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Regenerable and high-throughput surface plasmon resonance assay for rapid screening of anti-SARS-CoV-2 antibody in serum samples

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HIGHLIGHTS
- Sequential use of a five-channel SPR instrument for high-throughput analyses.
- Development of a rapid screening assay of anti-SARS-CoV-2 antibody.
- Optimization of ligand density for reproducible and sensitive measurements.
- Cost-effective assays obviating the use of enzymes.

GRAPHICAL ABSTRACT

Current serological antibody tests for severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) require enzyme or fluorescent labels, and the titer well plates cannot be reused. By immobilizing histidine (His)-tagged SARS-CoV-2 spike (S1) protein onto tris-nitrilotriacetic acid (tris-NTA) sensor and using the early association phase for mass-transfer-controlled concentration determination, we developed a rapid and regenerable surface plasmon resonance (SPR) method for quantifying anti-SARS-CoV-2 antibody. On a five-channel SPR instrument and with optimized S1 protein immobilization density, each of the four analytical channels is sequentially used for multiple measurements, and all four channels can be simultaneously regenerated once they have reached a threshold value. Coupled with a programmable autosampler, each sensor can be regenerated at least 20 times, enabling uninterrupted assays of more than 800 serum samples. The accuracy and speed of our method compare well with those of the enzyme-linked immunosorbent assay (ELISA), and the detection limit (0.057 μg mL⁻¹) can easily meet the requirement for screening low antibody levels such as those in convalescent patients. In addition, our method exhibits excellent channel-to-channel (RSD = 1.9%) and sensor-to-sensor (RSD = 2.1%) reproducibility. Obviation of an enzyme label drastically reduced the assay cost, rendering our method (<60 cents) much more cost effective than those of commercial ELISA kits ($4.4–11.4). Therefore, our method offers a cost-effective and high-throughput alternative to the existing methods for serological measurements of anti-SARS-CoV-2 antibody levels, holding great promise for rapid screening of clinical samples without elaborate sample pretreatments and special reagents.

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1. Introduction

The novel coronavirus or severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has created the COVID-19 pandemic that is still ravaging the world [1–3]. As of the end of March 2022, over 480 million confirmed cases of COVID-19 have been reported with more than 6 million deaths worldwide [4]. SARS-CoV-2 is a large single-strand positive-sense RNA virus with a spike protein (S) that facilitates its entry into different cells by binding the angiotensin-converting enzyme 2 (ACE2) receptor at the cell membrane. The S protein comprises two subunits (S1 and S2), with the S1 subunit encompassing the receptor binding domain [5,8]. Compared to the severe acute respiratory syndrome (SARS-CoV) and Middle East respiratory syndrome coronaviruses (MERS-CoV), SARS-CoV-2 exhibits a longer viral incubation period and much higher transmissibility even before the onset of symptoms, the latter of which is responsible to the infection of nearly 6% of the total world population thus far [7,8].

In addition to developing effective therapeutic cures and vaccines, the availability of and accessibility to facile and high-throughput screening methods for SARS-CoV-2 is key to contain this pandemic. Many methods have been reported [5–18] and some have been widely implemented for mass testing. The methods that can be categorized into two types: molecular tests for viral RNA detection and serological tests for antibodies. Real-time reverse transcription-polymerase chain reaction (RT-PCR) is the most widely accepted method for diagnosing SARS-CoV-2 [11,12]. However, as viruses are present in the respiratory tract and RNA molecules are unstable, accurate diagnoses are highly dependent on careful nasal and/or oropharyngeal sampling and proper implementation of the RT-PCR, which are often difficult to carry out in developing countries [19]. The relatively long analysis time of RT-PCR also causes a delay in quarantine notification, which is critical to counter an outbreak in populated areas [20,21]. In addition to the high false negative/positive rates (up to 38% [22,23]), there are the high false negative/positive rates (up to 38% [22,23]), there are

\[ R = R_{\text{max}} k_a f C \]

where \( R \) is the SPR signal at a short time \( t \) of the association phase, \( R_{\text{max}} \) is the maximum binding signal for a sensor covered with a fixed amount of ligand, \( k_a \) is the association rate constant, and \( C \) is the analyte concentration. Although signals at an earlier stage of the association phase are smaller than those at the equilibrium phases, each analysis can be shorter than 1 min and requires a much smaller sample solution. Furthermore, the analytical signals can be augmented by using a relatively high density of ligand immobilized on the sensor so that \( R_{\text{max}} \) in the above equation is greater. Measuring \( R \) at a short \( t \) is certainly a more viable approach for the purpose of serological antibody assays of samples from a huge populace.

By immobilizing histidine (His)-tagged protein G onto a dextran-based tris–nitroliatriacetic acid (tris-NTA) sensor and using a dual-channel SPR instrument, we recently demonstrated the feasibility of using the early association phase for facile assays of serological assays of total immunoglobulin G (IgG) antibodies [49]. The faster speed and capability of surface regeneration represent a step closer than many other studies for rapid analyses of numerous clinical samples. To the best of our knowledge, SPR and other related detection variants have not been used for rapid and high-throughput mass testing of samples for infectious diseases. However, three additional challenges are inherent in the sensor assays for SARS-CoV-2 antibodies. First, the antibody concentrations of SARS-CoV-2 are in the low to mid-\( mg\) range (vide supra), more than three orders of magnitude lower than the total IgG-type antibodies in serum (5.6–177 mg mL\(^{-1}\) [50]). The second difficulty confronting the serological antibody testing for SARS-CoV-2 is sample throughput. If an SPR sensor needs to be regenerated after only one or a few measurements, the sample throughput is undoubtedly compromised. Finally, sample delivery, sensor regeneration, and data collection should be as automated as possible. This will allow unattended measurements to be performed, affording not only high throughputs, but also accurate and reproducible results. We resolved these problems by using four analysis channels sequentially over a single tris-NTA sensor immobilized with a S1 protein. Using this method, at least 20 cycles of surface generations can be achieved on a single sensor for assaying at least 800 serum samples (the exact number of assays is dependent on the antibody concentrations in sera). With 96 filter plates preloaded with samples and placed in a pre-programmed autosampler, continuous analyses and surface regeneration can be realized in an unattended manner for days. In addition to the small channel-to-channel variability (1.9%), the variability between different...
sensors is less than 3%. The average time for assaying each sample is 6 min (including times needed for sample delivery and measurements and sensor regeneration in between) and the cost for each sample testing is less than 60 cents. The excellent correlation between our method and a commercial ELISA kit in assaying donors inculoted with different vaccines indicates that our method is accurate and reproducible. Thus, our approach represents a more rapid and cost-effective alternative to the current ELISA and CLIA methods and stands to offer important information for studies of the durability of vaccination, immunity, and suitability of convalescent serum samples for therapeutic treatments of patients.

2. Experimental section

2.1. Reagents and materials

N-(3-Dimethylaminopropyl)-N’-ethylcarbodiimide hydrochloride (EDC), N-hydroxysuccinimide (NHS), and ethanalamine hydrochloride were acquired from Sigma (St. Louis, MO, USA). K2HPO4, KH2PO4, NaCl, NaOH, NiCl2, ethylenediaminetetraacetic acid (EDTA), Tween-20 and heparin were obtained from Macklin Biochemical Technology (Shanghai, China). His-tagged SARS-CoV-2 S1 protein and its polyclonal antibody (anti-SARS-CoV-2, 95% purity, type rabbit) were purchased from Sangon Biotech (Shanghai, China). CM-dextran sensor chips were obtained from Biosensing Instrument Inc. (Tempe, AZ, USA). Tris-NTA sensor was prepared as described previously [49]. Other reagents were of analytical purity and used as received. Deionized water was purified by a Millipore system (Simplicity 185, Millipore Corp, Billerica, MA).

2.2. Serum sample collection

The blood samples from two unvaccinated and 113 vaccinated donors were collected by the Hospital of The University of Jinan. These samples were centrifuged for 10 min at 2000 rpm after clotting, and the supernatants were individually collected as serum samples. The 113 immunized donors all received two doses of vaccines made by Sinopharm Group (China), Sinovac Biotech Co., Ltd. (Beijing, China), or Moderna Therapeutics (Cambridge, MA, USA). The current study was in accordance with the Helsinki declaration and approved by the Medical Ethics Committee of the Hospital affiliated with The University of Jinan.

2.3. Surface plasmon resonance

All experiments were conducted on a five-channel instrument (SPR-4500, Biosensing Instrument Inc.). A programmable autosampler (Spark 7800, A/Emmen Inc., Netherlands) that can accommodate two deep 96-well microplates (VWR International LLC, Radnor, PA) was used in conjunction with the SPR instrument. The running buffer was 10 mM PBS (10 mM KH2PO4/K2HPO4, 150 mM NaCl, 0.005% Tween-20, pH 7.4) containing 50 μM EDTA.

For assays of the anti-SARS-CoV-2 antibody in serum samples, the His-tagged S1 protein was immobilized onto each tris-NTA sensor to 1.000 ng mm⁻² by injecting 250 μL of 5 μg mL⁻¹ solution at 10 μL min⁻¹, using a procedure similar to that described in our previous work [51]. A series of anti-SARS-CoV-2 antibody standards with varying concentrations (0.5, 1.0, 2.0, 4.0, 8.0, 16.0, 32.0, and 96.0 μg mL⁻¹) were injected at 80 μL min⁻¹ for 60 s to construct the calibration curve. Serum samples were diluted 2- to 1000-fold, and 60-μL aliquots of diluted samples were injected at 80 μL min⁻¹. Each sample was analyzed at least three times, and 20 mM NaOH was used to dissociate preformed bioconjugates for sensor regeneration.

2.4. ELISA for antibody levels in serum samples

An ELISA kit for quantifying the anti-SARS-CoV-2 antibody was purchased from T & J Biomedical (Beijing, China) and used according to the manufacturer’s instruction. Serum samples (100 μL each), pipetted into wells of a 96-well titer plate, were incubated at 37 °C for 60 min. Afterwards, the content in each well was decanted and each well was washed three times with PBS. Horseradish peroxidase (HRP)-conjugated secondary antibody (50 μL) was then added into each well. Following incubation at 37 °C for 30 min, each well was again rinsed with PBS to remove the unbound secondary antibody. Then 100 μL of 3,3’,5,5’-tetramethylbenzidine (TMB) solution was added into each well and the resulting mixture was incubated at room temperature for 15 min. The UV–vis absorbance was recorded at 450 nm with a multimode microplate reader (TECAN, Spark, Mannedorf, Switzerland).

3. Results and discussion

3.1. SPR assay development

Fig. 1A illustrates the sequential use of four fluidic channels for detecting the anti-SARS-CoV2 antibody captured by the His-tagged S1 protein (scheme shown in Fig. 1B), with the first channel serving as the reference. The programmable autosampler is used in conjunction with a six-port injector valve (V1) and a check valve (V2) for delivering samples from a deep 96-well plate to the sensor surface. In a typical configuration, channel I (CH I), without immobilized S1 protein, serves as the reference for background subtraction, whereas CH II–CH V are pre-immobilized with S1 protein. When the center port of the selector valve (VC) is connected to port 3, samples are injected from CH I to CH II. CH II is continuously used until the SPR signal reaches a threshold wherein analytes can no longer be captured quantitatively (cf. the determination of the threshold described below). At this moment, the program connects the center port of VC to port 4, opening CH III for quantitative analyses of the next samples. This process continues until all four channels have reached or surpassed the threshold value. For antibody measurements, with the center port staying connected to port 6, a single injection of 20 mM NaOH simultaneously regenerates all four channels and the sensor can be used in the next assay cycle.

We first examined the effectiveness of sensor regeneration after the antibody capture. In Fig. 2A, the first three consecutive injections of 20 mM NaOH all completely dissociated the conjugates formed between the antibody and S1 protein, as reflected by the full recovery of the original baseline. We found that after 60 re-generations the sensorgram signal for the same antibody concentration varied by less than 5%. Consequently, highly reproducible sensorgrams can be repeatedly collected. The antibody binding signal increased with the S1 protein density between 0.500 and 1.000 ng mm⁻², but decreased beyond 1.000 ng mm⁻². This trend suggests that 1.000 ng mm⁻² is the optimal immobilization density that offers the greatest number of binding sites without imposing steric hindrance to the antibody binding (Fig. 2B). The sensorgram in Fig. 2C was collected from a continuous injection of a relatively high antibody concentration (20 μg mL⁻¹) for 1 h. Although the signal is quite high, a steady-state signal was not obtained, indicating that concentration determination using SPR signals at the equilibrium phase is indeed time- and sample-consuming and often impractical. Thus, our method, by measuring the SPR signals at the earlier stage of the association phase, shortens the analysis time and cuts down the sample consumption. Adopting this facile method is particularly advantageous for relatively weak biomolecular interactions whose equilibria are too long to attain [48,49].
Based on the two criteria for constructing calibration curves [49], we found that 25 s is the right time to obtain the following calibration curve:

\[ R = 4.726[C] + 0.022, \]

with a coefficient of determination \( R^2 \) of 0.9999 (Fig. S1). The relative standard deviations (RSDs) of signals in Fig. S1 range from 0.2% to 4.1%, indicative of high reproducibility. The detection limit of our method was estimated to be 0.057 \( \mu \text{g} \text{mL}^{-1} \) \( C_0 \) (S/N = 3), a value more than adequate for quantifying the anti-SARS-CoV-2 antibody in sera of convalescent patients [13,14,27].

### 3.2. Method establishment using antibody-spiked serum samples

As mentioned in the Introduction, a viable assay of the anti-SARS-CoV-2 antibody in serum must be rapid, reproducible, and of high throughput. While the feasibility of regenerating the sensor surface in a single channel is apparent in Fig. 2A, the throughput will be much greater if all four channels are highly comparable and can be regenerated simultaneously. Along this line, we immobilized CHs II–V with the same S1 protein density (1.000 \( \text{ng mm}^{-2} \)), with CH I serving as the reference channel for background subtraction. Fig. 3A illustrates the entire measurement sequence comprising 41 injections. To obtain these sensorgrams, the center port of \( V_C \) was first connected to port 3 (cf. Fig. 1), opening CH II for assays while preserving the other three channels for samples to be analyzed later. After ten consecutive injections of an 8.0- \( \mu \text{g} \text{mL}^{-1} \) antibody standard, the cumulative SPR signal reached 400 RU where many of the immobilized S1 protein molecules have captured the antibody (Fig. 3B). The next injection, above 400 RU, led to a binding signal that is more than 5% lower than the expected value (see also Table S1 in the Supplementary Information). Such a deviation indicates that the binding sites in CH II have become limited once the threshold value is reached. Instead of regenerating the channel right away, we programmed \( V_C \) (cf. Fig. 1) to connect its center port to port 4 so that CH III was open for accurate measurements of...
additional samples. Because CHs II–V are connected in series, the downstream channels display similar signals to the upstream one(s), with only those below the threshold value being accurate. Fig. 3C shows that five consecutive injections of 16.0-µg mL⁻¹ antibody into CH III resulted in a cumulative SPR signal of 400 RU, beyond which the accuracy of the sixth measurement deviates from the true value by more than 5% (cf. Table S1). Because the antibody concentration injected into CH III is doubled, the number of injections to reach this threshold value is halved when compared to that analyzed by CH II. Two points are worth mentioning: The amounts of antibody captured in the upstream channel hardly changes the antibody concentration in the bulk and does not affect the quantitative measurements in the downstream channels, a fact elucidated in our previous work [51]. The uniformity of the immobilized S1 protein in CH II is the same as that in CH III. When the same threshold in CHs IV and V is reached, the subsequent measurements also become less reliable (cf. Fig. 3D and E as well as Table S1). Thus, 33 injections (i.e., 10, 5, 10, and 8 measurements in CHs II, III, IV, and V) generated signals below the threshold. After signal in the last channel (CH V) reached 400 RU, 20 mM NaOH was injected to regenerate CHs II–V, reverting the signals of all four channels back to the original baseline.

The regeneration is indeed effective and not perturbative to the S1 protein conformation, as evidenced by Fig. 3B and E. In the former, the sensorgrams before (solid black curve) and after (dashed red) the regeneration are almost congruent, while in the latter, after analyses of samples of randomized antibody concentrations, six injections of 16.0-µg mL⁻¹ produced a sensorgram that is essentially identical to that recorded in CH III. We found that the channel-to-channel variability is only 1.2%. This is consistent with the small RSD values manifested in Fig. S1. The 33 accurate measurements displayed in Fig. 3A (see the measured values in Table S1) require only 185 min. Thus, each sample can be analyzed in 5.6 min, and the total time per sample is 6 min if the time for sensor regeneration is included, a time even shorter than that required for the qualitative LFA method (~15 min) [52,53].
Before we applied our method to the analysis of the anti-SARS-CoV-2 antibody in real samples, we conducted a recovery experiment to ensure that the matrix of the serum samples does not affect the assay accuracy. Table 1 shows the actual amounts of different antibody standards spiked into serum samples of uninfected and unvaccinated donors, along with values measured with our method. With a serum sample diluted 10-fold by the running buffer, the differences between the spiked and measured values are exceedingly small and the recovery values are satisfactory. Moreover, the anti-SARS-CoV-2 antibody measurement is not affected by the presence of heparin (cf. Fig. S2) [54,55], based on the exceedingly small variance (<1%). As heparin is commonly used in treating Covid-19 patients, this small variance ensures accurate quantification of antibody levels in serum samples of patients under treatments. Thus, the matrix effects are essentially negligible, due to the excellent anti-fouling properties of the dextran-based tris-NTA sensor surface [49].

We also found that our sensor can be continuously used for at least 20 assay cycles of serum samples, as evidenced by the high comparability (RSD% less than 1.9%) between the sensograms of the same samples measured in the same channel and that between the sensograms of samples measured in different channels subject to different regeneration steps (Fig. 4A). The four channels of this sensor were used uninterruptedly for a total of 877 assays in 87.6 h or 3.65 days. If the same approach is extended to SPR instruments possessing parallel channels [56] or related techniques designed for simultaneous analyses of different samples, the throughput will be even greater. For example, SPR imaging (SPRI) [57–59], an SPR variant, is capable of monitoring reactions occurring simultaneously in 96 channels [60]. We should also note that, in addition to the excellent channel-to-channel consistency, the variability between different sensors is also quite small. In Fig. 4B, the RSD% of the signals at 25 s of the three sensograms, collected separately from three sensors, is 2.1%.

![Fig. 4](image.png)

3.3. Assays of clinical serum samples

With the method established for serum sample analysis, we programed the instrument for automated analyses of the serum antibody levels from donors who have received two doses of different vaccines, as well as those from donors who were unvaccinated but not infected by Covid-19 (confirmed by the PCR tests of their oropharyngeal swab samples). The SPR assays of the 115 clinical samples were performed parallelly with ELISA in a double-blind fashion. The results displayed in Table S2 are in excellent agreement with each other. The Passing–Bablok regression plot comparing SPR and ELISA methods afforded a slope of 1.010 and an intercept of −0.467 μg mL⁻¹, whose 95% confidence interval (CI) encompasses values 1 and 0, respectively (Fig. 5). This plot indicates that proportional and constant differences are insignificant between SPR and ELISA [61]. Moreover, the Bland–Altman plot revealed a mean bias of 0.0 μg mL⁻¹, further indicating that the two methods are equivalent (Fig. S3A). Between these two methods, only four outliers were observed within ±1.96 SD (Fig. S3A).

![Fig. 5](image.png)

**Table 1**

| Samples   | Antibody spiked (µg mL⁻¹) | Antibody measured (µg mL⁻¹) | Recovery (%) |
|-----------|---------------------------|-----------------------------|--------------|
| Serum     | 0.20                      | 0.21                        | 105.0        |
|           | 1.00                      | 1.01                        | 101.0        |
|           | 5.00                      | 5.20                        | 104.0        |
|           | 10.00                     | 10.34                       | 103.4        |
|           | 40.00                     | 40.07                       | 100.2        |
|           | 45.00                     | 44.75                       | 99.4         |

Sample antibody recovery in serum samples.
consistent with the normal error (Gaussian) distribution for a large yet finite number of measurements, as shown by the Shapiro–Wilk test (Fig. S3B). As can be seen from Table S2, all of the vaccinated donors have antibody levels substantially greater than the two unvaccinated donors. Moreover, the test results are considered to be antibody-positive because their concentrations are all greater than the cut-off value, 1.0 μg ml$^{-1}$ [24–26,62].

We should note that the cost for our homemade tris-NTA sensor is about $200, which constitutes the bulk of the expense for screening. Thus, each measurement costs about 25 cents or less. Even if the commercially available sensor chip ($\approx$500) [63–65] were used and costs of other reagents were included, the overall cost would be still quite low (<60 cents/sample). An ELISA kit based on the platform of 96-well plates ranges from $420 to $1100 [66], and the cost for testing one sample corresponds to a range between $4.4 and $11.4. Therefore, our SPR method is much more cost-effective than ELISA. In addition, to minimize trial-to-trial and plate-to-plate variations, constructing a new calibration curve for a different 96-well plate is generally recommended [67]. All the procedures and requirements of ELISA decrease its throughput. Our method obviates the use of enzymes and the incubation/washing steps, and the programmable autosampler and regenerable sensor chips facilitate uninterrupted assays for days (cf. Fig. 4A).

4. Conclusions

Towards the objective of rapid, accurate, and high-throughput screening of the anti-SARS-CoV-2 antibody, we coupled a programmable autosampler with a five-channel SPR instrument in conjunction with a regenerable S1 protein sensor surface for continuous and automated analyses of many serum samples. Different from commercial ELISA that requires enzyme-linked antibodies, our method is label-free and uses signals of the early association phase of the biomolecular interaction to reduce analysis time and enhance the throughput. Consequently, hundreds of samples can be analyzed on a single, renewable sensor, and the high channel-to-channel and sensor-to-sensor comparability ensures data quality and assay fidelity. It is worth noting that SPR has been used for detections of viral particles such as H1N1, H7N9, and even SARS-CoV-2 [68,69]. Therefore, by immobilizing different proteins on different channels of a single sensor, serological testing can provide additional information about immunity, convalescence, and immunization efficacy. As a result, SPR is a versatile and powerful tool for studying the prevalence and pathogenicity of SARS-CoV-2 and other related infectious diseases.

CRediT authorship contribution statement

Meng Jiang: Formal analysis. Tianbao Dong: Investigation. Chaowei Han: Investigation. Luyao Liu: Investigation. Tianzang Zhang: Resources. Qing Kang: Supervision, Funding acquisition. Pengcheng Wang: Investigation, Supervision, Writing — original draft, Writing — review & editing. Funding acquisition. Feimeng Zhou: Conceptualization, Methodology, Supervision, Writing — original draft, Writing — review & editing, Funding acquisition.

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.

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Appendix A. Supplementary data

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