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Double Stopband Bilayer Photonic Crystal Based Upconversion Fluorescence PSA Sensor

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Abstract

Precise detection on tumor markers is significant for the early diagnosis on cancer. Upconversion fluorescence probes are ideal for tumor marker sensing, but still subject to bottleneck problems of low emission efficiency/strength. In this work, a novel bilayer of poly(methyl methacrylate) opal photonic crystal (PMMA OPC) with double photonic stop bands was fabricated to match the excitation field and emission field of the upconversion nanoparticles (UCNPs) at the same time, and greatly enhance their upconversion fluorescence intensity. Herein, NaYF\(_4\):Yb\(^{3+}\),Er\(^{3+}\) UCNPs and NaYF\(_4\):Yb\(^{3+}\),Er\(^{3+}\)@NaYF\(_4\):Yb\(^{3+}\),Nd\(^{3+}\) UCNPs with a core-shell structure were employed as a core, which could be excited by 980 nm and 808 nm light, respectively. With the optimized geometry structure, the upconversion emission intensity was enhanced over 150 folds under the excitation of both 808 nm and 980 nm laser. Through the NaYF\(_4\)/OPC biochip and fluorescence resonant energy transfer (FRET) sensing strategy, the lowest detection limit of 0.01 ng/mL with good linear relationship was obtained ranging from 0.1 ng/mL to 10 ng/mL. The power density of the pumping light was as low as 1.7 W/cm\(^2\). This enhanced upconversion tumor marker biochip sensor provides a promising and powerful tool for facile early diagnosis on cancer markers.

Keywords

Upconversion fluorescence enhancement, double stop band, photonic crystal, tumor marker detection.
1. Introduction

Cancer is one of the main diseases that endanger human life and health. Conventional detection methods, such as magnetic resonance imaging, and computed tomography, are mainly applied in clinical diagnosis with high sensitivity but restricted by complex equipment, high cost and time-consuming.[1-7] It’s important to design convenient and efficient methods for early detection of tumors, especially for home-based self-screening. Note that the occurrence of cancer is closely associated with the abnormal expression of tumor markers, and the precise identification of potential cancer biomarkers in patient samples is a key factor in the early cancer diagnosis and an estimation of the cancer development stage.[8-12]

Recent years, different kinds of innovative tumor marker detection methods have been developed. Among them, biochip-based device for biomarker detection is an important assay not only in clinical diagnostic fields, but also in home-based self-screening, due to the portability, flexibility, and short processing time on samples. In addition, new technological progress in the biomarker analysis occurs with recent innovations on optical detection methods with fluorescence readout.[13, 14] Unlike traditional luminescent materials, such as quantum dots and dyes,[15-18] rare earth doped upconversion nanoparticles (UCNPs) are able to transform NIR light into visible light,[19-26] with less damage on biospecimen, meanwhile, avoids background fluorescence during detection in comparison with conventional fluorescence probes.[27-31] Furthermore, UCNPs can be applied as a suitable energy donor in fluorescence resonant energy transfer (FRET) for the detection on tumor markers, and lots of biochip-based diagnostics of cancer for point-of-care (POC) applications were developed. [13, 14]

The goal of the POC diagnosis is to detect biomarkers with the requisite sensitivity and accuracy to meet the needs of clinical diagnosis and home-based self-screening. However, low fluorescence quantum yield of UCNPs probe limits their clinical applications on biochip devices for tumor marker analysis. Nowadays, enhancing the intensity of upconversion fluorescence is a challenge to expand their biological applications. To regulate upconversion florescence, core-shell structure and
local field effect became the main means to enhance the intensity of upconversion fluorescence. Normally, the fluorescence intensity can be improved[32-36] by preparing passivation shell, which, however, may cause the increasing size of UCNPs. In biological detection, small size NPs with large specific surface area are preferred for the enough active sensing site, and meanwhile, low power density of the excitation light is applied on specimens.[37] both of which limit the emission intensity of UCNPs.[38, 39] To enhance the emission intensity, various methods based on local field effect are developed, including photonic crystal effect[40-42] and surface plasma effect[43-45]. They have been proved as effective ways to enhance fluorescence intensity, especially for small size NPs and with a low excitation power. For upconversion based biochips, local field effect with the advantage on the fluorescence enhancement, might greatly improve the detection performance and satisfy the clinical requirements for patient sample assays.

Since the discovery of the photonic crystal effect, it has received extensive attentions on optical regulation, especially for the intensity enhancement excited by the low-power light. Opal photonic crystal (OPC) with photonic stop bands can prevent or scatter the light that falls into the band gaps.[46] Once photonic stop band matches with the excitation light or emission light wavelength, upconversion emission intensity of UCNPs could increase over dozens of folds,[47] with putting UCNPs on the surface of OPCs. In our previous work, through combining PMMA OPC effect and surface plasma effect, the enhancement factor of upconversion emission of NaYF₄:Yb³⁺,Er³⁺ NPs reached 1000-fold[48]. The photonic crystal stop band coupling with the upconversion excitation field has been proved to greatly increase the upconversion fluorescence intensity. Until now, synchronous coupling the upconversion excitation and emission fields has not been reported yet.

In this work, in order to promote the potential clinical utility of upconversion based biochip devices, a novel bilayer PMMA OPC with double photonic stop band was designed. In this biochip, based on the previous research results, bilayer photonic crystal structure was used to regulate the stop band to match the excitation field and emission field of the upconversion fluorescence, respectively. Because
NaYF$_4$:Yb$^{3+}$,Er$^{3+}$ and core-shell structure NaYF$_4$:Yb$^{3+}$,Er$^{3+}$@NaYF$_4$:Yb$^{3+}$,Nd$^{3+}$ UCNPs were chose for the energy donor, which can be excited by 980 nm and 808 nm light, respectively. The double stop bands were regulated to 980 nm and 808 nm (excitation) and 545 nm (emission) band of UCNPs respectively (Fig. 1). Through the NaYF$_4$/OPC hybrid structure, the upconversion emission intensity was enhanced over 150 folds under excitation of both 808 nm and 980 nm, which was superior to the conventional monolayer OPC. The geometry of the dual photonic crystal was also investigated. Utilizing the optimized structure, an upconversion fluorescence sensor was fabricated to detect a tumor marker, prostate specific antigen (PSA). The detection limit of 0.01 ng/mL with good linear relationship in the range of 0.1~10 ng/mL was obtained under the excitation power density of 1.7 W/cm$^2$. Since the upconversion fluorescence is greatly enhanced, in the same condition, the excitation power density can be lowered with minimal damage on the sample. The remarkable performance of our PSA-detection biochip could lead to further development in clinical diagnostic applications, as well as in home-based self-screening.

Fig. 1 Schematic illustration of detection system. (a) UCNPs/bilayer OPC hybrid fluorescence film. (b) Fluorescence film modified by poly-dopamine film and capture anti-PSA, excited by 980/808 nm laser. (c) After connecting with PSA and detection anti-PSA modified gold nanoparticles, emission light intensity of UCNPs quenches.

2. Experimental

**Preparation of double stop band OPC**: PMMA OPC consisted of PMMA microsphere was prepared through a MMA polymerization process. Clean glass substrate was put into PMMA colloidal suspension (5% solid content) vertically and placed in 35°C for 24h, the PMMA microsphere would be assembled orderly on glass
substrate by surface tension along with the evaporation process. After this, OPC was put into 120°C for 1h to enhance its physical strength. The position of PSB could be controlled by changing the size of PMMA microsphere. In this work, three kinds of OPC with PSB at 545 nm, 808 nm, and 980 nm were prepared. Next, we fixed OPC in beaker vertically and injected deionized water into beaker slowly by micro syringe pump with rate of 10 mL/h, OPC film on glass substrate would be peeled by surface tension. After OPC was peeled, we used another OPC to fish it, and strengthened it in 60°C for 2h.

**Synthesis of NaYF₄:Yb⁺³, Er⁺³ core nanocrystals:** 0.78mmol YCl₃, 0.20mmol YbCl₃ and 0.02mmol ErCl₃ were added into a 100mL flask containing 6 mL oleic acid and 15 ml 1-octadecene. The mixture was stirred at 160°C for 60 min to form the lanthanide–oleate complexes and then cooled down to 40°C. Next, dropwise adding 6 mL of methanol solution containing 4mmol NH₄F and 2mmol NaOH into flask and the resultant solution was gently stirred for 30 min. After fully stirred, the solution was heated to 120°C to remove methanol. Finally, the solution was further heated to 300°C for 90min to obtain NaYF₄:Yb⁺³, Er⁺³ nanoparticles. The whole preparation took place with the nitrogen protection. Then the solution was cooled down to room temperature. The resulting NPs were collected by centrifugation in ethanol at 9,500 rpm for 15min, washed with absolute ethanol–cyclohexane (1: 3 v/v) two times and dispersed in cyclohexane for the following experiments.

**Synthesis of NaYF₄:Yb⁺³, Er⁺³@NaYF₄:Yb⁺³, Nd⁺³ core-shell nanocrystals:** 0.70mmol YCl₃, 0.15mmol YbCl₃ and 0.15mmol NdCl₃ were added into a 100mL flask containing 6 mL oleic acid and 15 ml 1-octadecene. The mixture was stirred at 160°C for 60 min to form the lanthanide–oleate complexes and then cooled down to 40°C. Next, 6 mL cyclohexane solution was added to coat 1mmol NaYF₄:Yb⁺³, Er⁺³ NPs. After fully stirred for 30min, the solution was heated to 90°C to remove methanol cyclohexane and then cooled down to 40°C. Next steps are the same with the last part.

**Preparation of gold nanoparticles:** Gold NPs were prepared through sodium citrate reduction method. Firstly, 50mL HAuCl₄ solution (1mM) was added into flask,
the solution was stirred and heated in oil bath to boiling, then rapidly added sodium citrate solution (39mM). The whole progress required condensation reflux. After a few minutes, the solution turned to wine red. Being heated 15min continually, and then the solution was cooled down to room temperature. Gold NPs prepared through sodium citrate reduction method could be modified with antibody through electrostatic adsorption. Take 1mL gold NPs solution, collected by centrifugation at 12,000 rpm for 15min and re-dispersed in 1mL deionized water. Then add 60µL 0.1 mg/mL detection anti-PSA. After shaking for 1h and storing at 4℃ for 10h, detection anti-PSA modified gold NPs were collected by centrifugation at 12,000 rpm for 15min again.

**Fabrication of PDA/NaYF₄/OPC hybrid film:** Glass and OPC films were vertically dropped into NaYF₄ cyclohexane solution (0.1 mg/mL) and placed at 28℃ for 6h. With the volatilization of cyclohexane, the NaYF₄ UCNPs will self-assemble onto the surface of glass and OPC films. Vertically putting hybrid films into dopamine TRIS solution (0.6 mg/mL, pH=8.0) for 1h and let dopamine self-polymerization at the surface of hybrid films for binding antibody.

**Detection Section:** PSA, capture antigen and detection antigen were purchased from Shanghai Linc-Bio science company(Basic information in Supplementary Material). The capture anti-PSA (0.1 mg/mL) was modified onto the surface of hybrid film. Next, the samples with different concentration of PSA were added onto the surface of hybrid film sensor, after incubation for 10 minutes, detection area was washed adequately by PBS buffer, then BSA blocking buffer and gold NPs modified with detection anti-PSA were orderly added and modified onto the surface of hybrid film sensor. It also needed 10 minutes for incubation in each step. Finally, detection area was washed by PBS buffer again. In next step, the emission spectra of detection area were recorded with fluorescence spectrometer equipped with 980 and 808 nm excitation lasers. For the tumor marker detection with confocal microscope, we just chose one detection area and added excessive capture anti-PSA, and then orderly added PSA sample and detection anti-PSA modified GNPs for multi times. The
amount of capture anti-PSA was calculated to be enough to meet the requirement of six experiments. 0, 0.01, 0.1, 2, 5, 12.5 ng/mL PSA was added to the same detection area and the GNPs with same amount was added after each PSA sample addition. The real time tumor marker detection images in the sensing area and the corresponding spectra were recorded and scanned with confocal microscope.

![Fig. 2](image.png)

**Fig. 2** (a-d) SEM image and transmitted spectra of 980/545 nm and 808/545 nm double stop band OPC. (e-f) TEM image of UCNPs NaYF₄:Yb⁺³, Er³⁺ and NaYF₄:Yb⁺³,Er⁺³@NaYF₄:Yb⁺³,Nd⁺³. (g) SEM image of UCNPs/OPC hybrid film. (h) Transmitted spectra of glass substrate, UCNPs/glass, OPC film, and UCNPs/OPC hybrid film. (i) TEM image of gold nanoparticles. (j) Absorption spectra of gold nanoparticles, they are modified by different amount of detection antibody and then adding NaCl solution. (k) SEM image of PDA/UCNPs/OPC hybrid film. (l) Fourier infrared spectra of PDA/UCNPs/OPC hybrid film. (m) FTIR spectra of poly-dopamine modified fluorescence film.

### 3. Results and discussions

PSA is an important tumor marker of prostatic cancer, and the value of healthy person is 4 ng/mL. Above this value, the risk of tumor increases, and accurate detection of this tumor marker is important for early screening of prostatic cancer.

Figure 1 shows the design of bilayer upconversion sensor and the principle of PSA detection. Through antibody-antigen-antibody combination, UCNP fluorescence donor can be linked with gold nanoparticles (NPs) receptor, and then the concentration of antigen can be quantified through measuring the decreased emission
intensity due to the fluorescence quenching. After modifying the UCNPs on the surface of the photonic crystal, poly-dopamine (PDA) was further modified on the surface of UCNPs film for connecting with capture anti-PSA. Next, gold NPs with absorption peak at 560 nm was prepared, which matched with green emission peak of Er³⁺, and the anti-PSA was further modified on gold NPs by electrostatic adsorption for detection.[49-51] In this system, we detected the concentration of PSA in PBS buffer solution under both 980 nm and 808 nm excitation. In the process, dropping capture anti-PSA, sample, BSA blocking buffer, and gold nanoparticles modified with detection anti-PSA onto fluorescence film in order, then testing the emission spectra of the detection area. BSA blocking buffer plays the function of anti-fouling to prevent the nonspecific adsorption of detection anti-PSA. Both capture anti-PSA and detection anti-PSA can target the PSA to form a sandwich structure that shortens the distance between the UCNPs and the gold NPs. Since the green emission from UCNPs and the plasma absorption of gold NPs overlap, efficient energy transfer can happen. The emission intensity of 545 nm light would reduce as the concentration of PSA increases.

Fig. 2a-d shows the SEM images and transmittance spectra of 980/545 nm and 808/545 nm double stop band OPC, respectively. The bilayer structure of OPC is composed of PMMA spheres with two different sizes. From the SEM images, it can be observed that PMMA spheres are uniform and regular in arrangement, and both of the layers maintain good integrity. In Fig. 2a and b, the photonic crystals with band gaps around 980 and 808 are both in the upper layer, and the photonic crystals with band gaps around 545 are at down layer. It should be mentioned that, through the photonic crystal grafting method mentioned in experiment section, the photonic crystals of different sizes can be geometrically arranged separately, and this method is also applicable for flexible substrate, such as polyethylene glycol terephthalate (PET) substrate.[48] The bilayer photonic crystal films can match both the excitation field and the emission field together, which is important for studying the interaction between the excitation field and the emission field with the photonic band gap, and exploring the optimal fluorescence enhancement mode. From transmittance spectra,
two photonic stop bands matching the excitation light and emission light, can be observed and the position of photonic stop band could be regulated by adjusting the size of PMMA spheres. The black and red lines correspond to the transmittance spectra of a single-layer photonic crystal, and the stop band position corresponds to the size of the PMMA. When the two photonic crystals are combined, transmission spectra (blue line) with double stop bands are obtained, which further suggests the successful preparation of double stop band bilayer photonic crystal structure. It is worth mentioning that, in the 980/545 nm bilayer photonic crystal structure, although the stop band of 545 nm is relatively shallow, it still provides contribution to the fluorescence enhancement.

From the TEM images of UCNPs (Fig. 2e, f), NaYF₄:Yb³⁺,Er³⁺ UCNPs are ~23 nm, and the NaYF₄:Yb³⁺,Er³⁺@NaYF₄:Yb³⁺,Nd³⁺ UCNPs are ~34 nm, both of which have good morphology of hexagonal, uniform sizes and good dispersion. The size of UCNPs can be tuned by changing the reaction temperature from 280°C to 320°C and the size of UCNPs increases with the increasing reaction temperature. The size of NaYF₄:Yb³⁺,Er³⁺@NaYF₄:Yb³⁺,Nd³⁺ UCNPs increased, due to the modification of the shell, and the thickness of the shell was about 5 nm. Both UCNPs can emit strong green light under the excitation of 808 and 980 nm lights. The 980 nm excitation can generate upconversion emission of Er³⁺ by the two-step sensitization of Yb³⁺ ions. The emission process under 808 nm excitation is more complicated, in which the Nd ions in the shell absorbs the 808 nm photon and then transfers energy to nearby Yb³⁺. Through the core-shell structure, this energy can be effectively transferred to the Yb³⁺ ions in the core and this migrated energy is finally transferred to the Er ions by two-step energy transfer, as shown in the schematic diagram. (Fig. 3)

The SEM image of the UCNPs/OPC hybrid films (Fig. 2g) shows that uniform distribution of UCNPs in the crevices between PMMA spheres. When the position of stop band matches with excitation wavelength, the excitation field can be amplified, and even with low-power excitation, upconversion fluorescence will be enhanced. Especially when UCNPs are located in the interstices between PMMA spheres, the luminescent intensity of UCNPs in those interstices could be further enhanced. The
PMMA photonic crystal prepared by the self-assembly method is very sensitive to water, and water can damage the photonic crystal structure. In this process, the surface of the photonic crystal is modified with a thin hydrophobic layer, which can protect the photonic crystal structure when detecting aqueous solutions. In order to verify the role of the hydrophobic protective layer, we tested the transmittance spectra of UCNPs/bilayer OPC hybrid film before and after placed in water for 4 hours. Benefit from hydrophobic oleic acid ligands on the surface of UCNPs (Fig. S1), UCNPs/bilayer OPC hybrid film shows good structural integrity after placed in water and the transmittance spectra of bilayer OPC base keeps intact, demonstrating the well water stability of UCNPs/bilayer OPC hybrid film. After modified with poly-dopamine, SEM image and Fourier Transform Infrared Spectroscopy (FTIR) spectra of hybrid film was tested (Fig. 2k, l). Contrast with Fig. 2g, a dense film in SEM image could be observed. In FTIR spectra, the new absorption peak between 3100 nm and 3500 nm is assigned to the N-H bond from poly-dopamine, which can be further used to connect with capture anti-PSA.

When studying the photonic crystal effect on the upconversion fluorescence enhancement, it is necessary to determine that the amount of UCNPs on the photonic crystal and the glass substrate are equal, so it is meaningful to compare the upconversion fluorescence enhancement. In the experimental process, same UCNPs solution were modified on the surface of photonic crystal and glass substrate at the same time, to ensure the comparability of their optical properties. In addition, we tested transmittance spectroscopy of glass substrate, UCNPs/glass, OPC film, and UCNPs/OPC hybrid film as shown in Fig. 2h. In the calculations of enhancement factor, it is necessary to avoid the influence of PSB, so the OPC with PSB of 780 nm is chosen and the transmittance at 980 nm is tested. Through Beer-Lambert law ($E_i=\log(1/T_i), i=1,2$), the values of extinction were calculated ($E_1=1.79$ and $E_2=1.97$). This result indicates that the amount of UCNPs on glass substrate and OPC film are quite close. Thus the influence of the amount of UCNPs on different substrates can be neglected on calculation of the upconversion enhancement factors.

As the energy acceptor, gold NPs in Fig. 2i show uniform size and good
dispersion. The absorption peak locates at 560 nm, which overlaps with the green emission of Er$^{3+}$ (Fig. S2a). To confirm the appropriate concentration of gold NPs, the concentration of prepared gold NPs was adjusted to be 0.002, 0.02, 0.2, 0.6, 1.0 mg/mL, then 4uL PBS and five kinds of adjusted gold NPs were respectively added onto fluorescence film and the emission spectrum of every detection area was tested. In our detection system, when the concentration of gold NPs progress was between 0.02 mg/mL to 0.2 mg/mL, most of emission light was quenched and the FRET progress reached saturation point (Fig. S2b, c). So gold NPs with concentration of 0.2 mg/mL was chose to process subsequent experiment. To demonstrate that the detection anti-PSA was connected with gold NPs, six controlled trials were conducted. Firstly, 0, 40µL, 60µL, 80µL, 100µL, and 120µL 0.1 mg/mL detection anti-PSA were added into 1mL prepared gold NPs solution, respectively. After uniformly oscillating for 1h, gold NPs were centrifuged and re-dispersed in 1mL deionized water. Finally adding 100µL 1 mg/mL NaCl solution in each sample and test their absorption spectra (Fig. 2j). After mixed with NaCl solution, the absorption spectra of pure gold NPs solution were broadened obviously due to the agglomeration of gold nanoparticles. More antibodies were connected with gold NPs as their concentration increased, which further weakened the agglomeration of gold NPs, so that the absorption spectra broaden disappeared gradually. This absorption trend indicates that the antibody was successfully modified on the surface of gold NPs.
In our previous work,[44, 48, 52] remarkable enhancement on the intensity of upconversion fluorescence was realized in the coupling of the photonic crystal stop band with the upconversion excitation field. Upconversion fluorescence is a multiphoton process, therefore the fluorescence efficiency is relatively low. In biological detection, it is important to increase the fluorescence intensity. In this work, we use the local field effect of PMMA OPC to increase the upconversion fluorescence intensity. The band gap of the photonic crystal structure can be adjusted by the size of the PMMA sphere. When the photonic stop band is consistent with the wavelength of the incident light, the light will scatter many times on the surface of OPC, especially in the crevices between PMMA spheres. Since UCNPs are placed in the crevices of photonic crystal microspheres, when the excitation light intensity on the surface of OPC is greatly enhanced, then the up-conversion fluorescence is greatly enhanced. According to this principle, the emission light of UCNPs can also be enhanced, as
long as the position of the upconversion emission light and the photonic crystal stop band overlaps. Therefore, we have designed double-layer photonic crystal structure, so that their stop bands can be respectively coupled with the excitation light and emission light of UCNPs, causing that the excitation field and emission field of UCNPs can be simultaneously enhanced, and the intensity of up-converted fluorescence can be further improved. We found that when the photonic stop band is coupled with the excitation field, the better fluorescence enhancement can be obtained, so the upper photonic crystal stop band is set at 980 and 808 nm, and the down layer is at 545 nm coupling with emission field, to obtain the best upconversion fluorescence.

As for the important factors related to fluorescence enhancement, it should be the position of the photonic band gap and the power of the excitation light. The position of the photonic band gap is controlled by the size of the PMMA microsphere. Once the optical band gap position mismatches with excitation light or emission light wavelength, the enhancement factor will decrease rapidly. Regarding excitation power and enhancement factor, we found that when the lower excitation power density is employed, greater enhancement factor can be obtained. In addition, the size of fluorescence nanoparticles also influences the enhancement factor. Bigger size of nanoparticles results in lower enhancement factor. Although clear quantitative relation is hard to get, we can build a more appropriate enhancement system for detection according to the variation trend. Normally, low power density of excitation light is meaningful and necessary for biological detection, which is needed to avoid damage to biological samples. Interestingly, our test system can obtain higher fluorescence enhancement when low power is used, showing that this system is suitable for biological detection which can protect biological samples and improve detection accuracy. In this work, we further introduced a bilayer photonic crystal and synchronously coupled the excitation and emission fields to enhance upconversion fluorescence. The work of synchronizing to couple the upconversion excitation and emission fields has not been reported yet. The difficulty lies in not only the fabrication, but also the spatial configuration and regularity, which are still unclear. The photonic crystal effect has a certain working distance, and UCNPs can obtain the maximum
enhancement on the surface of the photonic crystal. With this bilayer photonic crystal, it is necessary to study the spatial order structure. In this work, as mentioned above, different spatial order of photonic crystals is designed to optimize the enhancement of upconversion emission. Fig. 3a-d shows emission spectra of two kinds of UCNPs excited by 980 nm and 808 nm laser, tested by spectrograph. Emission spectra and enhancement factor of UCNPs films on glass substrate, 545 nm OPC, 980/808 nm OPC, and double stop band OPC was compared. From spectra, compared with glass substrate, 545 nm emission intensity increased 12, 98, 156 folds under 980 nm excitation and 28, 122, 171 folds under 808 nm excitation. Light of corresponding wavelength was prevented by photonic stop band and was scattered on the surface of OPC, especially in the crevices between PMMA spheres. Obviously this effect was more effective to excitation light than emission light, and double stop band OPC showed much stronger enhancement performance than monolayer OPC. To confirm the influence of the spatial sequence of bilayer OPC with PSB of 545 nm and 980/808 nm, controlled trial was conducted. From emission spectra, the trend of fluorescence enhancement of 980 nm part and 808 nm part is the same. When the 545 nm OPC located on the upper side and 980/808 nm OPC on the down side, the enhancement factor was the same level with single 980/808 nm OPC. When the 980/808 nm OPC was on the upper side and the 545 nm OPC was on the down side, the enhancement factor was much higher than another sample. To get available light signal on glass substrate, these spectra were tested under a relatively high excitation light of 5 W/cm².
Fig. 4 (a-e) upconversion emission spectra of detection area, linear fitting and fluorescence lifetime result under 980 nm and 808 nm excitation in antigen detection. (f) specificity test result of PSA and other common substance in the blood.

Fig. 4 shows the PSA detection results tested by spectrograph. We chose 545 nm emission peak of Er\(^{3+}\) as detection position, and the emission intensity decreases as the increase of the concentration of PSA in sample (Fig. 4a, c). The correlation between emission intensity and PSA concentration was also calculated (Fig. 4b, d), which showed satisfactory linear relationship within the range of 0.1~10 ng/mL and
the limit of detection (LOD) is 0.01 ng/mL. When sample concentration is 20ng/mL, signal response showed a saturation trend and deviated from the fitting curve. Note that the PSA content of normal people is below 4ng/mL, and if PSA concentration is above 10ng/mL, risk of prostate cancer will significantly increase, our detection system meets the requirement of clinical detection well. Fig. S3 shows the detection result in PBS and blood sample. To test blood sample with low PSA concentration, blood sample with PSA concentration of about 2ng/mL was diluted to 1/10 of its original concentration and blood sample with PSA concentration of 0.2, 2.2, 5.2, 10.2, 20.2ng/mL was prepared, respectively. Compared with PBS solution, detection result of blood sample also shows very good linear relationship with a little slop change in the range of 0~10ng/mL, indicating that the detection system is promising for PSA detection in blood sample. When sample concentration is 20ng/mL, signal response showed a saturation trend and deviated from the fitting curve, which was consistent with previous result (Fig. 4b, d). The signal response of blood sample was slightly higher than that in PBS solution, which can be attributed to that, in our detection system, antigen-antibody reaction in blood sample might be impeded by viscosity of blood. Compared with PBS solution sample at the same concentration, less PSA in blood sample can be attached with capture antibody due to the viscosity of blood, so that less fluorescence quenching occurred in detection, which resulted in the difference in signal response. These results were tested under excitation light of 1.7 W/cm². It should be mentioned that, the same amount of GNPs were added to each test, while excessive GNPs did not affect the results of the assay. The upconversion fluorescence intensity is only related to the amount of PSA, indicating that this sandwich structure is effective for narrowing the distance between the donor and the acceptor, and that other kind of contacts do not lead to energy transfer. Fig. 4e shows the fluorescence lifetime in each detection area. The lifetime decreases as the increase of the concentration of PSA in sample. However, compared with emission intensity, the descend range of lifetime is much smaller. The reason may be that under present testing condition, emission spectra could be tested in a very low excitation power, so only surface layer UCNPs were excited and quenched by gold nanoparticles. However,
the pulsed laser required in lifetime testing was more powerful, and when more and
deep UCNPs were excited, the change of statistical lifetime was not obvious since
the deeper UCNPs were out of the range of FRET. Fig. 4f shows the result of
specificity test, and the concentration of each control group is the same of 20 ng/mL.
From the result, PSA quenches fluorescence intensity significantly, indicating good
selectivity of this detection system.

This biochip for the detection of tumor marker is promising to be applied for
home self-screening, since the upconversion emission is greatly enhanced, and the
change in fluorescence under a fluorescence microscope can be clearly seen in the
range of 0.1~20 ng/mL.

**Fig. 5** (a) schematic diagram of detection process tested with confocal microscope. (b-c) emission
spectra and corresponding photograph tested by confocal microscope under 980 nm excitation
(Concentration of PSA in sample is 0, 0.01, 0.1, 2, 5, 12.5 ng/mL, respectively).

Fig. 5 shows the detection result tested by confocal microscope with 980 nm
excitation. We coupled a 980 nm laser to a confocal microscope, and the upconversion
emission can be observed through a microscope. In addition, the spectra can be tested
through a microscope. Fig. 5a is a schematic diagram of the overall sensing system. In
this sensing system, the results also show remarkable sensing performance with the
detection limit of 0.01 ng/mL, in the range of 0.1~12.5 ng/mL. We made a
comparison between our detection method and the reported methods within Table 1, which indicates the remarkable detection performance of this tumor marker sensor among the various methods. Fig. 5(b-c) shows the emission spectra and corresponding photograph tested by confocal microscope with 980 nm excitation. In the green fluorescent region of the photonic crystal (Fig. 5c), when the PSA is added, the brightness of the green fluorescent region is gradually decreased. Note that the color in the figure is pseudo color and visual image of the detection can be realized through the gray value. Since excessive GNPs made no difference to fluorescence quenching, in this section, the same area on the biochip was selected for the multi-detection of PSA. Enough amount of capture anti-PSA was added firstly, the amount of capture anti-PSA was calculated to be enough to meet the requirement of six experiments. After PSA and detection anti-PSA modified GNPs was subsequently added, through this sandwich detection method, PSA with different concentrations can be tested multiple times.

<table>
<thead>
<tr>
<th>Detection structure</th>
<th>Sensor principle</th>
<th>Linear range</th>
<th>LOD</th>
<th>Ref.</th>
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</thead>
<tbody>
<tr>
<td>MWCNT</td>
<td>ECL</td>
<td>2.25~11.3ng/mL</td>
<td>0.88ng/mL</td>
<td>[53]</td>
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<tr>
<td>Nanomagnetic beads</td>
<td>Fluorescent</td>
<td>0.05~100ng/mL</td>
<td>0.05ng/mL</td>
<td>[54]</td>
</tr>
<tr>
<td>CdTe QDs</td>
<td>Fluorescent</td>
<td>1~40ng/mL</td>
<td>0.4ng/mL</td>
<td>[55]</td>
</tr>
<tr>
<td>Gold NPs</td>
<td>SERS</td>
<td>4~10ng/mL</td>
<td>0.012ng/mL</td>
<td>[56]</td>
</tr>
<tr>
<td>MOFs</td>
<td>Electrochemical</td>
<td>0.1~100ng/mL</td>
<td>0.06ng/mL</td>
<td>[57]</td>
</tr>
<tr>
<td>Mesoporous silica</td>
<td>Electrochemical</td>
<td>1~300ng/mL</td>
<td>0.28ng/mL</td>
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</tr>
<tr>
<td>/gold film</td>
<td></td>
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<tr>
<td>gold film</td>
<td>SPR</td>
<td>0.1~50ng/mL</td>
<td>0.091ng/mL</td>
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<td>Gold nanodisk</td>
<td>LSPR</td>
<td>1.7~20.4ng/mL</td>
<td>1.49ng/mL</td>
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</tr>
<tr>
<td>OPC/UCNPs</td>
<td>FRET</td>
<td>0.1~10ng/mL</td>
<td>0.01ng/mL</td>
<td>Present work</td>
</tr>
</tbody>
</table>

4. Conclusion

In summary, a novel bilayer structure OPC is designed which simultaneously matches with the excitation wavelength and emission wavelength of the UCNPs. According to the emission spectra of UCNPs on different substrates which are prepared in the same condition, this novel structure shows significantly higher
upconversion enhancement factor than the conventional single OPC. With low excitation power density and enhanced emission intensity, the upconversion based PSA sensor was fabricated using NaYF₄:Yb³⁺, Er³⁺ UCNPs and NaYF₄:Yb³⁺, Er³⁺@NaYF₄:Yb³⁺ UCNPs, respectively. Under 980 nm and 808 nm laser excitation, this detection shows a good linear relationship within the range of 0.1 ng/mL~10 ng/mL and LOD of 0.01 ng/mL. Since the upconversion fluorescence was greatly enhanced, this system could excited by the NIR laser with low power density of 1.7 W/cm². This enhanced upconversion based strategy can be extended to design a universal biomaterial sensor, and applied in the areas of health assessment and disease diagnosis.

**Acknowledgements**

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