Impediometric Electrochemical Sensor Based on The Inspiration of Carnation Italian Ringspot Virus Structure to Detect an Attommolar of miR

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Electrochemical sensors are the tools to detect the accurate and sensitive miRs. There is the challenge to increase the power and sensitivity of the surface for the electrochemical sensor. We design a virus-like hollow structure of cuco$_2$o$_4$ that it holds the large amounts of p19 protein by mimicking of inherent virus (Carnation italian ringspot virus) to detect 21mir with the limit of detection (LOD = 1aM). The electrochemical measurements are performed between the potentials at $-0.3$ V and $+0.3$ V with 1 mM [Fe(CN)$_6$]$^{3-}/^{4-}$. After dropping the cuco$_2$o$_4$ on the SCPE (screen carbon printed electrode), the sensor is turned on due to the high electrochemical properties. Then, p19 proteins move into the hallow structure and inhibit the exchange of electrochemical reactions between the shells and the sensor is turned off. Then, adding the duplexes of RNA/miRs cause to increase the electrochemical property of p19 due to the change of p19 conformation and the system is turned on, again. So, for the first time, a virus-like hallow structure has been used to detect the 21miR in the human serum, MCF-7, Hella cells, with high sensitivity, specificity, and reproducibility in few minutes.

MicroRNAs are endogenous, single-stranded, non-coding small RNA with a length of ~22 nucleotide (nt). The alterations in the expression of these molecules have been linked to many diseases and cancers. So, these molecules have been important as diagnostic and prognostic tools in many cancers. A wide range of nanoparticles, nanorods, nanowires, nanoclusters, nanocomposites, etc, have been used in the electrochemical biosensor fabrication to detect of miRNA. Hollow nanostructures offer promising potential for advanced energy storage and electrochemical applications. The higher weight of the multi-shelled hallow nanostructures can cause to increase the energy density. They are designed with a hybrid of internal architectures such as core(s)-in-hollow-shell(s), multishell and multi-chambers. P19 protein is a suppressing factor in the silencing pathway as a common form of antiviral defense. This protein is used as a nanotechnology tool for micro RNA quantitation. It was demonstrated that p19 and other proteins staying in the virus based on the changes in the charges. There is some research about core-shell structures via virus mimicking for medical applications. In this research; we have constructed the nanomaterials based on the inspiration of the inherent structure of CIRV to use as biosensor applications. This structure consists of a mixture of metals such as cobalt and copper and oxygen (cuco$_2$o$_4$) which generate shells with positive and negative charges that they can create a strong bed surface. For the first time,

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these structures have been used in the design of electrochemical sensors to the identification of miR with high sensitivity. After dropping the CuCo2O4 hollow structures on the SCPE-GNP (screen carbon printed electron-gold nanoparticle) electrode, the chemical reaction has increased significantly due to the cobalt/copper-emitting conductive structure with inner shells and outer. Adding the p19 protein to this sensor cause to move the shells via pore by mimicking of CIR virus. These proteins prevent the electrochemical exchange of ions into the shell and outer space, and the sensor resistance is reduced and switched off. Then, we add a duplex of 21 miR/RNA which they moved into the shells and connect with p19 tightly. The change in p19 conformation induced more electrochemical reactions and the sensor is on, again. Because of the high p19 content, the sensitivity of this sensor is very high (1 μM). We use this sensor to detect 21miR in a real sample (human serum, MCF-7, Hella cells which they have the differential expression of 21 miR as model cases). So this virus-like sensor can identify the miRs with high sensitivity, reproducibility, and specificity.

**Experimental**

All of the chemical materials such as analytical grade potassium ferrocyanide, potassium ferricyanide, sulfuric acid, hydroponic, sodium chloride, and potassium chloride and cobalt and copper were purchased from chimeric co. Glutaraldehyde and Trizol were provided from nano alvand Co. Synthetic oligonucleotides were prepared by MW GE biotech, Ebersberg, Germany. We used as extraction RNA kit and commercially human serum from cinnamon Co (IRAN). The following sequences were used for RNA sensing experiments Probe: 5′-ACAACAUCAGUCGAACUGAU-3′ (21) miR-21; 5′-UAGCUUACUGACUGUUGC-3′ (RNA) mismatch RNA: 5′-GUA GACAAAGCGACUACCUG-3′ (RNA). P19 was prepared from New England Bio Labs Inc and used without further purification. P19 siRNA Binding Protein (10 units/ml) was stored at −20 °C16.

Screen-printed carbon electrode modified with gold nanoparticles on the ceramic substrate was purchased from DropSens Inc. This electrode consisted of a GNP-carbon working electrode; a carbon counter electrode and a silver reference electrode.

**Methods**

In this study, we used as SP-300 Instruments (SP-300) Texas to detect all of electrochemical analyzes such as electrochemical impedance spectroscopy (EIS) and differential pulse voltammetry (DPV). DPV measurements were performed in the presence of 1 mM [Fe(CN)6]3−/4− in phosphate Buffer (PS) buffer in the potential window −0.3 V to +0.3 V at a scan rate 50mVs1. The frequency range used for impedance measurement was 100 kHz to 1 Hz of [Fe(CN)6]3−/4− in PBS pH = 7.417. Analysis of EIS spectra was done using equivalent circuits using ZSimpWin 3.22 (Princetin Applied Research) and results reported in Nyquist plots. The AFM and TEM images were gathered using TecnaiG20 instruments from FEI Company and Hillsboro, USA. Particle sizes and zeta potentials were achieved from Horiba nanoparticle size analyzer, Malvern nano SZ-100 which uses green laser light at wavelength 532 nm.

Preparing CuCo2O4 as multi-shelled nanoporous materials. Synthesis of the onion-like nanoporous mixed copper-cobalt oxide hollow spheres was synthesized by a facile self-templating method. In a normal procedure, 0.060 mmol of Co(NO3)2.6H2O and 0.030 mmol of Cu(NO3)2.6H2O and 0.08 mmol of isophthalic acid (H2IPA) were added to a mixture of 10 mL N,N-dimethylformamide (DMF); acetone (1:1 v/v) to achieve a clear homogeneous solution by stirring for 6 h. The mixture was transferred into a teflon-lined stainless-steel autoclave and the reaction was kept at 160 °C for 4 h18. The resultant product was washed by ethanol several times and separated by centrifugation. The Onion-like CuCo2O4 hollow spheres are generated through thermal treatment in air at the temperature of 500 °C for 10 min with a heating rate of 5 °C min−1.

Fabrication the sandwiched of cuco2o4/p19/21miR-RNA on the SCPE/GNP. CuCo2O4 was prepared by dissolving in ethanol at the ratio 1:1, dried, rehydrated with buffer (pH = 7.4) and this mixture became a centrifuge at 10,000 rpm for 10 min to prepare supernatant. So, 1.2 μl of this supernatant was dropped on the surface of SCPE/GNP and stored at 4 °C for 1 h. The electrode was dried and carefully rinsed with water, dried again at room temperature, and then used for the electrochemical test. We use p19 to detect the sensitivity of miR. So, we use ten ml of 1:10 (v/v) dilute p19 protein solution were shaken at 37 °C for a 45 min. Then 0.5 μl of p19 was dropped on the cuco2o4/SCPE/GNP and incubated at 37 °C for 1 h. To identification the electrochemical behavior of p19 relative to the duplex of miR21-probe, we added the duplex of miR21/RNA. So, 6 mg/ml RNA probe and 3 mg/ml miR21 were blended in a vial containing Tris-EDTA Buffer (TEB). The target was put into the thermal shaker which was set to 65 °C and 250 rpm mixing speed and kept there for 1 h16. Then, 1 mM of 2 ml hybrid of miR21 was added on the surface of the electrode and incubated at 37 °C for 2 h in a dark without shaking. Finally, DPV and EIS measurements were carried between the potentials at +0.3 V and −0.3 V in PBS.

Identification of the specificity and sensitivity of the cuco2o4/p19/21SCPE/GNP sensor. Identification of the specificity of the sensor was done for three electrodes. So, three separate electrodes were incubated with 1 μM of the RNA probe (for detection miR21) with 1 μM of the miR-21 and 1 μM of RNA (mismatch) and one electrode as bare which they incubated in the buffer for 24 h at 4 °C under similar experimental conditions. DPV and EIS techniques were used for monitoring the electrochemical reactions in PBS buffer (pH 7.4) in the presence of the [Fe(CN)6]3−/4− redox couple. To evaluation the sensitivity of miR, increasing concentrations (1 μM to 500 μM) of miR21, were incubated with 1 μM of RNA probe in a 100 μL total volume of PBS at 37 °C for 1 h after incubation of 10 μL of the reaction mixture with cuco2o4/p19 sensor for 1 h at room temperature, and then washing 3 times with 500 μL of phosphate buffer (PB) and electrochemical reactions were done16. To examine the effect of the amount of p19 in limit of detection of 21 miR, we dropped 0.4 μl of p19 on the sensor with the increasing concentrations (1 μM to 500 μM) as the same of conditions.
Optimization of the cuco$_2$O$_4$/p19/SCPE/GNP sensor. In this study, we determinate the electrochemical power of the cuco$_2$O$_4$ hollow structure in the sensor, and also the reproducibility of the sensor because of staying of p19 into the cuco$_2$O$_4$ structure by mimicking the native virus-like structure. To analysis the stability of this sensor, DPV was done in one electrode for 30 cycles, and the change of behavior was recorded.

Cell culture and RNA extraction. We used as metastatic breast-cancer cell lines (MCF-7) to determine the miR-21 levels and Hella cell (a human cervix-adenocarcinoma cell line) as a control cell with no increase 21miR. MCF-7 is shown as estrogen-receptor-positive and Hella cell used as negative control cell lines, respectively. Cell lines including MCF-7, MCF-10A, Hella cells were cultured with 1% penicillin/streptomycin and growth in 5% CO$_2$ at 37°C. After collection of $10^7$ cells of conditioned medium, they centrifuged at 2000 $\times$ g for 30 min to eliminate cell debris. For RNA extraction of these cells, $\beta$-mercaptoethanol was used to the lysis to protect against the RNase activity. Then, we isolated the total RNA by purification kit (Norgen BioTek Corp., Canada) according to the protocol. The sample was divided two sample volumes of the lysis solution and 0.2 mL of the separation matrix. Next, we vortexed the mixture for 30 s, and incubated at 60 °C for 10 min. Three volumes of 99% ethanol were added to blending for 30 s. Subsequently, the mixture was centrifuged. The binding solution was then added to the pellet and incubated at 60 °C for 10 min. After adding 0.3 mL of 99% ethanol, the slurry was washed of the bound RNA to remove the remaining proteins and other impurities. The purified total miRNA was then eluted in 60 $\mu$L RNase free water.

Results

Electrochemical behavior of the cuco$_2$O$_4$/p19/SCPE/GNP sensor. Cuco$_2$O$_4$ structure showed the different electrochemical behavior in presence of 1 mM [Fe(CN)$_6$]$^{3-}/4-$ redox probe in PBS buffer on the GNP-SCPE (Fig. 1A,B). Bare electrode showed the $\Delta$E(Epa -Epc) 78 mV in the [Fe (CN)$_6$]$^{3-}/4-$ redox probe. The electrochemical behavior of cuco$_2$O$_4$ showed the peak currents decreased from 14.2 to 13.5 mA and the $\Delta$Ep increased from 78 to 81 mV, due to the facilitate electrolyte penetration into the interior surfaces by the nanoporous shells (Fig. 1B). This cuco$_2$O$_4$ structure can provide more active sites for electrochemical reactions by nano pores and a high specific surface area that they can facilitate the access of the electrolyte. In the previous studies, the addition of p19 protein to the substrates was via synthetic or non-synthetic linkers. For example, in the ramnani’s research showed that the binding of p19 on the CNT/FET surface was via PBASE which it caused to reduce the resistance of the electrochemical substrate by covering the electrochemical reactions. In this study, a virus-like cuco$_2$O$_4$ structure mimics of the intrinsic structure of CIRV. After adding p19 on this sensor, peak current increases from 13.5 to 14.1 mA and the $\Delta$Ep decreased from 81 to 70 mV (Fig. 1B). The interesting point of our study is the placement of the p19 into the cuco$_2$O$_4$ structure from pores by mimicking the CIR virus. In our study, the cuco$_2$O$_4$ pores were the hot-spot site for electrochemical exchanges between the internal and external charges. So the large amounts of p19s pass through these pores. The pH-induced in the core-shells can play a role to retain p19 in the space of shells. On the other hands, placement of p19 can inhibit the path of electron and decrease the resistance. The electrochemical behavior of the cuco$_2$O$_4$/p19/SCPE/GNP was examined by electrochemical impedance spectroscopy (EIS), which was an effective method for the characteristic of surface-modified electrodes. Nyquist
The current concentration curve showed a linear relationship in the range of 0.1 μl with a sensitivity of 0.2 μl linearly with increasing the concentration of miR-21 ranged from 100 fM to 1aM when it was used as a sensor in this stage is very high, it may become saturated in every CuCo2O4 structure. Therefore, it can connect with frequencies varying from 0.1 Hz to 100 kHz. As shown in Fig. 1A (a), the bare electrode illustrated a Rct value of approximately 210 Ω. When CuCo2O4 was added on the electrode, the Rct value increased by about 330 Ω (Fig. 1A(b)). Rct value was continuously increased to 380 Ω when the p10 protein was immobilized on the electrode surface. These results were in good agreement with that of differential pulse voltammetry.

**Model of sensing 21miR in the CuCo2O4/p19/SCPE/GNP sensor based on the virus-like structure.**
Adding duplex of 21miR/RNA on the CuCo2O4/p19/SCPE/GNP showed the ΔE(Epa-Epc) 92 mV in the [Fe(CN)6]3-/4 redox probe (Fig. 1B). The electrochemical behavior of CuCo2O4 showed the peak currents decreased from 14.1 to 12.1 mA and the ΔE increased and the Ret value increased by about 400Ω due to the stable interactions of duplexes of 21miR/RNA with p19, followed by the change of p19 conformational and enhancement the electrochemical reaction (Fig. 1A). Recent studies showed that the pH-dependent mechanisms can cause to maintain the proteins in the virus shell. So, the micro RNA sensing mechanism can be explained by the virus-like model due to enhance the electron transfer of the change in p19 conformation with the redox probe [Fe(CN)6]3-/4 to the electrode surface when it connected with the miR/RNA duplex. Since the resistance of the sensor in this stage is very high, it may become saturated p19 in every CuCo2O4 structure. Therefore, it can connect with the duplexes of 21miR/RNA. Results of the unit value of the electrochemical tests were reported in the different stages in Table (s.1). On the hands, there are many CuCo2O4 structures on the SCPE-GNP which they can behave as electrochemical reservists (Fig. 4).

**Surface morphology of the sandwiched CuCo2O4/p19 nanocomposite.**
The morphology of bimetallic (CuCo2O4) was examined by field-emission scanning electron microscopy (FESEM) and transmission electron microscopy (TEM). Figure 2A,B show (FESEM) image of the microspheres (CuCo2O4). The morphology and detailed structure of the samples were observed using the TESCAN Mira3 XMU field emission scanning electron microscope (FESEM) at accelerating voltage from 1-30kV and Philips CM30 transmission electron microscope (TEM) at 300 Kv.

Figure 2. (A,B) FESEM and (C) TEM images of morphology of bimetallic (CuCo2O4). Figure A and B show (FESEM) Image of the Microspheres (CuCo2O4). The morphology and detailed structure of the samples were observed using the TESCAN Mira3 XMU field emission scanning electron microscope (FESEM) at accelerating voltage from 1-30kV and Philips CM30 transmission electron microscope (TEM) at 300 Kv.
A regression equation of \( y = 14.476x + 23.475 \) \((R^2 = 0.976)\) was obtained, where \( y \) is the \( j \) value in \( \mu \text{A cm}^{-2} \) and \( x \) is the logarithmic concentration of 21miR in \( \text{aM} \). As shown in Fig. 3B the relative standard deviation (RSD) values were between 9% and 0.1% and the limit of detection (LOD) was 1 aM. It was done from 3(Sb/m), where Sb is introduced as the standard deviation of the measurement signal for the blank and m detected the slope of the analytical curve in the linear region. On the other hand, with the concentration of 0.4 \( \mu \text{l} \) p19 at the same range of the concentration of miR-21(100 fM to 1aM), there is no change in the limit of detection (Fig. 3C,D). Results showed in Fig. 3C that the \( j \) value increases linearly with increasing the concentration of miR-21 ranged from 100 fM to 1 aM when it was used as 0.4 \( \mu \text{l} \) p19. A regression equation of \( y = 13.876x + 23.548 \) \((R^2 = 0.976)\) was obtained.
where $y$ is the $j$ value in $\mu L/cm^2$ and $x$ is the logarithmic concentration of miR21 in aM. As shown in Fig. 3D the relative standard deviation (RSD) values were between 8% and 0.3% and the limit of detection (LOD) was 1 aM.

The best reason for these results is the saturation of p19 in the low concentration with a duplex of 21miR/RNA. In the past research about p19-based biosensor, the limit of detections was the most of the time <1fM, when the concentrations of p19 were 0.5 $\mu L/cm^2$. These results may be due to the qualification of p19 connection with RNA/miR. Because of the p19 is stayed into the cuco2o4 structure in this model, the probability of the connection between the duplex and p19 is more than other p19-biosensors when they fix on the sensor (Table 1).

### Table 1. Comparison of sensitivity of the different of liposomal substrates in electrochemical biosensor.

| Lipid–platforms | Materials/linker | specificity | sensitivity | reproducibility | Complicated design | Ref |
|-----------------|-----------------|-------------|-------------|----------------|-------------------|-----|
| Electronic Detection of MicroRNA at Attomolar Level with High Specificity | CNT-FET-p19/PBASE as linker in bed | ++ | ++++ | ++++ | ++++++ | 26 |
| Magnetobiosensors Based on Viral Protein p19 for MicroRNA Determination in Cancer Cells and Tissues | Magnetic-p19 Not attach in bed | ++ | +++ | +++ | +++ | 27 |
| Three-Mode Electrochemical Sensing of Ultralow MicroRNA Levels | AuNP/P19 Not attach in bed | +++ | ++++ | ++ | ++++++ | 28 |
| Sequential or Multiplex Electrochemical Detection of miRs Based on the p19 Function Relative to three Sandwiches of Different Structural Hybrids on the Liposomal Sensor | DOTAP-DOPE/p19 Not attach in bed | +++ | ++ | ++ | ++++++ | 4 |
| exosomal bed by using multi covalent attachment p19 | Esosome/p19 Attach in bed with no linker | ++ | ++++ | ++++ | +++ | 4 |
| A new insight into electrochemical microRNA detection: A molecular caliper, p19 protein | P19/ Not attach in bed | ++ | +++ | ++ | ++ | 29 |
| A New Insight of Virus-like Hallow Structure in Electrochemical Biosensor to Detect an Attomolar of miR | P19/ preserved into the hallow structure | ++ | ++++ | ++++ | + | This study |

Figure 4. Schematic drawing of Detection of 21 miRs by the cuco2o4/p19/SCPE/GNP sensor mimicking CIR virus. (A) A fabrication of cuco2o4 hallow structure in the SPCE-GNP caused an increase in the resistance measured by EIS. (B) Adding p19 into the shells of cuco2o4 caused a decrease in the resistance measured by EIS. (C) Adding duplexes of 21miR/RNA caused an increase more resistance range.
Optimization of the reproducibility and stability of the cuco$_2$O$_4$/p19/SCPE/GNP sensor. We evaluate the reproducibility of the cuco$_2$O$_4$/p19/ miR/SCPE/GNP sensor with ten different modified sensors which were prepared to detect 0.1 nM of 21miR target sequence. As shown in Fig. 5(A), the results of relative standard deviation of measurements for the 10 repeats showed the 4.11%, which confirmed the high reproducibility for them. To detect the stability of the hallow structure biosensor, the cuco$_2$O$_4$/p19/ miR/SCPE/GNP sensor
was first stored in the refrigerator at 4°C for over 4 months and evaluated via EIS after its hybridization with 0.1 nM RNA/21miR sequence. In Fig. 5(B), we showed that there were the same signal occurred for 60 days. Afterward, the DPV signals was significantly decreased to 50.9%, and 90.2%, which corresponded to 90 days, respectively. We haven’t seen the signal response when this sensor was stored for 111 days because the biological activity of the probe might be destroyed.

**Electrochemical detection of miRNAs in real samples.** We used cuco_{2o}/p19/SCPE/GNP to detect 21miRs in commercially human serum and MCF-7 / MCF-10A cells / HeLa cells. Before DPV experiments, an excess of yeast tRNA was added to a human serum sample to protect the endogenous miRNAs from nucleases. We used 1–5 mg of total raw RNA (RNA) isolated from these cells. The hallow structure biosensor response was significantly different for MCF-7 and MCF-10A cells, human RNA, thus indicating that miR-21 is highly expressed only in the metastatic cell lines (MCF-7), and not in the MCF-10A and HeLa cell lines (Fig. 6A). Based on the previous findings, miR-21 was overexpressed in MCF-7 cell line relate to the expression in the MCF-10A cell line. On the other hand, we optimize the amount of RNA with amperometric signal. The best amount of extracted RNA from cells was about 0.5 mg (Fig. 3). Our result obtained for five different samples as above described. RNA extracts were (22.1) fmol of miR-21 per microgram of RNA, extracted from MCF-7 cells (RSD n = 5 = 12.6%). We also evaluated the discrimination efficiency towards the tested sequence 1 m by spiking RNA (1.0 mg) extracted from MCF-10A cells and with the synthetic target (2 nm; ca. 18 fmol) and 1 m oligonucleotides (2 nm; ca. 18 fmol) and MCF-7 cells RNA with the (17.4) fmol of miR-21 per microgram of RNA and 16.1 fmol of miR-21 per microgram of RNA extracted from HeLa cells. Results show the best signal detection of cuco_{2o}, biosensor is for human serum RNA > MCF-7 cells RNA > Synthetic 21 miR > MCF-10 cells RNA = HeLa cells (Fig. 6C). The impedance spectra of the electrochemical tests were reported in the different cells and human cells are in Fig. 6B.

**Discussion**

The large studies were researched about the design of the electrochemical sensors for identification micro RNAs. The design of appropriate beds is important to provide a good response in the electrochemical reactions due to the low amount of miRs in the body fluids. P19 protein is an important nanobiotechnology tool that increases the sensitivity of miR detection in the electrochemical sensors. In the recent research, this protein is bound to the electrode via a linker on the electrode, so the probability of connection with the duplexes of RNA/miR is being challenged in every step. So, p19-biosensors shows the low sensitivity especially in the real samples. In our research we used as a structure which it prompt the sensitivity of p19. For the first time, we designed the virus-like hallow structure which it can increase the intensity of electrochemical reactions. The placement of various copper metals along with cobalt has provided many conductive and electrochemical properties. On the other hand, every hallow structure can mimic of virus-like CRIV and provide the reservoir of p19 proteins that they can connect with the more of duplex of RNA/miR without any linkers. Our sensor shows the best sensitivity (LOD = 1 μM) related to the sensitivity of other p19-biosensor. On the hand, we evaluated this sensor in the real samples and we get the same acceptable results. These structures can open a new window for a wide range of medical purposes. As a result, the sensor has been able to detect miR with very high sensitivity and stability and high repeatability. In the next research, the other hallow structure should be studied as biosensor applications in the diagnosis and therapy.

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Author contributions
We confirmed that E. Ghazizadeh and negin daneshmand wrote the main manuscript text and Seyed Ebrahim Mosavifarad analysis about electrochemical reactions and Saeid kamari kaverlavani prepared figures and tables. We confirm that Seyed Ebrahim Moosavifarad is also co-corresponding, too.

Competing interests
The authors declare no competing interests.

Additional information
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