Black phosphorus integrated tilted fiber grating for ultrasensitive heavy metal sensing

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Research Highlights:

(1) The first black phosphorus (BP)-fiber chemical sensor for heavy metal (Pb\textsuperscript{2+} ions) detection.
(2) An integrated BP-tilted fiber grating configuration is proposed for strong light-matter interaction.
(3) A new in-situ layer-by-layer technique is developed for BP nanosheets deposition on fiber surface.
(4) Unique optical tunable features including coating thickness-dependent polarization and polarization-selective coupling have been experimentally observed.
(5) Significant performance for Pb\textsuperscript{2+} detection with ultrahigh sensitivity (0.5×10\textsuperscript{-3} dB/ppb), lower limit of detection (0.25 ppb), and wider concentration range (0.1 ppb to 1.5×10\textsuperscript{7} ppb).
(6) Limit of detection of 0.25 ppb is 40 times lower than WHO’s permissible limit of 10 ppb for lead in drinking-water. The detectable Pb\textsuperscript{2+} concentration range is 4 orders of magnitude larger than that of BP-FET based lead sensor.

ABSTRACT

We propose an integrated black phosphorus (BP)-tilted fiber grating (TFG) configuration as an enhanced light-matter interaction platform for heavy metal chemical sensing. We synthesized BP nanosheets by using a liquid-phase exfoliation method and developed an in-situ layer-by-layer (i-LbL) technique for BP nanosheets deposition. These approaches secured high-quality BP coating on specific fiber cylindrical surface with strong adhesion as well as a prospective thickness control. By taking advantage of i-LbL deposition, the unique optical tunable features including coating thickness-dependent polarization and polarization-selective coupling have been experimentally observed. Furthermore, the BP-TFG was exploited as the first BP-fiber optic chemical sensor for heavy metal Pb\textsuperscript{2+} ions detection, demonstrating
significant performance of ultrahigh sensitivity up to $0.5 \times 10^3$ dB/ppb, much lower limit of detection down to 0.25 ppb, and extremely wide concentration range from 0.1 ppb to $1.5 \times 10^7$ ppb. We believe that this BP-fiber optic architecture opens the path as an optical platform for superior chemical sensing and biomedical applications.

**Keywords:** Black phosphorus; Nanomaterials; Fiber grating; Optical sensor; Chemical sensor; Heavy metal.

1. Introduction

Two-dimensional (2D) layered materials, which include graphene, transition metal dichalcogenides (TMDs), and boron/carbon nitride, have attracted significant attentions [1-4]. Due to their exceptional properties of large surface-to-volume ratio, atomic-thin layer structure and high carrier mobility, 2D materials show a wide range of applications in electronics, light processing, energy storages, and chemical and biochemical sensors [5-8]. Very recently, black phosphorus (BP) sparks intense interest of scientific communities in physics, chemistry, materials, and biology [9-16]. BP has a puckered lattice configuration along the armchair direction and a bilayer structure along the zigzag direction [11]. Thanks to the unique puckered orthorhombic structure and in-plane anisotropy, BP possesses much larger surface-to-volume ratio, extremely high hole mobility and superior molecular adsorption energy than those of other 2D materials [9,15,16]. The extraordinary properties of BP make it a very promising nanomaterial for widespread applications such as BP field-effect transistors (FETs), modulators, photodetectors, chemical sensors, bioimaging and biomedical therapy [9-11,16-20].

The novel lab-on-fiber technology has been proposed for optical sensing and optical communications [21,22]. Besides the inherent merits of optical fiber, such as compact size, light weight, and electromagnetic interference immunity, in-fiber grating technology provides advantages of real-time, multiplex, high sensitivity, label-free, and in-line determination [23-28]. Both long period fiber grating (LPG) and tilted fiber grating (TFG) can couple the light from fiber core to the cladding yielding an evanescent wave penetrating outside of the cladding. As the evanescent wave can be affected by changes in concentration or composition of the surrounding medium or by molecular interactions occurring on the cladding surface, it is a good candidate for chemical and biochemical sensing [23-24]. In contrast to conventional LPG, TFG owns narrow band resonances demonstrating much higher Q-factor hence high accuracy in optical signal detection [25-28]. In particular, the excessively tilted fiber grating shows much lower thermal and tension cross-talk effects and higher sensitivity in lower refractive index (RI) region [25].

Techniques have been developed to improve the sensitivity of fiber devices by fiber tapering, cladding etching and side polishing. Instead of sacrificing the mechanical integrity, thin film deposition is an alternative elegant approach. To date, various materials including carbon nanotubes (CNTs), titanium dioxide, zinc oxide, and graphene have been employed to enhance the RI sensitivity of fiber sensors [29-32]. However, the grand challenge of film deposition on optical fiber is from its cylindrical geometry and such a thin diameter. The recent exploitation reveals that 2D layered materials could modulate light with superior performance for realistic applications in ultrafast lasers, broadband polarizer, gas detection, humidity sensor, and biosensing [6, 7, 33-36].

Heavy metals are usually considered to be toxic and carcinogenic with the harmful effects on the environment, drinking-water and living organisms. It needs to be recognized that lead is exceptional compared with other chemical hazards. Exposure to lead is associated with a wide range of effects, including various neuro developmental effects, cardiovascular diseases, impaired renal function, hypertension, and impaired fertility. According to the guidelines for drinking-water quality of World Health Organization (WHO), the maximum permissible limit of lead in drinking-water is 10 ppb [37], which represents the lead concentration does not exceed tolerable risk to the health. Therefore, the detection of heavy metal becomes a major public concern in modern days.

In this work, we report an integrated BP-fiber optic configuration as an optical tunable platform for the enhancement of light-matter interaction and heavy metal detection. As the schematic illustration shown in Fig. 1, the light passes
down the core of a fiber and encounters a 81°-TFG, certain wavelength of the light are forward-coupled into the cladding yielding an evanescent wave penetrating into BP coating, demonstrating strong light-matter interaction. We synthesized BP nanosheets by using a liquid-phase exfoliation method and developed an in-situ layer-by-layer (i-LbL) deposition technique. These approaches secured high-quality BP overlay deposited on specific cylindrical surface of fiber with strong adhesion as well as a prospective thickness control. The unique optical tunable properties have been experimentally observed. Furthermore, the BP-TFG was exploited as the first BP-fiber chemical sensor for heavy metal detection with significant performance.

2. Materials and methods

2.1. Fabrication of 81°-tilted fiber grating

The 81°-TFG with length of 12 mm was UV-inscribed in hydrogen loaded Corning SMF-28 fiber by using a frequency-doubled Ar laser and mask scanning technique. A custom-designed mask with a pitch of 6.6 μm was purchased from Edmund Optics Ltd. The mask was rotated in the plane of incidence to realize grating fringes tilted by 81° with respect to the fiber axis. Multiple scans were applied to achieve a maximum forward cladding mode coupling at the wavelength determined by the following phase-matching condition:

\[ \lambda_{\text{eff}} = \left( n_{i,\text{eff}}^\text{core} - n_{m,\text{eff}}^\text{cladding} \right) \frac{\Lambda_g}{\cos \theta} \]

where \( n_{i,\text{eff}}^\text{core} \) is the effective index of fiber core, \( i \) represents transverse-magnetic (TM) or transverse-electric (TE) polarization state, \( n_{m,\text{eff}}^\text{cladding} \) is the effective index of cladding, \( m \) means \( m \)th mode, \( \Lambda_g \) is the normal period of grating, and \( \theta \) is the tilted angle.

2.2. Synthesis of BP nanosheets

The BP nanosheets were synthesized by using a liquid-phase exfoliation method, which was similar to our previous work [20], except the solvent in this work was changed with absolute ethanol. As shown in Fig. 2: (i) 25 mg bulk BP was cut and grinded into small pieces then added to 25 mL absolute ethanol; (ii) The BP dispersion was sonicated by a cell crasher at 25 kHz and 1200 W for 3h to break the van-der Waals stack of BP, where the temperature of solution was kept below 277 K with an ice bath; (iii) The as-prepared dispersion was centrifuged at 5000 rpm for 15 min; (iv) the supernatant containing few-layered BP nanosheets was decanted gently.

2.3. BP deposition on fiber surface

For BP deposition, we developed an i-LbL technique, which was based on chemical surface modification in conjunction with in-situ layer-by-layer assembly. The deposition process consists of: (v) The section of fiber over TFG region was rinsed with acetone to clean the surface from any contaminants. Thereafter, an alkaline treatment was carried out by immersing the fiber device in 1.0 M NaOH solution for 1 h to enrich the number of -OH groups on the fiber surface, then washed with DI water and ethanol thoroughly; (vi) A silanization procedure was followed by incubating the fiber in freshly-prepared 5% (3-Aminopropyl)triethoxysilane (APTES) for 2 h to form Si-O-Si bonding; (vii) The fiber was fixed into a homemade microchannel container where 30 μL BP dispersion was carefully added. Due to positively-charged amino groups on the APTES-silanized fiber surface, the negatively-charged BP nanosheets were adhered to fiber by electrostatic force. After ethanol solvent fully evaporated, the fiber surface was dried and coated with BP nanosheets (Fig. S1). Subsequently, the second deposition cycle was conducted. In each cycle, BP dispersion (30 μL) was added in microchannel container where ethanol solvent was evaporating gradually and totally, such drying process enhanced the physisorption of BP nanosheets on fiber surface. After 8 cycles, the
BP-coated TFG was rinsed by absolute ethanol to remove non-adhered BP nanosheets, dried and stored in a vacuum environment to enhance the layer-by-layer adhesion before further uses. The transmission spectra of BP-TFG were monitored in real-time during the entire BP deposition process, which will be discussed later.

3. Results and discussion

3.1. Surface morphological characterization

The surface morphological characterization was investigated by using digital microscope (Keyence VHX-2000), Raman spectroscopy (Horiba Jobin-Yvon Lab Ram HR VIS, 633 nm), scanning electron microscope (SEM, Zeiss SUPRA 55), and atomic force microscopy (AFM, Veeco Instruments Inc.).

The initial microscope image Fig. 3(a) shows a successful overlay coated on fiber surface. Subsequently, Raman spectroscopy was used to characterize the coating material. Raman spectrum in Fig. 3(b) presents three characteristic peaks, which include $A_{1g}^1$ (at ~362 cm$^{-1}$), $B_{2g}$ (at ~440 cm$^{-1}$) and $A_{2g}^2$ (at ~466 cm$^{-1}$), indicating the presence of black phosphorus [11]. The $A_{1g}^1$ corresponds to atoms oscillate out-of-plane while $B_{2g}$ and $A_{2g}^2$ correspond to atoms vibrate within the plane. The surface coverage of BP was further examined by SEM with the magnification of 1000× (Fig. 3c) and 10000× (Fig. 3d). It can be seen that the BP nanosheets are densely covered on the fiber and BP overlay is quite homogeneous over the entire cylindrical surface demonstrating the high-quality BP deposition.

The thickness of BP overlay depends on the conditions of deposition. With the flexibility of i-LbL deposition technique, the overlay thickness can be prospectively controlled by the number of coating cycles. Fig. 3(e) and Fig. 3(f) plot AFM tapping mode topographic images of BP overlay on fiber surface after 8 coating cycles, in which the thickness of BP overlay was measured as ~323.1 nm.

3.2. Observation of optical tunable properties

The optical properties of BP-TFG were experimentally investigated by monitoring BP-TFG transmission spectra over entire BP deposition process. For a 81°-TFG, the excessively tilted plane of index-modulated grating breaks the symmetry of mode coupling, inducing the significant polarization dependent coupling characteristics. 81°-TFG exhibits two sets of comb-like split resonances in transmission spectra, which suggest that the light have been coupled into two sets of birefringence modes corresponding to two orthogonal polarization modes, TM and TE modes [27]. The polarization modes can be fully excited by launching the linearly polarized light. Spectra (i) in Fig. 4(a) show a zoomed-pair of polarization modes, in which TM and TE modes have been fully excited, respectively. The transmission spectra of BP-TFG were monitored during deposition process and recorded as (ii), (iii), (iv) and (v) in Fig. 4(a) after 2nd, 4th, 6th and 8th coating cycle, respectively.

It is apparent that the polarization depends on BP overlay thickness. As can be seen from Fig. 4, optical signals (both TM- and TE-polarized resonances) show a significant red-shift in wavelength and a decrease in intensity against the increase of overlay thickness which was determined by number of coating cycles. The wavelength shift was attributed to the increase of cladding effective refractive index induced by high RI coating layer, and consequently the phase-matching condition Eq. (1) was tuned.
leaky radiation from cladding to coating layer could reduce the coupling coefficient between core and cladding modes hence decrease the resonant intensity.

It should be emphasized that the extinction ratio has been extremely enhanced with the increase of BP overlay thickness, meanwhile a polarization-selective coupling feature has been clearly observed. As shown in Fig. 4, the variation of TE mode is always larger than that of TM mode. After 8 cycles of BP deposition, TE mode has 16.9 nm in wavelength shift (Fig. 4b) and 4.35 dB in intensity decrease (Fig. 4c), which are 1.2 times and 2.1 times larger than those of TM mode, respectively. The separation between two polarized resonances of BP-TFG has been increased to 8.38 nm, giving an estimated birefringence of $-3.0 \times 10^{-7}$ which is 150% enhancement that of bare 81°-TFG. In such BP-TFG architecture, TE-polarized radiative light can be strongly perturbed by BP overlay in terms of coupling and guiding, leaving the TM-polarized light passes through the fiber.

3.3. Strong ability of Pb$^{2+}$ adsorption and ultrahigh sensitivity

The BP-TFG was exploited for the implementation of BP strong ability of lead ions adsorption and ultrahigh sensitivity for heavy metal chemical sensing.

The experiments were conducted with the setup illustrated in Fig. 5. The light from broadband source was launched into BP-TFG and the output was monitored by an optical spectrum analyzer (OSA). By adjusting the polarization controller, the TM-polarized resonance was fully excited and selected for optical signal monitoring. To avoid thermal and tension cross-talk effects, the BP-TFG was mounted on a homemade mini-bath with two ends fixed and all the experiments were implemented at room temperature. A series of aqueous solutions containing Pb$^{2+}$ ions with concentrations of 0.1, 1.0, 10, 100, 1000, 1×10$^5$, 1×10$^6$, 1×10$^7$ and 1.5×10$^7$ ppb were prepared for the detection. The BP-TFG was firstly immersed into Pb$^{2+}$ solution for 120 s, the solution was subsequently withdrawn by careful pipetting to expose the BP-TFG in the air for 180 s when the transmission spectrum achieved steady and was captured by OSA. Afterwards, the BP-TFG was washed with ethanol thoroughly to remove the adsorbed lead ions prior to next concentration measurement.

The transmission spectra at different Pb$^{2+}$ concentrations are plotted in Fig. 6(a). By increasing Pb$^{2+}$ concentration, the peak shows apparent intensity decrease indicating the strong optical absorption occurred between BP and lead ions. The slight wavelength red-shift could be caused by the increase of effective refractive index of cladding. Fig. 6(b) depicts the intensity change as a function of ion concentrations, revealing a non-linear relationship with piecewise sensitivities of 0.5×10$^{-3}$ dB/ppb, 7.7×10$^{-7}$ dB/ppb and 2.3×10$^{-8}$ dB/ppb for Pb$^{2+}$ concentration ranges of 0–100 ppb, 10$^5$–10$^6$ ppb, and 10$^6$–10$^7$ ppb, respectively. The proposed BP-TFG can effectively detect Pb$^{2+}$ ions concentration over a wide range from 0.1 ppb to 1.5×10$^7$ ppb with no saturation effect, which is four orders of magnitude larger than that of BP-FET based lead sensor [17]. The reported BP-FET sensor limited its detection for Pb$^{2+}$ concentrations lower than 10 ppb and faced the saturation problem for concentrations higher than 1×10$^7$ ppb.

The limit of detection (LOD) is generally defined as three times the standard deviation of its blank measurement. The blue curve in Fig. 6(b) provides the best logistic fitting of the experimental data, whereas the dashed red line represents intensity change three times the standard deviation. Therefore, the LOD of BP-TFG is 0.25 ppb for Pb$^{2+}$ ions, which is 4-fold better than CNTs functionalized heavy metal sensor [38]. Meanwhile, this LOD of 0.25 ppb is only one fortieth of WHO’s permissible limit of 10 ppb for lead in drinking-water [37]. For comparison, the similar detection was also conducted for light metal Na$^+$ ions with the results plotted as green symbols in Fig. S2, showing a very limited sensitivity. It reveals that BP exhibits strong adsorption of heavy metal ions hence the ultrahigh sensitivity for lead detection. In addition, the proposed BP-TFG has been demonstrated its excellent reusability for lead ions detection (Fig. S3), showing the potential to develop a more cost-effective and reusable chemical sensor.

3.4. Heavy metal adsorption – induced optical signal change

We discuss the basic mechanism underpinning the unique optical properties of BP-TFG with the adsorption of heavy metal ions. It has been reported that BP band gap can be widely tunable by thickness, strain, electric field and dopant [12-14]. As the pristine BP behaves as a p-type semiconducting characteristic, its majority charge carriers are holes. The adsorption of heavy metal ions denoted charges and enhanced hole carrier concentration in BP. With increasing ion density, the electric field from the ionized donors gradually reduced band gap owing to the giant Stark effect and tuned the electronic state of BP from a semiconductor to a Dirac semimetal [12]. Due to its ultrahigh carrier mobility and superior molecular adsorption energy, BP has strong capability to attract heavy metal ions, which is more efficient for charge transfer between BP and heavy metal ions. When the carrier concentration was large enough, correlation effects became significant and resulted in an increase of carrier-induced refractive index change. Consequently, the fiber guided wave was modulated by perturbing phase condition and optical absorption coefficient.
. Conclusions

In conclusion, we have reported an integrated BP-TFG configuration for enhancement of light-matter interface and for chemical sensing. The developed i-LbL deposition technique secured high-quality BP overlay and prospective thickness control. We have experimentally observed the unique optical properties including overlay thickness-dependent polarization and polarization-selective coupling. With BP deposition, both phase condition and light coupling coefficient have been modulated with the corresponding resonant wavelength shift and intensity change. The BP-TFG has been implemented for Pb^{2+} ions detection, demonstrating ultrahigh sensitivity up to 0.5×10^{-4} dB/ppb, much lower LOD of 0.25 ppb, and extremely wide concentration range from 0.1 ppb to 1.5×10^{7} ppb. The achievable LOD is 40 times lower than WHO’s permissible limit for lead in drinking-water [37].

The wide detection range is 4-order of magnitude larger than that of BP-FET based lead sensor [17]. The proposed BP-fiber optic architecture provides a remarkable platform for chemical and biochemical applications.

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References


Biographies

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Fig. 1. Schematic of BP-TFG configuration as an optical tunable platform for strong light-matter interaction.
Fig. 2. Schematic representation of synthesis of BP nanosheets (i-iv) and deposition process of BP nanosheets on fiber device surface (v-vii).
Fig. 3. Surface morphology of BP-coated fiber. (a) Microscope image (scale bar: 50 µm), (b) Raman spectrum, SEM images with magnification of (c) 1000× (bar: 10 µm) and (d) 10000× (bar: 1 µm), Atomic force micrograph topographic (e) 3D image and (f) profile of BP coating boundary (inset: height profile of BP overlay).
Fig. 4. (a) Spectral evolution during BP-deposition process revealing strong light-matter interface and optical tunable features. (b) Wavelength shift and (c) intensity change of TM- and TE-polarized resonances against number of deposition cycles.
Fig. 5. Experiment setup for heavy metal chemical sensing.
Fig. 6. (a) Transmission spectra of TM-resonance showing a clear up-shift as Pb$^{2+}$ concentration increases. (b) Resonant intensity change against Pb$^{2+}$ concentrations. The blue line is the best logistic fitting curve and the dashed horizontal red line represents three times the standard deviation of the blank measurement.