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# Rapid Detection of COVID 19 via Surface Enhanced Raman Spectroscopy

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**Abstract:** Coronavirus Disease-2019, known as COVID-19, has been declared a public health emergency of worldwide concern [1-4]. This paper reports a new design of Surface Enhance Raman Spectroscopy (SERS) substrate that can deliver higher sensitivity and the closest interaction to the target structure of the hot spots for the immunomagnetic capturing based concentration method. This substrate contains Polydimethylsiloxane (PDMS) inverted nanocavity arrays, which can funnel the light to the top where plasmonic gold nanorods occur. A high reflective and low loss layer is deposited at the outer surface, which reflects the photons multiple times and hence the number of photons is increased. After monitoring the consistent pattern and geometric shape of inverted cavities, the hot spots of the suggested Surface Enhance Raman Spectroscopy (SERS) substrate might have improved influence of  $10^{10}$  or even higher, which could be sufficient to discover the immunomagnetically densified virus.

Raman spectroscopy is a promising candidate in virus diagnosis given its fast and label-free nature, so long as the concentration of the pathogen is high enough to provide reasonable sensitivity [5-9].

**Keyword:** COVID 19, Plasmonic hot carrier collection, Gold nanorods, SERS, nanocavity arrays, PDMS.

## I. INTRODUCTION

Surface Enhanced Raman Spectroscopy (SERS) technique is an operational biosensors technique that could make the sensitivity of the pathogen more rapid [5-9]. Scattering photons deliver chemical information in the form of vibrational rays to other chemical bonds with the highest spatial resolution [10]. However, as the inelastic scattering signals are very weak, Raman scattering cannot be ideal for the identification of chemicals unless these signals are enhanced effectively. The vital idea in creating SERS is to generate the ideal metal surface. In addition to this, it has also been discovered that SERS mainly depends on the kind of metal and the roughness of the surface of that metal, and consequently, the major concern of this process is to prepare the substrate [10].

The main purpose of imposing SERS into the minimum concentration of bio-sensing is to enhance the intensity level of the vibration mode,  $I_{SERS}$  through increasing the electric field at the location of the sensing [[10]].

$$I_{SERS} \propto \left| \frac{E(\lambda)}{E_0(\lambda)} \right|^2 \left| \frac{E(\lambda_s)}{E_0(\lambda_s)} \right|^2 \quad (1)$$

Where  $\lambda$  is the incident wavelength and  $\lambda_s$  represents the Raman scattering wavelength,  $E$  signifies the local electric field intensity;  $E_0$  is incident electric field intensity. As the frequency shift due to Raman scattering is smaller than the incident wavelength, the enhancement factor (EF) is shown as [[11]]:

$$E \propto \left| \frac{E(\lambda)}{E_0(\lambda)} \right|^4 \quad (2)$$

With the provided order of magnitude enhancement on the light field, the signal output of a substrate is:

$$I \propto EF \times N_{molecule} \quad (3)$$

Where,  $N_{molecule}$  characterizes the number of molecules in relation to the plasmonic "hot-spot".

The most common method to enhance the EF factor of a substrate is to use plasmonic nanomaterial that can provide the localized surface plasmon attribute (LSPR) [7]. In LSPR, the surface plasmon will connect with the photon, which is too close to the surface of the metal, restraining the light field within a small range and increasing light field intensity.

Within the typical detection, the pathogen concentration is as small as 1-100 CFU/mL, and the pathogen must be enhanced by  $10^5$  to accomplish Raman measurements [[12], [13]]. So according to this situation, the performance of the Raman enhancement approach is to put control on the local electromagnetic environment of plasmonic deposition, along with the intention of increasing the concentration of photons through which the metal deposition can take.

Weisheng et al. [[15]] suggest placing the gold dimers on a metallic surface. The metal surface creates the plasmon and restrains the light field. The dimers receive the photons from the focused light field and the EF is about  $10^5$ . Masahiko et al. [[15]] use Au disc array on a dielectric-metal-quartz substrate. The periodic disc array scatters lights into wave pack whose wavevector matches the plasmon mode between dielectric and metal, and photons are trapped and delivered to the discs. In this work, the EF rises to  $10^6$ . Dongzhi et al. [[16]] have said to place the metallic film on oxidized porous Al. The  $Al_2O_3$  layer between the substrate and metallic film will perform as the function of the resonator that records the light, which enhances the EF created by the metal film at the topmost structure, resulting in an EF of  $10^4$ . Still, the enhancement of a single nanorod can reach  $10^6$  if associated accurately with incident light.

Immunomagnetic capturing is one of the famous methods to concentrate pathogens, which permits two to three orders of magnitude upgrading of the pathogen. Moreover, the pathogen will be connected by the antibiotic modified magnetic bead. The magnetic field can be utilized to collect the pathogen-bead clusters [18-20]. Further, for the purpose of performing rapidly the non-culturing discovery of the pathogen, Raman sensor sensitivity should be upgraded in a more effective way. But the purpose of Raman measurements cannot be considered to fulfill this sensitivity gap. The trapped pathogen will be transmitted to the Raman substrate as in a substrate approach [[14]-[16]], or can be directly reconnected to the plasmonic nanoparticles recognized as a colloidal method [[17]-[19]]. The colloidal method is intended to stimulate the particles of LSPR, whose EF is restricted to  $10^6$ . Moreover, with the support of an ordered surface structure, the EF contained in the substrate can be barely increased more than  $10^6$ .

Within the colloidal approach, bead, nano metallic particles, and pathogens are rarely connected to each other, and incident light can be restricted by those target cells, mostly if they are covered by the target cells. In the substrate method, pathogens will stand on the surface of plasmonic and incident light will be transmitted from the bottom to the top. Along with that, a huge portion of the incident light will be blocked/absorbed by the target cells and small number of photons can reach out to the plasmonic metal.

## II. FDTD SIMULATION

The SERS substrate determined in this paper is to make series transparent inverted nanocavity arrays from which a radius reduces beside the path of light. A reflectively high and bottom layer of the nanocavity series is covered with a reflected particle. In this context, these particles are multi-layered or reflected, which are known as photons. These are increased in density by an outside layer with a density of 5-10 times each. When this higher concentration of "photon gas" reaches the plasmonic metal, the Raman signal enhancement can be as high as  $10^8$ . Furthermore, the substrate is premeditated, which allows the light particles to transparently enter the bottom surface along with the non-photon particles to interrupt the process.

The incident photons are multi-reflected inside the cavity, known as cavity resonance. The intended resonance can capture light and concentrate it. In this work, the capturing of photons inside the structured cavity is utilized to demonstrate SERS along with the concentrated absorption of the photon. The numerical method used in this paper is Finite-Difference Time-Domain (FDTD) method, conducted by FDTD Solution (Lumerical Inc.) with 3nm cubic mesh. In the simulation, 2-D arrays of meshes have been utilized as observing planes, and the SERS refers to the maximum field enhancement in the observing plane. The ground surface of the substrate can be stated as Polydimethylsiloxane (PDMS) along with the light material used from top to bottom. A coated film in this procedure is utilized to enhance the substrate while the photons reflections can be enabled at EF  $10^{10}$ . The effected enhancement occurs only when the inverted cavity top surface is decreased due to the sloping of the cavity. Similarly, the gap between the two inverted cavity series must be enough to cover the magnetic beads, which can create the contact lesser.

By pouring liquid PDMS onto the structure and letting it cool down, the microstructure pattern can be transferred into a new structure. The geometric features of inverse cavity are bottom diameter 500 nm, top diameter 200 nm, height 1000 nm and periodicity 840 nm. Note this set of geometry is subject to change in the second section of the designing process [21, 22].

In the FDTD simulation, all structures are surrounded by water, and materials' optical models are obtained from Palik's book [23].

## III. RESULTS AND DISCUSSION

A 20 nm film is made on the surface of the inverse cavity, except the top. A plane wave source illuminates normally from the bottom base to the top. The incident electric field is parallel to the axis of the nanorod to excite LSPR. Because of this, the vertical axis electronic field is rounded with the nanorod so that the LSPR is not prompted with the entire structure; thus this transformation would not be comprehended. A 15nm diameter along with the length of the axis at 45nm Au nanorod will be used in the simulation, as it was previously used [24, 25, 26]. We are particularly interested in the field enhancement at the top of cavity, as in a real bio-sensing experiment, the top of the inverse cavity will be the most possible touching point of the structure and analyte. To better elucidate the role of the cavity's photon capturing to enhance the nanorod's LSPR, there is only one nanorod at the top of inverse cavity.



Distribution of the electromagnetic field in a cross section of structures along the axis of the nanorod will also be used in this work. In the flat PDMS surface and nanorod case, a layer of Au nanorods has been deposited at a flat PDMS surface, with the separation of every nanorod at 850 nm. A flat PDMS surface case stands for the commercial Raman substrate and will serve as the standard of comparison.

In order to prompt the resonated cavity, a higher level reflected interior cavity is needed. According to Fresnel's equations, a high reflection occurs when the refractive indexes of two interface materials are large [27]. Si and Au are chosen as coated films.

Because the refractive index for PDMS in the visible region is around 1.4, Au with an index around 0.3 and Si with an index around 4 are chosen as coating material. But due to the metallic nature of Au, the optical loss of the Au coated cavity can be high. The power dissipating profile of Si and Au deposited cavity are shown in Figure 1a and Figure 1b, in the cross section. The power dissipation can be defined as [28], which is directly correlated with light energy density [29]:

$$w(x, z) = \frac{1}{2} \epsilon_0 \omega \epsilon''(x, z) |\vec{E}(x, z)|^2 \quad (4)$$

In Fig. 1a and Fig. 1b, photon energy is mostly dissipated in the film for both Si and Au coated cases. But the intensity in the Au coated case is nearly 750 times larger than with Si film. Fig. 1c and Fig. 1d show the electric field intensity. In Fig. 1c, with Si coated, the electric field is enhanced around the top of cavity. With a gradually narrowing cavity and low optical loss, the photons are concentrated and delivered to the top of the cavity structure. In Fig. 1d, the electric field is enhanced at the side wall of the cavity, and the photons are dissipated and scattered at the cavity, before going to the top. Even photons are concentrated in a similar level in two cases, but the dissipation effect will create immense modification.

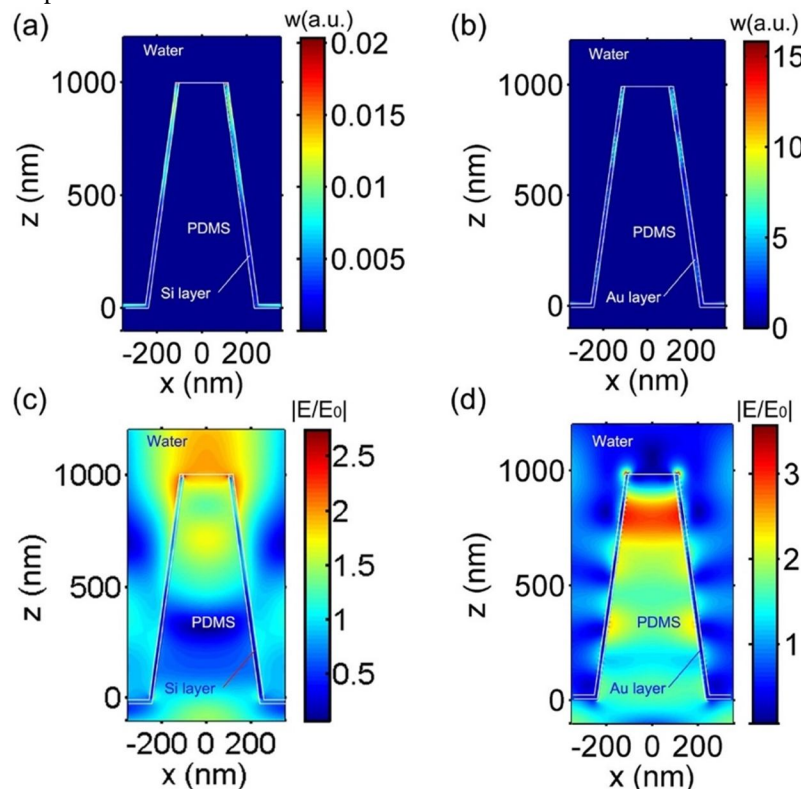


Figure 1: 758nm power dissipation distribution for cross sections of (a) SiCN without nanorod (b) AuCN without nanorod. 758nm electric distribution for cross sections of (c) SiCN without nanorod (d) AuCN without nanorod.

Figure 2a and Fig. 2b show the 758nm field distribution in Si and Au coated inverted nanocavity with nanorod at the tops. The cross section is illustrated in Fig. 2d. Note that the top is free of coating to ensure that concentrated photons will not be reflected by the interface. In Fig. 2a, the electric field is highly confined around the nanorod, the top of the inverted structure. Inside the cavity, the electric vectors exhibit a multi-reflection resonance pattern, exactly the cavity resonance mode. In Fig. 2b, the electric field is also highly confined around the nanorod. In Fig. 2a and Fig. 2b, it can be seen that the enhancement patterns are exactly same, but the intensities are different.

But with the help of Si film, the electric field enhancement can be improved nearly 5 times that of the Au coated case. Fig. 2c shows the SERS as a function of wavelength at the top observing plane for Si coated, Au coated and flat surface cases. One can see at 758nm, the SERS reaches  $10^8$  for the Si coated case,  $10^6$  for the flat surface case and  $10^3$  for the Au coated case. With the help of a low-loss photon-concentrated Si coated cavity, the SERS can be improved by  $10^2$ . However, even with the assistance of the cavity, the Au loss reduces the SERS a lot, making field enhancement lower than isolated LSPR.

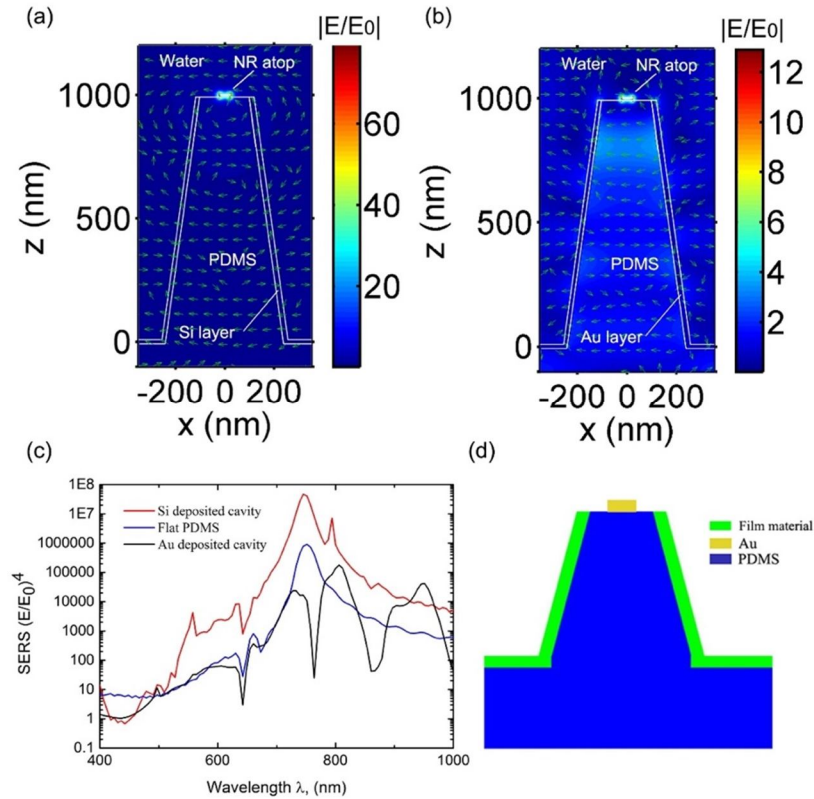


Figure 2: 758nm electric field in cross sections of (a) Si (b) Au deposited inverse cavity without nanorod. (c) SERS spectrum for 3 related structures. (d) Illustration of cross section of the proposed structure.

In order to analyze the optimum bottom and slope inverted nanocavity diameter, the comparative concentration  $I_{relative}$  has a correlation [[10]]:

$$I_{relative} \propto \sum EF_{top} \times N_{molecule} \tag{5}$$

$N_{molecule}$  is the molecules number at the top surface, and is comparative high range  $S_{top}$ . The enhancement at the top  $EF_{top}$  is certainly connected to the inverted cavity slope whereas  $S_{top}$  is adversely linked. For finding the tradeoff amid  $EF_{top}$  and  $S_{top}$ , a function is described as:

$$f_{inverted\ cavity} = \frac{EF_{top} \times S_{top}}{\sum S_i} \tag{6}$$

Where  $\sum S_i$  is the total area that a single cavity averagely takes. This function is actually the ratio between signal intensity of inverted nanocavity arrays to flat surface under the same rod deposition density. For the flat top area  $f = 1$ , and for our first harvesting nano cavity prepared combination  $f = 1$ . The determination is to max out  $f$ . The  $EF_{top}$  is premeditated via:

$$\sqrt{EF_{top}} = \frac{S_{top} + (\pi D^2 - S_{top}) \times \bar{R} (0.5\pi - \theta)}{S_{top}} \tag{7}$$

$\bar{R}$  is the polarization averaged reflection of the film, obtained from Fresnel's equation. In the calculation, given that the film thickness is 15nm and much smaller than the wavelength in the film, thin film limit is introduced where wavevectors in PDMS and water are close to free space and wave vector in film is nearly independent of radius  $a$  [30-33].

#### IV. CONCLUSION

In this work, an SERS substrate is proposed for the rapid diagnosis of COVID-19. The high sensitivity originates from the combination of photon funneling, dense packing and an optimized inverted nanocavity. 15nm thick Si film is deposited at the surface of the inverted cavity structure, which has high reflection and a low loss concentrating of photons. The Raman enhancement factor is increased by 2 orders of magnitude, while only reducing contact area by 1 order and making the overall sensitivity 1 order higher as compared with the rough surface approach. The Si coated structure proposed in this paper has the advantages of low-cost and large-scale productivity, and fits immunomagnetic pathogen collecting well. This work will facilitate the development of novel optical bio-substrate along with fast and precise pathogen diagnosis.

#### V. ACKNOWLEDGMENTS

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