carried out in 0.1 mol L⁻¹ phosphate buffer at pH 3 by differential pulse voltammetry with a pulse amplitude of 100 mV, a pulse width of 0.4 ms, and a potential scan speed of 100 mV s⁻¹.

2.6 Preparation of real samples

2.6.1 Milk sample. To precipitate of proteins and extract the remaining drugs, 50 mL of the milk sample was heated to 40 °C in a water bath. The pH of the sample solution was adjusted to 3 by 0.5 M HCl. After cooling the resulting suspension, it was centrifuged at 4000 rpm for 5 min to remove the precipitated proteins. The supernatant was filtered using a 0.45 μ m filter. Sample solution was stored at 4 °C before used.⁴⁴

2.6.2 Water sample. The surface water was diluted five times with phosphate buffer (0.1 mol L^{-1} , pH 3). Solutions of CIP at different concentrations were prepared with diluted surface water and the pH was adjusted to pH 3 with phosphate buffer. The samples were treated with the developed method. Recoveries of CIP were obtained based on the standard addition method.

2.6.3 Urine sample. A urine sample was collected from a healthy volunteer and centrifuged for 15 min at 2000 rpm without any further pretreatment. Then, it was filtered using a 0.45 mm filter, 5 times diluted with phosphate buffer solution (pH = 3), and transferred to the electrochemical cell for analysis. The standard addition method was used for the determination of CIP.⁴⁵

Results and discussion

3.1 Characterization

3.1.1 FTIR analysis. FTIR of the synthesized MOF (Fig. 1A(a)) shows a broad peak at 3398 cm $^{-1}$ which is related to the stretching vibration of the OH bond and may originate from physically or chemically adsorbed water molecules on the MOF. The characteristic peaks at 1612 cm $^{-1}$ and 1371 cm $^{-1}$ are attributed to the asymmetric and symmetric stretching vibrations of the carboxylate functional group, respectively. The absorption peak at 1556 cm $^{-1}$ is assigned to the stretching vibration of the C–O bond of H_3BTC . The absorption peaks at 530–760 cm $^{-1}$ correspond to Ce–O stretching vibrations. These observations proved that the metal ions were successfully included in the MOF structure and coordinated with the 1,3,5- H_3BTC ligand. 46,47

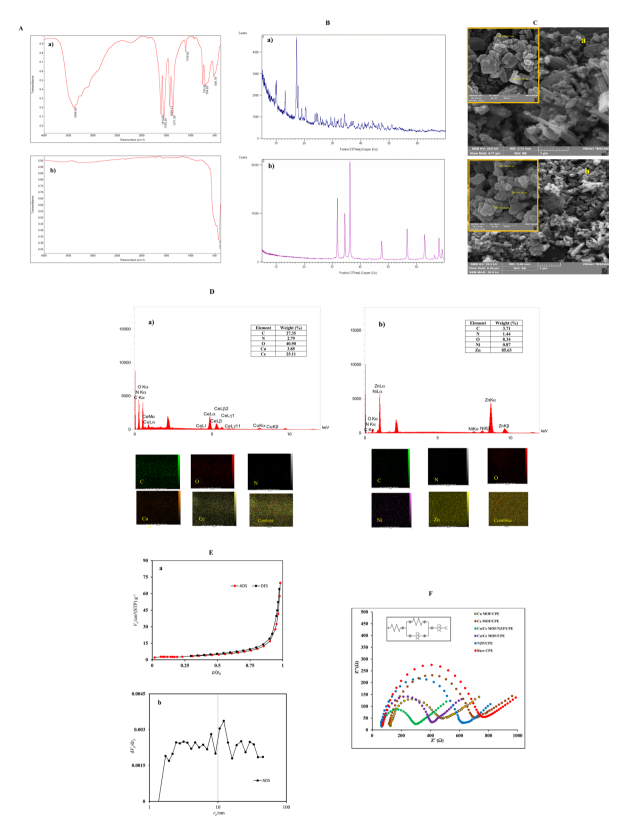


Fig. 1 (A) FT-IR spectra of (a) Cu/Ce-MOF, and (b) NZP; (B) XRD patterns for (a) Cu/Ce-MOF, and (b) NZP; (C) SEM images at two magnifications of (a) Cu/Ce-MOF, and (b) NZP; (D) EDX spectra of (a) Cu/Ce-MOF, and (b) NZP (inset shows the EDX and mapping analysis); (E) BET analyses (a) N_2 adsorption/desorption isotherm and (b) the BJH pore volume distribution of the synthesized Cu/Ce-MOF, and (F) the Nyquist plots of the bare CPE, NZP/CPE, Cu/Ce-MOF/CPE and Cu/Ce-MOF/NZP/CPE in 5.0 mmol L⁻¹ [Fe(CN)₆]^{3-/4-} solution containing 0.10 mol L⁻¹ KCl (Inset shows the equivalent Randles circuit).

The FTIR spectrum of NZP (Fig. 1A(b)) shows high intensity peaks at 550 and 434 cm $^{-1}$ which are related to the stretching vibrations of Zn–O. 43 The broad absorption peak located at 3458 cm $^{-1}$ corresponds to the O–H stretching vibration of $\rm H_2O$ absorbed on the surface of the Ni-doped ZnS nanocomposite. The absorption peak at 2340 cm $^{-1}$ is attributed to the presence of CO $_2$ in air. The stretching vibrations of Ni–O bond at 870 cm $^{-1}$ and 434 cm $^{-1}$ in NZP confirm the inclusion of nickel ions in ZnO matrix.

3.1.2 XRD analysis. Fig. 1B(a) shows the powder XRD pattern of MOF with strong peaks located at 2θ values of 5.0°, 9.96°, 10.1°, 13.3°, 17.3°, 17.8°, 19.1°, 19.8°, 20.6° corresponding to (200), (220), (222), (400), (331), (333), (420), and (442) reflection planes of the Ce(BTC)(H₂O)₆ MOF nanocrystals with octahedral geometry which agreed well with that reported in literature. The identified peaks with no additional peaks in the XRD pattern of Cu/Ce (BTC) MOF demonstrates a good purity and crystallinity of the Cu/Ce-MOF structure and revealed that Cu doped didn't affect the crystal structure of Ce-MOF. The XRD pattern of as-prepared Ni doped ZnO (NZP) (Fig. 1B(b)) shows diffraction peaks at 2θ value of 31.89°(100), 34.53° (002), 36.37° (101), 43.27, 47.66° (102), 56.72° (110), 62.98° (103), 66.49° (200), 68.06° (112) and 69.18° (201) which are consistent with the reported references for NZP. 42,49

3.1.3 FE-SEM analysis. The morphology of the synthesized Cu/Ce-MOFs was studied using FE-SEM as shown in Fig. 1C(a). The images show the nano rod structure of the as-synthesized MOF with an average diameter size of 52 nm and length of 1– 2 μ m. Fo Also, the FE-SEM images of NZP sample in Fig. 1C(b) shows a spherical morphology with an average particle size of 50 nm.

3.1.4 EDX and MAPPING analysis. The chemical composition of the synthesized NZP and Cu/Ce-MOF was determined by energy dispersive X-ray spectroscopy (EDX) as shown in Fig. 1D. EDX analysis indicated that Cu/Ce-MOF is composed of N, O, C, Ce and Cu elements with a molar ratio of Ce to Cu of 3:1. Mapping analysis in Fig. 1D(a) revealed that Ce and Cu elements are homogeneously distributed in MOF nanostructure.

Elemental mapping analysis of pristine NZP in Fig. 1D(b) also revealed that N, O, C, Ni, Zn elements were homogeneously distributed throughout in the nanoparticles structure.

3.1.5 BET analysis. The specific surface area of MOF was determined by Brunauer–Emmett–Teller (BET) analysis. The isotherm of Cu/Ce-MOF is type IV with a typical H3 hysteresis loop that appears at relative pressure above 0.3 (Fig. 1E(a and b)). The specific surface area ($S_{\rm BET}$) was estimated from the adsorption isotherm and obtained 10.738 m² g⁻¹. The average pore diameter of Cu/Ce-MOF was evaluated by BJH method and found to be 40.169 nm, which is promising for electron transfer and analyte diffusion. Such high surface area and porous structure of Cu/Ce-MOF has the potential to provide more reaction sites and improve the performance of electroactive material.

3.2 Electrochemical characterization of modified electrodes

The electroactivity of synthetic materials was tested by cyclic voltammetry (CV) and electrochemical impedance spectroscopy

(EIS). The electrochemical active surface areas of the electrodes were explored in electrochemical probe solution of 5 mM K₃Fe(CN)₆/K₄Fe(CN)₆ containing 0.1 M KCl by CV. Fig S1† shows the CV curves recorded on bare CPE, NZP/CPE, Cu/Ce-MOF/CPE and Cu/Ce-MOF/NZP/CPE in the potential window range of -1000 mV to 1000 mV with different potential scan rates (ν) from 5 to 200 mV s⁻¹. The voltammograms show a pair of well redox peaks corresponding to the redox reaction between K_3 Fe(CN)₆ and K_4 Fe(CN)₆. It is obvious that after modification of the bare CPE with Cu/Ce-MOF/NZP material, peak-to-peak separation potential decreased, while the current intensity of the redox peaks increased, indicating the enhanced electron transfer ability in the modified electrodes. In addition, the effect of the scan rate on the peaks' current was studied to evaluate the electrochemical active surface areas of the modified electrodes according to the Randles-Sevcik eqn (1):

$$i_{\rm p} = (2.65 \times 10^5) A D^{1/2} n^{3/2} v^{1/2} C^*$$
 (1)

The linearly dependence of the redox peak currents (i_p) on the square root of scan rates $(v^{1/2})$ indicates the diffusion controlled nature of the reaction occurred at modified CPEs. The electroactive surface area of bare CPE, NZP/CPE, Cu/Ce-MOF/CPE, and Cu/Ce-MOF/NZP/CPE was evaluated from the slope of the corresponding linear plots i_p - $v^{1/2}$, and found to be 0.184 cm², 0.197 cm², 0.215 cm² and 0.292 cm², respectively, which are larger than the geometric area (0.125 cm^2) of bare CPE. These observations implied that incorporation of Cu/Ce-MOF and NZP modifiers into the CPE provides more active sites, thereby improving the effective surface of the modified CPEs compared to the bare electrode, and enhancing the electrochemical activity of the Cu/Ce-MOF/NZP. Therefore, Cu/Ce-MOF/NZP/CPE was used for subsequent electrochemical measurements of CIP.

The interfacial behavior of electrodes has been extensively studied by EIS analysis. Commonly, an electrochemical system can be described by an analog Rundles circuit containing resistance, capacitance, and inductance. Commercial software is used to fit the experimental impedance data to a simple circuit model. The impedance of an electrochemical cell is examined with the Nyquist plot, which is a plot of the imaginary impedance against real impedance. In order to further prove the benefit of the Cu/Ce-MOF/NZP as an electrocatalyst in the electrooxidation of CIP into CPE, EIS measurements were performed for bare CPE, NZP/CPE, Cu-MOF/CPE, Ce-MOF/CPE, Cu/Ce-MOF/CPE, and Cu/Ce-MOF/NZP/CPE. The fitted corresponding Nyquist plots and equivalent circuit are illustrated in Fig. 1F. The values of the fitting parameters are summarized in Table 1, where R_s represents the electrolyte resistance, Rct is the charge transfer resistance, and constant phase elements CPE₁ and CPE₂ are the respective capacitances at the electrode/electrolyte interface and faradaic processes. As shown in Table 1, among the modified electrodes, the lowest Rct value obtained for CPE modified with NZP and Cu/Ce-MOF (Cu/Ce-MOF/NZP/CPE), indicating a decrease in resistance to the charge transport of redox probe at the Cu/Ce-MOF/NZP/ CPE and the increase in electron-transfer kinetics at this electrode surface.

Table 1 Impedance parameters for the Nyquist plots of each prepared modified CPE electrodes in $[Fe(CN)_6]^{3-/4-}$ using the equivalent Randles circuit

Electrode	$R_{ m s}\left(\Omega ight)$	$R_{\mathrm{ct}}\left(\Omega ight)$	CPE ₁ (μMho)	CPE ₂ (mMho)
Bare CPE	55	607	$Y_0 = 1.53 \ n_1 = 0.921$	$Y_0 = 3.05 \ n_2 = 0.28$
Ce-MOF/CPE	98.9	514	$Y_0 = 2.61 \ n_1 = 0.894$	$Y_0 = 3.21 \ n_2 = 0.274$
NZP/CPE	53	509	$Y_0 = 1.53 \ n_1 = 0.944$	$Y_0 = 4.63 \ n_2 = 0.322$
Cu-MOF/CPE	55	262	$Y_0 = 3.91 \ n_1 = 0.918$	$Y_0 = 2.98 \ n_2 = 0.233$
Cu/Ce-MOF/CPE	52	300	$Y_0 = 2.49 \ n_1 = 0.941$	$Y_0 = 4.26 \ n_2 = 0.33$
Cu/Ce-MOF/NZP/CPE	53	184	$Y_0 = 3.67 \ n_1 = 0.926$	$Y_0 = 4.14 \ n_2 = 0.282$

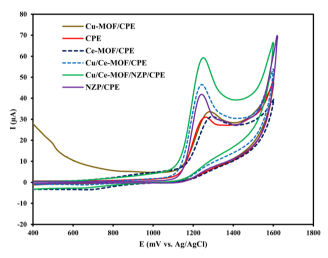


Fig. 2 Cyclic voltammograms of 50 μ mol L $^{-1}$ of target analyte in 0.10 mol L $^{-1}$ phosphate buffer solution at the bare CPE, NZP/CPE, Ce-MOF/CPE, Cu-MOF/CPE, Cu/Ce-MOF/CPE and Cu/Ce-MOF/NZP/CPE.

The inhomogeneity of the electrode surface often causes heterogeneous electron transfer and deviation from the ideal response of the resistance and capacitor, resulting in frequency dispersion of the interfacial capacitance and the appearance of a constant phase element (CPE). The impedance of CPE (Z_Q) can be expressed by eqn (2):

$$Z_{\rm Q} = \frac{1}{Y_0(j\omega)^n} \tag{2}$$

where Y_0 is a constant representing the charge separation at the interface, and n corresponds to the roughness of the electrode surface with values ranging from 0 to 1. Other symbols have their common meanings. Obviously, the Cu/Ce-MOF/NZP/CPE with the highest CPE₁ possesses the largest active electrode surface with highest active sites.

3.3 Electrochemical behavior of target analyte on modified electrodes

Cyclic voltammograms of 1×10^{-4} mol L $^{-1}$ CIP in 0.1 mol L $^{-1}$ phosphate buffer solution (pH = 3) at bare CPE, NZP/CPE, Cu/Ce-MOF/CPE, and Cu/Ce-MOF/NZP/CPE were recorded (Fig. 2). The voltammograms for CIP at Cu/Ce-MOF/CPE, and Cu/Ce-MOF/NZP/CPE showed two well defined anodic peaks situated at 125 mV and 486 mV νs . Ag/AgCl corresponding to the oxidation of

Cu to Cu⁺ and Cu²⁺, respectively, and two cathodic peaks in the reverse scan located at 273 mV and -44 mV, corresponding to the reduction of Cu²⁺ to Cu⁺ and Cu, respectively (Fig. 3). These peaks didn't observe in voltammograms for CIP at bare CPE and NZP/CPE. This redox event suggests that the Cu/Ce-MOF is electroactive due to the presence of the Cu²⁺ state in this framework with a partially metallic state at the Fermi level.⁵¹ A new peak for CIP oxidation (*vs.* Ag/AgCl) was observed at 1.25 V in all voltammograms which can be attributed to the irreversible redox behavior of CIP. The highest anodic peak current of CIP oxidation on the electrode surface of Cu/Ce-MOF/NZP/CPE compared to other CPEs clearly indicates that the combination of Cu/Ce-MOF and NZP nanoparticles improve the electron transfer for CIP oxidation and revealing the catalytic effect of the modifiers on the oxidation of target analyte.

3.4 Effect of scan rate and kinetics of oxidation process

Study the effect of scan rate on the peak current can be useful for elucidating the electrochemical mechanism. Therefore, cyclic voltammograms of 50 $\mu \rm mol~L^{-1}$ CIP at Cu/Ce-MOF/NZP/CPE were recorded at different scan rates from 5 to 160 mV s $^{-1}$. From Fig. 3, it is evident that the current gradually increases with the increase of the scan rates along with the shift of the oxidation peak potential to positive values. The anodic peak current ($i_{\rm p}$) showed a linear relationship with the square root of scan rates ($v^{1/2}$) by a regression equation of $i_{\rm p}$ ($\mu \rm A$) = 1.20 $v^{1/2}$ (mV s $^{-1}$) $^{1/2}$ + 5.57 and the correlation coefficient R^2 = 0.9887. In addition, the plot of log $i_{\rm p}$ against log v showed a linearity with a slope of 0.32, which is very close to the theoretical value of 0.5, confirming that the oxidation of CIP on the surface of Cu/Ce-MOF/NZP/CPE is a diffusion-controlled process.

The value of the electron transfer coefficient (α) in an irreversible system is between 0.3 and 0.7. For an anodic reaction, α can be calculated from the slope of Tafel plot using data from the initial part of the voltammetric wave of the current–potential curve for the oxidation of 50 μ mol L⁻¹ CIP at a potential scan rate of 20 mV s⁻¹. Considering an arbitrary value of 0.5 for α , the number of electrons in the rate-determining step can be evaluated from the eqn (3):

$$\alpha = \frac{RT}{F} \frac{\mathrm{d}(\ln i)}{\mathrm{d}E} \tag{3}$$

Taking into account that $\alpha=\frac{2.303RT/F}{\text{slope}}$ with the slope of the linear regression of the Tafel plot in Fig. S2,† the value of

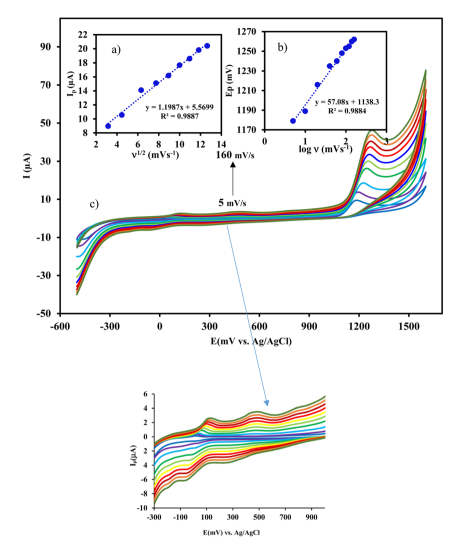


Fig. 3 Effect of scan rate (from 5 to 160 mV s⁻¹) on the response of CIP at Cu/Ce-MOF/NZP/CPE. Inset shows the plot of (a) I_p vs. $\nu^{1/2}$, and (b) E_p vs. log ν .

 α is estimated to be 0.476. The anodic peak potential (mV) of cyclic voltammogram varies linearly as a function of the logarithm of the scan rate (mV s⁻¹) with the regression equation $E_{\rm pa}$ (mV) = 57.1 log ν + 1138.3 (R^2 = 0.9884) (inset of Fig. 3), according to Laviron equation for irreversible redox process (4):

$$E_{\text{Pa}} = E^{\circ \prime} + \frac{(2.303RT)}{(1-\alpha)n_{\alpha}F} \log \frac{(RTK_{\text{s}})}{(1-\alpha)n_{\alpha}F} + \frac{(2.303RT)}{(1-\alpha)n_{\alpha}F} \log \nu \quad (4)$$

where $E^{\circ\prime}$, n_{α} , α and $k_{\rm s}$ are the standard electrode potential, the number of electrons involve in the rate determining step, charge transfer coefficient, and the rate constant of the electrochemical reaction. Other symbols have their common meanings. By substituting the α value (obtained from the Tafel plot) in the slope of $E_{\rm pa}-\log\nu$ plot, the n_{α} value of 1.976 was obtained, suggesting that two electrons are involved in the rate determining step of the CIP electro-oxidation reaction at Cu/Ce-MOF/NZP/CPE. Also, the value of rate constant ($k_{\rm s}$) at a scan rate of 100 mV s⁻¹ was calculated from eqn (5), and obtained 1.9 × $10^{-3}~{\rm s}^{-1}$.

$$\ln K_{\rm s} = \alpha \ln(1-\alpha) + (1-\alpha) \ln \alpha - \ln \left(\frac{RT}{n_{\alpha}F\nu}\right) - (1-\alpha) \frac{\alpha n_{\alpha}F\Delta E}{RT}$$
(5)

3.5 Chronoamperometric study

Chronoamperometric measurement was carried out for the determination of diffusion coefficient (D) of the target analyte by setting Cu/Ce-MOF/NZP/CPE at a fixed potential of 1.1 V in 50 s for different CIP concentrations in phosphate buffer (Fig. 4). Using the slope of the straight line of the currents against $t^{-1/2}$ plot at each concentration, and the linear relationship of the slopes with the CIP concentrations (insets of Fig. 4), the value of the diffusion coefficient (D) of CIP was calculated ($D = 7.68 \times 10^{-4} \ {\rm cm}^2 \ {\rm s}^{-1}$) according to the Cottrell eqn (6):

$$I = nFACD^{1/2}\pi^{-1/2}t^{-1/2}$$
(6)

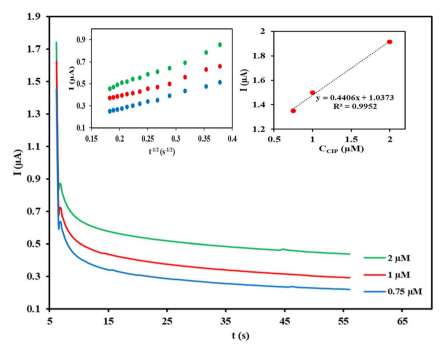


Fig. 4 Chronoamperometry of CIP in 0.1 mol L $^{-1}$ phosphate buffer (pH = 3) at three different concentrations (0.75, 1.0, 2.0 μ mol L $^{-1}$) with a potential step at 1.1 V at Cu/Ce-MOF/NZP/CPE. Insets show the plot of / (μ A) vs. $t^{-1/2}$ (s $^{-1/2}$) obtained from chronoamperograms, and plot of slopes of the linear lines *versus* CIP concentrations.

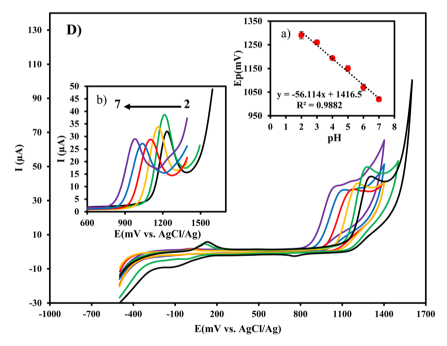
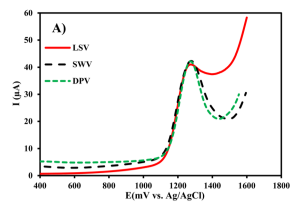


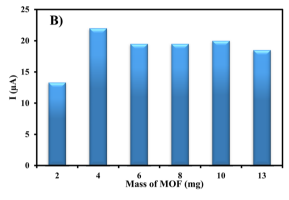
Fig. 5 Effect of pH on the response of Cu/Ce-MOF/NZP/CPE for CIP detection.

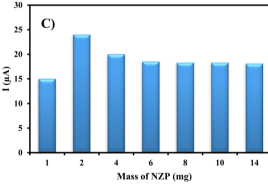
3.6 Effect of pH

The effect of pH on the response of Cu/Ce-MOF/NZP/CPE electrode in CIP solution (10 $\mu mol~L^{-1})$ was investigated by cyclic voltammetry and the results are shown in Fig. 5. It can be seen that the oxidation current of CIP reached to the highest value at pH 3 and then decreased with increasing pH which might be

due to the participation of protons in the oxidation process. Moreover, the anodic peak potential shifted towards less positive values with increasing solution pH. The relationship between the anodic peak potentials and the pH values of the CIP solution (inset of Fig. 5) is linear with the regression equation $E_{\rm pa}$ (V) = -0.0561 pH + 1.4165 ($R^2=0.9882$). The closeness of the slope value to the theoretical value of 59 mV per pH decade,







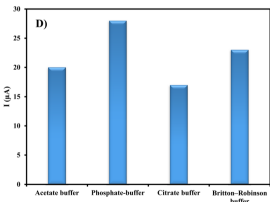


Fig. 6 Influence of the (A) type of technique, (B) Cu/Ce-MOF mass, (C) NZP mass, and (D) type of buffer on the anodic response of the Cu/Ce-MOF/NZP/CPE to CIP detection.

implies that the number of protons and electrons in CIP oxidation at Cu/Ce-MOF/NZP/CPE is equal. Thus, the mechanism of CIP oxidation at Cu/Ce-MOF/NZP/CPE is probably

through the oxidation of the –NH group to hydroxylamine (–N–OH) pathway by two protons and two electrons as presented in Fig. S3.† CIP contains a carboxylic acid and an amine functional group in piperazinyl ring (p $K_{a1}=6.1$, p $K_{a2}=8.7$), indicating that at pH below 6.1 and above 8.7, cationic and anionic forms of CIP predominate, respectively. Therefore, the electrostatic repulsive interactions between anionic form of CIP and carboxylate group on the electrode surface reduce oxidation peak current. Further, π – π interactions between the linker and the CIP molecule may facilitate the diffusion of CIP to the electrode surface and enhance the response of the electrode to the target analyte.

3.7 Optimization

3.7.1 Optimizations of the experimental parameters. To increase the electroanalytical performance of Cu/Ce-MOF/NZP/CPE, the influence of several parameters such as the type of voltammetry techniques, the amounts of modifiers, the pH of the sample solution, and the type of buffer as well as the instrumental parameters on the electrode response were investigated.

Since a high electrochemical oxidation peak current is necessary to determine the CIP at Cu/Ce-MOF/NZP/CPE, voltammetric techniques including LSV, SWV, and DPV were evaluated. DPV was chosen because of its higher signal and smaller peak width than the other studied techniques (Fig. 6A).

The content of Cu/Ce-MOF in the CPE composition affects the oxidation current of CIP. For this reason, six electrodes containing different amounts of Cu/Ce-MOF (2, 4, 6, 8, 10 and 13 mg) were prepared and investigated for the detection of 50 μ mol L⁻¹ CIP under the same conditions. As can be seen from Fig. 6B, the peak current increases up to 4 mg (2.7%, w/w) of Cu/Ce-MOF in the CPE composition and decreases with further increase of the modifier amount.

In addition, the effect of the mass of NZP nanoparticles on the anodic current of CIP oxidation was investigated. Seven electrodes containing different amounts of NZP (1, 2, 4, 6, 8, 10 and 14 mg; from 0.7% to 9.0% w/w of the CPE composition) were prepared and examined for the detection of 50 μ mol L $^{-1}$ CIP. It was observed that the peak current increases up to 1.3% of NZP in CPE composition, and then decreases (Fig. 6C). As a result, NZP and Cu/Ce-MOF with 1.3% (w/w) of and 2.7% (w/w) in CPE mass, respectively, were considered as the optimal amounts of modifiers for CIP quantification in subsequent studies.

To investigate the maximum response of the target analyte at pH 3, various buffers such as Britton–Robinson, acetate, citrate and phosphate buffers at 0.1 mol L^{-1} ionic strength were used. The maximum current response with a well-defined peak shape was obtained in phosphate buffer, which was used to detect the target analyte in subsequent studies (Fig. 6D).

The effect of potential scan rate (ν) on the performance of Cu/Ce-MOF/NZP/CPE was also studied in the range of 5–160 mV s⁻¹. Increasing the scan rate resulted in a higher oxidation peak current without changing the peak width, so 100 mV s⁻¹ was chosen as the optimal scan rate in further studies.

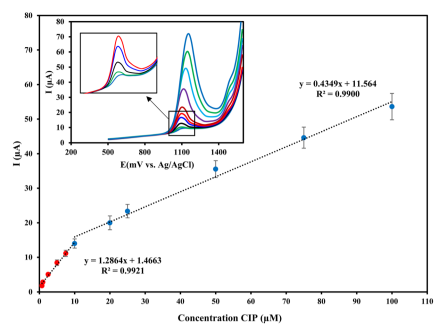


Fig. 7 The calibration curve and linear range of Cu/Ce-MOF/NZP/CPE for determining CIP in 0.1 mol L^{-1} phosphate buffer (pH = 3) under optimal conditions (scan rate: 100 mV s⁻¹, pulse amplitude: 100 mV). Inset shows the differential pulse voltammograms of CIP.

Table 2 Comparisons of analytical performance of Cu/Ce-MOF/NZP/CPE for CIP with other reported electrodes modified with different materials^a

Electrode	Linear range (μM)	Technique	$\mathrm{LOD}\left(\mu M\right)$	Real sample	Ref.
ds-DNA-BDD	0.5-60	DPV	0.44	Urine	19
MWCNT/GCE	3-1200	LSV	0.9	Urine, plasma, tablet	52
MWCNT-GCE	40-1000	Chronoamperometry	6	Urine, serum	53
Boron-doped diamond electrode	0.74-20	DPV	0.6	Urine	54
DNA/GCE	1-10	DPV	0.12	_	55
TiO ₂ /PB/AuNPs/CMK-3/Nafion/GE	1-10	CV	0.10	Environmental water	56
CRGO/GC	6-40	SWV	0.21	Drugs and milk	57
Cu/Ce-MOF/NZP/CPE	0.75-100	DPV	0.14	Water, milk, urine	This work

^a CPE – carbon paste electrode; GCE – glassy carbon electrode; GE – graphite electrode, GO – graphene oxide, NiONP – nickel oxide nanoparticles; SWV – square wave voltammetry; CRGO – chemically-reduced graphene-oxide.

3.8 Electrochemical detection of CIP

The electrochemical response of Cu/Ce-MOF/NZP/CPE to CIP with different concentrations in PB (pH 3) under optimal conditions at an oxidation potential of 1.25 V was determined by DPV technique. As shown in Fig. 7, the oxidation peak current increases with increasing CIP concentration, so that the plot of current responses against CIP concentrations is bilinear within the ranges from 0.75 $\mu mol~L^{-1}$ to 10 $\mu mol~L^{-1}$ and from 10 $\mu mol~L^{-1}$ to 100 μ mol L⁻¹ with linear regressions of $I_{pa} = 1.2864C_{CIP} + 1.4663$ ($R^2 =$ 0.9921) and $I_{pa} = 0.4349C_{CIP} + 11.564$ ($R^2 = 0.9900$), respectively. According to the slopes of the calibration curves, the sensitivity of the method in the low dynamic range is $1.2864 \, \mu A \, \mu mol^{-1} \, L$ which is about 3 times higher than that of the higher concentration range. According to the IUPAC guideline, the limit of detection (LOD = $3\sigma/m$) and limit of quantitation (LOQ = $10\sigma/m$) of the proposed method are estimated to be 0.142 μ mol L⁻¹ and 0.47 μ mol L⁻¹, respectively.

The present method is compared with reported electrochemical methods for the CIP quantification. ^{19,52-57} As can be seen in Table 2, the performance of the designed electrode is comparable or better than other reported electrodes for the detection of CIP in terms of linear range, detection limit, and type of real sample tested. Besides, the proposed electrochemical electrode

Table 3 The effect of interferents on the determination of CIP

Interferents	Tolerance limit (mol mol ⁻¹)		
Ca ²⁺ , NO ₃ ⁻ , Mg ²⁺ , Cl ⁻ , Na ⁺ , PO ₄ ³⁻ , K ⁺ , Cu ²⁺ , NH ₄ ⁺ , urea, lysine, creatinine	1000		
SO_4^{2-}	800		
Citric acid, sucrose, glucose	500		
Ascorbic acid, histidine	100		

Table 4 Determination of CIP in real samples at Cu/Ce-MOF/NZP/CPE and comparison with fluorometric method^a

Sample Milk		Suggested method			Fluorometric method		
	Amount added $(\mu \text{mol L}^{-1})$	Amount found $(\mu \text{mol L}^{-1})$	Recovery RSD ^b (%)		Amount found $(\mu \text{mol L}^{-1})$	Recovery RSD^{b} (%)	
			_	_	_	_	
	2.5	2.9	114	3.3	2.6	102.2	2.5
	5.0	5.4	107	0.4	4.5	90.2	1.9
	7.5	7.4	98.0	3.7	6.4	85.1	1.2
	10	9.9	99.0	3.8	8.2	82.3	5.6
Urine	0	_	_	_			
	2.5	2.4	94.0	1.8	2.6	105.6	3.3
	5.0	5.5	109	2.5	4.8	96.5	12.6
	7.5	7.5	99.0	1.9	6.9	91.6	8.5
	10	9.4	93.0	4.1	8.7	86.7	8.4
Water	0	_	_	_	_	_	
	2.5	2.6	104	2.2	3.3	134	9.4
	5.0	5.4	108	2.9	5.3	106	8.1
	7.5	7.4	98.0	2.8	8.4	112	3.3
	10	9.7	96.0	2.5	10.5	105	0.6

 $^{^{}a}$ t_{crit} . $(p = 0.05, n - 1 = 3) = 3.18; t_{\text{exp. milk}} = 2.57; t_{\text{exp. urine}} = 1.36; t_{\text{exp. water}} = 1.73; F_{\text{crit, 2, 2}} = 19.$ Based on triplicate analysis.

has a renewable surface, simple preparation and cost-effective compared to some reported electrodes that provide almost the same analytical characteristics, making it a desirable sensing platform for the target analyte.

3.9 Repeatability, reproducibility, stability and selectivity of the method

The repeatability of Cu/Ce-MOF/NZP/CPE was evaluated in terms of relative standard addition (RSD%) for seven replicated CIP response measurements at a concentration of 10 $\mu mol \ L^{-1}$ and was found to be 2.7%. The reproducibility of the sensor was assessed by measuring the oxidation peak current using seven individual sensors and was 3.6%. The results confirm the efficient performance of the sensor with satisfactory reproducibility. The long-term stability of the developed sensor toward CIP was studied for one month and found that the initial response reached 90% after 2 weeks, which suggests that the developed sensor is stable over a long period.

The selectivity of Cu/Ce-MOF/NZP/CPE was evaluated using the DPV technique by measuring the electrode response to CIP in the presence of interfering substances commonly present in water, urine and milk samples such as ions, amino acids, sugars and drugs. In this study, the tolerance limit is defined as the maximum concentration of the coexisting substance that affects the response of Cu/Ce-MOF/NZP/CPE to CIP (10 $\mu mol\ L^{-1}$) by more or less than 5%. The results in Table 3 indicate that most of the examined substances did not interfere in CIP measurement. Among them, nalidixic acid, glutamic acid, naproxen and ofloxacin probably interfere with analyte measurement due to their similar structures.

3.10 Analysis of CIP in urine, water, and milk

Under optimum conditions, the Cu/Ce-MOF/NZP/CPE was applied to determine CIP in urine, water, and milk samples by employing DPV. The recoveries of CIP in the samples were

calculated using standard addition method. From the results in Table 4, the recoveries obtained for analysis of CIP in urine, water, and milk samples were found in the ranges of 93–109%, 96–105%, and 98–114%, respectively. The precision of the method as percent relative standard deviation (RSD%) for each sample is summarized in Table 4. The RSD of each sample is less than 5%, implying that the practical ability of the developed sensor for rapid detection of CIP in complex matrices with high precision. Also, the results obtained by the proposed method were compared with fluorometric method due to fluorescence property of CIP. Although the results showed low recoveries of analyte in the studied samples by flourometry, but statistical analysis using Student's t-test and variance ratio t-test at 95% confidence level (t = 0.05) indicated no significant difference between the performance of the two methods (Table 4).

4. Conclusions

Cu/Ce-MOF and NZP were used as modifiers to fabricate a selective and robust modified electrode for CIP detection by differential pulse voltammetric technique. Characterization of Cu/Ce-MOF modifier showed a relatively high surface area and porous structure which provided more reaction sites and facilitated electron transfer and analyte diffusion to the electrode surface. The carbon paste electrode modified with Cu/Ce-MOF and NZP modifiers showed significant sensitivity and selectivity for the electrochemical determination of CIP. In comparison with other electrochemical sensors for detection of CIP, the proposed sensor showed a large linear concentration range $(0.75-100 \, \mu \text{mol L}^{-1})$, acceptable limit of detection $(0.142 \,$ μmol L⁻¹; 47 ng mL⁻¹), good reproducibility (RSD% < 5), and high selectivity. It was successfully applied to determine CIP in serum, urine, and water samples. This sensor has been shown to be promising for the determination of CIP in complex matrices.

Conflicts of interest

There are no conflicts to declare.

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Paper

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