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Systematic Study of Various Functionalization Steps for Ultrasensitive Detection of SARS-CoV-2 with Direct Laser-Functionalized Au-LIG Electrochemical Sensors

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Abstract

The 2019 coronavirus (COVID-19) pandemic impaired global health, disrupted society, and slowed down the economy. Early detection of the infection using highly sensitive diagnostics is crucial in preventing the disease's spread. In this paper, we demonstrate electrochemical sensors based on laser induced graphene (LIG) functionalized directly with gold (Au) nanostructures for the detection of SARS-CoV-2 with an outstanding limit of detection (LOD) of ~1.2 $ag.mL^{-1}$. To optimize the sensors, we explored various functionalization parameters to elucidate their impact on LOD, sensitivity, and linearity. Specifically, we investigated the effect of (i) gold precursor concentration, (ii) crosslinker chemistry, (iii) crosslinker and antibody incubation conditions, and (iv) antigen-sensor interaction (diffusion-dominated incubation vs. pipette-mixing), as no systematic study on these parameters have been reported thus far. Our benchmarking analysis highlights the critical role of the antigen-sensor interaction and crosslinker chemistry. We showed that pipette-mixing enhances sensitivity and LOD by more than 1.6-fold and 5.5-fold, respectively, and also enables multimodal readout compared to diffusion-dominated incubation. Moreover, the PBA/Sulfo-NHS: EDC crosslinker improves the sensitivity and LOD compared to PBASE. The sensors demonstrate excellent selectivity against other viruses, including HCoV-229E, HCoV-OC43, HCoV-NL63, and influenza H5N1. Beyond the ability to detect antigen fragments, our

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Supporting Information. Comparison of surface modification, limit of detection, sensitivity, and selectivity among other graphenebased COVID-19 sensing platforms; electrochemical characterization of 1-pass LIG, 2-pass LIG, and L-Au/LIG; studies on effect of crosslinker and antibody incubation environment on antibody surface coverage density; studies on effect of mixing method for E-Au/LIG sensors; calibration curve values of sensors in response to antigen in buffer and artificial saliva, as well as virus mimics; cross-reactivity data for the optimized sensor's response to SARS-CoV S1 and MERS-CoV S1.

Graphical Abstract



Keywords

SARS-COV-2; biosensor; laser induced graphene; antigen; electrochemical

1. Introduction

The 2019 coronavirus (COVID-19) pandemic led to a drastic loss of human life and presented unprecedented challenges to global health, society, and the economy. As of 2024, at least 6.8 million people have died from COVID-19 globally.¹ At least 10% of infected individuals experience long COVID symptoms² including heart problems,³ blood clot-related issues,⁴ diabetes, chronic fatigue syndrome, and postural orthostatic tachycardia syndrome (POTS)—affecting 65 million people worldwide. Moreover, the pandemic pushed an estimated 90 million people into extreme poverty in 2020, the largest increase in global inequality and poverty since 1990.⁵ Increased testing availability enabled tracking of disease spread,⁶ but several diagnostic challenges remain.

Conventional polymerase chain reaction (PCR) tests, while accurate, are slow, expensive, and require trained personnel, causing strain on healthcare systems as they struggled to meet testing demands.⁷ Commercial rapid point-of-care (POC) antigen tests offer a faster alternative but concerns about their sensitivity and high rates of false negatives— especially for asymptomatic individuals with low viral load—remain a hurdle to disease containment and treatment access.⁸ Moreover, these rapid tests provide a qualitative binary outcome—positive or negative—without providing information on the viral load. To address these challenges, electrochemical antigen sensors are a viable solution.⁹ These sensors enable quantitative measurements of viral load to facilitate assessment of disease severity, monitoring of disease progression, and evaluation of treatment efficacy.

Sensors detecting COVID antigens have advantages over antibody tests due to their capability to identify earlier stages of illness when viral loads are typically higher, facilitating timely intervention and containment methods. Antibodies often only become detectable during the late and recovery stages of infection, which can lead to false negatives.¹⁰ In antigen sensors, antigens bind to biorecognition molecules (such as immobilized antibodies, aptamers, etc.) on the sensor surface, leading to a measurable change in the sensor signal, such as Faradaic current,¹¹⁻¹³ impedance,¹⁴⁻¹⁶ or surface potential/charge.¹⁷⁻¹⁹ An effective electrochemical sensor must have a large surface area for antigen immobilization and excellent electron transfer properties. Graphene supports both characteristics, making it a promising material for developing POC virus diagnostic devices.²⁰⁻²⁴ Various studies have capitalized on graphene's excellent electrical, physical, and mechanical properties for electrochemical antigen detection. Quasi-free-standing epitaxial graphene-based biosensors, developed by Kim et al., showed real-time ultrasensitive detection capability with a LOD of 1 ag. mL^{-1} for spike protein antigen.²⁵ Graphene ink printed on disposable paper substrates reported by Jaewjaroenwattana et al. offered a more cost-effective solution with high specificity and stability, achieving an LOD of 2.0 fg. mL^{-1} .²⁶

Recently, laser-induced graphene (LIG) has gained significant attention in electrochemical sensing. LIG is a scaffold-like porous material made using readily available films, such as polyimide, through a simple CO₂ laser processing, which carbonizes the substrate into graphene through photothermal and photochemical effects.²⁷ Several diagnostic devices based on LIG have been developed to detect COVID-19. Torrente-Rodríguez et al. demonstrated a LIG-based multiplexed telemedicine platform for viral antigen nucleocapsid protein, IgM and IgG antibodies, and the inflammatory biomarker C-reactive protein.²⁸ Cui et al. developed a field-effect transistor with a porous graphene channel to detect antigens at a concentration of 1 pg. mL^{-1} .²⁹ Other studies have explored LIG biosensors functionalized with gold nanostructures to improve LOD. Alafeef et al. developed an electrochemical biosensor chip using graphene/paper functionalized with gold nanoparticles (AuNPs).³⁰ The presence of AuNPs significantly improved the sensor's ability to detect the virus compared to without them, achieving a sensitivity of 231 copies. μL^{-1} and a LOD of 6.9 copies. μL^{-1} .³⁰ Sadique *et al.* harnessed electrodeposited AuNPs to improve the electrochemical properties of graphene oxide for the detection of SARS-CoV-2 antigen, achieving a limit of detection of 3.99 ag. mL^{-1} .³¹ The literature employs various graphene modification methods, including AuNP functionalization, choice of crosslinkers, and parameters such as humidity, temperature, and incubation duration as summarized in Table S1. However, a direct comparison of these functionalization parameters is missing.

In this study, we optimized the functionalization process of Au-LIG electrochemical sensors for SARS-CoV-2 antigen detection. We explored the functionalization of LIG with AuNPs via direct CO_2 laser processing (L-Au/LIG) as a means of improving sensor sensitivity, limit of detection (LOD), and selectivity. The study also investigates the impact of different functionalization parameters and compares two commonly used crosslinkers in graphenebased devices (PBASE vs. PBA/Sulfo-NHS: EDC) on the sensor's sensitivity. The effect of crosslinker and antibody incubation conditions is also studied. Moreover, we evaluated

the LOD and sensitivity of the sensors using two antigen-sensor interaction/incubation techniques (diffusion-dominated incubation vs. pipette-mixing), showing that the pipette-mixing not only improves the sensitivity and LOD, but also enables multimodal readout for more reliable testing. The sensor performance in artificial saliva is also demonstrated. Finally, as a more accurate model for the real infection, we tested the sensors with virion-like SARS-CoV-2 mimics and showed the ability to detect down to ~5 particles. mL^{-1} .

2. Results and Discussion

A. Characterization of material morphology, surface chemistry, and kinetic parameters

Fig. 1 depicts the fabrication steps of the Au-LIG SARS-CoV-2 antigen sensor and the overview of the parameters that are studied in order to optimize the sensor response (sensitivity, LOD, linearity, and noise). We first examine the direct laser functionalization of LIG with different concentrations of HAuCL₄ (gold precursor solution) in DI water on the sensor's electrochemical properties. We also evaluate the influence of two commonly used crosslinkers in graphene-based devices (PBASE vs. PBA/Sulfo-NHS: EDC) on the sensor's sensitivity. Moreover, we study the effect of crosslinker and antibody incubation conditions (humidity, temperature, duration) and antigen-sensor incubation process (diffusion-dominated incubation vs. pipette-mixing).

First, we investigate the effect of Au precursor concentration on the electrochemical properties of LIG functionalized with AuNPs via CO₂ laser processing. In this fabrication method, the CO₂ laser reduces Au³⁺ ions in the HAuCL₄ precursor solution, forming nanoparticles. The porous structure of LIG increases the surface area available for electrochemical reactions and protein immobilization, enhancing its effectiveness for sensing applications. To understand the effect of Au functionalization on the available surface area and kinetics of charge-transfer, we calculate electrochemically active surface area (ECSA) and the heterogeneous electron transfer rate constant, k^0 , as plotted in Fig. 2a and Fig. 2b for 1-pass LIG, 2-pass LIG, and L-Au/LIG functionalized with different concentrations of HAuCl₄ in DI water. ECSA and k^0 of the working electrode (WE), with a geometric area of π mm² are derived from cyclic voltammetry (CV) measurement data shown in Fig. S1a using the Randal-Sevcik equations (Eq. 1 and Eq. 2):

$$i_{peak} = 0.4463 n^{1.5} F^{1.5} CA \sqrt{\frac{Dv}{RT}},$$

Eq. 1

$$\psi = k_0 \sqrt{\frac{\pi DnFv}{RT}} = \frac{-0.6288 + 0.0021(\Delta E_{peak}n)}{1 - 0.017(\Delta E_{peak}n)},$$
Eq. 2

where i_{peak} is the peak current, *n* is the number of transferred electrons in the redox event (here n = 1), *F* is Faraday's constant, *D* is the diffusion coefficient of the redox probe (here

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 $D = 7.63e^{-6}\frac{cm^2}{s}$), *R* is the universal gas constant, *T* is the absolute temperature, *C* is the redox probe concentration, *A* is the electrochemically active surface area, and ΔE_{peak} is the oxidation-reduction peak separation in *mV*.

The direct laser functionalization process involves two passes of the laser over the WE, resulting in a double-layer LIG WE. We measured the electrochemical properties of the double-written WE LIG (2-pass LIG) without the AuNPs, aiming to determine whether the observed enhancement in ECSA and k^0 of L-Au/LIG sensors primarily resulted from the 2-pass LIG or the presence of AuNPs. Our findings indicate that the enhancement primarily originates from the presence of AuNPs rather than the double-written structure itself. Despite the addition of the second lasing, the ECSA and k^0 did not exhibit significant changes compared to the single lasing WE sensor. This suggests that improvements can be attributed to the AuNPs rather than the 2-pass WE structure.

ECSA increases by a factor of 3.3 with bare LIG compared to the geometric area. The enhancement ranges from a factor of 4.0 to 4.5 (compared to the geometric area) for L-Au/LIG functionalized with different concentrations of HAuCl₄ solution as shown in Table 1—the exception being 1:100 concentration of HAuCl₄: DI water, which reduces the ECSA and interestingly has the highest k^0 . L-Au/LIG shows a $1.3 \times$ to $1.9 \times$ enhancement of the rate transfer constant compared to 2-pass LIG. These results confirm that the direct laser functionalization of AuNPs significantly improves the electrochemical properties of LIG—in particular the available surface area for immobilization of the biorecognition molecules. A 1:100 concentration of HAuCl₄: DI water solution was chosen in the final design of the L-Au/LIG sensor given its most improved ECSA.

We also characterized the electrode surface morphology and chemical composition with scanning electron microscopy (SEM), Raman spectroscopy, and X-ray photoelectron spectroscopy (XPS). SEM reveals the porous 3D structure of LIG in Fig. 3a, a result of the rapid liberation of gaseous products during laser scribing.²⁷ The average Raman spectrum of 1-pass and 2-pass LIG samples in Fig. 3b depicts the D peak at ~1,350 cm^{-1} induced by defects or bent sp²-carbon bonds, the first order G peak at ~1,580 cm^{-1} , and the 2D peak at ~2,700 cm^{-1} originating from second order zone-boundary phonons.³² 1-pass LIG produces a more crystalline graphene with less structural disorder and defects compared to the 2-pass LIG, as confirmed by its higher 2D band intensity and lower D peak intensity.³³ The presence of Au 4f in the XPS spectrum in L-Au/LIG confirms the presence of Au on the graphene surface (Fig. 3c). In our previous work, X-ray diffraction (XRD) analysis of the L-Au/LIG material shows a slight peak of Au in the (111) plane, confirming the formation of crystalline gold structures. ³⁴

Sensor optimization: Studying the effect of various functionalization

parameters on sensitivity.—When more antigen molecules are immobilized on the surface of the electrode, the electron transfer kinetics change, resulting in a lower differential pulse voltammetry (DPV) current³⁵ and a higher charge-transfer resistance, R_{c} which is derived from the electrochemical impedance spectroscopy (EIS).³⁶ Thus, the DPV peak

current should decrease with increasing concentrations of antigen, and R_{ct} should increase with increasing the concentration of antigen. The data shown in Fig. 4a and 4b depict the DPV curves at each stage of the functionalization process using PBASE and PBA/Sulfo-NHS: EDC crosslinkers, respectively. Fig. 4c plots the DPV response of a representative sensor incubated with antigen concentrations ranging from 3 ag. mL^{-1} to 300 ag. mL^{-1} . As expected, the decreasing peak current after each functionalization step suggests that 1) both PBASE and PBA/Sulfo-NHS:EDC support activation of the carboxyl groups on the electrode surface; 2) the antibody cross-links to the carboxyl groups; 3) BSA blocks unbound sites on the electrode surface; and 4) the spike protein antigen is immobilized on the sensor surface. Fig. 4d and 4e depict R_{ct} after each functionalization step derived from the EIS Nyquist plots (an example shown in Fig. S1b). R_{ct} is extracted from the Nyquist plot by modeling the electrodesolution using an equivalent circuit shown in Fig. S1c. An increase in the R_{ct} after each stage of the functionalization process indicates surface modification at each step of the process, supporting the conclusions drawn from the DPV measurements.

To optimize the L-Au/LIG sensor, we first investigate how functionalization parameters -humidity, temperature, and duration-of the PBASE crosslinker and SARS-CoV-2 antibody affect the density of surface-linked antibodies. Maximizing the attachment of receptor antibodies enhances sensor sensitivity. Incubation duration is a crucial parameter to study because it affects the crosslinker and amine-terminated protein (i.e. antibody) binding efficiency,³⁷ thereby influencing sensor performance. In general, shorter incubation periods in bioconjugation might not provide sufficient binding time,³⁸ which could lead to weakened sensor sensitivity. Conversely, longer incubation times allow for more complete binding³⁸ but increase the risk of degradation (hydrolysis of crosslinkers or denaturing of antibodies³⁹), both of which may lead to weakened sensitivity. Incubation humidity and temperature also play critical roles in crosslinker and antibody binding. High humidity/low temperature prevents a 10 µL crosslinker/antibody aliquot from evaporating before it can diffuse through the working electrode and bind to graphene. Moreover, antibodies are more stable at lower temperatures.⁴⁰ From the DPV curves of L-Au/LIG modified with PBASE, SARS-CoV-2 antibody, and BSA (example shown in Fig. 4a and 4b), the amount of surface linked antibody can be evaluated using Eq. 3:³⁷

 $\Gamma = \frac{Q}{nFA},$

Eq. 3

Where *Q* is the exchanged charge in the redox event, *n* is the number of exchanged electrons (here n = 1), *F* is Faraday's constant, and *A* is the ECSA (here $A = 13.98 \text{ mm}^2$).

We first compare two common antibody incubation times from the literature: overnight (12 hours) and 3 hours (results summarized in Table S2). The PBASE crosslinker is incubated for an hour in ambient conditions, followed by incubation of the SARS-CoV-2 antibody for 3 hours/overnight and BSA for 1.5 hours at room temperature. Incubating the antibody for 3 hours compared to overnight increased the attachment density by 139% from $3.03 \pm 1.23 \times 10^{-10} \text{ mol.} mm^{-2}$ to $7.25 \pm 1.67 \times 10^{-10} \text{ mol.} mm^{-2}$. In the next series of tests, we compare

incubating PBASE for one hour under ambient conditions to a humidity chamber (97% humidity). There was no significant difference observed between them; however, we favored the 74% decrease in standard deviation of the humidity chamber. A similar experiment is performed to compare the incubation of SARS-CoV-2 antibodies at ambient room temperature versus refrigeration at 4°C. Incubating the antibody at 4°C increased attachment density by 1.55-fold. Finally, we examined various crosslinker incubation durations—1 hour, 1.5 hours, and 3 hours. Both 1 hour and 1.5 hours of incubation resulted in the highest antibody attachment densities. While there was no statistically significant difference between 1 hour and 1.5 hours of incubation, we favored the 2.85-fold lower standard deviation of 1.5 hours.

Using the optimized crosslinker/antibody incubation humidity/duration, we systematically studied the effect of crosslinker chemistry (PBASE vs. PBA/Sulfo-NHS) on antibody surface attachment density, given the lack of direct comparison in the literature. These crosslinkers are among the most studied functionalization methods in graphene-based sensors.^{18, 28, 41-46} In both crosslinkers, the aromatic pyrene group binds to the basal plane of graphene through pi-pi interactions.^{28, 45} In PBASE, the amine-reactive succinimide group covalently bonds to the antibody.⁴⁵ PBA does not inherently contain this succinimide ester group. Instead, EDC first activates the graphene's carboxyl group.⁴⁷ Sulfo-NHS then reacts with the activated carboxyl group to form a Sulfo-NHS ester group that enables amide bonding to antibodies.⁴⁷ These succinimide ester groups are prone to hydrolysis. which decreases their effective concentration and thus hinders the amidization process.⁴⁷ Because the humidity chamber did not have a statistically significant effect on antibody attachment density for the PBASE crosslinker but reduced noise, we continued using a humid environment for PBA/Sulfo-NHS:EDC. However, due to the sensitivity of the succinimide ester to hydrolysis, we optimized the incubation duration of the Sulfo-NHS: EDC. An incubation time of 2 hours yielded the highest antibody attachment density of 1.86 $\pm 0.22 \times 10^{-9}$ mol.mm⁻² (refer to Table S2), 78% higher than that of 1 hour—which may have not been enough time for complete bonding-and 142% higher than 3 hours-where hydrolysis evidently occurred.

We then evaluated the sensitivity and LOD of L-Au/LIG sensors functionalized with the optimized PBASE and PBA/Sulfo-NHS: EDC humidity/duration conditions showing a distinct advantage of PBA/Sulfo-NHS:EDC for graphene sensors. In Fig. 5a, present the R_{cr} (normalized w.r.t. R_{cr} of samples after BSA treatment) in response to varying concentrations of SARS-CoV-2 antigen (30—3 × 10³ ag.m L^{-1}). PBA/Sulfo-NHS: EDC yields 34% better sensitivity and 22% lower LOD than the sensors functionalized with PBASE (results are summarized in Table S3). The superior performance of the sensor functionalized with PBA/Sulfo-NHS stems from the higher antibody attachment density achieved by this crosslinker; this higher density likely results from its lower susceptibility to hydrolysis compared to PBASE. These findings underscore the critical impact of crosslinker selection and functionalization conditions on the sensitivity of electrochemical sensors.

Building upon the optimized functionalization parameters discussed above, we next investigate the effect of antigen-sensor interaction/incubation: diffusion-dominated

incubation vs. pipettemixing. We hypothesize that the pipette-mixing would expedite the transport of antigens to the working electrode surface where the antibodies immobilize them. Our results show that diffusion-dominated incubation yields 42% better sensitivity and 13% lower LOD than pipette-mixing for 1-pass LIG sensors (Fig. 5b). On the other hand, the pipette-mixing method demonstrates 1.6-fold improvement in sensitivity and 5.5-fold lower LOD for L-Au/LIG sensors (Fig. 5c). Importantly, pipette-mixing not only significantly improves sensitivity and LOD of L-Au/LIG sensors compared to diffusion-dominated incubation, but it also yields a more linear response with lower noise (refer to Table S3). Ultimately, L-Au/LIG sensors utilizing pipette-mixing yield the highest sensitivity

 $\left(56 \frac{\%}{\log(particles .mL^{-1})}\right)$ of all compared sensor conditions and an impressively low LOD of

~1.2 ag. mL^{-1} .

We believe diffusion-dominated incubation yields a better sensor response for 1-pass LIG and pipette-mixing yields a better response for L-Au/LIG because of the differences in how antibody-antigen immunocomplexes interact with their surface morphology. Antibodies bind to antigens through weak hydrogen bonds, electrostatic interactions and Van der Waals forces.⁴⁸ Excessive mechanical disruption—via pipette mixing—can physically dislodge antigens from the sensor's graphene surface, resulting in worsened sensitivity. On the other hand, graphene functionalized with AuNPs, which has a more complex topography and porosity, seems to benefit from mechanical mixing. L-Au/LIG has lower sensitivity than 1-pass LIG when relying on transporting antigens to the sensor surface with diffusion-dominated incubation alone, despite its larger ECSA and increase in available binding sites —perhaps due to its more complex topography and porosity. Pipette mixing may allow antigens to overcome the obstacles of the topography and better reach the available binding sites. Moreover, antibody-antigen immunocomplexes attached to this more complex surface may be more resilient to the mechanical agitation of pipette mixing.

It should be noted that we also initially investigated another Au-functionalization method, electroless deposition (E-Au/LIG). E-Au/LIG differs from L-Au/LIG in its method of AuNP formation. The LIG sensor is first submersed in a 1:100 ratio of HAuCl₄: ethanol, causing Au³⁺ ions to adsorb to the sensor's surface.⁴⁹ Adding ascorbic acid to the mixture reduces the Au³⁺ ions to Au nanoparticles.⁴⁹ Interestingly, while E-Au/LIG exhibits poor sensitivity and low yield (with 0 out of 4 sensors exhibiting good sensitivity) using the diffusion-dominated incubation (Fig. S2a), their performance was retrieved with 50% yield using pipette-mixing (Fig. S2b). These results underscore the importance of how antigens interact with sensors which is not usually clearly discussed in immunosensor literature. Moving forward, the results are collected using the optimum process, i.e. direct laser-functionalized AuNP LIG carboxylated with PBA/Sulfo-NHS: EDC and utilizing pipette-mixing during antigen incubation.

B. Testing in spiked saliva, selectivity analysis, interrogation with virion-like SARS-CoV-2 mimics

To demonstrate the suitability of the developed sensor for non-invasive detection of infection using saliva samples, we performed experiments using spiked artificial saliva. Human saliva

naturally contains electrolytes, which could potentially disrupt molecular sensing.⁵⁰ The artificial saliva mimicked the electrolyte concentration found in human saliva.⁵¹ Despite the presence of the electrolytes, the sensor maintained high sensitivity (S) and a strong linear correlation coefficient (\mathbb{R}^2), as evidenced by Fig. 6a.

To demonstrate the selectivity of the sensors toward SARS-CoV-2, R_{ci} response against non-target molecules is illustrated in Fig. 6b and S4. We examined the specific binding of the SARS-CoV-2 antigen in comparison to similar coronaviruses with varying degrees of sequence homology, such as Influenza A subtype H5N1 (n.d.), HCoV-229E (30%), HCoV-OC43 (33%), HCoV-NL63 (28%), SARS-CoV S1 (76-79%), and MERS-CoV (42-50%).^{52,53} Notably, there was no significant cross-reaction with HCoV-229E, HCoV-OC43, HCoV-NL63—which are responsible for a large proportion of seasonal common cold infections.⁵⁴ There is also no significant cross-reactivity with Influenza A subtype H5N1. However, we observed a measurable signal when testing with SARS-CoV S1 and MERS-CoV antigens, which is associated to a significant homology with SARS-CoV-2 spike protein (up to ~80% homology with SARS-CoV spike proteins and up to 50% homology with MERS-CoV). That said, such cross-reactivity is not a major cause for concern because there is not a current outbreak of these viruses.⁵⁵

In addition, to validate the effectiveness in detecting the target virus particles (i.e. virions), we performed electrochemical tests using negatively-charged virus mimics made of Spike antigen-coated polymer beads with diameter of 100 *nm* (similar in size and charge to SARS-CoV-2 virus – See Zeta potential data in Fig. S3a). These mimics closely emulate the virus structure and functionality, offering a more accurate representation of the sensor's performance compared to just the antigen fragments. As shown in Fig. 6c and 6d and summarized in Table S4, ultralow counts of the virus down to ~5 particles/mL can be detected using the 2-pass LIG. On the other hand, the L-Au/LIG sensor has poor linearity (r^2 =0.26), and thus its sensitivity and LOD cannot be quantitatively determined. To elucidate why the response of the 2-pass LIG sensor to virus mimics is better than the L-Au/LIG sensor, we measured the Zeta potential of the two materials. Fig. S3b illustrates the more negatively charged surface of the L-Au/LIG compared to LIG is believed to be due to electrostatic repulsion between negatively charged virion mimics and the Au-LIG surface.

Overall, as summarized in Table S5, the obtained LOD of the optimized Au-LIG sensor is superior compared to previously reported sensors for detecting antigen fragments. Moreover, compared to the existing reports which mostly just focus on antigen fragments, our work takes a step further by testing the sensors using virion-like particle mimics which better represent the real infection.

3. Conclusion

In this study, we demonstrate electrochemical sensors based on LIG functionalized with Au nanostructures using direct laser processing for the rapid and selective detection of SARS-CoV-2. By systematically exploring various functionalization parameters—such as gold precursor concentration, crosslinker chemistry, and incubation and antigen

mixing conditions—we optimized the sensitivity and LOD of our sensors and obtained 56 $\frac{\%}{\log(ag.mL^{-1})}$ and ~1.2 $ag.mL^{-1}$, respectively. We found that a HAuCl₄: DI water

precursor concentration of 1:100 enhances the ECSA by 4.5-fold compared to the geometric area. Additionally, we found that PBA/Sulfo-NHS: EDC improves sensitivity and LOD by 34% and 22% compared to PBASE. Moreover, using pipette-mixing for antigen incubation (compared to diffusion-dominated mixing), we achieved sensitivity enhancement of 1.6-fold, LOD enhancement of 5.5-fold, and multimodal EIS and DPV readout. To demonstrate the ability to analyze real samples, we tested the sensors with artificial saliva as well as virion-like SARS-CoV-2 mimics (polymer beads coated with SARS-CoV-2 Spike protein and with a similar size and charge as the SARS-CoV-2 virions), which more accurately model the real infection compared to antigen fragments. This study offers valuable guidelines for researchers to optimize the fabrication of various graphene-based sensors (beyond just electrochemical devices) that utilize the immobilization of capture molecules (antibodies, DNA, fragments, aptamers, etc.) for a wide range of biological analytes (antigens, proteins, nucleic acids, neurotransmitters, etc.)

4. Experimental details

A. Materials and Reagents

Polyimide (PI) sheets were purchased from American Durafilm Co., Inc. (Kapton HN, 500 mils). Gold (III) chloride solution (CAS: 16903-35-8), sodium chloride (CAS: 7647-14-5), L-ascorbic acid (CAS: 50-81-7), calcium chloride (CAS: 10043-52-4), potassium chloride (CAS: 7447-40-7), citric acid (CAS: 77-92-9), potassium thiocyanate (CAS: 333-20-0), ammonium chloride (CAS: 12125-02-9), 1-Pyrenebutyric acid (CAS: 3443-45-6), Sulfo-NHS (N-Hydroxysulfosuccinimide sodium salt) (CAS: 106627-54-7), Bovine Serum Albumin (CAS: 9048-46-8), Tween 20 (CAS: 9005-64-5), N,N-Dimethylformamide (DMF) (CAS: 66-12-2), MES (CAS: 145224-94-8), Glycine (CAS: 56-40-6), and Sodium Azide (CAS: 26628-22-8) were purchased from Sigma Aldrich. EDC (1-ethyl-3-(3dimethylaminopropyl) carbodiimide hydrochloride) (CAS: 22980) was purchased from Thermofisher. PBASE (1-Pyrenebutanoic Acid Succinimidyl Ester) was purchased from Chem Cruz. 2-propanol (IPA) was purchased from J.T. Baker (CAS: 67-63-0). DPBS (Dulbecco's Phosphate buffered saline) (CAS: 20-030-CV) was purchased from Corning. Ethanol (Koptec, 200 proof) was purchased from Decon Labs (CAS: V1016). SARS-CoV-2 (2019 nCoV) Spike S2 Antibody Affinity Purified (CAS: 40590-T62), Influenza A H5N1 (A/Hubei/1/2010) Neuraminidase / NA (His Tag) (CAS: 40018-V07H), Human coronavirus (HCoV-229E) Spike/S1 Protein (S1 Subunit, His Tag) (CAS: 40601-V0BH), Human coronavirus (HCoV-OC43) Spike S1 Protein (His Tag) (CAS: 40607-V08H1), Human coronavirus (HCoV-NL63) Spike/S1 Protein (S1 Subunit, HIS Tag) (CAS: 40600-V08H), SARS-CoV Spike/S1 Protein (S1 Subunit, His Tag) (CAS: 40150-V08B1), MERS-CoV Spike/S1 Protein (S1 Subunit, aa 1-725, His Tag) (CAS: 40069-V0BH) were purchased from Sino Biological. SARS-CoV-2 S Protein HIS Tag (CAS: 103871-150) was purchased from VWR International, LLC. CML Latex Beads, 4% w/v, 0.1 um (CAS: C37479) were purchased from ThermoFisher Scientific (CAS: F8803). Ecoflex 5, Smooth-On, Inc.

was purchased from Amazon. Silver conductive epoxy adhesive was purchased from MG Chemicals (CAS:8331D-14G).

B. Sensor preparation and functionalization of laser induced graphene (LIG)

We designed a three-electrode graphene sensor pattern with a working electrode (WE; L-Au/ LIG), a counter electrode (CE; LIG), and a reference electrode (RE; Ag/AgCl paste on LIG) in AutoCAD. A 30-W CO₂ laser engraving machine (VSL2.30, Universal Laser Systems) converts PI sheets to LIG. To prepare the polyimide sheet for laser writing, we rinsed it with 2-propanol (IPA), dried it with Kimwipes, and then fixed it to a glass substrate using double-sided adhesive tape. The laser ablates the PI sheets using the following fabrication parameters: 12.6% power, 5.5% speed, 1000 points per inch (PPI) resolution, in raster mode at a focused height of 0.040. For the fabrication of L-Au/LIG, different concentrations (1:1000, 5:1000, 1:100, 5:100) of HAuCl₄ solution in DI water were prepared. After patterning the WE with the CO₂ laser, we drop-cast a 5 µL aliquot of HAuCl₄: DI water mixture onto the WE and dried with a fan for 10 minutes. Gold nanoparticles form as an additional laser scribing step prints a second layer of the WE over the first layer. We also patterned the CE and RE in this step. For the fabrication of E-Au/LIG, we submersed the three electrode LIG sensor in a 1:100 ratio of HAuCl₄ in ethanol for 30 minutes. After adding ascorbic acid to the mixture to bring the concentration to 1 mM, we left the LIG sensor for an additional hour. Finally, we applied silicone (Ecoflex 5, Smooth-On, Inc.) between the sensor region and the electrical contact pads to passivate the sensor.

C. Characterization of the functionalized LIG electrodes

Scanning electron microscopy (SEM) micrographs were taken using a ThermoFisher Q250 instrument with a $6,500 \times$ magnification. X-ray photoelectron spectroscopy (XPS) measurements were carried out using a Physical Electronics VersaProbe II instrument (Chanhassen, MN). An Al Ka X-ray source was used at 45° takeoff angle. Charge neutralization was achieved using low energy (<5 eV) electrons and Ar ions. Raman spectroscopy was carried out using a Horiba LabRam instrument (Kyoto, Japan) with a 50× objective and 300 gr/mm grating. A 532 nm laser operating at 25% of 110mW was used. Raman spectra were analyzed using the LabSpec 6 software and XPS spectra were analyzed using CasaXPS.

D. Zeta potential of LIG electrodes

To determine the surface charge of the LIG and L-Au/LIG, graphene powder was prepared by scraping 2-pass LIG or L-Au/LIG from the polyimide surface, as previously described,⁵⁶ and then suspended in DIW. 1 mL of the graphene powder suspension was injected into a capillary cell and its zeta potential measured using a Zetasizer Nano ZS apparatus (Malvern Instruments Ltd.) with an equilibrium time of 2 minutes at room temperature.

E. Electrode functionalization with PBASE crosslinker

To activate carboxylic groups on the electrode surface,³² we drop-cast 10 μ L of 2.0 mM PBASE in methanol on the WE. After incubating the sensor for 1.5 hours in a humid chamber (97% humidity), we rinsed it with phosphate-buffered saline (PBS) to remove

the unbound reagent. Next, we prepared a 1:2000 ratio of SARS-CoV-2 (2019 nCoV) Spike S2 Antibody to 0.1% Tween 20 detergent dissolved in PBS. We drop-cast 10 μ L of this antibody solution onto the WE, and incubated the sensor for 3 hours at 4 °C. After incubation, we rinsed the sensors with PBS to remove excess and unbound proteins. We drop-cast 10 μ L of 0.1 % BSA and 0.1% Tween 20 detergent in PBS on the WE and incubated the sensor for 1.5 hours in ambient room temperature to block unbound free sites on the surface of the electrode. We again rinsed the electrodes with PBS. Finally, the sensors were ready for limit of detection and sensitivity studies.

F. Electrode functionalization with PBA and Sulfo-NHS: EDC

To activate carboxylic groups on the electrode surface,⁵⁷ we drop-cast 10 μ L of 5.0 mM PBA in DMF on the graphene surface and incubated for 2 hours at room temperature in a humid chamber (97% humidity). After rinsing with DMF, IPA, and deionized (DI) water and then drying under nitrogen airflow, we drop-cast a 10 μ L aliquot of a solution containing 0.4 M EDC and 0.1 M sulfo-NHS in 0.025 M MES (pH 6.5) on the WE. We incubated the sensor in a humid chamber for 35 minutes. The sensors were then functionalized with antibodies and BSA as described in the preceding section.

G. Preparation of artificial saliva

The artificial saliva was composed of 5 mM NaCl, 1 mM of $CaCl_2$, 15 mM of KCL, 1 mM of citric acid, 1.1 mM of KSCN, and 4 mM of NH₄Cl in DI water. HCL and NaOH were used to bring the pH to 6.7 (the pH of human saliva).⁵⁸

H. Preparation of SARS-CoV-2 antigen-coated polymer mimics

We prepared virus mimics by first dissolving 10 µL of SARS-CoV-2 S protein with 600 μ L. mL^{-1} concentration in 720 μ L of 50 mM MES buffer (pH 6.0) to make a 0.1 mg. mL^{-1} protein solution. We then added 270 µL of a 2% aqueous suspension of 100 nm carboxylate-modified microspheres to the protein solution and then left to incubate for 15 minutes. After incubation, we added 10 mg of EDAC into the mixture and then mixed initially via vortex and then for 2 h on a rocker orbital shaker. To mitigate agglomeration of the microsphere particles, we adjusted the pH to 6.5. At this point, we added glycine to achieve a concentration of $100 \, mM$, which quenches the reaction. We then incubated the mixture for 30 minutes. Next, we centrifuged the protein-labeled microsphere particles from unreacted protein at $25,000 \times g$ for 30-60 minutes. After resuspending the pellet in 1 mL of 50 mM PBS by gentle vortex, we centrifuged again at $25,000 \times g$ for 30-60 minutes. This washing step was repeated twice more. After washing, the protein-conjugated microspheres were resuspended in 100 µL of 50 mM PBS with 1% BSA to form a stable suspension. Finally, we added sodium azide — a preservative⁵⁹ — to achieve a concentration of 2 mMand stored the protein-conjugated microspheres at 4°C until use. The successful construction of the SARS-CoV-2 mimics were confirmed using Flow Cytometry (for details, see⁶⁰).

I. Electrochemical testing

To prepare for electrochemical measurements with the multi-channel potentiostat (MultiPalmSens4, PalmSens), we painted Ag/AgCl paste on the RE. We electrochemically

characterized the sensors using cyclic voltammetry (CV) with an applied window of -1 V to 1 V and a scan rate of 30 mV / s; differential pulse voltammetry (DPV) measurements applied a potential window of -0.5 V to 1 V and a scan rate of 30 mV / s; and electrochemical impedance spectroscopy (EIS) measurements applied a frequency range of 10 to 10,000 Hz. We used 5 mM ferricyanide-ferrocyanide in DI water (12.9 M Ω . cm) as the redox probe.

The sensors were tested using varying concentrations of SARS-CoV-2 spike protein antigen or virion-like mimics via serial dilution. We exposed the sensors to the lowest concentration of the antigen first. 100 μ *L* of the lowest concentration antigen solution was drop-casted on the working electrode of each sensor and was either left to incubate statically for 15 minutes (i.e. diffusion-dominated incubation) or was pipette-mixed for 30 *s* every 5 minutes for a total of 15 minutes to transport the antigens to the WE surface. After 15 minutes, the sensor was rinsed with phosphate-buffered saline (PBS) to remove unbound antigens. Next, DPV and EIS tests were performed. Each test was scanned three times for replicability, as the current response/impedance response can vary between scans. After these electrochemical tests, the sensors were rinsed with PBS to remove electrolyte residue. Then, the next highest concentration of antigen was tested. Because the previous concentrations of antigen that the sensor was exposed is a minimum of one order of magnitude smaller than the concentration being tested, they do not interfere with the electrochemical tests.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1. Fabrication of L-Au/LIG.

First, a CO₂ laser carbonizes polyimide into laser induced graphene (LIG), patterning the working electrode (WE). Au is directly doped into LIG through a second lasing step to form L-Au/LIG. Effect of different Au concentrations is studied. Carboxylic groups are activated on the WE surface with PBASE or PBA/Sulfo-NHS: EDC as crosslinker. Effect of incubation humidity is studied. SARS-CoV Spike S2 antibody covalently bonds to the carboxylic groups. The effect of incubation temperature for this step is studied. Bovine serum albumin (BSA) is then used to block unbound sites, followed by testing with the virus samples. Specifically, two different antigen mixing techniques are investigated in this study.





(a) Electrochemically active surface area (ECSA) of LIG and L-Au/LIG functionalized with different concentrations of HAuCl₄ (the gold precursor solution). Data are presented as mean (n=9). Error bars are standard error of the mean. (b) k^0 of LIG and L-Au/LIG functionalized with different concentrations of HAuCl₄. Data are presented as mean (n=9). Error bars are S.E.M.





(a) SEM image of the LIG confirms the 3D porous structure of the material (with $6,500 \times$ magnification). Scale bar is 20 µm. (b) Average Raman spectra of the 1-pass (n=24) and 2-pass LIG sensor (n=8). Error bands are standard deviation (STD). (c) XPS results of 1-pass LIG (n=4), 2-pass LIG (n=4), and L-Au/LIG sensors (n=4) showing their chemical components. Error bands are STD.



Figure 4. L-Au/LIG electrochemical sensor performance.

(a) Differential pulse voltammetry (DPV) curves after each step of functionalization process with PBASE crosslinker. Antigen concentration is 30 ng. mL^{-1} . (b) Representative differential pulse voltammetry (DPV) curves after each step of functionalization process with PBA/Sulfo-NHS: EDC crosslinker. Antigen concentration is 30 ng. mL^{-1} . (c) Representative DPV curves in response to varying concentrations of SARS-CoV-2 antigen. (d) The charge-transfer resistance (R_{ct}) after each step of functionalization process for PBASE crosslinker. Antigen concentration is 30 ng. mL^{-1} . Data are presented as mean with n = 3. Error bars are standard error of the mean. (e) The charge-transfer resistance (R_{ct}) after each step of functionalization process for PBA/Sulfo-NHS: EDC crosslinker. Antigen concentration is 30 ng. mL^{-1} . Data are presented as mean with n = 3. Error bars are standard error of the mean.



Figure 5. Studying the effect of crosslinker chemistry and antigen-sensor interaction (mixing method).

(a) Charge transfer resistance (R_{cr}) of L-Au/LIG sensors (normalized w.r.t. R_{cr} of samples after BSA treatment) in response to various concentrations of SARS-CoV-2 antigen in PBS using PBASE vs. PBA/Sulfo-NHS: EDC crosslinkers. Sensors are incubated with antigen using the diffusion method. Data are presented as mean (n=12). Error bars are standard error of the mean (S.E.M.). (b) Normalized R_{cr} of 1-pass LIG and 2-pass LIG sensors (with PBA linker chemistry) in response to various concentrations of SARS-CoV-2 antigen in PBS using diffusion-dominated incubation vs. pipette-mixing methods. Data are presented as mean (n=12). Error bars are S.E.M. (c) Normalized R_{cr} of L-Au/LIG sensors in response to various concentrations of SARS-CoV-2 antigen in PBS using diffusion-dominated incubation vs. Data are presented as mean (n=12). Error bars are S.E.M. (c) Normalized R_{cr} of L-Au/LIG sensors in response to various concentrations of SARS-CoV-2 antigen in PBS using diffusion-dominated incubation vs. Data are presented as mean (n=12). Error bars are S.E.M. (c) Normalized R_{cr} of L-Au/LIG sensors in response to various concentrations of SARS-CoV-2 antigen in PBS using diffusion-dominated incubation vs. Pipette-mixing methods. Data are presented as mean (n=12). Error bars are S.E.M.



Figure 6. Investigation of the sensor performance in saliva and with virion-like mimics. (a) Charge transfer resistance (R_{cl}) derived from EIS data (normalized w.r.t. R_{cl} of samples after BSA treatment) after incubation with various concentrations of SARS-CoV-2 antigen in artificial saliva (3—300 ag. mL^{-1}). Data are presented as mean (n=9). Error bars are standard error of the mean (S.E.M.). (b) Selective response of L-Au/LIG sensors against 30 ag. mL^{-1} concentration of SARS-CoV-2 and non-target antigens. Data are presented as mean (n=9). Error bars are S.E.M. (c) R_{cl} after incubation with various concentrations of SARS-CoV-2 virus mimics in PBS ($2 \times 10^2 - 2 \times 10^4 \text{ particles} \cdot mL^{-1}$) using 2-pass WE LIG sensors. Data are presented as mean (n=12). Error bars are S.E.M. (d) R_{cl} after incubation with various concentrations of SARS-CoV-2 virus mimics of SARS-CoV-2 virus mimics are presented as mean (n=12). Error bars are S.E.M. (d) R_{cl} after incubation with various concentrations of SARS-CoV-2 virus mimics in PBS ($2 \times 10^2 - 2 \times 10^4 \text{ particles} \cdot mL^{-1}$) using L-Au/LIG sensors. Data are presented as mean (n=12). Error bars are S.E.M.

Table 1.

Calculated electrochemically active surface area (ECSA) and k^0 .

The calculated ECSA and the heterogeneous electron transfer rate constant (k^0) and comparison between 1-pass LIG, 2-pass LIG, and L-Au/LIG. Data are presented as mean with n=9.

1-pass LIG	10.38	0.208
2-pass LIG	9.283	0.224
L-Au/LIG 1:1000	8.48	0.386
L-Au/LIG 5:1000	12.52	0.359
L-Au/LIG 1:100	13.98	0.309
L-Au/LIG 5:100	12.64	0.269