Sensitivity enhanced biosensor using graphene-based one-dimensional photonic crystal

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A B S T R A C T

In this paper, we propose and analytically demonstrate a biosensor configuration based on the excitation of surface electromagnetic waves in a graphene-based one-dimensional photonic crystal (1D PC). The proposed graphene-based 1D photonic crystal consists of alternating layers of high (graphene) and low (PMMA) refractive index materials, which gives a narrow angular reflectivity resonance and high surface fields due to low loss in PC. A differential phase-sensitive method has been used to calculate the sensitivity of the configuration. Our results show that the sensitivity of the proposed configuration is 14.8 times higher compared to those of conventional surface plasmon resonance (SPR) biosensors using gold thin films. This novel and effective graphene-based 1D PC will serve as a promising replacement of metallic thin film as a sensor head for future biosensing applications.

1. Introduction

In the past decade, surface plasmon (SP) concept has found tremendous applications in variety of areas such as biosensors, solar cells, lithography, etc. [1–4]. The SP based biosensors for instance has enabled high speed and high sensitive biomolecular detection. This is mainly because they eliminate time-consuming labeling process and reduce molecular binding disturbance compared to commonly employed fluorescence based optical biosensors. However, the damping of light intensity caused by the metal absorption loss affects the sensitivity of biosensors for various applications [5,6]. In this context, excitation of surface electromagnetic waves (SEWs) in one-dimensional (1D) photonic crystals (PC) has been investigated for various potential applications such as sensors, fluorescence emission enhancement, and enhancement of the Goos–Hänchen effect [7–10]. Surface electromagnetic waves are non-radiative electromagnetic modes that appear on the surface of semi-infinite 1D photonic crystal, which is considered to be an alternative to surface plasmon [11,12]. The low loss in photonic crystals may provide higher intensity of SEWs, which leads to much sharper resonance dips and high surface electromagnetic fields. More recently, our group has been experimentally demonstrated the excitation of surface electromagnetic waves in a graphene-based 1D PC using prism coupling technique [13].

Since graphene is a single two-dimensional plane of carbon atoms forming a hexagonal lattice, the optical properties of graphene has been extensively investigated both theoretically and experimentally for many photonics and optoelectronics applications due to its strong interaction with light in a broad wavelength interval [14–17]. In particular, graphene exhibits universal optical conductivity from visible to infrared frequencies due to interband transition [18]. Recently, graphene-based photonic crystals at THz frequencies have been proposed and theoretically investigated for the development of frequency filters and waveguides [19]. The photonic crystal waveguide analogy of graphene nanoribbons have been theoretically investigated by Benisty [20]. More recently, Yan, et al. has reported the fabrication of tunable infrared plasmonic devices using photonic crystal like structure (graphene/insulator stacks) [21]. These reported works may provide an effective way in designing graphene-based optoelectronics devices. In addition, graphene has been recently started using in surface plasmon resonance biosensor to enhance the adsorption of biomolecules and hence to improve the sensitivity of the biosensors [22–25]. Here, we propose a biosensor configuration based on the excitation of surface electromagnetic waves in a graphene-based 1D photonic crystal and analytically investigate the sensitivity of the configuration.

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Fig. 1. Schematic diagram of (a) proposed biosensor configuration and (b) surface plasmon resonance biosensor. Graphene-based 1D photonic crystal consists of alternating layers of graphene and PMMA layers with an additional graphene termination layer.

2. Theory

The schematic diagram of proposed biosensor configuration based on prism coupling technique is shown in Fig. 1(a). The graphene-based 1D PC is attached via index matching fluid on the top of the BK7 glass prism. The binding layer of thickness 3 nm and refractive index 1.462 (n2) is assumed as ssDNA layer with a density of 0.028 g/cm², which is prepared in water and covers the whole PC [23]. The graphene-based 1D PC consists of alternating layers of high and low refractive index materials such as graphene and poly methyl methacrylate (PMMA), respectively deposited on the BK7 glass slide. The wavelength of excitation light source is taken as 633 nm. Here, we assume that the graphene is a dielectric material with isotropic refractive index and light polarization does not affect on graphene refractive index because of the universal opacity of graphene at optical frequencies. The refractive index of graphene calculated at 633 nm wavelength is n1 = 3 + 1.149i [26]. The refractive index of BK7 prism (n0), PMMA (n2) and water (n4) is taken as 1.515, 1.49 and 1.33, respectively. The thicknesses of graphene and PMMA are taken as 0.34 × L nm (d1) (where L = 1, number of graphene layer) and 470 nm (d2), respectively. The proposed periodic structure represents a Bragg grating with a Bragg wavelength of 1.4 μm (calculated using λB = 2(n1d1 + n2d2)). The designed 1D PC consists of 9 layers of graphene/PMMA with an additional graphene termination layer. It is possible to move the surface mode dispersion through the photonic bands from one edge to the other when the termination layer of the photonic crystal is varied [27]. In order to compare the results of proposed configuration, conventional surface plasmon resonance biosensor geometry is also investigated (as shown in Fig. 1(b)). In Fig. 1(b), the metal is taken as Au film with thickness 45 nm. The refractive index of Au at 633 nm wavelengths is 0.1968 + 3.09i [28]. The remaining parameters are same as those of proposed configuration (Fig. 1(a)).

In order to determine the sensitivity of both configurations, a differential phase sensitive method has been employed [29,30]. In this method, a standard Mach–Zehnder interferometer setup can be used to extract the phase difference between the interfering signal of TM-polarization (signal) and TE-polarization (reference). The phase difference between TM- and TE-polarization is given by, Δφ = φTM − φTE. The values of φTM and φTE can be extracted from the Fresnel’s reflection coefficients of TM- and TE-polarization, rTM = |rTM|exp(iφTM) and rTE = |rTE|exp(iφTE), respectively. The reflectance (|rTM,TE|^2 = R) of TM- and TE-polarization can be calculated by solving the Fresnel’s equations for N-layer model using transfer matrix method (TMM). Here, the reflectance (R) is serial product of the interface matrix ljk (j = 0, 1, 2, 3 and k = j + 1), and layer matrix Lj

\[
R = \left| \frac{M_{12}}{M_{22}} \right|^2
\]

with

\[
M = \begin{bmatrix}
M_{11} & M_{12} \\
M_{21} & M_{22}
\end{bmatrix}, \quad l_{jk} = \begin{bmatrix}
1 & r_{jk} \\
r_{jk} & 1
\end{bmatrix} \quad \text{and} \quad L_j = \begin{bmatrix}
e^{ik_jd_j} & 0 \\
0 & e^{-ik_jd_j}
\end{bmatrix}
\]

where d_j is the thickness of jth layer, and

\[
k_{jk} = \sqrt{s_j \left( \frac{2\pi}{\lambda} \right)^2 - k_x^2}
\]

In the case of TM polarization

\[
r_{jk} = \frac{(k_{jk}/\epsilon_j) - (k_{jk}/\epsilon_k)}{(k_{jk}/\epsilon_j) + (k_{jk}/\epsilon_k)}
\]

In the case of TE polarization

\[
r_{jk} = \frac{(k_{jk}/\sqrt{\epsilon_j\epsilon_k}) - (k_{jk}/\sqrt{\epsilon_j\epsilon_k})}{(k_{jk}/\sqrt{\epsilon_j\epsilon_k}) + (k_{jk}/\sqrt{\epsilon_j\epsilon_k})}
\]

In Eq. (3), k_s is the parallel wave vector, and given by k_s = n_0(2π/λ)sinθ, where λ is excitation wavelength and θ is the incident angle.

3. Results and discussion

Based on the TMM method, the reflectance data is simulated separately for both configurations. The excitation of surface electromagnetic wave is recognized as the angle at which minimum reflected intensity is occurred (ATR minimum) and this ATR minimum angle is taken as the resonance angle. Fig. 2(a) represents the reflectance diagram of proposed configuration using TM-polarization. From the figure, it is evident that the obtained resonance angle is 79.2°, which is greater than the angle for total internal reflection (TIR) between prism and water (61.3°). Hence this deepest dip at resonance angle corresponds to the non-radiative SEW propagating at the 1D PC/binding layer boundary. Fig. 2(b) shows the reflectance curve obtained for surface plasmon resonance biosensor and observed resonance angle is 75.25°. It is visible from the Fig. 2 that the proposed biosensor configuration provides a deepest, narrow dip as compared to surface plasmon resonance biosensor. It shows that the graphene-based 1D PC has low loss and maximum surface mode excitation compared to surface plasmon resonance biosensor.

The sensitivity of the biosensor is defined as, S = (Δφ/Δn3)E, where the first term (Δφ/Δn3) represents the ratio of the change in the differential phase between TM- and TE-polarization (Δφ) to the change in the refractive index of the binding layer (sensing medium) (Δn3). And the second term, E is the adsorption efficiency of the target biomolecule on the binding layer. We assume that
the refractive index of binding layer \( n_1 \) increases from 1.462 to 1.466 in an interval of 0.0001, when the adsorption of biomolecules occurs and hence the total refractive index change is given by, \( \Delta n_3 = 0.004 \). The relative differential phase change between TM- and TE-polarization with refractive index of sensing medium is shown in Fig. 3. Fig. 3(a) and (b), respectively represent the differential phase change curve of proposed biosensor and surface plasmon resonance biosensor, and both curves linearly increase when the refractive index of the sensing medium increases. In the case of proposed configuration, relative differential phase change varies from 135° to 183° for the total refractive index change of 0.004, while the surface plasmon resonance biosensor has a slight variation of 99.7° to 103° for the same refractive index change. Hence the calculated sensitivity for proposed biosensor and surface plasmon resonance biosensor are \( 1.2 \times 10^4 \)/RIU and \( 8.25 \times 10^2 \)/RIU, respectively. It shows that the sensitivity of the proposed configuration is 14.8 times more than those of a surface plasmon resonance biosensor.

The influence of number of graphene layers in the 1D PC on sensitivity enhancement is also investigated and plotted in Fig. 4(a). It is found that the maximum sensitivity is observed when the 1D PC consists of single layer graphene and there is a sudden decrease in sensitivity for bilayer graphene. After that the sensitivity is almost constant when the graphene layers in the PC increases. It can be attributed presumably to the reduction in light intensity due to the increase of graphene layers (2.3% light absorption in the case of monolayer graphene). It shows that the proposed configuration can provide maximum sensitivity when the photonic crystal consists of single layer graphene. The sensitivity variation of the surface plasmon resonance biosensor is also analyzed by varying the thickness of the gold thin film (as shown in Fig. 4(b)). In this case, the Au film thickness is varied from 30 nm to 60 nm and maximum sensitivity is observed for 45 nm. This thickness (45 nm) represents the optimum thickness for maximum sensitivity and the corresponding sensitivity value is almost equal to those of a 4 layer graphene-based proposed sensor. In addition, the sensitivity of biosensor configuration has been studied by varying the number of bilayers (graphene/PMMA) and note that maximum sensitivity is obtained for 9 bilayers. Further, we observed that the sensitivity of the proposed configuration decreases when the excitation wavelengths increases. It could be due to the increase of graphene extinction coefficient with increase in excitation wavelengths.

In order to confirm the existence of surface electromagnetic waves in the proposed biosensor configuration, the surface electromagnetic field distribution is simulated using finite difference time domain (FDTD) method. In the simulation, the graphene thickness is assumed as 1 nm and hence the smallest spatial grid size of 0.1 nm is used for the iteration to maintain the accuracy and stability of FDTD calculations. Here we assume that the light is incident at a resonant coupling angle of 79.2° through the prism. The electric field distribution and its magnitude as a function of depth along the 1D PC is shown in Fig. 5. Note that the output electric field is normalized with respect to the incident excitation field. It is visible from the figure that the intensity is decaying toward the prism/1D PC interface and field oscillates many times throughout the periodic structure. It is also clear from the figure that when the surface mode is excited, the electromagnetic field enhancement occurs within the last layer of the 1D PC (1D PC/binding layer interface). It shows that the mode at the band edge has less attenuation and thus the decaying evanescent filed can penetrate much
further into 1D PC. Maximum field enhancement factor is observed at PC/binding layer interface, which is due to the tighter confinement of mode to the surface. This large enhancement factor is responsible for the observed enhanced sensitivity of the proposed graphene-based 1D PC biosensor. In addition, the excitation of SEW in the proposed structure can be confirmed by studying the surface dispersion diagram [13]. As reported in our previous work [13], the SEW exists in the non-radiative region of the surface dispersion diagram and away from the air light line. It means that SEWs cannot radiate either into the air side or into the 1D PC. The parallel wave vector of the mode calculated at 633 nm incident wavelength using the expression \((2\pi/\lambda)\eta_0 \sin \theta\) is 1.476 \times 10^7 \text{ rad/m}. We found that this value occur close to the second band edge and appear away from the air light line, which further confirm the excitation of SEW in the proposed graphene-based biosensor configuration.

4. Conclusion

A biosensor configuration based on the excitation of surface electromagnetic waves in a graphene-based 1D photonic crystal is presented and the results are then compared with that of a conventional surface plasmon resonance biosensor. The existence of SEW in the proposed biosensor configuration is numerically investigated for enhanced sensitivity. About 15% increment in sensitivity is obtained compared to surface plasmon resonance biosensor and it is also evident from the results that the number of graphene layer is an important parameter to be considered for improving the sensitivity when designing efficient biosensor geometries. The experimental study is needed to further support the proposed concept.

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