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Photobleaching on Photonic Crystal Enhanced Fluorescence Surfaces

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Abstract The effect of resonant fluorescent enhancement from a photonic crystal surface upon the fluorescent photobleaching rate of Cyanine-5 labeled protein has been investigated. We show that the enhanced excitation mechanism for photonic crystal enhanced fluorescence, in which the device surface resonantly couples light from an excitation laser, accelerates photobleaching in proportion to the coupling efficiency of the laser to the photonic crystal. We also show that the enhanced extraction mechanism, in which the photonic crystal directs emitted photons approximately normal to the surface, does not play a role in the rate of photobleaching. We show that the photobleaching rate of dye molecules on the photonic crystal surface is accelerated by 30x compared to an ordinary glass surface, but substantial signal gain is still evident, even after extended periods of continuous illumination at the resonant condition.

Keywords Photonic crystal · Enhanced fluorescence · Photobleaching

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Introduction

The development of nanostructured surfaces for enhancing the electric field exposure of adsorbed fluorescent dve molecules has been a topic of intense research interest due to the potential for such surfaces to increase the detection sensitivity of surface-based fluorescent assays by over two orders of magnitude [1, 2]. Enhanced fluorescence (EF) can be applied to a broad range of assays, including gene expression microarrays [3], protein biomarker detection [4, 5], next-generation DNA sequencing [6], single molecule biophysics [7], and cell microscopy. One route towards EF has been the use of nanostructures fabricated from gold or silver, which are capable of coupling external light sources through surface plasmons to generate localized regions with electric fields exceeding the intensity of the illumination source by $10-15 \times [8-10]$. EF using metal nanostructures is associated with relatively low resonant quality factors (Q~ 15), due to optical loss in the metal material, and the possibility for fluorescent quenching when dye molecules are <10 nm from the metal surface [10].

Recently, surfaces comprised entirely of dielectric materials have been demonstrated as an alternative approach for EF. Specifically, photonic crystal (PC) surfaces that are comprised of a periodically modulated grating structure that is coated over with a thin film of high refractive index material have been demonstrated to produce high quality (Q~1000) resonances and a lack of quenching effects [11]. Through selection of the grating period, a PC surface may be designed to function as an optical resonator at any desired wavelength, from ultraviolet to infrared [12, 13], and thus can be designed to interact with any fluorescent dye molecule or quantum dot [1, 2]. PC surfaces with an optical resonance designed to coincide with the wavelength of an excitation laser will

generate an enhanced electric field that is strongly confined to the PC surface and the media (air or water) immediately adjacent to the PC surface. PC "enhanced excitation" results in exposure of surface-adsorbed fluorophores to a greater electric field strength than they would otherwise experience upon an ordinary glass surface, and can be considered to be equivalent to increasing the intensity of the illumination source, but only in a region within ~100 nm of the device surface. PC surfaces may also be designed to simultaneously exhibit an optical resonance that coincides with the emission wavelength of a fluorophore. In this case, called PC "enhanced extraction," emitted photons, which would ordinarily exit the surface distributed uniformly in all directions, are spatially biased away from the PC surface at an (approximately) normal angle, so they may be gathered more efficiently by imaging objectives and/or sensors [2]. It has been shown that the processes for PC enhanced excitation and enhanced extraction operate independently, and that their overall effects multiply. Importantly, PC enhanced fluorescence (PCEF) surfaces may be produced inexpensively over large surface areas by nanoreplica molding on plastic substrates or by nanoimprint lithography on guartz substrates [14-16] so that a PC-active surface may cover an entire 1×3 in² slide or the bottom surface of an entire microplate.

While PCEF has been applied to gene expression microarrays and protein biomarker detection [5], the relationship between fluorescence enhancement and the rate of photobleaching has not yet been characterized. Although photobleaching may not be of critical importance for single-measurement biomolecular assays, knowledge of photobleaching effects on PC surfaces will become important as researchers attempt to apply PCs to cell-based assays and single-molecule measurements of molecular machines using FRET (Forster Resonance Energy Transfer) probes [17]. The relationship between photobleaching and fluorescent enhancement factor is not completely straightforward for PCEF because only enhanced excitation is expected to participate in photobleaching, while the enhanced extraction effect is only redirecting emitted photons for more efficient collection. Further, the angle of incidence of a single wavelength collimated illumination source is extremely important in determining the degree of enhanced excitation of the resonant modes of the PC [18]. Therefore, our goal is to characterize the rate of photobleaching from a PC surface using a collimated illumination source and to study its dependence on the angle of illumination (and thus the extent of enhanced excitation). We study PCEF using a fluorescent microscope that incorporates a collimated monochromatic light source that can be precisely tuned to the resonant illumination angle to excite the PC resonance. With this instrument, we are able to match the illumination conditions (of incident wavelength and incident angle) to satisfy the resonant condition of the PC, so the illumination is called "on-resonance." By adjusting the incident angle for a fixed wavelength so that the resonant coupling condition of the PC is not satisfied, the illumination may also be supplied in an "off-resonance" manner. We observe that the rate of photobleaching is substantially reduced when off-resonance illumination is used, and that the photobleaching rate is strongly correlated with the coupling efficiency between the PC and the illumination source, indicating that indeed only the enhanced excitation mechanism participates in modulating the photobleaching rate.

Photonic Crystal Enhanced Fluorescence

The confinement of evanescent fields to a small volume region adjacent to a surface has been previously exploited in Total Internal Reflection Fluorescence (TIRF) microscopy to reduce the levels of observable background fluorescence from the sample volume [19]. More recently, it was shown that a corrugated waveguide structure demonstrating "evanescent resonance" could enhance fluorescence excitation of dyes immobilized on the surface of a glass slide [3]. thus providing elevated electric field levels to complement the surface localization of TIRF. Since that time, device structures have evolved to provide enhanced excitation with greater magnitude through resonant surfaces with narrower wavelength bands [20], combination of enhanced excitation with enhanced extraction [1], and detection instruments that more efficiently couple light into and out of the structure [18].

The PCs used in this work are nanostructures comprised of a periodically modulated low refractive index plastic/ SiO₂ surface structure that is coated with a high refractive index dielectric thin film of TiO2. The purpose of the structure is to provide an efficient optical resonator, as described in previous research [1, 2, 5, 14, 21], but summarized briefly here. The periodically modulated dielectric structure of the PC functions as a resonantly reflective optical filter, where only particular wavelength/ incident angle combinations interact strongly with the structure, resulting in highly efficient reflection, while all other wavelength/incident angle combinations are transmitted through. Unlike a conventional diffraction grating, the grating period is smaller than the wavelength of light, so only the zeroth-order reflected and transmitted waves are allowed, resulting in evanescent fields with a profile that extend into the medium in contact with the PC. The resonance phenomenon is easily observed by illuminating the structure with a broad band of wavelengths and measuring the narrow bands of wavelengths that are reflected with high efficiency. The resonant condition is also dependent upon the incident angle of the external illumination, so each incident angle can have one or more distinct resonant wavelengths. This resonance effect has been exploited to design highly efficient narrowband optical filters [22] as well as label-free optical biosensors [16]. PC resonant wavelength/angle combinations are obtained for specific polarizations of light. Transverse magnetic (TM) modes are excited when the electric field vector of illumination is oriented perpendicular to the grating lines, while transverse electric (TE) modes are excited when the electric field vector is oriented parallel to the grating lines.

The resonant modes are highly localized within and in direct proximity to the PC surface, and a large energy density is observed at these locations in the form of enhanced electromagnetic fields. The intensity of a fluorophore's emission is proportional to the electric field intensity (which is proportional to the square of the electric field) of the light exciting the molecule, so an enhanced electric field will enhance the excitation of fluorophores close to the device surface. Enhanced excitation works selectively in regions within close proximity to the PC surface due to the exponential decay of the electric field intensity from the PC into the superstrate region above the PC. Thus, enhanced excitation of the PC shares one of the advantages of TIRF microscopy because fluorophores close the substrate surface are selectively excited [9]. Rather than utilizing a TIRF microscope, a conventional confocal microarray scanner or a modified fluorescence microscope can be utilized for rapid imaging of large areas without a coupling prism. Using PC enhanced excitation, we have demonstrated fluorescence enhancement from the fluorescent dye Cyanine-5 [21] and detailed the spectral characteristics of the PC-fluorophore interaction [21] as well as the dependence on distance from the PC on enhanced excitation [11].

While enhanced excitation can be observed when the laser light incident on the PC is spectrally aligned with the resonance wavelength, another enhancement effect can be observed when a resonant mode of the PC coincides with the fluorophore emission wavelength. Enhanced extraction occurs when light emitted by fluorophores couples into this mode and is re-radiated such that it can be more efficiently gathered by the measurement instrumentation. Essentially, the PC causes some proportion of photons that would normally radiate through the substrate to be reoriented in a direction that is approximately normal to the device surface.

To perform enhanced excitation and enhanced extraction simultaneously, the PCs used in this study are designed with two separate resonances. The first is a TM resonance that coincides with the excitation wavelength of λ = 632.8 nm. The TM resonance is designed for a narrow band of wavelengths, because the electric field enhancement is proportional to the resonant quality factor, defined as Q= $\lambda/\Delta\lambda$ [23]. The second resonance is a TE mode that spectrally overlaps with the emission wavelengths of the fluorophore, where ideally the center wavelength of the TE mode coincides with the wavelength of maximum emission. The combination of these two effects has been previously used to enhance the fluorescence from semiconductor quantum dots [1] with a magnification factor of 8x for the excitation effect and $13 \times$ for the extraction effect, for an overall sensitivity enhancement of $108 \times [1]$ and additional nanostructures have been used to obtain PCEF enhancements as high as $588 \times [20]$.

Experimental

Photonic Crystal Substrate

In order to study photobleaching on substrates with fluorescence enhancement capability, a PC surface with periodic modulation in one direction was fabricated as shown in the cross-sectional diagram (not to scale) of Fig. 1 (a). The surface grating structure was formed in a ultraviolet curable polymer (UVCP) on a polyethyleneterephthalate (PET) substrate, and the polymer grating surface was coated with a 300 nm SiO₂ spacer layer followed by a high refractive index dielectric layer of TiO₂, which functions as a wave confinement layer. In order to achieve a resonance at $\lambda = 632.8$ nm, electromagnetics simulation software (DiffractMOD, RSoft Design Group) based on the rigorous coupled-wave analysis (RCWA) technique was used to design the PC, resulting in a desired grating period of 360 nm and grating depth d=50 nm. Figure 1 (b) shows a top-down SEM image of a device fabricated to these design dimensions

Fabrication of the device was performed using a plasticbased nanoreplica molding process [14]. Briefly, a silicon wafer with a negative surface volume image of the desired grating pattern was fabricated using deep-UV lithography and reactive ion etching. A viscous liquid that contains an uncured monomer and a UV-activated polymerization initiator is sandwiched between a PET sheet and the silicon master wafer to enable the liquid to fill the silicon surface structure subsequent to curing with a high intensity UV lamp (Xenon, Inc). The hardened polymer grating preferentially adheres to the PET substrate, and thus can be easily peeled away from the silicon. After the molding step, the replica was cut and attached to a 1×3 in² microscope slide. An evaporated SiO₂ intermediate layer (t_{SiO2})=300 nm; ebeam evaporation, Denton Inc.) on the grating surface was deposited to reduce autofluorescence from the underlying polymer material due to dielectric sputtering. After the SiO₂ deposition, ~120 nm of TiO₂ was deposited by a RF sputtering system (PVD 75, Kurt Lesker) using an in-situ process monitor to accurately achieve a resonance condition





Fig. 1 Schematic cross section diagram of the photonic crystal structure (a), with a top view electron microscope image (b), and a photograph of the device attached to a standard glass microscope slide (c)

that nominally results in λ =633 nm wavelength resonantly coupling to the PC surface at an incident angle of 2°. Figure 1 (c) shows the final device after all fabrications steps were completed.

J Fluoresc (2011) 21:707-714



Fig. 2 Detection instrument used for photonic crystal enhanced fluorescence photobleaching study. A HeNe laser beam is expanded and collimated before reflection from an angle-tunable mirror that illuminates the photonic crystal device from below. The fluorescence emission is gathered by a conventional microscope objective and measured using a cooled EM-CCD camera

For this study we designed the PC with a TM-polarized resonance close to the Cyanine-5 (Cy5) excitation wavelength of λ =625 nm and a TE-polarized resonance spectrally overlapping the Cy5 emission spectrum centered



Fig. 3 Angular transmission spectrum for the TM enhanced excitation mode showing the choice of various angles of excitation **a** to generate optimal on-resonance excitation at an incident angle of 2.03°. Transmission efficiency versus wavelength **b** showing dips in transmission intensity at normal incidence for the TE (λ =690 nm) and TM (λ =625 nm) modes. Wavelengths of maximum resonance reflection correspond to wavelengths of minimum transmission

Fig. 4 Images of

for photonic crystal on-

crystal off-resonance

for direct comparison



at λ =690 nm. The TM resonance increases the excitation of the fluorophore through enhanced electric field intensities, while the TE resonance redirects a proportion of emitted light toward the detection instrumentation. Figure 3 (b) shows the transmission spectra of the PC measured using white light illumination at normal incidence, in which dips in the transmission spectra indicate a resonance.

Detection Instrumentation

The detection system used in our study is a modified backilluminated fluorescent microscope (Olympus BX51WI) shown schematically in Fig. 2. The microscope is equipped with a $4 \times$ objective (N.A.=0.1) and an electron multiplying CCD (C9100-13 EM-CCD; Hammamatsu Inc.) for imaging. The EM-CCD provides control over the gain settings and integration times used during the imaging process. All results reported here were measured with a sensitivity gain of 161, an analog gain of five and an exposure time of 400 ms. A 35-mW, λ =632.8 nm HeNe laser was chosen as an excitation source that was aligned with the absorption spectrum of Cy5. A high-resolution motorized gimbalmounted mirror and beam-expanded laser provides collimated illumination at a user-selectable incident angle to the PC. In order to maintain a constant illumination area on the device, the gimbal-mounted mirror sits on top of a motorized linear stage and moves as the mirror rotates. As the collimated light at a fixed wavelength is incident on the PC surface, the angle of incidence can be tuned to allow the laser to couple with the PC resonance, thereby allowing maximum field coupling into the TM mode of the PC. The excitation illumination was TM polarized by passing the laser light through a half-wave plate.

Method

A detection experiment using a Cy5-labeled protein was carried out on the PC surface and a reference glass slide in order to directly compare the rates of photobleaching. The PC surface and the glass slide were precleaned with O₂ plasma for 5 min. Following the cleaning, both PC and glass were functionalized by overnight incubation in an enclosed glass container with 5% 3-glycidoxypropyldimethylethoxysilane in dry toluene at 100 °C. After incubation the silanized devices were cleaned by sonication in toluene, methanol and deionized (DI) water and then dried under a nitrogen stream. Cy5 conjugated streptavidin (GE Healthcare) at $10 \mu g/ml$ was spotted onto the slides by a piezo dispenser (Piezorray, Perkin Elmer) to produce 4×4 arrays of labeled protein spots of ~500 µm diameter. After overnight incubation, the devices were washed by gently dipping them in a protein blocking buffer (Phosphate buffered saline at pH 7.4 with Kathon antimicrobial agent) solution for 60 s. followed by DI water rinse.

To perform measurements using the detection system in Fig. 2, we first tuned the excitation laser to the resonant angle without exposing the fluorophores. Next, we opened a shutter and captured 400 image frames in sequence, exposing the sample each time for 400 ms. The shutter was closed and reopened before each measurement so as to avoid unnecessary exposure to light. All measurements were taken on the same sample from identical spots. In order to discern the effect of enhanced excitation on the photobleaching rate, we recorded data from the angle of maximum excitation (on-resonance), minimum excitation



Fig. 5 Plot of the measured fluorescence intensity versus time for continuous exposure at different angles of incidence for the excitation light

Table 1 Photobleaching decayrates and signal intensity gain aspertaining to excitation angle

Device	Angle of excitation	Photobleaching decay rate 'A'	Signal intensity gain 'B'
PC	2.03°	0.0030	346.71
PC	2.23°	0.0020	279.65
PC	2.46°	0.0010	223.40
PC	20.00°	0.0001	9.13
Glass	20.00°	0.0001	1

(off-resonance) and several intermediate angles. The transmission efficiency of the selected points is shown in Fig. 3 (a). We selected angles of 20° , 2.46° , 2.23° and 2.03° where the transmission efficiency of the PC is 100%, 76%, 52% and 26% (minimum transmission) respectively.

Results

Figure 4 shows initial (first exposure) fluorescent images of Streptavidin-Cy5 spots on the PC at 2.03° (on-resonance), 20° (off-resonance) and on the glass slide illuminated at 20° . The spots shown in Fig. 4 clearly indicate enhanced excitation and enhanced extraction effects that also become evident when we analyze the numerical data.

Figure 5 shows the fluorescent intensities collected for each angle over a period of 159 s. Each curve was

measured at a different angle of excitation (and same excitation wavelength of λ =632.8 nm), starting with a fresh, unexposed array of spots. The output intensity on a set of 16 spots was recorded and averaged for each frame. The background value for each frame was then subtracted to give the final averaged spot intensity. The final data are plotted on the same scale for comparison. The highest raw signal value is for the angle pertaining to the lowest transmission efficiency, corresponding to maximum resonant coupling of the excitation laser with the PC surface. As expected the curve corresponding to the 52% transmission (2.23°) has the next highest raw signal value followed by the curve corresponding to 76% transmission (2.46°). Note that even when the PC is illuminated at an angle that is far from the resonant condition (20°), we still obtain greater signal output compared to a glass surface. This high signal output is due to the enhanced extraction effect.



Fig. 6 Electric field intensity plots generated using rigorous coupled wave analysis for excitation conditions corresponding to 2.03° (a), 2.23° (b), 2.46° (c) and 20° (d). The electric field in the medium

directly adjacent to the photonic crystal surface is modeled to be more than two orders of magnitude higher in the on-resonance (2.03°) condition, compared to the off-resonance (20°) condition

In order to compare the rates of photobleaching on the PC, we normalized each data set to our unpatterned glass control and fitted each resulting curve to an exponential function. The fitting equation used was as follows:

Signal_Intensity =
$$B^*e^{-A^*t}$$
 (1)

where A and B are fitting parameters indicative of the photobleaching decay rate and signal intensity gain.

Table 1 lists the values of parameters A and B for different excitation angles. We observed that as we moved from the on-resonance case to the off-resonance case, there was a steady decrease in signal intensity gain and the photobleaching decay rate. Thus for higher degrees of enhancement we see a greater rate of photobleaching. This trend can be attributed to the strength of the electric fields present close to the surface at various angles.

In order to further investigate the effect more fully, we analyzed spatial plots of electric field intensity (close to the surface of the PC) obtained by rigorous coupled wave analysis (RCWA) electromagnetic field computer simulation (Rsoft, DiffractMod). Figure 6 shows cross section plots of the electric field at the four different angles of excitation. We observe that the field intensity is also directly related to the degree of coupling with the resonant mode (highest field for the on-resonance case). The increase in the electric field intensity helps explain the reason for a higher enhancement and subsequent faster rate of photobleaching. For a higher electric field intensity, a greater amount of energy will be delivered to the fluorophores located near the surface thus producing a higher fluorescence signal output.

It is important to note here that even though the rate of photobleaching is $30 \times$ higher for the PC on resonance when compared to unpatterened glass (as indicated in Table 1), after hundreds of individual exposures, a $186 \times$ enhancement factor is still observed relative to measuring the same fluorophore on a glass surface after the final scan, compared to a $346 \times$ enhancement factor for the initial scan. In fact, in order to extrapolate to mathematically determine how long of an exposure would be required to eliminate the benefits of the PC, we can equate the respective fitting equations (Eq. 1) for the PC on resonance and glass cases:

$$B_{2.03^{\circ}} * e^{-A_{2.03^{\circ}} * t} = B_{glass} * e^{-A_{glass} * t}$$
(2)

Inputting the respective values for constants A and B from Table 1 in the cases for the PC on resonance and the glass, we find that for the fluorescence signal for the PC on resonance to equal the fluorescence signal on the glass slide, both substrates would have to be subjected to constant exposure from the excitation laser for over 2,000 s. Thus for all practical experimental time frames,

the fluorescence signal for the PC on-resonance will be higher than the fluorescence signal from an unpatterened glass substrate.

An interesting point to highlight here is that even though the signal intensity for the off-resonance case is $9 \times$ higher than the case for glass, the rate of photobleaching remains unchanged. This is a direct consequence of the extraction effect of a PC surface. The ability of the PC to allow emitted light to couple to a resonant mode and be directed towards the collection lens allows for a high degree of enhancement of the signal without adversely affecting the rate of photobleaching.

Conclusion

In this work, we have correlated the rate of fluorescent photobleaching in PC with the level of resonant fluorescent enhancement. Accelerated fluorescent photobleaching rates in a resonantly excited PC are a direct consequence of the enhancing of the surface localized electric fields exposed to adsorbed fluorescent dye molecules. This mechanism of enhanced excitation for PCEF, accelerates photobleaching in proportion to the coupling efficiency of the laser to the photonic crystal. The enhanced extraction mechanism of the PC, however, provides no contribution to the rate of photobleaching but still has significant contribution to the net signal enhancement.

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