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Graphite nanocrystals coated paper-based electrode for detection of SARS-Cov-2 gene using DNA-functionalized Au@carbon dot core–shell nanoparticles

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ARTICLE INFO

Keywords:
- Au@CD core–shell NPs
- Electrochemical genosensor
- Graphite nanocrystals
- SARS-CoV-2 RdRP

ABSTRACT

Currently, the development of biosensors is an urgent need due to the rapid spread of SARS-CoV-2 and the limitations of current standard methods for the diagnosis of COVID-19. Hence, many researchers have focused on the design of high-performance biosensors for measuring coronavirus genes. In this study, a voltammetric genosensor was developed for the determination of SARS-CoV-2 RdRP gene based on the format of cDNA probe/Au@CD core–shell NPs/graphite nanocrystals (GNCs)/paper electrode. For the first time, graphite nanocrystals were used in the electrochemical biosensor design. This genosensor was exposed to different concentrations of virus gene and then the hybridization between cDNA probe and RdRP gene was monitored by redox-active toluidine blue (TB). With increasing the RdRP concentration, the reduction peak current of TB enhanced in a linear range of 0.50 pM-12.00 nM according to the regression equation of I (μA) = 7.60 log C_{RdRP} (pM) + 25.78. The repeatability with a RSD of 2.2% clearly exhibited that the response of modified electrode is stable because of the high adhesion of GNC layer on the paper substrate and the high stability of cDNA-Au@CD bioconjugates. The spike-and-recovery studies showed the acceptable recoveries for the sputum samples (>95%).

1. Introduction

The pandemic of coronavirus disease 2019 (COVID-19) caused by the SARS-CoV-2 still remains a global threat [1]. Millions of people have died from this disease, and many of them continue to suffer from symptoms long after their initial infection is gone. Regarding the high rate of transmission of this viral infection and the involvement of all countries in the world, it is necessary to make fundamental decisions to manage this fatal viral disease. The most important practical step in this regard is to break the chain of infection transmission between infected and susceptible people in the community, which requires fast and reliable diagnosis methods for the accurate viral detection. At present, the qualitative reverse transcription-polymerase chain reaction (RT-PCR) assay as a nucleic acid amplification test is the most common method for the detection of different virus genes, especially nucleocaspid (N), envelope (E) and RNA-dependent RNA polymerase (RdRP) genes [2–4]. However, the possibility of both false-negative and false-positive errors should be considered in the interpreting of RT-PCR tests. The false-positive results can arise from contamination with previously amplified DNA [5]. While, the false-negative errors can result in the viral RNA degradation during transport or storage, poor sample collection and low virus concentrations [6]. The large number of both false-negative and false-positive errors reduce the reliability of this method. Therefore, it is necessary to refine the current available tests for the sensitive and reliable diagnosis of SARS-CoV-2, aiming to control and prevent large outbreak of COVID-19.

The electrochemical genosensors are an alternative method for detection of viral genes, due to excellent sensitivity, high selectivity, simplicity and low limit of detection (LOD). More importantly, they don’t require amplification processes as the RT-LAMP or RT-PCR [7]. Combining the sensing strategies and nanotechnology can make these analytical diagnostic tools more efficient. In particular, the use of AuNPs and carbon-based nanomaterials in the design of electrochemical genosensors leads to the sensing technology evaluation. Numerous research articles published annually on metal NPs and carbon nanomaterials as the electrocatalysts or electron-transfer accelerators confirms this claim [8–11].

The unique characteristics of AuNPs to provide a large surface area

https://doi.org/10.1016/j.microc.2022.107585
Received 23 March 2022; Received in revised form 9 May 2022; Accepted 10 May 2022
Available online 11 May 2022
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for the loading of bioreceptors without losing their bioactivity and to facilitate electron transfer on the electrode surface have led to the widespread utility of these nanoparticles for improving the analytical performance of electrochemical sensors [12–14]. However, small sized AuNPs tend to form larger nanoparticle aggregates [15]. Thus, it is critical to prevent the agglomeration of particles using the appropriate capping (stabilizing) agents during the synthesis of AuNPs [16]. The suitable choice of stabilizing agent not only affects the shape and size of nanoparticles but can also play an important role in the electrochemical performance of AuNPs. The semiconductor carbon quantum dots (CQDs) or carbon dots (CDs) have been well documented as the stabilizing and capping (stabilizing) agents during the synthesis of AuNPs [16] . The semiconductor carbon quantum dots (CQDs) or carbon dots (CDs) have been well documented as the stabilizing and reducing agents [16,17]. They have been recognized as a novel generation of carbon-based nanomaterials with high solubility, low toxicity, great electronic features, large surface areas and lots of edge sites for functionalization [18]. An effective approach for overcoming the inherent limitations of AuNPs and CDs as the nanoplatforms in the fabrication of electrochemical biosensors is to combine them in the nanocomposite form [19]. These Au-C nanocomposites provide enhanced sensitivity and low LOD in the electrochemical nanobiosensors especially paper-based sensors [19].

Paper-based biosensors suggest low-cost substrates for detection of biological species in the clinical samples. Despite the unique advantages such as low-cost, disposability and biocompatibility, the electrical conductivity of the cellulose paper is not sufficient for use in electrochemical sensors [20]. A number of methods have been utilized to deposit conductive materials on a substrate including filtration, sputtering, printing methods, etc [21]. Here for the first time, a conductive coating was coated on the paper substrate by ion beam sputtering deposition (IBSD) method. Due to the monoenergetic and highly collimated ion beam, the obtained coating is durable, stable and high-quality films by this method compared to the vapor deposition technology [22]. According to the results obtained in this study, the formed GNC layer has a high capacity to increase the paper surface as well as increase its electrical conductivity.

2. Experimental

2.1. Reagents and chemicals

Dried borago (starflower) herb was prepared from Ramsar (Mazandaran, Iran) and powdered prior to use. Thiourea, bovine serum albumin (BSA), toluidine blue (TB), N-hydroxy succinimidine (NHS), disodium hydrogen phosphate (NaH2PO4), potassium ferricyanide (K3Fe(CN)6), sodium tetrachloroaurate (NaAuCl4), 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide (EDC), sodium dihydrogen phosphate (NaH2PO4) and potassium ferrocyanide (K2Fe(CN)6·3H2O) were also obtained from Sigma-Aldrich.

The interfering and target oligonucleotides, consisting the SARS RdRP gene (5′-C CAGGT GGAAC ATCAT CCGGT GATGC-3′), SARS-CoV-2 envelope gene (5′-ACAC TAGCC ATCCT TACTG CGCTT CG-3′), SARS-CoV-2 RdRP gene (5′-CAGGT GGAAAC CTCTG CAGGA GATGC-3′), and single-base mismatch strand (5′-CAGGT GGAAAC CTCTG CAGGA GATGC-3′) were purchased by Faza Pajooh (Tehran, Iran) according to the previous reports [23–26].

2.2. Fabrication of GNC/paper electrode

At the beginning of the experiment, the filter paper was oven-dried for 3 h at 105°C. Then, a thin film of GNC was coated on the paper surface by the sputtering of SIGRACELL® bipolar plates produced from high purity graphite (>99%, 12 cm × 15 cm). The main characteristic of natural graphite is high resistance to chemicals and good electrical conductivity. The specific angle between the paper and graphite plate was about 45°. The pressure was kept constant during deposition at 2 × 10⁻⁵ Torr by the introduction of high purity argon gas (99.999% purity). To achieve the desired pressure, the vacuum pump of 200 W rating should be run for approximately 1 h. During this time, no sputtering takes place and the power is only expended on creating a vacuum in the chamber. The ion current, accelerator voltage and deposition time were 25 mA, 2200 V and 45 min, respectively. After deposition process, the coated paper substrate was cooled in an argon atmosphere to room temperature. Finally, for preparation of working electrode, the GNC/paper was cut to 10 mm × 6 mm dimensions [27].

2.3. Green N, S-co-doped CD synthesis

Typically, the powered borage herb was utilized as a carbon source for preparation of N,S-doped CDs with thiourea as a nitrogen and sulfur source. 0.5 g of borago powder, 1.0 g of thiourea and 40 mL of distilled water were transferred into a Teflon-lined stainless steel autoclave and sealed. Then, the autoclave was put into a preheated (180°C) oven for 6 h similar to another reports in which citric acid or dopamine were used as raw materials under the hydrothermal route [28,29]. After that, the resulting brown solution centrifuged and filtered. Then, the solvent was eliminated by a rotary evaporator.

2.4. Production of Au@CD core–shell NPs

For preparation of a stock solution (2 mg mL⁻¹), the powder N,S-doped CDs were dis-solved into the distilled water. Then, the Au@CD core–shell NPs were synthesized following a procedure reported in the previous article with minor modifications [16]. Briefly, 1 mL of the N,S-doped CDs stock solution was added dropwise to 10 mL of NaAuCl4 solution (1.0 mM) at 95°C under vigorous stirring to form the reddish brown Au@CD core–shell NPs. The N,S-doped CDs as the mild reducing agents can be lead to the reduction of Au(III).

2.5. Development of signal-on genosensor

The Au@CD core–shell NPs were deposited on the GNC/paper electrode to obtained Au@CD NPs/GNC/paper. The obtained modified electrode was oven-dried for 3 h at 60°C. Then, a solution including 1 mM of EDC, 2 mM of NHS and 5 mM was dropped on the Au@CD NPs/GNC/paper electrode and remained for 30 min at room temperature. Finally, the modified paper-based electrode was dipped in the aminated cDNA (10 μM) solution for 1 h. In this step, the functional groups of carboxyl on the CDs were activated by an EDC/NHS cross-linker [30]. To decrease the non-specific binding, a 0.25% BSA solution was pipetted on the modified electrode and dried at room temperature.

The electrochemical sensing assay for RdRP detection is based on a signal-on mechanism. The cDNA/Au@CD NPs/GNC/paper electrode was exposed to different amounts of RdRP standards at room temperature for 40 min and then rinsed with distilled water to separate the unhybridized strands. To detect the DNA hybridization, TB as a heterocyclic aromatic dye which binds to nucleic acids was used in the genosensor fabrication. This hybridization indicator interacts electrostatically with negatively charged backbone phosphate groups. In addition, TB interacts with small grooves of dsDNA or ssDNA via hydrophobic effects, while it intercalates into the large grooves of dsDNA or dsDNA through π–π stacking and forms hydrogen bonds with bases in the major grooves [31]. So, TB can be used for the discrimination of hybridization reaction. It has stronger affinity with dsDNA compared to the ssDNA [31–33]. Here, 1 mM of TB solution was placed on the RdRP/Au@CD NPs/GNC/paper electrode and incubated for 20 min. Then, the voltammetry currents of the designed genosensor were taken in 0.1 M PBS (pH 7.4) for detection of target strands.

The differential pulse voltammetry (DPV) measurements were carried out using a PGSTAT 302 N electrochemical workstation (Autolab, Netherlands) in a three-electrode cell consisting of a Pt auxiliary
3. Results and discussion

3.1. Characterization of GNC/paper electrode

To determine morphology, the scanning electron microscope (SEM) images of pristine paper (Fig. 2A and 2B) and GNC deposited on the paper (Fig. 2C and 2D) were conducted with various magnification levels using an EVO 18 microscope (ZEISS, Germany). These images revealed the formation of graphite nanoflakes on the cellulose fibers of filter paper. In addition, the SEM study of deposited GNC film showed a morphology similar to that of nanocrystals synthesized in previous papers [34,35].

Raman spectroscopy, as one of the most common analytical techniques, determines the structure and the bonding nature of carbonaceous nanomaterials. Here, dispersive Raman microscope (DRM) system (Bruker, Germany) was applied for the characterization of GNCs. The confocal depth resolution of 2 µm allows individual layers of GNCs to be analyzed separately.

In general, the spectrum of all carbon nanomaterials consists of two Raman bands (D and G), except for natural graphite which has a single Raman active mode (G) at 1580 cm⁻¹. The G-band (1550–1570 cm⁻¹) is related to the stretching vibration of C–C (sp²) on unsaturated compounds such as aromatic rings or olefinic chains (both rings and chains) [36,37], while the disorder induced D-band (1370 cm⁻¹) as a breathing mode is strongly related to the six-membered aromatic rings [38]. Therefore, no rings, no D peak [39]. The analysis of Raman spectrum of as-prepared carbon nanomaterial indicates the two distinct bands D (1370 cm⁻¹) and G (~1600 cm⁻¹) with the intensity ratio (I_D/I_G) of 2 (Fig. 3). According to Ferrari’s amorphization trajectory (Three-stage model of the variation of the Raman G position and the I_D/I_G with enhancing disorder) [39,40], the produced nanomaterial which is in the end of stage 1 corresponds to graphite nanocrystals. The evolution of the Raman spectrum in stage 1 is as follows: D band becomes clear and I_D/I_G increases (inset of Fig. 3). Although the D peak is absent in ordered graphite, it appears with decreasing grain size and consequently disorder increasing. The crystallization of graphite leading to decrease in grain size [41]. In this stage, the G band position moves from 1580 cm⁻¹ to 1600 cm⁻¹. The G-peak wavenumber or frequency shifts upward as the grain size decreases [42].

3.2. Characterization of N, S-co-doped CDs

To survey the optical characteristics of N,S-doped CDs, the UV–vis absorption and fluorescence spectra of the as-synthesized CDs were taken on UV-1700 spectrophotometer (Shimadzu, Japan) and LS 55 photo luminescence spectrometer (Perkin Elmer, USA) at the room temperature, respectively. The UV–vis absorption of the as-synthesized N,S-doped CDs demonstrated a band at about 345 nm (Fig. S1A), which is characteristic of the CDs [43]. In addition, the maximal fluorescence emission intensity was detected at an emission wavelength of 412 nm and an excitation wavelength of 320 nm (Fig. S1B). The N,S-doped CD solution under a laser lamp appeared blue (inset of Fig. S1B), due to Tyndall effect [44]. To investigate the morphology and size of the N,S-doped CD solution, transmission electron microscope (TEM) image of CD solution was recorded on an EM10C instrument (ZEISS, Germany). As seen in Fig. S1C, the N,S-CDs were spherical with an average size of ~10 nm. The particle size of N,S-doped CD solution was also confirmed by detection of hydrodynamic diameter (D_h) using Nano ZS90 (Malvern Instruments, UK). The size distribution showed a sharp peak around 10 nm (inset of Fig. S1C). To identify molecular structure such as bond and functional group, the Fourier transform infrared (FT-IR) spectrum of the as-synthesized N,S-doped CD was recorded (Fig. S1D) using a Vector 22 spectrometer (Bruker Corporation, Germany). The absorption band at 3456.2 cm⁻¹ was related to the O–H bond stretching vibrations, while the strong peaks at 2850.5 and 2923.8 cm⁻¹ show the presence of CH₂ groups. These results confirms the chain structure of the N,S-doped CDs. The observed intense band at 1743.5 cm⁻¹ is due to the stretching vibration of C=O functional group. In addition, the characteristic absorption band at 1168.7 cm⁻¹ originates from C-N vibration.
The absorption at 1548.7 cm\(^{-1}\) is attributed to the skeleton vibration of C = N in the triazine ring \([45]\). The strong peak at 1639.3 cm\(^{-1}\) is assigned to the stretching vibration of C = C. The C-S stretching vibration is related to a weak peak at 621.0 cm\(^{-1}\) \([46]\). Other peaks at 715.5 and 1461.9 cm\(^{-1}\) reflect the absorption of C–H bond vibration. Our results are consistent with those of the previous published literature \([42]\).

3.3. Characterization of Au@CD core–shell NPs

The morphology of N,S-doped CDs was surveyed by TEM. As seen in Fig. S2A, the average size of quasi-spherical CDs was approximately 30 nm. The DLS intensity-based size distribution histogram confirmed this particle size. The DLS result showed an average \(D_H\) of ~ 30 nm (inset of Fig. S2A). Unlike N,S-doped CDs, a wide range of size distribution was observed for Au@CD NPs due to the use of a mild reducing agent. In general, strong reducing agents such as NaBH\(_4\) induces instant nucleation, while CDs as a mild reducing agent leads to producing the AuNPs with a wide range of size distribution \([16]\). The UV–vis absorbance spectrum of AuNPs was surveyed (Fig. S2B) and showed a characteristic surface plasmon peak at ~ 525 nm. This result suggests the formation of red AuNPs (inset of Fig. S2B) through the reduction of AuCl\(_4\) by N,S-doped CDs. The elemental composition of nanoparticles was investigated by EDX technique. The obtained spectrum indicates the presence of Au, S and N (Fig. S2C).

3.4. Characterization of Au@CD NPs/GNC/paper electrode

SEM was utilized to provide information concerning the morphology...
of Au@CD NPs/GNC/paper electrode surface. Fig. S3 showed that the described approach could lead to homogeneous formation of Au@CD NPs on the GNC modified electrode surface.

3.5. Electrochemical behaviors of different modified electrodes

The electrochemical response of electrode was investigated after each modification step using the cyclic voltammetry (CV) in the Fe(CN)$_6^{3-/4-}$ solutions as the electroactive indicator (Fig. 4A). As observed, the peak current of Au@CD NPs/GNC/paper electrode (curve b) increased significantly compared to the GNC/paper electrode (curve a). This confirms that the deposition of Au@CD NPs on the modified electrode increases the electrical conductivity and thus encourages the electron transfer rate [31,47]. So, the Au@CD core–shell NPs can greatly amplify the biosensor signals. The peak current clearly decreased when the cDNA was added to the electrode surface (curve c) due to the electrostatic repulsion. The negative charge of DNA strands caused by the presence of phosphate groups in their backbones leads to repel of signal probe from the electrode surface [46]. In addition, DNA as a non-conductive oligonucleotide chain creates an insulting layer on the electrode surface [31,48]. With the introduction of negatively-charged RdRP strands (12 nM), the peak current further increased (curve d), while the conjugation of positively-charged TB (1 mM) with DNA-DNA duplex led to dramatic increase in the peak current and negative shift of peak potential (curve e). The obtained result is explained by electrostatic attraction between TB and negatively-charged Fe(CN)$_6^{3-/4-}$.

Electrochemical impedance spectroscopy (EIS) is also a convenient approach for investigation of surface changes of electrode during the modification process. To support the CV results, the Nyquist plots were collected at a bias potential of 0.18 V (Fig. 4B). The impedance of the bare GNC/paper electrode was acquired (curve a), which represents a small semicircle. When the Au@CD NPs were casted on the electrode surface, the $R_{ct}$ reduced significantly (curve b) due to the high electrical conductivity of metallic nanoparticles. The immobilization of cDNA on the Au@CD NPs/GNC/paper electrode led to a considerable increase in $R_{ct}$ (curve c). The non-conductive cDNA strands acted as a mass-transfer blocking barrier, thus insulating the electrode surface and preventing the electroactive indicator from reaching the electrode surface [48]. After the hybridization of cDNA with the RdRP gene, the $R_{ct}$ indicated a further enhancement (curve d) due to the thickening of the insulting film and the electrostatic repulsion between dsDNA and signal probe. Finally, the $R_{ct}$ decreased significantly upon the interaction of electroactive TB with the dsDNA (curve e), as a result of electrostatic attraction between positively-charged TB and negatively-charged Fe(CN)$_6^{3-/4-}$.

3.6. Optimization of genosensor performance

To survey the influence of experimental effective parameters on the genosensor signal, the DPV current of the TB (1 mM) intercalated into the cDNA-RdRP (10 pM) duplex was measured under the following conditions: (I) deposition time varied in the range of 30 min to 1 h, while the accelerator voltage and the ion current were kept constant (2200 V and 25 mA, respectively); (II) the Au@CD NPs/GNC/paper electrode was dipped into the aminated cDNAs (bioreceptor strands) with different concentrations from 1 µM to 12 µM; (III) 1 mM of TB solution as the hybridization indicator was placed on the RdRP/cDNA/Au@CD NPs/GNC/paper electrode and incubated for different times from 5 min to 20 min.

The bar charts of DPV currents of the accumulated hybridization indicator on the modified electrode indicated the optimum desorption
time of 45 min (Fig. S4A). In less times, the graphite nanocrystals have not yet formed and the main material is in the form of graphite. At higher times, the G peak shifts to lower wavenumbers[49]. The blue shift of G band caused by reducing the number of ordered aromatic rings contributes to the passing from graphite nanocrystals to amorphous carbon[50]. With increasing desorption time from 45 min to 1 h, graphite nanocrystals were formed in a bed of amorphous carbon. So, the TB current decreased due to the reduction of electrode surface and electrical conductivity.

The peak current of hybridization indicator intensified with enhancing the concentration of immobilized cDNA strands on the surface of Au@CD NPs/GNC/paper electrode up to 10 μM (Fig. S4B). However, the higher amounts of cDNA strands resulted in the saturation of electrode surface and thus steric resistance[48].

The peak current of TB intercalated into the dsDNA enhanced with overtime from 5 to 20 min (Fig. S4C). Further increase of the accumulation time did not change the indicator signal.

3.7. Investigation of analytical figure of merits

The important figures of merit to be considered for validating SARS-CoV-2 RdRP biosensor were investigated by the DPV method in the potential interval of 0.4–0 V. To achieve this aim, the proposed genosensor was exposed to different concentrations of RdRP and then the hybridization between the immobilized cDNA probe and target gene was monitored by a redox-active indicator. As seen in Fig. 5A, a very small peak was recorded in the absence of target sequences due to the inter- action of TB (1 mM) with small grooves of probe ssDNA through hydrophobic effects. With the addition of RdRP sequences (0.5 pM) to the cDNA/Au@CD NPs/GNC/paper electrode, a significant peak was recorded because of TB intercalation into the large grooves of cDNA-RdRP duplex through Л-Л stacking. These observations show that TB has a stronger affinity with dsDNA compared to the ssDNA. The addition of different amounts of SARS-CoV-2 RdRP to the cDNA/Au@CD NPs/ GNC/paper electrode surface led to different increases in the peak current of the TB indicator. Based on such signal-on strategy, the developed genosensor showed the acceptable responses to different concentrations of RdRP gene in a dynamic linear range (DLR) of 0.50 pM-12.00 nM with a regression equation of \(I(\mu A) = 7.60 \log C_{RdRP}(pM) + 25.78\) \((R^2 = 0.9799(\sum = 6))\) (Fig. 5B). The limit of detection (LOD) and limit of quantification (LOQ) assessed as 3σb/m and 10σb/m were 0.15 pM and 0.50 pM, respectively.

The peak current stability of TB/RdRP/cDNA/Au@CD NPs/GNC/paper electrode was surveyed in 0.1 M PBS solution (pH 7.4). To achieve this aim, 5 repetitive DPV peaks of TB reduction were recorded. The relative standard deviation (RSD) for the voltammogram currents was calculated as 2.2%. Thus, this outcome obviously exhibited that the response of modified electrode is stable because of the high adhesion of GNCs on the paper substrate and the high stability of cDNA-Au@CD NPs biocoujugate.

For estimation of reproducibility, 5 modified electrodes were fabricated with the same protocol and utilized to measure 10 pM of RdRP gene. The RSD for 5 independent measurements was evaluated at 8.1%. The specificity of the cDNA sequence immobilized on the Au@CD NPs/GNC/paper electrode towards different sequences including SARS RdRP gene (100 pM), SARS-CoV-2 E gene (100 pM) and SARS-CoV-2...
RdRP gene (10 pM) was evaluated in the presence of TB (1 mM) using DPV method. Compared to curve a, a significant increase in TB current was observed for curve b, while negligible changes were recorded for curves c and d (Fig. 5C). The response of CDNA/Au@CD NPs/GNC/ paper electrode in the mixture of target (10 pM) and interfering (10-fold) genes indicated a slight increase compared with that of modified electrode for SARS-CoV-2 RdRP (curve e). These achievements confirm the genosensor selectivity towards the SARS-CoV-2 RdRP gene in the presence of SARS RdRP and SARS-CoV-2 E genes. In addition, the selectivity of the current genosensor in the presence of single-base mismatch strand was investigated in the SI.

### 3.8. Determination of RdRP gene in sputum samples

To validate the accuracy of this protocol, the SARS-COV-2 RdRP sequences were measured in the spiked human sputum samples. These samples were prepared from a healthy volunteer and centrifuged at 5000 rpm for 15 min. Then, the standard addition method or spike-and-recovery was utilized to detect RdRP genes in the diluted sputum samples. To achieve this aim, different amounts of SARS-CoV-2 RdRP gene (0.5 pM, 10.0 pM and 1000.0 pM) were added to the diluted sputum samples. Each spiked sputum sample was quantified 4 times and the obtained data were presented in Table 1. These outcomes showed the acceptable recoveries for the sputum samples (>95%).

#### 3.9. Performance comparison of the designed RdRP sensing device with the similar genosensors

For better comparison with the performance of other genosensors for the SARS-CoV-2 detection, Table 2 presents figures of merit of the previously reported nanomaterial-based genosensors and the current strategy [26,51–60].

Compared to the reported articles, the analytical performance of developed genosensor in this study is acceptable. Unlike the signal-off trends, our proposed genosensor does not suffer from limitations imposed by the background current [61]. It becomes clear that the performance of electrochemical biosensors depends on the electrode materials and nanoplatforms. In this study, GNC produced by IBSD technique not only increase the effective electrode surface area but also lead to greater sensor stability. These nanocrystals as the efficient interfaces possess excellent adhesion. In fact, the adhesion of coating is best when the carbon film is deposited by IBSD technique [22]. Another advantage of this biosensor is that the Au@CD NPs can be efficiently applied as the signal amplification nanoplatform. More importantly, the paper-based electrodes are very attractive components for the disposable, nontoxic, low-cost and flexible biosensors especially wearable biosensors which require characteristics of paper-based electrodes.

### Table 1

| Sputum samples | Added gene | Found gene | RSD (%) | Recovery (%) |
|---------------|------------|------------|---------|--------------|
| 1             | 10.0 pM    | 9.7        | 2.2     | 97.0         |
| 2             | 50.0 pM    | 48.9       | 1.7     | 97.8         |
| 3             | 500 pM     | 490.3      | 1.4     | 98.1         |

* Calculated as a mean of four measurements.

### Table 2

| Detection method | Format of biosensing strategy | LOD (pM) | DLR (10^-5 M) | Real sample | Total time | Ref. |
|------------------|-------------------------------|----------|---------------|-------------|------------|------|
| EC               | CPE-HT18C6(Ag)/CHT/SiQDs/PAMAM/cDNA/RdRP gene | 0.3 | 1.0 | Sputum | 35 min | [26] |
| FET              | Si/SiO2/RGO/AuNPs/cDNA/RdRP gene | 0.37 | 10 | Throat swab | 35 min | [52] |
| ECL              | DEP-Chips/GONCs/cDNA/RdRP gene | 3.99 | 10^-6 | Serum | 75 min | [53] |
| ECL              | DEP-Chips/GONCs/cDNA/RdRP gene | 186 x 10^-5 | 10^-10 | Sputum | 210 min | [54] |
| ECL              | SPE/Au@Fe3O4/HT-ORF1lab gene/GCN/AuNP-TL-AP | 3 | 10^-17 | Serum | 90 min | [55] |
| ECL              | SPE/Ni/MoS2NSs/GNP, CHT/AuNPs/S/ | 4.4 x 10^-5 | 1 | Artificial saliva | 110 min | [56] |
| ECL              | GCE/AuNP-g-C3N4/DNA1(Y1, Y2 and Y3)/Hairpin1/ Hairpin2/DNA-Ru/RdRP gene | 1.28 | 10^-5 | Pharyngeal swab | 180 min | [57] |
| ECL              | GCE/AuNP-g-C3N4/DNA1(Y1, Y2 and Y3)/Hairpin1/ Hairpin2/DNA-Ru/RdRP gene | 59 | 10^-5 | Pharyngeal swab | 120 min | [58] |
| ECL              | GCE/Nafion/Au@TiO2@Pt-Ru(dehyd1/2)/Hairpin DNA/Swarming arm/RdRP gene/Nb.BvCl/DNA-AgNCs | 0.21 | 1 | Serum | 14 min | [59] |
| LSPR             | siDNA-functionalized AuNls/Aln/RdRP gene | 0.22 | 10^-15 | Oropharyngeal swab | 10 min | [60] |
| NEC              | GDN/AuNPs or N gene | 0.18 | —— | —— | 75 min | This work |
| EC               | GCE/AuNP/sDNA/RdRP gene | 0.15 | 0.50 | Sputum | 90 min | [51] |

* LOD: Limit of detection; DLR: Detection limit range; Total time: Total time for the sample preparation and analysis; Ref: Reference.
4. Conclusion

In this study, the cDNA-functionalized Au@CD core–shell NPs were designed on the coated paper electrode for the SARS-CoV-2 RdRP detection. The deposited GNC film on the paper substrate allowed the development of a voltammetric genosensor by enhancing the electrical conductivity of substrate as well as increasing the effective electrode surface. Based on the stronger binding affinity of TB to cDNA-RdRP than to cDNA, this redox-active indicator was applied to monitor nucleic acid hybridization. With increasing RdRP concentration, the reduction peak current of TB enhanced in a linear range of 0.50 \(\pm\)12.00 nM. The developed signal on genosensor exhibited excellent LOD, excellent selectivity, good stability and high recovery for spiked spumut samples. Despite these strengths, the modification of electrode involves a series of time-consuming steps.

CRediT authorship contribution statement

Mohammad Ali Farzin: Investigation, Writing – original draft. Hassan Abdoos: Supervision, Project administration, Writing – review & editing. Reza Saber: Supervision.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.microc.2022.107585.

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