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Peptide-based simple detection of SARS-CoV-2 with electrochemical readout

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Highlights

- First peptide-based impedimetric biosensor for Spike protein detection in only 15 min.
- Direct immobilization of synthetic peptide on SPAuE electrodes.
- No cross-reactivity with Spike proteins of MERS and SARS-CoV.
- High selectivity towards RBD protein in S1 region of target transmembrane protein.
- Sensing of the target in spiked buffered solutions, viral particles and clinical samples.

Abstract

Coronavirus disease 2019 (COVID-19) caused by Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-CoV-2) is considered one of the worst pandemic outbreaks worldwide. This ongoing pandemic urgently requires rapid, accurate, and specific testing devices to detect the virus. We report a simple electrochemical biosensor based on a highly specific synthetic peptide to detect SARS-CoV-2 Spike protein. Unlike other reported electrochemical biosensors involving nanomaterials or complex approaches, our electrochemical platform uses screen-printed gold electrodes functionalized with the thiolated peptide, whose interaction with the Spike protein is directly followed by Electrochemical Impedance Spectroscopy. The electrochemical platform was Spike protein concentration-dependent, with high sensitivity and reproducibility and a limit of detection of 18.2 ng/mL when tested in Spike protein commercial solutions and 0.01 copies/mL in lysed SARS-CoV-2 particles. The label-free biosensor successfully detected the Spike protein in samples from infected patients straightforwardly in only 15 min. The simplicity of the proposed format combined with an on-demand designed peptide opens the path for detecting other pathogen-related antigens.

Article history:
Received 14 December 2021
Received in revised form 15 February 2022
Accepted 16 March 2022

Keywords:
SARS-CoV-2
Synthetic peptide
Impedimetric biosensor
Infectious disease

1. Introduction

The rapid expansion of the coronavirus 2019 (COVID-19), a disease caused by the Severe Acute Respiratory Syndrome coronavirus 2 (SARS-CoV-2), was declared a pandemic in 2020 and continues today to threaten public health systems worldwide [1–4]. The current diagnostic recommended by the World Health Organization (WHO) is based on the detection of the SARS-CoV-2 viral RNA by reverse transcriptase real-time polymerase chain reaction (RT-PCR) [5]. Although the RT-PCR is the gold standard method, it is time-consuming and requires specialized testing facilities with highly qualified personnel, which decreases the high number of daily
samples essential to slow its spread. Besides, immunoassays based on the detection of viral structural proteins and serological methods that detect antibodies have some limitations related to low sensitivity, reduced efficiency and cross-reactivity. Therefore, developing accurate, specific and sensitive devices for the simple and rapid detection of the SARS-CoV-2 and early diagnosis of COVID-19 has a pivotal role in controlling the viral outbreak [5–7].

Previous reports have found that Electrochemical Impedance Spectroscopy (EIS)-based biosensors rapidly detect bioreceptors associated with SARS-CoV-2 infection, with high sensitivity, specificity and reliability [8–17]. In such biosensors, bioreceptors are anchored on the working electrode surface to directly bind to target molecules specifically, generating changes in the interfacial properties in a concentration-dependent manner [18–20]. The SARS-CoV-2 has also been detected electrochemically by immuno-sensors [14,21–32] and genosensors [33–36]. Furthermore, outstanding features of biosensors have been exploited in the detection of other viruses [37,38], pathogens [39], other infections [18], cancer biomarkers [40–42] and other diseases-related biomarkers [43] demonstrating to be promising analytical tools that can help to solve current diagnosis limitations [44].

Here, we report the first peptide-based biosensor for simple monitoring of SARS-CoV-2 (Spike protein) and SARS-CoV-2 viral particles in COVID-19 positive patients by EIS. Peptides are specific sequences of amino acids with similar selectivity, specificity and chemical nature as proteins but smaller sizes, higher stability against denaturation, cost-effectiveness, accessibility, easiness of modification, and more extensive chemical versatility [45,46]. Therefore, peptides are ideal candidates to substitute proteins as bioreceptor in (bio)recognition and biosensing events. The biosensor uses a synthetic thiolated peptide bioreceptor chemisorbed at the working electrode of a screen-printed gold electrode (SPAuE) that directly interacted with the Spike protein and whose interaction was detected and quantified by EIS. The resultant device showed high sensitivity and reproducibility, low limit of detection (LOD) and a linear dynamic range of clinical relevance. Besides, the peptide-based biosensor was highly specific for SARS-CoV-2 and detected the Spike protein from COVID-19 positive patients in clinical samples. This is the first time the peptide has been incorporated in a transducer platform for SARS-CoV-2 sensing purposes of remarkable analytical performance. Performance, simplicity, and low cost not only underline the great potential of our biosensor for detecting the virus at the clinical level but for the on-demand design of peptide-based sensing platforms for detecting other viruses, pathogens, and disease biomarkers.

2. Materials and methods

2.1. Reagents

A targeting synthetic peptide with 23 amino acids length, (MPA)[PEG4]IEEQAKTLDKFNHAEDLFQYS sequence, purity ≥95%, MW: 3137.42 g/mol and a thiol-linker was synthesized on-demand by GenScript U.S.A. Co. (New Jersey, U.S.A.). SARS-CoV-2 (2019-nCoV) Spike recombinant protein (protein S, purity 90% by SDS-PAGE) (MW: 76.5 kDa) was acquired by SinoBiological (Pennsylvania, USA); recombinant human beta-1,4-galactosyltransferase-5 glycoprotein (β-1,4-GALT-5) (cod Ab160437) was purchased from Abcam (Cambridge, UK); receptor binding domain Spike protein (RBD) was acquired by Active Motif (California, USA); potassium ferricyanide ([Fe(CN)6]3−/4−), potassium hexacyanoferrate (II) tribromide (K3[Fe(CN)6]3·H2O), tris(2-carboxyethyl) phosphine hydrochloride (TCEP) and 6-mercaptop-1-hexanol (MCH) were purchased from Merck Millipore (Darmstadt, Germany). Dipotassium hydrogen phosphate (KH2PO4), disodium hydrogen phosphate (Na2HPO4) and potassium nitrate (KNO3) were acquired from PanReac AppliChem (Darmstadt, Germany). Potassium dihydrogen phosphate (KH2PO4), potassium chloride (KCl) and sodium chloride (NaCl) were obtained from J.T.Baker® (Xalostoc, Mexico). Sulphuric acid (H2SO4) was purchased from Honeywell FlukaTM (Seelze, Germany). RIPA (Radioimmunoprecipitation) lysis buffer system composed of protease inhibitor cocktail in dimethyl sulfoxide (DMSO), 100 mM Sodium orthovanadate in water, 200 mM phenylmethylsulfonyl fluoride (PMSF) in DMSO and 1X lysis buffer pH 7.4 were acquired from ChemCruz (Dallas, U.S.A.). Sterile and nuclease-free water was acquired from VWR Life Science (Ohio, USA) used to dilute each sample. All reagents and commercial samples were used without purification and the solutions were prepared using deionized water of 18.3 mΩ cm.

2.2. Devices and equipment

All electrochemical measurements were performed with a three-electrode cell configuration SPAuE (ref. 220BT, from Metrohm) in a potentiostat galvanostat VersaSTAT 3 with a single channel with an impedance mode. The spectra were analyzed using VersaStudio 2.61.3 software. The chips consist of a 4 mm gold working electrode, a gold counter electrode, and a silver pseudo-reference electrode, respectively printed on the same strip.

2.3. Assembly of the peptide-based biosensor

SPAuEs were pre-treated according to the previously reported protocol to clean and activate the surface, i.e., cyclic voltammetry (CV) in 50 μl of 0.1 M H2SO4, scanning at a potential between 1.6 and −0.2 V (vs. Ag pseudo-reference electrode) at a scan rate of 0.1 V/s until the voltammogram becomes stable (approximately 10 consecutive cycles). The effective area and roughness factor were estimated by considering that the gold surface’s reductive peak has a charge of 400 μC/cm2 [47] and the gold electrode has a geometry area of 12.56 mm2.

1.59 mM of synthetic peptide stock solution was prepared by dissolving 1 mg of the peptide in 200 μl of nuclease-free water. TCEP was added at a final concentration of 0.5 mM to reduce thiol groups from the peptide that may be oxidized by incubation at 37 °C and 1100 rpm for 30 min. The pre-treated SPAuE was incubated in 6 μl of the TCEP-pretreated peptide solution for 1 h at room temperature, followed by 6 μl 0.01 mM MCH for 10 min to orient the peptide reactive places and block nonspecific sites. The modified SPAuE was then incubated in a Spike protein solution at 37 °C and 50 rpm for 15 min in a wet chamber in a Thermo Scientific MAXQ 4450 shaker for the peptide-protein interaction. The peptide biosensor assembly steps and detection mechanism are illustrated in Scheme 1.

2.4. Electrochemical measurements

The electrochemical performance of the as-functionalized platform was evaluated by CV in the 5 mM [Fe(CN)6]3−/4−/KNO3 redox pair in a potential window between +0.4 V and −0.2 V, at a scanning rate of 0.05 V/s for 5 consecutive cycles. This measurement was made initially with the bare electrode and later after each step of the functionalization process.

Electrochemical impedance spectroscopy (EIS) was carried out in the 5 mM [Fe(CN)6]3−/4−/KNO3 redox pair by applying a potential of +0.19 V in frequencies ranging from 50 kHz to 0.05 Hz at 0.010 V of amplitude. The impedance data were fitted with the EIS Spectrum Analyzer software using the Levenberg-Marquardt algorithm. The impedance measurements were performed after each step of the biosensor development, i.e., functionalization of the SPAuE
surface by chemisorption of the peptide, blocking the electrodes with MCH and incubation with Spike protein. Calibration curves were built with increasing concentrations of Spike protein and SARS-CoV-2 viral particles and the limit of detection (LOD) was calculated as LOD = 3σ_{blank}/slope.

2.5. Selectivity and specificity studies and quantification of spike protein in human samples

The selectivity of the biosensor was assessed by comparing the response against supernatants of the target (SARS-CoV-2 viral particles inactivated by UV-light) and supernatants of other coronaviruses such as Middle East Respiratory Syndrome (MERS) and SARS-CoV at equivalent concentrations. We also evaluated the specificity of the peptide-based biosensor against region binding domain (RBD) protein solutions and glycoproteins such as beta-1,4-galactosyltransferase-5 (b-1,4-GALT-5).

All viral supernatant samples were prepared at a concentration of 10^6 copies/mL dissolved in a lysis buffer containing 10 μl of PMSF solution, 10 μl of sodium orthovanadate solution and 20 μl of cocktail solution of inhibitor protease per ml of 1X RIPA lysis buffer. The samples were vortexed for 1 min and then sonicated for 1.5 min to promote the disruption of the nuclear membrane and favor the shedding of Spike protein molecules on the surface of the viral particles. The lysed solutions were dropped on the modified electrodes and incubated at 37 °C and 50 rpm for 15 min in a wet chamber. The biosensor response was interrogated by EIS, as mentioned above. Paired t-test and a 1-way ANOVA with a 99% level of statistical significance (ANOVA) were performed using Excel software to evaluate the statistical significance between the samples.

Nasopharyngeal swab clinical sample solutions in universal transport medium (UTM) from three COVID-19 negative and nine positive patients were donated by the Tropical Diseases Study and Control Program (PECET - by its Spanish acronym) laboratory after testing by RT-PCR in a biosafety lab (BSL)-2 facility following established biosafety and ethical standards and inactivation by UV-light. The RT-PCR protocol is briefly described in the next subsection. Before analyzing the samples with the peptide-based biosensor, they were lysed according to the protocol mentioned above. It facilitated the viral protein release and diluted 100-fold its concentration and interferent species from the samples to fit well in the calibration curve.

2.6. Quantification of viral RNA from clinical samples by RT-PCR

Samples were obtained by nasopharyngeal swabs and immediately deposited in a UTM, followed by RNA extraction from 200 to 300 μl of samples on the same day. The extraction was done by automated extraction using the king Fisher Flex robot and the MagMAXTM Viral/Pathogen II (MVP II) Nucleic Acid Isolation kit (Thermo Fisher) or using a manual extraction with the Quick RNA Viral Kit (Zymo Research). The amplification reaction was done using the Berlin protocol, with modifications as detailed in the Supporting Information section (SI). The Human Research Bioethics Committee from the University of Antioquia governed by Resolution 008430, October 4, 1993, Ministry of Health from Colombia endorsed the project ‘Nanobiosensors for rapid detection of SARS-CoV-2’ in an approval certificate number 20-109-897.

3. Results and discussion

3.1. Development of the detection platform

We selected a thiolated synthetic peptide as a bioreceptor to develop the biosensor reported herein, inspired by the high interaction between the RBD region of the SARS-CoV-2 and the
angiotensin-converting enzyme (ACE)-2 host receptor protein [48,49]. Recent crystallographic studies revealed that the amino acid residues, mainly through polar residues present in the z1 helix of the peptidase domain of the ACE-2 protein, are the key to the interfacial interaction with the RBD protein of SARS-CoV-2 [4,50]. It has been reported that multiple variants of the virus are related to variations in the hydrogen bonds and amino acid residues involved in the RBD region-ACE-2 protein interaction. However, we propose a peptide-based biosensor concept that can be readily re-designed on-demand based on new changes of the amino acid content of the virus by conserving a similar length chain, terminal functional groups, and optimized functionalization conditions before assembling in a device.

Therefore, a synthetic peptide was designed considering the previously reported peptide sequence [4] but changing the N-terminus of the peptide for a thiol group to anchor the peptide to the SPE surface. The amino acids of length and composition are homologous to the sequence of the SPE surface. The amino acids of length and composition are homologous to the sequence of the z1 helix of the peptidase domain of the ACE-2 protein that expects to interact specifically with the RBD while minimizing unspecific interactions with other proteins. The peptide-modified electrode detected the Spike protein expressed at the outermost surface of viral nanoparticles in a spiked buffered solution and lysates from SARS-CoV-2 viral particles, as shown below. The structure of this synthetic peptide consists of 23 amino acids that specifically recognize the Spike protein, a segment of polyethylene glycol (PEG) as a spacer and a C-terminal modification of mercaptopropionic acid (MPA) to link to the transducer platform. The synthetic peptide is highly specific for a side branch of the RBD region of the SARS-CoV-2 Spike protein [51]. We assembled the impedimetric biosensor by chemisorbing the thiolated peptide on an SPAuE surface previously activated by CV in acidic conditions. The activated electrodes showed an effective area and roughness factor of 231.62 ± 11.42 mm² and 18.44 ± 0.91, respectively. An efficient, reproducible, and well-controlled self-assembled monolayer of the peptide was formed at the SPAuE surface, followed by treatment with MCH characterized by CV and EIS. Incubation with MCH allowed to block nonspecific binding sites on the SPAuE surface and better orient the synthetic peptide perpendicular to the surface to point to the RBD region of the Spike protein [52–54].

Fig. 1 shows comparative CV and EIS plots from the step-by-step electrode modification process. For example, Fig. 1A shows the current intensity decreased after chemisorbing the peptide at the SPAuE due to the formation of a peptide layer that limits the electrons transfer at the electrode-solution interface as compared with the bare electrode. Yet, the incorporation of MCH favored the interaction with the Spike protein noticeably. Fig. 1B shows differences in electron transfer of the transducer platform after each modification step represented by the Nyquist plot (-Z” vs. Z”). Fitting electrochemical parameters from EIS by a Randles equivalent circuit (Inset Fig. 1B) was used to estimate the charge-transfer resistance (Rct), the electrolyte resistance (Ron), the constant phase element (CPE), which depends on a pre-exponential factor (P) and an exponent (n) [55–58], as summarized in Table 1.

The EIS from the bare electrode in inset Fig. 1B shows that the semicircle diameter in the Nyquist plot is very similar to the charge transfer resistance, Rct = 1.3 ± 0.2 kΩ and that the Ron = 35.1 ± 1.6 Ω is very low. The bare electrode also shows Warburg diffusion at high frequencies, lost after modification with the peptide (Inset Fig. 1B). In contrast, Rct increased three orders of magnitude (up to 25.0 ± 0.8 kΩ) after immobilizing the peptide at the electrode surface due to an increased interface thickness that insulated the conductive support and blocked the electron transfer from the electrochemical probe and the surface. This change also indicates the successful chemisorption of the thiolated peptide at the electrode surface. Treatment of the SPAuE with MCH induced a decrease in the insulation and Rct value (Fig. 1B) related to the reorganization of the amino acid chain of the peptide that went away from the transducer generating more free spaces for easier access of the electrochemical probe. We further evaluated the interaction of the peptide-modified electrode with synthetic Spike protein. When the peptide-electrode was incubated with a protein solution at 37 °C for 15 min, the protein binding event induced a decrease of the Rct to a value of 10.9 ± 0.6 kΩ due to re-orientation of the synthetic peptide end segment during the Spike protein recognition event [51]. P values in Table 1 showed a decrease in the capacitive behavior of the system, and in the meantime, the n values for modified electrodes were less than one, consistent with a double layer pseudo-interfacial capacitance at the electrode/electrolyte interface [59].

Although the use of the theoretical model of the Randles circuit for the adjustment of experimental data is widely extended; it is crucial to recognize that this model does not consider various factors involved in the biosensing process of the Spike protein, such as binding affinity, leaching, denaturation, transport phenomena, surface defects, among others; that may affect the results. Nevertheless, the very small values of Chi-squared function (χ2) for the best-fitted equivalent circuit, lower than 1.6 × 10⁻², demonstrated that the curve showed herein and the experimental data from the mathematical fitting agreed (See S.I Fig. S1 A-C). Table 1 summarizes all the fitted electrochemical parameters. These results suggest that the anchorage of the peptide and the interaction with the Spike protein decreased the capacitive current on modified-electrodes and decreased the charge transfer resistance, which indicates that the peptide-Spike protein recognition event was successful.
3.2. Optimization of the analytical conditions

Once the functionality of the biosensor platform was established, the peptide concentration was optimized as a function of the increase in the difference of the charge transfer resistance (ΔRct) (Figure 2). The behavior of the functionalized electrode was evaluated with different concentrations of the peptide after blocking with MCH and incubation with 1.0 mg/mL of a Spike protein solution. The ΔRct for 0.25 and 0.7 mM concentrations were lower (Figure 2A-C) than for 0.50 mM of peptide solution (Figure 2B). These differences may be associated with the organization degree of the peptide at the modified-electrode surface. Accordingly, too low concentrations of peptides seem to generate poor self-assembly of the monolayers, thus affecting the Spike protein recognition event. In contrast, too high concentration of peptide molecules could generate overlapping of peptide multilayers that limit the interaction of the specific segment of the peptide and the Spike protein. Figure 2D shows the comparative results where chemisorption of the bioreceptor at 0.50 mM produced the highest values of ΔRct. At this concentration, a suitable amount of peptide should be immobilized on the electrode surface and be accessible for interaction with the Spike protein in solution. Furthermore, a high peptide density can improve surface blocking and decrease nonspecific adsorption of proteins; therefore, such concentration was selected for the following experiments.

The peptide incubation time was evaluated in a range from 30 to 120 min and overnight, finding a higher ΔRct value for 60 min incubation time (See Figure S2A in SI), in correlation with the formation of a well-organized monolayer at the electrode surface. The concentration and blocking time with MCH were also systematically optimized after modifying the electrode-anchored peptide and comparing its interaction with the Spike protein. The difference in charge transfer resistance was greater for concentrations of 0.01 mM MCH after 10 min of blocking (See Figure S2 B–C in SI). Concentrations higher than 0.01 mM MCH may generate overreorganization of the monolayer and displacement of the peptide molecules that may move away from the transducer, affecting the Spike protein recognition event.

Finally, although the changes in ΔRct studied at different incubation times at 37 °C showed a maximum value at 30 min, 15 min produced comparable results and better than the longer time at 25 °C (See Figure S2D in SI). Therefore, the incubation temperature selected for the subsequent tests was 37 °C.

At lower temperatures, the RBD protein has a closed conformation that disables binding to the peptide due to its burying receptor-binding residues [59]. In summary, the optimized conditions are the incubation of the SPAuE in 0.5 mM of peptide for 60 min, blocking with 0.01 mM MCH for 10 min and interaction with Spike protein solutions for 15 min at 37 °C, indicating the easiness of the biosensor assembly of the format proposed and the

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### Table 1

| Electrode                     | Rct (kΩ) | Rs (Ω) | CPE | P x 10⁻⁵ | n   | $\chi^2$ x 10⁻⁵ |
|-------------------------------|----------|--------|-----|----------|-----|-----------------|
| SPAuE                         | 1.3 ± 0.2| 33.1 ± 1.6| 3.4 | 0.867⁵ | 1.6 | 0.3             |
| Peptide                       | 25.0 ± 0.8| 28.0 ± 2.8| 2.0 | 0.845⁵ | 1.3 | 1.0             |
| Peptide/MCH                   | 21.9 ± 0.1| 35.7 ± 0.9| 1.0 | 0.925⁵ | 1.0 | 0.7             |
| Peptide/MCH/Spike protein 0.1 μg/mL | 10.9 ± 0.6| 34.2 ± 1.5| 1.1 | 0.866⁵ | 1.0 | 0.7             |

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Fig. 2. Nyquist plots recorded using the redox probe 5 mM [Fe(CN)₆]³⁻/⁴⁻/KNO₃. Frequency from 50 kHz to 0.05 Hz and amplitude 10 mV. Peptide concentration: A) 0.25, B) 0.50 and C) 0.70 mM. D) The difference in the charge transfer resistance for the different concentrations of the synthetic peptide, with 0.01 mM MCH and interaction with 1.0 μg/mL Spike protein.
possibility of a reliable interrogation of the Spike protein concentration in only 15 min once the electrodes are sensitized.

3.3. Direct detection of spike protein using the impedimetric biosensor

Having the analytical parameters optimized, we evaluated the analytical performance of the peptide-based biosensor by analyzing different dilutions of the Spike protein ranging from 0.05 to 3.0 µg/mL as detailed in the Materials and Methods. Remarkably, charge transfer resistance (Rct) decreased (Fig. 3A) and the difference in charge transfer resistance (ΔRct) increased (Fig. 3B) with increasing concentrations of Spike protein solution due to the reorganization of the peptide structure during the Spike protein biorecognition event, as mentioned above.

The data in Fig. 3B show a trend line, with the inset showing the linear dependence described by the equation ΔRct = 10017 [Spike] + 9702 and a correlation coefficient R² = 0.9944. The peptide-based biosensor showed a high dependence on the concentration of Spike protein with a linear response of 0.05–1.0 µg/mL, a sensitivity of 10.2 kΩ mL/µg and a LOD of 18.2 ng/mL, whose LOD are higher than 1 fg/mL [60] and 2.9 ng/mL and 14.6 µA mL/ng [61], but comparable with 66 pg/mL and 14.9%/nM [62]. Remarkably, the signal of our peptide-based biosensor can detect Spike protein in solution and viral particles at clinically relevant concentrations as will be shown below.

In an attempt to approach a real scenario, we construct a calibration curve estimating the ΔRct for viral particles of SARS-CoV-2 inactivated by UV-light (Fig. 3C and D). The data showed a linear correlation between 10² to 10³ copies/mL with LOD of 0.01 copies/mL described by the equation ΔRct = 15308.5 log C – 27759.1 and a correlation coefficient R² = 0.9801 (Inset Fig. 3D). Considering the Spike protein levels [63,64] and viral particles on human samples [14], our results indicate that the peptide-based biosensor can detect Spike protein concentration-dependent after only 15 min sample incubation.

3.4. Specificity, storage stability, reproducibility, and repeatability studies

The specificity of the peptide-based biosensor is essential to reduce or avoid a false positive assignment. We considered herein 0.1 µg/mL of solutions that contain the Spike protein and compared it with those with RBD or β-1,4-GALT-5 glycosylated protein (Fig. 4A). The Spike protein is a transmembrane protein containing two subunits, S1 and S2. S1 mainly contains an RBD responsible for recognizing the ACE-2 host cell surface receptor, and the S2 region contains essential elements needed for membrane fusion once in the host cell [65]. The platform’s resultant ΔRct against the Spike protein and RBD, measured by EIS, exhibited a similar response with no statistically significant differences (p values are greater than 0.01). It means the RBD-peptide interaction of free RBD and RBD in the S1 region of the Spike protein are comparable. In contrast, the response was differential concerning β-1,4-GALT-5 and negative control with a very low β-1,4-GALT-5-peptide interaction, which could be explained by the presence of sialic acid in the β-1,4-GALT-5 [66] known as a receptor that binds to the S1 region of the Spike protein [65]. Subsequently, we estimated the specificity of the synthetic peptide-based biosensor against lysates from MERS and SARS-CoV, showing a high response when interrogated with 10⁴ copies/µL of lysed SARS-CoV-2 viral particles respect raw SARS-CoV-2, lysed MERS and SARS-CoV, with statistically significant differences (p < 0.01) (Fig. 4B). In contrast, raw SARS-CoV-2, SARS-CoV, and negative control showed a similar response with no statistically significant differences (p > 0.01), indicating the need for lysis to facilitate the peptide-Spike protein interaction. Yet, overall, the results demonstrate the high specificity of the peptide-sensitized transducer platform only for the Spike protein (in lysed solutions).

Long-term stability is also a significant feature in developing biosensors, especially when thinking about a translational approach. Next, the stability of the resulting peptide-based...
Five peptide-based biosensors were prepared independently and their impedimetric responses were measured, three times each, on the 3σ criteria (See Fig. S3A in SI). The stability was estimated by potting a control chart considering the first day of the study as the initial value. The upper and lower control limits were set at three times this value’s standard deviation (3σ). The finding confirms that the response of the peptide-based biosensors was in between the control limits and retained 73.6% of its initial response when tested up to 20 days of storage, which reveals the relative long-term stability of the biosensors related to the mixed self-assembled monolayer at the modified-electrodes surface.

Finally, we checked the reproducibility and repeatability of the developed biosensor by interrogating a 0.1 μg/mL Spike solution. Five peptide-based biosensors were prepared independently and their impedimetric responses were measured, three times each, under identical experimental conditions (See Fig. S3B in SI). The intra- and inter-electrode relative standard deviation (RSD) was estimated to be 4.1 (n = 5) and 2.2% (n = 3), which indicates the high repeatability and reproducibility of the biosensors developed, respectively.

The biosensor was challenged with nasopharyngeal swab samples from nine COVID-19 positive patient samples and three negative samples. The clinical diagnostic feasibility of the peptide-based biosensor was evaluated by analyzing the lysed diluted samples with different CT (cycle threshold) values, tested by RT-PCR, as reported in Tables 2 and SI. It is important to remark that RT-PCR is the gold standard technique for diagnosing COVID-19 based on detecting viral RNA from SARS-CoV-2 and used herein to estimate the samples’ viral loading, i.e., lower the CT higher the viral loading. Rather than RNA detection, our peptide-based device was designed to detect the Spike protein from viral swab particles (Table 2 in SI). Therefore, although results from RT-PCR can’t be directly compared with ours, they are clear evidence of a positive or negative sample. Fig. 5 shows that the biosensor can differentiate between positive and negative nasopharyngeal swab samples concerning the controls and differentiate the viral loading, i.e., high, middle, or low. It was evident that the biosensor is responding to the presence of Spike protein in COVID-19 positive samples with statistically significant differences (p < 0.01) concerning the negative samples and the negative control and no statistically significant differences between the negative samples and the negative control (p > 0.01). These results agree with those from RT-PCR, thus demonstrating the biosensor’s great potential for detecting the Spike protein from SARS-CoV-2 and its correlation with a rapid diagnosis of COVID-19.

3.5. Performance of peptide-based biosensor in clinical samples

The biosensor was challenged with nasopharyngeal swab samples from nine COVID-19 positive patient samples and three negative samples. The clinical diagnostic feasibility of the peptide-based biosensor was evaluated by analyzing the lysed diluted samples with different CT (cycle threshold) values, tested by RT-PCR, as reported in Tables 2 and SI. It is important to remark that RT-PCR is the gold standard technique for diagnosing COVID-19 based on detecting viral RNA from SARS-CoV-2 and used herein to estimate the samples’ viral loading, i.e., lower the CT higher the viral loading. Rather than RNA detection, our peptide-based device was designed to detect the Spike protein from viral swab particles (Table 2 in SI). Therefore, although results from RT-PCR can’t be directly compared with ours, they are clear evidence of a positive or negative sample. Fig. 5 shows that the biosensor can differentiate between positive and negative nasopharyngeal swab samples concerning the controls and differentiate the viral loading, i.e., high, middle, or low. It was evident that the biosensor is responding to the presence of Spike protein in COVID-19 positive samples with statistically significant differences (p < 0.01) concerning the negative samples and the negative control and no statistically significant differences between the negative samples and the negative control (p > 0.01). These results agree with those from RT-PCR, thus demonstrating the biosensor’s great potential for detecting the Spike protein from SARS-CoV-2 and its correlation with a rapid diagnosis of COVID-19.

4. Conclusion

We developed the first peptide-based impedimetric biosensor that detected and quantified Spike protein and viral particles of SARS-CoV-2 in human samples, straightforwardly in only 15 min. The sensing platform uses a synthetic peptide inspired by the interaction of the RBD region of the SARS-CoV-2 and the ACE-2 host receptor protein minimizing unspecific interactions with other proteins. The high performance of the label-free peptide-based biosensor detected a concentration of Spike protein and viral particles of SARS-CoV-2 of clinical relevance. The resultant device recognized Spike proteins in clinical swab samples and discriminated patients diagnosed with COVID-19 and healthy individuals. Overall, the peptide-based biosensor can sensitively and selectively detect the SARS-CoV-2 Spike protein and viral particles from clinical samples. Since the method can be performed with portable voltammetric analyzers, it can provide significant potential as point-of-care systems in the COVID-19 testing and be employed by mobile healthcare teams.
Declarations of competing interest

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: Dayana Soto reports financial support was provided by the University of Antioquia - Colombia Ministry of Science Technology and Innovation. Jahir Orozco reports financial support was provided by the University of Antioquia - Colombia Ministry of Science Technology and Innovation.

Acknowledgments

JO and DS thank the financial support provided by MINCICIANES, the University of Antioquia and the Max Planck Society through the Cooperation agreement 566-1, 2014; and EPM and Ruta N for hosting the Max Planck Tandem Groups. The authors acknowledge MINCICIANES for partial support from the project Nanobiosensors to detect SARS-CoV-2 rapidly (Cod. 1115101576765). The authors also thank the Tropical Diseases Study and Control Program (PECT) laboratory, the Immunovirology and the Gastrohepatology Groups from the University of Antioquia for donating the clinical samples and for letting us use the BSL-2 facilities, respectively.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.aca.2022.339739.

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