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Country	Germany - IIII SIR Ranking of Germany	45	
Subject Area and Category	Biochemistry, Genetics and Molecular Biology Biochemistry Biophysics Biotechnology Materials Science Nanoscience and Nanotechnology	Hindex	
Publisher	Springer Verlag		
Publication type	Journals		
ISSN	15571955		
Coverage	2006-ongoing		
Scope	Plasmonics publishes peer-reviewed articles that both advance and report on the knowledge and practice of the interactions of free-metal electrons, Plasmons. Coverage includes theory, physics, and applications of surface plasmons in metals, and rapidly emerging areas of nanotechnology, biophotonics, sensing, biochemistry and medicine. The journal covers the theory, synthesis and optical properties of noble metal nanostructures, patterned surfaces, continuous or grated surfaces and devices. Applications include surface-enhanced spectroscopic properties, such as Raman scattering or fluorescence, as well developments in techniques such as surface plasmon resonance and near-field scanning optical microscopy.		
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Absorption Enhancement in Peridinin–Chlorophyll–Protein Light-Harvesting Complexes Coupled to Semicontinuous Silver Film

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Received: 5 July 2011 / Accepted: 17 August 2011 / Published online: 24 August 2011 © The Author(s) 2011. This article is published with open access at Springerlink.com

Abstract We report on experimental and theoretical studies of plasmon-induced effects in a hybrid nanostructure composed of light-harvesting complexes and metallic nanoparticles in the form of semicontinuous silver film. The results of continuous-wave and time-resolved spectroscopy indicate that absorption of the light-harvesting complexes is strongly enhanced upon coupling with the metallic film spaced by 25 nm of a dielectric silica layer. This conclusion is corroborated by modeling, which confirms the morphology of the silver island film.

Keywords Light-harvesting complex · Chlorophyll · Absorption enhancement · Plasmon excitation · Fluorescence · Time-resolved fluorescence · Artificial light harvesting · Hybrid nanostructure

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Introduction

Plasmon excitations in metallic nanoparticles provide an excellent way to control the optical properties of matter at the nanoscale [1]. Hybrid nanostructures comprising metallic nanoparticles have been studied in many fields such as optoelectronics [2], biosensors [3], cancer therapy [4], or photovoltaics [5]. One of the prerequisites for controlling the properties at the nanoscale using plasmonic excitations is the ability to fabricate an on-demand hybrid plasmonic nanostructure with well-defined characteristics. As the strength of the interaction between plasmons and other nanostructures depends upon the spectral properties of both components as well as the geometry of a hybrid nanostructure, it is required to research the fundamental interactions between plasmons localized in metallic nanoparticles with optically active molecules. Placing a fluorescent molecule in the vicinity of a metallic nanoparticle may either quench or enhance the fluorescence [6] depending on the interplay between the plasmon-induced increase of both fluorescence quantum yield and the absorption rate and non-radiative energy transfer from the molecule to the nanoparticle. Typically, the latter process becomes dominant for distances below 5 nm: the emission is efficiently quenched. On the other hand, the largest enhancements of the fluorescence have been observed for distances between 10 and 30 nm between the metallic nanoparticle and the fluorescent molecule [7].

While spectroscopy studies of hybrid plasmonic nanostructures comprising only one type of fluorescentmolecule are currently in an advanced stage, extending both the techniques and understanding toward more complicated systems such as multi-pigment biomolecules still remains a challenge. A major difficulty concerns the interactions between the pigments in multi-chromophoric systems, for instance fluorescence resonant energy transfer, and the preparation of biological systems in a state that is suitable for spectroscopic studies of plasmoninduced effects.

Recently, several reports have focused on the effect of local electromagnetic field generated in metallic nanoparticles on the optical properties of chlorophyll-containing biological light-harvesting complexes [8–12]. When peridinin–chlorophyll–protein (PCP) and Photosystem I (PSI) were coupled to a highly inhomogeneous ensemble of metallic nanoparticles, the fluorescence [8, 9] and/or absorption [10] can be increased. The influence of plasmonic interaction on the protein function has also been studied for the latter PSI light-harvesting system [11] and the fluorescence enhancement for allophycocyanin deposited on silver island film has been observed [12].

In this work, we attempt to optimize the geometry of a plasmonic hybrid nanostructure such that the electromagnetic field generated in metallic nanoparticles leads exclusively to the enhancement of absorption of the lightharvesting complex. The structure comprises a layer of semicontinuous silver film with a relatively narrow size distribution that is separated from the light-harvesting complexes by a 25-nm-thick SiO₂ layer. Large energy separation between the absorption and fluorescence of the light-harvesting complex, PCP, combined with a properly designed metallic nanostructure, with well-defined plasmon resonance, offers the unique opportunity to separate the influence of plasmonic excitation on the excitation rate from that on the emission. The fluorescence excitation spectra show a very strong absorption band in the spectral range corresponding to the plasmon resonance of the silver islands, which could be modeled by the electromagnetic field enhancement generated in such a hybrid nanostructure. On the other hand, fluorescence decay times change only slightly upon placing the light-harvesting complexes on the silver island film. We conclude that the design of our structure provides an efficient way to plasmonically enhance the absorption of light-harvesting complexes.

Samples and Experiment

Light-harvesting complexes, or antennas, enhance and broaden the absorption of photosystems for the efficient use of sunlight in photosynthesis. PCP is one of many such complexes. It is a water-soluble biomolecule employed by *Dinoflagellates* as an antenna external to the membrane [13]. The structure of the PCP complex has been determined with 1.3-A resolution using X-ray crystallography. The native form of PCP consists of two chlorophyll *a* (Chl) and eight peridinin (Per) molecules [13]. All pigments are embedded in a protein matrix. As the ratio of Per to Chl is >>1, in contrast to most of other light-harvesting complexes, PCP utilizes the carotenoid, Per, and not the Chls as its main light-harvesting pigment. The PCP complex can be reconstituted with other Chl dervatives which, for example, exhibit different optical properties [14, 15]. One example is [3-acetyl]-chlorophyll *a* (acChl). Chemically, it differs from Chl only by the C-3 substituent, but the absorption and fluorescence spectra of acChl–PCP, the PCP complex reconstituted with acChl, is red-shifted as compared with the PCP complex containing Chl. At the same time, the Per absorption in the blue-green spectral range is affected very slightly [14]. The advantage of using a red-shifted pigment stems from the better spectral separation between the emission of the pigment and the Per absorption to which the plasmon resonance frequency of the metallic nanoparticles used to fabricate the plasmonic hybrid nanostructure is matched.

The peridinin–chlorophyll–protein complexes used in our experiments were prepared as described [16]. In order to obtain PCP complexes containing Chl, the N-terminal apoprotein was reconstituted with an appropriate mixture of Per and, as desired, Chl or acChl. The reconstituted monomeric PCP was equilibrated with TRIS buffer and kept frozen until used. The absorption (and fluorescence excitation) spectrum of the acChl–PCP has an intense, broad band between 400 and 550 nm that is due to Per and two Chl bands at 440 nm (Soret) and 690 nm (Q_Y), the emission spectrum peaks at 690 nm (Fig. 1a) [16].

The semicontinuous silver films were deposited onto a glass coverslip utilizing the e-beam evaporation technique [17]. The thickness of the silver layer was nominally 7 nm. In order to protect the metallic layer and to diminish the impact of the non-radiative energy transfer from the PCP complexes to the silver islands, a 25-nm-thick layer of silica was deposited using e-beam evaporation [17, 18]. The e-beam evaporation method allows for a much more homogeneous distribution of the size of silver islands compared, in particular, with the methods used previously [8, 9, 19]. The absorption spectrum of such a silver island film (Fig. 1b) shows the plasmon resonance around 450 nm that matches very well the Per absorption of the PCP complex (Fig. 1a). Previous SEM studies of the identically prepared semicontinuous silver film [17] have demonstrated the relatively good homogeneity of the plasmonic structure.

A 2-µl drop of the acChl–PCP solution was deposited directly onto the metallic substrate and then slowly dried at room temperature in darkness to inhibit the degradation of the light-harvesting complexes. In comparison with previous experiments [8, 9], where a polymer layer was used, the present approach assures that most of the PCP complexes are at the SiO₂ surface; thus, their distance between the PCP complexes and metallic nanoparticles is very uniform and defined by the thickness of the SiO₂ layer. For the reference, a similar sample was prepared with PCP complexes deposited directly on a clean glass coverslip.



Fig. 1 a Absorption (*black*) and fluorescence spectra (*red*, λ_{exc} = 532 nm) of acChl–PCP complexes. *Arrows* mark the wavelengths used for time-resolved experiment. b Absorption spectrum of the semicontinuous silver film (*red*) compared to the absorption spectrum of the glass substrate (*black*)

The fluorescence and fluorescence excitation spectra of both structures were measured using the FluoroLog 3 spectrofluorimeter (Jobin Yvon) equipped with a specially designed mount suitable for holding planar samples. A xenon lamp source with a double-grating monochromator was used for excitation, and the signal was detected with a thermoelectrically cooled photomultiplier tube characterized by a dark current of <100 cps. The experiments were carried out in a backscattering geometry. The decay times of fluorescence were measured using time-correlated singlephoton counting. The experimental setup was based on the SPC-150 card (Becker & Hickl). For the excitation, 3-ps lasers were used with the wavelengths of 405, 485, and 640 nm, each operating at an 80-MHz repetition rate; the average excitation power was about 2 mW. The detection was carried out using an idQ-100-50-ULN silicon avalanche photodiode detector (idQuantique). The time resolution of the setup was around 250 ps. The fluorescence of the acChl-PCP complexes was extracted using a band-pass filter HQ690/20 (Chroma).

The three excitation wavelengths (marked by arrows in Fig. 1a) provide a means to study the behavior of Chl fluorescence for qualitatively different conditions of plasmon coupling. Excitation at 405 nm corresponds mainly to the Soret band of acChl absorption, excitation at 485 nm

corresponds to the Per absorption, and excitation at 640 nm corresponds to $Q_{\rm Y}$ absorption of the acChls. In the latter, Per molecules are not excited. Moreover, whereas the two first lasers efficiently excite plasmon resonance in the silver island film, the wavelength of 640 nm overlaps very weakly with the plasmon spectrum of the silver nanoparticles.

Results and Discussion

The effect of plasmon excitations in metallic nanoparticles on the optical properties of nearby emitters is commonly monitored by its fluorescence intensity. In the case of plasmon-induced enhancement, the fluorescence intensity of such a hybrid structure is increased [6]. When, on the other hand, non-radiative energy transfer from the emitter to the metallic nanoparticles plays the dominant role, the emission is efficiently quenched. This gives only incomplete information, however, because the fluorescence depends upon the spectral properties of the emitter and their relation to the plasmon resonances in the metallic nanoparticles. Additional information about plasmonic interactions can be extracted by combining fluorescence excitation spectroscopy [20] and time-resolved fluorescence spectroscopy; these two experimental techniques provide a way to separate the plasmon-induced increase of the radiative rate from an induced increase of the absorption. The acChl-PCP complex is well suited for such a study because of its large Stokes shift when the carotenoids are excited (Fig. 1).

In Fig. 2 we compare the fluorescence excitation spectra of acChl -PCP deposited on the semicontinuous silver film and on the glass substrate, also shown is the emission spectrum (λ_{exc} =532 nm). The excitation spectrum for the reference sample looks very similar to previously published data [14]; it features a strong absorption due to Per in the spectral range from 400 to 550 nm and corresponds roughly to the absorption spectrum (Fig. 1a). This indicates that the sample preparation process affects neither the protein nor the pigments. In contrast, the maximum is blue-shifted by ~40 nm for the PCP complexes deposited on the silver island film and separated from the metallic nanostructures by a 25-nm-thick SiO₂ layer. After subtracting both curves (Fig. 2b), we find that the difference has a shape of the band with a maximum at 407 nm and linewidth of about 35 nm. We conclude that the observed enhancement is due to plasmon excitations in the metallic layer: this observation is a central result of this work.

The fluorescence excitation spectra displayed in Fig. 2a have not been in any way adjusted or normalized. They are very similar to each other for wavelengths longer than 475 nm, in particular in the absorption range of low-energy



Fig. 2 a Fluorescence excitation spectra measured for the acChl–PCP complexes deposited on the semicontinuous silver film substrate (*red*) compared with that of the reference sample (*green*). The detection wavelength was 690 nm. b Difference between the two spectra displayed in (a)

Per molecules. This suggests that the number of PCP complexes probed in both experiments is almost identical, which makes the estimation of the enhancement factor remarkably straightforward. Importantly, the fluorescence spectrum of the acChl–PCP complexes deposited on the semicontinuous silver film is identical, for all excitation wavelengths, to that of the reference structure, which supports our previous observation that the preparation of the hybrid nanostructure has no measureable effect on the protein or pigment structure.

The time-resolved fluorescence of the two samples is compared in Fig. 3 for excitation wavelengths of (a) 405 nm, (b) 485 nm, and (c) 640 nm. In this way, we probe the fluorescence dynamics when plasmons in the semicontinuous silver film are weakly excited (640-nm excitation) and when they are efficiently excited (405 and 485 nm). The decay kinetics of the control sample on glass are $3.25\pm$ 0.28 ns irrespective of the excitation wavelength (Table 1). A similar value is also obtained for the sample on the silver islands with an excitation at 640 nm. The lifetime is, in contrast, decreased to 2.28 ± 0.2 ns when plasmons in the silver island film are excited. While measurable, this change is relatively small compared with previous results on fluorescent dyes [21] and light-harvesting complexes [8, 9],



Fig. 3 Fluorescence kinetics for the acChl–PCP complexes deposited on the metallic substrate (*red*) compared with the ones obtained for the reference sample on glass (*black*). The excitation wavelengths were 405 nm (a), 485 nm (b), and 640 nm (c)

where order-of-magnitude changes have been measured. The small change of the fluorescence lifetimes in the case of the PCP complexes coupled to the silver island film supports our conclusion that the enhancement measured in the fluorescence excitation (Fig. 1) is predominantly due to the enhancement of the excitation rate in the light-harvesting complex. At the same time, the dependence of the fluorescence decay on the excitation wavelength for the PCP complexes deposited onto the metallic islands suggests that the non-plasmon-related influence of the metallic film upon the studied biomolecule is negligible.

In order to model the observed enhancement of both the absorption and emission of the PCP complexes deposited onto the silver island film, we used the boundary element method [22]. This approach allows for calculating the total

Table 1 Fluorescence decay times measured for acChl-PCP complexes deposited on glass and semicontinuous silver substrates

Excitation wavelength (nm)	405	485	640
acChl-PCP on glass substrate (ns)	3.18	3.02	3.56
acChl-PCP on SIF substrate (ns)	2.28	2.33	3.1
Change (%)	28	23	13

The percentage change for the three excitation wavelengths used in the experiment is given. The error for estimating the values of fluorescence decay times is approximately 0.25 ns

electric field intensity, E_{tot} , on the top surface of the silicon dioxide spacer for a predefined geometry of the metallic nanoparticle layer. This total electric field is a sum of an electric component of incident laser light, E_{inc} , and the field scattered by the silver nanostructures, E_{scatt} . We define the enhancement factor κ as [23, 24]:

$$\kappa \sim rac{<\left|E_{tot}
ight|^2>}{\left|E_{inc}
ight|^2}$$

where $\langle |E_{tot}|^2 \rangle$ corresponds to the average total electric field on the top surface of the silica spacer. We introduce the average value of the electric field for the purpose of estimating the electromagnetic enhancement not only for isotropic (spherical) nanoparticles but also for anisotropic geometries. Indeed, we carried out the analysis for three qualitatively different arrangements of metallic nanoparticles: uncoupled spherical islands, coupled dimers of spherical nanoparticles, and elongated nanoparticles, that is nanorods.

While for a single metallic nanosphere the resonance frequency is independent of the direction, the other two geometries have a reduced symmetry. This leads to two characteristic frequencies; therefore, the calculated enhancement factor must be averaged over two polarizations of incident electric field (along the symmetry axes of the respective metallic nanoparticle arrangement). For all cases, the sizes of silver nanostructures were chosen to approximately match the actual size of the silver islands as measured using the SEM technique [17]. The diameter (width) of spherical (elongated) nanoparticles was in the range of 5–20 nm. The results for all three cases are displayed in Fig. 4, where spectral dependence of κ is shown.

In the case of spherical nanoparticles (Fig. 4a), we obtained a single electric dipolar resonance at $\lambda \approx 400$ nm, which matches well with the spectral dependence of the fluorescence enhancement observed experimentally for the hybrid nanostructure comprising light-harvesting complexes. The spectral position of this resonance is virtually independent of the size of the nanoparticle in the range studied.

For dimers of spherical nanoparticles (Fig. 4b), we observe two effects: firstly, the energies of calculated resonances strongly depend upon the polarization of the



Fig. 4 Enhancement factor κ of electric field intensity calculated on the surface of a 25-nm-thick silica spacer as a function of incident light wavelength for three morphologies of plasmonic structures: isolated spherical nanoparticle (**a**), coupled spherical nanoparticles (**b**), and metallic nanorods (**c**). Calculations for coupled metallic nanoparticles and nanorods were carried out for three cases of separation and length, as indicated. The insets show the distribution of the electric field in the cross section of the structures calculated for the resonance wavelengths

incident light, and secondly, the splitting between the two resonances increases with a larger separation between the two spherical nanoparticles [25, 26]. The first effect was taken into account when estimating the total electromagnetic field enhancement by averaging over the two directions. On the other hand, by gradually increasing the distance between the two nanoparticles forming a dimer from 2 to 10 nm, we induce blue shifts of the resonance frequencies of up to 50 nm, which reflects the weaker coupling of the nanoparticles at longer distances. The results obtained for metallic nanorods (Fig. 4c) are in a way more extreme than those of coupled spherical nanoparticles. We assumed a constant radius of 5.5 nm of the endcaps that were modeled as hemispheres, while the length of the nanorod changed from 13 to 21 nm. In this case, we observe a strong polarization selectivity of the two plasmon resonances, with the lower-energy one yielding much larger enhancement factors and shifting as far as to 700 nm for the longest nanorods. The energy of the strong, red-shifted resonance depends linearly upon the length of the nanorod. On the other hand, the weaker resonance appears around 400 nm and is almost independent of the nanorod length [27].

We now consider the implications of the effects observed for the three geometries of plasmonic nanostructures for understanding the impact of plasmon resonances in the silver island film on the optical properties of the PCP complexes. While the nanorods feature strong resonances in the spectral region comparable to the fluorescence emission range of the PCP complexes, coupling to such structures would result in a strong enhancement of the radiative rate, hence substantial shortening of the fluorescence decay time. In contrast, the enhancement in the 400- to 500-nm spectral range is minute for metallic nanorods. Therefore, we conclude that such geometry is highly unlikely in the studied structure. On the other hand, a moderate, approximately twofold enhancement observed experimentally can be well explained by assuming the presence of isolated and coupled spherical nanoparticles in the silver film structure. Indeed, they exhibit plasmon resonances in the range of 400-450 nm. Furthermore, since the strength of the interaction between metallic nanoparticles is not homogeneous, some distribution of the spectral shifts toward longer wavelengths is expected. These can cause broadening of the enhancement curve observed experimentally for the studied hybrid nanostructure.

Conclusions

We studied the electromagnetic field enhancement in a hybrid nanostructure composed of light-harvesting complexes deposited on a semicontinuous silver film separated by a 25-nm-thick silica spacer. The increase of the acChl– PCP absorption in the spectral range comparable to the plasmon resonances in metallic nanoparticles indicates strong enhancement of absorption due to plasmon interactions. The combination of fluorescence excitation and time-resolved fluorescence spectroscopy allows for separating contributions of absorption and fluorescence rate enhancements. A comparison between the experimental results and calculations for various morphologies indicates that the semicontinuous silver film consists predominantly of isolated islands and/or coupled ones. Acknowledgments Research in Poland has been supported by the WELCOME project "Hybrid Nanostructures as a Stepping Stone towards Efficient Artificial Photosynthesis" funded by the Foundation for Polish Science and EUROCORES project "BOLDCATS" funded by the European Science Foundation. THPB acknowledges support from the US Department of Energy (DE-SC001035), which established the Photosynthetic Antenna Research Center (PARC).

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